Title: Disparities in availability of essential medicines to treat non-communicable diseases in Uganda: a Poisson analysis using the Service Availability and Readiness Assessment

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Abstract

Objective Although the WHO-developed Service Availability and Readiness Assessment (SARA) tool is a comprehensive and widely applied survey of health facility preparedness, SARA data have not previously been used to model predictors of readiness. We sought to demonstrate that SARA data can be used to model availability of essential medicines for treating non-communicable diseases (EM-NCD).

Methods We fit a Poisson regression model using 2013 SARA data from 196 Ugandan health facilities. The outcome was total number of different EM-NCD available. Basic amenities, equipment, region, health facility type, managing authority, NCD diagnostic capacity, and range of HIV services were tested as predictor variables.

Findings In multivariate models, we found significant associations between EM-NCD availability and region, managing authority, facility type, and range of HIV services. For-profit facilities’ EM-NCD counts were 98% higher than public facilities (p<.001). General hospitals and referral health centers had 98% (p=.004) and 105% (p=.002) higher counts compared to primary health centers. Facilities in the North and East had significantly lower counts than those in the capital region (p=0.015; p=0.003). Offering HIV care was associated with 35% lower EM-NCD counts (p=0.006). Offering HIV counseling and testing was associated with 57% higher counts (p=0.048).

Conclusion We identified multiple within-country disparities in availability of EM-NCD in Uganda. Our findings can be used to identify gaps and guide distribution of limited resources. While the primary purpose of SARA is to assess and monitor health services readiness, we show that it can also be an important resource for answering complex research and policy questions requiring multivariate analysis.
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Authors Contributions  SK and JIS developed the concept for the study. MAH led the data analysis, literature review, writing, and design of tables and figures. SK and JIS led the data interpretation and contributed significantly to the literature review and writing. JIS led the final editing of the manuscript. SB and GM facilitated access to the data set used in the analysis. All authors contributed to reviewing and finalizing the manuscript.

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Conflicts of Interest  The authors do not declare any conflicts of interest related to this manuscript.

Ethics Committee Approval  Given the nature of this secondary analysis of health facility-level data, no ethics committee approval was required.
Introduction

Background/Rationale

The World Health Organization (WHO) defines Essential Medicines (EM) as drugs considered critical to meeting the needs of the population and expects them to be accessible. To qualify as accessible, drugs must be available and affordable.[1] Yet EM used to treat non-communicable diseases (EM-NCD) remain poorly accessible to the populations of low- and middle-income countries (LMIC)[2-5], where non-communicable diseases (NCD) such as cardiovascular disease, diabetes, chronic lung disease, and mental health disorders are the leading causes of mortality. [1,6-8]

WHO has called for an 80% availability target for EM-NCD as part of a Global Action Plan, making EM-NCD a global priority.[9] However, aggregate estimates of availability at the country level may disguise stark disparities. To our understanding, the extent to which disparities for EM-NCD availability exist within individual LMIC has not previously been studied.

We sought to develop a scalable strategy for identifying within-country availability disparities from routinely collected data that could be compared across multiple LMIC. The WHO Service Availability and Readiness Assessment (SARA) is a widely endorsed methodology used to collect health facility-level data on essential medicines, technologies, and human resources.[10] This comprehensive survey of health system preparedness is intended to be performed annually and provides a national sampling of drug availability, among other indicators. At the time of publication, 11 LMIC have conducted 17 SARA surveys.[10,11] Data from SARA surveys have
been used in country reports and published articles, but these have relied solely on descriptive
statistics.[12-15]

In this analysis, we use SARA data to model internal disparities in the availability of EM-NCD
in Uganda. Our objective was to model meaningful associations between EM-NCD availability
and facility-level characteristics in a sample of Ugandan health facilities. While the primary
purpose of SARA is to assess and monitor health services readiness rather than produce ready-to-
analyze data for research, we show that SARA can also be an important resource for answering
more complex research and policy questions using statistical methods.

Methods

Study Design and Setting

In 2013, the Ugandan Ministry of Health used the WHO SARA methodology to survey 209
health facilities in 10 districts. Healthcare in Uganda, a low-income country with a growing NCD
burden[16], is delivered in three sectors: public, private-not-for-profit (PNFP), and private-for-
profit (PFP). Within each sector, health facilities are divided into levels. These include health
center (HC) I, II, III, IV, general hospital, and regional/national referral hospital. Each facility
type varies by population served, functionality, and leadership. The HC-I level represents the
community health worker program rather than facility-based services, and thus is not included in
the SARA sampling.
In 2013, the Ugandan Ministry of Health, with support from WHO Country Office-Uganda, systematically sampled from facilities across these layers to conduct the SARA survey. Survey personnel visited a stratified sample of 209 Ugandan health facilities across 10 districts over a two-week period. Each health facility was assessed in one day. The presence of each medicine, equipment, or other supply was visually confirmed by the surveyor.

Exclusions
While the complete SARA dataset for Uganda includes 13 national and regional referral hospitals, we excluded these facilities from our analysis. These referral facilities were sampled from outside the 10-district geographic frame of the other 196 facilities, which posed problems for modeling several predictor variables of interest. After excluding the referral hospitals, 196 facilities remained, including HC-II, HC-III, HC-IV, and general hospitals.

Outcome Variable
The 2013 Uganda SARA collected availability data on 20 EM, called “tracer medicines.” We identified 10 of these tracer medicines as EM-NCD. All but one of these, simvastatin, also appear on the Uganda Essential Medicines List (EML), which designates the lowest-level health facility at which each medicine is expected to be stocked (Table 1). The outcome variable, EM-NCD availability, was measured as a count score of these medicines ranging from 0 to 10. The score represents how many of the ten EM-NCD a particular facility had in stock on the day of the SARA survey.

Independent Variables
The independent variables of interest include geographic location, facility characteristics and the presence of other services or equipment. The basic amenities domain score for each facility is the proportion of the list of basic amenities available at a given site. The basic amenities included in the domain score were a consultation room, adequate sanitation facilities, emergency transportation, improved water source, communication equipment, power, and a computer with internet and email. Similarly, the basic equipment domain score is a proportion on the list of basic equipment available at a given facility. The basic equipment included in the domain score were as follows: adult scale, child scale, thermometer, stethoscope, blood pressure apparatus, and light source. Finally, NCD diagnostic capacity is a simple count of facility capabilities using the following tracer items: hemoglobin, blood glucose, urine dipstick (protein), urine dipstick (glucose), urine pregnancy test, and dried blood spot (DBS) collection.

If the facility offered HIV counseling and testing at the time of the survey, it was coded 1 for HIV counseling and testing (HCT). If counseling and testing were not available, the facility was coded 0. Similarly, if the facility offered HIV care and support services at the time of the survey, it was coded 1 for HIV care and support services. If HIV care and support services were not available, the facility was coded 0.

We divided Uganda into four commonly accepted regions: West, North, East, and South. The South region includes Kampala, the capital city. We then assigned each facility to a region according to its recorded district in the SARA dataset. Because Kampala is generally acknowledged to have the greatest concentration of medical resources, we used the South region was used as the reference region.
Finally, each facility in the SARA data is identified by its managing authority, or sector. These include public, PNFP, or PFP, as defined above. In the current analysis, public facilities are the reference category to which PNFP and PFP facilities are compared. The remaining facilities were coded as HC-II, HC-III, HC-IV or General Hospital. HC-IVs offer the most services outside hospitals, while HC-II facilities offer the fewest services.

**Analysis**

We fit a series of Poisson regression models using the GENMOD procedure in SAS University Edition (SAS Institute, Inc.). Beginning with a baseline model predicting NCD score by basic amenities domain score, we added independent variables hypothesized to associate with NCD score in a stepwise fashion. With the addition of each new independent variable, we assessed whether model fit was improved relative to the increased number of parameters using the Akaike information criterion (AIC). If model fit improved with the addition of a variable, we retained the variable and added the next one. Using this forward selection strategy, we reached a full “saturated” model. We then used backward elimination to remove independent variables with non-significant parameter estimates, limited contribution to model fit, or limited clinical significance until we reached our final model. All analyses were scaled to correct for over-dispersion.

To account for SARA’s complex sampling design, we weighted all our analyses using the WEIGHT option in the SAS GENMOD procedure and the sampling weights provided in the SARA dataset. Once we reached the final model, we performed diagnostics for fit and robustness.
with particular attention to the possibility that the SARA sampling design might result in sparse
data for certain types of facilities. We checked the quality of the model fit to the data using the
model deviance and degrees of freedom (see method from SAS Proceedings Paper 247-26). Our
test of the null hypothesis that there was a better fitting model than our final model returned a
nonsignificant p-value, indicating that our final model was a good fit to the data. Finally, we
checked the deviance and Pearson residuals for our final model and performed sensitivity
analyses by removing the two observations with the greatest residuals, then assessed their impact
on parameter estimates. As there was little impact, these observations were added back to the
main analysis.

Results

Descriptive Data

The count of different EM-NCD present at each facility was highly skewed; scores clustered at 0,
the lowest possible score, with a long tail towards 10, the highest possible score (Fig 1). More
than a third of the facilities surveyed (37%) had no EM-NCD on site at all.

Figure 1. Distribution of EM-NCD counts in sampled facilities from the 2013 Uganda
SARA survey

Table 1 describes the ten EM-NCD by category, lowest level facility expected to stock[17], and
percentage of facilities stocking among the sample of facilities. No facility had all ten EM-NCD
in stock. Furthermore, availability varied considerably by medicine. The least available medicine
was the beclomethasone inhaler, which was only present at 3 of 196 (1.5%) facilities. The most
widely available medicine, amitriptyline, was present at 93 facilities (48%). Presence of a given
EM-NCD did not strongly correlate to the level facility at which it was expected. For example,
ACE inhibitors were expected only in referral hospitals but were present at 33 lower-level facilities (17%). Conversely, injectable insulin was expected at lower-level facilities but was only observed in 11% (22) of facilities.

Table 1. Essential medicines for treating non-communicable diseases (EM-NCD) included in the 2013 Uganda SARA survey

<table>
<thead>
<tr>
<th>Essential medicine</th>
<th>Disease Category</th>
<th>Lowest level facility expected</th>
<th>Facilities stocking</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nifedipine cap/tab</td>
<td>Cardiovascular</td>
<td>HC-III</td>
<td>64 (32.7%)</td>
</tr>
<tr>
<td>Enalapril cap/tab or alternative ACE inhibitor</td>
<td>Cardiovascular</td>
<td>Regional referral hospital</td>
<td>33 (16.8%)</td>
</tr>
<tr>
<td>Atenolol cap/tab</td>
<td>Cardiovascular</td>
<td>Hospital</td>
<td>40 (20.4%)</td>
</tr>
<tr>
<td>Metformin cap/tab</td>
<td>Diabetes</td>
<td>HC-IV</td>
<td>46 (23.5%)</td>
</tr>
<tr>
<td>Glibenclamide cap/tab</td>
<td>Diabetes</td>
<td>HC-IV</td>
<td>50 (25.5%)</td>
</tr>
<tr>
<td>Insulin regular</td>
<td>Diabetes</td>
<td>HC-IV</td>
<td>22 (11.2%)</td>
</tr>
<tr>
<td>Salbutamol inhaler</td>
<td>Asthma/Chronic Obstructive Lung Disease</td>
<td>HC-IV</td>
<td>39 (19.9%)</td>
</tr>
<tr>
<td>Beclomethasone inhaler</td>
<td>Asthma/Chronic Obstructive Lung Disease</td>
<td>HC-IV</td>
<td>3 (1.5%)</td>
</tr>
<tr>
<td>Amitriptyline cap/tab</td>
<td>Mental health/Depression</td>
<td>HC-III</td>
<td>93 (47.5%)</td>
</tr>
<tr>
<td>Simvastatin cap/tab</td>
<td>Cardiovascular</td>
<td>Excluded from Uganda EML</td>
<td>6 (3.1%)</td>
</tr>
</tbody>
</table>

Main Results
In bivariate analyses, region, facility type, managing authority, availability of HCT, and availability of HIV care were significantly associated with EM availability (Table 2). In the preferred multivariate model (Table 3), facilities under different types of managing authority perform significantly differently in terms of EM-NCD availability. The parameter estimate for PFP facilities compared to public facilities is 0.6837; in other words, PFP facilities have EM-NCD counts that are 98% higher on average—nearly double—those of public facilities (p<.001) even after adjusting for facility level. PNFP facilities also perform significantly better than public facilities in this model, but not nearly as well as the PFP facilities. Adjusting for the other variables, PNFP facilities have average EM-NCD counts that are 47% higher on average than public facilities (p<.014).
Table 2. Distribution of study variables and their association with availability of NCD medicines

<table>
<thead>
<tr>
<th>Variable</th>
<th>N (%)</th>
<th>None present</th>
<th>1 – 3 present</th>
<th>4 or more present</th>
<th>p*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Region</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>West</td>
<td>23 (11.7)</td>
<td>12 (52.2)</td>
<td>5 (21.7)</td>
<td>6 (26.1)</td>
<td>0.04</td>
</tr>
<tr>
<td>North</td>
<td>63 (32.1)</td>
<td>17 (27.0)</td>
<td>29 (46.0)</td>
<td>17 (27.0)</td>
<td></td>
</tr>
<tr>
<td>East</td>
<td>64 (32.7)</td>
<td>32 (50.0)</td>
<td>21 (32.8)</td>
<td>11 (17.2)</td>
<td></td>
</tr>
<tr>
<td>South</td>
<td>46 (23.5)</td>
<td>12 (26.1)</td>
<td>21 (45.7)</td>
<td>13 (28.3)</td>
<td></td>
</tr>
<tr>
<td><strong>Facility type</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&lt;.001</td>
</tr>
<tr>
<td>General hospital</td>
<td>17 (8.7)</td>
<td>0 (0)</td>
<td>1 (5.9)</td>
<td>16 (94.1)</td>
<td></td>
</tr>
<tr>
<td>HC-IV</td>
<td>17 (8.7)</td>
<td>2 (11.8)</td>
<td>4 (25.5)</td>
<td>11 (64.7)</td>
<td></td>
</tr>
<tr>
<td>HC-III</td>
<td>68 (34.7)</td>
<td>6 (8.8)</td>
<td>50 (73.5)</td>
<td>12 (17.7)</td>
<td></td>
</tr>
<tr>
<td>HC-II</td>
<td>94 (48.0)</td>
<td>65 (69.2)</td>
<td>21 (22.3)</td>
<td>8 (8.5)</td>
<td></td>
</tr>
<tr>
<td><strong>Managing authority</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Public</td>
<td>125 (63.8)</td>
<td>60 (48.0)</td>
<td>47 (37.6)</td>
<td>18 (14.4)</td>
<td></td>
</tr>
<tr>
<td>Private non-profit</td>
<td>43 (21.9)</td>
<td>6 (14.0)</td>
<td>16 (37.2)</td>
<td>21 (48.8)</td>
<td></td>
</tr>
<tr>
<td>Private for-profit</td>
<td>28 (14.3)</td>
<td>7 (25.0)</td>
<td>13 (46.4)</td>
<td>8 (28.6)</td>
<td></td>
</tr>
<tr>
<td>HCT^ available</td>
<td>152 (77.6)</td>
<td>41 (27.0)</td>
<td>65 (42.8)</td>
<td>46 (30.3)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>HIV care services available</td>
<td>113 (57.7)</td>
<td>30 (26.6)</td>
<td>48 (42.5)</td>
<td>35 (31.0)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Percentages may not sum to 100% due to rounding.

* P-value for χ² test. ^HCT=HIV Counseling and Testing
Table 3. Poisson regression model predicting greater availability of NCD essential medicines

<table>
<thead>
<tr>
<th>Variable</th>
<th>Adjusted β (SE)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Managing authority</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Public</td>
<td>Reference</td>
<td>---</td>
</tr>
<tr>
<td>Private non-profit</td>
<td>0.3882 (0.1573)</td>
<td>0.014</td>
</tr>
<tr>
<td>Private for-profit</td>
<td>0.6837 (0.1866)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Facility type</td>
<td></td>
<td></td>
</tr>
<tr>
<td>General hospital</td>
<td>0.6811 (0.2372)</td>
<td>0.004</td>
</tr>
<tr>
<td>HC-IV</td>
<td>0.7154 (0.2271)</td>
<td>0.002</td>
</tr>
<tr>
<td>HC-III</td>
<td>0.2165 (0.1498)</td>
<td>0.148</td>
</tr>
<tr>
<td>HC-II</td>
<td>Reference</td>
<td>---</td>
</tr>
<tr>
<td>Region</td>
<td></td>
<td></td>
</tr>
<tr>
<td>West</td>
<td>-0.0892 (0.2261)</td>
<td>0.693</td>
</tr>
<tr>
<td>North</td>
<td>-0.4217 (0.1727)</td>
<td>0.015</td>
</tr>
<tr>
<td>East</td>
<td>-0.4782 (0.1629)</td>
<td>0.003</td>
</tr>
<tr>
<td>South (Kampala)</td>
<td>Reference</td>
<td>---</td>
</tr>
<tr>
<td>Basic amenities score</td>
<td>1.0580 (0.3679)</td>
<td>0.004</td>
</tr>
<tr>
<td>Basic equipment score</td>
<td>1.3451 (0.4904)</td>
<td>0.006</td>
</tr>
<tr>
<td>NCD diagnostic capacity</td>
<td>0.2240 (0.0410)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>HIV counseling &amp; testing*</td>
<td>0.4530 (0.2295)</td>
<td>0.048</td>
</tr>
<tr>
<td>HIV services*</td>
<td>-0.4340 (0.1586)</td>
<td>0.006</td>
</tr>
</tbody>
</table>

*Dichotomous variable; reference category is 0.
The facility type parameter estimates indicate that general hospitals had EM-NCD availability scores nearly twice as high as the lowest level facilities (98% higher, p=.004). HC-IV facilities performed even better than general hospitals, with EM-NCD scores 105% higher than HC-II (p=.002). On average, HC-III do not have significantly greater EM-NCD availability than HC-II; these two facility types were the least likely to have any EM-NCD essential medicines on hand at all.

On average, and adjusting for the other predictors, facilities in the North and East have EM-NCD availability scores 34% lower (parameter estimate = -0.4217, p = 0.015) and 38% lower (parameter estimate = -0.4782, p = 0.003), respectively, than facilities in the Kampala region.

Finally, the two dichotomous variables indicating the availability of different types of HIV-related services indicate a complex set of interrelationships between HIV/AIDS services and the availability of essential medicines for NCDs. Offering HIV care and support services was associated with 35% lower average EM-NCD counts (parameter estimate = -0.4340, p = 0.006). However, offering HIV counseling and testing was associated with 57% higher EM-NCD counts (parameter estimate = 0.4530, p = 0.048).

**Other Analyses**

Due to concerns about sparse data, we considered and rejected a zero-inflated Poisson model. While a zero-inflated model was not appropriate in the case of Uganda, researchers interested in using SARA data to analyze health systems where some types of facilities are expected to never have any essential medicines available should consider these types of mixed models. Other
models that were considered and rejected, including a multilevel mixed model, are described in the Appendix.

Discussion

Our findings support previous work that demonstrates that Ugandan health facilities are poorly prepared to address the growing burden of NCD.\[12,16,18\] We extend this previous work by identifying and quantifying clear within-country disparities in preparedness. We found significant associations between EM-NCD availability and geographic region, managing authority, health facility type, and the range of HIV services. The availability of EM-NCD was substantially higher in PFP facilities than in public facilities and strikingly lower in the North and East regions. Availability of EM-NCD had a mixed relationship to availability of care and counseling for HIV. On the one hand, facilities that offer HIV care and support had lower average EM-NCD availability. However, facilities that offer HIV counseling and testing were associated with 57% higher EM-NCD availability counts.

Our model suggests that PFP health facilities are responding most quickly to the burgeoning need for EM-NCD. Adjusting for the other variables such as facility type and amenities, PFP facilities had EM-NCD counts nearly twice as high as public facilities. However, PFP facilities are often out of financial reach for most Ugandans. For example, a controller medicine for asthma, such as beclomethasone inhaler, costs approximately seven US dollars, the equivalent of three days wages, based on the per capita gross domestic product.
Facility type also had a sizable effect on EM-NCD availability in our model, though the facilities offering the most sophisticated services—general hospitals—do not necessarily have the greatest availability. Adjusting for region and other facility characteristics, the HC-IV facilities outperformed even general hospitals. Primary care HC-II and HC-III facilities, on the other hand, are likely to have few, if any, EM-NCD on hand. It may not be surprising that facility type has a significant effect on predicted EM-NCD count. However, consistent, long-term access to these medicines is critical for the effective and uninterrupted treatment of patients with chronic conditions. Individual countries adapt the WHO Essential Medicines List (EML) based on local disease prevalence, cost-effectiveness, and other national priorities. Countries also determine the lowest-level health facilities that are expected to stock each EM (see Table 1). Based on 2014 census data and hypertension prevalence data from the 2014 National Non-Communicable Disease Risk Factor Survey, an estimated 4.5 million Ugandan adults have hypertension.[19,20] Given the high prevalence, a reanalysis of these distribution guidelines would be prudent. Limiting the supply of anti-hypertensive medicines to higher level health facilities is incongruent with the provision of high quality, chronic care for persons with hypertension. Lower level health facilities, where the population is expected to receive primary health care, should be expected to stock EM for NCDs such as hypertension.

There is also evidence of clear regional disparities in EM-NCD. While the West region is not significantly different from the Kampala region, facilities in the North and the East have significantly lower counts of EM-NCD than those in Kampala, even controlling for other predictors of availability. On average and adjusting for the other predictors, facilities in the North have scores 34% lower and those in the East have scores 38% lower than facilities in the
Kampala region. One possible explanation is that the supply routes running East-West are of higher quality than those running North-South. However, in recent years, the Ugandan highway infrastructure has improved greatly and there are equally high quality highways spanning East-West as there are North-South. Certainly, further research is warranted towards understanding such in-country regional disparities.

Finally, the two HIV-related findings deserve special attention. We initially hypothesized that the availability of services for communicable diseases such as HIV/AIDS might be diverting resources and attention away from NCDs, resulting in lower average counts for facilities with HIV/AIDS services. However, the preferred model suggests a more complex set of interrelationships between HIV/AIDS services and the availability of EM-NCD. As hypothesized, offering HIV care and support services was associated with lower average NCD medicines counts. But offering HIV counseling and testing (HCT) was associated with higher counts of NCD essential medicines. It is plausible that facilities that are able to offer HCT have dispensary managers who are more attuned to the need to maintain chronic disease medicines. Or possibly these facilities have more sophisticated processes in place for monitoring and replenishing their medicine stock. Certainly, this is a result that we find compelling and in need of further study.

SARA data are collected using a complex, non-representative sampling strategy that must be corrected for using sample weights. In addition, SARA sample sizes are neither intentionally, nor necessarily, powered to provide significant estimates in regression models. This has been an impediment to wider use of these important data. Both the openly available country SARA
reports and all prior published research using SARA data have relied only on descriptive statistics, reporting simple unadjusted proportions rather than associations. We have shown that, despite these perceived barriers, researchers can use SARA data to develop regression models by applying straightforward corrections and diagnostic checks. By conducting the first Poisson analysis using SARA data, we have identified multiple disparities in availability of EM-NCD within Uganda.

Our approach had some limitations. First, like any cross-sectional design, ours is unable to infer causality. Longitudinal research is needed to better understand the sources of availability disparities like those we describe. Second, the SARA tool does not collect data on EM cost, thereby limiting its utility for directly addressing access, which is a function of both availability and cost. Further, like other EM availability surveys, SARA data reflect stock on the pharmacy shelf on a single day. This approach fails to account for variability in stock over time, which could be substantial and might particularly influence estimates of geographic disparity. Finally, though the public-facing data summary was available via the WHO[21], obtaining the raw dataset for analysis was challenging. These limitations point to the unmet need for technologies that provide real-time, hyper-local data to help spotlight and redress disparities in access faster -- and to map, measure and monitor disparities in access to care. Overlaying such insights with disease prevalence, population density, and health determinants such as traffic patterns and household income would further increase utility for decision-makers.

To deepen our understanding of variation in EM-NCD availability within LMIC, future research should aim to understand facility- and system-level barriers and facilitators to EM-NCD
availability. As more LMIC conduct SARA surveys, these datasets represent a largely untapped empirical resource for global health researchers and policymakers. We demonstrate that data generated by the SARA tool may be used to generate a robust, informative statistical model by applying well-recognized techniques to correct for some of the most common challenges inherent in these data. The results of such analyses can guide operational research and inform decision-making, investment, and priority-setting.
Supplemental Analyses

Given the complex sampling strategy and the possibility that health facilities in the same district may influence one another with regard to availability of EM-NCD, we also fit a multilevel mixed model to supplement our primary analysis. There was little evidence of need for a multilevel model and the parameter estimates of the multilevel mixed model were in general agreement with those of the easier-to-interpret Poisson model presented in the main analysis.

We also considered an alternative model including the presence of other essential medicines as a predictor, which was rejected because of evidence of serious multicollinearity.
REFERENCES


