





# 17 Doctoral Candidate Positions within the MSCA Doctoral Network MC4DD – Macrocycles for Drug Discovery

Subject to the successful signing of the EU Grant Agreement, a call for 17 DC (Doctoral Candidate) positions is open within the context of the Doctoral Network (DN) project MC4DD.

The Doctoral Network **MC4DD** "MC4DD – Macrocycles for Drug Discovery", funded within the framework of the Marie Skłodowska-Curie Actions (MSCA), follows an interdisciplinary and cross-sectoral approach by bringing together leading experts in macrocyclic drug discovery from academia and industry from the fields of organic synthesis, medicinal, high-throughput and computational chemistry, pharmacological and structural analytics, and modelling.

Eight academic research groups and five industrial partners, coordinated by Technische Universität Darmstadt in Germany, join forces in **MC4DD** to create a mobility and training platform for young scientists by means of cross-site, interdisciplinary research projects. The DCs will work on individual research projects to expand the opportunities of macrocycles as next-generation drug modalities.

Eligibility criteria (see also MSCA guide for applicants)

**Supported researchers** must be doctoral candidates, i.e. not already in possession of a doctoral degree at the date of the recruitment.

**Mobility Rule:** researchers must not have resided or carried out their main activity (work, studies, etc.) in the country of the recruiting beneficiary >12 months in the 36 months preceding their recruitment date.

#### Monthly allowances (employee gross):

Living allowance\*: 3400 € Mobility allowance: 600 € Family allowance\*\*: 660 €



\*adjusted by country correction coefficient \*\*if applicable

Further information: <u>www.mc4dd.com</u>

#### What we offer

The project offers research and training excellence in medicinal chemistry and drug discovery. The partners of MC4DD are leading research groups in this field and their research institutes actively promote young researchers. The 8 academic research groups and 5 industry partners join forces in MC4DD to create a platform of intersectoral and multidisciplinary mobility and training. To complement the academic and scientific goals of the DCs, the project offers customized research projects, structured interdisciplinary local and network-wide transferable skills training activities, and secondments at top-ranking European universities and industry partners.

#### **Candidate profile:**

- MSc or equivalent in Chemical, Pharmaceutical or Life Sciences with a solid chemistry training
- Research experience with organic, medicinal, analytical, computational or bio-chemistry prior experience with macrocycles and beyond-Rule-of-5 compounds is advantageous
- Scientific interest, dedication to research, and career goal to work in drug discovery
- Appreciation for interdisciplinarity and proactive drive to collaborate across fields
- Proficiency in English, good communication skill and social competence



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### **Applications:**

- Applications are written in **English** and include a **Cover Letter** indicating the motivation for the project, a **CV**, copies of **transcripts of degrees** obtained, as well as contact details of **two references** (ideally at least one of them with extensive EU experience). The inclusion of a **two-page project proposal** is highly encouraged and will be an evaluation benefit.
- Applications should be submitted via the **procedure specified for each project** as a **single pdf** file by **31.08.2024**, unless specified otherwise for the specific project (see below)
- Applicants should apply to only one project
- The individual DC projects are expected to start between 01.11.2024 and 31.05.2025

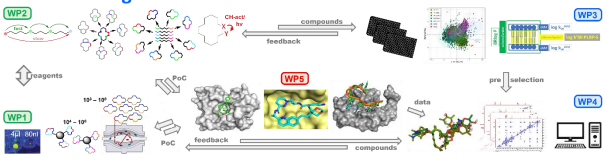
### Network (Main supervisors and sites):



### **Research areas:**

EPFL <sup>1,3</sup> LU <sup>1</sup> TUDa <sup>1</sup> UORSY <sup>2</sup>	Roche <sup>2,3</sup> AstraZ <sup>2</sup>	UU <sup>1</sup> ABBV <sup>2</sup> UniTO <sup>1</sup> TUDa <sup>1</sup> AZ <sup>3</sup> SANOFI <sup>3</sup> Roche <sup>2,3</sup>
WP1 Improved access to screenable MC libraries		WP3 Generation & mining of large MC-data sets
	WP5 Proof of concept for industry-relevant targets	
WP2 Synthetic access to diverse low-peptidic MCs		WP4 In-depth analysis of model macrocycles
TUDa <sup>1</sup> UOY <sup>1,3</sup> UCL <sup>1,3</sup> Sanofi <sup>2</sup>	TUDa <sup>1</sup> Sanofi <sup>2</sup> ABBV <sup>2</sup>	UniTO <sup>1</sup> UU <sup>1</sup> ETHZ <sup>1,3</sup> Roche <sup>2,3</sup>
Enamine DE <sup>4</sup> Analyticon <sup>4</sup> Bayer <sup>4</sup> WP6 Training Selvita <sup>4</sup> Novo <sup>4</sup> UCam <sup>5</sup> GUF <sup>5</sup>		

## Scientific Integration:











# Individual DC projects:

Fellow: DC1	Host institution: TU Darmstadt	Apply to: kornelia.graefing@tu-darmstadt.de
	diversification and conformational control	
Objectives: (1) Develop ne and diversification; (2) Ver conformational design; (4) Methodology: (1) Synthesi scaffolds; (3) Explore diver selectivity and binding mod Expected Results: 1) Acce Identification of improved I Planned secondments (Ho (1) Abbvie (Germany), S. V (2) EPFL (Switzerland), C. (3) ETHZ (Switzerland), C. (3) ETHZ (Switzerland), S. Further information: http Fellow: DC2 Supervisor: <i>F. Hausch</i> Project Title: Synthesis an Objectives: (1) Explore po MetStab) and comparison t applicable building blocks; Methodology: (1) Stereose incorporation into FKBP-pr selectivity (FKBPs), intrace Expected Results: 1) Vers insights & improved unders Planned secondments (Ho (1) Analyticon Discoveries	w macrocyclization chemistry & post-massatile approaches to conformational contro Discovery of next-generation of improved ze and test pairs of mutually reactive build sity-generating reactions applicable to rap le and test for ternary complex formation a ss to new classes of macrocycles and tailo FKBP and cyclophilin ligands; (3) Identifi st institution, host mentor & secondment to <i>/uklevic</i> : ADME studies of synthesized ma <i>Heinis</i> : Adaption macrocycle generation <i>Riniker</i> : Computational Modelling of advo os://www.chemie.tu-darmstadt.de/hausch/g Host institution: TUDa Co-Supervisor: K. Schmitz (TUDa) d exploration of multifunctionalized linke lar linker side chains in MCs to improve k o simple linkers; (2) Develop a tool box for (3) Extension to macrocyclic molecular g lective generation of hydroxylated and flu referring scaffolds; (3) High-throughput m ellular binding (nanoBRET), and binding to atile method & tool set for macrocyclization st institution, host mentor & secondment to (Germany), L. Haustedt: Adaption to ext	Apply to:       kornelia.graefing@tu-darmstadt.de         Industry Mentor:       S. Ruf (Sanofi)         roup_members_hausch/join_us_hausch/jinty, via sembly of functionalized linkers and sets of broady
Planned secondments (Ho	st institution, host mentor & secondment t	opic):
<ul><li>(2) University of Cambrid</li><li>(3) Sanofi (Germany), <i>M. M.</i></li></ul>	ge (United Kingdom), D. Spring: Incorpo Mendez & S. Ruf: Combination with photo	ration of additional reactions /electrochemistry.
Further information: http Notes: Application Deadli		roup_members_hausch/join_us_hausch/index.en.jsp
Fellow: DC3	Host institution: Uppsala University	Apply to: <u>uu.varbi.com/en/what:job/jobID:739380/</u>
Supervisor: M. Erdelyi	Co-Supervisor: J. Kihlberg	Industry Mentor: S. Vucelić (Abbvie)
	(P3): Predicting and understanding MC ce	
<b>Objectives:</b> (1) Use machine learning to develop fast and accurate classification and regression models for prediction of the Caco-2 cell permeability of macrocycles; (2) Use and improve state-of-the-art machine learning technologies for identification of the most important descriptors contributing to the output from the models. <b>Methodology:</b> (1) Determine cell permeability of selected 1000 macrocycles across Caco-2 cell monolayers; (2) Use principle component analyses (PCA) and other machine learning methods to select the most informative 2D and 3D descriptors for model building; (3) Build models for cell permeability using different machine learning methods; (4) Use data analysis approaches such as partial least squares regression, principle component analyses, and clustering to provide insight into the effective set of descriptors governing the output from the models. (5) Determine to what extent selected macrocycles behave as		
molecular chameleons.	not models for an il dian of d 1	ability of MCs of diverse structures (2) 1
		eability of MCs of diverse structures; (2); machine llow facile use to design macrocycles; (3) Dissemination
<b>Planned secondment(s):</b> (1) <b>ABBV</b> , <i>S. Vukelic/C. Hoft</i> : Determination of cell permeability across Caco-2 cells; (2) UniTO, <i>G. Caron</i> : Characterize MCs chromatographically to assess lipophilicity, polarity & chameleonicity.		
Fellow: DC4	Host institution: Uppsala University	Apply to: <u>uu.varbi.com/en/what:job/jobID:739380/</u>
Supervisor: M. Erdelyi	<b>Co-Supervisor:</b> J. Kihlberg	Industry Mentor: W. Czechtizky
Project Title (related to W	(P4): NMR characterization of macrocycl	e conformational dynamics in solution at atomic level
<b>Objectives:</b> Development of of (1) orientational (residua paramagnetic relaxation en Development of a paramag <b>Methodology</b> : (1) Determi	of a solution NMR algorithm for the atomi l dipolar coupling, scalar coupling) and di hancement) restrains; (2) Identification of netic NMR method for determination of b	c level description of solution ensembles by combined use stance (nuclear Overhauser effect, pseudocontact shift, the conformational ensemble of macrocycles; (3)
(alignment media or param		paramagnetic relaxation rates (paramagnetic tagging); (2)







Computational generation of a theoretical conformational pool; (3) Order matrix analysis by singular value decomposition, identification of solution conformers and their populations.

Expected Results: (1) Development of algorithms to handle a series of isotropic and anisotropic NMR observables simultaneously; (2) Improvement of NMR-based characterization of small and midsize macrocycles' structure and dynamics; (3) Understanding of selected macrocycles' conformation in solution, in membranes and in their protein binding sites.
 Planned secondment(s): (1) ABBV, *S. Vuklevic/C. Hoft*: Determination of cell permeability across Caco-2 cells; (2) ETHZ *S. Riniker*: Computational generation of conformational ensembles, method development.

Fellow: DC5	Host institution: University of Torino	Apply to: via web link below
Supervisor: G. Caron	Co-Supervisor: G. Ermondi	Industry Mentor: S. Kostrun (Selcia)
	<b>3&amp;4):</b> Physicochemical profiling of macroo	
		nicity and solubility descriptors to characterize the
		nd local lipophilicity predictors and local polarity,
chameleonicity and solubili	ity models; (3) In depth analysis of chamele	onicity data for drug design purposes.
Methodology: (1) Potentio	metry; (2) Chromatographic methods based	on different stationary/mobile phases combination; (3)
Shake-flask to measure the	rmodynamic solubility; (4) Machine learnin	g to build models and (5) molecular modeling tools for
the implementation of chan		
		(2) Setting-up of lipophilicity calculators; (3) Polarity, eleonicity in drug design; (all specific for MCs
		al studies by NMR to highlight chameleonic behavior;
		<b>Selvita</b> , S. Kostrun, M31-33: intracellular/lysosomal
	s high-throughput imaging assay	civita, 5. Rostiuli, 1951-55. Intracential/1980solilar
	s://www.cassmedchem.unito.it/en/content/n	nc/dd
Notes: Application Deadlin		
Notes: Application Deadlin	10 30.09.2024	
Eallarry DC(	Heat in stitutions Hainsanites of Terring	A marter for the mark limb hole and
Fellow: DC6	Host institution: University of Torino	Apply to: via web link below
Supervisor: G. Caron	Co-Supervisor: G. Ermondi	Industry Mentor: C. Hoft (Abbvie)
	<b>(P4):</b> Identification of druggable regions in	
		of descriptors; (2) Locate the different classes of MCs
		gions using descriptors also obtained in WP3 and WP4;
		ploration and b) regions to obtain new MCs with a
favorable drug-like profile.		
		nical space and (2) infographic tools to generate
	e with the industrial partners.	
Expected Results: (1) MC	C chemical space; (2) Indication for future N	IC synthetic efforts
Planned secondment(s): ()	1) ABBV, C. Hoft, M16-18: High throughp	ut cell permeability measurements; (2) ETHZ, S.
Riniker, M36-38: In silico n	nodelling of selected macrocycles	
Further information: http	s://www.cassmedchem.unito.it/en/content/n	nc4dd
Notes: Application Deadlin	ne 30.09.2024	
Fellow: DC7	Host institution: Uppsala University	Apply to: <u>uu.varbi.com/en/what:job/jobID:739380/</u>
Supervisor: M. Erdelyi	Co-Supervisor: J. Kihlberg	Industry Mentor: NJ. Hempel (NovoNordisk)
		cell permeable large macrocyclic peptides (MCPs)
		e (MW >1000 Da) macrocylic peptides based on a potent
		nd target binding; (2) Determination of the solution
		as in a membrane-like, low dielectric environment. A
		e studied; (3) Determination of the solution ensembles of
	ptide in aqueous and membrane-like enviror	
		esis; (2) Get experience of the determination of cell
		(3) Determination of NOE build-up rates, scalar
		act shifts (alignment media or paramagnetic tagging); (4)
Computational generation of	of a theoretical conformational pool; (5) Ide	ntification of solution conformers and their populations,
using NMR data and the co		
		s conformation in aqueous solution, in membranes and in
		ation of large macrocyclic peptides' structure and
		ptides cross cell membranes, and what structural features
that are important.	0 0 F	. ,
	1) Novo Nordisk. NJ. Hemnel: Synthesis	and characterization of macrocyclic pentides (2) ETHZ
Planned secondment(s): (		and characterization of macrocyclic peptides (2) <b>ETHZ</b> es studied by NMR spectroscopy.
Planned secondment(s): (	1) <b>Novo Nordisk</b> , NJ. Hempel: Synthesis as for one or two of the macrocyclic peptide	
Planned secondment(s): ( S. Riniker,: MD simulation	is for one or two of the macrocyclic peptide	es studied by NMR spectroscopy.
Planned secondment(s): (	s for one or two of the macrocyclic peptide Host institution: UORSY/Enamine	
Planned secondment(s): ( S. Riniker,: MD simulation	is for one or two of the macrocyclic peptide Host institution: UORSY/Enamine Germany	es studied by NMR spectroscopy.



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**Project Title (related to WP1):** Design and producing of building blocks for the synthesis of macrocycles using non-amidic disconnections

**Objectives:** (1) design several sets of building blocks for the synthesis of macrocycles based on non-amidic disconnections (Grubbs reaction etc); (2) elaborate efficient cost-effective procedures for designed building blocks producing and determine their scope and limitation; (3) generate the database of building blocks based on common starting materials and elaborated procedures and synthesize the necessary examples in 1-10 gram amounts for further combinatorial chemistry uses.

**Methodology**: (1) The design will perform using our in-hose experience in the synthesis of polyfunctional small molecules with several orthogonal or protected functionalities; (2) The elaboration of the methodologies and determination of their scope and limitations will provide using various synthetic chemistry technics depends on their efficacy for the appropriate scheme with further functional group protection/deprotection/interconversion; (3) Based on a predicted structural features and elaborated procedures for combinatorial synthesis the set of building blocks will be synthesized in 1-10 g amount.

**Expected Results:** (1) Available sets of target compounds appropriated for project needs; (2) Available a 10-20 methodologies with determined scope for cherry picking and generation of final BB set; (3) 100-200 Compounds for combinatorial libraries' synthesis are produced and ready for further utilization.

**Planned secondment(s):** (1) TUDa, F. Hausch, M16-21: Incorporation of building blocks into FKBP- and Cyp-focussed libraries; (2) LU, S. Pomplun, M22-24: Adaption of high-throughput chemistry to ASMS library generation and deconvolution. **Further information:** This project is conducted in close collaboration between UORSY (Kiyv, Ukraine, <u>www.uorsy.com</u>) and Enamine Germany (Frankfurt, Germany, <u>www.enamine.de</u>). The research is performed at the Enamine-Germany site (Frankfurt).

Fellow: DC9	Host institution: Leiden University	Apply to: Link via https://pomplunlab.com/open-
		positions/
Supervisor: S. Pomplun	Co-Supervisor: C. Heinis	Industry Mentor: W. Czechtizky

**Project Title (related to WP1 ):** Ultrahigh throughput property screening of self-encoded MC libraries **Objectives:** (1) Generate a general synthesis and screening workflow for combinatorial macrocycle libraries (incl. tandem MSbased decoding strategy). (2) Produce a diverse set of tandem MS encoded MC libraries (10<sup>^3</sup> – 10<sup>^8</sup> members). (3) Combinatorial screen of building block related properties with the 1000-member libraries (cell penetration via PAMPA and proteolytic stability). (4) Affinity enrichment of binders for disease relevant RNA-binding proteins.

**Methodology**: (1) Establish solid phase synthesis and macrocyclization of the libraries. (2) Develop robust quality control assays for the in-solution libraries (3) Develop MSMS based decoding (combining nanoLC-MS/MS on an Orbitrap and *de novo* MS decoding software such as PEAKS or Sirius). (4) Establish property screening assays (e.g. PAMPA with 1000 compounds at a time) (5) Perform affinity selection against RNA binding properties and validate hit compounds

**Expected Results:** (1) Improved understanding which building blocks and chemistries in macrocycles lead to 'druglike' properties (2) High affinity binders for disease relevant RNA binding proteins and immunophilins

**Planned secondment(s):** (1) UCL, *A. Tabor*, M15-16: Multi-protection chemistry for library generation; **EPFL**, *C. Heinis*, M29-M30: MC chemistry for libraries; (2) **AZ**, *W. Czechtizky*, M31-M32: Test MCs on AZ-relevant RNA-protein targets.

Fellow: DC10	Host institution: Sanofi	Apply to: <a href="mailto:sven.ruf@sanofi.com">sven.ruf@sanofi.com</a>
Supervisor: S. Ruf	Co-Supervisor: F. Hausch	Industry Mentor: M. Mendez Perez (Sanofi)

**Project Title (related WP):** Late-stage functionalization methodologies for the synthesis and derivatization of MCs (WP2) **Objectives:** (1) Systematic exploration of C-H activation, photochemical and electrochemical methods for the synthesis of MC, especially the macrocyclization step; (2) Selective modification of aliphatic or heteroaromatic residues in MCs and introduction of side chains by late-stage functionalization; (3) Deliver matched molecular pairs of MCs based on variations in linkers to assess the impact on ADME properties, exemplified for FKBP and Cyp-based scaffolds.

**Methodology:** (1) Photoinduced decarboxylative radical addition; (2) Photo-redox Ni catalyzed C-N cross coupling; (3) Novel electrochemical reaction protocols; (4) Late-stage functionalization methodologies via electro-/photochemical approaches to introduce CF<sub>3</sub>-substituents; (5) Introduction of methyl groups, fluorine atoms and explore C-H activation by Rh-catalyzed additions of carbene residues; (6) In-silico analyses of conformational ensembles of the MCs.

**Expected Results:** (1) Novel late-stage functionalization toolbox of photo-/electrochemical reactions for the rapid construction and derivatization of MCs; (2) Improved macrocyclic inhibitors for FKBPs or Cyps; (3) Transfer of methods to other target classes.

 Planned secondment(s): (1) UoY, W. Unsworth, M11-12: Combination of photo/electrochemical synthesis with cascade-ring expansion approaches; (2) GUF, S. Knapp, M23-24: Application of photo/electrochemistry to macrocyclic kinase inhibitors; (3) TUDa, F. Hausch, M37-38: Testing and co-crystallization of synthesized MCs for binding to FKBPs & Cyps

 Further information:
 Sanofi: global healthcare and pharmaceutical company

Notes: Application Deadline 31.12.2024, mail application with subject header "MC4DD" to sven.ruf@sanofi.com

Fellow: DC11	Host institution: AstraZeneca	Apply to: to be provided, check for updates
Supervisor: W.	Co-Supervisor: J. Kihlberg	Industry Mentor: S. Schiesser (AstraZeneca)
Czechtizky		
Project Title (related to WP5): Expanding the druggable space of MCs to protein/RNA interactions – identify suitable		

protein/RNA interfaces and discover MCs which can bind and modulate disease-associated RNAs



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**Objectives:** (1) Discover and optimize macrocyclic inhibitors of disease-associated RNA/protein interactions. (2) Investigate binding mode and cellular potency of MCs.

**Methodology**: (1) Select RNAs with peptide/protein interfaces, where the protein/peptide can be mimicked by a MC; (2) *De novo* synthesis of MCs, VS within AZ and collaborator's MC library, or phage display selection of macrocyclic binders to selected RNA structures; (3) Validation of discovered binders via SPR/ITC/ASMS; (4) Investigation of MC/RNA binding site using NMR and/or X-ray crystallography; (5) Affinity optimization of macrocyclic hit to RNA; (6) Biologic effect assay(s) and benchmarking against state-of-the-art methodologies to modulate RNA biology.

**Expected Results:** (1) Learn how to analyze peptide/protein-RNA interfaces regarding binding motifs and pharmacophore patterns; (2) Understand if we can *de novo* design or experimentally identify MCs that mimic proteins or peptides in their binding to RNA; (3) Demonstrate binding and functional effect of macrocyclic binders to RNA; (4) Generate a better understanding if we can design macrocyclic modulators of RNA function.

**Planned secondment(s):** (1) **TUDa**, *F. Hausch*, M34-35: Affinity optimization of discovered MCs; (2) **UU**, *M. Erdelyi/J. Kihlberg*, M36-42: Elucidation of NMR structure of MC-RNA complexes & solution conformation of MCs.

Fellow: DC12	Host institution: Abbvie	Apply to: <u>https://careers.abbvie.com/de</u>
Supervisor: F. Pohlki         Co-Supervisor: F. Hausch(TUDa)         Industry Mentor: C. Hoft (Abbvie)		Industry Mentor: C. Hoft (Abbvie)
Project Title (related to WP3): Brain penetrant MCs for CNS Drug Discovery		

**Objectives:** (1) Systematically explore the effect of key structural and physicochemical parameters such as ring size, ring conformation, hydrogen bond donors and acceptor count and lipophilicity on brain penetration of macrocyclic model substrates; (2) Generate a comprehensive set of in silico and in vitro parameters for a set of MCs and selected acyclic analogues; (3) Investigate brain pharmacokinetics of selected compounds with appropriate properties.

**Methodology**: (1) Define a set of macrocyclic model substrates spanning a range of key structural and physicochemical parameters; (2) Characterize structural and physicochemical properties in silico; (3) Generate in vitro data on permeability and efflux substrate properties using cellular models: Caco2, MDCK (+MDR1 & BCRP efflux transporters); (4) for a selected subset of compounds, use mouse PK cassette studies to determine in vivo brain/plasma ratio, absolute brain and plasma compound levels, and determine unbound free fraction in brain homogenate to calculate Kp,uu,brain. (5) Determine the desired property space for brain penetrant macrocycles by analyzing relationships between endpoints determined above, e.g. SAR (relationship between structural or physicochemical properties and in vitro permeability / efflux or in vivo brain penetration), and in vitro – in vivo (relationship between in vitro permeability / efflux and and in vivo brain penetration)

**Expected Results:** (1) Improved understanding of structural and physicochemical parameters, as well as in vitro properties determining brain penetration of macrocyclic drugs; (2) dataset for efflux properties of MCs

**Planned secondment(s):** (1) **UniTO**, *G. Caron*, M16-18: Measurement of chameleonicity of brain-permeable MCs. (2) **TUDa**, *F. Hausch*, M31-33: SAR (structure activity relationship) analyses of brain-permeable MCs.

**Further information:** application through AbbVie's job portal: <u>https://careers.abbvie.com/de</u> (search for "student" and filter location to Ludwigshafen). We are looking for a student with biology, biochemistry or life sciences background

Fellow: DC13	Host institution: EPFL	Apply to: <u>christian.heinis@epfl.ch</u>
Supervisor: C. Heinis	Co-Supervisor: S. Pomplun	Industry Mentor: W. Czechtizky
Project Title (related to WP1): Synthesis and screening of large MC compounds libraries		

**Objectives:** (1) Identify MCs for challenging targets (e.g. FKBPs, Cyps, RNA). (2) Optimize obtained MCs into drug leads. **Methodology**: (1) Design and synthesis of target-tailored libraries comprising ten-thousands of MCs using an approach recently developed in the Heinis group (combinatorial synthesis of MCs at a nanomole-scale using acoustic reagent transfer, and screening of crude reactions); (2) Screening the MC libraries against relevant model targets using biochemical assays; (3) Characterize the MCs (affinity, selectivity, membrane permeability, cellular activity).

**Expected Results:** (1) Validation/improvement of methods for combinatorial synthesis of large MC compound libraries; (2) High affinity inhibitors of multiple protein-protein interaction targets; (3) Drug leads for important disease targets. **Planned secondment(s):** (1) **AZ**, *S. Schiesser*, M16-M18: Testing of macrocyclic libraries for RNA binding; (2) **TUDa**, *K. Schmitz*, M33-M36: Structural & biochemical characterization of macrocyclic libraries & hits for FKBPs and cyclophilins.

Fellow: DC14	Host institution: ETH Zürich,	Apply to: Link provided on <u>http://www.riniker.ethz.ch</u>
	Switzerland	
Supervisor: S. Riniker	<b>Co-Supervisor:</b> <i>M. Erdelyi (U. Uppsala)</i>	Industry Mentor: C. Kroll (Roche)
Project Title (related to WP4): Computational characterization of macrocycles conformations and connection to permeability		

#### Objectives:

(1) Characterization of the structure-permeability relationship of macrocycles using molecular dynamics simulations and comparison with NMR data

(2) Development of novel conformation- and environment-dependent 3D descriptors based on the MD simulations

(3) Development and refinement of machine learning models for permeability prediction for macrocycles

**Requirements**: Applicants should hold a Master degree in chemistry, computational chemistry, or physics. Experiences with machine learning and/or biomolecular simulation, and strong programming skills (Python) are highly advantageous.

#### Planned secondments:

(1) Roche, Switzerland, C. Kroll: Experimental determination of PAMPA permeability coefficients for macrocyclic library

(2) U. Uppsala, Sweden, *M. Erdelyi*: Conformational studies by NMR experiments

(3) Bayer, Germany, D. Barber: Conformational sampling of macrocycles



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Fellow: DC15	Host institution: Roche	Apply to: to be provided, check for updates		
Supervisor: C. Kroll	Co-Supervisor: S. Riniker (ETHZ)	Industry Mentor: S. Schadt (Roche)		
•	<b>P3&amp;4):</b> Refinement of <i>in silico</i> methods f	for permeability prediction of peptidic MCs through		
experimental validation				
		3; (2) Apply models and methods that aid optimization of		
		oughput of computational methods, e.g. by combining		
		rivileged properties; (4) Increase predictivity of		
	(e.g. 3D-PSA) to act as surrogates for actu			
		(e.g. from WP1) and assess permeability in assays such		
		ability; (2) Combine obtained data with short MD		
		lity prediction; (3) Iteratively improve predictions and		
	) Generate novel molecules based on newl			
		biological activity, ideally derive generalizable design		
principles that hold across s				
		nation-based computational methods. (2) UU, J. Kihlberg		
& M. Erdelyi, M25-27: Det	fine design principles & experimental analy	ysis of conformation.		
	1			
Fellow: DC16	Host institution: University of York	Apply to: william.unsworth@york.ac.uk		
Supervisor: W. Unsworth	Co-Supervisor: A. Tabor	Industry Mentor: S. Vuklelić (Abbvie)		
	<b>(P2):</b> High throughput synthesis and scree			
	ascade ring expansion methods to prepare			
		ional groups to prepare macrocycles containing, e.g.		
		f cascade ring expansion conditions with focus on broad		
		assembly on substrates and MCs; (4) Generation of		
	5) Optimization after property (WP3 & 4)			
		rsification; (2) Library of high-quality MCs as a resource		
	ngs; (3) Novel cell-permeable MCs with in			
		measurements for cascade ring-expanded MCs; (2)		
		stry into cascade ring expansion syntheses.		
	os://unsworthlab.weebly.com/			
https://jobs.york.ac.uk/vacancy/marie-curie-early-stage-researcher-chemistry-563469.html				
Fellow: DC17	Host institution: University College	Apply to: <u>a.b.tabor@ucl.ac.uk</u>		
	London			
Supervisor: A. Tabor	Co-Supervisor: W. Unsworth	Industry Mentor: S. Güssregen (Sanofi)		
		d on privileged scaffolds for cell permeability		
		alysis of low-peptide, cyclophilin-targeted MCs		
Methodology: (1) Synthesis of orthogonally protected building blocks containing thioether linkages to give Cyp-targeted				
scaffolds by solid phase macrocyclization; (2) synthesis of precursors with functional groups to give sangliferin-inspired Cyp-				
targeted scaffolds; (3) solid-phase synthesis of scaffolded MC displaying tripeptide and depsipeptide sequences, including Cyp-				
preferring piperazic acid and $\Psi$ Pro; (4) generation and screening of diverse scaffolded MC libraries.				
<b>Expected Results:</b> (1) novel, versatile MC libraries combining privileged non-peptidic scaffolds with minimal peptide display;				
(2) improved Cyp ligands (resource for WP5); (3) datasets with structural and ADME data for WP3.				

**Planned secondment(s):** (1) **UU**, *M. Erdelyi*, MD simulations & NMR characterization of selected MC, M23-M24; (2) **Sanofi**, *S. Güssregen*, Modelling of constraint MCs, M29-M30; (3) **LU**, *S. Pomplun*, Scaffold-MCs for ASMS, M31-M32.

