Local approximation order strategy in a DGTD method for the simulation of nanoscale light-matter interactions

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Monthly stipend: ≈ 500 €

Nanostructuring of materials has opened up a number of new possibilities for manipulating and enhancing light-matter interactions, thereby improving fundamental device properties. Low-dimensional semiconductors, like quantum dots, enable one to catch the electrons and control the electronic properties of a material, while photonic crystal structures allow to synthesize the electromagnetic properties. These technologies may, e.g., be employed to make smaller and better lasers, sources that generate only one photon at a time for applications in quantum information technology, or miniature sensors with high sensitivity. The incorporation of metallic structures into the medium add further possibilities for manipulating the propagation of electromagnetic waves. In particular, this allows subwavelength localisation of the electromagnetic field and, by subwavelength structuring of the material, novel effects like negative refraction, e.g. enabling super lenses, may be realized. Nanophotonics is the recently emerged, but already well defined field of science and technology, which aims at establishing and using the peculiar properties of light and light-matter interaction in various nanostructures.

Numerical modeling of nanoscale light-matter interaction requires to solve the system of Maxwell equations possibly coupled to appropriate models of physical dispersion such as the Drude and Drude-Lorentz models. The Finite Difference Time-Domain (FDTD) method introduced by K.S. Yee in 1996 is a widely used approach for solving the resulting system of partial differential equations (PDEs). In this method, the whole computational domain is discretized using a structured (cartesian) grid. Due to the possible straightforward implementation of the algorithm and the availability of computational power, FDTD is often the method of choice for the simulation of time-domain nanoscale light-matter interaction problems. However, the space and time scales that characterize the underlying physical phenomena, in addition to the geometrical characteristics of the considered nanostructures, are particularly challenging for an accurate and efficient application of the FDTD method. During the last ten years, numerical methods formulated on unstructured meshes have drawn a lot of attention in computational electromagnetics with the aim of dealing with irregularly shaped structures and heterogeneous media. In particular, the discontinuous Galerkin time-domain (DGTD) method has progressively emerged as a viable alternative to the well established FDTD method.

Nachos is a common project-team between Inria and the J.A. Dieudonné Mathematics Laboratory at University Nice Sophia Antipolis. The team gathers applied mathematicians and computational scientists who are collaboratively undertaking research activities aiming at the design, analysis, development and application of innovative numerical methods for systems of partial differential equations (PDEs) modeling nanoscale light-matter interaction problems. In this context, the team is developing the DIOGENeS software suite (https://diogenes.inria.fr/), which implements several Discontinuous Galerkin (DG) type methods tailored to the systems of PDEs modeling of time- and frequency-domain nanophotonics. In the time-domain case, the DGTD method is a discontinuous finite element type that relies on a high order interpolation of the electromagnetic field components within each cell of discretization mesh. This piecewise polynomial numerical approximation is allowed to be discontinuous from one mesh cell to another, and the consistency of the global approximation is obtained thanks to the definition of appropriate numerical traces for imposing the continuity of the tangential fields on faces shared by two neighboring cells. Time integration is achieved using an explicit scheme and, as a result of the discontinuity of the approximation, no global mass matrix inversion is required to advance the solution at each time step. Such a DGTD method, which is formulated on an unstructured tetrahedral mesh, has been designed in the context of the Ph.D thesis of Jonathan Viquerat for the simulation...
of nanoscale light-matter interaction problems [Viq16]. This DGTD method has been recently refactored and implemented in the object-oriented framework of the DIOGENeS software suite.

In its most usual form, the formulation of the DGTD method is derived assuming a uniform distribution of the polynomial order in the cells of the underlying tetrahedral mesh. However, in the case of a mesh showing large variations in cell size, the time step imposed by the smallest cells can represent a serious hindrance when trying to exploit high approximation orders. Indeed, a potentially large part of the CPU time is spent in the update of the physical field inside small cells where high polynomial orders might not be essential, while they are necessary in the larger cells. To overcome this limitation, several strategies can be considered. In [Viq16] (see also [VL16]), a first approach has been investigated that relies on the use of non-uniform distribution of the polynomial order in the framework of DGTD method that makes use of a global time step that is constrained by a CFL condition. By imposing low orders in small cells and high orders in large cells, it is possible to significantly alleviate both the global number of degrees of freedom and the time step restriction with a minimal impact on the method accuracy. Strategies exploiting locally adaptive formulations usually combine both $h$- and $p$-adaptivity (where $h$ denotes the discretization parameter in space and $p$ the degree of the interpolation of the field components) in order to concentrate the computational effort in the areas of high field variations.

Starting from this preliminary work on a local approximation order DGTD method, the objectives of this internship will be the following:

1. The DGTD method currently implemented in the DIOGENeS software suite assumes a uniform distribution of the polynomial interpolation order. The first objective will thus be to extend this implementation in order to allow for a non-uniform distribution of the polynomial interpolation order coupled to the assignment strategy proposed in [VL16]. Preliminary validations will be performed by simulating a model electromagnetic wave propagation problem;

2. Starting from the strategy proposed in [VL16], the second objective will be to investigate alternative a priori local adaptation strategies. This part of the work will include a bibliographical review of related existing contributions on this topic. The implementation work undertaken in the first part will be extended to take into account the selected or proposed a priori local adaptation strategies.

3. Finally, the third objective of this study will consist in the numerical validation and (accuracy and performance) evaluation of the developed a priori local adaptation strategies by considering test problems relevant to nanophotonics.

Required background:

- Master in applied mathematics or scientific computing;
- Knowledge of finite element type methods for solving PDE;
- Software development skills, preferably in Fortran 200x.

References
