



ID-EPTRI a new research infrastructure that will facilitate the future development of better paediatric medicines

Roma, January 2018 - **ID-EPTRI** (European Paediatric Translational Research Infrastructure) is a project coordinated by the Consorzio per Valutazioni Biologiche e Farmacologiche (CVBF-TEDDY) that granted with 3 million of Euros from funding of the European Commission. The main objective of the project lasting 24 months, is to **design the framework for a new European Paediatric Research Infrastructure (RI)**.

Minors represent 20% of the European population and their care is one of the most important priorities and challenges for Europe. It's essential the development of evidence-based paediatric medicines and treatment strategies. Nowadays around the 50% of the medicines addressed to children and young patients have not been tested specifically for them. For this reason, it's strategic the development of the suitable research infrastructure that can solve this problem, studying the paediatric research from the early phases to the paediatric formulation. Children and young patients cannot be never compared with adults as they are growing up and their metabolism is different. For this reason, the only way to develop better medicines for children and young patients is studying them specifically for this type of "special" population. A dedicated infrastructure integrating the different basic research networks addressed to paediatric population will help in the process to reduce time and increase the number of projects. On the other hand, it also can help to a fast translation into the clinical practice.

This new research infrastructure, EPTRI is complementary to the other existing Biomed Research Infrastructures acting as a 'Paediatric Common Service' in the [ESFRI](#) (European Strategy Forum on Research Infrastructures) Scenario. The project involves 26 partners (listed in the **Appendix 1**) from EU and non-EU countries including consolidated research infrastructures, top-level universities, scientific and clinical centers of excellence in Europe and aims to create a Conceptual Design Report (CDR) to set-up the European Paediatric RI.



In order to set up the new RI within the European landscape, there will be three different phases:

- 1) a context analysis, aiming to acquire the information needed to complete a consistent CDR;
- 2) an operational phase focused more specifically on the design of the whole RI;
- 3) a feasibility phase in which selected pilot experiences will allow testing a limited number of services and tools delivered by EPTRI.

The involvement of children and young people is included in the development of the project, with the aim to ensure that their needs are addressed in the conceptual design report for the development of the new infrastructure, that will join and cover all the different domains previous to the clinical research in drug development: 1. Pediatric Medicines Discovery and Preclinical Studies; 2. Biomarkers; 3. Paediatric Pharmacology, 4. Formulation Science, 5. Underpinning Paediatric Studies.

EPTRI will allow to

- Cover the current existing gaps connecting the different steps in research from early stage.
- Enable and prepare researchers in many methodological areas to conduct research that effectively underpins the development of paediatric medicines.
- Increase the global competitiveness of the European RI also in favor of children, young people and their families.

To test this new RI, in the second year a feasibility phase is proposed to develop virtual exercises simulating the operation of the RI. In this phase four types of experiments will be developed: 1. Feasibility studies with scientists; 2. Feasibility studies with governments; 3. To test the interest of patients' associations and YPAGs and 4. Commons services with RI feasibility studies.

The Kick-off Meeting of the project will be held in Rome next 15th and 16th of January, with the involvement of all the partners. The agenda of the event is included in the Appendix 2. All the means of communication are welcome to attend to the meeting. In case of interest to plan a meeting with the coordinators of the project, please contact in advance with us. The venue is going to be:

Ministry of Education, Universities and Research
Sala della Comunicazione
Viale Trastevere, 76/a
Rome, Italy

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About Consorzio per Valutazioni Biologiche e Farmacologiche (CVBF)

Consorzio per Valutazioni Biologiche e Farmacologiche (CVBF) is a not-for-profit organisation, founded in 2000 with the mission to perform research and provide scientific, economic and regulatory consultancy for innovation in the health sector at European level. The main fields of interest are life sciences and biotechnologies, drug development for small populations (pediatric and rare diseases), research management and methodology, monitoring, statistics, regulatory, ethics and pharmacovigilance.

Learn more: <https://www.cvbf.net>



Appendix 1.**List of participants**

Participant No	Participant organisation name	Acronym	Country
1 Coordinator	CONSORZIO PER VALUTAZIONI BIOLOGICHE E FARMACOLOGICHE	CVBF	Italy
2	FONDAZIONE PENTA-FOR THE TREATMENT AND CARE OF CHILDREN WITH HIV-ONLUS	PENTA	Italy
3	EUROPEAN INFRASTRUCTURE FOR TRANSLATIONAL MEDICINE	EATRIS-ERIC	Netherlands
4	UNIVERSITY COLLEGE LONDON	UCL	UK
5	ATHINA-EREVNITIKO KENTRO KAINOTOMIAS STIS TECHNOLOGIES TIS PLIFOFORIAS, TON EPIKOLNONION KAI TIS GNOSIS	ATHENA	Greece
6	BIOBANKS AND BIOMOLECULAR RESOURCES RESEARCH INFRASTRUCTURE CONSORTIUMN (BBMRI-ERIC)	BBMRI-ERIC	Austria
7	THE CYPRUS FOUNDATION FOR MUSCULAR DYSTROPHY RESEARCH	CING	Cyprus
8	ASSISTANCE PUBLIQUE - HOPITAUX DE PARIS	AP-HP	France
9	STICHTING KATHOLIEKE UNIVERSITEIT	RUMC	Netherlands
10	THE UNIVERSITY OF LIVERPOOL	ULIV	UK
11	ROMANIAN ANGEL APPEAL	RAA	Romania
12	INSTYTUT POMNIK CENTRUM ZDROWIA DZIECKA	IPCZD	Poland
13	OSPEDALE PEDIATRICO BAMBINO GESU	OPBG	Italy
14	FYZIOLOGICKY USTAV AKADEMIE VED CESKE REPUBLIKY VEREJNA VYZKUMNA INSTITUCE (VVI)	IPHYS	Czech Republic
15	FUNDACIO SANT JOAN DE DEU	FSJD	Spain



16	NIZHEGORODSKIY GOSUDARSTVENNIY UNIVERSITET IM N.I. LOBACHEVSKOGO	UNN	Russia
17	SERVICIO MADRILENO DE SALUD	SERMAS-HULP	Spain
18	ECRIN EUROPEAN CLINICAL RESEARCH INFRASTRUCTURE NETWORK	ECRIN	France
19	QENDRA SPITALORE UNIVERSITARE NENE TEREZA TIRANE	UHCT	Albania
20	TECHNION - ISRAEL INSTITUTE OF TECHNOLOGY	TECHNION	Israel
21	TARTU ULIKOOL	UTARTU	Estonia
22	UNIVERSITATSKLINIKUM ERLANGEN	UKER	Germany
23	SWISS CLINICAL TRIAL ORGANISATION VEREIN	SCTO	Switzerland
24	STATE "INSTITUTE OF PEDIATRICS OBSTETRICS AND GYNECOLOGY NATIONAL ACADEMY OF MEDICAL SCIENCES OF UKRAINE"	UKR	Ukraine
25	VASTRA GOTALAND LANS LANDSTING	VGR	Sweden
26	UNIVERSITAIR MEDISCH CENTRUM UTRECHT	UMCU	Netherlands



Appendix 2. Agenda Kick off meeting

Day 1, January 15th, 2018

08.30	REGISTRATION	
08:45	OPENING SESSION	
	Welcome address from the supporting Italian Institutions	
	Ministry of Education, University and Research (MIUR)	L. Nicolais
	Italian Medicines Agency (AIFA)	S. Vella
	Apulia Region – Director of University Paediatric Hospital	A. Del Vecchio
	University of Bari “Aldo Moro”	L. Margari
	Regional Agency for Technology and Innovation (ARTI)	A. Monterisi
	Opening remarks	D. Bonifazi
09:15	SESSION 1 - The role of RIs to strengthening research outcomes and to improve patients' health	
	Chair: A. Ceci, I. Lutsar	
	How to implement a new RI: lessons from the Executive Master in Management of Research Infrastructures	M. Lavitrano
	BBMRI ERIC: Biobanking and BioMolecular resources Research Infrastructure	E. Steinfelder
	EATRIS ERIC: European infrastructure for translational medicine	G. Migliaccio
	ECRIN ERIC: European clinical research infrastructure network	J. Demotes
	The role of ERNs in the European scenario and interactions with RIs	L. Sangiorgi
11.00	COFFEE BREAK	
11.30	SESSION 2 - The EPTRI project and the involved Partners	
	Chairs: P. Macheras, C. Altomare	
	ID-EPTRI: overview of the project (planned activities, expected results and impact, timelines)	D. Bonifazi



	<p>Introduction round and presentation of Project Partners and involved staff (5 min. for each representative of the EPTRI General Assembly member)</p> <ul style="list-style-type: none"> • Introduction of each Beneficiary and team members • Information on the organisation • Competences related to the project • Expectations of the project 	All the Partners
13.30	LUNCH	
14.30	SESSION 3 - The Global Scenario for EPTRI Chairs: M. Turner, C. Giaquinto	
	How European Biological and Medical Sciences Research Infrastructures boost Innovation	J. Demotes
	Relationship with the existing RIs and other organization groups	A. Ceci
	Map of units and context analysis	S. Wimmer
	Ethical issues and set up of the Project Ethical Advisory Board	M. Migdal
	Governance and sustainability: key strategies to design, establish and govern a sustainable European research infrastructure	F. De Man
16.30	SESSION 4 - Project management and coordination Chairs: L. Mangiarini, M. Lupo	
	ID-EPTRI General Assembly and Bodies designation	D. Bonifazi
	ID-EPTRI Project management, requirements and timelines	G. Vecchia
	Horizon 2020: Administrative, Legal and Financial Issues	S. Faggion
	Financial reporting in Horizon 2020	M. Montanaro
18.30	END OF THE DAY	



Day 2, January 16th, 2018

8.30	SESSION 5 - Paediatric Medicines Discovery (WP5) and Biomarkers (WP6) Chairs: O. Mukvich, J. Kindblom	
	Design of the thematic platform supporting paediatric medicines discovery and early drug development	E. Mikros
	Run in a Proof-of-concept study simulation	H. Kubova
	Design of the thematic platform supporting biomarkers in paediatric medicines development	M. Lavitrano
	Feasibility study for the development of biomarkers in paediatrics	M. Kleanthous
10.00	COFFEE BREAK	
10.30	SESSION 6 - Paediatric Pharmacology (WP7) and Formulation Science (WP8) Chairs: V. Kazantsev, S. Stasenko	
	Design of the thematic platform on paediatric pharmacology	E. Jacqz Aigrain
	Physiologically based PK/PD and modelling	S. de Wildt
	Design of the thematic platform on formulation science	C. Tuleu
	Feasibility study on use of in vitro / in vivo tools for Taste Assessment	
	Techniques employed for taste masking of pharmaceuticals	N. De Nora
12.00	SESSION 7 - Underpinning Paediatric Studies (WP9) Chairs: D. Nika, P. Wenger	
	Design of the thematic platform to relate work that underpins medicines development to paediatric clinical studies	M. Turner
	Certification of paediatric clinical research centres	A. Simonetti
	The PedCRIN project to integrate paediatric tools in an existing RI	J. Demotes
13.00	LUNCH	
14.00	SESSION 8 - Concept design, IT structure and feasibility assessment of the infrastructure Chairs: M. Mellado, O. Della Pasqua	
	Organisation and technical design (model) of the infrastructure	M. Felisi



Information Technologies (IT) supporting EPTRI activities	F. Bonifazi
Design model for collaboration with the existing RIs in Biomed field	G. Migliaccio
Paediatric Data use and reuse	I. Wong
Common data models for re-use of health care data	M. Sturkenboom
SESSION 9 - Communication, networking and patients involvement	
15.30	Chair: F. Kalambayi, D. Bonifazi
Communication and dissemination plan in the project	M. Lupo
Communication and dissemination materials in the project	J. Claverol
Project website	L. Mangiarini
Plan for patients participation	B. Nafria
The patients' perspective and the European RIs landscape	Patients Association Representatives
17.00	GOODBYE COCKTAIL
18.00	END OF THE DAY

