Handling of the cluster 3D Fortran90-code

Instructions and status reprot

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1 Installation and usage

1.1 Installation

The following steps assume that you have successfully unpacked the code and that you are now in the sub-directory 'source_f90'.

Before compilation, one should update some settings (for detailed explanations of the parameters see section 2):

- Edit 'define.h' to choose the wanted code options.
- Edit 'params. F90' if you need to change some limiting values (rarely required).
- Edit 'makefile' and insert your compiler with its appropriate options. Some lines for that are provided and presently commented out. Fill the lines and remove the comments if needed.
- Finally execute 'make'. The executable will be copied to the working directory which is one level below the sub-directory 'source_f90'. Go back to the working directory. The last file called 'essai*' is the new executable.

1.2 Basic input structure

The cluster 3D code has six entries for options:

compile time			
define.h	variants of the code		
	run time		
for005. <name></name>	general input for settings, static and dynamics		
for005ion. <name></name>	ionic configuration of cluster		
for005surf. <name></name>	atomic configuration of substrate (optional)		
for005	defines the qualifier <name> for the other for005 files</name>		

The first two entries have to be set before compilation. The other four are read in for an actual run and can be varied from run to run. The input structure for these files is summarized in section 2.

1.3 Some practical advices

Important compile-time settings:

You have to chose the wnated options in 'define.h'.

Save and restart:

The parameters 'isave', 'istat', and 'irest' allow to switch saving wavefunctions and restarting from them.

For ismax>0 and isave>1, the static wavefunctions are saved on rsave after the static iterations. These can be used in two ways. Setting istat=1 and ismax>0 continues static iteration from rsave. Setting ismax=0, istat=1, irest=0, and, of course, itmax>0 starts a dynamical run at time zero with the static wavefunctions from rsave.

Dynamical configurations are saved on save. <name> after every isave time steps. Setting irest=1 will continue the dynamical calculation from the stage saved in save. <name>.

Diagonalization amongst occupied states:

The run time option ifhamdiag=1 activates the diagonalization of the mean-field Hamiltonian amongst the active wavefunctions in each static iteration step. This option can accelerate the convergence of the static solution significantly. *However:* At present, this method works safely only if the number of active states nstate equals the actual number of electrons. This has to be checked by the user. It may work in other cases, but may also induce oscillating iteration which nevers converges.

2 Input files

Compile time settings in define.h				
version control:				
IVERSION	define your own version number			
grid representat	grid representation of kinetic energy:			
gridfft	FFT			
findiff	finite diffences 3. order (yet unsafe)			
numerov	finite diffences 5. order (yet unsafe)			
Variants of the	Coulomb solver (for gridfft=1):			
coufou	FALR (standard)			
coudoub	exact boundary conditions			
parallele version	1:			
parayes	use parallelization for wavefunctions			
parano	produce serial code			
simpara	pseudo-parallel code, runs different inputs simultaneously			
functional and	handling of electrons:			
gunnar	Gunnarson & Lundquist			
exonly	only exchange in LDA (requires gunnar=1)			
pw92	Perdew & Wang (mostly used standard)			
fullspin	with or without explicit electron spin			
versions of SIC	for electrons:			
kli	enables KLI (incompatible with exchange and fullsic)			
directenergy	direct computation of energy (neede for kli)			
exchange	exact exchange (incompatible with kli and fullsic)			
fullsic	old full SIC (incompatible with exchange and kli)			
symmcond	old full SIC with double set technique			
twostsic	new full SIC from PhD Messud			
	Compile time settings in define.h – part 2			
options for substrate:				
raregas	enables substrates			

	Namelist GLOBAL	in for005. <nam< th=""></nam<>
	choice of system	
kxbox	nr. of grid points in x direction	
kybox	nr. of grid points in y direction	
kzbox	nr. of grid points in z direction	
	box sizes must fulfill kxbox≥kybox≥ k	zbox
kstate	maximum nr. of s.p. states	
nclust	number of QM electrons	
nion	number of cluster ions	
nspdw	number of spin down electrons	
nion2	selects type of ionic background	
	$0 \rightarrow \text{jellium background}$	
	$1 \rightarrow \text{background from ionic pseudo-pot}$	tentials
	$2 \rightarrow \text{background read in from potion.}$	
radjel	Wigner-Seitz radius of jellium backgro	
surjel	surface thickness of jellium background	
bbeta	quadrupole deformation of jellium back	
gamma	triaxiality of jellium background	
dx,dy,dz,	grid spacing (in Bohr) for the 3D nume	erical grid
	the grid size is defined before compilat	
imob	global switch to allow ionic motion (if	-
isurf	switch for Ar or MgO surface (isurf=1	,
nc	number of O cores in MgO(001)	
nk	number of Mg cations in MgO(001)	
rotclustx,y,z	vector fo angle of initial rotation of ion	ıs
	initialization of wave functions	
b2occ	deformation for initial harmonic oscilla	tor wf's
gamocc	triaxiality for initial harmonic oscillato	r wf's
deocc	shift of inital Fermi energy (determines	
shiftWFx	shift of initial wavefunctions in x direc	,
ishiftCMtoOrigin	switch to shift center of mass of cluster	to origin
ispinsep	initialize wavefunctions with some spin	<u> </u>
	convergence issues	
e0dmp	damping paramter for static solution o	f Kohn-Shahm equatio
	(typically about the energy of the lowe	st bound state)
epswf	step size for static solution of Kohn-Sh	ahm equations
-	(of order of 0.5)	_
epsoro	required variance to terminate static it	eration
-	(order of 10^{-5})	

Namelist DYNAMIC in for005.<				
numerical and physical parameters for statics and dynamics				
dt1 time step for propagating electronic wavefunctions				
ismax	maximum number of static iterations			
idyniter	switch to s.p. energy as E0DMP for 'iter>idyniter'			
ifhamdiag	diagonalization of m.f. Hamiltonian in static step			
	(presently limited to fully occupied configu	rations)		
itmax	number of time steps for electronic propaga	ation		
ifexpevol	exponential evolution 4. order instead of T	N splitting		
iffastpropag	accelerated time step in TV splitting			
	(for pure electron dynamics, interplay with	absorbing b.c. ??)		
irest	switch to restart dynamics from file 'save'			
istat switch to read wavefunctions from file 'rsave'				
	it continues static iteration for 'ismax>0)'		
	it starts dynamics from these wf's for 'is	$\max=0$		
isave	saves results after every 'isave' steps			
	on file 'rsave' in and after static iteration			
	on file 'save' in dynamic propagation			
ipseudo	switch for using pseudo-densities to represe	ent substrate		
	atoms			
ifsicp	selects type of self-interaction correction			
	0 = pure LDA, 1 = SIC-GAM, 2 = ADSIC	C; 3 = SIC-Slater;		
	4 = SIC-KLI; 5 = exact exchange; 6 = old	SIC (?);		
	7 = GSlat; 8 = full SIC.			
	for activation see switches kli, exchange,	fullsic, twostsic.		
icooltyp	type of cooling (0=none, 1=pseudo-dynam	ics,		
	2=steepest descent, 3=Monte Carlo)			
ifredmas	switch to use reduced mass for ions in dyn	amics		
ionmdtyp	ionic propagation (0=none, 1=leap-frog, 2=	=velocity Verlet)		
ntref	nr. time step after which absorbing bounds			
nabsorb	number of absorbing points on boundary (0 switches off)		

power of absorbing boundary conditions

switch to spherical mask in absorbing bounds

powabso

ispherabso

	Namelist DYNAMIC	in for005. <name< th=""></name<>
	way of excitation	
centfx	initial boost of electronic wavefuncfu	tions in x-direction
centfy	initial boost of electronic wavefuncfu	tions in x-direction
centfz	initial boost of electronic wavefuncfu	tions in x-direction
tempion	initial temperature of cluster ions	
ekmat	initial kinetic energy of substrate at	om (boost in x , in eV)
itft	choice of shape of laser pulse	
	1 = ramp laser pulse, sine switching	on/off
	2 = gaussian laser pulse	·
	$3 = \cos^2 \text{ pulse}$	
tnode	time (in fs) at which pulse computa	tion starts
deltat	length of ramp pulse (itft = 1), in	fs
tpeak	time (in fs, relative to tnode) at wh	ich peak is reached
	(for itft $= 1$ and 2 , pulse length be	ecomes 2*tpeak)
omega	laser frequency (in Ry)	
e0	laser field strength in Ry/Bohr	
e1x,e1y,e1z	orientation of pulse	
iexcit	modus of excitation (0=shifts, 1=ro	tation)
iangmo	switch to compute angular momentu	ım
irotat	axis of rotation for excitation (x=1,	y=2,z=2,xyz=4)
phirot	angle of rotation for excitation (in u	inits of degree)
phangle	angle of "rotation" into a $1ph$ state	
phphase	phase of "rotation" into a $1ph$ state	
nhstate, npstate	nr. of hole and particle state for 1ph	n excitation
_	this $1ph$ option can only be run from	n istat=1

Namelist DYNAMIC in for00			
flags for observables			
iemomsRel calculates multipole momentes of electron density			
	relative to origin (0) or c.m. of cluster (1)		
istinf	modulus for printing information in static iteration	tion	
ifspemoms	switch to compute and print spatial s.p. momen	nts	
iftransme	switch to compute and print transition m.eleme	ents	
$ifrhoint_time$	switch to slices of integrated densities for all tir	nes	
jstinf	modulus for printing information in dynamic		
jinfo	modulus for printing dynamical information on	infosp. <name></name>	
jdip	modulus for printing dipole moments on pdip.	<name></name>	
jquad	modulus for printing quadrupole moments on p	quad. <name></name>	
jesc	modulus for printing ionization pescel. <name></name>		
jenergy	modulus for printing energy information on per	ergies. <name></name>	
iflocaliz	activates computation of Becke's localization		
jelf	modulus for analyzing and printing electron loc	alization in dynamics	
	various files are written of the form pelf*. <nam< td=""><td>ne></td></nam<>	ne>	
iflocaliz	modulus for analyzing and printing electron loc	alization in statics	
jstinf	modulus for printing s.p. energies and variances	8	
jpos	modulus for printing ionic positions on pposior	n. <name></name>	
jvel	modulus for printing ionic velocities on pvelion	n. <name></name>	
jstateoverlap	switch to compute overlap of static state with		
	the state directly after dynamical initialization		

	Namelist SURFACE	in for005. <name></name>
ivdw	handling of Van-der-Waals with substrate atoms	
	$0 \Longrightarrow \text{no VdW}$	
	$1 \Longrightarrow \text{enables full computation of VdW}$	
	$2 \Longrightarrow$ enables effective VdW th	rough PsP parameters
ifadiadip	switch to adiabatic treatment of substrate dipoles global shift in x for all substrate atoms	
shiftx		
shifty,shiftz	as shiftx for y and z direction	L
mion	mass of surface anion (16 for O	in $MgO(001)$)
mkat	mass of surface kation (24.3 for	Mg in MgO(001)
me	mass of valence shell	
cspr	spring constant for interaction	between core and valence shell
chgc0	charge of (anion) core	
chge0	charge of valence shell	
chgk0	charge of cation	
sigmak	gauss width of cation	
sigmac	gauss width of core	
sigmav	gauss width of valence shell	
iUseCell	switch for reading/building latt	tice of substrate atoms
	$0 \Longrightarrow \text{lattice atoms are read in}$	from input file 'for005surf.*'
	$1 \Longrightarrow \text{lattice is built from replied}$	cating unit cell and
	lattice parameters rlattve	ec are read in (see md.F)
iPotFixed	switch for Madelung summation	,
	read/write electrostatic potenti	
	so that their run-time calculation	on can be skipped
	$0 \Longrightarrow \text{do not read}$; calculate full	ll potential at each iteration
	$1 \Longrightarrow \text{read in potFixedIon}() \text{ from}$	om previously prepared file
	$-1 \Longrightarrow \text{calculate potFixedIon}()$	
	be later read in by option	1, stop after that
	$2 \Longrightarrow \text{calculate potFixedIon()}$	at the beginning, do not write
ifmdshort	includes short range interaction	n electron—substrate
isrtyp(i,j)	type of interaction between the	different kinds of particles
	$0 \rightarrow \text{no short range interaction}$	
	$1 \to GSM \text{ core}$	
	$2 \rightarrow \text{GSM valence shell} = 1 \Longrightarrow$	Born-Mayer type
	$3 \to \text{GSM kation} = 2 \Longrightarrow \text{Argor}$	n case
	$4 \rightarrow \text{Na core}$	
	$5 \to \mathrm{DFT}$ electron	
unfixCLateralRadx	radius of cylinder with mobile of	cores
unfixELateralRadx	radius of cylinder with mobile	
fixCBelowx	fixes cores which lay below give	
iDielec switch to dielectic support		
xDielec	x below which dielectric zone is	s activated
epsDi	dielectric constant in the dielec	

Ι	Namelist PERIO	in for005. <name< th=""></name<>
ch	effective charge of ion	
amu	mass of ion in units of hydrogen	mass
dr1,dr2	radii of soft local PsP	
prho1,prho2	strenghts of soft local PsP	
crloc	radius for local part of Goedecker	r PsP
cc1,cc2	strengths for local part of Goedeo	cker PsP
r0g,r1g,r2g	radii for non-local parts of Goede	ecker PsP
h0_11g,h0_22g,h0_33g	strenghts for non-local parts of G	loedecker PsP
h1_11g,h1_22g,h2_11g	strenghts for non-local parts of G	loedecker PsP
radiong	carrier radius for projecteor in no	on-local Goedecker PsF
nrow	"row" of element \longrightarrow defines leve	el of projectors

	Namelist FSIC	in for005. <name></name>
step step size in iteration of localizing or symmetry cond		
precis	symmetry condition	
SymUtBegin nr. iteration where symmetry co		ion starts
	for pure localizing step set SymUtBeg	ginį ismax
radmaxsym	limiting value in radius division for a	actucal step

A Open ends and to-be-dones

Status of Fortran90 code development:

- All common blocks have been replaced by modules and corresponding USE command. The then appearing dependences are mapped in the makefile.
- All code is now genuinly double precision and can be complied without the autodouble option. Only exception if the FFT package fftpack.F90 which still requires the autodouble, as handled explicitely in the Makefile.
 Note that the precision is set at the header of params.F90 and used as a KIND parameter in typical Fortran90 fashion.
- The somewhat dangerous practice of reusing workspace has been abandoned. Workspace is now associated dynamically with the ALLOCATE/DEALLOCATE mechanisms.
- The compiled code works now for all box sizes and number of s.p. states as long as memory allows. The box size and maximum number of states is now entered in for005. <name> in namelist GLOBAL.

Next in Fortran90 code development:

- Remove numbered labels and GOTO in favour of CYCLE or EXIT switches.
- Exploit compact vector operations to simplify long (and nested) DO loops.
- The access USE kinetic has been given too generously. Confine that to routines which really need it.
- The module params.F90 collects practically all global variables. It should be disentangled to more specific modules with restricted access.
- There are still problems with running substrates. This case has to be tested.
- Full SIC has not yet been checked.
- The code should be slowly moved to IMPLICIT NONE.

Open problems of general nature:

- The implementation of GSlat and full SIC needs to be checked and updated if necessary.
- Check PES and PAD for the option parayes.
- Option iaddCluster is presently questionable. It may be extended to allow for initialization of cluster collisions.
- The computation of pseudo-potentials from the substrates valence electrons should be separated from the slower atomic (ionic) parts. This concerns routine calcpseudo.
- The setting for the valence-electron mass in 'vstep' may be wrong for the case of MgO.
- Check proper setting of 'time' in outputs.
- Exponential propagation should yet be certified to cooperate with ionic motion.
- Subgrids for Gaussian pseudo-densities have fixed grid size of ± 7 points. This should be made more flexible to accommodate mesh size in relation for PsP radius.
- Although not necessary for performance, one may replace DO loops by the Fortran 95 SUM construct. This will make the code more transparent.
- Present parallele version still needs to specify the number of nodes at compile time. This should be changed to allow dynamical adjustment of number of nodes.