

DAO Office Note 96-02

Office Note Series on Global Modeling and Data Assimilation

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Documentation of the Physical-space Statistical Analysis System (PSAS) Part I: The Conjugate Gradient Solver Version PSAS-1.00

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February 1996

Abstract

This document describes Version 1 of the conjugate gradient solver component of DAO's Physical-space Statistical Analysis System (PSAS). An overview of the general PSAS algorithm is presented, followed by an outline of the pre-conditioned conjugate gradient algorithm, and its implementation in PSAS. A description of the main Fortran 90 subroutines related to the conjugate gradient solver is given, with the source code listed in the Appendix.

This Office Note focuses on a particular aspect of the PSAS algorithm, namely the conjugate gradient solver. The details of the observation and forecast error covariance modeling, the strategies for parallelization and domain decomposition, data flow and user interface will be described in subsequent DAO Office Notes. The emphasis of this document is on software design and implementation, and not on the scientific aspects of PSAS which will be documented elsewhere. An on-line version of this document can be obtained from

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Contents

Abstract	iii
1 Introduction	1
2 Overview of PSAS	1
3 Overview of the Conjugate Gradient Algorithm	3
3.1 General Search Directions	3
3.2 The Steepest Descent Algorithm	4
3.3 Conjugate Gradients	5
4 Choice of pre-conditioner in PSAS	6
5 Fortran 90 implementation of the PSAS Conjugate Gradient Solver	8
5.1 The main PSAS driver: getA1all()	9
5.2 Getting ready for the conjugate gradient: solve4x()	9
5.3 The main conjugate gradient driver: cg_main()	10
5.4 Pre-conditioner level 2: cg_level2()	10
5.5 Pre-conditioner level 1: cg_level1()	10
6 Concluding remarks	11
Acknowledgments	11
A Appendix: PSAS Conjugate Gradient Solver prologues and source code	12
Appendix: PSAS CG prologue and source code	12
A.1 getA1all()	12
A.2 getA1all0()	21
A.3 solve4x()	27
A.4 solve4x0()	32
A.5 cg_main()	34
A.6 cg_level2()	42
A.7 cg_level1()	51
A.8 cg_blocks()	59
A.9 cg_level0()	62
References	65

1 Introduction

The central mission of the Data Assimilation Office (DAO) is to develop a state-of-the-art Data Assimilation System capable of assimilating relevant remotely-sensed data from the Earth Observing System (EOS) platforms, as well as global atmospheric data from the other observing systems. The Physical-space Statistical Analysis System (PSAS) is a component of the Goddard EOS Data Assimilation System (GEOS/DAS) which implements a global statistical interpolation algorithm in physical rather than spectral space. This analysis system is a successor to our current *Optimal Interpolation*-based system (Pfaendtner *et al.* 1995) used to produce the GEOS-1 Multiyear assimilation (Schubert *et al.* 1993, 1995a,b). An overview of PSAS and comparisons with the Optimal Interpolation System used in Version 1 of GEOS/DAS can be found in da Silva *et al.* (1995), while some computational aspects of PSAS are discussed in Guo and da Silva (1995).

The purpose of this report is to document the software implementation of the global conjugate gradient solver in PSAS. The details of the observation and forecast error covariance modeling, the strategies for parallelization and domain decomposition, data flow and user interface will be described in subsequent DAO Office Notes.

The organization of this document is as follows. In section 2 the mathematical formulation of PSAS is introduced, with a brief overview of the whole algorithm. Section 3 describes the numerical aspects of the pre-conditioned conjugate gradient algorithm adopted in PSAS. The actual pre-conditioners used in PSAS are introduced in section 4, while section 5 provides an overview of the tasks performed in each major conjugate gradient routine. The actual Fortran 90 source code along with prologues appear in the Appendix. In the *acknowledgments* we present brief historical notes on PSAS design and development at DAO.

2 Overview of PSAS

One of the main design goals of PSAS is to provide a flexible analysis system for the assimilation of several new data types available during the EOS period. In addition, PSAS must provide the framework to test advanced forecast error covariance models, such as generic anisotropic models, and to support research on approximate Kalman filtering and smoothing at DAO. In view of this, PSAS is designed with very few assumptions on the structure of the innovation covariance matrix. Although the current implementation uses a horizontal correlation model which is homogeneous and isotropic, the numerical algorithm takes no advantage of this simplification. In contrast, most current variational systems [ECMWF's 3D-VAR (Courtier *et al.* 1993), NMC's SSI (Parrish and Derber 1992)] depend heavily on this assumption for computational feasibility. Other design goals are the elimination of data selection, and a fully global analysis system which could easily handle non-conventional data types such as satellite radiances.

PSAS implements the statistical analysis equations in physical rather spectral space. The computational advantage of a spectral formulation is tied to the assumption of isotropic horizontal error correlation structures, an assumption we would like to relax in the near future. In addition, PSAS analyses are compatible with the GEOS General Circulation Model which is formulated in grid-point space.

Formulation

Although a non-linear version of PSAS is planned, we focus our discussion on the linear aspects of the algorithm. The non-linear PSAS algorithm in consideration consists of iterations based on linear PSAS solutions.

A statistical interpolation scheme attempts to obtain an *optimal* estimate of the state of the system by combining observations with a forecast model first guess. Under a requirement of optimality the analysis equation is shown to be (*e.g.*, Daley 1991)

$$w_a = w_f + K (w_o - H w_f) \quad (1)$$

$$K = P^j H^T (H P^j H^T + R)^{-1} \quad (2)$$

where $w_a \in \mathbb{R}^n$ is a vector representing the analyzed field, $w_f \in \mathbb{R}^n$ denotes the model forecast first guess, and $w_o \in \mathbb{R}^p$ is the observational vector. The operator H is a generalized interpolation operator which transforms model variables into observables. The matrix $K = P^j H^T (H P^j H^T + R)^{-1}$ is the so-called *gain* or the weights of the analysis. Typically, the number of model degrees of freedom is $n \sim 10^6$ and the current observing system has $p \sim 10^5$. The analysis equations are solved approximately by our OI system: for each grid point the weights in eq. (2) are computed with a reduced number of gridpoints $p' \ll p$, and eq. (1) is used to obtain the analyzed field. This method is clearly not feasible if all observations are to be retained. The algorithm in PSAS consists of solving one $p \times p$ linear system for the quantity y

$$(H P^j H^T + R) y = w_o - H w_f \quad (3)$$

and subsequently obtaining the analyzed state w_a from the equation

$$w_a = w_f + P^j H^T y \quad (4)$$

which is a matrix-vector multiply plus a vector addition, requiring no iterations. The intermediate vector y will be referred to as the *partially weighted innovations*. The linear system (3) is solved by a conjugate gradient algorithm which is documented in subsequent sections.

For typical correlation models the innovation matrix $M = H P^j H^T + R$ is not sparse, although entries associated with grid points over several correlation lengths are negligibly small. In order to introduce some sparseness in M and save computational effort, zeros are introduced in M for entries corresponding to observational points distant by more than 6,000 km. For computational convenience, the sphere is divided in N regions, and matrix blocks associated with regions distant by more than 6,000 km are set to zero. For the sake of consistency and numerical stability, the tail of the correlation function must be adjusted to exactly go to zero beyond a certain distance, usually 6,000 km. For information on the construction of spatially limited correlation functions see Gaspari and Cohn (1996).

Clearly, a linear system of size $10^5 \times 10^5$ can only be solved by iterative methods. The system (3) is solved by a standard pre-conditioned conjugate gradient (CG) algorithm (Golub and van Loan, 1989). First, each row of M is normalized by the innovation variance (*i.e.*, we solve the problem with a correlation matrix instead of a covariance matrix). The system is pre-conditioned by solving another CG problem subject to observations confined within the boundaries of each one of the N regions. These smaller CG problems are in turn pre-conditioned by solving smaller block-diagonal systems which are designed to include full vertical observational profiles, as described in section 4. These block-diagonal systems

are directly solved using the Linear Algebra PACKage's (LAPACK, Anderson *et al.* 1992) Cholesky solver. In the serial implementation of PSAS, the normalized matrix M is provided as an operator, and the elements of M are recomputed each CG iteration. In the parallel implementation of PSAS being developed at the Jet Propulsion Laboratory (R. Ferraro, personal communication), blocks of the matrix M are pre-computed and stored in memory. Details of the serial implementation of PSAS are given in sections 3 and 5.

As a convergence criterion for the CG solver we specify that the residual must be reduced by 1 or 2 orders of magnitude. Experiments with reduction of the residual beyond 2 orders of magnitude produced differences much smaller than the expected analysis errors. This is mainly because of the filtering properties of the operator $P^J H^T$ in (4) which attenuates the small scale details in the linear system variable y .

3 Overview of the Conjugate Gradient Algorithm

This section describes the pre-conditioned conjugate gradient algorithm from a numerical point of view; the algorithm adopted is given in Table 3. The choice of pre-conditioner in PSAS is discussed in the next section, followed by a discussion of the current Fortran 90 implementation. Readers familiar with the conjugate gradient algorithm should proceed directly to section 4.

Let $M \equiv H P^J H^T + R$ be the innovation *covariance* matrix. We start by normalizing the linear system by the diagonal of M ,

$$(D^{-1} M D^{-1}) (Dy) = D^{-1} (w^o - H w^f) \quad (5)$$

or

$$\boxed{Cx = b} \quad (6)$$

where $D_{ij} = \sqrt{M_{ij}} \delta_{ij}$. In this equation C is the innovation *correlation* matrix. Following Golub and van Loan (1989, hereafter referred to as GvL) we outline the standard pre-conditioned conjugate gradient algorithm as implemented in PSAS.

We want to solve the linear system (6) where

$$b, x \in \mathbb{R}^p \quad (7)$$

$$C \in \mathbb{R}^{p \times p} \quad (8)$$

with $p \sim 10^5$ being the number of observations. Since C is *positive definite*, solving $Cx = b$ is equivalent to finding x which minimizes the functional

$$J(x) = \frac{1}{2} x^T C x - x^T b \quad (9)$$

The general strategy is to devise an iteration which converges to the minimum of $J(x)$ as fast as possible.

3.1 General Search Directions

Consider the iteration k ,

$$x_k = x_{k-1} + \alpha_k p_k \quad (10)$$

where the step size $\alpha \in \mathbb{R}$ is a scalar and $p_k \in \mathbb{R}^p$ is a vector defining a *search direction* to be determined. It is easy to show that to minimize $J(x_{k-1} + \alpha p_k)$ with respect to α , we merely set

$$\alpha = \alpha_k = \frac{p_k^T r_{k-1}}{p_k^T C p_k} \quad (11)$$

where r_k is the *residual*

$$r_k = b - Cx_k \quad (12)$$

For this choice of α we can show that

$$J(x_{k-1} + \alpha_k p_k) = J(x_{k-1}) - \frac{1}{2} (p_k^T r_{k-1})^2 p_k^T C p_k \quad (13)$$

Notice that to ensure the reduction of J we must insist on p_k not be orthogonal to r_{k-1} .

3.2 The Steepest Descent Algorithm

The gradient of $J(x) = \frac{1}{2}x^T Cx - x^T b$ with respect to x is given by

$$\nabla J|_{x=x_k} = Cx_k - b \equiv -r(x_k) \quad (14)$$

The *steepest descent* algorithm looks for the minimum in the direction in which $J(x_k)$ decreases most rapidly, i.e, down-gradient

$$p_k = -\nabla J|_{x_{k-1}} = r(x_{k-1}) \quad (15)$$

GvL give an algorithm for finding the minimum of $J(x)$ by the steepest descent method which is reproduced in Table 1.

Table 1: Steepest descent search direction algorithm (Golub and van Loan, 1989)

```

k = 0; x_0 = 0; r_0 = b
while r_k ≠ 0
  k = k + 1
  q_{k-1} = Cr_{k-1}
  α_k = r_{k-1}^T r_{k-1} / r_{k-1}^T q_{k-1}
  x_k = x_{k-1} + α_k r_{k-1}
  r_k = r_{k-1} - q_{k-1} α_k
end

```

A known drawback of this algorithm is that convergence is too slow for matrices with large condition numbers ($\kappa_2(C) = \lambda_{max}/\lambda_{min}$, where λ is the eigenvalue of C); in this case the contours of J are elongated hyperellipsoids, and we are forced to travel back and forth *across* a valley rather than *down* a valley (there is a good discussion in Press et al. 1992). The conjugate gradient algorithm addresses this deficiency of the steepest descent method.

Table 2: Conjugate gradient algorithm (Golub and van Loan, 1989)

```

k = 0; x0 = 0; r0 = b
while rk ≠ 0
  k = k + 1
  if k = 1 { p1 = r0 }
  else { βk = rk-1Trk-1/rk-2Trk-2
        pk = rk-1 + βkpk-1 }
  qk = Cpk
  αk = rk-1Trk-1/pkTqk
  xk = xk-1 + αkpk
  rk = rk-1 - αkqk
end

```

3.3 Conjugate Gradients

Recall that

$$J(x_{k-1} + \alpha_k p_k) = J(x_{k-1}) - (1/2) (p_k^T r_{k-1})^2 p_k^T C p_k$$

To avoid the problems we encountered with the *steepest descent* algorithm, we would like to make sure we always travel in a direction perpendicular to the directions already traveled. Mathematically, we would like

$$p_j^T C p_k = 0, \quad j < k \quad (16)$$

and, of course, we must have $p_k^T r_{k-1} \neq 0$ to ensure that J decreases in each iteration (see eq. 13). The following choice has this property

$$p_k = r_{k-1} - \frac{p_{k-1}^T C r_{k-1}}{p_{k-1}^T C p_{k-1}} p_{k-1} \quad (17)$$

It can be shown that

$$J(x_k) = \min\{J(x) | x \in \text{span}\{p_1, \dots, p_k\}\} \quad (18)$$

which guarantees global convergence and finite termination. Using a few identities (see GvL) we arrive at the algorithm given in Table 2.

Pre-conditioned Conjugate Gradients

The conjugate gradient converges as follows

$$\|x - x_k\|_C \leq 2 \|x - x_0\|_C \left(\frac{\sqrt{\kappa} - 1}{\sqrt{\kappa} + 1} \right)^k \quad (19)$$

where $\|x\|_C^2 = x^T C x$, and $\kappa = \lambda_{max}/\lambda_{min}$ is the condition number. So, convergence can be slow for large condition numbers¹. In order to improve convergence we seek a transformation

¹In practice, the early convergence rate depends on an *effective* condition number which is related to the *smoothness* of the RIIS.

Table 3: The pre-conditioned conjugate algorithm as implemented in PSAS (Golub and van Loan, 1989)

```

k = 0; x0 = 0; r0 = b
while rk ≠ 0
  solve  $\hat{C}z_k = r_k$  !  $\hat{C} = A^2$ : preconditioner
  matrix
  k = k + 1
  if k = 1 { p1 = z0 }
  else {  $\beta_k = r_{k-1}^T z_{k-1} / r_{k-2}^T z_{k-2}$ 
        pk = zk-1 +  $\beta_k p_{k-1}$  }
  qk = Cpk
   $\alpha_k = z_{k-1}^T r_{k-1} / p_k^T q_k$ 
  xk = xk-1 +  $\alpha_k p_k$ 
  rk = rk-1 -  $\alpha_k q_k$ 
end

```

of the original matrix C of the form,

$$\bar{C} \equiv A^{-1}CA^{-1} \quad (20)$$

where the matrix A is to be determined. Rather than solving $Cx = b$ we solve

$$(A^{-1}CA^{-1})Ax = A^{-1}b \quad \text{or} \quad \bar{C}\bar{x} = \bar{b} \quad (21)$$

If $A^2 \sim C$ then $\bar{C} \sim I$, and the conjugate gradient converges very fast because $\kappa(\bar{C}) \sim 1$. However, $\hat{C} \equiv A^2$ must be *simple* enough for the algorithm to be cost-effective. Usually the pre-conditioner is obtained by solving a simplified version of the problem. The pre-conditioned conjugate gradient algorithm implemented in PSAS is given in Table 3. The pre-conditioner amounts to solve an extra linear system $A^2z_k = r_k$ every iteration. Notice that the major cost of each iteration is the matrix vector multiply operation Cp_k . Therefore, the flop counts for this algorithm scales as $\sim p^2$, *i.e.*, it scales as the square of the number of observations.

The choice pre-conditioners implemented in PSAS is discussed in the next section.

4 Choice of pre-conditioner in PSAS

The first step consists of dividing the globe into N non-overlapping geographic regions, and sorting the observations by region and data-type. For the Cray C-90 implementation we divide the globe in 80 equal-area regions using a *icosahedral* grid (Pfaendtner 1996). In the Massive Parallel implementation of PSAS being developed at JPL the globe is divided in 256 or 512 geographically irregular regions, each having approximately the same number of observations. This strategy is necessary to achieve load balance. The domain decomposition in PSAS is user specified and the different options will be documented elsewhere.

A good pre-conditioner must have two important characteristics: 1) it must be cheap to compute, and 2) it must retain the essentials of the original problem if it is to effectively improve

the convergence rate of the algorithm. In fact, when we normalized the original problem by the innovation standard deviations, we indeed performed an implicit pre-conditioning. In this case the pre-conditioner approximates the original matrix by its diagonal.

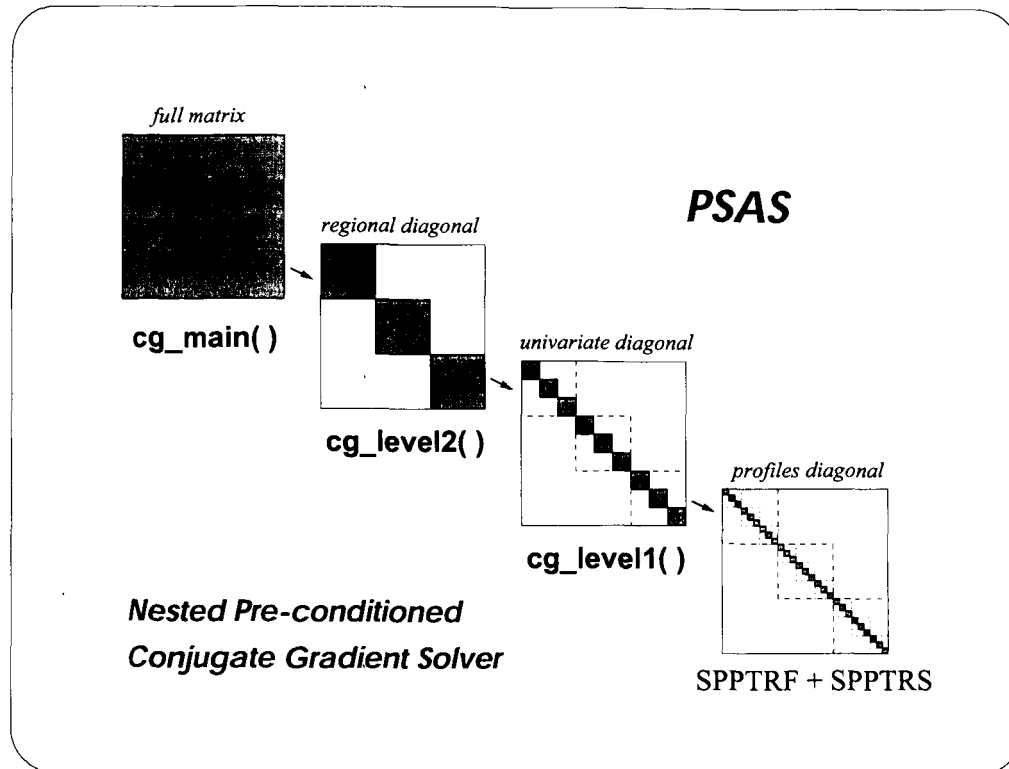


Figure 1: PSAS nested pre-conditioned conjugate gradient solver. Routine `cg_main()` contains the main conjugate gradient driver. This routine is pre-conditioned by `cg_level2()`, which solves a similar problem for each region. This routine is in turn pre-conditioned by `cg_level1()` which solves the linear system univariately. See text for details.

For the statistical interpolation problem that PSAS implements, a natural candidate for pre-conditioner is an OI-like approximation, in which the problem is solved separately for each of the N regions we used to partition the data. With $p \sim 100,000$ observations and $N \sim 80$ regions, each of these regional problems would have on average more than 1,000 observations, still too many observations for an efficient pre-conditioner. These regional problems are also solved by a pre-conditioned conjugate gradient (CG) algorithm; internally we refer to this solver as the *CG level 2*. As a pre-conditioner for *CG level 2* we solve the same problem univariately for each data type, *i. e.*, observations of u -wind, v -wind, geopotential height, etc., are treated in isolation. However, these univariate problems are still too large to be efficiently solved by direct methods and another iterative solver is used; this is the *CG level 1* algorithm. As a pre-conditioner for *CG level 1* we use LAPACK (Anderson *et al.* 1992) to perform a direct Cholesky factorization of diagonal blocks of the *level 1* correlation sub-matrix. These diagonal blocks are typically of size 32, and are carefully chosen to include full vertical profiles, a desirable feature for the implementation of new data types. These nested pre-conditioned conjugate gradient solvers are illustrated in Figure 1.

5 Fortran 90 implementation of the PSAS Conjugate Gradient Solver

In this section we discuss the main Fortran 90 drivers implementing PSAS's nested conjugate gradient solver. Intentionally, we will not discuss the details of the covariance matrix-vector multiply, i.e., the step $q_k = Cp_k$ in the algorithm shown in Table 3; this complex aspect of the PSAS algorithm will be documented in a separate Office Note. A block diagram of the

PSAS Fortran 90 Driver

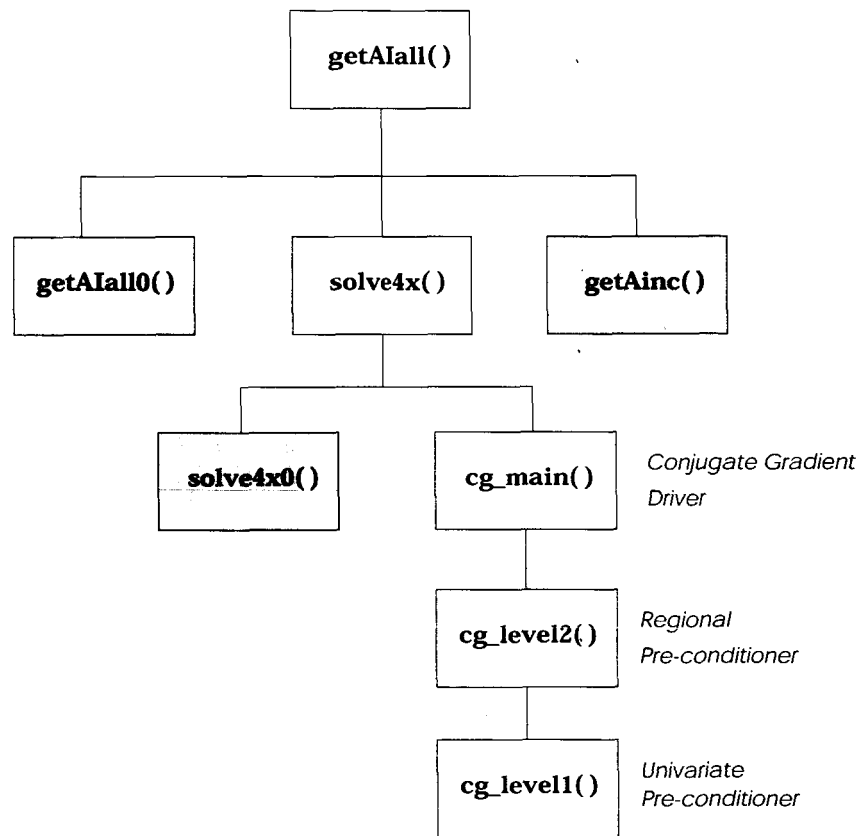


Figure 2: Block diagram of the higher level PSAS modules. The shaded blocks are not discussed in detail in this document.

modules discussed in this document is given in Figure 2; source code listing and prologue appear in the Appendix. The shaded blocks in Figure 2 are shown only for completeness; their description requires details of the covariance modeling sub-system which we do not discuss in this document.

As our starting point, we will assume that quality-controlled innovations are available, and we will discuss how the partially weighted innovations y (eq. 3) are computed. The actual calculation of the analysis increments requires the matrix-vector multiply $P^J H^T y$ (eq. 4) which cannot be discussed without going into the details of the error covariance modeling. For this reason, the module `getAinc()` shown in Figure 2 will be discussed in a separate

Office Note.

5.1 The main PSAS driver: `getAIall()`

This current version of this routine starts by performing a number of pre-processing tasks which eventually should be moved to the data ingestion section of GEOS/DAS; we have isolated this code segment inside the internal routine `psas0()`. Among these tasks are the partition of observations into regions and sorting (routine `sort()`). The following keys are used in the sorting of data

- region index
- data type index (`kt`)
- data source index (`kx`)
- latitude
- longitude
- level

After sorting, the first segment of the observation vectors will have data for region 1, then region 2, up to region N (although some regions may be empty). Inside each region all data with $kt = 1$ will be grouped together, then $kt = 2$, and so on. This sorting of the data is dictated by the strategy used for pre-conditioning described in the previous section. All routines below this point assume this sorting of the data.

Note: The current PSAS interface to GEOS/DAS is based on a customization of routine `getAIall` which processes observations and produces analysis increments in 3 separate batches, namely

surface: sea level pressure and surface winds, routine `getAIpuv()`

upper-air wind/mass: geopotential height and winds, routine `getAIzuv()`

upper-air moisture: mixing ratio, routine `getAImix()`

Because the focus of this document is on the conjugate gradient solver, we have chosen to start the PSAS driver from `getAIall()`. This interface is currently only used in the stand-alone PSAS implementation, and will eventually become the preferred GEOS/DAS interface.

5.2 Getting ready for the conjugate gradient: `solve4x()`

The internal routine `solve4x0()` performs several initializations, including

- Computes (x, y, z) cartesian coordinates on the unity sphere corresponding to the (lat, lon) of the input observations. These cartesian coordinates are used by the covariance modeling subsystem to compute horizontal distances.

- Computes the sounding index of the observations (da Silva and Redder 1995).
- Set interpolation indices and weights.
- Normalizes observation and forecast error standard deviations (by the innovation standard deviation).

This routine also performs normalization by the innovation standard deviation to transform the system to the form $Cx = b$ which is then handled by the conjugate gradient solver `cg_main()`.

5.3 The main conjugate gradient driver: `cg_main()`

This routine does a straightforward implementation of the pre-conditioned conjugate gradient algorithm given in Golub and van Loan (1989) and reproduced in Table 3; even variable names have been chosen to closely follow the book notation (with the exception perhaps, of the matrix name which we use C instead of A). The *Basic Linear Algebra Subprograms* (BLAS), which are often hand-coded in assembler and provided by several vendors, are used to perform the basic linear algebra operations such as dot products, norms, vector additions, etc. The pre-conditioner for this routine is implemented in routine `cg_level2()`. The most costly portion of this routine is the global correlation matrix-vector multiply (routine `sCxpy`) which will be documented in a separate Office Note.

5.4 Pre-conditioner level 2: `cg_level2()`

This routine has a structure very similar to `cg_main()`. The main difference is how the pre-conditioner is invoked. Recall that as a result of the data sorting, within each region the observations are sorted by data-type (*e.g.*, sea level pressure, heights, u-wind, etc. are all grouped together). The pre-conditioner for this routine is implemented in routine `cg_level1()` which acts on each of these (univariate) data-type vector segments independently. In order to achieve multi-tasking on the Cray C90, this routine includes compiler directives to perform pre-conditioner operations for each data-type segments in parallel.

5.5 Pre-conditioner level 1: `cg_level1()`

The general structure of this routine is again similar to `cg_main()`. However, at this level the correlation block sub-matrices are explicitly computed and stored (see internal routine `cg_blocks()`). The pre-conditioner is now implemented in `cg_level0()`. This internal routine identifies blocks of the correlation sub-matrix which contain full vertical profiles. The number of profiles is user specified; typical values are 2 or 3. A direct Cholesky solver is performed on these blocks using LAPACK (Anderson *et al.* 1992). This Cholesky solver is typically performed on matrix of size 32×32 .

6 Concluding remarks

As of this writing the PSAS system is undergoing major revisions in its fundamental modules. In particular, the error covariance modeling sub-system is being updated to allow more general models (for example, non-homogeneous, non-separable correlation models), and an infra-structure for dealing with complex data-types (*e.g.*, radiances, total precipitable water) is being developed. In this document we have concentrated on the conjugate gradient solver component of PSAS. Although some revisions in these modules will be necessary as we expand some of the data structures, they will almost certainly only involve interface changes. The general structure of the algorithm appears robust and is not expected to change.

Acknowledgments

The original proposal for a global, physical-space statistical analysis system to replace DAO's OI was made by S. Cohn (1991, manuscript notes). A Fortran 77 version of PSAS was designed and implemented by the late Jim Pfaendtner during 1992-93 on his workstation. Jim Searl implemented a preliminary (univariate) version of the error covariance routines. Meta Sienkiewicz wrote the original wind-mass covariance routines and implemented the moisture analysis. David Lamich wrote the main interface to PSAS on the GEOS/DAS end (internally referred to as the "plug-version"). We would like to acknowledge their contribution and consistent encouragement during the course of this project. Thanks also to Ricky Rood (head of DAO) for overall support, and to Jim Stobie for his continued encouragement of our documentation efforts.

A formal technical review of this document was conducted on February 26, 1996 at the Data Assimilation Office. We would like to thank Meta Sienkiewicz (review leader), Genia Brin (recorder), David Lamich and Peter Lyster (reviewers) for valuable suggestions. Thanks also to Ricardo Todling for proofreading the manuscript.

A Appendix: PSAS Conjugate Gradient Solver prologues and source code

A.1 getAIall()

Given innovation (observation minus forecast) data, this routine returns the analysis increments (analysis minus first guess) using the Global conjugate gradient algorithm implemented in PSAS. Basically, the calculation is performed in 2 stages. First, a global, pre-conditioned conjugate gradient solver is used to solve for y in the equation

$$(HP^f H^T + R)y = w^o - Hw^f$$

where $w^o - Hw^f$ is the innovation. Notice that y is defined in observation locations. Subsequently, the gridded analysis increments δw_a are computed from y by the matrix-vector multiply

$$\delta w_a = P^f H^T y$$

CALLING SEQUENCE:

```
      call getAIall ( nobs, lat, lon, pres,
&                  time, kx, kt, dels,
&                  sig_F, sig_0,
&                  im, jnp, mlev, pres_lev,
&                  psl_sigF, usl_sigF, vsl_sigF,
&                  z_sigF, u_sigF, v_sigF, mix_sigF,
&                  psl_inc, usl_inc, vsl_inc,
&                  z_inc, u_inc, v_inc, mix_inc,
&                  psl_sigA, usl_sigA, vsl_sigA,
&                  z_sigA, u_sigA, v_sigA, mix_sigA )
```

INPUT PARAMETERS:

```
use OEclass_tbl, only : nlev_oe, plev_oe

implicit NONE

integer      nobs           ! number of observations
real         lat(nobs)     ! latitude (deg) of each obs
real         lon(nobs)     ! longitude (deg) of each obs
real         pres(nobs)    ! pressure level (hPa) of obs
real         time(nobs)    ! time (minutes) from central
                        ! synoptic time
integer      kx(nobs)      ! GEOS/DAS data source index
```

```

integer      kt(nobs)          ! GEOS/DAS data type index
real         dels(nobs)       ! innovations (O-F)
real         sig_F(nobs)      ! forecast error stdv
real         sig_0(nobs)      ! observation error stdv (no longer
                             ! used (t. b. r.))

```

```

! -----
! NOTE: nobs, kx, kt, dels, sig_F & sig_0 are updated during
!       the super-obing (routine proxel() ).
! -----

```

```

integer      im                ! no. of zonal grid-points
integer      jnp               ! no. of meridional gridpoints
integer      mlev              ! no. of vertical grid-points
real         pres_lev(mlev)    ! list of vertical levels (hPa)

! The arrays below with suffix
! _sigF are gridded forecast error
! standard deviations for:
! o sea level pressure (hPa)
real         psl_sigF(im,jnp)
! o surface u-wind (m/s)
real         usl_sigF(im,jnp)
! o surface v-wind (m/s)
real         vsl_sigF(im,jnp)
! o upper-air u-wind (m/s)
real         u_sigF(im,jnp,mlev)
! o upper-air v-wind (m/s)
real         v_sigF(im,jnp,mlev)
! o geopotential height (m/s)
real         z_sigF(im,jnp,mlev)
! o mixing ratio (g/kg)
real         mix_sigF(im,jnp,mlev)

```

OUTPUT PARAMETERS:

```

! The arrays below with suffix
! _inc are gridded analysis
! increments for:
! o sea level pressure (hPa)
real         psl_inc(im,jnp)
! o surface u-wind (m/s)
real         usl_inc(im,jnp)
! o surface v-wind (m/s)
real         vsl_inc(im,jnp)
! o upper-air u-wind (m/s)
real         u_inc(im,jnp,mlev)
! o upper-air v-wind (m/s)
real         v_inc(im,jnp,mlev)
! o geopotential height (m/s)
real         z_inc(im,jnp,mlev)
! o mixing ratio (g/kg)
real         mix_inc(im,jnp,mlev)

! The arrays below with suffix
! _sigA are gridded analysis error
! standard deviations for:
! o sea level pressure (hPa)
real         psl_sigA(im,jnp)
! o surface u-wind (m/s)
real         usl_sigA(im,jnp)

```



```

real    vsl_sigA(im,jnp)      ! o surface v-wind (m/s)
real    u_sigA(im,jnp,mlev)  ! o upper-air u-wind (m/s)
real    v_sigA(im,jnp,mlev)  ! o upper-air v-wind (m/s)
real    z_sigA(im,jnp,mlev)  ! o geopotential height (m/s)
real    mix_sigA(im,jnp,mlev) ! o mixing ratio (g/kg)

```

Return status:

<none> : The subroutine may exit with a non-zero status through a call to PSASexit() when an error condition is detected. In such cases execution is aborted.

BUGS:

The super-obing alters the value of observations in violation of the ODS standard. No known side effects, but this should be fixed.

SEE ALSO:

```

solve4x()  interface to conjugate gradient routines.
stdio.h    include file defining standard I/O units
BLAS       basic linear algebra sub-programs

```

SYSTEM ROUTINES:

```

getenv(3f)  UNIX interface returning the value of an
            environment variable (PSASRC here).

```

FILES USED:

```

stdrc      a unit number allocated when the subroutine is in use,

```

for the input of control parameters and data tables.

REVISION HISTORY:

ddmm95	Lamich/Guo	Interface design.
ddmm95	Guo	Initial code.
04Jan96	da Silva	Revised prologue, major clean-up. Removed IFDEFs about dynamic allocation. Code now requires Fortran 90 for portability. Introduced getAIall0() as internal routine.

SOURCE CODE:

```
character*8 myname          ! Name of routine for error messages
parameter (myname='getAIall')

! Local functionality controls
! -----
logical want_usl
logical want_vsl
logical want_psl
logical want_u
logical want_v
logical want_z
logical want_mix
parameter(want_usl=.true.)
parameter(want_vsl=.true.)
parameter(want_psl=.true.)
parameter(want_u  =.true.)
parameter(want_v  =.true.)
parameter(want_z  =.true.)
parameter(want_mix=.true.)

real sigFmiss
parameter(sigFmiss=1.e+15)

integer  nnobs
integer  n,ier
integer  nprox

! Experiment ID and date/time: debris t.b.r.
! -----
character*9  c9date
```

```

character*8  c8time

!
! Control parameters for conjugate gradient iterations
! -----
include      'bands.h'

!
! Control parameters for output.
! -----
logical      verbose
parameter ( verbose = .true.)

!
! Basically debris from left over from JimPf time
! -----
integer      idelprb
integer      idelpre
integer      idelpri
parameter ( idelprb = 250 ) ! beg to print dels
parameter ( idelpre = 20000 ) ! end to prind dels
parameter ( idelpri = 250 ) ! increment to print del
logical      prtdat1
parameter ( prtdat1 = .false.)
integer      ntwidth
parameter ( ntwidth = 30000 )

!
! Size parameters for database
! -----
include      'maxreg.h'      ! maximum number of regions
include      'kxmax.h'      ! maximum numer of data sources
include      'ktmax.h'      ! maximum number of data types
include      'ktwanted.h'   ! data structure defining data
! types for which we produce
! analysis increments.

!
! Regional (domain) decomposition maps used in PSAS
! -----
integer iregbeg(maxreg)      ! pointers to beginning of regions
integer ireglen(maxreg)      ! the no. of obs. in each region
integer ityplen(ktmax,maxreg) ! sizes of type blocks

!
! Storage for data items (dynamic allocation)
! -----
real         sig_Ou(nobs)     ! spatially uncorrelated portion of
! obs error stdv
real         sig_Oc(nobs)     ! spatially correlated portion of
! obs error stdv
real         xvec(nobs)       ! Conjugate gradient solution
! at obs location

```

```

logical      kl(nobs)          ! debris t. b. r.

include 'lvmax.h'          ! maximum no. of levels for internal tables
include 'levtabl.h'       ! vertical level tables for interpolation
                                ! of correlation functions, etc.

include      'stdio.h'     ! standard I/O units

integer      l, i
integer      n2grd, n3grd
integer      stdrc
integer      luavail, lnblnk
external     luavail, lnblnk

external     psasrcbd      ! a blockdata unit
include     'psasrc.h'    ! a default psasrc file name

```

!.....

! Initialize PSAS, sort data, assign regions, etc...

! call getAIall0() ! internal routine

! COMPUTATIONAL SECTION
! -----

! Up to this point we have done a bunch of pre-processing
! to prepare the internal data structures (forecast and
! observation correlation tables, etc). Next we actually
! do some real calculations for a change.
!

! First, solve

$$(HP^TfH^T + R) x = w^o - Hw^f$$

! for the vector x defined in observation locations.
! -----

```

call ZEITBEG('solve4x ')
call SOLVE4X ( maxreg, iregbeg, ireglen, ityplen,
&             nobs, kx, lat, lon,pres,
&             sig_Ou, sig_Oc, sig_F, 1,
&             nobs, dels, xvec )
call ZEITEND

!
call OBSTAT ( stdout, nobs, kx, kt, pres, xvec,
&            nlev_oe,plev_oe,'getAIall*SolutionVector')
```

```

!      Next, obtain the gridded analysis increments from
!
!      \delta w_a = P^f H^T x
!
-----
call ZEITBEG ('getAinc')
call getAinc ( verbose, stdout, nbandcg,
&             nobs, iregbeg, ireglen, ityplen, xvec,
&             lat, lon, pres, sig_F,
&             im, jnp, mlev, pres_lev,
&             usl_inc, vsl_inc, psl_inc,
&             u_inc, v_inc, z_inc, mix_inc,
&             ktwanted(ktus),
&             ktwanted(ktvs),
&             ktwanted(ktslp),
&             ktwanted(ktuu),
&             ktwanted(ktvv),
&             ktwanted(ktHH),
&             ktwanted(ktqq),
&             ier)
call ZEITEND      ! getAinc

!      Error handling
!      -----
if(ier.ne.0) then
  write(stderr, '(2a,i4)') myname,
&      ': error from getAinc(), ', ier
  call PSASexit ( 2, myname )
end if

!      Scale the normalized analysis increments returned by getAinc()
!      -----
if(ktwanted(ktus )) call QVMV (usl_inc,usl_inc,usl_sigF,n2grd)
if(ktwanted(ktvs )) call QVMV (vsl_inc,vsl_inc,vsl_sigF,n2grd)
if(ktwanted(ktslp)) call QVMV (psl_inc,psl_inc,psl_sigF,n2grd)
if(ktwanted(ktuu )) call QVMV (u_inc,u_inc,u_sigF,n3grd)
if(ktwanted(ktvv )) call QVMV (v_inc,v_inc,v_sigF,n3grd)
if(ktwanted(ktHH )) call QVMV (z_inc,z_inc,z_sigF,n3grd)
if(ktwanted(ktqq )) call QVMV (mix_inc,mix_inc,mix_sigF,n3grd)

!      Print summary (means/std/min/max) of several grids
!      -----
if(ktwanted(ktus).or.ktwanted(ktvs).or.ktwanted(ktslp)) then
  write(stdout, '(/2a)') myname,
&      ': Analysis-Increments of Surface Variables:'

```

```

    if(ktwanted(ktus)) call LVSTAT (stdout,im,jnp,usl_inc,
&      0.,'WIND','SRFC',1.e+15,'USL')
    if(ktwanted(ktvs)) call LVSTAT (stdout,im,jnp,vsl_inc,
&      0.,'WIND','SRFC',1.e+15,'VSL')
    if(ktwanted(ktslp)) call LVSTAT (stdout,im,jnp,psl_inc,
&      0.,'PRES','SRFC',1.e+15,'SLP')

```

```
end if
```

```

if(ktwanted(ktuu).or.ktwanted(ktvv).or.
&   ktwanted(ktHH).or.ktwanted(ktqq)) then

```

```

    write(stdout,'(/2a)') myname,
&      ': Analysis-Increments of Upper-Air Variables:'

```

```

    if(ktwanted(ktuu)) call GDSTAT (stdout,im,jnp,mlev,
&      u_inc,pres_lev,'WIND','PRES',1.e+15,'A-Inc of UWND',1)
    if(ktwanted(ktvv)) call GDSTAT (stdout,im,jnp,mlev,
&      v_inc,pres_lev,'WIND','PRES',1.e+15,'A-Inc of VWND',1)
    if(ktwanted(ktHH)) call GDSTAT (stdout,im,jnp,mlev,
&      z_inc,pres_lev,'HGHT','PRES',1.e+15,'A-Inc of HGHT',1)
    if(ktwanted(ktqq)) call GDSTAT (stdout,im,jnp,mlev,
&      mix_inc,pres_lev,'MIXR','PRES',1.e+15,'A-Inc of MIXR',1)

```

```
end if
```

```

! Assign sigA values here. They are initialized to zeroes for
! now. The operation must be conditional since the memory may
! not be available for some calls.
!

```

```

-----
call ZEITBEG ( 'getsigA' )
if(ktwanted(ktus)) call SSCAL (n2grd,0.,usl_sigA,1)
if(ktwanted(ktvs)) call SSCAL (n2grd,0.,vsl_sigA,1)
if(ktwanted(ktslp)) call SSCAL (n2grd,0.,psl_sigA,1)
if(ktwanted(ktuu)) call SSCAL (n3grd,0.,u_sigA,1)
if(ktwanted(ktvv)) call SSCAL (n3grd,0.,v_sigA,1)
if(ktwanted(ktHH)) call SSCAL (n3grd,0.,z_sigA,1)
if(ktwanted(ktqq)) call SSCAL (n3grd,0.,mix_sigA,1)
call ZEITEND

```

```

! All done
!

```

```

l=len(psasname)+len('*')+len(myname)+len('(): normal return')
write(stdout,'(/80a)') ('=',i=1,l)
write(stdout,'(5a)') psasname,'*',myname,'(): normal return'
write(stdout,'(80a)') ('=',i=1,l)

```

```
return
```

```
CONTAINS
```


A.2 getAIall0()

This INTERNAL routine initializes several aspects of PSAS, including:

- Opens resource file and initializes several tables necessary for the error covariance modeling subsystem.
- Assigns a region number to each observation and set the relevant internal pointers.
- Sorts observations by region, data-type, data-source, latitude, longitude and level.
- Performs super-obing.
- Prints out lots of informational output, if specified.

CALLING SEQUENCE:

```
call getAIall0()
```

INPUT PARAMETERS:

Explicitly none, but this routine inherits all data from its parent getAIall().

OUTPUT PARAMETERS:

Explicitly none, but this routine resets most of the input parameters to getAIall().

BUGS:

Most of the complexity level of this routine is due to its provisional nature. Eventually most of these tasks will be moved to the data ingestion level of the data assimilation system.

SEE ALSO:

getAIall() parent routine.

FILES USED:

stdrc a unit number allocated when the subroutine is in use,
for the input of control parameters and data tables.

REVISION HISTORY:

12feb96 da Silva Moved from main body of getAIall().

SOURCE CODE:

```
! Hello, world!
! -----
! l=len(psasname)+len('*')+len(myname)+
& len('(): Version_')+lnblnk(version)
!
! write(stdout,'(/80a)') ('=',i=1,1)
! write(stdout,'(5a)') psasname,'*',myname,'(): Version ',version
! write(stdout,'(80a)') ('=',i=1,1)
!
! Total number of 2-D and 3-D gid-points
! -----
! n2grd = im * jnp
! n3grd = im * jnp * mlev
!
! Open resource file and initialize INPAK77
! -----
! stdrc=luavail()
! call GETENV ( 'PSASRC', psasrc ) ! Unix extension
```

```

if(psasrc.eq.' ') psasrc=def_psasrc      ! default name
call OPNINPK (stdrc,psasrc,ier)
l=max(1,lnblnk(psasrc))
if(ier.ne.0) then
  write(stderr,'(4a,i4)') myname,': error from opninpk(',
&      psasrc(1:1),')', iostat = ',ier
  call PSASexit(2,myname)
else
  write(stdout,'(4a)') myname,': using ',psasrc(1:1),
&      ' for runtime parameter input'
end if

!      Initialize observation related information
!      -----
call initRSRC

!      List initialized information.  Need rewrite pardisp(), since
!      so many changes have been made.  A lot of information listed by
!      pardisp() is no longer relevant, while some thing important is
!      not even listed.
!      -----
c9date='01-apr-99'  ! talking about debris...
c8time='000000'
call PARDISP ( STDOUT,
&      myname,   c9date, c8time,
&      nobs,    kxmax,  ktmax,
&      verbose, stdout, idelprb, idelpre, idelpri,
&      '*****', -99, 0, 0, ntwidht,
&      nbands,  msmall,
&      cgname,  seplim, criter, minmax, maxpass  )

!      Print a summary of all observations.
!      -----
if(verbose) call OBSSMRY ( stdout, nobs, kx, kt )

!      Reset ktwanted according to the mask for this call.
!      -----
ktwanted(ktus )=ktwanted(ktus ).and.want_usl
ktwanted(ktvs )=ktwanted(ktvs ).and.want_vsl
ktwanted(ktslp)=ktwanted(ktslp).and.want_psl
ktwanted(ktuu )=ktwanted(ktuu ).and.want_u
ktwanted(ktvv )=ktwanted(ktvv ).and.want_v
ktwanted(ktHH )=ktwanted(ktHH ).and.want_z
ktwanted(ktqq )=ktwanted(ktqq ).and.want_mix

!      Print out informational summaries
!      -----
if(ktwanted(ktus).or.ktwanted(ktvs).or.ktwanted(ktslp)) then
  write(stdout,'(/2a)') myname,

```

```

&           ': Sigma-F of Surface Variables:'
  if(ktwanted(ktus)) call lvstat(stdout,im,jnp,usl_sigF,
&           0.,'WIND','SRFC',sigFmiss,'USLE')
  if(ktwanted(ktvs)) call lvstat(stdout,im,jnp,vsl_sigF,
&           0.,'WIND','SRFC',sigFmiss,'VSLE')
  if(ktwanted(ktslp)) call lvstat(stdout,im,jnp,psl_sigF,
&           0.,'PRES','SRFC',sigFmiss,'SLPE')
end if

if(ktwanted(ktuu).or.ktwanted(ktvv).or.
&   ktwanted(ktHH).or.ktwanted(ktqq)) then

  write(stdout,'(/2a)') myname,
&           ': Sigma-F of Upper-Air Variables:'

  if(ktwanted(ktuu)) call GDSTAT(stdout,im,jnp,mlev,
&   u_sigF,pres_lev,'WIND','PRES',sigFmiss,'Sigma-F of UWND',1)
  if(ktwanted(ktvv)) call GDSTAT(stdout,im,jnp,mlev,
&   v_sigF,pres_lev,'WIND','PRES',sigFmiss,'Sigma-F of VWND',1)
  if(ktwanted(ktHH)) call GDSTAT(stdout,im,jnp,mlev,
&   z_sigF,pres_lev,'WIND','PRES',sigFmiss,'Sigma-F of HGHT',1)
  if(ktwanted(ktqq)) call GDSTAT(stdout,im,jnp,mlev,
&   mix_sigF,pres_lev,'WIND','PRES',sigFmiss,
&   'Sigma-F of MIXR',1)

end if

! Restrict observations only to those 'within' at least one of
! 'hyper-boxes', defined by lat/lon/pres/kx/kt/time. Remove data
! outside the 'hyper-boxes' by pushing them to the end of the list
! and reset 'nobs' to the size of the front part of the list.
! -----
call ZEITBEG ('restrict')
call RESTRICT ( verbose, stdout, nobs, prtdat1,
&             lat, lon, pres,kx, kt,
&             dels, sig_0, sig_F,
&             time,nobs )
nobs = nobs ! completely redefine the whole data record.
call ZEITEND

! Sort observations in the order of:
!
!           region(lat,lon)-kt-kx-lat-lon-pres
!
! Also, define pointer/size information of each region and type
! by set arrays iregbeg, ireglen, and ityplen.
! -----
call ZEITBEG ('sort')

call SORT ( myname, verbose, stdout, nobs,
&         lat, lon, pres,
&         kx, kt, dels,

```

```

&          sig_0, sig_F, time,
&          maxreg, ktmax, iregbeg, ireglen, ityplen )

call ZEITEND

!          Remove 'duplicates' in the observations and adjust iregbeg,
!          ireglen, and ityplen accordingly.
!          -----
call ZEITBEG ('dupelim')
call DUPELIM (verbose, stdout,
&          nobs, kx, kt, kl,
&          lat, lon, pres,
&          dels, sig_0, sig_F, time,
&          maxreg, iregbeg, ireglen, ktmax, ityplen )
call ZEITEND

!          'Superob' observations that are within a given range. Quit
!          searching loop if nothing to 'superob', or have looped 5 times.
!          Iregbeg, ireglen, and ityplen arrays are adjusted accordingly.
!          -----
call ZEITBEG ('proxel')
nprox=0
n=1
do while(n.eq.1 .or. nprox.ne.0.and.n.le.5)
  call PROXEL ( verbose, stdout,
&          nobs, kx, kt, kl,
&          lat, lon, pres,
&          dels, sig_0, sig_F,
&          time, maxreg, iregbeg, ireglen,
&          ktmax, ityplen, nprox )
  n=n+1
end do
call ZEITEND

!          Reset the levels of the surface variables to 1000.
!          This way the surface analysis will use the same error
!          characteristics at the surface and at 1000 hPa.
!          -----
do n=1,nobs
  if( kt(n).eq.ktslp .or.
&          kt(n).eq.ktus .or.
&          kt(n).eq.ktvs) then
    pres(n)=1000.
  end if
end do

!          Set the grid parameters. There apparently a good here for it
!          to be defined only now.
!          -----
call GRIDXXO

```

```

! Merge in observation levels
! -----
call ZEITBEG('setcors')
call SETPLEVS ( mlev,pres_lev,nobs,pres,
&             MXveclev,nveclev,pveclev)
call SET_oeCHH
call SET_fecHH
call SET_fecQQ
call SETfecW      ! naming inconsistency
call ZEITEND

! Create observation error stdv. NOTE: in the original
! PSAS design the observation error standard deviation
! came along with the data stream. Due to the increasing
! complexity of the observation error modeling, the
! observation error is now derived from parameters in the
! resource file. Next we overwrite whatever came in...
! -----
call INTP_sig0 ( nobs, kx, kt, pres, sig_0c, sig_0u )

! More informational output. This time prints a summary of the
! observations actually used in analysis
! -----
if(verbose) call OBSSMRY ( stdout, nobs, kx, kt )

call OBSTAT ( stdout, nobs,
&             kx,kt,pres, sig_F,
&             nlev_oe, plev_oe,
&             'getAIall*FcstErr*sigF' )

call OBSTAT ( stdout, nobs,
&             kx,kt,pres, dels,
&             nlev_oe,plev_oe,'getAIall*InnovVector')

call OBSTAT ( stdout, nobs,
&             kx, kt, pres, sig_0c,
&             nlev_oe, plev_oe, 'getAIall*ObsErr*sig0c')

call OBSTAT ( stdout, nobs,
&             kx, kt, pres, sig_0u,
&             nlev_oe, plev_oe,'getAIall*ObsErr*sig0u')

return
end subroutine getAIall0

```

A.3 solve4x()

Given innovation (observation minus forecast) data, this routine returns the vector y solution of the linear system of equations

$$(HP^f H^T + R)y = w^o - Hw^f$$

where $w^o - Hw^f$ is the innovation. (The notation follows da Silva and Guo 1996, DAO Office Note 9602). Notice that y is defined at observation locations. A pre-conditioned conjugate gradient algorithm is used to solve this linear system. This routine can handle multiple RHS vectors, a feature needed for the calculation of analysis error variances by means of randomized trace estimates.

CALLING SEQUENCE:

```
      call SOLVE4X ( nkr, kr_beg, kr_len, kt_len,
&                  nobs, kx, rlat, rlon, rlev,
&                  sigU, sig_Oc, sig_F, nvecs,
&                  nobs_d, rhs, Xvec )
```

INPUT PARAMETERS:

```
implicit NONE

include 'ktmax.h'           ! maximum no. of data types
integer nkr                 ! number of regions
integer kr_beg(nkr)        ! beginning of each region
integer kr_len(nkr)        ! no. of obs. in each region
integer kt_len(ktmax,nkr)  ! no. of obs. of a given data
                           ! in each region

integer nobs                ! number of observations
integer kx(nobs)            ! GEOS/DAS data sources
real rlat(nobs)             ! latitudes (deg) of obs.
real rlon(nobs)            ! longitudes (deg) of obs.
real rlev(nobs)            ! pressure levels (hPa) of obs.
real sig_Ou(nobs)          ! spatially uncorrelated portion
                           ! of obs. error stdv
real sig_Oc(nobs)          ! spatially correlated portion
                           ! of obs. error stdv
real sig_F(nobs)           ! forecast error stdv

integer nvecs              ! number of RHS vectors
integer nobs_d             ! leading dimension of RHS vector
                           ! as declared in calling program.
```

```

! Usually nobs_d = nobs.

real      rhs(nobs_d,nvecs) ! RHS vectors. For the convetional
! PSAS analysis system 'rhs' will
! contain the innovations (O-F).
! However, multiple RHS will be
! necessary for implementation
! analysis error variances by
! randomized trace estimates.

```

```

----
NOTE: All input arrays indexed by 'nobs' or 'nobs_d' are assumed
----
sorted by region. Within each region, data is assumed
sorted by data type (kt). Within each data-type, data
is assumed sorted by latitude, longitude and finally by
levels.

```

OUTPUT PARAMETERS:

```

real      Xvec(nobs_d,nvecs) ! solution vectors.

```

SEE ALSO:

```

cg_main()      top level conjugate gradient routine.

```

REVISION HISTORY:

```

ddmmm93 Pfaendtner Original code.
28may93 Searl      Modification for dynamic storage on CRAY.
07jan94 Sienkiewicz Added pass of trig lat/lon.
03oct94 da Silva   Implemented CRAY specifics with IFDEFs.
                Eliminated calls to conjgr3 . conjgr4.
                Input parameter 'nbandmx' is now obsolete.
04oct94 da Silva   Introduced parameter nband, and call to CONJGR.
19Jan95 Guo        Added wobs tables to pass pindx2() values to
                ??cor1() and ??corx() routines. One could use

```

rlevs for the same purpose to reduce the overhead, since rlevs has no real purpose in this subroutine and subsequent routines.
 02Feb95 Guo Changed CRAY to _UNICOS for consistency and to follow the guide lines.
 05Feb96 da Silva Revised prologue and major clean-up.
 Removed IFDEFs about dynamic allocation. Code now requires Fortran 90 for portability.
 Introduced internal routine solve4x0().

SOURCE CODE:

```

character*7 myname
parameter(myname='solve4x')

!
! Conjugate gradient data structure
! -----
include      'bands.h'

!
! Dynamic allocation
! -----
real      sig_del(nobs)      ! innovation (O-F) stdv
real      nsig_Ou(nobs)     ! normalized sig_Ou = sig_Ou/sig_del
real      nsig_Oc(nobs)     ! normalized sig_Oc = sig_Oc/sig_del
real      nsig_F(nobs)      ! normalized sig_F = sig_F /sig_del

! Cartesian coordinates (on the
! unity sphere) of unit vectors
! of the spherical coordinate system
real      qr_x(nobs)        ! o x-coord of radial      unit vector
real      qr_y(nobs)        ! o y-coord of radial      unit vector
real      qr_z(nobs)        ! o z-coord of radial      unit vector
real      qm_x(nobs)        ! o x-coord of meridional  unit vector
real      qm_y(nobs)        ! o y-coord of meridional  unit vector
real      qm_z(nobs)        ! o z-coord of meridional  unit vector
real      ql_x(nobs)        ! o x-coord of longitudinal unit vector
real      ql_y(nobs)        ! o y-coord of longitudinal unit vector
! NOTE: ql_z is not needed.

! Interpolation indices/weights:
integer   ktab(nobs)        ! o vertical  interpolation index
real      wtab(nobs)        ! o vertical  interpolation weights
integer   jtab(nobs)        ! o meridional interpolation index
real      vtab(nobs)        ! o meridional interpolation weights

integer   ks(nobs)          ! sounding index
  
```



```

real      bvec(nobs,nvecs) ! normalized RHS = RHS / sig_del

! Levels for correlation tables, etc.
! -----
include 'lvmax.h'
include 'levtabl.h'
include 'hfecW.h'

include 'stdio.h'          ! standard i/o

! Local variables
! -----
integer  i                  ! data index
integer  ivec               ! vector index
integer  ierr               ! error code

real     var

! .....

! Compute cartesian coordinates and set interpolation indices
! -----
call solve4x0()

! Normalize the the RHS vectors
! -----
do ivec=1,nvecs
do i=1,nobs
      bvec(i,ivec) = rhs(i,ivec) / sig_del(i)
end do
end do

! Use conjugate gradient algorithm to solve the normalized
! linear system based on the innovation CORRELATION matrix,
! i.e., the CG solver works on the system
!
!
!          C x = b
!
! where C is the innovation correlation matrix and b is
! (usually) the innovation normalized by its standard deviation
! -----
call CG_MAIN ( cgverb(nbandcg),
&             nkr, kr_beg, kr_len, kt_len,
&             nobs, ks, nsig_Ou, nsig_Oc, nsig_F,
&             qr_x, qr_y, qr_z,
&             qm_x, qm_y, qm_z,
&             ql_x, ql_y, ktab,
&             wtab, jtab, vtab,
&             nvecs, nobs, bvec, nobs, Xvec, ierr )

```

```

!   Error handling
!   -----
      if ( ierr .ne. 0 ) then
          write(stderr,'(2a,i3)') myname,
&          ' : error from cg_main(), ',ierr
          call PSASexit ( 2, myname )
      end if

!   Scale solution by the innovation standard deviation
!   -----
      do ivec = 1, nvecs
          do i = 1, nobs
              Xvec(i,ivec) = sig_del(i) * Xvec(i,ivec)
          end do
      end do

!   All done
!   -----

      return

!   CONTAINS
!   -----

```

A.4 solve4x0()

This INTERNAL Fortran 90 routine initializes several internal parameters relevant to the conjugate gradient solver, including

- Computes (x, y, z) cartesian coordinates on the unity sphere corresponding to the (lat,lon) of the input observations. These cartesian coordinates are used by the covariance modeling subsystem to compute horizontal distances.
- Computes the sounding index of the observations.
- Set interpolation indices and weights.
- Normalizes observation and forecast error standard deviations (by the innovation standard deviation).

CALLING SEQUENCE:

```
call solve4x0()
```

INPUT PARAMETERS:

Explicitly none, but this routine inherits all data from its parent solve4x().

OUTPUT PARAMETERS:

Explicitly none, but this routine sets several quantities of relevance to the conjugate gradient solver.

SEE ALSO:

solve4x() parent routine.

REVISION HISTORY:

12feb96 da Silva Moved from main body of solve4x().

SOURCE CODE:

```
!      Compute x,y,z coordinates of observations
!      -----
      call LL2QVEC ( nobs,rlat,r lon,
&                  qr_x,qr_y,qr_z,qm_x,qm_y,qm_z,ql_x,ql_y)

!      Set sounding index of observations
!      -----
      call SETPIX ( nobs, kx, rlat, rlon, ks )

!      Set tables for vertical/horizontal interpolation
!      -----
      call SLOGTAB (.true., nveclev,pveclev,nobs,rlev,ktab,wtab)
      call SLINTAB (.true., nHlat,Hlat,nobs,rlat,jtab,vtab)

!      Compute normalized error stdv
!      -----
      do i=1,nobs
        var=sig_Ou(i)*sig_Ou(i)+sig_Oc(i)*sig_Oc(i)+sig_F(i)*sig_F(i)
        sig_del(i) = 1. / sqrt(var)
        nsig_Ou(i) = sig_Ou(i) / sig_del(i)
        nsig_Oc(i) = sig_Oc(i) / sig_del(i)
        nsig_F(i)  = sig_F(i)  / sig_del(i)
      end do

      return

      end subroutine SOLVE4X0
```

A.5 cg_main()

Solves the linear system of equations

$$Cx = b$$

where C is the innovation correlation matrix, and b is a set of multiple RHS. When performing a global analysis with PSAS, the RHS is simply the innovation (O-F) normalized by its standard deviation. The multiple RHS are necessary to estimate analysis error variances by means of randomized trace estimates.

The *Pre-conditioned Conjugate Gradient* algorithm is standard and closely follows

Golub, G. H. and C. F. van Loan, 1989: *Matrix Computations*, 2nd Edition, The John Hopkins University Press, 642pp.

and is reproduced below.

```
k = 0; x0 = 0; r0 = b
while r_k ≠ 0
  solve  $\hat{C}z_k = r_k \Rightarrow$  call cg_level2()
  k = k + 1
  if k = 1 { p1 = z0 }
  else {  $\beta_k = r_{k-1}^T z_{k-1} / r_{k-2}^T z_{k-2}$ 
         $p_k = z_{k-1} + \beta_k p_{k-1}$  }
   $q_k = Cp_k \Rightarrow$  call sCxy()
   $\alpha_k = z_{k-1}^T r_{k-1} / p_k^T q_k$ 
   $x_k = x_{k-1} + \alpha_k p_k$ 
   $r_k = r_{k-1} - \alpha_k q_k$ 
end
```

Notice that `cg_level2()` implements the pre-conditioner which consists of solving the same problem using only regional diagonal blocks of the the correlation matrix C .

The practical implementation below stops the iteration before exact convergence. Indeed, the iteration stops if we exceed a pre-determined maximum number of iterations or the residual is reduced by a specified number of orders of magnitude. These options are selected via the PSAS resource file (usually named `psas.rc`).

CALLING SEQUENCE:

```
call CG_MAIN ( verbose,
&               nkr, kr_beg, kr_len, kt_len,
&               nobs, ks, nsig_Ou, nsig_Oc, nsig_F,
&               qr_x, qr_y, qr_z, qm_x, qm_y, qm_z, ql_x, ql_y,
&               ktab, wtab, jtab, vtab,
&               nvecs, nobs_d, b, x, ierr )
```

INPUT PARAMETERS:

```

implicit NONE

logical verbose           ! if .true. prints out all kind
                          ! of informational output to stdout.

include 'ktmax.h'        ! maximun no. of data types
integer nkr               ! number of regions
integer kr_beg(nkr)      ! beginning of each region
integer kr_len(nkr)      ! no. of obs. in each region
integer kt_len(ktmax,nkr) ! no. of obs. of a given data type
                          ! in each region

integer nobs              ! number of observations
integer ks(nobs)          ! sounding index

                          ! Observation/forecast errors stdv
                          ! normalized by innovation (O-F) stdv:
real nsig_Ou(nobs)       ! o normalized spatially uncorrelated
                          ! observation error stdv
real nsig_Oc(nobs)       ! o normalized spatially correlated
                          ! observation error stdv
real nsig_F(nobs)        ! o normalized forecast error stdv

                          ! Cartesian coordinates (on the
                          ! unity sphere) of unit vectors
                          ! of the spherical coordinate system
real qr_x(nobs)           ! o x-coord of radial unit vector
real qr_y(nobs)           ! o y-coord of radial unit vector
real qr_z(nobs)           ! o z-coord of radial unit vector
real qm_x(nobs)           ! o x-coord of meridional unit vector
real qm_y(nobs)           ! o y-coord of meridional unit vector
real qm_z(nobs)           ! o z-coord of meridional unit vector
real ql_x(nobs)           ! o x-coord of longitudinal unit vector
real ql_y(nobs)           ! o y-coord of longitudinal unit vector
                          ! NOTE: ql_z is not needed.

integer ktab(nobs)        ! Interpolation indices/weights:
                          ! o vertical interpolation index
real wtab(nobs)           ! o vertical interpolation weights
integer jtab(nobs)        ! o meridional interpolation index
real vtab(nobs)           ! o meridional interpolation weights

```

```

integer  nvecs          ! Number of RHS vectors
integer  nobs_d         ! leading dimension of RHS vector
                        ! as declared in calling program.
                        ! Usually nobs_d = nobs.

real     b(nobs_d,nvecs) ! RHS vectors normalized by
                        ! innovation stdv. For the conventional
                        ! PSAS analysis system 'b' will
                        ! contain the normalized innovations
                        ! (O-F). However, multiple RHS will be
                        ! necessary for implementation
                        ! analysis error variances by
                        ! randomized trace estimates.

```

OUTPUT PARAMETERS:

```

real     x(nobs_d,nvecs) ! Solution vectors.
integer  ierr            ! error return code. All is well
                        ! if ierr=0.

```

SEE ALSO:

```

cg_level2()  Pre-conditioner routine.
stdio.h      Include file defining standard I/O units
BLAS         Basic linear algebra sub-programs

```

REVISION HISTORY:

```

03apr93  Pfaendtner  Original code
04jun93  Pfaendtner  Modification for dynamic storage on CRAY
07jan94  Sienkiewicz Added pass of trig lat/lon to subroutine
14feb94  da Silva    Fixed search direction bug
09apr94  Pfaendtner  Added prologue
13apr94  Pfaendtner  Added use of libsci routines
03oct94  da Silva    Implemented CRAY specifics with IFDEFs.
04oct94  da Silva    Routine changed name from CONJGR5 to

```

simply CONJGR. Introduced parameter nbandmx.
 19Jan95 Guo Added wobs tables to pass pindx2() values to ??cor1() and ??corx() routines. One could use rlevs for the same purpose to reduce the overhead, since rlevs has no real purpose in this subroutine and subsequent routines.
 02Feb95 Guo Changed CRAY to _UNICOS for consistency and to follow the guide lines.
 11Oct95 Guo Summary of changes since 02Feb95:
 + some structural changes for multitasking on C90, including now handling all regions in one conjgr2() call.
 + modified to accept multi vectors;
 + replaced multbyC() call to sCxy() call;
 06Feb96 da Silva Revised prologue, and several minor changes for readability:
 o name change: from conjgr() to cg_main()
 o removed static allocation IFDEFs; code now requires Fortran 90 for portability.
 o simplified main loop
 o several variable name changes to conform to notation in Golub and van Loan; comments are straight quotation from book.
 o introduction of f90 assignments whenever possible.

SOURCE CODE:

```

character*7 myname
parameter (myname='cg_main')

! Local storage (dynamic allocation)
! -----
include      'mypass.h'           ! max dimension for sizerr
real        x_k(nobs,nvecs)      ! solution at kth iteration
real        r_k(nobs,nvecs)      ! residual at kth iteration
real        z_k(nobs,nvecs)      ! pre-conditioner at kth iteration
real        p_k(nobs,nvecs)      ! search direction at kth iteration
real        Cp_k(nobs,nvecs)     ! Correlation matrix * p_k
real        r_norm(0:mypass,nvecs) ! residual norm

real        zTr_new(nvecs)       ! z' * r (new)
real        zTr_old              ! z' * r (old)
! where z' = transpose(z)
  
```



```

!   Defines kind of covariance matrices
!   -----
integer kind_mat, kind_cov
include 'kind_mats.h'
include 'kind_covs.h'

!   Convergence control parameters.
!   -----
include      'bands.h'

include      'stdio.h'

!   BLAS functions
!   -----
real        sdot, snrm2
external    sdot, snrm2

!   Minor local variables not worth commenting
!   -----
real        alpha_k, beta, tol
integer     k, kn, ivec, k_max
integer     i

!.....

      call ZEITBEG (cname(nbandcg))   ! starts timing

!   Initialization: k=0; x_0=0; r_0=b
!   -----
k = 0
x_k = 0.
do ivec=1,nvecs
  r_k(1:nobs,ivec) = b(1:nobs,ivec)
  r_norm(k,ivec) = SNRM2(nobs,r_k(1,ivec),1)
end do
kn = 0

!   Iterate...
!   -----
k_max = maxpass(nbandcg)
tol = criter(nbandcg)
DO WHILE ( k .le. k_max .and.
&         (r_norm(kn,1)/r_norm(0,1)) .gt. tol )

      k = k + 1

!   Pre-conditioner step: Solve  $\hat{C} z_k = r_k$ 
!   -----

```

```

call CG_LEVEL2 ( verbose.and.cgverb(2), kind_cov0.or.kind_covF,
&               nkr, kr_beg, kr_len, kt_len,
&               nob, ks, nsig_Ou, nsig_Oc, nsig_F,
&               qr_x, qr_y, qr_z, qm_x, qm_y, qm_z, ql_x, ql_y,
&               ktab, wtab, jtab, vtab,
&               nvecs, nob, r_k, z_k, ierr )

!
! Error handling
! -----
if ( ierr .ne. 0 ) then
  if ( ierr .lt. 0 ) then
    write(stderr,*) myname,
&       ': insufficient working space in cg_level2(), ',
&       'size = ',-ierr
  else
    write(stderr,'(3a,i6)') myname,
&       ': unexpected return from cg_level2(), ',
&       'err = ',ierr
  end if
  call PSASexit(2,myname)
end if

!
! Set search direction, p_k.
! -----
do ivec=1,nvecs

  if k = 1 { p_1 = z_0 }
  -----
  if( k.eq.1 ) then

    zTr_new(ivec) = SDOT(nob,r_k(1,ivec),1,z_k(1,ivec),1)
    call SCOPY(nob,z_k(1,ivec),1,p_k(1,ivec),1)

  else { beta_k = r_{k-1}^T z_{k-1} / r_{k-1}^T z_{k-2}
        p_k = z_{k-1} + \beta_k p_{k-1} }
  -----
  else

    zTr_old = zTr_new(ivec)
    zTr_new(ivec) = SDOT(nob,r_k(1,ivec),1,z_k(1,ivec),1)
    beta = zTr_new(ivec) / zTr_old
    call SAXPY(nob,beta,p_k(1,ivec),1,z_k(1,ivec),1)
    call SCOPY(nob,z_k(1,ivec),1,p_k(1,ivec),1)

  end if

end do      ! loop over RHS vectors

!
! q_k = C p_k
! -----

```

```

kind_mat=nbandcg
call sCxy ( kind_mat, kind_cov0 .or. kind_covF,
&          nkr, kr_beg, kr_len, kt_len,
&          nobs, ks, nsig_Ou, nsig_Oc, nsig_F,
&          qr_x, qr_y, qr_z, qm_x, qm_y, qm_z, ql_x, ql_y,
&          ktab, wtab, jtab, vtab,
&          nvecs, nobs, p_k, nobs, Cp_k,
&          ierr )

!
! Error handling
! -----
if ( ierr .ne. 0 ) then
  if(ierr.lt.0) then
    write(stderr,'(3a,i10)') myname,
&          ': insufficient working space in sCxy(), ',
&          'size = ',-ierr
  else
    write(stderr,'(3a,i3)') myname,
&          ': unexpected return from sCxy(), ',
&          'err = ',ierr
  end if
  call PSASexit(2,myname)
end if

!
! For each RHS vector
! -----
do ivec=1,nvecs

!
!   alpha_k = z_{k-1}^T r_{k-1} / p_k^T q_k
!   -----
&   alpha_k = zTr_new(ivec) /
&             SDOT(nobs,p_k(1,ivec),1,Cp_k(1,ivec),1)

!
!   x_k = x_{k-1} + alpha_k p_k
!   -----
  call SAXPY(nobs, +alpha_k, p_k(1,ivec),1,x_k(1,ivec),1)

!
!   r_k = r_{k-1} - alpha_k q_k
!   -----
  call SAXPY(nobs,-alpha_k,Cp_k(1,ivec),1,r_k(1,ivec),1)

end do

!
! Residual norm at end of this iteration
! -----
kn = kn + 1
if ( kn .gt. MXPASS ) kn = 1  ! cyclic storage
do ivec=1,nvecs
  r_norm(kn,ivec) = SNRM2(nobs,r_k(1,ivec),1)
end do

```

```

END DO ! end of CG iteration

! Convergence achieved
! -----
if( (r_norm(kn,1)/r_norm(0,1)) .le. tol .and. verbose ) then
    write(stdout,'(2a)') myname,': convergence achieved'

! Maximum number of iterations exceeded
! -----
else if ( verbose ) then
    write(stdout,'(2a)') myname,
&          ': maximum number of iterations exceeded'
end if

! Print summary
! -----
if ( verbose ) then
    call CGNORM ( myname, criter(nbandcg), mxpass,
&              k, nvecs, r_norm, nob)
end if

! Return kth iterate as solution
! -----
do ivec=1,nvecs
    x(1:nobs,ivec) = x_k(1:nobs,ivec)
end do

! All done
! -----
call ZEITEND

return
end

```

A.6 cg_level2()

Solves the linear system of equations

$$\tilde{C}x = b$$

where \tilde{C} is a simplified version of innovation covariance matrix, and b is a set of multiple RHS. The matrix \tilde{C} consists of regional diagonal blocks of the the correlation matrix C . This routine is meant to be a pre-conditioner for routine `cg_main()`. When performing a global analysis with PSAS, the RHS is simply the innovation (O-F) normalized by its standard deviation. The multiple RHS are necessary for the estimate of analysis error variances by means of randomized trace estimates.

The *Pre-conditioned Conjugate Gradient* algorithm is standard and closely follows

Golub, G. H. and C. F. van Loan, 1989: *Matrix Computations*, 2nd Edition, The John Hopkins University Press, 642pp.

and is reproduced in the prologue of routine `cg_main()`. The pre-conditioner for this routine is implemented in `cg_level1()`. This pre-conditioner solves a similar problem, this time univariately.

CALLING SEQUENCE:

```
call CG_LEVEL2 ( verbose, kind_cov,
&               nkr, kr_beg, kr_len, kt_len,
&               nobs, ks, nsig_Ou, nsig_Oc, nsig_F,
&               qr_x, qr_y, qr_z, qm_x, qm_y, qm_z, ql_x, ql_y,
&               ktab, wtab, jtab, vtab,
&               nvecs, nobs_d, b, x, ierr )
```

INPUT PARAMETERS:

logical	verbose	! if .true. prints out all kind ! of informational output to stdout.
integer	kind_cov	! specifies the kind of covariance ! matrix.
include	'ktmax.h'	! maximum no. of data types
integer	nkr	! number of regions
integer	kr_beg(nkr)	! beginning of each region
integer	kr_len(nkr)	! no. of obs. in each region
integer	kt_len(ktmax,nkr)	! no. of obs. of a given data type

```

! in each region

integer nobs ! number of observations
integer ks(nobs) ! sounding index

! Observation/forecast errors stdv
! normalized by innovation (O-F) stdv:
real nsig_Ou(nobs) ! o normalized spatially uncorrelated
! observation error stdv
real nsig_Oc(nobs) ! o normalized spatially correlated
! observation error stdv
real nsig_F(nobs) ! o normalized forecast error stdv

! Cartesian coordinates (on the
! unity sphere) of unit vectors
! of the spherical coordinate system
real qr_x(nobs) ! o x-coord of radial unit vector
real qr_y(nobs) ! o y-coord of radial unit vector
real qr_z(nobs) ! o z-coord of radial unit vector
real qm_x(nobs) ! o x-coord of meridional unit vector
real qm_y(nobs) ! o y-coord of meridional unit vector
real qm_z(nobs) ! o z-coord of meridional unit vector
real ql_x(nobs) ! o x-coord of longitudinal unit vector
real ql_y(nobs) ! o y-coord of longitudinal unit vector
! NOTE: ql_z is not needed.

! Interpolation indices/weights:
integer ktab(nobs) ! o vertical interpolation index
real wtab(nobs) ! o vertical interpolation weights
integer jtab(nobs) ! o meridional interpolation index
real vtab(nobs) ! o meridional interpolation weights

integer nvecs ! Number of RHS vectors
integer nobs_d ! leading dimension of RHS vector
! as declared in calling program.
! Usually nobs_d = nobs.

real b(nobs_d,nvecs) ! Normalized (by innovation stdv)
! RHS vectors. For the conventional
! PSAS analysis system 'b' will
! contain the innovations (O-F).
! However, multiple RHS will be
! necessary for implementation
! analysis error variances by
! randomized trace estimates.

```

OUTPUT PARAMETERS:

```
real      x(nobs_d,nvecs)  ! Solution vectors.
integer   ierr             ! error return code. All is well
                                ! if ierr=0.
```

SEE ALSO:

```
cg_level1()  Pre-conditioner routine.
stdio.h      Include file defining standard I/O units
BLAS         Basic linear algebra sub-programs
```

REVISION HISTORY:

```
03apr93  Pfaendtner  Original code
04jun93  Pfaendtner  Modification for dynamic storage on CRAY
07jan94  Sienkiewicz Added pass of trig lat/lon to subroutine
14feb94  da Silva    Fixed search direction bug
09apr94  Pfaendtner  Added prologue
13apr94  Pfaendtner  Added use of libsci routines
03oct94  da Silva    Implemented CRAY specifics with IFDEFs.
04oct94  da Silva    Routine changed name from CONJGR5 to
                    simply CONJGR. Introduced parameter
                    nbandmx.
19Jan95  Guo         Added wobs tables to pass pindx2() values to
                    ??cor1() and ??corx() routines. One could use
                    rlevs for the same purpose to reduce the over-
                    head, since rlevs has no real purpose in this
                    subroutine and subsequent routines.
02Feb95  Guo         Changed CRAY to _UNICOS for consistency and
                    to follow the guide lines.
11Oct95  Guo         Summary of changes since 02Feb95:
                    + some structural changes for multitasking on
                      C90, including now handling all regions in
                      one cg_level2() call.
                    + modified to accept multi vectors;
                    + replaced multbyC() call to sCxpy() call;
06Feb95  da Silva    Revised prologue, and several minor changes
                    for readability:
                    o name change: from conjgr2() to cg_level2()
                    o removed static allocation IFDEFs;
```

- code now requires Fortran 90 for portability.
- o simplified main loop
- o several variable name changes to conform to notation in Golub and van Loan;
- o comments are straight quotation from book.
- o introduction of f90 assignments whenever possible.

SOURCE CODE:

```

character*9 myname
parameter (myname='cg_level2')

! Local storage (dynamic allocation)
! -----
include      'mxpass.h'          ! max dimension for sizerr
real        x_k(nobs,nvecs)     ! solution at kth iteration
real        r_k(nobs,nvecs)     ! residual at kth iteration
real        z_k(nobs,nvecs)     ! pre-conditioner at kth iteration
real        p_k(nobs,nvecs)     ! search direction at kth iteration
real        Cp_k(nobs,nvecs)    ! Correlation matrix * p_k
real        r_norm(0:mxpass,nvecs) ! residual norm

real        zTr_new(nvecs)      ! z' * r (new)
real        zTr_old              ! z' * r (old)
! where z' = transpose(z)

integer     kt_beg(ktmax,nkr)
integer     lblkerr(ktmax*nkr)

! Minor local variables
! -----
real        alpha_k, beta, tol
integer     k, kn, ivec, k_max
integer     ibeg, ireg, ilen, kt, ik0x, ikFx
integer     i, ier, blk

! Convergence control parameters.
! -----
include      'bands.h'

include      'stdio.h'

! Defines kind of covariance matrices
! -----

```



```

integer    kind_mat
include    'kind_mats.h'
include    'kind_covs.h'

!         BLAS functions
!         -----
real       sdot, snrm2
external  sdot, snrm2

!.....

      call ZEITBEG (curname(2))

!         Initialization: k=0; x_0=0; r_0=b
!         -----
      k = 0
      x_k = 0.
      do ivec=1,nvecs
         r_k(1:nobs,ivec) = b(1:nobs,ivec)
         r_norm(k,ivec) = SNRM2(nobs,r_k(1,ivec),1)
      end do
      kn = 0

!         Iterate...
!         -----
      k_max = maxpass(2)
      tol = criter(2)
      DO WHILE ( k .le. k_max .and.
&             (r_norm(kn,1)/r_norm(0,1)) .gt. tol )

         k = k + 1

!         Loop over kt-blocks across regions. The data are sorted by
!         regions, and within each region the obs are sorted by data type
!         (kt). The loop here is over these kt-blocks...
!         -----
      do lblk = 1, ktmax*nkr

         lblkerr(lblk)=0

         ireg = (lblk-1)/ktmax+1
         kt   = mod(lblk-1,ktmax)+1

         ibeg = kt_beg(kt,ireg)
         ilen = kt_len(kt,ireg)

         ik0x=1
         if((kind_cov.and.kind_cov0).ne.0) ik0x=ibeg
         ikFx=1
         if((kind_cov.and.kind_covF).ne.0) ikFx=ibeg

```

```

!       If the kt-block is not empty...
!       -----
!       if ( ilen.gt.0 ) then
!
!           Invoke the pre-conditioner for each of these
!           univariate kt-blocks
!           -----
!           call CG_LEVEL1 ( verbose.and.cgverb(1), kind_cov,
!           &               ireg, kt, ilen, ks(ik0x),
!           &               nsig_Cu(ik0x), nsig_Oc(ik0x), nsig_F(ikFx),
!           &               qr_x(ibeg), qr_y(ibeg), qr_z(ibeg),
!           &               qm_x(ibeg), qm_y(ibeg), qm_z(ibeg),
!           &               ql_x(ibeg), ql_y(ibeg),
!           &               ktab(ibeg), wtab(ibeg), jtab(ikFx), vtab(ikFx),
!           &               nvecs, nob, r_k(ibeg,1),
!           &               z_k(ibeg,1), ier )
!
!           Error handling. Notice that zeitend() is not balanced,
!           but who cares, since there is a much more serious problem
!           -----
!           if(ier.ne.0) then
!               if(ier.lt.0) then
!                   write(stderr,'(3a,i10)') myname,
!                   &               ': insufficient working space in cg_level1(), ',
!                   &               'size = ',-ier
!               else
!                   write(stderr,'(2a,2(a,i3))') myname,
!                   &               ': unexpected return from cg_level1(), ',
!                   &               'err = ',ier,' with kt = ',kt
!               end if
!               lblkerr(lblk)=ier
!           end if
!
!           end if          ! kt-block is not empty
!
!       end do            ! loop over kt-blocks
!
!
!       Additional error handling. This apparently redundant
!       step is only necessary on a parallel enviroment
!       -----
!       ierr=0
!       do lblk=1,ktmax*nkr
!           if(lblkerr(lblk).ne.0) then
!               ierr=lblkerr(lblk)
!               return
!           end if
!       end do
!
!       Set search direction, p_k.
!       -----

```

```

do ivec=1,nvecs
!
!   if k = 1 { p_1 = z_0 }
!   -----
!   if( k.eq.1 ) then
!
!       zTr_new(ivec) = SDOT(nobs,r_k(1,ivec),1,z_k(1,ivec),1)
!       call SCOPY(nobs,z_k(1,ivec),1,p_k(1,ivec),1)
!
!   else { beta_k = r_{k-1}^T z_{k-1} / r_{k-1}^T z_{k-2}
!         p_k = z_{k-1} + \beta_k p_{k-1} }
!   -----
!   else
!
!       zTr_old = zTr_new(ivec)
!       zTr_new(ivec) = SDOT(nobs,r_k(1,ivec),1,z_k(1,ivec),1)
!       beta      = zTr_new(ivec) / zTr_old
!       call SAXPY(nobs,beta,p_k(1,ivec),1,z_k(1,ivec),1)
!       call SCOPY(nobs,z_k(1,ivec),1,p_k(1,ivec),1)
!
!   end if
!
! end do      ! loop over RHS vectors
!
! q_k = C p_k
! -----
! kind_mat=kind_Rmat
! call sCxy ( kind_mat, kind_cov,
&           nkr, kr_beg, kr_len, kt_len,
&           nobs, ks,nsig_Ou, nsig_Oc, nsig_F,
&           qr_x, qr_y, qr_z, qm_x, qm_y, qm_z, ql_x, ql_y,
&           ktab, wtab, jtab, vtab,
&           nvecs, nobs, p_k, nobs, Cp_k,
&           ierr )
!
! Error handling
! -----
! if ( ierr .ne. 0 ) then
!   if(ierr.lt.0) then
!     write(stderr,'(3a,i10)') myname,
&           ': insufficient working space in sCxy(), ',
&           'size = ',-ierr
!   else
!     write(stderr,'(3a,i3)') myname,
&           ': unexpected return from sCxy(), ',
&           'err = ',ierr
!   end if
!   call PSASexit(2,myname)
! end if

```

```

!       For each RHS vector
!       -----
!       do ivec=1,nvecs
!
!           alpha_k = z_{k-1}^T r_{k-1} / p_k^T q_k
!           -----
!           alpha_k = zTr_new(ivec) /
&               SDOT(nobs,p_k(1,ivec),1,Cp_k(1,ivec),1)
!
!           x_k = x_{k-1} + alpha_k p_k
!           -----
!           call SAXPY(nobs,+alpha_k, p_k(1,ivec),1,x_k(1,ivec),1)
!
!           r_k = r_{k-1} - alpha_k q_k
!           -----
!           call SAXPY(nobs,-alpha_k,Cp_k(1,ivec),1,r_k(1,ivec),1)
!
!       end do
!
!       Residual norm at end of this iteration
!       -----
!       kn = kn + 1
!       if ( kn .gt. MXPASS ) kn = 1   ! cyclic storage
!       do ivec=1,nvecs
!           r_norm(kn,ivec) = SNRM2(nobs,r_k(1,ivec),1)
!       end do
!
!       END DO ! end of CG iteration
!
!       Convergence achieved
!       -----
!       if( (r_norm(kn,1)/r_norm(0,1)) .le. tol .and. verbose ) then
!           write(stdout,'(2a)') myname,': convergence achieved'
!
!       Maximum number of iterations exceeded
!       -----
!       else if ( verbose ) then
!           write(stdout,'(2a)') myname,
&               ': maximum number of iterations exceeded'
!
!       end if
!
!       Prints summary
!       -----
!       if ( verbose ) then
!           call CGNORM ( myname, criter(2), mxpass, k, nvecs, r_norm, nobs )
!       end if

```

```
!   Return kth iterate as solution
!   -----
do ivec=1,nvecs
    x(1:nobs,ivec) = x_k(1:nobs,ivec)
end do

!   All done
!   -----
call ZEITEND

return
end
```

A.7 cg_level1()

Solves the linear system of equations

$$\hat{C}x = b$$

where \hat{C} is a simplified version of innovation covariance matrix, and b is a set of multiple right-hand-sides. The matrix \hat{C} consists of regional diagonal blocks of the the correlation matrix C . This routine is meant to be a pre-conditioner for routine `cg_level2()`. When performing a global analysis with PSAS, the RHS is simply the innovation (O-F) normalized by its standard deviation. The multiple RHS are necessary for the estimate of analysis error variances by means of randomized trace estimates.

The *Pre-conditioned Conjugate Gradient* algorithm is standard and closely follows

Golub, G. H. and C. F. van Loan, 1989: *Matrix Computations*, 2nd Edition, The John Hopkins University Press, 642pp.

and is reproduced in the prologue of routine `cg_main()`. The pre-conditioner for this routine is implemented using LAPACK's Cholesky solver [routines `spptrf()` and `spptrs()`]. This pre-conditioner solves a much smaller problem, considering only diagonal blocks of C with a "couple" of profiles.

CALLING SEQUENCE:

```
      call CG_LEVEL1 ( verbose, kind_cov,
&                    ireg, kt,
&                    nobs, ks, nsig_Ou, nsig_Oc, nsig_F,
&                    qr_x, qr_y, qr_z, qm_x, qm_y, qm_z, ql_x, ql_y,
&                    ktab, wtab, jtab, vtab,
&                    nvecs, nobs_d, b, x, ierr )
```

INPUT PARAMETERS:

```
implicit NONE

logical  verbose           ! if .true. prints out all kind
                          ! of informational output to stdout.

integer  kind_cov         ! specifies the kind of covariance
                          ! matrix

integer  ireg             ! PSAS region index
integer  kt              ! GEOS/DAS data-type index
```

```

integer  nobs          ! number of observations
integer  ks(nobs)     ! sounding index

                                ! Observation/forecast errors stdv
                                ! normalized by innovation (O-F) stdv:
real     nsig_Ou(nobs) ! o normalized spatially uncorrelated
                                ! observation error stdv
real     nsig_Oc(nobs) ! o normalized spatially correlated
                                ! observation error stdv
real     nsig_F(nobs)  ! o normalized forecast error stdv

                                ! Cartesian coordinates (on the
                                ! unity sphere) of unit vectors
                                ! of the spherical coordinate system
real     qr_x(nobs)    ! o x-coord of radial      unit vector
real     qr_y(nobs)    ! o y-coord of radial      unit vector
real     qr_z(nobs)    ! o z-coord of radial      unit vector
real     qm_x(nobs)    ! o x-coord of meridional  unit vector
real     qm_y(nobs)    ! o y-coord of meridional  unit vector
real     qm_z(nobs)    ! o z-coord of meridional  unit vector
real     ql_x(nobs)    ! o x-coord of longitudinal unit vector
real     ql_y(nobs)    ! o y-coord of longitudinal unit vector
                                ! NOTE: ql_z is not needed.

                                ! Interpolation indices/weights:
integer  ktab(nobs)    ! o vertical  interpolation index
real     wtab(nobs)    ! o vertical  interpolation weights
integer  jtab(nobs)    ! o meridional interpolation index
real     vtab(nobs)    ! o meridional interpolation weights

integer  nvecs         ! Number of RHS vectors
integer  nobs_d         ! leading dimension of RHS vector
                                ! as declared in calling program.
                                ! Usually nobs_d = nobs.

real     b(nobs_d,nvecs) ! Normalized (by innovation stdv)
                                ! RHS vectors. For the convetional
                                ! PSAS analysis system 'b' will
                                ! contain the innovations (O-F).
                                ! However, multiple RHS will be
                                ! necessary for implementation
                                ! analysis error variances by
                                ! randomized trace estimates.

```

OUTPUT PARAMETERS:

```
real      x(nobs_d,nvecs)    ! Solution vectors.
integer   ierr               ! error return code. All is well
                                ! if ierr=0.
```

SEE ALSO:

```
stdio.h   Include file defining standard I/O units
LAPACK    Linear Algebra PACKage
BLAS      Basic linear algebra sub-programs
```

REVISION HISTORY:

```
03apr93   Pfaendtner   Original code
04jun93   Searl        Modification for dynamic storage on CRAY
07jan94   Sienkiewicz  Added pass of trig lat/lon to subroutine
14feb94   da Silva     Fixed search direction bug
09apr94   Pfaendtner   Added prologue
13apr94   Pfaendtner   Added use of libsci routines
03oct94   da Silva     Implemented CRAY specifics with IFDEFs.
04oct94   da Silva     Routine changed name from CONJGR5 to
                        simply CONJGR. Introduced parameter
                        nbandmx.
19Jan95   Guo          Added wobs tables to pass pindx2() values to
                        ??cor1() and ??corx() routines. One could use
                        rlevs for the same purpose to reduce the over-
                        head, since rlevs has no real purpose in this
                        subroutine and subsequent routines.
02Feb95   Guo          Changed CRAY to _UNICOS for consistency and
                        to follow the guide lines.
11Oct95   Guo          Summary of changes since 02Feb95:
                        + some structural changes for multitasking on
                          C90, including now handling all regions in
                          one cg_level2() call.
                        + modified to accept multi vectors;
                        + replaced multbyC() call to sCxpy() call;
06Feb95   da Silva     Revised prologue, and several minor changes
                        for readability:
                        o name change: from conjgr1() to cg_level1()
                        o removed static allocation IFDEFs;
```


- code now requires Fortran 90 for portability.
- o simplified main loop
- o several variable name changes to conform to notation in Golub and van Loan; comments are straight quotation from book.
- o introduction of f90 assignments whenever possible.

SOURCE CODE:

```

character*9 myname
parameter (myname='cg_level1')

! Local storage (dynamic allocation)
! -----
include 'mxpass.h'

real corr(nobs*(nobs+1)/2) ! Temporary correlation matrix
real corrM(nobs*(nobs+1)/2) ! Innovation correlation matrix
real corrI(nobs*(nobs+1)/2) ! Inverse of corrM
real x_k(nobs,nvecs) ! solution at kth iteration
real r_k(nobs,nvecs) ! residual at kth iteration
real z_k(nobs,nvecs) ! pre-conditioner at kth iteration
real p_k(nobs,nvecs) ! search direction at kth iteration
real Cp_k(nobs,nvecs) ! Correlation matrix * p_k
real r_norm(0:mxpass,nvecs) ! residual norm

real zTr_new(nvecs) ! z' * r (new)
real zTr_old ! z' * r (old)

! Minor local storage (static allocation)
! -----
integer ivec
integer k_max
real tol
integer begin_blk, next_blk, begin_sav
real endqrx
logical next

character*1 Mtyp
integer ij

real alpha_k, beta
integer N_diverg
integer k, m, i, j, kn, km
logical converging

```

```

logical    solved
integer    nshift
integer    mshift
parameter  (mshift=10)
real      dshift
parameter  (dshift=.1/mshift)

include    'bands.h'          ! Convergence control parameters
include    'stdio.h'          ! standard I/O
include    'realvals.h'       ! machep look-alike
include    'kind_covs.h'      ! kind of covariance matrices

logical    setCorF
parameter  (setCorF=.true.)

! BLAS functions
! -----
real      sdot, snrm2
external  sdot, snrm2

integer    lnblk, luavail
external  lnblk, luavail

! .....

call ZEITBEG (curname(1))

! Initialization: k=0; x_0=0; r_0=b
! -----
k = 0
x_k = 0.
do ivec=1,nvecs
  r_k(1:nobs,ivec) = b(1:nobs,ivec)
  r_norm(k,ivec) = SNRM2(nobs,r_k(1,ivec),1)
end do
kn = 0

! Compute the block matrix corrM to work on
! -----
corr = 0.
corrI = 0.
corrM = 0.
call CG_BLOCKS()              ! this an internal routine

```

```

! Iterate...
! -----
k_max = maxpass(1)
tol = criter(1)
N_diverg = 0
converging = .true.
DO WHILE ( converging .and.
&         k .le. k_max .and.
&         (r_norm(kn,1)/r_norm(0,1)) .gt. tol )

    k = k + 1

! Preconditioner for level 1 (one region, one kt) is direct
! solver on diagonal sub-blocks. It makes sure that
! soundings are kept together (group by qr_x)
! -----
call CG_LEVEL0()          ! this an internal routine

! if k = 1 { p_1 = z_0 }
! -----
if( k.eq.1 ) then
    do ivec=1,nvecs
        zTr_new(ivec) = SDOT(nobs,r_k(1,ivec),1,z_k(1,ivec),1)
        call SCOPY(nobs,z_k(1,ivec),1,p_k(1,ivec),1)
    end do

! else { beta_k = r_{k-1}^T z_{k-1} / r_{k-1}^T z_{k-2}
!       p_k = z_{k-1} + \beta_k p_{k-1} }
! -----
else
    do ivec=1,nvecs
        zTr_old = zTr_new(ivec)
        zTr_new(ivec) = SDOT(nobs,r_k(1,ivec),1,z_k(1,ivec),1)
        beta = zTr_new(ivec) / zTr_old
        call SAXPY(nobs,beta,p_k(1,ivec),1,z_k(1,ivec),1)
        call SCOPY(nobs,z_k(1,ivec),1,p_k(1,ivec),1)
    end do
end if

! For each RHS vector
! -----
do ivec=1,nvecs

!     q_k = C p_k
!     -----
!     call SSPMV('U',nobs, 1.,corrM,p_k(1,ivec),1,
&              0., Cp_k(1,ivec),1)

!     alpha_k = z_{k-1}^T r_{k-1} / p_k^T q_k
!     -----

```

```

      alpha_k = zTr_new(ivec) /
&          SDOT(nobs,p_k(1,ivec),1,Cp_k(1,ivec),1)

!      x_k = x_{k-1} + alpha_k p_k
!      -----
!      call SAXPY(nobs,+alpha_k, p_k(1,ivec),1,x_k(1,ivec),1)

!      r_k = r_{k-1} - alpha_k q_k
!      -----
!      call SAXPY(nobs,-alpha_k,Cp_k(1,ivec),1,r_k(1,ivec),1)

      end do

!      Residual norm at end of this iteration
!      -----
      km = kn
      kn = kn + 1
      if ( kn .gt. MXPASS ) kn = 1   ! cyclic storage
      do ivec=1,nvecs
          r_norm(kn,ivec) = SNRM2(nobs,r_k(1,ivec),1)
      end do

!      Detect divergence: one iteration is termed "divergent" if the
!      residual increases instead of decreasing. N_diverg
!      records how many times this happens
!      -----
      if ( r_norm(kn,ivec) .ge. r_norm(km,ivec) ) then
          N_diverg = N_diverg + 1
      end if

!      The CG process is called "divergent" if the number
!      of divergent iterations exceeds a pre-determined
!      number (minmax(1))
!      -----
      converging = N_diverg .lt. minmax(1)

      END DO ! end of CG iteration

!      Convergence achieved
!      -----
      if( (r_norm(kn,1)/r_norm(0,1)) .le. tol .and. verbose ) then
          write(stdout,'(2a)') myname,': convergence achieved'

!      Divergence detected
!      -----
      else if ( .not. converging .and. verbose ) then
          write(stdout,'(2a)') myname,

```

```

&          ': conjugate gradient is not converging. '

!   Maximum number of iterations exceeded
!   -----
!   else if ( verbose ) then
!       write(stdout,'(2a)') myname,
&          ': maximum number of iterations exceeded'

!   end if

!   Prints summary
!   -----
!   if ( verbose ) then
!       call CGNORM ( myname, criter(1), mxpass, k, nvecs, r_norm, nobs )
!   end if

!   Return kth iterate as solution
!   -----
!   do ivec=1,nvecs
!       x(1:nobs,ivec) = x_k(1:nobs,ivec)
!   end do

!   All done
!   -----
!   call ZEITEND

!   return

!   CONTAINS
!   -----

```

A.8 cg_blocks()

Computes innovation correlation blocks. This is an internal routine of CG_LEVEL1().

CALLING SEQUENCE:

```
call cg_blocks()
```

INPUT PARAMETERS:

none.

OUTPUT PARAMETERS:

None explicitly, but corrM is calculated here.

SEE ALSO:

cg_level1() parent routine.

REVISION HISTORY:

06Feb96 da Silva Moved from body of CG_LEVEL1 for readability.

SOURCE CODE:

```

Mtyp='Z'
if ((kind_cov.and.kind_cov0).ne.0) then

!       Construct spatially correlated observation error correlation
!       matrix
!       -----
!       call DiagCor0 ( kt,nobs,ks,qr_x,qr_y,qr_z,ktab,wtab,
&                   Mtyp,corr,ierr)

!       Error handling
!       -----
!       if(ierr.ne.0) then
!         write(stderr,'(a,2(a,i3))') myname,
&           ': unexpected variable type for diagcor0(), kt = ',kt,
&           ', ierr =',ierr
!         return
!       end if

!       If ( Mtyp.eq.'U' .or. Mtyp.eq.'u' ) then
!         do j=1,nobs
!           ij=j*(j-1)/2
!           do i=1,j
!             corrM(ij+i)=nsig_0c(i)*corr(ij+i)*nsig_0c(j) + corrM(ij+i)
!           end do
!         end do
!       else if ( Mtyp .eq. 'I' .or. Mtyp .eq. 'i' ) then
!         do j=1,nobs
!           ij=j*(j+1)/2
!           corrM(ij)=nsig_0c(j)*nsig_0c(j) + corrM(ij)
!         end do
!       end if

!       Construct uncorrelated observation error correlation
!       -----
!       call DiagCorU (kt,nobs,ks,ktab,wtab,Mtyp,corr,ierr)

!       Error handling
!       -----
!       if(ierr.ne.0) then
!         write(stderr,'(a,2(a,i3))') myname,
&           ': unexpected variable type for diagcorU(), kt = ',kt,
&           ', ierr =',ierr
!         return
!       end if

```

```

      If(Mtyp.eq.'U'.or.Mtyp.eq.'u') then
        do j=1,nobs
          ij=j*(j-1)/2
          do i=1,j
            corrM(ij+i)=nsig_Du(i)*corr(ij+i)*nsig_Du(j) + corrM(ij+i)
          end do
        end do
      elseif(Mtyp.eq.'I'.or.Mtyp.eq.'i') then
        do j=1,nobs
          ij=j*(j+1)/2
          corrM(ij)=nsig_Du(j)*nsig_Du(j) + corrM(ij)
        end do
      end if

    end if

    Mtyp='Z'
    if ((kind_cov.and.kind_covF).ne.0) then

      call DiagCorF ( kt,nobs,qr_x,qr_y,qr_z,qm_x,qm_y,qm_z,
&                    ql_x,ql_y,ktab,wtab,
&                    Mtyp,corr,ierr)

!      Error handling
!      -----
      if(ierr.ne.0.or.Mtyp.eq.'E') then
        write(stderr,'(a,2(a,i3))') myname,
&          ': unexpected variable type for diagcorF(), kt = ',kt,
&          ', ierr = ',ierr
        return
      end if

      if(Mtyp.eq.'U'.or.Mtyp.eq.'u') then
        do j=1,nobs
          ij=j*(j-1)/2
          do i=1,j
            corrM(ij+i)=nsig_F(i)*corr(ij+i)*nsig_F(j) + corrM(ij+i)
          end do
        end do
      end if
    end if

!    Al done
!    -----
    return

  end subroutine CG_BLOCKS

```


A.9 `cg_level0()`

Implements the pre-conditioner for `cg_level1()`. The pre-conditioner for level 1 (one region, one kt) is direct solver on diagonal sub-blocks. It makes sure that soundings are kept together (group by `qr_x`). This is an internal routine of `CG_LEVEL1()`.

CALLING SEQUENCE:

```
call cg_level0()
```

INPUT PARAMETERS:

none.

OUTPUT PARAMETERS:

None explicitly, but `z_k` is calculated here.

SEE ALSO:

`cg_level1()` parent routine.

REVISION HISTORY:

06Feb96 da Silva Moved from body of `CG_LEVEL1` for redability.

SOURCE CODE:

```

!      Make a copy of the current residual
!      -----
do ivec=1,nvecs
  call SCOPY(nobs,r_k(1,ivec),1,z_k(1,ivec),1)
end do

begin_blk = 1
begin_sav = 1
DO WHILE ( begin_blk .le. nobs )

!      It (next_blk) is actually the end-of-this-block
!      -----
next_blk = min(begin_blk+msmall-1,nobs)
endqrx = qr_x(next_blk)

!      Search for end of this sounding (at end of msmall sized
!      block) and set block break where soundings change
!      Tests are made in sequence to avoid qr_x(nobs+1) ever
!      being referenced.
!      -----
next=.true.
do while ( next )
  next_blk = next_blk + 1

  next=next_blk.le.nobs
  if(next) next=qr_x(next_blk).eq.endqrx
end do

m = next_blk - begin_blk

if(k.eq.1) then

  nshift=0
  solved=.false.
  call smex(corrM,nobs,begin_blk,m,corrI(begin_sav))

  do while(.not.solved)
    call SPTRF('U',m,corrI(begin_sav),ierr)

    if( ierr.ne.0 ) then

      write(stdout,'(a,5(a,i3),a,i5)') myname,
&          ': SPTRF() error ',ierr,
&          ': nshift=',nshift,
&          ' region=',ireg,
&          ' type=',kt,
&          ' msmall=',m,
&          ' begblk=',begin_blk

```

```

nshift=nshift+1
if(nshift.gt.mshift) then
  write(stderr,'(a,2(a,i4),a)') myname,
  &       ': err = ',ierr,' in SPTRF() after ',nshift,
  &       ' tries'
  return
end if

call smex(corrM,nobs,begin_blk,m,corrI(begin_sav))
call smexsh(corrI(begin_sav),m,nshift*dshift)

else
  solved=.true.
end if      ! error
end do      ! .not.solved
end if      ! k.eq.1

call SPPTRS('U',m,nvecs,corrI(begin_sav),z_k(begin_blk,1),
&          nobs,ierr)

if(ierr.ne.0) then      ! if it ever happens.
  write(stderr,'(a,2(a,i2))') myname,
  &       ': err = ',ierr,' from SPPTRS() with m = ',m,
  &       ' and begin_blk = ',begin_blk
  return
end if

begin_blk = next_blk
begin_sav = begin_sav+m*(m+1)/2

end do      ! next block (starting from begin_blk)?

! All done
! -----
return
end subroutine CG_LEVEL0

!.....

end subroutine CG_LEVEL1

```

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