

Dr IOANNIS SKARMOUTSOS

Born in Vasilika Fthiotidas, Greece

28 March 1977

Nationality: Greek

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iskarmoutsos@uoi.gr,
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ACADEMIC EDUCATION

- 2003-2006 Ph. D. in Physical Chemistry**, National and Kapodistrian **University of Athens**.
Department of Chemistry, Physical Chemistry Laboratory, Supervisor: Prof. Jannis Samios
“Theoretical-Computational investigation of the influence of thermodynamic parameters on the macroscopic, structural and dynamic properties of supercritical molecular systems”.
<https://phdtheses.ekt.gr/eadd/handle/10442/20859>
- 2003 M.Sc. in Physical Chemistry**, National and Kapodistrian **University of Athens**.
Department of Chemistry, Physical Chemistry Laboratory, Supervisor: Dr. Jannis Samios
“A statistical mechanical investigation of the properties of supercritical carbon dioxide and of the liquid mixture cis-trans N-Methylformamide via MD simulation techniques.”
- 2000 Diploma in Chemistry**, National and Kapodistrian University of Athens, Dpt of Chemistry.
Catalytic reactions with transition metal complexes in aqueous solutions.”
Scientific Advisor: Dr George Papadogiannakis (Laboratory of Industrial Chemistry)

WORKING EXPERIENCE

April 2022-	Assistant Professor (Tenure Track), University of Ioannina, Department of Chemistry Field: Theoretical Physical Chemistry
April 2019- April 2022	National Hellenic Research Foundation (NHRF) (In Collaboration with the Aristotle University of Thessaloniki (AUTH) and FORTH/ICE-HT) Duties: <i>Computational modeling of pillared graphene-based porous materials for energy storage applications</i>
January 2019- March 2019	HPC-Europa 3 Research Fellow, Universidad Pablo de Olavide, Seville, Spain Duties: <i>i) Computational modeling of mixtures of ionic liquids with organic battery electrolytes ii) MOF-based materials for the removal of organic contaminants from water.</i>
October 2016- December 2018	Research Associate, Institut Charles Gerhardt (UMR, CNRS 5253), University of Montpellier 2, Montpellier, France Duties: <i>Computational modeling of MOF-based materials for gas storage and separation.</i>
June 2016- July 2016	Visiting Researcher, Universitat Politècnica de Catalunya (UPC), Spain. Department of Physics Duties: <i>Computational modeling of the plastic phase of water.</i>

May 2016- June 2016	Visiting Researcher, University of Athens, Greece Department of Chemistry – Laboratory of Physical Chemistry Duties: <i>Computational modeling of liquids at elevated pressures.</i>
Apr. 2015- May 2016	Research Associate. University College London (UCL), London, UK. Department of Earth Sciences – Department of Physics and Astronomy Funding: ERC European project Duties: <i>Computational modeling of aqueous systems using ab initio molecular dynamics simulations.</i>
Oct. 2013- March 2015	Research Engineer (Ingénieur Chercheur). CEA, Grenoble, France. Institute for Nanoscience and Cryogenics (INAC-SPrAM) Funding: ANR ALIBABA project Duties: <i>Computational modeling of organic electrolytes with applications in battery technology.</i>
Sept.2012 – August 2013	Research Associate position at University of Crete, Greece. Materials Modeling & Design Group, Department of Chemistry Funding: THALES Research Project Duties: <i>Computational modeling of adsorption and separation of gas mixtures in carbon-based nanoporous materials.</i>
Apr.2012 – June 2012	HPC-Europa 2 Research Fellow. Univ. of Sassari, Italy. Funding: HPC-Europa European Union fellowship Duties: <i>To use the High Performance Computing facilities of the supercomputing center of CINECA to study the behavior of water and aqueous solutions by means of ab initio molecular dynamics simulations.</i>
Sept.2009 – Sept 2011	Research Associate, Imperial College, London, UK Dept. of Chemistry, Webpage: http://www.huntresearchgroup.org.uk Funding: ERC European Project. Duties: <i>Investigation of the thermodynamical, structural and dynamic properties of ionic liquids.</i>
Apr.2008 – Apr. 2009	Post-Doctoral research fellow, Univ. Politècnica de Catalunya (UPC), Spain. Computer Simulation in Cond. Matt. Res. Group, http://simcon.upc.edu/people Funding: Fellowship of the UPC Duties: <i>Investigation of local structural effects, characteristics of hydrogen bonding networks and related dynamics in supercritical fluids and liquid aqueous solutions.</i>

OBLIGATORY MILITARY SERVICE (FEBRUARY 2007 – FEBRUARY 2008)

SCIENTIFIC ACHIEVEMENTS

Google Scholar Profile: <https://scholar.google.com/citations?hl=en&user=CRCpodIAAAAJ>

Metrics Corresponding to 23-02-2025

● **53 Publications** in International Peer-Reviewed Journals (See Appendix)

- **h-index: 22** (source: Google Scholar)
- **i10-index: 34** (Source: Google Scholar)
- **1** publication in a scientific book (see Appendix)
- **4** Papers in conference proceedings (see Appendix)
- **2** Republished papers (see Appendix)
- **19** Invited Lectures (see Appendix)
- **19** Poster Presentations (see Appendix)
- **22** Talks in conferences (See Appendix)
- **4** Awarded scientific projects
- **1** Awarded Research Fellowship

AWARDED SCIENTIFIC PROJECTS (Before my election as Assistant Professor)

1. **HPC – Europa 2 project** to study the properties of water and aqueous solutions using ab initio molecular dynamics simulations.

Project awarded by the European Union (2012).

Principal Investigator: Dr Ioannis Skarmoutsos

Co-Investigator: Dr Marco Masia, University of Sassari, Department of Chemistry and Pharmacy.

2. **A joint experimental and theoretical study of supercritical mixtures.**

Project awarded by the Science and Technology Facilities Council (STFC), United Kingdom (UK) (2016).

Principal Investigator: Dr Sarantos Marinakis, Queen Mary University of London

Co-Investigator: Dr Ioannis Skarmoutsos

Co-Investigator: Professor Alan K. Soper, Science and Technology Facilities Council, UK

3. **FPMDCISTRANS: First Principles Molecular Dynamics studies of Cis- and Trans-N-Methylformamide liquid mixtures.**

Project Awarded by the Greek Research and Technology Network (GRNET) (2016).

Principal Investigator: Dr Ioannis Skarmoutsos

Principal Investigator: Professor Jannis Samios, University of Athens, Greece, Department of Chemistry, Director of Laboratory of Physical Chemistry

4. **HPC – Europa 3 project: “Ab Initio Molecular Dynamics of Mixtures of Ionic Liquids with Organic Carbonate Electrolytes “.** Project awarded by the European Union (2018).

Principal Investigator: Dr Ioannis Skarmoutsos

Co-Investigator: Professor Sofia Calero, Universidad Pablo de Olavide, Sevilla, Spain, Department of Physical, Chemical and Natural Systems.

AWARDED SCIENTIFIC PROJECTS (After my election as Assistant Professor)

Participation in European collaborative research projects as scientific partner

1. **MUSICODE project** (<https://musicode.eu/>), ” *An experimentally-validated multi-scale materials, process and device modeling & design platform enabling non-expert access to open innovation in the organic and large area electronics industry*”

European collaborative research and innovation project led by a multi-disciplinary consortium coordinated by University Ioannina. I started participating in the project on 2022.

Participation in Greek collaborative research projects as scientific partner

2. Emblematic actions in interdisciplinary scientific areas of special interest for the connection with the productive network. Research Project Name: “*7.1 Advanced Materials for Energy: Development of efficient third-generation photovoltaic materials and devices to enhance the competitiveness of the green energy production sector*” Grant Approved on 2023

<https://gsri.gov.gr/istoselides-enischyomenon-ergon/>

SCIENTIFIC TRAINING

2016 Annual CP2K-UK Users Meeting, <i>United Kingdom</i>	Third annual CP2K-UK users meeting organized by University College London, Kings College London and EPCC (EPSRC funded).
2013 Summer School <i>Italy</i>	International School of Physics “Enrico Fermi”: Water: Fundamental as a basis for understanding the environment and promoting technology.
2012 GPU workshop <i>Cyprus Institute</i>	Organized by the Computation-based Science and Tech. Research Centre (CaSToRC), in the framework of LinkSCEEM-2 and Cy-Tera projects.
2005 Summer School <i>Cardiff University- UK.</i>	CCP5 and Marie Curie Actions: Methods in Molecular Simulation Summer School, Advanced Courses in “ <i>First Principles Simulations</i> ”
2002 Winter School <i>Germany-Netherlands</i>	Quantum Simulations of Complex Many-Body Systems: From Theory to Algorithms, John von Neumann Institute for Computing (NIC)
2002 Summer School <i>Greece</i>	NATO Advanced Study Institute (ASI): Novel Approaches to the Structure and Dynamics of Liquids: Experiments, Theories and Simulations.

COMPUTING SKILLS

Programming	Fortran 77/90, Python, experience in Unix systems, Linux, Mac OS, Windows.
Computing	Use of High Performance Computing Systems (HPC Imperial College, Grace-UCL, Mare Nostrum BSC Barcelona, CINECA, Julich Germany, Titan-USA)
MD packages	Extensive knowledge of molecular simulation packages DL_POLY, Moscito, Moldy, Moliq-Dynamo and working experience with the ab initio codes CPMD, cp2k, Gaussian (GaussView) and Materials Studio.
MC packages	Very good working experience with the Monte-Carlo codes RASPA and CADSS.
Theoretical Chemistry Development	Development of utility software , for analysis of molecular trajectories of MD simulations (constructed for the Moscito, Moldy, DL_POLY, cp2k and CPMD, RASPA codes), to investigate structural and dynamic properties of fluid molecular systems.

AWARDED PERSONAL RESEARCH FELLOWSHIPS

- Universitat Politècnica de Catalunya (UPC), Research Fellowship, **Project:** “*Theoretical-computational investigation of the structural and dynamic properties of molecular systems in condensed phases via computer simulations.*”, Fellowship awarded by the Department of Physics and Nuclear Engineering, 2008

TEACHING EXPERIENCE

Before my appointment as assistant professor

Teaching in pre-graduate students (Chemistry-Pharmacy), Lab. of Phys. Chemistry, Athens (**2 academic years**).

Teaching in pre-graduate students (Chemistry), Theoretical Chemistry Laboratory (Imperial College London, UK) (**2 academic years**),

Teaching of application of Informatics in Chemistry in pre-graduate students, Department of Chemistry, University of Athens (**1 academic year**),

Teaching of statistical mechanics and molecular simulation techniques in post-graduate students, Department of Physics, Technical University of Catalonia (**1 academic year**)

After my appointment as assistant professor at the University of Ioannina

Teaching of the Physical Chemistry I course (Classical Thermodynamics) in pre-graduate students, Department of Chemistry, University of Ioannina (**Academic years 2022-2023, 2024-2025**)

Teaching of Informatics for Chemistry (Introduction to Python programming) in pre-graduate students, Department of Chemistry, University of Ioannina (**Academic years 2022-2023, 2023-2024, 2024-2025**)

Teaching at the Physical Chemistry Laboratory Course for pre-graduate students, Department of Chemistry, University of Ioannina, (**Academic years 2022-2023, 2023-2024, 2024-2025**)

Introduction of Computational Physical Chemistry Lab Exercises in the Physical Chemistry Laboratory Course for pre-graduate students (**Academic years 2023-2024, 2024-2025**)

Participation in the teaching of the Optional (Elective) Course: “Concepts of Chemistry” for 4th year students of the department of Chemistry (**Academic year 2023-2024**)

Official Academic Supervisor of pre- and post-graduate students:

Supervisor of the Master Thesis of 1 MSc student (Mr Ilias Karvounis, Department of Chemistry, University of Ioannina, Thesis Defended: September 2024)

Supervisor of the Diploma Thesis of 7 pre-graduate students (4 Diploma theses defended, 3 in progress).

Official Member of the 3-member advisory committee for 2 PhD Candidates

Official Member of the 7-member advisory committee for 3 PhD Candidates

Official Member of the 3-member advisory committee for 1 MSc Candidate.

Co- Supervisor of 1 post-doctoral researcher (in the framework of the Musicode project, <https://musicode.eu/>), co-supervised with Prof. Eleftherios Lidorikis)

OTHER SCIENTIFIC ACTIVITIES

Reviewer for 32 international scientific journals: *Journal of Physical Chemistry, Chemical Physics Letters, Chemical Physics, Physical Chemistry Chemical Physics, Fuel, Journal of the Brazilian Chemical Society, Journal of Molecular Liquids, International Journal of Modern Physics B, International Journal of Heat and Mass Transfer, Microporous and Mesoporous Materials, Advanced Theory and Simulations, Journal of Molecular Graphics and Modelling, New Journal of Chemistry, ACS Energy Letters, Fluid Phase Equilibria, Journal of Industrial and Engineering Chemistry, Journal of Photochemistry and Photobiology A: Chemistry, Journal of Chemical Information and Modeling, Chinese Physics B, Entropy, Applied Surface Science, Journal of Inorganic and Organometallic Polymers and Materials, Applied Thermal Engineering, Journal of CO₂ Utilization, Frontiers in Physics-Condensed Matter Physics, Materials Today Communications, ACS Omega, Langmuir, Computational and Theoretical Chemistry, Separation and Purification Technology, Physical Review Research, Scientific Reports*

Reviewer for the Greek Research & Technology Network (GRNET) High Performance Computing Services

Reviewer for the French National Research Agency (ANR, <https://anr.fr/en/>) to evaluate research proposals submitted for funding.

Member of the *European Molecular Liquids Group* (EMLG) .

Member of the editorial board of the *Journal of Theoretical Chemistry* (Hindawi Publishing Corporation) from 28-11-2012 to 23-07-2017.

Member of the editorial board of *Frontiers in Physics, Condensed Matter Physics* (Frontiers Media, Lausanne, Switzerland) from 07-11-2020.

LANGUAGES

Greek (native), English (fluent, working experience), Spanish (very good knowledge, working experience), French (higher intermediate level, working experience)

REFERENCES

References may be asked from the following academics:

Prof. Jannis Samios

National & Kapodistrian University of Athens, Department of Chemistry, Physical Chemistry Laboratory, Panepistimiopolis 15771, Athens, Greece

Email: isamios@chem.uoa.gr

Prof. Elvira Guardia

Departament de Física i Enginyeria Nuclear, Universitat Politècnica de Catalunya, Edifici B4-B5, Campus Nord, Jordi Girona 1-3, 08034 Barcelona, Spain

Email: elvira.guardia@upc.edu

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Dr Stefano Mossa

INAC/SPrAM (UMR 5819 UJF, CNRS, CEA), CEA-Grenoble, 17 Rue des Martyrs, 38054 Grenoble, France

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Prof. Guillaume Maurin

Institut Charles Gerhardt (UMR, CNRS 5253), University of Montpellier 2, Montpellier, France

Email: guillaume.maurin@univ-montp2.fr

Prof. Giancarlo Franzese

Departament de Física de la Matèria Condensada & Institut de Nanociència i Nanotecnologia (IN2UB), Universitat de Barcelona

Email: gfranzese@ub.edu

Web page : <https://spcmub.wordpress.com/>

Prof. George E. Froudakis

Department of Chemistry, University of Crete, Voutes, Heraklion, 71003 Crete, Greece

Email: frudakis@chemistry.uoc.gr

Dr Patricia Hunt

Department of Chemistry, Exhibition Rd, Imperial College of Science, Technology and Medicine, South Kensington, London, SW7 2AZ, United Kingdom

Email: p.hunt@imperial.ac.uk

Web page : <http://www.huntresearchgroup.org.uk/index.html>

Prof. Tom Welton

Dean of the Faculty of Natural Sciences, Imperial College of Science, Technology and Medicine, Exhibition Rd, South Kensington, London, SW7 2AZ, United Kingdom

Email: t.welton@imperial.ac.uk

Dr Marco Masia

Former Assistant Professor of Theoretical Chemistry, Dipartimento di Chimica, Università degli studi di Sassari, Via Vienna 2, 07100, Sassari, Italy

Currently: Head of Entrepreneurship, University of Vienna, Austria

Email: marco.masia@univie.ac.at

APPENDIX

Publications in International Journals with referees

53) **Solvation structure and dynamics of the thiocyanate anion in mixed N,N-Dimethylformamide-water solvents: A molecular dynamics approach.**

Ioannis Skarmoutsos and Ilias G. Karvounis, *ChemPhysChem* <https://doi.org/10.1002/cphc.202400732> (2024)

52) **A comprehensive molecular dynamics simulation of plastic and liquid succinonitrile: Structural, dynamic, and dielectric properties.**

Manel Canales, Ioannis Skarmoutsos and Elvira Guardia, *J. Chem. Phys.* **161**, 174503 (2024)

51) **Substantial breakdown of the hydrogen-bonding network, local density inhomogeneities and fluid-liquid structural transitions in supercritical octanol-1: A molecular dynamics investigation.**

Ioannis Skarmoutsos, *J. Chem. Phys.* **161**, 044506 (2024)

50) **Pillared graphene oxide frameworks for the adsorption and separation of polar protic and aprotic liquid solvents: The cases of pure water, methanol, dimethyl sulfoxide, and dimethyl sulfoxide–water mixtures.**

Ioannis Skarmoutsos, Emmanuel N. Koukaras and Emmanuel Klontzas, *J. Chem. Phys.* **160**, 184701 (2024)

49) **Solvation structure and dynamics of favipiravir in supercritical CO₂. A molecular dynamics investigation.**

Ioannis Skarmoutsos and Jannis Samios, *J. Mol. Liq.* **392**, 123512 (2023)

48) **Hydration structure and dynamics, ultraviolet–visible and fluorescence spectra of caffeine in ambient liquid water. A combined classical molecular dynamics and quantum chemical study.**

Ioannis Skarmoutsos, Demeter Tzeli and Ioannis D. Petsalakis, *J. Mol. Liq.* **391**, 123220 (2023)

47) **Local intermolecular structure, hydrogen bonding and related dynamics in the liquid cis/trans N-methylformamide mixture: A density functional theory based Born-Oppenheimer molecular dynamics study.**

Ioannis Skarmoutsos, Ricardo L. Mancera, Stefano Mossa and Jannis Samios, *J. Mol. Liq.* **365**, 120085 (2022)

46) **Fingerprints of the Crossing of the Frenkel and Melting Line on the Properties of High-Pressure Supercritical Water.**

Ioannis Skarmoutsos, Jannis Samios and Elvira Guardia *J. Phys. Chem. Lett.* **13**, 7636 (2022)

45) **Using Car-Parrinello simulations and microscopic order descriptors to reveal two locally favored structures with distinct molecular dipole moments and dynamics in ambient liquid water.**

Ioannis Skarmoutsos, Giancarlo Franzese and Elvira Guardia, *J. Mol. Liq.* **364**, 119936 (2022)

44) **A Computational Study on Phenylidiboronic Acid-Pillared Graphene Oxide Frameworks for Gas Storage and Separation.**

Ioannis Skarmoutsos, Emmanuel N. Koukaras, and Emmanuel Klontzas, *ACS Appl. Nano Mater.* **5**, 9286 (2022)

43) **CF₄ Capture and Separation of CF₄–SF₆ and CF₄–N₂ Fluid Mixtures Using Selected Carbon Nanoporous Materials and Metal–Organic Frameworks: A Computational Study.**

Ioannis Skarmoutsos, Emmanuel N. Koukaras, and Emmanuel Klontzas, *ACS Omega* **7**, 6691 (2022)

42) **On the Different Faces of the Supercritical Phase of Water at a Near-Critical Temperature: Pressure-Induced Structural Transitions Ranging from a Gaslike Fluid to a Plastic Crystal Polymorph**

Ioannis Skarmoutsos, Andres Henao, Elvira Guardia and Jannis Samios, *J. Phys. Chem. B* **125**, 10260 (2021)

41) The Polar Cosolvent Effect on Caffeine Solvation in Supercritical CO₂–Ethanol Mixtures: A Molecular Modeling Approach.

Ioannis Skarmoutsos, Ioannis D. Petsalakis and Jannis Samios, *Ind. Eng. Chem. Res.* **60**, 11834 (2021)

40) The Impact of Ionic Liquid Loading in Three-Dimensional Carbon Nanotube Networks on the Separation of CO₂/CH₄ Fluid Mixtures: Insights from Molecular Simulations.

Ioannis Skarmoutsos, Emmanuel N. Koukaras, and Emmanuel Klontzas, *J. Phys. Chem. C* **125**, 13508 (2021)

39) Confinement effects on the properties of polar hydrogen-bonded fluids: A showcase on methanol adsorbed in three-dimensional pillared graphene and carbon nanotube networks.

Ioannis Skarmoutsos, Emmanuel N. Koukaras, George E. Froudakis, Guillaume Maurin and Emmanuel Klontzas, *J. Phys. Chem. C* **124**, 22959 (2020)

38) Hydration structure and dynamics of the favipiravir antiviral drug: A molecular modelling approach.

Ioannis Skarmoutsos, Guillaume Maurin, Elvira Guardia and Jannis Samios, *Bull. Chem. Soc. Jpn.* **93**, 1378 (2020)

37) Porous carbon nanotube networks and pillared graphene materials exhibiting high SF₆ adsorption uptake and separation selectivity of SF₆/N₂ fluid mixtures: A comparative molecular simulation study.

Ioannis Skarmoutsos, Emmanuel N. Koukaras, Costas Galiotis, George Froudakis, Emmanuel Klontzas *Micropor. Mesopor. Mat.* **307**, 110464 (2020)

36) Solvation Structure and Dynamics of the Dimethylammonium Cation Diluted in Liquid Water: A Molecular Dynamics Approach

Ioannis Skarmoutsos and Elvira Guardia *J. Chem. Phys.* **152**, 234501 (2020)

35) Highly efficient rare-earth based metal–organic frameworks for water adsorption: A molecular modelling approach.

Ioannis Skarmoutsos, Mohamed Eddaoudi and Guillaume Maurin (2019) *J. Phys. Chem. C* **123**, 26989 (2019)

34) A study of Ar-N₂ supercritical mixtures using neutron scattering, molecular dynamics simulations and quantum mechanical scattering calculations.

Alan K. Soper, Ioannis Skarmoutsos, Jacek Klos, Jannis Samios and Sarantos Marinakis (2019) *J. Mol. Liq.* **290**, 111168 (2019)

33) On the interplay between the local structure and dynamics in low concentration mixtures of H₂O and HOD in the [Emim⁺][TF₂N⁻] room temperature ionic liquid.

Ioannis Skarmoutsos, Leonidas Spyrogiannopoulos, Emmanouil Kainourgiakis and Jannis Samios *J. Mol. Liq.* **289**, 111135 (2019)

32) The effect of polymorphism on the structural, dynamic and dielectric properties of plastic crystal water: A molecular dynamics simulation perspective.

Ioannis Skarmoutsos, Stefano Mossa and Elvira Guardia (2019) *J. Chem. Phys.* **150**, 124506 (2019)

31) Highly tunable sulfur hexafluoride separation by interpenetration control in metal organic frameworks.

Ioannis Skarmoutsos, Mohamed Eddaoudi and Guillaume Maurin, *Micropor. Mesopor. Mat.* **281**, 44 (2019)

30) Solvent and Salt Effect on Lithium Ion Solvation and Contact Ion Pair Formation in Organic Carbonates: A Quantum Chemical Perspective.

Veerapandian Ponnuchamy, Stefano Mossa and Ioannis Skarmoutsos *J. Phys. Chem. C* **122**, 25930 (2018)

29) Peculiar Molecular Shape and Size Dependence of the Dynamics of Fluids confined in a Small-Pore Metal-Organic Framework.

Ioannis Skarmoutsos, Mohamed Eddaoudi and Guillaume Maurin, *J. Phys. Chem. Lett.* **9**, 3014 (2018)

- 28) **CO₂ capture using the SIFSIX-2-Cu-i metal-organic framework: A computational approach.**
Ioannis Skarmoutsos, Youssef Belmabkhout, Karim Adil, Mohamed Eddaoudi and Guillaume Maurin, *J. Phys. Chem. C*, **121**, 27462 (2017)
- 27) **Local structural fluctuations, hydrogen bonding and structural transitions in supercritical water.**
Ioannis Skarmoutsos, Elvira Guardia and Jannis Samios, *J. Supercrit. Fluids* **130**, 156 (2017)
- 26) **Local Structure and Translational Dynamics of NMF (N-Methylformamide)– DMF (N, N-Dimethylformamide) Mixtures via Molecular Dynamics Simulation.**
Nikolaos Elpidoforou, Ioannis Skarmoutsos, Emmanuel Kainourgiakis, Vasilios Raptis and Jannis Samios, *J. Mol. Liq.*, **226**, 16 (2017)
- 25) **Structure and dynamics of liquid CS₂: Going from ambient to elevated pressure conditions.**
Ioannis Skarmoutsos, Stefano Mossa and Jannis Samios, *J. Chem. Phys.*, **145**, 154505 (2016)
- 24) **The Anion Effect on Li⁺ Ion Coordination Structure in Ethylene Carbonate Solutions**
Bo Jiang, Veerapandian Ponnuchamy, Yuneng Shen, Xueming Yang, Kaijun Yuan, Valentina Vetere, Stefano Mossa, Ioannis Skarmoutsos, Yufan Zhang and Junrong Zheng, *J. Phys. Chem. Lett.*, **7**, 3554 (2016)
- 23) **Highly selective separation and adsorption-induced phase transition of SF₆-N₂ fluid mixtures in three-dimensional carbon nanotube networks.**
Ioannis Skarmoutsos, George Tamiolakis and George E. Froudakis *J. Supercrit. Fluids* **113**, 89 (2016)
- 22) **Structural and dipolar fluctuations in liquid water: A Car-Parrinello molecular dynamics study.**
Ioannis Skarmoutsos, Elvira Guardia and Marco Masia *Chem. Phys. Lett.*, **648**, 102 (2016)
- 21) **Li⁺ solvation in pure, binary and ternary mixtures of organic carbonate electrolytes.**
Ioannis Skarmoutsos, Veerapandian Ponnuchamy, Valentina Vetere and Stefano Mossa *J. Phys. Chem. C*, **119**, 4502 (2015)
- 20) **Hydrogen bonding and related properties in liquid water: A Car-Parrinello molecular dynamics simulation study.**
Elvira Guardia, Ioannis Skarmoutsos and Marco Masia *J. Phys. Chem. B*, **119**, 8926 (2015)
- 19) **The Importance of Timescale for Hydrogen Bonding in Imidazolium Chloride Ionic Liquids.**
Ioannis Skarmoutsos, Tom Welton and Patricia A. Hunt *Phys. Chem. Chem. Phys.*, **16**, 3675 (2014)
- 18) **Separation of CO₂ / N₂ mixtures in 3D carbon-based porous nanotube networks: A molecular dynamics investigation.**
Ioannis Skarmoutsos, George Tamiolakis and George E. Froudakis, *Phys. Chem. Chem. Phys.*, **16**, 876 (2014)
- 17) **Carbon-based nanoporous networks as media for the separation of CO₂/CH₄ mixtures: A molecular dynamics approach.**
Ioannis Skarmoutsos, George Tamiolakis and George E. Froudakis, *J. Phys. Chem. C*, **117**, 19373 (2013)
- 16) **Hydrogen Bonding in 1-Butyl- and 1-Ethyl-3-Methylimidazolium Chloride Ionic Liquids**
Ioannis Skarmoutsos, Dimitris Dellis, Richard P. Matthews, Tom Welton and Patricia A. Hunt, *J. Phys. Chem. B*, **116**, 4921 (2012)
- 15) **Solvation structure and Dynamics of cis- and trans- 1,2 Dichloroethene Isomers in Supercritical Carbon Dioxide. A molecular dynamics simulation study.**
Dimitris Dellis, Ioannis Skarmoutsos and Jannis Samios, *J. Phys. Chem. B*, **115**, 12098 (2011)
- 14) **Structural and dynamic properties of the new alternative refrigerant 2,3,3,3-Tetrafluoro-1-propene (HFO-1234yf) in the liquid state.**
Ioannis Skarmoutsos and Patricia A. Hunt, *J. Phys. Chem. B*, **114**, 17120 (2010)

13) **Hydrogen bond, electron donor acceptor dimer and residence dynamics in supercritical CO₂-ethanol mixtures and the effect of hydrogen bonding on single reorientational and translational dynamics. A molecular dynamics simulation study.**

Ioannis Skarmoutsos, Elvira Guardia and Jannis Samios, *J. Chem. Phys.*, **133**, 014504 (2010)

12) **Effect of the local hydrogen bonding network on the reorientational and translational dynamics in supercritical water.**

Ioannis Skarmoutsos and Elvira Guardia, *J. Chem. Phys.*, **132**, 074502 (2010)

11) **Molecular simulations of benzene and hexafluorobenzene using new optimized effective potential models: Investigation of the liquid, vapor-liquid coexistence and supercritical fluid phases.**

Dimitris Dellis, Ioannis Skarmoutsos and Jannis Samios, *J. Mol. Liq.*, **153**, 25 (2010)

10) **On ion and molecular polarization of halides in water.**

Elvira Guardia, Ioannis Skarmoutsos and Marco Masia, *J. Chem. Theory Comput.*, **5**, 1449 (2009)

9) **Local structural effects and related dynamics in supercritical ethanol. 2. Hydrogen bonding network and its effect on single reorientational dynamics.**

Ioannis Skarmoutsos and Elvira Guardia, *J. Phys. Chem. B*, **113**, 8898 (2009)

8) **Local structural effects and related dynamics in supercritical ethanol. 1. Mechanisms of local density reorganization and residence dynamics.**

Ioannis Skarmoutsos and Elvira Guardia, *J. Phys. Chem. B*, **113**, 8887 (2009)

7) **The effect of intermolecular interactions on local density inhomogeneities and related properties in pure supercritical molecular fluids. A comparative molecular dynamics study.**

Ioannis Skarmoutsos, Dimitris Dellis and Jannis Samios, *J. Phys. Chem. B*, **113**, 2783 (2009)

6) **Investigation of the local composition enhancement and related dynamics in supercritical CO₂-cosolvent mixtures via computer simulation. The case of ethanol in CO₂.**

Ioannis Skarmoutsos, Dimitris Dellis and Jannis Samios, *J. Chem. Phys.*, **126**, 224503 (2007)

5) **Local Density Augmentation and Dynamic Properties of Hydrogen- and non Hydrogen- Bonded Supercritical Fluids: A Molecular Dynamics Study**

Ioannis Skarmoutsos and Jannis Samios, *J. Chem. Phys.*, **126**, 044503 (2007)

4) **Local Density Inhomogeneities and Dynamics in Supercritical Water: A molecular dynamics simulation approach.**

Ioannis Skarmoutsos and Jannis Samios, *J. Phys. Chem. B*, **110**, 21931 (2006)

3) **Local intermolecular structure and dynamics in binary supercritical solutions. A molecular dynamics simulation study of methane in carbon dioxide.**

Ioannis Skarmoutsos and Jannis Samios, *J. Mol. Liq.*, **125**, 181-186 (2006)

2) **Investigation of the vapor-liquid equilibrium and supercritical phase of pure methane via computer simulations.**

Ioannis Skarmoutsos, Leonidas I. Kampanakis and Jannis Samios, *J. Mol. Liq.*, **117**, 33-41 (2005)

1) **Molecular dynamics of cis/trans N-methylformamide (NMF) liquid mixture using an all atom optimized rigid force field**

Ioannis Skarmoutsos and Jannis Samios, *Chem. Phys. Lett.*, **384**, 108-113 (2004)

Republished Papers:

1) **Local Density Augmentation and Dynamic Properties of Hydrogen- and non Hydrogen- Bonded Supercritical Fluids: A Molecular Dynamics Study**

Ioannis Skarmoutsos and Jannis Samios, **republished in the** *Virtual Journal of Biological Physics Research*, **13 (Issue 3)** (2007)

2) **Investigation of the local composition enhancement and related dynamics in supercritical CO₂-cosolvent mixtures via computer simulation. The case of ethanol in CO₂.**

Ioannis Skarmoutsos Dimitris Dellis and Jannis Samios, **republished in the *Virtual Journal of Biological Physics Research*, 13 (Issue 12) (2007)**

Publications in Scientific Books:

1) **Molecular Dynamics Simulation of cis-trans N-Methylformamide (NMF) liquid mixture. Structure and Dynamics.**

Ioannis Skarmoutsos and Jannis Samios, *Lecture Series on Computer and Computational Sciences* (VSP International), **1**, 479 (2004)

Papers in Conference Proceedings:

1) **Molecular Dynamics simulation studies of supercritical carbon dioxide using available potential models. Investigation of the bulk thermodynamical, transport and dynamical properties.**

Ioannis Skarmoutsos and Jannis Samios, Winter School: Quantum Simulations of Complex Many-Body Systems: From Theory to Algorithms, *Publication Series of the John von Neumann Institute for Computing (NIC Series)*, **Vol. 11**, 20, (2002)

2) **Molecular dynamics studies of cis/trans N-methylformamide (NMF) liquid mixture using a new optimized all atom force field.**

Ioannis Skarmoutsos and Jannis Samios, NATO Advanced Study Institute (ASI): Novel Approaches to the Structure and Dynamics of Liquids: Experiments, Theories and Simulations. page 151 (2002)

3) **Investigation of the Local Density Inhomogeneities and Dynamics in Neat Supercritical Fluids using MD simulation techniques: Comparison between Hydrogen- and Non Hydrogen-Bonded fluids.**

Ioannis Skarmoutsos Nikolaos Elpidoforou and Jannis Samios, Winter School: Computational Nanoscience: Do It Yourself, *Publication Series of the John von Neumann Institute for Computing (NIC Series)*, page 11, (2006)

4) **Computational Modelling of Nanoporous Materials for Sustainable Energy and Environmental Applications.**

Ioannis Skarmoutsos, Emmanuel Klontzas, Emmanuel N. Koukaras and Guillaume Maurin, *Conference Proceedings, 12th Panhellenic Conference of Chemical Engineering (2019)*.

PRESENTATIONS

INVITED LECTURES

19) **Molecular Simulations: A computational toolbox to predict, understand and rationally tailor the physicochemical properties of liquid and supercritical solvents and materials for sustainable chemical applications.**

Ioannis Skarmoutsos, Invited Lecture at the Institute of Physics, Polish Academy of Sciences, Warsaw, Poland (2024)

18) Using classical and ab initio molecular dynamics, statistical mechanical theories and structural order parameters to study the phase-behavior of molecular systems: The unique case of water at ambient-pressure liquid up to extreme-pressure supercritical thermodynamic conditions.

Ioannis Skarmoutsos, Invited Lecture at the Department of Materials Science and Engineering, University of Ioannina, Workshop on Computational Materials Science organized by Department of Materials Science and Engineering, University of Ioannina and the Hellenic Society for the Science and Technology of Condensed Matter (HSSTCM).

17) Molecular Simulations: A computational physical chemistry toolbox to predict, understand and rationally tailor the properties of molecular/ionic solvents and materials for sustainable chemical applications

Ioannis Skarmoutsos, Invited Lecture at the Chemistry Department, University of Athens, Greece (2021)

16) Molecular modelling and simulation methods to predict and understand the properties of condensed matter: A physicochemical approach to develop modern chemical, environmental and engineering applications

Ioannis Skarmoutsos, Invited Lecture at the Chemistry Department, University of Ioannina, Greece (2020)

15) Understanding the properties of aqueous systems and their interactions with nanoporous materials and sustainable ionic liquid solvents to develop environmental, energy storage and geological applications: A computational modelling perspective.

Ioannis Skarmoutsos, Invited Lecture at the National Hellenic Research Foundation, Athens, Greece (2019).

14) Computational Modeling of liquid electrolytes and porous materials for sustainable energy applications.

Ioannis Skarmoutsos, Invited Lecture at the Department of Physical, Chemical, and Natural Systems, Universidad Pablo de Olavide, Seville, Spain (2019).

13) Computational modeling of modern electrolytes for energy applications

Ioannis Skarmoutsos, Invited Lecture at the Department of Chemistry of the École Normale Supérieure de Lyon, France (2018).

12) Gas adsorption and separation using metal-organic frameworks: A computational approach.

Ioannis Skarmoutsos, Invited Lecture at the Department of Physics, Technical University of Catalonia (UPC), Spain (2018)

11) Computational modeling of modern electrolytes for energy applications: Structure and dynamics in ionic liquids and their mixtures with water.

Ioannis Skarmoutsos, Invited Lecture at the Department of Physics, University of Barcelona (UB), Spain (2018)

10) Gas adsorption and separation using the SIFSIX-2-Cu-i and SIFSIX-2-Cu metal-organic frameworks: A computational approach.

Ioannis Skarmoutsos, Invited Lecture at the Physical Chemistry Laboratory, Department of Chemistry, National and Kapodistrian University of Athens, Greece (2018)

9) Local structural and dipolar fluctuations in liquid water: Insights from ab initio molecular dynamics simulations.

Ioannis Skarmoutsos, Invited lecture at the University of Barcelona (UB) - Physics Department, Barcelona, Catalonia, Spain (2016)

8) Local structural and dipolar fluctuations in liquid water: Insights from ab initio molecular dynamics simulations.

Ioannis Skarmoutsos, Invited lecture at the Institut Laue-Langevin (ILL), Grenoble, France (2016)

7) Hydrogen Bonding Interactions in Ionic Liquids: Insights from molecular simulation.

Ioannis Skarmoutsos, Invited lecture at INAC/SPrAM, CEA-Grenoble, France (2013)

6) Hydrogen Bonding interactions and their effect on the properties of self-associating liquids: Insights from molecular simulations.

Ioannis Skarmoutsos, Invited lecture at the University of Crete, Department of Materials Science and Technology, Iraklio, Greece (2012)

5) Molecular Dynamics Simulation Techniques: A very useful tool to investigate the properties of condensed matter.

Ioannis Skarmoutsos, Invited lecture at the University of Crete, Department of Chemistry, Iraklio, Greece (2012)

4) Hydrogen Bonding interactions and their effect on the properties of self-associating liquids: Insights from molecular simulations.

Ioannis Skarmoutsos, Invited lecture at the University of Sassari, Department of Chemistry and Pharmacy, Sassari, Italy (2012)

3) Molecular Dynamics Simulation: A useful tool to investigate the properties of gases confined in nanoporous materials.

Ioannis Skarmoutsos, Invited lecture at the *European Technical School on Hydrogen and Fuel Cells*, Heraklion, Greece (2012).

2) Supercritical Fluids: An introduction to the physicochemical properties of a “peculiar” state of matter.

Ioannis Skarmoutsos, Invited Lecture at the Department of Chemical Engineering, Aristotle University of Thessaloniki, Greece (2011)

1) Supercritical Fluids: Investigation of local density inhomogeneities and related properties using computer simulations.

Ioannis Skarmoutsos, Invited lecture at the “Day of the research” of the Department of Physics & Nuclear Engineering (Technical University of Catalonia- UPC), Barcelona, Spain (2009)

TALKS IN CONFERENCES

22) The Unique Structural Features of Water, Ranging from Ambient Liquid up to Supercritical, Extreme-Pressure Conditions: Insights from Classical and ab initio Molecular Dynamics Simulations.

23rd Panhellenic Chemistry Conference, Athens-Greece (2024)

21) Pillared graphene oxide frameworks for the adsorption and separation of dimethylsulfoxide-water mixtures.

European/Japanese Molecular Liquids Group-EMLG/JMLG Conference entitled: *"Structure and Dynamics of Hydrogen-Bonded Systems."* in Trieste- Italy (2024)

20) The unique structural features of water, ranging from ambient liquid up to supercritical, extreme-pressure conditions: Insights from classical and ab initio molecular dynamics simulations.

2nd International Conference on Physical Chemistry and Spectroscopy (ICPCS), Patras, Greece (2023)

19) Solvation structure and dynamics of pharmaceutical compounds in liquid and supercritical solvents: The cases of caffeine and favipiravir.

European/Japanese Molecular Liquids Group-EMLG/JMLG Conference entitled: "Understanding Solvation in Molecular and Ionic Fluids: Towards a Sustainable Future." in Bordeaux- France (2023)

18) The unique structural features of water, ranging from ambient liquid up to supercritical, extreme-pressure conditions: Insights from classical and ab initio molecular dynamics simulations.

European Conference of Computational & Theoretical Chemistry, EuChems CompChem in Thessaloniki, Greece (2023)

17) The unique structural features of water, ranging from ambient liquid up to supercritical, extreme-pressure conditions: Insights from classical and ab initio molecular dynamics simulations.

European/Japanese Molecular Liquids Group-EMLG/JMLG Conference entitled: "Molecular liquids at interfaces." in Barcelona- Spain (2022)

16) Understanding the interactions of water with nanoporous materials to develop environmental 1) Gas storage and separation using three-dimensional pillared graphene oxide frameworks with phenyldiboronic acid linkers: A computational approach.

Graphene 2021 Conference, Grenoble, France (2021)

15) SF₆ Capture and Separation of SF₆-N₂ Fluid Mixtures Using Porous Carbon Nanotube Networks and Pillared Graphene Materials

35th Panhellenic Conference on Solid State Physics and Materials Science, Athens, Greece (2021)

14) The different faces of the supercritical phase of water at a near-critical temperature.

1st International Conference on Physical Chemistry and Spectroscopy (ICPCS), Patras, Greece (2021)

13) Understanding the interactions of water with nanoporous materials to develop environmental applications: The case of water adsorption in metal-organic frameworks.

CECAM Workshop: WaterEurope: Multiscale simulations and coarse-grained models for water and aqueous systems, CECAM Headquarters, EPFL, Lausanne, Switzerland (2019)

12) Computational Modelling of Nanoporous Materials for Sustainable Energy and Environmental Applications.

12th Panhellenic Conference of Chemical Engineering, Athens-Greece (2019)

11) On the Peculiar Molecular Shape and Size Dependence of the Dynamics of Fluids Confined in a Small-Pore Metal-Organic Framework

Energy Landscapes 2018 Conference (organized by the University of Cambridge), Kalamata, Greece (2018)

10) Local structural fluctuations, hydrogen bonding and structural transitions in supercritical water.

WaterSpain 2017 Conference, The Zaragoza Scientific Center for Advanced Modeling (ZCAM), Zaragoza, Spain (2017)

9) Local structural and dipolar fluctuations in liquid water: Insights from ab initio molecular dynamics simulations.

European Molecular Liquids Group-EMLG Conference entitled: "Recent Progresses on the Experimental & Theoretical-Computational Techniques for the Study of Liquids and Supercritical Fluids." in Crete-Greece (2016)

8) Computational modeling of battery electrolytes: insights from molecular simulations.

European Molecular Liquids Group-EMLG Conference entitled: "Molecular Liquids and Soft Matter: From Fundamentals to Applications." in Roma Tre University- Italy (2014)

7) Local Structural Inhomogeneities, Hydrogen Bonding and Tetrahedral Structure in Supercritical Water: Insights from Molecular Simulations.

International School of Physics “Enrico Fermi”, Varenna, Italy (2013)

6) Local density inhomogeneities and related properties in supercritical fluids. A computer simulation study.

Ioannis Skarmoutsos, Dimitris Dellis* and Jannis Samios, European Molecular Liquids Group-EMLG Conference entitled: "Understanding solvation from liquid to supercritical conditions." in Lisboa- Portugal (2008)

5) MD Simulation of cis-trans N-Methylformamide (NMF) liquid mixture. Structure and Dynamics.

Ioannis Skarmoutsos Ricardo L. Mancera and Jannis Samios, CCP5 and Marie Curie Actions: Methods in Molecular Simulation Summer School, Cardiff University- UK (2005)

4) Molecular Dynamics Simulation of cis-trans N-Methylformamide (NMF) liquid mixture. Structure and Dynamics.

Ioannis Skarmoutsos, Ricardo L. Mancera and Jannis Samios, International Conference of Computational Methods in Sciences and Engineering (ICCMSE), organized by the **European Society of Computational Methods in Sciences and Engineering (ESCMSE)**, Athens-Greece (2004)

3) Is the hydrogen-bonding network around the cis and trans N-Methylformamide (NMF) conformers responsible for the observation of different diffusion coefficients? A temperature dependent molecular dynamics approach.

Ioannis Skarmoutsos, Ricardo L. Mancera and Jannis Samios, European Molecular Liquids Group-EMLG Conference entitled: "Complex Liquids: Fundamental Properties to Industrial Applications." in Sheffield Hallam University- UK (2004)

2) Molecular Dynamics simulations of cis-trans N-Methylformamide liquid mixture. Structure, dynamics and hydrogen bonding analysis.

Ioannis Skarmoutsos and Jannis Samios, European Molecular Liquids Group-EMLG Conference entitled: "Molecular Liquids. Routes from Local Order to Large-Scale Cooperativity." in Castelvechio Pascoli- Italy (2003)

1) Molecular dynamics studies of cis/trans N-methylformamide (NMF) liquid mixture using a new optimized all atom force field.

Ioannis Skarmoutsos and Jannis Samios, NATO Advanced Study Institute (ASI): Novel Approaches to the Structure and Dynamics of Liquids: Experiments, Theories and Simulations. Rhodes- Greece (2002)

POSTER PRESENTATIONS

19) Fingerprints of the crossing of the Frenkel and Melting lines in high-pressure supercritical water.

Elvira Guardia, Ioannis Skarmoutsos, Jannis Samios

European/Japanese Molecular Liquids Group-EMLG/JMLG Conference entitled: "Molecular liquids at interfaces." in Barcelona- Spain (2022)

18) Confined Ionic Liquids (IL) in carbon-based materials for CO₂/CH₄ separation: insights from computer simulations.

Ioannis Skarmoutsos, Emmanuel Klontzas, Emmanuel N. Koukaras

Graphene 2021 Conference, Grenoble, France (2021)

17) F-gas adsorption in pillared graphene materials. Insights from molecular simulations.

Ioannis Skarmoutsos, Emmanuel Klontzas, Emmanuel N. Koukaras

Graphene 2021 Conference, Grenoble, France (2021)

16) Role of the Loading of Ionic Liquid [EMIM]⁺[BF₄]⁻ on the Separation of CO₂/CH₄ in 3D Carbon Nanotube Networks

I. Skarmoutsos, R. Lingas, E.N. Koukaras, E. Klontzas

35th Panhellenic Conference on Solid State Physics and Materials Science, Athens, Greece (2021)

15) Molecular Dynamics Study of H₂O and HOD dissolved in the Room Temperature Ionic Liquid Emim⁺ Tf₂N⁻: Dynamical and Structural properties

Leonidas Spyrogiannopoulos, Emmanouil Kainourgiakis, Ioannis Skarmoutsos and Jannis Samios

Athens Conference of Advances in Chemistry (acac2018), organised on the occasion of the centennial anniversary of the Chemistry Department of the National and Kapodistrian University of Athens.

14) CO₂ capture and separation from methane using the SIFSIX-2-Cu-i metal-organic framework: A computational approach.

Ioannis Skarmoutsos, Youssef Belmabkhout, Karim Adil, Mohamed Eddaoudi and Guillaume Maurin

11th International Symposium on the Characterization of Porous Solids (COPS-XI), Avignon-France (2017)

13) Structure and dynamics of liquid CS₂: Going from ambient to elevated pressure conditions.

Ioannis Skarmoutsos, Stefano Mossa and Jannis Samios

European Molecular Liquids Group-EMLG Conference entitled: "Recent Progresses on the Experimental & Theoretical-Computational Techniques for the Study of Liquids and Supercritical Fluids." in Crete-Greece (2016)

12) Short range order and rotational dynamics in plastic crystal phases of water.

Andres Henao, Ioannis Skarmoutsos, Luis Carlos Pardo and Elvira Guardia

European Molecular Liquids Group-EMLG Conference entitled: "Recent Progresses on the Experimental & Theoretical-Computational Techniques for the Study of Liquids and Supercritical Fluids." in Crete-Greece (2016)

11) Local Structural Inhomogeneities, Hydrogen Bonding and Local Orientational Structure in Supercritical Water: Insights from Molecular Simulations.

Ioannis Skarmoutsos, Elvira Guardia and Jannis Samios

European Molecular Liquids Group-EMLG Conference entitled: "Recent Progresses on the Experimental & Theoretical-Computational Techniques for the Study of Liquids and Supercritical Fluids." in Crete-Greece (2016)

10) Carbon-based nanoporous networks as media for the separation of CO₂/CH₄ mixtures: A molecular dynamics approach.

Ioannis Skarmoutsos, George Tamiolakis and George E. Froudakis

30th Panhellenic Conference on Solid-State Physics and Materials Science, Heraklion, Crete-Greece (2014)

9) Local Structural Inhomogeneities, Hydrogen Bonding and Tetrahedral Structure in Supercritical Water: Insights from Molecular Simulations.

International School of Physics "Enrico Fermi", Varenna, Italy (2013)

8) Carbon based nanoporous networks as media for the separation of CO₂/CH₄ mixtures: A molecular dynamics approach.

International School of Physics "Enrico Fermi", Varenna, Italy (2013)

7) Solvation properties of alkali and halide ions in water from Car-Parinello molecular dynamics simulations

Ausias-March Calvo, Marco Masia, Ioannis Skarmoutsos and Elvira Guardia

CPMD meeting 2011: *Extending the limits of Ab Initio Molecular Dynamics Simulations for Chemistry, Materials Science and Biophysics, Barcelona-Spain* (2011)

6) Investigation of the Local Density Inhomogeneities and Dynamics in Neat Supercritical Fluids using MD simulation techniques: Comparison between Hydrogen- and Non Hydrogen-Bonded fluids.

Ioannis Skarmoutsos Nikolaos Elpidoforou and Jannis Samios, Winter School: *Computational Nanoscience: Do It Yourself*, John von Neumann Institute for Computing (NIC) Julich-Germany (2006)

5) Local density inhomogeneities and dynamic properties in supercritical fluids. A MD simulation study of neat sc CO₂ and of the binary mixture CH₄-CO₂.

Ioannis Skarmoutsos and Jannis Samios, *CCP5 and Marie Curie Actions: Methods in Molecular Simulation Summer School*, Cardiff University- UK (2005)

4) Is the hydrogen-bonding network around the cis and trans N-Methylformamide (NMF) conformers responsible for the observation of different diffusion coefficients? A temperature dependent molecular dynamics approach.

Ioannis Skarmoutsos, Ricardo L. Mancera and Jannis Samios, European Molecular Liquids Group-EMLG Conference entitled: *"Complex Liquids: Fundamental Properties to Industrial Applications."* in Sheffield Hallam University- UK (2004)

3) Molecular Dynamics simulations of cis-trans N-Methylformamide liquid mixture. Structure, dynamics and hydrogen bonding analysis.

Ioannis Skarmoutsos and Jannis Samios, European Molecular Liquids Group-EMLG Conference entitled: *"Molecular Liquids. Routes from Local Order to Large-Scale Cooperativity."* in Castelvechchio Pascoli- Italy (2003)

2) Supercritical Methane. Investigation of the bulk thermodynamic, structural and dynamic properties.

Ioannis Skarmoutsos and Jannis Samios, European Molecular Liquids Group-EMLG Conference entitled: *"Molecular Liquids. Routes from Local Order to Large-Scale Cooperativity."* in Castelvechchio Pascoli- Italy (2003)

1) Molecular dynamics studies of cis/trans N-methylformamide (NMF) liquid mixture using a new optimized all atom force field.

Ioannis Skarmoutsos and Jannis Samios, NATO Advanced Study Institute (ASI): Novel Approaches to the Structure and Dynamics of Liquids: Experiments, Theories and Simulations. Rhodes- Greece (2002)