

Laboratory of Theoretical Chemistry – Unit of Theoretical and Structural Physical-Chemistry – Namur Institute of Structured Matter – UNamur Chemistry supporting simulation and new materials

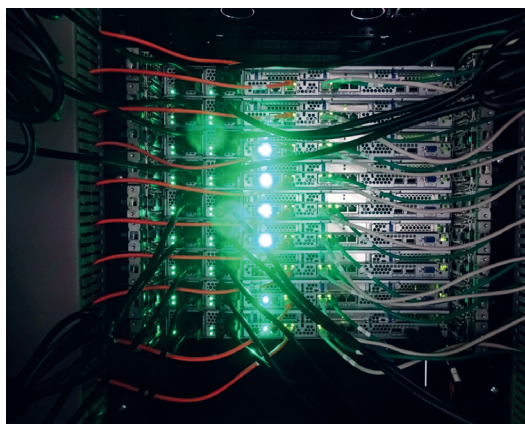
With about ten researchers, fifteen international publications a year and as many presentations at international congresses, the Laboratory of Theoretical Chemistry (LCT) of UNamur is currently leading about ten research projects.

Focused on multidisciplinary research and training for researcher, the LCT is developing theoretical chemistry methods and implement them in simulation codes/programs in order to understand the properties of matter, to deduce property-structure relationships, and to design new materials with target properties.

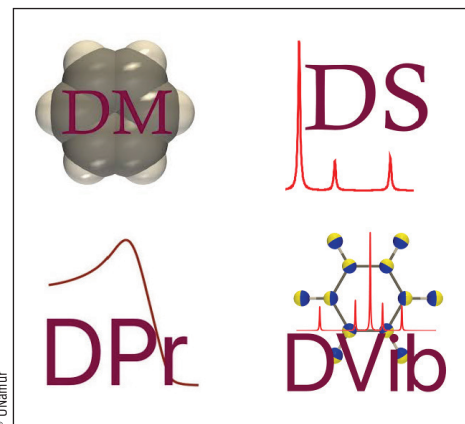
For this purpose, the main lines of research are molecules/materials for nonlinear optics, vibrational spectroscopies, electronics and molecular photonics. Within the context of a sustained collaboration with the University of Bordeaux (France), the LCT is studying molecular switches which have the potential to be active elements in sensors and in logical devices (at a molecular scale). The collaboration revealed methods of triggering molecular switches and generating high second-order nonlinear optical contrasts, opening the way to multiple applications in data storage and treatment (writing, reading without erasing, and erasing operations). In addition to the transformations brought about by the variation in pH or irradiation, the researchers have shown the capacity of intelligent molecules to recognize and distinguish cations (a prediction that is awaiting experimental confirmation). On the other hand, the demonstration that chiral vibrational spectroscopies can provide information on polymer chain structures has been confirmed by experiments.

Within the context of an ARC inter-university collaboration with the teams of UNamur and UCLouvain, LCT researchers are studying the phenomena of solid state photochromism and thermochromism. At national level, its participation in three successive projects in the context of the Belspo Interuniversity Attraction Poles, has led to a thesis under joint supervision with UHasselt. The subject is TADF (Thermally Activated Delayed Fluorescence) and it aims at enhancing the efficiency of organic light emitting diodes. They are also involved with a team from KULeuven working on fluorescent proteins, used for medical imagery.

The LCT has numerous research international partners: Bordeaux and Angers (France), Osaka (Japan), Krakow (Poland), Sherbrooke (Canada), Sfax and Tunis (Tunisia), Bologna (Italy), Calcutta (India), etc. The collaboration with Osaka University has proved to be particularly productive. First, it enabled the researchers to forecast in theory that compounds with intermediate diradical character should present high third-order nonlinear optical responses – which was confirmed by experiments some years later. From then on, it



Hercules2, HPC server of the PTCl (Technological platform for High-Performance Computing) of UNamur, is an integral part of the CÉCI (Consortium of Equipment of High-Performance Computing) of the Walloon-Brussels Federation.



DrawMol, DrawSpectrum, DrawProfile, and DrawVib graphical interfaces for representing and analyzing the structures and properties of molecules and crystals, as obtained from theoretical chemistry calculations

became possible to apply a magnetic field to modify the electronic and optical properties of these materials that are still not well known, even if the aspects of stability and implementation of these new phenomena still need to be fine-tuned.

At the moment the LCT is facing two challenges: first, that of taking into account several levels of approximation, both of size and time, in simulations and second the development of simulation codes that could be used reliably by non-specialists. This is the reason for the series of software programs DrawMol, DrawSpectrum, etc. developed by Dr V. Liégeois (FNRS researcher) and available on Mac platforms. Using this powerful graphic interface, you can visualize and construct molecular structures. This is a boon for extending scientific research to ordinary citizens!



Namur Institute of Structured Matter

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