1	Title:
2	Subnational burden estimation in Tuberculosis: generation and application of a new tool in Indonesia
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35 **Abstract** Setting In many high tuberculosis (TB) burden countries, there is substantial geographical heterogeneity 36 in TB burden. In addition, decisions on TB funding and policy are highly decentralised. Subnational 37 38 estimates of burden however are usually unavailable for planning and target-setting. 39 40 Objective and Design We developed SUBsET to distribute national TB incidence through a weighted score using selected variables, and applied for the 514 districts in Indonesia, which have substantial 41 42 policy and budgetary autonomy in TB. Estimated incidence was compared to reported facility and 43 domicile-based notifications to estimate the case detection rate (CDR). Local stakeholders led model 44 development and dissemination. 45 46 **Results** The final SUBsET model included district population size, level of urbanisation, socio-economic 47 indicators (living floor space and high school completion), HIV prevalence and air pollution. We estimated district-level TB incidence between 201 and 2,485/100,000/year. The facility-based CDR 48 49 varied between 0 and 190% with high variation between neighbouring districts, e.g. suggesting strong cross-district health utilisation, which was confirmed by domicile-based CDR estimation. SUBsET results 50 51 informed district-level TB action plans across Indonesia. 52 Conclusion Applying SUBsET to estimate the subnational burden can be important for high-burden 53 countries and inform TB policy-setting at the relevant, decentralised administrative level. 54 55

INTRODUCTION

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57 TB remains the leading cause of death from a single infectious agent and funding to fight the disease 58 remains limited. The burden of TB is widely assumed to be heterogeneously distributed within 59 60 countries,² and policy decision-making, including setting TB care and prevention planning and budgeting, 61 often takes place at the subnational level. To inform decision making at this level, and tailoring of TB 62 care and prevention efforts to local epidemiology, subnational estimates of TB burden are key. 63 64 While many high TB burden countries have conducted national TB prevalence surveys to obtain a better estimate of their TB burden, these surveys do not provide estimates on relevant subnational 65 administrative levels. Various studies have reported subnational estimation of disease burden, 3-10 though 66 few in TB, which often used complex methods that cannot be easily understood by local policy makers. 11-67 ¹⁶ As such, subnational policy makers are usually left without estimates to inform planning. Data on TB 68 69 notifications is usually available at subnational level, but provide a poor reflection of disease burden.² 70 Indonesia, with a total population of around 260 million people in 2017, consists of 34 provinces and 514 71 districts. 17,18 Since 1999, local (i.e. Provincial and District) governments have full autonomy to manage 72 health, financing, planning, and budgeting. 19 Health care is provided by the public and a large private 73 sector.²⁰ Although TB notification is mandatory, only 53% of all estimated incident cases were notified to 74 the National TB Program (NTP).¹ 75 76 77 Following a recent inventory study, Indonesia is estimated to have approximately 842,000 incident TB cases a year in 2017. To achieve ambitious targets for ending the TB epidemic by 2030, the Indonesia 78 NTP has encouraged local governments to develop a district action plan, ²¹ that is linked to the National 79 80 Strategic Plan but tailored to the local challenges, including estimated local burden and health system 81 utilisation. 82 83 Our aim was to develop a tool to estimate district-level incidence and health system utilization, balancing 84 detail and granularity with simplicity, so both method and result could be effectively disseminated to local government, and adapted for other high burden countries. We describe the development, findings and 85

dissemination of the SUBsET (SUBnational Burden Estimation for Tb) tool.

METHODS 88 89 Principle of method To promote acceptability and application of the results by policy makers, we worked from the principle 90 91 that the model should be as simple as possible, use widely available software, and involve a limited 92 number of calculation steps while still utilising available data in an efficient way. Data to inform the 93 model was required to be available in 95% of districts and have an association with TB burden. 94 95 No separate ethics approval was obtained as all data were publicly available or anonymised at time of 96 analysis. Model development, including the selection of variables, was inclusive, with direct input from 97 the NTP, relevant partners and representatives from local academia. Taking into account that program 98 indicators and milestones for the End TB strategy were set on incidence rather than prevalence, we chose TB incidence as our outcome.²² 99 100 101 Data 102 Burden estimates The national level incidence estimate from WHO Global TB report was used as the starting point. In 103 104 2014, the prevalence survey found substantial differences in burden between 3 regions (Sumatera, Java-Bali, and Others, i.e. regions other than Sumatera and Java-Bali). 23 We applied the same distribution to 105 106 the national incidence estimate. 107 108 Variables for model 109 Population size for each district was based on estimates from the Central Statistics Agency (BPS) that released a 2010-2020 district population projection for each province based on 2010 National Population 110 survey.²⁴ 111 112 Additional variables were extracted from the National Socio-Economic Survey, an annual socio-113 demographic survey which covers the whole nation and is powered for district-level estimates.²⁵ We 114 identified urbanisation, floor space, and education level (see table 1 for definitions), which were also 115 measured in the prevalence survey. We also included HIV burden, ^{26,27} and air pollution levels, ^{28–30}. 116 117 118 To inform current health system performance or utilisation, and to check estimated values of burden, the NTP provided both domicile-based (according to patient's address) and health-facility-based (according 119 to facility address) notification data for each district.³¹ 120 121

122 Model

- The SUBsET tool combined all available data to distribute National TB burden through a weighted score
- for each of the 514 districts, through the steps outlined below.

125

- 126 Step 1: Regional incidence
- 127 Incidence estimate of those three regions was calculated by applying the distribution of absolute TB
- prevalence across the respective regions among 2017 Indonesia population in the respective regions:

$$I_r^{(case)} = \frac{P_r^{(case)}}{P^{(case)}} \times I^{(case)}$$

- 130 where:
- 131 $I_r^{(case)}$ = Estimated TB incident cases in region r
- 132 $Pr^{(case)}$ = TB prevalent cases (absolute value) in region r
- 133 $I^{(case)}$ = National TB incident cases (absolute value)
- 134 $P^{(case)}$ = National TB prevalent cases (absolute value)

135

- 136 Step 2: Variable weight
- For the socioeconomic variables, through conducting multivariable logistic regression we were able to
- estimate the relative risk directly, by region, from the 2014 prevalence survey.²³ For HIV prevalence and
- air pollution, values from the literature were used. 27–30

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- We then calculated a weight for each variable by multiplying the regional relative risk with the proportion
- in that district (e.g. proportion living in an urban area):

$$S_d^{(v)} = \left(Pr\left(v_d\right) \times RR_r^{(v)}\right) + \left(1 - Pr\left(v_d\right)\right)$$

- where,
- 145 $S_d^{(v)}$ = weight for variable (v) in district d
- 146 $Pr(v_d)$ = proportion variable (v) among population in district d
- 147 $RR_r^{(v)}$ = TB relative risk ratio for variable (v) in region r
- 148 $1 Pr(v_d) = 1 \text{proportion of variable } (v) \text{ in district } d$

- 150 Step 3: Calculation of total weight score per district
- 151 A total score for each district was calculated by multiplying all variable weights with the population size:

$$S_d = N_d \times s_d^{(floor/kapita < 8m^2)} \times s_d^{(urban)} \times s_d^{(low\ education)} \times s_d^{(HIV)} \times s_d^{(air\ pollution)}$$

where:

154 S_d = total score for district d

155 N_d = number of population in district d

156 $s_d^{(floor/kapita < 8m^2)}$ = weight score for variable living floor space in district d

157 $s_d^{(urban)}$ = weight score for variable level of urbanisation in district d

158 $s_d^{(low\ education)}$ = weight score for variable junior high school completion in district d

159 $s_d^{(HIV)}$ = weight score for variable HIV prevalence in district d

160 $s_d^{(air\ pollution)}$ = weight score for variable air pollution prevalence in district d

161

Step 4: Distribution of burden

163 Total weight score per region was calculated by adding up the total weight score per district by respective

region, and then distributing the estimated burden across districts-based total on district score from step 3:

$$I_d^{(case)} = \frac{S_d}{S_r} \times I_r^{(case)}$$

where:

167 $I_d^{(case)}$ = Estimated TB incident cases in district d

168 S_d = Total weight score in district d

169 $I_r^{(case)}$ = Estimate TB incident cases in region r

170 S_r = Total weight score all districts in region r

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Calculation of district-level Case Detection Rate (CDR)

173 To estimate the district-level CDR, the estimated burden in each district was compared to both domicile-

and health-facility-based reported notifications. Comparing both domicile- and health-facility-based

notifications within and between surrounding districts allowed assessment of district health system

performance and cross-district health utilisation.

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Validation of SUBsET results

While model validation with data is desirable,³² neither the prevalence survey or inventory study enabled

a district-level comparison. The prevalence survey did not cover complete districts, and the inventory

study was powered to provide a national, not district-level estimates. An attempt to use inventory study

data at the district level would lead to extremely wide uncertainty intervals around the therefore non-

informative point estimates.

184	
185	Dissemination and adoption of model
186	The model was disseminated and discussed at provincial and district levels, followed by a round of
187	revisions during a national-level stakeholder meeting. The final development step resulted in the addition
188	of two variables to capture strong heterogeneity in HIV prevalence, and measured air pollution between
189	districts.
190	
191	Uncertainty intervals
192	Uncertainty intervals were calculated by generating 10,000 random draws from the distribution for both
193	the regional incidence estimate as well as relative risks for included variables. ^{23,27-30}
194	
195	Sensitivity analysis
196	To understand the heterogeneity captured by our model, we compared the results of our calculation with
197	an estimate based on regional incidence and population size alone. We also performed a calculation where
198	we removed each individual variable and compared the results with the full model.
199	
200	The model was set up in Microsoft Excel, multivariate analyses for region specific TB relative risks were
201	conducted in STATA version 14. We used spmap ado file in STATA version 14 to create the maps which
202	visualise the distribution of the district TB burden estimates and CDR throughout Indonesia, particularly
203	within provinces, thus allowing us to better understand the connection or relationship between one area to
204	another.
205	
206	RESULTS
207	Model
208	Relative risks for the model variables used in step 2 are shown in Table 2.
209	
210	The range of values across districts for each risk factor was wide (see Table 2, column 4 and 5). When the
211	relative risks were combined with the data for each risk factor, differences in population weight for
212	districts were found in each region i.e. median (range) relative weights Sumatera 2.52 (2.29-2.75), Java-
213	Bali 1.50 (1.37-1.64), and Others 2.10 (1.91-2.29).
214	
215	District-level TB Incidence
216	Fig 1 shows the distribution of the SUBsET estimated TB incidence across the 514 districts in Indonesia.
217	The estimated point values for TB incidence ranged between 201 and 2,485/100,000/year. The estimated

218 TB incidence rates was lowest at Java-Bali region (average median 242/100,000, range 201-787) 219 compared to Sumatera (373/100,000, 295-918) and Others (350/100,000, 280-2485). However, considering that 58% of the total population of Indonesia resides in Java-Bali, ²⁰ this region has the highest 220 absolute number of TB cases.²³ 221 222 223 **District-level CDR** 224 Fig 2 shows the distribution of the estimated facility-based CDR throughout all districts. While some 225 districts have very low CDR (0-20%, dark red colour) some others have very high CDR (>100%, green 226 colour) with a range of 0 to 190%. Among 24 (5%) districts with an estimated facility-based CDR of 227 228 more than 100%, 15 were urban and suburban districts, surrounded by rural districts, which usually have fewer or lower quality TB services (Fig 2, pull outs). Twenty-one districts (4%) had an estimated facility-229 230 based CDR between 80 and 100%. 231 232 For domicile-based CDR, 9 (2%) districts had an estimated CDR of more than 100%. A further 24 (5%) districts had a domicile-based CDR between 80% and 100% and 51 (10%) had a domicile-based CDR 233 234 below 20%. At the district level, there was considerable contrast between facility and domicile-based 235 CDR. As an example, for the year 2017, Salatiga city, Surakarta and Magelang city had 121%, 129% and 236 170% facility-based CDR while the domicile-based CDR were only 32%, 39% and 33% respectively (Fig. 237 2, pulls out). 238 239 **Uncertainty analyses** Uncertainty analyses provided ranges for incidence rate per 100,000/year population at district level as 240 241 well as at regional level. For Sumatera Region, this resulted in value (95% uncertainty interval) of 413.4 (305.3-530.8), for Java-Bali 268.0 (212.3-321.0), and for Others 380.1 (277.8-495.9). District-level 242 243 uncertainty intervals are shown in figure 3. 244 245 Sensitivity analyses Figure 3 shows the additional variation in estimated incidence introduced by the variables in our model, 246 by comparing with a model including population size and regional differences in prevalence. We found 247 that 30% of the districts had a higher and 70% had a lower point estimate for TB incidence rates 248 249 compared to previous estimates. The newly estimated TB incidence rates were more than 10% different 250 (higher or lower) from the previous TB incidence estimate for 73% of the districts.

252 Removing a single variable had no relevant impact on the distribution of the estimated burden in the 253 model which shows that there is no single model variable that dominates the differentiation between 254 districts. Considering the dominant influence of population size in the burden distribution across districts, 255 a lower or higher value of a relative risk in a single variable would lead to a lower or higher value of the 256 uncertainty interval. 257 258 **Model dissemination** The district- and provincial-level TB burden estimates were fed into the development of District and 259 Provincial Action Plans, particularly to inform policy decisions on budget, resource allocation, and 260 intervention planning. Estimates were also incorporated in the 2016-2020 TB National Strategic Plan, and 261 have fed into joint AIDS, TB and Malaria policy meetings at the national level.³³ 262 263 264 **DISCUSSION** The SUBsET tool approach was found to provide an accessible and intuitive model for subnational 265 266 burden estimation. Our final model used five variables to distribute TB incidence from three regional estimates across 514 districts in Indonesia. The model provided substantial differentiation, estimating an 267 incidence ranging between 201 and 2,485/100,000/year. The facility-based CDR varied between 0 and 268 269 190%, highlighting low-performing districts, and cross-district health utilisation. Dissemination of the 270 SUBsET tool showed rapid uptake and acceptance of results. 271 On district-level, the SUBsET facilitated the comparison of facility-based and domicile-based-CDR 272 273 which highlighted previously unrecognised cross-district health system utilization. These insights encouraged such districts to improve their own health care system and case detection, as well as improve 274 275 collaboration with neighbouring districts. 276 277 Limitations 278 Our work has several limitations. Both the regional distribution of incidence and associations between TB 279 burden and socioeconomic variables are based on the 2014 national TB prevalence survey, not on directly measured incidence. While those associations may be slightly different if directly calculated for 280 incidence, we feel they are a reasonable approximation and the limited bias is outweighed by the ability to 281 282 calculate the relative risks directly for the population and time period. For HIV, the association matches the range of the relative risk of developing TB in HIV-positive infected persons in concentrated and low-283 284 level HIV prevalence area; likewise, the association between air pollution and risk of developing TB

corresponds with results found in various studies from low to middle income settings. 27-30

286 287 Second, we acknowledge the likelihood that there may be a residual or uncaptured variation of TB 288 incidence beyond that captured by the model, e.g. due to differential levels of malnutrition, or in 289 additional sub-categories within the variables included, but data was not available to include in the model. 290 291 Third, we recognize the inability of conducting results validation due to unavailability of data. This prevents the assessment of consistency between the results of our model and other evidence and/or the 292 true burden at district level; however, with future availability of data, the model can be continuously 293 294 updated and be validated. 295 296 **Advantages** 297 Within these limitations we achieved our main aim to keep the SUBsET tool simple and intuitive, 298 enabling the rapid dissemination and further country-led adaptation of the model. Using publicly available 299 data also helped the results to be acceptable to the autonomous District Health Office staff. While it is theoretically possible that a more complicated (and effectively 'black box' model^{11,13}) approach could 300 301 have been equally successful as our intuitive and open approach, input from Indonesian stakeholders at 302 the start, and local feedback throughout the process, suggests our judged approach was correct. 303 304 Through the above, SUBsET filled an urgent need within the Indonesia NTP to help inform with- and 305 between-districts discussions. Furthermore, adding variables, or new districts is relatively easy, and shows how SUBsET provides a template for other countries to consider when looking for subnational 306 307 advocacy, provided data are available. 308 309 CONCLUSIONS The transparent modelling approach applied in SUBsET enabled understanding, ownership, and 310 acceptance among the sub-national decision makers in Indonesia. Our approach shows how local data can 311 312 be utilised to estimate subnational burden, thus providing a template for adaptation in other high burden countries to enable them to inform TB policy at the relevant, decentralised administrative level. 313 314 315 Acknowledgements 316 We would like to express our gratitude to all TB staffs, policy makers, and academicians who attended 317 the workshop both at National and Provincial level and contributed to model development and results 318 disseminations. We also gratefully acknowledge Rizka Nurfadila who had prepared the data required for 319 this study.

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- 322 Edine Tiemersma contributed in editing the manuscript. All authors provided critical feedback on
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Table 1. Variable sources and definitions

Variable	Definition	Range	Data Source
Population size	Number of individuals per	13,763 to 5,682,911	Projected Population of
	district		Regency/City 2010-2020,
			Statistics Indonesia
Level of	Proportion of population	0% to 100%	National Socio-Economic
urbanisation	that lives in urban area		Survey 2017
Living floor space	Proportion of individuals	0% to 92%	National Socio-Economic
	who live in a house with		Survey 2017
	less than 8m ² /person		
Junior high school	Proportion of individuals	29% to 76%	National Socio-Economic
completion	who did not complete		Survey 2017
	junior high school or less		
HIV	Proportion of individuals	0% to 23%	National AIDS
	with HIV infection		programme 2012
Air pollution	Proportion of individuals	5% to 100%	Meteorological,
	with air pollution exposure		Climatological, and
			Geophysical Agency
			(BMKG) 2017

410 Table 2. Results from multivariate analysis of 2013/2014 TB Prevalence Survey

RISK FACTORS-TB ASSOCIATIONS							
Variable	Region	Relative Risk	Lower*	Upper*			
Living in urban	Sumatera	1.72	1.22	2.44			
area	Java-Bali	1.32	0.93	1.88			
	Others	1.30	0.92	1.82			
Living in a house	Sumatera	1.50	1.03	2.19			
less than	Java-Bali	1.30	0.83	2.06			
8m²/person	Others	1.15	0.79	1.65			
Not completing	Sumatera	1.11	0.78	1.60			
junior high school	Java-Bali	1.34	0.90	2.00			
	Others	1.61	1.10	2.36			
HIV prevalence	All regions	30	20	45			
Air pollution	All	1.47	1.20	1.80			

^{*} Lower and upper bounds reflect 95% confidence interval. Note, relative risks for HIV prevalence and air pollution

⁴¹² were not available by region, but came from literature. ^{26–29}

The estimates of TB incidence rate per 100k population at district level based on the model, 2017

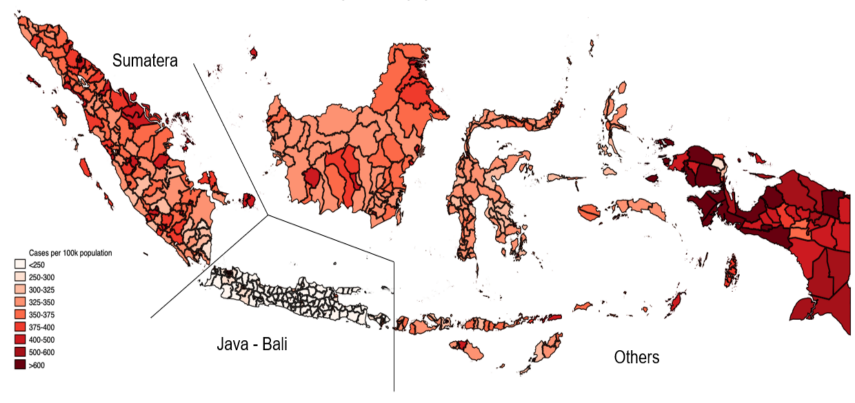


Fig 1. Estimated incidence per 100 000/year by district.

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Figure shows three regions (solid lines) and 514 districts with their estimated incidence per 100,000 population.

The estimated facility-based case detection rate (CDR) by district, 2017

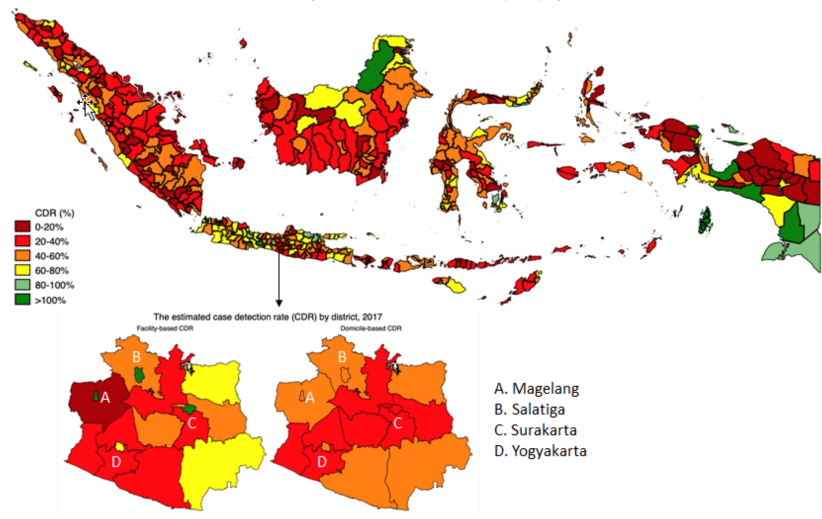


Fig 2. The distribution of the estimated facility-based case detection rate (CDR).

National map shows distribution of estimated facility-based CDR across the 514 districts. Pull-out figure shows very high facility-based CDR (more than 100%, green colour) in central urban districts, and low facility-based CDRs in surrounding districts. When viewed as domicile-based CDR, these differences in CDR are no longer present, highlighting cross-district health system utilisation.

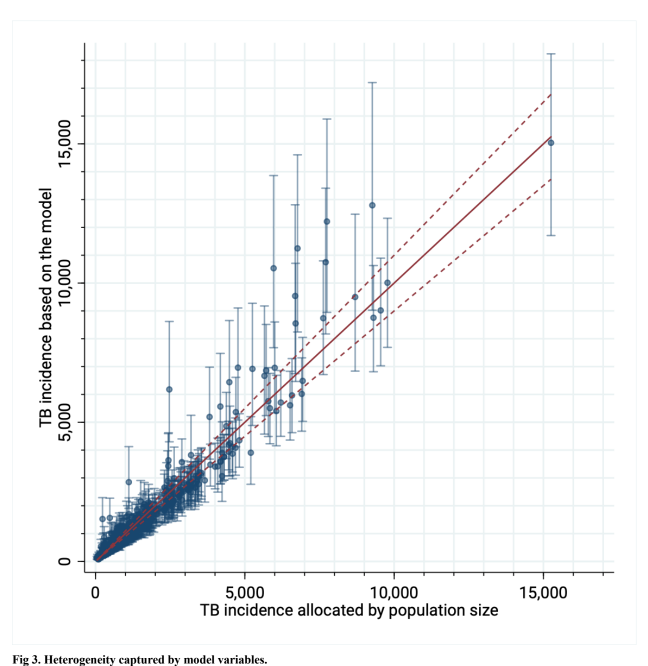


Figure shows change in estimated absolute incidence with 95% uncertainty interval from a model with population size and regional differences in prevalence only (X-axis), and a model from SUBsET (Y-axis). Markers above/lower straight red line indicate districts with a higher/lower estimate based on the full model compared to the simple model.