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multicentre, randomised controlled trial and text message intervention targeting Effect of a brief motivational interview tuberculosis treatment outcomes in adult patients with tuberculosis: a medication adherence to improve tobacco smoking, alcohol use and of the ProLife programme in South Africa

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> 2022;**12**:e056496. doi:10.1136/ outcomes in adult patients with improve tuberculosis treatment intervention targeting tobacco Kanaan M, Morojele NK, et al. Effect of a brief motivational interview and text message tuberculosis: a multicentre. randomised controlled trial South Africa. BMJ Open smoking, alcohol use and of the ProLife programme medication adherence to To cite: Louwagie G,

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treatment success, medication adherence, alcohol use and interviewing (MI) sessions, augmented with short message trial where participants were assigned (1:1) to the ProLife sensitive pulmonary TB who smoked tobacco or reported Interventions The intervention, delivered by lay health Design Multicentre, individual, randomised controlled behavioural intervention, ProLife, on tuberculosis (TB) Participants 574 adults starting treatment for drugworkers (LHWs), consisted of three brief motivational Setting 27 primary care clinics in South Africa. harmful/hazardous alcohol use. intervention or usual care. tobacco smoking.

additional supplemental material Prepublication history and

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(AUDIT) score, improved TB and antiretroviral therapy (ART)

reduction in the Alcohol Use Disorder Identification Test biochemically confirmed sustained smoking cessation,

months, from TB records. Secondary outcomes were successful versus unsuccessful TB treatment at 6-9

Outcome measures The primary outcome was

alcohol use and tobacco smoking.

who had bacteriology-confirmed TB at baseline, from TB

Results Between 15 November 2018 and 31 August

2019, 574 participants were randomised to receive

6 months by questionnaires; and cure rates in patients

adherence and ART initiation, each measured at 3 and

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Strengths and limitations of this study

Objective To investigate the effectiveness of a complex

ABSTRACT

- The use of motivational interviewing combined with short text messaging to address the effect of multiple risk behaviours (smoking, drinking and poor adherence) on tuberculosis treatment outcomes is a novel and much needed intervention.
- Our study design was strong: this was a multisite, individually randomised controlled trial with a large sample size and a high follow-up rate for the primary outcome.
- We used validated measurement tools; furthermore, assessment were blinded, thereby limiting measurement bias. analysis and primary outcome data
- However, the study was underpowered for secondary outcomes, and low intervention uptake may have diluted any potential intervention effects. lack

service (SMS) messages, targeting medication adherence,

0.76, 95% Cl 0.35 to 1.63), TB medication adherence (OR 1.22, 95% CI 0.52 to 2.87; OR 0.89, 95% CI 0.26 to 3.07), 0.55, 95% CI -1.01 to 2.11; -0.04, 95% CI -2.0 to 1.91) taking ART (OR 0.79, 95% Cl 0.38 to 1.65; OR 2.05, 95% 0.9, 95% CI 0.64% to 1.27%). There was no evidence of an effect at 3 and 6 months, respectively, on continuous Cl 0.80 to 5.27) or AUDIT scores (mean score difference smoking abstinence (OR 0.65, 95% Cl 0.37 to 1.14; OR between intervention (67.8%) and control (70.1%; OR TB treatment success rates did not differ significantly

either the intervention (n=283) or usual care (n=291).

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and adjusting for baseline values. Cure rates were not significantly higher (OR 1.16,95% Cl 0.83 to 1.63).

Conclusions Simultaneous targeting of multiple health risk behaviours with MI and SMS using LHWs may not be an effective approach to improve TB outcomes.

Trial registration number ISRCTN62728852.

INTRODUCTION

Tuberculosis (TB) is among the most common chronic infectious diseases in the world today. In 2019, 1.4 million deaths worldwide were attributed to TB, and the majority of these occurred in low-income and middle-income countries (LMICs). Not only has South Africa one of the highest TB burdens in the world but also it is faced with high TB treatment interruption and loss to follow-up rates. It also has a high prevalence of HIV coinfection in patients with TB and a relatively high mortality in these coinfected patients. Studies of interventions to advance the goal of ending the TB epidemic and improving treatment outcomes are therefore research priorities in South Africa and in other LMICs.

Mortality and morbidity from TB are strongly associated with health risk behaviours, particularly smoking and hazardous or harmful alcohol use, both of which are prevalent and often co-occur in patients with TB. 3-10 Strategies are also required to improve TB medication adherence in patients with TB and adherence to TB medication and antiretroviral therapy (ART) in patients coinfected with TB and HIV, both of which may be negatively influenced by excessive alcohol use. There is very limited research on how to concurrently tackle these three risk behaviours—namely, smoking, harmful alcohol use and poor medication adherence—in patients with TB, particularly in LMICs.

Motivational interviewing (MI) has been shown to support reduced drinking, smoking cessation in patients with TB, and TB treatment and/or ART medication adherence. 11-13 MI interventions can be effectively delivered by lay health workers (LHWs). 14 The more widespread use of LHWs and the increased use of mobile health (mHealth) digital technologies represent promising ways to increase the scalability of MI interventions. Indeed, the WHO has called for researchers to capitalise on advances in mobile phone technology, network coverage and the increased use of common and widely available digital technologies (including the mobile phone short message service (SMS)) to improve TB care. There is evidence that mHealth technologies can have modest beneficial effects on a range of health outcomes, including medication adherence. 16 17 Mobile phone messaging also shows a modest effect in improving TB treatment success rates. 18 19 The evidence is, however, stronger for two-way messaging and interactive systems for which smart phones are required. 18 These are often not available to patients with TB in Africa.²⁰

A limitation of existing MI and mHealth interventions is that they have been studied in the context of modifying a single lifestyle factor. Integrated interventions are likely

to be better accepted and more effective than multiple interventions targeting different health risk factors. ^{21 22} In the case of TB, there is a need for an intervention that has the flexibility to target multiple lifestyle factors as appropriate and in line with patient preferences. This could be achieved through increased integration of TB and noncommunicable disease services. ²³

Recent re-engineering of primary healthcare in South Africa has seen the introduction of municipal ward-based primary healthcare outreach teams of community health workers (CHWs). CHWs work in an integrated, teambased manner, supported by nurses, and take responsibility for health education and promotion, counselling and support for a range of health conditions. Task shifting in this context has been shown to improve population health in LMICs, and these teams can be trained and supported to take responsibility for TB/HIV care. Integrated interventions could be implemented within this framework in a feasible and scalable way to improve outcomes for patients with TB across South Africa and beyond.

Building on previous successes with MI and mHealth interventions, we developed a complex behavioural intervention (ProLife) comprising MI-based counselling and SMS, targeting three lifestyle risk behaviours for poor TB outcomes (smoking, hazardous/harmful alcohol consumption and poor medication adherence) and delivered by LHWs. We then conducted a randomised controlled trial (RCT) to assess the effectiveness of the ProLife intervention on improving TB treatment outcomes, smoking abstinence, reducing alcohol consumption, and improving adherence to TB and ART medication compared with usual care. The cost-effectiveness of the intervention was also assessed, but only the costing results will be presented in this paper.

METHODS

Study design and participants

This was a prospective, two-arm, multicentre, individual RCT which took place across 27 primary care clinics in three districts in South Africa (Lejweleputswa in the Free State province, Bojanala in the North West province and Sedibeng in Gauteng province). Adult patients (18 years or older) were eligible for the study if they had drug-sensitive pulmonary tuberculosis (PTB) and were initiating TB treatment or had been on TB treatment for less than a month for this treatment episode (both 'new' and 'retreatment patients'). They had to be tobacco smokers (defined as smoking daily or non-daily in the last 4 weeks on the Global Adult Tobacco Survey questionnaire)²⁸ and/or hazardous/harmful drinkers who were not alcohol dependent (Alcohol Use Disorder Identification Test (AUDIT) score of ≥ 8 for men or ≥ 7 for women but <20).²⁹ They also had to have access to a mobile phone and understand one of the four languages used for the trial (English, IsiZulu, SeSotho and Setswana). Potential participants were recruited consecutively at the



participating clinics between 15 November 2018 and 31 August 2019. Trained field workers identified those interested in the study and screened them for eligibility. If eligible and willing to be enrolled into the trial, written informed consent was obtained.³⁰

Randomisation and blinding

Patients were centrally randomised (1:1) to the ProLife intervention or control group using a randomised sequence generator by the trial statistician (MK) who was blinded to the arm allocation. We used block randomisation with varying block sizes stratified by clinic so as to achieve equal numbers in intervention and control groups within each clinic. Fieldworkers used sequentially numbered, sealed, opaque envelopes to allocate participants to intervention or control. ProLife involved a complex behavioural intervention; therefore, LHWs and participants could not be blinded to the intervention. However, the determination of the primary outcome was done by the TB nurses who were blinded to the intervention status of the participants, based on routinely collected data. The statistician (MK) was blinded to the intervention or control arm allocation of participants during the analysis.

Intervention and procedures

The ProLife intervention was developed based on a conceptual framework, following a review of pre-existing evidence.³¹ This framework assumed that smoking cessation, reducing harmful alcohol use and improved adherence to TB and HIV treatment would result in improved TB treatment outcomes.³⁰ The intervention consisted of three brief MI counselling sessions, lasting 15–20 min, 1 month apart, delivered by trained LHWs at their TB clinic. The first MI session took place immediately or shortly after the randomisation and involved prioritisation and agenda setting, wherein the participant determined which factor should be prioritised (either a plan to quit tobacco smoking or to reduce or quit drinking, or to deal with barriers relating to ART or TB medication adherence). The second and third sessions built on the previous one until all relevant behavioural problems had been addressed. These sessions were reinforced with follow-up SMS text messages, two times per week over 12 weeks.³⁰ Study patients received 10 TB-related messages followed by seven alcohol reduction-related and/or seven smoking cessation-related messages, as appropriate. Messages were aimed at giving information and augmenting motivation or behavioural skills (we refer to the feasibility paper for more details).³¹ Applicable SMS messages were automatically activated after the first MI had taken place. Thereafter, remaining messages were delivered even if the participant did not attend the second or third MI session.

Participants randomised to the ProLife intervention also received the same 'usual care' as those in the control group. The control group received the usual care and routine treatment and support offered to patients with

TB in South Africa, which vary by district but include health education, dietetic input, social support, point of care biochemical testing, and HIV testing with pretest and post-test HIV test counselling.

Data were collected at baseline and 3 and 6 months and were recorded by fieldworkers equipped with mobile phones with the ProLife mobile data collection application (built with CommCare)³² installed. They used a standardised electronic case report form (CRF) and followed standard operating procedures to ensure quality. Details of data collection, protection and storage procedures were reported elsewhere.³⁰

Patient and public involvement

Patients or the public were not involved in the design, conduct, reporting or dissemination plans of our research.

Outcomes

Primary outcome

The primary outcome of TB treatment success at 6-9 months of follow-up (depending on when it was recorded) was as per the WHO definitions adopted in South Africa, ¹⁰ that is, either successful treatment (cured or treatment completed) or failed treatment, death, acquired drug resistance, loss to follow-up (defined as treatment interruption of more than 2 months) or outcome not evaluated. It was measured using the routinely collected TB treatment outcomes in patients' individual files.

Secondary outcomes

For those participants with bacteriologically confirmed PTB at baseline (either sputum acid-fast bacilli-positive, culture positive or GeneXpert-positive PTB), sputum conversion at the end of treatment ('cure rate') was measured as a secondary outcome. 10 Continuous smoking abstinence was assessed at 3 and 6 months of follow-up in those participants who were current cigarette smokers at baseline. It was defined as having quit smoking completely and a self-report of not smoking more than five cigarettes from the start of the study, in addition to a negative biochemical test (exhaled carbon monoxide (CO) <7 ppm). 33 34 Changes in alcohol consumption were computed using the AUDIT questionnaire scores measured at 3 and 6 months of follow-up in those participants who were hazardous/harmful drinkers at baseline.

HIV-positive participants were asked about ART status at baseline and 3 and 6 months using standardised questions on the CRF and change in ART status as measured at the two follow-up times.

TB and ART medication adherence was measured using modified versions of the AIDS Clinical Trials Group Adherence Questionnaire, a validated tool for measuring adherence specifically to ART. 35 Adherence was measured using an adherence index calculated by the formula (using the 4-day recall): [total number of doses taken/ total number of doses prescribed]×100. Patients with at least 95% adherence were classed as having optimal adherence, and those with less than 95% were classed as

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having low (or suboptimal) adherence. This was assessed at 3 and 6 months.

During COVID-19 lockdown (in the second term of 2020), we switched to telephonic follow-up of participants using a shortened questionnaire whereby only strictly needed information for the measurement of outcomes was inquired about.

Training and intervention fidelity monitoring

The training and intervention fidelity monitoring is described in more detail in previous papers. ^{30 31} In brief, 18 LHWs, 3 district coordinators and 1 research assistant who focused on counselling supervision underwent MI training over 5 days. LHWs completed a postsession semistructured form onto which they indicated the extent to which they implemented each element of MI, as well as their general qualitative impressions of that particular session. In addition, we assessed MI intervention fidelity based on ratings of the counsellors' recorded MI sessions, as described further and in footnotes to the online supplemental table 4. SMS-message delivery was also assessed.

Economic evaluation

The ProLife intervention costs consisted of the costs of training and the delivery of the ProLife intervention, including relevant personnel involvement (trainers and LHWs), materials used, travel, accommodation and refreshments, and digital infrastructure for the intervention. These were estimated based on research team records. Usual care costs consisted of TB medication costs, biochemical investigations and ART costs if applicable. These were estimated based on information obtained through routine records. The country-specific version of EuroQol with five dimensions and three levels of response categories (EQ-5D-3L) for South Africa was administered to participants at baseline and 3 and 6 months of follow-up to measure health-related quality of life. ^{36 37}

Statistical analysis

The sample size was estimated at 696 in total (348 participants per arm) to detect a 10% difference in TB treatment success rates (0.86 vs 0.76) in the ProLife arm (intervention) versus the control arm with 80% power, a significance level of 0.05% and 25% attrition rate. The assumed success rates in the control group were based on actual success rates in patients with TB in the studied provinces obtained from TB managers at the time of the grant application for this study.

We summarised baseline data descriptively by trial arm. For the primary outcome, we conducted statistical analysis on an intention-to-treat basis. We used binary logistic regression to compare the main outcome (TB treatment success rate) between the intervention and the usual care arm. Where treatment outcome data were missing, the outcome was coded as unsuccessful. TB treatment outcomes recorded by the TB nurse were taken on face value as inconsistencies in the dates of bacteriological

results did not permit us to verify the correctness of the nurse assessment. We carried out similar statistical analyses for the secondary outcomes with appropriate regression techniques. For the reduction in harmful or hazardous drinking, we used linear regression to estimate the difference in total AUDIT score between control and intervention groups accounting for the baseline AUDIT score as covariate. Separate analyses at 3 and 6 months were performed.

For our main analyses, we adjusted for baseline characteristics if these differed between trial arms at baseline. The covariates that we controlled for in each model are specified when a model is presented. The statistical packages Stata³⁸ and R³⁹ were used to carry out the analyses, with a p value of <0.05 considered statistically significant.

The validated Motivational Interviewing Treatment Integrity (MITI) coding tool V.4.2.1 was used to assess MI intervention fidelity. The coding entailed making 'global ratings' (on four dimensions: cultivating change talk, softening sustain talk, partnership and empathy) and 'behaviour' counts (with respect to the items giving information, persuade, persuade with permission, question, simple reflection, complex reflection, affirm, seeking collaboration, emphasising autonomy and confront). A score was assigned to each of these items, and the scores were compared against the competency and proficiency thresholds that are specified in the MITI manual.

For the analysis of the costs, all costs were collected in South Africa Rand (ZAR) except for the data management system subscription. Results are presented in both ZAR and US dollar using the 2019 Organisation for Economic Co-operation and Development (OECD) exchange rate (US\$1=14.448 rand). No South African specific valuation set was available for EQ-5D-3L. The valuation set of Argentina, based on a Visual Analogue Scale, was used to derive utility values, because the Gross Domestic Product (GDP) per capita in international dollars was the closest between the two countries at the time of analysis. Unality-adjusted life years (QALYs) were derived from the utility values at the time points by calculating the area under the curve. No missing data imputation was performed.

Data were stored in the institutional data repository at Sefako Makgatho Health Sciences University. Data will be embargoed until 30 June 2023 after which they will be freely accessible. 43

RESULTS

Participant enrolment and follow-up

A total of 2099 patients with TB were screened for eligibility, out of which 574 consenting and eligible participants were randomised: 291 to control and 283 to intervention. Trial recruitment was terminated on 31 August 2019 before the planned sample size was reached because of budget and time constraints. In the intervention arm, 227 (80.2%) participants completed the first MI (MI 1) session; 199 (70.3%) completed MI 2; and

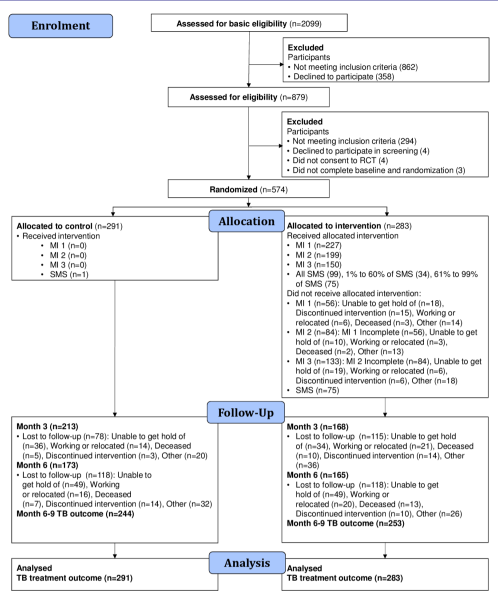


Figure 1 Consolidated Standards of Reporting Trials flow diagram. RCT, randomised controlled trial; SMS, short message service: TB. tuberculosis.

150 (53.0%) completed MI 3. In the intervention arm, at least one message was delivered to 208 (73.5%) participants, while 99 (35.0%) received all messages. Of those randomised to the control and intervention groups, the primary outcome was recorded in 244 (83.8%) and 253 (89.4%) participants, respectively (figure 1).

Baseline participant characteristics of the intervention and control arms

Baseline characteristics were distributed similarly in the intervention and control arms for most variables but with some imbalances in educational level. A total of 513 (91.3%) participants were new patients with TB, 129 (22.5%) women, and nearly all had PTB (International Classification of Diseases-10 A15) without extra-PTB manifestations (553, 98.9%). About half of the participants were HIV positive (305, 53.2%), of whom 204 (65.4%) were on cotrimoxazole and 257 (82.4%) were

on ART (table 1). Details of marital status, employment, wealth, depression status and comorbidities are presented in online supplemental table 1.

There were 372 current smokers (298 daily, 74 less than daily). Seventy-eight participants (26.8 %) in the control arm were dual smokers and drinkers compared with 114 (40.3 %) in the intervention arm. In the control arm, 110 (37.8%) were hazardous/harmful drinkers only and 103 (35.4%) were smokers only, compared with 92 (32.5%) and 77 (27.2), respectively, in the intervention arm (table 2). More details of smoking and drinking history, forms of tobacco use, addiction and quit attempts are presented in online supplemental table 2.

Primary outcome

Overall, 396 (70%) of participants were classified as treated successfully (treatment completed or cured). The remainder either interrupted treatment, failed treatment,

	Control (N=291) n (%)*	Intervention (N=283) n (%)*	Total (N=574) n (%)*
Age (years), mean (SD)	39.37 (12.60)	38.56 (11.15)	
Female sex	69 (23.7)	60 (21.2)	129 (22.5)
Education			
No education	7 (2.4)	5 (1.8)	12 (2.1)
Grades 1–5	23 (7.9)	20 (7.1)	43 (7.5)
Grades 6-7	32 (11.0)	35 (12.4)	67 (11.7)
Grades 8–11	96 (33.0)	128 (45.2)	224 (39.0)
Grade 12	87 (29.9)	70 (24.7)	157 (27.4)
Higher	24 (8.2)	8 (2.8)	32 (5.6)
Declined to answer†	22 (7.6)	17 (6.0)	39 (6.8)
TB patient category			
New patient	264 (92.3)	249 (90.2)	513 (91.3)
Relapse	10 (3.5)	9 (3.3)	19 (3.4)
Retreatment after default	9 (3.1)	14 (5.1)	23 (4.1)
Retreatment after failure	1 (0.3)	2 (0.7)	3 (0.5)
Other	2 (0.7)	2 (0.7)	4 (0.7)
TB site of disease pulmonary only (International Classification of Diseases, ICD-10 A15)	281 (98.9)	272 (98.9)	553 (98.9)
TB sputum smear, Gene XPert or culture result available (N)	236	227	463
At least one sputum smear, Gene XPert or culture result positive	208 (88.1)	195 (85.9)	403 (87.0)
HIV status			
Negative	118 (40.7)	125 (44.2)	243 (42.4)
Positive	163 (56.2)	142 (50.2)	305 (53.2)
Unknown	9 (3.1)	16 (5.7)	25 (4.4)
HIV-positive patients			
Using cotrimoxazole	104 (63.8)	100 (67.1)	204 (65.4)
Using antiretroviral therapy	139 (85.3)	118 (79.2)	257 (82.4)

^{*}Frequencies and (percentages) are presented unless otherwise stated.

developed drug resistance, were transferred out or had an unknown treatment outcome (online supplemental table 3). The percentage of successful TB treatment did not differ significantly between the control and intervention arm (70.1% vs 67.8%), OR for successful TB treatment of 0.90 (95% CI 0.64 to 1.27) comparing the intervention arm to the control arm, and was similar to adjusted ORs (tables 3 and 4).

Secondary outcomes

Cure rates

Among the 403 participants who had at least one positive bacteriological result at baseline, 168 (41.7%) were recorded as cured; of these, 83/205 (39.9%) were in the control arm compared with 85/195 (43.6%) in the intervention arm. The OR of being cured was 1.16 (95% CI

0.83 to 1.63) in the intervention vs the control arm and was similar to the adjusted OR (tables 3 and 4).

Continuous smoking abstinence

Among those who identified as cigarette smokers at baseline (345 (60.1%)), 27 had information (self-report plus biochemical verification) to enable the identification of continuous abstinence at 6 months, of which 22 had continuously abstained from smoking. These were similarly distributed across the two study arms: 10 (5.59%) participants in the intervention arm compared with 12 (7.23%) in the control arm (OR 0.76, 95% CI 0.35 to 1.63) (tables 3 and 4). At the 3-month follow-up, 20 (11.2%) participants in the intervention arm compared with 27 (16.3%) in the control arm continuously abstained from smoking (OR 0.65, 95% CI 0.37 to 1.14) (tables 3 and 5).

[†]More variables with the option 'declined to answer' are listed in online supplemental table 1.



Table 2 Baseline descriptive alcohol and smoking characteristics by study arm

	Control (N=291) n (%)*	Intervention (N=283) n (%)*	Total (N=574) n (%)*
In the past month, smoked tobacco			
Not at all†	110 (37.8)	92 (32.5)	202 (35.2)
Daily	149 (51.2)	149 (52.7)	298 (51.9)
Less than daily	32 (11.0)	42 (14.8)	74 (12.9)
Had a drink in the past 12 months	208 (71.5)	223 (78.8)	431 (75.1)
AUDIT score (men): mean (SD) (max: 19)‡	12.27 (3.98)	13.02 (3.78)	12.66 (3.89)
AUDIT score (women): mean (SD) (max: 19)‡	11.32 (4.02)	10.98 (4.02)	11.15 (4.0)
Hazardous/harmful drinking and smoking combined	(constructed)		
Hazardous/harmful drinking only§	110 (37.8)	92 (32.5)	202 (35.2)
Smoking only	103 (35.4)	77 (27.2)	180 (31.4)
Smoking and hazardous/harmful drinking§	78 (26.8)	114 (40.3)	192 (33.4)

^{*}Frequencies and (percentages) are presented unless otherwise stated.

§Harmful/hazardous drinking is defined as having an AUDIT scores of ≥8 for men or ≥7 for women but <20.

AUDIT, Alcohol Use Disorders Identification Test.

Change in harmful/hazardous drinking

AUDIT scores were about four points lower at both follow-up times than at baseline, independent of the intervention (table 3). In the intervention arm, participants had, on average, a reduction of 0.04 points (95% CI –2.0 to 1.91) on the AUDIT score at 6 months, compared with those in the control arm controlling for baseline scores, whereas an average increase of 0.55 (95% CI –1.01 to 2.11) was observed at 3 months (tables 4 and 5).

Medication adherence and ART uptake

At 6 months, the OR of taking ART medication was 2.05 (95% CI 0.80 to 5.27) comparing the intervention arm to the control arm and controlling for ART baseline medication status, whereas it was 0.79 (95% CI 0.38 to 1.65) at 3 months. The proportion of participants who had optimal TB medication adherence was 90.2% (120/133) at 6 months and 91.7% (319/348) at 3 months. Suboptimal TB medication adherence ORs were 0.89 (95% CI 0.26 to 3.07) and 1.22 (95% CI 0.52 to 2.87) comparing intervention arm to the control arm at 6 and 3 months, respectively. The proportions of participants on ART who had optimal ART medication adherence were high at both 3 months (165/167, 98.8%) and 6 months (139/143, 97.2%) of follow-up. Suboptimal ART medication adherence ORs were 1.17 (95% CI 0.14 to 9.94) and 1.58 (95% CI 0.10 to 26.12) comparing the intervention arm to the control arm at 6 and 3 months, respectively. (tables 3-5)

Intervention fidelity

MI fidelity

The recordings of 17 counsellors (one each) were transcribed verbatim and then assessed. In terms of the global ratings, the LHWs' counselling sessions were above proficiency levels on all items, namely, cultivating change talk, softening sustain talk, partnership and empathy (as the mean scores were all above 2). In terms of the summary measures, the LHWs' counselling sessions did not achieve the basic proficiency threshold of 3.5 for the relational component (partnership+empathy) as their mean score was 3.1 (SD 1.19). However, their mean score on the technical component (cultivating change talk+softening sustain talk) of 3.3 (SD 0.97) was above the threshold of 3. For behavioural counts, 'asking questions' had the highest mean score (24.2, SD 10.42), followed by 'affirm', with a mean score of 5.5 (SD 3.7). The counsellors were least likely to engage in the following: persuade with permission and emphasising autonomy. The mean reflections to questions ratio was 0.23 (SD 0.24). The LHWs made on average 9.3 (SD=4.74) MI adherent (affirm, emphasise autonomy and seek collaboration) and 1.2 (SD 2.28) MI non-adherent (confront and persuade) statements per session (online supplemental table 4).

SMS delivery

Of the total number of information–motivation–behaviour messages triggered, 3583 (80.4%) were delivered. All due SMS messages were delivered to 95 (41.9%)

[†]Non-smokers were included only if they were harmful or hazardous drinkers.

[‡]Only hazardous/harmful drinkers and/or current smokers were included in the study. Therefore, patients with TB were excluded if they were non-current smokers and had an AUDIT score of <7 (women) or <8 (men) or 19; however, they were included if they were smokers independent of whether they had a drink in the past year and therefore independent of the AUDIT score. These AUDIT scores are thus representative of the mean AUDIT scores in the entire study sample and differ from the AUDIT score in the harmful/hazardous drinkers whose change in AUDIT score was measured at 3 and 6 months of follow-up.

	Baseline n (%)*			3 months of follow-up n (%)*			6 months of follow-up n (%)*		
	Control	Intervention	Total	Control	Intervention	Total	Control	Intervention	Total
TB treatment status†									
Successful‡							204 (70.1)	192 (67.8)	396 (69.0)
Not Successful							87 (29.9)	91 (32.16)	178 (31.01)
Cured†,§									
Yes							83 (39.9)	85 (43.6)	168 (41.7)
No							125 (60.1)	110 (56.4)	235 (58.3)
Continuous smoking abstinence¶									
Yes				27 (16.3)	20 (11.2)	47 (13.6)	12 (7.2)	10 (5.6)	22 (6.4)
No				139 (83.7)	159 (88.8)	298 (86.4)	154 (92.8)	169 (94.4)	323 (93.6)
Harmful/hazardous drinkers** (N)	188††	206††	394††	141††	130††	271††	112††	127††	239††
AUDIT score, mean (SD)	12.76 (3.42)	13.12 (3.47)	12.94 (3.45)	8.28 (6.18)	8.84 (5.38)	8.55 (5.81)	8.79 (6.66)	8.70 (5.83)	8.74 (6.22
Difference from baseline, mean (SD)				-4.61 (6.26)	-4.07 (5.33)	-4.35 (5.83)	-4.25 (6.56)	-4.17 (6.61)	-4.21 (6.57
HIV-positive patients	163†† (56.2)	142†† (50.2)	305†† (53.2)	122††	83††	205††	100††	83††	183††
Taking ART medication if HIV positive‡‡	139 (85.3)	115 (81.0)	254 (83.3)	91 (74.6)	58 (69.9)	149 (72.7)	80 (80.0)	74 (89.2)	154 (84.2)
ART medication adherence¶									
Optimal adherence				101 (99.0)	64 (98.5)	165 (98.8)	75 (97.4)	64 (97.0)	139 (97.2)
Suboptimal adherence				1 (1.0)	1 (1.54)	2 (1.2)	2 (2.6)	2 (3.0)	4 (2.8)
TB medication adherence									
Optimal adherence				181 (92.3)	138 (90.8)	319 (91.7)	61 (89.7)	59 (90.8)	120 (90.2)
Suboptimal adherence				15 (7.6)	14 (9.2)	29 (8.3)	7 (10.3)	6 (9.2)	13 (9.8)

Table 3 Descriptive statistics for primary and secondary outcomes by study arm at baseline (where available), 3 months (where available) and 6 months

^{*}Frequencies and (percentages) are presented unless otherwise stated.

[†]Only assessed at 6-months.

[‡]Primary outcome: this is a binary variable defined as either successful treatment (cured or treatment completed) or failed treatment, death, acquired drug resistance, loss to follow-up or treatment interrupted for more than 2 months, or outcome not evaluated/unknown.

[§]Based on having a cured treatment outcome among those who were bacteriologically positive at baseline.

[¶]Assessed at 3 and 6 months; this table refers to cigarette smokers only (other forms of tobacco smoking are excluded).

^{**}Hazardous/harmful drinkers who are not alcohol dependent=AUDIT score of ≥8 for men or ≥7 for women but <20; *! Important distinction at baseline for eligibility purposes.

^{††}Denominator for the mean (SD) or denominator for %.

^{‡‡}Information on HIV positivity was obtained from information from TB records combined with patient self-report at baseline. True HIV-positivity rates may have been higher. ART, antiretroviral therapy; AUDIT, Alcohol Use Disorders Identification Test; TB, tuberculosis.



Table 4 Regression analysis results for the primary and secondary outcomes at 6 months Crude OR (95% CI)* P value* Adjusted OR (95% CI)* P value* Primary outcome TB treatment status: successful (ref: not 0.548 0.90 (0.64 to 1.27) 0.86† (0.60 to 1.24) 0.421 successful) Secondary outcomes Cured (ref: not cured) 1.16 (0.83 to 1.63) 0.374 1.07† (0.76 to 1.51) 0.684 Continuous smoking abstinence (ref: no)‡ 0.76 (0.35 to 1.63) 0.482 TB medication adherence (ref: optimal) 0.89 (0.26 to 3.07) 0.849 ART medication adherence (ref: optimal) 1.17 (0.14 to 9.94) 0.884 Taking ART medication (ref: no) 2.05§ (0.80 to 5.27) 0.136 **AUDIT** -0.04¶ (-2 to 1.91) 0.966 0.02** (-1.55 to 1.6) 0.976

of the participants who completed the first MI (see online supplemental table 5 for more details).

Costs and health-related quality of life

Unit costs used to estimate the mean costs are presented in online supplemental table 6. Incremental cost:utility ratios are not presented since the intervention was not clinically effective. The mean cost of the ProLife intervention was ZAR 2601 (SD 6) (\$180.02, SD \$0.42) per participant in the intervention arm (n=283). The mean cost of usual care was ZAR 681 (SD 357) (\$47.13, SD \$24.71)

in the intervention arm (n=122) vs ZAR 706 (SD 302) (\$48.86, SD \$20.90) in the control arm (n=131). The total mean cost of care including the intervention was ZAR 3285 (SD 357) (\$227.37, SD 24.71) in the intervention arm (n=122). EQ-5D-3L data were available at the three time points for 137 intervention and 159 control arm participants. The mean QALYs estimated over 6 months were 0.442 (SD 0.061) in the intervention arm vs 0.430 (SD 0.074) in the control arm (adjusted mean difference 0.006, 95% CI –0.001 to 0.013).

Table 5 Regression analysis results for secondary outcomes measured at 3 months				
Secondary outcome	Crude OR (95% CI)*	P value*	Adjusted OR (95% CI)*	P value*
Continuous smoking abstinence (ref: no)†	0.65 (0.37 to 1.14)	0.135		
TB medication adherence (ref: optimal)	1.22 (0.52 to 2.87)	0.641		
ART medication adherence (ref: optimal)	1.58 (0.10 to 26.12)	0.750		
Taking ART medication (ref: no)	0.79‡ (0.38 to 1.65)	0.53	0.74§ (0.35 to 1.58)	0.443
AUDIT	0.55¶ (–1.01 to 2.11)	0.474	0.74** (-0.62 to 2.1)	0.273

^{*}Analyses accounted for clustering.

†Given the limited number of those who were identified as continually abstained, we were only able to adjust for one additional variable at a time. Adding one of the following variables: heaviness of smoking, type of drinker at baseline, age when started smoking and the duration of smoking at baseline, the adjusted OR of continuous abstinence comparing the intervention to the control arm ranged between 0.63 and 0.66 with similar confidence limits as for the crude estimate.

^{*}Analyses accounted for potential clustering by centre.

[†]Adjusted for district, sex, and smoking/drinking status and HIV status at baseline. It is worth noting that of the variables in the adjusted model, the only statistically significant result is for the district variable.

[‡]Given the limited number of those who were identified as continually abstained, we were only able to adjust for one additional variable at a time. Adding one of the following variables: heaviness of smoking, type of drinker at baseline, age when started smoking and the duration of smoking at baseline, the adjusted OR of continuous abstinence comparing the intervention to the control arm ranged between 0.73 and 0.76 with similar confidence limits as for the crude estimate.

[§]Adjusting for ART status at baseline.

[¶]Controlling for the AUDIT baseline values; the values represent the study arm regression coefficient.

^{**}Controlling for the AUDIT baseline values and adjusted for district, sex, and smoking/drinking status and HIV status at baseline; the values represent the study arm regression coefficient.

ART, antiretroviral therapy; AUDIT, Alcohol Use Disorders Identification Test; TB, tuberculosis.

[‡]Adjusting for art status at baseline.

^{\$}Adjusted for art status at baseline, district, sex, and smoking/drinking status and HIV status at baseline.

[¶]Controlling for the AUDIT baseline values; the values represent the study arm regression coefficient.

^{**}Controlling for the AUDIT baseline values and adjusted for district, sex, and smoking/drinking status and HIV status at baseline; the values represent the study arm regression coefficient.

ART, antiretroviral therapy; AUDIT, Alcohol Use Disorders Identification Test; TB, tuberculosis.

DISCUSSION

This RCT did not provide evidence for improved TB treatment success rates in those receiving the ProLife intervention compared with those receiving usual care. We could also not demonstrate significant beneficial effects on any of the secondary outcomes, that is, smoking, alcohol consumption, medication adherence and ART initiation. To our knowledge, there are no other published studies of similar complex interventions that aim to improve TB treatment outcomes in patients who smoke or drink to harmful or hazardous extent. Interventions evaluated by other studies were either complex interventions or SMSbased interventions aimed at improving TB outcomes through the pathway of increasing adherence, but without an alcohol or smoking intervention component 44 45 or focused on a single behaviour, namely, smoking or drinking. 46 47 Of the latter studies, a brief smoking cessation intervention was effective in inducing smoking cessation in patients with TB but did not improve TB outcomes. 46 Conversely, in another study in India, intensive counselling for alcohol disorders led to significantly better TB treatment outcomes in the intervention group compared with the control group.⁴⁷ Smoking cessation also led to better TB treatment outcomes in a secondary analysis of a large tobacco cessation trial in patients with TB in Bangladesh and Pakistan. 48 Our non-significant result for smoking-related outcomes is not consistent with findings from our previous TB study, which used a single MI session and found that the chance of sustained smoking cessation was twice as high in the MI intervention group compared with the control group, ¹⁴ although with a less stringent exhaled CO cut-off point. Evidence on the effectiveness of MI for smoking abstinence in non-TB settings has been equivocal.⁴⁹ Self-reported alcohol consumption decreased with about 4 points in both intervention and control arms in our study at both follow-up times. Answering questions on drinking in brief intervention trials may alter subsequent self-reported behaviour: exposing non-intervention control groups to an integral component of the intervention may therefore underestimate the effect of the intervention.⁵⁰ There have been few previous studies looking at MI and SMS interventions for the modification of hazardous/harmful drinking in the context of TB. A previous trial of a brief counselling intervention to reduce alcohol consumption in patients with TB did not find a significant effect on alcohol reduction.⁵¹ Outside a TB setting, results have been mixed. A meta-analysis showed a small but significant improvement in outcomes when MI was used in conjunction with cognitive behavioural therapy for comorbid alcohol use and depression.⁵² Self-reported TB and ART medication adherence was high overall in our study population, which is consistent with other studies conducted in South Africa. 53 54 It is possible that we did not find a difference in treatment adherence due to a ceiling effect.

There were several key strengths in this RCT. This was an individual RCT with a relatively large sample size and a high follow-up rate (87%) for the primary outcome.

Primary outcome assessment was blinded. This was a novel intervention, which built on previous successes with both MI and mHealth interventions and was aligned with the WHO's call to increase the use of digital technologies to improve TB care. 15 We used a validated alcohol consumption questionnaire (AUDIT)²⁹ and a 4-day timeline follow-back for medication adherence to reduce recall bias as self-reports tend to under-report drinking while overestimating adherence behaviour. 35 55 Smoking cessation was confirmed with exhaled CO using strict cut-off points. Overall, the quality of the counselling was acceptable. The results of our MI analyses suggest that the LHWs trained as counsellors were more proficient in MI than during the feasibility stage, as observed by their global rating scores on cultivating change talk, softening sustain talk, partnership and empathy (online supplemental table 4). These results were achieved by ongoing monitoring and training of LHWs during the trial and adapting the training based on feedback from the feasibility stage. Extra counsellors were also appointed to minimise travel distances to clinics. There were some limitations associated with this RCT. Trial recruitment had to be terminated before the planned sample size because of funding and time constraints. Nevertheless, the calculation of sample size was based on an anticipated 25% Loss To Follow Up for the primary outcome, while in reality, only 13.4 %, of the TB outcomes were not available. As a result, we achieved a slightly higher power to detect the a 10% difference in primary outcomes than that we had aimed for (83% vs 80%). The smaller sample size did, however, reduce the power to detect a difference for secondary outcomes for which the LTFU was much higher than 25%. Also, the calculated sample size was not powered for subgroup analysis, which was the case for outcomes relating to smoking, drinking, ART and cure rates. In addition, due to the COVID-19 lockdown in March 2020, we had to switch to telephonic follow-up of participants using a shortened questionnaire (22 participants) and could not access clinics to retrieve outstanding TB treatment outcomes. The low intervention uptake meant that half of the participants received only one or two MI sessions combined with SMS messages. SMS messages were only used for the first half of the study period, and one-quarter of participants did not receive their messages, a commonly occurring problem in LMICs. 20 56 It could be argued that in the absence of ongoing text messages, the MI and associated text messages were not enough to keep participants focused for the second 3 months of the trial. The two-arm study design did not permit the untangling of the individual effects of SMS and MI. Understanding their separate effects could have important cost implications as SMS communication would be cheaper and easier to organise than individual counselling.

The lack of effectiveness of our intervention on the primary outcome (TB treatment success) can have a number of possible explanations. Although intervention uptake was high (80.2%) for the first counselling session, many participants did not return for the second



(29.7%) and third (47%) sessions. As a result of this, only about half of the intervention arm participants received all three MI sessions. Furthermore, about one-quarter of all participants did not receive any SMS messages. Low intervention uptake leads to a dilution of any potential effects. The lack of effectiveness on TB treatment success could perhaps also be explained by the complexity of the ProLife intervention itself: counsellors had to address multiple behaviours, namely, medication adherence, tobacco smoking and hazardous/harmful drinking. Despite having established the feasibility and acceptability of this approach³¹ and ongoing on-site performance monitoring and feedback of counsellors, it is possible that MI for multiple behaviour change in the ProLife study was counterproductive as counsellors may have ended up not focusing on any of the behaviours at optimal levels. Similarly, patients might have found it difficult to change multiple behaviours simultaneously, especially because smoking and drinking are mutually reinforcing. This integrated approach was nevertheless adopted to avoid the need for multiple vertical counselling services (in addition to TB treatment and HIV treatment), to allow the different elements of the programme to reinforce one another, and to improve the affordability, feasibility and acceptability for a future roll-out of the programme. It is also possible that sequential interventions may be better, at least for smoking cessation.⁵⁷ More intensive counselling (more sessions) or a modified counselling method may have been more appropriate, even more so since a recent review of reviews of MI casts doubt on its efficacy. 49 58 59 For example, more emphasis on increasing patient knowledge, in addition to increasing self-efficacy, may have been more effective. 60 The cause of the mhealth message delivery problems (such as poor network coverage and no electricity to charge phones)⁶¹ would need to be investigated in order to increase the effectiveness of future mHealth interventions. Messages may also have to be intensified or modified to be more interactive and/or tailored to specific circumstances of each individual. This would improve the personal value of the intervention to the individual, which is likely to increase the chances of their participation in the intervention.⁶² Consistent with the normalisation process theory, 63 cognitive participation in the intervention might have been higher had we been deliberate in the implementation to ensure the TB nurse, who would have routinely seen the participants, provided additional support and motivated participants to attend MI sessions with the counsellor. In this way, the intervention would have gained 'legitimacy', but this would have led to unblinding of the nurses to the intervention arm.

In conclusion, we could not demonstrate that the ProLife intervention was effective in improving TB treatment outcomes. This may be due to the lack of effect of the intervention, but the study may also have been underpowered for the intermediary secondary outcomes. Valuable lessons were learnt on challenges relating to training LHWs in MI counselling and delivery, SMS delivery in a

challenging socioeconomic context and the reasons for loss to follow-up of TB participants with multiple health problems. Further research is needed to provide answers on how to increase intervention uptake in poor resource settings and whether our complex intervention should have been more intensive. Other important questions are whether another counselling method would have been more effective. Lastly, in the light of the already existing evidence of SMS and the costs and implementation challenges relating to MI, intervention studies limited to an mHealth intervention but using different intensities, duration and type of interventions (one-way, two-way and interactive) are needed.

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Competing interests OAA-Y, the principal investigator, received a research grant (MRC-RFA-02: TB -05-2015) which was allocated to Sefako Makgatho Health Sciences University to pay for the expenses related to the research project, and received support for attending meetings and for travel by Tax Justice Network Africa. He is the vice chair of the Standing Committee on Health, Academy of Science, SA (not remunerated). As coprincipal investigator, KS received part



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Patient consent for publication Not applicable.

Ethics approval This study involves human participants and was approved by the research ethics committees of Sefako Makgatho Health Sciences University (SMU) (ref: SMUREC/D/234/2017: IR); the South African Medical Research Council (with the SMU Ethics Committee serving as the Research Ethics Committee Record); the University of Pretoria (Ref: 434/2017); the University of Witwatersrand (M160455); and the University of York (no reference number, approval date 15 January 2017 (29). Participants gave informed consent to participate in the study before taking part. The trial protocol was previously published.

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Data availability statement Data are available in a public, open access repository. The study protocol was previously published (ImPROving TB Outcomes by Modifying LIFE-style Behaviours through a Brief Motivational Intervention Followed by Short Text Messages (ProLife): study protocol for a randomised controlled trial). The deidentified participant and SMS data sets are stored in labelled Stata files and are accompanied by a statistical analysis plan and metadata explaining each variable. Data will be embargoed for data analysis until 30 June 2023. Thereafter, permission must be obtained from the principal investigators (OAA-Y and KS) for any data analysis not yet performed by the primary research group. Data are stored in the institutional data repository at Sefako Makgatho Health Sciences University called Discover research (https://smu-za.figshare.com/) with a CC-BY 4.0 (Attribution) license (Creative Commons — Attribution 4.0 International—CC BY 4.0). The statistical analysis plan is available as supplementary material.

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SUPPLEMENTARY TABLES

Marital status	Supplementary table 1 Detailed baseline desc	riptive statistics fo	r socio-demogr	aphic, socio-
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	•			
NUMBER OF BYO-TYGOTMONE CMACY TACEC		0 (0.0)	2 (0.7)	۷ (۵.۵)
Number of pre-treatment smear tests recorded?	•			

One	e 197 (92.1	1) 169 (87.6) 366 (89.9)
Two	o 17 (7.9)	24 (12.4)	41 (10.1)
Number of Gene XPert results recorded			
One	e 225 (97.8	3) 210 (97.2)) 435 (97.5)
Two	o 5 (2.2)	6 (2.8)	11 (2.5)
Number of culture results recorded on the T	В		
Treatment record			
One	e 54 (94.7)	42 (97.7)	96 (96.0)
Two	o 3 (5.3)	1 (2.3)	4 (4.0)
Co-morbidities			
Hypertension	n 19 (6.93)	11 (4.1)	30 (5.54)
Diabete	s 5 (1.84)	4 (1.49)	9 (1.66)
Epileps	y 3 (1.09)	4 (1.49)	7 (1.29)
Mental illnes	s 3 (1.09)	0 (0)	3 (0.55)
Liver disease	e 1 (0.36)	1 (0.38)	2 (0.37)
Renal insufficience	y 1 (0.36)	1 (0.38)	2 (0.37)
Allergie	s 2 (0.76)	0 (0)	2 (0.38)
Othe	er 1 (0.36)	1 (0.38)	2 (0.37)
Depression score (CESD 10): mean (SD)	8.44 (4.3	8) 8.74 (4.8)	8.59 (4.59)

^{*}Counts and percentages unless otherwise indicated

	Control	Intervention	Total
	n (%)*	n (%)*	n (%)*
Smoking History (current smokers only) N=	181	191	372
On the days that you smoke, how soon after			
you wake up do you have your first cigarette? After 60 minutes	30 (16.6)	28 (14.7)	58 (15.6)
31-60 minutes	24 (13.3)	16 (8.4)	40 (10.8)
6- 30 minutes	60 (33.1)	65 (34.0)	125 (33.6
Within 5 minutes	67 (37.0)	82 (42.9)	149 (40.1
Duration of smoking in months: mean (SD)	212.09	224.93	218.68
	(134.03)	(127.82)	(130.86)
Duration of smoking in months: median (IQR)	186	206	200.5
	(110, 282)	(135, 294)	(123, 287
Age started smoking in years: mean (SD)	19.2 (6.3)	19.3 (6.3)	19.3 (6.3)
Age started smoking in years: median (IQR)	18 (15-20)	18 (16-21)	18 (15.5-20.
Form of tobacco used			(2010 2011
Manufactured cigarettes (Yes)	166 (91.7)	179 (93.7)	345 (92.7
Number of days you smoked in the past 7days: mean (SD)	5.3 (2.65)	5.53 (2.33)	5.42 (2.49
Average number of cigarettes smoked daily: mean (SD)	6.18 (6.43)	6.48 (8.21)	6.34 (7.39
Hand-rolled cigarettes (Yes)	14 (7.7)	21 (11.0)	35 (9.4)
Number of days you smoked in the past 7days: mean (SD)	4.71 (2.84)	3.71 (2.95)	4.11 (2.92
Average number of handrolled cigarettes smoked daily: mean (SD)	3.71 (3.97)	3.81 (2.82)	3.77 (3.2)
Pipe (Yes)	4 (2.2)	2 (1.0)	6 (1.6)
Number of days in the past 7days you smoked: median (IQR)	1 (0, 2.5)	4.5 (2, 7)	2 (0, 3)
Average number of daily sessions: median (IQR)	1 (0, 3.5)	3 (1, 5)	1.5 (0, 5)
Length of one session (on average) in minutes : median (IQR)	90 (60, 107.5)	60 (30, 90)	90 (30, 90
Cigars, cheroots or cigarillos (Yes)	1 (0.55)	0 (0)	1(0.27)
Water pipe (Yes)	3 (1.7)	2 (1.0)	5 (1.3)
Other	10 (5.5)	4 (2.1)	14 (3.8)
Heaviness of smoking index >= 4	134(74.03)	158(82.72)	292(78.4
Smoking inside your home restrictions			
Total: Not allowed	96 (53.0)	108 (56.5)	204 (54.8)
Some rules: where/when it is allowed	61 (33.7)	58 (30.4)	119 (32.0)

Attempts to quit smoking (current smokers only)			
Ever attempted to quit in the past (Yes)	52 (28.7)	64 (33.5)	116 (31.2)
Number of attempts to quit: mean (SD)	2.46 (2.98)	2.64 (1.62)	2.56 (2.32)
Time elapsed since attempt to quit last time, in months: mean (SD)	36.65 (94.62)	25.13 (37.91)	30.29 (69.22)
Longest duration abstinent in previous quit attempts: mean (SD)	6.15 (13.48)	4.22 (8.9)	5.09 (11.18)
Likelihood to TRY TO QUIT smoking completely and permanently in the next three months			
Definitely will not	6 (3.3)	5 (2.6)	11 (3.0)
Probably will not	10 (5.5)	12 (6.3)	22 (5.9)
Probably will	104 (57.5)	109 (57.1)	213 (57.3)
Definitely will	61 (33.7)	65 (34.0)	126 (33.9)
Likelihood that I WILL QUIT smoking completely and permanently in the next three months			
Definitely will not	6 (3.3)	5 (2.6)	11 (3.0)
Probably will not	11 (6.1)	13 (6.8)	24 (6.5)
Probably will	103 (56.9)	104 (54.5)	207 (55.6)
Definitely will	61 (33.7)	69 (36.1)	130 (34.9)
Ever used any methods to help you stop smoking tobacco in the past 3-months? (Yes)	23 (12.7)	16 (8.4)	39 (10.5)
Smokeless tobacco use (all participants)			
In the past month, have you used smokeless tobacco (Snuff) on a daily basis			
Not at all	275 (94.5)	275 (97.2)	550 (95.8)
Daily	12 (4.1)	6 (2.1)	18 (3.1)
Less than Daily	4 (1.4)	2 (0.7)	6 (1.0)
Duration of using ST in months: mean(SD)	113.69 (112.7)	152 (91.16)	126.46 (105.61)
Age started using ST in years: mean (SD)	27.56 (10.57)	25.88 (13.43)	27 (11.33)
Form of ST used (for SLT users)	(20.07)	(23, 13)	(11.00)
Snuff (by mouth)	2 (12.5)	0 (0.0)	2 (8.3)
Snuff (by nose)	11 (68.8)	5 (62.5)	16 (66.7)
Chewing tobacco leaves	0(0)	0(0)	0(0)
Other	1 (6.3)	0 (0.0)	1 (4.2)
Help to stop drinking (N= , had a drink in past 12 months)	208	223	431
Ever used any methods to stop drinking alcohol in the past 3-months	21 (11.2)	22 (10.7)	43 (10.9)

^{*} Frequencies and (percentages) are presented unless otherwise stated

Supplementary table 3 Detailed descrimonths	ptive statistics fo	or primary outcome	by study arm at 6-
TB treatment status detailed	Control	Intervention	Total
	n (%)	n (%)	n (%)
Cured	108 (37.1)	105 (37.1)	213 (37.1)
Treatment completed	96 (33.0)	87 (30.7)	183 (31.9)
Treatment interrupted > 2 months	15 (5.2)	29 (10.2)	44 (7.7)
Treatment failure	5 (1.7)	2 (0.7)	7 (1.2)
Acquired drug resistance	1 (0.3)	4 (1.4)	5 (0.87)
Died	11 (3.8)	15 (5.3)	26 (4.5)
Transfer out	8 (2.75)	11 (3.9)	19 (3.31)
Unknown	42 (14.4)	25 (8.8)	67 (11.7)

Supplementary table 4 MI treatment fidelity scores of counselling sessions delivered by 17 lay health workers*

Ratings		Mean (SD)	Range
Global ratings**			
	Cultivating change talk	3.2 (1.17)	1-5
	Softening sustain talk	3.4 (0.96)	1-4
	Partnership	3.4 (1.12)	2-5
	Empathy	2.9 (1.34)	1-5
Behaviour counts ^{&}			
	Giving information	4.8 (8.13)	0-5
	Persuade	0.8 (1.48)	0-5
	Persuade with permission	0.1 (0.24)	0-1
	Question	24.2 (10.42)	12-51
	Simple reflection	4.1 (3.21)	0-10
	Complex reflection	1.6 (2.69)	0-9
	Affirm	5.5 (3.47)	1-12
	Seeking collaboration	2.4 (1.46)	0-6
	Emphasising autonomy	1.4 (1.37)	0-4
	Confront	0.4 (0.74)	0-3
Summary measures			
	Total MI non-adherent †	1.2 (2.28)	0-7
	Total MI adherent†	9.3 (4.74)	1-16
	Technical global ratings**	3.3 (0.97)	1.5-4.5
	Relational global ratings**	3.1 (1.19)	1.5-5
	Reflection to question ratio††	0.23 (0.24)	0-0.83
	Percentage of complex reflections***	20.4 (21.9)	0-67

^{*}The recordings of 17 counsellors (one each) were transcribed verbatim and then assessed. In order to assess the fidelity of the counsellors' delivery of motivational interviewing during the trial, we are used the validated Motivational Interviewing Treatment Integrity Coding Manual 4.2.1 (MITI) tool. The table shows the results of the fidelity assessment that was conducted by one rater who is proficient in English, seSotho and seTswana and in motivational interviewing. The rater listened to the recordings and coded a randomly selected 20-min portion of the written transcript. In the case of shorter counselling sessions, the entire recording was assessed.

^{**}The "global ratings" involve assessing, on a 5-point Likert scale (1 for low and 5 for high), how well or poorly the counsellor adheres to the MI practice. Ratings are conducted on four items, two each making up Technical Components (Cultivating change talk and Softening sustain talk) and Relational Components (Partnership and Empathy).

^{**}Scores on Cultivating Change Talk and Softening Sustain Talk are averaged to obtain the Technical global scores.

^{***}Scores on Partnership and Empathy are averaged to obtain Relational global scores. The basic competency threshold scores for fair and good proficiency are 3 and 4, respectively for Technical scores and 3.5 and 4, respectively for Relational scores

[&]amp;The "behaviour counts" involve counting 10 verbal behaviours of the counsellor during the intervention.

^{† &}quot;MI adherence" is determined by adding up the following verbal behaviours: Seeking Collaboration, Affirm and Emphasising autonomy. "MI non-adherence" is determined by summing instances of Confront and Persuade. No thresholds for MI adherence or non-adherence are specified in the MITI 4.2.1.

^{††}The Reflection-to-Question (R:Q) ratio is the total reflections divided by the total questions asked. One reflection to each question is considered a "fair" practice level while two reflections to each question is considered a "good" practice level.

^{***} The Percentage of Complex Reflections (% CRs) is calculated by dividing the number of complex reflections by the sum of complex reflections and simple reflections. A fair and a good % CRs are 40% and 50%, respectively.

Supplementary table 5 SMS delivery					
			Interven	tion (N=283)	Control (N=291)
No. of participants who received ALL due IMB messages INDEPENDER completed MI 1	NT OF WHETHER	they	99/283 (35%)	1/291 (0.3%)
Completion of first MI and initiation of SMS-sequence					
Completed first MI			227/283	(80.2%)	0/291 (0%)
No. of participants who received ALL due messages after receipt of I	MI1		95/227 (41.9%)	0/0 (0%)
SMS delivery for participants for whom the SMS-sequence was initiated (after receipt of first MI)					
	Mean (SD)	Median Range	(IQR)	Mean (SD)	Median (IQR) Range
Average no. adherence messages received per participant (n=227)	7.9 (3.5)	10 (8-10 0-11)	0	0
Average no. tobacco-related messages received (n=153)	5.4 (2.5)	7 (5-7) 0-7		0	0
Average no. alcohol related messages received (n=171)	5.5 (2.5)	7 (5-7) 0-7		0	0
Average no. IMB messages received (n= 227)	15.7 (7.3)	17 (13-2 0-25	2)		

Supplementary table 6 Ur	nit costs used in the analysis (Presented in Rand as sources, \$1=R14.448 in 2019)
Item	Unit costs
Personnel	LHWs: R3000/month, 160 hours/month, R18.75/hour; District coordinators: R7500/month, 160 hours/month, R46.88/hour;
	MRC appointed supervisor: R205.79/hour; Trainers: R373.17/hour; Administrative staff: R333.24/hour
Materials	Training manual: R3250; Printing: R3000; Additional printing & stationary: R500
Accommodation / Travel	LHWs accommodation: R540 000; Trainers accommodation/travel: R36 000; Supervision accommodation/travel: R30 000;
/ Refreshments	Refreshment: R12 500
SMS system	Monthly subscription at R433.44 (\$30) for 3-months: R1300
MI sessions	Session 1 (17 minutes): R5.31; Session 2 (16 minutes): R5.00; Session 3 (17 minutes): R5.31
Biochemical	Cost of TB smear microscopy: R28.37; culture: R79.22 and GeneXpert: R201.56 (Source: 2019/20 National Health Laboratory
investigations	Service pricing schedule)*
ART	Atroiza: R3.78/dose; Dumiva: R5.55/dose; Tenemine: R2.34/dose; Zovilam: R1.72/dose; Kavimun: R1.89/dose; Ricovir:
	R1.36/dose; Zidomat: R1.38/dose; Lazena: R0.55/dose; Efrin:R0.63/dose; Efamat: R0.63/dose; Acriptaz: R0.61/dose (Source:
	Western Cape Department of Health. Antiretroviral and TB Stockmaster Worksheet. 2019)
TB medication	Month's supply of RHZE (intensive) @ R65.80; RH (continuation) @ R55.56 (Source: Western Cape Department of Health.
	Antiretroviral and TB Stockmaster Worksheet. 2019)

PROLIFE Trial

Jan 2020

PROLIFE

Improving TB outcomes by modifying *life*-style behaviours through a brief motivational intervention followed by short text messages (Phase II)

A multi-centre randomised controlled trial looking at the effect of a complex behavioural intervention on TB and lifestyle-related outcomes in South Africa

Version 1.2

Trial Registration number/registry: ISRCTN62728852/ISRCTN registry http://www.isrctn.com/ISRCTN62728852

Department of Health Sciences University of York

University of York York, YO10 5DD **Version date: 14 Feb 2020** 25 Oct 2020

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Trial Coordinator: Goedele M Louwagie

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	University/ University of Pretoria
Funder	Newton Fund/MRC
Funder Reference Number	MRC-RFA-02: TB -05-2015

PROLIFE Trial

Jan 2020

STATISTICAL ANALYSIS PLAN V1.2

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This analysis plan deals only with the statistical analysis of the trial.

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1. Definition of terms

Abbreviation	Expansion
ACTG	AIDS Clinical Trials Group
AE	Adverse Event
ART	Anti-retroviral therapy
AUDIT	Alcohol Use Disorders Identification Test
CI	Confidence Interval
СО	Carbon monoxide
CONSORT	Consolidated Standards of Reporting Trials
CRF	Case Report Form
ЕРТВ	Extra-pulmonary TB
HIV	Human immunodeficiency virus
LHW	Lay health worker
MI	Motivational interviewing
PTB	Pulmonary TB
SMS	Short Message Service
SA	South Africa
SADHS	South Africa Demographic and Health Survey
SAE	Serious Adverse Event
SD	Standard deviation
SOPs	Standard Operating Procedures
ТВ	Tuberculosis

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PROLIFE Trial

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2. Trial Objectives

The PROLIFE model is a complex behavioural intervention comprised of a brief motivational interviewing (MI) counselling strategy augmented with subsequent SMS messaging. To be delivered in three brief sessions, the MI intervention will target three main areas, as appropriate:

- Tobacco smoking
- Alcohol drinking
- Tuberculosis (TB) and Anti-retroviral therapy (ART) adherence or ART initiation

Primary objective:

 To assess the <u>effectiveness</u> of the PROLIFE model delivered by lay health workers (LWH) compared to usual care in improving Pulmonary TB (PTB) treatment outcomes

Secondary objective: (this element will not be addressed in this SAP)

• To estimate the cost-effectiveness of the PROLIFE model

PROLIFE Trial

Jan 2020

3. Design

This is a pragmatic, prospective, multicentre, two-arm, parallel, individual RCT taking place in 27 purposively selected primary care clinics with the highest TB case-load in three districts in South Africa: Welkom in the Free State; Bojanala in the North West province; and Sedibeng in Gauteng province. The intervention will be delivered by LHWs and three district coordinators who will each cover 1–2 clinics.

This is a pragmatic parallel superiority individually randomised controlled trial. There are two treatment arms:

<u>The control arm (Arm1):</u> Intervention arm – participants will receive the PROLIFE programme;

<u>The intervention arm (Arm2):</u> Control arm – participants will receive usual treatment and support provided to TB patients in TB treatment clinics in South Africa ('usual care').

Full details of the background and design of the trial are presented in the protocol (version 1.2 Prolife Protocol_15 Dec 17_with markup) and the published protocol in Moriarty et al (2019) https://doi.org/10.1186/s13063-019-3551-9.

Participants

The inclusion criteria for participants are:

- adult patients (aged ≥ 18 years)
 - with drug-sensitive (bacteriologically or clinically confirmed) PTB;
- initiating TB treatment or on TB treatment for < 1 month (these include both 'new' and 'retreatment' patients);
- current smokers and/or
- hazardous/harmful drinkers who are not alcohol dependent (Alcohol Use Disorders Identification Test [AUDIT] score ≥ 8 for men or ≥ 7 for women but < 20);
- access to a functional mobile phone; and
- understand one of the four languages used for the trial (Sesotho, Setswana, Isizulu or English).

Exclusion criteria:

- alcohol-dependent participants (AUDIT score ≥ 20);
- Extrapulmonary TB without PTB; or
- Resistance to one or more TB drugs at baseline

PROLIFE Trial

Jan 2020

4. Sample Size

We will recruit 696 participants (348 per study arm). The sample size calculations were based on the following assumptions:

- Detection a 10% difference in TB treatment success rates (0.86 vs 0.76) in the ProLife group versus the control group
- 80% power,
- a significance level of 0.05, and
- 25% attrition.

The sample size per clinic was in the range of 14–74 participants per clinic with a median of 24. The assumed success rates in the control group are based on actual success rates in TB patients in the studied provinces that were available at the time of sample size calculations in 2015.

5. Randomisation

Patients will be randomised using a randomised sequence generator performed by the trial statistician (MK) who will remain blind to the arm allocation. We will use block randomisation with varying block sizes stratified by the clinic to achieve equal numbers in intervention and control groups within each clinic. Allocation concealment will be done with consecutively numbered, sealed, opaque envelopes.

Lay health workers delivering the intervention, field researchers, and participants cannot be blinded to the intervention. However, the determination of the primary outcome will be completed by TB nurses who are blinded to the intervention status of the participants based on routinely collected data.

The statistician will be blinded to the intervention or control arm allocation of participants during the analysis stage.

PROLIFE Trial

Jan 2020

6. Outcomes

6.1 Primary outcome

The primary outcome is TB treatment success at six to nine months of follow-up. This is a binary variable defined as

- Success: cured or treatment completed
- Failure: failed treatment, death, acquired drug resistance, loss to follow-up or 'default', or not outcome evaluated.

The different mutually exclusive treatment outcomes are summarised here

Treatment outcome	Definition
Cure	Patient in whom baseline smear or culture was positive at beginning of treatment AND is smear/culture negative in the last month of treatment and on at least one previous occasion at least 30 days prior According to local protocol, a patient who is diagnosed using Gene Xpert and is sputum negative for TB at 11 and 23 weeks is considered 'Cured'.
Treatment completed	Patient whose baseline smear or culture was positive at the beginning and has completed treatment but does not have a negative smear/culture in the last month of treatment and on at least one previous occasion > 30 days prior. Patients diagnosed with PTB whose baseline smear (or culture) result was negative and who started treatment based on clinical and radiological findings who have shown clinical improvement and completed the prescribed course of treatment. N.B. The smear examination may not have been done or the results may not be available at the end of treatment.
Treatment failure	Patient whose baseline smear or culture was positive and remains or becomes positive again at 5 months or later during treatment. Patients who were negative at baseline but were later found to be positive. N.B. This definition excludes those patients who are diagnosed with RR-TB or MDR-TB during treatment.
Died	Patient who dies for any reason during the course of TB treatment.
Treatment default	Patient whose treatment was interrupted for two consecutive months or more during the treatment period.
Transfer out	Patient who was referred to a facility in another district to continue treatment and for whom the treatment outcome is not known.
Acquired resistance	Participants who are subsequently referred for MDR treatment.

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PROLIFE Trial

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6.2 Secondary outcomes

The following outcome measures will be recorded at the six-month follow-up:

- sputum conversion at the end of treatment in the group of participants who had bacteriology confirmed PTB at baseline ¹
- continuous smoking abstinence for identified smokers at baseline²

Whereas, the following will be assessed at three and six months follow-up:

- reduction in harmful or hazardous drinking³
- TB and ART medication adherence will be measured using a modified version of the AIDS Clinical Trials Group (ACTG) Adherence Questionnaire⁴; using an adherence index calculated by the formula (using the four-day recall table):

[Total number of doses taken/Total number of doses prescribed] x 100

Patients with at least 95% of adherence will be considered as having optimal adherence otherwise will be considered as having low (or suboptimal) adherence.

• increase in proportion of HIV-positive participants on ART at three and six months from baseline using standardised questions on the CRF.

6.2.1 Monitoring adverse events

Adverse events (AE) and serious adverse events (SAE) will be defined *apriori* and relevant information will be collected.

A **Serious Adverse Event (SAE)** is actually a special case of an adverse event where adverse outcomes are severe. It includes following events: Death of any of the participants associated with a clinical trial. **Examples of events**: Death, a lifethreatening adverse event, inpatient hospitalization or prolongation of existing hospitalization, a persistent or significant.

An adverse event (AE) is any untoward medical occurrence in a patient or clinical investigation subject administered a pharmaceutical product and which does not necessarily have a causal relationship with this treatment.

The events are reported to ethics committee within 72 hours

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 $^{^{1}}$ i.e. cure rates in intervention group versus control group for participants who initially had sputum AFB-positive, culture-positive or GeneXpert-positive PTB

 $^{^2}$ defined as a self-report of not smoking > 5 cigarettes six months from the start of the abstinence period, supported by a negative biochemical test CO < 7 ppm

³ alcohol use will be measured using the AUDIT questionnaire. The questionnaire will be administered at screening (which will take place on the same day or shortly after the baseline assessment) and again at three months and six months.

⁴ The questionnaire is a validated tool for measuring adherence specifically to ART and we will use an adapted version to also measure TB medication adherence $[\underline{40}]$.

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The following information were collected to report the events:

Participants identification number, Gender, Age, Date of Enrollment, Arm (Control or intervention), Date of death notification to staff, Date of death , If death is related or not related to study.

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6.3 Trial assessment schedule

Table 2 details the trial assessment schedule

Table 2: Trial assessment schedule								
A	Items	Pre randomisation	ation Timel		ine (post randomisation)			
Assessment		Eligibility (Day 0)	Baseline (Day 0)	2 months	3 months	6-9 months	Trial end	
ELIGIBILITY								
Smoking status		X						
Smoking profile			X					
Alcohol profile		X						
Medical eligibility		X						
Eligible, consenting		X						
MEASURES								
Trial ID, visit date			X					
Socio-demographic history			X					
Depression screen			X		X	X		
Clinical review of TB Treatment record for disease information			X				X	
Smoking history			X					
Smoking abstinence (self-report)					X	X		
Exhaled CO					X	X		
Record sputum culture or smear or Gene Expert result			X	X	X	X		
HIV Status			X					
ART Status (if HIV positive)			X		X	X		
AUDIT		X			X	X		
Modified ACTG (Follow up)					X	X		
Economic evaluation			X		X	X		

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6.4 Other important information

In addition to the above, the following information was collected at baseline. Socio-demographic history included age, gender, marital status, education, employment status, and comorbidities. For details about these variables, see Table 3.

6.5 Fidelity of the intervention

The main fidelity analysis will be published somewhere else. Some descriptive statistics regarding fidelity will be added here once the main analysis is completed and the statistician is unblinded to the treatment arm.

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7. Data

7.1 Data collection methods

The fieldworkers will screen all TB patients for eligibility immediately after the TB nurse at the clinic has initiated TB treatment and opened the TB "blue card". Consent will be obtained for this screening phase as the alcohol related questions are sensitive and the fieldworkers must gain insight in the patient files.

Eligible and consenting patients will be enrolled in the trial and the baseline questionnaire and record review completed. Patients will be given a unique Trial Number that will be used on all research documents. Data will be collected and recorded by field workers equipped with Android phones with a mobile data collection application installed.

Participants in the control arm will continue with the routine TB care. Intervention arm study participants will be referred by the fieldworker to the lay counsellor for motivational interviewing. The first MI session will be on the same day of the completion of the screening questionnaire, where possible (with a 2-week window period). The second and third MI session will be scheduled 4 weeks and 8 weeks from the first counselling session respectively each time with a 2-week window period.

MI counselling and data collection will take place in a well-ventilated private area inside or outside the clinic, and audio-recorded after consent obtained. Fieldworkers and LHWs will be provided with high particulate respirator masks to minimise the risk of infection.

Fieldworkers will follow-up all participants in both arms at 3 and 6 months within a window period of 2-weeks before and 2-weeks after the ideal 3 and 6-month visit. Participants will receive SMS reminders 3 days before each planned visit. Participants will also be in a position to send "please call me "messages to the fieldworkers or district coordinators, who will then call the participant to solve problems that may have arisen with the appointment.

Patients who did not return for the planned 3 and 6 months visit will be contacted by telephone up to 3-times, as needed. Home visits will also be undertaken by existing clinic "tracer teams" or Community Based Outreach teams -where feasible - for participants who cannot be traced by telephone. The data collection process is illustrated in Table 3.

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Time-point	Information required at time-point	Data collection method
Baseline	1. Socioeconomic and demographic status (to	
interview	include history of mine work)	
	• Age	TB Treatment
	• Gender	Record
	Marital status	Record
	Educational level achieved	
		CDE Questions
	• Employment	CRF – Questions
	 Mine work/type of mine work 	to participant –
		demographic
	2. Clinical information:	details
	 Patient category (First episode vs recurrence) 	
	 Site of disease 	
	 Results of sputum smear, culture and Gene 	
	Xpert	CRF – Questions
	HIV status	to participant and
	ART information	information from
	 Co-morbidities 	the TB Treatment
	Co morei uiu es	Record (as
	3. Current smoking status and quit history, second hand smoke exposure	indicated in CRF
	4. Alcohol history	
	5. Depression	
		Questions adapted
		from Global Adu
		Tobacco Survey
		Questionnaire
		Baseline AUDIT
		score
		CES-D
3 months	1. ART information	CRF – Question t
	1. The momunon	patient
	TB and ART medication adherence	patient
		ACTC
	(modified ACTCG)	ACTG
	2 41 1 11' (() 41'DIT	questionnaire for
	3. Alcohol history (repeat AUDIT)	both TB
		medication and
	4. Smoking history	ART
	5. SLT use	
		AUDIT score at 3
		month

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	6. Depression	
	•	Follow-up questions as per Russell's Standard and exhaled CO
		CES-D
6 months	1. TB treatment status (<u>Primary outcome</u>)	TB treatment outcome from TB Treatment Record
	2. Sputum smear or culture result	combined with information from
	3. ART information	TB record on cultures and smear
	4. TB and ART medication adherence	results.
	5. Alcohol history (repeat AUDIT)	CRF- Question to patient
	6. Smoking history and exhaled CO	Follow-up ACTG questionnaire at 6
	7. Depression	months
		AUDIT at 6 months
		Questions as per Russell's Standard and exhaled CO
		CES-D

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7.2 Data management

Sefako Makgatho University (SMU) appointed a data manager who will utilise an electronic platform for data collection, ensuring data quality, and facilitating the SMS messages.

Fieldworkers collecting research data will be equipped with Android mobile phones, which will have a mobile application installed on them to allow for data collection in areas with poor internet connectivity. The electronic data captured will be stored on secure and password protected storage servers and mobile phones, which ensure data privacy through only allowing authorised research staff access to the data.

The electronic data collection system used for the study requires an SMS gateway to send and receive messages to the research participants. Consenting participants' phone numbers, participant IDs, and associated SMS messages will be stored on the SMS gateway's secured and password-protected server.

Data quality will be ensured by providing fieldworkers with standard operating procedures (SOPs), training, and ongoing support on the importance of data quality, data collection, and data collection problem-solving. The data manager will continuously monitor the captured data for missing variables and inconsistencies in order to resolve any data problems.

The data manager will export the data from the secured server, conceal the participants study arm allocation, and de-identify the data before sharing the data in STATA and R compatible formats. The exported de-identified data will be stored in Dropbox, a secure cloud storage platform, for sharing with the lead trial statistician at the University of York for analysis.

All research data and documents referring to the PROLIFE trial will be stored and maintained in a secured storage space at SMU for a minimum of 15 years from the end of the PROLIFE trial. Study materials will be destroyed 15 years after the study.

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8. Analysis

The computer packages STATA 16 (StataCorp. 2019) and R 3.5.3 (ref) will be used. Significance tests will be two-sided and the significance level is set at 0.05. The statistician will remain blind to allocation until results are finalised. We will follow the CONSORT statement guidelines in reporting.

Below, we detail the analyses that we will carry out for the data collected at baseline, the primary outcome, the secondary outcomes, and adverse events. We also list the sensitivity analyses that we might perform and subgroup analyses.

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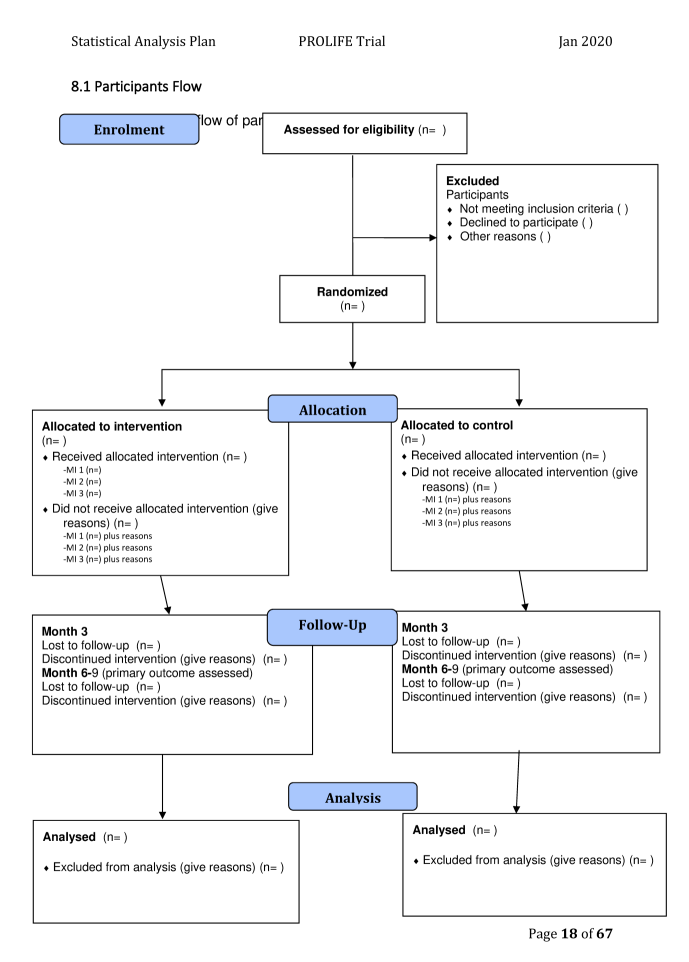


Figure 1 CONSORT DIAGRAM SHOWING FLOW OF PARTICIPANTS

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8.2 Baseline data

Baseline data Analysis Plan:

Baseline data including demographic variables will be summarised descriptively by trial arm, but no formal statistical comparisons will be undertaken. Continuous measures will be reported as means and standard deviations (SD) while categorical data will be reported as counts and percentages, see Tables 1 to 4 in Appendix A. For skewed continuous measures, we will also provide medians and interquartile ranges.

Baseline data results:

Consent:

In the control arm (Arm1) 286 gave written consent and 5 verbal consent for participation in the study and for access to their medical records

In the intervention arm (Arm2) 279 gave written consent and 4 verbal consent for participation in the study and for access to their medical records

Baseline Imbalances

Education:

It seems that for education there is imbalance between the two arms for Grades 8-11/Grade 12/Higher; the control arm (Arm1) had a higher educated group the difference in percentage points 8.2% as opposed to 2.8% in the intervention arm, see Table 2 for further details. There are 9 participants who are not literate in the study (7 in the control arm (Arm1) and 5 in the intervention arm (Arm2)).

Drinking

In the intervention arm, 223 (78.8%) had a drink in the past 12-months compared to 208 (71.5%) in the control arm, see Table 3.

In the control arm (Arm1), 110 (37.8%) were drinkers only, 83 (28.5%) were smokers only, and 98 (33.7%) were smokers and drinkers compared to 92 (32.5%), 60 (21.2%), and 131 (46.3%) in the intervention arm (Arm2), respectively.

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Assets:

The vast majority had a radio (84%), a television (89%), a refrigerator (85%), an electric or gas stove (86%), and a microwave (67%), however, only 50% had a washing machine. Furthermore, a minority had a landline telephone (7.5%), a desktop or laptop computer (20%:), and a vacuum cleaner or floor polisher (15%). Figure 2 is a boxplot of the total number of assets by study arm. The spread in the control arm (Arm1) is greater than that in the intervention arm (Arm2). However, the mean number of assets is similar across the two groups of 5 (SD: 1.87).

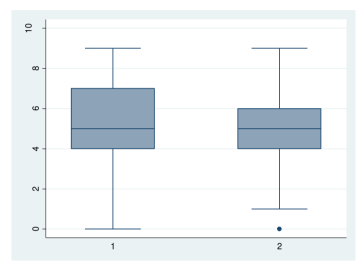


Figure 2: A boxplot of the total number of assets by study arm

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Medical History:

There were 305 (53.2%) **HIV positive** participants out of which 204 (65.38%) were using Cotrimoxazole and 257 (82.37%) were undergoing anti-retroviral therapy, see Table 2.

TB History: The vast majority of participants were New TB patients 513(91.3%), with Pulmonary only (ICD-10 A15) being the site of disease for the vast majority 553 (98.9%). Among those with results known, the majority had one pre-treatment smear result 366 (89.9%), one Gene XPert recorded 435 (97.5%), and one culture result recorded 96 (96%).

Among those with results available, 220 (58.51%) had at least **one positive smear** result,362 (87.23%) had at least **one positive Gene XPert result**, and 35 (47.95%) at least **one positive culture result**.

However, 85 (53.46%) in the control arm had at least one positive smear result within 60 days of the TB treatment start date compared to 96 (61.15%) in the intervention arm.

In addition, only 34 had their culture results within 60 days from the TB start dates of which 21 had positive results; 11 in the control arm compared to 10 in the intervention arm.

The vast majority of participants did not have any **co-morbidities** 525 (96.3%) and 18 (3.3%) had one.

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Baseline Smoking Related variables:

Thirty five percent (202) did not smoke in the past month, whereas 52% smoked daily and the remaining 13% smoked in the past month but less frequently than on a daily basis, see Table 3.

In the past 30 days, the vast majority were exposed to smoke for seven days or less at: home 462 (80.5%); cafes/restaurants 464 (80.8%); Shebeens, bars or clubs 432 (75.3%); Bus/train/taxi/ vehicle 448 (78.0%); and Shops/shopping mall 461 (80.3%) and where applicable at the workplace 173 (66.8%).

Among those who smoked in the past month 372: 181 in the control arm (Arm1) vs 191 in the intervention arm (Arm2),

345 (92.7%) **smoked manufactured cigarettes** of which 225(65%) did so on a daily basis in the past seven days whereas 40 (11.6%) did not smoke in the past week. The mean number of days smoked was 5.42 days in the past week (SD: 2.49) with 6.34 cigarettes (SD: 7.39) smoked daily on average.

Hand-Rolled cigarettes were used by 35 participants of which 16 (45.7%) smoked daily in the past week. They smoked on average for 4.11 (SD:2.91) days in the past week and on average smoked 3.77 (SD: 3.27) hand-rolled cigarettes per day.

There was only one person who exclusively smoked waterpipe, two who exclusively smoked pipe, and seven who exclusively used other formats of tobacco other than the ones that are listed here.

For the vast majority the total number of cigarettes smoked is based on their answer to manufactured cigarettes. In 27 cases, they supplemented this with other sources and 8 used only other forms to report the number smoked on average per day.

Over the past 3 months, they **spent** on average 174.58 (SD: 181.61) Rands per week on tobacco products.

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Smoking restrictions

No smoking was allowed **inside home** for 204(54.8%) participants, whereas 119(32.0%) had some rule where/when it is allowed and 49(13.2%) had no rules in place.

Quit Attempts:

Among smokers, 116 (31.2%) made an **attempt to quit**; the mean number of attempts to quit was 2.56 (SD: 2.32).

Furthermore, 213 (57.3%) said they will probably **try to quit** smoking completely and permanently in the next three months and 126 (33.9%) said that they definitely will.

Whereas, 207 (55.6%) said they will probably **quit** smoking completely and permanently in the next three months and 130 (34.9%) said that they definitely will.

Only 39 (10.5%) have ever used any methods in the past 3 months to help them stop smoking tobacco. These spent on average 136.64 (SD: 205.36) Rands on methods to help you stop smoking in the past 3 months.

Smokeless tobacco was used by only 24 participants (4.2%). They have been using ST for an average of 10.5 years (SD: 8.8) and have started using it at the age of 27 (SD: 11.33) years on average.

Heaviness of smoking:

In the control arm (Arm1), 67(37.0%) of smokers reported smoking within 5 minutes of waking up whereas 82(42.9%) did so in the intervention arm (Arm2).

Among those who smoked and who reported the number of cigarettes/pipes/cigars they used on average per day, 134(74.03%) in the control arm (Arm1) and 158(82.72%) in the intervention arm (Arm2) were considered as heavy smokers.

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8.3 Primary analysis

8.3.1 Primary outcome definition:

As per section 6.1, we would define the primary outcome as those who had a successful treatment versus not. Where successful treatment is considered if the patient is considered to have been cured or treatment completed; all other categories will be deemed as not successful. The successful treatment⁵ categories are defined as follows:

Cure	Patient in whom baseline smear or culture was positive at beginning of
	treatment AND is smear/culture negative in the last month of treatment
	and on at least one previous occasion at least 30 days prior
	According to local protocol, a patient who is diagnosed using Gene
	Xpert and is sputum negative for TB at 11 and 23 weeks is considered
	'Cured'.
Treatment	Patient whose baseline smear or culture was positive at the beginning
completed	and has completed treatment but does not have a negative
	smear/culture in the last month of treatment and on at least one
	previous occasion > 30 days prior. Patients diagnosed with PTB whose
	baseline smear (or culture) result was negative and who started
	treatment based on clinical and radiological findings who have shown
	clinical improvement and completed the prescribed course of
	treatment.
	N.B. The smear examination may not have been done or the results
	may not be available at the end of treatment.

8.3.2 Primary outcome Analysis

For the primary outcome, we will conduct analysis on an intention-to-treat basis. We will use binary logistic regression to compare the main outcome between the

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⁵ Following discussion with the team, we will take this at face value as it is not possible to query some of the anomalies found in the recording of dates of tests.

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intervention and the usual care arm. We will also investigate any potential clustering at the centre level and account for it. We will present the results for this analysis in Table 6.

We will also adjust for HIV status, sex, alcohol versus tobacco versus both, and district; if these differ between trial arms at baseline.

8.3.3 Primary outcome Analysis Results

Table 4 gives the descriptive statistics for the initial outcome and the derived - dichotomised outcome; these indicate that overall, 69% were classified as successful treatment based on the medical professional assessment (cured/treatment completed). This percentage was similar for the two arms with the control arm (Arm1) having a slightly higher percentage of success of 70.1% compared to 67.8% in the intervention arm (Arm2)⁶. Table 9 gives the distribution per centre by study arm. Generally, these are fairly balanced any imbalance observed is most probably due to the early termination of the study.

For 203 participants, the TB treatment outcome date was not available, with those participants more likely to not have been cured (59% not cured), however, the percentage was similar across the two study arms for those with missing TB treatment outcome date.

The odds of successful treatment is 0.9 (95% CI: (0.64,1.27)) in the intervention arm (Arm2) compared to the control arm (Arm1). This estimate is very similar to the estimate adjusting for district and drinking/smoking status. This is also the case if you further adjust for sex and HIV status at baseline where the OR is 0.86 (95% CI: (0.60,1.24)); see Table 6 for further details.

⁶ Primary outcome Control: 204/291, 95% CI proportion 0.70 (0.64,0.75) Primary outcome Intervention:192/283, 95% CI proportion 0.68 (0.62,0.73)

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8.4 Secondary analyses

Secondary analyses plan

In the group of participants who had bacteriology confirmed at baseline, we will use logistic regression to compare cured versus not cured, as indicated by the outcome at the end of treatment, between the two study arms. We will also control for baseline characteristics and other covariates such as sex, alcohol use, HIV-status, district, and account for any potential clustering by centre.

We will use a similar approach for the six-months continuous smoking abstinence outcome. This analysis will be performed on the group of participants who were current tobacco smokers at baseline. We will also control for baseline characteristics and other covariates such as age, duration of smoking, alcohol problem (hazardous, harmful, non-drinker/light drinker), heaviness of smoking index⁷, depression, and potentially HIV-status.

For the reduction in harmful or hazardous drinking, we will use linear regression to measure difference in total AUDIT score between control and intervention groups accounting for the baseline AUDIT scores. Separate analyses for the AUDIT at 3 and 6 months will be performed.

The <u>AUDIT is a 10-items questionnaire</u> with a range between 0 and 40 where higher values indicate higher dependency.⁸ It is worth mentioning that a score of 8 or more

⁷ [Goedele's Comment on an earlier Version]Definitely a measure of severity of smoking and duration of smoking. For example Heaviness of Smoking Index which can be derived as follows:

HSI=Heaviness of smoking index \ge 4, calculated based on sum of time to first cigarette (0: 61+min, 1:31-60 min, 2: 6-30 min, 3: \le 5 min) and number of cigarettes smoked per day (0: 0-10 cigarettes per day [CPD]

Also: age, duration of smoking, alcohol problem (hazardous, harmful, non-drinker/light drinker), maybe HIV-status, (adding this may reduce your sample size too much, because of missing HIV-status, unless you include HIVstatus unknown as a category of HIV-status). Depression

⁸ Scoring the audit

Scores for each question range from 0 to 4, with the first response for each question (eg never) scoring 0, the second (eg less than monthly) scoring 1, the third (eg monthly) scoring 2, the fourth (eg weekly) scoring 3, and the last response (eg. daily or almost daily) scoring 4. For questions 9 and 10, which only have three responses, the scoring is 0, 2 and 4 (from left to right).

A score of 8 or more is associated with harmful or hazardous drinking, a score of 13 or more in women, and 15 or more in men, is likely to indicate alcohol dependence.

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is associated with harmful or hazardous drinking, a score of 13 or more in women, and 15 or more in men, is likely to indicate alcohol dependence. Eligibility criteria for our study is an AUDIT score ≥ 8 for men or ≥ 7 for women but <20. However, if assumptions of linear regression are not met we will either transform the data or use alternative regression analyses such as ordinal logistic regression.

Adherence to TB and ART medication will be measured using an adherence index based on a modified version of the AIDS Clinical Trials Group (ACTG) Adherence Questionnaire; where patients with at least 95% of adherence will be considered as having optimal adherence otherwise will be considered as having low (or suboptimal) adherence. We will use logistic regression to model patient's characteristics (age, sex, alcohol, smoking status, depression) that might influence adherence at 6-month; we will also compare adherence between study arms. Similar to the previous outcomes, we will account for any potential clustering by centre.

We will also report the proportion of HIV-positive participants on ART at six months and compare these to the baseline using standardised questions on the CRF.

8.4.1 Secondary outcome definitions that involve defining a positive baseline test:

To operationalise the above we need to define how we determine if someone has a positive baseline smear or culture. Each participant might have up to two tests of the following: smear test, GeneExpert test, and culture test.

If a test was administered two times then to be considered negative both tests should be negative, otherwise it is considered positive. If a test was administered only once, then the result of that instance is taken as is.

¹Saunders JB, Aasland OG, Babor TF et al. Development of the alcohol use disorders identification test (AUDIT): WHO collaborative project on early detection of persons with harmful alcohol consumption — II. Addiction 1993, 88: 791–803

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If for a patient more than one test was administered, then to be considered negative the result should be negative under both tests.

This is illustrated in the following table.

Baseline	P=+, N = -		
		XXXX Test at run 2 (Result 2)	
XXXX Test run 1 (Result 1)		N	Р
	N	NN	NP
	Р	PN	PP
XXXX=Smear or Culture or GeneXpert			
Smear test at baseline	N1	NN	
	P1	NP, PN, PP	
Genexpert	N2	NN	
	P2	NP, PN, PP	
culture	N3	NN	
	P3	NP, PN, PP	
Positive at baseline	PB	P1 or P2 or P3	
Negative at baseline	NB	N1 and N2 and N3	

To define a conversion among those who were positive at baseline, we used the primary outcome response category cured to indicate a negative result at month 6 per the nurse's assessment.

Among the 403 participants who were positive at baseline 168(41.69%) were recorded as cured by 6-month of these 83 (39.9%) in the control arm compared to 85 (43.59%) see Table 4. The odds ratio of conversion is 1.16 (95% CI: (0.83,1.63)) comparing the intervention arm to the control arm. When adjusting for district, sex, and smoking/drinking status and HIV status at baseline, the OR reduces to 1.07 (95% CI: (0.76,1.51))

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8.4.2 6-months Continuous Abstinence

For this outcome, those who smoked cigarettes at baseline were considered as the analytic sample. In addition, those where the following could not be ascertained: self-report of not smoking > 5 cigarettes six months from the start of the abstinence period and supported by a negative biochemical test CO < 7 ppm were considered as smokers for the analysis of this variable. The number of participants who identified as cigarette smokers at baseline were 345 (60.1%)⁹.

23 (85.19%) out of 27 with three measurements available ¹⁰ managed to abstain continuously for six months. These were similarly distributed across the two study arms, see Table 4. Among those who identified as cigarette smokers, 10 (5.59%) participants in the intervention arm continuously abstained. In the control arm there were 12 (7.23%) who continuously abstained for 6-months.

The crude odds of 6-months continuous abstinence is 0.76 (95% CI: (0.35,1.63)) in the intervention arm compared to the control arm among baseline cigarette smokers. Given the limited number of those who were identified as continually abstained, we were only able to adjust for one additional variable at a time. Adding one of the following variables: heaviness of smoking, type of drinker at baseline, age when started smoking, and the duration of smoking at baseline, the adjusted odds ratio of continuous abstinence comparing the intervention to the control arm did not differ much from the crude estimate of 0.76. The adjusted estimate for the various models ranged between 0.73 and 0.76 with similar confidence limits as for the crude estimate. Furthermore, we did not have evidence that any of the adjusting variables were statistically significantly correlated to continuous abstinence in these models.

We carried an additional analysis where those who died and were smokers at baseline, 22 in total, 20 were cigarette smokers and were removed from the analytical sample for the continuous abstinence outcome. This resulted in a crude

⁹ Any type of tobacco smoking at baseline 372 (64.8%). The numbers reflect those who used manufactured cigarettes which were the vast majority.

¹⁰ We had only 27 participants who had self-report of not smoking at 3 months, 6 months and a carbon monoxide reading at 6 months. Continuous abstinence was defined as a self-report of not smoking > 5 cigarettes six months from the start of the abstinence period and supported by a negative biochemical test CO < 7 ppm

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OR of 0.78 which is similar to the OR when the larger analytic sample was considered; this was also the case for the associated 95% CI. Similar estimates were observed when adjusting for the aforementioned variables.

We carried an additional analysis where the analytic sample was all those who were smokers regardless of the type used. There were a total of 372 that identified as smokers. The crude odds ratio in this case changes to 0.86 (95% CI: (0.38,1.95)). The 95% CI is similar to that of the smaller analytic sample. When adjusting for the heaviness of smoking, type of drinker at baseline, age when started smoking, and the duration of smoking at baseline, the adjusted odds ratio of continuous abstinence comparing the intervention to the control arm did not differ much from the crude estimate.

3-months Continuous Abstinence

Among those who identified as cigarette smokers, 20 (11.17%) participants in the intervention arm continuously abstained for 3-months while in the control arm there were 27 (16.27%) who continuously abstained for 3-months.

The crude odds of 3-months continuous abstinence is 0.65 (95% CI: (0.37,1.14)) in the intervention arm compared to the control arm among baseline cigarette smokers. Given the limited number of those who were identified as continually abstained, we were only able to adjust for one additional variable at a time. Adding one of the following variables: heaviness of smoking, type of drinker at baseline, age when started smoking, and the duration of smoking at baseline, the adjusted odds ratio of continuous abstinence comparing the intervention to the control arm did not differ much from the crude estimate of 0.65. The adjusted estimate for the various models ranged between 0.63 and 0.66 with similar confidence limits as for the crude estimate.

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Point Abstinence

There were 57 participants who indicated that they stopped smoking tobacco completely at the 3-month follow-up, of which one had more than 5 cigarettes in the past 3 months. There were 81 participants who indicated that they stopped smoking tobacco completely at the 6-month follow-up, of which three had more than 5 cigarettes in the past 3 months. Only 30 participants had information for the entirety of the 6-month period, of which none consumed more than 5 cigarettes over the past 6-month period. Of these, 23 had a confirmed CO < 7 ppm, 4 had these levels >= 7, and 3 were missing.

182 responded that they continued smoking either as usual or at a reduced rate but regularly at month 3; whereas 133 done so at month 6 of these we had 97 that had measurements at both time points thus resulting in a total of 218 where they have smoked on a regular basis over the past 6 month period. 11 had responded as not smoking in the past three months at month 6 but had missing information for the first three months; 9 of these had carbon monoxide readings available at month 6. Of these nine, two had their CO >= 7 (in fact these were 10 & 10.1).

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8.4.4 Change in harmful or hazardous drinking at month 3 and 6 follow-ups

Alcohol use was measured using the AUDIT questionnaire. We will assess whether there has been a reduction in alcohol consumption three months and six months following recruitment. This analysis will be performed on the group of participants who were harmful or hazardous drinkers at baseline. Figure YYY provides histograms of the AUDIT score at baseline, 3-months and 6-months, respectively, among those who were considered as hazardous or harmful drinkers. It also presents scatterplots of the 3-months and 6-months scores versus the baseline scores for this group by trial arm, respectively.

Among those who were harmful or hazardous drinkers at baseline those in the intervention arm had on average a reduction of 0.04 points (95% CI: (-2,1.91)) on the AUDIT score at 6-months compared to those in the control arm controlling for their baseline score, see Table 6. However, when additionally adjusting for district, sex, and smoking/drinking status (which effectively flags smokers/non-smokers who are also drinkers) and HIV status at baseline; the intervention arm had an average increase of 0.02 points on the AUDIT score (95% CI: (-1.55,1.6)) compared to the control arm. It is worth noting that of the variables in the adjusted model; the only statistically significant result is for the district variable. It seems that those in district "S" score on average 5.8 points less than those in "B" (95% CI: (-11.26,-0.35)); similarly those in "L" score 5 points less than those in "B" on AUDIT but we do not have evidence that this difference is statistically significant (95% CI: (-10.35,0.26)), see Table 6 for further details.

At 3-month, the estimates were an average increase of 0.55 (95% CI: (-1.01,2.11)) on the AUDIT score in the intervention arm compared to the control arm when only accounting for the baseline scores; whereas it increased to 0.74 (95% CI: (0.62,2.1)) when adjusting for other covariates in the model, for further details see Table 7.

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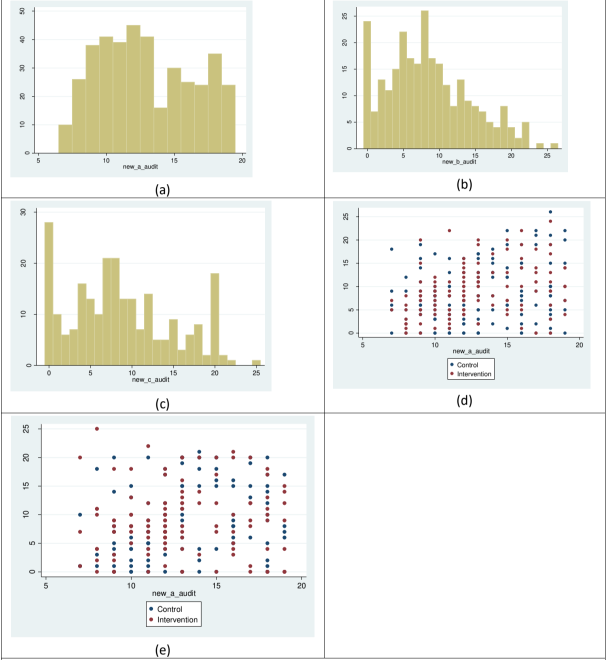


Figure YYY: Graphs (a), (b), and (c) are histograms of the AUDIT score at baseline, 3-months and 6-months, respectively, among those who were considered as hazardous or harmful drinkers. (d) and (e) are scatterplots of the 3-months and 6-months scores versus the baseline scores for this group, respectively.

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8.4.5 Proportion of ART uptake of HIV-positive participants at month 3 and 6 follow-ups

We will assess whether there has been an increase in proportion of HIV-positive participants on ART at three and six months from baseline using standardised questions on the CRF.

There were 171 HIV-positive participants whose baseline ART medication status was known and whose ART medication status was known as well at 6-months. Of these, 123 remained on their medication, 19 took up medication at 6-months compared to not taking medication at baseline, whereas 29 stopped taking their medication at 6-months but were on medication at baseline, see Table 6 for further details.

There were 10 who had an unknown status in terms of medication at baseline and no information was available about them at 6-months, 12 who were initially of unknown medication status who took up medication at 6-months (these were equally distributed between the two arms). Furthermore, there were 10 who were not taking medication at baseline and 102 who were taking medication at baseline whose 6-months medication status was not recorded.

At 6-months, the odds ratio of taking medication at 6-months was 2.05 (95% CI: (0.80,5.27)) in the intervention arm compared to the control arm, controlling for ART baseline medication status.

There were 188 HIV-positive participants whose baseline ART medication status was known and whose ART medication status was known as well at 3-months. Of these, 122 remained on their medication, 16 took up medication at 3-months compared to not taking medication at baseline, whereas 50 stopped taking their medication at 3-months but were on medication at baseline, see Table 7 for further details.

There were 9 who had an unknown status in terms of medication at baseline and no information was available about them at 3-months, 11 who were initially of unknown

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medication status who took up medication at 3-months, and two who were initially of unknown medication status who were not taking medication at 3-months. Furthermore, there were 9 who were not taking medication at baseline and 82 who were taking medication at baseline whose 3-months medication status was not recorded. Furthermore, there were 4 who carried on not taking medication at 3-months.

At 3-months, the odds ratio of taking medication at 6-months was 0.79 (95% CI: (0.38,1.65)) in the intervention arm compared to the control arm, controlling for ART baseline medication status.

Medicine Adherence:

At 3-months 165 (98.8%) of 167 participants had optimal ART medication adherence, whereas 139(97.2%) of 143 had optimal ART medication adherence. These were similar across the two arms.

Similarly, at 3-months 319(91.67%) of 348 participants had optimal TB medication adherence, whereas 120(90.23%) of 133 had optimal TB medication adherence. These were similar across the two arms.

8.5 Subgroup analyses (See above analyses)

We will conduct subgroup analyses to determine whether TB treatment outcomes differ between subgroups, as follows: HIV-positive versus HIV-negative participants; participants with an alcohol problem only versus smokers only versus participants who are conjoint smokers and drinkers; and participants who were GeneXpert positive versus participants who were GeneXpert negative at baseline.

8.5 Sensitivity analyses

In case of missing data, multiple imputations and appropriate sensitivity analyses will be conducted. As it is likely that more than one variable will have missing data we will use multiple imputations using chained equations (**MICE**). A minimum of 10 imputations will be performed; however, the final number of imputations will depend on the missing in the data. We will report the decisions that we make with regard to the number of imputations and the

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variables we use in the imputations. We will also conduct a sensitivity analysis to explore the implications of the missing at random assumption [21p,22p].

8.6 Adverse events

Analysis of adverse events and serious AE will explore whether these differ by treatment arm using Chi-square tests.

8.7 Planned interim review and analyses

No interim analysis is planned. The main analyses will be completed after three months of the data closing.

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8.8 List of Tables and Graphs

Measures of central tendency and percentages will be reported to two decimal places whereas measures of variability and p-values will be reported to three decimal places.

The following is a list of suggested tables and graphs; the templates are included in Appendix A.

- TABLE 1: Numbers in the study at Baseline and follow-ups at month 3, and 6 by centre and study arm.
- TABLE 2: Descriptive statistics for socio-demographic and socio-economic characteristics at baseline and as analysed by study arm. Frequencies and (percentages) are presented unless otherwise stated
- TABLE 3: Descriptive statistics for smoking history, alcohol history, clinical characteristics and depression score at baseline and as analysed by study arm. Frequencies and (percentages) are presented unless otherwise stated.
- TABLE 4: Descriptive statistics for primary and secondary outcomes by study arm at baseline (where available), 3 month (where available) and 6 month. Frequencies and (percentages) are presented unless otherwise stated.
- TABLE 5: Number and type of adverse events at month 2, 3, and 6 by centre and study arm.
- TABLE 6: Regression analysis results for the primary and secondary outcomes at 6 month. Estimates presented with corresponding 95% CI. Crude and adjusted estimates are provided.
- TABLE 7: Regression analysis results for the secondary outcomes that are measured at 3 month. Estimates presented with corresponding 95% CI. Crude and adjusted estimates are provided.

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9. SAP amendment log

Amendment/addition to SAP and reason for change	New version number, name and date
SAP completed and signed-off	V1.0,
Updated verion	V1.1, May 2020

10. Signatures of approval

Sign-off of the final approved version of the Statistical Analysis Plan by the principle investigator and trial statistician(s) (can also include Trial Manager/Co-ordinator)

<u>Name</u>	<u>Trial Role</u>	<u>Signature</u>	<u>Date</u>

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11. References

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12. Appendix A (Results tables)

Table 1: Numbers in the study at Baseline and follow-ups at month 3 and 6 by centre and study arm. (SEE FLOWCHART)

Centre Baseline			Month 3		Month 6	
	Intervention	Control	Intervention	Control	Intervention	Control
Total						

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TABLE 2: Descriptive statistics for socio-demographic, socio-economic, and clinical characteristics at baseline by study arm. Frequencies and (percentages) are presented unless otherwise stated

TABLE 2: DESCRIPTIVE STATISTICS FOR SOCIO-DEMOGRAPHIC, SOCIO-ECONOMIC, AND CLINICAL CHARACTERISTICS AT BASELINE AND AS ANALYSED BY STUDY ARM. FREQUENCIES AND (PERCENTAGES) ARE PRESENTED UNLESS OTHERWISE STATED Baseline N1/N2 Control Intervention Total Arm 1 Arm 2 291/283 38.56 (11.15) Age in years: mean (SD) 39.37 (12.60) Age in years: median (IQR) Gender 69 (23.7) 60 (21.2) 129 (22.5) Female 445 (77.5) Male 222 (76.3) 223 (78.8) Do not want to disclose Marital status Married or living together 102 (35.1) 95 (33.6) 197 (34.3) Divorced/separated 20 (6.9) 18 (6.4) 38 (6.6) Widowed 10 (3.4) 7 (2.5) 17 (3.0) 150 (53.0) Never married and never lived together 144 (49.5) 294 (51.2) Declined to answer 15 (5.2) 13 (4.6) 28 (4.9) **Education** 7 (2.4) 5 (1.8) 12 (2.1) No education 23 (7.9) 20 (7.1) 43 (7.5) Grades 1-5 Grades 6-7 32 (11.0) 35 (12.4) 67 (11.7)

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Grades 8-11	96 (33.0)	128 (45.2)	224 (39.0)
Grade 12	87 (29.9)	70 (24.7)	157 (27.4)
Higher	24 (8.2)	8 (2.8)	32 (5.6)
Declined to answer	22 (7.6)	17 (6.0)	39 (6.8)
Employment			
Self-employed (full-time)	30 (10.3)	36 (12.7)	66 (11.5)
Employed full-time (30 hrs a week or more)	62 (21.3)	54 (19.1)	116 (20.2)
Employed part-time (less than 30 hrs a week)	19 (6.5)	29 (10.2)	48 (8.4)
Retired	17 (5.8)	16 (5.7)	33 (5.7)
Unemployed (but able to work)	125 (43.0)	120 (42.4)	245 (42.7)
Unable to work because of long-term disability or ill health	9 (3.1)	8 (2.8)	17 (3.0)
Full-time student	12 (4.1)	4 (1.4)	16 (2.8)
Caring from my home and family/doing household work/housewife	0 (0.0)	2 (0.7)	2 (0.3)
Occasional work ("piece job")	17 (5.8)	12 (4.2)	29 (5.1)
Declined to answer	0 (0.0)	2 (0.7)	2 (0.3)
Ever worked or spent time in mines			
No	244 (83.8)	237 (83.7)	481 (83.8)
Yes	46 (15.8)	45 (15.9)	91 (15.9)
Socioeconomic status			

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Household items (Yes is displayed)			
A radio	249 (85.6)	234 (82.7)	483 (84.1)
A television	255 (87.6)	255 (90.1)	510 (88.9)
A landline telephone	21 (7.2)	22 (7.8)	43 (7.5)
A desktop or laptop computer	67 (23.0)	48 (17.0)	115 (20.0)
A refrigerator	248 (85.2)	240 (84.8)	488 (85.0)
A vacuum cleaner or floor pol	49 (16.8)	35 (12.4)	84 (14.6)
A microwave oven	198 (68.0)	189 (66.8)	387 (67.4)
An electric or gas stove	254 (87.3)	238 (84.1)	492 (85.7)
A washing machine	153 (52.6)	136 (48.1)	289 (50.3)
Total number of assets: mean (SD)	5.14 (1.96)	4.94 (1.77)	5.04 (1.87)
In the past month, number of days you or people in			
the household went to bed hungry because there			
was no food to eat			
0 days	244 (83.8)	238 (84.1)	482 (84.0)
1-7 days	45 (15.5)	34 (12.0)	79 (13.8)
More than 7 days	2 (0.7)	9 (3.2)	11 (1.9)
Declined to answer	0 (0.0)	2 (0.7)	2 (0.3)
TB and medical history			
TB history			
Patient category			
New patient	264 (92.3)	249 (90.2)	513 (91.3)
Relapse	10 (3.5)	9 (3.3)	19 (3.4)
Re-treatment after default	9 (3.1)	14 (5.1)	23 (4.1)
Re-treatment after failure	1 (0.3)	2 (0.7)	3 (0.5)
Other	2 (0.7)	2 (0.7)	4 (0.7)
Site of disease			
Pulmonary and Extra Pulmonary (ICD-10 A17-A19)	3 (1.1)	3 (1.1)	6 (1.1)

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Pulmonary only (ICD-10 A15)		281 (98.9)	272 (98.9)	553 (98.9)
Number of pre-treatment smear results				
One		197 (92.1)	169 (87.6)	366 (89.9)
Two		17 (7.9)	24 (12.4)	41 (10.1)
Smear result	N =	196	180	376
At least one positive smear result		111 (56.63)	109 (60.56)	220 (58.51)
Number of Gene XPert results recorded				
One		225 (97.8)	210 (97.2)	435 (97.5)
Two		5 (2.2)	6 (2.8)	11 (2.5)
Gene XPert result	N =	211	204	415
At least one positive Gene XPert result		184 (87.2)	178 (87.25)	362 (87.23)
Number of culture results recorded on the TB Treatment record				
One		54 (94.7)	42 (97.7)	96 (96.0)
Two		3 (5.3)	1 (2.3)	4 (4.0)
Culture result	N =	41	32	73
At least one positive culture result		20 (48.78)	15 (46.88)	35 (47.95)
Co-morbidities				
Hypertension		19 (6.93)	11 (4.1)	30 (5.54)
Diabetes		5 (1.84)	4 (1.49)	9 (1.66)
Epilepsy		3 (1.09)	4 (1.49)	7 (1.29)
Mental illness		3 (1.09)	0 (0)	3 (0.55)
Liver disease		1 (0.36)	1 (0.38)	2 (0.37)
Renal insufficiency		1 (0.36)	1 (0.38)	2 (0.37)
Allergies		2 (0.76)	0 (0)	2 (0.38)

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Other	1 (0.36)	1 (0.38)	2 (0.37)
Total Number of comorbidities			
0	265 (96.0)	260 (96.7)	525 (96.3)
1	10 (3.6)	8 (3.0)	18 (3.3)
2	0 (0.0)	1 (0.4)	1 (0.2)
5	1 (0.4)	0 (0.0)	1 (0.2)
HIV status			
Negative	118 (40.7)	125 (44.2)	243 (42.4)
Positive	163 (56.2)	142 (50.2)	305 (53.2)
Unknown	9 (3.1)	16 (5.7)	25 (4.4)
HIV positive patients			
CD4 Count: mean(SD)			
Using Cotrimoxazole	104 (63.8)	100(67.11)	204 (65.38)
Using anti-retroviral	139 (85.28)	118 (79.19)	257 (82.37)

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TABLE 3: Descriptive statistics for smoking history, alcohol history and depression score at baseline by study arm. Frequencies and (percentages) are presented unless otherwise stated.

		Baseline	
	Control	Intervention	Total
	Arm 1	Arm 2	Total
n the past month, smoked tobacco			
Not at all	110 (37.8)	92 (32.5)	202 (35.2)
Daily	149 (51.2)	149 (52.7)	298 (51.9)
Less than Daily	32 (11.0)	42 (14.8)	74 (12.9)
lad a drink in the past 12-months	208 (71.5)	223 (78.8)	431 (75.1)
UDIT Score (males): mean (SD) [max :19] [min = 8 if drinkers only]	12.27 (3.98)	13.02 (3.78)	12.66 (3.89)
UDIT Score (females): mean (SD) [max :19] [min = 7 if drinkers only]	11.32 (4.02)	10.98 (4.02)	11.15 (4)
Orinking and Smoking Combined (Constructed)			
Drinkers Only	110 (37.8)	92 (32.5)	202 (35.2)
Smokers Only	83 (28.5)	60 (21.2)	143 (24.9)
Smokers and Drinkers	98 (33.7)	131 (46.3)	229 (39.9)

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	Arm 1	Arm 2	Total
Smoking History (current smokers only)	181	191	372
On the days that you smoke, how soon after you wake up do you have your first cigarette?			
After 60 minutes	30 (16.6)	28 (14.7)	58 (15.6)
31-60 minutes	24 (13.3)	16 (8.4)	40 (10.8)
6- 30 minutes	60 (33.1)	65 (34.0)	125 (33.6)
Within 5 minutes	67 (37.0)	82 (42.9)	149 (40.1)
Duration of smoking in months : mean (SD)	212.09 (134.03)	224.93 (127.82)	218.68 (130.86)
Duration of smoking in months: median (IQR)	186 (110, 282)	206 (135, 294)	200.5 (123, 287)
Age started smoking in years : mean (SD)	19.2 (6.3)	19.3 (6.3)	19.3 (6.3)
Age started smoking in years : median (IQR)	18 (15-20)	18 (16-21)	18 (15.5-20.5)
Form of tobacco used			
Manufactured cigarettes (Yes)	166 (91.7)	179 (93.7)	345 (92.7)
Number of days in the past 7days you smoked: mean (SD)	5.3 (2.65)	5.53 (2.33)	5.42 (2.49)
Average number of cigarettes smoked daily: mean (SD)	6.18 (6.43)	6.48 (8.21)	6.34 (7.39)
Hand-rolled cigarettes (Yes)	14 (7.7)	21 (11.0)	35 (9.4)
Number of days in the past 7days you smoked: mean (SD)	4.71 (2.84)	3.71 (2.95)	4.11 (2.91)
Average number of handrolled cigarettes smoked daily: mean (SD)	3.71 (3.97)	3.81 (2.82)	3.77 (3.27)
Pipe (Yes)	4 (2.2)	2 (1.0)	6 (1.6)
Number of days in the past 7days you smoked: median (IQR)	1 (0, 2.5)	4.5 (2, 7)	2 (0, 3)
Average number of daily sessions: median (IQR)	1 (0, 3.5)	3 (1, 5)	1.5 (0, 5)
Length of one session (on average) in minutes: median (IQR)	90 (60, 107.5)	60 (30, 90)	90 (30, 90)
Cigars, cheroots or cigarillos (Yes)	1 (0.55)	0 (0)	1(0.27)

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		5 (1.3)
10 (5.5)	4 (2.1)	14 (3.8)
134(74.03)	158(82.72)	292(78.49)
96 (53.0)	108 (56.5)	204 (54.8)
61 (33.7)	58 (30.4)	119 (32.0)
24 (13.3)	25 (13.1)	49 (13.2)
52 (28.7)	64 (33.5)	116 (31.2)
129 (71.3)	127 (66.5)	256 (68.8)
2.46 (2.98)	2.64 (1.62)	2.56 (2.32)
36.65 (94.62)	25.13 (37.91)	30.29 (69.22)
6.15 (13.48)	4.22 (8.9)	5.09 (11.18)
6 (3.3)	5 (2.6)	11 (3.0)
10 (5.5)	12 (6.3)	22 (5.9)
104 (57 5)	109 (57.1)	213 (57.3)
61 (33.7)	65 (34.0)	126 (33.9)
Control	Intervention	
t	134(74.03) 96 (53.0) 1	134(74.03) 158(82.72) 1 96 (53.0) 108 (56.5) 1 61 (33.7) 58 (30.4) 2 4 (13.3) 25 (13.1) 2 5 129 (71.3) 127 (66.5) 2 2 46 (2.98) 2.64 (1.62) 3 6.65 (94.62) 25.13 (37.91) 6 .15 (13.48) 4.22 (8.9) 1 0 (5.5) 12 (6.3) 1 104 (57.5) 109 (57.1) 6 1 (33.7) 65 (34.0)

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	Arm 1	Arm 2	Total
Likelihood that I WILL QUIT smoking completely and permanently in the next three months			
definitely will not	6 (3.3)	5 (2.6)	11 (3.0)
probably will not	11 (6.1)	13 (6.8)	24 (6.5)
probably will	103 (56.9)	104 (54.5)	207 (55.6)
definitely will	61 (33.7)	69 (36.1)	130 (34.9)
Ever used any methods to help you stop smoking tobacco in the past 3 months? (Yes)	23 (12.7)	16 (8.4)	39 (10.5)
Out of your pocket spend (in Rands) on methods to help you stop smoking in the past 3 months: mean (SD)	134.87 (237.58)	139.19 (155.07)	136.64 (205.36)
Average spend per week on cigarettes over the past 3 months: mean (SD)	168.77 (178.15)	180.29 (185.27)	174.58 (181.61)
Smokeless tobacco use (all participants)			
In the past month, have you used smokeless tobacco (Snuff) on a daily basis			
Not at all	275 (94.5)	275 (97.2)	550 (95.8)
Daily	12 (4.1)	6 (2.1)	18 (3.1)
Less than Daily	4 (1.4)	2 (0.7)	6 (1.0)
Duration of using ST in months: mean(SD)	113.69 (112.7)	152 (91.16)	126.46 (105.61)
Age started using ST in years : mean (SD)	27.56 (10.57)	25.88 (13.43)	27 (11.33)
Form of ST used (for SLT users)			
Snuff (by mouth)	2 (12.5)	0 (0.0)	2 (8.3)
Snuff (by nose)	11 (68.8)	5 (62.5)	16 (66.7)
Chewing tobacco leaves	0(0)	0(0)	0(0)
Other	1 (6.3)	0 (0.0)	1 (4.2)
	Arm 1	Arm 2	Total

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	Control	Intervention	
Tobacco smoke exposure: number of days in the past 30 days, you were in a place where someone smoked close to you (all participants?)			
Home			
7 days or less	233 (80.1)	229 (80.9)	462 (80.5)
More than 7 days	58 (19.9)	54 (19.1)	112 (19.5)
Workplace (Missing 315; probably Not Applicable)			
7 days or less	84 (65.6)	89 (67.9)	173 (66.8)
More than 7 days	11 (21 1)	42 (32.1)	86 (33.2)
Cafes/restaurants			
7 days or less	230 (79.0)	234 (82.7)	464 (80.8)
More than 7 days	61 (21.0)	49 (17.3)	110 (19.2)
Shebeens, bars or clubs			
7 days or less	219 (75.3)	213 (75.3)	432 (75.3)
More than 7 days	72 (24 7)	70 (24.7)	142 (24.7)
Bus/train/taxi/ vehicle			
7 days or less	229 (78.7)	219 (77.4)	448 (78.0)
More than 7 days	62 (21.3)	64 (22.6)	126 (22.0)
Shops/shopping mall			
7 days or less	235 (80.8)	226 (79.9)	461 (80.3)
More than 7 days	56 (19.2)	57 (20.1)	113 (19.7)
·			
	Arm 1	Arm 2	Total

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Supplemental material

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Had a drink in the past 12-months	208 (71.5)	223 (78.8)	431 (75.1)
Help to stop drinking (drinkers only)			
Ever used any methods to stop drinking alcohol in the past 3 months	21 (11.2)	22 (10.7)	43 (10.9)
Average spend in Rands per week on alcohol over the past 3 months: mean (SD)	363.76 (531.56)	337.03 (387.94)	349.79 (461.66)
Depression: CESD 10 how often you felt or behaved this way during the past week			
I was bothered by things that usually don't bother me.			
Rarely or None of the Time (Less than 1 day)	180 (61.9)	159 (56.2)	339 (59.1)
Some or a Little of the Time (1-2 days)	71 (24.4)	78 (27.6)	149 (26.0)
Occasionally or a Moderate Amount of the Time (3-4 days)	24 (8.2)	31 (11.0)	55 (9.6)
Most or All of the Time (5-7 days)	16 (5.5)	15 (5.3)	31 (5.4)
I had trouble keeping my mind on what I was doing.			
Rarely or None of the Time (Less than 1 day)	150 (51.5)	136 (48.1)	286 (49.8)
Some or a Little of the Time (1-2 days)	93 (32.0)	85 (30.0)	178 (31.0)
Occasionally or a Moderate Amount of the Time (3-4 days)	31 (10.7)	38 (13.4)	69 (12.0)
Most or All of the Time (5-7 days)	17 (5.8)	24 (8.5)	41 (7.1)
I felt depressed.			
Rarely or None of the Time (Less than 1 day)	152 (52.2)	171 (60.4)	323 (56.3)
Some or a Little of the Time (1-2 days)	94 (32.3)	70 (24.7)	164 (28.6)
Occasionally or a Moderate Amount of the Time (3-4 days)	37 (12.7)	30 (10.6)	67 (11.7)
Most or All of the Time (5-7 days)	8 (2.7)	12 (4.2)	20 (3.5)
I felt that everything I did was an effort.			

Rarely or None of the Time (Less than 1 day)	140 (48.1)	117 (41.3)	257 (44.8)
Some or a Little of the Time (1-2 days)	77 (26.5)	87 (30.7)	164 (28.6)
Occasionally or a Moderate Amount of the Time (3-4 days)	38 (13.1)	43 (15.2)	81 (14.1)
Most or All of the Time (5-7 days)	36 (12.4)	36 (12.7)	72 (12.5)
I felt hopeful about the future.			
Rarely or None of the Time (Less than 1 day)	85 (29.2)	84 (29.7)	169 (29.4)
Some or a Little of the Time (1-2 days)	78 (26.8)	70 (24.7)	148 (25.8)
Occasionally or a Moderate Amount of the Time (3-4 days)	53 (18.2)	56 (19.8)	109 (19.0)
Most or All of the Time (5-7 days)	75 (25.8)	73 (25.8)	148 (25.8)
I felt fearful.			
Rarely or None of the Time (Less than 1 day)	175 (60.1)	159 (56.2)	334 (58.2)
Some or a Little of the Time (1-2 days)	82 (28.2)	79 (27.9)	161 (28.0)
Occasionally or a Moderate Amount of the Time (3-4 days)	25 (8.6)	32 (11.3)	57 (9.9)
Most or All of the Time (5-7 days)	9 (3.1)	13 (4.6)	22 (3.8)
My sleep was restless.			
Rarely or None of the Time (Less than 1 day)	142 (48.8)	135 (47.7)	277 (48.3)
Some or a Little of the Time (1-2 days)	95 (32.6)	89 (31.4)	184 (32.1)
Occasionally or a Moderate Amount of the Time (3-4 days)	31 (10.7)	33 (11.7)	64 (11.1)
Most or All of the Time (5-7 days)	23 (7.9)	26 (9.2)	49 (8.5)
I was happy.			
Rarely or None of the Time (Less than 1 day)	85 (29.2)	74 (26.1)	159 (27.7)
Some or a Little of the Time (1-2 days)	70 (24.1)	65 (23.0)	135 (23.5)
Occasionally or a Moderate Amount of the Time (3-4 days)	73 (25.1)	86 (30.4)	159 (27.7)
Most or All of the Time (5-7 days)	63 (21.6)	58 (20.5)	121 (21.1)
I felt lonely.			

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Rarely or None of the Time (Less than 1 day)	158 (54.3)	161 (56.9)	319 (55.6)
Some or a Little of the Time (1-2 days)	89 (30.6)	83 (29.3)	172 (30.0)
Occasionally or a Moderate Amount of the Time (3-4 days)	23 (7.9)	26 (9.2)	49 (8.5)
Most or All of the Time (5-7 days)	21 (7.2)	13 (4.6)	34 (5.9)
I could not get "going".			
Rarely or None of the Time (Less than 1 day)	168 (57.7)	174 (61.5)	342 (59.6)
Some or a Little of the Time (1-2 days)	78 (26.8)	69 (24.4)	147 (25.6)
Occasionally or a Moderate Amount of the Time (3-4 days)	35 (12.0)	29 (10.2)	64 (11.1)
Most or All of the Time (5-7 days)	10 (3.4)	11 (3.9)	21 (3.7)
Total CESD 10: mean (SD)	8.44 (4.38)	8.74 (4.8)	8.59 (4.59)

Table 4: Descriptive statistics for primary and secondary outcomes by study arm at baseline (where available), 3 month (where available) and 6 month. Frequencies and (percentages) are presented unless otherwise stated

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TABLE 4: DESCRIPTIVE STATISTICS FOR PRIMARY AND SECONDARY OUTCOMES BY STUDY ARM AT BASELINE (WHERE AVAILABLE), 3 MONTH (WHERE AVAILABLE) AND 6 MONTH. FREQUENCIES AND (PERCENTAGES) ARE PRESENTED UNLESS OTHERWISE STATED.

	Baseline		Fol	Follow-up 3-month			Follow-up 6-month		
	Control	Intervention	Total	Control	Interventio n	Total	Control	Intervention	Total
	Arm1	Arm2	Total	Arm1	Arm2	Total	Arm1	Arm2	Total
TB treatment status detailed									
Cured							108 (37.11)	105 (37.1)	213 (37.11)
Treatment completed							96 (33.0)	87 (30.74)	183 (31.88)
Treatment default							15 (5.2)	29 (10.25)	44 (7.67)
Treatment failure							5 (1.7)	2 (0.7)	7 (1.2)
Acquired drug resistance							1 (0.34)	4 (1.41)	5 (0.87)
Died							11 (3.78)	15 (5.30)	26 (4.53)
Transfer out							8 (2.75)	11 (3.89)	19 (3.31)
Unknown							42 (14.43)	25 (8.83)	67 (11.67)
Missing							5 (1.72)	5 (1.77)	10 (1.74)
TB treatment status binary (Primary outcome**)									
Not Successful							87 (29.9)	91 (32.16)	178 (31.01)
Successful							204 (70.10)	192 (67.84)	396 (68.99)
At least one positive smear result	85 (53.46)	96 (61.15)	181 (57.28)						
At least one positive Gene XPert result	184 (87.20)	178 (87.25)	362 (87.23)						
At least one positive culture result	11 (57.89)	10 (66.67)	21 (61.76)						

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Sputum smear, Gene XPert, or culture result									
Negative	29 (12.45)	32 (14.16)	61 (13.29)						
Positive	204 (87.55)	194 (85.84)	398 (86.71)						
Conversion from positive to negative***									
Yes							83(39.9)	85(43.59)	168(41.69)
No							125(60.1)	110(56.41)	235(58.31)
Continuous smoking abstinence									
among cigarette smokers at baseline									
Yes				27(16.27)	20(11.17)	47(13.62)	12(7.23)	10(5.59)	22(6.38)
No				139(83.73)	159(88.83)	298(86.38)	154(92.77)	169(94.41)	323(93.62)
Harmful or hazardous drinking ¹									
Had a drink in the past 12-months	208 (71.5)	223 (78.8)	431 (75.1)						
AUDIT score: mean (SD)	12.03 (4)	12.53 (3.93)	12.29 (3.96)						
Harmful or hazardous drinkers out of those who had a drink in the past 12-months at baseline (%)	188(90.38)	206(92.38)	394(91.42)	141	130	271	112	127	239
AUDIT Score (males): mean (SD) [max :19; min = 8]*!	13.14 (3.31)	13.61 (3.29)	13.39 (3.31)	8.20(6.08)	9.08(4.97)	8.63(5.58)	9.21(6.58)	8.24(5.41)	8.69(5.99)
AUDIT Score (females): mean (SD) [max:19; min = 7]*!	11.73 (3.52)	11.55 (3.6)	11.64 (3.54)	8.5(6.52)	8.15(6.44)	8.33(6.44)	7.67(6.84)	9.97(6.79)	8.89(6.86)
AUDIT score: mean (SD)	12.76 (3.42)	13.12 (3.47)	12.94 (3.45)	8.28(6.18)	8.84(5.38)	8.55(5.81)	8.79(6.66)	8.70(5.83)	8.74(6.22)
Difference from baseline				-4.61 (6.26)	-4.07(5.33)	-4.35(5.83)	-4.25(6.56)	-4.17(6.61)	-4.21(6.57)

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Taking ART medication among HIV	[163 (56.2)]	[142 (50.2)]	[305 (53.2)]	[122]	[83]	[205]	[100]	[83]	[183]
Positive patients									
Yes	139 (85.28)	115 (80.99)	254 (83.28)	91 (74.6)	58 (69.9)	149 (72.7)	80 (80.0)	74 (89.2)	154 (84.2)
ART medication adherence									
Optimal adherence				101(99.02)	64(98.46)	165(98.8)	75(97.4)	64(96.97)	139(97.2)
Suboptimal adherence				1(0.98)	1(1.54)	2(1.2)	2(2.6)	2(3.03)	4(2.8)
TB medication adherence									
Optimal adherence				181(92.35)	138(90.79)	319(91.67)	61(89.71)	59(90.77)	120(90.23)
Suboptimal adherence				15(7.65)	14(9.21)	29(8.33)	7(10.29)	6(9.23)	13(9.77)
1h									

¹ hazardous/harmful drinkers who are not alcohol dependent= AUDIT score ≥ 8 for men or ≥ 7 for women but < 20

^{**}Primary Outcome: in the published protocol paper "This is a binary variable defined as either successful treatment (cured or treatment completed) or failed treatment, death, acquired drug resistance, loss to follow-up or 'default', or not outcome evaluated.

^{***} Conversion from positive to negative: this was based on having a cured treatment outcome among those who were positive at baseline.

^{*!} Important distinction at baseline for eligibility purposes.

TABLE 5: Number and type of adverse events at month 2, 3, and 6 by centre and study arm (see other document)

TABLE 5: Number and type of adverse events at month 2, 3, and 6 by centre and study arm								
Centre		Month 2		Month 3		Month 6		
	TYPE	Intervention	Control	Intervention	Control	Intervention	Control	

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TABLE 6: Regression analysis results for the primary and secondary outcomes at 6 months. Estimates presented with corresponding 95% CI. Crude and adjusted estimates are provided.

TABLE 6: Regression analysis results for the primary and secondary outcomes at 6 months. Estimates presented with corresponding 95% CI. Crude and adjusted estimates are provided.

	Crude Odds Ratio (95% CI)*	P Value*	Adjusted Odds Ratio (95% CI)*	P Value*	
Primary outcome					
TB treatment status: Successful (Ref: Not successful)	0.9 (0.64,1.27)	0.548	0.86 (0.60,1.24)	0.421	
Secondary outcomes					
Sputum smear or culture result: converted from positive to negative (Ref: Not converted)	1.16 (0.83,1.63)	0.374	1.07‡ (0.76,1.51)	0.684	
Six-month continuous smoking abstinence among cigarette smokers at baseline (Ref: No)	0.76 (0.35,1.64)	0.482			
Taking ART medication among HIV positive patients!	2.05 (0.80,5.27)	0.136			
TB medication adherence (Reference: Optimal)	0.89 (0.26,3.07)	0.849			
ART medication adherence (Reference: Optimal)	1.17 (0.14,9.94)	0.884			

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	Study arm regression coefficient		Study arm regression coefficient		
AUDIT for those who were harmful or hazardous drinkers at baseline**:	-0.04 (-2,1.91)	0.966	0.02 (-1.55,1.6)	0.976	

^{*} analyses accounted for potential clustering by centre.

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[†] Number of participants whose outcome was treatment successful among the total number in the group.

 $[\]ddagger$ Adjusted for district, sex, and smoking/drinking status and HIV status at baseline

^{**}Controlling for the AUDIT baseline values.

[!] Controlling for the AUDIT baseline values and adjusted for district, sex, and smoking/drinking status and HIV status at baseline

^{!!} Adjusting for art status at baseline

TABLE 7: Regression analysis results for secondary outcomes measured at 3-months. Estimates presented with corresponding 95% CI. Crude and adjusted estimates are provided.

TABLE 7: Regression analysis results for secondary outcomes measured at 3-months. Estimates presented with corresponding 95% CI. Crude and adjusted estimates are provided.

	Crude Odds Ratio (95% CI)	P Value	Adjusted Odds Ratio (95% CI)	P Value	
Secondary outcome					
3-months continuous smoking abstinence among cigarette smokers at baseline	0.65 (0.37,1.14)	0.135			
Taking ART medication among HIV positive patients!!	0.79 (0.38,1.65)	0.53	0.74‡ (0.35,1.58)	0.443	
TB medication adherence (Reference: Optimal)	1.22 (0.52,2.87)	0.641			
ART medication adherence (Reference: Optimal)	1.58 (0.10,26.12)	0.750			
	Study arm regression coefficient		Adjusted estimates		
AUDIT for those who were harmful or hazardous drinkers at baseline**:	0.55 (-1.01,2.11)	0.474	0.74 (-0.62,2.1)	0.273	

^{*} analyses accounted for clustering.

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[‡] Adjusted for district, sex, and smoking/drinking status and HIV status at baseline

^{**}Controlling for the AUDIT baseline values.

[!] Controlling for the AUDIT baseline values and adjusted for district, sex, and smoking/drinking status and HIV status at baseline

^{!!} Adjusting for art status at baseline

Additional tables SMS-fidelity

Variable	Intervention (N=) (=no. of participants allocated to	Control (N=) (=no. participants allocated to
	intervention group, for example 245)	control group, for example 248)
No. of participants who received ALL due IMB messages INDEPENDENT OF WHETHER they completed MI 1 (i.o.w this is ITT analysis)	For example 120/245 (49 %)	For example 2/248 (%)
Com	pletion of first MI and initiation o	f SMS-sequence
Completed first MI	n/N (%) (=No. who completed first MI in intervention arm/ no. participants allocated to intervention arm [%] For example 170/245 [69%])	n/N (%) (=No. who completed first MI in control/ no. participants allocated to control [%] For example 2/248)
No. of participants who received ALL due messages after receipt of MI1 and SMS sequence was generated (this is a type of Per Protocol analysis:	For example 120/170 (71%)	2/2

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denominator limited						
to those who had						
MI1)						
SMS delivery for participants for whom the SMS-sequence was initiated (after receipt of first MI)						
	Mean (SD)	Median (IQR)	Mean (SD)	Median (IQR)		
		Range				
Average no.	For example		Expected to be	Expected to be		
adherence	8 (3.4)	9 (7-10)	0 or close to it	0 or close to it		
messages received						
per participant		Range 0-10				
(n=) $(n = the no. of$						
participants who completed first MI,						
in this example 170)						
Average no.	For example		Expected to be	Expected to be		
tobacco-related	7 (3.4)	7 (6-7)	0 or close to it	0 or close to it		
messages received						
(n=) (n=no of		Range 0-7				
participants who						
completed first MI						
AND were current						
tobacco users at baseline, for						
example 90)						
Average no. alcohol	For example		Expected to be	Expected to be		
related messages	7 (3.4)	7 (6-7)	0 or close to it	0 or close to it		
received						
(n=) (n=no.		Range 0-7				
participants who						
completed first MI						

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AND were drinkers at baseline=for example 100)				
Average no. IMB	For example		Expected to be	Expected to be
messages received	16 (4.5)	15 (10-20)	0 or close to it	0 or close to it
(n= total no who				
completed first MI)		Range 0-24		

Other variables not yet analysed:

CESD at 3 and 6-month FU

Other smoking related questions at 3 and 6 month (quit intentions etc) but also SLT use at 3 and 6 month. (important because participants may have switched from tobacco smoking to SLT)
Other non-HE questions not yet analysed

13. Additional descriptive Statistics

TABLE 8: Additional Descriptive statistics for characteristics at baseline by study arm. Frequencies and (percentages) are presented unless otherwise stated

		Baseline		
	N1/N2	Intervention	Control	Total
		Arm 1	Arm 2	
TYPE OF MINE WORK	46/45			91
Coal		7 (15.2)	3 (6.7)	10 (11.0)
Diamond		3 (6.5)	1 (2.2)	4 (4.4)
Gold		13 (28.3)	17 (37.8)	30 (33.0)
Platinum and palladium		24 (52.2)	17 (37.8)	41 (45.1)
Chromium		11 (23.9)	12 (26.7)	23 (25.3)
Uranium		1 (2.2)	1 (2.2)	2 (2.2)
Manganese		0(0)	0(0)	0(0)
Other		0(0)	0(0)	0(0)
Total Types of Mines worked				
in				
1	73			
2	12			
3	3			
4	1			

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Exact Distribution for number smoked in the past seven days			
0	24 (14.5)	16 (8.9)	40 (11.6)
1	4 (2.4)	4 (2.2)	8 (2.3)
2	4 (2.4)	3 (1.7)	7 (2.0)
3	5 (3.0)	13 (7.3)	18 (5.2)
4	13 (7.8)	8 (4.5)	21 (6.1)
5	5 (3.0)	17 (9.5)	22 (6.4)
6	1 (0.6)	3 (1.7)	4 (1.2)
7	110 (66.3)	115 (64.2)	225 (65.2)
new_a_positive_TB			
0	29 (11.98)	33 (14.35)	62 (13.14)
1	122 (50.41)	104 (45.22)	226 (47.88)
2	80 (33.06)	81 (35.22)	161 (34.11)
3	11 (4.55)	12 (5.22)	23 (4.87)

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