# Transitioning from Visual Setup to text-based VSim simulations 



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## A brief introduction to me...

- Senior Research Scientist, 10.5 years at Tech-X
- Ph.D. @ Princeton/PPPL (2007), developing numerical methods for gyrokinetic PIC simulation
- Postdoc @ UW-Madison, working on RF/MHD coupling for electron cyclotron current drive in fusion plasmas
- Current research interests:
- methods for speeding up particle-in-cell simulations (SLPIC)
- modeling RF sheaths/impurity sputtering in fusion devices
- kinetic theory - wave/particle interactions, etc.
- PIC modeling of low-temperature plasmas
- Website, where this talk and many other talks/papers/presentations are posted:
http://nucleus.txcorp.com/~tgjenkins


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## This talk focuses on how VSim works 'under the hood'

- VSim's Visual Setup interface is designed to quickly bring new users up the VSim learning curve
- Allows common actions to be done quickly and systematically, with visual cues
-defining grids
-importing shapes
-applying boundary conditions
-adding particle species
- Shows users options consistent with their previous choices, while hiding others
-electrostatic vs. electromagnetic -direct vs. iterative matrix solve
- Visual Setup tools are adequate for many user needs (and we welcome suggestions for their improvement and development).
- Not everything that users want to do can be done in visual setup.
- Exercise experimental or developing code features
- Verify that the equations being solved are the ones the user intended
- ?


## My objective for this talk



The "New User" approach


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## My objective for this talk



- Understand how numerical discretization techniques are implemented in text input files
- Understand how to add to/edit text input files to get the result you want
- Some complexity unavoidable! But we'll look at things in stages, and periodically review and regroup, to make things easier.
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## Useful resources for this talk

- VSim online documentation:
https://www.txcorp.com/images/docs/vsim/latest/VSimDocumentation.html
- Slides for this talk:
http://nucleus.txcorp.com/~tgjenkins/pres/TWSSTalk2020.pdf
- Download page for the VSim input files I will use in this talk:
http://nucleus.txcorp.com/~tgjenkins/TWSS2020.html


## Choose a simple electrostatics problem: 1D Poisson

Physics problem

$$
\frac{d^{2} \phi(x)}{d x^{2}}=-\frac{\rho(x)}{\epsilon_{0}} ; \phi(x=0)=\phi^{\text {left }}, \quad \phi(x=L)=\phi^{\text {right }} ; x \in[0, L]
$$

## Numerical approach: discretize on a grid with $N$ cells.

$$
\text { Define the grid: } \quad \Delta x=\frac{L}{N} \quad ; \quad x_{n}=n \Delta x \quad \forall n=0,1, \ldots, N
$$

Use a finite-difference approximation to the second derivative, at interior gridpoints:

$$
-\epsilon_{0}\left[\frac{\phi_{j+1}-2 \phi_{j}+\phi_{j-1}}{\Delta x^{2}}\right]=\rho_{j} \quad \forall \quad j=1,2, \ldots, N-1
$$

Apply boundary conditions, at edge gridpoints:

$$
\begin{aligned}
\phi_{0} & =\phi^{l e f t} \\
\phi_{N} & =\phi^{r i g h t}
\end{aligned}
$$

## Solution error scales as $1 / N^{2}$

Physics problem:
exact
solution

$$
\frac{d^{2} \phi(x)}{d x^{2}}=-\frac{\rho_{0} \sin \left(\frac{\pi x}{L}\right)}{\epsilon_{0}} ; \quad \phi(x=0)=\phi^{\text {left }}, \quad \phi(x=L)=\phi^{\text {right }} \text { on }[0, L]
$$ has exact solution

$$
\phi(x)=\phi^{l e f t}+\left(\phi^{\text {right }}-\phi^{l e f t}\right) \frac{x}{L}+\frac{\rho_{0} L^{2}}{\epsilon_{0} \pi^{2}} \sin \left(\frac{\pi x}{L}\right)
$$

Numerical discretization: approximate solution

On the discrete grid, we have

$$
\phi_{j}^{e x a c t}=\phi^{l e f t}+\left(\phi^{\text {right }}-\phi^{l e f t}\right) \frac{j}{N}+\frac{\rho_{0} L^{2}}{\epsilon_{0} \pi^{2}} \sin \left(\frac{\pi j}{N}\right) ; \rho_{j}^{\text {exact }}=\rho_{0} \sin \left(\frac{\pi j}{N}\right)
$$

Putting these functions into the discretized Poisson equation yields

$$
\begin{aligned}
&-\frac{\rho_{0}}{\epsilon_{0}} \sin \left(\frac{\pi j}{N}\right)\left\{\frac{2 N^{2}}{\pi^{2}}\left[1-\cos \left(\frac{\pi}{N}\right)\right]\right\} \approx-\frac{\rho_{0}}{\epsilon_{0}} \sin \left(\frac{\pi j}{N}\right) \\
&\left\{\frac{2 N^{2}}{\pi^{2}}\left[1-\left(1-\frac{\pi^{2}}{2 N^{2}}+\frac{\pi^{4}}{24 N^{4}}+\cdots\right)\right]\right\} \approx 1
\end{aligned}
$$

## What does this look like in VSim?

Let's set up a basic simulation with Visual Setup and run it for one step:

Visual
Setup
input
file

Parameters (5)
VLEFT $=0$
VRIGHT $=1$
LX $=1$
NX $=10$
RHOZERO $=20$

Basic Settings (4)
number of steps = 1
steps between dumps $=1$
dimensionality $=1$
field solver $=$ electrostatic

```
SpaceTimeFunctions (1)
RHOxt=RHOZERO*sin(PI*x/LX)
```

```
Grids (3)
xMin =0
xMax = LX
xCells = NX
```

Field Dynamics: Fields (1)
Background Charge Density RHO=RHOxt
Field Dynamics: FieldBoundaryConditions (2) Dirichlet on lower x: VLEFT
Dirichlet on upper x: VRIGHT
Field Dynamics: PoissonSolver (2) preconditioner $=$ no preconditioner solver $=$ SuperLU

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# $\checkmark$ Sim generates .pre and in files from the Visual Setup .sdf file, when we Save and Setup 


.pre file - an intermediate object not of much immediate use to us (if generated by Visual Setup from a .sdf file)
.in file - the text input file we want to learn how to work with

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## Looking at vsim.in - input blocks

Frontmatter
<Grid globalGrid>
</Grid>
<Decomp decomp>
</Decomp>
<MultiField NAME_OF_MULTIFIELD>
<Field NAME_OF_FIELD>
</Field>
<FieldUpdater NAME_OF_FIELDUPDATER>
</FieldUpdater>
<InitialUpdateStep NAME_OF_INITIALUPDATESTEP>
</InitialUpdateStep>
<UpdateStep NAME_OF_UPDATESTEP>
</UpdateStep>
updateStepOrder $=$ [NAME_OF_UPDATESTEP_1 NAME_OF_UPDATESTEP2 ...] </MultiField>

Key VSim concept 0: block structures

Or very generally, <OBJECT objectName>
object features
</OBJECT>

## Looking at vsim.in - overall structure

Frontmatter
<Grid globalGrid>
</Grid>

## Key VSim concept 1: the MultiField block

<Decomp decomp>
</Decomp>
<MultiField NAME_OF_MULTIFIELD>

<FieldUpdater NAME_OF_FIELDUPDATER>
</FieldUpdater>
<InitialUpdateStep NAME_OF_INITIALUPDATESTEP>
</InitialUpdateStep>
<UpdateStep NAME_OF_UPDATESTEP> </UpdateStep>
define mathematical operations on field objects: e.g. taking the gradient of a scalar field and directing the output to a vector field
f define initial conditions - done only once at simulation outset
C Call the previously defined FieldUpdaters to manipulate the fields
updateStepOrder = [NAME_OF_UPDATESTEP_1 NAME_OF_UPDATESTEP2 ...] </MultiField>
in a specified ordered sequence of operations
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## Looking at vsim.in - Frontmatter, Grid block

| defines |
| :---: |
| some |
| global |
| simulation |
| parameters |

```
nsteps = 1 number of steps in simulation
dumpPeriodicity = 1 
dt = 1.0 < timestep (in units of seconds)
dimension = 1 4 1D simulation
floattype = double
verbosity = 127 How detailed the VSim output should
copyHistoryAtEachDump = 0
be (= 2}\mp@subsup{2}{}{M}-1) ; larger M = more detail
useGridBndryRestore = False
constructUniverse = False
```

defines
spatial grid properties
<Grid globalGrid>
verbosity $=127$
numCells $=\left[\begin{array}{lll}10 & 11 & 12\end{array}\right]$
lengths $=\left[\begin{array}{lll}1.0 & 1.0 & 1.0\end{array}\right]$
startPositions $=\left[\begin{array}{lll}0.0 & -0.5 & 0.0\end{array}\right]$
maxCellXings $=1$
</Grid>

3 D gid: default $\mathrm{y}, \mathrm{z}$ values
$\Delta x=1 / 10 ; \Delta y=1 / 11 ; \Delta z=1 / 12$ (extra $y, z$ dimensions are not used in this 1D computation, but may still be present in several parts of the input file)

## Looking at vsim.in - Field blocks

```
defines a
scalar or
vector field
to be used
    in the
simulation
```

<Field E>
numComponents \(=3\) offset \(=\) edge kind \(=\) regular overlap = [11] labels = \(\left[\begin{array}{lll}{\left[\begin{array}{l}x \\ E\end{array}\right]} & E_{-} z\end{array}\right]\) </Field>
<Field Phi> numComponents \(=1\)
offset \(=\) none kind = regular
\(\qquad\) lives on grid points overlap = [12] labels \(=[\) Phi]
</Field>
<Field ChargeDensity> numComponents =1 offset $=$ none $\longleftarrow$ lives on grid points kind $=$ depField $\longleftarrow$ special VSim field type, built from particle data overlap = [12] $\qquad$ labels = [ChargeDensity] </Field> nclude data from guard cells in a different way

## Looking at vsim.in - FieldUpdater blocks

defines a mathematical
operation on
Field objects
<FieldUpdater gradPhi> kind = gradVecUpdater factor $=-1.0$
lowerBounds = $\left[\begin{array}{lll}0 & 0 & 0\end{array}\right]$
upperBounds = [10 11
readFields $=[$ Phi]
writeFields = [E]
</FieldUpdater>
in other words,
$\vec{E}=-\vec{\nabla} \phi$.
inclusive)
exclusive scalar input, vector output for this FieldUpdater kind.

```
<FieldUpdater RHO> built-in operation that manipulates SpaceTimeFunction objects
    kind = STFuncUpdater
    operation = add < adds (subtracts, multiplies, etc.) the specified
    lowerBounds = [llll}000] (inclusive)
    upperBounds = [llllll}11212 13 [ (exclusive)
    writeFields = [ChargeDensity] scalar
    component = 0
    cellsToUpdateAboveDomain = [False False False]
    <STFunc f>
        kind = expression
```



```
        </STFunc>
```

    </FieldUpdater>
    
## Looking at vsim.in - InitialUpdateStep blocks

These updates are performed only once, at the simulation outset.

```
<InitialUpdateStep RHOInitStep>
    alsoAfterRestore = True Also do this step when restarting a simulation
    updaters = [RHO
                                4 Previously defined field updater, defines ChargeDensity field
    messageFields = []
    </InitialUpdateStep>
    <InitialUpdateStep esSolveInitStep>
    alsoAfterRestore = True
    updaters = [esSolve]
    messageFields = [Phi]
```



```
                            for phi field
</InitialUpdateStep>
<InitialUpdateStep gradPhilnitStep>
    alsoAfterRestore = True
    updaters = [gradPhi]
    Previously defined field updater, computes E from phi.
    messageFields = [E]
```

    </InitialUpdateStep>
    
## Looking at vsim.in - UpdateStep blocks

## Apply various FieldUpdater operations to <br> Field objects, <br> in a given <br> sequence

These updates are performed at every timestep in the simulation.
<UpdateStep RHOStep>

<UpdateStep gradPhiStep> toDtFrac $=1.0$ updaters = [gradPhi] messageFields = [E]
</UpdateStep>
...

Previously defined field updater, computes E from phi (just as in InitialUpdateStep call).

UpdateSteps can appear in the input file in any order you like, the updateStepOrder determines which ones will be called when.

Tech ${ }^{\text {undatestepOrder }=[\text { RHOStep esSolveStep gradPhistep }]}$ SIMULATIONS EMPOWERING INNOVATION

## Regroup and Review

So far, we have:
-built an .sdf file in VSim, using Visual Setup, that solves the 1D Poisson equation
-found the .in text input file that VSim built from our initial .sdf file
-looked at the general block structure of that in file

| -looked at some typical blocks that live in the larger MultiField block, and their contents |  |
| :--- | :--- |
| *Field | *FieldUpdater |
| *InitialUpdateStep | *UpdateStep |

Now, we'll do a bit of a deeper dive into how VSim solves the Poisson equation, and learn a bit more about how data is organized 'under the hood' in VSim.

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## Electrostatic solves, without VSim

Physics problem

VSim solves the Poisson equation

$$
\frac{d^{2} \phi(x)}{d x^{2}}=-\frac{\rho(x)}{\epsilon_{0}} ; \quad \phi(x=0)=\phi^{\text {left }}, \quad \phi(x=L)=\phi^{\text {right }} ; x \in[0, L]
$$

with Fields and FieldUpdaters and UpdateSteps.

Numerical
discretization

Let's build a discretized version of this problem "by hand", to see what kinds of things we might expect VSim to be doing:

N-cell grid:
Discrete Poisson equation:

Boundary
conditions:

$$
\Delta x=\frac{L}{N} \quad ; \quad x_{n}=n \Delta x \quad \forall n=0,1, \ldots, N
$$

$$
-\epsilon_{0}\left[\frac{\phi_{j+1}-2 \phi_{j}+\phi_{j-1}}{\Delta x^{2}}\right]=\rho_{j} \quad \forall \quad j=1,2, \ldots, N-1
$$

$$
\begin{aligned}
& \phi_{0}=\phi^{l e f t} \\
& \phi_{N}=\phi^{\text {right }}
\end{aligned}
$$

Result: a linear system of equations $\phi_{N}=\phi^{\text {right }} \quad$ for the unknown $\phi_{j}$ values.


## Constructing the matrix - interior points

$$
\begin{array}{r}
-\epsilon_{0}\left[\frac{\phi_{j+1}-2 \phi_{j}+\phi_{j-1}}{\Delta x^{2}}\right]=\rho_{j} \quad \forall j=1,2, \ldots, N-1 \\
\text { becomes a matrix of form }
\end{array}
$$

$$
\left(-\frac{\epsilon_{0}}{\Delta x^{2}}\right)\left[\begin{array}{ccccccccc}
1 & -2 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & -2 & 1 & 0 & 0 & 0 & 0 & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
0 & 0 & 1 & -2 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & -2 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & -2 & 1 & 0 & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
0 & 0 & 0 & 0 & 0 & 1 & -2 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & -2 & 1
\end{array}\right]\left[\begin{array}{c}
\phi_{0} \\
\phi_{1} \\
\phi_{2} \\
\vdots \\
\phi_{j-1} \\
\phi_{j} \\
\phi_{j+1} \\
\vdots \\
\phi_{N-2} \\
\phi_{N-1} \\
\phi_{N}
\end{array}\right]=\left[\begin{array}{c}
\rho_{0} \\
\rho_{1} \\
\rho_{2} \\
\vdots \\
\rho_{j-1} \\
\rho_{j} \\
\rho_{j+1} \\
\vdots \\
\rho_{N-2} \\
\rho_{N-1} \\
\rho_{N}
\end{array}\right]
$$

This doesn't work for the first/last rows of matrix. Instead, we must use boundary conditions there.

## Constructing the matrix - boundary conditions

$$
\begin{gathered}
\phi_{0}=\phi^{\text {left }} \\
\phi_{N}=\phi^{\text {right }} \\
\left(\frac{-\epsilon_{0}}{\Delta x^{2}}\right)\left[\begin{array}{ccccccccc}
-\gamma \Delta x^{2} / \epsilon_{0} & 0 & 0 & 0 & 0 & 0 & & 0 & 0 \\
1 & -2 & 1 & 0 & 0 & 0 & 0 \\
0 & & 1 & -2 & 1 & 0 & 0 & 0 & 0 \\
0 \\
\vdots & \vdots & \vdots & & \vdots & \vdots & \vdots & 0 & 0 \\
0 & \vdots & \vdots \\
0 & 0 & 1 & & -2 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & & 1 & -2 & 1 & 0 \\
0 & 0 & 0 & & 0 & 1 & -2 & & 0 \\
\vdots & 1 & 0 & 0 \\
\vdots & \vdots & \vdots & & \vdots & \vdots & \vdots & & \vdots \\
0 & 0 & 0 & & 0 & 0 & 1 & -2 & 1 \\
0 & & & 0 \\
0 & 0 & 0 & & 0 & 0 & 0 & 1 & -2 \\
0 & 0 & 0 & & 0 & 0 & 0 & 0 & 0 \\
\hline
\end{array}\right]\left[\begin{array}{c}
\phi_{0} \\
\phi_{1} \\
\phi_{2} \\
\vdots \\
\phi_{-1} \\
\phi_{j} \\
\phi_{j+1} \\
\vdots \\
\phi_{N-2} \\
\phi_{N-1} \\
\phi_{N}
\end{array}\right]=\left[\begin{array}{c}
\gamma \phi^{l e f t} \\
\rho_{1} \\
\rho_{2} \\
\vdots \\
\rho_{j-1} \\
\rho_{j} \\
\rho_{j+1} \\
\vdots \\
\rho_{N-2} \\
\rho_{N-1} \\
\mu \phi^{r i g h t}
\end{array}\right]
\end{gathered}
$$

Changes in the right-hand side vector (charge density) are necessary to implement the BCs.
Rescaling factors $\gamma, \mu$ can be used to adjust the matrix condition number.
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Canonical form: $\mathrm{Ax}=\mathrm{b}$.

# linearSolveUpdater - solving the Poisson equation 

Now let's look at how this is done in the vsim.in file.

One of VSim's built-in FieldUpdater blocks is the linearSolveUpdater, which solves equations of the form $A x=b$.

## Looking at vsim.in - linearSolveUpdater

## A FieldUpdater object <br> (mathematical <br> operation) that <br> solves a matrix <br> equation $A x=b$.

```
<FieldUpdater esSolve>
kind = linearSolveUpdater
lowerBounds = [0] (inclusive)
upperBounds = [11] (exclusive)
readFields = [ChargeDensity]
readComponents = [0]
writeFields = [Phi]
writeComponents = [0]
writeEquationToFile = 0
<MatrixFiller interiorFiller>
    kind = stFuncStencilFiller
    verbosity = 127
    minDim = 1
    lowerBounds = [[\begin{array}{lll}{1}&{1}&{1}\end{array}]\quad (inclusive)
    upperBounds = [lllll
    component = 0
<STFunc coeff>
    kind = expression
    expression = -8.854187817591624e-12}=-\mp@subsup{\epsilon}{0}{
</STFunc>
```

MatrixFiller blocks do just what they sound like - filling rows in the matrix.

## linearSolveUpdater - StencilElements

## Inside the MatrixFiller block, we have various StencilElements:

```
<STFuncStencilElement phi_dxp>
value = -100.0
value = -100.0 No offset
cellOffset = [00 0 0
functionOffset = [0.5 0.0.]
    rowFieldIndex = 0
    columnFieldIndex = 0
</STFuncStencilElement>
<STFuncStencilElement phi_npx>
    value = 100.0
    minDim = 1
+1 cell
    cellOffset = [llll
    functionOffset = [0.5 0. 0.]
    rowFieldIndex = 0
    columnFieldIndex = 0
    </STFuncStencilElement>
</STFuncStencilElement> diagonal matrix element
\(1 / \Delta x^{2}\)
```

$-1 / \Delta x^{2}$
$n, d=$ non-diagonal or
functionOffset is irrelevant for node-centered fields
<STFuncStencilElement phi_nmx>
value $=100.0$
minDim $=1$
-1 cell
celloffset $=\left[\begin{array}{lll}-1 & 0 & 0\end{array}\right]$
functionOffset $=\left[\begin{array}{lll}-0.5 & 0 . & 0 .\end{array}\right]$
rowFieldIndex $=0$
columnFieldIndex $=0$
</STFuncStencilElement>
columnFieldIndex $=0$
</STFuncStencilElement>
m, $\mathrm{p}=-\mathrm{/}+$
cell/function offset

```
<STFuncStencilElement phi_dxm>
```

<STFuncStencilElement phi_dxm>
value = -100.0 No offset
value = -100.0 No offset
value = -100.0 No offset
value = -100.0 No offset
cellOffset = [l0}0000
cellOffset = [l0}0000
    functionOffset = [-0.5 0. 0.]
    functionOffset = [-0.5 0. 0.]
    rowFieldIndex = 0
    rowFieldIndex = 0
    columnFieldIndex = 0
    columnFieldIndex = 0
</STFuncStencilElement>
</STFuncStencilElement>
value = -100.0 No offset

```
value = -100.0 No offset
```

A generic interior row in the 1D Poisson matrix is

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coefficient $\cdot\left[\begin{array}{lllllll}\cdots & 0 & p h i_{n m x} & \left(p h i_{d x m}+p h i_{d x p}\right) & p h i_{n p x} & 0 & \cdots\end{array}\right]$

## linearSolveUpdater - boundary conditions



## linearSolveUpdater - the linearSolver block

```
<LinearSolver linearSolver>
    kind = directSolver \ Solve Ax=b by computing A-1 directly.
    solverType = superLU Simplest VSim solver option (by the
    verbosity = 127 length-of-input-file metric, at least), but
</LinearSolver> not useful if your matrix is too large.
```

All other VSim solver types are iterative:

- generalized minimum residual
- conjugate gradient
- biconjugate gradient
- etc.

Iterative solvers can be sped up by appropriate multigrid preconditioners (for which many options are available in VSim).

## Let's look at the matrix VSim creates

- Edit the vsim.in file so that writeEquationToFile $=1$.
- NOTE: If you now hit the "Save" button, VSim Composer will
- re-read the vsim.sdf file, and
- generate a new .in file from the information it finds there.
- This will overwrite the change you just made, since the sdf file defaults to writeEquationToFile $=0$.
- Therefore: if you want to do text-based problem setup starting from a Visual Setup file, you'll need to generally do something like the following:
- Generate the initial .in file from the sdf file with the "Save" button
- Using your computer's file management utilities, copy the in file to a .pre file with a different prefix name, e.g. vsim.in becomes vsimTextBased.pre
- Edit this new .pre file in the way you want to
- Open the modified .pre file in VSim, and run VSim as normal (the visual setup utilities will no longer work, but the physics engine will still parse and run the input file that you've modified)


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## Assuming $A x=b, A$ is in esSolveMatrix.mtx

\%\%MatrixMarket matrix coordinate real general 111129
$111.7708375635183248 \mathrm{e}-09$ $21-8.8541900000000002 \mathrm{e}-10$ $221.7708380000000000 \mathrm{e}-09$ $23-8.8541900000000002 \mathrm{e}-10$ 3 2 -8.8541900000000002e-10 $331.7708380000000000 \mathrm{e}-09$ 3 4-8.8541900000000002e-10 4 3-8.8541900000000002e-10 $441.7708380000000000 \mathrm{e}-09$ $45-8.8541900000000002 \mathrm{e}-10$ $54-8.8541900000000002 \mathrm{e}-10$ 55 1.77083800000000000e-09 5 6-8.8541900000000002e-10 $65-8.8541900000000002 \mathrm{e}-10$ $661.7708380000000000 \mathrm{e}-09$ $67-8.8541900000000002 \mathrm{e}-10$ $76-8.8541900000000002 \mathrm{e}-10$ $771.7708380000000000 \mathrm{e}-09$ 78 -8.85419000000000002e-10 87-8.8541900000000002e-10 $881.7708380000000000 \mathrm{e}-09$ 89-8.8541900000000002e-10 $98-8.8541900000000002 \mathrm{e}-10$ $991.7708380000000000 \mathrm{e}-09$ $910-8.8541900000000002 \mathrm{e}-10$ 109 -8.85419000000000002e-10 $10101.7708380000000000 \mathrm{e}-09$ 1011 -8.8541900000000002e-10 1111 1.7708375635183248e-09
\%\%MatrixMarket matrix coordinate real genera
111129
11 2*eps0/dx^2
21 -eps0/dx^2
$222^{*} \mathrm{eps} 0 / \mathrm{dx} \mathrm{n}^{\wedge}$
23 -eps0/dx^2
32 -eps0/dx^2
33 2*eps0/dx^2
34 -eps0/dx^2
43 -eps0/dx^2
$442^{*} \mathrm{eps} 0 / \mathrm{dx}^{\wedge} 2$
45 -eps0/dx^2
54 -eps0/dx^2
$552^{*}$ epso/dx^2
56 -eps0/dx^2
$65-\mathrm{eps} 0 / \mathrm{dx}^{\wedge} 2$
$662^{*}$ eps0/dx^2
67 -eps0/dx^2
76 -eps0/dx^2
$772^{*} \mathrm{eps} 0 / \mathrm{dx} \mathrm{n}^{2}$
78 -eps0/dx^2
87 -eps0/dx^2
88 2*eps0/dx^2
89 -eps0/dx^2
98 -epso/dx^2
$992^{*}$ eps $0 / \mathrm{dx}^{\wedge} 2$
910 -eps $0 / d x^{\wedge} 2$
109 -eps0/dx^2
$10102^{*} \mathrm{eps} 0 / \mathrm{dx}^{\wedge} 2$
1011 -eps0/dx^2
$11112^{*} \mathrm{eps} 0 / \mathrm{dx}^{\wedge} 2$

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## Assuming $A x=b, x$ and $b$ are esSolve vectors

esSolveWriteVector.mtx (b)

\%\%MatrixMarket matrix array real general

111
$0.0000000000000000 \mathrm{e}+00$ $6.1803398874989481 e+00$ $1.1755705045849464 \mathrm{e}+01$ $1.6180339887498949 \mathrm{e}+01$ $1.9021130325903069 \mathrm{e}+01$ $2.0000000000000000 \mathrm{e}+01$ $1.9021130325903069 \mathrm{e}+01$ $1.6180339887498949 \mathrm{e}+01$ $1.1755705045849465 \mathrm{e}+01$ $6.1803398874989499 \mathrm{e}+00$ $1.7708375635183248 \mathrm{e}-09$

$$
\begin{aligned}
& =\frac{2 \epsilon_{0}}{\Delta x^{2}} \cdot \phi^{l e f t} \\
& =20 \sin \left(\frac{\pi x_{j}}{L}\right)=\rho_{j} \\
& =\frac{2 \epsilon_{0}}{\Delta x^{2}} \cdot \phi^{\text {right }}
\end{aligned}
$$

esSolveReadVector.mtx (x)
\%\%MatrixMarket matrix array real general
111
$0.0000000000000000 \mathrm{e}+00 \quad=\phi^{l e f t}$
$7.1308064483412903 e+10$
$1.3563599878269949 e+11$
$1.8668693648958243 e+11$
$2.1946365612852356 e+11$
$2.3075774401243900 e+11$
$2.1946365612872348 e+11$
$1.8668693648998233 \mathrm{e}+11$
$1.3563599878329944 \mathrm{e}+11$
$7.1308064484212875 e+10$
$1.0000000000000000 \mathrm{e}+00$
$=\phi^{r i g h t}$

## Regroup and Review

So far, we have:
-solved the discrete 1D Poisson equation 'by hand' and looked at the matrix and the vectors involved in that process
-looked at how VSim builds this matrix and these vectors with a FieldUpdater (of kind linearSolveUpdater), using MatrixFiller and StencilElement and LinearSolver blocks
-seen how to modify the .in file
-seen how to examine the matrix and vectors VSim builds.
But:
-most interesting problems are not 1D
-most interesting problems involve particles, complicated geometries, and/or complicated boundary conditions
Let's add some interesting features to our input file, and see how the .in file changes.

## Moving to 2D

Let's copy the simulation we had before into a new simulation, and add:

Parameters
LY = 1
$\mathrm{NY}=15$
RHOZERO $=2.0 \mathrm{e}-10$


SpaceTimeFunctions
RHOxt=RHOZERO*sin(PI*x/LX)*sin(PI*y/LY)
LINEARPHIxt=VLEFT+(VRIGHT-VLEFT)*x/LX
FieldBoundaryConditions
TOPBC: Dirichlet, LINEARPHIxt, upper y BOTTOMBC: Dirichlet, LINEARPHIxt, lower y

Basic Settings
dimensionality $=2$

Grid
$y \mathrm{Min}=0$
yMax=LY
yCells=NY
save as vsim2D.sdf

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## Matrix is larger, no longer tridiagonal

Now $176 \times 176[176=11 * 16=(N X+1) *(N Y+1)]$ and band-structured

$\rho$ and $\phi$ arrays are now representing 2D quantities in a vector, e.g.

$$
\left[\begin{array}{c}
\rho_{1,1} \\
\vdots \\
\rho_{1, N} \\
\rho_{2,1} \\
\vdots \\
\rho_{2, N} \\
\vdots \\
\rho_{M, N}
\end{array}\right]
$$

The same approach generalizes to 3D also; we will have large sparse matrices.
XTech-X
In general this 2D input file looks pretty similar to the 1D version. SIMULATIONS EMPOWERING INNOVATION

## Additional StencilElements relevant in 2D/3D

```
\Deltay
Typical stencil elements:
```



```
In 2D, general matrix row is coefficient \(\cdot\left[\begin{array}{lllllllllll}\cdots & 0 & p h i_{n m y} & \cdots & p h i_{n m x} & \left(p h i_{d x m}+p h i_{d x p}+p h i_{d y m}+p h i_{d y p}\right) & p h i_{n p x} & \cdots & p h i_{n p y} & 0 & \cdots\end{array}\right]\)

\section*{Adding GridBoundary geometric features}

Let's modify our simulation some more, to add geometric features:

\section*{Materials}

PEC: add to simulation

\section*{Geometries}

CSG: Add Primitive: cylinder material \(=\) PEC length \(=0.5\)
radius \(=0.1\)
x position \(=0.5\)
y position \(=0.5\)
\(z\) position \(=-0.25\)
axis direction \(\mathrm{x}=0.0\)
axis direction \(\mathrm{y}=0.0\)
axis direction \(z=1.0\)
save as vsim2Dcyl.sdf

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\section*{New: Material and GridBoundary blocks}
```

<EmMaterial PEC>
kind = conductor
resistance = 0.0
</EmMaterial>
<GridBoundary cylinder0>
    kind = gridRgnBndry
    calculateVolume = 1
    dmFrac = 0.5
    polyfilename = cylinder0.stl
    flipInterior = True
    scale = [1.0 1.0 1.0]
    printGridData = False
    mappedPolysfile = cylinder0_mapped.stl
</GridBoundary>
```

See documentation...
https://www.txcorp.com/images/docs/vsim/latest/VSimReferenceManual/vsimComposerMaterials.html and
https://www.txcorp.com/images/docs/vsim/latest/VSimReferenceManual/blocks_gridboundary.html

\section*{XTech-X}

\section*{New: GridBoundary MatrixFillers}
```

<MatrixFiller CYLINDERFiller>
kind = nodeStencilFiller
gridBoundary = cylinder0
rowInteriorosity = [cutByBoundary outsideBoundary]
colInteriorosity = [cutByBoundary outsideBoundary]
component = 0
minDim = 1
lowerBounds = [llll}
upperBounds = [llo 15 12]
<StencilElement ident>
    value = 5.7552220814345554e-09
    minDim = 1
    cellOffset = [ 0 0 0
    rowFieldIndex = 0
    columnFieldIndex = 0
    </StencilElement>
```
</MatrixFiller> See documentation..
As before, we could go and look at the matrix again, to see how these operations changed it, and get a sense for what VSim is doing behind-the-scenes.

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## Adding particles to an input file

- Instead of doing this through the visual setup, let's just open an example and test our developing .in-file-reading skills.
- File > New From Example > VSim for Plasma Discharges > Capacitively Coupled Plasma > Turner Case 2
- l'll show a quick movie of this discharge so that you have a sense for what we'll be looking at: available here:
http://nucleus.txcorp.com/~tgjenkins/movies/ShortCCPmovie.mov
- Neutral gas is contained between two parallel plates; one plate is grounded and the other biased with RF. The motion of free electrons creates plasma between the plates, and the formation of plasma sheaths is observed. The long-time steady state of the discharge is a balance between collisional ionization (source) and wall losses (sink).


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## Looking at the Turner .in file - ScalarDepositors

- Some familiar things: Fields, FieldUpdaters, UpdateSteps, MultiFields, etc.
- Some new things: ScalarDepositor, Species, Fluid, History, collisional physics, etc.

ScalarDepositors act like "buckets" that collect particle charge on the simulation grid.

```
<ScalarDepositor ChargeDensityDep>
    kind = areaWeighting « charge-collecting algorithm
    </\mp@code{lepField = esMultiField.ChargeDensity }
```

When all particles have been put into the bucket, its contents are then put into the specified depField.

## Looking at the Turner in file - Species

- Some familiar things: Fields, FieldUpdaters, UpdateSteps, MultiFields, etc.
- Some new things: ScalarDepositor, Species, Fluid, History, collisional physics, etc.

Species = almost everything having to do with the particle-in-cell aspects of VSim. Details beyond the scope of this already-long talk, but the principles are the same - nested block structures that describe objects and their interactions with other objects.

```
<Species electrons>
    kind = nonRelBoris
    charge = -1.6021766208e-19
    mass = 9.10938215e-31
    .••
    <ParticleSource particleLoaderE>
        <PositionGenerator posGen>
            ...
        </PositionGenerator>
        <VelocityGenerator velGen>
            ...
        </VelocityGenerator>
    </ParticleSource>
    <ParticleSink leftElecAbsorber>
    </ParticleSink>
</Species>
```

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## Looking at the Turner in file - History

- Some familiar things: Fields, FieldUpdaters, UpdateSteps, MultiFields, etc.
- Some new things: ScalarDepositor, Species, Fluid, History, collisional physics, etc.

History blocks create a record of various physics events on a per-timestep basis, rather than on a per-dump-step basis (e.g. current entering a wall, number of particles in the simulation, etc.)

```
<History numElec>
    kind = speciesNumberOf
                count the number of macroparticles in a species
    species = [electrons]
```

$\qquad$

``` specify the species
</History>
<History leftIonCurr>
```

```
                measure absorbed particle current from a species
    kind = speciesCurrAbs
    species = [He1] \longleftarrow
    ptclAbsorbers = [leftIonAbsorber] \longleftarrow}\mathrm{ specify the absorbing region
</History>
```

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## Exercises to test your text-file reading skills

- Look at the Turner.in file and see if you can identify how a species identifies the electric and magnetic fields that its particles respond to.
- Look in the VSim documentation at the different "kinds" of particle species (besides nonRelBoris) that are available. Could a nonRelES species work equally well for the Turner example?
- See if you can determine how particles are loaded in velocity space in the Turner.in input file. How would you add a mean flow to the particles?
- See if you can add a history block to Turner.in that records various properties (kinetic energy, velocity, loss time, etc.) of electrons that strike the left wall of the simulation and are lost.


## Useful resources: reminder

- VSim online documentation:
https://www.txcorp.com/images/docs/vsim/latest/VSimDocumentation.html
- Slides for this talk:
http://nucleus.txcorp.com/~tgjenkins/pres/TWSSTalk2020.pdf
- Download page for the VSim input files I used in this talk:
http://nucleus.txcorp.com/~tgjenkins/TWSS2020.html
- Also potentially of interest: Slides for other VSim talks I've given in the past (visualization, plasma sheath modeling, RF antenna simulations, CCPs, etc.):
http://nucleus.txcorp.com/~tgjenkins/informal.html


## Summary/Overview

- This talk was only a top-level view of the kinds of things you'll see if you edit text-based VSim input files... it's certainly possible to dig deeper.
- But if you're comfortable with the idea of block structures, and with digging into the documentation, I've hopefully given you enough information that you can start to tackle your own problems.
- Nevertheless, it's good to ask questions if you get stuck - please feel free to do so as you're figuring this stuff out. There are quite a few "power-users" of VSim who have probably had to wrestle with many of the problems you'll run into.
- Thanks for your attention!


## XTech-X

