



5th Training course on "Quality assurance, variant interpretation and data management in the NGS diagnostics era"

18-20 Oct 2023, Warsaw, Poland

Tentative PROGRAM – v0805 -

EJP RD European Joint Programme on Rare Diseases

H2020-SC1-2018-Single-Stage-RTD
SC1-BHC-04-2018
Rare Disease European Joint Programme Cofund



Grant agreement number 825575

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Pillar 3 - WP14.2 - Standards and quality of genetics/genomics data in laboratory and clinical practice

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Day 1 - 18 October 2023

		Speaker	Affiliation
	Arrivals		
11:00-	Registration Desk		
13:00			
12:00	Lunch		
13:00	Opening & Welcome	Krystyna	
		Chrzanowska	
Session 1	NGS diagnostics		
13:15	The importance of NGS in daily clinical practice: the impact of a	Saskia	Paracelsus Medical University,
	genetic diagnosis for treatment and beyond	Wortmann	Salzburg, Austria
14:15	Efficient analysis of whole genome sequencing data	Tomasz	University of Bergen, Norway
		Stokowy	
15:15	NIPT	Eftychia or	
		Beata	
16:00	Coffee Break		
Session 2 - Hands-on: Variant filtering and prioritisation		Leslie	CNAG-CRG
		Matalongo,	
		Steven Laurie	
16:30	Short introduction on data filtering, prioritisation and		
	classification + GPAP		
17:00	Case demo introducing key concepts		
17:30	Explanation and use of variant classification following ACMG		
	guidelines		
18:00	End Day 1		





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Day 2 - 19 October 2023

		Speaker	Affiliation
Session 3	- Quality in NGS diagnostics		
9:00	Diagnostic Laboratory quality assurance and management	Gert Matthijs	KU Leuven, Belgium
9:30	The Polish experience on the external quality assurance of genetic testing.	Beata Nowakowska	Institute of Mother and Child, Warsaw, Poland
10:00	Recommendations for whole genome sequencing in diagnostics for rare diseases	Erika Souche	KU Leuven, Belgium
10:30	Coffee Break		
Session 5	– Clinical and functional validation		
11:00	The importance of (reverse) phenotyping, communication and functional validation for variant interpretation	Saskia Wortmann	Paracelsus Medical University, Salzburg, Austria
11:45	Multi OMICS data in Mendelian disease diagnostics	Holger Prokisch	Technical University of Munich, Germany
12:30	Lunch		
Session 4	- Innovation and research		
13:30	Novel genomic applications in preimplantation genetic testing and prenatal diagnosis	Eftychia Dimitriadou	KU Leuven, Belgium
14:15	Long read sequencing for rare diseases	Erika Souche	KU Leuven, Belgium
15:00	Free slot –		
15:45	Coffee Break		
Session 6	- Hands-on (part 2) -: Variant filtering and prioritisation		
16:15	Title to be decided	Leslie Matalongo, Steven Laurie	
17:30	End Day 2		





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Day 3 - 20 October 2023

		Speaker	Affiliation
Session 4	- Validation of NGS tools and pipelines		
9:00	Validation of NGS variant pipeline/WGS	Erika Souche	KU Leuven, Belgium
9:30	Validation of PGT pipelines	Eftychia Dimitriadou	KU Leuven, Belgium
10.00	NGS techniques for gene discovery: detection and variant interpretation	Rafał Płoski	Warsaw Medical University, Poland
10:30	Coffee Break		
Session 8	- Future prospects		
11:00	How exome sequencing changes the prenatal diagnostics	Beata Nowakowska	Institute of Mother and Child, Warsaw
11:30	Free slot -		
12:15	Closing remarks	Gert / Krystyna	
12:30	Lunch		
13:30	End Day 3		