FELIX-1.0: A finite element solver for the time dependent generator coordinate method with the Gaussian overlap approximation

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Abstract
We describe the software package FELIX that solves the equations of the time-dependent generator coordinate method (TDGCM) in \(N\)-dimensions \((N \geq 1)\) under the Gaussian overlap approximation. The numerical resolution is based on the Galerkin finite element discretization of the collective space and the Crank-Nicolson scheme for time integration. The TDGCM solver is implemented entirely in C\textsuperscript{++}. Several additional tools written in C\textsuperscript{++}, Python or bash scripting language are also included for convenience. In this paper, the solver is tested with a series of benchmarks calculations. We also demonstrate the ability of our code to handle a realistic calculation of fission dynamics.

Keywords: FELIX; Finite element method; Generator coordinate method; Gaussian overlap approximation; Nuclear fission;

PROGRAM SUMMARY/NEW VERSION PROGRAM

Program Title: FELIX-1.0

Journal Reference:
Catalogue identifier:
Licensing provisions:
Programming language: C\textsuperscript{++}
Computer: Intel Xeon, Intel Core
Operating system: LINUX

RAM: Memory usage depends on the number of nodes in the calculation mesh as well as on the degree of the interpolation polynomials. For a 1D calculation with linear polynomials on a mesh with 600 nodes, memory usage is approximately 3.3 MB; in a realistic simulation of fission on a 2D mesh with quadratic polynomials and 1.3 \(10^7\) nodes, it reaches 1.5 GiB.

Number of processors used:
The code is multi-threaded based on the OpenMP API specification for parallel programming. Any number of threads may be specified by the user.

Keywords: FELIX; Finite element method; Generator coordinate method; Gaussian overlap approximation; Nuclear fission;

Classification: 17.23 Fission and Fusion Processes

External routines/libraries: The solver itself requires the BLAS and LAPACK libraries, and a Fortran compiler with OpenMP support. Building the documentation requires DoxyGen-1.8.6 or higher. Building the full set of tools also requires GSL, PETSc, SLEPc and Boost. In particular, environment variables PETSC\_DIR, PETSC\_ARCH, SLEPC\_DIR and SLEPC\_ARCH must be set.

Nature of problem:
Nuclear fission is a relatively slow process compared to the typical timescale of the intrinsic motion of the nucleons. In the adiabatic approximation, it can be described as a large amplitude collective motion driven by only a few collective degrees of freedom. In the time-dependent generator coordinate method (TDGCM), the nuclear wave-function is thus described as a time-dependent, linear superposition of basis functions in this collective space. Further assuming a Gaussian overlap approximation (GOA) for the basis functions, the time-dependent Schrödinger equation can be reduced into a local, time-dependent, Schrödinger-like equation in collective space. This is the TDGCM+GOA equation. Scission configurations are defined as a hyper-surface in the \(N\)-dimensional collective space. Fission fragment distributions are then computed by integrating over time the flux of the collective wave-packet across the scission hyper-surface. This microscopic approach to fission fragment distributions is fully quantum-mechanical.

Solution method:
FELIX solves the TDGCM+GOA equation by using the Galerkin finite element method to discretize the \(N\)-dimensional collective space, and the Crank-Nicolson scheme to solve for the time evolution. At each time step, this procedure requires solving a linear system of equation involving sparse, complex, symmetric matrices. FELIX employs an iterative QMR algorithm to perform matrix inversion.

Restrictions:
Although the program can operate in an arbitrary number of dimensions \(N\), it has only been tested in practice on 1, 2 and 3 dimensional meshes.

Unusual features:

Additional comments:
The code has checkpointing capabilities: the collective wave-function, norm \(a\) and energy kernels are stored on disk every \(n\) iterations, ensuring that the program can resume where it stops.

Running time:
Running time grows linearly with the number of time-steps requested by the user. It is also highly dependent on the number of nodes in the...
space mesh. Two periods of a 1D harmonic oscillator (600 nodes, 800 time steps) are typically computed in a few seconds on one thread of an Intel(R) Core(TM) i5 CPU. A 2-dimensional realistic case of fission (10^3 nodes, 10^5 time steps) requires roughly 10 hours on 10 threads of an Intel Xeon EP X5660 processor.

1. Introduction

Induced nuclear fission plays an essential role in important societal applications ranging from stockpile science to critical assemblies for new generation nuclear reactors [1]. It is also one of the leading mechanisms determining the stability of super-heavy elements and the end point of nucleosynthesis [2]. Many of these applications require the detailed knowledge of fission product yields (FPY), which may include the charge, mass, kinetic energy and excitation energy distribution of the fission fragments. In spite of recent technological advances, FPY measurements are not always possible, especially in very short-lived neutron-rich or heavy nuclei. Predictions based on theoretical models of fission are thus unavoidable.

While there exists a number of powerful phenomenological or semi-microscopic models on the market, the consensus is that a truly predictive theory of fission should ultimately be based only on our knowledge of nuclear forces and quantum many-body methods. In this context, most of the effort in the last decades has focused on describing the dynamics of induced fission in the time-dependent generator coordinate method (TDGCM) associated with the Gaussian overlap approximation (GOA) [3, 4, 5]. Under these assumptions, the original time-dependent many-body Schrödinger equation is reduced to a local Schrödinger-like equation that depends on only a few relevant collective variables. This approach was able to predict the characteristic times of low-energy induced fission [6, 7]. More recently, it was also successfully used to provide the first estimate of the mass and kinetic energy distributions of fission fragments for neutron-induced fission in the actinides region [8, 9, 10].

Until now, the aforementioned calculations have been based on the discretization of the TDGCM equations using finite differences in a regularly meshed hyper-cube. Given the computational resources required by this simple scheme, fission dynamics has only been studied in 2-dimensional collective spaces. Yet, it is well-known from both semi-phenomenological and fully microscopic approaches that at least four or five collective variables play a role in the dynamics of fission [11, 12, 13, 14]. Although possible in theory, extending the current scheme to $N > 2$ collective spaces would be prohibitively computationally. Similarly, increasing the fidelity of the calculation from $N_0$ to $N$ points for all $N$ dimensions scales approximately like $(n/N_0)^N$. A lot of this increase in computational cost would be wasted in regions of the collective space far away from the scission configurations, where the resolution does not need to be very high. For these reasons, it is highly desirable to move to a more flexible, more scalable discretization scheme.

In this work, we thus introduce the code FELIX, which implements the Galerkin finite element method for the discretization of the TDGCM equation. This well-known method allows the use of irregularly-spaced meshes in the collective space, which results in turn in calculations scaling much more efficiently with the number of collective variables. In addition, standard p-refinement techniques, i.e., the use of higher-degree polynomial bases in each finite element, give a better control on the numerical precision of the calculations. Finally, there are virtually no restrictions in the number of collective variables used. The code FELIX has been tested up to $N = 3$.

After a brief introduction to the TDGCM+GOA equations in section 2, we present in section 3 the numerical implementation in the code FELIX. The implementation is validated through a series of simple benchmarks discussed in section 4. The convergence of a realistic calculation of the fission of $^{240}$Pu is demonstrated in section 5. Finally, sections 6 and 7 give practical information on how to install and use the code FELIX.

2. Fission dynamics in the TDGCM+GOA approach

In this section, we briefly recall how to obtain a collective, Schrödinger-like equation to describe low-energy nuclear dynamics. We also explain how fission fragment distributions can be extracted from the integration of the collective flux across the hyper-surface defining the scission configurations.

2.1. The TDGCM+GOA equation

We recall that the time evolution of a many-body quantum system is given by the time-dependent Schrödinger equation, which is obtained from the variation $\delta S[\Psi] = 0$ of the quantum mechanical action given by [15]

$$S[\Psi] = \int_0^t \langle \Psi(t) | \left[ \hat{H} - \frac{\hbar^2}{2 m} \right] | \Psi(t) \rangle, \quad (1)$$

where $|\Psi(t)\rangle$ is the full many-body wave-function for the system. In most nuclear physics applications, the nuclear Hamiltonian $\hat{H}$ contains an effective two-body potential such as, e.g., the Skyrme or Gogny interaction. In the time-dependent generator coordinate method (TDGCM), the nuclear many-body wave function $|\Psi(t)\rangle$ takes the form $[3, 4, 5]$

$$|\Psi(t)\rangle = \int dq \ f(q, t)|\Psi(q)\rangle. \quad (2)$$

The functions $|\Psi(q)\rangle$ are known many-body states parametrized by a vector of collective variables $q$. In the context of fission, the $|\Psi(q)\rangle$ are chosen as the solutions to the static Hartree-Fock-Bogoliubov (HFB) equations under a set of constraints $q$. These constraints, which are the collective variables driving the fission process, can be expectation values of multipole moments, quantities related to pairing such as particle number fluctuations $\Delta N^2$, etc. Recall that the HFB solutions at point $q$ are entirely characterized by the one-body density matrix $\rho$ and two-body pairing tensor $\kappa$.

Inserting the ansatz of Eq.(2) in the variational principle (1) yields the (time-dependent) Hill-Wheeler equation. In contrast to the static case, there has been no attempt so far to solve
the time-dependent Hill-Wheeler equation numerically, as the computational resources needed are beyond current capabilities. Instead, a widespread approach consists in assuming that the norm kernels \( \langle \Psi(q)|\Psi(q') \rangle \) can be approximated by a Gaussian form factor \[ \[16\]. Inserting this Gaussian overlap approximation (GOA) into the Hill-Wheeler equation (using a second order expansion in \( q - q' \)) leads to a local, time-dependent, Schrödinger-like equation in the space \( Q \) of collective coordinates \( q \).

\[
i \hbar \frac{\partial}{\partial t} g(q, t) = \left[ -\frac{\hbar^2}{2} \sum_{kl} \frac{\partial}{\partial q_k} B_{kl}(q) \frac{\partial}{\partial q_l} + V(q) \right] g(q, t),
\]

(3)

where

- The function \( g(q, t) \) is complex. It is related to the weight function \( f(q, t) \) appearing in Eq.\( (2) \) and contains all the information about the dynamics of the system. Moreover, the quantity \( |g(q, t)|^2 \) can be interpreted as the probability density for the system to be in the state \( |\Psi(q)\rangle \) at time \( t \); see also section 2.2 below.

- The real scalar field \( V(q) \) and the real symmetric tensor field \( B_{kl}(q) \) are fully determined by the knowledge of the effective Hamiltonian \( \hat{H} \) and the generator states \( |\Psi(q)\rangle \). They reflect the static nuclear properties of the system under study.

Throughout this paper, equation (3) will be referred to as the TDGCM+GOA equation.

2.2. Collective flux and fission fragment distributions

The TDGCM+GOA equation implies a continuity equation for the probability density \( |g(q, t)|^2 \),

\[
\frac{\partial}{\partial t} |g(q, t)|^2 = -\nabla \cdot J(q, t).
\]

(4)

The real vector field \( J(q, t) \) is thus a current of probability. It can be expressed formally as a function of the collective wave-function,

\[
J(q, t) = \frac{\hbar}{2i} B(q) \left[ g^*(q, t) \nabla g(q, t) - g(q, t) \nabla g^*(q, t) \right].
\]

(5)

Specifically, the coordinates of the current of probability read

\[
J_k(q, t) = \frac{\hbar}{2i} \sum_{l=1}^N B_{kl}(q) \left[ g^*(q, t) \frac{\partial g_k(q, t)}{\partial q_l} - g(q, t) \frac{\partial g^*_k(q, t)}{\partial q_l} \right].
\]

(6)

As our system evolves in time, its density probability will flow starting from the area of the collective space \( Q \) where the initial wave-function was localized. This evolution is driven by the Hamiltonian \( \hat{H} \) through the inertia tensor \( B_{kl}(q) \) and the potential energy surface \( V(q) \).

In the case of fission, the potential energy surface is computed up to the points \( q \) where the nuclear geometry corresponds to two well-separated fragments. One can thus partition the space \( Q \) into a region where the nucleus is whole – the internal region, and another where it has split in two fragments – the external region. The hyper-surface separating the two regions corresponds to the set of scission configurations. The rigorous definition and accurate determination of these scission configurations are themselves challenging problems, which go beyond the scope of this paper; see Refs.\[ \[17, 18, 19, 12, 20, 14\] for additional discussions. For practical calculations of fission fragment distributions with FELIX we will simply assume the existence of such a scission hyper-surface.

In general, the local, one-body density matrix \( \rho(r) \) in each of the scission points in the collective space \( Q \) is characterized by two high-density regions separated by a thin neck. Assuming the neck is located along the \( z \)-axis of the intrinsic reference frame, the charge and mass of each fragment can be obtained by simple integration of \( \rho(r) \) over the domains \( z \in [-\infty, z_N] \) and \( z \in [z_N, +\infty] \); see, e.g., \[ \[19, 12, 14\]. According to this procedure, one can associate with each point \( q \) of the scission hyper-surface a pair of fragment masses. It follows that the flux of the probability current (6) through the scission hyper-surface gives a very good estimate of the relative probability of observing a given pair of fragments at time \( t \). We thus define the integrated flux \( F(\xi, t) \) through an oriented surface element \( \xi \) as

\[
F(\xi, t) = \int_{t=0}^t d\xi \int_{q(\xi,t)} J(q, t) \cdot dS.
\]

(7)

Following \[ \[7, 10\], the fission fragment mass yield for mass \( A \) is defined formally as

\[
Y(A) \propto \sum_{\xi} \lim_{t \to +\infty} F(\xi, t),
\]

(8)

where \( A \) is the set of all oriented hyper-surfaces \( \xi \) belonging to the scission hyper-surface such that one of the fragments has mass \( A \). In practice, our calculation of the fragments mass number produces non integer values. Moreover, one elementary surface \( \xi \) may contain several fragmentations. In this work, we equally distribute the flux component \( F(\xi, t) \) between the masses calculated at the vertices of the edge \( \xi \):

\[
Y(A) = C \sum_{\xi \in A} \frac{1}{N} \sum_{r \in R(\xi)} \lim_{t \to +\infty} F(\xi, t),
\]

(9)

The sum on \( \xi \) runs on the whole scission hyper-surface. The set \( R(\xi) \) contains the vertices of \( \xi \) at which one of the fragments has a mass in the interval \([A - 1/2; A + 1/2]\). The normalization constant \( C \) is chosen as usual such that

\[
\sum_{A=0}^{\text{num}} Y(A) = 200.
\]

(10)

In practice, the flux is only integrated from \( t = 0 \) to \( t = t_{\text{max}} \). Equations (6)-(10) show how to extract fission fragment yields from the knowledge of the collective wave function \( g(q, t) \) solution to the TDGCM+GOA equations.

3. Numerical methods

In this section, we detail the numerical methods implemented in the code FELIX to solve the TDGCM+GOA equation (3) and calculate the flux defined in Eq. (7).
3.1. Restriction to a finite domain of space

FELIX solves Eq. (3) in a finite domain $\Omega$ of the collective space $Q$. To ensure the uniqueness of the solution, Dirichlet conditions are imposed at the boundary $\partial \Omega$ of the domain,

$$\forall q \in \partial \Omega : \quad g(q, t) = 0. \tag{11}$$

Imposing this condition is justified as long as the actual solution $g(q, t)$ is well confined inside the domain $\Omega$ during the whole time evolution of the system. In practice, this may require choosing an excessively large domain $\Omega$. In the case of fission for example, only the internal region discussed in section 2.2 and its interface with the external zone present a physical interest. However, we cannot limit $\Omega$ to this area because the probability to observe the fissioning system outside of this configuration subset is not negligible.

To circumvent this issue, FELIX defines an absorption band along the boundary $\partial \Omega$. This band artificially simulates the leakage of the wave packet $g(q, t)$ outside of the calculation domain. Formally, absorption is taken into account by introducing a new imaginary term in the evolution equation,

$$\forall q \in \Omega, \ t \in [0, t_{\text{max}}] : \quad \imath h \frac{\partial}{\partial t} g(q, t) = \left[ -\frac{\hbar^2}{2} \sum_{kl} \frac{\partial}{\partial q_k} B_{kl}(q) \frac{\partial}{\partial q_l} + V(q) - \imath \hbar A(q) \right] g(q, t). \tag{12}$$

The real scalar field $A(q)$ is non zero only in the absorption band. In this region, $A(q)$ is taken as a simple polynomial increasing smoothly from 0 on the inner border of the band and reaching its maximum at the boundary of the domain,

$$A(q) = 4r \left(1 - \frac{x(q)}{w}\right)^3. \tag{13}$$

The quantity $x(q)$ is the minimal Euclidean distance between the point $q$ and the boundary $\partial \Omega$. The parameters $r$ and $w$ correspond to the average absorption rate and width of the absorption band respectively. These two parameters can be tuned by the user as a function of the problem characteristics to ensure optimal absorption.

3.2. Space discretization

As mentioned earlier, we use the Galerkin finite element method [21, 22] to discretize the collective space $Q$. The main reasons for choosing this approach are its capability to manage non regular meshes and the possibility to apply h-refinement and p-refinement techniques to improve computational efficiency. In this section, we show how to formally derive a linear system of equation from the discretization of Eq. (12).

As customary in quantum mechanics, we note $\langle \cdot | \cdot \rangle$ the scalar product in the space $L^2(\Omega, \mathbb{C})$ of complex-valued, square-integrable functions,

$$\langle \phi | \phi \rangle = \int_{\Omega} dq \ \phi^*(q) \phi(q). \tag{14}$$

With this definition, Eq. (12) can be recast into

$$\forall \phi \in L^2(\Omega, \mathbb{C}), \ \forall t \in [0, t_{\text{max}}] : \quad \langle \phi | r(t) \rangle = 0, \tag{15}$$

with the residual $r(q, t)$ defined as

$$r(q, t) = \left[ -\frac{\hbar^2}{2} \sum_{kl} \frac{\partial}{\partial q_k} B_{kl}(q) \frac{\partial}{\partial q_l} + V(q) - \imath \hbar A(q) \right] g(q, t). \tag{16}$$

Following the standard approach of the finite elements method, the domain $\Omega$ is first partitioned into a mesh. In our case, each cell of the mesh is a $N$-dimensional simplex (triangle if $N = 2$, tetrahedron if $N = 3$, etc.). We note $S$ the set of all simplices in the domain. Inside every simplex of the mesh, we assume a polynomial form for the numerical solution of Eq. (12). At any time $t$ and in any simplex $s \in S$, we thus define the local interpolating polynomial $P_s$.

$$\forall s \in S, \forall q \in s : \quad g(q, t) = P_s(q). \tag{17}$$

For each simplex $s \in S$, we select the degree $d_s$ of the interpolating polynomial. The space $P_s$ of all interpolating polynomials in the simplex $s \in S$ is a vector space. Its dimension $D_s$ is given by the binomial coefficient,

$$D_s = \binom{N + d_s}{d_s}. \tag{18}$$

In order to discretize Eq. (12), we now build a convenient basis of the space $P_s$. First, we define for each simplex $s \in S$ a finite set of specific points $q_i \in s$ called nodes. Next, we introduce a set of real polynomials $\phi_{s,i}$ associated with the simplex $s$. For all nodes $i$ of the simplex $s$, the polynomial $\phi_{s,i}$ is defined by the requirement

$$\phi_{s,i}(q_j) = \begin{cases} 1, & \text{if } i = j \\ 0, & \text{if } i \neq j \end{cases} \tag{19}$$

In other words, the $\phi_{s,i}$ are the usual Lagrange polynomials. The total number of nodes in each simplex $s \in S$ is equal to $D_s$, so that the set $\{\phi_{s,i}\}_{i=1}^{D_s}$ forms a basis of $P_s$. The total number of nodes in the entire domain $\Omega$ is noted $m$.

With the help of these local bases, we can define for each of the $m$ nodes $i$ of the domain $\Omega$ a function $\phi_i$ such that

$$\forall q \in \Omega : \quad \phi_i(q) = \begin{cases} \phi_{s,i}(q), & \text{if } q \in s \text{ and } i \text{ is a node of } s \\ 0, & \text{otherwise} \end{cases} \tag{20}$$

The functions $\{\phi_i\}_{i=1}^m$ form a basis of our solution space. The solution of Eq. (12) can thus be expanded as

$$g(q, t) = \sum_{i=1}^{m} g(q_i, t) \phi_i(q). \tag{21}$$

Applying the Galerkin finite element method, we search for a numerical solution $g(q, t)$ of the form (21) that verifies

$$\forall t \in [0, t_{\text{max}}], \forall i \in [1, m] : \quad \langle \phi_i(q) | r(q, t) \rangle = 0. \tag{22}$$
This process yields a discretized system of $m$ equations with the $m$ coefficients $g(q_j,t)$ as the unknown. It can be written in the condensed form

$$i\hbar M \frac{\partial G(t)}{\partial t} = [H - i\hbar A]G(t),$$  \hspace{1cm} (23)$$

where $G(t)$ denotes the $m$-dimensional vector of coefficients $g(q_j,t)$ at every node $j$ of the domain $\Omega$. The $m \times m$ matrices $M, H$ and $A$ are defined by

$$M_{ab} = \langle \phi_a(q) | \phi_b(q) \rangle,$$

$$A_{ab} = \langle \phi_a(q) | A(q) | \phi_b(q) \rangle,$$

$$H_{ab} = \langle \phi_a(q) \rangle \left[ -\frac{\hbar^2}{2} \sum_{kl} \frac{\partial}{\partial q_k} B_{ik}(q) \frac{\partial}{\partial q_l} + V(q) \right] | \phi_b(q) \rangle.$$  \hspace{1cm} (24)$$

All matrix elements can be computed by applying the basis expansion (21) to the fields $V(q)$, $B_{ik}(q)$ and $A(q)$, that is,

$$F(q) = \sum_{c=1}^{m} F(q_c) \phi_c(q) \quad \text{with } F = V, B_{kl}, \text{ or } A.$$  \hspace{1cm} (25)$$

The double derivative term in $H_{ij}$ can be integrated by parts using the Dirichlet conditions imposed on the boundary $\partial \Omega$. The final expression for the matrix elements is

$$M_{ab} = \int_\Omega dq \; \phi_a(q) \phi_b(q),$$

$$A_{ab} = \sum_{c=1}^{m} A(q_c) \int_\Omega dq \; \phi_a(q) \phi_b(q) \phi_c(q),$$

$$H_{ab} = \sum_{c=1}^{m} V(q_c) \int_\Omega dq \; \phi_a(q) \phi_b(q) \phi_c(q)$$

$$- \sum_{c=1}^{m} \sum_{kl} B_{ik}(q_c) \int_\Omega dq \; \phi_a(q) \frac{\partial \phi_b(q)}{\partial q_k} \frac{\partial \phi_c(q)}{\partial q_l}.$$  \hspace{1cm} (26)$$

Since the basis functions $\phi_a(q)$ are simple polynomials of $q$, integrations can be performed analytically, and no quadrature or numerical integration scheme is needed. Note that the three matrices obtained here are real, symmetric and sparse. The sparsity comes from the fact that the overlap between any two basis functions is zero unless at least one element was defined with both the corresponding nodes.

The values of each field at each node are inputs of the calculation. To compute the matrix elements, FELIX relies on a formal representation of the polynomials. Basis elements $\phi_{sij}$ are first derived from Eq. (19). This step requires inverting a small dense linear system for each simplex. Then derivatives, multiplications and integrations of the polynomials involved in (Eq. 26) can all be performed formally, so that these operations do not generate other errors than those related to the accuracy of the polynomial coefficients.

### 3.3. Time discretization

The Crank-Nicolson scheme is used to discretize Eq. (23) in time [23]. We recall that the Crank-Nicolson scheme gives the following prescriptions for the function and its time-derivative,

$$\frac{\partial G}{\partial t} \approx \frac{G(t + \Delta t) - G(t)}{\Delta t}, \quad G(t) \approx \frac{G(t) + G(t + \Delta t)}{2}.$$  \hspace{1cm} (27)$$

Starting from Eq. (23), this numerical scheme yields the fully discretized equation

$$R \times G(t + \Delta t) = b(t),$$  \hspace{1cm} (28)$$

with

$$R = M + \frac{\Delta t}{2} A + i \frac{\Delta t}{2\hbar} H,$$

$$b(t) = \begin{bmatrix} M - \frac{\Delta t}{2} A - i \frac{\Delta t}{2\hbar} H \end{bmatrix} G(t).$$  \hspace{1cm} (29)$$

Using this time discretization scheme, we can show that both the norm of the collective wave-function and the average energy computed from the numerical solution are both constants in time if the absorption term is set to zero

$$\|g(q,t)\|_2 = \left[ \int_\Omega \left| g^\ast(q,t) g(q,t) \right| \right]^{1/2} = \text{cst},$$

$$\int_\Omega \left| g^\ast(q,t) \left[ -\frac{\hbar^2}{2} \sum_{kl} \frac{\partial}{\partial q_k} B_{ik}(q) \frac{\partial}{\partial q_l} + V(q) \right] g(q,t) \right| = \text{cst}.$$  \hspace{1cm} (30)$$

These properties can be used to test the validity of the numerical implementation.

### 3.4. Inversion of a linear system

The collective wave-function at time $t$ is obtained by solving the fully-discretized Eq. (28) for the vector $G(t)$. This requires inverting at each time step a complex, sparse $m \times m$ matrices. In FELIX these inversions are computed with the iterative QMR algorithm without look-ahead as described in [24]. The numerical solution at iteration $n$ is used as the initial guess for iteration $n + 1$. The convergence criterion to stop the iterations for each inversion is defined by

$$\|R \times G(t + \Delta t) - b\|_2 < \epsilon \|b\|_2,$$  \hspace{1cm} (31)$$

with the tolerance $\epsilon$ specified by the user. In order to accelerate the inversion, a Jacobi preconditioner is applied to the system before the first time iteration.

Since no look-ahead statement is implemented, the QMR algorithm may occasionally fail to converge at the level of precision required. In such case, the system is rewritten in the $(2m \times 2m)$ real form

$$\begin{bmatrix} \begin{bmatrix} M + \frac{\Delta t}{2} A \end{bmatrix} & -\frac{\Delta t}{\hbar} H \\ \frac{\Delta t}{\hbar} H & \begin{bmatrix} M + \frac{\Delta t}{2} A \end{bmatrix} \end{bmatrix} \begin{bmatrix} \Re \{G(t + \Delta t)\} \\ \Im \{G(t + \Delta t)\} \end{bmatrix} = \begin{bmatrix} \Re \{b(t)\} \\ \Im \{b(t)\} \end{bmatrix},$$  \hspace{1cm} (32)$$

where $\Re$ and $\Im$ refer to the real and imaginary parts, respectively. This real system is then solved with the Bi-conjugate Gradient Stabilized Method as described in [25].

In practice, the small numerical errors caused by these matrix inversions accumulate over time, and can lead to violations of the properties (30). However, this numerical error is very small, especially if the time span of the time iterations is reasonable and the tolerance $\epsilon$ is small enough.
3.5. Calculation of the flux

In FELIX, hyper-surfaces are defined as the union of oriented faces of an arbitrary list of simplices in the mesh. Note that the hyper-surfaces thus defined are not necessarily connected, as the simplices need not be adjacent. Given such a hyper-surface provided by the user, the code can compute the flux $F(\xi)$ as defined in Eq. (7) through each of the faces $\xi$.

The instantaneous elementary flux $f$ going through an oriented simplex face $\xi$ at time $t$ is calculated as

$$f(\xi, t) = \sum_{k=1}^{N} n_k^{(\xi)} \cdot \int_{q \in \xi} J_k(q, t) dS$$  \hspace{1cm} (33)

where $n_k^{(\xi)}$ is the unit vector normal to the simplex face $\xi$, $J_k(q, t)$ is the probability current (6) and $N$ is the dimension of the collective space \(^1\). In order to compute the instantaneous flux, we expand the collective wave function $\Psi(q, t)$ and the inertia tensor field $I(q, t)$ on the FE basis using Eqs. (21)-(25). The integral in the flux becomes

$$\int_{q \in \xi} J_k(q, t) dS = h \sum_{l=0}^{N} \sum_{w, v, u} B_{kl}(q_w) \times [\Re(G_w(t))\Im(G_v(t)) - \Im(G_w(t))\Re(G_v(t))] I_{\xi,u,v,w,l},$$  \hspace{1cm} (34)

with

$$I_{\xi,u,v,w,l} = \int_{q \in \xi} \phi_u(q) \phi_v(q) \frac{\partial \phi_w(q)}{\partial q_l} dS.$$  \hspace{1cm} (35)

The integral $I_{\xi,u,v,w,l}$ is the numerical tolerance $\epsilon$ for matrix inversions; see Eq. (31);

- the mesh size $h$, which provides an estimate of the “spatial” resolution of the domain $\Omega$;
- the degree $d_s$ of basis polynomials;
- the time step $\delta t$ used in time integration.

4.1. Conservation of the Norm

The conservation of the norm expressed by Eq.(30) is the simplest test of our implementation. In this benchmark, the calculation domain $\Omega$ is a 3D cube of size 10 arbitrary units (a.u.). The mesh is built by creating a regular grid of equidistant vertices with a mesh size $h = 1$ a.u. The position of each vertex is then randomly perturbed. A new coordinate $q'_k$ is sampled uniformly in the interval $[q_k - fh; q_k + fh]$, where $fh$ is a fluctuation factor set to 15%, and $q_k$ is the associated old coordinate. Once the vertices are defined, the simplices are determined by Delaunay triangulation. In each simplex, a polynomial basis of degree $d_s = 1$ is used and the nodes are positioned exactly at the vertices.

Based on this mesh, a free wave packet is propagated during a time $t_{\text{max}} = 15$ a.u.. The inertia tensor is diagonal, inversely proportional to a mass $m = 1.3$ a.u. and independent of space. The initial wave packet is a 3D isotropic Gaussian centered in the middle of the simulation domain and characterized by the width $\sigma = \sqrt{\hbar/1.04}$ a.u.. Finally, we use the time step $\delta t = 5 \times 10^{-4}$ a.u., and the numerical tolerance for matrix inversions is set to its default value of $10^{-15}$.

The error $e_{\text{norm}}$ of the norm is defined as

$$e_{\text{norm}} = \frac{||g(q, t)||_2 - ||g(q, 0)||_2}{||g(q, 0)||_2}$$  \hspace{1cm} (36)

This quantity is computed at different times of the simulation and is plotted in figure 2. We note that the error is maintained below $10^{-12}$ during the whole simulation with this choice for the numerical tolerance of matrix inversions. This is consistent with the expected property of Eq. (30). As discussed in section 3.4 the error $e_{\text{norm}}$ comes from the accumulation of errors.

\(^1\)This definition is only valid if the collective space has a dimension strictly superior to one. The flux calculation is not enabled in FELIX in the case of 1-dimensional spaces.
4.2. Harmonic oscillator potential

We now turn to the dynamics of a quantum system in an isotropic harmonic oscillator (HO) potential in \( N = 1 \) and \( N = 2 \). The advantage of the HO potential is that it provides analytical solutions that can be used to test the implementation.

\[
\begin{array}{cccc}
m & \omega & \Omega & f_h \\
1.3 & 0.8 & [-20;20]^N & 0.15
\end{array}
\]

Table 1: Characteristics of the 1D and 2D harmonic oscillators used in this study.

In the following calculations, the inertia tensor is always diagonal, inversely proportional to a mass \( m \) and independent of space, \( B_{ij}(q) = \delta_{ij}/m \). The HO potential being isotropic, it is characterized by a single frequency \( \omega \),

\[
V(q) = \frac{1}{2}m\omega q^2. \tag{37}
\]

The numerical values adopted for the HO potential are listed in Table 1. All calculations are performed in a domain \( \Omega = [-20;20]^N \). The same procedure as described in section 4.1 is used to build an irregular mesh. Dirichlet boundary conditions are enforced at the boundaries \( \partial\Omega \) of the domain. On the other hand, we find that the analytic expressions for the first two eigenstates of the HO verify

\[
\forall q \in \partial\Omega : \frac{|g(q)|}{\max |g(q)|}_{q \in \Omega} < 2.10^{-13}. \tag{38}
\]

Therefore, the numerical error coming from the finite size of the domain of resolution \( \Omega \) is completely negligible compared to the other sources of error under study.

4.2.1. Ground state of the 1D HO

For any eigenstate of the potential, the modulus \(|g(q,t)|\) of the wave function is independent of time. We first test this property for the ground state of the 1D HO, \( q \equiv q \). The initial wave function is taken as

\[
g(Q, t = 0) = \exp\left(-\frac{Q^2}{2}\right), \tag{39}
\]

where the reduced coordinate \( Q \) is defined as

\[
Q = \sqrt{\frac{m\omega}{\hbar}}q. \tag{40}
\]

Calculation is performed up to \( t_{\text{max}} = 32 \) a.u., which is slightly larger than two periods of the complex function \( g(q, t) \). The deviation of the modulus of the numerical solution at time \( t \) from its initial value is measured by the error \( e_{\text{mod}} \),

\[
e_{\text{mod}} = \frac{||G(t_{\text{max}})|| - ||G(t = 0)||}{||G(t = 0)||}, \tag{41}
\]

where the infinity norm \( || \cdot ||_{\infty} \) for a vector \( G \) refers to the maximum absolute value of its elements. This error has been computed from a set of calculations with different mesh sizes \( h \) and time steps \( \delta t \). Results are presented in Table 2.

| \( \delta t/h \) | 1.0   | 0.1   | 0.01  |
|----------------|-------|-------|-------|
| \( 10^{-4} \)  | 8.089.10^{-4} | 1.039.10^{-5} | 7.460.10^{-7} |
| \( 10^{-1} \)  | 3.780.10^{-3} | 7.616.10^{-6} | 1.780.10^{-6} |
| \( 10^{-2} \)  | 3.688.10^{-3} | 2.603.10^{-5} | 2.601.10^{-6} |
| \( 10^{-3} \)  | 3.662.10^{-3} | 2.626.10^{-5} | 2.461.10^{-6} |
| \( 10^{-4} \)  | 3.661.10^{-3} | 2.619.10^{-5} | 2.601.10^{-6} |

Table 2: Error \( e_{\text{mod}} \) on the modulus \( |g(q,t)| \) for the ground-state of a 1D HO.

We verify that the error \( e_{\text{mod}} \) decreases when refining the calculation in both time and space. For a given mesh size \( h \), the error slightly fluctuates with \( \delta t \) before it finally reaches a converged value. The major part of the error is clearly driven by space discretization.

4.2.2. Sum of two eigenstates for the 1D HO

In this benchmark, the initial state of the system is a sum of the first two eigenstates of the 1D HO, and can thus be written

\[
g(Q, t = 0) = \exp\left(-\frac{Q^2}{2}\right)(1 + Q). \tag{42}
\]

The second eigenstate is characterized by a frequency three times larger than in the ground-state. Therefore, the motion is periodic with the same period as in section 4.2.1. In this case, the modulus of the wave function is not constant anymore, since the system is not in an eigenstate, but it oscillates between two
positions. The full time-dependent solution is given analytically by

\[ g(Q, t) = \exp\left(-\frac{Q^2}{2}\right)e^{-i\omega t/2}(1 + Qe^{-i\omega t}). \]  

We compare the real part of the numerical solution \( \Re \left( g(q, t) \right) \) to its analytic expression using the error \( e_R \) defined as

\[ e_R = \frac{\|\Re (g_{\text{num}}(t)) - \Re (g_{\text{the}}(t))\|_{\infty}}{\|\Re (g_{\text{the}}(t))\|_{\infty}}. \]

This error is computed for several time and space refinements and is plotted as a function of time step in figure 3. As in the previous benchmark, we observe that the numerical solution converges to its analytic value as time and space are refined.

\[ e_R \]

**Figure 3:** Error \( e_R \) on the real part of the solution for a periodic motion in a 1D HO.

### 4.2.3. **Sum of two eigenstates in 2D**

In this section, the previous study is generalized to the case of a 2D HO, \( q \equiv (q, q') \). This benchmark of a 2D case allows us to compare the numerical calculation of the flux with its analytic expression. The initial wave function reads

\[ g(Q, Q', t = 0) = \exp\left(-\frac{Q^2 + Q'^2}{2}\right)(1 + Q), \]  

where \( Q' \) is the reduced coordinate associated with \( q' \). Starting from this state, the full time-dependent solution reads

\[ g(Q, Q', t) = \exp\left(-\frac{Q^2 + Q'^2}{2}\right)e^{-i\omega t/2}(1 + Qe^{-i\omega t}). \]  

The system oscillates from one side of the line \( q = 0 \) to the other side with a period \( 2\pi/\omega \approx 31.4 \) a.u. The derivatives of the wave function read

\[ \frac{\partial g}{\partial q} = \sqrt{\frac{m\omega}{\hbar}}\exp\left(-\frac{Q^2}{2}\right)e^{-i\omega t}\left(Qe^{-i\omega t} - e^{-i\omega t}Q\right), \]

\[ \frac{\partial g}{\partial q'} = -\sqrt{\frac{m\omega}{\hbar}}\exp\left(-\frac{Q^2}{2}\right)e^{-i\omega t}Q'(1 + Qe^{-i\omega t}). \]

The instantaneous flux of Eq. (33) through an oriented segment \([AB]\), as depicted in figure 4, is given by

\[ f([AB], t) = -\sqrt{\frac{\hbar}{2m}}\sin(\theta)\sin(\omega t)\]  

\[ \times \left[ \exp\left(\frac{t}{2Q_{\theta}\sin\theta + Q'_{\theta}\cos\theta}\right)\right]^{2} \]  

\[ \times \left[ \frac{\theta}{t/2Q_{\theta}\sin\theta + Q'_{\theta}\cos\theta}\right]. \]

with the error function defined as usual by

\[ \text{erf}(z) = \frac{2}{\sqrt{\pi}}\int_{0}^{z} e^{-x^2} dx. \]  

**Figure 4:** Definition of the segment \([A,B]\)

**Figure 5:** Domain of resolution \( \Omega \) for the 2D HO.

To test our implementation, we first define an arbitrary frontier following an oscillating path around the line \( q = 0 \); see figure 5. The dynamics of the system is computed up to \( t_{\text{max}} = 16 \) a.u. At each vertex of the frontier, we calculate the quantities \( \Re (g) \) and \( \partial m(g)/\partial q \) at the end of the time evolution. In addition, our code provides the numerical value of the instantaneous flux through each element of the frontier. These three vectors of results obtained at the frontier are compared to their respective analytic expressions based on the error

\[ e_{\text{front}}(v) = \frac{\|v_{\text{num}} - v_{\text{the}}\|_{\infty}}{\|v_{\text{the}}\|_{\infty}}. \]  

These errors have been estimated for a time step \( \delta t = 5.10^{-4} \) and for the following set of different space discretizations,
• $h = 2, 1, 0.5, 0.2, 0.1$, with a polynomial basis of degree $d_e = 1$. Decreasing the mesh size $h$ is known as $h$-
refinement in the FE approach;

• $h = 2, 1, 0.5$, with a polynomial basis of degree $d_e = 2$. For these calculations, each simplex has 6 nodes, of which three are positioned at the vertices and the three others in the middle of each edge.

Note that for a same mesh size $h$, going from linear to quadratic polynomials doubles the number of nodes in the mesh. This is an example of $p$-
refinement in the FE approach.

![Image](6.png)

Figure 6: Numerical errors for the real part of the solution, $\Re(q)$, the spatial derivative of the imaginary part of the solution, $\partial \Im(q)/\partial q$, and the instantaneous flux as a function of the mesh size $h$ for an isotropic 2D HO.

The results obtained are summarized in figure 6. Although the results are only presented for one value of the time step $\delta t$, we obtained similar results with another series of calculations using $\delta t = 1.10^{-4}$. For the three quantities tested, the convergence to their analytic expression is numerically confirmed. We note that the convergence is much faster when using polynomials of degree two in the basis. Moreover, the error associated with the derivative of the solution is much larger than the one associated with the solution itself. In particular, it seems that the convergence rate of the flux is limited by the error on the derivative.

4.3. Stokes theorem

The goal of this benchmark is to test the calculation of the flux in a case where no analytic expression of the solution is available. This will be achieved by the use of the Stokes theorem of differential geometry. As is well known, if we define an enclosed volume $V \subset \Omega$, the continuity equation (4) yields the following conservation relation

$$\varepsilon_{\text{Stokes}} = \int_{t=0}^{t_{\\text{max}}} dt \int_{q \in \Omega} J(q, t) dS - \left[ \int_{q \in V} \left| g(q, t) \right|^2 dq \right]_{t=0}^{t_{\\text{max}}} = 0.$$  \hspace{1cm} (51)

We tested this property in the case of a 2-dimensional free wave packet. The simulation domain, the mesh, the frontier, and the inertia are the same as in section 4.2.3. The initial state is a Gaussian function centered at $q = (-5; 0)$ and having a width $\sigma = \sqrt{\hbar}/1.04$ a.u. This function is then multiplied by a plane wave of impulsion $k = (1; 0)$ and normalized to one.

We define the volume $V$ as one half of $\Omega$ delimited by the frontier $\mathcal{F}$ and containing the major part of the initial wave packet. In this configuration, the Dirichlet condition imposed on the boundary of the simulation domain imposes

$$\int_{q \in \partial V} J(q, t) dS = \sum_{\xi \in \mathcal{F}} F(\xi).$$  \hspace{1cm} (52)

The error $\varepsilon_{\text{Stokes}}$ on Eq. (51) can therefore be computed at any time from the flux and the numerical solution produced by FELIX.

In this benchmark, we run the simulation up to $t_{\\text{max}} = 3$ a.u.. At this time, most of the wave packet has crossed the frontier. The calculation is performed with a time step $\delta t = 10^{-4}$ and repeated for several space discretizations. The results are shown in figure 7. The property (51) is verified up to $0.1\%$ of the norm.

![Image](7.png)

Figure 7: Absolute deviation from the Stokes theorem $|\varepsilon_{\text{Stokes}}|$.

5. Application to the fission of $^{240}$Pu

To demonstrate the capability of FELIX, we show in this section the results obtained by solving the TDGCM+GOA equation for the neutron-induced fission of a $^{239}$Pu target. In particular, we emphasize the convergence of such a calculation in a realistic setting as well as several features used to increase computational efficiency.

5.1. Description of the calculation

We solve the TDGCM+GOA equation in the 2D collective space spanned by the quadrupole ($q_{20}$) and octupole ($q_{30}$) moments of the fissioning system. In the following, $q_{20}$ and $q_{30}$ are expressed in fm$^2$ and fm$^3$, respectively. The potential energy surface $V(q)$ of the compound nucleus $^{240}$Pu in this collective space is obtained by solving the HFB equations with a Skyrme energy density in the particle-hole channel and a surface-volume, density- dependent pairing energy density; see [14] for details. The potential field $V(q)$ is initially computed...
in a domain $\Omega_0$ characterized by $(q_{20}, q_{30}) \in [500; 60.10^3] \times [-92.10^3; +92.10^3]$ (in the units above). In practice, the HFB calculation did not converge for every point in the domain; the initial grid is thus irregularly spaced and contains 2784 fully converged points. At each point, the collective inertia tensor \( B^{(ij)}_{ij} \) is computed using the perturbative cranking approximation of the adiabatic time-dependent Hartree-Fock (ATDHF) theory [26]. We also determine at each point the expectation value \( q_N \) of the Gaussian neck operator, which will be used to determine the scission hyper-surface.

The initial collective state \( g(q, 0) \) is defined as the product of Gaussian functions in the \( q_{20^{-}} \) and \( q_{30^{-}} \) directions,

\[
g(q_{20}, q_{30}, 0) = \frac{1}{\sigma_{20} \sqrt{2\pi}} e^{\left(\frac{(q_{20} - q_{20}^{(g.s.)})^2}{2\sigma_{20}^2}\right)} \times \frac{1}{\sigma_{30} \sqrt{2\pi}} e^{\left(\frac{(q_{30} - q_{30}^{(g.s.)})^2}{2\sigma_{30}^2}\right)},
\]

where \( \sigma_{20} = 2800 \text{ fm}^2 \) and \( \sigma_{30} = 6000 \text{ fm}^3 \) are the widths of the Gaussian functions, and \( q_{20}^{(g.s.)} = 3000 \text{ fm}^2 \), \( q_{30}^{(g.s.)} = 0 \text{ fm}^3 \) the coordinates of the ground-state. The advantage of such a choice is that the initial wave-packet is given analytically and does not yield additional numerical errors.

The resulting wave packet is then multiplied by a plane wave characterized by a wave vector \( k = (2.6 \times 10^{-3}, 0) \). This last step gives the initial state an initial momentum toward positive elongations. It also boosts the average energy of the initial state up to roughly 500 keV above the first fission barrier.

In this work, the isoline \( q_N = 3.5 \) mass units defines the scission hyper-surface in the collective space. The width of the absorption zone is set to \( w = 8 \times 10^3 \) in the system of units adopted. In this area, we impose an average absorption rate of \( r = 20 \times 10^{22} \text{ s}^{-1} \). We solve Eq.(3) up to a time \( t_{\text{max}} = 60.10^{-22} \text{ s} \). For the time step values used in table 3 shown in section 5.3, this corresponds to 60000 and 120000 time iterations. At the end of the simulation, the probability for the system to populate post-scission configurations is approximately 30\%. We checked that stopping the time iterations after \( t_{\text{max}} \) would not significantly change the fission fragments yields.

5.2. Construction of the spatial mesh

To minimize the computational cost, the time-evolution is not performed on a regularly meshed hypercube of the collective space. Instead, we use several techniques offered by the finite element method to generate a more efficient partition of the domain \( \Omega \). We list below the various steps followed to produce the mesh:

1. Delaunay triangulation - We start with the initial rectangular, irregularly spaced domain \( \Omega_0 \) mentioned in Sec. 5.1. We generate a first set of regularly spaced vertices in this domain with \( h \equiv h_{30} = 3.1 \times h_{30} \) the resolution of this new mesh \( \Omega_1 \). A Delaunay triangulation provides a partition of \( \Omega_1 \). For every simplex, we choose an interpolation polynomial of degree one. The continuity of each field is ensured by placing the three required nodes at the vertices. The values of the input fields \( \{V(q), B_{ij}^{(k)}\} \) are then evaluated at each node by linear interpolation in \( \Omega_0 \).

2. Absorption areas - The mesh \( \Omega_1 \) is then extended with 60 new spatial steps in the lower \( q_{20} \) values, 40 new spatial steps in the upper \( q_{20} \) values and 20 in both \( q_{30} \) directions. These extensions define the absorption band. By default, the input fields in the absorption band are extrapolated as constants based on their values at the edges of the mesh \( \Omega_1 \). The one exception is the potential field in the lower \( q_{20} \) region of the absorption band. In this case, we evaluate \( V(q) \) as a convex parabola continuously connected at \( q_{20} = 500 \text{ fm}^2 \). This simple extrapolation prohibits the system to explore oblate shapes during the dynamics. This extended mesh is denoted \( \Omega_2 \).

3. h-refinement and coarsening - We take advantage of the flexibility of the finite element method to refine or coarsen the mesh \( \Omega_2 \) depending on its relevance in the time evolution of the system and the flux calculation. This step is needed to improve the numerical precision of the calculation while keeping the computational cost as low as possible. After this series of refinement, the new mesh is \( \Omega_3 \).

- In the regions of the domain \( \Omega_2 \) where the potential takes very large values, the collective wave-function \( g(q, t) \) will remain very small during the entire time evolution. We use the criteria \( V > V_{GS} + 35 \text{ MeV} \), \( q_{20} < 1.0 \text{ and } A_H > 170 \) to automatically detect such regions, where we locally coarsen the mesh by discarding two out of three vertices.

- By contrast, other regions of the domain \( \Omega_2 \) are important to the physics of fission, such as near the ground-state (where the initial collective state is defined) or near the saddle points. In these regions, we apply an additional h-refinement step: each simplex is divided in four new elements by adding a vertex in the middle of each of its edges. At the boundaries of this refinement zone, some simplices may be divided differently to ensure the continuity of the fields in the new mesh.

- The most critical region of the potential energy surface is the scission hyper-surface, where one calculates the total flux. In this area, it is essential to maximize the accuracy of the calculation of the function \( g(q, t) \) and its derivatives. Therefore, we apply two successive h-refinement steps near the scission hyper-surface.

- Up to now the domain \( \Omega_2 \) remains rectangular. However, the areas past and far away the scission hyper-surface are totally irrelevant to the calculation. We thus crop the mesh to retain only the regions of interest. This is done with the following criterion: a simplex in the external region is kept only if its distance to the scission hyper-surface is lower than \( 10^3 \).

4. p-refinement - We apply one p-refinement step to the whole mesh \( \Omega_3 \), that is, after local h-refinement and re-creening and cropping. As a result, the degree of interpolation polynomials in each simplex increases to two. This requires three new nodes per simplex, which are positioned...
in the middle of the simplex edges. The resulting, and final mesh, is denoted \( \Omega \). Starting with \( h = 397 \) (in fm units), it contains a list \( L \) of about 106000 different nodes.

5. Evaluation of fields - Finally, the input fields are again evaluated at each node of \( \Omega \). To do so, results from the original calculation on the mesh \( \Omega_0 \) are linearly interpolated as discussed in the first step.

The figure 8 shows the final mesh \( \Omega \) obtained after all the aforementioned steps. The main advantage of mesh optimization is to reduce significantly the total number of nodes required to achieve a given numerical precision. As an example, let us consider a two-dimensional, regularly-spaced hypercube mesh \( \mathcal{M}_{26} \) with \( h = 476 \) and degree 2 polynomials. We compare it with an optimized version \( \Omega \) built from the same initial spatial step. The total number of nodes has been decreased by more than 25% even though the mesh is locally up to four times more refined in the physically relevant areas.

![Figure 8: Optimized mesh \( \Omega \) obtained with a parameter \( h = 1190 \). The red line represents the scission hyper-surface, defined arbitrarily by the isoline \( q_Y = 3.5 \). This figure shows only the region \( q_{11} > 0 \text{ fm}^3 \) and \( q_{20} > -10^3 \text{ fm}^2 \).](image)

5.3. Results

We performed a series of calculations with different values of the spatial and time resolution parameters \( h \) and \( \delta t \). The parameter values \( h = 793, 595, 476, 397, 340, 298, 264 \) allow us to analyze the convergence in space, while the different time-steps \( \delta t = 10^{-3}, 5.10^{-4} \) (in \( 10^{-22} \text{ s} \)) control the convergence in time.

We first consider the total cumulative flux that crosses the scission hyper-surface during the whole evolution time. This total flux \( F(t_{\text{max}}) \) reads

\[
F(t_{\text{max}}) = \sum_\xi F(\xi, t_{\text{max}})
\]  

The figure 9 shows the rate of convergence of the flux as a function of spatial resolution. The relative difference between the values obtained for \( h = 298 \) and 264 is less than \( 10^{-3} \). We can also notice that the calculation is fully converged in time: the error on the flux is mostly driven by the spatial resolution.

![Figure 9: Total flux crossing the scission hyper-surface in the interval \([0; t_{\text{max}}] \) for two different time-steps \( \delta t = 3 \times 10^{-3}, 5 \times 10^{-4} \) (in \( 10^{-22} \text{ s} \)). The flux is computed for several meshes characterized by the parameter \( h = 793, 595, 476, 397, 340, 298, 264 \). Note that the x-axis is in log scale.](image)

Last but not least, we computed the fission fragment mass yields as a function of both time and space resolutions. As discussed in [9], the mass of the fission fragments along the scission hyper-surface is not necessarily an integer, since both the compound nucleus and the fission fragment are described in the Hartree-Fock-Bogoliubov approximation, where particle number is not conserved. As a result, one must take into account the fluctuations in particle numbers when comparing the yields with experimental data. Following [9], we have convoluted the yields coming out of the flux calculation with a Gaussian with a width of \( \sigma = 3.5 \) mass units. As customary, the resulting values are then normalized to 200. To measure the convergence of the yields, we define the quantity \( e_Y \)

\[
e_Y = \|Y(A) - Y_{\text{ref}}(A)\|_\infty.
\]

The most accurate calculation was obtained with \( h = 264 \) and \( \delta t = 5 \times 10^{-4} \) and is chosen as the reference \( Y_{\text{ref}}(A) \).

| \( \delta t \) | \( h \)   | 793 | 595 | 476 | 397 | 340 | 298 | 264 |
|-------------|---------|-----|-----|-----|-----|-----|-----|-----|
| 10^{-3}     | 1.10    | 0.69| 0.28| 0.26| 0.13| 0.07| 0.03|
| 5.10^{-4}   | 1.08    | 0.67| 0.27| 0.24| 0.11| 0.05| 0.00|

Table 3: Deviation \( e_Y \) as a function of time and space resolutions. The parameters \( \delta t \) and \( h \) are expressed in \( 10^{-22} \text{ s} \) and Fermi units respectively.

We compute the deviation \( e_Y \) for the different values of our numerical parameters. The results are summarized in table 3. We first note that the differences on the deviation \( e_Y \) caused by the time resolution are of the order of 0.02. They are much smaller than the variations induced by the spatial resolution. The evolution of \( e_Y \) as a function of \( h \) shows the convergence...
in space of our calculation. The values obtained for \( h = 298 \) typically correspond to a numerical error of a few percents for masses with \( Y(A) > 1\% \).

6. Program FELIX

The package FELIX is composed of the following directories and files:

- README: contains detailed instructions to build the solver, the tools and their dependencies, and to run the code with the examples provided;
- Makefile: a standard GNU makefile to build the solver and the tools;
- src/: C++ source files of the TDGCM solver and of the tools;
- tools/: additional C++ source files, python and shell scripts to handle the inputs and outputs of the TDGCM solver;
- benchmarks/: a few preset inputs and their corresponding outputs;
- doc/: documentation of the package in DoxyGen format.

The full Felix package depends on several standard Open Source libraries:

- The TDGCM solver itself requires BLAS, LAPACK, and a Fortran compiler with OpenMP support;
- The documentation requires DoxyGen-1.8.6 or higher;
- In order to build the full set of tools included in this release, the user must also install GSL, PETSc, SLEPc and Boost. The versions GSL-0.16, PETSc-3.5.2, SLEPc-3.5.3 and Boost-1.54 have been used during development.

6.1. Compilation

The program is shipped with a Makefile containing a preset configuration assuming compilation with the GNU gcc compiler on a standard LINUX distribution. The user should expect to have to change this Makefile to match his/her own system configuration. The Makefile contains some instructions to help the user with this configuration step. The different components of the package can be compiled separately by typing the following commands:

- make Doc: generate the DoxyGen documentation in the directory, which is located in doc/DoxygenDoc.
- make Solver: build the TDGCM+GOA solver executable. Its name is tdgcmFE and it is located by default in src.
- make Tests: build the executable for the full suite of tests included in the package. The name of the executable is tdgcmFEtest and is located in tests/src/.
- make Tools: compile all tools in the directory tools/.

6.2. Running the solver

There are two different ways to run the FELIX solver with specific input data.

- If an option file input.opt is available, the user may simply type
  
  ./src/tdgcmFE input.opt

- Otherwise, the list of options can directly be passed via the command line as:
  
  ./src/tdgcmFE --option0 [value0] --option1 [value1] ...

The available options are discussed below in section 7.2.

By default, the solver uses every available core on the machine. As usual, the number of OpenMP threads can be controlled by setting the environment variable OMP_NUM_THREADS.

6.3. System of units

By default, the value of the reduced Planck constant \( h \) is set to

\[
 h = 6.58211928 \times 10^{-22} \text{MeV.s). (56)}
\]

This imposes a consistency relation between the energy and time units that can be used in the code. The most natural choice is to set the value of energies in MeV so that time is given in units of \( 10^{-22} \) s. This is recommended and is the default setting for FELIX.

If needed, the value of the reduced Planck constant can be changed in the file ./src/defines.h, which allows the user to define his/her own set of physical units. Note that the code must be entirely re-compiled for such changes to take effect. Also, special attention must be paid to setting the units for the inertia tensor and coordinates \( q \) in a consistent manner.

7. Inputs and outputs

7.1. Input files

FELIX reads most of its input from several ASCII files. The names of these files begin with the same user-defined prefix, and have a specific extension, as for instance, example.coor, example.val, example.geo, etc. The following mandatory files are needed by the solver:

- example.coor: This file contains an unsorted list \( L \) of points in the domain \( \Omega \) defined by their coordinates \( q \) (in their appropriate unit). In the other inputs files, a point is labeled by its index in the list \( L \). Each line of this file contains a series of \( N \) double numbers separated by a space;
- example.geo: This file defines the list of all simplices of the mesh. Every line contains a series of \( \text{int} \) integers separated by spaces: the first integer is the degree of the interpolating polynomial used in the simplex; it is followed by the list of the \( N + 1 \) vertices of each simplex, followed by all the nodes used to define the interpolating polynomials. In this version of FELIX the only limitation on the
geometry comes from the boundary condition Eq. 11. Internally, this condition is imposed by setting the values of the coefficients \( g(q_{\text{f}}, t) \) to zero for every node lying on the boundary \( \partial \Omega \). The user must therefore design the boundary elements so that the aforementioned property implies Eq. 11.

- `example.val`: This file contains the values of several fields at each point of the list \( L \). The first line starts with the special character `#`, and contains a list of string keys that define the fields. Mandatory fields are the potential \( V(q) \) and the lower part of the inertia tensor. The key for the potential is simply \( v \); for the inertia tensor: in 2D, the keys are \( B00, B10, B11; \) in 3D, they are \( B00, B10, B20, B21, B22; \) etc. Optional fields recognized by the code are \( qN, tN \) – the expectation value of the Gaussian neck operator, and \( AH \) – the mass of the heaviest fragment.

- `example.init`: It contains the values of the initial wave function \( g(q, t = 0) \) at each point of the list \( L \). Each line is made of two double corresponding to the real and imaginary parts of the wave function.

The user may also specify additional features and options. They are handled through the option list described in section 7.2 below. Some of these options require one or several additional input files:

- `example.front`: This file contains a list of oriented hyper-surfaces for which the solver will calculate the flux. The surfaces are simplex edges and are defined by \( N \) vertices. The additional vertex "vOpposite" shown in figure 1 must be specified to set up the orientation.

- `example.matM` and `example.matH`: These two files store the sparse matrices \( M \) and \( H \) defined in Eq. (26).

- `example.opt`: A file containing the list of options described in section 7.2.

7.2. Options

The user can pass a number of options to the solver. Most of these options control the numerical parameters of the calculation and the frequency at which output data is written on disk.

- `--help (flag)`
  If present, the code only prints a help message and stops the execution. Default: Absent.

- `--version (flag)`
  If present, the code only prints the version number and stops the execution. Default: Absent.

- `--file (string)`
  Prefix name for the input files. All input files must be named as `prefix.ext` where `ext` is one of the extensions described in section 7.1. Default: `input`.

- `--outputDir (string)`
  Name of the output directory. If no existing directory is found, a new directory is created. Default: `./results`.

- `--dump (integer)`
  Number of time iterations between two prints of the solution and the flux. Default: 100.

- `--refresh (integer)`
  Number of time iterations between two displayed lines in the standard output. Default: 100.

- `--max (integer)`
  Maximum number of time iterations for the calculation. Default: -1 (no maximum).

- `--step (double)`
  Time step of the calculation. By default, the unit of the time step is \( 10^{-22} \) s. Default: \( 10^{-4} \).

- `--inversionTol (double)`
  Numerical tolerance for matrix inversion; see Eq. (31). Default: \( 10^{-15} \).

- `--limit (integer)`
  Maximum number of iterations in matrix inversions. Default: \( 10^5 \).

- `--absRate (double)`
  Average absorption rate per time unit in the absorption zone. For instance, `--absRate 20` specifies an absorption rate of \( 20 \times 10^{22} \) s\(^{-1}\), since the basis time unit is \( 10^{-22} \) s. Default: 0.

- `--absWidth (double)`
  Width of the absorption zone. The unit depends on the units of the collective coordinates \( q \). For any node of the mesh, the euclidean distance \( d \) to the boundary \( \partial \Omega \) is computed. A node is included in the absorption zone if and only if \( d < \text{absWidth} \). A negative value will lead to no absorption. Example: Consider a rectangular 1D space with the axial quadrupole moment given in b. Assume the domain is \( q_{20} \in [0, 600] \) b and \( \text{absWidth}=10 \). Then, all points with \( q_{20} \in [0, 10] \cup [590, 600] \) b will be included in the absorption band. Default: -1.

- `--calcEnergy (flag)`
  If this flag is present, the code prints the average energy of the solution every dump time iteration. Default: Absent.

- `--dumpMat (flag)`
  If present, the sparse real matrices \( M \) and \( H \) are stored in the files `input.matM` and `input.matH` respectively. Default: Absent.

- `--readMat (flag)`
  If present, the code reads the \( M \) and \( H \) matrices from the files `input.matM` and `input.matH`. Note that these matrices depend only on the particular mesh of the collective space, but
not on time \( t \): for a given mesh, they can be pre-calculated, stored on disk using the \(--dumpMat\) flag, and re-used in a different run. Default: Absent.

\(--ISMethod\) (string='file', 'impulsed' or 'wavePacket')
Method of determination of the initial state \( g(q, t = 0) \).

- If set to 'file', the solver reads the initial state from the file \texttt{input.init}. 
- If set to 'impulsed', the code reads the initial state contained in the file \texttt{input.init} and multiplies it by a plane wave. We note \( \mathbf{k} = \hat{\mathbf{k}} \) the wave vector characterizing the plane wave, with \( \hat{\mathbf{k}} \) the unit vector indicating its direction and \( \mathbf{k} \) the modulus. The user must specify the coordinates of \( \mathbf{k} \) in a file \texttt{input.k}. The modulus \( k \) is computed by the solver so that the average energy of the initial state matches the value provided with the option \(--ISEnergy\). The user may provide an initial guess of the parameter \( k \) via the option \(--ISLambdaGuess\).
- If set to 'wavePacket', the solver builds a linear combination of states provided by the user \( g(q, t = 0) = \sum_k \alpha_k g_k(q) \). The user provides the states \( g_k(q) \) as a set of files \texttt{state}_0, \texttt{state}_1, etc. A single directory, the name of which is set with the option \texttt{ISSstatesDir}, must contain all the files. The format for the files \texttt{state}_k is the same as for \texttt{input.init}. In the expansion of the initial state, all weights have a Gaussian dependency on the expectation value \( E_k \) of the energy of each state: \( \alpha_k \propto \exp(-E_k - \langle E \rangle)^2/2\sigma^2) \). The user can tune the parameter \( \sigma \) via the option \(--ISSigma\). The code determines the parameter \( \langle E \rangle \) so that the energy of the wave packet matches the value provided in the option \(--ISEnergy\). Default: 'file'.

\(--ISEnergy\) (double)
Requested energy of the initial state for the methods 'impulsed' and 'wavePacket'. Default: 0.

\(--ISSigma\) (double)
Gaussian width of the initial state for the method 'wavePacket'. Default: 1.

\(--ISSstatesDir\) (string)
Directory containing the files \texttt{state}_k required for the construction of the initial state with the method 'wavePacket'. Default: '.\eigenStates'.

\(--ISLambdaGuess\) (double)
Initial guess for the modulus \( k \) of the plane wave multiplying the initial state with the method 'impulsed'. If this value is negative, the code will initialize \( k \) with a default value. Default: -1.

\(--ISNorm\) (flag)
If present, the initial state will be normalized to 1 before starting time iterations. Default: Absent.

\(--frontier\) (string='qN' or 'file')
Method of determination of the frontier. If set to 'file', the frontier will be read from the corresponding input file. If no such file can be found, the frontier is empty and the instantaneous flux is zero. If set to 'qN', the frontier will be computed on the fly as an isoline of the field \( qN \). This field must then be present as an additional column in the .val file, with the key \( qN \). Default:'file'.

\(--frontIso\) (double)
Value of the field \( qN \) used to define the scission hyper-surface. Default: 1.

\(--fluxInst\) (flag)
If present, the instantaneous flux through the frontier is recorded every dump time iteration. Default: Absent.

\(--lumpedMass\) (flag)
If present, the lumped mass approximation is applied when calculating the \( M \) matrix. Default: Absent.

\(--pCoeff\) (double)
Arbitrary multiplicative factor applied to the potential field \( V(q) \). Default: 1.

\(--bCoeff\) (double)
Arbitrary multiplicative factor applied to the inertia tensor field \( B_m(q) \). Default: 1.

7.3. Output files
All outputs are recorded in a directory that can be specified via the option \texttt{outputDir} described in the previous section. Upon successful execution of the solver, this directory should contain

- The file \texttt{example.opt}: It contains the list of all options used for the run. It could be re-used as an input file for any other calculation;
- The directory \texttt{gFunction/:} Each file \texttt{gFunction.xxx.log} in this directory contains the solution \( g \) after \( xxx \) time iterations. These files are formatted in the same way as the input file \texttt{example.init}. For example, the file \texttt{gFunction.000000000.log} is a copy of \texttt{example.init}, unless a renormalization has been requested by the user through the use of option \texttt{norm} described above. Thanks to this identical format, these files can be used in subsequent runs as initial wave-functions by simply copying them in place of whatever \texttt{.init} file was used. The program FELIX thus has basic checkpointing capabilities;
- The file \texttt{normDeviation}: It contains the deviation \( \|g(t)\|_2 - \|g(t = 0)\|_2 \) as a function of the number of time iterations;
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9. References

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