



Tutorial - Writing Advanced Components

Introduction

This tutorial is a continuation of the *Writing Component Libraries* tutorial. It explains how to write more advanced components based on acausal equations and physical connections. It also covers how to re-write the equations for the impedance variables, which are a consequence of the transmission line element method.

Two components are explained; a variable pump component with linear equations, and a translational mass component with second order dynamics.

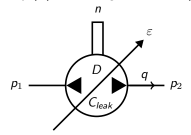
Requirements

It is necessary to know how to write and compile components for Hopsan, either using the built-in editor in Hopsan or a third-party tool. This is covered by the *Writing Component Libraries* tutorial.

Hydraulic Pump

A hydraulic pump is a component that transforms an angular velocity on a shaft into hydraulic flow. In this case we will use a simple model with no dynamics, where the angular velocity is assumed to be constant at all time. This is similar to the Q-Type Variable Displacement Pump component in the Hopsan default library.

The component is shown in the figure below. It consists of two hydraulic ports for inlet and outlet, respectively. It also has four input variables; displacement (D), displacement setting (ε), angular velocity (ω) and leakage coefficient (c_{leak}). All of these



Equations

The flow from a hydraulic pump can be modelled with the following equations:

$$\left\{ egin{aligned} q - q_{ ext{leak}} &= arepsilon D\omega/2\pi \ q_{ ext{leak}} &= C_{ ext{leak}}\Delta p \end{aligned}
ight.$$

Here q is the generated flow and q_{leak} the leakage flow. Δp is the pressure difference over the pump. In Hopsan we will have one flow and pressure at each port. Thus, we need to introduce the variables p_1 and q_1 for the inlet port, and p_2 and p_3 for the outlet. Flow is always defined as positive outwards from Q-type components. Therefore, the inlet flow will be the same as the outlet flow but negative. The equations now become:



$$\left\{egin{aligned} q_2 - \mathit{C}_{\mathit{leak}}(\mathit{p}_1 - \mathit{p}_2) = arepsilon \mathit{D}\omega/2\pi \ q_1 = -\mathit{q}_2 \end{aligned}
ight.$$

However, Hopsan uses the transmission line element method. For this reason we will not get the pressure variables as explicit input variables. Instead, we get a *wave variable* and an *impedance*. The TLM equations are then used to calculate the pressures from the flow variables, according to the two new equations below:

$$\left\{egin{aligned} q_2 - \mathcal{C}_{\textit{leak}}(p_1 - p_2) &= arepsilon D\omega/2\pi \ q_1 &= -q_2 \ p_1 &= c_1 + q_1 Z_{c,1} \ p_2 &= c_2 + q_2 Z_{c,2} \end{aligned}
ight.$$

It is now possible to rearrange the equations so that the can be solved analytically. First we replace the pressure variables in the first equation with the TLM equations:

$$\begin{cases} q_2 - \textit{C}_{\textit{leak}}(\textit{c}_1 - \textit{q}_2 \textit{Z}_{\textit{c},1} - \textit{c}_2 - \textit{q}_2 \textit{Z}_{\textit{c},2}) = \varepsilon \textit{D}\omega/2\pi \\ q_1 = -q_2 \\ p_1 = \textit{c}_1 + \textit{q}_1 \textit{Z}_{\textit{c},1} \\ p_2 = \textit{c}_2 + \textit{q}_2 \textit{Z}_{\textit{c},2} \end{cases}$$

Finally, we rearrange the first equation to break out the flow variable:

$$\begin{cases} q_2 = (\varepsilon D\omega/2\pi - \textit{C}_{\textit{leak}}(\textit{c}_1 - \textit{c}_2))/(1 - \textit{C}_{\textit{leak}}(\textit{Z}_{\textit{c},1} - \textit{Z}_{\textit{c},2})) \\ q_1 = -q_2 \\ p_1 = \textit{c}_1 + q_1 \textit{Z}_{\textit{c},1} \\ p_2 = \textit{c}_2 + q_2 \textit{Z}_{\textit{c},2} \end{cases}$$

We now have a linear equation system that can be solved step-by-step.

C++ Code

First, we create the skeleton for the new component, as described in the previous tutorial. We need two hydraulic ports and four input variables (ω , D_p , C_{leak} and ε). The resulting code for the class members and the creator functions is shown below:

```
#include "ComponentEssentials.h"
3
   namespace hopsan {
5
   class HydraulicFixedDisplacementPump : public ComponentQ
6
7
   private:
        double *mpND_p1, *mpND_q1, *mpND_c1, *mpND_Zc1;
9
        double *mpND_p2, *mpND_q2, *mpND_c2, *mpND_Zc2;
        double *mpW, *mpDp, *mpCleak, *mpEps;
10
11
       Port *mpP1, *mpP2;
12
13
       static Component *Creator()
14
15
            return new HydraulicFixedDisplacementPump();
16
17
```



The only thing needed in the configuration() function is to create ports and input variables:

```
void configure()
1
2
         mpP1 = addPowerPort("P1", "NodeHydraulic");
3
         mpP2 = addPowerPort("P2", "NodeHydraulic");
4
5
              addInputVariable("eps", "Displacement setting", "", 1.0, &mpEps);
6
         addInputVariable("w_p", "Angular Velocity", "rad/s", 250.0, &mpW); addInputVariable("D_p", "Displacement", "m^3/rev", 0.00005, &mpDp);
7
8
         addInputVariable("C_leak", "Leakage Coeff.", "(m^3/s)/Pa", 0.0, &mpCleak);
9
10
```

The initialize() function will only need the getSafeDataNodePtr() function calls:

```
void initialize()
1
2
3
        mpND_p1 = getSafeNodeDataPtr(mpP1, NodeHydraulic::Pressure);
        mpND_q1 = getSafeNodeDataPtr(mpP1, NodeHydraulic::Flow);
mpND_c1 = getSafeNodeDataPtr(mpP1, NodeHydraulic::WaveVariable);
4
5
        mpND_Zc1 = getSafeNodeDataPtr(mpP1, NodeHydraulic::CharImpedance);
6
        mpND_p2 = getSafeNodeDataPtr(mpP2, NodeHydraulic::Pressure);
8
        mpND_q2 = getSafeNodeDataPtr(mpP2, NodeHydraulic::Flow);
9
10
        mpND_c2 = getSafeNodeDataPtr(mpP2, NodeHydraulic::WaveVariable);
        mpND_Zc2 = getSafeNodeDataPtr(mpP2, NodeHydraulic::CharImpedance);
11
12
```

Now it is time for the interesting part, to write the actual equations in the simulateOneTimestep() function. The final code is presented below. Some of the steps below will already be done if you generate the code from the Hopsan graphical interface.

1. Create local variables

We begin with creating local variables for the port variables (line 3-4 below).

2. Read input variables

Assign all input variables them with their corresponding pointers (line 6-16).

3. Write the pump equations

The equations must be converted to C++ syntax (line 18-22).

4. Check for cavitation

In order to make the model physically correct, we must make sure that negative pressures are not allowed. For this we create a boolean variable for cavitation and assign it with false (line 25). Then we check if pressure p_1 or p_2 is smaller than zero. If they are, we set the cavitation boolean to true and assign c_1 and c_1 with zero (line 26-37).

5. Handle the cavitation

If the cavitation boolean is true, at least one of the pressures was negative. In this case we need to recalculate the flow and the pressure variables (line 39-46).

6. Write output variables

Finally, the output variables must be written to the nodes (line 48-52).



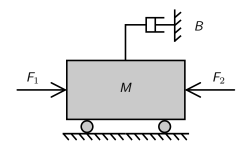
```
void simulateOneTimestep()
2
        //Declare local variables
3
        double p1, q1, c1, Zc1, p2, q2, c2, Zc2, w, dp, Cleak, eps;
4
5
        //Read input variables
6
7
        w = (*mpW);
        dp = (*mpDp);
8
        Cleak = (*mpCleak);
9
10
        eps = (*mpEps);
11
12
        //Get variable values from nodes
13
        c1 = (*mpND_c1);
        Zc1 = (*mpND_Zc1);
14
        c2 = (*mpND_c2);
15
        Zc2 = (*mpND_Zc2);
16
17
        //Fixed Displacement Pump equations
18
        q2 = (dp*eps*w/(2.0*pi) + Cleak*(c1-c2)) / ((Zc1+Zc2)*Cleak+1);
19
20
        q1 = -q2;
        p1 = c1 + Zc1*q1;
21
        p2 = c2 + Zc2*q2;
22
23
        //Check for cavitation
24
25
        bool cav = false;
26
        if (p1 < 0.0)
27
        {
28
            c1 = 0.0;
            Zc1 = 0.0;
cav = true;
29
30
31
        if (p2 < 0.0)
32
33
34
            c2 = 0.0;
            Zc2 = 0.0;
35
36
            cav = true;
37
38
39
        //Handle cavitation
40
        if (cav)
41
42
            q2 = (dp*eps*w/(2.0*pi) + Cleak*(c1-c2)) / ((Zc1+Zc2)*Cleak+1);
            q1 = -q2;

p1 = c1 + Zc1 * q1;
43
44
            p2 = c2 + Zc2 * q2;
45
46
47
        //Write new values to nodes
48
        (*mpND_p1) = p1;
49
        (*mpND_q1) = q1;
        (*mpND_p2) = p2;
51
52
        (*mpND_q2) = q2;
53 }
```



Translational Mass

This part of the tutorial will explain how to write a translational mass component with second order dynamics. It will contain a mass and a damping coefficient, as shown in the figure below. We leave out dry friction and spring coefficient for simplicity.



Equations

The fundamental equation in the component will be Newton's second law of motion:

$$M\ddot{x} + B\dot{x} = \sum F$$

The Hopsan component will have two ports, which will be defined in opposite directions. It is also necessary to provide equivalent mass variables, which are required by some other components in Hopsan:

$$\begin{cases} M\ddot{x_2} + B\dot{x_2} = F_1 - F_2 \\ x_1 = -x_2 \\ \dot{x}_1 = -\dot{x}_2 \\ m_{e,1} = M \\ m_{e,2} = M \end{cases}$$

Finally, we must add the TLM equations in order to calculate the force from the wave variables and impedances:

$$\begin{cases} M\ddot{x_2} + B\dot{x_2} = F_1 - F_2 \\ x_1 = -x_2 \\ \dot{x}_1 = -\dot{x}_2 \\ m_{e,1} = M \\ m_{e,2} = M \\ F_1 = c_1 + Z_{c,1}\dot{x}_1 \\ F_2 = c_2 + Z_{c,2}\dot{x}_2 \end{cases}$$

The equations of the mass is more complicated than the ones for the pump, since it contains time derivatives (\ddot{x} and \dot{x}). Hopsan uses bilinear transforms to solve such equations. In order to use these, we must convert the first equation into a Laplace transform. We use a first order transform to calculate the velocity. Then the position can be obtained by integrating the velocity. By adding the impedance directly to the damping, we can use the wave variables as the external force. In this way we can calculate x and v without first calculating the forces.



$$\begin{cases} v_2 = \frac{1}{Ms + B}(F_1 - F_2) = \frac{1}{Ms + (B + Z_{c,1} + Z_{c,2})}(c_1 - c_2) \\ \dot{x}_2 = v_2 \\ v_1 = -v_2 \\ x_1 = -x_2 \\ F_1 = c_1 + Z_{c,1}v_1 \\ F_2 = c_2 + Z_{c,2}v_2 \end{cases}$$

C++ Code

As with the previous component, we begin by creating a code skeleton. We want two mechanical ports, a constant parameter for the mass and input variables for damping and minimum and maximum position. Note that we also need to include the file ComponentUtilities.h, which contain classes for the transfer function and the integration.

```
#include "ComponentEssentials.h"
   #include "ComponentUtilities.h"
3
   namespace hopsan {
6
   {\tt class} \ {\tt MechanicTranslationalMass} \ : \ {\tt public} \ {\tt ComponentQ}
7
   private:
8
9
        Port *mpP1, *mpP2;
10
        double *mpP1_f, *mpP1_x, *mpP1_v, *mpP1_c, *mpP1_Zc, *mpP1_me;
        double *mpP2_f, *mpP2_x, *mpP2_v, *mpP2_c, *mpP2_Zc, *mpP2_me;
11
12
        double *mpB, *mpXMin, *mpXMax;
13
        double mMass;
14
15
        FirstOrderTransferFunction mTF;
16
        Integrator mIntegrator;
17
        double mNum[2], mDen[2];
18
   public:
19
20
        static Component *Creator()
21
        {
             return new MechanicTranslationalMass();
22
```

There is nothing special with the configure() function. All we need to do here is to create the ports, the constant and the input variables.

```
void configure()
1
        mpP1 = addPowerPort("P1", "NodeMechanic");
mpP2 = addPowerPort("P2", "NodeMechanic");
3
4
6
         addConstant("m",
                                        "Mass",
                                                                  "kg",
                                                                           100.0, mMass);
8
         addInputVariable("B",
                                       "Viscous Friction",
                                                                  "Ns/m", 10.0, &mpB);
9
         addInputVariable("x_min", "Minimum Position",
                                                                  "m",
                                                                           0.0,
                                                                                    &mpXMin);
10
         addInputVariable("x_max", "Maximum Position",
                                                                            1.0,
                                                                                    &mpXMax);
11
```

The initialize() function is a little more advanced than for the pump. We need to initialize the transfer function and the integrator with start values for f, v and x. We also need to define the coefficients for the transfer function. This is done with two arrays called mDen and mNum, according to this:

```
\frac{mNum[1]s + mNum[0]}{mDen[1]s + mNum[0]}
```



Finally, we also need to initialize the equivalent mass variables in the nodes.

```
void initialize()
1
2
3
          mpP1_f = getSafeNodeDataPtr(mpP1, NodeMechanic::Force);
         mpP1_x = getSafeNodeDataPtr(mpP1, NodeMechanic::Position);
mpP1_v = getSafeNodeDataPtr(mpP1, NodeMechanic::Velocity);
4
5
          mpP1_c = getSafeNodeDataPtr(mpP1, NodeMechanic::WaveVariable);
6
         mpP1_Zc = getSafeNodeDataPtr(mpP1, NodeMechanic::CharImpedance);
mpP1_me = getSafeNodeDataPtr(mpP1, NodeMechanic::EquivalentMass);
7
8
9
10
          mpP2_f = getSafeNodeDataPtr(mpP2, NodeMechanic::Force);
         mpP2_x = getSafeNodeDataPtr(mpP2, NodeMechanic::Position);
mpP2_v = getSafeNodeDataPtr(mpP2, NodeMechanic::Velocity);
11
12
          mpP2_c = getSafeNodeDataPtr(mpP2, NodeMechanic::WaveVariable);
13
         mpP2_Zc = getSafeNodeDataPtr(mpP2, NodeMechanic::CharImpedance);
mpP2_me = getSafeNodeDataPtr(mpP2, NodeMechanic::EquivalentMass);
14
15
16
17
          //Read node variables
18
          double f1, f2, x2, v2;
19
          f1 = (*mpP1_f);
20
          f2 = (*mpP2_f);
21
          x2 = (*mpP2_x);
          v2 = (*mpP2_v);
22
23
24
          //Initialization code
25
          mNum[0] = 1.0;
          mNum[1] = 0.0;
26
27
          mDen[0] = (*mpB);
          mDen[1] = mMass;
28
29
          mTF.initialize(mTimestep, mNum, mDen, f1-f2, v2);
30
31
          mIntegrator.initialize(mTimestep, x2, v2);
33
          //Write values to nodes
34
          (*mpP1_me) = mMass;
          (*mpP2_me) = mMass;
35
36
```

Last but not least, we need to write the simulateOneTimeStep() function. This is explained step-by-step:

1. Define local variables

Define local variables for input variables and node values (line 3-4)

2. Read from input variables

Assign input variables to local variables (line 6-9)

3. Read from nodes

Assign node variables to local variables (line 11-15)

4. Update the damping coefficient

Update the coefficient in the transfer function by adding the impedance variables to the damping factor (line 17-19)

5. Calculate velocity

Use the transfer function to calculate v_2 from c_1-c_2 (line 21-22)

6. Calculate position

Use the integrator to calculate x_2 as a function of v_2 (line 24-25)

7. Handle limitations

Reset variables and re-initialize the transfer function and the integrator if x_2 is outside limits (line 27-41)



8. Calculate the remaining variables

Calulate velocity and position for node 1 (v_1, x_1) , and then calculate, F_1 and F_2 using the TLM equations (line 43-47)

9. Write values to the nodes

Write back the values of the local variables to the node variables (line 49-57)

```
void simulateOneTimestep()
2
3
        //Declare local variables
        double f1, x1, v1, c1, Zc1, f2, x2, v2, c2, Zc2, B, xmin, xmax;
4
5
6
        //Read input variables
        B = (*mpB);
7
        xmin = (*mpXMin);
        xmax = (*mpXMax);
9
10
11
        //Get variable values from nodes
        c1 = (*mpP1_c);
12
        Zc1 = (*mpP1_Zc);
13
        c2 = (*mpP2_c);
14
15
        Zc2 = (*mpP2_Zc);
16
        //Mass equations
17
        mDen[0] = B+Zc1+Zc2;
18
19
        mTF.setDen(mDen);
20
21
        //Calculate velocity
        v2 = mTF.update(c1-c2);
22
23
24
        //Calculate position
25
        x2 = mIntegrator.update(v2);
26
27
        //Handle position limits
        if(x2<xmin)
28
29
            x2=xmin;
30
            v2=0.0;
31
32
            mTF.initializeValues(c1-c2, v2);
            mIntegrator.initializeValues(v2, xmin);
33
34
        }
35
        if(x2>xmax)
36
37
            x2=xmax;
38
            v2=0.0;
39
            mTF.initializeValues(c1-c2, v2);
40
            mIntegrator.initializeValues(v2, xmax);
41
42
43
        //Calculate remainig varaibles
44
        v1 = -v2;
45
        x1 = -x2;
        f1 = c1 + Zc1*v1;
46
47
        f2 = c2 + Zc2*v2;
48
        //Write new values to nodes
49
50
        (*mpP1_f) = f1;
        (*mpP1_x) = x1;
51
        (*mpP1_v) = v1;
52
53
        (*mpP2_f) = f2;
        (*mpP2_x) = x2;
54
        (*mpP2_v) = v2;
55
        (*mpP1_me) = mMass;
56
        (*mpP2_me) = mMass;
57
  }
58
```