Model Predictive Control Toolbox

For Use with MATLAB®

Manfred Morari N. Lawrence Ricker

Computation

Visualization

Programming



User's Guide Version 1

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Model Predictive Control Toolbox User's Guide

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Preface

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Tutorial

Introduction

The Model Predictive Control (MPC) Toolbox is a collection of functions (commands) developed for the analysis and design of model predictive control (MPC) systems. Model predictive control was conceived in the 1970s primarily by industry. Its popularity steadily increased throughout the 1980s. At present, there is little doubt that it is the most widely used multivariable control algorithm in the chemical process industries and in other areas. While MPC is suitable for almost any kind of problem, it displays its main strength when applied to problems with:

- A large number of manipulated and controlled variables
- Constraints imposed on both the manipulated and controlled variables
- Changing control objectives and/or equipment (sensor/actuator) failure
- · Time delays

Some of the popular names associated with model predictive control are Dynamic Matrix Control (DMC), IDCOM, model algorithmic control, etc. While these algorithms differ in certain details, the main ideas behind them are very similar. Indeed, in its basic unconstrained form MPC is closely related to linear quadratic optimal control. In the constrained case, however, MPC leads to an optimization problem which is solved on-line in real time at each sampling interval. MPC takes full advantage of the power available in today's control computer hardware.

This software and the accompanying manual are not intended to teach the user the basic ideas behind MPC. Background material is available in standard textbooks like those authored by Seborg, Edgar and Mellichamp (1989)¹, Deshpande and Ash (1988)² and the monograph devoted solely to this topic authored by Morari and coworkers (Morari et al., 1994)³. This section provides a basic introduction to the main ideas behind MPC and the specific form of implementation chosen for this toolbox. The algorithms used here are consistent with those described in the monograph by Morari et al. Indeed, the software is meant to accompany the monograph and vice versa. The routines included in the MPC Toolbox fall into two basic categories: routines which use

- D.E. Seborg, T.F. Edgar, D.A. Mellichamp; Process Dynamics and Control; JohnWiley & Sons, 1989
- P.B. Deshpande, R.H. Ash; Computer Process Control with Advanced Control Applications, 2nd ed., ISA, 1988
- 3. M. Morari, C.E. Garcia, J.H. Lee, D.M. Prett; *Model Predictive Control*, Prentice Hall, 1994 (In the process of being written.)

a step response model description and routines which use a state-space model description. In addition, simple identification tools are provided for identifying step response models from plant data. Finally, there are also various conversion routines which convert between different model formats and analysis routines which can aid in determining the stability of the unconstrained system, etc. All MPC Toolbox commands have a built-in usage display. Any command called with no input arguments results in a brief description of the command line. For example, typing mpccon at the command line gives the following:

```
usage: Kmpc = mpccon(model, ywt, uwt, M, P)
```

The following sections include several examples. They are available in the tutorial programs mpctut. m, mpctutid. m, mpctutst. m, and mpctutss. m. You can copy these demo files from the mpctool s/mpcdemos source into a local directory and examine the effects of modifying some of the commands.

Target Audience for the MPC Toolbox

The package is intended for the classroom and for the practicing engineer. It can assist in communicating the concepts of MPC to a student in an introductory control course. At the same time it is sophisticated enough to allow an engineer in industry to evaluate alternate control strategies in simulation studies.

System Requirements

The MPC Toolbox assumes the following operating system requirements:

- MATLAB® is running on your system.
- If nonlinear systems are to be simulated, Simulink® is required for the functions nl cmpc and nl mpcsi m.
- If the *theta* format from the System Identification Toolbox is to be used to create models in the MPC *mod* format (using the MPC Toolbox function, th2mod), then the System Identification Toolbox function polyform and the Control System Toolbox function append are required.

The MPC Toolbox analysis and simulation algorithms are numerically intensive and require approximately 1MB of memory, depending on the number of inputs and outputs. The available memory on your computer may limit the size of the systems handled by the MPC Toolbox.

Note: there is a pack command in MATLAB that can help free memory space by compacting fragmented memory locations. For reasonable response times, a computer with power equivalent to an 80386 machine is recommended unless only simple tutorial example problems are of interest.

MPC Based on Step Response Models

Step Response Models

Step response models are based on the following idea. Assume that the system is at rest. For a linear time-invariant single-input single-output (SISO) system let the output change for a unit input change Δv be given by

$$\{0, s_1, s_2, \ldots, s_n, s_n, \ldots\}$$

Here we assume that the system settles exactly after n steps. The step response $\{s_1, s_2, \ldots, s_n\}$ constitutes a complete model of the system, which allows us to compute the system output for any input sequence:

$$y(k) = \sum_{i=1}^{n} s_{i} \Delta v(k-i) + s_{n} v(k-n-1)$$

Step response models can be used for both stable and integrating processes. For an integrating process it is assumed that the slope of the response remains constant after n steps, i.e.,

$$s_n - s_{n-1} = s_{n+1} - s_n = s_{n+2} - s_{n+1} = \dots$$

For a multi-input multi-output (MIMO) process with n_{v} inputs and n_{y} outputs, one obtains a series of step response coefficient matrices

$$S_{i} = \begin{bmatrix} s_{1, 1, i} & s_{1, 2, i} & \dots & s_{1, n_{v}, i} \\ s_{2, 1, i} & & & & \\ \vdots & & & & & \\ s_{n_{y}, 1, i} & s_{n_{y}, 2, i} & \dots & s_{n_{y}, n_{v}, i} \end{bmatrix}$$

where $s_{l,m,i}$ is the i^{th} step response coefficient relating the m^{th} input to the l^{th} output.

The MPC Toolbox stores step response models in the following format:

$$\text{plant} = \begin{bmatrix} S_1 \\ S_2 \\ \vdots \\ S_n \\ \text{nout}(1) & 0 & \dots & 0 \\ \text{nout}(2) & 0 & \dots & 0 \\ \vdots & \vdots & & & \vdots \\ \text{nout}(\mathbf{n}_y) & 0 & \dots & 0 \\ n_y & 0 & \dots & 0 \\ \text{delt2} & 0 & \dots & 0 \end{bmatrix}_{(n \cdot n_v + n_v + 2) \times n_v}$$

where delt2 is the sampling time and the vector nout indicates if a particular output is integrating or not:

nout(i) = 0 if output i is integrating.
nout(i) = 1 if output i is stable.

The step response can be obtained directly from identification experiments, or generated from a continuous or discrete transfer function or state-space model. For example, if the discrete system description (sampling time T = 0.1) is

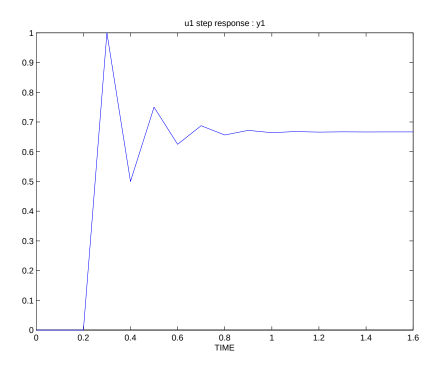
$$y(k) = -0.5y(k-1) + v(k-3)$$

then the transfer function is

$$g(z) = \frac{z^{-3}}{1 + 0.5z^{-1}}$$

The following commands (see mpctut. m) generate the step response model for this system and plot it:

```
num = 1;
den = [1 0.5];
delt1 = 0.1;
del ay = 2;
g = poly2tfd(num, den, delt1, delay);
% Set up the model in tf format
tfinal = 1.6;
delt2 = delt1;
nout = 1;
plant = tfd2step(tfinal, delt2, nout, g);
% Calculate the step response
plotstep(plant) % Plot the step response
```



Alternatively, we could first generate a state-space description applying the command tf2ss and then generate the step response with ss2step. In this case, we need to pad the numerator and denominator polynomials to account for the time delay.

```
num = [0 0 0 num];
den = [den 0 0];
[phi,gam,c,d] = tf2ss(num,den); % Convert to state-space
plant = ss2step(phi,gam,c,d,tfinal,delt1,delt2,nout);
% Calculate step response
```

We can get some information on the contents of a matrix in the MPC Toolbox via the command mpci nfo. For our example, mpci nfo(pl ant) returns:

```
This is a matrix in MPC Step format.

sampling time = 0.1

number of inputs = 1

number of outputs = 1

number of step response coefficients = 16

All outputs are stable.
```

Model Identification

The identification routines available in the MPC Toolbox are designed for multi-input single-output (MISO) systems. Based on a historical record of the output $y_l(k)$ and the inputs $v_1(k)$; $v_2(k)$, . . . , $v_{n_v}(k)$,

$$yy_l = \begin{bmatrix} y_l(1) \\ y_l(2) \\ y_l(3) \\ \vdots \end{bmatrix} \qquad v = \begin{bmatrix} v_1(1) & v_2(1) & \dots & v_{n_v}(1) \\ v_1(2) & v_2(2) & \dots & v_{n_v}(2) \\ v_1(3) & v_2(3) & \dots & v_{n_v}(3) \\ \vdots & \vdots & & & \end{bmatrix}$$

the step response coefficients

$$\begin{bmatrix} s_{l,\,1,\,1} & s_{l,\,2,\,1} & \cdots & s_{l,\,n_{v},\,1} \\ s_{l,\,1,\,2} & s_{l,\,2,\,2} & \cdots & s_{l,\,n_{v},\,2} \\ \vdots & & & & \\ s_{l,\,1,\,i} & s_{l,\,2,\,i} & \cdots & s_{l,\,n_{v},\,i} \\ \vdots & & & & \vdots \\ \vdots & & & & \vdots \\ \end{bmatrix}$$

are estimated. For the estimation of the step response coefficients we write the SISO model in the form ${\bf r}$

$$\Delta y(k) = \sum_{i=1}^{n} h_i \Delta v(k-i)$$

where

$$\Delta y(k) = y(k) - y(k-1)$$

and

$$h_i = s_i - s_{i-1}$$

 h_i are the impulse response coefficients. This model allows the designer to present all the input (v) and $\operatorname{output}(y_i)$ information in deviation form, which is often desirable. If the particular output is integrating, then the model

$$\Delta(\Delta y(k)) = \sum_{i=1}^{n} \Delta h_i \Delta v(k-i)$$

where

$$\Delta(\Delta y(k)) = \Delta y(k) - \Delta y(k-1)$$

$$\Delta h_i = h_i - h_{i-1}$$

should be used to estimate Δh_i , and thus h_i and s_i are given by

$$h_i = \sum_{k=1}^{i} \Delta h_k$$

$$s_i = \sum_{j=1}^{i} h_j = \sum_{j=1}^{i} \sum_{k=1}^{j} \Delta h_k$$

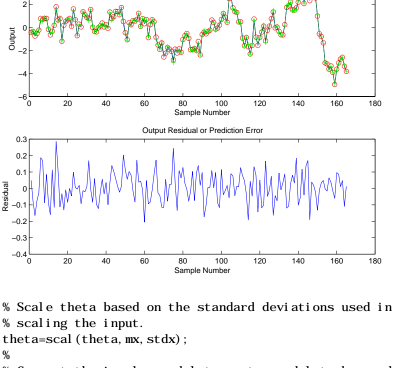
For parameter estimation it is usually recommended to scale all the variables such that they are the same order of magnitude. This may be done via the MPC Toolbox functions autosc or scal. Then the data has to be arranged into the form

$$Y = X\Theta$$

where Y contains all the output information $(\Delta y(k))$ for stable and $\Delta(\Delta y(k))$ for integrating outputs) and X all the input information $(\Delta v(k))$ appropriately arranged. Θ is a vector including all the parameters to be estimated (h_i) for stable and Δh_i for integrating outputs). This rearrangement of the input and output information is handled by wrtreg. The parameters Θ can be estimated via multivariable least squares regression (ml r) or partial least squares (pl sr). Finally, the step response model is generated from the impulse response coefficients via i mp2step. The following example (see mpctutid) illustrates this procedure.

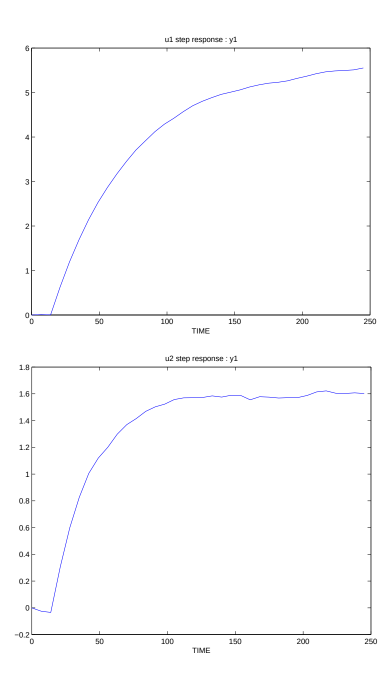
Example:

```
% Load the input and output data. The input and output
% data are generated from the following transfer
% functions and random zero-mean noises.
% TF from input 1 to output 1: g11=5.72exp(-14s)
\% (60s+1)
% TF from input 2 to output 1: g12=1.52exp(-15s)
% (25s+1)
% Sampling time of 7 minutes was used.
% load mlrdat
% Determine the standard deviations for input data using
% autosc.
[ax, mx, stdx] = autosc(x);
% Scale the input data by their standard deviations only.
mx=0*mx;
sx=scal(x, mx, stdx);
% Put the input and output data in a form such that they
% can be used to determine the impulse response
% coefficients. 35 coefficients (n) are used.
n=35:
[xreg, yreg] = wrtreg(sx, y, n);
% Determine the impulse response coefficients via mlr.
% By specifying plotopt=2, two plots-plot of predicted
% output and actual output, and plot of the output
% residual (or prediction error)-are produced.
ni nput=2; pl otopt=2;
[theta, yres] = mlr(xreg, yreg, ni nput, pl otopt);
```



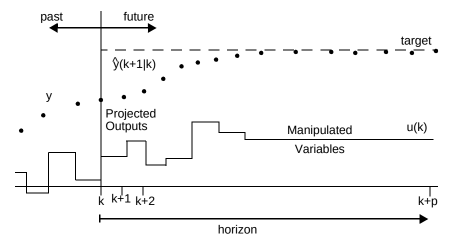
Actual value (o) versus Predicted Value (+)

```
% scaling the input.
theta=scal(theta, mx, stdx);
%
% Convert the impulse model to a step model to be used
% in MPC design.
% Sampling time of 7 minutes was used in determining
% the impulse model.
% Number of outputs (1 in this case) must be specified.
nout=1;
delt=7;
model=imp2step(delt, nout, theta);
%
% Plot the step response.
plotstep(model)
```



Unconstrained Model Predictive Control

The MPC control law can be most easily derived by referring to the following figure.



For any assumed set of present and future control moves $\Delta u(k)$, $\Delta u(k+1)$, . . . , $\Delta u(k+m-1)$ the future behavior of the process outputs $y(k+1 \mid k)$, $y(k+2 \mid k)$, . . . , $y(k+p \mid k)$ can be predicted over a horizon p. The m present and future control moves $(m \leq p)$ are computed to minimize a quadratic objective of the form

$$\min_{\Delta u(k)...\Delta u(k+m-1)} \sum_{l=1}^{p} \left\| \Gamma_{l}^{y}([y(k+l|\mathbf{k}) - r(k+l)]) \right\|^{2} + \sum_{l=1}^{m} \left\| \Gamma_{l}^{u}[\Delta u(k+l-1)] \right\|^{2}$$

Here Γ_l^y and Γ_l^u are weighting matrices to penalize particular components of y or u at certain future time intervals. r(k+l) is the (possibly time-varying) vector of future reference values (setpoints). Though m control moves are calculated, only the first one $(\Delta u(k))$ is implemented. At the next sampling interval, new values of the measured output are obtained, the control horizon is shifted forward by one step, and the same computations are repeated. The resulting control law is referred to as "moving horizon" or "receding horizon."

The predicted process outputs $y(k+1|k),\ldots,y(k+p|k)$ depend on the current measurement $\hat{y}(k)$ and assumptions we make about the unmeasured disturbances and measurement noise affecting the outputs. The MPC Toolbox assumes that the unmeasured disturbances for each output are steps passing through a first order lag with time constant tfilter(2,:). For rejecting measurement noise, the time constant of an exponential filter tfilter(1,:) can be specified by the user. (It can be shown that this procedure is optimal for white noise disturbances passed through an integrator and a first order lag, and white measurement noise). For conventional Dynamic Matrix Control (DMC) the disturbance time constant is assumed to be zero (tfilter(2,:) = zeros(1,ny)), i.e., the unmeasured disturbances have the form of steps, and the noise filter time constant is also set to zero (tfilter(1,:) = zeros(1,ny)), i.e., there is no measurement noise filtering for doing the prediction.

Under the stated assumptions, it can be shown that a linear time-invariant feedback control law results

$$\Delta u(k) = K_{MPC} E_p(k+1 \mid k)$$

where $E_p(k+1 \mid k)$ is the vector of predicted future errors over the horizon p which would result if all present and future manipulated variable moves were equal to zero $\Delta u(k) = \Delta u(k+1) = \ldots = 0$.

For open-loop stable plants, nominal stability of the closed-loop system depends only on K_{MP} C which in turn is affected by the horizon p, the number of moves m and the weighting matrices Γ_l^y and Γ_l^u . No precise conditions on m, p, Γ_l^y and Γ_l^u exist which guarantee closed-loop stability. In general, decreasing m relative to p makes the control action less aggressive and tends to stabilize a system. For p=1, nominal stability of the closed-loop system is guaranteed for any finite m, and time-invariant input and output weights. More commonly, Γ_l^u is used as a tuning parameter. Increasing Γ_l^u always has the effect of making the control action less aggressive.

The noise filter time constant tfilter(1,:) and the disturbance time constant tfilter(2,:) do not affect closed-loop stability or the response of the system to setpoint changes or measured disturbances. They do, however, affect the robustness and the response to unmeasured disturbances.

^{1.} See cmpc in the "Reference" section for details on how to specify tfilter.

Increasing the noise filter time constant makes the system more robust and the unmeasured disturbance response more sluggish. Increasing the disturbance time constant increases the lead in the loop, making the control action somewhat more aggressive, and is recommended for disturbances which have a slow effect on the output.

All controllers designed with the MPC Toolbox track steps asymptotically error-free (Type 1). If the unmeasured disturbance model or the system itself is integrating, ramps are also tracked error-free (Type 2).

Example: (see mpctutst. m)

```
% Plant transfer function: g=5.72exp(-14s)/(60s+1)
% Disturbance transfer function: gd=1.52exp(-15s)/
% (25s+1)
% Build the step response models for a sampling period
% of 7.
del t1=0;
del av1=14;
num1=5.72;
den1=[60 1];
g=poly2tfd(num1, den1, delt1, del ay1);
tfinal =245;
delt2=7:
nout 1=1;
pl ant=tfd2step(tfi nal, del t2, nout1, g);
del ay2=15;
num2=1.52;
den2=[25 1];
gd=poly2tfd(num2, den2, delt1, delay2);
delt2=7:
nout2=1;
dpl ant=tfd2step(tfi nal, del t2, nout2, gd);
% Calculate the MPC controller gain matrix for
% No plant/model mismatch,
% Output Weight=1, Input Weight=0
% Input Horizon=5, Output Horizon=20
model =pl ant;
ywt=1; uwt=0;
```

```
M=5; P=20;
Kmpc1=mpccon(model, ywt, uwt, M, P);
% Simulate and plot response for unmeasured and measured
% step disturbance through dplant.
tend=245;
r=[ ]; usat=[ ]; tfilter=[ ];
dmodel =[ ];
dstep=1;
[y1, u1] = mpcsi m(pl ant, model, Kmpc1, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
dmodel =dpl ant; % measured disturbance
[y2, u2] = mpcsi m(plant, model, Kmpc1, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
plotall([y1, y2], [u1, u2], delt2);
pause; % Perfect rejection for measured disturbance case.
                           Outputs
8.0
0.6
0.4
0.2
             50
                                  150
                                             200
                                                        250
                            Time
                        Manipulated Variables
-0.1
-0.2
-0.3
-0.4
-0.5
-0.6
-0.7 L
                       100
                                  150
                                             200
                            Time
% Calculate a new MPC controller gain matrix for
% No plant/model mismatch,
% Output Weight=1, Input Weight=10
```

```
% Input Horizon=5, Output Horizon=20
model =pl ant;
ywt=1; uwt=10;
M=5; P=20;
mpc2=mpccon(model, ywt, uwt, M, P);
% Simulate and plot response for unmeasured and measured
% step disturbance through dplant.
tend=245;
r=[ ]; usat=[ ]; tfilter=[ ];
dmodel = [ ];
dstep=1;
[y3, u3] = mpcsi m(plant, model, Kmpc2, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
dmodel =dpl ant; % measured disturbance
[y4, u4] = mpcsi m(plant, model, Kmpc2, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
plotall([y3, y4], [u3, u4], delt2);
pause;
                              Outputs
 0.8
 0.6
 0.4
 0.2
 -0.2 L
              50
                         100
                                    150
                                                200
                                                           250
                              Time
                         Manipulated Variables
-0.05
 -0.1
-0.15
 -0.2
-0.25
 -0.3
-0.35 L
                         100
                                    150
                                                200
                                                           250
                              Time
```

```
% Simulate and plot response for unmeasured
% step disturbance through dplant with uwt=0,
% with and without noise filtering.
tend=245;
r=[ ]; usat=[ ]; dmodel=[
tfilter=[ ];
dstep=1;
[y5, u5] = mpcsi m(pl ant, model, Kmpc1, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
tfilter=20; % noise filtering time constant=20
[y6, u6] = mpcsim(plant, model, Kmpc1, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
plotall([y5, y6], [u5, u6], delt2);
pause;
                             Outputs
0.8
0.6
0.4
0.2
 0
0
             50
                        100
                                               200
                                   150
                                                          250
                             Time
                         Manipulated Variables
-0.1
-0.2
-0.3
-0.4
-0.5
-0.6
-0.7
                        100
                                    150
                                               200
                             Time
```

```
% Simulate and plot response for unmeasured
% step disturbance through dplant with uwt=0,
% with and without unmeasured disturbance time constant
% being specified.
tend=245;
r=[ ]; usat=[ ]; dmodel=[ ];
tfilter=[ ];
dstep=1;
[y7, u7] = mpcsim(plant, model, Kmpc1, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
tfilter=[0;25]; % unmeasured disturbance time constant=25
[y8, u8] = mpcsi m(plant, model, Kmpc1, tend, r, usat, tfilter, ...
    dpl ant, dmodel, dstep);
plotall([y7, y8], [u7, u8], delt2);
pause;
                            Outputs
0.8
0.6
0.4
0.2
 ٥L
                        100
                                              200
                                                         250
                                   150
                             Time
                        Manipulated Variables
-0.5
             50
                        100
                                   150
                                              200
                                                         250
                             Time
```

Closed-Loop Analysis

Apart from simulation, other tools are available in the MPC Toolbox to analyze the stability and performance of a closed-loop system. We can obtain the state-space description of the closed-loop system with the command <code>mpccl</code> and then determine the pole locations with <code>smpcpole</code>.

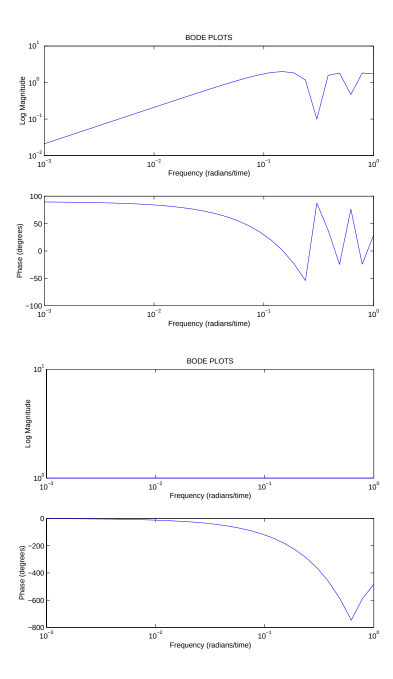
Example: (mpctutst. m)

```
% Construct a closed-loop system for no disturbances
% and uwt = 0. Determine the poles of the system.
cl mod = mpccl(plant, model, Kmpc1);
poles = smpcpole(cl mod);
maxpole = max(poles)
Result is: maxpole = 1.0
```

The closed-loop system is stable if all the poles are inside or on the unit-circle. Furthermore we can examine the frequency response of the closed-loop system. For multivariable systems, singular values as a function of frequency can be obtained using svdfrsp.

Example: (mpctutst. m)

```
% Calculate and plot the frequency response of the % sensitivity and complementary sensitivity functions. freq = [-3,0,30]; ny = 1; in = [1:ny]; % input is r for comp. sensitivity out = [1:ny]; % output is yp for comp. sensitivity [frsp, eyefrsp] = mod2frsp(cl mod, freq, out, in); plotfrsp(eyefrsp); % sensitivity pause; plotfrsp(frsp); % complementary sensitivity pause; % Magnitude = 1 for all frequencies chosen.
```



Constrained Model Predictive Control

The control action can also be computed subject to hard constraints on the manipulated variables and the outputs.

Manipulated variable constraints:

$$u_{min}(l) \le u(k+l) \le u_{max}(l)$$

Manipulated variable rate constraints:

$$|\Delta u(k+l)| \leq \Delta u_{max}(l)$$

Output variable constraints:

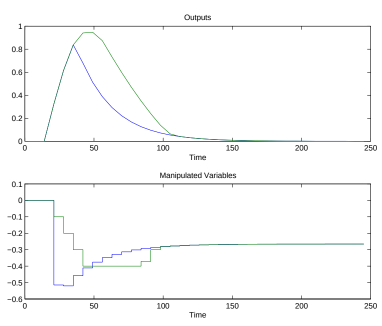
$$y_{min}(l) \le y(k+1|k) \le y_{max}(l)$$

When hard constraints of this form are imposed, a quadratic program has to be solved at each time step to determine the control action and the resulting control law is generally nonlinear. The performance of such a control system has to be evaluated via simulation.

Example: (mpctutst. m)

```
% Simulate and plot response for unmeasured step
% disturbance through dplant with and without
% input constraints.
% No plant/model mismatch,
% Output Weight = 1, Input Weight = 0
% Input Horizon = 5, Output Horizon = 20
% Minimum Constraint on Input = -0.4
% Maximum Constraint on Input = inf
\% Delta Constraint on Input = 0.1
model = plant;
ywt = 1; uwt = 0;
M = 5; P = 20;
tend = 245:
r = 0:
ulim = [];
ylim = [ ]; tfilter = [ ]; dmodel = [ ];
dstep = 1;
[y9, u9] = cmpc(plant, model, ywt, uwt, M, P, tend, r, ...
   ulim, ylim, tfilter, dplant, dmodel, dstep);
ulim = [-0.4, inf, 0.1]; % impose constraints
```

 $\label{eq:continuous} \begin{array}{ll} [y10,\,u10] &=& cmpc(pl\,ant,\,model\,,\,ywt,\,uwt,\,M,\,P,\,tend,\,r,\,\ldots\,\\ &u\,l\,i\,m,\,yl\,i\,m,\,tfi\,l\,ter,\,dpl\,ant,\,dmodel\,,\,dstep)\,;\\ pl\,otal\,l\,(\,[y9,\,y10]\,,\,[u9,\,u10]\,,\,del\,t2)\,; \end{array}$



Application: Idle Speed Control

Process Description

An idle speed control² system should maintain the desired engine rpm with automatic transmission in neutral or drive. Despite sudden load changes due to the actions of air conditioning, power steering, etc., the control system should maintain smooth stable operation. Because of varying operating conditions and engine-to-engine variability inevitable in mass production, the system dynamics may change. The controller must be designed to be *robust* with respect to these changes. Two control inputs, bypass valve (or throttle) opening and spark advance, are used to maintain the engine rpm at a desired idle speed level. For safe operation, spark advance should not change by more than 20 degrees. Also, spark advance should be at the design point at steady-state for fuel economy. Thus, spark advance is viewed both as a manipulated input and a controlled output.

Control Problem Formulation

Here we consider two different operating conditions (transmission in neutral and drive positions) and the models for the two plants are taken from the paper by Hrovat and Bodenheimer. The goal is to design a model predictive controller such that the closed loop performance at both operating conditions is good in the presence of the input constraint specified above. There is no synthesis method available which systematically generates a controller design which guarantees *robust* performance (or even just robust stability) in the presence of constraints. Thus, we must rely on both design and simulation tools to determine achievable performance objectives when there are both constraints and robustness requirements. The toolbox helps us toward this objective.

Consider the following system:

$$\begin{bmatrix} y_1 \\ y_2 \end{bmatrix} = \begin{bmatrix} G_{11} & G_{21} \\ 0 & 1 \end{bmatrix} \begin{bmatrix} u_1 \\ u_2 \end{bmatrix} + \begin{bmatrix} G_d \\ 0 \end{bmatrix} W$$

- 2. More detail on the problem formulation can be found in the paper by Williams et al., "Idle Speed Control Design Using an H_∞ Approach," *Proceedings of American Control Conference*, 1989, pages 1950–1956.
- 3. D. Hrovat and B. Bodenheimer, "Robust Automotive Idle Speed Control Design Based on μ-Synthesis," *Proceedings of American Control Conference*, 1993, pages 1778–1783.

where y_1 is engine rpm, y_2 and u_2 are spark advance, u_1 is bypass valve, w is torque load (unmeasured disturbance), and G_{11} , G_{21} and Gd are the corresponding transfer functions. After scaling, the constraints on spark advance become ± 0.7 , i.e., $|u_2| \le 0.7$.

Plant #1 corresponds to operation in drive at 800 rpm and a load of 30 Nm and the transfer functions are given by

$$G_{11} = \frac{9.62 e^{-0.16s}}{s^2 + 2.4s + 5.05}$$

$$G_{21} = \frac{15.9(s+3)e^{-0.04s}}{s^2 + 2.4s + 5.05}$$

$$G_d = \frac{-19.1(s+3)}{s^2 + 2.4s + 5.05}$$

Plant #2 corresponds to operation at 800 rpm in neutral with zero load and the transfer functions are given by

$$G_{11} = \frac{20.5 e^{-0.16s}}{s^2 + 2.2 s + 12.8}$$

$$G_{21} = \frac{47.6(s + 3.5) e^{-0.04s}}{s^2 + 2.2 s + 12.8}$$

$$G_d = \frac{-19.1(s + 3.5)}{s^2 + 2.2 s + 12.8}$$

The goal is to design a model predictive controller such that the closed-loop performance is good for plants #1 and #2 when subjected to an unmeasured torque load disturbance.

Simulations

Since the toolbox handles only discrete-time systems, the models are discretized using a sampling time of 0.1. We approximate each of the discrete transfer functions with 40 step response coefficients. The function cmpc is used to generate the controller and to simulate the closed-loop response because it determines $\it optimal$ changes of the manipulated variables subject to constraints. For comparison (Simulation # 4), we also use the functions mpccon for controller design and mpcsi m for simulating closed-loop responses. On-line computations are simpler, but the resulting controller is linear and the constraints are not handled in an optimal fashion. The following additional functions from the toolbox are also used: tfd2step and pl otal l. The MATLAB code for the following simulations can be found in the file i dl ectr. m in the directory mpcdemos.

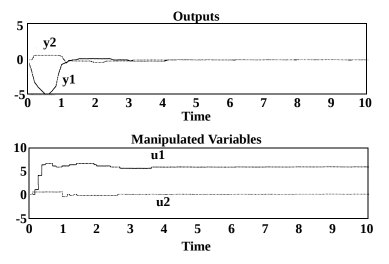


Figure 2-1 Responses to a Unit Torque Disturbance for Plant #1 (no model/plant mismatch)

Simulation #1. No model/plant mismatch. The following parameters are used:

$$M = 10$$
, $P = inf$, $ywt = [5 1]$, $uwt = [0.5 0.5]$, $tfilter = []$

The larger weight on the first output (engine rpm) is to emphasize that controlling engine rpm is more important than controlling spark advance. Figure 2-1 and Figure 2-2 show the closed-loop response for a unit step torque

load change. No model/plant mismatch is introduced, i.e., we use Plant #1 and Plant #2 as the nominal model for simulating the closed loop response for Plant #1 and Plant #2, respectively.

As we can see, both controllers perform well for their respective plants. Because of the infinite output horizon, i.e., $P = i \, nf$, nominal stability is guaranteed for both systems. In some sense, the performance observed in Figure 2-1 and Figure 2-2 is the best which can be expected, when the spark advance constraint is invoked and there is no model/plant mismatch. Obviously, if we want to control Plant #1 and Plant #2 with the same controller the nominal performance for each plant will deteriorate.

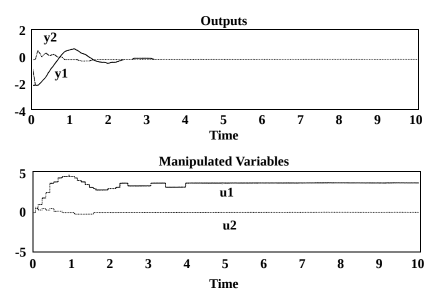


Figure 2-2 Responses to a Unit Torque Disturbance for Plant #2 (no model/plant mismatch)

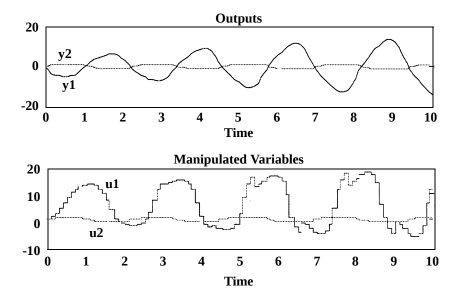


Figure 2-3 Responses to a Unit Torque Disturbance for Plant #1 (nominal model = Plant #2)

Simulation #2. Model/plant mismatch. All parameters are kept the same as in Simulation #1. Shown in Figure 2-3 is the response to a unit torque disturbance for Plant #1 using Plant #2 as the nominal model. Figure 2-4 depicts the response to a unit torque disturbance for Plant #2 using Plant #1 as the nominal model. As one can see, both systems are unstable. Therefore, the controllers must be detuned to improve robustness if one wants to control both plants with the same controller.

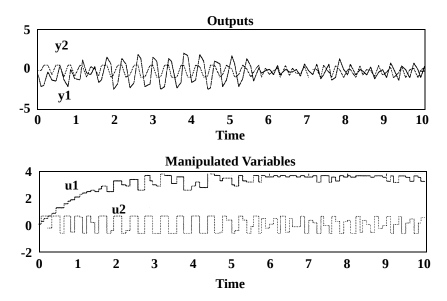


Figure 2-4 Responses to a Unit Torque Disturbance for Plant #2 (nominal model = Plant #1)

Simulation #3. The input weight is increased to [10 20] to improve robustness. All other parameters are kept the same as in Simulation #1. Plant #1 is used as the nominal model. The simulation results depicted in Figure 2-5 and Figure 2-6 seem to indicate that with an input weight of [10 20] the controller stabilizes both plants. However, we must point out that the design procedure guarantees global asymptotic stability only for the nominal system, i.e., Plant # 1. Because of the input constraints, the system is nonlinear. The observed stability for Plant # 2 in Figure 2-6 should not be mistaken as an indication of global asymptotic stability.

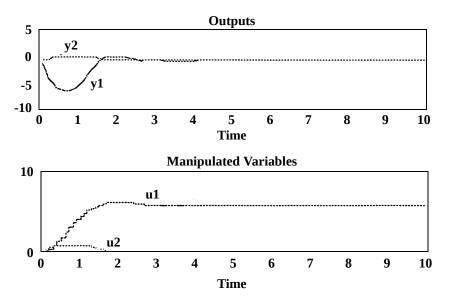


Figure 2-5 Responses to a Unit Torque Disturbance for Plant #1 (nominal model = Plant #1)

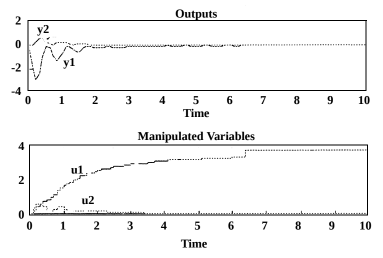


Figure 2-6 Responses to a Unit Torque Disturbance for Plant #2 (nominal model = Plant #1)

As expected, the nominal performance for both Plant #1 and Plant #2 has deteriorated when compared to the simulations shown in Figure 2-1 and Figure 2-2. A similar effect would be observed if we had detuned the controller which uses Plant #2 as the nominal model.

Simulation #4. The parameter values are the same as in Simulation #3. Instead of using cmpc, we use mpccon and mpcsi m for simulating the closed loop responses. Figure 2-2 compares the responses for Plant #1 using mpccon and mpcsi m, and cmpc. As we can see, for this example and these tuning parameters, the improvement obtained through the on-line optimization in cmpc is small. However, the difference could be large, especially for ill-conditioned systems and other tuning parameters. For example, by reducing the output horizon to P=80 while keeping the other parameters the same, the responses for Plant # 1 found with mpccon and mpcsi m are significantly slower than those obtained with cmpc (Figure 2-8).

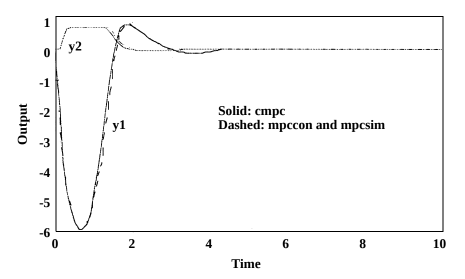


Figure 2-7 Comparison of Responses From ${\rm cmpc},$ and ${\rm mpccon}$ and ${\rm mpcsi}\,{\rm m}$ for Plant #1 P = i nf

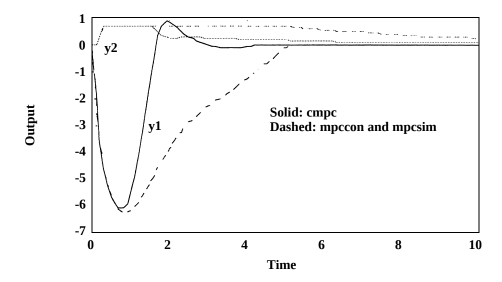


Figure 2-8 Comparison of Responses From ${\it cmpc}$, and ${\it mpccon}$ and ${\it mpcsim}$ for Plant #1 (P = 80)

Application: Control of a Fluid Catalytic Cracking Unit

Process Description

Fluid Catalytic Cracking Units (FCCUs) are widely used in the petroleum refining industry to convert high boiling oil cuts (of low economic value) to lighter more valuable hydrocarbons including gasoline. Cracking refers to the catalyst enhanced thermal breakdown of high molecular weight hydrocarbons into lower molecular weight materials. A schematic of the FCCU studied⁴ is given in Figure 2-9. Fresh feed is contacted with hot catalyst at the base of the riser and travels rapidly up the riser where the cracking reactions occur. The desirable products of reaction are gaseous (lighter) hydrocarbons which are passed to a fractionator and subsequently to separation units for recovery and purification. The undesirable byproduct of cracking is coke which is deposited on the catalyst particles, reducing their activity. Catalyst coated with coke is transported to the regenerator section where the coke is burned off thereby restoring catalytic activity and raising catalyst temperature. The regenerated catalyst is then transported to the riser base where it is contacted with more fresh feed. Regenerated catalyst at the elevated temperature provides the heat required to vaporize the fresh feed as well as the energy required for the endothermic cracking reaction.

^{4.} A detailed problem description and the model used for this study can be found in the paper by McFarlane et al., "Dynamic Simulator for a Model IV Fluid Catalytic Cracking Unit," Comp. & Chem. Eng., 17(3), 1993, pages 275–300

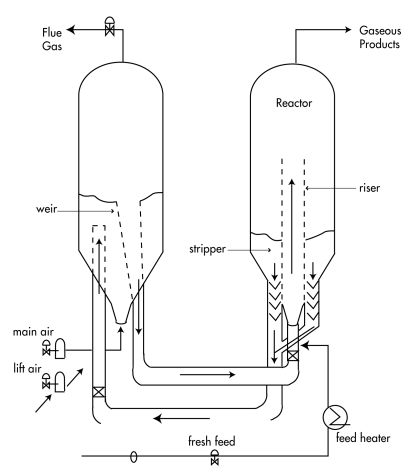


Figure 2-9 Fluid Catalytic Cracking Unit Schematic

Product composition, and therefore the economic viability of the process, is determined by the cracking temperature. The bulk of the combustion air in the regenerator section is provided by the main air compressor which is operated at full capacity. Additional combustion air is provided by the lift air compressor, the throughput of which is adjustable by altering compressor speed. By maintaining excess flue gas oxygen concentration, it is possible to ensure essentially complete coke removal from the catalyst.

Control Problem Formulation

The open loop system is modeled as follows:

$$y = Gu + G_d d$$

where

$$u = \left[V_{fg}V_{lift}\right]^{T} \quad y = \left[C_{O_{2sa}}T_{I}\Delta F_{la}\right]^{T} \quad d = \left[d_{1}d_{2}d_{3}\right]^{T}$$

G is the plant model and G_d is the disturbance model. The variables are:

- Controlled variables
 - Cracking temperature (T_r)
 - Flue gas oxygen concentration (C_{O_2} , σ
- Associated variable
 - Lift Air Compressor Surge Indicator (ΔF_{la})
- · Manipulated variables
 - Lift air compressor speed (V_{lift})
 - Flue gas valve opening (V_{fg})
- Modeled disturbances
 - Variations in ambient temperature affect compressor throughput (d_1)
 - Fluctuations in heavy oil feed composition to the FCCU (d_2)
 - Pressure upset in down stream units propagating back to the FCCU (d_3)

In addition to the controlled variables there are many process variables that need not be maintained at specific setpoints but which need to be within bounds for safety or economic reasons. These variables are called *associated variables*. For example, compressors must not surge during process upsets i.e., the suction flow rate must be greater than some minimum flow rate (surge flow rate).

The control objective is to maintain the controlled variables (cracking temperature and flue gas oxygen concentration) at pre-determined setpoints in the presence of typical process disturbances while maintaining safe plant operation.

Simulations

State-space realizations of the plant and disturbance models are available in the MATLAB file fcc_{dat} mat in the directory mpcdemos. A MATLAB script detailing the simulations is also included ($fcc_{demo.\ m}$). The following table gives the parameters used for controller design and examination of the closed loop response:

Table 2-1 FCCU Simulation Parameters

Simulation Time(s)	tend = 2500
# Step Response Coefficients	60
Process Sampling Time	delt2 = 100
Output Weights	ywt = [3 3 0]
Input Weights	uwt = [0 2]
Prediction Horizon (steps)	P = 12
# manipulated variable moves	M = 3
input constraints	$u_i \in [-1,1], i = 1,2$
output constraints	$y_i \in [-1,1]$, $i = 1,2$ $y_3 \check{\mathbf{S}} - 1$ (hard constraint)

Step Response Model

Figure 2-10A shows the plant open loop step response to a unit step in V_{fg} . Although the plant is stable the settling time is large (1 day). The time scale of interest for control purposes is on the order of one hour — which corresponds to the initial plant response, Figure 2-10B. For time scales of one hour, the process can be approximated by an integrating system. In deriving the step response model, the plant is therefore assumed to be an integrating process.

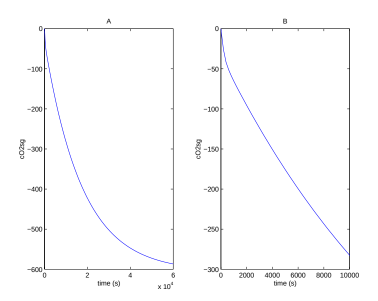


Figure 2-10 Open Loop Step Response to $u = [1 \ 0]$

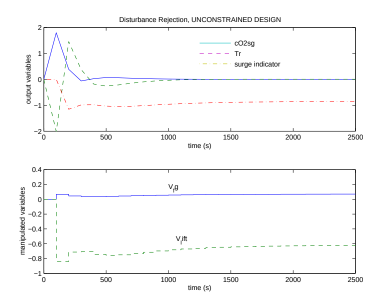


Figure 2-11 Unconstrained Closed Loop Response to d = [0 - 0.5 - 0.75]

Associated Variables

As mentioned previously, associated variables need not be at any setpoint as long as they are within acceptable bounds. Deviations of associated variables from the nominal value do not appear in overall objective function to be minimized and the output weight corresponding to the associated variable is set to zero in Table 2-1.

Unconstrained Control Law

Figure 2-11 shows the closed loop response to a disturbance d = [0 - 0.5 - 0.75] at t = 0. The controller gain matrix is derived using mpccon and the closed loop response is examined using mpcsi m. Note the following:

- At the first time step (t=100s) the controlled variables are outside their allowed limits. The onset of the disturbance at t=0 is unknown to the controller at t=0 since there is no disturbance feedforward loop. Thus from t=0 to t=100s there is no control action and the process response is the open loop response with no control action. Only after t=100s is corrective action implemented.
- At t = 200s (2^{nd} time step) riser temperature is outside the allowed limits.
- The lift air compressor surges during the interval $200s \le t \le 800s$ which is unacceptable. Compressor surging will result in undesirable vibrations in the compressor leading to rapid wear and tear.

Constrained Control Law

It is clear that the unconstrained control law generated using mpcsi m is physically unacceptable since hard output constraints are violated. Figure 2-12 shows the closed loop response of the nominal plant to the same disturbance taking process constraints explicitly into account. The closed loop response is determined using the command cmpc.

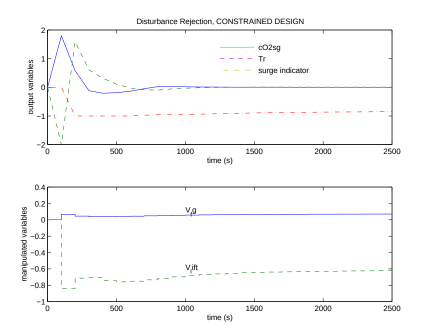


Figure 2-12 Constrained Closed Loop Response to d = [0.0.5.0.75]

The output limit vector is:

$$ylim = \begin{bmatrix} -1 & -1 & -1 & 1 & 1 & Inf \\ -1 & -1 & -1 & 1 & 1 & Inf \\ -Inf & -Inf & -Inf & Inf & Inf & Inf \end{bmatrix}$$

Note the following:

- At the first time step (t=100s) the controlled variables are outside their allowed limits. In fact the outputs are identical to the outputs for the unconstrained case at t=100s. This should be expected as there is no control action from t=0 to t=100s for both constrained and unconstrained designs.
- At t = 200s (2^{nd} time step) riser temperature (y_2) is still outside the allowed limits. This is because the constrained QP solved at t = 100s assumes that disturbances are constant for t > 100s which is not the case for this process. Thus while the manipulated variable move made at t = 100s ensures that the predicted $y_2 = 1$ at t = 200s, the actual output at t = 200s exceeds one.
- The lift air compressor does not surge during the disturbance transient, Figure 2-13.

The constrained control law therefore ensures safe operation while rejecting process disturbances. If no constraints are violated, mpccon and mpcsi m, and cmpc will give identical closed loop responses. Note that the disturbance $d = \begin{bmatrix} 0 & -0.5 & -0.75 \end{bmatrix}$ was specifically chosen to illustrate the possibility of constraint violations during disturbance transient periods.

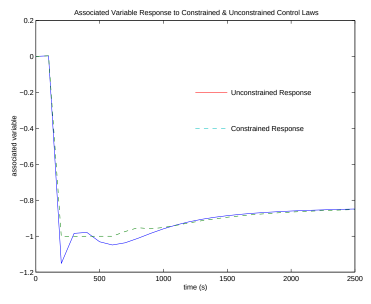
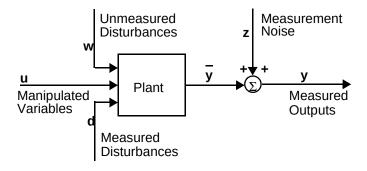


Figure 2-13 Comparison of Constrained and Unconstrained Response of ΔF_{la} to d = [0 - 0.5 - 0.75]

MPC Based on State-Space Models

State-Space Models



Consider the process shown in the above block diagram. The general discrete-time linear time invariant (LTI) state-space representation used in the MPC Toolbox is:

$$x(k+1) = \Phi x(k) + \Gamma_u u(k) + \Gamma_d d(k) + \Gamma_w w(k)$$

$$y(k) = \overline{y}(k) + z(k)$$

$$= Cx(k) + D_u u(k) + D_d d(k) + D_w w(k) + z(k)$$

where x is a vector of n state variables, u represents the n_u manipulated variables, d represents n_d measured but freely-varying inputs (i.e., measured disturbances), w represents n_w unmeasured disturbances, y is a vector of n_y plant outputs, z is measurement noise, and Φ , Γ_u , etc., are constant matrices of appropriate size. The variable $\bar{y}(k)$ represents the plant output before the addition of measurement noise. Define:

$$\Gamma = \left[\Gamma_u \, \Gamma_d \, \Gamma_w \right]$$
$$D = \left[D_u \, D_d \, D_w \right]$$

In many applications, all outputs are measured. In some cases, however, one has n_{ym} measured and n_{yu} unmeasured outputs in y, where $n_{ym} + n_{yu} = n_y$. If so, the MPC Toolbox assumes that the y vector and the y0 matrices are arranged such that the measured outputs come first, followed by the unmeasured outputs.

Mod Format

The MPC Toolbox works with state-space models in a special format, called the **mod** format. The **mod** format is a single matrix that contains the state-space Φ , Γ , C, and D matrices plus some additional information (see mod format in the "Command Reference" chapter for details). The MPC Toolbox includes a number of commands that make it easy to generate models in the **mod** format. The following sections illustrate the use of these commands.

SISO Continuous-Time Transfer Function to Mod Format

The MPC Toolbox uses a format called the tf format. Let the continuous-time transfer function be

$$G(s) = \frac{b_0 s^n + b_1 s^{n-1} + \dots + b_n}{a_0 s^n + a_1 s^{n-1} + \dots + a_n} e^{-T_d s}$$

where T_d is the time delay. The tf format is a matrix consisting of three rows:

row 1: The *n* coefficients of the numerator polynomial, b_0 to b_n .

row 2: The *n* coefficients of the denominator polynomial, a_0 to a_n .

row 3: column 1: The sampling period. *This must be zero for a continuous system.* (It must be positive for discrete transfer functions — see next section).

column 2: The time delay in time units. It must satisfy $Td \ge 0$.

The tf matrix will always have at least two columns, since that is the minimum width of the third row.

You can either define a model in the tf format directly or use the command pol y2tfd. The general form of this command is

For example, consider a SISO system modeled by the transfer function

$$G(s) = \frac{-13.6s + 1}{54.3s^2 + 113.5s + 1}e^{-5.3s}$$

To create the tf format directly you could use the command

$$G = [0 - 13.6 1; 54.3 113.5 1; 0 5.3 0];$$

which defines a matrix consisting of three rows and three columns. Note that all rows must have the same number of columns so you must be careful to insert zeros where appropriate. The pol y2tfd command is more convenient since it does that for you automatically:

$$G = poly2tfd([-13.6 1], [54.3 113.5 1], 0, 5.3);$$

Either command would define a variable G in your workspace, containing the matrix

To convert this to the \boldsymbol{mod} format, use the command tfd2mod, which has the form

$$model = tfd2mod(delt, ny, g1, g2, g3, ..., gN)$$

where:

del t The sampling period. tfd2mod will convert your continuous time transfer function(s) g1, ..., gN to

discrete-time using this sampling period.

ny is the number of output variables in the plant you are

modeling.

g1, g2, ..., gN A sequence of N transfer functions in the tf format, where $N \ge 1$. tfd2mod assumes that these are the individual elements of a transfer-function matrix:

$$\begin{bmatrix} g_{1,\,1} & g_{1,\,2} & \cdots & g_{1,\,n_u} \\ g_{2,\,1} & g_{2,\,2} & \cdots & g_{2,\,n_u} \\ \vdots & \vdots & \ddots & \vdots \\ g_{n_y,\,1} & g_{n_y,\,2} & \cdots & g_{n_y,\,n_u} \end{bmatrix}$$

Thus N must be an integer multiple (n_u) of the number of outputs, n_y . You supply the transfer functions in *column-wise* order. In other words, you first give the n_y transfer functions for input 1 $(g_{1,\,1}$ to $g_{n_y,\,1})$, then the n_y transfer functions for input 2 $(g_{1,\,2}$ to $g_{n_y,\,2})$, etc.

Suppose you want to convert the SISO model defined above to the **mod** format with a sampling period of 2.1 time units. The appropriate command would be

$$mod = tfd2mod(2.1, 1, G);$$

This would define a variable called mod in your workspace that would contain the discrete-time state-space description of your system.

SISO Discrete-Time Transfer Function to Mod Format

Suppose you have a transfer function in discrete-time format (in terms of the forward-shift operator, *z*):

$$G(q) = \frac{b_0 + b_1 z^{-1} + \dots + b_n z^{-n}}{a_0 + a_1 q^{-1} + \dots + a_n z^{-n}} z^{-d}$$

where d is an integer (≥ 0) and represents the sampling periods of pure delay. The corresponding tf format is the same as for the continuous-time case except for the definition of the third row:

- column 1 is the sampling period for which G(z) was created. It must be positive (in contrast to the continuous-time case described above).
- column 2 is the *periods* of pure delay, d, which must be an $integer \ge 0$. Contrast this to the continuous case, where the delay is given in time units.

As in the previous section, you can use poly2tfd followed by tfd2mod to get such a transfer function in **mod** format. For example, the discrete-time representation of the SISO system considered in the previous section is

$$G(z) = \frac{-0.1048 + 0.1215z^{-1} + 0.0033z^{-2}}{1 - 0.9882z^{-1} + 0.0082z^{-2}}z^{-3}$$

If you had this to begin with, you could convert it to the **mod** format as follows:

$$G = poly2tfd([-0.1048 \ 0.1215 \ 0.0033], [1 -0.9882 \ 0.0082], 2.1, 3);$$

 $mod = tfd2mod(2.1, 1, G);$

Note that both the poly2tfd and tfd2mod commands specify the same sampling period (del t=2. 1). This would be the usual case, but you have the option of converting a discrete-time model in the tf format to a different sampling period in the mod format.

MIMO Transfer Function Description to Mod Format

Suppose you have a transfer-function matrix description of your system in the form

$$\begin{bmatrix} g_{1,\,1} & g_{1,\,2} & \cdots & g_{1,\,n_u} \\ g_{2,\,1} & g_{2,\,2} & \cdots & g_{2,\,n_u} \\ \vdots & \vdots & \ddots & \vdots \\ g_{n_y,\,1} & g_{n_y,\,2} & \cdots & g_{n_y,\,n_u} \end{bmatrix}$$

where $g_{i,j}$ is the transfer function of the i^{th} output with respect to the j^{th} input. If all n_y outputs are measured and all n_u inputs are manipulated variables, the default mode of tfd2mod will give you the correct **mod** format. For example, consider the 2-output, 3-input system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} & \frac{3.8e^{-8s}}{14.9s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} & \frac{4.9e^{-3s}}{13.2s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \\ u_3(s) \end{bmatrix}$$

The following sequence of commands would convert this to the equivalent **mod** format with a sampling period of T = 4:

```
g11 = poly2tfd(12.8, [16.7 1], 0, 1);

g21 = poly2tfd(6.6, [10.9 1], 0, 7);

g12 = poly2tfd(-18.9, [21.0 1], 0, 3);

g22 = poly2tfd(-19.4, [14.4 1], 0, 3);

g13 = poly2tfd(3.8, [14.9 1], 0, 8);

g23 = poly2tfd(4.9, [13.2 1], 0, 3);

pmod = tfd2mod(4, 2, g11, g21, g12, g22, g13, g23);
```

Suppose, however, that the third input were actually an unmeasured disturbance, i.e., the system were

$$\begin{bmatrix} y_1 s \\ y_2 s \end{bmatrix} = \begin{bmatrix} \frac{12.8 e^{-s}}{16.7 s + 1} & \frac{-18.9 e^{-3s}}{21.0 s + 1} \\ \frac{6.6 e^{-7s}}{10.9 s + 1} & \frac{-19.4 e^{-3s}}{14.4 s + 1} \end{bmatrix} \begin{bmatrix} u_1 s \\ u_2 s \end{bmatrix} + \begin{bmatrix} \frac{3.8 e^{-8s}}{14.9 s + 1} \\ \frac{4.9 e^{-3s}}{13.2 s + 1} \end{bmatrix} w(s)$$

In this case you would need to override the default mode of tfd2mod by specifying the number of inputs in each of the three categories described at the beginning of this section, i.e., manipulated variables, measured disturbances, and unmeasured disturbances. This and other information about the system is contained in the first 7 columns of row 1 of the **mod** format, as follows:

- column 1 T, the sampling period.
 - 2 *n*, the number of states.
 - 3 n_{u} , the number of manipulated variable inputs.
 - 4 n_d , the number of measured disturbances.
 - 5 n_{W} , the number of unmeasured disturbances.
 - 6 n_{ym} , the number of measured outputs.
 - 7 n_{yu} , the number of unmeasured outputs.

For example, if you had defined pmod using the default mode of tfd2mod as shown above, the contents of row 1, columns 1 to 7 of pmod would be:

You could override this to set $n_u = 2$ and $n_w = 1$ as follows:

$$pmod(1, 3) = 2;$$

 $pmod(1, 5) = 1;$

Note that in the original transfer function matrix description, the first n_u columns must be for the manipulated variables, the next n_d for the measured disturbances (if any), and the last n_w for the unmeasured disturbances (if any). Similarly, the first n_{ym} outputs must be measured and the last $n_{yu} \ (\ge 0)$ unmeasured.

Continuous or Discrete State-Space to Mod Format

If you have a continuous-time state-space model, you may convert it to **mod** format by first using the function c2dmp (continuous to discrete-time state-space), followed by ss2mod(discrete-time state-space to **mod** format). Of course, if you are starting with a discrete-time state-space model you can skip the first step.

For example, suppose a, b, c, and d are matrices describing a continuous-time system. To convert to the **mod** format using a sampling period of T = 1.5, you could use the following commands:

```
[phi, gam] = c2dmp(a, b, 1.5);
mod = ss2mod(phi, gam, c, d, 1.5);
```

If your system is complicated, i.e., it contains disturbance inputs and/or unmeasured outputs, you will need to override the default mode of ss2mod. See the "Command Reference" section for more details.

Identification Toolbox ("Theta") Format to Mod Format

The System Identification Toolbox identifies discrete-time transfer-function models from input/output data. The result is a model in a special form called the *theta* format (see the *System Identification Toolbox User's Guide* for details). In general, each *theta* model describes the response of a single output to one or more inputs (MISO model).

The MPC Toolbox function, th2mod, converts one or more such models to the **mod** format. Suppose, for example, that

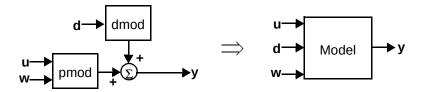
- th1 is the *theta* model describing the response of output y_1 to inputs u_1 and u_2 .
- th2 is the *theta* model describing the response of output y_2 to inputs u_1 and u_2 .

Then the following command would provide the equivalent **mod** format with $n_v = 2$ and $n_u = 2$:

```
mod = th2mod(th1, th2);
```

Combination of Models in Mod Format

The functions addmod, addmd, addumd, appmod, paramod, and sermod allow you to combine simple models in the **mod** format to generate more complex plant structures. For example, addmd includes the effect of one or more measured disturbances in an existing model, as shown in the following schematic:



pmod gives the effect of one or more manipulated variables, u(k), and optional unmeasured disturbance(s), w(k), on the output(s), y(k). dmod gives the effect of the measured disturbance(s), d(k), on the same outputs. Once you have defined pmod and dmod (e.g., starting from transfer functions as illustrated above), you can use the command addmd to generate the composite, model:

```
model = addmd(pmod, dmod);
```

Please see Chapter 4, "Command Reference" for more details on the various model-building commands.

Converting Mod Format to Other Model Formats

The function mod2ss converts a model in the **mod** format to the standard discrete-time state-space format:

```
[phi, gam, c, d, minfo] = mod2ss(mod);
```

Here, phi, gam, c, and d are the coefficient matrices of

$$x(k+1) = \Phi x(k) + \Gamma u(k)$$

$$y(k) = Cx(k) + Du(k)$$

The vector mi nfo contains the first 7 columns of the first row in mod. The section "MIMO Transfer Function Description to Mod Format" gives the significance of this information.

Once you have phi, gam, c, and d, you can use d2cmp, ss2tf2, and other functions to convert from discrete state-space to other model forms.

The function mod2step uses a model in the **mod** format to generate a stepresponse model in the **step** format as required by the functions mpccon, mpcsi m, etc., discussed in Chapter 2, "MPC Based on Step Response Models". See the Chapter 4, "Command Reference" for details on the use of mod2step.

Unconstrained MPC Using State-Space Models

Once you have described your system by generating state-space models in the **mod** format you can use the commands:

smpccon to calculate the unconstrained controller gain matrix.

smpcest to design a state estimator (optional).

smpcsim to simulate the response of the closed-loop system to one or more specified inputs.

plotal 1 (or ploteach) to plot the response(s).

In addition, you can analyze certain properties of the closed-loop system using the commands:

smpccl to generate a model of the closed-loop system (plant plus controller).

smpcgain to calculate the closed-loop gain matrix.

smpcpol e to calculate the closed-loop poles.

mod2frsp (and plotfrsp) to calculate and plot the closed-loop frequency response.

svdfrsp to calculate the singular values of the frequency response.

Note: smpcgai n, smpcpol e and mod2frsp also work with open-loop models in the **mod** format.

Example: (see mpctutss. m)

The following example (mpctutss. m) illustrates the basic procedures. The example process has 2 measured outputs, 2 manipulated variables, and an unmeasured disturbance:

$$\begin{bmatrix} y_1 s \\ y_2 s \end{bmatrix} = \begin{bmatrix} \frac{12.8 e^{-s}}{16.7 s + 1} & \frac{-18.9 e^{-3s}}{21.0 s + 1} \\ \frac{6.6 e^{-7s}}{10.9 s + 1} & \frac{-19.4 e^{-3s}}{14.4 s + 1} \end{bmatrix} \begin{bmatrix} u_1 s \\ u_2 s \end{bmatrix} + \begin{bmatrix} \frac{3.8 e^{-8s}}{14.9 s + 1} \\ \frac{4.9 e^{-3s}}{13.2 s + 1} \end{bmatrix} w(s)$$

We first define the model in the **mod** format. The following commands use a sampling period of T = 2 time units (chosen arbitrarily):

```
del t = 2;
ny = 2;
g11 = poly2tfd(12.8, [16.7 1], 0, 1);
g21 = poly2tfd(6.6, [10.9 1], 0, 7);
g12 = poly2tfd(-18.9, [21.0 1], 0, 3);
g22 = poly2tfd(-19.4, [14.4 1], 0, 3);
umod = tfd2mod(del t, ny, g11, g21, g12, g22);
% Defines the effect of u inputs
g13 = poly2tfd(3.8, [14.9 1], 0, 8);
g23 = poly2tfd(4.9, [13.2 1], 0, 3);
dmod = tfd2mod(del t, ny, g13, g23);
% Defines the effect of w input
pmod = addumd(umod, dmod); % Combines the two models.
```

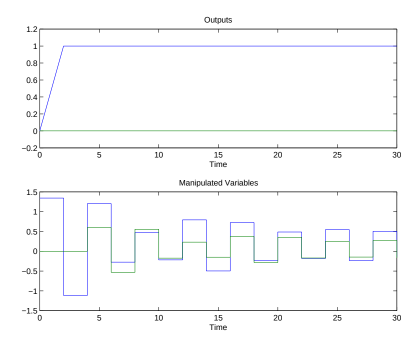
We now design an unconstrained MPC controller. The design parameters are essentially the same as for the functions based on step response models (see Chapter 2). In this case, start by choosing design parameters such that we get the *perfect controller*:

```
i mod = pmod; % assume perfect modeling
ywt = [ ]; % default (unity) weights on both outputs
uwt = [ ]; % default (zero) weights on both inputs
P = 5; % prediction horizon
M = P; % control horizon
Ks = smpccon(i mod, ywt, uwt, M, P);
```

We check the design by running a simulation for a step increase in the setpoint of output y_1 :

```
tend=30; % time period for simulation. r = [1 \ 0]; % setpoints for the two outputs. [y, u] = smpcsim(pmod, imod, Ks, tend, r); plotall(y, u, delt)
```

Note that there is no model error since we used the same model to represent the *plant* (pmod) as that used to design the controller (i mod). The results are:



Note that we get perfect tracking of the specified setpoint change ($y_1 = 1$, $y_2 = 0$), but the manipulated variables are *ringing*. You could have anticipated this by calculating the poles of the controller:

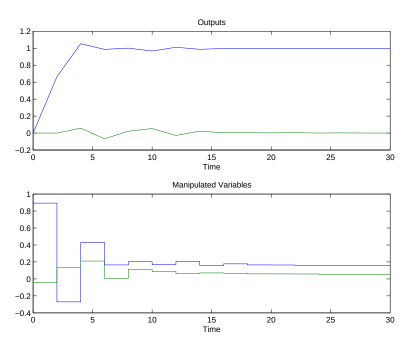
```
[cl mod, cmod] = smpccl (pmod, i mod, Ks);
smpcpol e(cmod)
```

The result shows that one of the poles is at -0.9487 and another is at -0.9223. In general, such negative-real poles cause ringing.

One way to minimize ringing is to make the prediction horizon significantly larger than the control horizon:

```
P = 10;
M = 3;
Ks = smpccon(i mod, ywt, uwt, M, P);
[y, u] = smpcsi m(pmod, i mod, Ks, tend, r);
plotall(y, u, delt)
```

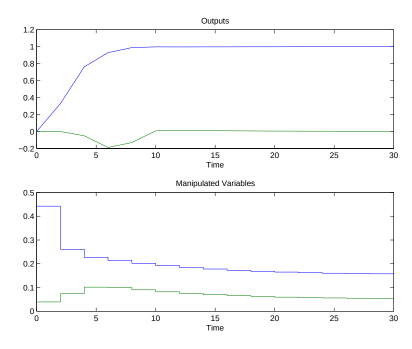
This results in the following improved responses:



Another (often more effective) way is to use *blocking*. In the case of blocking, each element of the vector M indicates the number of steps over which $\Delta u = 0$ during the optimization. For example, $M = [2\ 3]$ indicates that u(k+1) = u(k) or $\Delta u(k+1) = 0$ and u(k+4) = u(k+3) = u(k+2) (or $\Delta u(k+3) = \Delta u(k+4) = 0$):

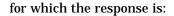
```
\label{eq:main_main} \begin{split} M &= [2\ 3\ 4]; \ \% \ Defines \ 3 \ blocks \ of \ control \ moves \\ Ks &= smpccon(i \ mod, ywt, uwt, M, P); \\ [y, u] &= smpcsi \ m(pmod, i \ mod, Ks, tend, r); \\ plotall(y, u, delt) \\ pause \end{split}
```

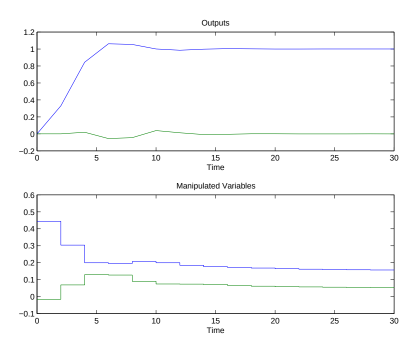
This completely eliminates the ringing, as shown in the following responses, at the expense of a more sluggish servo response and a larger disturbance in y_2 .



A third approach is to increase the weights on the manipulated variables:

```
uwt = [1 1]; % increase input weighting
P = 5; % original prediction horizon
M = P; % original control horizon
Ks = smpccon(i mod, ywt, uwt, M, P);
[y, u] = smpcsim(pmod, i mod, Ks, tend, r);
plotall(y, u, delt)
```

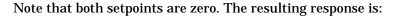


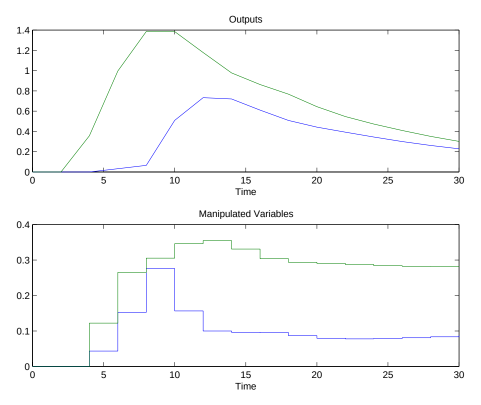


In general, you must choose the horizons and weights by trial-and-error, using simulations to judge their effectiveness.

The servo-response of the last controller looks good. Let's see how it responds to a unit-step in the unmeasured disturbance, w(k):

```
ulim = []; % default (no) constraints on u variables.
Kest = []; % default (DMC) state estimator.
r = [0 0]; % Both output setpoints at zero.
z = []; % default (zero) measurement noise.
v = []; % default (zero) measured disturbances.
w = [1]; % unit-step in unmeasured disturbance.
[y, u] = smpcsim(pmod, i mod, Ks, tend, r, ulim, Kest, z, v, w);
plotall(y, u, delt)
pause
```

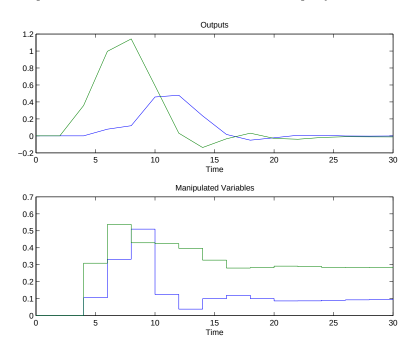




The regulatory response has a rather long transient. Let's see if we can improve it by using a state estimator other than the default (DMC) estimator:

```
[Kest, newmod] = smpcest(i mod, [15 15], [3 3]);
Ks = smpccon(newmod, ywt, uwt, M, P);
[y, u] = smpcsim(pmod, newmod, Ks, tend, r, ulim, Kest, z, v, w);
plotall(y, u, delt)
```

See the detailed description of the smpcest function for a discussion of the estimator design parameters. The results show that the controller now compensates for the disturbance much more rapidly:



State-Space MPC with Constraints

The function scmpc handles problems with inequality constraints on the manipulated variables and/or outputs. The recommended procedure is to first use the tools described in the section section Unconstrained MPC Using State-Space Models to find values of the prediction horizon, P, control horizon, M, input and output weights, and a state-estimation strategy that work well for the unconstrained version of your problem. Then define the constraints and solve the problem using scmpc. The following example illustrates the use of scmpc.

Example: (see mpctutss. m)

We use the same example process as in the previous section, but use a sampling period of 1 and omit the unmeasured disturbance input:

```
T = 1;

g11 = poly2tfd(12.8, [16.7 1], 0, 1);

g21 = poly2tfd(6.6, [10.9 1], 0, 7);

g12 = poly2tfd(-18.9, [21.0 1], 0, 3);

g22 = poly2tfd(-19.4, [14.4 1], 0, 3);

i mod = tfd2mod(2, T, g11, g21, g12, g22);
```

The following statements specify parameters required in both the constrained and unconstrained cases, and calculate the gain for the unconstrained controller.

```
nhor = 10; % Prediction horizon.
ywt = [ ]; % Unity weighting on output tracking errors
% (default).
uwt = [ ]; % Zero weighting on man. variable moves
% (default).
blks = [2 3 5]; % Allows 3 moves of manipulated variables.
K = [ ]; % DMC-type state estimation (default).
Ks = smpccon(i mod, ywt, uwt, blks, nhor);
```

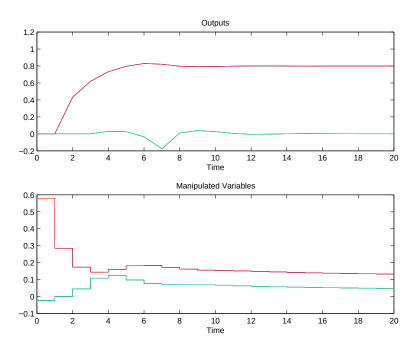
Let's first verify that the constrained and unconstrained solutions will be the same when the constraints are *loose* i.e. inactive. The following statements define upper and lower bounds on u(k) at $-\infty$ and ∞ , respectively, and bounds on $\Delta u(k)$ at 10 (both u_1 and u_2). Also, bounds on y(k) are set at the default values of $\pm \infty$.

```
ulim = [-inf -inf inf inf 10 10];
ylim = [ ]; % Default -- no limits on outputs.
```

For the simulation we will make a step change of 0.8 in the setpoint for y_1 . We will also assume a perfect model, i.e., use the same model for the plant as was used to design the controller.

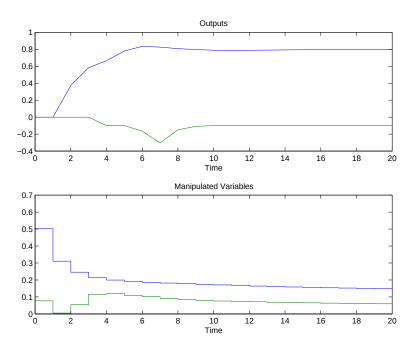
The above plotall command plots the results from smpcsim and scmpc on the same graph. Since the constraints were loose, there should be no difference. In the following plots, you can only distinguish two curves, i.e., the two simulations give the same values of y and u, as expected.

^{1.} Finite bounds on Δu are required by scmpc. Here they are chosen large enough so that they have no effect.



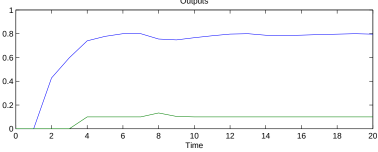
Now let's add some constraints to the problem. Suppose we want the maximum value of y_2 to be -0.1. In the previous case it goes slightly above zero (dashed line in the *Outputs* plot). The following statements define a hard upper limit of y_2 = -0.1, starting at the 4th step in the prediction horizon. This accounts for the minimum delay of 3 sampling periods before y_2 can be affected by either u_1 or u_2 , i.e., it is important to leave y_2 unconstrained for the first 3 steps in the prediction horizon. In this case, since the initial condition is y_2 = 0, it is impossible to make $y_2 \le$ -0.1 prior to t = 4. If you were to attempt to do so, you would get an error message stating that the problem is infeasible. Note also that the upper bound on y_2 supersedes the setpoint, which is still specified as zero. The controller thus maximizes the value of y_2 at steady state.

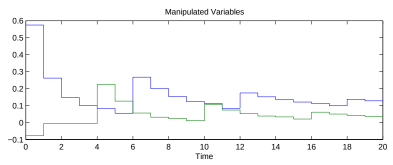




If you use a long prediction horizon with constraints, the calculations can be time-consuming. You can minimize this by *turning off* constraints that are far out in the prediction horizon. The following example defines bounds on y_1 and y_2 , then turns them off beyond the 4th point in the prediction horizon. The calculations are much faster than would be the case if only the first 4 rows of yl i m had been used (try it). Also, since there is neither model error nor unmeasured disturbances, the solution satisfies all constraints for t 4 in any case. In general, output constraints must be chosen carefully to avoid infeasibilities and maximize the speed of the calculations.

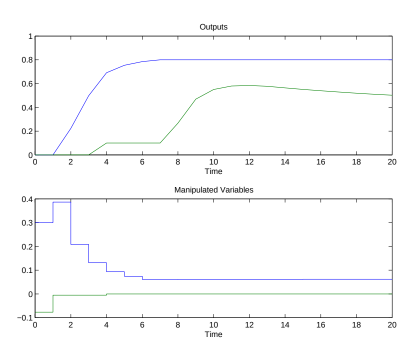
```
 [y,u] = scmpc(pl \ ant, i \ mod, ywt, uwt, bl \ ks, nhor, tend, ... \\ setpts, ul \ i \ m, yl \ i \ m, K); \\ pl \ otal \ l \ (y,u,T)
```





As a final example we impose bounds on the manipulated variables:

Again, to save computer time the constraints apply only for the first block in the prediction horizon, i.e., the constraints are turned off for periods 2 through P=10. The following plot shows that the upper bound of $u_2 \le 0$ and $\Delta u_1 \le 0.3$ are the most restrictive. The former prevents y_2 from coming back to the minimum allowed value of 0.1.



Application: Paper Machine Headbox Control

Ying et al. $(1992)^2$ studied the control of composition and liquid level in a paper machine headbox, a schematic of which is shown in Figure 3-1. The process model is given by a set of ordinary differential equations (ODEs) in bilinear form. Using their nomenclature, the states are $x^T = [H_1 \ H_2 \ N_1 \ N_2]$, where H_1 is the liquid level in the feed tank, H_2 is that in the headbox, N_1 is the consistency (percentage of pulp fibers in suspension) in the feed tank, and N_2 is that in the headbox. All states except H_1 are measured, i.e., the measured outputs are $y^T = [H_2 \ N_1 \ N_2]$. The primary control objective is to hold H_2 and N_2 (y_1 and y_3) at specified setpoints.

There are two manipulated variables: $u^T = [G_p \ G_w]$, where G_p is the flowrate of stock entering the feed tank, and G_w is that of the recycled *white water*. There is a single measured disturbance: $v = [N_p]$, the consistency of the stock entering the feed tank, and a single unmeasured disturbance: $d = [N_w]$, the consistency of the white water. All variables are normalized such that they are zero at the nominal steady state, and variations about the steady-state are of the same order of magnitude. The process is open-loop stable.

Ying, Y., M. Rao, and Y. Sun, "Bilinear Control Strategy for Paper-Making Process," Chem. Eng. Comm. 1992, 111, 13–28.

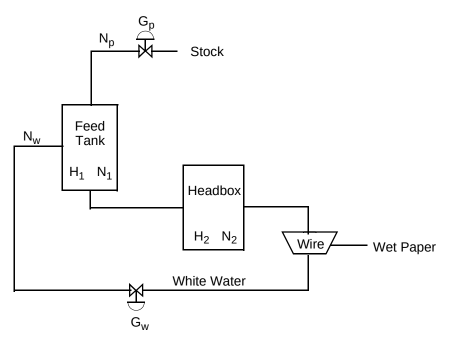


Figure 3-1 Schematic of Paper Machine Headbox Control Problem

MPC Design Based on Nominal Linear Model

The standard MPC design methods require a linear model of the plant. We therefore linearize the bilinear model at the nominal steady-state condition (x = 0; u = 0; v = 0; d = 0). Since the model is simple, one can linearize it analytically to obtain:

$$x_m = Ax_m + B_0 u + B_v v + B_d d_m$$

where x_m , y_m , and d_m are the model states, outputs, and disturbances, respectively. The desired closed-loop response time is of the order of 10 minutes, so we choose a sampling period of $T_s = 2$ minutes. The file pm_l i n. m in the directory mpcdemos contains the code for all the computations in this section of the manual. The following commands define the linear model and plot the response of the outputs to a unit step in each manipulated variable:

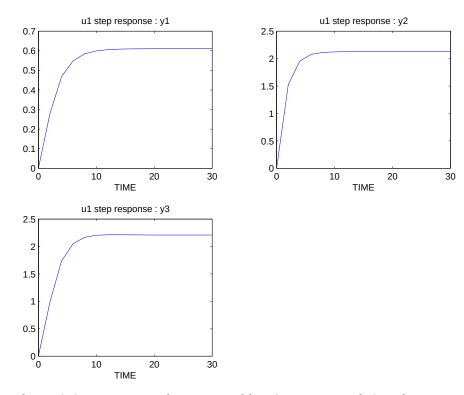


Figure 3-2 Responses of Paper Machine Outputs to Unit Step in u_1

```
% Matrices= of the linearized paper machine model
```

```
A = [-1.93 \ 0 \ 0]; \ .394 \ -.426 \ 0]; \ 0 \ 0 \ .63 \ 0; \ .82 \ -.784
   . 413 - . 426];
```

 $B = [1.274 \ 1.274 \ 0 \ 0; 0 \ 0 \ 0; 1.34 \ -.65 \ .203 \ .406; 0 \ 0 \ 0];$

 $C = [0 \ 1 \ 0 \ 0; \ 0 \ 0 \ 1 \ 0; \ 0 \ 0 \ 0 \ 1];$

D = zeros(3, 4);

% Discretize the linear model and save in mod form.

dt = 2;

```
[PHI, GAM] = c2dmp(A, B, dt);
  minfo = [dt, 4, 2, 1, 1, 3, 0];
 i mod = ss2mod(PHI, GAM, C, D, mi nfo);
 plotstep(mod2step(i mod, 30))
             u2 step response : y1
                                                               u2 step response : y2
 0.7
 0.6
                                                  -0.2
 0.5
                                                 -0.4
 0.4
                                                 -0.6
 0.3
                                                 -0.8
 0.2
                                                   -1
 0.1
                                                 -1.2
 0
0
                                                 -1.4 <sup>L</sup>
               10
                           20
                                       30
                                                                 10
                                                                                         30
                                                                             20
                    TIME
                                                                      TIME
             u2 step response : y3
 0.4
 0.2
  0
-0.2
-0.4
-0.6
-0.8
 -1 <u>-</u>
               10
                           20
                                       30
                    TIME
```

Figure 3-3 Responses of Paper Machine Outputs to Unit Step in u_2

The step responses (Figure 3-2 and Figure 3-3) show that there are large interactions in the open-loop system. Adjustments in u_1 and u_2 have strong effects on both y_1 and y_3 . Also, the $u_2 \rightarrow y_3$ step exhibits an inverse response. We begin by attempting a controller design for good response of both y_1 and y_3 :

```
% Define controller parameters
P = 10; % Prediction horizon
M = 3: % Control horizon
ywt = [1, 0, 1]; % Equal weighting of y(1) and y(3),
% no control of y(2)
uwt = 0.6*[1\ 1]; % Equal weighting of u(1) and u(2).
ulim = [-10*[1 \ 1] \ 10*[1 \ 1] \ 2*[1 \ 1]];% Constraints on u
ylim = [ ]; % No constraints on y
Kest = [ ]; % Use default estimator
% Simulation using scmpc -- no model error
pmod=i mod; % plant and internal model are identical
setpts = [1 \ 0 \ 0];
% servo response to step in y(1) setpoint
tend = 30; % duration of simulation
[y, u, ym] = scmpc(pmod, i mod, ywt, uwt, M, P, tend, ...
   setpts, ulim, ylim, Kest);
plotall(y, u, dt)
```

The prediction horizon of 10 sampling periods (20 minutes) extends well past the desired closed-loop response time. Preliminary trials suggested that longer horizons increased the computational load but provided no advantage in setpoint tracking. The use of M < P is not required in this case, but helps to reduce the computational load and inhibit ringing of the manipulated variables. Note the equal penalties on setpoint tracking errors for y_1 and y_3 (ywt variable), reflecting our desire to track both setpoints accurately. There is no penalty on y_2 , since it does not have a setpoint. The listed uwt penalties were determined by running several trials. Figure 3-4 shows smooth control of y_1 with the desired 10-minute response time, but there is a noticeable disturbance in y_3 .

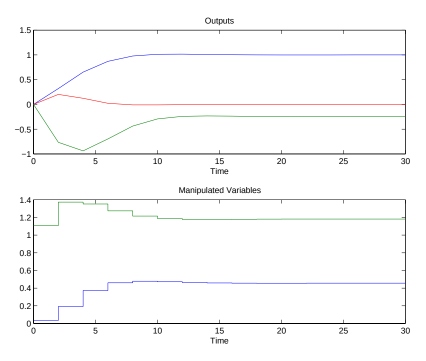


Figure 3-4 Response of closed-loop system to unit step in y_1 setpoint for equal output weighting. Output y_2 is uncontrolled, and the y_3 setpoint is zero

One could repeat the simulation for a step change in the y_3 setpoint as follows:

```
setpts = [0\ 0\ 1]; \ \% \ servo \ response \ to \ step \ in \ y(3) \ setpoint \\ [y, u, ym] = scmpc(pmod, i mod, ywt, uwt, M, P, tend, ... \\ setpts, ulim, ylim, Kest); \\ plotall(y, u, dt)
```

Normally, control of consistency (y_3) is more important than control of liquid level, y_1 . We can achieve better control of y_3 if we allow larger tracking errors in y_1 . For example, an alternative controller design uses unequal weights on the controlled outputs:

As shown in Figure 3-5, a y_1 setpoint change causes a much smaller disturbance in y_3 than before (compare with Figure 3-4). The disadvantage is that the response time of y_1 has increased from about 8 to 25 minutes. Similarly, a step change in the y_3 setpoint would cause a larger disturbance in y_1 than in the original design. Overall, however, the controller with unequal weighting gives better nominal performance and will be used as the basis for subsequent designs.

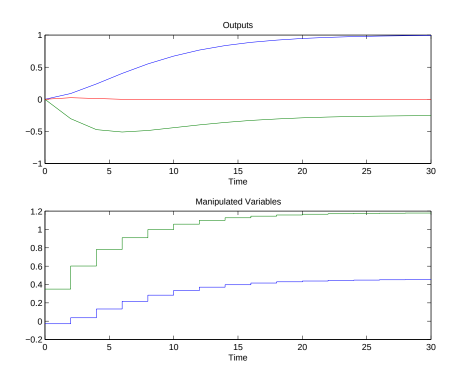
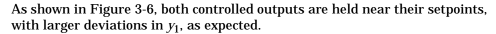


Figure 3-5 Response of closed-loop system to Unit step in y_1 setpoint for unequal output weighting. Output y_2 is uncontrolled, and the y_3 setpoint is zero.

We now evaluate the response of the above controller to a unit step in the measured disturbance, v (i.e., feedforward control). The commands required for this are:

```
setpts = [0\ 0\ 0]; \ \% \ output setpoints \ z = [\ ]; \ \% \ measurement \ noise v = 1; \ \% \ measured \ disturbance d = 0; \ \% \ unmeasured \ disturbance \ [y, u, ym] = scmpc(pmod, i mod, ywt, uwt, M, P, tend, ... setpts, ulim, ylim, Kest, z, v, d); plotall(y, u, dt)
```



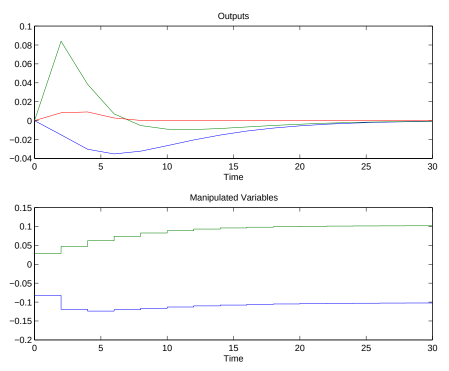


Figure 3-6 Response of closed-loop system to unit step in measured disturbance, v. unequal output weighting with y_1 and y_3 setpoints at zero.

Finally, we check the response to a step in the unmeasured disturbance. The required commands are:

```
setpts = [0 0 0]; % output setpoints
v = 0; % measured disturbance
d = 1; % unmeasured disturbance
[y, u, ym] = scmpc(pmod, i mod, ywt, uwt, M, P, tend, . . .
    setpts, ulim, ylim, Kest, z, v, d);
plotall(y, u, dt)
```

As shown in Figure 3-7, the unmeasured disturbance causes significant deviations in both controlled outputs. In fact, the higher-priority output, y_3 , exhibits the larger tracking error of the two controlled variables.

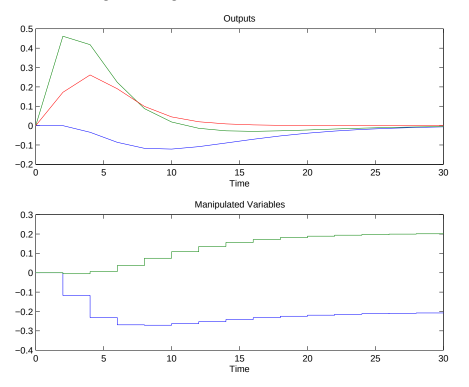
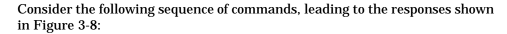


Figure 3-7 Response of closed-loop system (with default state estimator) to unit step in unmeasured disturbance, d. unequal output weighting with y_1 and y_3 setpoints at zero.

The closed-loop response to unmeasured disturbances can often be improved by a change in the state estimator. In the previous trials, we were using the default estimator, which assumes that disturbances are independent, random steps at each output. In fact, the *known* unmeasured disturbance, d, has no effect on y_1 , and its effects on y_2 and y_3 are approximately first order with time constants of 3 and 5 minutes, respectively. One way to exploit this knowledge is to specify an expected covariance for d and a measurement noise covariance for y, then use the Kalman gain for the modeled disturbance characteristics.



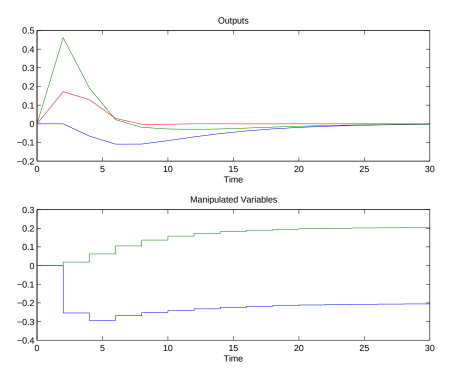


Figure 3-8 Response of closed-loop system to unit step in unmeasured disturbance, d. with Kalman Estimator, unequal output weighting with y_1 and y_3 setpoints at zero.

```
% Estimator design
Q = 30;
R = 1*eye(3);
Kest = smpcest(i mod, Q, R);
% Simulation using scmpc -- no model error
setpts = [0 0 0]; % servo response to step in y1 setpoint
d = 1; % unmeasured disturbance
[y, u, ym] = scmpc(pmod, i mod, ywt, uwt, M, P, tend, ...
    setpts, ulim, ylim, Kest, z, v, d);
plotall(y, u, dt)
```

The resulting Kest matrix is:

```
Kest =
    0    00
    0.0000 - 0.00000.0000
    0.0000    0.74460.0732
    -0.0000    0.29850.0851
    0.0000 - 0.00000.0000
    -0.0000    0.77770.2934
    0.0000    0.29340.1977
```

We have specified equal measurement noise for each output (R is diagonal with a rank equal to the number of outputs, and equal elements on the diagonal). This makes the Kalman estimator give equal weight to each output measurement. The dimension of Q must equal the number of elements in d (unity in this case). A relatively large value of Q(i,i) signifies an important disturbance mode. In practice, the elements of Q and R are tuning parameters, and one adjusts the relative magnitudes to achieve the desired balance of fast disturbance rejection (usually promoted by making Q relatively large) and robustness.

For the chosen Q and R, and the disturbance model in i mod, the elements of column 1 of Kest (shown above) are essentially zero. Thus, the measurement of y_1 provides no information regarding the effect of d on the process states. Output y_2 , on the other hand, provides large corrections to the state estimates. If it were not available, rejection of d would degrade.⁴

Figure 3-8 shows that although the revised estimator reduced the disturbance in y_3 , it is still significant (compare to Figure 3-7). A key limiting factor is the use of a 2-minute sampling period. As shown in Figure 3-8, the controller does not respond to the disturbance until it is first detected at t=2 minutes. You can verify that reducing the sampling period to 0.25 minutes (holding all other parameters constant) greatly reduces the disturbance in y_3 . Such a change would also speed up the setpoint tracking in the nominal case. It may cause robustness problems, however, so we defer further consideration of the sampling period to tests with the nonlinear plant (see next section).

This application is unusual in that the characteristics of the unmeasured disturbance are known. When this is not the case, the *output disturbance* form

^{3.} If a measurement were known to be inaccurate, its R(i, i) value should be relatively large.

^{4.} You can see how serious the degradation would be by setting $R(2,\,2)$ to a large value, e.g., 10000.

of the estimator simplifies the design procedure. It requires only a rough idea of the characteristic times for the disturbances, and the signal-to-noise ratio for each output. For example, you can verify that the following design rejects the d disturbance almost as well as the *optimal* Kalman design:

```
% Alternative estimator design -- output disturbances
taus = [5 5 5];
signoise = [10 \ 10 \ 10];
[Kest, newmod] = smpcest(i mod, taus, si gnoi se);
% Simulation using scmpc -- no model error
[y, u, ym] = scmpc(pmod, newmod, ywt, uwt, M, P, tend, . . .
    setpts, ulim, ylim, Kest, z, v, d);
plotall(y, u, dt)
```

MPC of Nonlinear Plant

We are now ready to test the controller design on the *real* (nonlinear) plant. A special version of the scmpc function (called scmpcnl) is available for this purpose. It uses a nonlinear plant model in the S-function format required by Simulink. (See the Simulink documentation for more information on how to write such models.) The model of the paper machine is in the file pap_mach. m. Simulations with Simulink involving nonlinear models usually take much longer (by an order of magnitude) than linear simulations of a plant of comparable complexity. This is especially likely if the plant model is in the form of an .M file, as is the case here. If such models are to be used extensively, it may be worthwhile to code them as a . mex file (see MATLAB documentation). To see how well the MPC design rejects the *d* disturbance of Figure 3-8, we could use the commands found in the file pm_nonl . m in the directory mpcdemos. The only differences between these commands and those for the original linear simulation are:

- We have defined the initial values of the plant state and manipulated variables (x0 and u0, respectively).
- A step size for numerical integration has been specified. The value of 0.05 minutes provides reasonable accuracy in this application. In general, one must choose the step size to fit the problem (or use a variable step-size integration method, as provided by Simulink).

You can verify that the results are nearly identical to those shown in Figure 3-8. In other words, the nonlinearities in the plant have caused negligible

performance degradation. Very similar results are also obtained for the setpoint change of Figure 3-4.

As the magnitude of the disturbance (or setpoint change) increases, nonlinear effects become significant. For example, Figure 3-9 is for a step in d of 7 units. If the plant were linear, the curves in Figure 3-9 would be the same shape as those in Figure 3-8, but scaled by a factor of 7. Although this is approximately true, there are some qualitative differences. For example, at t = 8 minutes in Figure 3-9, y_2 has gone below y_1 , whereas in Figure 3-8, $y_2 > y_1$ at all times.

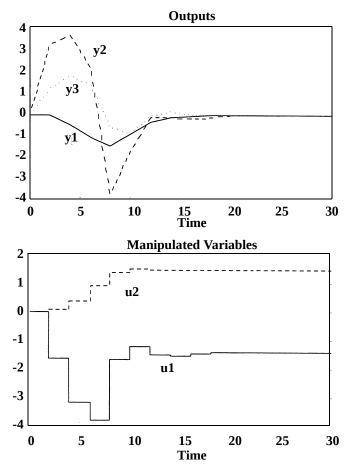


Figure 3-9 As for Figure 3-8, but With Nonlinear Plant, and Step in d of 7 Units

If d is increased to 8, control quality degrades dramatically and the maximum tracking error in y_3 goes to about -10⁵ (not shown). This is caused by changes in the plant characteristics as it moves away from the nominal state (i.e., causing errors in the MPC's linear model).

Sensitivity to modeling error can often be reduced by *de-tuning* the controller. A common approach is to increase the magnitudes of the uwt parameters. When nonlinear effects are severe, however, it may be impossible for *any* time-invariant, linear controller to provide stable, offset-free performance. In that case, if the nonlinear effects are predictable, one might try MPC based on a nonlinear model (e.g., Gattu and Zafiriou, 1992). Scripts for this purpose can be developed using the functions in this toolbox.

As a final test, let's repeat the simulation of Figure 3-8 with a controller sampling period of 0.25 minutes (recall that the original sampling period was 2 minutes). Results appear in Figure 3-10. Compared to Figure 3-8, which had no model error (i.e., linear plant), we reduced the disturbance in y_3 by a factor of 3. Thus, a reduction in sampling period may not lead to robustness problems, and should be tested more thoroughly. You can verify that it works well for other combinations of *small* disturbances and setpoint changes.

Gattu, G. and E. Zafiriou, "Nonlinear Quadratic Dynamic Matrix Control with State Estimation," Ind. Eng. Chem. Research, 1992, 31, 1096–1104.

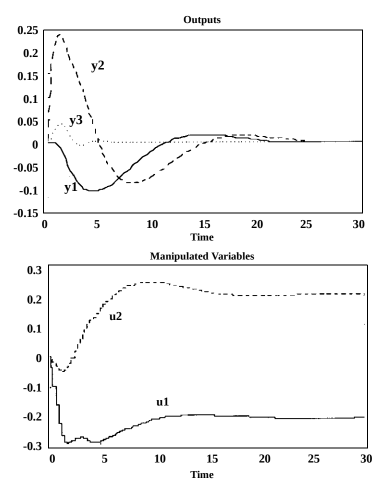


Figure 3-10 As for Figure 3-8, (d=1) but With Nonlinear Plant, Sampling Period of 0.25 Minutes

Command Reference

Commands Grouped by Function

Identification	
autosc	Automatically scales a matrix by its means and standard deviations.
imp2step	Combines MISO impulse response models to form MIMO models in MPC step format.
ml r	Calculates MISO impulse response model via multivariable linear regression.
pl sr	Calculates MISO impulse response model via partial least squares regression.
rescal	Converts scaled data back to its original form.
scal	Scales a matrix by specified means and standard deviations.
val i dmod	Validates a MISO impulse response model using new data.
wrtreg	Writes data matrices used for regression.

Plotting and Matrix Information	
mpci nfo	Outputs matrix type and attributes of system representation.
plotall	Plots outputs and inputs from a simulation run on one graph.
plotfrsp	Plots the frequency response of a system as a Bode plot.
ploteach	Makes separate plots of outputs and/or inputs from a simulation run.
plotstep	Plots the coefficients of a model in MPC step form.

Model Conversions	
c2dmp	Converts state-space model from continuous time to discrete-time. (Equivalent to c2d in Control System Toolbox)
cp2dp	Converts from a continuous to a discrete transfer function in poly format.
d2cmp	Converts state-space model from discrete-time to continuous time. (Equivalent to d2c in Control System Toolbox)
mod2mod	Changes sampling period of a model in MPC mod format.
mod2ss	Converts a model in MPC mod format to a state-space model.
mod2step	Converts a model in MPC mod format to MPC step format.
poly2tfd	Converts a transfer function in poly format to MPC <i>tf</i> format.
ss2mod	Converts a state-space model to MPC mod format.
ss2step	Converts a state-space model to MPC step format.
ss2tf2	Converts state-space model to transfer function. (Equivalent to ss2tf in Control System Toolbox)
tf2ssm	Converts transfer function to state-space model. (Equivalent to tf2ss in Control System Toolbox)
tfd2mod	Converts a model in MPC tf format to MPC mod format.
tfd2step	Converts a model in MPC tf format to MPC step format.
th2mod	Converts a model in theta format (System Identification Toolbox) into MPC mod format.

Model Building — MPC mod format	
addmd	Adds one or more measured disturbances to a plant model.
addmod	Combines two models such that the output of one adds to the input of the other.
addumd	Adds one or more unmeasured disturbances to a plant model.
appmod	Appends two models in an unconnected, parallel structure.
paramod	Puts two models in parallel such that they share a common output.
sermod	Puts two models in series.

Controller Design and Simulation — MPC step format	
стрс	Solves the quadratic programming problem to simulate performance of a closed-loop system with input and output constraints.
mpccl	Creates a model in MPC mod format of a closed-loop system with an unconstrained MPC controller.
mpccon	Calculates the unconstrained controller gain matrix for MPC.
mpcsi m	Simulates a closed-loop system with optional saturation constraints on the manipulated variables.
nl cmpc	Simulink S-function block for MPC controller with input and output constraints (solves quadratic program).
nl mpcsi m	Simulink S-function block for MPC controller with optional saturation constraints.

Controller Design and Simulation — MPC mod format	
scmpc	Solves the quadratic programming problem to simulate performance of a closed-loop system with input and output constraints.
smpccl	Creates a model in MPC mod format of a closed-loop system with an unconstrained MPC controller.
smpccon	Calculates the unconstrained controller gain matrix for MPC.
smpcest	Designs a state estimator for use in MPC.
smpcsi m	Simulates a closed-loop system with optional saturation constraints on the manipulated variables.

Analysis	
mod2frsp	Calculates frequency response for a system in MPC mod format.
smpcgai n	Calculates steady-state gain matrix of a system in MPC mod format.
smpcpol e	Calculates poles of a system in MPC mod format.
svdfrsp	Calculates singular values of a frequency response.

Utility Functions	
abcdchkm	Checks dimensional consistency of (A,B,C,D) set. (Equivalent to abcdchk in Control System Toolbox)
dantzgmp	Solves quadratic programs.
darei ter	Solves discrete Riccati equation by an iterative method.
di mpul sm	Generates impulse response of discrete-time system. (Equivalent to di mpul se in Control System Toolbox)
dl qe2	Calculates state-estimator gain matrix for discrete systems.
dl si mm	Simulates discrete-time systems. (Equivalent to dl si m in Control System Toolbox)
mpcaugss	Augments a state-space model with its outputs.
mpcparal	Puts two state-space models in parallel.
nargchkm	Checks number of M-file arguments. (Equivalent to nargchk in Control System Toolbox)
mpcstai r	Creates the <i>stairstep</i> format used to plot manipulated variables.
vec2mat	Converts a vector to a matrix.

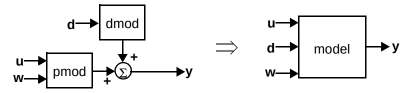
Adds one or more *measured* disturbances to a plant model in the MPC **mod** format. Used to allow for feedforward compensation in MPC.

Syntax

model = addmd(pmod, dmod)

Description

The disturbance model contained in dmod adds to the plant model contained in pmod to form a composite, model, with the structure given in the following block diagram:



pmod, dmod and model are in the MPC mod format (see mod in the online $MATLAB\ Function\ Reference$ for a detailed description). You would normally create pmod and dmod using either the tfd2mod, ss2mod or th2mod functions.

addmd is a specialized version of paramod. Its main advantage over paramod is that it assumes all the inputs to dmod are to be measured disturbances. This saves you the trouble of designating the input types in a separate step.

Example

See ss2mod for an example of the use of this function.

Algorithm

addmd converts pmod and dmod into their state-space form, then uses the mpcparal function to build the composite model.

Restrictions

- pmod and dmod must have been created with equal sampling periods and number of output variables.
- pmod must not include measured disturbances, i.e., its **mod** format must specify $n_d = 0$.
- All inputs to dmod must be classified as manipulated variables. (They will be reclassified automatically as measured disturbances in model.) So the **mod** format of dmod must specify $n_d = n_w = 0$ (which is the default for all model creation functions).

See Also

addmod, addumd, appmod, paramod, sermod

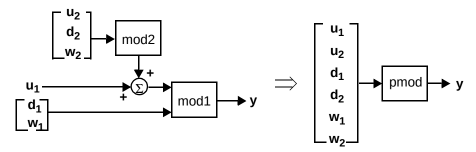
Combines two models in the MPC **mod** format such that the output of one combines with the *manipulated* inputs of the other. This function is specialized and rarely needed. Its main purpose is to build up a model of a complex structure that includes the situation shown in the diagram below.

Syntax

pmod = addmod(mod1, mod2)

Description

The output(s) of mod2 add to the manipulated variable(s) of mod1 to form a composite system, pmod, with the structure given in the following block diagram:



pmod, mod1 and mod2 are in the MPC **mod** format (see mod in the online *MATLAB Function Reference* for a detailed description). You would normally create mod1 and mod2 using either the tfd2mod, ss2mod or th2mod functions.

The different input *types* associated with mod1 and mod2 will be retained in pmod and will be ordered as shown in the diagram.

Example

See mod2ss for an example of the use of this function.

Restrictions

- \bullet mod1 and mod2 must have been created with equal sampling periods.
- The number of manipulated variables in mod1 must equal the number of output variables in mod2.

See Also

addmd, addumd, appmod, paramod, sermod

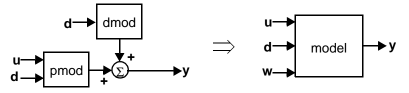
Adds one or more *unmeasured* disturbances to a plant model in MPC **mod** format. Used for simulation of disturbances and for design of state estimators in MPC.

Syntax

model = addumd(pmod, dmod)

Description

The disturbance model contained in dmod adds to the plant model contained in pmod to form a composite, model , with the structure given in the following block diagram:



pmod, dmod and model are in the MPC **mod** format (see mod in the online *MATLAB Function Reference* for a detailed description). You would normally create pmod and dmod using either the tfd2mod, ss2mod or th2mod functions.

addumd is a specialized version of paramod. Its main advantage over paramod is that it assumes all the inputs to dmod are to be unmeasured disturbances. This saves you the trouble of designating the input types in a separate step.

Example

See ss2mod for an example of the use of this function.

Algorithm

addumd converts pmod and dmod into their state-space form, then uses the mpcparal function to build the composite model.

Restrictions

- pmod and dmod must have been created with equal sampling periods and number of output variables.
- pmod must not include unmeasured disturbances, i.e., its **mod** format must specify $n_w = 0$.
- All inputs to dmod must be classified as manipulated variables. (They will be reclassified automatically as unmeasured disturbances in model.) So the **mod** format of dmod must specify $n_d = n_w = 0$ (which is the default for all model creation functions).

See Also

addmod, addmd, appmod, paramod, sermod, smpcest

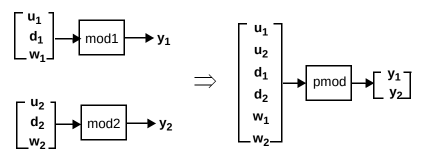
Appends two models to form a composite model that retains the inputs and outputs of the original models. In other words, for models in the MPC **mod** format appmod replaces the append function of the Control Toolbox.

Syntax

pmod = appmod(mod1, mod2)

Description

The two input models combine as shown in the following block diagram:



mod1, mod2 and pmod are in the MPC **mod** format (see mod in the online *MATLAB Function Reference* for a detailed description). You would normally create mod1 and mod2 using either the tfd2mod, ss2mod, or th2mod function.

Restriction

mod1 and mod2 must have been created with equal sampling periods.

See Also

addmod, addmd, addumd, paramod, sermod

Scales a matrix automatically or by specified mean and standard deviation.

Syntax

```
[ax, mx, stdx] = autosc(x)

sx = scal(x, mx)
```

sx = scal (x, mx, stdx) rx = rescal (x, mx) rx = rescal (x, mx, stdx)

Description

autosc scales an input matrix or vector x by its column means (mx) and standard deviations (stdx) automatically and outputs mx and stdx as options. By using scal, the input can also be scaled by some specified means and/or standard deviations. rescal converts scaled data back to original data.

Output mx is a row vector containing the mean value for each column of x while st dx is a row vector containing the standard deviation for each column. Outputs ax and sx are obtained by dividing the difference of each column of x and the mean for the column by the standard deviation for the column, i.e., ax(:,i) = (x:,i) - mx(i)/st dx(i). Output rx is determined by multiplying each column of x by the corresponding standard deviation and adding the corresponding mean to that product.

If only two arguments are specified in scal or rescal, x is scaled by specified means (mx) only.

Example

See ml r for an example of the use of these functions.

See Also

mlr, plsr, wrtreg

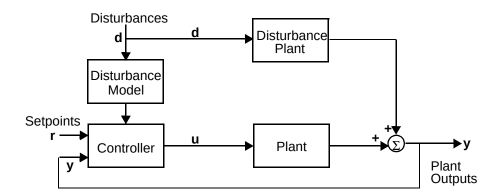
Simulates closed-loop systems with hard bounds on manipulated variables and/or outputs using models in the MPC *step* format. Solves the MPC optimization problem by quadratic programming.

Syntax

yp = cmpc(plant, model, ywt, uwt, M, P, tend, r)
[yp,u,ym] = cmpc(plant,model,ywt,uwt,M,P,tend,...

r,ulim,ylim,tfilter,dplant,dmodel,dstep)

Description



cmpc simulates the performance of the type of system shown in the above diagram when there are bounds on the manipulated variables and/or outputs. Measurement noise can be simulated by treating it as an unmeasured disturbance.

The required input variables are as follows:

plant

Is a model in the MPC *step* format that represents the plant.

model

Is a model in the MPC *step* format that is to be used for state estimation in the controller. In general, it can be different from pl ant if you want to simulate the effect of plant/controller model mismatch.

ywt

Is a matrix of weights that will be applied to the setpoint tracking errors. If $ywt=[\]$, the default is equal (unity) weighting of all out- puts over the entire prediction horizon. If $ywt+[\]$, it must have n_y columns, where n_y is the number of outputs. All weights must be ≥ 0 .

You may vary the weights at each step in the prediction horizon by including up to P rows in ywt. Then the first row of n_y values applies to the tracking errors in the first step in the prediction horizon, the next row applies to the next step, etc. See mpccon for details on the form of the optimization objective function.

If you supply only *nrow* rows, where $1 \le nrow < P$, cmpc will use the last row to fill in any remaining steps. Thus if you want the weighting to be the same for all P steps, you need only specify a single row.

uwt

Same format as ywt, except that uwt applies to the changes in the manipulated variables. If you use uwt = [], the default is zero weighting. If uwt +[], it must have n_{μ} columns, where n_{μ} is the number of manipulated variables.

M

There are two ways to specify this variable:

If it is a scalar , cmpc interprets it as the input horizon (number of moves) as in DMC.

If it is a *row vector* containing n_b elements, each element of the vector indicates the number of steps over which $\Delta u = 0$ during the optimization and cmpc interprets it as a set of n_b blocking factors. There may be $1 \le n_b \le P$ blocking factors, and their sum must be $\le P$

If you set M=[] and P + Inf, the default is M=P, which is equivalent to M=ones(1, P). The default value for M is 1 if P=Inf.

P

The number of sampling periods in the prediction horizon. If P=I nf, the prediction horizon is infinite.

tend

Is the desired duration of the simulation (in time units).

 \mathbf{r}

Is a setpoint matrix consisting of N rows and n_y columns, where n_y is the number of output variables, y:

$$\mathbf{r} = \begin{bmatrix} r_1(1) & r_2(1) & \dots & r_{n_y}(1) \\ r_1(2) & r_2(2) & \dots & r_{n_y}(2) \\ \vdots & \vdots & \dots & \vdots \\ r_1(N) & r_2(N) & \dots & r_{n_y}(N) \end{bmatrix}$$

where $r_i(k)$ is the setpoint for output j at time t = kT, and T is the sampling period (as specified in the step format of pl ant and model). If tend > NT, the setpoints vary for the first N periods in the simulation, as specified by r, and are then held constant at the values given in the last row of r for the remainder of the simulation.

In many simulations one wants the setpoints to be constant for the entire time, in which case r need only contain a single row of n_v values.

If you set $r=[\]$, the default is a row of n_v zeros.

The following input variables are optional. In general, setting one of them equal to an empty matrix causes cmpc to use the default value, which is given in the description.

ul i m

Is a matrix giving the limits on the manipulated variables. Its format is as follows:

$$\begin{aligned} & \text{ulim} = & \begin{bmatrix} u_{min,\,1}(1) & \dots & u_{min,\,n_u}(1) \\ u_{min,\,1}(2) & \dots & u_{min,\,n_u}(2) \\ \vdots & \dots & \vdots \\ u_{min,\,1}(N) & \dots & u_{min,\,n_u}(N) \end{bmatrix} \\ & \begin{bmatrix} u_{max,\,1}(1) & \dots & u_{max,\,n_u}(1) \\ u_{max,\,1}(2) & \dots & u_{max,\,n_u}(2) \\ \vdots & \dots & \vdots \\ u_{max,\,1}(N) & \dots & u_{max,\,n_u}(N) \end{bmatrix} \\ & \begin{bmatrix} \Delta u_{max,\,1}(1) & \dots & \Delta u_{max,\,n_u}(N) \\ \Delta u_{max,\,1}(2) & \dots & \Delta u_{max,\,n_u}(2) \\ \vdots & \dots & \vdots \\ \Delta u_{max,\,1}(N) & \dots & \Delta u_{max,\,n_u}(N) \end{bmatrix} \end{bmatrix}$$

Note that it contains three matrices of N rows. In this case, the limits on N are $1 \le N \le n_b$, where n_b is the number of times the manipulated variables are to change over the input horizon. If you supply fewer than n_b rows, the last row is repeated automatically.

The first matrix specifies the *lower bounds* on the n_u manipulated variables. For example, $u_{min'j}(2)$ is the lower bound for manipulated variable j for the second move of the manipulated variables (where the first move is at the start of the prediction horizon). If $u_{min'j}(k) = -inf$, manipulated variable j will have no lower bound for that move.

The second matrix gives the *upper bounds* on the manipulated variables. If $u_{max,j}(k) = inf$, manipulated variable j will have no upper bound for that move.

The lower and upper bounds may be either positive or negative (or zero) as long as $u_{min',j}(k) \le u_{max,j}(k)$.

The third matrix gives the limits on the rate of change of the manipulated variables. In other words, cmpc will force $|u_j(k) - u_j(k-1)| \le \Delta u_{max,j}(k)$. The limits on the rate of change must be nonnegative and *finite*. If you want it to be unbounded, set the bound to a large number (but not too large — a value of 10^6 should work well in most cases).

The default is $u_{min} = -inf$, $u_{max} = inf$ and $\Delta u_{max} = 10^6$

yl i m

Same idea as for ulim, but for the lower and upper bounds of the outputs. The first row applies to the first point in the prediction horizon. The default is $y_{min} = -inf$, and $y_{max} = inf$.

tfilter

Is a matrix of time constants for the noise filter and the unmeasured disturbances entering at the plant output. The first row of n_y elements gives the noise filter time constants and the second row of n_y elements gives the time constants of the lags through which the unmeasured disturbance steps pass. If tfilter only contains one row, the unmeasured disturbances are assumed to be steps. If you set tfilter=[] or omit it, the default is no noise filtering and steplike unmeasured disturbances.

dpl ant

Is a model in MPC *step* format representing all the disturbances (measured and unmeasured) that affect pl ant in the above diagram. If dpl ant is provided, then input dstep is also required. For output step disturbances, set dpl ant=[]. The default is no disturbances.

dmodel

Is a model in MPC *step* format representing the measured disturbances. If dmodel is provided, then input dstep is also required. If there are no measured disturbances, set dmodel=[]. For output step disturbances, set dmodel=[]. If there are both measured and un-measured disturbances, set the columns of dmodel corresponding to the unmeasured disturbances to zero. The default is no measured disturbances.

dstep

Is a matrix of disturbances to the plant. For output step disturbances (dpl ant=[] and dmodel=[]), the format is the same as for r. For disturbances through step-response models (dpl ant only or both dpl ant and dmodel nonempty), the format is the same as for r, except that the number of columns is n_d rather than n_v The default is a row of zeros.

Notes

- You may use a different number of rows in the matrices r, ulim, ylim and dstep, should that be appropriate for your simulation.
- The ulim constraints used here are fundamentally different from the usat constraints used in the mpcsim function. The ulim constraints are defined relative to the beginning of the prediction horizon, which moves as the simulation progresses. Thus at each sampling period, k, the ulim constraints apply to a block of calculated moves that begin at sampling period k and extend for the duration of the input horizon. The usat constraints, on the other hand, are relative to the fixed point t = 0, the start of the simulation.

The calculated outputs are as follows (all but yp are optional):

yp

Is a matrix containing M rows and n_y columns, where M = max(fix(tend=T) + 1, 2). The first row will contain the initial condition, and row k-1 will give the values of the plant outputs, y (see above diagram), at time t = kT.

u

Is a matrix containing the same number of rows as yp and n_u columns. The time corresponding to each row is the same as for yp. The elements in each row are the values of the manipulated variables, u (see above diagram).

ym

Is a matrix of the same structure as yp, containing the values of the predicted output from the state estimator in the controller. These will, in general, differ from those in yp if model |p| ant and/or there are unmeasured disturbances. The prediction includes the effect of the most recent measurement, i.e., it is $\hat{y}(k|k)$.

For unconstrained problems, cmpc and mpcsi m should give the same results. The latter will be faster because it uses an analytical solution of the QP problem, whereas cmpc solves it by iteration.

Examples

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

The following statements build the model and set up the controller in the same way as in the mpcsi m example.

```
g11=poly2tfd(12.8, [16.7 1], 0, 1);

g21=poly2tfd(6.6, [10.9 1], 0, 7);

g12=poly2tfd(-18.9, [21.0 1], 0, 3);

g22=poly2tfd(-19.4, [14.4 1], 0, 3);

delt=3; ny=2; tfinal=90;

model=tfd2step(tfinal, delt, ny, g11, g21, g12, g22);

plant=model;

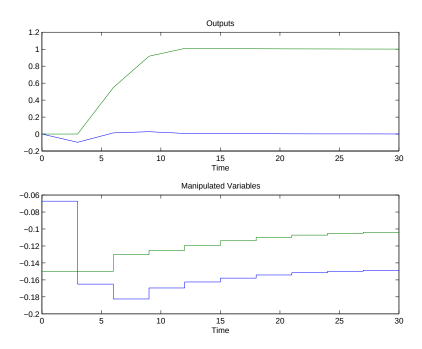
P=6; M=2; ywt=[]; uwt=[1 1];

tend=30; r=[0 1];
```

Here, however, we will demonstrate the effect of constraints. First we set a limit of 0.1 on the rate of change of u_1 and a minimum of -0.15 for u_2 .

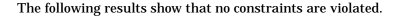
```
\label{eq:limit} $$ ulim=[-inf -0.15 inf inf 0.1 100]; $$ ylim=[\ ]; $$ [y,u]=cmpc(plant,model,ywt,uwt,M,P,tend,r,ulim,ylim); $$ plotall(y,u,delt), pause $$
```

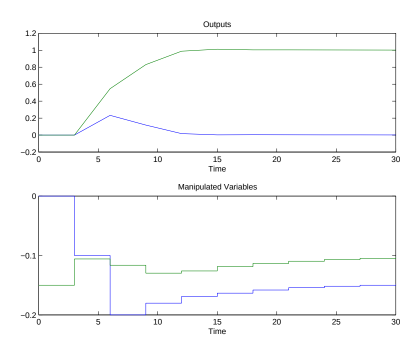
Note that Δu_2 has a large (but finite) limit. It never comes into play.



We next apply a lower bound of zero to both outputs:

```
ulim=[-inf -0.15 inf inf 0.1 100];
ylim=[0 0 inf inf];
[y,u]=cmpc(plant, model, ywt, uwt, M, P, tend, r, ulim, ylim);
plotall(y, u, delt), pause
```





Restriction

Initial conditions of zero are used for all the variables. This simulates the condition where all variables represent a deviation from a steady-state initial condition.

Suggestion

Problems with many inequality constraints can be very time consuming. You can minimize the number of constraints by:

- Using small values for P and/or M.
- Leaving variables unconstrained (limits at $\pm inf$) intermittently unless you think the constraint is important.

See Also

plotall, ploteach, mpccl, mpccon, mpcsi \boldsymbol{m}

Converts a single-input-single-output, continuous-time transfer function in standard MATLAB polynomial form (including an optional time delay) to a sampled-data transfer function.

Syntax

```
[numd, dend] = cp2dp(num, den, delt)
[numd, dend] = cp2dp(num, den, delt, delay)
```

Description

num and den are the numerator and denominator polynomials of the continuous-time system (in the standard Control Toolbox polynomial format), del t is the sampling period, and del ay is the (optional) time delay (in time units). If you omit del ay, cp2dp assumes zero delay. The calculated results are numd and dend, the numerator and denominator polynomials of the corresponding discrete-time transfer function. cp2dp adds a zero-order hold at the input of the continuous-time system during the conversion to discrete-time.

cp2dp accounts properly for the effect of a time delay that is a nonintegral multiple of the sampling period. If del ay=0, cp2dp is equivalent to the MATLAB commands:

```
[a, b, c, d]=tf2ss(num, den);

[phi, gam]=c2dmp(a, b, delt);

[numd, dend]=ss2tf2(phi, gam, c, d, 1);
```

Example

See poly2tfd, poly format for an example of the use of this function.

Algorithm

cp2dp first converts num and den to the equivalent discrete state-space form. It then accounts for the fractional time delay (if any) using the formulas in Åström and Wittenmark (1984), pages 40–42. Finally, it converts the discrete state-space model to a discrete transfer-function model, simultaneously accounting for the whole periods of delay (if any).

Reference

Åström, K. J.; Wittenmark, B. *Computer Control Systems Theory and Design*, Prentice-Hall, Englewood Cliffs, N.J., 1984.

Restriction

The order of num must be \leq that of den.

See Also

poly2tfd, poly format

Solves the discrete Riccati equation by an iterative method to determine the optimal steady-state gain (and optional covariance matrices) for a discrete Kalman filter or state estimator.

Syntax

$$\begin{array}{ll} k = dl \, qe2(phi\,,\,gamw,\,c,\,q,\,r) \\ [\,k,\,m,\,p\,] = dl \, qe2(phi\,,\,gamw,\,c,\,q,\,r) \end{array}$$

Description

Filter form:

Consider the state-space description:

$$x(k+1) = \Phi x(k) + \Gamma_u u(k) + \Gamma_d d(k) + \Gamma_w w(k)$$
$$y(k) = \overline{y}(k) + z(k)$$
$$= Cx(k) + Du(k) + z(k)$$

where x is a vector of n state variables, u contains n_u known inputs, \bar{y} is a vector of n_y measured outputs, y is the *noise-free output*, w is a vector of n_w unmeasured disturbance inputs, z is a vector of n_y measurement noise inputs, and Φ , Γ_w , Γ_w , C and D are constant matrices. We assume that w and z are stationary random-normal signals (white noise) with covariances

$$E\{w(k)w^{T}(k)\} = Q$$

$$E\{w(k)z^{T}(k)\} = R_{12} = 0$$

$$E\{z(k)z^{T}(k)\} = R$$

The steady-state Kalman filter is

$$\hat{x}(k|k) = \hat{x}(k|k-1) + K[y(k) - C\hat{x}(k|k-1) - Du(k)]$$

$$\hat{x}(k+1|k) = \Phi \hat{x}(k|k) + \Gamma_u u(k)$$

$$\hat{y}(k|k) = C\hat{x}(k|k) + Du(k)$$

where $\hat{x}(k|k)$ is the estimate of x(k) based on the measurements available at period k, $\hat{x}(k|k-1)$ is that based on the measurements available at period

k-1, etc. Note that \hat{y} is an estimate of the noise-free output, \bar{y} . The steady-state Kalman gain, K, is the solution of

$$K = MC^{T}[R + CMC^{T}]^{-1}$$

$$M = \Phi P \Phi^T + \Gamma_W Q \Gamma_W^T$$

$$P = M - KCM$$

where M and P may be interpreted as the expected covariance of the errors in the state estimates before and after the measurement update, respectively, i.e.,

$$M = E\{x(k|k-1)x(k|k-1)^T\}$$

$$P = E\{x(k|k)x(k|k)^T\}$$

where, by definition,

$$X(k|k) = X(k) - \hat{X}(k|k)$$

$$X(k|k-1) = X(k) - \hat{X}(k|k-1)$$

The dl qe2 function takes Φ , Γ_{U} , C, R, and Q as inputs and calculates K, M, and P. The last two output arguments are optional.

Note that the input and output arguments are identical to those for dl qe in the Control Toolbox. The advantage of dl qe2 is that it can handle a singular state-transition matrix (Φ) , e.g., for systems with time delay.

Predictor form:

You can also use dl qe2 to calculate a state-estimator in the *predictor* form:

$$\hat{x}(k+1|k) = \Phi\hat{x}(k|k-1) + \Gamma_u u(k) + K_p e(k)$$

$$\hat{y}(k|k-1) = C\hat{x}(k|k-1) + Du(k)$$

$$e(k) = y(k) - \hat{y}(k|k-1)$$

The relationship between K_p , the estimator gain for the predictor form, and K as calculated by $\mathrm{dl}\,\mathrm{qe}2$ is:

$$K_D = \Phi K$$

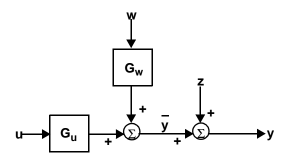
The matrix M calculated by dl qe2 is the expected covariance of the errors in $\mathfrak{X}(k|k-1)$.

Algorithm

 $dl\ qe2\ calls\ darei\ ter^1,$ which solves the discrete algebraic Riccati equation using an iterative doubling algorithm.

Example

Consider a system represented by the block diagram:



^{1.} We gratefully acknowledge Kjell Gustafsson, Department of Automatic Control, Lund Institute of Technology, Lund, Sweden, who provided this function.

where G_u and G_w are first-order, discrete-time transfer functions.

$$G_u(z) = \frac{0.20}{1 - 0.8z^{-1}}$$
 $G_w(z) = \frac{0.3}{1 - 0.95z^{-1}}$

and the statistics of the unmeasured inputs are Q = 2, R = 1.

We use the appropriate MPC Toolbox functions to build a model of the system, then calculate the optimal gain:

```
\begin{array}{l} \mbox{del}\,t\!=\!2; & ny\!=\!1; \\ \mbox{gu=pol}\,y2tfd(0.\,2,\,[\,1\,\,\text{-}\,0.\,8\,],\,del\,t\,)\,; \\ \mbox{Gw=pol}\,y2tfd(0.\,3,\,[\,1\,\,\text{-}\,0.\,95\,],\,del\,t\,)\,; \\ \mbox{[phi,gam,c,d]=mod2ss(tfd2mod(del\,t,\,ny,\,gu,\,Gw))\,;} \\ \mbox{k=dl}\,qe2(phi,\,gam(:\,,\,2)\,,\,c,\,2,\,1) \end{array}
```

The result is:

$$k = 0$$
1.0619

Note that the gain for the first state is zero since this corresponds to the state of G_{uv} which is unaffected by the disturbance, w. Also notice that in the composite system, the second column of gam is Γ_{uv} . This is because of the order in which gu and G_{uv} were specified as inputs to the tfd2mod function.

See Also smpcest

imp2step

Description

Purpose Constructs a multi-input multi-output model in MPC *step* format from

multi-input single-output impulse response matrices.

Syntax plant = i mp2step(delt, nout, theta1, theta2, ...,

theta25)

Given the impulse response coefficient matrices, theta1, theta2, etc., a model in MPC step format is constructed. Each thetai is an n-by- n_u matrix corresponding to the impulse response coefficients for output i. n is the number

of the coefficients and n_u is the number of inputs.

del t is the sampling interval used for obtaining the impulse response coefficients. nout is the output stability indicator. For stable systems, this argument is set equal to number of outputs, n_y . For systems with one or more integrating outputs, this argument is a column vector of length n_y with nout (i) =0 indicating an integrating output and nout (i) =1 indicating a stable

output.

Example See ml r and pl sr for examples of the use of this function.

Restriction The limit on the number of impulse response matrices thetai is 25.

See Also ml r, pl sr

Determines impulse response coefficients for a multi-input single-output system via Multivariable Least Squares Regression or Ridge Regression.

Syntax

```
[theta, yres] = mlr(xreg, yreg, ni nput)
[theta, yres] = mlr(xreg, yreg, ni nput, plotopt, wtheta, ...
    wdel theta)
```

Description

xreg and yreg are the input matrix and output vector produced by routines such as wrtreg. ni nput is number of inputs. Least Squares is used to determine the impulse response coefficient matrix, theta. Columns of theta correspond to impulse response coefficients from each input. Optional output yres is the vector of residuals, the difference between the actual outputs and the predicted outputs.

Optional inputs include pl otopt, wtheta, and wdel theta. No plot is produced if pl otopt is equal to 0 which is the default; a plot of the actual output and the predicted output is produced if pl otopt=1; two plots — plot of actual and predicted output, and plot of residuals — are produced for pl otopt=2. Penalties on the squares of theta and the changes in theta can be specified through the scalar weights wtheta and wdel theta, respectively (defaults are 0). theta is calculated as follows:

$$theta1 = (X^T X)^{-1} X^T Y$$

where

$$X = \begin{bmatrix} xreg \\ wtheta \times I \\ wdeltheta \times delI \end{bmatrix}$$

$$Y = egin{bmatrix} yreg \ 0 \ dots \ 0 \end{bmatrix}$$

where *I* is identity matrix of dimension $n * n_u$

$$deII = \begin{bmatrix} -1 & 1 & 0 & \dots & 0 \\ 0 & -1 & 1 & \dots & 0 \\ & & \vdots & & \\ 0 & \dots & 0 & -1 & 1 \\ 0 & \dots & 0 & 0 & -1 \end{bmatrix}$$

dimension of *delI* is $n * n_n$ by $n * n_n$

then

$$theta = [theta1(1:n) \quad theta1(n+1:2n)... \\ theta1(nu*(n-1)+1:nu*n)]$$

Example

Consider the following two-input single-output system:

$$y(s) = \left[\frac{5.72e^{-14s}}{60s+1} \frac{1.52e^{-15s}}{25s+1}\right] \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

Load the input and output data. The input and output data were generated from the above transfer function and random zero-mean noise was added to the output. Sampling time of 7 minutes was used.

load mlrdat:

Determine the standard deviations for input data using the function autosc.

```
[ax, mx, stdx] = autosc(x);
```

Scale the input data by their standard deviations only.

```
mx = [0, 0];

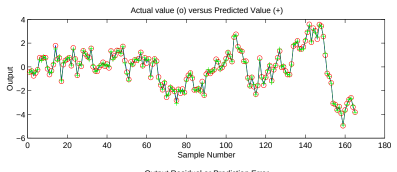
sx = scal(x, mx, stdx);
```

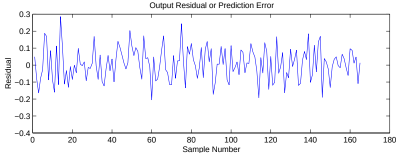
Put the input and output data in a form such that they can be used to determine the impulse response coefficients. 35 impulse response coefficients (n) are used.

```
n = 35;
[xreg, yreg] = wrtreg(sx, y, n);
```

Determine the impulse response coefficients via ml r. By specifying pl ot opt=2, two plots — plot of predicted output and actual output, and plot of the output residual (or predicted error) — are produced.

```
ni nput = 2;
pl otopt = 2;
[theta, yres] = mlr(xreg, yreg, ni nput, pl otopt);
```





Scale theta based on the standard deviations used in scaling the input.

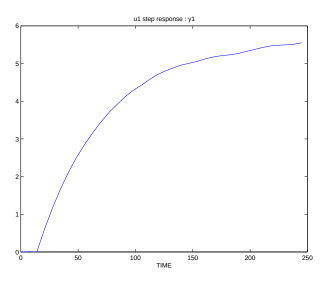
```
theta = scal(theta, mx, stdx);
```

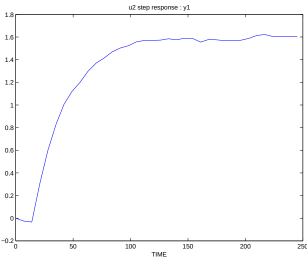
Convert the impulse model to a step model to be used in MPC design. Recall that a sampling time of 7 minutes was used in determining the impulse model. Number of outputs (1 in this case) must be specified.

```
nout = 1;
delt = 7;
model = imp2step(delt, nout, theta);
```

Plot the step response coefficients.

plotstep(model)



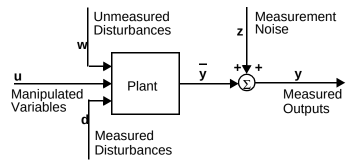


See Also

plsr, validmod, wrtreg

The MPC **mod** format is a compact way to store the model of a linear system for subsequent use in the MPC Toolbox functions.

Description



Consider the process shown in the above block diagram. Its *discrete-time* LTI state-space representation is:

$$x(k+1) = \Phi x(k) + \Gamma_u u(k) + \Gamma_d d(k) + \Gamma_w w(k)$$

$$y(k) = \overline{y}(k) + z(k)$$

$$= Cx(k) + D_u u(k) + D_d d(k) + D_w w(k) + z(k)$$

where x is a vector of n state variables, u represents the n_u manipulated variables, d represents n_d measured but freely-varying inputs (i.e., measured disturbances), w represents n_w unmeasured disturbances, y is a vector of n_y plant outputs, z is measurement noise, and Φ , Γ_{up} etc., are constant matrices of appropriate size. The variable $\bar{y}(k)$ represents the plant output before the addition of measurement noise. Define:

$$\Gamma = [\Gamma_u \Gamma_d \Gamma_w]$$

$$D = [D_u D_{ud} D_w]$$

In some cases one would like to include n_{ym} measured and n_{yu} unmeasured outputs in y, where $n_{ym} + n_{yu} = n_y$. If so, the **mod** format assumes that the y vector and the C and D matrices are arranged such that the measured outputs come first, followed by the unmeasured outputs.

The **mod** format is a single matrix that contains the Φ , Γ , C, and D matrices, plus some additional information. Let M be the **mod** representation of the above system. Its overall dimensions are:

- Number of rows = $n + n_v + 1$
- Number of columns = $max(7, 1 + n + n_u + n_d + n_w)$

The mi nfo vector is the first seven elements of the first row in *M*. The elements of mi nfo are:

- \min nfo (1) T, the sampling period used to create the model.
 - (2) n, the number of states.
 - (3) n_{u} , the number of manipulated variable inputs.
 - (4) n_{d} , the number of measured disturbances.
 - (5) n_w the number of unmeasured disturbances.
 - (6) n_{vm} , the number of measured outputs.
 - (7) n_{yu} , the number of unmeasured outputs.

The remainder of M contains the discrete state-space matrices:

$$\Phi \text{ in rows 2 to } n+1$$
 columns 2 to $n+1$ columns $n+2$ to $n+n_u+n_d+n_w+1$ columns $n+2$ to $n+n_u+n_d+n_w+1$ columns 2 to $n+1$ columns $n+2$ to $n+n_u+n_d+n_w+1$ columns $n+2$ to $n+n_u+n_d+n_w+1$

Notes

Since the minfo vector requires seven columns, this is the minimum possible number of columns in the **mod** format, regardless of the dimensions of the state-space matrices.

Also, the first column is reserved for other uses by MPC Toolbox routines. Thus the state-space matrices start in column 2, as described above.

In order for the mpci nfo routine to recognize matrices in the MPC **mod** format, the (2,1) element is set to NaN (Not-a-Number).

mod format

Example See ss2mod for a **mod** format example.

See Also mod2ss, mod2step, step format, mpcinfo, ss2mod, step, tfd2mod, tf format,

th2mod, theta format

Purpose

Calculates the complex frequency response in *varying* format of a system in MPC **mod** format.

Syntax

frsp = mod2frsp(mod, freq)
[frsp, eyefrsp] = mod2frsp(mod, freq, out, in, balflg)

Description

mod2frsp calculates the complex frequency response of a system (mod) in MPC mod format. The desired frequencies are given by the input freq, a *row vector* of 3 elements specifying the lower frequency as a power of 10, the upper frequency as a power of 10, and the number of frequency points.

Optional inputs out and in are *row vectors* that specify the outputs and inputs for which the frequency response is to be generated. If these variables are omitted or empty, the default is to use all outputs and inputs.

Optional input bal fl g indicates whether the system's Φ matrix should be balanced (using the MATLAB bal ance command). If bal fl g is nonzero, balancing is performed. Balancing improves the conditioning of the problem, but may cause errors in the frequency response. If bal fl g=[] or is omitted, no balancing is performed.

Output frsp is the frequency response matrix given in *varying* format. Let $F(\omega)$ denote a matrix-valued function of the independent variable ω . Then the N sampled values $F(\omega_1), \ldots, F(\omega_N)$ are contained in frsp as follows:

$$\operatorname{frsp} = \begin{bmatrix} F(\omega_1) & \omega_1 \\ \vdots & \vdots \\ F(\omega_i) & \omega_N \\ \vdots & 0 \\ F(\omega_N) & \vdots \\ & 0 \\ 0...0N & inf \end{bmatrix}$$

If the dimension of each submatrix $F(\omega_i)$ is n by m, then the dimensions of frsp is $n \cdot N + 1$ by m + 1.

Optional output eyefrsp is in *varying* format and represents $I - F(\omega_i)$ at each frequency. This output can only be specified for square submatrices and may be useful in computing the frequency responses of both the sensitivity and complementary sensitivity functions.

Example

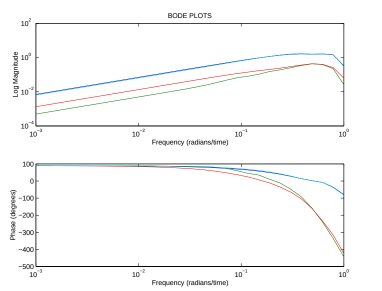
Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

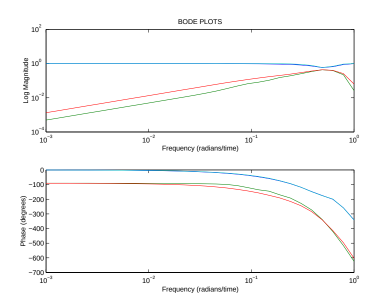
See the mpccl example for the commands that build the a closed-loop model for this process using a simple controller. However for this example, $\det t$ =6 and tfi nal =90 are used to reduce the number of step response coefficients.

Now we will calculate and plot the frequency response of the sensitivity and complementary sensitivity functions.

```
freq = [-3,0,30];
in = [1:ny]; % input is r for comp. sensitivity
out = [1:ny]; % output is yp for comp. sensitivity
[frsp, eyefrsp] = mod2frsp(cl mod, freq, out, in);
plotfrsp(eyefrsp); % Sensitivity
pause;
```

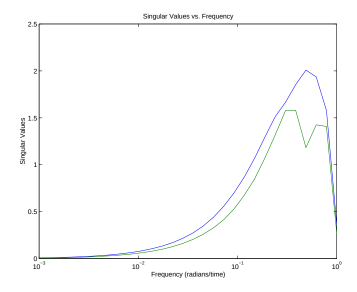


plotfrsp(frsp); % Complementary Sensitivity pause;



Calculate and plot the singular values for the sensitivity function response.

```
[sigma, omega] = svdfrsp(eyefrsp);
clg;
semilogx(omega, sigma);
title('Singular Values vs. Frequency');
xlabel('Frequency (radians/time)');
ylabel('Singular Values');
```



Algorithm

The algorithm to calculate the complex frequency response involves a matrix inverse problem which is solved via a Hessenberg matrix.

Reference

A.J. Laub, "Efficient Multivariable Frequency Response Computations," *IEEE Transactions on Automatic Control*, Vol. AC–26, No. 2, pp.407–408, April, 1981.

See Also

mod, mpccl, plotfrsp, smpccl, svdfrsp

mod2mod

Purpose Changes the sampling period of a model in MPC **mod** format.

Syntax newmod = mod2mod(ol dmod, del t2)

Description Input ol dmod is the existing model in MPC mod format. Input del t2 is the new

sampling period for the model. mod2mod returns newmod, which is the system in

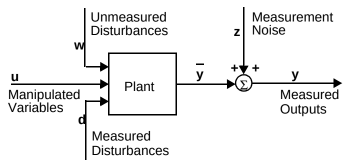
mod format converted to the new sampling time.

See Also mod, ss2mod

Extracts the standard discrete-time state-space matrices and other information from a model stored in the MPC **mod** format.

Syntax

Description



Consider the process shown in the above block diagram. mod2ss assumes that mod is a description of the above process in the MPC **mod** format (see mod in the online *MATLAB Function Reference* for more details). An equivalent state-space representation is:

$$x(k+1) = \Phi x(k) + \Gamma_u u(k) + \Gamma_d d(k) + \Gamma_w w(k)$$

$$y(k) = \overline{y}(k) + z(k)$$

$$= Cx(k) + D_u u(k) + D_d d(k) + D_w w(k) + z(k)$$

where x is a vector of n state variables, u represents the n_u manipulated variables, d represents n_d measured but freely-varying inputs (i.e., measured disturbances), w represents n_w unmeasured disturbances, y is a vector of n_y plant outputs, z is measurement noise, and Φ , Γ_u , etc., are constant matrices of appropriate size. The variable \bar{y} (k) represents the plant output before the addition of measurement noise. Define:

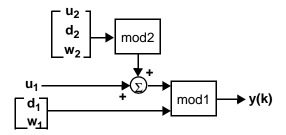
$$\Gamma = [\Gamma_u \Gamma_d \Gamma_w]$$

$$D = [D_u D_d D_w]$$

mod2ss extracts the Φ , Γ , C, and D matrices from the input variable, mod. It also extracts the vector mi nfo, which contains additional information about the sampling period, number of each type of input and output, etc. see mod in the online MATLAB Function Reference for more details on mi nfo.

Examples

- 1 See the example in the description of dl qe2.
- 2 Suppose you have a plant with the structure



where the inputs and outputs are all scalars, and you have constructed mod1 and mod2 using the commands:

Now you want to calculate the response to a step change in d_2 , which is the *fourth* input to the composite system, pmod. One way to do it is:

```
[phi, gam, c, d, minfo] = mod2ss(pmod);
nstep=10;
ustep=[zeros(nstep, 3) ones(nstep, 1) zeros(nstep, 2)];
% Define step in d2
y=dlsimm(phi, gam, c, d, ustep);
% simulate response to step input
plot([0:nstep-1], y)
```

The results of the mod2ss call are:

c= 0.2000 - 0. 4000 0 d=

1.0000

0 0 0 0 0 0 mi nfo= 3 2 1 2 2 1

0

- . 5000

0

0

0

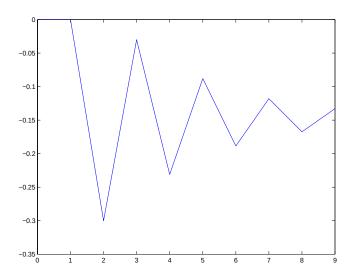
0

0

0.2000

And the step response is:

0



See Also

mod, ss2mod

Uses a model in the **mod** format to calculate the step response of a SISO or MIMO system in MPC *step* format.

Syntax

```
pl ant = mod2step(mod, tfi nal)
[pl ant,dpl ant] = mod2step(mod, tfi nal, del t2, nout)
```

Description

The input variable mod is assumed to be a model in the **mod** format (see mod in the online *MATLAB Function Reference* for a description). You would normally create it using ss2mod, tfd2mod, or th2mod. The input variable tfi nal is the time at which you would like to end the step response.

The optional input variable del t2 is the desired sampling period for the step response. If you use del $t2=[\]$ or omit it, the default is equal to the sampling period of mod (contained in the mi nfo vector of mod).

The optional input variable nout is the output stability indicator. For stable systems, set nout equal to the number of outputs, n_y . For systems with one or more integrating outputs, nout is a column vector of length n_y with nout (i) =0 indicating an integrating output and nout (i) =1 indicating a stable output. If you use nout=[] or omit it, the default is nout= n_y (only stable outputs).

pl ant and dpl ant are matrices in MPC *step* format containing the calculated step responses. pl ant is the response to the manipulated variables, and dpl ant is the response to the disturbances (if any), both measured and unmeasured. The overall dimensions of these matrices are:

```
plant n-by-n_y + n_y + 2 rows, n_u columns.

dpl ant n-by-n_y + n_y + 2 rows, n_d + n_w columns.

where n = round (tfi nal /del t2)
```

It is assumed that stable step responses are nearly constant after n sampling periods, while integrating responses increase with a constant slope after n – 1 sampling periods.

Each column gives the step response with respect to the corresponding input variable. Within each column, the first n_y elements are the response for each output at time t = T, the next n_y elements give each output at time t = 2T, etc.

mod2step, step format

The last n_y + 2 rows contain nout, n_y and del t2, respectively (all in column 1 — any remaining elements in these rows are set to zero). In other words, for pl ant the arrangement is as follows:

$$\text{plant} = \begin{bmatrix} S_1 & & & & \\ & S_2 & & & \\ & \vdots & & & \\ & S_n & & & \\ & & & \\ & & & S_n & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & &$$

where

$$S_{i} = \begin{bmatrix} S_{1, 1, i} & S_{1, 2, i} & \dots & S_{1, n_{w} i} \\ S_{2, 1, i} & S_{2, 2, i} & \dots & S_{2, n_{w} i} \\ \vdots & & & & \\ S_{n_{y}, 1, i} & S_{n_{y}, 2, i} & \dots & S_{n_{y}, n_{w} i} \end{bmatrix}$$

 S_{kji} is the \emph{i}^{th} step response coefficient describing the effect of input \emph{j} on output \emph{k} .

The arrangement of dpl ant is similar; the only difference is in the number of columns.

Example

The following process has 3 inputs and 4 outputs:

```
phi =di ag([0.3, 0.7, -0.7]);

gam=eye(3);

c=[1 0 0; 0 0 1; 0 1 1; 0 1 0];

d=[1 0 0; zeros(3,3)];
```

We first calculate its step response for 4 samples (including the initial condition) with respect to each of the inputs using the Control Toolbox function, dstep:

```
nstep=4; del t=1.5;
yu1=dstep(phi, gam, c, d, 1, nstep)
yu2=dstep(phi, gam, c, d, 2, nstep)
yu3=dstep(phi, gam, c, d, 3, nstep)
```

The results are:

	Response to u ₁				Response to u ₂				Response to u ₃			
Time	<i>y</i> ₁	<i>y</i> ₂	<i>y</i> ₃	<i>y</i> ₄	<i>y</i> ₁	<i>y</i> ₂	<i>y</i> ₃	<i>y</i> ₄	<i>y</i> ₁	<i>y</i> ₂	<i>y</i> ₃	<i>y</i> ₄
0	1	0	0	0	0	0	0	0	0	0	0	0
Т	2	0	0	0	0	0	1	1	0	1	1	0
2T	2.3	0	0	0	0	0	1.7	1.7	0	0.3	0.3	0
3T	2.39	0	0	0	0	0	2.19	2.19	0	0.79	0.79	0

We then use mod2step to do the same job:

```
plant=mod2step(ss2mod(phi, gam, c, d, delt), (nstep-1)*delt)
```

mod2step, step format

obtaining the results:

pl ant =		
2. 0000	0	0
0	0	1. 0000
0	1. 0000	1. 0000
0	1. 0000	0
2. 3000	0	0
0	0	0. 3000
0	1. 7000	0. 3000
0	1. 7000	0
2. 3900	0	0
0	0	0. 7900
0	2. 1900	0. 7900
0	2. 1900	0
1. 0000	0	0
1. 0000	0	0
1. 0000	0	0
1. 0000	0	0
4. 0000	0	0
1. 5000	0	0

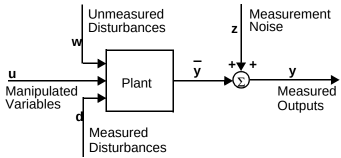
See Also

plotstep, ss2step, tfd2step

Differences the states of a system and augments them with the output variables. Mainly used as a utility function for setting up model predictive controllers.

Syntax

Description



Consider the process shown in the above block diagram. A state-space representation is:

$$z(k+1) = \Phi x(k) + \Gamma_u u(k) + \Gamma_d d(k) + \Gamma_w w(k)$$
$$y(k) = \overline{y}(k) + D_u u(k) + D_d d(k) + D_w w(k) + z(k)$$
$$= \overline{y}(k) + z(k)$$

where x is a vector of n state variables, u is a vector of n_u manipulated variables, d is a vector of n_d measured disturbances, w is a vector of n_w unmeasured disturbances, y is a vector of n_y plant outputs, z is measurement noise, and Φ , Γ_{u^*} Γ_{d^*} Γ_{w^*} etc., are constant matrices of appropriate size. The variable y(k) = Cx(k) represents the plant output before the addition of the direct contribution of the inputs $[D_u u(k) + D_d v(k) + D_w w(k)]$ and the measurement noise [z(k)]. (The variable \bar{y} is the output before addition of the measurement noise). Define:

$$\Delta u(k) = u(k) - u(k-1)$$

$$\Delta x(k) = x(k) - x(k-1)$$

etc. Then equations 4.28 and 4.29 can be converted to the form

$$x_a(k+1) = \Phi_a x_a(k) + \Gamma_{ua} \Delta u(k) + \Gamma_{da} \Delta(k) + \Gamma_{wa} \Delta w(k)$$

$$y(k) = C_a x_a(k) + D_u u(k) + D_d d(k) + D_w w(k) + z(k)$$

where, by definition,

$$\begin{split} x_{a}(k) &= \begin{bmatrix} \Delta x(k) \\ y(k) \end{bmatrix} \\ \Phi_{a} &= \begin{bmatrix} \Phi & 0 \\ C\Phi & I \end{bmatrix} \quad \Gamma_{a} = \begin{bmatrix} \Gamma_{ua} & \Gamma_{da} & \Gamma_{wa} \end{bmatrix} \\ \Gamma_{ua} &= \begin{bmatrix} \Gamma_{u} \\ C\Gamma_{u} \end{bmatrix} \quad \Gamma_{da} = \begin{bmatrix} \Gamma_{d} \\ C\Gamma_{d} \end{bmatrix} \quad \Gamma_{wa} = \begin{bmatrix} \Gamma_{w} \\ C\Gamma_{w} \end{bmatrix} \\ C_{a} &= \begin{bmatrix} 0 & I \end{bmatrix} \quad D_{a} = \begin{bmatrix} D_{u} & D_{d} & D_{w} \end{bmatrix} \end{split}$$

The mpcaugss function takes the matrices Φ , Γ (= [$\Gamma_u \Gamma_d \Gamma_w$]), C as input, and creates the augmented matrices Φ_a , Γ_a , C_a and D_a in the form shown above. The D input matrix is optional. If you include it, mpcaugss assumes it has the form $D = [D_u D_d D_w]$. If you omit it, the default is zero. Note that all MPC design routines require $D_u = D_d = 0$.

The last output variable, na, is the order of the augmented system, i.e., $n_a = n + n_v$. It is optional.

Example

The following system has 2 states, 3 inputs, and 2 outputs.

```
phi =di ag([0.8, -0.2]);
gam=[1 -1 0; 0 2 -0.5];
c=[0.4 0; 0 1.5];
```

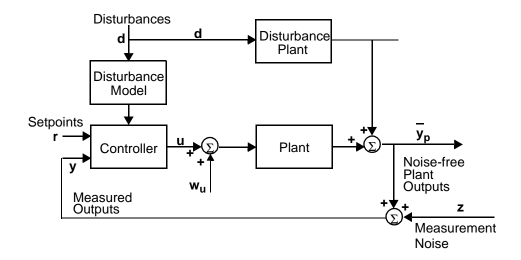
Here is the augmentation command, followed by the calculated results:

Combines a plant model and a controller model in MPC *step* format, yielding a closed-loop system model in the MPC **mod** format. This can be used for stability analysis and linear simulations of closed-loop performance.

Syntax

```
[cl mod] = mpccl(pl ant, model, Kmpc)
[cl mod, cmod] = mpccl(pl ant, model, Kmpc, tfilter, ...
dpl ant, dmodel)
```

Description



plant

Is a model (in *step* format) representing the plant in the above diagram.

model

Is a model (in *step* format) that is to be used to design the MPC controller block shown in the diagram. It may be the same as plant (in which case there is no "model error" in the controller design), or it may be different.

Kmpc

Is a controller gain matrix, which was calculated by the function mpccon.

tfilter

Is a (optional) matrix of time constants for the noise filter and the unmeasured disturbances entering at the plant output. If omitted or set to an empty matrix, the default is no noise filtering and steplike unmeasured disturbances. See the documentation for the function mpcsi m for more details on the design and proper format of tfilter.

dpl ant

Is a (optional) model (in *step* format) representing all the disturbances (measured and unmeasured) that affect pl ant in the above diagram. If omitted or set to an empty matrix, the default is that there are no disturbances.

dmodel

Is a (optional) model (in *step* format) representing the measured disturbances. If omitted or set to an empty matrix, the default is that there are no measured disturbances. See the documentation for the function mpcsi m for more details on how disturbances are handled when using step-response models.

mpccl

Calculates a model of the closed-loop system, cl mod. It is in the **mod** format and can be used, for example, with analysis functions such as smpcgain and smpcpole, and with simulation routines such as mod2step and dl si mm. mpccl also calculates (as an option) a model of the controller element, cmod.

The closed-loop model, cl mod, has the following state-space representation:

$$x_{cl}(k+1) = \Phi_{cl}x_{cl}(k) + \Gamma_{cl}u_{cl}(k)$$

 $y_{cl}(k) = C_{cl}x_{cl}(k) + D_{cl}u_{cl}(k)$

where x_{cl} is a vector of n state variables, u_{cl} is a vector of input variables, y_{cl} is a vector of outputs, and Φ_{ch} Γ_{ch} C_{ch} and D_{cl} are matrices of appropriate size. The expert user may want to know the significance of the state variables in x_{cl} . They are (in the following order):

- The n_p states of the plant (as specified in pl ant),
- \bullet The n_i state estimates (based on the model specified in model),
- n_d integrators that operate on the Δd signal to yield a d signal. If there are no disturbances, these states are omitted.

- n_u integrators that operate on the Δw_u signal to yield a w_u signal.
- n_u integrators that operate on the Δu signal produced by the standard MPC formulation to yield a u signal that can be used as input to the plant and as a closed-loop output.

The closed-loop input and output variables are:

$$u_{cl}(k) = \begin{bmatrix} r(k) \\ z(k) \\ w_{u}(k) \\ d(k) \end{bmatrix} \text{ and } y_{cl}(k) = \begin{bmatrix} \bar{y}_{p}(k) \\ u(k) \\ \hat{y}(k|k) \end{bmatrix}$$

where $\hat{y}(k|k)$ is the estimate of the noise-free plant output at sampling period k based on information available at period k. This estimate is generated by the controller element.

The state-space form of the controller model, cmod, can be written as:

$$x_c(k+1) = \Phi_c x_c(k) + \Gamma_c u_c(k)$$

$$y_c(k) = C_c x_c(k) + D_c u_c(k)$$

where

$$u_c(k) = \begin{bmatrix} r(k) \\ y(k) \\ d(k-1) \end{bmatrix} \text{ and } y_c(k) = u(k-1)$$

and the controller states are the same as those of the closed loop system *except* that the n_p plant states are not included.

Example Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

We build the step response model using the MPC Toolbox functions poly2tfd and tfd2step.

```
\begin{array}{l} g11 = pol\ y2tfd(12.\ 8,\ [16.\ 7\ 1],\ 0,\ 1)\,;\\ g21 = pol\ y2tfd(6.\ 6,\ [10.\ 9\ 1],\ 0,\ 7)\,;\\ g12 = pol\ y2tfd(-18.\ 9,\ [21.\ 0\ 1],\ 0,\ 3)\,;\\ g22 = pol\ y2tfd(-19.\ 4,\ [14.\ 4\ 1],\ 0,\ 3)\,;\\ del\ t = 3;\ ny = 2;\ tfi\ nal\ =\ 60;\\ model\ = tfd2step(tfi\ nal\ ,\ del\ t,\ ny,\ g11,\ g21,\ g12,\ g22)\,;\\ pl\ ant = model\ ;\ \%\ No\ pl\ ant / model\ mi\ smatch \end{array}
```

Now we design the controller. Since there is delay, we use M < P: We specify the defaults for the other tuning parameters, uwt and ywt, then calculate the controller gain:

```
P=6; % Prediction horizon.
M=2; % Number of moves (input horizon).
ywt=[]; % Output weights (default - unity
% on all outputs). uwt=[]; % Man. Var weights (default - zero
% on all man. vars).
Kmpc=mpccon(model, ywt, uwt, M, P);
```

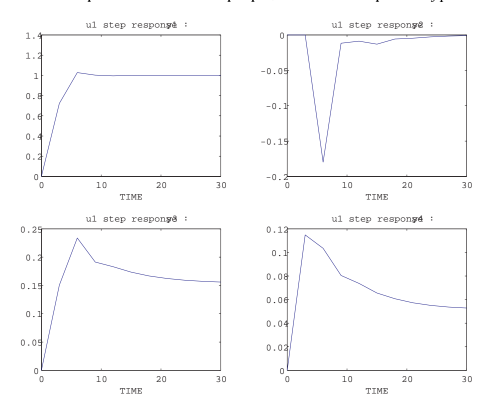
Now we can calculate the model of the closed-loop system:

```
cl mod=mpccl (pl ant, model, Kmpc);
```

You can use the closed-loop model to calculate and plot the step response with respect to all the inputs. The appropriate commands are:

```
tend=30;
clstep=mod2step(clmod, tend);
plotstep(clstep)
```

Since the closed-loop system has m = 6 inputs and p = 6 outputs, only one of the plots is reproduced here. It shows the response of the first 4 closed-loop outputs to a unit step in the first closed-loop input, which is the setpoint for y_1 :



Closed-loop outputs y_1 and y_2 are the true plant outputs (noise-free). Output y_1 goes to the new setpoint quickly with a small overshoot. This causes a small, short-term disturbance in y_2 . The plots for y_3 and y_4 show the required variation in the manipulated variables.

Restriction

model and plant must have been created using the same sampling period.

See Also

cmpc, mod2step, step format, mpccon, mpcsi m, smpcgai n, smpcpol e

Purpose

Calculates MPC controller gain using a model in MPC *step* format.

Syntax

```
Kmpc = mpccon(model)
```

Kmpc = mpccon(model, ywt, uwt, M, P)

Description

Combines the following variables (most of which are optional and have default values) to calculate the MPC gain matrix, Kmpc.

model

is the model of the process to be used in the controller design (in the *step* format).

The following input variables are optional:

ywt

Is a matrix of weights that will be applied to the setpoint tracking errors. If you use $ywt=[\]$ or omit it, the default is equal (unity) weighting of all outputs over the entire prediction horizon. If you use $ywt+[\]$, it must have n_y columns, where n_y is the number of outputs. All weights must be ≥ 0 .

You may vary the weights at each step in the prediction horizon by including up to P rows in ywt. Then the first row of n_y values applies to the tracking errors in the first step in the prediction horizon, the next row applies to the next step, etc.

If you supply only nrow rows, where $1 \le nrow < P$, mpccon will use the last row to fill in any remaining steps. Thus if you wish the weighting to be the same for all P steps, you need only specify a single row.

uwt

Same format as ywt, except that uwt applies to the changes in the manipulated variables. If you use uwt = $[\]$ or omit it, the default is zero weighting. If uwt $[\]$, it must have n_u columns, where n_u is the number of manipulated variables.

M

There are two ways to specify this variable:

- If it is a scalar, mpccon interprets it as the input horizon (number of moves) as in DMC.
- If it is a *row vector* containing n_b elements, each element of the vector indicates the number of steps over which $\Delta u = 0$ during the optimization and cmpc interprets it as a set of n_b blocking factors. There may be $1 \le n_b \le P$ blocking factors, and their sum must be $\le P$.

If you set $M=[\]$ or omit it and P+I nf, the default is M=P, which is equivalent to M=ones(1,P). The default value for M is 1 if P=I nf.

P

The number of sampling periods in the prediction horizon. If you set P=I nf or omit it, the default is P=1. If P=i nf, the prediction horizon is infinite.

If you take the default values for all the optional variables, you get the "perfect controller," i.e., a model-inverse controller. This controller is not applicable when one or more outputs can not respond to the manipulated variables within one sampling period due to time delay. In this case, the plant-inverse controller is unrealizable. For nonminimum phase discrete plants, this controller is unstable. To counteract this you can penalize changes in the manipulated variables (variable uwt), use blocking (variable M), and/or make P>>M The model-inverse controller is also relatively sensitive to model error and is best used as a point of reference from which you can progress to a more robust design.

Algorithm

The controller gain is a component of the solution to the optimization problem:

Minimize
$$J(k) = \sum_{j=1}^{p} \sum_{i=1}^{n_y} (ywt_i(j)[r_i(k+j) - \hat{y}_i(k+j)])^2 + \sum_{j=1}^{n_b} \sum_{i=1}^{n_u} (uwt_i(j)\Delta \hat{u}_i(j))^2$$

with respect to $\Delta u_j(j)$ (a series of current and future moves in the manipulated variables), where $\hat{y}_i(\mathbf{k}+j)$ is a prediction of output i at a time j sampling periods into the future (relative to the current time, k), which is a function of

 $\Delta \hat{u}_i(j)$, $r_i(k+j)$ is the corresponding future setpoint, and n_b is the number of blocks or moves of the manipulated variables.

Example

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

See the mpccl example for the commands that build the model and a simple controller for this process.

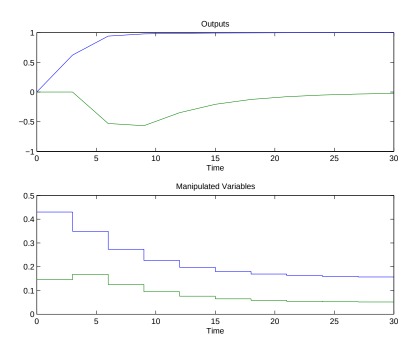
Here is a slightly more complex design with blocking and time-varying weights on the manipulated and output variables:

```
\begin{array}{l} P=6; \ \ M=[2\ 4]; \\ uwt=[1\ 0;\ 0\ 1]; \\ ywt=[1\ 0.\ 1;\ 0.\ 8\ 0.\ 1;\ 0.\ 1\ 0.\ 1]; \\ Kmpc=mpccon(model, ywt, uwt, M, P); \\ tend=30; \ \ r=[1\ 0]; \\ [y,u]=mpcsim(plant, model, Kmpc, tend, r); \end{array}
```

There is no particular rationale for using time varying weights in this case it is only for illustration. The manipulated variables will make 2 moves during the prediction horizon (see value of M, above). The uwt selection gives u_1 a unity weight and u_2 a zero weight for the first move, then switches the weights for the second move. If there had been any additional moves they would have had the same weighting as the second move.

mpccon

The ywt value assigns a constant weight of 0.1 to y_2 , and a weight that decreases over the first 3 periods to y_1 . The weights for periods 4 to 6 are the same as for period 3. The resulting closed-loop (servo) response is:



See Also cmp

cmpc, mpccl, mpcsi m

Purpose

Determines the type of a matrix and returns information about the matrix.

Syntax:

mpci nfo(mat)

Description

mpci nfo returns information about the type and size of the matrix, mat. The information is determined from the matrix structure. The matrix types include MPC *step* format, MPC **mod** format, *varying* format and constant. mpci nfo returns text output to the screen.

If the matrix is in MPC *step* format, the output includes the sampling time used to create the model, number of inputs, number of outputs and number of step response coefficients; it also indicates which outputs are stable and which outputs are integrating.

If the matrix is in MPC **mod** format, the output includes the sampling time used to create the model, number of states, number of manipulated variable inputs, number of measured disturbances, number of unmeasured disturbances, number of measured outputs and number of unmeasured outputs.

For a matrix in *varying* format, as formed in mod2frsp, the number of independent variable values, and the number of rows and number of columns of each submatrix are output.

For a constant matrix, the text output consists of the number of rows and number of columns.

Examples

1 MPC *step* format: After running the mod2step example mpci nfo(pl ant) returns:

```
This is a matrix in MPC Step format. sampling time = 1.5 number of inputs = 3 number of outputs = 4 number of step response coefficients = 3 All outputs are stable.
```

2 MPC **mod** format: After running the ss2mod example mpci nfo(pmod) returns:

```
This is a matrix in MPC Mod format.

minfo = [2 3 1 1 1 1 0 ]

sampling time = 2

number of states = 3

number of manipulated variable inputs = 1

number of measured disturbances = 1

number of measured outputs = 1

number of unmeasured outputs = 0
```

3 *varying* format: After running the mod2frsp example mpci nfo(eyefrsp) returns:

```
varying: 30 pts 2 rows 2 cols
```

See Also

mod, step, mod2frsp, varying format

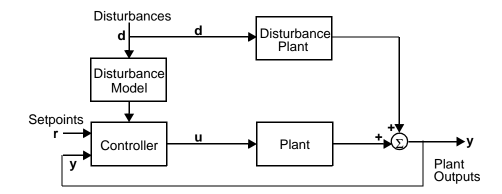
Purpose

Simulates closed-loop systems with saturation *constraints* on the manipulated variables using models in the MPC *step* format. Can also be used for open-loop simulations.

Syntax

```
yp = mpcsim(plant, model, Kmpc, tend, r)
[yp, u, ym] = mpcsim(plant, model, Kmpc, tend, r, usat, ...
tfilter, dplant, dmodel, dstep)
```

Description



mpcsi m provides a convenient way to simulate the performance of the type of system shown in the above diagram. Measurement noise can be simulated by treating it as an unmeasured disturbance. The required input variables are as follows:

pl ant

Is a model in the MPC *step* format that is to represent the plant.

model

Is a model in the MPC *step* format that is to be used for state estimation in the controller. In general, it can be different from pl ant if you want to simulate the effect of plant/controller model mismatch. Note, however, that model should be the same as that used to calculate Kmpc.

Kmpc

Is the MPC controller gain matrix, usually calculated using the function mpccon.

If you set Kmpc to an empty matrix, mpcsi m will do an open-loop simulation. Then the inputs to the plant will be r (which must be set to the vector of manipulated variables in this case) and dstep.

tend

Is the desired duration of the simulation (in time units).

r

Is normally a setpoint matrix consisting of N rows and n_y columns, where n_y is the number of output variables, y:

$$\mathbf{r} = \begin{bmatrix} r_1(1) & r_2(1) & \dots & r_{n_y}(1) \\ r_1(2) & r_2(2) & \dots & r_{n_y}(2) \\ \vdots & \vdots & \dots & \vdots \\ r_1(N) & r_2(N) & \dots & r_{n_y}(N) \end{bmatrix}$$

where $r_i(k)$ is the setpoint for output j at time t = kT, and T is the sampling period (as specified in the step format of pl ant and model). If tend > NT, the setpoints vary for the first N periods in the simulation, as specified by r, and are then held constant at the values given in the last row of r for the remainder of the simulation.

In many simulations one wants the setpoints to be constant for the entire time, in which case r need only contain a single row of n_y values.

If you set $r=[\]$, the default is a row of n_y zeros.

For open-loop simulations, r specifies the manipulated variables and must contain n_{ij} columns.

The following input variables are optional. In general, setting one of them equal to an empty matrix causes mpcsi m to use the default value, which is given in the description.

usat

Is a matrix giving the limits on the manipulated variables. Its format is as follows:

$$\text{usat} = \begin{bmatrix} u_{min, 1}(1) & \dots & u_{min, n_u}(1) \\ u_{min, 1}(2) & \dots & u_{min, n_u}(2) \\ \vdots & \dots & \vdots \\ u_{min, 1}(N) & \dots & u_{min, n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} u_{max, 1}(1) & \dots & u_{max, n_u}(1) \\ u_{max, 1}(2) & \dots & u_{max, n_u}(2) \\ \vdots & \dots & \vdots \\ u_{max, 1}(N) & \dots & u_{max, n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} \Delta u_{max, 1}(1) & \dots & \Delta u_{max, n_u}(N) \\ \Delta u_{max, 1}(2) & \dots & \Delta u_{max, n_u}(2) \\ \vdots & \dots & \vdots \\ \Delta u_{max, 1}(N) & \dots & \Delta u_{max, n_u}(N) \end{bmatrix}$$

Note that it contains three matrices of Nrows. N may be different than that for the setpoint matrix, r, but the idea is the same: the saturation limits will vary for the first N sampling periods of the simulation, then be held constant at the values given in the last row of usat for the remaining periods (if any).

The first matrix specifies the *lower bounds* on the n_u manipulated variables. For example, $u_{min^ij}(k)$ is the lower bound for manipulated variable j at time t = kT in the simulation. If $u_{min^ij}(k) = -inf$, manipulated variable j will have no lower bound at t = kT.

The second matrix gives the *upper bounds* on the manipulated variables. If $u_{max,j}(k) = inf$, manipulated variable j will have no upper bound at t = kT.

The lower and upper bounds may be either positive or negative (or zero) as long as $u_{min^{-}j}(k) \le u_{max,j}(k)$.

The third matrix gives the limits on the rate of change of the manipulated variables. In other words, mpcsi m will force

 $|u_j(k) - u_j(k-1)| \le \Delta u_{max,j}(k)$. The limits on the rate of change must be nonnegative.

The default is no saturation constraints, i.e., all the u_{min} values will be set to – inf, and all the u_{max} and Δu_{max} values will be set to inf.

Note: Saturation constraints in mpcsi m are enforced by simply *clipping* the manipulated variable moves so that they satisfy all constraints. This is a nonoptimal solution that, in general, will differ from the results you would get using the ulim variable in cmpc.

tfilter

Is a matrix of time constants for the noise filter and the unmeasured disturbances entering at the plant output. The first row of n_y elements gives the noise filter time constants and the second row of n_y elements gives the time constants of the lags through which the unmeasured disturbance steps pass. If tfilter only contains one row, the unmeasured disturbances are assumed to be steps. If you set tfilter=[] or omit it, the default is no noise filtering and steplike unmeasured disturbances.

dpl ant

Is a model in MPC *step* format representing all the disturbances (measured and unmeasured) that affect pl ant in the above diagram. If dpl ant is provided, then input dstep is also required. For output step disturbances, set dpl ant=[]. The default is no disturbances.

dmodel

Is a model in MPC *step* format representing the measured disturbances. If dmodel is provided, then input dstep is also required. If there are no measured disturbances, set dmodel=[]. For output step disturbances, set dmodel=[]. If there are both measured and un-measured disturbances, set the columns of dmodel corresponding to the unmeasured disturbances to zero. The default is no measured disturbances.

dstep

Is a matrix of disturbances to the plant. For output step disturbances $(dpl\ ant=[\])$ and $dmodel=[\])$, the format is the same as for r. For disturbances through step-response models $(dpl\ ant\ only\ or\ both\ dpl\ ant\ and\ dmodel$ nonempty), the format is the same as for r, except that the number of columns is n_d rather than n_v . The default is a row of zeros.

Note: You may use a different number of rows in the matrices r, usat and dstep, should that be appropriate for your simulation.

The calculated outputs are as follows (all but yp are optional):

уp

Is a matrix containing M rows and n_y columns, where M = round(tend/delt2) + 1 and delt2 is the sampling time. The first row will contain the initial condition, and row k-1 will give the values of the plant outputs, y (see above diagram), at time t=kT.

u

Is a matrix containing the same number of rows as yp and n_u columns. The time corresponding to each row is the same as for yp. The elements in each row are the values of the manipulated variables, u (see above diagram).

ym

Is a matrix of the same structure as yp, containing the values of the predicted outputs from the state estimator in the controller. ym will, in general, differ from yp if model |p| ant and/or there are unmeasured disturbances. The prediction includes the effect of the most recent measurement, i.e., $\hat{y}(k|k)$.

Examples

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

The following statements build the model and calculate the MPC controller gain:

```
g11=poly2tfd(12.8, [16.7 1], 0, 1);

g21=poly2tfd(6.6, [10.9 1], 0, 7);

g12=poly2tfd(-18.9, [21.0 1], 0, 3);

g22=poly2tfd(-19.4, [14.4 1], 0, 3);

delt=3; ny=2; tfinal=90;

model=tfd2step(tfinal, delt, ny, g11, g21, g12, g22);

plant=model;

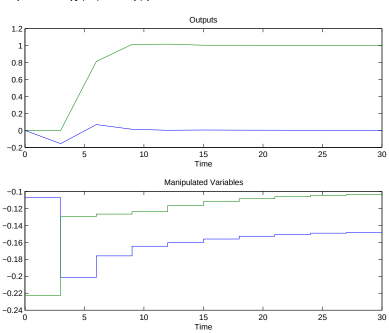
P=6; M=2;

ywt=[]; uwt=[1 1];

Kmpc=mpccon(i mod, ywt, uwt, M, P);
```

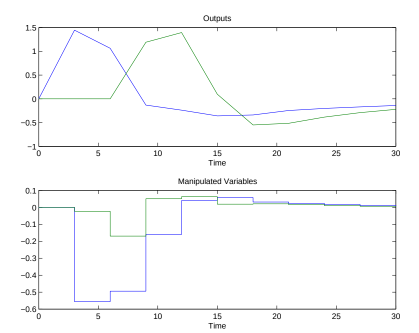
Simulate and plot the closed-loop performance for a unit step in the setpoint for y_2 , occurring at t = 0.

```
tend=30; r=[0 1];
[y, u] = mpcsim(plant, model, Kmpc, tend, r);
plotall(y, u, delt), pause
```



Try a pulse change in the disturbance that adds to u_1 :

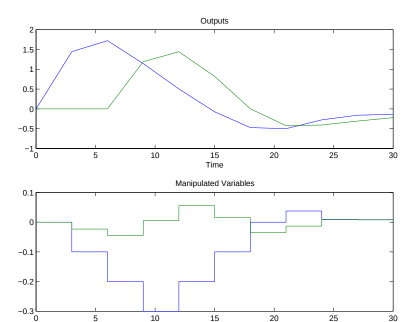
```
r=[ ]; usat=[ ]; tfilter=[ ]; dmodel=[ ];
dplant=plant;
dstep=[ 1 0; 0 0];
[y,u]=mpcsim(plant, model, Kmpc, tend, r, usat, tfilter, . . .
    dplant, dmodel, dstep);
plotall(y, u, delt), pause
```



For the same disturbance as in the previous case, limit the rates of change of both manipulated variables.

```
usat=[-inf -inf inf inf 0.1 0.05];
[y, u]=mpcsim(plant, model, Kmpc, tend, r, usat, tfilter, . . .
```

dpl ant, dmodel, dstep);
pl otall(y, u, delt), pause



Restriction

Initial conditions of zero are used for all the variables. This simulates the condition where all variables represent a deviation from a steady-state initial condition.

See Also

plotall, ploteach, cmpc, mpccl, mpccon

nlcmpc

Purpose

Model predictive controller for simulating closed-loop systems with hard bounds on manipulated variables and/or controlled variables using linear models in the MPC *step* format for *nonlinear* plants represented as Simulink S-functions.

Description

nl cmpc is a Simulink S-function block and can be invoked by typing nl mpcl i b at the MATLAB prompt. Its usage is identical to other Simulink blocks. The input to nl cmpc includes both the variables controlled by nl cmpc and measured disturbances. The first n_y elements of the input are treated as the controlled variables while the rest is taken as the measured disturbances. The output from nl cmpc are the values of the manipulated variables. Initial conditions for the manipulated variables and the measured disturbances must be specified. The controlled variables sent to nl cmpc and the manipulated variables returned by nl cmpc are actual variables; they are not deviation variables.

Because of the limit on the number of masked variables that can be specified for a Simulink block, model and dmodel are put together as "one" variable, r, ywt, and uwt as "one" variable, and ylim and ulim as "one" variable. m and p should be entered as one *row vector*. u0 and d0 should also be entered as one *row vector*. The required input variables are as follows:

model pd

Equals [model dmodel]. model is a *linear* model in the MPC *step* format that is to be used for state estimation in the controller. In general, it is a linear approximation of the nonlinear plant. dmodel is a model in MPC *step* format representing the effect of the measured disturbances. The default is no measured disturbances. Note that the truncation time for model and dmodel must be the same and the number of outputs for model and dmodel must be the same.

ryuwt

Equals [r ywt uwt]. r is a setpoint matrix consisting of N rows and n_y columns, where n_y is the number of output variables, y:

$$\mathbf{r} = \begin{bmatrix} r_1(1) & r_2(1) & \dots & r_{n_y}(1) \\ r_1(2) & r_2(2) & \dots & r_{n_y}(2) \\ \vdots & \vdots & \dots & \vdots \\ r_1(N) & r_2(N) & \dots & r_{n_y}(N) \end{bmatrix}$$

Where $r_i(k)$ is the setpoint for output i at time t = kT, and T is the sampling period (as specified in the step format of model). If the simulation time is larger than NT, the setpoints vary for the first N periods in the simulation, as specified by r, and are then held constant at the values given in the last row of r for the remainder of the simulation. In many simulations one wants the setpoints to be constant for the entire time, in which case r need only contain a single row of n_v values.

ywt

Must have n_y columns, where n_y is the number of outputs. All weights must be ≥ 0 .

You may vary the weights at each step in the prediction horizon by including up to P rows in ywt. Then the first row of n_y values applies to the tracking errors in the first step in the prediction horizon, the next row applies to the next step, etc. See mpccon for details on the form of the optimization objective function.

If you supply only nrow rows, where $1 \le nrow < P$, nl cmpc will use the last row to fill in any remaining steps. Thus if you wish the weighting to be the same for all P steps, you need only specify a single row.

uwt

Has the same format as ywt, except that uwt applies to the changes in the manipulated variables. If $uwt + [\quad]$, it must have n_u columns, where n_u is the number of manipulated variables.

Notice that the number of rows for r, ywt, and uwt should be the same. If not, one can enter the variable as parpart (r, ywt, uwt). The function parpart

appends extra rows to r, ywt, and/or uwt so that they have the same number of rows. The default is r=y0, where y0 is the initial condition for the output, equal (unity) weighting of all outputs over the entire prediction horizon and zero weighting of all input.

mp

Equals [M P]. P equals the last element of MP. There are two ways to specify M: If it is a scalar, nl cmpc interprets it as the input horizon (number of moves) as in DMC; if it is a row vector containing n_b elements, each element of the vector indicates number of the steps over which $\Delta u(k) = 0$ during the optimization and nl cmpc interprets it as a set of n_b blocking factors. There may be $1 \le n_b \le P$ blocking factors, and their sum must be $1 \le P$. If you set M=[1], the default is M=P, which is equivalent to M=ones(1, P). P is the number of sampling periods in the prediction horizon.

yul i m

Equals [ylimulim]. ulimis a matrix giving the limits on the manipulated variables. Its format is as follows:

$$\begin{aligned} & \text{ulim} = & \begin{bmatrix} u_{min,\,1}(1) & \dots & u_{min,\,n_u}(1) \\ u_{min,\,1}(2) & \dots & u_{min,\,n_u}(2) \\ \vdots & \dots & \vdots \\ u_{min,\,1}(N) & \dots & u_{min,\,n_u}(N) \end{bmatrix} \\ & \begin{bmatrix} u_{max,\,1}(1) & \dots & u_{max,\,n_u}(1) \\ u_{max,\,1}(2) & \dots & u_{max,\,n_u}(2) \\ \vdots & \dots & \vdots \\ u_{max,\,1}(N) & \dots & u_{max,\,n_u}(N) \end{bmatrix} \\ & \begin{bmatrix} \Delta u_{max,\,1}(1) & \dots & \Delta u_{max,\,n_u}(1) \\ \Delta u_{max,\,1}(2) & \dots & \Delta u_{max,\,n_u}(2) \\ \vdots & \dots & \vdots \\ \Delta u_{max,\,1}(N) & \dots & \Delta u_{max,\,n_u}(N) \end{bmatrix} \end{aligned}$$

Note that it contains three matrices of N rows. In this case, the limits on N are $1 \le N \le n_b$, where n_b is the number of times the manipulated variables are to change over the input horizon. If you supply fewer than n_b rows, the last row is repeated automatically.

The first matrix specifies the *lower bounds* on the n_u manipulated variables. For example, $u_{min'j}(2)$ is the lower bound for manipulated variable j for the second move of the manipulated variables (where the first move is at the start of the prediction horizon). If $u_{min'j}(k) = -inf$, manipulated variable j will have no lower bound for that move.

The second matrix gives the *upper bounds* on the manipulated variables. If $u_{max,j}(k) = inf$, manipulated variable j will have no upper bound for that move.

The lower and upper bounds may be either positive or negative (or zero) as long as $u_{min,j}(k) \le u_{max,j}(k)$.

The third matrix gives the limits on the rate of change of the manipulated variables. In other words, cmpc will force $|u_j(k) - u_j(k-1)| \le \Delta u_{max,j}(k)$. The limits on the rate of change must be nonnegative and *finite*. If you want it to be unbounded, set the bound to a large number (but not too large — a value of 10^6 should work well in most cases).

 $yl\,i\,m$ has the same format as $ul\,i\,m$, but for the lower and upper bounds of the outputs. The first row applies to the first point in the prediction horizon.

Note that the number of rows for ylim and ulim should be the same. If the number of rows for ylim and ulim differs, one can use parpart (ylim, ulim). The function parpart appends extra rows to ylim or ulim so that they have the same number of rows. If you set yulim = [], then $u_{min} = -inf$, $u_{max} = inf$, $\Delta u_{max} = 10^6$, $y_{min} = -inf$ and $y_{max} = inf$.

tfilter

Is a matrix of time constants for the noise filter and the unmeasured disturbances entering at the plant output. The first row of n_y elements gives the noise filter time constants and the second row of n_y elements gives the time constants of the lags through which the unmeasured disturbance steps pass. If tfilter only contains one row, the unmeasured disturbances are assumed to be steps. If you set tfilter= [], no noise filtering and steplike unmeasured disturbances are assumed.

nlcmpc

ud0

Equals [u0 d0]. u0 are initial values of the manipulated variables arranged in a *row vector* having n_u elements; n_u is the number of the manipulated variables computed by nl cmpc. d0 are initial values of the measured disturbances arranged in a *row vector* having n_d elements; n_d is the number of the measured disturbances. The default is u0 = 0 and d0 = 0.

Notes

- Initial conditions for the manipulated variables that are calculated by nl cmpc are specified through nl cmpc while initial conditions for the controlled variables are specified through the S-function for the nonlinear plant.
- You may use a different number of rows in the matrices r, ulim and ylim, should that be appropriate for your simulation.
- The ulim constraints used here are fundamentally different from the usat constraints used in the nlmpcsimblock. The ulim constraints are defined relative to the beginning of the prediction horizon, which moves as the simulation progresses. Thus at each sampling period, k, the ulim constraints apply to a block of calculated moves that begin at sampling period k and extend for the duration of the input horizon. The usat constraints, on the other hand, are relative to the fixed point t = 0, the start of the simulation.
- For unconstrained problems, nl cmpc and nl mpcsi m should give the same results. The latter will be faster because it uses an analytical solution of the QP problem, whereas nl cmpc solves it by iteration.

Example

See the examples for nl mpcsi m with one modification: replace the block nl mpcsi m with nl cmpc. Clearly, additional variables should be defined appropriately.

See Also

cmpc, nl mpcsi m

Purpose

Model predictive controller for simulating closed-loop systems with *saturation constraints* on the manipulated variables using linear models in the MPC *step* format for nonlinear plants represented as Simulink S-functions.

Description

nl mpcsi m is a Simulink S-function block and can be invoked by typing nl mpcl i b at the MATLAB prompt. Its usage is identical to other Simulink blocks. The input to nl mpcsi m includes both the variables controlled by nl mpcsi m and measured disturbances. The first n_y elements of the input are treated as the controlled variables while the rest is taken as the measured disturbances. The output from nl mpcsi m are the values of the manipulated variables. Initial conditions for the manipulated variables and the measured disturbances must be specified. Both the controlled variables sent to nl mpcsi m and the manipulated variables returned by nl mpcsi m are the actual variables; they are not deviation variables.

Because of the limit on the number of masked variables that can be specified for a Simulink block, model and dmodel are put together as one variable. u0 and d0 should be entered as one *row vector*. The required input variables are as follows:

model pd

Equals [model dmodel]. model is a linear model in the MPC *step* format that is to be used for state estimation in the controller. In general, it is a linear approximation for the nonlinear plant. Note, however, that model should be the same as that used to calculate Kmpc. dmodel is a model in MPC *step* format representing the measured disturbances. If dmodel = [], the default is no measured disturbances. Note that the truncation time for model and dmodel should be the same and the number of outputs for model and dmodel should be the same.

Kmpc

Is the MPC controller gain matrix, usually calculated using the function mpccon.

nImpcsim

 \mathbf{r}

Is a setpoint matrix consisting of N rows and n_y columns, where n_y is the number of controlled variables, y:

$$\mathbf{r} = \begin{bmatrix} r_1(1) & r_2(1) & \dots & r_{n_y}(1) \\ r_1(2) & r_2(2) & \dots & r_{n_y}(2) \\ \vdots & \vdots & \dots & \vdots \\ r_1(N) & r_2(N) & \dots & r_{n_y}(N) \end{bmatrix}$$

Where $r_i(k)$ is the setpoint for output i at time t = kT, and T is the sampling period (as specified in the step format of model). If the simulation time is larger than NT, the setpoints vary for the first N periods in the simulation, as specified by r, and are then held constant at the values given in the last row of r for the remainder of the simulation.

In many simulations one wants the setpoints to be constant for the entire time, in which case r need only contain a single row of n_v values.

Note that r is the actual setpoint. If you set $r=[\]$, the default is y0.

usat

Is a matrix giving the saturation limits on the manipulated variables. Its format is as follows:

$$usat = \begin{bmatrix} u_{min, 1}(1) & \dots & u_{min, n_u}(1) \\ u_{min, 1}(2) & \dots & u_{min, n_u}(2) \\ \vdots & \dots & \vdots \\ u_{min, 1}(N) & \dots & u_{min, n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} u_{max, 1}(1) & \dots & u_{max, n_u}(1) \\ u_{max, 1}(2) & \dots & u_{max, n_u}(2) \\ \vdots & \dots & \vdots \\ u_{max, 1}(N) & \dots & u_{max, n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} \Delta u_{max, 1}(1) & \dots & \Delta u_{max, n_u}(1) \\ \Delta u_{max, 1}(2) & \dots & \Delta u_{max, n_u}(2) \\ \vdots & \dots & \vdots \\ \Delta u_{max, 1}(N) & \dots & \Delta u_{max, n_u}(N) \end{bmatrix}$$

Note that it contains three matrices of Nrows. Nmay be different from that for the setpoint matrix, r, but the idea is the same: the saturation limits will vary for the first Nsampling periods of the simulation, then be held constant at the values given in the last row of usat for the remaining periods (if any).

The first matrix specifies the *lower bounds* on the n_u manipulated variables. For example, $u_{min'j}(k)$ is the lower bound for manipulated variable j at time t = kT in the simulation. If $u_{min'j}(k) = -inf$, manipulated variable j will have no lower bound at t = kT.

The second matrix gives the *upper bounds* on the manipulated variables. If $u_{max,j}(k) = inf$, manipulated variable j will have no upper bound at t = kT.

The lower and upper bounds may be either positive or negative (or zero) as long as $u_{min'j}(k) \le u_{max,j}(k)$.

The third matrix gives the limits on the rate of change of the manipulated variables. In other words, mpcsi m will force $|u_j(k) - u_j(k-1)| \le \Delta u_{max,j}(k)$. The limits on the rate of change must be nonnegative.

If usat = [], then all the u_{min} values will be set to -inf, and all the u_{max} and u_{max} values will be set to inf.

Note: Saturation constraints are enforced by simply *clipping* the manipulated variable moves so that they satisfy all constraints. This is a nonoptimal solution that, in general, will differ from the results you would get using the ulim variable in cmpc or nl cmpc.

tfilter

Is a matrix of time constants for the noise filter and the unmeasured disturbances entering at the plant output. The first row of n_y elements gives the noise filter time constants and the second row of n_y elements gives the time constants of the lags through which the unmeasured disturbance steps pass. If tfilter only contains one row, the unmeasured disturbances are assumed to be steps. If you set tfilter= [], no noise filtering and steplike unmeasured disturbances are assumed.

ud0

Equals [u0 d0]. u0 are initial values of the manipulated variables arranged in a *row vector* having n_u elements; n_u is the number of the manipulated variables computed by nl mpcsi m. d0 are initial values of the measured disturbances arranged in a *row vector* having n_d elements; n_d is the number of the measured disturbances. The default is u0 = 0 and d0 = 0.

Note: You may use a different number of rows in the matrices r and usat, should that be appropriate for your simulation.

Examples

Let us now demonstrate the use of the controller nl mpcsi m. Since the plant used in Example 1 is linear, using mpcsi m would be much faster. The point, however, is to show how masked variables are specified for nl mpcsi m.

1 The plant is linear with two inputs and two outputs. It is represented by

$$\frac{dx}{dt} = \begin{bmatrix} -1.2 & 0 \\ 0 & -\frac{1}{1.5} \end{bmatrix} x + \begin{bmatrix} 0.2 \\ 1 \end{bmatrix} u + \begin{bmatrix} 50 \\ 0 \end{bmatrix}$$
$$y = x$$

The Simulink S-function for this plant is in mpcpl ant. m. The nominal steady-state operating condition is $y0 = [58.3\ 1.5]$ and $u0 = [100\ 1]$. The Simulink block to simulate this plant using nl mpcsi m is in nl mpcdm1. m and shown in Figure 1-1.

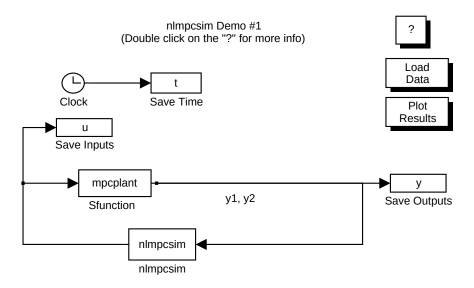


Figure 1-1 Simulink Block for Example 1

The following statements build the step response model and specify the parameter values. Note that model does not equal the plant model stored in

mpcpl ant. m. The important thing to notice is that both r and usat are actual variables. They are not deviation variables.

```
g11=poly(0.4, [1 2]);
g21 = poly2tfd(0, 1);
g12 = poly2tfd(0, 1);
g22=poly2tfd(1, [1 1]);
tfinal = 8;
del t=0.2:
nout=2;
model =tfd2step(tfinal, delt, nout, g11, g21, g12, g22);
ywt = [1 \ 1];
uwt = [0 \ 0];
M=4:
P=10:
r=[68.32];
usat=[100 1 200 3 200 200];
tfilter=[];
Kmpc = mpccon(model, ywt, uwt, M, P);
dmodel = [];
```

There are two ways to simulate the closed loop system. We can set the simulation parameters and click on **Start** under **Simulation** or via the following statements.

```
plant='nlmpcdm1'; y0=[58.3 1.5];
u0=[100 1];
tfsim = 2;
tol=[1e-3];
minstep=[];
maxstep=[];
[t,yu]=gear(plant,tfsim,[y0 u0],[tol,minstep,maxstep]);
```

Figure 1-2 shows the response for the setpoint change.

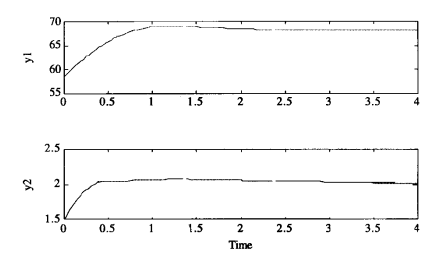


Figure 1-2 Output responses for a setpoint change for Example 1

2 The plant is the paper machine headbox discussed in the section, "Application: Paper Machine Headbox Control" in Chapter 3. The nonlinear plant model is represented as a Simulink S-function and is in pap_mach. m. The plant has two inputs, three outputs, four states, one measured disturbance, and one unmeasured disturbance. All these variables are zero at the nominal steady-state. Since the model for nl mpcsi m must be linear, we linearize the nonlinear plant at the nominal steady-state to obtain a linear model. Since the model is simple, we can linearize it analytically to obtain A, B, C, and D.

The Simulink block to simulate this nonlinear plant using nl mpcsi m is in nl mpcdm2. m and shown in Figure 1-3.

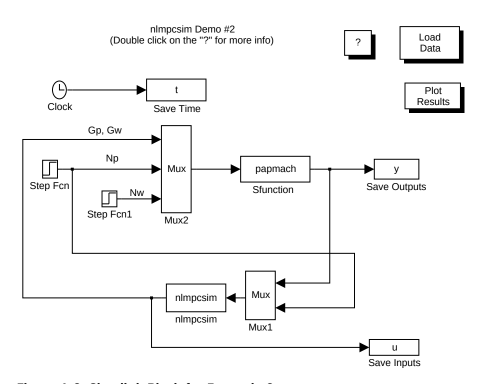


Figure 1-3 Simulink Block for Example 2

The following statements build the step response model and specify the parameter values.

```
[model,dmodel] = mod2step(i mod,20);\\ m=5;\\ p=20;\\ ywt=[1\ 0\ 5]; \ \% \ unequal\ weighting\ of\ y1\ and\ y3,\ no\ control \% \ of\ y2\\ uwt=[1\ 1]; \ \% \ Equal\ weighting\ of\ u1\ and\ u2\\ ulim=[-10\ -10\ 10\ 10\ 2\ 2]; \ \% \ Constraints\ on\ u\\ ylim=[\ ]; \ \% \ No\ constraints\ on\ y\\ usat=ulim;\\ tfilter=[\ ];\\ y0=[0\ 0\ 0];\\ u0=[0\ 0];\\ r=[0\ 0\ 0];\\ Kmpc=mpccon(model,ywt,uwt,M,P);
```

Figure 1-4 shows the output responses for a unit-step measured disturbance Np = 1 and a step unmeasured disturbance with Nw = 5.

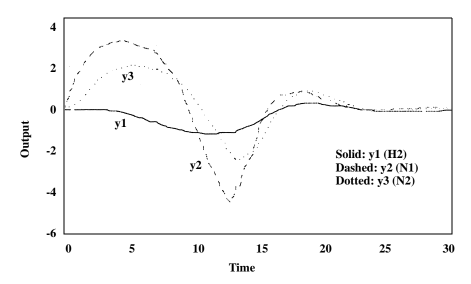


Figure 1-4 Output responses for a unit-step measured disturbance Np=1 and a step unmeasured disturbance Nw=5

See Also mpcsi m, nl cmpc

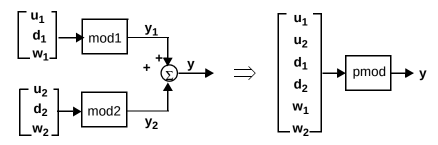
Purpose

Puts two models in parallel by connecting their outputs. Mimics the utility function <code>mpcparal</code>, except that <code>paramod</code> works on models in the MPC <code>mod</code> format.

Syntax

pmod = paramod(mod1, mod2)

Description



mod1 and mod2 are models in the MPC mod format (see mod in the online MATLAB Function Reference format section for a detailed description). You would normally create them using either the tfd2mod, ss2mod or th2mod functions.

paramod combines them to form a composite system, pmod, as shown in the above diagram. It is also in the \mathbf{mod} format. Note how the inputs to $\mathrm{mod}1$ and $\mathrm{mod}2$ are ordered in pmod .

Restriction

mod1 and mod2 must have been created with equal sampling periods and they must have the same number of measured and unmeasured outputs.

See Also

addmd, addmod, addumd, appmod, sermod

Purpose

Plots outputs and manipulated variables from a simulation, all on one "page."

Syntax

plotall(y, u) plotall(y, u, t)

Description

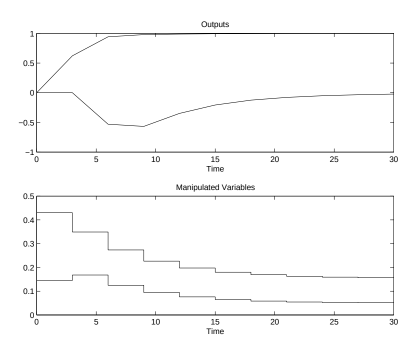
Input variables y and u are matrices of outputs and manipulated variables, respectively. Each row represents a sample at a particular time. Each column shows how a particular output (or manipulated) variable changes with time.

Input variable t is optional. If you supply it as a scalar, plotall interprets is as the sampling period, and calculates the time axis for the plots accordingly. It can also be a column vector, in which case it must have the same number of rows as y and u and is interpreted as the times at which the samples of y and u were taken. If you do not supply t, plotall uses a sampling period of 1 by default.

pl ot all 1 plots all the outputs on a single graph. If there are multiple outputs that have very different numerical scales, this may be unsatisfactory. In that case, use pl ot each.

pl otal l plots all the manipulated variables in "stairstep" form (i.e., assuming a zero-order hold) on a single graph. Again, pl ot each may be the better choice if scales are very different.

Example output: (mpccon example)



See Also

ploteach, plotstep, plotfrsp

Purpose

Plots outputs and manipulated variables from a simulation on separate graphs, up to four per page.

Syntax

```
pl oteach(y)
pl oteach(y, u)
pl oteach([ ], u)
pl oteach(y, [ ], t)
pl oteach([ ], u, t)
pl oteach(y, u, t)
```

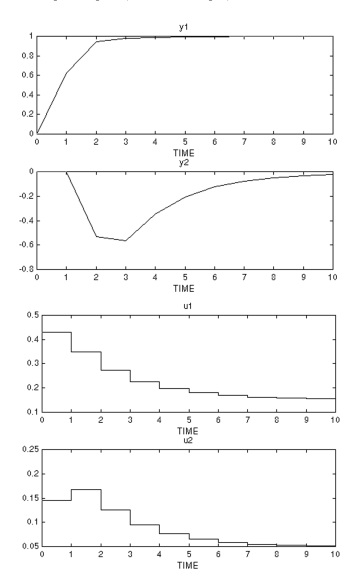
Description

Input variables y and u are matrices of outputs and manipulated variables, respectively. Each row represents a sample at a particular time. Each column shows how a particular output (or manipulated) variable changes with time. As shown above, you may supply both y and u, or omit either one of them.

Input variable t is optional. If you supply it as a scalar, ploteach interprets is as the *sampling period*, and calculates the time axis for the plots accordingly. It can also be a column vector, in which case it must have the same number of rows as y and u and is interpreted as the times at which the samples of y and u were taken. If you do not supply t, pl ot each uses a sampling period of 1 by default.

pl ot each plots the manipulated variables in "stairstep" form (i.e., assuming a zero-order hold).

Example output: (mpccon example)



See Also

plotall, plotfrsp, plotstep

Plots the frequency response generated by mod2frsp as a Bode plot.

Syntax

```
plotfrsp(vmat)
plotfrsp(vmat, out, i n)
```

Description

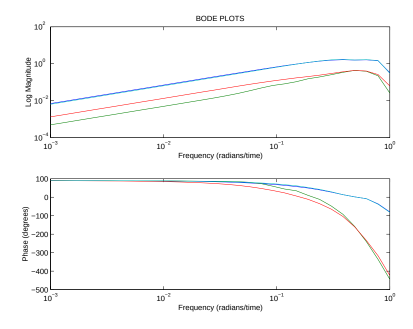
vmat is a *varying* matrix which contains the data to be plotted.

Let $F(\omega)$ denote the matrix (whose entries are functions of the independent variable ω) whose sampled values $F(\omega_1), \ldots, F(\omega_N)$ are contained in vmat.

pl of frsp(vmat) will generate Bode plots of all elements of $F(\omega)$.

Optional inputs out and in are *row vectors* which specify the row and column indices respectively of a submatrix of $F(\omega)$. plotfrsp will then generate Bode plots of the elements of the specified submatrix of $F(\omega)$.

Example Output: (mod2frsp example)



See Also

mod2frsp, varying format

plotstep

Purpose

Plots multiple step responses as calculated by mod2step, ss2step or tfd2step.

Syntax

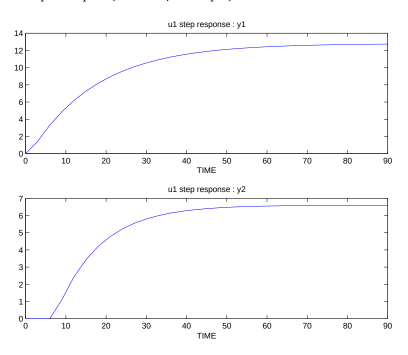
pl otstep(pl ant)
pl otstep(pl ant, opt)

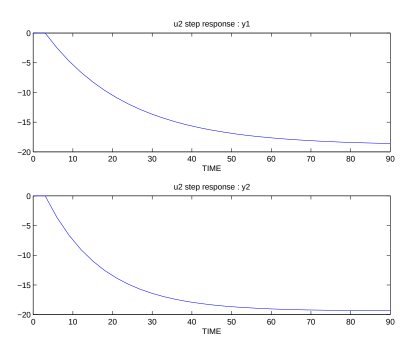
Description

pl ant is a step-response matrix in the MPC step format created by mod2step, ss2step or tfd2step.

opt is an optional scalar or *row vector* that allows you to select the outputs to be plotted. If you omit opt, plotstep plots every output. If, for example, plant contains results for 6 outputs, setting opt=[1, 4, 5] would cause only y_1 , y_4 and y_5 to be plotted.

Example output: (tfd2step example)





See Also i mp2step, mod2step, step format, plotall, ploteach, plotfrsp, ss2step, tfd2step

Determine the impulse response coefficients for a multi-input single-output system via Partial Least Squares (PLS).

Syntax

```
[theta, w, cw, ssqdif, yres] = plsr(xreg, yreg, ni nput, lv)
[theta, w, cw, ssqdif, yres] = plsr(xreg, yreg, ni nput, lv, plotopt)
```

Description

Given a set of regression data, xreg and yreg, the impulse response coefficient matrix, theta, is determined via PLS. Column i of theta corresponds to the impulse response coefficients for input i. Only a single output is allowed. The number of inputs, ni nput, and the number of latent variables, l v, must be specified.

Optional output w is a matrix of dimension n (number of impulse response coefficients) by $l\ v$ consisting of orthogonal column vectors maximizing the cross variance between input and output. Column vector cw (optional) contains the coefficients associated with each orthogonal vector for calculating theta (theta=w*cw).

Optional output ssqdi f is an l v-by-2 matrix containing the percent variances captured by PLS. The first column contains information for the input; the second column for the output. Row i of ssqdi f gives a measure of the variance captured by using the first i latent variables.

The output residual or prediction error (yres) is also returned (optional).

No plot is produced if plotopt is equal to 0, which is the default; a plot of the actual output and the predicted output is produced if plotopt=1; two plots — plot of actual and predicted output, and plot of output residual — are produced for plotopt=2.

Example

Consider the following two-input single-output system:

$$y(s) = \left[\frac{5.72e^{-14s}}{60s+1} \frac{1.52e^{-15s}}{25s+1}\right] \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

Load the input and output data. The input and output data were generated from the above transfer function and random zero-mean noise was added to the output. Sampling time of 7 minutes was used.

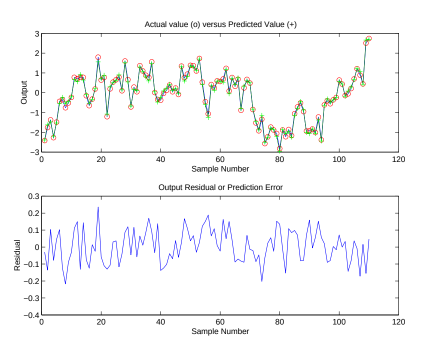
load plsrdat;

Put the input and output data in a form such that they can be used to determine the impulse response coefficients. 30 impulse response coefficients (n) are used.

```
n = 30;
[xreg, yreg] = wrtreg(x, y, n);
```

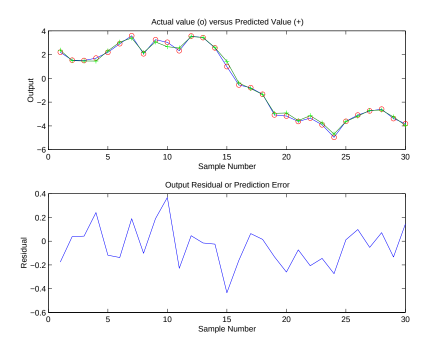
Determine the impulse response coefficients via pl sr using 10 latent variables. By specifying pl ot opt=2, two plots — plot of predicted output and actual output, and plot of the output residual (or predicted error) — are produced.

```
ni nput = 2;
lv = 10;
pl otopt = 2;
theta = pl sr(xreg, yreg, ni nput, lv, pl otopt);
```



Use a new set of data to validate the impulse model.

```
[newxreg, newyreg] = wrtreg(newx, newy, n);
yres = validmod(newxreg, newyreg, theta, plotopt);
```

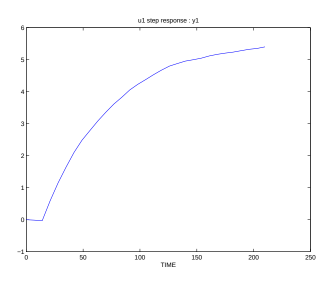


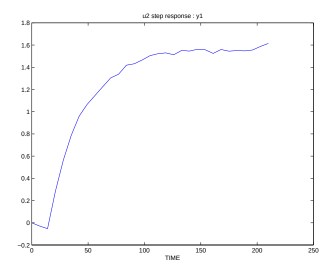
Convert the impulse model to a step model to be used in MPC design. Sampling time of 7 minutes was used in determining the impulse model. Number of outputs (1 in this case) must be specified.

```
nout = 1;
delt = 7;
model = imp2step(delt, nout, theta);
```

Plot the step response coefficients.

plotstep(model)





See Also

mlr, validmod, wrtreg

poly2tfd, poly format

Purpose

pol y2tfd converts a transfer function (continuous or discrete) from the standard MATLAB poly format into the MPC tf format.

Syntax

Description

Consider a continuous-time (Laplace domain) transfer function such as

$$G(s) = \frac{b_0 s^n + b_1 s^{n-1} + \dots + b_n}{a_0 s^n + a_1 s^{n-1} + \dots + a_n}$$

or a discrete-time transfer function such as

$$G(z) = \frac{b_0 + b_1 z^{-1} + \dots + b_n z^{-n}}{a_0 + a_1 z^{-1} + \dots + a_n z^{-n}}$$

where *z* is the forward-shift operator. Using the MATLAB *poly* format, you would represent either of these as a numerator polynomial and a denominator polynomial, giving the coefficients of the highest-order terms first:

If the numerator contains leading zeros, they may be omitted, i.e., the number of elements in num can be \leq the number of elements in den.

pol y2tfd uses num and den as input to build a transfer function, *g*, in the MPC *tf* format (see tf section for details). Optional variables you can include are:

del t

The sampling period. If this is zero or you omit it, poly2tfd assumes that you are supplying a continuous-time transfer function. If you are supplying a discrete-time transfer function you must specify del t. Otherwise g will be misinterpreted when you use it later in the MPC Toolbox functions.

del ay

The time delay. For a continuous-time transfer function, del ay should be in time units. For a discrete-time transfer function, del ay should be specified as

the integer number of sampling periods of time delay. If you omit it, poly2tfd assumes a delay of zero.

Examples

Consider the continuous-time transfer function:

$$G(s) = 0.5 \frac{3s-1}{5s^2 + 2s + 1}$$

It has no delay. The following command creates the MPC *tf* format:

Now suppose there were a delay of 2.5 time units:

G(s) =
$$0.5 \frac{3s-1}{5s^2+2s+1}e^{-2.5s}$$
 . You could use:

Next let's get the equivalent transfer function in discrete form. An easy way is to get the correct poly form using cp2dp, then use pol y2tfd to get it in the *tf* form. Here are the commands to do it using a sampling period of 0.75 time units:

```
delt=0.75; [numd, dend]=cp2dp(0.5*[3 -1], [5 2 1], delt, rem(2.5, delt)); g=poly2tfd(numd, dend, delt, fix(2.5/delt));
```

Note that cp2dp is used to handle the fractional time delay and the integer number of sampling periods of time delay is an input to pol y2tfd. The results are:

See Also

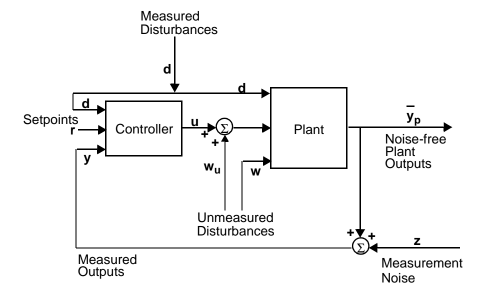
cp2dp, tf, th2mod, tfd2step

Simulates closed-loop systems with hard bounds on manipulated variables and/or outputs using models in the MPC **mod** format. Solves the MPC optimization problem by quadratic programming.

Syntax

```
yp = scmpc(pmod, i mod, ywt, uwt, M, P, tend, r)
[yp, u, ym] = scmpc(pmod, i mod, ywt, uwt, M, P, tend, ...
r, ulim, ylim, Kest, z, d, w, wu)
```

Description



scmpc simulates the performance of the type of system shown in the above diagram when there are bounds on the manipulated variables and/or outputs.

The required input variables are as follows:

pmod

Is a model in the MPC **mod** format that is to represent the plant.

i mod

Is a model in the MPC **mod** format that is to be used for state estimation in the controller. In general, it can be different from pmod if you want to simulate the effect of plant/controller model mismatch.

ywt

Is a matrix of weights that will be applied to the setpoint tracking errors (optional). If $ywt=[\]$, the default is equal (unity) weighting of all outputs over the entire prediction horizon. If $ywt+[\]$, it must have n_y columns, where n_y is the number of outputs. All weights must be ≥ 0 .

You may vary the weights at each step in the prediction horizon by including up to P rows in ywt. Then the first row of n_y values applies to the tracking errors in the first step in the prediction horizon, the next row applies to the next step, etc. See smpccon for details on the form of the optimization objective function.

If you supply only nrow rows, where $1 \le nrow < P$, scmpc will use the last row to fill in any remaining steps. Thus if you wish the weighting to be the same for all P steps, you need only specify a single row.

uwt

Is as for ywt, except that uwt applies to the *changes* in the manipulated variables. If you use uwt=[], the default is zero weighting. If uwt [], it must have n_u columns, where n_u is the number of manipulated variables.

M

There are two ways to specify this variable:

If it is a *scalar*, scmpc interprets it as the input horizon (number of moves) as in DMC.

If it is a *row vector* containing n_b elements, each element of the vector indicates the number of steps over which $\Delta u = 0$ during the optimization and scmpc interprets it as a set of n_b blocking factors. There may be $1 \le n_b \le P$ blocking factors, and their sum must be $\le P$.

If you set $M=[\]$, the default is M=P, which is equivalent to M=ones(1,P).

P

The number of sampling periods in the prediction horizon.

tend

Is the desired duration of the simulation (in time units).

 \mathbf{r}

Is a setpoint matrix consisting of N rows and n_y columns, where n_y is the number of output variables, y:

$$\mathbf{r} = \begin{bmatrix} r_1(1) & r_2(1) & \dots & r_{n_y}(1) \\ r_1(2) & r_2(2) & \dots & r_{n_y}(2) \\ \vdots & \vdots & \dots & \vdots \\ r_1(N) & r_2(N) & \dots & r_{n_y}(N) \end{bmatrix}$$

where $r_i(k)$ is the setpoint for output j at time t = kT, and T is the sampling period (as specified by the minfo vector in the **mod** format of pmod and i mod). If tend > NT, the setpoints vary for the first N periods in the simulation, as specified by r, and are then held constant at the values given in the last row of r for the remainder of the simulation.

In many simulations one wants the setpoints to be constant for the entire time, in which case r need only contain a single row of n_V values.

If you set $r=[\]$, the default is a row of n_v zeros.

The following input variables are optional. In general, setting one of them equal to an empty matrix causes scmpc to use the default value, which is given in the description.

ul i m

Is a matrix giving the limits on the manipulated variables. Its format is as follows:

$$\text{ulim} = \begin{bmatrix} u_{min,1}(1) & \dots & u_{min,n_u}(1) \\ u_{min,1}(2) & \dots & u_{min,n_u}(2) \\ \vdots & \dots & \vdots \\ u_{min,1}(N) & \dots & u_{min,n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} u_{max,1}(1) & \dots & u_{max,n_u}(1) \\ u_{max,1}(2) & \dots & u_{max,n_u}(2) \\ \vdots & \dots & \vdots \\ u_{max,1}(N) & \dots & u_{max,n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} \Delta u_{max,1}(1) & \dots & \Delta u_{max,n_u}(1) \\ \Delta u_{max,1}(2) & \dots & \Delta u_{max,n_u}(2) \\ \vdots & \dots & \vdots \\ \Delta u_{max,1}(N) & \dots & \Delta u_{max,n_u}(N) \end{bmatrix}$$
 Note that it contains these vectors as of Newey, In this

Note that it contains three matrices of N rows. In this case, the limits on N are $1 \le N \le n_b$, where n_b is the number of times the manipulated variables are to change over the input horizon. If you supply fewer than n_b rows, the last row is repeated automatically.

The first matrix specifies the *lower bounds* on the n_u manipulated variables. For example, $u_{min'j}(2)$ is the lower bound for manipulated variable j for the second move of the manipulated variables (where the first move is at the start of the prediction horizon). If $u_{min'j}(k) = -inf$, manipulated variable j will have no lower bound for that move.

The second matrix gives the *upper bounds* on the manipulated variables. If $u_{max,j}(k) = inf$, manipulated variable j will have no upper bound for that move.

The lower and upper bounds may be either positive or negative (or zero) as long as $u_{min;j}(k) \le u_{max,j}(k)$.

The third matrix gives the limits on the rate of change of the manipulated variables. In other words, cmpc will force $|u_j(k) - u_j(k-1)| \le \Delta u_{max,j}(k)$. The limits on the rate of change must be nonnegative and *finite*. If you want it to be unbounded, set the bound to a large number (but not too large — a value of 10^6 should work well in most cases).

The default is $u_{min} = -inf$, $u_{max} = inf$ and $\Delta u_{max} = 10^6$

ylim

Same format as for ul i m, but for the lower and upper bounds of the outputs. The first row applies to the first point in the prediction horizon. The default is $y_{min} = -inf$, and $y_{max} = inf$.

Kest

Is the estimator gain matrix. The default is the DMC estimator. See smpcest for more details.

\mathbf{z}

Is measurement noise that will be added to the outputs (see above diagram). The format is the same as for r. The default is a row of n_v zeros.

А

Is a matrix of measured disturbances (see above diagram). The format is the same as for r, except that the number of columns is n_d rather than n_y The default is a row of n_d zeros.s

w

Is a matrix of unmeasured disturbances (see above diagram). The format is the same as for r, except that the number of columns is n_w rather than n_y The default is a row of n_w zeros.

wu

Is a matrix of unmeasured disturbances that are added to the manipulated variables (see above diagram). The format is the same as for r, except that the number of columns is n_u rather than n_y The default is a row of n_u zeros.

Notes

• You may use a different number of rows in the matrices r, z, d, w and wu, should that be appropriate for your simulation.

• The ulim constraints used here are fundamentally different from the usat constraints used in the smpcsim function. The ulim constraints are defined relative to the beginning of the prediction horizon, which moves as the simulation progresses. Thus at each sampling period, k, the ulim constraints apply to a block of calculated moves that begin at sampling period k and extend for the duration of the input horizon. The usat constraints, on the other hand, are relative to the fixed point t = 0, the start of the simulation.

The calculated outputs are as follows (all but yp are optional):

yp

Is a matrix containing M rows and n_y columns, where M = max(fix(tend/T) + 1, 2). The first row will contain the initial condition, and row k-1 will give the values of the noise-free plant outputs, \bar{y}_p (see above diagram), at time t = kT.

u

Is a matrix containing the same number of rows as yp and n_u columns. The time corresponding to each row is the same as for yp. The elements in each row are the values of the manipulated variables, u (see above diagram).

Note The u values are those coming from the controller before the addition of the unmeasured disturbance, w_{u} .

ym

Is a matrix of the same structure as yp, containing the values of the predicted output from the state estimator in the controller. These will, in general, differ from those in yp if i mod | pmod and/or there are unmeasured disturbances. The prediction includes the effect of the most recent measurement, i.e., it is $\hat{y}(k|k)$.

For unconstrained problems, scmpc and smpcsi m should give the same results. The latter will be faster because it uses an analytical solution of the QP problem, whereas scmpc solves it by iteration.

Examples

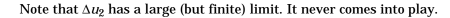
Consider the linear system:

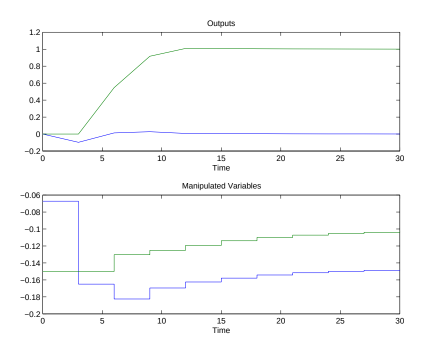
$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

The following statements build the model and set up the controller in the same way as in the smpcsi m example.

Here, however, we will demonstrate the effect of constraints. First we set a limit of 0.1 on the rate of change of u_1 and a minimum of -0.15 for u_2 .

```
ulim=[-inf -0.15 inf inf 0.1 100];
ylim=[ ];
[y,u]=scmpc(pmod,imod,ywt,uwt,M,P,tend,r,ulim,ylim);
plotall(y,u,delt), pause
```

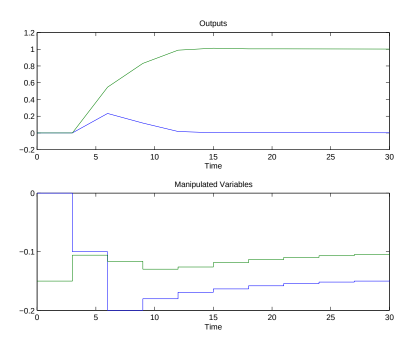




We next apply a lower bound of zero to both outputs:

```
\label{eq:ulim=[-inf-0.15 inf inf 0.1 100];} $$ylim=[0\ 0\ inf\ inf];$$ [y,u]=scmpc(pmod,imod,ywt,uwt,M,P,tend,r,ulim,ylim);$$ plotall(y,u,delt), pause
```

The following results show that no constraints are violated.



Restrictions

- Initial conditions of zero are used for all states in i mod and pmod. This simulates the condition where all variables represent a deviation from a steady-state initial condition.
- The first $n_u + n_d$ columns of the D matrices in i mod and pmod must be zero. In other words, neither u nor d may have an immediate effect on the outputs.

Suggestions

Problems with many inequality constraints can be very time consuming. You can minimize the number of constraints by:

- $\bullet\,$ Using small values for P and/or M
- Leaving variables unconstrained (limits at $\pm inf$) intermittently unless you think the constraint is important.

See Also

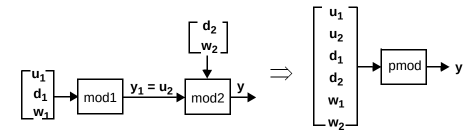
plotall, ploteach, smpccl, smpccon, smpcest, smpcsi ${\tt m}$

Puts two models in series by connecting the output of one to the input of the other. Mimics the series function of the Control System Toolbox, except that sermod works on models in the MPC **mod** format.

Syntax

pmod = sermod(mod1, mod2)

Description



mod1 and mod2 are models in the MPC mod format (see mod in the online MATLAB Function Reference for a detailed description). You would normally create them using either the tfd2mod, ss2mod or th2mod functions.

sermod combines them to form a composite system, pmod, as shown in the above diagram. It is also in the **mod** format. Note how the inputs to mod1 and mod2 are ordered in pmod.

Restrictions

- mod1 and mod2 must have been created with equal sampling periods.
- The number of *measured* outputs in mod1 must equal the number of manipulated variables in mod2.

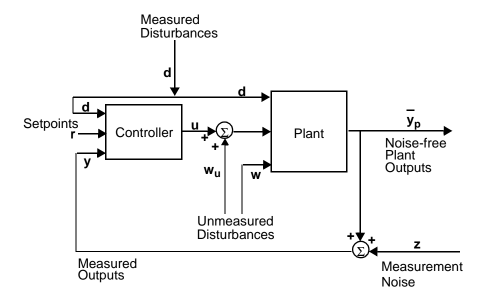
See Also

addmd, addmod, addumd, appmod, paramod

Combines a plant model and a controller model in the MPC **mod** format, yielding a closed-loop system model in the MPC format. This can be used for stability analysis and linear simulations of closed-loop performance.

Syntax

Description



pmod

Is a model (in the **mod** format) representing the pl ant in the above diagram.

i mod

Is a model (in the same format) that is to be used to design the MPC controller block shown in the diagram. It may be the same as pmod (in which case there is no *model error* in the controller design), or it may be different.

Ks

Is a controller gain matrix, which must have been calculated by the function smpccon.

Kest

Is an (optional) estimator gain matrix. If omitted or set to an empty matrix, the default is to use the *DMC estimator* index DMC estimator. See the documentation for the function smpcest for more details on the design and proper format of Kest.

smpccl

Calculates a model of the closed-loop system, cl mod. It is in the **mod** format and can be used, for example, with analysis functions such as smpcgain and smpcpole, and with simulation routines such as mod2step and dlsimm. smpccl also calculates a model of the controller element, cmod.

The closed-loop model, cl mod, has the following state-space representation:

$$x_{cl}(k+1) = \Phi_{cl}x_{cl}(k) + \Gamma_{cl}u_{cl}(k)$$
$$y_{cl}(k) = C_{cl}x_{cl}(k) + D_{cl}u_{cl}(k)$$

where x_{cl} is a vector of n state variables, \mathbf{u}_{cl} is a vector of input variables, y_{cl} is a vector of outputs, and Φ_{ch} Γ_{ch} C_{ch} and D_{cl} are matrices of appropriate size. The expert user may want to know the significance of the state variables in x_{cl} . They are (in the following order):

- The n_p states of the plant (as specified in pmod),
- The n_i changes in the state estimates (based on the model specified in i mod and the estimator gain, Kest),
- The n_y estimates of the *noise-free* plant output $\hat{y} = (k|k-1)$ (from the state estimator),
- n_u integrators that operate on the Δu signal produced by the standard MPC formulation to yield a u signal that can be used as input to the plant and as a closed-loop output, and
- n_d differencing elements that operate on the d signal to produce the Δd signal required in the standard MPC formulation. If there are no measured disturbances, these states are omitted.

The closed-loop input and output variables are:

$$u_{cl}(k) = \begin{bmatrix} r(k) \\ z(k) \\ w_{u}(k) \\ d(k) \\ w(k) \end{bmatrix} \text{ and } y_{cl}(k) = \begin{bmatrix} \overline{y}_{p}(k) \\ u(k) \\ \hat{y}(k|k) \end{bmatrix}$$

where $\hat{y} = (k|k)$ is the estimate of the noise-free plant output at sampling period k based on information available at period k. This estimate is generated by the controller element.

Note that u_{cl} will include d and/or w automatically whenever pmod includes measured disturbances and/or unmeasured disturbances. Thus the length of the u_{cl} vector will depend on the inputs you have defined in pmod and i mod. Similarly, ycl depends on the number of outputs and manipulated variables. Let m and p be the lengths of u_{cl} and v_{cl} respectively. Then

$$m = 2n_y + n_u + n_d + n_w$$

$$p = 2n_y + n_u$$

The state-space form of the controller model, cmod, can be written as:

$$x_c(k+1) = \Phi_c x_c(k) + \Gamma_{cl} u_c(k)$$
$$y_c(k) = C_c x_c(k) + D_c u_c(k)$$

where

$$u_c(k) = \begin{bmatrix} r(k) \\ y(k) \\ d(k) \end{bmatrix}$$
 and $y_c(k) = u(k)$

and the controller states are the same as those of the closed loop system except that the n_p plant states are not included.

Examples

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

We build this model using the MPC Toolbox functions poly2tfd and tfd2mod.

```
g11=poly2tfd(12.8, [16.7 1], 0, 1);
g21=poly2tfd(6.6, [10.9 1], 0, 7);
g12=poly2tfd(-18.9, [21.0 1], 0, 3);
g22=poly2tfd(-19.4, [14.4 1], 0, 3);
delt=3; ny=2;
i mod=tfd2mod(delt, ny, g11, g21, g12, g22);
pmod=i mod; % No plant/model mismatch
```

Now we design the controller. Since there is delay, we use M < P: We specify the defaults for the other tuning parameters, uwt and ywt, then calculate the controller gain:

```
P=6; % Prediction horizon.
M=2; % Number of moves (input horizon).
ywt=[]; % Output weights (default - unity on
% all outputs).
uwt=[]; % Man. Var weights (default - zero on
% all man. vars).
Ks=smpccon(i mod, ywt, uwt, M, P);
```

Now we can calculate the model of the closed-loop system and check its poles for stability:

```
cl mod=smpccl (pmod, i mod, Ks);
maxpol e=max(abs(smpcpol e(cl mod)))
```

The result is:

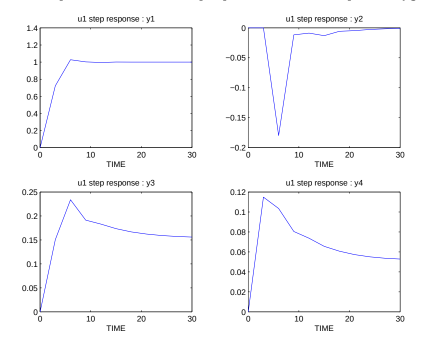
```
maxpole = 0.8869
```

Since this is less than 1, the plant and controller combination will be closed-loop stable. (The closed-loop system has 20 states in this example).

You can also use the closed-loop model to calculate and plot the step response with respect to *all* the inputs. The appropriate commands are:

```
tend=30;
cl step=mod2step(cl mod, tend);
pl otstep(cl step)
```

Since the closed-loop system has m = 6 inputs and p = 6 outputs, only one of the plots is reproduced here. It shows the response of the first 4 closed-loop outputs to a step in the first closed-loop input, which is the setpoint for y_1 :



Closed-loop outputs y_1 and y_2 are the true plant outputs (noise-free). Output y_1 goes to the new setpoint quickly with a small overshoot. This causes a small, short-term disturbance in y_2 . The plots for y_3 and y_4 show the required variation in the manipulated variables.

The following commands show how you could use dl si mm to calculate the response of the closed-loop system to a step in the setpoint for y_1 , with added random measurement noise.

```
r=[ones(11, 1) zeros(11, 1)];
z=0.1*rand(11, 2);
wu=zeros(11, 2);
d=[];
w=[];
w=[];
ucl=[r z wu d w];
[phi cl, gamcl, ccl, dcl]=mod2ss(cl mods);
ycl=dlsimm(phi cl, gamcl, ccl, dcl, ucl);
y=ycl(:,1:2); u=ycl(:,3:4); ym=ycl(:,5:6);
```

Restrictions

- i mod and pmod must have been created using the same sampling period, and an equal number of outputs, measured disturbances, and manipulated variables.
- Both i mod and pmod must be strictly proper, i.e., the D matrices in their state-space descriptions must be zero. *Exception:* the last n_w columns of the D matrices may be nonzero, i.e., the unmeasured disturbance may have an immediate effect on the outputs.

See Also

mod2step, scmpc, smpccon, smpcest, smpcgain, smpcpole, smpcsim

Calculates MPC controller gain using a model in MPC **mod** format.

Syntax

```
Ks = smpccon(i mod)
```

Ks = smpccon(i mod, ywt, uwt, M, P)

Description

Combines the following variables (most of which are optional and have default values) to calculate the state-space MPC gain matrix, Ks.

i mod is the model of the process to be used in the controller design (in the **mod** format).

The following input variables are optional:

ywt

Is a matrix of weights that will be applied to the setpoint tracking errors. If you use $ywt=[\]$ or omit it, the default is equal (unity) weighting of all outputs over the entire prediction horizon. If $ywt+[\]$, it must have n_y columns, where n_y is the number of outputs. All weights must be ≥ 0 .

You may vary the weights at each step in the prediction horizon by including up to P rows in ywt. Then the first row of n_y values applies to the tracking errors in the first step in the prediction horizon, the next row applies to the next step, etc.

If you supply only nrow rows, where $1 \le nrow < P$, smpccon will use the last row to fill in any remaining steps. Thus if you wish the weighting to be the same for all P steps, you need only specify a single row.

uwt

Same format as ywt, except that uwt applies to the *changes* in the manipulated variables. If you use $uwt=[\]$ or omit it, the default is zero weighting. If $uwt+[\]$, it must have n_u columns, where n_u is the number of manipulated variables.

M

There are two ways to specify this variable:

If it is a *scalar*, smpccon interprets it as the input horizon (number of moves) as in DMC.

If it is a *row vector* containing n_b elements, each element of the vector indicates the number of steps over which $\Delta u = 0$ during the optimization and smpccon interprets it as a set of n_b blocking factors. There may be $1 \le n_b \le P$ blocking factors, and their sum must be $\le P$.

If you set $M=[\]$ or omit it, the default is M=P, which is equivalent to M=ones(1,P).

P

The number of sampling periods in the prediction horizon. If you set $P=[\]$ or omit it, the default is P=1.

If you take the default values for all the optional variables, *y*ou get the "perfect controller," i.e., a model-inverse controller. This controller is not applicable in the following situations:

- When one or more outputs cannot respond to the manipulated variables within 1 sampling period due to time delay, the plant-inverse controller is unrealizable. To counteract this you can penalize changes in the manipulated variables (variable uwt), use blocking (variable M), and/or make P>>M.
- When i mod contains transmission zeros outside the unit circle the plant-inverse controller will be unstable. To counteract this, you can use blocking (variable M), restrict the input horizon (variable M), and/or penalize changes in the manipulated variables (variable uwt).

The model-inverse controller is also relatively sensitive to model error and is best used as a point of reference from which you can progress to a more robust design.

Algorithm

The controller gain is a component of the solution to the optimization problem:

Minimize
$$J(k) = \sum_{j=1}^{p} \sum_{i=1}^{n_{y}} (ywt_{i}(j)[r_{i}(k+j) - \hat{y}_{i}(k+j)])^{2} + \sum_{j=1}^{n_{b}} \sum_{i=1}^{n_{u}} (uwt_{i}(j)\Delta \hat{u}_{i}(j))^{2}$$

with respect to $\Delta u_i(j)$ (a series of current and future moves in the manipulated variables), where $\hat{y}_i(\mathbf{k}+j)$ is a prediction of output i at a time j sampling periods into the future (relative to the current time, k), which is a function of

 $\Delta \hat{u}_i(j)$, $r_i(k+j)$ is the corresponding future setpoint, and n_b is the number of blocks or moves of the manipulated variables.

References

Ricker, N. L. "Use of Quadratic Programming for Constrained Internal Model Control," *Ind. Eng. Chem. Process Des. Dev.*, 1985, 24, 925–936.

Ricker, N. L. "Model-predictive control with state estimation," I & ECRes., 1990, 29, 374.

Example

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

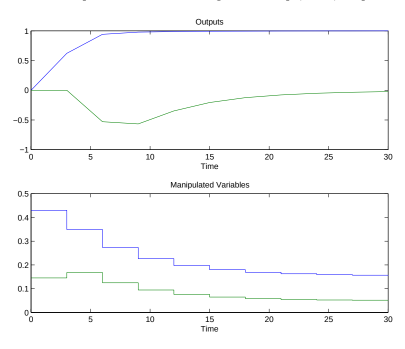
See the smpccl example for the commands that build the model and a simple controller for this process.

Here is a slightly more complex design with blocking and time-varying weights on the manipulated and output variables:

```
P=6; M=[2 4];
uwt=[1 0; 0 1];
ywt=[1 0.1; 0.8 0.1; 0.1 0.1];
Ks=smpccon(i mod, ywt, uwt, M, P);
tend=30; r=[1 0];
[y, u]=smpcsi m(pmod, i mod, Ks, tend, r);
```

There is no particular rationale for using time varying weights in this case—it is only for illustration. The manipulated variables will make 2 moves during the prediction horizon (see value of M, above). The uwt selection gives u_1 a unity weight and u_2 a zero weight for the first move, then switches the weights for the second move. If there had been any additional moves they would have had the same weighting as the second move.

The ywt value assigns a constant weight of 0.1 to y_2 , and a weight that decreases over the first 3 periods to y_1 . The weights for periods 4 to 6 are the same as for period 3. The resulting closed-loop (servo) response is:



See Also

scmpc, smpccl, smpcsim

Sets up a state-estimator gain matrix for use with MPC controller design and simulation routines using models in MPC **mod** format. Can use either a disturbance/noise model that you specify, or a simplified form in which each output is affected by an independent disturbance (plus measurement noise).

Syntax

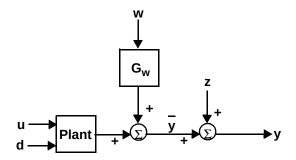
For the general case:

```
[Kest] = smpcest(i mod, Q, R)
```

For simplified disturbance modeling:

```
[Kest, newmod] = smpcest(i mod)
[Kest, newmod] = smpcest(i mod, tau, si gnoi se)
```

Description



In the above block diagram, u is a vector of n_u manipulated variables $(n_u \ge 1)$, d is a vector of n_d measured disturbances $(n_d \ge 0)$, w is a vector of unmeasured disturbances, z is measurement noise, y is a vector of outputs, and \bar{y} represents these outputs before the addition of measurement noise. The objective of the state estimator in MPC is to estimate the present and future values of \bar{y} , rejecting as much of the measurement noise as possible. The inputs u and d are assumed perfectly measurable, whereas w and z are unknown and must be inferred from the measurements. G_w is a transfer function matrix representing the effect of each element of w on each output in y.

General Case

i mod

Is the model (in **mod** format) to be used as the basis for the state estimator. It should be the same as that used to calculate the controller gain (see smpccon). It must include a model of the disturbances, i.e., the G_W element in the above

diagram. You could, for example, use addumd to combine a plant and disturbance model, yielding a composite model in the proper form.

Q

Is a symmetric, positive semi-definite matrix giving the covariances of the disturbances in w. It must be n_w by n_w where n_w (\geq 1) is the number of unmeasured disturbances in i mod (i.e., the length of w).

R

Is a symmetric, positive-definite matrix giving the covariances of the measurement noise, z. It must be n_{ym} by n_{ym} , where n_{ym} (\geq 1) is the number of measured outputs in i mod.

The calculated output variable is:

Kest

The estimator gain matrix. It will contain $n + n_y$ rows and n_{ym} columns, where n is the number of states in i mod, and n_y is the total number of outputs (measured plus unmeasured).

Simplified disturbance modeling

For the *simplified disturbance/noise model* we make the following assumptions:

- The vectors w, z, y and \bar{y} are all length n_v
- G_W is diagonal. Thus each element of W affects one (and only one) element of Y. Diagonal element G_{Wi} has the discrete (sampled-data) form:

$$G_{wi}(q) = \frac{1}{q - a_i}$$

where $a_i = e^{-T/\tau^i}$, $0 \le \tau_i \le \infty$, and T is the sampling period.

As $\tau_i \to 0$, $\mathit{Gwi}(q)$ approaches a unity gain, while as $\tau_i \to \infty$, G_{wi} becomes an integrator.

- Element *i* of Δw is a stationary white-noise signal with zero mean and standard deviation σ_{wi} (where $w_i(k) = w_i(k) w_i(k-1)$).

The input variables are then as follows:

i mod

Is the model (in **mod** format) to be used as the basis for the state estimator. It should be the same as that used to calculate the controller gain (see smpccon).

tau

Is a *row vector*, length n_y giving the values of τ_i to be used in eq. 1. Each element must satisfy: $0 \le \tau_i \le \infty$. If you use tau=[], smpcest uses the default, which is n_v zeros.

si gnoi se

Is a *row vector*, length n_y giving the signal-to-noise ratio for the each disturbance, defined as $\gamma_i = \sigma_{wi} = \sigma_{zi}$. Each element must be nonnegative. If omitted, smpcsi m uses an infinite signal-to-noise ratio for each output.

The calculated output variables are:

Kest

The estimator gain matrix.

newmod

The modified version of i mod, which must be used in place of i mod in any simulation/analysis functions that require Kest (e.g., smpccl, smpcsi m, scmpc).

If i mod contains n states, and there are n_1 outputs for which $\tau_i > 0$, then newmod will have $n + n_1$ states. The optimal gain matrix, Kest, will have $n + n_1 + n_y$ rows and n_{ym} columns. The first n rows will be zero, the next n_1 rows will have the gains for the estimates of the n_1 added states (if any), and the last n_v rows will have the gains for estimating the noise-free outputs, \bar{y} .

Examples

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix} + \begin{bmatrix} \frac{3.8e^{-8s}}{14.9s+1} \\ \frac{4.9e^{-3s}}{13.2s+1} \end{bmatrix} w(s)$$

The following statements build two models: pmod, which contains the model of the disturbance, *w*, and i mod, which does not.

```
g11=poly2tfd(12.8, [16.7 1], 0, 1);

g21=poly2tfd(6.6, [10.9 1], 0, 7);

g12=poly2tfd(-18.9, [21.0 1], 0, 3);

g22=poly2tfd(-19.4, [14.4 1], 0, 3);

delt=1; ny=2;

imod=tfd2mod(delt, ny, g11, g21, g12, g22);

gw1=poly2tfd(3.8, [14.9 1], 0, 8);

gw2=poly2tfd(4.9, [13.2 1], 0, 3);

pmod=addumd(imod, tfd2mod(delt, ny, gw1, gw2));
```

Calculate the gain for a typical MPC (unconstrained) controller

```
P=6; M=2;
ywt=[ ]; uwt=[1 1];
Ks=smpccon(i mod, ywt, uwt, M, P);
```

Next design an estimator using the G_W model in pmod. The choices of Q and R are arbitrary. R was made relatively small (since measurement noise will be negligible in the simulations).

```
Kest1=smpcest(pmod, 1, 0.001*eye(ny));
Ks1=smpccon(pmod, ywt, uwt, M, P);
```

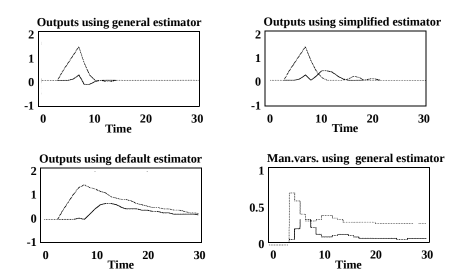
Now design another estimator using a simplified disturbance model in which each output is affected by a disturbance with a first-order time constant of 10 and a signal-to-noise ratio of 3.

```
tau=[10 10]; signoise=[3 3];
[Kest2, newmod]=smpcest(i mod, tau, signoise);
Ks2=smpccon(newmod, ywt, uwt, M, P);
```

Compare the performance of these two estimators to the default (DMC) estimator when there is a unit step in *w*:

```
r=[ ]; ulim=[ ]; z=[ ]; d=[ ]; w=[1]; wu=[ ]; tend=30; [y_1, u_1]=smpcsim(pmod, pmod, Ks1, tend, r, ulim, Kest1, z, d, w, wu); [y_2, u_2]=smpcsim(pmod, newmod, Ks2, tend, r, ulim, Kest2, z, d, w, wu); [y_3, u3]=smpcsim(pmod, imod, Ks, tend, r, ulim, [ ], z, d, w, wu);
```

The solid lines in the following plots are for y_1 (or u_1) and the dashed lines are for y_2 (or u_2). Both outputs have setpoints at zero. You can see that the default estimator is much more sluggish than the others in counteracting this type of disturbance. The simplified disturbance design does nearly as well as that using the exact model of the disturbances. The main difference is that it allows more error in y_1 following the disturbance in y_2 .



The first 14 states in both i mod and pmod are for the response of the outputs to *u*. Since the unmeasured disturbance has no effect on them, their gains are

zero. pmod contains 10 additional disturbance states and there are 2 outputs, so the last 12 rows of Kest 1 are nonzero:

Kest1(15:26,:) =	
8.8659	
7. 1499	
5. 1314	
2. 7748	
0.0411	
- 0. 0182	
- 0. 0008	
0.0001	
0.0000	
- 0. 0000	
0.0000	
0. 9925	

and the last 4 rows of Kest 2 are nonzero:

Algorithm

In the general case, smpcest uses dl qe2 to calculate the optimal estimator gain, Kest. In the simplified case, it uses an analytical solution of the discrete Riccati equation (which is possible to obtain in this case because the disturbances are independent with low-order dynamics).

The number of rows in Kest is larger than that in newmod because the MPC analysis and simulation functions augment the model states with the outputs (see mpcaugss), and Kest must be set up to account for this.

If all $\tau_i = 0$ and all $\gamma_i = \infty$, we get the DMC *estimator*, which has n rows of zeros followed by an identity matrix of dimension n_y . This is the default for all of the MPC analysis and simulation routines that require an estimator gain as input.

smpcest

Important note: smpcest decides whether you are using the general case or the simplified approach by checking the number of output arguments you have supplied. If there is only one, it assumes you want the general case. Otherwise, it proceeds as for the simplified case. It checks the dimensions of your input arguments to make sure they are consistent with this decision.

If you get unexpected results or an error message, make sure you have specified the correct number of output arguments.

See Also

scmpc, smpccl, smpccon, smpcsi m

Purpose Calculates steady-state gain matrix or poles for a system in the MPC **mod**

format.

Syntax g = smpcgai n(mod)

poles = smpcpole(mod)

Description mod is a dynamic model in the MPC **mod** format. smpcgai n and smpcpol e convert it to its equivalent state-space form:

 $x(k+1) = \Phi x(k) + \Gamma v(k)$

$$y(k) = Cx(k) + Dv(k)$$

where v includes all of the inputs in mod. smpcgain then calculates the gain matrix:

$$G = C(I - \Phi)^{1}\Gamma + D$$

which contains n_y rows, corresponding to each of the outputs in mod, and $n_u + n_d + n_w$ columns, corresponding to each of the inputs.

smpcpol e calculates the poles, i.e., the eigenvalues of the $\boldsymbol{\Phi}$ matrix.

Restriction If mod is not asymptotically stable, smpcgain terminates with an error message.

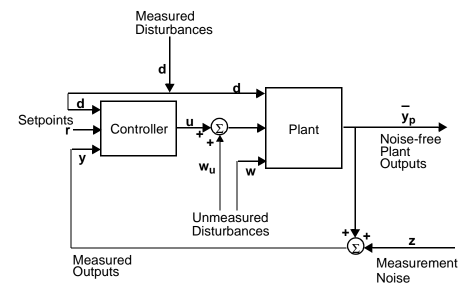
See Also mod

Simulates closed-loop systems with *saturation constraints* on the manipulated variables using models in the MPC **mod** format. Can also be used for open-loop simulations.

Syntax

```
yp = smpcsi m(pmod, i mod, Ks, tend, r)
[yp, u, ym] = smpcsi m(pmod, i mod, Ks, tend, r, usat, ...
Kest, z, d, w, wu)
```

Description



smpcsi m provides a convenient way to simulate the performance of the type of system shown in the above diagram. The required input variables are as follows:

pmod

Is a model in the MPC **mod** format that is to represent the plant.

i mod

Is a model in the MPC **mod** format that is to be used for state estimation in the controller. In general, it can be different from pmod if you wish to simulate the effect of plant/controller model mismatch. Note, however, that i mod should be the same as that used to calculate Ks.

Ks

Is the MPC controller gain matrix, usually calculated using the function smpccon.

If you set Ks to an empty matrix, smpcsi m will do an open-loop simulation. Then the inputs to the plant will be r (which must be set to the vector of manipulated variables in this case), d, w, and wu. The measurement noise input, z, will be ignored.

tend

Is the desired duration of the simulation (in time units).

 \mathbf{r}

Is *normally* a setpoint matrix consisting of N rows and n_y columns, where n_y is the number of output variables, y:

$$\mathbf{r} = \begin{bmatrix} r_1(1) & r_2(1) & \dots & r_{n_y}(1) \\ r_1(2) & r_2(2) & \dots & r_{n_y}(2) \\ \vdots & \vdots & \dots & \vdots \\ r_1(N) & r_2(N) & \dots & r_{n_y}(N) \end{bmatrix}$$

where $r_i(k)$ is the setpoint for output j at time t = kT, and T is the sampling period (as specified by the minfo vector in the **mod** format of pmod and i mod). If tend > NT, the setpoints vary for the first N periods in the simulation, as specified by r, and are then held constant at the values given in the last row of r for the remainder of the simulation.

In many simulations one wants the setpoints to be constant for the entire time, in which case r need only contain a single row of n_v values.

If you set $r=[\]$, the default is a row of n_y zeros.

For *open-loop* simulations, r specifies the *manipulated variables* and must contain n_{ij} columns.

The following input variables are optional. In general, setting one of them equal to an empty matrix causes smpcsi m to use the default value, which is given in the description.

usat

Is a matrix giving the saturation limits on the manipulated variables. Its format is as follows:

$$\text{usat} = \begin{bmatrix} u_{min, 1}(1) & \dots & u_{min, n_u}(1) \\ u_{min, 1}(2) & \dots & u_{min, n_u}(2) \\ \vdots & \dots & \vdots \\ u_{min, 1}(N) & \dots & u_{min, n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} u_{max, 1}(1) & \dots & u_{max, n_u}(1) \\ u_{max, 1}(2) & \dots & u_{max, n_u}(2) \\ \vdots & \dots & \vdots \\ u_{max, 1}(N) & \dots & u_{max, n_u}(N) \end{bmatrix}$$

$$\begin{bmatrix} \Delta u_{max, 1}(1) & \dots & \Delta u_{max, n_u}(1) \\ \Delta u_{max, 1}(2) & \dots & \Delta u_{max, n_u}(2) \\ \vdots & \dots & \vdots \\ \Delta u_{max, 1}(N) & \dots & \Delta u_{max, n_u}(N) \end{bmatrix}$$

Note that it contains three matrices of Nrows. Nmay be different than that for the setpoint matrix, r, but the idea is the same: the saturation limits will vary for the first N sampling periods of the simulation, then be held constant at the values given in the last row of usat for the remaining periods (if any).

The first matrix specifies the *lower bounds* on the n_u manipulated variables. For example, $u_{min'j}(k)$ is the lower bound for manipulated variable j at time t = kT in the simulation. If $u_{min'j}(k) = -inf$, manipulated variable j will have no lower bound at t = kT.

The second matrix gives the *upper bounds* on the manipulated variables. If $u_{max,j}(k) = inf$, manipulated variable j will have no upper bound at t = kT.

The lower and upper bounds may be either positive or negative (or zero) as long as $u_{min:j}(k) \le u_{max,j}(k)$.

The third matrix gives the limits on the rate of change of the manipulated variables. In other words, smpcsi m will force $|u_j(k) - u_j(k-1)| \le \Delta u_{max,j}(k)$. The limits on the rate of change must be nonnegative.

The default is no saturation constraints, i.e., all the umin values will be set to -inf, and all the u_{max} and Δu_{max} values will be set to inf.

Note: Saturation constraints are enforced by simply "clipping" the manipulated variable moves so that they satisfy all constraints. This is a nonoptimal solution that, in general, will differ from the results you would get using the ulim variable in scmpc.

Kest

Is the estimator gain matrix. The default is the DMC estimator. See smpcest for more details.

Z

Is measurement noise that will be added to the outputs (see above diagram). The format is the same as for r. The default is a row of n_v zeros.

d

Is a matrix of measured disturbances (see above diagram). The format is the same as for r, except that the number of columns is n_d rather than n_y The default is a row of n_d zeros.

W

Is a matrix of unmeasured disturbances (see above diagram). The format is the same as for r, except that the number of columns is n_w rather than n_y The default is a row of n_w zeros.I

wu

Is a matrix of unmeasured disturbances that are added to the manipulated variables (see above diagram). The format is the same as for r, except that the number of columns is n_u rather than n_v . The default is a row of n_u zeros.

Note: You may use a different number of rows in the matrices r, usat, z, d, w and wu, should that be appropriate for your simulation.

The calculated outputs are as follows (all but yp are optional):

уp

Is a matrix containing M rows and n_y columns, where M = max(fix(tend=T)+1, 2). The first row will contain the initial condition, and row k-1 will give the values of the plant outputs, y (see above diagram), at time t=kT.

u

Is a matrix containing the same number of rows as yp and n_u columns. The time corresponding to each row is the same as for yp. The elements in each row are the values of the manipulated variables, u (see above diagram).

Note: The u values are those coming from the controller *before* the addition of the unmeasured disturbance, w_{ur}

ym

Is a matrix of the same structure as yp, containing the values of the predicted output from the state estimator in the controller. These will, in general, differ from those in yp if i mod|pmod and/or there are unmeasured disturbances. *The prediction includes the effect of the most recent measurement, i.e.*, it is $\hat{y}(k|k)$.

Examples

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

The following statements build the model and calculate the MPC controller gain:

```
g11=poly2tfd(12.8, [16.7 1], 0, 1);

g21=poly2tfd(6.6, [10.9 1], 0, 7);

g12=poly2tfd(-18.9, [21.0 1], 0, 3);

g22=poly2tfd(-19.4, [14.4 1], 0, 3);

delt=3; ny=2;

i mod=tfd2mod(delt, ny, g11, g21, g12, g22);

pmod=i mod;

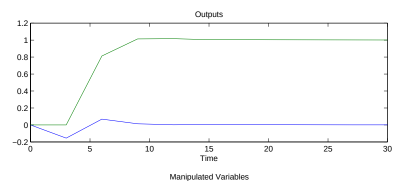
P=6; M=2;

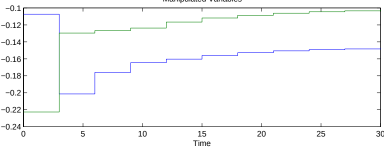
ywt=[]; uwt=[1 1];

Ks=smpccon(i mod, ywt, uwt, M, P);
```

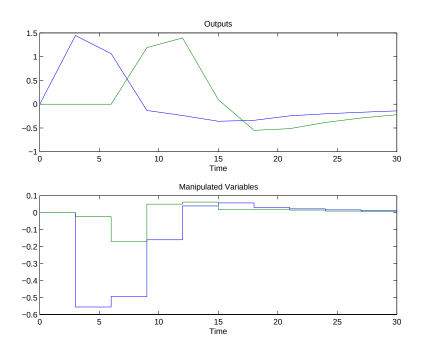
Simulate and plot the closed-loop performance for a unit step in the setpoint for y_2 , occurring at t = 0.

```
tend=30; r=[0\ 1];
[y, u]=smpcsim(pmod, i mod, Ks, tend, r);
plotall(y, u, delt), pause
```



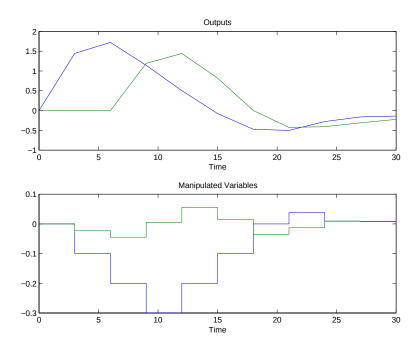


Try a pulse change in the disturbance that adds to u_1 :



For the same disturbance as in the previous case, limit the rates of change of both manipulated variables.

```
usat=[-inf -inf inf inf 0.1 0.05]; [y, u]=smpcsim(pmod, i mod, Ks, tend, r, usat, Kest, z, d, w, wu); plotall(y, u, delt), pause
```



Restrictions

- Initial conditions of zero are used for all states in i mod and pmod. This simulates the condition where all variables represent a deviation from a steady-state initial condition.
- The first $n_u + n_d$ columns of the D matrices in pmod and i mod must be zero. In other words, neither u nor d may have an immediate effect on the outputs.

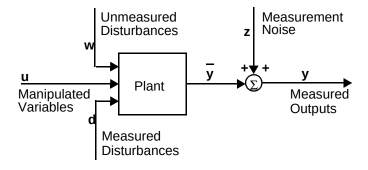
See Also

plotall, ploteach, scmpc, smpccl, smpccon, smpcest

Converts a discrete-time state-space system model into the MPC **mod** format.

Syntax

Description



Consider the process shown in the above block diagram. ss2mod assumes the following state-space representation:

$$x(k+1) = \Phi x(k) + \Gamma_u u(k) + \Gamma_d d(k) + \Gamma_w w(k)$$

$$y(k) = \overline{y}(k) + z(k)$$

$$= Cx(k) + D_u u(k) + D_d d(k) + D_w w(k) + z(k)$$

where x is a vector of n state variables, u represents n_u manipulated variables, d represents n_d measured disturbances, w represents n_w unmeasured disturbances, y is a vector of n_y plant outputs, z is measurement noise, and Φ , Γ_{up} etc., are constant matrices of appropriate size. The variable $\bar{y}(k)$ represents the plant output before the addition of measurement noise. We further define:

$$D = [D_u D_d D_w]$$
$$\Gamma = [\Gamma_u \Gamma_d \Gamma_w]$$

ss2mod uses the Φ , Γ , C, and D matrices you supply to build a model, pmod, in the MPC **mod** format. See the mod section for more details.

You can also divide the outputs into n_{ym} measured outputs and n_{yu} unmeasured outputs, where $n_{ym} + n_{yu} = n_y$. Then the first n_{ym} elements in y and the first n_{ym} rows in C and D are assumed to be for the measured outputs, and the rest are for the unmeasured outputs.

mi nfo is an optional variable that allows you to specify certain characteristics of the system. The general form is a *row vector* with 7 elements, the interpretation of which is as follows:

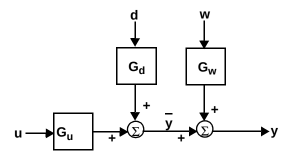
- minfo(1) *T*, the sampling period used to create the model.
 - (2) n, the number of states.
 - (3) n_{ip} the number of manipulated variable inputs.
 - (4) n_d , the number of measured disturbances.
 - (5) n_w the number of unmeasured disturbances.
 - (6) n_{vm} the number of measured outputs.
 - (7) n_{vv} the number of unmeasured outputs.

If you specify mi nfo as a scalar, ss2mod takes it as the sampling period and sets the remaining elements of mi nfo as follows:

```
\min \text{nfo}(2) = \# \text{ rows in phi}, \quad \min \text{nfo}(3) = \# \text{ columns in gam},
\min \text{nfo}(4) = \min \text{nfo}(5) = 0, \quad \min \text{nfo}(6) = \# \text{ rows in c}, \quad \min \text{nfo}(7) = 0.
```

In other words, the default is to assume that all inputs are manipulated variables and all outputs are measured. If you omit minfo, ss2mod sets the sampling period to 1 and uses the defaults for the remaining elements.

Example



Suppose you have the situation shown in the above diagram where u, d, w, and y are scalar signals, and the three transfer functions are first-order:

$$G_u(z) = \frac{0.7}{1 - 0.9z^{-1}}$$
 $G_d(z) = \frac{-1.5}{1 - 0.85z^{-1}}$ $G_w(z) = \frac{1}{1 + 0.6z^{-1}}$

The sampling period is T = 2.

One way to build the model of the complete system is to convert these to state-space form and use ss2mod:

```
[ phi u, gamu, cu, du] = tf2ss(0.7, [1 -0.9]);
[ phi d, gamd, cd, dd] = tf2ss(-1.5, [1 -0.85]);
[ phi w, gamw, cw, dw] = tf2ss(1, [1 0.6]);
[ phi, gam, c, d] = mpcparal (phi u, gamu, cu, du, phi d, gamd, cd, dd);
[ phi, gam, c, d] = mpcparal (phi, gam, c, d, phi w, gamw, cw, dw);
del t = 2;
mi nfo = [ del t 3 1 1 1 1 0];
pmod = ss2mod(phi, gam, c, d, mi nfo)
```

You must be careful to build up the parallel structure in the correct order. For example, the columns corresponding to Γ_u must always come first in Γ .

Another, more foolproof way is to use the addmd and addumd functions:

```
ny=1;
gu=poly2tfd(0.7,[1 -0.9], delt);
gd=poly2tfd(-1.5,[1 -0.85], delt);
gw=poly2tfd(1,[1 0.6], delt);
pmod=tfd2mod(delt, ny, gu);
pmod=addmd(pmod, tfd2mod(delt, ny, gw));
pmod=addumd(pmod, tfd2mod(delt, ny, gw))
```

Using either approach, the result is:

```
pmod =
2.0000
          3.0000
                    1.0000
                              1.0000
                                        1.0000
                                                  1.0000
                                                              0
NaN
          0.9000
                    0
                              0
                                        1.0000
                                                              0
                                                  0
0
          0
                    0.8500
                              0
                                        0
                                                  1.0000
                                                              0
0
          0
                             - 0. 6000
                                        0
                                                  0
                                                              1.0000
0
          0.7000
                  - 1. 5000
                              1.0000
                                        0
                                                  0
                                                              0
```

See Also

mod format, mod2ss

Uses a model in state-space format to calculate the step response of a SISO or MIMO system, in MPC *step* format.

Syntax

```
plant = ss2step(phi, gam, c, d, tfinal)
plant = ss2step(phi, gam, c, d, tfinal, delt1, delt2, nout)
```

Description

The input variables *phi*, *gam*, *c*, and *d* are assumed to be a state-space model of a process. The model can be either continuous time:

$$\dot{x}(t) = \Phi x(t) + \Gamma u(t)$$

$$y(t) = Cx(t) + Du(t)$$

or discrete time:

$$x(k+1) = \Phi x(k) + \Gamma u(k)$$

$$y(k) = Cx(k) + Du(k)$$

where x is a vector of n state variables, u is a vector of n_u inputs (usually but not necessarily manipulated variables), y is a vector of n_y plant outputs, and Φ , Γ , etc., are constant matrices of appropriate size. The ss2step function calculates the step responses of all the outputs of this process with respect to all the inputs in u, and puts this information into the variable plant in MPC step format. The section for mod2step describes the step format in detail.

The input variable tfinal is the time at which you would like to end the step response calculation, and delt1 is the sampling period. For continuous systems, use delt1=0. If you do not specify delt1, the default is delt1=0.

The optional input variable del t2 is the desired sampling period for the step response model. If you use del $t2=[\]$ or omit it, the default is del t2=del t1 if delt1 is specified and del t1 neq 0; otherwise, the default is del t2=1.

The optional input variable nout is the output stability indicator. For stable systems, set nout equal to the number of outputs, n_y . For systems with one or more integrating outputs, nout is a column vector of length n_y with nout (i) =0 indicating an integrating output and nout (i) =1 indicating a stable output. If you use nout=[] or omit it, the default is nout= n_y (only stable outputs).

Example

The following process has 3 inputs and 4 outputs (and is the same one used for the example in the mod2step section):

```
phi =di ag([0.3, 0.7, -0.7]);
gam=eye(3);
c=[1 0 0; 0 0 1; 0 1 1; 0 1 0];
d=[1 0 0; zeros(3,3)];
```

The following command duplicates the results obtained with mod2step:

```
delt1=1.5; tfinal=3*1.5;
plant=ss2step(phi, gam, c, d, tfinal, delt1)
```

See Also

plotstep, mod2step, tfd2step

svdfrsp

Purpose Calculates the singular values of a varying matrix, for example, the frequency

response generated by mod2frsp.

Syntax [sigma, omega] = svdfrsp(vmat)

Description vmat is a *varying* matrix which contains the sampled values $F(\omega_1), \dots, F(\omega_N)$ of a matrix function $F(\omega)$.

If the smaller dimension of $F(\omega_i)$ is m, and if $\sigma_I(\omega_i), \ldots, \sigma_m(\omega_i)$ are the singular values of $F(\omega_i)$, in decreasing magnitude, then the output sigma is a matrix of singular values arranged as follows:

$$\operatorname{sigma} = \begin{bmatrix} \sigma_1(\omega_1) & \sigma_2(\omega_1) & \dots & \sigma_m(\omega_1) \\ \sigma_1(\omega_2) & \sigma_2(\omega_2) & \dots & \sigma_m(\omega_2) \\ \vdots & \vdots & & \vdots \\ \sigma_1(\omega_N) & \sigma_2(\omega_N) & \dots & \sigma_m(\omega_N) \end{bmatrix}$$

The output omega is a column vector containing the frequencies ω_1 , . . . , ω_{N^c}

Example See mod2frsp, varying format for an example of the use of this function.

See Also mod2frsp

tfd2mod converts a transfer function (continuous or discrete) from the MPC *tf* format into the MPC **mod** format, converting to discrete time if necessary.

Syntax

$$\verb|model| = \verb|tfd2mod(delt2|, n_y, g1, g2, g3, ..., g25)|$$

Description

Consider a transfer function such as

$$G(s) = \frac{b_0 s^n + b_1 s^{n-1} + \dots + b_n}{a_0 s^n + a_1 s^{n-1} + \dots + a_n}$$

or

$$G(z) = \frac{b_0 + b_1 z^{-1} + \dots + b_n z^{-n}}{a_0 + a_1 z^{-1} + \dots + a_n z^{-n}}$$

The MPC *tf* format is a matrix consisting of three rows:

row 1 The n coefficients of the numerator polynomial, b_0 to b_n .

row 2 The *n* coefficients of the denominator polynomial, a_0 to a_n .

row 3 column 1: The sampling period. This must be zero if the coefficients in the above rows are for a continuous system. It must be positive otherwise.

column 2: The time delay. For a continuous-time transfer function, it is in time units. For a discrete-time transfer function, it is the integer number of sampling periods of time delay.

The *tf* matrix will always have at least two columns, since that is the minimum width of the third row.

The input arguments for tfd2mod are:

del t2

The sampling period for the system. If any of the transfer functions $g1, \ldots, gN$ are continuous-time or discrete-time with sampling period not equal to del t2, tfd2mod will convert them to discrete-time with this sampling period.

ny

The number of output variables in the plant you are modeling.

$$g1, g2, \dots gN$$

A sequence of N transfer functions in the tf format described above, where $N \ge 1$. These are assumed to be the individual elements of a transfer-function matrix:

$$\begin{bmatrix} g_{1,\,1} & g_{1,\,2} & \cdots & g_{1,\,n_u} \\ g_{2,\,1} & g_{2,\,2} & \cdots & g_{2,\,n_u} \\ \vdots & \vdots & \ddots & \vdots \\ g_{n_{j},\,1} & g_{n_{j},\,2} & \cdots & g_{n_{j},\,n_u} \end{bmatrix}$$

Thus it should be clear that N must be an integer multiple (n_u) of the number of outputs, n_v

Also, tfd2mod assumes that you are supplying the transfer functions in a *column-wise* order. In other words, you should first give the n_y transfer functions for input 1 ($g_{1,1}$ to g_{n_y} 1), then the n_y transfer functions for input 2 ($g_{1,2}$ to g_{n_y} 2), etc.

tfd2mod converts the transfer functions to discrete-time, if necessary, and combines them to form the output variable, model, which is a composite system in the MPC mod form.

Example

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix} + \begin{bmatrix} \frac{3.8e^{-8s}}{14.9s+1} \\ \frac{4.9e^{-3s}}{13.2s+1} \end{bmatrix} w(s)$$

The following commands build separate models of the response to the manipulated variables, u, and the unmeasured disturbance, w, all for a sampling period T=3 then combines them using addumd to get a model of the entire system (the pmod variable):

```
\begin{array}{l} g11 = pol\ y2tfd(12.\ 8,\ [16.\ 7\ 1],\ 0,\ 1)\ ;\\ g21 = pol\ y2tfd(6.\ 6,\ [10.\ 9\ 1],\ 0,\ 7)\ ;\\ g12 = pol\ y2tfd(-\ 18.\ 9,\ [21.\ 0\ 1],\ 0,\ 3)\ ;\\ g22 = pol\ y2tfd(-\ 19.\ 4,\ [14.\ 4\ 1],\ 0,\ 3)\ ;\\ del\ t = 3;\ ny = 2;\\ umod = tfd2mod(del\ t,\ ny,\ g11,\ g21,\ g12,\ g22)\ ;\\ gw1 = pol\ y2tfd(3.\ 8,\ [14.\ 9\ 1],\ 0,\ 8)\ ;\\ gw2 = pol\ y2tfd(4.\ 9,\ [13.\ 2\ 1],\ 0,\ 3)\ ;\\ wmod = tfd2mod(del\ t,\ ny,\ gw1,\ gw2)\ ;\\ pmod = addumd(umod,\ wmod)\ ;\\ \end{array}
```

Restriction

The current limit on the number of input transfer functions is N = 25.

See Also

mod, poly2tfd, tfd2step

Calculates the MIMO step response of a model in the MPC *tf* format. The resulting step response is in the MPC *step* format.

Syntax

```
plant = tfd2step(tfinal, delt2, nout, g1)
plant = tfd2step(tfinal, delt2, nout, g1, ..., g25)
```

Description

The input variables are as follows:

tfi nal

Truncation time for step response.

del t2

Desired sampling period for step response.

nout

Output stability indicator. For stable systems, this argument is set equal to the number of outputs, n_y For systems with one or more integrating outputs, this argument is a column vector of length n_y with nout (i) =0 indicating an integrating output and nout (i) =1 indicating a stable output.

$$g1, g2, \dots gN$$

A sequence of N transfer functions in the tf format (see tf format section), where $N \ge 1$. These are assumed to be the individual elements of a transfer-function matrix:

$$\begin{bmatrix} g_{1,\,1} & g_{1,\,2} & \cdots & g_{1,\,n_u} \\ g_{2,\,1} & g_{2,\,2} & \cdots & g_{2,\,n_u} \\ \vdots & \vdots & \ddots & \vdots \\ g_{n_{j},\,1} & g_{n_{j},\,2} & \cdots & g_{n_{j},\,n_u} \end{bmatrix}$$

Thus it should be clear that N must be an integer multiple (n_u) of the number of outputs, n_v

tfd2step assumes that you are supplying the transfer functions in a *column-wise* order. In other words, you should first give the n_y transfer functions for input 1 ($g_{1,1}$ to g_{ny} 1), then the n_y transfer functions for input 2 ($g_{1,2}$ to g_{ny} 2), etc.

The output variable pl ant is the calculated step response of the n_y outputs with respect to all inputs. The format is as described in the step section.

Example

Consider the linear system:

$$\begin{bmatrix} y_1(s) \\ y_2(s) \end{bmatrix} = \begin{bmatrix} \frac{12.8e^{-s}}{16.7s+1} & \frac{-18.9e^{-3s}}{21.0s+1} \\ \frac{6.6e^{-7s}}{10.9s+1} & \frac{-19.4e^{-3s}}{14.4s+1} \end{bmatrix} \begin{bmatrix} u_1(s) \\ u_2(s) \end{bmatrix}$$

which is the same as that considered in the mpcsi m example. We build the individual *tf* format models, then calculate and plot the MIMO step response.

```
g11=poly2tfd(12.8, [16.7 1], 0, 1);
g21=poly2tfd(6.6, [10.9 1], 0, 7);
g12=poly2tfd(-18.9, [21.0 1], 0, 3);
g22=poly2tfd(-19.4, [14.4 1], 0, 3);
delt=3; ny=2; tfinal=90;
plant=tfd2step(tfinal, delt, ny, g11, g21, g12, g22, gw1, gw2);
plotstep(plant)
```

The plots should match the example output in the plotstep description.

Restriction

The current limit on the number of input transfer functions is N = 25.

See Also

mod2step, plotstep, ss2step

th2mod, theta format

Purpose

Converts a SISO or MISO model from the *theta* format (as used in the System Identification Toolbox) to one in the MPC **mod** format. Can also combine such models to form a MIMO system.

Syntax

Description

The System Identification Toolbox allows you to identify single-input, single-output (SISO) and multi-input, single-output (MISO) transfer functions from data. The MISO form relating an output, y, to m inputs, u_1 to u_m , and a noise input, e, is:

$$A(z)y(k) = \frac{B_1(z)}{F_1(z)}u_1(k) + \frac{B_2(z)}{F_2(z)}u_2(k) + \dots + \frac{B_m(z)}{F_m(z)}u_m(k) + \frac{C(z)}{D(z)}e(k)$$

where A, B_i , C, D, and F_i are polynomials in the forward-shift operator, z.

The System Identification Toolbox automatically stores such models in a special format, the *theta* format. See the *System Identification Toolbox User's Guide* for details.

th2mod converts one or more MISO *theta* models into the MPC **mod** format, which you can then use with the MPC Toolbox functions. If you supply a single input argument, th, and a single output argument, umod, then umod will model the response of a single output, y, to m inputs, u_1 to u_m , where $m \ge 1$. The value of m depends on the number of inputs included in the input model, th. Note that umod will reflect the values of the A(z), B(z), and F(z) polynomials in eq. 1.

If you supply a second output argument, emod, it will model the response of y to the noise, e, i.e., the A(z), C(z) and D(z) polynomials in eq. 1.

If you supply p input models $(1 \le p \le 8)$, tfd2mod assumes that they define a MIMO system in the following form:

$$\begin{split} A_1(z)y_1(k) &= \frac{B_{11}(z)}{F_{11}(z)}u_1(k) + \ldots + \frac{B_{1m}(z)}{F_{1m}(z)}u_m(k) + \frac{C_1(z)}{D_1(z)}e_1(k) \\ &\vdots & \vdots & \vdots \\ A_p(z)y_p(k) &= \frac{B_{p1}(z)}{F_{p1}(z)}u_1(k) + \ldots + \frac{B_{pm}(z)}{F_{pm}(z)}u_m(k) + \frac{C_p(z)}{D_p(z)}e_p(k) \end{split}$$

The p output variables have independent noise inputs. In this case, each of the p input models must include the same number of inputs, m. The p outputs are arranged in parallel in the resulting umod output model (and the emod model, if used).

If the th models are auto-regressive (i.e., m = 0), then umod will be set to an empty matrix and only emod will be nonempty.

Example

The following commands create three SISO *theta* models using the mktheta command (System Identification Toolbox), then converts them to the equivalent **mod** form:

```
 \begin{array}{l} th1=mktheta([1\ 0\ -.\ 2],[0\ 0\ -1]);\\ th2=mktheta([1\ -.\ 8\ .\ 1],[0\ -.\ 5\ .\ 3],1,1,1);\\ th3=mktheta([1\ -.\ 2],[0\ 1],[1\ 2\ 0],[1\ -1.\ 2\ .\ 3],\ 1);\\ [umod,emod]=th2mod(th1,th2,th3) \end{array}
```

th2mod, theta format

Tri.	1		
INA	resul	TC	are.

umod =						
1.0000	5. 0000	1. 0000	0	0	3.0000	0
NaN	0	0. 2000	0	0	0	1.0000
0	1. 0000	0	0	0	0	0
0	0	0	0.8000	- 0. 1000	0	1.0000
0	0	0	1.0000	0	0	0
0	0	0	0	0	2.0000	1.0000
0	0	- 1. 0000	0	0	0	0
0	0	0	- 0. 5000	0. 3000	0	0
0	0	0	0	0	1. 0000	
emod =						
	s 1 througl					
1.0000	7. 0000	3. 0000	0	0	3. 0000	0
NaN	0	0. 2000	0	0	0	0
0	1. 0000	0	0	0	0	0
0	0	0	0.8000	- 0. 1000	0	0
0	0	0	1. 0000	0	0	0
0	0	0	0	0	1. 4000	- 0. 5400
0	0	0	0	0	1. 0000	0
0	0	0	0	0	0	1.0000
0	0	0. 2000	0	0	0	0
0	0	0	0.8000	- 0. 1000	0	0
0	0	0	0	0	3. 4000	- 0. 5400
Col um	ns 8 throug	gh 11				
0	0	0	0			
0	1.0000	0	0			
0	0	0	0			
0	0	1. 0000	0			
0	0	0	0			
0.0600	0	0	1. 0000			
0	0	0	0			
0	0	0	0			
0	1. 0000	0	0			
0	0	1. 0000	0			
0. 00	600	0	0 1.0	0000		

th2mod, theta format

Restriction The System Identification Toolbox must be installed to use this function.

See Also mod

validmod

Purpose Validates an impulse response model for a new set of data.

Syntax yres = validmod(xreg, yreg, theta)

yres = validmod(xreg, yreg, theta, plotopt)

Description Model validation is a very important part of building a model. For a new set of

data, xreg and yreg, the impulse response model is tested by calculating the output residual, yres. theta consists of impulse response coefficients as

determined by routines such as pl sr or ml r.

Optional input, pl otopt, can be supplied to produce various plots. No plot is produced if pl otopt is equal to 0 which is the default; a plot of the actual output and the predicted output is produced if pl otopt=1; two plots — plot of actual

and predicted output, and plot of output residual — are produced for

plotopt=2.

Example See pl sr for an example of the use of this function.

See Also ml r, pl sr

Writes input and output data matrices for a multi-input single-output system such that they can be used in regression routines ${\tt ml}$ r and ${\tt pl}$ s for determining impulse response coefficients.

Syntax

$$[xreg, yreg] = wrtreg(x, y, n)$$

Description

x is the input data of dimension N by n_u where N is number of data points and n_u is number of inputs. y is the output of dimension N by 1. n is number of impulse response coefficients for all inputs. x is rearranged to produce xreg of dimension (N-n-1) by $n*n_u$ while yreg is produced by deleting the first n rows of y. This operation is illustrated as follows:

$$\mathbf{x} = \begin{bmatrix} x_1(1) & \dots & x_{n_u}(1) \\ x_1(2) & \dots & x_{n_u}(2) \\ \vdots & & \vdots \\ x_1(N) & \dots & x_{n_u}(N) \end{bmatrix}$$

$$y = \begin{bmatrix} y(1) \\ \vdots \\ y(N) \end{bmatrix}$$

then

$$\operatorname{xreg} = \begin{bmatrix} x_1(n) & \dots & x_1(1) & \dots & x_{n_u}(n) & \dots & x_{n_u}(1) \\ x_1(n+1) & \dots & x_1(2) & \dots & x_{n_u}(n+1) & \dots & x_{n_u}(2) \\ \vdots & & \vdots & & \vdots & & \vdots \\ x_1(N-1) & \dots & x_1(N-n) & \dots & x_{n_u}N-(1) & \dots & x_{n_u}(N-n) \end{bmatrix}$$

$$yreg = \begin{bmatrix} y(n+1) \\ \vdots \\ y(N) \end{bmatrix}$$

A single sampling delay is assumed for all inputs. y must be a column vector, i.e., only one output can be specified.

wrtreg

Example See $ml \ r$ and $pl \ sr$ for examples of the use of this function.

See Also mlr, plsr

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