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#!/bin/bash
# Developed by: Darren E. Holland, Bryan V. Egner, Julie V. Logan
# Modified by Darren Holland 2020-11-02
# This wrapper creates the geometry (CreateMat.py and CreateGeo.inp), compiles
# the Geant code to calculate the mass, runs Geant, and
# calls the analysis script (Analyze.py) which contains the surrogate model and
# evaluation.
# Inputs to ./SurrWrapper.sh $1 $2 $3
# $1 is the folder name
# $2 is the geometry increment number/identifier
# $3 is the results file to return to Dakota
#************************************
*****
# Settings
#***********************
******
#Job
threads=2
                    # Total number of processors
                    # Number of processors to use per group (On Bridgman use
numproc=2
16 aka all of a node)
# G4MPI (Geant parallel processing) directory
MPIDIR=/app/afit/geant4/geant4.10.03.p02-install-vis/G4MPI/lib64/G4mpi-10.3.2
# Output spectrum binning
nbins=664
                    # Number of spectrum bins
                         # Lowest bin energy (MeV)
lhins=0
# Output spectrum energy cut-offs for detector response curve creation (Use full
energy peak only)
ubins=0.663
                    # Highest bin energy (MeV)
# Source
                    # Start source at voxel edge for voxel at this angle
StartSource=1
(degrees)
                 # Example: 25 with deltaphi = 12 starts source at
int(25/12)=2 * 12=24 degrees
                    # Number of theta sampling subdivisions (Nyquist
s_subdiv=2
criterion)
s_dist=100.
                    # Source distance (cm)
SourceEnergyType=none
                         # Source energy spectrum, none = monoenergetic
source
SEnergy=0.662
                    # Source energy (MeV)
SPart=gamma
                    # Particle type (gamma or neutron)
# Geometry (note origin is at detector's geometric center):
StartGeo=0
                    # Phi position for starting geometry (voxel edge)
MaskMaterial=PMMA # Mask Material, Options: PMMA
MaskMinThick=0.81 # Mask's minimum thickness must be greater than zero to
connect mask elements (cm)
DH=1.27
                         # Detector HALF height (cm)
DR=1.27
                         # Detector radius (cm)
SOR=2.69875
                    # Sleeve outer radius (cm)
                    # Sleeve height above origin (cm)
SH=2.69875
SB=55
                    # Sleeve bottom (cm)
                    # (0) Use Tesselated Solid elements
RSMTet=0
# Code Directories
Dakotadir=$(pwd) # Dakota working directory
cd ../../
homedir=$(pwd)
                    # Location of Neutron Mask Optimization Code (aka this
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code)
Method="MOGA"
# The following variables are set in the Dakota call (to calculate number of
variables)
*****
# Set attenuation coefficients for PMMA surrogate model
if [[ $MaskMaterial == "PMMA" ]]
    maskdensity=1.19
    #see PMMALinAtten.pdf for NIST table 0.6 MeV 8.701E-02 cm^2/g
                                                                       0.8
7.641E-02
            so 0.662 0.08372
    LinAtten=0.09975 #/cm assuming 1.19 g/cc density
    DetAtten=0.2829 #/cm assuming 3.667 g/cc density
fi
#READ DAKOTA DISCRITIZATION
finthick=$(echo "$(grep ' finthick' "$Dakotadir/$1.MOGApara")" | grep -Eo "[-+]?
[0-9]*\.?[0-9]+([eE][-+]?[0-9]+)?.")
wallthick=$(echo "$(grep ' wallthick' "$Dakotadir/$1.MOGApara")" | grep -Eo "[-
+]?[0-9]*\.?[0-9]+([eE][-+]?[0-9]+)?.")
deltatheta=$(echo "$(grep ' deltatheta' "$Dakotadir/$1.MOGApara")" | grep -Eo
"[-+]?[0-9]*\.?[0-9]+([eE][-+]?[0-9]+)?.")
deltaphi=$(echo "$(grep 'deltaphi' "$Dakotadir/$1.MOGApara")" | grep -Eo "[-+]?
[0-9]*\.?[0-9]+([eE][-+]?[0-9]+)?.")
finwidth=$(echo "$(grep ' finwidth' "$Dakotadir/$1.MOGApara")" | grep -Eo "[-+]?
[0-9]*\.?[0-9]+([eE][-+]?[0-9]+)?.")
wallwidth=$(echo "$(grep ' wallwidth' "$Dakotadir/$1.MOGApara")" | grep -Eo "[-
+]?[0-9]*\.?[0-9]+([eE][-+]?[0-9]+)?.")
# Set phi limits for geometry (geostart to geofinal) and measurements (phistart
to phifinal)
phistart=$(awk -v dd=$StartSource -v ee=$deltaphi 'BEGIN {printf "%.4f\
n",int(dd/ee)*ee}')
geostart=$(awk -v dd=$StartGeo -v ee=$deltaphi 'BEGIN {printf
"%.4f\n",int(dd/ee)*ee}')
                             # Phi final limit (uncomment to set manually)
phifinal=155
geofinal=$(awk -v dd=$geostart -v ee=170 -v ff=$deltaphi 'BEGIN {printf "%.4f\
n",int((ee-dd)/ff)*ff+dd}')
#RSMmaxsize is depreciated for this code (set to one so no effect)
RSMmaxsize=1
                             # Maximum RSM radius (cm)
# Copy source code into project and working directories
filedir="$homedir/$1/GeantRuns"
workdir="$Dakotadir"
export PYTHONPATH=$PYTHONPATH:$filedir
mkdir "$workdir/bld"
cp "$homedir/VolCalc.cc" "$workdir/bld"
cp "$homedir/CMakeListsVolCalc.txt" "$workdir/bld"
if [[ ! -d "$filedir" ]]
then
      # Create working directory and copy files
    mkdir "$filedir"
    mkdir "$filedir/res"
      cp -R "$homedir/src/" "$filedir"
      cp -R "$homedir/include/" "$filedir"
      cp "$homedir/AnalyzeSurr.py" "$filedir"
      cp "$homedir/CreateGeo.py" "$filedir"
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cp "$homedir/CreateMat.pv" "$filedir"
     cp "$homedir/intersectLineCylinder.py" "$filedir"
cp "$homedir/intersectLinePlane.py" "$filedir"
cp "$homedir/linePosition3d.py" "$filedir"
      cp "$homedir/submitGeant.pbs" "$filedir"
fi
cp "$homedir/Settings.cc" "$workdir/bld/Settingstemp.txt"
cp "$homedir/Geoinp.py" "$workdir/bld/Geoinptemp.py"
# Set contant geometry inputs
sed -i -e "s?RSMmaxsize=20?RSMmaxsize=$RSMmaxsize?g"
"$workdir/bld/Geoinptemp.py"
sed -i -e "s?MaskMinThick=0.1?MaskMinThick=$MaskMinThick?g"
"$workdir/bld/Geoinptemp.py"
sed -i -e "s?sleeve_bottom=55?sleeve_bottom=$SB?g" "$workdir/bld/Geoinptemp.py"
sed -i -e "s?sleeve_height=5.0?sleeve_height=$SH?g"
"$workdir/bld/Geoinptemp.py"
sed -i -e "s?sleeve_outer_rad=4.1275?sleeve_outer_rad=$$OR?g"
"$workdir/bld/Geoinptemp.py"
sed -i -e "s?sleeve_inner_rad=3.81?sleeve_inner_rad=$DR?g"
"$workdir/bld/Geoinptemp.py"
sed -i -e "s?det_rad=3.81?det_rad=$DR?g" "$workdir/bld/Geoinptemp.py"
sed -i -e "s?det_height=7.62?det_height=$DH?g" "$workdir/bld/Geoinptemp.py"
sed -i -e "s?Settings.cc?$workdir/bld/Settingstemp.txt?q"
"$workdir/bld/Geoinptemp.py"
#*********************
* * * * * * * * * * * * * * *
# Implement Geant settings
*****
# Set spectrum binning
lbinsmev=$(awk -v dd=$lbins 'BEGIN {printf "%.4f\n",dd/1000}')
ubinsmev=$(awk -v dd=$ubins 'BEGIN {printf "%.4f\n",dd/1000}')
           "s?nofBins(10)?nofBins($nbins)?g" "$workdir/bld/Settingstemp.txt"
sed -i -e
            # Number of bins
# Source info
sed -i -e "s?AmBe.mac?$SourceEnergyType?g" "$workdir/bld/Settingstemp.txt"
              # Source spectrum
sed -i -e "s?SourceDiv(2)?SourceDiv($s_subdiv)?g"
"$workdir/bld/Settingstemp.txt"
                                         # Number of measurement substeps
sed -i -e "s?nParts2Run(50000)?nParts2Run($nParts2Run)?q"
                                          # Number of source particles
"$workdir/bld/Settingstemp.txt"
sed -i -e "s?gamma?$SPart?g" "$workdir/bld/Settingstemp.txt"
                    # Source particle
sed -i -e "s?SourceDist(86.36)?SourceDist($s_dist)?g"
"$workdir/bld/Settingstemp.txt"
                                   # Source distance (cm)
sed -i -e "s?numEnergies(1)?numEnergies(1)?g" "$workdir/bld/Settingstemp.txt"
                  # Number of source energies (currently commented out)
sed -i -e "s?energiesMeV(0.662)?energiesMeV($SEnergy)?g"
"$workdir/bld/Settingstemp.txt"
                                  # Source particle energy
# Geometry discretization
sed -i -e "s?deltatheta(10)?deltatheta($deltatheta)?g"
"$workdir/bld/Settingstemp.txt"
                                 # Theta increments
sed -i -e "s?deltaphi(10)?deltaphi($deltaphi)?g"
"$workdir/bld/Settingstemp.txt"
                                          # Phi increments
sed -i -e "s?phifinal(170)?phifinal($phifinal)?g"
"$workdir/bld/Settingstemp.txt"
                                          # Final phi position
# Geometry (in cm)
sed -i -e "s?RSMTet(0)?RSMTet($RSMTet)?g" "$workdir/bld/Settingstemp.txt"
                # TesselatedSolids (newer designs)
sed -i -e "s?DetRad(3.81)?DetRad($DR)?g" "$workdir/bld/Settingstemp.txt"
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# Detector radius
           "s?DetHeight(3.81)?DetHeight($DH)?g" "$workdir/bld/Settingstemp.txt"
sed -i -e
                  # Detector HALF height
sed -i -e "s?SleeveOuterRad(4.1275)?SleeveOuterRad($SOR)?g"
"$workdir/bld/Settingstemp.txt"
                                   # Sleeve radius
sed -i -e "s?SleeveHeight(5)?SleeveHeight($SH)?g"
"$workdir/bld/Settingstemp.txt"
                                          # Top of sleeve (extends past
detector)
sed -i -e "s?SleeveBottom(55)?SleeveBottom($SB)?g"
"$workdir/bld/Settingstemp.txt"
                                          # Total length of sleeve
# Set design number
cd "$workdir/bld"
if [[ $2 -lt 10 ]]
then
      geonum=""0000$2""
fi
if [[ $2 -lt 100 ]] && [[ $2 -ge 10 ]]
then
     geonum=""000$2""
fi
if [[ $2 -lt 1000 ]] && [[ $2 -ge 100 ]]
then
     geonum=""00$2""
fi
if [[ $2 -lt 10000 ]] && [[ $2 -ge 1000 ]]
then
     geonum=""0$2""
fi
if [[ $2 -ge 10000 ]]
then
     geonum=""$2""
# Create Geometry
# Overwrite template values and filenames
cp "$workdir/bld/Geoinptemp.py" "$workdir/bld/Geoinp.py"
sed -i -e "s?EigMat.txt?DesignMat$Method$2.txt?g" "$workdir/bld/Geoinp.py"
Geofile="$workdir/bld/DesignMat$Method$2.txt"
Geofile_repeat="../"$Method"bld."$2/bld/DesignMat$Method$2.txt
inodes="$workdir/"$1"Nodes"$Method$geonum".inp"
ielements="$workdir/"$1"Ele"$Method$geonum".inp"
sed -i -e "s?s_dist=86.36?s_dist=$s_dist?g" "$workdir/bld/Geoinp.py"
# Node and element filenames
sed -i -e "s?Nodesfile='/nodes.inp'?Nodesfile='$inodes'?g"
"$workdir/bld/Geoinp.py"
sed -i -e "s?Elemfile='/elem.inp'?Elemfile='$ielements'?g"
"$workdir/bld/Geoinp.py"
# Create nodes and elements
Bad_Geo=$(python3 $workdir/bld/Geoinp.py $finthick $wallthick $deltatheta
$deltaphi $finwidth $wallwidth $geostart $geofinal $DR $DH)
# If invalid geometry skip evaluation
if [[ $Bad_Geo == 0 ]]
then
    echo "Bad_Geo 0" > $Dakotadir/$3
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# Calculate variance reduction using cone angle created in geometry file
     coneangle=$(grep 'coneangle(' "$workdir/bld/Settingstemp.txt" | cut -c
32-)
     coneangle="${coneangle%)*}"
     VRval=$(awk -v dd=$coneangle 'BEGIN {printf "%.6e\n",(1 - cos(dd*atan2(0,-
1)/180)) / 2} )
     # Read nodes and elements into Geant
     cp "$workdir/bld/Settingstemp.txt" "$workdir/bld/Settings.cc"
     sed -i -e "s?RSM_nodes.inp?$inodes?g" "$workdir/bld/Settings.cc"
     # Nodes
     sed -i -e "s?RSM_elements.inp?$ielements?g" "$workdir/bld/Settings.cc"
     # Elements
     sed -i -e "s?Ofile?"$1$Method$geonum"?q" "$workdir/bld/Settings.cc"
Geometry output file
     # Create copy of settings file for records
     cp "$workdir/bld/Settings.cc" "$workdir/bld/Settings"$Method$geonum".txt"
     # Calculate the design's mass using Geant
       cp "$homedir/VolCalc.cc" "$workdir/bld"
     cp "$homedir/CMakeListsVolCalc.txt" "$workdir/bld/CMakeLists.txt"
     cmake -DFILE_DIR=$filedir -DExeName=VolCalc -DExeName2="VolCalc$geonum" -
DSettings=Settings .
       make
     $workdir/bld/VolCalc$geonum
#******************************
       # Analyze results and return to Dakota
#*********************************
     # Move files and clean up directory
     mkdir $filedir/res/$Method$geonum
     find "$workdir/bld/" -maxdepth 1 -name "WallPos.inp" -exec mv -t
"$filedir/res/$Method$geonum" {} +
     find "$workdir/bld/" -maxdepth 1 -name "FinPos.inp" -exec mv -t "$filedir/
res/$Method$geonum" {} +
     find "$workdir/bld/" -maxdepth 1 -name "x_c.txt" -exec mv -t
"$filedir/res/$Method$geonum" {} +
     find "$workdir/bld/" -maxdepth 1 -name "y_c.txt" -exec mv -t
"$filedir/res/$Method$geonum" {} +
     find "$workdir/bld/" -maxdepth 1 -name "z_c.txt" -exec mv -t
"$filedir/res/$Method$geonum" {} +
     # Analyze the results
     echo "python3 ./AnalyzeSurr.py ./$3 ./res/$Method$geonum/Results $SPart
$s_subdiv $deltaphi $phifinal $deltatheta ./res/$Method$geonum $VRval
$maskdensity $phistart $Geofile_repeat $LinAtten 1 $wallwidth $finwidth
$geostart $geofinal $DR $DH $wallthick $finthick $DetAtten" >
$filedir/AnalyzeCommand$Method$geonum.sh
       chmod u+x $filedir/AnalyzeCommand$Method$geonum.sh
     python3 $filedir/AnalyzeSurr.py $Dakotadir/$3 $filedir/res/$Method$geonum/
Results $SPart $s_subdiv $deltaphi $phifinal $deltatheta
$filedir/res/$Method$geonum $VRval $maskdensity $phistart $Geofile $LinAtten 0
$wallwidth $finwidth $geostart $geofinal $DR $DH $wallthick $finthick $DetAtten
else
  #echo "Skipping invalid geometry..."
  echo "Bad_Geo "$Bad_Geo >> $Dakotadir/$3
fi
echo $Dakotadir/$3
# Change to initial run directory
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cd \$Dakotadir