An Enhanced Global Spectral Model

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This report describes the development of a vectorized, multiprocessing global spectral model (GSM) with enhanced physical parameterizations. The starting point was the Phillips Laboratory GSM, in its GL90 version, containing an enhanced suite of physical parameterizations. The latitude tasking scheme for multiprocessing the loop over latitude in the calculation of the spectral tendencies and adjusted model variables was implemented, using the general truncation version of the hydrodynamics code. Wavenumber calculations were vectorized over wavenumber and multiprocessed over vertical level. All gridpoint calculations were vectorized over longitude, and the physics packages were brought into closer compliance with plug-compatibility rules. Speedups due to the optimization were demonstrated in single- and multiprocessed timing tests on a dedicated Cray 2 and Cray Y-MP. The enhanced physics GSM was evaluated in forecast tests, and contrasted with the simpler physics GSM used at GWC. Minimization of errors in the computation of the horizontal pressure gradient force was investigated, using a perturbation temperature instead of full temperature in the integration of the hydrostatic equation. Several schemes were tested in an idealized model atmosphere.
TABLE OF CONTENTS

1. Introduction ................................................................. 1

2. Description of model code .............................................. 3
   2.1 Hydrodynamics and overall design ............................... 5
   2.2 Radiation .................................................................... 6
   2.3 Planetary Boundary Layer ........................................... 6
   2.4 Adjustment Physics .................................................... 7

3. Timing Results ............................................................... 8
   3.1 Single-Processor Timings ........................................... 8
   3.2 Multi-Processor Timings ............................................ 12
   3.3 Overall Speedups ..................................................... 14

4. Forecast Results ............................................................ 15

5. Minimization of pressure gradient force errors .................. 27
   5.1 Pressure-gradient force errors ........................................ 27
   5.2 Error minimization ................................................... 28
      5.2.1 Schemes .......................................................... 31
         5.2.1.1 The PG - scheme ......................................... 31
         5.2.1.2 The MPG -scheme ......................................... 32
         5.2.1.3 The null scheme - an isothermal case .......... 32
      5.2.2 Code modification ................................................. 32
         5.2.2.1 Model equations .......................................... 34
1. Introduction

This report describes the work performed by AER under contract F19628-89-C-0112 for the US Air Force Phillips Laboratory, Geophysics Directorate (hereafter referred to simply as PL), entitled "An Enhanced Global Spectral Model", for the entire period of the contract, 27 September 1989 to 27 November 1991. The purpose of the work was the development of a vectorized, multiprocessing global spectral model (GSM) with enhanced physical parameterizations. The starting point was the Phillips Laboratory (then called Geophysics Laboratory, or GL) GSM, in its GL90 version.

The global spectral model developed by PL has evolved from the GSM used at NMC as described in (Sela, 1980). The hydrodynamics were completely redesigned (Brenner et al., 1982, 1984). This model has been used by the Air Force Global Weather Central (GWC) with the physics routines taken almost intact from NMC (circa 1983). More recently, an enhanced suite of physical parameterizations has been developed by different investigators and integrated by PL personnel. The new physics routines consist of a coupled boundary layer and soil model (Mahrt et al., 1984, 1987), a radiation parameterization (Liou et al., 1984; Ou and Liou, 1988), and a modified Kuo convection parameterization (Soong et al., 1985 and Norquist and Yang, 1990). The new physics routines have been used in a research mode in a number of studies by PL personnel which demonstrated their potential for improving forecast skill.

As originally implemented, the new physics package required significantly more CPU time than the baseline GWC physics. Thus, before the potentially better forecast skill could be realized in the operational environment, the execution time of the enhanced global spectral model had to be reduced.

Implementation of the code on the Cray 2 offers two major avenues for reducing wall-clock execution time: vectorization and parallelism. Vectorization reduces the CPU time (and, hence, the wall-clock time) spent by a single processor by performing identical (or similar) operations on a number of storage locations in a pipeline fashion. This technique was
pioneered by Cray Research, and extensive support tools for vectorizing scalar codes are now available as part of compilers on Cray and other vector machines. To take full advantage of the potential speed-ups, however, it is necessary to take into account vectorization considerations during the design phase of program development.

A more recent trend in computer technology is the construction of machines composed of a number of processors linked together, allowing multiple simultaneous computations. Such multiprocessing computers contain two or more, in some cases tens of thousands, of individual processing units. These processors may operate in lock step or they may be nearly autonomous. The former case, single instruction multiple data (SIMD), is particularly relevant for image processing applications, while the latter case, multiple instruction multiple data (MIMD), is of more general interest and includes the Cray 2 and Y-MP. Multiprocessor computers give a substantial increase in computing speed for large complex numerical problems. Thus, they provide an opportunity to reduce significantly the computing time for GCM studies and NWP. However, a special effort is required to exploit this opportunity. According to the recommendations of GARP Special Report No. 43 (WMO, 1985), "Increasing computer power and changes in computer architecture...[have] always played a major part in determining integration techniques and will continue to do so if advances in computer technology are to be fully exploited. Therefore, study to determine the most efficient or appropriate algorithms will continue to be necessary, taking into account, for example, the future use of multiprocessors with a very large storage. Development costs of such optimized computer codes will be high, and they should be designed from the outset to be widely usable."

As an add-on to the original contract, we investigated ways of minimizing errors in the computation of the pressure-gradient force in the sigma-coordinate system. This work concentrated on the use of perturbation temperature in the integration of the hydrostatic equation, and was prompted by work of Simmons and Chen (1991).

The design of the optimized version of the GSM is described in detail in the interim technical report of this contract (Nehrkorn et al., 1990;
referred to as I in the following). The main points of the design, and
differences in the final implementation from the original design, are
discussed in Section 2. Detailed user documentation in the form of a user's
guide and an annotated list of model variables are provided as Appendix A
and B, respectively; the appendices are also included as plain text files in
machine-readable form, with the delivered model code. Section 3 describes
the results of timing tests of the original and optimized version of the GSM
with enhanced physics, along with a version of the GSM with simple
physics (similar to what is used operationally at the US Air Force Global
Weather Central - GWC). Section 4 contains the results from 4-day
forecasts, using the GSM with enhanced and simple physics. Section 5
describes results of the study to minimize errors in the computation of the
horizontal pressure gradient force. A summary and concluding remarks
form Section 6, and Section 7 contains the references.

2. Description of model code

The starting point for our optimization effort was the GL90 version of
the GSM. The physics modules of this version consisted of the radiation
module with a one-cloud deck structure, the PBL scheme with globally
uniform soil and vegetation characteristics, and the Kuo convection
scheme. Simultaneous with our recoding for computational efficiency,
testing and tuning of the physics packages were carried out by PL
personnel, and we were able to incorporate some of the resulting changes
into our optimized design. In particular, we used a three-deck version of
the radiation for optimization, and this was incorporated both into the
"original" and optimized code used in the forecast and timing tests.
Additional minor changes in the Kuo convection and PBL scheme were also
incorporated at the same time. We refer to this version of the optimized
model code as version 5.1 (all modules of the optimized code are archived
with the UNIX Source Code Control System (SCCS), and all are identified by
the SCCS sequence number 5.1).

Further modifications were made to the radiation parameterization
that could not be incorporated into the model in time for the forecast and
timing tests. These changes involved the use of latitudinally varying
climatologies of ozone, and replacement of the Geleyn cloud parameteriza-
tion by the scheme of Slingo (1987). The latter also required some minor code changes to the convection and hydrodynamics modules. For our work with the GSM under another contract, these changes were incorporated into the optimized GSM, and the resulting model version was archived as version 5.2. Both versions of the code have been delivered, and the user's guide document (Appendix A) is applicable to both versions.

There have been other improvements made to the physics modules by PL personnel that have not yet been incorporated into the optimized model code. Most notably, these are the addition of a gravity wave drag scheme, the replacement of the Kuo convection scheme by a mass flux scheme, and the use of geographical databases of soil and vegetation characteristics.

Our changes to the original code were guided by two general strategies.

*Enhance clarity.* Clear, direct coding is best. Coding which makes special use of a particular machine architecture and compiler is in the long run counterproductive. Eventually, and probably sooner than later, the compiler will be updated or the code will be transported to other machines. Generally, as time goes on, compilers get more and more capable and what seemed to be necessary last year is now a hindrance. Complicated coding strategies to squeeze the maximum horsepower out of the CRAY2 will be restricted to very specific and isolated procedures. For the most part these procedures will be standard library routines for which vanilla versions are readily available. An example is the FFT software.

*Avoid impacts.* Do not make changes which will have a big impact on the results. Changes which result in small changes in the tendencies, which are not less correct are fine. Some differences in the calculated tendencies are unavoidable. It is expected that different versions of the model, or the same version compiled differently will usually result in a different order of arithmetic and hence in different round off errors. On the other hand changes which result in large differences in the tendencies, even though they may be as correct as the original, should be avoided. For example, one could argue that a speedup is possible by using a T150 truncation instead of a R120 truncation. However, demonstrating that this is
indeed the case, would require a substantial suite of test cases, well beyond the scope of the present effort.

2.1 Hydrodynamics and overall design

The latitude tasking scheme described in I was implemented, using the general truncation version of the hydrodynamics code developed under a previous contract. The multitasking was implemented through Cray microtasking directives (these are treated as comments if Cray multitasking is deactivated, or if other compilers are used). Separation of data into shared (global) and private (i.e., local to each latitude task) was accomplished with Cray Fortran TASK COMMON statements - on single-processor computers, these would be replaced by simple COMMON statements, and on other multi-processing computers with language extensions analogous to the Cray TASK COMMON statement.

The general truncation version (documented in Kaplan et al., 1985) supports any pentagonal truncation, including the frequently used rhomboidal and triangular truncations. In addition, computations may be performed over only one hemisphere or a section of the globe. Array dimensions are given in terms of PARAMETER statements. The parameter statements and the common blocks are stored in include files, which are incorporated into the code at compile time. Edit symbols are retained, however, to allow to choose between the hemispheric and global versions of the code, and to use double precision for certain, sensitive calculations (useful for use of the code on 32-bit machines; see Nehrkorn, 1990). This version of the code also makes use of Hoffman's (1985) semi-implicit time stepping scheme, which uses full temperature (as opposed to perturbation temperature) throughout. This does not preclude introducing a basic state and perturbation temperature for the integration of the hydrostatic equation. In general, the basic state temperature used for the hydrostatic equation and that used for the semi-implicit time scheme will be different. Therefore it is best to use full temperature throughout and introduce linearizations where needed. In our study described in Section 5, the basic state temperature is a function of pressure (which depends on latitude, longitude, and time for each sigma layer), and the hydrostatic calculation is integrated in grid point space; the semi-implicit scheme still uses a
linearization about a globally uniform basic state temperature as a function of sigma.

The input/output was adapted to the PL practice, with minor changes and corrections (see Appendix A for details).

The gridpoint calculation in the hydrodynamic part of the code (in subroutine NLPROD) were completely recoded such that longitude was the innermost loop index, resulting in much improved vectorization.

2.2 Radiation

The GL90 version of the radiation, modified for a three-deck cloud structure, was the starting point for this work. The code was redesigned for plug-compatibility and vectorization as described in I for version 5.1 of the GSM. In particular, algorithms were redesigned to allow efficient vectorization, with longitude as the inner-most loop index. The holding file of radiative heating rates (needed for time steps when radiation computations are not performed) has been replaced by in-core storage in a global array. On the Cray 2, this did not lead to memory problems, even for the high-resolution runs (rhomboidal truncation at wavenumber 120, R120). However, we have included code to store these data in a direct access file which could be accessed simultaneously by the different latitude tasks.

As discussed above, a version of the radiation code was delivered (as version 5.2 of the model code) that included several enhancements: global climatology of ozone, and Slingo (1987) cloud parameterizations. This version of the code was not used in the forecast and timing tests reported below, however.

2.3 Planetary Boundary Layer

We implemented the design proposed in I by reordering all loops such that longitude was the innermost index. Vertical loop limits that were dependent on longitude had to be replaced by flags that indicated regions where calculations would or would not be performed. Similarly, surface computations that depend on the surface type (snow, land, or ocean) are now handled by flags. Computations that were unnecessary
were eliminated (commented out) - most of these involve calculations of boundary layer clouds, which were not used anywhere in the original code.

 Diagnostic calculations were isolated in a separate entry point, which is called after all the latitude task computations are completed. Values needed for diagnostic calculations are stored in internal common blocks. In the original code, the diagnostic computations used algorithms that are not identical to those used in the actual tendency calculations; we have noted these inconsistencies, but have not made any substantive changes to the diagnostic computations.

 The plug-compatibility rules are adhered to for the most part, except that we still use common blocks (in ground.blk) to pass values of boundary data sets between the PBL and the rest of the GSM.

 2.4 Adjustment Physics

 The adjustment physics consists of three separate, plug-compatible modules: dry adiabatic adjustment, large-scale precipitation, and Kuo convection. The cover routine for these routines performs spectral transforms to and from gridpoint space, and was adapted from the general truncation version of the GSM.

 The large-scale precipitation module was already vectorized in the original code, and we only changed the interface to the cover routine in order to bring it into compliance with the plug-compatibility rules.

 The dry adiabatic adjustment was redesigned for vectorization over longitude, as described in detail in I (the final version of the code differs slightly from the listing reproduced in I, however). The interface was also changed for plug-compatibility.

 We had originally developed a vectorized design for the Kuo convection (as described in I), but did not pursue its implementation and testing to completion when it became evident that the final physics package assembled by PL would not contain a Kuo scheme. Instead, we implemented the original code, with changes only to the interface and the initialization of physical constants.
3. Timing Results

3.1 Single-Processor Timings

Three versions of the GSM were used for timing tests. The first, denoted GWC in the following, consists of the optimized GSM code for the hydrodynamics and physics modules that emulate current GWC operational practice. The GWC physics does not contain any radiative parameterization, and a very simple drag-law type boundary layer parameterization. The adjustment physics consists of the same large-scale precipitation and dry adiabatic package as the full physics GSM, but the Kuo convection is disabled at most points through the use of high threshold values and various criteria that must be met before moist convection is allowed to take place. In our tests, we emulated this by using a dummy routine for Kuo convection. The second version is the original code of the full physics GSM, before any optimization. Finally, the third version is the optimized version of the full physics GSM. This version is identified by its SCCS sequence number as version 5.1. Thus, comparisons between the GWC and optimized timings will give an indication of the computational cost of the enhanced physics packages, while comparisons between the original and optimized code show the speedup due to optimization.

All timing tests were performed as history restarts from a previous 24-hour forecast of the GSM, to eliminate any spin-up that would distort the timing information. Each timing test consists of a three-hour forecast; since radiation is computed every three hours, timings can thus be extrapolated to multiples of this forecast length. All timing tests use the 18-layer structure used extensively by PL (e.g., Norquist and Yang, 1990). Table 1 contains the timings from our single-processor timing tests for the three model versions on a Cray 2, for a model resolution of rhomboidal 40 (R40) with 18 layers in the vertical. The differences in CPU time between these runs are summarized in table 2. All runs (except run 1c) were conducted in a dedicated mode, and CPU and elapsed (wall-clock) times are essentially identical.
Table 1: Cray 2 timings of single processor runs

<table>
<thead>
<tr>
<th>Run number</th>
<th>Model version</th>
<th>Compiler optimization</th>
<th>CPU time</th>
<th>elapsed time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>original</td>
<td>vectorized</td>
<td>253.5</td>
<td>257.2</td>
</tr>
<tr>
<td>1b</td>
<td>optimized</td>
<td>vectorized</td>
<td>133.9</td>
<td>134.7</td>
</tr>
<tr>
<td>1c</td>
<td>GWC</td>
<td>vectorized</td>
<td>91.2</td>
<td></td>
</tr>
<tr>
<td>2a</td>
<td>original</td>
<td>scalar</td>
<td>676.2</td>
<td>678.4</td>
</tr>
<tr>
<td>2b</td>
<td>optimized</td>
<td>scalar</td>
<td>525.4</td>
<td>527.3</td>
</tr>
</tbody>
</table>

Table 2: Cray 2 single processor run speedups

<table>
<thead>
<tr>
<th>Run numbers</th>
<th>Description</th>
<th>speedup</th>
</tr>
</thead>
<tbody>
<tr>
<td>1b - 1a</td>
<td>Effect of recoding for optimization</td>
<td>1.9</td>
</tr>
<tr>
<td>1c - 1b</td>
<td>Cost of advanced physics</td>
<td>1.5</td>
</tr>
<tr>
<td>1a - 2a</td>
<td>Vectorization of original code</td>
<td>2.7</td>
</tr>
<tr>
<td>1b - 2b</td>
<td>Vectorization of optimized code</td>
<td>3.9</td>
</tr>
</tbody>
</table>

It is evident from table 2 that there is about a two-fold speedup due to the optimization of the original code. Comparison of the timings for runs.
with vectorization turned on and off indicates that this is due mostly to the better vectorization speedup in the optimized code (3.9 vs. 2.7), although there is also a slight speedup in the scalar run. The computational cost of the enhanced physics is only a 50% increase in CPU time for the optimized code (recall that the GWC code in run 1c uses the optimized hydrodynamics code). In current operational practice at GWC, only 12 layers are used in the vertical; if this is taken into account, the computational cost is approximately 120%.

A further breakdown of timings for the original and optimized model code is shown in table 3. The timings shown are derived from a repeat of runs 1a and 1b, in which the flowtrace option was enabled. The flowtrace utility on the Cray 2 results in estimates of CPU time spent in each subroutine (along with the number of times it is called). The numbers in table 3 were obtained by adding the CPU times of the appropriate subroutines (in cases where the same subroutine was called from different parts of the calculation, the total times were divided among the tasks). A comparison of the original and optimized timings shows that the major speedup stems from the optimization of the physics packages (radiation and pbl), with further speedups due to the adiabatic nonlinear products computation (nlprod), and more efficient use of the FFTs. As was discussed earlier, the Kuo scheme, which is the most important part of the adjustment physics, was not optimized, and there is consequently little change in the CPU time between the original and optimized code. The Legendre transforms indicate a speedup in the case of the Gaussian quadrature, and a slowdown for the Legendre sums, resulting in a slight overall speedup. The time stepping computation was also greatly accelerated, but it was only a minor contributor even in the original code.

Considering just the results of the optimized code from table 3, one can identify further areas of potential improvement. Aside from the existing Kuo scheme, which will be replaced by a different algorithm, the spectral transforms should be examined for further speedups, since they now consume more than half of the CPU time. The all-fortran version of the FFTs should be replaced by whatever optimized routines exist for a particular platform, and ways of rearranging the Legendre transforms to
Table 3. Flowtrace CPU times from 3-hour single-processor forecast on the Cray 2, for the original and optimized model code.

<table>
<thead>
<tr>
<th>Calculation:</th>
<th>original</th>
<th>optimized</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>sec ( % of total)</td>
<td>sec ( % of total)</td>
</tr>
<tr>
<td>Total:</td>
<td>250.000 (100.0 %)</td>
<td>132.000 (100.0 %)</td>
</tr>
<tr>
<td>1. Transforms:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.1. FFTs:</td>
<td>101.313 (40.5 %)</td>
<td>79.719 (60.4 %)</td>
</tr>
<tr>
<td>1.2. Gaussian quadrature:</td>
<td>39.542 (15.8 %)</td>
<td>21.107 (16.0 %)</td>
</tr>
<tr>
<td>1.3. Legendre sums:</td>
<td>32.584 (13.0 %)</td>
<td>24.557 (18.6 %)</td>
</tr>
<tr>
<td>1.4. Pmns:</td>
<td>27.909 (11.1 %)</td>
<td>33.615 (25.5 %)</td>
</tr>
<tr>
<td>1.5. Pmns:</td>
<td>1.278 (0.5 %)</td>
<td>0.440 (0.3 %)</td>
</tr>
<tr>
<td>2. Gridpoint calculations:</td>
<td>118.601 (47.4 %)</td>
<td>48.572 (36.8 %)</td>
</tr>
<tr>
<td>2.1. Radiation:</td>
<td>20.611 (8.2 %)</td>
<td>4.669 (3.5 %)</td>
</tr>
<tr>
<td>2.2. PBL:</td>
<td>43.947 (17.6 %)</td>
<td>4.249 (3.2 %)</td>
</tr>
<tr>
<td>2.3. Adjustment physics:</td>
<td>29.503 (11.8 %)</td>
<td>27.678 (21.0 %)</td>
</tr>
<tr>
<td>2.4. adiabatic nlprod:</td>
<td>24.540 (9.8 %)</td>
<td>11.976 (9.1 %)</td>
</tr>
<tr>
<td>3. time stepping:</td>
<td>29.946 (12.0 %)</td>
<td>3.192 (2.4 %)</td>
</tr>
<tr>
<td>4. Miscellaneous:</td>
<td>0.140 (0.0 %)</td>
<td>0.517 (0.4 %)</td>
</tr>
</tbody>
</table>
take advantage of optimized linear algebra routines should be considered. Other minor speedups may be possible throughout the code, and some have been identified in comment cards in the optimized physics packages. The results from these further modifications are bound to be machine-specific, however, and are only cost effective once a particular platform has been selected for extended use of the code.

3.2 Multi-Processor Timings

We conducted a number of other timing tests, all designed to test how well the optimized code can take advantage of multiple processors, and how its performance scales with increased resolution. The timings are shown in table 4; again, all runs were conducted on a dedicated Cray 2, except the 8-processor run, which was done on a dedicated Cray Y-MP. The R40 runs were conducted with the code compiled for multiprocessing, with 1, 2, and 4 processors. Comparison of the 1-processor run with the corresponding run in table 1 indicates that the overhead for multiprocessing is negligible (less than a 1% increase in elapsed time). The 2-processor run is about 1.87 times faster than the single-processor run, which corresponds to an efficiency of 93%. Here efficiency is defined as the ratio of speedup to the number of processors - a linear speedup would correspond to a 100% efficiency. For 4 processors, the speedup is only 3.1 (an efficiency of 78%). However, for R80, the 4-processor efficiency is 86% (with a speedup of 3.4), and for R120, it is 90% (with a speedup of 3.6). The length of the time step was decreased for the higher resolution runs (we used 15 minutes for R40, 7.5 minutes for R80, and 5 minutes for R120). Finally, we reran the R80 timing tests on an 8-processor Cray Y-MP. The 8-processor run shows a speedup of 5.3 (an efficiency of 66%).
Table 4. Elapsed times for single processor (s.p.) and multiprocessor (m.p.) runs on the Cray 2 (C2) and Cray Y-MP (Y-MP). NCPUs is the number of processors of the m.p. run, speedup the ratio of s.p. to m.p. elapsed time, and efficiency the ratio of speedup to NCPUs.

<table>
<thead>
<tr>
<th>Resolution</th>
<th>Computer</th>
<th>NCPUs</th>
<th>s.p. time</th>
<th>m.p. time</th>
<th>Speed up</th>
<th>Efficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>R40</td>
<td>C2</td>
<td>2</td>
<td>135.84</td>
<td>72.77</td>
<td>1.9</td>
<td>93%</td>
</tr>
<tr>
<td>R40</td>
<td>C2</td>
<td>4</td>
<td>135.84</td>
<td>43.31</td>
<td>3.1</td>
<td>78%</td>
</tr>
<tr>
<td>R80</td>
<td>C2</td>
<td>4</td>
<td>1426.63</td>
<td>415.69</td>
<td>3.4</td>
<td>86%</td>
</tr>
<tr>
<td>R120</td>
<td>C2</td>
<td>4</td>
<td>6645.33</td>
<td>1852.70</td>
<td>3.6</td>
<td>90%</td>
</tr>
<tr>
<td>R80</td>
<td>Y-MP</td>
<td>8</td>
<td>727.10</td>
<td>138.00</td>
<td>5.3</td>
<td>66%</td>
</tr>
</tbody>
</table>

In interpreting the multiprocessing efficiencies, it is useful to consider the four sources of sub-linear speedup: incomplete parallelization of the computation, multiprocessing overhead, load imbalances among the different tasks, and memory contention problems. In the timings reported here, the overhead is negligible, as discussed above. The other sources of inefficiency are not easily isolated. Incomplete parallelization of the computation implies that there is a certain fraction of the code that is executed in serial, not parallel, mode. Load imbalances can occur if the computational work cannot be evenly divided among the processors. If all latitude tasks required the same amount of computations, the load imbalance could easily be estimated from the mismatch between the number of processors and number of latitude tasks. This kind of analysis was successfully performed for an adiabatic version of the GSM by Hoffman and Nehrkorn (1989). However, in the present case with full physics, the latitude tasks will require different amounts of CPU time, and load imbalances are not
easily determined without additional diagnostics. Memory contention problems can occur when two latitude tasks need to increment the same global accumulators. Again these could only be detected with the aid of special diagnostics. We have not implemented any diagnostics.

All the sources of inefficiency become more important as the number of processors increases relative to the problem size. Incomplete parallelization becomes more important as the number of processors is increased, because the parallel portion of the code requires a smaller fraction of the total wall-clock time. Load imbalances can become more severe, because as the size of the tasks assigned to the processors becomes smaller, the fractional differences between the loads increase. Memory contention becomes more likely as more and more processors try to access the same number of storage locations. This explains the clear trend in table 3 that for a given problem size, the efficiency decreases with increasing number of processors, and conversely for a given number of processors, the efficiency increases with increasing problem size.

We have indicated possible future enhancements in comments in the code, which might mitigate the memory contention problems. They would require calls to machine-specific parallel processing library routines, and should only be considered once it is determined that a particular platform will be available in dedicated mode for a large enough time to make it worth the extra effort.

3.3 Overall Speedups

To assess the overall effectiveness of our optimization effort, one has to combine the speedups due to improved single-processor execution speed with the speedups due to multiprocessing. Compared with the original code (at R40), the optimized, multiprocessed code exhibits a 5.9-fold speedup when run on 4-processor Cray 2. The computational cost of the enhanced physics evaluated in the context of the optimized code (a factor of 2.2 if one considers the different number of vertical levels) is more than made up by the multiprocessing speedup on even a 4-processor Cray 2 (a factor of 3.1). Thus, the goal of making the implementation of the enhanced physics package feasible was achieved by our optimization techniques.
Furthermore, since the R80 resolution 8-processor time on a Cray Y-MP is only 2% larger than the R40 single-processor time on the Cray 2, a doubling of the resolution would be possible at the same turn-around time if the computing platform were upgraded from a Cray 2 to a Cray Y-MP. The recently announced Cray C90 would present further opportunities for increases in resolution.

4. Forecast Results

We conducted several 4-day forecasts, using a FGGE 3b analysis for 12 UTC 12 January 1979 as the initial condition. This analysis was produced at ECMWF using a resolution of 1.875°. Boundary data sets were prepared using routines and databases supplied by PL. Forecasts were produced using the GSM with GWC physics, as described in Section 3, with a resolution of R40. The enhanced physics GSM (referred to as APGSM in the remainder of this section) was used for forecasts with a R40, R80, and R120 resolution. The size of the transform grid was determined, for each truncation, from the minimum requirements of exact transforms of quadratic terms, and, in the restrictions imposed by the FFTs (the number of longitudes must be a multiple of 2, 3, and 5). The resulting grid sizes are 128 longitudes by 102 latitudes for R40, 250 by 202 for R80, and 384 by 302 for R120. The vertical structure of the GSM was the same 18-layer structure for all forecasts, with sigma interfaces at 0, 0.05, 0.1, 0.15, 0.2, 0.25, 0.3, 0.35, 0.4, 0.45, 0.546, 0.642, 0.735, 0.820, 0.893, 0.948, 0.973, 0.990, and 1.0. The length of the time step was decreased in accordance with the increase in resolution (20 minutes was used for the R30 runs reported in I, and we used here 15 minutes for R40, 7.5 minutes for R80, and 5 minutes for R120). The other numerical constants that needed adjusting were the diffusion coefficients for the horizontal 4th order diffusion. At R30, a value of 6x1015 m^4/s was used in I. For the higher resolutions, we chose a value that would result in the same fractional damping per timestep of the highest wavenumber (approximately 6%). The corresponding values are 2.55x1015 m^4/s for R40, 3.23x1014 m^4/s for R80, and 9.60x1012 m^4/s for R120. The remaining numerical constants were left unchanged from one resolution to the next (for the timestepping, a time-filter coefficient of 0.04 and a timestep coefficient of 0.5, corresponding to a semi-implicit step, were used).
It should be noted that for a more realistic assessment of resolution on forecast skill, the physical parameterization would have to be retuned, since they all contain a number of adjustable parameters that have been tuned to the R30 resolution used most extensively by PL. In addition, the vertical structure should also be reconsidered, particularly in the case of the high resolution (R120) run. The results reported here should thus be considered a preliminary exploration of the effect of higher horizontal resolution.

Forecasts are verified against the FGGE 3b analyses in the following discussion. Global root mean square (rms) and mean (bias) errors for the GWC physics GSM are shown in Figure 1, at forecast lead times of 24, 48, 72, and 96 hours. Because we used the same 18-layer structure as in the APGSM forecasts, the drag-law scheme of the GWC GSM affected only a very thin bottom layer, leading to somewhat exaggerated forecast errors in the boundary layer (below sigma=0.9). These large low-level errors are apparent in all panels of Figure 1. In the free atmosphere, the most striking feature of the error structure is the large positive temperature bias which grows with time (up to 3 K by day 4), presumably due to the lack of radiative cooling of the atmosphere in this model. This is also reflected in a sizeable geopotential height bias. Temperature rms errors also increase with forecast lead time to 4.8 K throughout most of the troposphere by day 4. The geopotential height rms error is dominated by the bias, reaching a maximum value of 137 m by day 4 at the tropopause (disregarding values for the top level of the model). Errors of the rotational wind exhibit maxima at the surface, at the top of the model, and at the jet level. At the jet level, the maximum values grow from 7.9 m/s to 19.9 m/s from day 1 to 4. Errors of the divergent wind are not shown, because the quality of the verifying analysis is uncertain.
Fig. 1: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) for the GWC GSM forecast with R40 resolution. Errors are shown at day 1 (solid line), day 2 (dotted line), day 3 (dashed line), and day 4 (dash-dotted line).
Fig. 1: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) for the GWC GSM forecast with R40 resolution. Errors are shown at day 1 (solid line), day 2 (dotted line), day 3 (dashed line), and day 4 (dash-dotted line).
Fig. 1: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) for the GWC GSM forecast with R40 resolution. Errors are shown at day 1 (solid line), day 2 (dotted line), day 3 (dashed line), and day 4 (dash-dotted line).
The corresponding figure for the APGSM R40 forecast (Figure 2) reveals sharp decreases in boundary layer errors, and the elimination of the positive temperature bias (which is now slightly negative in the upper troposphere, reaching -39 m after 4 days). The rms errors of temperature and height are decreased as well, to less than 4 K throughout most of the troposphere by day 4 in the case of temperature, and less than 80 m in the case of height. The rotational wind errors are smaller throughout the atmosphere, with jet level wind errors between 7.8 m/s (at day 1) and 16.5 m/s (at day 4).

For a direct comparison of day 4 forecast errors, the curves for the GWC GSM and the APGSM R40 forecasts are plotted together in Figure 3. Also shown in Figure 3 are the error curves for the APGSM R80 and R120 forecasts, allowing an assessment of physics and resolution changes. Errors of all variables are most affected by the change in physics packages, and only to a lesser degree by increases in resolution. The most obvious effect of the higher resolution is a reduction of the negative temperature and height bias in the upper troposphere. Rms errors of these quantities are also reduced overall by increases in resolution, but not consistently at all levels. Beneficial effects of increased resolution are not obvious for the rotational wind, which shows errors that are essentially identical for all three truncations.

It is not uncommon that increases in forecast skill, as measured by the kind of global statistics presented here, are hard to detect when model resolution is increased (Simmons, 1991, personal communication). The primary reason for this is that while low resolution forecasts will be missing small-scale features present in the verifying analyses (and the atmosphere), high resolution model forecasts may not predict them at the proper location (or time). In that kind of scenario, the high resolution forecast would be penalized more severely in terms of rms statistics. In addition, the verifying analysis used here is not high resolution, so small scales present in the R120, and, to some extent, the R80 forecasts would appear as errors. It is thus not surprising that in the tests reported here, where the high resolution model was also not properly tuned, the results are somewhat ambiguous.
Fig. 2: Global mean (a, c) and rms (b, d, e) error of temperature (a, b),
geopotential height (c, d), and rotational wind (c) for the APGSM GSM
forecast with R40 resolution. Errors are shown at day 1 (solid line), day
2 (dotted line), day 3 (dashed line), and day 4 (dash-dotted line).
Fig. 2: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) for the APGSM GSM forecast with R40 resolution. Errors are shown at day 1 (solid line), day 2 (dotted line), day 3 (dashed line), and day 4 (dash-dotted line).
Fig. 2: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) for the APGSM GSM forecast with R40 resolution. Errors are shown at day 1 (solid line), day 2 (dotted line), day 3 (dashed line), and day 4 (dash-dotted line).
Fig. 3: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) at day 4. Errors are shown for the GWC GSM forecast with R40 resolution (solid line), and the APGSM forecasts with R40 (dotted line), R80 (dashed line), and R120 (dash-dotted line) resolution.
Fig. 3: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) at day 4. Errors are shown for the GWC GSM forecast with R40 resolution (solid line), and the APGSM forecasts with R40 (dotted line), R80 (dashed line), and R120 (dash-dotted line) resolution.
Fig. 3: Global mean (a, c) and rms (b, d, e) error of temperature (a, b), geopotential height (c, d), and rotational wind (e) at day 4. Errors are shown for the GWC GSM forecast with R40 resolution (solid line), and the APGSM forecasts with R40 (dotted line), R80 (dashed line), and R120 (dash-dotted line) resolution.
5. Minimization of pressure gradient force errors

5.1 Pressure-gradient force errors

The low accuracy of geopotential calculations in spectral numerical weather prediction models introduces substantial errors, and therefore affects the overall forecast error. The order of magnitude of the errors commonly reaches 100 m at 500 mb level (in non-dissipative systems, Phillips, 1974), and generates erroneous systematic cooling of the atmosphere above the high mountains regions. This error is the most noticeable in conjunction with the calculations of the Pressure-Gradient Force (PGF)

\[
- \nabla_\sigma \Phi - RT \nabla ln p_s
\]  

(5.1.1)

which involves horizontal differencing of the geopotential, \( \Phi \) (\( p_s \) is the surface pressure, \( R \) is universal gas constant, \( T \) is temperature and \( \sigma \) is the normalized pressure coordinate). Over sloping terrain the two terms of (5.1.1) tend to be large in absolute values and to have opposite signs. The cancellation, however, is difficult to achieve in the models, and a 1% error in temperature (2-3° C) will result in a 10% error in the pressure gradient force (Sundqvist, 1975). An example of the PGF existing at the model level is given in Figure 4 (a) (see Section 5.3 for data description). This force is produced by the spectrally represented orography as demonstrated by the figure. The PGF is clearly shown over the high mountain regions; Himalayas, Andes and Greenland (max vector intensity plotted is \( 10^{-2} \text{ m/s}^2 \)). With an increase of the model spectral resolution, this PGF decreases as the mountains are represented better.

Together with the truncation error, the erroneous estimates of geopotential, \( \Phi \), are a persistent source of spurious PGF in the spectral models (Gary, 1973). The low accuracy of geopotential calculations is mainly caused by the vertical differencing scheme used to compute \( \Phi \) from the model hydrostatic equation. If we integrate the hydrostatic equation in
the $\sigma$-coordinate system

$$\Phi(\sigma) = \Phi(1) - \int_1^\sigma RTd(ln\sigma)$$ (5.1.2)

we see that the geopotential at any $\sigma$-level is only a function of temperature at and below that level. Therefore, the derivative $\partial \Phi / \partial (ln\sigma)$ should not be approximated by a higher order finite-difference scheme in order to preserve the continuity of the hydrostatic equation (Janjic, 1977). As a result, the geopotential is calculated with a low accuracy that leads to erroneous estimates of (5.1.1). These inadequacies have been investigated in part by Brenner (1988), who showed that all commonly used schemes produced substantial height errors at the upper model levels. An example of the errors produced by the current model vertical differencing scheme (equation (15) in Brenner et al., 1982) is illustrated by Figure 4 (b). It shows the difference of the PGF intensity, at the model level 2, between the analytically determined and numerically determined $\Phi$. In both cases the same spectrally represented orography and surface pressure are used. Again, the additional PGF, caused by the model vertical differencing scheme, exists in the high mountain regions. Note that, in this particular case, the difference of the PGF intensity is of order $1\% \times (10^{-3} \text{ m/s}^2)$ of the PGF in diagram (a).

5.2 Error minimization

These errors can be reduced if the standard stratification approximation is used, i.e., if the thermodynamic model variable is not the temperature $T$, but the deviation of temperature $T'$, from some reference temperature $T_r$

$$T' = T - T_r$$ (5.2.1)

The reference temperature, $T_r$, can be defined to be a function that removes most of the cancellation in the calculations of term (5.1.1). It is determined
Figure 4. (a) Spurious pressure gradient force for a standard atmosphere due to spectral representation of mountains at R20 truncation. (b) Difference in spurious pressure gradient force at model level 2, due to model vertical differencing scheme for the same spectral truncation (R20).
by the stratification factor $C_0$ defined as

$$C_0^2 = \frac{R^2 T_r}{g} \left( \frac{g}{c_p} + \frac{dT_r}{dz} \right) \tag{5.2.2}$$

where, $g$ is the gravity constant, $c_p$ is the specific heat at constant pressure and $z$ is the vertical coordinate. This factor can be chosen to be

$$C_0 = \begin{cases} \text{const.} \\ f(p) \\ f(p, \phi) \end{cases}$$

i.e., constant, function of pressure only, or function of pressure and latitude $\varphi$ (Zhang et al., 1990). The model development at the ECMWF (Simmons and Chen, 1991) is based on defining the stratification factor as a function of vertical coordinate only, i.e., pressure or $\sigma$. The work of Zhang et al. includes a more general case of a reference atmosphere varying with latitude.

Two possible approaches in applying this approximation to the model have been investigated at the ECMWF. The first approach (called a TR scheme; Simmons and Chen [1991]) is based on the assumption that the reference atmosphere is removed from the spectrally represented thermodynamic variable. In this approach a substantial modification of the model code is required since the spectral variable is changed from the temperature itself to the perturbation of temperature, $T'$. A second approach, that requires less extensive modifications (identified as the PG-scheme), is based on using the standard stratification approximation in the model calculations that involve the PGF calculations only.
5.2.1 Schemes

In the present study we have tested three schemes: the PG-scheme originally used in the ECMWF study, a modified version of the PG-scheme called MPG-scheme, and the null scheme which consists of an isothermal reference profile. Table 5 summarizes the differences in parameter values for the three schemes.

Table 5. Parameter values used in the three schemes: the PG, the MPG and null (an isothermal case).

<table>
<thead>
<tr>
<th>Scheme</th>
<th>$T_0$ [°K]</th>
<th>$p_0$ [kPa]</th>
<th>$\Lambda$ [°K/km]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PG</td>
<td>288</td>
<td>1013.2</td>
<td>6.5</td>
</tr>
<tr>
<td>MPG</td>
<td>270</td>
<td>1013.2</td>
<td>2.5</td>
</tr>
<tr>
<td>null</td>
<td>300</td>
<td>1013.2</td>
<td>0.0</td>
</tr>
</tbody>
</table>

5.2.1.1 The PG-scheme

This scheme is based on a reference atmosphere defined as

$$T_r(p) = T_0 \left( \frac{p}{p_0} \right)^\alpha$$  \hspace{1cm} (5.2.3)

where $T_0$ and $p_0$ are the surface values of the temperature and pressure respectively, $\alpha=\Delta R/g$ and $\Lambda$ is the lapse rate. This reference atmosphere is subtracted from the model virtual temperature for the purpose of calculating the geopotential (see Section 5.2.2 for a description of the code modification).
We used this scheme in the present study since the idealized tests of the ECMWF model showed that the PG-scheme accounted for an error reduction similar to the reduction obtained by the more extensive TR scheme. The parameters were chosen to be the same as those reported in Simmons and Chen (1991) (the first row of table 5). In Figure 5, diagram (a) illustrates this reference temperature in respect to the standard atmospheric profile described in Section 5.3.

5.2.1.2 The MPG -scheme

We also considered, in the present study, a modified PG-scheme (the MPG -scheme) which is based on the same reference profile (5.2.3) but uses different parameter values (the second row of table 5). As Figure 5 (b) shows, this profile resembles a linear approximation of the standard temperature profile with an unrealistic lapse rate. The motivation for considering this modification was to estimate what impact the variation of reference atmosphere would have on the present model error. Further modifications of this scheme are not presented here since the improvements that we found were insignificant (see Section 5.4).

5.2.1.3 The null scheme - an isothermal case

Figure 5 (c) illustrates the case in which the reference atmosphere is isothermal. The value of the surface temperature $T_0$ was chosen to be 300°K which is the same as the reference temperature used in the model semi-implicit time integration scheme. This profile was used as a reference for an earlier version of the model when the thermodynamic variable was a perturbation from a uniform temperature profile.

5.2.2 Code modification

The implementation of the PG - scheme (5.2.3) requires a minimal modification of the code (Simmons and Chang, 1990). It involves only the equations that include PGF terms and, if applied to the grid-point thermodynamic variable, it is localized to a very few subroutines of the multi - tasking model code.
Figure 5. Reference temperature profiles compared to the temperature profile of the standard atmosphere, used in: (a) the PG scheme, (b) the MPG scheme and (c) the isothermal case.
5.2.2.1 Model equations

The model equation that directly involves PGF calculations is the divergence equation (Kaplan et al., 1985)

\[
\frac{\partial D}{\partial t} = -\frac{1}{a \cos^2 \phi} \left[ \frac{\partial B}{\partial \lambda} - \cos \phi \frac{\partial A}{\partial \phi} \right] - \nabla^2 [E + \Phi] + \nabla \cdot F_h 
\]

(5.2.4)

where the notation is identical to Brenner et al. (1982). When the temperature variable, T, is replaced by the expression (5.2.1), the equation retains the same form for the temperature perturbation T', with additional terms due to a reference atmosphere

\[
- \left[ \frac{1}{a^2 \cos^2 \phi} \frac{\partial}{\partial \lambda} \left( R_T \frac{\partial \ln p_*}{\partial \lambda} \right) + \frac{1}{a^2 \cos \phi} \frac{\partial}{\partial \phi} \left( R_T \cos \phi \frac{\partial \ln p_*}{\partial \phi} \right) \right] + \nabla^2 \int c_p \kappa T_r d(ln \sigma). \quad (5.2.5)
\]

When compared to (5.1.1) we see that the later expression represents the PGF due to a reference atmosphere, T_r, in the model coordinate system. Furthermore, if T_r is replaced by the expression (5.2.3), the above expression reduces to one term

\[
- \nabla^2 \frac{R T_0}{\alpha} \left( \frac{p_*}{p_0} \right)^\alpha. \quad (5.2.6)
\]

This term can be easily combined with the geopotential calculations for the perturbation temperature under the Laplacian operator in equation (5.2.4). Therefore, the major cancellation of the PGF, due to a reference atmosphere, can be removed from the calculations in the divergence equation if the geopotential calculations in the hydrostatic equation are modified by the term (5.2.6). Indeed, this has been done by Simmons and Chen (1990), who modified the surface geopotential (equation
15) in the ECMWF hydrostatic equation by the term identical to the expression under the Laplacian operator in (5.2.6).

5.2.2.2 Code changes

For the experiments in this part of the study we used the dry-adiabatic version of the code. The present code structure enabled a relatively easy implementation of the changes described above. The changes were made in a grid-point space, which eliminated the need for additional corrections of the rest of the model equations and the semi-implicit time integration scheme. (In the case of implementing the schemes in the spectral space, it would be necessary to make quite radical model code changes.)

The first change that was made was to calculate the reference temperature in grid-point space. This was done in the subroutine NLPROD, that estimates nonlinear terms, where grid-point values of the surface pressure, \( p^* \), are available. The reference temperature was then subtracted from the temperature field and added back to the field after the evaluation of the nonlinear terms.

The second and most significant change was to integrate the model hydrostatic equation in the grid-point space. (The current model formulation involves hydrostatic integrations in the spectral space.) Therefore, the subroutine consisting of the hydrostatic equation has been recoded and called before the nonlinear term evaluation. During this process, the surface geopotential was also modified by the corresponding portion of the term (5.2.6).

The third type of modification mostly involved the changes related to the postprocessing of model integrations. In particular, for each model run we have created evolution files for model tendencies, we have developed routines for calculations of the pressure gradient force, maximal grid-point tendencies and spectral modes that have the maximal amplitude at each model level (this enabled us to check if any of the schemes exhibit spectral biases). Finally, we have modified the postprocessing package to plot the
forecast fields at the model σ-levels instead of the mandatory pressure levels. The latter was needed in order to eliminate further contribution of the hydrostatic equation to the temperature (and geopotential) errors in the post-processing cycle.

5.3 Initial Data

Most of the results presented here are from idealized tests. As in Simmons and Chen, idealized tests were based on initial data describing a dry atmosphere at rest. They consisted of an analytically prescribed temperature profile which is a function of pressure alone, and which is of a form such that the balanced surface pressure could be computed analytically given the basic model orography, $\Phi_s$. In our study, we used the analytical profiles defined by Phillips (1974)

$$T\left(\ln \frac{P}{P_0}\right) = \frac{1110}{R} \left\{72.43 - \ln \frac{P}{P_0} \left[2(-6.9) - 3\ln \frac{P}{P_0}\right]\right\}$$

(5.3.1)

An example of this temperature profile is shown in Figure 5. This temperature profile has been extensively used in numerical studies related to spurious PGF. The balanced surface pressure, $p_s$, was computed from

$$\Phi_s\left(\ln \frac{p_s}{P_0}\right) = 1110\left\{0.95 - \ln \frac{p_s}{P_0} \left[72.43 - \ln \frac{p_s}{P_0} \left(-6.9 - \ln \frac{p_s}{P_0}\right)\right]\right\}$$

(5.3.2)

To calculate the surface pressure we used the surface geopotential from the gridded FGGE 3b data set. The orography data was smoothed horizontally first, and then truncated spectrally at the needed resolution.
5.4 Tests

The idealized tests were created with the purpose of investigating how well the PGF-scheme minimizes present model error. Simmons and Chang found that the errors were significantly reduced, especially for longer time integrations, at truncation T43. The degree of forecast improvement was such that the accuracy of T43 model runs became equivalent to the unmodified T106 model runs. These are significant corrections that can affect the cost of model integrations substantially. The purpose of this study is to determine if the similar improvements of the forecasts can be achieved by the present model at R40 truncation and compared to the forecasts with R80 truncation.

Before starting the long, ten-days model runs at R40 and R80 truncations, we tested the minimization schemes at a lower spectral truncation, R20 and R40. This was based on the fact that the PGF errors are so tightly related to the spectral truncation error (as described in Section 5.1.) that the effects of the schemes should be noticeable even at the lower spectral resolution. In general, we produced four types of model runs. The Control runs were based on an unmodified model code and used as a reference for the "PG" and the "MPG" runs, which utilized the PG and the MPG schemes, respectively. Finally, "Isothermal" runs were based on an isothermal reference atmosphere. We also tested the schemes' performance in respect to distribution of the model vertical levels (equally-spaced levels vs. current vertical structure) and the model diffusion. The results are presented in terms of the model initial tendencies and the one-day and ten-days forecasts.

We limited the investigation to the cases described above, and we reviewed the results of the idealized runs only. We did not find the substantial improvements of the forecasts by the minimization schemes considered, although the model vorticity dynamics showed some sensitivity to the presence/absence of the reference atmosphere.
5.4.1 Scheme comparison

The performance of the three schemes are best illustrated by the initial tendencies presented in Figure 6. Two diagrams on the left-hand side show max grid-point divergence tendency at R20 (upper panel) and R40 (lower panel) model truncation. It is noticeable that the PG and MPG schemes do not differ very much from the unmodified code, while the "isothermal" scheme produces the largest initial tendencies. Two panels on the right-hand side of the figure show the corresponding vorticity tendencies. Although the differences between the schemes are not large, some sensitivity of the vorticity is noticeable. In particular, for the R20 case the "isothermal" scheme produces the smallest tendencies, while in the R40 case (the lower panel), the PG-scheme produces the smallest tendencies. Also, in the case of the PG-scheme, it appears that the maximum tendency decreases slightly with height (with exception of the model level 15) in R20 truncation, but increases slightly with height (except at the model level 13) in R40 truncation.

Since the differences in Figure 6 appear to be very small, we considered the most sensitive, the velocity fields, for the forecast comparison. Figure 7 shows the differences of the global RMS of the velocity field. Again, the upper two panels represent R20 truncation, and the lower two panels represent R40 truncation. The differences between the forecasts are of order of 0.1 cm/s. In the case of the divergent velocity component (two panels on the left-hand side of the figure) the PG and the MPG produce larger values than the control (solid line patterns) at R20 truncation. At R40 truncation, both of the schemes produce larger values at the upper model levels (except the PG-scheme at the level 3), and slightly smaller values at the lower model levels. The rotational wind component (two panels on the right-hand side of the figure) shows that both of the schemes produce larger values at all model levels except the uppermost levels (5-1 and 3-1, respectively).

The geographical distribution of the winds at level $\sigma = 0.124$ for R20 forecasts is shown in the Figure 8. Diagram (a) shows a distribution of the wind speed obtained in the Control case after 10 days of integrations. The
Figure 6. Vertical distribution of maximum initial tendencies for two model truncations: R20 (upper panels) and R40 (lower panels). Divergence tendencies are shown on the left-hand side and vorticity on the right-hand side.
**Figure 7.** Vertical distribution of the velocity RMS. The upper two panels apply to R20 truncation, and the lower two panels apply to R40 truncation. Differences between the PG, MPG and Control runs are plotted. Two panels on the left-hand side are derived for the divergent component of the velocity, while panels on the right-hand side are derived for the rotational part of the velocity.
Figure 8. Geographical distribution of the isotachs of horizontal wind for R20 model truncation. Diagram (a) shows the wind speed after 10-days integration in the Control case. Diagram (b) shows the differences between wind forecasts obtained with the PG-scheme and the Control case.
wind maxima are noticeable over the Himalayas, Greenland and the Antartica. The lower diagram (b) shows the difference in wind speed between the Control forecast and the PG-scheme (the PG-scheme produced smaller wind speeds).

Figure 9 shows the winds for R40 truncation at $\sigma = 0.594$. Diagram (a) shows the horizontal wind field after 1-day of integrations in the Control case (maximum wind does not exceed 5m/s). The rotational component of the wind is pronounced over the mountains. Again, diagram (b) shows the difference in the wind speed between the Control case and the PG-scheme. It is noticeable from the last two figures that the differences are very small and localized above the mountains.

Comparison of the global mean values of the differences between forecasted and initial temperature and geopotential fields corresponding to Figure 7 is shown in Figure 10. In general, the 10-days forecasts at R20 produced slightly increased temperature (of the order of $10^{-2}$) at the upper model levels ($\sigma = 0.497$ to $\sigma = 0.021$) and decreased temperature at the lower levels ($\sigma = 0.995$ to $\sigma = 0.594$). In these forecasts, the schemes tended to produce smaller increase at the upper levels, and almost no difference in the forecasts at the lower levels (diagrams on the left of Figure 10). This is reflected in difference between the geopotential forecasts to be of the order of $10^{-2}$ m. Similar behavior was obtained at R40 truncation for one day forecasts (two lower panels).

The comparison of the vertical distribution of the spectral modes, that have the largest amplitudes, did not show any particular bias of the schemes. This is illustrated by table 6.
Figure 9. Geographical distribution of the horizontal wind for R40 model truncation. Diagram (a) shows the winds at model level 11 after 1-day integration in the Control case. Diagram (b) shows the difference between the wind speed forecasts obtained with the PG-scheme and unmodified code.
Figure 10. Vertical distribution of global mean values of the differences between initial data and forecasts. Two panels on the left-hand side are derived for the temperature, while panels on the right hand-side are derived for the geopotential. The upper two panels apply to 10-days forecast R20 truncation, and the lower two panels apply to 1-day forecasts at R40 truncation. Differences between the PG, MPG and Control runs are plotted. (Note difference between the horizontal axes of the upper and the lower panels).
### Table 6. Vertical distribution of the vorticity spectral wave number (m,n) that has max amplitude after one day of integrations.

<table>
<thead>
<tr>
<th>Model truncation</th>
<th>R20</th>
<th>R40</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scheme</td>
<td>Control</td>
<td>PG</td>
</tr>
<tr>
<td>σ</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.021</td>
<td>(0,12)</td>
<td>(0,12)</td>
</tr>
<tr>
<td>0.074</td>
<td>(1,16)</td>
<td>(1,16)</td>
</tr>
<tr>
<td>0.124</td>
<td>(3,21)</td>
<td>(3,21)</td>
</tr>
<tr>
<td>0.174</td>
<td>(1,14)</td>
<td>(1,14)</td>
</tr>
<tr>
<td>0.225</td>
<td>(1,20)</td>
<td>(1,20)</td>
</tr>
<tr>
<td>0.275</td>
<td>(1,20)</td>
<td>(1,20)</td>
</tr>
<tr>
<td>0.325</td>
<td>(7,25)</td>
<td>(7,25)</td>
</tr>
<tr>
<td>0.375</td>
<td>(7,25)</td>
<td>(7,25)</td>
</tr>
<tr>
<td>0.425</td>
<td>(8,25)</td>
<td>(8,25)</td>
</tr>
<tr>
<td>0.497</td>
<td>(8,25)</td>
<td>(8,25)</td>
</tr>
<tr>
<td>0.594</td>
<td>(0,5)</td>
<td>(0,5)</td>
</tr>
<tr>
<td>0.688</td>
<td>(3,17)</td>
<td>(3,17)</td>
</tr>
<tr>
<td>0.777</td>
<td>(3,17)</td>
<td>(3,17)</td>
</tr>
<tr>
<td>0.856</td>
<td>(10,28)</td>
<td>(10,28)</td>
</tr>
<tr>
<td>0.920</td>
<td>(10,28)</td>
<td>(10,28)</td>
</tr>
<tr>
<td>0.960</td>
<td>(10,28)</td>
<td>(10,28)</td>
</tr>
<tr>
<td>0.981</td>
<td>(10,28)</td>
<td>(10,28)</td>
</tr>
<tr>
<td>0.995</td>
<td>(10,28)</td>
<td>(10,28)</td>
</tr>
</tbody>
</table>

#### 5.4.2 Vertical distribution of the model levels

As indicated by the results reviewed in the previous section, some variability of the schemes with the height exists. To examine the impact of the model level distribution on error minimization schemes, we repeated the Control, the PG and the MPG runs with equally spaced model levels at R20 truncation. The results are shown in Figure 11.
Figure 11. Vertical distribution of maximum initial tendencies for the model truncation R20, and different distribution of the model levels. The results shown on the left-hand side are obtained with the current distribution, and the results shown on the right-hand side are obtained with equally-spaced model levels.
Two panels on the left-hand side of the figure are equivalent to the upper two panels of Figure 6. They represent the differences between the maximum grid-point initial tendencies obtained with the current distribution of model vertical levels. The other two panels, on the right-hand side of the figure, are corresponding tendencies that were obtained with the equally-spaced model levels. It is noticeable, from the figure, that the performance of the minimization schemes is not affected by the level distribution in respect to the control runs. Slight differences are noticeable in the vorticity field (the lower two panels), but not of a significant value.

5.4.3 The model diffusion

Since the differences reviewed in earlier two sections were small, we compared them with the amount of horizontal diffusion applied to the Laplacian term in the equation (5.2.4) that involves geopotential estimates. The results of this comparison are given in Figure 12.

An evolution of total divergence and vorticity tendencies during 1-day forecasts at R20 truncation are plotted on the diagrams. Diagrams on the left-hand side of the figure represent the Control and the PG runs without diffusion, while the diagrams on the right-hand side represent the corresponding runs with applied horizontal diffusion. The upper two diagrams represent the divergence tendencies, and the lower two diagrams represent the vorticity tendencies. It is quite noticeable that the scheme follows the Control run quite closely independently on the model horizontal diffusion. Differences appear to be slightly larger for the divergence tendencies than for the vorticity tendencies. The horizontal axis represents the number of time steps 30 min long.
Figure 12. Evolution diagrams of total divergence tendencies (the upper diagrams) and total vorticity tendencies (the lower diagrams) from the runs with and without horizontal diffusion. Model truncation is R20, and the length of forecasts is one day.
5.5 Conclusions

In the present study we investigated schemes for minimization of the PGF errors in the current version of the global spectral model. We tested three schemes, named the PG-scheme, the MPG-scheme and the Isothermal case against the Control case that consisted of the unmodified model code. The forecasts were made at the R20 and R40 model truncation and were of durations varying from 1-hour to 1-day to 10 days. The initial data described a dry standard atmosphere that justified use of dry adiabatic model code without any physics.

We tested the error minimization against the various scheme parameters (the temperature and the lapse rate), the model spectral resolution (R20 and R40), the distribution of the model vertical levels (current vs. equally-spaced levels) and the model horizontal diffusion.

The idealized tests showed that no substantial improvements of the forecasts were achieved by the PG-scheme. Some sensitivity was found in the fields related to the model vorticity dynamics (the rotational velocity and the vorticity). The changes found did not exceed 1% of the Control forecasts obtained by the current model code. We did not find this justifiable for extending the study to the integrations at higher spectral resolution (i.e., R80). The results obtained at the lower resolution are qualitatively in agreement with the results of idealized tests of Simmons and Chen (1990). The only difference is that the ECMWF orography is more realistic and spectrally fitted to T106, while the orography available for the runs with the current model version has been smoothed by the low spectral truncation. This resulted in smaller initial tendencies than the ones obtained by the ECMWF.

Although substantial improvements of the forecasts by the PG-scheme were not found, the integrations with the real data and more realistic orography may provide improved forecasts. Simmons and Chen have found improvements in real data forecasts which were described later by specific vorticity dynamics of the ECMWF model (personal communication with Simmons). In our idealized tests, it was found that the additional
PGF force has been created by the vertical differencing scheme at the same geographical positions where the ECMWF forecast improvements were located. Together with the sensitivity of the rotational wind and the vorticity field, which we also diagnosed, this can be indicative of possible forecast improvements with the real data and orography, similar to the ones obtained in the ECMWF study.
6. Summary

The development of a vectorized, multiprocessing global spectral model (GSM) with enhanced physical parameterizations is documented. The starting point was the Phillips Laboratory GSM, in its GL90 version, containing an enhanced suite of physical parameterizations. The latitude tasking scheme for multiprocessing the loop over latitude in the calculation of the spectral tendencies and adjusted model variables was implemented, using the general truncation version of the hydrodynamics code. Wavenumber calculations were vectorized over wavenumber and multiprocessed over vertical level. All gridpoint calculations were vectorized over longitude, and the physics packages were brought into closer compliance with plug-compatibility rules.

Speedups due to the optimization were demonstrated in single- and multiprocessing timing tests on a dedicated Cray 2 and Cray Y-MP. Compared with the original code (at R40), the optimized code runs 1.9 times faster on a single processor Cray 2. When run on 4-processor Cray 2, the optimized code exhibits a 5.9-fold speedup compared to the original version. The computational cost of the enhanced physics evaluated in the context of the optimized code (a factor of 2.2 if one considers the different number of vertical levels) is more than made up by the multiprocessing speedup on even a 4-processor Cray 2 (a factor of 3.1). Thus, the goal of making the implementation of the enhanced physics package feasible was achieved by our optimization techniques.

Furthermore, since the R80 resolution 8-processor time on a Cray Y-MP is only 2% larger than the R40 single-processor time on the Cray 2, a doubling of the resolution would be possible at the same turn-around time if the computing platform were upgraded from a Cray 2 to a Cray Y-MP. The recently announced Cray C90 would present further opportunities for increases in resolution.

The advanced physics GSM (APGSM) was evaluated in forecast tests, and contrasted with the simpler physics GSM used at GWC. Use of the APGSM leads to sharp decreases in boundary layer forecast errors, and the
elimination of the positive temperature bias of the GWC GSM (which is now slightly negative in the upper troposphere, reaching -39 m after 4 days). The rms errors of temperature and height are decreased as well, and the rotational wind errors are smaller throughout the atmosphere. Forecast errors of all variables are most affected by the change in physics packages, and only to a lesser degree by increases in resolution. The most obvious effect of the higher resolution is a reduction of the negative temperature and height bias in the upper troposphere. Rms errors of these quantities are also reduced overall by increases in resolution, but not consistently at all levels. Beneficial effects of increased resolution are not obvious for the rotational wind, which shows errors that are essentially identical for all three truncations.

Minimization of errors in the computation of the horizontal pressure gradient force was investigated, using a perturbation temperature instead of full temperature in the integration of the hydrostatic equation. Several schemes were tested in an idealized model atmosphere. The idealized tests showed no substantial improvements of the forecasts were achieved by any of the schemes we tested. Some sensitivity was found in the fields related to the model vorticity dynamics (the rotational velocity and the vorticity). However, the changes found did not exceed 1% of the control forecasts obtained by the current model code.

7. References


Appendix A: User's Guide

The document reproduced on the following pages is included as a plain text file (named "users.guide") in the software deliverables of this contract.
Users's Guide to the Global Spectral Model (GSM)

1. Overview

This document describes the Global Spectral Model (GSM), in its general truncation version, as configured for the GL (new) physics with vectorization and multiprocessing. A technical report (GL-TR-90-0309, NTIS ADA232123) describes the algorithms and software design of this version. For a scientific documentation, refer to the GL technical reports, and the references contained therein (hydrodynamics: AFGL-TR-82-0393, NTIS ADA129283; AFGL-TR-84-0308, NTIS ADA160370; PBL: AFGL-TR-84-0063, NTIS ADA144224; AFGL-TR-87-0246, NTIS ADA199440; PI-TR-91-2031, NTIS ADA235310; radiation: AFGL-TR-84-0217, NTIS ADA148015; AFGL-TR-88-0018, NTIS ADA193369; convection: AFGL-TR-85-0160, NTIS ADA170137; GL-TR-90-0285, NTIS ADA241684). This code employs what is known as the "GL90" physics with the following changes. The 3-decker radiation scheme is used in version 5.1; version 5.2 uses an improved radiation scheme ("rad5") along with the tuned Slingo parameterization of stratiform and convective cloud cover (Slingo, 1987: QJRMS, 113, 889-927). This document applies to both version 5.1 and version 5.2; differences between the two versions exist in the boundary data set format, and are noted in the appropriate section. Minor changes and bug fixes have been made to pbl and Kuo schemes. In the following, the installation and running of the GSM are discussed.

2. Portability

The code itself is strictly FORTRAN 77 (ANSI X3.9-1978), with the following exceptions:

- INCLUDE statements are used throughout the code to include common blocks and parameter statements; the syntax of the statements (and filenames) may differ from one machine to the next, but (almost) all machines support this Fortran extension

- Multiprocessing (refer to C<mp> comment lines) is implemented using CRAY autotasking directives (CMIC$ comment lines). These are comments in standard fortran. In addition the Cray extension task common is used to identify private common blocks. In a standard single processor version of the code these would be ordinary common blocks.

- Module tcheck.f uses machine-dependent system routines for obtaining the time and date, and CPU time

- Some of the modules contain edit symbols that need to be replaced before compilation. (Edit symbols are strings enclosed by <> brackets.)

Along with the code several files are provided for compiling, linking, and running the code. These are developed for a UNICOS operating system, and are easily adapted to different machines using UNIX. They are:

A-1
makefile - used by the UNIX make facility for compilation and linking of the gsm (executable stored in gsm.x) and the utility program asize (stored in asize.x). The replacement of the edit symbols is included in the compilation rule where appropriate - the UNIX sed facility is used for this purpose.

Changes needed for porting: the directory pathnames, and the compiler statements and flags.

gsm.sed - sed script used by the makefile.

Changes needed for porting: for 32-bit machines, edit symbols <DOUBLE>, <DP> should be set for double precision, for 64-bit machines they should be set for single precision.

rungsm - c-shell script for running the gsm

Changes needed for porting: directory pathnames

3. Configuration

3.1. Arrangement of code

The source code for the GSM is stored in a number of different files. The fortran code is stored in files with extension "f". Modules containing edit symbols are stored in files with extension "fed"; edited versions of these modules (with the edit symbols replaced by running the sed script gsm.sed) are stored in files with extension "f". The fortran modules contain include statements for the inclusion of common and parameter statements to facilitate consistent use of commons and easy changes to array dimensions. Include blocks are stored in files with extension "blk". Include blocks containing edit symbols are stored in files with extension "bed"; after replacement of edit symbols, the edited include blocks are stored in files with extension "blk". A complete list of the file names is appended to this document. The makefile also contains the current list of source and object file names.

Aside from the GSM code proper, the FFTs are loaded in separately as a library. Since these may be more readily changed from one installation to the next, they are maintained in a separate directory (and with a separate makefile). They do not contain edit symbols, and do not use any of the GSM include blocks.

Edit symbols in the code (strings enclosed by <> ) are used to allow two options that cannot be accommodated with standard fortran:

- the model can be run in a hemispheric or a global version; this requires use of the edit symbols "<GLO>" and "<HEM>" , which serve to activate lines of code. For the global version, "<GLO>" is replaced by five blank spaces, and "<HEM>" by "C   ", and vice versa for the hemispheric version.
certain sensitive calculations can be performed in single or double precision. Generally, single precision should be used on 64-bit machines, and double precision on 32-bit machines. To select single precision, the symbols "<DOUBLE>" and "<DP>" are replaced by "REAL" and "E"; for double precision, they are replaced by "DOUBLE PRECISION" and "D".

The replacement of the edit symbols is handled automatically by the makefile (using the sed script gsm.sed). To change any one of the above two options, this script must be modified accordingly. The editing step may be eliminated if the options are not expected to be changed - changes in resolution will not require edit symbol replacements.

3.2. Resolution changes

The program allows great flexibility in the choice of spectral truncation and vertical resolution. Changing either requires changing parameter values in the include blocks containing the parameters (gsmdim.blk and param.blk), in the work space include block (work0.blk), and, if applicable, changing the data statement for the vertical levels (in block data vrtst0, in gdata.f). In addition, there are some vertical resolution dependent constants in the radiation scheme which must be changed in the initialization entry point of that module.

Although it is relatively painless to change the resolution in the GSM we caution the reader that the parameterizations have been tuned for rhomboidal 40 truncation, using 18 layers as described by Norquist (GL-TR-90-0285). Increasing resolution may require retuning or possibly even reformulating the physics. Otherwise it is possible that increasing resolution will degrade forecast skill. In addition constants describing the time step and diffusion coefficients should be changed to be compatible with the new resolution. Finally the input boundary value data sets will need to be recreated at the new resolution. These aspects of changing the model resolution are not addressed in this document.

Any pentagonal spectral truncation can be selected by choosing the following three parameters (in param.blk):

- **NMAX** the maximum value of the meridional index n
- **MMAX** the maximum value of the zonal wavenumber m
- **NMAX0** the maximum value of n at m=0

The popular rhomboidal (R) and triangular (T) truncations are selected by letting

- rhomboidal (e.g., R30): NMAX0=MMAX (=30), NMAX=NMAX0+MMAX (=60)
- triangular (e.g., T30): NMAX0=MMAX=NMAX (=30).

In addition, the following two parameters allow the computational domain to be a subregion of the globe

- **PER** the periodicity in the zonal direction (=1 for a global domain)
For a global domain (covering both hemispheres and all longitudes), PER=1 and NHEM=2. The value of NHEM must be consistent with the choice of the <HEM> and <GLO> edit symbols in gsm.sed.

The vertical resolution is controlled by the two parameters (in gsmdim.blk):

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>KPDIM</td>
<td>number of layers</td>
</tr>
<tr>
<td>KHDIM</td>
<td>number of layers which carry humidity.</td>
</tr>
</tbody>
</table>

The arrangement of the levels in the vertical is set by a data statement in block data vrtst0, in gdata.f, for the array SI(H) sigma at level H.

This data statement must be consistent with a top-down vertical structure, i.e. SI(1)=0 and SI(KP+1)=1.

Based on the selection of these seven resolution-determining parameters (NMAX through KHDIM), several other parameters related to the number of spectral coefficients and the size of the transform grid must be determined. A standalone program (asize.f) exists to compute values for remaining parameters that need to be specified in gsmdim.blk, param.blk and work0.blk. The variables described here, as well as all other important variables in the GSM, are all described in the comments of the file containing the common blocks in which they are stored. Further a complete global list of these variables is presented in the appendix to this guide.

3.3. Input and output files

The input and output to and from the GSM involves files of name 'tapenn', where nn is the unit number (the only exception to this rule is output to unit IOTERM: no file name is specified for this unit, allowing the use of "standard out" if the appropriate value for IOTERM is used). The c-shell script (rungsm) provides the link between these internal file names and user-chosen file names. The unit numbers are stored in cntrl.blk, and initialized in cntrl0 (in gdata.f). The following lists the variable name along with the current value of the input and output units:

**INPUT:**
- IOCNTL=99: control input - read in by CTRLID
- IOGSM1=11: GSM initial conditions spectral coefficients input data set (either for initial start - read in by RDINIT, or for a restart - read in by RHIST)
- IOGSM7=17: GSM surface data input data set (not needed for restart) - read in by RDINIT

**OUTPUT:**
- IOTERM=6: (std out) printed output for console
- IOLIST=16: printed output for list file
- IOGSM2=12: GSM forecast spectral coefficients output data set - written out by WDATA, or WDATA2 (if only one time step and not an NMI run), or WNMI (if an NMI run)
IOGS11=21: GSM restart output data set (written by WHIST)

In addition, there are the following internal files, and diagnostic output files (some of which are not currently written to):

- IOGS15=15: GSM holding file for radiation
- IOGS25=25: GSM diagnostic output file for radiation
- IOGS26=26: GSM diagnostic output file for Kuo convection
- IOGS27=27: GSM diagnostic output file for Kuo convection
- IOGS50=50: GSM diagnostic output file for pbl.

The file structure of the input and output files is described in more detail in the following. The file structure conforms with current GL practice, with the following exceptions:

- control input:

  Istart greater than or less than zero indicates a restart or NMI calculation.

  Ntime less than or equal to zero indicates preliminary values or tendencies from the first time step will be output.

  Floating point values are used for the record 2 variables (trms, hfinc, hrsinc) described below. If hrsinc is less than zero, the history restart output file is appended to rather than rewound each time a restart record is output. This option was absent from the GL GSM.

- spectral coefficient data set:

  The precipitation record described below is written out entirely whenever the spectral arrays are written.

- history restart data set:

  The header record contains the value of nstep.

  This data set may be appended to (see description of hrsinc).

3.3.1. Control input

Read in by CTRLID (in gmain.fed)

The control input file allows setting a number of values for the current GSM without having to recompile it. They are read in using list-directed I/O from an ASCII file, in the following order:

Record 1:

  istart: restart/nmi run flag ( >0 for restart, <0 for nmi run, =0 for normal initial start )

  ntime: no. of time steps for this run (integer)

  ntime = or < 0 implies that tendencies or preliminary values will be written after the first, unfiltered time step, either by wnmi (if istart<0) or wdata2 (if istart=0)

  ntime= 0, -1, -2, -3 implies that STEP1 will return after the
To do a restart, set istart=number of filtered time steps of the restart data set, and ntime=number of timesteps left that are necessary to finish the forecast:

\[
ntime(\text{restart fcst}) = ntime(\text{total fcst}) - \text{istart}.
\]

3.3.2. Spectral coefficient data set

Read in by RDINIT; written out by WDATA, WDATA2.

This is a binary data file, consisting of the following:

<table>
<thead>
<tr>
<th>Record 1:</th>
<th>(Header record)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Four variables: nstep (integer), htime (real), itime (character<em>8), idate (character</em>8)</td>
<td></td>
</tr>
</tbody>
</table>

They have the following meaning:

- **nstep**: number of filtered time steps completed (a negative number is used to denote an unfiltered data set)
- **htime**: forecast hour of data set
- **itime(1:4)**: initial state time in 'hhz' format
- **idate(1:8)**: initial state date in 'ddmonyy' format (where mon is the 3-letter, lower case month designator; other formats require changes to the tempus subroutine in gmain.fed)

Initial data sets (as read by RDINIT) must have htime=0 and nstep=0.

At present, the only time an unfiltered data set is written out in this format is by WDATA2, which is called if ntime = or < 0 and istart=0 is specified in the input (see the discussion under control input): depending on the value of ntime, all or part of the first time step is completed, and either tendencies or unfiltered values are written out. In any case, nstep in the file is -1, and htime=dt.

Records 2 - 6: (spectral coefficients of data)

The spectral coefficients of absolute vorticity, divergence, temperature, and specific humidity (all at KP layers), and of...
ln(surface pressure/1000 mb), corresponding to nstep and htime.

NOTE:
For each layer, all spectral coefficients are present for a rhomboidal truncation with m<= MMAX, (n-m)<= NMAXO, with m increasing along the diagonals. On input, the inactive spectral coefficients are ignored; on output, they are output as zeros.

Record 7: (surface geopotential)
The spectral coefficients of the surface geopotential.

Record 8: (Precip - only present for nstep > 0)
This record contains 3 global arrays (on the transform grid - nlont*nhem*nlatt values) plus 4 global accumulators (nstep+1 values each, one for each time step) - all are floating point arrays:

geprev - total precip at time htime
gecpv - convective precip at time htime
evpmp - scaled evaporation at current time step
geptot,cvptot - total, convective global precip at each time step
evpot - total evaporation at each time step.

NOTE:
To avoid memory contention problems, the actual dimension and number of locations used in many arrays are two separate parameters. For example, the number of longitudes of the transform grid (NLONT) may be 128. On the CRAY2 using NLONT to dimension arrays would be disastrous in this case. Therefore the actual array dimension (NLONTD) is taken to be 129. However, file storage is always in terms of the actual number (NLONT). Therefore, depending on the values chosen for NLONT and NLONTD, the internal storage of the gridpoint arrays may differ from the storage in file - this is handled automatically by the I/O routines. Thus, the value of NLONTD (but not of NLONT) may be changed without having to modify the input files.

3.3.3. NMI data set
Written out by WNMI.
This is a binary data file, consisting of the following:

Record 1: (Header record)
Four variables: nstep (integer), htime (real),
itime (character*8), idate (character*8)
They have the following meaning:
nstep - number of filtered time steps completed (a negative number is used to denote an unfiltered data set)
htime - forecast hour of data set
itime(1:4) - initial state time in 'hhZ' format
idate(1:8) - initial state date in 'ddmonyy' format (where mon is the 3-letter, lower case month designator; other formats require changes to the tempus subroutine in gmain.fed)
At present, the only time a data set is written out in this format is by WNMI, which is called if ntime = or < 0 and istart < 0 is specified in the input (see the discussion under control input): depending on the value of ntime, all or part of the first time step is completed, and either tendencies or unfiltered values are written out. In any case, nstep in the file is -1, and htime=dt.

Records 2 - 6: (spectral coefficients of initial data)

These records are in the same format as records 2-6 of the spectral coefficient data set, except that the values of the initial data are written out (valid at htime=0).

Records 7 - 11: (spectral coefficients of tendencies or unfiltered values)

These records are in the same format as records 2-6 of the spectral coefficient data set, except that the values of the tendencies or unfiltered values are written out (valid at htime=dt).

Record 12: (surface geopotential)

The spectral coefficients of the surface geopotential.

3.3.4. Surface data set

Read in by RDINIT

This is a binary data file, consisting of the following:

Record 1: (Header record)

Four variables: nstep (integer), htime (real), itime (character*8), idate (character*8)

These are not used inside the program (other than being echoed to the list file), and are used only for descriptive purposes.

for version 5.1:

Records 2 - 10: (global fields of data)

for version 5.2:

Records 2 - 5 and 9 - 13: (global fields of data)

These records contain values of the following fields on the transform grid (nlont*nhem*nlatt values), at 1 or 2 levels (nlev=1 or 2):

\texttt{nlev}

\begin{verbatim}
1 ts - surface (skin) temperature  
1 cdr - drag coefficient  
1 salb - solar albedo  
1 esd - equivalent snow depth  
2 stc - soil temperature  
1 ws - surface moisture  
2 smc - soil moisture  
1 canopy - vegetation cover
\end{verbatim}
1. *shtbot* - temperature below lowest soil layer
   
   for version 5.2 only:
   
   Records 6 - 8: Fixed fields used by radiation
   
   These are zonal mean fields, at nhem*nlatt latitudes and npmrad levels
   (npmrad = 41 at present) (ordered as j=1,nhem*nlatt, k=1,npmrad):
   
   - clpr - pressure levels for the next 2 fields
   - clh2o- climatological values of water vapor
   - clo3 - climatological values of Ozone

3.3.5. Restart data set

Read in by RHIST; written out by WHIST.

This is a binary data file, consisting of the following:

Record 1: (Header record)

Four variables: nstep (integer), htimel (real),

*itime* (character*8), *idate* (character*8)

They have the following meaning:

- nstep - number of filtered time steps completed (this must
  be a positive number)
- htimel - forecast hour of the filtered time step
- itime(1:4) - initial state time in 'hhZ' format
- idate(1:8) - initial state date in 'ddmonyy' format (where mon is
  the 3-letter, lower case month designator; other formats
  require changes to the tempus subroutine in gamin.fed)

for version 5.1:

Records 2 - 10: Identical to records 2-10 of the surface data set

for version 5.2:

Records 2 - 13: Identical to records 2-13 of the surface data set

Records 14 - 16: Global fields of convection parameters used by

radiation

The global fields are written as (i=1,nlont, j=1,nhem*nlatt)

- cnvpre - convective precipitation rate
- kcnvtop- layer index of convective cloud top
- kcnvbot- layer index of convective cloud bottom

The record numbers in the following are correct for version 5.1. For

version 5.2, they have to be incremented by 6.

Records 11 - 10+(nlatt+1)/2: (radiative heating rates)

These (nlatt+1)/2 records contain the radiative heating rates and the

surface flux. Each record contains nlont*nhem*(kp+1) values. These data

are necessary for a restart at a time step without a full

radiation calculation. The internal storage of the data differs from

the storage in the input/output data set. The file format was chosen

for compatibility with the previous version of the GL GSM.

NOTE: in the original GL version, there were only nlatt/2 records

Records 11+(nlatt+1)/2 - 15+(nlatt+1)/2: (spectral coefficient data at

A-9
These records are in the same format as records 2-6 of the spectral coefficient data set.

Record 16+(nlatt+1)/2: (surface geopotential)

The spectral coefficients of the surface geopotential as record 7 of the spectral coefficient data set.

Record 17+(nlatt+1)/2: (Precip valid at time htimel)

This record contains 5 global arrays (on the transform grid - nlont*nhem*nlatt values):

- geprev - total precip at time htimel
- gecpv - convective precip at time htimel
- gevmp - scaled evaporation for current time step
- devmp - accumulated total dew
- etotbl - accumulated total evaporation

Records 18+(nlatt+1)/2 - 22+(nlatt+1)/2: (spectral coefficient data at time htime2)

These records are in the same format as records 2-6 of the spectral coefficient data set, except they contain data for the unfiltered time step (valid time htime2=htime+dt, time step number nstep+1)

Record 23+(nlatt+1)/2: (Precip valid at time htime2 plus global accumulators)

This record contains 2 global arrays (on the transform grid - nlont*nhem*nlatt values) plus 4 global accumulators (nstep+1 values each, one for each time step) - all but nkuo are floating point arrays:

- geshem - total precip at time htime2
- gecnv - convective precip at time htime2
- geptot,cvptot - total, convective global precip at each time step
- nkuo - number of gridpoints with Kuo convection at each time step
- ** integer array - note difference from spectral coefficient data set **
- evptot - total evaporation at each time step.

3.4. Multiprocessing

The code is designed to be multiprocessed on a shared memory machine. The GSM contains comments of the form "C<mp> ... " in several modules. These comments are meant to help in the implementation of the multiprocessing, and provide inline documentation after implementation.

We have implemented multiprocessing as Cray microtasking directives, and task common statements. The CRAY autotasking directives are activated by invoking cf77 with the -Zp option. Generally speaking autotasking automatically makes maximum use of available resources. However the UNICOS command :;ctenv NCPUS may be used to restrict the GSM to fewer than the full set of available processors.
All common blocks in the GSM are global and should be shared among the processors, except for the following, which are local to each task and are identified in the code as task common. These task common blocks contain data for a single latitude.

/aleg/, in aleg.bed  - Legendre functions at a single latitude
/grdpt/, in grdpt.blk - gridpoint values of variables at one latitude pair
/prod/, in prod.blk  - gridpoint values of nonlinear terms at one latitude pair
/work1/, /worksp/, in workO.blk  - workspace common
/pblval/, in pblvar.blk - pbl variables local to each latitude
/abci/, in gpbl.f    - pbl variables local to each latitude

Results with multiprocessing are not strictly reproducible because the order of addition in the Gaussian sums (i.e. the grid point to spectral transforms) is arbitrary. Differences between runs due to this are initially the size of round off error, but may amplify exponentially. Runs using a single processor are reproducible. In addition, for testing purposes only, we have included some code to enforce the order of the sums. This code, named spin.f, is commented out and is extremely inefficient. We have ourselves used it only to make sure that the differences between multiprocessing runs is in fact due to differences in the order of the sums and not some other problem.

We note that the radiation parameterization performs a full calculation only for odd latitudes. The results for an even latitude are simply taken to be that calculated for the neighboring, equatorward odd latitude. Consequently the multiprocessing for the main latitude loop in laloop, parcels out the latitudes in pairs. Changes to the radiation parameterization which increase or decrease the coupling between latitudes will generally have implications for the multiprocessing strategy. For example if the radiation calculation was performed in full for each latitude, then the multiprocessing in laloop could be done latitude by latitude singly.

4. List of filenames

The following is a complete list of the file names of the GSM code modules currently in use:

4.1. Include blocks with edit symbols:

- aleg.bed  - Legendre functions at a single latitude
- bl.bed    - surface geopotential
- cpcnst.bed - computational and physical constants
- dtime.bed - spectral coefficients of time level 3
- ipmn.bed  - constants used for Pmns calculation
- tme1.bed  - spectral coefficients of time level 1
- tme2.bed  - spectral coefficients of time level 2

4.2. Include blocks without edit symbols

- char.blk  - character constants
- clock.blk - variables related to time stepping and valid time

A-11
cntrl.blk - control parameters
ctauw.blk - global array used by modkuo
czalb.blk - albedo and internal radiation array
dercot.blk - common used by dercot.f
fftn0.blk - constants used by FFTs
grdim.blk - parameter statements used in radiation package
grdpt.blk - gridpoint values of variables at one latitude pair
ground.blk - global arrays of surface parameters
gsmdim.blk - PARAMETER statements, included in param.blk
param.blk - PARAMETER statements
pblvar.blk - common used in pbl package
prod.blk - gridpoint values of nonlinear terms at one latitude pair
ridiag.blk - common used in radiation package
tconst.blk - constants for time scheme
vrtst.blk - vertical structure information
waven.blk - wavenumber arrays
wavvec.blk - arrays needed for gtranv (vectorizing Gaussian sums)
work0.blk - work space common

For version 5.2:
mstrad.blk - convection parameters used by radiation

4.3. Fortran modules with edit symbols

gconvec.fed - Module referenced by gmdkuo.f (Kuo convection)
gcphys.fed - Cover routine for adjustment physics
gmain.fed - Main routine and time step driver and I/O routines
gtend.fed - tendency calculation (hydrodynamics) routines
gtrans.fed - version of gtranv.fed for completely general truncation
(Gaussian sums will not vectorize)
gtranv.fed - spectral transform routines: includes cover routine
for FFTs, but not the FFTs themselves
gtstep.fed - time stepping routines
gutil.fed - utility subroutines

4.4. Fortran modules without edit symbols

dercot.f - Derickson and Cotton code for saturation vapor pressure
dumphy.f - dummy routine for convection
dumrad.f - dummy routine for radiation
gbidum.f - dummy routine for PBL
gdata.f - block data statements
gfttet.f - dry adiabatic adjustment
glrgsc.f - large scale precipitation
gmdkuo.f - Kuo convection
gpbl.f - pbl package
grad.f - radiation package
gradia.f - radiation package
rdsdum.f - dummy version of rdsf-cl.g.l
rdsf-cl.g.l - module to read in boundary data for pbl and radiation
spin.f - module to enforce same order of Gaussian sums
svp.f - module for computing saturation vapor pressure
tcheck.f - checkpointing and error handling routines
work0.f - work space allocation routine
4.5. Other related files

- gsm.sed - sed script for replacing edit symbols
- makefile - makefile for compiling and linking GSM
- rungsm - c-shell script for running gsm
- asize.f - standalone program for computing dimension parameters

0. Define outline-mode:

::: Local Variables: ***
::: mode:outline ***
::: outline-regexp: "\((0-9)+\./\)" ***
::: End: ***
Appendix B: List of Variables

The document reproduced on the following pages is included as a plain text file (named "variables.list") in the software deliverables of this contract.
Variable dictionary

Alphabetical list of commons, include file names (without .blk), variables in commons, and parameters. Also shown are the include file names where they are defined, and a short description. Descriptions that apply to several variables are only shown for the first one in this list - others are cross-referenced. Additional descriptions can be found in the include files and the fortran source code itself. This list is applicable to version 5.1.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Include file</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>prod</td>
<td>a = Z*U + ... Eq. (3)</td>
</tr>
<tr>
<td>ae</td>
<td>param</td>
<td>radius of the earth.</td>
</tr>
<tr>
<td>aleg</td>
<td>aleg</td>
<td>aleg contains values of the associated Legendre functions, their meridional derivatives, etc., and the constants needed in the quadrature at a single latitude. All variables are defined at a single Northern Hemisphere latitude.</td>
</tr>
<tr>
<td>alfa</td>
<td>pblvar</td>
<td>alpha constant which determines if time scheme is backwards implicit (alpha=1) or semi-implicit (alpha=0.5)</td>
</tr>
<tr>
<td>alpha</td>
<td>tconst</td>
<td>A matrix Eq. (B4)</td>
</tr>
<tr>
<td>amtrx</td>
<td>tconst</td>
<td>beta: time filter factor</td>
</tr>
<tr>
<td>andre</td>
<td>tconst</td>
<td>A matrix Eq. (B4)</td>
</tr>
<tr>
<td>b</td>
<td>prod</td>
<td>b = Z*V + ... Eq. (4)</td>
</tr>
<tr>
<td>beta</td>
<td>pblvar</td>
<td>time filter factor</td>
</tr>
<tr>
<td>bl</td>
<td>bl</td>
<td>D matrix appearing in Eq. (B11) as the form A<em>C + R</em>T0*(transpose DEL) where DEL appears in the text as r.</td>
</tr>
<tr>
<td>bmatrx</td>
<td>tconst</td>
<td>D matrix appearing in Eq. (B11) as the form A<em>C + R</em>T0*(transpose DEL) where DEL appears in the text as r.</td>
</tr>
<tr>
<td>bowen</td>
<td>pblvar</td>
<td>canopy - canopy moisture</td>
</tr>
<tr>
<td>canopy</td>
<td>ground</td>
<td>ybar(L,IHEM) the vertical integral from sigma = 0 to 1 at longitude L in hemisphere IHEM of Y = C and D where C is the advection of Q, i.e. velocity* gradient(Q).</td>
</tr>
<tr>
<td>cbar</td>
<td>prod</td>
<td></td>
</tr>
<tr>
<td>char</td>
<td>char</td>
<td>contains character variables</td>
</tr>
<tr>
<td>clock</td>
<td>clock</td>
<td>contains various parameters related to the forecast valid time</td>
</tr>
<tr>
<td>cmtrx</td>
<td>tconst</td>
<td>C matrix defined following Eq. (B3) contains variables defining forecast time parameters, and the I/O units, including, the specific heat of dry air at constant pressure</td>
</tr>
<tr>
<td>cntrl</td>
<td>cntrl</td>
<td></td>
</tr>
<tr>
<td>cp</td>
<td>param</td>
<td></td>
</tr>
<tr>
<td>cpbinm</td>
<td>pblvar</td>
<td>contains computational and physical constants and all the values of the weights, GWGT, and the sines of the quadrature latitudes, GLAT. The transform grid (GLAT,GWGT) is always Gaussian.</td>
</tr>
<tr>
<td>cpcnst</td>
<td>cpcnst</td>
<td>contains a grid point common</td>
</tr>
<tr>
<td>ctauw</td>
<td>ctauw</td>
<td></td>
</tr>
</tbody>
</table>
block of horizontal convergence. It is computed in nlpred, and used in modkuo.

$c\theta 1(K), c\theta 2(K)$ constants for computing level temperatures used in Eq. (39).

$c\theta 1 vrtst$

$c\theta 2 vrtst$

$moisgl$

$czalb$

$czalb$

contains gridpoint arrays of surface parameters needed by the PBL routine.

$d x(L,K,IHEM)$ the value at longitude $L$ in hemisphere $IHEM$ in layer $K$ of $X = Z, D, T, W, U,$ and $V$ (W defined only for moisture-carrying layers).

$dlb$ the IBth spectral coefficient in layer $K$ of $X = Z, D, T$ and $W$. ($n=m$ even)

$dlc$ the ICth spectral coefficient in layer $K$ of $X = Z, D, T$ and $W$. ($n=m$ odd)

$ddtb$ the IBth spectral coefficient tendency in layer $K$ of $X = Z, D, T$ and $W$. ($n-m=even$)

$ddtc$ the ICth spectral coefficient tendency in layer $K$ of $X = Z, D, T$ and $W$. ($n-m=odd$)

$del$ thickness of layer $K$ Fig. (1)

$delta tconst$ difference in time between time levels 1 and 3 (s)

$dercot$ Common block for the saturation vapor pressure calculation

$dew$ accumulated total dew

$dewmis$ diffusion coefficient for $D$

$dkh$ the value at longitude $L$ in hemisphere $IHEM$ of the derivative of $Q$ w.r.t. $Y = L$ and $P$ which are lambda and phi, i.e. the longitude and latitude respectively:

$cos(\phi) \cdot \text{gradient}(Q) = (dqdl, dqdp)$

$dqdl$ the IBth spectral coefficient tendency of $Q$ ($n=m$ even)

$dpd$ see under dqdl

$dqdtb$ the IBth spectral coefficient tendency of $Q$ ($n-m=even$)
<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dqtetc</td>
<td>dtime</td>
</tr>
<tr>
<td>dsoil</td>
<td>pblvar</td>
</tr>
<tr>
<td>dt</td>
<td>tconst</td>
</tr>
<tr>
<td>dtdt</td>
<td>pblvar</td>
</tr>
<tr>
<td>dtdtb</td>
<td>dtime</td>
</tr>
<tr>
<td>dtdtc</td>
<td>dtime</td>
</tr>
<tr>
<td>dtime</td>
<td>dtime</td>
</tr>
<tr>
<td>dtimeb</td>
<td>dtime</td>
</tr>
<tr>
<td>dtsodt</td>
<td>pblvar</td>
</tr>
<tr>
<td>dudt</td>
<td>pblvar</td>
</tr>
<tr>
<td>dvdt</td>
<td>pblvar</td>
</tr>
<tr>
<td>dwdtb</td>
<td>dtime</td>
</tr>
<tr>
<td>dwdtc</td>
<td>dtime</td>
</tr>
<tr>
<td>dwdotd</td>
<td>pblvar</td>
</tr>
<tr>
<td>dzdtb</td>
<td>dtime</td>
</tr>
<tr>
<td>dzdtc</td>
<td>dtime</td>
</tr>
<tr>
<td>e</td>
<td>prod</td>
</tr>
<tr>
<td>ehh</td>
<td>cpcnst</td>
</tr>
<tr>
<td>ep</td>
<td>pblvar</td>
</tr>
<tr>
<td>eps1</td>
<td>ipmn</td>
</tr>
<tr>
<td>epslon</td>
<td>ipmn</td>
</tr>
<tr>
<td>epslr</td>
<td>cpcnst</td>
</tr>
<tr>
<td>esatdc</td>
<td>dercot</td>
</tr>
<tr>
<td>escl</td>
<td>cpcnst</td>
</tr>
<tr>
<td>esc2</td>
<td>cpcnst</td>
</tr>
<tr>
<td>esdc</td>
<td>dercot</td>
</tr>
<tr>
<td>etotbl</td>
<td>moisgl</td>
</tr>
<tr>
<td>evap</td>
<td>pblvar</td>
</tr>
<tr>
<td>evppmp</td>
<td>moisgl</td>
</tr>
<tr>
<td>evptot</td>
<td>moisgl</td>
</tr>
<tr>
<td>fdown</td>
<td>pblvar</td>
</tr>
<tr>
<td>fftn0</td>
<td>fftn0</td>
</tr>
<tr>
<td>freqrd</td>
<td>param</td>
</tr>
<tr>
<td>gdc</td>
<td>dercot</td>
</tr>
<tr>
<td>gecnv</td>
<td>moisgl</td>
</tr>
<tr>
<td>ge cvp</td>
<td>moisgl</td>
</tr>
<tr>
<td>ge prev</td>
<td>moisgl</td>
</tr>
<tr>
<td>geptot</td>
<td>moisgl</td>
</tr>
<tr>
<td>geshem</td>
<td>moisgl</td>
</tr>
</tbody>
</table>
gflx  pblvar   gflx pblvar

glat  cpcnst  glat(LA) sine of Gaussian
    latitudes (transform grid)

gninv  tconst  stored matrix inverses needed by
          semi-implicit time scheme. (See
          comments in IMPLCT.)

grav  param  g, acceleration of gravity
grddim  grddim  Defines several parameters for
          the dimensions of arrays and for i/o
          units. The important thing is that the
          radiation calculation is performed on a
          separate grid with ip longitudes and jp
          latitudes.

grdpt  grdpt  contains grid point values for a
          single Gaussian latitude of the
          variables:

ground  ground  contains gridpoint arrays of
          surface and ground parameters needed
          by the PBL routine. These arrays are read
          in at the start of, and carried along
          during the forecast.

gs2rd  cpcnst  gs2rd, GS7E5 boundary layer flux
          constants:
          \[ GS2RD = \sqrt{2} \left( \frac{g \sigma(KP)}{R \Delta(KP)} \right) \]
          \[ GS7E5 = 7 \times 10^{-5} \left( \frac{g \sigma(KP)}{R \Delta(KP)} \right) \]
          where KP is the layer next to the
          surface.

gs7e5  cpcnst  see under gs2rd

gsmdim  gsmdim  Parameters of array dimensions
          used in plug-compatible modules, and in
          param.blk:

gwgt  cpcnst  gwgt(LA) Gaussian quadrature
         weights (transform grid)

hab  aleg  hab(IB) ILAPB(IB)*HB/AE

hac  aleg  hac(IC) ILAPC(IC)*HC/AE PAB and HAB are
         used to calculate UMS and VMA, PAC and
         HAC are used to calculate UMA and VMS,
         in Eqs. (32) and (33).

hb  aleg  hb(IB) the value of the meridional
        derivative of PB: HB =
        \[ -\cos(\phi) \frac{d(PB)}{d(\phi)} \]

hc  aleg  hc(IC) the value of the meridional
        derivative of PC: HC =
        \[ -\cos(\phi) \frac{d(PC)}{d(\phi)} \]

hdc  dercot

hdf  pblvar

heat  pblvar

hfinc  cntrl  Forecast output increment

hflon  cntrl  Forecast length (total)

hfzero  cntrl  =1 to output initial
              conditions on IOGSM2

ho  clock  ho, idy, mo, iyr - hour (hh), day (dd),
          month (mm), and year (yy) of forecast
          initialization time (GMT)

hpbl  pblvar

B-4
hpinc  cntrl  Precipitation output increment (NOT USED)
hpzero cntrl  Precipitation accumulator zeroing increment
hrsinc cntrl  Restart output increment
htime1 cntrl  Forecast time of variables in /TIME1/
htime2 cntrl  Forecast time of variables in /TIME2/
ib00  cntrn  value of index IB for which m=0 and n=0
ib0n  waven  array of array indices IB for which m=MB(IB)=0
ic01  cntrn  value of index IC for which m=0 and n=1
ic0n  waven  array of array indices IC for which m=MC(IC)=0
icl  ridiag
idate  char  idate,ITIME  Initial state date/time
idum1 cntrl  idum9  IDUM1  Dummy entries for additional I/O units, this allows units to be added without changing the common block size
idum2 cntrl  see under idum1
idum3 cntrl  see under idum1
idum4 cntrl  see under idum1
idum5 cntrl  see under idum1
idum6 cntrl  see under idum1
idum7 cntrl  see under idum1
idum8 cntrl  see under idum1
idum9 cntrl  see under idum1
idy  clock  see under ho
iendw0 work0  last element of work array allocated
ifax  fftn0
iheat  ridiag
iheati  ridiag
iheats  ridiag
ilapb waven ilapb(IB)  inverse Laplacian operator for n=NB(IB)
ilapc waven ilapc(IC)  inverse Laplacian operator for n=NC(IC)
iocnt1 param  Control input
iogs11 cntrl  GSM restart output data set
iogs15 cntrl  GSM holding file for radiation
iogs25 cntrl  GSM holding file for radiation
iogs26 cntrl  GSM diagnostic output file for Kuo convection
iogs27 cntrl  GSM diagnostic output file for Kuo convection
iogs50 cntrl  GSM diagnostic output file for pbl
iogsm0 cntrl  GSM tendencies s. c. output data set (NOT USED)
iogsm1 cntrl  GSM initial conditions spectral coefficients input data set (either for
iogsm2 cntrl initial start or for restart
GSM forecast spectral coefficients output data set
iogsm7 cntrl GSM surface data input data set
iogsm8 cntrl GSM precipitation output data set (NOT USED)
iolist param List output
ioterm param Terminal output
ipdim grddim
contains the constant parts of the PMNS computation. Refer to routines INIPMN and PMNS (in GTRANS) for more details.
ipmn

isfcf ridiag
isoi1 pblvar
istrw0 work0
istrw0(I) first element of WORK0 allocated at subroutine level I

itldc dercot see under idate
it2dc dercot
itime char
iuerr grddim see under ho
iuout grddim

iyr clock corresponding julian day

jpblp pblvar
jpdim grddim

julday clock

kflux ridiag
kh param

khl param

khdim gsmdim
kh: no of moist sigma layers
lower index of humidity loop, i.e.
there is moisture in layers
K=KHL, ..., KP

kp param
number of layers (=kpdim
from gsmdim.blk)

kpdim grddim
kp: no of sigma layers

lapb waven lapb(IB) Laplacian operator corresponding to NB(IB)
lapc waven lapc(IC) Laplacian operator corresponding to NC(IC)
latpbl pblvar

lbdiag wavvec lbdiag(IBON) - Number of even spectral coefficients along the diagonal
n-m=2*(IBON-1)
lcdiag wavvec lcdiag(ICON) - Number of odd spectral coefficients along the diagonal
n-m=2*(ICON-1) + 1

lonpbl pblvar
maxstp moisgl
mb waven mb(IB) zonal wavenumber m corresponding to the 18th even spectral coefficient
mbldim gsmdim maximum no of sigma layers used in pbl
mc waven mc(IC) zonal wavenumber m corresponding to the ICth odd spectral coefficient
mmax param the maximum value of m
mmb waven mmb(IB) index in the Fourier
coefficient arrays that corresponds to
the wavenumber m=MB(IB)
mnc waven mnc(IC) index in the Fourier
coefficient arrays that corresponds to
the wavenumber m=MC(IC)
mo clock see under ho
moisgl moisgl contains gridpoint arrays of
precip etc., and precip counters
moist moisgl
mpdim grddim
mzbl pblvar
nb waven nb(IB) meridional index n corresponding
to the IBth even spectral coefficient

nb0n param length of IBON array (in /WAVEN/)
c waven nc(IC) meridional index n corresponding
to the ICth odd spectral coefficient
nc0n param length of IBON array (in /WAVEN/)
ndiff cpcnst minimum n wavenumber to which diffusion
operator is applied for variables other
than D

ndimi0 work0 size of array ISTRWO
ndimw0 work0 size of work array
nhdim gsmdim nhem: no of hemispheres
nem param 1 for hemispheric, =2 for the global
version (=nhdim from gsmdim.blk)
nkdim grddim
nkuo moisgl number of gridpoints with Kuo
convection at each time step
nlatt param number of Gaussian latitudes in one
hemisphere of transform grid, equator
and pole never included. (=npdim/nhem,
npdim from gsmdim.blk) (must be >=
(2*NMAX0+NMAX+1)/4 for MMAX
<=2*(NMAX-NMAX0) and >= (3*NMAX+.1)/4
for MMAX >= 2*(NMAX-NMAX0) )
nlev0 work0 last element of ISTRWO used, number of
subroutine levels in use

nlon0 fftn0 number of longitudes in transform grid.
(nmust be >=3*NUMZW-2)
nlont param longitude dimension of transform grid
arrays (+mpdim from gsmdim.blk) (must
be >=NLONT )
nmax param the maximum value of n
nmax0 param the maximum value of n at m=0
npbdim pblvar
npbl pblvar
npdim grddim
nrstp cntrl
nsodim gsmdim
nsoil pblvar
timestep clock

nhem*nlatt: no of latitudes of transform grid
Time step to begin forecasting
no of soil layers used in pl (nsoil) + 1
time step number; during STIPN, the
unfiltered forecast is complete for NSTEP steps, and the unfiltered forecast for step NSTEP+1 (NT) is being computed; after STEPN, the filtered forecast for nstep is complete (and stored in time level 1), and the unfiltered one for NSTEP+1 is complete (and stored in time level 2). The corresponding forecast hours are stored in /CNTRL/ as HTIME1 and HTIME2. (During STEPL nstep=0; there is no filtering on the initial time (HTIME1=0), but the data is treated the same way as a filtered forecast after STEPL)

\begin{align*}
  \text{nt} & \quad \text{clock} \\
  \text{numb} & \quad \text{param} \\
  \text{numbd} & \quad \text{param} \\
  \text{numbs} & \quad \text{param} \\
  \text{numc} & \quad \text{param} \\
  \text{numcd} & \quad \text{param} \\
  \text{numcs} & \quad \text{param} \\
  \text{numzw} & \quad \text{param} \\
  \text{omega} & \quad \text{param} \\
  \text{pab} & \quad \text{aleg} \\
  \text{pac} & \quad \text{aleg} \\
  \text{param} & \quad \text{param} \\
  \text{pb} & \quad \text{aleg} \\
  \text{pbl} & \quad \text{pblvar} \\
  \text{pblag} & \quad \text{pblvar} \\
  \text{pblval} & \quad \text{pblvar} \\
  \text{pblvar} & \quad \text{pblvar} \\
  \text{pc} & \quad \text{aleg} \\
  \text{per} & \quad \text{param} \\
  \text{phisc} & \quad \text{bl} \\
  \text{phisc} & \quad \text{bl} \\
  \text{precip} & \quad \text{pblvar} \\
  \text{prod} & \quad \text{prod} \\
  \text{psbl} & \quad \text{grdpt} \\
  \text{psfc} & \quad \text{grdpt} \\
  \text{qlb} & \quad \text{pblvar} \\
  \text{qlc} & \quad \text{timel}
\end{align*}

unfiltered forecast is complete for NSTEP steps, and the unfiltered forecast for step NSTEP+1 (NT) is being computed; after STEPN, the filtered forecast for nstep is complete (and stored in time level 1), and the unfiltered one for NSTEP+1 is complete (and stored in time level 2). The corresponding forecast hours are stored in /CNTRL/ as HTIME1 and HTIME2. (During STEPL nstep=0; there is no filtering on the initial time (HTIME1=0), but the data is treated the same way as a filtered forecast after STEPL)

\begin{align*}
  \text{nt} & \quad \text{clock} \\
  \text{numb} & \quad \text{param} \\
  \text{numbd} & \quad \text{param} \\
  \text{numbs} & \quad \text{param} \\
  \text{numc} & \quad \text{param} \\
  \text{numcd} & \quad \text{param} \\
  \text{numcs} & \quad \text{param} \\
  \text{numzw} & \quad \text{param} \\
  \text{omega} & \quad \text{param} \\
  \text{pab} & \quad \text{aleg} \\
  \text{pac} & \quad \text{aleg} \\
  \text{param} & \quad \text{param} \\
  \text{pb} & \quad \text{aleg} \\
  \text{pbl} & \quad \text{pblvar} \\
  \text{pblag} & \quad \text{pblvar} \\
  \text{pblval} & \quad \text{pblvar} \\
  \text{pblvar} & \quad \text{pblvar} \\
  \text{pc} & \quad \text{aleg} \\
  \text{per} & \quad \text{param} \\
  \text{phisc} & \quad \text{bl} \\
  \text{phisc} & \quad \text{bl} \\
  \text{precip} & \quad \text{pblvar} \\
  \text{prod} & \quad \text{prod} \\
  \text{psbl} & \quad \text{grdpt} \\
  \text{psfc} & \quad \text{grdpt} \\
  \text{qlb} & \quad \text{pblvar} \\
  \text{qlc} & \quad \text{timel}
\end{align*}
The document page contains a list of variables and their descriptions. The text is as follows:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>q2b</td>
<td>time2</td>
<td>q2b(n) the nth spectral coefficient of Q. (n-m-even)</td>
</tr>
<tr>
<td>q2c</td>
<td>time2</td>
<td>q2c(n) the nth spectral coefficient of Q. (n-m-odd)</td>
</tr>
<tr>
<td>qml</td>
<td>pblvar</td>
<td>qml ground moisture</td>
</tr>
<tr>
<td>qs</td>
<td>ground</td>
<td>qs (1-2) time spectral coefficient of Q. (n-m-odd)</td>
</tr>
<tr>
<td>radintd</td>
<td>ridiag</td>
<td>rdate, RTIME Run date/time (time forecast run on computer)</td>
</tr>
<tr>
<td>rdate</td>
<td>char</td>
<td>rdate(K), REX2(K) ratios of Exner function in layer K to that in layers K+1 and K+2. These constants are used in the dry adiabatic adjustment.</td>
</tr>
<tr>
<td>rex1</td>
<td>vrtst</td>
<td>see under rex1</td>
</tr>
<tr>
<td>rex2</td>
<td>vrtst</td>
<td>see under rex1</td>
</tr>
<tr>
<td>rgas</td>
<td>param</td>
<td>R, gas constant for dry air Virtual temperature effects are not included in this model.</td>
</tr>
<tr>
<td>rh</td>
<td>pblvar</td>
<td>Non-flux part of thermodynamic equation. Eq. (29) (RI in the code is I in Brenner et al. (1982).)</td>
</tr>
<tr>
<td>ri</td>
<td>prod</td>
<td>see under rh</td>
</tr>
<tr>
<td>rdiag</td>
<td>rdiag</td>
<td>see under rdiag</td>
</tr>
<tr>
<td>rkappa</td>
<td>cpcnst</td>
<td>kappa, R/Cp</td>
</tr>
<tr>
<td>rt0</td>
<td>tconst</td>
<td>R*T0 (m/s)**2</td>
</tr>
<tr>
<td>rtemp</td>
<td>cpcnst</td>
<td>cosine(alpha)</td>
</tr>
<tr>
<td>rtime</td>
<td>char</td>
<td>see under rtime</td>
</tr>
<tr>
<td>rtsn</td>
<td>cpcnst</td>
<td>sine(alpha) (alpha is the rotation angle between the surface wind and the wind in the lowest layer.)</td>
</tr>
<tr>
<td>salb</td>
<td>cvalb</td>
<td>solar albedo</td>
</tr>
<tr>
<td>si</td>
<td>vrtst</td>
<td>see under si</td>
</tr>
<tr>
<td>sikap</td>
<td>vrtst</td>
<td>see under sikap</td>
</tr>
<tr>
<td>sinlat</td>
<td>aleg</td>
<td>see under sinlat</td>
</tr>
<tr>
<td>sl</td>
<td>vrtst</td>
<td>see under sl/K</td>
</tr>
<tr>
<td>sikap</td>
<td>vrtst</td>
<td>see under sikap(K)</td>
</tr>
<tr>
<td>snoflux</td>
<td>pblvar</td>
<td>see under snoflux</td>
</tr>
<tr>
<td>snow</td>
<td>pblvar</td>
<td>see under snow</td>
</tr>
<tr>
<td>ss</td>
<td>pblvar</td>
<td>see under ss</td>
</tr>
<tr>
<td>t0</td>
<td>vrtst</td>
<td>see under t0(K)</td>
</tr>
<tr>
<td>t1b</td>
<td>time1</td>
<td>see under t1b(K)</td>
</tr>
<tr>
<td>t1c</td>
<td>time1</td>
<td>see under t1c(K)</td>
</tr>
<tr>
<td>t2b</td>
<td>time2</td>
<td>see under t2b(K)</td>
</tr>
<tr>
<td>t2c</td>
<td>time2</td>
<td>see under t2c(K)</td>
</tr>
<tr>
<td>taux</td>
<td>pblvar</td>
<td>see under taux</td>
</tr>
<tr>
<td>taoy</td>
<td>pblvar</td>
<td>see under taoy</td>
</tr>
<tr>
<td>taw</td>
<td>ctauw</td>
<td>see under taw</td>
</tr>
<tr>
<td>tbl</td>
<td>gridpt</td>
<td>see under tbl(K)</td>
</tr>
<tr>
<td>tbot</td>
<td>gridpt</td>
<td>tbot ground temperature below lowest soil layer</td>
</tr>
</tbody>
</table>

Note: The text contains abbreviations and symbols that are typical in meteorological and atmospheric science contexts. The descriptions provide context for the variables, which are likely part of a model or dataset used in atmospheric science.
tconst  tconst  contains constants needed for the implicit time scheme.
timel  timel  contains prognostic variables at time level 1 (the time level preceding the central time):
time2  time2  contains prognostic variables at time level 2 (the central time):
tml  pblvar  surface (skin) temperature
trigs  fftn0  soil temperature
ts  ground  see under d
tsoflx  pblvar  see under tbl
tsoil  pblvar  u
tso1  pblvar  grdpt
u  grdpt  ut = U*T : eastward temperature flux.
inground  see under d
uflx  pblvar  uw = U*W : eastward moisture flux.
tml  pblvar  Eq. (16)
tsl  pblvar  Eq. (17)
tml  pblvar  v
ubl  grdpt  see under d
ubl  grdpt  see under tbl
vt  prod  vt = V*T : northward temperature flux.
vml  pblvar  Eq. (16)
vml  pblvar  Eq. (17)
w  grdpt  w
wlb  time  see under dlb
wlb  time  see under tbl
wlc  time  see under dlc
wlc  time  see under tbl
w2b  time2  see under d2b
w2c  time2  see under d2c
waven  waven  contains the zonal wavenumber m, meridional index n, and related quantities.
wavvec  wavvec  This common block is used by the vectorizing version of the transform routines (gtranv.fed). It assumes that the spectral truncation is pentagonal, and all spectral coefficients within the pentagon and with the given periodicity PER are active (i.e., the "arbitrary truncation" of AER TM NWP-1 is not supported). Furthermore, the following routines and this common block assume that the spectral coefficients are stored with m increasing along the diagonals defined by n-m-i, where i goes from 0 to 2*(NBON-1) (even coeff.) or 1 to 2*(NCON-1)+1 (odd coeff.): GQB, GQC, LSUMB, LSUMC, ADVU, ADVV.
<table>
<thead>
<tr>
<th>Variable</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>wbl</td>
<td>grdpt</td>
<td>see under tbl</td>
</tr>
<tr>
<td>wdf</td>
<td>pblvar</td>
<td>the quadrature weight: GWGT(LA) or AWGT(LA)</td>
</tr>
<tr>
<td>wgtlat</td>
<td>aleg</td>
<td>wj = non-flux transport of moisture. Eq. (30) (WJ in the code is J in Brenner et al. (1982).)</td>
</tr>
<tr>
<td>wj</td>
<td>prod</td>
<td>prod</td>
</tr>
<tr>
<td>wkt</td>
<td>pblvar</td>
<td>contain the work array and related variables.</td>
</tr>
<tr>
<td>work0</td>
<td>work0</td>
<td>work array</td>
</tr>
<tr>
<td>work1</td>
<td>work0</td>
<td>contain the work array and related variables.</td>
</tr>
<tr>
<td>workap</td>
<td>work0</td>
<td>soil moisture</td>
</tr>
<tr>
<td>wsoflx</td>
<td>pblvar</td>
<td></td>
</tr>
<tr>
<td>wsoil</td>
<td>ground</td>
<td></td>
</tr>
<tr>
<td>wsoml</td>
<td>pblvar</td>
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<tr>
<td>xl</td>
<td>pblvar</td>
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<td>z</td>
<td>grdpt</td>
<td>see under d</td>
</tr>
<tr>
<td>z0</td>
<td>ground</td>
<td>roughness length</td>
</tr>
<tr>
<td>z1b</td>
<td>time1</td>
<td>see under d1b</td>
</tr>
<tr>
<td>z1c</td>
<td>time1</td>
<td>see under d1c</td>
</tr>
<tr>
<td>z2b</td>
<td>time2</td>
<td>see under d2b</td>
</tr>
<tr>
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<td>time2</td>
<td>see under d2c</td>
</tr>
<tr>
<td>zlev</td>
<td>pblvar</td>
<td></td>
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</tbody>
</table>