







GALAHAD EH

USER DOCUMENTATION

GALAHAD Optimization Library version 5.3

1 SUMMARY

This package uses an **exponential-penalty function to find a (local) minimizer of a differentiable objective function** $f(\mathbf{x})$ **of** n **variables x, subject to** m **general constraints** $\mathbf{c}^{\mathsf{L}} \leq \mathbf{c}(\mathbf{x}) \leq \mathbf{c}^{\mathsf{U}}$ and simple bounds $\mathbf{x}^{\mathsf{L}} \leq \mathbf{x} \leq \mathbf{x}^{\mathsf{U}}$ on the **variables.** Here, any of the components of the vectors of bounds \mathbf{c}^{L} , \mathbf{c}^{U} , \mathbf{x}^{L} and \mathbf{x}^{U} may be infinite. The method offers the choice of direct and iterative solution of the key trust-region subproblems, and is most suitable for large problems. First derivatives are required, and if second derivatives can be calculated, they will be exploited—if the product of second derivatives with a vector may be found but not the derivatives themselves, that may also be exploited.

ATTRIBUTES — Versions: GALAHAD_EPF_single, GALAHAD_EPF_double. Uses: GALAHAD_CLOCK, GALAHAD_NLPT, GALAHAD_SYMBOLS, GALAHAD_USERDATA, GALAHAD_SPECFILE, GALAHAD_SMT, GALAHAD_BSC, GALAHAD_MOP, GALAHAD_SSLS, GALAHAD_TRU, GALAHAD_GLTR, GALAHAD_STRINGS, GALAHAD_SPACE, GALAHAD_NORMS, GALAHAD_BLAS_interface, and GALAHAD_LAPACK_interface. Date: may 2025. Origin: N. I. M. Gould, Rutherford Appleton Laboratory. Language: Fortran 95 + TR 15581 or Fortran 2003.

2 HOW TO USE THE PACKAGE

The package is available with single, double and (if available) quadruple precision reals, and either 32-bit or 64-bit integers. Access to the 32-bit integer, single precision version requires the USE statement

USE GALAHAD_EPF_single

with the obvious substitution GALAHAD_EPF_double, GALAHAD_EPF_quadruple, GALAHAD_EPF_single_64, GALAHAD_EPF_double_64 and GALAHAD_EPF_quadruple_64 for the other variants.

If it is required to use more than one of the modules at the same time, the derived types SMT_type, GALAHAD_userdata_type, EPF_time_type, EPF_control_type, EPF_inform_type, EPF_data_type and NLPT_problem_type, (Section 2.4) and the subroutines EPF_initialize, EPF_solve, EPF_terminate, (Section 2.5) and EPF_read_specfile (Section 2.9) must be renamed on one of the USE statements.

2.1 Basic terminology

The exponential penalty function is defined to be

$$\psi(\mathbf{x}, \mathbf{w}, \boldsymbol{\mu}, \mathbf{v}, \mathbf{v}) = f(x) + \sum_{i} \mu_{i}^{L} w_{i}^{L} \exp[(c_{i}^{L} - c_{i}(\mathbf{x})) / \mu_{i}^{L}] + \sum_{j} \mu_{i}^{U} w_{i}^{U} \exp[(c_{i}(\mathbf{x}) - c_{i}^{U}) / \mu_{i}^{U}]
+ \sum_{j} \mathbf{v}_{j}^{L} \mathbf{v}_{j}^{L} \exp[(x_{j}^{L} - x_{j}) / \mathbf{v}_{j}^{L}] + \sum_{j} \mathbf{v}_{j}^{U} \mathbf{v}_{j}^{U} \exp[(x_{j} - x_{j}^{U}) / \mathbf{v}_{j}^{U}],$$
(2.1)

where c_i^L , c_i^U and $c_i(\mathbf{x})$ are the *i*-th components of \mathbf{c}^L , \mathbf{c}^U and $\mathbf{c}(\mathbf{x})$, and \mathbf{c}_j^L , c_j^U and x_j are the *j*-th components of \mathbf{x}^L , \mathbf{x}^U and \mathbf{x}_j , respectively. Here the components of $\boldsymbol{\mu}^L$, $\boldsymbol{\mu}^U$, $\boldsymbol{\nu}^L$ and $\boldsymbol{\nu}^U$ are separate **penalty parameters** for each lower and upper, general and simple-bound constraint, respectively, while those of \mathbf{w}^L , \mathbf{w}^U , \mathbf{v}^L , \mathbf{v}^U are likewise separate **weights** for the same. The algorithm iterates by approximately minimizing $\boldsymbol{\psi}(\mathbf{x}, \mathbf{w}, \boldsymbol{\mu}, \mathbf{v}, \boldsymbol{v})$ for a fixed set of penalty parameters and weights, and then adjusting these parameters and weights. The adjustments are designed so the sequence of approximate minimizers of $\boldsymbol{\psi}$ converge to that of the specified constrained optimization problem.

Key constructs are the **gradient** of the objective function

$$\mathbf{g}(\mathbf{x}) \stackrel{\text{def}}{=} \nabla_{x} f(\mathbf{x}), \tag{2.2}$$

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the Jacobian of the vector of constraints,

$$\mathbf{J}(\mathbf{x}) \stackrel{\text{def}}{=} \nabla_{x} c(\mathbf{x}), \tag{2.3}$$

and the gradient and Hessian of the Lagrangian function

$$\mathbf{g}_{L}(\mathbf{x}, \mathbf{y}) \stackrel{\text{def}}{=} \mathbf{g}(\mathbf{x}) - \mathbf{J}^{T}(\mathbf{x})\mathbf{y} - \mathbf{z} \text{ and } \mathbf{H}_{L}(\mathbf{x}, \mathbf{y}) \stackrel{\text{def}}{=} \nabla_{xx} \left[f(\mathbf{x}) - \sum_{i} y_{i} \nabla^{2} c_{i}(\mathbf{x}) \right]$$
(2.4)

for given vectors y and z.

The required solution \mathbf{x} necessarily satisfies the primal optimality conditions

$$\mathbf{c}^{\mathsf{L}} \le \mathbf{c}(\mathbf{x}) \le \mathbf{c}^{\mathsf{U}} \text{ and } \mathbf{x}^{\mathsf{L}} \le \mathbf{x} \le \mathbf{x}^{\mathsf{U}},$$
 (2.5)

the dual optimality conditions

$$\mathbf{g}(\mathbf{x}) = \mathbf{J}^{T}(\mathbf{x})\mathbf{y} + \mathbf{z} \tag{2.6}$$

where

$$y = y^{L} - y^{U}, \ z = z^{L} - z^{U}, \text{ and } (y^{L}, y^{U}, z^{L}, z^{U}) \ge 0,$$
 (2.7)

and the complementary slackness conditions

$$(\mathbf{c}(\mathbf{x}) - \mathbf{c}^{\mathsf{L}})^T \mathbf{y}^{\mathsf{L}} = 0, \ (\mathbf{c}(\mathbf{x}) - \mathbf{c}^{\mathsf{U}})^T \mathbf{y}^{\mathsf{U}} = 0, \ (\mathbf{x} - \mathbf{x}^{\mathsf{L}})^T \mathbf{z}^{\mathsf{L}} = 0 \text{ and } (\mathbf{x} - \mathbf{x}^{\mathsf{U}})^T \mathbf{z}^{\mathsf{U}} = 0,$$
 (2.8)

where the vectors \mathbf{y} and \mathbf{z} are known as the Lagrange multipliers for the general linear constraints, and the dual variables for the bounds, respectively, and where the vector inequalities hold component-wise.

2.2 Matrix storage formats

The matrices J(x) and $H_L(x,y)$ (as required and when available) may be stored in a variety of input formats.

2.2.1 Dense storage format

The matrix \mathbf{J} is stored as a compact dense matrix by rows, that is, the values of the entries of each row in turn are stored in order within an appropriate real one-dimensional array. Component n*(i-1)+j of the storage array J%val will hold the value $\mathbf{J}_{i,j}$ for $i=1,\ldots,m,\ j=1,\ldots,n$. Since $\mathbf{H}_{\mathbb{L}}$ is symmetric, only the lower triangular part (that is the part $\mathbf{H}_{\mathbb{L}ij}$ for $1 \le j \le i \le n$) should be stored. In this case the lower triangle will be stored by rows, that is component i*(i-1)/2+j of the storage array H%val will hold the value $\mathbf{H}_{\mathbb{L}ij}$ (and, by symmetry, $\mathbf{H}_{\mathbb{L}ij}$) for $1 \le j \le i \le n$.

2.2.2 Sparse co-ordinate storage format

Only the nonzero entries of the matrices are stored. For the l-th entry of J, its row index i, column index j and value J_{ij} are stored in the l-th components of the integer arrays J%row, J%col and real array J%val. The order is unimportant, but the total number of entries J%ne is required. The same scheme is applicable to H_L (thus requiring integer arrays H%row, H%col, a real array H%val, and an integer value H%ne), except that only the entries in the lower triangle should be stored.

2.2.3 Sparse row-wise storage format

Again only the nonzero entries are stored, but this time they are ordered so that those in row i appear directly before those in row i+1. For the i-th row of \mathbf{J} , the i-th component of the integer array $\mathsf{J}\$ptr$ holds the position of the first entry in this row, while $\mathsf{J}\$ptr$ (m+1) holds the total number of entries plus one. The column indices j and values \mathbf{J}_{ij} of the entries in the i-th row are stored in components $l=\mathsf{J}\$ptr(i),\ldots,\mathsf{J}\ptr (i+1)-1 of the integer array $\mathsf{J}\$col$, and real array $\mathsf{J}\$val$, respectively. The same scheme is applicable to \mathbf{H}_L (thus requiring integer arrays $\mathsf{H}\$ptr$, $\mathsf{H}\$col$, and a real array $\mathsf{H}\$val$), except that only the entries in the lower triangle should be stored.

For sparse matrices, this scheme almost always requires less storage than its predecessor.

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2.2.4 Diagonal storage format

If \mathbf{H}_{L} is diagonal (i.e., $\mathbf{H}_{Lij} = 0$ for all $1 \le i \ne j \le n$) only the diagonal entries \mathbf{H}_{Lii} for $1 \le i \le n$ need be stored, and the first n components of the array H%val may be used for the purpose. There is no sensible equivalent for the non-square matrix \mathbf{J} .

2.3 Real and integer kinds

We use the terms integer and real to refer to the fortran keywords REAL(rp_) and INTEGER(ip_), where rp_ and ip_ are the relevant kind values for the real and integer types employed by the particular module in use. The former are equivalent to default REAL for the single precision versions, DOUBLE PRECISION for the double precision cases and quadruple-precision if 128-bit reals are available, and correspond to rp_ = real32, rp_ = real64 and rp_ = real128 respectively as defined by the fortran iso_fortran_env module. The latter are default (32-bit) and long (64-bit) integers, and correspond to ip_ = int32 and ip_ = int64, respectively, again from the iso_fortran_env module.

2.4 The derived data types

Seven derived data types are accessible from the package.

2.4.1 The derived data type for holding matrices

The derived data type SMT_TYPE is used to hold the Jacobian matrix J and Hessian matrix H_L if these are available. The components of SMT_TYPE used here are:

- m is a scalar component of type INTEGER (ip_), that holds the row dimension of the matrix.
- n is a scalar component of type INTEGER (ip_), that holds the column dimension of the matrix.
- ne is a scalar variable of type INTEGER (ip_), that holds the number of matrix entries.
- type is a rank-one allocatable array of type default CHARACTER, that is used to indicate the matrix storage scheme used. Its precise length and content depends on the type of matrix to be stored (see §2.4.2).
- is a rank-one allocatable array of type REAL (rp_) and dimension at least ne, that holds the values of the entries. Each pair of off-diagonal entries $h_{ij} = h_{ji}$ of the *symmetric* matrix \mathbf{H}_L is represented as a single entry (see §2.2.1–2.2.3). Any duplicated entries that appear in the sparse co-ordinate or row-wise schemes will be summed.
- row is a rank-one allocatable array of type INTEGER (ip_), and dimension at least ne, that may hold the row indices of the entries. (see §2.2.2).
- is a rank-one allocatable array of type INTEGER(ip_), and dimension at least ne, that may hold the column indices of the entries (see §2.2.2–2.2.3).
- ptr is a rank-one allocatable array of type INTEGER (ip_), and dimension at least m + 1, that may hold the pointers to the first entry in each row (see §2.2.3).

2.4.2 The derived data type for holding the problem

The derived data type NLPT_problem_type is used to hold the problem. The relevant components of NLPT_problem_type are:

n is a scalar variable of type INTEGER (ip_-), that holds the number of optimization variables, n.

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- is a scalar variable of type INTEGER (ip_), that holds the number of general constraints, m.
- H is scalar variable of type SMT_TYPE that holds the Hessian matrix of the Lagrangian, \mathbf{H}_{L} . The following components are used here:

H*type is an allocatable array of rank one and type default CHARACTER, that is used to indicate the storage scheme used. If the dense storage scheme (see Section 2.2.1) is used, the first five components of H*type must contain the string DENSE. For the sparse co-ordinate scheme (see Section 2.2.2), the first ten components of H*type must contain the string COORDINATE, for the sparse row-wise storage scheme (see Section 2.2.3), the first fourteen components of H*type must contain the string SPARSE_BY_ROWS, and for the diagonal storage scheme (see Section 2.2.4), the first eight components of H*type must contain the string DIAGONAL.

For convenience, the procedure SMT_put may be used to allocate sufficient space and insert the required keyword into H%type. For example, if nlp is of derived type EPF_problem_type and involves a Hessian we wish to store using the co-ordinate scheme, we may simply

```
CALL SMT_put( nlp%H%type, 'COORDINATE' )
```

See the documentation for the GALAHAD package SMT for further details on the use of SMT_put.

- H%ne is a scalar variable of type INTEGER(ip_), that holds the number of entries in the **lower triangular** part of **H** in the sparse co-ordinate storage scheme (see Section 2.2.2). It need not be set for any of the other three schemes
- H%val is a rank-one allocatable array of type REAL(rp_), that holds the values of the entries of the **lower triangular** part of the Hessian matrix **H** in any of the storage schemes discussed in Section 2.2.
- H%row is a rank-one allocatable array of type INTEGER (ip_), that holds the row indices of the **lower triangular** part of **H** in the sparse co-ordinate storage scheme (see Section 2.2.2). It need not be allocated for any of the other three schemes.
- H%col is a rank-one allocatable array variable of type INTEGER(ip_), that holds the column indices of the **lower triangular** part of **H** in either the sparse co-ordinate (see Section 2.2.2), or the sparse row-wise (see Section 2.2.3) storage scheme. It need not be allocated when the dense or diagonal storage schemes are used.
- H%ptr is a rank-one allocatable array of dimension n+1 and type INTEGER (ip_), that holds the starting position of each row of the **lower triangular** part of **H**, as well as the total number of entries plus one, in the sparse row-wise storage scheme (see Section 2.2.3). It need not be allocated when the other schemes are used.
- is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the gradient **g** of the objective function. The *j*-th component of G, j = 1, ..., n, contains \mathbf{g}_j .
- is a scalar variable of type REAL (rp_), that holds the value of the objective function.
- J is scalar variable of type SMT_TYPE that holds the Jacobian matrix J(x) (if it is available). The following components are used here:
 - J*type is an allocatable array of rank one and type default CHARACTER, that is used to indicate the storage scheme used. If the dense storage scheme (see Section 2.2.1) is used, the first five components of J*type must contain the string DENSE. For the sparse co-ordinate scheme (see Section 2.2.2), the first ten components of J*type must contain the string COORDINATE, and for the sparse row-wise storage scheme (see Section 2.2.3), the first fourteen components of J*type must contain the string SPARSE_BY_ROWS.

For convenience, the procedure SMT_put may be used to allocate sufficient space and insert the required keyword into J%type. For example, if nlp is of derived type EPF_problem_type and involves a Jacobian we wish to store using the co-ordinate scheme, we may simply

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```
CALL SMT_put( nlp%J%type, 'COORDINATE' )
```

See the documentation for the GALAHAD package SMT for further details on the use of SMT_put.

- Jine is a scalar variable of type INTEGER (ip_), that holds the number of entries in J in the sparse co-ordinate storage scheme (see Section 2.2.2). It need not be set for any of the other two schemes.
- J%val is a rank-one allocatable array of type REAL (rp_), that holds the values of the entries of the Jacobian matrix **J** in any of the storage schemes discussed in Section 2.2.
- J%row is a rank-one allocatable array of type INTEGER (ip_), that holds the row indices of **J** in the sparse co-ordinate storage scheme (see Section 2.2.2). It need not be allocated for any of the other two schemes.
- J%col is a rank-one allocatable array variable of type INTEGER (ip_), that holds the column indices of **J** in either the sparse co-ordinate (see Section 2.2.2), or the sparse row-wise (see Section 2.2.3) storage scheme. It need not be allocated when the dense scheme is used.
- J%ptr is a rank-one allocatable array of dimension m+1 and type INTEGER (ip_), that holds the starting position of each row of **J**, as well as the total number of entries plus one, in the sparse row-wise storage scheme (see Section 2.2.3). It need not be allocated when the other schemes are used.
- is a rank-one allocatable array of dimension m and type REAL (rp_), that holds the constraint function $\mathbf{c}(\mathbf{x})$. The *i*-th component of C, j = 1, ..., m, contains $\mathbf{c}_j(\mathbf{x})$.
- C_1 is a rank-one allocatable array of dimension m and type REAL (rp_), that holds the vector of lower bounds \mathbf{c}^L on the constraints. The *i*-th component of C_1, i = 1, ..., m, contains c_i^L . Infinite bounds are allowed by setting the corresponding components of C_1 to any value smaller than -infinity, where infinity is a component of the control array control (see Section 2.4.3).
- C_u is a rank-one allocatable array of dimension m and type REAL (rp_), that holds the vector of upper bounds \mathbf{c}^{U} on the constraints. The *i*-th component of C_u, i = 1, ..., m, contains c_i^{U} . Infinite bounds are allowed by setting the corresponding components of C_u to any value larger than that infinity, where infinity is a component of the control array control (see Section 2.4.3).
- X_{-1} is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the vector of lower bounds \mathbf{x}^L on the the variables. The j-th component of X_{-1} , j = 1, ..., n, contains x_j^L . Infinite bounds are allowed by setting the corresponding components of X_{-1} to any value smaller than -infinity, where infinity is a component of the control array control (see Section 2.4.3).
- X_u is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the vector of upper bounds \mathbf{x}^{U} on the variables. The j-th component of X_u, j = 1, ..., n, contains x_j^{U} . Infinite bounds are allowed by setting the corresponding components of X_u to any value larger than that infinity, where infinity is a component of the control array control (see Section 2.4.3).
- GL is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the gradient \mathbf{g}_L of the Lagrangian function. The j-th component of GL, $j = 1, \ldots, n$, contains \mathbf{g}_{Lj} .
- X is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the values **x** of the optimization variables. The *j*-th component of X, j = 1, ..., n, contains x_j .
- is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the values **x** of the optimization variables. The *j*-th component of X, j = 1, ..., n, contains x_j .
- Y is a rank-one allocatable array of dimension m and type REAL (rp_), that holds the values y of the Lagrange multipliers. The *i*-th component of Y, i = 1, ..., m, contains y_i .
- is a rank-one allocatable array of dimension n and type REAL (rp_), that holds the values **x** of the dual variables. The *j*-th component of \mathbb{Z} , j = 1, ..., n, contains z_j .

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pname is a scalar variable of type default CHARACTER and length 10, which contains the "name" of the problem for printing. The default "empty" string is provided.

- VNAMES is a rank-one allocatable array of dimension n and type default CHARACTER and length 10, whose *j*-th entry contains the "name" of the *j*-th variable for printing. This is only used if "debug" printing control*print_level > 4) is requested, and will be ignored if the array is not allocated.
- CNAMES is a rank-one allocatable array of dimension m and type default CHARACTER and length 10, whose *i*-th entry contains the "name" of the *i*-th constraint for printing. This is only used if "debug" printing control%print_level > 4) is requested, and will be ignored if the array is not allocated.

2.4.3 The derived data type for holding control parameters

The derived data type EPF_control_type is used to hold controlling data. Default values may be obtained by calling EPF_initialize (see Section 2.5.1), while components may also be changed by calling GALAHAD_EPF_read_spec (see Section 2.9.1). The components of EPF_control_type are:

- error is a scalar variable of type INTEGER (ip_), that holds the stream number for error messages. Printing of error messages in EPF_solve and EPF_terminate is suppressed if error ≤ 0 . The default is error = 6.
- out is a scalar variable of type INTEGER (ip_), that holds the stream number for informational messages. Printing of informational messages in EPF_solve is suppressed if out < 0. The default is out = 6.
- print_level is a scalar variable of type INTEGER(ip_), that is used to control the amount of informational output which is required. No informational output will occur if print_level ≤ 0 . If print_level = 1, a single line of output will be produced for each iteration of the process. If print_level ≥ 2 , this output will be increased to provide significant detail of each iteration. The default is print_level = 0.
- start_print is a scalar variable of type INTEGER (ip_), that specifies the first iteration for which printing will occur in EPF_solve. If start_print is negative, printing will occur from the outset. The default is start_print = -1.
- stop_print is a scalar variable of type INTEGER(ip_), that specifies the last iteration for which printing will occur in EPF_solve. If stop_print is negative, printing will occur once it has been started by start_print. The default is stop_print = -1.
- print_gap is a scalar variable of type INTEGER (ip_). Once printing has been started, output will occur once every print_gap iterations. If print_gap is no larger than 1, printing will be permitted on every iteration. The default is print_gap = 1.
- max_it is a scalar variable of type INTEGER (ip_), that holds the maximum number of iterations which that be allowed in EPF_solve. The default is max_it = 100.
- max_eval is a scalar variable of type INTEGER(ip_), that holds the maximum number of function evaluations that will be allowed in EPF_solve. The default is max_eval = 10000.
- alive_unit is a scalar variable of type <code>INTEGER(ip_)</code>. If alive_unit > 0, a temporary file named alive_file (see below) will be created on stream number alive_unit on initial entry to <code>GALAHAD_EPF_solve</code>, and execution of <code>GALAHAD_EPF_solve</code> will continue so long as this file continues to exist. Thus, a user may terminate execution simply by removing the temporary file from this unit. If alive_unit ≤ 0 , no temporary file will be created, and execution cannot be terminated in this way. The default is alive_unit = 60.
- update_multipliers_itmin is a scalar variable of type INTEGER(ip_), that holds the smallest iteration number for which a Lagrange multipliers/dual variables update will be attempted. Up until this value, only penalty parameter reductions will be allowed. The default is update_multipliers_itmin = 0.

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- update_multipliers_tol is a scalar variable of type REAL(rp_), that is used to specify the minimum value the dual infeasibility is allowed to be before Lagrange multipliers/dual variables updates will be attempted. The default is update_multipliers_tol = 10¹⁹.
- infinity is a scalar variable of type REAL(rp_), that is used to specify which constraint bounds are infinite. Any bound larger than infinity in modulus will be regarded as infinite. The default is infinity = 10^{19} .
- stop_abs_p and stop_rel_p are scalar variables of type REAL(rp_), that hold the required absolute and relative accuracy for the primal infeasibility (see Section 4). The absolute value of each component of the primal infeasibility on exit is required to be smaller than the larger of stop_abs_p and stop_rel_p times a "typical value" for this component. The defaults are stop_abs_p = stop_rel_p = $u^{1/3}$, where u is EPSILON(1.0) (EPSILON(1.0D0) in GALAHAD_EPF_double).
- stop_abs_d and stop_rel_d are scalar variables of type REAL (rp_), that hold the required absolute and relative accuracy for the dual infeasibility (see Section 4). The absolute value of each component of the dual infeasibility on exit is required to be smaller than the larger of stop_abs_p and stop_rel_p times a "typical value" for this component. The defaults are stop_abs_d = stop_rel_d = $u^{1/3}$, where u is EPSILON(1.0) (EPSILON(1.0D0) in GALAHAD_EPF_double).
- stop_abs_c and stop_rel_c are scalar variables of type REAL (rp_), that hold the required absolute and relative accuracy for the violation of complementary slackness (see Section 4). The absolute value of each component of the complementary slackness on exit is required to be smaller than the larger of stop_abs_p and stop_rel_p times a "typical value" for this component. The defaults are stop_abs_c = stop_rel_c = $u^{1/3}$, where u is EPSILON(1.0) (EPSILON(1.0D0) in GALAHAD_EPF_double).
- stop_s is a scalar variable of type REAL (rp_), that is used to specify the minimum acceptable correction step s relative to the current estimate of the solution x The algorithm will be deemed to have converged if $|s_i| \le \text{stop}_s * \max(1, |x_i|)$ for all i = 1, ..., n. The default is stop_s = u, where u is EPSILON(1.0) (EPSILON(1.0D0) in GALAHAD_EPF_double).
- initial_mu is a scalar variable of type REAL (rp_), that holds the required initial value of the penalty parameter. If initial_radius ≤ 0 , the initial penalty parameter will be chosen automatically by GALAHAD_EPF_solve The default is initial_radius = 0.1.
- mu_reduce is a scalar variable of type REAL (rp_), that holds the amount by which the penalty parameter is reduced at the end of an iteration. The default is mu_reduce = 0.5.
- obj_unbounded is a scalar variable of type default REAL (rp_), that specifies smallest value of the objective function that will be tolerated before the problem is declared to be unbounded from below. The default is potential_unbounded = $-u^{-2}$, where u is EPSILON(1.0) (EPSILON(1.0D0) in GALAHAD_EPF_double).
- try_advanced_start is a scalar variable of type default REAL(rp_), that specifies the largest value of the KKT residuals before an advanced start will be attempted. The default is try_advanced_start = 10^{-2} .
- try_sqp_start is a scalar variable of type default REAL (rp_), that specifies the largest value of the KKT residuals before an advanced SQP start will be attempted. The default is try_sqp_start = 10^{-4} .
- stop_advance_start is a scalar variable of type default REAL(rp_), that specifies the smallest value of the KKT residuals that an advanced start will be attempted. The default is stop_advance_start = 10^{-8} .
- cpu_time_limit is a scalar variable of type REAL(rp_), that is used to specify the maximum permitted CPU time.

 Any negative value indicates no limit will be imposed. The default is cpu_time_limit = 1.0.
- clock_time_limit is a scalar variable of type REAL(rp_), that is used to specify the maximum permitted elapsed system clock time. Any negative value indicates no limit will be imposed. The default is clock_time_limit = 1.0.

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hessian_available is a scalar variable of type default LOGICAL, that should be set .TRUE. if the user will provide second derivatives (either by providing an appropriate evaluation routine to the solver or by reverse communication, see Section 2.7), and .FALSE. if the second derivatives are not explicitly available. The default is hessian_available = .TRUE..

- subproblem_direct is a scalar variable of type default LOGICAL, that should be set .TRUE. if a direct (factorization) method is desired when solving for the step, and .FALSE. if an iterative method suffices. The default is subproblem_direct = .FALSE..
- space_critical is a scalar variable of type default LOGICAL, that must be set .TRUE. if space is critical when allocating arrays and .FALSE. otherwise. The package may run faster if space_critical is .FALSE. but at the possible expense of a larger storage requirement. The default is space_critical = .FALSE..
- deallocate_error_fatal is a scalar variable of type default LOGICAL, that must be set .TRUE. if the user wishes to terminate execution if a deallocation fails, and .FALSE. if an attempt to continue will be made. The default is deallocate_error_fatal = .FALSE..
- alive_file is a scalar variable of type default CHARACTER and length 30, that gives the name of the temporary file whose removal from stream number alive_unit terminates execution of GALAHAD_EPF_solve. The default is alive_unit = ALIVE.d.
- prefix is a scalar variable of type default CHARACTER and length 30, that may be used to provide a user-selected character string to preface every line of printed output. Specifically, each line of output will be prefaced by the string prefix (2:LEN(TRIM(prefix))-1), thus ignoreing the first and last non-null components of the supplied string. If the user does not want to preface lines by such a string, they may use the default prefix = "".
- BSC_control is a scalar variable of type BSC_control_type whose components are used to control the formation of the Hessian matrix of the penalty function, as performed by the package GALAHAD_BSC. See the specification sheet for the package GALAHAD_BSC for details, and appropriate default values.
- SSLS_control is a scalar variable of type SSLS_control_type whose components are used to control the linear solve aspects of the calculation, as performed by the package GALAHAD_SSLS. See the specification sheet for the package GALAHAD_SSLS for details, and appropriate default values.
 - GLTR_control_type whose components are used to control the iterative trust-region step calculation (if any), performed by the package GALAHAD_GLTR. See the specification sheet for the package GALAHAD_GLTR for details, and appropriate default values (but note that value of GLTR_control%unitm may be changed by GALAHAD_EPF_solve).
- TRU_control is a scalar variable of type TRU_control_type whose components are used to control the minimization of the penalty function, performed by the package GALAHAD_TRU. See the specification sheet for the package GALAHAD_TRU for details, and appropriate default values (but note that value for TRU_control%hessian_available, will be overridden by GALAHAD_EPF_solve).

2.4.4 The derived data type for holding timing information

The derived data type <code>EPF_time_type</code> is used to hold elapsed CPU and system clock times for the various parts of the calculation. The components of <code>EPF_time_type</code> are:

total is a scalar variable of type default REAL, that gives the CPU total time spent in the package.

preprocess is a scalar variable of type REAL (rp_), that gives the CPU time spent reordering the problem to standard form prior to solution.

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- analyse is a scalar variable of type REAL(rp_), that gives the CPU time spent analysing required matrices prior to factorization.
- factorize is a scalar variable of type REAL (rp_), that gives the CPU time spent factorizing the required matrices.
- solve is a scalar variable of type REAL (rp_), that gives the CPU time spent using the factors to solve relevant linear equations.
- clock_total is a scalar variable of type default REAL, that gives the total elapsed system clock time spent in the package.
- clock_preprocess is a scalar variable of type REAL (rp_), that gives the elapsed system clock time spent reordering the problem to standard form prior to solution.
- clock_analyse is a scalar variable of type REAL(rp_), that gives the elapsed system clock time spent analysing required matrices prior to factorization.
- clock_factorize is a scalar variable of type REAL(rp_), that gives the elapsed system clock time spent factorizing the required matrices.
- clock_solve is a scalar variable of type REAL (rp_), that gives the elapsed system clock time spent using the factors to solve relevant linear equations.

2.4.5 The derived data type for holding informational parameters

The derived data type EPF_inform_type is used to hold parameters that give information about the progress and needs of the algorithm. The components of EPF_inform_type are:

- status is a scalar variable of type INTEGER (ip_), that gives the exit status of the algorithm. See Sections 2.7 and 2.8 for details.
- alloc_status is a scalar variable of type INTEGER(ip_-), that gives the status of the last attempted array allocation or deallocation. This will be 0 if status = 0.
- bad_alloc is a scalar variable of type default CHARACTER and length 80, that gives the name of the last internal array for which there were allocation or deallocation errors. This will be the null string if status = 0.
- n_free is a scalar variable of type INTEGER (ip_), that holds the number of variables that are free from their bounds.
- iter is a scalar variable of type INTEGER (ip_), that holds the number of iterations performed.
- factorization_status is a scalar variable of type INTEGER(ip_), that gives the return status from the matrix factorization.
- max_entries_factors is a scalar variable of type INTEGER(int64), that gives the maximum number of entries in any of the matrix factorizations performed during the calculation.
- factorization_max is a scalar variable of type INTEGER (ip_), that gives the largest number of factorizations required during a subproblem solution.
- factorization_integer is a scalar variable of type default INTEGER (ip_), that gives the amount of integer storage used for the matrix factorization.
- factorization_real is a scalar variable of type INTEGER(ip_), that gives the amount of real storage used for the matrix factorization.

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f_eval is a scalar variable of type INTEGER(ip_), that gives the total number of objective function evaluations performed

- g_eval is a scalar variable of type INTEGER(ip_), that gives the total number of objective function gradient evaluations performed.
- h_eval is a scalar variable of type INTEGER(ip_), that gives the total number of objective function Hessian evaluations performed.
- obj is a scalar variable of type REAL (rp_), that holds the value of the objective function at the best estimate of the solution found.
- primal_infeasibility is a scalar variable of type REAL(rp_), that holds the norm of the violation of primal optimality (see Section 2.4.4) at the best estimate of the solution found.
- dual_infeasibility is a scalar variable of type REAL(rp_), that holds the norm of the violation of dual optimality (see Section 2.4.4) at the best estimate of the solution found.
- complementary_slackness is a scalar variable of type REAL(rp_), that holds the norm of the violation of complementary slackness (see Section 2.4.4) at the best estimate of the solution found.
- mu is a scalar variable of type REAL (rp_), that holds the current value of the penalty parameter.
- factorization_average is a scalar variable of type REAL (rp_), that gives the average number of factorizations per subproblem solved.
- time is a scalar variable of type EPF_time_type whose components are used to hold elapsed elapsed CPU and system clock times for the various parts of the calculation (see Section 2.4.4).
- BSC_inform is a scalar variable of type BSC_inform_type whose components give information about the formation of the Hessian matrix of the penalty function, as performed by the package GALAHAD_BSC. See the specification sheet for the package GALAHAD_BSC for details.
- SSLS_inform is a scalar variable of type SSLS_inform_type whose components give information about the progress and needs of the linear-solve stages of the algorithm performed by the package GALAHAD_SSLS. See the specification sheet for the package GALAHAD_SSLS for details.
- GLTR_inform is a scalar variable of type GLTR_inform_type whose components give information about the progress and needs of the iterative solution stages of the algorithm performed by the package GALAHAD_GLTR. See the specification sheet for the package GALAHAD_GLTR for details.
- TRU_inform is a scalar variable of type TRU_inform_type whose components give information about the progress and needs of the algorithm used to minimize the penalty function, as performed by the package GALAHAD_TRU. See the specification sheet for the package GALAHAD_TRU for details.

2.4.6 The derived data type for holding problem data

The derived data type EPF_data_type is used to hold all the data for a particular problem, or sequences of problems with the same structure, between calls of EPF procedures. This data should be preserved, untouched (except as directed on return from GALAHAD_EPF_solve with positive values of inform%status, see Section 2.7), from the initial call to EPF_initialize to the final call to EPF_terminate.

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2.4.7 The derived data type for holding user data

The derived data type GALAHAD_userdata_type is available from the package GALAHAD_userdata to allow the user to pass data to and from user-supplied subroutines for function and derivative calculations (see Section 2.6). Components of variables of type GALAHAD_userdata_type may be allocated as necessary. The following components are available:

integer is a rank-one allocatable array of type INTEGER (ip_).

real is a rank-one allocatable array of type default REAL (rp_)

complex is a rank-one allocatable array of type default COMPLEX (double precision complex in GALAHAD_EPF_double).

character is a rank-one allocatable array of type default CHARACTER.

logical is a rank-one allocatable array of type default LOGICAL.

integer_pointer is a rank-one pointer array of type INTEGER (ip_).

real_pointer is a rank-one pointer array of type default REAL (rp_)

complex_pointer is a rank-one pointer array of type default COMPLEX (double precision complex in GALAHAD_EPF_-double).

character_pointer is a rank-one pointer array of type default CHARACTER.

logical_pointer is a rank-one pointer array of type default LOGICAL.

2.5 Argument lists and calling sequences

There are three procedures for user calls (see Section 2.9 for further features):

- 1. The subroutine EPF_initialize is used to set default values, and initialize private data, before solving one or more problems with the same sparsity and bound structure.
- 2. The subroutine EPF_solve is called to solve the problem.
- 3. The subroutine EPF_terminate is provided to allow the user to automatically deallocate array components of the private data, allocated by EPF_solve, at the end of the solution process. It is important to do this if the data object is re-used for another problem with a different structure since EPF_initialize cannot test for this situation, and any existing associated targets will subsequently become unreachable.

We use square brackets [] to indicate OPTIONAL arguments.

2.5.1 The initialization subroutine

Default values are provided as follows:

```
CALL EPF_initialize( data, control, inform )
```

data is a scalar INTENT (INOUT) argument of type EPF_data_type (see Section 2.4.6). It is used to hold data about the problem being solved.

control is a scalar INTENT (OUT) argument of type EPF_control_type (see Section 2.4.3). On exit, control contains default values for the components as described in Section 2.4.3. These values should only be changed after calling EPF_initialize.

inform is a scalar INTENT (OUT) argument of type EPF_inform_type (see Section 2.4.5). A successful call to EPF_initialize is indicated when the component status has the value 0. For other return values of status, see Section 2.8.

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2.5.2 The minimization subroutine

The minimization algorithm is called as follows:

```
CALL EPF_solve( nlp, control, inform, data, userdata[, eval_FC, eval_GJ, eval_HLPROD] )
```

nlp is a scalar INTENT (INOUT) argument of type NLPT_problem_type (see Section 2.4.2). It is used to hold data about the problem being solved. For a new problem, the user must allocate all the array components, and set values for nlp%n and the required integer components of nlp%H if second derivatives will be used. Users are free to choose whichever of the matrix formats described in Section 2.2 is appropriate for **H** for their application.

The component nlp%X must be set to an initial estimate, \mathbf{x}^0 , of the minimization variables. A good choice will increase the speed of the package, but the underlying method is designed to converge (at least to a local solution) from an arbitrary initial guess.

On exit, the component nlp%X will contain the best estimates of the minimization variables \mathbf{x} , while nlp%G will contain the best estimates of the dual variables \mathbf{z} .

```
Restrictions: nlp%n > 0 and nlp%H%type \in \{'DENSE', 'COORDINATE', 'SPARSE_BY_ROWS', 'DIAGONAL'\}.
```

- control is a scalar INTENT (IN) argument of type EPF_control_type (see Section 2.4.3). Default values may be assigned by calling EPF_initialize prior to the first call to EPF_solve. Note that value for TRU_control%hessian_available, will be overridden by GALAHAD_EPF_solve.
- inform is a scalar INTENT (INOUT) argument of type EPF_inform_type (see Section 2.4.5). On initial entry, the component status must be set to the value 1. Other entries need note be set. A successful call to EPF_solve is indicated when the component status has the value 0. For other return values of status, see Sections 2.7 and 2.8.
- data is a scalar INTENT (INOUT) argument of type EPF_data_type (see Section 2.4.6). It is used to hold data about the problem being solved. With the possible exceptions of the components eval_status and U (see Section 2.7), it must not have been altered by the user since the last call to EPF_initialize.
- userdata is a scalar INTENT(INOUT) argument of type GALAHAD_userdata_type whose components may be used to communicate user-supplied data to and from the OPTIONAL subroutines eval_FC, eval_GJ, eval_HL and eval_HLPROD (see Section 2.4.7).
- eval_FC is an OPTIONAL user-supplied subroutine whose purpose is to evaluate the value of the objective function $f(\mathbf{x})$ and constraints $\mathbf{c}(\mathbf{x})$ at a given vector \mathbf{x} . See Section 2.6.1 for details. If eval_FC is present, it must be declared EXTERNAL in the calling program. If eval_FC is absent, GALAHAD_EPF_solve will use reverse communication to obtain function values (see Section 2.7).
- eval_GJ is an OPTIONAL user-supplied subroutine whose purpose is to evaluate the value of the gradient of the objective function $\mathbf{g}(\mathbf{x})$ in (2.2) and the Jacobian of the constraints $\mathbf{J}(\mathbf{x})$ in (2.3) at a given vector \mathbf{x} . See Section 2.6.2 for details. If eval_GJ is present, it must be declared EXTERNAL in the calling program. If eval_GJ is absent, GALAHAD_EPF_solve will use reverse communication to obtain gradient values (see Section 2.7).
- eval_HL is an OPTIONAL user-supplied subroutine whose purpose is to evaluate the value of the Hessian of the Lagrangian function $\mathbf{H}_{L}(\mathbf{x},\mathbf{y})$ in (2.4) at a given vectors \mathbf{x} and \mathbf{y} . See Section 2.6.3 for details. If eval_H is present, it must be declared EXTERNAL in the calling program. If eval_H is absent, GALAHAD_EPF_solve will use reverse communication to obtain Hessian values (see Section 2.7).
- eval_HLPROD is an OPTIONAL user-supplied subroutine whose purpose is to evaluate the value of the product $\mathbf{H}_L(\mathbf{x},\mathbf{y})\mathbf{v}$ of the Hessian of the Lagrangian function with a given vector \mathbf{v} . See Section 2.6.4 for details. If eval_HLPROD is present, it must be declared EXTERNAL in the calling program. If eval_HLPROD is absent, GALAHAD_EPF_solve will use reverse communication to obtain Hessian-vector products (see Section 2.7).

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2.5.3 The termination subroutine

All previously allocated arrays are deallocated as follows:

```
CALL EPF_terminate( data, control, inform )
```

data is a scalar INTENT (INOUT) argument of type EPF_data_type exactly as for EPF_solve, which must not have been altered by the user since the last call to EPF_initialize. On exit, array components will have been deallocated.

control is a scalar INTENT (IN) argument of type EPF_control_type exactly as for EPF_solve.

inform is a scalar INTENT (OUT) argument of type EPF_inform_type exactly as for EPF_solve. Only the component status will be set on exit, and a successful call to EPF_terminate is indicated when this component status has the value 0. For other return values of status, see Section 2.8.

2.6 Function and derivative values

2.6.1 objective function and constraint values via internal evaluation

If the argument eval_FC is present when calling GALAHAD_EPF_solve, the user is expected to provide a subroutine of that name to evaluate the values of the objective function $f(\mathbf{x})$ and/or the constraints $\mathbf{c}(\mathbf{x})$.

The routine must be specified as

```
SUBROUTINE eval_FC( status, X, userdata, f, C)
```

whose arguments are as follows:

- status is a scalar INTENT (OUT) argument of type INTEGER (ip_), that should be set to 0 if the routine has been able to evaluate the objective function and constraints as required, and to a non-zero value if the evaluation has not been possible.
- X is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector x.
- userdata is a scalar INTENT (INOUT) argument of type GALAHAD_userdata_type whose components may be used to communicate user-supplied data to and from the subroutines eval_FC, eval_GJ, eval_HL and eval_HLPROD (see Section 2.4.7).
- is an OPTIONAL scalar INTENT (OUT) argument of type REAL (rp_), that should be set to the value of the objective function $f(\mathbf{x})$ evaluated at the vector \mathbf{x} input in X if f is present.
- C is an optional rank-one INTENT (OUT) argument of type REAL (rp_), whose components should be set to the values of the constraints c(x) evaluated at the vector x input in X if C is present.

2.6.2 Gradient and Jacobian values via internal evaluation

If the argument eval_GJ is present when calling GALAHAD_EPF_solve, the user is expected to provide a subroutine of that name to evaluate the value of the gradient the objective function $\nabla_x f(\mathbf{x})$. The routine must be specified as

```
SUBROUTINE eval_GJ( status, X, userdata, G, J\_val )
```

whose arguments are as follows:

status is a scalar INTENT (OUT) argument of type INTEGER (ip_), that should be set to 0 if the routine has been able to evaluate the gradient and Jacobian if required, and to a non-zero value if the evaluation has not been possible.

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- is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector x.
- userdata is a scalar INTENT (INOUT) argument of type GALAHAD_userdata_type whose components may be used to communicate user-supplied data to and from the subroutines eval_FC, eval_GJ, eval_HL and eval_HLPROD (see Section 2.4.7).
- G is a rank-one INTENT (OUT) argument of type REAL (rp_), whose components should be set to the values of the gradient of the objective function $\nabla_x f(\mathbf{x})$ evaluated at the vector \mathbf{x} input in X if G is present.
- J_val is a scalar INTENT (OUT) argument of type REAL (rp_), whose components should be set to the values of the Jacobian J(x) evaluated at the vector x input in X if J_val is present. The values should be input in the same order as that in which the array indices were given in nlp%J.

2.6.3 Hessian values via internal evaluation

If the argument eval_HL is present when calling GALAHAD_EPF_solve, the user is expected to provide a subroutine of that name to evaluate the values of the Hessian of the Lagrangian function $\mathbf{H}_{\scriptscriptstyle L}(\mathbf{x},\mathbf{y})$. The routine must be specified as

```
SUBROUTINE eval_HL( status, X, Y, userdata, H\_val )
```

whose arguments are as follows:

- status is a scalar INTENT (OUT) argument of type INTEGER (ip_), that should be set to 0 if the routine has been able to evaluate the Hessian of the Lagrangian function, and to a non-zero value if the evaluation has not been possible.
- X is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector x.
- y is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector y.
- userdata is a scalar INTENT (INOUT) argument of type GALAHAD_userdata_type whose components may be used to communicate user-supplied data to and from the subroutines eval_FC, eval_GJ, eval_HL and eval_HLPROD (see Section 2.4.7).
- H_val is a scalar INTENT (OUT) argument of type REAL (rp_), whose components should be set to the values of the Hessian of the Lagrangian function $\mathbf{H}_L(\mathbf{x},\mathbf{y})$ in (2.4) evaluated at the vectors \mathbf{x} and \mathbf{y} input in X and Y. The Hessian values should be input in the same order as that in which the array indices were given in nlp%H.

2.6.4 Hessian-vector products via internal evaluation

If the argument <code>eval_HLPROD</code> is present when calling <code>GALAHAD_EPF_solve</code>, the user is expected to provide a subroutine of that name to evaluate the sum $\mathbf{u} + \mathbf{H}_{\scriptscriptstyle L}(\mathbf{x},\mathbf{y})\mathbf{v}$ involving the product of the Hessian of the Lagrangian function $\mathbf{H}_{\scriptscriptstyle L}(\mathbf{x},\mathbf{y})$ with a given vector \mathbf{v} . The routine must be specified as

```
SUBROUTINE eval_HLPROD( status, X, Y, userdata, U, V, got_h)
```

whose arguments are as follows:

- status is a scalar INTENT (OUT) argument of type INTEGER (ip_), that should be set to 0 if the routine has been able to evaluate the sum $\mathbf{u} + \mathbf{H}_{L}(\mathbf{x}, \mathbf{y})\mathbf{v}$, and to a non-zero value if the evaluation has not been possible.
- X is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector x.
- Y is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector y.

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- userdata is a scalar INTENT (INOUT) argument of type GALAHAD_userdata_type whose components may be used to communicate user-supplied data to and from the subroutines eval_FC, eval_GJ, eval_HL and eval_HLPROD (see Section 2.4.7).
- U is a rank-one INTENT (INOUT) array argument of type REAL (rp_) whose components on input contain the vector \mathbf{u} and on output the sum $\mathbf{u} + \nabla_{xx} f(\mathbf{x}) \mathbf{v}$.
- V is a rank-one INTENT (IN) array argument of type REAL (rp_) whose components contain the vector v.
- got_h is an OPTIONAL scalar INTENT (IN) argument of type default LOGICAL. If the Hessian has already been evaluated at the current and y, got_h will be PRESENT and set .TRUE.; if this is the first time the Hessian is to be accessed at x, either got_h will be absent or PRESENT and set .FALSE.. This gives the user the opportunity to reuse "start-up" computations required for the first instance of x to speed up subsequent products.

2.7 Reverse Communication Information

A positive value of inform%status on exit from EPF_solve indicates that GALAHAD_EPF_solve is seeking further information—this will happen if the user has chosen not to evaluate function or derivative values internally (see Section 2.6). The user should compute the required information and re-enter GALAHAD_EPF_solve with inform%status and all other arguments (except those specifically mentioned below) unchanged.

Possible values of inform%status and the information required are

- 2. The user should compute the objective function value $f(\mathbf{x})$ and the constraint values $\mathbf{c}(\mathbf{x})$ at the point \mathbf{x} indicated in nlp%X. The required values should be set in nlp%f and nlp%c, and data%eval_status should be set to 0. If the user is unable to evaluate $f(\mathbf{x})$ or $\mathbf{c}(\mathbf{x})$ —for instance, if the function is undefined at \mathbf{x} —the user need not set nlp%f or $\mathbf{c}(\mathbf{x})$, but should then set data%eval_status to a non-zero value.
- 3. The user should compute the gradient $\mathbf{g}(\mathbf{x})$ of the objective function and the Jacobian $\mathbf{J}(\mathbf{x})$ of the constraints at the point \mathbf{x} indicated in nlp%X. The value of the i-th component of the gradient should be set in nlp%G(i), for $i=1,\ldots,n$ and those for the Jacobian should be set in nlp%J%val (in the same order as that in which the array indices were given in nlp%J, and $data\%eval_status$ should be set to 0. If the user is unable to evaluate a component of $\mathbf{g}(\mathbf{x})$ or $\mathbf{J}(\mathbf{x})$ —for instance, if a component of the gradient is undefined at \mathbf{x} —the user need not set nlp%G and nlp%J%val, but should then set $data\%eval_status$ to a non-zero value.
- 4. The user should compute the Hessian $\mathbf{H}_{\text{L}}(\mathbf{x},\mathbf{y})$ of the Lagrangian function at the point \mathbf{x} and \mathbf{y} indicated in nlp%X and nlp%Y, respectively. The value l-th component of the Hessian stored according to the scheme input in the remainder of nlp%H (see Section 2.4.2) should be set in nlp%H%val(1), for $l=1,\ldots,$ nlp%H%ne, and data%eval_status should be set to 0. If the user is unable to evaluate a component of $\mathbf{H}_{\text{L}}(\mathbf{x},\mathbf{y})$ —for instance, if a component of the Hessian is undefined at \mathbf{x} and \mathbf{y} —the user need not set nlp%H%val, but should then set data%eval_status to a non-zero value.
- The user should compute the product $\mathbf{H}_{\text{L}}(\mathbf{x},\mathbf{y})\mathbf{v}$ of the Hessian of the Lagrangian function $\mathbf{H}_{\text{L}}(\mathbf{x},\mathbf{y})$ at the point \mathbf{x} and \mathbf{y} indicated in nlp%Y and nlp%Y with the vector \mathbf{v} and add the result to the vector \mathbf{u} . The vectors \mathbf{u} and \mathbf{v} are given in data%U and data%V respectively, the resulting vector $\mathbf{u} + \mathbf{H}_{\text{L}}(\mathbf{x},\mathbf{y})\mathbf{v}$ should be set in data%U and data%eval_status should be set to 0. If the user is unable to evaluate the product—for instance, if a component of the Hessian is undefined at \mathbf{x} and \mathbf{y} —the user need not set nlp%H%val, but should then set data%eval_status to a non-zero value.

2.8 Warning and error messages

A negative value of inform%status on exit from EPF_solve or EPF_terminate indicates that an error has occurred. No further calls should be made until the error has been corrected. Possible values are:

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-1. An allocation error occurred. A message indicating the offending array is written on unit control%error, and the returned allocation status and a string containing the name of the offending array are held in inform%alloc_status and inform%bad_alloc, respectively.

- -2. A deallocation error occurred. A message indicating the offending array is written on unit control%error and the returned allocation status and a string containing the name of the offending array are held in inform%alloc_status and inform%bad_alloc, respectively.
- -3. The restriction nlp%n > 0 or requirement that nlp%H_type contains its relevant string 'DENSE', 'COORDINATE', 'SPARSE_BY_ROWS' or 'DIAGONAL' has been violated.
- -4. The bound constraints are inconsistent.
- -7. The objective function appears to be unbounded from below on the feasible set.
- -9. The analysis phase of the factorization failed; the return status from the factorization package is given in the component inform%factor_status.
- -10. The factorization failed; the return status from the factorization package is given in the component inform%factor_status.
- -11. The solution of a set of linear equations using factors from the factorization package failed; the return status from the factorization package is given in the component inform%factor_status.
- -15. The preconditioner P(x) appears not to be positive definite.
- -16. The problem is so ill-conditioned that further progress is impossible.
- -17. The step is too small to make further impact.
- -18. Too many iterations have been performed. This may happen if control%maxit is too small, but may also be symptomatic of a badly scaled problem.
- -19. The elapsed CPU or system clock time limit has been reached. This may happen if either control%cpu_time_limit or control%clock_time_limit is too small, but may also be symptomatic of a badly scaled problem.
- -82. The user has forced termination of GALAHAD_EPF_solve by removing the file named control%alive_file from unit control%alive_unit.

2.9 Further features

In this section, we describe an alternative means of setting control parameters, that is components of the variable control of type EPF_control_type (see Section 2.4.3), by reading an appropriate data specification file using the subroutine EPF_read_specfile. This facility is useful as it allows a user to change EPF control parameters without editing and recompiling programs that call EPF.

A specification file, or specifie, is a data file containing a number of "specification commands". Each command occurs on a separate line, and comprises a "keyword", which is a string (in a close-to-natural language) used to identify a control parameter, and an (optional) "value", which defines the value to be assigned to the given control parameter. All keywords and values are case insensitive, keywords may be preceded by one or more blanks but values must not contain blanks, and each value must be separated from its keyword by at least one blank. Values must not contain more than 30 characters, and each line of the specifie is limited to 80 characters, including the blanks separating keyword and value.

The portion of the specification file used by EPF_read_specfile must start with a "BEGIN EPF" command and end with an "END" command. The syntax of the specifile is thus defined as follows:

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```
( .. lines ignored by EPF_read_specfile .. )
BEGIN EPF
    keyword value
    .....
    keyword value
END
( .. lines ignored by EPF_read_specfile .. )
```

where keyword and value are two strings separated by (at least) one blank. The "BEGIN EPF" and "END" delimiter command lines may contain additional (trailing) strings so long as such strings are separated by one or more blanks, so that lines such as

```
BEGIN EPF SPECIFICATION

and

END EPF SPECIFICATION
```

are acceptable. Furthermore, between the "BEGIN EPF" and "END" delimiters, specification commands may occur in any order. Blank lines and lines whose first non-blank character is ! or * are ignored. The content of a line after a ! or * character is also ignored (as is the ! or * character itself). This provides an easy manner to "comment out" some specification commands, or to comment specific values of certain control parameters.

The value of a control parameters may be of three different types, namely integer, logical or real. Integer and real values may be expressed in any relevant Fortran integer and floating-point formats (respectively). Permitted values for logical parameters are "ON", "TRUE", ".TRUE.", "T", "YES", "Y", or "OFF", "NO", "N", "FALSE", ".FALSE." and "F". Empty values are also allowed for logical control parameters, and are interpreted as "TRUE".

The specification file must be open for input when EPF_read_specfile is called, and the associated device number passed to the routine in device (see below). Note that the corresponding file is REWINDed, which makes it possible to combine the specifications for more than one program/routine. For the same reason, the file is not closed by EPF_read_specfile.

2.9.1 To read control parameters from a specification file

Control parameters may be read from a file as follows:

```
CALL EPF_read_specfile( control, device )
```

control is a scalar INTENT (INOUT) argument of type EPF_control_type (see Section 2.4.3). Default values should have already been set, perhaps by calling EPF_initialize. On exit, individual components of control may have been changed according to the commands found in the specifile. Specifile commands and the component (see Section 2.4.3) of control that each affects are given in Table 2.1 on the following page.

device is a scalar INTENT (IN) argument of type INTEGER (ip_), that must be set to the unit number on which the specifile has been opened. If device is not open, control will not be altered and execution will continue, but an error message will be printed on unit control%error.

2.10 Information printed

If control*print_level is positive, information about the progress of the algorithm will be printed on unit control*cout. If $control*print_level = 1$, a single line of output will be produced for each iteration of the process. This will include the values of the objective function and the norm of its gradient, the ratio of actual to predicted decrease following the step, the radius of the trust-region and the time taken so far. In addition, if a direct solution of the

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command	component of control	value type
error-printout-device	%error	integer
printout-device	%out	integer
print-level	%print_level	integer
start-print	%start_print	integer
stop-print	%stop_print	integer
iterations-between-printing	%print_gap	integer
maximum-number-of-iterations	%max_it	integer
maximum-number-of-evaluations	%max_eval	integer
alive-device	%alive_unit	integer
update-multipliers-from-iteration	%update_multipliers_itmin	real
update-multipliers-feasibility-tolerance	%update_multipliers_tol	real
infinity-value	%infinity	real
absolute-primal-accuracy	%stop_abs_p	real
relative-primal-accuracy	%stop_rel_p	real
absolute-dual-accuracy	%stop_abs_d	real
relative-dual-accuracy	%stop_rel_d	real
absolute-complementary-slackness-accuracy	%stop_abs_c	real
relative-complementary-slackness-accuracy	%stop_rel_c	real
minimum-relative-step-allowed	%stop_s	real
initial-penalty-parameter	initial_mu	real
penalty-parameter-reduction-factor	mu_reduce	real
minimum-objective-before-unbounded	obj_unbounded	real
try-advanced-start-tolerance	try_advanced_start	real
try-sqp-start-tolerance	try_sqp_start	real
stop-advanced-start-tolerance	stop_advance_start	real
maximum-cpu-time-limit	%cpu_time_limit	real
maximum-clock-time-limit	%clock_time_limit	real
hessian-available	%hessian_available	logical
sub-problem-direct	%subproblem_direct	logical
space-critical	%space_critical	logical
deallocate-error-fatal	%deallocate_error_fatal	logical
alive-filename	%alive_file	character

Table 2.1: Specfile commands and associated components of control.

subproblem has been attempted, the Lagrange multiplier from the secular equation and the number of factorizations used will be recorded, while if an iterative solution has been used, the numbers of phase 1 and 2 iterations will be

If control*print_level ≥ 2 this output will be increased to provide significant detail of each iteration. This extra output includes residuals of the linear systems solved, and, for larger values of control%print_level, values of the variables and gradients. Further details concerning the attempted solution of the models may be obtained by increasing control%TRU_control%print_level, control%SSLS_control%print_level and control%GLTR_control%print-_level, while details about factorizations are available by increasing control%SSLS_control%print_level. See the specification sheets for the packages GALAHAD_GLTR, GALAHAD_SSLS and GALAHAD_TRU for details.

3 **GENERAL INFORMATION**

Use of common: None.

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Workspace: Provided automatically by the module.

Other routines called directly: None.

Other modules used directly: EPF_solve calls the GALAHAD packages GALAHAD_CLOCK, GALAHAD_NLPT, GALAHAD_SYMBOLS, GALAHAD_USERDATA, GALAHAD_SPECFILE, GALAHAD_SMT, GALAHAD_BSC, GALAHAD_MOP, GALAHAD_SSLS, GALAHAD_TRU, GALAHAD_GLTR, GALAHAD_STRINGS, GALAHAD_SPACE, GALAHAD_NORMS, GALAHAD_BLAS_interface, and GALAHAD_LAPACK_interface.

Input/output: Output is under control of the arguments control%error, control%out and control%print_level.

Restrictions: nlp%n > 0 and nlp%H_type \(\inf \' DENSE', 'COORDINATE', 'SPARSE_BY_ROWS', 'DIAGONAL' \).

Portability: ISO Fortran 95 + TR 15581 or Fortran 2003. The package is thread-safe.

4 METHOD

To be updated ... A trust-region method is used. In this, an improvement to a current estimate of the required minimizer, \mathbf{x}_k is sought by computing a step \mathbf{s}_k . The step is chosen to approximately minimize a model $m_k(\mathbf{s})$ of $f(\mathbf{x}_k + \mathbf{s})$ within the intersection of the bound constraints $\mathbf{x}^L \leq \mathbf{x} \leq \mathbf{x}^U$ and a trust region $\|\mathbf{s}_k\| \leq \Delta_k$ for some specified positive "radius" Δ_k . The quality of the resulting step \mathbf{s}_k is assessed by computing the "ratio" $(f(\mathbf{x}_k) - f(\mathbf{x}_k + \mathbf{s}_k))/(m_k(\mathbf{0}) - m_k(\mathbf{s}_k))$. The step is deemed to have succeeded if the ratio exceeds a given $\eta_s > 0$, and in this case $\mathbf{x}_{k+1} = \mathbf{x}_k + \mathbf{s}_k$. Otherwise $\mathbf{x}_{k+1} = \mathbf{x}_k$, and the radius is reduced by powers of a given reduction factor until it is smaller than $\|\mathbf{s}_k\|$. If the ratio is larger than $\eta_v \geq \eta_d$, the radius will be increased so that it exceeds $\|\mathbf{s}_k\|$ by a given increase factor. The method will terminate as soon as $\|\nabla_x f(\mathbf{x}_k)\|$ is smaller than a specified value.

Either linear or quadratic models $m_k(\mathbf{s})$ may be used. The former will be taken as the first two terms $f(\mathbf{x}_k) + \mathbf{s}^T \nabla_x f(\mathbf{x}_k)$ of a Taylor series about \mathbf{x}_k , while the latter uses an approximation to the first three terms $f(\mathbf{x}_k) + \mathbf{s}^T \nabla_x f(\mathbf{x}_k) + \frac{1}{2} \mathbf{s}^T \mathbf{B}_k \mathbf{s}$, for which \mathbf{B}_k is a symmetric approximation to the Hessian $\nabla_{xx} f(\mathbf{x}_k)$; possible approximations include the true Hessian, limited-memory secant and sparsity approximations and a scaled identity matrix. Normally a two-norm trust region will be used, but this may change if preconditioning is employed.

The model minimization is carried out in two stages. Firstly, the so-called generalized Cauchy point for the quadratic subproblem is found—the purpose of this point is to ensure that the algorithm converges and that the set of bounds which are satisfied as equations at the solution is rapidly identified. Thereafter an improvement to the quadratic model on the face of variables predicted to be active by the Cauchy point is sought using either a direct approach involving factorization or an iterative (conjugate-gradient/Lanczos) approach based on approximations to the required solution from a so-called Krlov subspace. The direct approach is based on the knowledge that the required solution satisfies the linear system of equations $(\mathbf{B}_k + \lambda_k \mathbf{I})\mathbf{s}_k = -\nabla_x f(\mathbf{x}_k)$, involving a scalar Lagrange multiplier λ_k , on the space of inactive variables. This multiplier is found by uni-variate root finding, using a safeguarded Newton-like process, by GALAHAD_TRS or GALAHAD_DPS (depending on the norm chosen). The iterative approach uses GALAHAD_GLTR, and is best accelerated by preconditioning with good approximations to \mathbf{B}_k using GALAHAD_PSLS. The iterative approach has the advantage that only matrix-vector products $\mathbf{B}_k \mathbf{v}$ are required, and thus \mathbf{B}_k is not required explicitly. However when factorizations of \mathbf{B}_k are possible, the direct approach is often more efficient.

The iteration is terminated as soon as the Euclidean norm of the projected gradient,

$$\|\min(\max(\mathbf{x}_k - \nabla_x f(\mathbf{x}_k), \mathbf{x}^{\scriptscriptstyle L}), \mathbf{x}^{\scriptscriptstyle U}) - \mathbf{x}_k\|_2$$

is sufficiently small. At such a point, $\nabla_x f(\mathbf{x}_k) = \mathbf{z}_k$, where the *i*-th dual variable z_i is non-negative if x_i is on its lower bound x_i^L , non-positive if x_i is on its upper bound x_i^L , and zero if x_i lies strictly between its bounds.

References:

The generic bound-constrained trust-region method is described in detail in

A. R. Conn, N. I. M. Gould and Ph. L. Toint (2000). Trust-region methods. SIAM/MPS Series on Optimization.

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5 EXAMPLES OF USE

Suppose we wish to minimize the objective function $f(\mathbf{x}) = x_1^2 + x_2^2$ subject to the constraints

$$x_1 + x_2 \ge 1$$

$$x_1^2 + x_2^2 \ge 1$$

$$px_1^2 + x_2^2 \ge 9$$

$$x_1^2 - x_2 \ge 0$$

$$x_2^2 - x_1 \ge 0 \text{ and}$$

$$-50 \le x_1, x_2 \le 50$$

when the parameter p takes the value 9. Starting from the initial guess $\mathbf{x} = (3, 1)$, we may use the following code:

```
PROGRAM GALAHAD_EPF_EXAMPLE ! GALAHAD 5.3 - 2025-05-29 AT 16:45 GMT.
  USE GALAHAD EPF double
                                               ! double precision version
  IMPLICIT NONE
  INTEGER, PARAMETER :: rp = KIND( 1.0D+0 )
                                               ! set precision
  TYPE ( NLPT_problem_type ):: nlp
  TYPE ( EPF_control_type ) :: control
  TYPE ( EPF_inform_type ) :: inform
  TYPE ( EPF_data_type ) :: data
  TYPE ( GALAHAD_userdata_type ) :: userdata
  EXTERNAL :: FC, GJ, HL
  INTEGER :: s
  INTEGER, PARAMETER :: n = 2, m = 5, j_ne = 10, h_ne = 2
  REAL ( KIND = rp ), PARAMETER :: p = 9.0_{rp}
  REAL (KIND = rp), PARAMETER :: infinity = 10.0_rp ** 20
                                                                ! infinity
! start problem data
  nlp%pname = 'HS23'
                                               ! name
  nlp%n = n ; nlp%m = m ; nlp%H%ne = h_ne
                                              ! dimensions
  ALLOCATE( nlp%X(n), nlp%G(n), nlp%X_1(n), nlp%X_u(n))
  ALLOCATE( nlp%C( m ), nlp%C_1( m ), nlp%C_u( m ) )
  nlp%X(1) = 3.0_rp ; nlp%X(2) = 1.0_rp
  nlp%X_1 = -50.0_rp ; nlp%X_u = 50.0_rp
                                               ! variable bounds
  nlp\C_l = 0.0_rp ; nlp\C_u = infinity
                                               ! constraint bounds
! sparse row-wise storage format for the Jacobian
  CALL SMT_put( nlp%J%type, 'SPARSE_BY_ROWS', s ) ! Specify sparse row storage
  ALLOCATE( nlp%J%val( j_ne ), nlp%J%col( j_ne ), nlp%H%ptr( m + 1 ) )
  nlp%J%col = (/ 1, 2, 1, 2, 1, 2, 1, 2, 1, 2 /) ! Jacobian J
  nlp%J%ptr = (/ 1, 3, 5, 7, 9, 11 /)
 sparse co-ordinate storage format for the Hessian
  CALL SMT_put( nlp%H%type, 'COORDINATE', s ) ! Specify co-ordinate storage
  ALLOCATE( nlp%H%val( h_ne ), nlp%H%row( h_ne ), nlp%H%col( h_ne ) )
  nlp%H%row = (/ 1, 2 /)
                                     ! Hessian H
  nlp%H%col = (/ 1, 2 /)
                                      ! NB lower triangle
! problem data complete
  ALLOCATE ( userdata%real ( 1 ) )
                                               ! Allocate space for parameter
  userdata%real( 1 ) = p
                                               ! Record parameter, p
  CALL EPF_initialize( data, control, inform ) ! Initialize control parameters
  control%subproblem_direct = .TRUE.
  control\mbox{max\_it} = 20
  control%max_eval = 100
  control%print_level = 1
  control%stop_abs_p = 1.0D-5
  control % stop_abs_d = 1.0D-5
```

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```
control%stop_abs_c = 1.0D-5
! control%tru_control%print_level = 1
  inform%status = 1
                                              ! set for initial entry
  CALL EPF_solve( nlp, control, inform, data, userdata, eval_FC = FC,
                 eval_GJ = GJ, eval_HL = HL ) ! Solve problem
  IF ( inform%status == 0 ) THEN
                                              ! Successful return
    WRITE( 6, "( ' EPF: ', IO, ' major iterations -',
                                                                             δ
      ' optimal objective value =',
   δ
                                                                             δ
           ES12.4, /, ' Optimal solution = ', ( 5ES12.4 ) )" )
   &
    inform%iter, inform%obj, nlp%X
                                              ! Error returns
    WRITE( 6, "( ' EPF_solve exit status = ', I6 ) " ) inform%status
  END IF
  CALL EPF_terminate( data, control, inform ) ! delete internal workspace
  DEALLOCATE( nlp%X, nlp%H%, nlp%H%val, nlp%H%row, nlp%H%col, userdata%real )
  DEALLOCATE ( nlp%J%val, nlp%J%col, nlp%J%ptr )
  DEALLOCATE( nlp%C, nlp%X_l, nlp%X_u, nlp%C_l, nlp%C_u )
  END PROGRAM GALAHAD_EPF_EXAMPLE
  SUBROUTINE FC ( status, X, userdata, F, C )
  USE GALAHAD_USERDATA_double
  INTEGER, PARAMETER :: rp = KIND( 1.0D+0 )
  INTEGER ( KIND = ip_ ), INTENT( OUT ) :: status
  REAL ( kind = rp ), DIMENSION( : ), INTENT( IN ) :: X
  REAL ( kind = rp ), OPTIONAL, INTENT( OUT ) :: F
  REAL ( kind = rp ), DIMENSION( : ), OPTIONAL, INTENT( OUT ) :: C
  TYPE ( GALAHAD_userdata_type ), INTENT( INOUT ) :: userdata
  REAL ( kind = rp ) :: r
  r = userdata real(1)
  f = X(1) ** 2 + X(2) ** 2
  C(1) = X(1) + X(2) - 1.0_{rp}
  C(2) = X(1) ** 2 + X(2) ** 2 - 1.0_rp
  C(3) = r * X(1) ** 2 + X(2) ** 2 - rp
  C(4) = X(1) ** 2 - X(2)
  C(5) = X(2) ** 2 - X(1)
  status = 0
  END SUBROUTINE FC
  SUBROUTINE GJ ( status, X, userdata, G, J_val )
  USE GALAHAD_USERDATA_double
  INTEGER, PARAMETER :: rp = KIND( 1.0D+0 )
  INTEGER, INTENT( OUT ) :: status
  REAL ( KIND = rp ), DIMENSION( : ), INTENT( IN ) :: X
  REAL ( KIND = rp ), DIMENSION( : ), OPTIONAL, INTENT( OUT ) :: G
  REAL ( KIND = rp ), DIMENSION( : ), OPTIONAL, INTENT( OUT ) :: J_val
  TYPE ( GALAHAD_userdata_type ), INTENT( INOUT ) :: userdata
  REAL ( kind = rp ) :: r
  r = userdata real(1)
  G(1) = 2.0 rp * X(1)
  G(2) = 2.0 rp * X(2)
  J_val(1) = 1.0_rp
  J_val(2) = 1.0_rp
  J_val(3) = 2.0_rp * X(1)
  J_val(4) = 2.0_rp * X(2)
```

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```
J_val(5) = 2.0_rp * r * X(1)
J_val(6) = 2.0_rp * X(2)
J_val(7) = 2.0_rp * X(1)
J_val(8) = -1.0_rp
J_val(9) = -1.0_rp
J_val(10) = 2.0_rp * X(2)
END SUBROUTINE GJ
SUBROUTINE HL ( status, X, Y, userdata, H_val )
USE GALAHAD_USERDATA_double
INTEGER, PARAMETER :: rp = KIND( 1.0D+0 )
INTEGER, INTENT( OUT ) :: status
REAL ( KIND = rp ), DIMENSION( : ), INTENT( IN ) :: X, Y
REAL ( KIND = rp ), DIMENSION( : ), INTENT( OUT ) ::H_val
TYPE ( GALAHAD_userdata_type ), INTENT( INOUT ) :: userdata
REAL ( kind = rp ) :: r
r = userdata%real(1)
H_val(1) = 2.0_rp - 2.0_rp * (Y(2) + r * Y(3) + Y(4))
H_val(2) = 2.0_rp - 2.0_rp * (Y(2) + Y(3) + Y(5))
END SUBROUTINE HL
```

Notice how the parameter p is passed to the function evaluation routines via the real component of the derived type userdata. The code produces the following output:

```
Problem: HS23 (n = 2, m = 5): EPF stopping tolerance = 1.0000E-05
 iter f
                 pr-feas du-feas cmp-slk max mu inner stop cpu time
   1 2.36612966E+01 0.0E+00 1.4E-09 1.8E+01 7.4E+01 7 0 0.00
   2q 2.36612966E+01 0.0E+00 9.7E+00 1.8E+01 3.7E+01
                                                    1
                                                         0 0.02
   2 1.11923551E+00 1.7E+00 1.4E-07 2.5E+00 3.7E+01 20 0 0.02
   3q 1.11923551E+00 1.7E+00 2.1E+00 2.5E+00 1.8E+01 1 0 0.02
   3 9.17057619E-01 2.6E+00 3.6E-08 1.1E-01 1.8E+01 9 0 0.02
   4 1.25713283E+00 2.2E+00 1.4E-08 3.8E-02 9.2E+00 6 0 0.02
   5 1.65916893E+00 1.5E-01 3.5E-07 1.3E-02 4.6E+00 3 0 0.02
   6 1.69374369E+00 1.4E-01 9.5E-11 1.7E-02 2.3E+00 3 0 0.02
   7 1.73335862E+00 1.2E-01 9.0E-09 3.8E-02 1.2E+00
                                                   3 0 0.02
   8 1.87972432E+00 5.3E-02 5.6E-06 1.4E-02 5.8E-01
                                                   4
                                                         0 0.02
   9 1.99742853E+00 1.2E-03 2.1E-07 2.3E-05 2.9E-01
                                                    5
                                                          0
                                                               0.02
||r|| = 2.5307E-01
primal, dual, comp = 1.1817E-03 2.6492E-01 2.5508E-03
  10a 1.99742853E+00 1.2E-03 2.6E-01 2.6E-03 1.4E-01
                                                     1
                                                          0
                                                                0.03
  10 2.00002417E+00 0.0E+00 2.8E-11 2.0E-05 1.4E-01
                                                     3
                                                          0
                                                                0.03
  11q 2.00002417E+00 0.0E+00 9.9E-16 2.0E-05
                                           7.2E-02
                                                     2
                                                          0
                                                                0.03
                                                         0
  11 1.99999986E+00 5.7E-08 2.1E-06 2.2E-23 7.2E-02
                                                   1
                                                                0.03
# function evaluations = 65
# gradient evaluations = 54
# Hessian evaluations = 53
# major iterations
                   = 11
Terminal objective value = 1.99999986147732E+00
Terminal gradient norm = 2.1350E-06
Total time = .03 seconds
EPF: 11 major iterations - optimal objective value = 2.0000E+00
Optimal solution = 1.0000E+00 1.0000E+00
```

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