

Reduced Order Modelling, Simulation and Optimization of Coupled systems

Coupled parameterized reduced order modelling of thermo-mechanical phenomena arising in blast furnace

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Authors:	Nirav Vasant Shah, ESR10, Mathematical Analysis, Modelling, and Applications, mathLab, SISSA Dr. Michele Girfoglio, Post doctoral researcher, Mathematical Analysis, Modelling, and Applications, mathLab, SISSA Dr. Patricia Barral, Associate Professor, Applied Mathematics, University of Santiago de Compostela, ITMATI Prof. Peregrina Quintela, Full Professor, Applied Mathematics, University of Santiago de Compostela, ITMATI Prof. Gianluigi Rozza, Full professor, Mathematical Analysis, Modelling, and Applications, mathLab, SISSA Ing. Alejandro lengomin, R&D Engineer, Global Research and Development Center, AMIII
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Abstract

The benchmark cases related to coupled thermomechanical phenomena arising in blast furnace have been addressed by using open-source libraries. In this document, we provide details about the numerical implementation of the benchmark tests to verify, validate and reproduce the results of numerical experiments by aiming towards smooth transition for next developers and students interested in detailed investigation of our work.

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List of Acronyms

ESR Early Stage Researcher

ITMATI Technological Institute of Industrial Mathematics
 SISSA Scuola Internazionale Superiore di Studi Avanzati
 AMIII ArcelorMittal Innovación Investigación e Inversion S.L.

1 Introduction

In our previous work [1], we focused on the physical problem, mathematical formulation, statement of the benchmark problems and design of numerical experiments. We now provide the code for numerical experiments, using open source libraries, with the objective of validating or reproducing the results and smooth handover of numerical software to future developers as per best practices [2].

2 Prerequisites

We use python 3.6.9 as the programming language. In this project we use the libraries:

- FEniCS 2019.1.0 ([3],[4],[5],[6], www.fenicsproject.org)
- RBniCS 0.1.dev1 ([7],www.rbnicsproject.org)
- Matplolib 3.1.2 ([8],www.matplotlib.org)
- numpy 1.17.4 ([9],www.numpy.org)

```
from dolfin import * #FEniCS library
from mshr import * #mshr - mesh generation component of FEniCS
from rbnics import * #RBniCS library
import matplotlib.pyplot as plt #Matplotlib library
import numpy as np #Numpy library
```

The solutions are stored in .pvd format, which can later be viewed with Paraview (www.paraview.org).

3 Installation

Simply clone the public repository:

```
$ git clone https://github.com/ROMSOC/benchmark_thermomechanical_model
```

4 Running the benchmark cases

Source codes for input data are provided in the folder *source_files*. Source codes for running the benchmark are provided in folder *source*. After running the benchmark, results are stored in folder *result_files*.

Run required .py file e.g. file_name.py as,

```
python3 file_name.py
```

5 Benchmark cases

5.1 Reading the mesh

We first construct the polygonal domain using the mshr tool.

The domain is divided into triangular 30 subdomains. We define the mapping from the reference domain to the parametrized domain. As an example, we take the subdomain 1 whose coordinates on reference domain are (0,0), (4.25,0), (0,2.365). It is deformed to the parametrized subdomain with coordinates



 $(0,0), (\mu_6,2), (0,\mu_0)$. The μ_6 and μ_0 are the 6th and 0th parameter of zero-indexed tuple Ξ . This mapping can be defined as,

```
{
    ("0", "0"): ("0", "0"),
    ("4.25", "0"): ("mu[6]/2", "0"),
    ("0", "2.365"): ("0", "mu[0]")
}, # subdomain 1
```

The RBniCS computes the mapping for each subdomain and uses it during affine transformation of operators. Similarly, the mappings and subdomains are created for other 29 subdomains.

Next, we set the subdomain marker:

The mesh and subdomains are created based on subdomain markers.

```
# Create mesh
mesh = generate_mesh(domain, 30) #30 specifies the mesh size.
# Create subdomains
subdomains = MeshFunction("size_t", mesh, 2, mesh.domains())
```

We now set the boundary markers. The domain boundaries are shared across 20 subdomains, hence we define 20 classes and set markers for each of these boundaries. For example, for the boundary γ_s , we define the class

```
class Gamma_s(SubDomain):
   def inside(self, x, on_boundary):
     return x[0] < DOLFIN_EPS and on_boundary</pre>
```

and create an instance of this class and set boundary marker as 1.

```
gamma_s = Gamma_s()
gamma_s.mark(boundaries, 1)
```

Unlike γ_s , the bottom boundary γ_- is shared by 5 subdomains, we define 5 different classes and correspondingly 5 different markers.



```
gamma_minus1 = Gamma_minus1()
gamma_minus1.mark(boundaries, 2)
gamma_minus2 = Gamma_minus2()
gamma_minus2.mark(boundaries, 3)
gamma_minus3 = Gamma_minus3()
gamma_minus3.mark(boundaries, 4)
gamma_minus4 = Gamma_minus4()
gamma_minus4 = Gamma_minus5()
gamma_minus5 = Gamma_minus5()
gamma_minus5.mark(boundaries, 6)
```

In a similar manner, markers are set for other boundaries and the mesh, the boundary markers, the subdomain markers and affine maps are stored.

At the beginning of any benchmark test, we first read the mesh, the subdomains and also read the boundary markers. The volume of each subdomains and length of each boundary are measured. Notice that all boundary markers with same boundary condition are combined.

```
# Read the mesh file from specified path.
mesh = Mesh("../../benchmarks/data_files/mesh_data/hearth.xml")
domains = MeshFunction('size_t', mesh, mesh.topology().dim()) # Read domain marker
subdomains = MeshFunction("size_t", mesh, "../../benchmarks/data_files/mesh_data/
                                                                                                                                        hearth_physical_region.xml") # Read
                                                                                                                                         subdomain markers
boundaries = MeshFunction("size_t", mesh, "../../benchmarks/data_files/mesh_data/
                                                                                                                                        hearth_facet_region.xml") # Read boundary
                                                                                                                                        markers
dx = Measure('dx', domain = mesh, subdomain_data = domains) # Volume measure
ds = Measure('ds', domain = mesh, subdomain_data = boundaries) # Boundary measure
n = as_vector(FacetNormal(mesh)) # Edge unit normal vector
d_{bottom} = ds(2) + ds(3) + ds(4) + ds(5) + ds(6) # Markers of bottom boundary \gamma_{-}
d_{out} = ds(7) + ds(8) + ds(9) + ds(10) + ds(11) # Markers of outer boundary <math>g_{out} = ds(7) + ds(8) + ds(9) + ds(9) + ds(10) + ds(11) + ds(11)
d_sf = ds(13) + ds(14) + ds(15) + ds(16) + ds(17) + ds(18) + ds(19) + ds(20) # Markers of
                                                                                                                                            inner boundary \gamma_{sf}
```

5.2 Thermal model

We define the norm for $\psi \in H_r^1(\omega)$,



```
#Computation of H^1_r(\omega) norm

def compute_hlr_norm(psi,mesh):
    r = SpatialCoordinate(mesh)[0]
    dx = Measure('dx', domain = mesh)
    a = inner(psi,psi)*r*dx + inner(grad(psi),grad(psi))*r*dx
    A = assemble(a)
    return sqrt(A)
```

Next, we define the range of polynomials degree for measuring p-convergence and an object to store the relative error.

```
error_T_vector = [] #List to store error in temperature w.r.t. polynomial degree
p = range(1,4) # List of polynomial degrees
```

Next, we specify the relevant physical parameters, which are than later used in weak form.

```
k = 10. # Thermal conductivity
h_fluid = 200. # Convection coefficient on \gamma_{sf}
h_right = 2000. # Convection coefficient on \gamma_{out}
h_bottom = 2000. # Convection coefficient on \gamma_{-}
```

For every polynomial degree, we define the relevant "Lagrange" function space of a particular degree. We also define the solution field and test function in this space.

```
# Define function space
VT = FunctionSpace(mesh, "CG",i) # Function space for temperature
psi, T_ = TestFunction(VT), TrialFunction(VT) # Evaluate trial and test function
T = Function(VT, name = "temperature increase")
x = list()
x.append(Expression("x[0]", element=VT.ufl_element())) #r coordinate
x.append(Expression("x[1]", element=VT.ufl_element())) #y coordinate
```

We then define and solve the equation in weak formulation.

```
# solving weak form of energy equation
a_T = k * inner(grad(psi),grad(T_)) * x[0] * dx + \
h_fluid * psi * T_ * x[0] * d_sf + h_right * psi * T_ * x[0] * d_out + \
h_bottom * psi * T_ * x[0] * d_bottom # Bilinear side

1_T = h_fluid * psi * (x[0] * x[0] * x[1] + k/h_fluid * (2 * x[0] * x[1] * n[0] + x[0] * x[0] * n[1])) * x[0] * d_sf + \
h_right * psi * (x[0]*x[0]*x[1]+2*x[0]*x[1]*k/h_right) * x[0] * d_out + \
h_bottom * psi * (x[0] * x[0] * x[1] - x[0] * x[0] * k / h_bottom) * x[0] * d_bottom + \
-4 * k * x[1] * psi * x[0] * dx + psi * k * x[0] * x[0] * x[0] * ds(12) # Linear side

solve(a_T == 1_T, T) # Solve the variational form
```

After performing the computations, we store the data in format compatible with paraview for further visualization. Also, we plot the p-convergence.



```
# Plotting and printing convergence tests
plt.figure(figsize=[10,8])
a = plt.semilogy([1,2,3],error_T_vector,marker='o',linewidth=4)
plt.xticks([1,2,3],fontsize=18)
plt.yticks(fontsize=18)
plt.xlabel('Polynomial degree',fontsize=24)
plt.ylabel('Relative error',fontsize=24)
plt.axis('tight')
plt.savefig("../../benchmarks/result_files/thermal_model/convergence_test")
plt.show()
```

5.3 Mechanical model

We define the norm for $\overrightarrow{\phi} \in \mathbb{U}$.

```
#Computation of \mathbb{U} norm

def compute_U_norm(phi,mesh):
    x = SpatialCoordinate(mesh)
    dx = Measure('dx', domain = mesh)
    a = inner(phi,phi)*x[0]*dx + inner(grad(phi),grad(phi))*x[0]*dx + (phi[0]**2)/x[0]*dx
    A = assemble(a)
    return sqrt(A)
```

Next, we define the axisymmetric stress and strain tensor.

Since, our aim is to assess p—convergence, we define the range of polynomial degrees and measure corresponding relative errors.

```
error_u_vector = [] #List for absolute error in displacement
p = range(1,4) #Polynomial degrees
```

We introduce the physical properties.

```
E = Constant(5e9) # Young's modulus
nu = Constant(0.2) # Poisson's ratio
mu = E/2/(1+nu) # Lam\'e parameter
lmbda = E*nu/(1+nu)/(1-2*nu) # Lam\'e parameter
```

We now define the function space for displacement and stress. We also initialize variables for the test function and the solution field.

```
# Define function space
VM = VectorFunctionSpace(mesh, "CG",i) # Function space for displacement
x = Expression(("x[0]","x[1]"), element=VM.ufl_element())
VS = FunctionSpace(mesh, "CG", max(i-1,1))
# Function space for Von Mises stress NOTE: when i=1, the VS is of degree 1 and not 0.
phi, u_ = TestFunction(VM), TrialFunction(VM)
u = Function(VM, name = "Displacement") # u[0] = u_r and u[1] = u_y
```



We define the variables related to the source term and the boundary data.

```
# Dirichlet boundary data
bcs_M = [DirichletBC(VM.sub(0), Constant(0.), 'x[0] < DOLFIN_EPS and on_boundary'), \
     DirichletBC( VM.sub(1), Constant(0.), 'near(x[1],0) and on_boundary')]
 # Set Dirichlet boundary. Note that only functions which satisfy zero normal displacement
                                                                                                                                         on \gamma_s \cup \gamma_- are admissible.
 # Source term and relevant boundary data
f0_r = - (2*E*nu*1e-4*x[0]/(1-2*nu)/(1+nu)+2*E*1e-4*x[0]/(1+nu))
f0_y = - (4 \times E \times 1e - 4 \times x[1] / (1+nu) + 4 \times E \times 1e - 4 \times x[1] \times nu / (1-2 \times nu) / (1+nu))
g_plus_r = 2*E*1e-4*x[0]*x[1]/(1+nu)
q_plus_y = E / (1-2*nu) / (1+nu) * (2*nu*1e-4*x[1]*x[1]+(1-nu)*1e-4*x[0]*x[0])
q_minus_r = -q_plus_r
q_sf_r = E / (1-2*nu) / (1+nu) * (1e-4 * x[1] * x[1] + nu * 1e-4 * x[0] * x[0]) * n[0] +
                                                                                                                                       2 * E * 1e-4 * x[0] * x[1] / (1 + nu) * n[1]
q_sf_y = 2 * E * 1e-4 * x[0] * x[1] / (1 + nu) * n[0] + E / (1-2*nu) / (1+nu) * (2 * nu * nu) + (1+nu) * (2 * nu) + (1+nu) * (1
                                                                                                                                         1e-4 * x[1] * x[1] + (1 - nu) * 1e-4 * x[0]
                                                                                                                                          * x[0]) * n[1]
g_out_r = E / (1-2*nu) / (1+nu) * (1e-4 * x[1] * x[1] + nu * 1e-4 * x[0] * x[0])
g_{out_y} = 2*E*1e-4*x[0]*x[1]/(1+nu)
```

The equation in weak form is then defined and solved for computing displacement. Based on this displacement, the Von Mises stress is computed.

The data is stored in format compatible to paraview. We also plot the polynomial degree vs the relative error.



```
error_stress = Function(VS) #Function for absolute error in stress tensor
error_stress.vector()[:] = abs(project(sigma_vm, VS).vector().get_local() - project(
                                              sigma_vm_analytical, VS) .vector() .get_local()
File("../../benchmarks/result_files/mechanical_model/von_mises_stress_computed.pvd") <<
                                             project(sigma_vm, VS)
File("../../benchmarks/result_files/mechanical_model/von_mises_stress_analytical.pvd") <<
                                              project(sigma_vm_analytical, VS)
File("../../benchmarks/result_files/mechanical_model/von_mises_stress_error.pvd") <<
                                             project (error_stress, VS)
#Convergence tests
plt.figure(figsize=[10,8])
a = plt.semilogy([1,2,3],error_u_vector,marker='o',linewidth=4)
plt.xticks([1,2,3],fontsize=18)
plt.yticks(fontsize=18)
plt.xlabel('Polynomial degree', fontsize=24)
plt.ylabel('Relative error', fontsize=24)
plt.axis('tight')
plt.savefig("../../benchmarks/result_files/mechanical_model/convergence_test")
plt.show() # To show the plots
print("Relative error in U norm: "+ str(error_u_vector))
```

5.4 Coupled model

Similar to the Thermal model (Section 5.2), we first solve the energy equation in weak form.

```
# Define function space
VT = FunctionSpace(mesh, "CG", 3) # Function space for temperature
psi, T_ = TestFunction(VT), TrialFunction(VT) # Evaluate trial and test function
T = Function(VT, name = "temperature increase")
# Known analytical solution, Thermal material properties and Boundary data
T_{analytical} = Expression('x[0]*x[0]*x[1]', degree = 3)
VT_analytical = FunctionSpace(mesh, "CG", 3) #Space for analytical solution
T_analytical = project(T_analytical, VT_analytical)
k = 10. # Thermal conductivity
h_fluid = 200. # Convection coefficient on \gamma_{sf}
h_right = 2000. # Convection coefficient on \gamma_{out}
h_bottom = 2000. # Convection coefficient on \gamma_{-}
x = list()
x.append(Expression("x[0]", element=VT.ufl_element())) #r coordinate
x.append(Expression("x[1]", element=VT.ufl_element())) #y coordinate
# solving weak form of energy equation
a_T = k * inner(grad(psi), grad(T_)) * x[0] * dx + 
    h_fluid * psi * T_ * x[0] * d_sf + h_right * psi * T_ * x[0] * d_out + \
   h_bottom * psi * T_ * x[0] * d_bottom # Bilinear side
1_T = h_fluid * psi * (x[0] * x[0] * x[1] + k/h_fluid * (2 * x[0] * x[1] * n[0] + x[0] * x[1] * n[0] * x[0] * x[
                                                                                                           x[0] * n[1] ) ) * x[0] * d_sf + 
    h_{right} * psi * (x[0]*x[0]*x[1]+2*x[0]*x[1]*k/h_{right}) * x[0] * d_{out} + 
    -4 \ * \ k \ * \ x[1] \ * \ psi \ * \ x[0] \ * \ dx \ + \ psi \ * \ k \ * \ x[0] \ * \ x[0] \ * \ x[0] \ * \ ds(12) \ \# \ \textit{Linear side}
solve(a_T == l_T, T) # Solve the variational form
```

Also, similar to mechanical model (section 5.3), we define the \mathbb{U} norm, stress and strain tensor. Additionally, we define the thermomechanical stress tensor.



```
# Define \mathbb{U} norm
def compute_U_norm(phi, mesh):
 x = SpatialCoordinate(mesh)
 a = inner(phi, phi) *x[0] *dx + inner(qrad(phi), qrad(phi)) *x[0] *dx + (phi[0] **2/x[0]) *dx
 A = assemble(a)
 return sqrt (A)
# Axisymmetric strain tensor definition. Alternative could be to express strain as vector
                                              using Voigt notation.
def eps(u):
 return \
   sym(as\_tensor([[u[0].dx(0), u[0].dx(1), 0.], \
    [u[1].dx(0), u[1].dx(1), 0.], \
    [0., 0., u[0]/x[0]]))
# Axisymmetric thermo-mechanical stress tensor definition. Alternative could be to
                                             express as vector using Voigt notation.
def sigma(u,T):
 return lmbda * tr(eps(u)) * Identity(3) + 2.0 * mu * eps(u) - (2 * mu + 3 * lmbda) *
                                             alpha * (T - T_0) * Identity(3)
# Axisymmetric mechanical stress tensor definition. Alternative could be to express as
                                             vector using Voigt notation.
def sigma2(u):
 return lmbda * tr(eps(u)) * Identity(3) + 2.0 * mu * eps(u)
```

Next, the physical data are specified.

```
T_0 = 298 # Reference temperature for zero thermal stress
E = Constant(5e9) # Young's modulus
nu = Constant(0.2) # Poisson's ratio
mu = E/2/(1+nu) # Lame\'e parameter
lmbda = E*nu/(1+nu)/(1-2*nu) # Lame\'e parameter
alpha = Constant(1e-6) # Thermal expansion coefficient
```

Similar to the mechanical model (section 5.3), we solve the weak form.

```
# Define function space for displacement
VM = VectorFunctionSpace(mesh, "CG",i) # Function space for displacement
x = Expression(("x[0]","x[1]"), element=VM.ufl_element())
phi, u_{-} = TestFunction(VM), TrialFunction(VM)
 u = Function(VM, name = "Displacement") # u[0] = u_r and u[1] = u_y
VS = FunctionSpace(mesh, "CG", max(i-1,1)) # Function space for shear component of stress
  # Dirichlet boundary data
bcs_M = [DirichletBC(VM.sub(0), Constant(0.), 'x[0] < DOLFIN_EPS and on_boundary'),
                                                                                                                                                                                                                                                                                                                                                                                      DirichletBC( VM.sub(1), Constant(0.), 'near(
                                                                                                                                                                                                                                                                                                                                                                                      x[1],0) and on_boundary')]
   #Boundary and source terms
  \texttt{f0\_r} \ = \ - \ (2 \times \texttt{E} \times \texttt{nu} \times \texttt{1e} - 4 \times \texttt{x[0]} / (\texttt{1} - 2 \times \texttt{nu}) / (\texttt{1} + \texttt{nu}) + 2 \times \texttt{E} \times \texttt{1e} - 4 \times \texttt{x[0]} / (\texttt{1} + \texttt{nu}) - 2 \times \texttt{E} \times \texttt{x[0]} \times \texttt{x[1]} \times \texttt{alpha} / (\texttt{1} - 2 \times \texttt{nu}) / (\texttt{1}
                                                                                                                                                                                                                                                                                                                                                                                     n11))
  \texttt{f0\_y} \ = \ - \ (4 \times \texttt{E} \times \texttt{1e} - 4 \times \texttt{x} \texttt{[1]} / (1 + \texttt{nu}) + 4 \times \texttt{E} \times \texttt{1e} - 4 \times \texttt{x} \texttt{[1]} \times \texttt{nu} / (1 - 2 \times \texttt{nu}) / (1 + \texttt{nu}) - \texttt{E} \times \texttt{x} \texttt{[0]} \times \texttt{x} \texttt{[0]} \times \texttt{alpha} / (1 - 2 \times \texttt{nu}) / (1 - 2 \times \texttt{nu}
                                                                                                                                                                                                                                                                                                                                                                                      ) )
 g_plus_r = 2*E*1e-4*x[0]*x[1]/(1+nu)
 g_plus_y = E / (1-2*nu) / (1+nu) * (2*nu*1e-4*x[1]*x[1]+(1-nu)*1e-4*x[0]*x[0]) - E*alpha/
                                                                                                                                                                                                                                                                                                                                                                                      (1-2*nu)*(x[0]*x[0]*x[1] - T_0
 g_minus_r = -g_plus_r
```



```
q_sf_r = (E / (1-2*nu) / (1+nu) * (1e-4 * x[1] * x[1] + nu * 1e-4 * x[0] * x[0]) - E*
                                             alpha/(1-2*nu)*(x[0]*x[0]*x[1] - T_0)) * n[0]
                                              ] + 2 * E * 1e-4 * x[0] * x[1] / (1 + nu) *
                                             n[1]
g_sf_y = 2 * E * 1e-4 * x[0] * x[1] / (1 + nu) * n[0] + (E / (1-2*nu) / (1+nu) * (2 * nu)
                                              * 1e-4 * x[1] * x[1] + (1 - nu) * 1e-4 * x[0]
                                               \times x[0] - E*alpha/(1-2*nu)*(x[0]*x[0]*x[1] 
                                             - T_0)) * n[1]
g_{out} = E / (1-2*nu) / (1+nu) * (1e-4 * x[1] * x[1] + nu * 1e-4 * x[0] * x[0]) - E*
                                             alpha/(1-2*nu)*(x[0]*x[0]*x[1] - T_0)
q_out_y = 2*E*1e-4*x[0]*x[1]/(1+nu)
# solving weak form of momentum equation
# This is not bilinear side as terms related to thermal stress are included.
a_M1 = inner(sigma(u_T), eps(phi)) * x[0] * dx
# This is not linear side as terms related to thermal stress are not included.
l_M1 = (phi[0] * f0_r + phi[1] * f0_y) * x[0] * dx + (phi[0] * g_plus_r + phi[1] *
                                             g_plus_y) * x[0] * ds(12) + 
 (phi[0] * g_minus_r) * x[0] * d_bottom + (phi[0] * g_sf_r + phi[1] * g_sf_y) * x[0] * \\
                                             d_sf + 
(phi[0] * g_out_r + phi[1] * g_out_y) * x[0] * d_out
F = a_M1 - l_M1
a_M = lhs(F) # Now a_M is bilinear form
l_M = rhs(F) # Now l_M is linear form
solve(a_M == 1_M, u, bcs_M) # Solve equation
# Compute \mathbb{U} norm of error
error_u = compute_U_norm(u_analytical-u, mesh)/compute_U_norm(u_analytical, mesh)
error_u_vector.append(error_u)
print("Relative error in U-norm : ", str(error_u))
```

We compute the relevant stress fields.

Finally, we store the solution field for further visualization.



```
File("../../benchmarks/result_files/coupled_model/displacement_absolute_error.pvd") <</pre>
                                           project(u_analytical - u,VM)
File("../../benchmarks/result_files/coupled_model/von_mises_computed_coupling.pvd") <</pre>
                                           project(sigma_vm, VS)
File("../../benchmarks/result_files/coupled_model/von_mises_analytical_coupling.pvd") <<
                                           project(sigma_vm_analytical, VS)
error_stress_von_mises = Function(VS)
error_stress_von_mises.vector()[:] = abs(project(sigma_vm, VS).vector().get_local() -
                                           project(sigma_vm_analytical, VS).vector().
                                           get_local())
File("../../benchmarks/result_files/coupled_model/von_mises_stress_error_coupling.pvd") 
                                            < project(error_stress_von_mises, VS)</pre>
File("../../benchmarks/result_files/coupled_model/difference_in_spherical_stress.pvd") <<
                                            project(sigma_spherical -
                                            sigma_spherical_non_thermal, VS)
(-(2 * mu + 3 * lmbda) * alpha * (T - T_0),
error_stress_spherical = Function(VS)
error_stress_spherical.vector()[:] = abs(project(sigma_spherical,VS).vector().get_local()
                                            - project(sigma_spherical_non_thermal - (2
                                            * mu + 3 * lmbda) * alpha * (T - T_0), VS).
                                           vector().get_local())
File("../../benchmarks/result_files/coupled_model/absolute_error_spherical_stress.pvd") <
                                           < error_stress_spherical</pre>
#Convergence tests
plt.figure(figsize=[10,8])
a = plt.semilogy([1,2,3],error_u_vector,marker='o',linewidth=4)
plt.xticks([1,2,3],fontsize=18)
plt.yticks(fontsize=18)
plt.xlabel('Polynomial degree', fontsize=24)
plt.ylabel('Relative error', fontsize=24)
plt.axis('tight')
plt.savefig('../../benchmarks/result_files/coupled_model/
                                           Convergence_coupling_displacement_benchmark_comparison
                                            .png')
plt.show()
```

5.5 Reduced basis method

For the affine geometric parametrization, we use 2 decorators: one for the affine shape parametrization and the other for transfer of operators between reference domain and parametrized domain.

```
@PullBackFormsToReferenceDomain() #Decorator for operator transformation between
parameterized domain to reference domain
@AffineShapeParametrization("../../benchmarks/data_files/mesh_data/
hearth_vertices_mapping.vmp") #Decorator for
shape parametrization with mapping defined
in specified file
```

To compute the temperature field required to compute displacement for coupling model, we use another decorator:

```
\verb§ExactParametrizedFunctions() # Decorator for computing temperature field required for linear side
```



Considering that the solution computed by finite element method is used as benchmark for assessing the accuracy of the reduced basis method, we refer to the solution computed by finite element method as "Truth solution".

5.5.1 Thermal system

We first define the class *HearthThermal*, inherited from *EllipticCoerciveProblem*, for the thermal system.

```
class HearthThermal(EllipticCoerciveProblem):
```

The default initialization involves all the parameters related to the problem.

```
# Default initialization of members
def __init__(self, V, **kwargs):
 # Call the standard initialization
 EllipticCoerciveProblem.__init__(self, V, **kwargs)
  # ... and also store FEniCS data structures for assembly
 assert "subdomains" in kwargs
 assert "boundaries" in kwargs
 assert "mesh" in kwargs
 assert "h_cf" in kwarqs
 assert "h_out" in kwargs
 assert "h_bottom" in kwargs
 self.subdomains, self.boundaries = kwargs["subdomains"], kwargs["boundaries"]
 self.u = TrialFunction(V)
 self.v = TestFunction(V)
 self.dx = Measure("dx")(subdomain_data=subdomains)
 self.ds = Measure("ds")(subdomain data=boundaries)
 self.subdomains = subdomains
 self.boundaries = boundaries
 self.reference_mesh = kwarqs["mesh"]
 self.h_cf = kwargs["h_cf"]
 self.h_out = kwarqs["h_out"]
 self.h_bottom = kwargs["h_bottom"]
 self.x0 = Expression("x[0]", element=V.ufl_element())
```

Firstly, the affine mulitplicative terms and next the weak formulation are defined.

```
# Return theta multiplicative terms of the affine expansion of the problem.
def compute_theta(self, term):
 mu = self.mu
 if term == "a":
   theta_a0 = mu[10]
   theta_a1 = 1.0
   return (theta_a0, theta_a1)
 elif term == "f":
   theta f0 = 1.0
   return (theta_f0, )
   raise ValueError("Invalid term for compute_theta().")
# Return forms resulting from the discretization of the affine expansion of the problem
                                            operators.
def assemble_operator(self, term):
 u = self.u
 v = self.v
 reference_mesh = self.reference_mesh
 dx = self.dx
 ds = self.ds
 h_cf = self.h_cf
```



```
h_out = self.h_out
h_bottom = self.h_bottom
r = self.x0
d_bottom = ds(2) + ds(3) + ds(4) + ds(5) + ds(6)
d_{out} = ds(7) + ds(8) + ds(9) + ds(10) + ds(11)
d_sf = ds(13) + ds(14) + ds(15) + ds(16) + ds(17) + ds(18) + ds(19) + ds(20)
if term == "a":
  a0 = inner(grad(u), grad(v))*r*dx
  a1 = h\_bottom*u*v*r*d\_bottom + h\_out*u*v*r*d\_out + h\_cf*u*v*r*d\_sf
  return (a0, a1)
elif term == "f":
  \texttt{f0} = \texttt{h\_bottom} * 313 * \texttt{v} * \texttt{r} * \texttt{d\_bottom} + \texttt{h\_out} * 313 * \texttt{v} * \texttt{r} * \texttt{d\_out} + \texttt{h\_cf} * 1773 * \texttt{v} * \texttt{r} * \texttt{d\_sf}
  return (f0, )
elif term == "inner_product":
  x0 = u*v*r*dx + inner(grad(u), grad(v))*r*dx
  return (x0,)
else:
  raise ValueError("Invalid term for assemble_operator().")
```

Using the *HearthThermal* class, we now perform POD-Galerkin approximation of the thermal problem. The function space is defined first. Next, an instance of *HearthThermal* class, *hearth_problem_thermal* is created and model order reduction is performed.

```
# 2A. Create Finite Element space (Lagrange P1)
VT = FunctionSpace(mesh, "Lagrange", 1) # For temperature
# 3A. Allocate an object of the Hearth class
hearth_problem_thermal = HearthThermal(VT, subdomains=subdomains, boundaries=boundaries,
                                             mesh=mesh, h_cf=200., h_out=2000., h_bottom=
                                              2000.)
#specify and set range of each parameter
mu_range = [(2.3, 2.4), (0.5, 0.7), (0.5, 0.7), (0.4, 0.6), (3.05, 3.35), (13.5, 14.5), (8.3, 8.5)
                                              7), (8.8,9.2), (9.8,10.2), (10.4,10.8), (9.8
                                              ,10.2), (2.08e9,2.08e9), (1.39e9,1.39e9), (
                                             1e-6, 1e-6)]
hearth_problem_thermal.set_mu_range(mu_range)
# 4A. Prepare reduction with a POD-Galerkin method
#NOTE : truth_problem attribute is FEM problem and reduced_problem is RB problem
pod_galerkin_method_thermal = PODGalerkin(hearth_problem_thermal)
pod_galerkin_method_thermal.set_Nmax(100) #Maximum size of reduced basis space
pod_galerkin_method_thermal.set_tolerance(1e-4) #Maximum eigenvalue tolerance
```

Using pod_galerkin_method_thermal, we perform the offline phase for the thermal system.

Next, we perform the error analysis, compute the time taken for truth solution and reduced basis solution.



```
testing_set_speedup_analysis = pod_galerkin_method_thermal.testing_set
pod_galerkin_method_thermal._patch_truth_solve(True) #To enable cahce reading
truth_timer = Timer("parallel") #Timer for computation of FEM solution
time_thermal_truth = np.empty(len(testing_set_speedup_analysis)) #Storage of time taken
                                             for solving FEM equation. It is a vector of
                                             size of number of speedup analysis
                                             parameters
# Iteration over speedup analysis parameters for measuring time taken for FEM solution
for (mu_index, mu_test) in enumerate(testing_set_speedup_analysis):
 print (TextLine(str(mu_index), fill="#"))
 pod_galerkin_method_thermal.truth_problem.set_mu(mu_test) #Set the parameter
 truth_timer.start()
  pod_galerkin_method_thermal.truth_problem.solve() #Solve the FEM problem
 truth_time_thermal = truth_timer.stop()
  print("Truth time thermal : ",truth_time_thermal)
  time_thermal_truth[mu_index] = truth_time_thermal #Save time taken for truth solve
np.save("time_thermal_truth",time_thermal_truth) #Save time taken for computation of FEM
                                             solution
pod_galerkin_method_thermal._undo_patch_truth_solve(True) #To enable cache reading
# 8A2. Perform a speedup analysis - Compute time for reduced solutions
pod_galerkin_method_thermal._patch_truth_solve(True) #To disable cache reading
reduced_timer = Timer("serial") #Timer for computation of RB solution
max_basis_function = reduced_hearth_problem_thermal.N #Size of reduced basis space
time_thermal_reduced = np.empty([max_basis_function,len(testing_set_speedup_analysis)]) #
                                             Storage of time taken for solving RB
                                             equation. It is a matrix of size size of
                                             reduced basis space \times number of speedup
                                              analysis parameters
# Iteration over speedup analysis parameters for measuring time for RB solution
for basis_size in range(1, max_basis_function+1):
  for (mu_index, mu_test) in enumerate(testing_set_speedup_analysis):
   print(TextLine(str(mu_index), fill="#"))
    pod_galerkin_method_thermal.reduced_problem.set_mu(mu_test) #Set parameter
    reduced_timer.start()
    pod_galerkin_method_thermal.reduced_problem.solve(basis_size) #Solve the RB problem
   rb_time_thermal = reduced_timer.stop()
   print("Reduced time thermal: ", rb_time_thermal)
    time_thermal_reduced[basis_size-1, mu_index] = rb_time_thermal #Save time taken for RB
                                              solve
pod_galerkin_method_thermal._undo_patch_truth_solve(True) #To disable cache reading
np.save("time_thermal_reduced",time_thermal_reduced) #Save time taken for computation for
                                              RB solution
```

For any new parameter *online_mu*, the truth solution and the reduced basis solution can be performed using these classes.



5.5.2 Mechanical system

We first define the class *HearthMechanical*, inherited from *EllipticCoerciveProblem*, for the mechanical system.

```
class HearthMechanical(EllipticCoerciveProblem):
```

We specify the default initialization for this class.

```
# Default initialization of members
def __init__(self, V, **kwargs):
  # Call the standard initialization
 EllipticCoerciveProblem.__init__(self, V, **kwargs)
  # ... and also store FEniCS data structures for assembly
 assert "subdomains" in kwargs
 assert "boundaries" in kwargs
 assert "mesh" in kwargs
 self.normal = as_vector(FacetNormal(kwargs["mesh"]))
 self.subdomains, self.boundaries = kwargs["subdomains"], kwargs["boundaries"]
 self.u = TrialFunction(V)
 self.v = TestFunction(V)
 self.dx = Measure("dx")(subdomain_data=subdomains)
 self.ds = Measure("ds")(subdomain_data=boundaries)
 self.subdomains = subdomains
 self.boundaries = boundaries
 self.x0 = Expression("x[0]", element=V.sub(0).ufl_element())
 self.x1 = Expression("x[1]", element=V.sub(1).ufl_element())
```

Next, the affine multiplicative terms, weak forms and strain tensors are defined.

```
# Return theta multiplicative terms of the affine expansion of the problem.
def compute_theta(self, term):
 mu = self.mu
 if term == "a":
   theta_a0 = mu[11]
   theta_a1 = 2*mu[12]
   return (theta_a0, theta_a1, )
 elif term == "f":
   theta_f0 = 1.0
    return (theta_f0, )
 else:
   raise ValueError("Invalid term for compute_theta().")
# Return strain tensor
def strain(self,u):
 r = self.x0
 return sym(as_tensor([[u[0].dx(0), u[0].dx(1), 0.], [u[1].dx(0), u[1].dx(1), 0.], [0
                                           u[0]/r]
```



```
# Return forms resulting from the discretization of the affine expansion of the problem
                                                                                                               operators.
def assemble_operator(self, term):
    u = self.u
    v = self.v
    dx = self.dx
    ds = self.ds
    r = self.x0
    x1 = self.x1
    n = self.normal
    d_{bottom} = ds(2) + ds(3) + ds(4) + ds(5) + ds(6)
    d_{out} = ds(7) + ds(8) + ds(9) + ds(10) + ds(11)
    d_sf = ds(13) + ds(14) + ds(15) + ds(16) + ds(17) + ds(18) + ds(19) + ds(20)
    if term == "a":
         a0 = (u[0].dx(0)+u[1].dx(1)+u[0]/r)*(v[0].dx(0)+v[1].dx(1)+v[0]/r)*r*dx
         a1 = (u[0].dx(0)*v[0].dx(0) + u[1].dx(1)*v[1].dx(1) + (u[0]*v[0])/(r)**2 + 0.5*(u[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v
                                                                                                             ].dx(1)+u[1].dx(0))*(v[0].dx(1)+v[1].dx(0)))
                                                                                                                * r * dx
         return (a0, a1,)
    elif term == "f":
          f0 = - dot(v, 7460*9.81*(7.265-x1)*n) * r * d_sf
          return (f0,)
    elif term == "inner_product":
         x0 = inner(u,v) * r * dx + inner(self.strain(u),self.strain(v)) * r * dx
          return (x0,)
    elif term == "dirichlet_bc":
         bc0 = [DirichletBC(self.V.sub(0), Constant(0.), self.boundaries, 1),
              DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 2),
               DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 3),
               DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 4),
              \label{eq:constant} \mbox{DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 5),}
              DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 6),]
         return (bc0,)
    else:
          raise ValueError("Invalid term for assemble_operator().")
```

The function space and an instance *hearth_problem_mechanical* of *HearthMechanical* class are defined.

```
# 2B. Create Finite Element space (Lagrange P1)
VM = VectorFunctionSpace(mesh, "Lagrange",1) # For mechanical
# 3B. Allocate an object of the HearthThermoMechanical class
hearth_problem_mechanical = HearthMechanical(VM, subdomains=subdomains, boundaries= boundaries, mesh=mesh)
# specify and set range of each parameter
mu_range = [(2.3,2.4), (0.5,0.7), (0.5,0.7), (0.4,0.6), (3.05,3.35), (13.5,14.5), (8.3,8.7), (8.8,9.2), (9.8,10.2), (10.4,10.8), (10.10.1), (1.9e9,2.5e9), (1.2e9,1.8e9), (1e-6,1e-6)]
hearth_problem_mechanical.set_mu_range(mu_range)
```

Next, reduction with POD-Galerkin method is initialised and offline phase is performed.

```
# 4B. Prepare reduction with a POD-Galerkin method

#NOTE: truth_problem attribute is FEM problem and reduced_problem is RB problem

pod_galerkin_method_mechanical = PODGalerkin(hearth_problem_mechanical)

pod_galerkin_method_mechanical.set_Nmax(100) #Maximum size of reduced basis space

pod_galerkin_method_mechanical.set_tolerance(1e-4) #Maximum eigenvalue tolerance

# 5B. Perform the offline phase

pod_galerkin_method_mechanical.initialize_training_set(1000) #Initialize training set

with specified number of training parameters
```



Error analysis is performed and time taken for the computation of truth solution and reduced basis solution are measured.

```
# 7B. Perform an error analysis
pod_galerkin_method_mechanical.initialize_testing_set(50) #Initialize error analysis set
                                           with specified number of parameters
\verb"pod_galerkin_method_mechanical.error_analysis"() \textit{ \#Perform error analysis}
# 8B1. Perform a speedup analysis - Compute time for truth solutions
pod_galerkin_method_mechanical.initialize_testing_set(50) #Initialize speedup analysis
                                            set with specified number of parameters
testing_set_speedup_analysis = pod_galerkin_method_mechanical.testing_set
pod_galerkin_method_mechanical._patch_truth_solve(True) # To disable cache reading
truth_timer = Timer("parallel") #Timer for computation of FEM solution
taken for solving FEM equation. It is a
                                            vector of size of number of speedup analysis
                                             parameters
# Iteration over speedup analysis parameters for measuring time taken for FEM solution
for (mu_index, mu_test) in enumerate(testing_set_speedup_analysis):
 print (TextLine(str(mu_index), fill="#"))
 pod_galerkin_method_mechanical.truth_problem.set_mu(mu_test) #Set the parameter
 truth_timer.start()
 pod_galerkin_method_mechanical.truth_problem.solve() #Solve the FEM problem
 truth_time_mechanical = truth_timer.stop()
 print("Truth time mechanical : ",truth_time_mechanical)
 time_mechanical_truth[mu_index] = truth_time_mechanical #Save time taken for truth
pod_galerkin_method_mechanical._undo_patch_truth_solve(True) #To enable cache reading
np.save("time_mechanical_truth", time_mechanical_truth) #Save numpy array of time taken
                                            for FEM solution
# 8B2. Perform a speedup analysis - Compute time for reduced solutions
pod_galerkin_method_mechanical._patch_truth_solve(True) #To disable cache reading
reduced_timer = Timer("serial") #Timer for computation of reduced solution
max_basis_function = reduced_hearth_problem_mechanical.N # Size of reduced basis space
time_mechanical_reduced = np.empty([max_basis_function,len(testing_set_speedup_analysis)]
                                            ) #Storage of time taken for solving RB
                                            equation. It is a matrix of size size of
                                            reduced basis space \times number of speedup
                                             analysis parameters
# Iteration over speedup analysis parameters for measuring time taken for RB solution
for basis_size in range(1, max_basis_function+1):
  for (mu_index, mu_test) in enumerate(testing_set_speedup_analysis):
   print (TextLine(str(mu_index), fill="#"))
   pod_galerkin_method_mechanical.reduced_problem.set_mu(mu_test) #Set the parameter
   reduced_timer.start()
   pod_galerkin_method_mechanical.reduced_problem.solve(basis_size) #Solve the RB
                                            problem
   rb_time_mechanical = reduced_timer.stop()
   print("Reduced time mechanical: ",rb_time_mechanical)
```



For any new parameter *online_mu*, the truth solution and the reduced basis solution are computed as:

```
# 6B. Perform an online solve
pod_galerkin_method_mechanical.reduced_problem.set_mu(online_mu) #Set parameter
u_rb = pod_galerkin_method_mechanical.reduced_problem.solve() #Reduced problem solve
pod_galerkin_method_mechanical.reduced_problem.export_solution(filename="
                                             reference_domain_mechanical_rb") #Save
                                             solution for visualization with paraview
u_rb = pod_galerkin_method_mechanical.reduced_problem.basis_functions * u_rb #RB solution
                                              projected back to FEM space
pod_galerkin_method_mechanical.truth_problem.set_mu(online_mu) #Set parameter
u = pod_galerkin_method_mechanical.truth_problem.solve() #FEM problem solve
pod_galerkin_method_mechanical.truth_problem.export_solution(filename="
                                             reference_domain_fem") #Save solution for
                                             visualization with paraview
pod_galerkin_method_mechanical.truth_problem.mesh_motion.move_mesh() #Deform mesh as per
                                             geometric parameters
File("HearthMechanical/reference_domain_mechanical_spatial_error.pvd") << pre>project(u-u_rb,
                                             VM) #Spatial error
pod_galerkin_method_mechanical.truth_problem.mesh_motion.reset_reference() #Restore mesh
                                            to reference configuration
```

5.5.3 Coupling system

For the coupling system, we define the class *HearthThermoMechanical*.

```
class HearthThermoMechanical(EllipticCoerciveProblem):
```

The default initialization is specified,

```
# Default initialization of members
def __init__(self, V, **kwargs):
  # Call the standard initialization
 EllipticCoerciveProblem.__init__(self, V, **kwargs)
  # ... and also store FEniCS data structures for assembly
 assert "subdomains" in kwargs
 assert "boundaries" in kwargs
 assert "mesh" in kwargs
 assert "hearth_problem_thermal" in kwargs
 assert "ref_temperature" in kwargs
 self.subdomains, self.boundaries = kwargs["subdomains"], kwargs["boundaries"]
 self.u = TrialFunction(V)
 self.v = TestFunction(V)
 self.dx = Measure("dx")(subdomain_data=subdomains)
 self.ds = Measure("ds")(subdomain_data=boundaries)
 self.subdomains = subdomains
 self.boundaries = boundaries
  self.hearth_problem_thermal = kwargs["hearth_problem_thermal"]
  self.T_0 = kwargs["ref_temperature"]
  self.x0 = Expression("x[0]", element=V.sub(0).ufl_element())
```

Similar to mechanical system, we define the affine mutiplicative terms, weak formulation and strain tensor.



```
# Return theta multiplicative terms of the affine expansion of the problem.
def compute_theta(self, term):
   mu = self.mu
    if term == "a":
       theta_a0 = mu[11]
       theta_a1 = 2*mu[12]
       return (theta_a0, theta_a1,)
    elif term == "f":
       theta f0 = (2 * mu[11] + 3 * mu[12]) * mu[13]
       return (theta f0,)
        raise ValueError("Invalid term for compute_theta().")
# Return strain tensor
def strain(self,u):
   r = self.x0
   return sym(as_tensor([[u[0].dx(0), u[0].dx(1), 0.], [u[1].dx(0), u[1].dx(1), 0.], [0])
                                                                                         \cdot, 0., u[0]/r]]))
# Return forms resulting from the discretization of the affine expansion of the problem
                                                                                           operators.
def assemble_operator(self, term):
   u = self.u
    v = self.v
   dx = self.dx
   ds = self.ds
    T_0 = self.T_0
   T = self.hearth_problem_thermal._solution
    r = self.x0
   d_{bottom} = ds(2) + ds(3) + ds(4) + ds(5) + ds(6)
    d_{out} = ds(7) + ds(8) + ds(9) + ds(10) + ds(11)
    d_sf = ds(13) + ds(14) + ds(15) + ds(16) + ds(17) + ds(18) + ds(19) + ds(20)
   if term == "a":
       a0 = (u[0].dx(0)+u[1].dx(1)+u[0]/r)*(v[0].dx(0)+v[1].dx(1)+v[0]/r)*r*dx
       a1 = (u[0].dx(0)*v[0].dx(0) + u[1].dx(1)*v[1].dx(1) + (u[0]*v[0])/(r)**2 + 0.5*(u[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0]*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v[0])**(u[0)*v
                                                                                         ].dx(1)+u[1].dx(0))*(v[0].dx(1)+v[1].dx(0)))
                                                                                           * r * dx
        return (a0, a1,)
    elif term == "f":
        f0 = (T-T_0) * (v[0].dx(0) + v[1].dx(1) + v[0]/r) * r * dx
        return (f0,)
    elif term == "inner_product":
        x0 = inner(u, v) * r * dx + inner(self.strain(u), self.strain(v)) * r * dx
        return (x0,)
    elif term == "dirichlet_bc":
       bc0 = [DirichletBC(self.V.sub(0), Constant(0.), self.boundaries, 1),
            DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 2),
            DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 3),
           DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 4),
           DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 5),
           DirichletBC(self.V.sub(1), Constant(0.), self.boundaries, 6),]
        return (bc0,)
    else:
        raise ValueError("Invalid term for assemble_operator().")
```

We define an instance of *HearthThermoMechanical* class and initialise reduction with POD-Galerkin method.



```
hearth_problem_thermal, ref_temperature=T_0)
#specify and set range of each parameter

mu_range = [(2.3,2.4), (0.5,0.7), (0.5,0.7), (0.4,0.6), (3.05,3.35), (13.5,14.5), (8.3,8.7), (8.8,9.2), (9.8,10.2), (10.4,10.8), (9.8,10.2), (1.9e9,2.5e9), (1.2e9,1.8e9), (0.8e-6,1.2e-6)]

hearth_problem_thermo_mechanical.set_mu_range(mu_range)

# 4C. Prepare reduction with a POD-Galerkin method
#NOTE: truth_problem_attribute is FEM problem and reduced_problem is RB problem
pod_galerkin_method_thermo_mechanical = PODGalerkin(hearth_problem_thermo_mechanical)
pod_galerkin_method_thermo_mechanical.set_Nmax(100) #Maximum size of reduced basis space
pod_galerkin_method_thermo_mechanical.set_tolerance(1e-4) #Maximum eigenvalue tolerance
```

Next, we perform the offline phase.

Before performing the reduction of thermal problem, we perform the truth solution computation related operations. This is due to the reason that, we want to use temperature field computed by finite element method for the truth solution of displacement. We compute the truth solution at few parameters to perform error analysis and to measure time taken for computing the truth solution.

```
# 6C. Perform a truth solve : Reference domain
online_mu = (2.365, 0.6, 0.6, 0.5, 3.2, 14.10, 8.50, 9.2, 9.9, 10.6, 10., lame1, lame2,
                                             1e-6)
pod_galerkin_method_thermo_mechanical.truth_problem.set_mu(online_mu)
u_ref = pod_galerkin_method_thermo_mechanical.truth_problem.solve()
pod_galerkin_method_thermo_mechanical.truth_problem.export_solution(filename="
                                             reference_domain_fem")
# 6C. Perform a truth solve : Parametrized domain
online_mu = (2.365, 0.6, 0.6, 0.45, 3.2, 14.10, 8.30, 9.2, 9.9, 10.6, 10., lame1, lame2,
                                              1e-6
pod_galerkin_method_thermo_mechanical.truth_problem.set_mu(online_mu)
u_par = pod_galerkin_method_thermo_mechanical.truth_problem.solve()
pod_galerkin_method_thermo_mechanical.truth_problem.export_solution(filename="
                                             parametric_domain_fem")
# 7C1. Perform an error analysis - Compute truth solutions
pod_galerkin_method_thermo_mechanical.initialize_testing_set(50) #Initialize error
                                             analysis set with specified number of
                                             parameters
testing_set_error_analysis = pod_galerkin_method_thermo_mechanical.testing_set
truth_solution_thermo_mechanical = list()
# Iteration over error analysis parameters for measuring time taken for FEM solution
for (mu_index, mu_test) in enumerate(testing_set_error_analysis):
 print (TextLine(str(mu_index), fill="#"))
 pod_galerkin_method_thermo_mechanical.truth_problem.set_mu(mu_test) #Set parameter
 truth_solution_thermo_mechanical.append(pod_galerkin_method_thermo_mechanical.
                                             truth_problem.solve()) #Solve and store FEM
                                             solution
# 8C1. Perform a speedup analysis - Compute time for truth solutions
```



```
pod_galerkin_method_thermo_mechanical.initialize_testing_set(50) #Initialize truth
                                          solution with specified number of parameters
testing_set_speedup_analysis = pod_galerkin_method_thermo_mechanical.testing_set
pod_galerkin_method_thermo_mechanical._patch_truth_solve(True) #To disable cache reading
truth_timer = Timer("parallel") #Timer for computation of FEM solution
time_thermo_mechanical_truth = np.empty(len(testing_set_speedup_analysis)) #Storage of
                                          time taken for solving FEM equation. It is a
                                           vector of size of number of speedup
                                          analysis parameters
# Iteration over speedup analysis parameters for measuring time taken for RB solution
for (mu_index, mu_test) in enumerate(testing_set_speedup_analysis):
 print(TextLine(str(mu_index), fill="#"))
 pod_galerkin_method_thermo_mechanical.truth_problem.set_mu(mu_test) #Set the parameter
 truth_timer.start()
 pod_galerkin_method_thermo_mechanical.truth_problem.solve() #Solve the RB problem
 truth_time_thermo_mechanical = truth_timer.stop()
 print("Truth time thermomechanical : ",truth_time_thermo_mechanical)
 for reduced basis solution
pod_galerkin_method_thermo_mechanical._undo_patch_truth_solve(True) #To disable cache
                                          reading
np.save("time_thermo_mechanical_truth",time_thermo_mechanical_truth) #Save numpy array of
                                           time taken for RB solution
```

Now, since the operations related to the truth solution are performed, we can reduce the thermal system. The reduced basis solution of temperature field is used for computing reduced solution at few parameters. Also we compute the reduced basis solution for error analysis and time taken for computing the reduced basis solution.

```
#6C. Perform an online solve : Reference domain
online_mu_reference = ( 2.365, 0.6, 0.6, 0.5, 3.2, 14.10, 8.50, 9.2, 9.9, 10.6, 10.,
                                             lame1, lame2, 1e-6)
online_mu = online_mu_reference
pod_galerkin_method_thermo_mechanical.reduced_problem.set_mu(online_mu)
u_rb = pod_galerkin_method_thermo_mechanical.reduced_problem.solve()
pod_galerkin_method_thermo_mechanical.reduced_problem.export_solution(filename="
                                             reference_domain_thermomechanical_rb")
\verb"u"rb" = \verb"pod_galerkin_method_thermo_mechanical.reduced_problem.basis_functions * u"rb"
pod_galerkin_method_thermo_mechanical.truth_problem.mesh_motion.move_mesh()
File("HearthThermoMechanical/reference_domain_thermomechanical_spatial_error.pvd") <<
                                             project(u_ref-u_rb,VM)
pod_galerkin_method_thermo_mechanical.truth_problem.mesh_motion.reset_reference()
# 6C. Perform an online solve : Parametrized domain
online_mu_parametrized = (2.365, 0.6, 0.6, 0.45, 3.2, 14.10, 8.30, 9.2, 9.9, 10.6, 10.,
                                             lame1, lame2, 1e-6)
online_mu = online_mu_parametrized
pod_galerkin_method_thermo_mechanical.reduced_problem.set_mu(online_mu)
u_rb = pod_galerkin_method_thermo_mechanical.reduced_problem.solve()
pod_galerkin_method_thermo_mechanical.reduced_problem.export_solution(filename="
                                             parametric_domain_thermomechanical_rb")
u_rb = pod_galerkin_method_thermo_mechanical.reduced_problem.basis_functions * u_rb
pod_galerkin_method_thermo_mechanical.truth_problem.mesh_motion.move_mesh()
File("HearthThermoMechanical/parametric_domain_thermomechanical_spatial_error.pvd") <<
                                             project(u_par-u_rb,VM)
\verb|pod_galerkin_method_thermo_mechanical.truth_problem.mesh_motion.reset_reference()|
```



```
# 7C2. Perform an error analysis - Compute reduced basis solution
dx = Measure("dx")(subdomain_data=subdomains) #Volume measure
r = Expression("x[0]", element=VM.sub(0).ufl_element()) #
max_basis_function = reduced_hearth_problem_thermo_mechanical.N # Size of reduced basis
                                             space
error_thermo_mechanical = np.empty([max_basis_function,len(testing_set_error_analysis)])
                                             # Numpy array of size of reduced basis space
                                              \times number of error analysis parameters
                                             for storing error
# Iteration over error analysis parameters for measuring time taken for RB solution
for basis_size in range(1, max_basis_function+1):
  for (mu_index, mu_test) in enumerate(testing_set_error_analysis):
    print(TextLine(str(mu_index), fill="#"))
    pod_galerkin_method_thermo_mechanical.reduced_problem.set_mu(mu_test) #Set parameter
    rb_dofs = pod_galerkin_method_thermo_mechanical.reduced_problem.solve(basis_size) #
                                             Compute reduced basis degrees of freddom
    rb_solution = reduced_hearth_problem_thermo_mechanical.basis_functions[:basis_size] *
                                              rb_dofs #RB solution projected back to FEM
                                             space
    # Absolute and relative error measurement
    absolute_error = assemble(inner(truth_solution_thermo_mechanical[mu_index] -
                                             rb_solution,truth_solution_thermo_mechanical
                                             [mu_index] - rb_solution) * r * dx + inner(
                                             hearth_problem_thermo_mechanical.strain(
                                             truth_solution_thermo_mechanical[mu_index] -
                                              rb_solution),
                                             hearth_problem_thermo_mechanical.strain(
                                             truth_solution_thermo_mechanical[mu_index] -
                                              rb_solution)) * r * dx)
    error_thermo_mechanical[basis_size-1,mu_index] = np.sqrt(absolute_error / assemble(
                                             inner(truth_solution_thermo_mechanical[
                                             mu_index],truth_solution_thermo_mechanical[
                                             mu_index]) * r * dx + inner(
                                             hearth_problem_thermo_mechanical.strain(
                                             truth_solution_thermo_mechanical[mu_index]),
                                              hearth_problem_thermo_mechanical.strain(
                                             truth_solution_thermo_mechanical[mu_index]))
                                              \star r \star dx))
np.save("HearthThermoMechanical/error_analysis/error_thermo_mechanical",
                                             error_thermo_mechanical)
# 8C2. Perform a speedup analysis - Compute time for reduced solutions
pod_galerkin_method_thermo_mechanical._patch_truth_solve(True) #To disable cache reading
reduced_timer = Timer("serial") #Timer for computation of RB solution
time_thermo_mechanical_reduced = np.empty([max_basis_function,len(
                                             testing_set_speedup_analysis)]) #Storage of
                                             time taken for solving RB equation. It is a
                                             matrix of size size of reduced basis space \
                                             times number of speedup analysis parameters
# Iteration over speedup analysis parameters for measuring time taken for RB solution
for basis_size in range(1, max_basis_function+1):
  for (mu_index, mu_test) in enumerate(testing_set_speedup_analysis):
    print (TextLine(str(mu_index), fill="#"))
    pod_galerkin_method_thermo_mechanical.reduced_problem.set_mu(mu_test) #Set parameter
    reduced timer.start()
    pod_galerkin_method_thermo_mechanical.reduced_problem.solve(basis_size) #Solve the RB
                                              problem
```



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