

Curriculum Vitae et Studiorum

Ciro Achille Guido - Ph.D.

INFORMAZIONI PERSONALI

COGNOME	GUIDO
NOME	CIRO ACHILLE
DATA DI NASCITA	26/06/1983
POSIZIONE ATTUALE	RICERCATORE TEMPO DET. LETT. B S.S.D.CHIM/02 - CHIMICA FISICA
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Responsabile dell'unità "LIME lab" di Fotofisica e Fotochimica Teorica per il Gruppo di Chimica Fisica NanoMAT, PI del progetto PRIN "EnvELOP" finanziato dal MUR.

- Titoli Accademici:

- 10/05/2019: **Abilitazione Scientifica Nazionale Italiana Professore di seconda fascia S. C. 03/A2**
- 20/09/2011: **Diploma di Perfezionamento (Ph.D.) in Chimica**, conseguito presso la **Scuola Normale Superiore** di Pisa, con voto **70/70 e lode**.
Titolo della tesi: "*TD-DFT and TD-DFT/PCM approaches to molecular electronic excited states in gas phase and in solution*"
Relatori: Prof. Jacopo Tomasi e Prof. Benedetta Mennucci.
- 11/10/2007: **Laurea Magistrale in Chimica – Curriculum Chimica-Fisica** conseguita presso l'Università di Pisa con voto **110/110 e Lode**.
Titolo della tesi: "*Metodi Quantistici Per Il Calcolo Dello Scattering Raman Risonante Di Molecole In Soluzione*".
Relatori: Prof. Jacopo Tomasi e Prof. Benedetta Mennucci.
- 21/10/2005: **Laurea triennale in Chimica** conseguita presso l'Università di Pisa con voto **110/110 e lode**.
Titolo della tesi: "*Studio termodinamico delle interazioni polimero-tensioattivo in acqua*". Relatore: Prof. Paolo Gianni.

- **Organizzazione, direzione e coordinamento di centri o gruppi di ricerca nazionali e internazionali o partecipazione agli stessi e altre attività di ricerca quali la direzione o la partecipazione a comitati editoriali di riviste**

- Direzione di progetti di ricerca ottenuti su finanziamento:

- **19/06/2023**: **PI progetto PRIN EnvELOP su finanziamento del Ministero dell'Università e Ricerca**: 216,736 €
- **27/10/2022**: **Vincitore del finanziamento F-CUR Università di Siena**: 35,000 € (offerta declinata per trasferimento RTD-B in altro ateneo)
- **11/07/2017 - 10/07/2018**: **Vincitore del finanziamento LUMOMAT:** "Fully Consistent Polarization Response, FCPol-Resp", 60.000 €, finanziato da Région des Pays de la Loire (France) e Comunità Europea. **P.I.:** Dr. Ciro A. Guido e Prof. D. Jacquemin. Université de Angers e Université de Nantes (Francia).
- **11/07/2016 - 10/07/2017**: **Vincitore del finanziamento LUMOMAT:** "The Fate of Excitation Energy in solution, EE-Fate", 60.000 €, finanziato da Région des Pays de la Loire (France) e Comunità Europea., **P.I.:** Dr. Ciro A. Guido e Prof. D. Jacquemin. Université de Angers e Université de Nantes (Francia).
- 25/08/2009 – 31/07/2010: **finanziamento Erasmus placement per la mobilità internazionale** degli studenti della **Scuola Normale Superiore di Pisa**. Studente Ph.D. in scambio presso **École Normale Supérieure de Paris** e ricerca presso il gruppo MSC (Modélisation de Systèmes Complexes) del Prof. Carlo Adamo presso Ecole Nationale Supérieure de Chimie de Paris-Chimie Paristech (Parigi, Francia)

- Partecipazione a progetti di ricerca ottenuti su finanziamento:

- **01/12/2021 - 27/10/2022.** Programma “Dipartimento d’eccellenza 2018-2022” progetto “Sviluppo, valutazione ed applicazione di software per lo studio della spettroscopia e fotoreattività di cromofori incorporati in cavità proteiche”. **Ricercatore Tempo Determinato lett. A.**
- **01/10/2021 - 30/11/2021.** Progetto “**ProID**, Ultrafast Raman Technologies for Protein Identification and Sequencing” H2020-EU.1.2.1 ID:964363, € 2988513,75. Coord. IIT-Genova. Partecipante. Università di Padova (€ 250125). Referente: Stefano Corni. Ricercatore: Ciro A. Guido
- **01/10/2020 - 30/09/2021.** Progetto “**TAME-Plasmons**, a Theoretical chemistry Approach to tiME-resolved molecular Plasmonics”, ERC-CoG-2015 - **ERC Consolidator Grant project: GA 681285 2016-2021**, 1,432,890 €. P.I. Prof. S. Corni (Università di Padova, Italia). Researcher: Ciro A. Guido
- **01/10/2018 - 30/09/2020.** Progetto “Simulation of energy and electron transfer for molecules close to plasmonic nanostructures, **PlasmoChem**” MIUR, “**progetti FARE**” grant number **R164LZWZ4A, 2016-2020** 152,313 € P.I. Prof. S. Corni (Università di Padova, Italia). Researcher: Ciro A. Guido.
- **01/07/2014 - 30/06/2016.** Progetto “**EnLight**, The interplay between quantum coherence and environment in the photosynthetic electronic energy transfer and light-harvesting: a quantum chemical picture”, **ERC Starting Grant project: 277755 2011-2016**, 1,300,000 €. PI: Prof. B Mennucci (Università di Pisa, Italia). Researcher: Ciro A. Guido.
- **01/09/2012 – 19/06/2014.** Progetto “**Dinf DFT**, Développement et implementation de Nouvelles fonctionnelles d’échange et corrélation en théorie de la fonctionnelle de la densité” finanziato dalla **Agenzia Nazionale della Ricerca Francese, ANR** project. ANR 2010 BLANC n. 0425, 360,000 €. P.I. Prof. Pietro Cortona (École Centrale Paris) e Prof. Carlo Adamo (Chimie-Paristech). Researcher: Ciro A. Guido.
- **01/05/2011 – 31/07/2012.** Progetto “**Advanced atomistic simulations for carbon capture and sequestration**”, grant number 200021-132081 from **Swiss National Science Foundation**. 2010-2012. amount: 294,510 CHF. P.I. Prof. Wanda Andreoni (EPFL). Researcher: Ciro A. Guido.

- Attività di formazione o ricerca presso qualificati istituti italiani:

- 28/10/2022: **Ricercatore Tempo Determinato lett. B Chim/02** Università del Piemonte Orientale “A. Avogadro”.
- 01/12/2021 – 27/10/2022: **Ricercatore Tempo Determinato lett. A Chim/06** (Dipartimento d’Eccellenza 2018-2022). Università di Siena.
- 01/10/2021 – 30/11/2021: **assegnista di ricerca presso il gruppo del Prof. S. Corni** (H2020 project ProID). Università di Padova.
- 01/10/2020 – 30/09/2021: **assegnista di ricerca presso il gruppo del Prof. S. Corni** (ERC project TAME-Plasmon). Università di Padova.
- 01/10/2018 – 30/09/2020: **assegnista di ricerca presso il gruppo del Prof. S. Corni** (MIUR FARE project Plasmochem). Università di Padova.
- 01/07/2014 – 30/06/2016: **assegnista di ricerca presso il gruppo della Prof. B. Mennucci (ERC EnLight)**. Dipartimento di Chimica e Chimica Industriale, Università di Pisa, Italia.
- 01/01/2011 – 30/04/2011: **Borsa di studio e ricerca presso il gruppo della Prof. Benedetta Mennucci**. Dipartimento di Chimica e Chimica Industriale – Università di Pisa.
- 01/01/2008 – 31/12/2010: **Perfezionando della Scuola Normale Superiore di Pisa e membro del gruppo PCM dei Professori Jacopo Tomasi e Benedetta Mennucci**. Scuola Normale Superiore di Pisa – Dip. Chimica e Chimica Industriale. Università di Pisa.

- Attività di formazione o ricerca presso qualificati istituti stranieri:

- 11/07/2017 – 10/07/2018: **ricercatore a contratto presso dipartimento CEISAM Université de Nantes** (CO-PI progetto LUMOMAT FCPol-Resp).

- 11/07/2016 – 10/07/2017: ricercatore a contratto presso dipartimento CEISAM Université de Nantes (CO-PI progetto LUMOMAT EE-Fate).
- 01/09/2012 – 19/06/2014: Ricercatore CNRS presso il gruppo del Prof. Pietro Cortona. École Centrale Paris.
- 01/05/2011 – 31/07/2012: Ricercatore Postdoc presso il gruppo della Prof. Wanda Andreoni. École Polytechnique Fédérale de Lausanne (EPFL) – Centre Européen de Calcul Atomique et Moléculaire.
- 25/08/2009 – 31/07/2010: Studente Ph.D. in scambio presso il gruppo MSC (Modélisation de Systèmes Complexes) del Prof. Carlo Adamo e École Normale Supérieure de Paris. **Finanziamento Erasmus placement per la mobilità internazionale** degli studenti della Scuola Normale Superiore di Pisa. École Nationale Supérieure de Chimie de Paris-Chimie Paristech (Parigi, Francia)

- Organizzazione di congressi e attività per la divulgazione scientifica (terza missione)

- 16/06/2016: **Workshop internazionale “EnLight Workshop 2016”**, Università di Pisa, Comitato organizzatore: Benedetta Mennucci, Ciro A. Guido, Stefano Caprasecca, Sandro Jurinovich.
- 27/09/2016: **“Bright, la Notte dei Ricercatori in Toscana”**, coordinatore per lo stand del gruppo Molecolab della Prof. B. Mennucci.

- Partecipazione a comitati editoriali di riviste:

- 07/10/2022 – in corso: **Review Editor** della rivista open access “Frontiers in Chemistry”. Editore Frontiers
- 01/03/2021 – in corso: **Topic Editor** della rivista open access “Compounds” (ISSN 2673-6918). Editore MDPI https://www.mdpi.com/journal/compounds/topic_editors

- Attività di referee per riviste internazionali (peer-reviewed):

- The Journal of Physical Chemistry Letters
- The Journal of Chemical Physics
- Journal of Chemical Theory and Computation
- The Journal of Physical Chemistry A,B,C
- Journal of Computational Chemistry
- Physical Chemistry Chemical Physics
- Chemical Physics Letters
- Journal of Molecular Structure
- Theoretical Chemistry Accounts
- New Journal of Chemistry
- Scientific Reports
- RSC Advances
- Chirality

- Didattica, didattica integrativa e servizio agli studenti

- Didattica: corsi di laurea e laurea specialistica in Chimica:

- A.A. 2023/2024: Titolare del Corso di Chimica Teorica e Computazionale, Corso di Laurea Specialistica in Scienze Chimiche, Università del Piemonte Orientale. 6 CFU.
- A.A. 2022/2023: Titolare del Corso di Modellistica Chimica, Corso di Laurea Specialistica in Scienze Chimiche, Università del Piemonte Orientale. 6 CFU.
- A.A. 2022/2023: Titolare del Corso di Spettroscopie Ottiche, Corso di Laurea Specialistica in Scienze Chimiche, Università del Piemonte Orientale. 6 CFU.
- A.A. 2021/2022: dal 02/03/2002 al 27/04/2022: **Docente** del corso di Laboratorio di Chimica Organica mod. B. **48 ore**. Corso di Laurea in Chimica. Università di Siena. <https://www.unisi.it/ugov/person/207774>
- A.A. 2015/2016: Ven 13/05/2016 14:30-16:30, **2 ore** di seminario didattico: “Eccitazioni elettroniche e risposta del solvente” all’interno del corso “Modellistica Molecolare di Sistemi Complessi”, Università di Pisa (corso della Prof. Benedetta Mennucci).
- A.A. 2015/2016: dal 12/04/2016 al 21/04/2016 e dal 10/05/2016 al 12/05/2016. Esercitazioni di Laboratorio del corso di Chimica Fisica II, **Università di Pisa** (Prof. B. Mennucci): **36 ore**, (3 volte a settimana, 3 settimane, 4h per ogni lezione).

- A.A. 2015/2016: dal 3/12/2014 al 10/12/2014. Esercitazioni di Laboratorio del corso di Modellistica Molecolare Di Biomolecole, **Università di Pisa** (Prof. B. Mennucci): **12 ore** di laboratorio, (3 lezioni, 4h per ogni lezione).
- A.A. 2014/2015: dal 18/03/2015 al 03/04/2015. Esercitazioni di Laboratorio del corso di Chimica Fisica II, **Università di Pisa** (Prof. B. Mennucci): **36 ore** di laboratorio, (3 volte a settimana, 3 settimane, 4h di lezione).
- A.A. 2014/2015: dal 3/12/2014 al 10/12/2014. Esercitazioni di Laboratorio del corso di Modellistica Molecolare Di Biomolecole, **Università di Pisa** (Prof. B. Mennucci): **12 ore** di laboratorio, (3 lezioni, 4h per ogni lezione).
- A.A. 2010/2011: dal 10/01/2011 al 28/03/2011. Esercitatore e supervisore corso di Principi di Chimica Quantistica, **Scuola Normale Superiore di Pisa** (in collaborazione con il Prof. Ivo Cacelli). **24 ore** di lezione (2h a settimana, 12 settimane).
- A.A. 2009/2010: dal 4/01/2010 al 24/05/2010. Esercitatore e supervisore corso di Principi di Chimica Quantistica, **Scuola Normale Superiore di Pisa** (in collaborazione con il Prof. R. Moccia). **24 ore** di lezione (2h a settimana, 12 settimane).

- Attività di relatore di tesi, supervisione e mentoring:

- 2023: Membro del Consiglio dei docenti del Corso di Dottorato in Chemistry and Biology, Università del Piemonte Orientale.
- 2020/2021: **Relatore di Tesi dello studente** Jonas Nijssen, tesi di laurea triennale, B.Sc., KU Leuven University, Belgium. Titolo: "Optical properties of two amino benzothiadiazoles in solution: a computational study"
- 2018/2019: *Assistenza alla* supervisione della studentessa Giulia Dall'Osto, tesi di laurea, Università di Padova. Relatore Prof. S. Corni.
- 2017/2018: *Assistenza alla* supervisione della studentessa Ph.D. P. Verità, Università di Nantes (Francia). Relatore Prof. D. Jacquemin.
- 2015/2016: *Assistenza alla* supervisione della tesi di laurea dello studente Andrea Pasti, Università di Pisa. Relatrice Prof. B. Mennucci e Dr. Tarita Biver.

- Conseguimento di premi e riconoscimenti nazionali e internazionali per attività di ricerca.

- Fellowship:

- **30/11/2021. Maria Zambrano Fellowship from Spanish government and EU (NextGenerationEU).** MARIA ZAMBRANO GRANTS FOR THE ATTRACTION OF INTERNATIONAL TALENT: **1 classificato (score 100/100)** nel settore Health Sciences per la call Maria Zambrano bandita dal Ministerio de Universidades Spagnolo e l'Università di Barcelona. Declined (<https://seu.ub.edu/documentPublic/download/274629>).
- **07/09/2018. Marie Skłodowska-Curie Actions-COFUND Fellowship "Beatriu de Pinos".** 92000 euro di budget. Vincitore fellowship per "outstanding postdoctoral researchers" e host organization in Catalonia (<https://euraxess.ec.europa.eu/worldwide/brazil/postdoctoral-research-grants-catalonia-beatriu-de-pin%C3%B3s-programme-2019-msca>).
Titolo del Progetto: "Development of computational models for describing environment polarization and excited state dispersion interactions in photosynthetic light harvesting proteins".

- Abilitazioni Scientifiche:

- **15/03/2021. Abilitazione Scientifica Nazionale Sistema Universitario della Catalunya Professor Lector.** Qualifica numero 1TQLV664H. 2a Convocatoria Lector (2020).
- **10/05/2019. Abilitazione Scientifica Nazionale Italiana Professore di seconda fascia S. C. 03/A2** La commissione ha riconosciuto che il candidato presenta una maturità scientifica richiesta per le funzioni di professore di II fascia per il settore concorsuale 03/A2.
- **09/02/2018. Abilitazione Scientifica Nazionale Francese alla funzione di Maîtres de Conférences** (Insegnante-Ricercatore/Professore Associato). Qualifica numero: 14231260035D. Periodo 2018-2022. La commissione ha riconosciuto che il candidato presenta una maturità scientifica richiesta per le funzioni di Maîtres de Conférences per la classe 31 (chimica teorica, fisica ed analitica).
- **07/02/2014. Abilitazione Scientifica Nazionale Francese alla funzione di Maîtres de Conférences** (Insegnante-Ricercatore/Professore Associato). Qualifica numero: 14231260035P. Periodo 2014-2018.

La commissione ha riconosciuto che il candidato presenta una maturità scientifica richiesta per le funzioni di Maîtres de Conférences per la classe 31 (chimica teorica, fisica ed analitica).

- Relatore a congressi e convegni di interesse nazionale e internazionale:

1. **INVITED TALK.** "Dispersion Interactions At The Excited State: Influence On Light-Responsive Properties Of Biosystems" CT4OPTO, PSI-K and CECAM Workshop, organized by CNR-Nano, Modena, (Italy). 3-5/07/2023.
2. **INVITED TALK.** "Ab initio simulation of single amino acid SERS of proteins near plasmonic nanoparticles". ProID workshop. IIT-Genova (Italy). 27/01/2022
3. "Exploring the spatial features of electronic transitions in biomolecular systems by swift electrons", Psi-K workshop "Principles of light-induced charge transfer for optogenetics", CNR Nano, Modena (Italy). 14-16 June 2021.
4. "Non-equilibrium dynamical effects on excited states within Polarizable Continuum models", CECAM workshop "Nonequilibrium dynamical solvent effects on excited states: from spectroscopy to photoreactivity", CECAM, Lausanne (Switzerland). 14-17 June 2021.
5. **INVITED TALK.** "Simulating advanced excitation energy loss spectroscopies of molecular excitations", Workshop day: *Giornata della Chimica Teorica*, University of Padua, Padua (Italy). 25/06/2019
6. "Control of Coherences and Optical Responses of Pigment-Protein Complexes by Plasmonic Nanoantennae", *Plasmonica 2019*, International Workshop on Plasmonics, Naples (Italy), 19/06/2019 - 21/06/2019.
7. **INVITED TALK.** "Excited State Gradients for a State-Specific Continuum Solvation Approach: a Lagrangian TDDFT Formulation and Implementation of VEM". Workshop: *Journées Scientifiques SolvATE*. 16/05/2018 – 18/05/2018.
8. **INVITED TALK.** "Polarizable schemes for simulating excited state processes in multichromophoric systems: from continuum to discrete environment description" *French Network of Theoretical Chemistry, annual day workshop of West-France*. Nantes (France), 10/11/2017.
9. "Excited State Gradients for a State-Specific Continuum Solvation Approach: a Lagrangian TDDFT Formulation and Implementation of VEM" *Coding in Solvation* workshop (WATOC satellite meeting), Livorno, Italy, 22-25 August 2017.
10. "On the role of charge transfer states of LH2 system: a state-specific MMPol-TDDFT approach coupled to excitonic models." *DCTC 2015*, Italian Chemical Society, Rome, Italy, 14-16 December 2015.
11. "Charge-transfer like excitations in solution: a critical assessment of TDDFT/Continuum Models", **ACS Spring 2015** National Meeting, Denver, CO, 22-25 March 2015. Meeting Abstract 202, Volume 249.
12. **INVITED TALK.** "Charge-transfer like excitations in solution: a critical assessment of TDDFT/Continuum Models" *Solution for Solvation*, International Workshop to Celebrate Jacopo Tomasi, Scuola Normale Superiore-CNR-Unipi, 31 August – 1 September 2014.
13. "Orbital based tools for TDDFT computational spectroscopy" 9th Congress on Electronic Structure: Principles and Applications (ESPA 2014). Badajoz, 2 - 4 July 2014.
14. "Understanding molecular excitation character made simpler" TD-DFT Conference, Université de Nantes, 23-26 April 2013.
15. "The time dependent problems in solvent model: the PCM point of view" Ecole Nationale Supérieure de Chimie de Paris – Paris, 24 November 2009.
16. "A systematic study of TD-DFT performance in the determination of excited state geometries and properties" COTAW09, FUNDP - Namur (Belgium), 16-19 September 2009.

Attività di ricerca:

L'attività di ricerca è principalmente (ma non esclusivamente) focalizzata sullo sviluppo di modelli teorici e computazionali e relativa implementazione in codici di calcolo, per la **simulazione di proprietà ottiche ed elettroniche in sistemi molecolari e supra-molecolari in presenza di un ambiente esterno**. Tale ambiente può rappresentare un solvente isotropico, un ambiente anisotropo discreto quale una cavità proteica e/o la presenza di nanoparticelle metalliche soggette ad eccitazione plasmonica.

All'attività di sviluppo ed implementazione è al contempo accoppiata un'attività di tipo applicativo, tramite collaborazioni con gruppi sperimentali, quali quello dei Professori Pescitelli e Di Bari (Università di Pisa) o del Dr. Vincenzo Grillo dell'Istituto CNR-Nano (Modena).

Una concisa rappresentazione dei diversi interessi e risultati conseguiti nell'ambito della chimica teorica e computazionale è riportata di seguito (in inglese).

- Principali risultati conseguiti:

- Starting from the 1 December 2021, I spent less than 1 year as Assistant Professor at Siena University, performing research in the field of computational Photochemistry and Photobiology, collaborating with Prof. Massimo Olivucci, developing new theoretical/computational methodologies to describe the **polarization and dispersion** environment effects on the photochemistry of Rhodopsins, in a multireference (CASSCF/XMS-CASPT2) QM/MM protocol and interfaced with the automated Rhodopsin Model (ARM) recently developed in the lab.
- My work at University of Padova have been mainly focused on the reformulation of the time-dependent problem of the QM description of molecular systems in the presence of a polarizable external environment and/or metallic nanoparticles in terms of **open quantum systems** theory. By using a non-Markovian stochastic Schrödinger equation (SSE) formulation, a comprehensive picture of the electronic energies and the coupling between solute and solvent electronic dynamics is provided. This is indeed a new formulation of the coupling between QM methods and continuum solvation models which will allow **to increase the accuracy and achieve a complete description of the different timescales in ultrafast spectroscopies, for instance in molecular experiments involving plasmonic effects of metal nanoparticles**.
Publications: *J. Chem. Phys.*, **152**, 174114 (2020); *J. Chem. Phys.*, **153**, 200901 (2020); *Phys. Chem. Chem. Phys.*, **22**, 16734 (2020). Funding: ERC TAME-Plasmon and MIUR FARE Plasmochem.
- In collaboration with the experimental group of Vincenzo Grillo at CNR-Nano Modena, we conceived a new kind of experiment that extends the technology of electron energy loss spectroscopy (EELS) to probe (supra-)molecular systems coupled to the analysis of the single components of orbital angular momentum (OAM) of the outgoing electrons: I theoretically demonstrated that this new spectroscopy provides information on **the multipolar components and even the chirality of molecular electronic transitions, superseding the usual optical spectroscopies for those cases that are problematic, such as dipole-forbidden transitions, at a very high spatial resolution**.
Publication: *J. Chem. Theory Comput.*, **17**, 2364 (2021). Funding: ERC TAME-Plasmon and MIUR FARE Plasmochem.
- As CO-PI of two LUMOMAT projects in Nantes, I developed new QM methods to describe molecular electronic transition in solution that consider the correct solvent polarization response due to the electronic excited state of interest. In this framework, I developed and implemented a **Lagrangian formulation of the Vertical Excitation Method** (introduced during my Ph. D in collaboration with the group of Prof. Thrular at Minnesota University, *Chem. Sci.* **2**, 2143, 2011, **160 citations**) and its **analytical gradients**, opening the way to simulate with high accuracy emission spectra, quantum yields or dipole moments. On the same foot, in collaboration with X. Blase and I Duchemin of CNRS, we introduced, for the first time, the extension of **continuum models to the many body perturbation theory within the GW-BSE framework (the GW-BSE/PCM approach)**.
Publications: **101 total citations**. *Chem. Sci.* **9**, 4430 (2018); *Int. J. Quantum Chem.* **119**, e25711 (2019); *J. Chem. Theory Comput.*, **14**, 1544 (2018); *Phys. Chem. Chem. Phys.*, **21**, 2307 (2019); *J. Chem Phys.* **146**, 204106 (2017). Funding: LUMOMAT projects FCPol-Resp ed EE-Fate.
- During my postdoc for the ERC Project "EnLight" in the group of Prof. Mennucci, we developed a **QM/MM excitonic model to describe energy transfer processes in pigment-protein systems**, such as Light Harvesting complexes in plants and bacteria. The developed model was also extended to the calculation of different spectroscopies of multichromophoric systems, such as the **Excitonic Circular Dichroism (ECD) spectra in a gauge invariant formulation** and implemented in a Python software (EXAT) which allows the calculation of excitonic properties starting from results obtained through the Gaussian software (G09 and G16). These **strategies have been also coupled to a plasmonic description of metal nanorods, deeply analyzing the different radiative and non-radiative channels and the possibility of tuning the fluorescence by metal nanoparticles**.

Publications: **309 total citations**. *J. Phys. Chem. Lett.* **9**, 6892 (2018); *J. Comput. Chem.* **39**, 279 (2017); *J. Phys. Chem. Lett.*, **7**, 2189 (2016). *Chem. Comm.* **51**, 10498 (2015); *JCTC*, **11**, 5782 (2015); *JCTC*, **9**, 2209 (2013). Funding: ERC project EnLight

- The quantum mechanical description of systems based on DFT was the subject of my work at École Centrale Paris, working with Prof. Pietro Cortona and Prof. Carlo Adamo. **We proposed a new functional, PBE0-1/3, and introduced two new metrics for a quantitative description of molecular excited states.** The new hybrid functional generally improves performances in the cases of barrier heights and of CT excitations and different studies from the groups of Brédas and Galli showed the very good performances in the description of the electronic structure of organic charge-transfer molecular crystals. PBE0-1/3 is now implemented in many quantum chemistry and solid physics codes. The two new metrics have been used to evaluate the CT character of organic chromophores, to rationalize the singlet-triplet gap in organic light-emitting diode (OLED) materials, and a relationship with the nonlinear optical properties of push-pull systems was also provided [*JCTC*, **11**, 4182 (2015)]. Their definition has been also extended to tight binding TD-DFT, and implemented in an open-source code, *Multiwfn*. **On this subject we already received more than 250 citations.** Publications: **322 total citations**. *J. Chem. Phys.* **140**, 104101 (2014); *J. Chem. Theory Comput.* **9**, 3118 (2013); *J. Chem. Phys.* **138**, 021104 (2013). Funding: ANR project "Dinf-DFT"
- I performed ab-initio Car-Parrinello molecular dynamics simulations to study the fundamental reactions accompanying the capture of carbon dioxide in amine solutions at fully atomistic level, during my first postdoctoral experience at the Centre Européen de Calcul Atomique et Moléculaire (CECAM) in Lausanne, working with Prof. Andreoni. I investigated the role of the zwitterionic carbamate, a species long proposed as intermediate in the formation of a stable carbamate in a dilute aqueous solution. **We have shown that CO₂ release and deprotonation are competitive routes for its dissociation and that water molecules play a crucial role in both pathways resulting in large entropic effects.** Publication: **40 citations** *J. Chem. Theory Comput.* **9**, 28 (2013). Funding: Swiss National Science Foundation project n. 132081.
- Other research interests, during my PhD at **Scuola Normale Superiore di Pisa**, include the benchmarking of TDDFT performances (*JPCA*, **114**, 13402, 2010; *PCCP* **12**, 8016, 2010, **166 citations**) in collaboration with Prof. Carlo Adamo at Chimie-Paristech, and the implementation of a fully automated algorithm for the **anharmonic second order perturbative correction to the infrared absorption intensities** in the Gaussian suite of programs (*CPL* **496**, 157, 2010, **103 citations**), in collaboration with Prof. V. Barone, past director of Scuola Normale.

- Pubblicazioni più rappresentative su nuovi sviluppi teorici e computazionali.

- "Exploring the spatial features of electronic transitions in molecular and biomolecular systems by swift electrons"
C. A. Guido*, E. Rotunno, M. Zanfognini, S. Corni and V. Grillo, *J. Chem. Theory Comput.*, **17**, 2364 (2021). *Corresponding Author.

In this contribution a conceived new kind of experiment is proposed, that extends the technology of electron energy loss spectroscopy (EELS) to probe (supra-)molecular systems. Indeed, understanding the electronic structure of matter is a formidable task that largely made use of optical spectroscopies and their corresponding selection rules, but not all the electronic transitions can be probed: for instance, a long debate in the literature is still ongoing on the possible role of charge transfer (CT) states in photosynthetic mechanisms: being dark, it can only be indirectly probed. On the other hand, electron-beam spectroscopies are now emerging as probing techniques to study optical excitations with combined space, energy, and time resolution.

Performed in a scanning transmission electron microscope, EELS is based on inelastic scattering of fast electrons in a thin specimen and, very recently, new electron optics configuration has been introduced, opening the way to the analysis of the single components of orbital angular momentum (OAM) of the outgoing electrons. **Physical insight into the proposed experiment is provided by means of a rigorous model to obtain the transition rate and the selection rule.** Numerical simulations of DNA G-quadruplexes and other biomolecular systems, based on time dependent density functional theory calculations, point out that **the conceived new technique can probe the multipolar components and even the chirality of molecular transitions, superseding the usual optical spectroscopies for those cases that are problematic, such as dipole-forbidden transitions, at a very high spatial resolution.**

The work has been selected for ACSLiveSlides.

- "An Open Quantum System Theory for Polarizable Continuum Models "
C. A. Guido, M. Rosa, R. Cammi and S. Corni, *J. Chem. Phys.* **152**, 174114 (2020).

In this work the time dependent (TD) problem of a solute described by Quantum Chemistry within an environment represented as a polarizable continuum model (PCM) has been reconsidered in terms of the open quantum systems (OQS) theory.

In particular, by using a non-Markovian stochastic Schrödinger equation (SSE) formulation, a comprehensive picture of the electronic energies and the coupling between solute and solvent electronic dynamics is provided: **it proves to be a unifying theoretical framework able to describe the delayed solvent response due to the solute charge density reorganization evolving in a time regime, naturally including polarization and dispersion interactions** (due to the solute fluctuations and solvent polarization response and vice versa). The **OQS-PCM accounts, by construction, for the proper coupled time scales of solute and solvent electronic dynamics and covers all the possible relative time scales of electronic solute and solvent responses**, from the limit where the solute electronic dynamics is much faster than the solvent one to the opposite limit where the solvent electronic response is considered faster than any electronic dynamics taking place in the solute.

- "Excited State Gradients for State-Specific Continuum Solvation Models: the Vertical Excitation Model within a Lagrangian TDDFT

formulation”

C. A. Guido*, G. Scalmani, B. Mennucci and D. Jacquemin, *J. Chem Phys.* 146, 204106 (2017). *Corresponding Author.

This study presents the analytical expression of the gradients of the Vertical Excitation Model approach, for which I used a Lagrangian formulation in the TD-DFT framework. The method here presented opens the way to calculate excited state (ES) properties and optimize ES molecular structures using a state-specific (SS) solvation model that represents the correct polarization response scheme. The SS corrections become of fundamental importance for electronic transitions involving a very large electron density rearrangement. **This is the first work on ES geometry optimization by SS TDDFT analytical gradients in literature.** The method here presented can be extended to other types of polarizable response of the environment, therefore also to discrete models, such as polarizable molecular mechanics (MM) in QM/MM methods.

• “The Bethe-Salpeter formalism with polarizable continuum embedding: combining state-specific and linear-response features”
I. Duchemin, **C. A. Guido**, D. Jacquemin and X. Blase, *Chem. Sci.* **9**, 4430 (2018).

This work introduces, for **the first time, the combination of the GW and Bethe–Salpeter equation (BSE) formalism with the PCM approach.** In particular, I contribute to this work by demonstrating that **GW/BSE-PCM straightforwardly includes, on the same footing, both linear-response (LR) and state-specific (SS) contributions to the solvatochromic shifts.**

This result was established on analytic grounds as well as by comparing the numerical results obtained for several molecules or complexes using BSE + PCM or TD-DFT + PCM approaches, the latter being performed with both the LR and corrected-LR formalisms. It is important to remark that, later on, by using the framework of the open quantum system theory, I pointed out that GW-BSE/PCM can include also regimes where the solvent response is much faster than the solute electrons (see JCP, 152, 174114, 2020).

• “On the metric of charge transfer molecular excitations: a simple chemical descriptor”
C. A. Guido*, P. Cortona, B. Mennucci and C. Adamo, *J. Chem. Theory Comput.* **9**, 3118 (2013). *Corresponding Author.

In this contribution, a new index is defined with the aim of further exploring the metric of excited electronic states in the framework of the time-dependent density functional theory. This descriptor, called Δr , is based on the charge centroids of the orbitals involved in the excitations and can be interpreted in terms of the hole–electron distance. The tests carried out on a set of molecules characterized by a significant number of charge-transfer excitations well illustrate its ability in discriminating between short ($\Delta r \leq 1.5$ Å) and long-range ($\Delta r \geq 2.0$ Å) excitations. **Based on the well-known pitfalls of TD-DFT, its values can be then associated to the functional performances in reproducing different type of transitions and allow for the definition of a “trust radius” for GGA and hybrid functionals.** The study of some well-known difficult cases for other metric descriptors gives further evidence of **the high discrimination power of the proposed index, that received a very large number of citations (more than 200)** and have been implemented also in different software (G16 development versions, Multiwfn and NTOBuilder just cite few).

- **Competenze tecniche:**

- **Competenze Informatiche:**

- Sistemi Operativi: GNU/Linux e UNIX per workstations, Apple IOS and MS Windows.
- Linguaggi di Programmazione: FORTRAN 77 and 95, Python, OpenMP, MPI, awk and bash scripting.
- Software per quantum chemistry and modeling: GAUSSIAN 16, GAMESS, Molcas, OpenMolcas, GaussView, CPMD, AMBER, VMD. **Sviluppo ed implementazione di subroutine per GAUSSIAN G09 e G16.**
- Altri software: Matlab, Grace, Gnuplot, Origin, MS Office.

- **Software e moduli computazionali sviluppati:**

- **OAM-EELS** (C. A. Guido, Università di Padova, 2020).

Programma Fortran per il calcolo di probabilità di perdita di energia elettronica risolte nel momento angolare per interazione molecola – elettroni veloci.

- **Gaussian 09, development version I09:** implementazione del Vertical Excitation Method (VEM) e relativi gradienti analitici. Collaborazione con Giovanni Scalmani, Gaussian Inc.

- **Gaussian 16:** implementazione del Vertical Excitation Method (VEM) e relativi gradienti analitici. Collaborazione con Giovanni Scalmani, Gaussian Inc.; Implementazione delle metriche di caratterizzazione degli stati eccitati

- **EXAT** (S. Jurinovich, L. Cupellini, C. A. Guido e B. Mennucci, Università di Pisa, 2014).

Python software per il calcolo di proprietà eccitoniche. <https://zenodo.org/record/4282773>

- **NTOBuilder** (L. Cupellini, S. Caprasecca e C. A. Guido, Università di Pisa, 2018).

<https://zenodo.org/record/3948078>

Python software per la caratterizzazione di eccitazioni elettroniche ottenute con approcci TDDFT e Coupled Cluster.

- Lingue:

- Italiano: madrelingua.
- Inglese: scritto e parlato. Livello: Ottimo
- Francese: scritto e parlato. Livello: Ottimo
- Spagnolo: scritto e parlato. Livello: Base.

- Altre Informazioni Accademiche:

- Insegnamenti di Perfezionamento presso la Scuola Normale Superiore di Pisa:

- “*Chimica Computazionale*”, tenuto dal Prof. Michele Parrinello. **Voto 30/30**. Scuola Normale Superiore di Pisa.
- “*Chimica Supramolecolare e Magnetochimica*” tenuto dal Prof. Dante Gatteschi. **Voto 28/30**. Scuola Normale Superiore di Pisa.
- “*Metodi avanzati in Chimica Teorica*”, tenuto dal Prof. Renato Colle. **Voto 29/30**. Scuola Normale Superiore di Pisa.
- “*Molecular Modeling*”, tenuto dal Prof. Vincenzo Barone. Scuola Normale Superiore di Pisa.

- Collaborazioni (nazionali ed internazionali):

- Stefano Corni (Padova University, Italy)
- Benedetta Mennucci (Pisa University, Italy)
- Carles Curutchet (Barcelona University, Spain)
- Gennaro Pescitelli (Pisa University, Italy)
- Lorenzo di Bari (Pisa University, Italy)
- Denis Jacquemin (Nantes University, France)
- Carlo Adamo (Chimie Paristech, France)
- Laura Zanetti-Polzi (CNR-Nano, Italy)
- Giovanni Scalmani (Gaussian Inc., U.S.A.)
- Pietro Cortona (École Centrale Paris, France)
- Xavier Blase (CNRS Grenoble, France)
- Jacob Kongsted (Copenhagen University, Denmark)
- Vincenzo Grillo (CNR-Nano, Modena, Italy)
- Enzo Rotunno (CNR-Nano, Modena, Italy)
- Francesco De Angelis (IIT, Genova, Italy)
- Nicolò Maccaferri (Umeå University, Umeå, Sverige)

LISTA DELLE PUBBLICAZIONI

35 publications: 34 journal articles, 1 book chapter

Citation Metrics

Web of Science (WoS): **Ciro A. Guido** - D-3533-2012

WoS ResearcherID: AAG-4460-2020

Scopus author ID: 15072889800

ORCID number: 0000-0003-1924-2862

h-index: **23** (fonte: Scopus e WebOfScience)

Times cited: **1972** from 1604 articles.

Corresponding author (*).

• **Journal articles:**

1. "CPL calculations of [7]helicenes with alleged exceptional emission dissymmetry values"
C. A. Guido*, F. Zinna and G. Pescitelli *J. Mater. Chem. C*, DOI: 10.1039/D3TC01532G (2023). IF: 8.067 Q1 Chemistry and Material Sciences.
2. "A Simple Protocol for Capturing both Linear-Response and State-Specific Effects in Excited-State Calculations with Continuum Solvation Models"
C. A. Guido*, A. Chrayteh, G. Scalmani, B. Mennucci and D. Jacquemin *J. Chem. Theory Comput*, **17**, 5155 (2021).
IF: 6.006 Q1 Physical and Theoretical Chemistry.
3. "Exploring the spatial features of electronic transitions in molecular and biomolecular systems by swift electrons"
C. A. Guido* E. Rotunno, M. Zanfognini, S. Corni and V. Grillo, *J. Chem. Theory Comput*, **17**, 2364 (2021).
IF: 6.006. Q1 Physical and Theoretical Chemistry. **Selected for ACSLiveSlides**
4. "Hybrid theoretical models for molecular nanoplasmonics".
E. Coccia, J. Fregoni, **C. A. Guido**, M. Marsili, S. Pipolo and S. Corni *J. Chem. Phys.* **153**, 200901 (2020)
IF: 3.488. Q1 Physical and Theoretical Chemistry
5. "Investigating ultrafast two-pulse experiments on single DNQDI fluorophores: a stochastic quantum approach".
G. Dall'Osto, E. Coccia, **C. A. Guido** and S. Corni, *Phys. Chem. Chem. Phys.* **22**, 16734 (2020).
IF: 3.676. Q1 Physical and Theoretical Chemistry.
6. "An Open Quantum System Theory for Polarizable Continuum Models"
C.A. Guido, M. Rosa, R. Cammi and S. Corni, *J. Chem. Phys.* **152**, 174114 (2020)
IF: 3.488. Q1 Physical and Theoretical Chemistry.
7. "On the description of the environment polarization response to electronic transitions" **Invited Perspective**.
C. A. Guido* and S. Caprasecca, *Int. J. Quantum Chem.* **119**, e25711 (2019).
IF: 2.444. Q1 Physical and Theoretical Chemistry.
8. "First-principles investigation of the double ESIPT process in a thiophene-based dye"
P. V erit e, **C. A. Guido** and D. Jacquemin, *Phys. Chem. Chem. Phys.*, **21**, 2307 (2019).
IF: 3.676. Q1 Physical and Theoretical Chemistry
9. "Coupling to Charge Transfer States Is the Key to Modulate the Optical Bands for Efficient Light-Harvesting in Purple Bacteria"
L. Cupellini, S. Caprasecca, **C. A. Guido**, F. M uh, T. Renger and B. Mennucci. *J. Phys. Chem. Lett.* **9**, 6892 (2018). IF: 6.475. : Q1 Material Science (miscellaneous)
10. "Excited State Dipole Moments in Solution: Comparison Between State-Specific and Linear-Response TD-DFT Values"
C. A. Guido*, B. Mennucci, G. Scalmani and D. Jacquemin, *J. Chem. Theory Comput.*, **14**, 1544 (2018).
IF: 6.006. Q1 Physical and Theoretical Chemistry

11. "Density-dependent formulation of dispersion interactions in hybrid QM/MM models"
C. Curutchet, L. Cupellini, J. Kongsted, S. Corni, L. Frediani, A. Hykkerud Steindal, **C. A. Guido**, G. Scalmani, B. Mennucci, *J. Chem. Theory Comput.*, **14**, 1671 (2018).
[IF: 6.006](#). Q1 Physical and Theoretical Chemistry
12. "The Bethe-Salpeter formalism with polarizable continuum embedding: combining state-specific and linear-response features"
I. Duchemin, **C. A. Guido**, D. Jacquemin and X. Blase, *Chem. Sci.* **9**, 4430 (2018).
[IF: 9.825](#). Q1 Chemistry.
13. "EXAT: EXcitonic Analysis Tool"
S. Jurinovich, L. Cupellini, **C. A. Guido** and B. Mennucci, *J. Comput. Chem.* **39**, 279 (2017).
[IF: 3.376](#). Q1 Physical and Theoretical Chemistry
14. "Metrics for Molecular Electronic Excitations: A Comparison between Orbital- and Density-Based Descriptors"
M. Savarese, **C. A. Guido**, E. Brémond, I. Ciofini and C. Adamo, *J. Phys. Chem A* **121**, 7543 (2017).
[IF: 2.781](#). Q1 Physical and Theoretical Chemistry
15. "Excited State Gradients for State-Specific Continuum Solvation Models: the Vertical Excitation Model within a Lagrangian TDDFT formulation"
C. A. Guido*, G. Scalmani, B. Mennucci and D. Jacquemin, *J. Chem Phys.* **146**, 204106 (2017).
[IF: 3.488](#). Q1 Physical and Theoretical Chemistry
16. "Control of Coherences and Optical Responses of Pigment-Protein Complexes by Plasmonic Nanoantennae."
S. Caprasecca, **C. A. Guido**, and B. Mennucci. *J. Phys. Chem. Lett.*, **7**, 2189 (2016).
[IF: 6.475](#). Q1 Material Science (miscellaneous)
17. "Circularly Polarized Luminescence from Axially Chiral BODIPY DYEmers: an Experimental and Computational Study" Selected as cover article.
F. Zinna, T. Bruhn, **C. A. Guido**, J. Ahrens, M. Bröring, L. Di Bari, and G. Pescitelli. *Chem. Eur. J.* **22**, 16089 (2016).
[IF: 5.236](#). Q1 Chemistry (miscellaneous)
18. "An Ab Initio Description of the Excitonic Properties of LH2 and Their Temperature Dependence"
L. Cupellini, S. Jurinovich, M. Campetella, S. Caprasecca, **C. A. Guido**, S. M. Kelly, A. T. Gardinier, R. Cogdel, and B. Mennucci. *J. Phys. Chem. B*, **120**, 11348 (2016).
[IF: 2.991](#). Q1 Physical and Theoretical Chemistry
19. "Electronic Excitations in Solution: The Interplay between State Specific Approaches and a TD-DFT Description"
C. A. Guido*, D. Jacquemin, C. Adamo and B. Mennucci. *J. Chem. Theory Comput.* **11**, 5782 (2015).
Top 20 monthly most read articles.
[IF: 6.006](#). Q1 Physical and Theoretical Chemistry
20. "Negative Solvatochromism of Push-pull Biphenyl Compounds: A Theoretical Study"
S. Meng, S. Caprasecca, **C. A. Guido**, S. Jurinovich, and B. Mennucci. *Theor. Chem. Acc.* **134**, 150 (2015).
[IF: 1.702](#). Q2 Physical and Theoretical Chemistry
21. "Circular Dichroism and TDDFT Investigation of Chiral Fluorinated ArylBenzyl Sulfoxides"
R. Berardozzi, **C. A. Guido**, M. A. M. Capozzi, C. Cardellicchio, L. Di Bari and G. Pescitelli. *Eur. J. Org. Chem.* **25/2015**, 5554 (2015).
[IF: 3.021](#). Q1 Organic Chemistry
22. "The role of magnetic-electric coupling in the exciton-coupled ECD spectra. The case of bis-phenanthrenes."
S. Jurinovich, **C. A. Guido**, T. Bruhn, G. Pescitelli and B. Mennucci, *Chem. Comm.* **51**, 10498 (2015).
[IF: 6.222](#). Q1 Chemistry (miscellaneous)
23. "Plasmon Enhanced Light-Harvesting: Multiscale Modeling of the FMO Protein Coupled With Gold Nanoparticles".
O. Andreussi, S. Caprasecca, L. Cupellini, I. Guarnetti Prandi, **C. A. Guido**, S. Jurinovich, L. Viani, and B. Mennucci, *J. Phys. Chem. A* **119**, 5197 (2015).
[IF: 2.781](#). Q1 Physical and Theoretical Chemistry
24. "Effective Electron Displacements: a tool for TD-DFT Computational Spectroscopy"

- C. A. Guido***, P. Cortona and C. Adamo, *J. Chem. Phys.* 140, 104101 (2014).
IF: 3.488. Q1 Physical and Theoretical Chemistry
25. "On the metric of charge transfer molecular excitations: a simple chemical descriptor"
C. A. Guido*, P. Cortona, B. Mennucci and C. Adamo, *J. Chem. Theory Comput.* 9, 3118 (2013).
Top 20 monthly most read articles
IF: 6.006. Q1 Physical and Theoretical Chemistry
26. "Benchmarking TD-DFT for excited state geometries of organic molecules in gas-phase and in solution"
C. A. Guido*, S. Knecht, J. Kongsted and B. Mennucci, *J. Chem. Theory Comput.* 9, 2209 (2013).
Top 20 monthly most read articles
IF: 6.006. Q1 Physical and Theoretical Chemistry
27. "One Third: a new recipe for the PBE0 paradigm"
C. A. Guido, E. Brémond, C. Adamo and P. Cortona *J. Chem. Phys.* 138, 021104 (2013).
Top 20 most read communications of 2013
IF: 3.488. Q1 Physical and Theoretical Chemistry.
28. "The Fate of a Zwitterion in Water from Ab-Initio Molecular Dynamics: Monoethanolamine (MEA)-CO₂"
C. A. Guido, F. Pietrucci, G. A. Gallet and W. Andreoni, *J. Chem. Theory Comput.* 9, 28 (2013).
Top 20 monthly most read articles
IF: 6.006. Q1 Physical and Theoretical Chemistry
29. "Practical computation of electronic excitation in solution: vertical excitation model"
A. V. Marenich, C. J. Cramer, D. G. Truhlar, **C. A. Guido**, B. Mennucci, G. Scalmani and M. J. Frisch, *Chemical Science*, 2, 2143 (2011).
IF: 9.825. Q1 Chemistry
30. "On the TD-DFT Accuracy in Determining Single and Double Bonds in Excited-State Structures of Organic Molecules"
C. A. Guido*, D. Jacquemin, C. Adamo and B. Mennucci, *J. Chem. Phys. A*, 114, 13402 (2010).
Top 20 monthly most read articles
IF: 2.781. Q1 Physical and Theoretical Chemistry
31. "A fully automated implementation of VPT2 Infrared intensities "
V. Barone, J. Bloino, **C. A. Guido** and F. Lipparini, *Chem. Phys. Lett.* 496, 157 (2010).
Top 20 monthly most read articles
IF: 2.328. Q2 Physical and Theoretical Chemistry
32. "Planar vs. twisted intramolecular charge transfer mechanism in Nile Red: new hints from theory"
C. A. Guido, B. Mennucci, D. Jacquemin and C. Adamo, *Phys. Chem. Chem. Phys.* 12, 8016 (2010).
IF: 3.676. Q1 Physical and Theoretical Chemistry
33. "Structures and properties of electronically excited chromophores in solution from the polarizable continuum model coupled to the Time-Dependent Density Functional Theory "
B. Mennucci, C. Cappelli, **C. A. Guido**, R. Cammi and J. Tomasi, *J. Phys. Chem. A* 113, 3009 (2009).
IF: 2.781. Q1 Physical and Theoretical Chemistry
34. "Calorimetric investigation of the aggregation of lithium perfluorooctanoate on poly(ethyleneglycol) oligomers in water "
P. Gianni, L. Bernazzani, **C. A. Guido** and V. Mollica, *THERMOCHIMICA ACTA*, 451, 73 (2006).
IF: 3.115. Q2 Physical and Theoretical Chemistry
- **Book chapters:**
1. "Computational studies of environmental effects and their interplay with experiment"
B. Mennucci, S. Caprasecca and **C. A. Guido**, *Advances in Physical Organic Chemistry*, Volume 50, 2016, Elsevier. DOI: 10.1016/bs.apoc.2016.04.001.

Comunicazioni tramite poster:

1. Ciro A. Guido, Enzo Rotunno, Matteo Zanfrognini, Vincenzo Grillo, Stefano Corni
"Simulating advanced excitation energy loss spectroscopies of molecular excitations" ISTCP-X, Tromso (Norway), 11-17 July 2019.
2. "Excited State Gradients for a State-Specific Continuum Solvation Approach: a Lagrangian TDDFT Formulation and Implementation of VEM"
WATOC 2017, Munich (Germany), 27 August - 1 September 2017.
3. Guido Ciro Achille, Jurinovich Sandro, Loco Daniele, Pescitelli Gennaro and Mennucci Benedetta
"A QM excitonic scheme for simulating electronic circular dichroism spectra of multichromophoric systems."
International Congress of Quantum Chemistry 2015, IAQMS, Beijing (China), 8-13 June 2015.
4. Guido Ciro Achille, Cortona Pietro, Mennucci Benedetta and Adamo Carlo
"On the metric of charge transfer molecular excitations: a simple chemical descriptor".
DFT 2013, Durham University (U.K.), 9th -13th September 2013.
5. Guido Ciro Achille, Brémond Eric, Adamo Carlo and Cortona Pietro
"One Third: a new recipe for the PBE0 paradigm".
GDR CoDFT 2013, Guidel Plage (France), 21th – 24th May 2013.
6. Guido Ciro Achille, Mennucci Benedetta, Jacquemin Denis and Adamo Carlo
"Planar versus twisted intramolecular charge transfer mechanism in Nile Red: new hints from theory".
Italian National Congress of Physical Chemistry, Società Chimica Italiana, Stresa (Italy), 20-24 September, 2010.
7. Guido Ciro Achille, Mennucci Benedetta, Jacquemin Denis and Adamo Carlo
"Planar versus twisted intramolecular charge transfer mechanism in Nile Red: new hints from theory".
IX Girona Seminar: Electron Density, Density Matrices, and Density Functional Theory, University of Girona, Girona (Spain), 5th – 8th July 2010
8. Guido Ciro Achille, Mennucci Benedetta, Jacquemin Denis and Adamo Carlo
"Planar versus twisted intramolecular charge transfer mechanism in Nile Red: new hints from theory".
Winter Modeling 2010, Scuola Normale Superiore di Pisa, Pisa (Italy), 26th February 2010
9. Guido Ciro Achille, Mennucci Benedetta and Adamo Carlo, "A systematic study of TD-DFT performance in the determination of excited state geometry and properties".
DFT09, Université Claude Bernard Lyon 1, Lyon (France) 31th August – 4th September 2009
10. Guido Ciro Achille, Mennucci Benedetta, Cappelli Chiara, Cammi Roberto, Tomasi Jacopo, "TDDFT-IEFPCM study of electronic excited state structures and properties in solution".
Winter Modeling '08. Pisa, 17th December 2008
11. Guido Ciro Achille, Cappelli Chiara, Mennucci Benedetta, Cammi Roberto, Tomasi Jacopo,
"Julolidine-Malononitrile and IndolineDimethine-MalonoNitrile: a TDDFT-IEFPCM study of electronic excited state structures and properties in solution".
XIV ESCMQC. Isola d'Elba (Italy), 8th -12th September 2008

Data

07/07/2023

Luogo

Alessandria