
GRAMPC documentation

Release 2.3

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INTRODUCTION

This manual describes the model predictive control tool GRAMPC (gradient-based MPC - [græmp'si:]) for nonlinear continuous-time systems subject to (possibly nonlinear) state and control constraints. The optimization algorithm underlying GRAMPC consists of an augmented Lagrangian scheme in connection with a tailored gradient method. GRAMPC is implemented as C code with an additional user interface to C++, Matlab/Simulink, and dSpace. GRAMPC allows one to cope with (embedded) MPC problems of nonlinear and highly dynamical systems with sampling times in the (sub)millisecond range.

The presented framework is a fundamental revision of version 1.0 of the MPC toolbox GRAMPC¹ that was originally presented for nonlinear systems with pure input constraints. Beside “classical” nonlinear MPC, GRAMPC can be used for MPC on shrinking horizon, general optimal control problems, moving horizon estimation, and parameter optimization problems.

The documentation is outlined as follows. *Installation and structure of GRAMPC* describes the installation of GRAMPC for use in C and Matlab and gives a brief overview on the software structure. The formulation of the optimization problem is shown in *Problem formulation and implementation*. Furthermore, the chapter describes the available parameters and the implementation of the problem as C functions. *Optimization algorithm and options* summarizes the optimization algorithm and provides a detailed description of the available options. The usage of GRAMPC in C and Matlab is explained in *Usage of GRAMPC*, which includes initialization, setting of parameters and options, compiling and running as well as the interfaces to Matlab and Simulink. *Tutorials* describes several example problems illustrating the application of GRAMPC to model predictive control, optimal control and moving horizon estimation. Valuable hints for tuning the software to a specific optimization problem are given at multiple places in the documentation, see especially the description of the options in *Optimization algorithm and options*, the provided plot functions in *Plot functions* and the tutorials in the last chapter.

Note that the PDF version of this documentation provides many hyperlinks to quickly jump to the definition of parameters, options and functions. In addition, the descriptions of parameters and options are repeated in the *Appendix*.

¹ B. Käpernick and K. Graichen. The gradient based nonlinear model predictive control software GRAMPC. In *Proceedings of the European Control Conference (ECC)*, 1170–1175. Strasbourg (France), 2014.

CHANGELOG

2.1 Changes in version 2.3 of GRAMPC:

Major changes:

- Added `grampc.param` as an additional input for the `probfct` signatures
 - In addition the parameter ordering of `dfdx_vec`, `dfdu_vec`, `dfdp_vec` and `dHdxdt` changed
 - `xdes` and `udes` are no longer passed to the cost functions as they are available inside `grampc.param`
- The explicit time dependency of the system dynamics was removed. The dynamics are now called with the MPC internal time. If the global time is needed, one can access `t0` in the `grampc.param` struct
- Added support for discrete-time systems
- Added config reader which reads a `.cfg` file for GRAMPC parameters and options
- Added `CMakeLists.txt` files for the GRAMPC library and every example
- Added online documentation
- Added gradient checker and an utility function via finite-differences for computing the gradients in the `probfct`
- Added Python interface

Minor changes:

- Added timer helper functions for Windows and Linux OS
- Added C++ example
- Added Matlab example for comparing MPC and MHE with continuous-time and discrete-time dynamics
- Simplified `grampc_run_Sfct.c`
- Bugfix: Rodas-Flags were not passed in the Matlab-Interface
- Bugfix: fixed typo in option `trapezoidal`
- Bugfix: fixed error in explicit line search strategy for problems without control optimization as, for example, MHE
- Bugfix: fixed error in Rodas integrator with analytical derivatives with respect to time (`IFCN=1` and `IDFX=1`)
- Bugfix: fixed gradients in the following examples:
 - `DAE_Integrator`: `dfdx` and `dfdxtrans` were specified row-wise instead of column-wise
 - `Crane_2D`: `dHdx_vec` was incorrect
 - `Reactor_PDE`: `dVdx` was incorrect

- VTOL: Vfct was missing a POW() statement and dfdu_vec was incorrect

2.2 Changes in version 2.2 of GRAMPC:

Major changes:

- Switch from GNU Lesser General Public License (LGPL) to BSD-3-Clause License, see LICENSE.txt
- Support for fixed-size arrays instead of dynamic memory allocation, see Section 5.1.5 in documentation
- Added userparam as additional input to Simulink models
- Function for estimation of PenaltyMin is exposed in C++ interface

Minor changes:

- Bugfix: Consider scaling for calculating terminal constraints in grampc_update_plot_pred.m
- Bugfix: Computation of augmented cost function correctly adds equality constraints
- Bugfix: Removed wrong flag “-o” from “ar” commands and added “-rcs” flags, which fixes problems on Windows and MacOS
- Bugfix: Provide two output arguments to calls of grampc_estim_penmin_Cmex as expected by the function
- Bugfix: OCP solution with startOCP does not abort after one iteration if convergence check is deactivated

INSTALLATION AND STRUCTURE OF GRAMPC

The following subsections describe the installation procedure of GRAMPC for use in C and Matlab. *Structure of GRAMPC* presents the principal structure of the toolbox.

3.1 Installation of GRAMPC for use in C with Cygwin

A convenient way to use GRAMPC in C/C++ under MS Windows is the Linux environment Cygwin. To install Cygwin, download the setup file from the web page <http://www.cygwin.com/> and follow the installation instructions. In the installation process when packages can be selected, you have to choose the gcc compiler and make. If Cygwin is properly installed, open a Cygwin terminal and perform the following steps:

1. Download the current version of GRAMPC from <https://github.com/grampc/grampc>.
2. Unpack the archive to an arbitrary location on your computer. After the unpacking procedure, a new directory with the following subfolders is created:
 - **cpp**: Interface of GRAMPC to C++.
 - **doc**: Contains this GRAMPC documentation.
 - **examples**: This folder contains several executable MPC, MHE and OCP problems as well as templates for implementing your own problems.
 - **include**: The header files of the GRAMPC project are located in this folder.
 - **libs**: This folder is only available after compiling the GRAMPC toolbox and contains the GRAMPC library.
 - **matlab**: Interface of GRAMPC to Matlab/Simulink, also see *Installation of GRAMPC for use in Matlab*.
 - **src**: The source files of the GRAMPC project are located in this folder.

The GRAMPC directory additionally contains a makefile for building the GRAMPC toolbox. For the remainder of this manual, the location of the created GRAMPC folder will be denoted by `<grampc_root>`.

3. Compile GRAMPC by running the following commands in a terminal:

```
$ cd <grampc_root>
$ make clean all
```

The dollar symbol indicates the line prompt of the terminal. The make command compiles the source files and generates the GRAMPC library within `<grampc_root>/libs`, which can now be used to solve a suitable problem in C. The additional argument `clean all` removes previously installed parts of GRAMPC.

3.2 Installation of GRAMPC for use in C/C++ with CMake

GRAMPC also supplies `CMakeLists.txt` files for building the GRAMPC library and the examples. From the command line, if `cmake` is installed, type

```
mkdir build
cd build
cmake ../
cmake --build .
```

and the toolbox and the examples are built. If only the toolbox shall be build, one can issue

```
cmake --build . --target grampc
```

which only compiles the GRAMPC library.

3.3 Installation of GRAMPC for use in Matlab

GRAMPC requires a C compiler that is supported by Matlab for a direct use. Details on supported compilers for the current Matlab version as well as previous releases can be found via the Mathworks homepage. The correct linkage of the compiler to Matlab can be checked by typing

```
>> mex -setup
```

in the Matlab terminal window and subsequently selecting the corresponding C compiler. The symbol `>>` denotes the Matlab prompt. The GRAMPC installation under Matlab proceeds in two steps:

1. After downloading and unpacking GRAMPC as described in *Installation of GRAMPC for use in C with Cygwin*, go to the Matlab directory

```
>> cd <grampc_root>/matlab
```

which contains the following subfolders:

- **bin**: This folder is only available after compiling the GRAMPC toolbox and contains the object files of GRAMPC.
- **include**: The header files of the GRAMPC project for the Matlab interface.
- **mfiles**: Various auxiliary functions for the Matlab interface.
- **src**: C sources files of the Mex files which provide the interface between GRAMPC and Matlab.

In addition, the subfolder contains the m-file `make.m` to start the building process.

2. Build the necessary object files for GRAMPC by executing the `make` function. The compilation of the source files can be performed with the following options:
 - `>> make clean` removes all previously built GRAMPC files,
 - `>> make` creates the necessary object files to use GRAMPC,
 - `>> make verbose` the object files are created in verbose mode, i.e. additional information regarding the building process are provided during the compilation,
 - `>> make debug` the debug option creates the object files with additional information for use in debugging,
 - `>> make debug verbose` activates the debug option as well as the verbose mode.

Similar to the compiling procedure in C as described in *Installation of GRAMPC for use in C with Cygwin*, the `make` command compiles the source files to generate object files within `<grampc root>/matlab/bin`, which can now be used to solve a suitable problem in Matlab.

3.4 Installation of GRAMPC for use in Python

Added in version v2.3.

The Python interface of GRAMPC uses pybind11 <https://github.com/pybind/pybind11> and Eigen 3.4 https://eigen.tuxfamily.org/index.php?title=Main_Page. First, download Eigen 3.4 and follow the steps in INSTALL, so its header files are available through the `find_package()` command in CMake. Make sure the run the installation for Eigen with administration rights. Then, the interface is installed through

```
pip install .
```

assuming you are within `<grampc root>`, or directly from github with

```
pip install git+https://github.com/grampc/grampc .
```

The Python interface can then be imported with

```
import pygrampc
```

Attention

If working on Windows, make sure to use Microsoft Visual Studio Compiler (MSVC), since Python for Windows is compiled with MSVC.

3.5 Structure of GRAMPC

The aim of GRAMPC is to be portable and executable on different operating systems and hardware devices without the use of external libraries. After the installation procedure as described in *Installation of GRAMPC for use in C with Cygwin* and *Installation of GRAMPC for use in Matlab*, the GRAMPC structure shown in Fig. 3.5.1 is available to cope with problems from optimal control, model predictive control, moving horizon estimation, and parameter optimization. As illustrated in Fig. 3.5.1, the GRAMPC project is implemented in plain C with a user-friendly interface to C++, Matlab/Simulink, and dSpace.

A specific problem can be implemented in GRAMPC using the C template `probfc_t_TEMPLATE.c` included in the folder `<grampc root>/examples/TEMPLATES`. A more detailed discussion about this step can be found in *Problem formulation and implementation*. The workspace of a GRAMPC project as well as algorithmic options and parameters are stored by the structure variable `grampc`. While several parameter settings are problem specific and need to be provided, most values are set to their respective default value, see *Optimization algorithm and options*. The GRAMPC structure can be manipulated through a generic interface, e.g. in order to set algorithmic options or parameters for a specific problem without the need to recompile the `grampc` project every time.

A specific example contained in the folder `<grampc root>/examples` can be compiled in C as well as in Matlab and linked against the GRAMPC toolbox. A more detailed discussion on this step can be found in *Usage of GRAMPC*.

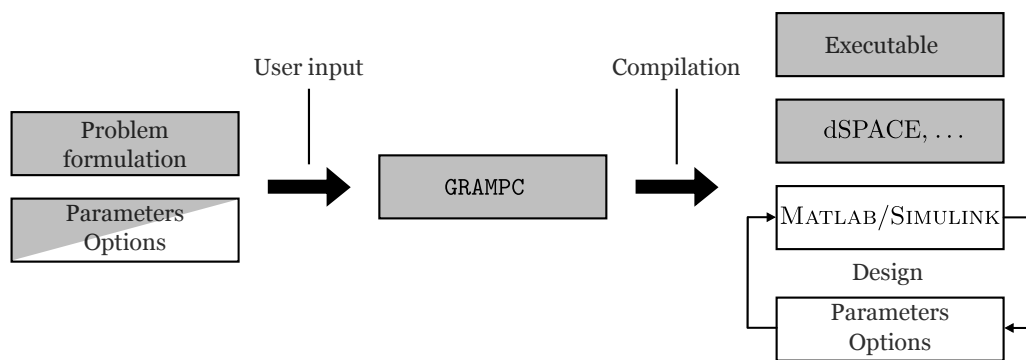


Fig. 3.5.1: General structure of GRAMPC (gray - C code, white - Matlab code).

PROBLEM FORMULATION AND IMPLEMENTATION

GRAMPC provides a solver for nonlinear input and state constrained optimal control problems. It is in particular tailored to the application of real-time MPC with a moving or shrinking horizon with focus on a memory and time efficient implementation. Other applications concern the problem of moving horizon estimation and parameter estimation.

This chapter describes the underlying optimization problem and how it can be implemented for a specific application in GRAMPC. In addition, the parameter structure of GRAMPC is introduced.

4.1 Optimization problem and parameters

GRAMPC allows one to cope with optimal control problems of the following type

$$\begin{aligned}
 \min_{\mathbf{u}, \mathbf{p}, T} \quad & J(\mathbf{u}, \mathbf{p}, T; \mathbf{x}_0) = V(\mathbf{x}(T), \mathbf{p}, T) + \int_0^T l(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) dt \\
 \text{s.t.} \quad & \mathbf{M}\dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t), \quad \mathbf{x}(t_0) = \mathbf{x}_0 \\
 & \mathbf{g}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) = \mathbf{0}, \quad \mathbf{g}_T(\mathbf{x}(T), \mathbf{p}, T) = \mathbf{0} \\
 & \mathbf{h}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) \leq \mathbf{0}, \quad \mathbf{h}_T(\mathbf{x}(T), \mathbf{p}, T) \leq \mathbf{0} \\
 & \mathbf{u}(t) \in [\mathbf{u}_{\min}, \mathbf{u}_{\max}] \\
 & \mathbf{p} \in [\mathbf{p}_{\min}, \mathbf{p}_{\max}], \quad T \in [T_{\min}, T_{\max}]
 \end{aligned} \tag{4.1.1}$$

in the context of model predictive control, moving horizon estimation, and/or parameter estimation. The cost functional $J(\mathbf{u}, \mathbf{p}, T; \mathbf{x}_0)$ to be minimized consists of the continuously differentiable terminal cost (Mayer term) $V : \mathbb{R}^{N_x} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}$ and integral cost (Lagrange term) $l : \mathbb{R}^{N_x} \times \mathbb{R}^{N_u} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}$ with the state variables $\mathbf{x} \in \mathbb{R}^{N_x}$, the control variables $\mathbf{u} \in \mathbb{R}^{N_u}$, the parameters $\mathbf{p} \in \mathbb{R}^{N_p}$, and the end time $T \in \mathbb{R}$.

The cost functional $J(\mathbf{u}, \mathbf{p}, T; \mathbf{x}_0, t_0)$ is minimized with respect to the optimization variables $(\mathbf{u}, \mathbf{p}, T)$ subject to the system dynamics $\mathbf{M}\dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t)$ with the mass matrix $\mathbf{M} \in \mathbb{R}^{N_x \times N_x}$, the continuously differentiable right hand side $\mathbf{f} : \mathbb{R}^{N_x} \times \mathbb{R}^{N_u} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}^{N_x}$, and the initial state \mathbf{x}_0 . GRAMPC allows one to formulate terminal equality and inequality constraints $\mathbf{g}_T : \mathbb{R}^{N_x} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}^{N_{g_T}}$ and $\mathbf{h}_T : \mathbb{R}^{N_x} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}^{N_{h_T}}$ as well as general equality and inequality constraints $\mathbf{g} : \mathbb{R}^{N_x} \times \mathbb{R}^{N_u} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}^{N_g}$ and $\mathbf{h} : \mathbb{R}^{N_x} \times \mathbb{R}^{N_u} \times \mathbb{R}^{N_p} \times \mathbb{R} \rightarrow \mathbb{R}^{N_h}$. In addition, the optimization variables are limited by the box constraints $\mathbf{u}(t) \in [\mathbf{u}_{\min}, \mathbf{u}_{\max}]$, $\mathbf{p} \in [\mathbf{p}_{\min}, \mathbf{p}_{\max}]$ and $T \in [T_{\min}, T_{\max}]$.

The terminal cost V , the integral cost l , as well as the system dynamics \mathbf{f} and all constraints $(\mathbf{g}, \mathbf{g}_T, \mathbf{h}, \mathbf{h}_T)$ contain an explicit time dependency with regard to the internal time $t \in [0, T]$. In the context of MPC, the internal time is distinguished from the global time $t_0 + t \in [t_0, t_0 + T]$ where the initial time t_0 and initial state \mathbf{x}_0 correspond to the sampling instant t_k that is incremented by the sampling time $\Delta t > 0$ in each MPC step k .

A detailed description about the implementation of the optimization problem (4.1.1) in C code is given in [Problem implementation](#). In addition, some parts of the problem can be configured by parameters (cf. the GRAMPC data structure `param`) and therefore do not require repeated compiling. A list of all parameters with types and allowed

values is provided in the appendix (Table 8.1.1). Except for the horizon length `Thor` and the sampling time `dt`, all parameters are optional and initialized to default values. A description of all parameters is as follows:

- `x0`: Initial state vector $x(t_0) = x_0$ at the corresponding sampling time t_0 .
- `xdes`: Desired (constant) setpoint vector for the state variables x .
- `u0`: Initial value of the control vector $u(t) = u_0 = \text{const.}, t \in [0, T]$ that is used in the first iteration of GRAMPC.
- `udes`: Desired (constant) setpoint vector for the control variables u .
- `umin, umax`: Lower and upper bounds for the control variables u .
- `p0`: Initial value of the parameter vector $p = p_0$ that is used in the first iteration of GRAMPC.
- `pmin, pmax`: Lower and upper bounds for the parameters p .
- `Thor`: Prediction horizon T or initial value if the end time is optimized.
- `Tmin, Tmax`: Lower and upper bound for the prediction horizon T .
- `dt`: Sampling time Δt of the considered system for model predictive control or moving horizon estimation. Required for prediction of next state `grampc.sol.xnext` and for the control shift, see [Control shift](#).
- `t0`: Current sampling instance t_0 that is provided in the `grampc.param` structure.
- `userparam`: Further problem-specific parameters, e.g. system parameters or weights in the cost functions that are passed to the problem functions via a void-pointer in C or `typeRNum` array in MATLAB.

Although GRAMPC uses a continuous-time formulation of the optimization problem (4.1.1), all trajectories are internally stored in discretized form with `Nhor` steps (cf. [Numerical Integration](#)). This raises the question of whether all constraints are evaluated for the last trajectory point or only the terminal ones. In general, the constraints should be formulated in such a way that there are no conflicts. However, numerical difficulties can arise in some problems if constraints are formulated twice for the last point. Therefore, GRAMPC does not evaluate the constraints g and h for the last trajectory point if terminal constraints are defined, i.e. $N_{g_T} + N_{h_T} > 0$. In contrast, if no terminal constraints are defined, the functions g and h are evaluated for all points. Note that the opposite behavior is easy to implement by including g and h in the terminal constraints g_T and h_T .

4.1.1 Problem implementation

Regardless of what kind of problem statement is considered (i.e. model predictive control, moving horizon estimation or parameter estimation), the optimization problem (4.1.1) must be implemented in C, cf. [Fig. 3.5.1](#). For this purpose, the C file template `probfc_t_TEMPLATE.c` is provided within the folder `<grampc_root>/examples/TEMPLATE`, which allows one to describe the structure of the optimization problem (4.1.1), the cost functional J to be minimized, the system dynamics f , and the constraints g, g_T, h, h_T . The number of C functions to be provided depends on the type of dynamics f of the specific problem at hand.

Problems involving explicit ODEs

A special case of the system dynamics f and certainly the most important one concerns ordinary differential equations (ODEs) with explicit appearance of the first-order derivatives

$$\dot{x}(t) = f(x(t), u(t), p, t),$$

corresponding to the identity mass matrix $M = I$ in OCP (4.1.1). In this case, the following functions of the C template file `probfc_t_TEMPLATE.c` have to be provided:

- **ocp_dim**: Definition of the dimensions of the considered problem, i.e. the number of state variables N_x , control variables N_u , parameters N_p , equality constraints N_g , inequality constraints N_h , terminal equality constraints N_{g_T} , and terminal inequality constraints N_{h_T} .
- **ffct**: Formulation of the system dynamics function f .
- **dfdx_vec**, **dfdx_vec**, **dfdp_vec**: Matrix vector products $(\frac{\partial f}{\partial x})^\top v$, $(\frac{\partial f}{\partial u})^\top v$ and $(\frac{\partial f}{\partial p})^\top v$ for the system dynamics with an arbitrary vector v of dimension N_x .
- **Vfct**, **lfct**: Terminal and integral cost functions V and l of the cost functional J .
- **dVdx**, **dVdp**, **dVdT**, **dldx**, **dldu**, **dldp**: Gradients $\frac{\partial V}{\partial x}$, $\frac{\partial V}{\partial p}$, $\frac{\partial V}{\partial T}$, $\frac{\partial l}{\partial x}$, $\frac{\partial l}{\partial u}$, and $\frac{\partial l}{\partial p}$ of the cost functions **Vfct** and **lfct**.
- **hfct**, **gfct**: Inequality and equality constraint functions h and g as defined in (4.1.1).
- **hTfct**, **gTfct**: Terminal inequality and equality constraint functions h_T and g_T as defined in (4.1.1).
- **dhdxd_vec**, **dhdxd_vec**, **dhdxd_vec**: Matrix products $(\frac{\partial h}{\partial x})^\top v$, $(\frac{\partial h}{\partial u})^\top v$, and $(\frac{\partial h}{\partial p})^\top v$ for the inequality constraints with an arbitrary vector v of dimension N_h .
- **dgdxd_vec**, **dgdxd_vec**, **dgdxd_vec**: Matrix product functions $(\frac{\partial g}{\partial x})^\top v$, $(\frac{\partial g}{\partial u})^\top v$, and $(\frac{\partial g}{\partial p})^\top v$ for the equality constraints with an arbitrary vector v of dimension N_g .
- **dhTdx_vec**, **dhTdx_vec**, **dhTdx_vec**: Matrix product functions $(\frac{\partial h_T}{\partial x})^\top v$, $(\frac{\partial h_T}{\partial p})^\top v$, and $(\frac{\partial h_T}{\partial T})^\top v$ for the terminal inequality constraints with an arbitrary vector v of dimension N_{h_T} .
- **dgTdx_vec**, **dgTdx_vec**, **dgTdx_vec**: Matrix product functions $(\frac{\partial g_T}{\partial x})^\top v$, $(\frac{\partial g_T}{\partial p})^\top v$, and $(\frac{\partial g_T}{\partial T})^\top v$ for the terminal equality constraints with an arbitrary vector v of dimension N_{g_T} .

The respective problem function templates only have to be filled in if the corresponding constraints and cost functions are defined for the problem at hand and depending on the actual choice of optimization variables (u , p , and/or T). For example, if only the control u is optimized, the partial derivatives with respect to p and T are not required.

The gradients $\frac{\partial V}{\partial x}$, $\frac{\partial V}{\partial p}$, $\frac{\partial V}{\partial T}$, $\frac{\partial l}{\partial x}$, $\frac{\partial l}{\partial u}$, and $\frac{\partial l}{\partial p}$ as well as the matrix product functions listed above appear in the partial derivatives H_x , H_u and H_p of the Hamiltonian H within the gradient method (see [Projected gradient method](#)). The matrix product formulation is chosen over the definition of Jacobians (e.g. $(\frac{\partial f}{\partial x})^\top v$ instead of $\frac{\partial f}{\partial x}$) in order to avoid unnecessary zero multiplications for sparse matrices or alternatively the usage of sparse numerics.

Problems with discrete-time systems

Added in version v2.3.

GRAMPC also allows to consider discrete-time system dynamics of the form

$$x_{k+1} = f(x_k, u_k, p, t_k)$$

with $x_k = x(t_k)$ and $u_k = u(t_k)$. For this case the same functions as in [Problems involving explicit ODEs](#) are used to implement the problem description but the option **Integrator** is set to **discrete**. Since the discrete **ffct** computes the system dynamics for a fixed sampling time Δt (Parameter **dt**), the end time T (Parameter **Thor**) of the optimization problem must satisfy

$$T = (N_{\text{hor}} - 1)\Delta t$$

with the number of discretization points along the horizon N_{hor} (Option **Nhor**). As a consequence it is no longer possible to consider a free end time, where T is an optimization variable, since this would change the sampling time Δt . Note that for continuous-time system dynamics the sampling time Δt can be chosen independently of the end time T and the number of discretization points N_{hor} , because it is only used for predicting the next state `grampc.sol.xnext` and for the control shift.

Note

For discrete-time systems the option `Integrator` is set to `discrete`, the settings `Nhor`, `Thor` and `dt` need to satisfy the relation `Thor = (Nhor-1)*dt` and the option `OptimTime` must be set to `off`.

An MPC example that compares continuous-time and discrete-time versions of a double integrator is included in `<grampc_root>/examples/Continuous_vs_Discrete`. Furthermore, a discrete-time formulation of the helicopter example is provided in `<grampc_root>/examples/Continuous_vs_Discrete`.

Problems involving semi-implicit ODEs and DAEs

Beside explicit ODEs, GRAMPC supports semi-implicit ODEs with mass matrix $M \neq I$ and DAEs with M being singular. The underlying numerical integrations of the dynamics f is carried out using the integrator RODAS¹². In this case, additional C functions must be provided and several RODAS-specific options must be set, cf. *Integration of semi-implicit ODEs and DAEs (RODAS)* and *Setting options and parameters*. Especially, the option `IMAS = 1` must be set to indicate that a mass matrix is given. The numerical integrations performed with RODAS can be accelerated by providing partial derivatives. In summary, the following additional C functions are used by GRAMPC for semi-implicit ODEs and DAE systems:

- `Mfct`, `Mtrans`: Definition of the mass matrix M and its transpose M^T , which is required for the adjoint dynamics, cf. *Projected gradient method* in the projected gradient algorithm. The matrices must be specified column-wise. If the mass matrix has a band structure, only the respective elements above and below the main diagonal are specified. This only applies if the options `IMAS = 1` and `MLJAC < N_x` are selected. Non-existent elements above or below the main diagonal must be filled with zeros so that the same number of elements is specified for each column.
- `dfdx`, `dfdxtrans`: The Jacobians $\frac{\partial f}{\partial x}$ and $(\frac{\partial f}{\partial x})^T = \frac{\partial^2 H}{\partial x \partial \lambda}$ are provided by these functions if the option `IJAC = 1` is set. This allows one to evaluate the right hand sides of the canonical equations time efficiently. The Jacobians must be implemented in vector form by arranging the successive columns for $\frac{\partial f}{\partial x}$ and $(\frac{\partial f}{\partial x})^T$. If the option `MLJAC < N_x` is set to exploit the band structure of the Jacobians, only the corresponding elements above and below the main diagonal must be specified.
- `dfdt`, `dHdxdt`: The partial derivatives $\frac{\partial f}{\partial t}$ and $\frac{\partial^2 H}{\partial x \partial t}$ allow for evaluating the right hand sides of the canonical equations time efficiently, if the problem explicitly depends on time t . These functions must only be provided if the options `IFCN = 1` and `IDFX = 1` are used.

An MPC example with a semi-implicit system dynamics is included in `<grampc_root>/examples/Reactor_PDE`. The problem formulation is derived from a quasi-linear diffusion-convection-reaction system that is spatially discretized using finite elements.

Finite differences and gradient check

Added in version v2.3.

For the purpose of rapid prototyping, the partial derivatives can also be approximated by forward finite differences. To this end, the header `finite_diff.h` provides a helper function

```
void finite_diff_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
  ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam,
  typeNum *memory, ctypeNum step_size, const typeFiniteTarget *target,
  ctypeInt func_out_size, const typeFiniteDiffFctPtr func);
```

¹ E. Hairer and G. Wanner. *Solving Ordinary Differential Equations: Stiff and Differential-Algebraic Problems*. Springer, Heidelberg, Germany, 1996.

² RODAS. Webpage. <http://www.unige.ch/~hairer/software.html>, Accessed 01-December-2018.

where the first eight arguments are the same as for typical functions in `probfc.h` and the other arguments are:

- **memory**: Sufficient user-provided memory of size at most $3 \cdot \max\{N_x, N_u, N_p, N_g, N_h, N_{gT}, N_{hT}\}$ that is needed for storing intermediate values. If used within a `probfc`, this memory should be provided via the `userparam` pointer to avoid dynamic memory allocation during the optimization.
- **step_size**: Small value used as step size for the finite differences.
- **target**: Argument with respect to which the finite differences are computed. Options are `DX` for states, `DU` for inputs, `DP` for parameters and `DT` for time.
- **func_out_size**: Size of the function's out argument, e.g. `Nh` for `hfct`.
- **func**: Pointer to the function that is differentiated. For `Vfct`, `gTfct` and `hTfct` wrappers are provided with an unused dummy argument in place of the control `u`.

The intended usage is shown in the template file `probfc_TEMPLATE_finite_diff.c`. In addition, the helper function

```
void finite_diff_dfdx(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
    const typeGRAMPCparam *param, typeUSERPARAM *userparam,
    typeNum *memory, ctypeNum step_size, ctypeInt ml, ctypeInt mu,
    typeBoolean transpose);
```

approximates the partial derivatives of the `ffct` for semi-implicit ODEs and DAEs with the arguments:

- **memory**: Sufficient allocated memory for storing intermediate values of size $3 \cdot N_x$. If used within a `probfc`, this memory should be provided via the `userparam` pointer to avoid dynamic memory allocation during the optimization.
- **step_size**: Small value used as step size for the finite differences.
- **ml**: Number of lower non-zero diagonals if banded. Set to `Nx` for full matrix.
- **mu**: Number of upper non-zero diagonals if banded. Set to `Nx` for full matrix.
- **transpose**: Set to `0` for approximating `dfdx` and to `1` for approximating `dfdxtrans`.

Finally, it is recommended to validate the user-supplied analytic gradients by comparison to finite differences. For this task, GRAMPC provides a helper function

```
void grampc_check_gradients(typeGRAMPC *grampc, ctypeNum tolerance, ctypeNum step_size);
```

that checks the gradients at the initial point defined by `t0`, `x0`, `u0` and `p0`. The tolerance is applied element-wise to the relative error

$$\left| \frac{\nabla f_{fd,i} - \nabla f_i}{\max(1, \nabla f_i)} \right|$$

where $\nabla f_{fd,i}$ is the finite difference approximation and ∇f_i the user-supplied derivative. If the tolerance is violated, the function prints messages like

```
dfdx_vec: element (out_index,vec_index) exceeds supplied tolerance with 4.234546e-5
```

so that one is able to correct the wrong derivative. Recommended settings for the gradient check include a step size of $\sqrt{\text{eps}}$ where `eps` is the floating point machine precision, and a tolerance of `1e-6`. Nevertheless, the gradient checker can produce false positives, so every flagged gradient should be carefully checked.

Example: Ball-on-plate system

The appropriate definition of the C functions of the template `probfcn_TEMPLATE` is described for the example of a ball-on-plate system³ in the context of MPC. The problem is also included in `<grampc_root>/examples(BallOnPlate)`. The underlying optimization problem reads as

$$\begin{aligned}
 \min_{u(\cdot)} \quad & J(u; \mathbf{x}_0) = \frac{1}{2} \Delta \mathbf{x}^\top(T) \mathbf{P} \Delta \mathbf{x}(T) + \frac{1}{2} \int_0^T \Delta \mathbf{x}^\top(\tau) \mathbf{Q} \Delta \mathbf{x}(\tau) + R \Delta u^2 d\tau \\
 \text{s.t.} \quad & \begin{bmatrix} \dot{x}_1 \\ \dot{x}_2 \end{bmatrix} = \begin{bmatrix} 0 & 1 \\ 0 & 0 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} + \begin{bmatrix} -0.04 \\ -7.01 \end{bmatrix} u, \quad \begin{bmatrix} x_1(0) \\ x_2(0) \end{bmatrix} = \begin{bmatrix} x_{k,1} \\ x_{k,2} \end{bmatrix} \\
 & \begin{bmatrix} -0.2 \\ -0.1 \end{bmatrix} \leq \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} \leq \begin{bmatrix} 0.01 \\ 0.1 \end{bmatrix}, \quad |u| \leq 0.0524
 \end{aligned} \tag{4.1.2}$$

The cost functional in (4.1.2) penalizes the state and input error $\Delta \mathbf{x} = \mathbf{x} - \mathbf{x}_{\text{des}}$ and $\Delta u = u - u_{\text{des}}$ in a quadratic manner using the weights

$$\mathbf{P} = \mathbf{Q} = \begin{bmatrix} 100 & 0 \\ 0 & 10 \end{bmatrix}, \quad R = 1.$$

The system dynamics in (4.1.2) describes a simplified linear model of a single axis of a ball-on-plate system³. An optimal solution of the optimization problem has to satisfy the input and state constraints present in (4.1.2).

The user must provide the C functions and to describe the general structure and the system dynamics of the optimization problem, also see *Problem implementation*. As shown in Listing 4.1.1, the C function `ocp_dim` is used to define the number of states, control inputs, and number of (terminal) inequality and equality constraints. Note that GRAMPC uses the generic type `typeInt` for integer values. The word size of this integer type can be changed in the header file `grampc_macro.h` within the folder `<grampc_root>/include`. This is particularly advantageous with regard to implementing GRAMPC on embedded hardware.

Listing 4.1.1: Settings of the general structure of optimization problem (4.1.2).

```

/** OCP dimensions */
void ocp_dim(typeInt *Nx, typeInt *Nu, typeInt *Np, typeInt *Ng, typeInt *Nh,
             typeInt *NgT, typeInt *NhT, typeUSERPARAM *userparam)
{
    *Nx = 2;
    *Nu = 1;
    *Np = 0;
    *Nh = 4;
    *Ng = 0;
    *NhT = 0;
    *NgT = 0;
}

```

The system dynamics are described by the C function `ffcn` shown in Listing 4.1.2. The example is given in explicit ODE form, for which the functions `Mfcn` and `Mtrans` for the mass matrix M are not required. Similar to the generic integer type `typeInt`, the data type `typeRNum` is used to address floating point numbers of different word sizes, e.g. float or double (cf. the header file `grampc_macro.h` included in the folder `<grampc_root>/include`).

³ S. Richter. *Computational Complexity Certification of Gradient Methods for Real-Time Model Predictive Control*. PhD thesis, ETH Zürich, 2012.

Listing 4.1.2: Formulation of the system dynamics of optimization problem (4.1.2).

```
/** System function f(t,x,u,p,grampc.param,userparam) */
void ffct(ctypeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u,
          ctypeRNum *p, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    out[0] = x[1]-0.04*u[0];
    out[1] = -7.01*u[0];
}
```

The cost functions are defined via the functions lfct and Vfct, cf. Listing 4.1.3. Note that the input argument userparam is used to parametrize the cost functional in a generic way.

Listing 4.1.3: Formulation of the cost functional of optimization problem (4.1.2).

```
/** Integral cost l(t,x(t),u(t),p,grampc.param,userparam) */
void lfct(ctypeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u,
          ctypeRNum *p, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    ctypeRNum* pCost = (ctypeRNum*)userparam;
    ctypeRNum* xdes = param->xdes;
    ctypeRNum* udes = param->udes;
    out[0] = (pCost[0] * (x[0] - xdes[0]) * (x[0] - xdes[0])
              + pCost[1] * (x[1] - xdes[1]) * (x[1] - xdes[1])
              + pCost[2] * (u[0] - udes[0]) * (u[0] - udes[0])) / 2;
}

/** Terminal cost V(T,x(T),p,grampc.param,userparam) */
void Vfct(ctypeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p,
          const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    ctypeRNum* pCost = (ctypeRNum*)userparam;
    ctypeRNum* xdes = param->xdes;
    out[0] = (pCost[3] * (x[0] - xdes[0]) * (x[0] - xdes[0])
              + pCost[4] * (x[1] - xdes[1]) * (x[1] - xdes[1])) / 2;
}
```

Listing 4.1.4 shows the formulation of the inequality constraints $h(x(t), u(t), p, t) \leq 0$. For the sake of completeness, Listing 4.1.4 also contains the corresponding functions for equality constraints $g(x(t), u(t), p, t) = 0$, terminal inequality constraints $h_T(x(T), p, T) \leq 0$ as well as terminal equality constraints $g_T(x(T), p, T) = 0$, which are not defined for the ball-on-plate example. Similar to the formulation of the cost functional, the input argument userparam is used to parametrize the inequality constraints.

Listing 4.1.4: Formulation of the state constraints of optimization problem (4.1.2).

```
/** Inequality constraints h(t,x(t),u(t),p,grampc.param,userparam) <= 0 */
void hfct(ctypeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u,
          ctypeRNum *p, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    ctypeRNum* pSys = (ctypeRNum*)userparam;
```

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```

    out[0] = pSys[5] - x[0];
    out[1] = -pSys[6] + x[0];
    out[2] = pSys[7] - x[1];
    out[3] = -pSys[8] + x[1];
}

/** Equality constraints  $g(t,x(t),u(t),p,grampc.param,userparam) = 0$  */
void gfct(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u,
          ctypeRNum *p, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
}

/** Terminal inequality constraints  $hT(T,x(T),p,grampc.param,userparam) \leq 0$  */
void hTfct(typeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p,
            const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
}

/** Terminal equality constraints  $gT(T,x(T),p,grampc.param,userparam) = 0$  */
void gTfct(typeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p,
            const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
}

```

The Jacobians of the single functions defined in Listing 4.1.2 to Listing 4.1.4 with respect to state x and control u are required for evaluating the optimality conditions of optimization problem (4.1.2) within the gradient algorithm, see *Projected gradient method*. If applicable, the Jacobians of the above-mentioned functions are also required with respect to the optimization variables p and T . Listing 4.1.5 shows the corresponding Jacobians for the ball-on-plate example. For the matrix product functions `dfdx_vec`, `dfdu_vec`, and `dhdxd_vec`, the pointer to a generic vector `vec` is passed as input argument that corresponds to the adjoint state, respectively a vector that accounts for the Lagrange multiplier and penalty term of state constraints, cf. *Optimization algorithm*. Note that `vec` is of appropriate dimension for the respective matrix product function, i.e. of dimension N_x or N_h . A complete C function template and further examples concerning the problem formulation are included in the GRAMPC software package.

Listing 4.1.5: Jacobians of the system dynamics and inequality constraint.

```

/** Jacobian  $df/dx$  multiplied by vector  $vec$ , i.e.  $(df/dx)^T * vec$  or  $vec^T * (df/dx)$  */
void dfdx_vec(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p,
              ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    out[0] = 0;
    out[1] = vec[0];
}

/** Jacobian  $df/du$  multiplied by vector  $vec$ , i.e.  $(df/du)^T * vec$  or  $vec^T * (df/du)$  */
void dfdu_vec(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p,
              ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    out[0] = (typeRNum)(-0.04)*vec[0] - (typeRNum)(7.01)*vec[1];
}

```

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```

/** Gradient dl/dx */
void dldx(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
         const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    ctypeNum* pCost = (ctypeNum*)userparam;
    ctypeNum* xdes = param->xdes;

    out[0] = pCost[0] * (x[0] - xdes[0]);
    out[1] = pCost[1] * (x[1] - xdes[1]);
}

/** Gradient dl/du */
void dlldu(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
         const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    ctypeNum* pCost = (ctypeNum*)userparam;
    ctypeNum* udes = param->udes;

    out[0] = pCost[2] * (u[0] - udes[0]);
}

/** Gradient dV/dx */
void dVdx(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p,
         const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    ctypeNum* pCost = (ctypeNum*)userparam;
    ctypeNum* xdes = param->xdes;

    out[0] = pCost[3] * (x[0] - xdes[0]);
    out[1] = pCost[4] * (x[1] - xdes[1]);
}

/** Jacobian dh/dx multiplied by vector vec, i.e. (dh/dx)^T*vec or vec^T*(dg/dx) */
void dhdx_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
              ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    out[0] = -vec[0] + vec[1];
    out[1] = -vec[2] + vec[3];
}
...

```


OPTIMIZATION ALGORITHM AND OPTIONS

This chapter summarizes the optimization scheme that is used to solve the optimization problem in *Problem formulation and implementation*. This includes the basic algorithm as well as the options that can be adjusted by the user. Note that all options as well as their corresponding default values are listed in Table *Algorithmic Options*. The setting of the options is detailed in *Usage of GRAMPC* for usage in C and Matlab.

5.1 Optimization algorithm

The optimization algorithm of GRAMPC is based on an augmented Lagrangian formulation with an inner projected gradient method as minimization step and an outer multiplier and penalty update. This section gives a brief sketch of the algorithm. Note that a more detailed description is given in¹.

5.1.1 Augmented Lagrangian method

GRAMPC implements the augmented Lagrangian approach to handle the equality and inequality constraints of the OCP. The constraints are adjoined to the integral cost function using the time-dependent multipliers $\mu = [\mu_g^\top, \mu_h^\top]^\top$ and penalties $c = [c_g^\top, c_h^\top]^\top$. Similarly, multipliers $\mu_T = [\mu_{g_T}^\top, \mu_{h_T}^\top]^\top$ and penalties $c_T = [c_{g_T}^\top, c_{h_T}^\top]^\top$ are used for the terminal constraints. Where appropriate, the syntax $\bar{\mu} = (\mu_g, \mu_h, \mu_{g_T}, \mu_{h_T})$ and $\bar{c} = (c_g, c_h, c_{g_T}, c_{h_T})$ is used to denote all multipliers and penalties. The algorithm requires a reformulation of the inequality constraints that leads to the transformed functions (see¹ for details)

$$\begin{aligned}\bar{h}(x, u, p, t, \mu_h, c_h) &= \max \{h(x, u, p, t), -C_h^{-1} \mu_h\} \\ \bar{h}_T(x, p, T, \mu_{h_T}, c_{h_T}) &= \max \{h_T(x, p, T), -C_{h_T}^{-1} \mu_{h_T}\}\end{aligned}$$

with the component-wise **max**-function and the diagonal matrix syntax $C = \text{diag}(c)$. The augmented Lagrangian function is defined as

$$\bar{J}(u, p, T, \bar{\mu}, \bar{c}; x_0) = \bar{V}(x, p, T, \mu_T, c_T) + \int_0^T \bar{l}(x, u, p, t, \mu, c) dt \quad (5.1.1)$$

with the augmented terminal cost term

$$\begin{aligned}\bar{V}(x, p, T, \mu_T, c_T) &= V(x, p, T) + \mu_{g_T}^\top g_T(x, p, T) + \frac{1}{2} \|g_T(x, p, T)\|_{C_{g_T}}^2 \\ &\quad + \mu_{h_T}^\top \bar{h}_T(x, p, T, \mu_{h_T}, c_{h_T}) + \frac{1}{2} \|\bar{h}_T(x, p, T, \mu_{h_T}, c_{h_T})\|_{C_{h_T}}^2\end{aligned}$$

¹ T. Englert, A. Völz, F. Mesmer, S. Rhein, and K. Graichen. A software framework for embedded nonlinear model predictive control using a gradient-based augmented Lagrangian approach (GRAMPC). *Optimization and Engineering*, 20(3):769–809, 2019. doi.org/10.1007/s11081-018-9417-2.

and the augmented integral cost term

$$\begin{aligned}\bar{l}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t, \boldsymbol{\mu}, \mathbf{c}) = & l(\mathbf{x}, \mathbf{u}, \mathbf{p}, t) + \boldsymbol{\mu}_g^\top \mathbf{g}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t) + \frac{1}{2} \|\mathbf{g}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t)\|_{\mathbf{C}_g}^2 \\ & + \boldsymbol{\mu}_h^\top \bar{\mathbf{h}}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t, \boldsymbol{\mu}_h, \mathbf{c}_h) + \frac{1}{2} \|\bar{\mathbf{h}}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t, \boldsymbol{\mu}_h, \mathbf{c}_h)\|_{\mathbf{C}_h}^2.\end{aligned}$$

Instead of solving the original problem, the algorithm solves the max-min-problem

$$\begin{aligned}\max_{\bar{\boldsymbol{\mu}}} \min_{\mathbf{u}, \mathbf{p}, T} \quad & \bar{J}(\mathbf{u}, \mathbf{p}, T, \bar{\boldsymbol{\mu}}, \bar{\mathbf{c}}; \mathbf{x}_0) \\ \text{s.t.} \quad & \mathbf{M} \dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t), \quad \mathbf{x}(0) = \mathbf{x}_0 \\ & \mathbf{u}(t) \in [\mathbf{u}_{\min}, \mathbf{u}_{\max}], \quad t \in [0, T] \\ & \mathbf{p} \in [\mathbf{p}_{\min}, \mathbf{p}_{\max}], \quad T \in [T_{\min}, T_{\max}],\end{aligned}\tag{5.1.2}$$

whereby the augmented Lagrangian function \bar{J} is maximized with respect to the multipliers $\bar{\boldsymbol{\mu}}$ and minimized with respect to the controls \mathbf{u} , the parameters \mathbf{p} and the end time T . Note that the full set of optimization variables $(\mathbf{u}, \mathbf{p}, T)$ is considered in what follows for the sake of completeness. The max-min-problem (5.1.2) corresponds to the dual problem of (4.1.1) in the case of $\bar{\mathbf{c}} = \mathbf{0}$. The maximization step is performed by steepest ascent using the constraint residual as direction and the penalty parameter as step size. See^{Page 21, 1} for a detailed description of the augmented Lagrangian algorithm.

5.1.2 Projected gradient method

GRAMPC uses a projected gradient method to solve the inner minimization problem subject to the dynamics $\mathbf{f}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t)$ as well as the box constraints $\mathbf{u}(t) \in [\mathbf{u}_{\min}, \mathbf{u}_{\max}]$ and $\mathbf{p} \in [\mathbf{p}_{\min}, \mathbf{p}_{\max}]$. The algorithm is based on the first-order optimality conditions that can be compactly stated using the Hamiltonian

$$H(\mathbf{x}, \mathbf{u}, \mathbf{p}, \boldsymbol{\lambda}, t, \boldsymbol{\mu}, \mathbf{c}) = \bar{l}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t, \boldsymbol{\mu}, \mathbf{c}) + \boldsymbol{\lambda}^\top \mathbf{f}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t)$$

with the adjoint states $\boldsymbol{\lambda}$. The canonical equations are then given by

$$\begin{aligned}\mathbf{M} \dot{\mathbf{x}} &= \mathbf{f}(\mathbf{x}, \mathbf{u}, \mathbf{p}, t), & \mathbf{x}(0) &= \mathbf{x}_0, \\ \mathbf{M}^\top \dot{\boldsymbol{\lambda}} &= -H_{\mathbf{x}}(\mathbf{x}, \mathbf{u}, \mathbf{p}, \boldsymbol{\lambda}, t, \boldsymbol{\mu}, \mathbf{c}), & \mathbf{M}^\top \boldsymbol{\lambda}(T) &= \bar{\mathbf{V}}_{\mathbf{x}}(\mathbf{x}(T), \mathbf{p}, T, \boldsymbol{\mu}_T, \mathbf{c}_T)\end{aligned}\tag{5.1.3}$$

consisting of the original dynamics $\dot{\mathbf{x}}$ and the adjoint dynamics $\dot{\boldsymbol{\lambda}}$. The canonical equations can be iteratively solved in forward and backward time for given initial values of the optimization variables. In each iteration and depending on the optimization variables of the actual problem to be solved, the gradients

$$\begin{aligned}d_{\mathbf{u}} &= H_{\mathbf{u}}(\mathbf{x}, \mathbf{u}, \mathbf{p}, \boldsymbol{\lambda}, t, \boldsymbol{\mu}, \mathbf{c}) \\ d_{\mathbf{p}} &= \bar{\mathbf{V}}_{\mathbf{p}}(\mathbf{x}(T), \mathbf{p}, T, \boldsymbol{\mu}_T, \mathbf{c}_T) + \int_0^T H_{\mathbf{p}}(\mathbf{x}, \mathbf{u}, \mathbf{p}, \boldsymbol{\lambda}, t, \boldsymbol{\mu}, \mathbf{c}) dt \\ d_T &= \bar{\mathbf{V}}_T(\mathbf{x}(T), \mathbf{p}, T, \boldsymbol{\mu}_T, \mathbf{c}_T) + H(\mathbf{x}(T), \mathbf{u}(T), \mathbf{p}, \boldsymbol{\lambda}(T), T, \boldsymbol{\mu}(T), \mathbf{c}(T))\end{aligned}$$

with respect to the controls \mathbf{u} , parameters \mathbf{p} , and end time T are used to formulate a line search problem

$$\min_{\alpha} \bar{J}(\psi_{\mathbf{u}}(\mathbf{u} - \alpha d_{\mathbf{u}}), \psi_{\mathbf{p}}(\mathbf{p} - \gamma_{\mathbf{p}} \alpha d_{\mathbf{p}}), \psi_T(T - \gamma_T \alpha d_T); \bar{\boldsymbol{\mu}}, \bar{\mathbf{c}}, \mathbf{x}_0)$$

with projection functions $\psi_{\mathbf{u}}$, $\psi_{\mathbf{p}}$ and ψ_T and, finally, to update the optimization variables according to

$$\mathbf{u} \leftarrow \psi_{\mathbf{u}}(\mathbf{u} - \alpha d_{\mathbf{u}}), \quad \mathbf{p} \leftarrow \psi_{\mathbf{p}}(\mathbf{p} - \gamma_{\mathbf{p}} \alpha d_{\mathbf{p}}), \quad T \leftarrow \psi_T(T - \gamma_T \alpha d_T).$$

See^{23Page 21, 1} for a detailed description of the projected gradient algorithm. GRAMPC provides two methods for the approximate solution of the line search problem, which are explained in [Line search](#).

² K. Graichen and B. Käpernick. A real-time gradient method for nonlinear model predictive control. In T. Zheng, editor, *Frontiers of Model Predictive Control*, pages 9–28. InTech, 2012.

³ B. Käpernick and K. Graichen. The gradient based nonlinear model predictive control software GRAMPC. In *Proceedings of the European Control Conference (ECC)*, 1170–1175. Strasbourg (France), 2014.

5.1.3 Algorithmic structure

i Algorithm 1 (Basic algorithmic structure of GRAMPC.)

```

Optional: Shift trajectories by sampling time  $\Delta t$ 

For  $i = 1$  to  $i_{max}$  do
    For  $j = 1$  to  $j_{max}$  do
        If  $i > 1$  and  $j = 1$  then
            Set  $\mathbf{x}^{i|j} = \mathbf{x}^{i-1}$ 
        else
            Compute  $\mathbf{x}^{i|j}$  by forward time integration of system dynamics
            Evaluate all constraints
        End If
        Compute  $\boldsymbol{\lambda}^{i|j}$  by backward time integration of adjoint system
        Evaluate gradients  $\mathbf{d}_u^{i|j}$ ,  $\mathbf{d}_p^{i|j}$  and  $\mathbf{d}_T^{i|j}$ 
        Solve line search problem to determine step size  $\alpha^{i|j}$ 
        Update controls  $\mathbf{u}^{i|j+1}$ , parameters  $\mathbf{p}^{i|j+1}$  and end time  $T^{i|j+1}$ 
        If minimization is converged then
            Break inner loop
        End If
    End For
    Set  $\mathbf{u}^i = \mathbf{u}^{i|j+1}$ ,  $\mathbf{p}^i = \mathbf{p}^{i|j+1}$  and  $T^i = T^{i|j+1}$ 
    Compute  $\mathbf{x}^i$  by forward time integration of system dynamics
    Evaluate all constraints
    Update multipliers  $\bar{\boldsymbol{\mu}}^{i+1}$  and penalties  $\bar{\mathbf{c}}^{i+1}$ 
    If minimization is converged & constraint thresholds are satisfied then
        Break outer loop
    End If
End For
Compute cost  $J$  and norm of constraints

```

The basic structure of the algorithm that is implemented in the main calling function `grampc_run` is outlined in *Algorithm 1*. The projected gradient method is realized in the inner loop and consists of the forward and backward integration of the canonical equations as well as the update of the optimization variables based on the gradient and the approximate solution of the line search problem. The outer loop corresponds to the augmented Lagrangian method consisting of the solution of the inner minimization problem and the update of the multipliers and penalty parameters.

As an alternative to the augmented Lagrangian framework, the user can choose external penalty functions in GRAMPC that handle the equality and inequality constraints as “soft” constraints. In this case, the multipliers $\bar{\boldsymbol{\mu}}$ are fixed at zero and only the penalty parameters are updated in the outer loop. Note that the user can set the options `PenaltyIncreaseFactor` and `PenaltyDecreaseFactor` to 1.0 in order to keep the penalty parameters at the initial value `PenaltyMin`. The single steps of the algorithm and the related options are described in more detail in the

following sections.

The following options can be used to adjust the basic algorithm. The corresponding default values are listed in [Table 8.2.1](#) in the appendix.

- **MaxMultIter**: Sets the maximum number of augmented Lagrangian iterations $i_{\max} \geq 1$. If the option **ConvergenceCheck** is activated, the algorithm evaluates the convergence criterion and terminates if the inner minimization converged and all constraints are satisfied within the tolerance defined by **ConstraintsAbsTol**.
- **MaxGradIter**: Sets the maximum number of gradient iterations $j_{\max} \geq 1$. If the option **ConvergenceCheck** is activated, the algorithm terminates the inner loop as soon as the convergence criterion is fulfilled.
- **EqualityConstraints**: Equality constraints $\mathbf{g}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) = \mathbf{0}$ can be disabled by the option value **off**.
- **InequalityConstraints**: To disable inequality constraints $\mathbf{h}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) \leq \mathbf{0}$, set this option to **off**.
- **TerminalEqualityConstraints**: To disable terminal equality constraints $\mathbf{g}_T(\mathbf{x}(T), \mathbf{p}, T) = \mathbf{0}$, set this option to **off**.
- **TerminalInequalityConstraints**: To disable terminal inequality constraints $\mathbf{h}_T(\mathbf{x}(T), \mathbf{p}, T) \leq \mathbf{0}$, set this option to **off**.
- **ConstraintsHandling**: State constraints are handled either by means of the augmented Lagrangian approach (option value **auglag**) or as soft constraints by outer penalty functions (option value **extpen**).
- **OptimControl**: Specifies whether the cost functional should be minimized with respect to the control variable \mathbf{u} .
- **OptimParam**: Specifies whether the cost functional should be minimized with respect to the optimization parameters \mathbf{p} .
- **OptimTime**: Specifies whether the cost functional should be minimized with respect the horizon length T (free end time problem) or if T is kept constant.

5.2 Numerical Integration

GRAMPC employs a continuous-time formulation of the optimization problem. However, internally, all time-dependent functions are stored in discretized form with N_{hor} elements and numerical integration is performed to compute the cost functional and to solve the differential equations.

5.2.1 Integration of cost functional and explicit ODEs

The approximate line search method requires the evaluation of the cost functional. To this end, the integral cost is integrated numerically with the trapezoidal rule, Simpson rule or discrete summation and the terminal cost is added. Note that the discrete summation does not compute the Riemann sum but the plain sum with $\sum_k l(\mathbf{x}_k, \mathbf{u}_k, \mathbf{p}, t_k)$. The cost values are additionally evaluated at the end of the optimization as an add-on information for the user.

The gradient algorithm involves the sequential forward integration of the system dynamics and backward integration of the adjoint dynamics. To this end, GRAMPC provides the following four explicit Runge-Kutta (ERK) methods with fixed step size and the Butcher tableaux

$$\text{erk1 (Euler's first-order method)} : \begin{array}{c|c} 0 & \\ \hline & 1 \end{array}$$

$$\text{erk2 (Heun's second-order method)} : \begin{array}{c|cc} 0 & & \\ 1 & 1 & \\ \hline & 1/2 & 1/2 \end{array}$$

$$\begin{array}{l}
 \text{erk3 (Kutta's third-order method) :} \\
 \begin{array}{c|ccc}
 0 & & & \\
 1/2 & 1/2 & & \\
 1 & -1 & 2 & \\
 \hline
 & 1/6 & 2/3 & 1/6
 \end{array} \\
 \\
 \text{erk4 (Kutta's fourth-order method) :} \\
 \begin{array}{c|cccc}
 0 & & & & \\
 1/2 & 1/2 & & & \\
 1/2 & 0 & 1/2 & & \\
 1 & 0 & 0 & 1 & \\
 \hline
 & 1/6 & 1/3 & 1/3 & 1/6
 \end{array}
 \end{array}$$

In addition, a 4th-order explicit Runge-Kutta method with variable step size is available (option value `ruku45`). For discrete-time system dynamics, the integrator is set to `discrete`. For semi-implicit ODEs and DAEs, the Rosenbrock solver RODAS¹² with variable step size has to be used (option value `rodas`).

Note

For discrete-time systems the option `Integrator` is set to `discrete`, the settings `Nhor`, `Thor` and `dt` need to satisfy the relation `Thor = (Nhor-1)*dt` and the option `OptimTime` must be set to `off`. See also *Problems with discrete-time systems*.

The following options can be used to adjust the numerical integrations:

- `Nhor`: Number of discretization points within the time interval $[0, T]$.
- `IntegralCost`, `TerminalCost`: Indicate if the integral and/or terminal cost functions are defined.
- `IntegratorCost`: This option specifies the integration scheme for the cost functionals. Possible values are `trapezoidal`, `simpson` and `discrete`.

Changed in version v2.3: Fixed typo in `trapezoidal` option. Renamed `euler` to `erk1` and `heun` to `erk2`. Removed `modeuler`. Added `erk3` and `erk4`.

- `Integrator`: This option specifies the integration scheme for the system and adjoint dynamics. Possible values are `erk1`, `erk2`, `erk3`, `erk4`, `discrete` with fixed step size and `ruku45`, `rodas` with variable step size.
- `IntegratorMinStepSize`: Minimum step size for RODAS and the Runge-Kutta integrator.
- `IntegratorMaxSteps`: Maximum number of steps for RODAS and the Runge-Kutta integrator.
- `IntegratorRelTol`: Relative tolerance for RODAS and the Runge-Kutta integrator with variable step size. Note that this option may be insignificant if the minimum step size is chosen too high or the maximum number of steps is set too low.
- `IntegratorAbsTol`: Absolute tolerance for RODAS and the Runge-Kutta integrator with variable step size. Note that this option may be insignificant if the minimum step size is chosen too high or the maximum number of steps is set too low.

¹ E. Hairer and G. Wanner. *Solving Ordinary Differential Equations: Stiff and Differential-Algebraic Problems*. Springer, Heidelberg, Germany, 1996.

² RODAS. Webpage. <http://www.unige.ch/~hairer/software.html>, Accessed 01-December-2018.

5.2.2 Integration of semi-implicit ODEs and DAEs (RODAS)

GRAMPC supports problem descriptions with ordinary differential equations in semi-implicit form as well as differential algebraic equations using the solver RODAS for numerical integration (see [Problems involving semi-implicit ODEs and DAEs](#)). If a semi-implicit problem or differential algebraic equations are considered, the mass matrix M and its transposed version M^T must be defined by the C functions `Mfct` and `Mtrans`. The numerical integration can be accelerated by additionally providing the C functions `dfdx`, `dfdxtrans`, `dfdt` and `dHdxdt`. See [Problem implementation](#) for a detailed description of the problem implementation.

The integration with RODAS is configured by a number of flags that are passed to the solver using the vector `FlagsRodas` with the elements `[IFCN, IDFX, IJAC, IMAS, MLJAC, MUJAC, MLMAS, MUMAS]`. See [Page 25, 2](#) [Page 25, 1](#) for a detailed description of these flags. The default values `[0, 0, 0, 0, Nx, Nx, Nx, Nx]` correspond to an autonomous system with an identity matrix as mass matrix. The following options can be adjusted via `FlagsRodas`:

- **IFCN**: Specifies if the right hand side of the system dynamics $f(x, u, p, t)$ explicitly depends on time t (`IFCN = 1`) or if the problem is autonomous (`IFCN = 0`).
- **IDFX**: Specifies how the computation of the partial derivatives $\frac{\partial f}{\partial t}$ and $\frac{\partial^2 H}{\partial x \partial t}$ is carried out. The partial derivatives are computed internally by finite differences (`IDFX = 0`) or are provided by the functions `dfdt` and `dHdxdt` (`IDFX = 1`) as described in [Problem implementation](#).
- **IJAC**: Specifies how the computation of the Jacobians $\frac{\partial f}{\partial x}$ and $(\frac{\partial f}{\partial x})^T = \frac{\partial^2 H}{\partial x \partial \lambda}$ is carried out for numerically solving the canonical equations. The Jacobians are computed internally by finite differences (`IJAC = 0`) or are provided by the functions `dfdx` and `dfdxtrans` (`IJAC = 1`), also see [Problem implementation](#).
- **IMAS**: Gives information on the mass matrix M , which is either an identity matrix (`IMAS = 0`) or is specified by the function `Mfct` (`IMAS = 1`). Note that the adjoint dynamics requires the transposed mass matrix that has to be provided by the function `Mtrans`.
- **MLJAC**: Gives information on the banded structure of the Jacobian $\frac{\partial f}{\partial x}$ and $(\frac{\partial f}{\partial x})^T$, respectively. The Jacobian is either a full matrix (`MLJAC = Nx`) or is of banded structure. In the latter case, the number of non-zero diagonals below the main diagonal are specified by $0 \leq \text{MLJAC} < N_x$.
- **MUJAC**: Specifies the number of non-zero diagonals above the main diagonal of the Jacobian $\frac{\partial f}{\partial x}$. This flag needs not to be defined if `MLJAC = Nx`. Since the partial derivative of the right hand side of the adjoint dynamics with respect to the adjoint state λ is given by $(\frac{\partial f}{\partial x})^T$, the meaning of the flags `MLJAC` and `MUJAC` switches in this case.
- **MLMAS and MUMAS**: Both options have the same meaning as `MLJAC` and `MUJAC`, but refer to the mass matrix M .

If a semi-implicit problem (option `IMAS = 1`) with Mayer term (option `TerminalCost = on`) is considered, the terminal conditions of the adjoint system must be specified in a specific form. To provide RODAS [Page 25, 1](#) [Page 25, 2](#) the proper terminal condition $\lambda(T)$, the function `dVdx` must be specified as follows

$$\lambda(T) = \underbrace{\left(M^T\right)^{-1} \bar{V}_x(x(T), p, T, \mu_T, c_T)}_{\text{dVdx}}$$

cf. Equation (5.1.3). In the case of a DAE system, the mass matrix is singular and therefore only the elements of the mass matrix for the differential equations are inverted. For an example of using RODAS and the respective options, take a look at the MPC problem `Reactor_PDE` in the folder `<grampc_root>/examples`.

5.3 Line search

The projected gradient method as part of the GRAMPC algorithm *Algorithm 1* requires the solution of the line search problem²

$$\min_{\alpha} \bar{J} \left(\psi_{\mathbf{u}} \left(\mathbf{u}^{i|j} - \alpha \mathbf{d}_{\mathbf{u}}^{i|j} \right), \psi_{\mathbf{p}} \left(\mathbf{p}^{i|j} - \gamma_{\mathbf{p}} \alpha \mathbf{d}_{\mathbf{p}}^{i|j} \right), \psi_T \left(T^{i|j} - \gamma_T \alpha \mathbf{d}_T^{i|j} \right); \bar{\boldsymbol{\mu}}, \bar{\mathbf{c}}, \mathbf{x}_0 \right). \quad (5.3.1)$$

GRAMPC implements two efficient strategies to solve this problem in an approximate manner. The following options apply to both line search methods that are detailed below (*Adaptive line search* and *Explicit line search*):

- **LineSearchType:** This option selects either the adaptive line search strategy (value `adaptive`) or the explicit approach (value `explicit1` or `explicit2`).
- **LineSearchExpAutoFallback:** If this option is activated, the automatic fallback strategy is used in the case that the explicit formulas result in negative step sizes.
- **LineSearchMax:** This option sets the maximum value α_{\max} of the step size α .
- **LineSearchMin:** This option sets the minimum value α_{\min} of the step size α .
- **LineSearchInit:** Indicates the initial value $\alpha_{\text{init}} > 0$ for the step size α . If the adaptive line search is used, the sample point α_2 is set to $\alpha_2 = \alpha_{\text{init}}$.
- **OptimParamLineSearchFactor:** This option sets the adaptation factor $\gamma_{\mathbf{p}}$ that weights the update of the parameter vector \mathbf{p} against the update of the control \mathbf{u} .
- **OptimTimeLineSearchFactor:** This option sets the adaptation factor γ_T that weights the update of the end time T against the update of the control \mathbf{u} .

5.3.1 Adaptive line search

An appropriate way to determine the step size is the adaptive line search approach from³, where a polynomial approximation of the cost \bar{J} is used and an adaptation of the search intervals is performed. More precisely, the cost functional is evaluated at three sample points $\alpha_1 < \alpha_2 < \alpha_3$ with $\alpha_2 = \frac{1}{2}(\alpha_1 + \alpha_3)$, which are used to construct a quadratic polynomial of the cost according to

$$\begin{aligned} \bar{J} \left(\psi_{\mathbf{u}} \left(\mathbf{u}^{i|j} - \alpha \mathbf{d}_{\mathbf{u}}^{i|j} \right), \psi_{\mathbf{p}} \left(\mathbf{p}^{i|j} - \gamma_{\mathbf{p}} \alpha \mathbf{d}_{\mathbf{p}}^{i|j} \right), \psi_T \left(T^{i|j} - \gamma_T \alpha \mathbf{d}_T^{i|j} \right); \bar{\boldsymbol{\mu}}, \bar{\mathbf{c}}, \mathbf{x}_0 \right) \\ \approx \Phi(\alpha) = p_0 + p_1 \alpha + p_2 \alpha^2. \end{aligned} \quad (5.3.2)$$

Subsequently, a step size α^j is computed by minimizing the cost approximation (5.3.2). If necessary, the interval $[\alpha_1, \alpha_3]$ is adapted for the next gradient iteration in the following way

$$\begin{aligned} [\alpha_1, \alpha_3] &\leftarrow \begin{cases} \kappa [\alpha_1, \alpha_3] & \text{if } \alpha \geq \alpha_3 - \varepsilon_{\alpha}(\alpha_3 - \alpha_1) \text{ and } \alpha_3 \leq \alpha_{\max} \text{ and } |\Phi(\alpha_1) - \Phi(\alpha_3)| > \varepsilon_{\phi} \\ \frac{1}{\kappa} [\alpha_1, \alpha_3] & \text{if } \alpha \leq \alpha_1 + \varepsilon_{\alpha}(\alpha_3 - \alpha_1) \text{ and } \alpha_1 \geq \alpha_{\min} \text{ and } |\Phi(\alpha_1) - \Phi(\alpha_3)| > \varepsilon_{\phi} \\ [\alpha_1, \alpha_3] & \text{otherwise} \end{cases} \\ \alpha_2 &\leftarrow \frac{1}{2}(\alpha_1 + \alpha_3) \end{aligned} \quad (5.3.3)$$

² T. Englert, A. Völz, F. Mesmer, S. Rhein, and K. Graichen. A software framework for embedded nonlinear model predictive control using a gradient-based augmented Lagrangian approach (GRAMPC). *Optimization and Engineering*, 20(3):769–809, 2019. doi:10.1007/s11081-018-9417-2. doi:10.1007/s11081-018-9417-2.

³ K. Graichen and B. Käpelnick. A real-time gradient method for nonlinear model predictive control. In T. Zheng, editor, *Frontiers of Model Predictive Control*, pages 9–28. InTech, 2012.

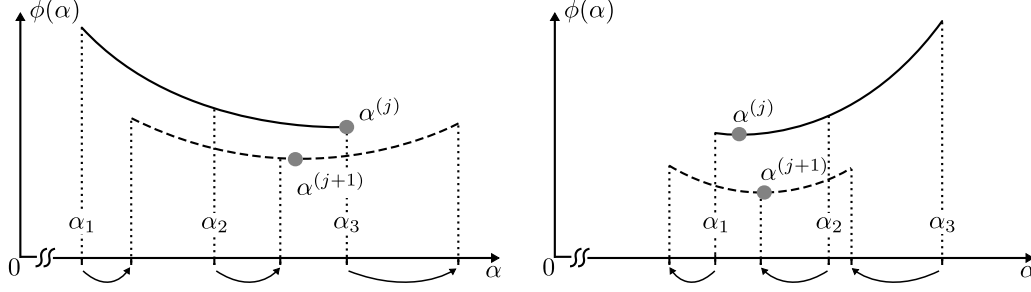


Fig. 5.3.1: Adaptation of line search interval.

with the adaptation factor $\kappa > 1$, the interval tolerance $\varepsilon_\alpha \in (0, 0.5)$, the absolute cost tolerance $\varepsilon_\phi \in [0, \infty)$ for adapting the interval and the interval bounds $\alpha_{\max} > \alpha_{\min} > 0$. The modification (5.3.3) of the line search interval tracks the minimum point α^j of the line search problem in the case when α^j is either outside of the interval $[\alpha_1, \alpha_3]$ or close to one of the outer bounds α_1, α_3 , as illustrated in Fig. 5.3.1. The adaptation factor κ accounts for scaling as well as shifting of the interval $[\alpha_1, \alpha_3]$ in the next gradient iteration, if α^j lies in the vicinity of the interval bounds $[\alpha_1, \alpha_3]$ as specified by the interval tolerance ε_α . This adaptive strategy allows one to track the minimum of the line search problem (5.3.1) over the gradient iterations j and MPC steps k , while guaranteeing a fixed number of operations in view of a real-time MPC implementation. The absolute tolerance ε_ϕ of the difference in the (scaled) costs at the interval bounds $|\Phi(\alpha_1) - \Phi(\alpha_3)|$ avoids oscillations of the interval width in regions where the cost function \bar{J} is almost constant.

The following options apply specifically to the adaptive line search strategy:

- **LineSearchAdaptAbsTol:** This option sets the absolute tolerance ε_ϕ of the difference in costs at the interval bounds α_1 and α_2 . If the difference in the (scaled) costs on these bounds falls below ε_ϕ , the adaption of the interval is stopped in order to avoid oscillations.
- **LineSearchAdaptFactor:** This option sets the adaptation factor $\kappa > 1$ in (5.3.3) that determines how much the line search interval can be adapted from one gradient iteration to the next.
- **LineSearchIntervalTol:** This option sets the interval tolerance $\varepsilon_\alpha \in (0, 0.5)$ in (5.3.3) that determines for which values of α the adaption is performed.
- **LineSearchIntervalFactor:** This option sets the interval factor $\beta \in (0, 1)$ that specifies the interval bounds $[\alpha_1, \alpha_3]$ according to $\alpha_1 = \alpha_2(1 - \beta)$ and $\alpha_3 = \alpha_2(1 + \beta)$, whereby the mid sample point is initialized as $\alpha_2 = \alpha_{\text{init}}$.

5.3.2 Explicit line search

An alternative way to determine the step size in order to further reduce the computational effort for time-critical problems is the explicit line search approach originally discussed in⁴ and adapted in⁵ for the optimal control case. The motivation is to minimize the difference between two consecutive control updates $u_k^{i|j}(\tau)$ and $u_k^{i|j+1}(\tau)$ in the uncon-

⁴ J. Barzilai and J. M. Borwein. Two-point step size gradient methods. *SIAM Journal on Numerical Analysis*, 8(1):141–148, 1988.

⁵ B. Käpernick and K. Graichen. Model predictive control of an overhead crane using constraint substitution. In *Proceedings of the American Control Conference (ACC)*, 3973–3978. 2013.

strained case and additionally assuming the same step size $\alpha^{i|j}$, i.e.

$$\begin{aligned}
 \alpha^{i|j} &= \arg \min_{\alpha > 0} \left\| \mathbf{u}^{i|j+1} - \mathbf{u}^{i|j} \right\|_{L_m^2[0,T]}^2 + \left\| \mathbf{p}^{i|j+1} - \mathbf{p}^{i|j} \right\|_2^2 + \left| T^{i|j+1} - T^{i|j} \right| \\
 &= \arg \min_{\alpha > 0} \left\| \underbrace{\mathbf{u}^{i|j} - \mathbf{u}^{j-1}}_{=: \Delta \mathbf{u}^{i|j}} - \alpha \underbrace{\left(\mathbf{d}_u^{i|j} - \mathbf{d}_u^{j-1} \right)}_{=: \Delta \mathbf{d}_u^{i|j}} \right\|_{L_m^2[0,T]}^2 + \left\| \underbrace{\mathbf{p}^{i|j} - \mathbf{p}^{j-1}}_{=: \Delta \mathbf{p}^{i|j}} - \gamma_p \alpha \underbrace{\left(\mathbf{d}_p^{i|j} - \mathbf{d}_p^{j-1} \right)}_{=: \Delta \mathbf{d}_p^{i|j}} \right\|_2^2 \\
 &\quad + \left\| \underbrace{T^{i|j} - T^{j-1}}_{=: \Delta T^{i|j}} - \gamma_T \alpha \underbrace{\left(d_T^{i|j} - d_T^{j-1} \right)}_{=: \Delta d_T^{i|j}} \right\|_2^2
 \end{aligned} \tag{5.3.4}$$

with $\|z\|_{L_m^2[0,T]}^2 = \langle z, z \rangle := \int_0^T z^\top(t) z(t) dt$.

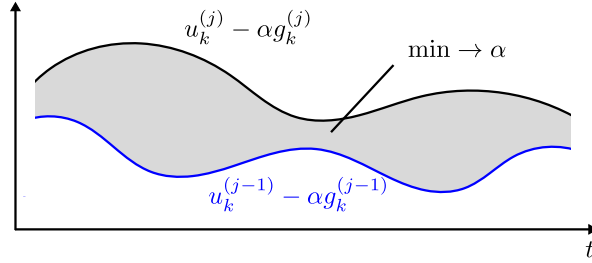


Fig. 5.3.2: Motivation for the explicit line search strategy.

Fig. 5.3.2 illustrates the general idea behind (5.3.4). To solve (5.3.4), consider the following function

$$\begin{aligned}
 q(\alpha) &:= \left\| \Delta u_k^j - \alpha \Delta d_k^j \right\|_{L_m^2[0,T]}^2 \\
 &= \int_0^T \left(\Delta u_k^j - \alpha \Delta d_k^j \right)^\top \left(\Delta u_k^j - \alpha \Delta d_k^j \right) dt \\
 &= \int_0^T \left(\Delta u_k^j \right)^\top \Delta u_k^j dt + \alpha^2 \int_0^T \left(\Delta d_k^j \right)^\top \Delta d_k^j dt - 2\alpha \int_0^T \left(\Delta u_k^j \right)^\top \Delta d_k^j dt.
 \end{aligned} \tag{5.3.5}$$

The minimum has to satisfy the stationarity condition

$$\frac{\partial q(\alpha)}{\partial \alpha} = 2\alpha \int_0^T \left(\Delta d_k^j \right)^\top \Delta d_k^j dt - 2 \int_0^T \left(\Delta u_k^j \right)^\top \Delta d_k^j dt = 0.$$

A suitable step size α^j then follows to

$$\alpha^j = \frac{\int_0^T \left(\Delta u_k^j \right)^\top \Delta d_k^j dt}{\int_0^T \left(\Delta d_k^j \right)^\top \Delta d_k^j dt} = \frac{\langle \Delta u_k^j, \Delta d_k^j \rangle}{\langle \Delta d_k^j, \Delta d_k^j \rangle}.$$

Another way to compute an appropriate step size for the control update can be achieved by reformulating (5.3.5) in the following way:

$$q(\alpha) = \left\| \Delta u_k^j - \alpha \Delta d_k^j \right\|_{L_m^2[0,T]}^2 = \alpha^2 \left\| \frac{1}{\alpha} \Delta u_k^j - \Delta d_k^j \right\|_{L_m^2[0,T]}^2 =: \alpha^2 \bar{q}(\alpha).$$

In the subsequent, the new function $\bar{q}(\alpha)$ is minimized w.r.t. the step size leading to a similar solution

$$\alpha^j = \frac{\langle \Delta u^j, \Delta u^j \rangle}{\langle \Delta u^j, \Delta d_k^j \rangle}.$$

For the original problem (5.3.4), the solution

$$\alpha^{i|j} = \frac{\langle \Delta \mathbf{u}^{i|j}, \Delta \mathbf{d}_{\mathbf{u}}^{i|j} \rangle + \gamma_{\mathbf{p}} \langle \Delta \mathbf{p}^{i|j}, \Delta \mathbf{d}_{\mathbf{p}}^{i|j} \rangle + \gamma_T \Delta T^{i|j} \Delta d_T^{i|j}}{\langle \Delta \mathbf{d}_{\mathbf{u}}^{i|j}, \Delta \mathbf{d}_{\mathbf{u}}^{i|j} \rangle + \gamma_{\mathbf{p}}^2 \langle \Delta \mathbf{d}_{\mathbf{p}}^{i|j}, \Delta \mathbf{d}_{\mathbf{p}}^{i|j} \rangle + \gamma_T^2 (\Delta d_T^{i|j})^2} \quad (5.3.6)$$

$$\alpha^{i|j} = \frac{\langle \Delta \mathbf{u}^{i|j}, \Delta \mathbf{u}^{i|j} \rangle + \gamma_{\mathbf{p}} \langle \Delta \mathbf{p}^{i|j}, \Delta \mathbf{p}^{i|j} \rangle + \gamma_T (\Delta T^{i|j})^2}{\langle \Delta \mathbf{u}^{i|j}, \Delta \mathbf{d}_{\mathbf{u}}^{i|j} \rangle + \gamma_{\mathbf{p}}^2 \langle \Delta \mathbf{p}^{i|j}, \Delta \mathbf{d}_{\mathbf{p}}^{i|j} \rangle + \gamma_T^2 \Delta T^{i|j} \Delta d_T^{i|j}} \quad (5.3.7)$$

follows.

In the GRAMPC implementation, both approaches (5.3.6) and (5.3.7) are available. In addition, the step size α^j is bounded by the upper and lower values $\alpha_{\max} > \alpha_{\min} > 0$. However, if the originally computed step size α^j is less than zero¹, either the initial step size $\alpha^j = \alpha_{\text{init}}$ or the automatic fallback strategy that is detailed in the next subsection is used in order to achieve a valid step size. The fallback strategy is set with the following option (only available for the explicit line search strategies):

- **LineSearchExpAutoFallback**: If this option is activated, the automatic fallback strategy is used in the case that the explicit formulas result in negative step sizes.

5.3.3 Fallback strategy for explicit line search

While the initial step size can be used as fallback solution if the explicit step size computation yields negative values, it often requires problem-specific tuning of α_{init} for achieving optimal performance. As alternative, GRAMPC implements an automatic fallback strategy that is based on the idea of using at most 1% of the control range defined by u_{\max} and u_{\min} . For this purpose, the maximum absolute value $\mathbf{d}_{\mathbf{u},\max}^{i|j} = \|\mathbf{d}_{\mathbf{u}}^{i|j}(t)\|_{L^\infty}$ of the search direction $\mathbf{d}_{\mathbf{u}}^{i|j}(t)$ over the horizon is determined. Subsequently, the step size

$$\alpha^{i|j} = \frac{1}{100} \cdot \min_{k \in \{1, \dots, N_{\mathbf{u}}\}} \left\{ \frac{u_{\max,k} - u_{\min,k}}{\mathbf{d}_{\mathbf{u},\max,k}^{i|j}} \right\}$$

follows as the minimal step size required to perform a step of 1% with respect to the range of at least one control in at least one time step. Additionally, the step size is limited to 10% of the maximum step size α_{\max} . Since this strategy requires reasonable limits for the controls, it is only executed if **LineSearchExpAutoFallback** is activated and if these limits are defined by the user. Furthermore, this strategy can only be used if **OptimControl** is switched on. In all other case, the initial step size $\alpha^j = \alpha_{\text{init}}$ will be used as fallback solution.

5.4 Update of multipliers and penalties

GRAMPC handles general nonlinear constraints using an augmented Lagrangian approach or, alternatively, using an external penalty method. The key to the efficient solution of constrained problems using these approaches are the updates of the multipliers in the outer loop for $i = 1, \dots, i_{\max}$.

¹ It can be shown that the step size α is negative if the cost function is locally non-convex.

5.4.1 Update of Lagrangian multipliers

The outer loop of the GRAMPC algorithm in [Algorithm 1](#) maximizes the augmented Lagrangian function with respect to the multipliers $\bar{\mu}$. This update is carried out in the direction of steepest ascent that is given by the constraint residual. The penalty parameter is used as step size, as it is typically done in augmented Lagrangian methods.

For an arbitrary equality constraint $g^i = g(x^i, \dots)$ with multiplier μ_g^i , penalty c_g^i , and tolerance ε_g , the update is defined by

$$\mu_g^{i+1} = \zeta_g(\mu_g^i, c_g^i, g^i, \varepsilon_g) = \begin{cases} \mu_g^i + (1 - \rho)c_g^i g^i & \text{if } |g^i| > \varepsilon_g \wedge \eta^i \leq \varepsilon_{\text{rel,u}} \\ \mu_g^i & \text{else.} \end{cases}$$

The update is not performed if the constraint is satisfied within its tolerance ε or if the inner minimization is not sufficiently converged, which is checked by the maximum relative gradient η^i (see (5.5.1) for the definition) and the threshold $\varepsilon_{\text{rel,u}}$. Similarly, for an inequality constraint $\bar{h}^i = \bar{h}(x^i, \dots)$ with multiplier μ_h^i , penalty c_h^i , and tolerance ε_h , the update is defined by

$$\mu_h^{i+1} = \zeta_h(\mu_h^i, c_h^i, \bar{h}^i, \varepsilon_h) = \begin{cases} \mu_h^i + (1 - \rho)c_h^i \bar{h}^i & \text{if } (\bar{h}^i > \varepsilon_h \wedge \eta^i \leq \varepsilon_{\text{rel,u}}) \vee \bar{h}^i < 0 \\ \mu_h^i & \text{else.} \end{cases}$$

Similar update rules are used for the terminal equality and terminal inequality constraints.

GRAMPC provides several means to increase the robustness of the multiplier update, which may be required if few iterations j_{max} are used for the suboptimal solution of the inner minimization problem. The damping factor $\rho \in [0, 1)$ can be used to scale the step size of the steepest ascent and the tolerance $\varepsilon_{\text{rel,u}}$ can be used to skip the multiplier update in case that the minimization is not sufficiently converged. Furthermore, the multipliers are limited by lower and upper bounds $\mu_g \in [-\mu_{\text{max}}, \mu_{\text{max}}]$ for equalities and $\mu_h \leq \mu_{\text{max}}$ for inequalities, respectively, to avoid unlimited growth. A status flag is set if one of the multipliers reaches this bound and the user should check the problem formulation as this case indicates an ill-posed or even infeasible optimization problem.

The following options can be used to adjust the update of the Lagrangian multipliers:

- **MultiplierMax**: Upper bound μ_{max} and lower bound $-\mu_{\text{max}}$ for the Lagrangian multipliers.
- **MultiplierDampingFactor**: Damping factor $\rho \in [0, 1)$ for the multiplier update.
- **AugLagUpdateGradientRelTol**: Threshold $\varepsilon_{\text{rel,u}}$ for the maximum relative gradient of the inner minimization problem.
- **ConstraintsAbsTol**: Thresholds $(\varepsilon_g, \varepsilon_h, \varepsilon_{g_T}, \varepsilon_{h_T}) \in \mathbb{R}^{N_c}$ for the equality, inequality, terminal equality, and terminal inequality constraints.

5.4.2 Update of penalty parameters

The penalty parameters \bar{c} are adapted in each outer iteration according to a heuristic rule that is motivated by the LANCELOT package¹³. A carefully tuned adaptation of the penalties can speed-up the convergence significantly and is therefore highly recommended (also see [Estimation of minimal penalty parameter](#) and the tutorial in [Model predictive control of a PMSM](#)). Note that the penalty parameters are also updated if external penalties are used instead of the augmented Lagrangian method, i.e. `ConstraintsHandling` is set to `extpen`. In order to keep the penalty parameters at the initial value `PenaltyMin`, the options `PenaltyIncreaseFactor` and `PenaltyDecreaseFactor` can be set to 1.0, which basically deactivates the penalty update.

¹ A. R. Conn, G.I.M. Gould, and P. L. Toint. *LANCELOT: A Fortran Package for Large-Scale Nonlinear Optimization (Release A)*. Springer-Verlag, Berlin, Germany, 1992.

³ J. Nocedal and S. Wright. *Numerical Optimization*. Springer Science & Business Media, New York, USA, 2006.

For an arbitrary equality constraint $g^i = g(x^i, \dots)$ with penalty c_g^i and tolerance ε_g , the update is defined by

$$c_g^{i+1} = \xi_g(c_g^i, g^i, g^{i-1}, \varepsilon_g) = \begin{cases} \beta_{\text{in}} c_g^i & \text{if } |g^i| \geq \gamma_{\text{in}} |g^{i-1}| \wedge |g^i| > \varepsilon_g \wedge \eta^i \leq \varepsilon_{\text{rel,u}} \\ \beta_{\text{de}} c_g^i & \text{else if } |g^i| \leq \gamma_{\text{de}} \varepsilon_g \\ c_g^i & \text{else.} \end{cases} \quad (5.4.1)$$

The penalty c_g^i is increased by the factor $\beta_{\text{in}} > 1$ if the (sub-optimal) solution of the inner minimization problem does not generate sufficient progress in the constraint, which is rated by the factor $\gamma_{\text{in}} > 0$ and compared to the previous iteration $i - 1$. This update is skipped if the inner minimization is not sufficiently converged, which is checked by the maximum relative gradient η^i and the threshold $\varepsilon_{\text{rel,u}}$. The penalty c_g^i is decreased by the factor $\beta_{\text{de}} < 1$ if the constraint g^i is sufficiently satisfied within its tolerance, whereby currently the constant factor $\gamma_{\text{de}} = 0.1$ is used. The setting $\beta_{\text{in}} = \beta_{\text{de}} = 1$ can be used to keep the penalty constant, i.e., to deactivate the penalty adaptation. Similarly, for an inequality constraint $\bar{h}^i = \bar{h}(x^i, \dots)$ with penalty c_h^i and tolerance ε_h , the update is defined by

$$c_h^{i+1} = \xi_h(c_h^i, \bar{h}^i, \bar{h}^{i-1}, \varepsilon_h) = \begin{cases} \beta_{\text{in}} c_h^i & \text{if } \bar{h}^i \geq \gamma_{\text{in}} \bar{h}^{i-1} \wedge \bar{h}^i > \varepsilon_h \wedge \eta^i \leq \varepsilon_{\text{rel,u}} \\ \beta_{\text{de}} c_h^i & \text{else if } \bar{h}^i \leq \gamma_{\text{de}} \varepsilon_h \\ c_h^i & \text{else.} \end{cases} \quad (5.4.2)$$

Similar update rules are used for the terminal equality and inequality constraints. In analogy to the multiplier update, the penalty parameters are restricted to upper and lower bounds $c_{\text{max}} \gg c_{\text{min}} > 0$ in order to avoid unlimited growth as well as negligible values.

The following options can be used to adjust the update of the penalty parameters:

- **PenaltyMax**: This option sets the upper bound c_{max} of the penalty parameters.
- **PenaltyMin**: This option sets the lower bound c_{min} of the penalty parameters.
- **PenaltyIncreaseFactor**: This option sets the factor β_{in} by which penalties are increased.
- **PenaltyDecreaseFactor**: This option sets the factor β_{de} by which penalties are decreased.
- **PenaltyIncreaseThreshold**: This option sets the factor γ_{in} that rates the progress in the constraints between the last two iterates.
- **AugLagUpdateGradientRelTol**: Threshold $\varepsilon_{\text{rel,u}}$ for the maximum relative gradient of the inner minimization problem.
- **ConstraintsAbsTol**: Thresholds $(\varepsilon_g, \varepsilon_h, \varepsilon_{g_T}, \varepsilon_{h_T}) \in \mathbb{R}^{N_c}$ for the equality, inequality, terminal equality, and terminal inequality constraints.

5.4.3 Estimation of minimal penalty parameter

In real-time or embedded MPC applications, where only a limited number of iterations per step is computed, it is crucial that the penalty parameter is not decreased below a certain threshold c_{min} . This lower bound should be large enough that an inactive constraint that becomes active is still sufficiently penalized in the augmented Lagrangian cost functional. However, it should not be chosen too high to prevent ill-conditioning. A suitable value of c_{min} tailored to the given MPC problem therefore is of importance to ensure a high performance of GRAMPC.

In order to support the user, GRAMPC offers the routine `grampc_estim_penmin` to compute a problem-specific estimate of c_{min} . The basic idea behind this estimation is to determine c_{min} such that the actual costs J are in the same order of magnitude as the squared constraints multiplied by c_{min} , see equation (5.1.1). This approach requires initial values for the states x , controls u , and cost J . If the GRAMPC structure includes only default values, i.e. zeros (cold start), the estimation function `grampc_estim_penmin` can be called with the argument `rungrampc=1` to perform one optimization or MPC step, where the possible maximum numbers of gradient iterations `MaxGradIter` and augmented Lagrangian iterations `MaxMultIter` are limited to 20. Afterwards, the estimated value for `PenaltyMin` is set as detailed below and, if `rungrampc=1`, the initial states x , controls u and costs J are reset.

Based on the initial values, a first estimate of the minimal penalty parameter is computed according to

$$\hat{c}_{\min}^I = \frac{2|J|}{\|g(x(t), u(t), p, t)\|_{L_2}^2 + \|h(x(t), u(t), p, t)\|_{L_2}^2 + \|g_T(x(T), p, T)\|_2^2 + \|h_T(x(T), p, T)\|_2^2} \quad (5.4.3)$$

However, if the inequality constraints are initially inactive and are far away from their bounds (i.e. large negative values are returned), the estimate \hat{c}_{\min}^I may be too small.

To deal with these cases, a second estimate for the minimal penalty parameter

$$\hat{c}_{\min}^{II} = \frac{2|J|}{T(\|\epsilon_g\|_2^2 + \|\epsilon_h\|_2^2) + \|\epsilon_{g_T}\|_2^2 + \|\epsilon_{h_T}\|_2^2} \quad (5.4.4)$$

is computed in the same spirit using the constraint tolerances (see `ConstraintsAbsTol`, ϵ_g , ϵ_h , ϵ_{g_T} and ϵ_{h_T}) instead of the constraint values². Since the norms of the tolerances are summed, more conservative values for c_{\min} are estimated and therefore instabilities can be avoided. Note that it is recommended to scale all constraints so that they are in the same order of magnitude, see e.g. the PMSM example in *Model predictive control of a PMSM*.

Finally, the minimal penalty parameter

$$\hat{c}_{\min} = \min \left\{ \max \{ \hat{c}_{\min}^I, \kappa \hat{c}_{\min}^{II} \}, \frac{c_{\max}}{500} \right\}$$

is chosen as the maximum of (5.4.3) and (5.4.4) and additionally limited to 0.2% of the maximum penalty parameter c_{\max} . This limitation ensures reasonable values even with very small constraint tolerances. The relation factor κ has been determined to 10^{-6} on the basis of various example systems. Please note that this estimation is intended to assist the user in making an initial guess. Problem-specific tuning of `PenaltyMin` can lead to further performance improvements and is therefore recommended. All MPC example problems in `<grampc_root>/examples` contain an initial call of `grampc_estim_penmin` to estimate \hat{c}_{\min} and, as alternative, manually tuned values that can further enhance the performance of GRAMPC for fixed numbers of iterations.

5.5 Convergence criterion

While the usage of fixed iteration counts i_{\max} and j_{\max} for the outer and inner loops is typical in real-time MPC or MHE applications, GRAMPC also provides an optional convergence check that is useful for solving optimal control or parameter optimization problems, or if the computation time in MPC is of minor importance.

The inner gradient loop in *Algorithm 1* evaluates the maximum relative gradient

$$\eta^{i|j+1} = \max \left\{ \frac{\|u^{i|j+1} - u^{i|j}\|_{L_2}}{\|u^{i|j+1}\|_{L_2}}, \frac{\|p^{i|j+1} - p^{i|j}\|_2}{\|p^{i|j+1}\|_2}, \frac{|T^{i|j+1} - T^{i|j}|}{T^{i|j+1}} \right\} \quad (5.5.1)$$

in each iteration $i|j$ and terminates if

$$\eta^{i|j} \leq \epsilon_{\text{rel,c}}. \quad (5.5.2)$$

Otherwise, the inner loop is continued until the maximum number of iterations j_{\max} is reached. The last value $\eta^i = \eta^{i|j+1}$ is returned to the outer loop and used for the convergence check as well as for the update of multipliers and penalties.

The augmented Lagrangian loop is terminated in iteration i if the inner loop is converged, that is $\eta^i \leq \epsilon_{\text{rel,c}}$, and all constraints are sufficiently satisfied, i.e.

$$\begin{bmatrix} |g^i(t)| \\ \max\{0, \bar{h}^i(t)\} \end{bmatrix} \leq \begin{bmatrix} \epsilon_g \\ \epsilon_h \end{bmatrix} \quad \forall t \in [0, T] \quad \wedge \quad \begin{bmatrix} |g_T^i| \\ \max\{0, \bar{h}_T^i\} \end{bmatrix} \leq \begin{bmatrix} \epsilon_{g_T} \\ \epsilon_{h_T} \end{bmatrix}, \quad (5.5.3)$$

² The integration behind the L^2 -norm can be replaced by a multiplication by the horizon length T , as the constraint tolerances ϵ_g and ϵ_h are no functions of time.

whereby the notation $|\cdot|$ denotes the component-wise absolute value. Otherwise, the outer loop is continued until the maximum number of iterations i_{\max} is reached. The thresholds $\varepsilon_g, \varepsilon_h, \varepsilon_{g_T}, \varepsilon_{h_T}$ are vector-valued in order to rate each constraint individually.

The following options can be used to adjust the convergence criterion:

- **ConvergenceCheck**: This option activates the convergence criterion. Otherwise, the inner and outer loops always perform the maximum number of iterations, see the options **MaxGradIter** and **MaxMultIter**.
- **ConvergenceGradientRelTol**: This option sets the threshold $\varepsilon_{\text{rel},c}$ for the maximum relative gradient of the inner minimization problem that is used in the convergence criterion. Note that this threshold is different from the one that is used in the update of multipliers and penalties.
- **ConstraintsAbsTol**: Thresholds $(\varepsilon_g, \varepsilon_h, \varepsilon_{g_T}, \varepsilon_{h_T}) \in \mathbb{R}^{N_c}$ for the equality, inequality, terminal equality, and terminal inequality constraints.

5.6 Scaling

Scaling is recommended for improving the numerical conditioning when the states \mathbf{x} and the optimization variables $(\mathbf{u}, \mathbf{p}, T)$ of the given optimization problem differ in several orders of magnitude. Although GRAMPC allows one to scale a specific problem automatically using the option **ScaleProblem=1**, it should be noted that this typically increases the computational load due to the additional multiplications in the algorithm, cf. the tutorial on controlling a permanent magnet synchronous machine in *Model predictive control of a PMSM*. This issue can be avoided by directly formulating the scaled problem within the C file template `probfc_t_TEMPLATE.c` included in the folder `examples/TEMPLATE`, also see *Problem implementation*.

The scaling in GRAMPC is performed according to

$$\begin{aligned}\bar{\mathbf{x}}(t) &= (\mathbf{x}(t) - \mathbf{x}_{\text{offset}}) ./ \mathbf{x}_{\text{scale}} \\ \bar{\mathbf{u}}(t) &= (\mathbf{u}(t) - \mathbf{u}_{\text{offset}}) ./ \mathbf{u}_{\text{scale}} \\ \bar{\mathbf{p}} &= (\mathbf{p} - \mathbf{p}_{\text{offset}}) ./ \mathbf{p}_{\text{scale}} \\ \bar{T} &= \frac{T - T_{\text{offset}}}{T_{\text{scale}}},\end{aligned}\tag{5.6.1}$$

where $\mathbf{x}_{\text{offset}} \in \mathbb{R}^x$, $\mathbf{u}_{\text{offset}} \in \mathbb{R}^u$, $\mathbf{p}_{\text{offset}} \in \mathbb{R}^p$ and $T_{\text{offset}} \in \mathbb{R}$ denote offset values and $\mathbf{x}_{\text{scale}} \in \mathbb{R}^x$, $\mathbf{u}_{\text{scale}} \in \mathbb{R}^u$, $\mathbf{p}_{\text{scale}} \in \mathbb{R}^p$ and $T_{\text{scale}} \in \mathbb{R}$ are scaling values. The symbol $./$ in (5.6.1) denotes element-wise division by the scaling vectors.

Furthermore, GRAMPC provides a scaling factor J_{scale} for the cost functional as well as scaling factors $\mathbf{c}_{\text{scale}} = [\mathbf{c}_{\text{scale},g}, \mathbf{c}_{\text{scale},h}, \mathbf{c}_{\text{scale},g_T}, \mathbf{c}_{\text{scale},h_T}] \in \mathbb{R}^{N_c}$ for the constraints. The scaling of the cost functional is relevant as the constraints are adjoined to the cost functional by means of Lagrangian multipliers and penalty parameters and the original cost functional should be of the same order of magnitude as these additional terms.

The following options can be used to adjust the scaling:

- **ScaleProblem**: Activates or deactivates scaling. Note that GRAMPC requires more computation time if scaling is active.
- **xScale, xOffset**: Scaling factors $\mathbf{x}_{\text{scale}}$ and offsets $\mathbf{x}_{\text{offset}}$ for each state variable.
- **uScale, uOffset**: Scaling factors $\mathbf{u}_{\text{scale}}$ and offsets $\mathbf{u}_{\text{offset}}$ for each control variable.
- **pScale, pOffset**: Scaling factors $\mathbf{p}_{\text{scale}}$ and offsets $\mathbf{p}_{\text{offset}}$ for each parameter.
- **TScale, TOffset**: Scaling factor T_{scale} and offset T_{offset} for the horizon length.
- **JScale**: Scaling factor J_{scale} for the cost functional.
- **cScale**: Scaling factors $\mathbf{c}_{\text{scale}}$ for each state constraint. The elements of the vector refer to the equality, inequality, terminal equality and terminal inequality constraints.

5.7 Control shift

The principle of optimality for an infinite horizon MPC problem motivates to shift the control trajectory $\mathbf{u}(t)$, $t \in [0, T]$ from the previous MPC step $k - 1$ by the sampling time Δt before the first GRAMPC iteration in the current MPC step k , cf. *Algorithm 1*. The last time segment of the shifted trajectory is hold on the last value of the trajectory. Shifting the control can lead to a faster convergence behavior of the gradient algorithm for many MPC problems.

If the control shift is activated for a problem with free end time T , GRAMPC assumes a shrinking horizon problem, because time optimization is unusual in classical model predictive control. The principle of optimality then motivates to subtract the sampling time from the horizon T after each MPC step, which corresponds to a control shift for the end time.

- **ShiftControl**: Activates or deactivates the shifting of the control trajectory and the adaptation of T in case of a free end time, i.e., if **OptimTime** is active.

5.8 Status flags

Several status flags are set in the solution structure `grampc.sol.status` during the execution of *Algorithm 1*. These flags can be printed as short messages by the function `grampc_printstatus` for the levels error, warn, info and debug.

The following status flags are printed on the level `STATUS_LEVEL_ERROR` and require immediate action:

- **STATUS_INTEGRATOR_INPUT_NOT_CONSISTENT**: This flag is set by the integrator `rodas` if the input values are not consistent. See¹ for further details.
- **STATUS_INTEGRATOR_MAXSTEPS**: This flag is set by the integrators `ruku45` or `rodas` if too many steps are required.
- **STATUS_INTEGRATOR_STEPS_TOO_SMALL**: This flag is set by the integrator `rodas` if the step size becomes too small.
- **STATUS_INTEGRATOR_MATRIX_IS_SINGULAR**: This flag is set by the integrator `rodas` if a singular Jacobian $\frac{\partial \mathbf{f}}{\partial \mathbf{x}}$ or $\left(\frac{\partial \mathbf{f}}{\partial \mathbf{x}}\right)^T$ is detected. See¹ for further details.
- **STATUS_INTEGRATOR_H_MIN**: This flag is set by the integrator `ruku45` if a smaller step size than the minimal allowed value is required.

The following flags are printed in addition to the previous ones on the level `STATUS_LEVEL_WARN`:

- **STATUS_MULTIPLIER_MAX**: This flag is set if one of the multipliers $\bar{\mu}$ reaches the upper limit μ_{\max} or the lower limit $-\mu_{\max}$. The situation may occur for example if the problem is infeasible or ill-conditioned or if the penalty parameters are too high.
- **STATUS_PENALTY_MAX**: This flag is set if one of the penalty parameters \bar{c} reaches the upper limit c_{\max} . The situation may occur for example if the problem is infeasible or ill-conditioned or if the penalty increase factor β_{in} is too high.
- **STATUS_INFEASIBLE**: This flag is set if the constraints are not satisfied and one run of *Algorithm 1* does not reduce the norm of the constraints. The situation may occur in single runs, if few iterations i_{\max} and j_{\max} are used for a suboptimal solution. However, if the flag is set in multiple successive runs, it is a strong indicator for an infeasible optimization problem.

The following flags are printed in addition to the previous ones on the level `STATUS_LEVEL_INFO`:

- **STATUS_GRADIENT_CONVERGED**: This flag is set if the convergence check is activated and the relative tolerance $\varepsilon_{\text{rel},c}$ is satisfied for the controls \mathbf{u} , the parameters \mathbf{p} , and the end time T .

¹ RODAS. Webpage. <http://www.unige.ch/~hairer/software.html>, Accessed 01-December-2018.

- **STATUS_CONSTRAINTS_CONVERGED:** This flag is set if the convergence check is activated and the absolute tolerances $\varepsilon_g, \varepsilon_h, \varepsilon_{g_T}, \varepsilon_{h_T}$ are satisfied for all constraints.
- **STATUS_LINESEARCH_INIT:** This flag is set if the gradient algorithm uses the initial step size α_{init} as fallback for the explicit line search strategy in one iteration.

The following flags are printed in addition to the previous ones on the level **STATUS_LEVEL_DEBUG**:

- **STATUS_LINESEARCH_MAX:** This flag is set if the gradient algorithm uses the maximum step size α_{max} in one iteration.
- **STATUS_LINESEARCH_MIN:** This flag is set if the gradient algorithm uses the minimum step size α_{min} in one iteration.
- **STATUS_MULTIPLIER_UPDATE:** This flag is set if the relative tolerance $\varepsilon_{\text{rel},u}$ is satisfied for the controls \mathbf{u} , the parameters \mathbf{p} and the end time T and therefore the update of the multipliers $\bar{\boldsymbol{\mu}}$ and the penalty parameters $\bar{\mathbf{c}}$ is performed, cf. *Update of multipliers and penalties*.

USAGE OF GRAMPC

This chapter describes how to use GRAMPC in C/C++ and via the interface to Matlab/Simulink and Python. Note that the implementation of the problem formulation is described in *Problem implementation* and therefore not repeated here.

6.1 Using GRAMPC in C

A high level of portability is provided by GRAMPC in view of different operating systems and hardware devices. The main components are written in plain C without the use of external libraries, also see the discussion in *Structure of GRAMPC*. This section describes the usage of GRAMPC in C including initialization, compilation and running the MPC framework.

6.1.1 Main components of GRAMPC

As illustrated in Figure Fig. 6.1.1, GRAMPC contains initializing as well as running files, different integrators for the system and adjoint dynamics (*Problem formulation and implementation* and *Optimization algorithm and options*) and functions to alter the options and parameters (also see *Problem formulation and implementation* and *Optimization algorithm and options*).

In more detail, GRAMPC comprises the following main files (see also Appendix *GRAMPC function interface* for the function interface):

- `grampc_init.c`: Functions for initializing GRAMPC.
- `grampc_alloc.c`: Functions for memory allocation and deallocation.
- `grampc_fixedsize.c`: Replaces the functions in `grampc_alloc.c` in the fixed-size mode without dynamic memory allocation.
- `grampc_run.c`: Functions for running GRAMPC including the implemented augmented Lagrangian algorithm and the underlying gradient algorithm. Further functions of this file are concerned with the line search strategies and the update steps of the primal and dual variables as described in *Optimization algorithm and options*.
- `grampc_setparam.c`: Provides several functions for setting problem-related parameters, see the description in *Problem formulation and implementation* and the list of parameters in Table Table 8.1.1.
- `grampc_setopt.c`: Provides several functions for setting algorithmic options, see the description in *Optimization algorithm and options* and the list of options in Table Table 8.2.1.
- `grampc_configreader.c`: Provides an alternative method for reading parameters and options from a text file.
- `grampc_mess.c`: Function for printing information regarding the initialization as well as execution of GRAMPC (e.g. errors, convergence behavior of the augmented Lagrangian algorithm or status of integrators).

- `grampc_util.c`: Auxiliary functions for the GRAMPC toolbox such as memory management and trajectory interpolation. It also contains the function `grampc_estim_penmin` to compute an estimate of the minimal penalty parameter, cf. *Estimation of minimal penalty parameter*.
- `simpson.c`: Function for integrating the integral cost by means of the Simpson rule.
- `trapezoidal.c`: Function for integrating the integral cost by means of the trapezoidal rule.
- `grampc_erk.c`: Explicit Runge-Kutta methods of order 1, 2, 3, 4 with fixed step size.
- `rodas.c`: Semi-implicit Rosenbrock integration scheme with variable step size. The code follows from an f2c conversion of the original Fortran files of RODAS¹ and are directly included in the GRAMPC source files.
- `ruku45.c`: Runge-Kutta integration scheme of order 4 with variable step size.
- `discrete.c`: Discrete integrator.
- `timing.c`: Optional functions for measuring the execution time.
- `finite_diff.c`: Finite difference utility functions and gradient checker.

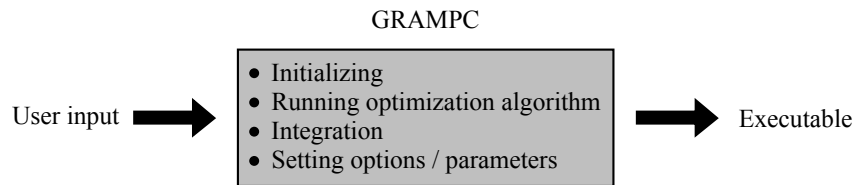


Fig. 6.1.1: Main components of GRAMPC

6.1.2 Initialization of GRAMPC

The global initialization of GRAMPC is done via the routine

```
void grampc_init(typeGRAMPC **grampc, typeUSERPARAM *userparam)
```

where the overall structure variable contains the following substructures (see *GRAMPC data types* for the definition of the data type):

- `sol` (data type `typeGRAMPCsol`): Contains the optimization variables $(\mathbf{u}^{i+1}, \mathbf{p}^{i+1}, T^{i+1})$ and the interpolated state \mathbf{x}^{i+1} in the next MPC step as well as the corresponding cost values $\bar{J}(\mathbf{u}^{i+1}, \mathbf{p}^{i+1}, T^{i+1}, \boldsymbol{\mu}^{i+1}, \mathbf{c}^{i+1}; \mathbf{x}_0)$ and $J(\mathbf{u}^{i+1}, \mathbf{p}^{i+1}, T^{i+1}; \mathbf{x}_0)$, respectively. The control \mathbf{u}^{i+1} and the state \mathbf{x}^{i+1} refer to the corresponding trajectories evaluated at the sampling time Δt . In addition, the substructure `sol` contains the evaluated functions of the defined state constraints, some status information, and an array in which the number of gradient iterations are stored in each augmented Lagrangian step.
- `param` (data type `typeGRAMPCparam`): Contains the parameter structure of GRAMPC. A detailed description of all parameters is given in *Problem formulation and implementation*.
- `opt` (data type `typeGRAMPCopt`): Contains the option structure of GRAMPC. A detailed description of all options is given in *Optimization algorithm and options*.
- `rws` (data type `typeGRAMPCrws`): Contains the real-time workspace of GRAMPC including calculations of the augmented Lagrangian algorithm and the gradient algorithm along the prediction horizon.
- `userparam` (data type `typeUSERPARAM`): Can be used to define parameters, e.g. to parametrize the cost functional, the system dynamics or the state constraints.

¹ RODAS. Webpage. <http://www.unige.ch/~haider/software.html>, Accessed 01-December-2018.

The definition of these data types is given in Appendix *GRAMPC data types*. The deallocation of is done by means of the function

```
void grampc_free(typeGRAMPC **grampc)
```

6.1.3 Setting options and parameters

The single options in Table *Algorithmic Options*. are set via the functions

```
void grampc_setopt_real(const typeGRAMPC *grampc, const typeChar *optName, ctypeRNum,
↳ optValue)
void grampc_setopt_real_vector(const typeGRAMPC *grampc, const typeChar *optName,
↳ ctypeRNum *optValue)

void grampc_setopt_int(const typeGRAMPC *grampc, const typeChar *optName, ctypeInt,
↳ optValue)
void grampc_setopt_int_vector(const typeGRAMPC *grampc, const typeChar *optName,
↳ ctypeInt *optValue)

void grampc_setopt_string(const typeGRAMPC *grampc, const typeChar *optName, const,
↳ typeChar *optValue)
```

for option values with floating point, integer and string type, respectively. An overview of the current options can be displayed by using

```
void grampc_printopt(typeGRAMPC *grampc)
```

Example (Setting options in C)

The number of gradient iterations, the integration scheme, and the relative tolerance of the integrator can be set in the following way:

```
...
/***** declaration *****/
typeGRAMPC *grampc;
...

/***** option definition *****/
/* Basic algorithmic options */
ctypeInt MaxGradIter = 2;

/* System integration */
const char* Integrator = "ruku45";
ctypeRNum IntegratorRelTol = 1e-3;
...

/***** grampc init *****/
grampc_init(&grampc, userparam);
...
```

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(continued from previous page)

```

/***** setting options *****/
grampc_setopt_int(grampc, "MaxGradIter", MaxGradIter);

grampc_setopt_string(grampc, "Integrator", Integrator);
grampc_setopt_real(grampc, "IntegratorRelTol", IntegratorRelTol);
...

```

Similar to setting the GRAMPC options, the parameters in Table *Problem-specific parameters* are set according to their data type with the following functions:

```

void grampc_setparam_real(const typeGRAMPC *grampc, const typeChar *paramName, ctypeRNum_
↳ paramValue);

void grampc_setparam_real_vector(const typeGRAMPC *grampc, const typeChar *paramName, _
↳ ctypeRNum *paramValue);

```

An overview of the parameters can be displayed by using

```

void grampc_printparam(const typeGRAMPC *grampc);

```

Example (Setting parameters in C)

The sampling time, the prediction horizon, and the initial conditions for a system with two states and one control input can be set in the following way:

```

...
/***** parameter definition *****/
ctypeRNum dt = (typeRNum)0.001;
ctypeRNum Thor = 6.0;

ctypeRNum x0[NX] = {-1,-1};
ctypeRNum u0[NU] = {0};

/***** setting parameters *****/
grampc_setparam_real(grampc, "dt", dt);
grampc_setparam_real(grampc, "Thor", Thor);
grampc_setparam_real_vector(grampc, "x0", x0);
grampc_setparam_real_vector(grampc, "u0", u0);
...

```

Config Reader

Added in version v2.3.

GRAMPC also supplies a config reader which reads .cfg files for setting the parameters and options inside the grampc struct. The config file looks like this

Listing 6.1.1: Config file for Crane-2D example

```
# GRAMPC configuration file for example CRANE_2D
#####

# GRAMPC parameter
[GRAMPC parameter]

# Initial values and setpoints of the states
x0 = [-2.000,0.000,2.000,0.000,0.000,0.000]
xdes = [2.000,0.000,2.000,0.000,0.000,0.000]

# Initial values, setpoints and limits of the inputs
u0 = [0.000,0.000]
udes = [0.000,0.000]
umax = [2.000,2.000]
umin = [-2.000,-2.000]

# Time variables
Thor = 2.00
dt = 0.002
t0 = 0.000

# GRAMPC options
[GRAMPC options]

# Basic algorithmic options
Nhor = 20

# Cost integration
TerminalCost = off

# Type of constraints' consideration
ConstraintsAbsTol = [1e-4, 1e-3, 1e-3]
```

The structure is as follows: Whitespace or tabs are trimmed. Therefore one can indent the parameter names or include spaces in the .cfg file. A parameter name follows a = sign, which then follows the parameter value. String values are plain text without whitespace. Vectors are specified with square brackets. The delimiter for the values is either a , or whitespace.

The parameters must be listed under [GRAMPC parameter]. However the order is not important. The same applies for the options which must be listed under [GRAMPC option].

Calling the config reader is simple with

```
/****** Get and set options and parameter from configuration file *****/
const char *fileName = "config_CRANE_2D.cfg";
```

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```
grampc_get_config_from_file(grampc, fileName);
```

and it sets all defined parameters and options for GRAMPC. For further examples please see the Crane-2D, Crane-3D, DAE-Integrator, Double-Integrator and the Template folder.

6.1.4 Compiling and calling GRAMPC

GRAMPC can be integrated into an executable program after the problem formulation (cf. *Problem formulation and implementation*) as well as options and parameters are provided. A Makefile for compilation purposes is provided in the folder `<grampc_root>/examples/TEMPLATES`. The main calling routine for GRAMPC is

```
void grampc_run(const typeGRAMPC *grampc)
```

The following code demonstrates how to integrate GRAMPC within an MPC loop. For a full implementation please refer to the examples inside `<grampc_root>/examples/`.

Example (C code for running GRAMPC within an MPC loop)

```
...
/*initialize grampc struct and set parameters and options*/
...
// estimate minimum penalty parameter
grampc_estim_penmin(grampc, 1);

/* MPC loop */
for (iMPC = 0; iMPC <= MaxSimIter; iMPC++) {
    grampc_run(grampc);

    /* check solver status */
    if (grampc->sol->status > 0) {
        if (grampc_printstatus(grampc->sol->status, STATUS_LEVEL_ERROR)) {
            myPrint("at iteration %i:\n -----\n", iMPC);
        }
    }

    /* reference integration or interface to real world measurements */
    get_new_state(xnext);

    /* update state and time */
    t = t + dt;
    grampc_setparam_real_vector(grampc, "x0", xnext);
}
...
```

GRAMPC is repetitively executed until a defined simulation time is reached. Note that the estimate of the minimal penalty value c_{\min} is determined via the function `grampc_estim_penmin` before the augmented Lagrangian algorithm is executed, also see the discussion in *Estimation of minimal penalty parameter*.

For a more convenience MPC design, there are several status flags that can be printed after each MPC step. As shown in the above example, the variable `grampc->sol->status` is used to check whether new status informations are

available. Subsequently, the function `grampc_printstatus` is used to visualize the corresponding informations on the console. In the current GRAMPC version, there are status informations concerning the integration scheme, the update of Lagrange multipliers, convergence properties of the augmented Lagrangian and gradient algorithm, and the line search method, see *Status flags* for more details. In addition, the GRAMPC examples provide functions to print key variables such as the system state or the controls into text files.

The examples can be compiled by running the following commands in a (Cygwin) terminal:

```
$ cd <grampc_root>/examples/*
$ make
```

The `make` command compiles the file `main_*.c` and links it against the GRAMPC toolbox, i.e., against the GRAMPC library within `<grampc_root>/libs`. Note that compiling an application example in the folder `<grampc_root>/examples/` requires the previous compilation of the GRAMPC toolbox as described in *Installation and structure of GRAMPC*. As a result, the executable `startMPC` is generated, which can now be used to solve and/or design the MPC problem.

With CMake, one can specify the build target with

```
cmake --build . --target *
```

Note that CMake outputs the executables for the examples inside `<grampc_root>/build/examples/`.

6.1.5 Using GRAMPC without dynamic memory allocation

GRAMPC restricts the usage of dynamic memory allocation, i.e. `calloc` and `realloc`, to initialization and option setting, such that no allocations are performed while the MPC is running. However, some microcontrollers and embedded devices do not support dynamic memory allocation at all. Using GRAMPC on these devices requires to replace all dynamically-sized arrays by fixed-size arrays and to remove all functions that involve dynamic memory allocation.

To this end, a header file `fixedsize_settings.h` must be placed in the search path of the compiler. This header file defines several constant parameters such as the number of states `NX`, the number of controls `NU`, as well as several constant options, e.g. the number of discretization steps `NHOR`, the number of gradient iterations `MAXGRADITER`, and the number of multiplier iterations `MAXMULTITER`. Note that these options cannot be changed during run-time, but are fixed at compile-time. A template file is provided in the folder `<grampc_root>/examples/TEMPLATES`.

In addition, the GRAMPC structures must be created on the stack instead of the heap. To this end, the macro `TYPE_GRAMPC_POINTER` is provided that allows to use the same code both with and without dynamic memory allocation:

```
int main()
{
    TYPE_GRAMPC_POINTER(grampc);
    ...
    grampc_init(&grampc, userparam);
    ...
    grampc_free(&grampc);
}
```

The makefile can be called with the parameter `FIXEDSIZE=1`, which automatically defines the required preprocessor macro `FIXEDSIZE`. Thus compiling and running the ball-on-plate example without dynamic memory allocation is done by executing the following commands in the terminal:

```
$ cd <grampc_root>/examples/*
$ make FIXEDSIZE=1
$ ./startMPC_fixedsize
```

Note that in this case a separate GRAMPC library is created in the problem folder that depends on the constants defined in `fixedsize_settings.h`. Changing these settings requires to recompile both the library and the application.

With CMake one can directly specify a fixed-size target in the respective `CMakeLists.txt` file. For the GRAMPC examples a `_fixedsize` is appended to the target name like

```
cmake --build . --target *_fixedsize
```

which directly compiles the main and GRAMPC library with the fixed-size settings.

6.2 Using GRAMPC in C++

The C++ interface provides a single class for managing the `grampc` struct and all related functions as outlined in *Using GRAMPC in C*. The main difference is the probfct handling which is described below.

6.2.1 Interface from C++ to C

The constructor of the class `Grampc` takes a pointer to a specific problem description. This problem description is of type `ProblemDescription` and holds all methods which are needed to implement the OCP. The constructor of `Grampc` now passes the `ProblemDescription*` as `userparam` to `grampc_init`. Whenever GRAMPC calls a problem specific function like `ffct`, the `userparam` pointer gets dereferenced as a `ProblemDescription*` which calls the corresponding method.

Listing 6.2.1: Example how the bridge between C -> C++ is made.

```
void ffct(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_  
↪ typeGRAMPCparam *param, typeUSERPARAM *userparam)  
{  
    ((grampc::ProblemDescription*)userparam)->ffct(out, t, x, u, p, param);  
}
```

6.2.2 Handling of Problem Descriptions

The C++ interface provides a virtual base class `ProblemDescription` under the namespace `grampc`. It provides virtual methods which map the functions from `probfct.h`.

Note

`userparam` is not needed, because problem specific quantities can be defined as class fields.

For an example please see the folder `<grampc_root>/cpp/examples/MassSpringDamper` with an implementation of the mass spring damper example. Additionally, compare with the C implementation in `<grampc_root>/examples/MassSpringDamper`.

6.3 Using GRAMPC in Matlab/Simulink

As already mentioned, the main components of GRAMPC are implemented in plain C to ensure a high level of portability. However, GRAMPC also provides a user-friendly interface with Matlab/Simulink to allow for a convenient MPC design.

6.3.1 Interface to Matlab

Each main component of GRAMPC (cf. *Main components of GRAMPC*) has a related Mex routine that is included in the directory `<grampc_root>/matlab/src`, also see Fig. 6.3.1. This allows one to run GRAMPC in Matlab/Simulink as well as altering options and parameters without recompilation.

A makefile to compile GRAMPC for use in Matlab/Simulink is provided in `<grampc_root>/matlab`. The makefile compiles the source files to generate object files within `<grampc_root>/matlab/bin`. In order to obtain an actual Mex compilation for a given problem, the object files must be linked against the object file obtained by compiling the problem function, since at least some of these functions depend on the actual problem formulation, e.g. the state dimension N_x .

The m-file `startMPC.m` in each of the examples under `<grampc_root>/examples` contains a flag `compile`. Setting this flag to 1 leads to a compilation of the problem file and to the generation of the Mex files. Setting `compile` to 2 leads to an additional recompilation of the toolbox by calling the makefile in the directory `<grampc_root>/matlab/src`. The Mex files are stored in the local subfolder `+CmexFiles`. The folder name begins with a plus sign allowing the user to call the functions stored in this folder using the command `CmexFiles.<function name>`. The S-function files are stored in the local application directory, since Simulink requires the S-functions to be in the Matlab path.

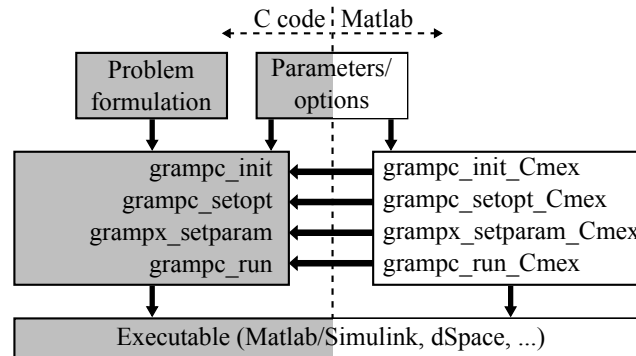


Fig. 6.3.1: Interface of GRAMPC to Matlab/Simulink (gray - C code, white - Matlab code)

The structure of the main components of GRAMPC in C and Matlab for setting options and parameters are slightly different, as it is not allowed (or at least not very elegant) to manipulate the input argument of Mex routines. Consequently, each Mex routine returns the manipulated structure variable `grampc` as an output argument. If, for example, the initial condition x_0 should be set to a specific value in C, the function `grampc_setparam_real_vector` must be used, as already discussed in *Initialization of GRAMPC*. In Matlab, however, this is done using the Mex routine `grampc_setparam_Cmex` with the structure variable `grampc` as an input argument. The manipulated structure variable `grampc` is returned as an output argument including the initial condition x_0 . For instance, the parameter setting in C

```
ctypeRNum x0[NX] = {-1,-1};
grampc_setparam_real_vector(grampc, "x0", x0);
```

would read as follows in Matlab:

```
grampc = grampc_setparam_Cmex(grampc, 'x0', [-1;-1]);
```

Note that the Mex routine `grampc_setparam_Cmex` does not distinguish between vectors and scalars, but handles the different dimensions of parameters internally. The same applies to the Mex routine `grampc_setopt_Cmex` for changing algorithmic options in GRAMPC. The data type of the parameter or option to be set can either be double or the corresponding data type in the parameter structure `param` or option structure `opt`, see also Table [Table 8.1.1](#) or Table [Table 8.2.1](#).

In order to simplify changing parameters and options in Matlab, GRAMPC also provides the routine `grampc_update_struct_grampc(grampc,user)` included in `<grampc_root>/matlab/mfiles`. The purpose of this function is to allow the user to define the options and parameters to be set as structure variable instead of requiring to call the functions `grampc_setparam_Cmex` and `grampc_setopt_Cmex` manually for each chosen parameter/option. In detail, the structure variable `user` must contain the substructures `param` and `opt` that define the parameters and options to be set. The corresponding function call under Matlab is as follows:

```
...
%% Parameter definition
% Initial values of the states
user.param.x0 = [-1;-1];
...

%% Option definition
% Basic algorithmic options
user.opt.Nhor = 10;
...

%% User parameter definition
% e.g. system parameters or weights for the cost function
userparam = [100, 10, 1, 100, 10, -0.2, 0.01, -0.1, 0.1];

%% Grampc initialization
grampc = CmexFiles.grampc_init_Cmex(userparam);

%% Update grampc struct while ensuring correct data types
grampc = grampc_update_struct_grampc(grampc,user);
...
```

Similar to *Initialization of GRAMPC* and *Compiling and calling GRAMPC*, the following lines show the main steps to run the ball-on-plate example in Matlab. An executable version of this example within Matlab can be found in the folder `<grampc_root>/examples/BallOnPlate`. In analogy to the C implementation, the simulation loop and the evaluation are implemented in the main file `startMPC.m`. The parameters and options are defined in the separate file `initData.m` that is called within `startMPC.m` for the sake of readability and to use the settings directly in the Simulink model, see *Interface to Matlab/Simulink*. Please note a template file can be found in the folder `<grampc_root>/examples/TEMPLATES`.

Example (Matlab code for setting options and parameters, see `initData.m`)

```
%% Parameter definition
user.param.x0    = [ 0.1, 0.01];
user.param.xdes  = [-0.2, 0.0];
```

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```

% Initial values, setpoints and limits of the inputs
user.param.u0    = 0;
user.param.udes  = 0;
user.param.umax  = 0.0524;
user.param.umin  = -0.0524;

% Time variables
user.param.Thor  = 0.3;           % Prediction horizon

user.param.dt    = 0.01;          % Sampling time
user.param.t0    = 0.0;           % time at the current sampling step

%% Option definition
user.opt.Nhor     = 10;           % Number of steps for the system integration
user.opt.MaxMultIter = 3;        % Maximum number of augmented Lagrangian iterations

% Constraints thresholds
user.opt.ConstraintsAbsTol = 1e-3*[1 1 1 1];

%% User parameter definition, e.g. system parameters or weights for the cost function
userparam = [100, 10, 180, 100, 10, -0.2, 0.2, -0.1, 0.1];

%% Grampc initialization
grampc = CmexFiles.grampc_init_Cmex(userparam);

%% Update grampc struct while ensuring correct data types
grampc = grampc_update_struct_grampc(grampc,user);

%% Estimate and set PenaltyMin (optional)
grampc = CmexFiles.grampc_estim_penmin_Cmex(grampc,1);ot_stat(vec,grampc,phpS);
...

```

Example (Matlab code for running GRAMPC within an MPC loop, see startMPC.m)

```

...
%% Initialization
[grampc,Tsim] = initData;
CmexFiles.grampc_printopt_Cmex(grampc);
CmexFiles.grampc_printparam_Cmex(grampc);

% init solution structure
vec = grampc_init_struct_sol(grampc, Tsim);

% init plots and store figure handles
phpP = grampc_init_plot_pred(grampc,figNr);    figNr = figNr+1;
phpT = grampc_init_plot_sim(vec,figNr);        figNr = figNr+1;
phpS = grampc_init_plot_stat(vec,grampc,figNr); figNr = figNr+1;

%% MPC loop

```

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```

i = 1;
while 1
    % set current time and current state
    grampc = C_mexFiles.grampc_setparam_Cmex(grampc, 't0', vec.t(i));
    grampc = C_mexFiles.grampc_setparam_Cmex(grampc, 'x0', vec.x(:,i));

    % run MPC and save results
    [grampc, vec.CPUtime(i)] = C_mexFiles.grampc_run_Cmex(grampc);
    vec = grampc_update_struct_sol(grampc, vec, i);

    % print solver status
    printed = C_mexFiles.grampc_printstatus_Cmex(grampc.sol.status, 'Error');

    % check for end of simulation
    if i+1 > length(vec.t)
        break;
    end

    % simulate system
    [~, xtemp] = ode45(@C_mexFiles.grampc_ffct_Cmex, vec.t(i)+[0 double(grampc.param.dt)],
        vec.x(:,i), odeopt, vec.u(:,i), vec.p(:,i), grampc.param, grampc.userparam);
    vec.x(:,i+1) = xtemp(end,:);

    % evaluate time-dependent constraints
    vec.constr(:,i) = C_mexFiles.grampc_ghfct_Cmex(vec.t(i), vec.x(:,i), vec.u(:,i),
        vec.p(:,i), grampc.param, grampc.userparam);

    % update iteration counter
    i = i + 1;

    % plot data
    grampc_update_plot_pred(grampc, phpP);
    grampc_update_plot_sim(vec, phpT);
    grampc_update_plot_stat(vec, grampc, phpS);
end

```

Similar to the C example in *Compiling and calling GRAMPC*, the structure variable `grampc` is initialized before the options as well as optional parameters are set. In addition, the plot functions (see *Plot functions*) are initialized before GRAMPC is started within an MPC loop, where the current state of the system (new initial condition) is provided to GRAMPC. After computing the new controls, the status of GRAMPC is printed, see *Status flags* for more details. Subsequently, a reference integration of the system is performed, and the constraints are evaluated before the plots are updated.

6.3.2 Interface to Matlab/Simulink

GRAMPC also allows a Matlab/Simulink integration via the S-function `grampc_run_Sfct.c` (also included in the directory `<grampc_root>/matlab/src`). A corresponding Simulink template can be found in the folder `<grampc_root>/examples/TEMPLATES` for a number of Matlab versions. The directory also contains the m-file `initData_TEMPLATE.m` which can be used for initializing GRAMPC's options and parameters, as mentioned in the previous subsection. The build procedure of the Mex routines additionally compiles the S-function for the Simulink block.

The Matlab/Simulink model of GRAMPC is shown in Fig. 6.3.2. The block MPC-Subsystem contains algorithmic components of GRAMPC (implemented within the S-function `grampc_run_Sfct.c`). The block Click to init `grampc` must be executed by a double click in order to initialize the structure variable that is required by the Matlab/Simulink model. This generates also the Simulink-specific structure variable `grampc_sdata`. For the sake of convenience, the blocks Click to compile toolbox and Click to compile `probfcn` are included in the model to be able to compile the GRAMPC toolbox and the specific problem directly from Matlab/Simulink.

The block `System` function is a Matlab Function block which implements the system dynamics in order to numerically integrate the system dynamics after each MPC step k for the sampling time Δt and to return the new state value x_{k+1} corresponding to the next sampling instant t_{k+1} that is fed back to the MPC block.

Furthermore, the S-function `grampc_run_Sfct.c` satisfies the additional restrictions of the Matlab code generator. Therefore, the block can be used in models implemented for running on various hardware platforms, such as dSpace real-time systems. Please note that especially in case of dSpace applications, the include folders `<grampc_root>/include` and `<grampc_root>/matlab/include` as well as all C source files in `<grampc_root>/src`, the source file of the S-function `<grampc_root>/matlab/src/grampc_run_Sfct.c` and the problem function must be listed as additional build information in the Model Configuration Parameters of the Simulink model under Code Generation / Custom Code. It is recommended to use absolute paths at least for the S-function file.

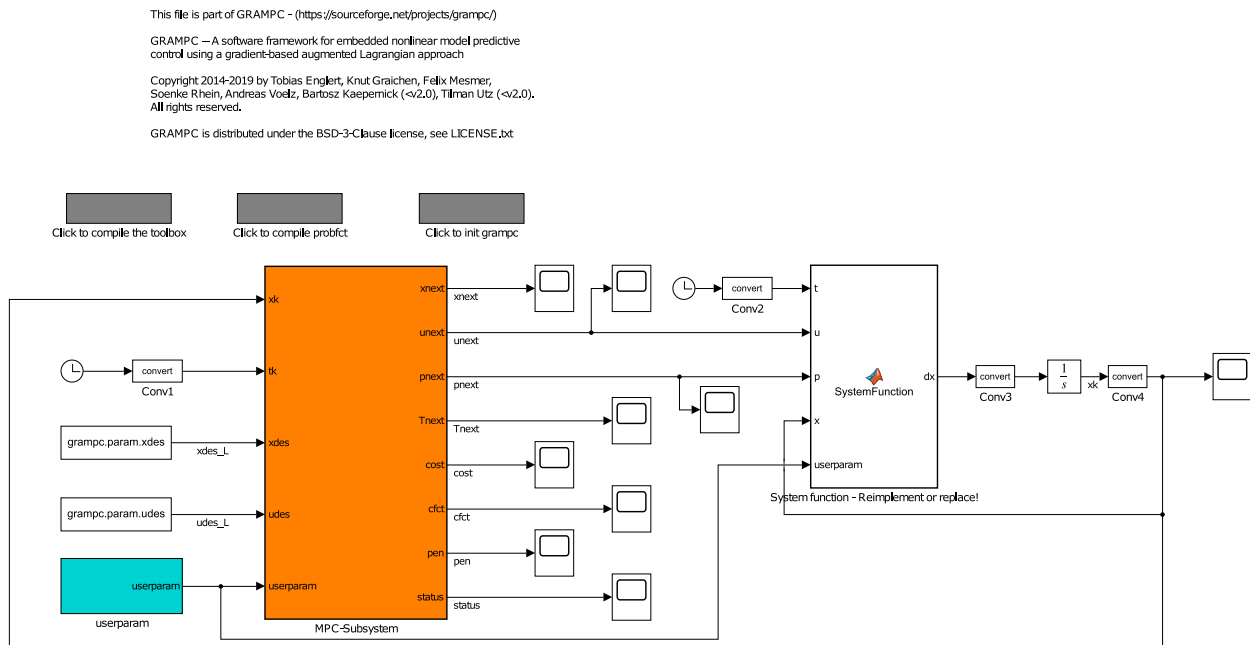


Fig. 6.3.2: Matlab/Simulink model of GRAMPC.

6.3.3 Plot functions

GRAMPC offers various plot functions in the folder `<grampc_root>/matlab/mfiles`. Each plot must be initialized at first using the m-files `grampc_init_plot_*.m`. During the simulation the plots can be updated by the m-files `grampc_update_plot_*.m`. Beside the trajectories of the simulated system dynamics (file ending `*=sim`) and trajectories along the prediction horizon (file ending `*=pred`), also some statistics (file ending `*=stat`) can be plotted. When solving OCPs instead of MPC problems, the plot along the prediction horizon shows the actual results. The plotted quantities depend on the parameter and option settings of GRAMPC, i.e. whether constraints are considered or not. The available plots are listed in more detail in the following lines. (Also see the example problems under `<grampc_root>/examples` for code samples on how to use the plot routines.)

System dynamics plot (`plot_sim`)

- **States:** This plot illustrates the trajectories of the state x along the simulation time.
- **Adjoint states:** This plot illustrates the trajectories of the adjoint state λ along the simulation time.
- **Controls:** This plot illustrates the trajectories of the controls u along the simulation time.
- **Constraints:** This plot appears only if equality and/or inequality constraints are defined (N_g and/or N_h is larger than zero as specified in `ocp_dim`). The plot shows the trajectories of the constraints g and h along the simulation time.
- **Lagrange multipliers:** This plot appears only if equality and/or inequality constraints are defined. The plot shows the trajectories of the multipliers μ_g and μ_h along the simulation time. If any Lagrange multiplier reaches the limit μ_{\max} (specified by `MultiplierMax`), it indicates that the penalty parameters are too high or that the problem is not well-conditioned or that the costs are badly scaled.
- **Penalty parameters:** This plot appears only if equality and/or inequality constraints are defined. The plot shows the trajectories of the penalties c_g and c_h along the simulation time. If any penalty reaches the maximum value c_{\max} , set by `PenaltyMax`, it indicates that either the limit is not high enough or the update is too aggressive.

Prediction plot (`plot_pred`)

- **Predicted states:** This plot illustrates the trajectories of the state x along the prediction horizon.
- **Predicted adjoint states:** This plot illustrates the trajectories of the adjoint state λ along the prediction horizon.
- **Predicted controls:** This plot illustrates the trajectories of the controls u along the prediction horizon.
- **Predicted constraints:** This plot appears only if (terminal) equality and/or inequality constraints are defined. The plot shows the predicted violation of the equality constraints g and inequality constraints $\max(h, 0)$ along the prediction horizon and the predicted violation of the terminal equality constraints g_T and inequality constraints $\max(h_T, 0)$ at the end of the prediction horizon. Please note that except for OCPs, these are not the actual but predicted internal constraint violations of the current GRAMPC iteration.
- **Lagrange multipliers:** This plot appears only if (terminal) equality and/or inequality constraints are defined. The plot shows the trajectories of the multipliers μ_g and μ_h along the prediction horizon and the multipliers μ_{g_T} and μ_{h_T} . If any Lagrange multiplier reaches the limit μ_{\max} , set by `MultiplierMax`, it indicates that the penalty parameters are too high or that the problem is not well-conditioned or that the costs are badly scaled.
- **Penalty parameters:** This plot appears only if (terminal) equality and/or inequality constraints are defined. The plot shows the trajectories of the penalties c_g and c_h along the prediction horizon and the penalties c_{g_T} and c_{h_T} . If any penalty reaches the maximum value c_{\max} set by `PenaltyMax`, it indicates that either the limit is not high enough or the update is too aggressive, see also [Update of multipliers and penalties](#).

Statistics plot (plot_stat)

- **Costs:** This plot illustrates the costs J along the simulation time or along the augmented Lagrangian iterations. If constraints are defined, the augmented costs \bar{J} are shown as well.
- **Computation time:** This plot illustrates the computation time of one MPC or optimization step of GRAMPC along the simulation time or along the augmented Lagrangian iterations. The time measurement is done in the function `grampc_run_Cmex.c` using operating system specific timer functions. Consequently, the time excludes the overhead resulting from the Mex interface as well as the time consumed by the plot functions.
- **Line search step size:** This plot illustrates the step size α of the last gradient iteration along the simulation time or along the augmented Lagrangian iterations. If the adaptive line search is used (see [Adaptive line search](#)), the plot also illustrates the three corresponding sample points α_1 , α_2 , and α_3 . A step size equal to the maximum or minimum value α_{\max} or α_{\min} indicates that these values may have to be adapted or the problem may have to be scaled. Additionally, if the explicit line search is chosen and the fallback strategy is not activated (see [Explicit line search](#) and [Fallback strategy for explicit line search](#)), a frequent use of the initial value α_{init} indicates an ill-conditioned problem.
- **Gradient iterations:** This plot appears only if the option `ConvergenceCheck` is set to `on`. It illustrates the number of executed gradient iterations along the simulation time or along the augmented Lagrangian iterations. In particular, the plot depicts whether the maximum number of gradient iterations j_{\max} is reached or the convergence check caused a premature termination of the minimization.
- **Prediction horizon:** This plot appears only if the option `OptimTime` is set to `on`. It illustrates the prediction horizon T along the simulation time or along the augmented Lagrangian iterations. In shrinking horizon applications the value should decrease linearly after a short settling phase.
- **Norm of constraints over horizon:** This plot appears only if constraints are defined. It illustrates the norm $\frac{1}{T} \sqrt{\|g\|_{L_2}^2 + \|\max(h, 0)\|_{L_2}^2 + \|g_T\|_2^2 + \|\max(h_T, 0)\|_2^2}$ over all predicted constraints plotted over the simulation time or the augmented Lagrangian iterations. Especially when solving OCPs, the value should decrease continuously.
- **Norm of penalty parameters over horizon:** This plot appears only if the number of equality N_g , inequality N_h , terminal equality N_{g_T} or terminal inequality N_{h_T} constraints is not zero. It illustrates the norm $\frac{1}{T} \sqrt{\|\bar{c}\|_{L_2}}$ over all predicted penalty parameters along the simulation time or along the augmented Lagrangian iterations.
- **Terminal constraints:** This plot appears only if terminal constraints are defined. It illustrates the violation of the terminal equality constraints g_T and inequality constraints $\max(h_T, 0)$ along the simulation time or along the augmented Lagrangian iterations in case of OCPs.
- **Terminal Lagrangian multipliers:** This plot appears only if terminal constraints are defined. It illustrates the multipliers μ_{g_T} and μ_{h_T} along the simulation time or along the augmented Lagrangian iterations in case of OCPs.
- **Terminal penalty parameters:** This plot appears only if terminal constraints are defined. It illustrates the penalties c_{g_T} and c_{h_T} along the simulation time or along the augmented Lagrangian iterations in case of OCPs.

6.4 Using GRAMPC in Python

Added in version v2.3.

The Python interface for GRAMPC is similar to the Matlab interface. It furthermore offers the possibility to write the problem formulation in Python code without a compilation step for rapid prototyping.

6.4.1 Problem Formulation in C++

The first step to implement an MPC in Python is writing the problem definition. Doing so in C++ is the preferred way, since the computation speed of GRAMPC can be leveraged. Writing the problem definition itself does not differ greatly from the C++ interface. Instead of array pointers, e.g. `typeRNum *out`, matrices from Eigen are used. They are defined by

```
typedef Eigen::Matrix<typeRNum, Eigen::Dynamic, 1> Vector;
typedef Eigen::Matrix<typeRNum, Eigen::Dynamic, Eigen::Dynamic> Matrix;
typedef Eigen::Ref<Vector> VectorRef;
typedef const Eigen::Ref<const Vector>& VectorConstRef;
```

and offer a direct conversion to numpy for the Python interface. Accessing an element of `Vector`, `VectorRef` or `VectorConstRef` is the same as for an array pointer.

The class itself has to inherit from `PyProblemDescription`. An example of the problem specific header is given in `<grampc root>/python/examples/Crane2D` by

```
#include "pygrampc_problem_description.hpp" // <pybind11/pybind11.h> is already included,
↪ here
#include <vector>
#include <pybind11/stl.h>
#include <cmath>

using namespace grampc;

class Crane2D : public PyProblemDescription // must be inherited with public
{
    public: // writing the class fields public reduces overhead code for writing getters,
    ↪ and setters
        std::vector<typeRNum> Q_;
        std::vector<typeRNum> R_;
        typeRNum ScaleConstraint_;
        typeRNum MaxConstraintHeight_;
        typeRNum MaxAngularDeflection_;

        public:
            Crane2D(std::vector<typeRNum> Q, std::vector<typeRNum> R, ctypeRNum,
    ↪ ScaleConstraint, ctypeRNum MaxConstraintHeight, ctypeRNum MaxAngularDeflection);

            ~Crane2D() {}
```

Differently from C and the C++ interface, no `ocp_dim()` function is present to define the problem dimensions. Instead, they are passed in the constructor of `PyProblemDescription` with e.g. for `Crane2D`:


```
Crane2D::Crane2D(std::vector<typeRNum> Q, std::vector<typeRNum> R, ctypeRNum
→ ScaleConstraint, ctypeRNum MaxConstraintHeight, ctypeRNum MaxAngularDeflection)
: PyProblemDescription(/*Nx*/ 6, /*Nu*/ 2, /*Np*/ 0, /*Ng*/ 0, /*Nh*/ 3, /*NgT*/ 0, /
→ /*NhT*/ 0),
  Q_(Q),
  R_(R),
  ScaleConstraint_(ScaleConstraint),
  MaxConstraintHeight_(MaxConstraintHeight),
  MaxAngularDeflection_(MaxAngularDeflection)
{}

```

After writing the problem specific code, the Python wrapper with pybind11 has to be written. The binding code is wrapped in the macro PYBIND11_MODULE(module_name, module_reference) which is the entry point for the Python interpreter when importing a pybind11 module. For the Crane2D example it is given by

```
PYBIND11_MODULE(crane_problem, m)
{
    /** Imports pygrampc so this extensions module knows of the type_
→ grampc::PyProblemDescription, otherwise an import error like
    * ImportError: generic_type: type "Crane2D" referenced unknown base type
→ "grampc::PyProblemDescription"
    * may occur, if the problem description is imported before pygrampc.
    */
    pybind11::module_::import("pygrampc");

    pybind11::class_<Crane2D, PyProblemDescription, std::shared_ptr<Crane2D>>(m, "Crane2D
→ ")
    ...

```

At first, pygrampc is imported so our derived problem definition knows the already bound type PyProblemDefinition. This is just a convenience. After that, the Python class definition is written with pybind11::class_<>. Here, we start with our custom class, then the parent we are inheriting from, and also the used capsule for reference counting. A shared pointer is advisable since otherwise Crane2D can be prematurely destructed if no reference in Python is present. In the end of the line, we pass the module reference, here m and the class name in Python.

The __init__() function for Python is defined by

```
...
    .def(pybind11::init<std::vector<typeRNum>, std::vector<typeRNum>, typeRNum, typeRNum,
→ typeRNum>())
    ...

```

where a type list of the C++ constructor is supplied. Note that for binding std::vector<> the header pybind11/stl.h has to be included.

This is the theoretic bare minimum needed to initialize the C++ class in Python and pass to GRAMPC. It is convenient to also expose class fields or functions to Python. This can be done with

```
...
    .def_readonly("Nx", &Crane2D::Nx)
    .def_readonly("Nu", &Crane2D::Nu)
    .def_readonly("Np", &Crane2D::Np)
    .def_readonly("Ng", &Crane2D::Ng)

```

(continues on next page)

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```

.def_readonly("Nh", &Crane2D::Nh)
.def_readonly("NgT", &Crane2D::NgT)
.def_readonly("NhT", &Crane2D::NhT)

// make your custom fields available from python
.def_readwrite("Q", &Crane2D::Q_)
.def_readwrite("R", &Crane2D::R_)
.def_readwrite("MaxAngularDeflection", &Crane2D::MaxAngularDeflection_)
.def_readwrite("ScaleConstraint", &Crane2D::ScaleConstraint_)
.def_readwrite("MaxConstraintHeight", &Crane2D::MaxConstraintHeight_);
}

```

where in this case only fields are exposed. For a more detailed guide please look into the pybind11 documentation.

The next step involves writing the CMakeLists.txt file for compiling our problem definition. We start with configuring the languages and finding Python, pybind11 and Eigen.

```

cmake_minimum_required(VERSION 3.15)
project(crane_problem_project LANGUAGES CXX C)

find_package(Python REQUIRED COMPONENTS Interpreter Development.Module)
find_package(pybind11 CONFIG REQUIRED)

# Eigen 3.4 is required
find_package(Eigen3 3.4 REQUIRED NO_MODULE)

```

Then the path to the GRAMPC and PyGRAMPC header files has to be defined:

```

# Path to <grampc_root>
set(GRAMPC_ROOT
    ../../../../
)

include_directories(
    ${GRAMPC_ROOT}include # include directory of GRAMPC
    ${GRAMPC_ROOT}python/include # include directory of PyGRAMPC
)

```

Finally the pybind11 module is defined with

```

pybind11_add_module(crane_problem MODULE Crane2D.cpp)
target_link_libraries(crane_problem PRIVATE Eigen3::Eigen) # linking against Eigen is
↪ necessary
install(TARGETS crane_problem DESTINATION .) # copy the .pyd (Windows) or .so (Linux) to
↪ the correct installation folder where Python finds the extension

```

Note that pybind11_add_module is similar to add_executable from CMake. Here all source files and the target is defined. The install command is necessary to move the compiled module into site-packages, so Python can directly import the problem definition.

Important

The CMake target in pybind11_add_module and the module name in PYBIND11_MODULE has to be same!

To actually make the problem definition importable by Python, it needs a proper Python module definition. This is done by supplying a `pyproject.toml` file given by

```
[build-system]
requires = ["scikit-build-core>=0.10", "pybind11"]
build-backend = "scikit_build_core.build"

[project]
name = "crane_problem"
version = "1.0"
description="Compiled Crane2D problem for GRAMPC"

[tool.scikit-build]
wheel.expand-macos-universal-tags = true
```

In `[build-system]` the requirements for building this Python module are defined, namely `pybind11` and `scikit-build-core`. `scikit-build-core` takes care of building the extension with CMake and also installs the module into site-packages.

With that, the Crane2D C++ problem definition can be installed with pip with

```
$ cd <grampc_root>/python/examples/Crane2D
$ pip install .
```

and then imported by

```
from crane_problem import Crane2D
```

For an example of the project layout, please refer to `<grampc root>/python/examples/Crane2D`.

6.4.2 Problem Formulation in Python

The Python interface also allow to write the problem definition in Python.

Attention

The full speed of GRAMPC via Python is only reachable when writing the problem definition in C++! Writing the problem definition in Python can result in 100x longer computation times or more.

This leverages rapid prototyping without a compilation step, but should not be used for evaluating the computation times of GRAMPC. PyGRAMPC provides the class `ProblemDescription` which redirects the C function calls to Python. An example for defining the problem definition in Python is given in the `DoubleIntegrator` example by

Listing 6.4.1: Example for the `__init__` method in Python on the basis of the `DoubleIntegrator` example.

```
from pygrampc import ProblemDescription

class DoubleIntegrator(ProblemDescription);
    def __init__(self)
        ProblemDescription.__init__(self, Nx=2, Nu=1, Np=0, Ng=0, Nh=0, NgT=2, NhT=0)

        self.CostIntegral = 0.1
        self.CostTerminal = 1.0
```

Like in C++, the problem dimensions are set in the constructor of `ProblemDescription`. Note that every field has to be set. Just like in the C++ interface, only `__init__()`, `ffct()` and `dfdx_vec()` are mandatory to implement.

A sample function implementation looks like

```
def ffct(self, out, t, x, u, p, param):
    out[0] = x[1]
    out[1] = u[0]
```

where `t` is a float and `param` the `grampc_param` struct. The other parameters are numpy arrays, which point to memory allocated by GRAMPC. Thus, the results must explicitly write into the allocated memory so statements like

```
out[0] = ...
out[:] = ...
```

correctly set values to `out`. Note that `out = out + 1` computes `out + 1` and saves the result in the new variable `out`, which does not point to the allocated memory from GRAMPC. For a correct usage, please refer to the examples in `<grampc root>/python/examples`.

6.4.3 Usage in Python

The usage of GRAMPC in Python is described via the `DoubleIntegrator.py` example in `<grampc root>/python/examples/DoubleIntegrator`. We first start with importing the relevant packages

```
from pygrampc import ProblemDescription, Grampc, GrampcResults
import matplotlib.pyplot as plt
from scipy.integrate import solve_ivp

# alternatively define your problem definition
# as a C++ extension module or in a Python file
# and import the corresponding problem definition
```

Here, the `solve_ivp` function from Scipy is used for the reference integration. Scipy is easily installed with `pip install scipy` into the current environment. In this example, the problem definition is directly written in the same file for rapid prototyping. Thus, it does not need to be imported. After that, GRAMPC is initialized:

```
if __name__ == "__main__":

    ...

    options = "DoubleIntegrator.json"

    # initialize problem and GRAMPC
    Problem = DoubleIntegrator()
    grampc = Grampc(Problem, options, plot_prediction=False)
```

We start with a so-called import guard with `if __name__ == "__main__":` which only executes if the current file is run as a main file. With that, the problem definition in this file can be imported without executing e.g. our test code defined here. Then, the `DoubleIntegrator` class is initialized. After that, GRAMPC is initialized with `Problem` and also with a path to a json file with problem specific options. This options file is optional. We can also turn on prediction plots for debugging the current MPC formulation.

Like in Matlab, the result struct `GrampcResults` is supplied, which mimics the result and statistic plots available in Matlab. It is constructed by

```
# construct solution structure
vec = GrampcResults(grampc, Tsim, plot_results=True, plot_statistics=True)
```

The specific MPC loop for the Double-Integrator is given by

```
dt = grampc.param.dt

for i, t in enumerate(vec.t):
    vec.CPUtime[i] = grampc.run()
    vec.update(grampc, i)

    if i + 1 > len(vec.t) or vec.t[i] > Tsim:
        break

    # simulate system
    sol = solve_ivp(grampc.ffct, [t, t + dt], grampc.param.x0,
                    args=(grampc.sol.unext, grampc.sol.pnext, grampc.param))

    # set current time and state
    grampc.set_param({"x0": sol.y[:, -1],
                     "t0": t + dt})

    if grampc.sol.Tnext <= grampc.param.Tmin + grampc.param.dt and bool(grampc.opt.
    OptimTime):
        grampc.set_opt({"OptimTime": "off"})
        Tsim = vec.t[i + 1]

    # plots of the grampc predictions
    if i % plotSteps == 0:
        grampc.plot()
        vec.plot()
        if plotPause:
            input("Press Enter to continue...")
```

First, we call `grampc.run()` which returns the computation time in milliseconds. After that, the solution struct is updated with the current results. With `solve_ivp` the reference integration is carried out, with the wrapped `ffct` like in Matlab. Here a custom reference integration or model can be implemented. `Grampc` also provides `set_param` and `set_opt` to change parameters and options in GRAMPC. You have to always pass a Python dict with the respective key-value pairs like

```
parameters = {
    "x0": [1.05, 2.0],
    "u0": [0.05,]
}
grampc.set_param(parameters)
```

In the end of the MPC loop, the prediction, results and statistic plots are plotted, if set to `True`.

To run the example, either use your preferred python code editor, or run directly from the terminal with

```
$ cd <grampc_root>/python/examples/DoubleIntegrator
$ python DoubleIntegrator.py
```

For the usage in Python code, please refer to the examples in `<grampc root>/python/examples`.

TUTORIALS

This chapter presents some application examples of GRAMPC and how to tune GRAMPC to improve the performance compared to the default settings. In addition to a model predictive control and an optimal control problem, a moving horizon estimation example is presented. Please note that the plot functions, described in [Plot functions](#), and the status of GRAMPC, see [Status flags](#), also provide important information for tuning GRAMPC.

7.1 Model predictive control of a PMSM

The torque or current control of a permanent magnet synchronous machine (PMSM) is a challenging example for nonlinear constrained model predictive control. The following subsections illustrate the problem formulation as well as useful options of GRAMPC to improve the control performance. Corresponding m and C files can be found in the folder `<grampc_root>/examples/PMSM`.

7.1.1 Problem formulation

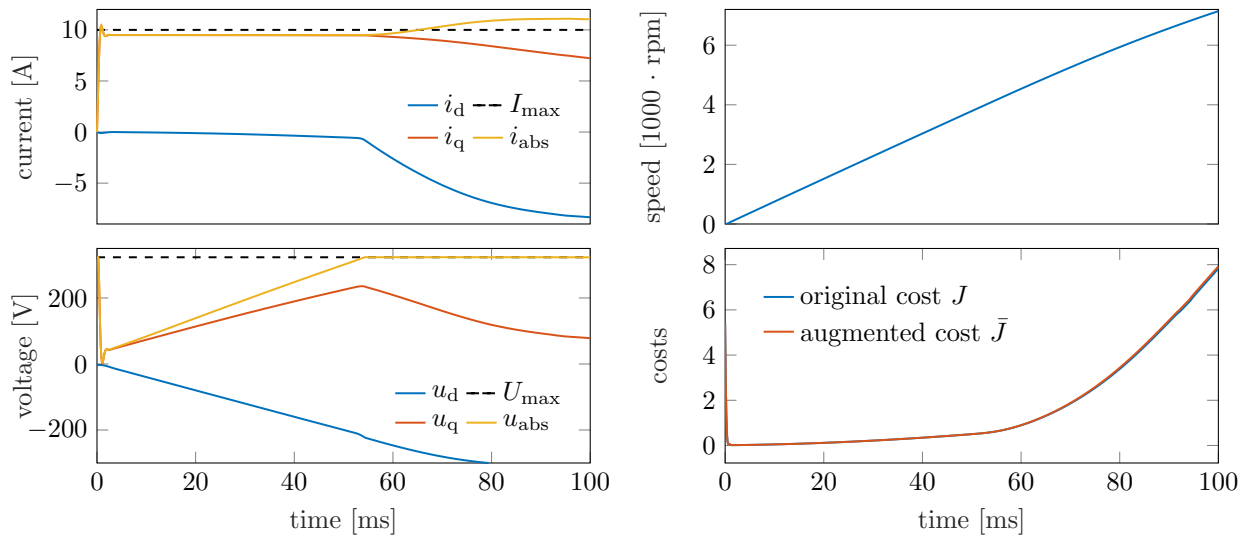


Fig. 7.1.1: Simulated MPC trajectories for the PMSM example with default settings.

The system dynamics of a PMSM¹

$$\begin{aligned}
 L_d \frac{d}{dt} i_d &= -R i_d + L_q \omega i_q + u_d \\
 L_q \frac{d}{dt} i_q &= -R i_q - L_d \omega i_d - \omega \psi_p + u_q \\
 J \frac{d}{dt} \omega &= \left(\frac{3}{2} z_p (\psi_p i_q + i_d i_q (L_d - L_q)) - \frac{\mu_f}{z_p} \omega - T_L \right) z_p \\
 \frac{d}{dt} \phi &= \omega
 \end{aligned} \tag{7.1.1}$$

is given in the dq-coordinates. The system state $\mathbf{x} = [i_d, i_q, \omega, \phi]^T$ comprises the dq-currents, the electrical rotor speed as well as the electrical angle. The dq-voltages serve as controls $\mathbf{u} = [u_d, u_q]^T$. Further system parameters are the stator resistance $R = 3.5 \Omega$, the number of pole-pairs $z_p = 3$, the permanent magnet flux $\psi_p = 0.17 \text{ V s}$, the dq-inductivities $L_d = L_q = 17.5 \text{ mH}$, the moment of inertia $J = 0.9 \text{ g m}^2$ as well as the friction coefficient $\mu_f = 0.4 \text{ mN m s}$.

The magnitude of the dq-currents is limited by the maximum current $I_{\max} = 10 \text{ A}$, i.e.

$$i_{\text{abs}} = i_d^2 + i_q^2 \leq I_{\max}^2, \tag{7.1.2}$$

in order prevent damage of the electrical components. Another constraint concerns the dq-voltages. Through the modulation stage between the controller and the voltage source inverter, the dq-voltages are limited inside the circle

$$u_{\text{abs}} = u_d^2 + u_q^2 \leq U_{\max}^2 \tag{7.1.3}$$

with the maximum voltage $U_{\max} = 323 \text{ V}$.

The optimal control problem

$$\begin{aligned}
 \min_{\mathbf{u}} \quad & J(\mathbf{u}; \mathbf{x}_k) = \int_0^T l(\mathbf{x}(\tau), \mathbf{u}(\tau)) d\tau \\
 \text{s.t.} \quad & \dot{\mathbf{x}}(\tau) = \mathbf{f}(\mathbf{x}(\tau), \mathbf{u}(\tau)), \quad \mathbf{x}(0) = \mathbf{x}_k \\
 & h_1(\mathbf{x}(\tau)) = x_1(\tau)^2 + x_2(\tau)^2 - I_{\max}^2 \leq 0 \\
 & h_2(\mathbf{u}(\tau)) = u_1(\tau)^2 + u_2(\tau)^2 - U_{\max}^2 \leq 0 \\
 & \mathbf{u}(\tau) \in [\mathbf{u}_{\min}, \mathbf{u}_{\max}]
 \end{aligned}$$

is subject to the system dynamics given by (7.1.1) and the constraints given by (7.1.2) and (7.1.3). The control constraints (7.1.3) are nonlinear and are therefore handled by the augmented Lagrangian framework and not by the projection gradient method itself. It is therefore reasonable to add the box constraints for \mathbf{u} with $\mathbf{u}_{\min} = [-U_{\max}, -U_{\max}]^T$ and $\mathbf{u}_{\max} = [U_{\max}, U_{\max}]^T$ to the OCP formulation to enhance the overall robustness of the algorithm. The cost functional consists of the integral part

$$l(\mathbf{x}, \mathbf{u}) = q_1 (i_d - i_{d,\text{des}})^2 + q_2 (i_q - i_{q,\text{des}})^2 + (\mathbf{u} - \mathbf{u}_{\text{des}})^T \mathbf{R} (\mathbf{u} - \mathbf{u}_{\text{des}}),$$

with the setpoints for the states $i_{d,\text{des}}, i_{q,\text{des}}$ and controls $\mathbf{u}_{\text{des}} \in \mathbb{R}^2$ respectively. The weights are set to $q_1 = 8 \text{ A}^{-2}$, $q_2 = 200 \text{ A}^{-2}$ and $\mathbf{R} = \text{diag}(0.001 \text{ V}^{-2}, 0.001 \text{ V}^{-2})$. The example considers a startup of the motor from standstill by defining the setpoints $i_{d,\text{des}} = 0 \text{ A}$ and $i_{q,\text{des}} = 10 \text{ A}$, corresponding to a constant torque demand of 7.65 N m . The desired controls are set to $\mathbf{u}_{d,\text{des}} = [0 \text{ V}, 0 \text{ V}]^T$.

The resulting OCP is solved by GRAMPC with the sampling time $\Delta t = 125 \mu\text{s}$ (parameter `dt`) and the horizon $T = 5 \text{ ms}$ (parameter `Thor`) using standard options almost exclusively. Only the number of discretization points `Nhor=11`, the number of gradient iterations `MaxGradIter=3` and the number of augmented Lagrangian iterations `MaxMultIter=3` are adapted to the problem. In addition, the constraints tolerances `ConstraintsAbsTol` are set to 0.1% of the respective limit, i.e. 0.1 A^2 and 104.5 V^2 . `PenaltyMin` is set to 2.5×10^{-7} by the estimation method of GRAMPC, see [Estimation of minimal penalty parameter](#).

¹ T. Englert and K. Graichen. Nonlinear model predictive torque control of PMSMs for high performance applications. *Control Engineering Practice*, 81:43 – 54, 2018.

Fig. 7.1.1 illustrates the simulation results. The setpoints are reached very fast and are stabilized almost exactly. However, an overshoot can be observed, which also leads to a small violation of the dq-current constraint by 0.45 A. With increasing rotor speed, the voltage also increases until the voltage constraint becomes active. While the voltage constraint is almost exactly hold, the dq-current constraint is clearly violated. The figure shows that at the end of the simulation the dq-current constraint violation is more than 1 A or 10%. Though a larger number of iterations might be used to reduce the constraint violation, the main reason for this deviation is that the nonlinear voltage and current constraints differ in several orders of magnitude. The next section therefore shows how to scale the problem in GRAMPC.

Also note that the increase of the cost functional does not indicate instability, but can be explained by the increasing speed, which affects the control term in the cost with the control setpoints ($\mathbf{u}_{d,des} = [0 \text{ V}, 0 \text{ V}]^T$). Moreover, the current setpoints ($i_{d,des} = 0 \text{ A}$, $i_{q,des} = 10 \text{ A}$) cannot be hold due to the constraints (7.1.2) and (7.1.3), which leads to an additional cost increase.

7.1.2 Constraints scaling

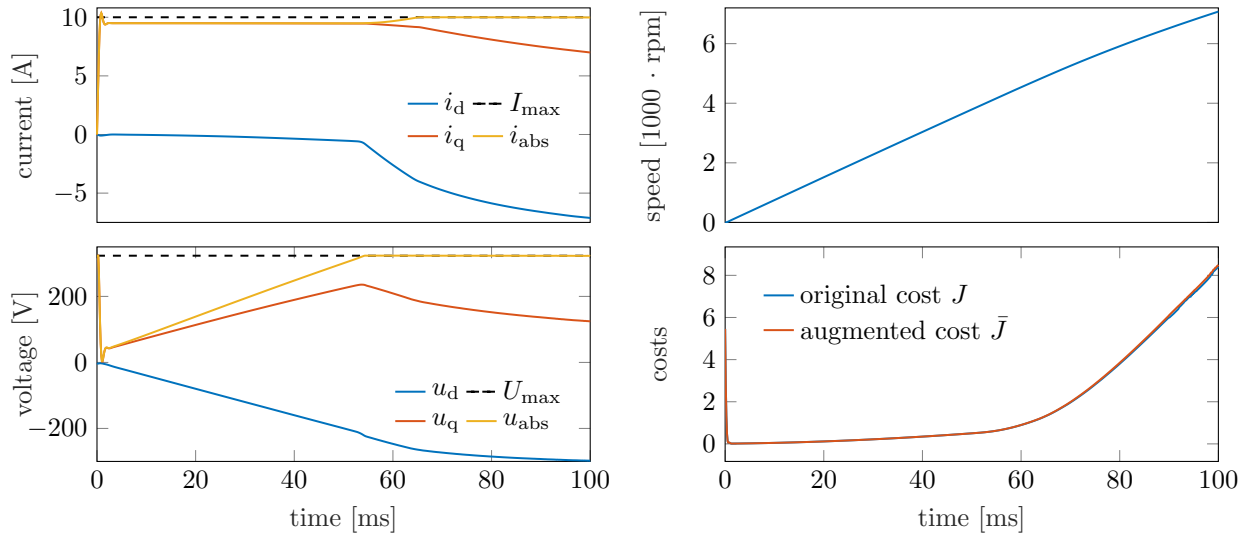


Fig. 7.1.2: Simulated MPC trajectories for the PMSM example with scaled constraints.

The two spherical constraints (7.1.2) and (7.1.3) lie in very different orders of magnitude, i.e. $I_{\max}^2 = 100 \text{ A}^2$ and $U_{\max}^2 = 104329 \text{ V}^2$. Consequently, the two constraints should be scaled by the maximum value

$$\frac{i_d^2 + i_q^2}{I_{\max}^2} - 1 \leq 0, \quad \frac{u_d^2 + u_q^2}{U_{\max}^2} - 1 \leq 0.$$

This scaling can either be done by hand directly in the problem function or by activating the option `ScaleProblem` and setting `cScale = [I_max^2, U_max^2]`. The scaling option of GRAMPC, however, causes additional computing effort (approx. 45% for the PMSM problem). Hence, this option is suitable for testing the scaling, but eventually should be done manually in the problem formulation to achieve the highest computational efficiency. In accordance with the scaling of the constraints, the tolerances are also adapted to 1×10^{-3} corresponding to 0.1% of the scaled constraints limits.

Besides the scaling and `PenaltyMin` that is set to 2×10^3 by the estimation routine of GRAMPC, all parameters and options are the same as in the last subsection. Please note that the estimation method for `PenaltyMin` strongly depends on reasonable constraint tolerances. In general, the method returns rather conservative values, which may lead to constraint violations if the order of magnitude of the constraints is very different.

Fig. 7.1.2 shows a clear improvement in terms of the dq-current constraint that is now fully exploited. The only violation results from the overshoot at the beginning, which is in the same range as in the unscaled case (approx. 0.35 A). Further

improvements, e.g. reduction of the overshoot, can be achieved by optimizing the penalty update as described in the next subsection.

7.1.3 Optimization of the penalty update

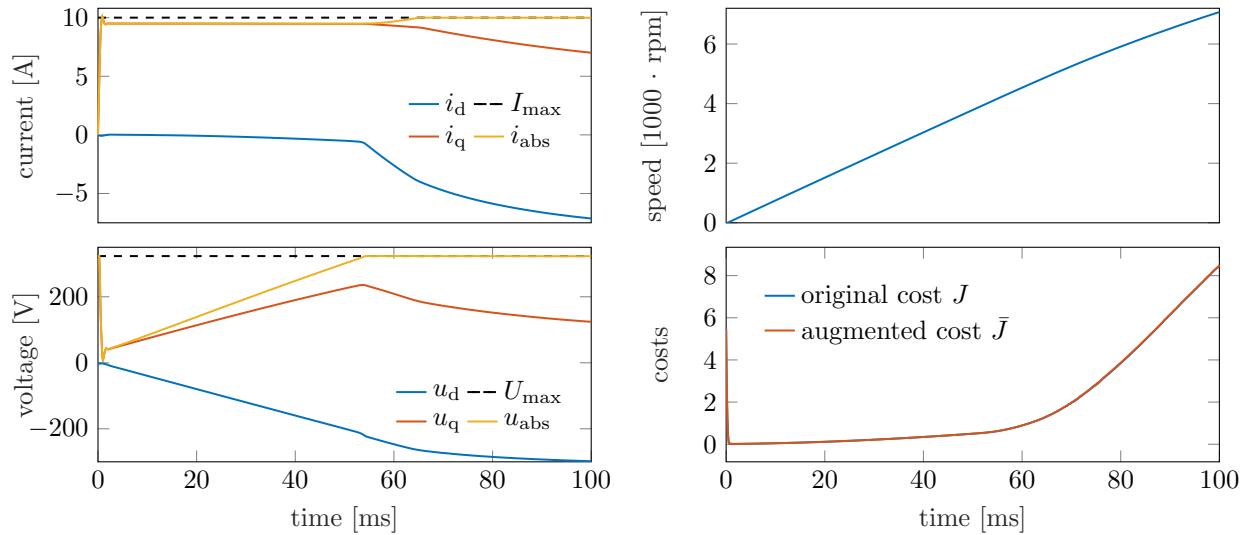


Fig. 7.1.3: Simulated MPC trajectories for the PMSM example with scaled constraints and optimized penalty update.

In order to improve compliance with the dq-current constraint at the beginning of the simulation, the number of augmented Lagrangian updates is increased. To this end `AugLagUpdateGradientRelTol` is raised to 1, which means that in every outer iteration an update of the multipliers and penalties is performed, even if the inner minimization is not converged. Furthermore `PenaltyMin` is raised to 1×10^4 compared to the estimated value of 2×10^3 . In addition, the plot of the step size, see the plot functions described in [Plot functions](#), shows that the maximum value α_{\max} is often used. Consequently, setting the maximum step size `LineSearchMax` to 10 allows larger optimization steps, especially at the beginning and at the end of the simulation. All other parameters and options, in particular the scaling options, are the same as in the previous subsection.

Fig. 7.1.3 illustrates the simulation result with the optimized penalty update. The initial dq-current overshoot is further reduced and the constraint is only violated by less than 0.07 A. Furthermore, no oscillations occur in the costs and the augmented and original cost are almost the same, which indicates that GRAMPC is well tuned.

The computation time on a Windows 10 machine with Intel(R) Core(TM) i5-5300U CPU running at 2.3 GHz using the Microsoft Visual C++ 2013 Professional (C) compiler amounts to 0.032 ms. On the dSpace real-time hardware DS1202, the computation time is 0.13 ms.

7.2 Optimal control of a double integrator

This section describes how GRAMPC can be used to solve OCPs by considering the example of a double integrator problem. The problem formulation includes equality and inequality constraints. Both the control variable u and the end time T serve as optimization variables. In addition to the problem formulation of the OCP, one focus of the following discussion will be the appropriate convergence check of the augmented Lagrangian and gradient method, and the optimization of the end time.

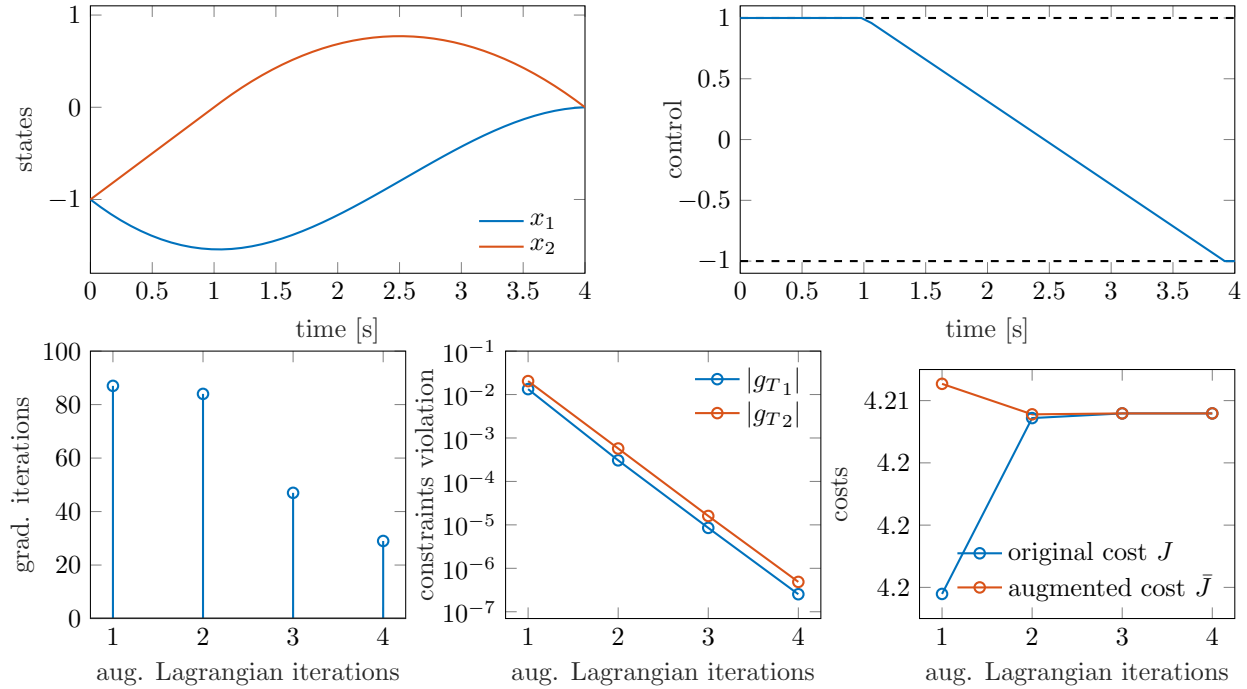


Fig. 7.2.1: Numerical OCP solution of the double integrator problem with fixed end time.

7.2.1 Problem formulation

The double integrator problems reads as

$$\begin{aligned}
 \min_{u, T} \quad & J(u, T; \mathbf{x}_0) = T + \int_0^T q_1 u^2(t) dt \\
 \text{s.t.} \quad & \dot{x}_1(t) = x_2(t), \quad x_1(0) = x_{1,0} \\
 & \dot{x}_2(t) = u(t), \quad x_2(0) = x_{2,0} \\
 & \mathbf{g}_T(\mathbf{x}(T)) = [x_1(T), x_2(T)]^\top = \mathbf{0} \\
 & h(\mathbf{x}(t)) = x_2(t) - 0.5 \leq 0 \\
 & u(t) \in [u_{\min}, u_{\max}], \quad T \in [T_{\min}, T_{\max}]
 \end{aligned} \tag{7.2.1}$$

including two terminal equality constraints $\mathbf{g}_T(\mathbf{x}(T))$ and one general inequality constraint $h(\mathbf{x}(t))$. The control task consists in a setpoint transition from the initial state $x_{1,0} = x_{2,0} = -1$ to the origin $x_1(T) = x_2(T) = 0$. The cost functional $J(u, T; \mathbf{x}_0)$ represents a trade-off between time and energy optimality depending on the weight $q_1 = 0.1$.

The problem is formulated in GRAMPC using the C file `probfc_t_DOUBLE_INTEGRATOR`, which can be found in `<grampc_root>/examples/Double_Integrator`. In particular, the terminal equality constraints are formulated in GRAMPC via the functions `gTfct`, `dgTdx_vec` and `dgTdT_vec`. The number of terminal equality constraints is set to `NgT=2` in the function `ocp_dim`. Similarly, the inequality constraint is formulated by means of the functions `hfct`, `dhdxd_vec` and `dhdu_vec` and setting to `Nh=1` in `ocp_dim`. More details on implementing the OCP can be found in [Problem implementation](#) and in the example provided in the folder `<grampc_root>/examples/Double_Integrator`.

The options `OptimControl` and `OptimTime` are activated to optimize not only the control variable u but also the end time T . The lower and upper bounds of the optimization variables are set to $u \in [-1, 1]$ and $T \in [1, 10]$ using the parameters `umin`, `umax`, `Tmin` and `Tmax`, cf. [Problem formulation and implementation](#). Note that the option `ShiftControl` is deactivated, as the shifting of the control trajectory typically only applies to MPC problems.

The option `ConvergenceCheck` is activated to terminate the augmented Lagrangian algorithm as soon as the state constraints are fulfilled with sufficient accuracy and the optimization variable has converged to an optimal value. To this end, the convergence criteria (5.5.2) and (5.5.3) are evaluated after each gradient and augmented Lagrangian step using the thresholds $\varepsilon_{g_T} = [1e-6, 1e-6]^T$ and $\varepsilon_h = 1e-6$, cf. the option `ConstraintsAbsTol`. The threshold for checking convergence of the optimization variable $\varepsilon_{rel,c}$ is set to $1e-9$ via the option `ConvergenceGradientRelTol`. Note that the activation of the convergence check using the option `ConvergenceCheck` causes the gradient method to be aborted when the convergence condition $\eta^{i|j+1} \leq \varepsilon_{rel,c}$ defined by (5.5.3) is reached.

7.2.2 Optimization with fixed end time

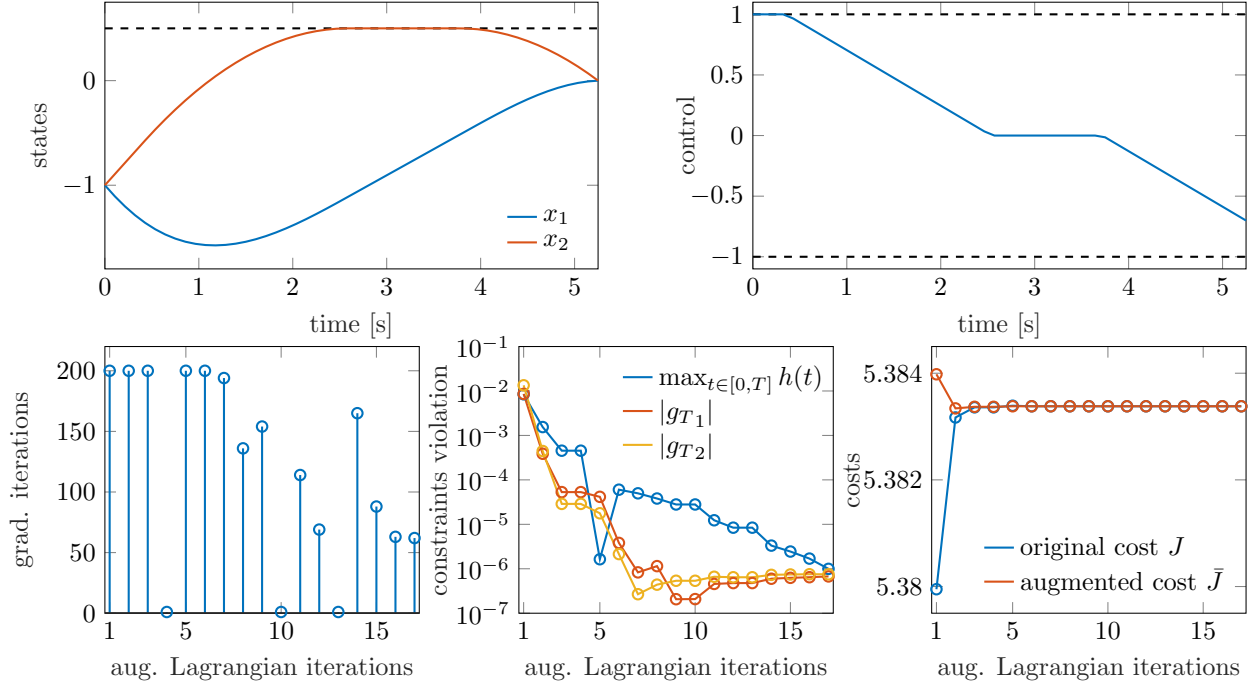


Fig. 7.2.2: Numerical OCP solution of the double integrator problem with fixed end time and state constraint.

In a first scenario, the OCP (7.2.1) is numerically solved with the fixed end time $T = 4$ s, i.e., the option `OptimTime` is set `off`, and without the inequality constraint, i.e., the option `InequalityConstraints` is also set `off`. The time interval $[0, T]$ is discretized using `Nhor=50` discretization points. During the augmented Lagrangian iterations, a decrease of the penalty parameters is prevented by setting the adaptation factor $\beta_{de} = 1$ (see option `PenaltyDecreaseFactor`). The increase factor β_{in} of the penalty parameter update (5.4.2) is set to the value 1.25. These settings ensure a fast convergence, since the very low tolerances ε_{g_T} require high penalty parameters. However, starting with high penalties can lead to instabilities, as high penalties distort the optimization problem.

As shown in Fig. 7.2.1, the state variables $x(t)$ are transferred to the origin as specified by the terminal state constraints. Note that only 5 augmented Lagrangian steps are required in total for numerically solving the state constrained optimization problem in accordance with the formulated convergence criterion for the terminal equality constraints and the optimization variable u . The number of gradient iterations varies in each augmented Lagrangian step as shown in Fig. 7.2.1. The violation of the formulated terminal equality constraints continuously decreases below the specified thresholds $\varepsilon_{g_T} = [1e-6, 1e-6]^T$. As a result, the augmented cost functional and the original cost functional converge to the same value, i.e. the so-called duality gap is zero.

In a second scenario, Fig. 7.2.2 shows the optimal solution of OCP (7.2.1) with activated inequality constraint using the fixed end time $T = 5.25$ s. Further problem settings are identical to the first simulation scenario. Again, the terminal equality constraints are satisfied by the optimal solution and the state variables $x(t)$ are transferred to the origin. The

control variable u is slightly adapted compared to the first simulation scenario in Fig. 7.2.1 in order to comply with the inequality constraint. In view of the additional inequality constraint, 17 augmented Lagrangian steps are required to be able to solve the optimization problem with sufficient accuracy.

The number of gradient iterations varies in each augmented Lagrangian step as shown in Fig. 7.2.2. The violation of the state constraints is almost continuously decreased below the specified thresholds $\varepsilon_{g_T} = [1e-6, 1e-6]^T$ and $\varepsilon_h = 1e-6$, respectively. As before, the augmented cost functional and the original cost functional converge to the same value. The computation time for solving the problem on a Windows 10 machine with an Intel(R) Core(TM) i5-5300U CPU running at 2.3GHz using the Microsoft Visual C++ 2013 Professional (C) compiler amounts to 1.1ms and 14.6ms, respectively.

7.2.3 Optimization with free end time

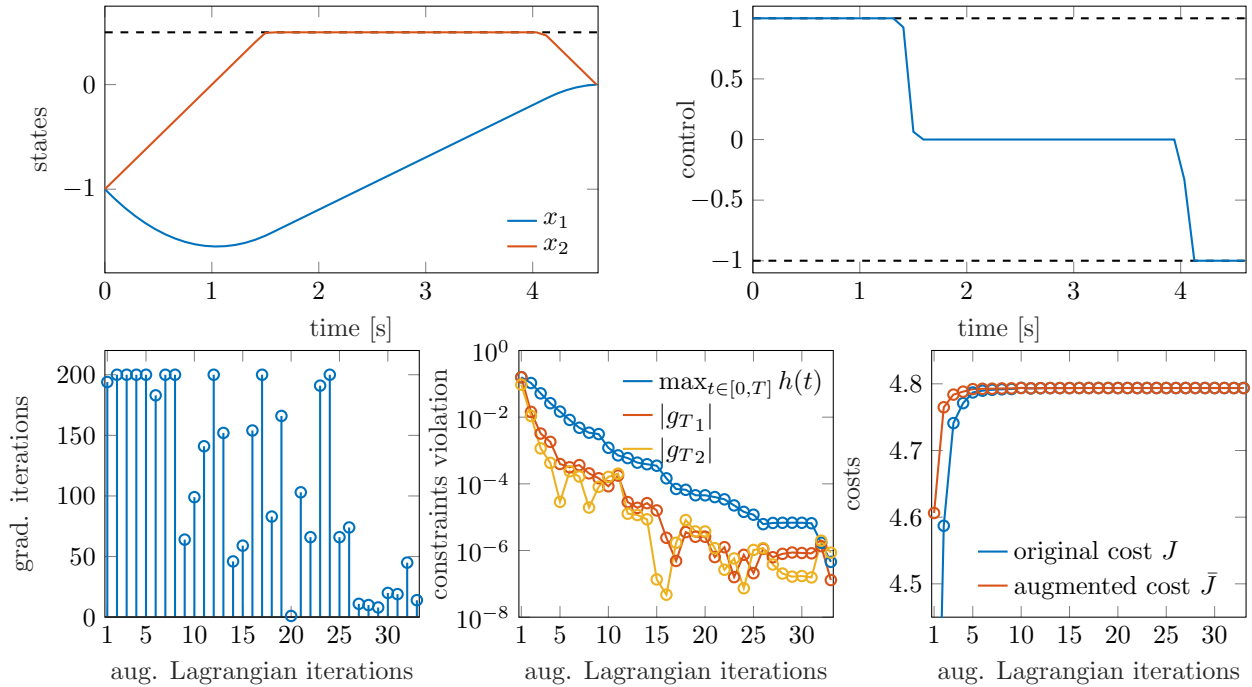


Fig. 7.2.3: Numerical OCP solution of the double integrator problem with free end time and state constraint.

In a third simulation scenario, the OCP (7.2.1) is numerically solved in the free end time setting. The initial end time is set to $T = 5.25$ s as given before. To weight the update of the end time T against the control update when updating the optimization variables according to (5.3.1), the adaptation factor γ_T is set using the option `OptimTimeLineSearchFactor`. The value $\gamma_T = 1.75$ is used in the scenario. Note, however, that values $\gamma_T < 1$ typically increase the algorithmic stability at the expense of the calculation time and vice versa.

The numerical results for the free end time case are shown in Fig. 7.2.3. In contrast to the first two simulation scenarios, the reduction of the end time below 4.6 s allows one to carry out the setpoint transition with a significantly more aggressive control trajectory u . The free end time optimization is a more challenging problem than before and is accompanied by a higher computational effort. This can be observed both in the larger number of augmented Lagrangian steps and gradient steps as well as in terms of the slower improvement of the violation of the state constraints.

Nevertheless, the state constraints are fulfilled at the last augmented Lagrangian step in accordance with the thresholds $\varepsilon_{g_T} = [1e-6, 1e-6]^T$ and $\varepsilon_h = 1e-6$. The improvement of the control performance when optimizing the end time compared to a fixed end time can be specified by the lower value of the cost functional, cf. Fig. 7.2.2 and Fig. 7.2.3. However, this results in a slightly increased computation time of 21.96ms compared to 14.66ms with a fixed end time on the same Windows 10 machine.

7.3 Moving horizon estimation of a CSTR

Continuous stirred tank reactors (CSTR) are a popular class of systems when it comes to the implementation of advanced nonlinear control methods. In this subsection, a CSTR model is used for an example implementation of a moving horizon estimation (MHE) used in closed loop with MPC. Corresponding m and C files can be found in the folder `<grampc_root>/examples/Reactor_CSTR`.

7.3.1 Problem formulation

The system dynamic are given by¹

$$\begin{aligned}\dot{c}_A &= -k_1(T)c_A - k_2(T)c_A^2 + (c_{in} - c_A)u_1 \\ \dot{c}_B &= k_1(T)c_A - k_1(T)c_B - c_B u_1 \\ \dot{T} &= -\delta(k_1(T)c_A\Delta H_{AB} + k_1(T)\Delta H_{BC} + k_2(T)c_A^2\Delta H_{AD}) \\ &\quad + \alpha(T_C - T) + (T_{in} - T)u_1 \\ \dot{T}_C &= \beta(T - T_C) + \gamma u_2,\end{aligned}$$

where the monomer and product concentrations c_A and c_B , respectively, as well as the reactor and cooling temperature T and T_C form the state vector $\mathbf{x} = [c_A, c_B, T, T_C]$. The two functions $k_1(T)$ and $k_2(T)$ are of Arrhenius type

$$k_i(T) = k_{i0} \exp\left(\frac{-E_i}{T/^\circ\text{C} + 273.15}\right), \quad i = 1, 2.$$

The measured quantities are the two temperatures $\mathbf{y} = [T, T_C]^\top$. The controls $\mathbf{u} = [u_1, u_2]$, i.e. the normalized flow rate and the cooling power, are assumed to be known as well. Table 7.3.1 gives the parameters of the system that are passed to the GRAMPC problem functions via `userparam`. A more detailed description can be found in¹.

Table 7.3.1: Parameters of the CSTR model¹.

Parameter	Value	Unit	Parameter	Value	Unit
α	30.828	h^{-1}	k_{20}	9.043×10^6	$\text{m}^3 \text{mol}^{-1} \text{h}^{-1}$
β	86.688	h^{-1}	E_1	9785.3	
δ	3.522×10^{-4}	K kJ^{-1}	E_2	8560.0	
γ	0.1	kh^{-1}	ΔH_{AB}	4.2	kJ mol^{-1}
T_{in}	104.9	$^\circ\text{C}$	ΔH_{BC}	-11.0	kJ mol^{-1}
c_{in}	5.1×10^3	mol m^{-3}	ΔH_{AD}	-41.85	kJ mol^{-1}
k_{10}	1.287×10^{12}	h^{-1}			

The control task at hand is the setpoint change between the two stationary points

$$\mathbf{x}_{\text{des},1} = [1370 \frac{\text{kmol}}{\text{m}^3}, 950 \frac{\text{kmol}}{\text{m}^3}, 110.0^\circ\text{C}, 108.6^\circ\text{C}]^\top, \mathbf{u}_{\text{des},1} = [5.0 \text{ h}^{-1}, -1190 \text{ kJ h}^{-1}]^\top$$

and

$$\mathbf{x}_{\text{des},2} = [2020 \frac{\text{kmol}}{\text{m}^3}, 1070 \frac{\text{kmol}}{\text{m}^3}, 100.0^\circ\text{C}, 97.1^\circ\text{C}]^\top, \mathbf{u}_{\text{des},2} = [5.0 \text{ h}^{-1}, -2540 \text{ kJ h}^{-1}]^\top.$$

The cost functional is designed quadratically

$$J(\mathbf{u}, \mathbf{x}_k) := \Delta \mathbf{x}(T)^\top \mathbf{P} \Delta \mathbf{x}(T) + \int_0^T \Delta \mathbf{x}(t)^\top \mathbf{Q} \Delta \mathbf{x}(t) \Delta \mathbf{u}(t)^\top \mathbf{R} \Delta \mathbf{u}(t)$$

¹ R. Rothfuss, J. Rudolph, and M. Zeitz. Flatness based control of a nonlinear chemical reactor model. *Automatica*, 32(10):1433–1439, 1996.

in order to penalize the deviation of the state and control from the desired setpoint $(\mathbf{x}_{\text{des},1}, \mathbf{u}_{\text{des},1})$ with $\Delta \mathbf{x} = \mathbf{x} - \mathbf{x}_{\text{des}}$ and $\Delta \mathbf{u} = \mathbf{u} - \mathbf{u}_{\text{des}}$. The weight matrices are chosen as

$$\mathbf{P} = \text{diag}(0.2, 1.0, 0.5, 0.2), \quad \mathbf{R} = \text{diag}(0.5, 5.0\text{e} - 3), \quad \mathbf{Q} = \text{diag}(0.2, 1.0, 0.5, 0.2).$$

The control task will be tackled by MPC using GRAMPC. In addition, an MHE using GRAMPC is designed to estimate the current state $\hat{\mathbf{x}}_k$ w.r.t. the measured temperatures $\mathbf{y} = [T, T_C]^\top$.

In analogy to the MPC formulation (4.1.1), moving horizon estimation is typically based on the online solution of a dynamic optimization problem

$$\begin{aligned} \min_{\hat{\mathbf{x}}_k} \quad & J(\hat{\mathbf{x}}_k; \mathbf{u}, \mathbf{y}) = \int_{t_k-T}^{t_k} \|\hat{\mathbf{y}}(t) - \mathbf{y}(t)\|^2 dt \\ \text{s.t.} \quad & \mathbf{M}\dot{\hat{\mathbf{x}}}(t) = \mathbf{f}(\hat{\mathbf{x}}(t), \mathbf{u}(t), t), \quad \hat{\mathbf{x}}(t_k) = \hat{\mathbf{x}}_k \\ & \hat{\mathbf{y}}(t) = \boldsymbol{\sigma}(\hat{\mathbf{x}}(t)) \end{aligned} \tag{7.3.1}$$

that depends on the history of the measured outputs $\mathbf{y}(t)$ and controls $\mathbf{u}(t)$ in the past time window $[t_k - T, t_k]$. The solution of (7.3.1) yields the estimate of the state $\hat{\mathbf{x}}_k$ such that the difference between the measured output $\mathbf{y}(t)$ and the estimated output function $\hat{\mathbf{y}}(t) = \boldsymbol{\sigma}(\hat{\mathbf{x}}(t))$ is minimal in the sense of (7.3.1). GRAMPC solves this MHE problem by means of parameter optimization. To this end, the state at the beginning of the optimization horizon is defined as optimization variable, i.e. $\mathbf{p} = \hat{\mathbf{x}}(t_k - T)$.

Both MHE and MPC use a sampling rate of $\Delta t = 1$ s. A prediction horizon of $T = 20$ min with 40 discretization points is used for the MPC, while a prediction horizon of $T = 10$ s with 10 discretization points is used for the MHE. The MPC implementation uses three gradient iterations per sampling step, i.e. $(i_{\text{max}}, j_{\text{max}}) = (1, 3)$, while the implementation of the MHE uses only a single gradient iteration, i.e. $(i_{\text{max}}, j_{\text{max}}) = (1, 1)$. Note that because the MHE and MPC problems are defined without state constraints, the outer augmented Lagrangian loop causes no computational overhead, as GRAMPC skips the multiplier and penalty update. As the implementation of the MHE is not quite as straightforward as the MPC case, the next subsection describes the implementation process in more detail.

7.3.2 Implementation aspects

The following lines describe the implementation of the MHE problem with GRAMPC, the corresponding simulation results are shown in the next subsection. In a first step, the MHE problem (7.3.1) has to be transformed in a more suitable representation that can be tackled with the parameter optimization functionality of GRAMPC. To this end, a coordinate transformation

$$\tilde{\mathbf{x}}(\tau) = \hat{\mathbf{x}}(t_k - T + \tau) - \mathbf{p}, \quad \tilde{\mathbf{u}}(\tau) = \mathbf{u}(t_k - T + \tau), \quad \tilde{\mathbf{y}}(\tau) = \mathbf{y}(t_k - T + \tau) \tag{7.3.2}$$

is used together with the corresponding time transformation from $t \in [t_k - T, t_k]^\top$ to the new time coordinate $\tau \in [0, T]$. In combination with the optimization variable $\mathbf{p} = \hat{\mathbf{x}}(t_k - T)$ and the homogeneous initial state $\tilde{\mathbf{x}}(0) = \mathbf{0}$, the optimization problem can be cast into the form

$$\begin{aligned} \min_{\mathbf{p}} \quad & J(\mathbf{p}; \tilde{\mathbf{u}}, \tilde{\mathbf{y}}) = \int_0^T \|\hat{\mathbf{y}}(\tau) - \tilde{\mathbf{y}}(\tau)\|^2 d\tau \\ \text{s.t.} \quad & \dot{\tilde{\mathbf{x}}}(\tau) = \mathbf{f}(\tilde{\mathbf{x}}(\tau) + \mathbf{p}, \tilde{\mathbf{u}}(\tau), t_k - T + \tau), \quad \tilde{\mathbf{x}}(0) = \mathbf{0} \\ & \hat{\mathbf{y}}(\tau) = \boldsymbol{\sigma}(\tilde{\mathbf{x}}(\tau) + \mathbf{p}). \end{aligned} \tag{7.3.3}$$

The implementation of this optimization problem still requires to access the measurements $\tilde{\mathbf{y}}$ in the integral cost term. This is achieved by appending the measurements to `userparam` (see `startMHE.m` in `<grampc_root>/examples/Reactor_CSTR`)

```
% init array of last MHE-Nhor measurements of the two temperatures
xMeas_array = repmat(grampcMPC.param.x0(3:4), 1, grampcMHE.opt.Nhor);
grampcMHE.userparam(end-2*grampcMHE.opt.Nhor+1:end) = xMeas_array;
```


The measurements are updated in each iteration of the MPC/MHE loop, e.g.

```
% set values of last MHE-Nhor measurements of the two temperatures
xMeas_temp = xtemp(end,3:4) + randn(1,2)*4; % measurement noise
xMeas_array = [xMeas_array(3:end), xMeas_temp];
grampcMHE.userparam(end-2*grampcMHE.opt.Nhor+1:end) = xMeas_array;
```

When the number of discretization points and the horizon length is known, the measurements can easily be accessed in the problem description file in the following way:

```
typeRNum* pSys = (typeRNum*)userparam;
typeRNum* pCost = &pSys[14];
typeRNum* pMeas = &pSys[20];
typeInt index = (int)floor(t / 2.77777777777778e-04 + 0.00001);
typeRNum meas1 = pMeas[2 * index];
typeRNum meas2 = pMeas[1 + 2 * index];
```

The pointer `pMeas` is set to the 20th element of the `userparam` vector, since the first 14 are the system parameters given in Table 7.3.1 and the next six elements are the weights of the cost function. Also note that $2.778e - 4 \text{ h}^{-1} \approx \Delta t$. Since the order of magnitude of the individual states and controls differs a lot, the scaling option `ScaleProblem` with

$$\begin{aligned} \mathbf{x}_{\text{scale}} &= [500 \frac{\text{kmol}}{\text{m}^3}, 500 \frac{\text{kmol}}{\text{m}^3}, 50^\circ\text{C}, 50^\circ\text{C}]^T & \mathbf{x}_{\text{offset}} &= [500 \frac{\text{kmol}}{\text{m}^3}, 500 \frac{\text{kmol}}{\text{m}^3}, 50^\circ\text{C}, 50^\circ\text{C}]^T \\ \mathbf{p}_{\text{scale}} &= [500 \frac{\text{kmol}}{\text{m}^3}, 500 \frac{\text{kmol}}{\text{m}^3}, 50^\circ\text{C}, 50^\circ\text{C}]^T & \mathbf{p}_{\text{offset}} &= [500 \frac{\text{kmol}}{\text{m}^3}, 500 \frac{\text{kmol}}{\text{m}^3}, 50^\circ\text{C}, 50^\circ\text{C}]^T \\ \mathbf{u}_{\text{scale}} &= [16 \text{ h}^{-1}, 4500 \text{ kJh}^{-1}]^T & \mathbf{u}_{\text{offset}} &= [19 \text{ h}^{-1}, -4500 \text{ kJh}^{-1}]^T \end{aligned}$$

is activated.

7.3.3 Evaluation

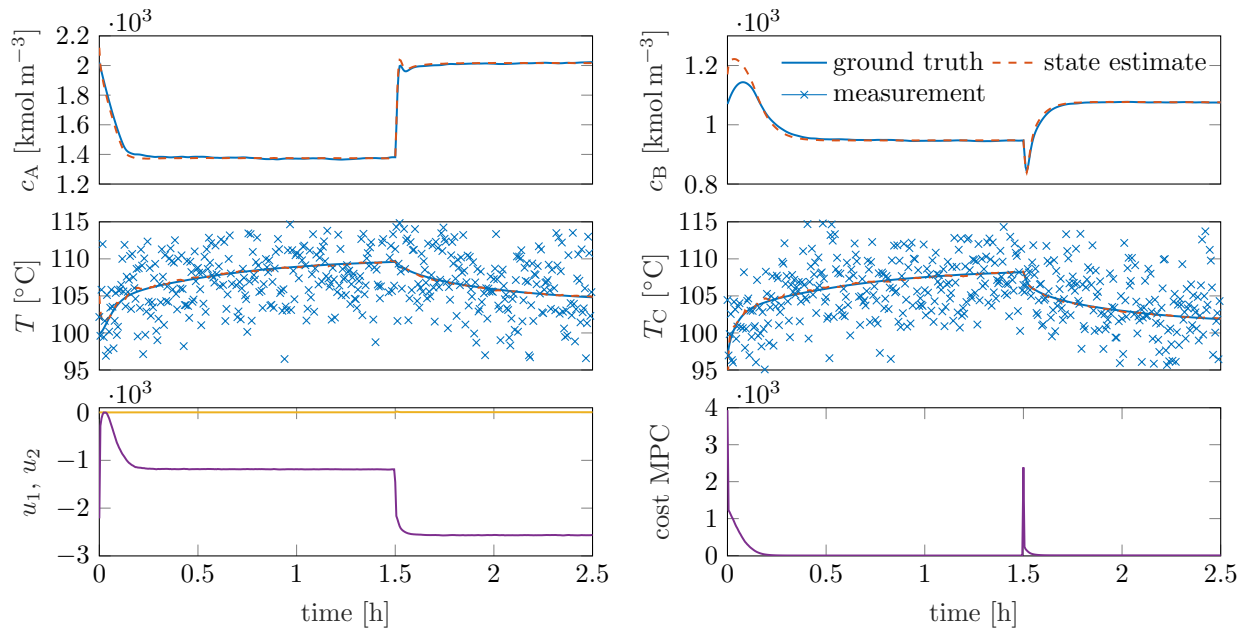


Fig. 7.3.1: Simulated MHE/MPC trajectories for the CSTR reactor example.

The moving horizon estimator is evaluated in conjunction with the MPC. The state estimates are initialized with an initial disturbance $\delta \mathbf{p} = [100 \frac{\text{kmol}}{\text{m}^3}, 100 \frac{\text{kmol}}{\text{m}^3}, 5^\circ\text{C}, 7^\circ\text{C}]^\top$. For a more realistic setting, white Gaussian noise with zero mean and a standard deviation of 4°C is added to the measurements $\mathbf{y} = [T, T_C]^\top$.

Fig. 7.3.1 shows the simulation results from the closed loop simulation of the MHE in conjunction with the MPC. The estimates $\hat{\mathbf{x}}_k$ quickly converge to the actual states (ground truth), as e.g. can be seen in the upper right plot. Furthermore, the cost of the MPC quickly converges to zero after each setpoint change at $t = 0$ h and $t = 1.5$ h, respectively, which shows the good performance of the combined MPC/MHE problem.

The corresponding computation times of GRAMPC amount to $58 \mu\text{s}$ and $11 \mu\text{s}$ per MPC and MHE step, respectively, on a Windows 10 machine with Intel(R) Core(TM) i5-5300U CPU running at 2.3GHz using the Microsoft Visual C++ 2013 Professional (C) compiler.

7.4 Differential algebraic equations

This section first introduces the DAE-system, before implementation aspects regarding the dedicated DAE-solver RODAS are considered. Finally, the example is evaluated.

7.4.1 Problem formulation

The problem at hand is a toy example to illustrate the functionality of GRAMPC with regard to the solution of DAEs. It consists of two differential integrator states and one algebraic state. In addition, this algebraic state is subject to an equality constraint. The corresponding MPC problem is given by

$$\begin{aligned} \min_{\mathbf{u}} \quad & J(\mathbf{x}, \mathbf{u}) = \int_0^T \frac{1}{2} (\Delta \mathbf{x} \mathbf{Q} \Delta \mathbf{x}^\top + \mathbf{u} \mathbf{R} \mathbf{u}^\top) dt \\ \text{s.t.} \quad & \dot{x}_1(t) = u_1(t), \quad x_1(0) = x_{1,0} \\ & \dot{x}_2(t) = u_2(t), \quad x_2(0) = x_{2,0} \\ & 0 = x_1(t) + x_2(t) - x_3(t) \\ & g(\mathbf{x}(t)) = x_3(t) - 1 = 0 \\ & \mathbf{u}(t) \in [\mathbf{u}_{\min}, \mathbf{u}_{\max}], \end{aligned} \tag{7.4.1}$$

where $\Delta \mathbf{x} = \mathbf{x} - \mathbf{x}_{\text{des}}$ and the weight matrices are chosen as $\mathbf{Q} = \text{diag}(500, 0, 0)$ and $\mathbf{R} = \text{diag}(1, 1)$, respectively. The target of the MPC formulation is to steer the first differential state to the desired value, while remaining on the manifold defined by $x_1(t) + x_2(t) = 1$. Note that this equation results from substituting the algebraic equation into the constraint $g(\mathbf{x}(t))$. Even though it would be possible to do this substitution and solve the resulting problem, the purpose of this example is to illustrate the solution of a DAE.

The DAE given in (7.4.1) can be rewritten with a mass matrix \mathbf{M} , i.e.

$$\underbrace{\begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 0 \end{pmatrix}}_{\mathbf{M}} \dot{\mathbf{x}} = \underbrace{\begin{pmatrix} u_1 \\ u_2 \\ x_1 + x_2 - x_3 \end{pmatrix}}_{\mathbf{f}(\mathbf{x}, \mathbf{u})}.$$

Clearly, the mass matrix is different from the identity matrix and singular, i.e. the inverse does not exist and therefore the solver RODAS is used to integrate the system dynamics as well as the corresponding adjoint dynamics.

7.4.2 Implementation aspects

To solve the MPC problem for a DAE, the integrator RODAS has to be used and some additional options have to be set. Furthermore, some additional functions have to be implemented in the `probfcf`-file.

The options are described in *Integration of semi-implicit ODEs and DAEs (RODAS)*. In the example at hand, the right hand side of the system dynamics is not explicitly dependent on the time t and therefore `IFCN` is set to zero. The next option concerns the calculation of $\frac{\partial f}{\partial t}$ and $\frac{\partial^2 H}{\partial x \partial t}$. It can either be set to zero (i.e. `IDFX` = 0) and finite differences are utilized or set to one (i.e. `IDFX` = 1) and the analytical solutions implemented in the functions `dfdt` and `dHdxdt` are called. The third option determines if the numerical (i.e. finite differences) or the analytical solution (i.e. `dfdx` and `dfdxtrans`) is used to compute the Jacobians $\frac{\partial f}{\partial x}$ and $(\frac{\partial f}{\partial x})^T = \frac{\partial^2 H}{\partial x \partial \lambda}$. The next option (`IMAS`) determines if the mass matrix is equal to the identity matrix (i.e. `IMAS` = 0) or if it is specified by the functions `Mfct` and `Mtrans` (i.e. `IMAS` = 1). In the current example, the mass matrix is singular (not the identity matrix) and therefore the option is set to one. The remaining options regard the size of the Jacobian and the mass matrix. The number of non-zero lower and upper diagonals of the Jacobian are given by `MLJAC` and `MUJAC`, respectively. In our case, we have a full matrix and therefore set both options to the system dimension, i.e. N_x . The only non-zero entries of the mass matrix lie on the main diagonal. Thus, the corresponding options (i.e. `MLMAS` and `MUMAS`) are set to zero. Note that one has to be careful, if the Jacobian or mass matrix are sparse, since the lower and upper diagonals are padded with zeros. This is shown for the example matrix in [Fig. 7.4.1](#) with the corresponding code in the following Example.

Listing 7.4.1: Example (C-Code for the mass function illustrated in [Fig. 7.4.1](#))

```
void Mfct(typeNum *out, const typeGRAMPCparam *param, typeUSERPARAM *userparam)
{
    /* row 1 */
    out[0] = 0;
    out[1] = k;
    out[2] = k;
    out[3] = k;
    /* row 2 */
    out[4] = k;
    out[5] = k;
    out[6] = k;
    out[7] = k;
    /* row 3 */
    out[8] = k;
    out[9] = k;
    out[10] = k;
    out[11] = k;
    /* row 4 */
    out[12] = k;
    out[13] = k;
    out[14] = k;
    out[15] = 0;
    /* row 5 */
    out[16] = k;
    out[17] = k;
    out[18] = 0;
    out[19] = 0;
};
```

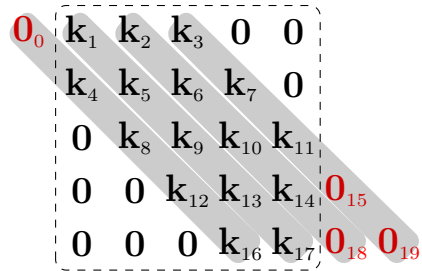


Fig. 7.4.1: Example mass matrix with the index i showing at which position in the output array (i.e. $\text{out}[i]$) the corresponding value has to be written, cf. the example above.

7.4.3 Evaluation

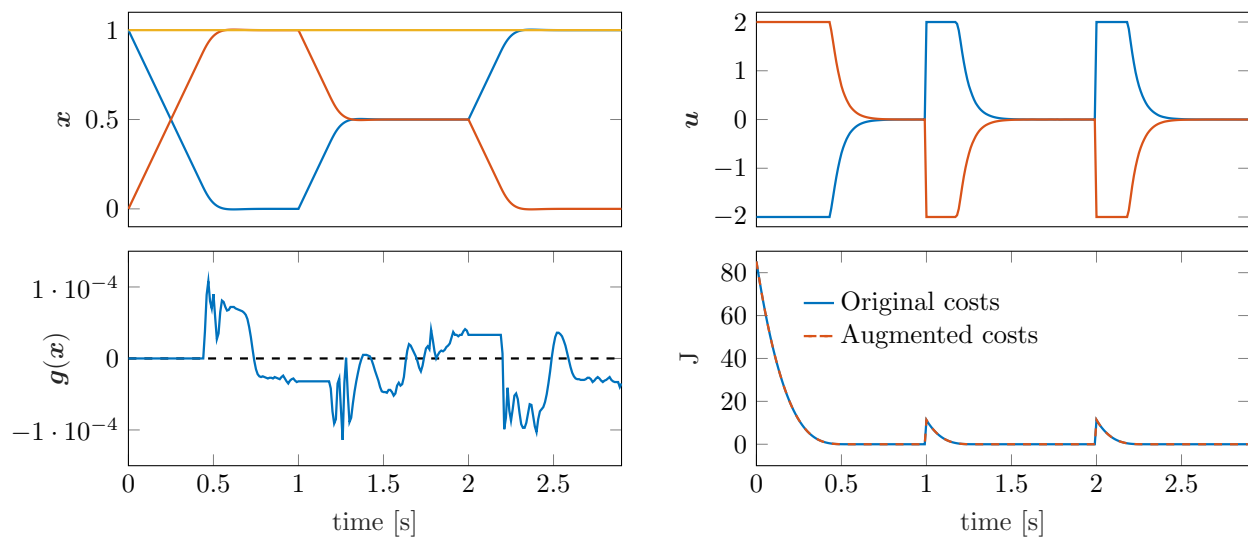


Fig. 7.4.2: Simulated MPC trajectories for the DAE-example

Fig. 7.4.2 shows the simulated trajectories for three set point changes. The setpoint of the first state x_1 changes from 1 to 0 at 0s, from 0 to 0.5 after 1s, and finally from 0.5 to 1 after 2s. Due to the algebraic state and the equality constraint, the trajectory of the second state x_2 has to be the mirror image of x_1 around 0.5, which can be observed in the upper left plot. The corresponding controls in the upper right plot are also mirrored. The constraint violation during the simulation is shown in the lower left plot of Fig. 7.4.2. The allowed constraint violation was set to 1×10^{-4} , which is approximately met. Lastly, the lower right plot shows the original costs and the augmented costs. Both of which quickly converge to zero after each set point change (after 0, 1 and 2s, respectively).

7.5 Constraint Tuning

This section shows how the options of GRAMPC can be adjusted in such a way that the performance in terms of computation time and optimality is improved. To this end, a specific OCP problem is considered. However, the approach can serve as template for different problems.

7.5.1 Problem formulation

The system at hand is a double integrator with one inequality constraint and a terminal equality constraint for each state¹. The OCP problem is then defined as

$$\begin{aligned}
 \min_u \quad & J(\mathbf{x}, u) = \int_0^T r u^2 dt \\
 \text{s.t.} \quad & \dot{x}_1(t) = x_2(t), \quad x_1(0) = x_{1,0} \\
 & \dot{x}_2(t) = u(t), \quad x_2(0) = x_{2,0} \\
 & h(\mathbf{x}(t)) = x_1(t) - 0.1 \leq 0 \\
 & g_{T,1}(\mathbf{x}(T)) = x_1(T) = 0 \\
 & g_{T,2}(\mathbf{x}(T)) = x_2(T) + 1 = 0,
 \end{aligned} \tag{7.5.1}$$

with the weight $r = 0.5$ and the initial state $\mathbf{x}(0) = [0, 1]^\top$. The target of the problem is to steer the double integrator states to the terminal state $\mathbf{x}(T) = [0, -1]^\top$ without violating the inequality constraint $h(\mathbf{x}(t))$. The time in which the set point change should be executed is set to $T = 1$ s.

7.5.2 Tuning approach

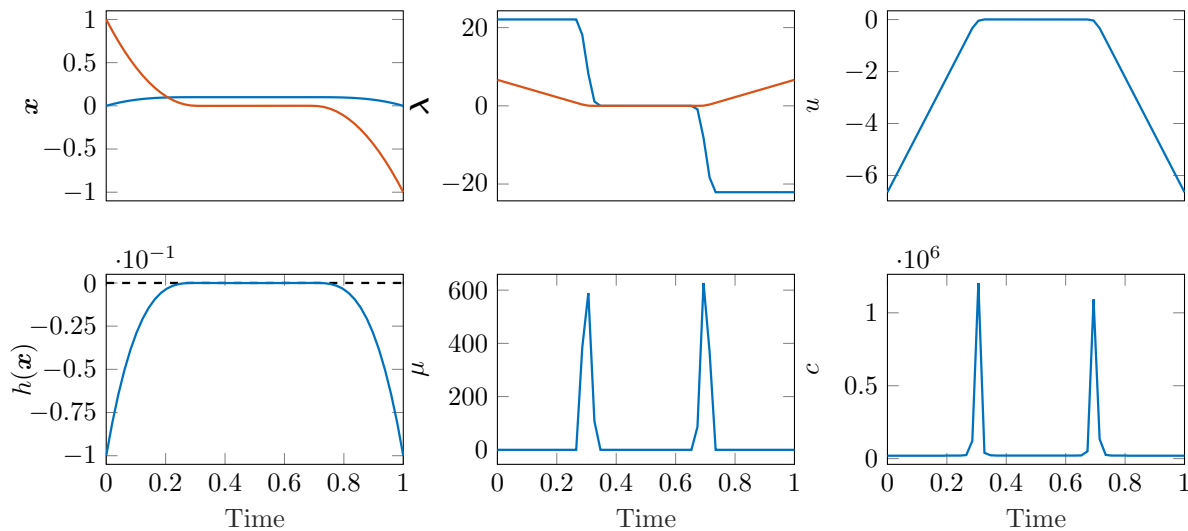


Fig. 7.5.1: The prediction plot of GRAMPC for the double integrator example in (7.5.1).

In the listing below, the step by step approach of tuning the parameters of the augmented Lagrangian algorithm are detailed. The computation time as well as the number of outer and inner iterations are shown in [Table 7.5.1](#) along with the corresponding option that was added or changed in each step.

¹ A. E. Bryson and Y.-C. Ho. *Applied Optimal Control*. Blaisdell, New York, USA, 1969.

Table 7.5.1: Computation time and outer / inner iteration count for the different settings. The last column shows which additional parameter was set different from the initial values in each step.

Step	Time	Multlter	Gradlter (mean)	Additional option
0.	343 ms	3883	10.3	Default settings
1.	129 ms	637	22.4	<code>grampc_estim_penmin</code>
2.	59 ms	171	42.0	<code>PenaltyIncreaseFactor</code> = 1.5
3.	36 ms	98	46.6	<code>PenaltyIncreaseThreshold</code> = 0.75
4.	22 ms	42	61.2	<code>LineSearchInit</code> = $5e-7$
4.	22 ms	42	61.2	<code>PenaltyMin</code> = $2e4$

1. The choice of the initial penalty parameter is crucial for numerical conditioning and therefore convergence. A value that is too high will initially put an unnecessary amount of weight on constraint satisfaction and mostly ignore the optimality. If the value is too low, the opposite will happen, i.e. at first the cost function is decreased at the cost of constraint violation. This will be especially detrimental in MPC applications. If no prior knowledge is available, it is recommended to use the function `grampc_estim_penmin` (or the corresponding Cmx interface `grampc_estim_penmin_Cmx`). For the example at hand, this reduces the computation time by approximately a third. Fig. 7.5.1 shows the simulation results using the estimated value for `PenaltyMin`.
2. While the convergence speed is significantly increased, the penalty parameters at 0.3s and 0.7s are several magnitudes greater than the initial value. Since this huge penalty parameter occurs only at two points during the simulation, it is advisable to set the `PenaltyIncreaseFactor` to a bigger value. This again reduces the computation time by more than half, since fewer increases of the penalty parameter during the outer iterations are necessary. The value should not be chosen too big, as this will have an adverse effect on the numerical conditioning and convergence.
3. In accordance with the previous step, the threshold to increase the penalty parameter, i.e. `PenaltyIncreaseThreshold` is lowered in order to increase the penalty parameter more aggressively. This step almost doubles the convergence speed. Note that this step and the previous step are interchangeable.
4. Another common tuning possibility is the initial step size of the line search, especially if one of the explicit methods is used, cf. [Explicit line search](#). Note that the initial value `LineSearchInit` is used in the case that the explicit formula results in a negative step size. One approach is to use the `LineSearchExpAutoFallback` option. However, problem specific tuning (mostly trial and error) can result in a significant performance boost. In the example at hand, this results in approximately 40% faster convergence speed.
5. To further optimize the parameters, the estimation function for the minimal penalty parameter can be deactivated again and a better value for `PenaltyMin` be used (note that the parameter is increased until there is no further improvement or an decrease in performance). This results in an additional 15% decrease of computation time.

APPENDIX

In addition to a list of all parameters and algorithmic options of GRAMPC, this appendix contains a short description of the structure variable `grampc`. Essential C functions of the GRAMPC project are also listed.

8.1 List of parameters

Table 8.1.1 gives an overview of the problem-specific parameters of GRAMPC in terms of the parameter name and type as well as the admissible range and default values (if available), whereby the symbol $\mathbf{e}_n = [1, \dots, 1]^T \in \mathbb{R}^n$ denotes an n -dimensional column vector.

Table 8.1.1: Problem-specific parameters.

Parameter name	Type	Allowed values	Default
<code>x0</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$0 \cdot \mathbf{e}_{Nx}$
<code>xdes</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$0 \cdot \mathbf{e}_{Nx}$
<code>u0</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$0 \cdot \mathbf{e}_{Nx}$
<code>udes</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$0 \cdot \mathbf{e}_{Nx}$
<code>umax</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$+\infty \cdot \mathbf{e}_{Nx}$
<code>umin</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$-\infty \cdot \mathbf{e}_{Nx}$
<code>p0</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$0 \cdot \mathbf{e}_{Nx}$
<code>pmax</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$+\infty \cdot \mathbf{e}_{Nx}$
<code>pmin</code>	<code>typeNum*</code>	$(-\infty, \infty)$	$-\infty \cdot \mathbf{e}_{Nx}$
<code>Thor</code>	<code>typeNum</code>	$(0, \infty)$	To be provided
<code>Tmax</code>	<code>typeNum</code>	$(0, \infty)$	10^8
<code>Tmin</code>	<code>typeNum</code>	$(0, \infty)$	10^{-8}
<code>dt</code>	<code>typeNum</code>	$(0, \infty)$	To be provided
<code>t0</code>	<code>typeNum</code>	$(0, \infty)$	0
<code>userparam</code> (in C)	<code>void*</code>	User-defined	NULL
<code>userparam</code> (in MATLAB)	<code>typeNum*</code>	$(-\infty, \infty)$	[]

A description of all parameters is as follows (see also *Problem formulation and implementation*):

- `x0`: Initial state vector $\mathbf{x}(t_0) = \mathbf{x}_0$ at the corresponding sampling time t_0 .
- `xdes`: Desired (constant) setpoint vector for the state variables \mathbf{x} .

- **u0**: Initial value of the control vector $\mathbf{u}(t) = \mathbf{u}_0 = \text{const.}, t \in [0, T]$ that is used in the first iteration of GRAMPC.
- **udes**: Desired (constant) setpoint vector for the control variables \mathbf{u} .
- **umin, umax**: Lower and upper bounds for the control variables \mathbf{u} .
- **p0**: Initial value of the parameter vector $\mathbf{p} = \mathbf{p}_0$ that is used in the first iteration of GRAMPC.
- **pmin, pmax**: Lower and upper bounds for the parameters \mathbf{p} .
- **Thor**: Prediction horizon T or initial value if the end time is optimized.
- **Tmin, Tmax**: Lower and upper bound for the prediction horizon T .
- **dt**: Sampling time Δt of the considered system for model predictive control or moving horizon estimation. Required for prediction of next state `grampc.sol.xnext` and for the control shift, see [Control shift](#).
- **t0**: Current sampling instance t_0 that is provided in the `grampc.param` structure.
- **userparam**: Further problem-specific parameters, e.g. system parameters or weights in the cost functions that are passed to the problem functions via a void-pointer in C or `typeRNum` array in MATLAB.

8.2 List of options

Table 8.2.1 gives an overview of the algorithmic options of GRAMPC in terms of the option name and type as well as the allowed and the default values (if available).

Table 8.2.1: Algorithmic Options.

Option name	Type	Allowed values	Default
Nhor	typeInt	$[2, \infty)$	30
MaxGradIter	typeInt	$[1, \infty)$	2
MaxMultIter	typeInt	$[1, \infty)$	1
ShiftControl	typeChar*	on/ off	on
IntegralCost	typeChar*	on/ off	on
TerminalCost	typeChar*	on/ off	on
IntegratorCost	typeChar*	trapezoidal/ simpson/ discrete	trapezoidal
Integrator	typeChar*	erk1/ erk2/ erk3/ erk4/ discrete/ ruku45/ rodas	erk2
IntegratorRelTol	typeRNum	$(0, \infty)$	10^{-6}
IntegratorAbsTol	typeRNum	$(0, \infty)$	10^{-8}
IntegratorMinStepSize	typeRNum	$(0, \infty)$	<i>eps</i>
IntegratorMaxSteps	typeInt	$[1, \infty)$	10^8
FlagsRodas	typeInt*	see description	see description
LineSearchType	typeChar*	adaptive/ explicit1/ explicit2	explicit2
LineSearchExpAutoFallback	typeChar*	on/ off	on
LineSearchMax	typeRNum	$(0, \infty)$	0.75
LineSearchMin	typeRNum	$(0, \infty)$	10^{-10}
LineSearchInit	typeRNum	$(0, \infty)$	10^{-4}
LineSearchAdaptAbsTol	typeRNum	$[0, \infty)$	10^{-6}
LineSearchAdaptFactor	typeRNum	$(1, \infty)$	3/2
LineSearchIntervalTol	typeRNum	$(0, 0.5)$	0.1
LineSearchIntervalFactor	typeRNum	$(0, 1)$	0.85

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Table 8.2.1 – continued from previous page

Option name	Type	Allowed values	Default
OptimControl	typeChar*	on/ off	on
OptimParam	typeChar*	on/ off	off
OptimParamLineSearchFactor	typeRNum	$(0, \infty)$	1.0
OptimTime	typeChar*	on/ off	off
OptimTimeLineSearchFactor	typeRNum	$(0, \infty)$	1.0
ScaleProblem	typeChar*	on/ off	off
xScale	typeRNum*	$(-\infty, \infty)$	e_{Nx}
xOffset	typeRNum*	$(-\infty, \infty)$	$0 \cdot e_{Nx}$
uScale	typeRNum*	$(-\infty, \infty)$	e_{Nu}
uOffset	typeRNum*	$(-\infty, \infty)$	$0 \cdot e_{Nu}$
pScale	typeRNum*	$(-\infty, \infty)$	e_{Np}
pOffset	typeRNum*	$(-\infty, \infty)$	$0 \cdot e_{Np}$
TScale	typeRNum	$(0, \infty)$	1.0
TOffset	typeRNum	$(-\infty, \infty)$	0.0
JScale	typeRNum	$(0, \infty)$	1.0
cScale	typeRNum*	$(-\infty, \infty)$	e_{Nc}
EqualityConstraints	typeChar*	on/ off	on
InequalityConstraints	typeChar*	on/ off	on
TerminalEqualityConstraints	typeChar*	on/ off	on
TerminalInequalityConstraints	typeChar*	on/ off	on
ConstraintsHandling	typeChar*	extpen/ auglag	auglag
ConstraintsAbsTol	typeRNum*	$(0, \infty)$	$10^{-4}e_{Nc}$
MultiplierMax	typeRNum	$(0, \infty)$	10^6
MultiplierDampingFactor	typeRNum	$[0, 1]$	0.0
PenaltyMax	typeRNum	$(0, \infty)$	10^6
PenaltyMin	typeRNum	$(0, \infty)$	10^0
PenaltyIncreaseFactor	typeRNum	$[1, \infty)$	1.05
PenaltyDecreaseFactor	typeRNum	$[0, 1]$	0.95
PenaltyIncreaseThreshold	typeRNum	$[0, \infty)$	1.0
AugLagUpdateGradientRelTol	typeRNum	$[0, 1]$	10^{-2}
ConvergenceCheck	typeChar*	on/ off	off
ConvergenceGradientRelTol	typeRNum	$[0, 1]$	10^{-6}

A description of all options is as follows:

- **Nhor**: Number of discretization points within the time interval $[0, T]$.
- **MaxMultIter**: Sets the maximum number of augmented Lagrangian iterations $i_{\max} \geq 1$. If the option **ConvergenceCheck** is activated, the algorithm evaluates the convergence criterion and terminates if the inner minimization converged and all constraints are satisfied within the tolerance defined by **ConstraintsAbsTol**.
- **MaxGradIter**: If the option **ConvergenceCheck** is activated, the algorithm terminates the inner loop as soon as the convergence criterion is fulfilled.
- **ShiftControl**: Activates or deactivates the shifting of the control trajectory and the adaptation of T in case of a free end time, i.e., if **OptimTime** is active.
- **IntegralCost**, **TerminalCost**: Indicate if the integral and/or terminal cost functions are defined.
- **IntegratorCost**: This option specifies the integration scheme for the cost functionals. Possible values are trapezoidal, simpson and discrete.

Changed in version v2.3: Fixed typo in trapezoidal option. Renamed euler to erk1 and heun to erk2. Removed modeuler. Added erk3 and erk4.

- **Integrator**: This option specifies the integration scheme for the system and adjoint dynamics. Possible values are erk1, erk2, erk3, erk4, discrete with fixed step size and ruku45, rodas with variable step size.
- **IntegratorMinStepSize**: Minimum step size for RODAS and the Runge-Kutta integrator.
- **IntegratorMaxSteps**: Maximum number of steps for RODAS and the Runge-Kutta integrator.
- **IntegratorRelTol**: Relative tolerance for RODAS and the Runge-Kutta integrator with variable step size. Note that this option may be insignificant if the minimum step size is chosen too high or the maximum number of steps is set too low.
- **IntegratorAbsTol**: Absolute tolerance for RODAS and the Runge-Kutta integrator with variable step size. Note that this option may be insignificant if the minimum step size is chosen too high or the maximum number of steps is set too low.
- **FlagsRodas**: Vector with the elements [IFCN, IDFX, IJAC, IMAS, MLJAC, MUJAC, MLMAS, MUMAS] that is passed to the integrator RODAS, see [Integration of semi-implicit ODEs and DAEs \(RODAS\)](#) for a description of the single entries.
- **LineSearchType**: This option selects either the adaptive line search strategy (value adaptive) or the explicit approach (value explicit1 or explicit2).
- **LineSearchExpAutoFallback**: If this option is activated, the automatic fallback strategy is used in the case that the explicit formulas result in negative step sizes.
- **LineSearchMax**: This option sets the maximum value α_{\max} of the step size α .
- **LineSearchMin**: This option sets the minimum value α_{\min} of the step size α .
- **LineSearchInit**: Indicates the initial value $\alpha_{\text{init}} > 0$ for the step size α . If the adaptive line search is used, the sample point α_2 is set to $\alpha_2 = \alpha_{\text{init}}$.
- **LineSearchAdaptAbsTol**: This option sets the absolute tolerance ε_ϕ of the difference in costs at the interval bounds α_1 and α_2 . If the difference in the (scaled) costs on these bounds falls below ε_ϕ , the adaption of the interval is stopped in order to avoid oscillations.
- **LineSearchAdaptFactor**: This option sets the adaptation factor $\kappa > 1$ in (5.3.3) that determines how much the line search interval can be adapted from one gradient iteration to the next.
- **LineSearchIntervalTol**: This option sets the interval tolerance $\varepsilon_\alpha \in (0, 0.5)$ in (5.3.3) that determines for which values of α the adaption is performed.
- **LineSearchIntervalFactor**: This option sets the interval factor $\beta \in (0, 1)$ that specifies the interval bounds $[\alpha_1, \alpha_3]$ according to $\alpha_1 = \alpha_2(1 - \beta)$ and $\alpha_3 = \alpha_2(1 + \beta)$, whereby the mid sample point is initialized as $\alpha_2 = \alpha_{\text{init}}$.
- **OptimControl**: Specifies whether the cost functional should be minimized with respect to the control variable \mathbf{u} .
- **OptimParam**: Specifies whether the cost functional should be minimized with respect to the optimization parameters \mathbf{p} .
- **OptimParamLineSearchFactor**: This option sets the adaptation factor $\gamma_{\mathbf{p}}$ that weights the update of the parameter vector \mathbf{p} against the update of the control \mathbf{u} .
- **OptimTime**: Specifies whether the cost functional should be minimized with respect the horizon length T (free end time problem) or if T is kept constant.
- **OptimTimeLineSearchFactor**: This option sets the adaptation factor γ_T that weights the update of the end time T against the update of the control \mathbf{u} .

- **ScaleProblem**: Activates or deactivates scaling. Note that GRAMPC requires more computation time if scaling is active.
- **xScale, xOffset**: Scaling factors $\mathbf{x}_{\text{scale}}$ and offsets $\mathbf{x}_{\text{offset}}$ for each state variable.
- **uScale, uOffset**: Scaling factors $\mathbf{u}_{\text{scale}}$ and offsets $\mathbf{u}_{\text{offset}}$ for each control variable.
- **pScale, pOffset**: Scaling factors $\mathbf{p}_{\text{scale}}$ and offsets $\mathbf{p}_{\text{offset}}$ for each parameter.
- **TScale, TOffset**: Scaling factor T_{scale} and offset T_{offset} for the horizon length.
- **JScale**: Scaling factor J_{scale} for the cost functional.
- **cScale**: Scaling factors $\mathbf{c}_{\text{scale}}$ for each state constraint. The elements of the vector refer to the equality, inequality, terminal equality and terminal inequality constraints.
- **EqualityConstraints**: Equality constraints $\mathbf{g}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) = \mathbf{0}$ can be disabled by the option value `off`.
- **InequalityConstraints**: To disable inequality constraints $\mathbf{h}(\mathbf{x}(t), \mathbf{u}(t), \mathbf{p}, t) \leq \mathbf{0}$, set this option to `off`.
- **TerminalEqualityConstraints**: To disable terminal equality constraints $\mathbf{g}_T(\mathbf{x}(T), \mathbf{p}, T) = \mathbf{0}$, set this option to `off`.
- **TerminalInequalityConstraints**: To disable terminal inequality constraints $\mathbf{h}_T(\mathbf{x}(T), \mathbf{p}, T) \leq \mathbf{0}$, set this option to `off`.
- **ConstraintsHandling**: State constraints are handled either by means of the augmented Lagrangian approach (option value `auglag`) or as soft constraints by outer penalty functions (option value `extpen`).
- **ConstraintsAbsTol**: Thresholds $(\varepsilon_g, \varepsilon_h, \varepsilon_{g_T}, \varepsilon_{h_T}) \in \mathbb{R}^{N_c}$ for the equality, inequality, terminal equality, and terminal inequality constraints.
- **MultiplierMax**: Upper bound μ_{max} and lower bound $-\mu_{\text{max}}$ for the Lagrangian multipliers.
- **MultiplierDampingFactor**: Damping factor $\rho \in [0, 1)$ for the multiplier update.
- **PenaltyMax**: This option sets the upper bound c_{max} of the penalty parameters.
- **PenaltyMin**: This option sets the lower bound c_{min} of the penalty parameters.
- **PenaltyIncreaseFactor**: This option sets the factor β_{in} by which penalties are increased.
- **PenaltyDecreaseFactor**: This option sets the factor β_{de} by which penalties are decreased.
- **PenaltyIncreaseThreshold**: This option sets the factor γ_{in} that rates the progress in the constraints between the last two iterates.
- **AugLagUpdateGradientRelTol**: Threshold $\varepsilon_{\text{rel,u}}$ for the maximum relative gradient of the inner minimization problem.
- **ConvergenceCheck**: This option activates the convergence criterion. Otherwise, the inner and outer loops always perform the maximum number of iterations, see the options `MaxGradIter` and `MaxMultIter`.
- **ConvergenceGradientRelTol**: This option sets the threshold $\varepsilon_{\text{rel,c}}$ for the maximum relative gradient of the inner minimization problem that is used in the convergence criterion. Note that this threshold is different from the one that is used in the update of multipliers and penalties.

8.3 GRAMPC data types

The following lines list the data types of GRAMPC. Note that these data types define the structure variable `grampc` including the substructures `sol`, `param`, `opt`, `rws`, and `userparam`, also see *Initialization of GRAMPC*.

Listing 8.3.1: GRAMPC main structure.

```
typedef struct
{
    typeGRAMPCparam *param;
    typeGRAMPCopt *opt;
    typeGRAMPCsol *sol;
    typeGRAMPCrws *rws;
    typeUSERPARAM *userparam;
} typeGRAMPC;
```

Listing 8.3.2: GRAMPC parameter structure.

```
typedef struct
{
    typeInt Nx;
    typeInt Nu;
    typeInt Np;
    typeInt Ng;
    typeInt Nh;
    typeInt NgT;
    typeInt NhT;
    typeInt Nc;

    typeRNum *x0;
    typeRNum *xdes;

    typeRNum *u0;
    typeRNum *udes;
    typeRNum *umax;
    typeRNum *umin;

    typeRNum *p0;
    typeRNum *pmax;
    typeRNum *pmin;

    typeRNum Thor;
    typeRNum Tmax;
    typeRNum Tmin;

    typeRNum dt;
    typeRNum t0;
} typeGRAMPCparam;
```

Listing 8.3.3: GRAMPC option structure.

```
typedef struct
{
```

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```

typeInt Nhor;
typeInt MaxGradIter;
typeInt MaxMultIter;
typeInt ShiftControl;

typeInt TimeDiscretization;

typeInt IntegralCost;
typeInt TerminalCost;
typeInt IntegratorCost;

typeInt Integrator;
typeRNum IntegratorRelTol;
typeRNum IntegratorAbsTol;
typeRNum IntegratorMinStepSize;
typeInt IntegratorMaxSteps;
typeInt *FlagsRodas;

typeInt LineSearchType;
typeInt LineSearchExpAutoFallback;
typeRNum LineSearchMax;
typeRNum LineSearchMin;
typeRNum LineSearchInit;
typeRNum LineSearchIntervalFactor;
typeRNum LineSearchAdaptFactor;
typeRNum LineSearchIntervalTol;

typeInt OptimControl;
typeInt OptimParam;
typeRNum OptimParamLineSearchFactor;
typeInt OptimTime;
typeRNum OptimTimeLineSearchFactor;

typeInt ScaleProblem;
typeRNum *xScale;
typeRNum *xOffset;
typeRNum *uScale;
typeRNum *uOffset;
typeRNum *pScale;
typeRNum *pOffset;
typeRNum TScale;
typeRNum TOffset;
typeRNum JScale;
typeRNum *cScale;

typeInt EqualityConstraints;
typeInt InequalityConstraints;
typeInt TerminalEqualityConstraints;
typeInt TerminalInequalityConstraints;
typeInt ConstraintsHandling;
typeRNum *ConstraintsAbsTol;

```

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```
typeNum MultiplierMax;
typeNum MultiplierDampingFactor;
typeNum PenaltyMax;
typeNum PenaltyMin;
typeNum PenaltyIncreaseFactor;
typeNum PenaltyDecreaseFactor;
typeNum PenaltyIncreaseThreshold;
typeNum AugLagUpdateGradientRelTol;

typeInt ConvergenceCheck;
typeNum ConvergenceGradientRelTol;

} typeGRAMPCopt;
```

Listing 8.3.4: GRAMPC solution structure.

```
typedef struct
{
    typeNum *xnext;
    typeNum *unext;
    typeNum *pnext;
    typeNum Tnext;
    typeNum *J;
    typeNum cfct;
    typeNum pen;
    typeInt *iter;
    typeInt status;
} typeGRAMPCsol;
```

Listing 8.3.5: GRAMPC real workspace structure.

```
typedef struct
{
    typeNum *t;
    typeNum *tls;

    typeNum *x;
    typeNum *adj;
    typeNum *dcdx;

    typeNum *u;
    typeNum *uls;
    typeNum *uprev;
    typeNum *gradu;
    typeNum *graduprev;
    typeNum *dcdu;

    typeNum *p;
    typeNum *pls;
    typeNum *pprev;
    typeNum *gradp;
    typeNum *gradpprev;
```

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```

typeNum *dcdp;

typeNum T;
typeNum Tprev;
typeNum gradT;
typeNum gradTprev;
typeNum dcdt;

typeNum *mult;
typeNum *pen;
typeNum *cfct;
typeNum *cfctprev;
typeNum *cfctAbsTol;

typeNum *lsAdapt;
typeNum *lsExplicit;
typeNum *rwsScale;
typeInt  lrwsGeneral;
typeNum *rwsGeneral;

typeInt  lworkRodas;
typeInt  liworkRodas;
typeNum *rparRodas;
typeInt  *iparRodas;
typeNum *workRodas;
typeInt  *iworkRodas;
} typeGRAMPCrws;

```

8.4 GRAMPC function interface

The main C functions for the usage of GRAMPC are listed below.

Listing 8.4.1: File `grampc_init`:

```

void grampc_init(typeGRAMPC **grampc, typeUSERPARAM *userparam);
void grampc_free(typeGRAMPC **grampc);

```

Listing 8.4.2: File `grampc_run`:

```

void grampc_run(const typeGRAMPC *grampc);

```

Listing 8.4.3: File `grampc_setparam`:

```

void grampc_setparam_real(const typeGRAMPC *grampc, const typeChar *paramName, ctypeNum_
↳ paramValue);

void grampc_setparam_real_vector(const typeGRAMPC *grampc, const typeChar *paramName, _
↳ ctypeNum *paramValue);

void grampc_printparam(const typeGRAMPC *grampc);

```

Listing 8.4.4: File grampc_setopt:

```

void grampc_setopt_real(const typeGRAMPC *grampc, const typeChar *optName, ctypeNum
↳optValue);

void grampc_setopt_int(const typeGRAMPC *grampc, const typeChar *optName, ctypeInt
↳optValue);

void grampc_setopt_string(const typeGRAMPC *grampc, const typeChar *optName, const
↳typeChar *optValue);

void grampc_setopt_real_vector(const typeGRAMPC *grampc, const typeChar *optName,
↳ctypeNum *optValue);

void grampc_setopt_int_vector(const typeGRAMPC *grampc, const typeChar *optName,
↳ctypeInt *optValue);

void grampc_printopt(const typeGRAMPC *grampc);

```

Listing 8.4.5: File grampc_mess:

```

/** estimates PenaltyMin on basis of the first MPC iteration */
typeInt grampc_printstatus(ctypeInt status, ctypeInt level);

```

Listing 8.4.6: File grampc_util:

```

/** estimates PenaltyMin on basis of the first MPC iteration */
typeInt grampc_estim_penmin(typeGRAMPC *grampc, ctypeInt rungrampc);

```

Listing 8.4.7: File probfct:

```

/** OCP dimensions: states (Nx), controls (Nu), parameters (Np), equalities (Ng),
    inequalities (Nh), terminal equalities (NgT), terminal inequalities (NhT) */
void ocp_dim(typeInt *Nx, typeInt *Nu, typeInt *Np, typeInt *Ng, typeInt *Nh, typeInt
↳NgT, typeInt *NhT, typeUSERPARAM *userparam);

/** System function f(t,x,u,p,param,userparam)
    ----- */
void ffct(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p, const
↳typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian df/dx multiplied by vector vec, i.e. (df/dx)^T*vec or vec^T*(df/dx) */
void dfdx_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
↳ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian df/du multiplied by vector vec, i.e. (df/du)^T*vec or vec^T*(df/du) */
void dfdu_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
↳ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian df/dp multiplied by vector vec, i.e. (df/dp)^T*vec or vec^T*(df/dp) */
void dfdp_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
↳ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);

```

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```

/** Integral cost  $l(t,x(t),u(t),p,param,userparam)$ 
    ----- */
void lfct(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_
↳ typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Gradient  $dl/dx$  */
void dldx(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_
↳ typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Gradient  $dl/du$  */
void dlldu(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_
↳ typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Gradient  $dl/dp$  */
void dldp(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_
↳ typeGRAMPCparam *param, typeUSERPARAM *userparam);

/** Terminal cost  $V(T,x(T),p,param,userparam)$ 
    ----- */
void Vfct(typeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p, const typeGRAMPCparam_
↳ *param, typeUSERPARAM *userparam);
/** Gradient  $dV/dx$  */
void dVdx(typeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p, const typeGRAMPCparam_
↳ *param, typeUSERPARAM *userparam);
/** Gradient  $dV/dp$  */
void dVdp(typeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p, const typeGRAMPCparam_
↳ *param, typeUSERPARAM *userparam);
/** Gradient  $dV/dT$  */
void dVdT(typeRNum *out, ctypeRNum T, ctypeRNum *x, ctypeRNum *p, const typeGRAMPCparam_
↳ *param, typeUSERPARAM *userparam);

/** Equality constraints  $g(t,x(t),u(t),p,param,userparam) = 0$ 
    ----- */
void gfct(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_
↳ typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian  $dg/dx$  multiplied by vector  $vec$ , i.e.  $(dg/dx)^T * vec$  or  $vec^T * (dg/dx)$  */
void dgdxd_vec(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p,
↳ ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian  $dg/du$  multiplied by vector  $vec$ , i.e.  $(dg/du)^T * vec$  or  $vec^T * (dg/du)$  */
void dgdu_vec(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p,
↳ ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian  $dg/dp$  multiplied by vector  $vec$ , i.e.  $(dg/dp)^T * vec$  or  $vec^T * (dg/dp)$  */
void dgdp_vec(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p,
↳ ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);

/** Inequality constraints  $h(t,x(t),u(t),p,param,userparam) \leq 0$ 
    ----- */
void hfct(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p, const_
↳ typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian  $dh/dx$  multiplied by vector  $vec$ , i.e.  $(dh/dx)^T * vec$  or  $vec^T * (dh/dx)$  */
void dhdx_vec(typeRNum *out, ctypeRNum t, ctypeRNum *x, ctypeRNum *u, ctypeRNum *p,
↳ ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);

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/** Jacobian dh/du multiplied by vector vec, i.e. (dh/du)^T*vec or vec^T*(dg/du) */
void dhdu_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
    ↪ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian dh/dp multiplied by vector vec, i.e. (dh/dp)^T*vec or vec^T*(dg/dp) */
void dhdp_vec(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
    ↪ctypeNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);

/** Terminal equality constraints gT(T,x(T),p,param,userparam) = 0
    ----- */
void gTfct(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, const typeGRAMPCparam
    ↪*param, typeUSERPARAM *userparam);
/** Jacobian dgT/dx multiplied by vector vec, i.e. (dgT/dx)^T*vec or vec^T*(dgT/dx) */
void dgTdx_vec(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, ctypeNum *vec,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian dgT/dp multiplied by vector vec, i.e. (dgT/dp)^T*vec or vec^T*(dgT/dp) */
void dgTdp_vec(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, ctypeNum *vec,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian dgT/dT multiplied by vector vec, i.e. (dgT/dT)^T*vec or vec^T*(dgT/dT) */
void dgTdT_vec(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, ctypeNum *vec,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);

/** Terminal inequality constraints hT(T,x(T),p,param,userparam) <= 0
    ----- */
void hTfct(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, const typeGRAMPCparam
    ↪*param, typeUSERPARAM *userparam);
/** Jacobian dhT/dx multiplied by vector vec, i.e. (dhT/dx)^T*vec or vec^T*(dhT/dx) */
void dhTdx_vec(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, ctypeNum *vec,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian dhT/dp multiplied by vector vec, i.e. (dhT/dp)^T*vec or vec^T*(dhT/dp) */
void dhTdp_vec(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, ctypeNum *vec,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian dhT/dT multiplied by vector vec, i.e. (dhT/dT)^T*vec or vec^T*(dhT/dT) */
void dhTdT_vec(typeNum *out, ctypeNum T, ctypeNum *x, ctypeNum *p, ctypeNum *vec,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);

/** Additional functions required for semi-implicit systems
    M*dx/dt(t) = f(t,x(t),u(t),p,param,userparam) using the solver RODAS
    ----- */
/** Jacobian df/dx in vector form (column-wise) */
void dfdx(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p, const
    ↪typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian df/dx in vector form (column-wise) */
void dfdxtrans(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,
    ↪const typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian df/dt */
void dfdt(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p, const
    ↪typeGRAMPCparam *param, typeUSERPARAM *userparam);
/** Jacobian d(dH/dx)/dt */
void dHdxdt(typeNum *out, ctypeNum t, ctypeNum *x, ctypeNum *u, ctypeNum *p,

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→ctypeRNum *vec, const typeGRAMPCparam *param, typeUSERPARAM *userparam);  
/** Mass matrix in vector form (column-wise, either banded or full matrix) **/  
void Mfct(ctypeRNum *out, const typeGRAMPCparam *param, typeUSERPARAM *userparam);  
/** Transposed mass matrix in vector form (column-wise, either banded or full matrix) **/  
void Mtrans(ctypeRNum *out, const typeGRAMPCparam *param, typeUSERPARAM *userparam);
```