

# User Manual of *ParNMPC* by Examples

*A Code Generation Toolbox for Parallel Nonlinear Model  
Predictive Control with OpenMP*

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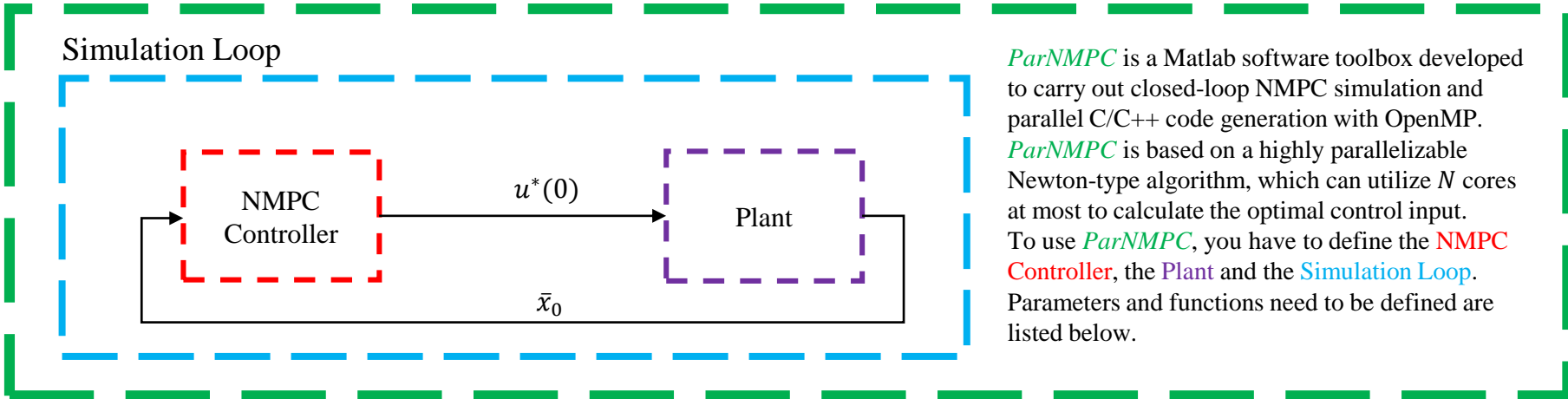
- *Highly Parallelizable*
- *Fast Rate of Convergence*
- *Parallel C/C++ Code Generation*
- *Closed-loop Simulation*
- *Easy to Use*

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# Introduction

## ParNMPC



*ParNMPC* is a Matlab software toolbox developed to carry out closed-loop NMPC simulation and parallel C/C++ code generation with OpenMP. *ParNMPC* is based on a highly parallelizable Newton-type algorithm, which can utilize  $N$  cores at most to calculate the optimal control input. To use *ParNMPC*, you have to define the **NMPC Controller**, the **Plant** and the **Simulation Loop**. Parameters and functions need to be defined are listed below.

### NMPC Controller

$$\begin{aligned} \min_{x(\cdot), u(\cdot)} \int_0^T L(u(t), x(t), p(t)) dt \\ \text{s.t. } \dot{x}(t) = f(u(t), x(t), p(t)), \\ x(0) = \bar{x}_0, \\ C(u(t), x(t), p(t)) = 0, t \in (0, T]. \end{aligned}$$

#### Functions:

- $f$ : dynamic model for controller design
- $L$ : cost function
- $C$ : constraint

#### Parameters:

- $x$ : state vector
- $u$ : input vector
- $p$ : parameter vector
- $\bar{x}_0$ : initial state vector
- $T$ : prediction horizon
- $N$ : number of discretization grids

### Plant

$$\begin{aligned} \dot{x}_{sim} &= f_{sim}(u_{sim}, x_{sim}, p_{sim}) \\ \bullet \quad &f_{sim}: \text{dynamic model of the plant} \\ \bullet \quad &p_{sim}: \text{parameter in the plant model} \end{aligned}$$

### Simulation Loop

#### Simulation parameters:

- Sampling interval  $T_s$
- Simulation length

# Introduction

- In practice, *ParNMPC* solves the discretized optimal control problem (OCP) for a given  $\bar{x}_0$ , and apply  $u_1^*$ .

$$\begin{aligned} \min_{x(\cdot), u(\cdot)} \int_0^T L(u(t), x(t), p(t)) dt \\ \text{s.t. } \dot{x}(t) = f(u(t), x(t), p(t)), \\ x(0) = \bar{x}_0, \\ C(u(t), x(t), p(t)) = 0, t \in (0, T]. \end{aligned}$$

Discretization with the  
implicit Euler method  $\longrightarrow$

$$\begin{aligned} \min_{X, U} \sum_{i=1}^N L(u_i, x_i, p_i) \Delta\tau \\ \text{s.t. } x_i = x_{i-1} + f(u_i, x_i, p_i) \Delta\tau, \\ x_0 = \bar{x}_0, \\ C(u_i, x_i, p_i) = 0, i = 1, \dots, N, \\ \text{where } X = (x_1, x_2, \dots, x_N), \\ U = (u_1, u_2, \dots, u_N), \\ \Delta\tau = T/N. \end{aligned}$$

- More specifically, *ParNMPC* solves the KKT conditions of the discretized OCP, i.e., the following nonlinear algebraic equations with respect to the costates  $\{\lambda_i\}_{i=1}^N$ , the multipliers  $\{\mu_i\}_{i=1}^N$ , the inputs  $\{u_i\}_{i=1}^N$  and the states  $\{x_i\}_{i=1}^N$ :

$$\begin{bmatrix} x_{i-1} - x_i + f(u_i, x_i, p_i) \Delta\tau \\ C(u_i, x_i, p_i) \Delta\tau \\ H_u^T(\lambda_i, \mu_i, u_i, x_i, p_i) \Delta\tau \\ \lambda_{i+1} - \lambda_i + H_x^T(\lambda_i, \mu_i, u_i, x_i, p_i) \Delta\tau \end{bmatrix}_{i=1}^N = 0 \in \mathbb{R}^{N(2n_x + n_\mu + n_u)}$$

Here,  $H(\lambda, \mu, u, x, p) := L(u, x, p) + \lambda^T f(u, x, p) + \mu^T C(u, x, p)$  denotes the Hamiltonian,  $\lambda \in \mathbb{R}^{n_x}$ ,  $\mu \in \mathbb{R}^{n_\mu}$ ,  $u \in \mathbb{R}^{n_u}$ ,  $x \in \mathbb{R}^{n_x}$ ,  $p \in \mathbb{R}^{n_p}$ ,  $x_0 = \bar{x}_0$  and  $\lambda_{N+1} = 0$ .

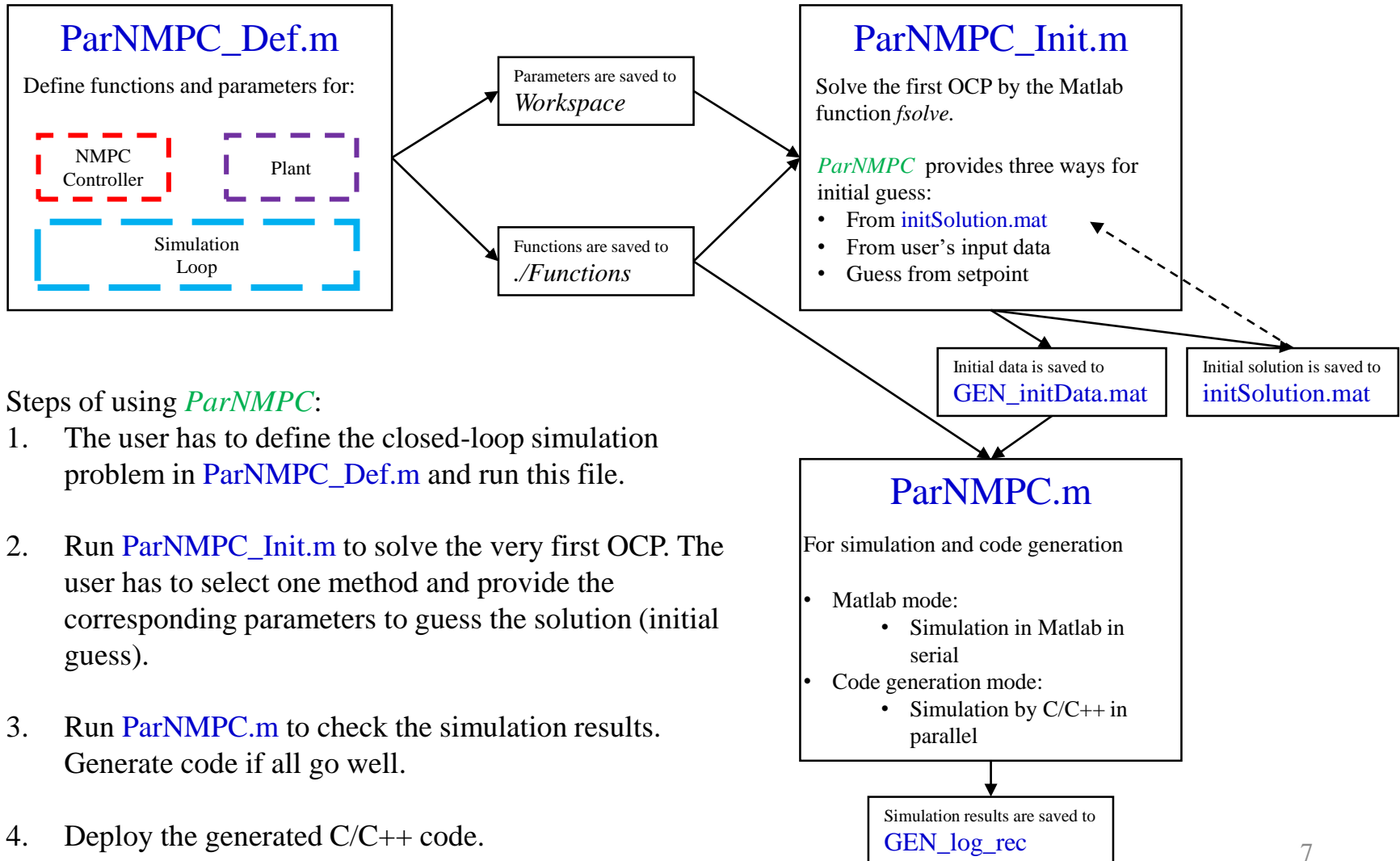
# Installation

1. Download [ParNMPC\\_Toolbox.rar](#)
  2. Extract files (5 folders):
    - *CSTR*
    - *DoubleInvertedPendulum*
    - *Quadrotor*
    - *SemiActiveDamper*
    - *UserManual*
  3. Go into one of the application folders, e.g., *Quadrotor*
  4. Files and folders in *Quadrotor*:
    - Folders
      - *codegen* (Pre-generated C code)
      - *Functions* (Functions and scripts)
    - Files:
      - [ParNMPC\\_Def.m](#) (Define the simulation problem)
      - [ParNMPC\\_Init.m](#) (Solve the first OCP)
      - [ParNMPC.m](#) (For simulation and code generation)
      - [initialSolution.mat](#) (Solution for the first OCP)
- ParNMPC is application-oriented and each application folder is self-contained.

# System Requirements

- MATLAB 2016a or later
  - MATLAB Coder
  - Optimization Toolbox
  - Parallel Computing Toolbox
  - Symbolic Math Toolbox
- Compiler
  - C/C++ Compilers supporting OpenMP for code generation
    - See [Supported and Compatible Compilers](#)
- Tested Environments:
  - Windows 10 + MATLAB 2016a + Microsoft Visual C++ 2015 Professional
  - Windows 10 + MATLAB 2017b + Microsoft Visual C++ 2017 Community

# Flow of *ParNMPC*



## Steps of using *ParNMPC*:

1. The user has to define the closed-loop simulation problem in *ParNMPC\_Def.m* and run this file.
2. Run *ParNMPC\_Init.m* to solve the very first OCP. The user has to select one method and provide the corresponding parameters to guess the solution (initial guess).
3. Run *ParNMPC.m* to check the simulation results. Generate code if all go well.
4. Deploy the generated C/C++ code.

# Getting Started

## *Semi-Active Damper*

### Simulation Loop:

- Sampling interval: 0.01 s
- Simulation length: 20 s

### Plant:

- Same as in NMPC:

$$f_{sim}(u, x, p_{sim}) = \begin{bmatrix} x_2 \\ ax_1 + bx_2u_1 \end{bmatrix}$$

### NMPC Controller:

- Dynamics:

$$f(u, x, p) = \begin{bmatrix} x_2 \\ ax_1 + bx_2u_1 \end{bmatrix}, a = b = -1$$

- Constraints (Converted into equality constraint by introducing dummy input  $u_2$  (see *ref*)):

$$-1 \leq u_1 \leq 1 \longrightarrow C(u, x, p) = u_1^2 + u_2^2 - 1$$

- Stage cost function:

$$L(u, x, p) = \frac{1}{2} (\|x - x_{ref}\|_Q^2 + \|u - u_{ref}\|_R^2 - r \cdot u)$$

$$Q = \text{diag}(10, 10), R = \text{diag}(0.01, 0.01), r = [0, 0.1]$$

$$x_{ref} = [0, 0]^T, u_{ref} = [0, 1]^T$$

- Initial state:  $\bar{x}_0 = [1, 0]^T$
- Prediction horizon: 2 s
- Number of discretization grids:  $N = 36$
- Max number of iterations: 5
- Tolerance: 1e-5



# Getting Started

## *Semi-Active Damper – Problem Definition*

Step 1:  
Define functions and parameters  
as follows in `ParNMPC_Def.m`  
and run `ParNMPC_Def.m`.

### Simulation Loop:

```
Ts = 0.01;           % sampling interval
simuLength = 20;    % simulation length
```

### Plant:

```
% Definition of dimension
pSimDim = 0; % dimension of the parameter in the
simulation model

% Definition of system dynamics f_sim(u,x,p_sim)
fSim([u;x;pSim]) =
    [x(2); a * x(1) + b * x(2) * u(1)];

% init pSim
pSimVal = zeros(pSimDim,1);
```

### NMPC Controller:

```
% Definition of dimensions
xDim = 2; % dimension of states
uDim = 2; % dimension of inputs including dummy inputs
muDim = 1; % dimension of constraints
pDim = 0; % dimension of parameters for the controller

% Definition of system dynamics f(u,x,p)
a = -1;
b = -1;
f([u;x;p]) = [x(2); a * x(1) + b * x(2) * u(1)];

% Definition of equality constraint C(u,x,p)
uMax = 1;
uMin = -1;
uBar = (uMax + uMin)/2;
C([u;x;p]) = (u(1) - uBar)^2 + u(2)^2 - (uMax - uBar)^2;

% Definition of cost function L(u,x,p)
Q = diag([10, 10]);
R = diag([0.01, 0.01]);
r = [0,0.1];
xRef = [0;0];
uRef = [0;uMax];
L([u;x;p]) = 0.5*(x-xRef).'*Q*(x-xRef)...
             +0.5*(u-uRef).'*R*(u-uRef)...
             -0.5*r*u;

% Definition of NMPC settings
x0Value = [1;0]; % init state
T = 2; % prediction horizon
N = 24; % number of discretization grids
pVal = zeros(pDim,N); % parameters
MaxIterNum = 5; % max num of iterations per update
tolerance = 1e-5; % tolerance of the KKT conditions to terminate
```

# Getting Started

## *Semi-Active Damper – Initialization*

Step 2:

Define the initialization method by `initMethod = INIT_BY_FILE` in [ParNMPC\\_Init.m](#) and run [ParNMPC\\_Init.m](#), solution will be initialized from file [initialSolution.mat](#).

- If [initialSolution.mat](#) is not available, another method should be selected by either `initMethod = INIT_BY_INPUT` or `INIT_BY_REF_GUESS`, and the corresponding parameters should be provided.
- The optimal solution will be automatically saved to [initialSolution.mat](#).

# Getting Started

## *Semi-Active Damper – Simulation & Code Generation*

Step 3:

1. Run [ParNMPC.m](#) to validate the closed-loop simulation. The results will be saved to [GEN\\_log\\_rec.mat](#).
  - This step is not in parallel. Only the generated C/C++ code is in parallel.
  - If the simulation doesn't go well, it is most likely that the first OCP is not well solved, or the measured state and time-dependent parameter change dramatically.
2. Code generation:
  1. Set up your compiler by *mex -setup*.
  2. Open the MATLAB Coder app.
  3. Select [ParNMPC.m](#).
  4. Click Generate.

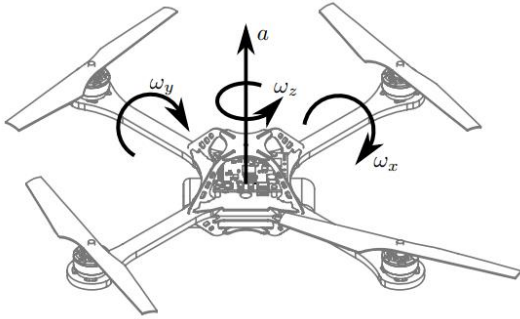
# Getting Started

## *Semi-Active Damper – Deployment*

Step 4:

1. Open Visual Studio.
2. Create an empty Win32 Console Application project.
3. Change to Release x64 mode.
4. Add \*.h and \*.c files in `\codegen\lib\ParNMPC` to project.
5. Add \*.h and \*.c files in `\codegen\lib\ParNMPC\examples` to project.
6. Add the folder `\codegen\lib\ParNMPC` to *Properties* → *C/C++* → *General* → *Additional Include Directories*.
7. *Properties* → *C/C++* → *Language* → *Open MP Support: Yes (/openmp)*.
8. Compile and run.

# Getting Started *Quadrotor*



## Simulation Loop:

- Sampling interval: 0.01 s
- Simulation length: 20 s

## Plant:

- Same as in NMPC:

$$\begin{aligned}\ddot{X} &= a(\cos \gamma \sin \beta \cos \alpha + \sin \gamma \sin \alpha) \\ \ddot{Y} &= a(\cos \gamma \sin \beta \sin \alpha - \sin \gamma \cos \alpha) \\ \ddot{Z} &= a \cos \gamma \cos \beta - g \\ \dot{\gamma} &= (\omega_X \cos \gamma + \omega_Y \sin \gamma) / \cos \beta \\ \dot{\beta} &= -\omega_X \sin \gamma + \omega_Y \cos \gamma \\ \dot{\alpha} &= \omega_X \cos \gamma \tan \beta + \omega_Y \sin \gamma \tan \beta + \omega_Z\end{aligned}$$

## NMPC Controller:

- States:

$$x = [X \ \dot{X} \ Y \ \dot{Y} \ Z \ \dot{Z} \ \gamma \ \beta \ \alpha]^T \in \mathbb{R}^9$$

- Dynamics:

$$\begin{aligned}\ddot{X} &= a(\cos \gamma \sin \beta \cos \alpha + \sin \gamma \sin \alpha) \\ \ddot{Y} &= a(\cos \gamma \sin \beta \sin \alpha - \sin \gamma \cos \alpha) \\ \ddot{Z} &= a \cos \gamma \cos \beta - g \\ \dot{\gamma} &= (\omega_X \cos \gamma + \omega_Y \sin \gamma) / \cos \beta \\ \dot{\beta} &= -\omega_X \sin \gamma + \omega_Y \cos \gamma \\ \dot{\alpha} &= \omega_X \cos \gamma \tan \beta + \omega_Y \sin \gamma \tan \beta + \omega_Z\end{aligned}$$

- Constraints (Converted to equality constraints by introducing dummy inputs  $u_5, u_6, u_7, u_8$ ):

$$\begin{bmatrix} 0 \\ -1 \\ -1 \\ -1 \end{bmatrix} \leq \begin{bmatrix} a \\ \omega_X \\ \omega_Y \\ \omega_Z \end{bmatrix} \leq \begin{bmatrix} 11 \\ 1 \\ 1 \\ 1 \end{bmatrix} \rightarrow C(u, x, p) = \begin{bmatrix} (a - 5.5)^2 + u_5^2 - 5.5^2 \\ \omega_X^2 + u_6^2 - 1 \\ \omega_Y^2 + u_7^2 - 1 \\ \omega_Z^2 + u_8^2 - 1 \end{bmatrix}$$

- Stage cost function:

$$L(u, x, p) = \frac{1}{2} (\|x - x_{ref}\|_Q^2 + \|u - u_{ref}\|_R^2 - r \cdot u)$$

$$Q = \text{diag}(10, 1, 2, 1, 10, 1, 1, 1, 1), R = \text{blkdiag}(I_4, 0.01 \cdot I_4)$$

$$r = [\text{zeros}(1, 4), 0.1 \cdot \text{ones}(1, 4)]$$

$$u_{ref} = [g, 0, 0, 0, 3.4167, 1, 1, 1]^T$$

$$x_{ref}(t) = \text{zeros}(9, 1), t \in [0, 10]$$

$$x_{ref}(t) = [1.5, 0, 1.5, 0, 1.5, 0, 0, 0]^T, t \in (10, 20]$$

- Initial state:  $\bar{x}_0 = [1, 0, 1, 0, 1, 0, 0, 0]^T$
- Prediction horizon: 1 s
- Number of discretization grids:  $N = 24$
- Max number of iterations: 5
- Tolerance: 5e-3

# Getting Started

## Quadrotor – Problem Definition

Step 1:  
Define functions and parameters as follows in `ParNMPC_Def.m` and run `ParNMPC_Def.m`.

### Simulation Loop:

```
Ts = 0.01; %sampling interval
simuLength = 10; %simulation length in seconds
```

### Plant:

```
% Definition of dimensions
pSimDim = 0; % dimension of the parameter in the simulation model

% Definition of system dynamics f_sim(u,x,p_sim)
fSim([u;x;pSim]) = [x(2);...
    u(1)*(cos(x(7))*sin(x(8))*cos(x(9)) + sin(x(7))*sin(x(9)));...
    x(4);...
    u(1)*(cos(x(7))*sin(x(8))*sin(x(9)) - sin(x(7))*cos(x(9)));...
    x(6);...
    u(1)*cos(x(7))*cos(x(8)) - g;...
    (u(2)*cos(x(7)) + u(3)*sin(x(7)))/cos(x(8));...
    -u(2)*sin(x(7)) + u(3)*cos(x(7));...
    u(2)*cos(x(7))*tan(x(8)) + u(3)*sin(x(7))*tan(x(8)) + u(4)];

% Init pSim
pSimVal = zeros(pSimDim,1);
```

### NMPC Controller:

```
% Definition of dimensions
xDim = 9; % dimension of states
uDim = 8; % dimension of inputs including dummy inputs
muDim = 4; % dimension of constraints
pDim = xDim+uDim; % dimension of parameters for the controller

% Definition of system dynamics f(u,x,p)
g = 9.81;
f([u;x;p]) = [x(2);...
    u(1)*(cos(x(7))*sin(x(8))*cos(x(9)) + sin(x(7))*sin(x(9)));...
    x(4);...
    u(1)*(cos(x(7))*sin(x(8))*sin(x(9)) - sin(x(7))*cos(x(9)));...
    x(6);...
    u(1)*cos(x(7))*cos(x(8)) - g;...
    (u(2)*cos(x(7)) + u(3)*sin(x(7)))/cos(x(8));...
    -u(2)*sin(x(7)) + u(3)*cos(x(7));...
    u(2)*cos(x(7))*tan(x(8)) + u(3)*sin(x(7))*tan(x(8)) + u(4)];

% Definition of equality constraint C(u,x,p)
uMax = [1;1;1;1];
uMin = [0;-1;-1;-1];
uBar = (uMax + uMin)/2;
C([u;x;p]) = [(u(1) - uBar(1))^2 + u(5)^2 - (uMax(1)-uBar(1))^2;...
    (u(2) - uBar(2))^2 + u(6)^2 - (uMax(2)-uBar(2))^2;...
    (u(3) - uBar(3))^2 + u(7)^2 - (uMax(3)-uBar(3))^2;...
    (u(4) - uBar(4))^2 + u(8)^2 - (uMax(4)-uBar(4))^2];

% Definition of cost function L(u,x,p)
Q = diag([10, 1, 2, 1, 10, 1, 1, 1, 1]);
R = diag([1, 1, 1, 1, ...
    0.01, 0.01, 0.01, 0.01]);
r = [0,0,0,0,...
    1,1,1,1]*0.1;
L([u;x;p]) = 0.5*(x-p(1:xDim)).'*Q*(x-p(1:xDim))...
    +0.5*(u-p(pDim-uDim+1:pDim)).'*R*(u-p(pDim-uDim+1:pDim))...
    -0.5*r*u;

% Definition of NMPC settings
x0Value = [1;0;1;0;1;0;0;0;0]; % initial state
T = 1.0; % prediction horizon
N = 24; % number of discretization grids
xRefConstant = [0;0;0;0;0;0;...
    0;0;0;0];
uRefConstant = [g;0;0;0;...
    sqrt(-(g - uBar(1))^2 + (uMax(1)-uBar(1))^2);...
    uMax(2);uMax(3);uMax(4)];
pVal = repmat([xRefConstant;uRefConstant],1,N); % init parameters
MaxIterNum = 5; % max num of iterations per update
tolerance = 5e-3; % tolerance of the KKT conditions to terminate
```

# Getting Started

## *Quadrotor – Initialization*

- Step 2: refer to [\*Semi-Active Damper – Initialization\*](#)

## *Quadrotor– Simulation & Code Generation*

- Step 3: refer to [\*Semi-Active Damper – Simulation & Code Generation\*](#)

## *Quadrotor– Deployment*

- Step 4: refer to [\*Semi-Active Damper – Deployment\*](#)

# Main Reasons Why *ParNMPC* Fails

1. The first OCP is not well solved. *ParNMPC* utilizes the warm-start strategy and highly depends on the solution of the first OCP. *ParNMPC* may fail even when the norm of the KKT conditions (first-order optimality) for the first OCP is small enough. In general, a good solution is smooth enough. In order to find a good solution to the first OCP, the method of continuation can be employed. For example, the solution of the first OCP can be initialized from the solution of an easy-to-solve problem, which has less active constraints or a shorter prediction horizon.
2. Dramatic changes of the measured state or time-dependent parameter. *ParNMPC* converges locally and requires that the current solution is close to the last solution. The current version of *ParNMPC* doesn't contain any globalization techniques such as line-search and trust-region methods. Make sure that the state and parameter don't vary too much with respect to time.
3. Ill-conditioned problem. It is usually caused by the settings of the weighting matrices.



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