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Hierarchical Non-Linear Bond Graphs: A Unified Methodology for Modeling Complex Physical Systems

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Abstract
Bond graphs have been around for a quarter of a century. While originally intended for modeling mechanical systems, they have meanwhile found widespread applications in many areas of physical system modeling. However, bond graphs, like all graphical techniques, become unwieldy when applied to complex systems. Also, bond graphs were traditionally used to model predominantly linear systems. This talk introduces a new concept for modeling complex physical systems through hierarchical bond graphs which can include arbitrary non-linearities. It introduces a software tool that can be used to implement these hierarchical non-linear bond graphs. Finally, a new application area for bond graphs will be discussed. It will be demonstrated how these hierarchical non-linear bond graphs can be used to model chemical reaction kinetics and chemical thermodynamics together in very general terms also farther away from equilibrium than traditional approaches would permit.

Introduction
It has been known for a long time that the mathematics behind the dynamical behavior of systems from different disciplines of physical sciences have much in common. In particular, it has been recognised that mechanical systems can be treated by making use of electro–mechanical analogies, i.e. by treating masses, springs, and frictions as inductors, capacitors, and resistors.

This methodology was formalised in the early sixties through the introduction of so-called bond graphs (Paynter 1961). A bond is a directed path that denotes the flow of power from one point of the system to another.

It turned out that, in all physical systems, power can be written as a product of two variables, one of which is an across variable, while the other is a through variable. Across variables around a node assume the same value, whereas through variables into a “node” add up to zero. The bond graph name of a node is a 0-junction. In an electrical circuit, the potentials around a node assume the same value, whereas the currents into a node add up to zero. Thus, if we identify the node of the electrical circuit with the 0-junction of the bond graph, the potentials will become across variables, while the currents will become through variables.

The electrical power flowing through a circuit element can be expressed as the product of the potential drop \( V \), and the current \( I \):

\[ P_{el} = V \cdot I \]

In bond graph terminology, across variables are called effort variables, while through variables are called flow variables. The symbol \( e \) is used to denote an effort, and the symbol \( f \) is used to denote a flow. A bond is symbolized by a harpoon with the hook pointing to the left. The effort is placed at the side of the hook, while the flow is shown on the opposite side.

The denomination of the across and through variables is arbitrary. If we had identified an electrical mesh with the 0-junction instead of the electrical node, the roles of the potentials and currents in the bond graph would have been interchanged, yet the power would still be the product of one across variable and one through variable.

Since electrical circuits, and all other physical systems, are described through a set of node (cutset) equations and a set of mesh (loop) equations, the bond graph must contain two types of junctions. The effort variables across a 0-junction assume the same value, whereas the flow variables into the 0-junction add up to zero. The flow variables across a 1-junction assume the same value, whereas the effort variables into the 1-junction add up to zero.

We are now ready to apply these concepts to a simple RLC-circuit:

The bond graph for this circuit is shown below:

\[ L_1 = 1.5 \text{ mH} \]
\[ R_1 = 100 \Omega \]
\[ R_2 = 20 \Omega \]
\[ C_1 = 0.1 \mu \text{F} \]

\[ V_0 = 10 \text{ V} \]

\[ I_1 \]
\[ I_L \]
\[ i_{L1} \]
\[ i_{L0} \]
\[ v_{R1} \]
\[ v_{R2} \]
\[ v_0 \]
\[ i_{C1} \]

\[ \text{SE: } V_0 = 10 \text{ V} \]
\[ \text{L: } L_1 = 1.5 \text{ mH} \]
\[ \text{R: } R_1 = 100 \Omega \]
\[ \text{R: } R_2 = 20 \Omega \]

\[ \text{C: } C_1 = 0.1 \mu \text{F} \]

\[ \text{SE: } V_0 = 10 \text{ V} \]

\[ \text{L: } L_1 = 1.5 \text{ mH} \]
\[ \text{R: } R_1 = 100 \Omega \]
\[ \text{R: } R_2 = 20 \Omega \]

\[ \text{C: } C_1 = 0.1 \mu \text{F} \]

\[ \text{SE: } V_0 = 10 \text{ V} \]
The rules for constructing a bond graph of an electrical circuit are very simple. We start by representing each circuit node by a 0-junction except for the reference node which is drawn like in the circuit diagram. We then represent each branch of the circuit diagram by a pair of bonds connecting two 0-junctions with a 1-junction between them. We let the harpoons point in the same direction that we picked for the branch currents. Finally, we attach the network elements to the 1-junctions with the harpoons away from the junction for passive network elements, and directed towards the junction for sources.

Contrary to other graphical abstractions such as block diagrams and signal flow graphs, the bond graph obviously preserves the geometric topology of the physical system. What destroys the topology in a block diagram or a signal flow graph is the fact that voltages and currents that participate in a power flow get separated from each other. However, efforts and flows aren’t tradable goods in a physical system. The only two types of merchandise that exist in a physical system are power and mass. The bond graph preserves the structural topology of the physical system since it reflects physical trades.

Since the potential of the reference node $v_0$ is normalized to zero, the power into and out of the reference node is zero, and consequently, we can eliminate the power bonds that connect the circuit to the reference node. In our example, this leads to a number of junctions with only two bonds attached to them. Such junctions can be eliminated. The two attached bonds are thereby amalgamated into one. The resulting simplified bond graph for the above example is shown below:

$$I : L_1$$
$$R : R_1$$
$$= 1.5 \text{ mH}$$
$$= 100 \Omega$$

$$\overrightarrow{v_0} \overleftarrow{i_0} \overrightarrow{v_1} \overleftarrow{i_1} \overrightarrow{v_2} \overleftarrow{i_2} \overrightarrow{v_3} \overleftarrow{i_3}$$

$\overrightarrow{u_R} \overleftarrow{i_R} \overrightarrow{u_C} \overleftarrow{i_C} \overrightarrow{u_I} \overleftarrow{i_I}$

The bond graph shows the dissipation of power from the effort source $SE$ into the various passive circuit elements of type resistance $R$, capacitance (or compliance $C$, and inductance (or iner- tance $I$).

If it should happen that ever there are two junctions of the same sex adjacent to each other and connected by a simple bond, the two junctions can be merged into one. Thus, 0-junctions and 1-junctions always toggle in any bond graph model.

Two variables are associated with each of the bonds. Consequently, we need two equations in the resulting simulation program to evaluate them. It turns out that, in all bond graphs, one of these variables is always evaluated at each of the two ends of the bond. We can denote this fact by a little vertical stroke attached to one of the two ends of the bond. By convention, the stroke denotes the end where the flow variable is evaluated. In bond graph terms, this is called assigning a (computational) causality to the bond graph.

Sources have mandated causalities as shown below:

$$\overrightarrow{u_C} \overleftarrow{i_C} \overrightarrow{u_I} \overleftarrow{i_I}$$

Since the effort of an effort source is determined at the source, the flow must be determined at the other end, thus, the little stroke is away from the source. In the flow source, the situation is opposite. The thin arrows denote signal paths. They symbolize the fact that these variables are somehow determined from outside the system. A source as drawn in the circuit example above is actually a non-physical element. Power cannot be generated, only transported and converted. However, a "system" never denotes the whole of the universe. It denotes a piece of the universe. Sources are interfaces between the system and the universe around it.

Capacitances and inductances have recommended causalities as shown below:

$$\overrightarrow{u_C} \overleftarrow{i_C} \overrightarrow{u_I} \overleftarrow{i_I}$$

By declaring that we wish to compute the effort at the capacitance, we generate an equation of the type:

$$\frac{du_C}{dt} = \frac{1}{C} \cdot i_C$$

i.e., we end up with a state equation in the state variable $u_C$ which can be integrated. The reverse causality would have produced an integral equation for the variable $i_C$ which would have forced us to compute the variable $u_C$ somewhere else in the circuit by means of numerical differentiation.

The resistance has an arbitrary causality:

$$\overrightarrow{u_R} \overleftarrow{i_R} \overrightarrow{u_R} \overleftarrow{i_R}$$

The causality shown to the left leads to the equation:

$$i_R = \frac{1}{R} \cdot u_R$$

whereas the causality shown to the right leads to the equation:

$$u_R = R \cdot i_R$$

Capacitances and inductances store the energy that flows into them. Resistors dissipate the energy. Since power cannot be dissipated, also the $R$ element is basically a non-physical element. It denotes another interface to the universe. The dissipated power is converted into heat. Since our simple model reflects only the electrical properties of reality, the generated heat is part of the universe, and not part of the system.

Since we add up the flows in a 0-junction, we can generate only one equation for the flows into a 0-junction. Consequently, only one of the bonds attached to a 0-junction can have its stroke at the side of the junction. Similarly, all bonds but one attached to a 1-junction must have their strokes at the side of the 1-junction.

These causality rules suffice to determine a unique causality for our simple RLC-circuit:

$$\overrightarrow{u_C} \overleftarrow{i_C} \overrightarrow{u_I} \overleftarrow{i_I}$$

In this example, we were able to satisfy all causality constraints in a unique manner. This is the preferred situation. If not all mandated causality constraints can be satisfied, we are confronted with a non-causal system. This case occurs for example if we try to
parallel connect two voltage sources with different voltage values. If we cannot satisfy all recommended causality constraints, i.e., if we run into wrong causality at either $C$ or $I$ elements, we are confronted with a degenerate system, i.e., the true system order is lower than the number of energy storages makes us believe. If we have a choice in assigning causality without offending any of the causality constraints, we have a system with one or several algebraic loops.

Mechanical systems can be modeled in a similar manner. Let us look at a simple translational system:

In a translational system, the forces have been made the effort variables, and the velocities have been made the flow variables. The mechanical power of translation can be expressed as the product of a force $f_i$ and a velocity $v_i$:

$$ P_{trans} = f_i \cdot v_i $$

The rules for constructing the bond graph are as simple as in the electrical case. We start by identifying all free moving bodies. We place 1-junctions for each of their velocities. Where ever two bodies interact with each other, we connect their junctions with branches consisting of two bonds and one 0-junction inbetween, and attach all interacting elements to that 0-junction. Newton's law (or rather the d'Alembert principle) is formulated at the 1-junctions themselves. The bond graph for the above translational system is shown below:

Rotational systems can be modeled in exactly the same way. Here, the effort variables are the torques $r_i$, and the flow variables are the rotational velocities $\omega_i$. The mechanical power of a rotation can be expressed as:

$$ P_{rot} = r_i \cdot \omega_i $$

Beside from the two basic quantities effort $e$ and flow $f$, we often make use of two additional derived quantities, namely the generalized momentum:

$$ p = \int_0^t e \, dt $$

and the generalized displacement:

$$ q = \int_0^t f \, dt $$

In electrical systems, the generalized momentum is the flux through a coil, and the generalized displacement is the charge in a capacitor. In translational mechanical systems, these are the momentum and the displacement (bond graphs were invented by mechanical engineers), and in rotational mechanical systems, they are the angular momentum and the angular position.

All these quantities are common to a large variety of other physical systems as well, as are the two Kirchhoff laws. Hydraulic, pneumatic, and acoustic systems operate similarly to the electrical and mechanical ones. In all these systems, the pressure is defined as the effort variable, while the volume flow rate is defined as the flow variable. The derived quantities are the pressure momentum and the volume.

The element laws, however, may look different for different types of systems. In particular, it may be noted that the equivalent to Ohm's law for these types of systems is often non-linear. E.g., the relation between effort $p$ and flow $q$ in a pipe or valve is quadratic:

$$ \Delta p \propto q^2 $$

Table 1 presents a summary of the four generic variables for the most commonly used types of physical systems.

<table>
<thead>
<tr>
<th>Effort</th>
<th>Flow</th>
<th>Generalised Momentum</th>
<th>Generalised Displacement</th>
</tr>
</thead>
<tbody>
<tr>
<td>$e$</td>
<td>$f$</td>
<td>$p$</td>
<td>$q$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Electrical</th>
<th>voltage</th>
<th>current</th>
<th>flux</th>
<th>charge</th>
</tr>
</thead>
<tbody>
<tr>
<td>$w , [V]$</td>
<td>$i , [A]$</td>
<td>$\Phi , [V \cdot s]$</td>
<td>$q , [A \cdot s]$</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Translational</th>
<th>force</th>
<th>velocity</th>
<th>momentum</th>
<th>displacement</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F , [N]$</td>
<td>$v , [m/s]$</td>
<td>$J , [N \cdot m]$</td>
<td>$x , [m]$</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Rotational</th>
<th>torque</th>
<th>angular velocity</th>
<th>twist</th>
<th>angle</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T , [N \cdot m]$</td>
<td>$\omega , [rad \cdot s^{-1}]$</td>
<td>$T , [N \cdot m \cdot s]$</td>
<td>$\phi , [rad]$</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Hydraulic</th>
<th>pressure</th>
<th>volume flow</th>
<th>pressure momentum</th>
<th>volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>$p , [N \cdot m^{-2}]$</td>
<td>$q , [m^3 \cdot s^{-1}]$</td>
<td>$\Gamma , [N \cdot m^{-2} \cdot s]$</td>
<td>$V , [m^3]$</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chemical</th>
<th>chemical potential</th>
<th>molar flow</th>
<th>number of moles</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mu , [J \cdot mol^{-1}]$</td>
<td>$\nu , [mol \cdot s^{-1}]$</td>
<td>$n , [mol]$</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Thermo-dynamical</th>
<th>flow</th>
<th>temperature</th>
<th>entropy</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T , [^\circ K]$</td>
<td>$\frac{dW}{dt} , [W \cdot ^\circ K^{-1}]$</td>
<td>$S , [J \cdot ^\circ K^{-1}]$</td>
<td></td>
</tr>
</tbody>
</table>
at one end must come out in one or more different form(s) at the other. A "lossless" energy transducer may e.g. transform electrical energy into mechanical energy. In reality, every energy transducer "loses" some energy, but the energy does not really disappear—it is simply transformed into heat.

There are exactly two ways how the above energy conservation law can be satisfied in an ideal energy transducer (i.e., an energy transducer in which only two forms of energy are involved). One is the ideal transformer. It is governed by the following set of relationships:

\[ e_1 = m \cdot e_2 \]
\[ f_2 = m \cdot f_1 \]

The ideal transformer is placed between two junctions. There are two types of causalities possible as shown below:

\[ \frac{e_1}{f_1} \rightarrow TF \rightarrow \frac{e_2}{f_2} \]

\[ e_2 = \left( \frac{1}{m} \right) e_1 \]
\[ f_2 = \left( \frac{1}{m} \right) f_1 \]

Examples of transformers are the electrical transformer, the mechanical gear, and the mecano-hydraulic pump:

The other type of energy transducer is the ideal gyrator. Its behavior is governed by the equations:

\[ e_1 = r \cdot f_2 \]
\[ e_2 = r \cdot f_1 \]

Also the ideal gyrator exhibits two forms of causalities as shown below:

\[ \frac{e_1}{f_1} \rightarrow GY \rightarrow \frac{e_2}{f_2} \]

\[ e_2 = r \cdot f_1 \]
\[ e_1 = r \cdot f_2 \]

Examples of gyators are most electro-mechanical converters, e.g., the DC-motor:

Notice that there is no real difference between the two transducer types. If the effort and flow variables in the mechanical system were toggled, the DC-motor would in fact become a transformer.

Let us go through an example. We want to model an armature controlled DC-motor with constant field which drives a translational load connected to the motor through a slip clutch and a gear. The resulting bond graph is shown below:

The electrical power generated by the effort source is partly stored in the armature inductance \( L_a \), and is partly dissipated in the armature resistance \( R_a \). The remaining power is available for conversion into mechanical power. On the rotational side, the power is partly stored in the internal motor inertia \( J_i \), and partly dissipated through friction \( B_1 \). The slip clutch reduces the angular velocity from \( \omega_1 \) to \( \omega_2 \). It contains the friction \( B_1 \). The remaining energy is partly stored in the external motor inertia \( J_e \) which contains the primary cog of the gear. There is also a spring \( k_1 \) which prevents the axle from rotating too far. The gear itself is represented by a transformer. In this example, it converts the remaining rotational energy into translational energy. The load consists of a mass \( m \), another spring \( k_2 \), a friction \( B_2 \), and the gravity force \( m \cdot g \) (another source).

The bond graph shows clearly that the power available for conversion from the electrical side to the rotational mechanical side is:

\[ P_{el} = u_i \cdot i_a = P_{rot} = r \cdot \omega_1 \]

The electrical power available for conversion is the product of the induced voltage \( u_i \) and the armature current \( i_a \). This power is equal to the generated rotational power which can be described as the product of the motor torque \( r \) and the angular velocity of the motor \( \omega_1 \). This fact would not have been so easy to read out of a block diagram or a signal flow graph.

Notice the problem with the causality assignment. After satisfying all mandated causalities at the sources and junctions, we are confronted with a conflict. The translational inerance (the mass) \( m \) has the wrong causality. Thus, while this system exhibits six different energy storages, it is in fact a fifth order system.

**BOND GRAPH MODELING IN DYMOLA**

The first bond graph simulation language developed in the early seventies was ENPORT-4 (Rosenberg 1974). This software used an approach similar to SPICE, i.e., it did not request causalities to be specified, and it transformed the topological input description into a branch admittance matrix which could then be solved employing similar techniques to those used in SPICE. Consequently, ENPORT-4 was able to handle structurally singular problems. Unfortunately, ENPORT's input language was not very user-friendly, and the code did not operate too well when we used it about ten years ago. To our knowledge, ENPORT-4 is no longer in use.

In the late seventies, another bond graph simulation language was developed at Twente University in the Netherlands, called THTSIM in Europe, and TUTSIM in the United States (van Dijkhoorn 1982). TUTSIM translated bond graphs into a state-space representation. The user is requested to specify the causalities, and structurally singular systems cannot be handled. TUTSIM's simulation engine is somewhat poor in comparison with other state-
space solvers such as ACSL. Its input language is better than that of ENPORT-4. TUTSIM is currently still in use.

The third product on the market is CAMP (Granda 1982, 1985), a preprocessor to ACSL (Mitchell and Gauthier 1985), which translates bond graphs into ACSL programs. CAMP has the same limitations as TUTSIM, i.e., it does not handle algebraic loops or structural singularities, but it has the better simulation engine (ACSL). The input format is topological (as for the two other products), and therefore, it is not truly flexible with respect to handling non-standard network elements. None of the three products is able to handle hierarchically structured systems, or arbitrary non-linearities.

For these reasons, we decided to go a different route. DYMOLA (Emlqvist 1978) is a general purpose hierarchical modular modeling software for continuous-time systems. As we noticed, elements — such as resistors — which allow different causalities to be applied lead to different equations depending on their causality assignment. Thus, it is not sufficient that a modeling software can sort equations (as most CSSL-type languages will do), but in addition, it is necessary that the software can solve equations for any variable. E.g., the software must be able to turn an equation of the type:

\[ u = R \cdot i \]

into:

\[ i = \frac{1}{R} \cdot u \]

when needed. DYMOLA provides for this capability.

DYMOLA also supports the concept of across and through variables. The DYMOLA statement:

```plaintext
cut A(v/i)
```

defines an electrical wire with the potential \( v \) and the current \( i \). Cuts are hierarchical data structures (similar to PASCAL records) that enable the user to group individual wires into buses or cables, and cables into trunks. A cut is like a plug or a socket. It defines an interface to the outside world. The DYMOLA statement:

```plaintext
connect z : A at y : B
```

plugs the cut \( A \) of model \( z \) into the socket \( B \) of model \( y \). Thereby, all the across variables (to the left of the slash separator) are set equal, and all the through variables (to the right of the slash operator) are summed up to zero. The DYMOLA preprocessor automatically generates the necessary coupling equations.

Nodes are a convenient means to organize connections. They act like your power distributor. You can plug several appliances into one such distributor. The above statement could also have been coded as:

```plaintext
node n
cut A(v/i)
```

DYMOLA’s nodes can be used as 0-junctions in a bond graph model. There is no DYMOLA equivalent for 1-junctions, but, as we explained before, 1-junctions are the same as 0-junctions with the effort and flow variables interchanged. Therefore, we created a model type “bond” which simply exchanges the effort and flow variables:

```plaintext
model type bond
cut A(y/x) B(x/y)
main cut C[A B]
main path P < A - B >
```

The bond acts just like a null-modem for a computer. Since neighboring junctions are always of opposite sex, they can both be described by regular DYMOLA “nodes” if they are connected with a “bond”.

Notice that my “bond” model type is actually a gyration with \( r = 1.0 \). This special gyration has sometimes been called symplectic gyration in the bond graph literature (Breedveld 1982, 1984).

Since we don’t want to maintain different types of \( R, C, L, TF, \) and \( GY \) elements, we add one additional rule: in DYMOLA, all elements (except for the bonds) can be attached to 0-junctions only. If they need to be attached to a 1-junction, we simply must place a bond inbetween.

The following DYMOLA model types suffice to describe simple bond graphs.

```plaintext
model type SE
cut A(e/f)
terminal E0
E0 = e
end
```

```plaintext
model type SF
cut A(f/j - f)
terminal F0
F0 = f
end
```

```plaintext
model type R
cut A(e/f)
parameter R = 1.0
R * f = e
end
```

```plaintext
model type C
cut A(e/f)
parameter C = 1.0
C * C = f
end
```

```plaintext
model type I
cut A(e/f)
parameter I = 1.0
I * C = f
end
```

```plaintext
model type TF
cut A(e/f) B(e2 - f2)
main cut C[A B]
main path P < A - B >
parameter m = 1.0
e1 = m * e2
f2 = m * f1
end
```

```plaintext
model type GY
cut A(e/f) B(e2 - f2)
main cut C[A B]
main path P < A - B >
parameter r = 1.0
e1 = r * e2
e2 = r * f1
end
```

With these modeling elements, we can e.g. formulate a bond graph description of our simple RLC network. Remember to expand the bond graph in such a way that all elements are attached to 0-junctions only. This is shown below.
The enter model statement reads in the model and immediately expands the set of equations by the coupling equations. The outfile statement specifies the name of the output file, and the output equations statement writes the generated equations to the output file. The generated equations are shown in the next code segment.

```
R : R1 = 100Ω
I : L1 = 1.5 mH

\[ U_0 \rightarrow U_R, R \rightarrow U_R \]
\[ \dot{Q} = U \]
\[ U_{R1} \rightarrow R \]
\[ L \rightarrow U \]
\[ R \rightarrow U_R \]
\[ U_R \rightarrow R \]
\[ C \rightarrow U \]
\[ C_{12} \rightarrow U \]

C : C = 0.1μF
```

I did not mark down the causalities here since DYMOLA is perfectly able to handle the causality assignment by itself (although no structural singularities yet).

The expanded bond graph can be immediately coded in DYMOLA as shown below:

```
@bond.bnd
@se.bnd
@r.bnd
@c.bnd
@i.bnd

model RLC

submodel (SE) U0
submodel (R) R1(R = 100.0), R2(R = 20.0)
submodel (L) L1(L = 1.5S - 3)
submodel (C) C1(C = 0.1E - 6)
submodel (bond) B1, B2, B3
node u1, i1, v1, i2, v2
input u
output y1, y2

connect U0 at u1
connect L1 at i1
connect R1 at v1
connect R2 at v2
connect C1 at i2
connect B1 from u1 to i1
connect B2 from i1 to v1
connect B3 from i1 to v1

U0.E0 = u
y1 = C.e
y2 = R2.f
```

The first six equations were extracted from the models. The remaining equations are automatically generated coupling equations.

We can now execute the algorithm which assigns the causalities, i.e., which determines what variable to compute from each of the equations. In DYMOLA, this is achieved with the following set of instructions:

```
> partition
> outfile rlc.sor
> output sorted equations
```

which results in the following answer:

```
RLC [U0.E0 = u]
U0 E0 = [v]
RLC U0.e = [L1.e]
L1.e = [B1.x]
C1.e = [B2.y]
B3.y + B2.y = B1.x
RLC R.f = [v]
RLC R1.f = [B3.x]
[ B2.x] = B5.x
B1.y = B2.x
[R2.e] = C.e
RLC C.e = [v]
L1 L = [v]
RLC C = [v]
RLC y1 = C.e
y2 = R2.f
```

This time, we decided to print out the sorted equations. The variables enclosed in "[ ]" are the variables for which each equation must be solved. This set of equations contains many trivial equations of the type \( \alpha = \beta \). DYMOLA is capable of throwing those out. This is accomplished through the following set of instructions:

```
> partition eliminate
> outfile rlc.rs2
> output sorted equations
```

which results in the following answer:

```
```

Let us see how the DYMOLA compiler preprocesses this code. DYMOLA is actually a program generator. At the current time, DYMOLA can generate either DESIRE (Korn 1989) or SIMNON (Elmqvist 1975) simulation programs. We enter DYMOLA, and specify the model to be compiled as follows:

```
$ dymola
  > enter model
  - @rlc.dym
  > outfile rlc.eq
  > output equations
```

The interpretation of this code is straightforward. DYMOLA's @ operator corresponds to the include statement of most programming languages. It includes the element definitions which were stored on separate files.
which is a much reduced set of equivalent equations. The next step will be to actually perform the symbolic manipulation on the equations. In DYMOLA, this is done in the following way:

```
> out file rlc.sov
> output solved equations
```

which results in the following answer:

```
R2   y2 = y1/R
R1   B3.y = u - y1
R2   B3.x = B3.y/R
RLC  C1.f = B3.x - y2
C1   dere = f/C
L1   der f = u/L
```

We are now ready to add the experiment description to the model. We can for instance use the following experiment:

```
emdef
simtime 2E - 5
step 2E - 7
commpoints 101
input 1, u,pendepend, 10.0)
```

```
cutblock
scale = 1
XCCC = 1
label TRY
Dr u |if XCCC < 0 then XCCC = -XCCC
scale = 2 * scale
go to TRY
else proceed
```

```
cend
```

```
outblock
OUT
disp y1, y2
```

```
end
```

This portion of code is specific for each of the target languages. The here shown version is the one required for DESIRE (Korn 1989). The cutBlock set of statements instructs DESIRE to automatically scale the run-time display. XCCC is a DESIRE variable which is set to -1 whenever the DESIRE program is interrupted with an “overflow”. This happens when one of the displayed variables hits either the top or the bottom of the displayed window. At this time, the plot is simply rescaled, and rerun with a new Dr u statement. Since DESIRE is so fast, it is not worth the effort to store the results of the previous attempt, instead, we simply rerun the entire simulation.

The set of DYMOLA instructions:

```
> enter experiment
    > file rlc.des
    > output desire program
```

```
which tells DYMOLA to generate the following DESIRE program:
```

--- CONTINUOUS SYSTEM RLC
--- STATE y1 L1$f
--- DER dC1$e d$L1$f
--- PARAMETERS and CONSTANTS:
R1$E = 100.0
C = 1.5E - 6
L = 1.5E - 3
R2$E = 20.0
--- INITIAL VALUES OF STATES:
y1 = 0
L1$f = 0
u = 10.0
---
TMAX = 2E - 5 | DT = 2E - 7 | N = 101
scale = 1
XCCC = 1
label TRY
Dr u |if XCCC < 0 then XCCC = -XCCC
scale = 2 * scale
go to TRY
else proceed
```

DYNAMIC
---
--- Submodel : R2
y2 = y1/R2$E
--- Submodel : RLC
B3$y = u - y1
--- Submodel : R1
B3$x = B3$y/R1$E
--- Submodel : RLC
C1$f = B3$x - y2
--- Submodel : C1
d/dt y1 = C1$f/C
--- Submodel : L1
d/dt L1$f = u/L
```

```
OUT
disp y1, y2
```

```
/---
/PIC ’rlc_PRC’
/---
```

which can be executed at once using the following instructions:

```
> stop
$ desire
> load ’rlc.des’
> run
```

which will immediately (within less than a second) produce the desired output variables u0, and iR2 on the screen. Both DYMOLA (Elmqist 1978) and DESIRE (Korn 1989) are currently running alternatively on VAX/VMS or PC/MS-DOS. UNIX versions are in the planning for both of these languages. It is also planned to add an ACSL interface to DYMOLA.

THERMODYNAMIC BOND GRAPHS

Thermal conductive or convective flow of heat can be described by the heat equation:

\[
\frac{\partial T}{\partial t} = \sigma \cdot \nabla^2 T
\]

Heat flow in one space direction can be modeled by a simple RC chain as shown below:
Using the bond graph methodology, the $RC$ chain can be represented as follows:

\[
\begin{align*}
\dot{T}_1 &= R \Delta T_1 \\
\Delta T_2 &= R \Delta T_2 \\
\Delta T_3 &= R \Delta T_3 \\
\cdots \\
\Delta T_{n-1} &= R \Delta T_{n-1} \\
\dot{T}_n &= C \Delta T_n \\
\end{align*}
\]

This bond graph shows that the approach does not work. When modeling electrical or mechanical phenomena, we could afford to model the dissipation of power through resistive elements. However, we have seen already that this is basically a non-physical concept. Power cannot be dissipated, only transported and converted. When modeling thermodynamic systems, the idea becomes absurd. What does it mean when we say that thermal power is being dissipated? This is obviously meaningless. Heat gets "absorbed" by the resistor, but it is immediately "generated" again by the same resistor.

To overcome this difficulty, Thoma introduced a new bond graph element called a resistive source (Thoma 1975). The heat which is absorbed by the resistive element is routed through the resistive source, and is immediately re-introduced at the next junction. The modified bond graph is shown below:

\[
\begin{align*}
\dot{T}_1 &= R \Delta T_1 \\
\Delta T_2 &= R \Delta T_2 \\
\Delta T_3 &= R \Delta T_3 \\
\cdots \\
\Delta T_{n-1} &= R \Delta T_{n-1} \\
\dot{T}_n &= C \Delta T_n \\
\end{align*}
\]

The enhanced bond graph still represents the heat equation, but while the previously suggested bond graph models the temperature flow only, the enhanced bond graph models also the power flow correctly which can be written as:

\[
P_{\text{therm}} = T \cdot S
\]

Thermal power is the product of the temperature $T$ and the entropy flow $S$.

In DYMOLO, the $RS$ element can be represented as follows:

\[
\begin{align*}
\text{model type rs} \\
\text{cut } A(e1/f1) B(e2/-f2) \\
\text{main cut } C[A B] \\
\text{main path } P < A - B > \\
\text{parameter } R = 1.0 \\
R \cdot f1 = e1 \\
e1 \cdot f1 = e2 \cdot f2 \\
\end{align*}
\]

There is however still one problem to be considered. Contrary to the electrical and mechanical systems, the $R$ and $C$ element of thermodynamics are not constant. The $R$ element is proportional to the temperature:

\[
R = \theta \cdot T
\]

whereas the $C$ element is inverse proportional to the temperature:

\[
C = \frac{1}{T}
\]

I.e., thermodynamic $R$ and $C$ elements are non-linear. To overcome this problem, earlier bond graph software systems introduced the heat flow as the flow variable instead of the entropy flow. This makes the $R$ and $C$ elements linear, but, at the same time, it makes it difficult to interface the so-called thermodynamic bond graphs with other bond graphs in a mixed energy system, since the product of temperature and heat flow does not represent power.

In DYMOLO, we don't have this problem, since we can model non-linearities easily. We simply introduce two new bond graph elements, the modulated resistive source: $mRS$, and the modulated capacitance: $mC$.

\[
\begin{align*}
\text{model type mRS} \\
\text{cut } A(e1/f1) B(e2/-f2) \\
\text{main cut } C[A B] \\
\text{main path } P < A - B > \\
\text{parameter } \theta = 1.0 \\
R = \theta \cdot e2 \\
R \cdot f1 = e1 \\
e1 \cdot f1 = e2 \cdot f2 \\
\end{align*}
\]

Notice that, in the $mRS$ model, $\theta$ must be multiplied by $e_3$ and not by $e_1$ since $e_2$ denotes an absolute temperature whereas $e_1$ denotes a temperature difference.

Let me now explain how the bond graph modeling concept can be used in a hierarchical fashion. For this purpose, we shall study a considerably more complex example: a solar heated house. The overall configuration is shown below:

One or several collectors act as black bodies which absorb incoming solar radiation. Consequently, the temperature inside the collectors raises. The collectors can be filled with any material with a large heat capacity. Usually, it is simply air. Inside the collectors,
there is a water pipe which meanders back and forth between the two ends of the collector to maximize the exposed pipe surface. We shall call this a “water spiral”. A (mostly conductive) heat exchange takes place between the collector chamber and the water pipe, thereby heating the water in the pipe. A pump circulates the water from the collectors to the storage tank, thereby transporting the heat convectively from the collectors to the tank. We call this the “collector water loop”. The water spirals in the various collectors can be either series connected, or they can be connected in parallel. The pump is usually driven by a solar panel. In the panel, the solar light is converted to electricity which drives the pump. Thereby, the pump circulates the water only while the sun is shining which is exactly what we want. In addition, a freeze protection device is often installed which also switches the pump on whenever the outside temperature falls below 5°C.

The storage tank is often realized simply as a large and well insulated water container (a water heater). However, we shall assume that a solid body storage tank is used together with another water spiral which deposits the heat in the storage tank just the same way as it was picked up in the collectors. Consequently, the water from the collector loop and from the heater loop never mix.

Inside the storage tank, there is a second water spiral which belongs to the “heater water loop”, and which can pick up the heat from the storage tank. There is also an additional electrical heater installed which heats the storage tank electrically whenever the storage tank temperature falls below a critical value.

The heater water loop is driven by another pump which is switched on whenever the room temperature falls below 20°C during the day or 18°C during the night, and which is switched off whenever the room temperature raises beyond 22°C during the day or 20°C during the night.

In the house, we use one or several “radiators” (more water spirals) which, contrary to what their name suggests, exchange heat with the room in a partly conductive and partly convective manner.

Below, the collector is shown in more detail.

Each cell is described by a DYMOLA model type called cfd.dym which, from now on, can be used as an additional bond graph element. The correct causalities have been marked on the graph. The mGS element is a “modulated conductive source”. It is modulated with temperature (as always in thermal systems), but, in addition, it is also modulated with the water velocity in the pipe as shown below:

Since the conductance changes linearly with the water velocity \( v_w \), I preferred to model this element through its conductance rather than through its resistance. The cfd model references three submodels: a temperature modulated capacitance \( mc \), a temperature and water velocity modulated conductive source \( mGS \), and finally the regular bond submodel.

The exchange of heat across the border of two media is modeled by a heat exchanger as shown below:

The heat exchanger is used here to model the transfer of heat from the collector chamber to the water spiral.

The water spiral is modeled through a series connection of several cfd elements with heat exchangers attached in between. Below, the water spiral is shown:

We decided to cut the spiral into three discrete links. Obviously, this is an approximation of a process with distributed parameters.

Notice that the newly introduced bond graph symbol representing the water spiral is a 3-port element.
We need to model also the loss from the collector chamber to the environment. This loss is partly conductive and partly convective. Below, the loss element (a 1-port element) is shown:

\[
\begin{array}{c}
mG \\
\text{0} \rightarrow 1 \\ \
\text{SE}
\end{array} \quad \Rightarrow \quad \text{0} \rightarrow \text{Loss}
\]

The effort source denotes the outside temperature. The \( mG \) element denotes the heat dissipation to the environment. The dissipated heat is proportional to the difference in temperatures between the inside and the outside. \( mG \) is a modulated conductance similar to the \( mGS \) element found earlier, but this time, the secondary port (the environment) is not modeled, and the modulation is now with respect to the wind velocity \( v_{\text{wind}} \) rather than with respect to the water velocity \( v_{\text{w}} \).

We are now ready to model the overall collector. It is shown below:

\[
\begin{array}{c}
mC \\
\text{SF} \rightarrow 0 \\ 
\text{Spi} \rightarrow 0
\end{array} \quad \Rightarrow \quad \text{0} \rightarrow \text{Coll} \rightarrow 0
\]

The \( mC \) element is the (temperature modulated) heat capacity of the collector chamber. The \( SF \) element denotes the heat input from solar radiation.

We used the hierarchical cut concept of Dymola to combine the two cuts (i.e., bonds), \( \text{inwater} \) and \( \text{outwater} \), into one hierarchical cut, \( \text{water} \). This can be pictorially represented by a double bond. This aggregated bond graph representation has, of course, the disadvantage that causalities can no longer be depicted.

We shall now model the transport of heat from the collector to the storage tank, i.e., the collector water loop. We model each of the pipes by a series of one-dimensional cells, and we shall assume that the pipes are thermally well insulated, i.e., that no heat is lost to the environment on the way. The water loop is depicted below:

\[
\begin{array}{c}
\text{0} \rightarrow \text{1D} \rightarrow \text{0} \rightarrow \text{1D} \\ 0 \rightarrow \text{1D} \rightarrow \text{0} \rightarrow \text{1D}
\end{array} \quad \Rightarrow \quad \text{0} \rightarrow \text{Coll} \rightarrow 0
\]

The \( mC \) element denotes the heat capacity of the storage tank. The flow source together with the \( mRS \) element denote the electrical backup heater. The primary side of the resistive source is electrical while the secondary side is thermic.

This is another 4-port. This time, we shall combine the cut \( \text{inwater1} \) with the cut \( \text{outwater1} \) to the hierarchical cut \( \text{inwater} \), and the cut \( \text{outwater2} \) with the cut \( \text{inwater2} \) to the hierarchical cut \( \text{outwater} \). We shall again declare a main path water which creates a logical bridge from the hierarchical cut \( \text{inwater} \) to the hierarchical cut \( \text{outwater} \).

The storage tank contains two water spirals, one which belongs to the collector water loop, and one which belongs to the heater water loop. In addition, there has been installed an electrical resistance heater as a backup device. Below, the storage tank is shown:

\[
\begin{array}{c}
mC \\
\text{SF} \rightarrow mRS \rightarrow 0 \\
0 \\
\text{ST}
\end{array} \quad \Rightarrow \quad \text{0} \rightarrow \text{ST} \rightarrow 0
\]

The \( mC \) element denotes the heat capacity of the storage tank. The flow source together with the \( mRS \) element denote the electrical backup heater. The primary side of the resistive source is electrical while the secondary side is thermic.

The heater water loop is modeled in exactly the same manner as the collector water loop.

The house itself is a little more tricky. We can model the house through a number of three-dimensional cells with loss elements attached to the walls, and heat exchangers denoting the heat input through the radiators. However, space limitations will prevent me from carrying this example all the way through to the end. More details can be found in a forthcoming book of mine (Cellier 1990).

Finally, we can connect all the pieces together and create a model for the overall house:

\[
\text{Coll} \rightarrow \text{WL} \rightarrow \text{ST} \rightarrow \text{WL} \rightarrow \text{HE} \rightarrow \text{House}
\]

This concludes the description of DYMOLA's modular hierarchical non-linear bond graph modeling capability.

**CHEMICAL REACTION KINETICS AND CHEMICAL THERMODYNAMICS**

Chemical power can be expressed as the product of the chemical potential \( \mu_i \) of a chemical species \( i \) multiplied by its molar flow rate \( \nu_i \):

\[
P_{\text{chem}} = \mu_i \cdot \nu_i
\]

Here, \( \mu_i \) is the effort variables, while \( \nu_i \) is the flow variable.

Traditionally, the reaction kinetics equations are expressed in terms of the involved number of moles and the molar flow rates only. The chemical potential does not appear in these (highly non-linear) differential equations. E.g. the reaction system:
\[ BR_2 \xrightarrow{h_1} 2Br^* \]
\[ 2Br^* \xrightarrow{h_2} Br_2 \]
\[ Br^* + H_2 \xrightarrow{h_3} HBr + H^* \]
\[ HBr + H^* \xrightarrow{h_4} Br^* + H_2 \]
\[ Br_2 + H^* \xrightarrow{h_5} HBr + Br^* \]

can be described by the set of differential equations:

\[ \frac{d}{dt} n_{Br_2} = -\nu_{k1} + \nu_{k2} - \nu_{k5} \]
\[ \frac{d}{dt} n_{Br^*} = 2\nu_{k1} - 2\nu_{k2} - \nu_{k3} + \nu_{k4} + \nu_{k5} \]
\[ \frac{d}{dt} n_{Br^*} = -\nu_{k3} + \nu_{k4} \]
\[ \frac{d}{dt} n_{H^*} = \nu_{k3} - \nu_{k4} + \nu_{k5} \]
\[ \frac{d}{dt} n_{HBr} = \nu_{k4} - \nu_{k5} \]

where \( n_i \) denotes the number of moles of the component gas \( i \), and \( \nu_{ki} \) is the reaction flow rate of the reaction \( k_i \):

\[ \nu_{k1} = k_1 \cdot n_{Br_2} \]
\[ \nu_{k2} = k_2 \cdot \left( \frac{Br^*}{V} \right) \]
\[ \nu_{k3} = k_3 \cdot \left( \frac{n_{Br^*} \cdot n_{Br^*}}{V} \right) \]
\[ \nu_{k4} = k_4 \cdot \left( \frac{n_{H^*} \cdot n_{Br^*}}{V} \right) \]
\[ \nu_{k5} = k_5 \cdot \left( \frac{n_{H^*} \cdot n_{Br^*}}{V} \right) \]

The chemical thermodynamics equations, on the other hand, i.e., the equations which can be used to determine the chemical potential, aren't commonly handled by means of differential equations at all. Chemical thermodynamics in fact are a misnomer. Chemical thermostatics would be a more appropriate term. Thermodynamics, if at all, are only treated through small signal aberrations from the steady-state. This is because the true thermodynamics equations are still poorly understood.

Thus, the "kineticists" deal with mass flow only. They ignore the power flow altogether. The "thermodynamicists" deal with the energy balance under equilibrium conditions, and they actually ignore both the mass flow and the power flow.

How come that this decomposition is possible? E.g., in electrical systems, we weren't able to decompose the dynamics into one set of equations involving currents only, and another set involving potentials exclusively. Why is it that both thermal and chemical systems exhibit this decomposition property? Remember that also the heat equation can be formulated either in terms of temperature or in terms of entropy alone. I discovered that this astounding property can be derived from the fact that neither thermal nor chemical systems have an inertia element. Due to the fact that there exists only one type of energy storage element, the equations become decoupled. However, we now have to deal with two separate forms for power flow through the system. In a chemical system, the two forms are:

\[ P_{therm} = \mu \cdot n \]
\[ P_{chem} = \dot{\mu} \cdot n \]

and in a thermal system, they are:

\[ P_{therm} = T \cdot \dot{S} \]
\[ P_{therm} = T \cdot S \]

As in the case of the thermal systems, this seemingly convenient decomposition property has a considerable disadvantage. Such models cannot easily be connected to other subsystems of different energy type. Most chemical reactions occur either exothermically or endothermically, and often, either the pressure or the volume changes as well. Thus, we should study chemical reactions, their thermodynamics, and their hydraulic/pneumatic flows in unison.

I wish to discuss how the concepts of bond graph modeling can help us to gain an improved understanding of what is happening in these highly complicated systems. We start by noticing that our combined dynamics will contain six types of variables: (i) the temperature \( T \), (ii) the entropy \( S \), (iii) the hydraulic/pneumatic pressures \( p \), (iv) the volume \( V \), (v) the chemical potential \( \mu \), and (vi) the number of moles \( n \). Consequently, we need six types of equations to describe the dynamics of this system in their entirety.

We have met one type of equation so far: the mass flow equations (reaction kinetics equations). It is also well known that there exists a power balance equation which can e.g. be written as:

\[ p \cdot \dot{V} = T \cdot \dot{S} + \sum_{i} \mu_i \cdot n_i \]

This is one way to express the combined effects of the first two laws of thermodynamics. However, I have discovered that chemical systems obey a second type of power balance equation as well:

\[ \dot{p} \cdot V = \dot{T} \cdot S + \sum_{i} \dot{\mu}_i \cdot n_i \]

The detailed derivation of this formulae is in my forthcoming book (Cellier 1990).

Furthermore, we realize that energy is traded between the chemical system and its thermal and hydraulic/pneumatic environments. Thus, there must exist some sort of transformer with bonds into the thermal and hydraulic/pneumatic "worlds". Since we know that, for each bond, there is a variable computed at each of the two ends of the bond, we realize that among the variables \( T \) and \( S \), one must be computed on the chemical side, while the other is computed on the thermal side of the bond. Similarly for \( p \) and \( V \). Thus, within the chemical system, we actually need only four different types of equations rather than six as previously assumed. The other two are dictated from the outside.

This fact is commonly reflected in chemical engineering by an assumption of e.g. isothermic and isobaric or isothermal and isochoric conditions. I.e., the chemical engineer makes his life easier by holding one of the thermal variables and any one of the hydraulic/pneumatic variables at a constant value. However, these are just special cases of a more general truth.

Finally, we need one more equation. For this purpose, we shall use the so-called equation of state. Every system has one such equation. E.g. an ideal gas reaction exhibits the following property:

\[ p \cdot V = n \cdot R \cdot T \]

This gives us all equations needed to describe the dynamics of this system. Unfortunately, there is a problem with the last of these equations. The equation of state is actually a steady-state equation, i.e., it is only valid in a strict sense under equilibrium conditions. It should be replaced by a more general equation which is true also far from equilibrium. Unfortunately, I haven't been able to find such an equation yet.

The current literature on chemical thermodynamics confounds wildly equations which describe structural (i.e., physical) properties of chemical reaction systems with other equations which are balance equations that are valid only under equilibrium conditions. The assumptions made are hardly ever explicitly stated.
Even worse, the chemical thermodynamics literature is full of myths. Examples of such myths are given in my forthcoming book (Cellier 1990) in which I tried carefully to make all assumptions explicitly known.

We are now ready to set up a chemical bond graph. In this paper, I shall deal with the simplest of all cases only: the isothermic and isobaric reaction. In this case, the second power balance equations degenerates, and the chemical potentials become constants. Below, the isothermic and isobaric bond graph of the hydrogen/bromine reaction system is shown. The fourth (and least important) reaction $k_4$ has been eliminated to keep the bond graph planar.

The $CS$ element is a newly introduced capacitive source. It models the mass storage. One $CS$ element is used for each species. It computes the chemical potential of the component species, i.e., it is a source of chemical potential. It also stores the molar mass. In our simple situation, it can be modeled as follows:

```plaintext
model type CS
  cut chem(mu/nu)
  terminal n, mu0
  nu = mu0
  den(n) = nu
end
```

The capacitive source is attached to a 0-junction which models the chemical mass balance, i.e., it converts the reaction flow rates into component flow rates.

Each reaction is represented by a new bond graph element of type chemical reactor. A chemical reactor is a three-legged transformer. It contains the first power balance equation, the equations for the reaction flow rates, and the equation of state expressed in terms of partial volume flows, not decomposed into an individual component gas, but decomposed into an individual reaction. My book (Cellier 1990) provides the details of the calculations. The DYMOOLA version of e.g. the third chemical reactor is as follows:

```plaintext
model type Ch3.k3
  cut chem(nu.k3/ - nu.k3)
  cut therm(T / - S.dot.k3), pneum(p/q.k3)
  terminal n.H2, n.Br, V
  parameter R = 8.314
  local k3, nue.k3
  k3 = (10 * 11.43) * exp(-82400/(R + T))
  nue.k3 = 0.0
  p * q.k3 = T * S.dot.k3 + nu.k3 * mu.k3
  p * q.k3 = nue.k3 * R + T
  nu.k3 = k3 * n.H2 * n.Br/V
end
```

Due to the high degree of non-linearity and modulation, it is necessary to define a model type for each of the reactors. Each of the reactors is attached to a 1-junction which models the chemical energy balance, i.e., it converts the chemical potentials of the component species into "reaction potentials" which are expressions for the Gibbs free energy of the reaction. The reaction power:

$$P_{\text{react}} = \mu_{n_i} \cdot \nu_{n_i}$$

is the power that is available to be converted into either thermal or hydraulic/pneumatic power.

I discovered that there exists an interesting symmetry between the mass balance and the power balance equations:

$$\begin{pmatrix}
\nu_{BR} \\
\nu_{H2} \\
\nu_{HBr}
\end{pmatrix} =
\begin{pmatrix}
-1 & 1 & 0 & 0 & -1 \\
2 & -2 & -1 & 1 & 1 \\
0 & 0 & -1 & 1 & 0
\end{pmatrix}
\begin{pmatrix}
\nu_{BR} \\
\nu_{H2} \\
\nu_{HBr}
\end{pmatrix}
= \begin{pmatrix}
\nu_{H2} \\
\nu_{HBr}
\end{pmatrix}
\begin{pmatrix}
\nu_{BR} \\
\nu_{H2} \\
\nu_{HBr}
\end{pmatrix}
= \begin{pmatrix}
\nu_{BR} \\
\nu_{H2} \\
\nu_{HBr}
\end{pmatrix}
\begin{pmatrix}
\mu_{n_2} \\
\mu_{n_3} \\
\mu_{n_4}
\end{pmatrix}
= \begin{pmatrix}
-1 & 2 & 0 & 0 & 0 \\
1 & -2 & 0 & 0 & 0 \\
0 & 0 & -1 & 1 & -1
\end{pmatrix}
\begin{pmatrix}
\mu_{n_2} \\
\mu_{n_3} \\
\mu_{n_4}
\end{pmatrix}
= \begin{pmatrix}
\mu_{BR} \\
\mu_{H2} \\
\mu_{HBr}
\end{pmatrix}
$$

The two matrices are the transpose of each other. Moreover, they are singular (in most cases, they are not even square). Consequently, the component flow rates can be computed from the reaction flow rates, and the "reaction potentials" can be computed from the component potentials, but not vice versa. My book (Cellier 1990) provides an explanation for this symmetry.

I realize that space limitations forced me to reduce this section to a mere skeleton. My book describes the chemical reaction kinetics problem on more than 60 pages. However, I wanted to present at least the essence of these still unpublished results in this paper.

REFERENCES


**AUTHOR’S BIOGRAPHY**

FRANÇOIS E. CELLIER received his B.S. degree in Electrical Engineering from the Swiss Federal Institute of Technology (ETH) Zürich in 1972, his M.S. degree in Automatic Control in 1973, and his Ph.D. degree in Technical Sciences in 1979, all from the same university. Following his Ph.D., Dr. Cellier worked as a Lecturer at ETH Zürich. He joined the University of Arizona in 1984 as an Associate Professor. Dr. Cellier’s main scientific interests concern modeling and simulation methodology, and the design of advanced software systems for simulation, computer-aided modeling, and computer-aided design. He has designed and implemented the GASP–V simulation package, and he was the designer of the COSY simulation language a modified version of which under the name of SYMOSD has meanwhile become a standard by the British Ministry of Defence. Dr. Cellier has authored or co-authored more than forty technical publications, and he has edited two books. He served as a chairman of the National Organizing Committee (NOC) of the Simulation’75 conference, and as a chairman of the International Program Committee (IPC) of the Simulation’77 and Simulation’80 conferences, and he has also participated in several other NOC’s and IPC’s. He is associate editor of several simulation related journals, and he served as vice-chairman of two committees on standardization of simulation and modeling software.