



UPPSALA  
UNIVERSITET

# Annual Report 2015

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Department of Pharmaceutical  
Biosciences

Fastställd av styrelsen 2016-05-24

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The scientific reports were written by the different research groups at the department.

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## Introduction

This annual report highlights the research activities in the Department of Pharmaceutical Biosciences during 2015. The research areas cover many different areas, including research on basic mechanisms of drug action, drug dependency and adverse health effects, as well as studies on drug metabolism, pharmacokinetics and pharmacodynamics. In addition, there is also a significant focus on pharmaceutical bioinformatics and proteomics, as well as pharmacometric modeling and simulations used in drug development. Some research groups are developing tools directly applicable to the pharmaceutical industry, while others are studying issues such as drug dependency and environmental contaminants with important socioeconomically implications. A special commitment was made during 2015 to build up an in vitro platform by investing in a new HCA/HCR equipment (Image Xpress from Molecular Devices) for high content analysis/screening. This in vitro platform will be used by several research groups at the department in different types of projects studying different types of end-points in cellular systems (including tissues). This report confirms that the research groups continue to have national and international collaborations within their research areas. Another important activity is the teaching we promote, and most of our junior lecturers, senior lecturers, and professors, and also some of our researchers, are heavily engaged in our education on various programs (focusing on pharmacy students). Most of our education is research-based and it is our aim to ensure that there is a synergy between teaching and research. Besides providing professional pharmacy education, the department also offers graduate pharmacy students specialization in clinical pharmacy.

### *Funding*

All research activities require funding from national and international research councils, pharmaceutical companies, and the government. For example, in 2015 major grants were obtained from various national and international research councils such as the Swedish Foundation for Strategic Research (VR and FORMAS), EU Innovative Medicines Initiative, Research and Innovation for Sustainable Growth (VINNOVA), the Swedish Association of the Pharmaceutical Industry (LIF), Kjell and Märta Beijers Foundation, Carl Tryggers Foundation, Science for Life Laboratory (SciLife), Swedish Council for Working Life and Social Research, Swedish Insitute, Uppsala University, the Alcohol Research Council of the Swedish Alcohol Retailing Monopoly, Berzeli Centre for Biotechnological Research, the Research Council of Swedish Criminal Care, the Swedish Governmental Agency for Innovation Systems, Agricultural Sciences and Spatial Planning, Genentech, EU FP7 Health, Innovative Medicines Initiative (IMI), Swedish Cancer Society, the National Board of Health and Welfare, National Institute of Health (NIH), AFA Försäkring, Svenska Spels forskningsråd, Hjärnfonden, eSSENCE, SeRC, Japanese Society for Promotion of Science, Fredrik och Ingrid Thulins stiftelse, and the Facias Foundation. Many projects also have financial support from pharmaceutical companies such as AstraZeneca, Janssen Pharmaceuticals, Novartis, Pfizer and Roche.

### *Organisation and financial review*

The department is organised on the basis of our main core activities, i.e., research, teaching and the collected support activities (i.e., management, finance/staff administration, education administration and infrastructure). All support activities are funded by a percentile overhead on salaries and operating costs. The government provides the basic funding for teaching and research. The peer-reviewed scientific publications, PhD examinations and the external grants provide the foundation for any decisions concerning the allocation of university funding of research. The revenues from research councils, EU, foundations and pharmaceutical companies are very important to increase the research activities at the department. Below follows a short summary of incomes and expenditures 2015.

<i>Major incomes 2015</i>	<i>(kSEK)</i>
Research and graduate education- government	34 153
Research grants – research councils	38 369
Research – commissioned	9 608
Education – basic and advanced level –government	41 901

<i>Major expenditures 2015</i>	<i>(kSEK)</i>
Staff costs	68 554
Operating expenses etc.	12 599
Premises	14 015
University/faculty support activities	13 909
Library	2 730
Depreciation	4 049
Travels	3 301

### *Core facilities and other engagements*

Apart from being involved in regular research and teaching activities, our department also host two major core facilities, NRMSI and UUBF. We are also involved in the platform UPPNEX hosted by the SciLifeLab. The department is also engaged in “the third mission”, mainly by hosting and leading the activities of U-FOLD. In addition, the department of Pharmaceutical Biosciences is also one of the participants in the EU-supported education project SafeSciMet, involving both the Academy and Pharmaceutical Industry in several European countries, and we are also one of the participants in SweTox, a national resource centre for academic interdisciplinary collaboration within toxicology sciences.

The National Resource for Mass Spectrometry Imaging (NRMSI) was founded at the Uppsala University in 2010 with funds provided by the Swedish Research Council-Research Infrastructure in 2010 and Uppsala University-Infrastructure in 2013. The mission of the resource (previously named NCMSI) is to accelerate the adoption of

MALDI (matrix-assisted laser desorption ionization) and DESI (desorption electrospray ionization)-MSI technology in the biological and medical research community. NRMSI provides access to advanced technologies and promotes interactions between scientists who are experts in these technologies and researchers conducting focused biological investigations. Such collaborations provide new insights into the targeted biological problems and a better understanding of health and disease at the molecular level. Another important activity of the NRMSI is training, informing scientists about the capabilities and limitations of the technology, good practices, and to established methods and protocols. NRMSI uses state-of-the-art mass spectrometry technologies and cutting edge research to help projects in e.g., drug discovery and development, neuroscience, oncology and pathology applications. NRMSI is located at Dr. Per Andrén's Medical Mass Spectrometry research group at the Dept. of Pharmaceutical Biosciences. The NRMSI was 2015 awarded an Infrastructure Research Fellow grant from the Swedish Foundation for Strategic Research for 2016-2020.

Uppsala University Behavioral Facility (UUBF) was established in 2011 through strategic funding from the Faculty of Medicine and Pharmacy. UUBF functions as a non-profit core facility that administers and organizes experimental behavioral testing of mice, rats and fish with the aim to optimize access to well validated tests for research groups at Uppsala University as well as for external researchers, and promote the further development of high quality behavioural tests. Services offered include experimental design, statistical processing and interpretation of results, rental of equipment including advanced recording equipment, and full performance of experiments. During the last years the number of users has increased from about 15 to 60 users distributed over a number of research groups and a recent survey shows that our users value the services offered.

E-mail: [uubf@farmbio.uu.se](mailto:uubf@farmbio.uu.se). Homepage: <http://www.farmbio.uu.se/Corefacility/uubf/>

The Bioinformatics Compute and Storage facility (UPPNEX) at Science for Life Laboratory (SciLifeLab) is headed from the Department of Pharmaceutical Biosciences. The facility provides high-performance computing and storage resources, maintain relevant bioinformatics software and data (e.g. reference genomes), and offer educational courses and user support. The facility had over 800 active projects during 2015 with computational resources hosted at UPPMAX - Uppsala University's resource for high-performance computing and a node in the Swedish National Infrastructure for Computing (SNIC). The facility had 3 FTE during 2015 and is funded by SciLifeLab and NBIS. The primary contact person at our department is: Ola Spjuth, Head of the Bioinformatics Compute and Storage facility and Co-Director of UPPMAX. E-mail: [ola.spjuth@farmbio.uu.se](mailto:ola.spjuth@farmbio.uu.se) Homepage: <http://www.scilifelab.se/facilities/uppnex/>.

U-FOLD is a forum for research on addiction to medical products and illegal drugs. This network is the link between research, prevention and treatment activities in Uppsala regarding different types of addiction. The network brings broad experience and different perspectives and the major aim of the activities within U-FOLD is to contribute to greater understanding and a common approach regarding the underlying chain of events leading to addiction. Within the network U-FOLD can create new methods to prevent drug addiction, including preventive measures, care and legal

actions. U-FOLD gather today twenty participants from academia, government, and nonprofit organizations. During 2015, the project on children in abusive environments initiated by Regionförbundet and U-FOLD, and the study on prevention strategies against pathological gambling supported by the Public Health Agency, Sweden were continued. During the year, a number of meetings and seminars were arranged by U-FOLD, most of them focused on hot topics related to the area of addiction. More information (still only in Swedish) can be found on <http://www.ufold.uu.se>

SafeSciMET is a European education and training network, developing and establishing a comprehensive education program in safety sciences for medicines. Teachers at 15 European universities and experts from the pharmaceutical industry and regulatory authorities have together developed 20 courses in the area. The courses are especially designed to support scientists and professionals in medicines research and development and address the needs in small and large pharmaceutical companies, regulatory authorities, academic institutes and health care. Professor Eva Brittebo is responsible for the SafeSciMET Student Office. A novel attractive website (<http://www.safescimet.eu/>) was launched for all kind of devices in the beginning of 2015. The focus of the new website is to give an detailed presentation of the courses for prospective students and staff from pharmaceutical companies.

The Swedish toxicology sciences research center (Swetox) is based on a consortium of eleven Swedish universities. Swetox was established as a national resource for academic interdisciplinary collaboration within toxicology sciences in 2013 and it promotes research, education and societal interactions related to chemicals, health and environment. A Swetox hub at Södertälje has been established as a research centre in modern laboratory facilities for safety assessment of pharmaceuticals. These facilities were closed-down by AstraZeneca in 2012. A vice chancellor council and a board govern the Swetox consortium. The board consists of representatives from all consortium universities. Professor Eva Brittebo is the representative from Uppsala University.

### *Future development*

The years ahead promise many changes in terms of research and education and one can only hope that we will be successful in our attempts to get our different types of activities financially supported by both governmental funding of undergraduate teaching and research, as well as external fundings, so that we can keep the high level of research and teaching also in the future. What we do know is that a couple of professors, teachers and members of our technical and administrative staff retired during 2014/2015, and that more staff members will retire during 2016. It is indeed a real challenge to find equally good replacements at all positions, but at the same time, it will also provide many possibilities for renewal.

A new professor in pharmacology (Robert Fredriksson) was recruited during 2015. Robert, who came from Department of Neurosciences at the Medical faculty Uppsala University, will start up his group and activities at our department in the beginning of January 2016. A new professor under recruitment is one in pharmaceutical cell biology,

which hopefully will be at place during 2016. One senior lecturer/associate professor in pharmacometrics (Lena Friberg) was promoted to professor during 2015, and our hope is that there will be more senior lecturers that will be promoted to professors during 2016. During the year we also employed three new senior lecturers at the department (Maria Kjellsson, Jörgen Bengtsson and Maria Norlin), and at least two new positions as senior lecturers will be announced during 2016. Seven PhD students at the department defended their thesis during 2015 (Jonathan Alvarsson, Anna Jonsson, Anders Kristoffersson, Brigitte Lacroix, Shima Momeni, Sara Palm and Anneli Wennman).

Uppsala May 24, 2016

*Björn Hellman*

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## Organisation 2015

### *Chairman*

Björn Hellman

### *Deputy chairman*

Mats Karlsson

### *Department board*

Björn Hellman, *chairman*

Mats Karlsson, *teacher representative*

Margareta Hammarlund-Udenaes, *teacher representative*

Maria Kjellsson, *teacher representative*

Siv Jönsson, *teacher representative*

Jörgen Bengtsson, *teacher representative*

Mathias Hallberg, *teacher representative*

Lena Bergström, *teacher representative, deputy*

Karin Tjäder, *technical/administrative representative*

Elisabeth Jonsson *technical/administrative representative*

Patrik Källback, *graduate student representative*

Ida Netterberg, *graduate student representative, deputy*

Björn Clausson, *student representative*

Desirée Legeth, *student representative, deputy*

Marina Rönngren, *personnel coordinator, adjunct*

Agneta Hortlund, *economical coordinator, adjunct*

Mikaela Andersson, *secretary, adjunct* (from August 2015)

Stina Silander, *secretary, adjunct* (until July 2015)

### *Professors*

Georgy Bakalkin (until May 2015)

Eva Brittebo

Lena Friberg (from July 2015)

Margareta Hammarlund-Udenaes

Björn Hellman

Mats Karlsson

Ingrid Nylander

Ernst Oliw (until January 2015)

Jarl Wikberg

Kjell Wikvall

### *Professor emeriti*

Lennart Paalzow

Lennart Dencker

### *Senior Professors*

Georgy Bakalkin (from June 2015)

Fred Nyberg

Ernst Oliw (from February 2015)

### *Adjunct Professors*

Jan Kehr

*Senior lecturers*

Per Andrén, *Associate professor*  
Jörgen Bengtsson  
Lena Bergström, *Associate professor*  
Agneta Freijs  
Mathias Hallberg, *Associate professor*  
Ronnie Hansson (until July 2015)  
Andrew Hooker  
Oskar Karlsson\*, *Associate professor*  
Maria Kjellsson, *Associate professor*  
Elisabet Nielsen  
Maria Norlin  
Ulrika Simonsson, *Associate professor*  
Anne-Lie Svensson  
Erika Roman, *Associate professor*

*Lecturer*

Ola Spjuth, *Associate professor*

*Adjunct Lecturer*

Markus Friden

*Junior lecturers*

Ann-Marie Falk  
Lena Klarén  
Emelie Lefvert\*  
Viktoria Lind  
Emma Lundkvist  
Samra Srdic\*  
Maria Swartling

*Directors of undergraduate studies*

Lena Bergström  
Jörgen Bengtsson  
Eva Brittebo  
Ann-Marie Falk  
Mathias Hallberg  
Björn Hellman  
Elisabet Nielsen  
Anne-lie Svensson  
Maria Swartling  
Jarl Wikberg  
Kjell Wikvall

*Postdocs, Researchers and PhD students*

Listed in the scientific reports

*Laboratory staff*

Jessica Dunhall  
Britt Jansson (until May 2015)  
Lena Norgren

*Administration and service*

Mikaela Andersson  
Ulrica Bergström  
Magnus Efverström  
Martin Grenzeliuss (from November 2015)  
Agneta Hortlund  
Elisabeth Jonsson  
Marina Rönngren  
Stina Silander\*  
Johanna Svensson  
Karin Tjäder

*Safety officers*

Magnus Efverström (from August 2015)  
Ronnie Hansson (until July 2015)  
Lena Norgren  
Marina Rönngren  
Henrik Wadensten  
Sviatlana Yahorava  
Kjell Åkerlund (until July 2015)

*The work environment group*

Björn Hellman, *chairman*  
Mikaela Andersson, *secretary*  
Jessica Dunhall  
Patrik Källback  
Lena Norgren  
Ernst Oliw  
Marina Rönngren  
Karin Tjäder

*Working group on post-graduate studies*

Margareta Hammarlund-Udenaes, *chairman*  
Ari Brekkan  
Maria Norlin  
Marina Rönngren, *secretary*

*Working group on equal opportunities*

Elin Svensson, *chairman and gender equality representative*  
Mikaela Andersson  
Mattias Hallberg  
Marina Rönngren\*  
Ronnie Hansson (until July 2015), *adjunct*  
Sebastian Axelsson

*\*Temporary position*

# Scientific Reports 2015

## Biochemical Pharmacology

Ernst H. Oliw

<http://farmbio.uu.se/research/researchgroups/biokemfarm/>

Arachidonic acid and a few other polyunsaturated fatty acids are bioactivated in humans by enzymatic oxygenation to prostaglandins, leukotrienes, epoxides (EETs) and other local hormones, which contribute to fever, pain, inflammation and cancer development, and to regulation of physiological processes. Common drugs such as aspirin, acetaminophen (paracetamol, APAP) and ibuprofen inhibit biosynthesis of prostaglandins and reduce symptoms of disease, but may also cause side effects related to their actions. Other drugs are based on leukotriene receptor antagonists (e.g., montelukast), which are used for treatment of bronchial asthma.

Bioactivation of polyunsaturated fatty acids also occurs in plants and fungi where oxygenation of linoleic and linolenic acids is important for the plant-pathogen interaction and for fungal reproduction and pathogenicity. The goal of our research is to investigate the mechanism of oxygenation and bioactivation of fatty acids and to determine their biological function.

We investigate mainly three groups of enzymes: (i) cytochromes P450, (ii) lipoxygenases, and (iii) DOX-CYP fusion proteins with dioxygenase (DOX) and cytochrome P450 (CYP) domains. These enzyme classes occur in man but also in important fungal pathogens, e.g., *Aspergillus fumigatus* causing farmer's lung disease and *Magnaporthe oryzae*, causing rice blast disease and destruction of 25% of the rice crop of Japan. Our goal is to understand how the enzymes work in order to understand their physiological functions.

(i) DOX-CYP fusion proteins: The DOX-CYP family consists of at least six subfamilies. LDS oxidize oleic and linoleic acids to a series of vicinal diols (e.g., 5,8-dihydroxy-, 7,8-dihydroxyoctadecadienoic acids) and hydroperoxides (e.g., 8-hydroperoxy- and 10-hydroperoxylinoleic acid), which likely function as sporulation hormones. Recent discoveries are the 8*R*- and 8*S*-dioxygenase and allene oxide synthase of human and plant pathogens causing Valley fever and septoria tritici blotch.

(ii) Lipoxygenases: All lipoxygenases contain a catalytic metal, iron in humans and plants. We focus our basic research on the first described manganese-lipoxygenases, which are important for *Gäumannomyces graminis*, an important pathogen of wheat, and its structurally similar lipoxygenases of *Magnaporthe salvinii* and *M. oryzae*, and the catalytic convergence of iron- and manganese-lipoxygenases by a single amino acid substitution. We have now determined the 3D structures of manganese lipoxygenase from *M. oryzae* and from *G. graminis*, and these are the only known structures of lipoxygenases with catalytic manganese.

### Members of the group during 2015

Ernst H. Oliw, MD, PhD, Senior professor

Anneli Wennman, PhD student

Linda Sooman, scientist

Yang Chen, scientist

### Publications 2013 – 2015

1. Hoffmann, I., Jerneren, F., and Oliw, E. H. (2013) Expression of fusion proteins of *Aspergillus terreus* reveals a novel allene oxide synthase. *J. Biol. Chem.* 288, 11459-11469
2. Hoffmann, I., and Oliw, E. H. (2013) 7,8- and 5,8-Linoleate diol synthases support the heterolytic scission of oxygen-oxygen bonds by different amide residues. *Arch. Biochem. Biophys.* 539, 87-91
3. Hoffmann, I., and Oliw, E. H. (2013) Discovery of a linoleate 9S-dioxygenase and an allene oxide synthase in a fusion protein of *Fusarium oxysporum*. *J. Lipid Res.* 54, 3471-3480
4. Wennman, A., and Oliw, E. H. (2013) Secretion of two novel enzymes, manganese 9S-lipoxygenase and epoxy alcohol synthase, by the rice pathogen *Magnaporthe salvinii*. *J. Lipid Res.* 54, 762-775
9. Hoffmann, I., and Oliw, E. H. (2013) Discovery of a linoleate 9S-dioxygenase and an allene oxide synthase in a fusion protein of *Fusarium oxysporum*. *J. Lipid Res.* 54, 3471-3480
10. Wennman, A., Oliw, E. H., and Karkehabadi, S. (2014) Crystallization and a preliminary crystallographic analysis of manganese-lipoxygenase. *Acta Crystallogr. F* 70, 522-525.
11. Wennman, A., Karkehabadi, S., and Oliw, E. H. (2014) Kinetic investigation of the rate-limiting step of manganese- and iron-lipoxygenases. *Arch. Biochem. Biophys.* 555-556, 9-15
12. Oliw, E. H., and Wennman, A. (2014) Chiral phase-HPLC separation of hydroperoxyoctadecenoic acids and their biosynthesis by fatty acid dioxygenases. *Methods Mol. Biol.* 1208, 85-95
13. Oliw, E. H., and Wennman, A. (2015) Chiral phase-HPLC separation of hydroperoxyoctadecenoic acids and their biosynthesis by fatty acid dioxygenases. *Methods Mol. Biol.* 1208, 85-95
14. Sooman, L., and Oliw, E. H. (2015) Discovery of a novel linoleate dioxygenase of *Fusarium oxysporum* and linoleate diol synthase of *Colletotrichum graminicola*. *Lipids* 50, 1243-1252
15. Wennman, A., Jernerén, F., Magnuson, A., and Oliw, E. H. (2015) Expression and characterization of manganese lipoxygenase of the rice blast fungus reveals prominent sequential lipoxygenation of alpha-linolenic acid. *Arch. Biochem. Biophys.* 583, 87-95
16. Wennman, A., Magnuson, A., Hamberg, M., and Oliw, E. H. (2015) Manganese lipoxygenase of *Fusarium oxysporum* and the structural basis for biosynthesis of distinct 11-hydroperoxy stereoisomers. *J. Lipid Res.* 56, 1606-1615
17. Sooman, L., Wennman, A., Hamberg, M., Hoffmann, I., and Oliw, E. H. (2016) Replacement of two amino acids of 9R-dioxygenase-allene oxide synthase of *Aspergillus niger* inverts the chirality of the hydroperoxide and the allene oxide. *Biochim. Biophys. Acta* e-pub 11 Nov

### Dissertations 2015

1. Anneli Wennman. *The structural basis of the catalytic specificity of manganese lipoxygenases. 3D structure of analysis of the lipoxygenase of Magnaporthe oryzae.* 2015. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, 204. ISSN 1651-6192.

### Agencies that supported the work/Funding 2015

The Swedish Research Council Medicine.

## Projects

(I) *Characterization of heme-containing fatty acid dioxygenases and P450 fusion enzymes of human and plant pathogens (Yang Chen, Fredrik Jernerén, Ernst Oliw)*

Fungi are severe pathogens of man and can be devastating for important crops. *Aspergillus* causes farmer's lung disease and invasive aspergillosis of immunocompromized patients. Rice blast disease is caused by *Magnaporthe oryzae*, and destroys ~25% of rice crops worldwide. *Aspergillus* and *M. oryzae* contain cyclooxygenase-related enzymes, diol synthases and dioxygenases, which transform linoleic acid into hydroperoxides and dihydroxy fatty acids. Our aim is to characterize the enzymes by enzyme expression, gene targeting and by studies on their biological importance.

(i) Characterization of 8R- and 8S-dioxygenases linked to allene oxide synthases of the human pathogen *Coccidioides immitis* and the plant pathogen *Zymospetoria tritici*. We are now investigating the reaction mechanisms and the mechanisms of biosynthesis of allene oxides.

(ii) Characterization of all unique DOX-CYP fusion enzymes of the 10 top pathogens in molecular biology. Three have already been characterized and we now focus on *Z. tritici* and the mechanism of biosynthesis of jasmonic acid.

(iii) EPR analysis of the protein radicals of CYP-DOX fusion enzymes. The Tyr residues of recombinant dioxygenases with a catalytic Tyr radical will be labeled with deuterium to conclusively identify that a Tyr radical is formed during catalysis.

(II) *Characterization of the 3D structures of manganese-lipoxygenases with substrate (Yang Chen and Ernst Oliw)*

Our aim is to study the reaction mechanisms of lipoxygenases by determination of the the 3D structure with linoleic acid in the active site. We will focus on the novel lipoxygenases of *M. oryzae*. The latter forms crystals in a reproducible way.

## Drug Safety and Toxicology

Eva Brittebo

<http://farmbio.uu.se/research/researchgroups/dst/>

This research group is focusing on mechanisms of adverse effects of environmental contaminants as well as pharmaceuticals and traditionally used medicinal plants, by the use of various *in vitro* and *in vivo* models. The major objective of our studies is to increase the knowledge about mechanisms of actions of toxicants in order to improve predictions of hazards and risks for humans. The studies are presently focusing on the following research areas: Neurotoxicology, Genetic toxicology, and Developmental toxicology.

The neurotoxicity studies aim to elucidate mechanisms of long-term cognitive impairments and neurodegeneration in the adult brain following neonatal exposure to neurotoxins. The concept of early life stage exposure and developmental toxicity manifested late in life has become a basal paradigm in current developmental toxicology studies. Another neurotoxicology project focus on a troublesome adverse effect (dyskinesia) of L-DOPA pharmacotherapy for patients with Parkinson disease. These studies are based on several molecular techniques including MALDI –TOF imaging mass spectrometry (MALDI IMS) for the topographical elucidation of proteins, neuropeptides and neurotransmitters and their changing concentrations in the brain during physiological and pathophysiological events. In addition, the direct delivery of therapeutic agents to the brain via intranasal administration is being examined.

The studies on genetic toxicology are based on the Comet Assay for evaluating DNA-strand breaks in individual cells. An important issue in the safety evaluation of drugs is if there are threshold doses or not for agents found to be genotoxic *in vitro* or *in vivo*.

The developmental toxicology is one the most challenging areas in predictive toxicology as the developmental period of organisms is much more sensitive to toxicants than the adult period. Some of these studies are directed towards the development of *in vitro* system for prenatal cardiotoxicity. Other studies aim to elucidate mechanisms of developmental neurotoxicity during the postnatal synaptogenic sensitivity period.

### Members of the group during 2015

Eva Brittebo, Professor

Björn Hellman, Professor

Lennart Dencker, Professor emeritus

Malin Andersson, Associate Professor

Oskar Karlsson, PhD, Associate Professor (20 %)

Michael Stigson, PhD, Associate Professor, Researcher

Mats Nilsson, PhD Researcher

Birger Scholz, PhD, Researcher

Lisa Ersson, PhD student

Daiane Cattani, PhD student (exchange program from Brazil)

Elena Piras, PhD student (on parental leave)

Jasem A Shemali, PhD student

Hans Lindén, senior project coordinator

Lena Norgren, Technician

Linnea Blomberg, Research assistant

Emelie Karlsson, Research assistant

## Publications 2013 – 2015

1. Karlsson O, Jiang L, Ersson L, Malmström T, Ilag LL, Brittebo EB. (2015) Environmental neurotoxin interaction with proteins: Dose-dependent increase of free and protein-associated BMAA ( $\beta$ -N-methylamino-L-alanine) in neonatal rat brain. *Sci Rep.* 5:15570.
2. Karlsson O, Berg AL, Hanrieder J, Arnerup G, Lindström AK, Brittebo EB. Intracellular fibril formation, calcification, and enrichment of chaperones, cytoskeletal, and intermediate filament proteins in the adult hippocampus CA1 following neonatal exposure to the nonprotein amino acid BMAA. *Arch Toxicol.* 89:423-36.
3. Shariatgorji M, Nilsson A, Källback P, Karlsson O, Zhang X, Svenningsson P, Andren PE. (2015) Pyrylium Salts as Reactive Matrices for MALDI-MS Imaging of Biologically Active Primary Amines. *J Am Soc Mass Spectrom.* 26:934-9.
4. Jernerén F, Söderquist M, Karlsson O. (2015) Post-sampling release of free fatty acids - effects of heat stabilization and methods of euthanasia. *J Pharmacol Toxicol Methods* 71:13-20.
5. Karlsson O, Colombo G, Roman E. (2015) Low copulatory activity in selectively bred Sardinian alcohol-nonpreferring (sNP) relative to alcohol-preferring (sP) rats. *Ups J Med Sci.* 120:181-9.
6. Sun, Y., Jergil, M., Stockling, K., Hellmold, H., Dencker, L. et al. (2014). Short-Term Embryonic Stem Cell Exposure Assay As a Tool to Predict Developmental Toxicity. *Birth defects research. Clinical and molecular teratology*, 100:402-402.
7. Stingl, C., Söderquist, M., Karlsson, O., Boren, M., Luider, T. (2014) Uncovering ex-vivo degradation of peptides and neuropeptides during sample extraction in rat brain tissue by oxygen-18 labelling. *J Proteome Res.* 13:2807-2817
8. Hanrieder, J., Karlsson, O., Brittebo, EB, Malmberg, P., and Ewing, AG. (2014) Probing the Lipid Chemistry of Neurotoxin Induced Hippocampal Lesions Using ToF-SIMS Imaging. *Surf Interf Anal.* 46:375–378
9. Karlsson O, Bergquist J, Andersson M. (2014) Quality measures of imaging mass spectrometry aids in revealing long-term striatal protein changes induced by neonatal exposure to the cyanobacterial toxin  $\beta$ -methylamino-L-alanine (BMAA). *Mol Cell Proteomics.* 13:93-104.
10. Ritchie H, Svensson CH, Nilsson MF, Webster WS. (2014) A comparison of drug-induced cardiotoxicity in rat embryos cultured in human serum or protein free media. *J Pharmacol Toxicol Methods.* 70:276-82
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### Agencies that supported the work/Funding 2015

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 Carl Tryggers Stiftelse  
 Fredrik och Ingrid Thuring's Stiftelse

## Other commitments/assignments of group members 2015

Eva Brittebo: Member of the Quality Advisory Board on issues of Uppsala University quality management. Member of the Faculty of Pharmacy Recruitment Group. Member of the Executive Committee for the SafeSciMET education and training program in safety sciences for medicines. Member of the Swetox Consortium Board. Study director in toxicology.

Björn Hellman: Head of Department; Member of the local committee for scholarships at the Faculty of Pharmacy; Study director in toxicology.

## Projects

*(I) Neurodegeneration following neonatal exposure to neurotoxins (Eva Brittebo, Oskar Karlsson, Lisa Ersson, Daiane Cattani, Malin Andersson, Per Andrén in collaboration with Marie Andersson and Ingvar Brandt, Dept Environmental Toxicology, Uppsala university; Mikael Enskog and Curt Petterson, Dept of Medicinal Chemistry Uppsala University; Anna-Lena Berg, AstraZenca, Södertälje; Jörg Hanrieder, University of Gothenburg, and Leopold Ilag, Lying Jiang, Stockholm University, Ariane Zamoner, Universidade Federal de Santa Catarina, Brazil)*

Neurodegenerative disease is a major health problem with increasing incidences in an ageing population. Such disease has a multi-factorial etiology involving interplay between genetic and environmental risk factors e.g. neurotoxic contaminants. BMAA (beta-N-methylamino-L-alanine) is a neurotoxic amino acid that is produced by cyanobacteria. This neurotoxin has been suggested to contribute to a particular neurodegenerative disease in humans. Our studies demonstrated a poor transfer of radiolabelled BMAA across the blood-brain barrier (BBB) with no specific localisation in discrete brain regions in adult mice. In neonatal mice, however, there is an efficient transport across the BBB and a selective uptake of radioactivity in the hippocampus and striatum. Neonatal exposure to BMAA also induces cognitive impairments such as reduced spatial learning and memory abilities in adulthood. In addition, neonatal rat pups treated with a high dose BMAA show acute neuronal cell death in the hippocampus, retrosplenial and cingulate cortices. These brain areas are all important for cognitive function. Histopathological and ultrastructural analysis identified intracellular fibril formation, neuronal degeneration, cell loss, calcium deposits and astrogliosis in the hippocampus of adult animals following a short-term neonatal exposure. Lower doses of BMAA caused distinct impairments in learning and memory function in adult animals following neonatal exposure without acute morphological changes in the brain.

Proteomic analysis of the adult hippocampus demonstrated an enrichment of chaperones, cytoskeletal and intermediate filament proteins, and proteins involved in the antioxidant defense system in the damaged area. Several of the most enriched proteins (plectin, glial fibrillar acidic protein, vimentin, Hsp 27, and ubiquitin) are known to form complex astrocytic inclusions. Moreover, neonatal exposure to BMAA results in increased protein ubiquitination in the adult hippocampus indicating that BMAA may induce protein aggregation. MALDI-TOF imaging mass spectrometry demonstrated that developmental exposure to BMAA induces changes in the expression of S100 $\beta$ , histones, calcium and calmodulin-binding proteins as well as guanine nucleotide-binding proteins in the damaged hippocampus. In addition, time-of-flight secondary ion mass spectrometry based imaging revealed BMAA-induced localization of phosphatidylcholine lipids in the damaged hippocampus. In addition, mass spectrometric quantification of the relative levels of endogenous neuropeptides in the neonatal striatum revealed that 25 peptides from 13 neuropeptide precursors were significantly changed. In another study we characterized changes of major intermediary metabolites in serum following neonatal exposure to BMAA using a non-targeted metabolomic approach. NMR data indicated that metabolites associated with energy metabolism and amino acid metabolism, were changed in serum of BMAA-treated neonatal rats.

Autoradiographic imaging of <sup>14</sup>C-labelled BMAA demonstrated a distinct uptake of radioactivity that was retained following acid extraction in tissues with a high rate of cell turnover and/or protein synthesis. Ultra-high performance liquid chromatography-tandem mass spectrometry analysis demonstrated a dose-dependent increase of protein-associated BMAA in neonatal rat tissues. BMAA was also associated to proteins in the brain, especially in the hippocampus. The level in the brain was, however, considerably lower compared to the liver that is not a target organ for BMAA. In contrast to the liver there was a significantly increased level of protein-association of BMAA in the hippocampus and other brain areas

following repeated administration suggesting that the degradation of BMAA-associated proteins may be lower in neonatal brain than in the liver. The association of BMAA to rat proteins suggests that BMAA may be misincorporated into proteins. However, protein-associated BMAA seems to be cleared over time, as none of the samples from adult rats have any detectable free or protein-associated BMAA. Another study suggests that, secretion of radiolabelled BMAA into milk is an elimination pathway in lactating mothers. Following secretion of radiolabelled L-BMAA into milk, the levels of radioactivity in the brains of the suckling neonatal mice significantly exceeded the levels in the maternal brains. *In vitro* studies using the mouse mammary epithelial HC11 cell line confirmed a more efficient influx and efflux of L-BMAA than of D-BMAA in cells, suggesting enantiomer-selective transport. Competition experiments with other amino acids suggest that the amino acid transporters LAT1 and LAT2 may contribute to the transport of L-BMAA into milk.

Glyphosate is the most widely used broad-spectrum herbicide in the world. A research group at the Universidade Federal de Santa Catarina, Brazil has recently demonstrated that exposure to a glyphosate formulation during the development induces an acute excitotoxicity in the neonatal rodent hippocampus. In collaboration with the Brazilian research group we are examining the long-term effects in the adult brain following neonatal exposure to a glyphosate formulation using MALDI-TOF imaging mass spectrometry. The effects of glyphosate on differentiating neuronal cells are also studied.

*(II) Nasal transfer of therapeutic agents (Eva Brittebo, Elena Piras, in collaboration with Moa Fransson and Angelica Loskog, Dept. of Immunology, Genetics and Pathology, Uppsala University)*

The nasal olfactory pathway is a potential route of delivery of therapeutic agents that do not easily pass the blood-brain barrier. The olfactory neurons have direct contact with the external environment via dendrites in the nasal mucus and with the brain via axons that reach the olfactory bulb without synaptic connections. The olfactory transfer of therapeutic agents into the brain is a novel principle for drug delivery. Intranasal administration of CNS-targeting modified CD4+ T cells significantly reduced disease symptoms as well as decreased IL-12 mRNA in a mouse model of brain inflammation - experimental autoimmune encephalomyelitis (EAE). Immunohistochemical markers for myelination and reactive astrogliosis confirmed recovery in mice treated with engineered Tregs. Furthermore, Treg-treated symptom-free mice recovering from EAE were rechallenged with a second EAE-inducing inoculum but remained healthy, demonstrating a sustained effect of engineered Tregs. In another study, intranasal administration of engineered mesenchymal stromal cells (MSCs) expressing a CNS-targeting receptor significantly reduced disease symptoms of EAE. Mice treated with CNS-targeting MSCs were resistant to further EAE induction whereas non-targeted MSCs did not give such persistent effects. This demonstrates that intranasal delivery of central nervous system-retargeted human mesenchymal stromal cells can prolong treatment efficacy in experimental autoimmune encephalomyelitis.

*(III) Evaluation of genotoxic/antigenotoxic effects of dietary anti-oxidants, traditionally used medicinal plants and other compounds of naturally origin (Björn Hellman Rikard Åsgård, Jasem A Shemali, Lena Norgren, in collaboration with Ulf Göransson, Dept Medicinal Chemistry, Uppsala University, Abdu Adem, Thomas Edward Adrian, UAE University Al Ain)*

In this project, we have mainly been evaluating the genotoxic and anti-genotoxic effects of some plants used in traditional medicine in Ethiopia and other countries, and in these studies we also include fractions of extracts and/or pure compounds from extracts. Following up clinical studies showing that intake of  $\beta$ -carotene and other antioxidants from the diet was found to be associated with a lower level of oxidative DNA damage in mononuclear leukocytes, and we have also recently published a study on the effect of  $\beta$ -carotene on catechol-induced DNA damage in mouse lymphoma cells. Our most recent study on compounds of natural origin is a study on Frondoside A (a triterpenoid glycoside isolated from the Atlantic sea cucumber *Cucumaria frondosa*) which was found to enhance the anti-cancer effects of gemcitabine, a finding, which in the future may turn out to be of clinical benefit for patients with pancreatic cancer.

*(IV) Evaluation of the potential genotoxicity of drinking water (Björn Hellman, Lena Norgren in collaboration with Agneta Oskarsson and Karin Winberg, Swedish Agricultural University)*

In our most recent project evaluating the potential genotoxicity of water samples intended to be used as drinking water, we have introduced HepG2-cells as an additional cell line to be used in our tests. The latter cells are liver cells of human origin, commonly used when screening for toxicological profiles of xenobiotics *in vitro*. The main objective of the drinking water project (which we only are a small part of)

is to develop methods for assessment of hazardous chemicals in drinking water by integrating chemical analysis and *in vitro* toxicity testing. Read more about the “SafeDrink project” on <http://www.slu.se/en/departments/aquatic-sciences-assessment/research/safedrink/>

*(V) Imaging Mass Spectrometry study of basal ganglia levels of neuropeptides in L-DOPA-induced dyskinesia in experimental Parkinson’s disease (Malin Andersson in collaboration with Anna Ljungdahl, Madelene Svedin, Kristen Burnum (PNLL), Jonas Bergquist, Uppsala University)*

In this project we study Parkinson’s disease (PD), which is characterized by degeneration of dopaminergic neurons and accompanied by a dramatic loss of dopamine in the striatum, particularly in the putamen. About 1% of the population over 65 years suffers from PD and the cardinal symptoms that include akinesia, bradykinesia, muscle rigidity, and tremor. Dopamine replacement therapy with L-DOPA is currently the most effective pharmacotherapy for patients with PD, however with PD disease progression and long-term L-DOPA treatment complications occur in many patients. The main complications are troublesome motor complications such as “wearing off” fluctuations and L-DOPA-induced dyskinesia. Despite large efforts in the field of dyskinesia research, the difficulties we face today remain as how to dissociate the molecular changes that arise by the motor execution of dyskinesia from causative changes that induce or predispose to dyskinesias. Our group studies the brain structures of the basal ganglia for molecular correlates of the incidence and severity of dyskinesia in an experimental model of Parkinson’s disease. We focus on neuropeptides and proteins involved in the development of L-DOPA-induced dyskinesias in experimental Parkinson’s disease. Here we use MALDI-TOF imaging mass spectrometry (IMS) for unbiased assessment and topographical elucidation of neuropeptides and proteins in the basal ganglia of high and low dyskinetic animals. MALDI-IMS has the unique advantage of high sensitivity, high molecular specificity, and the detection of hundreds of molecular species in a single tissue section. Indeed, several dynorphins and enkephalins have been detected in these studies, including dynorphin B, dynorphin A(1-8), alpha-neoendorphin, MetEnkRF, MetEnkRGL, PEnk (198-209, 219-229).

*(VI) Imaging MALDI mass spectrometry characterization of opioid peptides after a single dose cocaine or morphine (Malin Andersson in collaboration with Emma Gustafsson, Jonas Bergquist, Jan Rodriguez Parkitna and Ryszard Przewlocki (Polish Academy of Science)*

Drugs of abuse causes rapid changes in neurotransmission, for example release followed by synthesis of opioid peptides in different nuclei of the basal ganglia. In this study we examine the localization and time course of opioid peptides after an acute dose of either cocaine or morphine.

*(VII) MALDI mass spectrometry based molecular phenotyping of CNS and PNS glial cells for prediction in mammalian brain tissue (Malin Andersson in collaboration with Jörg Hanrieder, Grzegorz Wicher, Karin Forsberg Nilsson, Ping Sui, Jonas Bergquist, Åsa Fex-Svenningsen SDU, Denmark)*

These are several similar studies that examine the use of differential protein expression profiling of mammalian neural cells by means of MALDI TOF MS. MALDI MS profiling analysis is rapid, sensitive, robust and specific for large biomolecules in complex matrices. A new straightforward methodology was developed for direct characterization of rodent CNS glial cells using MALDI MS based intact cell mass spectrometry. Molecular phenotyping enables the characterization of cell growth stages, (stem) cell differentiation as well as probing cellular responses towards different stimulations. On-going work includes profiling of glioblastoma multiforme-derived cell lines for the prediction of cancer severity to facilitate diagnosis.

*(IX) In vitro system development for prenatal cardiotoxicity (Mats Nilsson in collaboration with Helen Ritchie, William S. Webster, University of Sydney, Australia)*

The embryo is not protected from pharmaceuticals and environmental pollutants. Intended and unintended pharmacological effects of drugs are often exerted in the conceptus as well. Various therapeutic drugs may affect the functioning of the early embryonic heart resulting in periods of bradycardia, arrhythmia and/or heart block. Such an effect would result in periods of hypoxia in the embryo and may result in spontaneous abortion, fetal growth retardation or in rare cases birth defects. In this project we examine the effect on rat embryonic cardiac function *in vitro* of pharmaceutical drugs with various ion channel-blocking properties.

*(X) Developmental neurotoxicity during the postnatal synaptogenic sensitivity period (Birger Scholz, Henrik Alm)*

Brain development includes key neurodevelopmental prenatal and postnatal stages where environmental stimuli (such as neurotoxicants) can be particularly efficient in inducing long lasting changes in neurodevelopmental trajectories. During the first two to three postnatal weeks of rodent life, there is a synaptogenic sensitivity period, corresponding to the first ~2-3 years of human life. DNT exposure within (but not outside) this period commonly leads to potentially irreversible alterations in rodent adult brain function. Substances as chemically diverse as metals, environmental chemicals (PCBs, Bisphenol A and polybrominated flame retardants (PBDEs)) and medical drugs (GABA type A agonists and NMDA antagonists) lead all to similar long lasting behavioral effects in animals when exposed during this sensitivity period. An important challenge is therefore to identify the molecular basis of the disruptions that lead to irreversible changes in adult behaviour and the nature of how the effects from such dissimilar substances can converge into similar phenotypes.

This research project aims at identifying how neurotoxicants (both environmental toxicants and medical drugs) induce both structural and epigenomic changes during the sensitivity period (postnatal day 10 in rodents) and how these effects become propagated to adult age. We have so far focused on Polybrominated diphenyl ether (PBDE), Bisphenol A and Ketamine induced effects on neural DNA methylation, gene expression and neurosignalling (peptidomic) changes in the cortex and hippocampus.

*(XI) Developmental contingencies and Developmental Neurotoxicology in vitro (Birger Scholz)*

The issue of in-vitro extrapolation to in-vivo conditions is of great importance in predictive toxicology considering how resource-intensive animal studies are. Unfortunately, the majority of today's *in silico* and *in vitro* assays suffer from weak predictive power for more complex toxicological endpoints. The mechanistic information from the postnatal sensitivity project is to be applied to the development of a range of more representative and specific DNT in vitro tests. To this end, this project investigates if and under what in-vitro conditions the observed developmental sensitivity period specific gene regulation changes can be replicated in cell lines (immortalized somatic neural cell lines and teratocarcinoma derived cell lines). In order to make the *in vitro* tests as relevant as possible for possible future hazard identification, they are conducted with and optimized for High Content Analysis/High Content Screening

## Medical Mass Spectrometry

Per Andrén

<http://farmbio.uu.se/research/researchgroups/mms>

The research group focus on new approaches in mass spectrometry imaging (MSI), i.e. matrix-assisted laser desorption ionization (MALDI) and desorption electrospray ionization (DESI)–MSI of biological tissue sections, and peptidomics, the comprehensive study of endogenous peptides.

Mass spectrometry imaging is a novel technique used to determine the spatial distribution of molecular species in biological tissue sections *in situ*. The technology allows analysis and visualization of endogenous proteins, peptides and neurotransmitters, as well as drugs and their metabolites, in their native biochemical states within the same tissue section with high molecular specificity. Molecular images are created by rasterizing over the sample while collecting MS or tandem MS (MS/MS) spectra from every position at a chosen spatial resolution. The localization pattern from individual molecular species present on the tissue surface can then be extracted and positioned on the original histological image with the abundances represented by a concentration dependent color scale.

Peptidomics involves the comprehensive analysis of the endogenous peptide content of a certain cell, organ, body fluid, or organism. It complements molecular biology approaches in its ability to characterize the processing of translation products, including changes in expression or posttranslational modifications (PTMs) of peptides and small proteins. By comparing the proteins and peptides in samples of diseased tissue with those in normal tissue, differential expression patterns can be detected that may lead to the identification of novel biomarkers.

The objective of our research is to utilize MSI and peptidomics approaches to study neurochemical processes in Parkinson's disease (PD) and specifically L-Dopa-induced dyskinesias (LID). The aim is to define neuropeptides, proteins and neurotransmitters that are differentially expressed in the basal ganglia complex of animals with striatal dopamine depletions, and to determine which of these molecules are regulated by loss of dopamine signaling, as well as to investigate their expression patterns in subjects with and without LID symptoms.

Understanding the relationship between pharmacokinetics and pharmacodynamics is crucial in the development of effective drug therapies. Current technologies only provide information on the total amount of drug in the whole tissue with no possibility to address micro-environmental localization of the compound or metabolic derivatives. The applications of MSI in drug discovery studies provide a new tool to accurately measure local tissue concentrations of drug/drug metabolite in context with efficacy achieved with the targeted biology or in the context of safety.

Our laboratory is equipped with the latest separation and MS technologies, i.e., three MALDI-MSI instruments; two Ultraflextreme TOF/TOF (Bruker Daltonics) and one MALDI/ESI Q-TOF Synapt G2si (Water Corp.), and two electrospray ionization mass spectrometers, LTQ (Thermo Scientific) and a high-resolution Q-TOF mass spectrometer (Maxis Impact, Bruker Daltonics).

### Members of the group during 2015

Per Andrén, Associate Prof.  
 Anna Nilsson, Researcher  
 Henrik Wadensten, Researcher  
 Mohammadreza Shariatgorji, Researcher  
 Patrik Källback, PhD student  
 Theodosia Vallianatou, PhD student

### Publications 2013-2015

1. Nilsson CL, Berven F, Selheim F, Liu H, Moskal JR, Kroes RA, Sulman EP, Conrad CA, Lang FF, Andrén PE, Nilsson A, Carlsohn E, Lilja H, Malm J, Fenyö D, Subramaniam D, Wang X, Gonzales-Gonzales M, Dasilva N, Diez P, Fuentes M, Végvári Á, Sjödin K, Welinder C, Laurell T, Fehniger TE, Lindberg H, Rezeli M, Edula G, Hober S, Marko-Varga G. (2013) Chromosome 19 annotations with disease speciation: a first report from the Global Research Consortium. *J Proteome Res.* 12:135-50.
2. Karlsson O, Kultima K, Wadensten H, Nilsson A, Roman E, Andren PE, Brittebo E (2013) Neurotoxin-induced neuropeptide perturbations in striatum of neonatal rats. *J Proteome Res.* 12:1678-90.
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4. Malm J, Fehniger TE, Danmyr P, Végvári A, Welinder C, Lindberg H, Appelqvist R, Sjödin K, Wieslander E, Laurell T, Hober S, Berven FS, Fenyö D, Wang X, Andrén PE, Edula G, Carlsohn E, Fuentes M, Nilsson CL, Dahlbäck M, Rezeli M, Erlinge D, Marko-Varga G (2013) Developments in biobanking workflow standardization providing sample integrity and stability. *J Proteomics.* 95:38-45.
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6. Bourdenx M, Nilsson A, Wadensten H, Fälth M, Li Q, Crossman AR, Andrén PE, Bezard E (2014) Abnormal structure-specific peptide transmission and processing in a primate model of Parkinson's disease and l-DOPA-induced dyskinesia. *Neurobiol Dis.* 62:307-312.
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4. Nilsson A, Goodwin RJ, Shariatgorji M, Vallianatou T, Webborn PJ, Andrén PE (2015) Mass Spectrometry Imaging in Drug Development. *Anal Chem.* 87:1437-1455.

### Agencies that supported the work/Funding 2015

The Swedish Research Council (VR); NT (2014–6215), MH (2008-5597 and 2013–3105) and RFI (2009-6050); VINNOVA; Japan Society for the Promotion of Science (JSPS) Joint Projects; AstraZeneca Global DMPK; FP7-PEOPLE-2013- Marie Curie Initial Training Networks (ITN); Uppsala University – Infrastructure.

### Other commitments/assignments of group members 2015

Per Andrén: Swedish Academy of Pharmaceutical Sciences, Section for Drug Analysis (member of the board); Swedish Proteomics Society (member of the board); Journal of Proteomics (editorial board); European Proteomics Association (EuPA) Open Proteomics (editorial board), Cooperation in Science and Technology (COST) Action; Mass Spectrometry Imaging (member of the management committee).

## Projects

*(I) Functional neuroproteomic changes associated with L-Dopa-induced dyskinesia in Parkinson's disease (Collaboration with Per Svenningsson, Karolinska Institutet, Erwan Bezard, Univ. of Bordeaux 2, France)*

The main objective of the present research is to study neurochemical processes in Parkinson disease and specifically L-Dopa-induced dyskinesias (LID). No treatment exists yet for the management of LID, a debilitating complication of L-dopa therapy for PD. The aim is to define neuropeptides, proteins, neurotransmitters and endogenous metabolites that are differentially expressed in the basal ganglia complex of animals with striatal dopamine depletions, with and without LID. Project supported by VR-MH grant 2013–3105.

*(II) Mass Spectrometry Imaging in Drug Discovery – Targeting drugs, metabolites, peptides and neurotransmitters*

General objectives include development of MALDI-MSI in the following health research and drug discovery areas, a) ADME, high-resolution imaging the spatial distribution and absolute concentration of unlabeled drugs within tissue micro-compartments, b) Safety Toxicology, monitoring of molecular toxicology in organs (kidney), c) Blood-Brain Barrier (BBB), determine BBB profile for drug candidates in the early stages of the drug development, d) Neuroscience, absolute quantitation of neurotransmitters directly in tissue sections, e) msiQuant, MSI software development for quantitation of drugs, metabolites and endogenous compounds directly in tissue sections. The application of these methods will provide new tools aiding pharmaceutical development, as well as for clinical and experimental model systems. Project supported by VR-NT grant 2014–6215.

*(III) Identification and functional characterization of protein-protein interactions in cerebrospinal fluid and brain tissue from Parkinson's disease (PD) patients and experimental PD models (Collaboration with Per Svenningsson, Karolinska Institutet)*

Using surface plasmon resonance technique (Biacore 3000) coupled to mass spectrometry technology, new protein partners of  $\alpha$ -synuclein and parkin have been captured and identified in cerebrospinal fluid or post-mortem human tissue from PD patients. In addition to using native  $\alpha$ -synuclein and parkin, mutated forms of these proteins seen in familiar forms of PD will be immobilized on the sensor chip and used as baits. Project supported by VR-MH grant 2013–3105 and 2011-4722.

*(IV) Fine mapping the spatial distribution and concentration of unlabeled drugs within tissue micro-compartments using imaging mass spectrometry (Collaboration with AstraZeneca)*

In respiratory inhalation drug discovery projects, one key objective is to optimize retention of compound in the lung and consequently achieve duration of effect. Current technologies only provide information on the total amount of compound in the whole lung with no possibility to address micro-environmental localization of the compound or metabolic derivatives. The application of MALDI-MSI in such studies would provide a new tool to accurately measure local tissue concentrations of drug/drug metabolite in context with efficacy achieved with the targeted biology or in the context of safety. Such technology could provide new and important information. Project supported by collaboration agreement with AstraZeneca.

*(V) Characterization of drug-induced kidney toxicity using MALDI-MSI (Collaboration with AstraZeneca)*

The project is aimed at developing a routine methodology for the use of MALDI imaging in problem solving of toxicity questions. It will include preparation of tissue material, interpretation of the data generated and structure elucidation of new analytes. Project supported by collaboration agreement with AstraZeneca.

## Neuropharmacology and Biological Research on Addiction

### Ingrid Nylander

The research group Neuropharmacology and Biological Research on Addiction have three different research areas focusing on the role of basic neurology, and physiological and pathophysiological mechanisms leading to addiction and neurodegenerative diseases. The research group is divided into the three following subgroups:

- Neuropharmacology, Addiction and Behavior
- Biological Research on Drug Dependence
- Molecular Neuropsychopharmacology

### Neuropharmacology, Addiction and Behaviour

Ingrid Nylander, Lena Bergström, Lova Segerström, Erika Roman, Anne-Lie Svensson

<http://farmbio.uu.se/research/researchgroups/nab>

The research in the group is devoted to studies on basic neurobiology, physiological and pathophysiological mechanisms in the brain, and neuropharmacology of relevance for substance use disorders and neurodegenerative diseases. Current projects include studies of neurobiological substrates for individual differences in addiction processes, especially vulnerability for risk consumption of alcohol and alcohol use disorders, and responses to drugs used in treatment of addiction. Alcohol addiction is a complex trait and the phenotype related to vulnerability for addiction is based on the interaction of multiple genes and environmental factors. Of particular interest for us is the impact of early environmental factors, such as rearing environment and the consequences of adolescent drug intake, and the association between behavioural characteristics, such as risk taking behaviour, and later drug consumption. It is hypothesized that disruption of early developmental processes in transmitter networks either by rearing factors or drug intake early in life, causes long-term changes in brain function and behaviour that, in turn, affects alcohol consumption later in life. We collaborate with Drs E Comasco, L Oreland (Department of Neuroscience) and K Nilsson (Centre for Clinical Research, Västerås) in projects that include investigation of how epigenetic processes are involved in long-term consequences of exposure to various early-life environmental factors.

Another line of research investigates the role of cannabinoids and neurosteroids and endocannabinoids for neurogenesis and for interactive processes that are ongoing in neurodegenerative disorders, like Alzheimer's disease (AD). Since AD is associated with excitotoxicity, oxidative stress and neuroinflammation, the research line emphasis on neuroprotective properties of neurosteroids and endocannabinoids against different toxic insults in *in vitro* cell models.

In the projects, experimental models are used in combination with extensive evaluation of neurobiological and behavioural consequences of different early environmental conditions. A number of behavioural models within the field of neuroscience and neuropharmacology are employed including tests for assessment of neonatal development, exploratory behaviour, locomotor activity, risk taking behaviour, self-administration, learning and memory and a multivariate test arena (the multivariate concentric square field™, MCSF) in combination with multivariate data analysis approaches. Neurobiological assessment includes methods to analyse effects on receptors, transmitters and mRNA in tissue samples and brain slices but also analysis of transmitter release and re-uptake patterns using *in vivo* chronoamperometric analysis using Fast Analytical Sensing Technology (FAST). Specific research activities within the group are described shortly under projects.

### Members of the group during 2015

Ingrid Nylander, Professor  
 Lena Bergström, Associate Professor, Senior Lecturer  
 Erika Roman, Associate Professor, Senior Lecturer  
 Anne-Lie Svensson, Senior Lecturer  
 Viktoria Lind, Junior Lecturer  
 Lova Segerström, Researcher  
 Sara Palm, Researcher  
 Sarah Holst, Researcher  
 Shima Momeni, PhD student  
 Linnea Granholm, PhD student  
 Stina Lundberg, PhD student  
 Bengt J Meyerson, Professor Emeritus (deceased October 2015)  
 Marita Berg, Technician  
 Jenny Gustavsson, Research assistant  
 Matilda Persson, Research assistant  
 Laila Nauman, Research assistant

### Publications 2013-2015

1. Daoura L, Nylander I, Roman E. Qualitative differences in pup-retrieval strategies in a maternal separation paradigm. *JBBS* (2013) 3 603-616
2. Palm S, Daoura L, Roman E, Nylander I. Effects of rearing conditions on behavior and endogenous opioids in rats with alcohol access during adolescence. *Plos One* (2013) 8 (10):e76591
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4. Rosén A, Lund I, Lundeborg T, Nylander I. Antinociceptive effects of sensory stimulation involve dynorphin B supraspinally in rats. *Acupuncture Rel Ther* (2013) 35-41
5. Karlsson O, Kultima K, Wadensten H, Nilsson A, Roman E, Andrén PE, Brittebo EB. Neurotoxin-induced neuropeptide perturbations in striatum of neonatal rats. *J Proteome Res* (2013) 12 1678–1690
6. Roman E, Karlsson O. Increased anxiety-like behavior but no cognitive impairments in adult rats exposed to constant light conditions during perinatal development. *Ups J Med Sci* (2013) 118 222-227
7. Meyerson BJ, Jurek B, Roman E. A rank-order procedure applied to an ethoexperimental behavior model – the multivariate concentric square field™ (MCSF) test. *JBBS* (2013) 3 350-361
8. Momeni S, Sharif M, Ågren G, Roman E. Individual differences in risk-related behaviors and voluntary alcohol intake in outbred Wistar rats. *Behavioural Pharmacology* (2014) 25 206-215
9. Palm S, Momeni S, Lundberg S, Nylander I, Roman E. Risk-assessment and risk-taking behavior predict potassium- and amphetamine-induced dopamine response in the dorsal striatum of rats. *Frontiers in Behavioral Neuroscience* (2014) 8 236
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12. Granholm L, Roman E, Nylander I. Single housing during early adolescence causes time-, area- and peptide-specific alterations in endogenous opioids of rat brain. *Br J Pharmacol* (2015) 172 606-614

13. Karlsson O, Colombo G, Roman E. Low copulatory activity in selectively bred Sardinian alcohol-nonpreferring (sNP) relative to alcohol-preferring (sP) rats. *Ups J Med Sci* (2015) 120 181-189
14. Magara S, Holst S, Lundberg S, Roman E, Lindskog M. Altered explorative strategies and reactive coping style in the FSL rat model of depression. *Front Behav Neurosci* (2015) 9 89
15. Momeni S, Segerström L, Roman E. Supplier-dependent differences in intermittent voluntary alcohol intake and response to naltrexone in Wistar rats. *Front Neurosci* (2015) 9 424
16. Karlsson O, Roman E. Dose-dependent effects of alcohol administration on behavioural profiles in the MCSF test. *Alcohol* (2015) doi: 10.1016/j.alcohol.2015.10.003
17. Palm S and Nylander I. Alcohol-induced changes in opioid peptide levels in adolescent rats are dependent on housing conditions. *Alcohol Clin Exp Res* (2015) 38 (12) 2978-2987
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19. Comasco E, Todkar A, Granholm L, Nilsson KW, Nylander I. Alpha 2a-Adrenoceptor Gene Expression and Early Life Stress-Mediated Propensity to Alcohol Drinking in Outbred Rats. *Int. J. Environ. Res. Public Health* (2015) 12 7154-7171
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21. Vrettou M, Granholm L, Todkar A, Nilsson KW, Wallén-Mackenzie Å, Nylander I, Comasco E. Ethanol affects limbic and striatal presynaptic glutamatergic and DNA methylation gene expression in outbred rats exposed to early-life stress (2015) *Addiction Biol* doi:10.1111/adb.12331
22. Bendre M, Comasco E, Nylander I, Nilsson KW, Effect of voluntary alcohol consumption on *Maoa* expression in the mesocorticolimbic brain of adult male rats previously exposed to prolonged maternal separation. *Transl Psychiatry* (2015) 5, e690; doi:10.1038/tp.2015.186

### Reviews 2013-2015

1. Nylander I, Roman E. Is the rodent maternal separation model a valid and effective model for studies on the early-life impact on ethanol consumption? *Psychopharmacology* (2013) 229 555-569

### National publications (Text books) 2013-2015

1. Franck J & Nylander I (Eds.) *Beroendemedicin* (2nd ed.), Studentlitteratur, 2015
2. Nylander I. *Beroendemekanismen*. In *Beroendemedicin*, 2nd ed. (Franck & Nylander, Eds.), Studentlitteratur, 2015
3. Roman E. *Djurexperimentell metodik*. In *Beroendemedicin*, 2nd ed. (Franck & Nylander, Eds.), Studentlitteratur, 2015
4. Svensson A, *Nikotin*. In *Beroendemedicin*, 2nd ed. (Franck & Nylander, Eds.), Studentlitteratur, 2015

### Dissertations 2013-2015

1. Shima Momeni *Profiling of risk-related behaviors in relation to voluntary alcohol intake and neurobiology*, 2014. Faculty of Pharmacy, Uppsala University Licentiate Theses, 47.
2. Shima Momeni *Individual differences in behavior, neurochemistry and pharmacology associated with voluntary alcohol intake*, 2015. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, 205. ISSN 1651-6192.
3. Sara Palm *Early environment, adolescent alcohol drinking and neurobiological responses to drugs*, 2014. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, 190. ISSN 1651-6192.
4. Linnea Granholm *Neurobiological Consequences of Social Conditions and Alcohol Exposure in Adolescent rats*, 2015. Faculty of Pharmacy, Uppsala University Licentiate Theses, 48.

## Agencies that supported the work/Funding 2015

The Swedish Research Council (Nylander); The Alcohol Research Council of the Swedish Alcohol Retailing Monopoly (two separate projects; Nylander and Roman); The Facias Foundation (Roman); Svenska Spels forskningsråd (Roman).

## Other commitments/assignments of group members 2015

Lena Bergström: Member of the Academic Senate, Uppsala University

Ingrid Nylander: Grant committee member Alcohol Research Council of the Swedish Alcohol Retailing Monopoly; Member of the Faculty of Pharmacy committee; Vice chairman, the recruitment committee Faculty of Pharmacy; Chairman, the committee for undergraduate education Faculty of Pharmacy; Vice Dean and board member, Disciplinary Domain of Medicine and Pharmacy

Erika Roman: International Adjunct Associate, Department of Psychology, Indiana University, USA; Review Editor: Frontiers in Addictive Disorders and Behavioural Dyscontrol; External mentor Salvatore Magara, KI; Member of the board: The Society for Swedish Alcohol and Drug Research; Member of the board of Uppsala University Laboratory Animal Resources; Member of the Animal Welfare Body, Uppsala University; Member of the Postgraduate Programs Committee (KUF), Uppsala University; Approved Supervisor by the Swedish Board of Agriculture; Member of the expert panel, Swedish Centre for Animal Welfare; One of three coordinators of Uppsala University Behavioural Facility (UUBF), Uppsala University

Anne-Lie Svensson: Member of the committee for undergraduate education (GRUFF) at the Faculty of Pharmacy, Uppsala University; Member of the gender equality committee at the Disciplinary Domain of Medicine and Pharmacy, Uppsala University, Member of the Academic Senate, Uppsala University.

## Projects

*(I) The impact of early life environment on the brain and the stress axis (Sara Palm, Linnea Granholm, Stina Lundberg, Lova Segerström, Ingrid Nylander)*

*Early life adversity*: We have previously shown that adverse experiences early in life cause long-term neurobiological and behavioural alterations. Currently, we study the mechanisms underlying long-term consequences of early life stress. A rodent maternal separation (MS) model is used to simulate rearing conditions that are associated with resilience (after short MS) or vulnerability (after longer MS) in terms of adult risk consumption. We investigate transmitter networks relevant for reward and reinforcement, for example, endogenous opioid, glutamate and monoamine circuits. In collaboration with Dr E Comasco (Nylander co-supervisor to PhD students Megha Bendre and Maria Vrettou in these studies), we assess the effects of early-life stress on expression and methylation of genes involved in regulation of stress and reward.

*Adolescent drug intake*: We study the long-term effects of adolescent drug exposure on opioid networks and on drug consumption in adulthood. The impact of social deprivation in single housing drinking paradigms has been evaluated. Alcohol-induced effects are evaluated both after voluntary drinking during adolescence and with administration of a given dose alcohol. Currently we examine the effects of adolescent alcohol and also the combined alcohol and nicotine exposure on opioid networks.

*(II) In vivo and in vitro studies of drug-induced effects in the brain (Sara Palm, Anne-Lie Svensson, Ingrid Nylander)*

Chronoamperometry with Fast Analytical Sensing Technology (FAST) is currently used for *in vivo* analysis of dopamine in the brain. This technique enables *in vivo* electrochemical detection of dopamine in anaesthetized or awake animals. Microelectrodes are used to measure electrochemically active substances like dopamine. FAST offers unique advantages as compared to *in vivo* microdialysis: high temporal resolution and close spatial orientation enables a second-by-second analysis instead of several minutes, high sensitivity allows measurement of resting levels, high specificity and the microelectrodes cause minimal brain damage. We analyse effects of adolescent alcohol exposure on dopamine dynamics and have so far shown differences in the dorsal striatum, a brain area involved in transition from controlled to compulsive drug taking behaviour. It is also used to correlate *in vivo* dopamine dynamics with behaviour, e.g. risk taking or risk assessment.

*(III) Method development (Jenny Gustavsson, Lova Segerström, **Ingrid Nylander**)*

Post-mortem metabolism is a major concern in the analysis of opioid peptides in biological samples and we have used the bench top Stabilizer T1 instrument (Denator AB, Gothenburg, Sweden) for heat stabilization of tissue samples to establish a procedure for rapid and efficient enzyme inactivation. We were able to optimize the measurements of opioid peptides in the brain and achieve detection of peptide levels more similar to the *in vivo* concentration with a reliable, reproducible and easy to use procedure to avoid peptide metabolism in the handling of fresh tissues.

*(IV) Behavioural profiling of animals exposed to early environmental stress and adolescent alcohol consumption (Sara Palm, Linnea Granholm, Stina Lundberg, **Erika Roman, Ingrid Nylander**)*

Current experiments analyse the short- and long-term behavioural consequences of rearing in different environmental settings and of long-term alcohol consumption. The project comprises development of animal experimental models to assess maternal behaviour, interactions between the dam and offspring and play behaviour. In addition, the MCSF test is employed to examine behavioural profiles after different rearing conditions and before and after adolescent alcohol consumption. Individual behavioural profiling in animals subjected to various rearing conditions and alcohol early in life in controlled experimental groups enables distinction between rearing-induced and drug-induced consequences for behaviour later in life. It is also examined whether and how altered behaviour relate to vulnerability for drug addiction.

*(V) Development and validation of the MCSF test (**Erika Roman, Stina Lundberg, Matilda Persson, Svante Winberg, Bengt J Meyerson**)*

The multivariate concentric square field™ (MCSF) test is a technique with an ethoexperimental approach that in one and the same test situation includes a variety of physical situations meaning risk, safety, exploratory incentives etc. that permit measurements of general activity, exploration, approach/avoidance in open versus sheltered areas and the motivation for seeking reinforcers. The purpose of the multivariate design is to gather information that taken together should illustrate a personality profile including how this profile is altered under various forms of challenges. The MCSF test is under continuous development. On-going work aims to i) validate the test arena set up for zebrafish, ii) study the impact of pharmacological substances on behavioural profiles, and iii) develop plug-in units for studies of motivated behaviours in rodents and expanding the use of the MCSF for studies of learning and memory.

*(VI) Behavioural profiling of selectively bred alcohol-preferring and alcohol-avoiding rodent lines (**Erika Roman, Robert Stewart, Nicholas Grahame, Tiebing Liang, Giancarlo Colombo, Petri Hyttiä, Lawrence, Lumeng**)*

Genetic aspects of alcohol use disorders have been modelled using rats and mice selectively bred for extremes of alcohol preference and voluntary alcohol intake. Several different lines of rodents have been selectively bred for high and low oral alcohol preference and intake. These lines show similar alcohol drinking phenotypes but have different genetic and environmental backgrounds and may therefore display diverse behavioural traits as seen in human alcohol dependent individuals. This project involves behavioural characterization of selectively bred alcohol preferring and non-preferring rat and mouse lines as well as congenic rat lines using the MCSF test. Moreover, the response to natural reinforcing stimuli is characterized in selectively bred alcohol preferring and non-preferring rat lines. This extensive behavioural characterization enables a deeper understanding of behavioural traits of importance for understanding of alcohol use disorders and related addiction processes.

*(VII) Ethoexperimental studies of appetitive and consummatory mechanisms related to natural rewarding stimuli and drugs of abuse (**Erika Roman, Bengt J Meyerson**)*

The project aims at exploring basic mechanisms of reinforcing stimuli with special focus on differentiating appetite for seeking reinforcers such as food, sexual activity and drugs of abuse from consummatory behaviours. On-going studies assess the animal's motivation for passing the risk area and reach reinforcement by increasing the resistance of passing. The association between natural rewards, such as sexual activity and food intake, and drugs of abuse, i.e. alcohol, is subject for examination. The hypothesis is that reward motivated behaviours are different in animals with different voluntary intake of drugs of abuse.

*(VIII) The role of individual differences in drug-seeking and drug-intake behaviour and associated neurobiological effects of relevance to vulnerability for addiction (Shima Momeni, **Lena Bergström, Erika Roman**)*

We use experimental methods to examine the neurobiological basis for individual differences in risk-related behaviours and the association between voluntary alcohol intake, addiction processes and response to treatment. With regard to neurobiology, focus is on the cannabinoid, opioid and dopamine systems. A multivariate behavioural approach with an ethological foundation that incorporates several aspects of the behavioural repertoire and evolutionary conserved behaviours is used. Risk-related behaviours are of importance for liability for excessive alcohol intake and also affect the response to drug treatment. The impact of individual differences in risk-related behaviours on voluntary alcohol intake and CB1 and opioid receptor density are investigated. We also study the effects of alcohol on FAAH and MAGL enzyme activity, i.e. enzymes metabolizing the endogenous cannabinoids anandamide and 2-AG.

*(IX) Experimental studies of gambling disorder (Sarah Holst, Laila Nauman, **Erika Roman**)*

There is an urgent need for valid animal models for studying the neurobiological and behavioral associations between behavioral traits and decision-making of relevance for gambling disorder and other addictive disorders. This project combines multivariate behavioral profiling, brain network analysis with tasks targeting the actual deficit of gambling disorder with the specific aim to produce a valid translational model.

*(X) Neurosteroids and Alzheimer's disease: Mechanistic studies of neuroprotection and amyloid- $\beta$ -modulation (Anne-Lie Svensson)*

Neurosteroids are endogenous modulators of neuronal functions responsible for many biological and pathophysiological effects. Plausible links between neurosteroids and neurodegenerative disorders, like Alzheimer's disease (AD), are suggested. Accumulation of amyloid- $\beta$  induced by toxic events in cells might be able to reduce the synthesis of neuroprotective neurosteroids, thus favour/support neurodegenerative processes. In ongoing studies the neuroprotective properties of neurosteroids and their metabolites, against amyloid- $\beta$ -induced toxicity are investigated. Since neurosteroids most likely affects neuronal and glial cells differently, their effects on amyloid- $\beta$ -induced toxicity are also investigated in numerous cell types.

*(XI) Neuroprotective properties of endocannabinoids against different toxic insults (Anne-Lie Svensson)*

The endocannabinoid system is widespread in the central nervous system and involved in many neurophysiological processes. Neurodegenerative disorders, such as Alzheimer's disease (AD), are associated with excitotoxicity, oxidative stress and neuroinflammation. Endocannabinoids have been demonstrated to affect the progression of neurodegeneration. In ongoing studies the neuroprotective properties of different endocannabinoids against toxic insults are investigated in numerous cell types.

## Biological Research on Drug Dependence

Mathias Hallberg and Fred Nyberg

<http://farmbio.uu.se/research/researchgroups/brdd/biolbero>

The first objective of our research is to improve the understanding of brain mechanisms of relevance for the etiology of drug addiction and to develop strategies for relevant treatment. Studies of drug effects on brain circuits involved in reward and dependence are combined with attempts to develop peptidomimetics reducing the expression of opioid withdrawal, an important aspect in the treatment of opioid addiction. Studies are also aimed to find strategies to repair drug-induced brain damages, i.e. using peptidomimetics or growth factors counteracting apoptosis and stimulating neurogenesis. As animal models do not reflect all complexity of addiction current research also involves epigenetic adaptations and regulation of plastic neuropeptide genes in human brain. The second objective of our research is a) to discover new classes of analgesics with potential use in the treatment of neuropathic pain and b) to discover unique cognitive enhancers. These two subprojects are derived from findings in the tachykinin and renin-angiotensin systems.

### Members of the group during 2015

Mathias Hallberg, PhD Associate Professor  
 Fred Nyberg, Senior Professor  
 Alfhild Grönbladh, PhD  
 Anna Carlsson PhD student  
 Erika Brolin, PhD student  
 Erik Nylander, PhD student  
 Shanti Diwakarla, Postdoc  
 Anna Lesniak, Postdoc  
 Stina Silander, Project administrator  
 Myron Zaluha, Project manager

### Publications 2013-2015

1. Grönbladh A, Johansson J, Nyberg F, Hallberg M (2013) Recombinant human growth hormone affects the density and functionality of GABA(B) receptors in the male rat brain. *Neuroendocrinology*. 97:203-211.
2. Johansson J, Grönbladh A, Nyberg F, Hallberg M. (2013) Application of *in vitro* [<sup>35</sup>S]GTPγ-S autoradiography in studies of growth hormone effects on opioid receptors in the male rat brain. *Brain Res Bull*. 90:100-106.
3. Grönbladh A, Johansson J, Nöstl A, Nyberg F, Hallberg M. (2013) Growth hormone improves spatial memory and reverses certain anabolic androgenic steroid-induced effects in intact rats. *J Endocrinology*. 216:31-41.
4. Bazov I, Kononenko O, Watanabe H, KuntiĆ V, Sarkisyan D, Taqi MM, Hussain MZ, Nyberg F, Yakovleva T, Bakalkin G (2013) The endogenous opioid system in human alcoholics: molecular adaptations in brain areas involved in cognitive control of addiction. *Addict Biol*. Jan;18(1):161-9.
5. Enhamre-Brolin E, Carlsson A, Hallberg M, Nyberg F. (2013). Growth hormone reverses streptozotocin-induced cognitive impairments in male mice. *Behav Brain Res*. 238:273-278.
6. Rhodin A, Grönbladh A, Ginya H, Nilsson KW, Rosenblad A, Zhou Q, Enlund M, Hallberg M, Gordh T, Nyberg F. (2013) Combined analysis of circulating beta-endorphin with gene polymorphisms in OPRM1, CACNAD2 and ABCB1 reveals correlation with pain, opioid

- sensitivity and opioid-related side effects. *Molecular Brain*. 2013;6:8.
7. Fransson R, Sköld C, Kratz J, Svensson R, Artursson P, Nyberg F, Hallberg M, Sandström A. (2013) Constrained H-Phe-Phe-NH<sub>2</sub> analogues with high affinity to the substance P 1-7 binding site and with improved metabolic stability and cell permeability. *J Med Chem*. 56:4953-4965.
  8. Hallberg M, Nyberg F (2013) Fortfarande oklart om steroider framkallar eget beroende. *Läkartidningen*. 110:1736-1739.
  9. Grönbladh A, Johansson J, Bergquist J, Hallberg M. (2013) The impact of nandrolone decanoate and growth hormone on plasma steroid levels in rodents. *Steroids*. 78:1192-1199.
  10. Ali MA, Kazzam E, Amir N, Nyberg F, Adem A. (2013) Effects of dehydration and blockade of angiotensin II AT<sub>1</sub> receptor on stress hormones and anti-oxidants in the one-humped camel. *BMC Vet Res*. 2013 doi: 10.1186/1746-6148-9-232.
  11. Adem A, Al Haj M, Benedict S, Yasin J, Nagelkerke N, Nyberg F, Yandle TG, Frampton CM, Lewis LK, Nicholls MG, Kazzam E. (2013) ANP and BNP responses to dehydration in the one-humped camel and effects of blocking the renin-angiotensin system. *PLoS One*. 2013;8(3):e57806.
  12. Grönbladh A, Johansson J, Nyberg F, Hallberg M. (2013) Administration of growth hormone and nandrolone decanoate alters gene expression in the hypothalamus and pituitary. *Growth Hormone and IGF-1 Res*. Growth Hormone and IGF-1 Res. Jan 16, S1096-6374(14)00003-3.
  13. Johansson J, Grönbladh A, Hallberg M. (2013) Gamma-hydroxybutyric acid (GHB) induces cognitive deficits and affects GABA<sub>B</sub>- and IGF-1 receptors in male rats. *Behav Brain Res*. S0166-4328(14)00252-6.
  14. Righard L, Carlsson-Jonsson A, Nyberg F. (2014) Enhanced levels of immunoreactive  $\beta$ -casomorphin-8 in milk of breastfeeding women with mastitis. *Peptides*. 51:54-8.
  15. Borhade S, Rosenström U, Sävmarker J, Lundbäck T, Jenmalm-Jensen A, Sigmundsson K, Axelsson H, Svensson F, Konda V, Sköld C, Larhed M, Hallberg M. (2014) Inhibition of Insulin-regulated aminopeptidase (IRAP) by arylsulfonamides. *Chemistry Open*. 3(6):256-63.
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  17. Heddini U, Bohm-Starke N, Grönbladh A, Nyberg F, Nilsson KW, Johannesson U. Serotonin receptor gene (5HT-2A) polymorphism is associated with provoked vestibulodynia and comorbid symptoms of pain. *J Sex Med*. 11(12):3064-71.
  18. Carlsson-Jonsson A, Gao T, Hao JX, Fransson R, Sandström A, Nyberg F, Wiesenfeld-Hallin Z, Xu XJ. (2014) N-terminal truncations of substance P 1-7 amide affect its action on spinal cord injury-induced mechanical allodynia in rats. *Eur J Pharmacol*. 738:319-25.
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  21. Wallinder C, Sköld C, Botros M, Guimond MO, Hallberg M, Gallo-Payet N, Karlén A, Alterman M. (2014) Interconversion of Functional Activity by Minor Structural Alterations in Nonpeptide AT<sub>2</sub> Receptor Ligands. *ACS Med Chem Lett*. 2014 6(2):178-82.
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  23. Muresanu DF, Sharma A, Lafuente JV, Patnaik R, Tian ZR, Nyberg F, Sharma HS. (2015) Nanowired Delivery of Growth Hormone Attenuates Pathophysiology of Spinal Cord Injury and Enhances Insulin-Like Growth Factor-1 Concentration in the Plasma and the Spinal Cord. *Mol Neurobiol*. 52(2):837-45
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3. Nyberg F. (2014) Structural plasticity of the brain to psychostimulant use. *Neuropharmacology.* 87:115-24.
4. Pope HG Jr, Wood RI, Rogol A, Nyberg F, Bowers L, Bhasin S. (2014) Adverse health consequences of performance-enhancing drugs: an Endocrine Society scientific statement. *Endocr Rev.* 35(3):341-75.
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1. Anna Jonsson, *The Impact of the Neuropeptide Substance P (SP) Fragment SP1-7 on Chronic Neuropathic Pain*, 2015. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, 198. ISSN 1651-6192

### Agencies that supported the work/Funding 2015

Swedish Research Council - Medicine and Health; Hjärnfonden; Kjell and Märta Beijers Foundation; Carl Tryggers Foundation; Science for Life Laboratory (SciLifeLab); Swedish Council for Working, Life and Social Research; Precision Science System; Swedish Foundation for Strategic Research; Berzelii Centre for Biotechnological Research; Swedish Institute, Visby Program; Disciplinary Domain of Medicine and Pharmacy; The Alcohol Research Council of the Swedish Alcohol Retailing Monopoly; The Research Council of Swedish Criminal Care.

## Other commitments/assignments of group members 2015

Fred Nyberg: Member of the Governmental Advisory Board for Addictive drugs (ANDT-S Advisory Board). Member of the Uppsala University Center for Studies of the Religion in the Society since 2006 and the National Center for Mens Violence against women since 2006. Member of the Board for the Medical Committee of the Swedish Criminal Care. Member of the Board of the Alcohol Research Council of the Swedish Alcohol Retailing Monopoly. Member of the Board of The Research Council of the Swedish Criminal Care. President of the International Narcotics Research Conference (INRC) from 2014-2017. Member of Editorial Board of Scientific journals: Peptides, Open J Endocrinology (Editor in Chief), Pharmacology-on-line, J Musc Skel. Pain. PI at the Uppsala Berzelii Technology Center for Neurodiagnostics (100 milj, SEK 2006-2015); PI at the Linne project Impact of Religion: Challenges for Society, Law and Democracy (50 milj. SEK 2008-2017); PI at the FAS supported project on alcohol effects on cognitive functions (12 milj. 6 years). Swedish Research Council/Medicine and health for peptidergic mechanism in the development of drug dependence: 3 milj SEK for 2015-2017.

Mathias Hallberg: Deputy Chairman of the Board for Education (GRUFF), Faculty of Pharmacy, Member of the Steering Committee/Reference group for the development of the Language Workshop, Uppsala University, Deputy Chairman of the Student Recruitment Committee, Faculty of Pharmacy. Referee: Curr Protein Pept Sci. and The Open Biochemistry Journal.

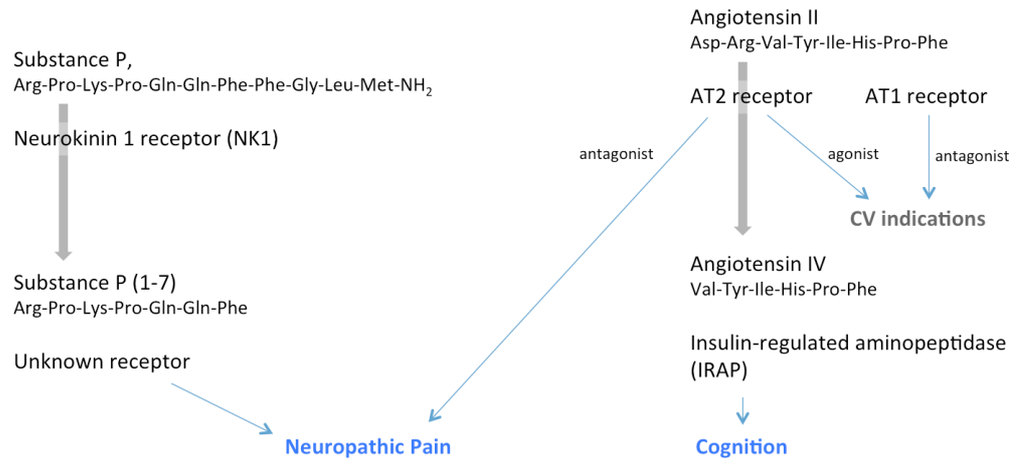
## Projects

(I) Studies on neuropeptides, neurohormones and steroids in relation to opioid sensitivity and chronic pain (including animal experimental models, in vitro cell cultures and clinical studies). In clinical studies the project directs to assessment of genetic polymorphism in various transmitter systems combined with quantification of neuropeptide levels in various body fluids. Peptides chosen for analysis includes the tachykinins and opioid peptides. In animal studies models for nociceptive, neuropathic pain are chosen.

(II) Studies on neuropeptides, neurohormones and steroids (in particular anabolic androgenic steroids = AAS) in relation to drug dependence (including experimental animal models, in vitro cell cultures and clinical studies). In clinical studies the project directs to assessment of genetic polymorphism in various transmitter systems combined with quantification of neuropeptide levels in various body fluids. Peptides chosen for analysis includes the tachykinins and opioid peptides. In animal studies models to investigate opiate tolerance and withdrawal and drug self-administration (in collaboration with other laboratories) are used. Endogenous peptides with high potency to attenuate withdrawal reactions have been identified and serve as basis for design and synthesis of peptides and non-peptides that may be further developed to act as drugs in the treatment of opiate addiction. In studies of effects of AAS on the brain neurochemical technologies (radioimmunoassays, autoradiography, Western blot, etc.) are combined with various behavioural assays.

(III) Studies on the functions of growth hormone (GH) and prolactin (PRL) and their receptors in the central nervous system (including experimental animal models, in vitro cell cultures and clinical studies). Receptors for GH have been identified in the brain in areas of relevance for many of the known effects of GH on the central nervous system (CNS). Beneficial effects of GH on cognitive functions are recorded by the assessment of memory and cognition using the Water maze in conjunction with various neurobiological techniques.

(IV) Studies on synthetic compounds acting on the angiotensin II AT2 receptor and the angiotensin IV receptor, insulin-regulated aminopeptidase. Receptor assays animal models are used to guide synthesis and design of peptide and non-peptide analogues. The objective is a) to discover new classes of analgesics with potential use in the treatment of neuropathic pain and b) to discover unique cognitive enhancers. These two subprojects are derived from findings in the tachykinin and renin-angiotensin systems (RAS) that we have studied for a long period. The substance P metabolite substance P (1-7) alleviates pain and drug-like substances mimicking substance P (1-7) have been discovered that powerfully alleviate pain in animal models. In parallel selective drug-like angiotensin II AT2 receptor antagonists are made as potential analgesics.



**Figure:** Degradation of bioactive neuropeptides may lead to fragments with different biological activities as compared to the parent peptide (Hallberg, 2014 Med Res Rev).

Angiotensin IV (angiotensin 3-8), a fragment derived from the hypertensive angiotensin II, improves memory after i.c.v. injection in rats. Potent drug-like compounds that mimic the effects of angiotensin IV and e.g. increase dendritic spine density in primary hippocampal neuronal cultures and that display neuroprotective effects have been identified. The density and morphological architecture of dendritic spines correlate with parameters associated with cognition. These compounds, aimed at serving as potential cognitive enhancers inhibit the proposed molecular target for angiotensin IV, insulin-regulated aminopeptidase (IRAP). At present considerable research efforts are devoted to these drug discovery programs that now involve both national and international collaborations.

## Molecular Neuropsychopharmacology

Georgy Bakalkin

<http://farmbio.uu.se/research/researchgroups/brdd/molneupsyfarm/>

The main goal for the research group on Molecular Neuropsychopharmacology is to increase the understanding of molecular and epigenetic mechanisms underlying the development of alcoholism and substance addiction, and to develop treatment strategies. Identification of biomarkers and new therapeutic targets for the treatment of developmental psychiatric disorders is also in focus. Specific research activities within the group are described under projects.

### Members of the group during 2015

Georgy Bakalkin, PhD, Senior professor  
 Tatiana Yakovleva, PhD, Senior scientist  
 Hiroyuki Watanabe, PhD, Research scientist  
 Daniil Sarkisyan, PhD, Research scientist  
 Lada Stålhandske, PhD, Research scientist  
 Wei Sun, Ph.D., Postdoctoral scientist  
 Xingwu Zhou, PhD, Postdoctoral scientist  
 Igor Bazov, PhD, Postdoctoral scientist  
 Olga Kononenko, PhD student

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2. Walker BM, Valdez GR, McLaughlin JP, Bakalkin G. Targeting dynorphin/kappa opioid receptor systems to treat alcohol abuse and dependence. *Alcohol.* 2012 Jun;46(4):359-70. doi: 10.1016/j.alcohol.2011.10.006. Epub 2012 Mar 27. Review.

## Agencies that supported the work/Funding 2015

The Swedish Council for Working Life and Social Research; The Swedish Governmental Agency for Innovation Systems; The Swedish Institute (Visby grant for collaboration with Ukraine); The Swedish Research Council; The Swedish Research Council for Environment, Agricultural Sciences and Spatial Planning.

## Other commitments/assignments of group members 2015

Georgy Bakalkin: Editor in *Addiction Biology* journal

## Projects

### *(I) Integrated genetic and epigenetic approach to developmental psychiatric disorders: analysis of human blood and brain*

Environmental stimuli influence the developmental trajectories of neural circuits from birth through adolescence. Exposure to harmful environmental stimuli during these developmental stages may result in increased vulnerability to psychiatric disorders. These effects are suggested to be partly dependent on genotype and mediated by epigenetic mechanisms. We aim to identify biomarkers and new therapeutic targets for the treatment of developmental psychiatric disorders, primarily alcohol dependence. We will perform genome-wide analysis of DNA methylation in blood from 2000 adolescents part of the IMAGEN study on factors that influence mental health in adolescents (<http://www.imagen-europe.com/en/the-imagen-study.php>), and of genotype, DNA methylation and gene expression in brain from circa 400 controls at different developmental stages and adult alcohol dependents. Loci associated with phenotypic traits relevant to alcohol dependence, DNA methylation and gene expression in the IMAGEN sample and diagnosis, DNA methylation and gene expression in the brain sample will be considered candidate biomarkers for alcohol dependence. Mechanisms underlying these associations will be considered candidate therapeutic targets for the treatment of alcohol dependence.

### *(II) Alcohol binge drinking – induced cognitive impairment in health and working life: genetic vulnerability and novel pharmacotherapy*

Alcohol binge drinking patterns and excessive alcohol consumption have severe effects on public health and working life in Sweden and its Nordic neighbors. Delayed impairment of cognitive capabilities and executive functions are the most deteriorating consequence of heavy drinking. We examine the novel pathophysiological mechanism of this impairment, and develop clinically manageable strategies to identify subpopulations at risk and effective pharmacotherapy to treat alcohol-produced cognitive impairment. According to the mechanism, cycles of alcohol intoxication and withdrawal impair cognition through dysregulations in neurotransmission that is controlled by the endogenous opioid systems (EOS) including kappa-opioid receptor (KOPr) and its ligands dynorphins, prodynorphin (PDYN) products. 1) We examine whether genetic variants of the KOPr/*OPRK1* and *PDYN* genes contribute to alcohol use, abuse and dependence, and to alcohol-induced cognitive impairments. Population-based cohorts of young adult monozygotic and dizygotic twin pairs and office workers/social drinkers are analysed. 2) By conducting human post-mortem studies, we are identifying and characterizing dysregulations in the opioid and glutamate neurotransmitter systems, potential targets for pharmacological interference, in human heavy alcohol drinkers; and analyze underlying epigenetic mechanism. We focus on epigenetic modifications that may cause long-lasting behavioral alterations. 3) We plan to use animals models to mimic binge alcohol - induced impairment of cognitive functions and evaluate if these detrimental effects are reversed by available and novel pharmacological means. Characterization of the novel mechanism of alcohol binge drinking - induced cognitive impairment will open new possibilities for identification of subpopulations at risk, and for therapeutic intervention with opioid or glutamate antagonists.

*(III) Pathogenic mechanisms of human neuropeptide mutations: implications for regulation of the anti-reward system*

Much of our knowledge about functions of genes and proteins in human brain has come from analysis of rare human and mouse mutations. Such mutations represent entry points into novel signaling pathways and molecular mechanisms. In collaboration with Dr. Dineke Verbeek, Dept. Genetics, Univ. Med. Center Groningen, The Netherlands, we have identified missense mutations in the human prodynorphin gene (*PDYN*) coding for a precursor to opioid peptides dynorphins (Bakalkin et al., 2010). These mutations cause profound neurodegeneration in the cerebrum and cerebellum underlying the development of spinocerebellar ataxia type 23, a dominantly inherited neurodegenerative disorder. Remarkably, seven out of eight mutations are located in dynorphins, which also have non-opioid neurodegenerative activities. This is the first finding on neuropeptide mutations underlying human neuropathology. The best-established function of the dynorphins in the brain is regulation of the anti-reward system by inducing dysphoria that counterbalances positive hedonic effects of endorphins and enkephalins, and addictive substances. Analysis of *PDYN* mutations could uncover novel molecular and cellular mechanisms of anti-reward - reward regulation that are generally unknown. We focus on two mechanisms. First, the mutations may impair correct folding of *PDYN* molecules in the endoplasmic reticulum, resulting in *PDYN* aggregation, cellular stress and the unfolded protein response. Long-lasting activation of the unfolded protein response by mutant *PDYN*s, or by wild-type - *PDYN* excessively produced under pathological conditions such as substance addiction, may lead to neuronal dysfunction, atrophy and neurodegeneration. Second, *PDYN* mutations may enhance non-opioid pathogenic activities of dynorphins. Our previous analysis suggests that dynorphins may mediate communications between neurons through non-receptor excitatory mechanism. This mechanism may be engaged in pathogenic effects of upregulated wild-type - dynorphins and mutant peptides. Our general goal is to understand pathogenic mechanisms of *PDYN* mutations and, in the following studies to evaluate whether wild-type - *PDYN* and dynorphins engage these mechanisms to regulate reward circuits under normal conditions and in substance addiction when these neuropeptides are excessively produced. We explore pathogenic mechanisms underlying actions of wild-type- and mutant-*PDYN* in cellular and in vitro biochemical/ biophysical studies, and are planning to use in vivo transgenic mice expressing human wild-type- or pathogenic mutant-*PDYN* that have been produced by Dr. Verbeek. Generalized pathological changes including cerebral cortical and subcortical atrophy, and agenesis of corpus callosum in patients carrying *PDYN* mutations emphasize the fundamental role of neuropeptides in brain functions, and propose that the essential molecular mechanisms are affected; the phenomenon that requires detailed investigation. Dissection of these mechanisms is critically important for understanding of reward – anti-reward, substance addiction, depression and chronic pain, all neuropathological conditions in which dynorphins play a critical role, as well as for the neuropeptide and neurodegeneration fields in general because of the novelty of mechanisms identified.

*(IV) Pharmacotherapy of chronic pain. A novel approach that targets the ubiquitin-proteasome system*

Chronic pain including neuropathic pain is an extremely disabling condition with the enormous cost for society and affected individuals and loss in work productivity. This pain is resistant to standard treatment protocols and thus represents a significant unmet medical and social need. We in collaboration with Prof. Frank Porreca group (Dept. Pharmacol., University of Arizona, USA) discovered a critical role of the ubiquitin-proteasome system (UPS), the specialized system for protein degradation, in the development and maintenance of neuropathic pain. This study provides experimental background for novel molecular concept that states that the development and maintenance of neuropathic pain critically depends on regulated protein degradation. We also demonstrated strong pain-killing effects of the UPS inhibitors. This is an especially promising possibility, because proteasome inhibitor velcade (bortezomib) has been recently approved in the US and Europe for the treatment of cancer. We now focus on the selection of the most potent and safe UPS inhibitors for further medical applications, and on molecular and cellular mechanisms of chronic pain. Actions of the UPS inhibitors are apparently mediated through pronociceptive sensory neuropeptides including dynorphins and CGRP.

## Pharmaceutical Bioinformatics

Jarl Wikberg and Ola Spjuth

<http://farmbio.uu.se/research/researchgroups/pb>

During the year, studies for enabling large-scale predictive modeling on high-performance computing and cloud computing resources were carried out. We reported on the utility of Amazon Cloud for predictive modeling, and also established a public catalogue for virtual machine images in bioinformatics (BioImg.org). We evaluated the Big Data frameworks Apache Hadoop for analysis of next-generation sequencing data, and the Apache Spark framework for distributed machine learning. We extended together with collaborators at AstraZeneca (AZ) the Conformal Prediction (CP) applications for drug discovery, and developed methods for interpreting non-linear QSAR-modeling in terms of chemical substructures. With collaborators at AZ and SweTox we also developed the first Apache Spark implementation of CP, which we will further develop and benchmark during the coming years for large-scale predictive modeling in pharmaceutical bioinformatics. Studies in predictive metabolism were advanced, and the XMetDB open access database for xenobiotic metabolism was further developed.

During the year eight novel ring-opened pharngalin limonoids and one novel libiguin type of limonoid were isolated from *Neobegonia mahafalensis* and structurally determined. One of the ring-opened limonoids was found to induce sleep in mice and was named Dodoguin.

In 2015, the PhenoMeNal H2020 project was launched where the group leads WP5: “*Operation and maintenance of PhenoMeNal grid/cloud*” with the aim to provide PhenoMeNal and researchers with the capability to spawn secure Virtual Research Environments with easy access to scalable, interoperable data and tools for data analysis in primarily large-scale metabolomics. A joint project with Uppsala Academic Hospital on the translation of on long-read single molecule sequencing for analysis of mutations in BCR-ABL1 for chronic myelogenous leukemia (CML) was expanded.

### Members of the group during 2015

Jarl Wikberg, Professor  
 Ola Spjuth, Assistant Professor  
 Maris Lapins, PhD, Researcher  
 Wesley Schaal, PhD, Researcher  
 Martin Eklund, PhD, Researcher  
 Iryna Shutava, PhD, Post-Doc  
 Jonathan Alvarsson, PhD student  
 Marco Capuccini, PhD student  
 Samuel Lampa, PhD student  
 Martin Dahlö, PhD student  
 Arvid Berg, Software engineer  
 Valentin Georgiev, PhD, Software engineer  
 Polina Georgieva, PhD  
 Aleh Yahorau, Technician  
 Staffan Arvidsson, Software engineer

## Publications 2013-2015

1. M. Dahlö, F. Haziza, A. Kallio, E. Korpelainen, E. Bongcam-Rudloff, and O. Spjuth. (2015) BioImg.org: A catalogue of virtual machine images for the life sciences. *Bioinformatics and Biology Insights* 2015:9 125-128
2. Eklund, M., Norinder, U., Boyer, S., Carlsson, L. (2015) The application of conformal prediction to the drug discovery process. *Annals of Mathematics and Artificial Intelligence*, vol. 74, ss. 117-132
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11. Alvarsson, J., Eklund, M., Engkvist, O., Spjuth, O., Carlsson, L., Wikberg, J. E., & Noeske, T. (2014). Ligand-based target prediction with signature fingerprints. *Journal of chemical information and modeling*, 54(10), 2647-2653.
12. A. Siretskiy and O. Spjuth. (2014) HTSeq-Hadoop: Extending HTSeq for Massively Parallel Sequencing Data Analysis using Hadoop. In *e-Science, 2014 IEEE 10th International Conference on*, vol. 1, pp. 317–323.
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14. P. Kohonen, R. Ceder, I. Smit, V. Hongisto, B. Fadeel, O. Spjuth, and R. Grafström. (2014) The fields of Cancer Biology, Toxicology and Alternative Methods Development Go Hand-in-Hand. *Basic Clin Pharmacol Toxicol.* 115(1):50-8.
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- & Wikberg, J. E. (2014). Illuminating the origins of spectral properties of green fluorescent proteins via proteochemometric and molecular modeling. *Journal of computational chemistry*, 35(27), 1951-1966.
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### Reviews 2013-2015

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2. A. Claesson and O. Spjuth (2013) On Mechanisms of Reactive Metabolite Formation from Drugs. *Mini Rev Med Chem*. 13(5)
3. Lampa, S., Dahlo, M., Olason, P., Hagberg, J., & Spjuth, O. (2013). Lessons learned from implementing a national infrastructure in Sweden for storage and analysis of next-generation sequencing data. *GigaScience*, 2(1), 9.

### Dissertations 2015

1. Jonathan Alvarsson, *Ligand-based Methods for Data Management and Modelling*, 2015. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, ISSN 1651-6192 ; 200

## Agencies that supported the work/Funding 2015

The Swedish Research Council; Swedish Institute; eSENCE, SeRC

## Other commitments/assignments of group members 2015

Ola Spjuth: Co-Director, UPPMAX High Performance Computing center. Head of Bioinformatics Compute and Storage Facility at Science for Life Laboratory.

## Projects

### *(I) Pharmacology of the libiguins (Jarl Wikberg et al.)*

Studies on the mechanisms of action for the effects of libiguins on sexual behaviour; in part a collaboration with Philippe Rasoanaivo, IMRA, Antananarivo, Madagascar, Gunnar Antoni, PET-centre, Uppsala University and Aigars Jirgensons, IOS, Riga, Latvia.

### *(II) Isolation, structural determination and pharmacology of novel natural and semi-synthetic compounds (Jarl Wikberg et al.)*

Studies devoted to the isolation, structural determination of novel natural compounds and semi-synthetic derivatives therefrom; collaborations with Torgils Fossen, Centre for Pharmacy, Department of Chemistry University of Bergen, Bergen, Norway, Philippe Rasoanaivo, IMRA, Antananarivo, Madagascar, and Aigars Jirgensons, IOS, Riga, Latvia.

### *(III) Proteochemometrics (Jarl Wikberg et al.)*

Studies directed to further development of proteochemometrics: Large scale proteochemometrics and integration of 3D-molecular modelling with proteochemometric modelling. Wet laboratory experimentations in relation to development of antivirals directed towards dengue and Japanese encephalitis viruses; in part a collaboration Mahidol University, Salaya, Thailand.

### *(IV) The Bioclipse Workbench (Ola Spjuth et al.)*

Development of the Bioclipse workbench for e-Science. Main focus is on consumption and visualization of predictive models in drug discovery applications. Collaboration with Lars Carlsson at AstraZeneca R&D.

### *(V) Automated large-scale predictive modeling (Ola Spjuth et al.)*

Studies on predictive modeling applied to large pharmaceutical data sets, requiring high-performance computing and cloud computing infrastructures. Collaboration with Andreas Hellander and Salman Toor at the Department of Information Technology, UU as well as Lars Carlsson, Ernst Ahlberg, AstraZeneca R&D and Ulf Norinder, SweTox.

### *(VI) Predictive toxicology and site-of-metabolism (Ola Spjuth et al.)*

Studies on informatics and predictive modeling in toxicology and site-of-metabolism. Collaboration with Lars Carlsson at AstraZeneca R&D; Egon Willighagen at Maastricht University NL; Nina Jeliaskova at Ideaconconsult BG; Catrin Hasselgren, UCB and Leadscope, US; Roland Grafström at Karolinska Institutet and the OpenTox consortium ([www.opentox.org](http://www.opentox.org)).

### *(VII) Translational Bioinformatics (Ola Spjuth et al.)*

Studies devoted to translating bioinformatics-based analysis of long-read amplicon sequencing data to clinical diagnostics with applications in chronic myelogenous leukemia (CML) and multi-drug resistant bacteria. Collaboration with Åsa Melhus at Department of Medical Sciences and Uppsala Academic Hospital, Adam Ameer at National Genomics Infrastructure, Lucia Cavelier at Department of Genetics and Pathology, and Johan Rung at Clinical Diagnostics, Science for LifeLaboratory, Uppsala University.

## Pharmacometrics

**Mats Karlsson**

<http://farmbio.uu.se/research/researchgroups/pharmacometrics>

Pharmacometric research focuses on nonlinear mixed effects ("population") models. Such models describe data, generally the response-time profiles observed in a clinical trial, by a basic model, accounting for the general structure of the underlying system, and a set of hierarchical variability components, accounting for variability between subjects, within subjects over time and remaining between observation variability. Research at the pharmacometrics group can be divided into four main areas. First, development and evaluation of methods for efficient and robust model building. This involves development of estimation algorithms, methods for model diagnosis and sequential procedures for model building. The result of the research, when applicable, is made available as free software. Second, so-called platform models are being developed for the use in specific therapeutic areas or for particular therapeutic/pharmacological principles. Such a model may involve the time-course of a biomarker or a system of biomarkers during normal, diseased and/or provoked situations. The third research area concerns utilization of the developed models for the purpose of designing studies, deciding upon dosing strategies and other developmental decisions. Fourth, we also do analyses of dose-concentration-response data from trials to understand therapies with existing drugs with the aim of allowing improved therapy.

### Members of the group during 2015

Mats O Karlsson, Professor  
 Lena Friberg, Professor  
 Andrew Hooker, Senior Lecturer  
 Elisabet Nielsen, Senior Lecturer  
 Ulrika Simonsson, Senior Lecturer, Docent  
 Martin Bergstrand, Researcher  
 Nick Holford, Researcher  
 Siv Jönsson, Researcher  
 Kristin Karlsson, Researcher  
 Maria Kjellsson, Researcher, Docent  
 Joakim Nyberg, Researcher  
 Elodie Plan, Researcher  
 Sebastian Ueckert, Researcher  
 Jonas Häggström, Researcher  
 Kajsa Harling, System Developer  
 Rikard Nordgren, System Developer  
 Gunnar Yngman, Research assistant  
 Britt Jansson, Lab Engineer  
 Yasunori Aoki, Post-doctoral fellow  
 Chenhui Deng, Post-doctoral fellow  
 Margreke Brill, Post-doctoral fellow  
 Thomas Dorlo, Post-doctoral fellow  
 Gopichand Gottipati, Post-doctoral fellow  
 Elham Haem, Post-doctoral fellow  
 Anna Largajolli, Post-doctoral fellow  
 Wanchana Ungphakorn, Post-doctoral fellow  
 Sven von Dijkman, Post-doctoral fellow  
 Sebastian Wicha, Post-doctoral fellow

Gülbeyaz Yildiz, Post-doctoral fellow  
 João Abrantes, PhD Student  
 Oskar Alskär, PhD Student  
 Brendan Bender, PhD Student  
 Henrik Bjugård Nyberg, PhD Student  
 Salim Bouchene, PhD Student  
 Ari Brekkan, PhD Student  
 Chunli Chen, PhD student  
 Steve Choy, PhD student  
 Oskar Clewe, PhD Student  
 Anne-Gaëlle Dosne, PhD student  
 Benjamin Guiastrennec, PhD Student  
 Mendel Jansen, PhD Student  
 Ana Kalezic, PhD student  
 David Khan, PhD Student  
 Anders Kristoffersson, PhD Student  
 Brigitte Lacroix, PhD Student  
 Jennifer Leohr, PhD student  
 Siti Maisharah Sheikh Ghadzi, PhD Student  
 Moustafa Mahmoud Abdellatif Ibrahim, PhD student  
 Rikke Meldgaard, PhD Student  
 Ida Netterberg, PhD Student  
 Jesmin Permal, PhD Student  
 Patanjali Ravva, PhD Student  
 Emilie Schindler, PhD Student  
 Marina Senek, PhD student  
 Eric Strömberg, PhD Student  
 Elin Svensson, PhD Student  
 Robin Svensson, PhD Student  
 Anders Thorsted, PhD Student  
 Shijun Wang, PhD Student  
 Gustaf Wellhagen, PhD Student  
 Anneke Hesselning, visiting Professor  
 Juha Vakkalainen, visiting Scientist  
 Vincent Aranzana-Climent, visiting PhD Student  
 Tore Bjerregaard Stage, visiting PhD Student  
 Bruna Gaelzer Silva Torres, visiting PhD Student  
 Rasmus Vestergaard Juul, visiting PhD Student

### Publications 2013-2015

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- Neuroimaging I. Improved utilization of ADAS-cog assessment data through item response theory based pharmacometric modeling. *Pharm Res.* 2014;31(8):2152-65.
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### Dissertations 2015

1. Brigitte Lacroix. *Pharmacometric Modeling in Rheumatoid Arthritis*, 2015. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, 199. ISSN 1651-6192.
2. Anders Kristoffersson. *Study Design and Dose Regimen Evaluation of Antibiotics based on Pharmacokinetic and Pharmacodynamic Modelling*, 2015. Digital Comprehensive Summaries of Uppsala Dissertations from the Faculty of Pharmacy, 206. ISSN 1651-6192.

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### Other commitments/assignments of group members 2015

Lena Friberg: Deputy Editor-in-chief, CPT: Pharmacometrics & Systems Pharmacology; Organizing committee, chair scientific program, PAGE conference, Crete 2015 and Lisbon 2016; Executive committee, WCoP 2016 conference; Scientific committee, 1<sup>st</sup> international workshop on dose optimization strategies for targeted drugs, Amsterdam 2015; Faculty of Pharmacy: Board member

Andrew Hooker: Organizing committee, PODE conference

Siv Jönsson: Department Board Member; Organizing committee, chair scientific program, PAGE conference, Crete 2015 and Lisbon 2016; Organizing committee for The 13th Symposium on Pharmacokinetics and Drug Metabolism: Oligonucleotide-based therapeutics– New challenges in evaluation pharmacokinetic, pharmacodynamics and safety properties; Gothenburg 2015; Board Member: Swedish Academy of Pharmaceutical Sciences, Section for Pharmacokinetics and Drug Metabolism

Mats Karlsson: Deputy Head of Department; Department Board Member; Equal Opportunities Board Member; Editor Journal of Pharmacokinetics and Pharmacodynamics; Editorial Board on Clin Pharmacol Ther, Eur J Pharm Sci, Basic Clin Pharmacol Toxicol, CPT: Pharmacometrics & Systems Pharmacology

Maria Kjellsson: Department Board Member, Board Member: Swedish Academy of Pharmaceutical Sciences, Section for Pharmacokinetics and Drug Metabolism

Elisabet Nielsen: Executive Committee member, EPASG, ESCMID PK/PD Study Group

Ulrika Simonsson: Board Member: CPTR Regulatory Science Consortium, Critical Path to TB Drug Regimens. Clinical Disease Progression Modeling Workgroup. <http://cptrinitiative.org/>; Board member of IF's stiftelse, Swedish Academy of Pharmaceutical Sciences.

## Projects

### Methodological research

*(I) Optimal design/Clinical trial design (Martin Bergstrand, Andrew Hooker, Kristin Karlsson, Mats Karlsson, Joakim Nyberg)*

There are two principle ways in which models can be used to evaluate and optimize clinical and pre-clinical experiments. The first is by simulation of a set of proposed designs from a model (or set of models), followed by evaluation of those resulting data sets using a metric of interest. The simulations, repeated many times with different random seeds, provide information about the expected results of various different designs (for example, measures of the precision and bias of parameter estimates, or the power to detect a drug effect of a specific size). With this methodology we have investigated, for example, differences in randomization schemes for dose-finding trials where it was found that dose-randomized trials are more powerful in characterizing the underlying relation compared to concentration-randomized trials. This increase in power can be achieved with, in most instances, a similar or lower number of observed side effects.

The second way of evaluating and optimizing trial designs is through the use of optimal experimental design methodologies. These methods often rely on calculations of an Information Matrix (e.g. the Fisher Information Matrix), which characterizes the information content of any possible design. Each design evaluation is much quicker than clinical trial simulation, thus one can investigate the landscape of possible designs (within constraints) potentially available for an experiment, and even optimize a design based on this information. We have developed methods and software (PopED) that utilize these methods with both local and global design criteria (e.g. E-family optimal designs, which take into account the underlying uncertainty in a pharmacometric model description of a biological system. Additionally, while optimal design has previously focused on optimization of sampling times in an experiment, we have extended the methodology to apply to other aspects of trial designs, such as the dose administered or the length of run-in, treatment and wash-out phases of an experiment. Further, we have extended optimal design methodology to optimize a study for power, as opposed to the traditional optimization based on model parameter estimation uncertainty.

The two methods of evaluating and optimizing trial designs can be combined to evaluate and explore adaptive optimal designs. In these types of trial designs, interim analyses are used to update models used to describe the system being investigated and then to use this information to re-optimize the next cohort of patients coming into a study. With combined simulation/optimization one can explore the adaptation and optimization rules one will use in an adaptive trial. We are currently developing such a tool ([MBAOD](#)), and are investigating the use of such designs in, for example, pediatric bridging studies.

*(II) Model building and parameter estimation (Andrew Hooker, Mats Karlsson)*

Pharmacometric models are based on (patho-)physiological and pharmacological knowledge. The complexity and heterogeneity of biological data makes the knowledge about, and development of, statistical data analysis methods a central part of this scientific field. There are many benefits to using pharmacometric models in the analysis of data from clinical trials, for example the ability to handle sparse data and to integrate different types of observations into one model; however, these models are complex and intrinsically non-linear which presents technical challenges in model building and estimation.

One main challenge is to reduce the time it takes to develop these models. With complex, non-linear models and data from a clinical trial that can have thousands of data points from hundreds of patients with multiple response variables, computer runtimes become non-ignorable. Generally, run-times can be divided into short (minutes), intermediate (hours to days) and long (days to months). The number of runs in a complete analysis tends to range between 30 and many hundred. We are investigating the implementation and automation of important modelling tasks through the use of new algorithms developed in our research group. Additionally, we are developing new methods of model building and

new algorithm development that can shorten run times and the number of steps needed in the model building process.

Other areas of active research include the influence on parameter estimates of single observations and rational and statistically correct algorithms for adding explanatory variables, .i.e. covariates, to the models.

*(III) Diagnostic tools (Andrew Hooker, Mats Karlsson, Siv Jönsson)*

A main problem for complex pharmacometric models and data is to evaluate how well the models fit the data. For the assessment of an adequate description of data, methods of evaluations can be based on model predictions, residuals, simulations from the model, simulations followed by evaluation and simulations followed by full re-estimation. We are developing diagnostic tools based on all of these principles and for both continuous and categorical type data. Knowing about model and parameter uncertainty is often crucial in model-informed decision-making. We are developing methods for diagnosing existing methods of uncertainty estimates as well as developing new methods for both model and parameter uncertainty and how these can be applied in decision making.

*(IV) Software development (Andrew Hooker, Mats Karlsson, Kajsa Harling, Rikard Nordgren)*

One integral part of all of our research activities is the implementation of the methods developed in freely available software to facilitate a wider and consistent use of the new algorithms. Software developed by the group is PopED (<http://poped.sf.net>), PsN (<http://psn.sf.net>) and Xpose (<http://xpose.sf.net>).

*(V) Pharmacodynamic modelling of discrete outcomes (Andrew Hooker, Mats Karlsson, Elodie Plan)*

For many diseases, the main outcome is of discrete nature: stage category, symptom severity, number of events, or occurrence of events. In pharmacometrics, we distinguish non-ordered categorical data, ordered categorical data, count data, and (repeated) time-to-event data. Models handling this type of data are based on probabilities.

In this project we aim to describe disease progression and treatment exposure-response, and to develop new models for simulations of future studies. The time course of sleep stages and its relation to placebo and drug effects has been analysed using Markov models in patients with insomnia. Pain scores rated on a Likert scale by neuropathic patients have been modelled by including features for under-dispersion and serial correlation. Daily numbers of seizures have been used in the investigation of over-dispersion and Markov patterns in count data. Simultaneous characterization of drug effect on severity and time to acid reflux events has been made possible by a new type (repeated time-to-categorical event) of model and resulted in good simulation properties and high power to detect a drug effect. Parametric time-to-event models have been compared to semi-parametric Cox proportional hazard models implemented in NONMEM and methodology to simulate large scale unbiased (repeated) time-to-event data has been developed.

In parallel we also investigate the performance of available estimation methods with discrete models. We have pointed out the fact that the Laplacian estimation method in NONMEM and NLMIXED results in biased parameter in situations with non-even distributions of the response categories. In another study the Laplace method produced accurate parameter estimation for Poisson models, with or without Markov elements and mixture distribution, whereas we identified a small bias in the random effect of zero-inflated Poisson, generalized Poisson and negative binomial models. The performance of the SAEM and importance sampling have been shown to be generally higher than Laplace in repeated time-to-events models where the frequency of individuals with events was low, while at high frequencies all methods were equal in performance. We have also conducted a study investigating all methods available in NONMEM version 7 for all types of discrete models, where we highlighted the overall fast and robust results obtained with Laplace.

Currently we are interested in handling the within-individual variability for discrete models. More specifically we adapted methods to handle inter-occasion variability in count models. Since count data are often collected on a regular basis (e.g. daily) in clinical trials, the time and length of potential occasions is not predefined but can be estimated through dynamic implementations. Alternatively the noise present in these types of records can be characterised using stochastic differential equations. Additionally we study ways to characterize underlying discrete processes with the implementation of hidden Markov models (HMM) in NONMEM. When there are suspected transient or definite transitions from an unobserved state to another one (low to high epileptic activity for example), these models permit to estimate the transition probability, on which time and drug effects can be investigated. Finally the individual sequences can be retrieved post-hoc via a Viterbi algorithm.

*(V) Mechanism-based pharmacokinetic models (Martin Bergstrand, Mats Karlsson)*

Clinical pharmacokinetic experiments typically measure drug concentrations in plasma only. As a consequence, pharmacokinetic models, used in drug development, aim to describe observations of drug concentration in plasma with minimum model complexity. Such models have limited capacity for extrapolations and to predict concentration-time profiles in tissues and organs. Also, mechanistic insight about drug disposition dependence of factors related to individual organs and tissues may not be possible to incorporate in a fully satisfactory manner. Physiologically-based pharmacokinetic (PBPK) models, which have a structure based on anatomy, can provide predictions in tissues and organs. However, because of their complexity, such models are traditionally not used for analysing clinical data. We have demonstrated that PBPK model parameters can be estimated based on clinical observations and are currently investigating the possibilities for improving such a combined “bottom-up and top down approach”. We have showed that for a relatively simple PBPK model such analyses can become feasible by using informative prior information about physiology and drug-related parameters. To further improve such an approach we are combining information about (co-)variability in organ and tissue properties from a data base representing physiological values for about 30000 subjects, tissue composition models and models for relating drug molecular properties and *in vitro* data to expected behaviour in tissues and organs.

A mechanism-based approach has also been applied to better understand the processes involved in oral absorption. Several factors influencing oral absorption vary along the GI tract, e.g. pH, active influx and efflux transporters, gut wall metabolism. Models characterizing gastro intestinal transit and absorption properties along the GI region can be especially useful in the case of modified release formulations and/or substances with slow dissolution rate. Mechanism based models have been used for prospective population predictions of plasma concentrations based on *in vitro* dissolution data.

*(VI) Pharmacometric modelling of biologic medicinal products (Siv Jönsson, Andrew Hooker, Mats Karlsson)*

Biological medicinal products are becoming an important contributor in the treatment of many diseases, e.g. multiple sclerosis, rheumatoid arthritis, cancer, psoriasis. Characterization of biologics benefit from pharmacometric modelling, since they exhibit complex disposition characteristics, quite different to the processes and pathways utilized for small molecules, e.g. monoclonal antibodies exhibit target mediated drug disposition (TMDD).

Available TMDD models describe the formation of one complex (a dimer), but in reality further complexes may be formed (trimers, hexamers, etc), as described for IgE and omalizumab. We aim to explore and develop alternative TMDD models for the interaction between a target and drug, taking into account formation of different complexes. Furthermore, to explore study design options for studies in different stages of drug development, optimal design methodology is applied to TMDD models.

## Applied research/Disease areas

### (I) Antibiotics (Lena Friberg, Mats Karlsson, Elisabet Nielsen)

Antibiotics are considered one of the greatest discoveries of modern therapeutic medicine and have turned previously fatal diseases into treatable minor illnesses. Today, treatment failures due to multidrug-resistant bacteria are becoming more frequently observed. The evolution of resistance is a natural phenomenon; however, the use and misuse of antibiotics will accelerate this phenomenon. We aim to advance the understanding of pharmacokinetic-pharmacodynamic relationships for antibiotics of value for improving dosing recommendations and minimizing resistance development.

Today, dosing regimens are typically selected based on PK/PD indices that discard information on dynamic changes in the drug-bacteria interaction. Mechanism-based models describing time-kill curves from *in vitro* experiments form the basis for the modelling. The models have shown to be applicable across drugs and bacteria strains (including clinical isolates), for both static and dynamic concentration experiments, for different sizes of start inocula, for mixtures of wild-type and resistant bacteria and for drug combinations, and for predicting competition experiments of wild-type and mutants. Optimal experimental design techniques are applied to find experimental protocols that increase the efficiency of both static and dynamic time-kill curve experiments and in clinical studies of colistin.

For colistin and meropenem, PKPD models based on *in vitro* data have been able to successfully predict previously determined pre-clinical *in vivo* PK/PD indices. Simulations have shown the PK/PD indices to be sensitive to experimental design, misspecification of the MIC, and differences in PK characteristics. The use of a mechanism-based PKPD modelling approach in dose selection has been suggested for increased robustness and extrapolation potential, especially for special patient populations.

Colistin has regained interest in recent years as a promising drug to overcome antibiotic drug resistance. With an in-house developed LC-MS-MS method we can quantify colistin and its prodrug CMS in both clinical plasma samples and in samples from *in vitro* experiments. Developed pharmacokinetic models have been applied to suggest loading doses of 6-12 MU. Whole-body Physiology-based Pharmacokinetic (WBPBK) models for CMS, colistin and ciprofloxacin have been developed based on data from various sources, including patients, healthy volunteers and several animal species. Such a model can be used to understand the time-courses of the antibiotics, and thereby the bacterial killing, in different tissues.

### (II) Infectious diseases (Martin Bergstrand, Mats Karlsson, Ulrika Simonsson)

Plasmodium falciparum, the human immunodeficiency virus (HIV), and Mycobacterium tuberculosis (TB) are three devastating pathogens in tropical areas. Due to the geographical overlap of malaria, HIV and TB prevalence, the diseases are likely to co-exist in a great number of individuals. For these individuals, there is an obvious need for concomitant use of antimalarial, antiretroviral and antitubercular drugs. Drug-drug interactions may result from concurrent administration of drugs leading to diminished therapeutic efficacy of or increased toxicity from one or more of the administered drugs. Drug-drug interactions are an important concern in the management of patients with HIV because of the large number of antiretroviral drugs and other drugs that are required by these patients for the management of co-morbidities and opportunistic infections. Combination therapy has also been introduced in the management of malaria and TB, to overcome drug resistance. The exposure response relationships are not fully understood for the management of malaria and TB whereas there is more knowledge in the field of HIV. Better understanding of the exposure response relationships in malaria and TB is urgently needed. Our research focus also in suggesting dosing recommendations for children based on scaling from adult data or evaluation of pediatric data. Limited information is available in the literature on drug-drug interactions between the artemisinin antimalarial drugs and other drugs such as antiretrovirals or antitubercular drugs; consequently, the extent of such interactions is not fully known.

Activities of CYP enzymes and consequently drug-drug interactions occurring due to their inhibition or induction can be studied by using probe drugs. The pharmacokinetics of probes and drugs under investigation can be described by mathematical models in order to characterise and quantify the interaction.

We have developed enzyme turnover models to describe the time course of induction of different CYP450 enzymes by different artemisinin derivatives. One focus of the work has been to compare the potential for drug-drug interactions among the artemisinin drugs to choose a derivative that is suitable for combination therapy from a drug-drug interaction perspective.

One commonly used antitubercular drug is rifampicin which is known to induce CYP450 enzymes. Our work has involved quantitative analysis by modelling the pharmacokinetics of other drugs metabolised by these enzymes and which are used in HIV treatment. The need for potential dose adjustment has been evaluated for example nevirapine and lopinavir. For example, the population pharmacokinetics of nevirapine in HIV-infected patients taking nevirapine-based antiretroviral therapy in the presence and absence of the antitubercular drug rifampicin has been evaluated.

Pharmacokinetic drug-drug interactions can possibly be compensated for by dose adjustment of the target drug. The developed nevirapine model was used for simulations of different doses of nevirapine which revealed that increasing the dose of nevirapine to 300 mg twice daily elevated nevirapine concentrations above sub-therapeutic levels in most patients, with minimum exposure above the recommended maximum concentration. We have also investigated the population pharmacokinetics of lopinavir in TB/HIV co-infected children taking lopinavir/ritonavir in a ratio of 1:1 in the presence of the antitubercular drug rifampicin, with that of lopinavir in HIV-infected children taking lopinavir/ritonavir in a ratio of 4:1. Increasing the ritonavir dose in the TB/HIV co-infected children resulted in model predicted lopinavir trough concentrations above the recommended minimum therapeutic concentration.

Malaria was estimated to cause 800,000 deaths and 225 million cases worldwide in 2010. The mortality has recently been decreasing and is expected to decrease further due to more widespread use of effective treatment with drugs from the artemisinin class. However, a possible emerging resistance to these drugs might counteract this positive development. Drug resistance has appeared repeatedly within the area of malaria chemotherapy and drastically hampered our ability to fight the disease. It has been hypothesised that such development could have been avoided and or delayed with a better treatment regimen. We are conducting research with the aim of optimising anti-malarial treatment regimens with regards to both short and long-term outcome. Pharmacometric models have been used for translational simulations of expected treatment outcome in vulnerable populations such as children and pregnant and to optimize the treatment regimen.

### *(III) Type 2 Diabetes (Mats Karlsson, Maria Kjellsson)*

Diabetes is a chronic disease, affecting more than 220 million people worldwide and the “diabetic epidemic” is projected to affect 366 million people by 2030, of which more than 90% will suffer from type 2 diabetes (T2D). This disease occurs when the body does not effectively respond to insulin and is unable to produce enough insulin to account for the inefficient use of insulin. This results in elevated blood glucose levels which is toxic and lead to complications; e.g. cardio-vascular diseases (CVD). The aim with most anti-diabetic treatment is to bring glucose levels down to healthy levels. Diagnosis of diabetes is mainly based on fasting plasma glucose (FPG) but also on glycosylated haemoglobin (HbA1c). The success of treatments is assessed on both FPG and HbA1c but also on dynamic glucose after provocation studies.

Provocation studies are used to characterize the functionality of the glucose-insulin system and could vary greatly in design from clamping of glucose or insulin by variable rate infusions, graded glucose infusions, intravenous bolus administration of glucose or insulin and oral administration of glucose solution or meals. We have developed several integrated models with simultaneous analysis of dynamic glucose and insulin after such provocations. These models, which include production, disposition and control (homeostatic) mechanisms of the system, have shown to realistically simulate outcomes of short dynamic glucose provocations at the raw data level. These models have been developed to describe both healthy subjects and patients with T2D as well as for the effect of incretin hormones on gastric emptying and insulin secretion, characterizing pre-hepatic insulin, mechanisms of oral glucose absorption as well as inclusion of exogenous insulin for insulin treated patients. We are focusing our current research on

including mechanism of glucagon release and glucagon effects on glucose and insulin as well as a fully mechanistic whole-body integrated glucose homeostasis model. As diabetes is a progressing disease we are also exploring the possibility of describing how glucose and insulin after provocation studies change over time for a pre-diabetic population.

Long-term clinical trials in T2D patients mainly focus on HbA1c. HbA1c is the fraction of the haemoglobin, in red blood cells (RBC), that has been glycosylated. This is a naturally occurring reaction depending on the amount of glucose in plasma; the higher the glucose concentration, the higher the HbA1c. As the life-span of RBCs ranges from 2 to 4 months, the HbA1c supplies a measurement of the sustained glycaemic control. Several models have been developed in-house to describe the relationship between HbA1c and FPG or mean plasma glucose (MPG) either using an empirical description or a mechanistic approach using knowledge about RBC life-span. Also models acknowledging the mechanism of insulin sensitivity, glucose production and disposition and changes in beta-cell mass or function in relation to weight loss has been developed. Currently we are investigating in what conditions which models perform the best in terms of power to detect drug effects and accuracy in estimates.

The overall endpoint of most anti-diabetic treatments is to lower the risk of long-term complications, such as CVD, retinopathy and chronic kidney disease. Long term studies commonly involve assessments of the risk of CVD in relation to elevated levels of HbA1c or FPG. We are developing parametric risk models, using registry data, quantifying the relationship between CVD and time-varying covariates such as HbA1c and other predictors of CVD, i.e. blood pressure, blood lipids, etc.

All models have been developed for the purpose of being used to quantify changes in the system following interventions (drug administration, diet changes, etc) and associate these changes with known or hypothesized mechanisms of impact of the system. Further the models are intended as tools for hypothesis generation regarding single or combined interventions as well as clinical trial design optimization.

#### *(IV) Oncology (Lena Friberg, Mats Karlsson)*

Within the oncology area, we are working on PK and PKPD models describing the time-courses of biomarkers drug-induced toxicity, tumour size measurements (SLD, sum of longest diameter), tumor activity measurements (SUV, standard uptake value) and overall survival. The models are aimed to be mechanism-based and be applicable across different drugs, and thereby valuable in development of new and existing drugs, including individualization of the therapy. By integrating information of different variables in to a modelling framework the variables' relations and predictive value can be tested, and a better overview of both desired and adverse effects from a changed dosing regimen can be obtained. The models can also be used to explore different concepts of study design in oncology. This type of modelling framework, including biomarkers, side-effects, tumor response and survival, has been developed for sunitinib in gastrointestinal stromal tumors and is being developed for axinitib in renal cell carcinoma. Different metrics of tumor size, both constant and time-varying, as well as one dimension (diameter) vs. three-dimensional (volume), is being compared for predicting overall survival.

Projects are on-going around extensions and applications of a semi-physiological model that can describe neutropenia for numerous anticancer drugs. As an example, the interaction between G-CSF and neutrophil, as well as the time-courses of IL-6 and CRP after initiation of chemotherapy, has been characterized based on generated data from a clinical study. In addition, the predictive value of frequent measurements of neutrophils, that would be possible with a device for home measurements, is being explored based on simulations from the semi-physiological model. The time-courses of thrombocytopenia have been characterized for abexinostat and TDM-1 using the semi-physiological model as basis. For TDM-1, an integrated model that in addition to platelets includes the liver enzymes ASAT and ALAT is being developed, where the driver for the toxicity is being explored.

Patient Reported Outcomes (PRO) is an increasingly used component for comparison of drug treatments during oncology drug development. Item Response Theory has shown to be a promising methodology to analyse and interpret these type of data where questions are grouped into different categories.

*(V) Progressive disorders (Lena Friberg, Andrew Hooker, Mats Karlsson, Elodie Plan, Ulrika Simonsson)*

Progressive neurological disorders such as Alzheimer's Disease and Parkinsons Disease represent challenges in many respects. For characterising drug effects, the slow progression may make it difficult to assess in what aspects the treatment impacts the disease, in a disease-modifying or symptomatic manner. How to best design and analyse trials in these diseases, but also diseases with other progression pattern such as Multiple Sclerosis and Schizophrenia and the subject of such evaluations. Further, the severity of most of these diseases are followed over time using clinical scoring scales which are formed from tests or questions probing individual aspects of the disease. Rather than using the overall clinical scale, we are developing methodology and models based on item response theory (IRT). In these each item (i.e. question or test) is modelled individually as a latent variable of the disease-related disability.

*(VI) Immunological disorders (Lena Friberg, Mats Karlsson)*

Multiple Sclerosis is both a complex and chronic neurological disease of the CNS. The natural course of MS is slow and difficult to monitor clinically. The overall aim of this project is to establish the first population, data driven, MS disease progression model in order to construct a mathematical modelling platform where the interplay between the majority of relevant aspects of the disease, such as time course of disability progression, relapse rate dynamics, time course of the imaging data, time course of lymphocytes and population characteristics are incorporated. Model building involves sequential development of (i) separate models for all components of interest (disease progression, relapse rate dynamics, MRI dynamics and lymphocytes including CD4+, CD8+) and (ii) the covariate model, which explore the underlying patient factors influencing within- and between-subject variability in treatment response. The final anticipated model shall enable simultaneous characterization of the interplay between relapse rate dynamics, total CD4+ and CD8+ lymphocytes and their ratio, and MRI readout dynamics in the evolution of disease and, most importantly, link the time-course of MRI and clinical outcome (relapse rate and disability) and lymphocyte data.

In rheumatoid arthritis a range of variables of the disease are summarized into a clinical endpoint for evaluation of drug response - the dichotomous ACR20 score. Integrated longitudinal transition models with dropout are useful for understanding the outcome of different dosing schedules and by expanding such models to also include the more stringent ACR70 criteria more information can be preserved. To increase the information on the concentration-effect relationship in the available data, a longitudinal transition model describing the probability of ACR20, ACR50 and ACR70 responses have been developed.

*(VII) Dose individualisation in hemophilia (Siv Jönsson, Elisabet Nielsen)*

Hemophilia is a group of hereditary genetic disorders impairing blood coagulation. Hemophilia A and B refer to the coagulation factor VIII and IX deficiency, respectively, whereof hemophilia A is the more common although rare: in Sweden 14 of 100 000 boys and men have hemophilia, whereof 80 % has hemophilia A. By substitution therapy with the coagulation factors, on-demand or prophylactically, the disease status can be controlled. In prophylactic treatment of factor VIII and IX, pharmacokinetic (PK) tailored dose individualization, i.e. the individual dose is based on Bayesian estimation using pharmacokinetic information and a population PK model, is promoted and appears to be a good approach for reducing the total doses administered.

We are involved in one project where the feasibility of PK tailored dosing for prophylactic FVIII treatment is evaluated in routine clinical practice. Our contribution in the project refers to the Bayesian estimation of the dose and in the future re-development of previously existing population PK models. Furthermore, for factor IX we have re-developed a population PK model and, based on the model, explored alternative clinical practically sampling schedules to be used in PK tailoring.

## **Steroid Biochemistry (previously called Steroid P450)**

**Maria Norlin & Kjell Wikvall**

<http://farmbio.uu.se/forskning/researchgroups/steroidbiokemi/>

Our research is focused on the properties and regulation of enzymatic processes involving steroids; in particular those mediated by cytochrome P450 enzymes. Steroids and steroid-related genes and enzymes, producing biologically active metabolites, are vital for human physiology. Many steroids or steroid-related compounds are used as drugs. We apply our expertise in biochemistry and molecular biology to obtain knowledge of relevance for drug discovery and development and drug management. In particular we focus on steroids. We study metabolic conversions, gene regulation and cellular effects of steroids.

Primary missions and goals are to find new drug targets, understand the mechanisms for effects of steroids and steroid-related enzymes, compare effects of endogenous molecules and drugs, examine effects of drugs and drug candidates on steroid-related processes and study molecular mechanisms behind adverse drug effects. Our research activities, focusing on basic research, provide important information that increases the understanding of mechanisms for steroid action. This should improve drug management (e.g. counteract/avoid adverse side-effects) and can be used for development of novel, better drugs.

### **Members of the group during 2015**

Kjell Wikvall, MD, PhD, Professor

Maria Norlin, PhD, Associate Professor

Ida Emanuelsson, PhD student

Mokhtar Almokhtar, PhD student

Ahmad Zayny, PhD student

Kerstin Rönqvist, research assistant

### **Publications 2013-2015**

1. Wicher G, Norlin M. Estrogen-mediated regulation of steroid metabolism in rat glial cells; effects on neurosteroid levels via regulation of CYP7B1-mediated catalysis. *J Steroid Biochem Mol Biol.* 145, 21-27 (2015)
2. Almokhtar M, Wikvall K, Ubhayasekera SJ, Bergquist J, Norlin M. Motor neuron-like NSC-34 cells as a new model for the study of vitamin D metabolism in the brain. *J Steroid Biochem Mol Biol.* 158:178-188 (2016) *Doi*10.1016/j.jsbmb.2015.12.010. *Epub* 2015 Dec 15.

### **Reviews 2013-2015**

1. Nebert DW, Wikvall K, Miller WL. Human cytochromes P450 in health and disease. *Philos Trans R Soc Lond B Biol Sci.* 2013 Jan 6;368(1612):20120431. doi: 10.1098/rstb.2012.0431. Print 2013 Feb 19.

### **Other commitments/assignments of staff members 2015**

Kjell Wikvall: Chair of the Scholarships committee for the Faculty of Pharmacy; Member of the Scholarships committee for Uppsala University; Study director in Pharmaceutical Biochemistry

Maria Norlin: Senior member of the Departmental Committee for PhD education (FUG).

## Projects

### *(I) Functions of steroids and steroid-metabolizing enzymes for hormonal signalling and cellular viability*

This research concerns steroids involved in hormonal signalling, sex hormone biosynthesis and brain function. The studies are focused on physiological and pharmacological control of steroid levels, effects of metabolic events and regulation of gene expression. The project concerns endogenous steroids, steroid drugs and drugs affecting steroid hormone receptors such as SERMs (selective estrogen receptor modulators) and vitamin D analogues.

The studies include mechanisms of importance for estrogenic and androgenic signalling. Some examples of steroids of interest are pregnenolone and dehydroepiandrosterone (DHEA). These steroids are well-known as precursors for androgens and estrogens but also play roles in brain function and in connection with cell growth and viability, e.g. in neurodegenerative processes. Tissue-specific metabolism of DHEA leads to a number of metabolites with differential effects on cell functions. Enzymes that regulate the concentration of neuroactive steroids in the brain may be future targets for therapy of importance for abnormal cell growth, immune function or in neurodegenerative conditions. Some of our current studies involve enzymes and genes of importance for the levels of neurosteroids such as DHEA and pregnenolone in neurons and glial cells. Regulation of hormone metabolism in the nervous system by endogenous steroids and pharmaceutical compounds is also studied.

Steroids may affect growth and differentiation in several tissues. Thus properties of steroids may be of interest in a wide range of normal and disease conditions, e.g. in neuroprotection or cancer therapy. We study effects of steroids such as enzymatically formed oxysterols (cholesterol derivatives), hormones and vitamin D-like compounds on cellular survival and growth. These studies particularly focus on cells of the central nervous system.

### *(II) Bioactivation and metabolism of vitamin D including vitamin D-mediated effects on cellular function*

This research is focused on enzymes and genes of importance for vitamin D bioactivation, metabolism and function. Effects on these processes by endogenous and pharmacological compounds are studied. Many drugs result in adverse side effects, including increased risk of bone disease. We study the mechanisms by which anti-inflammatory, anti-viral and anti-epileptic drugs affect cells, with particular focus on the bone and the central nervous system. Vitamin D is needed for regulation of calcium levels in the body and vitamin D deficiency leads to skeletal diseases such as rickets in children and osteomalacia/osteoporosis in adults. The biologically active form, 1,25-dihydroxyvitamin D, is formed through metabolic activation. The activated form of vitamin D blocks cell division and increases cell differentiation. Vitamin D analogues (synthetic compounds with vitamin D-like effects) are used in the treatment of psoriasis and are of interest in development of new cancer therapy. In addition, epidemiological data during recent years have indicated that vitamin D may have many more targets than previously known. Altered vitamin D levels in plasma have been linked to a number of different conditions including neuropsychiatric and neurodegenerative disease. For these reasons, it is important to obtain more knowledge about the enzymes and genes in activation and metabolism of vitamin D and the roles of the formed metabolites. Part of this project is focused on cellular effects of vitamin D and vitamin D analogues in order to explore previously unknown functions of these compounds.

## Translational Pharmacokinetics/Pharmacodynamics

Margareta Hammarlund-Udenaes

<http://farmbio.uu.se/research/researchgroups/tPKPD>

Our research focuses on understanding the fundamental and functional aspects of pharmacokinetics-pharmacodynamics (PKPD) in health and disease. Within the area of translational PKPD we address conversion of pharmacokinetic data from the preclinical to the clinical setting. This is strongly needed for optimizing drug discovery processes and promoting efficient treatment of CNS diseases. We are interested in brain drug delivery in relation to transport processes of both small and large molecules across the blood brain barrier (BBB), where a quantitative role of nanocarriers is specifically studied. Using the concepts and methodologies developed for the brain we are also exploring drug distribution in the lung. This is important for the design and evaluation of lung targeted drugs which may be administered by inhalation. For this purpose, methods are being developed to help analyzing the pharmacologically relevant unbound concentrations in brain and other tissues. Our work also emphasizes the importance of bridging the expertise within academia and pharmaceutical industry in order to seek excellence in method development for better therapeutics treating CNS and lung disorders.

### Members of the group during 2015

Jessica Dunhall, Laboratory Assistant  
 Sofia Gustafsson, MSc in Biomedicine, PhD student  
 Markus Fridén, Adjunct Lecturer  
 Margareta Hammarlund-Udenaes, PhD, Professor  
 Yang Hu, MSc in Pharmacy, PhD student  
 Britt Jansson, Laboratory Engineer  
 Annika Lindqvist, MSc in Pharmacy, PhD student  
 Irena Loryan, MD, PhD, Researcher  
 Erik Melander, MSc in Pharmacy, PhD student  
 Nebojsa Mihajlica, MSc in Pharmacy, PhD student  
 Johanna Sulku, MSc in Pharmacy, PhD student

### Publications 2013-2015

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### Reviews, books and book chapters 2013-2015

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8. de Lange EC and Hammarlund-Udenaes M. Translational aspects of blood-brain barrier transport and central nervous system effects of drugs: from discovery to patients. *Clin Pharmacol Ther* 97:380-394 (2015).

### **Agencies that supported the work/Funding 2015**

Janssen Pharmaceuticals; Swedish Research Council; AstraZeneca

### **Other commitments/assignments of staff members 2015**

Margareta Hammarlund-Udenaes: Associate Editor of the Journal *Pharmaceutical Research*, Editorial Advisory Board member of the *Journal of Pharmaceutical Sciences* and of *Fluids and Barriers of the CNS*, Member of the Executive Committee of the European Federation of Pharmaceutical Sciences with responsibility regarding Education and Training. Vice Chair of the Research Education Committee of the area of Medicine and Pharmacy.

### **Projects**

*(I) Concepts and method development regarding drug delivery to the brain (Irena Loryan and Margareta Hammarlund-Udenaes)*

Drug transport across the BBB and cellular barriers with subcellular distribution in the brain parenchyma are key processes of interest. Moreover, brain pharmacokinetic processes are also investigated at the level of its regions. In order to enhance the mechanistic understanding of brain target-site pharmacokinetics, several advanced methods are being developed by the group. Neuropharmacokinetic parameters such as unbound brain-to-plasma concentration ratio ( $K_p$ , uu, brain), unbound volume of distribution in brain ( $V_u$ , brain), and permeability clearance into the brain (CL<sub>in</sub>), are descriptors of BBB function and intracerebral distribution established by the group.  $K_p$ , uu, cell is another unique parameter characterizing the extent of cellular barrier transport. These parameters are identified by means of systematic PKPD analyses. For quantitative evaluation of BBB transport we are using cerebral microdialysis, recognized as the “gold standard” for measurement of unbound drug concentration in the brain. An alternative strategy is based on combination of high-throughput in vitro techniques, such as brain homogenate equilibrium dialysis and brain slices.

*(II) Translational aspects of brain drug distribution in health and disease (Sofia Gustafsson, Irena Loryan and Margareta Hammarlund-Udenaes in collaboration with Stina Syvänen and Martin Ingelsson, Dept Public Health and Caring Sci.)*

The interpretation and translation of neuropharmacokinetic data might be even more challenging during disease conditions, where a disrupted integrity and function of the BBB is apparent.

Recent findings even point towards dysfunctional BBB as being the cause of neurodegenerative disease etiology and progression. As a result, dysfunction in BBB processes might lead to altered brain pharmacokinetics of CNS drugs as well as peripherally acting drugs, which normally have a very low brain penetrance, resulting in unpredicted CNS effect or side effect profiles. Moreover, CNS disorder pathology usually affects certain areas of the brain which might result in regional differences of brain drug distribution and binding.

Hence, our research aims to investigate differences in drug distribution to and within the brain as well as drug binding in separate brain regions both in health and disease, preclinically and clinically. By combining and comparing in vitro and in vivo experiments with clinical studies the current project strives to increase the understanding of pharmacokinetics and disease implication on brain drug distribution. Results from in vitro techniques as well as microdialysis and non-invasive imaging techniques, such as positron emission tomography (PET), is integrated and used to address the current issues.

*(III) The role of pericytes for brain drug distribution (Nebojsa Mihajlica, Margareta Hammarlund-Udenaes in collaboration with Christer Betsholtz, Dept Immunol, Genet and Pathol).*

Endothelial cells of the BBB represent a key component of the neurovascular unit (NVU), which also includes other types of cells such as pericytes, astrocytes, vascular smooth muscle cells, microglia and neurons.

Although the anatomical relationship between pericytes and endothelial cells suggests their close interaction, the relative contribution of the pericytes is still not sufficiently elucidated. Interaction between cells within the NVU is essential for the normal functioning of the central nervous system, whilst impairments in their communication can result in the development of many pathological conditions.

The principal aim of this project is to improve our understanding of the mechanisms of the NVU regarding the drug distribution into brain parenchyma, with special emphasis on the contribution of pericytes. Hence, better understanding of pericyte role in BBB regulation and drug distribution into the CNS has a great potential for the improvement of current pharmacotherapy of neurodegenerative diseases and other pathological conditions associated with BBB impairment.

*(IV) Biomolecular drugs and nanocarriers (Annika Lindkvist, Erik Melander, Yang Hu and Margareta Hammarlund-Udenaes in collaboration with Drs Pieter Gaillard and Jaap Rip, to-BBB, and Ulf Göransson, Div Pharmacogenosy, UU).*

Peptides and proteins play a crucial role in the regulation of brain activity in health and disease conditions. They are therefore promising candidates in the development of new neurotherapeutics. Understanding the use of large molecules and their interaction with the barriers of the CNS is crucial in order to succeed in the clinic.

One promising group of peptides is the cyclic peptides originating from different plant families. These peptides exhibit exceptional stability due to a cyclic cysteine knot forming disulfide bonds. Their cyclic nature makes them less susceptible to degradation by proteases, causing a great interest from a drug development perspective.

Our research focuses on the pharmacokinetics of brain delivery of peptides and the possible quantitative benefits of using nanocarrier systems, including their effect on the pharmacodynamic outcome. In vivo preclinical studies are performed to estimate the modulation of penetration across the BBB using targeted liposomes. For authentication of the principles, physiologically based pharmacokinetics population modeling is applied.

*(V) Optimizing brain penetration of drug candidates (Irena Loryan and Margareta Hammarlund-Udenaes)*

The combinatorial mapping approach developed by our group can be used as a BBB screening toolbox for selection of candidate drugs in early drug discovery by pharmaceutical companies.

In addition, a present project is focused on identification of desirable physicochemical properties for CNS penetration using computational models, aiming to facilitate discovery and development of novel neurotherapeutics.

Integration of overall findings is directed towards the development of physiologically based mathematical models of brain drug disposition.

*(VI) Drug distribution in the lung (Erica Bäckström, Elin Boger (Marie Curie program IMPACT, University of Warwick), Markus Fridén (AstraZeneca), and Margareta Hammarlund-Udenaes)*

This is a new project started in 2013 in collaboration with AstraZeneca. The treatment of asthma and chronic obstructive pulmonary diseases (COPD) was revolutionized by the introduction of inhaled corticosteroids (ICS) and bronchodilators such as beta adrenergics and anti-muscarinics. For these drug classes topical delivery by inhalation has provided an efficient means of overcoming systemically mediated side-effects, that previously limited the dosing and therapeutic response.

Despite the historical success of inhalation medicines and the significant share (10 %) of the global drug market, there is very little known about the fundamental prerequisites for a drug molecule to be retained in and exert a localized effect in the lung when administered by inhalation.

Our research aims to study basic mechanisms of drug distribution in the lung which include non-specific tissue binding, lysosomal trapping and carrier-mediated membrane transport as well as the profound influence of blood perfusion. We are employing a lung slice methodology, equilibrium dialysis of lung tissue and in vivo methodologies to study the extent of drug distribution in the lung and the absorption half-life of inhaled drugs.

As an indicator of the unbound and pharmacologically active drug concentration we are collaborating with industrial and academic partners on measurement of target occupancy in the lung after inhalation using mass-spectrometry or positron emission tomography (PET). Along the same lines we are also developing physiologically based pharmacokinetic (PBPK) models to better explain and predict the possible advantage of the inhaled route of drug delivery.

*(VII) Clinical Pharmacy Research (Johanna Sulku, Elisabet Nielsen and Margareta Hammarlund-Udenaes in collaboration with Håkan Melhus, Dept Med Sci, Ulrika Gillespie, Uppsala Univ Hospital, and Hirsh Koyi, Gävle)*

We are interested in evaluating the impact of clinical pharmacist interventions in medical care.

The purpose of this research is to see whether, and in what way the contributions of clinical pharmacy services in the hospital ward changes patient treatment and status. This research area is new in Sweden and important for the development of clinical pharmacy in this country. A seminal paper was published in 2009 in Arch Intern Med, which received much attention. Here we showed that clinical pharmacist intervention reduced costs and decreased the number of readmissions to hospital. The research is further oriented towards appropriate use of medicines in chronic obstructive lung disease.

## Undergraduate Teaching 2015

Most of the teachers were involved in the continued work with revision of the Master of Science in Pharmacy program, including both renewal as well as improved training in oral and written communication, lab skills and professional training. A new support function started within the disciplinary domain of Medicine and Pharmacy consisting of three key teachers and a reference group. The main goals are to stimulate pedagogical development and dissemination of good ideas and to improve quality assurance in educational activities. The Faculty of Pharmacy is represented by Jörgen Bengtsson. Anne-Lie Svensson and Jörgen Bengtsson continued the faculty work with bachelor projects including half-time assessment, improved feedback and examination. The project has been evaluated and proved to result in higher quality of the projects. Two teachers, Jörgen Bengtsson and Emma Lundkvist, continued with their faculty assignments as coordinators of apotekare, receptarie and master programs, respectively.

The major part of the undergraduate teaching is within the two Pharmacy programs. During 2015 the extent of undergraduate teaching was 485 *hst* (full-time equivalents) representing 46% of the total number of *hst* within the Faculty of Pharmacy. In addition, the department is involved in teaching at the Master of Science in Chemical Engineering with specialization in drugs and the Biomedical program. Students attending internet-based courses during 2015 comprised 77 *hst*.

Pharmaceutical Biosciences comprises a number of courses, e.g. Drug development and Drug usage, Drug metabolism and safety, Gene technology, Infection Biology, Microbiology and immunology, Pharmaceutical Molecular Biology with Bioinformatics, Pharmaceutical Biochemistry, Pharmaceutical bioinformatics, Pharmacology, Pharmacokinetics, Pharmacotherapy, Physiology, Toxicology, Drug metabolism and Safety assessment. A mix of traditional teaching and problem-based learning with lectures, laboratory sessions, seminars, workshops and computer sessions characterizes the teaching. In addition, the teachers are involved in interdisciplinary training in laboratory practice, communication skills and professional development.

### *Teaching at the basic level*

The main teaching at the basic level is within the Bachelor of Science in Pharmacy program that comprises three years studies (180 hp) and the first years of the Master of Science in Pharmacy program, see below. Completed studies at the Bachelor program provide the necessary theoretical and practical competence that is required to apply for a pharmacist license as *receptarie*. The teachers also instruct in undergraduate projects (15 hp) at the basic level. These projects are individual and they are examined by an oral presentation and a written report. During 2015, the teachers within the department supervised 73% (29 students) of the total number of undergraduate projects within the Bachelor of Science in Pharmacy program.

Some basic level courses are open for other students than pharmacy students and attract both students at other programs at Uppsala University or other universities and also professionals.

### *Teaching at the advanced level*

The department gives courses at the Master of Science in Pharmacy program that comprises five years studies (300 hp). Completed studies at the programs provide the necessary theoretical and practical competence that is required to apply for a pharmacist license as *apotekare*. The teachers instruct in undergraduate projects (30 hp) at the

advanced level. These projects are individual and are examined by an oral presentation and a written report. The majority of the projects are laboratory-based and involve the student in ongoing research projects. During 2015, the teachers supervised 79% (55 students) of the total number of undergraduate projects within the Master of Science in Pharmacy program.

The teachers lead and teach a number of elective courses and single subject courses at the advanced level. These courses mirror research profiles within the department, such as Bioinformatics, Clinical pharmacy, Drug metabolism and safety, Drug addiction and Pharmacokinetics. The courses attract a large number of students, not only pharmacy students but also other students and professionals showing the proper prerequisites. During 2015, 75% (274 students) of the students within the Master of Science in Pharmacy program and 73% (70 students) of the students within the Bachelor of Science in Pharmacy program participated in elective courses given by the department.

The teachers also contribute to the teaching within several master programs within the Faculty of Pharmacy, Drug development, Drug management and safety and Clinical Pharmacy, and at master programs at the Faculty of Medicine, Infection biology and Forensic Science. The one-year post-graduate program in Clinical Pharmacy is the only one in the Nordic countries attracting students from all over Sweden and also from Norway.

Uppsala 2016-05-10

*Ingrid Nylander*

## Course List 2015

### List of courses on basic and advanced (second cycle) levels

Abuse and Addiction, 7,5 c  
 Acute Intoxications and Clinical Toxicology Second cycle, 7.5 c  
 Advanced Pharmacotherapy Second cycle, 7.5 c  
 Adverse Drug Reactions and Pharmacovigilance Second cycle, 7.5 c  
 Analytical Toxicology Second cycle, 30 c  
 Applied Pharmaceutical Bioinformatics Second cycle, 5 c  
 Applied Pharmaceutical Structural Bioinformatics Second cycle, 5 c  
 Applied Pharmacotherapy, Pharmacokinetics and Therapeutics Second cycle, 15 c  
 Biochemistry of Gene Regulation Second cycle, 7.5 c  
 Clinical Attachment and Service Development Second cycle, 18 c  
 Clinical Drug Trials with Applied Biostatistics Second cycle, 7.5 c  
 Clinical Pharmacokinetics and Pharmacodynamics Second cycle, 7.5 c  
 Clinical Pharmacy, 7.5 c  
 Degree Project in Drug Discovery and Development Second cycle, 30 c  
 Degree Project in Drug Management, 15 c  
 Degree Project in Drug Management Second cycle, 30 c  
 Degree Project in Pharmaceutical Biochemistry, 15 c  
 Degree Project in Pharmaceutical Biochemistry Second cycle, 30 c  
 Degree Project in Pharmaceutical Bioinformatics Second cycle, 30 c  
 Degree Project in Pharmaceutical Bioscience Second cycle, 20 c  
 Degree project in Pharmaceutical Pharmacology Second cycle, 30 c  
 Degree project in Pharmacokinetics, 15 c  
 Degree Project in Pharmacokinetics Scend cycle, 30 c  
 Degree project in Pharmacokinetics D Second cycle, 30 c  
 Degree Project in Pharmacology, 15 c  
 Degree Project in Pharmacotherapy, 15 c  
 Degree Project in Pharmacotherapy Second cycle, 30 c  
 Degree Project in Pharmacotherapy D Second cycle, 30 c  
 Degree Project in Toxicology, 15 c  
 Degree Project, Toxicology D Second cycle, 30 c  
 Drug Dependence Mechanisms, Prevention of Cannabis Abuse (Contract education) 7,5 c  
 Drug Development and Drug Usage, 7.5 c  
 Drug Management Second cycle, 7.5 c  
 Drugs and Dependence, Advanced Course Second cycle, 7.5 c  
 Drugs and the Elderly Second cycle, 7.5 c  
 Drugs during Pregnancy and Lactation, Second cycle, 7.5 c  
 Drug Safety and Pharmacovigilance, Second cycle, 7.5 c  
 Evidence Based Clinical Pharmaceutical Methods Second cycle, 12 c  
 Models for Biological Systems Second cycle, 7.5 c  
 Molecular Biology with Focus on Drug Therapy, 7.5c  
 Molecular Mechanisms for Enzymatic Activation Second cycle, 7.5 c  
 Molecular Pharmacology, 7.5 c  
 Neuropharmacology Second cycle, 7.5 c  
 Pharmaceutical Biochemistry, 9 c  
 Pharmaceutical Biochemistry and Cell Biology, 7.5 c  
 Pharmaceutical Bioinformatics Second cycle, 7.5 c  
 Pharmaceutical Bioinformatics (Contract Education), 7.5c  
 Pharmaceutical Molecular Biology with Bioinformatics, 7.5c  
 Pharmacokinetics, 3 c  
 Pharmacokinetics, 7.5 c  
 Pharmacokinetics and Statistics, 9 c  
 Pharmacology, 15 c  
 Pharmacology, 16.5 c  
 Pharmacology for engineering students, 7.5 c

Pharmacotherapy, 7.5 c  
Pharmacotherapy in Self-Treatment, 9 c  
Preclinical Safety Assessment and Pharmacovigilance, 7.5c  
Research Project in Clinical Pharmacy Second cycle, 15 c  
Toxicology B, 7.5 c  
Toxicology for Engineering Students, 7.5 c  
Toxicology, Advanced Course D Second cycle, 30 c  
Toxicology, Drug Metabolism and Safety Assessment, 4.5 c  
Toxicology, Drug Metabolism and Safety Assessment, 7.5 c  
Veterinary Pharmacology Second cycle, 7.5 c

## Research Education 2015

The Department has a high priority for research training. The aim is to prepare the PhD students to become independent researchers able to make significant contributions to academia and/or pharmaceutical industry, and to take on highly qualified professional tasks. The Department has two weekly seminar series for PhD students and young scientists – one series on pharmacokinetics/pharmacometrics and one series on basic biomedical research. Here, the PhD students can improve their oral presentation skills and their ability to discuss research, which is an important goal within the scientific training.

The 4-year PhD program consists of research work and a number of courses, with a total course requirement of 30 points (one semester full-time). Most PhD students take more courses than that. The PhD thesis is a doctoral thesis consisting of separately published articles with a comprehensive introductory summary. A PhD thesis typically contains 4 – 5 scientific papers, of which at least half are published in scientific journals at the time of the public thesis defence. Besides research activities, the PhD students participate at international conferences, of which one presentation at an international conference is obligatory during the program.

Many PhD students also perform important tasks in undergraduate teaching, comprising around 20 % of their time, making the time for a PhD exam be around 5 years in total. The teaching assignments give the students a broader knowledge base than their own research area. It also gives them leadership and communication skills that are of importance in further professional life. PhD students may also tutor master students in experimental research work. At the end of 2015 there were 48 PhD students registered at the department, and 7 PhD students who defended their theses during the year.

The Research Education Group at the Department actively structures the application procedure for PhD student positions, especially by contributing to the process of establishing new positions and selecting new PhD students. This has shown a decline in new positions during 2015. The group consists of two representatives from teachers and one PhD student. The group, through the Chair, also follows the training by requesting yearly follow up document from a meeting between each PhD student and supervisor, where issues regarding coursework, progression in the research project, possible change of the scope of research and thereby updates of the research plan, as well as how the communication between student and supervisor is functioning, are evaluated. The Chair and Erika Roman are also members of the Research Education Committee of the area of Medicine and Pharmacy, hereby connecting the Department to central decision making.

Uppsala 2016-04-29

*Margareta Hammarlund-Udenaes*

## Awards 2015

1. Per Andrén: Research Infrastructure Fellow grant from the Swedish Foundation for Strategic Research.
2. Margareta Hammarlund-Udenaes: Nagai International Woman Scientist Award from the Academy of Pharmaceutical Science and Technology in Japan.
3. Stina Lundberg: Bertil Göranssonstipendiet för unga alkoholforskare.
4. Fred Nyberg: ECAD Annual Award 2015 from European Cities Against Drugs.
5. Ingrid Nylander: CANs drogforskningspris 2015.