Waiting times, patient flow, and occupancy density in South African primary health care clinics: implications for infection prevention and control

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1 Abstract

2 Background

- 3 Transmission of respiratory pathogens, such as *Mycobacterium tuberculosis* and severe acute respiratory
- 4 syndrome coronavirus 2, is more likely during close, prolonged contact and when sharing a poorly ventilated
- 5 space. In clinics in KwaZulu-Natal (KZN) and Western Cape (WC), South Africa, we estimated clinic visit
- 6 duration, time spent indoors and outdoors, and occupancy density of waiting rooms.

7 Methods

8 We used unique barcodes to track attendees' movements in 11 clinics in two provinces, multiple imputation

- 9 to estimate missing arrival and departure times, and mixed-effects linear regression to examine associations
- 10 with visit duration.

11 Results

- 12 2,903 attendees were included. Median visit duration was 2 hours 36 minutes (interquartile range [IQR]
- 13 01:36–3:43). Longer mean visit times were associated with being female (13.5 minutes longer than males;
- 14 p<0.001) and attending with a baby (18.8 minutes longer than those without; p<0.01), and shorter mean
- 15 times with later arrival (14.9 minutes shorter per hour after 0700; p<0.001) and attendance for tuberculosis
- 16 or ante/postnatal care (24.8 and 32.6 minutes shorter, respectively, than HIV/acute care; p<0.01).
- 17 Overall, attendees spent more of their time indoors (median 95.6% [IQR 46–100]) than outdoors (2.5% [IQR
- 18 0–35]). Attendees at clinics with outdoor waiting areas spent a greater proportion (median 13.7% [IQR 1–
- 19 75]) of their time outdoors.
- In two clinics in KZN (no appointment system), occupancy densities of ~2.0 persons/m² were observed in
 smaller waiting rooms during busy periods. In one clinic in WC (appointment system), occupancy density did
 not exceed 1.0 persons/m² despite higher overall attendance.

23 Conclusions

- Longer waiting times were associated with early arrival, being female, and attending with a young child.
- 25 Attendees generally waited where they were asked to. Regular estimation of occupancy density (as patient
- 26 flow proxy) may help staff assess for risk of infection transmission and guide intervention to reduce time
- 27 spent in risky spaces.

28 Key words

- 29 tuberculosis; SARS-COV-2; COVID-19; transmission; airborne; nosocomial; infection prevention; healthcare-
- 30 associated infection; health services management

31

32 Background

33	Transmission of respiratory infections is a persistent problem in health care facilities, where proportions of
34	attendees who are infectious and susceptible are likely to be higher than in other settings. ^[1–3] As well as
35	creating risk for individuals attending for care, nosocomial transmission can 'institutionally amplify'
36	epidemics and represents a serious threat to health care worker (HCW) safety. ^[4–7] Pathogens such as
37	Mycobacterium tuberculosis (Mtb) ^[8,9] and severe acute respiratory syndrome coronavirus 2 (SARS-CoV-
38	2) ^[10,11] are more likely to be transmitted during close, prolonged contact and between individuals 'sharing
39	air' in a poorly ventilated space. ^[12,13]

40 Reducing overcrowding and time spent in health care facilities are recommended infection prevention and control (IPC) measures for tuberculosis (TB) and other respiratory infections.^[14,15] Several initiatives to 41 42 reduce frequency of clinic visits have been tested and deployed, primarily with the intention of providing 'differentiated care'.^[16,17] However, interventions that focus on the movement of people around the facility 43 ('patient flow') have received less attention than individual-focused IPC interventions, such as mask-wearing, 44 triage, and prompt initiation of treatment,^[18,19] and 'passive' interventions, such as structural changes to 45 46 improve ventilation.^[20] This is partly because of the complexity of intervening to change patient flow or waiting times in busy facilities, which can vary widely in size, layout, internal organisation, and patient 47 load.^[21] Considerations also differ for hospitals and primary health care (PHC) clinics; this article relates 48 49 mainly to operations at PHC level.

50 Estimating time spent in South African PHC clinics

The nearly 3,500 PHC clinics in South Africa function in diverse epidemiological, political, cultural, and climactic conditions and serve individuals with a wide range of needs.^[22] Long clinic waiting times have been documented over many years^[23,24] and are frequently cited as a concern for patients.^[25–27] The 'Ideal Clinic' initiative, devised and scaled up by the National Department of Health since 2013, aims to enable universal standards of practice, routine measurement of relevant metrics, and fair comparisons of performance between facilities.^[28] Regular estimation of waiting times is recommended by Ideal Clinic and is standard

practice in most clinics, though the methods used vary by province. National guidelines recommend that
 total visit times should be less than three hours.^[29]

59 Most approaches to estimating waiting times conceptualise the patient's journey through the clinic as linear, 60 with each individual passing through the clinic as quickly as possible while ensuring that the necessary 'touch 61 points' are accessed. In South Africa, waiting times are usually measured through the provision of a physical card to all or a selection of patients attending on the day.^[30] This card is time-stamped at the beginning and 62 end of each interaction with a service or 'touch point' (for example, when an individual has their blood 63 pressure measured or sees a clinician). This method is useful for estimating time spent waiting for services 64 65 and the efficiency of selected processes but is less useful in assessing risk of respiratory disease transmission, 66 as it does not describe where patients are waiting. This method also does not include non-patient attendees (e.g., parents accompanying children), any of whom may be susceptible to infection or have undiagnosed 67 68 disease, nor does it allow for estimation of staff exposure to 'high risk' areas.

This work was conducted as part of the *Umoya omuhle* study, a multidisciplinary initiative taking a 'whole systems' approach to TB IPC in South African PHC clinics.^[31] This study component aimed to develop and test a method to 1) estimate how long attendees spent in clinics, and determine why some individuals spent longer than others; 2) estimate how long attendees spend in outdoor and indoor clinic areas; 3) describe variation in occupancy of waiting areas over the clinic day; and 4) collect data for mathematical modelling of IPC interventions in clinics to reduce risk of *Mtb* transmission (McCreesh et al., in preparation).

75 Methods

The literature was reviewed to explore methods previously used to estimate waiting times, occupancy density, crowding, and patient flow (Appendix 1). The methods that allowed estimation of the broadest range of outcomes were waiting/working time surveys (either paper-based or using a real-time location system)^[23,27,32] and camera-based systems.^[33–35] Given the resources available to the study, as well as the ethical and logistical complications of using camera-based systems in multiple public clinics, an approach

81	based on waiting time surveys was used. As described below, unique barcodes and hand-held barcode
82	scanners were used instead of radio-frequency identification tags ^[36–38] or a paper-based system. ^[30]

83 Data collection

84 Data were collected in clinics in KwaZulu-Natal (KZN) province (coded KZN1–6) from 22 February to 14 85 March 2019 and in Western Cape (WC) province (coded WC1–6) from 14–22 May 2019. Clinics were visited 86 at least once prior to data collection to provide written information, discuss concerns with managers and 87 staff, and observe patient flow (Supplementary figure 1). On the day of data collection, a member of the 88 research team attended the morning staff meeting to answer additional questions and issue unique 89 barcodes to staff (including non-clinical staff). Each HCW who accepted a barcode was asked for their job 90 title and role that day. No other personal information was collected. Cards were used that could be easily 91 divided into two after completion to protect confidentiality (Supplementary figure 2). At the top of the card 92 was a unique one-dimensional (1D) barcode and brief instructions. The rest of the card contained a brief 93 questionnaire and the same 1D barcode. After questionnaire completion, the card was divided along a 94 perforated line, the bottom part returned to the researcher, and the top part retained by the participant (or 95 HCW) to be worn, on a lanyard, around their neck.

96 Researchers (usually 6–10 individuals, depending on the size of the clinic) were positioned at key points 97 throughout the premises, including facility entrance(s), the window where patient files were issued (filing 98 station), the triage/vitals station, and doorways of consultation rooms and waiting areas (Supplementary 99 figure 3). Each researcher carried a cordless, hand-held barcode scanner (OPN-2001, Opticon Limited, United 100 Kingdom); clinic staff in certain locations (most often consultation rooms) were also asked to carry scanners 101 and were instructed on how and when to use them. Scanners were cleaned of all data and time-102 synchronised before each day's data collection.

All individuals attending the clinic during the hours of data collection were asked to participate. A researcher
 approached attendees and explained the purpose of the study and that participation was voluntary and

anonymous. Numbers and details of individuals who refused were not recorded because the enrolment
process was time-sensitive. Individuals who agreed to participate were asked to complete the card
(recording their sex, age group, and reason for attendance) and to wear the barcode until they left the clinic.
If requested, the researcher completed the card on a participant's behalf. Individuals attending together had
their cards stapled together before storage; this was accounted for during data entry.

At the beginning of data collection, all individuals already in the clinic were asked to participate and their location noted by use of a designated scanner. Simultaneously, researchers positioned at the entrance(s) asked individuals who were entering the clinic to participate. Within 60–90 minutes of commencing data collection, all individuals were offered the opportunity to participate; this sometimes took longer for busier clinics.

115 Barcode scanners at doorways and other designated 'transition points' were used to scan every person with a barcode who passed through. Scanners at filing or vitals stations (where blood pressure and other 116 measurements were taken), consultation rooms, or at other service points were used to scan an individual's 117 118 arrival and departure from that station or room. Individuals leaving the clinic had their barcode scanned as 119 they left. For logistical reasons, data collection was stopped at 1400; at this point all individuals remaining in 120 the clinic were scanned, with designated scanners used to note their location. A questionnaire was 121 administered to the facility manager or nurse in charge to record information about staffing levels and other 122 factors that may have affected service provision. The dimensions of waiting areas were measured using a 123 Bosch PLR 40R digital laser measure (Bosch, Gerlingen, Germany; accuracy +/-2 mm).

124 Data management

Data from barcode scanners were transferred to a password-protected computer at the end of each day's
 data collection. Data from completed cards were entered into a Research Electronic Data Capture (REDCap)
 database,^[39,40] hosted at the Africa Health Research Institute, that was programmed to assign a 'group ID' to

128	all individuals attending	g together (denoted b	y cards having	g been stapled to	ogether). Data	from questionnaires
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administered to facility managers were entered into a password-protected Excel spreadsheet.

130 Analysis

- Analysis had three strands, examining 1) time spent in clinic and factors that influenced this; 2) the
- 132 proportion of each individual's time in clinic that was spent in indoor spaces (higher transmission risk) vs.
- 133 outdoors (lower risk, primarily due to better ventilation), and how this varied by clinic and reason for visit;
- and 3) in clinics with more than one indoor waiting area, occupancy density of each indoor waiting area and
- how this varied over the course of the day. Supplementary table 3 describes the clinics and numbers of
- 136 individuals included in each analysis.

137 Time spent in clinic

A large number of data were missing for arrival and departure times because several individuals were already in the clinic when data collection began or remained in the clinic when data collection ended and therefore did not have their arrival and/or departure time recorded. Multiple imputation was used to generate arrival and/or departure times for individuals in whom one or both was not recorded (see Appendix 2.2.1 for details).

143 Data were excluded from this analysis where clinic entry and exit were not recorded (because the research 144 team was too small to monitor all entrances and exits of the clinic). Using multiply-imputed data (n = 20 145 imputations), relationships between individual characteristics and time spent in clinic (continuous outcome) 146 were examined using a mixed-effects linear regression model with a random effect for clinic. Province was 147 included as fixed effect. The shape of the relationship between time of arrival and time spent in clinic was 148 examined using fractional polynomials regression with a set of defined powers (-2, -1, -0.5, 0.5, 1, 2, and149 In[x]) and a maximum of two power terms in the model. The differences in model deviances were compared: 150 the linear model was used if the improvement in fit was not statistically significant at p <0.05. Province, age 151 group, sex, and the ratio of patients to clinical staff were included in the multivariable model as a priori

confounders; other variables were included if they showed an important association (p <0.05) in the
 univariable model. Coefficients, representing the difference in mean time spent in clinic (in minutes), are
 reported with 95% confidence intervals (CIs).

155 **Proportion of time spent indoors vs. outdoors**

156 Non-imputed data were used from clinics where a scanner had been positioned at all facility entrance/s and 157 all indoor/outdoor doorways. Individuals with a total captured visit time of less than five minutes were excluded, as they were considered likely to have discarded their barcode. Each individual's pathway through 158 159 the clinic was mapped: for each barcode scan recorded, the individual's location in the time preceding the 160 scan was categorised as 'indoors', 'outdoors', or 'unknown' (if they appeared to have moved between two 161 unconnected locations, indicating a missing barcode scan) based on the location of their previous scan. Total 162 time spent in each type of location (as a proportion of the individual's overall recorded time in clinic) was 163 summarised by clinic and by self-reported reason for clinic attendance.

164 Occupancy density

Non-imputed data were used from clinics that had more than one indoor waiting area and where a barcode scanner had been positioned at all entrances and exits of at least two waiting areas. Data were divided into 10 second slices and entries and exits from each demarcated space noted for each 10 second period; the number of individuals within a space at the end of each 10 second period was divided by the floor area and volume of that space to give the occupancy density (in persons/m² and persons/m³, respectively) for that 10 second period.

Analyses were conducted in Stata versions 14 and 16 (Statacorp, College Station, Tx). Figures were created
 using Stata, Microsoft PowerPoint, Microsoft Excel, and Inkscape v0.92.4.^[41]

173 Ethical considerations

- 174 Identifiable data were not collected from participating individuals; written informed consent was not
- 175 requested. This study received ethical approval from the Biomedical Research Ethics Committee of the

University of KwaZulu-Natal (ref. BE082/18), the Human Research Ethics Committee of the Faculty of Health
Sciences of the University of Cape Town (ref. 165/2018), the Research Ethics Committee of Queen Margaret
University (ref. REP 0233), and the Observational/Interventions Research Ethics Committee of the London
School of Hygiene & Tropical Medicine (ref. 14872).

180 **Results**

181 Patient flow in study clinics was broadly organised around three key steps in the following order: 1) patient registration (file collection); 2) vital signs; and 3) HCW consultation. Individuals usually waited in different 182 183 parts of the clinics for each step. The pathway taken depended on the reason for visit (many individuals also 184 visited one or more of the in-clinic pharmacy, phlebotomist, and other specialist practitioners) and was 185 implemented variably in clinics based on their size, design, and organisation of care. In most clinics, individuals attending for TB care (i.e., those being investigated for TB or taking anti-TB treatment) were 'fast-186 187 tracked' and skipped steps 1 and 2 above. Clinics varied widely in size, population served, services offered, 188 and organisation of care. Importantly, some clinics routinely asked patients to wait in covered outdoor 189 waiting areas, whereas others had only indoor areas. All clinics in WC and no clinics in KZN operated a date-190 time appointment system for at least some patients (Supplementary table 5); no clinics had an active queue 191 management system.

192 Twelve datasets were available for analysis from 11 clinics: six in KZN and five in WC (Table 1; clinic WC4 193 could not be visited for logistical reasons and clinic KZN1 was visited for a second time to attempt better 194 coverage). Data were collected for 2,903 patients and visitors: 1,925 (66.3%) in KZN and 978 (33.7%) in WC). 195 Across clinics, a median 70% (interquartile range [IQR] 69%–74%) of clinic attendees were female. Most 196 individual characteristics were similar between provinces, with the only large differences seen in 'main 197 reason for clinic visit': in KZN clinics, a median 32.6% of clinic attendees reported attending for HIV care or 198 antiretroviral therapy (ART), compared with a median 3.5% in WC clinics. This was thought likely due, at least 199 in part, to an error during data collection in WC clinics, with 'acute care' consistently incorrectly marked by 200 those attending for HIV care (49% in WC vs. 29% in KZN). Because no identifying details of individuals were

- 201 collected, this could not be rectified, and the two categories were combined in analysis (but are shown
- 202 separately in Table 1).
- 203 Table 1. Characteristics of clinics, individuals attending, and staff working on the day of data collection, overall
- 204 and by province (N = 12 exercises at 11 clinics; N = 2,903 attendees)

205 Time spent in clinics

- 206 Data were excluded from clinic KZN4 (n = 269) as all entrances and exits had not been monitored. Data from
- 207 2,634 individuals attending 10 clinics (11 data collection exercises) underwent multiple imputation and were
- included in this analysis (1,063 [40%] missing time of arrival and 934 [35%] missing time of departure;
- 209 Supplementary table 4). Overall median time spent in clinic was 2 hours 36 minutes (IQR 01:36–3:43). This
- 210 was similar in each province (KZN 02:33 [IQR 01:35–3:40; n = 1,656]; WC 02:42 [IQR 01:37–03:49; n = 978]).
- 211 Visit durations by demographics and reason for visit are provided in Supplementary table 6.
- 212 In univariable analysis (Table 2), there was strong evidence of an increase in mean time spent in clinic for
- individuals who were female (p <0.001), attending with a baby (p <0.001), or attending with \geq 1 other person
- 214 (p <0.01). There was also strong evidence of differences by reason for visit (p <0.01): individuals attending
- for TB care and ante/post-natal care spent the shortest time in clinic. Mean time in clinic reduced by ~15
- 216 minutes for each hour that arrival was delayed after 0700 (p < 0.001).
- 217 Table 2. Results of univariable and multivariable mixed-effects linear regression using imputed data, showing
- effects of different factors on total time spent in clinic (n = 2,634; 11 exercises in 10 clinics)
- In multivariable analysis, longer mean times remained associated with being female (13.5 [95% CI 6–21]
- 220 minutes longer than males) and attending with a baby (18.8 [95% CI 8–30] minutes longer than those
- attending without). Reason for visit (p < 0.01) and time of arrival (p < 0.001) also remained important: those
- attending for TB care or ante/post-natal care spent a mean 24.8 (95% CI 9–41) minutes and 32.6 (95% CI 11–
- 54) minutes less in clinic, respectively, than those attending for HIV/acute care, and mean time in clinic

reduced by 14.9 (95% CI 13–17) minutes for each hour that arrival was delayed after 0700. The results of the fractional polynomial models showed that the linear model adequately described the relationship between the time at clinic and arrival time (Appendix 3.2.1).

227 Proportion of time spent indoors vs. outdoors

- The 2,190 clinic attendees included in this analysis (≥5 minutes captured; 10 visits; 9 clinics) spent a median
- 229 95.6% (IQR 45.6–100) of their time indoors and a median 2.5% (IQR 0–35.3) outdoors (Supplementary table
- 230 7), with the remainder in unknown locations. This varied by clinic (Figure 1A): in four clinics with an outdoor
- waiting area that was used as part of normal patient flow, individuals spent a median 13.7% (IQR 1.4–74.5; n
- = 1,362) of their time outdoors, compared with a median 0% (IQR 0–1.4; n = 828) outdoors among attendees
- at the five clinics without an outdoor waiting area or where the outdoor area was not used.
- **234** Figure 1. Box and whiskers plots showing proportions of time spent indoors and outdoors, by clinic (panel A)

and for two visits to clinic KZN1, by selected reasons for visit (panel B)

- 236 In clinics with outdoor waiting areas, the wide IQR (1.4–74.5) for estimated time spent outdoors reflects the
- 237 considerable variation seen among attendees to clinic KZN1, where the outdoor waiting area is used only by
- 238 individuals in the 'chronic' stream. For example, in the second exercise at KZN1, individuals attending for
- 'acute' care spent a median 89.8% (IQR 18.9–98.3; n = 118) of their time indoors, compared with those
- attending for HIV care, who spent a median 98.6% (IQR 92.8–100; n = 125) of their time *outdoors* (Figure 1B).
- 241 Estimates by reported reason for visit for each clinic are provided in Supplementary table 8.

242 Occupancy density of indoor spaces

Data from three clinics were sufficient to estimate occupancy density of at least three indoor spaces (Figure
2). In clinic KZN6 (Figure 2, panel 2), the occupancy density of area A consistently declined over the course of
the day as individuals moved into areas B and C. Because of its relatively large volume, the occupancy
density of area A never went above 0.9 persons/m². In contrast, in the smallest space (area C), occupancy

peaked at around 1200, with density around or above 2.0 persons/m² from 1000–1200. In clinic KZN2 (panel
3), the smaller overall numbers of attendees meant that although the spaces are of similar size to clinic
KZN6, density was generally lower. Overall occupancy was highest in clinic WC1, but the larger waiting
spaces in this clinic meant that occupancy density was never higher than 1.0 persons/m² (panel 4), even in
the smallest space. Clinic WC1 also had a well-functioning date-time appointment system, which is likely why
occupancy of these spaces was more evenly distributed over the day compared with the other two clinics.

- 253 Figure 2. Line graph and heat maps showing, respectively, total numbers of people in and approximate
- 254 occupancy density (in persons/m²) of three indoor waiting areas in each of clinics KZN2, KZN6, and WC1
- 255 between 0800 and 1345
- 256 Occupancy density by room volume (persons/m³) was calculated for the same spaces (Supplementary table
- 257 9). This is a more relevant measure of occupancy density for predominantly airborne pathogens, such as
- 258 *Mtb*. All assessed waiting spaces in clinics KZN2 and KZN6 had relatively low ceilings (maximum height 2.5–
- 259 2.7 m) and occupancy density was higher (median 0.21–1.02 persons/m³) than in spaces in clinic WC1, where
- 260 ceilings were higher (maximum height 4.2–5.9 m; median occupancy density 0.10–0.14 persons/m³).

261 **Discussion**

We tracked 2,903 clinic attendees at 11 PHC clinics in two provinces of South Africa. Median time spent in clinic was 2 hours 36 minutes (IQR 01:36–03:43). People who arrived early in the morning spent longer in clinic, as did women and individuals attending with babies. Individuals attending for TB and maternal care spent less time in clinic. People attending clinics that had outdoor covered waiting areas spent more of their visit time outdoors, though differences were also seen between individuals attending the same clinic based on how care was organised for different 'streams'. In clinics with multiple indoor waiting areas, occupancy was often not distributed evenly between areas or over time; periods of high occupancy density (>2

269 persons/m²) were observed in smaller waiting areas.

270 Time spent in clinic was below the national target maximum time^[29] of three hours for around 60% of clinic 271 attendees (ranging from 48% to 82% across clinics), but was over four hours for around 20% (range 7%–37%) 272 and over five hours for around 9% (range 4%–27%). Detailed comparison with other studies is challenging, given the variation in operational characteristics of PHC clinics and methods used (Supplementary table 10). 273 274 On crude comparison, median time spent in clinic in our study was slightly higher than seen in recent South African studies (Stime et al. [urban KZN, 2016],^[24] median 01:48 for sexually transmitted infection care and 275 median 02:46 for HIV care; Egbujie et al. [rural KZN, 2014],^[42] median 01:56 in nine PHCs) and slightly lower 276 277 than seen in older studies (Bachmann and Barron [urban WC, 1997],^[23] median 2.6 hours and 4.1 hours for 278 'preventive' and 'curative' care, respectively). Patterns in our data were also observed by previous investigators, including longer times for individuals who arrived earlier^[23,24,42] and the early arrival of the 279 majority of attendees, often before the clinic opened.^[30] A higher patient to nurse ratio was strongly 280 associated with longer waiting times in the study by Egbujie et al., ^[42] but not in our study, possibly because 281 282 our estimates of staff numbers included all clinical staff, not only nurses. We are not aware of any previous studies that estimated proportions of time spent indoors vs. outdoors or the occupancy density of waiting 283 284 areas.

Early arrival and queueing outside clinics is common in South Africa. It is influenced by the frequent absence of appointment and queue management systems; the organisation of services around the 'morning rush'; the lack of incentives for staff to change working patterns; and complex factors outside the health system, such as the availability of public transport and the community's trust in the system. Detailed exploration of these issues is beyond the scope of this paper, but some discussion can be found in the report of an *Umoya omuhle* workshop on patient flow that involved a range of South African experts.^[43]

The observed between-clinic and within-clinic variation in proportions of time spent indoors versus outdoors reflects the importance of both clinic design and the organisation of care in moderating the risk of respiratory disease transmission in these settings. The existence of an outdoor, 'low risk' waiting area is of little benefit if most individuals spend most of their time in poorly ventilated indoor spaces. However, the

- use of outdoor spaces may be less feasible in areas with lower temperatures. Thermal comfort and user
 acceptability are important considerations when planning changes to patient flow.
- In clinics where it may be impractical to wait outdoors, risk indoors can be moderated through more even distribution of occupancy throughout the available space. For example, during the busiest period in a small clinic like KZN6 (106 people in the clinic [Figure 3]), restricting occupancy of the smaller waiting spaces (B and C) to 20 and 16 individuals, respectively, would have left 70 individuals in the largest space and resulted in an occupancy density of around 1 person/m² in all three spaces. This is in line with 2014 WHO guidelines for spatial separation as part of IPC for 'epidemic- and pandemic-prone acute respiratory infections', which recommend maintaining a distance of at least 1 metre between patients.^[15]
- South African draft national guidelines^[44] suggest a number of potential interventions to reduce waiting
 times and improve patient flow. Some have been tested in South Africa and other similar settings and are
 discussed briefly below.

307 **Potential interventions**

308 Interventions to improve flow can be classified broadly as targeting two domains: 1) reducing the number of 309 individuals overall and/or in particular spaces; and 2) reducing the time spent by attendees overall and/or in 310 particular spaces. Most measures affect both domains, sometimes indirectly.

Initiatives to reduce numbers of attendees include the Central Chronic Medicine Dispensing and Distribution
(CCMDD) system, where certain groups of patients collect chronic medication from community-based
sites,^[17,45] and reducing the frequency of routine clinic visits for certain conditions, for example by increasing
the amount of medication provided (trials among people taking ART have shown promising results).^[16,46-49]
Measures to improve the overall efficiency of the clinic aim to move people through the facility as quickly as
possible and to reduce the likelihood of bottlenecks in flow. These include holistic approaches, such as

'Lean', ^[50,51] value-stream mapping, ^[52] and other quality improvement methods, ^[53] as well as more targeted
changes in staffing or resources at specific points in clinical pathways. ^[24]

319 Streaming and triage interventions focus on the movement of people once they enter a health facility. In line 320 with Ideal Clinic guidance,^[28] every clinic in our study operated a streaming system that allowed people 321 attending for TB care to bypass many of the steps in the pathway. This is partly intended to reduce the risk of 322 Mtb transmission and is made feasible by the relatively small numbers of people treated for TB at each clinic and because no additional triage process is required. Triage (broadly defined as the process of prioritising 323 patients for care based on their needs)^[54] has also been shown to reduce waiting times in a hospital in South 324 325 Africa, though it was less effective when used in two PHC clinics.^[55,56] Effective triage can be challenging and resource-intensive to sustain,^[57] and sub-optimal implementation of symptom-based triage for TB IPC has 326 been documented by several studies.^[58–61] Active queue management has also been tested: a qualitative 327 328 study around the use of a 'Fast Queue' in clinics in KZN found that the use of multiple, managed queues was 329 generally well-received by attendees, particularly if accompanied by smooth (i.e., unidirectional) flow and 330 effective communication with HCWs, though there were still those who experienced long waiting times.^[62]

331 Date-time appointment systems have been most widely used to reduce both numbers of people and time spent in clinics. Appointment systems have been shown to reduce waiting times in outpatient ART clinics in 332 Ethiopia^[63] and Kenya,^[64] antenatal clinics in Mozambique,^[65] and PHC clinics in South Africa,^[42] the last as 333 334 part of a suite of interventions that included streaming, training, and infrastructure upgrades. Investigators describe generally encouraging results, though they also highlight the considerable challenges involved in 335 336 standardising implementation at facilities that are differently organised. During the Umoya omuhle patient 337 flow workshop, discussions around appointment system implementation emphasised the importance of 338 support processes (such as pre-retrieval of files) and technological infrastructure in sustaining this complex intervention.^[43] 339

340 **Recommendations**

341 Building flexibility into the organisation of flow would allow a clinic to adapt to and absorb periods of 342 increased traffic without putting patients or staff at risk; for example, by moving people from an 343 overcrowded area to one that is relatively empty, or by activating 'overflow' covered outdoor waiting areas. 344 However, any such initiative would require 1) a queue management system, to ensure that individuals 345 moved between areas are not placed at a disadvantage, and 2) clinic managers to have a) easy access to real-time information about flow and b) the resources and freedom to try to improve flow.^[27] Patient flow 346 347 can be difficult to measure quickly: previous published descriptions focus on largely qualitative descriptions 348 of observed movement patterns.^[23,66] Occupancy density, however, is easy to measure (e.g., through manual headcounts) and, measured periodically across a clinic, could be used as a proxy estimate for flow. We 349 350 suggest that regular, light-touch ('diagnostic') approximation of this metric may have numerous potential 351 direct and indirect benefits, including improved efficiency; shorter waiting times; better clinic-specific decision-making; and a strengthened relationship between the clinic and its community.^[27,43,67] 352

353 Importantly, interventions intended to reduce attendance and waiting times may adversely affect the flow 354 around the which the clinic was designed and may therefore increase the rate of transmission to an 355 individual during the time they do spend in the clinic. Most clinics are designed with waiting areas that get 356 successively smaller as patients move through the pathway; as pathways diverge, patients 'diffuse' through 357 the clinic and one would expect occupancy to be lower. However, if the overall 'patient load' is greater than 358 the capacity of the clinic, or if different stages of the pathway are variably efficient, or if certain attendees 359 (e.g., those with appointments) are allowed to skip parts of the queue, bottlenecks can arise in areas that 360 are designed to hold fewer people, leading to higher than optimal occupancy of 'downstream' areas and/or 361 under-use of 'upstream' areas. Interventions to improve flow and reduce waiting times are acutely vulnerable to achieving "many small successes and one big failure"^[68] and should be undertaken with careful 362 363 consideration of potential effects on other parts of the pathway, possible increases in risk of disease

transmission, and adjustments that may be needed in resource allocation, ventilation, and the organisationof care.

366 Limitations

367 The method employed in this study was relatively inexpensive, built on methods already widely used in 368 South African PHCs, and included elements that could be incorporated into routine estimation of waiting 369 times and flow. Numbers of individuals who declined to participate were not recorded and we were 370 therefore unable to assess for selection bias introduced by the enrolment process. Starting data collection 371 after some individuals had arrived and stopping data collection at 1400 (because of logistical restrictions) 372 reduced the numbers of individuals whose data could be used to estimate total waiting time, requiring the 373 use of multiple imputation to deal with missing data. Multiple imputation assumes that the data are missing 374 at random, which means that the observed values can be used to predict the missing values. However, if the 375 assumption is incorrect, the results may be biased. Furthermore, the validity of results derived from multiply 376 imputed data depend on the appropriateness of the imputation model. Future exercises should, at a 377 minimum, continue to record clinic exits for as long as possible. Because of variability between and within 378 clinics, and because data were collected on only one day from almost all clinics, estimates presented here 379 should not be considered representative of the two provinces, types of clinics, or the clinics themselves. In 380 busy clinics in particular, many attendees' barcodes were not scanned every time at every scanning point, 381 and estimates of waiting area occupancy and time spent indoors or outdoors should be treated as 382 approximations. Even so, our headline findings are plausible and consistent with those from other studies.

383 **Conclusions**

Measuring patient flow is important for estimating clinic efficiency and disease transmission risk. In our study, women, individuals arriving early, and those attending with young children spent longer at clinic. Attendees generally waited where they were asked to: using outdoor waiting areas as part of patient pathways increased the proportion of visit time spent outdoors. Occupancy of indoor spaces varied considerably over the day and people often were not distributed evenly throughout the available space.

- Regular, light-touch estimation of occupancy density (as a proxy for patient flow) may help staff to assess for
- 390 the risk of nosocomial transmission and guide the use of interventions to reduce time spent in risky spaces.

391 List of abbreviations

- 392 ART: antiretroviral therapy; CI: confidence interval; HCW: health care worker; hh: hours; IPC: infection
- 393 prevention and control; IQR: interquartile range; KZN: Kwa-Zulu Natal; mm: minutes; *Mtb*: *Mycobacterium*
- 394 *tuberculosis*; NCD: noncommunicable disease; PHC: primary health care; REDCap: Research Electronic Data
- 395 Capture; ref.: reference; SARS-COV-2: severe acute respiratory syndrome coronavirus 2; TB: tuberculosis;
- 396 WC: Western Cape

397 **Declarations**

398 Ethics approval and consent to participate

- 399 Identifiable data were not collected from participating individuals and written informed consent was
- 400 therefore not requested. This study received ethical approval from the Biomedical Research Ethics
- 401 Committee of the University of KwaZulu-Natal (ref. BE082/18), the Human Research Ethics Committee of the
- 402 Faculty of Health Sciences of the University of Cape Town (ref. 165/2018), the Research Ethics Committee of
- 403 Queen Margaret University (ref. REP 0233), and the Observational/Interventions Research Ethics Committee
- 404 of the London School of Hygiene & Tropical Medicine (ref. 14872).

405 Consent for publication

406 Not applicable.

407 Availability of data and materials

408 Data will be available from LSHTM DataCompass. Questionnaires used are provided in the supplementary409 material.

410 **Competing interests**

- 411 The authors declare that they have no competing interests. All authors have completed the ICMJE uniform
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423 Authors' contributions

Conceptualisation	ADG, ASK, AV, HM, KK, NM, TAY
Methodology	ADG, ASK, AV, NM
Formal analysis	ASK, KB, NM
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Data curation	ASK
Writing – original draft	ASK
Writing – review & editing	All authors
Visualisation	ASK
Supervision	ADG, AV, KB
Project administration	ADG, ASK, IG
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424

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601

602

603 Tables and figures

Table 1. Characteristics of clinics, individuals attending, and staff working on the day of data collection,

605 overall and by province (N = 12 exercises at 11 clinics; N = 2,903 attendees)

Characteristic	All clinics, n	KZN province, n (row %)	WC province, n (row %)
Number of clinics	11	6 (54.5)	5 (45.5)
Number of data collection exercises	12	7 (58.3)	5 (41.7)
Hours of data collection, HH:MM	77:02	44:32 (57.8)	32:30 (42.2)
Patients & visitors included	2,903	1,925 (66.3)	978 (33.7)
On the day of data collection	Median (range) per exercise	Median (range) per exercise	Median (range) per exercise
Hours of data collection, HH:MM	06:15 (05:37–07:18)	06:15 (05:40–07:08)	06:15 (05:37–07:18)
Clinical staff working*, n	16 (3–45)	14 (8–45)	17 (3–45)
Administrative staff working*, n	11 (4–21)	10 (7–17)	16 (4–21)
Patients† per clinical staff*, n	14 (5–27)	14 (6–27)	14 (5–18)
Patients and visitors included, n	252 (69–417)	269 (170–417)	144 (69–337)
Proportion female, %	70.0 (56.3–789.7)	71.2 (68.4–74.8)	69.2 (56.3–79.7)
Proportion aged			
0–5 years, %	9.1 (0.7–30.8)	8.4 (7.1–10.1)	10.4 (5.6–30.8)
6–15 years, %	3.4 (0–9.4)	3.5 (1.5–6.2)	3.2 (0–8.3)
16–25 years, %	17.3 (6.3–25.7)	20.1 (15.9–25.3)	15.0 (13.1–17.4)
26–35 years, %	27.3 (19.4–36.2)	27.8 (25.8–32.0)	22.5 (19.4–36.2)
36–45 years, %	18.6 (15.0–35.7)	18.0 (15.3–24.8)	20.3 (15.0–21.1)
>45 years, %	18.0 (7.2–35.4)	18.4 (17.1–22.1)	16.9 (7.2–35.4)
Proportion attending with a baby or very young child, %	11.5 (0.7–36.2)	11.2 (0.7–17.9)	12.1 (10.4–36.2)
Proportion attending with ≥1 other person‡, %	24.9 (1.5–60.8)	22.9 (1.5–35.9)	27.9 (15.4–60.8)
Proportion attending for			
Acute care/minor problems, %	37.0 (9.3–53.5)	28.8 (9.3–41.7)	48.7 (34.8–53.5)
HIV care/ART, %	16.1 (0.8–85.9)	32.6 (14.2–85.9)	3.5 (0.8–7.1)¶
Tuberculosis, %	3.8 (0.6–16.3)	2.2 (0.6–16.3)	9.0 (2.1–13.3)
NCDs (including mental health), %	4.4 (0–16.9)	4.1 (0.4–5.3)	6.9 (0–16.9)
Mother & child§, %	12.4 (0.7–30.4)	14.7 (0.7–19.7)	7.7 (4.2–30.4)
Maternal & obstetric, %	2.9 (0–8.7)	3.2 (0–4.7)	2.7 (0–8.7)
Accompanying a patient, %	14.6 (1.5–22.5)	12.4 (1.5–17.1)	19.2 (10.1–22.5)
Attending for another person, %	2.6 (0–6.3)	2.9 (0–4.3)	1.7 (0–6.3)

*Based on questionnaire administered to manager or senior member of staff; data from clinic KZ04 (including number
 of staff) captured only for HIV/chronic unit

608 +Counted as those who reported a main visit reason that was not 'accompanying' or 'attending for another person'

609 ‡Not including babies and very young children

610 §Includes attendance for family planning

611 Two exercises conducted at clinic KZ01, roughly one month apart

612 ¶Likely due to an error during data collection (see text). 'HIV care/ART' combined with 'Acute care/minor problems' for

613 subsequent analysis.

614 ART: antiretroviral therapy; KZN: KwaZulu-Natal; NCD: noncommunicable disease; WC: Western Cape

615 Table 2. Results of univariable and multivariable mixed-effects linear regression using imputed data,

616 showing effects of different factors on total time spent in clinic (n = 2,634; 11 exercises in 10 clinics)

Variable		Univariable analysis*		Multivariable analysis*			
		Difference in time spent, <i>p</i> minutes (95% Cl)		Difference in time spent, <i>p</i> minutes (95% CI)			
Province							
KwaZulu-Natal	1,656	REF	0.000	REF	0 70 4		
Western Cape	978	-0.03 (-32.2, 32.2)	0.998	-5.1 (-34.8, 24.5)	0.734		
Sex							
Male	783	REF		REF			
Female	1,851	17.0 (9.4, 24.5)	<0.001	13.5 (6.0, 21.0)	<0.001		
Age group							
<16 years	381	REF		REF			
16–45 years	1,703	-7.0 (-17.8, 3.7)	0.250	-5.6 (-17.2, 5.9)	0.332		
≥46 years	550	-10.3 (-22.2, 1.6)		-9.9 (-22.7, 2.9)			
Patients ⁺ to clinical staff [‡] ratio							
<10:1	698	REF	0 724	REF	0.024		
≥10:1	1,936	-6.2 (-41.9, 29.5)	0.734	-3.8 (-36.9, 29.3)	0.821		
Attending with a baby or child aged less than ~15 months							
No	2,271	REF		REF			
Yes	344	25.6 (15.2, 36.0)	<0.001	18.8 (8.1, 29.6)	0.002		
Not recorded	19	17.0 (-23.1, 57.1)		10.0 (-28.8, 48.8)			
Attending with ≥1 other person§							
No	1,983	REF	0.004	REF	0.076		
Yes	651	12.6 (4.0, 21.2)	0.004	8.9 (-0.9, 18.7)	0.076		
Time of arrival							
Per hour later than 07h00	2,634	-15.1 (-17.1, -13.1)	<0.001	-14.9 (-16.9, -12.9)	<0.001		
Reason for visit							
Acute care/HIV care	1,526	REF		REF			
Tuberculosis				-24.8 (-40.6, -8.9)			
NCDs (incl. mental health)				-5.9 (-20.8, 9.0)			
Mother & child (incl. family planning)	297	9.5 (-2.6, 21.6)	0.002	-3.6 (-15.7, 8.5)	0.008		
Ante/post-natal	66	-22.1 (-44.4, 0.1)	0.002	-32.6 (-54.0, -11.2)			
Accompanying	360	2.9 (-7.9, 13.8)		-7.3 (-18.7, 4.2)			
Attending on another's behalf	79	-15.0 (-36.2, 6.3)		-11.6 (-32.0, 8.7)			
Not recorded	4	40.5 (-48.6, 129.5)		29.8 (-55.4, 115.1)			

617 *Mixed-effects linear regression with a random effect for clinic day (i.e., two visits to clinic 1 treated as separate

618 clusters).

⁶¹⁹ [†]Attendees who reported a main visit reason that was not 'accompanying' or 'attending for another person'

620 *Includes enrolled ('staff') and professional nurses, clinical nurse practitioners, clinical and enrolled nursing assistants,*

621 and doctors. Does not include lay-counsellors, peer navigators, community health workers, pharmacists, or

622 nursing/medical students.

623 §Not including babies and very young children

ART: antiretroviral therapy; CI: confidence interval; incl.: including; REF: reference; NCD: non-communicable diseases

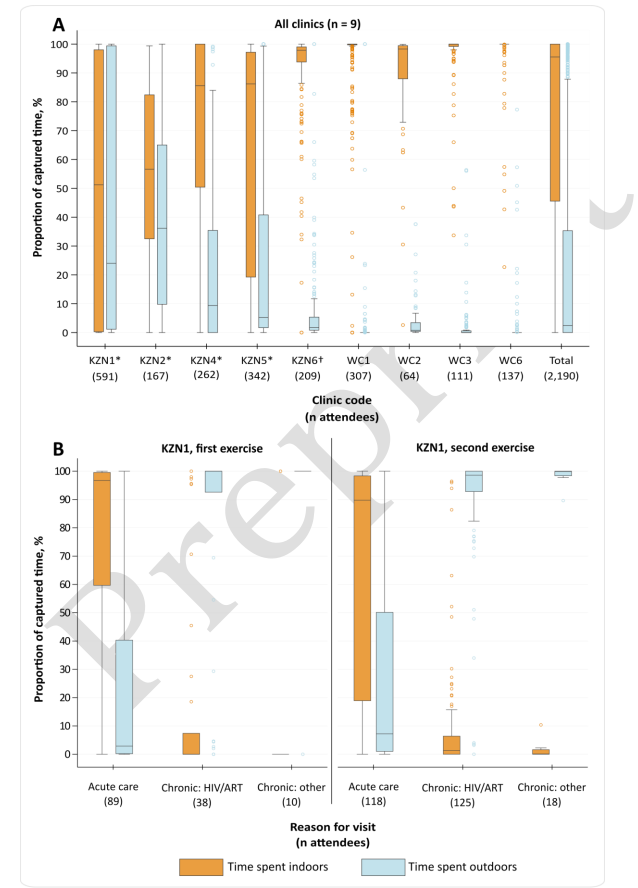
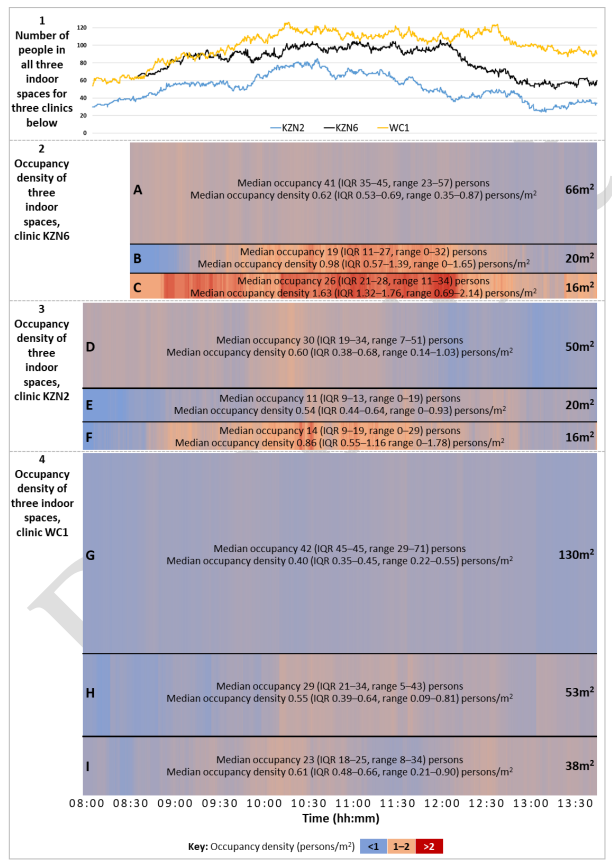


Figure 1. Box and whiskers plots showing proportions of time spent indoors and outdoors, by clinic (panel A) and for two visits to clinic KZN1, by selected reasons for visit (panel B)

627

- *Clinic has at least one outdoor waiting area that is part of the patient pathway.
- 629 [†]Clinic has at least one outdoor waiting area, but it is not part of the patient pathway.
- 630 The central horizontal line represents the median value; boxes represent the interquartile range (IQR); and whiskers
- 631 represent largest and smallest values within 1.5 IQR of the upper and lower quartiles, respectively. Time spent in
- 632 unknown locations was negligible for most clinics and is therefore not shown.
- 633 **Panel A:** Proportions are shown by clinic for all attendees at nine clinics with at least five minutes captured. Data from
- both data collection exercises at clinic KZN1 are shown combined.
- 635 **Panel B:** Proportions are shown by self-reported reason for attendance for individuals with at least five minutes
- 636 captured who were attending clinic KZN1 for the three selected reasons.
- 637 ART: antiretroviral therapy; KZN: KwaZulu-Natal; TB: tuberculosis; WC: Western Cape

Figure 2. Line graph (panel 1) and heat maps (panels 2–4) showing, respectively, total numbers of people
 in three indoor waiting areas and approximate occupancy density (in persons/m²) of each waiting area in
 each of clinics KZN2, KZN6, and WC1 from 0800–1345*



- 642 *Data available only from 0830 for clinic KZN6.
- Height of each row proportional to the area of the space. Each clinic was visited on a different day. See Supplementary
 table 8 for occupancy density relative to room volume (persons/m³).
- Total numbers (line graph) indicative only of numbers of people occupying the three spaces examined, not overall numbers of people in the entire clinic.
- 647 Spaces A, D, and G were the main (pre-filing +/- pre-vitals) formal waiting areas for their respective clinics; spaces B, C,
- 648 H, and I were formal (pre-vitals and/or pre-consultation) waiting areas; space E was a corridor used as a pre-
- 649 consultation waiting area; and space F was a combined pre-vitals waiting area, vitals administration area, and patient
 650 registration area.
- 651 hh: hours; IQR: interquartile range; mm: minutes