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Implementing the Kernel of the Australian Region Weather Prediction Model in SISAL

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The Australian Region Weather Prediction Model is an experimental FORTRAN code which uses a variable resolution nesting scheme to provide higher resolution predictions over important areas of the Australian continent such as cities and coastal fisheries. In this preliminary study we explore the performance of the SISAL implicit parallel programming language on a significant scientific application by recording the kernel subroutine of the Model in SISAL. Results are presented for a low end SPARC workstation, an entry level Cray Y-MP EL and a high end Cray C90.

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The Australian Region Weather Prediction Model is an experimental FORTRAN code which uses a variable resolution nesting scheme to provide higher resolution predictions over important areas of the Australian continent such as cities and coastal fisheries. In this preliminary study we explore the performance of the SISAL implicit parallel programming language on a significant scientific application by recoding the kernel subroutine of the Model in SISAL. Results are presented for a low end SPARC workstation, an entry level Cray Y-MP EL and a high end Cray C90.

1 Introduction

The Australian Region Weather Prediction Model (ARPE) was developed by the Australian Bureau of Meteorology Research Centre [1] for short-term weather forecasting up to 36 hours. ARPE draws upon the work of Arakawa, Lamb and Miyakoda [2][3] for its formulation and is intended to be a production code for the prediction of weather over the Australian region. This paper will concentrate on the implementation of the core subroutine of the ARPE in the SISAL language and readers are directed to reference [1] for a detailed description of the model. The work is part of a continuing long term international study of SISAL

being conducted in collaboration with the Lawrence Livermore National Laboratory.

2 The SISAL language

SISAL is a functional language for numerical computation [4]. The developers of SISAL have been able to demonstrate performance comparable with FORTRAN on a number of computing platforms including the Cray Research multiprocessors [5].

SISAL prohibits by design the ability to express constructions which lead to the side effects that make compilation for parallel computer systems extremely difficult. Examples of side effects include those which occur through the COMMON and EQUIVALENCE statements in FORTRAN and SISAL has neither of these constructs. SISAL is block structured and superficially resembles a number of modern languages. The single assignment nature of SISAL means variables have values assigned to them once. This requires some departure from a common style of programming where variables are re-used in programs sometimes for unrelated computations. Translation of FORTRAN programs into SISAL is not necessarily a simple process and can be complicated significantly if the program being re-expressed has been the subject of undisciplined maintenance or construction. This may be compounded if there is no original formulation of the mathematical model available. Direct transliteration of well written FORTRAN code can yield satisfactory results.

Most comparative studies to date have involved the complete recoding of an application in SISAL. In this study the mixed language facility of the current (V12.9.1) Optimising SISAL Compiler is used with an initial core subroutine being recoded.

3 The weather prediction model

The Weather Prediction Model code (ARPE) consists of some 10,000 lines of FORTRAN source code. Its pre-processors and ancillary code constitute perhaps another 5,000 lines of code. The code is generally well written with disciplined use of COMMON and EQUIVALENCE statements. The kernel routines make almost no use of subroutines although the structure of the code suggests they should be used. ARPE then is a reasonable example of a code where inlining has occurred from the outset in an attempt to obtain improved performance. It predates modern FORTRAN pre-processors which automatically inline selected subroutines.

4 FORTRAN

The Cray Research FORTRAN tool suite used [6] runs under X Windows and is a marked advance on those generally available only a few years ago. The tool set comprises: a profiler (flowview) which identifies key subroutines and subroutines which are candidates for inlining; a pre-processor which performs inlining and attempts to identify and annotate parallel regions; an assistant for explicit parallel annotation (atscope); and a parallelism estimator (atexpert).

Routine Name	Tot Time	Calls	Avg Time	Percentage	Accum%
INNER2	2.52E+01	9	2.80E+00	42.24	42.24
LIE	1.09E+01	24	4.53E-01	18.23	60.47
PHYS	6.15E+00	5	1.23E+00	10.31	70.79
LIEBIG	5.61E+00	12	4.68E-01	9.42	80.21
LIEH	5.50E+00	12	4.58E-01	9.23	89.44
LIEBH	1.69E+00	9	1.86E-01	2.84	92.28
SEMIMP	1.48E+00	9	1.64E-01	2.48	94.76
VMODES	1.08E+00	4	2.71E-01	1.82	96.57
INNER	9.56E-01	9	1.06E-01	1.60	98.18
DADADJ	4.40E-01	11476	3.84E-05	0.74	98.91
LAMLL	1.43E-01	2600	5.50E-05	0.24	99.15

Table 1: Execution Profile (5 iterations Y-MP EL)

The original program was profiled using flowtrace to identify the core subroutines. For reasons already stated flowtrace did not identify any subroutines eligible for inlining.

The INNER2 subroutine was chosen as the starting point for this study but as it represents only 42% of the run time contribution no significant speedup is to be expected. The LIE and PHYS subroutines will be translated in due course. Our interest here is to confirm that the run time is not adversely affected and that underlying concurrency is uncovered by the OSC compiler.

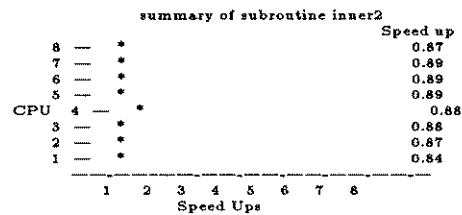


Figure 1: speedup of INNER2 predicted by atexpert

4.1 Results for FORTRAN

The automatic parallel annotator was used to annotate the INNER2 subroutine. No attempt was made to resolve data dependencies in the original FORTRAN in this part of the study although this is intended later. The atexpert measurement tool was used to examine individual DO loops for predicted speedup. Atexpert is claimed to accurately predict performance for dedicated systems. The tool provides parallelism profiles and allows routines associated with parallel or sequential regions to be examined and analysed interactively.

It can be seen in Figure 1 that fpp failed to discover significant parallel regions in INNER2.

5 SISAL

5.1 Mixed language compilation

 **
 ** The osc compiler compiles and links modules written in FORTRAN and SISAL. In this FORTRAN is invoking a SISAL function. To do this the original INNER2 subroutine was replaced by a FORTRAN shell. The shell initialises the array descriptors required by SISAL and calls the replacement INNER2 written in SISAL [7].

Fortunately the array descriptors may be re-used for other arrays which have an identical shape. The ability to specify an offset for returned data structures could be used to avoid the often clumsy process of dealing with boundary values. The current descriptor mechanism unfortunately sets to zero the elements not written to.

5.2 The transliteration process

Although the mathematical formulation was available it did not provide significant assistance in the transliteration process. The INNER2 subroutine was

directly transliterated into SISAL with no restructuring being attempted. A number of unintentional out of bound accesses were discovered in the FORTRAN program during this transliteration.

The transliteration process was significantly complicated by the size of the INNER2 subroutine. While the SISAL debugger (sdbx) gave some assistance there were many cases where sdbx was not able to determine the original source line causing the error. Other minor difficulties which would cause irritation for programmers used to imperative styles also arose. In this case even though the author has a reasonable understanding of SISAL the passage of time since writing his previous SISAL program still led him to be caught by the following:

```

for initial
....
k:=0;
while k < kz repeat
    k:= old k +1;
    .....u[k].....

```

Most programmers will expect k to be 1 when the variable u is accessed on the first loop iteration rather than zero as stated by the for initial clause.

Transliteration and debugging took approximately 35 hours.

5.3 Results for SISAL

The results for one call of INNER2 in FORTRAN and SISAL are shown in Table 2. In their current form both versions are several hundred lines long and the interleaving of initialisation, the calculation of primary meteorological variables and common working variables makes their inner workings difficult to comprehend (Appendices).

Language	Sparc	EL (1-cpu)	C90 (1-cpu)	C90 (4-cpu)
f77 -O	6.6+0.7			
c177 -Zp		3.01+0.48	0.39+0.01	
osc -O	7.2+1.0	6.57+0.35	1.06+0.01	0.29+0.01

Table 2: Run Times for FORTRAN and SISAL

It may be noted that although the run times on the SPARC workstation for FORTRAN and SISAL are comparable performance on the Cray systems is not as good. It is believed that the transliteration resulted in a SISAL style which caused difficulty for the SISAL optimisers; this is currently being resolved.

6 Conclusions

A modest amount of difficulty was encountered in the transliteration of the kernel INNER2 subroutine into SISAL. The run time for this first SISAL implementation relative to FORTRAN is acceptable. Good speedup has been achieved with the SISAL version's runtime falling below that for FORTRAN at four processors. Given this promising start the study will now refine the version of INNER2 and move to the other dominant kernel subroutines LIE and PHYS. The PHYS subroutine is dominated by conditionally executed code as are many other weather codes. It is anticipated that this will produce a more demanding test for SISAL.

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Appendices

INNER2.F

The original code of INNER2 has been stripped out and replaced with descriptor initialisation and call to sinner2.

```

SUBROUTINE INNER2
C
C INNER2 CALCULATES THE RH SIDES OF THE MAIN SEMI-
C IMPLICIT EQUATIONS
C
C include 'arpe.inc'
C PARAMETER
C (
C I2=I1+1, I3=I1+2, I4=I1+3, ILM=IL-1, ILN=IL-2
C J2=J1+1, J3=J1+2, J4=J1+3, JLM=JL-1, JLN=JL-2
C KZM1=KZ-1, KZP1=KZ+1
C CP=1.00464E7, G=980.6, HL=2.501E10, PBAR=1.E6, R=2.87E6
C RV=4.61E+6
C )
C
C COMMON
C /DTDS/ DT,DS,DTI,DSI,DSI2,DSSQ,TDSI,HDTDS,BET65,DTMAX
C + /INTGRL/ PRECP, PRECTA, CKS, EKE, PE, PS-
C BAR, TRHAT, VROMG
C + /KTAU/ KTAU
C
C COMMON
C /CDIFF/ CDIFF(IL,JL)
C + /CORP/ CORP(IL,JL)
C + /DNORM/ DNORM(KZ)
C + /DQ/ DQ(KZ)
C + /DTODQ/ DTODQ(KZ)
C + /EM/ EM(IL,JL)
C + /EMSQ/ EMSQ(IL,JL)
C + /EMSQI/ EMSQI(IL,JL)
C + /GAMA/ GAMA(KZ)
C + /OMEGA/ OMEGA(KZ,IL,JL)
C + /PHI/ PHI(KZ,IL,JL)
C + /PS/ PSM(IL,JL), PS(IL,JL), PSP(IL,JL)
C + /Q/ Q(KZ)
C + /QPH/ QPH(KZ)
C COMMON

```

```

+ /RM/ RM(KZ,IL,IL), RM(KZ,IL,IL), RMP(KZ,IL,IL)
+ /RTBAR/ RTBAR(KZ)
+ /SIGDOT/ SIGDOT(KZ,IL,IL)
+ /T/ TM(KZ,IL,IL), T(KZ,IL,IL), TP(KZ,IL,IL)
+ /TBAR/ TBAR(KZ)
+ /U/ UM(KZ,IL,IL), U(KZ,IL,IL), UP(KZ,IL,IL)
+ /V/ VM(KZ,IL,IL), V(KZ,IL,IL), VP(KZ,IL,IL)
+ /ZS/ ZS(IL,IL)
C
REAL Q
integer ik(100),iij(100),ikij(100)
DIMENSION RMPR(KZ,IL,IL)
DIMENSION TFLBV(KZ),ETFDQ(KZ),WVEL(KZ)
DIMENSION VADVU(KZP1),VADVU(KZP1),VADVVM(KZP1)
DATA VADVU/KZP1*0./,VADVVM/KZP1*0./,VADVVM/KZP1*0./
DATA OMG / 0.0 /
C
c SISAL array descriptors
c one dimension
ik(1)=0
ik(2)=0
ik(3)=0
c
ik(4)=1
ik(5)=kz
ik(6)=1
ik(7)=kz
ik(8)=1
c
c two dimensions
iij(1)=0
iij(2)=0
iij(3)=0
c
iij(4)=i1
iij(5)=i1
iij(6)=i1
iij(7)=i1
iij(8)=i1
c
iij(9)=j1
iij(10)=j1
iij(11)=j1
iij(12)=j1
iij(13)=1
c
c three dimensions
ikij(1)=0
ikij(2)=0
ikij(3)=0
c
ikij(4)=1
ikij(5)=kz
ikij(6)=1
ikij(7)=kz
ikij(8)=1
c
ikij(9)=i1
ikij(10)=i1
ikij(11)=i1
ikij(12)=i1
ikij(13)=1
c
ikij(14)=j1
ikij(15)=j1
ikij(16)=j1
ikij(17)=j1
ikij(18)=1
c
call sinner2(
+dt,ds,dsi,dsi2,tdsi,dmax,cks, eke, pe, psbar, trhat, vromg,
+ktau,
+cdiff,iij,
+corp,iij,
+dnorm,ik,
+dq,ik,
+dtodq,ik,
+em,iij,
+emsq,iij,
+emaq,iij,
+gama,ik,
+omega,ik,
+phi,ikij,
+psm, iij,ps,iij,
+q,ik,
+qph,ik,
+rmm, ikij,rm, ikij,rmp,ikij,
+rtbar,ik,
+sigdot,ikij,
+tm,ikij, t,ikij, tp,ikij,
+tbars,ik,
+um, ikij,u, ikij,up,ikij,
+vm, ikij,v, ikij,vp,ikij,
+zs, iij,
c
returne
+rm,ikij,
+sigdot,ikij,
+up,ikij,
+new'tp,ikij,
+new'rmp,ikij,
+vp,ikij,
+eke,
+cks,
+trhat,

```

```

+pe,
+psbar,
+vromg)
c
RETURN
END

```

inner2.sis

```

define sinner2
% G.K. Egan 1993

type OneDReal = array[real];
type TwoDReal = array[OneDReal];
type ThreeDReal = array[TwoDReal];

global log(a:real returns real)

global sqrt(a:real returns real)

function boundary'cell(i,i1,j,j1)integer returns boolean)
((i = i1) — (i = i1) — (j = j1) — (j = j1))
end function

function divergence'sums(
i,j,kz,i1,j1,jl:integer; dsi:real;
dq:OneDReal;u,v,t:ThreeDReal;emsq:TwoDReal
returns
real, real, OneDReal, OneDReal,
OneDReal, OneDReal, OneDReal)
for initial
sumu:=0.0;
sumv:=0.0;
sumx:=0.0;
k:=1;
while (k < kz) repeat
k:=old k +1;
sumu, sumv, sumx := (
if boundary'cell(i,i1,j,j1) then
old sumu, old sumv, old sumx
else
old sumu+dq[k]* (u[k,i+1,j]-u[k,i-1,j]
+v[k,i,j]+v[k,i+1,j]-v[k,i,j-1]-v[k,i+1,j-1]),
old sumv+dq[k]* (u[k,i,j]+u[k,i,j+1]
-u[k,i-1,j]-u[k,i-1,j+1]+v[k,i,j+1]-v[k,i,j-1]),
old sumx+dq[k]* (u[k,i,j]-u[k,i-1,j]+v[k,i,j]-v[k,i,j-1])
end if)
returns
value of sumu
value of sumv
value of sumx
array of sumu
array of sumv
array of sumx
array of (-emsq[i,j])*sumx*dsi
array of t[k,i,j]
end for
end function

function sinner2(
dt,ds,dsi,dsi2,tdsi,dmax,cks, eke, pe, psbar, trhat, vromg:real;
ktau:integer;
cdiff:TwoDReal;
corp:TwoDReal;
dnorm:OneDReal;
dq:OneDReal;
dtodq:OneDReal;
em:TwoDReal;
emsq:TwoDReal;
emaq:TwoDReal;
gama:OneDReal;
omega:OneDReal;
phi:ThreeDReal;
psm, ps:TwoDReal;
q:OneDReal;
qph:OneDReal;
rmm, rm, rmp:ThreeDReal;
rtbar:OneDReal;
tm, t, tp:ThreeDReal;
tbar:OneDReal;
um, u, up:ThreeDReal;
vm, v, vp:ThreeDReal;
zs:TwoDReal
returns
ThreeDReal,%new'rm
ThreeDReal,%new'sigdot
ThreeDReal,%new'up
ThreeDReal,%new'tp
ThreeDReal,%new'rmp
ThreeDReal,%new'vp
real,%new'eke
real,%new'cks
real,%new'trhat
real,%new'pe
real,%new'psbar
real,%new'vromg
)
let

```

```

ks:=15;
ii:=65;
jl:=40;
i1:=1;
j1:=1;
i2:=i1+1;
i3:=i1+2;
i4:=i1+3;
ilm:=i1-1;
j2:=j1+1;
j3:=j1+2;
j4:=j1+3;
jlm:=j1-1;
kzm1:=ks-1;
kzp1:=kz+1;
cp:=1.00464e7;
g:=980.6;
hl:=2.501e10;
pbar:=1.e6;
rv:=2.87e6;
rv:=4.61e+6;

dt:=
if (ktau = 1) then
  dt
else
  2.0 *dt
end if;

new'rm, rmpr:=
for k in 1,kz cross i in i1,il cross j in j1,jl
  t'rm,
  t'rmpr:=
  if (rm[k,i,j] > 0.0) then
    rm[k,i,j],
    rm[k,i,j] / (ps[i,j]+pbar)
  else
    0.0,
    0.0
  end if
returns
array of t'rm
array of t'rmpr
end for;

dmonp:=0.0;
emonp:=0.0;

new'ip, new'up, new'vp, new'rmp, new'sigdot,
new'cke, new'cks, new'that,
new'pe, new'psbar, new'romg:=
for i in i1,il cross j in j1,jl

  psijc:=ps[i,j]+pbar;
  psijci:=1.0/psijc;
  corf1,corf2:=

if boundary'cell(i,il,il,j,j1,jl) then
  0.0,0.0
else
  0.125*(corp[i,j]+corp[i+1,j]),
  0.125*(corp[i,j]+corp[i,j+1])
end if;

emthad := emsq[i,j]*psijci*tdei;
em2tps := emthad*psijci*r / cp;

emhad1,emhad2:=
if boundary'cell(i,il,il,j,j1,jl) then
  0.0,0.0
else
  0.25*tdei*(em[i,j]+em[i+1,j]),
  0.25*tdei*(em[i,j]+em[i,j+1])
end if;

amonp:=emonp; % 0.0 then cycle'emomp
bmonp:=dmonp; % 0.0 then cycle'dmonp

cycle'dmonp, fmonp:=
if boundary'cell(i,il,il,j,j1,jl) then
  0.0,0.0
else
  em[i+1,j]/(ps[i+1,j]+pbar),
  em[i,j+1]/(ps[i,j+1]+pbar)
end if;

new'bmonp:=
if (i = i2) & (^((j = j1)---(j = jl))) then
  em[i,j]*psijci
else
  cycle'dmonp
end if;

new'amonp:=
if (i = i2) & (^((j = j1)---(j = jl))) then
  0.25*(new'bmonp+fmonp+em[i-1,j] / (ps[i-1,j]+pbar)
  +em[i-1,j+1] / (ps[i-1,j+1]+pbar))
else
  emonp
end if;

cmonp:=new'bmonp+cycle'dmonp;
cycle'emomp, new'cmonp:=
if boundary'cell(i,il,il,j,j1,jl) then

```

```

0.0, cmonp
else
  0.25*(cmonp+em[i+1,j+1]/(ps[i+1,j+1]+pbar)+fmonp),
  0.25*(cmonp+em[i+1,j-1] / (ps[i+1,j-1]+pbar)
  +em[i,j-1] / (ps[i,j-1]+pbar))
end if;

pae, psn:=
if boundary'cell(i,il,il,j,j1,jl) then
  0.0, 0.0
else
  0.5*(ps[i,j]+ps[i+1,j])+pbar,
  0.5*(ps[i,j]+ps[i,j+1])+pbar
end if;

%
% extra variables for evaluating p.grad terms logarithmically
%
parmi, psrmj, psldi, psldj, zspdi, zspdj, emudsi, emvdsi, emrdsi:=
if boundary'cell(i,il,il,j,j1,jl) then
  0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0
else
  ps[i+1,j]-ps[i,j]-psm[i+1,j]+psm[i,j],
  ps[i,j+1]-ps[i,j]-psm[i,j+1]+psm[i,j],
  log(ps[i+1,j]+pbar)-log(psijc),
  log(ps[i,j+1]+pbar)-log(psijc),
  pbar*(sz[i+1,j]-sz[i,j]),
  pbar*(sz[i,j+1]-sz[i,j]),
  emsq[i,j]/(4.0*ds*pae),
  emsq[i,j]/(4.0*ds*psn),
  emsq[i,j]/(4.0*ds*psijc)
end if;

%
% compute total divergence
%
sumu, sumv, sumx, vadvu, vadvv, vadvr, wvel, flev:=
divergence'sums(i,j,kz,il,il,j1,jl,dsi,dq,u,v,t,emsq);

sigdot'k:=
for k in 1,kz
returns array of (
  if (k = 1) then
    0.0
  else
    wvel[k-1]-qph[k-1]*wvel[kz]
  end if)
end for;

vadvr'm'k, vadv'u'k, vadv'v'k:=
for l in 1,kz
  ll:=l+1;
  t'vadvr, t'vadvu, t'vadvv :=
  if (l = kz) then
    0.0, 0.0, 0.0
  else
    emrdsi*(qph[ll]*sumx-vadvr[ll])
    *(new'rm[ll,i,j]+new'rm[l,i,j]
    +2.0*sqrt(new'rm[ll,i,j]*new'rm[l,i,j])),
    emudsi*(qph[ll]*sumu-vadvu[ll])
    *(u[ll,i,j]+u[l,i,j]),
    emvdsi*(qph[ll]*sumv-vadvv[ll])
    *(v[ll,i,j]+v[l,i,j])
  end if
returns
array of t'vadvr
array of t'vadvu
array of t'vadvv
end for;

%
% set up temperature difference terms
%
dtfdq:=
for k in 1, kz
returns array of (
  if ((k = 1)---boundary'cell(i,il,il,j,j1,jl)) then
    0.0
  else
    if (k = kz) then
      dtmax*(flev[kz]-flev[kzm1])+dtodq[kz]
    else
      0.5*(flev[k+1]-flev[k-1]) / dq[k]+dtodq[k]
    end if
  end if)
end for;

psmue, psmuw, psmun, psmus, psmu, psmvn, psmv, psmvw, psmv:=
if ((j = j2)---boundary'cell(i,il,il,j,j1,jl)) then
  0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0
else
  if (i = ilm) then
    1.5*psm[i,j]-0.5*psm[ilm,j]+pbar
  else
    0.5*(psm[i+1,j]+psm[i+2,j])+pbar
  end if;

  0.5*(psm[i-1,j]+psm[i,j])+pbar,
  0.5*(psm[i,j+1]+psm[i+1,j+1])+pbar,
  0.5*(psm[i,j-1]+psm[i-1,j-1])+pbar,
  0.5*(psm[i,j]+psm[i+1,j])+pbar,

if (j = jlm) then
  1.5*psm[i,j]-0.5*psm[i,jlm]+pbar
else
  0.5*(psm[i,j+1]+psm[i,j+2])+pbar
end if;

```

```

0.5*(psm[i,j-1]+psm[i,j])+pbar,
0.5*(psm[i+1,j]+psm[i+1,j+1])+pbar,
0.5*(psm[i-1,j]+psm[i-1,j+1])+pbar,
0.5*(psm[i,j]+psm[i,j+1])+pbar
end if;
%
% commence vertical level loop
%
tp'k, up'k, vp'k, rmp'k, omega'k,
new'eke, new'ppe, new'pvromg, new'ptrhat:=
for k in 1,kz
%
% compute vertical advection contribs. in rhs of mtm. = ns.
%
% compute horizontal advection terms associated with rhs of mtm. = ns.
%
up'k:=
if ((j = j2)—boundary'cell(i,i1,i1,j1,j1)) then
up[k,i,j]
else
let
vat1:=
if (k = kz) then
0.0 %gke
else
(vadvu[k+1]-vadvu[k])/dq[k]
end if;
ub:=u[k,i,j]+u[k,i-1,j];
uc:=u[k,i,j]+u[k,i,j-1];
ud:=u[k,i,j]+u[k,i+1,j];
ue:=u[k,i,j]+u[k,i,j+1];
vb:=v[k,i,j]+v[k,i,j-1];
vc:=v[k,i,j]+v[k,i+1,j];
vd:=v[k,i,j]+v[k,i,j+1];
hadv1:=emhad1*( ud*ud*cyclc'dmonp-ub*ub*new'bmonp
+ue*ue*cyclc'emonp-uc*uc*new'cmonp);
%
% compute pressure gradient terms on rhs of mtm. = ns.
% logarithmically
pg1:= ((pse-pbar)*(phi[k,i+1,j]-phi[k,i,j])
+sspdi-ribar[k]*psrmi
+psc*psldi*( r*0.5*( t[k,i+1,j]+t[k,i,j])
+rtbar[k]))*dsi;
ct1:= corf1*(vc+ve);
ubdiff:=
if (dnorm[k] = 0.0) then
0.0
else
cdiff[i,j]*dnorm[k]*dsi2
*( um[k,i+1,j]/psmue+um[k,i-1,j]/psmuw
+um[k,i,j+1]/psmun+um[k,i,j-1]/psmus
-4.0*um[k,i,j]/psmu )*psmu
end if;
in
(ct1+vad1-hadv1-pg1+ubdiff)*dt+um[k,i,j]
end let
end if;
%
t'rmv'k, tp'k, omega'k:=
if ((i = i2)—boundary'cell(i,i1,i1,j1,j1)) then
rmp[k,i,j],
tp[k,i,j],
omega[k]
else
%
% calculate horizontal advection term in the temp. = ation
let
wvelav:=
if (k > 1) then
0.5*(wvel[k-1]+wvel[k])
else
0.5*wvel[1]
end if;
thadv1:= u[k,i,j]*(t[k,i+1,j]-t[k,i,j])
+u[k,i-1,j]*(t[k,i,j]-t[k,i-1,j]);
thadv2:= v[k,i,j]*(t[k,i,j+1]-t[k,i,j])
+v[k,i,j-1]*(t[k,i,j]-t[k,i,j-1]);
thadv:= emthadv*(thadv1+thadv2);
tfull:= tflev[k]+tbar[k];
phadv1:= u[k,i,j]*(ps[i+1,j]-ps[i,j])
+u[k,i-1,j]*(ps[i,j]-ps[i-1,j]);
phadv2:= v[k,i,j]*(ps[i,j+1]-ps[i,j])
+v[k,i,j-1]*(ps[i,j]-ps[i,j-1]);
t'omg:= em2tps*(phadv1+phadv2);
tpstar:= t'omg*tfull;
omg:= wvelav+q[k]*psijc*t'omg*cp / r;
gamapr:= (r / cp)*tfull / q[k]-dtfdq[k];
thdiff:=
if (dnorm[k] = 0.0) then
0.0
else
cdiff[i,j]*dnorm[k]*dsi2
*( tm[k,i+1,j]+tm[k,i-1,j]
+tm[k,i,j+1]+tm[k,i,j-1]-4.0*tm[k,i,j])
end if;
tp'k:= (tpstar-thadv+thdiff
+(wvelav*gamapr+q[k]*dtfdq[k]*wvel[kz])*psijc
-(wvelav*gamapr+q[k]*dtodq[k]*wvel[kz]) / pbar
)*dt+tm[k,i,j];
%
% moisture
rme:=rmpv[k,i,j]+rmpv[k,i+1,j];
rmw:=rmpv[k,i,j]+rmpv[k,i-1,j];
rma:=rmpv[k,i,j]+rmpv[k,i,j+1];
rmb:=rmpv[k,i,j]+rmpv[k,i,j-1];
rmvad:=
if (k = kz) then
0.0 %gke
else
-(vadvr[k+1]-vadvr[k]) / dq[k]
end if;
rmhad:=emsq[i,j]*tdsi*( u[k,i,j]*rme-u[k,i-1,j]*rmw
+v[k,i,j]*rma-v[k,i,j-1]*rmb);
rmme:= rmm[k,i+1,j]/(psm[i+1,j]+pbar);
rmmw:= rmm[k,i-1,j]/(psm[i-1,j]+pbar);
rmma:= rmm[k,i,j+1]/(psm[i,j+1]+pbar);
rmmj:= rmm[k,i,j-1]/(psm[i,j-1]+pbar);
rmdif:=
if (dnorm[k] = 0.0) then
0.0
else
cdiff[i,j]*dnorm[k]*dsi2
*(rmme+rmmw+rmma+rmmj-4.0*rmmj)
*(psm[i,j]+pbar)
end if;
t'rmv'k:= rmm[k,i,j]+dtt*(rmhad+rmvad+rmhdif);
%
% suppress negative mixing ratios
%
rmp'k:=
if (t'rmv'k < 1.0E-20) then
0.0
else
t'rmv'k
end if;
in
rmp'k,
tp'k,
omg
end let
end if;
%
vp'k,
new'eke'k,
new'ppe'k,
new'pvromg'k:=
if ((i = i2)—boundary'cell(i,i1,i1,j1,j1)) then
vp[k,i,j],
0.0, %eke
0.0, %ppe
0.0 %pvromg
else
let
ua:=u[k,i-1,j]+u[k,i-1,j+1];
ue:=u[k,i,j]+u[k,i,j+1];
va:=v[k,i,j]+v[k,i-1,j];
vb:=v[k,i,j]+v[k,i,j-1];
ve:=v[k,i,j]+v[k,i+1,j];
vf:=v[k,i,j]+v[k,i,j+1];
%
% calculate v velocity component logarithmically
ct2:= corf2*(ua+ue);
hadv2:= emhad2*( ue*ve*cyclc'emonp-ua*va*new'amonp
+vf*vf*fmop-vb*vb*new'bmonp);
pg2:= ((psn-pbar)*(phi[k,i,j+1]-phi[k,i,j])
+spdj-rtbar[k]*psrmi
+psn*psldj*( r*0.5*( t[k,i,j+1]+t[k,i,j])
+rtbar[k]))*dsi;
vbdiff:=
if (dnorm[k] = 0.0) then
0.0
else
cdiff[i,j]*dnorm[k]*dsi2
*( vm[k,i+1,j]/psmve+vm[k,i-1,j]/psmvw
+vm[k,i,j+1]/psmvn+vm[k,i,j-1]/psmvs
-4.0*vm[k,i,j]/psmv)*psmv
end if;
vat2:=
if (k = kz) then
0.0 %gke
else
-(vadvv[k+1]-vadvv[k])/dq[k]
end if;
t'vp:= (ct2+vad2-hadv2-pg2+vbdiff)*dt+vm[k,i,j];
%
% calculate contributions to integrals
in
t'vp,
dq[k]*psijc*( u[k,i,j]*u[k,i,j]+ v[k,i,j]*v[k,i,j]),
tflev[k]*dq[k],
(omega'k*omega'k)*dq[k]
end let
end if;
returna
array of tp'k
array of up'k
array of vp'k
array of t'rmv'k
array of omega'k

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value of sum (dq[k]*psijc*(up'k*up'k+vp'k*vp'k)) %new'cke'k
value of sum (tlev[k]*dq[k]) %new'ppe'k
value of sum (omega'k*omega'k*dq[k]) %new'pvromg'k
value of sum (t'rm'k*dq[k])%ptrhat
end for; % k
t'new'ptrhat := new'ptrhat+emsqi[i,j];
t'new'epe := new'ppe*psijc*emsqi[i,j];
t'new'vromg := new'pvromg*emsqi[i,j];
return
array of tp'k
array of up'k
array of vp'k
array of rmp'k
array of sigdot'k
value of sum new'cke
value of sum (emsqi[i,j]) %new'cks
value of sum (t'new'ptrhat*emsqi[i,j])
value of sum t'new'epe
value of sum ((psijc-0.988e6)*emsqi[i,j]) % psbar
value of sum t'new'vromg
end for % i,j
in
new'rm,
new'sigdot,
new'up,
new'vp,
new'rmp,
new'vp,
new'cke,
new'cks,
new'ptrhat,
new'pe,
new'psbar,
new'romg
end let
end function

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