celestine

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BOUNDARY CONDITIONS

```
celestine.boundary_conditions.dissolved_gas_BCs(dissolved_gas_centers, cfg: Config)
     Add ghost cells with BCs to center quantity
celestine.boundary_conditions.enthalpy_BCs(enthalpy_centers, cfg: Config)
     Add ghost cells with BCs to center quantity
celestine.boundary_conditions.gas_BCs(gas_centers, cfg: Config)
     Add ghost cells with BCs to center quantity
celestine.boundary_conditions.gas_fraction_BCs(gas_fraction_centers, cfg: Config)
     Add ghost cells with BCs to center quantity
celestine.boundary_conditions.liquid_fraction_BCs(liquid_fraction_centers, cfg: Config)
     Add ghost cells to liquid fraction such that top and bottom boundaries take the same value as the top and bottom
     cell center
celestine.boundary_conditions.liquid_salinity_BCs(liquid_salinity_centers, cfg: Config)
     Add ghost cells with BCs to center quantity
celestine.boundary_conditions.salt_BCs(salt_centers, cfg: Config)
     Add ghost cells with BCs to center quantity
celestine.boundary_conditions.temperature_BCs(temperature_centers, time, cfg: Config)
     Add ghost cells with BCs to center quantity
     Note this needs the current time as well as top temperature is forced.
```

TWO

ENTHALPY METHOD

class celestine.enthalpy_method.EnthalpyMethod(physical_params: PhysicalParams)

Template for an enthalpy method. To implement a new method overwrite the initializer to initialise the physical parameters and a suitable phase boundaries object. Then implement a calculate enthalpy method that takes a state and uses bulk enthalpy, salt and gas to return (temperature, liquid_fraction, gas_fraction, solid_fraction, liquid_salinity, dissolved_gas).

class celestine.enthalpy_method.FullEnthalpyMethod(physical_params: PhysicalParams)

THREE

FLUX

 ${\tt celestine.flux.calculate_conductive_heat_flux} ({\it temperature}, D_g)$

Calculate conductive heat flux as

$$-\frac{\partial \theta}{\partial z}$$

Parameters

- temperature (Numpy Array of size I+2) temperature including ghost cells
- **D_g** (Numpy Array) difference matrix for ghost grid

Returns

conductive heat flux

 $\verb|celestine.flux.calculate_diffusive_salt_flux|| \textit{liquid_salinity}, \textit{liquid_fraction}, \textit{D_g}, \textit{cfg})|$

Take liquid salinity and liquid fraction on ghost grid and interpolate liquid fraction geometrically

6 Chapter 3. Flux

CHAPTER FOUR

FORCING

8 Chapter 4. Forcing

FIVE

GRIDS

celestine.grids.add_ghost_cells(centers, bottom, top)

Add specified bottom and top value to center grid

Parameters

- centers (Numpy array) numpy array on centered grid (size I).
- **bottom** (*float*) bottom value placed at index 0.
- top (float) top value placed at index -1.

Returns

numpy array on ghost grid (size I+2).

celestine.grids.average(ghosts)

Returns arithmetic mean pairwise of first dimension of an array

This should get values on the ghost grid and returns the arithmetic average onto the edge grid celestine.grids.geometric(ghosts)

Returns geometric mean of the first dimension of an array

10 Chapter 5. Grids

CHAPTER	
SIX	

LOGGING CONFIG

SEVEN

PARAMS

Classes containing parameters required to run a simulation

The config class contains all the parameters needed to run a simulation as well as methods to save and load this configuration to a yaml file.

```
 \textbf{class} \ \ \textbf{celestine.params.BoundaryConditionsConfig} (\textit{far\_gas\_sat: float} = 1.0, \textit{far\_temp: float} = 0.1, \\ \textit{far\_bulk\_salinity: float} = 0)
```

values for bottom (ocean) boundary

```
class celestine.params.Config(name: str, physical_params: PhysicalParams =
```

PhysicalParams(expansion_coefficient=0.029, concentration_ratio=0.17, stefan_number=4.2, lewis_salt=inf, lewis_gas=inf, frame_velocity=0),

boundary_conditions_config: BoundaryConditionsConfig =

BoundaryConditionsConfig(far_gas_sat=1.0, far_temp=0.1, far_bulk_salinity=0), darcy_law_params: DarcyLawParams =

DarcyLawParams(B=100, bubble radius scaled=1.0,

pore_throat_scaling=0.5, drag_exponent=6.0, liquid_velocity=0.0),

forcing config: ForcingConfig =

ForcingConfig(temperature_forcing_choice='constant',

constant_top_temperature=- 1.5, offset=- 1.0, amplitude=0.75, period=4.0),

 $numerical\ params:\ NumericalParams = NumericalParams(I=50,$

timestep=0.0002, regularisation=1e-06, solver='LU'), $total_time$: float=4.0,

 $savefreq: float = 0.0005, data_path: str = 'data/')$

contains all information needed to run a simulation and save output

this config object can be saved and loaded to a yaml file.

```
\textbf{class} \ \ \textbf{celestine.params.DarcyLawParams} \ (B: float = 100, bubble\_radius\_scaled: float = 1.0, bubble\_radius\_sca
```

pore_throat_scaling: float = 0.5, drag_exponent: float = 6.0,

 $liquid_velocity: float = 0.0$)

non dimensional parameters for calculating liquid and gas darcy velocities

class celestine.params.ForcingConfig(temperature_forcing_choice: str = 'constant',

 $constant_top_temperature$: float = -1.5, offset: float = -1.0,

amplitude: float = 0.75, period: float = 4.0)

choice of top boundary (atmospheric) forcing and required parameters

class celestine.params.NumericalParams(I: int = 50, timestep: float = 0.0002, regularisation: float = 1e-06, solver: <math>str = 'LU')

parameters needed for discretisation and choice of numerical method

property Courant

This number must be <0.5 for stability of temperature diffusion terms

class celestine.params.PhysicalParams(expansion_coefficient: float = 0.029, concentration_ratio: float = 0.17, stefan_number: float = 4.2, lewis_salt: float = inf, lewis_gas: float = inf, frame_velocity: float = 0)

non dimensional numbers for the mushy layer

14 Chapter 7. Params

EIGHT

PHASE BOUNDARIES

class celestine.phase_boundaries.**FullPhaseBoundaries**(*physical_params*: PhysicalParams) calculates the phase boundaries when we include gas fraction in bulk enthalpy and bulk salinity.

 $\textbf{class} \ \ \textbf{celestine.phase_boundaries.PhaseBoundaries} (\textit{physical_params}: \ PhysicalParams)$

Template for phase boundary calculation.

Concrete implementations should use the state containing enthalpy, salt and gas to calculate the liquidus, enthalpy, solidus and saturation boundaries and then return masks for each possible phase of the system.

NINE

RUN SIMULATION

celestine.run_simulation.run_batch(list_of_cfg)

Run a batch of simulations from a list of configurations.

Each simulation name is logged, as well as if it successfully runs or crashes. Output from each simulation is saved in a .npz file.

Parameters

list_of_cfg (List[celestine.params.Config]) - list of configurations

TEN

STATE

Classes to store solution variables

State: store variables on cell centers StateBCs: add boundary conditions in ghost cells to cell center variables Solution: store primary variables at each timestep we want to save data

```
class celestine.state.Solution(cfg: Config)
    store solution at specified times on the center grid
    add_state(state: State, index: int)
        add state to stored solution at given time index
class celestine.state.State(cfg: Config, time, entited)
```

class celestine.state(cfg: Config, time, enthalpy, salt, gas, pressure=None)

Stores information needed for solution at one timestep on cell centers

class celestine.state.StateBCs(state: State)

Stores information needed for solution at one timestep with BCs on ghost cells as well

Note must initialise once enthalpy method has already run on State.

20 Chapter 10. State

ELEVEN

VELOCITIES

celestine.velocities.calculate_bubble_radius(liquid_fraction, cfg: Config)

Takes liquid fraction on edges and returns bubble radius parameter on edges

celestine.velocities.calculate_gas_interstitial_velocity(liquid_fraction, pressure, D_g, cfg: Config)

Calculate Vg from liquid fraction and pressure on ghost grid

Return Vg on edge grid

celestine.velocities.calculate_liquid_darcy_velocity(liquid_fraction, pressure, D_g)

Calculate liquid Darcy velocity as

$$W_l = -\Pi(\phi_l) \frac{\partial p}{\partial z}$$

Parameters

- liquid_fraction (Numpy Array (size I+2)) liquid fraction on ghost grid
- pressure (Numpy Array (size I+2)) pressure on ghost grid
- **D_g** (Numpy Array (size I+2)) difference matrix for ghost grid

Returns

liquid darcy velocity on edge grid

celestine.velocities.calculate_velocities(state_BCs, D_g, cfg: Config)

Inputs on ghost grid, outputs on edge grid

celestine.velocities.solve_pressure_equation($state_BCs$, new_state_BCs , timestep, D_e , D_g , cfg:

Config)

Calculate pressure on ghost grid from current and new state on ghost grid

Return new pressure on centers but easy to add boundary conditions

TWELVE

LAGGED SOLVER

class celestine.solvers.lagged_solver.LaggedUpwindSolver(cfg: Config)

Take timestep using upwind scheme with liquid velocity calculation lagged.

pre_solve_checks()

Optionally implement this method if you want to check anything before running the solver.

For example to check the timestep and grid step satisfy some constraint.

take_timestep(state: State)

advance enthalpy, salt, gas and pressure to the next timestep.

Note as of 2023-05-17 removed ability to have adaptive timestepping for simplicity.

Parameters

state (celestine.solvers.template.State) — object containing current time, enthalpy, salt, gas, pressure and surface temperature.

Returns

state of system (new enthalpy, salt, gas and pressure) after one timestep.

celestine.solvers.lagged_solver.take_forward_euler_step(quantity, flux, timestep, D_e)

Advance the given quantity one forward Euler step using the given flux

The quantity is given on cell centers and the flux on cell edges.

Discretise the conservation equation

$$\frac{\partial Q}{\partial t} = -\frac{\partial F}{\partial z}$$

as

$$Q^{n+1} = Q^n - \Delta t(\frac{\partial F}{\partial z})$$

THIRTEEN

TEMPLATE

Template for a solver concrete solvers should inherit and overwrite required methods

class celestine.solvers.template.SolverTemplate(cfg: Config)

generate_initial_solution()

Generate initial solution on the ghost grid

Returns

initial solution arrays on ghost grid (enthalpy, salt, gas, pressure)

pre_solve_checks()

Optionally implement this method if you want to check anything before running the solver.

For example to check the timestep and grid step satisfy some constraint.

abstract take_timestep(state: State) $\rightarrow State$

advance enthalpy, salt, gas and pressure to the next timestep.

Note as of 2023-05-17 removed ability to have adaptive timestepping for simplicity.

Parameters

state (celestine.solvers.template.State) — object containing current time, enthalpy, salt, gas, pressure and surface temperature.

Returns

state of system (new enthalpy, salt, gas and pressure) after one timestep.

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