Assistant professor (tenure track)

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Research Interests

- BIOSTATISTICS & BIOINFORMATICS
- Computational Biology & Medicine
- Data Mining & Machine Learning
- Multi-omics data analysis and integration
- Research Software Engineering

BRIEF DESCRIPTION

Alessia Visconti is an expert in bioinformatics and genetic epidemiology. Her research activity deals with the development and application of statistical and computational methods to identify multi-*omics* modifications influencing complex human phenotypes. 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 Computer Science , University of Turin. THESIS TITLE: Systems Biology: Knowledge Discovery and Reverse Engineering
Jul 2008	Master degree in Computer Science "summa cum laude", University of Turin. THESIS TITLE: SPOT: an algorithm for the extraction and the analysis of biological patterns
Mar 2006	Bachelor degree in Computer Science "summa cum laude", University of Turin. THESIS TITLE: The Haskell language

SKILLS

Language Skills	ITALIAN: native proficiency ENGLISH: professional working proficiency GREEK (MODERN): elementary proficiency
Computing Skills	PROGRAMMING LANGUAGES: bash, C, C++, JAVA, php, python, R, ruby OTHER LANGUAGES: CSS, IATEX, HTML, PyQt, XML STATISTICAL SOFTWARE: R, SAS DATABASE MANAGEMENT: MySQL, MariaDB VERSION CONTROL SYSTEMS & REPRODUCIBILITY : GIT, nextflow, docker, singularity 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

Assistant professor (tenure track) at the Center for Biostatistics, Epi-DEC 2023 - PRESENT demiology and Public Health, Department of Clinical and Biological Sciences, University of Turin, Italy RESEARCH ACTIVITY: Alessia Visconti is responsible for (a) the development and application of statistical and computational multi-omics approaches to understand the mechanisms and to improve the diagnosis and treatment of human diseases, (b) the design and deployment of Deep Learning approaches for monitoring and improving public health responses, and for identifying medical vulnerabilities that exist within communities, (c) the teaching of under and post-graduates courses, and (d) the preparation of research grants, of which she has also the scientific responsibility. Alessia Visconti remains lead the microbiome data analysis at the Department of Twin Research & Genetic Epidemiology, King's College London, UK, where she explores the effect of the gut microbiome on human health [J53, J54], with a particular interest on its phage component [J52, PP2], and continues her research on IgA glycosylation [PP1]. Senior bioinformatician at the Genomics Research Centre, Human Ост 2023 - Nov 2023 Technopole, Italy RESEARCH ACTIVITY: Alessia Visconti is involved in a project performing genome-wide association study (GWAS) on phenotypes acquired from cardiac MRIs via an unsupervised deep-learning approach. She remains a visitor at the Genomics Research Centre. Aug 2017 - Sep 2023 Research fellow at the Department of Twin Research & Genetic Epidemiology, King's College London, UK RESEARCH ACTIVITY: Alessia Visconti was responsible for (a) the development and application of statistical and computational multi-omics approaches to understand the mechanisms and to improve the diagnosis and treatment of human diseases, (b) the supervision of master and PhD students as well as postdocs, (c) the preparation of research grants, of which she has also the scientific responsibility From her previous role, Alessia Visconti continued to lead the bioinformatics analyses for a set of studies aiming at dissecting the aetiology of melanoma and its risk phenotypes [J21, J24, J25, J29] and at studying IgA Nephropathy [J38]. She was also responsible for several new projects aiming at investigating, among the others, the influence of the gut microbiome on human health [J22, J31, J41, J44, J45, J48, J49, J50], and predicting melanoma response to immunotherapy [J43, J46], and at studying thyroid diseases [J23], atopic dermatitis [J39, J51], cardiovascular diseases and their risk factors [J35], immune system modifications [J36], and X-inactivation [J47]. During the SARS-CoV-2 pandemic, she (a) developed the analysis pipeline for the daily data provided by more than four million users [J40], (b) led two studies investigating skin manifestations of SARS-CoV-2 [J33, J42], and (c) performed the bioinformatics analysis for several other studies [J26, J28, J30, J32, J34, J37]. Honorary research associate at CERN OpenLab, Switzerland Jun 2016 - Jun 2019 RESEARCH ACTIVITY: Alessia Visconti extended the ROOT library (https: //root.cern/), developed by the CERN OpenLab for enabling the storage and scientific analyses and visualisation of large amounts of data from particle physics experiments, to allow the efficient storage of genomic data. Apr 2015 - Jul 2017 Research associate at the Department of Twin Research & Genetic Epidemiology, King's College London, UK

	RESEARCH ACTIVITY: Alessia Visconti was mostly responsible for the develop- ment and application of statistical and computational multi-omics approaches to understand the mechanisms and to improve the diagnosis and treatment of human diseases. She also supervised master and PhD students and con- tributed to the writing of research grants. In this role, Alessia Visconti led the bioinformatics analyses for a set of projects aiming at dissecting the ae- tiology of melanoma, melanoma risk phenotypes, and their connection with ageing [J11, J15, J16, J19], and at studying IgA Nephropathy [J10], cognition and neurodevelopmental disease [J20], and epigenetic modification [J14]. She also reported on how to conduct metagenomic studies in microbiology and clinical research [J18] and developed a novel pipeline which ensures re- producibility in metagenomics research [J17].
Jan 2014 - Mar 2015	Research associate at the Department of Genomics of Common Disease, School of Public Health, Imperial College London, UK RESEARCH ACTIVITY: Alessia Visconti (a) developed and implemented a novel approach for the population and pedigree association testing for quantitative data [J13], and (b) conducted the bioinformatics analysis for [J7, J8, J9, J12].
Jan 2012 - Dec 2013	Research associate at the Department of Computer Science, University of Turin, Italy RESEARCH ACTIVITY: Alessia Visconti (a) developed and implemented a novel bi-clustering approach leveraging additional knowledge [J5], (b) contributed to the development of a novel exact algorithm for answering Maximum a Pos- teriori queries on tree structures [P5], (c) applied machine learning approaches for the prediction and interpretation of the lipophilicity of small peptides [J6], and (d) conducted the bioinformatics analysis to measure the ability of more than 200 compounds of acting as hydrogen bond donors [J4].
Jun 2011 - Dec 2011	Visiting researcher at the Center of Biological Sequence Analysis, Technical University of Denmark, Denmark RESEARCH ACTIVITY: Alessia Visconti (a) continued her PhD project develop- ing a new method for the reverse engineering of gene regulatory networks that uses a popular econometrics statistical hypothesis test, namely the Granger Causality [T3], and (b) developed an <i>ensemble</i> approach for the prediction of promoter activity.
Jan 2009 - Dec 2011	PhD student at Department of Computer Science, University of Turin, Italy RESEARCH ACTIVITY: Alessia Visconti developed and implemented: (a) a de novo framework (accompanied by a web interface) performing protein motifs identification and allowing the simultaneous associations between groups of protein sequences and groups of motifs thanks to a constrained co-clustering approach [P1], (b) a new methodology (accompanied by a web interface) that provides meaningful co-clusters whose discovery and interpretation are enhanced by embedding gene ontology (GO) annotations [BC1, P2], (c) a novel algorithm for the rewriting of the GO aiming at obtaining a more compact and informative ontology [J2, P3], (d) two algorithms for the reverse engineering of gene regulatory networks [P4, J3, T3], and (e) contributed to a modular framework for the analysis of metagenomics sequences leading the co-clustering module development [J1].

CAREER BREAKS

Aug 2022 - Apr 2023 Maternity leave

AWARDS

Summer 2019	Awarded a mini-grant $(\pounds 1000)$ to hire an undergraduate student through
	the "King's Undergraduate Research Fellowship (KURF)".
Summer 2018	Awarded a mini-grant $(\pounds 1000)$ to hire an undergraduate student through
	the "King's Undergraduate Research Fellowship (KURF)".

July 2018	Winner of one of the challenges of the "BioDataHack 2018 - Genomic,
	<i>Biodata and Improving Health Outcomes</i> ". The project presented by Alessia Visconti and the other members of her team (Jun Aruga, Oliver Giles, Ioannis
	Valasakis e Chen Zhang) advanced the vision of a device that will allow the
	constant monitoring of IBD by patients from the comfort of their own homes,
	ranked first on the ARM, Cavium, and Atos Challenge: <i>How can we use</i>
	mobile technology to transform biological data processing?. During the two-day BioData Hackathon, the team successfully ported the metagenomics pipeline
	developed by Alessia Visconti [J17] onto Arm's 64-bit architecture, where the
	data could be processed in a few hours, showing that the analysis of microbial
	data can be successfully taken out of centralised data centres.
	The solution also implemented a neural network that, receiving as input the
	microbial profile produced by the analysis pipeline, could predict the disease
	status. Even with a very small set of training data, the proposed approach
	was able to separate patients with Crohn's disease and ulcerative colitis (the
	two main forms of IBD) from healthy controls with over 90% accuracy.
	The team also proposed a derivative score from the neural network's output. Such a score, which could be tracked over time, would allow individuals to
	measure the effects of lifestyle/medication changes on their disease's progres-
	sion to assess their efficacy in almost real-time. The next step would be to use
	the data generated in this monitoring process to craft suggestive models, able
	to offer individuals advice and treatments based on what has been effective in
	patients with similar microbiome profiles.
Summer 2017	Awarded a mini-grant (£1000) to hire an undergraduate student through the "King's Undergraduate Research Fellowship (KURF)".
Jan 2012 - Dec 2013	Awarded two years postdoctoral fellowship awarded by the Italian Min- ister of Education, University and Research
Jan 2012 - Dec 2012	Awarded a postdoctoral training grant (€3000) awarded by the Regione
	Piemonte
Summer 2011	Participation to the "DREAM6 – Promoter Activity Prediction Challenge".
	The proposed approach (developed with Ali Altintas and Chris Workman)
	which combined results from two well-known machine learning approaches
	(regression trees and support vector machines for regression), ranked 8th out of 21 participants.
Summer 2010	Winner of one of the challenges of the "DREAM5 – Network Inference
Somment 2010	<i>Challenge</i> ". The ensemble approach [P4, J3] developed with the other team
	members (Roberto Esposito and Francesca Cordero), which was compared
	with other 35 approaches and which combines multiples approaches within
	a Naive Bayes classifier, despite its simplicity, ranked third for the reverse
	engineering of real organisms' gene regulatory networks and first for the re-
LAN 2000 Drg 2011	construction of the <i>Saccharomyces Cerevisiae</i> 's network. Awarded three years PhD fellowship awarded by the Italian Minister of
Jan 2009 - Dec 2011	Education, University and Research (best PhD project)

RESEARCH COLLABORATIONS

PROJECTS FOR WHICH ALESSIA VISCONTI HAS SCIENTIFIC RESPONSIBILITY

2022-2025 TITLE: "Challenging the Dogma of Homogeneity in Gestational Diabetes Mellitus" FUNDER: MRC Medical Research Council – BUDGET: £1,090,268 ROLE: In this project, which is now in the phase of data collection and aims at characterising pathophysiologically distinct subtypes of gestational diabetes (GDM), Alessia Visconti will develop and use bioinformatics and machine learning approaches to (a) evaluate similarities and differences in GDM subtypes in women of White European and South Asian descent, identifying variables (clinical/biochemical) that distinguish between subtypes, and (b) explore the relationships between subtypes and maternal/fetal/neonatal outcomes.

2022-2023	TITLE: "A Collaborative Approach to the Borne Uterine Mapping Programme (BUMP) Feasibility Study"
	FUNDER: BORNE – BUDGET: £500,000
	ROLE: In this project, which is now in the first phase of data collection, Alessia Visconti will develop (a) pipelines for the analysis of single-cell and single-nucleus RNA sequencing and spatial transcriptomic data, and (b) deep-learning approaches for modelling their interaction.
2021-2024	TITLE: "Understanding phenotype and mechanisms of spontaneous preterm birth in sub- Saharan Africa (PRECISE-SPTB)"
	FUNDER: MRC Medical Research Council – BUDGET: £458,80
2020-2021	ROLE: In this project, which aims at determining the epidemiological and contextual na- ture of spontaneous preterm birth in three sub–Saharan African countries (Kenya, The Gambia and Mozambique), while developing technical infrastructure and training re- search scientists, Alessia Visconti is responsible for the preparation and delivery of a set of workshops which cover both basic and specialistic skills, namely: shell programming, version control and collaboration with Git/GitHub, programming in R, programming in python, workflow development with Nextflow, machine-learning approaches for biomedi- cal data analysis, and metagenomic data analysis. Workshops delivered so far have been rated as "exceptional" by the attendees. TITLE: "A multi-omics study to dissect the role of the gut microbiome in IgA nephropathy
	risk"
	FUNDER: King's College London - Peking University Health Science Centre Joint Insti- tute for Medical Research — BUDGET: £74,000
	ROLE: Within this project, Alessia Visconti is responsible for the <i>in silico</i> characterisation and validation, using bioinformatics models, of microbes associated with IgA nephropathy and/or IgA glycosylation profiles.
2016-2018	TITLE: "A high-resolution map of copy number and structural variation in Qatari genomes and their contribution to quantitative traits and disease"
	FUNDER: Qatar Foundation – BUDGET: £160,521
	ROLE: Within this project, Alessia Visconti developed (a) an approach for the storage of

genomic data taking advantage of the ROOT library, and (b) an ensemble approach for the identification of structural variation. She also conducted the bioinformatics analysis for [J35] and supervised a research associate.

PROJECTS TO WHICH ALESSIA VISCONTI PARTICIPATES AS RESEARCHER

Alessia Visconti has a prominent role in all projects, as shown by the number of publications in which she is first/last author.

2021-2022	TITLE: "Predicting Response to Immunotherapy for Melanoma with gut Microbiome and metabolomics - The PRIMM Study"
	FUNDER: Seerave Foundation
	ROLE: Alessia Visconti led the bioinformatics analysis for the identification of glyco- markers [J46] (first author) and collaborated to the analysis of proteomic data [J43]
	aiming at finding novel biomarkers of response and survival to identify those patients with melanoma who are most likely to benefit from immune checkpoint inhibitors.
2019-2021	TITLE: "Dissecting the mechanisms of immune-mediated inflammation: a systems- immunology approach"
	FUNDER: MRC Medical Research Council
	ROLE: Alessia Visconti performed bioinformatics analyses aiming at (a) the reverse engineering of immune cell co-expression networks and their involvement in a set of autoimmune diseases, and (b) the identification of genetic variations and microbes/metabolites
	responsible for the development of such diseases.

2016-2018 TITLE: "Gut microbiome modulation of fasting glucose homeostasis and postprandial glycaemic response in TwinsUK and PREDICT: towards personalised diet for healthy aging"

FUNDER: Chronic Disease Research Foundation

ROLE: Alessia Visconti (a) developed a tool for the analysis of metagenomic data which ensures the reproducibility of the scientific results [J17] (first author), (b) performed the bioinformatics analysis of metagenomics and metabolomics data [J22, J31] (co-first author in the first study), (c) collaborated to further studies ([J48, J49, J50], or under revision), and (d) supervised a PhD student for the work described in [J44] (co-senior author).

- 2014-2016 TITLE: "An integrative genomics approach for non-invasive diagnostic biomarkers discovery in IgA nephropathy"
 FUNDER: MRC Medical Research Council
 ROLE: Alessia Visconti applied statistical and bioinformatics approaches for studying the role of IgA and its glycosylation profiles in the development of IgA nephropathy, as described in [J10, J38] (co-first author in both).
- 2013-2015 TITLE: "Senescence and melanoma An integrative systems biology approach to characterise the link between reduced biological senescence and melanoma susceptibility" FUNDER: British Skin Foundation ROLE: Alessia Visconti applied statistical and bioinformatics approaches for studying melanoma, melanoma risk phenotypes, and their connection with ageing, as described in [J11, J15, J16, J19, J21, J24, J29] (first author in three manuscripts).
- 2013-2018 TITLE: "Genomic analysis of Type 2 Diabetes in Qatar, towards diabetes personalized medicine"

FUNDER: Qatar Foundation

ROLE: Alessia Visconti (a) developed and implemented an approach for the population and pedigree association testing for quantitative data [J13] (first author), and (b) conducted the bioinformatics analysis for [J8] and [J14].

2012-2013 TITLE: "LIMPET – Isotropic And Anisotropic Lipophilicity To Model Permeability Of New Therapeutic Peptides"

FUNDER: Compagnia di San Paolo

ROLE: Alessia Visconti (a) evaluated the ability of some combinations of descriptors/algorithms to find the best model to predict the lipophilicity of small peptides [J6] (first author), and (b) performed the bioinformatics analyses to measure the ability of more than 200 compounds of acting as hydrogen bond donors [J4].

- 2007-2011 TITLE: "BioBITs Developing white and green biotechnologies by converging platforms from biology and information technology towards metagenomics" FUNDER: Regione Piemonte ROLE: Alessia Visconti contributed to the development of a modular framework for the analysis of metagenomics sequences, and was responsible for the co-clustering module [J1].
- 2004-2009 TITLE: "Realizzazione di modelli informatici per la valorizzazione della qualità e la tracciabilità delle produzioni in specie da frutto coltivate in Piemonte"
 FUNDER: Regione Piemonte
 ROLE: Alessia Visconti contributed to the development of computational approaches for the classification and traceability of fruits produced in Piemonte.

PROJECTS WITH INDUSTRIAL PARTNERS TO WHICH ALESSIA VISCONTI PARTICIPATES

- 2020 2022 PARTNER: ZOE Ltd https://health-study.joinzoe.com/) ROLE: Alessia Visconti (a) developed the analysis pipeline for the daily data provided by more than four million users [J40], (b) led two studies investigating skin manifestations of SARS-CoV-2 [J33, J42], and (c) performed the bioinformatics analysis for several other studies [J26, J28, J30, J32, J34, J37].
- 2019 2020 PARTNER: Sanofi (https://www.sanofi.com) ("Sanofi iAwards Europe 2019") ROLE: Alessia Visconti carried out part of the bioinformatics analyses and supervised a research associate
- 2018 2020 PARTNER: Danone Nutricia Research (https://www.nutriciaresearch.com) ROLE: Alessia Visconti performed the metagenomic data analyses.

REVIEW ACTIVITY

EDITORIAL BOARD
MEMBERSHIPSince September 2019, Alessia Visconti is Review Editor of Human and Medical Genomics
(speciality section of Frontiers in Genetics).REVIEWERAlessia Visconti serves as a reviewer for the following international journals: Bioinformat-
ics, BioData Mining, BMC Cancer, BMC Nutrition, BMC Supplements, Computational
and Structural Biotechnology Journal, Communications Biology, European Journal of
Human Genetics, Frontiers in Cellular and Infection Microbiology, GigaScience, Journal
of the European Academy of Dermatology and Venereology, Knowledge and Information
Systems (KAIS), - Machine Learning, PLoS Computational Biology, PLoS One

INVITED TALKS

17 Jun 2019	<i>"Reproducible shotgun metagenomic analysis with Nextflow and containers"</i> at the 1st London Bioinformatics Frontiers Frontiers conference, London, UK
23 Nov 2018	"Nextflow on the go" at the 2nd Nextflow workshop, Barcelona, Spain
6 Dec 2017	"YAMP: a framework enabling reproducibility in metagenomics research" at the Compu- tational Biology seminar series, Francis Crick Institute, London, UK
13 Dec 2017	"My reproducible adventure in (meta)genomics (A tale of workflow development, research reproducibility & open science)" at the Researc/hers code seminar series, London, UK
15 Sep 2017	"Simplifying shotgun metagenomics analysis with Nextflow" at the 1st Nextflow work- shop, Barcelona, Spain
7 Jul 2012	<i>"Knowledge-driven Co-clustering of Gene Expression Data"</i> at the Center for Biological Sequence Analysis, Technical University of Denmark, Lyngby, Denmark
8 May 2009	"Using a priori knowledge for the reverse engineering of gene regulatory networks" at the Computer Laboratory, Cambridge University, Cambridge, UK

DEVELOPMENT OF RESEARCH SOFTWARE

The software is available at https://github.com/alesssia or upon request.

AID-ISA	extracts biologically relevant biclusters from microarray gene expression data by leverag- ing additional knowledge (described in [J5])
CDoT	is a novel exact algorithm for answering Maximum a Posteriori queries on tree-structured Probabilistic Graphical Models (described in [P5] and developed in collaboration with
FAMCNV (v2.0)	Roberto Esposito) enables genome-wide association of copy number variants with quantitative phenotypes in families (developed in collaboration with Mario Falchi)
GOCLUST	performs a co-clustering of microarray gene expression data using Gene Ontology-derived constraints (described in [P2])
MotifsLinker	associates clusters of proteins with their frequent motifs
PicNic	extracts topologies and discovers patterns in sets of chemical compound
PopPAnTe	enables pairwise association testing in related samples (described in [J13])
RGO	is a reorganization of the Gene Ontology emphasising regulative information and provid- ing better structure for gene functional analysis (described in [J2])
SPOT	performs an exhaustive search of frequent motifs in sets of biological sequences (described in [T2])
YAMP	allows processing raw metagenomic sequencing data up to the functional annotation (described in [J17])

Jul $2017 - \text{present}$	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 Re-
	searc[her] Research community. These groups aim at promoting gender di-
	versity in the R and STEM community via meetups, mentorship and global
	collaboration. With Researc[her], Alessia Visconti gave speeches on repro-
	ducibility, and workflow development

TEACHING ACTIVITY

In English

To postgraduate students, postdoctoral researchers, and PIs

- 2019 PRESENT Instructor for several programming and data analysis workshops:
 - The Unix Shell
 - Version Control and collaboration with Git/GitHub
 - Python Programming
 - R Programming
 - Introduction to Working with Data
 - OpenRefine

These workshops are designed for PhD students and early career researchers but are open to researchers at every level, including PIs, are offered regularly (roughly twice a year), and have always been evaluated as "excellent" or "exceptional" by the attendees (details on some of the workshops can be found at https://kcl-carpentries.github.io/).

The workshops are centred around the idea of "live coding", where attendees code along with the instructors thus getting useful hands-on experience and improving their ability to explore the topics on their own.

- 2016 2023 **Co-organiser** of the Regulatory Genomics journal club at the Department of Twins Research & Genetic Epidemiology, King's College London. The journal club discusses papers on the latest achievements and methods in the fields of regulation of gene expression, epigenetics, splicing, evolution, and related topics.
- 2021- 2022 Instructor for The Unix Shell and the Version Control and collaboration with Git/GitHub workshops for students and researchers of the MRC Gambia and the AKU Nairobi Research Centres. Both workshops were rated as "exceptional" by the attendees. Instructor for the Metagenomics Data Analysis: Investigating the invisible world of microbes (details at https://alesssia.github.io/metagenomic_workshop/). The workshop, which included frontal lessons and hands-on sessions, was designed for researchers at every level (including but not limited to PhD students, post-doctoral researchers and PIs) without any previous knowledge of the topics and tools presented.

To postgraduate students

- 2013 2014 **Teaching assistant** for the *"Human Molecular Genetics"* MSc Department of Genomics of Common Diseases, Imperial College London. Alessia Visconti was offering support during the practical sessions on R programming as well as one-to-one meetings with students attending the following courses:
 - The Unix Shell
 - R Programming
 - Exploratory Data Analysis and Probability
 - Quantitative genetics
 - Next Generation Sequencing Data Analysis.

Outside academia

2021-2022 **Instructor** for a Version Control and collaboration with Git/GitHub workshop at the UK Health Security Agency (UKHSA)

IN ITALIAN

To medical residents

2023- 2024 Lecturer for *Introduction to Statistics* short course for medical residents at University of Turin.

To undergraduate students

2013- 2014 Lecturer for the "Data analysis" course, Department of Biological Science, University of Turin. Alessia Visconti was offering support during the practical sessions on R programming as well as one-to-one meetings with students.
 Lecturer for the "Operating System" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions on R

sity of Turin. Alessia Visconti was offering support during the practical sessions on R programming as well as one-to-one meetings with students.

- 2012 2013 Lecturer for the "Operating System and Networking" course, Interfaculty School of Strategic Studies, University of Turin. Alessia Visconti was the sole responsible for the practical sessions covering the basis of GNU/Linux, the Unix shell, and process management. She designed the final project, *i.e.*, the development of a basic client/server application in C (details at https://alesssia.github.io/sistemi_elab_info_I_2012_13) Lecturer for the "Operating System" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions as well as one-to-one meetings with students.
- 2011- 2012 Lecturer for the "Database" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions as well as one-to-one meetings with students.
- 2010 2011 Lecturer for the "Database" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions as well as one-to-one meetings with students.
 Lecturer for the "Formal Language" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions as well

as one-to-one meetings with students. Lecturer for the "Statistics and data mining with SAS" course, Department of Mathematics, University of Turin. Alessia Visconti prepared slides and recorded 3 hours of

ematics, University of Turin. Alessia Visconti prepared slides and recorded 3 hours of video lessons on SAS Enterprise Miner. She also prepared a self-evaluation questionnaire for the students.

- 2009 2010 **Lecturer** for the "Computer Science" course, Department of Letters and Philosophy, University of Turin. Alessia Visconti was the sole responsible for the practical sessions covering the MS Office suite and the students' evaluation (details at https://alesssia. github.io/lab_lettere_2009_10/).
- 2006 2007 **Teaching assistant** for the "*Program Languages JAVA*" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions.
- 2005 2006 **Teaching assistant** for the "*Program Languages JAVA*" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions,
- 2004 2005 **Teaching assistant** for the "*Program Languages C*" course, Department of Computer Science, University of Turin. Alessia Visconti was offering support during the practical sessions.

SUPERVISION ACTIVITY

Thanks to her multidisciplinary experience, which combines a BSc, MSc, and PhD in computer science with more than 10 years of research activity in the biomedical field, Alessia Visconti has been able to supervise students with different backgrounds (*e.g.*, computer science, bioinformatics, molecular biology, medicine, neurobiology, and engineering) offering a stimulating and engaging environment.

BSc & MSc Theses

Co-supervisor of Ms Darvina Magandran's MSc project in Microbiome in Health and
Disease at King's College London.
Co-supervisor of Ms Raphaela Joos's MSc project in <i>Microbiome in Health and Disease</i>
at King's College London.
Co-supervisor of Ms Petra Blackburn's MSc project in <i>Microbiome in Health and Disease</i> at King's College London.
Co-supervisor of Ms Natalie Falshaw's MSc project in <i>Microbiome in Health and Dis</i> -
ease at King's College London.
Co-supervisor of Ms Xinyu Huang's MSc project in Pharmacology at King's College
London.
Co-supervisor of Ms Miriam Margari's MSc project in Genomic Medicine at Imperial
College London.
Co-supervisor of Dr Robin Mesnage's MSc project in Bioinformatics at Birkbeck University of London
Co-supervisor of Mr George Powell's MSc project in Human Molecular Genetics at
Imperial College London.
Co-supervisor of Mr Marco Gallizio's bachelor thesis in Computer Science at the University of Turin

Post-master students

2019 - 2020 **Co-supervisor** of Mr Simon Couvreur in his PhD rotation project at King's College London.

Co-supervisor of Ms Helen King in her PhD rotation project at King's College London.

PhD students

- 2021 PRESENT **Co-supervisor** of Mr Roger Compte Boixader in his PhD project at King's College London.
- 2019 PRESENT **Co-supervisor** (unofficial) of Ms Xinyuan Zhang in her PhD project at King's College London (Publications: [J44]).
 - 2020 2022 Assistance with the supervision of Ms Karla Lee in her PhD project at King's College London (supervision limited at the multi-*omics* data analyses; publications: [J43, J46]).
 - 2016 2019 **Co-supervisor** (unofficial) of Mr Niccolò Rossi during his research visit at King's College London (Publications [J35]).
 - 2018 2020 **Supervisor** of Ms Giulia Piaggeschi during her research visit at King's College London (Publications: [J36]).
 - 2014 2020 **Co-supervisor** (unofficial) of Ms Marianna Sanna in her PhD project at King's College London and her research activity at Imperial College London (Publications: [J11, J19, J21, J24, J29]).
 - 2015 2019 Assistance with the supervision of Mr Taghi Aliyev during his PhD at the CERN OpenLab (supervision limited to the biomedical part of the project).

Postdoctoral researchers

- 2019 2023 Assistance with the supervision of Mr Niccolò Rossi during his postdoc at King's College London (already supervised during the PhD; publications [J43]).
- 2016 2018 Assistance with the supervision of Dr Harriet Cullen during her research fellowship at King's College London (Publications: [J20])

Interns

SUMMER 2019 Supervisor of Mr Yuhao Lin's summer project as part of the King's Undergraduate Research Fellowships (KURF) ${\bf Supervisor}$ of Ms Olivia Castellini Pérez's summer internship as part of the Erasmus+ program

- SUMMER 2018 **Supervisor** of Ms Lechun Huo's summer project as part of the King's Undergraduate Research Fellowships (KURF).
- SUMMER 2017 **Supervisor** of Ms Yuri Nemoto's summer project as part of the King's Undergraduate Research Fellowships (KURF).

Co-supervisor of Ms Fudi Wang's research visit at King's College London

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