Battery Single Particle Model v1.0

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PX915 Group Project - Group A

This is a simulation code for a half-cell single particle model (half-SPM model) written in Fortran90 and interfaced with Python. The code calculates the change in concentration in a single, spherical particle for a set simulation time from which a voltage profile can also be obtained. The relevant input parameters can be set in the Jupyter notebook 'Input_NetCDF.ipynb' file, which saves the parameters as a NetCDF file. This will be read by the programme, which runs the simulation and returns a visualisation of the concentration and (if chosen) of the voltage profile.

1.1 Getting started

A Jupyter notebook tutorial is provided to guide the user through the the main features of the simulation package. This is most easily accessed by cloning the GitHub repository to the machine that will be used to run the simulation. git clone https://github.com/HubertJN/PX915_Group_Project.git

The tutorial notebook can be accessed by moving to the directory containing the cloned repository and opening a new jupyter notebook environment.

```
cd [directory the repository has been cloned to] jupyter notebook
```

There is the option to either start a new simulation or to continue a simulation from a checkpoint file. In the first case, all the input parameters have to be set and a new input file in NetCDF format will be created. If the simulation is continued from the checkpoint file, all input parameters are kept the same as for the first run except for the simulation length and the number of simultion steps after which a new checkpoint file is created.

1.2 Setting the input parameters

The default parameters are for a positive NCM (Nickel-Cobalt-Manganese) electrode and were taken from Chen2020.

The notebook provides slider bars and an input cell that allow the user to change the input parameters within suggested ranges. There is also the option so set the parameters in a unrestricted input cell – however the user is advised to use this sensibly, since it otherwise might lead to unphysical behaviour or failure of the code.

Important consideration are:

- time step: if the time steps are set too high, the simulation might take too long to complete
- SOC and applied current: at 0% or 100% state of charge (SOC), the applied current should be set positive (charging) or negative (discharging/use), respectively
- If an anode material is being simulated, it will still be the positive electrode with respect to lithium in this half-cell SMP model and the parameters need to be set accordingly

1.3 Running the simulation

Once the input parameters are set and the 'SPM_input.nc' file has been created, the simulation can be run using the provided Makefile. To compile the code the command is:

To run the simulation, the command is:

make exe

To visualise the output, the command is:

make visual

1.4 Documention

For more information see our <u>documentation</u>. If the documentation needs to be re-generated, input the following into the command prompt.

make docs

1.5 Uncertainty Quantification

There are a variety of option to visualise the uncertainties involved with the simulation. The following commands represent the available uncertainty quantification options.

Perform sensitivity analysis and then display the results.

make sensitive

Display the results from the sensitivity analysis.

make vis_sens

Perform uncertainty propagation using random latin hypercube sampling and display the results.

make uncertain

Display the results from random latin hypercube sampling.

make uncer_vis

Perform sensitivity analysis and perform random latin hypercube sampling. Then calculate uncertainty from the standard deviations and random latin hypercube sampling. Then display results.

make sens_uncer

Visualise results of calculated uncertainties from the standard deviations and random latin hypercube sampling. make vis_sens_uncer

Calculates the uncertainty from the standard deviations and then, assuming random latin hypercube sampling has been performed, displays both.

make sens_uncer_sep

1.6 How to run the data fitting

Download the /venv/ directory, which contains all the necessary python libraries which the data fitting notebook, 'DataFit_int.ipynb', requires. The /venv/ directory is a virtual environment, and launching Jupyter notebook inside this virtual environment will allow Jupyter to use the required libraries without the user needing to install them. To activate the virtual environment, go to the directory where /venv/ is stored. This directory should also contain the solver used for data fitting, 'datafitPde.f90', and the notebook that runs the datafitting, 'DataFit_int.ipynb'. Once the /venv/ directory and the aforementioned programs are stored in the same location, the virtual environment needs to be launched. To launch the virtual environment, move to the location where it is stored in a terminal and type the following command:

source /venv/bin/activate

You can tell if the virtual environment is activated if you can see '(venv)' written before your current working directory in the terminal. Jupyter notebook can now be launched in this virtual environment by typing the following in the same terminal:

jupyter notebook

Once Jupyter is launched, find the notebook 'DataFit_int.ipynb' in the Jupyter file browser and open it. You can then run the cells accordingly to preform the optimisation on the diffusio

Finite Difference Formulation

This section outline the formulae used within the code in order to carry out the simulation. Whenever a variable or code snippet refers to the formulation section, this is the part of the code being referred to.

2.1 Matrix A

This is the left hand side coefficient matrix of the system of equations used within the simulation, of size $n \times n$ with elements given by

$$A_{i,j} = \begin{cases} -\frac{\Delta tD}{2\Delta r^2} + \frac{\Delta tD}{2r_i\Delta r} & j=i-1 \text{ for } 2\leqslant i\leqslant n-1\\ 1 + \frac{\Delta tD}{\Delta r^2} & j=i\\ -\frac{\Delta tD}{2r_i\Delta r} - \frac{\Delta tD}{2\Delta r^2} & j=i+1 \text{ for } 2\leqslant i\leqslant n-1\\ -\frac{\Delta tD}{\Delta r^2} & (i,j) = (1,2)\,, (n,n-1) \end{cases}$$

2.2 Matrix B

This is the right hand side coefficient matrix of the system of equations used within the simulation, of size $n \times n$ with elements given by

$$B_{i,j} = \begin{cases} \frac{\Delta tD}{2\Delta r^2} - \frac{\Delta tD}{2r_i\Delta r} & j=i-1 \text{ for } 2\leqslant i\leqslant n-1\\ 1 - \frac{\Delta tD}{\Delta r^2} & j=i\\ \frac{\Delta tD}{2r_i\Delta r} + \frac{\Delta tD}{2\Delta r^2} & j=i+1 \text{ for } 2\leqslant i\leqslant n-1\\ \frac{\Delta tD}{\Delta r^2} & (i,j)=(1,2)\,,(n,n-1) \end{cases}$$

Modules Index

3.1 Modules List

Here is a list of all modules with brief descriptions:

input_	_output_	_net	tcdf		 				 			 							 	 			1	11
pde s	solver				 				 			 								 			- 2	26

6 Modules Index

Data Type Index

4.1 Data Types List

Here are the data types with brief descriptions:	
volt calc	3

8 Data Type Index

File Index

5.1 File List

Here is a list of all files with brief descriptions:

input_ou	tput_netcdf.f90	
	Module file for I/O using NetCDF	35
main.f90		
	Main fortran file which calls other functions from other modules	37
pde.f90		
	Module file for solving the diffusion equation and voltage calculation	38

10 File Index

Module Documentation

6.1 input_output_netcdf Module Reference

Functions/Subroutines

• subroutine error_check (ierr)

Error checking subroutine for NetCDF.

• subroutine assign_int (file_id, act, vect, sing, var_name, var_id_in, var_id_out)

I/O subroutine that reads integers from a NetCDF file or writes integers to a NetCDF file.

• subroutine assign_real (file_id, act, vect, sing, var_name, var_id_in, var_id_out)

I/O subroutine that reads reals from a NetCDF file or writes reals to a NetCDF file.

• subroutine assign_exp_int (var, file_id, it, var_name, var_id_in)

I/O subroutine that writes integer arrays to a variable with an infinite dimension.

subroutine assign_exp_real (var, file_id, it, var_name, var_id_in)

I/O subroutine that writes real arrays to a variable with an infinite dimension.

• subroutine create_sing_var (var_name, var_typ, var_len, file_id, units, act, var_id_out)

I/O subroutine that creates a named variable with specific length and data type within a NetCDF file.

• subroutine create_exp_var (var_name, var_typ, var_len, file_id, units, act, var_id_out)

I/O subroutine that creates an expanding variable with specific dimensions and data type within a NetCDF file.

subroutine save_int (var_name, file_id, vect, sing, units, act, var_len)

I/O subroutine that writes an integer vector or single number to existing variable within a NetCDF file.

• subroutine save_real (var_name, file_id, vect, sing, units, act, var_len)

I/O subroutine that writes a real vector or single number to existing variable within a NetCDF file.

subroutine fin_in_out ()

Subroutine that closes NetCDF output and checkpoint files that are open.

subroutine import input (file name)

I/O subroutine that opens a NetCDF file, reads values, writes them to global variables and closes the file.

• subroutine initiate file (file name)

I/O subroutine that creates a NetCDF file, initiates input variables and saves variables that are available.

subroutine initiate_checkp (file_name)

I/O subroutine that creates a NetCDF file, initiates checkpoint specific variables and saves available variables.

• subroutine load_checkp (check_file_name, out_file_name, conc, volt_do)

I/O subroutine that reads from a checkpoint file and saves the concentration vector.

subroutine update_checkp (conc, step_num)

This subroutine overwrites the concentration vector (conc) and number of time steps (step_num) in the checkpoint NetCDF file.

Variables

```
    integer, parameter dp =kind(1.0D0)

    real(kind=real64), parameter farad = 96485.3321233100184_DP

      farad is the Faraday constant, F

    real(kind=real64), parameter gas_con = 8.31446261815324_DP

      gas_con is the ideal gas constant, R_g
• real(kind=real64) temp
      temp is the temperature of the simulation, T
• real(kind=real64) rad
      rad is the radius of the particle, R
• real(kind=real64) thick
• real(kind=real64) rr coef
      rr\_coef is the reaction rate coefficient, K
• real(kind=real64) dif_coef
      {\it dif\_coef} is the diffusion coefficient, D

    real(kind=real64) iapp

      iapp is the applied current density as a function of time, i_{app}

    real(kind=real64) init c

      init\_c is the initial concentration, c_0
• real(kind=real64) max c
      {\it max\_c} is the maximum concentration of the simulation, c_{\it max}

    real(kind=real64) dt

      dt is the time step
real(kind=real64) vol_per
      vol_per is the active material volume fraction, \epsilon_{actk}

    real(kind=real64) final_time

      final_time is the time of the final time step

    integer(kind=int32) sim steps

      sim_steps is the number of simulation steps
• integer(kind=int32) out_steps
      out steps is the number of steps before writing to output file

    integer(kind=int32) space_steps

      space_steps is the resolution of the radius
• integer(kind=int32) tot_steps
      tot_steps is the total number of simulation steps
• integer(kind=int32) output_id
      output_id is the id of the output file
• integer(kind=int32) check_id
      check_id is the id of the checkpoint file

    integer(kind=int32) volt out id

      volt_out_id is the variable id of the voltage in the output file
• integer(kind=int32) conc_out_id
      conc out id is the variable id of the concentration in the output ilfe
• integer(kind=int32) conc_check_id
• integer(kind=int32) time_check_id
      time_check_id is the id of the final time of the simulation in the checkpoint file

    integer(kind=int32) ts check id

      ts_check_id is the id of the final time step of the simulation in the checkpoint file

    logical volt_do
```

volt_do is the logical value which determines whether to calculate voltage to the outputfile

· logical checkpoint

checkpoint is the logical value which determines whether to restart simulation from the checkpoint

• integer(kind=int32) volt_do_int

volt_do_int is the binary representation of the volt_do variable

integer(kind=int32) checkpoint_int

checkpoint_int is the binary representation of the checkpoint_int

6.1.1 Function/Subroutine Documentation

6.1.1.1 assign_exp_int()

I/O subroutine that writes integer arrays to a variable with an infinite dimension.

This subroutine writes a 2D integer array 'var', to the variable named 'var_name' with variable id 'var_id_in' in the NetCDF file with id 'file_id' at position 'it'.

This should be used to write integer arrays 'var', to a variable 'var_name', with an infinite dimension

Parameters

in	var	2D integer array
in	var_name	name of variable to write
in	var_id_in	id of variable to write
in	file_id	id of NetCDF file to write to
in	it	position within NetCDF file to write to

Definition at line 318 of file input output netcdf.f90.

6.1.1.2 assign_exp_real()

I/O subroutine that writes real arrays to a variable with an infinite dimension.

This subroutine writes a 2D real array 'var', to the variable named 'var_name' with variable id 'var_id_in' in the NetCDF file with id 'file_id' at position 'it'.

This should be used to write real arrays 'var', to a variable 'var_name', with an infinite dimension

Parameters

in	var	2D real array
in	var_name	name of variable to write
in	var_id_in	id of variable to write
in	file_id	id of NetCDF file to write to
in	it	position within NetCDF file to write to

Definition at line 357 of file input_output_netcdf.f90.

6.1.1.3 assign_int()

I/O subroutine that reads integers from a NetCDF file or writes integers to a NetCDF file.

This subroutine reads or writes a single integer or a vector of integers, from or to a variable with a given name or variable id in a NetCDF file with a given NetCDF id.

What the subroutine does is dictated by the input arguments. One of the arguments sing and vect must be inputed into the subroutine as well as one of the arguments var_name and var_id_in.

Optionally you can save the variable id by inputting a variable to store it, var_id_out.

Parameters

in	act	subroutine action: 'r' for read and 'w' for write
in,out	sing	single integer to read/write, optional argument
in,out	vect	vector of integers to read/write, optional argument
in	var_name	name of variable to read from or write to, optional argument
in	var_id_in	id of variable to read from or write to, optional argument
in	file_id	NetCDF file id
out	var_id_out	id of variable to store var_id_in, optional argument

Definition at line 168 of file input_output_netcdf.f90.

6.1.1.4 assign_real()

I/O subroutine that reads reals from a NetCDF file or writes reals to a NetCDF file.

This subroutine reads or writes a single real or a vector of reals, from or to a variable with a given name or variable id in a NetCDF file with a given NetCDF id.

What the subroutine does is dictated by the input arguments. One of the arguments sing and vect must be inputed into the subroutine as well as one of the arguments var name and var id in.

Optionally you can save the variable id by inputting a variable to store it, var id out.

Parameters

in	act	subroutine action: 'r' for read and 'w' for write
in,out	sing	single real to read/write, optional argument
in,out	vect	vector of reals to read/write, optional argument
in	var_name	name of variable to read from or write to, optional argument
in	var_id_in	id of variable to read from or write to, optional argument
in	file_id	NetCDF file id
out	var_id_out	id of variable to store var_id_in, optional argument

Definition at line 247 of file input_output_netcdf.f90.

6.1.1.5 create_exp_var()

I/O subroutine that creates an expanding variable with specific dimensions and data type within a NetCDF file.

This subroutine creates an expanding variable called 'var_name' with dimensions 'var_len x undefined' and data type 'var_typ' (in this case f90_int or f90_double) in a NetCDF file with id 'file_id'

You can optionally prescribe units to the variable'units', and if you want to save the variable id you can input a variable to store it 'var id out'

If the file is NOT in definition mode, so already exists, you can use (act='add') to add the variable to an existing netcdf file with id 'file id'

Parameters

in	var_name	name of variable to be created in NetCDF file
in	var_len	length of variable to be created in NetCDF file
in	var_type	data type of variable to be created, f90_int or f90_double
in	file_id	id of NetCDF file where variable is being created
in	units	units of variable, optional argument
out	var_id_out	variable to store NetCDF variable id, optional argument
in	act set to 'add' to add variable to existing NetCDF file, optional argument	

Definition at line 457 of file input output netcdf.f90.

6.1.1.6 create sing var()

I/O subroutine that creates a named variable with specific length and data type within a NetCDF file.

This subroutine creates a variable called 'var_name' with length 'var_len' and data type 'var_typ' (in this case f90_int or f90_double) in a NetCDF file with id 'file_id'

You can optionally prescribe units to the variable'units', and if you want to save the variable id you can input a variable to store it 'var_id_out'

If the file is NOT in definition mode, so already exists, you can use (act='add') to add the variable to an existing netcdf file with id 'file id'

Parameters

in	var_name	name of variable to be created in NetCDF file
in	var_len	length of variable to be created in NetCDF file
in	var_type	data type of variable to be created, f90_int or f90_double
in	file_id	id of NetCDF file where variable is being created
in	units	units of variable, optional argument
out	var_id_out	variable to store NetCDF variable id, optional argument
in	act	set to 'add' to add variable to existing NetCDF file, optional argument

Definition at line 402 of file input output netcdf.f90.

6.1.1.7 error_check()

Error checking subroutine for NetCDF.

This subroutine takes in an integer error code from NetCDF (ierr), prints out the associated error, and stops the code. If there is no error, subroutine continues

Parameters

Definition at line 138 of file input_output_netcdf.f90.

6.1.1.8 fin_in_out()

```
subroutine input_output_netcdf::fin_in_out
```

Subroutine that closes NetCDF output and checkpoint files that are open.

Definition at line 575 of file input_output_netcdf.f90.

6.1.1.9 import_input()

I/O subroutine that opens a NetCDF file, reads values, writes them to global variables and closes the file.

As a test case, if no file_name is given a series of test values are prescribed instead

Parameters

```
in file_name name of NetCDF file to open
```

Definition at line 593 of file input_output_netcdf.f90.

6.1.1.10 initiate_checkp()

I/O subroutine that creates a NetCDF file, initiates checkpoint specific variables and saves available variables.

Parameters

in file_name name o	f NetCDF file to create
---------------------	-------------------------

Definition at line 714 of file input_output_netcdf.f90.

6.1.1.11 initiate_file()

I/O subroutine that creates a NetCDF file, initiates input variables and saves variables that are available.

Parameters

in file_name name of NetCDF1	file to create
------------------------------	----------------

Definition at line 659 of file input_output_netcdf.f90.

6.1.1.12 load_checkp()

I/O subroutine that reads from a checkpoint file and saves the concentration vector.

This subroutine reads from a checkpoint file named 'file_name' and saves the concentration vector to 'conc'. It extracts other variables to keep track of the total simulation steps and total simulation time. It also opens an old netcdf output file and gets variable ids for the variables it will write new data to.

Parameters

in	check_file_name	name of NetCDF checkpoint file to read
in	out_file_name	name of NetCDF file to write to
in,out	conc	concentration vector
in	volt_do	logic value to determine whether to write voltage

Definition at line 747 of file input_output_netcdf.f90.

6.1.1.13 save_int()

I/O subroutine that writes an integer vector or single number to existing variable within a NetCDF file.

This subroutine writes an integer vector 'vect' or single number 'sing' to an existing variable named 'var_name' in a NetCDF file with id 'file_id' if (act='new'), this assumes the variable does not already exist and will create an integer variable called 'var_name' with length 'var_len' and units 'units' and write the integer variable ('vect' or 'sing') to this variable.

Parameters

in	var_name name of variable to write to in NetCDF file		
in file_id id of NetCDF file where variable is being written		id of NetCDF file where variable is being written	
in,out	out vect integer vector to be written, optional argument		
in,out	sing	g single integer to be written, optional argument	
in	units units of variable, optional argument		
in	act	set to 'new' to create variable in NetCDF file, optional argument	
in	var_len	length of variable to be created in NetCDF file	

Definition at line 512 of file input_output_netcdf.f90.

6.1.1.14 save_real()

I/O subroutine that writes a real vector or single number to existing variable within a NetCDF file.

This subroutine writes a real vector 'vect' or single number 'sing' to an existing variable named 'var_name' in a NetCDF file with id 'file_id' if (act='new'), this assumes the variable does not already exist and will create an integer variable called 'var_name' with length 'var_len' and units 'units' and write the real variable ('vect' or 'sing') to this variable.

Parameters

in	var_name	name of variable to write to in NetCDF file
in	file_id	id of NetCDF file where variable is being written

Parameters

in,out	vect	integer vector to be written, optional argument	
in,out	out sing single integer to be written, optional argument		
in	units	units of variable, optional argument	
in	act	set to 'new' to create variable in NetCDF file, optional argument	
in	var_len	length of variable to be created in NetCDF file	

Definition at line 551 of file input_output_netcdf.f90.

6.1.1.15 update_checkp()

This subroutine overwrites the concentration vector (conc) and number of time steps (step_num) in the checkpoint NetCDF file.

I/O subroutine that overwrites the concentration vector and number of time steps in the checkpoint NetCDF file

Parameters

in,out	conc	concentration vector
in,out	step_num	number of time steps

Definition at line 782 of file input_output_netcdf.f90.

6.1.2 Variable Documentation

6.1.2.1 check id

```
int32 check_id
```

check_id is the id of the checkpoint file

Definition at line 125 of file input_output_netcdf.f90.

6.1.2.2 checkpoint

```
logical checkpoint
```

checkpoint is the logical value which determines whether to restart simulation from the checkpoint

Definition at line 127 of file input_output_netcdf.f90.

6.1.2.3 checkpoint_int

```
int32 checkpoint_int
```

checkpoint_int is the binary representation of the checkpoint_int

Definition at line 128 of file input_output_netcdf.f90.

6.1.2.4 conc_check_id

```
integer(kind=int32) conc_check_id
```

Definition at line 126 of file input_output_netcdf.f90.

6.1.2.5 conc_out_id

```
int32 conc_out_id
```

conc_out_id is the varialbe id of the concentration in the output ilfe

Definition at line 126 of file input_output_netcdf.f90.

6.1.2.6 dif_coef

```
real64 dif_coef
```

 dif coef is the diffusion coefficient, D

It has units of $m^2s^{-1}\,$

Definition at line 122 of file input_output_netcdf.f90.

6.1.2.7 dp

```
integer, parameter dp =kind(1.0D0)
```

Definition at line 119 of file input_output_netcdf.f90.

6.1.2.8 dt

```
real64 dt
```

dt is the time step

It has units of s

Definition at line 123 of file input_output_netcdf.f90.

6.1.2.9 farad

```
real64 farad = 96485.3321233100184_DP
```

farad is the Faraday constant, ${\cal F}$

It has units of $Cmol^{-1}$

Definition at line 120 of file input_output_netcdf.f90.

6.1.2.10 final_time

```
real64 final_time
```

final_time is the time of the final time step

It has units of \boldsymbol{s}

Definition at line 123 of file input_output_netcdf.f90.

6.1.2.11 gas_con

```
real64 gas_con = 8.31446261815324_DP
```

gas_con is the ideal gas constant, $R_{\it g}$

It has units of $JK^{-1}mol^{-1}$

Definition at line 121 of file input_output_netcdf.f90.

6.1.2.12 iapp

```
real64 iapp
```

iapp is the applied current density as a function of time, i_{app}

It has units of Am^{-2}

Definition at line 122 of file input_output_netcdf.f90.

6.1.2.13 init_c

```
real64 init_c
```

init_c is the initial concentration, c_0

It has units of $molm^{-3}$

Definition at line 123 of file input_output_netcdf.f90.

6.1.2.14 max_c

```
real64 max_c
```

max_c is the maximum concentration of the simulation, c_{max}

It has units of $molm^{-3}$

Definition at line 123 of file input_output_netcdf.f90.

6.1.2.15 out_steps

```
int32 out_steps
```

out_steps is the number of steps before writing to output file

Definition at line 124 of file input_output_netcdf.f90.

6.1.2.16 output_id

```
int32 output_id
```

output_id is the id of the output file

Definition at line 125 of file input_output_netcdf.f90.

6.1.2.17 rad

```
real64 rad
```

rad is the radius of the particle, R

It has units of m

Definition at line 122 of file input_output_netcdf.f90.

6.1.2.18 rr_coef

```
real64 rr_coef
```

 rr_coef is the reaction rate coefficient, K

It has units of $Am^2 \left(m^3 mol^{-1}\right)^{1.5}$

Definition at line 122 of file input_output_netcdf.f90.

6.1.2.19 sim_steps

```
int32 sim_steps
```

sim_steps is the number of simulation steps

Definition at line 124 of file input_output_netcdf.f90.

6.1.2.20 space_steps

```
int32 space_steps
```

space_steps is the resolution of the radius

Definition at line 124 of file input_output_netcdf.f90.

6.1.2.21 temp

```
real64 temp
```

temp is the temperature of the simulation, ${\cal T}$

It has units of K

Definition at line 122 of file input_output_netcdf.f90.

6.1.2.22 thick

```
real(kind=real64) thick
```

Definition at line 122 of file input_output_netcdf.f90.

6.1.2.23 time_check_id

```
int32 time_check_id
```

time_check_id is the id of the final time of the simulation in the checkpoint file

Definition at line 126 of file input output netcdf.f90.

6.1.2.24 tot_steps

```
int32 tot_steps
```

tot_steps is the total number of simulation steps

Definition at line 124 of file input_output_netcdf.f90.

6.1.2.25 ts_check_id

```
int32 ts_check_id
```

ts_check_id is the id of the final time step of the simulation in the checkpoint file

Definition at line 126 of file input_output_netcdf.f90.

6.1.2.26 vol_per

```
real64 vol_per
```

vol_per is the active material volume fraction, ϵ_{actk}

Definition at line 123 of file input_output_netcdf.f90.

6.1.2.27 volt_do

```
logical volt_do
```

volt_do is the logical value which determines whether to calculate voltage to the outputfile

Definition at line 127 of file input output netcdf.f90.

6.1.2.28 volt_do_int

```
int32 volt_do_int
```

volt do int is the binary representation of the volt do variable

Definition at line 128 of file input_output_netcdf.f90.

6.1.2.29 volt out id

```
int32 volt_out_id
```

volt_out_id is the variable id of the voltage in the output file

Definition at line 126 of file input_output_netcdf.f90.

6.2 pde solver Module Reference

Functions/Subroutines

• subroutine setup_crank_nicholson (A, B)

Subroutine to setup the matrices of the system of equations.

- real(real64) function, dimension(:), allocatable crank_nicholson (A, B, c_cur)
 - Crank-Nicholson function.
- real(real64) function u_scalar (x)

Function to calculate the positive electrode OCV curve, U(c).

real(real64) function, dimension(:), allocatable u_arr (x)

Array version fo the U_scalar function.

real(real64) function volt_scalar (cin)

Scalar voltage calculator.

• real(real64) function, dimension(:), allocatable volt_array (arrin)

Array voltage calculator.

Variables

```
• real(kind=real64) rhs_const 

Rescaled source at the flux boundary given by: \left(dt+\frac{dt}{dr}\right)\frac{200Ri_{app}}{3\epsilon_{actk}FL}.
• real(kind=real64) volt_con_ial 

This is a combination of parameters given by: \frac{100i_{app}R}{3\epsilon_{actk}L}.
• real(kind=real64) volt_con_rtf 

This is a combination of parameters given by: \frac{2R_gT}{F}.
• real(kind=real64) mod_dif 

The rescaled diffusion coefficient given by: \frac{D}{R^2}
```

6.2.1 Function/Subroutine Documentation

6.2.1.1 crank nicholson()

Crank-Nicholson function.

Solves the diffusion equation with a constant diffusion coefficient and a constant i_{app} using the Crank-Nicholson algorithm. The function uses the LAPACK library, calling the dgesv function for solving systems of linear equations to evolve the given state by one timestep.

Parameters

in	Α	The left hand side coefficient matrix of the system of equations (please refer to formulation section for specific details on the matrix elements)
in	В	The right hand side coefficient matrix of the system of equations (please refer to formulation section for specific details on the matrix elements)
in	c_cur	The current time concentration array inputed into the Crank-Nicholson solver.

Definition at line 115 of file pde.f90.

6.2.1.2 setup_crank_nicholson()

Subroutine to setup the matrices of the system of equations.

This subroutine takes in the corresponding matrices and assigns their elements as the appropriate coefficients in the discretised diffusion equation using the Crank-Nicholson scheme.

Parameters

in	Α	The left hand side coefficient matrix of the system of equations (please refer to formulation section for specific details on the matrix elements)
in	В	The right hand side coefficient matrix of the system of equations (please refer to formulation section for specific details on the matrix elements)

Definition at line 40 of file pde.f90.

6.2.1.3 u_arr()

```
real(real64) function, dimension(:), allocatable pde_solver::u_arr ( real(real64), dimension(:), intent(in) x)
```

Array version fo the U_scalar function.

Parameters

```
x Array version of the stoichiometry.
```

Definition at line 211 of file pde.f90.

6.2.1.4 u_scalar()

Function to calculate the positive electrode OCV curve, U(c).

OCV stands for Open Circuit Voltage. Please refer to the paper, Chang-Hui Chen et. al. 2020 J. Electrochem Soc. 167 080534

Parameters

in	Х	The stoichiometry

Definition at line 190 of file pde.f90.

6.2.1.5 volt_array()

Array voltage calculator.

Calculates array of voltages when given an array input of concentration.

Parameters

in <i>arrin</i>	array input of concentrations
-----------------	-------------------------------

Definition at line 268 of file pde.f90.

6.2.1.6 volt_scalar()

Scalar voltage calculator.

Calculates scalar voltage when given a scalar input of concentration

Parameters

in <i>cin</i>	input concentration
---------------	---------------------

Definition at line 242 of file pde.f90.

6.2.2 Variable Documentation

6.2.2.1 mod_dif

```
real64 mod_dif
```

The rescaled diffusion coefficient given by: $\frac{D}{R^2}$

Definition at line 25 of file pde.f90.

6.2.2.2 rhs_const

```
real64 rhs_const
```

Rescaled source at the flux boundary given by: $\left(dt+\frac{dt}{dr}\right)\frac{200Ri_{app}}{3\epsilon_{actk}FL}$

Definition at line 25 of file pde.f90.

6.2.2.3 volt_con_ial

real64 volt_con_ial

This is a combination of parameters given by: $\frac{100i_{app}R}{3\epsilon_{actk}L}.$

Definition at line 25 of file pde.f90.

6.2.2.4 volt_con_rtf

real64 volt_con_rtf

This is a combination of parameters given by: $\frac{2R_gT}{F}.$

Definition at line 25 of file pde.f90.

Data Type Documentation

7.1 volt_calc Interface Reference

Public Member Functions

- · volt_scalar
- volt_array

7.1.1 Detailed Description

Definition at line 15 of file main.f90.

7.1.2 Member Function/Subroutine Documentation

7.1.2.1 volt_array()

volt_array

7.1.2.2 volt_scalar()

volt_scalar

The documentation for this interface was generated from the following file:

• main.f90

File Documentation

8.1 Formula.md File Reference

8.2 input_output_netcdf.f90 File Reference

Module file for I/O using NetCDF.

Modules

· module input_output_netcdf

Functions/Subroutines

• subroutine error_check (ierr)

Error checking subroutine for NetCDF.

- subroutine assign_int (file_id, act, vect, sing, var_name, var_id_in, var_id_out)
 - I/O subroutine that reads integers from a NetCDF file or writes integers to a NetCDF file.
- subroutine assign_real (file_id, act, vect, sing, var_name, var_id_in, var_id_out)
 - I/O subroutine that reads reals from a NetCDF file or writes reals to a NetCDF file.
- subroutine assign_exp_int (var, file_id, it, var_name, var_id_in)
 - I/O subroutine that writes integer arrays to a variable with an infinite dimension.
- subroutine assign_exp_real (var, file_id, it, var_name, var_id_in)
 - I/O subroutine that writes real arrays to a variable with an infinite dimension.
- subroutine create_sing_var (var_name, var_typ, var_len, file_id, units, act, var_id_out)
 - I/O subroutine that creates a named variable with specific length and data type within a NetCDF file.
- subroutine create_exp_var (var_name, var_typ, var_len, file_id, units, act, var_id_out)
 - I/O subroutine that creates an expanding variable with specific dimensions and data type within a NetCDF file.
- subroutine save_int (var_name, file_id, vect, sing, units, act, var_len)
 - I/O subroutine that writes an integer vector or single number to existing variable within a NetCDF file.
- subroutine save_real (var_name, file_id, vect, sing, units, act, var_len)
 - I/O subroutine that writes a real vector or single number to existing variable within a NetCDF file.
- subroutine fin_in_out ()
 - Subroutine that closes NetCDF output and checkpoint files that are open.

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• subroutine import_input (file_name)

I/O subroutine that opens a NetCDF file, reads values, writes them to global variables and closes the file.

• subroutine initiate_file (file_name)

I/O subroutine that creates a NetCDF file, initiates input variables and saves variables that are available.

subroutine initiate_checkp (file_name)

I/O subroutine that creates a NetCDF file, initiates checkpoint specific variables and saves available variables.

subroutine load_checkp (check_file_name, out_file_name, conc, volt_do)

I/O subroutine that reads from a checkpoint file and saves the concentration vector.

• subroutine update_checkp (conc, step_num)

This subroutine overwrites the concentration vector (conc) and number of time steps (step_num) in the checkpoint NetCDF file.

Variables

```
    integer, parameter dp =kind(1.0D0)
```

real(kind=real64), parameter farad = 96485.3321233100184 DP

farad is the Faraday constant, F

real(kind=real64), parameter gas_con = 8.31446261815324_DP

gas_con is the ideal gas constant, R_g

real(kind=real64) temp

temp is the temperature of the simulation, T

real(kind=real64) rad

rad is the radius of the particle, R

- real(kind=real64) thick
- real(kind=real64) rr_coef

 rr_coef is the reaction rate coefficient, K

• real(kind=real64) dif_coef

 $\it dif_coef$ is the diffusion coefficient, D

• real(kind=real64) iapp

iapp is the applied current density as a function of time, i_{app}

real(kind=real64) init c

 $\mathit{init_c}$ is the initial concentration, c_0

• real(kind=real64) max_c

 max_c is the maximum concentration of the simulation, c_{max}

• real(kind=real64) dt

dt is the time step

real(kind=real64) vol_per

vol_per is the active material volume fraction, ϵ_{actk}

real(kind=real64) final_time

final time is the time of the final time step

• integer(kind=int32) sim steps

sim_steps is the number of simulation steps

integer(kind=int32) out_steps

out_steps is the number of steps before writing to output file

• integer(kind=int32) space_steps

space_steps is the resolution of the radius

• integer(kind=int32) tot_steps

tot steps is the total number of simulation steps

integer(kind=int32) output_id

output_id is the id of the output file

• integer(kind=int32) check_id

check_id is the id of the checkpoint file

• integer(kind=int32) volt out id

volt_out_id is the variable id of the voltage in the output file

integer(kind=int32) conc_out_id

conc_out_id is the varialbe id of the concentration in the output ilfe

- integer(kind=int32) conc_check_id
- integer(kind=int32) time check id

time_check_id is the id of the final time of the simulation in the checkpoint file

• integer(kind=int32) ts_check_id

ts check id is the id of the final time step of the simulation in the checkpoint file

· logical volt_do

volt_do is the logical value which determines whether to calculate voltage to the outputfile

· logical checkpoint

checkpoint is the logical value which determines whether to restart simulation from the checkpoint

• integer(kind=int32) volt do int

volt_do_int is the binary representation of the volt_do variable

integer(kind=int32) checkpoint_int

checkpoint_int is the binary representation of the checkpoint_int

8.2.1 Detailed Description

Module file for I/O using NetCDF.

This contains the subroutines necessary for reading and writing NetCDF files in the appropriate format.

8.3 main.f90 File Reference

Main fortran file which calls other functions from other modules.

Data Types

· interface volt calc

Functions/Subroutines

program main

8.3.1 Detailed Description

Main fortran file which calls other functions from other modules.

This main program calls the functions and subroutines from the pde.f90 file to solve the diffusion equation, given an input netcdf file. It contains a checkpoint system as a failsafe during simulation runtime so that the entire simulation is not lost in the event of an unforeseen error.

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8.3.2 Function/Subroutine Documentation

8.3.2.1 main()

program main

Definition at line 1 of file main.f90.

8.4 pde.f90 File Reference

Module file for solving the diffusion equation and voltage calculation.

Modules

module pde_solver

Functions/Subroutines

• subroutine setup_crank_nicholson (A, B)

Subroutine to setup the matrices of the system of equations.

real(real64) function, dimension(:), allocatable crank_nicholson (A, B, c_cur)

Crank-Nicholson function.

real(real64) function u_scalar (x)

Function to calculate the positive electrode OCV curve, U(c).

real(real64) function, dimension(:), allocatable u_arr (x)

Array version fo the U_scalar function.

• real(real64) function volt_scalar (cin)

Scalar voltage calculator.

real(real64) function, dimension(:), allocatable volt_array (arrin)

Array voltage calculator.

Variables

• real(kind=real64) rhs_const

Rescaled source at the flux boundary given by: $\left(dt+rac{dt}{dr}
ight)rac{200Ri_{app}}{3\epsilon_{actk}FL}$.

• real(kind=real64) volt_con_ial

This is a combination of parameters given by: $\frac{100i_{app}R}{3\epsilon_{actk}L}$.

• real(kind=real64) volt_con_rtf

This is a combination of parameters given by: $\frac{2R_gT}{E}$.

• real(kind=real64) mod dif

The rescaled diffusion coefficient given by: $\frac{D}{R^2}$

8.4.1 Detailed Description

Module file for solving the diffusion equation and voltage calculation.

This contains the subroutines necessary for evolving the state of the system under diffusion and calculating the voltage for each simulation timestep.

8.5 README.md File Reference

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