

## Instruction for CADLIVE Simulator

### Example: *E. coli* heat shock response system

## CMA CONVERSION

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### NOTICE

If there is anything wrong with calculation and you want to kill your job, access the site of <http://kurata01.bse.kyutech.ac.jp/TEST/Life/kill>, and push the kill simulator button.

## 0. Objectives

The problem for DAEs is to find the initial values that are necessary for solving their algebraic equations. We suggest that the steady state concentration of CMA with rapid equilibrium approximation is very close to those of DAEs. This instruction presents how one makes the initial values for DAEs from CMA simulation.

## 1. Login

Following the installation of the CADLIVE Simulator, open <http://kurata01.bse.kyutech.ac.jp/TEST/Life/index.html> on a PC browser to display the screen(Fig. 1-1). Input the user name, subsequently his/her password to start up simulation.

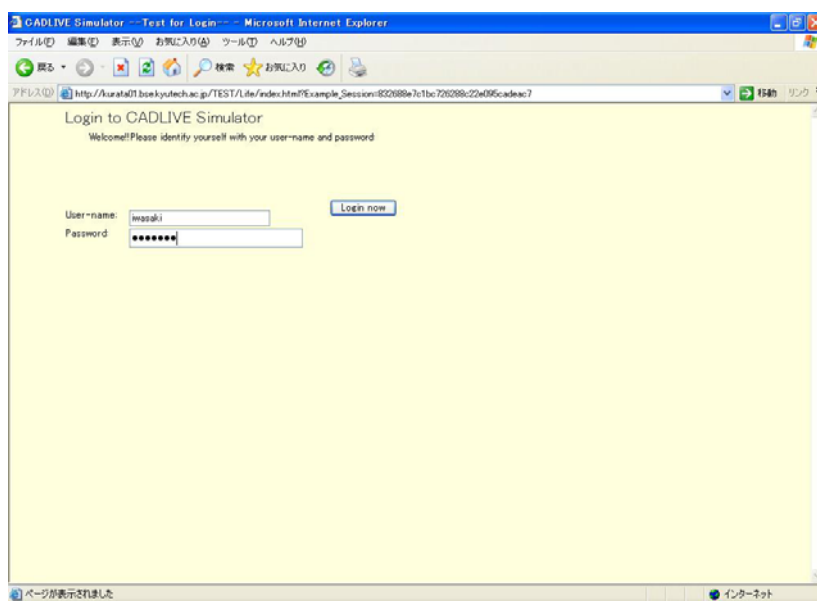


Fig. 1-1 Login screen

## 2. Startup simulator

Clicking the "Simulator" button on the left hand side (Fig. 2-1) changes the upper region of the screen, as shown in Fig.2-2

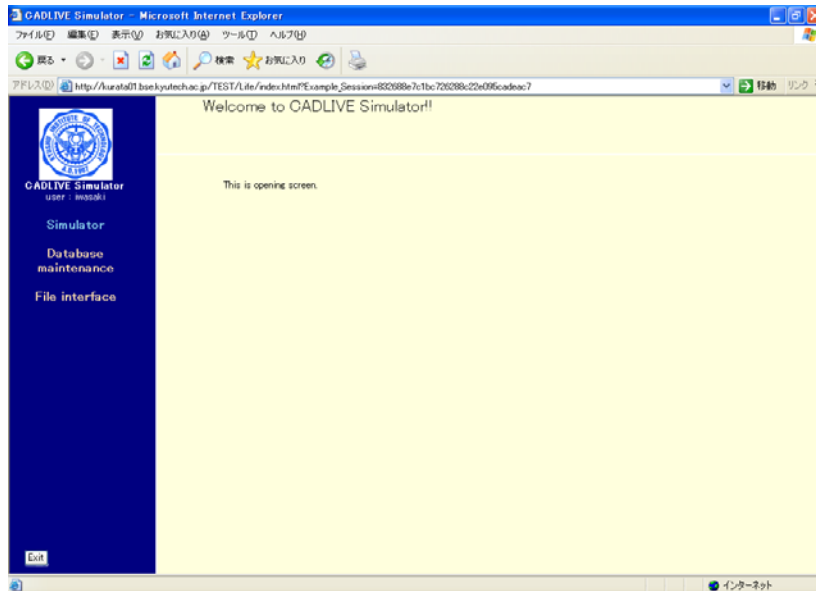


Fig. 2-1Startup menu

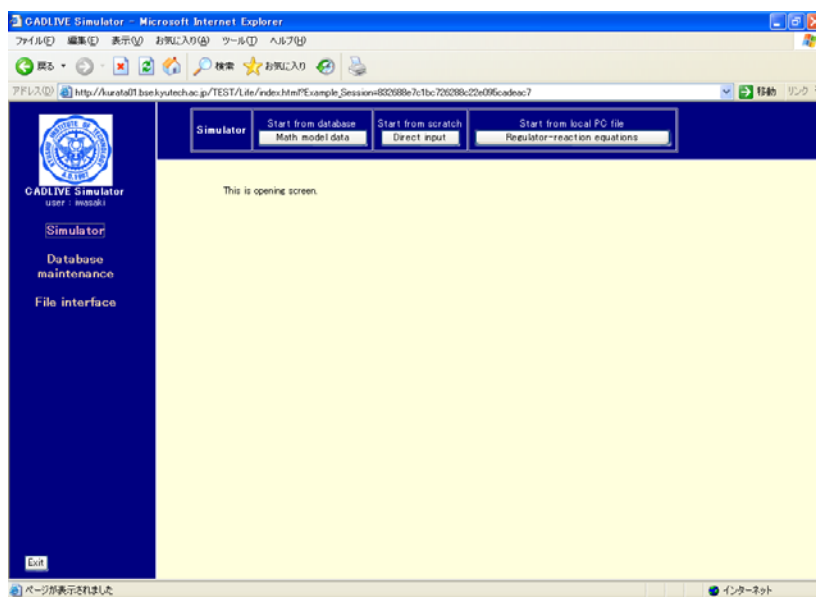


Fig. 2-2 Startup simulation

### 3. Upload of data for regulator-reaction equations

Clicking the [Regulator-reaction equations] button (Fig. 2-2) displays the new screen. Select the XML data file for the regulator-reaction equations (**HSR\_demo.xml**) from users' PC (Fig. 3-1). The data file should be described by using the CADLIVE Editors. Users are required to click the buttons within the screen of CADLIVE, not to carelessly click those of the browser.

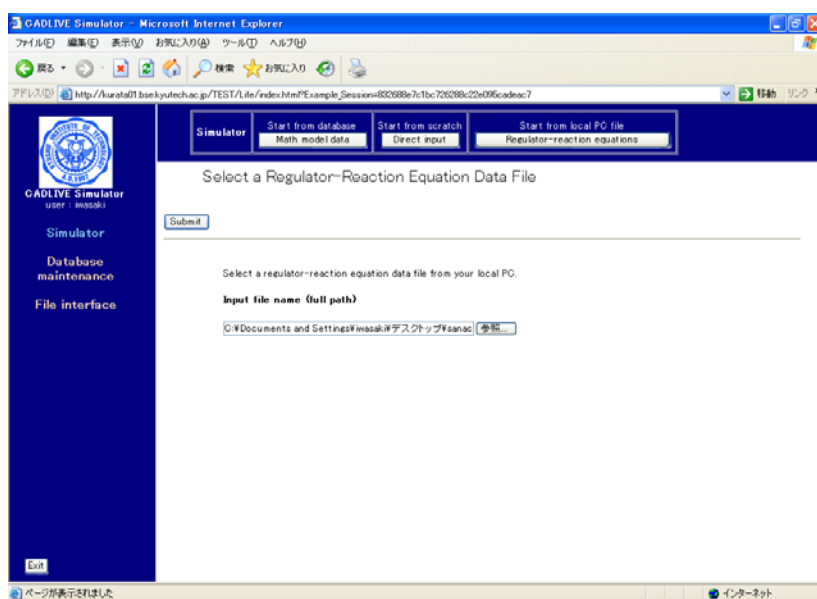


Fig. 3-1 Upload of a regulator-reaction equation file from a PC

#### 4. Selection of conversion methods

The table that indicates the selected regulator-reaction equations appears on the screen (Fig. 4-1). Users choose the conversion method with respect to gene-protein layer, because the heat shock response does not contain the metabolic layer. Select **CMA**.

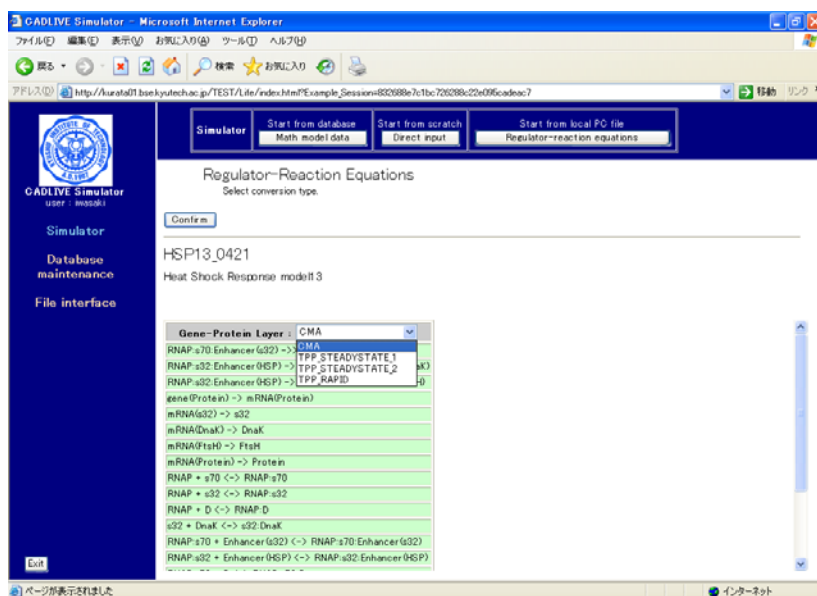


Fig. 4-1 Selection of conversion methods

Following the selection of the conversion method, clicking the [Confirm] button displays the confirmation screen (Fig. 4-2) that is similar to Fig. 4-1.

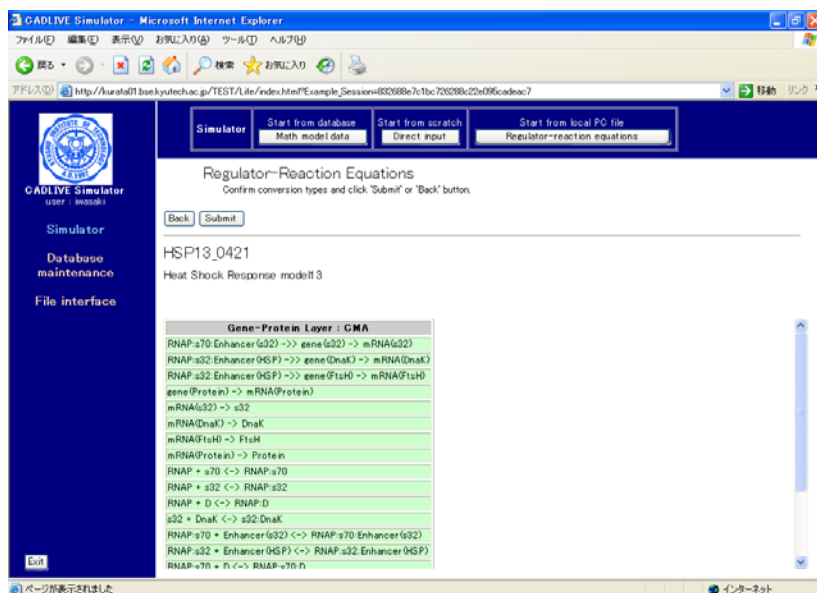


Fig. 4-2 Confirmation screen for the regulator-reaction equations

## 5. Editing Mathematical Model (CMA)

Clicking the [Submit] button on the confirmation screen (Fig. 4-2) parses the regulator-reaction equations, converting them into the mathematical model according to the selected conversion method. The resultant mathematical model is displayed as shown in Fig. 5-1.

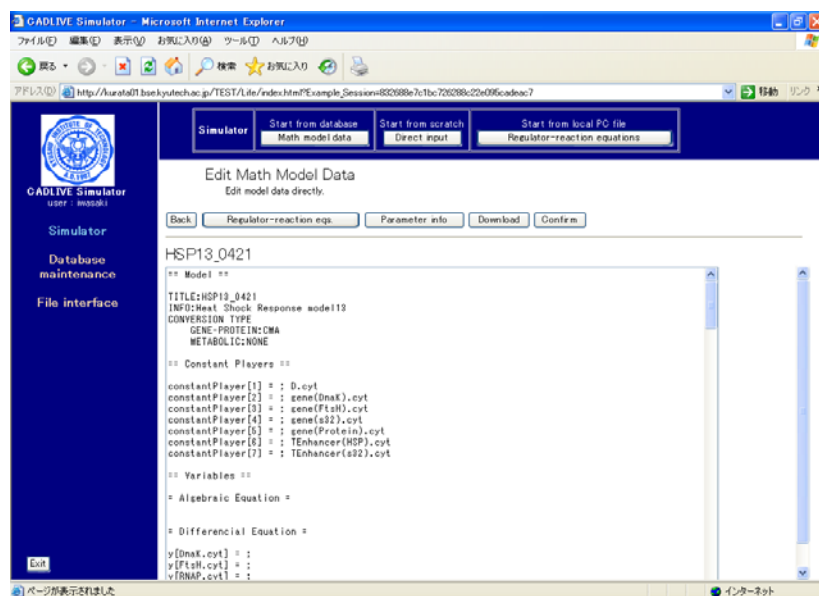


Fig. 5-1 Mathematical model that is obtained from regulator-reaction equations

This screen mainly has seven parts: the header, the definition of constant players, the definition of variables, the definition of the employed parameter labels, the definition of kinetic parameters, the definition of intermediate mathematical expressions, the definition of algebraic equations, and the definition of differential equations.

Clicking the [Regulator-reaction eqs.] button displays another window that shows regulator-reaction equations. Clicking the [Parameter info.] button shows the parameter information. The [Download] button enables users to download various data files to their local PC.

Clicking the [Confirm] button displays the confirmation screen (Fig. 5-2) that is similar to Fig. 5-1. If there is no problem, click the [Submit] button.

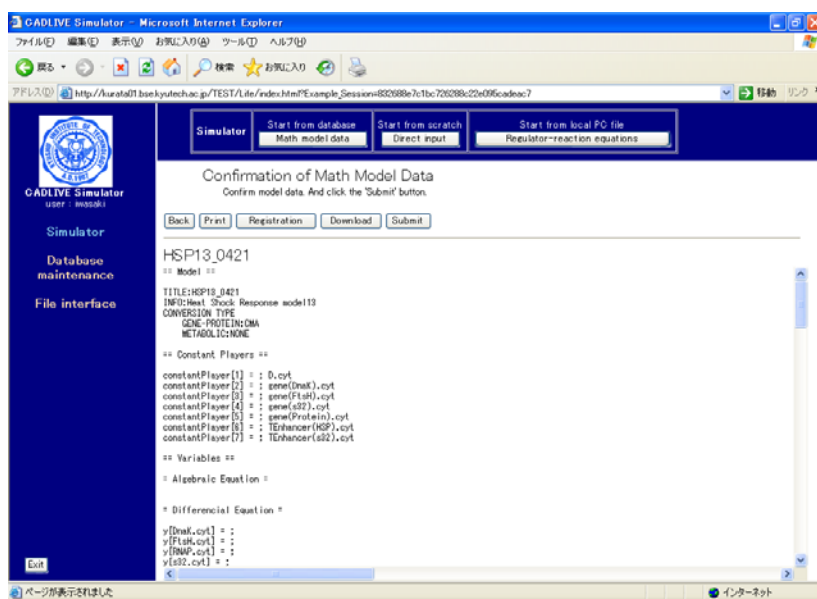


Fig. 5-2 Confirmation screen for a mathematical model

## 6. Selecting analysis types

Clicking the [Submit] button on the confirmation screen displays the screen (Fig. 6-1), where users select the method for numerical simulation. Select “**Dynamic**” as analysis type.

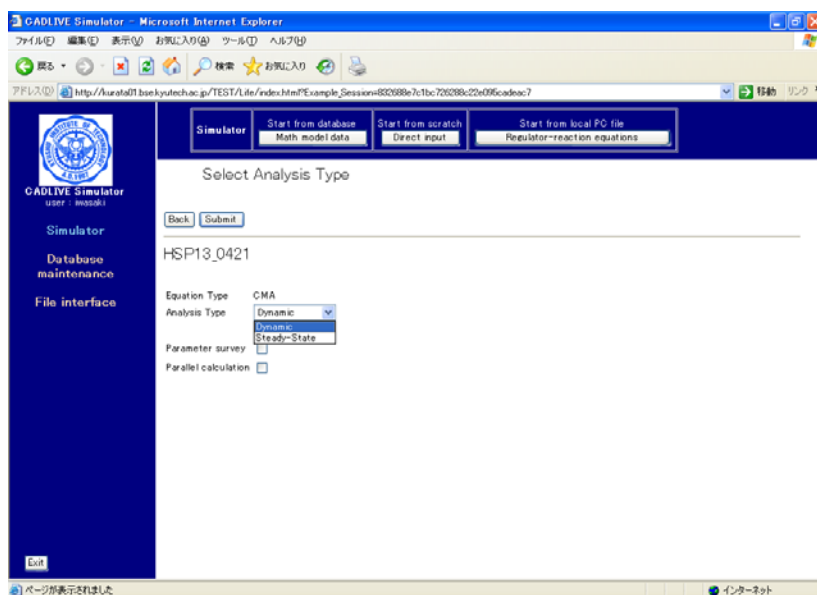


Fig. 6-1 Selection of analysis type

As "Analys type", users can choose either “Dynamic” or “Steady-state”. “Dynamic” simulates the time evolution of the concentrations by calculating DAEs, and “Steady-state” calculates the concentrations at steady state by solving algebraic equations. The checkbox of "Parameter survey" determines if the simulator surveys the parameter space. Checking the checkbox of

"Parallel calculation" carries out parallel calculation that employs the Message Passing Interface (MPI). Notice that the checkbox of "Parallel calculation" cannot be selected prior to checking the parameter survey.

## 7 Input for control data

Clicking the [Submit] button on the screen for selecting analytical type displays the screen for input of the control data (Fig. 7-1). Here, input the data, as provided by the following table

**Table Control parameters**

<b>Solver Type (tolerance)</b>		
NDF	relative: 1e-12	absolute 0.000001
<b>Set time span and time step-size.</b>		
Start time		0
End time		100
(Initial)time step-size		0.0001
Monitoring interval		1
<b>Other</b>		
G-value		<b>1.0</b>
Y default value		0.01

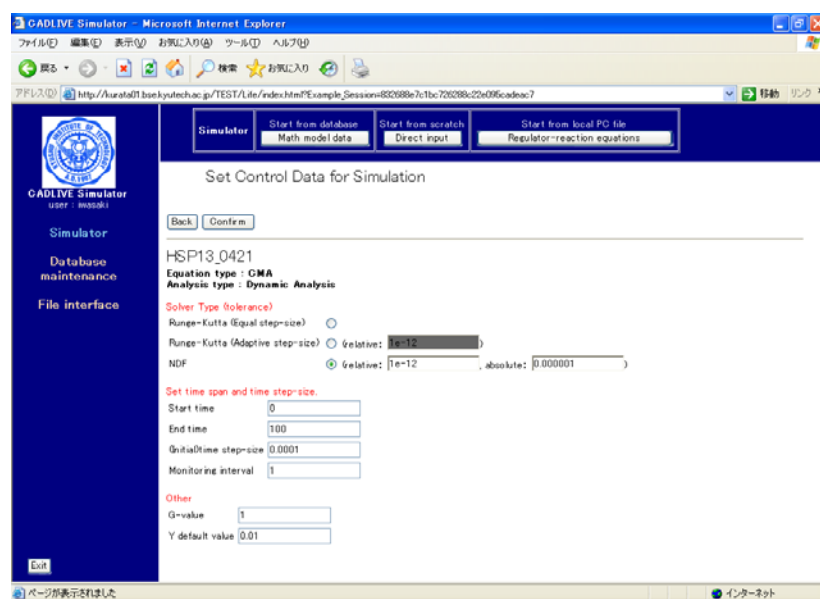


Fig. 7-1 Set control data for simulation

Following data input, clicking the [Confirm] button shows the confirmation screen (Fig. 7-2).



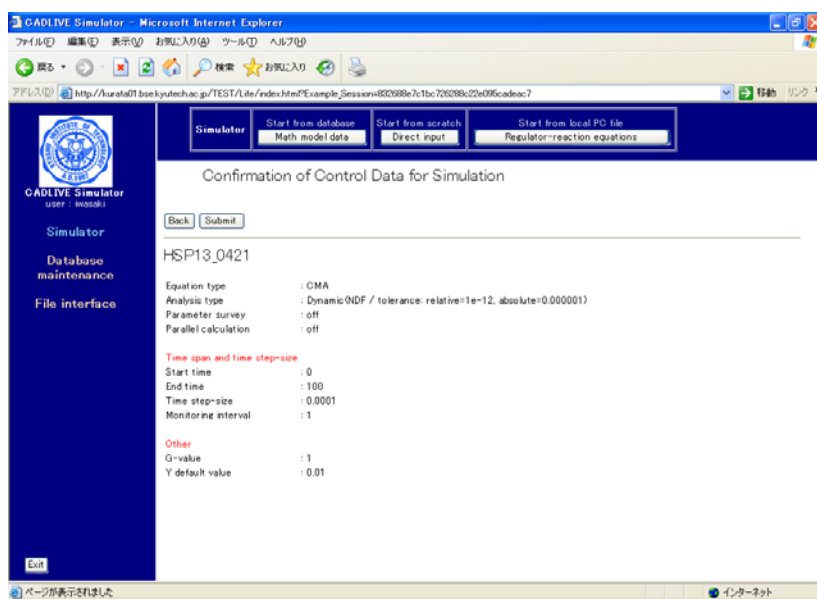


Fig. 7-2 Confirmation screen

## 8. Input of parameters

Following confirming the control data for simulation, clicking the [Submit] button on the screen (Fig. 7-2) displays the new screen (Fig. 8-1), where users input the kinetic parameters and initial values.

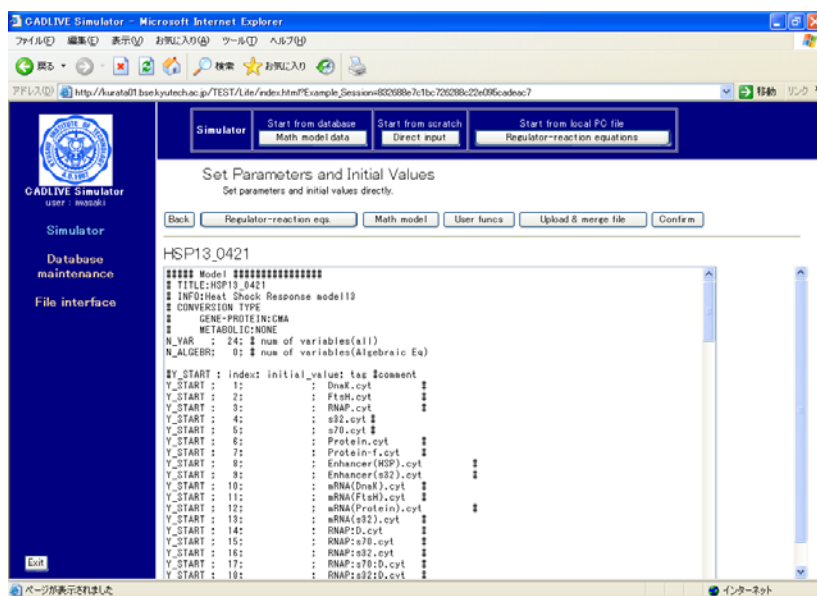


Fig. 8-1 Setting parameters and initial values

In this example, we presented the parameters and initial values that well simulated the heat shock response. Click the [Upload & merge File] button to display the screen (Fig. 8-2).

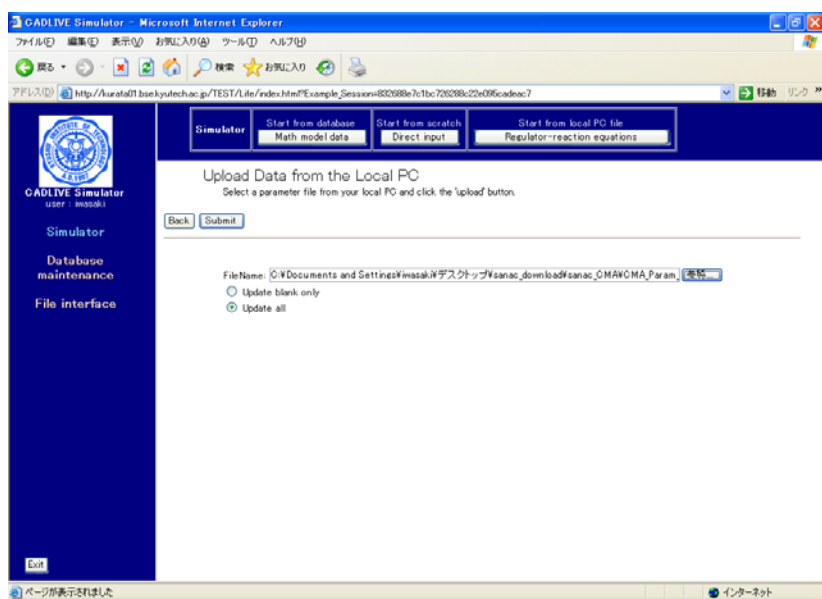


Fig. 8-2 Uploading a parameter file from the local PC

The parameter file that has been made (**CMA\_Param\_HSR.txt**) should be input in the box of "FileName". Check the box of "**Update all**" and click the [Submit] button. The parameters and initial values are automatically input into the mathematical model, as shown in Fig. 8-3

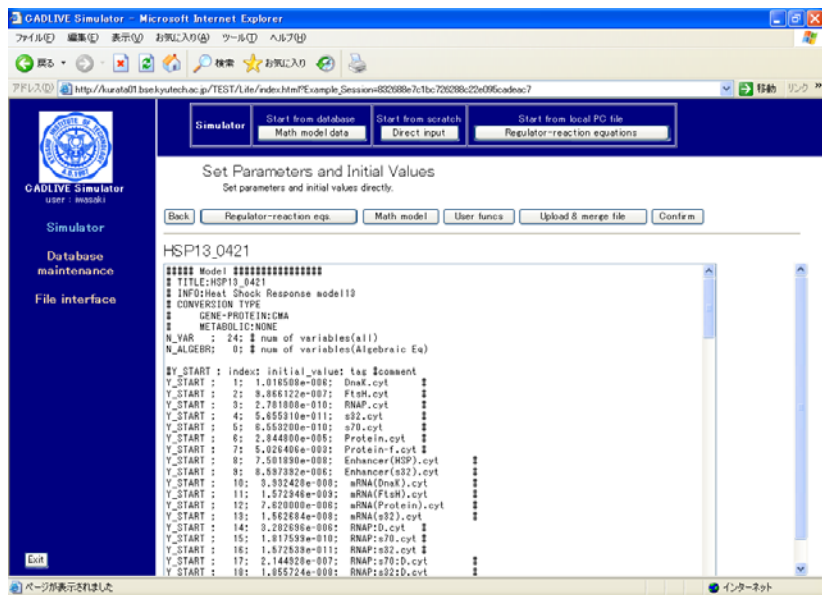


Fig. 8-3 Uploaded parameters and initial values

Following clicking the [Confirm] button, clicking the [Submit] button (Fig. 8-4) starts calculation.



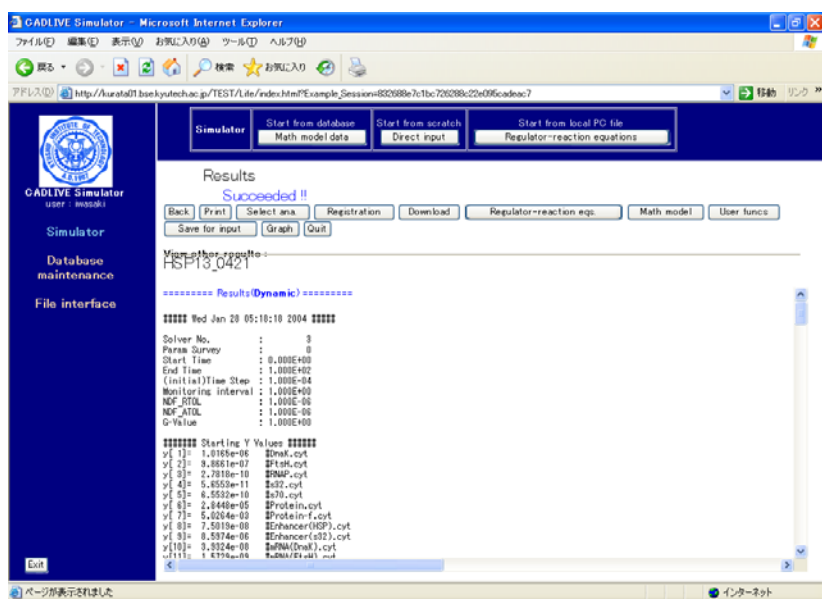


Fig. 9-1 Results

The screens for indicating the results (Fig. 9-1) have the [Registration] button. Clicking the [Registration] button shows the “Registration to the DB” screen (Fig. 9-2) that has also the button of [Registration]. Clicking it saves a series of simulation data as a mathematical model in the database.

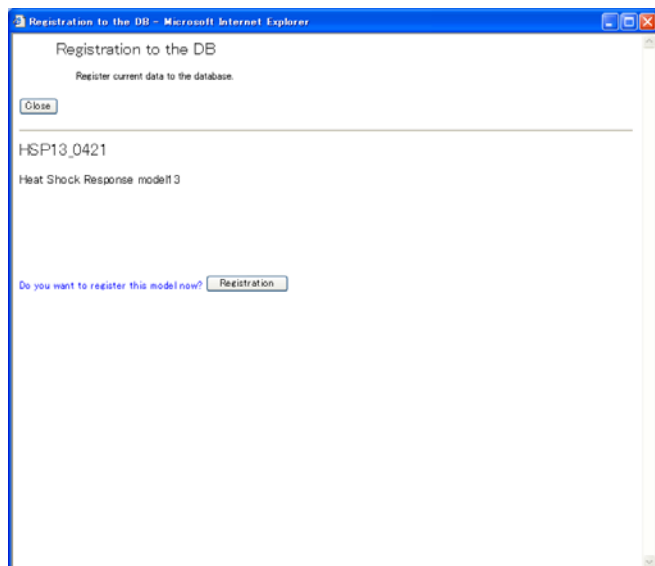


Fig. 9-2 Registration of Data in Database

The screen for results (Fig. 9-2) has the [Download] button. Clicking it displays the download screen (Fig. 9-3). The clicked files are downloaded to a local PC.

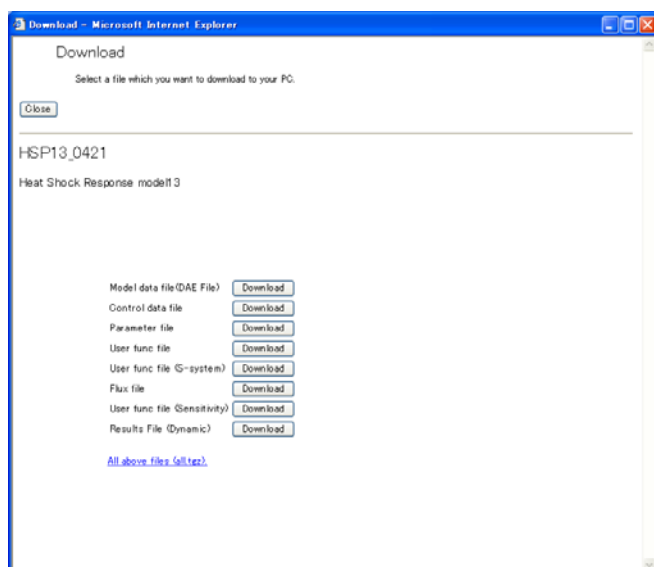


Fig. 9-3 Download

## 10. Graph visualization

In the screen (Fig. 9-1), clicking the [Graph] button displays the window of "Set Graph Data" (Fig. 10-1).

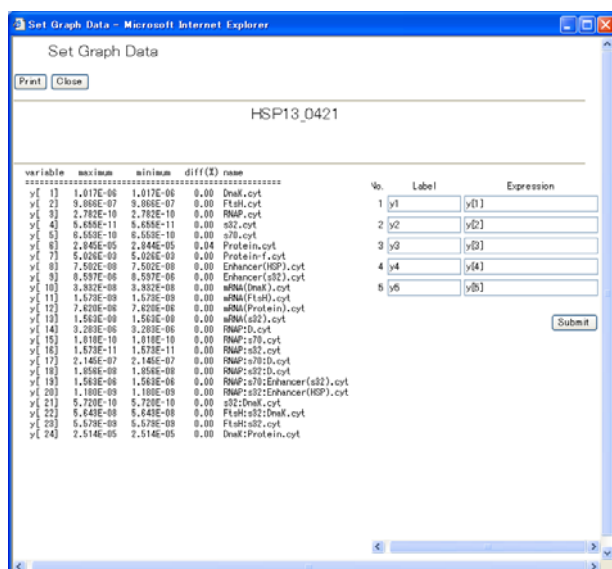


Fig. 10-1. Setting graph data

The right hand side of the screen defines the definition of data that will be plotted on a graph, and can indicate five series of labels and expressions. As default, y1 – y5 and y[1] – y[5] are set as labels and expressions, respectively. Empty labels and expressions are not allowed. The

variables  $y[i]$ , calculation symbols +, -, X, /, parentheses (,), and time T are allowed to be used for describing the mathematical expression. Here, set  $y[1]$ ,  $y[2]$ ,  $y[3]$ ,  $y[4]$ , and  $y[5]$ . After setting them, clicking the [Submit] button displays the screen (Fig. 10-2), where the scaling factor and the regions of X-axis and Y-axis are automatically calculated based on the expression and the minimum/maximum values to visualize the five time-evolution well.

Label	Expression	Maximum	Minimum	Scaling	Color	not draw
y1	y[1]	1.017E-06	1.017E-06	-----	black	<input checked="" type="checkbox"/>
y2	y[2]	9.066E-07	9.066E-07	-----	red	<input checked="" type="checkbox"/>
y3	y[3]	2.782E-10	2.782E-10	1	green	<input type="checkbox"/>
y4	y[4]	5.655E-11	5.655E-11	10	blue	<input type="checkbox"/>
y5	y[5]	5.553E-10	5.553E-10	1	magenta	<input type="checkbox"/>

Title : HSP13\_0421

	X	Y
Factor	100	1E-10
Maximum	1	7
Minimum	0	0
Tick Increment	0.1	1
Label	Time	Concentration
Unit	sec	mol/l
Unit factor	1	1

Fig. 10-2 Setting graph data

Users are able to change the labels, the scaling factors, and the color of the lines. Checking the checkbox of “not draw” omits its time evolution from the graph. The button [Re-calc] recalculates the scaling factors and the regions for the X/Y-axes. The table on the lower screen is set for the X/Y-axes. X indicates time, and Y the concentrations of variables. Clicking the [Draw graph] button (Fig. 10-2) displays the results of time-evolution data, as shown in Fig. 10-3.

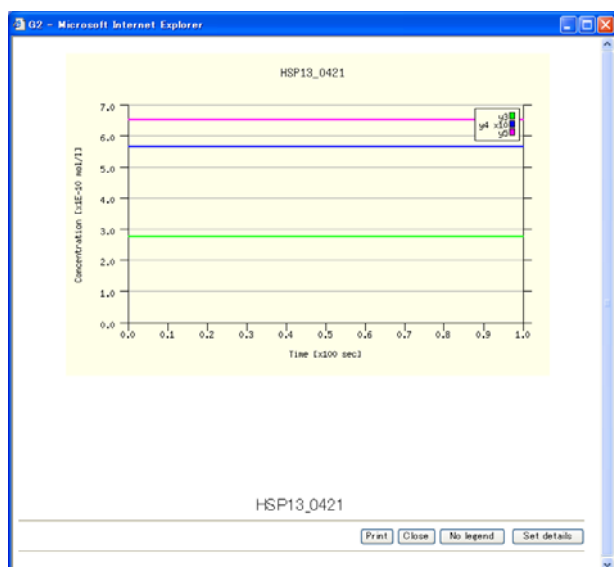


Fig. 10-3 Graph display

The [No legend] button omits the legends. The [Set details] button is able to change the color of the graph and other graph expressions. Clicking the [Set details] button displays the window (Fig.10-4). Users can change the style of the graph by inputting the value in the boxes. The values of the graph size and the location of the legends are determined by using pixel units. The location of the legend is determined by the coordinate of the left and upper legend.

Fig. 10-4 Detailed setting for a graph

## 11 Calculation of the initial values for DAEs (TPP)

The problem for DAEs is to find the initial values that are necessary for solving their algebraic equations. We suggest that the steady state concentration of CMA with rapid equilibrium approximation is very close to those of DAEs. This chapter instructs how one makes the initial values for DAEs from CMA simulation.

### 11.1 Saving the final points of CMA simulation

After the simulation for time courses for CMA, the button of [Save for input] stores the resultant data as the initial values for the subsequent simulation (Fig. 11-1, see Fig. 9-1), which is obtained by the [Initial val.] button.

In Fig. 11-2, click the [Initial val.] button to show the final points of the previously saved time-course data, which are very close to steady state (Fig. 11-2). This button appears only when the initial values are saved in the previous simulation.

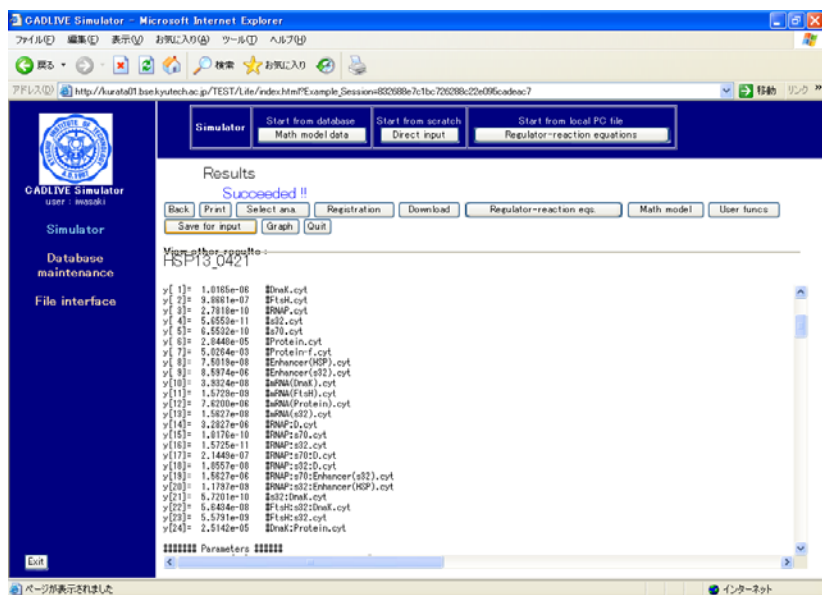


Fig. 11-1 Results of simulation



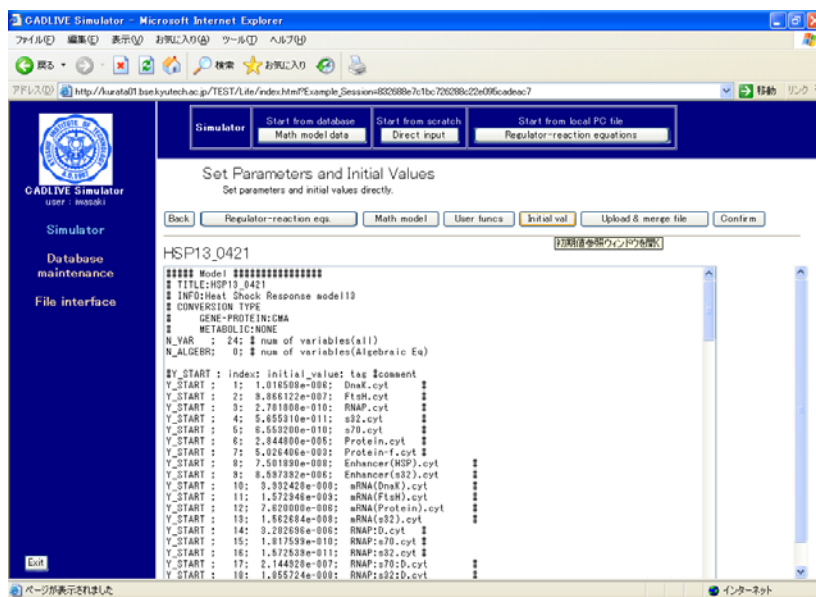


Fig. 11-2 Initial concentrations that are the resultant data of the previous simulation.

## 11.2 Conversion of the final concentration of CMA to the initial concentration for DAEs

Open the window for the initial values from previous simulation (Fig. 11-3). Calculate the total concentrations for specific species (monomer, modified)(Y\_START 25-30) and add them to the file, which can be employed as the initial values for DAEs that are equivalent to CMA.

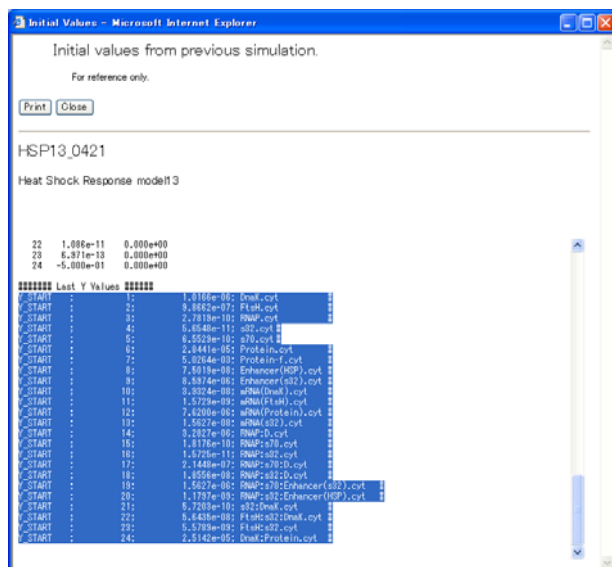


Fig. 11-3 Initial values from previous simulation.

Last Y Values #####		
Y_START :	1;	1.0186e-06; DnaK.cyt
Y_START :	2;	9.8662e-07; FtsH.cyt
Y_START :	3;	2.7818e-10; RNAP.cyt
Y_START :	4;	5.6548e-11; s32.cyt
Y_START :	5;	6.5529e-10; s70.cyt
Y_START :	6;	2.8441e-05; Protein.cyt
Y_START :	7;	5.0284e-03; Protein-f.cyt
Y_START :	8;	7.5019e-08; Enhancer(HSP).cyt
Y_START :	9;	8.5974e-06; Enhancer(s32).cyt
Y_START :	10;	3.9324e-08; mRNA(DnaK).cyt
Y_START :	11;	1.5729e-09; mRNA(FtsH).cyt
Y_START :	12;	7.6200e-06; mRNA(Protein).cyt
Y_START :	13;	1.5627e-08; mRNA(s32).cyt
Y_START :	14;	3.2827e-06; RNAP-D.cyt
Y_START :	15;	1.8176e-10; RNAP-s70.cyt
Y_START :	16;	1.5723e-11; RNAP-s32.cyt
Y_START :	17;	2.1446e-07; RNAP-s70-D.cyt
Y_START :	18;	1.8556e-08; RNAP-s32-D.cyt
Y_START :	19;	1.5627e-06; RNAP-s70:Enhancer(s32).cyt
Y_START :	20;	1.1797e-09; RNAP-s32:Enhancer(HSP).cyt
Y_START :	21;	5.7203e-10; s32:DnaK.cyt
Y_START :	22;	5.6435e-08; FtsH:s32:DnaK.cyt
Y_START :	23;	5.5789e-09; FtsH:s32.cyt
Y_START :	24;	2.5142e-05; DnaK:Protein.cyt
Y_START :	25;	5.000091e-06; TRNAP.cyt
Y_START :	26;	1.778017e-08; Ts70.cyt
Y_START :	27;	8.238990e-08; Ts32.cyt
Y_START :	28;	2.621561e-05; TDnaK.cyt
Y_START :	29;	1.048634e-06; TFtsH.cyt
Y_START :	30;	5.358300e-05; TProtein.cyt

Fig. 11-4. Addition of the calculated total concentrations for specific species Y\_START[25-30].