

Curriculum Vitae

Dr. Werner Dubitzky

Table of Contents

Table of Contents	1
Personal Information	2
Career and Profile Summary.....	2
Academic Education/Qualifications.....	3
Employment History	3
Research Interests and Scientific Leitmotiv	4
Acquired Research Funding.....	6
Activities as Evaluator of Research	6
Research Collaborations, Project and Management Experience	6
Editorial Boards.....	9
Editor	9
Edited Books and Volumes.....	9
Special Issues of Scientific Journals.....	10
Organization of Conferences and Workshops	10
PhD Supervision and Member of PhD Evaluation Committees.....	11
PhD Supervision	11
Member of PhD Evaluation Committees.....	11
Teaching.....	11
Publications	12
Peer-Reviewed Articles in Scientific Journals.....	12
Peer-Reviewed Articles in Conference Proceedings	14
Peer-Reviewed Articles in Edited Volumes.....	18
Other Publications	19
Appendix A: Research Interests in Data Science (Related to Biomedicine)	21
Statistical machine learning	21
Statistical learning as experimental science	22
Machine learning that matters.....	23
Machine learning for integration/analysis of complex biomedical data sets.....	25
Feature engineering and knowledge integration in machine learning.....	25
Intelligent e-biomedicine	25

Personal Information

Name **Werner Dubitzky**
Address: Stiefenhofen, Germany
E-Mail: werner@dubitzky.com



Google Scholar <https://scholar.google.com/citations?user=RZ-ncLQAAAAJ>

Career and Profile Summary

Within the restructuring framework of the education sector in the UK, I have availed of a voluntary redundancy offer and left the UK after more than 14 years as Professor of Bioinformatics at the University of Ulster. I am now looking for a new challenge in research, education, industry or government. Below I highlight some aspects of my professional qualifications and experience:

- PhD in computer science (artificial intelligence, machine learning, data mining) (UK, 1998) and MSc in electrical/telecommunication engineering (Germany, 1991).
- More than €8.5m research funding acquired from European (EC, ESF, COST Association) and national (UK, Germany, Japan) funding organizations.
- Author of 150 published research articles in international scientific journals and conference proceedings, editor of nine published books including the Springer Encyclopedia of Systems Biology, and editor of ten special issue in international scientific journals.
- 15 years' experience with strategic, organizational, financial and operational aspects of science management:
 - as coordinator and manager of large collaborative national and international research projects and networks as well as research grant proposals;
 - as formal evaluator and reviewer of large research grant proposals and funded research projects for European (EC, COST Association), national and international funding programs (in both biomed and ICT areas);
 - as Head of the formal Bioinformatics Research Group (20 team members) and as member of the Directorate of the Biomedical Science Research Institute (ca. 100 academic staff and ca. 200 PhD students) at the University of Ulster;
 - as adviser for national and international research funding organizations and governmental entities;
 - as manager of scientific communication and result exploitation activities in large European collaborative research projects.
- 25 years' experience as researcher and educator in ICT and bioinformatics areas with focus on data science (data and knowledge engineering, machine learning, algorithms)

and modeling and simulation of complex systems (systems biology, multiscale modeling and simulation).

- Member of editorial board and program committee of various high-caliber international scientific journals and conferences.
- Organization of large international conferences, workshops and symposia and chair of international program committees.
- Development/teaching of MSc course in computational systems biology.
- Experience in various programming languages including C/C++ (8 years), Java (8 years) and R (4 years).
- Experience with a large variety of software tools, including office and science applications and operating systems.
- Experience with spinning out an data science IT company in biotechnology sector.
- Experience as consultant in the area of data mining (in the biotechnology sector) and information and IT-security (specifically within the context of the international norms in the IEC/IOS 27000 series).
- Highly proficient in spoken and written English.

Academic Education/Qualifications

1998	PhD computer science (artificial intelligence and machine learning) at the University of Ulster, Jordanstown, UK
1991	MSc in electrical and telecommunications engineering at the University of Applied Sciences, Augsburg, Germany

Employment History

07/2016 – 09/2016	Senior consultant for IT security, ausecus GmbH, Augsburg, Germany
01/2002 – 04/2016	Chaired Professor for Bioinformatics at the University of Ulster, Coleraine, UK
03/2001 – 12/2001	IT consultant for databases and data mining for phase-it Intelligent Solutions AG, Heidelberg, Germany
01/2000 – 12/2001	Senior scientist and team leader (data mining, bioinformatics), German Cancer Research Center, Heidelberg, Germany
10/1999 – 12/1999	Senior scientist and team leader (data mining, knowledge managem), Research Institute for Knowledge Processing, Ulm, Germany
01/1999 – 09/1999	Lecturer in Informatics, University of Ulster, Jordanstown, UK
07/1993 – 12/1998	Research Fellow und Research Associate, University of Ulster, Jordanstown, UK
10/1991 – 07/1997	PhD candidate, University of Ulster, Jordanstown,UK
05/1997 – 09/1993	Software engineer for various companies in Germany (in total 18 months in the given period): Siemens-Nixdorf, Augsburg; Atlas Elektronik, Bremen; CIB Software, Munich; MEL Mikroelektronik,

Munich

07/1976 – 11/1979 Apprenticeship as industrial electronics technician, KUKA, Augsburg, Germany

Research Interests and Scientific Leitmotiv

Since I started my degree studies in electrical/telecommunications engineering in 1984, my career has been strongly influenced by my interest in natural science as well as computer science. Key science and information and communication technology (ICT) areas I have worked in since the start of my PhD in 1991 include the following:

- Object-oriented data and information modeling, management and programming
- Artificial intelligence, machine learning, statistics and data mining
- Modeling and simulation including multiscale modeling and simulation
- Computational science and e-science
- Distributed computing (grid/cloud computing)
- Computational biology, bioinformatics and systems biology

Reflecting the increasingly fuzzy boundaries between science and technology, the areas listed above could be viewed as the key elements in the emerging field of enhanced science or e-science. One of the key (and often overlooked) differences between e-science and other application areas where ICT is playing an increasingly important role (commerce, manufacturing, finance, government, agriculture, retail, and many others), is the requirement that ICT solutions in science need to facilitate the increase of scientific knowledge.

While e-science as an interdisciplinary framework (in its conventional form known as computational science or scientific computing) has been around for many years, there is still a bias towards compute-intensive modeling and simulation approaches (systems dynamics, dynamical and complex systems, control theory, and so on). The field of computational biology is a good example of this type of R&D. Recently, there has been a realization that data-intensive and intelligent technologies, such as machine learning and artificial intelligence, computational intelligence, computational creativity, could usefully complement the standard e-science framework. For instance, the field of bioinformatics and computational biology has a long tradition in incorporating intelligent techniques into the arsenal of tools, albeit usually not in the context of large-scale computing and very large data (this is changing, however). The data-intensive element in e-science is sometimes referred to as the »fourth paradigm« (complementing the first three science paradigms of experiment, theory and computational science). One could argue that modeling, analysis and simulation based on the methods and tools of intelligent technologies may develop into the »fifth paradigm« of the evolving scientific process. I refer to this emerging framework or scientific process as

intelligent e-science

which could be defined as follows:

1. Experiment (first paradigm)

2. Theory (second paradigm)
3. Computational science (third paradigm)
4. Data-intensive science (fourth paradigm) – nowadays called »data science«
5. Science enabled by intelligent technologies (fifth paradigm)

The intelligent e-science framework as outlined above could be viewed as a transdisciplinary scientific paradigm in which researchers work jointly using a shared conceptual framework and combined disciplinary-specific approaches to address complex R&D problems. Clearly, this paradigm will require considerable changes transcending the mind-set and culture of current science and education environments. The present scientific mind-set and culture is characterized by an interdisciplinary approach (where researchers work jointly but still from a disciplinary-specific basis) which has evolved from multidisciplinary science (researchers working in parallel or sequentially from disciplinary-specific base) in the past century. It is my view that intelligent e-science is likely to be at the heart of future transdisciplinary science.

Appendix A outlines some of my data science/machine learning research interests related to biomedicine in more detail.

Acquired Research Funding

Project Title (Acronym)	Role	Funder	From	To	£	€
Open Multiscale Systems Medicine (OpenMultiMed)	PI	COST	01.05.2016	30.04.2019		560,000
Multiscale Applications on European e-Infrastructures (MAPPER)	PI	EC-FP7	01.01.2010	30.09.2013		216,000
Robustness of Pathways to Invasion and Metastases in Breast Cancer	CI	Sasakawa	01.01.2009	31.07.2009	2,500	
Computational Systems Biology	PI	I.N.I.	13.12.2008	17.12.2008	368	
Bisociation Networks for Creative Information Discovery (BISON)	PI	EC-FP7	01.06.2008	31.05.2011		194,000
A Virtual Environment for Socially Aware knowledge Management (KnowledgeCraft)	PI	I.N.I.	30.01.2008	02.02.2008	630	
Computational Systems Biology	PI	I.N.I.	14.01.2008	28.02.2009	717	
SOLID -- Open, Service-Oriented Infrastructure for Language-Based Information Discovery	PI	I.N.I.	12.12.2007	14.12.2007	166	
Classification of Formalin-Fixed Paraffin-Embedded Tissue Data	PI	Almac	01.02.2007	30.09.2012	22,140	
SystemsBiologyGrid	PI	I.N.I.	27.11.2006	03.12.2006	1,662	
Mining of High-Throughput Data in Functional Genomics	CI	ESF	01.10.2006	30.09.2007	6,716	
Quasi-Opportunistic Supercomputing for Complex Systems in Grid Environments (QosCosGrid)	PI	EC-FP6	01.09.2006	31.05.2009		247,500
Bioinformatics Capability Funding	PI	D.E.L.	01.07.2006	31.07.2007	242,936	
Grid Services Based Environment to Enable Innovative Research (Chemomentum)	PI	EC-FP6	01.07.2006	31.03.2009		215,791
Advanced Methods and Technologies for Bioinformatics	CI	EC-ALFA	02.04.2006	01.12.2006	5,520	
Centre for Metabolomics	CI	D.E.L.	01.10.2005	31.03.2008	3,510,875	
Bioinformatics Capability Funding	PI	D.E.L.	31.08.2005	31.07.2006	234,000	
NUGO -- Impact of Food on Health -- Linking Genomics and Nutrition and Health Research	CI	I.N.I.	11.04.2005	30.04.2011	1,360	
A Regional Network for Post-Genomics and Systems Biology (SB(R)Net)	PI	EPSRC	01.10.2004	07.02.2008	58,182	
Data Mining Tools and Services for Grid Computing Environments (DataMiningGrid)	PI	EC-FP6	01.09.2004	30.09.2006		347,000
Systems Biology and Imaging: An Integrative Approach	CI	MRC/EPSRC	01.09.2004	31.08.2005	47,990	
Bioinformatics Capability Funding	PI	D.E.L.	01.08.2004	31.07.2007	502,000	
Bio 2004	PI	I.N.I.	02.06.2004	06.06.2004	5,995	
NUGO -- Impact of Food on Health -- Linking Genomics and Nutrition and Health Research	CI	EC-FP6	01.01.2004	30.04.2011	15,536	
Evaluation of the Prevalence of Vitamin D Deficiency ... and its Impact on Bone Health	CI	H.E.A.	01.10.2003	30.03.2006	72,816	
Medical Proteomics on the Grid	PI	I.N.I.	09.07.2003	17.17.2003	1,152	
Open Life Science GRID	PI	I.N.I.	20.11.2002	23.11.2002	2,235	
Information Integration of Life Science Data	PI	DWP	08.10.2002	30.10.2005	11,800	
Open Computing GRID for Molecular Science and Engineering (OpenMolGRID)	PI	EC-FP5	01.09.2002	28.02.2005	219,184	
ChemoGRID	PI	I.N.I.	02.02.2002	10.02.2002	280	
Miscellaneous	PI		01.01.2002	31.12.2011	1,916	
ESF COST Action 282: Knowledge and Exploration in Science and Technology (KnowEST)	PI	ESF	01.12.2001	30.06.2005		240,000
Bioinformatics System for Correlation of Clinical and Molecular-Genetic Data	CI	BMBF	2 years			450,000
					4,968,676	2,470,291

Key:

EC = European Commission
 Sasakawa = Great Britain Sasakawa Foundation
 D.E.L. = Northern Ireland Department of Education and Learning
 I.N.I. = Invest Northern Ireland
 EPSRC = Engineering and Physical Sciences Research Council, UK
 MRC = Medical Research Council, UK
 ESF = European Science Foundation
 BMBF = Ministry for Education and Research, Germany
 COST = COST Association

PI = Principal Investigator
 CI = Co-Investigator

Activities as Evaluator of Research

I have served as evaluator of research grant proposals and running research projects for the European Commission (FP4, FP5, FP6, FP7, H2020), the European Science Foundation (ESF), the COST Association, the Human Science Frontier Program (HSFP), and national research funding programs in various countries (Germany, Spain, Norway, Czech Republic, Republic of Ireland, UK, Cyprus, Poland, Netherlands, and Canada).

Research Collaborations, Project and Management Experience

Leading role in the initiation and execution of national and international research collaborations and projects:

- Initiator and Action-Chair: COST Action 15120 Open Multiscale Systems Medicine. Research network, partner organizations from over 20 countries.

- Initiator and member of the management committee: Multiscale Applications on European e-Infrastructures. European collaborative project (FP7, e-Infrastructures), eight partner organizations from six countries.
- Initiator und Lead-Editor-in-Chief der Springer Encyclopedia of Systems Biology. In total several hundred researchers globally.
- Initiator and member of the management committee: Bisociation Networks for Creative Information Discovery. European collaborative project (FP7, FET), seven partner organizations from six countries.
- Initiator and formal project coordinator/manager: A Virtual Environment for Socially Aware Knowledge Management (KnowledgeCraft). IP research proposal under FP7 with partner organizations from 18 countries¹.
- Initiator and formal project coordinator/manager: Quasi-Opportunistic Supercomputing for Complex Systems in Grid Environments. Europäisches Verbundprojekt (FP6), zehn Partnerorganisationen aus neun Ländern (einschließlich Israel und Australien).
- Initiator and member of the management committee: Grid Services Based Environment to Enable Innovative Research. European collaborative project (FP6), nine partner organizations from seven countries.
- Initiator and formal project coordinator/manager: A Regional Network for Post-Genomics and Systems Biology (EPSRC, UK). Research network with partner organizations from five countries.
- Initiator and formal project coordinator/manager: Data Mining Tools and Services for Grid Computing Environments. European collaborative project (FP6), five partner organizations from five countries including Israel.
- Initiator and member of the management committee: Open Computing GRID for Molecular Science and Engineering. European collaborative project (FP5), seven partner organizations from five countries.
- Initiator und Action-Co-Chair: COST Action 282: Knowledge Exploration in Science and Technology. Researcher network, partner organizations from more than ten countries.
- As editor of various edited volumes and special issues of scientific journals; member of editorial boards; organizer of conferences and workshops; and member of international program committees of international conferences and workshops I have cooperated with large number of organizations and scientists worldwide. (More details are found below)

In addition to the roles and activities listed above, I also held leading roles in the following:

- Head of the Bioinformatics Research Group (ca. 20 members), University of Ulster, Coleraine, UK
- Member of the Directorate of the Biomedical Sciences Research Institute (ca. 100 academics/researchers and 200 PhD students), University of Ulster, Coleraine, UK

¹ The proposal received 13 points in the remote evaluation and was ranked second among all large proposals in this call after the remote evaluation. After the hearing in Luxembourg the proposal was downgraded to 12 points for cost reasons.

- Leader of the data mining team at the German Cancer Research Center, Heidelberg, Germany
- Leader of the data mining and machine learning team at the Research Institute for Knowledge Processing, Ulm, Germany

I served as a representative of Northern Ireland (UK) at various meetings and events, including meetings with research funders from the European Commission and the National Science Foundation (NSF), USA.

As part of my research collaborations I spent significant periods of time at the following organizations :

- University of Amsterdam, Amsterdam, Netherland
- Jožef Stefan Institute, Ljubljana, Slovenia
- Universität Coimbra, Coimbra, Portugal
- Institute for Advanced Study, Budapest, Hungary
- Northwestern University, Chicago, USA
- University of Queensland, Brisbane, Australia
- Universidad de Talca, Talca, Chile
- Tokyo Institute of Technology, Tokyo, Japan
- Max-Planck-Institut für Astrophysik, Garching, Germany (in total ca. 3 months)
- Universität Konstanz, Konstanz, Germany (guest professorship 6 months)
- Hochschule Weihenstephan, Freising, Germany
- Hochschule Wildau, Wildau, Germany (ca. 2 months)
- Fraunhofer-Institut für Intelligente Analyse- und Informationssysteme IAIS, St. Augustin, Germany

Editorial Boards

2016 – 2016	<i>Advances in Bioinformatics</i> , Hindawi Publishing Corporation
2013 – 2016	<i>J. of Complex Systems</i> , Hindawi Publishing Corporation
2011 – present	<i>Network Modeling & Analysis in Health Inf. and Bioinformatics</i> , Springer
2009 – present	<i>Int'l J. of Next-Generation Computing</i> , Perpetual Innovation Media
2008 – 2016	<i>BMC Systems Biology</i> , BioMed Central
2006 – present	<i>Int'l J. of Comp. Intelligence in Bioinf. & Systems Biology</i> , Res. India
2004 – present	<i>OMICS A J. of Integrative Biology</i> , Mary Ann Liebert, Inc.
2003 – present	<i>Briefings in Bioinformatics</i> , Oxford University Press
2001 – present	<i>Online J. of Bioinformatics</i>
2008 – present	<i>The Open Biotechnology J.</i> , Bentham Open
2009 – 2012	<i>J. of Computational Science</i> , Elsevier
2005 – 2010	<i>Int'l J. of Data Mining and Bioinformatics (IJDMB)</i> , InderScience
2005 – 2011	<i>BMC Bioinformatics</i> , BioMed Central

Editor

Edited Books and Volumes

- Dubitzky W., Wolkenhauer O., Cho K-H., Yokota H. (editors-in-chief) (2013), *Encyclopedia of Systems Biology*, Springer
- Dubitzky W., Kurowski K., Schott B. (editors) (2012), *Large-Scale Computing Techniques for Complex System Simulations*, John Wiley and Sons, Inc.
- Dubitzky W., Southgate J., Fuß H. (editors) (2011), *Understanding the Dynamics of Biological Systems: Lessons Learned from Integrative Systems Biology*, Springer
- Dubitzky W. (editor) (2008), *Data Mining Techniques in Grid Computing Environments*, Wiley-Blackwell, ISBN: 978 0 470 51258 6.
- Dubitzky W., Schuster A., Sloot P., Schroeder M., Romberg M. (editors) (2007), *Proc. of Int'l Workshop Distributed, High-Performance and Grid Computing in Computational Biology (GCCB 2006)*, Eilat, January, 2007, *Lecture Notes in Bioinformatics LNBI 4360*, ISSN 0302-9743, Springer Berlin Heidelberg New York.
- Dubitzky W., Granzow M., Berrar D. (editors) (2007), *Fundamentals of Data Mining in Genomics and Proteomics*, Springer Berlin Heidelberg New York, ISBN: 978-0-387-47508-0.
- Bremer E.G., Hakenberg J., (Sam) Han E-H., Berrar D., Dubitzky W. (editors) (2006), *Proc. of Int'l Workshop on Knowledge Discovery in Life Science Literature (KDLL 2006)*, Singapore, April, 2006, *Lecture Notes in Bioinformatics LNBI 3886*, ISSN 0302-9743, Springer Berlin Heidelberg New York.
- Dubitzky W., Azuaje F. (editors) (2004), *Artificial Intelligence Methods and Tools for Systems Biology*, Kluwer Academic Publishers, Kluwer Academic Publishers,

Boston/Dordrecht/London. ISBN 1-4020-2859-8 (hardcover), 1-4020-2959-4 (paperback).

- López J.A., Benfenati E., Dubitzky W. (editors) (2004), Proc of Int'l Symposium Knowledge Exploration in Life Science Informatics 2004, Milan, Italy, 25-26 November 2004, Lecture Notes in Computer Science 3303, ISBN 3-540-23927-8 Springer Berlin Heidelberg New York.

Special Issues of Scientific Journals

- Dubitzky W. (2016). *Briefings in Bioinformatics*, 17(3). Special Issue: Computational Systems Biomedicine
- Dubitzky W. & Wang C. (2014). *Computation*. Special Issue: Multiscale Modeling and Simulation
- Gao J. & Dubitzky W. (2013), *IEEE J. of Biomedical and Health Informatics*
- Dubitzky W. (2010), *Briefings in Bioinformatics*, 10(4). Special Issue: Challenges in Bioinformatics and Computational Biology
- Dubitzky W. & Stankovski V. (2007), *Future Generation Computer Systems*, 23(1).
- Dubitzky W. (2006), *Briefings in Bioinformatics*, 7(4). Special Issue: Understanding the Computational Methodologies of Systems Biology
- Huang C-H., Lanza V., Rajasekaran S. & Dubitzky W. (2005), *J. of Clinical Monitoring and Computing*, 19(4-5)
- Lopez J.A., Azuaje F., Prank K. & Dubitzky W. (2004), *IEEE Transactions on Nano-Bioscience*, 3(3)
- Dubitzky W. (2004), *OMICS: A Journal of Integrative Biology*, 8(2)
- Dubitzky W. & Azuaje F.J. (2003), *Artificial Intelligence Review*, 20(1-2)

Organization of Conferences and Workshops

- Program Co-Chair: IEEE Int'l Conference on Bioinformatics & Biomedicine, Belfast, UK, 2-5 November 2014
- Program Co-Chair: Biomedical and Bioinformatics Challenges for Computer Science (BBC 2013) at Int'l Conference on Computational Science 2013 (ICCS 2013), Barcelona, Spain, 5-7 June 2013
- Program Co-Chair: IEEE Int'l Conference on Bioinformatics & Biomedicine, Philadelphia, USA, 4-7 October 2012
- Program Co-Chair: ESF Int'l Workshop on Mining of High-Throughput Data in Functional Genomics, University of Ulster, Coleraine, UK, May 8-9, 2007
- Program Co-Chair: Int'l Workshop on Distributed, High-Performance and Grid Computing in Computational Biology (GCCB 2006), Eilat, Israel, 2007
- Program Co-Chair: Int'l Workshop on Knowledge Discovery in Life Science Literature (KDLL 2006), Singapore, 2006

- Program Co-Chair: Int'l Program Committee of Int'l Symposium on Knowledge Exploration in Life Science Informatics (KELSI 2004), Milano, Italy, 2004
- Program Co-Chair: Int'l Conference on Artificial Intelligence 2003 (IC-AI'03), Las Vegas, Nevada, USA, 2003
- European Simulation Multi Conference, Track 8: Environment, Biology, Ecology, Sociology and Medicine, Darmstadt, 2002

Since 1999 I served as member of international program committees of over 100 international conferences and workshops.

PhD Supervision and Member of PhD Evaluation Committees

PhD Supervision

- Multiscale Modeling and Simulation in Systems Biology (PI)
- Computational Biology Study into of Genetic Causes of LQT Syndrome (PI)
- A Data Mining Approach to Protein Unfolding Simulation Data (Adviser)
- Data Exploration and Knowledge Extraction: Their Application to the Study of Endocrine-Disrupting Chemicals (Adviser)
- Analysis and Modelling of Interactions in Kinase/Phosphatase Systems (PI)
- Graph-Based Modeling and Reverse-Engineering of Biochemical Networks (PI)
- Automated Analysis of Biomedical Literature and its Application to Microarray Data Analysis and Interpretation (PI)
- Machine Learning Methods for Analyzing DNA Microarray Data (PI)
- Knowledge Discovery & Automated Decision-Making with Unstructured & Structured Data (PI)

Member of PhD Evaluation Committees

- UK (University of Ulster, Coleraine)
- Germany (Universität Konstanz)
- Switzerland (University of Geneva)
- Slovenia (Jožef Stefan Institute, Ljubljana)
- Spain (Universitat Pompeu Fabra, Barcelona)
- Portugal (University of Coimbra)

Teaching

I was the course director and sole developer of the PgCert/PgDip/MSc in Systems Biology course, which consists of eight taught modules plus a 60-credit research project. I have also experience in teaching C++ programming.

I designed and developed the full course which consists of eight taught modules and a research project. The course is designed around the various principles of systems, mathematical and computational thinking and the pedagogical concept of active learning. The course relies heavily on R as an analytical tool (analysis, visualization, modelling and simulation) and pedagogical tool facilitating active learning.

I have also supervised various local and international student dissertations and final year projects and as student adviser for local and international (Erasmus) students.

Publications

Peer-Reviewed Articles in Scientific Journals

Mizeranschi, A., Swain, M.T., Scona, R., Fazilleau, Q., Bosak, B., Piontek, T., Kopta, P., Thompson, P., Dubitzky, W. (2016). MultiGrain/MAPPER: A distributed multiscale computing approach to modeling and simulating gene regulation networks. *Future Generation Computing Systems*, 63: 1-14, Elsevier.

Mizeranschi, A., Zheng, H., Thompson, P., Dubitzky, W. (2016). A Two-Stage Inference Algorithm for Gene Regulation Network Models. *International Journal of Computational Biology and Drug Design*, 9:1/2, 6-24.

Mizeranschi, A., Zheng, H., Thompson, P., Dubitzky, W. (2015). Evaluating a common semi-mechanistic mathematical model of gene-regulatory networks. *BMC Systems Biology* 2015, 9(Suppl 5):S2.

Borgdorff J., Belgacem M.B., Bona-Casas C., Fazendeiro L., Groen D., Hoenen O., Mizeranschi A., Suter J.L., Coster D., Coveney P.V., Dubitzky W., Hoekstra A.G., Strand P., Chopard B. (2014), Performance of distributed multiscale simulations, *Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences*, 372(2021).

Wang C., Beyerlein P., Pospisil H., Krause A., Nugent C. & Dubitzky W. (2012), An efficient method for modeling kinetic behavior of channel proteins in cardiomyocytes. *IEEE/ACM Transactions on Computational Biology and Bioinformatics*, 9(1) (January 2012), 40-51. doi: 10.1109/TCBB.2011.84.

Dubitzky W. (2011), Toward a system-centric global knowledge management approach to discovering (organizing and sharing) scientific knowledge from large-scale data, *OMICS: A Journal of Integrative Biology*, 15(4): 243-246.

Hill T.R., Cotter A.A., Mitchell S., Boreham C.A., Dubitzky W., Murray L., Strain J.J., Flynn A., Robson P.J., Wallace J.M., Kiely M. & Cashman K.D. (2010), Vitamin D status and parathyroid hormone relationship in adolescents and its association with bone health parameters: analysis of the Northern Ireland Young Heart's Project, *Osteoporosis International*, 21(4): 695-700.

Swain M.T., Mandel, J.J. & Dubitzky, W. (2010), Comparative study of three commonly used continuous deterministic methods for modeling gene regulation networks. *BMC Bioinformatics* 11:459.

Swain M., Silva C.G., Loureiro-Ferreira N., Ostroptsyky V., Brito J., Riche O., Stahl F., Dubitzky W. & Brito R.M.M. (2010) P-found: Grid-enabling distributed repositories of protein folding and unfolding simulations for data mining. *Future Generation Computer Systems*, 26(3): 424-433

Stankovski, V., Swain, M., Kravtsov, V., Niessen, T., Wegener, D., Röhm, M., Trnkoczy, J., May M., Franke, J., Schuster, A. & Dubitzky, W. (2008), Digging deep into the data mine with DataMiningGrid, *IEEE Internet Computing*, 12(6), 69-76.

Cashman K.D., Hill T.R., Cotter A.A., Boreham C.A., Dubitzky W., Murray L., Strain J., Flynn A., Robson P.J., Wallace J.M. & Kiely M. (2008), Low vitamin D status adversely affects bone health parameters in adolescents, *Am J Clin Nutr.*, 87(4), 1039-1044.

Hill T.R., Cotter A.A., Mitchell S. & Boreham C.A., Dubitzky W., Murray L., Strain J.J., Flynn A., Robson P.J., Wallace J.M., Kiely M. & Cashman K.D. (2008), Vitamin D status and its determinants in adolescents from the Northern Ireland Young Hearts 2000 cohort, *Br J Nutr.*, 99(5), 1061-1067.

Stankovski V., Swain M., Kravtsov V., Niessen T., Wegener D., Kindermann J. & Dubitzky, W. (2008), Grid-enabling data mining applications with DataMiningGrid: An architectural perspective, *Future Generation Computer Systems*, 24(4), 259–279.

Fuß H., Dubitzky W., Downes C.S. & Kurth M.J. (2008), Src family kinases and receptors: analysis of three activation mechanisms by dynamic systems modeling, *Biophysical Journal*, 94(6), 1995-2006. Preprint, 30 Nov 2007: doi:10.1529/biophysj.107.115022. Open Access.

Fuß H., Dubitzky W., Downes C.S. & Kurth M.J. (2007), Deactivation of Src family kinases: Hypothesis testing using a Monte Carlo sensitivity analysis of systems-level properties, *Journal of Computational Biology*, 14(9), 1185–1200. doi: 10.1089/cmb.2007.0095

Mandel J.J., Fuss H., Palfreyman N.M. & Dubitzky W. (2007), Modeling biochemical transformation processes and information processing with Narrator, *BMC Bioinformatics*, 8:103. doi:10.1186/1471-2105-8-103.

Mandel J.J., Palfreyman N.M. & Dubitzky W. (2007), Modelling codependence in biological systems, *IET Systems Biology*, 1(1), 18-32. doi:10.1049/iet-syb:20060002

Gilbert D., Fuß H., Gu X., Orton R., Robinson S., Vyshemirsky V., Kurth M.J., Downes C.S. & Dubitzky W. (2006), Computational methodologies for modelling, analysis and simulation of signalling networks, *Briefings in Bioinformatics*, 7(4), 339-353. doi:10.1093/bib/bbl043.

Natarajan, J., Berrar, D., Dubitzky, W., Hack, C., Zhang, Y., DeSesa, C., Van Brocklyn, J.R. & Bremer, E.G. (2006), Text mining of full-text journal articles combined with gene expression analysis reveals a relationship between sphingosine-1-phosphate and invasivity of a glioblastoma cell line, *BMC Bioinformatics*, 7:373. doi:10.1186/1471-2105-7-373.

Fuß, H., Dubitzky, W., Downes, C.S. & Kurth, M.J. (2006), Bistable switching and excitable behaviour in the activation of Src at mitosis, *Bioinformatics*, 22(14), e158-e165. doi:10.1093/bioinformatics/btl201.

Berrar D., Bradbury I. & Dubitzky W. (2006), Instance-based concept learning from multiclass DNA microarray data, *BMC Bioinformatics*, 7:73. doi:10.1186/1471-2105-7-73.

Berrar D., Bradbury I. & Dubitzky W. (2006), Avoiding model selection bias in small-sample genomic data sets, *Bioinformatics*, 22(10), 1245-1250. doi:10.1093/bioinformatics/btl066.

Swain M., Hunniford T., Mandel J., Palfreyman N. & Dubitzky W. (2005), Reverse-engineering gene-regulatory networks using evolutionary algorithms and grid computing, *Journal of Clinical Monitoring and Computing*, 19(4-5), 329-337.

Berrar D., Stahl F., Goncalves Silva C.S., Rodrigues J.R., Brito R.M.M. & Dubitzky W. (2005), Towards data warehousing and mining of protein unfolding simulation data, *Journal of Clinical Monitoring and Computing*, 19(4-5), 307-317.

Fuß H., Dubitzky W., Downes S. & Kurth M.J. (2005), Mathematical models of cell cycle regulation, *Briefings in Bioinformatics*, 6(2), 1–15. doi:10.1093/bib/6.2.163.

Berrar D., Bradbury I., Sturgeon B., Downes C.S. & Dubitzky W. (2005), Survival trees for analyzing clinical outcome in lung adenocarcinomas based on gene expression profiles: Identification of neogenin and diacylglycerol kinase A expression as critical factors, *Journal of Computational Biology*, 12(5), 534–544.

Natarajan J., Berrar D., Hack C. & Dubitzky W. (2005), Knowledge discovery in biology texts: Applications, evaluation strategies, and perspectives, *Critical Reviews in Biotechnology*, 25(1-2), 31-52.

Mandel J., Palfreyman N., Lopez J. & Dubitzky W. (2004), Representing bioinformatic causality, *Briefings in Bioinformatics*, 5(3), 270-283. doi:10.1093/bib/5.3.270.

Dubitzky W., McCourt D., Galushka M., Romberg M. & Schuller B. (2004), Grid-enabled data warehousing for molecular engineering, *Parallel Computing*, 30(9-10), 1019-1035.

Brito R.M.M., Dubitzky W. & Rodrigues J.R. (2004), Protein folding and unfolding simulations: A new challenge for data mining, in W. Dubitzky (guest editor), Data mining meets integrative biology – a symbiosis in the making, *OMICS: A Journal of Integrative Biology*, 8(2), 153-166.

Berrar D., Dubitzky W., Solinas-Toldo S., Bulashevskaya S., Granzow M., Conrad C., Kalla J., Lichter P. & Eils R. (2001), Design and implementation of a database system for comparative genomic hybridization analysis, *IEEE Engineering in Medicine and Biology*, 20(4), 75-83.

Azuaje F., Dubitzky W., Black N.D. & Adamson K. (2001), Case retrieval strategies for CBR: A categorized bibliography, *The Knowledge Engineering Review*, 15 (4), 371-379.

Dubitzky W., Bulashevskaya S., Granzow M., Solinas-Toldo S., Joos S., Lichter P. & Eils R. (2001), A data mining approach to detect and analyse comparative genomic hybridization patterns of cancer cells, *Annales de Génétique*, 44(1), 160.

Azuaje F., Dubitzky W., Black N.D. & Adamson K. (2000), Discovering relevance knowledge in data: A growing cell structures approach, *IEEE Transactions on Systems, Man and Cybernetics. Part B: Cybernetics*, 30(3), 448-460.

Azuaje F., Dubitzky W., Black N.D. & Adamson K. (1999), Improving clinical decision support through case-based fusion, *IEEE Transactions on Biomedical Engineering*, 46(10), 1181-1185.

Dubitzky W., Büchner A.G., Hughes J.G. & Bell D.A. (1999), Towards concept-oriented databases, *Data and Knowledge Engineering*, 30, 23-55.

Dubitzky W., D. Mariotti, M. Hyland, J. McLaughlin & P. Maguire (1999), Analyzing plasma emission spectra using neural networks, *ISCA Int'l Journal of Computers and Their Applications*, 6, 88-94.

Patterson D., Anand S.S., Dubitzky W. & Hughes J.G. (1999), Towards automated case knowledge discovery in the M² case-based reasoning systems, *Knowledge and Information Systems: An Int'l Journal*, 1, 61-82.

Azuaje F., Dubitzky W., Lopes P., Black N.D., Adamson K., Wu X. & White J.A. (1999), Predicting coronary disease based on short-term electrocardiogram patterns: A neural network approach, *Artificial Intelligence in Medicine*, 15, 275-297.

Azuaje F., Dubitzky W., Lopes P., Black N.D., Adamson K., Wu X. & White J. (1998), Making clinical data meaningful: Knowledge discovery in coronary disease risk assessment, *The Irish Journal of Medical Science*, 167(4), 274.

Dubitzky W., Schuster A., Hughes J.G. & Bell A.D. (1997), An advanced case-knowledge architecture based on fuzzy objects, *Applied Intelligence: The Int'l Journal of Artificial Intelligence, Neural Networks, and Complex Problem-Solving Technologies*, 7, 187-204.

Dubitzky W., Hughes J.G. & Bell A.D. (1996), A generic, object-oriented case-knowledge representation scheme, and its integration into a wider information management scenario, *Expert Systems: The Int'l Journal of Knowledge Engineering and Neural Networks*, 13(3), 219-233, Blackwell Publishers, UK.

Dubitzky W., Carville F. & Hughes J.G. (1993), Case-level Knowledge modelling in CBR, *Irish Journal of Psychology*, 14(3), 478-479.

Peer-Reviewed Articles in Conference Proceedings

Mizeranschi, A., Zheng, H., Thompson, P., Dubitzky, W. (2014), A multi-model reverse-engineering algorithm for large gene regulation networks, *The IEEE Int'l Conference on Bioinformatics and Biomedicine 2014 (BIBM 2014)*, pp. 510-514, Belfast, UK, 2-5 November.

Mizeranschi, A., Kennedy, N., Zheng, H., Thompson, P., Dubitzky, W. (2014), Automated reverse-engineering of gene regulatory networks based on semi-mechanistic rate laws, *The 5th International Workshop on Integrative Data Analysis in Systems Biology*, Belfast, UK, 2-5 November 2014.

Mizeranschi A., Kennedy N., Thompson, P., Huiru Zheng, & Dubitzky W. (2014), The influence of network topology on reverse-engineering of gene-regulatory networks, *Procedia Computer Science, International Conference on Computational Science (ICCS 2014)*, 29(2014), 410-421.

Kennedy N., Mizeranschi A., Thompson, P., Zheng H. & Dubitzky W. (2013), Reverse-engineering of gene regulation models from multi-condition experiments, *IEEE Symposium Series on Computational Intelligence 2013 (SSCI 2013)*, Singapore, 16-19 April, pp. 112-119.

- Kennedy N., Thompson, P., Zheng H. & Dubitzky W. (2011), Multi-scale modelling of the bile acid and xenobiotic system, in Arabnia H.R. & Quoc-Nam T. (editors) Proc. of The 2011 Int'l Conference on Bioinformatics and Computational Biology (BIOCOMP'11), Volume II , 499-505.
- Kurowski, K., Back, W., Dubitzky, W., Gulyás, L., Kampis, G., Mamonski, M., Szemes, G. and Swain, M. (2009), Complex system simulations with QosCosGrid, Proc. of 9th Int'l Conference on Computational Science, Part I, G. Allen, J. Nabrzyski, E. Seidel, G.D. Albada, J. Dongarra and P.M. Sloot (eds.). Lecture Notes in Computer Science, vol. 5544. Springer-Verlag, Berlin, Heidelberg, 387-396.
- Kravtsov, V., Schuster, A., Carmeli, D., Kurowski, K. & Dubitzky, W. (2008), Grid-enabling complex system applications with QosCosGrid: An architectural perspective, in Proc. of Int'l Conference on Grid Computing and Applications (GCA'08), Las-Vegas, USA.
- Kravtsov, V., Carmeli, D., Dubitzky, W., Orda, A., Schuster, A., Silberstein, M. & Yoshpa, B. (2008), Quasi-opportunistic supercomputing in grid environments, in Proc. of Int'l Conference on Algorithms and Architectures, Cyprus, 233-244.
- Kravtsov, V., Swain, M., Dubin, U., Dubitzky, W. & Schuster A. (2008), A fast and efficient algorithm for topology-aware coallocation, in Proc. of Int'l Conference on Computational Science, Krakow, Poland, 274-283.
- Gulyas L., de Back W., Szemes G., Kurowski K., Dubitzky W. and Kampis G. (2008), Templates for distributed agent-based simulations, 20th European Modeling and Simulation Symposium, September 2008, Briatico, Italy.
- Swain, M., Mandel J.J. and Dubitzky, W. (2008), Comparing grid computing solutions for reverse-engineering gene regulatory networks, in Proc. of Int'l Conference on Computational Science (ICCS 2008), Poland, 106-115.
- Kravtsov., V., Swain, M., Schuster, A., Dubitzky, W. and Dubin, U. (2008), Grid computing solutions for distributed repositories of protein folding and unfolding simulations, in Proc. of Int'l Conference on Computational Science (ICCS 2008), Poland, 274-283.
- Swain, M., Ostropytskyy, V., Silva, C.G., Stahl, F., Riche, O., Brito, R.M.M. and Dubitzky, W. (2008), Grid computing solutions for distributed repositories of protein folding and unfolding simulations, Proc. of Int'l Conference on Computational Science (ICCS 2008), Krakow, Poland, 70-79.
- Silva C.G., Ostropytskyy V., Loureiro-Ferreira N., Berrar D., Swain M., Dubitzky W. & Brito R.M.M. (2006), P-found: The protein folding and unfolding simulation repository, in Proc. of the 2006 IEEE Symposium on Computational Intelligence in Bioinformatics and Computational Biology (CIBCB'06), 101-108.
- Berrar D. & Dubitzky W. (2006), Neural plasma, in M. Bramer (editor), Artificial Intelligence in Theory and Practice, Proc. of IFIP 19th World Computer Congress, TC 12: IFIP AI 2006 Stream, Series: IFIP Int'l Federation for Information Processing, Vol. 217, 159-168, XVI, Springer, Boston, ISBN: 0-387-34654-6.
- Bi Y. & Dubitzky W. (2006), An evidential approach in ensembles, in H. Haddad (editor), Proc. of the 2006 ACM Symposium on Applied Computing (SAC), , 1-6, Dijon, France, April 23-27, 2006. ACM 2006, ISBN 1-59593-108-2.
- Natarajan J., Haines C., Berglund B., DeSesa C., Hack C.J., Dubitzky W. & Bremer E.G. (2006), GetItFull – A tool for downloading and pre-processing full-text journal articles, in E.G., Bremer, J. Hakenberg, E.-H. S., Han, D. Berrar, W. Dubitzky (editors), Knowledge Discovery in Life Science Literature, Proc. of Int'l Workshop, KDLL 2006, Singapore, April 9, 2006, Lecture Notes in Bioinformatics, Vol. 3869, 139-145.
- Boreham C.A., Dubitzky W., Robson P.J., Wallace Julie M. & Bi Y. (2005), Three-dimension model for developing a nutrition, lifestyle and health database, Proc. of The 2005 Int'l Conference on Information and Knowledge Engineering, 151-157.
- Wang C., Krause A., Nugent C. & Dubitzky W. (2005), Focal activity in simulated LQT2 models at rapid ventricular pacing: Analysis of cardiac electrical activity using grid-based computation, Proc. of 6th Int'l Symposium on Biological and Medical Data Analysis (ISBMDA'05), 305-316, Portugal.

- Natarajan J., Mulay N., DeSesa C., Hack C.J., Dubitzky W. & Bremer E.G. (2005), A grid infrastructure for text mining of full text articles and creation of a knowledge base of gene relations, Proc. of 6th Int'l Symposium on Biological and Medical Data Analysis (ISBMDA'05), 101-108, Portugal.
- Hui W. & Dubitzky W. (2005), A flexible and robust similarity measure for analogy-based AI methods and analytical tasks, Proc. of the Int'l Joint Conference on Artificial Intelligence (IJCAI'05), 27-30, Edinburgh, Scotland.
- Stahl F., Berrar D., Goncalves Silva C.S., Rodrigues J.R., Brito R.M.M. & Dubitzky W. (2005), Grid warehousing of molecular dynamics protein unfolding data, in Proc. of 5th IEEE/ACM Int'l Symposium on Cluster Computing and the Grid, BioGrid 2005, Vol. 1, 496-503.
- Swain M., Hunniford T., Mandel J., Palfreyman N. & Dubitzky W. (2005), Modeling gene-regulatory networks using evolutionary algorithms and distributed computing, Proc. of the 5th IEEE/ACM Int'l Symposium on Cluster Computing and the Grid, BioGrid 2005, Vol. 1, 512- 519.
- Bremer E.G., Natarajan J., Zhang Y., DeSesa C., Hack C.J. & Dubitzky W. (2004), Text mining of full text articles and creation of a knowledge base for analysis of microarray data, in J.A. López, E. Benfenati & W. Dubitzky (editors), Proc. of Int'l Symposium Knowledge Exploration in Life Science Informatics 2004 (KELSI 2004), 84-95, Milan, Italy, 25-26 November 2004, Lecture Notes in Computer Science 3303, ISBN 3-540-23927-8 Springer Berlin Heidelberg New York, 2004.
- Bi Y., Bell D.A., Wang H., Guo G. & Dubitzky W. (2004), Classification decision combination for text categorization: An experimental study, Proc. of Database and Expert Systems Applications, 15th Int'l Conference (DEXA'04), 222-231, Lecture Notes in Computer Science 3180 Springer 2004, ISBN 3-540-22936-1.
- Stankovski V., May M., Franke J., Schuster A., McCourt D. & Dubitzky W. (2004), A service-centric perspective for data mining in complex problem solving environments, H.R. Arabnia & J. Ni (editors), Proc. of Int'l Conference on Parallel and Distributed Processing Techniques and Applications (PDPTA'04), Vol. 2, 780-787.
- Mazzatorta P., Benfenati E., Schuller B., Romberg M., McCourt D., Dubitzky W., Sild S., Karelson M., Papp A., Bágyi I. & Darvas F. (2004), OpenMolGRID: Molecular science and engineering in a grid context, H.R. Arabnia & J. Ni (editors), Proc. of Int'l Conference on Parallel and Distributed Processing Techniques and Applications (PDPTA'04), Vol II, 775-779.
- McCourt D., Lopez J., Benfenati E., Mazzatorta P., Romberg M., Schuller B. & Dubitzky W. (2003), Towards an intelligent data type for toxicity, Proc. of Int'l Conference on Artificial Intelligence, USA, 328-334.
- Ragg T., Granzow M., Menzel W. & Dubitzky W. (2003), Filtering nonlinear intensity dependencies in cDNA-microarray experiments, Proc. of Int'l Conference on Artificial Intelligence, 335-341.
- Berrar D., Downes C.S. & Dubitzky W. (2003), A probabilistic neural network for gene selection and classification of high-dimensional microarray data, Proc. of Int'l Conference on Artificial Intelligence, 342-349.
- Sturgeon B., McCourt D., Cowper J., Palmer F., McClean S. & Dubitzky W. (2003), Can the grid help to solve the data integration problems in molecular biology?, Proc. of the 3rd IEEE/ACM Int'l Symposium on Cluster Computing and the Grid, BioGrid 2003, 588-593.
- Berrar D., Downes C.S. & Dubitzky W. (2003), Multiclass cancer classification using gene expression profiling and probabilistic neural networks, Proc. of the Pacific Symposium on Biocomputing 8, 5-16.
- Berrar, D., Granzow M., Dubitzky W., Stilgenbauer S., Wilgenbus K., Döhner H., Lichter P. & Eils R. (2001), New insights in clinical impact of molecular genetic data by knowledge-driven data mining, Proc. 2nd Int'l Conference on Systems Biology, 275-281.
- Dubitzky W., Berrar D., Granzow M. & Eils R. (2001), Detecting broad-band and selective correlation patterns among gene expression and drug activity data, Proc. of Critical Assessment of Techniques for Microarray Data Analysis Techniques (CAMDA 2001), 17-22.
- Berrar D., Dubitzky W., Granzow M. & Eils R. (2001), Analysis of gene expression and drug activity data by knowledge-based association mining, Proc. of Critical Assessment of Techniques for Microarray Data Analysis Techniques (CAMDA 2001), 23-28.

- Dubitzky W., Krebs O. & Eils R. (2001), Minding, OLAPing, and mining biological data: Towards a data warehousing concept in biology, Proc. of Network Tools and Applications in Biology (NETTAB), CORBA and XML: Towards a Bioinformatics Integrated Network Environment, 78-82, Genoa, Italy.
- Schuster A., Dubitzky W., Azuaje F., Granzow M., Berrar D. & Eils R. (2000), Tumor identification by gene expression profiles: A comparison of five different clustering methods, Proc. of Critical Assessment of Techniques for Microarray Data Analysis (CAMDA 2000), 34-35.
- Dubitzky W., Granzow M., Berrar D., Bulashevskaya S., Conrad C., Gerlich D. & Eils R. (2000), A comparison of symbolic and subsymbolic machine learning approaches to molecular classification of cancer and gene identification, Proc. of Critical Assessment of Techniques for Microarray Data Analysis (CAMDA 2000), 12-13.
- Bulashevskaya S., Dubitzky W. & Eils R. (2000), Mining gene expression data using rough set theory, Proc. of Critical Assessment of Techniques for Microarray Data Analysis (CAMDA 2000), 4-5.
- Azuaje F., Dubitzky W., Black N.D. & Adamson K. (1999), Automated modelling of case retrieval structures using self-organising neural networks, Proc. of 1st Int'l Information Reuse and Integration Conference, 32-35.
- Lester N.G., Wilkie F.G., Dubitzky W. & Bustard D.W. (1999), A knowledge-guided retrieval framework for reusable software artefacts, Proc. of 17th Annual Association of Management / Int'l Association of Management Conf. on Computer Science, 206-211.
- Wang H., Dubitzky W., Düntsch I. & Bell A.D. (1999), A lattice machine approach to automated case base design: Marrying lazy and eager learning, Proc. of 16th Int'l Joint Conference on Artificial Intelligence, Sweden, Vol. 1, 254-259.
- Wu X.W., Dubitzky W., Azuaje F. & Black N.D. (1999), Discovering relevant knowledge for clustering through incremental growing cell structures, Proc. of 2nd Int'l. Conf. on Information Fusion, 46-52.
- Azuaje F., Dubitzky W., Black N.D. & Adamson K. (1999), Discovering and fusing relevant knowledge from databases based on an incremental unsupervised learning approach, Proc. 2nd Int'l. Conf. on Information Fusion, 31-38.
- Dubitzky W., Büchner A.G. & Azuaje F. (1999), Viewing knowledge management as a case-based reasoning application, Proc. of AAAI Workshop: Exploring Synergies of Knowledge Management and Case-Based Reasoning, AAAI Press, Technical Report WS-99-10, 23-27.
- Dubitzky W., Azuaje F., Lopes P. & Black N.D. (1999), McCullagh P., Song Y., On local and global feature weight discovery for case-based reasoning, Proc. of ISCA 14th Int'l Conference on Computers and their Applications, Mexico, 107-110.
- Azuaje F., Dubitzky W., Lopes P., Black N.D., Adamson K., Wu X., White J.A., Discovery of incomplete knowledge in electrocardiographic data, in Proc. of 3rd Int'l Conference of Neural Networks and Expert Systems in Medicine and Healthcare, Italy, E. Ifeachor, A. Sperduti, A. Starita (eds.), World Scientific, Singapore, 286-294, 1998.
- Dubitzky W., Mariotti D., Hyland M. & McLaughlin J. (1998), A neural network approach to plasma emission interpretation, Proc. of ISCA 11th Int'l Conference on Computers and their Applications in Industry and Engineering, 47-50.
- Dubitzky W., Lopes P., Hughes J.G. & Bell A.D. (1998), Representing incomplete knowledge in case-based reasoning, Proc. ISCA 11th Int'l Conference on Computers and their Applications in Industry and Engineering, USA, 133-136.
- Azuaje F., Dubitzky W., Lopes P., Black N.D., Adamson K., Wu X. & White J.A. (1998), Knowledge discovery in electrocardiographic data based on neural clustering algorithms, Proc. of 8th Mediterranean Conference on Medical and Biological Engineering and Computing, (electronic proceedings on CD), Cyprus, ISBN 9963-607-13-6.
- Patterson D., Dubitzky W., Anand S.S. & Hughes J.G. (1998), On the automation of case base development from large databases, Proc. of AAAI 1998 Workshop: Case-Based Reasoning Integrations, 126-130, USA.
- Azuaje F., Dubitzky W., Wu X., Lopes P., Black N.D., Adamson K. & White J.A. (1997), A neural network approach to coronary heart disease risk assessment based on short-term measurement of RR intervals, Proc. of Computers in Cardiology, Vol. 24, 53-56, Sweden.

Dubitzky W., Schuster A., Bell A.D. & Hughes J.G. (1997), How similar is VERY YOUNG to 43 years of age? On the representation and comparison of polymorphic properties, Proc. of 15th Int'l Joint Conference on Artificial Intelligence, 226-231, Japan.

Schuster A., Dubitzky W., Lopes P., Adamson K., Bell A.D., Hughes J.G. & White J.A. (1997), Aggregating features and matching cases on vague linguistic expressions, Proc. of 15th Int'l Joint Conference on Artificial Intelligence, 252-257, Japan.

Büchner A.G., Dubitzky W., Schuster A., Lopes P., Bell A.D., O'Doneghue P.G., Hughes J.G., Bell A.D., Adamson K, White J.A., Anderson J.M.C.C. & Mulvenna M.D. (1997), Corporate evidential decision making in performance prediction domains, Proc. of 13th Conference on Uncertainty in Artificial Intelligence, 38-45, USA.

Dubitzky W., Hughes J.G. & Bell A.D. (1997), Corporate multi-criteria decision making using evidential reasoning, Proc. of 26th Northeast Decision Sciences Institute Annual Meeting, 135-137, Annapolis, USA.

Schuster A., Dubitzky W., Adamson K., Bell A.D. & Hughes J.G. (1997), Processing similarity between a mix of crisply and fuzzily defined case properties, Proc. of 2nd Int'l ICSC Symposium on Fuzzy Logic and Applications, 247-253, Switzerland.

Dubitzky W., Hughes J.G. & Bell A.D. (1996), A multi-expert scheme for defining reliable case categories, Proc. of Int'l Conference on Knowledge Based Computer Systems, 91-102, India.

Dubitzky W., Hughes J.G. & Bell A.D. (1996), Case memory and the behaviouristic model of concepts, Proc. of Advances in Case-Based Reasoning, 3rd European Workshop, EWCBR-96, 120-134, Switzerland.

Dubitzky W., Lopes P., White J.A., Anderson J.M.C.C., Dempsey G.J., Hughes J.G. & Bell A.D. (1996), A holistic approach to coronary heart disease risk assessment using case-based reasoning, Proc. of 2nd Int'l Conference on Neural Networks and Expert Systems in Medicine and Healthcare, 97-103, UK.

Dubitzky W., Hughes J.G. & Bell A.D. (1996), Multiple opinion case-based reasoning and the theory of evidence, Proc. of 6th Int'l Conference on Information Processing and Management of Uncertainty in Knowledge-Based Systems, 447-452, Spain.

Dubitzky W., Hughes J.G. & Bell A.D. (1996), Knowledge management via case-knowledge and object-orientation, Proc. of 3rd World Congress on Expert Systems, 1150-1157, Korea.

Dubitzky W., Schuster A., Hughes J.G. & Bell A.D. (1995), Conceptual distance of numerically specified case features, Proc. of 2nd New Zealand Int'l Two-Stream Conference on Artificial Neural Networks and Expert Systems, 210-213, New Zealand.

Dubitzky W., Carville F. & Hughes J.G. (1993), Case-level knowledge modelling in CBR, Proc. of 6th Irish Conference on Artificial Intelligence and Cognitive Science, 217-227, UK.

Peer-Reviewed Articles in Edited Volumes

Mizeranschi, A., Groen, D., Borgdorff, J., Hoekstra, A.G., Chopard, B., Dubitzky, W. (2016), Chapter 17: Anatomy and physiology of multiscale modeling and simulation in systems medicine, in Ulf Schmits et al. (editors). Systems biology for medicine, Methods Mol Biology 2016; 1386, 375-404.

Berrar D. & Dubitzky W. (2013), Bootstrapping, in W. Dubitzky, O. Wolkenhauer, K.-H. Cho & H. Yokota (editors), Encyclopedia of systems biology, Springer, New York.

Berrar D. & Dubitzky W. (2013), Overfitting, in W. Dubitzky, O. Wolkenhauer, K.-H. Cho & H. Yokota (editors), Encyclopedia of systems biology, Springer, New York.

Berrar D. & Dubitzky W. (2013), Decision trees, in W. Dubitzky, O. Wolkenhauer, K.-H. Cho & H. Yokota (editors), Encyclopedia of systems biology, Springer, New York.

Kennedy N., Thompson P., Schmidt O., Zheng H. & Dubitzky W. (2013), Bile acid and xenobiotic system, in W. Dubitzky, O. Wolkenhauer, K.-H. Cho & H. Yokota (editors), Encyclopedia of systems biology, Springer, New York.

Dubitzky W., Kötter T., Schimdt O. & Berthold M.R. (2012), Towards creative information exploration based on Koestler's concept of bisociation, in M.R. Berthold (editor), *Bisociative knowledge discovery*, 11-32, Springer, LNCS 7250 2012.

Schmidt O., Kranjc J., Mozetič I., Thompson P. & Dubitzky W. (2012), Bisociative exploration of biological and financial literature with using clustering, in M.R. Berthold (editor), *Bisociative knowledge discovery*, 438-451, Springer, LNCS 7250 2012.

Swain M., Coti C., Mandel J. & Dubitzky W. (2011), A topology-aware evolutionary algorithm for reverse-engineering gene regulatory networks, in Dubitzky W., Kurowski K., Schott B. (eds.), *Large-scale computing techniques for complex system simulations*, 141-162, Wiley-VCH.

Sánchez A., Montes J., Dubitzky W., Valdés J.J., Pérez M.S. & de Miguel P. (2008), Data mining meets grid computing: Time to dance?, in W. Dubitzky (editor), *Data mining techniques in grid computing environments*, Wiley, 1-16.

Romberg M., Benfenati E. & Dubitzky W. (2007), Open computing grid for molecular sciences, in E-G. Talbi & A. Zomaya (editors), *Grids for bioinformatics and computational biology*, Wiley Book Series on Parallel and Distributed Computing. John Wiley & Sons, New York, 1-22. ISBN: 978-0-471-78409-8.

Berrar D., Sturgeon B., Bradbury I., Downes C.S. & Dubitzky W. (2004), Integration of microarray data for a comparative study of classifiers and identification of marker genes, in J.S. Shoemaker & S.M. Lin (editors), *Methods of microarray data analysis IV*, 147-162, Springer.

Dubitzky W., Granzow M., Downes C.S. & Berrar D. (2002), Introduction to Microarray Data Analysis, in D. Berrar, W. Dubitzky & M. Granzow (editors), *A Practical Approach to Microarray Data Analysis*, 1-46, Copyright 2002, Kluwer Academic Publishers, Boston/Dordrecht/London.

Dubitzky W., Granzow M. & Berrar D. (2001), Comparing symbolic and subsymbolic machine learning approaches to classification of cancer and gene identification, in S.M. Lin & K.F. Johnson (editors), *Methods of microarray data analysis: Papers from CAMDA'00*, 151-166, Kluwer Academic Publishers; ISBN: 0792375645.

Dubitzky W., Granzow M. & Berrar D. (2001), Data mining and machine learning methods for microarray analysis, in S.M. Lin & K.F. Johnson (editors), *Methods of microarray data analysis: Papers from CAMDA'00*, 5-22, Kluwer Academic Publishers; ISBN: 0792375645.

Dubitzky W. & Azuaje F. (2000), A genetic algorithm and a growing cell structures approach to learning case retrieval structures, in S.K. Pal, T.S. Dillon & D.S. Yeung (editors), *Soft Computing and Case-Based Reasoning*, 115-146, Springer.

Other Publications

Dubitzky W. (2010), Editorial: Challenges in bioinformatics and computational biology, *Briefings in Bioinformatics*, 343- 344, 10(4).

Dubitzky W., Stankovski V. (2007), Editorial: Data mining in grid and Web services computing environments: Challenges and applications, *Future Generation Computer Systems*, 23(1), 31-33. doi: 10.1016/j.future.2006.05.001.

Dubitzky W. (2006), Editorial: Understanding the computational methodologies of systems biology, *Briefings in Bioinformatics*, 7(4), 315-317. doi:10.1093/bib/bbl044.

Huang C-H., Lanza V., Rajasekaran S. & Dubitzky W. (2005), Editorial: *Journal of Clinical Monitoring and Computing*, 19(4-5), 259-262.

Lopez J.A., Azuaje F., Prank K. & Dubitzky W. (2004), Editorial: Molecular and sub-cellular systems biology, *IEEE Transactions on Nano-Bioscience*, 3(3), 141-143.

- Dubitzky W. (2004), Editorial: Data mining meets integrative biology – a symbiosis in the making, *OMICS: A Journal of Integrative Biology*, 8(2), 153-166.
- Dubitzky W. & Azuaje F.J. (2003), Editorial – AI in the life sciences, *Artificial Intelligence Review*, 20(1-2), 7-11.
- Dubitzky W. (2005), Review of Computational Modeling of Genetic and Biochemical Networks edited by Bower J.M. & Bolouri H. (2001), *BioMedical Engineering OnLine* 2005, 4:56, doi:10.1186/1475-925X-4-56
- Dubitzky W. (2003), Review of Advances in the Evolutionary Synthesis of Intelligent Agents edited by Patel M.J., Honavar V. & Balakrishnan K. (2001), *IEEE Transactions on Neural Networks*, 14(4), 970, July 2003.
- Berrar D.P., Sturgeon B., Bradbury I., Downes C.S. & Dubitzky W. (2003), Microarray data integration and machine learning techniques for lung cancer survival prediction, Critical Assessment of Microarray Data Analysis (CAMDA 2003), Oral and Poster Presenters Abstracts, 43-54, Durham, North Carolina, USA.
- Kochmann T.K., Dubitzky W., Flaig R.M. & Eils R. (2001), Towards a Hermeneutic knowledge management in science, Proc. of 222nd American Chemical Society National Meeting, CINF 36, Chicago, Illinois, USA. (Abstract)
- Dubitzky W., Kochmann T.K., Flaig R.M. & Eils R. (2001), Implementation of a global computing scenario in science, Proc. of 222nd American Chemical Society National Meeting, CINF 33, Chicago, Illinois, USA. (Abstract)
- Dubitzky W. (1998), Knowledge integration in case-based Reasoning: A concept-centered approach, PhD Thesis, University of Ulster, Northern Ireland, GB.
- Montazemi A.R., Bayless S.J., Dubitzky W., Gupta K.M. & Klahr P. (1996), Industrial application of case-based reasoning systems, Proc. of 25th Northeast Decision Sciences Institute Meeting, p587, USA.
- Granzow M., Berrar D., Dubitzky W., Schuster A., Azuaje F. & Eils R. (2001), Tumor classification by gene expression profiling: Comparison and validation of five clustering methods, in SIGBIO Newsletter Special Interest Group on Biomedical Computing of the ACM, ACM Press, 21(1), 16-22.
- Dubitzky W. (1992), An introduction to case-based reasoning, Expert. Newsletter of the Expert System Centre (University of Ulster, Jordanstown, GB), 5: 5-7, UK.

Appendix A: Research Interests in Data Science (Related to Biomedicine)

Below I outline various research areas related to data science/machine learning and biomedicine that I consider important. Essentially, these arise from my prior work which took me from an engineering perspective (computer science, artificial intelligence, machine learning, large-scale computing) to an experimental science perspective (bioinformatics and computational biology), and back (systems view, complexity, multiscale modeling, simulation and data science). These research interests center firmly on the integration and analysis of complex and heterogeneous biomedical data sets using statistical machine learning and modeling and simulation approaches.

In knowledge-rich domains like biomedicine, machine learning could offer a wealth of useful ways to approach problems that otherwise defy solution. Adopting machine learning in this area raises numerous new research challenges. First, partly because of the recent big data developments in the life sciences, we are witnessing a growing demand for novel statistical machine learning solutions. Second, there is a growing need to adopt a systematic experimental-science approach to machine learning research. Third, the bulk of machine learning research focuses on narrowly defined algorithmic and technical advances and data sets of limited scope, but the real-world impact of machine learning research is rarely assessed. Impact assessment is particularly important in biomedical applications. We need more research on machine learning that really matters, i.e. we need to develop and evaluate machine learning solutions in terms of their real-world impact. Fourth, emerging and future progress in biomedicine will increasingly rely on intelligent solutions from artificial intelligence, machine learning and other ICT areas – this is what I call *intelligent e-science*. We need to develop machine learning solutions in the context of the emerging scientific process in biomedical research. Fifth, systems medicine is an emerging interdisciplinary framework that aims to improve our understanding, prevention and treatment of complex diseases by integrating knowledge and data across multiple levels of biomedical organization. Such integration and analysis of complex, multiscale data requires a multiscale approach to systems medicine (multiscale machine learning, data science and modeling and simulation supported by multiscale computing). Sixth, feature engineering and representation learning are highly underestimated research areas in machine learning. Particularly, for scientific machine learning solutions in biomedicine these topics will become increasingly important in the near future.

Statistical machine learning

Machine learning was originally concerned with the automated acquisition, use and evolution of knowledge for intelligent systems that are able to perform complex tasks such as reasoning, design, diagnosis, problem solving, planning, and language understanding. This type of machine learning typically employs symbolic knowledge structures (e.g. production rules, decision trees and logical formulae) and data sets of small to moderate size. Statistical learning refers to a set of tools for modeling and understanding complex data sets. Statistical learning approaches commonly employ sub-symbolic knowledge structures and techniques (such as lasso and sparse regression, classification and regression trees, support vector machines, artificial neural networks) and are often based on large and very large data sets. Recent developments in the field of big data have led to a fast growth in the field of statistical learning. Nowadays, the distinction between statistical learning and machine learning is blurred and the term statistical machine learning is often used. Statistical machine learning could be viewed as a hybrid discipline borrowing techniques from statistical learning and machine learning. The adoption of statistical machine learning especially in knowledge-rich domains and in science poses many new machine learning research challenges. These challenges include, but are not limited to, high-dimensional data, sparsity, semi-supervised learning, the relation between computation and risk, and structured prediction. Some of these challenges are briefly highlighted below.

Most statistical machine learning theory is based on asymptotic approximations that allow the sample size n to grow large. When the number of variables p in the model is large, this theory can be problematic, however. The small- n -large- p problem is known as the curse of dimensionality. Unless certain strong assumptions hold, the sample size n needs to grow exponentially with p to achieve good model performance (statistical curse of dimensionality). One important challenge in statistical machine learning is to develop relevant theory and methods when p grows with n . Such a theory should yield useful insights for real data sets with moderate sample sizes but large number of variables. Associated with the statistical curse of dimensionality is the computational curse of dimensionality – the computational burden of algorithms typically grows exponentially with p . We need to develop novel statistical machine learning algorithms and tools that are able to scale gracefully with a growing number of variables.

In a typical statistical machine learning problem, getting raw data is relatively easy, but producing labeled examples is time-consuming and expensive, since the labeling may require expensive experiments, such as clinical trials or human experts. The challenge of semi-supervised learning is to somehow leverage large amounts of unlabeled data in order to improve upon a learning algorithm that uses only labeled data. While some work on this problem exists in the classical machine learning community, little attention has been given to this in the statistical learning field.

Statistical machine learning research is usually aimed at finding algorithms that minimize the expected loss or cost (or error) of the learner. But these algorithms usually ignore the computational costs of machine learning. We need to develop a new statistical machine learning theoretical framework that combines prediction error with computational complexity.

In statistical machine learning, a structured prediction problem can be thought of as a multi-class problem with a large number of class labels, typically exponential in the number of variables. Developing estimators and efficient algorithms for structured prediction problems, the structure of the problem must be taken into account. For example, in part-of-speech tagging in natural-language processing, the type of a word in a sentence depends strongly on the type of previous words in the sentence. More complex forms of structured prediction exist. As more complex problems are being tackled in knowledge-rich domains, more research is needed in the area of structured prediction problems.

Statistical learning as experimental science

Fundamental research in machine learning is inherently empirical, because the performance of machine learning algorithms is determined by how well their underlying assumptions match the structures and conditions in the real world. Hence, no amount of mathematical analysis can determine a priori whether a machine learning model will work well or not (this is known as the *no free lunch theorem* of machine learning). Hence, an experimental-science approach to machine learning is required. A machine learning researcher translates these assumptions into a set of learners (called the hypothesis space) and defines how the performance of a learner is to be evaluated. The researcher then implements these specifications using suitable computer software and hardware, and tests the performance on real-world data sets.

A great advantage of my excursion into life science is that I am very familiar with scientific process in experimental science. Machine learning researchers typically come from computer science, other engineering disciplines or mathematics. Therefore, they are normally not very familiar with the methodologies of experimental science. As a result, a great deal of machine learning research is conducted in a manner that does not fully embrace the principles and methodologies of experimental science. Typically, machine learning studies are designed around the following basic procedure: (1) Implement new algorithm. (2) Compare its performance (typically a kind of accuracy or error measure)

based on benchmark data sets to state-of-the-art algorithms. (3) Publish, if accuracy is higher (in most cases). This design of a machine learning study leaves a lot to be desired. First, it restricts assessment to a small selection of data and methods. Second, it limits assessment to only one criterion (typically a kind of accuracy score possibly associated with a p-value expressing statistical significance). Indeed, one of the main differences between machine learning and statistics is that machine learning places a strong emphasis on accuracy (of model predictions), whereas statistics typically emphasizes interpretability of models. Third, such a study design fails to take into account the idiosyncrasies (context) of many real-world problems. Arguably, the utility of this type of machine learning research methodology is limiting the progress of machine learning research.

Increasingly, because of the strong focus of machine learning on standard data sets (which are essentially viewed as matrices of numbers with their domain context ignored), performance measure typically report a kind of accuracy/error scalar associated with the p-value of a statistical significant test. Given the diversity of potentially interested people (particular researcher, the research community as a whole, end users of applications and, of course, referees for conference and journal papers), it is unreasonable to expect to capture all their concerns in a single scalar measure. With only the outcome of a significance test (usually: reject null hypothesis), we have no idea of the size of the actual difference between our measured values, however small the p-value. We need additional information. We can study the tables of results produced in a paper and make our own estimate of the difference, but this is not a statistically justified procedure. One way this problem could be addressed is by the use of confidence intervals. From these we can judge not statistical significance but also the size of the effect. Each researcher can then decide what effect is sufficiently large to make the approach interesting in the context of their work and problems. Another problem of requiring statistically significant results before a paper is published is that we do not see the whole picture. A survey of the literature would give an impression that there is stronger support for a particular algorithm than there actual is. Another issue is the strong reliance of machine learning research on standard benchmark data sets. The main advantage of commonly used machine learning benchmark data sets is our familiarity with them. However, this is also the major downside, as the very familiarity with these data sets potentially leads to over-fitting. Our knowledge encourages the writing of algorithms that are tuned to them. This is part of a larger concern about how well experimental results will generalize to other yet unseen problems.

Even when an experimental machine learning study has been carried out with great care, the conclusions that can be drawn from it are often very weak. Problematic components of the standard machine learning testing procedures include the performance measures used, the reliance on null hypothesis statistical testing, and the use of benchmark data sets. Hence, future machine learning research should adopt a much wider view in evaluating machine learning solutions. For example, from a *local* perspective, we would like to know how well our testing procedure predicts performance on future applications. From a *global* perspective, we would like to know how well our testing procedure encourages progress in our field. Present machine learning evaluation procedures are not as effective, from the local perspective, as people imagine. But it does not seem to be doing very well from the global perspective. Not only is it ineffective at filtering out dubious theories, the overly strong emphasis on testing discourages a broad, dynamic, and ultimately more fruitful dialogue within the machine learning community.

Machine learning that matters

A large portion of machine learning research is inspired by challenging real-world problems in medicine, education, science, engineering, environment, economy, society, and so on. And yet, we still see a plethora of published machine learning papers that evaluate new solutions on a handful of isolated benchmark data sets. While these data sets may have originated in the real world, the results from the evaluation of the solutions are rarely communicated back to the origin. Quantitative improvements in performance are rarely accompanied by an assessment of whether those gains

matter in the world outside of machine learning research. A large part of current machine learning research is too focused on benchmark data sets and abstract performance metrics, and lacks proper follow-through with the real-world user community.

A growing number of machine learning studies present the results of a new algorithm based on synthetic data and/or standard data sets, such as those available from the UCI machine learning repository. The main advantage of such an approach is better comparison with other algorithms. However, in practice comparisons fail, because there are no standards for reproducibility. Machine learning studies vary considerably in methodology (partitioning of data, performance metrics, parameter settings), implementations (tools, libraries, platforms, programming languages, hardware), and reporting (language, formal specification, tables, charts and plots, intermediate data). Furthermore, interpretation of the result in the context of the domain problem is practically never made. Does a certain increase of classification performance over a small set of state-of-the-art solutions matter in the domain from which the data set comes from? Which classes were predicted well and which not so well, and what does this mean in the context of the domain?

There are also problems with how we measure performance. Commonly used performance metrics include classification accuracy, various forms of errors, F-measure, lift, area under the ROC curve, etc. These metrics are highly abstract in that they do not contain problem-specific details. While this allows for results to be compared across solutions and domains, these metrics tell us nothing about the impact different performance has or may have in the problem domain. For example, a 82% correct classification rate might be sufficient for a bank to decide whether or not to approve a small credit loan application, but to classify a sample as *cancer* or *no cancer*, perhaps a 98% or higher accuracy is required. The assumption of cross-domain comparability is an illusion created by abstract metrics that have the same numeric range, but not the same meaning. Such measures tell us nothing at all useful about generalization of impact across different data sets and across different domains. Beyond abstract measures of performance (and statistical significance and effect size plus confidence interval), we need to measure the true impact a novel machine learning technique has in the real world.

With the strong focus on benchmark data and abstract performance measures, a considerable part of machine learning research in the last 20 years has concentrated on the following simplified study procedure: (1) Set machine learning task. (2) Identify standard data sets. (3) Select or generate features. (4) Choose or develop algorithm. (5) Choose performance metrics. (6) Conduct experiments. (7) Publish results in machine learning journal or conference. Emerging and future machine learning research should be more ambitious and meaningful and design a machine learning study around problems that promise an impact of machine learning in the real world. Future machine learning researchers should adopt a more challenging and potentially more rewarding procedure, which could be stated in simplified form as follows (impact aspects highlighted in *italics*): (1) Identify a challenging real-world problem for which machine learning could potentially make a crucial contribution. This is difficult and is likely to involve interdisciplinary collaboration. (2) Determine what data needs to be collected. (3) Select and generate relevant features from the data. (4) Choose or develop a relevant learning procedure or algorithm. (5) Choose existing implementation or implement the learning algorithm. (6) Select an evaluation method and procedure. (7) Perform the machine learning experiments. (8) Apply impact measures and interpret results by involving domain experts where necessary. (9) Publicize results both to the machine learning and domain-specific communities. (10) Try to convince users to adopt the new technique and tool. All these steps are a necessary component of any machine learning research program that seeks to have a real impact on the world outside of machine learning.

My experience in evaluating and reviewing EU projects tells me that particularly the ICT community sometimes does not distinguish clearly between innovation and impact. Given the recent (since FP7)

emphasis on impact, this is quite remarkable. To facilitate machine learning research that matters, we need complement traditional performance measures with evaluation measures that enable direct assessment of the impact of novel machine learning solutions. Such methods should measure the money saved, life preserved, disease burden lowered, time conserved, effort reduced, quality of life improved, loss of biodiversity decreased, crimes prevented/solved, environmental pollution reduced, level of poverty reduced, unemployment lowered, effects of catastrophes and crises mitigated, recycling of waste increased, waste of energy and resources reduced, educational learning outcomes improved, quality of goods and services enhanced, economic competitiveness improved, effects of aging populations mitigated, social inequality lowered, and so on. Such impact measures will serve to refocus and restructure machine learning research efforts in terms of the problems we tackle, the data sets we use, the way we design machine learning studies, and the objective functions we define and use. They will also motivate machine learning researchers to report how a novel feature or the performance of a new machine learning solution translates to impact in the originating problem domain.

Clearly, this type of *impact machine learning* research is inherently interdisciplinary (and ultimately transdisciplinary), and will require a change in the way we communicate and publish machine learning research results.

Machine learning for integration/analysis of complex biomedical data sets

Human health and disease are characterized by a complex interplay of multiple factors from the genome to the exposome. For many complex diseases, a sufficiently detailed understanding of the underlying mechanisms has remained elusive, and therefore the development of effective cures continues to be major challenge. As a result, the socioeconomic burden (morbidity, mortality, financial cost) of complex diseases remains high and is likely to grow within Europe's aging population. Systems medicine is an emerging interdisciplinary framework that aims to improve our understanding, prevention and treatment of complex diseases by integrating knowledge and data across multiple levels of biomedical organization. Such integration and analysis of complex, multiscale data requires a multiscale approach to machine learning, data science and modeling and simulation supported by multiscale computing. I have recently led a EU H2020 COST network grant proposal on Open Multiscale Systems Medicine which aims to integration research in this area.

Feature engineering and knowledge integration in machine learning

The selection of relevant features, and the elimination of irrelevant ones, is a central problem in machine learning. Before an induction algorithm can move beyond the training data to make predictions about novel test cases, it must decide which attributes to use for these predictions and which to ignore. Intuitively, one would like the learner to use only those attributes that are *relevant* to the target concept. In other words, the success of machine learning algorithms generally depends on (input) data representation – different data representations can entangle and hide more or less the different explanatory factors of variation behind the data. In knowledge-rich domains such as science, suitable data representations are often created by preprocessing the measured data with specific domain knowledge. This feature engineering approach is one of the most important ways to integrate domain knowledge into the machine learning process. Another approach that aims to automate the construction of relevant features is referred to as feature learning, representation learning or deep learning. I think in particular for complex scientific data sets (e.g. in biomedicine), representation learning research and feature engineering could potentially lead to fundamental breakthroughs.

Intelligent e-biomedicine

Since my degree in electrical/telecommunication engineering, my career has been strongly influenced by my interest in natural science as well as computer science. Reflecting on the increasingly fuzzy boundaries between science and technology, ICT areas such as artificial intelligence and machine learning could be viewed as the key components in the emerging field of enhanced science or e-

science (e.g. e-biomedicine). While e-science as an interdisciplinary framework (in its conventional form known as computational science or scientific computing) has been around for many years, there is still a bias towards compute-intensive modeling and simulation approaches (systems dynamics, dynamical and complex systems, control theory, and so on). The field of computational biology is a good example of this type of R&D. Recently, there has been a realization that data-intensive and intelligent technologies (machine learning, AI, computational intelligence, computational creativity, etc.) could usefully complement the standard e-science framework. For instance, the field of bioinformatics and computational biology has a long tradition in incorporating intelligent techniques into the arsenal of tools, albeit usually not in the context of large-scale computing and very large data (this is changing, however). The data-intensive element in e-science is sometimes referred to as the fourth paradigm (complementing the first three science paradigms of experiment, theory and computational science). One could argue that modeling, analysis and prediction based on the methods and tools of intelligent technologies may develop into the fifth paradigm of the evolving scientific process. I refer to this emerging framework or scientific process as: *intelligent e-science*, which could be defined as follows:

1. Experiment (first paradigm)
2. Theory (second paradigm)
3. Computational science (third paradigm)
4. Data-intensive science (fourth paradigm) – nowadays called data science
5. Science enabled by intelligent technologies (fifth paradigm)

The intelligent e-science framework as outlined above should be understood as a transdisciplinary scientific paradigm in which researchers work jointly using a shared conceptual framework and combined disciplinary-specific approaches to address complex R&D problems. Clearly, this paradigm will require considerable changes transcending the mind-set and culture of current science and education environments. The present scientific mind-set and culture is characterized by an interdisciplinary approach (where researchers work jointly but still from a disciplinary-specific basis) which has evolved from multidisciplinary science (researchers working in parallel or sequentially from disciplinary-specific base) in the past century.

It is my view that intelligent e-science – and its domain-specific versions like intelligent e-biomedicine – is likely to be at the heart of a future, highly transdisciplinary science. This type of science is likely to revolve around the processing and analyzing large amounts of data (big data). Research in the field of statistical machine learning will probably play a major role in developing this way of doing science. At present we are designing and constructing the foundation of this kind of future science. I would think that I have the qualification, experience and standing to make a constructive contribution in laying this foundation.