Documentation for laser-propagation

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1 Field propagation

1.1 Propagation equation

The propagation of the electric field is represented by the unidirectional pulse propagation equation written in the spectral domain as

$$\frac{\partial \mathcal{E}(z, k_{\perp}, \omega)}{\partial z} = ik_z(k_{\perp}, \omega)\mathcal{E}(z, k_{\perp}, \omega) + \frac{i\omega^2}{2\epsilon_0 c^2 k_{z,n}(k_{\perp}, \omega)} \left[\mathcal{P}(z, k_{\perp}, \omega) + i\frac{\mathcal{J}(z, k_{\perp}, \omega)}{\omega} \right]$$
(1)

where $k_z(k_{\perp}, \omega)$ describes how the spectral field amplitudes propagate linearly in the medium (both diffraction and linear absorption). The nonlinear properties of the medium are described by functions representing the nonlinear polarization P and/or nonlinear current J. The functional forms of k_z , P, and J are determine by the parameter file supplied by user at runtime. All of the available options are detailed in Section 2.

It's not ideal to solve Equation (1) directly since it still contains a fast oscillating term resulting from linear propagation. The is remedied by defining the spectral amplitude $A(k_{\perp}, \omega) = E(k_{\perp}, \omega) \exp[-ik_z z]$, whose evolution is the govern by the nonlinear sources only

$$\partial_z \mathcal{A}(z, k_\perp, \omega) = \frac{i\omega^2}{2\epsilon_0 c^2 k_z(k_\perp, \omega)} e^{-ik_z z} \left[\mathcal{P}(z, k_\perp, \omega) + i \frac{\mathcal{J}(z, k_\perp, \omega)}{\omega} \right]$$
 (2)

The variables $A(k_{\perp}, \omega)$ are the native variables of the simulation. Their evolution equation is a large ODE system that is solved using GSL's ode solver library. This library provides many methods of solving ODE's and includes adaptive step control. The ODE solver only requires the RHS of Eq. 2 and some error tolerance values to control the adaptive stepping algorithm.

1.2 Algorithm

At each step in solving Eq. (2), the solver invokes a call to the RHS with a proposed step size forward in z. Since nonlinear sources P and J are typically defined in real space, the RHS function has the job of spectrally transformed $\mathcal{A}(k_{\perp},\omega)$ to E(r,t), calling each nonlinearity, and transforming back to spectral amplitudes. The RHS function returns the change in spectral amplitudes $\partial_z \mathcal{A}(z,k_{\perp},\omega)$ to the solver, which determines whether the proposed step was small enough to evolve the amplitudes \mathcal{A} within a given error tolerance.

Explicitly, during each step of the ODE solver, the RHS is calculated the following way:

1. Linear propagate \mathcal{A} forward by Δz (ω/v_q term keeps the pulse centered in the computational box)

$$\mathcal{E}(k_{\perp}, \omega) = A(z, k_{\perp}, \omega) \exp\left[i\left(k_z\left(k_{\perp}, \omega\right) - \omega/v_a\right) \Delta z\right] \tag{3}$$

2. Transform to real space in radial coordinates using the Fourier-Hankel transform

$$E(r,t) = \text{Hankel}\{\mathcal{E}(k_{\perp})\} \text{FT}\{\mathcal{E}(\omega)\}$$
(4)

3. Calculate nonlinearities

$$P(r,t) = \sum_{i} P_{i}[E(r,t)]$$

4. Transform to spectral space

$$P(k_{\perp}, \omega) = \text{Hankel}\{P(r)\}\text{FT}^{-1}\{P(t)\}$$

5. Undo the propagation in Step 1

$$\mathcal{P}(k_{\perp}, \omega) = P(k_{\perp}, \omega) \exp\left[-i(k_z(k_{\perp}, \omega) - \omega/v_q)\Delta z\right]$$
(5)

6. Finally, multiply the nonlinearity by the coupling factor and return to ODE solver

$$\partial_z \mathcal{A}(k_\perp, \omega) = \frac{i\omega^2}{2\epsilon_0 c^2 k_z} \mathcal{P}(k_\perp, \omega) \tag{6}$$

1.3 Computational Grid

The computational grids in the simulation are three 2D arrays representing the fields $\mathcal{A}(k_{\perp},\omega)$ and E(r,t), and also the electron density $\rho(r,t)$ in cylindrical coordinates. There are also some auxillary arrays that are used for computing the spectral transforms. The spectral transformation of this grid involves a Fourier transform between time t and frequency ω and a Hankel transform between r and k_{\perp} .

To perform the Fourier transform the library FFTW is used. A norm-preserving Hankel transform was introduced by H. Fisk Johnson in 1986. The radial grid points are not equally spaced, but are proportional to the zeros of the Bessel $J_0(\alpha_i) = 0$, and are defined as $r_i = R_{max}\alpha_i/\alpha_N$ where N is the number of radial

grid points. The values of $k_{\perp,i} = \alpha_i/R_{max}$. With this coordinate transformation, the Hankel transform is simply a matrix multiplication with elements

$$H_{mn} = \frac{2J_0(\alpha_n \alpha_m / \alpha_N)}{J_1(\alpha_n)J_1(\alpha_m)\alpha_N}$$

The Hankel transform is its own inverse so multiplication twice is the identity: HHf = f. This multiplication is speed up by using the library Eigen, which generates SIMD instructions resulting in a speed up of 30-40% in freespace calculations.

The size of the spectral amplitudes \mathcal{A} (number of points in k_{\perp} and ω) is always smaller than the size of E, since only positive frequency components of \mathcal{A} are propagated. This saves quite a bit of memory since the number of active frequencies ω are less than half the size of the number of time points in the grid. Using a capillary waveguide with only a few number of modes (the number of points in k_{\perp} greatly reduces the size of \mathcal{A} as well. For example, a capillary waveguide simulation with a real-space grid size of $Nr \times Nt = 100 \times 16384 = 1.6M$ points is represented by the spectral amplitude grid-size of $N\omega \times Nk_{\perp} = 2199 \times 10 = 22k$ points when using 10 capillary modes and only propagating the active frequencies corresponding to the wavelengths between 250 nm - 3000 nm.

It's good to keep in mind the algorithmic scaling of the two transformations: FFT and Hankel. The FFT is performed for each radial grid point and scales as $\mathcal{O}(NrNt\log_2 Nt)$. FFT runs most efficiently with Nt being of a power of two: 2^k . The Hankel transform has a different scaling for freespace and capillary calculations. For freespace calculations a full matrix multiplication is done for each active frequency ω and thus scales as $\mathcal{O}(N\omega Nr^2)$. Due to the reduced Nk_{\perp} in capillary simulations this complexity can be reduced to $\mathcal{O}(N\omega Nk_{\perp}Nr)$. Since the number of capillary modes Nk_{\perp} is usually around 5-10, the Hankel transform in the capillary simulations scale linearly with the number of radial grid points.

2 Parameter file

The type of simulation, medium model, choice of nonlinearities, and output are controlled by a parameter file that is passed to the executable as an argument on the command line. All values in the parameter file are assumed to be in SI units. There are some values that are required in order to run any type of simulation (e.g. setting up the computational grid, initial conditions, etc.), but many values (e.g. selecting which nonlinearities to include) are optional and only need to be included in the parameter file if you wish to have them in the simulation.

2.1 Syntax

The syntax of the parameter file is similar to conf or ini styles, where key-value pairs are grouped into sections. Here's an example:

```
[section1]
key = value
x = y

# top level comment
[section2]
s = hello
a = 1.1
x = -2e-12  # inline comment
```

Whitespace surrounding the keys, values, sections, and = symbol are ignored. Comments can be placed basically anywhere since all text after a hash # symbol is ignored. The keys and values can be read and runtime to the program by the Parameters class defined in parameters.h & parameters.cc.

To read the parameter file input:

```
[time] N = 1024
```

```
time_min = -10e-15
time_max = 50e-15
```

instantiate the Parameters class with the filename and extract the values

```
Parameters::Parameters params("input");
int Nt = params.get<int>("time/N");
double time_min = p.get<double>("time/time_min");
double time_max = p.get<double>("time/time_max");
```

Internally, all keys and values are stored as strings and keys have had their surrounding section prepended to them using a /. The Parameters class has a template function get that uses stringstreams to interpret the type of value. This means that the syntax of the values is limited to readable C++ types: e.g. string, int, double, etc.

2.2 Setting up the computational grid

2.2.1 [time] (required)

This section defines the properties of the temporal and spectral domains of the field. For the temporal domain, the keys to define are the number of temporal points N, and the minimum time_min and maximum time_max values of the temporal range. For efficiency, N should be a power of 2. Additionally, only certain positive frequency components of the spectral field are propagated. To set the range of active frequencies, define wavelength_min and wavelength_max.

An example of this section is:

```
[time]
N = 2048
time_min = -100e-15
time_max = 200e-15
wavelength_min = 150e-9
wavelength_max = 4e-6
```

This defines a temporal box of 2048 points extending from -100fs to 200fs. The active frequencies in the simulation are only the ones that fall between 150nm and 4 microns.

2.2.2 [filtering] (optional)

This section defines temporal and spectral filtering. Filtering is crude since it amounts to multiplication of the temporal and spectral fields by \sin^2 mask before the forward and inverse Fourier transformations. The values defined here along with the ones defined in sections [time] determine how fast \sin^2 ramps go from 0 to 1. Setting a filter min/max value to a value outside the limits in [time] disables filtering on that particular side.

Reasonable filter values for the time domain in previous section could be:

```
[filtering]
time_filter_min = -90e-15
time_filter_max = 150e-15
wavelength_filter_min = 250e-9
wavelength_filter_max = 3e-6
```

Note: there is no filtering in r or k_{\perp} which leads to reflections if the field reaches the radial boundary. If interested in THz generation, the filter value on the long wavelength side should be set to a large value:

```
[filtering]
time_filter_min = -90e-15
time_filter_max = 150e-15
wavelength_filter_min = 250e-9
wavelength_filter_max = 1
```

2.2.3 [space] (required)

This section defines the spatial properties of the radial domain: N sets the number of radial points, and radius_max sets the radial extent of the domain. Since the radial spectral transform is the Hankel transform, there are no special values for N that are more efficient, unlike there for the time domain (due to the FFT).

An example of this section is:

```
[space]
N = 173
radius_max = 10e-3
```

This defines a radial domain of 173 points with a maximum extent of 10 mm.

2.3 Propagtion and ODE step control

2.3.1 [propagation] (required)

This section sets the physical starting and ending distances (starting_distance and ending_distance) of the simulation, and when data is written to the disk. The key num_reports_cheap sets the number of cheap diagnostics (fast to calculate or small in written data size) to perform between the starting and ending distances, and num_reports_expensive sets the number of expensive diagnostics. Whether a diagnostic is cheap or expensive is defined by the how the Observer was added in main.cc:initialize_observers via the function main.cc:conditionally_add.

An example of this section is:

```
[propagation]
starting_distance = 0
ending_distance = 4
num_reports_cheap = 40
num_reports_expensive = 4
```

The laser propagates from 0 to 4 meters and during propagation 40 cheap and 4 expensive diagnostics are performed.

2.3.2 [ode] (required)

This section defines the parameters for the ODE solver. Adjustment of these values has a great impact on the runtime of the simulation, since they force the solver to take small enough steps to satisfy these criteria. The values below are passed directly to GSL's ODE step control function. They form a two-parameter heuristic value $D_i = \epsilon_{abs} + \epsilon_{rel} |A_i|$ that is compared to the error calculated by the RK45 method $|Aerr_i|$.

For fast, but perhaps not the most accurate simulations, try:

```
[ode]
absolute_error = 1e10 # or 1e8
relative_error = 1e-4
first_step = 1e-3
```

This sets the two error parameters and the first attempted step of the simulation to be 1 mm. For production runs where the data should be reasonable converged, try:

```
[ode]
absolute_error = 1e6
relative_error = 1e-4
first_step = 1e-3
```

2.4 Linear properties

2.4.1 [medium] (required)

This section defines the linear properties of the medium in the simulation. The key type controls the functional form of the linear propagator (basically the formula for $k_z(k_{\perp},\omega)$), and can be set to either freespace, capillary, or diffractionless.

Selecting freespace sets

$$k_z(k_\perp,\omega) = \sqrt{\left(\frac{n(\omega)\omega}{c}\right)^2 - k_\perp^2}$$

Selecting capillary sets

$$k_z(k_\perp,\omega) = \sqrt{\left(\frac{n(\omega)\omega}{c}\right)^2 - k_\perp^2} + i\alpha(k_\perp)$$

where the absorption α due to the gas cladding interface is

$$\alpha = \frac{1}{2R} \left(\frac{k_{\perp} c}{\omega} \right)^2 \frac{n_{clad}^2 + 1}{\sqrt{n_{clad}^2 - 1}}$$

where R is the capillary radius and n_{clad} is the index of the cladding. If capillary is chosen, then the input file must contain the section [capillary] as described in 2.4.2.

Selecting diffractionless sets

$$k_z(k_\perp, \omega) = \frac{n(\omega)\omega}{c}$$

which is useful for verifying that temporal dispersion is computed properly.

The key index defines the index of refraction function $n(\omega)$. Currently implemented functions are: vacuum, air, argon, and ethanol. The index can also be defined using tabulated data from a file that will be linearly interpolated onto the points ω of the spectral domain. To do this set index equal to a filename ending with the .dat extension and contains three space-separated columns: $\omega \Re\{n(\omega)\}$ $\Im\{n(\omega)\}$.

Lastly, the key pressure is used to increase or decrease the index of refraction using the Lorentz-Lorenz relation.

An example of freespace propagation in air at a pressure of 1 atm is:

```
[medium]
type = freespace
index = air
pressure = 1
```

An example of capillary propagation in argon at 3.3 atm is:

```
[medium]
type = capillary
index = argon
pressure = 3.3
```

2.4.2 [capillary] (optional)

This section defines a capillary waveguide which sets the functional form of $k_z(k_{\perp},\omega)$ (adding absorption from gas-cladding interface) and also the number of propagating capillary modes. The index of the cladding is cladding.

```
[capillary]
cladding = 1.45
modes = 5
```

2.5 Initial conditions

2.5.1 [laser] (required)

This section defines the initial laser pulse. The key type can be gaussian, restart, or file. An example of a Gaussian beam is:

```
[laser]
type = gaussian
wavelength = 800e-9
length = 20e-15
waist = 4e-3
focus = 2
energy = 1e-3
chirp = 0
phase_deg = 0
delay = 0
```

Here the laser has a envelope shape of Gaussian in space and time. It's central wavelength is 800 nm, FWHM temporal length is 20 fs, $1/e^2$ radius is 4 mm, is focused at 2 m, has energy of 1 mJ, and has zero chirp, phase and temporal delay. Note: the starting_distance in [propagation] can be used to linear propagate forward in z assuming Gaussian beam propagation in space and a Gaussian-like propagation in time through a dispersive media (gvd and chirp are taken into account). This is useful for only performing a nonlinear simulation near the focus, while propagating linearly through a medium beforehand.

An example of restarting from an old simulation:

```
[laser]
type = restart
filename = A004.dat
wavelength = 800e-9
```

The filename must point to a spectral field file that was generated from a previous simulation, and you must specify the central wavelength (used for calculating values, such as n0, gvd, P_{cr}). Note: you might want to set the starting_distance in [propagation] to match the physical distance of the spectral field file.

An example of starting from a binary space-time field file:

```
[laser]
type = file
filename = init_field.dat
wavelength = 800e-9
```

Note: the field E(r,t) in init_field.dat must be stored in a binary file of doubles (8 bytes) where the real and imaginary values are stored as [real0 imag0 real1 imag1 ...]. The time axis t is the fast axis (convention of C, opposite of Fortran). The sampling of r and t must match the values are set in the sections [time] and [space].

2.6 Output data

2.6.1 [results] (optional)

This section defines which data is written to which files during propagation. All of the entries in this section are optional. Commenting or removing any line below results in no data being written for that particular item.

```
[results]
# field and density filename patterns and their associated distances
temporal_field = E
spectral_field = A
electron_density = Rho
```

```
distance = distance.dat

# coordinates
time = time.dat
radius = radius.dat
omega = omega.dat
kperp = kperp.dat
wavelength = wavelength.dat

# values along propagation
energy = energy.dat
max_intensity = max_intensity.dat
max_density = max_density.dat
```

The keys are fixed by the program, but the values can be changed by the user. The only special values are the ones for the field and density files (temporal_field = E, spectral_field = A, and electron_density = Rho) where their filename values contain no extension (e.g. dat or txt). This is because their filenames will be enumerated (e.g. E000.dat, E001.dat, E002.dat, etc.) and the distances at which they are written out is contained in distance.

2.7 Nonlinear properties

2.7.1 [kerr] (optional)

Defining this section adds Kerr nonlinearity to the simulation using

$$P_{\rm NL}(r,t) = \epsilon_0 \chi^{(3)} E^3(r,t)$$

where $\chi^{(3)} = \frac{4}{3}\epsilon_0 c n_2 n_0^2$. The only value to set is the nonlinear coefficient n_2 .

```
[kerr]
n2 = 8e-24
```

Note: The value of $\chi^{(3)}$ is multiplied by the value of pressure that is set in section [medium].

2.7.2 [ramankerr] (optional)

Defining this section add the Raman-Kerr nonlinearity to the simulation as:

$$P_{\rm NL}(r,t) = \epsilon_0 \chi^{(3)} \left[(1 - f_R) E^2(r,t) + f_R \int_{-\infty}^t R(t - t') E^2(t') \right] E(r,t)$$

where f_R is the Raman fraction, $R(t) = R_0 \exp(-\Gamma t/r) \sin(\Lambda t)$, $R_0 = (\Gamma^2/4 + \Lambda^2)/\Lambda$, $\chi^{(3)} = \frac{4}{3}\epsilon_0 c n_2 n_0^2$. The required keys are n2 for the nonlinear coefficient, fraction for the proportion of n2 associated with the delayed Raman response, and gamma & lambda for the frequency parameters of R(t).

This equation is solved using the exponential time differencing method. The discrete formula for the time integration for array index j from 0 to Nt is

$$P_{\rm NL}[j] = \epsilon_0 \chi^{(3)} \left[(1 - f_R) E[j]^2 + f_R Q[j] \right] E[j]$$

with an initial condition for j = 0

$$Q[0] = R_0 \frac{\Delta t}{2} E[0]^2$$

and for $j \geq 1$

$$Q[j] = \Im \left\{ e^{(-\Gamma/2 + i\Lambda)\Delta t} Q[j-1] + R_0 \frac{\Delta t}{2} \left[E[j]^2 + e^{(-\Gamma/2 + i\Lambda)\Delta t} E[j-1]^2 \right] \right\}$$

An example of this section is:

```
[ramankerr]
n2 = 22.3e-24
fraction = 0.65
gamma = 10e12
lambda 16e12
```

Note: the value of $\chi^{(3)}$ is multiplied by the value of pressure that is set in section [medium].

2.7.3 [ionization] (optional)

Defining this section adds a rate of ionization which populates the electron density $\rho(r,t)$ at each step of the simulation according to

$$\frac{\partial \rho}{\partial t} = W(E(t))(f\rho_{nt} - \rho)$$

where W is the rate of ionization and ρ_{nt} is the density_of_neutrals. This equation is solved using the exponential time differencing method

$$\rho(t + \Delta t) = \exp\left[-\int_{t}^{t + \Delta t} W(E(t'))dt'\right] \left\{\rho(t) + \frac{\Delta t}{2}\rho_{nt}W(E(t))\right\} + \frac{\Delta t}{2}\rho_{nt}W(E(t + \Delta t))$$

If the keys generate and formula are present the rate of ionization is calculated for the given central laser wavelength (defined in [medium]) and ionization_potential. The value of generate is the filename to which the rates are written (two-column data corresponding to the intensity $[W/m^2]$ and rate [1/s]). The options for formula are adk, mpi, tunnel, yudin, and ilkov.

The rates can also be read from a file by defining the key filename. This file should contain two-column data corresponding to the intensity $[W/m^2]$ and rate [1/s].

Defining [ionization] also adds nonlinear absorption

$$J_{\rm NA}(t) = \frac{W(E(t))}{I(t)} U_i \left(f \rho_{nt} - \rho(t) \right) \epsilon_0 c E(t)$$

where U_i is the ionization-potential and f is the fraction of neutrals that can be ionized (ionizing_fraction). Defining [ionization] also adds effects due to plasma according to

$$\frac{\partial J_{\rm PL}}{\partial t} + \frac{J_{\rm PL}(t)}{\tau_c} = \frac{e^2}{2m_e} \rho(t) E(t)$$

where τ_c is the collision_time. This equation is solved using the exponential time differencing method:

$$J_{\rm PL}(t+\Delta t) = e^{-\Delta t/\tau_c} \left\{ J_{\rm PL}(t) + \frac{\Delta t e^2}{2m_e} \rho(t) E(t) \right\} + \frac{\Delta t e^2}{2m_e} \rho(t+\Delta t) E(t+\Delta t)$$

```
[ionization]
filename = ../../data/ionization-argon.txt
ionizing_fraction = 1
density_of_neutrals = 2.5e25
collision_time = 190e-15
ionization_potential = 2.5250303e-18
```

2.7.4 [argon] (optional)

This is an experimental feature! Adding the section [argon] to the parameter file initializes a realistic SAE model for argon. This model is based on the work from "Calculations of strong field multiphoton processes...", Schafer et al., AIP Conf. Proc. **525**, 45 (2000).

There is an atom placed at each radial point, so the number of quantum systems in the simulation is equal to the value of N in [space]. The time-dependent nonlinear polarization (from the dipole moment) and ionization probability are computed for each atom as it interacts with the laser field.

Since the time to compute the response of a single atom is anywhere from 20-120 s (depending on the values that are chosen for the quantum system), it's highly recommended that the parallel MPI version of the executable main-mpi.out is used. Since there is no interaction between the individual argon atoms, parallelization is straightforward. The laser simulation run on processor 0, as well as a single argon atom for radial grid point 0. Every other processor holds a single argon atom for the radial grid points 1 to N. When the electron density or nonlinear polarization is calculated, the laser field for each radial point is distributed to every processor using MPI_Scatter. Everyone calculates the electron density and nonlinear dipole moment, then sends their results back to processor 0 using MPI_Gather. Since there is currently the limitation of one quantum system per radial grid point, it is necessary to launch the executable with the same number of threads as radial grid points defined in [space]. If this is not the case, the program signals an error.

A good set of parameters to start with for Argon gas at 15 C, 1 atm are the following:

```
[argon]
Nr = 300
Nl = 21
Nmask = 100
step_size = 1
potential = short-range-potential.dat
ionization_box_size = 40
density_of_neutrals = 2.55e25
collision_time = 190e-15
ionization_potential = 2.5250303e-18
```

where Nr is the number of grid points in radius for the wavefunction (the spacing in r, Δr is fixed at 0.25 in atomic units), N1 is the number of angular momentum states, Nmask is the number of radial grid points for the absorber, step_size is the time scale (in atomic units) on which the wave function evolves, potential is the filename that contains the short range potentials of argon for l=0,1,2 states. The parameter ionization_box_size specifies the number of radial grid points above which the wavefunction is considered ionized (i.e. the part of the wavefunction that is outside of this sphere is considered part of the free electron probability). Finally, for scaling the single atom response up to a macroscopic response, it's necessary to define the density of neutrals.

Nonlinear effects from plasma are also added and contribute a current:

$$\frac{\partial J_{\rm PL}}{\partial t} + \frac{J_{\rm PL}(t)}{\tau_c} = \frac{e^2}{2m_e} \rho(t) E(t)$$

where τ_c is the collision-time. Nonlinear absorption due to ionization is also added using: Nonlinear effects from plasma are also added:

$$J_{\rm NA}(t) = \frac{W(E(t))}{I(t)} U_i \left(f \rho_{nt} - \rho(t) \right) \epsilon_0 c E(t)$$

where U_i is the ionization_potential.