# Emidio Capriotti PhD

Name: Emidio Capriotti Nationality: Italian

Visiting

May 2009

Jun 2012 – Jul 2012

Leuven (Belgium)

Date of birth: February, 1973 Place of birth: Roma, Italy

Languages: Italian, English, Spanish

Positions			
Oct 2019	Associate Professor: Department of Pharmacy and Biotechnology (FaBiT). University of Bologna, Bologna, Italy		
2016-2019	Senior Assistant Professor (RTD type B): Department of Pharmacy and Biotechnology (FaBiT) and Department of Biological, Geological, and Environmental Sciences (BiGeA). University of Bologna, Bologna, Italy.		
2015-2016	Junior Group Leader: Institute of Mathematical Modeling of Biological Systems, University of Düsseldorf, Düsseldorf, Germany		
2012-2015	Assistant Professor: Division of Informatics, Department of Pathology, University of Alabama at Birmingham (UAB), Birmingham (AL), USA.		
2011-2012	Marie-Curie IOF: Contracted Researcher at the Department of Mathematics and Computer Science, University of Balearic Islands (UIB), Palma de Mallorca, Spain.		
2009-2011	Marie-Curie IOF: Postdoctoral Researcher at the Helix Group, Department of Bioengineering, Stanford University, Stanford (CA), USA.		
2006-2009	Postdoctoral Researcher in the Structural Genomics Group at Department of Bioinformatics and Genetics, Prince Felipe Research Center (CIPF) Valencia, Spain.		
2004-2006	Contract researcher at Department of Biology, University of Bologna, Bologna, Italy.		
2001-2003	Ph.D student in Physical Sciences at University of Bologna, Bologna, Italy.		
Education			
Sep 2004	Master in Bioinformatics (first level) University of Bologna, Bologna (Italy).		
Jun 2004	Ph.D. in Physical Sciences University of Bologna, Bologna (Italy).		
Jul 1999	Laurea (B.S.) Degree in Physical Sciences, score 106/110 University of Bologna, Bologna (Italy).		

Prof. Frederic Rousseau and Prof. Joost Schymkowitz, VIB Switch Laboratory, KU Leuven,

Prof. Francisco Melo group. Department of Molecular Genetics and Microbiology. Pontificia

Universidad Catolica de Chile, Santiago de Chile (Chile).

Jul 2008 - Aug 2008	Prof. Andrej Sali group, Departments of Biopharmaceutical Sciences and Pharmaceutical Chemistry, University of California at San Francisco (UCSF), San Francisco (CA).
Aug 2005 - Nov 2005	Prof. Jeffrey Skolnick group. Center of Excellence in Bioinformatics University of New York at Buffalo, Buffalo (NY)
Awards and Grants	
Jan 2022 - Dec 2023	ELIXIR Commissioned Service on Rare Disease and Federated European Phenome Genome Archive (EGA).
Sep 2019 - Mar 2023	Unit Coordinator: Research Projects of National Relevance (PRIN-201744NR8S) Ministry of Education, University and Research (MIUR). Integrative tools for defining the molecular basis of the diseases: Computational and Experimental methods for Protein Variant Interpretation.
Sep 2018	National Scientific Habilitation (ASN) for Full Professor in Biochemistry (BIO/10)
Sep 2017 – Sep 2018	University of Bologna Cooperation Grant: International Cooperation Project UNIBO-UCSD. Study and development of computational methods for cancer genome interpretation.
Dec 2017	Ministry of Education, Universities and Research: Fondo per il finanziamento delle attività base di ricerca 2017.
Jun 2012 – Jul 2012	Short term EMBO Fellowship for "Computational methods to predict the functional impact of protein variations on alpha-galactosidase and the efficacy of pharmacological chaperone therapy" at KU Leuven, Leuven (Belgium)
Sep 2011 – Aug 2012	Marie-Curie IOF at the Department of Mathematics and Computer Sciences, University of Balearic Islands (UIB).
Sep 2009 – Aug 2011	Marie-Curie IOF at the Helix Group, Department of Bioengineering, Stanford University
Oct 2006 – Aug 2009	Postdoc Research Fellowship. in the Structural Genomics Group at Department of Bioinformatics and Genetics, Prince Felipe Research Center (CIPF) Valencia, Spain
Jul 2008 – Aug 2008	Short-term research fellowship from the Valencian Government (Spain) for the development of a method for RNA structure prediction using MODELLER program. Project supervised by Prof. Andrej Sali, Departments of Biopharmaceutical Sciences and Pharmaceutical Chemistry, University of California, San Francisco (UCSF).
Sep 2004 - Sep 2006	Postdoc Research Fellowship. <i>BioSapiens</i> Network of Excellence, funded by the European Union's VI Framework Programme.
Aug 2005 - Dec 2005	Marco Polo Research Fellowship for "Implementation of new software for protein structure prediction" developed in the Center of Excellence in Bioinformatics University at Buffalo under the supervision of Prof. Jeffrey Skolnick
Jan 2003 - May 2004	National Institute for Biophysics and Biomaterials (INBB) Research Fellowship. Partially supported by MURST (FISR 2002) project <i>Hydrolases from Thermophiles: Structure, Function and Homologous and Heterologous Expression</i> .
Sep 2001 - Sep 2002	SPINNER Consortium (Regione Emilia-Romagna) Research Fellowship for Technology Transfer through a grant to the BioDec project.

# **Teaching activity**

At the University of Bologna, I am member of the committee of the PhD program in Data Science and Computation. I am also coordinator of the International Master in Bioinformatics and teacher of the second module of *Laboratory of Bioinformatics 1*. In the Bachelor in Genomics I am teacher of the course *Proteomes, Interactomes and Biological Networks*. I am responsible for the ERASMUS exchange program with 7 foreigner universities.

The teaching material is available at <a href="https://biofold.org/training.html">https://biofold.org/training.html</a>

From 2015 to 2016, at the University of Düsseldorf I gave 2 one-week modules as part of the master courses *Introduction* to *Molecular Systems Biotechnology* and *Biological Networks*. From 2014 to 2015 at the University of Alabama at Birmingham, in collaboration with Dr. Malay Basu, I was course master in two courses in the Graduate Biomedical Science program at UAB and organizer of the CB2 (Computational Biology and Bioinformatics) Journal Club at UAB. The list of courses given during last few years is reported below.

2016-2022 Module 2: Laboratory of Bioinformatics 1 (66563), International Master in Bioinformatics (60 hours/year).

University of Bologna.

2019-2021 Proteomes, Interactomes and Biological Networks (85306), Bachelor in Genomics (62/year hours).

University of Bologna.

2015-2016 Module: Introduction to Computational Biology and Bioinformatics. Introduction to Molecular Systems

Biotechnology course (M4453). Molecular Systems Biotechnology Master (20 hours). University of

Düsseldorf.

Module: Introduction to Protein-Protein Interaction Network. Biological Networks course (M4424).

2014-2015 GBSC 703-01E Computational Biology and Bioinformatics, Graduate Biomedical Science Program (40

hours). University of Alabama at Birmingham (USA).

2014-2015 GBSC 703-01A - Introduction to Scientific Computing, Graduate Biomedical Science Program (48

hours). University of Alabama at Birmingham (USA).

2013-2014 GBSC 703-01A - Introduction to Scientific Computing, Graduate Biomedical Science Program (30

hours). University of Alabama at Birmingham (USA).

I was also contracted professor for the following courses held by Prof. Rita Casadio at the University of Bologna (Italy):

**2004-2005** Bioinformatics - Degree in Biotechnology (30 hours) **2003-2004** Laboratory of Biophysics II - Degree in Biotechnology

Models for Biological Systems - Degree in Biotechnology (90 hours total)

2002-2003 Structural Biochemistry - Degree in Biotechnology (70 hours)
2001-2002 Laboratory of Biophysics II - Degree in Biotechnology (25 hours)

#### Mentoring activity

At the University of Bologna, I am member of the committee of the PhD program in Data Science and Computation. Currently, I am supervising the research activity of Andrea Cicconardi, research assistant in our group. I am also mentoring the master thesis of Cecilia Foglini, student of the International Master in Bioinformatics at the University of Bologna (Italy). I am co-mentoring the research activity Jaume Sastre Tomàs, PhD student at the Department of Mathematics and Informatics, University of Balearic Islands (Spain). In the past, I directed the thesis of:

Riccardo Ottalevi Title: Predicting the effect of single point mutations on protein-protein interactions

International Master in Bioinformatics at the University of Bologna (Italy).

Bologna 28 February 2022

Tommaso Mosca Title: Impact of protein modeling on the prediction of protein stability change upon mutation.

International Master in Bioinformatics at the University of Bologna (Italy).

Bologna 28 February 2022

Lucrezia Valeriani Title: Predicting the functional impact of single nucleotide variants in rare disease associated

genes.

International Degree in Genomics at the University of Bologna (Italy).

Bologna 11 September 2020

Alessandro Vinceti Title: Development of new tools for predicting the impact of genetic variants in the cancer

genome.

International Master in Bioinformatics at the University of Bologna (Italy).

Bologna, 25 September 2019.

Emina Merdan Title: Characterizing the impact of mutations at functional and network levels in Lung

Adenocarcinoma.

International Master in Bioinformatics at the University of Bologna (Italy).

Bologna, 26 September 2018.

Oronzo Tassiello Title: Computational methods for scoring genomes of Lung Adenocarcinoma

International Master in Bioinformatics at the University of Bologna (Italy).

Bologna, 9 March 2018.

Luigi Chiricosta Title: Detecting cancer causing genes and variants in Colon Adenocarcinoma.

International Master in Bioinformatics at the University of Bologna (Italy).

Bologna, 26 September 2017.

At the University of Alabama at Birmingham, I was supervising the research activity of one postdoc (Dr. Rui Tian) and one master student (Shivani Viradia).

Previously, I collaborated with Prof. Rita Casadio, Dr Mario Compiani and Dr Marc A. Marti-Renom to mentor the research activity of the following students:

Alberto Stizza BS thesis in Physical Sciences, Catholic University of Brescia (Italy)
Daniela Danesi BS thesis in Physical Sciences, Catholic University of Brescia (Italy)

Maria Procopio BS thesis in Physical Sciences, University of Bologna (Italy)
Remo Calabrese PhD thesis in Biotechnology, University of Bologna (Italy)

Giulia Gentile MS thesis in Bioinformatics, CRS4 Bioinformatics Laboratory Cagliari (Italy)

Stefania Bosi PhD student, University La Sapienza, Roma (Italy)
Miquel Oltra Sastre Master thesis, Open University of Catalonia (Spain)

## Reviewer activity

I am faculty member of the Faculty Opinions in the section of Bioinformatics, Biomedical Informatics & Computational Biology / Translational Bioinformatics. I am reviewer for the following journals: Nature Communications, Bioinformatics, Briefings in Bioinformatics, Nucleic Acids Research, The American Journal of Human Genetics, PLOS Computational Biology, Scientific Reports, Cancer Research, Oncotarget, BMC Bioinformatics, PLOS ONE, BMC Genomics, Proteins, Human Mutation, Human Genetics, Human Genomics, Amino Acids, BMC Structural Biology, Database, Current Bioinformatics, Current Protein and Peptide Science, Journal of Bioinformatics and Computational Biology, Neurocomputing, Information Fusion, IEEE/ACM Transactions on Computational Biology and Bioinformatics. I was reviewer of projects for the Medical Research Council of the United Kingdom and for the Austrian Academy of Sciences.

## Other scientific activity

As a member of the International Society of Computational Biology (ISCB). I served as co-chair in the organization of the Personal Genomics session at the Pacific Symposium of Biocomputing (PSB) 2011. Since 2011, in collaboration with Yana Bromberg, Hannah Carter and Antonio Rausell, I organized 10 editions of the VarI-SIG meeting (formerly SNP-SIG) in Vienna (Austria), Long Beach, (California), Berlin (Germany), Boston (Massachusetts), Dublin (Ireland), Orlando (USA), Prague (Czech Republic), Chicago (USA), Basel (Switzerland) and on online conference platform in 2020 and 2021. With Yana Bromberg and Hannah Carter, I was co-editor of 6 special issues on BMC Genomics which published the selected works presented at the VarI-SIG meetings from 2011 to 2018. More information about the VarI-SIG meeting is available at <a href="http://varisig.biofold.org">http://varisig.biofold.org</a>. In 2013, I was member of the Proceedings Papers Committee for the ISMB/ECCB Conference. In 2015 I was member of the Late Breaking Research Committee for the ISMB/ECCB Conference. In 2016 I was co-chair of the Disease track for the ISCB Latin American Conference in Buenos Aires (Argentina). In 2020 I was also member of the proceeding committee of the ISMB Conference for the "Genomic Variation Analysis" track.

In 2010, I was member of the Data Committee in the first edition of the Critical Assessment of Genome Interpretation (CAGI), an international community effort for the assessment of computational methods for genome interpretation (<a href="https://genomeinterpretation.org/">https://genomeinterpretation.org/</a>). Later in the 2018 and 2021 edition of CAGI, I was organizer of the Frataxin, Calmodulin and MAPKs challenges.

Since 2020 I am co-coordinator of the Service Bundle 1 (<a href="https://amp4rd.github.io">https://amp4rd.github.io</a>) of the Rare Disease Community of the ELIXIR leading European infrastructure for life science organizations (<a href="https://elixir-europe.org/">https://elixir-europe.org/</a>).

# Personal research projects

2017-2018 University of Bologna Cooperation Grant: International Cooperation Project UNIBO-UCSD. Study and

development of computational methods for cancer genome interpretation.

2017 Ministry of Education, Universities and Research: Fondo di Finanziamento per le Attività Base di Ricerca

(FFABR).

2009-2012 Marie Curie International Outgoing Fellowships for Career Development, New methods to evaluate the

impact of single point protein mutation on human health.

2012 EMBO Short Term Fellowship, Computational methods to predict the functional impact of protein

variations on alpha-galactosidase and the efficacy of pharmacological chaperone therapy.

2008 Short-term research fellowship from the Valencian Government (Spain), Development of a method for

RNA structure prediction using MODELLER program.

## Participation in research projects

Sep 2019 -	Integrative tools for defining the molecular basis of the diseases: computational and experimental methods for protein variant interpretation (MIUR-PRIN-201744NR8S) PI: Piero Fariselli Role: Unit Coordinator
2018-2020	Role of VpreB in immunoglobulin antigen binding site selection (NIH 1R21Al134027-01A1) PI: Harry Schroeder. Role: Consultant.
2016-2018	Bioinformatics applications in phylogenetics, metagenomics, systems biology and cancer genomics. (MEC: DPI2015-67082-P)
	PI: Francesc Andreu Rossello'. Role: Investigator.
2015-2017	The pre-BCR CDR-H3 sensing site and H chain selection (NIH: 1R21AI117703-01A1).
	PI: Harry Schroeder. Role: Co-Investigator.
2014-2015	Mechanisms of glucose mediated cardiac mitochondrial dysfunction (NIH: 3R00HL111322-04S1).
	PI: Adam Wende. Role: Co-Investigator.
2006-2008	Marie Curie Reintegration Grant - European Union, PI: Marc A. Marti-Renom.
	RNA Comparative Modeling.
2004-2006	VI Framework Programme - European Union, PI: Janet Thornton.
	Biosapiens Network of Excellence, A European Virtual Institute for Genome Annotation.
2003-2004	FISR2002 - Italian Ministry of Education Research and University (MIUR), PI: Rita Casadio.
	Hydrolases from Thermophiles: Structure, Function and Homologous and Heterologous Expression.

#### **Research Interests**

- Analysis and interpretation of cancer genome.
- Genome interpretation and prediction of disease-related protein mutations.
- Machine learning approaches in molecular biology.
- Protein-protein interactions.
- RNA structure comparison and prediction.
- Protein folding kinetics.
- Prediction of protein stability changes upon mutation.
- Protein structural prediction by threading methods and building by homology.
- Molecular dynamics of protein systems.

## **Developed Web Servers, Tools and Databases**

- ContrastRank: probabilistic method for cancer gene prioritization and cancer sample classification.
  - WEB: http://snps.biofold.org/contrastrank
- **DrCancer:** predictor of cancer causing non-synonymous single nucleotide polymorphisms.
  - WEB: http://snps.biofold.org/drcancer
- DDGun: untrained method for predicting of protein stability upon single and multiple site mutations.
  - WEB: <a href="http://folding.biofold.org/ddgun">http://folding.biofold.org/ddgun</a>
- Fido-SNP: predicts the impact of genetic variants in coding and non-coding regions of the dog genome.
  - WEB: http://snps.biofold.org/fido-snp
- I-Mutant1.0: Neural Network based method to predict the sign of free energy change of proteins upon single point
  mutation.
  - WEB: http://gpcr2.biocomp.unibo.it/cgi/predictors/I-Mutant/I-Mutant.cgi
- I-Mutant2.0: Support Vector Machine based method to predict the sign and the value of free energy change of proteins upon single point mutation.
  - WEB: http://folding.biofold.org/i-mutant
- K-Fold: Support Vector Machine based method to predict the mechanism and rate of protein folding kinetic.
   WEB: <a href="http://folding.biofold.org/k-fold">http://folding.biofold.org/k-fold</a>
- Meta-SNP: Meta-predictor of disease-causing variants that uses PANTHER, PhD-SNP, SIFT and SNAP.
   WEB: <a href="http://snps.biofold.org/meta-snp">http://snps.biofold.org/meta-snp</a>
- Omidios: Omidios, a database of pre-calculated likely impact of a Single Nucleotide Polymorphism in the human genome.
  - WEB: http://sqt.cnaq.cat/services/Omidios/
- PhD-SNP: Support Vector Machine based Method to discriminate between disease-related and neutral mutations in proteins.
  - WEB: http://snps.biofold.org/phd-snp

 PhD-SNP9: A gradient boosting-based method for predicting the impact of genetic variants in coding and non-coding regions.

WEB: http://snps.biofold.org/phd-snpg

• SARA: a tool for Structural Alignment of Ribonucleic Acids.

WEB: <a href="http://structure.biofold.org/sara">http://structure.biofold.org/sara</a>

SARA-Coffee: tool for RNA multiple structural alignment obtained merging SARA and T-Coffee.
 WEB: <a href="http://www.tcoffee.org/Projects/saracoffee/">http://www.tcoffee.org/Projects/saracoffee/</a>

• ThermoScan: tool for scanning biomedical publications to retrieve protein thermodynamic data.

WEB: https://folding.biofold.org/thermoscan

WebRASP: statistical potential for scoring the quality of RNA three-dimensional structure.

WEB: http://melolab.org/webrasp

• WS-SNPs&GO: predictor of human disease related mutations in proteins with functional annotation.

WEB: http://snps.biofold.org/snps-and-go

## International conferences meetings and schools

- Moscow Conference on Computational Molecular Biology (MCCMB), Moscow (Russian Federation), 30 July 2 August 2021.
- XXIX Intelligent Systems for Molecular Biology meeting and XX European Conference on Computational Biology (ECCB). Virtual Conference, 35-30 July 2021.
- XXVIII Intelligent Systems for Molecular Biology meeting (ISMB). Virtual Conference, 13-16 July 2020.
- Bologna Winter School 2020: What can we learn from protein structure? Bologna (Italy), 17-21 February 2020.
- Bologna Winter School 2019: Data Science for Bioinformatics. Bologna (Italy), 18-22 February 2019.
- XXVI Intelligent Systems for Molecular Biology meeting (ISMB), Chicago, (USA), 6-10 July 2018.
- V Critical Assessment of Genome Interpretation (CAGI). Chicago, (USA), 5-7 July 2018.
- The molecular basis of diseases: Can we infer phenotypes from protein variant analysis? FEBS Advanced Course.
   23-25 May 2018, Bologna, Italy
- Bologna Winter School 2018: Big Data and Bioinformatics. Bologna (Italy), 12-16 February 2018.
- XXV Intelligent Systems for Molecular Biology meeting (ISMB) and XVI European Conference on Computational Biology (ECCB), Prague, (Czech Republic), 21-25 July 2017
- Bologna Winter School 2017: Revisiting Bioinformatics Foundations. Bologna (Italy), 13-17 February 2017.
- XIV Intelligent Systems for Molecular Biology meeting (ISMB), Orlando, FL (USA), 8-12 July 2016
- Bologna Winter School 2016: In Silico Markers for Precision Medicine. Bologna (Italy) 22-26 February 2016.
- EMBL Conference on Cancer Genomics, Heidelberg (Germany), 1-4 November 2015.
- XXIII Intelligent Systems for Molecular Biology meeting (ISMB) and XIV European Conference on Computational Biology (ECCB), Dublin (Ireland), 12-14 July 2015
- UAB NHGRI IV Short Course on Next-Generation Sequencing; Technology and Statistical Methods. Birmingham (AL), 15-18 December 2014.
- XIII European Conference on Computational Biology (ECCB), Strasbourg (France), 7-10 September 2014.
- XXII Intelligent Systems for Molecular Biology meeting (ISMB), Boston, Massachusetts (USA), 13-15 July 2014.
- VarI-SIG meeting. Identification and annotation of genetic variants in the context of structure, function, and disease.
   Boston, Massachusetts (USA), 12 July 2014.
- XXI Intelligent Systems for Molecular Biology meeting (ISMB) and XII European Conference on Computational Biology (ECCB), Berlin (Germany), 21-23 July 2013
- SNP-SIG meeting. Identification and annotation of SNPs in the context of structure, function, and disease. Berlin (Germany), 19 July 2013
- Critical Assessment of Genome Interpretation (CAGI). Berlin (Germany), 17-18 July 2013.
- ESHG Course in Next Generation Sequencing, Bertinoro di Romagna (Italy), 17-20 May 2013
- Summit on Translational Bioinformatics (TBI), San Francisco, California (USA), 18-20 March 2013.
- XX Intelligent Systems for Molecular Biology meeting (ISMB), Long Beach, California (USA), 15-17 July 2012.
- SNP-SIG meeting. Identification and annotation of SNPs in the context of structure, function, and disease. Long Beach, California (USA), 14 July 2012.
- Bologna Winter School 2012 Proteins and their variants: structure and function prediction. Bologna (Italy) 13-17
  February 2012.
- XIX Intelligent Systems for Molecular Biology meeting (ISMB) and X European Conference on Computational Biology (ECCB), Vienna (Austria), 17-19 July 2011.
- SNP-SIG meeting. Identification and annotation of SNPs in the context of structure, function, and disease. Vienna (Austria), 15 July 2011.
- EMBO Young Scientist Forum. International Institute of Molecular and Cell Biology (IIMCB), Warsaw (Poland), June 30<sup>th</sup> July 1<sup>st</sup> 2011.
- Pacific Symposium on Biocomputing (PSB) 2011. Big Islands, Hawaii January 3-7 2011.
- Critical Assessment of Genome Interpretation (CAGI). University of California at Berkeley. Berkeley, California (USA), 10 December 2010.

- Biomedical Computation at Stanford (BCATS). Stanford University. Palo Alto, California (USA), 6 November 2010.
- Exploring the functional consequences of genomic variation (HGVS meeting), Washington DC (USA), 2 November 2010
- II Workshop on Annotation, Interpretation and Management of Mutations (AIMM) and IX European Conference on Computational Biology (ECCB), Ghent (Belgium), 26-29 September 2010.
- 4<sup>th</sup> Comprehensive Cancer Research Training Program (CCRTP) at Stanford University, Palo Alto California (USA), 13-17 September 2010.
- 9th International Conference on Computational Systems Bioinformatics (CSB). Stanford, Palo Alto, California (USA), 16-18 August 2010.
- XVIII Intelligent Systems for Molecular Biology meeting (ISMB), Boston (USA), 11-13 July 2010.
- Biomedical Computation at Stanford (BCATS). Stanford University. Palo Alto, California (USA), 7 November, 2009.
- Lipari International Summer School on BioInformatics and Computational Biology. RNAs: structure, function and therapy. Lipari (ME) 13-20 June, 2009.
- VII European Conference on Computational Biology (ECCB), Cagliari (Italy), 22-26 September 2008.
- Workshop on Applications of Protein Models in Biomedical Research, University of California San Francisco (UCSF), San Francisco (CA) 11-12 July, 2008
- III Course on Molecular Evolution, Phylogenetics and Phylogenomics, Valencia (Spain) 12-16 May 2008.
- Non-Coding RNAs: Computational Challenges and Applications. Antalya (Turkey) 28-30 April 2008.
- XV Intelligent Systems for Molecular Biology meeting (ISMB) and VI European Conference on Computational Biology (ECCB), Vienna (Austria), 21-25 July 2007.
- ISMB 3DSig Satellite Meeting Structural Bioinformatics and Computational Biophysics, Vienna (Austria), 19-20 July 2007.
- EMBO Workshop: Viral RNA: Structure Function and Targeting. EMBL Heidelberg (Germany) 5-7 March 2007.
- Bologna Winter School 2006. Applied Bioinformatics: The test case of Human Genome. Bologna (Italy), 13-17 February, 2006.
- Bologna Winter School 2005: *How Complex is Functional Genomics?* Bologna (Italy), 13-19 February 2005. XII Intelligent Systems for Molecular Biology (ISMB) and III European Conference on Computational Biology meeting (ECCB), Glasgow (Scotland) 31 July 4 August 2004.
- Bologna Winter School 2004: The State of the Art of Protein-Protein Interaction Networks. The role of the "in silico" approach, Bologna (Italy) 8-14 February 2004.
- Meeting Galileo Project. Marseille (France) 27-28 June 2003.
- Bologna Winter School 2003: Hot Topics in Structural Genomics Bologna (Italy) 9-15 February 2003.
- Bologna Winter School 2002: Predicting 3D Structure of Difficult Proteins. Bologna (Italy) 3-9 February 2002.
- Bologna Winter School 2001: In Silico Biomolecular Recognition. Bologna (Italy) 4-10 February 2001.
- Bologna Summer School: Biotechnology Protein Sequence Analysis in the Genomic Era. Bologna (Italy) 10-16 October 1999.

#### National conferences meetings and schools

- SIB 2019, 60th Congress, Lecce 18-20 September 2019.
- Computational and Translational Methods for Cancer Genomics, Bologna 29 May, 2018.
- German Conference on Bioinformatics (GCB) 2015, Dortmund (Germany), 27-30 September 2015.
- UAB Comprehensive Cancer Center, 15th Annual Research Retreat and Research Competition, Birmingham, Alabama (USA), October 29 2012.
- VIII Jornadas de Bioinformatica. Valencia (Spain), 13-15 February 2008.
- VI Meeting on Nucleic Acids and Nucleotides (RANN07), Valencia (Spain), 22-23 November 2007.
- VII Jornadas de Bioinformatica. Zaragoza (Spain), 20-22 November 2006.
- Bioinformatics Italian Society (BITS) Annual Meeting 2006. Bologna (Italy), 28-29 April 2006.
- Bioinformatics Italian Society (BITS) Annual Meeting 2004 Padova (Italy), 26-27 March 2004.
- Workshop Staminal Cells: *Properties and Perspectives*. Bressanone (Italy), 11-13 September 2003.
- XI National School of Biophysics: Biophysics of the Cell. Bressanone (Italy), 8-10 September 2003.
- IX National School of Biophysics: *Biophysics and Biomaterials*. Bressanone (Italy), 3-5 September 2001.
- XXXI National Congress of Physical Chemistry, Padova (Italy), 19-23 June 2001.

# Invited talks

07 Aug 2006	Centro de Investigacion Principe Felipe (CIPF), Valencia (Spain)
23 Oct 2007	Centro Nacional de Investigaciones Oncológicas (CNIO), Madrid (Spain)
06 May 2008	Département d'Informatique, Université Libre de Bruxelles (ULB), Bruxelles (Belgium)
23 Apr 2009	Departament de Ciències Matemàtiques i Informàtica, Universitat de les Illes Balears (UIB), Palma de
	Mallorca (Spain)
27 May 2010	Buck Institute, Novato (California, USA)

02 Jul 2010	Luxembourg Centre for System Biomedicine, Luxemburg University, Luxembourg
20 Sep 2010	Department of Genetics and Bioengineering, Yeditepe University, Istanbul (Turkey)
25 Jan 2011	Department of Medicinal Chemistry and Molecular Pharmacology, Purdue University, Lafayette (Indiana,
	USA).
18 Feb 2011	Lawrence Berkeley National Laboratory, Berkeley (California, USA).
21 Mar 2011	Department of Computer Sciences, Wayne State University, Detroit (Michigan, USA).
06 Apr 2011	J. Craig Venter Institute, San Diego (California, USA)
22 Apr 2011	Department of Bioengineering, University of Texas at Dallas (Texas, USA)
17 May 2011	Department of Pathology, University of Alabama at Birmingham (Alabama, USA)
26 May 2011	Department of Computer Science, Université Pierre et Marie Curie, Paris (France)
16 Jun 2011	Instituto Gulbenkian de Ciencia, Oeiras (Portugal)
29 Jun 2011	International Institute of Molecular and Cell Biology, Warsaw (Poland)
04 Oct 2011	Institut de Cancerologie Gustave Roussy, Villejuif (France)
06 Oct 2011	Karlsruhe Institute of Technology, Karlsruhe (Germany)
12 Jun 2012	Switch Lab, KU Leuven, Leuven (Belgium)
30 Apr 2013	Biomedical Informatics Day, Adelaide (SA, Australia)
16 Jul 2013	Institute for Medical and Human Genetics, Charité University, Berlin (Germany)
07 Oct 2013	CCNR, Northeastern University, Boston (Massachusetts, USA)
10 Dec 2013	Macromolecular Biochemistry Research Center (CRBM), CNRS, Montpellier (France)
14 Dec 2013	Computational Biology Institute (IBC), CNRS, Montpellier (France)
12 Feb 2014	Institute of Genetics and Molecular and Cell Biology (IGBMC), Strasbourg (France)
22 May 2014	Montpellier Cancer Research Institute, University of Montpellier, Montpellier (France)
06 Aug 2014	Izmir Biomedicine and Genome Center (IBG), Izmir (Turkey)
18 Sep 2014	Technical University of Munich (TUM), Munich (Germany)
26 Sep 2014	Pontificia Universidad Catolica de Chile, Santiago de Chile (Chile)
12 Sep 2016	Adam Mickiewicz University (AMU), Poznan (Poland)
17 Jul 2019	University of Lisboa, Lisboa (Portugal)
02 Aug 2021	Moscow Conference on Computational Molecular Biology, Moscow (Russian Federation)

# **Invited lectures**

01 Sep 2011

17 Feb 2012	Bologna Winter School 2012, Bologna (Italy)
08 Feb 2013	Anatomic Pathology Didactic Conference. University of Alabama at Birmingham (Alabama, USA)
15 Feb 2013	Genetics and Genomics Seminar Series. University of Alabama at Birmingham (Alabama, USA)
04 Mar 2013	GBM-722 Bioinformatics Course. University of Alabama at Birmingham (Alabama, USA)
07 Mar 2013	Biotechnology Professional Master. University of Alabama at Birmingham (Alabama, USA)
22 Apr 2013	GBS Structural Biology Course. University of Alabama at Birmingham (Alabama, USA)
14 May 2013	Laboratory Medicine Seminar. University of Alabama at Birmingham (Alabama, USA)
17 May 2013	European School of Genetic Medicine. European School of Genetic Medicine, Bertinoro (Forli', Italy)
02 Feb 2014	GBM-722 Bioinformatics Course. University of Alabama at Birmingham (Alabama, USA)
06 Mar 2014	Biotechnology Professional Master. University of Alabama at Birmingham (Alabama, USA)
03 Jul 2014	GBS-758 New Perspectives in Cardiovascular Biology. University of Alabama at Birmingham (Alabama,
	USA)
08 Oct 2014	Biotechnology Professional Master. University of Alabama at Birmingham (Alabama, USA)
18 Dec 2014	UAB NHGRI IV Short Course on Next-Generation Sequencing; Technology and Statistical Methods.
	University of Alabama at Birmingham (Alabama, USA)
25/26 Feb 2016	Bologna Winter School 2016, University of Bologna (Italy).
09 Jun 2017	PhD Program in Biochemistry, "La Sapienza" University, Roma (Italy)
06 Sep 2017	Special course on NGS data analysis. CRO National Cancer Institute, Aviano (Italy)
17 Jan 2019	Winter School University of Verona, Canazei (Italy)

Statistics and Genomics Seminar, University of California, Berkeley (California, USA)

# **Publications**

I published 52 research articles and 11 reviews in international peer-reviewed journals with impact factor. I also published 18 between book chapters (10) and meeting reports (8). Using Google Scholar my papers received more than 6,200 citations corresponding to an h-index of 30. According to Scopus, my papers received more than 4,300 citations corresponding to an h-index of 28. Using Web of Science my articles received more than 3,900 citations corresponding to h-index of 26.

Google Scholar: <a href="http://bit.do/ecapriotti-scholar">http://bit.do/ecapriotti-scholar</a>

Scopus: http://bit.do/ecapriotti-scopus
ORCID: http://bit.do/ecapriotti-orcid

ResearcherID: <a href="http://bit.do/ecapriotti-wos">http://bit.do/ecapriotti-wos</a>
ResearchGate: <a href="http://bit.do/ecapriotti-rgate">http://bit.do/ecapriotti-rgate</a>

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*Capriotti E, Fariselli P, Calabrese R, Casadio R* (2005). Predicting protein stability changes from sequences using support vector machines. **Bioinformatics**, 21 Suppl 2:ii54-ii58.

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Grandi F, Sandal M, Guarguaglini G, Capriotti E, Casadio R, Samorì B (2006). Hierarchical mechanochemical switches in Angiostatin. ChemBiochem, 7; 1774-1782.

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Pancotti C, Benevenuta S, Birolo G, Repetto V, Alberini V, Sanavia T, Capriotti E\*, Fariselli P\* (2022). Predicting protein stability changes upon single-point mutation: a thorough comparison of the available tools on a new dataset. Briefings in Bioinformatics. DOI:10.1093/bib/bbab555

Montanucci L<sup>†</sup>, Capriotti E<sup>†</sup>, Birolo G, Benevenuta S, Pancotti C, Lal D, Fariselli P\* (2022). DDGun: an untrained predictor of protein stability changes upon amino acid variants. **Nucleic Acids Research**. DOI: 10.1093/nar/gkac325

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Capriotti E, Marti-Renom MA (2008). Computational RNA structure prediction. Current Bioinformatics, 3; 32-45.

Liu T, Tang GW, Capriotti E\*. (2011). Comparative modeling: the state of the art and protein drug target structure prediction. Combinatorial Chemistry & High Throughput Screening. 14: 532-547.

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*Capriotti E\**, *Nehrt NL, Kann MG\**, *Bromberg Y\**. (2012). Bioinformatics for personal genome interpretation. **Briefings in Bioinformatics.** 13: 495-512.

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Sanavia T, Birolo G, Montanucci L, Turina P, Capriotti E\*, Fariselli P\*. (2020). Limitations and challenges in protein stability prediction upon genome varia-tions: towards future applications in precision medicine. Computational and Structural Biotechnology Journal. 18: 1968-1079.

Petrosino M, Novak L, Pasquo A, Chiaraluce R, Turina P, Capriotti E\*, Consalvi V\*. (2021). Analysis and interpretation of the impact of missense variants in cancer. International Journal of Molecular Sciences. 22:5416. DOI:10.3390/ijms22115416.

*Merlotti A, Menichetti G, Fariselli P, Capriotti E, Remondini D.* Network-based strategies for protein characterization. **Advances in Protein Chemistry and Structural Biology**. Elsevier 2021. DOI:10.1016/bs.apcsb.2021.05.001.

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Casadio R, Calabrese R, Capriotti E, Compiani M, Fariselli P, Marani P, Montanucci L, Martelli PL, Rossi I, Tasco G Machine learning and the prediction of protein structure: the state of the art. X International Conference on Information Processing and Management of Uncertainty in Knowledge-Based Systems (IPMU 2004). July 4-9 2004, Perugia Italy, Publisher: La Sapienza, Roma, pag 933-940.

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