





Management of DR-TB in children and adolescents – implementation considerations

Kerri Viney, WHO Global Programme on Tuberculosis and Lung Health Joint European regional workshop to plan the accelerated adoption and uptake of new WHO TB policies

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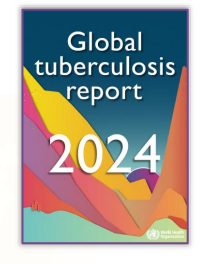
TB incidence and mortality in children and adolescents, 2023

10.8 million

TB among all ages in 2023



TB deaths in 2023



1.25 million

children (0-14 years) developed TB in 2023 (12% of all TB)



727 000 adolescents

(10-19 year-olds) developed TB in 2012 (Snow et al, 2018)

191 000

TB deaths in 2023 (15% of all TB deaths)



Among deaths in HIV-negative children and young adolescents 0–14

73% were in children <5 years



96% of deaths occurred in children who did not access TB treatment

(Dodd et al, 2017)



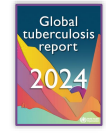
25 000

(14%) TB deaths in the 0–14 year age group were among children living with HIV

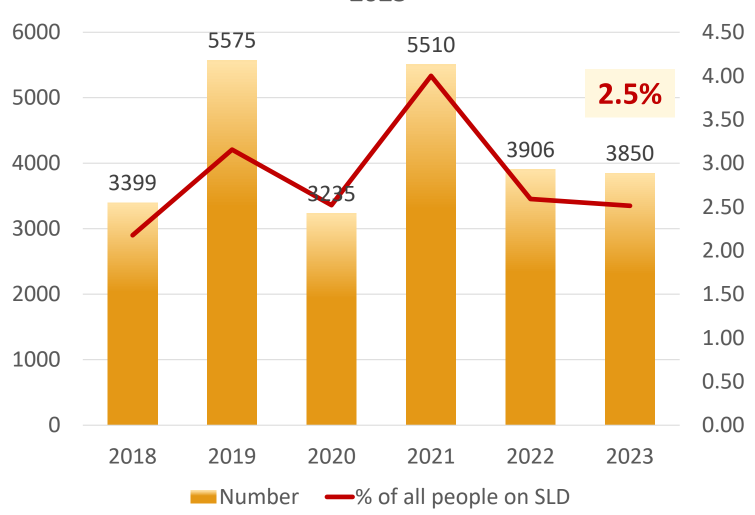




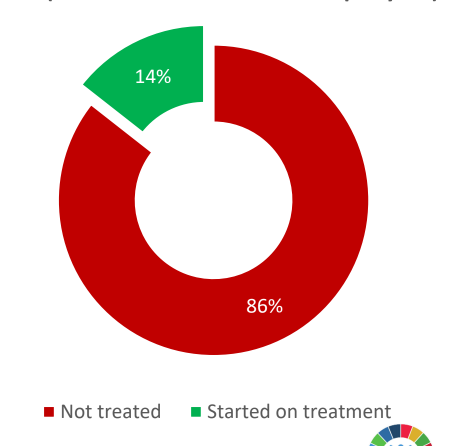
Treatment initiation in children with MDR/RR-TB (global)



Second-line treatment initiation in 0-14y, 2018-2023



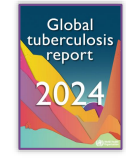
MDR/RR-TB treatment coverage in children and young adolescents, average for 2018-2023 (out of an estimated 30 000 per year)

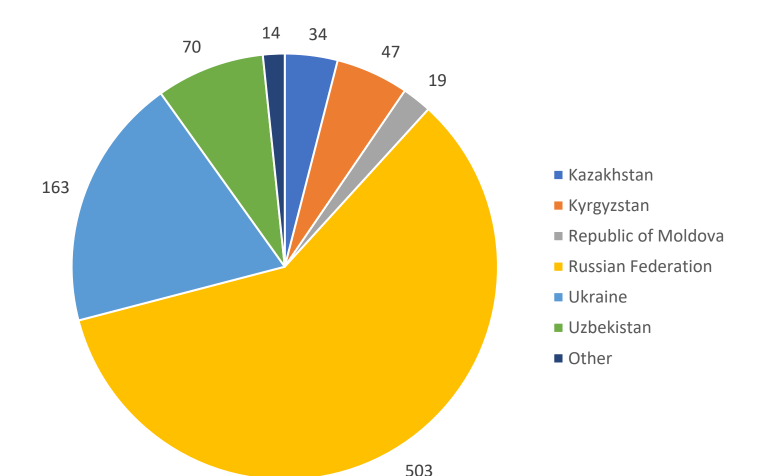




Second-line treatment initiation for DR-TB in <15 years in the European Region, 2023

MDR/RR-TB <15 years started on treatment, EUR





0-14 years =
2.5% of all MDR/RR-TB
initiated on second-line
treatment
(850 out of 33 986)

- Almost 60% in the Russian Federation
- Almost 20% in Ukraine



Other: countries with less than 10 children started on treatment: Armenia, Azerbaijan, Belarus, Georgia, Portugal, Tajikistan, Türkiye



Reasons for the treatment coverage gap



Characteristics of children and adolescents with multidrugresistant and rifampicin-resistant tuberculosis and their association with treatment outcomes: a systematic review and individual participant data meta-analysis

Anthony J Garcia-Prats*, Maria Garcia-Cremades*, Vivian Cox, Tamara Kredo, Rory Dunbar, H Simon Schaaf, James A Seddon, Jennifer Furin, Jay Achar, Kendra Radke, Tina Sachs, Amanzhan Abubakirov, Saman Ahmed, Onno W Akkerman, Nadia Abdulkareem Al Ani, Farhana Amanullah, Nafees Ahmad, Laura F Anderson, Meseret Asfaw, Funeka Bango, Torsten Bauer, Mercedes Becerra, Martin Boeree, Folke Brinkmann, Rosemary Brown, James Brust, Jonathon R Campbell, Anna Cristina Carvalho, Isabel Carvalho, J Peter Cegielski, Rosella Centis, Edward D Chan, Sandeep Chauhan, Silvia S Chiang, Pei-Chun Chan, Lia D'Ambrosio, Margareth Dalcolmo, Narine Daneilyan, Gerard de Vries, Heather R Draper, the endTB Study Group, Lee Fairlie, Joshua R Francis, Molly Franke, Medea Gegia, Camilo Gomez Restrepo, Annette Guenther, Tatyana Gureva, Brit Haecker, Elizabeth Harausz, Catherine Hewison, Robert M Hicks, Helena Huerga, Jennifer Hughes, Petros Isaakidis, Syed M Kadri, Mazhar Ali Khan, Tinatin Kotrikadze, Liga Kuksa, Nathalie Lachenal, Christoph Lange, Leonid Lecca, Elisa Lopez-Varela, Sheila Lucena, Andrei Mariandyshev, Sanjay Mattoo, Ana Mendez-Echevarria, Giovanni Battista Migliori, Carole Mitnick, Erika Mohr-Holland, Winston Mulanda, Totugul Murzabakova, Bakyt Myrzalieve, Norbert Ndjeka, Stefan Niemann, Iveta Ozere, Nesri Padayatchi, Malik Parmar, Nargiza Parpieva, Mohammad Manzur-Ul-Alam, Natasha Rybak, Kuldeep Singh Sachdeva, Kelly Salmon, Begoña Santiago-Garcia, Dagmar Schaub, Ira Shah, Sarita Shah, Vaibhav Shah, Sangeeta Sharma, Tae Shun Shim, Sonya Shin, Animesh Sinha, Alena Skrahina, Hardik Solanki, Belen P Solans, Antoni Soriano-Arandes, Atyrkul Toktogonova, Tjip van der Werf, Gustavo E Velásquez, Bhanu Williams, Jae-Joon Yim, Rada Savic†, Anneke Hesseling†

Summary

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See Comment page 78
*Contributed equally
†Contributed equally

Background There are few data on the treatment of children and adolescents with multidrug-resistant (MDR) or rifampicin-resistant (RR) tuberculosis, especially with more recently available drugs and regimens. We aimed to describe the clinical and treatment characteristics and their associations with treatment outcomes in this susceptible population.

Paediatric drug-resistant TB individual patient database:

- High % of adolescents
- High % of bacteriological confirmation



Suggesting:

- Young children with DR-TB not detected
- Treatment seldomly started in absence of bacteriological confirmation



Risk factors for MDR/RR-TB in children

Risk factors for MDR/RR-TB in children and adolescents

- Exposure to person with confirmed DR-TB
- Exposure to person who failed TB treatment or who died from TB
- Non-response to first-line TB treatment
- Previous TB treatment



Children with a decision to start treatment based on the treatment decision algorithms need to be assessed for risk of DR-TB

Contact investigation:

a critical intervention to identify children and adolescents exposed to DR-TB







Evaluation and diagnosis

- High index of suspicion needed
- Bacteriological testing critical
 - <u>Concurrent testing</u> with different samples (respiratory and stool) increases sensitivity (fewer false negative results)
- If bacteriological testing negative or cannot be done, a clinical diagnosis can be made (include detailed guidance in national guidelines)
- The DST pattern of the child/adolescent or the most likely source case informs treatment





Treatment of DR-TB in children – use of bdq & dlm in children

- WHO consolidated guidelines on tuberons Managara of aborton and adolescents.
- In children with MDR/RR-TB aged below 6 years, an all-oral treatment regimen containing bedaquiline may be used
- In children with MDR/RR-TB aged below 3 years, delamanid may be used as part of longer regimens

(NEW: both conditional recommendations, very low certainty of the evidence)

Remarks:

- Applies to and complements current WHO recommendations on shorter and longer regimens that contain bedaquiline
- Complements the current WHO recommendation on longer regimens that contain delamanid

These recommendations make it possible to build all oral regimens for children of all ages





Information notes on bedaquiline and delamanid





Use of bedaquiline in children and adolescents with multidrug- and rifampicin-resistant tuberculosis - Information note



To provide practical guidance on the administration of badequiline in children and adolescents in the context of the treatment of multidrug- and riferration-resistant suberculosis (MDR/RR-TB), in line with the latest World Health Organization (WHO) recommendations, dosing guidance and available formulations.

Doctors, clinicians, peedletricians, nurses, pharmacists, perents and caregivers of children with MDR/RR-TB, community health workers, programme managers, implementing partners and partners providing technical assistance.

WHO recommendations for bedaquiline in children and adolescents

The United States Food and Drug Administration granted accelerated approval for bedequaine in 2012 for the treatment. of adults aged 18 years and over with multidrug-resistant. pulmonary TB (MDR-TB) for whom an effective treatment. regimen could not otherwise be composed (1). This approval was based on phase IIb trial data and made bedaquiline the first medicine from a new dass approved with a TB indication

Since then, additional evidence has been generated on the use of bedequiline for the treatment of MDR/RR-TB in both adults and children. Bedequiline has played an increasingly important role in TB treatment as a component of both shorter and longer regimens, and has allowed the move away from injectable-containing regimens to all-oral regimens (2).

Bedaquiline - a key medicine in WHOrecommended regimens

- Bedequiline is now recommended by WHO for the treatment of MDR/RR-TB in adults and children of all ages (3).
- Bedequiline is a component of the 9-month all-oral regimen, which is the treatment of choice for eligible people aged under 14 years with MDR/RR-TB rather than longer (18 month) regimens.

- For people aged 14 years and over with MDR/RR-TB. WHO suggests the use of a 6-month treatment regimen. composed of bedaquiline, pretomanid, linezolid and modificracin (BPaLM) rather than the 9-month or longer. (18 month) regimens. In cases of documented resistance to fluoroquinolones, BPaL without moxifloracin would be initiated or continued (4).
- Bedaquiline is a group A medicine and a core component of longer individualized regimens for people who are not eligible for the 9-month all-oral or BPsLM/BPsL regimens.

Bedaguiline can be used as part of short and long all-oral WHO-recommended regimens for people with MDR/RR-TB of all ages.

- · Bedequiline is usually given for 6 months. This may be extended to the entire duration of the 9-month all-oral regimen if the initial phase of the regimen is extended from 4 to 6 months, if sputum is positive after 4 months
- When used as part of a longer regimen in people with fluoroguinolone resistance or with limited treatment options, the extension of bedaquiline beyond 6-9 months may be considered (off-label use), with strict baseline and follow-up monitoring. For children, this should be done in consultation with an expert in paediatric drug-resistant TB.

9-month all-ond regimen: Initial phase: 4-6 months of bedeguline, levoftosoin or mosflosoin, clofedwine, purchamide, other buto, high-dose

Continuation phase 5 months of levofloracin or modfloracin, clofazimine, pyrazinamide and ethambatol.

Group A medicines: Include involves in a world out in. bedaguiline and in codid. These would be swere found to be highly effective in improving treatment outcomes and reducing deaths. It is strongly recommended that they are used for all people with MDR/RRTB eligible for longer regimens unless there is a toxicity issue or drug resistance.

Longer individualized regimens: As a group A medicine, bedaquitne should be included in individualized MDR/RR-TB regimens for both fluoroquinolone-ausces tible and fluoroquinolone resistant treatment, unless bedaquiline resistance has been detected.

Possible individualized MDR/RR-TII regiment for children of all ages and adolescents can be found in Section 5.3.2.4 (Table 512) of the WHO Operational Handbook on Tuberculous, Module 5 Management of Tuberculous in Onlides and Adolescents (S).



DELAMANID

Use of delamanid in children and adolescents with multidrug- and rifampicin-resistant tuberculosis - Information note



To provide practical guidance on the administration of delamanid in children and addescents in the context of the bestment of multidrug- and rifempicin-resistant tuberculosis (MDR/RR-TB), in line with the latest World Health Organization (WHO) recommendations, dosing guidence and available formulations.

Doctors, clinicians, psedistricians, nurses, pharmacists, parents and caregivers of children with MDR/RR-TB, community health workers, programme managers, implementing partners and partners providing technical

WHO recommendations for delamanid in children and adolescents

The European Medicines Agency granted conditional approval to delamanid in 2014 "as part of an appropriate combination regimen for pulmonary multidrug-resistant tuberculosis in adult patients (≥18 years of age) when an effective treatment regimen cannot otherwise be composed for reasons of resistance or tolerability" (1). This made delamanid the second new medicine from a new class approved with a TB indication, following on from bedaquiline.

Since then, additional evidence has been generated on the use of delamanid for the treatment of MDR/RR-TB in both adults and children. Its use has expanded the list of medicines available to design all-oral longer individualized regimens for people with MDR/RR-TB, moving ewey from toxic injectable agents. The availability of delamanid is particularly important. for people, including children, with limited options due to a more extensive resistance profile.

Delamantd can be used as part of individualized longer regimens for people of all ages with MDR/RR-TB.

Delamanid - a medicine for people of all ages with limited treatment options

- Delamanid is now recommended by WHO for the treatment. of MDR/RR-TB in adults and children of all ages (2, 3).
- Delamanid is a group C medicine and can be used as part of longer individualized regimens for people with MDR/RR-TB, including children and adolescents, who are not eligible for the 9-month all-oral regimen or the 6-month regimen composed of bedaquiline, pretomanid and linezolid, with or without moxifloracin (BPaLM/BPaL).
- As a group C medicine, delamanid can be included in MDR/RR-TB regimens when a treatment regimen cannot be composed of group A or B agents alone, due to resistance or intolerance.

. Delamanid is usually given for 6 months. The duration may 6 months (when given alongside other medicines, including bedaquiline) is safe (4, 5).

be extended beyond 6 months (off-label use) in people, including children, with fluoroguinolone resistance or with limited treatment options. Studies undertaken between 2020 and 2022 showed that the use of delamanid beyond

Group A medicines: holade levollosacin or modificacin, bedaquiline and inspoid. Group B medicines; include clofazimine and dycloserine or terisidons.

Group C medicines: Include ethanitatal, defensarid, pyradinanide, imiperem-clastatin or meropenem in combination with classificacid, amiliacin or streptomych lionly used as salvage though in children and addissents aged under 16 years), othionamide or profitionamide and Plannin osalicylic acid Group C medicines are included in longer regimens if the regimen cannot be composed of Group A and B medicines alone.

Longor individual tend regimens: Exemples of individualized MDR/RR-TB regimens for children of all ages and adolescents can be found in Section 5.3.2.4 (Table 5.32) of the WHO Operational Handbook on Tuberculosis, Module 5: Management of Tuberculosis in Children and Adolescents (6).



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Treatment of DR-TB in children – updates

- **BEAT-TB trial** in South Africa 6-month Bdg-Lzd-Dlm-Lfx/Cfz (or both) vs Standard of Care
- New recommendation:

WHO suggests the use of a 6-month treatment regimen composed of bedaquiline, delamanid, linezolid (600 mg), levofloxacin, and clofazimine (BDLLfxC) in MDR/RR-TB patients with or without fluoroquinolone resistance

(Conditional recommendation, very low certainty of evidence)

FQ S FQ R Applies to (among others):

RR TB Diagnosed Assessed for Randomisation **Control Strategy BEAT Tuberculosis** South African RR TB (BDQ, DLM, LNZ, LVX, Standard of Care Diagnostic Diagnostic Triage Triage FQ sensitive FQ sensitivity not established FQ resistant FQ sensitive FQ sensitivity not established FQ resistant Individualised Continue BEAT Continue Standard of Continue Standard of regimen with BDQ. BDQ, DLM, LNZ, LVX BDQ, DLM, LNZ, CFZ **Tuberculosis** LNZ. DLM and other Regimen

- - PTB TB, including children, adolescents, PLHIV, pregnant and breastfeeding women
 - EPTB except CNS, osteoarticular, or disseminated forms of TB with multi-organ involvement
 - Children and adolescents without bacteriological confirmation of TB or DST but with a high likelihood of MDR/RR-TB (based on clinical signs and symptoms of TB, in combination with a history of contact with a patient with confirmed MDR/RR-TB)



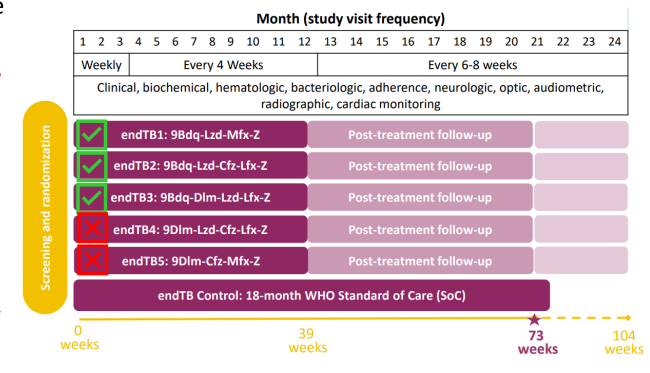


Treatment of DR-TB in children – updates

- endTB trial 9-month regimens vs Standard of Care
- New recommendation:

WHO suggests using the 9-month all-oral regimens (BLMZ, BLLfxCZ and BDLLfxZ) over currently recommended longer (>18 months) regimens in patients with MDR/RR-TB and in whom resistance to fluoroquinolones has been excluded. Amongst these regimens, using BLMZ is suggested over BLLfxCZ, and BLLfxCZ is suggested over BDLLfxZ

(Conditional recommendation, very low certainty of evidence)



FQ S

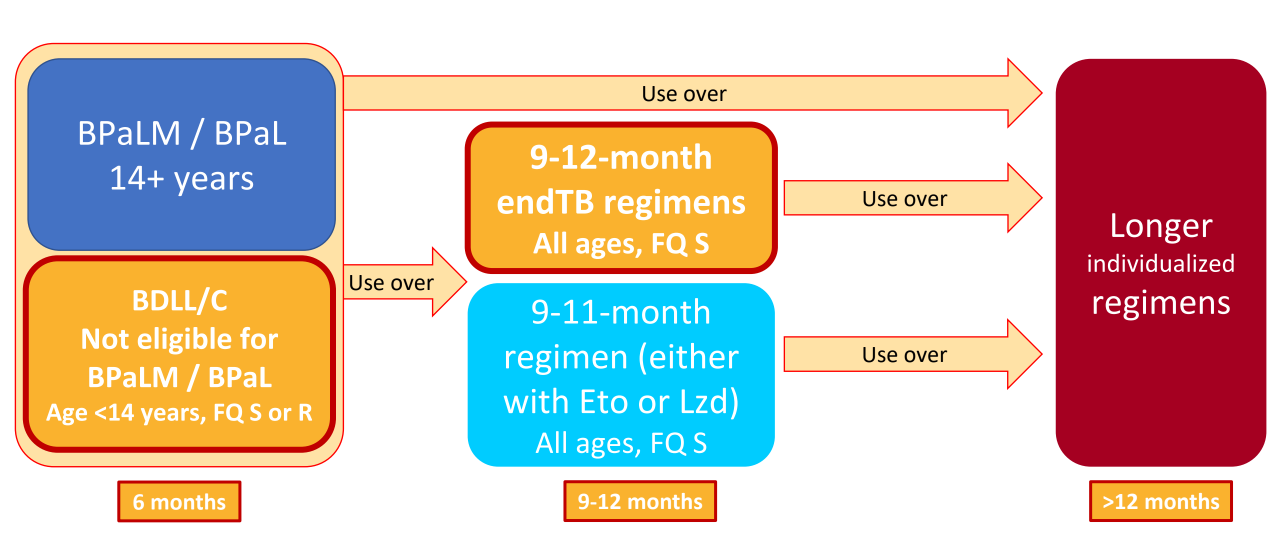
× FQ R

- Applies to (among others):
 - a. PTB TB, including children, adolescents, PLHIV, pregnant and breastfeeding women
 - b. EPTB except CNS, osteoarticular, or disseminated forms of TB with multi-organ involvement
 - c. Children and adolescents without bacteriological confirmation of TB or DST but with a high likelihood of MDR/RR-TB (based on clinical signs and symptoms of TB, in combination with a history of contact with a patient with confirmed MDR/RR-TB)



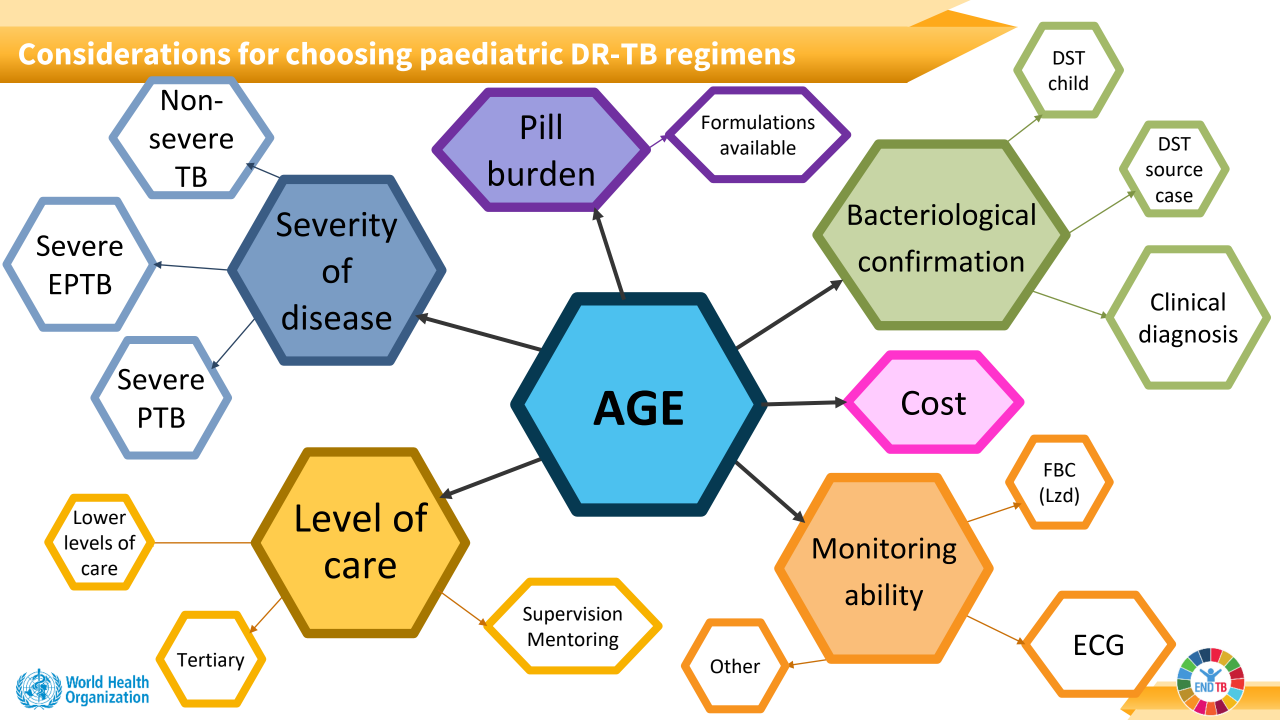


Updated mapping of DR-TB regimens – children & adolescents



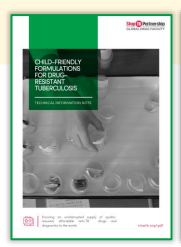






Child-friendly formulations: second-line medicines

- Child-friendly formulations of second-line medicines should be used whenever possible and included in funding requests
- New formulations available through GDF:
 - Bedaquiline 20 mg tab
 - Delamanid 25 mg disp tab
 - Linezolid 150 mg disp tab





WHO- RECOMMENDED GROUPING	MEDICINE	FORMULATION	PACK SIZE	SHELF-LIFE	STORE BELOW
	Levofloxacin 100mg	Dispersible tablet	100 in blister	36 months	30°C
А	Moxifloxacin 100mg	Dispersible tablet	100 in blister	24 or 36 months	30°C
	Bedaquiline 20mg	Tablet	60 in jar	36 months	30°C
	Linezolid 150mg	Dispersible tablet	100 in blister	24 months	30°C
В	Clofazimine 50mg	Tablet	100 in blister	36 months	30°C
	Cycloserine 125mg	Mini-Capsule	100 in blister	24 months	25°C
С	Ethambutol 100mg	Dispersible tablet	100 in blister	24 months	30°C
	Delamanid 25mg	Dispersible tablet	48 in blister	36 months	25°C
	Pyrazinamide 150mg	Dispersible tablet	100 in blister	36 months	30°C
	Ethionamide 125mg	Dispersible tablet	100 in blister	36 or 48 months	30°C
None	Isoniazid 100mg	Dispersible tablet	100 in blister	36 months	30°C

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Dosing guidance for second-line treatment

Annex to the Module 4 operational handbook: weight-based dosing of medicines used in MDR-TB regimens, adults and children

Group A medicines	Formulation (tablets, diluted in 10 mL of water, as applicable)	3-<5 kg	5-<7 kg	7-<10 kg	10-<16 kg	16-<24 kg	24-<30 kg	30-<36 kg	36-<46 kg	46-<56 kg	56-<70 kg	≥70 kg	Comments
Levofloxacin (Lfx)	100 mg dt (10 mg/mL)	5 mL (0.5 dt)	1	1.5	2	3	-	3 4					
	250 mg tab (25 mg/mL)	2 mL ^b	5 mL (0).5 tab) ^b	1	1.5	2			4			
	500 mg tab	_				1	1.5 2						
	750 mg tab	-					1 1.5						
Moxifloxacin (Mfx)	100 mg dt (10 mg/mL)	4 mL	8 mL	1.5	2	3	4	4	1		-		
	400 mg tab (40 mg/mL)	1 mL ^b	2 mL ^b	3 mL ^b	5 mL (0.5 tab) ^b	7.5 mL (0.75 tab) ^b	1	1					
	Standard dose												
	 400 mg tab high ose Dosing guidance available for children, adolescents and ad 				dults								

- 3 kg to >70 kg
- Age and weight-based approach for bedaquiline and delamanid
- Dosing provided using child-friendly formulations (preferred) but can also be given using adult formulations
- Final approach to dosing depending on formulations available in country



Dosing calculator in KSP app

← TUBERCULO	5: MANAGEMENT OF SIS IN CHILDREN AND DOLESCENTS
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AGE 1	
WEIGHT 9	
GROUP GROUP A	
MEDICINE BEDAQUILINE	
RESET	→ DOWNLOAD
Dosages for MDR-TI weighing 9 Kg, wit Bedaquiline	B patient aged 1 years, th drugs selected :
	Group A
DRUG : Bedaquili	ne
FORMULATION	DAILY DOSE
20 mg dt	4 od for 2 weeks; then 2 od M/W/F for 22 weeks
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Thank you for your attention!

vineyk@who.int



