Documentation for the RTA code in 3D

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Abstract

We present a RTA+TDLDA code on a cartesian 3D grid

Keywords: electron-electron collisions, electronic disspation, time-dependent density functional theory, metal cluster, plasmon, electron emission

1. The subroutines in detail

SUBROUTINE calc_Eref(occup,ispin,Ei,Eref)

```
occup(1:nstate) in occupation number for s.p. states. ispin(1:nstate) in spin assignement for s.p. states. Ei(1:nstate) in spin assignement for s.p. states. Eref(1:2) out sum of s.p. energies per spin.
```

Computes the weighted sum of s.p. energies as reference energy for DCMF. The sum is accumulated for each spin separately.

SUBROUTINE fermi1(ekmod,eref,occup,ispinact,T0i,T1i,T2,mu)

```
ekmod(1:kstate)
                      in
                             given s.p. energies, spin up block first, then spin down
eref
                             reference energy = wanted sum of s.p. energies
                      in
ispinact
                      in
                             spin for which routine is run
TOi, T1i
                      in
                             lower and upper temperature for search
occup(1:kstate)
                    in/out
                             occupation numbers, spin block-wise
                             final temperature for which Fermi distribution matches eref
T2
                     out
                     out
                             final chemical potential
mu
```

Determines thermal Fermi occupation such that given sum of s.p. energies eref and particle number is matched. Is done for each spin separately. Solution scheme is bracketing. Refers to SUBROUTINE OccT1 while iterating temperatur T2.

PGR2all: Nr. of spin-up/spin-down states comes through m_params. We should protocol all

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such entries. First step is to augment each USE by ONLY such that the explicitely communicated variables becomes visible. Important variables may then be listed explicitly.

PGR2all: Routine requires that arrays are sorted in continuous blocks of spin. Do we have an initial check for that? And we need to address that in the general part which explains the layout of arrays.

SUBROUTINE OccT1(occrefloc,enerloc,Etotloc,muloc,occtotloc,n,T,occuploc)

```
s.p. energies for actual spin
enerloc(1:n)
                        number of s.p. states treated here
                  in
n
Т
                        temperature
                  in
occrefloc
                  in
                        wanted total number of particles
                       thermal occupation numbers for given T and s.p. energies
occuploc(1:n)
                  out
                       chemical potential (Fermi energy)
muloc
                  out
                       final total number of particles
occtotloc
                  out
Etotloc
                  out
                        sum of s.p. energies
```

Determines by bracketing chemical potential muloc for given array of s.p. energies, temperature T, and wanted number of particles occrefloc with precision 1D-12. Delivers with it thermal occupation numbers and corresponding total particle number and sum of s.p. energies.

SUBROUTINE Calc_psi1(psi1,aloc,rhotot0,rhototloc,curr0,curr1,j,lambda,mu,lambdaj,muj,sumvar2,ecombined with internal SUBROUTINE calc_hamiltonien.

```
number of DCMF iteration, used here for print
                                in
j
                                       Lagrange parameter for density for spin up&down
lambda(1:kdfull2,1:2)
                                in
lambdaj(1:kdfull2,1:3)
                                in
                                       Lagrange parameter for current
mu, muj
                                in
                                       driving parameter for augmented Lagrangian
aloc(1:2*kdfull2)
                                       local potentials for spin up and down
                                in
rhoto0(1:kdfull2,1:2)
                                       initial density PGR2all: not used ??
                                in
                                       wanted current
curr0(1:kdfull2,1:3)
                                in
psi1(1:kdfull2,1:kstate)
                              in/out
                                       set of s.p. wavefunctions iterated
rhototloc(1:kdfull2,1:2)
                                out
                                       actual density according to psi1
                                       actual current from psi1
curr1(1:kdfull2,1:3)
                                out
                                       final s.p. energies
ekmod(1:nstate)
                                out
eal
                                       final sum of s.p. energies
                                out
                                       variance of s.p. energies
sumvar2
                                out
```

Performs one damped gradient step of with density & current constrained Hamiltonian. PGR2all: The density array distinguishes spin up/down while the current array does not. Reason?

PGRcommThe IN & OUT assignments in this subroutine have to be updated.

SUBROUTINE eqstate(psi,aloc,rho,psi1,occuporth,iterat)

```
iterat
                                 in
                                       actual iteration number (for printing)
                                       initial set of s.p. wavefunctions
psi(1:kdfull2,1:kstate)
                                 in
psi1(1:kdfull2,1:kstate)
                                out
                                       final set of s.p. wavefunctions
aloc(1:2*kdfull2)
                                       local part of potential, spin up/down stacked in blocks
                              in/out
rho(1:2*kdfull2)
                                       initial density, spin up/down stacked in blocks
                                 in
occuporth(1:kstate)
                                in
                                       occupation numbers for psi and still the same for psi1.
```

DCMF iterations by reapeatedly calling Calc_psi1, updating Lagrangian parameters for density & current constraints, and occassionally tuning temperature to achieve correct energy. The latter is done by calling fermi1. The local potential is kept constant during DCMF iteration and updated only at the very end.

PGR2all: Fetches nr. of spin up/down from m_params.

PGR2all: Lagrange parameters are started from scratch. May it be faster to recycle the previous Lagrange parameters?

PGR2all: Density rho is entered via list and still recomputed as rhotot0. Unnecessary doubling?

SUBROUTINE OccupTO(occloc, esploc, Estar)

```
esploc(1:nstate) in given s.p. energies
occloc(1:nstate) in given occupation numbers
Estar out excitation energy relative to T=0 distribution
```

Computes thermal excitation energy as difference of actual energy to the energy obtained by Fermi distribution for T=0. The latter distributions is computed for the given s.p. energies which are the same as used for the thermal state.

SUBROUTINE calcrhotot(rho,q0)

```
q0(1:kdfull2,1:kstate) in set of s.p. wavefunctions for which density is accumulated rho(kdfull2,2) out resulting density
```

Computes local density for set of wavefunctions q0. Note that two crucial information is communicated via module params, namely occup, the array of occupation numbers, ispin the array assigning spin top each s.p. state, and nstate, the number of s.p. states. PGR2all: Exploiting the sorting of spin in blocks of s.p. states, we could rewrite the code with to SUM statements.

```
SUBROUTINE calc_var(hpsi,psi1,sumvar2)
```

```
psi1(kdfull2,kstate) in set of s.p. states for which variance of s.p. energies of calculated hpsi(kdfull2,kstate) in/out array H \to \psi_{\alpha}, on input in k-space, on output in r-space sumvar2 out summed variance of s.p. energies
```

Computes the sum of variances of the s.p. energies, $\langle \hat{\Delta h}^2 | rangle$.

PGRcommThe routine projects from each $hath\psi_{\alpha}$ all s.p. states ψ_{β} from the pool of states. That is too much. The s.p. variance should be $\sum_{\alpha} \langle |\psi_{\alpha}| (\hat{h} - \varepsilon_{\alpha})^{2} |\psi_{\alpha}\rangle$ where $\varepsilon_{\alpha} = \langle |\psi_{\alpha}| \hat{h} |\psi_{\alpha}\rangle$.

SUBROUTINE forceTemp(amoy,occup,n,temp,mu)

```
amoy(1:n) in given s.p. energies
occup(1:n) in given thermal occupation
n in number of s.p. states
temp in temperature
mu out emerging chemical potential
```

Determines chemical potential for given s.p. energies and temperature by call to OccT1. PGR2all: Obsolete and never used.

```
SUBROUTINE fermi_init(ekmod,T,occup,ispinact)

ekmod(1:nstate) in given s.p. energies

T in given temperature

ispinact in actual spin

occup(1:kstate) in/out initial occupation and resulting Fermi distribution for T.
```

Determines Fermi distribution for given s.p. energies and temperature. Searches appropriate chemical potential mu by bracketing. Use for repreated calls to FUNCTION occ. PGR2all: This routine fermi_init and the related FUNCTION occ are never used, thus obsolete. May be removed.

SUBROUTINE srhomat(psi,aloc,psiorth,occuporth)

```
psi(1:kdfull2,1:kstate) in set of s.p. wavefunctions, not orth-normalized
psiorth(1:kdfull2,1:kstate) out ortho-normalized natural orbitals
aloc(1:2*kdfull2) in actual local potential
occuporth(1:kstate) out occupation numbers for ortho-normalized states
```

Computes the density matrix of initial state goiven by set of wavefunctions psi together with their occupations occup, the latter communicated through module params. Then diagonalizes the density matrix and computes on psiorth the new wavefunctions associated with diagonal representation of the density matrix.

Finally updates running transformation matrix psitophi which is communicated and stored through module params.

PGR2all: Usage and propagation of psitophi is somewhat hidden because it is handled through a module. Needs to be explained somwhere.

```
SUBROUTINE scalar(tab1,tab2,scal,ispin, mess)
```

```
tab1(1:kdfull2,1:kstate) in 1. set of s.p. wavefunctions
tab2(1:kdfull2,1:kstate) in 2. set of s.p. wavefunctions
ispin(1:nstate) in spin of s.p. states
mess in message for print inside routine
scal(nstate,nstate) out matrix of wavefunction overlaps
```

SUBROUTINE cdiagspin(mat, eigen, vect, N)

```
mat(N,N) in complex Hermitean matrix to be diagonalized
N in dimension of matrix
eigen(N) out resulting eigenbyalues
Vect(N,N) out resulting eigenstates
```

Driver routine for diagonalization of a complex Hermitean matrix of dimension N which consists in a two blocks for separate spin. Refers for each single block to routine cdiag and subsequent library routines contained therein.

SUBROUTINE indexx (n,arrin,indx)

```
n in length of array
arrin(1:n) in array to be sorted
indx(1:n) out pointer array
```

Evaluates sorting of an array in ascending order.

SUBROUTINE occupPerSpin(mess,Occ)

```
mess in character variable with comment printed inside routine Occ(1:2) out total number of particles in each spin
```

Computes number of particles in each sin block. Uses nstate and occupations occup from module params.

CorrectEnergy2(Wref,Eref,w,E,Wout,nloc)

```
W(1:nloc)
                   initial occupations numbers
              in
E(1:nloc)
                    given s.p. energies
                    reference particle number to be reached
Wref
Eref
              in
                    reference sum of s.p. energies to be reached
nloc
              in
                    actual number of states
              out
                  readjusted occupation numbers
Wout (nloc)
```

Final energy correction by one step along Fermi distribution (using Taylor expansion about actual distribution), see Eq. $(??)^1$

SUBROUTINE ordo_per_spin(psi)

```
psi(1:kdfull2,1:kstate) in/out s.p. wavefunctions before and after reordering Reorder states in two blocks of spin up and down. Applies that reshuffling to all relevant field of states, s.p. wavefunctions psi, spin per state ispin, and occupations occup. PGR2all: Routine has been rendered obsolete by new initialization of states which produces immediately the correct sorting. But routine should be kept for possible later use (e.g., mixing states from different sources.
```

SUBROUTINE temperature (mu, T)

```
mu out resulting chemical potentialT out resulting temperature
```

Takes s.p. energies amoy and occupations occup from module params and fits a Fermi distribution to it. Temperature and chemical potentials of the fitted distribution are returned via list. Calls a fitting routine lmdifl using subroutine ff as argument.

SUBROUTINE ff(m,n,X,FVEC,IFLAG)

```
    X(1:n) in array handling chemical potential and temperature
    Fvec(1:m) out array of mismatches of distributions
    n in number of parameters of model, actually 2
    m in number of entries in array
    iflag in flaf possibly written (actually not used)
```

Mismatch of occup (via modules params) from Fermi distribution to given chemical potential and temperature. To be used in fitting routine lmdef1.

¹This equation from the theory part, set to be written.

```
SUBROUTINE cproject(qin,qout,ispact,q0)
```

```
qin(1:kdfull2) in s.p. wvaefunction to be projected
```

q0(1:kdfull2,1:kstate) in set of s.p. wavefunctions which is projected out from qin

ispact in spin associated with qin qout(1:kdfull2) out projected s.p. wavefunction

Projects away from qin all contributions of the set q0.

PGR2all: This routine may become obsolete if we recode the the variance in routine calc_var to meet the standard definition.

2. Formula from Ann. Phys. paper

2.1. Mean-field propagation

The starting point and dominant feature of the dynamics is the propagation at the level of the mean field. In this paper, we are dealing with the electron dynamics in metal clusters and we describe it by time-dependent density functional theory at the level of the Time-Dependent Local-Density Approximation (TDLDA) treated in the real time domain [? ?]. It is augmented by a self-interaction correction (SIC) approximated by average-density SIC (ADSIC) [?] in order to attain correct ionization properties [?] in the course of the dynamical simulation. TDLDA is formulated within the usual Kohn-Sham picture in terms of a set of occupied single-particle (s.p.) wavefunctions $\{|\phi_{\alpha}\rangle, \alpha=1...N\}$. Their dynamics is described by the time-dependent Kohn-Sham equation

$$i\partial_t |\phi_\alpha\rangle = \hat{h}[\varrho] |\phi_\alpha\rangle \tag{1}$$

where \hat{h} is the Kohn-Sham mean-field Hamiltonian which is a functional of the instantaneous local density $\varrho(\mathbf{r},t) = \sum_{\alpha} |\phi_{\alpha}(\mathbf{r},t)|^2$ [? ?]. The time evolution delivered by Eq. (1) can be expressed formally by the unitary one-body time-evolution operator

$$\hat{U}(t,t') = \hat{\mathcal{T}}\exp\left(-i\int_{t}^{t'}\hat{h}(t'')dt''\right)$$
(2a)

where $\hat{\mathcal{T}}$ is the time-ordering operator. This yields a closed expression for the time-evolution of s.p. states

$$|\phi_{\alpha}(t)\rangle = \hat{U}(t, t')|\phi_{\alpha}(t')\rangle.$$
 (2b)

So far, TDLDA propagates pure states. Dissipation which we will add later on leads inevitably to mixed states. This requires to generalize the description from fully occupied s.p. wavefunctions to a one-body density operator $\hat{\rho}$. Its representation in configuration space, i.e. in terms of a given set of s.p. states $|\varphi_i\rangle$, reads in general $\hat{\rho} = \sum_{ij} |\varphi_i\rangle \rho_{ij} \langle \varphi_j|$. By appropriate transformation of the s.p. basis, one can diagonalize the density matrix ρ_{ij} which defines what are called natural orbitals. The natural orbitals representation of the one-body density operator then reads

$$\hat{\rho} = \sum_{\alpha=1}^{\infty} |\phi_{\alpha}\rangle W_{\alpha}\langle\phi_{\alpha}| \quad . \tag{3}$$

The weights W_{α} represent the probability with which a state $|\phi_{\alpha}\rangle$ is occupied. The mean-field propagation (1) then becomes

$$i\partial_t \hat{\rho} = \left[\hat{h}[\varrho], \hat{\rho}\right] \tag{4}$$

where $\tilde{h}[\varrho]$ is formally the same as before and the local density is now computed as

$$\varrho(\mathbf{r},t) = \sum_{\alpha} W_{\alpha} |\phi_{\alpha}(\mathbf{r},t)|^{2}.$$
 (5)

The (coherent) pure mean-field propagation (4) leaves the occupation weights W_{α} unchanged and propagates only the s.p. states. The mean-field propagation of an initial state (3) then reads

$$\hat{\rho}(t) = \sum_{\alpha=1}^{\infty} |\phi_{\alpha}(t)\rangle W_{\alpha}\langle\phi_{\alpha}(t)| = \hat{U}(t,0)\hat{\rho}(0)\hat{U}^{-1}(t,0)$$
(6)

where \hat{U} is the mean-field evolution operator (2a).

2.2. RTA in quantum-mechanical framework

The generalization of the one-body phase-space distribution $f(\mathbf{r}, \mathbf{p})$ to a quantum-mechanical mean-field theory is the one-body density operator $\hat{\rho}$, or one-body density matrix $\rho(\mathbf{r}, \mathbf{r}')$ respectively. The equation of motion for $\hat{\rho}$ including dynamical correlations reads in general [? ?]

$$i\partial_t \hat{\rho} - [\hat{h}, \hat{\rho}] = \hat{I}[\hat{\rho}] .$$
 (7)

The left hand side embraces the mean-field propagation. It may be time-dependent Hartree-Fock or the widely used LDA version of TDDFT. The right-hand side consists of the quantum-mechanical collision term. Motivated by the successful semi-classical RTA, we import Eq. (??) for the quantum case as

$$\partial_t \hat{\rho} = -i[\hat{h}, \hat{\rho}] - \frac{1}{\tau_{\text{relay}}} (\hat{\rho} - \hat{\rho}_{\text{eq}}[\varrho, \mathbf{j}, E]) ,$$
 (8)

where $\hat{\rho}_{eq}$ is the density operator of the thermal equilibrium for local density $\varrho(\mathbf{r},t)$, current distribution $\mathbf{j}(\mathbf{r},t)$ and total energy E(t) given at that instant of time t. These constraining conditions are, in fact, functionals of the actual state $\hat{\rho}$, i.e. $\varrho[\hat{\rho}]$, $\mathbf{j}[\hat{\rho}]$, and $E[\hat{\rho}]$. For the diagonal representation Eq.(3) of the density operator $\hat{\rho}$, they read

$$\varrho(\mathbf{r}) = \sum_{\alpha} |\phi_{\alpha}(\mathbf{r})|^{2} W_{\alpha} \quad , \quad \mathbf{j}(\mathbf{r}) = \sum_{\alpha} W_{\alpha} \phi_{\alpha}^{*}(\mathbf{r}) \frac{\overrightarrow{\nabla} - \overleftarrow{\nabla}}{2i} \phi_{\alpha}(\mathbf{r}) \quad . \tag{9}$$

The energy E(t) is taken as the total energy because the semi-classical concept of a local kinetic energy is ambiguous in a quantum system. This RTA equation (8) looks innocent, but is very involved because many entries depend in various ways on the actual state $\hat{\rho}(t)$. The self-consistent mean field is a functional of the actual local density, i.e. $\hat{h} = \hat{h}[\varrho]$. The instantaneous equilibrium density $\hat{\rho}_{eq}$ is the solution of the stationary, thermal mean-field equations with constraint on the actual $\varrho(\mathbf{r})$, $\mathbf{j}(\mathbf{r})$ and energy E, for details see Appendix ??.

The relaxation time τ_{relax} is estimated in semi-classical Fermi liquid theory, for details see appendix ??. For the metal clusters serving as test examples in the following, it becomes

$$\frac{\hbar}{\tau_{\rm relax}} = 0.40 \frac{\sigma_{ee}}{r_s^2} \frac{E_{\rm intr}^*}{N} \quad , \tag{10}$$

where E_{intr}^* is the intrinsic (thermal) energy of the system (appendix ??), N the actual number of particles, σ_{ee} the in-medium electron-electron cross section, and r_s the effective

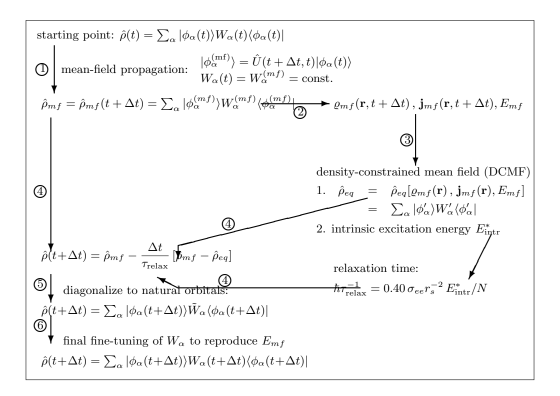


Figure 1: Sketch of the scheme for performing one large time step $t \longrightarrow t + \Delta t$ in solving the RTA equations. The numbers in open circles indicate the steps as outlined in the text.

Wigner-Seitz radius of the electron cloud. Note that r_s is tuned to the average density of the electron cloud (appendix ??), because a spatially varying $\tau_{\rm relax}$ would be very cumbersome to implement in a quantum mechanical context. This approximation is legitimate for metallic systems where the density remains generally close to the average.

2.3. Summary of the procedure

The solution of the RTA equations is rather involved. We explain the necessary steps here from a practical side and unfold details in the appendices. We briefly summarize the actual scheme for one step from t to $t+\Delta t$. Note that mean-field propagation (actually TDLDA) runs at a much faster pace than relaxation. We resolve it by standard techniques [? ?] on a time step δt which is much smaller (factor 10–100) than the RTA step Δt . We summarize this TDLDA propagation in the evolution operator \hat{U} from Eq. (2a) and discuss only one RTA step. Its sub-steps are sketched in Figure 1 and explained in the following whereby the label here correspond to the ones in the Figure:

- 1. We first propagate $\hat{\rho}$ by pure TDLDA. This means that the s.p. states in representation (3) evolve as $|\phi_{\alpha}(t)\rangle \to |\phi_{\alpha}^{(\mathrm{mf})}\rangle = \hat{U}(t+\Delta t,t)|\phi_{\alpha}(t)\rangle$, while the occupation weights W_{α} are kept frozen (pure mean-field propagation).
- 2. We compute density $\varrho(\mathbf{r}, t+\Delta t)$, current $\mathbf{j}(\mathbf{r}, t+\Delta t)$, and total energy $E_{\rm mf}$ associated to the TDLDA-propagated density matrix $\hat{\rho}_{\rm mf}$.

- 3. We determine the thermal mean-field equilibrium state $\hat{\rho}_{eq}$ constrained to the given ϱ , \mathbf{j} , and E_{mf} . This is achieved by Density-Constrained Mean Field (DCMF) iterations as outlined in Appendix ??. The actual equilibrium state $\hat{\rho}_{eq}$ is represented by new s.p. states $\{|\phi'_{\alpha}\rangle\}$ and new occupation numbers W'_{α} in diagonal form (3).
- 4. We compose the new density matrix from the TDLDA propagated state $\hat{\rho}_{\rm mf}$ and the equilibration driving term $\hat{\rho}_{\rm mf} \hat{\rho}_{\rm eq}$ with the appropriate weight $\Delta t/\tau_{\rm relax}$, as outlined in Appendix ??. The relaxation time Eq. (10) requires the actual intrinsic excitation energy $E_{\rm intr}^*$ which is also obtained from DCMF, see appendix ??.
- 5. We diagonalize the state emerging from step 4 to natural-orbital representation Eq. (3). This yields the s.p. states $\{|\phi_{\alpha}(t+\Delta t)\rangle\}$ for the next step and preliminary new occupations \tilde{W}_{α} .
- 6. After all these steps, the initial energy $E_{\rm mf} = E_{\rm TDLDA}(t)$ may not be exactly reproduced. We may remain with a small energy mismatch as compared to the goal $E_{\rm mf}$. We now apply a small iterative thermalization step to readjust the energy, as outlined in Appendix ??. This then yields the final occupation weights $W_{\alpha}(t+\Delta t)$ which comply with energy conservation.

The scheme can be used also in connection with absorbing boundary conditions [??]. The particle loss will be mapped automatically to loss of occupation weights in step 4. A word is in order about the choice of the time steps. The δt for propagation of TDLDA is limited by the maximal energy on the grid representation and thus very small (for Na clusters typically 0.005 fs). The stepping for the relaxation term needs only to resolve the changes in the actual mean field which is achieved already with $\Delta t \approx 0.5$ fs. We have tested a sequence of Δt and find the same results for all $\Delta t \leq 0.5$ fs. Changes appear slowly above that value. For reasons of efficiency, we thus use the largest safe value of $\Delta t = 0.5$ fs.

A word is in order about the range of applicability of the RTA for finite fermion systems. The relaxation time $\tau_{\rm relax}$ is allowed to depend on time which allows to accomodate changes of the dynamical state. But $\tau_{\rm relax}$ is at each instant if time one global number chosen according to the average electron density. This requires systems with only small density variations in the bulk as it holds typically for metallic bonds. The RTA is insensitive to many details of the VUU collision term as energy- and angle-dependent scattering cross sections or a broad spectrum of relaxation rates. However, these details are usually resolved only (if at all) for fast and energetic processes which are anyway deep in the regime of semi-classical VUU. The grossly averaged treatment of RTA is acceptable for not too fast and not too energetic processes in compact metallic systems.

2.4. Numerical representation and computation of relevant observables

The numerical implementation of TDLDA is done in standard manner [??]. The coupling to the ions is mediated by soft local pseudopotentials [?]. The Kohn-Sham potential is handled in the Cylindrically Averaged Pseudo-potential Scheme (CAPS) [??], which has proven to be an efficient and reliable approximation for metal clusters close to axial symmetry. Wavefunctions and fields are thus represented on a 2D cylindrical grid in coordinate space [?]. For the typical example of the Na₄₀ cluster, the numerical box extends up to 104 a₀ in radial direction and 208 a₀ along the z-axis, while the grid spacing is 0.8 a₀. To solve the (time-dependent) Kohn-Sham equations (1) we use time-splitting for time propagation [?] and accelerated gradient iterations for the stationary

solution [?]. The Coulomb field is computed with successive over-relaxation [?]. We use absorbing boundary conditions [??], which gently absorb all outgoing electron flow reaching the bounds of the grid and thus prevent artifacts from reflection back into the reaction zone. We take the exchange-correlation energy functional from Perdew and Wang [?].

A great manifold of observables can be deduced from the $\hat{\rho}(t)$ thus obtained. We will consider in the following the dipole signal, dipole spectrum, ionization, angular distribution of emitted electrons, and entropy. We focus here on the dipole moment along symmetry axis z, which is obtained from the local density as $\langle \hat{d}_z \rangle(t) = \int d^3r \, d_z(z) \varrho(r)$ where $d_z(z) = z$ is the (local) dipole operator. The dipole strength distribution is computed with the methods of spectral analysis [?]. It is attained by an instantaneous dipole-boost excitation, collecting $\langle \hat{d}_z \rangle(t)$ during propagation, and finally Fourier transforming $\langle \hat{d}_z \rangle(t)$ into frequency domain. The angular distribution of emitted electrons is obtained from recording the absorbed electrons as in TDLDA [??]. The angular distribution is characterized by the anisotropy parameter β_2 , the leading parameter in the photo-electron angular cross section $d\sigma/d\Omega \propto (1 + \beta_2 P_2(\cos(\theta) +)$ [??] where P_2 is the second order Legendre polynomial and θ the direction with respect to laser polarization axis (here z-axis in 2D cylindrical geometry). A specific quantity to track relaxation processes is the one-body entropy which is computed in diagonal representation (3) by the standard expression [?]

$$S = -\sum_{\alpha} \left[W_{\alpha} \log W_{\alpha} + (1 - W_{\alpha}) \log(1 - W_{\alpha}) \right] \tag{11}$$

in units of Boltzmann constant. It serves as a direct indicator of thermalization and allows to read off the typical time scale of relaxation processes.