

Numbat

High-resolution simulations of density-driven convective mixing in porous media

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1 Numbat

Version: 7118bc7 (15/06/2017)

1.1 High-resolution simulations of density-driven convective mixing in porous media

Numbat is a massively-parallel code for high-resolution simulations of density-driven convective mixing in porous media built using the MOOSE framework.

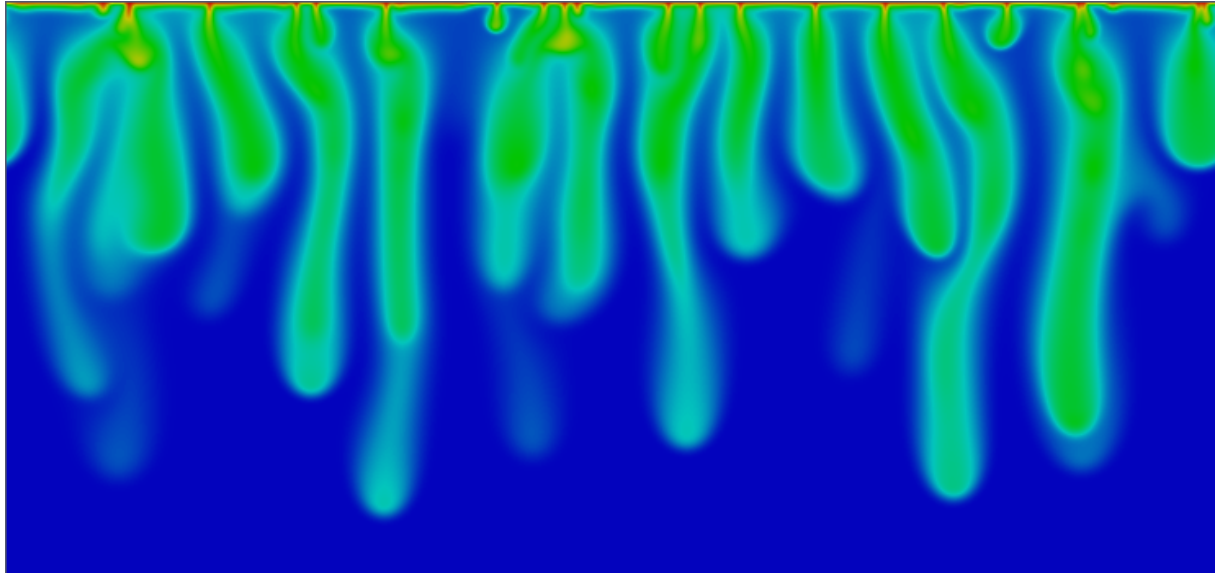


Figure 1.1: Density-driven convective mixing in a porous medium

2 Installation instructions

To install Numbat, follow these simple instructions.

2.1 Install MOOSE

Numbat is based on the MOOSE framework, so the first step is to install MOOSE. For detailed installation instructions depending on your hardware, see www.mooseframework.com.

2.2 Clone Numbat

The next step is to clone the Numbat repository to your local machine.

In the following, it is assumed that MOOSE was installed to the directory `~/projects`. If MOOSE was installed to a different directory, the following instructions must be modified accordingly.

To clone Numbat, use the following commands

```
cd ~/projects
git clone https://github.com/cpgr/numbat.git
cd numbat
git checkout master
```

2.3 Compile Numbat

Next, compile Numbat using

```
make -jn
```

where n is the number of processing cores on the computer. If everything has gone well, Numbat should compile without error, producing a binary named *numbat-opt*.

2.4 Test Numbat

Finally, to test that the installation worked, the test suite can be run using

```
./run_tests -jn
```

where n is the number of processing cores on the computer.

3 Background theory

3.1 Governing equations

Numbat implements the Boussinesq approximation to model density-driven convective mixing in porous media.

The governing equations for density-driven flow in porous media are Darcy's law

$$\mathbf{u} = -\frac{\mathbf{K}}{\mu} \left(\nabla P + \rho(c)g\hat{\mathbf{k}} \right), \quad (3.1)$$

where $\mathbf{u} = (u, v, w)$ is the velocity vector, \mathbf{K} is permeability, μ is the fluid viscosity, P is the fluid pressure, $\rho(c)$ is the fluid density as a function of solute concentration c , g is gravity, and $\hat{\mathbf{k}}$ is the unit vector in the z direction.

The fluid velocity must also satisfy the continuity equation

$$\nabla \cdot \mathbf{u} = 0, \quad (3.2)$$

and the solute concentration is governed by the convection - diffusion equation

$$\phi \frac{\partial c}{\partial t} + \mathbf{u} \cdot \nabla c = \phi D \nabla^2 c, \quad (3.3)$$

where ϕ is the porosity, t is time and D is the diffusivity.

Darcy's law and the convection-diffusion equations are coupled through the fluid density, which is given by

$$\rho(c) = \rho_0 + \frac{c}{c_0} \Delta \rho, \quad (3.4)$$

where c_0 is the equilibrium concentration, and $\Delta \rho$ is the increase in density of the fluid at equilibrium concentration.

The boundary conditions are

$$w = 0, \quad z = 0, -H, \quad (3.5)$$

$$\frac{\partial c}{\partial z} = 0, \quad z = -H, \quad (3.6)$$

$$c = c_0, \quad z = 0, \quad (3.7)$$

which correspond to impermeable boundary conditions at the top and bottom boundaries, given by $z = 0$ and $z = -H$, respectively, and a saturated condition at the top boundary.

Initially, there is no solute in the model

$$c = 0, \quad t = 0. \quad (3.8)$$

Numbat solves Eq's. (3.1) and (3.3) with density coupled to concentration as in Eq. (3.4).

3.2 Dimensionless formulation

The governing equations can also be solved using a streamfunction formulation in 2D and a vector potential formulation in 3D. As a result, we shall consider the two cases separately.

3.2.1 2D solution

If we consider an anisotropic model, with vertical and horizontal permeabilities given by k_z and k_x , respectively, we can non-dimensionalise the governing equations in 2D following [Ennis-King et. al \(2005\)](#). Defining the anisotropy ratio γ as

$$\gamma = \frac{k_z}{k_x}, \quad (3.9)$$

we scale the variables using

$$\begin{aligned} x &= \frac{\phi\mu D}{k_z\Delta\rho g\gamma^{1/2}}\hat{x}, & z &= \frac{\phi\mu D}{k_z\Delta\rho g}\hat{z}, & u &= \frac{k_z\Delta\rho g}{\mu\gamma^{1/2}}\hat{u}, & w &= \frac{k_z\Delta\rho g}{\mu}\hat{w} \\ t &= \left(\frac{\phi\mu}{k_z\Delta\rho g}\right)^2\hat{t}, & c &= c_0\hat{c}, & P &= \frac{\mu\phi D}{k_z}\hat{P}, \end{aligned} \quad (3.10)$$

where \hat{x} refers to a dimensionless variable. The governing equations in dimensionless form are then

$$\mathbf{u} = -(\nabla P + c\hat{\mathbf{k}}), \quad (3.11)$$

$$\mathbf{u} = 0, \quad (3.12)$$

$$\frac{\partial c}{\partial t} + \mathbf{u} \cdot \nabla c = \gamma \frac{\partial^2 c}{\partial x^2} + \frac{\partial^2 c}{\partial z^2}, \quad (3.13)$$

where we have dropped the hat on the dimensionless variables for brevity.

The dimensionless boundary conditions are

$$w = 0, \quad z = 0, -Ra, \quad (3.14)$$

$$\frac{\partial c}{\partial z} = 0, \quad z = -Ra, \quad (3.15)$$

$$c = 1, \quad z = 0, \quad (3.16)$$

where Ra is the Rayleigh number, defined as

$$Ra = \frac{k_z\Delta\rho g H}{\phi\mu D}. \quad (3.17)$$

In this form, the Rayleigh number only appears in the boundary conditions as the location of the lower boundary. Therefore, Ra can be interpreted in this formalism as a dimensionless model height, and can be varied in simulations by simply changing the height of the mesh.

Finally, the dimensionless initial condition is

$$c = 0, \quad t = 0. \quad (3.18)$$

For isotropic models, where $k_x = k_z$ and hence $\gamma = 1$, we recover the dimensionless equations given by [Slim \(2014\)](#).

The coupled governing equations must be solved numerically. To simplify the numerical analysis, we introduce the streamfunction $\psi(x, z, t)$ such that

$$u = -\frac{\partial\psi}{\partial z}, \quad w = \frac{\partial\psi}{\partial x}. \quad (3.19)$$

This definition satisfies the continuity equation, Eq. (3.12), immediately.

The pressure P is removed from Eq. (3.11) by taking the curl of both sides and noting that $\nabla \times \nabla P = 0$ for any P , to give

$$\nabla^2\psi = -\frac{\partial c}{\partial x}, \quad (3.20)$$

where we have introduced the streamfunction ψ using Eq. (3.19).

The convection-diffusion equation, Eq. (3.13) becomes

$$\frac{\partial c}{\partial t} - \frac{\partial\psi}{\partial z} \frac{\partial c}{\partial x} + \frac{\partial\psi}{\partial x} \frac{\partial c}{\partial z} = \gamma \frac{\partial^2 c}{\partial x^2} + \frac{\partial^2 c}{\partial z^2}. \quad (3.21)$$

The boundary conditions become

$$\frac{\partial\psi}{\partial x} = 0, \quad z = 0, -Ra, \quad (3.22)$$

$$\frac{\partial c}{\partial z} = 0, \quad z = -Ra, \quad (3.23)$$

$$c = 1, \quad z = 0, \quad (3.24)$$

while the initial condition is still given by Eq. (3.18).

In two dimensions, Numbat solves Eq's. (3.20) and (3.21).

3.2.2 3D solution

We now consider the case of a three-dimensional model. For simplicity, we consider the case where all lateral permeabilities are equal ($k_y = k_x$). The governing equations for the 3D model are identical to the 2D model. In dimensionless form, they are given by Eq's. (3.11) to (3.13), with boundary conditions given by Eq's. (3.14) to (3.16), and initial condition given by Eq. (3.18).

To solve these governing equations in 3D, a different approach must be used as the streamfunction ψ is not defined in three dimensions. Instead, we define a vector potential $\Psi = (\psi_x, \psi_y, \psi_z)$ such that

$$\mathbf{u} = \nabla \times \Psi. \quad (3.25)$$

It is important to note that the vector potential is only known up to the addition of the gradient of a scalar ζ as

$$\nabla \times (\Psi + \nabla\zeta) = \nabla \times \Psi \quad \forall \zeta, \quad (3.26)$$

as $\nabla \times \nabla \zeta = 0$ for any scalar ζ . This uncertainty is referred to as gauge freedom, and is common in electrodynamics. Taking the curl of Eq. (3.11) and substituting Eq. (3.25), we have

$$\nabla(\nabla \cdot \Psi) - \nabla^2 \Psi = \left(-\frac{\partial c}{\partial y}, \frac{\partial c}{\partial x}, 0 \right), \quad (3.27)$$

where we have again used the fact that $\nabla \times \nabla P = 0$. If we choose $\nabla \cdot \Psi = 0$ to specify the gauge condition, this simplifies to

$$\nabla^2 \Psi = \left(\frac{\partial c}{\partial y}, -\frac{\partial c}{\partial x}, 0 \right). \quad (3.28)$$

As shown in [E and Liu \(1997\)](#), $\nabla \cdot \Psi = 0$ is satisfied throughout the domain if

$$\psi_x = \psi_y = 0, \quad z = 0, -Ra, \quad \frac{\partial \psi_z}{\partial z} = 0, \quad z = 0, -Ra.$$

The governing equations are then

$$\nabla^2 \Psi = \left(\frac{\partial c}{\partial y}, -\frac{\partial c}{\partial x}, 0 \right), \quad (3.29)$$

$$\frac{\partial c}{\partial t} + \mathbf{u} \cdot \nabla c = \gamma \left(\frac{\partial^2 c}{\partial x^2} + \frac{\partial^2 c}{\partial y^2} \right) + \frac{\partial^2 c}{\partial z^2}, \quad (3.30)$$

where the continuity is satisfied automatically because $\nabla \cdot (\nabla \times \Psi) = 0$ for any Ψ .

Finally, it is straightforward to show that $\psi_z = 0$ in order to satisfy $\nabla^2 \psi_z = 0$ and $\frac{\partial \psi_z}{\partial z} = 0$, which means that the vector potential has only x and y components,

$$\Psi = (\psi_x, \psi_y, 0), \quad (3.31)$$

and therefore the fluid velocity $\mathbf{u} = (u, v, w)$ is

$$\mathbf{u} = \left(-\frac{\partial \psi_y}{\partial z}, \frac{\partial \psi_x}{\partial z}, \frac{\partial \psi_y}{\partial x} - \frac{\partial \psi_x}{\partial y} \right). \quad (3.32)$$

Note that if there is no y dependence, Eq's. (3.29) and (3.30) reduce to

$$\nabla^2 \Psi = \left(0, -\frac{\partial c}{\partial x}, 0 \right), \quad (3.33)$$

$$\frac{\partial c}{\partial t} + \mathbf{u} \cdot \nabla c = \gamma \frac{\partial^2 c}{\partial x^2} + \frac{\partial^2 c}{\partial z^2}. \quad (3.34)$$

It is simple to show that $\nabla^2 \psi_x = 0$ and $\psi_x = 0$ at $z = 0, -Ra$ are only satisfied if $\psi_x = 0$ in the entire domain. In this case, the governing equations reduce to the two-dimensional formulation, as expected.

In three dimensions, Numbat solves Eq's. (3.29) and (3.30).

4 Input file syntax

The input file for a Numbat simulation is a simple, block-structured text file.

A working example of a 2D problem can be found at <https://github.com/cpgr/numbat/blob/master/examples/2D/2Dddc.i>

A working example of a 3D problem can be found at <https://github.com/cpgr/numbat/blob/master/examples/3D/3Dddc.i>

4.1 Essential input

Details of the minimum input file requirements are given below.

4.1.1 Mesh

All simulations must feature a mesh. For the basic model with a rectangular mesh, the built-in MOOSE *GeneratedMesh* can be used to create a suitable mesh. In 2D, the input block looks like:

```
[Mesh]
  type = GeneratedMesh
  dim = 2
  xmax = 1000
  ymin = -200
  ymax = 0
  nx = 80
  ny = 20
  bias_y = 0.7
[]
```

This creates a 2D mesh from $x = 0$ to $x = 1000$ and $y = -200$ to $y = 0$ with 80 elements in the x -direction and 20 elements in the y -direction. It is useful to have a mesh that is more refined at the top of the model, to accurately capture the initially small structure of the convective fingers. This is achieved using the built-in *bias_y* parameter.

In 3D, the Mesh block would look like:

```
[Mesh]
  type = GeneratedMesh
  dim = 3
  xmax = 200
  ymax = 200
  zmin = -200
  zmax = 0
  nx = 10
  ny = 10
  nz = 10
  bias_z = 0.7
[]
```

Again, the mesh is refined at the top of the model using the *bias_z* parameter.

4.1.2 Variables

For a 2D model, the simulation must have two variables: *concentration* and *streamfunction*. This can be implemented in the input file using the following code:

```
[Variables]
[./concentration]
order = FIRST
family = LAGRANGE
  [./InitialCondition]
    type = NumbatPerturbationIC
    variable = concentration
    amplitude = 0.01
    seed = 1
  [../]
[../]
[./streamfunction]
order = FIRST
family = LAGRANGE
initial_condition = 0.0
[../]
[]
```

Initial conditions can also be specified in the *Variables* block. In this case, the initial concentration is perturbed using a *NumbatPerturbationIC* to seed the instability.

For a 3D model, three variables are required: one *concentration* variable and two *streamfunction* variables corresponding to the *x* and *y* components. This can be implemented in the input file using:

```
[Variables]
[./concentration]
[../]
[./streamfunctionx]
[../]
[./streamfunctiony]
[../]
[]
```

4.1.3 Kernels

Three kernels are required for a 2D model: a *NumbatDarcySF* kernel for the *streamfunction* variable, a *NumbatConvectionDiffusionSF* kernel for the *concentration* variable, and a *TimeDerivative* kernel also for the *concentration* variable. An example for an isotropic model is

```
[Kernels]
[./Darcy]
type = NumbatDarcySF
variable = streamfunction
concentration_variable = concentration
[../]
```

```

[./ConvectionDiffusion]
  type = NumbatConvectionDiffusionSF
  variable = concentration
  streamfunction_variable = streamfunction
  coeff_tensor = '1 0 0 0 1 0 0 0 1'
[../]
[./TimeDerivative]
  type = TimeDerivative
  variable = concentration
[../]
[]

```

The *coeff_tensor* parameter in each convective diffusion kernel can be modified. The format of this For 3D models, an additional *NumbatDarcySF* kernel is required for the additional stream function variable. An example of the kernels block for a 3D isotropic model is

```

[Kernels]
[./Darcy]
  type = NumbatDarcySF
  variable = streamfunctionx
  concentration_variable = concentration
  component = x
[../]
[./Darcy]
  type = NumbatDarcySF
  variable = streamfunctiony
  concentration_variable = concentration
  component = y
[../]
[./ConvectionDiffusion]
  type = NumbatConvectionDiffusionSF
  variable = concentration
  streamfunction_variable = 'streamfunctionx streamfunctiony'
  coeff_tensor = '1 0 0 0 1 0 0 0 1'
[../]
[./TimeDerivative]
  type = TimeDerivative
  variable = concentration
[../]
[]

```

In the 3D case, it is important to note that the *NumbatDarcySF* kernel must specify the component that it applies to, and that the *streamfunction_variable* keyword in the *NumbatConvectionDiffusionSF* kernel must contain both *streamfunction* variables ordered by the x component then the y component.

4.1.4 Initial condition

To seed the instability, a random perturbation to the initial concentration is prescribed using the *NumbatPerturbationIC* initial condition. This can be applied in the *Variables* block in the input file (see above) using

```
[./InitialCondition]
  type = NumbatPerturbationIC
  variable = concentration
  amplitude = 0.01
  seed = 1
[../]
```

The *NumbatPerturbationIC* initial condition applies the diffusive concentration profile to the nodes for (scaled) time $t = 1$,

$$c_d(z, t = 1) = 1 + \text{erf}(z/2), \quad (4.1)$$

for $z < 0$, where $\text{erf}(z)$ is the error function.

A uniform random perturbation is then added to the diffusive concentration profile, where the amplitude of the perturbation is specified by the input file value *amplitude*.

4.1.5 Boundary conditions

Appropriate boundary conditions must be prescribed. Typically, these will be constant concentration at the top of the model domain, periodic boundary conditions on the lateral sides (to mimic an infinite reservoir), and no-flow boundary conditions at the top and bottom surfaces.

In 2D, this can be achieved using the following input block:

```
[BCs]
[./conctop]
  type = DirichletBC
  variable = concentration
  boundary = top
  value = 1.0
[../]
[./streamfunctop]
  type = DirichletBC
  variable = streamfunction
  boundary = top
  value = 0.0
[../]
[./streamfunbottom]
  type = DirichletBC
  variable = streamfunction
  boundary = bottom
  value = 0.0
[../]
[./periodic]
[./x]
  variable = 'concentration streamfunction'
  auto_direction = x
[../]
[../]
```

[]

In this case, the *conctop* boundary condition is a Dirichlet condition at the top of the model that fixes the value of concentration to unity. It is useful to note that a MOOSE *GeneratedMesh* provides descriptive names for the sides of the model (top, bottom, left, right) which can be referenced in the input file. No-flow boundary conditions are prescribed on the top and bottom surfaces by holding the *streamfunction* variable constant (in this case 0). Finally, periodic boundary conditions are applied by the *periodic* block, which specifies that both the *concentration* and *streamfunction* variables are periodic on boundaries in the *x*-direction.

A similar boundary condition block is used in 3D, except that no-flow boundaries must be imposed on both streamfunction variables, see below:

```
[BCs]
  [./conctop]
    type = DirichletBC
    variable = concentration
    boundary = front
    value = 1.0
  [../]
  [./streamfunxtop]
    type = DirichletBC
    variable = streamfunctionx
    boundary = front
    value = 0.0
  [../]
  [./streamfunxbottom]
    type = DirichletBC
    variable = streamfunctionx
    boundary = back
    value = 0.0
  [../]
  [./streamfunytop]
    type = DirichletBC
    variable = streamfunctiony
    boundary = front
    value = 0.0
  [../]
  [./streamfunybottom]
    type = DirichletBC
    variable = streamfunctiony
    boundary = back
    value = 0.0
  [../]
  [./Periodic]
    [./xy]
      variable = 'concentration streamfunctionx streamfunctiony'
      auto_direction = 'x y'
    [../]
```

```
[../]  
[]
```

4.1.6 Executioner

Each MOOSE simulation must use an *Executioner*, which provides parameters for the solve. In both 2D and 3D models, a transient *Executioner* is used, an example of which is presented below:

```
[Executioner]  
  type = Transient  
  scheme = bdf2  
  dtmin = 0.1  
  dtmax = 200  
  end_time = 2000  
  solve_type = PJFNK  
  petsc_options_iname = '-ksp_type -pc_type -pc_sub_type'  
  petsc_options_value = 'gmres asm ilu'  
  [./TimeStepper]  
    type = IterationAdaptiveDT  
    dt = 1  
    cutback_factor = 0.5  
    growth_factor = 2  
  [../]  
[]
```

Executioners are a standard MOOSE feature that are well documented on the [MOOSE website](#), so no further detail is provided here.

4.1.7 Preconditioning

A default preconditioning block is used that provides all Jacobian entries to aid convergence. This is identical for both 2D and 3D models:

```
[Preconditioning]  
  [./smp]  
    type = SMP  
    full = true  
  [../]  
[]
```

This is a standard MOOSE feature that is documented on the [MOOSE website](#), so no further detail is provided here.

4.1.8 Outputs

To provide output from the simulation, an *Outputs* block must be specified. An example is

```
[Outputs]  
  [./console]  
    type = Console  
    perf_log = true
```



```

    output_nonlinear = true
  [../]
  [./exodus]
    type = Exodus
    file_base = filename
    execute_on = 'INITIAL TIMESTEP_END FINAL'
  [../]
[]

```

In this case, some output regarding the iterations is streamed to the console, while the results are provided in an Exodus file named *filename.e*. There are a large number of output options available in MOOSE, see the [MOOSE website](#) for further details.

4.2 Optional input

While the above required blocks will enable a Numbat simulation to run, there are a number of optional input blocks that will improve the simulations and increase the amount of results provided.

4.2.1 Mesh adaptivity

MOOSE features built-in mesh adaptivity that is extremely useful in Numbat simulations. This can be included using:

```

[Adaptivity]
  marker = combomarker
  max_h_level = 2
  initial_marker = boxmarker
  initial_steps = 1
  [./Indicators]
    [./gradjumpindicator]
      type = GradientJumpIndicator
      variable = concentration
    [../]
  [../]
  [./Markers]
    [./errormarker]
      type = ErrorToleranceMarker
      coarsen = 0.0025
      refine = 0.005
      indicator = gradjumpindicator
    [../]
    [./boxmarker]
      type = BoxMarker
      bottom_left = '0 -1.0 0'
      top_right = '1000 0 0'
      inside = refine
      outside = dont_mark
    [../]
    [./combomarker]
      type = ComboMarker

```

```

        markers = 'boxmarker errormarker'
    [../]
[../]
[]

```

For details about mesh adaptivity, see the [MOOSE website](#).

4.2.2 Flux at the top boundary

The flux over the top boundary is of particular interest in many cases (especially convective mixing of CO₂). This can be calculated at each time step using a *Postprocessor*:

```

[Postprocessors]
  [./boundaryfluxint]
    type = NumbatSideFlux
    variable = concentration
    boundary = top
  [../]
[]

```

The output of the *Postprocessor* can be saved to a csv file by including the following additional sub-block in the *Outputs* block:

```

[./csvoutput]
  type = CSV
  file_base = filename
  execute_on = 'INITIAL TIMESTEP_END FINAL'
[../]

```

4.2.3 Velocity components

The velocity components in the x and y directions (in 2D), and x , y , and z directions in 3D can be calculated using the auxiliary system. These velocity components are calculated using the streamfunction(s), see the governing equations for details.

In the 2D case, two auxiliary variables, u and w , can be defined for the horizontal and vertical velocity components, respectively. Importantly, these auxiliary variables **must** have constant monomial shape functions (these are referred to as *elemental* variables, as the value is constant over each mesh element). This restriction is due to the gradient of the streamfunction variable(s) being undefined for *nodal* auxiliary variables (for example, those using linear Lagrange shape functions). An example of the input syntax for the 2D case is

```

[AuxVariables]
  [./u]
    order = CONSTANT
    family = MONOMIAL
  [../]
  [./w]
    order = CONSTANT
    family = MONOMIAL

```

```
[../]
[]
```

For the 3D case, there is an additional horizontal velocity component (v), so the input syntax is

```
[AuxVariables]
  [./u]
    order = CONSTANT
    family = MONOMIAL
  [../]
  [./v]
    order = CONSTANT
    family = MONOMIAL
  [../]
  [./w]
    order = CONSTANT
    family = MONOMIAL
  [../]
[]
```

The velocity components are calculated by *NumbatDarcyVelocitySF* AuxKernels, one for each component. For the 2D case, the input syntax is

```
[AuxKernels]
  [./uAux]
    type = NumbatDarcyVelocitySF
    variable = u
    component = x
    streamfunction_variable = streamfunction
  [../]
  [./wAux]
    type = NumbatDarcyVelocitySF
    variable = w
    component = y
    streamfunction_variable = streamfunction
  [../]
[]
```

For the 3D case, three AuxKernels are required. Note that both streamfunction variables must be given, in the correct order (x then y). An example of the input syntax is

```
[AuxKernels]
  [./uAux]
    type = NumbatDarcyVelocitySF
    variable = u
    component = x
    streamfunction_variable = 'streamfunctionx streamfunctiony'
  [../]
```

```
[./vAux]
  type = NumbatDarcyVelocitySF
  variable = v
  component = y
  streamfunction_variable = 'streamfunctionx streamfunctiony'
[../]
[./wAux]
  type = NumbatDarcyVelocitySF
  variable = w
  component = z
  streamfunction_variable = 'streamfunctionx streamfunctiony'
[../]
[]
```

5 2D examples

5.1 Isotropic models

The first 2D example is for an isotropic porous medium ($\gamma = 1$). A working example of this example can be found at https://github.com/cpgr/numbat/blob/master/examples/2D/isotropic/2Dddc_isotropic.i.

5.1.1 Input file

The complete input file for this problem is

```
[Mesh]
  type = GeneratedMesh
  dim = 2
  xmax = 1000
  ymin = -500
  ymax = 0
  nx = 80
  ny = 25
  bias_y = 0.9
[]

[Adaptivity]
  marker = errormarker
  max_h_level = 1
  [./Indicators]
    [./gradjumpindicator]
      type = GradientJumpIndicator
      variable = concentration
    [../]
  [../]
  [./Markers]
    [./errormarker]
      type = ErrorToleranceMarker
      refine = 0.005
      indicator = gradjumpindicator
    [../]
  [../]
[]

[Variables]
  [./concentration]
    order = FIRST
    family = LAGRANGE
  [./InitialCondition]
    type = NumbatPerturbationIC
    variable = concentration
    amplitude = 0.1
    seed = 1
  [../]
```

```

[../]
[/streamfunction]
    order = FIRST
    family = LAGRANGE
    initial_condition = 0.0
[../]
[]

[Kernels]
[/Darcy]
    type = NumbatDarcySF
    variable = streamfunction
    concentration_variable = concentration
[../]
[/TwoDConvectionDiffusion]
    type = NumbatConvectionDiffusionSF
    variable = concentration
    streamfunction_variable = streamfunction
[../]
[/TimeDerivative]
    type = TimeDerivative
    variable = concentration
[../]
[]

[AuxVariables]
[/u]
    order = CONSTANT
    family = MONOMIAL
[../]
[/w]
    order = CONSTANT
    family = MONOMIAL
[../]
[]

[AuxKernels]
[/uAux]
    type = NumbatDarcyVelocitySF
    variable = u
    component = x
    streamfunction_variable = streamfunction
[../]
[/wAux]
    type = NumbatDarcyVelocitySF
    variable = w
    component = y
    streamfunction_variable = streamfunction

```

```

[../]
[]

[BCs]
[./conctop]
    type = DirichletBC
    variable = concentration
    boundary = top
    value = 1.0
[../]
[./streamfuntop]
    type = DirichletBC
    variable = streamfunction
    boundary = top
    value = 0.0
[../]
[./streamfunbottom]
    type = DirichletBC
    variable = streamfunction
    boundary = bottom
    value = 0.0
[../]
[./Periodic]
    [./x]
        variable = 'concentration streamfunction'
        auto_direction = x
    [../]
[../]
[]

[Executioner]
    type = Transient
    dtmax = 100
    end_time = 5000
    start_time = 1
    solve_type = PJFNK
    nl_abs_tol = 1e-10
    [./TimeStepper]
        type = IterationAdaptiveDT
        dt = 1
        cutback_factor = 0.5
        growth_factor = 2
    [../]
    [./TimeIntegrator]
        type = LStableDirk2
    [../]
[]

```

```

[Postprocessors]
  [./boundaryfluxint]
    type = NumbatSideFlux
    variable = concentration
    boundary = top
  [../]
  [./numdofs]
    type = NumDOFs
  [../]
[]

[Preconditioning]
  [./smp]
    type = SMP
    full = true
    petsc_options = -snes_ksp_ew
    petsc_options_iname = '-pc_type -sub_pc_type -ksp_atol'
    petsc_options_value = 'asm lu 1e-12'
  [../]
[]

[Outputs]
  [./console]
    type = Console
    perf_log = true
    output_nonlinear = true
  [../]
  [./exodus]
    type = Exodus
    file_base = 2Dddc_isotropic
    execute_on = 'INITIAL TIMESTEP_END'
  [../]
  [./csvoutput]
    type = CSV
    file_base = 2Dddc_isotropic
    execute_on = 'INITIAL TIMESTEP_END'
  [../]
[]

```

5.1.2 Running the example

This example can be run on the commandline using

```
numbat-opt -i 2Dddc_isotropic.i
```

Alternatively, this file can be run using the *Peacock* gui provided by MOOSE using

```
peacock -i 2Dddc_isotropic.i
```


in the directory where the input file *2Dddc_isotropic.i* resides.

5.1.3 Results

This 2D example should take only a few minutes to run to completion, producing a concentration profile similar to that presented in Figure 5.1, where several downwelling plumes of high concentration can be observed after 3528 s:

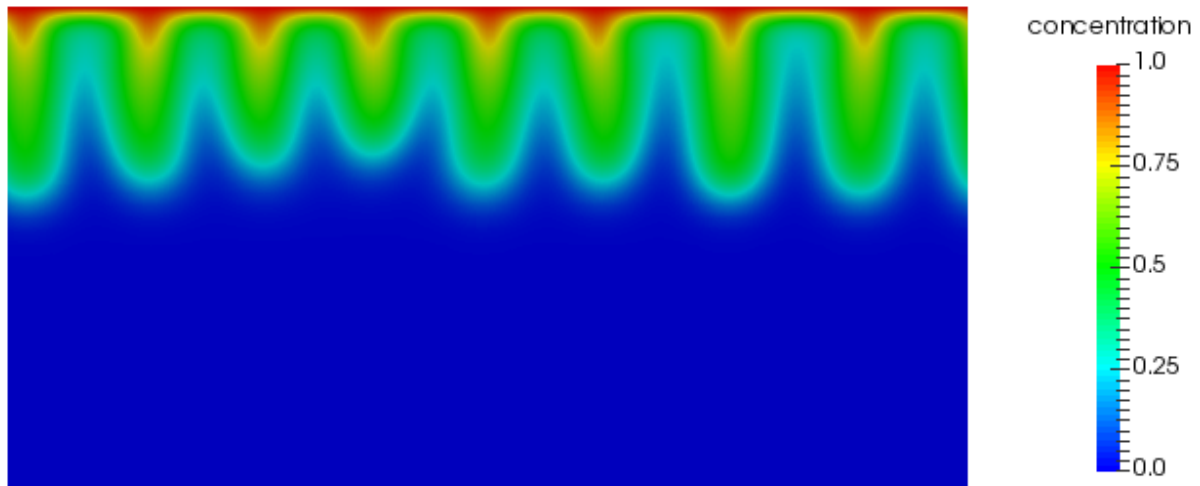


Figure 5.1: 2D concentration profile ($t = 3528$ s)

The flux per unit width over the top boundary is of particular interest in many cases (especially convective mixing of CO_2). This is calculated using the *boundaryfluxint* postprocessor in the input file, and presented in Figure 5.2.

Initially, the flux is purely diffusive, and scales as $1/\sqrt{(\pi t)}$, where t is time (shown as the dashed red line). After some time, the convective instability becomes sufficiently strong, at which point the flux across the top boundary rapidly increases (at a time of approximately 2000 seconds).

5.2 Anisotropic models

The second 2D example is for an anisotropic porous medium with $\gamma = 0.75$ (ie., the vertical permeability is three quarters of the horizontal permeability). A working example of this example can be found at https://github.com/cpgr/numbat/blob/master/examples/2D/anisotropic/2Dddc_anisotropic.i.

5.2.1 Input file

The complete input file for this anisotropic example is

```
[Mesh]
  type = GeneratedMesh
  dim = 2
  xmax = 1000
  ymin = -500
  ymax = 0
  nx = 80
  ny = 25
  bias_y = 0.9
```

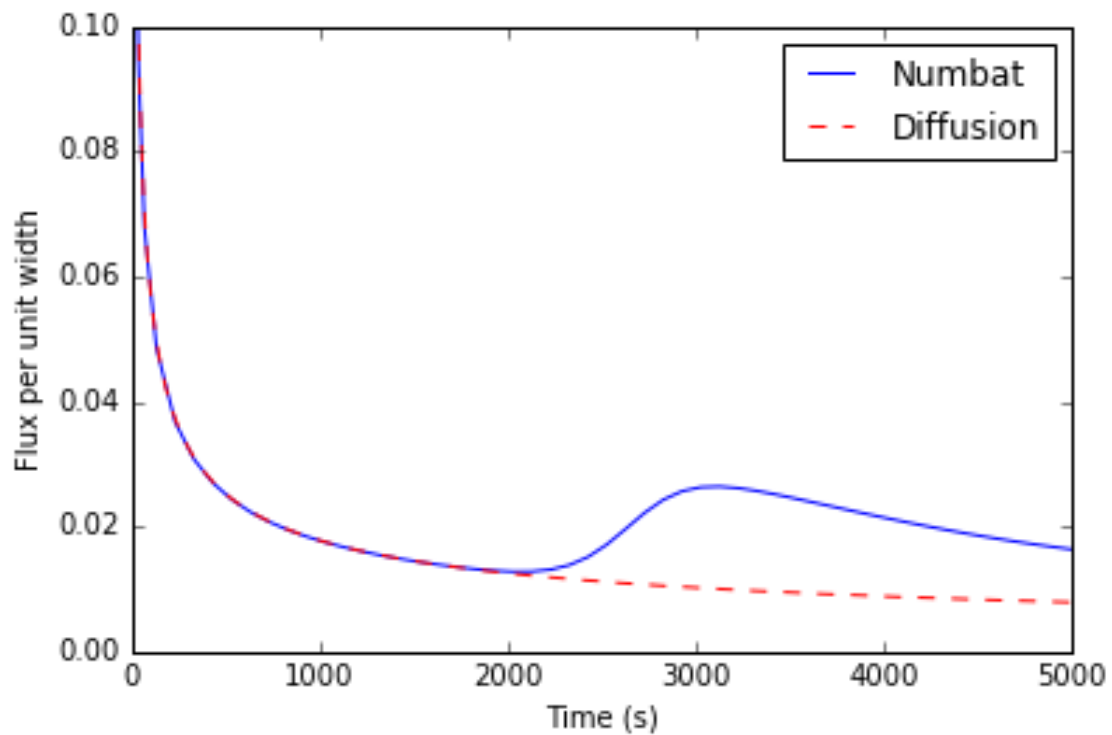


Figure 5.2: 2D flux across the top boundary

```
[
[Adaptivity]
  marker = errormarker
  max_h_level = 1
[./Indicators]
  [./gradjumpindicator]
    type = GradientJumpIndicator
    variable = concentration
  [./]
[./]
[./Markers]
  [./errormarker]
    type = ErrorToleranceMarker
    refine = 0.005
    indicator = gradjumpindicator
  [./]
[./]
[
[Variables]
  [./concentration]
    order = FIRST
```

```

family = LAGRANGE
[./InitialCondition]
    type = NumbatPerturbationIC
    variable = concentration
    amplitude = 0.1
    seed = 1
[../]
[../]
[./streamfunction]
    order = FIRST
    family = LAGRANGE
    initial_condition = 0.0
[../]
[]

[Kernels]
[./Darcy]
    type = NumbatDarcySF
    variable = streamfunction
    concentration_variable = concentration
    gamma = 0.75
[../]
[./ConvectionDiffusion]
    type = NumbatConvectionDiffusionSF
    variable = concentration
    streamfunction_variable = streamfunction
    anisotropic_tensor = '0.75 0 0 0 1 0 0 0 1'
[../]
[./TimeDerivative]
    type = TimeDerivative
    variable = concentration
[../]
[]

[AuxVariables]
[./u]
    order = CONSTANT
    family = MONOMIAL
[../]
[./w]
    order = CONSTANT
    family = MONOMIAL
[../]
[]

[AuxKernels]
[./uAux]
    type = NumbatDarcyVelocitySF
    variable = u

```

```

        component = x
        streamfunction_variable = streamfunction
    [../]
    [./wAux]
        type = NumbatDarcyVelocitySF
        variable = w
        component = y
        streamfunction_variable = streamfunction
    [../]
[]

[BCs]
    [./conctop]
        type = DirichletBC
        variable = concentration
        boundary = top
        value = 1.0
    [../]
    [./streamfuntop]
        type = DirichletBC
        variable = streamfunction
        boundary = top
        value = 0.0
    [../]
    [./streamfunbottom]
        type = DirichletBC
        variable = streamfunction
        boundary = bottom
        value = 0.0
    [../]
    [./Periodic]
        [./x]
            variable = 'concentration streamfunction'
            auto_direction = x
        [../]
    [../]
[]

[Executioner]
    type = Transient
    dtmax = 100
    end_time = 5000
    start_time = 1
    solve_type = PJFNK
    nl_abs_tol = 1e-10
    [./TimeStepper]
        type = IterationAdaptiveDT
        dt = 1
        cutback_factor = 0.5

```

```

    growth_factor = 2
[../]
[./TimeIntegrator]
    type = LStableDirk2
[../]
[]

[Postprocessors]
[./boundaryfluxint]
    type = NumbatSideFlux
    variable = concentration
    boundary = top
[../]
[./numdofs]
    type = NumDOFs
[../]
[]

[Preconditioning]
[./smp]
    type = SMP
    full = true
    petsc_options = -snes_ksp_ew
    petsc_options_iname = '-pc_type -sub_pc_type -ksp_atol'
    petsc_options_value = 'asm lu 1e-12'
[../]
[]

[Outputs]
[./console]
    type = Console
    perf_log = true
    output_nonlinear = true
[../]
[./exodus]
    type = Exodus
    file_base = 2Dddc_anisotropic
    execute_on = 'INITIAL TIMESTEP_END'
[../]
[./csvoutput]
    type = CSV
    file_base = 2Dddc_anisotropic
    execute_on = 'INITIAL TIMESTEP_END'
[../]
[]

```

Note that the permeability anisotropy is introduced in the kernels using the *gamma* and *anisotropic_tensor* input parameters.

5.2.2 Running the example

This example can be run on the commandline using

```
numbat-opt -i 2Dddc_anisotropic.i
```

Alternatively, this file can be run using the *Peacock* gui provided by MOOSE using

```
peacock -i 2Dddc_anisotropic.i
```

in the directory where the input file *2Dddc_anisotropic.i* resides.

5.2.3 Results

This 2D example should take only a few minutes to run to completion, producing a concentration profile similar to that presented in Figure 5.3, where several downwelling plumes of high concentration can be observed after 5000 s:

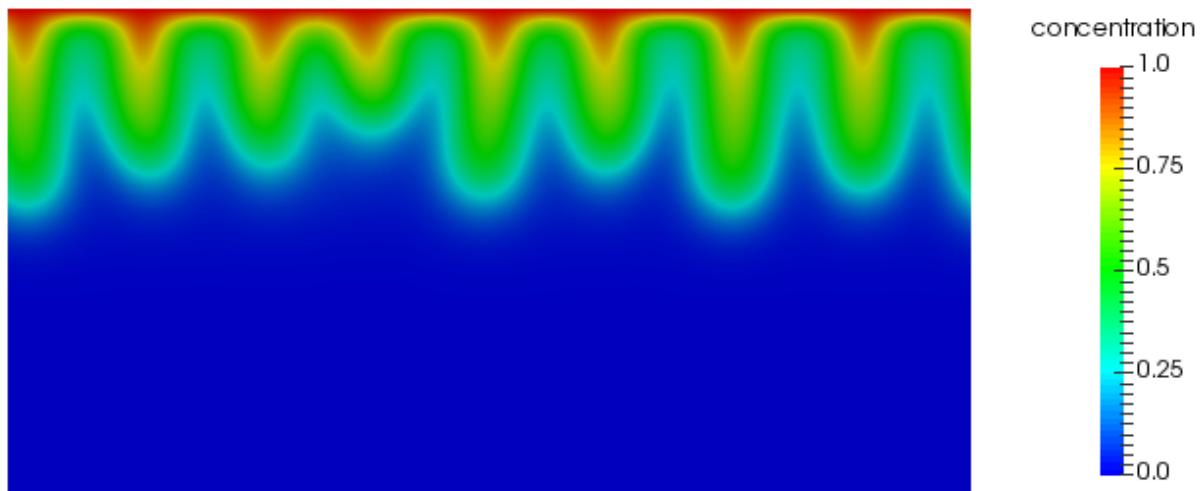


Figure 5.3: 2D concentration profile for $\gamma = 0.75$ ($t = 5000$ s)

In comparison to the isotropic example (with $\gamma = 1$) presented in Figure 5.1, we note that the concentration profile in the anisotropic example has only reached a similar depth after 5000 s (compared to 3528 s). The effect of the reduced vertical permeability in the anisotropic example slows the convective transport.

This observation can be quantified by comparing the flux per unit width over the top boundary of both examples, see Figure 5.4.

The inclusion of permeability anisotropy delays the onset of convection in comparison to the isotropic example, from a time of approximately 2000 seconds in the isotropic example to approximately 3500 seconds in the anisotropic example.

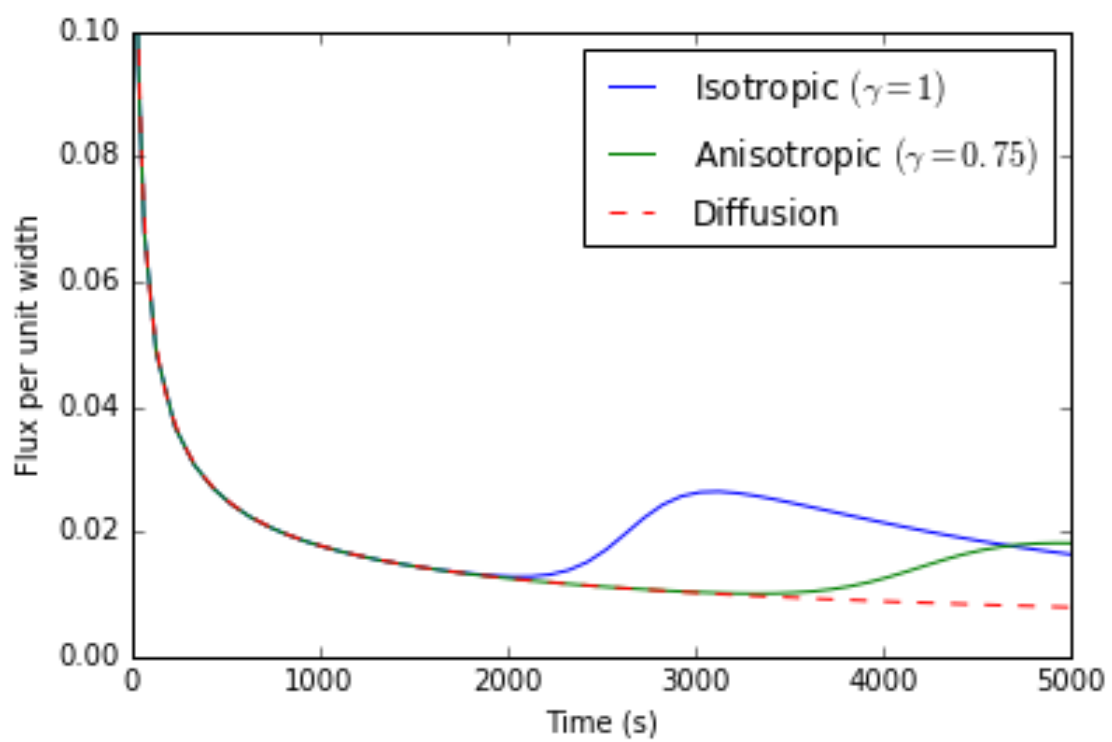


Figure 5.4: Comparison of the 2D flux across the top boundary

6 3D example

A working example of a 3D problem can be found at <https://github.com/cpgr/numbat/blob/master/examples/3D/isotropic/3Dddc.i>.

6.0.1 Input file

The complete input file for this problem is

```
[Mesh]
  type = GeneratedMesh
  dim = 3
  xmax = 500
  ymax = 500
  zmin = -200
  zmax = 0
  nx = 40
  ny = 40
  nz = 20
  bias_z = 0.7
[]

[Adaptivity]
  max_h_level = 1
  initial_marker = boxmarker
  initial_steps = 1
  marker = combomarker
  [./Indicators]
    [./gradjumpindicator]
      type = GradientJumpIndicator
      variable = concentration
    [../]
  [../]
  [./Markers]
    [./errormarker]
      type = ErrorToleranceMarker
      refine = 0.05
      indicator = gradjumpindicator
    [../]
    [./boxmarker]
      type = BoxMarker
      bottom_left = '0 0 -10'
      top_right = '500 500 0'
      inside = refine
      outside = dont_mark
    [../]
    [./combomarker]
      type = ComboMarker
      markers = 'boxmarker errormarker'
    [../]
  [../]
```



```
[]
```

```
[Variables]
```

```
  [./concentration]
    order = FIRST
    family = LAGRANGE
    [./InitialCondition]
      type = NumbatPerturbationIC
      variable = concentration
      amplitude = 0.02
      seed = 1
```

```
  [../]
```

```
[../]
```

```
  [./streamfunctionx]
    order = FIRST
    family = LAGRANGE
    initial_condition = 0.0
```

```
  [../]
```

```
  [./streamfunctiony]
    order = FIRST
    family = LAGRANGE
    initial_condition = 0.0
```

```
  [../]
```

```
[]
```

```
[Kernels]
```

```
  [./Darcy_x]
    type = NumbatDarcySF
    variable = streamfunctionx
    concentration_variable = concentration
    component = x
```

```
  [../]
```

```
  [./Darcy_y]
    type = NumbatDarcySF
    variable = streamfunctiony
    concentration_variable = concentration
    component = y
```

```
  [../]
```

```
  [./ConvectionDiffusion]
    type = NumbatConvectionDiffusionSF
    variable = concentration
    streamfunction_variable = 'streamfunctionx streamfunctiony'
    coeff_tensor = '1 0 0 0 1 0 0 0 1'
```

```
  [../]
```

```
  [./TimeDerivative]
    type = TimeDerivative
    variable = concentration
```

```
  [../]
```

```

[]

[AuxVariables]
  [./u]
    order = CONSTANT
    family = MONOMIAL
  [../]
  [./v]
    order = CONSTANT
    family = MONOMIAL
  [../]
  [./w]
    order = CONSTANT
    family = MONOMIAL
  [../]
[]

[AuxKernels]
  [./uAux]
    type = NumbatDarcyVelocitySF
    variable = u
    component = x
    streamfunction_variable = 'streamfunctionx streamfunctiony'
  [../]
  [./vAux]
    type = NumbatDarcyVelocitySF
    variable = v
    component = y
    streamfunction_variable = 'streamfunctionx streamfunctiony'
  [../]
  [./wAux]
    type = NumbatDarcyVelocitySF
    variable = w
    component = z
    streamfunction_variable = 'streamfunctionx streamfunctiony'
  [../]
[]

[BCs]
  [./conctop]
    type = DirichletBC
    variable = concentration
    boundary = front
    value = 1.0
  [../]
  [./streamfunxtop]
    type = DirichletBC
    variable = streamfunctionx

```

```

    boundary = front
    value = 0.0
[../]
[./streamfunxbottom]
    type = DirichletBC
    variable = streamfunctionx
    boundary = back
    value = 0.0
[../]
[./streamfunytop]
    type = DirichletBC
    variable = streamfunctiony
    boundary = front
    value = 0.0
[../]
[./streamfunybottom]
    type = DirichletBC
    variable = streamfunctiony
    boundary = back
    value = 0.0
[../]
[./Periodic]
    [./xy]
        variable = 'concentration streamfunctionx streamfunctiony'
        auto_direction = 'x y'
    [../]
[../]
[]

[Preconditioning]
    [./smp]
        type = SMP
        full = true
    [../]
[]

[Executioner]
    type = Transient
    dtmax = 100
    end_time = 2500
    start_time = 1
    solve_type = PJFNK
    nl_abs_tol = 1e-10
    petsc_options = -snes_ksp_ew
    [./TimeStepper]
        type = IterationAdaptiveDT
        dt = 1
        cutback_factor = 0.5
        growth_factor = 2

```

```

[../]
[./TimeIntegrator]
    type = LStableDirk2
[../]
[]

[Postprocessors]
[./boundaryfluxint]
    type = NumbatSideFlux
    variable = concentration
    boundary = front
[../]
[./numdofs]
    type = NumDOFs
[../]
[]

[Outputs]
[./console]
    type = Console
    perf_log = true
    output_nonlinear = true
    output_linear = true
[../]
[./exodus]
    type = Exodus
    file_base = 3Dddc
    execute_on = 'INITIAL TIMESTEP_END'
[../]
[./csvoutput]
    type = CSV
    file_base = 3Dddc
    execute_on = 'INITIAL TIMESTEP_END'
[../]
[]

```

6.0.2 Running the example

Note: This example should **not** be run on a laptop or workstation due to the large computational requirements. Do **not** run this using the *Peacock* gui provided by MOOSE.

Examples of the total run times for this problem on a cluster are over 27 hours for a single processor down to only 30 minutes using 100 processors in parallel.

6.0.3 Results

This 3D example should produce a concentration profile similar to that presented in Figure 6.1, where several downwelling plumes of high concentration can be observed:

Note that due to the random perturbation applied to the initial concentration profile, the geometry of the concentration profile obtained will differ from run to run.

The flux over the top surface is of particular interest in many cases (especially convective mixing of

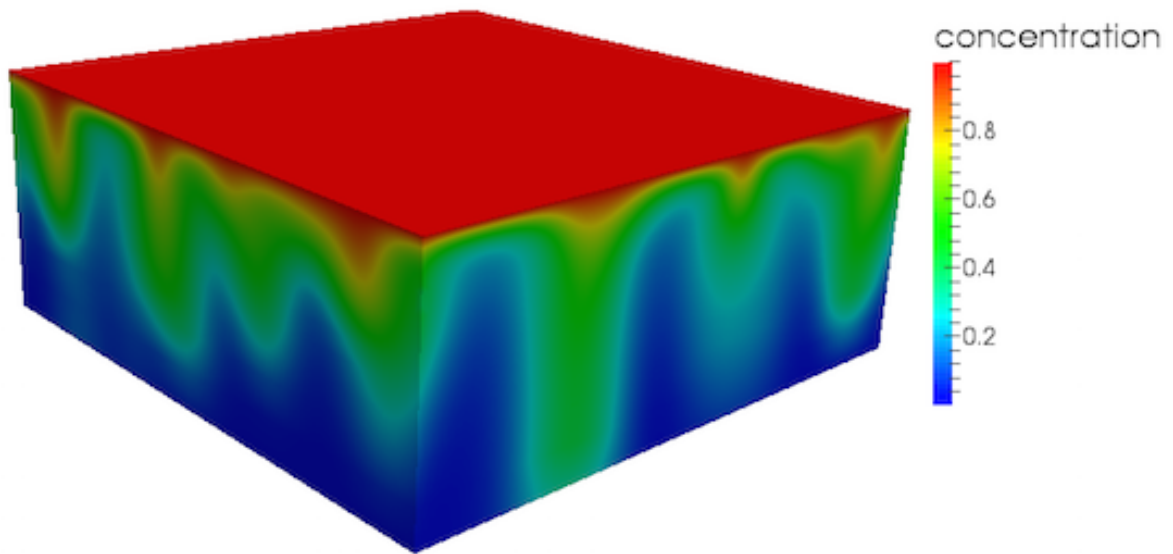


Figure 6.1: 3D concentration profile

CO₂). This is calculated in this example file using the *boundaryfluxint* postprocessor in the input file, and presented in Figure 6.2.

Initially, the flux is purely diffusive, and scales as $1/\sqrt{(\pi t)}$, where t is time (shown as the dashed green line). After some time, the convective instability becomes sufficiently strong, at which point the flux across the top boundary rapidly increases (at a time of approximately 1,700 seconds). Also shown for comparison is the flux for the 2D example. It is apparent that the 3D model leads in a slower onset of convection (the time where the flux first increases from the diffusive rate).

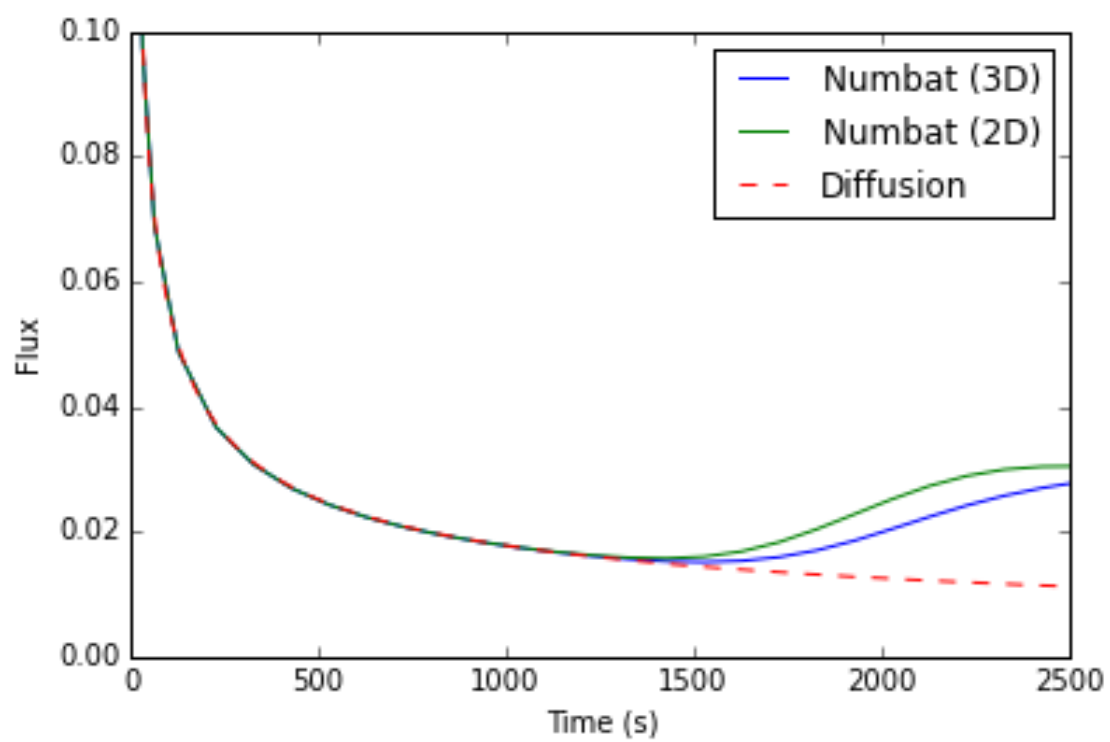


Figure 6.2: 3D flux across the top boundary

7 References

- E, W. and Liu, J. G., *Finite difference methods for 3D viscous incompressible flows in the vorticity-vector potential formulation on nonstaggered grids*, J. Comp. Phys., 138, 57–82 (1997)
- Ennis-King, J. and Paterson, L., *Role of convective mixing in the long-term storage of carbon dioxide in deep saline aquifers*, SPE J., 10, 349–356 (2005)
- Slim, A.C., *Solutal-convection regimes in a two-dimensional porous medium*, J. Fluid Mech., 741, 461–491 (2014)

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