

Curriculum Vitae of Daniele Marchisio

Place and date of birth	Savona (Italy), 21 December 1973
Citizenship	Italian
Work address	Dipartimento di Scienza Applicata e Tecnologia, Istituto di Ingegneria Chimica, Politecnico di Torino, C.so Duca degli Abruzzi 24, 10129 Torino (TO) - Italy
Telephone	0039-011-0904622
Email	daniele.marchisio@polito.it
Skype	daniele.marchisio.torino
Webpage	http://www.disat.polito.it/personale/scheda/(nominativo)/daniele.marchisio
Current position	<i>Full Professor</i>

Education

BS & MS (<i>cum laude</i>) in Chemical Engineering	1992 - 1997	Politecnico di Torino (Italy)
PhD in Chemical Engineering	1998 – 2001	Politecnico di Torino (Italy)
	1999 - 2000	Visiting Scholar at Iowa State University (USA)
Post-doc in Chemical Engineering	2001 – 2003	Iowa State University (USA)
Post-doc / academic guest in Chemical Engineering	2003 (Dec.) - 2004 (Apr.)	Eidgenössische Technische Hochschule (ETH) of Zurich (Switzerland)

Work & research experience

2004 – 2010	<i>Assistant Professor</i> at the Department of Material Science and Chemical Engineering of the Politecnico di Torino (Italy)
Summers of 2007 & 2008	<i>Visiting Professor</i> at the Department of Chemical Engineering of University College London (UK)
2010 – 2016	<i>Associate Professor</i> at the Department of Material Science and Chemical Engineering (now Applied Science and Technology) of the Politecnico di Torino (Italy)
2012 (Dec.) – 2013 (Feb.)	<i>Visiting Scientist</i> at CSIRO (CMIS – Clayton, Melbourne, VIC, Australia)
2016 – present	<i>Full Professor</i> at the Department of Applied Science and Technology of the Politecnico di Torino (Italy)
2016 – present	<i>Adjunct Visiting Professor</i> at the Beijing University of Chemical Technology (China)

International Awards & Recognition

1997	Laurea Prize “Vittorio de Bernochi” from the Association of Architects and Engineers of the Politecnico di Torino as the best graduate in Chemical Engineering in the year 1997
1998	Prize “Optime” from the Industrial Union of Turin as one of the best 100 students of the year
2001	United Engineering Foundation Conference Fellowship, for attending the Chemical Reaction Engineering Conference (24-29 June 2001, Barga, Italy)
2007	Most cited paper for Chemical Engineering Science: D.L. Marchisio, R.D. Vigil, R.O. Fox (2003) Implementation of the Quadrature Method of Moments in CFD codes for aggregation-breakage problems, Chemical Engineering Science, 58, 3337-3351
2007	Recipient of the International Incoming Short Visits fellowship funded by the Royal Society of the United Kingdom
2010	Scimedirect top 25 most downloaded article for: M. Hussain, R. Ceccarelli, D.L. Marchisio, D. Fino, N. Russo, F. Geobaldo (2010) Synthesis, characterization and photocatalytic application of novel TiO ₂ nanoparticles, Chemical Engineering Journal. 157, 45-51

Membership in Professional Organizations and Scientific Committees

- Member of the American Institute of Chemical Engineering (AIChE)
- Member of the Italian Association of Chemical Engineering (AIDIC)
- Member of the European Federation of Chemical Engineering (EFCE)
- Member of the Working Parties on Industrial Crystallization and Multiphase Flows of the EFCE
- *Consigliere* (Vice-president) of the GRICU (Gruppo Ingegneria Chimica dell'Università)

Membership in editorial boards of international journals, scientific committees of international conferences and reviewing activity

- Associate Editor for the Canadian Journal of Chemical Engineering [http://onlinelibrary.wiley.com/journal/10.1002/\(ISSN\)1939-019X/homepage/EditorialBoard.html](http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)1939-019X/homepage/EditorialBoard.html)
- Member of the Advisory Board of Chemical Engineering & Technology [http://onlinelibrary.wiley.com/journal/10.1002/\(ISSN\)1521-4125/homepage/2044_edbd.html](http://onlinelibrary.wiley.com/journal/10.1002/(ISSN)1521-4125/homepage/2044_edbd.html)
- Member of scientific/organizing committees of the International Symposium on Industrial Crystallization (ISIC), International Conference on Multiphase flows, European Conference on Mixing, Population Balance Modelling Conference
- Reviewer for the most important journals in his field (*Journal of Computational Physics, International Journal of Multiphase Flows, A.I.Ch.E. Journal, Chemical Engineering Science, Journal of Colloid and Interface Science, Chemical Engineering Journal, Industrial and Engineering Chemistry Research, Chemical Engineering Research and Design, Computers & Chemical Engineering, Physical Review E, Journal of Pharmaceutical Sciences, etc.*) as well as for the Cambridge University Press for book proposal reviewing
- Reviewer for grant proposals for Swiss National Science Foundation, German Science Foundation (DFG), Austrian Science Foundation, Beijing University of Chemical Technology
- Member of the Advisory Board of the CORAL EPSRC project <http://gow.epsrc.ac.uk/NGBOViewGrant.aspx?GrantRef=EP/N024915/1>

Invited Keynote/Plenary Lectures, Invited Seminars and Organization of Summer Schools

- Daniele Marchisio has delivered about **20+ invited/keynote/plenary lectures at important international conferences** (among which: BIRS-CMO Workshop: New Frontiers in Multiphase CFD for the 21st Century Energy Mix at Casa Matemática Oaxaca, Mexico, 2018; 14th International Conference on Multiphase Flow, Desenzano, Italy, 2017; 15th Multiphase Flow Conference and Short Course: Simulation, Experiment and Application, Dresden, Germany, 2017; Dynamics of Evolving Fluid Interfaces – DEFI; Lyon, France, 2016; 19th International Symposium of Industrial Crystallization, Toulouse, France, 2014; 5th International Conference on Population Balance Modelling, Bangalore, India, 2013; 9th European Congress of Chemical Engineering, The Hague, The Netherlands, 2013; North American Mixing Forum – Mixing XXIII, Cancun, Mexico, 2013; 50th European Two-Phase Flows Group Meeting, 2012 - 2nd Joint ETPFG-EFCE Multi-Phase Meeting, 2012, Udine, Italy; etc.)
- Daniele Marchisio has delivered **30+ invited seminars at public and private institutions** around the world (TU Darmstadt, TU Munich, University College London, CSIRO Melbourne, MIT, University of Warwick, CEA Marcoule, Warsaw Technical University, Eindhoven Technical University and Multiscale Institute, Aalto University, University of Alberta, Imperial College, etc.)
- Daniele Marchisio has **organized and lectured in 14 advanced doctorate summer schools** on several topics including: multiscale modelling, simulation of multiphase flows and population balance modelling (GRICU Summer School on Multiscale Modelling, Palermo, Italy, 2017; Multiscale modelling of flowing soft matter, CISM, Italy, 2016; Multiscale modelling and Multiphysics coupling in solid and fluids mechanics, TEC21, France, 2015; 3rd Summer School of the IMPRS Magdeburg, Germany, 2013; Computational Models for Polydisperse Particulate and Multiphase Systems, CSIRO – CMIS, Australia, 2013; Multiphase turbulent reacting flows, CISM, 2006; etc.)

Tutoring & Research Activity / Coordination of National and International Project

Daniele Marchisio has acted as **supervisor for 50+ master students and 20+ PhD students**.

Daniele Marchisio's research activity focuses on the development, validation and implementation of computational methods for multiscale modeling, with a particular focus on turbulent multiphase reacting systems. His early research interests focused on the treatment of very fast chemical reactions with Computational Fluid Dynamics (CFD) through the Reynolds-Averaged Navier-Stokes equation (RANS) and the Large Eddy Simulation (LES) approaches. More recently he has been interested in the description of the evolution of multiphase systems through Population Balance Models (PBM). He developed, investigated and validated an entire class of methods (Quadrature-Based Moments Methods, QBMM) that are now employed in commercial and open source CFD codes. These methods have been also applied to the atomistic description of fluids, by solving the Boltzmann equation. Lately the problem of coupling, following the multiscale approach, different scales, namely fully atomistic and coarse-grained molecular dynamics simulations (by using GROMACS and LAMMPS), mesoscale models, such as Dissipative Particle Dynamics (DPD) and continuum models (CFD), has been investigated.

Daniele Marchisio has acted as **Principal Investigator in the following projects** (selection of most relevant project in recent years):

- Research project funded by the European Commission (H202) titled: “SimDOME: Digital Ontology-based Modelling Environment for Simulation of materials”, 2019 – 2022, € 575.000
- Research project funded by the European Commission (H2020) titled “VIMMP: Virtual Materials (Modelling) Marketplace”, 2018-2021, € 650.000
- Consultancy Project titled “CFD simulation of precipitation processes” granted by BASF, 2017-2019, € 60.000
- Research Project titled “CFD simulation of foam formation in carbonated beverages” granted by Sidel/Tetrapack/Optimad, 2017-2020, € 30.000 + 85.000
- Research project titled “Simulation of industrial bubble columns under heterogeneous regime with CFD and PBM” funded by the IFP Energie Nouvelles, Lyon (France), 2015-2018, € 30.000
- Collaborative project funded by the European Commission (FP7) titled “MODENA: Modelling of morphology development of micro- and nano-structures”, 2014-2016, € 305.000

- Research project titled “CFD simulation of gas-liquid reactors with population balances” funded by BASF, Ludwigshafen (Germany), 2012-2014, € 120.000

Publications & Citation Report

Daniele Marchisio has published 140+ papers on international journals and 130+ papers on proceedings of international conferences. He has authored one book (Marchisio D.L., Fox R.O. (2013) Computational Models for Polydisperse Particulate and Multiphase Systems, Cambridge University Press: Cambridge; ISBN: 978-0-521-85848-9) and edited another one (Marchisio D.L., Fox R.O. (2007) Multiphase reacting flows: modelling and simulation, Springer: Wien; CISM Series, Vol. 492, ISBN: 978-3-211-72464-4).

January 2019	Scopus	Web of Science
Total number of documents	139	124
Total number of citations	4066	3571
Hirsch factor (<i>H-index</i>)	33	31

In particular **Daniele Marchisio** meets the requirement to act as “**Commissario all'Abilitazione Nazionale nel Settore 09/D2**”.

List of 25 most recent publications

1. Bazzano, M., Marchisio, D., Sangermano, M., Wörner, M., Pisano, R. A molecular dynamics approach to nanostructuring of particles produced via aerosol cationic photopolymerization (2019) *Chemical Engineering Science*, 195, pp. 1021-1027.
2. Salenbauch, S., Hasse, C., Vanni, M., Marchisio, D.L. A numerically robust method of moments with number density function reconstruction and its application to soot formation, growth and oxidation (2019) *Journal of Aerosol Science*, 128, pp. 34-49.
3. Castellano, S., Sheibat-Othman, N., Marchisio, D., Buffo, A., Charton, S. Description of droplet coalescence and breakup in emulsions through a homogeneous population balance model (2018) *Chemical Engineering Journal*, 354, pp. 1197-1207.
4. Droghetti, H., Pagonabarraga, I., Carbone, P., Asinari, P., Marchisio, D. Dissipative particle dynamics simulations of tri-block co-polymer and water: Phase diagram validation and microstructure identification (2018) *Journal of Chemical Physics*, 149 (18), art. no. 184903.
5. Barresi, A.A., Marchisio, D.L. Computational Fluid Dynamics data for improving freeze-dryers design (2018) *Data in Brief*, 19, pp. 1181-1213.
6. Marchisio, D.L., Galan, M., Barresi, A.A. Use of computational fluid dynamics for improving freeze-dryers design and process understanding. Part 2: Condenser duct and valve modelling (2018) *European Journal of Pharmaceutics and Biopharmaceutics*, 129, pp. 45-57.
7. Barresi, A.A., Rasetto, V., Marchisio, D.L. Use of computational fluid dynamics for improving freeze-dryers design and process understanding. Part 1: Modelling the lyophilisation chamber (2018) *European Journal of Pharmaceutics and Biopharmaceutics*, 129, pp. 30-44.
8. Gemello, L., Cappello, V., Augier, F., Marchisio, D., Plais, C. CFD-based scale-up of hydrodynamics and mixing in bubble columns (2018) *Chemical Engineering Research and Design*, 136, pp. 846-858.
9. Gemello, L., Plais, C., Augier, F., Cloupet, A., Marchisio, D.L. Hydrodynamics and bubble size in bubble columns: Effects of contaminants and spargers (2018) *Chemical Engineering Science*, 184, pp. 93-102.
10. Lavino, A.D., Banetta, L., Carbone, P., Marchisio, D.L. Extended Charge-On-Particle Optimized Potentials for Liquid Simulation Acetone Model: The Case of Acetone-Water Mixtures (2018) *Journal of Physical Chemistry B*, 122 (20), pp. 5234-5241.
11. Karimi, M., Marchisio, D., Laurini, E., Fermeiglia, M., Pricl, S. Bridging the gap across scales: Coupling CFD and MD/GCMC in polyurethane foam simulation (2018) *Chemical Engineering Science*, 178, pp. 39-47.
12. Falzone, S., Buffo, A., Vanni, M., Marchisio, D.L. Simulation of Turbulent Coalescence and Breakage of Bubbles and Droplets in the Presence of Surfactants, Salts, and Contaminants (2018) *Advances in Chemical Engineering*, 52, pp. 125-188.

13. Li, D., Buffo, A., Podgórska, W., Marchisio, D.L., Gao, Z. Investigation of droplet breakup in liquid–liquid dispersions by CFD–PBM simulations: The influence of the surfactant type (2017) *Chinese Journal of Chemical Engineering*, 25 (10), pp. 1369-1380.
14. Karimi, M., Droghetti, H., Marchisio, D.L. PU Foam : A novel open-source CFD solver for the simulation of polyurethane foams (2017) *Computer Physics Communications*, 217, pp. 138-148.
15. Coletto, M.M., Marchisio, D.L., Barresi, A.A. Mixing and segregation of wheat bran and vegetable pieces binary mixtures in fluidized and fluid-spout beds for atmospheric freeze-drying (2017) *Drying Technology*, 35 (9), pp. 1059-1074.
16. Li, D., Gao, Z., Buffo, A., Podgórska, W., Marchisio, D.L. Droplet breakage and coalescence in liquid–liquid dispersions: Comparison of different kernels with EQMOM and QMOM (2017) *AIChE Journal*, 63 (6), pp. 2293-2311.
17. Buffo, A., Vanni, M., Marchisio, D.L. Simulation of a reacting gas–liquid bubbly flow with CFD and PBM: Validation with experiments (2017) *Applied Mathematical Modelling*, 44, pp. 43-60.
18. Lavino, A.D., Di Pasquale, N., Carbone, P., Marchisio, D.L. A novel multiscale model for the simulation of polymer flash nano-precipitation (2017) *Chemical Engineering Science*, 171, pp. 485-494.
19. Crevacore, E., Boccardo, G., Grillo, A., Marchisio, D.L., Sethi, R. Pore-scale simulations of particle transport for groundwater remediation: The effect of gravitational settling (2017) *Chemical Engineering Transactions*, 60, pp. 193-198.
20. Salenbauch, S., Sirignano, M., Marchisio, D.L., Pollack, M., D'Anna, A., Hasse, C. Detailed particle nucleation modeling in a sooting ethylene flame using a Conditional Quadrature Method of Moments (CQMOM) (2017) *Proceedings of the Combustion Institute*, 36 (1), pp. 771-779.
21. Crevacore, E., Tosco, T., Sethi, R., Boccardo, G., Marchisio, D.L. Recirculation zones induce non-Fickian transport in three-dimensional periodic porous media (2016) *Physical Review E*, 94 (5), art. no. 053118.
22. Buffo, A., De Bona, J., Vanni, M., Marchisio, D.L. Simplified volume-averaged models for liquid–liquid dispersions: Correct derivation and comparison with other approaches (2016) *Chemical Engineering Science*, 153, pp. 382-393.
23. Buffo, A., Vanni, M., Marchisio, D.L. On the implementation of moment transport equations in OpenFOAM: Boundedness and realizability (2016) *International Journal of Multiphase Flow*, 85, pp. 223-235.
24. Coletto, M.M., Marchisio, D.L., Barresi, A.A. A new segregation index for solid multicomponent mixtures (2016) *Powder Technology*, 299, pp. 77-86.
25. Buffo, A., Vanni, M., Renze, P., Marchisio, D.L. Empirical drag closure for polydisperse gas–liquid systems in bubbly flow regime: Bubble swarm and micro-scale turbulence (2016) *Chemical Engineering Research and Design*, 113, pp. 284-303.