Last update 06/2024			Novelty					Development Stage							
AMR Accelerator Project	Asset Owner	Programme	New Class	New MoA	Mode of Action (MoA)	Description	Discovery (Pre)-H to Lead	lit Lead to d Candidate	Candidate to Phase I	Phase I	Phase 2a - alone or in combi- nation	Phase 2b - Dose ranging	Phase 2b - Regimen selection	Phase 2c - Duration rando- mization	
GNA NOW €31 m	NOSOPHARM	NOSO-502	1	1	Inhibition bacterial ribosome	NOSO-502 is the first clinical candidate in the novel antibiotic class called Odilhorhabdins, inhibiting the bacterial ribosome with a new mechanism of action.									
TRIC-TB €8 m	BioVersys and GSK	Alpibectir	1	1	Transcriptional modulator	Boosting Ethionamide efficacy and lowering the dose with small molecule transcriptional modulators to overcome multi-drug resistant tuberculosis infections and define a new place for Ethionamide in 1st-line tuberculosis treatments.					*				
AB-Direct €4 m		Gepotidacin tissue distribution	~	1	Topoisomerase type II inhibitor	Demonstrating penetration of gepotidacin in tonsillar and prostate tissues.									
ERA4TB €208 m	JANSSEN	ERA4TB-01	~	1	Cholesterol catabolism of mycobacteria	Molecule targeting cholesterol catabolism of mycobacteria.									
		ERA4TB-02	~	1	<i>Mycobacterium tuberculosis</i> tryptophan synthase	Compound targeting <i>Mycobacterium tuberculosis</i> tryptophan synthase, enzyme that catalyses the final two steps in the biosynthesis of tryptophan.									
		ERA4TB-03			Energy metabolism	Compounds targeting energy metabolism (electron chain transport).									
		ERA4TB-04	1	1	Lysine transfer RNA synthase	Compound targeting lysine transfer RNA synthase (Rv3598c), which is an essential gene as assessed by transposon mutagenesis.									
		ERA4TB-06	1	1	Mmpl3	Mycobacterial membrane protein Large 3 (Mmpl3) compounds with potent in vitro inhibitory and bactericidal activity against <i>Mycobacterium tuberculosis</i> .									
		ERA4TB-09	1	1	Not known	Natural product analogs active against <i>Mycobacterium tuberculosis</i> .									
		ERA4TB-10	1	1	DprE1	Piperazinobenzothiazinone derivative as anti-mycobacterial compound that targets and covalently inhibits the enzyme Decaprenyl-phosphoryl-ribose 2'-epimerase (DprE1).									
		ERA4TB-11	1	1	LeuRS	Small molecule oxaborole inhibitor of Mtb leucyl tRNA synthetase				Phase II of clinical development Phase I in ERA4TB					
		ERA4TB-13	1	1	Cholesterol catabolism of mycobacteria	Targets cholesterol catabolism of <i>Mycobacterium tuberculosis</i> (Mtb)									
		ERA4TB-14	~	1	Inhibits new target within a known pathway (Mtb energy metabolism) via a novel mode of inhibition	Small molecule compound that inhibits the mycobacterial cytochrome bc1 complex in the cellular respiration pathway, leading to the depletion of ATP, in three mycobacterial species, <i>M. tuberculosis</i> , <i>M. leprae</i> , and <i>M. ulcerans</i>									
		ERA4TB-15	~	1	Inhibits new target within a known pathway (Mtb mycolic acid biosynthesis) via a novel mode of inhibition Inhibits new target within a known	Novel class of small-molecule antibiotics shown to covalently inhibit the acyl transferase domain of Mtb Pks13, a polyketide synthase involved in the mycolic acid biosynthetic pathway Novel class of small-molecule antibiotics shown to covalently inhibit the									
		ERA4TB-16 ERA4TB-17	1	1	Inhibits a new target within a known Inhibits a new target within a known	acyl transferase domain of Mtb Pks13, a polyketide synthase involved in the mycolic acid biosynthetic pathway Novel class of small-molecule antibiotics that inhibits FadD32, a key enzyme									
		ERA4TB-17	√	1	H3D-012895 inhibits a new target within	at the interface between the fatty acid synthase and polyketide synthase biosynthetic pathways and involved in the synthesis of mycolic acid Novel class of small-molecule antibiotics that inhibits FadD32, a key enzyme									
		BC1 back up	~	1	a known pathway (Mtb mycolic acid biosynthesis) BC1	at the interface between the fatty acid synthase and polyketide synthase biosynthetic pathways and involved in the synthesis of mycolic acid Lead optimization program on BC1 inhibitors									
€9 m RespiriNTM €8 m	TBA	MenG	1	1	MenG	H2L medChem for novel menG inhibitors									
		BDQ LAI	√	1	ΑΤΡαse	Novel long acting injectable formulation of bedaquiline for Tb preventive									
		PASA	√		DHFR	therapy Novel PAS analogues									
		HDT	 Image: A start of the start of		Various	Exploring of known host directed therapies for TB treatment									
		Mtr		1	Mtr	Mtr target exploration.									
					not known	Progress novel assets (one FIH start) for Non-Tubercular Mycobacterium (NTM) that may act synergistically with Bedaquiline						 			
€8 m UNITE4TB €185 m	GSK	GSK656			LeuRS	A first-in-class investigational antitubercular agent which is being developed for the treatment of tuberculosis as part of a future combination regimen.						 			
0105111	Leibniz-HKI/ LMU	BTZ-043		1	Cell wall synthesis	New MoA/not regulatory approved product with this MoA. Suppresses protein synthesis in <i>Mycobacterium tuberculosis</i> (Mtb) by inhibiting the enzyme leucyl t-RNA synthetase (LeuRS). A first-in-class investigational antitubercular agent which is being developed for the treatment of tuberculosis as part of a future combination regimen.									
			1			New MoA/not regulatory approved product with this MoA. BTZ-043 inhibits an enzyme (BTZ-043) with is essential for cell wall synthesis in <i>Mycobacterium tuberculosis.</i>									

Accelerating scientific discoveries in the antimicrobial resistance (AMR) field



