FWO DMP Template - Flemish Standard Data Management Plan

Version KU Leuven

Project supervisors (from application round 2018 onwards) and fellows (from application round 2020 onwards) will, upon being awarded their project or fellowship, be invited to develop their answers to the data management related questions into a DMP. The FWO expects a **completed DMP no later than 6 months after the official start date** of the project or fellowship. The DMP should not be submitted to FWO but to the research co-ordination office of the host institute; FWO may request the DMP in a random check.

At the end of the project, the **final version of the DMP** has to be added to the final report of the project; this should be submitted to FWO by the supervisor-spokesperson through FWO's e-portal. This DMP may of course have been updated since its first version. The DMP is an element in the final evaluation of the project by the relevant expert panel. Both the DMP submitted within the first 6 months after the start date and the final DMP may use this template.

The DMP template used by the Research Foundation Flanders (FWO) corresponds with the Flemish Standard Data Management Plan. This Flemish Standard DMP was developed by the Flemish Research Data Network (FRDN) Task Force DMP which comprises representatives of all Flemish funders and research institutions. This is a standardized DMP template based on the previous FWO template that contains the core requirements for data management planning. To increase understanding and facilitate completion of the DMP, a standardized **glossary** of definitions and abbreviations is available via the following link.

	1. General Project Information
Name Grant Holder & ORCID	Brandán Pedre Pérez, 0000-0003-2158-2807
Contributor name(s) (+ ORCID) & roles	Peter Dedecker (supervisor) 0000-0002-1882-2075
	Elena Levtchenko (co-supervisor) 0000-0002-8352-7312
	Lambertus van den Heuvel (co-supervisor) 0000-0003-3917-6727
Project number ¹ & title	1276324N, Genetically-encoded cysteine and cystine biosensors for real-time monitoring and diagnostics
Funder(s) GrantID ²	FWO
Affiliation(s)	X KU Leuven
	☐ Universiteit Antwerpen
	☐ Universiteit Gent
	☐ Universiteit Hasselt
	☐ Vrije Universiteit Brussel
	☐ Other:
	ROR identifier KU Leuven: 3E230514

¹ "Project number" refers to the institutional project number. This question is optional. Applicants can only provide one project number.

² Funder(s) GrantID refers to the number of the DMP at the funder(s), here one can specify multiple GrantIDs if multiple funding sources were used.

Please provide a short project description

Cysteine is an essential amino acid that is required for proper cell functioning via its redox-active thiol group. Dysregulation of cysteine metabolism is implicated in diseases such as cystinosis, while insufficient cysteine supply triggers ferroptosis, a type of cell death with a promising potential in cancer therapy. Despite its seemingly well-established nature, many fundamental aspects in cysteine metabolism remain unknown due to a lack of tools that can efficiently monitor cysteine levels in the native environment, subcellular compartments, and in clinical samples.

This project responds to this technology gap by developing genetically-encoded fluorescent biosensors for cysteine and its oxidized form cystine. These biosensors will combine a fluorescent protein reporter with a specific cysteine or cystine-binding protein. The resulting biosensors can be expressed by cells and will convert the presence of these molecules into a change in fluorescence, that can be read using fluorescence microscopes or plate readers, ensuring compatibility with a wide range of settings. I will then use the sensors in cellulo to understand how cystine is transformed into cysteine intracellularly, as well as to develop a cheaper and faster diagnostic procedure to measure cystine levels in biological samples. These cysteine/cystine biosensors will set the basis for research breakthroughs in cysteine/cystine metabolism, from their use in this project and their use by other researchers.

2. Research Data Summary

List and describe all datasets or research materials that you plan to generate/collect or reuse during your research project. For each dataset or data type (observational, experimental etc.), provide a short name & description (sufficient for yourself to know what data it is about), indicate whether the data are newly generated/collected or reused, digital or physical, also indicate the type of the data (the kind of content), its technical format (file extension), and an estimate of the upper limit of the volume of the data ³.

				ONLY FOR DIGITAL DATA	ONLY FOR DIGITAL DATA	ONLY FOR DIGITAL DATA	ONLY FOR PHYSICAL DATA
Dataset	Description	New or Reused	Digital or	Digital Data Type	Digital Data	Digital Data	Physical Volume
Name			Physical		Format	Volume (MB, GB,	
						TB)	
DNA plasmids	New biosensor plasmids for	⊠ Generate new	☐ Digital	☐ Audiovisual		□ < 1 GB	In order of tens of microliters
	bacterial or	data	☐ Physical	☐ Images		□ < 100 GB	moromoro
	eukaryotic cell epxression;	☐ Reuse existing		☐ Sound		□ < 1 TB	
	obtained by cloning	data		☐ Numerical		□ < 5 TB	
	strategies			☐ Textual		□ > 5 TB	
				☐ Model		□NA	
				☐ Software			
				☐ Other:			
DNA and amino	DNA and amino	⊠ Generate new	☐ Digital	☐ Audiovisual	.fasta .ab1	⊠ < 1 GB	
acid sequences acid sequences	data	☐ Physical	☐ Images	.ab i	□ < 100 GB		
		☐ Reuse existing		☐ Sound		□ < 1 TB	
		data		⋈ Numerical		□ < 5 TB	
						□ > 5 TB	
				☐ Model		□NA	
				☐ Software			
				☐ Other:			
Plate reader fluorescence intensity values data	⊠ Generate new	□ Digital	☐ Audiovisual	.xlsx .txt	□ < 1 GB		
	data	☐ Physical	☐ Images	.txt	⊠ < 100 GB		
	☐ Reuse existing		☐ Sound		□ < 1 TB		
	data				□ < 5 TB		
						□ > 5 TB	
				☐ Model		□NA	

				☐ Software			
				☐ Other:			
Images	Images & videos	⊠ Generate new	☐ Digital	☐ Audiovisual	.tif, .avi, .mov	□ < 1 GB	
	acquired by fluorescence	data				□ < 100 GB	
	microscopy	☐ Reuse existing		☐ Sound		□ < 1 TB	
		data		⊠ Numerical		⊠ < 5 TB	
						□ > 5 TB	
				☐ Model		□ NA	
				☐ Software			
				☐ Other:			
Data analysis	Analysis, output, and	☑ Generate new	☐ Digital		.tif, .pxp, .pptx, .prism,	□ < 1 GB	
	data visualization	data	□ Physical		.mat, .xlsx, .pdf	□ < 100 GB	
		☐ Reuse existing		☐ Sound		□ < 1 TB	
		data		☐ Numerical		□ < 5 TB	
				☐ Textual		□ > 5 TB	
				☐ Model		□ NA	
				☐ Software			
				☐ Other:			
GUIDANCE:							
The data descript	ion forms the basis o	f your entire DMP, so make	e sure it is detai	led and complete. It in	cludes digital and phys	sical data and encompas	sses the whole spectrum
	•	nd analysed data including		•			-
		nical issues are associated.				nclude your own manus	cripts, theses and
presentations; do	cumentation is an int	egral part of your datasets	s and should de	scribed under docume	ntation/metadata.		

RDM Guidance on data

³ Add rows for each dataset you want to describe.

If you reuse existing data, please specify the source, preferably by using a persistent identifier (e.g. DOI, Handle, URL etc.) per dataset or data type.	NA NA
Are there any ethical issues concerning the creation and/or use of the data (e.g. experiments on humans or animals, dual use)? If so, refer to specific datasets or data types when appropriate and provide the relevant ethical approval number.	 Yes, human subject data; provide SMEC or EC approval number: Yes, animal data; provide ECD reference number: Yes, dual use; provide approval number: No Additional information:
Will you process personal data ⁴ ? If so, please	,
refer to specific datasets or data types when appropriate and provide the KU Leuven or UZ	
Leuven privacy register number (G or S number).	Additional information:
Does your work have potential for commercial	☐ Yes
valorization (e.g. tech transfer, for example spin-	⊠ No
offs, commercial exploitation,)?	If yes, please comment:
If so, please comment per dataset or data type	
where appropriate.	
Do existing 3rd party agreements restrict	□ Yes
exploitation or dissemination of the data you	⊠ No
(re)use (e.g. Material/Data transfer agreements,	If yes, please explain:
research collaboration agreements)?	
If so, please explain to what data they relate and	
what restrictions are in place.	

⁴ See Glossary Flemish Standard Data Management Plan

Are there any other legal issues, such as	☐ Yes
intellectual property rights and ownership, to be	⊠ No
managed related to the data you (re)use?	If yes, please explain:
If so, please explain to what data they relate and	
which restrictions will be asserted.	

3. Documentation and Metadata Clearly describe what approach will be followed - Electronic lab book written by the grant holder, clearly labeled with dates and experiment descriptions to capture the accompanying information - PowerPoint files summarizing project progress necessary to keep data understandable and readme.txt files for developed imaging methods (for end-users after publishing) usable, for yourself and others, now and in the - Protocols in .pdf format future (e.g. in terms of documentation levels and - DNA sequences and maps annotated and described on an online platform Benchling, within the project types required, procedures used, Electronic Lab file of the host group Notebooks, README.txt files, Codebook.tsv etc. where this information is recorded). RDM guidance on documentation and metadata. Will a metadata standard be used to make it ☐ Yes easier to find and reuse the data? \bowtie No If yes, please specify (where appropriate per dataset or data type) which metadata standard will be used: If so, please specify which metadata standard will be used. If not, please specify which metadata will be created to make the data If no, please specify (where appropriate per dataset or data type) which metadata will be created: easier to find and reuse. REPOSITORIES COULD ASK TO DELIVER METADATA IN A CERTAIN FORMAT, WITH SPECIFIED ONTOLOGIES AND VOCABULARIES, I.E. STANDARD LISTS WITH UNIQUE IDENTIFIERS.

	4. Data Storage & Back-up during the Research Project
Where will the data be stored?	☐ Shared network drive (J-drive)
Consult the interactive KIII amon storage guide to	□ Personal network drive (I-drive)
Consult the <u>interactive KU Leuven storage guide</u> to find the most suitable storage solution for your data.	☐ OneDrive (KU Leuven)
,	☐ Sharepoint online ☐ Sharepoint on-premis
	☐ Large Volume Storage
	☐ Digital Vault
	☐ Other:
How will the data be backed up?	⊠ Standard back-up provided by KU Leuven ICTS for my storage solution □ □
WHAT STORAGE AND BACKUP PROCEDURES WILL BE IN PLACE TO	□ Personal back-ups I make (specify)
PREVENT DATA LOSS?	☐ Other (specify)
Is there currently sufficient storage & backup	⊠ Yes
capacity during the project? If yes, specify	□ No
concisely. If no or insufficient storage or backup	
capacities are available, then explain how this will be taken care of.	If no, please specify:

How will you ensure that the data are securely stored and not accessed or modified by unauthorized persons?	Data stored in OneDrive (KU Leuven) is secured through two-step authentication process which requires my personal cellphone.
CLEARLY DESCRIBE THE MEASURES (IN TERMS OF PHYSICAL SECURITY, NETWORK SECURITY, AND SECURITY OF COMPUTER SYSTEMS AND FILES) THAT WILL BE TAKEN TO ENSURE THAT STORED AND TRANSFERRED DATA ARE SAFE. Guidance on security for research data	Data stored in personal network drive (I-drive) will be stored in a drawer locked by a key, whose access is restricted to grant holder, supervisor and lab manager.
What are the expected costs for data storage	Personal network drives will be purchased using the FWO bench fee.
and backup during the research project? How will these costs be covered?	Free data storage capacity provided by KU Leuven OneDrive is enough for this project.

5. Data Preservation after the end of the Research Project		
Which data will be retained for at least five	☑ All data will be preserved for 10 years according to KU Leuven RDM policy	
years (or longer, in agreement with other	\square All data will be preserved for 25 years according to CTC recommendations for clinical trials with	
retention policies that are applicable) after the	medicinal products for human use and for clinical experiments on humans	
end of the project? In case some data cannot be	\square Certain data cannot be kept for 10 years (explain)	
preserved, clearly state the reasons for this		
(e.g. legal or contractual restrictions,		
storage/budget issues, institutional policies).		
Guidance on data preservation		

Where will these data be archived (stored and curated for the long-term)? Dedicated data repositories are often the best place to preserve your data. Data not suitable for preservation in a repository can be stored using a KU Leuven storage solution, consult the interactive KU Leuven storage guide.	 □ KU Leuven RDR ☑ Large Volume Storage (longterm for large volumes) □ Shared network drive (J-drive) ☑ Other (specifiy): Physical data, such as plasmids, will be stored in freezers present in the host group.
What are the expected costs for data preservation during the expected retention period? How will these costs be covered?	Physical drives will be purchased during the project (~300EUR) and no maintenance cost is expected after the end of the project.

6. Data Sharing and Reuse		
Will the data (or part of the data) be made available for reuse after/during the project? Please explain per dataset or data type which data will be made available. Note that 'Available' does not necessarily mean that the data set becomes openly available, conditions for access and use may apply. Availability in this question thus entails both open & restricted access. For more information: https://wiki.surfnet.nl/display/standards/info-eu-repo/#infoeurepo-AccessRights	 Yes, as open data Yes, as embargoed data (temporary restriction) Yes, as restricted data (upon approval, or institutional access only) No (closed access) Other, please specify: 	
If access is restricted, please specify who will be able to access the data and under what conditions.		

Are there any factors that restrict or prevent the sharing of (some of) the data (e.g. as defined in an agreement with a 3rd party, legal restrictions)? Please explain per dataset or data type where appropriate.	 Yes, privacy aspects Yes, intellectual property rights Yes, ethical aspects Yes, aspects of dual use Yes, other No If yes, please specify:
Where will the data be made available?	☐ KU Leuven RDR
If already known, please provide a repository	☐ Other data repository (specify): Zenodo data repository
per dataset or data type.	□ Other (specify): Preprints, open access publications
When will the data be made available?	 ☑ Upon publication of research results ☑ Specific date (specify): 1 year embargo for green open access publications ☐ Other (specify)
Which data usage licenses are you going to provide? If none, please explain why.	 ⊠ CC-BY 4.0 (data) ⊠ Data Transfer Agreement (restricted data): In case of plasmids described in preprints
A DATA USAGE LICENSE INDICATES WHETHER THE DATA CAN BE REUSED OR NOT AND UNDER WHAT CONDITIONS. IF NO LICENCE IS GRANTED, THE DATA ARE IN A GREY ZONE AND CANNOT BE LEGALLY REUSED. DO NOTE THAT YOU MAY ONLY RELEASE DATA UNDER A LICENCE CHOSEN BY YOURSELF IF IT DOES NOT ALREADY FALL UNDER ANOTHER LICENCE THAT MIGHT PROHIBIT THAT. Check the RDR quidance on licences for data and software sources code or consult the License selector tool to help you choose.	☐ MIT licence (code) ☐ GNU GPL-3.0 (code) ☐ Other (specify)

Do you intend to add a PID/DOI/accession number to your dataset(s)? If already available, please provide it here.	 ⊠ Yes, a PID will be added upon deposit in a data repository □ My dataset already has a PID □ No □
INDICATE WHETHER YOU INTEND TO ADD A PERSISTENT AND UNIQUE IDENTIFIER IN ORDER TO IDENTIFY AND RETRIEVE THE DATA.	
What are the expected costs for data sharing? How will these costs be covered?	Data deposit in Zenodo repository is free of charge. Open access publication fees depend on type of open access chosen and available funds from the FWO bench fee. Green open access publication is usually free, while I estimate gold open access publication as 3000-4000EUR/publication. Expected cost used for publications ~6000-8000EUR.

7. Responsibilities	
Who will manage data documentation and metadata during the research project?	Grant holder: Brandán Pedre Pérez
Who will manage data storage and backup	Grant holder: Brandán Pedre Pérez
during the research project?	Supervisor: Peter Dedecker at the end of the research project to ensure its 10 year preservation.
Who will manage data preservation and	Grant holder: Brandán Pedre Pérez
sharing?	
Who will update and implement this DMP?	Grant holder: Brandán Pedre Pérez