CURRICULUM VITAE

GENERAL INFORMATION

Citizen:

Education: Laurea in Scienze dell'Informazione (University of Pisa)

PhD in Artificial Intelligent Systems (University of Ancona)

PROFESSIONAL ADDRESS:

Emanuela Merelli

Università di Camerino, Computer Science

e-mail:

home-page:

POSITION

Full Professor of Computer Science at Universit`a di Camerino (Unicam)

APPOINTMENTS

- · Member of Academic Senate of the University of Camerino
- · Vi-President of Italian Chapter of EATCS
- · Member of the Board of the CINI Laboratory of Big Data and Data Science
- · Coordinator of PhD in Computer Science and Mathematics, ISAS-University of Camerino (2010 202
- · Member of the Council of the European Association for Theoretical Computer Science (2017 2021)
- · Fellow member of the COST Action CA19122, EU Net for Gender Balance in Informatics (2020-2024)
- · Delegate to the European Mobility of the University of Camerino (2011 2012)
- · Head of Computer Science Div., School of Science and Tech., University of Camerino (2006 2012)
- · Fulbright Scholar at University of Oregon, Computer Science Dep., Eugene, US (2005 2006)
- · Visiting Researcher at University of East Anglia, School of Information System, Norwich, UK (1998)
- · Research Fellow at National Research Institute (CNR), Pisa, Italy (1987)
- · Delegate to ISO/WG6 for Italy, Editor of OSI/WG6-Presentation Layer (1986 1987)
- · Scientist Fellow at European Networking Center, IBM, Heidelberg, Germany (1985 1986)
- · Honorary Scholar at In . Olivetti S.p.A., Ivrea, Italy (1984 1985)

PROFESSIONAL CONTRIBUTIONS

- · Scientific Coordinator "Biomolecules computing cell behaviour" project funded by Diatech Pharmacogenetics srl, (2022-2024)
- Project Coordinator of the EC FP7, FET Proactive, "TOPDRIM: Topology Driven Methods for Modelling Complex Systems", GA N. 318121, (2012-2015)
- · Principal Investigator of the ERASMU+ KA2 HEI Cooperation for Innovation,
- "Data Science Pathways to Re-image Education", Grant n.2016-1-IT02-KA203-024645, (2014-2020)
- Principal Investigator of the COST Action on "Reversible Computaxitem ding horizon of computing" IC1405, (2015-2017)
- Principal Investigator of the IT Flagship, RITMARE, "Agent-oriented modelling for spacial and temporal multiscale analysis, simulation and prediction of a dynamics population in the marine ecosystem" project SP2-WP2-AZ2-UO05, (2012-2016)
- · Principal Investigator of the FESR 2007-2013, "AALISABETH: "Ambient-Aware LifeStyle tutoring, Aiming at a BETter Health" project, (2013-2015)
- · Principal Investigator of IT-MIUR, FIRB, "LITBIO: Interdisciplinary Laboratory of Bioinformatics Technologies", project, (2005-2009)
- · Scientific delegate at FP6, NoE, Virtual Physiological Human- Network of Excellence, (2009-2011)
- · Principal Investigator of IT-MIUR-Strategic Research Project, "O2I: Oncology over Internet -

Methodologies, Models, Techniques and Tools, per Information Extraction and Retrieval", (2002-200). Principal Investigator of CIPE MARCHE "SICOM: SIstemi COoperativi e Multiagente", (2003-2004)

EXPERT and EVALUATOR

- · European Commissio FP7, H2020, Horizon-EIC
- · Italian Ministry for Research and Education, MIUR
- · Fulbright US-Italy Program
- · Quatar National Research Fund, QNRF
- · Region Marche, Regione Lombardia, Provincia di Trento R&D&I and Technology Transfer
- · Galileo Program for the Italo-French Cooperation

TEACHING ACTIVITIES

- · Distributed Calculus and CoordinatioMultiagent Systems Lab Algorithms and Data Structures
- · Machine Learning Computability Theory Data Science & TDA · Information Systems and DB · Algorithms and Complexity · Operational Research · Non-linear Optimisation · Methods and Programmin Languages (1 current academic year)

MEMBERSHIPS

- · Member of EATCS (European Association for Theoretical Computer Science)
- · Member of ACM (Association for Computing Machinery)
- · Member of Bioinformatics Italian Society (BITS)
- · Member of Gruppo GNCS (Gruppo Nazionale per il Calcolo Scientifico)

FOUNDING MEMBER OF SPIN-OFF

- · NGB: New Generation Bioinformatics dell'Universit`a di Camerino (2010) http://www.engeebee.com
- · eLios: eLinking on line services S.r.l., Universit`a di Camerino (2007) http://www.e-lios.eu

SCIENTIFIC ACTIVITIES

REFEREE

ACM Transaction on Computational Biology and Bioinformatics · Acta Informatics, Springer · Americar Mathematical Society · Applied Mathematics and Computation, Elsevier · Bioinformatics, Oxford Journ · BioSystemsElsevier Journal · BMC Bioinformatics · Briefings in Bioinformatic ①xford Journal · Bulletin of Mathematical Biology · Computer and Industrial Engineering, Elsevier · Data and Knowledg Engineering, Elsevier · IEEE Transaction on Parallel and Distributed Computing · International Journal of Computer MathematicsInternationalJournal for Production ResearchInternationalJournal of Modelling, Identification and Control · International Journal on Software Tools for Technology Transfer Springer · Journal of Theoretica Biology, Elsevier · Scientific ReportNature · NeuralComputing and Applications, Springer Journal · PLOS, Computational Biology · PLOS ONE · Robotics and Computer-Integrated Manufacturingsevier · Simulation Modelling Practice and Theoretical

GUEST EDITOR

Journal of Theoretical Computer Science - Theory of Natural Computing, Essedied issues on:

Computer Science, Elsevier · Transactions on Computational Systems Biology, Springer

- · From Computer Science to Biology and Back6081, (2015)
- · Interaction between Computer Science, Complex Systems and Bio 687,, (2015)
- · Hybrid Automata and Oscillatory Behaviour in Biological System 1/0 issue 20, (2010)
- · Concurrent Systems Biology: Nadia Busi (1968-2007), Val. 0, issue 33-34.3037-3038, (2009)

Journal of Electronic Proceedings of Theoretical Computer Science dings of:

- · 5th CS2Bio Workshop Interaction between Computer Science and Biologo, (2014)
- · 4th CS2Bio Workshop Interaction between Computer Science and Biol299, (2013)
- · 2nd FBTC Workshop From Biology To Concurrency and Back220J.issue 1, (2009)
- · 1st FBTC Workshop From Biology To Concurrency and Back1941.issue 3, (2008)

ICALP Proceedings:

- · ICALP 2022, LIPIcs Vol. 229 (2022)
- · ICALP 2021, LIPICS Vol. 198 (2021)
- · ICALP 2020, LIPICS Vol. 168 (2020)

Springer Proceedings in ComplexPtyceedings of:

· ECCS 2014, ISBN 978-3-319-29228-1 (2016)

Transaction on Computational Systems Biology, SprSpeeial issue on:

• Model and Metaphors from Biology to Bioinformatics and Back73701(2005)

AWARDS

FET11: The European Future Technologies Conference and Exhibition.3rd Price for the Best Poster: MethodologicaBridges for Complex Systemsco-authors:Pietro Lio' and Nicola Paoletti. http://www.fet11.eu/awards

DISSEMINATION ACTIVITIES

- · Member of ICALP Steering Committee
- · Chair and co-chair of several events with an interdisciplinary character, among which the Internatio School on Concurrency and Complexity through Topology for Young Respectation for Theoretical Computer Science and the TOPDRIM EU-project.
- · Member of many program committees of workshops and conferences
- · Advisor and Mentor of several brilliant PhD Candidates among which Leonardo Mariani, Ezio Bartoco and Nicola Paoletti

RESEARCH ACTIVITIES

Author or co-author of bout a hundred peer-reviewed articles in internation alls and volumes. Most cited paper: "A tabu search method guided by shifting bottleneck for the job shop scheduling problem" E. Merelli, F. Pezzella European Journab f Operationa Research 120 (22,97-310,2000): ~ 329 Scholar (172 Scopus) citation 28 Scholar (18 Scopus), total citation 2244 Scholar (1110 Scopus)

MAIN FIELD OF RESEARCH

TopologicaField Theory ofData and S[B]models ofcomputationagent-oriented modelling & multi-levelcomplex systems; o-inspired formahethods. Computationabiology ofRNA Folding. n-body interaction for the Immune System metaphor of memory evalue in the Immune System metaphor of memory evalue.

MOST RELEVANT CONTRIBUTIONS

The most relevant contributions in the various areas are:

- i) a research program towards a new strategy for mining data through data language that turns out to be the shape language pological Field Theory of Data; [19-20]
- ii) an extension to n-body interactions of Parisi's idiotypic network model to study the memory evolution of human **Immune System** as a topological application of the S[B] paradigm; [27]

- iii) **the S[B] machine** a new model of computation supporting data-driven evolution of the model. characterized by two entangled levelses tription global and local: the environment, properly the semantic context where the computation takes place through a pool of interacting agents each of which behaves as a deterministic machine; [23]
- iv) An algebraic language for RNA structure comparison \$\delta \text{O}_3\s new bio-inspired formalethods such as BioAgent, **SHAPE Calculus**, BIOSHAPE, and Biological Oscillators synchronization language (BOSL), for modellingsimulating and analysing autonomous agents represented as geometric shapes whose internal behavior allows them to interact, bind and move in a Euclidian space; [42,43] v) the construction of lermes an agent-based middleware for mobile computing to basis of
- **Orion** an agent-based spatial simulator for MAS. vi) the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and implementation of jHoles algorithm based on the design and jHoles algori
- vii) **Persistent Entropy**, a new measure of complexity defined as a function of Betti barcodes strong related to the topological structure of the data space and a methodology to construct a persistent er automaton from topological data analysis; [25]
- ix) a wide study on bone remodelling as a multismalleilevelsystem based on Spatla Bystems, complex automata and BioShape, by combining process algebraic and stochastic approach and by melling some pathologies such as osteomyelitis and osteoporosis; [33-35]
- x) a model of cell cycle viewed as a model reduction of an hybrid system that captures the crucial feat of a dynamical system has of hybrid automata technology allows us to describe the cellular system by combining continuous behavior with discrete events; [38]
- xi) a new data model, Resourceome, that allows the managing of declarative and procedural knowled with a unique model whose actions connect the use of a resource to its domain; [61]
- xii) the design and development of DISPAS, Demersal fish Stock Probabilistic Agent-based Simulator, investigate and understand sustainability in the exploitation of fishery resources, by focusing on comsole (Solea solea) stock in the Northern Adriatic Sea; [30]
- xiii) a new heuristic method combines Tabu Search and Shifting Bottleneck and solves the optimizati problem of job shop scheduling in a better time [64].

INVITED TALKS

- 2022 May, Tampa, USA. "The topology of RNA folding"

 Workshop on Discrete and Topological Models in Molecular Biology
- 2019 August, Stony Brook, USA. "A topological Interpretation of Interactive Computation" Symposium in honor of Scott Smolka on the occasion of 65th birthday
- 2019 February Lugano, "Topology Driven Methods for Complex Systems an alternative approach to Machine Learning Theory and some Applications" IDSIA-Dalle Molle Institute for Artificial Intelliger
- 2019 November, Milan, "Interdisciplinamyobstacles course rising" Symposium in honour of G Mauri
- 2019 September, Como, "Present and future of Theoretical Computer Science (not only) in Italy" Panel-ICTCS
- 2017 June, Wien, "A topological view of Compositionality of Process Algebra", Open Problems in Concurrency Theory, IFIP Group
- 2016 December, Berkeley, USA. "Topological Approach to Compositionality in Complex Systems", Simon Institute
- 2016 September, Amsterdam, "On topological characterization of complex systems", DyM-CS- CCS
- 2016 September, Cagliari "Tutorial on Topological data ajihbd Jesisand persistent entropy" KBWeb
- 2016 July, Wien, "Topological Field Theory of Datarogram towards a novel strategy for mining data through data language", DataMOD

- 2015 December, New York, USA. "Topological Field Theory of Data", BICT
- 2015 June, London, "TOPDRIM a global vision of datan-locality, topology, formal languargems: tools to handle large data set", NGS
- 2015 March, Brussels, "Topological Field Theory of Data, the new science for computer science", EU FET seminar
- 2012 August, Taormin'aThe Immune System as a Metaphor for Topology Driven Patterns Formation in Complex Systems" at the I@bnference on Artificial Immune Systems"
- 2011 September, Palermo, IT. "Self-adaptive systems for Ageing and Quality of Life" at the AI*IA workshop on Technological Challenges and Scenarios of the Ageing Society"
- 2009 April 16, San francisco, USA. "Model Checking Biological Oscillators" at HSCB'09 on "Hybrid Systems Approaches to Computational Biology"
- 2007 September 18, Reykjavik, IS. "Agent-based Modelling and Simulation in Systems Biology" at Sc of Computer Science, Reykjavik University
 2007 October 25, Pis' Agent-based Modelling and Simulation in Systems Biology" at MiniWorkshop
- 2007 October 25, Pis'Agent-based Modelling and Simulation in Systems Biology" at MiniWorkshop on "Computational Approach to Biology" Scuola di dottorato Galileo Galileo Galileo Galileo Cannata and Luca Tesei)

2007 November 8, Stony Brook, USA. "Agent-based Modelling and Simulation in Systems Biology" at

- Dept. of Computer Science State University of New York
 2007 December 10, Milanorion Framework for systems Biology applications" at BioinfoGRID
- 2007 December 10, Milañorion Framework for systems Biology applications" at BioinfoGRID (Bioinformatics Grid Application for Life Science) Symposium 2007.
- 2005 February 1, Stanford Universityent in Bioinformatics" at Center for Biomedical Computation

LIST OF SELECTED PUBLICATIONS

- 1. D. Marchei, E. MerellRNA secondary structure factorization in prime tangles BMC Bioinformatics, 23, 345 (2022)
- 2. M. Quadrini, L. Tesei, E. Merellautomatic generation of pseudoknotted RNAs taxoBMfay. Bioinformatics, 23, 575 (2022)
- 3. S. Maestri, E. Merelli, M. Pettin and models for detecting the driving forces of biomolecular interaction cientific Reports, 12(1), 1878 (2022)
- 4. M. Belenchia G. Rocchetti, S. Maestri, A. Cimadamore R. Montironi, M. Santoni, E. Merelli: Agent-Based Learning Mod Redr the Obesity Paradox in RCCF rontiers in Bioengineering and Biotechnology, 9, 642760 (2021)
- 5. A. Mancini, L. Vito, E. Marcelli, S. Pucciarelli, E. Merelli: Machine learning models predicting multidrug-resistant urinary tract infections using dsaaS. BMC Bioinformatics, 21, 347 (2020)
- 6. M. Piangerelli, S. Maestri, E. Merelli: Visualising 2-simplex formation in metabolic reactions. Journal of Molecular Graphics and Modelling, 9701(2020)
- 7. M. Quadrini, L. Tesei, E. MerellASPRAlign: A tool for the alignment of RNA secondary structures with arbitrary pseudoknobis informatics, 36(11), pp. 78–3579 (2020)
- 8. M. Quadrini, E. Merelli, R. Piergalliniabel core for understanding RNA structunes-LNBI, vol. 12313 (2020)

- 9. S. Maestri, E. Merelli: Algebraic characterisation **ι**ννοί n-coding RNA.LNCS-LNBI, vol 12313 (2020)
- 10. S. Maestri, E. Merelli: Process calculinay reveal the equivalence lying at the heart of RNA and proteins Scientific Reports, vol. n. 1 (2019)
- 11. E. Merelli, A. Wasilewska: opological interpretation of interactive computation. LNCS, vol. 11500 (2019)
- 12. M. Quadrini, L. Tesei, E. Merelli: An algebraic languages fo RNA structure comparism C Bioinformatics, vo20, n.161 (2019)
- 13. M. Piangerelli L. Tesei, E. Merelli: A Persistent Entropy Automaton for the Dow Jones Stock Market. LNCS vol. 11761 (2019)
- 14. M. Quadrini, E. Merelli, R. Piergallinioop grammars to identify RNA structural patteros. of BIOINFORMATICS (2019)
- 15. L. Bettini, E. Merelli, F. TiezziX-Klaim is backLNCS vol. 11665 (2019)
- 16. M. Piangerelli, M. Rucco, L. Tesei, E. Merellippological classifier for detecting the emergence of epileptic seizure MC Research Notes, vol. (1) (2018)
- 17. M. Quadrini, E. MerelliLoop-loop interaction metrics on RNA secondary structures with pseudo-knots.BIOINFORMATICS (2018)
- 18. J. H. Johnson, L. Tesei, M. Piangerelli E. Merelli, R. Paci, N. Stojanovic P. Leit ad, Barbosa, M. Amador: Big Data: Business, Technology, Education, and Sciebioquity vol. 2 n. 1 (2018)
- 19. D. Cacciagrano, F. Corradini, E. Merelli, L. Tebriformity in Multiscale Modefsom Complex Automata to BioShape. Cellular Automata vol12 n.5 (2017)
- 20. M. Rucco, R. Gonz´alez-D´ıMz, J. Jim´enez N. Atienza, C. Cristalli, E. Concettoni A. Ferrante, E. Merelli: A new topological ntropy-based approach for measuring similarities among piecewise linear function Signal Processing vol 34 (2017)
- 21. M. Rasetti, E. Merelli: The Topologica Field Theory of Datamining data beyond complex networks. In Advances in Disordered System Andom Processes and Some Application Processes and So
- 22. M. Rasetti, E. MerelliThe Topological Field Theory of Dataprogram towards a novel strategy for mining data through data language hysics: Conf. Series, vol.626, (2015)
- 23. A.L. Mamuye, Merelli, L. Tesei: Graph Grammar for Modeling RNA Folding Evolution as a Self-Adaptive System Proceedings of Graphs as Models Electronic Proceedings in Theoretical Computer Science, Val31, (2016)
- 24. E. Merelli, M. Piangerelli, M. Rucco, D. Tollartopological approach for multivariate time series characterization epileptic brain EAI Endorsed TransSelf-Adaptive Systems 2(%): (2016)
- 25. E. Merelli, N. Paoletti, L. TeseAdaptability checking in complex systemience of Computing Programming, vol.115-116, (2016)

- 26. Rucco Matteo, Castiglione Filippo, Merelli Emanuela, Pettini Marco. Characterisation of the Idiotypic Immune Network Through Persistent Entropy. Proceedings of ECCS 2014. 117-128, Springer (2016)
- 27. E. Merelli, M. Rucco, P. Sloot and L. Tesei: Topologica Characterization of omplex Systems: Using Persistent Entropy, 17(10), (2015)
- 28. E. Merelli, I. Petre: From Computer Science to Biology and Bableor. Comput. Sci. vol. 608, (2015)
- 29. E. Merelli, M. Pettini, M. RasettTopology driven modelithe IS metaphor, *Natural Computing Journal*, *NACO*, 14(3), Springer, (2015)
- 30. P. Giannini, E. Merelli, A. Troina: Interactions between Computer Science and BioToggor. Comput. Sci. vol. 587, (2015)
- 31. J. Binchi, E. Merelli, M. Rucco, G. Petri, F. Vaccarj**Ho**les:A Tool for Understanding Biological Complex Networks via Clique Weight Rank Persistent Hom**&legty**... Notes Theor.Comput. Sci. vol.306, (2014)
- 32. Pierluigi Penna, Nicola Paoletti, Giuseppe Scarcella, Luca Tesei, Mauro Marini, Emanuela Merelli: DISPAS: An Agent-Based Toofor the Management offshing Effort. SEFM Workshops 2013: 362-367
- 33. E. Merelli, M. Rasetti: Non locality, topology, formal languages: new global tools to handle large data sets, Procedia Computer Science, vol.18, (2013)
- in Complex System Int. Conference on Artificial Immune Systems. Taormina, (2012)

34. E. Merelli, M. RasettiThe Immune System as a Metaphor for Topology Driven Patterns Formation

- 35. P. Li`o, N. Paoletti, M. A. Moni, K. Atwell, E. Merelli, M. Vicedotelling osteomyeliteMC Bioinformatics 13(S-145)12 (2012)
- 36. N. Paoletti, P. Lio', E. Merelli, M. Vicecontiultilevel Computational Modeling and Quantitative Analysis of Bone Remodeling EE/ACM Trans. Comput. Biology Bioinformatics, 9(5), (2012)
- 37. P. Li`o, E. Merelli, N. PaolettDisease processes as hybrid dynamical systems/orkshop on *Hybrid Systems*. Newcastle, (2012)
- 38. F. Buti, F. Corradini, E. Merelli, L. Tese Geometrical Refinement of Shape Calculus Enabling Direct Simulation Conference on Simulation, Meth. And Techn. And Appl. Rome, (2012)
- 39. E. Bartocci, P. Li`o, E. Merelli, N. Paoletti: Multiple Verification in Complex Biologic stems: The Bone Remodelling Case Studings. Computational Systems Biology vol. 14, (2012)
- 40. R. Alfieri, E. Bartocci, E. Merelli, L. Milane Modeling the cell cyclerom deterministic models to hybrid system sournal of Biosystems 105(1), 2011
- 41. F. Buti, D. Cacciagranof, Corradini, E. Merelli, L. Tesei: A Uniform Multiscale Meta-modell BioShapeElectr. Notes TheorComput.Sci. vol.277, (2011)
- 42. P. Li`o, E. Merelli, N. Paoletti, M. Viceconti: A Combined Process Algebraic and Stochastic Approach to Bone Remodelingectr. Notes Theor Comput. Sci. vol. 277, (2011)

- 43. E. Bartocci, F. Corradini, E. Merelli, L. Tesedetecting Synchronisation of Biological Oscillators by by Model Checking *Theoretical Computer Science* 411(20), (2010)
- 44. E. Bartocci, M.R. Di Berardini, D. Cacciagrano, E. Merelli, L. Tlesed Operational Semantics and Well-Formedness of Shape Calculustific Annals of Computer Science, vol.20, (2010)
- 45. E. Bartocci, M.R. Di Berardini, F. Corradini, E. Merelli, L. Tesei: Shape Calculus A Spatial Mobile Calculus for 3D Shapescientific Annals of Computer Science 2001 (2010)
- 46. E. Bartocci, D. Cacciagrano, F. Corradini, E. Merelli, and L. **WiRe**sourceome for the automation of in-silico biologic**a**kperimentsAppearing in proceedings of *Internation@onference on Computational and Systems Biology and Microbiology (BioSysCom)*, Cancum, Messico, (2010)
- 47. F. Buti, D. Cacciagrano, F. Corradini, E. Merelli, L.TestioShapea spatial shape-based scale-independent simulation environment for biologistems Appearing in proceedings of *International Conference on Computational Science (ICCS)*, Amsterdam, (2010)
- 48. F. Buti, D. Cacciagrano, F. Corradini, E. Merelli, M. Pani, L. Tæsne remodelling in BioShape. In CS2BIO 2010: Interactions between Computer Science and Biology, 1st International Workshop, (2010)
- 49. D. Cacciagranof. Corradini, E. Merelli. Bone Remodellinga Complex Automata-based model running in Bioshape *ACRI 2010: The Ninth International Conference on Cellular Automata for Research and Industry*, Ascoli Piceno (Italy), (2010)
- 50. E. Bartocci, F. Corradini, E. Merelli, L. Tesei: Detecting synchronisation of biologise illators by modelchecking. Theor. Comput. Sci. 411(20):1999-2018 (2010) heor. Comput. Science 411(20), (2010)
- 51. F. Buti, F. Corradini, E. Merelli, E. Paschini, P. Penna, L. Teseindividual-based Probabilistic Model for Fish Stock Simulation MCA-POP 37-55, (2010)
- 52. E. Bartocci, F. Corradini, E. Merelli, L. Tesei. Model Checking Biologic Scillators. *Electronic Notes in Theoretical Computer Science* 229(1), (2009)
- 53. N. Cannata, F. Corradini, E. Merelli, F. Piersigilli, L. Vito Towards Bioinformatics Resourceomes. *Biomedical Data and Applications* 13-36, (2009)
- 54. N. Cannata, F. Corradini, E. MerelMultiagent modelling and simulation of carbohydrate oxidation *International Journal of Modelling, Identification and Control* (IJMIC), 1(3), (2008)
- 55. E. Bartocci, F. Corradini, R. Grosu, E. Merelli, O. Riganelli, S. A. SmoßkanyCam: Formal Framework for Modeling, Analyzing and Regulating Cardiac My@oytes:ency, Graphs and Models, LNCS vol.5065, (2008)
- 56. E. Merelli et al Agents in Bioinformatics, Computational and Systems Bioinformatics, 8(1), (2007)
- 57. E. Merelli, M. Young. Validating MAS with mutation *International Journal of Multiagent and Grid Systems* 3(2), (2007)
- 58. N. Cannata, F. Corradini, E. Merelli. A Resourceomic Grid for Bioinformatic ** atternational Journal of Grid Computing: Theory, Methods and Applications: Future Generation Computer Systems Journal, 23(3), (2007)

- 59. E. Bartocci, F. Corradini, E. Merelli, L. ScortichBioWMS: a web-based Workflow Management System for BioinformaticBMC Bioinformatics Int. Journal, vol.8 suppl.1, (2007)
- 60. Bartocci, E., Cacciagrano, D., Cannata, N., (...), Milanesi, L., Roman agent-based multi-layer architecture for bioinformatics grids *EEE Transactions on Nanobioscience* 6(2), (2007)
- 61. F. Corradini, E. Merelli: Hermes: Agent-Based Middleware for Mobile Computing M 2005 LNCS Vol. 3465.234-270, (2005)
- 62. N. Cannata, F. Corradini, E. Merelli, A. Omicini, A. RArciagent-oriented conceptual framework for Systems Biology Transaction on Computational Systems Biology, LNBI Vol. 3737,105-122, Springer, (2005)
- 63. N. Cannata, E. Merelli, R.B. Altman. Time to organize the Bioinformatics Resource *Phos. Computational Biology*, Vol. 1, Num. 7, (2005)
- 64. M. Luck and E.Merelli. Agents in Bioinformatic The Knowledge Engineering Review, vol. 20, Num. 2 117-125, Cambridge University Press, (2005)
- 65. F. Corradini, L. Mariani, and E. Merelli. An agent-based approach to time tegration *Software Tools for Technology Transfer*, Vol. 6 Num. 3, 231-244, Springer Journal, (2004)
- 66. E. Merelli, F. Pezzella. A tabu search method guided by a shifting bottleneck for a job shop scheduling European Journal of Operational Research, vol. 120, (2000)
- 67. R. De Leone, R. Capparuccia, E. Mere**A**isuccessive overrelaxation back propagation algorithm for neural network trainin**te** *E Transactions on Neural-Networks*, 9(3), (1998)
- 68. F. Caneschi, E. MerellÄn architecture for an asn.1 encoder/decodeputer Networks, North Holland, vol.14, (1987)

Camerino, October 2023