Learning Based MPC

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Documentation for LBmpcTP template class

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Introduction

The aim of this report is to give a brief introduction to the LBmpcTP template class, a particular implemention of the interior point method solver taylored to the learning based MPC algorithm described in [1] with quadratic cost and affine oracle dynamics.

The report is structured as follows: First, the Learning Based MPC model is introduced. Second, the interface to the LBmpcTP class is presented. In the third section, some tweaks and hidden parameters are described.

1 The Learning Based MPC model

The learning based MPC model is taken from [1]. In our particular framework, the cost is assumed to be quadratic and the oracle dynamics affine in the oracle state \tilde{x} and input \tilde{u} . Furthermore, the feasible sets are assumed to be convex polyhedron.

Hence, we consider the following optimization problem:

$$\min_{c[\cdot],\theta} \quad (\tilde{x}[m+N] - x^{\star}[m+N])^{T} \tilde{Q}_{f}(\tilde{x}[m+N] - x^{\star}[m+N]) + \\
\sum_{i=0}^{N-1} \{ (\tilde{x}_{[}m+i] - x^{\star}[m+i])^{T} \tilde{Q}(\tilde{x}_{[}m+i] - x^{\star}[m+i]) + (\tilde{u}[m+i] - u^{\star}[m+i])^{T} R(\tilde{u}[m+i] - u^{\star}[m+i]) \}$$

$$\begin{split} \text{s.t.} \quad & \tilde{x}[m] = \hat{x}[m], \quad \bar{x}[m] = \hat{x}[m] \\ & \tilde{x}[m+i] = A\tilde{x}[m+i-1] + B\check{u}[m+i-1] + s + \mathcal{O}_m(\tilde{x}[m+i-1],\check{u}[m+i-1]), \quad \forall i \\ & \mathcal{O}_m(\tilde{x}[m+i-1],\check{u}[m+i-1]) = L_m\tilde{x}[m+i-1] + M_m\check{u}[m+i-1] + t_m, \quad \forall i \\ & \bar{x}[m+i] = A\bar{x}[m+i-1] + B\check{u}[m+i-1], \quad \forall i \\ & \check{u}[m+i-1] = K\bar{x}[m+i-1] + c[m+i-1], \quad \forall i \\ & F_{\bar{x}[m+i]}\bar{x}[m+i] \leq f_{\bar{x}[m+i]}, \quad F_{\check{u}[m+i]}\check{u}[m+i] \leq f_{\check{u}[m+i]}, \quad \forall i \\ & F_{x\theta}\bar{x}[m+i] + F_{\theta}\theta \leq f_{x\theta}, \quad i \in \{1,\dots,N\} \end{split}$$

The conditions on the matrices above are given in Tab. 1.

To solve (1), the optimization problem (1) is casted into the following quadratic program form:

$ ilde{Q}$	positive definite
$ ilde{Q}_f$	positive definite
R	positive semidefinite
$F_{\bar{x}[m+i]}$	full rank
$F_{\check{\mathbf{u}}[m+i]}$	full rank
F_{θ}	full rank

Tabelle 1: Assumptions on matrices.

$$\begin{split} \min_{z} & z^{T}Hz + g^{T}z \\ \text{s.t.} & Cz = b \\ & Pz \leq b, \end{split} \tag{2}$$

where z is the stacked vector:

$$z = \begin{pmatrix} c[m]^T & \bar{x}[m+1]^T & \tilde{x}[m+1]^T & \cdots & c[m+N-1]^T & \bar{x}[m+N]^T & \tilde{x}[m+N]^T & \theta^T \end{pmatrix}^T$$

To solve the QP, the LBmpcTP template class applies a primal barrier interior-point method. To this end, we add a log-barrier term to the cost function in (2), and get:

$$\min_{z} z^{T}Hz + g^{T}z - \kappa \sum_{i} \log(h - Pz)_{i}$$
s.t. $Cz = b$. (3)

A sequence of such problems is solved where at each step the cost penalty factor κ is reduced by a factor $\mu \in (0,1)$, i.e. $\{\kappa_0, \kappa_1, \kappa_2, \ldots\}$, $\kappa_i \triangleq \mu^i \kappa_0$. For each κ , (3) is solved using an infeasible Newton start method.

2 Using the LBmpcTP template class

The LBmpcTP template class is typically called in two seperate steps:

- 1. Definition of matrices such as $A,\ B,\ s,\ \tilde{Q},\ \tilde{Q}_f,\ R,\ K,\ \{F_{\bar{x}[m+i]}\}_i,\ \{f_{\bar{x}[m+i]}\}_i,\ \{F_{\tilde{u}[m+i]}\}_i,\ \{f_{\tilde{u}[m+i]}\}_i$ and scalars such as $\epsilon_{barrier},\ \epsilon_{nt},\ \epsilon_{rp},\ \alpha,\ \beta,\ \kappa_{0,I},\ \kappa_{0,II},\$ in MATLAB file Init.m. These values are written to a binary file, which is called ConstrParam.bin by default. The complete list of variables to be specified can be found in Tab. 2.
- 2. A C++-file (e.g. mainLBmpcTP.cpp) then reads the binary file ConstrParam.bin. mainLBmpcTP.cpp is the main function file and performs two tasks:
 - (a) It calls the constructor of the template class in LBmpcTP.h and instantiates an object of this template class, e.g. myObj.
 - (b) It performs the step-function myObj.step(.) which returns the optimal input u_opt. At each call of the step-function, the following parameters are needed: L_m , M_m , t_m , \hat{x} , $\{x^*[m+i]\}_i$.

The file can be compiled using the gcc-compiler and the following command: g++ -I /usr/local/include/eigen3/ -O3 mainLBmpcTP.cpp -o mainLBmpcTP.

In the following sections, both files and the variables are described in more detail.

2.1 MATLAB: Init.m

In this MATLAB-file, the parameters required for the instantiation of the LBmpcTP object are defined. More specifically, Init.m consists of two parts:

- User has to manually specify the parameters given in Tab. 2. See also (1).
- A binary file (default: ConstrParam.bin) containing the parameters in Tab. 2 is written by calling the m-file writeParam.m.

In appendix B.1, a typical implementation of the Init.m file is shown.

Remarks:

- The number of state constraints is assumed to be constant, i.e. the number of rows in $Fx\{i\}$ is constant for all i. Subsequently, the number of state constraints is denoted as $_nSt$.
- The number of input constraints is assumed to be constant, i.e. the number of rows in Fu{i} is constant for all i. Similarly, the number of input constraints is denoted as _nInp.
- The number of constraints involving θ in (1) is assumed to be _nF_xTheta.

2.2 C++: mainLBmpcTP.cpp

This file contains the main control routine which interacts with the LBmpcTP template class. An example file is provided in appendix B.2. The tasks of mainLBmpcTP.cpp include:

- Read values from ConstrParam.bin.
- Instantiate an object from the template class LBmpcTP, e.g. myObj.
- Update the oracle dynamics and retrieve L_m , M_m , t_m , $\hat{x}[m]$, $\{x^*[m+i]\}_i$ from an external source (not provided in this framework).
- Solve the optimization problem (3) by calling the step-function with the updated variables above, i.e. myObj.step(.), and obtain the optimal input u_opt.

Since LBmpcTP is a template class and makes use of the linear algebra template class Eigen, some template parameters must be adjusted manually. More specifically, the following steps must be completed (a description of the individual parameters can be found in Tab. 3):

- 1. SPECIFY parameters: _N, _m, _n, _nSt, _nInp, _nF_xTheta and _pos_omega.
- 2. mainLBmpcTP.cpp reads the parameters (with _arg appended) listed in Tab. 2 from the binary source file ConstrParam.bin.
- 3. The oracle matrices Lm_arg, Mm_arg, tm_arg, x_hat_arg, x_star_arg[_N] are updated. Note: Computation of these values is not provided by the LBmpcTP template class.
- 4. The optimization problem (1) is then solved with the updated oracle matrices by calling the member function myObj.step(.).

It should be noticed that the parameters in the MATLAB file Init.m and the C++ file mainLBmpcTP.cpp should be consistent with each other.

Tabelle 2: Key parameters in Init.m

MATLAB variable	description	typical range/value
N	length of MPC horizon	
m	number of inputs	
n	number of states	
A	linear dynamics matrix: $\bar{x}^+ = A\bar{x} + B\check{u} + s$	
В	input-state dynamics matrix: $\bar{x}^+ = A\bar{x} + B\check{u} + s$	
S	affine offset in state dynamics: $\bar{x}^+ = A\bar{x} + B\check{u} + s$	
K	feedback gain matrix, $\check{u} = K\bar{x} + c$, $A + BK$ is stable	
Q_tilde	p.d. weight matrix for state	
Q_tilde_f	p.d. weight matrix for final state	
R	p.s.d. weight matrix on input	
$Fx\{i\}$	$F_{\bar{x}[m+i]}\bar{x}[m+i] \leq f_{\bar{x}[m+i]}, \text{ full-rank}, i=1,\ldots,N$	
$fx{i}$	$F_{\bar{x}[m+i]}\bar{x}[m+i] \leq f_{\bar{x}[m+i]}$	
$\operatorname{\mathtt{Fu}}\{i\}$	$F_{\check{u}[m+i]}\check{u}[m+i] \leq f_{\check{u}[m+i]}, \text{ full-rank}, \ i=0,\ldots,N-1$	
$\mathtt{fu}\{i\}$	$F_{\check{\mathbf{u}}[m+i]}\check{\mathbf{u}}[m+i] \le f_{\check{\mathbf{u}}[m+i]}$	
F_xTheta	$F_{x\theta}\bar{x}[m+i] + F_{\theta}\theta \le f_{x\theta}$	
F_theta	$F_{x\theta}\bar{x}[m+i] + F_{\theta}\theta \le f_{x\theta}$, full-rank,	
f_xTheta	$F_{x\theta}\bar{x}[m+i] + F_{\theta}\theta \le f_{x\theta}$	
kappa_start_PhaseI	Phase I starting value of $\kappa = \kappa_0$ to find z_0 : $Pz_0 < h$	100
kappa_start_PhaseII	Phase II starting value of $\kappa = \kappa_0$ for which (3) is solved	[1e2, 1e9]
mu	rate at which κ is decreased in (3)	[1/50, 1/2]
eps_nt	threshold at which the Newton steps are stopped,	
	i.e. $\ ig(r_d^T r_p^Tig)\ \leq ext{ ext{eps_nt}}$	0.1
eps_normRp	threshold primal feasibility, i.e. $\ r_p\ \leq \text{eps_normRp}$	0.1
eps_barrier	bound on suboptimality of solution,	0.1
	i.e. $\kappa \cdot (\# \text{ constraints}) \leq \text{eps_barrier}$	(depends on J_{opt})
eps_ls	lower threshold on step size t in backtracking line search,	_
	e.g. $z^+ = z + t \cdot \Delta z$	1e-7
n_iter_PhaseI	max. number of Newton steps allowed to solve (3) for fixed κ ,	[# #0]
D. TT	(does not apply to the first step, i.e. $\kappa = \text{kappa_start_PhaseII}$, tbd)	[5, 50]
n_iter_PhaseII	similar to n_iter_Phase I, just for Phase I.	[10, 50]
alpha_ls	parameter in line search, i.e. $ r(t) \le (1 - t \cdot \texttt{alpha_ls}) r $, where $r = \begin{pmatrix} r_d^T & r_p^T \end{pmatrix}^T$	
beta_ls	decrease parameter of t in line search, i.e. $t^+ := \beta t$	[0.1, 0.8]
reg_PhaseI	regularization coefficient for PhaseI	0.1
reg_PhaseII	regularization coefficient for PhaseII	[0.001, 1]
weight_PhaseI	weight on the linear cost in PhaseI	1e3

variable	description	default
Туре	only "double" is supported	double
_N	length of MPC horizon	
_m	number of inputs	
_n	number of states	
_nSt	number of state constraints (constant over the horizon)	
_nInp	number of input constraints (constant over the horizon)	
_nF_xTheta	number of constraints involving θ in (1)	
_pos_Omega	index i in $F_{x\theta}\bar{x}[m+i] + F_{\theta}\theta \le f_{x\theta}$	
Lm_arg	oracle matrix, i.e. $\mathcal{O}_m(\tilde{x}[m+i], \check{u}[m+i]) = L_m \tilde{x}[m+i] + M_m \check{u}[m+i] + t_m$	
Mm_arg	oracle matrix, i.e. $\mathcal{O}_m(\tilde{x}[m+i], \check{u}[m+i]) = L_m \tilde{x}[m+i] + M_m \check{u}[m+i] + t_m$	
tm_arg	oracle matrix, i.e. $\mathcal{O}_m(\tilde{x}[m+i], \check{u}[m+i]) = L_m \tilde{x}[m+i] + M_m \check{u}[m+i] + t_m$	
x_hat_arg	current state estimate, i.e. $\tilde{x}[m] = \hat{x}[m], \bar{x}[m] = \hat{x}[m]$	
$x_star_arg[_N]$	array of desired states in cost function (1)	
u_opt	stores the optimal input from myObj.step(.)	

Tabelle 3: Key parameters in mainLBmpcTP.cpp

2.3 C++ template class: LBmpcTP.h

This section gives a rough overview of what happens inside the LBmpcTP class. Access to the class is possible through two methods, the constructor and the step(.) method. The main idea of the algorithm is based on [2]. The constructor initializes some of the private variables as discussed in the previous sections. The step(.)-method performs the following tasks:

• We recursively compute the sequence $\{u^*[m+i]\}_i$ from the given desired state sequence $\{x^*[m+i]\}_i$ by solving

$$x^*[m+i] = (A+L_m)x^*[m+i-1] + (B+M_m)u^*[m+i-1] + (s+t_m)$$

and taking the least-squared solution (SVD).

- Cast (1) into (2).
- If the warm start variable z_warm does not satisfy $P \cdot z_warm < h$, then the method PhaseI() computes a new z_warm which satisfies the above (strict) inequality, see App. A.
- Solve (2) by solving a sequence of (3). It can be shown that as κ approaches 0, the solution of (3) will converge to the solution of (2).
- Finally, it returns the control input based on the eps_barrier-suboptimal solution or if the number of Newton iterations exeeds n_iter.

3 Hidden Parameters, Tweaks

Some of the parameters given in Tab. 2 can be used to tweak the LBmpcTP template class if the algorithm does not work as desired:

- The problem cannot be solved with the default parameters. It either does not converge or the code returns nan.
- Convergence is too slow for the desired purpose, i.e. the optimization step needs too many Newton or backtracking line search steps.

• The exact solution is not desired and an approximate solution suffices to speed up algorithm.

The goal of this section is to share some experience of how to react to certain situations and give some general advice on how to choose the parameters.

Tab. 4 lists the tuning parameters from Tab. 2 and describes their role and influence in greater detail.

3.1 Compiling

Some compilers provide the option to generate optimized executable codes. For example, the gcc compiler allows the user to add the -03 option which reduces the size of the executable file and increases the performance of the generated at the expense of longer compiling time (more than 2 min) and more memory (more than 2 GB) usage: g++ -I /usr/local/include/eigen3/ -03 mainLBmpcTP.cpp -o mainLBmpcTP

3.2 Troubleshooting

In this section, some common errors are described. Possible sources for these errors are given and solutions are proposed.

- 1. Phasel: does not converge or requires too many Newton steps.
 - Many Newton steps may indicate that the matrices are badly conditioned. Try to increase kappa_start_PhaseI,
 i.e. start with a larger κ₀.
 - Try to decrease the value of reg_PhaseI to give the quadratic cost less weight.
 - Increase weight_PhaseI to give the linear cost (original cost) more weight.
 - If still a lot of Newton steps are required increase n_iter_PhaseI.
- 2. PhaseII: many Newton steps ($\gtrsim 100$) are required to solve (3), i.e. κ fixed. Solutions:
 - IF FOR $\kappa=\kappa$ rappa_start_PhaseII: Increase kappa_start_PhaseII. For example, increasing the value of kappa_start_PhaseI from $\kappa=100$ to $\kappa=1e9$ reduced the number of Newton iterations from 170 to 5 for a particular problem. However, the larger kappa_start_PhaseI, the more problems of form (3) must be solved, until κ is sufficiently small.
 - IF FOR $\kappa \neq \kappa$ Kappa_Start_PhaseII: This can be typically observed for small κ ($\kappa \lessapprox 0.1$), where many Newton steps with small step sizes ($t \approx 10^{-17}$) are performed. Since this is waste of computational time, the number of Newton iterations can be upperbounded by choosing a smaller n_iter_PhaseII. Depending on the problem setup, numbers as few as 5 iterations might be enough to produce usable results.
- Phasel/Phasell: obtained result is a nan-vector (not a number).
 Solutions:
 - This problem typically shows up during Phase II when z_{opt} lies on some face of the feasible set. Even though positive definiteness (and hence the existence of Cholesky decomposition) is theoretically guaranteed, this might not be true from a numerical point of view. Indeed, a nan often suggests that some of the eigenvalues numerically approach zero, ending up dividing by zero, leading to nan. To solve this, the cost function in (3) is regularized using the weight term reg_PhaseI or reg_PhaseII. Thus, choosing a largerreg_PhaseI/reg_PhaseII usually overcomes this problem, but may return inferior results.
- 4. **Choosing** eps_barrier: Difficult, since choosing it small when the optimal value is large is only a waste of computational power. However, if the order of the optimal value is known approximately, then eps_barrier can be chosen accordingly.

Tabelle 4: Tuning parameters defined in Init.m

tuning variable	influence	typical range/value
kappa_start_PhaseI	If Phase I needs a lot of Newton iterations to find a suitable z_warm, increasing kappa_start_PhaseI might reduce this number.	100
kappa_start_PhaseII	Small κ can lead to poorly conditioned matrices. If many iterations ($\gtrsim 100$) are needed to solve (3) for $\kappa = \text{kappa_start_PhaseII}$, starting with a larger kappa_start_PhaseII might help. Downside: a larger sequence of problems of form (3) must be solved.	[1e2, 1e9]
mu	The smaller mu, the fewer problems of form (3) has to be solved. However, if mu is too small, i.e. $\mu\kappa\ll\kappa$, the problems might become difficult to solve, resulting in nan. This is especially the case if the minimizer lies on a face of the feasible set.	[1/50, 1/2]
eps_nt	The smaller this value, the more accurately are the intermediate problems of form (3) solved. However, there is usually no need to exactly solve the intermediate problems.	0.1
eps_normRp	Should not be set too large, because primal feasibility ought to be achieved accurately.	0.1
eps_barrier	If the optimal solution J^* was known, then one could simply allow a deviation of several percentages. However, J^* is usually not known apriori, making it difficult to choose.	0.1
eps_ls	The lower bound is motivated by the fact that for very small step sizes, $z^+ = z + t\Delta z \approx z$. Hence, it is better to stop the backtracking line search and perform the next Newton step.	1e-7
n_iter_PhaseII	This parameter bounds the max. number of Newton steps. It is often observed that, especially for small κ , only the first few Newton steps have significant step size t . The following Newton steps often have step sizes of order 10^{-17} , having no effect on the updates of z . However, if \mathbf{n} _iter is set to be too small, the problems might be solved inaccurately, leading to poor, even no, solutions.	[5, 50]
n_iter_PhaseI	Similar idea as for PhaseII, typically larger than in PhaseI	[20, 50]
alpha_ls	typically 0.01	
beta_ls	If set very small, t decreases rapidly, thus missing the optimal t by far. This might lead to the need to perfom more Newton steps, which are expensive. However, if beta_ls ≈ 1 , too many $t^+ := \beta t$ steps are needed, slowing down the algorithm.	[0.1, 0.8]
reg_PhaseI	This regularization term is needed to cast the Phase I formulation (which is a linear program) into a quadratic program, which we can solve using the same tools as in (3). If the Phase I algorithm does not run as anticipated, decreasing reg_PhaseI, which influences the quadratic term, can help. Also see App. A	0.1
reg_PhaseII	It can be often observed that if the minimzer z_{opt} lies on the face of the polyhedron, the algorithm runs into numerical problem when computing the Cholesky decomposition. To avoid this, the quadratic cost H in (3) is regularized using the coefficient reg_PhaseII. Note that if many Newton steps are performed, the algorithm more likely runs into numerical problems, requiring a larger reg_PhaseII.	[0.001, 1]
weight_PhaseI	Weight on the linear cost during the PhaseI. See App. ??	1000

3.3 Additional Remarks

- So far, the algorithm only works for a minimum prediction horizon of 3.
- There are more parameters in the LBmpcTP.h file which can be used to improve the performance of the solver, such as the difference variable in Phasel or how exactly to regularize the cost functions in Phasel and Phasell. However, it usually suffices to tune the parameters given in Tab. (4).
- If the prediction horizon $N \geq 50$, then some variable definitions in the class file have to be changed. More precisely, the size of the preallocated array of the LLT-class must be increased. This is done easiest by searching for the term LLT in the variable definition in LBmpcTP.h.
- When an LBmpcTP object is instantiated, a class variable called z_warm representing the "warm start" from
 one time step to another is created. By default, it is the 0-vector. However, this can be changed easily
 to a more appropriate value.

A PHASE I 9

A Phase I

This section roughly describes how the Phase I algorithm works. Parts of the ideas are taken from [3]. We will provide a variation of the Phase I algorithm proposed there. We start by explaining Phase I and formulating the original problem setup. In the second section, the original problem setup is casted to fit our PhaseII framework.

Original Problem Formulation

Phase I is needed if the provided vector z does not satisfy Pz < h, a requirement arising from the log-barrier term in (3). Hence, the goal of Phase I is to find a z that satisfies the (strict) inequality Pz < h. Subsequently, the existence of such a point is assumed to exist. One way to compute such a point is to introduce a variable $s \in \mathbb{R}$ and solve the following linear program (LP) [3]:

$$\begin{aligned} \min_{z,s} & s \\ \text{s.t.} & Cz = b \\ & (Pz - h)_i \leq s \quad \forall i \end{aligned}$$

Assuming the polyhedron is not empty, then we can find a s < 0. The above problem can be solved using a LP-solver. However, we aspire to transform the equation above to our framework to exploit the structure of the problem. Simple augmentation of z and s to \hat{z} (in this case, the refers to Phase I and not to the state estimate) will not work because it will destroy the structure specific to our problem.

Modified Problem Setup

We introduce 2N+1 new variables s_0, \ldots, s_{2N} , where N is the prediction horizon. So, the aim is to solve the following optimization problem:

$$\min_{z,s} \qquad s_0 + s_{2N}$$
 s.t.
$$Cz = b$$

$$s_0 = s_1 = \ldots = s_{2N-1}$$

$$(Pz - h)_j \le s_i, \qquad \forall j \in \mathcal{J}_i, \forall i,$$

where $\mathcal{J}_i \subset \mathbb{N}$ is some set. We augment the vector z by $s \in \mathbb{R}^{2N+1}$ in a specific way and denote the augmented vector by \hat{z} . The new optimization problem is

$$\begin{aligned} & \min_{\hat{z}} & s_0 + s_{2N} \\ & \text{s.t.} & & \hat{C}\hat{z} = \hat{b} \\ & & & & \hat{P}\hat{z} - h \leq \hat{s}. \end{aligned} \tag{4}$$

The $\hat{}$ -matrices shall not be defined in more detail. To use our algorithm, we regularize (4) by adding a small quadratic term and get:

$$\min_{\hat{z}} \qquad \epsilon_{II} \hat{z}^T \hat{z} + w_{II} \cdot \hat{g} \hat{z}
\text{s.t.} \qquad \hat{C} \hat{z} = \hat{b}
\qquad \hat{P} \hat{z} - h \leq \hat{s}.$$
(5)

A PHASE I 10

The parameters ϵ_{II} and w_{II} correspond to reg_PhaseI and weight_PhaseI in Tab. 4, respectively. Since (5) corresponds to (2) and both problems have similar structures, the same interior point algorithm as in Phase II can be used.

Problems:

• Regularization does not guarantee that our the solution of the regularized QP will converge to the solution of original LP, hence making (5) a heuristic algorithm.

B Example Files

B.1 Init.m

60 $Fx\{1\} = [H; -H];$

```
1 %% Init.m
 2 % Writes relevant data to binary file.
 3 % author: Xiaojing ZHANG
 4 % date: October 17, 2011
 7 clc;
 8 clear all;
9 format('short');
11 %% MPC parameters:
12 N = 10;
                                 % MPC horizon
                                  % # input
13 \text{ m} = 2;
                                  % # states
14 n = 5;
15
16 %% Parameters for constructor
17 kappa_start_PhaseI = 1000; % barrier parameter for PhaseI - can be as high as 1e10
18 kappa_start_PhaseII = 1e9; % barrier parameter for PhaseII -
19 mu = 1/10; % decrease parameter of kappa, i.e. kappa := mu∗kappa
20 eps_nt = 0.1; % tolerance for residua norm([r_p; r_d]) in Newton method
21 eps_normRp = 0.1; % tolerance for primal residua norm(r_p)
22 eps_barrier = 0.1; % barrier parameter, suboptimality of the solution
23 eps_ls = 1e-7; % smallest t, s.t. z+=z+t*dz, nu+=nu+t*dnu
24 n_iter_PhaseI = 30; % maximum number of Newton iterations for fixed kappa in PhaseI
25 n.iter_PhaseII = 10; % maximum number of Newton iterations for a fixed kappa in PhaseII
26 alpha_ls = 0.01; % alpha parameter in line search, (0.01, 0.3)
27 beta_ls = 0.5; % 0.1 < beta_ls < 0.8
28 reg_PhaseI = 1e-6; % regularization Term in PhaseI
29 reg_PhaseII = 1e-6; % regularization Term in PhaseII
30 weight_PhaseI = 1e3; % weight for linear cost (i.e. the original PhaseI problem)
32
33 %% System dynamic parameters
35 A = [1 0 1.2 1.3 1]
               0.5 2.1 1 1 -0.3
                1 1 .2 1 -2
37
                0 1 0.3 1.4 -2
38
               0.4 -0.9 2 1.2 -.4];
39
40
41 B = [1 0]
         1.3 1
42
                0 1.2
43
                -0.11
44
             0.2 -1];
45
47 s = [0; 2; 1.4; 2; 1];
 49 \quad \text{K} = -[ \quad -0.687725010189527 \quad \quad 1.970370349984470 \quad -0.865901978685416 \quad \\ -3.069636538756281 \quad \quad 2.096473307971948 \quad \\ -3.069636538756281 \quad \\ -3.069636756281 \quad \\ -3.06963676281 \quad \\ -3.0696376281 \quad \\ -3.06967
            0.181027584433678 1.040671203681152 -0.344287251091615 0.362844179335401 -1.109614558033092];
52 %% cost and constraint matrices
0 Q_tilde = 1 \times eye(n);
54 Q_tilde_f = Q_tilde+1;
56 R = 1 \times \text{eye} (m);
58 % constraint matrices: constrained on
59 H = eye(n); k = 1000*ones(n,1);
```

```
61 \text{ Fx}\{2\} = [H; -H];
 62 \text{ Fx}{3} = [H; -H];
 63 Fx{4} = [H; -H];
 64 \text{ Fx}\{5\} = [H; -H];
 65 \text{ Fx}\{6\} = [H; -H];
 66 Fx\{7\} = [H; -H];
 67 \text{ Fx}\{8\} = [H; -H];
 68 \text{ Fx}{9} = [H; -H];
 69 Fx\{10\} = [H ; -H];
 70 \text{ fx}\{1\} = [k ; k]-3;
 fx(2) = [k ; k] - 0;
 72 \text{ fx}{3} = [k; k];
 73 fx{4} = [k ; k];
 74 \text{ fx}{5} = [k; k]-2;
 75 fx\{6\} = [k ; k]+3;
 76 \text{ fx}\{7\} = [k; k]+4;
 77 fx\{8\} = [k ; k]+2;
 78 fx{9} = [k ; k]-10;
 79 fx\{10\} = [k ; k]+10;
 80
 81
 82 H = eye(m); k = 100*ones(m,1);
 83 Fu\{1\} = [H; -H];
 84 \text{ Fu}\{2\} = [H; -H];
 85 \text{ Fu}\{3\} = [H; -H];
 86 \text{ Fu}\{4\} = [H; -H];
 87 \text{ Fu}{5} = [H; -H];
 88 \text{ Fu}\{6\} = [H; -H];
 89 \text{ Fu}\{7\} = [H; -H];
 90 Fu\{8\} = [H; -H];
 91 \text{ Fu}\{9\} = [H; -H];
 92 \text{ Fu}\{10\} = [H; -H];
 93 fu\{1\} = [k ; k+20]+1;
 94 \text{ fu}{2} = [k+4; k]-0;
95 fu\{3\} = [k ; k]+5;
 96 \text{ fu}{4} = [k; k]+10;
 97 \text{ fu}{5} = [k ; k-10];
98 fu\{6\} = [k ; k]+2;
    fu{7} = [k ; k]-7;
100 fu\{8\} = [k ; k]+10;
101 fu{9} = [k ; k]-10;
102 \text{ fu}\{10\} = [k+10; k];
103
104
_{105} F_xTheta = [ 1 1 1 0 0
                 -1 -1 -1 0 0
                   0 0 0 1 1
107
                   0 0 0 -1 -1
108
                   1 0 1 0 0
109
                  -1 0 -1 0 0
110
                   0 0 0 1 0
                   0 0 0 -1 0
112
                   0 0 -1 1 1
                   0 0 1 -1 -1];
114
115 F_{theta} = [1 0]
               -1 0
116
                  0 1
117
                 0 -1
                 0 1
119
                 0 -1
120
                  1 0
121
                  -1
122
                      0
123
                  0 1
                  0 - 1;
124
f_xTheta = 100 \times [20 \ 20 \ 20 \ 30 \ 30 \ 40 \ 40 \ 50 \ 50]';
```

B.2 mainLBmpcTP.cpp

```
1 // mainLBmpcTP.cpp
2 // example file to test simple examples
3 // date: October 28, 2011
4 // author: Xiaojing ZHANG
5 //
6 // horizon: N = 4
7 // states: n = 5
8 // input: m = 2;
10 // matrices are imported from binary file created by MATLAB
11
13 #include <iostream>
                       // I-O
14 #include <fstream>
                       // read binary data
15 #include <Eigen/Dense> // matrix computation
16 #include "LBmpcTP.h" // class template
18 using namespace Eigen;
19 using namespace std;
20
21 int main()
22 {
      // ----- SPECIFY parameters -----
23
      const int _{N} = 10; // MPC horizon
      const int _m = 2;
                           // #input
25
    26
27
28
    const int _nF_xTheta = 10; // # Omega constraints
      const int _pos_omega = 4;  // < _N</pre>
30
      // ----- SPECIFY sizes of matrices -----
32
    Matrix<double, _n , _n> A_arg; // n x n
33
                                   // n x m; resizng for non-square matrices doesn't work // n x 1
     Matrix<double, _n, _m> B_arg;
34
     Matrix<double, _n, 1> s_arg;
35
      Matrix<double, _n, _n> Q_tilde_arg; // n x n
     Matrix<double, _n, _n> Q_tilde_f_arg; // n x n
37
     Matrix<double, _m, _m> R_arg;
                                          // m x m
     Matrix<double, _m, _n> K_arg;
39
                                           // _nSt x n, [_N]
40
41
                                         // _nInp x m, [_N]
// _nInp x 1, [_N]
     Matrix<double, _nInp, _m> Fu_arg[_N];
42
      Matrix<double, _nInp, 1> fu_arg[_N];
43
44
      45
46
      Matrix<double, _nF_xTheta, 1> f_xTheta_arg; // _nF_xTheta x 1
47
      Matrix<double, _n, _n> Lm_arg;
                                      // n x n
49
     Matrix<double, _n, _m> Mm_arg;
Matrix<double, _n, 1> tm_arg;
50
                                      // n x m
                                      // n x 1
51
                                      // n x 1, state estimate
      Matrix<double, _n, 1> x_hat_arg;
52
      Matrix<double, _n, 1> x_star_arg[_N]; // n x 1, [_N], tracking
      Matrix<double, _m, 1> u_opt;
                                         // m x 1, optimal input is saved there
54
      // ----- no changes necessary -----
57
      double kappa_arg; // for PhaseII
      double kappa_PhaseI_arg;  // for PhaseI
59
      int n_iter_arg;
      int n_iter_PhaseI_arg;
61
     double mu_arg;
      double eps_barrier_arg;
```

```
double eps_nt_arg;
        double eps_normRp_arg;
65
        double eps_ls_arg;
66
        double alpha_ls_arg;
67
        double beta_ls_arg;
        double reg_arg;
                            // regularization term for PhaseII
69
        double reg_PhaseI_arg; // regularization term for PhaseI
70
        double weight_PhaseI_arg; // weight for linear cost in PhaseI
71
72
        // ----- read from binary file -----
73
        ifstream fin;
                                    // Definition input file object
74
        fin.open("ConstrParam.bin", ios::binary); // open file
75
76
        if (!fin.is_open())
77
            cout << "File open error \n";</pre>
78
            return 1;
79
        }
81
82
83
        fin.read((char *) &kappa_arg, sizeof(double));
        fin.read((char *) &kappa_PhaseI_arg, sizeof(double));
84
        fin.read((char *) &n_iter_PhaseI_arg, sizeof(int));
        fin.read((char *) &n_iter_arg, sizeof(int));
86
        fin.read((char *) &mu_arg, sizeof(double));
87
        fin.read((char *) &eps_barrier_arg, sizeof(double));
88
        fin.read((char *) &eps_nt_arg, sizeof(double));
89
        fin.read((char *) &eps_normRp_arg, sizeof(double));
90
        fin.read((char *) &eps_ls_arg, sizeof(double));
91
        fin.read((char *) &alpha_ls_arg, sizeof(double));
92
        fin.read((char *) &beta_ls_arg, sizeof(double));
93
        fin.read((char *) &reg_arg, sizeof(double));
94
        fin.read((char *) &reg_PhaseI_arg, sizeof(double));
95
        fin.read((char *) &weight_PhaseI_arg, sizeof(double));
96
97
98
        // read A_arg
        for (int i = 0; i \le _n-1; i++)
100
101
            for (int j = 0; j \le -n-1; j++)
102
103
            {
                fin.read((char *) &A_arg(j,i), sizeof(double));
            }
105
106
107
        // read B_arg
108
        for (int i = 0; i < -m-1; i++) // #columns
110
            for (int j = 0; j \le -n-1; j++) // #rows
111
112
                fin.read((char *) &B_arg(j,i), sizeof(double));
113
            }
114
        }
115
        // read s_arg
117
        for (int i = 0; i \le -n-1; i++) // #columns
118
119
            fin.read((char *) &s_arg(i,0), sizeof(double));
120
121
122
        // read Q_tilde_arg
123
        for (int i = 0; i \le -n-1; i++) // #columns
124
125
            for (int j = 0; j \le -n-1; j++) // #rows
126
127
                fin.read((char *) &Q_tilde_arg(j,i), sizeof(double));
128
            }
129
```

```
}
130
131
         // read Q_tilde_f_arg
132
         for (int i = 0; i \le _n-1; i++) // #columns
133
134
             for (int j = 0; j \le _n-1; j++) // #rows
135
136
                 fin.read((char *) &Q_tilde_f_arg(j,i), sizeof(double));
137
             }
138
139
140
         // read R_arg
141
142
        for (int i = 0; i \le -m-1; i++) // #columns
143
             for (int j = 0; j \le _m-1; j++) // #rows
144
145
                 fin.read((char *) &R_arg(j,i), sizeof(double));
146
             }
147
         }
148
149
         // read Fx_arg[]
150
151
        for (int k = 0; k \le -N-1; k++)
152
             for (int i = 0; i \le _n-1; i++) // #columns
153
154
                 for (int j = 0; j \le _nSt-1; j++) // #rows
155
156
                      fin.read((char *) &Fx_arg[k](j,i), sizeof(double));
157
158
             }
159
        }
160
161
         // read fx_arg[]
162
         for (int k = 0; k \le -N-1; k++)
163
164
             for (int i = 0; i \le _nSt-1; i++)
                                                  // #columns
165
166
                      fin.read((char *) &fx_arg[k](i,0), sizeof(double));
167
168
        }
169
170
         // read Fu_arg[]
171
172
         for (int k = 0; k \le N-1; k++)
173
             for (int i = 0; i < _m-1; i++) // #columns
174
             {
                 for (int j = 0; j \leq _nInp-1; j++) // #rows
176
177
178
                      fin.read((char *) &Fu_arg[k](j,i), sizeof(double));
179
180
             }
181
182
         // read fu_arg[]
183
         for (int k = 0; k \le -N-1; k++)
184
185
             for (int i = 0; i \le \_nInp-1; i++) // #columns
186
187
             {
                      fin.read((char *) &fu_arg[k](i,0), sizeof(double));
188
             }
189
        }
190
191
192
         // read F_xTheta_arg
         for (int i = 0; i \le -n-1; i++) // #columns
193
194
             for (int j = 0; j \leq _nF_xTheta-1; j++) // #rows
195
```

```
{
196
                  fin.read((char *) &F_xTheta_arg(j,i), sizeof(double));
197
             }
198
         }
199
         // read F_theta_arg
201
         for (int i = 0; i \le _m-1; i++) // #columns
202
203
             for (int j = 0; j < _nF_xTheta-1; j++) // #rows
204
205
                  fin.read((char *) &F_theta_arg(j,i), sizeof(double));
206
             }
208
         }
209
         // read f_xTheta_arg
210
         for (int i = 0; i \le _nF_xTheta-1; i++) // #columns
211
             fin.read((char *) &f_xTheta_arg(i,0), sizeof(double));
213
214
215
         // read K_arg
216
         for (int i = 0; i \le -n-1; i++) // #columns
217
218
             for (int j = 0; j \le _m-1; j++) // #rows
220
                  fin.read((char *) &K_arg(j,i), sizeof(double));
221
222
         }
223
224
         fin.close();
                                        // close file
225
226
227
         cout << "kappa_arg: " << kappa_arg << endl << endl;</pre>
228
         cout << "kappa_PhaseI_arg " << kappa_PhaseI_arg << endl << endl;</pre>
         cout << "n_iter_arg: " << n_iter_arg << endl << endl;</pre>
230
         cout << "n_iter_PhaseI_arg: " << n_iter_PhaseI_arg << endl << endl;</pre>
231
         \verb"cout << "mu\_arg: " << mu\_arg << endl << endl;
232
         cout << "eps_barrier_arg: " << eps_barrier_arg << endl << endl;</pre>
233
         cout << "eps_nt_arg: " << eps_nt_arg << endl << endl;</pre>
234
         cout << "eps_normRp_arg: " << eps_normRp_arg << endl << endl;</pre>
235
         cout << "eps_ls_arg: " << eps_ls_arg << endl << endl;</pre>
         cout << "alpha_ls_arg: " << alpha_ls_arg << endl << endl;</pre>
237
         cout << "beta_ls_arg: " << beta_ls_arg << endl << endl;</pre>
238
         cout << "n_iter_arg: " << n_iter_arg << endl << endl;</pre>
239
         cout << "reg_arg: " << reg_arg << endl << endl;</pre>
240
         cout << "reg_PhaseI_arg: " << reg_PhaseI_arg << endl << endl;</pre>
         \verb|cout| << "weight_PhaseI_arg: " << weight_PhaseI_arg << endl << endl;
242
         cout << "A_arg: " << endl << A_arg << endl << endl;</pre>
243
         cout << "B_arg: " << endl << B_arg << endl << endl;</pre>
244
         cout << "s_arg:" << endl << s_arg << endl << endl;</pre>
245
         \verb|cout| << "Q-tilde-arg: " << endl << Q-tilde-arg << endl << endl;
246
         cout << "Q_tilde_f_arg: " << endl << Q_tilde_f_arg << endl << endl;</pre>
247
         cout << "R_arg: " << endl << R_arg << endl << endl;</pre>
249
         for (int i = 0; i \le -N-1; i++)
250
251
         {
             cout << "Fx_arg[" << i << "]: " << endl << Fx_arg[i] << endl << endl;
252
253
         for (int i = 0; i \le -N-1; i++)
254
255
         {
             cout << "fx_arg[" << i << "]: " << endl << fx_arg[i] << endl << endl;</pre>
256
257
         for (int i = 0; i \le -N-1; i++)
259
             cout << "Fu_arg[" << i << "]: " << endl << Fu_arg[0] << endl << endl;</pre>
261
```

LITERATUR 18

```
for (int i = 0; i \le N-1; i++)
262
263
        {
            cout << "fu_arg[" << i << "]: " << endl << fu_arg[i] << endl << endl;</pre>
264
265
        cout << "F_xTheta_arg: " << endl << F_xTheta_arg << endl << endl;</pre>
266
        cout << "F_theta_arg: " << endl << F_theta_arg << endl << endl;</pre>
267
        cout << "f_xTheta_arg: " << endl << f_xTheta_arg << endl << endl;</pre>
268
        cout << "K_arg:" << endl << K_arg << endl << endl;
269
270
271
        // ----- object instantiation -----
272
         LBmpcTP<double, _n, _m, _N, _nSt, _nInp, _nF_xTheta, _pos_omega> myObj(
                                                                                                      // constructor
273
274
                                 kappa_arg, kappa_PhaseI_arg, n_iter_arg, n_iter_PhaseI_arg, mu_arg,
    eps_barrier_arg, eps_nt_arg, eps_normRp_arg, eps_ls_arg,
275
                                  alpha_ls_arg, beta_ls_arg, reg_arg, reg_PhaseI_arg, weight_PhaseI_arg, A_arg, B_arg, Q_
                                  fx_arg, Fu_arg, fu_arg, F_xTheta_arg, F_theta_arg, f_xTheta_arg, K_arg, s_arg);
276
277
        // ----- SPECIFY arguments for step() -----
278
        // ----- they are updated at each time step -----
279
280
        Lm_{arg} << 1, 2, 0, 1, 2,
281
              -2, 1.3, 1, 2, 2.3,
               1, 2, -1, 0, -1,
1, 2, 2, -2.3, 1,
283
284
              0, 0, 2, 1.4, -2;
285
286
        Mm\_arg \ll 1, 1.4,
287
           2, -1,
288
           1,
               2,
           Ο,
                0,
290
           2, -1;
291
292
        tm_arg << -1, 2, 1, 1, 2;
293
        x_{hat\_arg} << 3, 3, -2, 3, 4;
294
295
        for (int i = 0; i \le N-1; i++)
296
297
            x_star_arg[i].setZero();
298
300
301
        u_opt = myObj.step( Lm_arg, Mm_arg, tm_arg, x_hat_arg, x_star_arg);
302
        cout << "optimal input:" << endl << u_opt << endl << endl;</pre>
303
304
        return 0;
305
```

Literatur

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