

Interior Point Methods applied to Quadratic Programming*

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Abstract—Describe in a few sentences what the paper is about and why it is interesting to read it.

I. INTRODUCTION

Notation. Throughout this paper, we denote the set of all real numbers as \mathbb{R} and the set of all natural numbers as \mathbb{N} , respectively \mathbb{N}_0 if zero is included. Further we denote the Jacobian of a function $f : \mathbb{R}^n \rightarrow \mathbb{R}^m$, differentiated along direction v , evaluated at point $x \in \mathbb{R}^n$ as $J_v f(x)$. The index v is omitted, if it is contextually clear. For a collection of scalar valued functions $h_i : \mathbb{R}^n \rightarrow \mathbb{R}, i = 1, \dots, m$ with the same domain, we define

$$h(x) := \begin{pmatrix} h_1(x) \\ \vdots \\ h_m(x) \end{pmatrix}$$

for the stacked values of all h_i evaluated at $x \in \mathbb{R}^n$.

II. PROBLEM STATEMENT AND BACKGROUND

For theoretical discussions, we consider the convex constrained optimization problem

$$\begin{aligned} & \underset{x}{\text{minimize}} && f_0(x) \\ & \text{subject to} && f_i(x) \leq 0, \quad i = 1, \dots, m. \\ & && A_{\text{eq}} x = b_{\text{eq}}. \end{aligned} \quad (1)$$

with $f_0 : \mathbb{R}^n \rightarrow \mathbb{R}$ convex and twice differentiable, $f_i : \mathbb{R}^n \rightarrow \mathbb{R}$ for $i = 1, \dots, m$ convex and differentiable, $A_{\text{eq}} \in \mathbb{R}^{n \times p}$, $b_{\text{eq}} \in \mathbb{R}^p$ with equality and inequality constraints. For such an optimization problem, we call its Lagrangian $L : \mathbb{R}^n \times \mathbb{R}^m \times \mathbb{R}^p \rightarrow \mathbb{R}$ with

$$L(x, \lambda, \nu) = f_0(x) + \lambda^T f(x) + \nu^T (A_{\text{eq}} x - b_{\text{eq}}).$$

Further, we denote its dual problem by

$$\begin{aligned} & \underset{(\lambda, \nu)}{\text{maximize}} && g(\lambda, \nu) \\ & \text{subject to} && \lambda \geq 0, \nu \in \mathbb{R}^p \end{aligned} \quad (2)$$

with

$$g(\lambda, \nu) = \inf_{x \in \mathbb{R}^n} L(x, \lambda, \nu).$$

Moreover, we give a MATLAB-implementation of a primal-dual interiorpoint method for convex quadratic optimization problems. Quadratic problems are a subclass of

(1) and denote as

$$\begin{aligned} & \underset{x}{\text{minimize}} && f_0(x) = \frac{1}{2} x^T Q x + c^T x \\ & \text{subject to} && A_{\text{in}} x - b_{\text{in}} \leq 0, \quad A_{\text{in}} \in \mathbb{R}^{m \times n}, b_{\text{in}} \in \mathbb{R}^m \\ & && A_{\text{eq}} x - b_{\text{eq}} = 0, \quad A_{\text{eq}} \in \mathbb{R}^{p \times n}, b_{\text{eq}} \in \mathbb{R}^p \end{aligned} \quad (3)$$

with matrices $0 \prec Q \in \mathbb{R}^{n \times n}, c \in \mathbb{R}^n$.

III. MAIN RESULTS

A. Newton's Method

Newton's method is an iterative process to solve nonlinear equation systems

$$F(x) = 0 \quad (4)$$

for a differentiable map $F : \mathbb{R}^n \rightarrow \mathbb{R}^m$. The algorithm works as follows: At a given point x_k , the zero of the linear approximation of F around x_k is computed. This point is chosen as the next iterate x_{k+1} . In particular, a linear approximation of F in x_k is defined as

$$L(x) := F(x_k) + JF(x_k)(x - x_k) \text{ for } x \in \mathbb{R}^n, \quad (5)$$

where $JF(x_k)$ is the Jacobian of F at the point x_k . If $JF(x_k)$ is invertible, the point \tilde{x} with $L(\tilde{x}) = 0$ is exactly the solution of the linear equation $JF(x_k)x = -F(x_k)$. Technical conditions and proofs about convergence rates of Newton's method can be found in [1]. The procedure executing a Newton search is summarized in (1).

Algorithm 1: Newton's Method

Result: \tilde{x} , approximate solution of nonlinear system of equalities $F(x) = 0$, residual tolerance

$\epsilon_{\text{res}} > 0$, cauchy-tolerance $\epsilon_c > 0$

Data: Function $F : \mathbb{R}^n \rightarrow \mathbb{R}^n$, initial point x_0

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while  $\|x - x_{\text{last}}\| \geq \epsilon_c$  or  $\|F(x)\| \geq \epsilon_{\text{res}}$  do
    compute Newton direction  $\Delta x$  by solving
         $JF(x)\Delta x = -F(x)$ ;
    remember last iteration for checking term. crit.
         $x_{\text{last}} = x$ ;
    update current point by  $x = x + \Delta x$ ;
end
return  $\tilde{x} = x$ ;

```

Remark 1: The residual and the cauchy-criterion for termination should be combined for the newton method. Easy examples are known, where one of the criteria is satisfied even though the current iteration is far from the optimal point. For details, see [1]. For theoretical reasoning, or if $\nabla^2 f(x)^{-1}$ can be used explicitly, one can also use the

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decrease of $\lambda^2 = \nabla f(x)^T \nabla^2 f(x)^{-1} \nabla f(x)$ (so called newton-decrement) under a certain tolerance.

For the purpose of optimizing a convex, twice differentiable objective function f_0 we want to find a zero of the gradient ∇f_0 . Therefore we can apply the Newton Method to solve the non-linear equation

$$F(x) := \begin{pmatrix} \nabla f_0(x) \\ g(x) \end{pmatrix} = 0.$$

By convexity, satisfying $\nabla f_0(x^*) = 0$ is not only necessary, but also sufficient for x^* to be a global minimum of f_0 .

B. Concept of Barrier Methods

Convex optimization problems with no inequality constraints can be solved efficiently by using Newton's method. If inequality constraints are involved, Newton's method can not guarantee feasibility of a found solution. Hence it is desirable, to transform an inequality-constrained optimization problem into one, that is only equality-constrained. Therefore, we move the inequality constraints implicitly to the objective function. A simple and precise way to do this, would be to evaluate an indicator function

$$I_-(x) := \begin{cases} 0 & \text{for } u \neq 0 \\ \infty & \text{for } u > 0 \end{cases} \quad (6)$$

on the values of the inequality constraints $f_i, i = 1, \dots, m$. We obtain a problem in the following shape

$$\begin{aligned} \underset{x}{\text{minimize}} \quad & f_0(x) + \sum_{i=1}^m I_-(f_i(x)) \\ \text{subject to} \quad & A_{\text{eq}}x - b_{\text{eq}} = 0. \end{aligned} \quad (7)$$

This problem is equivalent to (1), since it yields an objective value of $+\infty$ for every infeasible point while it is the same problem for every feasible one. We now have a formulation without inequality constraints. However, it is clearly neither convex nor continuous and hence not differentiable. Since we need these properties to solve the optimization problem computationally, we approximate the indicator function I_- by the function

$$\hat{I}_-(u) = \begin{cases} \frac{1}{t} \log(-u) & \text{for } u < 0, \\ \infty & \text{for } u \geq 0, \end{cases} \quad (8)$$

The parameter $t > 0$ sets the approximation's accuracy. A higher value for t results in a better approximation of the indicator function. By replacing the indicator function by \hat{I}_- , we obtain an approximation

$$\begin{aligned} \underset{x}{\text{minimize}} \quad & f_0(x) - \sum_{i=1}^m \frac{1}{t} \log(-f_i(x)) \\ \text{subject to} \quad & A_{\text{eq}}x - b_{\text{eq}} = 0 \end{aligned} \quad (9)$$

of problem (1). Throughout this paper, we denote its Lagrangian by $L_t : \mathbb{R}^n \times \mathbb{R}^p \rightarrow \mathbb{R}$.

Note that $\frac{1}{t} \log(-u)$ is convex, increasing in u , and differentiable on the feasible set. Hence the entire function

$\sum_{i=1}^m \hat{I}_-(f_i(x))$ is convex and (9) is a convex Problem with differentiable objective function. These properties allow us to solve (9) computationally. We call an optimal point $x^*(t)$ of (9) with parameter t a central point and a solution to its dual problem $(\lambda^*(t), \nu^*(t))$ a dual central point. The set of (dual) solutions of (9) for all $t > 0$ we call the (dual) central path. Since for points x with $f_i(x) = 0$ for any $x \in \{1, \dots, m\}$, the objective of (9) is ∞ , all central points are in the interior of the set, satisfying the inequality constraints of (1). Thus this framework is named interior point method. One can show, that solutions $(x^*(t), \lambda^*(t), \nu^*(t))$ of (9) converge to the solution (x^*, λ^*, ν^*) of (1) for $t \rightarrow 0$. The proof can be found in [2].

C. Measure for the Approximation's quality

An immediately arising question is which conclusions about the solution (x^*, λ^*, ν^*) of (1) can be drawn from knowing a solution of (9) for a certain $t > 0$ about the value $f_0(x^*(t))$ of a central point $x^*(t)$, compared with the optimal value p^* of the original problem. For compactness, we denote the barrier term of the approximated problem as

$$\phi(x) = - \sum_{i=1}^m \log(-f_i(x)),$$

with its Jacobian and Hessian being

$$\nabla \phi(x) = \sum_{i=1}^m \frac{1}{-f_i(x)} \nabla f_i(x),$$

$$\nabla^2 \phi(x) = \sum_{i=1}^m \frac{1}{f_i(x)^2} \nabla f_i(x) \nabla f_i(x)^T + \sum_{i=1}^m \frac{1}{-f_i(x)} \nabla^2 f_i(x).$$

For the sake of simplifying notation, throughout this section we consider the problem

$$\begin{aligned} \underset{x}{\text{minimize}} \quad & t f_0(x) + \phi(x) \\ \text{subject to} \quad & A_{\text{eq}}x = b_{\text{eq}}. \end{aligned} \quad (10)$$

that is obtained by multiplying the objective in (9) with $t > 0$. The original and the obtained problem are equivalent. Any arbitrary $x^*(t)$ a strictly feasible point of (1). Since $x^*(t)$ solves (10), there exists $\hat{\nu} \in \mathbb{R}^p$, such that

TODO!check consistency

$$\begin{aligned} \nabla L_t(x^*(t), \hat{\nu}) &= t \nabla f_0(x^*(t)) + \nabla \phi(x^*(t)) + A_{\text{eq}}^T \hat{\nu} \\ &= t \nabla f_0(x^*(t)) \end{aligned} \quad (11)$$

$$+ \sum_{i=1}^m \frac{1}{-f_i(x^*(t))} \nabla f_i(x^*(t)) + A_{\text{eq}}^T \hat{\nu}.$$

$$= 0 \quad (12)$$

$$= 0 \quad (13)$$

holds. Note that the Lagrangian only depends on $(x, \hat{\nu})$, since there are no explicit inequality constraints involved. We keep in mind, that $x^*(t)$ minimizes (9). Using this insight, we know that there exists a dual feasible point $(x^*(t), \lambda^*(t), \nu^*(t))$ of the original problem (1). In particular, we choose

$$\lambda^*(t) = - \frac{1}{t f_i(x^*(t))} \text{ for } i = 1, \dots, m, \quad \nu^*(t) = \frac{\hat{\nu}}{t}.$$

Here, $\lambda^*(t) > 0$ follows from $f_i(x^*) < 0$ for all $i = 1, \dots, m$ since x^* is strictly feasible.

Note that (13) is the derivative of the Lagrangian

$$L(x, \lambda, \nu) = f_0(x) + \sum_{i=1}^m \lambda_i^*(t) f_i(x) + \nu^*(t)^T (A_{\text{eq}} x^*(t) - b_{\text{eq}})$$

divided by $t > 0$ of the original problem. The Lagrangian is convex in the first coordinate, hence we infer that $x^*(t)$ minimizes the Lagrangian of the original problem for any fixed (λ, ν) . For the dual function of the original problem, we obtain

$$\begin{aligned} g(\lambda^*(t), \nu^*(t)) &= f_0(x^*(t)) + \sum_{i=1}^m \lambda_i^*(t) f_i(x^*(t)) \\ &\quad + \nu^*(t)^T (A_{\text{eq}} x^*(t) - b_{\text{eq}}) \\ &= f_0(x^*(t)) - \frac{m}{t}. \end{aligned} \quad (14)$$

The second of the three summands adds up to $m \cdot 1$, because of the particular choice of $\lambda^*(t)$, fractions cancel out. The last summand equals zero, since $A_{\text{eq}} x^*(t) - b_{\text{eq}} = 0$.

By weak duality, this means that the optimum $x^*(t)$ of the approximated problem (9) has an objective value $f_0(x^*(t))$ that is maximally larger by $\frac{m}{t}$ (and hence worse) than the real optimal value p^* of the original problem. Thus, one can theoretically force a desired bound on the suboptimality $\epsilon > 0$ by choosing t large enough, in particular $t := \frac{m}{\epsilon}$. However, just solving (9) with a large choice of t does not work out in general, since numerical issues can make convergence of Newton's Method dependent on the choice of the initial point x_0 .

D. Algorithmic Use of the Barrier Concept

As already mentioned in section III-C, one can not solve (9) generally without a good guess of the initial value x_0 . So how to make use of the barrier concept? The idea of interior methods is, to find points along the problem's central path. Two methods are introduced in the following. Emphasis of the explanations as well as the implementation in MATLAB will be on the Primal-Dual Interior Point Method.

1) *Barrier Method*: As mentioned before, for large t a good initial point x_0 , meaning an initial point that is not far away from the actual minimum of (1), is crucial for avoiding large numerical errors. This can be achieved by starting with optimization of (9) for small $t = t_1$, which leads to a rather bad approximation of the original problem, but also to better numerical behavior. After finding $x^*(t_1)$ via Newton's method, t is increased to $t = t_2 > t_1$ by a certain rate and (9) is solved again with parameter $t = t_2$, with choice $x_0 = x^*(t_1)$ for the initial point.

We call finding the minimum $x^*(t)$ of (9) the centering step or outer iteration, while we call a single Newton step inside the centering step an inner iteration.

2) *Primal-Dual Interior Point Method*: Like the previously introduced algorithm, the Primal-Dual Interior Point method uses the barrier concept to handle inequality constraints. It is motivated by the following idea: Since the points generated by each outer iteration converge to the

Algorithm 2: Barrier Method with full Newton search

Result: $x^*(t)$, approximate solution of (1) with

$$f_0(x^*(t)) - p^* < \frac{m}{t}$$

initialization: Matrices $0 \prec Q \in \mathbb{R}^{n \times n}$, $c \in \mathbb{R}^n$.

defining the objective function, matrices

$A_{\text{eq}} \in \mathbb{R}^{m \times n}$, $b_{\text{eq}} \in \mathbb{R}^m$, $A_{\text{in}} \in \mathbb{R}^{m \times n}$, $b_{\text{in}} \in \mathbb{R}^m$ defining

constraints, initial point x , initial approximation

parameter $t > 0$, rate for increasing approx. param.

$\mu > 1$ tolerance ϵ ;

while $\frac{m}{t} \geq \epsilon$ **do**

 Compute $x^*(t)$ by solving (9) via Newton's Method, starting at x ;

 Update $x := x^*(t)$;

 Increase t by $t := \mu t$

end

desired optimum on the central path, one does not gain much advantage by computing the central points with a high level of accuracy. This results in many newton-steps being computed, without improving the convergence towards the optimum value of (1). Hence, it would be useful to reduce the accuracy of each outer iteration as much as possible, without losing convergence to the optimum. Therefore, in this method only one newton step will be computed for each parameter t in the approximated problem (9). Furthermore, this Newton step is computed differently. While in the barrier method with full newton search, the search directions are computed only considering the primal problem, the Primal-Dual Method also takes the dual problem of (9) into account. In particular Newton's method is applied to a system of residual terms, which all have to equal zero by the modified KKT-conditions, here presented like in [2].

Theorem 1 (Modified KKT-Conditions): For a convex Optimization Problem with a logarithmic barrier function (9), the following conditions on a primal-dual point $(x, \lambda, \nu) \in \mathbb{R}^n \times \mathbb{R}^m \times \mathbb{R}^p$ are necessary and sufficient for x being a solution to the primal problem and (λ, ν) being a solution to the dual problem:

$$f_i(x) \leq 0, \quad \text{for } i = 1, \dots, m \quad (15a)$$

$$A_{\text{eq}} x - b_{\text{eq}} = 0 \quad (15b)$$

$$\lambda_i \geq 0, \quad \text{for } i = 1, \dots, m \quad (15c)$$

$$-\lambda_i f_i(x) = \frac{1}{t}, \quad \text{for } i = 0, \dots, m \quad (15d)$$

$$\nabla f_0(x) + \sum_{i=1}^m \lambda_i \nabla f_i(x) + \sum_{i=1}^p \nu_i \nabla h_i(x) = 0. \quad (15e)$$

Stacked in one vector, this yields the system of equalities

$$\begin{aligned} r_\mu(x, \lambda, \nu) &= \begin{pmatrix} r_{\text{dual}} \\ r_{\text{cent}} \\ r_{\text{pri}} \end{pmatrix} \\ &= \begin{pmatrix} \nabla f_0(x) + Jf(x)^T \lambda + A_{\text{eq}}^T \nu \\ -\text{diag}(\lambda)f(x) - \mu \mathbb{1} \\ A_{\text{eq}}x - b_{\text{eq}} \end{pmatrix} \stackrel{!}{=} 0. \end{aligned} \quad (16)$$

to apply Newton's method on. For formulation of the linear Newton equality, we also compute the jacobian

$$\begin{aligned} J_{(x, \lambda, \nu)} r_\mu(x, \lambda, \nu) & \quad (17) \\ &= \underbrace{\begin{pmatrix} \nabla^2 f_0(x) + \sum_{i=1}^m \lambda_i \nabla^2 f_i(x) & Jf(x) & A_{\text{eq}}^T \\ -\text{diag}(\lambda)Jf(x) & -\text{diag}(f(x)) & 0 \\ A_{\text{eq}} & 0 & 0 \end{pmatrix}}_{:=M_{\text{KKT}}} \end{aligned} \quad (18)$$

of the residual and refer to it as M_{KKT} . Consequently, the Newton equality for finding the search direction $(\Delta x, \Delta \lambda, \Delta \nu)$ in each newton step is obtained by solving the linear equation

$$M_{\text{KKT}} \begin{pmatrix} \Delta x \\ \Delta \lambda \\ \Delta \nu \end{pmatrix} = b_{\text{KKT}} \quad (19)$$

with $b_{\text{KKT}} = -r_\mu(x, \lambda, \nu)$.

Unfortunately, adding the obtained step direction $(\Delta x, \Delta \lambda, \Delta \nu)$ to (x, λ, ν) , does not in general yield a feasible point. Therefore we compute a suitable step-size s^* via a backtracking-linesearch, such that a certain decrease of the residual and feasibility is guaranteed for the next iteration point

$$\begin{pmatrix} x^+ \\ \lambda^+ \\ \nu^+ \end{pmatrix} = \begin{pmatrix} x \\ \lambda \\ \nu \end{pmatrix} + s^* \begin{pmatrix} \Delta x \\ \Delta \lambda \\ \Delta \nu \end{pmatrix}.$$

The detailed procedure of the backtracking linesearch is displayed in Algorithm 3.

Finally, we can present the entire algorithm of the Primal-Dual Method.

Remark 2: If a strictly feasible primal variable $x \in \mathbb{R}^n$ is known, $\lambda = -1/f_i(x) \geq 0, \nu = 0$ is always a valid choice for the initial dual variables.

Especially when results with high accuracy are needed, the Primal-Dual Method allows to omit a lot of newton steps, that would be have computed in the Barrier Method. For such problems, the Primal-Dual Method provides better performance.

Remark 3: The concept of using a barrier method to approximate inequality constraints can also be used for solving optimization problems involving generalized inequalities. Therefore, the barrier function and constraints on the dual problem have to be adjusted. Doing this, the class of problems that apply the usage of barrier methods is widely enlarged. For example it can be used for solving linear matrix inequalities or single order cone problems.

Algorithm 3: Backtracking linesearch

Result: Stepsize s^* , s.t. $\lambda^+ > 0, f(x^+) < 0$ and r_μ decreases by certain amount.

Data: Problem matrices, current x, λ, ν , Newton direction $\Delta x, \Delta \lambda, \Delta \nu$, barrier parameter μ , backtracking parameters $\alpha \geq 0, \beta \in (0, 1)$.

Initial step-size set

$$s_{\text{max}} = \min\{1, \min_{i|\Delta \lambda_i < 0} -\lambda_i / \Delta \lambda_i\}$$

compute $r_\mu(x, \lambda, \nu)$;

$s = s_{\text{max}}$;

$found = false$;

while $found == false$ **do**

 set $s = \beta s$;

 compute (x^+, λ^+, ν^+) ;

 compute $r_\mu(x^+, \lambda^+, \nu^+)$ and $f(x^+)$;

if $f(x^+) < 0$ **and**

$\|r_\mu(x^+, \lambda^+, \nu^+)\| \leq (1 - \alpha s) \|r_\mu(x, \lambda, \nu)\|$ **then**

$found = true$

end

end

Algorithm 4: Primal-Dual Interior Point Method

Result: approximate optimizer \hat{x}^* , approx. opt. value \hat{p}^* , approx. dual optimizer $(\hat{\lambda}^*, \hat{\nu}^*)$, surrogate duality gap $\hat{\eta}^*$ as measure of optimality

Data: Problem matrices, primal-dual initial point

(x, λ, ν) with $f_i(x) < 0$ for all $i = 1, \dots, m$,

$\lambda > 0, \nu \in \mathbb{R}^p$ (initial point strictly feasible),

reduction factor $\gamma \in (0, 1)$, tolerances

$$\epsilon_{\text{feas}}, \epsilon_{\text{opt}} > 0$$

Initialization;

determine problem dimensions n, m, p ;

set $found = false$;

while $found == false$ **do**

 compute surrogate duality gap: $\hat{\eta} = -f(x)^T \lambda$;

 compute KKT residual vector $r_\mu(x, \lambda, \nu)$ via (16);

 compute search direction $(\Delta x, \Delta \lambda, \Delta \nu)$ by

 solving (19);

 determine suitable step size s via backtracking algorithm 3;

 update current primal and dual points:

$$(x, \lambda, \nu) = (x, \lambda, \nu) + (s \Delta x, s \Delta \lambda, s \Delta \nu);$$

end

return $\hat{x}^* = x, \hat{p}^* = f_0(\hat{x}^*), \hat{\lambda}^* = \lambda, \hat{\nu}^* = \nu, \hat{\eta}^* = \hat{\eta}$;

Further, a higher speed of convergence can be shown for the application on some special classes of problems, such as quadratic problems or single order cone problems (SOCs). Here the Primal-Dual Method can perform faster than with linear convergence.

E. How to find a feasible initial point

The Algorithms 2 and 4 both need a strictly feasible initial point to start. Since such a point is in general not trivial to find, one can formulate the search for the initial point as another convex optimization problem, that is easier to solve than the original one. For problem (1), one way to implement this, is solving

$$\begin{aligned} & \underset{x}{\text{minimize}} && s \\ & \text{subject to} && f_i(x) \leq s, \quad i = 1, \dots, m \\ & && A_{\text{eq}}x - b_{\text{eq}} = 0, \end{aligned} \quad (20)$$

via Newton's method. If a point with optimal value strictly smaller than zero for (20) is found, then this point is strictly feasible. Solving such a first, more simple problem is called a Phase I problem. More examples of such problems can be found in [2].

F. Complexity Analysis for the Barrier Method

Emphasis of this article is on implementation and idea of the algorithms, so we treat complexity analysis only by presenting results without proves. We keep this restricted to the barrier method with full newton search. We discuss the time complexity of the barrier method, meaning the total number of newton steps needed to solve (1). An upper bound of these iterations can be proven for problems with objectives that are self-concordant. While linear and quadratic functions satisfy selfconcordance in general, any other convex optimization problem can be rewritten as a self-concordant one, so this condition is not very restrictive. The upper bound

$$\frac{f(x) - p^*}{\gamma} + c \quad (21)$$

on the maximal number of newton iterations that is needed to get a newton decrement (see remark 1) smaller than ϵ_{nt} , where c depends on ϵ_{nt} by $\log_2 \log_2(1/\epsilon_{\text{nt}})$, p^* is the primal problem's optimal value and γ depends on choice of the backtracking parameters α, β with

$$\frac{1}{\gamma} = \frac{20 - 8\alpha}{\alpha\beta(1 - 2\alpha)^2}.$$

The derivation of this bound is shown in [2], section 9.

One can further show that this bound holds uniformly for any parameter t for all problems (9). Since there are exactly

$$\left\lceil \frac{\log(m/\epsilon t_0)}{\log \mu} \right\rceil$$

outer steps necessary to solve (9) with initial parameter $t = t_0$ and tolerance ϵ , the entire barrier method needs maximally

$$N = \left\lceil \frac{\log(m/\epsilon t_0)}{\log \mu} \right\rceil \left(\frac{m(\mu - 1 - \log \mu)}{\gamma} + c \right)$$

inner newton iterations, where m denotes the number of inequality constraints on (9). to yield a result with a sub-optimality of ϵ or smaller. Detailed reasoning can be found in [2], section 11.5.

IV. EXAMPLES

We give an implementation of the Primal-Dual Method (Algorithm 4) in MATLAB, along with two example problems. One with equality constraints, one without.

A. Numerical Examples

1) *Example 1:* The effectivity of the algorithm can be demonstrated with the following explicit example of (1). For the problem defined by

$$\begin{aligned} Q &= 2 \cdot \begin{pmatrix} 1 & 0 \\ 0 & 2 \end{pmatrix}; \quad c = \begin{pmatrix} 1 \\ 2 \end{pmatrix} \\ A_{\text{eq}} &= \begin{pmatrix} 1 & -1 \end{pmatrix}; \quad b_{\text{eq}} = 0; \\ A_{\text{in}} &= \begin{pmatrix} 0 & 1 \end{pmatrix}; \quad b_{\text{eq}} = 10, \end{aligned}$$

we choose backtracking parameters $\alpha = 0.05, \beta = 0.5$ as well as $\gamma = 0.1$ for the reduction of the approximation parameter. Initial points are set to $x_0 = (1, 1)^T, \lambda_0 = 1, \nu_0 = 0$ and tolerances to $\epsilon_{\text{feas}} = \epsilon_{\text{opt}} = 10^{-4}$.

With this setup, the solver terminates after 20 iterations and yields the following results:

$$\begin{aligned} x^* &= \begin{pmatrix} -1.0000 \\ -1.0000 \end{pmatrix}; \\ \lambda^* &= 5.6650 \cdot 10^{-6}; \\ \nu &= 2.7562 \cdot 10^{-6}; \\ f_0(x^*) &= -1.5000; \\ \hat{\eta} &= 6.2315 \cdot 10^{-6}. \end{aligned}$$

2) *Example 2 :* To test algorithm (4) on a problem without equality constraints, we moreover provide a second example, taken from [3]. The problem is defined

$$\begin{aligned} Q &= 2 \cdot \begin{pmatrix} 3 & 1 \\ 1 & 1 \end{pmatrix}; \quad c = \begin{pmatrix} 1 \\ 6 \end{pmatrix}; \\ A_{\text{in}} &= \begin{pmatrix} -2 & -3 \\ 0 & -1 \end{pmatrix}; \quad b_{\text{in}} = \begin{pmatrix} -4 \\ 0 \\ 0 \end{pmatrix}, \end{aligned}$$

while initial values are chosen as $x_0 = (1, 1)^T$ and $\lambda_0 = (0.2, 1, 1)^T$ (like suggested in remark 2). Tolerances and backtracking-parameters are chosen as in Example 1.

With termination after 20 iterations, the solver yields the results

$$x^* = \begin{pmatrix} 0.5000 \\ 1.0000 \end{pmatrix} \quad (22)$$

$$\lambda^* = \begin{pmatrix} 3.0000 \\ 0.0001 \\ 2.812 \cdot 10^{-5} \end{pmatrix} \quad (23)$$

$$f_0(x^*) = 9.2500 \quad (24)$$

V. CONCLUSIONS

Mainly, in this text we introduced the concept of barrier methods for convex optimization problems, which are applied to solve inequality-constraints that include inequality constraints. Therefore we approximate such problems by problems without inequality problems. Solving these problems yields points, that are only suboptimal in the sense of the original problem up to a certain bound. Via analyzing the Lagrangian of the original and the approximative problem, we verified this bound, dependent on the approximation parameter and the number of inequality constraints. The explanation of the barrier concept was followed by an explanation how it can be applied algorithmically by the barrier method. Further, we gave an idea how to find a feasible initial point and also a time complexity bound of the barrier method, without proof. As an improvement, especially when dealing with quadratic problems, the Primal-Dual Method was introduced, where only one newton step is executed before the approximation parameter is changed. Finally, we showed how quadratic programming can be applied on a linear MPC problem with zero terminal constraints by reformulating it as a quadratic problem.

APPENDIX

A. MATLAB-Implementation

In the following, the MATLAB code of the primal-dual interior point method algorithm is presented

Algorithm main function:

```
function [x,fval,lambda,nu,eta] =
    ipquad_pd(Q,c,Aineq,bineq,Aeq,beq,x0,
        lambda0,nu0,gamma,eps_feas,eps_opt,
        ls_alpha,ls_beta)
%IPQUAD_PD Quadratic optimization via
% primal-dual-interior-point-method.
% Convex Quadr. function f(x) = (1/2)x'
% Qx + c'x with linear equality and
% inequality constraints.
% -----
% Input Arguments:
% - Q,c define objective function
%   dim(Q) = nxn, dim c = nx1
% - Aineq, bineq define inequality
%   constraints Aineq*x <= bineq
%   dim(Aineq) = mxn, dim(bineq) = mx1
% - Aeq, beq define inequality
%   constraints Aeq*x == beq
%   dim(Aeq) = pxn, dim(beq) = px1
% - x0 init. value for the primal
%   problem. lambda0, nu0. lambda0 >= 0.
%   Aineq*x0 <= bineq.
%   initial values for the dual problem
% - gamma is a reduction factor for
%   reducing the barrier weight
%   mu_barrier
%   in each iteration. gamma in (0,1)
% - eps_feas > 0 specifies the tolerance
%   for the 2norms of the primal and
%   dual residual
% - eps_opt > 0 specifies a tolerance on
%   the surrogate duality gap
% - ls_alpha, ls_beta are parameters for
%   the backtracking linesearch,
%   performed in each iteration. Typical
%   choices: ls_alpha in [0.01,0.1].
%   ls_beta in [0.3,0.8]
% Outputs:
% - x is the best approximation on the
%   primal optimum found by the algor.
% - lambda, nu are the best
%   approximation on the dual optimum
% - fval is the objective's value eval.
%   at x
% - eta is the surrogate duality gap at
%   lambda, x
% -----
% Created: 24.06.20, Daniel Bergmann
% -----
```

```

33 x = x0;
34 lambda = lambda0;
35 nu = nu0;
36
37 % initialize dimensions
38 n = size(Q,1);
39 m = size(Aineq,1);
40 p = size(Aeq,1);
41
42 found = 0;
43 count = 0;
44
45 while found == 0
46     % compute surrogate duality gap
47     eta = -(Aineq*x - bineq)'*lambda;
48     mu_barrier = gamma*eta/m;
49     % Compute KKT residual vector
50     r_mu = res_kkt(x,lambda,nu,Q,c,Aineq, bineq, Aeq, beq, mu_barrier);
51     r_dual = r_mu(1:m);
52     % r_cent = r_mu((m+1):(m+p));
53     r_pri = r_mu((m+p+1):(m+p+n));
54
55     if norm(r_pri) <= eps_feas && norm(r_dual) <= eps_feas && eta <= eps_opt
56         found=1;
57         break;
58     else
59         % update barrier weighting
60         % parameter mu_barrier
61         mu_barrier = gamma*eta/m;
62         % compute search vector
63         [x, lambda, nu] = newtonquad_pd(Q, c, Aineq, bineq, Aeq, beq, ls_alpha, ls_beta, x, lambda, nu, mu_barrier);
64     end
65     count = count+1;
66     disp(['Iteration No. ', num2str(count), ', current norm of residual: ', num2str(norm([x; lambda; nu])), ', eta = ', num2str(eta)])
67 end
68
69 % evaluate obj. function at found x
70 fval = 0.5.*x'*Q*x + c'*x;
71 end
72
73 Newton step including line-search:
74
75 function [x_new, lambda_new, nu_new] = newtonquad_pd(Q, c, Aineq, bineq, Aeq, beq, ls_alpha, ls_beta, x, lambda, nu, mu_barrier)
76 %NEWTONQUAD_PD Computes search direction for pd-ip-algorithm via Newton's
77
78 method and step size
79 %via backtracking line search
80 % Based on a given primal-dual point x, lambda,nu, this functions returns new points
81 % x_new, lambda_new, nu_new with smaller kkt-residual.
82 % First, a search direction is determined by applying newton's method to
83 % the nonlinear equation system r = 0 with r the residual of the
84 % kkt-conditions, second a suitable step-size is determined via a
85 % backtracking linesearch
86 % -----
87 % Input Arguments
88 % - mu_barrier is the current weight on the barrier function
89 % - for other input arguments, see comments in ipquad_pd.m
90 % Outputs
91 % - x_new, lambda_new, nu_new are the new primal-dual point after adding
92 % the search direction obtained by newton's method, multiplied by the
93 % step size obtained by backtracking line search to the old primal-dual
94 % point
95 % -----
96 % Created: 24.06.20, Daniel Bergmann
97 % -----
98
99 % initialize dimensions
100 n = size(Q,1);
101 m = size(Aineq,1);
102 p = size(Aeq,1);
103
104 % Define Matrices for KKT-equality
105 M_kkt*deltar = b_kkt
106 if ~isempty(Aeq)
107     M_kkt = [Q
108             Aineq'
109             -diag(lambda)*Aineq
110             Aineq*x-bineq
111             Aeq
112             zeros(m)
113             zeros(p,
114             zeros(p,p) ];
115 else
116     M_kkt = [Q
117             Aineq'
118             -diag(lambda)*Aineq
119             Aineq*x-bineq];
120 end
121 b_kkt = -res_kkt(x,lambda,nu,Q,c,Aineq, bineq, Aeq, beq, mu_barrier);

```

```

39
40 % Solve KKT-equality to get search
    direction
41 deltaxln = M_kkt\b_kkt;
42
43
44 deltax = deltaxln(1:n);
45 deltalambda = deltaxln((n+1):(n+m));
46 deltanu = deltaxln((n+m+1):(n+m+p));
47
48 % Perform Backtracking linesearch for
    determining suitable step size
49
50 % compute maximal step size smax
51 lambdaquot = [];
52 for i = 1:length(lambda)
53     if deltalambda(i) < 0
54         lambdaquot = [lambdaquot, -
            lambda(i)/deltalambda(i)];
55     end
56 end
57
58 smax = min([1 lambdaquot]);
59
60 % Backtracking-Linesearch
61 % Typical Parameter choices:
62 % alpha in [0.01,0.1]; beta in [0.3,0.8]
63 s = 0.99*smax; %S4 0.99 smax??
64 found = 0;
65 while found == 0
66     s = s*ls_beta;
67     x_new = x + s.*deltax;
68     lambda_new = lambda + s.*deltalambda
        ;
69     nu_new = nu + s.*deltanu;
70     r_new = res_kkt(x_new,lambda_new,
        nu_new,Q,c,Aineq,bineq,Aeq,beq,
        mu_barrier);
71     r_old = res_kkt(x,lambda,nu,Q,c,
        Aineq,bineq,Aeq,beq,mu_barrier);
72     if all(Aineq*x_new - bineq < 0) && (
        norm(r_new) <= (1-ls_alpha*s)*
        norm(r_old))
73         found = 1;
74     end
75 end
76 disp(['Backtracking search yields step
    size s = ',num2str(s)])
77 end

```

Computing KKT residual vector from problem matrices and current points:

```

1 function r = res_kkt(x,lambda,nu,Q,c,
    Aineq,bineq,Aeq,beq,mu_barrier)
2 %RES_KKT compute the current KKT-
    residual of a quadratic
3 %optimization problem inside a primal-

```

```

    dual interior point method.
4 % For the quadr. convex opt. problem
    defined by Q,c,Aineq,bineq,Aeq,beq
5 % (for details see comments on ipquad_pd
    .m) and a current primal-dual point
6 % (x,lambda,nu), this function computes
    the current KKT-residual.
7 %
8 % -----
9 % Input Arguments: for detailed
    explanation see comments on ipquad_pd
    .m and
10 % newtonquad_pd.m.
11 % -----
12 % Created: 24.06.20, Daniel Bergmann
13 % -----
14
15 m = size(Aineq,1);
16 if ~isempty(Aeq)
17     % if there are equality constraints
18     r = ...
19         [Q*x + c + Aineq'*lambda + Aeq'*
20             nu;...
21             -diag(lambda)*(Aineq*x - bineq)
22             - mu_barrier.*ones(m,1);...
23             Aeq*x - beq];
24 else
25     % If there are no equality
26     constraints
27     r = [Q*x + c + Aineq'*lambda;...
28         -diag(lambda)*(Aineq*x - bineq)
29         - mu_barrier.*ones(m,1)];
30 end
31 end

```

B. Model Predictive Control

To illustrate the practical relevance of quadratic programming in e.g. control tasks, we here present the application of interior methods on a Model Predictive Control (MPC) problem. More precisely, a MPC of a linear system with zero terminal constraint (ZTC). We consider the discrete time linear system

$$x_{k+1} = A_d x_k + B_d u_k, \quad (25)$$

where $x_k \in \mathbb{R}^n$, $u_k \in \mathbb{R}^m$ for all $k \in \mathbb{N}_0$, with the constraints that $\|u_k\|_{\max} \leq l_u$ and $\|x_k\|_{\max} \leq l_x$ for all steps $k \in \mathbb{N}_0$. Further we assume that (25) has an equilibrium in $x = 0$ and the current state x_0 is given. The maximum-norm is defined as the maximum absolute value over all entries of the vector, $\|x\|_{\max} = \max_{1 \leq i \leq n} |x_i|$. Goal is to find a input signal that steers the internal state x to zero, while additionally keeping u as small as possibly. Therefore, we consider the next $N \in \mathbb{N}$ timesteps. We call N the prediction horizon. This leads to optimization of a certain objective function over

all possible predicted steering signals $\bar{u} = (\bar{u}_1, \dots, \bar{u}_{N-1})^T$. We simulate the system for the next N timestep, hence we consider the sequence of states arising from applying a predicted sequence of input signals \bar{u} . The sequence of predicted states we denote as $(\bar{x}_0, \dots, \bar{x}_N)^T$. We choose the quadratic objective function

$$\sum_{k=0}^{N-1} \underbrace{\bar{x}_k^T Q \bar{x}_k}_{=: \|\bar{x}_k\|_Q} + \underbrace{\bar{u}_k^T R \bar{u}_k}_{=: \|\bar{u}_k\|_R} \quad (26)$$

under the condition, that \bar{x}_N , the last state in the predicted time, equals zero, to minimize. The regarding predicted states directly follow from the system dynamics, in particular

$$\bar{x}_{k+1} = A_d \bar{x}_k + B_d \bar{u}_k.$$

After finding an optimal signal \bar{u} we apply \bar{u}_0 in the next time step. After this first step, we again start an optimization over the next N timesteps. So even though the optimization problem is computed considering all signals up to the prediction horizon, the result is only applied for one timestep, before the next optimization is executed.

We can summarize one optimization step as an optimization problem

$$\begin{aligned} & \underset{\bar{u}=(\bar{u}_0, \dots, \bar{u}_{N-1})^T}{\text{minimize}} && \sum_{k=1}^{N-1} \|\bar{x}_k\|_Q + \|\bar{x}_k\|_R \\ & \text{subject to} && \bar{x}_0 = x_0, \\ & && \bar{x}_N = 0, \\ & && \bar{x}_{k+1} = A_d \bar{x}_k + B_d \bar{u}_k \quad \text{for all } k = 0, \dots, N, \\ & && \|\bar{x}_k\| \leq l_x, \|\bar{u}_k\| \leq l_u \quad \text{for all } k = 0, \dots, N. \end{aligned} \quad (27)$$

This problem can be transcribed into the form of (1). We therefore optimize over the whole vector

$$\tilde{x} := \begin{pmatrix} \bar{x} \\ \bar{u} \end{pmatrix}.$$

The objective function (26) then can be written as

$$f_0(\tilde{x}) = \tilde{x}^T H \tilde{x},$$

with the block diagonal matrix

$$H = \text{diag}(\underbrace{Q, \dots, Q}_{N \cdot n \text{ blocks}}, \underbrace{0}_{n \times n}, \underbrace{R, \dots, R}_{N \cdot n \text{ blocks}})$$

We further formulate the constraints at the maximum norms of state and input as $A_{\text{in}} \tilde{x} \leq b_{\text{in}}$, with $b_{\text{in}} = \mathbb{1} \in \mathbb{R}^{N(n+m)+n \times 1}$ and

$$A_{\text{in}} = \begin{pmatrix} I_{n(N+1)} & 0 \\ -I_{n(N+1)} & 0 \\ 0 & I_{N \cdot m} \\ 0 & -I_{N \cdot m} \end{pmatrix}$$

where the indices of the identity-matrix I denote its dimensional size. Rearranging the system dynamics (25) and taking initial condition as well as zero terminal constraint into account can be written as the equality constraints $A_{\text{eq}} \tilde{x} = b_{\text{eq}}$ with $A_{\text{eq}} = \begin{pmatrix} A_{\text{eq}}^x & A_{\text{eq}}^u \end{pmatrix}$, where

$$A_{\text{eq}}^x = \begin{pmatrix} I_{n \times n} & 0 & \dots & \dots & 0 \\ A_d & -I_{n \times n} & \dots & \dots & \vdots \\ 0 & \ddots & \ddots & & 0 \\ \vdots & & \ddots & \ddots & 0 \\ \vdots & \dots & & A_d & -I_{n \times n} \\ 0 & \dots & \dots & 0 & I_{n \times n} \end{pmatrix}$$

$$A_{\text{eq}}^u = \begin{pmatrix} 0 & \dots & 0 \\ B_d & \ddots & \vdots \\ \vdots & \ddots & 0 \\ 0 & \dots & B_d \\ 0 & \dots & 0 \end{pmatrix}.$$

With these matrices, we rewrote (27) as an equivalent problem in shape of (1).

REFERENCES

- [1] Carsten Scherer *Vorlesungsskript Einführung in die Optimierung* 2019: Lehrstuhl für Mathematische Systemtheorie, Universität Stuttgart.
- [2] Stephen Boyd, Lieven Vandenberghe *Convex Optimization* 2004: Cambridge University Press.
- [3] Jack Heider, *Quadratic programming* 2015: online, https://optimization.mccormick.northwestern.edu/index.php/Quadratic_programming.