

1. MAKING THE EU A BEST PRACTICE REGION

1.1 BETTER EVIDENCE AND AWARENESS OF THE CHALLENGES OF AMR

CONCRETE ACTIVITIES	NCRETE ACTIVITIES TIMELINES AND DELIVERABLES						
 Strengthen One Health surveillance and reporting of AMR and antimicrobial use 	2017	2018	2019	2020	2021	2022	
Review EU implementing legislation on monitoring AMR in zoonotic and commensal bacteria in farm animals and food.	Mandate to EFSA for technical advice		EFSA opinion	Adoption of the new legislation			
Review EU implementing legislation on reporting communicable diseases in humans.		New implementing act with updated AMR case definitions					
Identify and assess under the Animal Health Law, resistant bacteria that cause transmissible animal diseases and, if necessary, develop harmonised rules for their surveillance.			Mandate to EFSA for technical advice once list of transmissible disease is adopted under Animal Health Law				
Improve AMR detection in the human health sector by providing EU support for networking collaboration and reference laboratory activities.			collabo	nme funding to suppor ration and reference lat activities in human healt	oratory		
Consider options for the harmonised monitoring of AMR in the environment.		Linked to the EU strate	egic approach to pharm	naceuticals in the enviro	nment (see action 1.4)		
 Benefit from the best evidence-based analysis and data 	2017	2018	2019	2020	2021	2022	
Provide evidence-based data on possible links between the consumption of antimicrobial agents and the occurrence of antimicrobial resistance in humans and food-producing animals.	2 nd JIACRA report	Mandate to EFSA/ ECDC/EMA for a 3rd JIACRA report			3rd JIACRA report		
Define a limited number of key outcome indicators for AMR and antimicrobial consumption.	EFSA/ECDC/EMA report on indicators						



CONCRETE ACTIVITIES

TIMELINES AND DELIVERABLES



1.2 BETTER COORDINATION AND IMPLEMENTATION OF EU RULES TO TACKLE AMR

 Improve the coordination of Member States' One Health responses to AMR 	2017	2018	2019	2020	2021	2022
Make available regular information on AMR in the context of the AMR One Health network, which gives an overview of the AMR epidemiological situation at Member State and EU level.		2nd meeting 3rd meeting	4th meeting 5th meeting		6th meeting	7th meeting*
Support the implementation of national One Health action plans through joint Commission and the ECDC visits to Member States upon request.	Joint One Health visits with ECDC	Joint One Health visits with ECDC	Joint One Health visits with ECDC			Joint One Health visits with ECDC*
Launch a joint action to tackle AMR and healthcare-associated infections to support collaborative activities and policy development by Member States.	Launch	Interim	reports		Final conference and report	



CONCRETE ACTIVITIES TIMELINES AND DELIVERABLES 2017 2018 2019 2020 2021 2022 Make increased use of the EU Health Security Committee and the Commission Working Group on AMR in the veterinary and food Regular exchanges on AMR in HSC** areas to strengthen coordination and to share information. Meeting of working Meetings of group on AMR working group on in food AMR in food Seek to co-fund and collaborate with the WHO on activities to help EU Member States develop and implement national One New WHO EURO activities co-funded by EU health programme Health action plans against AMR. Better implementation of EU rules ٠ Interim Overview Assess the effectiveness of the implementation of EU legislation Overview report report on, inter alia, monitoring AMR in food-producing animal populations and food by continuing to carry out regular audits in Member States. Audits Audits Develop training programmes on AMR for Member State competent authorities under the Better Training for Safer Food BTSF training activities (BTSF) initiative and for health professionals. SRSS support Advise Member States on the possibility to use the Structural presented to MSs in Reform Support Service (SRSS) funding to Member States for the AMR One Health designing and implementing policies against AMR. network



1.3 BETTER PREVENTION AND CONTROL OF AMR

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES	
 Strengthen infection prevention and control measures 	2017 2018 2019 2020 2021	2022
Support good practices in infection prevention and control in hospital environments.	Work by OECD supported by EU health programme to report on cost effectiveness of AMR/infection prevention measures including hand hygiene and hospital cleaning	
Support activities for infection prevention and control in vulnerable groups, in particular to tackle resistant tuberculosis strains.	Report from HA-REACT joint action from EU Health programme on HIV and co- infection prevention and harm reduction which includes activities to address tuberculosisReport from INTEGRATE joint action from EU health programm on integrating prevention, testin and link to care strategies across HIV, viral hepatitis TB and STIs in Europe*)
Promote the uptake of vaccination in humans to prevent infections and subsequent use of antimicrobials.	Part of Joint Action Vaccination* Part of policy initiative on vaccination (Council Recommendation on strengthened cooperati against vaccine preventable diseases)*	on



CONCRETE ACTIVITIES

TIMELINES AND DELIVERABLES

		2018 2019 2020 2021 Improve the microbiota of animals through autorisation of several feed additives H2020 research projects launched: DISARM, ROADMAP and HealthyLivestock USARM, ROADMAP and HealthyLivestock Implementation of existing policy on biosecurity AGRI action Regulations adopted (as of 2022) Drafting of the acts and consultation with MS and stakeholders*				
	2017	2018	2019	2020	2021	2022
Continue to promote animal husbandry, including aquaculture and livestock farming systems, and feeding regimes which support good animal health and welfare to reduce antimicrobial consumption.	AMR related workshop		projects launched: DISARM, ROADMAP and			
		Implen	nentation of existing po	licy on biosecurity AGR	l action	
 Promote the prudent use of antimicrobials 	2017	2018	2019	2020	2021	2022
Forthcoming veterinary medicinal products and medicated feed Regulations containing concrete restrictions for the prophylactic and metaphylactic use of antimicrobials. Moreover, work towards EU implementing and delegated acts established in these Regulations include a list of antimicrobials reserved for human use, drawing up a list of antimicrobials that cannot be used		adopted		and consultation with MS and	and consultation with MS and	
under 'cascade use', limits for residues of antimicrobials in feed, requirements for animals or products of animal origin exported from third countries and methods for data gathering and reporting on the sales and use of antimicrobials.		be adopted as per deadlines in those Regulations		stakeholders	implementing and delegated acts adopted and published	
Develop EU guidelines for the prudent use of antimicrobials in human medicine.	EU guidelines published					
Assist Member States implement EU guidelines for the prudent use of antimicrobials in veterinary medicine.	Fact-finding missions on prudent use in veterinary sector	Fact-finding missions on prudent use in veterinary sector	Overview report			
	Interim Overview report					



1.4 BETTER ADDRESSING THE ROLE OF THE ENVIRONMENT

CONCRETE ACTIVITIES	TIMELINES	AND DELIVER	ABLES			
	2017	2018	2019	2020	2021	2022
Encourage the EMA to review all available information on the benefits and risks of older antimicrobial agent	EMA finalised	Exploratory meeting		EMA review of medicines containing fosfomycin		
	a review of the vancomycin- containing medicines	with EMA on the topic of availability of veterinary antimicrobials and their prudent use		Regular update meetings with EMA on the topic of availability of veterinary antimicrobials and their prudent use		
Adopt an EU strategic approach to pharmaceuticals in the environment.			EU strategic approach to pharmaceuticals in the environment (PIE) adopted		EU strategic approach in the environment (PIE	
Maximise the use of data from existing monitoring to improve knowledge, including by using the Information Platform for Chemical Monitoring (IPCheM).					PCHeM in regulatory esses*	
Reinforce the role of the Scientific Committee on Health, Environmental and Emerging Risks (SCHEER) in providing scientific advice to the European Commission on environment-related AMR issues.		Video material describing the role of SCHEER circulated to relevant sectors in the Commission				



1.5 A STRONGER PARTNERSHIP AGAINST AMR AND BETTER AVAILABILITY OF ANTIMICROBIALS

CONCRETE ACTIVITIES	TIMELINES A	ND DELIVER	ABLES							
	AMR discussed in the plenary of the advisory group of	2018	2019	2020	2021	2022				
Engage with and support collaboration among key stakeholders in the human health, animal health, food, water and environmental	the food chain and animal and plant health EMA information session on AMR									
sectors to encourage the responsible use of antimicrobials and appropriate handling of waste material.	(EMA Working Party with Patients' and Consumers' Organisations (PCWP) and Healthcare Professionals'									
	Organisations (HCPWP))		BTSF training activities		1 1 1 1					



CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES							
	2017	2018	2019	2020	2021	2022		
Work with stakeholders to ensure the availability of human and veterinary antimicrobials and continued access to established products; provide incentives to increase the uptake of diagnostics, antimicrobial alternatives and vaccines.			Discussion in the IVD Technical Group (IVD TG) and possibly in the Medical Device Coordination Group on how to promote the uptake of diagnostics		١	Multi-stakeholder conference on the themes of availability of veterinary antimicrobials (old, new and future) and on their prudent use, at EU level but also in the context of broader international cooperation*		
		Implementation of	FRegulation (EU) 2017/	746 on in vitro diagnos	tic medical devices			
Reduce the scope for falsified medicines by assisting Member States and stakeholders in the successful implementation of the safety features (unique identifier).	Presidency Safer Europe Without Falsified Medicines Conference organised by industry and the Estonian Presidency							



CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES						
	2017	2018	2019	2020	2021	2022	
Reduce the scope for falsified medicines by assisting Member States and stakeholders in the successful implementation of the safety features (unique identifier).		Conference organised by DIA (Drug information Association) with pharmaceutical industry	Regular Commissior	i expert group on the in safety features*	nplementation of the		
			Update of the Q&A in on the safe	terpretative document ty features*			
Discuss the availability of veterinary antimicrobials to tackle AMR in the Veterinary Pharmaceutical Committee.			Exploratory meeting (see above mentioned multistakeholders conference)				

2. BOOSTING RESEARCH, DEVELOPMENT AND INNOVATION ON AMR

2.1 IMPROVE KNOWLEDGE ON DETECTION, EFFECTIVE INFECTION CONTROL AND SURVEILLANCE

CONCRETE ACTIVITIES	TIMELINES	AND DELIVER	ABLES			
	2017	2018	2019	2020	2021	2022
	Funding of research projects under the ERA- NET SusAn on animal production	Launch of One Health EJP which includes research on new intervention tools on AMR	Launch of Call topic on clinical management of AMR			
Support research into the development and assessment of interventions that prevent the development and spread of AMR.		Funding of projects coming out of Call topic SFS-46-2017 on alternative production systems to address anti- microbial drug usage, animal welfare and the impact on health				
	Funding decision on projects from the JPIAMR 5th call on prevention and intervention strategies to control AMR infections					
Support research into understanding the epidemiology of AMR, in particular the pathways of transmission between animals and humans, and their impact.	Funding decision on project addressing the clinical burden of Clostridium difficile infection under under IMI2, Call 9				Report of projects selected under 3rd JPIAMR ERA- NET Co-fund call to bridge the knowledge gap on AMR transmission mechanisms*	



CONCRETE ACTIVITIES	TIMELINES	AND DELIVER	ABLES			
	2017	2018	2019	2020	2021	2022
Support research into the development of new tools for early (real-time) detection of resistant pathogens in humans		Launch of One Health EJP which includes research on early signalling and assessing zoonotic threats		Reporting COMPARE, on the rapid identification, containment and mitigation of emerging infectious diseases and foodborne outbreaks*		
Support research into the development of new tools for early (real-time) detection of resistant pathogens in humans and animals.		Launch of Call topic SC1-BHC-13-2019, on mining of big data for early detection of infectious disease threats	Reporting DIAGORAS, on bedside diagnosis of oral and respiratory tract infections, and identification of antibiotic resistances for personalised monitoring and treatment			
Support research into new eHealth solutions to improve prescription practices, self-management of health, care solutions, and improve awareness of AMR.		Launch of Call topic SC1-DTH-10-2019- 2020, on digital solutions for health and care services				



2.2 DEVELOP NEW THERAPEUTICS AND ALTERNATIVES

CONCRETE ACTIVITIES	TIMELINES	AND DELIVER	ABLES	,		
	2017	2018	2019	2020	2021	2022
Support research into the development of new antimicrobials and alternative products for humans and animals as well as the	Funding decision on EDCTP2 projects on treatments for cryptococcal meningitis, malaria, HIV, tuberculosis and reproductive tract infections	Reporting of project PneumoNP on nanotherapeutics to treat antibiotic resistant Gram- Negative pneumonia	Launch of IMI2 call topics on development of new antimicrobials and alternative products for humans		IDFF to development of d alternative products*	
		Reporting of project NAREB on nanotherapeutics for antibiotic resistant emerging bacterial pathogens	: Reports of projects from 1st and 2nd join transnational calls of JPIAMR		Reporting of project anTBiotic progressing TB drug candidates to clinical proof of concept*	
Support research into the development of new antimicrobials and alternative products for humans and animals as well as the repurposing of old antimicrobials or the development of new combination therapies.		Launch of Call topics SC1-BHC-14-2019 on stratified host-directed approaches to improve prevention, treatment and/or cure of infectious diseases and SC2- SFS-11-2019- 2019 on antimicrobials and animal production		Reporting of IMI ND4BB projects on development of new antimicrobials and alternative products for humans*		
		Reporting of project FormAMP on innovative nanoformulation design of antimicrobial peptides				



CONCRETE ACTIVITIES	TIMELINES	AND DELIVER	ABLES					
	2017	2018	2019	2020	2021	2022		
Support SMEs in their R&D efforts towards innovative and/ or alternative therapeutic approaches for the treatment or prevention of bacterial infections.				on development of ne	from SME Instrument, ew antimicrobials and e products*			
Facilitate sharing of antimicrobial research data among relevant stakeholders.		Launch of IMI2 call topics on development of new antimicrobials including facilitating data sharing Reporting of IMI ND4BB project TRANSLOCATION on antibacterial drug discovery						
Support the establishment of a European-wide sustainable clinical research network.	Launch of Call topic SC1-HCO-08-2018 on the creation of a European wide sustainable clinical research network for infectious diseases				Reporting of IMI ND4BB project COMBACTE on the development of new antibacterial treatments*			
Support research and innovation to promote the use of digital technologies supporting the development of new therapeutics and alternatives.				Reporting of Biotechnology (LEIT) project DD-DeCaF on optimisation of -omics data				



2.3 DEVELOP NEW PREVENTIVE VACCINES

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES						
	2017	2018	2019	2020	2021	2022	
					Reporting of projects from Call topic SC1- PM-16- 2017 on in-silico trials for developing and assessing biomedical products*		
	Launch of Call Topic SC1-BHC-15-2018, on new anti- infective agents	Launch of Call Topics SC1- BHC-14-2019, on stratified host-directed approaches to improve prevention, treatment and/or cure of infectious diseases, SC2- SFS-31-2019 (ERA-			Reporting of the European AIDS Vaccine Initiative (EAVI) 2020*	Reporting of projects OptiMalVax and MultiViVax on Malaria vaccine development*	
Continue to support research into the development of new	(including vaccines) for prevention and/ or treatment of neglected infectious diseases			Reporting of projects TBVAC2020 and EMI- TB on advancing tuberculosis vaccine candidates	Reporting of project TracVac on Developing a Chlamydia Trachomatis vaccine*		
effective preventive vaccines for humans and animals.		NET) on veterinary vaccinology and SC2-SFS-12-2019 on swine fever vaccines	Reporting of project SAPHIR on novel vaccine strategies for animal production	Loans awarded under of new v	DFF to development vaccines*		
				Reporting MycoSynVac, on development of a Mycoplasma vaccine for animal use			



2.4 DEVELOP NOVEL DIAGNOSTICS

CONCRETE ACTIVITIES	TIMELINES							
	2017	2018	2019	2020	2021	2022		
Support increasing the knowledge base concerning the barriers that influence the wider use of vaccination in medical and veterinary practice.		Launch of Call Topic SFS-11-2018-2019, on antimicrobials and livestock production						
	Launch of IMI2 call, on diagnostics development and validation	Reporting of project VIROGENESIS on virus discovery and epidemic tracing	Reporting of poject Poc-ID on Point-of- Care diagnostics for Infectious Diseases		Reporting of project FAPIC on developping a fast assay for pathogen identification and characterisation			
Support research into the development of new diagnostic tools in particular on-site tests in humans and animals.		Launch of One Health EJP, incl. research on new diagnostic tools on AMR		Loans awarded under IDFF to development of new diagnostic tools*				
					Reporting of EDCTP2 projects, on diagnostic tools for poverty-related diseases*			
					Reporting PREPARE, on outbreak preparedness			
Support the use of IT solutions in developing tools for diagnosing human and animal infections.				Reporting of projects on IT solutions ar diagnosing human a				



2.5 DEVELOP NEW ECONOMIC MODELS AND INCENTIVES

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES					
	2017	2018	2019	2020	2021	2022
Encourage the uptake of diagnostics in medical and veterinary practice.	Launch of Call Topic SC1-HCO-12-2018, on Innovation Procurement					
Increase the evidence base for understanding the societal costs and benefits of different strategies for fighting AMR.		Reporting of IMI DRIVE-AB project				
Support research into the development of new economic models, exploring and analysing incentives to boost the development of		Reporting of IMI DRIVE-AB project				
new therapeutics, alternatives, vaccines and diagnostics.		TATFAR discussion on incentives				
Analyse EU regulatory tools and incentives – in particular orphan and paediatric legislation – to use them for novel antimicrobials and innovative alternative medicinal products that currently do not generate sufficient returns on investment.	Report from the European Commission to the European Parliament and the Council on the 10 years of the paediatric Regulation	Study on the economic impact of the supplementary protection certificate, pharmaceutical incentives and rewards in Europe	Study on orphan medicinal products legislation	Evaluation of the orphan and paediatric medicines legislation*		
Encourage Member States to explore results and recommendations of EU research projects on new economic business models.			Discussion with Member States in the Pharmacuetical Committee			
Develop new or improved methodological HTA approaches and foster methodological consensus-building.	Launch of Call Topic SC1-BHC-26-2018, on HTA research to support evidence- based healthcare					



2.6 CLOSE KNOWLEDGE GAPS ON AMR IN THE ENVIRONMENT AND ON HOW TO PREVENT TRANSMISSION

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES						
	2017	2018	2019	2020	2021	2022	
Support research into knowledge gaps on the release of resistant microorganisms and antimicrobials into the environment and their spread.		Launch of One Health EJP, incl. research on resistance spread	Reporting of project EFFORT on microbial drug resistance and transmission		Reports of projects from JPIAMR 3rd ERA- NET Co-fund call on transmission dynamics*		
Explore risk assessment methodologies to evaluate the risks to human and animal health from the presence of antimicrobials in the environment.		Launch of One Health EJP, incl. research on resistance spread			n of Commission strated euticals in the environm		
Support research into and the development of new tools for monitoring antimicrobials and microorganisms resistant against antimicrobials in the environment.				•	n of Commission strated euticals in the environm		
Support the development of technologies that enable efficient and rapid degradation of antimicrobials in wastewater and the environment and reduce the spread of AMR.			ERA-NET for JPI Water and JPIAMR				

European Commission 18

3. SHAPING THE GLOBAL AGENDA

3.1 A STRONGER EU GLOBAL PRESENCE

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES							
Continue to actively contribute to the normative work of the WHO, the OIE, the FAO, and the Codex Alimentarius on the development of ambitious international frameworks and standards/norms/ guidelines/methodologies related to AMR.	2017	2018	2019	2020	2021	2022		
	Signature of a letter of intent for reinforced cooperation between FAO and EC between FAO DG and V. Andriukaitis Commission	Involvement of FAO	in new BTSF activities	New CODEX guidelines on integrated monitoring and surveillance of foodborne AMR*				
		Review upcoming normative work (e.g. as outlined in resolution of the World Health Assembly) and assess EC involvement			Revised CODEX code of practice to minimize AMR*			
Reinforce technical cooperation with the WHO and its members in key areas of the WHO Global Action Plan on AMR.								



CONCRETE ACTIVITIES

TIMELINES AND DELIVERABLES

CONCRETE ACTIVITIES							
	2017	2018	2019	2020	2021	2022	
Boost support for the International Conference on the Harmonisation of Technical Requirements for the Registration of Pharmaceuticals for Human Use (ICH) and the Veterinary International Conference on the Harmonisation (VICH) on relevant international guidelines/standards/norms related to AMR.	Promotion of the new EU Action Plan against AMR during the IPRF meeting in the margins of the ICH meeting in Geneva in November 2017		Text on AMR in the UN political declaration on Universal Health Coverage	VICH guidelines an	to update existing d other possibilities armonisation*		
Work towards continued high-level political attention and commitment to AMR action, including in the United Nations forums, the G7 and the G20.	AMR featuring in the G7 Health Minister Communique and G20 Health Ministers' Declaration	AMR on G7 and G20 Agenda	AMR on G7 and G20 Agenda	AMR on G20 Agenda	AMR on G7 Agenda		
	Approval by G7 CVOs of a common approach on the definitions of therapeutic, responsible and prudent use of antimicrobials in animals						
Look for synergies with the UN Strategic Approach to International Chemicals Management's work on the emerging policy issue of pharmaceuticals in the environment.				EU Chemical Startegy for Sustainability		n of EU Chemical Sustainability*	
Analyse the feasibility of setting up a global AMR clinical studies network in collaboration with G7 members.							



CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES						
	2017	2018	2019	2020	2021	2022	
Continue and strengthen ongoing collaboration within the Transatlantic Taskforce on Antimicrobial Resistance (TATFAR), which includes the EU, the USA, Canada, and Norway.				Develop actions and deliverables for new TATFAR work plan 2021-2025*			
		Updated TATFAR work plan and deliverables				of the new TATFAR «plan*	
						TATFAR Conference*	
Promote international regulatory convergence between the EMA and other regulatory agencies such as the US Food and Drug Administration (FDA) and the Japan Pharmaceuticals and Medical Devices Agency (PMDA) on development plans for new promising antimicrobials.			4th tripartite meeting between PMDA, EMA, and FDA to discuss convergence on approaches for the evaluation of antibacterial drugs*				



3.2 STRONGER BILATERAL PARTNERSHIPS FOR STRONGER COOPERATION





IMPROVE KNOWLEDGE ON DETECTION, EFFECTIVE INFECTION CONTROL AND SURVEILLANCE

CONCRETE ACTIVITIES	TIMELINES	TIMELINES AND DELIVERABLES						
	2017	2018	2019	2020	2021	2022		
Engage with major global players and strategic countries (e.g. Brazil, China, India), contributing towards achieving objectives of the WHO global action plan on AMR. F2F outreach including AMR*	Seminars on AMR with Argentina, Brazil, Chile & Colombia	Seminars on AMR with Paraguay, Peru & Uruguay	AMR activities with China and India*	F2F outreach	including AMR*			
	Seminar with India on the use of veterinary medicines and AMR	Overview report regarding national policies and measures against AMR in third countries	Follow-up seminars in South East Asia including Australia and New Zealand*	AMR activities in South East Asia and South America*				
	Identification mission on AMR cooperation with South-American partner countries		AMR activities in Latin American countries under FPI project					
Support EU candidate countries, potential candidate countries and neighbouring countries to which the ENP applies in the alignment with, and capacity building for the implementation of EU legislation related to AMR and EU standards.			ECDC/EFSA and EU- Enlargement multi- country workshop / ECDC/EFSA and EU- ENP multi-country workshop					
Invite the European Parliament, Member States and stakeholders to share views on actions to be taken to ensure that efforts to combat AMR made by EU producers, including farmers, do not place them at a competitive disadvantage.								

3.3 COOPERATING WITH DEVELOPING COUNTRIES

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES								
	2017	2018	2019	2020	2021	2022			
Continue to contribute to reducing AMR in least developed countries through infectious disease programmes such as the Global Alliance for Vaccines and Immunisations (GAVI)			Implement	ation through Commiss	ion pledges*				
Assist in the development of AMR strategies in the areas of food safety and animal health through regional training workshops on AMR	New BTSF activities for non-EU countries								
Support partner countries' policy initiatives on AMR, where appropriate, through international cooperation and development instruments (e.g. Global Public Goods & Challenges, the European Development Fund).	undertaking surve	Through a pilot project on mapping the global threat of AMR, DEVCO supports WHO to develop a point prevalence protocol for undertaking surveys on prescribing and use of antimicrobial medicines in hospital setting, build capacity to implement antimicrobial stewardship programmes in hospitals in SSA, address the prevention, detection and response to substandard and falsified products							
Support the development of resilient health systems in partner countries			Supporting WHC) for strengthened healt instruments at coun	h systems through de htry and region levels	velopment funding			

3.4 DEVELOPING A GLOBAL RESEARCH AGENDA

CONCRETE ACTIVITIES	TIMELINES AND DELIVERABLES							
	2017	2018	2019	2020	2021	2022		
Improve global coordination of research activities.		Launch of G20 Research hub						
Support the establishment of a virtual research institute under JPIAMR.					Reporting of project EXEDRA on the expansion of the JPIAMR*			
			Reporting of EDCTP2 projects on research capacity development in support of the EVD response					
Continue collaborative research with Sub-Saharan Africa in the context of the European and Developing Countries Clinical Trial Partnership (EDCTP) in particular in relation to tuberculosis, HIV/ AIDS, malaria and neglected infectious diseases.			Reporting of EDCTP2 projects on maximising the impact of EDCTP research and translation of research results into policy and practice					
Foster international research collaboration on AMR in the animal health sector in the STAR-IDAZ International Research Consortium.								