# NLPQLP: A Fortran Implementation of a Sequential Quadratic Programming Algorithm with Distributed and Non-Monotone Line Search <br> - User's Guide, Version 3.1 - 

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#### Abstract

The Fortran subroutine NLPQLP solves smooth nonlinear programming problems by a sequential quadratic programming (SQP) algorithm. This version is specifically tuned to run under distributed systems controlled by an input parameter $(l)$. In case of computational errors as for example caused by inaccurate function or gradient evaluations, a non-monotone line search is activated. Numerical results are included which show that in case of noisy function values, a significant improvement of the performance is achieved compared to the version with monotone line search. Further stabilization is obtained by performing internal restarts in case of errors when computing the search direction due to inaccurate derivatives. The new version of NLPQLP successfully solves more than $90 \%$ of our 306 test examples subject to a stopping tolerance of $10^{-7}$, although at most two digits in function values are correct in the worst case and although numerical differentiation leads to additional truncation errors. In addition, automated initial and periodic scaling with restarts is implemented. The usage of the code is documented and illustrated by an example.


Keywords: SQP, sequential quadratic programming, nonlinear programming, non-monotone line search, numerical algorithm, distributed computing, Fortran code

## 1 Introduction

We consider the general optimization problem to minimize an objective function $f$ under nonlinear equality and inequality constraints,

$$
x \in \mathbb{R}^{n}: \begin{array}{ll} 
& \min f(x) \\
g_{j}(x)=0, & j=1, \ldots, m_{e}  \tag{1}\\
g_{j}(x) \geq 0, & j=m_{e}+1, \ldots, m \\
x_{l} \leq x \leq x_{u}
\end{array}
$$

where $x$ is an $n$-dimensional parameter vector. It is assumed that all problem functions $f(x)$ and $g_{j}(x), j=1, \ldots, m$, are continuously differentiable on the whole $\mathbb{R}^{n}$.

Sequential quadratic programming (SQP) is the standard general purpose method to solve smooth nonlinear optimization problems, at least under the following assumptions:

- The problem is not too large.
- Functions and gradients can be evaluated with sufficiently high precision.
- The problem is smooth and well-scaled.

The original code NLPQL of Schittkowski [49] is a Fortran implementation of a sequential quadratic programming (SQP) algorithm. The numerical algorithm is based on extensive comparative numerical tests, see Schittkowski [42, 46, 44], Schittkowski et al. [58], Hock and Schittkowski [26], and on further theoretical investigations published in $[43,45,47,48]$. The algorithm is extended to solve also nonlinear least squares problems efficiently, see [51] or [53], and to handle problems with very many constraints, see [56]. To conduct the numerical tests, a random test problem generator is developed for a major comparative study, see [42]. Two collections with together 306 test problems are published in Hock and Schittkowski [26] and in Schittkowski [50]. Fortran source codes and a test frame can be downloaded from the home page of the author,

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Many of them became part of the Cute test problem collection of Bongartz et al. [8]. About 80 test problems based on a Finite Element formulation are collected for a comparative evaluation in Schittkowski et al. [58]. A set of 1,300 least squares test problems solved by an extension of the code NLPQL to retain typical features of a Gauss-Newton algorithm, is described in [53]. Also these problems can be downloaded from the home page of the author together with an interactive software system called EASY-FIT, see [54].

Moreover, there exist hundreds of commercial and academic applications of NLPQL, for example

1. mechanical structural optimization, see Schittkowski, Zillober, Zotemantel [58] and Kneppe, Krammer, Winkler [29],
2. data fitting and optimal control of transdermal pharmaceutical systems, see Boderke, Schittkowski, Wolf [3] or Blatt, Schittkowski [7],
3. computation of optimal feed rates for tubular reactors, see Birk, Liepelt, Schittkowski, and Vogel [6],
4. food drying in a convection oven, see Frias, Oliveira, and Schittkowski [16],
5. optimal design of horn radiators for satellite communication, see Hartwanger, Schittkowski, and Wolf [24],
6. receptor-ligand binding studies, see Schittkowski [52],
7. optimal design of surface acoustic wave filters for signal processing, see Bünner, Schittkowski, and van de Braak [9].

Previous and present versions of NLPQLP are part of commercial libraries, modeling systems, or optimization systems like

- IMSL Library (Visual Numerics Inc., Houston) for general nonlinear programming (Version 1.0, 1981),
- ANSYS/POPT (CAD-FEM, Grafing) for structural optimization,
- DesignXplorer (ANSYS, Canonsburg) for structural design optimization,
- STRUREL (RCP, Munich) for reliability analysis,
- TEMPO (OECD Reactor Project, Halden) for control of power plants,
- Microwave Office Suit (Applied Wave Research, El Segundo) for electronic design,
- MOOROPT (Marintek, Trondheim) for the design of mooring systems,
- iSIGHT (Enginious Software/Dassault) for multi-disciplinary CAE,
- POINTER (Synaps, Atlanta) for design automation,
- EXCITE (AVL, Graz) for non-linear dynamics of power units,
- ModeFRONTIER (ESTECO, Trieste) for integrated multi-objective and multidisciplinary design optimization,
- TOMLAB/MathLab (Tomlab Optimization, Västeras, Sweden) for general nonlinear programming, least squares optimization, data fitting in dynamical systems,
- EASY-FIT (Schittkowski, Bayreuth) for data fitting in dynamical systems,
- OptiSLang (DYNARDO, Weimar), for structural design optimization,
- AMESim (IMAGINE, Roanne), for multidisciplinary system design,
- LMS OPTIMUS (NOESIS, Leuven, Belgium) for multi-disciplinary CAE,
- RADIOSS/M-OPT (MECALOG/Altair, Antony, France) for multi-disciplinary CAE,
- CHEMASIM (BASF, Ludwigshafen) for the design of chemical reactors.

Customers include, among many others, Airbus, AMD, Astrium, BASF, Bayer, Bell Labs, BMW, Chevron Research, DLR, Dow Chemical, DuPont, EADS, EMCOSS, ENSIGC, EPCOS, ESA-ESOC, Eurocopter, Fantoft Prosess, General Electric, Hoechst, Hidroelectrica Espanola, IABG, IBM, Institute for Energy Technology Halden, KFZ Karlsruhe, Kongsberg Maritime, Lockheed Martin, Loral Space Systems, MAN, Markov Processes, Marintek, MTU, NASA Langley, NASA Ames, Nevesbu, National Airspace Laboratory, Norsk Hydro Research, Norwegian Computing Center, Norwegian Defense Agency, OECD Halden, Philips, Polysar, ProSim, Rolls-Royce, Shell, Siemens, Sintef, Solar Turbines, Statoil, TNO, Transpower, USAF Research Lab, Wright R \& D Center and in addition dozens of academic research institutions all over the world.

The general availability of parallel computers and in particular of distributed computing in networks motivates a careful redesign of the original implementation NLPQL to allow simultaneous function evaluations. The resulting extensions are implemented and the code is called NLPQLP. An additional input parameter $l$ is introduced for the number of parallel machines, that is the number of function calls to be executed simultaneously. In case of $l=1$, NLPQLP is more or less identical to NLPQL besides of additional changes of the code. Otherwise, the line search procedure is modified to allow parallel function calls, which can also be applied for approximating gradients by difference formulae. The mathematical background is outlined, in particular the modification of the line search algorithm to retain convergence under parallel systems. It must be emphasized that distributed computation of function values is only simulated throughout the paper. It is up to the user to adopt the code to a particular parallel environment.

However, SQP methods are quite sensitive subject to round-off or any other errors in function and especially gradient values. If objective or constraint functions cannot be computed within machine accuracy or if the accuracy by which gradients are approximated is above the termination tolerance, the code could break down typically with the error message $\operatorname{IFAIL}=4$. In this situation, the line search cannot be terminated within a given number of iterations and the algorithm is stopped.

All new versions since 2.0 makes use of non-monotone line search in the error situation described above. The idea is to replace the reference value of the line search termination
check, $\psi_{r_{k}}\left(x_{k}, v_{k}\right)$, by

$$
\max \left\{\psi_{r_{j}}\left(x_{j}, v_{j}\right): j=k-p, \ldots, k\right\},
$$

where $\psi_{r}(x, v)$ is a merit function and $p$ a given parameter. The general idea is not new and for example described in Dai [12], where a general convergence proof for the unconstrained case is presented. The general idea goes back to Grippo, Lampariello, and Lucidi [19], and was extended to constrained optimization and trust region methods in a series of subsequent papers, see Bonnans et al. [5], Deng et al. [14], Grippo et al. [20, 21], Ke and Han [27], Ke et al. [28], Lucidi et al. [31], Panier and Tits [35], Raydan [40], and Toint $[60,61]$. However, there is a basic difference in the methodology: Our goal is to allow monotone line searches as long as they terminate successfully, and to apply a non-monotone one only in a special error situation.

Despite of strong analytical results, SQP methods do not always terminate successfully. Besides of the difficulties leading to the usage of non-monotone line search, it might happen that the search direction as computed from a certain quadratic programming subproblem, is not a downhill direction of the merit function needed to perform a line search. Possible reasons are again severe errors in function and especially gradient evaluations, or a violated regularity condition concerning linear independency of gradients of active constraints (LICQ). In the latter case, the optimization problem is not modeled in a suitable way to solve it directly by an SQP method. Our new version performs an automated restart as soon as a corresponding error message appears. The BFGS quasi-Newton matrix is reset to a multiple of the identity matrix and the matrix update procedure starts from there.

Scaling is an extremely important issue and an efficient procedure is difficult to derive in the general case without knowing too much about the numerical structure of the optimization problem. If requested by the user, the first BFGS update is started from a multiple of the identity matrix, which takes into account information from the solution of the initial quadratic programming subproblem. This restart can be repeated periodically with successively adapted scaling parameters.

In Section 2 we outline the general mathematical structure of an SQP algorithm, the non-monotone line search, and the modifications to run the code under distributed systems. Section 3 contains some numerical results obtained for a set of 306 standard test problems of the collections published in Hock and Schittkowski [26] and in Schittkowski [50]. They show the sensitivity of the new version with respect to the number of parallel machines and the influence of gradient approximations under uncertainty. Moreover, we test the non-monotone line search versus the monotone one, and generate noisy test problems by adding random errors to function values and by inaccurate gradient approximations. This situation appears frequently in practical environments, where complex simulation codes prevent accurate responses and where gradients can only be computed by a difference formula. The usage of the Fortran subroutine is documented in Section 4 and Section 5 contains an illustrative example.

## 2 Sequential Quadratic Programming Methods

Sequential quadratic programming or SQP methods belong to the most powerful nonlinear programming algorithms we know today for solving differentiable nonlinear programming problems of the form (1). The theoretical background is described e.g. in Stoer [59] and an excellent review is given by Boggs and Tolle [4]. From the more practical point of view, SQP methods are also introduced in the books of Papalambros, Wilde [36] and Edgar, Himmelblau [15]. Their excellent numerical performance is tested and compared with other methods in Schittkowski [42], and since many years they belong to the most frequently used algorithms to solve practical optimization problems.

To facilitate the notation of this section, we assume that upper and lower bounds $x_{u}$ and $x_{l}$ are not handled separately, i.e., we consider the somewhat simpler formulation

$$
\begin{align*}
& \min f(x) \\
x \in \mathbb{R}^{n}: & g_{j}(x)=0, \quad j=1, \ldots, m_{e}  \tag{2}\\
& g_{j}(x) \geq 0, \quad j=m_{e}+1, \ldots, m
\end{align*}
$$

It is assumed that all problem functions $f(x)$ and $g_{j}(x), j=1, \ldots, m$, are continuously differentiable on $\mathbb{R}^{n}$.

The basic idea is to formulate and solve a quadratic programming subproblem in each iteration which is obtained by linearizing the constraints and approximating the Lagrangian function

$$
\begin{equation*}
L(x, u):=f(x)-\sum_{j=1}^{m} u_{j} g_{j}(x) \tag{3}
\end{equation*}
$$

quadratically, where $x \in \mathbb{R}^{n}$ is the primal variable and $u=\left(u_{1}, \ldots, u_{m}\right)^{T} \in \mathbb{R}^{m}$ the multiplier vector.

To formulate the quadratic programming subproblem, we proceed from given iterates $x_{k} \in \mathbb{R}^{n}$, an approximation of the solution, $v_{k} \in \mathbb{R}^{m}$, an approximation of the multipliers, and $C_{k} \in \mathbb{R}^{n \times n}$, an approximation of the Hessian of the Lagrangian function. Then one has to solve the quadratic programming problem

$$
\begin{align*}
& \min \frac{1}{2} d^{T} C_{k} d+\nabla f\left(x_{k}\right)^{T} d \\
d \in \mathbb{R}^{n}: & \nabla g_{j}\left(x_{k}\right)^{T} d+g_{j}\left(x_{k}\right)=0, \quad j=1, \ldots, m_{e}  \tag{4}\\
& \nabla g_{j}\left(x_{k}\right)^{T} d+g_{j}\left(x_{k}\right) \geq 0, \quad j=m_{e}+1, \ldots, m
\end{align*}
$$

Let $d_{k}$ be the optimal solution and $u_{k}$ the corresponding multiplier of this subproblem. A new iterate is obtained by

$$
\begin{equation*}
\binom{x_{k+1}}{v_{k+1}}:=\binom{x_{k}}{v_{k}}+\alpha_{k}\binom{d_{k}}{u_{k}-v_{k}} \tag{5}
\end{equation*}
$$

where $\alpha_{k} \in(0,1]$ is a suitable steplength parameter.

Although we are able to guarantee that the matrix $C_{k}$ is positive definite, it is possible that (4) is not solvable due to inconsistent constraints. One possible remedy is to introduce an additional variable $\delta \in \mathbb{R}$, leading to a modified quadratic programming problem, see Schittkowski [49] for details.

The steplength parameter $\alpha_{k}$ is required in (5) to enforce global convergence of the SQP method, i.e., the approximation of a point satisfying the necessary Karush-Kuhn-Tucker optimality conditions when starting from arbitrary initial values, typically a user-provided $x_{0} \in \mathbb{R}^{n}$ and $v_{0}=0, C_{0}=I . \alpha_{k}$ should satisfy at least a sufficient decrease condition of a merit function $\phi_{r}(\alpha)$ given by

$$
\begin{equation*}
\phi_{r}(\alpha):=\psi_{r}\left(\binom{x}{v}+\alpha\binom{d}{u-v}\right) \tag{6}
\end{equation*}
$$

with a suitable penalty function $\psi_{r}(x, v)$. Implemented is the augmented Lagrangian function

$$
\begin{equation*}
\psi_{r}(x, v):=f(x)-\sum_{j \in J}\left(v_{j} g_{j}(x)-\frac{1}{2} r_{j} g_{j}(x)^{2}\right)-\frac{1}{2} \sum_{j \in K} v_{j}^{2} / r_{j} \tag{7}
\end{equation*}
$$

with $J:=\left\{1, \ldots, m_{e}\right\} \cup\left\{j: m_{e}<j \leq m, g_{j}(x) \leq v_{j} / r_{j}\right\}$ and $K:=\{1, \ldots, m\} \backslash J$, cf. Schittkowski [47]. The objective function is penalized as soon as an iterate leaves the feasible domain. The corresponding penalty parameters $r_{j}, j=1, \ldots, m$ that control the degree of constraint violation, must carefully be chosen to guarantee a descent direction of the merit function, see Schittkowski [47] or Wolfe [62] in a more general setting, i.e., to get

$$
\begin{equation*}
\phi_{r_{k}}^{\prime}(0)=\nabla \psi_{r_{k}}\left(x_{k}, v_{k}\right)^{T}\binom{d_{k}}{u_{k}-v_{k}}<0 \tag{8}
\end{equation*}
$$

Finally one has to approximate the Hessian matrix of the Lagrangian function in a suitable way. To avoid calculation of second derivatives and to obtain a final superlinear convergence rate, the standard approach is to update $C_{k}$ by the BFGS quasi-Newton formula, cf. Powell [38] or Stoer [59].

The implementation of a line search algorithm is a critical issue when implementing a nonlinear programming algorithm, and has significant effect on the overall efficiency of the resulting code. On the one hand we need a line search to stabilize the algorithm, on the other hand it is not desirable to waste too many function calls. Moreover, the behavior of the merit function becomes irregular in case of constrained optimization because of very steep slopes at the border caused by large penalty terms. Even the implementation is more complex than shown above, if linear constraints and bounds of the variables are to be satisfied during the line search.

Usually, the steplength parameter $\alpha_{k}$ is chosen to satisfy the Armijo [1] condition

$$
\begin{equation*}
\phi_{r}\left(\sigma \beta^{i}\right) \leq \phi_{r}(0)+\sigma \beta^{i} \mu \phi_{r}^{\prime}(0) \tag{9}
\end{equation*}
$$

see for example Ortega and Rheinboldt [34]. The constants are from the ranges $0<\mu<$ $0.5,0<\beta<1$, and $0<\sigma \leq 1$. We start with $i=0$ and increase $i$ until (9) is satisfied for the first time, say at $i_{k}$. Then the desired steplength is $\alpha_{k}=\sigma \beta^{i_{k}}$.

Fortunately, SQP methods are quite robust and accept the steplength one in the neighborhood of a solution. Typically the test parameter $\mu$ for the Armijo-type sufficient descent property (9) is very small. Nevertheless the choice of the reduction parameter $\beta$ must be adopted to the actual slope of the merit function. If $\beta$ is too small, the line search terminates very fast, but on the other hand the resulting stepsizes are usually too small leading to a higher number of outer iterations. On the other hand, a larger value close to one requires too many function calls during the line search. Thus, we need some kind of compromise, which is obtained by first applying a polynomial interpolation, typically a quadratic one, and use (9) only as a stopping criterion. Since $\phi_{r}(0), \phi_{r}^{\prime}(0)$, and $\phi_{r}\left(\alpha_{i}\right)$ are given, $\alpha_{i}$ the actual iterate of the line search procedure, we easily get the minimizer of the quadratic interpolation. We accept then the maximum of this value and the Armijo parameter as a new iterate, as shown by the subsequent code fragment implemented in NLPQLP.

Algorithm 2.1 Let $\beta$, $\mu$ with $0<\beta<1,0<\mu<0.5$ be given.
Start: $\alpha_{0}:=1$
For $i=0,1,2, \ldots d o$ :

1) If $\phi_{r}\left(\alpha_{i}\right)<\phi_{r}(0)+\mu \alpha_{i} \phi_{r}^{\prime}(0)$, then stop.
2) Compute $\bar{\alpha}_{i}:=\frac{0.5 \alpha_{i}^{2} \phi_{r}^{\prime}(0)}{\alpha_{i} \phi_{r}^{\prime}(0)-\phi_{r}\left(\alpha_{i}\right)+\phi_{r}(0)}$.
3) Let $\alpha_{i+1}:=\max \left(\beta \alpha_{i}, \bar{\alpha}_{i}\right)$.

Corresponding convergence results are found in Schittkowski [47]. $\bar{\alpha}_{i}$ is the minimizer of the quadratic interpolation, and we use the Armijo descent property for checking termination. Step 3 is required to avoid irregular values, since the minimizer of the quadratic interpolation could be outside of the feasible domain $(0,1]$. The search algorithm is implemented in NLPQLP together with additional safeguards, for example to prevent violation of bounds. Algorithm 4.1 assumes that $\phi_{r}(1)$ is known before calling the procedure, i.e., that the corresponding function values are given. We have to stop the algorithm, if sufficient descent is not observed after a certain number of iterations, say 10. If the tested stepsize falls below machine precision or the accuracy by which model function values are computed, the merit function cannot decrease further.

To outline the new approach, let us assume that functions can be computed simultaneously on $l$ different machines. Then $l$ test values $\alpha_{i}=\beta^{i-1}$ with $\beta=\epsilon^{1 /(l-1)}$ are selected, $i=1, \ldots, l$, where $\epsilon$ is a guess for the machine precision. Next we require $l$ parallel function calls to get the corresponding model function values. The first $\alpha_{i}$ satisfying a sufficient descent property (9), say for $i=i_{k}$, is accepted as the new steplength to set the
subsequent iterate by $\alpha_{k}:=\alpha_{i_{k}}$. One has to be sure that existing convergence results of the SQP algorithm are not violated.

The proposed parallel line search will work efficiently, if the number of parallel machines $l$ is sufficiently large, and works as follows, where we omit the iteration index $k$.

Algorithm 2.2 Let $\beta$, $\mu$ with $0<\beta<1,0<\mu<0.5$ be given.
Start: For $\alpha_{i}=\beta^{i}$ compute $\phi_{r}\left(\alpha_{i}\right)$ for $i=0, \ldots, l-1$.
For $i=0,1,2, \ldots$ do:
If $\phi_{r}\left(\alpha_{i}\right)<\phi_{r}(0)+\mu \alpha_{i} \phi_{r}^{\prime}(0)$, then stop.

To precalculate $l$ candidates in parallel at log-distributed points between a small tolerance $\alpha=\tau$ and $\alpha=1,0<\tau \ll 1$, we propose $\beta=\tau^{1 /(l-1)}$.

The paradigm of parallelism is SPMD, i.e., Single Program Multiple Data. In a typical situation we suppose that there is a complex application code providing simulation data, for example by an expensive Finite Element calculation in mechanical structural optimization. It is supposed that various instances of the simulation code providing function values, are executable on a series of different machines, so-called slaves, controlled by a master program that executes NLPQLP. By a message passing system, for example PVM, see Geist et al. [17], only very few data need to be transferred from the master to the slaves. Typically only a set of design parameters of length $n$ must to be passed. On return, the master accepts new model responses for objective function and constraints, at most $m+1$ double precision numbers. All massive numerical calculations and model data, for example the stiffness matrix of a Finite Element model in a mechanical engineering application, remain on the slave processors of the distributed system.

In both situations, i.e., the serial or parallel version, it is still possible that Algorithm 2.1 or Algorithm 2.2 breaks down because to too many iterations. In this case, we proceed from a descent direction of the merit function, but $\phi_{r}^{\prime}(0)$ is extremely small. To avoid interruption of the whole iteration process, the idea is to repeat the line search with another stopping criterion. Instead of testing (9), we accept a stepsize $\alpha_{k}$ as soon as the inequality

$$
\begin{equation*}
\phi_{r_{k}}\left(\alpha_{k}\right) \leq \max _{k-p(k)<=j<=k} \phi_{r_{j}}(0)+\alpha_{k} \mu \phi_{r_{k}}^{\prime}(0) \tag{10}
\end{equation*}
$$

is satisfied, where $p(k)$ is a predetermined parameter with $p(k)=\min \{k, p\}, p$ a given tolerance. Thus, we allow an increase of the reference value $\phi_{r_{j_{k}}}(0)$ in a certain error situation, i.e., an increase of the merit function value. To implement the non-monotone line search, we need a queue consisting of merit function values at previous iterates. In case of $k=0$, the reference value is adapted by a factor greater than 1 , i.e., $\phi_{r_{j_{k}}}(0)$ is replace by $t \phi_{r_{j_{k}}}(0), t>1$. The basic idea to store reference function values and to replace the sufficient descent property by a sufficient 'ascent' property in max-form, is for example
described in Dai [12], where a general convergence proof for the unconstrained case is presented. The general idea goes back to Grippo, Lampariello, and Lucidi [19], and was extended to constrained optimization and trust region methods in a series of subsequent papers, see Bonnans et al. [5], Deng et al. [14], Grippo et al. [20, 21], Ke and Han [27], Ke et al. [28], Lucidi et al. [31], Panier and Tits [35], Raydan [40], and Toint [60, 61]. However, there is a difference in the methodology: Our goal is to allow monotone line searches as long as they terminate successfully, and to apply a non-monotone one only in an error situation.

The final step of an SQP method consists of updating the quasi-Newton matrix $C_{k}$, e.g., by the BFGS formula

$$
\begin{equation*}
C_{k+1}:=C_{k}+\frac{q_{k} q_{k}^{T}}{p_{k}^{T} q_{k}}-\frac{C_{k} p_{k} p_{k}^{T} C_{k}}{p_{k}^{T} C_{k} p_{k}} \tag{11}
\end{equation*}
$$

where $q_{k}:=\nabla_{x} L\left(x_{k+1}, u_{k}\right)-\nabla_{x} L\left(x_{k}, u_{k}\right)$ and $p_{k}:=x_{k+1}-x_{k}$. Special safeguards guarantee that $p_{k}^{T} q_{k}>0$ and that thus all matrices $C_{k}$ remain positive definite provided that $C_{0}$ is positive definite. A possible scaling factor and restart procedure is to replace an actual $C_{k}$ by $\gamma_{k} I$ before performing the update (11), where $\gamma_{k}=\frac{p_{k}^{T} q_{k}}{p_{k}^{T} p_{k}}$ and where $I$ denotes the identity matrix, see for example Liu and Nocedal [30]. Scaled restarts are recommended, if, e.g., the convergence turns out to become extremely slow.

## 3 Performance Evaluation

### 3.1 The Test Environment

Our numerical tests use the 306 academic and real-life test problems published in Hock and Schittkowski [26] and in Schittkowski [50]. Part of them are also available in the Cute library, see Bongartz et. al [8], and their usage is described in Schittkowski [57].

Since analytical derivatives are not available for all problems, we approximate them numerically. The test examples are provided with exact solutions, either known from analytical precalculations by hand or from the best numerical data found so far.

First we need a criterion to decide whether the result of a test run is considered as a successful return or not. Let $\epsilon>0$ be a tolerance for defining the relative accuracy, $x_{k}$ the final iterate of a test run, and $x^{\star}$ the supposed exact solution known from the test problem collection. Then we call the output a successful return, if the relative error in the objective function is less than $\epsilon$ and if the maximum constraint violation is less than $\epsilon^{2}$, i.e., if

$$
f\left(x_{k}\right)-f\left(x^{\star}\right)<\epsilon\left|f\left(x^{\star}\right)\right|, \text { if } f\left(x^{\star}\right) \neq 0
$$

or

$$
f\left(x_{k}\right)<\epsilon, \text { if } f\left(x^{\star}\right)=0
$$

and

$$
r\left(x_{k}\right)=\left\|g\left(x_{k}\right)^{-}\right\|_{\infty}<\epsilon^{2}
$$

where $\|\ldots\|_{\infty}$ denotes the maximum norm and $g_{j}\left(x_{k}\right)^{-}=\min \left(0, g_{j}\left(x_{k}\right)\right), j>m_{e}$, and $g_{j}\left(x_{k}\right)^{-}=g_{j}\left(x_{k}\right)$ otherwise.

We take into account that a code returns a solution with a better function value than the known one, subject to the error tolerance of the allowed constraint violation. However, there is still the possibility that an algorithm terminates at a local solution different from the known one. Thus, we call a test run a successful one, if in addition to the above decision the internal termination conditions are satisfied subject to a reasonably small tolerance ( $I F A I L=0$ ), and if

$$
f\left(x_{k}\right)-f\left(x^{\star}\right) \geq \epsilon\left|f\left(x^{\star}\right)\right|, \text { if } f\left(x^{\star}\right) \neq 0
$$

or

$$
f\left(x_{k}\right) \geq \epsilon, \text { if } f\left(x^{\star}\right)=0
$$

and

$$
r\left(x_{k}\right)<\epsilon^{2} .
$$

For our numerical tests, we use $\epsilon=0.01$ to determine a successful return, i.e., we require a final accuracy of one per cent. Note that in all cases, NLPQLP is called with a termination tolerance of $10^{-7}$.

If gradients are not available in analytical form, they must be approximated in a suitable way. The three most popular difference formulae are the following ones:

1. Forward differences:

$$
\begin{equation*}
\frac{\partial}{\partial x_{i}} f(x) \approx \frac{1}{\eta_{i}}\left(f\left(x+\eta_{i} e_{i}\right)-f(x)\right) \tag{12}
\end{equation*}
$$

2. Two-sided differences:

$$
\begin{equation*}
\frac{\partial}{\partial x_{i}} f(x) \approx \frac{1}{2 \eta_{i}}\left(f\left(x+\eta_{i} e_{i}\right)-f\left(x-\eta_{i} e_{i}\right)\right. \tag{13}
\end{equation*}
$$

3. Fourth-order formula:

$$
\begin{equation*}
\frac{\partial}{\partial x_{i}} f(x) \approx \frac{1}{4!\eta_{i}}\left(2 f\left(x-2 \eta_{i} e_{i}\right)-16 f\left(x-\eta_{i} e_{i}\right)+16 f\left(x+\eta_{i} e_{i}\right)-2 f\left(x+2 \eta_{i} e_{i}\right)\right) \tag{14}
\end{equation*}
$$

Here $\eta_{i}=\eta \max \left(10^{-5},\left|x_{i}\right|\right)$ and $e_{i}$ is the $i$-th unit vector, $i=1, \ldots, n$. The tolerance $\eta$ depends on the difference formula and is set to $\eta=\eta_{m}{ }^{1 / 2}$ for forward differences, $\eta=\eta_{m}{ }^{1 / 3}$ for two-sided differences, and $\eta=\left(\eta_{m} / 72\right)^{1 / 4}$ for fourth-order formulae. $\eta_{m}$ is a guess for
the accuracy by which function values are computed, i.e., either machine accuracy in case of analytical formulae or an estimate of the noise level in function computations. In a similar way, derivatives of constraints are computed.

The Fortran implementation of the SQP method introduced in the previous section, is called NLPQLP. The code represents the most recent version of NLPQL which is frequently used in academic and commercial institutions. NLPQLP is prepared to run also under distributed systems, but behaves in exactly the same way as the serial version, if the number of simulated processors is set to one. Functions and gradients must be provided by reverse communication and the quadratic programming subproblems are solved by the primal-dual method of Goldfarb and Idnani [18] based on numerically stable orthogonal decompositions. NLPQLP is executed with termination accuracy $A C C=10^{-7}$ as mentioned already above, and a maximum number of iterations $M A X I T=500$.

In the subsequent tables, we use the notation

$$
\begin{aligned}
& n_{\text {succ }}-\text { number of successful test runs (according to above definition) } \\
& n_{\text {func }}-\text { average number of function evaluations } \\
& n_{\text {grad }}-\text { average number of gradient evaluations or iterations, respectively } \\
& f(x) \text { - final objective function value } \\
& r(x) \text { - final constraint violation } \\
& i_{\text {fail }} \text { - failure code }
\end{aligned}
$$

To get $n_{\text {func }}$ or $n_{\text {grad }}$, we count each evaluation of a whole set of function or gradient values, respectively, for a given iterate $x_{k}$, also in the case of several simulated processors, $l>1$. However, additional function evaluations needed for gradient approximations, are not counted for $n_{\text {func }}$. Their average number is $n_{\text {func }}$ for forward differences, $2 \times n_{\text {func }}$ for two-sided differences, and $4 \times n_{\text {func }}$ for fourth-order formulae. One gradient computation corresponds to one iteration of the SQP method.

The Fortran codes are compiled by the Intel Visual Fortran Compiler, Version 11.0, 64 bit, under Windows 7 and $\operatorname{Intel}(\mathrm{R})$ Core(TM) i7 CPU 860, 2.8 GHz , with 8 GB RAM.

### 3.2 Testing Distributed Function Calls

First we investigate the question, how parallel line searches influence the overall performance. Table 1 shows the number of successful test runs and the average number of iterations or gradient evaluations, $n_{i t}$, for an increasing number of simulated parallel calls of model functions denoted by $l$. The forward difference formula (12) is used for gradient approximations and non-monotone line search is applied with a queue size of $p=30$. Calculation time is about one second for solving all 306 test problems without random perturbations.
$l=1$ corresponds to the sequential case, when Algorithm 2.1 is applied to the line search consisting of a quadratic interpolation combined with an Armijo-type bisection strategy and a non-monotone stopping criterion.

| $l$ | $n_{\text {succ }}$ | $n_{\text {grad }}$ | $l$ | $n_{\text {succ }}$ | $n_{\text {grad }}$ |
| ---: | ---: | ---: | ---: | ---: | ---: |
| 1 | 306 | 23 | 8 | 299 | 39 |
| 3 | 224 | 177 | 9 | 301 | 32 |
| 4 | 242 | 170 | 10 | 302 | 31 |
| 5 | 268 | 114 | 15 | 303 | 23 |
| 6 | 289 | 75 | 20 | 303 | 22 |
| 7 | 297 | 45 | 50 | 303 | 22 |

Table 1: Performance Results for Parallel Line Search
In all other cases, $l>1$ simultaneous function evaluations are made according to Algorithm 2.2. To get a reliable and robust line search, one should use at least seven parallel processors. No significant improvements are observed, if we evaluate more than ten functions in parallel.

The most promising possibility to exploit a parallel system architecture occurs, when gradients cannot be calculated analytically, but have to be approximated numerically, for example by forward differences, two-sided differences, or even higher order methods. Then we need at least $n$ additional function calls, where $n$ is the number of optimization variables, or a suitable multiple of $n$.

### 3.3 Testing NLPQLP under Random Noise

For our numerical tests, we use the forward difference formula (12). To test the stability of the SQP code, we add randomly generated noise to all function values. Non-monotone line search is applied with a queue length of $p=40$ in error situations, and the line search calculation by Algorithm 2.1 is used. The BFGS quasi-Newton updates are restarted with $\rho I$ if a descent direction cannot be computed, with $\rho=10^{4}$.

To compare the different stabilization approaches, we apply three different scenarios how to handle error situations, which would otherwise lead to early termination,

- monotone line search, no restarts $(\rho=0, p=0)$,
- non-monotone line search, no restarts $(\rho=0, p=40)$,
- non-monotone line search and restarts $\left(\rho=10^{4}, p=40\right)$.

The corresponding results shown in Table 2, are evaluated for increasing random perturbations $\left(\epsilon_{\text {err }}\right)$. More precisely, if $\nu$ denotes a uniformly distributed random number between 0 and 1 , we replace $f\left(x_{k}\right)$ by $f\left(x_{k}\right)\left(1+\epsilon_{e r r}(2 \nu-1)\right)$ at each iterate $x_{k}$. In the same way, restriction functions are perturbed. The tolerance for approximating gradients, $\eta_{m}$, is set to the machine accuracy in case of $\epsilon_{e r r}=0$, and to the random noise level otherwise.

The numerical results are surprising and depend heavily on the new non-monotone line search strategy and the additional stabilization procedures. We are able to solve about $90 \%$ of the test examples in case of extremely noisy function values with at most

|  | $\rho=0, p=0$ |  |  |  | $\rho=0, p=40$ |  |  | $\rho=10^{4}, p=40$ |  |  |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | :---: |
| $\epsilon_{\text {err }}$ | $n_{\text {succ }}$ | $n_{\text {func }}$ | $n_{\text {grad }}$ | $n_{\text {succ }}$ | $n_{\text {func }}$ | $n_{\text {grad }}$ | $n_{\text {succ }}$ | $n_{\text {func }}$ | $n_{\text {grad }}$ |  |
| 0 | 306 | 36 | 22 | 306 | 37 | 22 | 306 | 37 | 22 |  |
| $10^{-12}$ | 304 | 40 | 23 | 306 | 66 | 26 | 306 | 66 | 26 |  |
| $10^{-10}$ | 301 | 45 | 24 | 305 | 72 | 28 | 306 | 72 | 29 |  |
| $10^{-8}$ | 276 | 58 | 26 | 302 | 103 | 31 | 304 | 120 | 32 |  |
| $10^{-6}$ | 248 | 77 | 30 | 295 | 167 | 40 | 302 | 222 | 49 |  |
| $10^{-4}$ | 178 | 97 | 30 | 273 | 220 | 43 | 295 | 379 | 62 |  |
| $10^{-2}$ | 92 | 163 | 36 | 224 | 308 | 48 | 279 | 630 | 78 |  |

Table 2: Test Results for 306 Academic Test Problems
one correct digit in partial derivative values. However, the stabilization process is costly. The more test problems are successfully solved, the more iterations, especially function evaluations, are needed.

### 3.4 Testing NLPQLP on Problems with Slow Convergence

In some situations, the convergence of an SQP method becomes quite slow, e.g., in case of badly scaled variables or functions, inaccurate derivatives, or inaccurate solutions of the quadratic program (4). In these situations, errors in the search direction or the partial derivatives influence the update procedure (11) and the quasi-Newton matrices $C_{k}$ are getting more and more inaccurate.

Scaled restarts as described in Section 3 are recommended, if convergence turns out to become extremely slow, especially caused by inaccurate partial derivatives. To illustrate the situation, we consider a few test runs where the examples are generated by discretizing a two-dimensional elliptic partial differential equation by the five-star formula, see Maurer and Mittelmann [32, 33]. The original formulation is that of an optimal control problem, and state and control variables are both discretized. The test problems possess a different numerical structure than those used in the previous section, i.e., a large number variables and weakly nonlinear, sparse equality constraints, and are easily scalable to larger sizes. First partial derivatives are available in analytical form.

From a total set of 13 original test cases, we select five problems which could not be solved by NLPQLP as efficiently as expected with standard solution tolerances, especially if we add some noise. Depending on the grid size, in our case 20 in each direction, we get problems with $n=722$ or $n=798$ variables, respectively, and $m_{e}=361$ or $m_{e}=437$ nonlinear equality constraints. There are no nonlinear inequality constraints. Table 3 contains a summary including the best known objective function values subject to an optimal solution vector $x^{\star}$. The maximum number of iterations is limited by 500 , and all other tolerances are thee same as before.

| problem | $n$ | $m_{e}$ | $f\left(x^{\star}\right)$ |
| :---: | :---: | :---: | :--- |
| EX1 | 722 | 361 | $0.45903 \cdot 10^{-1}$ |
| EX2 | 722 | 361 | $0.40390 \cdot 10^{-1}$ |
| EX3 | 722 | 361 | 0.11009 |
| EX4 | 798 | 437 | $0.75833 \cdot 10^{-1}$ |
| EX5 | 798 | 437 | $0.51376 \cdot 10^{-1}$ |

Table 3: Elliptic Control Problems

Note that the code NLPQLP is unable to take sparsity into account. With an SQPIPM solver being able to handle sparsity, it is possible to solve the same test problems successfully with a fine grid leading to 5.000 .000 variables and 2.500 .000 constraints, see Sachsenberg [41].

Table 4 shows the number of iterations or gradient evaluations, respectively, for three sets of test runs and different noise levels. Since we have analytical derivatives, we add the perturbations to function as we did for the first set of test runs, and to all partial derivative values. For uniformly distributed random numbers $\nu$ between 0 and 1 , we add $1+\epsilon_{\text {err }}(2 \nu-1)$ to $f\left(x_{k}\right)$ and $\partial f\left(x_{k}\right) / \partial x_{i}$ for each iterate $x_{k}$ and $i=1, \ldots, n$. In the same way, restriction functions and their derivatives are perturbed. We consider three different scenarios defined by $\epsilon_{e r r}=0, \epsilon_{e r r}=10^{-6}$, and $\epsilon_{e r r}=10^{-4}$. The obtained objective function values coincide with those of Table 3 to at least seven digits with one exception. For one test run without scaling, but noisy function and gradient values, the upper bound of 500 iterations is reached. But also in this case, four digits of the optimal function value are correct. The queue length for non-monotone line search is set to 40 and the parameter for internal restarts in error cases is set to $\rho=10^{4}$.

We apply different strategies for restarting the BFGS update, i.e., $C_{k}$,

- no scaling, i.e., $C_{0}=I$,
- initial scaling by the Oren-Luenberger procedure, i.e., update started at $\gamma_{1} I$,
- $C_{k}$ is reset if $\gamma_{k} \leq \sqrt{\epsilon_{t}}$, where $\epsilon_{t}$ is the termination accuracy,
- $C_{k}$ is reset every 7 th step,
- $C_{k}$ is reset every 15 th step.

For test runs without or only initial scaling, there are only marginal difference between unperturbed and perturbed function and derivative values. On the other hand, adaptive and periodic scaling reduces the number of iterations significantly. The best results are obtained for periodic scaling after 7 iterations. However, the number of test problems is too small and their mathematical structure is too special to retrieve general conclusions.

| noise | scaling | EX1 | EX2 | EX3 | EX4 | EX5 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\epsilon_{\text {err }}=0$ | none | 64 | 109 | 88 | 113 | 212 |
|  | initial | 64 | 108 | 75 | 108 | 202 |
|  | adaptive | 14 | 13 | 14 | 16 | 26 |
|  | 7 -step | 14 | 13 | 14 | 23 | 29 |
|  | 15-step | 20 | 20 | 21 | 25 | 38 |
| $\epsilon_{\text {err }}=10^{-6}$ | none | 64 | 109 | 88 | 110 | 385 |
|  | initial | 64 | 108 | 83 | 115 | 313 |
|  | adaptive | 14 | 17 | 18 | 60 | 35 |
|  | 7 -step | 17 | 13 | 15 | 45 | 31 |
|  | 15-step | 20 | 20 | 29 | 26 | 39 |
| $\epsilon_{e r r}=10^{-4}$ | none | 74 | 111 | 104 | 178 | 500 |
|  | initial | 63 | 116 | 102 | 171 | 397 |
|  | adaptive | 30 | 81 | 43 | 106 | 42 |
|  | 7 -step | 21 | 32 | 52 | 53 | 34 |
|  | 15-step | 48 | 30 | 29 | 25 | 57 |

Table 4: Elliptic Control Problems, Number of Iterations

## 4 Program Documentation

NLPQLP is implemented in form of a Fortran subroutine. The quadratic programming problem is solved by the code QL, an implementation of the primal-dual method of Goldfarb and Idnani [18] going back to Powell [39], see also Schittkowski [55] for more details about implementation and usage. Model functions and gradients must be provided by reverse communication. The user has to evaluate function and gradient values in the same program which executes NLPQLP, according to the following rules:

1. Choose starting values for the variables to be optimized, and store them in the first column of an array called X.
2. Compute objective and all constraint function values, store them in $\mathrm{F}(1)$ and the first column of G, respectively.
3. Compute gradients of objective function and all constraints, and store them in DF and DG, respectively. The J-th row of DG contains the gradient of the J-th constraint, $\mathrm{J}=1, \ldots, \mathrm{M}$.
4. Set $\operatorname{IFAIL}=0$ and execute NLPQLP.
5. If NLPQLP returns with IFAIL=-1, compute objective and constraint function values for all variables found in the first L columns of X , store them in F (first L positions) and G (first L columns), and call NLPQLP again.
6. If NLPQLP terminates with $\operatorname{IFAIL}=-2$, compute gradient values with respect to the variables stored in the first column of X, and store them in DF and DG. Only derivatives for active constraints, $\mathrm{ACT}(\mathrm{J})=$.TRUE., need to be computed. Then call NLPQLP again.
7. If NLPQLP terminates with $\operatorname{IFAIL}=0$, the internal optimality criteria are satisfied. In case of IFAIL $>0$, an error occurred.

If analytical derivatives are not available, simultaneous function calls can be used for gradient approximations, for example by forward differences ( $2 \mathrm{~N}>\mathrm{L}$ ), two-sided differences ( $4 \mathrm{~N}>\mathrm{L} \geq 2 \mathrm{~N}$ ), or even higher order formulae ( $\mathrm{L} \geq 4 \mathrm{~N}$ ).

## Usage:

| CALL | NLPQLP | L, | M, | ME, | MMAX, |
| :--- | ---: | ---: | ---: | ---: | ---: |

## Definition of the parameters:

L: Number of parallel systems, i.e., function calls during line search at predetermined iterates.
M : Total number of constraints.
ME : Number of equality constraints.
MMAX : Row dimension of array DG containing Jacobian of constraints. MMAX must be at least one and greater or equal to M.
N: $\quad$ Number of optimization variables.
NMAX: Row dimension of C. NMAX must be at least two and greater than N .

| MNN2: | Must be equal to M+N+N+2 when calling NLPQLP. |
| :--- | :--- |
| X(NMAX,L) : | Initially, the first column of X has to contain starting values |
| for the optimal solution. On return, X is replaced by the |  |
| current iterate. In the driving program the row dimension |  |
| of X has to be equal to NMAX. X is used internally to store |  |
| L different arguments for which function values should be |  |
| computed simultaneously. |  |
| On return, F(1) contains the final objective function value. |  |
| F is used also to store L different objective function values |  |
| to be computed from L sets of arguments stored in X. |  |
| On return, the first column of G contains the constraint |  |
| function values at the final iterate X. In the driving pro- |  |
| gram, the row dimension of G has to be equal to MMAX. |  |
| G is used internally to store L different sets of constraint |  |
| function values to be computed from L sets of arguments |  |
| stored in X. |  |

D (NMAX) : The elements of the diagonal matrix of the LDL decomposition of the quasi-Newton matrix are stored in the onedimensional array D, if LQL is false.
ACC: The user has to specify the desired final accuracy (e.g. 1.0D-7). The termination accuracy should not be much smaller than the accuracy by which gradients are computed.
ACCQP : The tolerance is needed for the QP solver to perform several tests, for example whether optimality conditions are satisfied or whether a number is considered as zero or not. If $A C C Q P$ is less or equal to zero, then the machine precision is computed by NLPQLP and subsequently multiplied by $1.0 \mathrm{D}+4$.
STPMIN : Minimum steplength in case of $\mathrm{L}>1$. Recommended is any value in the order of the accuracy by which functions are computed. The value is needed to compute a steplength reduction factor by STPMIN** (1/(L-1)). STPMIN should not fall below machine accuracy.
MAXFUN : The integer variable defines an upper bound for the number of function calls during the line search (e.g. 20). MAXFUN is only needed in case of $L=1$, and must not be greater than 50.

MAXIT : Maximum number of outer iterations, where one iteration corresponds to one formulation and solution of the quadratic programming subproblem, or, alternatively, one evaluation of gradients (e.g. 100).
MAXNM: Stack size for storing merit function values at previous iterations for non-monotone line search (e.g. 10). If MAXNM $=0$, monotone line search is performed. MAXNM should not be greater than 50 .
RHO : Parameter for performing a restart in case of IFAIL=2 by setting the BFGS-update matrix to RHO*I, where I denotes the identity matrix. The number of restarts is bounded by MAXFUN. A value greater than one is recommended. (e.g. 100).

IPRINT: Specification of the desired output level.
0 - No output of the program.
1 - Only final convergence analysis.
2 - One line of intermediate results for each iteration.
3 - More detailed information for each iteration.
4 - More line search data displayed.
Note that constraint and multiplier values are not displayed for $\mathrm{N}, \mathrm{M}>1,000$
MODE : The parameter specifies the desired version of NLPQLP.
0 - Normal execution (reverse communication!).
1 - Initial guess for multipliers in U and Hessian of the Lagrangian function in C and D provided.
In case of $\mathrm{LQL}=$. TRUE., D is ignored. Otherwise, the lower part of C has to contain the lower triangular factor of an LDL decomposition and D the diagonal part.
2 - Initial scaling (Oren-Luenberger) after first step, BFGS updates started from multiple of identity matrix.
3 - Scaled restart, if scaling parameter is less than square root of ACC.
$>3$ - Initial and repeated scaling every MODE steps, reset of BFGS matrix to multiple of identity matrix.
IOUT : Integer indicating the desired output unit number, i.e., all write-statements start with 'WRITE(IOUT,... '.
IFAIL: The parameter shows the reason for terminating a solution process. Initially, IFAIL must be set to zero. On return, IFAIL could contain the following values:
-2 - Compute new gradient values.
-1 - Compute new function values.
0 - Optimality conditions satisfied.
1 - Stop after MAXIT iterations.
2 - Uphill search direction.
3 - Underflow when computing new BFGS-update matrix.

4 - Line search exceeded MAXFUN iterations.
5 - Length of a working array too short.
6 - False dimensions, $\mathrm{M}>\mathrm{MMAX}, \mathrm{N} \geq$ NMAX, or $\mathrm{MNN} 2 \neq \mathrm{M}+\mathrm{N}+\mathrm{N}+2$.
7 - Search direction close to zero at infeasible iterate.
8 - Starting point violates lower or upper bound.
9 - Wrong input parameter, e.g., MODE, IPRINT, IOUT.
10 - Inconsistency in QP, division by zero.
$>100$ - Error message of QP solver.
WA(LWA) : WA is a double precision working array of length LWA. On return, the first N positions contain the best feasible iterate obtained, WA $(\mathrm{N}+1)$ the corresponding objective function value, and the subsequent M positions the constraint values. If no intermediate feasible solution exists, $\mathrm{WA}(\mathrm{N}+1)$ contains a large value, e.g., $1.0 \mathrm{D}+72$.
LWA: Length of WA, has to be at least at least $23^{*} \mathrm{~N}+4^{*} \mathrm{M}+3^{*} \mathrm{MMAX}+150$.
NOTE: The standard QP-solver coming together with NLPQLP (QL) needs additional memory for $3^{*}$ NMAX*NMAX $/ 2+10 *$ NMAX + MMAX $+\mathrm{M}+1$ double precision numbers.
KWA(LKWA) : KWA is an integer working array of length LKWA. On return, the first 5 positions contain the following information: KWA(1) - Number of function evaluations.
KWA(2) - Number of gradient evaluations.
KWA(3) - Iteration count.
KWA(4) - Number of QP's solved.
KWA(5) - Flag for better feasible, but non-stationary iterate $(=1)$ or not $(=0)$, see below.
LKWA: Length of KWA, has to be at least 20.
NOTE: The standard QP-solver coming together with NLPQLP (QL) needs additional memory for N integer numbers.

ACT(LACT) : The logical array indicates constraints, which NLPQLP considers to be active at the last computed iterate, i.e., $G(J, 1)$ is active, if and only if $A C T(J)$ is true for $J=1, \ldots, M$.
LACT: Length of ACT, has to be at least $2^{*} \mathrm{M}+10$.
LQL: If LQL is set to true in the calling program, the quadratic programming problem is solved proceeding from a full positive definite quasi-Newton matrix. Otherwise, a Cholesky decomposition (LDL) is performed and updated internally, so that matrix C always consists of the lower triangular factor and D of the diagonal.
QPSLVE: External subroutine to solve the quadratic programming subproblem. The calling sequence is
CALL QPSLVE(M,ME,MMAX,N,NMAX,MNN,C,D,A,B, / XL,XU,X,U,EPS,MODE,IOUT,IFAIL,IPRINT, / WAR,LWAR,IWAR,LIWAR)
For more details about the choice and dimensions of arguments, see [55].

Some of the termination reasons depend on the accuracy used for approximating gradients. If we assume that all functions and gradients are computed within machine precision and that the implementation is correct, there remain only the following possibilities that could cause an error message:

1. The termination parameter ACC is too small, so that the numerical algorithm plays around with round-off errors without being able to improve the solution. Especially the Hessian approximation of the Lagrangian function becomes unstable in this case. A straightforward remedy is to restart the optimization cycle again with a larger stopping tolerance.
2. The constraints are contradicting, i.e., the set of feasible solutions is empty. There is no way to find out, whether nonlinear and nonconvex constraints are feasible or not. Thus, the nonlinear programming algorithms will proceed until running in any of the mentioned error situations. In this case, the correctness of the model must be carefully checked.
3. Constraints are feasible, but active constraints are degenerate, e.g., redundant. One should know that SQP algorithms assume the satisfaction of the so-called linear independency constraint qualification, i.e., that gradients of active constraints are linearly independent at each iterate and in a neighborhood of an optimal solution. In this situation, it is recommended to check the formulation of the model constraints.

However, some of the error situations also occur if, because of wrong or non-accurate gradients, the quadratic programming subproblem does not yield a descent direction for the underlying merit function. In this case, one should try to improve the accuracy of function evaluations, scale the model functions in a proper way, or start the algorithm from other initial values.

Since Version 2.1, NLPQLP returns the best iterate obtained. In case of successful termination ( $\mathrm{IFAIL}=0$ ), this is always the last one. But it might be possible that in an exceptional situation, an intermediate iterate is feasible with a better objective function value than that one of the final iterate, but the KKT optimality conditions are not satisfied. In this case, the better feasible solution is stored at the first $n$ positions of the double precision working array and the corresponding objective function value at position $n+1$. Moreover, positions $n+2$ to $n+1+m$ contain the constraint values. Note that feasibility is tested by sum of constrained violations tested against ACC.

On successful return with IFAIL=0, KWA(5) is set to zero. If, however, a better feasible objective function value has been found during the first five iterations, then KWA(5) is set to 1 , the BFGS-update matrix is set to $\rho I$ with $\rho>0$, where $I$ denotes the identity matrix. The corresponding formal argument of NLPQLP is called RHO. Moreover, the multiplier approximation vector U is set to 0 . Thus, an immediate restart under control of the user is possible with $\mathrm{MODE}=1$. Some information is printed on the standard IO channel in case of IPRINT $>0$. For compatibility reasons with previous versions, RHO replaces TOLNM.

The QP solver is defined in form of an external subroutine to allow a replacement in case of exploiting special sparsity patterns. A typical example is the usage of NLPQLP for solving least squares problems, where artificially introduced equality constraints lead to a Jacobian which consist partially of the identity matrix, see Schittkowski [52, 53].

The internal scaling and restart option is borrowed from limited-memory quasi-Newton methods, see for example Liu and Nocedal [30]. If requested by the user, the quasi-Newton matrix is replaced by a scalar multiple of the identity matrix just before updating. Either an initial scaling or a reset of the whole matrix and computation of a new scaling parameter is performed depending on the input parameter MODE. A scaled restart is recommended, if, e.g., the convergence turns out to become extremely slow.

## 5 Examples

To give an example how to organize the code, we consider Rosenbrock's post office problem, i.e., test problem TP37 of Hock and Schittkowski [26].

$$
\begin{align*}
& \min -x_{1} x_{2} x_{3} \\
& x_{1}+2 x_{2}+2 x_{3} \geq 0 \\
& x_{1}, x_{2} \in \mathbb{R}: 72-x_{1}-2 x_{2}-2 x_{3} \geq 0  \tag{15}\\
& 0 \leq x_{1} \leq 42 \\
& 0 \leq x_{2} \leq 42 \\
& 0 \leq x_{3} \leq 42
\end{align*}
$$

NLPQLP comes with a couple of demo programs by which the following situations are to be illustrated:

| file name | comments |
| :---: | :--- |
| nlp_demoA.for | numerical differentiation and distributed function calls |
| nlp_demoB.for | numerical differentiation |
| nlp_demoC.for | analytical derivatives |
| nlp_demoD.for | warm and cold restarts |
| nlp_demoE.for | simultaneous function and gradient evaluation |
| nlp_demoF.for | active set strategy |
| nlp_demoG.for | active set strategy |

A Fortran source code for a typical situation is listed below. Gradients are approximated by forward differences. The function block inserted in the main program can be replaced by a subroutine call. Also the gradient evaluation is easily exchanged by an analytical one or higher order derivatives.

| IMPLICIT | NONE |
| :---: | :---: |
| INTEGER | NMAX, MMAX, LMAX, MNN2X, LWA, LKWA, LACTIV |
| PARAMETER ( | NMAX $=4$, |
| / | MMAX $=2$, |
| / | LMAX $=10$, |
| / | MNN2X $=$ MMAX + NMAX + NMAX + 2, |
| / | LWA $=1.5 *$ NMAX $*$ NMAX $+33 *$ NMAX $+9 *$ MMAX +200 , |
| / | LKWA $=$ NMAX + 10, |
| / | LACTIV $=2 *$ MMAX +10 ) |
| INTEGER | KWA (LKWA), N, ME, M, L, MNN2, MAXIT, MAXFUN, |
| / | IPRINT, MAXNM, IOUT, MODE, IFAIL, I, J, K, NFUNC |
| DOUBLE PRECISION | X (NMAX, LMAX), F(LMAX), G(MMAX,LMAX), DF(NMAX), |
| / | DG(MMAX, NMAX), U(MNN2X), XL(NMAX), XU(NMAX), |
| / | C (NMAX, NMAX), D(NMAX), WA (LWA), ACC, ACCQP, |
| / | STPMIN, EPS, EPSREL, FBCK, GBCK (MMAX), XBCK, |
| / | RHO |
| LOGICAL | ACTIVE(LACTIV), LQL |
| EXTERNAL | QL |
| Set some constants and initial values |  |
| IOUT $=6$ |  |
| $\mathrm{ACC}=1.0 \mathrm{D}-8$ |  |
| ACCQP $=1.0 \mathrm{D}-12$ |  |
| STPMIN $=1.0 \mathrm{D}-10$ |  |
| EPS $=1.0 \mathrm{D}-7$ |  |
| MAXIT $=100$ |  |
| MAXFUN $=10$ |  |

```
        MAXNM = 10
        RHO = 0.ODO
        LQL = .TRUE.
        IPRINT = 2
        N = 3
        L = N
        M = 2
        ME = 0
        MNN2 = M + N + N + 2
        MODE = 0
        IFAIL = 0
        NFUNC = 0
        DO I=1,N
        DO K=1,L
                X(I,K) = 10.0DO
            ENDDO
            XL(I) = 0.0DO
            XU(I) = 42.0D0
        ENDDO
        1 CONTINUE
=================================================================
C This is the main block to compute all function values
C simultaneously, assuming that there are L nodes.
C The block is executed either for computing a steplength
C or for approximating gradients by forward differences.
C
    DO K=1,L
        F(K) = -X (1,K)*X(2,K)*X(3,K)
        G(1,K) = X(1,K) + 2.0D0*X(2,K) + 2.0D0*X(3,K)
        G(2,K) = 72.0DO - X(1,K) - 2.0DO*X(2,K) - 2.0DO*X(3,K)
        ENDDO
C
C===================================================================
    NFUNC = NFUNC + 1
    IF (IFAIL.EQ.-1) GOTO 4
    IF (NFUNC.GT.1) GOTO 3
    2 CONTINUE
    FBCK = F(1)
    DO J=1,M
        GBCK(J) = G(J,1)
    ENDDO
    XBCK = X(1,1)
    DO I=1,N
        EPSREL = EPS*DMAX1(1.0D0,DABS(X(I,1)))
        DO K=2,L
            X(I,K) = X(I,1)
            ENDDO
            X(I,I) = X(I,1) + EPSREL
        ENDDO
        GOTO 1
    3 CONTINUE
    X(1,1) = XBCK
    DO I=1,N
        EPSREL = EPS*DMAX1(1.0D0,DABS(X(I,1)))
        DF(I) = (F(I) - FBCK)/EPSREL
        DO J=1,M
            DG(J,I) = (G(J,I) - GBCK(J))/EPSREL
        ENDDO
    ENDDO
    F(1) = FBCK
    DO J=1,M
        G(J,1) = GBCK(J)
```

[^0]4 CONTINUE

|  | CALL NLPQLP ( L, | M, | ME, | MMAX | N, |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | / NMAX, | MNN2, | X, | F | G, |
|  | 1 DF, | DG, | U, | XL | XU, |
|  | 1 C, | D, | ACC, | ACCQP | STPMIN, |
|  | / MAXFUN, | MAXIT, | MAXNM, | RHO | IPRINT, |
|  | / MODE, | IOUT, | IFAIL, | WA | LWA, |
|  | 1 KWA, | LKWA, | ACTIVE, | LACTIV | LQL, |
|  | / QL) |  |  |  |  |
|  | IF (IFAIL.EQ.-1) GOTO | 1 |  |  |  |
|  | IF (IFAIL.EQ.-2) GOTO | 2 |  |  |  |
| C |  |  |  |  |  |
|  | WRITE(IOUT, 1000) NFUNC |  |  |  |  |
| 1000 | FORMAT(, *** Number | of func | tion cal | 1s: ', |  |
| C |  |  |  |  |  |
|  | STOP |  |  |  |  |
|  | END |  |  |  |  |

The following output should appear on screen:


```
        11-0.34560000D+04 0.11D-13 1 1 0.10D+01 0.00D+00 0.25D-11
    --- Final Convergence Analysis at Last Iterate ---
        Best result at iteration: ITER = 11
        Objective function value: F(X) = -0.34560000D+04
        Solution values: X =
        0.24000000D+02 0.12000000D+02 0.12000000D+02
    Distances from lower bounds: X-XL =
        0.24000000D+02 0.12000000D+02 0.12000000D+02
        Distances from upper bounds: XU-X =
        0.18000000D+02 0.30000000D+02 0.30000000D+02
        Multipliers for lower bounds: U =
        0.00000000D+00 0.00000000D+00 0.00000000D+00
    Multipliers for upper bounds: U =
        0.00000000D+00 0.00000000D+00 0.00000000D+00
    Constraint values: G(X) =
        0.72000000D+02 -0.10658141D-13
    Multipliers for constraints: U
        0.00000000D+00 0.14400000D+03
    Number of function calls: NFUNC = 11
    Number of gradient calls: NGRAD = 11
    Number of calls of QP solver: NQL = 11
*** Number of function calls: 22
```

In case of $L=1$ and analytical derivative computations, the corresponding serial implementation of the main program is as follows:

```
IMPLICIT NONE
INTEGER NMAX, MMAX, MNN2X, LWA, LKWA, LACTIV
PARAMETER ( NMAX = 4,
/ MMAX = 2,
/ MNN2X = MMAX + NMAX + NMAX + 2,
/ LWA = 1.5*NMAX*NMAX + 33*NMAX + 9*MMAX + 200,
/ LKWA = NMAX + 10,
/ LACTIV = 2*MMAX + 10)
INTEGER KWA(LKWA), N, ME, M, L, MNN2, MAXIT, MAXFUN,
/ IPRINT, MAXNM, IOUT, MODE, IFAIL, I, J, NFUNC
DOUBLE PRECISION X(NMAX),F,G(MMAX),DF(NMAX), DG(MMAX,NMAX),
/ U(MNN2X), XL(NMAX), XU(NMAX), C(NMAX,NMAX),
/ D(NMAX), WA(LWA), ACC, ACCQP, STPMIN, RHO
LOGICAL ACTIVE(LACTIV), LQL
EXTERNAL QL
C Set some constants and initial values
```

```
IOUT = 6
```

IOUT = 6
ACC = 1.0D-10
ACC = 1.0D-10
ACCQP = 1.0D-12
ACCQP = 1.0D-12
STPMIN = 0.0
STPMIN = 0.0
MAXIT = 100
MAXIT = 100
MAXFUN = 10
MAXFUN = 10
MAXNM = 0
MAXNM = 0
RHO = O.ODO
RHO = O.ODO
LQL = .TRUE.
LQL = .TRUE.
IPRINT = 2
IPRINT = 2
N=3
N=3
M = 2
M = 2
ME = 0
ME = 0
MNN2 = M + N + N + 2

```
MNN2 = M + N + N + 2
```

C
C

```
        MODE = 0
        IFAIL = 0
        L = 1
        NFUNC = 0
        DO I=1,N
            X(I) = 10.0D0
            XL(I) = 0.0DO
            XU(I) = 42.0D0
        ENDDO
    1 CONTINUE
C This block computes all function values.
C
        F}=-X(1)*X(2)*X(3
        G(1) = X(1) + 2.0DO*X(2) + 2.0DO*X(3)
        G(2) = 72.0DO - X(1) - 2.0D0*X(2) - 2.0D0*X(3)
C
C=================================================================
        NFUNC = NFUNC + 1
        IF (IFAIL.EQ.-1) GOTO 4
    2 CONTINUE
C===============================================================
C This block computes all derivative values.
C
        DF(1) = -X (2)*X(3)
        DF(2) = -X(1)*X(3)
        DF(3) = -X(1)*X(2)
        DG(1,1) = 1.0DO
        DG(1,2) = 2.0DO
        DG(1,3) = 2.0DO
        DG(2,1) = -1.0D0
        DG(2,2) = -2.0DO
        DG(2,3) = -2.0DO
C
C==============
        CALL NLPQLP ( L, M, ME, MMAX, N,
        / NMAX, MNN2, X, F, G,
        / DF, DG, U, XL, XU,
        / MAXFUN, MAXIT, MAXNM, ACCQP, STPMIN,
    / MAXFUN, MAXIT, MAXNM, RHO, IPRINT,
    / MODE, IOUT, IFAIL, WA, LWA,
    / KWA, LKWA, ACTIVE, LACTIV, LQL,
        QL)
        IF (IFAIL.EQ.-1) GOTO 1
        IF (IFAIL.EQ.-2) GOTO 2
C
        WRITE(IOUT,1000) NFUNC
    1000 FORMAT(, *** Number of function calls: ',I3)
C
        STOP
        END
```

NLPQLP displays the following output:

## START OF THE SEQUENTIAL QUADRATIC PROGRAMMING ALGORITHM

## Parameters:

$\mathrm{N}=3$

```
\begin{tabular}{lcc} 
M & \(=\) & 2 \\
ME & \(=\) & 0 \\
MODE & \(=\) & 0 \\
ACC & \(=\) & \(0.1000 \mathrm{D}-09\) \\
ACCQP & \(=\) & \(0.1000 \mathrm{D}-11\) \\
STPMIN & \(=\) & \(0.1000 \mathrm{D}-09\) \\
MAXFUN & \(=\) & 10 \\
MAXNM & \(=\) & 0 \\
MAXIT & \(=\) & 100 \\
IPRINT & \(=\) & 2
\end{tabular}
Output in the following order:
    IT - iteration number
    F - objective function value
    SCV - sum of constraint violations
    NA - number of active constraints
    I - number of line search iterations
    ALPHA - steplength parameter
    DELTA - additional variable to prevent inconsistency
    KKT - Karush-Kuhn-Tucker optimality criterion
\begin{tabular}{|c|c|c|c|c|c|c|c|}
\hline IT & F & SCV & NA & I & ALPHA & DELTA & KKT \\
\hline 1 & -0.10000000D+04 & 0.00D+00 & 2 & 0 & \(0.00 \mathrm{D}+00\) & 0.00D+00 & \(0.44 \mathrm{D}+04\) \\
\hline 2 & -0.23625000D+04 & \(0.00 \mathrm{D}+00\) & 1 & 1 & 0.10D+01 & 0.00D+00 & 0.11D+04 \\
\hline 3 & -0.32507304D+04 & 0.36D-14 & 1 & 1 & \(0.10 \mathrm{D}+01\) & 0.00D+00 & \(0.69 \mathrm{D}+03\) \\
\hline 4 & -0.33041403D+04 & \(0.00 \mathrm{D}+00\) & 1 & 1 & \(0.10 \mathrm{D}+01\) & 0.00D+00 & \(0.36 \mathrm{D}+03\) \\
\hline 5 & -0.34527380D+04 & 0.00D+00 & 1 & 1 & \(0.10 \mathrm{D}+01\) & 0.00D+00 & \(0.58 \mathrm{D}+01\) \\
\hline 6 & -0.34559629D+04 & 0.00D+00 & 1 & 1 & 0.10D+01 & 0.00D+00 & \(0.76 \mathrm{D}-01\) \\
\hline 7 & -0.34560000D+04 & 0.00D+00 & 1 & 1 & \(0.10 \mathrm{D}+01\) & 0.00D+00 & 0.25D-04 \\
\hline 8 & -0.34560000D+04 & 0.36D-14 & 1 & 1 & 0.10D+01 & \(0.00 \mathrm{D}+00\) & 0.90D-10 \\
\hline
\end{tabular}
    --- Final Convergence Analysis at Last Iterate ---
    Objective function value: F(X) = -0.34560000D+04
    Solution values: X =
        0.24000003D+02 0.11999999D+02 0.11999999D+02
    Distances from lower bounds: X-XL =
        0.24000003D+02 0.11999999D+02 0.11999999D+02
    Distances from upper bounds: XU-X =
        0.17999997D+02 0.30000001D+02 0.30000001D+02
    Multipliers for lower bounds: U =
        0.00000000D+00 0.00000000D+00 0.00000000D+00
    Multipliers for upper bounds: U =
        0.00000000D+00 0.00000000D+00 0.00000000D+00
    Constraint values: G(X) =
        0.72000000D+02 -0.35527137D-14
    Multipliers for constraints: U =
        0.00000000D+00 0.14400000D+03
    Number of function calls: NFUNC = 8
    Number of gradient calls: NGRAD = 8
    Number of calls of QP solver: NQL = 8
*** Number of function calls: 8
```


## 6 Conclusions

We present a modification of an SQP algorithm designed for execution under a parallel computing environment (SPMD) and where a non-monotone line search is applied in
error situations. Numerical results indicate stability and robustness for a set of 306 standard test problems. It is shown that not more than 7 parallel function evaluation per iterations are required for performing a sufficiently accurate line search. Significant performance improvement is achieved by the non-monotone line search especially in case of noisy function values and numerical differentiation, and by restarts in a severe error situation. With the new version of NLPQLP, we are able to solve about $90 \%$ of a standard set of 306 test examples subject to an termination accuracy $10^{-7}$ in case of extremely noisy function values with relative accuracy of $1 \%$ and numerical differentiation. In the worst case, at most one digit of a partial derivative value is correct.

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