Limitations to current methods to estimate cause of death: a validation study of a verbal autopsy model [version 3; peer review: 2 approved, 1 approved with reservations]

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Abstract
Background: Accurate information on causes of death (CoD) is essential to estimate burden of disease, track global progress, prioritize cost-effective interventions, and inform policies to reduce mortality. In low-income settings, where a significant proportion of deaths take place at home or in poorly-resourced peripheral health facilities, data on CoD often relies on verbal autopsies (VAs). Validations of VAs have been performed against clinical diagnosis, but never before against an acceptable gold standard: the complete diagnostic autopsy (CDA).

Methods: We have validated a computer-coded verbal autopsy method -the InterVA- using individual and population metrics to determine CoD against the CDA, in 316 deceased patients of different age groups who died in a tertiary-level hospital in Maputo, Mozambique between 2013 and 2015.

Results: We found a low agreement of the model across all age groups at the individual (kappa statistic ranging from -0.030 to 0.232, lowest in stillbirths and highest in adults) and population levels
(chance-corrected cause-specific mortality fraction accuracy ranging from -1.00 to 0.62, lowest in stillbirths, highest in children). The sensitivity in identifying infectious diseases was low (0% for tuberculosis, diarrhea, and disseminated infections, 32% for HIV-related infections, 33% for malaria and 36% for pneumonia). Of maternal deaths, 26 were assigned to eclampsia but only four patients actually died of eclampsia.

**Conclusions**: These findings do not lead to building confidence in current estimates of CoD. They also call to the need to implement autopsy methods where they may be feasible, and to improve the quality and performance of current VA techniques.

**Keywords**
Validation, verbal autopsy, cause of death, complete diagnostic autopsy, Mozambique

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

**Grant information**: This work was supported by the Bill & Melinda Gates Foundation [OPP1067522 and OPP1128001]. This work is also supported by the Spanish Instituto de Salud Carlos III [FIS, PI12/00757; CM, Acciones CIBER]. ISGlobal is included in the CERCA Programme /Generalitat de Catalunya.

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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**How to cite this article**: Menéndez C, Quintó L, Castillo P et al. Limitations to current methods to estimate cause of death: a validation study of a verbal autopsy model [version 3; peer review: 2 approved, 1 approved with reservations] Gates Open Research 2021, 4:55 https://doi.org/10.12688/gatesopenres.13132.3

**First published**: 28 May 2020, 4:55 https://doi.org/10.12688/gatesopenres.13132.1
Introduction

Global and disease-specific health statistics are regularly published and constitute an essential tool to define priorities and goals, identify inequalities, and track progress, including the achievement of global targets such as the health-related Sustainable Development Goals (SDGs). Insufficient confidence in the accuracy of estimates, particularly in those related to cause of death (CoD) has been indicated as a constraint to reduce mortality globally\(^1,2\). This lack of precise information on CoD in many low-income settings is largely explained by the significant number of deaths that occur either at home or at poorly resourced health facilities, with significant limitations of both complete diagnostic autopsies as well as accurate diagnostic methods; but also to the very limited number of diagnostic autopsies performed partly due to the massive shortfall of trained pathologists\(^3,4\). As such, CoD in low-income settings, continues to rely on estimates based on clinical records and verbal autopsies (VAs).

Clinical errors, which are common even in well-equipped hospitals, are more frequent in resource-restricted settings\(^3,5,6\). On the other hand, VAs remain the most practical and commonly used approach to estimate CoD at the population level in low-income settings\(^6\). A verbal autopsy consists of a structured interview to witnesses of the death subsequently interpreted and coded by physicians or using computerized methods. The method has shown to provide inconsistent results over time and place\(^7\). In addition, its diagnostic accuracy depends on the CoD, being high when the disease has a characteristic and well defined set of signs and symptoms, but much lower for conditions with unspecific symptoms, notably, malaria and acute respiratory infection in children, or meningitis in all age groups\(^8\). This results in frequent misclassifications of the CoD, which in turn leads to inaccurate cause-specific mortality rates\(^8\). Computerized methods of interpretation of the VA questionnaire have been developed to overcome some of the limitations of the VA technique. These methods are based either on algorithms derived from deaths with a medically confirmed CoD, or on probabilistic analyses\(^8\).

Computerized VA methods have been validated against physician-certified VA and clinical records\(^6,8\). However, neither computer-coded VA, nor physician-certified VA techniques have been validated against the complete diagnostic autopsy (CDA), the true gold standard for CoD determination. We present herein the results of a validation study of a commonly used computer-coded VA method, the InterVA (Interpreting Verbal Autopsy) model against the CDA in a series of deaths occurring in Maputo, Mozambique.

Methods

Study design and setting

The study included 316 CDA performed to patients who died between 2013 and 2015 at the Maputo Central Hospital, a 1500-bed institution that serves as the referral center for other hospitals in Mozambique. All the patients included in this analysis fulfilled the following criteria: (1) a CDA requested by the clinician as part of the medical evaluation of the patient and (2) informed consent to perform the autopsy given by the relatives. The following exclusion criterion was established: death of traumatic origin. In order to select only two cases per day from among the daily CDA requests received at the department of pathology (between 5 and 12 per day) without introducing selection biases, the two patients with death recorded before and closest to the time of 8:00 a.m. were included in the study. All maternal deaths that occurred in the study period were included.

From the 316 cases, 18 (6%) were stillbirths, 41 (13%) were neonates, 54 (17%) were children 1 month-15 years of age, 91 (29%) were maternal deaths and 112 (35%) were other adults. Written informed consent to perform the autopsy was obtained from the relatives of the deceased patients. In Maputo province malaria transmission is reported to be low (3%) and HIV prevalence is high (22%)\(^1,2,10\).

This study received approval by the Clinical Research Ethics Committee of the Hospital Clinic of Barcelona, Spain (File 2013/8677) and the National Bioethics Committee of Mozambique (Ref. 342/CNBS/13).

Determination of the cause of death by the complete diagnostic autopsy

The methodology for CoD determination by the CDA has been described in detail elsewhere\(^14-18\). Briefly, a panel of experts evaluated the CDA macroscopic, microscopic and microbiologic data, as well as the clinical information and assign the CoD. All morbid conditions directly leading to death, any underlying and any other significant conditions possibly contributing to death were codified according to the international classification of diseases, tenth revision (ICD-10, ICD-10 MM for maternal deaths)\(^19\). When more than one severe diagnosis was identified, the disease most likely causing the death was considered the final diagnosis\(^16-17\).

Cause of death assignment by the Verbal Autopsy model

We used the InterVA probabilistic model because it is one of the most commonly implemented VA tools\(^20\) and has shown a generally good level of agreement with the physician-coded verbal autopsy approach; it has also the advantage of being a completely reproducible method, reliable and standardized to interpretation\(^21,22\) reducing subjectivity. The InterVA method is based on the Bayes’ theorem and calculates the probability of a set of CoD given the presence of indicators reported in VA interviews\(^23,24\). We used version 4.04 of the model (InterVA-4) since the most recent version (InterVA-5) had not been released.
yet. In this analysis, the information feeding the model was extracted by the attending physician at the hospital from the clinical record of the deceased individual and from the obstetric record in perinatal deaths (Extended data\textsuperscript{2}; Clinical and epidemiological data collection questionnaire), unified into the WHO 2012 VA standard format\textsuperscript{2}, converted into the 245 input indicators of the VA model, and processed with malaria prevalence set to “low”, and HIV prevalence set to “high” using the InterVA4 package version 1.7.5 implemented in R version 3.5.0 software\textsuperscript{26}. Of the 245 input indicators of the model, 43 could not be extracted from the medical records; 24 (56%) of them were secondary questions, which are not pertinent if certain events did not occur.

Validation of the model
To validate the VA model across a variety of CoD distributions, 500 cause compositions based on uninformative Dirichlet sampling were generated for each study group\textsuperscript{27}. The performance of the model at the individual level was estimated comparing the CoD established by the CDA with the most probable CoD provided by the model. The Kappa statistic and the chance corrected concordance (CCC) were used as measures of the overall performance of the model (Extended data\textsuperscript{2}; Table S1 and Figure S1)\textsuperscript{27–29}.

At the population level, cause-specific mortality fractions (CSMFs) were calculated for each CoD and method within each study group. Since the model estimates up to three CoD with associated likelihoods for each cause, all identified CoD were considered as proportional to their partial likelihoods in the rate calculations for the model. In contrast, only one CoD was considered for the CDA and consequently, the associated likelihood was assumed to be 1. The CSMF accuracy (CSMFA) and the chance-corrected CSMFA (CCCSMFA) were calculated to compare the CSMFs determined by the InterVA model with those determined by the CDA (Extended data\textsuperscript{2}; Table S1 and Figure S1)\textsuperscript{27}. All analyses were done in Stata version 15 (Stata Corp., College Station, TX, USA) and R version 3.5.0 (R Core Team, 2017) statistical packages.

Results
The VA model assigned one CoD in 267 (84%) cases and two CoD in 33 (10%) cases. In 16 (5%) cases the model resulted in a non-conclusive diagnosis. The average likelihood of the model in estimating the first CoD was 90% (range 89% to 99%), and for the second CoD it was 38% (range 35% to 46%) (Extended data\textsuperscript{2}; Table S2). Three of the 316 cases (1%) had a non-conclusive diagnosis in the CDA.

Assignment of the CoD at the individual level compared with the CDA
In 168/316 cases (53%) the two methods agreed in the CoD. Most of the agreement was in the first CoD, while only in 8 cases the agreement was in the second CoD with a mean likelihood of 38% [95%CI: (33–43)] (Extended data\textsuperscript{2}; Table S3). In 148/316 cases (47%), there was no agreement in the CoD between the two methods.

Overall, the performance of the VA method in assigning a CoD to individual deaths was low (Table 1). In stillbirths, the sensitivity of the model in identifying infections, fetal growth restriction, and intrapartum and intrauterine hypoxia was 0%. In neonates, the sensitivity was 93% for infectious CoD, while it was 0% and 25% for preterm complications and congenital malformations, respectively. In children, the sensitivity of the model in identifying an infectious disease as CoD was 83%, while it was 0% for the congenital malformations, tumors and other diseases. The sensitivity of the model in identifying maternal mortality causes was low for all conditions except for eclampsia (75%) and obstetric hemorrhage (75%). In other adults, the sensitivity of the model was highest for infectious diseases (68%) and lowest for malignant neoplasms (19%).

| CoD | Sensitivity of VA model | Sensitivity of CDA |
|-----|-------------------------|-------------------|
| Infectious diseases | 93% | 0% |
| Fetal growth restriction | 83% | 0% |
| Congenital malformations | 0% | 0% |
| Maternal mortality | 0% | 0% |
| Neonatal deaths | 0% | 0% |
| Stillbirths | 0% | 0% |

Table 2 shows the measures of overall concordance between the two methods corrected for chance by study group for all CoDs. The CCC ranged between -0.093 and 0.246, and Kappa statistic ranged from -0.030 to 0.232 (lowest in stillbirths and highest in other adults). Figure 1 presents the alluvial diagrams showing the differences in the assignment of individual CoD established in the two methods by study group.

Cause of death assignment of the model at the individual level among patients dying of infectious diseases
Table 3 shows the performance of the VA model in assigning CoD among cases who died of an infectious disease according to the CDA in all study groups by infection category. The sensitivity of the model in identifying an infectious disease as CoD was low for all infectious categories, being 0% for tuberculosis, diarrhea, disseminated and other infections. Figure 2 shows the alluvial diagram of the comparison of CoD assigned by both methods.

| CoD | Sensitivity of VA model | Sensitivity of CDA |
|-----|-------------------------|-------------------|
| Infectious diseases | 93% | 0% |
| Fetal growth restriction | 83% | 0% |
| Congenital malformations | 0% | 0% |
| Maternal mortality | 0% | 0% |
| Neonatal deaths | 0% | 0% |
| Stillbirths | 0% | 0% |

Table 4 shows the CSMFs estimated by the VA model and the CDA aggregated into broad categories of CoD. In stillbirths, the most frequent CoD assigned by the model was congenital malformation (39%); however, no case of congenital malformation was identified by the CDA. In addition, fetal growth restriction (FGR) was the most frequent CoD in stillbirths determined by the CDA (39%), but only one case was estimated as such by the model. Infectious diseases were responsible for 22% of stillbirths by the CDA, but no stillbirth was assigned to infectious diseases by the VA model. According to the VA model, no deaths were assigned to preterm complications in neonates, while these represented 12% of the neonatal deaths by the CDA. Among children, malignant neoplasms accounted for 13% of the deaths in the CDA, but no case was assigned to this CoD in the model. The model identified eclampsia as the second most prevalent cause of maternal mortality while only in four (4%) cases eclampsia was the cause of maternal death by the CDA. Complications of abortion was diagnosed in nine (10%) cases, none of them being identified by the VA method.
Table 1. Performance of the InterVA (Interpreting Verbal Autopsy) model compared to the complete diagnostic autopsy at individual-level prediction by study group and cause of death.

| Cause of death (CDA)          | n | Classification of cases | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|------------------------------|---|-------------------------|-----------------|-----------------|---------|---------|
|                              |   | TP  | TN  | FP  | FN  |         |       |       |         |       |         |       |
| **Stillbirths (N=18)**       |   |     |     |     |     |         |       |       |         |       |         |       |
| Infections                   | 4 | 0   | 14  | 0   | 4   | 0       | 100   | N/A  | 78      |       |         |       |
| Fetal growth restriction     | 7 | 0   | 10  | 1   | 7   | 0       | 91    | 0    | 59      |       |         |       |
| Intrapartum hypoxia          | 3 | 0   | 15  | 0   | 3   | 0       | 100   | N/A  | 83      |       |         |       |
| Intraterine hypoxia          | 2 | 0   | 16  | 0   | 2   | 0       | 100   | N/A  | 89      |       |         |       |
| Congenital malformations     | 0 | 0   | 11  | 7   | 0   | N/A     | 61    | 0    | 100     |       |         |       |
| Non-conclusive               | 2 | 1   | 7   | 9   | 1   | 50      | 44    | 10   | 88      |       |         |       |
| **Neonates (N=41)**          |   |     |     |     |     |         |       |       |         |       |         |       |
| Infections                   | 27| 25  | 2   | 12  | 2   | 93      | 14    | 68   | 50      |       |         |       |
| Congenital malformations     | 4 | 1   | 37  | 0   | 3   | 25      | 100   | 100  | 92      |       |         |       |
| Preterm complications        | 5 | 0   | 36  | 0   | 5   | 0       | 100   | N/A  | 88      |       |         |       |
| Intrapartum complication     | 3 | 1   | 36  | 2   | 2   | 33      | 95    | 33   | 95      |       |         |       |
| Other diseases               | 2 | 0   | 39  | 0   | 2   | 0       | 100   | N/A  | 95      |       |         |       |
| Non-conclusive               | 0 | 0   | 41  | 0   | 0   | N/A     | 100   | N/A  | 100     |       |         |       |
| **Children (N=54)**          |   |     |     |     |     |         |       |       |         |       |         |       |
| Infections                   | 42| 35  | 3   | 9   | 7   | 83      | 25    | 80   | 30      |       |         |       |
| Congenital malformations     | 2 | 0   | 52  | 0   | 2   | 0       | 100   | N/A  | 96      |       |         |       |
| Malignant neoplasms          | 7 | 0   | 47  | 0   | 7   | 0       | 100   | N/A  | 87      |       |         |       |
| Other diseases               | 3 | 0   | 44  | 7   | 3   | 0       | 86    | 0    | 94      |       |         |       |
| Non-conclusive               | 0 | 0   | 51  | 3   | 0   | N/A     | 94    | 0    | 100     |       |         |       |
| **Maternal deaths (N=91)**   |   |     |     |     |     |         |       |       |         |       |         |       |
| Infections*                  | 39| 11  | 39  | 13  | 28  | 28      | 75    | 46   | 58      |       |         |       |
| Abortion                     | 9 | 0   | 82  | 0   | 9   | 0       | 100   | N/A  | 90      |       |         |       |
| Eclampsia                    | 4 | 3   | 64  | 23  | 1   | 75      | 74    | 12   | 98      |       |         |       |
| Obstetric hemorrhage         | 16| 12  | 59  | 16  | 4   | 75      | 79    | 43   | 94      |       |         |       |
| Other obstetric complications| 6 | 0   | 85  | 0   | 6   | 0       | 100   | N/A  | 93      |       |         |       |
| Non-obstetric diseases**     | 16| 4   | 67  | 8   | 12  | 25      | 89    | 33   | 85      |       |         |       |
| Non-conclusive               | 1 | 0   | 89  | 1   | 1   | 0       | 99    | 0    | 99      |       |         |       |
| **Other adults (N=112)**     |   |     |     |     |     |         |       |       |         |       |         |       |
| Infections                   | 80| 54  | 20  | 12  | 26  | 68      | 62    | 82   | 43      |       |         |       |
| Malignant neoplasms          | 16| 3   | 95  | 1   | 13  | 19      | 99    | 75   | 88      |       |         |       |
| Other diseases               | 16| 10  | 66  | 30  | 6   | 62      | 69    | 25   | 92      |       |         |       |
| Non-conclusive               | 0 | 0   | 110 | 2   | 0   | N/A     | 98    | 0    | 100     |       |         |       |

CDA: complete diagnostic autopsy; n: number of cases; TP: true positives; TN: true negatives; FP: false positives; FN: false negatives; PPV: positive predictive value; NPV: negative predictive value; N/A: not applicable.

* Includes all infections, both obstetric and non-obstetric.

** Non-obstetric diseases do not include infections.
Table 2. Measures of performance of the InterVA (Interpreting Verbal Autopsy) model compared to the complete diagnostic autopsy at individual-level prediction for all causes of death by study group.

| Study group   | CCC   | Kappa   |
|---------------|-------|---------|
| Stillbirths   | -0.093| -0.030  |
| Neonates     | 0.119 | 0.142   |
| Children     | 0.020 | 0.020   |
| Maternal deaths | 0.179 | 0.159   |
| Other adults  | 0.246 | 0.232   |

CCC: Chance-corrected concordance calculated from 500 Dirichlet draws.

Figure 1. Alluvial diagrams of the differences in assignment of individual causes of death established by the Complete Diagnostic Autopsy (CDA) and InterVA (Interpreting Verbal Autopsy) model by study group. The stacked blocks represent the causes of death (CoDs) determined by the CDA (left) and by the InterVA model (right), and their size as proportional to the cause-specific mortality fractions (CSMFs). The branches between blocks represent differences in the composition of the CoDs between the CDA and the InterVA model, being their thickness proportional to the number of cases contained in both blocks connected by the branch. Each CoD is represented by a different color, which is the same in both diagnostic methods. The color of the branches is determined by the cause of actual death (CDA). The concordant cases between the CDA and the InterVA model are represented by branches connected to blocks of the same color. In contrast, misclassified cases are shown as branches connected to blocks of different color.
The model was less accurate in stillbirths (CSMFA of 0.11) than in the other groups (CSMFA ranging from 0.59 to 0.77). When corrected by chance, the accuracy of the model compared to the CDA was not better than that expected by chance in stillbirths (negative CCCSMFA), close to chance in maternal deaths (close to zero CCCSMFA) and better than that expected by chance but far from perfection in the other groups (Table 4).

**Table 3. Performance of the InterVA (Interpreting Verbal Autopsy) model at individual-level prediction among all cases who died of infectious diseases according to the complete diagnostic autopsy.**

| Cause of death (CDA)                  | n   | Classification of cases | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|---------------------------------------|-----|-------------------------|-----------------|-----------------|---------|---------|
|                                       |     | TP|TN|FP|FN| |
| Disseminated infections               | 51  | 0 |139|2 |51|0 |99 |0 |73 |
| Pneumonia                             | 36  | 13|102|54|23|36 |65 |19 |82 |
| Meningitis                            | 15  | 3 |173|4 |12|20 |98 |43 |94 |
| Tuberculosis                          | 7   | 0 |183|2 |7 |0 |99 |0 |96 |
| Diarrhoea                             | 2   | 0 |186|4 |2 |0 |98 |0 |99 |
| HIV/AIDS related infections           | 57  | 18|120|15|39|32 |89 |55 |75 |
| Malaria                               | 6   | 2 |181|5 |4 |33 |97 |29 |98 |
| Other infections                      | 18  | 0 |171|3 |18|0 |98 |0 |90 |

**CDA:** complete diagnostic autopsy; **n:** number of cases; **TP:** true positives; **TN:** true negatives; **FP:** false positives; **FN:** false negatives; **PPV:** positive predictive value; **NPV:** negative predictive value; **N/A:** not applicable

**Disseminated infections:** bacterial sepsis of newborn (n=21), puerperal sepsis (n=6), streptococcal sepsis (n=5) and other sepsis (n=19)

**HIV/AIDS related infections:** candidiasis (n=1), congenital viral diseases (n=1), cryptococcosis (n=11), cytomegaloviral disease (n=7), herpes simplex infection (n=1), miliary tuberculosis (n=20), salmonella infections (n=1), pneumocystosis (n=5), pulmonary tuberculosis (n=2), toxoplasmosis (n=7) and tuberculous meningitis (n=1)

**Other infections:** acute pericarditis (n=2), pyelonephritis (n=2), congenital viral diseases (n=2), chorioamnionitis (n=2), GBS infection (n=2), tetanus (n=1), puerperal sepsis (n=3), rickets (n=3) and zygomycosis (n=1)

Discussion

It is recognized that accurate information on what is causing deaths is essential to reduce mortality. In this study, we have assessed for the first time to our knowledge, the validity of a commonly used VA method in establishing the CoD compared with the gold standard (the CDA) in different age groups of patients dying at a tertiary-level hospital in Maputo, Mozambique. The agreement of the VA model was overall poor across all age groups and conditions, both at the individual and at the population level.

The two reference standards that have been used for validating computer-coded VA, i.e. physician-coded VA methods and health facility medical records, cannot be considered true gold standards. The comparison between computer-coded and physician-coded VA methods lacks an external reference or gold standard comparator. On the other hand, although health facility-derived information is considered as an appropriate reference standard for VA validation, reports from both high and low-income countries indicate that this information frequently contains clinical errors. It seems quite evident that if the main source of input to the VA tool is inaccurate, the output of the VA will not be precise either. Furthermore, if clinical errors are frequent even in well-equipped hospitals, it is expected that their frequency would be higher in VA data.

In this study, the performance of the VA model was overall poor in identifying CoDs in stillbirths. These findings disagree with those of a report from Pakistan using clinical data as reference standard, indicating that a physician-coded VA tool was valid to ascertain causes of stillbirths, specially congenital malformations. In neonates, the sensitivity of the model in identifying preterm complications as a CoD, was also very low (0%), which may be relevant for pre-term birth prevention programs. In contrast, the performance of the model in identifying infectious diseases as a cause of neonatal death had a high sensitivity, suggesting that it may be an adequate method to identify neonatal sepsis at the community. Among children, the sensitivity of the model was only high in detecting infectious diseases as a CoD but it did not identify deaths due congenital malformations and malignant neoplasms. In maternal deaths, the sensitivity of the model was high in assigning eclampsia as a cause of maternal mortality; however, the probability that a maternal death identified as eclampsia by the model was actually eclampsia was quite low. Although there were 26 maternal deaths assigned to eclampsia as the most probable CoD according to the VA model, only four were actually due to eclampsia (most misdiagnosed cases...
Figure 2. Alluvial diagrams of the differences in the individual cause of death as established by the complete diagnostic autopsy (CDA) and the InterVA (Interpreting Verbal Autopsy) model among patients who died of infectious diseases. The stacked blocks represent the causes of death (CoDs) determined by the CDA (left) and by the InterVA model (right), and their size as proportional to the cause-specific mortality fractions (CSMFs). The branches between blocks represent differences in the composition of the CoDs between the CDA and the InterVA model, being their thickness proportional to the number of cases contained in both blocks connected by the branch. Each CoD is represented by a different color, which is the same in both diagnostic methods. The color of the branches is determined by the cause of actual death (CDA). The concordant cases between the CDA and the InterVA model are represented by branches connected to blocks of the same color. In contrast, misclassification cases are shown as branches connected to blocks of different color. **Disseminated infections**: bacterial sepsis of the newborn (n=21), puerperal sepsis (n=6), streptococcal sepsis (n=5) and other sepsis (n=19) **HIV/AIDS related infections**: candidiasis (n=1), congenital viral diseases (n=11), cryptocoecosis (n=11), cytomegaloviral disease (n=7), herpes simplex infection (n=1), miliary tuberculosis (n=20), salmonella infection (n=1), pneumocystosis (n=5), respiratory tuberculosis bacteriologically and histologically confirmed (n=2), toxoplasmosis (n=7) and tuberculous meningitis (n=1) **Other infections**: acute pericarditis (n=2), pyelonephritis (n=2), congenital viral diseases (n=2), chorioamnionitis (n=2), GBS infection (n=2), tetanus (n=1), peritonitis (n=3), rabies (n=3) and zygomycosis (n=1) **Non-infectious diseases (by the InterVA model)**: congenital malformations (n=1), intrapartum complication (n=2), eclampsia (n=12), obstetric haemorrhage (n=10), non-obstetric diseases (n=5) and other diseases (n=29).
Table 4. Cause-Specific Mortality Fractions and associated measures of validation of the InterVA (Interpreting Verbal Autopsy) model compared with the complete diagnostic autopsy (CDA) at the population level.

| Group        | Cause of death       | InterVA | CDA | CSMF Accuracy |
|--------------|----------------------|---------|-----|---------------|
|              |                      | n*   | CSMF (%) | n | CSMF (%) | Uncorrected | Chance Corrected |
|              |                      |       |         |   |         |             |                |
| Stillbirths  | Infections           | 0    | 0.0     | 4 | 22.2    | 0.11        | -1.00          |
|              | Fetal growth restriction | 1  | 5.0     | 7 | 38.9    |             |                |
|              | Intrapartum hypoxia  | 0    | 0.0     | 3 | 16.7    |             |                |
|              | Intrauterine hypoxia | 0    | 0.0     | 2 | 11.1    |             |                |
|              | Congenital malformations | 7  | 38.9    | 0 | 0.0     |             |                |
|              | Non conclusive       | 10   | 56.1    | 2 | 11.1    |             |                |
|              | Overall              | 18   | 100.0   | 18| 100.0   |             |                |
| Neonates     | Infections           | 37   | 91.0    | 27| 65.9    | 0.73        | 0.50           |
|              | Congenital malformations | 1  | 2.4     | 4 | 9.8     |             |                |
|              | Preterm complications | 0    | 0.0     | 5 | 12.2    |             |                |
|              | Intrapartum complications | 2  | 4.5     | 3 | 7.3     |             |                |
|              | Other diseases       | 0    | 0.0     | 2 | 4.9     |             |                |
|              | Non-conclusive       | 1    | 2.1     | 0 | 0.0     |             |                |
|              | Overall              | 41   | 100.0   | 41| 100.0   |             |                |
| Children     | Infections           | 40   | 73.9    | 42| 77.8    | 0.77        | 0.62           |
|              | Congenital malformations | 0  | 0.0     | 2 | 3.7     |             |                |
|              | Malignant neoplasms  | 0    | 0.0     | 7 | 13.0    |             |                |
|              | Other diseases       | 7    | 13.7    | 3 | 5.6     |             |                |
|              | Non-conclusive       | 7    | 12.5    | 0 | 0.0     |             |                |
|              | Overall              | 54   | 100.0   | 54| 100.0   |             |                |
| Maternal deaths | Infections*         | 22   | 24.1    | 39| 42.9    | 0.59        | -0.05          |
|              | Abortion             | 0    | 0.0     | 9 | 9.9     |             |                |
|              | Eclampsia            | 25   | 27.5    | 4 | 4.4     |             |                |
|              | Obstetric hemorrhage | 26   | 28.1    | 16| 17.6    |             |                |
|              | Other obstetric complications | 0 | 0.4    | 6 | 6.6     |             |                |
|              | Non-obstetric diseases** | 11 | 12.0    | 16| 17.6    |             |                |
|              | Non-conclusive       | 7    | 7.9     | 1 | 1.1     |             |                |
|              | Overall              | 91   | 100.0   | 91| 100.0   |             |                |
| Other adults | Infections           | 65   | 58.2    | 80| 71.4    | 0.76        | 0.49           |
|              | Malignant neoplasms  | 4    | 3.7     | 16| 14.3    |             |                |
|              | Other diseases       | 33   | 29.8    | 16| 14.3    |             |                |
|              | Non-conclusive       | 9    | 8.4     | 0 | 0.0     |             |                |
|              | Overall              | 112  | 100.0   | 112| 100.0   |             |                |

CDA: Complete diagnostic autopsy; n*: sum of cases estimated by InterVA model (in cause 1 or 2) weighted by their associated likelihood. The residual likelihoods count as non-conclusive case fractions; n: sum of cases established by the CDA; CSMF: cause-specific mortality fractions; CSMF Accuracy: measures the quality at the population level, quantifying how closely the estimated CSMF values approximate the truth; Uncorrected: median cause-specific mortality fractions accuracy across 500 Dirichlet draws. It ranges from zero to one; Chance Corrected: Median Cause-Specific Mortality Fractions Accuracy for random allocation across 500 Dirichlet draws. A score of zero indicates predictive accuracy equal to random allocation.

* Includes all infections, both obstetric and non-obstetric.

** Non-obstetric diseases do not include infections.
died of infectious diseases), suggesting a significant overestimation of eclampsia as a cause of maternal mortality by this method. This is in agreement with a previous report where a high frequency of false positive clinical diagnosis of eclampsia compared to the CDA was also found, being most of them deaths from infectious diseases\(^3\). These findings are of relevance to eclampsia prevention programs, which may fail in reducing maternal mortality due to misdiagnosis. In adults, the sensitivity of the VA model was higher for infectious diseases compared to other CoD, but low in identifying malignant neoplasms as cause of mortality. According to these results, the model would underestimate malignant neoplasms as CoD in adults, which may be important for prevention programs of this condition in high mortality settings.

The performance of the model in identifying the specific infection CoD among patients who died of an infectious disease was overall low. The sensitivity of the model in identifying tuberculosis as a CoD was very low, which may be of public health relevance in high burden countries. Regarding malaria infection, the VA model and the CDA only agreed in two cases, while in the other four cases established by the CDA, the model assigned three of them to a non-infectious disease CoD and one as non-conclusive (Figure 2). Lack of precision at the individual level in assigning malaria infection as a CoD may be important to target malaria control efforts in the community and increasing programme’s effectiveness.

The main use of the VA information is to determine cause-specific mortality and distribution of CoD at the population level\(^7\); for this reason we also estimated the CSMF accuracy between the two methods. Both methods differed in the distribution of the proportion of the deaths assigned to several disease categories. When corrected by chance, the accuracy of the model in predicting in the population the CoD was poor, especially in stillbirths and maternal deaths and imperfect in the other groups.

A possible limitation of our study that might have influenced the predictions of the model, is that the indicators used to estimate the CoD by the VA model were extracted from medical records, since VAs were not done, which relates to the absence in the clinical records of some indicators of the model (43 indicators, 18% of the total). Nevertheless, most of these indicators (n=24) were secondary questions related to the duration of the event and therefore, not pertinent if the event did not occur. On the other hand, the most likely explanation for the lack of registration in the clinical record of the other 19 indicators (8%) is that they were not identified. The fact that the study is based in a large hospital might be seen as a limitation to extrapolate findings to deaths occurring in rural health-facilities or at home, since cause-composition of deaths in the community may be different to that of those occurring in a hospital. However, it is important to remember that this is a validation study and therefore, the objective was not that the deaths included were representative of those occurring in the community, but rather that the comparator of the VA was as true gold standard as possible. Thus, we needed a set of deaths, whose causes were established by the CDA, and therefore they had to occur in a hospital setting with autopsy facilities. On the other hand, to avoid that the cause-composition of deaths in that particular hospital and/or time-period affected the accuracy of the estimates of the VA, we created multiple test datasets with widely varying cause-compositions as it has been suggested\(^8\).

As explained in the methods section, we used InterVA version 4 because version 5 was not available at the time of this analysis. Even if the estimated CoD might differ between the two InterVA versions, a change in the group of CoD would not be expected. Otherwise it would mean that the two versions provide different results, requiring a revision of all published information using the previous InterVA model.

The post-2015 Development Agenda expects that high burden countries should have reliable information on number and CoD to reduce their main health problems\(^9\). However, this goal cannot continue to rely on imprecise measurement tools. The main shortcoming to achieve the SDGs is the imprecision of the currently used methods to establish CoD. These findings highlight the need of improving the quality and performance of current VA techniques by developing more precise tools for CoD ascertainment.

In conclusion, the “data revolution” of the post-2015 Development Agenda expects that high burden countries should have reliable information on number and causes of death in order to reduce main health problems through evidence-based decision-making, and target and monitor health programs\(^9\). However, this goal cannot continue to rely on imprecise measurement tools. The main shortcoming to achieve the SDGs, is not the scarce availability of physicians to carry out death certificates or VA codification, nor the solution is the available automated methods created to overcome some of the physician-coded VA limitations, but rather the imprecision of these methods to reliably establish causes of death. The findings of this study should serve to highlight the need to implement autopsy methods where they may be feasible, but even more importantly to improve the quality and performance of current VA techniques and to develop more precise CoD ascertainment tools.

Consent
Written informed consent for publication of the patients’ details and/or their images was obtained from the parents/guardian/relative of the patient.

Data availability
Underlying data
Study data cannot be shared in a public domain due to their sensitive nature and, being such as small sample, especially for some age-specific causes of death, it would be relatively easy to identify study individuals even if anonymized. However, deidentified data will be made available from the corresponding author on reasonable request. Requesters will be required to sign a letter of agreement detailing the mechanisms by which the data will be kept secure and access restricted to their study team.
The agreements will also state that the recipient will not share the data with anyone outside of their research team.

Extended data

Open Science Framework: Limitations to current methods to estimate cause of death: a validation study of a verbal autopsy model. https://doi.org/10.17605/OSF.IO/UMJV2

This project contains the following extended data:

- Clinical_questionnaire.pdf (Clinical and epidemiological data collection questionnaire)
- VA_validation_extended_data.pdf (PDF containing supplementary figures and tables)

Figure S1. Outline of statistical methods

Table S1. Description of metrics

Table S2. Study group and number of causes of death and their associated likelihoods as established by the InterVA method

Table S3. Number of cases and mean likelihood agreement between the InterVA’s predicted cause of death and that established by the CDAbY study group

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 thank the families of the deceased patients included in this study. The authors are grateful to all the members of the Department of Pathology of the Maputo Central Hospital, whose work and enthusiasm made this study possible, and also to the staff of the Centre de Investigação em Saúde de Maniça for their invaluable support. We would also like to thank Mrs Ariadna Sanz for her help in the preparation of the manuscript. ISGlobal is a member of the CERCA Programme, Generalitat de Cataluna.

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The manuscript is scientifically sound, well written and contributes to body of literature on the validity of VA in assigning CoDs. There are just few suggestions that I think will assist with better organization of the manuscript. There are a few areas where there are grammatical errors that need to be addressed.

There is inconsistence in the way InterVA model is being referred to in the manuscript as it is sometimes called the VA model and in other places model. I would recommend that the authors refer to the model by its name and not as 'VA model' as there are a number of other computer coded models for VA.

The authors indicate the number of CAD that were completed in the methods section and that number could go into the results section. I recommend the same for the paragraph describing the population of decedents, which can also go into results section.

The authors could start off the results section with a paragraph describing the population of decedents according to age groups. It would also be good to include a paragraph or two just describing the top CoDs assigned by the two methods for each study group before comparing them.

The heading of the second paragraph in the results could be changed to “Agreement/comparison of COD at individual level between the two methods” as its describing agreement and not just assignment.

The authors are not providing source data because of issues of confidentiality.

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?
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?
No

Are the conclusions drawn adequately supported by the results?
Yes

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 21 October 2020
https://doi.org/10.21956/gatesopenres.14319.r29737

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Clara Calvert
Department of Population Health, London School of Hygiene & Tropical Medicine, London, UK

General assessment:
Thank you for the opportunity to review this very clearly written article. As the first study to compare cause of death assignment from Verbal Autopsy (VA) based on automated methods to cause of death assignment based on complete diagnostic autopsy, this is a very useful contribution to the field. Overall, the authors make a coherent argument, though I have a few suggestions on how certain aspects could be clarified or improved.

Introduction:
Generally, this is very clearly written and provides a clear rationale for this research. I have just minor comments, as follows:

1. The final sentence of the first paragraph is slightly confusing, implying that certain conditions need to be tracked using clinical records and VAs, where I think it is all conditions in certain settings that require use of clinical records and/or VAs. I would suggest amending the final sentence of first paragraph to state that cause of death (CoD) distributions in many low-income settings rely on estimates based on clinical records.
2. You note that clinical errors mean that VA remains the most practical approach for estimating CoD at the beginning of the second paragraph; however, I think one of the main reasons for using VA is to capture deaths that occur in the community.

3. The definition of a verbal autopsy in this paper specifies that the interview is interpreted by a physician, but I would argue that this is one method to interpret the data from the VA and the other is automated methods. It would be good to clarify this.

4. When describing the diagnostic accuracy of VA interpreted using physician review in the second paragraph, you provide a reference to the study by Quigley et al.; this study looks only at children so it would be good to provide further reference to the study which showed poor diagnostic accuracy for meningitis in all age groups.

Methods:

1. It would be helpful to provide an indication of which deaths are likely to have a complete diagnostic autopsy (CDA) requested and possibly a rough idea of how many deaths receive a CDA request. My suspicion – which may be completely unfounded – is that these deaths may represent particularly tricky deaths to assign cause of death given that a CDA was requested by the clinician. This would have important implications on the interpretation of the results.

2. You write that InterVA-4 has the advantage of being “standardized to interpretation” – can you clarify what this means?

3. Could you add a citation for the R package for InterVA-4 in the methods?

4. You mention that the data were extracted from the clinical record, and it would be helpful to have some more clarification on this tool – initially I had assumed that this was a medical record in line with what was routinely used in the hospital; however, this looks much more like a questionnaire that was designed specifically as part of the project – it would be useful to know if this was the case and, if so, who completed this questionnaire.

5. In the validation of the model, you write that you generated 500 cause compositions across “each study group” – which study groups are you referring to? Do you mean by age and separating out maternal deaths? It would be good to include a statement that clarifies that “All analyses were undertaken by the following groups: stillbirth, neonates, children, pregnant and postpartum women and other adults.”

6. More generally, I felt that the “validation of the model” was lacking in detail. It would be helpful to make the rationale for generating 500 cause compositions clear. You also present sensitivity, specificity, PPV and NPV in the results but do not mention the calculation of these in the methods; at the very least you should mention that you calculate these and use the CDA as the gold standard.

Results:

1. As a very minor point, it would be better to use the term InterVA-4 throughout the paper, rather than referring to it as the VA model.
2. Change reference to “tumours” in text to “malignant neoplasms” to match terminology used in Table 1.

3. The tables and figures are very clear; I particularly like the alluvial diagrams which are a very nice way to present these findings. Note that there is a small typo in the title of Figure one: change “misclassification” to “misclassified”.

Discussion:
A possible additional point to make is that one of the key drivers of the outputs from the InterVA-4 algorithm is the Symptom-Cause Information (SCI) which was initially derived from a group of physicians; information that is generated from studies like this has the potential to inform future iterations of the SCI to improve the performance of the automated methods (and newer versions of the algorithms are increasing opportunities for studies to specify their own SCI).

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?
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?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: maternal health; verbal autopsy; HIV

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.
This article is very interesting as a scientific work that explains the validation of the verbal autopsy model. The research problems have been described quite clearly and the research methods were presented in detail and were accurate. The results are presented clearly and perfectly with proper and in-depth discussion using sufficient references. The conclusion drawn was adequately supported by the results and discussion. But, perhaps there should be provided recommendations for researchers and practitioners regarding the use of verbal autopsy data in determining the cause of death.

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?
I cannot comment. A qualified statistician is required.

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

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Public health, health demography, epidemiology, research methodology, adolescent reproductive health, maternal and child health.

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.
Llorenç Quintó, Barcelona Institute for Global Health (ISGlobal), Hospital Clinic of Barcelona, Universitat de Barcelona, Barcelona, Spain

We really appreciate the comments on this paper made by Prof Byass, whose opinions we always considered very highly.

In fact, since the beginning of this analysis (more than 2 years ago), we invited him to revise its findings and even participate as an author. Since time has passed he may have forgotten this but we will be happy to resend him these emails to refresh his memory. I think this answers his question number 4.

Version 4 of InterVA may be considered outdated by version 5, and future studies should use this latest version. However, this does not mean that all the results and decisions made using version 4 are invalid or incorrect. If that would be the case, should all manuscripts published, protocols, and health policies done based on InterVA4 findings be reconsidered? Indeed, this would be very complex to handle and disturbing. Furthermore, as discussed in the manuscript, assuming that InterVA4 and InterVA5 results do not completely match, it would be very concerning if they involved changes in the broad groups of causes of death used in the analysis. Finally, we want to clarify that the statistical analysis of this study was carried out at late 2017-early 2018. Therefore, it is evident that (1) version 5 of InterVA did not yet exist then and (2) we were able to use the WHO international standard Verbal Autopsy tool, WHO-2016, to format the data appropriately.

Regarding the comment on the grouping of the causes of death, the preliminary analysis was done according to the standard categories of cause of death as suggested by Prof Byass and the results were very similar to those presented in the manuscript. However, to facilitate interpretation of results we decided to regroup them. On the other hand, despite that we calculated concordance statistics corrected by chance, we wanted to avoid discordant results being misinterpreted as a consequence of the high number of categories.

Clara Menéndez
Llorenç Quintó

Competing Interests: No competing interests were disclosed.

Reader Comment 05 Jun 2020

Peter Byass, InterVA, Sweden

Although this unreviewed pre-print is potentially interesting, there are some aspects that make its findings questionable.

1. This pre-print was published on 28 May 2020, including the statement "We used version 4.04 of the model (InterVA-4) since the most recent version (InterVA-5) had not been released yet." In fact, InterVA-5.0 was released in April 2019, and a paper describing the update in detail was published in BMC Medicine in May 2019 (https://bmcmedicine.biomedcentral.com/articles/10.1186/s12916-019-
1333-6), but is not cited in this pre-print. A further minor technical update, InterVA-5.1, was issued in April 2020. All of this is documented on an open-source basis at www.interva.net and the supporting Github repository https://github.com/peterbyass/InterVA-5

2. Since the analysed deaths and autopsies took place in 2013-2015, it was clearly not possible for the current WHO international standard Verbal Autopsy tool, WHO-2016, to have been used, but it seems that even the WHO-2012 VA standard was not used directly for the interviews, which is a further potential limitation. The WHO-2016 international standard constituted a major revision over the WHO-2012 VA standard, as detailed in PLOS Medicine in January 2018 (https://journals.plos.org/plosmedicine/article?id=10.1371/journal.pmed.1002486). These are major methodological limitations, which would not have precluded the application of InterVA-5, but should have been carefully discussed in this pre-print.

3. Both the WHO-2012 and WHO-2016 verbal autopsy standards define a set of ICD-10 compliant cause-of-death categories, which are completely implemented in all the InterVA tools. These international standards for VA cause categories were not used at all in this pre-print. It would have been more informative to have presented the InterVA output according to these standard cause categories, and to have used the WHO VA standard documentation to have re-classified the autopsy results on the basis of the detailed ICD-10 that were presumably generated by the pathologists, as the basis of comparison.

4. The InterVA group is always willing to engage with users of its open-source public-domain tools for interpreting verbal autopsy, and it is unfortunate that we were not approached during the analysis of this potentially interesting, but potentially flawed, study.

**Competing Interests:** All InterVA tools are provided as public goods, on an entirely non-profit basis.