3D06_sphere_inflation_v0p8_A.Flowers_Comments

July 7, 2025

1 Inflation of a sphere by an internal pressure

1.0.1 Units

Length: mmMass: kg

• Time: s

• Force: milliNewtons

• Stress: kPa

1.0.2 Software:

• Dolfinx v0.8.0

In the collection "Example Codes for Coupled Theories in Solid Mechanics,"

By Eric M. Stewart, Shawn A. Chester, and Lallit Anand.

https://solidmechanicscoupledtheories.github.io/

2 Import modules

```
[1]: # Import FEnicSx/dolfinx
import dolfinx

# For numerical arrays
import numpy as np

# For MPI-based parallelization
from mpi4py import MPI
comm = MPI.COMM_WORLD
rank = comm.Get_rank()

# PETSc solvers
from petsc4py import PETSc

# specific functions from dolfinx modules
from dolfinx import fem, mesh, io, plot, log
```

```
from dolfinx.fem import (Constant, dirichletbc, Function, functionspace, u
 from dolfinx.fem.petsc import NonlinearProblem
from dolfinx.nls.petsc import NewtonSolver
from dolfinx.io import VTXWriter, XDMFFile
# specific functions from ufl modules
import ufl
from ufl import (TestFunctions, TrialFunction, Identity, grad, det, div, dev, u
→inv, tr, sqrt, conditional ,\
                gt, dx, inner, derivative, dot, ln, split)
# basix finite elements (necessary for dolfinx v0.8.0)
import basix
from basix.ufl import element, mixed_element, quadrature_element
# Matplotlib for plotting
import matplotlib.pyplot as plt
plt.close('all')
# For timing the code
from datetime import datetime
# Set level of detail for log messages (integer)
# Guide:
# CRITICAL = 50, // errors that may lead to data corruption
# ERROR = 40, // things that HAVE gone wrong
# WARNING = 30, // things that MAY go wrong later
# INFO = 20, // information of general interest (includes solver info)
# PROGRESS = 16, // what's happening (broadly)
# TRACE = 13, // what's happening (in detail)
# DBG
          = 10 // sundry
log.set_log_level(log.LogLevel.WARNING)
```

3 Define geometry

```
DEFINE GEOMETRY

# Thick-walled sphere dimensions

Ro = 11 # mm outer radius

Ri = 10 # mm inner radius
```

```
# Pull in the mesh *.xdmf file and read any named domains in the mesh.
with XDMFFile(MPI.COMM_WORLD, "meshes/spherical_shell.xdmf", 'r') as infile:
    domain = infile.read_mesh(name="Grid", xpath="/Xdmf/Domain")
    cell_tags = infile.read_meshtags(domain,name="Grid")

# Create facet to cell connectivity required to determine boundary facets.
domain.topology.create_connectivity(domain.topology.dim, domain.topology.dim-1)

# Read in facet tags from an *.xdmf file.
with XDMFFile(MPI.COMM_WORLD, "meshes/facet_spherical_shell.xdmf", "r") as xdmf:
    facet_tags = xdmf.read_meshtags(domain, name="Grid")

x = ufl.SpatialCoordinate(domain)
```

Print out the unique facet index numbers

```
[3]: top_imap = domain.topology.index_map(2)  # index map of 2D entities in_u domain (facets)

values = np.zeros(top_imap.size_global)  # an array of zeros of the same_u size as number of 2D entities

values[facet_tags.indices]=facet_tags.values # populating the array with facet_u tag index numbers

print(np.unique(facet_tags.values))  # printing the unique indices

# Surface numbering from gmsh:
# Physical Surface("yBot", 12)
# Physical Surface("zBot", 13)
# Physical Surface("xBot", 14)
# Physical Surface("inner_surf", 15)
```

[12 13 14 15]

Visualize reference configuration and boundary facets

```
[4]: import pyvista
    pyvista.set_jupyter_backend('html')
    from dolfinx.plot import vtk_mesh
    pyvista.start_xvfb()

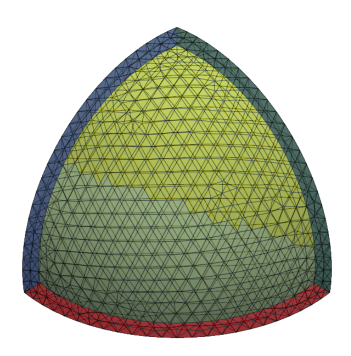
# initialize a plotter
plotter = pyvista.Plotter()

# Add the mesh -- I make the 3D mesh opaque, so that 2D surfaces stand out.
topology, cell_types, geometry = plot.vtk_mesh(domain, domain.topology.dim)
grid = pyvista.UnstructuredGrid(topology, cell_types, geometry)
plotter.add_mesh(grid, show_edges=True, opacity=0.5)
```

```
# Add colored 2D surfaces for the named surfaces
yBot = pyvista.UnstructuredGrid(*vtk_mesh(domain, domain.topology.
 →dim-1,facet_tags.indices[facet_tags.values==12]) )
zBot = pyvista.UnstructuredGrid(*vtk mesh(domain, domain.topology.

→dim-1,facet_tags.indices[facet_tags.values==13]) )
xBot = pyvista.UnstructuredGrid(*vtk mesh(domain, domain.topology.
dim-1,facet_tags.indices[facet_tags.values==14]) )
inner surf = pyvista.UnstructuredGrid(*vtk mesh(domain, domain.topology.
 dim-1,facet_tags.indices[facet_tags.values==15]) )
actor = plotter.add_mesh(yBot, show_edges=True,color="blue") # yBot face is blue
actor2 = plotter.add_mesh(zBot, show_edges=True,color="red") # zBot is red
actor3 = plotter.add_mesh(xBot, show_edges=True,color="green") # xBot is green
actor4 = plotter.add_mesh(inner_surf, show_edges=True,color="yellow") # inner_u
 ⇔surface is yellow
labels = dict(zlabel='Z', xlabel='X', ylabel='Y')
plotter.add_axes(**labels)
plotter.screenshot("mesh.png")
from IPython.display import Image
Image(filename='mesh.png')
```

[4]:





3.1 Define boundary and volume integration measure

```
[5]: # Define the boundary integration measure "ds" using the facet tags,
# also specify the number of surface quadrature points.
ds = ufl.Measure('ds', domain=domain, subdomain_data=facet_tags,
metadata={'quadrature_degree':4})

# Define the volume integration measure "dx"
# also specify the number of volume quadrature points.
dx = ufl.Measure('dx', domain=domain, metadata={'quadrature_degree': 4})

# Define facet normal
n = ufl.FacetNormal(domain)
```

4 Material parameters

-Arruda-Boyce model

5 Simulation time-control related params

6 Function spaces

```
[8]: # Define function space, both vectorial and scalar
    # dolfinx v0.8.0 syntax:
    U2 = element("Lagrange", domain.basix_cell(), 2, shape=(3,)) # For displacement
    P1 = element("Lagrange", domain.basix_cell(), 1) # For pressure
    #
    TH = mixed_element([U2, P1]) # Taylor-Hood style mixed element
    ME = functionspace(domain, TH) # Total space for all DOFs
    # Define actual functions with the required DOFs
    w = Function(ME)
    u, p = split(w) # displacement u, pressure p
    # A copy of functions to store values in the previous step
                 = Function(ME)
    u_old, p_old = split(w_old)
    # Define test functions
    u_test, p_test = TestFunctions(ME)
    # Define trial functions needed for automatic differentiation
    dw = TrialFunction(ME)
```

7 Initial conditions

- The initial conditions for degrees of freedom u and p are zero everywhere
- These are imposed automatically, since we have not specified any non-zero initial conditions.

8 Subroutines for kinematics and constitutive equations

```
[9]: # Deformation gradient
def F_calc(u):
    Id = Identity(3)
    F = Id + grad(u)
    return F

def lambdaBar_calc(u):
    F = F_calc(u)
    C = F.T*F
    Cdis = J**(-2/3)*C
    I1 = tr(Cdis)
    lambdaBar = sqrt(I1/3.0)
    return lambdaBar
```

```
def zeta_calc(u):
   lambdaBar = lambdaBar_calc(u)
   # Use Pade approximation of Langevin inverse
   z = lambdaBar/lambdaL
   z = conditional(gt(z, 0.95), 0.95, z) # Keep simulation from blowing up
   beta = z*(3.0 - z**2.0)/(1.0 - z**2.0)
   zeta = (lambdaL/(3*lambdaBar))*beta
   return zeta
# Generalized shear modulus for Arruda-Boyce model
def Gshear_AB_calc(u):
   zeta = zeta_calc(u)
   Gshear = Gshear_0 * zeta
   return Gshear
# Subroutine for calculating the Cauchy stress
#-----
def T_calc(u,p):
   Id = Identity(3)
   F = F_{calc}(u)
   J = det(F)
   B = F*F.T
   Bdis = J**(-2/3)*B
   Gshear = Gshear_AB_calc(u)
   T = (1/J)* Gshear * dev(Bdis) - p * Id
# Subroutine for calculating the Piola stress
def Piola_calc(u, p):
   Id = Identity(3)
   F = F_{calc}(u)
   J = det(F)
   T = T_{calc}(u,p)
   Tmat = J * T * inv(F.T)
   return Tmat
```

```
[]: ##A.Flowers Comments

# Deformation gradient
def F_calc(u):
    Id = Identity(3)
    F = Id + grad(u)
```

```
return F
##Calculation for deformation gradient tensor F
def lambdaBar_calc(u):
   F = F_{calc}(u)
   C = F.T*F
   Cdis = J**(-2/3)*C
   I1 = tr(Cdis)
   lambdaBar = sqrt(I1/3.0)
   return lambdaBar
##Scalar stretch measure used in hyperelasticity models
def zeta calc(u):
   lambdaBar = lambdaBar_calc(u)
##Isochoric stretch from deformation
   # Use Pade approximation of Langevin inverse
   z = lambdaBar/lambdaL
##Normalizes stretch from polymer network
   z = conditional(gt(z,0.95), 0.95, z) # Keep simulation from blowing up
##Prevents numeric instability; Langevin function because singular
   beta = z*(3.0 - z**2.0)/(1.0 - z**2.0)
##Pade approximation; used for stability
   zeta = (lambdaL/(3*lambdaBar))*beta
   return zeta
##Stress scalar from statistical mechanics model for polymers; accounts for \square
 finite chain extensibility. Stress tensors for nonlinear chain elasticity
# Generalized shear modulus for Arruda-Boyce model
def Gshear_AB_calc(u):
##Effective shear for nonlinear hyperelastic material
         = zeta_calc(u)
##Stretch dependent factor using inverse Langevin. Increasing of stretch-
 ⇔polymers stiffen
   Gshear = Gshear 0 * zeta
   return Gshear
##Shear module grows due to deformation
##This is important due to modeling with biological materials (i.e. tissue);
 ⇔nonlinear and stretch sensitive
#-----
# Subroutine for calculating the Cauchy stress
#-----
def T_calc(u,p):
   Id = Identity(3)
   F = F_{calc}(u)
##Deformation gradient
   J = det(F)
```

```
##Jacobian (volume change due to deformation)
   B = F*F.T
##Cauchy-Green tensor
   Bdis = J**(-2/3)*B
##Removes volumetric part to get isochoric (volume perserving aspect)
   Gshear = Gshear_AB_calc(u)
##Stretch dependent shear
   T = (1/J)* Gshear * dev(Bdis) - p * Id
   return T
##Cauchy stress calculation; shape change and pressure separated to obtain_
 ⇔deformed configuration
# Subroutine for calculating the Piola stress
def Piola_calc(u, p):
   Id = Identity(3)
   F = F_{calc}(u)
   J = det(F)
   T = T_{calc}(u,p)
   Tmat = J * T * inv(F.T)
   return Tmat
##Piola stress used in weak form of balance equation, with displacement \Box
 ⇔gradient; defined in terms of reference coordinates
```

9 Evaluate kinematics and constitutive relations

```
[10]: F = F_calc(u)
    J = det(F)
    lambdaBar = lambdaBar_calc(u)

# Piola stress
Tmat = Piola_calc(u, p)
```

```
[]: ##A.Flowers Comments

F = F_calc(u)
##F= deformation gradient tensor; u= displacement (unkown and solving for)
J = det(F)
##J= Jacobian determinant. Volume change during deformation
lambdaBar = lambdaBar_calc(u)
##Incompressible hyperelasticity; seperates volumetric deviatoric, shapeuschanging parts of deformation. Volume corrected stretch is calculated foruse in isochoric strain energy
```

```
# Piola stress
Tmat = Piola_calc(u, p)
##Computes Piola stress tensor from displacement field (u) and pressure (p)
```

10 Weak forms

```
[11]: # Residuals:
      # Res_0: Balance of forces (test fxn: u)
      # Res_1: Coupling pressure (test fxn: p)
      # Surface numbering from gmsh:
      # Physical Surface("yBot", 12)
      # Physical Surface("zBot", 13)
      # Physical Surface("xBot", 14)
      # Physical Surface("inner_surf", 15)
      # Cofactor of F
      Fcof = J*inv(F.T)
      # Create a constant for the pressure value
      pressRampCons = Constant(domain,PETSc.ScalarType(pressRamp(0)))
      # Configuration-dependent traction
      traction = - pressRampCons*dot(Fcof,n)
      # The weak form for the balance of forces
      Res_0 = inner(Tmat, grad(u_test) )*dx - dot(traction, u_test)*ds(15)
      # The weak form for the pressure
      fac_p = ln(J)/J
      Res_1 = dot( (p/Kbulk + fac_p), p_test)*dx
      # Total weak form
      Res = Res_0 + Res_1
      # Automatic differentiation tangent:
      a = derivative(Res, w, dw)
```

```
[]: ##A.Flowers Comments

# Residuals:
# Res_0: Balance of forces (test fxn: u)
# Res_1: Coupling pressure (test fxn: p)
```

```
# Surface numbering from qmsh:
# Physical Surface("yBot", 12)
# Physical Surface("zBot", 13)
# Physical Surface("xBot", 14)
# Physical Surface("inner_surf", 15)
# Cofactor of F
Fcof = J*inv(F.T)
##Cofactor matrix of deformation gradient; This is important for applying the
⇒pressure load and transforming elements between configurations
##Allows pressure loads to be applied on deforming boundaries
# Create a constant for the pressure value
pressRampCons = Constant(domain,PETSc.ScalarType(pressRamp(0)))
##Time-dependent or load-controlled simulation defined here; Able to modify⊔
⇔pressure constant due to time
# Configuration-dependent traction
traction = - pressRampCons*dot(Fcof,n)
##Traction vector (force/unit area) applied (pressure, tensile load)
##Pressure load in weak form of large deformation (non-linear) mechanics;
 Defines boundary traction caused by internal pressure in the hyperelastic
\hookrightarrow inflation
# The weak form for the balance of forces
Res_0 = inner(Tmat, grad(u_test) )*dx - dot(traction, u_test)*ds(15)
##Mechanical residual of weak form for nonlinear elasticity; used to build the
⇔residual vector. Used in FE for a deforming solid
##Internal work (stress times strain) minus the external work (traction times
⇔displacement); This is how force is blanced
# The weak form for the pressure
fac p = ln(J)/J
##Scalar factor used for compressible / incompressible materials due to \Box
 ⇔pressure / energy in nonlinear elasticity
Res_1 = dot( (p/Kbulk + fac_p), p_test)*dx
##Residual defined for pressure field in FE due to incompressible materials.
 →Differentiates volumetric strain energy
# Total weak form
Res = Res_0 + Res_1
##Defines total residual of weak form; from force balance (linear momentum / \sqcup
-mechanical equillibrium) and incompressibility (pressure equation)
# Automatic differentiation tangent:
```

```
a = derivative(Res, w, dw)
##Jacobian form to solve for nonlinear PDE
```

11 Set-up output files

```
[12]: # results file name
      results_name = "3D_sphere_inflate"
      # Function space for projection of results
      # v0.8.0 syntax:
      U1 = element("DG", domain.basix_cell(), 1, shape=(3,)) # For displacement
      PO = element("DG", domain.basix_cell(), 1) # For pressure
      V2 = fem.functionspace(domain, U1) #Vector function space
      V1 = fem.functionspace(domain, P0) #Scalar function space, must be
       ⇒discontinuous here since materials are discontinuous.
      # fields to write to output file
      u_vis = Function(V2)
      u_vis.name = "disp"
      p_vis = Function(V1)
      p_vis.name = "p"
      J_vis = Function(V1)
      J vis.name = "J"
      J_expr = Expression(J,V1.element.interpolation_points())
      lambdaBar_vis = Function(V1)
      lambdaBar_vis.name = "lambdaBar"
      lambdaBar_expr = Expression(lambdaBar, V1.element.interpolation_points())
      P11 = Function(V1)
      P11.name = "P11"
      P11_expr = Expression(Tmat[0,0],V1.element.interpolation_points())
      P22 = Function(V1)
      P22.name = "P22"
      P22_expr = Expression(Tmat[1,1],V1.element.interpolation_points())
      P33 = Function(V1)
      P33.name = "P33"
      P33_expr = Expression(Tmat[2,2],V1.element.interpolation_points())
      T = Tmat*F.T/J
      T0 = T - (1/3)*tr(T)*Identity(3)
      Mises = sqrt((3/2)*inner(T0, T0))
      Mises_vis= Function(V1,name="Mises")
```

```
Mises_expr = Expression(Mises, V1.element.interpolation_points())
# set up the output VTX files.
file_results = VTXWriter(
    MPI.COMM_WORLD,
    "results/" + results_name + ".bp",
    [ # put the functions here you wish to write to output
        u_vis, p_vis, J_vis, P11, P22, P33, lambdaBar_vis,
        Mises vis,
    ],
    engine="BP4",
def writeResults(t):
       # Output field interpolation
       u_vis.interpolate(w.sub(0))
       p_vis.interpolate(w.sub(1))
       J_vis.interpolate(J_expr)
       P11.interpolate(P11_expr)
       P22.interpolate(P22_expr)
       P33.interpolate(P33_expr)
       lambdaBar_vis.interpolate(lambdaBar_expr)
       Mises_vis.interpolate(Mises_expr)
       # Write output fields
       file results.write(t)
```

12 Infrastructure for pulling out time history data (force, displacement, etc.)

```
[13]: # infrastructure for evaluating functions at a certain point efficiently
    pointForDisp = np.array([Ri, 0, 0])
    bb_tree = dolfinx.geometry.bb_tree(domain,domain.topology.dim)
    cell_candidates = dolfinx.geometry.compute_collisions_points(bb_tree,u_opointForDisp)
    colliding_cells = dolfinx.geometry.compute_colliding_cells(domain,u_ocell_candidates, pointForDisp).array
```

13 Name the analysis step

```
[14]: # Give the step a descriptive name
step = "Stretch"
```

13.1 Boundary condtions

```
[15]: # Surface labels from qmsh:
      # Physical Surface("yBot", 12)
      # Physical Surface("zBot", 13)
      # Physical Surface("xBot", 14)
      # Physical Surface("inner_surf", 15)
      # Find the specific DOFs which will be constrained.
      xBtm_u1_dofs = fem.locate_dofs_topological(ME.sub(0).sub(0), facet_tags.dim,_u
       ⇒facet tags.find(14))
      yBtm_u2_dofs = fem.locate_dofs_topological(ME.sub(0).sub(1), facet_tags.dim,_u

¬facet_tags.find(12))
      zBtm_u3_dofs = fem.locate_dofs_topological(ME.sub(0).sub(2), facet_tags.dim,__

¬facet_tags.find(13))
      # building Dirichlet BCs
      bcs_1 = dirichletbc(0.0, xBtm_u1_dofs, ME.sub(0).sub(0)) # u1 fix - xBtm
      bcs_2 = dirichletbc(0.0, yBtm_u2_dofs, ME.sub(0).sub(1)) # u2 fix - yBtm
      bcs_3 = dirichletbc(0.0, zBtm_u3_dofs, ME.sub(0).sub(2)) # u3 fix - zBtm
      bcs = [bcs_1, bcs_2, bcs_3]
 []: ##A.Flowers Comments
```

```
# Surface labels from gmsh:
# Physical Surface("yBot", 12)
# Physical Surface("zBot", 13)
# Physical Surface("xBot", 14)
# Physical Surface("inner_surf", 15)
# Find the specific DOFs which will be constrained.
xBtm_u1_dofs = fem.locate_dofs_topological(ME.sub(0).sub(0), facet_tags.dim,_u

→facet_tags.find(14))
yBtm_u2_dofs = fem.locate_dofs_topological(ME.sub(0).sub(1), facet_tags.dim,_u

¬facet_tags.find(12))
zBtm u3 dofs = fem.locate dofs topological(ME.sub(0).sub(2), facet tags.dim,
→facet_tags.find(13))
##DoFs defined for boundary surface of mesh due to displacement field
# building Dirichlet BCs
bcs_1 = dirichletbc(0.0, xBtm_u1_dofs, ME.sub(0).sub(0)) # u1 fix - xBtm
bcs_2 = dirichletbc(0.0, yBtm_u2_dofs, ME.sub(0).sub(1)) # u2 fix - yBtm
bcs_3 = dirichletbc(0.0, zBtm_u3_dofs, ME.sub(0).sub(2)) # u3 fix - zBtm
##Pin sphere in place without over contraint
##Allows for symmetrical inflation under internal pressure
```

```
bcs = [bcs_1, bcs_2, bcs_3]
##Applies boundary conditions to the nonlinear solver
```

13.2 Define the nonlinear variational problem

```
[16]: # Set up nonlinear problem
      problem = NonlinearProblem(Res, w, bcs, a)
      # the global newton solver and params
      solver = NewtonSolver(MPI.COMM_WORLD, problem)
      solver.convergence_criterion = "incremental"
      solver.rtol = 1e-8
      solver.atol = 1e-8
      solver.max it = 50
      solver.report = True
      # The Krylov solver parameters.
      ksp = solver.krylov_solver
      opts = PETSc.Options()
      option_prefix = ksp.getOptionsPrefix()
      opts[f"{option_prefix}ksp_type"] = "preonly" # "preonly" works equally well
      opts[f"{option_prefix}pc_type"] = "lu" # do not use 'qamq' pre-conditioner
      opts[f"{option_prefix}pc_factor_mat_solver_type"] = "mumps"
      opts[f"{option_prefix}ksp_max_it"] = 30
      ksp.setFromOptions()
```

13.3 Start calculation loop

```
# Time-stepping solution procedure loop
while (round(t + dt, 9) <= Ttot):</pre>
    # increment time
    t += dt
    # increment counter
    ii += 1
    # update time variables in time-dependent BCs
    pressRampCons.value = pressRamp(t)
    # Solve the problem
    try:
        (iter, converged) = solver.solve(w)
    except: # Break the loop if solver fails
        print("Ended Early")
        break
    # Collect results from MPI ghost processes
    w.x.scatter_forward()
    # Write output to file
    writeResults(t)
    # Update DOFs for next step
    w_old.x.array[:] = w.x.array
    # Store time history variables at this time
    timeHistO[ii] = t # current time
    timeHist1[ii] = pressRamp(t) # time history of applied pressure
    timeHist2[ii] = w.sub(0).sub(0).eval([Ri, 0.0, 0.0],colliding_cells[0])[0]
 →# time history of displacement
    # Print progress of calculation
    if ii%1 == 0:
        now = datetime.now()
        current_time = now.strftime("%H:%M:%S")
        print("Step: {} | Increment: {}, Iterations: {}".\
              format(step, ii, iter))
        print("
                     Simulation Time: \{\} s of \{\} s".\setminus
              format(round(t,4), Ttot))
        print()
# close the output file.
```

```
file_results.close()
# End analysis
print("----")
print("End computation")
# Report elapsed real time for the analysis
endTime = datetime.now()
elapseTime = endTime - startTime
print("----")
print("Elapsed real time: {}".format(elapseTime))
print("----")
_____
```

```
Simulation Start
```

Step: Stretch | Increment: 1, Iterations: 3 Simulation Time: 0.1 s of 10.0 s

Step: Stretch | Increment: 2, Iterations: 3 Simulation Time: 0.2 s of 10.0 s

Step: Stretch | Increment: 3, Iterations: 3 Simulation Time: 0.3 s of 10.0 s

Step: Stretch | Increment: 4, Iterations: 3 Simulation Time: 0.4 s of 10.0 s

Step: Stretch | Increment: 5, Iterations: 3 Simulation Time: 0.5 s of 10.0 s

Step: Stretch | Increment: 6, Iterations: 3 Simulation Time: 0.6 s of 10.0 s

Step: Stretch | Increment: 7, Iterations: 3 Simulation Time: 0.7 s of 10.0 s

Step: Stretch | Increment: 8, Iterations: 3 Simulation Time: 0.8 s of 10.0 s

Step: Stretch | Increment: 9, Iterations: 3 Simulation Time: 0.9 s of 10.0 s

Step: Stretch | Increment: 10, Iterations: 3 Simulation Time: 1.0 s of 10.0 s

Step: Stretch | Increment: 11, Iterations: 3 Simulation Time: 1.1 s of 10.0 s

- Step: Stretch | Increment: 12, Iterations: 3 Simulation Time: 1.2 s of 10.0 s
- Step: Stretch | Increment: 13, Iterations: 3 Simulation Time: 1.3 s of 10.0 s
- Step: Stretch | Increment: 14, Iterations: 3 Simulation Time: 1.4 s of 10.0 s
- Step: Stretch | Increment: 15, Iterations: 3 Simulation Time: 1.5 s of 10.0 s
- Step: Stretch | Increment: 16, Iterations: 3 Simulation Time: 1.6 s of 10.0 s
- Step: Stretch | Increment: 17, Iterations: 3 Simulation Time: 1.7 s of 10.0 s
- Step: Stretch | Increment: 18, Iterations: 3 Simulation Time: 1.8 s of 10.0 s
- Step: Stretch | Increment: 19, Iterations: 3 Simulation Time: 1.9 s of 10.0 s
- Step: Stretch | Increment: 20, Iterations: 3 Simulation Time: 2.0 s of 10.0 s
- Step: Stretch | Increment: 21, Iterations: 3 Simulation Time: 2.1 s of 10.0 s
- Step: Stretch | Increment: 22, Iterations: 3 Simulation Time: 2.2 s of 10.0 s
- Step: Stretch | Increment: 23, Iterations: 3 Simulation Time: 2.3 s of 10.0 s
- Step: Stretch | Increment: 24, Iterations: 3 Simulation Time: 2.4 s of 10.0 s
- Step: Stretch | Increment: 25, Iterations: 3 Simulation Time: 2.5 s of 10.0 s
- Step: Stretch | Increment: 26, Iterations: 3 Simulation Time: 2.6 s of 10.0 s
- Step: Stretch | Increment: 27, Iterations: 3 Simulation Time: 2.7 s of 10.0 s

- Step: Stretch | Increment: 28, Iterations: 3 Simulation Time: 2.8 s of 10.0 s
- Step: Stretch | Increment: 29, Iterations: 3 Simulation Time: 2.9 s of 10.0 s
- Step: Stretch | Increment: 30, Iterations: 3 Simulation Time: 3.0 s of 10.0 s
- Step: Stretch | Increment: 31, Iterations: 3 Simulation Time: 3.1 s of 10.0 s
- Step: Stretch | Increment: 32, Iterations: 4 Simulation Time: 3.2 s of 10.0 s
- Step: Stretch | Increment: 33, Iterations: 4 Simulation Time: 3.3 s of 10.0 s
- Step: Stretch | Increment: 34, Iterations: 4 Simulation Time: 3.4 s of 10.0 s
- Step: Stretch | Increment: 35, Iterations: 4 Simulation Time: 3.5 s of 10.0 s
- Step: Stretch | Increment: 36, Iterations: 4 Simulation Time: 3.6 s of 10.0 s
- Step: Stretch | Increment: 37, Iterations: 4 Simulation Time: 3.7 s of 10.0 s
- Step: Stretch | Increment: 38, Iterations: 4 Simulation Time: 3.8 s of 10.0 s
- Step: Stretch | Increment: 39, Iterations: 4 Simulation Time: 3.9 s of 10.0 s
- Step: Stretch | Increment: 40, Iterations: 4 Simulation Time: 4.0 s of 10.0 s
- Step: Stretch | Increment: 41, Iterations: 4 Simulation Time: 4.1 s of 10.0 s
- Step: Stretch | Increment: 42, Iterations: 4 Simulation Time: 4.2 s of 10.0 s
- Step: Stretch | Increment: 43, Iterations: 4 Simulation Time: 4.3 s of 10.0 s

- Step: Stretch | Increment: 44, Iterations: 4 Simulation Time: 4.4 s of 10.0 s
- Step: Stretch | Increment: 45, Iterations: 4 Simulation Time: 4.5 s of 10.0 s
- Step: Stretch | Increment: 46, Iterations: 4 Simulation Time: 4.6 s of 10.0 s
- Step: Stretch | Increment: 47, Iterations: 4 Simulation Time: 4.7 s of 10.0 s
- Step: Stretch | Increment: 48, Iterations: 4 Simulation Time: 4.8 s of 10.0 s
- Step: Stretch | Increment: 49, Iterations: 4 Simulation Time: 4.9 s of 10.0 s
- Step: Stretch | Increment: 50, Iterations: 4 Simulation Time: 5.0 s of 10.0 s
- Step: Stretch | Increment: 51, Iterations: 4 Simulation Time: 5.1 s of 10.0 s
- Step: Stretch | Increment: 52, Iterations: 4 Simulation Time: 5.2 s of 10.0 s
- Step: Stretch | Increment: 53, Iterations: 4 Simulation Time: 5.3 s of 10.0 s
- Step: Stretch | Increment: 54, Iterations: 4 Simulation Time: 5.4 s of 10.0 s
- Step: Stretch | Increment: 55, Iterations: 4 Simulation Time: 5.5 s of 10.0 s
- Step: Stretch | Increment: 56, Iterations: 4 Simulation Time: 5.6 s of 10.0 s
- Step: Stretch | Increment: 57, Iterations: 4 Simulation Time: 5.7 s of 10.0 s
- Step: Stretch | Increment: 58, Iterations: 4 Simulation Time: 5.8 s of 10.0 s
- Step: Stretch | Increment: 59, Iterations: 4 Simulation Time: 5.9 s of 10.0 s

- Step: Stretch | Increment: 60, Iterations: 4 Simulation Time: 6.0 s of 10.0 s
- Step: Stretch | Increment: 61, Iterations: 4 Simulation Time: 6.1 s of 10.0 s
- Step: Stretch | Increment: 62, Iterations: 4 Simulation Time: 6.2 s of 10.0 s
- Step: Stretch | Increment: 63, Iterations: 4 Simulation Time: 6.3 s of 10.0 s
- Step: Stretch | Increment: 64, Iterations: 4 Simulation Time: 6.4 s of 10.0 s
- Step: Stretch | Increment: 65, Iterations: 4 Simulation Time: 6.5 s of 10.0 s
- Step: Stretch | Increment: 66, Iterations: 4 Simulation Time: 6.6 s of 10.0 s
- Step: Stretch | Increment: 67, Iterations: 4 Simulation Time: 6.7 s of 10.0 s
- Step: Stretch | Increment: 68, Iterations: 4 Simulation Time: 6.8 s of 10.0 s
- Step: Stretch | Increment: 69, Iterations: 4 Simulation Time: 6.9 s of 10.0 s
- Step: Stretch | Increment: 70, Iterations: 4 Simulation Time: 7.0 s of 10.0 s
- Step: Stretch | Increment: 71, Iterations: 4 Simulation Time: 7.1 s of 10.0 s
- Step: Stretch | Increment: 72, Iterations: 4 Simulation Time: 7.2 s of 10.0 s
- Step: Stretch | Increment: 73, Iterations: 4 Simulation Time: 7.3 s of 10.0 s
- Step: Stretch | Increment: 74, Iterations: 4 Simulation Time: 7.4 s of 10.0 s
- Step: Stretch | Increment: 75, Iterations: 4 Simulation Time: 7.5 s of 10.0 s

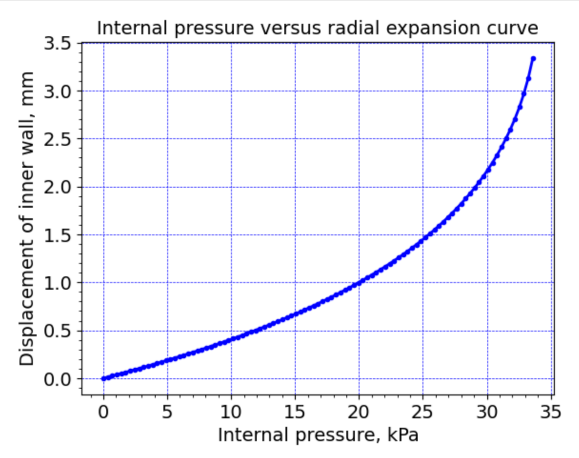
- Step: Stretch | Increment: 76, Iterations: 4 Simulation Time: 7.6 s of 10.0 s
- Step: Stretch | Increment: 77, Iterations: 4 Simulation Time: 7.7 s of 10.0 s
- Step: Stretch | Increment: 78, Iterations: 4 Simulation Time: 7.8 s of 10.0 s
- Step: Stretch | Increment: 79, Iterations: 4 Simulation Time: 7.9 s of 10.0 s
- Step: Stretch | Increment: 80, Iterations: 4 Simulation Time: 8.0 s of 10.0 s
- Step: Stretch | Increment: 81, Iterations: 4 Simulation Time: 8.1 s of 10.0 s
- Step: Stretch | Increment: 82, Iterations: 4 Simulation Time: 8.2 s of 10.0 s
- Step: Stretch | Increment: 83, Iterations: 4 Simulation Time: 8.3 s of 10.0 s
- Step: Stretch | Increment: 84, Iterations: 4 Simulation Time: 8.4 s of 10.0 s
- Step: Stretch | Increment: 85, Iterations: 4 Simulation Time: 8.5 s of 10.0 s
- Step: Stretch | Increment: 86, Iterations: 4 Simulation Time: 8.6 s of 10.0 s
- Step: Stretch | Increment: 87, Iterations: 4 Simulation Time: 8.7 s of 10.0 s
- Step: Stretch | Increment: 88, Iterations: 4 Simulation Time: 8.8 s of 10.0 s
- Step: Stretch | Increment: 89, Iterations: 4 Simulation Time: 8.9 s of 10.0 s
- Step: Stretch | Increment: 90, Iterations: 4 Simulation Time: 9.0 s of 10.0 s
- Step: Stretch | Increment: 91, Iterations: 4 Simulation Time: 9.1 s of 10.0 s

```
Step: Stretch | Increment: 92, Iterations: 4
     Simulation Time: 9.2 s of 10.0 s
Step: Stretch | Increment: 93, Iterations: 4
     Simulation Time: 9.3 s of 10.0 s
Step: Stretch | Increment: 94, Iterations: 4
     Simulation Time: 9.4 s of 10.0 s
Step: Stretch | Increment: 95, Iterations: 5
     Simulation Time: 9.5 s of 10.0 s
Step: Stretch | Increment: 96, Iterations: 5
     Simulation Time: 9.6 s of 10.0 s
Step: Stretch | Increment: 97, Iterations: 5
     Simulation Time: 9.7 s of 10.0 s
Ended Early
_____
End computation
Elapsed real time: 0:01:49.873143
```

14 Plot results

```
#plt.axis('tight')
plt.grid(linestyle="--", linewidth=0.5, color='b')
ax.set_xlabel(r'Internal pressure, kPa')
ax.set_ylabel(r'Displacement of inner wall, mm')
ax.set_title("Internal pressure versus radial expansion curve", size=14, weight='normal')
from matplotlib.ticker import AutoMinorLocator, FormatStrFormatter
ax.xaxis.set_minor_locator(AutoMinorLocator())
ax.yaxis.set_minor_locator(AutoMinorLocator())
plt.show()

fig = plt.gcf()
fig.set_size_inches(7,5)
plt.tight_layout()
plt.savefig("results/pressurise_sphere.png", dpi=600)
```



<Figure size 700x500 with 0 Axes>