PnPMPC Toolbox v. 0.9 - User manual

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March, 2013

\(^1\)The research leading to these results has received funding from the European Union Seventh Framework Programme [FP7/2007-2013] under grant agreement n° 257462 HYCON2 Network of excellence.
Chapter 1

Introduction

The PnPMPC toolbox is a GNU-licensed MatLab toolbox for the modeling of constrained Large Scale Systems (LSS) with linear time-invariant dynamics and for the implementation of the Plug-and-Play (PnP) Decentralized (De) Model Predictive Control (MPC) schemes described in [1]. The PnPMPC toolbox offers also several functionalities for handling zonotopes set and for computing invariant sets.

The realization of the toolbox has received funding from the European Union Seventh Framework Programme [FP7/2007-2013] under grant agreement no 257462 HYCON2 Network of excellence.

The last version of the PnPMPC toolbox can be downloaded at the webpage

http://sisdin.unipv.it/pnmpmc/pnmpmc.php

Please send bug reports, questions or comments to pnpmpc_toolbox@unipv.it

1.1 Notations

- \( \mathbb{R} \) is the set of real number.
- \( \mathbb{N} \) is the set of integers.
- \( a : b \) is the set of integer \( \{a, a + 1, \ldots, b\} \)
- \( A \geq 0 \) means that the matrix \( A \) is positive semi-definite. \( A > 0 \) means that matrix \( A \) is positive definite.
- \( A = \text{diag}(A_{11}, \ldots, A_{ss}) \) is the block diagonal matrix

\[
\begin{bmatrix}
A_{11} & 0 & 0 \\
0 & \ddots & 0 \\
0 & 0 & A_{ss}
\end{bmatrix}
\]  

\hspace{1cm} (1.1)

- \( x[i], u[i], y[i], d[i], u_{cen} \) are column vectors of suitable dimensions.
\( \mathbf{x} = (x_1, x_2, \ldots, x_s) \) means that \( \mathbf{x} \) is composed by column vectors \( x_i, i = 1 : s \) stacked in a single column, i.e.

\[
\mathbf{x} = \begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_s
\end{bmatrix}
\]  

(1.2)

- The symbol \( \oplus \) denotes the Minkowski sum, e.g. \( A = B \oplus C \) if and only if \( A = \{ a : a = b + c, \forall b \in B, \forall c \in C \} \).

- \( \bigoplus_{i=1}^s G_i = G_1 \oplus \ldots \oplus G_s \).

- \( \times \) indicates the cartesian product and \( \prod_{i=1}^s G_i = G_1 \times \ldots \times G_s \).

**Definition 1.** A polyhedron \( \mathbb{X} \) is the intersection of a finite number of closed half spaces. Therefore we can represent a polyhedron either as

\[
\mathbb{X} = \{ x \in \mathbb{R}^n : H x \leq K \}
\]

where \( H \in \mathbb{R}^{v \times n} \) and \( K \in \mathbb{R}^v \) (H-representation) or as a convex hull of vertices (V-representation).

**Definition 2.** A zonotope is a centrally symmetric convex polytope: given a vector \( p \in \mathbb{R}^n \) and a matrix \( \Xi \in \mathbb{R}^{n \times m} \), the zonotope \( \mathbb{X} \subseteq \mathbb{R}^n \) is the set \( \mathbb{X} = \{ x \mid x = p + \Xi d, \|d\|_{\infty} \leq 1 \} \), with \( d \in \mathbb{R}^m \). We will refer to this representation as G-representation.

### 1.2 Required toolboxes

The PnPMPC toolbox requires the following toolboxes to be installed.

- Control System Toolbox [2] which implements the class ss (state-space). We also use the function c2d for time-discretization.
- MPT toolbox [3] which implements the polytope class.
- YALMIP toolbox [4] which include the optimization function solvesdp that is called for solving optimization problems.
- GraphViz4MatLab toolbox [5] that allows one to plot the graph of a large-scale system composed by interconnected systems.

Please note that MPT, YALMIP and GraphViz4MatLab get automatically installed when installing PnPMPC (with an option to avoid their installation in case you already have them. One can check if all required toolboxes are correctly installed with the following instructions.

```
yalmiptest
mpt_init
```
1.3 Installed solvers

The following solvers get automatically installed when installing PnPMPMC.

- Sedumi [6].
- GLPK (GNU Linear Programming Kit) [7].
- SDPT3 [8].

1.4 Installation of the PnPMPMC toolbox

- **Step 1** Add the folder of the PnPMPMC toolbox and its subfolders in the MatLab path.
- **Step 2** Run `pnpmpc_toolbox_init`

**Remark 1:** if you already have installed MPT and/or YALMIP, `pnpmpc_toolbox_init` will give the possibility of skipping their installation. However, we guarantee that PnPMPMC works correctly only with the versions of MPT and YALMIP supplied with PnPMPMC. Therefore, if you want to be 100% sure that everything works, remove any previous copies of MPT and YALMIP from your disk before installing those supplied with PnPMPMC. Also remove any MPT and YALMIP directory from your Matlab path.

**Remark 2:** also for the solvers, `pnpmpc_toolbox_init` will give the possibility of skipping their installation. However, if you want to be 100% sure that everything works, remove any previous copies of the solvers from your disk before installing those supplied with PnPMPMC. Also remove any solver directory from your MatLab path.

**Remark 3:** we tested PnPMPMC-toolbox on MatLab R2009b and newer version for Windows, Linux and MacOS. For older versions of MatLab we cannot guarantee that everything works correctly.

1.5 Directories

The PnPMPMC toolbox consists of the following directories

- `./pnpmpc_toolbox/@epsilon` contains methods for computing outer-approximation of minimal robust positively invariant sets
- `./pnpmpc_toolbox/@localControlLyapunov` contains methods for computing control invariant sets
- `./pnpmpc_toolbox/@lss` contains methods for modeling of LSS
- `./pnpmpc_toolbox/@parameterizedRCI` contains methods for computing robust control invariant sets
- `./pnpmpc_toolbox/@pnpmpc` contains methods for design of PnPMPC controllers
- `./pnpmpc_toolbox/@polytope` contains functions of MPT toolbox improved to use zonotope sets
- `./pnpmpc_toolbox/@subss` contains methods for the modeling of a subsystem of an LSS
- `./pnpmpc_toolbox/@zonotope` contains methods for using zonotope sets
- `./pnpmpc_toolbox/@zonotopeRCI` contains methods for computing zonotopic robust control invariant sets
- `./pnpmpc_toolbox/docs` contains html documentation
- `./pnpmpc_toolbox/examples` contains examples demonstrating the functionalities of PnP MPC toolbox
- `./pnpmpc_toolbox/extra` contains extra useful functions
- `./pnpmpc_toolbox/solvers` contains solvers provided with PnP MPC toolbox
- `./pnpmpc_toolbox/toolbox` contains additional toolboxes provided with PnP MPC toolbox

For familiarizing quickly with PnP MPC toolbox, examples are supplied in the directory `./pnpmpc_toolbox/examples`. These .m files can also be used as templates for your personal experiments.
Chapter 2

Modeling of interconnected systems

In this chapter we describe the class of systems considered in the PnP-MPC toolbox. Features of subsystems and their interconnections will be discussed in details. The use of the PnP-MPC toolbox for modeling purposes will be illustrate in Chapter 3.

2.1 Large-scale linear systems

In this toolbox, we consider Multi-Input Multi-Output (MIMO), Linear Time-Invariant (LTI) systems either in discrete or continuous time. An LSS is composed by $s$ physically interconnected subsystems.

In the following, dependence of variables from time will be omitted, except where indicated. The dynamics of subsystem $i \in \mathcal{M} = 1 : s$ is

$$
x^+_{[i]} = A_{ii} x_{[i]} + B_{ii} u_{[i]} + \sum_{j \in \mathcal{N}_i} (A_{ij} x_{[j]} + B_{ij} u_{[j]}) \tag{2.1}
$$

where $x_{[i]} \in \mathbb{R}^{n_i}$ is the state at time $k$, $x^+_{[i]}$ is the state at time $k+1$ (for continuous-time systems, $x^+_{[i]}$ stands for $\frac{dx_{[i]}}{dt}$) and $u_{[i]} \in \mathbb{R}^{m_i}$ is the local input at time $k$. Moreover we have $A_{ii} \in \mathbb{R}^{n_i \times n_i}$, $A_{ij} \in \mathbb{R}^{n_i \times n_j}$, $B_{ii} \in \mathbb{R}^{n_i \times m_i}$ and $B_{ij} \in \mathbb{R}^{n_i \times m_j}$. Next, we define the index set $\mathcal{N}_i$ appearing in (2.1).

Definition 3. Subsystem $i$ is dynamically coupled to subsystem $j$ if $A_{ij} \neq 0$.

Definition 4. Subsystem $i$ is input coupled to subsystem $j$ if $B_{ij} \neq 0$.

It is possible to use graph theory to represent the coupling between the subsystems. The coupling graph of a system is directed graph where nodes are the subsystems, and the set of edges is given by $\xi = \{(i,j) : i \neq j, A_{ij} \neq 0 \text{ or } B_{ij} \neq 0\}$. An example is given in Figure 2.1.

Coupling allows us to define the predecessors of a subsystem.

Definition 5. The set of predecessors to subsystem $i$ is $\mathcal{N}_i = \{j : (i,j) \in \xi\}$. 
We further enhance our model by introducing centralized inputs $u$ from system $i$ to system $j$ means that $A_{ij} \neq 0$; a red arrow means $B_{ij} \neq 0$.

For example, from Figure 2.1 one has $\mathcal{N}_2 = \{1, 4\}$ since $A_{21}$, $B_{21}$ and $A_{24}$ are not null matrices. We also define the set of successors of subsystem $i$, i.e. the set of subsystems influenced by it.

**Definition 6.** The set of successors to system $i$ is $\mathcal{S}_i = \{j : (j,i) \in \xi\}$.

For example, from Figure 2.1 one has $\mathcal{S}_2 = \{4\}$ since $A_{42} \neq 0$.

The output of system $i$ is given by:

$$y_{[i]} = C_{ii}x_{[i]} + D_{ii}u_{[i]} + \sum_{j \in \mathcal{N}_i} (C_{ij}x_{[j]} + D_{ij}u_{[j]})$$

(2.2)

where $y_{[i]} \in \mathbb{R}^{p_i}$, $C_{ii} \in \mathbb{R}^{p_i \times n_i}$, $D_{ii} \in \mathbb{R}^{p_i \times m_i}$, $C_{ij} \in \mathbb{R}^{p_i \times n_j}$ and $D_{ij} \in \mathbb{R}^{p_i \times m_j}$.

**Definition 7.** Two systems $i$ and $j$ are output coupled if $C_{ij}$ or $D_{ij}$ are different from zero.

It is also possible to include exogenous signals that act on system $i$, by replacing (2.1) and (2.2) with:

$$x_{[i]} = A_{ii}x_{[i]} + B_{ii}u_{[i]} + \sum_{j \in \mathcal{N}_i} (A_{ij}x_{[j]} + B_{ij}u_{[j]}) + \sum_{j \in \mathcal{N}_i} M_{ij}d_{[j]}$$

(2.3)

$$y_{[i]} = C_{ii}x_{[i]} + D_{ii}u_{[i]} + \sum_{j \in \mathcal{N}_i} (C_{ij}x_{[j]} + D_{ij}u_{[j]}) + \sum_{j \in \mathcal{N}_i} N_{ij}d_{[j]}$$

(2.4)

where $d_{[j]} \in \mathbb{R}$ is an exogenous signal, $\mathcal{N}_{d,i} \subset \mathbb{N}$ is the set of exogenous signals that act on system $i$ and $M_{ij} \in \mathbb{R}^{n_i \times 1}$, $N_{ij} \in \mathbb{R}^{p_i \times 1}$.

We further enhance our model by introducing centralized inputs $u_{cen,[j]}$, so that the dynamics of subsystem $i$ becomes:

$$x_{[i]} = A_{ii}x_{[i]} + B_{ii}u_{[i]} + \sum_{j \in \mathcal{N}_i} (A_{ij}x_{[j]} + B_{ij}u_{[j]}) + \sum_{j \in \mathcal{N}_{u,i}} M_{ij}d_{[j]} + \sum_{j \in \mathcal{N}_{d,i}} B_{cen,ij}u_{cen,[j]}$$

(2.5)

$$y_{[i]} = C_{ii}x_{[i]} + D_{ii}u_{[i]} + \sum_{j \in \mathcal{N}_i} (C_{ij}x_{[j]} + D_{ij}u_{[j]}) + \sum_{j \in \mathcal{N}_{u,i}} N_{ij}d_{[j]} + \sum_{j \in \mathcal{N}_{d,i}} D_{cen,ij}u_{cen,[j]}$$

(2.6)

where $\mathcal{N}_{u,i} \subset \mathbb{N}$ is the set of centralized inputs $u_{cen,[j]}$, $j \in \mathbb{R}$ that act on system $i$ and $B_{cen,ij} \in \mathbb{R}^{n_i}$ and $D_{cen,ij} \in \mathbb{R}^{p_i}$. 

Figure 2.1: Example of a coupling graph of a system composed by four subsystems. A black arrow from system $i$ to system $j$ means that $A_{ij} \neq 0$; a red arrow means $B_{ij} \neq 0$. 
The difference between \( u[i] \) and \( u_{cen}[i] \) is that \( u[i] \) is a local input, i.e., the output of a local regulator for subsystem \( i \) while \( u_{cen}[i] \) is a global input that cannot be computed locally.

From models (2.5) and (2.6), the collective dynamics of the resulting LSS is

\[
x^+ = A + Bu + Md + B_{cen}u_{cen} \tag{2.7}
\]

\[
y = Cx + Du + Nd + D_{cen}u_{cen} \tag{2.8}
\]

where:

\[
x = (x[1], x[2], \ldots, x[s]) \in \mathbb{R}^n \tag{2.9}
\]

is the overall state and \( n = \sum_{i \in M} n_i \),

\[
u = (u[1], u[2], \ldots, u[s]) \in \mathbb{R}^m \tag{2.10}
\]

is the overall input and \( m = \sum_{i \in M} m_i \),

\[
y = (y[1], y[2], \ldots, y[s]) \in \mathbb{R}^p \tag{2.11}
\]

is the overall output and \( p = \sum_{i \in M} p_i \),

\[
d \in \mathbb{R}^{n_d} \tag{2.12}
\]

collects the exogenous signals acting on the overall system and

\[
u_{cen} \in \mathbb{R}^{n_u} \tag{2.13}
\]

collects the centralized inputs acting on the overall system.

Moreover one has:

\[
A = \begin{bmatrix}
A_{11} & \ldots & A_{1s} \\
\vdots & \ddots & \vdots \\
A_{s1} & \ldots & A_{ss}
\end{bmatrix} \in \mathbb{R}^{n \times n} \quad B = \begin{bmatrix}
B_{11} & \ldots & B_{1s} \\
\vdots & \ddots & \vdots \\
B_{s1} & \ldots & B_{ss}
\end{bmatrix} \in \mathbb{R}^{n \times m} \tag{2.14}
\]

All other matrices in (2.7) and (2.8) have a similar block structure. Moreover has \( C \in \mathbb{R}^{p \times n} \), \( D \in \mathbb{R}^{p \times m} \), \( M \in \mathbb{R}^{n \times n_d} \), \( N \in \mathbb{R}^{p \times n_d} \), \( B_{cen} \in \mathbb{R}^{n \times n_u} \) and \( D_{cen} \in \mathbb{R}^{p \times n_u} \).

The matrix \( A \) can also be split into matrices \( A_D \) and \( A_C \), defined as:

\[
A_D = \begin{bmatrix}
A_{11} & 0 & 0 \\
0 & \ddots & 0 \\
0 & 0 & A_{ss}
\end{bmatrix} \quad A_C = A - A_D \tag{2.15}
\]

Apparantly \( A_D \) collects all the local dynamics of the subsystems, while \( A_C \) collects coupling terms between subsystems. The same decomposition can be done for the matrices \( B \), \( C \), \( D \) appearing in (2.7) and (2.8).

Each subsystem can be equipped with the following constraints:

\[
x[i] \in X_i \subseteq \mathbb{R}^{n_i} \tag{2.16a}
\]

\[
u[i] \in U_i \subseteq \mathbb{R}^{m_i} \tag{2.16b}
\]

\[
y[i] \in Y_i \subseteq \mathbb{R}^{p_i} \tag{2.16c}
\]

We furthermore assume that \( X_i \), \( U_i \) and \( Y_i \) are polyhedra.
It is also possible to define coupling constraints between two or more subsystems. In the case of two subsystems \((i \text{ and } j)\) we have

\[(x_{[i]}, x_{[j]}) \in X_{ij} \subseteq \mathbb{R}^{n_i+n_j} \quad (2.17a)\]
\[(u_{[i]}, u_{[j]}) \in U_{ij} \subseteq \mathbb{R}^{m_i+m_j} \quad (2.17b)\]
\[(y_{[i]}, y_{[j]}) \in Y_{ij} \subseteq \mathbb{R}^{p_i+p_j} \quad (2.17c)\]

Constraints for the global lss system can be defined as follow

\[x \in X \subseteq \mathbb{R}^n \quad (2.18a)\]
\[u \in U \subseteq \mathbb{R}^m \quad (2.18b)\]
\[y \in Y \subseteq \mathbb{R}^p \quad (2.18c)\]

where

\[X = \left( \prod_{i \in \mathcal{M}} X_i \right) \cap \left( \bigcap_{i,j \in \mathcal{M}, i \neq j} X_{ij} \right) \quad (2.19a)\]
\[U = \left( \prod_{i \in \mathcal{M}} U_i \right) \cap \left( \bigcap_{i,j \in \mathcal{M}, i \neq j} U_{ij} \right) \quad (2.19b)\]
\[Y = \left( \prod_{i \in \mathcal{M}} Y_i \right) \cap \left( \bigcap_{i,j \in \mathcal{M}, i \neq j} Y_{ij} \right) \quad (2.19c)\]

Moreover we can define constraints for the exogenous signals and for the centralized inputs:

\[d \in D \subseteq \mathbb{R}^{n_d} \quad (2.20a)\]
\[u_{cen} \in U_{cen} \subseteq \mathbb{R}^{n_u} \quad (2.20b)\]

Recalling Definition 1 we have matrices \(H_x, H_u, H_d, H_{cen}\) and vectors \(K_x, K_u, K_d, K_{cen}\) of suitable dimensions such that

\[X = \{ x \in \mathbb{R}^n : H_x x \leq K_x \} \quad (2.21a)\]
\[U = \{ u \in \mathbb{R}^m : H_u u \leq K_u \} \quad (2.21b)\]
\[Y = \{ y \in \mathbb{R}^p : H_y y \leq K_y \} \quad (2.21c)\]
\[D = \{ d \in \mathbb{R}^{n_d} : H_d d \leq K_d \} \quad (2.21d)\]
\[U_{cen} = \{ u_{cen} \in \mathbb{R}^{n_u} : H_{cen} u_{cen} \leq K_{cen} \} \quad (2.21e)\]

Moreover, for each subsystem \(i \in \mathcal{M}\) there are matrices \(H_{x_i}, H_{u_i}, H_{y_i}\) and vectors \(K_{x_i}, K_{u_i}, K_{y_i}\) such that

\[X_i = \{ x_{[i]} \in \mathbb{R}^{n_i} : H_{x_i} x_{[i]} \leq K_{x_i} \} \quad (2.22a)\]
\[U_i = \{ u_{[i]} \in \mathbb{R}^{m_i} : H_{u_i} u_{[i]} \leq K_{u_i} \} \quad (2.22b)\]
\[Y_i = \{ y_{[i]} \in \mathbb{R}^{p_i} : H_{y_i} y_{[i]} \leq K_{y_i} \} \quad (2.22c)\]

In a similar way, for \(d_{[j]}, j \in \mathcal{J}\) we have

\[D_j = \{ d_{[j]} \in \mathbb{R} : H_{d,j} d_{[j]} \leq K_{d,j} \} \quad (2.23)\]

and for all \(u_{cen,[j]}, j \in \mathcal{J}\) it holds

\[U_{cen,j} = \{ u_{cen,[j]} \in \mathbb{R}^{n_u} : H_{cen,j} u_{cen,[j]} \leq K_{cen,j} \} \quad (2.24)\]

Coupling constraints have a totally analogous representation.

8
Chapter 3

Modeling of LSS: the lss and subss classes

3.1 The lss class

There are many MatLab toolboxes that offer facilities for modeling MIMO, LTI systems. However, most of them have not been designed to handle the interconnection of several subsystems. Therefore it is not immediate to manage an LSS, that means to add/remove subsystems, to extract a subsystem, to set coupling terms, etc. In the PnPMPC toolbox we have implemented methods for these tasks so as to ease the modeling of systems in the form (2.7) and (2.8). Moreover, the system object stores the constraints and some other useful pieces of information.

The properties of an lss object are:

- numSys: the number $s$ of subsystems composing the LSS;
- Ts: the sampling time;
- all matrices appearing in (2.7) and (2.8);
- $n_i$, $m_i$, $p_i$: vectors of $s$ elements which collect the numbers of states, inputs and outputs of every subsystem as in (2.9), (2.10) and (2.11);
- numExo, numICen are respectively $n_y$ in (2.12) and $n_u$ (2.13);
- all constraint matrices $H$ and $K$ appearing in (2.21);
- $H_{delta u}, K_{delta u}$ that will be explained in Section 3.2.4;
- coupling, name, that will be explained in Section 3.1;
- namex, nameu, namey, named, nameucen in order to assign names to variables (see Section 3.1).

Memory optimization

Taking into account that usually LSS have a sparse structure and therefore matrices in (2.7) and (2.8) have many zero elements, we decided to use of MatLab sparse matrices by default. This
usually allows one to save a considerable amount of memory. To visualize the matrices in the full format, is enough to use the MatLab command \texttt{full}.

To save further memory we also used the following trick. If in (2.8) one has $y = x$ then:

- the matrix $C$ is the identity matrix of order $n$
- $D$ is a matrix of zeros $\in \mathbb{R}^{n \times m}$
- there is no exogenous signal or centralized input acting on the output.

Therefore, we decided to not save these matrices. However if the user requires them with a \texttt{get} method, they will be generated upon demand.

In the following, this particular form of the output equation will be called the \textit{standard setting}.

\subsection*{Model discretization}

Continuous-time models can be discretized using the method \texttt{clss2dlss} (that means continuous LSS to discrete LSS), with this declaration:

$$
\texttt{dobjlss = objlss.clss2dlss(Ts, method)}
$$

where $Ts$ is the sampling time and \texttt{method} is the discretization technique. All the discretization techniques of the function \texttt{c2d} of the Control System Toolbox (which converts continuous-time systems to discrete-time models) are implemented, like \texttt{zoh} (zero order hold) or \texttt{Tustin} (see \texttt{help c2d} for more help). The \texttt{c2d} discretization methods are not always a good choice for LSS, because for exact discretization one has to compute the exponential of matrix $A$. Since $A$ is usually a large matrix with many zero elements, one has that:

- computing the exponential is time-consuming,
- elements that are zeros in the matrix $A$ of the continuous-time model can be different from zero in the exponential matrix. Therefore, exact discretization creates coupling between subsystems that were originally decoupled.

To avoid these problems we also implemented system-by-system discretization. That means that each subsystem in (2.5) and (2.6) is discretized considering states $x_{[j]}$, $j \neq i$ as exogenous inputs.

We can save computational time because we discretize sequentially subsystems that usually have low dimension and we do not create new couplings among subsystems. For using this technique, set \texttt{'subsystem'} as \texttt{method}.

\subsection*{Other utilities}

There are some other object’s properties that we have not explained yet.

- \texttt{coupling} is a matrix with size $\text{numSys} \times \text{numSys}$ composed by Boolean elements: element $(i, j)$ is 1 only if subsystem $i$ is dynamically coupled or input coupled to subsystem $j$, i.e. $(i, j) \in \xi$.
- \texttt{name} is a cell array of strings with size $1 \times \text{numSys}$, that associates a name to every subsystem in the lss object. It is possible to set name when calling the \texttt{addSystem} method.
If a name is not supplied to `addSystem`, a default name will be chosen and a warning appears. Every method that needs a subsystem as input, has been designed to use these names as an alternative to subsystem numbers. However, it is better to use numbers when possible, because MatLab is slower when working with strings.

- `namex`, `nameu`, `namey`, `named`, `nameucen` are cell arrays of strings, each with dimension $1 \times \text{numSys}$. The element in position $i$ stores information related to $i$-th subsystem. For example `namex(1,i)` is a matrix of strings of $n_i$ rows which stores the name associated to every state of $i$-th subsystem. Similar remarks can be applied to properties `nameu`, `namey`, `named`, `nameucen` that are related to system input, output, exogenous signals and centralized input. Names can be written with the LATEX syntax and are also used in plots of subsystems or time-evolution of individual variables.

### 3.2 How to use the lss class through examples

In the following sections, we provide several examples with comments that illustrate the modeling process with the PnPMPC toolbox. Section titles corresponds to m-files that can be found in the `./examples/lss` directory.

#### 3.2.1 example1lss.m

We show how to model the two coupled mass-spring-damper systems represented in Figure 3.1.

The model is

\[
\begin{bmatrix}
\dot{x}_1 \\
\dot{x}_2 
\end{bmatrix} = \begin{bmatrix}
A_{11} & A_{12} \\
A_{21} & A_{22}
\end{bmatrix} \begin{bmatrix}
x_1 \\
x_2
\end{bmatrix} + \begin{bmatrix}
u_1 \\
u_2
\end{bmatrix}
\]

(3.1)

where $x_{1,1}$ is composed by $x_{1,1}$, the displacement of first mass with respect to a given equilibrium position, and $x_{1,2}$, the velocity of the first mass. Similarly $x_{2,1}$ is composed by $x_{2,1}$ and $x_{2,2}$. Moreover $u_{1}$ and $u_{2}$ are forces applied to the first and second mass, respectively. We also have

\[
A = \begin{bmatrix}
A_{11} & A_{12} \\
A_{21} & A_{22}
\end{bmatrix}, \quad A_{11} = A_{22} = \begin{bmatrix}
0 & 1 \\
-\frac{k}{M} & -\frac{h}{M}
\end{bmatrix}, \quad A_{12} = A_{21} = \begin{bmatrix}
0 & 0 \\
\frac{h}{M} & \frac{h}{M}
\end{bmatrix}
\]

(3.2)

\[
B = \text{diag}(B_{11}, B_{22}), \quad B_{11} = B_{22} = \begin{bmatrix}
0 \\
\frac{100}{M}
\end{bmatrix}
\]

(3.3)

First of all we present the methods that will be used.
The lss method

`lss` is the constructor of the homonymous class, with this declaration

```plaintext
objlss = lss(A, B, ni, mi, C, D, pi, name, Ts)
```

where

- `A` and `B` are the matrices in (2.7);
- `ni` and `mi` are vectors storing the sizes of blocks `Aij` and `Bij` in matrices `A` and `B`;
- `C` and `D` are the matrices in (2.8);
- `pi` is a vector storing the sizes of blocks `Cij` and `Dij` in matrices `C` and `D`;
- `name` is an optional string defining the system name;
- `Ts` is the sampling time, that can be:
  - `Ts = []` for continuous-time system;
  - `Ts > 0` for discretized system with sampling time `Ts`;
  - `Ts = -1` for system already in the discrete-time form.

It is possible to introduce directly the matrices of the collective system system (2.7) and (2.8) but this operation is usually time consuming and error-prone. Hence in the example we will start from an empty object and add subsystems with `addSystem` method.

The addSystem method

The `addSystem` method adds to an `lss` object a single subsystem. The method declaration is

```plaintext
objlss = objlss.addSystem(Ai, Bi, Ci, Di, name, Ts)
```

where the subsystem matrices are `Ai`, `Bi`, `Ci`, `Di` as defined in (2.5) and (2.6). The arguments `name` and `Ts` have the same meaning as in the `lss` class. We also highlight that `addSystem` could also receive one input argument only: it can be a `subss` object (see Section 3.3) or a `ss` object (state-space object of System Control Toolbox) or a `tf` object (transfer function object of System Control Toolbox).

The addCoupling method

By using `addSystem` method only the block diagonal part of the matrices `A`, `B`, `C` and `D` in (2.7) and (2.8) can be set. If system `i` is coupled to system `j`, one or more blocks among `Aij`, `Bij`, `Cij` or `Dij` are non zero. The method `addCoupling` allows one to set them. The method declaration is

```plaintext
objlss = objlss.addCoupling(i, j, Aij, Bij, Cij, Dij)
```

where: `i` and `j` represent indexes of the the coupled subsystems. It is possible to use subsystem names instead of numbers `i` and `j`.

Code for modeling system (3.1)

With the three methods presented above it is possible create the model of coupled masses.
\[ M = 5; \quad \% \text{mass} \]
\[ k = 0.5; \quad \% \text{elastic constant of the springs} \]
\[ h = 5; \quad \% \text{damping coefficient} \]

\% creation of an empty lss object
modelCart = lss;

\[ A = \begin{bmatrix} 0 & 1; -k/M & -h/M \end{bmatrix}; \quad B = \begin{bmatrix} 0; 100/M \end{bmatrix}; \]

\% system 1
modelCart = modelCart.addSystem(A,B);

\% system 2
modelCart = modelCart.addSystem(A,B);

\% coupling among subsystems 1 and 2
\[ A_{ij} = \begin{bmatrix} 0 & 0; k/M & h/M \end{bmatrix}; \]
modelCart = modelCart.addCoupling(1,2,Aij);
modelCart = modelCart.addCoupling(2,1,Aij);

\% plot graph of subsystems
modelCart.plot

In general, the declaration of the methods contains many optimal parameters, that are optional and the user can provide only meaningful quantity. In the example above, parameters \( C_i, D_i, \text{name}, T_s \) are not supplied to addSystem method, because we are assuming \( y = x \), so we are in the “standard setting” discussed in Section 3.1. Moreover, we do not want to associate a name to the model and we are defining a model in continuous time, so that \( T_s \) is empty. Similar remarks apply to all methods of the toolbox.

### 3.2.2 example2lss.m

Now we will add an exogenous signal and a centralized input to the lss object modelCart through example1lss.m.

With reference to (2.7) and (2.8), we want to set

\[
M = \begin{bmatrix}
0 & 0 \\
1 & 0 \\
16 & 0 \\
2 & 3
\end{bmatrix}, \quad N = \begin{bmatrix}
0 & 0 \\
0 & 0 \\
0 & 0 \\
0 & 0
\end{bmatrix}, \quad B_{cen} = \begin{bmatrix}
0 \\
4 \\
0 \\
5
\end{bmatrix},
\]

(3.4)

Like in the previous section, we start with the description of the relevant methods.

**The addExoInput method**

The addExoInput method is used to set matrices \( M \) and \( N \).

The method declaration is

\[
\text{objlss} = \text{objlss.addExoInput}(\text{M,N},i,j)
\]
and it can be used in two ways:

- \( \text{objlss} = \text{objlss}.\text{addExoInput}(M,N) \), in this case we set the global \( M \in \mathbb{R}^{n \times nd} \) and \( N \in \mathbb{R}^{p \times nd} \) matrices of the LSS, according to (2.7) and (2.8).

- \( \text{objlss} = \text{objlss}.\text{addExoInput}(M,N,i,j) \), in this case we set only blocks \( M_{ij} \) and \( N_{ij} \) of matrices \( M \) and \( N \) (2.5) and (2.6), i.e. the relation among subsystem \( i \) and the exogenous signal \( j \), defined (2.5) and (2.6). The relation among all the other subsystems different from \( i \) and exogenous signal \( j \) is automatically set as zero if not already defined.

The addCentralizedInput method

The addCentralizedInput method is used to set matrices \( B_{cen} \) and \( D_{cen} \).

The method declaration is:

\[
\text{objlss} = \text{objlss}.\text{addCentralizedInput}(B_{cen},D_{cen},i,j)
\]

and it can be used in two ways:

- \( \text{objlss} = \text{objlss}.\text{addCentralizedInput}(B_{cen},D_{cen}) \), in this case we set the global \( B_{cen} \in \mathbb{R}^{n \times nu} \) and \( D_{cen} \in \mathbb{R}^{p \times nu} \) matrices of the LSS, according to (2.7) and (2.8).

- \( \text{objlss} = \text{objlss}.\text{addCentralizedInput}(B_{cen},D_{cen},i,j) \), in this case we set \( B_{cen,ij} \) and \( D_{cen,ij} \) of matrices \( B_{cen} \) and \( D_{cen} \) (2.5) and (2.6), i.e. the relation among subsystem \( i \) and the centralized input \( j \), defined (2.5) and (2.6). The relation among all the other subsystems different from \( i \) and centralized input \( j \) is automatically set as zero if not already defined.

```matlab
example1lss;
% specify how exogenous input number 1 affects system 1 (creation of M)
modelCart = modelCart.addExoInput ([0;1],[], 1,1);

% specify how exogenous input number 1 affects system 2 (creation of M)
modelCart = modelCart.addExoInput ([16;2],[],2,1);

% specify how exogenous input number 2 affects system 2 (creation of M)
modelCart = modelCart.addExoInput ([0;3],[],2,2);

% instead of calling addExoInput three times we can use
% modelCart = modelCart.addExoInput ([0 0; 1 0; 16 0; 2 3],[]);

% specify how centralized input number 1 affects system 1 (creation of Bcen)
modelCart = modelCart.addCentralizedInput ([0;4],[],1,1);

% specify how centralized input number 1 affects system 2 (creation of Bcen)
modelCart = modelCart.addCentralizedInput ([0;5],[],2,1);

% instead of the previous calls to addCentralizedInput we can use
% modelCart = modelCart.addCentralizedInput ([0;4;0;5],[]);
```
If, like in this case, the user does not need to set the \( N \) or \( Dcen \) matrix, it is possible to use empty matrices. Empty or not passed parameters are given default values. The same remark applies to all other methods in the toolbox.

### 3.2.3 example3lss.m

This example shows how to exploit modularity in the subsystem interconnection for quickly creating an LSS. The code below allows one to create a string of \( N \) mass-spring-damper systems (as shown in Figure 3.2) with different \( M \) (mass), different \( k \) (elastic constant of the springs) and different \( h \) (damping coefficient), verifying the bounds \( \min M \leq M \leq \max M \), \( \min k \leq k \leq \max k \) and \( \min h \leq h \leq \max h \). The system is characterized by the matrices:

![Large-scale system to model in example3lss.](image)

\( A_{i1} = \begin{bmatrix} 0 & 1 \\ -\frac{k(1)}{M(1)} & -\frac{h(1)}{M(1)} \end{bmatrix}, \quad A_{NN} = \begin{bmatrix} 0 & 1 \\ -\frac{k(N-1)}{M(N)} & -\frac{h(N-1)}{M(N)} \end{bmatrix}, \quad \forall i = 2 : N - 1 \)

\( A_{ii} = \begin{bmatrix} 0 & 1 \\ -\frac{k(i-1)+k(i)}{M(i)} & -\frac{h(i-1)+h(i)}{M(i)} \end{bmatrix}, \quad B_{ii} = \begin{bmatrix} 0 \\ \frac{1}{M(i)} \end{bmatrix} \)

\( A_{(i-1),i} = \begin{bmatrix} 0 & 1 \\ -\frac{k(i-1)}{M(i-1)} & -\frac{h(i-1)}{M(i-1)} \end{bmatrix}, \quad A_{i,(i-1)} = \begin{bmatrix} 0 & 1 \\ -\frac{k(i-1)}{M(i)} & -\frac{h(i-1)}{M(i)} \end{bmatrix} \)

\( C_{ii} = \begin{bmatrix} 1 & 0 \end{bmatrix}, \quad D_{ii} = 0; \)

The MatLab code for the model creation is when \( N = 1000 \) is

\[
N = 1000;
minM = 1;
maxM = 10;
mink = 0.1;
maxk = 0.9;
minh = 0.1;
maxh = 10;
\]

\[
M = \text{minM} + \text{rand}(1,N) * (\text{maxM} - \text{minM});
\]
k = mink + \text{rand}(i,N-1) * (maxk - mink);

h = minh + \text{rand}(1,N-1) * (maxh - minh);

modelCart = lss;
modelCart = modelCart.addSystem([0,1; -k(1)/M(1), -h(1)/M(1)],[0;100/M(1)],...
[1 0],0);
for i = 2:N-1
    modelCart = modelCart.addSystem([0,1; -(k(i-1)+k(i))/M(i),...
    -(h(i-1)+h(i))/M(i)],[0;100/M(i)],[1 0],0);
    modelCart = modelCart.addCoupling(i-1,i,[0,0;k(i-1)/M(i),...
    h(i-1)/M(i-1)],zeros(2,1));
    modelCart = modelCart.addCoupling(i,i-1,[0,0;k(i-1)/M(i),h(i-1)/M(i)],...
    zeros(2,1));
end
modelCart = modelCart.addSystem([0,1; -k(N-1)/M(N), -h(N-1)/M(N)],[0;100/M(N)],[1 0],0);
modelCart = modelCart.addCoupling(N,N-1,[0,0;k(N-1)/M(N),h(N-1)/M(N)],...
zeros(2,1));
modelCart = modelCart.addCoupling(N-1,N,[0,0;k(N-1)/M(N-1),h(N-1)/M(N-1)],...
zeros(2,1));

3.2.4 example4lss.m

This example shows how to add state and input constraints to an lss model. The addition of constraints on output, exogenous signal and centralized input can be done in a similar way. Relevant methods are described next.

Methods for adding constraints

The \text{addStateConstraint} method sets the matrices $H_x$ and $K_x$ in (2.22). The method declaration is:

\begin{verbatim}
objlss = objlss.addStateConstraint (sysindex,H,K)
\end{verbatim}

Where \text{sysindex} is the number or name of the subsystem to which constraints have to be added and \text{H} and \text{K} are the matrices $H_x$ and $K_x$ in (2.22). It is also possible to create constraints between two or more subsystems, in this case \text{sysindex} is a vector of scalars, or a cell array of names of the subsystems involved.

Methods that allow one to add constraints to inputs, outputs, exogenous signals and centralized inputs.

- \text{objlss = objlss.addInputConstraint(sysindex,H,K)}
- \text{objlss = objlss.addOutConstraint(sysindex,H,K)}. Note that if the user tries to insert a constraint on the output, but the system is in standard setting ($y = x$), the constraint will be converted in a state constraint and a warning will be displayed.
- \text{objlss = objlss.addExoConstraint(exoindex,H,K)}
- \text{objlss = objlss.addCentralizedInputConstraint(ceninputindex,H,K)
and works exactly as addStateConstraint.

The “double” methods

The double methods return matrices \( H \) and \( K \) related to constraints. The name “double” has been used for coherency with the MPT toolbox [3]. The method declaration is:

\[
[Hx,Kx] = objlss.doubleStateConstraint(index,selection)
\]

and it returns \( Hx \) and \( Kx \) matrix related to the states of the subsystem/subsystems expressed in \( index \). \( index \) can be a scalar or a string, but also a vector of scalars or a cell array of names, if one is interested in coupling constraints. \( selection \) is a Boolean flag with the following meaning:

- 0: exclusive (default). The state constraints only of the subsystem/subsystems indicated in \( index \) will be returned.
- 1: all. All state constraints which includes the subsystem/subsystems indicated in \( index \) will be returned.

For example, let \( c_1 \) be a constraint related to subsystem 1, \( c_3 \) be a constraint related to subsystem 3 and \( c_{13} \) be a coupling constraint between subsystem 1 and 3. If \( index = 3 \) and \( selection=0 \) (or not passed) \( H \) and \( K \) contain only \( c_3 \) because this is the only constraint that involves subsystem 3 only. If \( selection = 1 \), \( H \) and \( K \) will contain \( c_3 \) and \( c_{13} \), i.e. all constraints involving subsystem number 3. Another possible use of this method is:

\[
X = objlss.doubleStateConstraint(index,selection)
\]

with only one output argument. In this case, \( X \) is the polytope (i.e. an object of polytope class, see the documentation of the MPT toolbox for help) of system specified in \( index \) giving rise to a state constraint. Also if \( index \) contains more than one index, a polytope object will be returned.

Other double methods with similar meaning and use are:

- \([Hu,Ku] = objlss.doubleInputConstraint(index,selection)\)
- \([Hy,Ky] = objlss.doubleOutConstraint(index,selection)\)
- \([Hd,Kd] = objlss.doubleExoConstraint(index,selection)\)
- \([Hcen,Kcen] = objlss.doubleCenInputConstraint(index,selection)\)

Next, we provide the code for adding constraints (described in the code comments) to the model-Cart object.

```matlab
example2lss;
% 0(x_3)+1(x_4)\leq3
modelCart = modelCart.addStateConstraint (2,[0,1],3);
% 2(x_3)+1(x_4)\leq0 and 1(x_3)+3(x_4)\leq5
modelCart = modelCart.addStateConstraint (2,[2,1;1,3],[0;5]);
```
Input increment

In many systems, rather than constraining the input, it is preferable to bound the input increments \( \delta u_i = u_{i}(k+1) - u_{i}(k) \). With `addDeltaUConstraint` it is possible to define matrices \( H_{\delta u_i} \) and \( K_{\delta u_i} \) giving rise to the constraints \{\( H_{\delta u_i} \delta u_i \leq K_{\delta u_i} \}\).

The method declaration is

\[
\text{objlss} = \text{objlss}.\text{addDeltaUConstraint}(\text{sysindex}, H_{\delta u_i}, K_{\delta u_i})
\]

And the corresponding “double” method is

\[
[H_{\delta u_i}, K_{\delta u_i}] = \text{objlss}.\text{doubleDeltaUConstraint}(\text{index}, \text{selection})
\]

### 3.2.5 example5lss.m

All the methods described above are useful to add subsystems, couplings and constraints to an lss object. But if the user inserts wrong parameters it can be useful to remove system features in a selective way. This is the goal of the methods discussed next.

**The removeCoupling method**

\[
\text{objlss} = \text{objlss}.\text{removeCoupling}(i,j,\text{how})
\]

This method allows one to remove coupling between subsystems \( i \) and \( j \). The variable how can be 'a' or 'b' or 'c' or 'd' and specifies if we want to remove coupling terms in the matrices \( A, B, C, D \), respectively. If how is not passed blocks \((i,j)\) in all matrices \( A, B, C, D \) will be removed. See `help removeCoupling` for more information.

**The removeExoSignal method**

\[
\text{objlss} = \text{objlss}.\text{removeExoSignal}(i)
\]

This method allows one to remove the exogenous signal \( d_{i} \). It deletes the column \( i \) of matrices \( M \) and \( N \) and also the constraints related to this exogenous signals (if present). Note that a coupling
constraint related to exogenous signals $i$ and $j$, after the removal of the signal $i$ becomes a local constraint of signal $j$. For example, let $d_{[1]} + 2d_{[2]} \leq 10$ be a constraint among exogenous signals $d_{[1]}$ and $d_{[2]}$; the execution of $\text{objlss} = \text{objlss.removeExoSignal}(1)$ removes $d_{[1]}$ in the constraint, i.e. the new constraint is $2d_{[2]} \leq 10$. If one wants to remove all constraints where signal $d_{[i]}$ appears, the removeConstraint method can be used.

**The removeCentralizedInput method**

$$\text{objlss} = \text{objlss.removeCentralizedInput}(i)$$

This method allows one to deletes the centralized input number $i$. It eliminates the column $i$ of matrices $B_{cen}$ and $D_{cen}$ and also the constraints related to this centralized input (if present). Note that a coupling constraint related to centralized inputs $i$ and $j$, after the removal of $i$ becomes a local constraint of $j$. If one wants to remove all constraints where centralized input $u_{cen,[i]}$ appears, the removeConstraint method can be used.

**The removeConstraint method**

$$\text{objlss} = \text{objlss.removeConstraint}(\text{index}, \text{flag})$$

This method allows one to remove constraints. index can be, as usually, the subsystem number or name, if we want to remove local constraints. But it can also be a vector of scalars or a cell array of names, if we want to remove coupling constraints. flag can be `state' or `in' or `out' or `exo' or `cen' or `deltau'. In this way it is possible to remove, for instance, state constraints only. If flag is `exo' or `cen' we refer to the constraints to exogenous signals $d_{[i]}$ or centralized input, and in this case we have to specify the number of signal/signals interested. If flag is not specified all constraints on states, inputs and outputs of the system in index will be removed. Fore example if flag is empty and index is 1, then $X_1$, $U_1$ and $Y_1$ will be removed. See help removeConstraint for more information.

**The removeSystem method**

$$\text{objlss}=\text{objlss.removeSystem}(i)$$

This method allows one to remove the subsystem number $i$ from an lss object. Note that if subsystem $i$ is removed, this triggers the following changes.

- if an exogenous signal acts only on subsystem $i$ the method $\text{removeExoSignal}$ will be automatically called for removing the signal from the lss model and a warning appears on the screen. The same remark applies to centralized inputs.

- if two subsystems $i$ and $j$ are involved in a coupling constraint, and $i$ system is removed, the constraint becomes a local constraint of $j$. For example, let $u_{[1]} + u_{[2]} \leq 20$ be a constraint among subsystems 1 and 2; the execution of $\text{objlss} = \text{objlss.removeSystem}(1)$ removes $u_{[1]}$ in the constraint, i.e. the new constraint is $u_{[2]} \leq 20$.

In this example we illustrate the use of the removal methods. We start from the model created in example4lss.
example4lss;

  % remove exogenous signal number 2
  modelCart = modelCart.removeExoSignal(2);

  % remove centralized input number 1
  modelCart = modelCart.removeCentralizedInput(1);

  % remove coupling constraint on the state of system 1 and 2
  modelCart = modelCart.removeConstraint([1,2], 'state');

  % remove dynamic coupling between 1 and 2 so A12 = zeros
  modelCart = modelCart.removeCoupling(1,2, 'a');

  % remove subsystem number 2
  modelCart = modelCart.removeSystem(2);

3.2.6 Performances of the methods dedicated to modeling

We discuss here performances, in terms of memory occupation and execution time of the methods of the PnP-MPC toolbox dedicated to modeling. As a first case study, we use the string of N masses described in example1lss.m and example3lss.m with local constraints on states and inputs. In the left panel of Figure 3.3 one can see

- the execution time for creating the lss object (blue line),
- the execution time for discretizing the lss object using the system-by-system approach (red line)

as a function of the number of masses.

In the right panel of Figure 3.3 it is reported

- the memory occupation (in Kb) due to the continuous-time model (blue line),
- the memory occupation (in Kb) due to the discretized model (red line)

as a function of the number of masses.

These graphics have been obtained on a processor Intel Core i3 3.06 GHz, with 4GB of Ram 1.33 GHz running MatLab r2011a.

3.3 The subss class

This class allows one to model individual subsystems and it is useful for:

1. creating a subss object and pass it to an lss object with the addSystem method. In this way, dynamics and constraints of the subsystem will be added to the LSS.
2. extracting a subsystem from an lss object through the getSystem method.
Many methods of this class have the same name of methods of the lss class and work in the same way. In addition, many properties are the same as those of the lss class. Next we describe only the new ones referring the reader to the html documentation in the doc directory for a comprehensive description of all methods and properties. If we extract subsystem \(i\) from an lss object we lose coupling with other subsystems. Therefore we created the following properties to save:

- \(\text{couplingA, couplingB, couplingC, couplingD}\) are scalar vectors containing the indices of the subsystems in \(\mathcal{N}_i\) that are coupled with the extracted subsystem (i.e. subsystem \(i\)). \(\text{couplingA}\) will contain the indices of the subsystems coupled through the \(A\) matrix. \(\text{couplingB, couplingC and couplingD}\) have a similar meaning.

- \(A_{ij}, B_{ij}, C_{ij}, D_{ij}\). \(A_{ij}\) contains blocks \(A_{ij}\) where \(i\) is the number of the extracted subsystem and \(j \in \text{couplingA}\). Same remarks apply to the other properties.

### 3.3.1 example1subss.m

In this example we show how to add a subsystem to an lss object using a subss object.

```matlab
example2lss;
A = [0,1;-k/M,-h/M];
```
\[ B = \left[ \begin{array}{c} 0; 100/M \end{array} \right]; \]
\[ \text{sub} = \text{subss}(A,B); \]
\[ \text{sub} = \text{sub}.\text{addStateConstraint}([3,5;0,1],[1;2]); \]
\[ \text{modelCart} = \text{modelCart}.\text{addSystem}(	ext{sub}); \]

### 3.3.2 example2subss

After running example3lss, we can extract subsystem 237 as follows.

```matlab
example3lss;
sub = modelCart.getSystem(237)
```

With the previous command, the subsystem `sub` is an object of the class `subss`. In this case only local constraints are saved. For example constraints among subsystem 237 and 238 will not be saved. `sub` will have only the `couplingA` and `Aij` matrix, because the subsystem is dynamically coupled with 236 and 238, so

\[ \text{sub.couplingA} = [236 \ 238] \]
\[ \text{sub.Aij} = [A_{237,236}; A_{237,238}] \]

where \( A_{237,236} \) is the dynamic coupling among subsystems 237 and 236, and \( A_{237,238} \) is the dynamic coupling among subsystems 237 and 238.
Chapter 4

The pnpmpc class

4.1 Introduction

In this chapter, we introduce the pnpmpc class in order to design a PnPMPC controller for a single subsystem. PnPMPC controllers are described in [1].

In [1], the authors propose a PnP design procedure hinging on the notion of tube MPC [9] for handling coupling among subsystems, and aim at stabilizing the origin of the whole closed-loop system while guaranteeing satisfaction of constraints on local inputs and states. The model of subsystem $i$, $i \in M$ is given by (2.3) and (2.4) and we consider that each subsystem $i$ is input decoupled, i.e. $B_{ij} = 0$, $i \neq j$. Our control design procedure will hinge on the following assumptions.

**Assumption 1.** The matrix pairs $(A_{ii}, B_i)$ $\forall i \in M$ are controllable.

**Assumption 2.** Constraints $X_i$ and $U_i$, $i \in M$ are compact and convex polytopes containing the origin in their nonempty interior.

Let $Z_i$ be an RCI set for the nominal subsystem $i$ where the coupling term $\bigoplus_{j \in \mathcal{N}_i} A_{ij} X_j$ is treated as a disturbance. We design the following controller for subsystem $i$

$$C_i: u_i = v_i + \hat{r}_i(x_i - \hat{x}_i) \tag{4.1}$$

where $\hat{r}_i : Z_i \rightarrow U_i$ is any feedback control law guaranteeing $x_i(t) \in Z_i \Rightarrow x^+(t) \in Z_i$, $\forall x_{[i]} \in X_j$, $j \in \mathcal{N}_i$.

Following [9], we set in (4.1)

$$v_{[i]}(t) = v_{[i]}(0|t),$$

$$\hat{x}_{[i]}(t) = \hat{x}_{[i]}(0|t) \tag{4.2}$$

where $v_{[i]}(0|t)$ and $\hat{x}_{[i]}(0|t)$ are optimal values of the variables $v_{[i]}(0)$ and $\hat{x}_{[i]}(0)$, respectively.

---

**Robust Control Invariant (RCI) set**: Consider the discrete-time Linear Time-Invariant (LTI) system $x(t + 1) = Ax(t) + Bu(t) + w(t)$, with $x(t) \in \mathbb{R}^n$, $u(t) \in \mathbb{R}^m$, $w(t) \in \mathbb{W}$ and subject to constraints $u(t) \in \mathbb{U} \subseteq \mathbb{R}^m$ and $w(t) \in \mathbb{W} \subseteq \mathbb{R}^n$. The set $X \subseteq \mathbb{R}^n$ is an RCI set with respect to $w(t) \in \mathbb{W}$, if $\forall x(t) \in X$ then there exist $u(t) \in \mathbb{U}$ such that $x(t + 1) \in X$, $\forall w(t) \in \mathbb{W}$.

**Model of subsystem $i$ without coupling terms.**
appearing in the MPC-\(i\) problem
\[
P^N_i(x_{[i]}(t)) = \min_{\tilde{x}_{[i]}(0)} \sum_{k=0}^{N_i-1} \ell_i(\tilde{x}_{[i]}(k), v_{[i]}(k)) + V_{f_i}(\tilde{x}_{[i]}(N_i))
\]
(4.3a)
\[
x_{[i]}(t) - \tilde{x}_{[i]}(0) \in Z_i
\]
(4.3b)
\[
\dot{x}_{[i]}(k+1) = A_{ii}\tilde{x}_{[i]}(k) + B_i v_{[i]}(k), \quad k \in \{0 : N_i - 1\}
\]
(4.3c)
\[
\dot{\tilde{x}}_{[i]}(k) \in \tilde{X}_i, \quad k \in \{0 : N_i - 1\}
\]
(4.3d)
\[
v_{[i]}(k) \in V_i, \quad k \in \{0 : N_i - 1\}
\]
(4.3e)
\[
\dot{x}_{[i]}(N_i) \in \tilde{X}_f_i
\]
(4.3f)

In (4.3), \(N_i \in \mathbb{N}\) is the control horizon, \(\ell_i : \mathbb{R}^{n_i} \times m_i \rightarrow \mathbb{R}_+\) is the stage cost, \(V_{f_i} : \mathbb{R}^{n_i} \rightarrow \mathbb{R}_+\) is the final cost and \(\tilde{X}_f_i\) is the terminal set. Moreover, from (5.3c) and (5.3e), the tightened constraints \(\tilde{X}_i\) and \(V_i\) are defined respectively as
\[
\tilde{X}_i = X_i \odot Z_i, \quad V_i = U_i \odot \kappa_i(Z_i).
\]
(4.4)

Therefore, in order to design a PnPMPC controller for subsystem \(i\) we need to solve the following problem.

**Problem \(P_i\):** Compute nonempty RCI \(Z_i\) for the nominal subsystem \(i\) treating the coupling term as a disturbance. Compute sets \(\tilde{X}_i\) and \(V_i\).

We highlight that Problem \(P_i\) can be solved using efficient procedure proposed in [10] that requires the solution of a suitable LP problem. Moreover \(\tilde{X}_i\) and \(V_i\) can be computed using optimizers from the LP problem.

If Problem \(P_i\) cannot be solved, we declare that it is impossible to design a PnPMPC controller for subsystem \(i\).

The interested reader is refer to [1].

### 4.1.1 The pnpmpc method

Assume \texttt{subss} stores the model of subsystem \(i\) given by (2.3) and (2.4). Controller \(C_{[i]}\) is created through
\[
\text{objpnpmpc = pnpmpc (subss,N,Xj,k,options)}
\]
where \(N\) is the receding horizon, \(Xj\) is a cell array \(\{H_{xj}, K_{xj}\}\) such that neighbors states \(x_{[j]}\), \(j \in \mathcal{N}_i\) are in the set \(Xj = \{xj|H_{xj}xj \leq K_{xj}\}\). \(k\) is a parameter of Algorithm 1 in [1] and must be \(\geq\) of the controllability index of the pair \((A_{ii}, B_i)\). One can also set a cell array \(\{k_{min}, k_{max}\}\) in order to try different \(k\). \texttt{options} is used to set the \texttt{sdpsettings} for \texttt{yalmip} (see \texttt{yalmip manual} for details [4]).

We propose two methods to compute the terminal region and the terminal penalty.

### 4.1.2 The XFqpmax method

Using \texttt{XFqpmax} we compute a quadratic terminal region using procedures proposed in [11]. In particular we consider
\[
\ell_i(\tilde{x}_{[i]}(k), v_{[i]}(k)) = (\tilde{x}_{[i]}(k) - \tilde{z}_{[i]}^{ref}(k))^T Q(\tilde{x}_{[i]}(k) - \tilde{x}_{[i]}^{ref}(k)) + (v_{[i]}(k) - v_{[i]}^{ref}(k))^T R(v_{[i]}(k) - v_{[i]}^{ref}(k))
\]
where $Q = Q' \geq 0 \in \mathbb{R}^{n_i \times n_i}$, $R = R' > 0 \in \mathbb{R}^{m_i \times m_i}$ and $S = S' \geq 0 \in \mathbb{R}^{n_i \times n_i}$. \( \dot{x}_{i}^{r_{e}}(k) \) and \( v_{i}^{r_{e}}(k) \) are the state and input reference trajectories for tracking capabilities.

We can design the terminal penalty $S$ and the ellipsoidal terminal constraint executing the function

\[
\text{objpnpmpc = objpnpmpc.XFqpmax}(Q,R,\text{options})
\]

### 4.1.3 The zeroTerminal method

Using \text{zeroTerminal} we compute a zero terminal constraint as proposed in Chapter 2 in [12]. In particular we consider

\[
\ell_{i}(\tilde{x}_{i}^{r}(k),v_{i}^{r}(k)) = (\tilde{x}_{i}^{r}(k) - \dot{x}_{i}^{r_{e}}(k))^\prime Q(\tilde{x}_{i}^{r}(k) - \dot{x}_{i}^{r_{e}}(k)) + (v_{i}^{r}(k) - v_{i}^{r_{e}}(k))^\prime R(v_{i}^{r}(k) - v_{i}^{r_{e}}(k))
\]

\[
V_{f_{i}}(\tilde{x}_{i}(N_{i})) = 0
\]

\[
\hat{X}_{f_{i}} = \dot{x}_{i}^{r_{e}}(N_{i})
\]

where $Q = Q' \geq 0 \in \mathbb{R}^{n_i \times n_i}$ and $R = R' > 0 \in \mathbb{R}^{m_i \times m_i}$. \( \dot{x}_{i}^{r_{e}}(k) \) and \( v_{i}^{r_{e}}(k) \) are the state and input reference trajectories for tracking capabilities.

We can design the terminal penalty and the ellipsoidal terminal constraint executing the function

\[
\text{objpnpmpc = objpnpmpc.zeroTerminal}(Q,R)
\]

### 4.1.4 The uRH method

In order to compute the control action \( u_{i}^{r} \) we need to solve the optimization problem (4.3). The control action is computed executing the function

\[
[ \ u \ \text{diagnostics} \ ] = \text{objpnpmpc.uRH}(x0,din,x crefin,v crefin, options)
\]

- \( x0 \) is the initial state \( x_{i}^{r}(0) \), a vector of dimension \( n_i \times 1 \) for a decentralized implementation of \( PnPMPC \) controllers. For a distributed implementation (as in Section 5 of [1]), we consider \( x0 \) as a cell array \( x_{i}^{r}(0) \), \( \tilde{x}_{i}^{r}(0) \) where \( x_{i}^{r}(0) \) is the initial state for the current subsystem and \( x_{i}^{r}(0) \) is the state of predecessor subsystems.

- \( x crefin \) means the reference state of nominal system \( (\tilde{x}_{r_{e}}) \) over the control horizon; it can be expressed as a vector \( (n_i \times 1) \), if reference is constant over the control horizon) or a matrix \( (n_i \times N) \), the columns 1 refer to the values at the time instant 1 within the control horizon, columns 2 refer to the values at time 2, etc.). If this parameter is empty or not passed it is setted to zero by default.

- \( v crefin \) means the reference input of nominal system \( (v_{r_{e}}) \) over the control horizon; it can be expressed as a vector \( (m_i \times 1) \), if reference is constant over the control horizon) or a matrix \( (m_i \times N) \), the columns 1 refer to the values at the time instant 1 within the control horizon, columns 2 refer to the values at time 2, etc.). If this parameter is empty or not passed it is setted to zero by default.

- \( options \) \( \text{sdpsettings} \) object for \( \text{yalmip} \) options.
As output we have:

- $u$ control input value;
- *diagnostics* information about the optimization problems.

### 4.2 Design of a PnPMPC controller for a large-scale system

PnPMPC controllers $C_i$ for each subsystem in an lss object are created with the method `createCtrlPnPMPC` described next.

#### 4.2.1 The `createCtrlPnPMPC` method

This method acts on an lss object, and it is called as follows:

```matlab
ctrl = createCtrlPnPMPC(objlss,N,k,option)
```

where $N, \text{options}$ have the same meaning as in the `pnpmpc` method. The input $k$ is a cell array of two elements $\{k_{\text{min}}, k_{\text{max}}\}$: both $k_{\text{min}}$ and $k_{\text{max}}$ can be scalars and then they will be used for all execution of `pnpmpc` function for the $i$-subsystem or they can be vectors of dimension $1 \times \text{numSys}$, where $k_{\text{min}}(i)$ and $k_{\text{max}}(i)$ are used for the execution of `pnpmpc` function for the $i$-th subsystem.

This method extracts a subsystem, calls the method `pnpmpc` and creates the controller. The output $ctrl$ is an array of dimension $1 \times \text{numSys}$ and in the position $i$ it is stored the controller for subsystem $i$. After the creation of the controllers, we can run `XFqpsmax` or `zeroTerminal` methods, for setting terminal constraints, and then the `uRH` method, to compute the control inputs.

### 4.3 Examples

As an example of real application, in Chapter 5, we describe PnPMPC controllers for power network systems and run simulations using methods proposed in this chapter. In this section we propose simpler example.

#### 4.3.1 Example 1D

We consider a large-scale system composed by $N$ masses coupled as in Figure 4.1. Each mass $i \in \mathcal{M} = 1 : N$, is a subsystem with state variables $x_{[i]} = (x_{[i,1]}, x_{[i,2]})$ and input $u_{[i]} = u_{[i,1]}$.

![Figure 4.1: Array of masses in 1D.](image-url)
where $x_{i,1}$ is the displacements of mass $i$ with respect to a given equilibrium position, $x_{i,2}$ is the horizontal velocity of the mass $i$ and $100u_{i,1}$ is the force applied to mass $i$ in the horizontal direction. The values of $m_i$ have been extracted randomly in the interval $[5,10]$ while spring constants and damping coefficients are identical and equal to 0.5. Subsystems are equipped with the state constraints $||x_{i,1}||_{\infty} \leq 1.5$, $||x_{i,2}||_{\infty} \leq 0.8$, $i \in \mathcal{M}$ and with the input constraints $|u_{i,j}| \leq \beta_i$, where $\beta_i$ have been randomly chosen in the interval $[1,1.5]$. We obtain models $\Sigma_{i}^{\text{q}}$ by discretizing continuous-time models with 0.2 sec sampling time, using zero-order hold discretization for the local dynamics and treating $x_{j,i}$, $j \in \mathcal{N}_i$ as exogenous signals. At all time steps $t$, the control action $u_{i}(t)$ computed by the controller $C_i$, for all $i \in \mathcal{M}$, is kept constant during the sampling interval and applied to the continuous-time system.

In order to create the large-scale system composed by $N$ masses, we can use the function `makeModelTrucks1D` as follows

```matlab
N = 4;
[modelTrucks1Dc modelTrucks1Dd] = makeModelTrucks1D(N);
```

where $N$ is the number of trucks, `modelTrucks1Dc` and `modelTrucks1Dd` are the lss objects of the continuous-time and discrete-time models, respectively in form of lss objects.

We can design local controllers executing the function

```matlab
options = sdpsettings('verbose',0);
Trucks1Dpnp = makePnpmpcTrucks1D(modelTrucks1Dd,options);
```

whose instructions are described next.

The Controllers, $C_i$, $i \in \mathcal{M}$ are generated as follows. `Trucks1Dpnp` is an array of $N$ `pnpmpc` objects.

```matlab
NMPC = 20; \% prediction horizon
options = sdpsettings('verbose',0);
Trucks1Dpnp = modelTrucks1Dd.createCtrlPnPMPC(NMPC,\{1,30\},options);
```

Then we add a zero terminal constraint to each controller using the following instructions.

```matlab
Q = 10*eye(2); R = 1;
parfor i=1:modelTrucks1Dd.numSys
    Trucks1Dpnp(i) = Trucks1Dpnp(i).zeroTerminal(Q,R);
end
```

We note that the functions `makePnpmpcTrucks1D` or `modelTrucks1Dd.createCtrlPnPMPC` could take some minutes to be completely executed if $N$ is large. If you want take less time for the execution, we recommend to install CPLEX optimizer [13] that is free available for academic use.

To start a simulation, we can use the function `runSimTrucks1D` in the following way

```matlab
options = sdpsettings('verbose',0,'solver','sdpt3');
x0 = repmat([1 0]',N,1);`
This function computes the control action using the method $uRH$ of the pnpmpc class. We used sdpt3 solver, that is installed with the toolbox, but we really recommend the use of CPLEX.

4.3.2 Example 2D

We consider a large-scale system composed by $N$ masses coupled as in Figure 4.3.2 (for the case of $N = 1024$) where the four edges connected to a point correspond to springs and dampers arranged as in Figure 4.2. Hereafter we assume that $N = 2^z$ for some $z \in \mathbb{N}, z > 0$. Each mass $i \in \mathcal{M} = 1 : N$, is a subsystem with state variables $x_{[i]} = (x_{[i,1]}, x_{[i,2]}, x_{[i,3]}, x_{[i,4]})$ and input $u_{[i]} = (u_{[i,1]}, u_{[i,2]})$, where $x_{[i,1]}$ and $x_{[i,3]}$ are the displacements of mass $i$ with respect to a given equilibrium position on the plane (equilibria lie on a regular grid), $x_{[i,2]}$ and $x_{[i,4]}$ are the horizontal and vertical velocity of the mass $i$, respectively, and $100u_{[i,1]}$ (respectively $100u_{[i,2]}$) is the force applied to mass $i$ in the horizontal (respectively, vertical) direction. The values of $m_i$ have been extracted randomly in the interval $[5, 10]$ while spring constants and damping coefficients are identical and equal to 0.5. Subsystems are equipped with the state constraints $\|x_{[i,j]}\|_\infty \leq 1.5, \ j = 1,3, \ |x_{[i,4]}|_\infty \leq 0.8, \ i \in \mathcal{M}, \ l = 2,4$ and with the input constraints $\|u_{[i]}\|_\infty \leq \beta_i$, where $\beta_i$ have been randomly chosen in the interval $[1,1.5]$. We obtain models $\Sigma_{[i]}$ by discretizing continuous-time models with 0.2 sec sampling time, using zero-order hold discretization for the local dynamics and treating $x_{[j]}, \ j \in \mathcal{N}_i$ as exogenous signals. At all time steps $t$, the control action $u_{[i]}(t)$ computed by the controller $C_i$, for all $i \in \mathcal{M}$, is kept constant during the sampling interval and applied to the continuous-time system.
In order to create the large-scale system composed by $N$ masses, we can use the function `makeModelTrucks2D` as follows:

```matlab
N = 4;
[modelTrucks2Dc modelTrucks2Dd] = makeModelTrucks2D(N);
```

where $N$ is the number of trucks, `modelTrucks2Dc` and `modelTrucks2Dd` are the continuous-time and discrete-time large-scale models, respectively in form of lss objects.

We can design local controllers executing the function:

```matlab
options = sdpsettings('verbose',0);
Trucks2Dpnp = makePnpmpcTrucks2D(modelTrucks2Dd,options);
```

whose instructions are described next.

The Controllers $C_i$, $i \in \mathcal{M}$ are generated as follows. `Trucks2Dpnp` is an array of $N$ `pnpmpc` objects.

```matlab
NMPC = 20; % prediction horizon
options = sdpsettings('verbose',0);
Trucks2Dpnp = modelTrucks2Dd.createCtrlPnPMPC(NMPC,[2,30],options);
```

Then, we add a zero terminal constraint to each controller using the following instructions:

```matlab
Q = 10*eye(4); R = eye(2);
parfor i=1: modelTrucks2Dd.numSys
    Trucks2Dpnp(i) = Trucks2Dpnp(i).zeroTerminal(Q,R);
end
```
We note that the functions `makePnpmpcTrucks2D` or `modelTrucks2Dd.createCtrlPnPMPC` could take some minutes to be completely executed. If you want take less time for the execution, we recommend to install CPLEX optimizer [13] that is free available for academic use.

To start a simulation, we can use the function `runSimTrucks2D` in the following way

```matlab
options = sdpsettings('verbose',0);
x0 = repmat([1 0 -1 0]',N,1);
Tsim = 20;
[x u] = runSimTrucks2D(Tsim,x0,modelTrucks2Dc,modelTrucks2Dd,Trucks2Dpnp,options);
```

This function computes the control action using the method `uRH` of the pnpmpc class. We did not specify a solver in order to be as general as possible. We really recommend the use of CPLEX.

In [1], we propose the case of $N = 1024$ trucks and we give the results in terms of computational times.
Chapter 5

Hycon2 Benchmark: Power Network System

Note: for MatLab simulations in this chapter, we recommend the use of CPLEX optimizer [13].

5.1 Introduction

An example of a real application that can benefit of decentralized and distributed control schemes is the regulation of a Power Network System (PNS). Here we describe the PNS proposed as a benchmark exercise [1] within the HYCON2 project [14].

We consider a PNS as composed by several power generation areas coupled through tie-lines [15]. The aim is to design the Automatic Generation Control (AGC) layer for frequency control with the goal of:

- keeping the frequency approximately at the nominal value;
- controlling the tie-line powers in order to reduce power exchanges between areas. In the asymptotic regime each area should compensate for local load steps and produce the required power.

We consider thermal power stations with single-stage turbines. The dynamics of an area equipped with primary control and linearized around equilibrium value for all variables can be described by the following continuous-time LTI model [15]

\[
\Sigma^C_{[i]}: \quad \dot{x}_{[i]} = A_{ii} x_{[i]} + B_i u_{[i]} + L_i \Delta P_L + \sum_{j \in \mathcal{N}_i} A_{ij} x_{[j]} \quad (5.1)
\]

where \(x_{[i]} = (\Delta \theta_i, \Delta \omega_i, \Delta P_m, \Delta P_v)\) is the state, \(u_{[i]} = \Delta P_{ref_i}\) is the control input of each area, \(\Delta P_L\) is the local power load and \(\mathcal{N}_i\) is the sets of neighboring areas, i.e. areas directly connected.
to $\Sigma^C_{[i]}$ through tie-lines. The matrices of system (5.1) are defined as
\[
A_{ii}(\{P_{ij}\}_{j \in N_i}) = \begin{bmatrix}
0 & 1 & 0 & 0 \\
-\sum_{j \in N_i} P_{ij} & -\frac{D_i}{2H_i} & -\frac{2H_i}{T_{ti}} & 0 \\
0 & 0 & -\frac{1}{R_i T_{ti}} & -\frac{1}{T_{gi}} \\
0 & 0 & 0 & 0 \\
\end{bmatrix}
B_i = \begin{bmatrix}
0 \\
0 \\
0 \\
1 \\
\end{bmatrix}
\]
\[
A_{ij} = \begin{bmatrix}
P_{ij} & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
\end{bmatrix}
L_i = \begin{bmatrix}
0 \\
0 \\
0 \\
0 \\
\end{bmatrix}
(5.2)
\]

For the meaning of constants as well as some typical parameter values we refer the reader to Table 5.1.

| \(\Delta \theta_i\) | Deviation of the angular displacement of the rotor with respect to the stationary reference axis on the stator |
| \(\Delta \omega_i\) | Speed deviation of rotating mass from nominal value |
| \(\Delta P_{m_i}\) | Deviation of the mechanical power from nominal value (p.u.) |
| \(\Delta P_{v_i}\) | Deviation of the steam valve position from nominal value (p.u.) |
| \(\Delta P_{ref_i}\) | Deviation of the reference set power from nominal value (p.u.) |
| \(\Delta P_{L_i}\) | Deviation of the nonfrequency-sensitive load change from nominal value (p.u.) |
| \(H_i\) | Inertia constant defined as \(H_i = \frac{\text{kinetic energy at rated speed}}{\text{machine rating}}\) (typically values in range \([1 - 10]\) sec) |
| \(R_i\) | Speed regulation |
| \(D_i\) | Defined as \(\%\) change in load change in frequency |
| \(T_{ti}\) | Prime mover time constant (typically values in range \([0.2 - 2]\) sec) |
| \(T_{gi}\) | Governor time constant (typically values in range \([0.1 - 0.6]\) sec) |
| \(P_{ij}\) | Slope of the power angle curve at the initial operating angle between area \(i\) and area \(j\) |

Table 5.1: Variables of a generation area with typical value ranges [15]. (p.u.) stands for “per unit”.

We note that model (5.1) is input decoupled since both \(\Delta P_{ref_i}\) and \(\Delta P_{L_i}\) act only on subsystem \(\Sigma^C_{[i]}\). Moreover, subsystems \(\Sigma^C_{[i]}\) are parameter dependent since the local dynamics depends on the quantities \(-\sum_{j \in N_i} P_{ij}\).

In the following we introduce three scenarios corresponding to different interconnection topologies of generation areas. The model parameters and constraints on \(\Delta \theta_i\) and on \(\Delta P_{ref_i}\) for systems in all Scenarios are given in Table 5.2. We highlight that all parameter values are within the range of those used in Chapter 12 of [15]. We define \(M\) as the number of areas in the power network. For each scenario, discrete-time models \(\Sigma_{[i]}\) with \(T_s = 1\) sec sampling time are obtained from \(\Sigma^C_{[i]}\) using two alternative discretization schemes.

- Exact discretization of the overall system (acronym \(D\));
- Discretization system-by-system, i.e. exact discretization for each area treating \(u_{[i]}\), \(\Delta P_{L_i}\) and \(x_{[j]}, j \in N_i\) as exogenous inputs (acronym \(Dss\)).

In particular, we note that \(Dss\) preserves the input-decoupled structure of \(\Sigma^C_{[i]}\) while \(D\) does not.

### 5.1.1 Scenario 1

We consider four areas interconnected as in Figure 5.1. We will simulate Scenario 1 using the load steps specified in Table 5.3.
### Table 5.2: Model parameters and constraints for systems $\Sigma_i$, $i \in 1, \ldots, 5.$

<table>
<thead>
<tr>
<th>$\Delta \theta_i$</th>
<th>$|x_{[1,1]}|_\infty \leq 0.1$</th>
<th>$|x_{[2,1]}|_\infty \leq 0.1$</th>
<th>$|x_{[3,1]}|_\infty \leq 0.1$</th>
<th>$|x_{[4,1]}|_\infty \leq 0.1$</th>
<th>$|x_{[5,1]}|_\infty \leq 0.1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta P_{ref,i}$</td>
<td>$|u_{[1]}|_\infty \leq 0.5$</td>
<td>$|u_{[2]}|_\infty \leq 0.65$</td>
<td>$|u_{[3]}|_\infty \leq 0.65$</td>
<td>$|u_{[4]}|_\infty \leq 0.55$</td>
<td>$|u_{[5]}|_\infty \leq 0.5$</td>
</tr>
</tbody>
</table>

\[
P_{12} = 4 \quad P_{23} = 2 \quad P_{34} = 2 \quad P_{45} = 3 \quad P_{25} = 3
\]

Figure 5.1: Power network system of Scenario 1

Table 5.3: Load of power $\Delta P_{L,i}$ (p.u.) for simulation in Scenario 1. $+\Delta P_{L,i}$ means a step of required power, hence a decrease of the frequency deviation $\Delta \omega_i$ and therefore an increase of the power reference $\Delta P_{ref,i}$.

<table>
<thead>
<tr>
<th>Step time</th>
<th>Area $i$</th>
<th>$\Delta P_{L,i}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>1</td>
<td>+0.15</td>
</tr>
<tr>
<td>15</td>
<td>2</td>
<td>-0.15</td>
</tr>
<tr>
<td>20</td>
<td>3</td>
<td>+0.12</td>
</tr>
<tr>
<td>40</td>
<td>3</td>
<td>-0.12</td>
</tr>
<tr>
<td>40</td>
<td>4</td>
<td>+0.28</td>
</tr>
</tbody>
</table>

5.1.2 Scenario 2

We consider the power network proposed in Scenario 1 and add a fifth area connected as in Figure 5.2. We will simulate Scenario 2 using the load steps specified in Table 5.4.
5.1.3 Scenario 3

We consider the power network described in Scenario 2 and disconnect the area 4, hence obtaining the areas connected as in Figure 5.3. We will simulate Scenario 3 using load steps specified in Table 5.5.

Table 5.4: Load of power $\Delta P_{L_i}$ (p.u.) for simulation in Scenario 2. $+\Delta P_{L_i}$ means a step of required power, hence a decrease of the frequency deviation $\Delta \omega_i$ and therefore an increase of the power reference $\Delta P_{ref_i}$.
Table 5.5: Load of power $\Delta P_{L_i}$ (p.u.) for simulation in Scenario 3. $+\Delta P_{L_i}$ means a step of required power, hence a decrease of the frequency deviation $\Delta \omega_i$ and therefore an increase of the power reference $\Delta P_{ref_i}$.

We can create lss objects for each scenario, running `makeScenariosPNS`. For the first scenario we create 3 files: `pnsC1` (continuous-time lss object), `pnsD1` (discrete-time lss object), `pnsD1ss` (discrete-time system-by-system lss object). Similar files are created for scenario 2 and 3.

### 5.2 Design of the AGC layer for a power network using MPC

The goal of the Benchmark is to design the AGC layer for the scenarios introduced in Section 5.1. Different control schemes will be compared with the centralized MPC scheme described next. For a given Scenario, at time $t$ we solve the centralized optimization problem

$$
P_N^\pi(x(t)) : \min_{u(t:t+N-1)} \sum_{k=t}^{t+N-1} (||x(k) - x^O||_Q + ||u(k) - u^O||_R) + ||x(t+N) - x^O||_S)$$

(5.3a)

$$\begin{align}
x(k+1) &= Ax(k) + Bu(k) + L\Delta P_{L_i}(t) & k \in 0 : N - 1 \\
x(k) &\in X & k \in 0 : N - 1 \\
u(k) &\in U & k \in 0 : N - 1 \\
x(N) &\in X_f
\end{align}$$

(5.3b-5.3f)

and then apply $\Delta P_{ref_i} = u(0)$. We note that the cost function depend upon $x^O$ and $u^O$ that are defined as $x_{i^O}^O = (0, 0, \Delta P_{L_i}, \Delta P_{L_j})$ and $u_{i^O}^O = \Delta P_{L_i}$. The constraints $X$ and $U$ in (5.3d) and (5.3e) are obtained from constraints listed in Table 5.2. In the cost function (5.3b) we set $N = 15$, $Q = \text{diag}(Q_1, \ldots, Q_M)$ and $R = \text{diag}(R_1, \ldots, R_M)$, where

$$Q_i = \begin{bmatrix}500 & 0 & 0 & 0 \\ 0 & 0.01 & 0 & 0 \\ 0 & 0 & 0.01 & 0 \\ 0 & 0 & 0 & 10\end{bmatrix}$$

and $R_i = 10$.

Weights $Q_i$ and $R_i$ have been chosen in order to penalize the angular displacement $\Delta \theta_i$ and to penalize slow reactions to power load steps. Since the power transfer between areas $i$ and $j$ is
given by
\[ \Delta P_{\text{tie}ij}(k) = P_{ij}(\Delta \theta_i(k) - \Delta \theta_j(k)) \] (5.4)
the first requirement also penalizes huge power transfers.

In order to guarantee the stability of the closed loop system, we design the matrix \( S \) and the terminal constraint set \( \mathcal{X}_f \) in three different ways.

- **S is full (MPCfull):** we compute the symmetric positive-definite matrix \( S \) and the static state-feedback auxiliary control law \( K_{\text{aux}}x \), by maximizing the volume of the ellipsoid described by \( S \) inside the state constraints while fulfilling the matrix inequality \((A + BK_{\text{aux}})'S(A + BK_{\text{aux}}) - S \leq -Q - K_{\text{aux}}'RK_{\text{aux}}.\)

- **S is block diagonal (MPCdiag):** we compute the decentralized symmetric positive-definite matrix \( S \) and the decentralized static state-feedback auxiliary control law \( K_{\text{aux}} = \text{diag}(K_1, \ldots, K_M) \) by maximizing the volume of the ellipsoid described by \( S \) inside the state constraints while fulfilling the matrix inequality \((A + BK_{\text{aux}})'S(A + BK_{\text{aux}}) - S \leq -Q - K_{\text{aux}}'RK_{\text{aux}}.\)

- **Zero terminal constraint (MPCzero):** we set \( S = 0 \) and \( \mathcal{X}_f = x^O. \)

### 5.2.1 Performance criteria

We propose the following performance criteria for evaluating different control schemes.

- **\( \eta \)-index**

  \[ \eta = \frac{1}{T_{\text{sim}}} \sum_{k=0}^{T_{\text{sim}}-1} \sum_{i=1}^M (||x_i(k) - x^O_i(k)||_{Q_i} + ||u_i(k) - u^O_i(k)||_{R_i}) \] (5.5)

  where \( T_{\text{sim}} \) is the time of the simulation. From (5.5), \( \eta \) is a weighted average of the error between the real state and the equilibrium state and between the real input and the equilibrium input.

- **\( \Phi \)-index**

  \[ \Phi = \frac{1}{T_{\text{sim}}} \sum_{k=0}^{T_{\text{sim}}-1} \sum_{i=1}^M \sum_{j \in \mathcal{N}_i} |\Delta P_{\text{tie}ij}(k)|T_s \] (5.6)

  where \( T_{\text{sim}} \) is the time of the simulation and \( \Delta P_{\text{tie}ij} \) is the power transfer between areas \( i \) and \( j \) defined in (5.4). This index gives the average power transferred between areas. In particular, if the \( \eta \)-index is equal for two regulators, the best controller is the one that has the lower value of \( \Phi \).

### 5.3 Control Experiments

We applied the centralized MPC schemes introduced in the previous section to scenarios 1, 2 and 3. Furthermore, for each scenario we discretized the continuos system with both discretization schemes \( D \) and \( D_{ss} \). At time \( t \) we solve the optimization problem (5.3) and then apply the control action to the continuos-time system, keeping the value constant between time \( t \) and \( t + 1 \). If at time \( t \) the power load increases or decreases, we assume the controller can use this information at time \( t \). This means at time \( t \) the controller knows exactly the value of \( \Delta P_L \) hence can use
it. We highlight that violation of this assumption can impact considerably on the index $\eta$. In all experiments we use $T_{\text{sim}} = 100$. In Table 5.8 and 5.9 the values of the performance parameters $\eta$ and $\Phi$, respectively, are reported for each control experiment.

<table>
<thead>
<tr>
<th>Scenario</th>
<th>$D$</th>
<th>$D_{ss}$</th>
<th>$D$</th>
<th>$D_{ss}$</th>
<th>$D$</th>
<th>$D_{ss}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$MPC_{\text{full}}$</td>
<td>0.0249</td>
<td>0.0249</td>
<td>0.0346</td>
<td>0.0347</td>
<td>0.0510</td>
<td>0.0511</td>
</tr>
<tr>
<td>$MPC_{\text{diag}}$</td>
<td>0.0249</td>
<td>0.0249</td>
<td>0.0346</td>
<td>0.0347</td>
<td>0.0510</td>
<td>0.0511</td>
</tr>
<tr>
<td>$MPC_{\text{zero}}$</td>
<td>0.0249</td>
<td>0.0249</td>
<td>0.0346</td>
<td>0.0347</td>
<td>0.0510</td>
<td>0.0511</td>
</tr>
</tbody>
</table>

Table 5.6: Values of the performance parameter $\eta$ using different centralized MPC schemes for the AGC layer.

<table>
<thead>
<tr>
<th>Scenario</th>
<th>$D$</th>
<th>$D_{ss}$</th>
<th>$D$</th>
<th>$D_{ss}$</th>
<th>$D$</th>
<th>$D_{ss}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$MPC_{\text{full}}$</td>
<td>0.0030</td>
<td>0.0029</td>
<td>0.0063</td>
<td>0.0060</td>
<td>0.0060</td>
<td>0.0058</td>
</tr>
<tr>
<td>$MPC_{\text{diag}}$</td>
<td>0.0030</td>
<td>0.0029</td>
<td>0.0063</td>
<td>0.0061</td>
<td>0.0060</td>
<td>0.0058</td>
</tr>
<tr>
<td>$MPC_{\text{zero}}$</td>
<td>0.0030</td>
<td>0.0028</td>
<td>0.0063</td>
<td>0.0059</td>
<td>0.0059</td>
<td>0.0058</td>
</tr>
</tbody>
</table>

Table 5.7: Values of the performance parameter $\Phi$ using different centralized MPC schemes for the AGC layer.

### 5.4 Supporting MatLab files

In terms of PnP-MPC controllers, running file `makePnpmpcControllersPNS`, we can create controllers with different terminal regions. Then we can run each simulation, executing file `scenario[number scenario][Full—Zero][De—Di]`. For example if we want run the simulation for Scenario 2 using PnP-MPC controllers in a distributed fashion with zero terminal constraints, we should run

```
scenario2ZeroDi (options)
```

then a file `scenario2ZeroDiData` is created with all results of the simulation. We note that using PnP-MPC controllers we refer to `full` when using a quadratic terminal region for controller $C_i$, $i \in \mathcal{M}$. Moreover we consider discretization system-by-system only. `options` is a `sdpsettings` object for `yalmip`. We highlight that different performances can be achieved using different solvers. We also highlight that each simulation could take some minutes to be completely executed. If you want take less time for the execution without numerical errors, we recommend to install CPLEX optimizer [13] that is free available for academic use. We note that most of the functions can be used in a parallel fashion, then the interested user can run `MatLabpool` before the simulations. For our simulations results, we refer the interested reader to [1]. One can also execute all files for modeling and designed PnP-MPC controllers running `makeAllPNSfiles` and can delete all files for PnP-MPC controllers running `clearAllPNSfiles`. 37
For each control experiment we provide a file .mat of the simulation that contains

- lss object of the continuous linear system (pnsCn, where n is the number of the scenario);
- parameters of the control experiment T_{sim} and deltaPL, where deltaPL corresponds to \Delta P_L;
- the results of the control experiment x, deltaPref, \eta and \Phi, where deltaPref corresponds to \Delta P_{ref}.

For each Scenario we included also a Simulink model. In particular, one can load the file .mat of a control experiment and simulate the power network system given the power load steps and the power reference computed through centralized MPC or PnPMPC controllers.

5.4.1 Example of simulation

In the following we illustrate how to use the files .mat and the Simulink models for designing centralized and pnpmpc controllers.

- **Step 1** We can simulate different scenarios using the Simulink models present in the folder of each scenario. For Scenario 2 we then open the file simulatorPNS\_AGC\_2.mdl. This step is performed with the MatLab command:

  ```matlab
  open('simulatorPNS\_AGC\_2')
  ```

- **Step 2**

  **Centralized case** In the subfolders of each scenario there are MatLab files for centralized simulations as scenario[number scenario][D—Dss][Diag—Full—Zero]/Data. Assume we want to simulate Scenario 2 using the discretization Dss and centralized MPC with zero terminal constraint (MPCzero). We need to load MatLab file scenario2DssZeroData. This operation can be performed with the MatLab command:

  ```matlab
  load scenario2DssZeroData
  ```

  **PnPMPC case** In the subfolders of each scenario there are MatLab files for PnPMPC simulations as scenario[number scenario][Full—Zero][De—Di]/Data. Assume we want to simulate Scenario 2 in a distributed fashion with zero terminal constraint. We need to load MatLab file scenario2ZeroDiData. This operation can be performed with the MatLab command:

  ```matlab
  load scenario2ZeroDiData
  ```

- **Step 3** Start a simulation from Simulink will produce the results of the control experiments.
5.5 Summary of results

In this section, we summarize the simulation results using centralized and plug-and-play MPC.

In Figure 5.4 we compare the performance of the PnP scheme with the performance of the centralized MPC controller for Scenario 1. In the control experiment, step power loads $\Delta P_L_i$ are specified in Table 5.3 and they account for the step-like changes of the control variables in Figure 5.4. We highlight that the performance of decentralized and centralized MPC are totally comparable, in terms of frequency deviation (Figure 5.4(a)), control variables (Figure 5.4(b)) and power transfers $\Delta P_{tie}_{ij}$ (Figure 5.5). The values of performance parameter $\eta$ and $\Phi$ using different controllers are reported in Table 5.8 and Table 5.9, respectively. In terms of parameter $\eta$, plug-and-play controllers with decentralized and distributed online implementation are equivalent to centralized controller, however the performance of PnP-DeMPC are such that each area can absorb the local loads by producing more power locally ($\Delta P_{ref,i}$) instead of receiving power from predecessor areas: for this reason, PnP-DiMPC has performance more similar to centralized MPC.

The values of performance parameter $\eta$ and $\Phi$ using different controllers for Scenario 2 and 3 are reported in Table 5.8 and Table 5.9, respectively. We refer the interested reader to [1] for an in-depth analysis of the results for all scenarios.

<table>
<thead>
<tr>
<th>Scenario 1</th>
<th>Scenario 2</th>
<th>Scenario 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cen-MPC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.0249</td>
<td>0.0249</td>
<td>0.0346</td>
</tr>
<tr>
<td>De-PnPMPC</td>
<td>+2.81%</td>
<td>+3.61%</td>
</tr>
<tr>
<td>Di-PnPMPC</td>
<td>+3.61%</td>
<td>+2.15%</td>
</tr>
</tbody>
</table>

Table 5.8: Value of the performance parameter $\eta$ for centralized MPC (first line) and percentage change using decentralized and distributed MPC schemes for the AGC layer. Best values for PnP controllers are in bold.

<table>
<thead>
<tr>
<th>Scenario 1</th>
<th>Scenario 2</th>
<th>Scenario 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cen-MPC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.0030</td>
<td>0.0029</td>
<td>0.0063</td>
</tr>
<tr>
<td>De-PnPMPC</td>
<td>-26.66%</td>
<td>-31.25%</td>
</tr>
<tr>
<td>Di-PnPMPC</td>
<td>+0.00%</td>
<td>-8.62%</td>
</tr>
</tbody>
</table>

Table 5.9: Value of the performance parameter $\Phi$ for centralized MPC (first line) and percentage change using decentralized and distributed MPC schemes for the AGC layer. Best values for PnP controllers are in bold.
(a) Frequency deviation in each area controlled by the DePnP MPC (bold line) and centralized MPC (dashed line).

(b) Load reference set-point in each area controlled by the DePnP MPC (bold line) and centralized MPC (dashed line).

Figure 5.4: Simulation Scenario 1: 5.4(a) Frequency deviation and 5.4(b) Load reference in each area.
Figure 5.5: Simulation Scenario 1: tie-line power between each area controlled by the DePnP-MPC (bold line) and centralized MPC (dashed line).
Chapter 6

Invariant sets

In this chapter, we present methods to compute invariant sets. For more details we refer to the help of each function. First of all, we give some definitions.

**Definition 8 (Robust Positively Invariant (RPI) set).** The set $X \subseteq \mathbb{R}^n$ is RPI for $x(t + 1) = f(x(t), w(t))$, $w(t) \in W \subseteq \mathbb{R}^m$ if $x(t) \in X \Rightarrow f(x(t), w(t)) \in X$, $\forall w(t) \in W$.

**Definition 9 (Minimal Robust Positively Invariant (mRPI) set).** The RPI set $X$ is minimal if every other RPI $X$ verifies $X \subseteq X$. The RPI set $X(\delta)$ is a $\delta$-outer approximation of the minimal RPI $X$ if

$x \in X(\delta) \Rightarrow \exists \bar{x} \in X$ and $\bar{x} \in B_\delta(0) : x = \bar{x} + \bar{x}$

where, for $\delta > 0$, $B_\delta(v) = \{x \in \mathbb{R}^n \| x - v \| < \delta \}$.

**Definition 10 (\(\lambda\)-contractive control invariant set).** The set $X \subseteq \mathbb{R}^n$ is a $\lambda$-contractive set for $x(t + 1) = f(x(t), u(t))$, $u(t) \in U \subseteq \mathbb{R}^m$, if $\forall x(t) \in X$ there exist $u(t) \in U$ such that $x(t + 1) \in \lambda X$.

**Definition 11 (Robust Control Invariant (RCI) set).** The set $X \subseteq \mathbb{R}^n$ is an RCI set for $x(t + 1) = f(x(t), u(t), w(t))$, $u(t) \in U \subseteq \mathbb{R}^m$ and $w(t) \in W \subseteq \mathbb{R}^p$ if $\forall x(t) \in X$ there exist $u(t) \in U$ such that $x(t + 1) \in X$, $\forall w(t) \in W$.

### 6.1 \(\epsilon\)-mRPI

In this section, we explain how to use the `epsilon_mRPI` class in order to compute an $\epsilon$ outer approximation of the mRPI set for a linear constrained systems.

Our implementation is based on the algorithm proposed in [16]. We propose two examples.

#### 6.1.1 example1epsilon_mRPI.m

The code below shows how to compute an $\epsilon$-mRPI set for the LTI system $x^+ = Ax + Bu + w$ where $x \in \mathbb{R}^2$, $u = K x$ and $w$ lies in the polytope $W$. Non constraint on the state is assumed and hence $\mathbb{X} = \mathbb{R}^2$.

```matlab
% matrices of the LTI system
A = [ 1 1 ; 0 1 ];
```
\[ B = [ 1 ; 1 ]; \]
\[ K = -[1.17 1.03]; \]

% bounded disturbances as polytope object
\[ W = \text{polytope}([ \text{eye}(2) ; -\text{eye}(2) ],[ 1 1 1 1 ]'); \]

% approximation
epsilon = 5*10^{-5};

% create an object F that is a \( \epsilon \)-mRPI for the closed loop LTI system
% with dynamic \( A BK \) and disturbances bounded in \( W \) (\( W \) is a polytope object)
\[ F = \text{epsilon}_m\text{RPI}(A+B*K,W,\text{epsilon}) \]

% Otherwise one can create an object F that is a \( \epsilon \)-mRPI for the
% closed loop LTI system with dynamic \( A BK \) and disturbances bounded in \( W \)
% (\( W \) is a zonotope object)
% \[ W = \text{zonotope}([-0.5 0.1]',\text{eye}(2)); \]
% \[ F = \text{epsilon}_m\text{RPI}(A+B*K,W,\text{epsilon}); \]

% plot the approximation of the mRPI
\[ F.\text{plot} \]

Other useful methods for the \text{epsilon}_m\text{RPI} class are

\text{isinside}  \hspace{1em} \text{Test if a point is inside the \( \epsilon \)-mRPI set} \( F \).

\[ x = [ 1;1 ]; \]
\[ \text{test} = F.\text{isinside}(x) \]

\text{doubleHK}  \hspace{1em} \text{If} \( W \) \text{is a polytope object, the function returns the H-representation of the mRPI.}
\hspace{1em} \text{Moreover a polytope object of the mRPI is saved in} \( F.\text{epsilon} \).

\[ [ H K F ] = F.\text{doubleHK} \]
\[ \text{FP} = F.\text{F_epsilon} \]

6.1.2 \text{example2epsilon}_m\text{RPI.m}

This example shows how to compute an \( \epsilon \)-mRPI set in the case of constraints on the state variables,
\hspace{1em} i.e. \( x \in X \).

\[ X = \text{polytope}([ \text{eye}(2) ; -\text{eye}(2) ],[ 2 2 2 2 ]'); \]
\[ F = \text{epsilon}_m\text{RPI}(A+B*K,W,\text{epsilon},X) \]
6.2 \(\lambda\)-contractive control invariant sets

In this section, we explain how to use the `localControlLyapunov` class in order to compute a \(\lambda\)-contractive control invariant set for a discrete-time LTI system. Our implementation is based on the algorithm proposed in [10].

We illustrate the use of the class `localControlLyapunov` in the following example.

6.2.1 example1localControlLyapunov.m

```matlab
% matrices of the LTI system
A = [ 1 1 ; 0 1 ];
B = [ 0 ; 1 ];

% matrices for constraints on state and input variables
cx = [ eye(2) ; -eye(2) ];
dx = [ 2 2 1.5 2 ]';
cu = [ 1 ; -1 ];
du = [ 3 ; 3 ];

% parameter of the algorithm, \(\geq\) controllability index
k = 3;

% x0 is a parameter of the algorithm
% As x0 we consider the vertices of polytope = \(\{ x | cx \cdot x \leq dx \}\)
x0 = [2 2 ;-1.5 2 ;-1.5 -2;2 -2];

% create the lambda contractive control invariant set
L = localControlLyapunov(A,B,k,{cx,dx},{cu,du},x0)

% plot the lambda contractive control invariant set
L.plot

% Contractiveness
lambda = L.lambda
```

Other useful methods for the `localControlLyapunov` class are

- `isinside` Test if a point is inside the \(\lambda\)-contractive control invariant set \(L\).

```matlab
x = [1;1]
test = L.isinside(x)
```

- `uInv` Given \(x \in \mathcal{X}\), compute the control action \(u(x)\) such that \(x(t+1) \in \lambda\mathcal{X}\).

```matlab
x = zeros(2,1)
```
The function returns the H-representation of a \( \lambda \)-contractive control invariant set. Moreover a polytope object of the set can be obtained through the attribute LP.

\[
[ H K L ] = L \text{.double} \\
LP = L \text{.LP}
\]

6.3 Parameterized robust control invariant sets

In this section, we explain how to use the `parameterizedRCI` class in order to compute a parameterized robust control invariant set for a linear constrained systems. Our implementation is based on the algorithm proposed in [10].

We illustrate the use of the class `parameterizedRCI` in the following example.

6.3.1 `example1parameterizedRCI.m`

```matlab
% matrices of the LTI system
A = [ 1 1 ; 0 1 ];
B = [ 0 ; 1 ];

% matrices for constraints on state and input variables
cx = [ eye(2) ; -eye(2) ];
dx = [ 2 2 1.5 2 ];
cu = [ 1 ; -1 ];
du = [ 3 ; 3 ];

% parameter of the algorithm, greater than the controllability index of (A,B)
k = 5;

% x0 is a parameter of the algorithm such that W \subseteq hull(x0)
x0 = [ 0.99 0.99 ; 0.99 -0.01 ; -0.01 -0.01 ; -0.01 0.99 ];

% create the parameterized robust control invariant set
R = parameterizedRCI(A,B,k,[cx,dx],[cu,du],x0)

% plot the parameterized robust control invariant set
R.plot
```

Other useful methods for the `parameterizedRCI` class are

`isinide`  Test if a point is inside the parameterized robust control invariant set \( R \).
Given $x \in X$, compute the control action $u(x)$ such that $x(t + 1) \in X, \forall w \in W$.

```matlab
x = zeros(2,1)
u = R.uInv(x)
```

The function returns the H-representation of the parameterized robust control invariant set. Moreover a polytope object of the set can be obtained through the attribute `RP`.

```matlab
[H K R] = R.double
RP = R.RP
```

### 6.4 Zonotopic robust control invariant sets

In this section, we explain how to use the `zonotopeRCI` class in order to compute a zonotopic robust control invariant set for a linear constrained systems. Our implementation is based on the algorithm proposed in [17].

We illustrate the use of the class `zonotopeRCI` in the following example.

#### 6.4.1 example1zonotopeRCI.m

```matlab
% matrices of the LTI system
A = [ 1 1; 0 1 ];
B = [ 0; 1 ];

% matrices for constraints on state and input variables
% X = \{ x | cx \cdot x \leq dx \}
% U = \{ u | cu \cdot u \leq du \}
% 
% x = [ 1 0 ; -1 0 ; 0 -1 ; 0 1 ; -1 -1 ];
dx = [ 1.85 3 3 3 2.2 ]';
cx = [ 1 ; -1 ];
du = [ 3 ; 3 ];

% parameter of the algorithm, greater than the number of states
k = 5;

% matrices of the zonotope for the description of set W = \{ w=f+Ed , |d| \leq \inf\}
E = [ 1 0 ; 0 1 ];
f = [ 0 ; 0 ];

% create the zonotopic robust control invariant set
```
qa = 1 ; qb = 1 ; qp = 1 ;
Z = zonotopeRCI(A,B,{cx,dx},{cu,du},{f,E},k,[qa,qb,qp]);

% plot the zonotopic robust control invariant set
Z.plot

Other useful methods for the zonotopeRCI class are

isinside Test if a point is inside the zonotope robust control invariant set Z.

x = [1;1]
test = Z.isinside(x)

uInv Given $x \in X$, compute the control action $u(x)$ such that $x(t+1) \in X, \forall w \in W$.

x = zeros(2,1)
u = Z.uInv(x)
Chapter 7

PnPMPC and others toolboxes

7.1 Integration with the toolbox WIDE [18]

7.1.1 The lsmode2lss function

In WIDE a large-scale system is described by an LModel object. We use the function lsmode2lss to convert an LModel object to an lss object. The function declaration is

```
objlss = lsmode2lss( objlsmodel , m , p )
```

The arrays m and p are used to select the external inputs and external outputs in the LModel object. The k-th element of the array m is an integer number to classify the k-th external input of the LModel object: if m(k)=0 the k-th external input is a centralized input for the lss object, if m(k)=-1 the k-th external input is an exogenous input for the lss object and if m(k)=i the k-th external input is a local input for the i-th subsystem of the lss object. Similarly, the k-th element of the array p is an integer number to classify the k-th external output of the LModel object: if p(k)=i the k-th external output is a local input for the i-th subsystem of the lss object.

7.1.2 The createCtrlPnPMP4LModel function

Given an LModel object, we can design PnPMPC controllers using the createCtrlPnPMP4LModel function. This function calls the lsmode2lss function and the createCtrlPnPMPC method of the lss object of the function and returns an array of pnpmpc objects. The function declaration is

```
[ ctrl objlss ] = createCtrlPnPMP4LModel(objlsmodel,N,k,m,p,options)
```

For the meaning of the input arguments we defer the reader to Section 7.1.1 and Section 4.2.1.

7.1.3 Example

We create an LModel object of the power network system proposed in Scenario 1 in Chapter 5. We can create the LModel object executing the file makePNSLModel.m. Executing the
We can design PnP MPC controllers for the power network described by an LS model. We show below the essential part of the code.

```matlab
Q = diag([500 0.01 0.01 10]);
R = 10;
N = 15;
kmin = [8 8 8 6];
kmax = [20 20 20 20];

% m and p array for lsmodel2lss
% the odd elements of m are the power references of each area, therefore
% they are the local inputs of each subsystem; the even elements of m are
% the power loads of each area, therefore they are the exogenous signals
% of each subsystem
m = [1 -1 2 -1 3 -1 4 -1];
% the elements of p are the outputs of each subsystem, therefore they are
% the states of each area
p = [1 1 1 1 2 2 2 2 3 3 3 3 4 4 4 4];

% convert LSmodel to lss
objlss = lsmodel2lss(pnsD1ssLSmodel,m,p);
% design pnp controllers
ctrlPnpMpcFromLSmodel = objlss.createCtrlPnPMPC(N,{kmin,kmax},sdpsettings('verbose',0));
% one can also use the following instruction
% [ctrlPnpMpcFromLSmodel objlss ]=createCtrlPnPMPC4lsmode(pnsD1ssLSmodel,
% N , {kmin,kmax} , m , p , sdpsettings('verbose',0) )

% zero terminal constraint for each pnpmpc controller
ctrlPnpMpcZeroFromLSmodel = ctrlPnpMpcFromLSmodel;
for i=1:size(ctrlPnpMpcFromLSmodel,2)
    ctrlPnpMpcZeroFromLSmodel(i)=ctrlPnpMpcFromLSmodel(i).zeroTerminal(Q,R)
end
```
Chapter 8

Zonotope and Polytope classes

In PnPMPC-toolbox the zonotope class has been developed as an integration of MPT2 [3]. New methods have been proposed also for the polytope class. In the following we describe how to generate zonotope sets. Since several functions have the same meaning of functions of polytope class, we defer the reader to the MPT2 manual. These functions implement the standard operations between zonotope and polytope sets: Minkowski sum ($\oplus$), Pontryagin difference ($\ominus$), intersection ($\cap$), union ($\cup$) and relational operators ($\subset$, $\subseteq$, $\supset$, $\supseteq$, $=$ and $\neq$). Zonotope arrays are managed as polytope arrays in MPT2.

In the following sections, we introduce some useful operators for zonotope sets. Most of the definition are from [19].

8.1 Creating a zonotope

A zonotope in PnPMPC-toolbox is created by a call to the zonotope constructor

\[
Z = \text{zonotope}(p, G)
\]

This instruction creates a zonotope $Z$ centered in $p$ and described by generators $G$. Example

```
p = [ 1 ; 1 ];
G = [ 1 2 4 ; 3 1 4 ];
Z = \text{zonotope}(p, G)
\text{plot}(Z)
```

The zonotope is shown in Figure 8.1.

In order to access the G-representation of zonotope $Z$, one can use the command `doublePG`

```
[ p G ] = \text{doublePG}(Z)
```

In order to access the H-representation of zonotope $Z$, one can use the command `doubleHK`

```
[ H K Zo ] = \text{doubleHK}(Z)
```
Since the computation of the H-representation of zonotope $Z$ requires the computation of the V-representation both representation are stored in the output variable $zo$ for future uses.

### 8.2 Functions for zonotope objects

#### 8.2.1 Bisection and Split operators

The operator $\text{Bisect}_k(\cdot)$ generates two sub-zonotopes from one zonotope. In particular, given a zonotope $Z$ the operator $\text{Bisect}_k(Z)$ generates the two sub-zonotopes

$$Z^L = (p - \frac{g_k}{2}) \oplus \left[ g_1 \ldots \frac{g_k}{2} \ldots g_m \right] d, \ ||d||_{\text{inf}} \leq 1$$

$$Z^R = (p + \frac{g_k}{2}) \oplus \left[ g_1 \ldots \frac{g_k}{2} \ldots g_m \right] d, \ ||d||_{\text{inf}} \leq 1$$

where $g_k$ is the $k$-th column of $G$ and is the biggest generator, i.e. we split the $k$-th column in the middle. With the operator $\text{Split}_k(Z, \alpha)$ it is possible to split the $k$-th column of $G$ in a desired position $\alpha$, i.e., $\text{Split}_k(Z, \alpha)$ generates two sub-zonotopes

$$Z^L = (p - g_k(1 - \alpha)) \oplus \left[ g_1 \ldots g_k \alpha \ldots g_m \right] d, \ ||d||_{\text{inf}} \leq 1$$

$$Z^R = (p + g_k \alpha) \oplus \left[ g_1 \ldots g_k(1 - \alpha) \ldots g_m \right] d, \ ||d||_{\text{inf}} \leq 1$$

where $g_k$ is a column of $G$ matrix and the parameter $\alpha \in [0, 1]$. Figure 8.2 shows an example of the operator $\text{Bisect}_k(Z)$ applied to the zonotope $Z$ in Figure 8.1. This example shows that the operator $\text{Bisect}_k(Z)$ can generate two sub-zonotopes which intersect in their interior. The reason of the overlapping of $Z^L$ and $Z^R$ is that the line segment generators $g_1, \ldots, g_m$ are not linearly independent. Given $G \in \mathbb{R}^{n \times m}$, with $\text{Rank}(G) = n$, then, the bisection is complete, i.e., $\text{Bisect}_k(Z)$ provides two sub-zonotopes that do not overlap. The operators $\text{Bisect}_k(\cdot)$ and $\text{Split}_k(\cdot)$ have been implemented in PnPMPC-toolbox in the following two instructions
The following code shows an example where zonotope $Z$ is splitted.

```matlab
p = [1; 1];
G = [1 2 4; 3 1 4];
alpha = 0.2;
k = 2;
Z = zonotope(p,G)
Znb = bisect(Z)
Zns = split(Z,k, alpha)

figure(1)
plot(Z)
figure(2)
plot(Znb, struct('shade',0.6))
figure(3)
plot(Zns, struct('shade',0.6))
```

Figure 8.2: An example of split. $Bisect(Z)$.

### 8.2.2 Bounding box operator

The smallest interval vector containing $Z$ and having its same center is given by the operator $rs(\cdot)$ (row sum). Given a matrix $G \in \mathbb{R}^{n \times m}$, $rs(G)$ is a $n \times n$ diagonal matrix

$$rs(G)_{ii} = \sum_{j=1}^{m} |G_{ij}|, \; i = 1, \ldots, n.$$
The operator $rs(\cdot)$ has been implemented in PnP-MPC-toolbox in the following instruction

\[ BB = \text{bounding\_box}(Z) \]

where $BB$ is a zonotope object computed using operator $rs(\cdot)$. The following code shows an example where zonotope $Z$ is bounded by the smallest interval vector.

```matlab
p = [ 1 ; 1 ; -1 ];
G = [ 1 2 4 2.4 5 ; 3 1 4 5 6 ; 2 4 5 1 5 ];
Z = zonotope(p,G)
BB = bounding_box(Z)
figure
hold on
plot(BB)
plot(Z,'b')
```

Figure 8.3: An example of bounding box of a set $Z$.

### 8.2.3 Reduction operator

Another important operator is the reduction operator, whose purpose is to outer bound a given zonotope with a zonotope of reduced complexity, i.e., a reduced number of line segment generators. Given the zonotope $Z$, the reduction operator $\text{red}_n gen(Z)$ produces a lower complexity zonotope generated by a maximum of $ngen$ line segment generators. The procedure consists of first sorting the columns of $G$ with respect to decreasing Euclidean norm

\[ G = [g_1, g_2, \ldots, g_m], \quad \|g_i\| \geq \|g_{i+1}\|, \quad i = 1, \ldots, m - 1. \]
Then, denoting by $G_{ngen}$ the matrix describing $\text{red}_{ngen}(Z)$, we define

$$G_{gen} = \begin{cases} G, & \text{if } m \leq ngen \\ G_{ngen} = [g_1, \ldots, g_{ngen-n}, g_r], & \text{if } m > ngen. \end{cases}$$

It is important to mention that $Z \subseteq \text{red}_{ngen}(Z)$. Figure 8.4 shows the application of the reduction operator. As one can see, a reduction in the number of generators yields a more conservative zonotope. The operator $\text{red}_{ngen}()$ has been implemented in PnPMPC-toolbox in the following instruction

```
Zo = reduce(Z,ngen)
```

The following code shows an example where the number of generators of zonotope $Z$ is reduced.

```
p = [ 1 ; 1 ];
G = [ 1 2 4 2.4 5 ; 3 1 4 5 6 ];
Z = zonotope(p,G)
negen = 3;
Zo = reduce(Z,negen)
figure(1)
hold on
plot(Zo,’r’)
plot(Z,’b’)
```

Figure 8.4: The reduction operator $\text{red}_{ngen}(Z)$ is applied to the green zonotope yielding a more conservative approximation (red zonotope).
8.2.4 The support function of polytope/zonotope sets

The support function of a polyhedral set is defined as

\[ sup = \max_{x \in X} c'x \]  
(8.1)

For polytope and zonotope objects we developed the function \texttt{supportFunction}. One can compute \( sup \) using the following instruction

\[
\texttt{sup = supportFunction(X, c)}
\]

where \( X \) is a polytope or zonotope object.

8.2.5 Zonotope ↔ Polytope

- Polytope \( P \) is a zonotope?

\[
\texttt{bool = iszonotope(P)}
\]

- Polytope \( P \) to Zonotope \( Z \)

\[
\texttt{Z = polytope2zonotope(P)}
\]

- Zonotope \( Z \) to Polytope \( P \)

\[
\texttt{P = zonotope2polytope(Z)}
\]

8.3 Outer approximation of a nonlinear function

The function \texttt{outerApproximation}\(^1\) allows one to compute an outer approximation of a nonlinear function evaluated on a zonotope \( Z \). We compute the outer approximation of the nonlinear function using the methods proposed in [20, 19]. In [20], an algorithm to compute zonotope outer approximations for nonlinear systems was proposed. The authors suggest to create an image of a zonotope through a nonlinear function using DC programming, which is based on DC functions. A DC function \( f(x) : \mathbb{R}^n \rightarrow \mathbb{R} \) is a function that can be expressed as the difference of two convex functions, i.e., \( f(x) = g(x) - h(x) \) where \( g(x) \) and \( h(x) \) are convex functions. In order to compute a tighter outer approximation, in [19] the authors proposed an algorithm in order to split the zonotope set \( Z \) where the function \( f(x) \) is “more nonlinear” and then the outer approximation is obtained as the union of the outer approximations obtained using the DC programming on each zonotope of the splitting.

In the following, we propose three examples that can be found in the PnPMPC-toolbox.

\(^1\)The function \texttt{outerApproximation} needs Symbolic Math Toolbox and Partial Differential Equation Toolbox.
8.3.1 example1outerApproximation.m

```matlab
% definition of function f, set X and sampling of f(X)
npoints = 500;
X = zonotope(zeros(2,1),[2 0;0 3]);
xf = zeros(dimension(X),npoints);
f = @(x,w)([ 1+0.1*x(1)+0.5*x(2)-exp(0.1*x(1)^2) ;
0.1+0.9*x(1)-0.1*x(2)-0.1*cos(x(2))+0.05*x(2)^2 ]); for i=1:size(xf,2)
    xx = randpoint(X);
    xf(:,i) = f(xx);
end

% options for outerApproximation function
options.split.ngenerators = 2;
options.split.alpha = 0.5;
options.split.max_zono = 5;

% compute zonotope approximation of f(X)
% Zu approximation using 1 zonotope
% ZuVec approximation using options.split.max_zono zonotopes
% Xs zonotope X splitted
% J jacobian and H hessian are outputs variables for future computations
% The third and fourth input argument are empty, that means the user does
% not know convex functions gf and hf, such that f = gf - hf. Therefore the
% function outerApproximation computes gf and hf. We highlight that the
% IntLab toolbox http://www.ti3.tuhh.de/rump/intlab/ is needed.
[ Zu ZuVec Xs J H ] = outerApproximation(X,f,[],[],[],options);

% plots
figh = figure(1);
subplot(1,2,2)
hold on
plot(Zu,'y')
plot(ZuVec,'r')
plot(xf(1,:),xf(2,:),b.)
box on
subplot(1,2,1)
plot(Xs,'g')
box on
```

8.3.2 example2outerApproximation.m

```matlab
% definition of function f, set X and sampling of f(X)
npoints = 500;
```
% Figure 8.5: Results of example1outerApproximation.m.

\[
X = \text{zonotope(zeros(2,1),[2 0; 0 3])};
\]

\[
xf = \text{zeros(dimention(X),npoints)};
\]

\[
f = \emptyset(x,w) (\begin{bmatrix}
1+0.1\times x(1)+0.5\times x(2)-\exp(0.1\times x(1)^2) \\
0.1+0.9\times x(1)-0.1\times x(2)-0.1\times \cos(x(2))+0.05\times x(2)^2
\end{bmatrix})
\]

\[
\text{for } i = 1:\text{size}(xf,2)
xx = \text{randpoint}(X);
xf(:,i) = f(xx);
\]

\[
g = \emptyset(x,w) (\begin{bmatrix}
0.5\times x(2) \\
0.1+0.9\times x(1)-0.1\times \cos(x(2))+0.05\times x(2)^2
\end{bmatrix})
\]

\[
h = \emptyset(x,w) (\begin{bmatrix}
-1-0.1\times x(1)+\exp(0.1\times x(1)^2) \\
0.1\times x(2)
\end{bmatrix})
\]

\[
\% definition of g and h such that f=g-h where g and h are convex functions
\]

\[
g = \emptyset(x,w) (\begin{bmatrix}
0.5\times x(2) \\
0.1+0.9\times x(1)-0.1\times \cos(x(2))+0.05\times x(2)^2
\end{bmatrix})
\]

\[
h = \emptyset(x,w) (\begin{bmatrix}
-1-0.1\times x(1)+\exp(0.1\times x(1)^2) \\
0.1\times x(2)
\end{bmatrix})
\]

\[
\% options for outerApproximation function
\]

\[
\text{options.split.ngenerators = 2;}
\]

\[
\text{options.split.alpha = 0.5;}
\]

\[
\text{options.split.max_zono = 5;}
\]

\[
\% compute zonotope approximation of f(X)
\]

\[
\% Zu approximation using 1 zonotope
\]

\[
\% ZuVec approximation using options.split.max_zono zonotopes
\]

\[
\% Xs zonotope X splited
\]

\[
\% J jacobian and H hessian are outputs variables for future computations
\]

\[
\text{[ Zu ZuVec Xs J H ] = outerApproximation(X,f,[],g,h,options);}
\]

\[
\% plots
\]

\[
\text{figh = figure(1);}
\]

\[
\text{subplot(1,2,2)}
\]
8.3.3 example3outerApproximation.m

```matlab
% definition of function f, set X and sampling of f(X)
% differently from example2outerApproximation we want to split only along % x(1), therefore x(2) is substituted by w(1)

npoints = 500;
X1 = zonotope(0,2);
W = zonotope(0,3);
X = X1*W;
xf = zeros(dimension(X),npoints);

f = @(x,w)([ 1+0.1*x(1)+0.5*w(1)-exp(0.1*x(1).^2) ;
            0.1+0.9*x(1)-0.1*w(1)-0.1*cos(w(1))+0.05*w(1).^2 ]);

for i=1:size(xf,2)
    xx = randpoint(X);
    xf(:,i) = f(xx(1),xx(2));
end
```
% definition of g and h such that f=g-h where g and h are convex functions

\[
\begin{align*}
    g &= @(x,w) ([0.5\cdot w(1); 0.1+0.9\cdot x(1)-0.1\cdot \cos(w(1))+0.05\cdot w(1)^2]); \\
    h &= @(x,w) ([ -1-0.1\cdot x(1)+\exp(0.1\cdot x(1)^2); 0.1\cdot w(1)]);
\end{align*}
\]

% options for outerApproximation function

options.split.ngenerators = 2;
options.split.alpha = 0.5;
options.split.max_zono = 5;

% compute zonotope approximation of f(X)
% Zu approximation using 1 zonotope
% ZuVec approximation using options.split.max_zono zonotopes
% Xs zonotope X splitted
% J jacobian and H hessian are outputs variables for future computations
[ Zu ZuVec Xs J H ] = outerApproximation(X1,f,W,g,h,options);

% plots
h=figure(1);
subplot(1,2,2)
hold on
plot(Zu,'y')
plot(ZuVec,'r')
plot(xf(:,1),xf(:,2),'b')
box on
subplot(1,2,1)
plot(Xs,'g')
box on
Figure 8.7: Results of example example3outerApproximation.m.
Bibliography


