F.10	Pyrazinamide 500mg tablet	
Does the application adequately address the issue of the public health need for the medicine?		☐ Yes ☐ No ☑ Not applicable Comments:
Briefly summarize the role of the proposed medicine(s) relative to other therapeutic agents currently included in the Model List, or available in the market.		This application proposes a formulation amendment to the core list in section 6.2.5 Antituberculosis medicines as per the latest edition of the core EML (21st edition). The medicine is already listed in the current WHO EML as a 400mg formulation. This application requests the addition to 500mg formulation.
Have all important studies and all relevant evidence been included in the application?		 ☐ Yes ☐ No ☒ Not applicable If no, please provide brief comments on any relevant studies or evidence that have not been included:
evidence of e	ication provide adequate fficacy/effectiveness of the the proposed indication?	☐ Yes ☐ No ☑ Not applicable Briefly summarize the reported benefits (e.g. hard clinical versus surrogate outcomes) and comment, where possible on the actual magnitude and clinical relevance of benefit associated with use of the medicine(s). The main reason for the addition of a 500mg pyrazinamide tablet is to reduce pill-burden, to the benefit of adherence to treatment. Is there evidence of efficacy in diverse settings (e.g. low-resource settings) and/or populations (e.g. children, the elderly, pregnant patients)? n/a
evidence of th	ication provide adequate ne safety and adverse ated with the medicine?	☐ Yes ☐ No ☐ Not applicable Comments:
	adverse effects of nat may require special	☐ Yes ☐ No ☐ Not applicable Comments:
the overall be	arize your assessment of nefit to risk ratio of the favourable, uncertain,	Not applicable

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Briefly summarize your assessment of the overall quality of the evidence for the medicine(s) (e.g. high, moderate, low etc.)	Not applicable
Are there any special requirements for the safe, effective and appropriate use of the medicine(s)? (e.g. laboratory diagnostic and/or monitoring tests, specialized training for health providers, etc)	☐ Yes ☑ No ☐ Not applicable Comments:
Are you aware of any issues regarding the registration of the medicine by national regulatory authorities? (e.g. accelerated approval, lack of regulatory approval, off-label indication)	☐ Yes ☑ No ☐ Not applicable Comments:
Is the proposed medicine recommended for use in a current WHO Guideline approved by the Guidelines Review Committee? (refer to: https://www.who.int/publications/who-guidelines)	 ✓ Yes ☐ No ☐ Not applicable Comments: Different formulations are already listed in the current WHO EML and WHO EMLc.
Briefly summarize your assessment of any issues regarding access, cost and affordability of the medicine in different settings.	Different formulations and strengths are available and affordable across different countries.
Any additional comments	None
Based on your assessment of the application, and any additional evidence / relevant information identified during the review process, briefly summarize your proposed recommendation to the Expert Committee, including the supporting rationale for your conclusions, and any doubts/concerns in relation to the listing proposal.	The reviewer recommends the inclusion of the pyrazinamide 500mg tablet formulation for the indication of TB on the WHO EML. Different countries have access to different formulations (in terms of registration, affordability, and supply) and adding options may increase-availability and the pool of suppliers.
References (if required)	