Alessia Visconti, PhD

Department of Twin Research, King's College London St Thomas' Hospital Campus, 3rd Floor South Wing Block D Westminster Bridge Road, London SE1 7EH

E-MAIL: alessia.visconti@kcl.ac.uk

Research Interests

- Computational Biology & Medicine
- Data Mining & Machine Learning
- Big Data
- Research Software Engineering

Brief Synopsis of Research

Alessia Visconti's research activity deals with the study of the genomics and epigenomics of human diseases. Her work focuses on the identification of genetic and epigenetics modifications influencing the risk of common diseases or of related phenotypes. She is also interested in the human microbiome and its connections to human health and diseases. She has also worked on the problem of knowledge discovery in biological data, developing new approaches tailored to solve biological tasks, and on the reverse engineering of gene regulatory networks.

EDUCATION

Jul 2012 PhD in Science and High Technology, University of Torino.

Thesis title: Systems Biology: Knowledge Discovery and Reverse Engineering

Advisors: prof. Marco Botta, Dr. Roberto Esposito

Jul 2008 Master degree in Computer Science "summa cum laude", University of Torino.

Thesis title: SPOT: an algorithm for the extraction and the analysis of biological

patterns

Advisor: prof. Marco Botta

MAR 2006 Bachelor degree in Computer Science "summa cum laude", University of Torino.

THESIS TITLE: The Haskell language Advisor: prof. Viviana Bono

SKILLS

LANGUAGE SKILLS ITALIAN: native proficiency

ENGLISH: full professional proficiency

Computing Skills Programming Languages: bash, C, C++, JAVA, php, python, R, ruby

OTHER LANGUAGES: CSS, LATEX, HTML, PyQt, XML

STATISTICAL SOFTWARE: R, SAS

DATABASE MANAGEMENT: MySQL, MariaDB

VERSION CONTROL SYSTEMS & REPRODUCIBILITY: GIT, nextflow, docker BIOINFORMATICS & GENETIC ANALYSIS: BBmap, BEDTools, DESeq2, GCTA, GWAMA, LDAK, limma, lmekin, metal, Merlin, PLINK, QTDT,

samtools, vcftools, ...

STRUCTURAL EQUATION MODELLING: openMX, Mplus

Data visualisation: dot, ggplot2

RESEARCH ACTIVITY

Aug 2017 - Present	Research fellow at the Department of Twin Research & Genetic Epidemiology, King's College London, UK
Jun 2016 - Jun 2019	Honorary research associate at the CERN Openlab at CERN, Geneva, Switzerland
APR 2015 - JUL 2017	Research associate at the Department of Twin Research & Genetic Epidemiology, King's College London, UK
Jan 2014 - Mar 2015	Research associate at the Department of Genomics of Common Disease, School of Public Health, Imperial College London, UK
Jan 2012 - Dec 2013	Research associate at the Department of Computer Science, University of Torino, Italy
Jun 2011 - Dec 2011	Visiting researcher at the Regulatory Genomics Group at the Center of Biological Sequence Analysis, Systems Biology Department, Technical University of Denmark, Denmark
Jan 2009 - Dec 2011	Awarded PhD candidate at the School of Science and High Technology, University of Torino, Italy
SEP 2008 - DEC 2008	Research assistant at the Department of Computer Science, University of Torino and in collaboration with the Department of Arboriculture and Pomology, University of Torino, Italy

TEACHING ACTIVITY

Dec 2019/Present	Instructor	for	several	Carpentries	workshops	organised	by	King's	College
	London								

- The Unix Shell
- Version Control with Git
- Programming with Python & R
- Introduction to Working with Data
- OpenRefine

Nov 2016/PRESENT **Co-organiser** of the journal club on genetic and epigenetic regulation of gene expression at the Department of Twins Research & Genetic Epidemiology, King's College London

A.Y. 2013/2014 Teaching assistant for the "Human Molecular Genetics" MSc

Department of Genomics of Common Diseases, Imperial College London

- Unix command line and ${\bf R}$
- Exploratory Data Analysis and Probability
- Quantitative genetics
- Next Generation Sequencing.

(Italian only)

A.Y. 2013/2014 Lecturer for the "Data analysis" course

Department of Biological Science, University of Torino

Lecturer for the "Operating System" course

Department of Computer Science, University of Torino

A.Y. 2012/2013 Lecturer for the "Operating System and Networking" course

Interfaculty School of Strategic Studies, University of Torino

Lecturer for the "Operating System" course

Department of Computer Science, University of Torino

A.Y. 2011/2012 **Lecturer** for the "Database" course

Department of Computer Science, University of Torino

A.Y. $2010/2011$	Lecturer for the "Database" course
	Department of Computer Science, University of Torino
	Lecturer for the "Formal Language" course
	Department of Computer Science, University of Torino
	Lecturer for the "Statistics and data mining with SAS" course
	Department of Mathematics, University of Torino
A.Y. 2009/2010	Lecturer for the "Computer Science" course Department of Letters and Philosophy, University of Torino
A.Y. 2006/2007	Teaching assistant for the "Program Languages - JAVA" course Department of Computer Science, University of Torino
A.Y. 2005/2006	Teaching assistant for the "Program Languages - JAVA" course Department of Computer Science, University of Torino
A.Y. 2004/2005	Teaching assistant for the "Program Languages - C" course Department of Computer Science, University of Torino

SUP

A.Y. 2004/2005	Teaching assistant for the "Program Languages - C" course Department of Computer Science, University of Torino
PERVISION ACTIVI	TY
A.Y. 2019/PRESENT	Assistance with the supervision of Ms Xinyuan Zhang in her PhD project at King's College London. Xinyuan's PhD project aims at studying the interplay between the gut metagenome, medication, and diseases.
A.Y. 2019/2020	Assistance with the supervision of Mr Simon Couvreur in his project at King's College London. Simon's project aimed at studying the human glycome.
A.Y. 2019/2020	Co-supervisor of Ms Xinyu Huang in her MSc project in Pharmacology at King's College London. Xinyu's project aimed at studying the interaction between medications and the gut microbiome.
A.Y. 2019/2020	Co-supervisor of Ms Helen King in her PhD rotation project at King's College London. Helen's project aimed at dissecting the interaction between gut bacterial and lipids level in blood.
A.Y. 2018/2020	Supervisor of Ms Giulia Piaggeschi during her research visit at King's College London. Giulia's project, which is part of her PhD, aimed at studying cell-specific modification in peripheral blood and their interaction with lifestyle factors.
A.Y. 2014/2020	Assistance with the supervision of Ms Marianna Sanna in her PhD project at King's College London and in her research activity at Imperial College London. Marianna's work aimed at dissecting the aetiology of melanoma and of melanoma risk phenotypes.
A.Y. 2018/2019	Supervisor of Mr Yuhao Lin's summer project as part of the King's Undergraduate Research Fellowships (KURF). Yuhao's project aimed at studying the human gut microbiome in healthy and diseased twins.
A.Y. 2018/2019	Supervisor of Ms Olivia Castellini Pérez's summer project as part of the Erasmus+ program. Olivia's project aimed at studying epigenetic plasticity triggered by tobacco smoking.
A.Y. 2018/2019	Co-supervisor of Ms Miriam Margari's master thesis in Genomic Medicine at Imperial College London, entitled: "Identification of novel genomic imprinting effects on gene expression in human tissues".
A.Y. 2016/2019	Co-supervisor of Mr Niccolò Rossi during his research visit at King's College London. Niccolò's PhD project aimed at identifying and characterising the causes of lipid-metabolism disruption in patients with severe and unexplained familial dyslipidemia.

A.Y. 2017/2018	Supervisor of Ms Lechun Huo's summer project as part of the King's Undergraduate Research Fellowships (KURF). Lechun's project aimed at studying the human gut microbiome in twins.
A.Y. 2017/2018	Co-supervisor of Dr Robin Mesnage's master thesis in Bioinformatics at Birkbeck University of London, entitled: "A Metagenome-wide association study suggests that glycome composition associates with pathogenic bacteria abundance in the gut microbiome".
A.Y. 2016/2018	Assistance with the supervision of Dr Harriet Cullen during her research fellowship at King's College London. Harriet's project aimed at dissecting the genetic and epigenetic basis of pre-term delivery.
A.Y. 2016/2017	Supervisor of Ms Yuri Nemoto's summer project as part of the King's Undergraduate Research Fellowships (KURF). Yuri's project aimed at identifying connection between human gut microbiome and lipid profiles in twins.
A.Y. 2016/2017	Co-supervisor of Ms Fudi Wang's research visit. Fudi's project aimed at identifying connections between facial ageing features and a set of genomic loci.
A.Y. 2015/2016	Co-supervisor of Ms Esther Kok's summer internship at the CERN. Esther's project aimed at developing an efficient workflow to detect structural variants in DNA sequence data.
A.Y. 2013/2014	Co-supervisor of Mr Marcin Świstak's internship at Imperial College London. Marcin's project aimed at dissecting the genetic basis of melanoma and its connection with ageing.
A.Y. 2013/2014	Co-supervisor of Mr George Powell's master thesis in Human Molecular Genetics entitled: "Enrichment of Genomic Runs of Homozygosity for Copy Number Variation in Population Cohorts and Family Trios"
A.Y. 2010/2011	Co-supervisor of Mr Marco Gallizio's bachelor thesis in Computer Science entitled: "A web interface for querying the Restructured Gene Ontology", University of Torino.

OTHER ACTIVITIES

PARTICIPATION IN RESEARCH PROJECTS

"Dissecting the mechanisms of immune-mediated inflammation: a systems-immunology approach", funded by MRC-2019-2021

Role: Researcher

"A high resolution map of copy number and structural variation in Qatari genomes and their contribution to quantitative traits and disease", funded by Qatar Foundation – 2016-2018

Role: Researcher, Contribution to project proposal

"An integrative genomics approach for non-invasive diagnostic biomark-- ers discovery in IgA nephropathy", funded by MRC – 2014-2016 ROLE: Researcher

"Senescence and melanoma – An integrative systems biology approach to characterise the link between reduced biological senescence and melanoma susceptibility", funded by British Skin Foundation – 2013-2015

Role: Researcher

"Genomic analysis of Type 2 Diabetes in Qatar, towards diabetes personalized medicine", funded by Qatar Foundation – 2013-2018

Role: Researcher

"LIMPET – Isotropic And Anisotropic Lipophilicity To Model Permeability Of New Therapeutic Peptides", funded by Compagnia di San Paolo – 2012-2013

Role: Researcher

"BioBITs - Developing white and green biotechnologies by converging platforms from biology and information technology towards metagenomics", funded by Regione Piemonte - 2007-2011

Role: Researcher

"Realizzazione di modelli informatici per la valorizzazione della qualità e la tracciabilità delle produzioni in specie da frutto coltivate in Piemonte",

funded by Regione Piemonte – 2004-2009

Role: Developer

PARTICIPATION IN INTERNATIONAL CHALLENGES

"BioDataHack 2018 - Genomic, Biodata and Improving Health Out-

RANK: 1st on the ARM, Cavium, and Atos Challenge: How can we use mobile technology to transform biological data processing?

"DREAM5 - Network Inference Challenge"

- Rank: 15th out of 29 participants when both synthetic and real network are considered, 3rd on real networks, and 1st on S. Cerevisiae's network

"DREAM6 - Promoter Activity Prediction Challenge"

Rank: 8th out of 21 participants

Administrative tasks

- Nov 2016 present: co-organiser of the journal club on genetic and epigenetic regulation of gene expression at the Department of Twins Research & Genetic Epidemiology, King's College London
- Jan 2012 Dec 2013: faculty member as representative of postdoctoral fellows at the Department of Computer Science, University of Torino
- Jan 2009 Dec 2011: faculty member as representative of PhD students fellows at the Department of Computer Science, University of Torino

Extracurricular ACTIVITIES

- Jul 2017 present: member and mentor of the Artificial Intelligence Club for Gender Minorities, which aims at promoting gender diversity in the artificial intelligence and scientific community via meetups, and mentorship. Alessia Visconti organised workshops on collaborative data science via Git and GitHub. Since May 2018, she is also co-organising the group monthly journal club.
- Mar 2016 Sep 2017: member, tutor, and mentor of the RLadies London community and of Researc/her/ Research community. These groups aim at promoting gender diversity in the R and STEM community via meetups, mentorship and global collaboration. With Researc/her, Alessia Visconti gave speeches on reproducibility, and workflow development.

• -omics of human diseases.

Advances in -omics data collection created an opportunity to identify factors influencing the risk of common diseases. Alessia Visconti is mainly involved in a set of projects aiming at dissecting the aetiology of melanoma, melanoma risk phenotypes, and their connection with ageing [J11, J12, J15, J16, J19, J21, J24, J25, J29, A5]. She also used systems biology and bioinformatics approaches to study cognition and neurodevelopmental disease [J7, J20], reading and language disabilities [J9], epigenetic plasticity [A6] and modification [J14], thyroid diseases [J23], and IgA Nephropathy [J10]. Alessia Visconti developed a framework for pairwise association testing in related samples [J13], that has been used to perform the first epigenome-wide association study in an Arab population [J8]. She also reported on how to conduct metagenomic studies in microbiology and clinical research [J18], and developed a novel pipeline which ensure reproducibility in metagenomics research [J17] which she use to study the interplay between the human gut microbiome and the host metabolism [J22], while collaborating in other metagenomic studies [J31].

• REVERSE ENGINEERING OF GENE REGULATORY NETWORKS.

The reverse engineering problem, *i.e.*, the inference of gene regulatory networks from data, is a cardinal task on the biological research agenda. Alessia Visconti worked on two approaches that allow the reverse engineering of gene regulatory networks and applied them to several organisms (*E. Coli*, *S. Cerevisiae*, *S. Pombe*). In the first approach a Naive Bayes-based framework has been developed. It merges multiple pieces of information derived from microarray experiments [J3, P4]. In the second approach a method aiming at deciphering temporal influences among genes and proteins has been proposed [T2].

• Data mining techniques for biological data analysis.

Data mining allows the extraction of previously unknown interesting knowledge from large data sets. Alessia Visconti worked on two data mining methods: coclustering and biclustering. Specifically, she developed techniques that allow the exploitation of domain knowledge and multiple data sources to improve clustering results. These approaches have been used: *i*) to identify protein sequences characterised by common patterns [T1, P1, A2], *ii*) to study synthenies in microorganisms [J1], *iii*) to analyse RNA secondary structure [A1], and *iv*) to discover groups of genes showing similar expression profiles under the same set of experimental conditions [J5, P2, BC1, T2].

• Machine learning techniques for solving biological problems.

Machine learning focuses on the development of algorithms that improve through experience. An important application of Machine Learning is the prediction of new knowledge from patterns learnt from data. Alessia Visconti leveraged and combined machine learning approaches to deal with several biological problems. Past works based on this approach comprise: *i*) the prediction of promoter activities from promoter sequences, *ii*) the identification of pharmacogenes [A3, T2], and *iii*) the study of peptide-based drugs [J4, J6, A4].

• GENE ONTOLOGY RESTRUCTURATION.

Gene Ontology represents a collaborative effort to provide a structured vocabulary for consistent gene descriptions. Although Gene Ontology facilitates information retrieval, its structure may hide some useful knowledge, such as gene cooperation. Alessia Visconti worked on a restructuration of Gene Ontology, called RGO, that is useful for emphasising genes involved in the same biological process and for inferring new pieces of information about gene functions and localisation. The RGO enhances some kinds of automated analysis, such as gene profiling and clustering, statistical enrichment, as well as the evaluation of gene functional similarities [J2, P3, T2].

• Probabilistic Graphical Models.

Probabilistic Graphical Models (PGMs) sport a rigorous theoretical foundation and provide an abstract language for modeling application domains. Answering Maximum a Posteriori queries over a PGM entails finding the assignment to the graph variables that *globally* maximises the probability of an observation. Alessia Visconti contributed to the development of a novel exact algorithm for answering Maximum a Posteriori queries on tree-structured PGMs [P5].

Publications

International Journal

- [J32] Hopkinson N.S.[†], Rossi N.[†], El-Sayed Moustafa J., Laverty A.A., Quint J.K., Freidin M., **Visconti** A., Murray B., Modat M., Ourselin S., Small K., Davies R., Wolf J., Spector T.D., Steves C.J.[‡], and Falchi M.[‡], Current smoking and COVID-19 risk: results from a population symptom app in over 2.4 million people, Thorax, 2021, doi:10.1136/thoraxjnl-2020-216422
- [J31] Bar N.[†], Korem T.[†], Weissbrod O., Zeevi D., Rothschild D., Leviatan S., Kosower N., Lotan-Pompan M., Weinberger A., Le Roy C.I., Menni C., **Visconti A.**, Falchi M., Spector T.D., The IMI DIRECT consortium, Adamski J., Franks P.W., Pedersen O. and Segal E., *A reference map of potential determinants for the human serum metabolome*, Nature, 2020, doi:10.1038/s41586-020-2896-2
- [J30] Zazzara M.B.[†], Penfold R.S.[†], Roberts A.L.[†], Lee, K.A., Dooley H., Sudre C.H., Welch C., Bowyer R.C.E, **Visconti A**, ..., Martin F.C., Steves C.J.[‡], Lochlainn M.N.[‡], *Probable delirium is a presenting symptom of COVID-19 in frail, older adults: a cohort study of 322 hospitalised and 535 community-based older adults*, Age and Ageing, 2020, doi:10.1093/ageing/afaa223
- [J29] Sanna M.[†], Li X.[†], **Visconti A.**, Freidin M. B., Sacco C., Ribero S., Hysi P., Bataille V., Han J.[‡], and Falchi M.[‡], Looking for Sunshine: Genetic Predisposition to Sun-Seeking in 265,000 Individuals of European Ancestry, Journal of Investigative Dermatology, 2020, doi:10.1016/j.jid.2020.08.014
- [J28] Lee K.A.[†], Ma W.[†], Sikavi D.R., ..., **Visconti A.**, ..., Ourselin S., Spector T.D., and Chan A.T., COPE consortium, *Cancer and Risk of COVID-19 Through a General Community Survey*, Oncologist, 2020, doi:10.1634/theoncologist.2020-0572
- [J27] Scarfi F., Orozco A.P., Visconti A., and Bataille V., An Aggressive Clinical Presentation of Familial Leiomyomatosis Associated with a Fumarate Hydratase Gene Variant of Uncertain Clinical Significance, Acta Dermato-venereologica, 2020, doi:10.2340/00015555-3573
- [J26] Menni, C.[†], Valdes, A. M.[†], Freidin, M. B., Sudre, C. H., Nguyen, L. H., Drew, D. A., Ganesh, S., Varsavsky, T., Cardoso, M. J., El-Sayed Moustafa, J. S., Visconti, A., Hysi, P., Bowyer, R. C. E., Mangino, M., Falchi, M., Wolf, J., Ourselin, S., Chan, A. T., Steves, C. J.[‡], and Spector, T. D.[‡], Real-time tracking of self-reported symptoms to predict potential COVID-19, Nature Medicine, 2020, doi:10.1038/s41591-020-0916-2
- [J25] Landi M.T., Bishop D.T., MacGregor S., ..., Visconti A., ..., Shi J., Iles M.M. and Law M.H., Genome-wide association meta-analyses combining multiple risk phenotypes provide insights into the genetic architecture of cutaneous melanoma susceptibility, Nature Genetics, 2020, doi:10.1038/s41588-020-0611-8
- [J24] Visconti A., Sanna M., Bataille V., and Mario F., Genetics plays a role in nevi distribution in women, Melanoma Management, 2020, doi:10.2217/mmt-2019-0019 [Invited editorial]
- [J23] Martin T.C., Illieva K.M., Visconti A., Beaumont M., Kiddle S.J., Dobson R.J.B., Mangino M., Lim E.M., Pezer M., Steves C.J., Bell J.T., Wilson S.G., Lauc G., Roederer M., Walsh J.P., Spector T.D.[‡], Karagiannis S.N.[‡], Dysregulated Antibody, Natural Killer Cell and Immune Mediator Profiles in Autoimmune Thyroid Diseases, MDPI Cells, 2020, doi:10.3390/cells9030665
- [J22] Visconti A.[†], Le Roy C.I.[†], Rosa F., Rossi N., Martin T.C., Mohney R.P., Li W., de Rinaldis E., Bell J.T., Venter J.C., Nelson K.E., Spector T.D.[‡], and Falchi M.[‡], *Interplay between the human gut microbiome and host metabolism*, Nature Communications, 2019, doi:10.1038/s41467-019-12476-z
- [J21] Visconti A., Ribero S., Sanna M., Spector T.D., Bataille V., and Mario F., Body site-specific genetic effects influence naevus count distribution in women, Pigmented Cell & Melanoma Research, 2019, doi:10.1111/pcmr.12820
- [J20] Cullen H., Krishnan M.L., Selzam S., Ball G., Visconti A., Saxena A., Counsell S.J., Hajnal J., Breen G., Plomin R., and Edwards, A.D. Polygenic risk for neuropsychiatric disease and vulnerability to abnormal deep grey matter development, Scientific Reports, 2019, doi:10.1038/s41598-019-38957-1

[†] indicates that the authors contributed equally to the work

[‡] means that the authors jointly supervised the work

- [J19] Duffy D., Zhu G., Li X., ..., Visconti, A., ..., Falchi M., Han J.[‡], Martin N.G.[‡], Melanoma GWAS Consortium Novel pleiotropic risk loci for melanoma and nevus density implicate multiple biological pathways, Nature Communications, 2018, doi: 10.1038/s41467-018-06649-5
- [J18] Martin T.C.[†], **Visconti A**[†], Tim D. Spector, and Falchi M., Conducting metagenomic studies in microbiology and clinical research, Appl Microbiol Biotechnol, 2018, doi: 10.1007/s00253-018-9209-9
- [J17] Visconti A, Martin T.C., and Falchi M., YAMP: a containerised workflow enabling reproducibility in metagenomics research, GigaScience, 2018, doi: 10.1093/gigascience/giy072
- [J16] Visconti, A., Duffy, D., Liu, F., Zhu, G., ..., Han, J., Bataille, V., and Falchi, M., Genome-wide association study in 176,678 Europeans reveals genetic loci for tanning response to sun exposure, Nature Communications, 2018, doi: 10.1038/s41467-018-04086-y
- [J15] Hysi, P.G.[†], Valdes, A.M.[†], Liu, F.[†], Furlotte, N.A., Evans, D.M., Bataille, V., **Visconti, A.**, ..., Kayser, M.[‡], and Spector, T.D.[‡], Genome-wide association meta-analysis of individuals of European ancestry identifies new loci explaining a substantial fraction of hair color variation and heritability, Nature Genetics, 2018, doi: 10.1038/s41588-018-0100-5
- [J14] Zaghlool, S.B., Mook-Kanamori, D.O., Kader,S., Stephan, N., Halama, A., Engelke, R., Sarwath, H., Al-Dous, E. K., Mohamoud, Y. A., Roemisch-Margl, W., Adamski, J., KastenmÃ¹/₄ller, G., Friedrich, N., Visconti, A., ..., Malek, J.A., and Suhre K, Deep molecular phenotypes link complex disorders and physiological insult to CpG methylation, Human Molecular Genetics, 2018, doi: 10.1093/hmg/ddy006
- [J13] Visconti A., Al-Shafai M., Al Muftah W.A., Zaghlool S.B., Mangino M., Suhre K., and Falchi M., PopPAnTe: population and pedigree association testing for quantitative data, BMC Genomics, doi: 10.1186/s12864-017-3527-7
- [J12] Puig-Butille J.A., Gimenez-Xavier P., Visconti A., Nsengimana J., Garcia-Garcia F., Tell-Marti G., Escamez M.J., Newton-Bishop J.A., Bataille V., Del Rio M., Dopazo J., Falchi M, and Puig S., Genomic expression differences between cutaneous cells from red hair colour individuals and black hair colour individuals based on bioinformatic analysis., Oncotarget, 2016, doi:10.18632/oncotarget.14140
- [J11] Ribero S.[†], Sanna M.[†], Visconti A., Navarini A., Aviv A., Glass D., Spector T.D., Smith C., Simpson M., Barker J., Mangino M., Falchi M.[‡], and Bataille V.[‡], Acne and telomere length. A new spectrum between senescence and apoptosis pathways, The Journal of Investigative Dermatology, 2016, doi:10.1016/j.jid.2016.09.014
- [J10] Lomax-Browne H.J.[†], **Visconti A.**[†], Pusey C.D., Cook H.T., Spector T.D., Pickering M.C[‡], and Falchi M[‡], *IgA Glycosylation is Heritable in Healthy Twins*, Journal of the American Society of Nephrology, 2016, doi:10.1681/ASN.2016020184
- [J9] Gialluisi A., Visconti A., Willcutt E.G., Smith S.D., Pennington B.F. Falchi M., DeFries J.C., Olson R.K., Francks C., and Fisher S.E., Investigating the effects of copy number variants on reading and language performance, Journal of Neurodevelopmental Disorders, 2016, doi:10.1186/s11689-016-9147-8
- [J8] Al Muftah W.A.[†], Al-Shafai M.[†], Zaghlool S.B., **Visconti A.**, Tsai P.C., Kumar P., Spector T., Bell J., Falchi M.[‡], and Suhre K.[‡], *Epigenetic associations of type 2 diabetes and BMI in an Arab population*, Clinical Epigenetics, 2016, doi:10.1186/s13148-016-0177-6
- [J7] Johnson M.R., Shkura K., Langley S.R., ..., Visconti A., ..., Kaminski R.M., Deary I.J., and Petretto E., Systems genetics identifies a convergent gene network for cognition and neurodevelopmental disease, Nature Neuroscience, 2015, doi:10.1038/nn.4205
- [J6] Visconti A., Ermondi G., Caron G., and Esposito R., Prediction and Interpretation of the Lipophilicity of Small Peptides. Journal of Computer-Aided Molecular Design, 2015, pp. 1-10
- [J5] Visconti A., Cordero F., and Pensa R.G., Leveraging additional knowledge to support coherent bicluster discovery in gene expression data. Intelligent Data Analysis, 18:5, 2014, pp. 837-855
- [J4] Ermondi G., **Visconti A.**, Esposito R., and Caron G., The Block Relevance (BR) analysis supports the dominating effect of solutes hydrogen bond acidity on $\Delta \log P_{oct\text{-}tol}$. European Journal of Pharmaceutical Sciences, 53, 2014, pp. 50-45

- [J3] Marbach D., Costello J.C., Küffner R., Vega N., Prill R.J., Camacho D., Allison K.R., ..., Visconti A., ..., Kellis M., Collins J.J., and Stolovitzky G., Wisdom of crowds for robust gene network inference. Nature Methods, Vol. 9, 2012, pp. 796-804
- [J2] Visconti A., Esposito R., and Cordero F., Restructuring the Gene Ontology to Emphasize Regulative Pathways and to Improve Gene Similarity Queries. Int. J. Computational Biology and Drug Design, Vol. 4, No. 3, 2011, Inderscience Publishers, pp. 220-238
- [J1] Bonfante P., Cordero F., Ghignone S., Ienco D., Lanfranco L., Leonardi G., Meo R., Montani S., Roversi L., and Visconti A., A Modular Database Architecture Enabled to Comparative Sequence Analysis. LNCS Transactions on Large-Scale Data- and Knowledge-Centered Systems - TLDKS IV, LNCS 6990, 2011, Springer, pp. 124-147

In proceeding

- [P5] Esposito R, Radicioni D.P., and Visconti A., CDoT: optimizing MAP queries on trees. In proceedings of AI*IA 2013: Advances in Artificial Intelligence, XIIIth Int. Conf. of the Italian Association for Artificial Intelligence, Torino, December 4-6, 2013, LNAI 8249, pp. 481–492. Springer
- [P4] Visconti A., Esposito R., and Cordero F., Tackling the DREAM Challenge for Gene Regulatory Networks Reverse Engineering. In Proceedings of AI*IA 2011: Artificial Intelligence Around Man and Beyond, XIIth Int. Conf. of the Italian Association for Artificial Intelligence Palermo, September 15-17, 2011, LNAI 6934, pp. 373-383, Springer
- [P3] Visconti A., Cordero F., Botta M., Calogero R.A., Gene Ontology rewritten for computing gene functional similarity. In Proceedings of the Fourth International Conferences on Complex, Intelligent and Software Intensive Systems, February 15-18, 2010, IEEE Computer Society Press, pp. 694-699
- [P2] Cordero F., Pensa R.G, Visconti A., Ienco D. and Botta M., Ontology-driven Co-clustering of Gene Expression Data. In proceedings of AI*IA 2009: Emergent Perspectives in Artificial Intelligence, XI Int. Conf. of the Italian Association for Artificial Intelligence - Reggio Emilia, December 9-12, 2009, LNAI 5883, pp. 426-435, Springer
- [P1] Cordero F., Visconti A., and Botta M., A new protein motif extraction framework based on constrained co-clustering. In Proceedings of the 24th Annual ACM Symposium on Applied Computing March 8-12, 2009, ACM Press, pp. 776-781

BOOK CHAPTERS

[BC1] Visconti A., Cordero F., Ienco D., and Pensa R.G., Coclustering under Gene Ontology Derived Constraints for Pathway Identification. Biological Knowledge Discovery Handbook: Preprocessing, Mining and Postprocessing of Biological Data, Mourad Elloumi and Albert Y. Zomaya (Eds.), 2014, John Wiley & Sons, pp. 625-642

Pre Prints

[PP1] Bataille V.†, **Visconti A.**†,, Rossi N., Murray B., Bournot A., Wolf J., Ourselin S., Steves C., Spector T.D.‡, and Falchi M.‡, *Diagnostic value of skin manifestation of SARS-CoV-2 infection*, medRxiv, 2020, doi:10.1101/2020.07.10.20150656

Selected Abstracts and Posters

- [A6] Carnero-Montoro E., **Visconti A.**, Sacco C, Tsai P.C, Spector T.D, Falchi M., and Bell J.T., Environmentally-induced epigenetic variability is associated with metabolic traits, American Society of Human Genetics, October 2016
- [A5] Świstak M., Visconti A., Falchi M., Bataille V., and Spector T.D., Differential expression and coexpression analysis across multiple tissues in twins. EMBO Young Scientists Forum, Warsaw, July 2015, pp. 192

- [A4] Ermondi G., Esposito R., Visconti A., Visentin S., Vallaro M., Rinaldi L, and Caron G., Application of in-silico "classical" drug discovery tools to peptide research. NovAliX Conference 2013, Biophysics in drug discovery, Strasbourg, October 2013, pp. P14
- [A3] Visconti A., Calogero R.A, and Cordero F., Improving biomarker discovering for chemosensitivity prediction using an integrated approach. 9th Annual Meeting of the Italian Society of Bioinformatics (BITS), EMBnet.journal, Supplement A, April 2012, pp. 24
- [A2] Cordero F., Visconti A., and Botta M., A web interface to extract protein motif by constrained co-clustering. RECOMB Regulatory Genomics 2008, Boston, October 23-November 3, 2008.
- [A1] Cordero F., Visconti A., and Botta M., A motif extraction framework applied on RNA secondary structure. Alternative Splicing Workshop Milano, October 3, 2008.

THESIS

- [T2] Visconti A., Systems Biology: Knowledge Discovery and Reverse Engineering. PhD Thesis, Doctoral School in Science and High Technology, University of Turin, 2012.
- [T1] Visconti A., SPOT: an algorithm for the extraction and the analysis of biological patterns. Master Thesis, Department of Computer Science, University of Turin, 2008.

Software

The software is available at http://compbio.di.unito.it, http://twinsuk.ac.uk/resources-for-researchers/software/, https://github.com/alesssia, or upon request.

AID-ISA extracts biologically relevant biclusters from microarray gene expression data by leveraging additional knowledge

CDoT is a novel exact algorithm for answering Maximum a Posteriori queries on tree-structured Probabilistic Graphical Models

FAMCNV (v2.0) enables genome-wide association of copy number variants with quantitative phenotypes in families

GOCLUST performs a coclustering of microarray gene expression data by means of Gene Ontology-derived constraints

MOTIFSLINKER associates clusters of proteins with their frequent motifs

PopPAnTE enables pairwise association testing in related samples

RGO is a reorganization of the Gene Ontology emphasing regulative information and providing better structure for gene functional analysis

SPOT performs an exhaustive search of frequent motifs in sets of biological sequences

YAMP allows processing raw metagenomic sequencing data up to the functional annotation

London, January 6, 2021