

A comment on “Does Forest Loss Increase Human Disease? Evidence from Nigeria”*

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Abstract

[Berazneva and Byker \(2017\)](#) research the effect of deforestation on child disease incidence in Nigeria for 2008 and 2013. They find that the deforestation in the previous year has a positive effect on malaria incidence, but not on diarrhea and cough. Although we are unable to reproduce their results for malaria using the original processed data and adding an extra year to the panel, we also found that there is no consistent relationship between deforestation on diarrhea or cough. Our results also show that the original model is sensitive and overspecification may drive original findings. Future replications could examine heterogeneous effects.

KEYWORDS: Environmental Degradation, Deforestation, Malaria, Infant Mortality, Childrens Health, Nigeria

JEL CODES: I12, I15, J13, O13, O15, Q23

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1 Introduction

[Berazneva and Byker \(2017\)](#) investigate the impact of deforestation on three diseases in Nigerian children under 5: fever (malaria), diarrhea, and respiratory infections (cough). They aimed to expand the literature on the health impacts of environmental degradation by estimating the causal impact of forest loss on infectious disease incidence in young children using temporal and spatial variation in Nigeria using the Demographic and Health Surveys (DHS) for 2008 and 2013.

The authors construct a two-year panel of Nigerias second smallest administrative units (local government areas (LGAs)) using reported disease incidence in the two weeks before the survey, as well as various demographic indicators. They built their main regressor, tree loss, by using a high-resolution geospatial raster dataset of global forest change based on time-series analysis of Landsat images. The authors define forest loss as a change from a forest to non-forest state (indicator). To match the DHS data and the forest loss, the authors create a 5-kilometer buffer zone around each rural DHS cluster and a 2-kilometer buffer zone around each urban cluster. They then calculate the share of pixels within each buffer that experienced forest loss for the year of and up to three years preceding the DHS interview.

They present three different estimations and find that the one with the most demographic and spatial controls provides evidence that the first deforestation lag has a positive and significant impact on malaria, with an increase of 1% causing a 2 percentage point increase in malaria incidence. They do not find any relationship between forest loss and diarrhea or cough incidence. These results demonstrate a causal link between forest loss and malaria and provide information on the timing of exposure to malaria relative to tree loss, motivating future work on the longer-term consequences of forest loss in Nigeria and elsewhere on the continent.

In the present paper, we investigate whether [Berazneva and Byker's \(2017\)](#) main results are reproducible and perform additional robustness checks. Although we did our best to follow the authors' instructions to process the original data, we are unable to reproduce the results for malaria incidence in magnitude or significance. We also find that there is no relationship between forest loss and diarrhea or cough. We successfully reproduce Table 1 from [Berazneva and Byker \(2017\)](#) using their analysis data, but we are not able to replicate processing data from the original source.

We uncover one minor coding error in which the authors accidentally use incorrect values to de-normalize survey weights when constructing their panel. Specifically, they use incorrect figures for the population of women in reproductive age (WRA) and for the surveyed number of individuals in this group in both years. Correcting this error does not significantly alter any of the main results based on

their analysis data. Nevertheless, we use the correct de-normalized weights in our robustness tests.

We attempt to replicate the original results and conduct two robustness tests to evaluate the sensitivity of the papers main findings. Specifically, the original study uses the 2008 and 2013 waves of the Nigerian DHS. We (1) reproduce the results using the original data from the Nigerian DHS program (processed data), (2) expand the dataset by including an additional survey wave from 2018 (extended data), and (3) estimate a model with a reduced set of control variables to limit the risk of over-specification.

The results of our replication differ from the original study in direction, significance, and magnitude. The authors instructions did not allow us to fully reproduce their analysis dataset from the original source. Although we followed their procedures, discrepancies between their analysis data and our processed raw data yield different estimates for the effect of deforestation. Specifically, we find that the effect is present in the current year rather than in the first-year lag, as reported in the original paper. Additionally, our estimates vary in magnitude across lags: the contemporaneous effect is larger and decreases over time (excluding the third lag), suggesting that deforestation may contribute to a marginal reduction in malaria incidence. The main source of the discrepancy appears to be the distribution of forest loss: while Nigeria shows a clear pattern of deforestation concentrated in the south, the authors data indicate forest loss distributed across the entire country.

When we omit different variables to prevent a model overspecification caused by multicollinearity between spatial and social demographic data, we find that deforestation has no effect in any of the diseases explored, showing the sensitivity of the original model. This is found using the author’s analysis data, as well as our processed and extended data.

Finally, although we are unable to reproduce the original results, future replication studies could explore heterogeneity analysis by focusing on LGAs with substantial forest cover, as tree cover needs to be relatively significant to provide protection against diseases ([Laporta et al. 2021](#)). In addition, future analyses could restrict the sample to poorer and/or rural populations to account for the greater vulnerability of these groups and to assess whether deforestation has heterogeneous effects on disease outcomes ([Bauhoff and Busch 2020](#)).

2 Computational Reproducibility

[Berazneva and Byker \(2017\)](#) provide a replication package with the final dataset, code, and documentation of the data preparation process. We follow this documentation closely, reconstructing the dataset from the original sources.

We extend the analysis by adding an additional survey wave and an updated version of the forest loss dataset. The replication relies on four main sources: the 2008 and 2013 Nigerian Demographic and Health Surveys (DHS), High-Resolution Global Maps of 21st-Century Forest Cover Change, the DMSP-OLS Nighttime Lights Time Series, and georeferenced soil data.

In what follows, we describe the geolinking of these sources and the construction of the panel at the level of Nigerian local government areas (LGA).

2.1 Data Replication

Demographic and Health Surveys (DHS) The analysis is based on health data from the Nigeria Demographic and Health Surveys (DHS), nationally representative cross-sectional, geo-referenced surveys for 2008 and 2013 (NPC and ICF, 2009, 2014). Access to the data is granted following a brief request. Mapping the variables used in this analysis to the survey was relatively straightforward, although identifying and incorporating control variables presented more of a challenge. We were able to retrieve all relevant characteristics directly from the Childrens Recode dataset and did not require any information from the Household Recode. The Childrens Recode data were further extended with geographic information on cluster coordinates, allowing for geo-enrichment such as altitude data. The number of observations and mean disease incidence were fairly consistent across survey rounds. For robustness and replication with updated data, we extend the analysis to include the 2018 DHS survey (NPC and ICF, 2019).

Geolinking To link environmental data with health outcomes, we spatially matched DHS survey clusters to multiple geospatial datasets. Each cluster, defined as the centroid of census-based enumeration areas, is subject to random coordinate displacement by DHS to protect confidentiality: rural clusters are displaced 05 km (with 1% displaced up to 10 km), and urban clusters 02 km, in a random direction and distance (Perez-Haydrich et al. 2013). To account for this, we constructed 5 km buffers for rural clusters and 2 km buffers for urban clusters. Within each buffer, we extracted raster values for soil characteristics, forest loss, and luminosity. Administrative boundaries at the LGA level are taken from the map of Subnational Administrative Boundaries for Nigeria provided by the Humanitarian Data Exchange (HDX), published by the UN Office for the Coordination of Humanitarian Affairs (OCHA) country office in Nigeria. All datasets were reprojected from EPSG:4326 (geographic coordinates, WGS84) to EPSG:3857 (projected coordinates in meters) to ensure accurate distance-based calculations. For each cluster, raster values were averaged, and results were subsequently aggregated by computing means

across clusters within each LGA. This procedure provides a consistent framework for enriching DHS microdata with spatially referenced environmental indicators.

Deforestation Data The regressor of interest, tree loss, is drawn from the high-resolution Global Forest Change dataset, which is based on time-series analysis of Landsat imagery at a spatial resolution of one arc-second (approximately 30 meters at the equator), with annual allocation of forest loss (Hansen et al. 2013, version 1.2).

Forest loss is defined as a transition from forest to non-forest (stand-replacement disturbance). The analysis uses the year of gross forest cover loss event dataset, which disaggregates total forest loss to annual scales. Values are encoded as 0 (no loss) or 114, corresponding to loss detected in 20012014, respectively. This measure captures gross forest loss, so forest gain in the same year is not excluded. In addition, the authors use the tree canopy cover for the year 2000 dataset, defined as canopy closure for vegetation taller than 5 m. Values are encoded as percentages per output grid cell, ranging from 0 to 100.

The forest data are distributed across four large GeoTIFF files organized into tiles. This structure required scripts to identify whether a DHS cluster polygon was located within a single tile or spanned multiple tiles. We extract all pixel values across the relevant tiles and compute the mean proportion of forest loss for each cluster in a given year.

Figure 1 compares the replicated dataset with that of the authors. We find substantial discrepancies, notably because most forest loss occurred in southern Nigeria, whereas only limited loss is observed in the north. This raises concerns about the handling of the forest loss data in the original study.

For robustness and replication with updated data, we extend the analysis to include the 2018 DHS survey. To match this period, we use the Global Forest Change 20002019 dataset (Hansen et al. 2013, version 1.7) to update the regressor.

Soil Data The authors also make use of soil fertility indicators, namely Soil Organic Carbon (*SOC*), Soil pH (measured in H_2O), and Cation Exchange Capacity (*CEC*). The underlying data are provided as raster files that map soil properties across Africa at a 250 m resolution and for different depth intervals (Hengl et al. 2015). For each DHS cluster, we extract soil property values by averaging all raster cell values within a 5 km buffer for rural locations and a 2 km buffer for urban locations. We then compute the mean across all DHS clusters within a given LGA.

Soil values are retrieved at depths of 05 cm and 515 cm, and subsequently averaged to obtain values for the 015 cm layer. We do not observe any statistically significant differences in means or distributions when applying either a two-sided

t-test or the KolmogorovSmirnov (KS) test. Summary statistics are presented in Table 1.

Luminosity Data Finally, the authors use nighttime lights data as a proxy for economic activity (Chen and Nordhaus 2011). The data, capturing persistent lighting from cities, towns, and other sites, are obtained from the National Oceanic and Atmospheric Administration National Geophysical Data Center (US Air Force Weather Agency 2009).

Satellite F16 provides coverage for 2005-2008, and F18 for 2009-2013. We use Version 4 of the DMSP-OLS Nighttime Lights Time Series, constructed in smoothed spatial resolution mode at 30 arc-seconds, which corresponds to approximately 1,000 meters (1 km) at the equator.

For each DHS cluster, we retrieve values by averaging all raster cells within the respective buffer zone, and then take the average across clusters within a given LGA. We do not find significant differences in means or distributions using either a two-sided t-test or the KolmogorovSmirnov (KS) test. The descriptive statistics are reported in Table 1.

2.2 Discrepancies Between Analysis and Raw Data

For the summary statistics, we followed the authors methodology by identifying LGAs containing at least one cluster per DHS wave (2008 and 2013) and subsetting the data accordingly. We then merged the variables into a panel dataset for comparison.

Our replication dataset contains slightly more observations, likely due to differences in how survey clusters are geolinked to administrative boundaries. The main discrepancies between our data and the authors arise from variations in spatial linkage procedures and spatial data preprocessing. In particular, the authors accidentally included 40 observations corresponding to LGAs with missing values, which we exclude in our replication.

Table 1 reports test statistics (two-sided t -tests and KolmogorovSmirnov tests) comparing the dependent variables from the original and replication datasets. For survey-based variables, we fail to reject the null hypothesis of equal means and distributions, indicating no significant differences between the datasets. In contrast, for spatial variables, both means and distributions differ significantly, with discrepancies especially pronounced in the forest loss measures (see Figure ??).

While the original authors identify LGAs with at least one cluster per DHS wave, their code counts 40 observations without LGA identifiers as valid cases. In our replication, these missing values are excluded to ensure consistency. Minor differences also arise in the spatial join between DHS cluster locations and the Nigeria

LGA shapefile: the original dataset covers 409 LGAs, whereas our cleaned dataset includes 408, reflecting the exclusion of clusters with missing spatial information. Importantly, cluster locations coincide across both datasets, indicating alignment in the underlying DHS coordinates. For reference, Nigeria contains a total of 774 LGAs.

2.3 Regression Model

For our analysis, we rely on the same specifications as those used in the paper. We first estimate a balanced pooled OLS for each disease incidence in young children reported by each surveyed cluster (unit level) in lags for forest loss (3), then add fixed effects of LGA and year and the time-varying controlled variables selected by the authors. See the original study for additional details. We estimate the following equation:

$$Y_{itc} = \alpha_0 + \sum_{j=0}^3 \beta_j loss_{tc}^j + \mathbf{X}'_{itc} \gamma + \mathbf{LGA}'_c \pi + \mu' month_{itc} + \mathbf{DHSyear}'_t \theta \quad (1)$$

$$+ \sum_{m=1}^{12} \lambda_m month_{tc}^m \times Region_c \times DHSyear_t + \epsilon_{itc},$$

2.4 Results

We start by using the newly cleaned raw data (see Section 2.1) and correcting the de-normalized weights using the procedure used by the authors but with corrected survey and population values for women in reproductive age (15-49) (see Appendix Section 6.1). Column 1 in Table 3 shows the original study estimates for their most complete specification, and columns 2 and 3 present the results of our robustness tests, which include the use of processed raw data and the addition of an extra year of analysis (2018). Columns 4 to 6 focus on preventing overspecification in the original dataset, as well as in the processed raw and extended data (see Table 4 and Table 5 for the results for diarrhea and cough).

When using the processed raw data (column 2), we find that the point estimate for fever, which is the key variable in the original study, is significant in the current year rather than in the first-year lag. This suggests that deforestation in the current year is associated with a marginal increase in malaria incidence of 2.954 percentage points. When including the additional survey wave from 2018 (column 3), the contemporaneous effect increases and remains positive, while the lagged effects are smaller and generally insignificant.

Since the authors include several variables in their most complete model, we estimate the effect of deforestation by omitting variables that may be correlated

with deforestation or with sociodemographic characteristics. When applying this specification to the original analysis data, coefficients remain at a similar level of significance, although their magnitude decreases (column 4). Using the same variables with the processed raw data, we find no significant relationship (column 5). For the extended dataset, the results are very similar (column 6), with the contemporaneous effect of deforestation showing a marginal positive effect on malaria incidence.

Our replication results for the relationship between forest loss and malaria, using the processed raw and extended datasets, differ from the original study in direction, significance, and magnitude. Although some differences may reflect improvements in data collection and updates to key inputs since the original publication, our findings suggest that the effect of deforestation on malaria incidence is not clearly evident. We also find no consistent evidence of a relationship between deforestation and increases in the incidence of diarrhea or cough. While the original paper reported a large impact of forest loss on malaria, with a dynamic pattern consistent with a temporary ecological disturbance described in the tropical medicine literature, our replication does not confirm this result, and we do not find clear causal evidence supporting this link.

2.4.1 Replication with Processed Raw Data As discussed in Section 2.1, the instructions provided by the authors did not allow us to fully reproduce their analysis dataset from the original source. Although we followed their procedures, discrepancies between their analysis data and our processed raw data yield different estimates for the effect of deforestation (see columns 12 in Table 3). First, we find that the effect is present in the current year rather than in the first-year lag, as reported in the original paper. Second, our estimates vary in magnitude across lags: the contemporaneous effect is larger and decreases over time (excluding the third lag), suggesting that deforestation may contribute to a marginal reduction in malaria fever. The primary source of the discrepancy appears to be the distribution of forest loss in the datasets. While Nigeria exhibits a clear pattern of deforestation concentrated in the south, the authors data indicate forest loss distributed across the entire country (see Figure 1).

2.4.2 Extending the Time Period In 2018, the DHS conducted a new survey wave in the country, allowing us to expand our sample and examine how including an additional year affects the sign, magnitude, and statistical significance of deforestation on childrens disease incidence. We incorporated updated data on forest loss and calculated forest loss lags using the same procedure as the original authors (see Section 2.1). Estimating Equation 1 with the extended dataset, we find that

the impact of deforestation is significant in the current year and larger than using previous waves, reflecting an increase in disease incidence (see Table 1). Coefficients remain positive across lags, however, including the additional year does not improve the significance of the results for diarrhea or respiratory infections (cough), which remain largely insignificant.

2.4.3 Prevent for Overspecification Our final robustness check consists of revising the variables included in the original model’s most complete specification (Columns 3, 6, and 9 in Table 1 of the original paper) to ensure that the authors’ results, as well as our own, are robust. We exclude interaction variables already present in the model (region \times month \times year) and decide not to use the variables related to soil or luminosity, as they are very likely to be collinear with deforestation [Angelsen et al. \(2014\)](#), [Burgess et al. \(2012\)](#). For the demographic variables, we only include those focused on the children’s characteristics, the household head’s education, and socioeconomic status, avoiding those that could be collinear with each other. Table 6 in the Appendix shows the variables included in this robustness check to avoid overspecification and multicollinearity [Clarke \(2005\)](#).

Table 3 presents results for the analysis data, processed raw data, and extended dataset (columns 4-6). The original model is sensitive to the omission of potentially collinear variables, though some coefficients gain significance, nonetheless, we find no meaningful effects of deforestation on malaria. In contrast, results from our robustness checks are consistent and less sensitive to variable selection. Using the replication dataset, the contemporaneous effect of deforestation on malaria is positive and marginally significant at the 10-percent level, with a 1 percent loss of forest cover in the current year associated with a three percentage point increase in malaria incidence. R-squared values remain similar across fever (malaria), diarrhea, and respiratory infections (cough). Overall, we cannot identify a general relationship between deforestation and changes in childrens incidence of malaria, diarrhea, or respiratory infections.

3 Conclusion

We replicate the results of [Berazneva and Byker \(2017\)](#) and compare them with those obtained using the processed raw data and an extended dataset that includes an additional survey wave from 2018. We are unable to reproduce the original results for malaria, as our estimates show sensitivity in the current year rather than in the first-year lag, as reported by the authors. When controlling for potential overspecification, we find that deforestation has no significant effect on malaria, indicating that the original model is highly sensitive to changes in specification.

We also find no consistent evidence of a relationship between deforestation and increases in the incidence of diarrhea or cough. While the original paper reported a large impact of forest loss on malaria, with a dynamic pattern consistent with a temporary ecological disturbance described in the tropical medicine literature, our replication does not fully confirm this result, and we find no clear causal evidence supporting this link.

4 Figures



Figure 1: Geographic Variation of Forest Loss in Nigeria by LGA (2001-2012)

Panel (a) displays the estimates from [Berazneva and Byker \(2017\)](#) for the average total tree cover loss between 2001 and 2012 across the 409 LGAs included in their analysis sample. Each LGA is shaded according to the average level of tree cover loss observed across the surveyed clusters. LGAs not observed in either the 2008 or 2013 DHS waves are left unshaded. Panel (b) presents our replication, where we estimate for each cluster the share of forest loss between 2001 and 2012 and then compute the average by LGA.

5 Tables

Table 1: Descriptive Statistics Comparison Original and Replication Datasets

Variable	Replication Dataset			Original Dataset			Test Statistics	
	Observations	Mean	Std	Observations	Mean	Std	t-stat	KS-stat
Malaria (fever)	41,737	0.144	0.352	41,409	0.144	0.351	0.135	0.000
Diarrhea	41,786	0.106	0.308	41,458	0.106	0.307	0.173	0.000
Respiratory (cough)	41,682	0.110	0.312	41,354	0.110	0.313	-0.111	0.000
Forest loss								
This year	47,039	0.001	0.002	46,654	0.001	0.004	-17.524***	0.047***
1 year ago	47,039	0.001	0.002	46,654	0.001	0.003	-13.245***	0.052***
2 years ago	47,039	0.001	0.002	46,654	0.001	0.003	-16.803***	0.053***
3 years ago	47,039	0.000	0.001	46,654	0.001	0.002	-17.412***	0.061***
Luminosity								
This year	47,039	-0.098	1.193	46,654	0.087	1.492	-20.860***	0.185***
1 year ago	47,039	0.356	1.362	46,654	0.317	1.565	4.049***	0.113***
2 years ago	47,039	-0.274	1.989	46,654	-0.392	2.436	8.143***	0.085***
Soil Organic Carbon	47,039	12.712	7.512	46,654	12.936	7.879	-4.453***	0.036***
Soil pH	47,039	59.876	3.698	46,654	59.801	3.855	3.010***	0.038***
Cation Exchange Capacity	47,039	10.454	3.905	46,654	10.472	4.236	-0.665	0.058***
Tree cover in 2000	47,039	7.883	9.989	46,654	10.820	13.360	-38.089***	0.113***

The table presents descriptive statistics for the original and replication datasets, restricted to observations included in both the 2008 and 2013 DHS survey waves. Differences in distributions are evaluated using two-sided t-tests and KolmogorovSmirnov tests. Statistical significance is indicated by $p < 0.10$ (*), $p < 0.05$ (**), and $p < 0.01$ (***).

Table 2: Summary Statistics of child Illnesses by DHS Year

DHSYEAR	Malaria (Fever)			Diarrhea			Respiratory (Cough)		
	Observations	Mean	Std	Observations	Mean	Std	Observations	Mean	Std
2008	16,892	0.1578	0.3646	16,910	0.0994	0.2992	16,874	0.1197	0.3246
2013	22,237	0.1348	0.3415	22,260	0.1090	0.3117	22,200	0.1047	0.3062
2018	18,286	0.2393	0.4267	18,287	0.1279	0.3340	18,292	0.1687	0.3745
Total	57,415	0.1749	0.3799	57,457	0.1122	0.3156	57,366	0.1295	0.3357

The table presents descriptive statistics for the main dependent variables, malaria (fever), diarrhea, and respiratory illness (cough), using the original and replication datasets, restricted to observations included in the 2008, 2013, and 2018 DHS survey waves.

Table 3: Replication Results for Malaria (Fever)

Malaria (Fever)	Full Specification			Overspecification Correction		
	(1)	(2)	(3)	(4)	(5)	(6)
Forest Loss						
This year	0.466 (1.027)	2.954* (1.621)	3.344* (1.642)	0.422 (1.003)	2.823 (1.605)	3.280* (1.627)
1 year ago	2.042* (0.960)	1.091 (1.507)	1.042 (1.492)	1.979* (0.961)	1.331 (1.510)	1.246 (1.497)
2 years ago	0.954 (0.950)	-0.117 (2.319)	0.148 (2.342)	1.096 (0.985)	-0.044 (2.270)	0.088 (2.315)
3 years ago	-3.434 (2.758)	1.554 (3.701)	1.499 (3.696)	-3.097 (2.628)	0.805 (3.666)	0.858 (3.662)
Constant	0.122 (0.343)	0.255** (0.089)	0.235** (0.088)	0.136 (0.138)	0.174** (0.055)	0.166** (0.055)
Observations	40,675	40,459	37,885	40,880	40,620	38,038
R-squared	0.082	0.081	0.082	0.080	0.080	0.081

Notes: The table reports estimation results for fever (malaria) using the full specification from the original study in columns (1)(3), where the dependent variable is the reported disease incidence among children under five. Column (1) reproduces the original results, while columns (2) and (3) present estimates using the processed raw data and the extended dataset including the 2018 DHS wave, respectively. Columns (4)(6) show results from a reduced model estimated to address potential overspecification. Column (4) uses the original analysis data, and columns (5) and (6) use the processed raw data and the extended dataset, respectively. The variables excluded to prevent overspecification and multicollinearity are listed in Table 6 in the Appendix (Clarke 2005). Robust standard errors clustered at the LGA level are reported in parentheses. Statistical significance is denoted by $p < 0.10$ (*), $p < 0.05$ (**), and $p < 0.01$ (***)

Table 4: Replication Results for Diarrhea

Diarrhea	Full Specification			Overspecification Correction		
	(1)	(2)	(3)	(4)	(5)	(6)
Forest Loss						
This year	-0.608 (0.591)	-1.182 (1.068)	-1.259 (1.056)	-0.695 (0.576)	-1.381 (1.027)	-1.399 (1.016)
1 year ago	0.633 (0.617)	2.829* (1.236)	2.674* (1.235)	0.620 (0.600)	2.787* (1.207)	2.689* (1.207)
2 years ago	0.244 (0.942)	2.258 (2.622)	2.168 (2.441)	0.394 (0.964)	2.775 (2.468)	2.603 (2.318)
3 years ago	0.067 (1.131)	1.761 (1.539)	1.784 (1.516)	0.406 (1.062)	1.324 (1.556)	1.456 (1.537)
Constant	-0.183 (0.234)	0.157* (0.070)	0.166* (0.072)	0.186** (0.064)	0.003 (0.026)	0.004 (0.027)
Observations	40,715	40,508	37,926	40,922	40,669	38,079
R-squared	0.084	0.084	0.085	0.082	0.082	0.084

Notes: The table reports estimation results for diarrhea using the full specification from the original study in columns (1)(3), where the dependent variable is the reported disease incidence among children under five. Column (1) reproduces the original results, while columns (2) and (3) present estimates using the processed raw data and the extended dataset including the 2018 DHS wave, respectively. Columns (4)(6) show results from a reduced model estimated to address potential overspecification. Column (4) uses the original analysis data, and columns (5) and (6) use the processed raw data and the extended dataset, respectively. The variables excluded to prevent overspecification and multicollinearity are listed in Table 6 in the Appendix (Clarke 2005). Robust standard errors clustered at the LGA level are reported in parentheses. Statistical significance is denoted by $p < 0.10$ (*), $p < 0.05$ (**), and $p < 0.01$ (***)

Table 5: Replication Results for Respiratory (Cough)

Respiratory (Cough)	Full Specification			Overspecification Correction		
	(1)	(2)	(3)	(4)	(5)	(6)
Forest Loss						
This year	-1.161 (1.123)	-0.818 (1.760)	-0.787 (1.800)	-0.992 (1.089)	-0.111 (1.703)	0.019 (1.744)
1 year ago	0.425 (0.782)	-0.002 (1.466)	-0.093 (1.464)	0.582 (0.774)	0.613 (1.497)	0.489 (1.495)
2 years ago	-1.085 (0.830)	-4.479* (2.161)	-4.814* (2.180)	-1.329 (0.840)	-5.565** (2.126)	-5.992** (2.152)
3 years ago	0.574 (2.271)	3.160 (3.261)	3.545 (3.274)	0.417 (2.157)	2.602 (3.237)	3.064 (3.253)
Costant	0.081 (0.238)	0.164* (0.079)	0.146 (0.080)	0.286*** (0.083)	0.141** (0.054)	0.136* (0.054)
Observations	40,618	40,407	37,833	40,823	40,568	37,986
R-squared	0.093	0.092	0.093	0.091	0.090	0.091

Notes: The table reports estimation results for respiratory (cough) using the full specification from the original study in columns (1)(3), where the dependent variable is the reported disease incidence among children under five. Column (1) reproduces the original results, while columns (2) and (3) present estimates using the processed raw data and the extended dataset including the 2018 DHS wave, respectively. Columns (4)(6) show results from a reduced model estimated to address potential overspecification. Column (4) uses the original analysis data, and columns (5) and (6) use the processed raw data and the extended dataset, respectively. The variables excluded to prevent overspecification and multicollinearity are listed in Table 6 in the Appendix (Clarke 2005). Robust standard errors clustered at the LGA level are reported in parentheses. Statistical significance is denoted by $p < 0.10$ (*), $p < 0.05$ (**), and $p < 0.01$ (***)

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6 APPENDIX

6.1 Correct weight de-normalization

The authors recommend de-normalizing survey weights when using multiple waves of DHS data, which is a good practice. However, the authors use incorrect population figures for women of reproductive age (WRA), which is between 15 and 49 years old. Furthermore, the links cited for these figures are no longer active. We have used the correct values for 2008 (35,882,027), 2013 (41,018,918), and 2018 (47,146,386) based on data from the [United Nations \(2025\)](#). Furthermore, the authors also used an incorrect value for WRA in the survey itself. While this error does not lead to significantly different point estimates, it can produce biased estimators with wider standard errors. The details of the correction are in the do files in the repository. We used the equation proposed by the authors:

$$NewWeight_{year} = \frac{PopWRA_{year}^{weight}}{100,000} \times \frac{PopWRA_{year}^{real}}{PopWRA_{year}^{survey}}$$

6.2 Variables used in each specification

Table 6: Model Comparison

Variable	Description	Full Specification	Overspecification Correction	Reason for Exclusion
forest_loss_0	Present deforestation	Yes	Yes	
orest_loss_1-forest_loss_3	Deforestation 1st, 2nd and 3rd lag	Yes	Yes	
i.DHSYEAR	Year fixed effects	Yes	Yes	
i.LGA	LGA fixed effects	Yes	Yes	
i.month#i.region#i.DHSYEAR	Month X region X year fixed effects	Yes	No	Correlated with LGA and Month fixed effects
i.month	Month fixed effects	Yes	No	
i.month#i.region	Month X region fixed effects	No	Yes	
treecover_2000	Tree cover mean in 2000 (%)	Yes	Yes	
cec_ave_pt	Soil CEC 2.5 & 10cm ave	Yes	No	
ph_ave_pt	Soil pH 2.5 & 10cm ave	Yes	No	Correlation with deforestation
oce_ave_pt	Soil organic carbon 2.5 & 10cm ave	Yes	No	
luminosity_chg_0_-2	Change in luminosity coverage	Yes	No	
altitude	Altitude (m)	Yes	Yes	
no_HH_members	# of household members (hv009)	Yes	Yes	
no_kids_under_5	# of children under 5 in the HH (hv014)	Yes	Yes	
time_to_water	Minutes to nearest water source (hv204)	Yes	Yes	
HH_head_edu_years	Years of education by HH head (hv108_01)	Yes	Yes	
head_HH_age	Age of household head in years (hv220)	Yes	No	Not relevant
toilet	1 HH has a flush toilet, 0 if no (hv205)	Yes	No	
firewood	1 if source of fuel is firewood, straw, 0 if other (hv226)	Yes	No	Correlated with rural and poverty
floor	1 if floor made of earth/sand/dung, 0 if other (hv213)	Yes	No	
rural	1 if rural, 0 if urban (hv025)	Yes	Yes	
poorest	1 if in poorest wealth quintile, 0 if richer (hv270)	Yes	Yes	
age	Child's current age(b8)	Yes	Yes	
age_resp	Age of respondent in years (v012)	Yes	No	Not relevant
edu_years	Respondent years of education (v133)	Yes	No	
no_child_total	# of children ever born (v201)	Yes	No	Correlated with rural and poverty
no_child_living	# of living children (v218)	Yes	No	
resp_slept_net	1 if respondent slept under bednet, 0 if no (v461)	Yes	No	
resp_works	1 if respondent works, 0 if no (v714)	Yes	No	
livewith	# of children respondent lives with (v202 + v203)	Yes	Yes	
christian	1 if respondent is Christian, 0 if not (v130)	Yes	Yes	
muslim	1 if respondent is Muslim, 0 if not (v130)	Yes	Yes	
yoruba	1 if respondent ethnicity is Yoruba, 0 if not (v131)	Yes	Yes	
igbo	1 if respondent ethnicity is Igbo, 0 if not (v131)	Yes	Yes	
hausa	1 if respondent ethnicity is Hausa, 0 if not (v131)	Yes	Yes	
married	1 if respondent is married, 0 if otherwise (v501)	Yes	Yes	
pregnant	1 if respondent is pregnant at interview time, 0 if not (v213)	Yes	Yes	

Notes: This table lists the variables excluded from the full specification of Equation 1 to address potential overspecification caused by multicollinearity between spatial and sociodemographic variables. The final column provides the reason for each exclusion.

7 Appendix Tables

Table 7: Replication Package Contents and Reproducibility

Replication Package Item	Fully	Partial	No
Raw data provided			✓
Analysis data provided	✓		
Cleaning code provided			✓
Analysis code provided	✓		
Reproducible from raw data		✓	
Reproducible from analysis data	✓		

Notes: This table summarizes the replication package contents contained in [Berazneva and Byker \(2017\)](#).