



## DELIVERABLE D2.1 V1.1

REPORT DESCRIBING THE EATRIS-PLUS FAIRIFICATION TEMPLATE AND WORKFLOWS

WP2 – Data Stewardship and integration of omic research in Personalised Medicine

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Contractual Delivery Date: 31 December 2021 [M24]

Actual Delivery Date: v1 21. December 2021; v1.1 22 June 2023

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Grant agreement no. 871096

Horizon 2020

H2020-INFRADEV-3

Type of Action: RIA



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## EXECUTIVE SUMMARY

The key scientific output of the EATRIS-Plus project is to develop a Multi-omic Toolbox available for researchers in order to have a better understanding of the molecular profiles in personalised medicine.

Within work packages 1, 2, and 3, we implement practices based on the FAIR (Findability, Accessibility, Interoperability, and Reusability) principles to ensure reproducibility of our work and to make the multi-omics cohort data reusable for future translational research. The aim of this deliverable (D2.1) is to describe the EATRIS-Plus FAIRification process, and list standards, formats, and tools used in this process. Metadata templates developed and used in this project are shared.

## PROJECT OBJECTIVES

Our flagship project EATRIS-Plus aims to build further capabilities and deliver innovative scientific tools to support the long-term sustainability strategy of EATRIS as one of Europe's key European research infrastructures for Personalised Medicine.

The main goals of the EATRIS-Plus will be to:

- Consolidate EATRIS capacities in the field of Personalised Medicine (particularly omics technologies) to better serve academia and industry and augment the number of EATRIS Innovation Hubs with large pharma;
- Drive patient empowerment through active involvement in the infrastructure's operations;
- Expand strategic partnerships with research infrastructures and other relevant stakeholders, and
- Further strengthen the long-term sustainability of the EATRIS financial model.
- Develop a Multi-omic Toolbox for researchers.

## DETAILED REPORT ON THE DELIVERABLE

### BACKGROUND

Transparency about samples, experimental methods, data, and data analyses are crucial for reproducibility of translational research and to enable and promote reuse of omics data. We aim to increase the value of data generated within EATRIS-Plus and conducted research by applying FAIR principles to data, metadata, and analysis workflows. This should set an important example and provide guidelines for other research projects in the field of personalized health and personalized medicine.

### DESCRIPTION OF WORK

As part of this deliverable (D2.1), we describe the EATRIS-Plus FAIRification strategy (Appendix 1: EATRIS-Plus FAIRification strategy) to make the generated multi-omics cohort data reusable for future translational research.



We identified relevant data and metadata standards to report and describe data generated by different omics technologies. The existing ISA (Investigation, Study, Assay) metadata framework (<https://isa-tools.org/>)<sup>1</sup> is used to capture experimental metadata. ISA-Tab metadata is used and supported by a wide range of single-omics data repositories (<https://www.isacommons.org/>). We collected relevant fields needed to describe different omics experiments among the project partners. Based on this collection, a Jupyter notebook was developed to generate ISA-Tab and ISA-JSON files for the multi-omics data set that complies with reporting guidelines for individual omics types. Importantly, the ISA framework enables the use of controlled vocabularies and ontologies supporting both human-readability and machine-readability. The ISA-Tab template files being developed in EATRIS-Plus will serve as templates for collecting the required metadata for integration of multi-omics in the context of research projects around personalized health and personalized medicine. The ISA templates as well as workflow (Jupyter notebook) to create them are available at <https://github.com/EATRIS-Plus/eatris-plus-isa>. They are also shared as part of the EATRIS-Plus multi-omics toolbox.

In addition to the experimental metadata, the multi-omics data set is accompanied by phenotypic information. We chose the GA4GH Phenopackets model to capture Phenotype information in a FAIR format. Fields were mapped to ontology terms and a Phenopacket json files was created programmatically. The workflow is available at [https://github.com/EATRIS-Plus/phenopackets\\_template](https://github.com/EATRIS-Plus/phenopackets_template).

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## COMMUNITY OUTREACH

We organized a workshop on FAIRification of multi-omics metadata (New Trends in Bioinformatics, workshop 2; <https://eccb2022.org/ntb-w02/>) as part of the 21st European Conference on Computational Biology (ECCB2022) to reach out to other initiatives, share the EATRIS-Plus approach, and discuss challenges and solutions for multi-omics data FAIRification with the research community.

## NEXT STEPS

We keep monitoring developments in different omics communities and collaborating with other initiatives. The EATRIS-Plus FAIRification templates and corresponding Jupyter notebook will be updated accordingly and shared with the research community as part of the EATRIS-Plus multi-omics toolbox.

## ABBREVIATIONS

FAIR acronym for Findability, Accessibility, Interoperability, and Reusability

GA4GH Global Alliance for Genomic & Health

ISA Investigation, Study, Assay (metadata framework)

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<sup>1</sup> Sansone, S.-A., Rocca-Serra, P., Field, D., Maguire, E., Taylor, C., Hofmann, O., Fang, H., Neumann, S., Tong, W., Amaral-Zettler, L., Begley, K., Booth, T., Bougueleret, L., Burns, G., Chapman, B., Clark, T., Coleman, L.-A., Copeland, J., Das, S., ... Hide, W. (2012). Toward interoperable bioscience data. *Nature Genetics*, 44(2), 121–126. <https://doi.org/10.1038/ng.1054>



## DELIVERY AND SCHEDULE

The first version of the deliverable was submitted within the contractual delivery date. Following the review of the second reporting period and the updates requested the deliverable is resubmitted on 22 June 2023.

## ADJUSTMENTS MADE

The updates to this version of the deliverable include descriptions of and links to the EATRIS-Plus FAIRification templates for multi-omics metadata and phenotype data. These are publicly available. Additionally, a section on community outreach was included. Appendix 1 (EATRIS-Plus FAIRification strategy) was updated to include FAIRification of computational workflows.

## APPENDICES

- Appendix 1: EATRIS-Plus FAIRification strategy



## APPENDIX 1: EATRIS-PLUS FAIRIFICATION STRATEGY

The EATRIS-Plus multi-omics cohort data set comprises different types of molecular profiles (genomic variation and DNA methylation, mRNA, miRNA, protein, and metabolite levels) and phenotypic information of >100 healthy individuals. Guided by the FAIR principles that were first introduced in 2016 (Wilkinson et al., 2016), we implement data management practices to ensure transparency and reproducibility of our research, and reuse of the data generated within EATRIS-Plus. Our FAIRification strategy is aligned with international standards and guidelines related to data stewardship in general and to specific omics platforms.

The acronym FAIR stands for Findability, Accessibility, Interoperability, and Reusability. Different recommended practices (<https://www.go-fair.org/fair-principles/>) contribute to improve each of these aspects. The addition of rich metadata improves both findability and reusability. The use of standardized vocabularies, ontologies and unique and persistent identifiers for metadata and features enables machine-readability and thereby enhances interoperability. This is important for both integration of different types of omics data (mapping of identifiers) as well as integration with other data sets.

The ISA (Investigation, Study, Assay) metadata framework (Sansone et al., 2012) allows to capture experimental metadata. This includes metadata on samples and sample processing, omics measurements including sample preparation, measurement protocols, and analysis protocols. Different fields used to describe the metadata are presented in the EATRIS-Plus example ISA-Tab files. We also develop a Jupyter notebook employing the ISA-API (Johnson et al., 2021) which can be used to create the ISA-Tab and ISA-JSON files, and will be made publicly available as part of the EATRIS-Plus multi-omics toolbox. Minimal information standards and guidelines for submission to omics data repositories are being followed so that data submission is facilitated and the reuse of data in those repositories is enhanced. The standards and guidelines are described below.

For **metabolomics**, we follow the guidelines of the EBI MetaboLights repository (Haug et al., 2019) which adheres to Metabolomics Standard Initiative (MSI) standards (Salek et al., 2015; Spicer et al., 2017; van Rijswijk et al., 2017). These include reporting sample processing such as extraction and labeling, measurement specifications such as chromatography and mass spectrometry, and data analysis protocols. We link to the applied protocols (SOPs) using unique and persistent identifiers.

For **RNAseq** (mRNA and microRNA sequencing), we follow MINSEQE (Minimum Information About a Next-generation Sequencing Experiment) guidelines (Brazma et al., 2012) to provide information about the sequencing experiment.

For **WGS**, we follow the General Guide on ENA Data Submission (<https://ena-docs.readthedocs.io/en/latest/submit/general-guide.html>) to provide information about the sequencing experiment.

For **proteomics**, we follow guidelines of the HUPO-PSI, specifically MIAPE – The Minimum Information About a Proteomics Experiment (Taylor et al., 2007) in following parts, MIAPE: Mass Spectrometry version 2.98, MIAPE: Column Chromatography version 1.1 and MIAPE: Mass Spectrometry Informatics version 1.1. and mzTab v1.0.0. We have described our sample preparation protocol in our applied SOPs (Scherer, 2021) in detail.



For **microRNA qRT-PCR** determination, we follow the general recommendations for qPCR included in the MIQE guideline (Bustin et al., 2009). Since we used the miRCURY LNA miRNA Focus PCR Panel from Qiagen in serum samples, we follow Qiagen's guidelines for profiling miRNAs in body fluids (Guidelines for Profiling Biofluid miRNAs – QIAGEN (Qiagen, 2019)). For data normalization, we follow the recommendations described by Mestdagh et al., 2009 and Marabita et al., 2016 (Marabita et al., 2016; Mestdagh et al., 2009). These references describe procedures for each of the main steps required for the miRNA determination by qPCR including sample obtaining and processing, RNA isolation from fluids, cDNA generation by reverse transcription, determination of miRNAs panel by quantitative PCR, quality controls and data normalization.

For **DNA methylation sequencing (EM-seq)**, we follow the ENA Metadata Model to provide meta data about the library and sequencing experiment: <https://ena-docs.readthedocs.io/en/latest/submit/general-guide/metadata.html>

For reporting of measured omics data, we use community standard formats recommended by the aforementioned reporting guidelines. These are listed in Table 1. Database identifiers to report which molecular features were measured are summarized in Table 2.

**Table 1: Used formats for raw and processed data per omics technology.**

Omics technology	Raw data format	Processed data format(s)	Format used for multi-omics data integration
Targeted metabolomics	.raw (proprietary format), .mzML (open format)	.tsv - metabolite annotation file (MAF)	.tsv - metabolite annotation file (MAF)
Proteomics	.RAW file (Thermo XCalibur)	mzTab (proteomic standard data format)	.txt with pre-processed protein lists exported from ProteomeDiscoverer
mRNA sequencing	.fastq.gz	.bam, .bam.bai additional file formats in the pre-processed data are: .html, .txt, .zip, .csv, .sf, .tsv, .pdf, .r and .gtf	.tsv
microRNA sequencing	.fastq.gz	.bam additional file formats in the pre-processed data are: .bai, .html, .txt, .zip, .gz, .csv, .tsv, .pdf, .gff and .gtf	.tsv
microRNA qRT-PCR	.txt	Normalized Ct (Crossing Threshold)  .xls	.txt and .xls following NCBI GEO database standard
DNA methylation sequencing	.fastq	.bam .bedGraph	.bedgraph
WGS	.fastq.gz	.bam, .vcf	.tsv

Table 2: Database identifiers for molecular features per omics technology.

Omics technology	Entity	Database identifier(s)
Targeted metabolomics	Metabolite	CHEBI, HMDB
Untargeted lipidomics	Lipid	LipidBlast name
Proteomics	Peptide Protein	UniProtKB Sequence UniProtKB Accession Number
mRNA sequencing	Transcript	Ensembl ID
microRNA sequencing	microRNA	miRBase ID and Accession number
microRNA qRT-PCR	microRNA	miRBase ID (release 20)
DNA methylation sequencing	Genomic coordinate	Unique identifier based on chromosome (referred to by GenBank ID and version) and genomic coordinate (GRCh38) corresponding to CpG site
WGS	Gene	HUGO Gene Nomenclature (HGNC), Ensembl Gene ID in combination with a genomic coordinates (GRCh38)

The original FAIR principles apply to research data. However, in order to fully ensure reproducibility of conducted research, data analysis workflows need to be FAIR as well. We follow best practices recommended (Gruening et al., 2018; Jiménez et al., 2017) by different initiatives to improve FAIRness of computational research workflows as well. This includes version-control (git), documentation, citation, adding license, open source code sharing (<https://github.com/>), containerization (Docker, Singularity), workflow management and registration (). Metadata about computational workflows will be captured using RO-Crate and workflows will be published on WorkflowHub and shared via the Multi-omics Toolbox. An example workflow annotated with machine-readable metadata was developed in collaboration with The Netherlands X-omics Initiative (<https://doi.org/10.1101/2023.06.07.543986>).

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