1 SUMMARY

Let $K$ be an $n \times n$ sparse symmetric saddle-point matrix of the form

$$K = \begin{pmatrix} A & B^T \\ B & -C \end{pmatrix},$$

where $A$ is $(n-m) \times (n-m)$ symmetric positive definite, $B$ is rectangular $m \times (n-m)$ and of full rank ($m < n$), and $C$ is $m \times m$ symmetric positive semi-definite. HSL_M130 computes a signed incomplete Cholesky factorization. That is, a factorization of the form $LDL^T$, where $L$ is lower triangular and $D$ is diagonal with $n-m$ positive entries and $m$ negative entries. The matrix $K$ is optionally reordered, scaled and, if necessary, shifted to avoid breakdown of the factorization so that the $LDL^T$ incomplete factorization of the matrix

$$\tilde{K} = SQ^T \begin{pmatrix} A & B^T \\ B & -C \end{pmatrix} QS + \begin{pmatrix} \alpha(1)I & 0 \\ 0 & -\alpha(2)I \end{pmatrix}$$

is computed, where $Q$ is a permutation matrix, $S$ is a diagonal scaling matrix and $\alpha(1 : 2)$ are non-negative shifts.

The incomplete factorization may be used for preconditioning when solving the saddle-point system $Kx = b$. A separate entry performs the preconditioning operation

$$y = Pz$$

where $P = (L D T^T)^{-1}$, with $L = QS^{-1}L$, is the incomplete signed Cholesky factorization preconditioner. An option exists to use $P = (L | D | T^T)^{-1}$ as the preconditioner.

The incomplete factorization is based on a matrix decomposition of the form

$$K = (L + R) \begin{pmatrix} D & 0 \\ 0 & -E \end{pmatrix} \begin{pmatrix} L + R & 0 \\ 0 & E \end{pmatrix} - E,$$

(1.1)

where $L$ is lower triangular with unit diagonal entries, $R$ is a strictly lower triangular matrix with small entries that is used to stabilize the factorization process, $D$ is a diagonal matrix, and $E$ has the form

$$E = RDR^T + F + FT^T,$$

(1.2)

where $F$ is strictly lower triangle. $E$ is not computed explicitly and all terms in $F$ are ignored, while the matrix $R$ is used in the computation of $L$ but is then discarded. The user controls the dropping of small entries from $L$ and $R$ and the maximum number of entries within each column of $L$ and $R$ (and thus the amount of memory for $L$ and the intermediate work and memory used in computing the incomplete factorization).

Note: If an incomplete Cholesky factorization preconditioner for a symmetric positive-definite system is required, HSL_M128 should be used.

ATTRIBUTES — Version: 1.3.1 (2 April 2015). Interfaces: Fortran, MATLAB. Types: Real (single, double). Calls: KB07, MC61, HSL_MC64, HSL_MC68, HSL_MC69, MC77, copy and (optionally) METIS_NODEND. Language: Fortran 2003 subset (F95+TR155581). Date: March 2014. Origin: J. A. Scott, STFC Rutherford Appleton Laboratory and M. Tůma, Institute of Computer Science, Academy of Sciences of the Czech Republic. Remark: The development of this package was partially supported by EPSRC grant EP/I013067/1 and by Grant Agency of the Czech Republic grant P201/13-06684S.

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2 HOW TO USE THE PACKAGE

2.1 Calling sequences

Access to the package requires a USE statement

Single precision version
  use hsl_mi30_single
Double precision version
  use hsl_mi30_double

If it is required to use more than one module at the same time, the derived types (see Section 2.2) must be renamed in one of the USE statements.

The following procedures are available to the user:

(a) mi30_factorize computes an incomplete signed Cholesky factorization.

(b) mi30_precondition performs the preconditioning operation \( y = Pz \), where \( P \) is the incomplete factorization preconditioner computed by mi30_factorize.

(c) mi30_solve solves the system \( \bar{L}Dy = SQ^Tz \) (or \( \bar{L}|D|y = SQ^Tz \) or \( \bar{L}^TS^{-1}Q^Ty = z \)), where \( \bar{L} \) is the incomplete factor computed by mi30_factorize and \( |D| \) has entries \( |d_{ij}| \).

(d) mi30_finalise frees memory that has been allocated by mi30_factorize.

2.2 The derived data types

For each problem, the user must employ the derived types defined by the module to declare scalars of the types mi30_keep, mi30_control and mi30_info. The following pseudocode illustrates this.

```
use hsl_mi30_double
...
type (mi30_keep) :: keep
type (mi30_control) :: control
type (mi30_info) :: info
...
```

The components of mi30_control and mi30_info are explained in Sections 2.5 and 2.6. The components of mi30_keep are used to pass data between the subroutines of the package and must not be altered by the user.

2.3 METIS

The HSL_MI30 package optionally uses the METIS graph partitioning library available from the University of Minnesota website. If METIS is not available, the user must link with the supplied dummy subroutine METIS_NodeND. In this case, the METIS ordering option will not be available to the user and, if selected, mi30_factorize will return with an error.

Important: At present, HSL_MI30 only supports METIS version 4, not the latest version 5 releases.
2.4 Argument lists and calling sequences

2.4.1 Optional arguments

We use square brackets [ ] to indicate OPTIONAL arguments. In each call, optional arguments follow the argument info. Since we reserve the right to add additional optional arguments in future releases of the code, we strongly recommend that all optional arguments be called by keyword, not by position.

2.4.2 Integer and package types

INTEGER denotes default INTEGER and INTEGER(long) denotes INTEGER(kind=selected_int_kind(18)). We use the term package type to mean default real if the single precision version is being used and double precision real for the double precision version.

2.4.3 To compute a signed incomplete Cholesky factorization

To compute a signed incomplete Cholesky factorization, the lower triangular part of the matrix $K$ must be held in compressed column storage and a call of the following form must be made:

```fortran
    call mi30_factorize(n, m, ptr, row, val, lsize, rsize, keep, control, info[, scale, perm])
```

- $n$ is an INTENT(IN) scalar of type INTEGER that must hold the order of the matrix $K$. Restriction: $2 \leq n$.

- $m$ is an INTENT(IN) scalar of type INTEGER that must hold the order of the $(2, 2)$ block $-C$. Restriction: $1 \leq m \leq n-1$.

- $ptr$ is an INTENT(INOUT) rank-1 array of type INTEGER and size $n+1$. $ptr(j)$ must be set by the user so that $ptr(j)$ is the position in $row$ of the first entry in column $j$ and $ptr(n+1)$ must be set to one more than the number of matrix entries being input by the user. $ptr$ is only changed on exit if control%check is set to .true. (the default) and duplicates and/or out-of-range indices are found.

- $row$ is an INTENT(INOUT) rank-1 array of type INTEGER and size at least $ptr(n+1)-1$. It must hold the row indices of the entries of the lower triangular part of $K$ with the row indices for the entries in column 1 preceding those for column 2, and so on. Within each column, the row indices must be in increasing order (so that all entries of the column in the $(1,1)$ block $A$ must precede those in the $(2,2)$ block $-C$). The diagonal entry in the first $n-m$ columns must be present (but may have value zero). If control%check is set to .true. (the default), $row$ is checked for errors and duplicates and out-of-range indices are removed; otherwise, $row$ is unchanged.

- $val$ is an INTENT(INOUT) rank-1 array of package type and size at least $ptr(n+1)-1$. $val(k)$ must hold the value of the entry in $row(k)$. If control%check is set to .true. (the default), on exit duplicates are summed and out-of-range indices removed; otherwise, $val$ is unchanged.

- $lsize$ is an INTENT(IN) scalar of type INTEGER that determines the maximum number of fill entries within each column of the incomplete factor $L$. In general, increasing $lsize$ improves the quality of the preconditioner but increases the time to compute and then apply the preconditioner (see Section 4). Values less than 0 are treated as 0.

- $rsize$ is an INTENT(IN) scalar of type INTEGER that determines the maximum number of entries within each column of the strictly lower triangular matrix $R$ that is used in the computation of the preconditioner. A rank-1 array of type INTEGER and a rank-1 array of package type each of size $rsize \times n$ are allocated internally to hold $R$. Thus the amount of memory used, as well as the amount of work involved in computing the preconditioner, depends on $rsize$. Setting $rsize > 0$ generally leads to a higher quality preconditioner than using $rsize = 0$ (and $rsize \geq lsize$ is generally recommended). Values less than 0 are treated as 0.
keep is an INTENT(OUT) scalar of type mi30_keep. It is used to hold data about the problem being solved and must be passed unchanged to mi30_precondition. The following components may be of interest to the user:

- **fact_dinv** is an allocatable rank-1 array of package type. On exit, it is allocated to have size n and fact_dinv(k) holds the value of the k-th diagonal entry of D−1.

- **fact_ptr** is an allocatable rank-1 array of type INTEGER(long). On exit, it is allocated to have size n+1, fact_ptr(j) holds the position in fact_row of the first entry in column j of the computed factor L and ptr(n+1) is set to one more than the number of entries in L.

- **fact_row** is an allocatable rank-1 array of type INTEGER. On exit, the first fact_ptr(n+1)-1 entries hold the row indices of the entries of the computed factor L, with the row indices for the entries in column 1 preceding those for column 2, and so on.

- **fact_val** is an allocatable rank-1 array of package type. On exit, fact_val holds the values of the entries in the computed factor L such that fact_val(k) is the value of the entry in fact_row(k).

- **scale** is an allocatable rank-1 array of package type. On exit, if control%iscale > 0, it is allocated to have size n and holds the scaling factors for A.

- **invp** is an allocatable rank-1 array of type INTEGER. On exit, if control%iorder > 0, it is allocated to have size n and specifies the permutation such that the j-th column of the permuted matrix Q^T K Q is the invp(j)-th column of K (that is, invp(j) is the index of the j-th pivot).

- **perm** is an allocatable rank-1 array of type INTEGER. On exit, if control%iorder > 0, it is allocated to have size n and specifies the elimination ordering such that perm(i) holds the position of i-th column of K in the elimination order.

**control** is an INTENT(IN) scalar of type mi30_control (see Section 2.5).

**info** is an INTENT(OUT) scalar of type mi30_info. Its components provide information about the execution of the subroutine, as explained in Section 2.6.

- **scale** is an optional INTENT(IN) rank-one array of package type and size n that must be present if control%order=5. In this case, scale must be set by the user to hold scaling factors for A.

- **perm** is an optional INTENT(IN) rank-one array of type INTEGER and size n that must be present if control%iorder=3. In this case, the user must supply an elimination ordering such that perm(i) holds the position of the i-th column of K in the elimination order. A column with index j > n - m should only be ordered after all the columns with index i ≤ n - m for which k_{ij} ≠ 0 have been ordered; if this condition is not satisfied, a modified ordering will be used (and returned in keep%perm).

**2.4.4 To perform preconditioning operations**

The signed incomplete Cholesky factorization preconditioner may be applied to compute y = Pz by making a call as follows.

```fortran
    call mi30_precondition(job, n, keep, z, y, info)
```

**job** is an INTENT(IN) scalar of type INTEGER that must be set as follows:

1. If P = (L D L^T)^{-1} is to be used as the preconditioner (L = Q S^{-1} L).
2. If P = (L |D| L^T)^{-1} is to be used as the preconditioner.

n, keep: must be unchanged since the call to mi30_factorize.
z is an INTENT(IN) rank-1 array of package type and size n. It must be set by the user to hold the vector z to which
the incomplete factorization preconditioner P is to be applied.

y is an INTENT(OUT) rank-1 array of package type and size n. On exit, y contains Pz.

info is an INTENT(INOUT) scalar of type mi30_info. Only the component info%flag is accessed (see Section 2.6).

2.4.5 To perform solve operations

The system $L Dy = SQ^T z$, or $L |D| y = SQ^T z$ or $L^T S^{-1} Q^T y = z$ may be solved by making a call as follows.

call mi30_solve(job, n, keep, z, y, info)

job is an INTENT(IN) scalar of type INTEGER that must be set as follows:

1 if the solution of $L Dy = SQ^T z$ is required,
2 if the solution of $L |D| y = SQ^T z$ is required,
3 if the solution of $L^T S^{-1} Q^T y = z$ is required.

n, keep: must be unchanged since the call to mi30_factorize.

z is an INTENT(IN) rank-1 array of package type and size n. It must be set by the user to the right-hand side vector z.

y is an INTENT(OUT) rank-1 array of package type and size n. On exit, y contains the solution vector y.

info is an INTENT(INOUT) scalar of type mi30_info. Only the component info%flag is accessed (see Section 2.6).

2.4.6 The finalisation subroutine

Once all other calls are complete for a problem or after an error return, a call should be made to free memory allocated
by hsl_mi30_factorize using a call to mi30_finalise.

call mi30_finalise(keep, info)

keep is an INTENT(INOUT) scalar of type mi30_keep that must be passed unchanged. On exit, allocatable components
will have been deallocated.

info is an INTENT(INOUT) scalar of type mi30_info. Only the components info%flag and info%stat are accessed (see Section 2.6).

2.5 The control derived data type

The derived data type mi30_control is used to hold controlling data; it is used by mi30_factorize only. The components are automatically given default values in the definition of the type.

Components that control printing

unit_error is a scalar of type INTEGER with default value 6 that is used as the output stream for error messages. If it
is negative, these messages will be suppressed.

unit_warning is a scalar of type INTEGER with default value 6 that is used as the output stream for warning messages.
If it is negative, these messages will be suppressed.
Components that control the initial and subsequent choice of the shifts $\alpha(1:2)$.
Note that the aim is to choose the shifts to be as small as possible to avoid breakdown of the Cholesky factorization process (see Section 4).

- **alpha** is a rank-1 array of package type and size 2 with default values (0.0, 0.0) that holds the initial shifts $\alpha(1:2)$. Values less than zero are treated as zero.

- **lowalpha** is a scalar of package type with default value 0.001 that controls the choice of the shift in the event of a breakdown. Values less than or equal to zero are treated as the default.

- **maxshift** is a scalar of type INTEGER with default value 3 that controls the maximum number of times the shift can be decreased after a successful factorization with a positive shift. See Section 4 for details. Limiting maxshift may reduce the factorization time but may result in a poorer quality preconditioner.

- **shift_factor** is a scalar of package type with default value 4.0 that controls how rapidly a shift is increased after a breakdown. See Section 4 for details. Increasing shift_factor may reduce the factorization time but may result in a poorer quality preconditioner. Values less than one are treated as the default.

- **shift_factor2** is a scalar of package type with default value 4.0 that controls how rapidly a shift is decreased after a successful factorization with a positive shift. See Section 4 for details. Values less than one are treated as the default.

- **small** is a scalar of type REAL. Any pivot whose modulus is less than small is treated as zero and, if such a pivot is encountered, the factorization breaks down, a shift is increased and the factorization restarted. The default in the double version is $10^{-20}$ and in the single version is $10^{-12}$.

Components that control the dropping of small entries

- **tau1** and **tau2** are scalars of package type with default values 0.001 and 0.0001. They control the dropping of entries from $L$ and $R$. In the computation of the incomplete factorization, entries of magnitude less than $|\text{tau1}|$ are dropped from $L$; those that are at least $|\text{tau2}|$ but less than $|\text{tau1}|$ may be included in $R$ while those less than $|\text{tau2}|$ are dropped from $R$. If $|\text{tau2}| > |\text{tau1}|$, tau2=0.0 is used.

Other components

- **check** is a scalar of type LOGICAL with default value .true.. If .true., the matrix data is checked for errors and the cleaned matrix (duplicates are summed, out-of-range entries discarded and, within each column, the entries are ordered by increasing row index) overwrites the user-supplied data in ptr, row and val. Otherwise, no checking of the matrix data is carried out (it is important to note that any out-of-order entries or out-of-range entries or duplicates may cause HSL_MI30 to fail in an unpredictable way) and so it is recommended that the matrix data is checked.

- **iorder** is a scalar of type INTEGER with default value 6 that indicates the ordering that is required. The chosen ordering is computed and then post-processed (see Section 4). Options available are:
  
  - $\leq 0$ no ordering.
  - 1 A reverse Cuthill-McKee (RCM) ordering (computed using MC61) is used.
  - 2 An approximate minimum degree (AMD) ordering (computed using HSL_MG68) is used.
  - 3 User-supplied ordering is used.
  - 4 The rows are ordered by ascending degree.
METIS (nested dissection) ordering with default settings is used. If METIS is not supplied and this option is requested, the routine will return immediately with an error.

6 A Sloan profile reduction ordering (computed using MC61) is used. This is the default.

If iorder > 6, the default is used.

iscale is a scalar of type INTEGER with default value 1 that indicates the scaling that is required. Options available are:

\[ \leq 0 \] No scaling.
\( 1 \) Scaling generated using the \( l_2 \)-norm of the columns of \( A \). This is the default.
\( 2 \) Scaling generated by applying the iterative method of the package MC77 for one iteration in the infinity norm and three iterations in the one norm (equilibration ordering).
\( 3 \) Scaling generated from a weighted bipartite matching using the package HSL_MC64.
\( 4 \) Diagonal scaling is used.
\( 5 \) User-supplied scaling is used. The user must supply scaling factors for \( A \).

If iscale > 5, the default is used.

rrt is a scalar of type LOGICAL with default value .false. that is used to control whether the entries of \( RR^T \) (see (1.2)) that cause no additional fill-in in (1.1) are allowed (rsize > 0 only). Allowing such entries can improve the quality of the preconditioner (although this is not guaranteed) but at some additional computational cost in the factorization process. If rrt = .true. such entries are allowed; otherwise, they are not allowed.

2.6 The derived data type for holding information

The derived data type mi30_info is used to hold information from the execution of mi30_factorize. The components are:

alpha is a rank-1 array of package type and size 2 that holds the final shifts (it is set to zero if no shifts are used).

band_after is a scalar of type INTEGER. If control%iorder=1 or 6, band_after holds the semibandwidth of \( A \) after reordering; otherwise, it is set to 0.

band_before is a scalar of type INTEGER. If control%iorder=1 or 6, band_before holds the semibandwidth of \( A \) before reordering; otherwise, it is set to 0.

dup is a scalar of type INTEGER that holds the number of duplicated indices removed from row.

flag is a scalar of type INTEGER that gives the exit status of the algorithm (details in Section 2.7).

flag61 is a scalar of type INTEGER that holds the exit status on return from MC61 (and is set to 0 if MC61 is not used).

flag64 is a scalar of type INTEGER that holds the exit status on return from HSL_MC64 (and is set to 0 if HSL_MC64 is not used).

flag68 is a scalar of type INTEGER that holds the exit status on return from HSL_MC68 (and is set to 0 if HSL_MC68 is not used).

flag77 is a scalar of type INTEGER that holds the exit status on return from MC77 (and is set to 0 if MC77 is not used).

nrestart is a scalar of type INTEGER that holds the number of restarts (after reducing a shift).
nshift is a scalar of type INTEGER that holds the number of non-zero shifts used.

oor is a scalar of type INTEGER that holds the number of out-of-range indices removed from row.

profile_after is a scalar of package type. If control%:iorder=1 or 6, profile_after holds the profile of A after reordering; otherwise, it is set to 0.0.

profile_before is a scalar of package type. If control%:iorder=1 or 6, profile_before holds the profile of A before reordering; otherwise, it is set to 0.0.

size_r is a scalar of type INTEGER(long) that holds the size of the integer and real arrays that are used during the factorization to hold \( R \).

stat is a scalar of type INTEGER that holds the Fortran stat parameter.

### 2.7 Warning and error messages

A successful return from a subroutine in the package is indicated by info%:flag having the value zero. A negative value is associated with an error message that by default will be output on unit control%:unit_error.

Possible negative values are:

-1 memory allocation failed. The stat parameter is returned in info%:stat.

-2 The array row is too small.

-3 The array val is too small.

-4 Either \( n \) or \( m \) is out of range (\( n<2, m<1 \) or \( m>n/2 \)).

-5 Error in the array ptr.

-6 One or more diagonal entries in the (1,1) \( A \)-block is missing.

-7 Unexpected error returned by MC77. The MC77 exit status is returned in info%:flag77.

-8 Unexpected error returned by HSL_MC64. The HSL_MC64 exit status is returned in info%:flag64.

-9 HSL_MC64 has found that \( K \) is structurally singular.

-10 The optional argument scale is not present when it should be.

-11 The optional argument perm is either not present when it should be or it does not hold a permutation.

-12 Unexpected error returned by MC61. The MC61 exit status is returned in info%:flag61. Note that, if the matrix has not been checked for errors and there are duplicated or out-of-range entries in row, mc61 will return an error flag of \(-4\) and the computation will terminate.

-13 Unexpected error returned by HSL_MC68. The HSL_MC68 exit status is returned in info%:flag68. Note that this error is returned if METIS ordering has been requested (control%:iorder=5) but METIS is not linked.

-14 Memory deallocation failed. The stat parameter is returned in info%:stat.

-15 All entries in one or more columns are out of range.

-16 One or more of the diagonal entries of the (2,2) block \(-C\) is positive.
Positive values for info%flag are associated with a warning and can only be returned by mi30_factorize. Possible positive values are:

+1 Out-of-range indices have been removed from row. The number of such entries is given in info%oor.

+2 Duplicated entries were found in row; these have been removed and the corresponding entries in val have been summed. The number of such entries is given in info%dup.

+3 A warning has been issued by HSL_MC64 that the computed scaling factors are large and may cause overflow when used to scale the matrix. No scaling is used.

+4 A warning has been issued by MC61. The MC61 exit status is returned in info%flag61.

3 GENERAL INFORMATION

Input/output: Error messages on unit control%lp and warning and diagnostic messages on units control%wp and control%wp, respectively. These have default value 6; printing of these messages is suppressed if the relevant unit number is negative or if print_level is negative.

Restrictions: 1\leq m< n.

4 METHOD

mi30_factorize starts by optionally checking the matrix data for errors; this is done using HSL_MC69. Checking removes out-of-range entries, sums duplicates, and reorders the entries within each column by increasing row index. A scaling and/or ordering is then optionally computed; HSL packages are used to do this. Unless a problem is known to be well scaled, scaling is highly recommended. We impose a constraint on the ordering: a pivot corresponding to a variable i in the (2,2) block \(-C\) can only be eliminated once all the variables that corresponding to the entries in column j with row index \(j \leq n - m\) have been eliminated (in graph terms, a C-node can only be eliminated once all its A-node neighbours have been eliminated). Thus once an ordering has been computed using, for example, the Sloan algorithm, it is modified to satisfy the above constraint before the factorization begins.

A left-looking sparse Cholesky algorithm is used to compute the signed incomplete factorization, one column at a time. The parameters lsize and rsize control the amount of memory used as well as the amount of work involved in computing the factorization. lsize controls the number of entries in the computed incomplete factor L (at most lsize fill entries are permitted in each column of L) and rsize limits the number of entries in each column of the matrix R. If rsize=0 and control%tau1=0, the incomplete factorization is essentially that of [1]. However, it is generally advantageous (in terms of the quality of the preconditioner) to use rsize>0. Increasing lsize and/or rsize increases the cost of the factorization (in terms of time and memory). Furthermore, increasing lsize leads to a denser incomplete factorization (but one that is, in general, a better preconditioner), increasing the cost of each call to mi30_precondition and mi30_solve. Values of lsize and rsize equal to 10 is often a reasonable choice but, if the preconditioner is to be used for many problems, it may be worthwhile to experiment with a range of values to try and get the best overall performance; smaller values may be used if the memory available is limited or larger values may be used to try and obtain a higher quality preconditioner.

Dropping parameters control%tau1 and control%tau2 are used to further sparsify L and R, respectively. As each column of L is computed, entries of absolute value less than control%tau1 are dropped. These may be included in R but entries less than control%tau2 are dropped from R.

In the event of breakdown within the factorization (that is, a pivot is encountered that is smaller in absolute value than control%small), a diagonal shift is used. If the breakdown occurs for a pivot in the A-block, a shift \(\alpha(1)\) is used; if breakdown occurs in the (2,2) block \(-C\), a shift \(-\alpha(2)\) is used (\(\alpha(1:2) > 0\)). It is important to try and use
as small a shift as possible but also to limit the number of breakdowns. The user can supply initial shifts $\alpha_0(1:2)$. If breakdown occurs, a larger shift
\[
\alpha_1(j) = \max(control\%lowalpha, \alpha_0(j) \times control\%shift\_factor),
\]
is tried, where $j = 1$ if breakdown is in the $A$-block and $j = 2$ otherwise. The process continues until an incomplete factorization of
\[
\bar{K} = SQ^T \begin{pmatrix} A & B^T \\ B & -C \end{pmatrix} QS + \begin{pmatrix} \alpha(1)I & 0 \\ 0 & -\alpha(2)I \end{pmatrix}
\]
is successful. If breakdown occurs at the same (or nearly the same) stage of the factorization for two successive shifts, $\alpha(j)$ is increased by a factor of $2 \times control\%shift\_factor$. Conversely, if $\alpha_k(j) = control\%lowalpha(j)$, to prevent an unnecessarily large shift from being used, the shift is decreased by setting
\[
\alpha_{k+1}(j) = \alpha_k(j)/control\%shift\_factor2,
\]
and applying the incomplete factorization algorithm to
\[
\bar{K}_{k+1} = SQ^T \begin{pmatrix} A & B^T \\ B & -C \end{pmatrix} QS + \begin{pmatrix} \alpha_{k+1}(1)I & 0 \\ 0 & -\alpha_{k+1}(2)I \end{pmatrix}.
\]
If this factorization is also breakdown free, the process is repeated (up to $control\%maxshift$ times). In all cases, the values of the final shifts are returned to the user in info\%alpha(1:2), along with the number of shifts tried and the number of restarts (info\%nrestart).

For further details, see [1] and [2].

References:


5 EXAMPLE OF USE

Suppose we wish to use preconditioned GMRES to solve the linear system $Kx = b$ with
\[
K = \begin{pmatrix}
4 & 0 & 1 & -1 \\
0 & 3 & 0 & 2 \\
1 & 0 & 4 & 0 \\
-1 & 2 & 0 & -1
\end{pmatrix}
\]
and
\[
b = \begin{pmatrix}
4 \\
5 \\
5 \\
0
\end{pmatrix}.
\]

We may use the following code:

```fortran
program mi30_spec_double
    use hsl_mi30_double
    implicit none
    K = reshape((/ 4, 0, 1, -1, 0, 3, 0, 2, 1, 0, 4, 0, -1, 2, 0, -1 /), shape(K))
    b = reshape((/ 4, 5, 5, 0 /), shape(b))
end program mi30_spec_double
```
integer, parameter :: wp = kind(1.0d0)
integer, parameter :: long = selected_int_kind(18)

type(mi30_control) :: control
type(mi30_info) :: info
type(mi30_keep) :: keep

integer, allocatable :: ptr(:), row(:)
real(wp), allocatable :: h(:, :), val(:), w(:, :)

! Arrays and scalars required by the GMRES code mi24
real(wp) :: resid
real(wp) :: cntl24(4), rsave24(9)
integer :: icntl24(8), isave24(17), info24(4)
logical :: lsave24(4)

integer :: iact, locy, locz, lsize, m, m_restart, n, nz, rsize

! Read in the matrix data
read (5, *) n, m, nz

! Choose restart parameter for GMRES
m_restart = 10

! Allocate arrays for matrix and for GMRES
allocate (ptr(n+1), row(nz), val(nz), &
        w(n, m_restart+7), h(m_restart+1, m_restart+2))
read (5, *) ptr(1:n+1)
read (5, *) row(1:nz)
read (5, *) val(1:nz)
read (5, *) w(1:n, 1) ! Right-hand side array

! Choose lsize and rsize
lsize = 1
rsize = 1

control%iorder = 0 ! use supplied order
control%iscale = 0 ! do not scale

! Compute the preconditioner
call mi30_factorize(n, m, ptr, row, val, lsize, rsize, keep, control, info)

if (info%flag.lt.0) then
    write (*, '(a,i4)') ' Unexpected error from mi30_factorize. flag = ', info%flag
    call mi30_finalise(keep, info)
    stop
end if

! Prepare to use the GMRES code mi24 with preconditioning
call mi24id(icntl24, cntl24, isave24, rsave24, lsave24)
icntl24(3) = 2 ! right preconditioning

iact = 0
do
  call MI24AD(iact, n, m_restart, w, size(w, 1), locy, locz, h, &
    size(h, 1), resid, icntl24, cntl24, info24, isave24, &
    rsave24, lsave24)
  select case(iact)
    case(-11) ! Error
      write (*,'(a,i4)') ' Unexpected error from mi24. flag = ',info24(1)
      exit
    case(1) ! convergence achieved
      write (*,'(a,i3,a)') &
        ' GMRES Convergence in ',info24(2),' iteration(s)'
      write (*,'(a)') ' Solution = '
      write (*,'(5es12.4)') w(1:n,2)
      exit
    case(2) ! Form y = Kz
      call mxmult(n,ptr,row,val,w(1:n,locz),w(1:n,locy))
    case(4) ! Preconditioner
      call mi30_precondition(1, n, keep, w(1:n,locz), w(1:n,locy), info)
  end select
  end do
  call mi30_finalise(keep,info)
contains
!*****************************************************************************
! sparse matrix-vector multiplication y=K*x.
! Lower triangle of sparse matrix K held.

subroutine mxmult(n,ptr,row,val,x,y)
  real(wp), parameter :: zero = 0.0_wp

  integer, intent(in) :: n
  integer, intent(in) :: ptr(n+1),row(:)
  real(wp), intent(in) :: val(:),x(n)
  real(wp), intent(out) :: y(n)

  integer:: i,j,k
  real(wp) :: sum

  y = zero
do i = 1,n
    sum = zero
    do j = ptr(i),ptr(i+1)-1
        k = row(j)
        if (k.ne.i) y(k) = y(k) + val(j)*x(i)
        sum = sum + val(j)*x(k)
    end do
    y(i) = y(i) + sum
end do
end subroutine mxmult

end program mi30_spec_double

With the input data:

4 1 7
1 4 6 7 8
1 3 4 2 4 3 4
4. 1. -1. 3. 2. 4. -1.
4. 5. 5. 0.

we obtain the following output:

GMRES Convergence in 1 iteration(s)
Solution =
1.0000E+00 1.0000E+00 1.0000E+00 1.0000E+00