

A new quadratic programming strategy for efficient sparsity exploitation in SQP-based nonlinear MPC and MHE^{*}

Janick V. Frasch^{*,**} Milan Vukov^{*} Hans Joachim Ferreau^{***}
Moritz Diehl^{*}

^{*} *Department of Electrical Engineering, KU Leuven, Belgium (e-mail: janick.frasch@esat.kuleuven.be).*

^{**} *Department of Mathematics, Magdeburg University, Germany*

^{***} *ABB Corporate Research, Baden-Dättwil, Switzerland*

Abstract: A large class of algorithms for nonlinear model predictive control (MPC) and moving horizon estimation (MHE) is based on sequential quadratic programming and thus requires the solution of a sparse structured quadratic program (QP) at each sampling time. We propose a novel algorithm based on a dual two-level approach involving a nonsmooth version of Newton's method that aims at combining sparsity exploitation features of an interior point method with warm-starting capabilities of an active-set method. We address algorithmic details and present the open-source implementation qpDUNES. The effectiveness of the solver in combination with the ACADO Code Generation tool for nonlinear MPC is demonstrated based on set of benchmark problems, showing significant performance increases compared to the established condensing-based approach, particularly for problems with long prediction horizons.

Keywords: Predictive Control, Optimization algorithms, Open-Source Software

1. INTRODUCTION

Model predictive control (MPC) is an approach to obtain a feedback control law taking physical process models and problem-inherent constraints into account, cf. Rawlings and Mayne (2009). It relies on the online solution of an optimization problem in each feedback-generating iteration. For high predictive accuracy and good stability properties it is desirable to design MPC controllers that can deal with detailed process descriptions to reduce the prediction-inherent uncertainty, resulting in big dynamic systems and long prediction horizons. Particularly the long prediction horizons, however, can render the underlying optimization problem very challenging for a real-time implementation.

Most nonlinear MPC algorithms require the direct solution of an (in general) *nonlinear programming problem* (NLP) online. In this paper we focus on approaches that base on sequential quadratic programming (SQP) like the real-time iteration (RTI) scheme, see Diehl et al. (2002). Particularly in combination with automatic code generation, the RTI scheme can lead to very short computation times while retaining sufficient accuracy (see, e.g., Houska et al. (2011)) and therefore seems to be a well suited approach for embedded nonlinear MPC of high-speed applications.

^{*} This research was supported by Research Council KUL: PFV/10/002 Optimization in Engineering Center OPTEC, GOA/10/09 MaNet and GOA/10/11 Global real-time optimal control of autonomous robots and mechatronic systems. Flemish Government: FWO: PhD/postdoc grants; IWT: PhD Grants, projects: Eurostars SMART; Belgian Federal Science Policy Office: IUAP P7 (DYSCO, Dynamical systems, control and optimization, 2012-2017); EU: FP7-SADCO (MC ITN-264735), FP7-TEMPO(MC ITN-607957), ERC HIGHWIND (259 166).

Due to the structural equivalence, *moving horizon estimation* (MHE) can be cast into the same framework and solved efficiently using the RTI scheme, see Kühn et al. (2011). We subsume both problem classes, MHE and MPC, under the term MPC for clarity of the presentation.

A crucial step when using SQP-based algorithms on problems featuring long prediction horizons is the efficient solution of the *highly structured quadratic programming problem* (QP) in each iteration, which is usually performed by tailored algorithms based on interior point methods (e.g., Domahidi et al. (2012)), active-set methods (e.g., Ferreau et al. (2008)), or fast-gradient methods (e.g., Patrinos and Bemporad (2014)). Both classes of methods, interior-point and fast-gradient methods on the one hand, and active-set methods on the other hand, have significant shortcomings in the context of MPC and MHE. While the former ones generally cannot exploit the knowledge about similarity between the solutions of two subsequent QPs in the MPC context, the latter ones typically require a so-called condensing routine to benefit from the problem-inherent sparsity. Even though recent results from Andersson et al. (2013) lead to an improved performance of the condensing step, its overall runtime complexity is still quadratic in the horizon length and cannot be expected to be accelerated further.

In this paper, we present a new idea for solving strictly convex quadratic subproblems in the context of nonlinear MPC and MHE. It is based on ideas for a band-structured QP solver that were introduced in Ferreau et al. (2012). Based on original ideas from Li and Swetits (1997) the stage coupling constraints of the MPC problem are dual-

ized and the resulting QP is solved in a two level approach, using a non-smooth Newton method in the multipliers of the stage coupling constraints on the higher level, and a primal active-set method in the decoupled parametric QPs of each stage on the lower level. The advantage of this so-called *dual Newton strategy* is that it combines structure exploitation capabilities of interior point methods with the warm-starting capabilities of active set methods; in particular it comes with only a linear runtime complexity in the horizon length. Note that in contrast to classical active-set methods this approach permits several active-set changes in each Newton-type iteration. Still, the resulting algorithm has the flavor of an active-set method in the sense that the exact optimal solution is obtained.

In this paper, we address the core ideas of the dual Newton strategy and show how it can be applied efficiently for nonlinear MPC and MHE. Most importantly, we present qpDUNES, an open-source implementation of the *DUAL NEWTON STRATEGY* that is now available as structure-exploiting QP solver in the open-source ACADO toolkit (see Houska et al. (2011)) for nonlinear MPC and MHE of high-speed applications. We demonstrate the effectiveness of qpDUNES in comparison against a code-generated condensing/qpOASES approach first described in Houska et al. (2011); Ferreau et al. (2008) based on a selection of challenging nonlinear MPC benchmark problems.

2. QP SOLUTION METHOD

2.1 Quadratic subproblem description

We assume familiarity of the reader with the RTI scheme from Diehl et al. (2002). Within the RTI scheme we repeatedly need solve the following subproblem, that can be interpreted as a linear MPC problem. Here, we group the optimization variables, state increments $\Delta x_k \in \mathbb{R}^{n_x}$ and control increments $\Delta u_k \in \mathbb{R}^{n_u}$, in stage variables $z_k = [\Delta x_k^\top \ \Delta u_k^\top]^\top \in \mathbb{R}^{n_z}$ for each stage $k = 0, \dots, N-1$, and $z_N = [\Delta x_N \ 0]$ for the terminal stage. Throughout this paper we are consequently interested in repeatedly solving the following problem in an efficient manner:

$$\min_z \sum_{k=0}^N \left(\frac{1}{2} z_k^\top H_k z_k + g_k^\top z_k \right) \quad (1a)$$

$$\text{s.t. } E_{k+1} z_{k+1} = C_k z_k + c_k \quad \forall k = 0, \dots, N-1 \quad (1b)$$

$$\underline{d}_k \leq D_k z_k \leq \bar{d}_k \quad \forall k = 0, \dots, N. \quad (1c)$$

We assume positive definite second-order terms $0 \prec H_k \in \mathbb{R}^{n_z \times n_z}$ and $g_k \in \mathbb{R}^{n_z}$ for each $k \in \mathcal{S} := \{0, \dots, N\}$ throughout the paper. Two subsequent stages $k \in \mathcal{S}$ and $k+1 \in \mathcal{S}$ are coupled by first-order terms $C_k, E_{k+1} \in \mathbb{R}^{n_x \times n_z}$ and a constant term c_k . We assume that all C_k have full row rank; each E_k has the structure $E_k = [I \ 0]$, with an identity matrix $I \in \mathbb{R}^{n_x \times n_x}$ complemented by zeros. Vectors $\underline{d}_k, \bar{d}_k \in \mathbb{R}^{n_d}$, and a matrix $D_k \in \mathbb{R}^{n_d \times n_z}$ of full row rank denote stage constraints.

2.2 Dual decomposition

The main idea of our proposed QP solution algorithm is to decouple the QP stages by dualizing constraints (1b).

Introducing $\lambda := [\lambda_1^\top \ \lambda_2^\top \ \dots \ \lambda_N^\top]^\top \in \mathbb{R}^{N n_x}$ we can express (1a) and (1b) by the partial Lagrangian function

$$\begin{aligned} \mathcal{L}(z, \lambda) &= \sum_{k=0}^N \left(\frac{1}{2} z_k^\top H_k z_k + g_k^\top z_k \right. \\ &\quad \left. + \begin{bmatrix} \lambda_k \\ \lambda_{k+1} \end{bmatrix}^\top \begin{bmatrix} -E_k \\ C_k \end{bmatrix} z_k + \lambda_{k+1}^\top c_k \right) \\ &=: \sum_{k=0}^N L_k(z_k, \lambda_k, \lambda_{k+1}), \end{aligned}$$

where we define zero matrices $E_0 := C_N := 0 \in \mathbb{R}^{n_x \times n_z}$ and redundant multipliers $\lambda_0 := \lambda_{N+1} := 0 \in \mathbb{R}^{n_x}$ only for notational convenience.

By elementary Lagrangian duality theory the primal QP (1) is equivalent to

$$\begin{aligned} \max_{\lambda} \min_z \sum_{k=0}^N L_k(z_k, \lambda_k, \lambda_{k+1}) \\ \text{s.t. } \underline{d}_k \leq D_k z_k \leq \bar{d}_k \quad \forall k = 0, \dots, N \end{aligned}$$

Observe that this Problem is separable in the stage variables z_k . Problem (1) can thus equivalently be written as

$$\max_{\lambda} f^*(\lambda) := \max_{\lambda} \sum_{k=0}^N f_k^*(\lambda), \quad (2)$$

where

$$\begin{aligned} f_k^*(\lambda) &:= \min_{z_k} \frac{1}{2} z_k^\top H_k z_k + m_k(\lambda)^\top z_k + \lambda_{k+1}^\top c_k \\ \text{s.t. } \underline{d}_k &\leq D_k z_k \leq \bar{d}_k \quad (\text{QP}_k) \end{aligned}$$

with $m_k(\lambda) := g_k^\top + \begin{bmatrix} \lambda_k \\ \lambda_{k+1} \end{bmatrix}^\top \begin{bmatrix} -E_k \\ C_k \end{bmatrix}$. From this definition of the dual function f^* we can particularly see that $z(\lambda) := [z_0(\lambda), \dots, z_N(\lambda)]$ is optimal for each choice of λ .

2.3 Non-smooth Newton approach

Under the assumption that a feasible solution for (1) exists, one can show that $f^*(\lambda)$ exists on $\mathbb{R}^{N n_x}$ and further is a concave, piecewise quadratic, and once continuously differentiable function, cf. Ferreau et al. (2012); Fräsch et al. (2013) and the references therein. We solve the unconstrained piecewise-quadratic program (2) by employing a non-smooth Newton method, as originally proposed in Li and Swetits (1997). The second derivative $\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda^i)$ is unique everywhere but on a null set (the “seams” of f^*), where in general a jump occurs. The proposed QP solution method is given by Algorithm 1, where we denote the Lagrange multipliers of constraints (1c) by $\mu_k \in \mathbb{R}^{2n_d}$ for each $k \in \mathcal{S}$. We refer to Fräsch et al. (2013) for a proof of convergence of this algorithm to the (unique) optimal dual solution λ^* , that is characterized by stationary of $f^*(\lambda)$ and implicitly defines $z^* := z(\lambda^*)$ via (QP_k).

3. ALGORITHMIC DETAILS OF THE DUAL NEWTON STRATEGY

Due to its temporal coupling, Problem (2) possesses a specific structure that can be exploited for its efficient

Algorithm 1: Dual Newton Strategy

Input: Initial guess λ^0 , termination criteria $n_{\max\text{It}}$, ϵ_λ

Output: Optimal solution (z^*, λ^*, μ^*)

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1  $i := 0$ 
2 while  $i < n_{\max\text{It}}$  do
3   Solve all  $\text{QP}_k(\lambda^i)$  to obtain  $[z_k^*(\lambda^i), \mu_k^*(\lambda^i)]$ 
4   Compute  $\left[-\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda^i)\right]$  and  $\left[\frac{\partial f^*}{\partial \lambda}(\lambda^i)\right]$ 
   analytically
5   if  $\left\|\frac{\partial f^*}{\partial \lambda}(\lambda^i)\right\| \leq \epsilon_\lambda$  then
6     return  $[z_k^*(\lambda^i), \lambda^i, \mu_k^*(\lambda^i)]$ 
7   Solve Newton system  $-\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda^i) \Delta \lambda = \frac{\partial f^*}{\partial \lambda}(\lambda^i)$ 
8   Compute step size  $\alpha$ 
9   Update the current iterate  $\lambda^{i+1} := \lambda^i + \alpha \Delta \lambda$ 
10   $i := i+1$ 

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solution. We analyze details for the individual steps of Algorithm 1 in the following.

3.1 Structure-exploiting solution of the Newton System

The dual gradient $\frac{\partial f^*}{\partial \lambda}(\lambda) \in \mathbb{R}^{Nn_x}$ (as a column vector) is easily seen to only depend on two neighboring stages in each block λ_k . It holds

$$\frac{\partial f^*}{\partial \lambda}(\lambda) := \begin{bmatrix} \frac{\partial f_0^*}{\partial \lambda_1} + \frac{\partial f_1^*}{\partial \lambda_1} \\ \frac{\partial f_1^*}{\partial \lambda_2} + \frac{\partial f_2^*}{\partial \lambda_2} \\ \vdots \\ \frac{\partial f_{N-1}^*}{\partial \lambda_N} + \frac{\partial f_N^*}{\partial \lambda_N} \end{bmatrix}(\lambda).$$

The dual Hessian $\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda) \in \mathbb{R}^{Nn_x \times Nn_x}$ possesses a block tri-diagonal structure, as only neighboring multipliers λ_k, λ_{k+1} can have a joint contribution to f^* . We have

$$\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda) := \begin{bmatrix} \frac{\partial^2 f^*}{\partial \lambda_1^2} & \frac{\partial^2 f^*}{\partial \lambda_1 \lambda_2} & & & \\ \frac{\partial^2 f^*}{\partial \lambda_2 \lambda_1} & \frac{\partial^2 f^*}{\partial \lambda_2^2} & & & \\ & & \ddots & & \\ & & & \ddots & \\ & & & & \frac{\partial^2 f^*}{\partial \lambda_{N-1} \lambda_N} \\ & & & & \frac{\partial^2 f^*}{\partial \lambda_N \lambda_{N-1}} & \frac{\partial^2 f^*}{\partial \lambda_N^2} \end{bmatrix}(\lambda).$$

3.2 Dual function evaluation

Each stage problem (QP_k) has a fixed second order term H_k and a parametric first-order term $p_k(\lambda)$. An efficient method to solve such parametric problems repeatedly for changing parameter values λ is the so-called online active-set strategy (see Ferreau et al. (2008)); a well-established implementation of this algorithm that we propose to employ here is the open-source QP solver qpOASES, cf. Ferreau et al. (2008).

In the special case where H_k is a diagonal matrix and $D_k = I$ is an identity matrix (i.e., only bounds on states and controls of the MPC problem exist) the optimal solution z_k^* can conveniently be computed by component-wise

“clipping” of the unconstrained solution, here denoted by z_k , as it was presented in Ferreau et al. (2012):

$$z_k^* = \max(\underline{d}_k, \min(z_k, \bar{d}_k)) \quad (3)$$

3.3 Explicit derivative computation

Due to the strict positive definiteness of H_k the derivative of each dual function summand f_k^* with respect to the multiplier contribution $\begin{bmatrix} \lambda_k \\ \lambda_{k+1} \end{bmatrix}$ exists for all $k \in \mathcal{S}$, and is given by (see Bertsekas and Tsitsiklis (1989), App. C for a formal derivation)

$$\begin{bmatrix} \frac{\partial f_k^*}{\partial \lambda_k} & \frac{\partial f_k^*}{\partial \lambda_{k+1}} \end{bmatrix}^\top = z_k^{*\top} \begin{bmatrix} -E_k^\top & C_k^\top \end{bmatrix} + \begin{bmatrix} 0 & c_k \end{bmatrix}. \quad (4)$$

Observe that Equation (4) links the equivalence of primal feasibility and stationarity of the dual function f^* , justifying the termination criterion in Step 5 of Algorithm 1.

Differentiating (4) once more with respect to λ the non-zero dual Hessian blocks read (cf. Frasch et al. (2013))

$$\begin{aligned} \frac{\partial^2 f^*}{\partial \lambda_k \lambda_{k+1}} &= \frac{\partial}{\partial \lambda_k} \left(\frac{\partial f_k^*}{\partial \lambda_{k+1}} + \frac{\partial f_{k+1}^*}{\partial \lambda_{k+1}} \right) \\ &= \frac{\partial z_k^*}{\partial \lambda_k} C_k^\top + \underbrace{\frac{\partial z_{k+1}^*}{\partial \lambda_k} E_{k+1}^\top}_{=0} = E_k P_k^* C_k^\top \end{aligned}$$

and

$$\begin{aligned} \frac{\partial^2 f^*}{\partial \lambda_k \lambda_k} &= \frac{\partial}{\partial \lambda_k} \left(\frac{\partial f_{k-1}^*}{\partial \lambda_k} + \frac{\partial f_k^*}{\partial \lambda_k} \right) \\ &= \frac{\partial z_{k-1}^*}{\partial \lambda_k} C_{k-1}^\top - \frac{\partial z_k^*}{\partial \lambda_k} E_k^\top \\ &= -C_{k-1} P_{k-1}^* C_{k-1}^\top - E_k P_k^* E_k^\top. \end{aligned}$$

Here $P_k^* := Z_k^*(Z_k^{*\top} H_k Z_k^*)^{-1} Z_k^{*\top} \in \mathbb{R}^{n_z \times n_z}$ is an elimination matrix for the nullspace of the optimal active set of QP_k , with a nullspace basis matrix $Z_k^* \in \mathbb{R}^{n_z \times (n_z - n_{\text{act}})}$ suitably chosen to project the Hessian matrix H_k onto the active set of the (unique) optimal solution $z_k^*(\lambda)$ of QP_k , where we use n_{act} to denote the number of active constraints. When employing a null-space QP solver like qpOASES, Z_k^* and a Cholesky factor R for $R^\top R = Z_k^{*\top} H_k Z_k^*$ are directly available, making the computation of P_k^* rather cheap. In the case where Equation (3) is used to solve the stage subproblems, P_k^* boils down to a diagonal matrix with entries either from H_k^{-1} or 0, depending on whether the corresponding variable bound is inactive or active, cf. Ferreau et al. (2012).

3.4 Solution of the Newton system

The dual function f^* is concave, but not strictly concave everywhere, so $-\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda^i)$ is positive semi-definite. In case $-\frac{\partial^2 f^*}{\partial \lambda^2}(\lambda^i)$ is rank-deficient, we propose to add a small regularization term $\epsilon \Delta \lambda$ for $\epsilon > 0$ in Step 7 of Algorithm 1. The resulting, possibly regularized, system $\mathcal{M}(\lambda^i) \Delta \lambda = \frac{\partial f^*}{\partial \lambda}(\lambda^i)$ therefore has unique solution, and a Cholesky factorization $\mathcal{M}(\lambda^i) = LL^\top$ always exists. The

fact that L possesses the same structural zero-blocks as $\mathcal{M}(\lambda^i)$ below the diagonal can be exploited in the factorization algorithm by skipping all redundant blocks left and below the subdiagonal block $-\frac{\partial^2 f^*}{\partial \lambda_{i+1} \lambda_i}$ of each block column i . Utilizing this modified Cholesky decomposition the solution of the (possibly regularized) Newton system requires only $O(Nn_x^3)$ floating point operations (FLOPs).

3.5 Step size selection

The convergence of Algorithm 1 requires a globalization strategy (Step 8 in Algorithm 1) that finds an (approximate) solution to

$$\arg \max_{\alpha \in (0,1]} f^*(\lambda^i + \alpha \Delta \lambda). \quad (5)$$

Note that due to the piecewise quadratic nature of $f^*(\lambda)$ the exact solution of (5) is computationally tractable. Still, we propose to employ an approximate line search strategy for increased computational efficiency. It is important to observe that, based on the fact that Newton's method minimizes a local quadratic model of f^* , we are able to state a lower bound for (5) based on the first primal active-set change in search direction. Readers may convince themselves of this by recalling that first-order kinks f^* are caused by primal active-set changes. A formal proof can be found in Frasch et al. (2013). Step sizes α at which an active set changes occurs are obtained for free when employing an online active set strategy to solve each (QP_k) , cf. Ferreau et al. (2008). In cases where the solution to (QP_k) can be computed by Equation (3), points of active-set changes can still be obtained by a ratio test of the unconstrained and the clipped solution in each component.

3.6 Warm-starting for series of QPs

A key advantage of the dual Newton strategy in the context of nonlinear MPC and MHE are its warm-starting capabilities. While interior point methods typically cannot be warmstarted efficiently, and the active set of a *condensed* QP can, even in the nominal case, change quite significantly from one sampling time to the next due to shifted state constraints, the optimization variables λ can be shifted alongside with the sampling time in Algorithm 1. It is noteworthy that Newton's method guarantees a 1-step convergence in this context if the shifted λ -guess is in the correct quadratic region, i.e., if the optimal primal active set is consistent with the shifted one, even if the QP data changes (e.g., through re-linearization of the nonlinear problem in the RTI framework).

In detail, we suggest to shift from the optimal dual vector of the QP at sampling time s , λ^* , to an initial guess λ^0 of the subsequent QP at sampling time $s+1$ as follows:

$$\begin{aligned} \lambda_k^0 &:= \lambda_{k+1}^* & \forall k = 1, \dots, N-1 \\ \lambda_N^0 &:= \lambda_N^*. \end{aligned}$$

In the nominal case, this shift ensures that λ^0 already lies in the correct quadratic region (and thus 1-step convergence) if the primal terminal stage variables z_N lie in a stable active set (e.g., given by a steady state).

4. OPEN-SOURCE SOFTWARE IMPLEMENTATION

4.1 The structure-exploiting QP solver qpDUNES

The dual Newton strategy has been implemented in the open-source software package qpDUNES, which is available for download at qpDUNES (2013). It is a plain, self-contained C code written according to the C90 standard to enlarge compatibility with embedded hardware platforms. It comes with its own linear algebra module and efficient data storage formats to better exploit the problem intrinsic structures. Memory allocation is performed on a global scale to enable reusability of memory blocks and to enable switching between dynamic memory allocation for maximum flexibility and static memory allocation for increased performance and deployment on embedded hardware. A code generation routine for the linear algebra modules tailored to the structure and dimensions of a specific problem instance for even higher efficiency is currently under development. Application of such code generation techniques has lead to significant performance increases in related areas like interior point solvers, cf. Domahidi et al. (2012).

4.2 The ACADO Code Generation tool

The ACADO Toolkit (available at ACADO (2009–2013)) is an open source software for modeling, simulation and control of nonlinear dynamic processes. It is particularly suited to set up nonlinear MPC and MHE problems. It was recently extended by the ACADO Code Generation tool, see Houska et al. (2011), which allows to export a lean RTI scheme tailored to a specific problem's structure. It is essentially based on Bock's multiple shooting discretization, see Bock and Plitt (1984), and a Gauss-Newton method for the solution of the resulting NLP. For the evaluation of the dynamic system constant step-size explicit and implicit Runge-Kutta integrators are available, cf. Quirynen et al. (2012). In its default configuration, the solution of the structured quadratic subproblems is based on an optimized condensing routine, cf. Andersson et al. (2013) for reduction of the problem size. The dense QPs are then solved using the online QP solver qpOASES, see Ferreau et al. (2008). In the latest release, an alternative interface for a direct solution of the sparse quadratic subproblems via qpDUNES has been added, that avoids the condensing step and to exploits the problem-inherent sparsity structure directly as detailed above.

5. NUMERICAL TESTS

We provide a comparison of these two QP solution strategies within the ACADO Code Generation tool in the following. All simulations were performed on a 3.4GHz Intel i7 based desktop computer, running the 64-bit version of Ubuntu Linux 13.04. All codes are compiled with Clang 3.2.1, using the flag `-O3` and execution times are measured with Linux function `clock_gettime()`.

The first classes of benchmark problems deals with stabilizing a strongly deflected chain of M masses connected by springs after an initial perturbation, as described in Wirsching et al. (2006) and also used for benchmarking in Ferreau et al. (2008) and Vukov et al. (2013). One end of

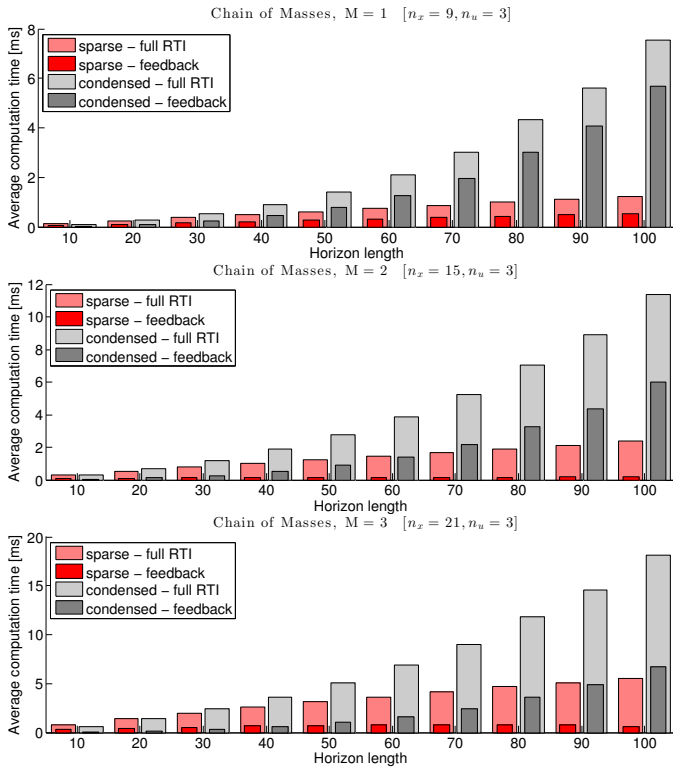


Fig. 1. Average computation time benchmark for three chain-of-masses test cases

the chain is attached to a fixed point, while the velocity of the other end can be controlled. Each mass is described by its position and velocity coordinates, resulting in a total of $n_x = 6M + 3$ states and $n_u = 3$ control inputs. We chose a sampling time of $T_s = 200$ ms and varying prediction horizon lengths $N \in [10, \dots, 100]$. For integration of the nonlinear system dynamics an implicit Gauss-Legendre integrator of order four was used, with two integration step per discretization interval.

In the RTI scheme each iteration is split in a preparation and a feedback phase, which we consider separately (basic familiarity of the reader with the RTI scheme again is assumed here; otherwise we kindly refer to Diehl et al. (2002)). In the condensing-based approach, the preparation phase consists of the linearization of the NLP and the condensing routine that yields the reduced-size QP, both of which are of significant computational effort. The time spent in the feedback phase is dominated by the solution of the condensed QP by qpOASES. In the sparse (qpDUNES-based) approach, no condensing routine is needed, so the preparation phase is dominated only by the effort for the linearization of the NLP. Almost all time of the feedback phase is then spent in the solution of the sparse QP.

We present average computation times for one RTI in Figure 1 and highlight the time spent in the feedback phase (note that the preparation phase factually has to be shorter in the sparse QP strategy). It can be seen that for a moderate number of states n_x , the sparse approach is already competitive on short horizon lengths, while it clearly outperforms the condensing-based approach both in terms of feedback time and in terms of total iteration time on longer horizon lengths due to its lower per-iteration computational complexity.

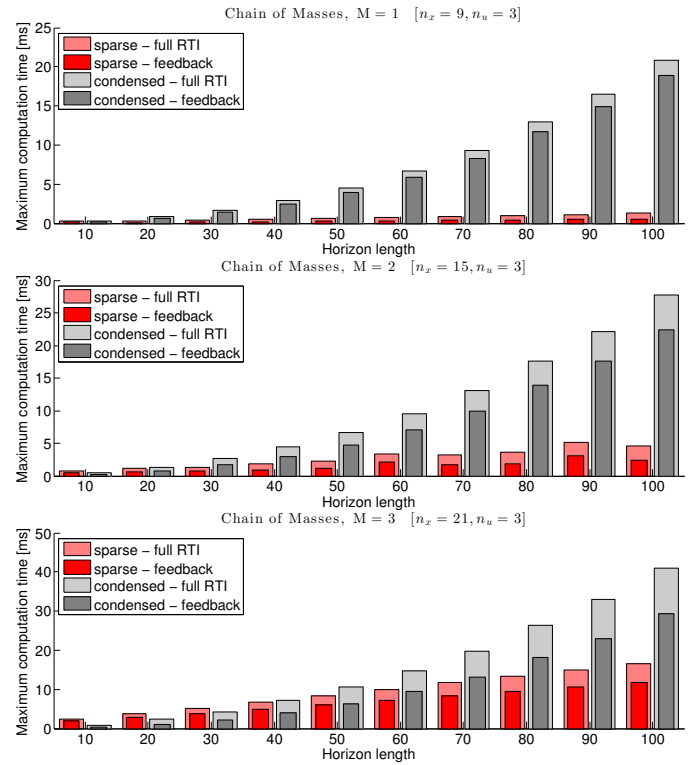


Fig. 2. Maximum computation time benchmark for three chain-of-masses test cases

A similar pattern can be observed when regarding maximum computation times over all simulation steps in Figure 2. For a small to medium number of states, the sparse approach dominates already on relatively short horizon lengths, both in the feedback phase and (thus even more) in the total iteration time. Obviously the computational efforts for integration and condensing are problem-data independent and thus the relative gap between both approaches decreases a bit for long horizons in the consideration of maximum computation times, as observed in the test case of $M = 3$.

The rather large gap between average and maximum computation times of qpDUNES can partly be explained by the wrong initialization of the QP solver after the chain is deflected; already excluding the first iteration from considerations lead to significantly shorter maximum computation times. Obviously the active set method nature of the dual Newton strategy is also reflected here.

While the first benchmark problem is purely academic (but well scalable), we use a real-world motivated second benchmark problem. Nonlinear MPC is used to prevent the occurrence of surge in centrifugal compressors (see Cortinovis et al. (2012) for details). Centrifugal compressors are widely used in gas extraction plants or gas pipelines to extract and transport natural gas from the source to the consumer. As compressing is an energy-intensive process, efficient compressor operation is desirable. This means to operate them at working points that are close to surge, an instable system state that can cause severe damage to the compressor and piping system. We describe the compressor by a nonlinear ODE model similar to the one presented in Cortinovis et al. (2012). It comprises $n_x = 6$ differential states and $n_u = 2$ control inputs: the opening of the recycle

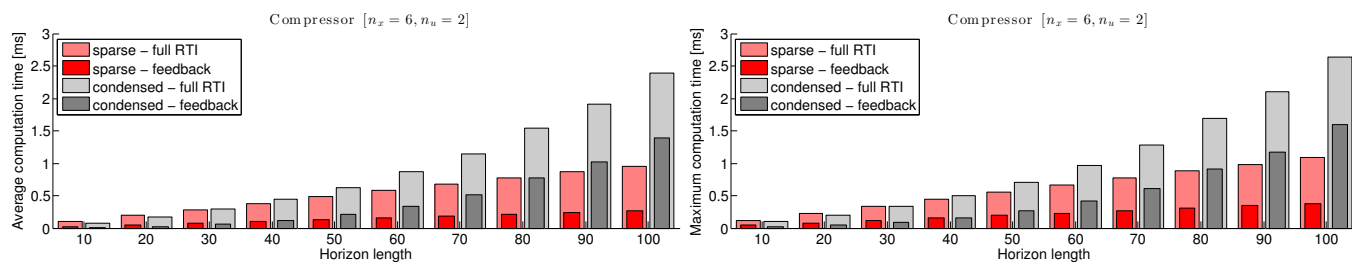


Fig. 3. Average and maximum computation time benchmark for the compressor test case

valve as well as the torque of the compressor's drive (which are subject to physical limitations). Our nonlinear MPC aims at tracking a given operating point in the event of a simulated sudden closure of the compressor's outlet valve.

The anti-surge controller is running at a sampling time of 25 ms on prediction horizons of length N as given in Figure 3. The sparse approach performs competitively on all considered horizon lengths, tying for horizon lengths around $N = 30$. For longer prediction horizons significant computational savings can be achieved applying the sparse approach. Due to the practical relevance we also include maximum computation times in the right part of Figure 3, which essentially confirm the just-made observations.

6. CONCLUSIONS AND FUTURE WORK

We presented a new quadratic programming strategy for an efficient solution of sparse quadratic subproblems in and nonlinear MPC and MHE of long horizon problems. Numerical results on several test cases showed the competitiveness and the potential of this approach in comparison with existing, already well-tuned methods. Currently ongoing research addresses parallelization aspects and code-generation of the linear algebra routines.

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