Global TB data collection form 2022

SECTION 0: (ONGOING, NOT PART OF THE ANNUAL FORM)

Section 0 questions are only for countries in the high TB burden list and other regional priority countries (dc tbcases monthly display = 1)

Ongoing reporting of the latest provisional numbers of new and relapse TB cases

Please provide the provisional number of people with new and relapse TB cases (all forms, including people with drug-resistant TB) that were notified each month or quarter as soon as the month or quarter has ended. WHO will use this ongoing reporting to monitor the impact of the disruptions caused by the COVID-19 pandemic on essential TB services and on TB incidence and mortality.

Note that these data are published instantly on the WHO TB data web page at https://www.who.int/teams/global-tuberculosis-programme/data

At what frequency can you report?	70 Monthly 71 Quarterly	report_frequency
Report coverage Please explain in the remarks below if these preliminary data do not include all reporting units in your country	7 111 011110	report_coverage

(if frequency = monthly):

2022

m.1	January	m_01
m.2	February	m_02
m.3	March	m_03
m.4	April	m_04
m.5	May	m_05
m.6	June	m_06
m.7	July	m_07
m.8	August	m_08
m.9	September	m_09

m.10	October	m_10
m.11	November	m_11
m.12	December	m_12
2023		
2020		
m.1	January	
m.2	February	
(if frequency	v = quarterly):	
2022		
q.1	January - March	q_1
q.2	April - June	q_2
q.3	July - September	q_3
q.4	October - December	q_4
Please	e tick the box if data are not av	vailable for empt
Remarks		

view_TME_master_data_collection

SECTION 1: IDENTIFICATION

National TB control programme manager (NTP) or equivalent

1.1	Name (as you would like it to appear in the acknowledgement section Report) Please do not use honorifics or titles such as "Dr", "Professor".	n of the WHO Global TB
	Given (first) name Family name Preferred order Given name - family name Family name - given name	ntp_name
1.2	Functional title	ntp_title
1.3	Telephone (including country and city codes)	ntp_phone
1.4	E-mail	ntp_email
	rson responsible for entering data on the WHO globers (if different from the NTP manager)	bal TB data collection
1.5	Name (as you would like it to appear in the acknowledgement section Report) Please enter only one name here. If you want us to acknowledge more names please remarks" section. Note: People with accounts to use the WHO global TB data collection system will also Global TB Report	enter them in the "General
	Given (first) name Family name Preferred order Given name - family name Family name - given name	rep_name
1.6	E-mail	rep_email

General remarks

remarks_general

Note that remarks made under individual sections are all combined into another field called <code>remarks_sections</code>

SECTION 2: DIAGNOSIS AND TREATMENT

view_TME_master_notification

TB notifications by history, site, diagnostic method and by age and sex, 2021 calendar year

Please report **all** people diagnosed with TB and eligible for TB treatment (including those diagnosed with drug-resistant TB), regardless of whether treatment was started or not. People who died or were lost before treatment start should be notified as they are important to include for surveillance purposes and, from a public health perspective, may have contacts that require tracing and follow up.

		Previous anti-TB	treatment status
		(i) New, or previous treatment history unknown	(ii) Relapse
2.1	Pulmonary TB cases, bacteriologically confirmed (positive by WHO-recommended rapid diagnostics such as Xpert MTB/RIF, Ultra, Truenat MTB, MTB Plus, TB-LAMP or LF-LAM; culture positive; smear positive)	new_labconf	ret_rel_labconf
2.2	Pulmonary TB cases, clinically diagnosed (Not bacteriologically confirmed as positive for TB, but diagnosed with active TB by a clinician or another medical practitioner who has decided to give the patient a full course of TB treatment)	new_clindx	ret_rel_clindx
2.3	Extrapulmonary TB cases, bacteriologically confirmed or clinically diagnosed Cases with both pulmonary and extrapulmonary TB are classified as pulmonary TB cases	new_ep	ret_rel_ep
	Total	c_newunk	

	Total new and relapse	c_newinc
2.4	Previously treated patients, <u>excluding relapse cases</u> (pulmonary or extrapulmonary, bacteriologically confirmed or clinically diagnosed) ('treatment after failure', 'treatment after lost to follow-up' and cases whose outcome after their most recent course of treatment is unknown or undocumented)	ret_nrel
	Total cases notified	c_notified

		Previous anti-TB treatment status			
		(i) New	(ii) Previously treated (including relapses)	(iii) Previous treatment history unknown	
2.5	Among the bacteriologically confirmed pulmonary TB cases reported in question 2.1 and question 2.4,	<pre>pulm_labconf_new</pre>	pulm_labconf_ret	pulm_labconf_unk	

numbers by previous anti- TB treatment status
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2.6 Among the cases reported in questions 2.1 – 2.4, total number of TB cases reported among foreign-born individuals (or among non-citizens if that is the criterion used in your country)

notif foreign

New and relapse TB cases by age and sex, 2021 calendar year

Time-changes in the distribution of cases by age and sex are analyzed by WHO to understand trends in disease burden and gaps in the performance of TB surveillance

2.7 For which age groups can you provide notifications disaggregated by age group and sex?

220 0-4, 5-9, 10-14, 15-19, 20-24, 25-34, 35-44, 45-54, 55-64, 65+ (if you have a national electronic case-based database (i.e. holding separate records for each TB case) for all TB patients)

221 0-4, 5-14, 15-24, 25-34, 35-44, 45-54, 55-64, 65+

222 0-14, 15-24, 25-34, 35-44, 45-54, 55-64, 65+

[223] (and last option only for the 2 countries which were not able to give full age/sex breakdowns for all new and relapse cases last year in 2021: Gambia, Mozambique)

0-4, 5-14, 15+

2.8 Are all relapse cases included in table 2.9 below?

2.9 New and relapse TB cases (pulmonary or extrapulmonary, bacteriologically confirmed or clinically diagnosed, drug-susceptible or drug-resistant)

(The table shown below has different column depending on the answer to 2.7 above

		Age					
	0–4	5–9	10–14	15–19	20-24	25-34	
Male	newrel_m04	newrel_m59	newrel_m1014	newrel_m1519	newrel_m2024	newrel_m2534	
Female	newrel_f04	newrel_f59	newrel_f1014	newrel_f1519	newrel_f2024	newrel_f2534	

	Age						
	35–44	45–54	55–64	>65	Unknown		
Male	newrel_m3544	newrel_m4554	newrel_m5564	newrel_m65	newrel_mu		
Female	newrel_f3544	newrel_f4554	newrel_f5564	newrel_f65	newrel_fu		
Total	(auto calc.)						

2022: WHO TB data collection form 2022-04-14

Use of WHO-recommended rapid diagnostic tests

WHO-recommended rapid diagnostic tests employ molecular techniques or biomarker-based techniques to detect TB. These are currently Xpert MTB/RIF (including Ultra), Truenat MTD and MTB Plus, TB-LAMP and lateral flow urine lipoarabinomannan assay (LF-LAM).

2.10	using a WHO-recommended	
	O No	rdx_data_available
	Yes, available from our routine surveillance system Yes, estimated from a review of a random sample of medical reconstitution, representative of the national TB patient population Not applicable (because there were no TB cases)	ords or treatment cards of TB
2.11	(if yes from routine surveillance) Number of new and relapse cases reported in questions 2.1 – 2.3 tested using a WHO-recommended rapid diagnostic (for example Xpert MTB/RIF) as the initial diagnostic test (regardless of test result) ^b	newinc_rdx
2.12	(if yes from survey) Number of new and relapse cases reported in questions 2.1 – 2.3 whose medical records or treatment cards were included in the survey ^b	rdxsurvey_newinc_
2.13	(if yes from survey) Among the cases reported in 2.12, the number tested using a WHO-recommended rapid diagnostic (such as Xpert MTB/RIF) as the initial diagnostic test (regardless of test result)	rdxsurvey_newinc_rdx
^b Pulmo	onary or extrapulmonary, bacteriologically confirmed or clinically diagnosed	
☐ Ple Rema	ease tick the box if data are not available for empty cells above. rks:	

2022: WHO TB data collection form 2022-04-14

Drug-resistant TB, 2021 calendar year

Calculating indicators related to drug-resistant TB detection and enrolment on treatment requires data on notified TB cases recorded in the basic management unit (BMU) TB register, not from laboratory registers

Diagnosis and enrolment on treatment of TB patients with laboratoryconfirmed rifampicin resistance and no evidence of fluoroguinolone resistance

2.14	Among all people diagnosed with pulmonary or extrapulmonary TB in 2021 (reported in questions 2.1 – 2.4), number with laboratory-confirmed rifampicin resistance and no evidence of fluoroquinolone resistance (susceptible or tests not done) This should not include pre-XDR-TB and XDR-TB patients.	conf_rr_nfqr
2.15	Number of patients with laboratory-confirmed rifampicin resistance and	conf_rr_nfqr_tx
	no evidence of fluoroquinolone resistance (susceptible or tests not done) started on MDR-TB treatment in 2021 Pulmonary or extrapulmonary. Also include patients diagnosed before 2021 but started on treatment in 2021. This should not include pre-XDR-TB and XDR-TB patients treated for pre-XDR-TB or XDR-TB.	
2.16	Number of patients without laboratory confirmation of rifampicin resistance started on MDR-TB treatment in 2021 Pulmonary or extrapulmonary. Also include patients diagnosed before 2021 but started on treatment in 2021. This should not include pre-XDR-TB and XDR-TB patients treated for pre-XDR-TB or XDR-TB.	unconf_rr_nfqr_tx
2.17	Total number of patients who started treatment for MDR-TB in 2021	(auto calc.)
2.18	Among patients in 2.17 who started treatment for MDR-TB in 2021, the number who were children aged 0-14 years Pulmonary or extrapulmonary. Also include patients diagnosed before 2021 but started on treatment in 2021. This should not include pre-XDR-TB and XDR-TB patients treated for pre-XDR-TB or XDR-TB.	rrmdr_014_tx

Diagnosis and enrolment on treatment of TB patients with laboratory-confirmed rifampicin resistance and also fluoroquinolone resistance (pre-XDR-TB or XDR-TB)

TB resistant to rifampicin and to any fluoroquinolone is now called pre-XDR-TB. Pre-XDR-TB that is also resistant to at least one of bedaquiline or linezolid is now called XDR-TB.

These new definitions were published by WHO in January 2022. See https://www.who.int/publications/i/item/meeting-report-of-the-who-expert-consultation-on-the-definition-of-extensively-drug-resistant-tuberculosis.

2.19	Number with laboratory-confirmed rifampicin resistance and also fluoroquinolone resistance (i.e. pre-XDR-TB or XDR-TB) identified in 2021 Pulmonary or extrapulmonary. Also include patients diagnosed with rifampicin resistance before 2021 and then with fluoroquinolone resistance in 2021.	conf_rr_fqr
2.20	Number with laboratory-confirmed rifampicin resistance and also fluoroquinolone resistance (i.e. pre-XDR-TB or XDR-TB) who started treatment for pre-XDR-TB or XDR-TB in 2021 Pulmonary or extrapulmonary. Also include patients diagnosed before 2021 but started on treatment in 2021.	conf_rr_fqr_tx

Treatment regimens

2.21	Had any TB patients been started on Bedaquiline for the treatment of MDR-TB, pre-XDR-TB or XDR-TB by the end of 2021?	mdrxdr_bdq_used
	1 Yes 0 No 3 Don't know	
2.22	(If yes to 2.21) Number of patients started on Bedaquiline in 2021	mdrxdr_bdq_tx
2.23	Had any patients been started on all oral longer MDR-TB treatment regimens by the end of 2021? Longer MDR-TB regimens are those used for the treatment of MDR/RR-TB. These last 18 months or more and may be standardized or individualized. These regimens are usually designed to include a minimum number of second-line TB medicines considered to be effective based on patient history or drug-resistance patterns. 1 Yes 0 No 3 Don't know	mdrxdr_alloral_used
2.24	(If yes to 2.23) Number of patients started on all oral longer MDR-TB treatment regimens in 2021	mdrxdr_alloral_tx
2.25	Had any patients been started on all oral shorter MDR-TB treatment regimens (duration of 12 months or less) by the end of 2021? 1 Yes 0 No 3 Don't know	mdr_alloral_short_used
2.26	(If yes to 2.25) Number of patients started on all oral shorter MDR-TB treatment regimens (duration of 12 months or less) in 2021.	mdr_alloral_short_tx
2.27	Number of patients who started treatment for MDR-TB, pre-XDR-TB or XDR-TB in 2021 who are also being actively monitored for adverse events This refers to the active and systematic clinical and laboratory assessment of patients on treatment with new anti-TB drugs, novel MDR-TB regimens or XDR-TB regimens to detect, manage and report suspected or confirmed drug toxicities. See.the WHO aDSM Implementation Framework	mdr_tx_adsm
☐ PI	ease tick the box if data are not available for empty cells above. urks:	

2022: WHO TB data collection form 2022-04-14

Anti-tuberculosis drug resistance: Surveillance

Diagnostic testing for drug resistance in bacteriologically confirmed pulmonary TB patients, 2021 calendar year

Note: Questions below are for reporting **all** bacteriologically confirmed pulmonary drug resistant cases notified in the country in 2021. To report the results of a drug resistance survey (i.e., a study using a specially-designed sample of patients that is representative of the national or a subnational TB population), please go to the drug resistance survey section.

Results of first-line drug testing

Data reported below should only include results from specimens taken at the start of a treatment course or within 2 weeks of starting treatment. For patients changing treatment course after failure, data should only include results from specimens taken before the start of the subsequent treatment course or within the first 2 weeks of starting the subsequent treatment course.

Rifampicin testing:

		Previous anti-TB treatment status		
		(i) New	(ii) Previously treated (including relapses) ^a	Total ^b
	Bacteriologically confirmed pulmonary TB patients reported in 2.5	pulm_labconf_new	pulm_labconf_ret	
2.28	Among bacteriologically confirmed pulmonary TB patients reported in 2.5, number of patients with test results for rifampicin	r_rlt_new	r_rlt_ret	(auto calc.)
2.29	Among patients with test results for rifampicin reported in 2.28, number of patients with resistance to rifampicin (RR-TB)	rr_new	rr_ret	(auto calc.)

^aPrevious anti-TB treatment: > 1 month of treatment with combined anti-TB drugs excluding preventive chemotherapy.

The following two questions will only be shown to countries in the high MDR-TB burden list

Rifampicin testing among relapse cases only:

	Bacteriologically confirmed pulmonary relapse TB patients reported in 2.1(ii)	ret_rel_labconf
2.28b	Among bacteriologically confirmed pulmonary relapse TB patients reported in 2.1(ii), number of patients with test results for rifampicin	r_rlt_rel
2.29b	Among relapse patients with test results for rifampicin reported in 2.28b, number of patients with resistance to rifampicin (RR-TB)	rr_rel

^bExcluding cases with unknown treatment history

Rifampicin and isoniazid testing among <u>new</u> patients in 2.28(i):

		(i) Resistant to isoniazid	(ii) Susceptible to isoniazid
2.30	Resistant to rifampicin	a	b
2.31	Susceptible to rifampicin	С	d
2.32	Total new patients tested for both rifampicin and isoniazid	(auto calc.)	

Rifampicin and isoniazid testing among *previously treated (including relapses)* patients in 2.28(ii):

		(i) Resistant to isoniazid	(ii) Susceptible to isoniazid
2.33	Resistant to rifampicin	е	f
2.34	Susceptible to rifampicin	g	h
2.35	Total previously treated (including relapse) patients tested for both rifampicin and isoniazid	(auto calc.)	

Note that 2.30 - 2.35 are equivalent to the following table used in previous years:

	Previous anti-T	B treatment status	
	New	Previously treated (including relapses)	Total
(a) Among patients with test results for rifampicin reported in 2.28, number of patients with test results for isoniazid	dst_rlt_new	dst_rlt_ret	(auto calc.)
(b) Among patients reported in (a) with test results for rifampicin and isoniazid, number of patients with resistance to isoniazid (regardless of result for rifampicin)	dst_rlt_hr_new	dst_rlt_hr_ret	(auto calc.)
(c) Among patients reported in (a) with test results for rifampicin and isoniazid, number of patients with resistance to rifampicin (regardless of result for isoniazid)	dst_rlt_rr_new	dst_rlt_rr_ret	(auto calc.)
(d) Among patients reported in (a) with test results for rifampicin and isoniazid, number of patients with resistance to both rifampicin and isoniazid (MDR-TB)	mdr_new	mdr_ret	(auto calc.)

```
dst_rlt_new = a + b + c + d

dst_rlt_hr_new = a + c

dst_rlt_rr_new = a + b

mdr_new = a

dst_rlt_ret = e + f + g + h

dst_rlt_hr_ret = e + g

dst_rlt_rr_ret = e + f

mdr_ret = e
```

Results of second-line drug testing

TB resistant to rifampicin and to any fluoroquinolone is now called pre-XDR-TB. Pre-XDR-TB that is also resistant to at least one of levofloxacin, moxifloxacin, bedaquiline or linezolid is now called XDR-TB.

These new definitions were published by WHO in January 2022. See https://www.who.int/publications/i/item/meeting-report-of-the-who-expert-consultation-on-the-definition-of-extensively-drug-resistant-tuberculosis.

Fluoroquinolone testing:

2.36	Among RR-TB patients reported in 2.29, number of patients with test results for any fluoroquinolone	rr_dst_rlt_fq
2.37	Among patients with test results for fluoroquinolones reported in 2.36, number of patients with resistance to any fluoroquinolone (pre-XDR-TB)	rr_fqr

Bedaquiline and linezolid testing among pre-XDR-TB patients in 2.37

		(i) Resistant to bedaquiline (XDR-TB)	(ii) Susceptible to bedaquiline	(iii) Unknown resistance to bedaquiline
2.38	Resistant to linezolid (XDR-TB)	rr_fqr_bdqr_lzdr	rr_fqr_bdqs_lzdr	rr_fqr_bdqu_lzdr
2.39	Susceptible to linezolid	rr_fqr_bdqr_lzds	rr_fqr_bdqs_lzds	rr_fqr_bdqu_lzds
2.40	Unknown resistance to linezolid	rr_fqr_bdqr_lzdu	rr_fqr_bdqs_lzdu	rr_fqr_bdqu_lzdu
2.41	Total number with XDR-TB (2.38 + 2.39(i) + 2.40(i))	(auto calc.)		
2.42	Total pre-XDR-TB cases tested in 2.38 + 2.39 + 2.40	(auto calc.)		

Notes: to determine testing coverage and to calculate prevalence of XDR among pre-XDR patients use the inner top left 2x2 square with results for BDQ and LZD

Testing coverage numerator = 2.38(i) + 2.38(ii) + 2.39(i) + 2.39(ii) Testing coverage denominator = 2.37

Prevalence of XDR among pre-XDR numerator = 2.38(i) + 2.38(ii) + 2.39(i)Prevalence of XDR among pre-XDR denominator = 2.38(i) + 2.38(ii) + 2.39(i) + 2.39(ii)

The total of all cells in the table (2.42) should be equal to 2.37

Please tick the box if data are not available for empty cells above
Remarks:

TB/HIV, 2020 calendar year

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To update TB/HIV data for 2020 and earlier years, please go to the update page

Number of new and relapse TB patients notified in 2020 tested for HIV at the time of TB diagnosis or with known HIV status at the time of TB diagnosis.	newrel_hivtest
Among new and relapse TB patients reported in 2020, the number recorded as HIV positive	newrel_hivpos
Among HIV-positive new and relapse TB patients reported in 2020, the number started or continued on antiretroviral therapy (ART)	newrel_art

TB/HIV, 2021 calendar year

	definitions and reporting framework? 1 Yes 0 No ("No" means all TB cases have been included in 2.44 – 2.46 according to the pre-2013 re	eporting framework)
	Total number of notified new and relapse cases in 2.1 - 2.3	c_newinc
2.44	Number of new and relapse TB patients notified in 2021 tested for HIV at the time of TB diagnosis or with known HIV status at the time of TB diagnosis. This should include TB patients who were known to be HIV-positive (e.g. documented evidence of enrolment in HIV care such as enrolment in the pre-ART register or in the ART register once started on ART) or with documented negative HIV test conducted at the time of TB diagnosis. If the patient's HIV status is subsequently determined, he or she should be reclassified accordingly.	newrel_hivtest
2.45	Among new and relapse TB patients reported in 2.44, the number recorded as HIV positive	newrel_hivpos
2.46	Among HIV-positive new and relapse TB patients reported in 2.45, the number who started or continued on antiretroviral therapy (ART) (Number of HIV-positive new and relapse TB patients started on TB treatment during the reporting period who were already on antiretroviral therapy or started on antiretroviral therapy during TB treatment)	newrel_art

The next questions will only be shown to countries in the high TB/HIV burden list, and all countries in AMR

2.43b	Do you have a case-based surveillance system that and relapse TB cases in children aged 0-14 years?	allows you to report on TB	HIV indicators for new
	1 Yes 0 No	tbhiv_014_flg	

(if yes from	2.43b):)	
2.44b	Number of new and relapse TB patients notified in 2021 aged 0-14 years tested for HIV at the time of TB diagnosis or with known HIV status at the time of TB diagnosis. This should include TB patients who were known to be HIV-positive (e.g. documented evidence of enrolment in HIV care such as enrolment in the pre-ART register or in the ART register once started on ART) or with documented negative HIV test conducted at the time of TB diagnosis. If the patient's HIV status is subsequently determined, he or she should be reclassified accordingly.	newrel_hivtest_014
2.45b	Among new and relapse TB patients aged 0-14 years reported in 2.44b, the number recorded as HIV positive	newrel_hivpos_014
2.46b	Among HIV-positive new and relapse TB patients aged 0-14 years reported in 2.45b the number who started or continued on antiretroviral therapy (ART)	newrel_art_014

	Please tick	the box if	data a	are not	available	for e	mpty	cells	above
Re	emarks:								

Global AIDS Monitoring 2021 data imported from UNAIDS (https://aidsreportingtool.unaids.org/), as supplied by National AIDS Programme respondents:

		2021
GAM.7.7 Numerator	Number of HIV-positive new and relapse TB patients started on TB treatment during the reporting period who were already on antiretroviral therapy or started on antiretroviral therapy during TB treatment within the reporting year	hiv_tbtx_art

		2021
GAM. 7.8 Numerator	Total number of people living with HIV newly enrolled in HIV treatment who have active TB disease during the reporting period	hiv_tbdetect
GAM. 7.8 Denominator	Total number of people newly enrolled in HIV treatment (i.e., those who registered for antiretroviral therapy during the reporting period)	hiv_reg_new2

		2021
GAM 7.9 Numerator	Total number of people living with HIV newly enrolled on antiretroviral therapy who also started tuberculosis preventive treatment during the same reporting period	hiv_new_tpt
GAM 7.9 Denominator	Total number of people living with HIV newly enrolled on antiretroviral therapy (i.e., those registered for antiretroviral therapy during the reporting period) This denominator should be the same as the denominator of indicator 7.8	hiv_new

GAM 7.9 Numerator (alternative)	Total number of people living with HIV currently enrolled on antiretroviral therapy who started tuberculosis preventive treatment during the reporting period	hiv_all_tpt
GAM 7.9 Denominator (alternative)		hiv_all

		2020
GAM. 7.10 Numerator		hiv_all_tpt_c ompleted
GAM. 7.10 Denominator	1 1 1 1 2	hiv_all_tpt_s tarted

Note that the following additional variables concerning data imported from UNAIDS appear only in

dcf.latest notification

and are not transferred to the master view:

Date extracted from UNAIDS reporting system:

hiv_unaids_date_exported

GAM.7.7 notes:

Indicator reported as not relevant or not available.

Reason data not imported

Reported remarks:

hiv_tbrx_art_NA
hiv_tbrx_art_NI_reason
hiv_tbrx_art_remarks

GAM.7.8 notes:

Indicator reported as not relevant or not available.

hiv tbdetect hiv reg new2 NA

Reason data not imported

hiv_tbdetect_hiv_reg_new2_NI_reason

Reported remarks:

hiv tbdetect hiv reg new2 remarks

GAM.7.9 notes:

Indicator reported as not relevant or not available.

hiv tpt eligible start NA

Reason data not imported

hiv_tpt_eligible_start_NI_reason

Reported remarks:

hiv_tpt_eligible_start_remarks

GAM.7.10 notes:

Indicator reported as not relevant or not available.

Reason data not imported

Reported remarks:

hiv_tpt_completed_NA

hiv_tpt_completed_reason

hiv_tpt_completed_remarks

Treatment outcomes for TB patients registered in 2020 calendar year for drug-susceptible TB treatment

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WHO is now collecting data using the revised treatment outcome definitions applicable to patients treated using any regimen regardless of drug-resistance status, type of drugs used or duration of treatment. The definitions were published in April 2021 at https://www.who.int/publications/i/item/9789240022195

Treatment failed: A patient whose treatment regimen needed to be terminated or permanently changed to a new regimen or treatment strategy. Reasons for the change include:

- evidence of additional drug resistance to medicines in the regimen;
- adverse drug reactions; or
- no clinical response and/or no bacteriological response.

Cured: A pulmonary TB patient with bacteriologically confirmed TB at the beginning of treatment who completed treatment as recommended by the national policy, with evidence of bacteriological response and no evidence of failure.

Treatment completed: A patient who completed treatment as recommended by the national policy, whose outcome does not meet the definition for cure or treatment failure.

Died: A patient who died for any reason before starting treatment or during the course of treatment.

Lost to follow-up: A patient who did not start treatment or whose treatment was interrupted for 2 consecutive months or more.

Not evaluated: A patient for whom no treatment outcome was assigned. This includes cases "transferred out" to another treatment unit and those whose treatment outcome is unknown; however, it excludes those lost to follow-up

2.47a Are outcome categories in questions 2.48 to 2.54 for both drug-susceptible and drug-resistant TB in line with the revised definitions published by WHO in April 2021 at https://www.who.int/publications/i/item/9789240022195?

1	Yes
0	No

used 2021 defs flg

2.47b	Are outcomes of relapse cases incl	uded in row 2.48 below	
	1 Yes	rel with new flg	
	0 No	roi_wikii_now_iig	
	("No" means relapse cases have	e been included in row 2.49	according to the pre-2013 reporting framework

		Number of			Treatment outcor	ne	
		patients registered in 2020	Cured or treatment completed	Treatment failed	Died	Lost to follow-up	Not evaluated
2.48	Patients treated for drug- susceptible TB who were registered as new, relapse or previous treatment history unknown patients (bacteriologically confirmed or clinically diagnosed, pulmonary or extrapulmonary)	newrel_coh	newrel_succ	newrel_fail	newrel_died	newrel_lost	c_newrel_neval
2.49	Patients treated for drug- susceptible TB who were registered as 'treatment after failure', 'treatment after lost to follow up' patients or patients whose outcome after their most recent course of treatment is unknown or undocumented (bacteriologically confirmed or clinically diagnosed, pulmonary or extrapulmonary)	ret_nrel_coh	ret_nrel_succ	ret_nrel_fail	ret_nrel_died	ret_nrel_lost	c_ret_nrel_neval
2.50	Among the patients in 2.48, all HIV-positive TB patients treated for drug-susceptible TB (bacteriologically confirmed or clinically diagnosed, pulmonary or extrapulmonary)	tbhiv_coh	tbhiv_succ	tbhiv_fail	tbhiv_died	tbhiv_lost	c_tbhiv_neval

Please tick the box if data are not available for empty cells above. Remarks:

Treatment outcomes for children aged 0-14 years registered in 2020 calendar year for drug-susceptible TB treatment

2.51	Do you have a case-based surveillance system that allows you to report on treatment outcomes for children aged 0-14 years?							
	1	Yes						
	0	No	outcomes_014_flg					

(if yes from 2.51)

				ries. Only cohort size and cured/completed shown for other countries Treatment outcome					
		Number of patients registered in 2020	Cured or treatment completed	Treatment failed	Died	Lost to follow-up	Not evaluated		
2.52	Children aged 0- 14 years treated for drug- susceptible TB who were registered as new, relapse and previous treatment history unknown patients		newrel_014_succ	newrel_014_fail	newrel_014_died	newrel_014_lost	c_newrel_014_neval		

These questions are only for countries in any one of the 3 high TB burden lists

Treatment outcomes for females registered in 2020 calendar year for drug-susceptible TB treatment

2.51b		ase-based surveillance system that allows you to report on treatment outcomes disaggregated by sex?				
	1 Yes 0 No	outcomes_sex_flg				

(if yes from 2.51b)

		Number of patients	Treatment outcome					
	Number of patients registered in 2020		Cured or treatment completed	Treatment failed	Died	Lost to follow-up	Not evaluated	
4	Females (all ages) treated for drug- susceptible TB who were registered as new, relapse and previous treatment history unknown patients		newrel_f_succ	newrel_f_fail	newrel_f_died	newrel_f_lost	c_newrel_f_neval	

Please tick the box if data are not available for empty cells above. Remarks:

		Number of patients		Treatment outcome					
		started on drug- resistant TB treatment in 2019	Cured or treatment completed	Treatment failed	Died	Lost to follow-up	Not evaluated		
2.53	Patients with laboratory-confirmed RR-TB/MDR-TB treated for MDR-TB and not treated for pre- XDR or XDR-TB	mdr_coh	mdr_succ	mdr_fail	mdr_died	mdr_def	c_mdr_neval		
2.54	Patients with laboratory-confirmed pre-XDR-TB/XDR-TB treated for pre-XDR or XDR-TB Do not include these cases in 2.53. If you cannot report outcomes for XDR-TB cases separately from RR-/MDR-TB cases, include all cases in 2.53 and add a note in the remarks below.	xdr_coh	xdr_succ	xdr_fail	xdr_died	xdr_def	c_xdr_neval		

Please tick the box if data	a are not available	for empty	cells	above
Remarks:				

SECTION 3: SURVEYS AND SERVICES

Recent drug resistance survey or patient cost survey (2020 and 2021 calendar years)

Variables 3.1a – 3.1c do not appear in any view; countries that answer yes are listed in

Note: The question below is only about drug resistance <u>surveys</u>. Survey results do not need to be reported to WHO more than once.

3.1a Was a drug resistance survey (i.e., a study using a specially-designed sample of patients that is representative of the national or a subnational TB patient population) completed in 2020 or 2021?

1 Yes	drs cmplt
0 No	ars_empre

If you have responded 'yes', you will be contacted by WHO to provide information about the methods and results of the survey

3.1b Was a patient costs survey completed in 2020 or 2021?

1	Yes	pat_costs_cmplt
0	No	

If you have responded 'yes', you will be contacted by WHO to provide information about the methods and results of the survey

3.1c Was a data quality review or inventory study completed in 2020 or 2021 to estimate the number of diagnosed TB cases that were not reported (i.e. not included in the TB surveillance system)?

1	Yes	dqr cmplt
0	No	agr_ompro

If you have responded 'yes', you will be contacted by WHO to provide information about the methods and results of the study

Non-routine surveillance of HIV prevalence in TB patients, 2021

3.2		-routine surveillance of	HIV prevalence in TB	patier view_TME_m	aster_strategy		
	sentinel sites in 2021? 1 Yes 0 No		surveil				
3.3	If yes, what so	urces of data were used	?				
		Estimated prevalence (%)	Year of estimate	95% Confid	ence interval upper limit		
	Nationwide surveys based on a representative sample of TB patients	tbhiv_surv_prev	tbhiv_surv_yr	tbhiv_surv_cil	tbhiv_surv_ciu		
	Sentinel sites	tbhiv_sentin_prev	tbhiv_sentin_yr	tbhiv_sentin_cil	tbhiv_sentin_ci		
PI Rema		x if data are not availabl	e for empty cells abo	ve.			
Digi	tal systems						
3.4	Are the numbers of TB cases reported in Sections 2.1-2.9 derived from a national, electronic, case-based database with separate records for each TB patient? A case-based system can show a list of individual cases and allows you to make specific calculations such as the "number of new extrapulmonary cases in males aged 29 notified during the first week of May". Using a spreadsheet such as Excel to store individual records does not meet the definition of a case-based database. 2 Yes, all TB patients in the country Yes, but only for MDR-TB patients countrywide Partially (transition to a case-based system is under way) No						

Contact investigation and TB preventive treatment, 2021

TPT.1 Do you have any data on the number of household contacts of TB cases who were evalu for TB?				
			screen_data_available	
	No Yes, available from our routine surveillance systems Yes, estimated from a review of a random samp of TB patients, representative of the national TB patie Not applicable (because there were no TB cases	le of n		
TPT.2	(if yes from routine surveillance) Number of household contacts of bacteriologically- confirmed pulmonary new and relapse TB cases notified in 2021	new	inc_con	
TPT.3	(if yes from routine surveillance) Among the household contacts reported in TPT.2, the number who were evaluated for TB disease and TB infection	new	inc_con_screen	
TPT.4	(if yes from review) Number of bacteriologically-confirmed pulmonary new and relapse TB cases notified in 2021 whose medical records or treatment cards were included in the review	rev	_newinc	
TPT.5	(if yes from review) Number of household contacts identified of the TB cases reviewed in TPT.4	rev _.	_newinc_con	
TPT.6	(if yes from review) Among the household contacts reported in TPT.5, the number who were evaluated for TB disease and TB infection	rev	_newinc_con_screen	
TPT.7	Do you have any data on the number of household copreventive treatment?	ntacts	s of TB cases started on TB	
			prevtx data_available	
	No Yes, available from our routine surveillance system Yes, estimated from a review of a random sample of medical records or treatment cards of TB patients, representative of the national TB patient population Not applicable (because there were no TB cases)			
TPT.8	(if yes from routine surveillance) Number of household contacts of bacteriologically- confirmed pulmonary new and relapse TB cases notified in 2021 who were started on TB preventive treatment	new	inc_con_prevtx	
TPT.9	(if yes from routine surveillance) Among the household contacts started on TB preventive treatment reported in TPT.8, number who were children aged under 5 years.	new	inc_con04_prevtx	
	(if was from ravious)	n+	muon noning	
121.10	(if yes from review) Number of bacteriologically-confirmed pulmonary new and relapse TB cases notified in 2021 whose	ptsi	rvey_newinc	

	medical records or treatment cards were included in the review	
TPT.11	(if yes from review) Number of household contacts of the TB cases reviewed in TPT.10 who were started on TB preventive treatment,	<pre>ptsurvey_newinc_con_prevtx</pre>
TPT.12	(if yes from review) Among the household contacts started on TB preventive treatment reported in TPT.11, number who were children aged under 5 years.	<pre>ptsurvey_newinc_con04_prevtx</pre>

Shorter TB preventive treatment regimens

TPT.13 Were any shorter TB preventive treatment regimes containing rifampicin or rifapentine used at least once in 2021?

		tpt_short_regimens_used
1	Yes	
0	No	
3	Don't know	

TPT.14 (if TPT13 is yes)

Were any of the shorter TB preventive treatment regimens below used at least once in 2021?

1 month daily rifapentine and isoniazid (1HP)	tpt_1hp
	1 Yes 0 No 3 Don't know
3 month weekly rifapentine and isoniazid (3HP)	tpt_3hp
	1 Yes 0 No 3 Don't know
3 month daily rifampicin and isoniazid (3HR)	tpt_3hr
	1 Yes 0 No 3 Don't know
44 month daily rifampicin (4R)	tpt_4r
	1 Yes 0 No 3 Don't know
TPT.15 (if TPT13 is yes) Total number of individuals started on shorter TB preventive treatment regimens containing rifampicin or rifapentine in 2021	<pre>prevtx_short_rifamycin</pre>

(please leave empty if the number is not available) (please also include data from national HIV/AIDS programme and research
projects)

Completion of TB preventive treatment, 2020			
TPT.16	Do you have any data on the number of household co TB preventive treatment in 2020 and who completed t		
	□ No	prevtx_cmplt_data_available	
	Yes Not applicable (because there were no TB cases	s)	
TPT.17	Number of household contacts of bacteriologically- confirmed pulmonary new and relapse TB cases who were started on TB preventive treatment in 2020 (as reported to WHO in 2021)	newinc_con_prevtx	
TPT.18	Among household contacts reported in TPT.17, number who completed the course of TB preventive treatment	newinc_con_prevtx_cmplt	
	Please tick the box if data are not available for empty cells above. Remarks:		

TB infection control

For long form countries only (dc_shortform=0)		
Data on TB in health workers is typically available from occupational health programmes.		
If infection control is effective, the annual TB incidence rates in health workers (relative to the total number of health workers) should not on average exceed annual TB incidence rates in the general population of the same age and sex groups.		
IC.1 How many workers at health care facilities (including non-medical staf such as drivers) had TB in 2021 (regardless of job position)?	hcw_tb_infected	
IC.2 How many workers at health care facilities (including non-medical staf such as drivers) were working in the country in the public and private sector in 2021?	hcw_tot	
Please tick the box if data are not available for empty cells above. Remarks:		

Laboratory diagnostic services

For all countries in AMR and EUR
For long form countries only (dc_shortform=0) in other regions

See Framework of indicators and targets for laboratory strengthening under the End TB Strategy

TB infection tests used in 2021

- LAB.1 Which tests of TB infection were used in the public or private sector before starting TB preventive treatment for any population at risk in 2021?
 - 175 Interferon Gamma Release Assays (IGRA)

176 Tuberculin Skin Test (TST)

177 IGRA and TST

194 Not used

3 Don't know

infection_tests_used

Molecular WHO-recommended rapid diagnostic testing in 2021

LAB.2 Total number of diagnostic tests performed using molecular WHO-recommended rapid diagnostics

These are currently Xpert MTB/RIF (including Ultra), Truenat MTB and MTB Plus, and TB-LAMP.

LAB.3 Number of positive results among the diagnostic tests performed using molecular WHO-recommended rapid diagnostics in LAB.2

Sites performing TB diagnostic testing at the end of 2021

LAB.4 Total number of sites providing laboratory diagnostic testing for TB at the end of 2021

Include all sites contributing to the diagnosis of TB, including laboratories within or outside the public health sector.

dx_test_sites

		(i) Number of sites providing these services at the end of 2021 a
LAB.5	Smear microscopy (including fluorescent)	smear
LAB.6	Culture	culture
LAB.7	Molecular WHO-recommended rapid diagnostics These are currently Xpert MTB/RIF (including Ultra), Truenat MTB and MTB Plus, and TB-LAMP.	m_wrd
LAB.8	Molecular tests for detection of isoniazid resistance (Low or moderate complexity automated nucleic acid amplification tests or first-line line probe assays)	m_inh

LAB.9	Molecular tests for detection of fluoroquinolone resistance (Low complexity automated nucleic acid amplification tests or second-line line probe assays)	m_fq
LAB.10	Drug susceptibility testing for pyrazinamide (MGIT or high complexity reverse hybridisation nucleic acid amplification tests)	dst_naat_pza
LAB.11	Phenotypic drug susceptibility testing for moxifloxacin and/or levofloxacin	dst_moxlev
LAB.12	Phenotypic drug susceptibility testing for bedaquiline	dst_bqd
LAB.13	Phenotypic drug susceptibility testing for linezolid	dst_1zd

^aPlease include all sites contributing to the diagnosis of patients notified by the NTP (including laboratories within or outside the public health sector).

Quality management systems among the sites performing TB testing at the end of 2021

LAB.14	Number of sites with ISO15189 accreditation	iso15189_accredited
LAB.15	Number of sites that are not accredited and that are in the process of establishing a formal quality management system towards achieving accreditation Implementation includes baseline assessment of the laboratory's quality management system using a recognized checklist based on ISO 15189, developing an action plan for quality improvements and starting to implement recommendations.	qms_pending

Data reporting among the sites performing TB testing at the end of 2021

LAB.16	Number of sites which record TB specimen reception and test results in a formal computer-based laboratory information system Using a spreadsheet such as Excel does not meet the definition of a formal computer-based laboratory information system.	lmis
LAB.17	Among the sites using molecular WHO-recommended rapid diagnostics reported in LAB.7, number of sites which transmit results automatically to clinicians and to an information management system. Electronic data connectivity solutions are able to rapidly make test results available to clinicians and information management systems (including a laboratory information management system or an electronic register, or both) via the Internet, mobile data networks or text messaging (SMS). See GLI Quick Guide to TB Diagnostics Connectivity Solutions	m_wrd_etrans
Please Remarks:	tick the box if data are not available for empty cells above.	

Public-Private Mix (PPM)

Fo	r 28	<pre>countries where dc_ppm_display = 1</pre>	
PF	PM.1	Number of TB cases contributed by private non-NTP providers through referral / diagnosis / notification in 2021 Include all contributions from private individual and institutional providers, corporate/business sector providers, mission hospitals, and other clinics/hospitals managed by non-governmental organizations and faith-based organizations	priv_new_dx
PF	PM.2	Number of TB cases contributed by public non-NTP providers through referral / diagnosis / notification in 2021 Include all contributions from public hospitals, public medical colleges, prisons/detention centres, military facilities, railways, public health insurance organizations etc	pub_new_dx
Re] Pleas	se tick the box if data are not available for empty cells above.	

Community Engagement

For 81 countries where dc_engage_community_display = 1		
CE.1	How many TB basic management units (BMUs) were there in 2021? A basic management unit (BMU) is defined in terms of management, supervision and monitoring responsibility. A BMU for the TB programme may have several treatment facilities, one or more laboratories and one or more hospitals. The defining aspect is the presence of a manager or coordinator who oversees TB control activities for the unit and who maintains a master register of all TB patients being treated. Typically, the units correspond to the government's second subnational administrative division, which might be called, for example, a "district" or "county".	bmu
CE.2	How many BMUs were implementing community-based referrals or any form of community treatment adherence support in 2021?	bmu_community_impl
CE.3	Do you have data on community-based referrals or any form of community treatment adherence support in 2021? 1 Yes 0 No	community_data_available

Referrals by community health workers / community volunteers in 2021

CE.4	Number of BMUs with data on referrals by community health workers / community volunteers in 2021	bmu_ref_data
CE.5	Total number of new and relapse TB cases notified in the BMUs of CE.4 in 2021	notified_ref
CE.6	Number of new and relapse cases referred by community health workers / community volunteers* among the cases in CE.5	notified_ref_community

Treatment adherence support from community health workers / community volunteers for patients who started TB treatment in 2020

CE.7	Number of BMUs with data on community treatment adherence support	bmu_rxsupport_data
CE.8	Total number of patients who started TB treatment in the BMUs of CE.7 in 2020	bmu_rxsupport_data_coh

CE.9	Total number of patients who started TB treatment in 2020 and who received any form of treatment adherence support from community health workers / community volunteers* in the BMUs of CE.7	rxsupport_community_coh
CE.10	Number of TB cases who were cured or who completed treatment among the cases in CE.9	rxsupport_community_succ
governme organization Community health ser Community through a	by health workers are people with some formal education who are given travices, and their time is often compensated by incentives in kind or in cash. By volunteers are community members who have been systematically sens short, specific training scheme or through repeated, regular contact sessions tick the box if data are not available for empty cells above	n-based organizations and patient-based nining to contribute to community-based nitized about TB prevention and care, either ons with professional health workers.

SECTION 4: FINANCE

Budget fiscal year 2022

For low/middle-income countries only (dc_finance_display=1)

		(i) Drug- susceptible TB treatment	(ii) MDR- TB treatment	(iii) pre-XDR / XDR-TB treatment	(iv) TB preventive treatment
4.1	Number of patients expected to start treatment in 2022	tx_dstb ^a	tx_mdr	tx_xdr	tx_tpt b
4.2	Average cost of drugs budgeted per patient, excluding buffer stock (US Dollars)	budget_cpp_d stb ^c	budget_cp p_mdr ^d	budget_cpp_ xdr ^e	budget_cpp _tpt ^f
4.3	Expected cost of drugs in 2022, excluding buffer stock (US Dollars)	(auto calc.)	(auto	calc.)	(auto calc.)

^a Include all patients receiving first-line drugs, including children and retreatment cases.

Please report all financial data in US Dollars. Please leave data items empty if their values are not known. Enter 0 only if the true value is zero.

	Budget line item	Budget required d	Expected funding ^e	Gap
4.4	Laboratory infrastructure, equipment and supplies Building, maintaining, and renovating TB laboratories, laboratory equipment purchase and maintenance, consumables for all tests (including TB screening for people living with HIV/AIDS and diagnosis of latent TB infection), quality assurance, retooling and the transportation of specimens.	budget_lab	cf_lab	gap_lab
4.5	National TB Programme staff (central unit staff and subnational TB staff) Salaries and incentives of those working only on TB activities at central and peripheral levels (for example provincial TB coordinators, district TB coordinators, etc.). Do not include primary health care personnel working on other diseases in addition to TB.	budget_staff	cf_staff	gap_staff
4.6	Drug-susceptible TB: drugs	budget_fld	cf_fld	gap_fld

b Include all people receiving TB preventive treatment

^C This can be estimated as the annual budget for first-line drugs (excluding any buffer stock) divided by the expected number all new and retreatment patients (adults and children).

^d This can be estimated as the annual budget for second-line drugs (excluding any buffer stock) divided by the expected number of patients who will be started on treatment for MDR-TB.

^e This can be estimated as the annual budget for pre-XDR/XDR-TB drugs (excluding any buffer stock) divided by the expected number of patients who will be started on treatment for pre-XDR/XDR-TB.

^f This can be estimated as the annual budget for drugs for TB preventive treatment (excluding any buffer stock) divided by the expected number of people who will be started on TB preventive treatment.

Drugs for patients being treated for drug- susceptible TB. Include children, re-treatment cases and buffer stock. 4.7 Drug-susceptible TB: programme costs The management and supervision of the TB control programme, training, policy development, meetings, visits for supervision, purchase of office equipment/vehicles, construction of buildings for use by programme staff, routine surveillance, advocacy and communication, public-private mix activities, community engagement, activities, community engagement activities finding, infection control, management of TB drug procurement and distribution, and programme activities linked to contact investigation for TB preventive treatment 4.8 Drug-resistant TB: drugs Drugs to treat drug-resistant TB (FR-TB, MDR- TB, pro-XDR-TB or XDR-TB). Include drugs to deal with adverse events 4.9 Drug-resistant TB: programme costs Management of drug-resistant TB services. Subcliding drugs. Exemples as removation of Committee, conducting an MDR situation assessment, default and contact tracing, palliative care. 4.10 TB preventive treatment: drugs Drugs for TB preventive treatment, as per latest WHO guidance (H-J, SH, AR, SHR and Levolioxacin) 4.11 Collaborative TB/HIV activities aimed at reducing the impact of HIV-related TB. Activities include TB/HIV coordinating bodies, join TB/HIV training and planning, HIV testing for TB patients, Irll Surveillance among TB patients, co- utmoxazole preventive threapy (CPT), joint TB/HIV training and planning, HIV testing for TB patients, educational and emotional support to patient or other in-kind benefits given to TB screening for people king with HIV/AIDS is included under (Lab infrastructure, equipment, and supplies). 4.12 Patient support Cash transfers, food packages, transportation vuchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (airtime or device for V.O.T), medications monitors (digital box, 39 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, dru					
The management and supervision of the TB control programme, training, policy development, meltings, visits for supervision, purchase of office equipment/vehicles, construction of buildings for use by programme staff, routine surveillance, advocacy and communication, public private mix activities, community engagement, active case-finding, infection control, management of TB drug procurement and distribution, and programme activities linked to contact investigation for TB preventive treatment 4.8 Drug-resistant TB: drugs Drugs to treat drug-resistant TB (RR-TB, MDR-TB, pre-XPR-TB or XPR-TB), Include drugs to deal with adverse events 4.9 Drug-resistant TB: programme costs Management of drug-resistant TB services, excluding drugs. Examples are renovation of MDR-TB wards, support for the Green Light Committee, conducting an MDR situation assessment, default and contact tracing, palliative care. 4.10 TB preventive treatment: drugs Drugs for TB preventive treatment; drugs Drugs for TB preventive treatment; drugs Drugs for TB preventive treatment, as per latest WHO guidance (6H, 9H, 4R, 3HR and Levofloxacin) 4.11 Collaborative TB/HIV activities Collaboration between TB and HIV programmes aimed at reducing the impact of HIV-related TB. Activities include TB/HIV activities (TB/HIV coordinating bodies, joint TB/HIV ratining and planning, HIV estings for TB patients, HIV surveillance among TB patients, co-trimoxacole preventive therapy (CPT), joint TB/HIV education/ communication, and antiretroviral teatment therapy (CPT), joint TB/HIV education and emotional support to patient or other in-this benefits given to TB patients, mobile phone (aitime or device for V.O.T), medications monitors (digital box, 99 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, drug resistance, patient classtrophic cost; routine surveillance (epidemiology review, inventory studies, pharmacoviglance, systematic assessment of the					
Drugs to treat drug-resistant TB. (RR-TB, MDR-TB, pre-XDR-TB or XDR-TB). Include drugs to deal with adverse events 4.9 Drug-resistant TB: programme costs Management of drug-resistant TB services, excluding drugs. Examples are renovation of MDR-TB wards, support for the Green Light Committee, conducting an MDR situation assessment, default and contact tracing, palliative care. 4.10 TB preventive treatment: drugs Drugs for TB preventive treatment, as per latest WHO guidance (6H, 9H, 4R, 3HR and Levofloxacin) 4.11 Collaborative TB/HIV activities Collaboration between TB and HIV programmes aimed at reducing the impact of HIV-related TB. Activities include TB/HIV condinating bodies, joint TB/HIV training and planning. HIV testing for TB patients, HIV surveillance among TB patients, co-trimoxazole preventive therapy (CPT), joint TB/HIV education/ communication, and antiretroviral treatment for TB patients. TB screening for people living with HIV/AIDS is included under (Lab infrastructure, equipment, and supplies). 4.12 Patient support Cash transfers, food packages, transportation vouchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (aitime or device for V.O.T), medications monitors (digital box, 99 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovygliance, systematic assessment of the	4.7	The management and supervision of the TB control programme, training, policy development, meetings, visits for supervision, purchase of office equipment/vehicles, construction of buildings for use by programme staff, routine surveillance, advocacy and communication, public-private mix activities, community engagement, active casefinding, infection control, management of TB drug procurement and distribution, and programme activities linked to contact investigation for TB	budget_prog	cf_prog	gap_prog
Management of drug-resistant TB services, excluding drugs. Examples are renovation of MDR-TB wards, support for the Green Light Committee, conducting an MDR situation assessment, default and contact tracing, palliative care. 4.10 TB preventive treatment: drugs Drugs for TB preventive treatment, as per latest WHO guidance (6H, 9H, 4R, 3HR and Levofloxacin) 4.11 Collaborative TB/HIV activities Collaboration between TB and HIV programmes aimed at reducing the impact of HIV-related TB. Activities include TB/HIV coordinating bodies, joint TB/HIV training and planning, HIV testing for TB patients, HIV surveillance among TB patients, co-trimoxazole preventive therapy (CPT), joint TB/HIV education/ communication, and antiretroviral treatment for TB patients. TB screening for people living with HIV/AIDS is included under (Lab infrastructure, equipment, and supplies). 4.12 Patient support Cash transfers, food packages, transportation vouchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (airtime or device for V.O.T), medications monitors (digital box, 99 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovigilance, systematic assessment of the	4.8	Drugs to treat drug-resistant TB (RR-TB, MDR- TB, pre-XDR-TB or XDR-TB). Include drugs to	budget_sld	cf_sld	gap_sld
Drugs for TB preventive treatment, as per latest WHO guidance (6H, 9H, 4R, 3HR and Levofloxacin) 4.11 Collaborative TB/HIV activities Collaboration between TB and HIV programmes aimed at reducing the impact of HIV-related TB. Activities include TB/HIV coordinating bodies, joint TB/HIV training and planning, HIV testing for TB patients, HIV surveillance among TB patients, cotrimoxazole preventive therapy (CPT), joint TB/HIV education/ communication, and antiretroviral treatment for TB patients. TB screening for people living with HIV/AIDS is included under (Lab infrastructure, equipment, and supplies). 4.12 Patient support Cash transfers, food packages, transportation vouchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (airtime or device for V.O.T), medications monitors (digital box, 99 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovigilance, systematic assessment of the	4.9	Management of drug-resistant TB services, excluding drugs. Examples are renovation of MDR-TB wards, support for the Green Light Committee, conducting an MDR situation assessment, default and contact tracing, palliative	budget_mdrmgt	cf_mdrmgt	gap_mdrmgt
Collaboration between TB and HIV programmes aimed at reducing the impact of HIV-related TB. Activities include TB/HIV coordinating bodies, joint TB/HIV training and planning, HIV testing for TB patients, HIV surveillance among TB patients, co-trimoxazole preventive therapy (CPT), joint TB/HIV education/ communication, and antiretroviral treatment for TB patients. TB screening for people living with HIV/AIDS is included under (Lab infrastructure, equipment, and supplies). 4.12 Patient support Cash transfers, food packages, transportation vouchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (airtime or device for V.O.T), medications monitors (digital box, 99 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovigilance, systematic assessment of the	4.10	Drugs for TB preventive treatment, as per latest WHO guidance (6H, 9H, 4R, 3HR and	budget_tpt	cf_tpt	gap_tpt
Cash transfers, food packages, transportation vouchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (airtime or device for V.O.T), medications monitors (digital box, 99 DOTS) 4.13 Operational research and surveys Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovigilance, systematic assessment of the	4.11	Collaboration between TB and HIV programmes aimed at reducing the impact of HIV-related TB. Activities include TB/HIV coordinating bodies, joint TB/HIV training and planning, HIV testing for TB patients, HIV surveillance among TB patients, cotrimoxazole preventive therapy (CPT), joint TB/HIV education/ communication, and antiretroviral treatment for TB patients. TB screening for people living with HIV/AIDS is included under (Lab infrastructure, equipment,	_	cf_tbhiv	gap_tbhiv
Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovigilance, systematic assessment of the	4.12	Cash transfers, food packages, transportation vouchers, educational and emotional support to patient or other in-kind benefits given to TB patients, mobile phone (airtime or device for V.O.T), medications monitors (digital box, 99	budget_patsup	cf_patsup	gap_patsup
	4.13	Periodic surveys (prevalence, drug resistance, patient catastrophic cost); routine surveillance (epidemiology review, inventory studies, pharmacovigilance, systematic assessment of the	budget_orsrvy	cf_orsrvy	gap_orsrvy
4.14 All other budget lines budget_oth cf_oth gap_oth	4.14	All other budget lines	budget_oth	cf_oth	gap_oth

	Please explain this amount in the "Remarks" box below.			
4.15	Total	budget_tot	cf_tot	gap_tot

^dTotal budget required should be in line with your annual national strategic plan. Indicate total amount that is necessary to carry out the National plan, not just the expected disbursements from funding partners.

Please enter the **total expected funding** for the budget required shown above:

	Funding source	Expected funding
4.16	Domestic (including loans)	cf_tot_domestic
4.17	Global Fund	cf_tot_gf
4.18	USAID	cf_tot_usaid
4.19	Other sources	cf_tot_grnt
4.20	Total expected funding	cf_tot_sources

Please tick the box if data are not available for empty cells above.	
Remarks:	

^e Funding from both the central and peripheral government, Global Fund, USAID, and other grants. The amount should be for the relevant fiscal year only and not the total amount of the grants or commitments over several fiscal periods.

Expenditure, fiscal year 2021

For low/middle-income countries only (dc_finance_display=1)

4.21	Average cost of drugs spent per patient starting first-line TB treatment, excluding buffer stock (US Dollars) This can be estimated as the annual expenditure for first-line drugs (excluding any buffer stock) divided by the total number of notifications of all new and retreatment patients (adults and children).	exp_cpp_dstb
4.22	Average cost of drugs spent per patient starting second-line treatment for MDR-TB, excluding buffer stock (US Dollars) This can be estimated as the annual expenditure for second-line drugs (excluding any buffer stock) divided by the number of patients enrolled on treatment for MDR-TB.	exp_cpp_mdr
4.23	Average cost of drugs spent per patient starting pre-XDR / XDR-TB treatment, excluding buffer stock (US Dollars) This can be estimated as the annual expenditure for XDR-TB drugs (excluding any buffer stock) divided by the number of patients enrolled on treatment for pre-XDR-TB / XDR-TB.	exp_cpp_xdr
4.24	Average cost of drugs spent per person starting TB preventive treatment	exp_cpp_tpt

Please report all financial data in US Dollars. Please leave data items empty if their values are not known. Enter 0 only if the true value is zero.

		Actual expenditure ^a	Received Funding ^b
4.25	Laboratory infrastructure, equipment and supplies	exp_lab	rcvd_lab
4.26	National TB Programme staff (central unit staff and subnational TB staff)	exp_staff	rcvd_staff
4.27	Drug-susceptible TB: drugs	exp_fld	rcvd_fld
4.28	Drug-susceptible TB: programme costs	exp_prog	rcvd_prog
4.29	Drug-resistant TB: drugs	exp_sld	rcvd_sld
4.30	Drug-resistant TB: programme costs	exp_mdrmgt	rcvd_mdrmgt
4.31	TB preventive treatment: drugs	exp_tpt	rcvd_tpt
4.32	Collaborative TB/HIV activities	exp_tbhiv	rcvd_tbhiv
4.33	Patient support	exp_patsup	rcvd_patsup
4.34	Operational research and surveys	exp_orsrvy	rcvd_orsrvy
4.35	All other budget lines for TB	exp_oth	rcvd_oth
4.36	TOTAL	exp_tot	rcvd_tot

^a Report the amounts that were actually spent on each line item during your last fiscal year. The total in this column might be lower than the total funds received, but not higher.

^b Report the funds actually received from each source of funding. The total amount from all sources might be higher than the expenditure reported, but not lower.

Please enter the **total received funding** for the actual expenditures in the table above:

	Source	Received funding
4.37	Domestic (including loans)	rcvd_tot_domestic
4.38	Global Fund	rcvd_tot_gf
4.39	USAID	rcvd_tot_usaid
4.40	Other sources	rcvd_tot_grnt
4.41	Total received funding	rcvd_tot_sources

Please tick the box if data are not available for empty cells above.
Remarks:

exp res

For all WB high income countries and the 30 HBCs only $\,$

TB research expenditure

RES.1 National expenditure on tuberculosis research, **excluding** expenditure already reported in question 4.34, in fiscal year 2021 (US Dollars)

Comprehensive spending on TB research includes epidemiological surveillance, operations research, patient cost surveys, health systems research, clinical research and fundamental (basic science) research. This information may be available from the Ministry of Health and/or national science and technology council or its equivalent.

	Please tick the box if data are not available for empty cells above.
Rer	marks:

	Patients starting first-line TB treatment	Patients starting MDR-TB / pre- XDR-TB / XDR-TB treatment
UTL.1 Typical number of visits to a health facility after diagnosis The average number of visits per patient to any health facility during TB treatment, for example for observed treatment (DOT), collection of drugs, smear monitoring, etc. after the patient has been diagnosed with TB, in view of your treatment guidelines. For example, if a TB patient on first-line treatment receives directly observed treatment daily in the intensive phase at clinics and, in the continuation phase 4 visits are required (one per month for collection of drugs), the total would be 60+4=64.	hcfvisit_dstb	hcfvisit_mdr
UTL.2 Estimated percentage of cases that are hospitalized (%) If the actual percentage of hospitalisations is available from the basic management unit register, please report. If not, please report the approximate percentage of patients hospitalized for TB treatment (for any duration of stay), in view of your treatment guidelines. For example, if your policy or general practice is to admit all TB patients for 2 months, the figure will be 100%.	hospd_dstb_prct	hospd_mdr_prct
UTL.3 Estimated average duration of stay if hospitalized (days) If the actual duration of stay is available from the basic management unit register, please report. If not, please estimate the number of days a patient would spend in hospital "on average".	hospd_dstb_dur	hospd_mdr_dur

UTL.4 If MDR-TB patients are hospitalized, in which type of facility are they most often treated?

140	Primary-level hospital	hosp_type_mdr
141	Secondary-level hospital	1-71 -
142	Tertiary-level hospital	
2	Not applicable	

Primary-level hospital (or 'district hospital' or 'first-level referral'): has few specialities, mainly internal medicine, obstetrics-gynecology, pediatrics, and general surgery, or only general practice; limited general laboratory services; 30-200 beds.

Secondary-level hospital (or 'provincial hospital'): highly differentiated by function with five to ten clinical specialities; 200-800 beds.

Tertiary-level hospital (or 'central' or 'regional' hospital): highly specialized staff and technical equipment, e.g., cardiology, ICU and specialized imaging units; clinical services are highly differentiated by function; may have teaching activities; 300-1500 beds.

may have teaching activities, 500 1000 beas	•
Source: WHO guide to cost effectiveness and	<u>alysis</u> p215
Please tick the box if data are not available t Remarks:	or empty cells above.

SECTION 5: MULTI-SECTORAL ACCOUNTABILITY

Only for countries that did not answer 'yes' to 5.1 and 5.2 in the 2020 or 2021 data collection form and also the 30 high TB burden countries (dc_unhlm_display=1)

Response to the political declaration of the 2018 UN High Level Meeting on TB		
Ann	ual report	
5.1	Does the National TB Programme (or equivalent) produce a publicly available annual report about the status of the TB epidemic and progress in response efforts? 1 Yes	annual_report_published
	0 No	
Review mechanism		
5.2	Is there a national multi-sectoral and multi-stakeholder accountability and review mechanism, under high-level leadership, to monitor and review progress towards ending TB? See https://www.who.int/tb/publications/MultisectoralAccountability/en/ Yes No	ms_review
5.3	(If yes to 5.2) Do representatives of civil society and affected communities participate in the multi-sectoral accountability and review mechanism? 1 Yes 0 No	ms_review_civil_soc
Ques	tions 5.4 to 5.9 only for the 30 high TB burden	countries
Soc	ial protection	
5.4	Is there a national policy to specifically provide social protection services to TB-affected individuals?	social_protn
	1 Yes 0 No	
If yes to 5.4, please indicate which of the following apply:		

5.4.1	Free access to medical services Yes No Don't know	free_access
5.4.2	Enablers to adhere to TB treatment Yes No Don't know	enable_tx_adherence
5.4.3	Conditional cash-transfers 1 Yes 0 No 3 Don't know	cond_cash_trans
5.4.4	Measures to ensure food security 1 Yes 0 No 3 Don't know	food_security
5.4.5	Other (please describe briefly or leave empty if not applicable)	other_social_protn
Prote	ection from stigma and discrimination	
	indicate which aspects of life of people with TB are protected from T nination through national laws and regulations:	B stigma and
5.5	Employment (e.g. not being dismissed because of a TB diagnosis) Yes No Don't know	protect_employment
5.6	Housing (e.g. not being evicted from housing facilities because of a TB diagnosis) 1 Yes 0 No 3 Don't know	protect_housing

5.7	Parental rights (e.g. not having parental rights over children affected in any way as a result of TB diagnosis of a parent) 1 Yes 0 No 3 Don't know	protect_parenting
5.8	Freedom of movement (e.g. no restriction to access any congregate setting or transit through any geographical area because of a TB diagnosis) 1 Yes 0 No 3 Don't know	protect_movement
5.9	Freedom of association (e.g. no compulsory isolation because of a TB diagnosis) 1 Yes 0 No 3 Don't know	protect_association