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c-----
      subroutine GPPUSHF2L(ppart,fxy,kpic,ncl,ihole,qbm,dt,ek,idimp,
     1nppmx,nx,ny,mx,my,nxv,nyv,mx1,mxy1,ntmax,irc)
c for 2d code, this subroutine updates particle co-ordinates and
c velocities using leap-frog scheme in time and first-order linear
c interpolation in space, with periodic boundary conditions.
c also determines list of particles which are leaving this tile
c OpenMP version using guard cells
c data read in tiles
c particles stored segmented array
c 44 flops/particle, 12 loads, 4 stores
c input: all except ncl, ihole, irc, output: ppart, ncl, ihole, ek, irc
c equations used are:
c vx(t+dt/2) = vx(t-dt/2) + (q/m)*fx(x(t),y(t))*dt,
c vy(t+dt/2) = vy(t-dt/2) + (q/m)*fy(x(t),y(t))*dt,
c where q/m is charge/mass, and
c x(t+dt) = x(t) + vx(t+dt/2)*dt, y(t+dt) = y(t) + vy(t+dt/2)*dt
c fx(x(t),y(t)) and fy(x(t),y(t)) are approximated by interpolation from
c the nearest grid points:
c fx(x,y) = (1-dy)*((1-dx)*fx(n,m)+dx*fx(n+1,m)) + dy*((1-dx)*fx(n,m+1)
c   + dx*fx(n+1,m+1))
c fy(x,y) = (1-dy)*((1-dx)*fy(n,m)+dx*fy(n+1,m)) + dy*((1-dx)*fy(n,m+1)
c   + dx*fy(n+1,m+1))
c where n,m = leftmost grid points and dx = x-n, dy = y-m
c ppart(1,n,m) = position x of particle n in tile m
c ppart(2,n,m) = position y of particle n in tile m
c ppart(3,n,m) = velocity vx of particle n in tile m
c ppart(4,n,m) = velocity vy of particle n in tile m
c fxy(1,j,k) = x component of force/charge at grid (j,k)
c fxy(2,j,k) = y component of force/charge at grid (j,k)
c that is, convolution of electric field over particle shape
c kpic(k) = number of particles in tile k
c ncl(i,k) = number of particles going to destination i, tile k
c ihole(1,:,k) = location of hole in array left by departing particle
c ihole(2,:,k) = destination of particle leaving hole
c ihole(1,1,k) = ih, number of holes left (error, if negative)
c qbm = particle charge/mass
c dt = time interval between successive calculations
c kinetic energy/mass at time t is also calculated, using
c ek = .125*sum((vx(t+dt/2)+vx(t-dt/2))**2+(vy(t+dt/2)+vy(t-dt/2))**2)
c idimp = size of phase space = 4
c nppmx = maximum number of particles in tile
c nx/ny = system length in x/y direction
c mx/my = number of grids in sorting cell in x/y
c nxv = first dimension of field arrays, must be >= nx+1
c nyv = second dimension of field arrays, must be >= ny+1
c mx1 = (system length in x direction - 1)/mx + 1
c mxy1 = mx1*my1, where my1 = (system length in y direction - 1)/my + 1
c ntmax = size of hole array for particles leaving tiles
c irc = maximum overflow, returned only if error occurs, when irc > 0
c optimized version
      implicit none
      integer idimp, nppmx, nx, ny, mx, my, nxv, nyv, mx1, mxy1, ntmax
      integer irc

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real qbm, dt, ek
real ppart, fxy
integer kpic, ncl, ihole
dimension ppart(idimp,npptmx,mxy1), fxy(2,nxv,nyv)
dimension kpic(mxy1), ncl(8,mxy1)
dimension ihole(2,ntmax+1,mxy1)
c local data
integer MXV, MYV
parameter(MXV=33,MYV=33)
integer noff, moff, npp
integer i, j, k, ih, nh, nn, mm
real qtm, dxp, dyp, amx, amy
real x, y, dx, dy, vx, vy
real anx, any, edgelx, edgely, edgerx, edgery
real sfxy
dimension sfxy(2,MXV,MYV)
c dimension sfxy(2, mx+1, my+1)
double precision sum1, sum2
qtm = qbm*dt
anx = real(nx)
any = real(ny)
sum2 = 0.0d0
c error if local array is too small
c if ((mx.ge.MXV).or.(my.ge.MYV)) return
c loop over tiles
!$OMP PARALLEL DO
!$OMP& PRIVATE(i,j,k,noff,moff,npp,nn,mm,ih,nh,x,y,dxp,dyp,amx,amy,dx,dy
!$OMP& ,vx,vy,edgelx,edgely,edgerx,edgery,sum1,sfxy)
!$OMP& REDUCTION(:sum2)
do 50 k = 1, mxy1
noff = (k - 1)/mx1
moff = my*noff
noff = mx*(k - mx1*noff - 1)
npp = kpic(k)
nn = min(mx,nx-noff)
mm = min(my,ny-moff)
edgelx = noff
edgerx = noff + nn
edgely = moff
edgery = moff + mm
ih = 0
nh = 0
c load local fields from global array
do 20 j = 1, mm+1
do 10 i = 1, nn+1
sfxy(1,i,j) = fxy(1,i+noff,j+moff)
sfxy(2,i,j) = fxy(2,i+noff,j+moff)
10 continue
20 continue
c clear counters
do 30 j = 1, 8
ncl(j,k) = 0
30 continue
sum1 = 0.0d0

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c loop over particles in tile
do 40 j = 1, npp
c find interpolation weights
  x = ppart(1,j,k)
  y = ppart(2,j,k)
  nn = x
  mm = y
  dxp = x - real(nn)
  dyp = y - real(mm)
  nn = nn - noff + 1
  mm = mm - moff + 1
  amx = 1.0 - dxp
  amy = 1.0 - dyp
c find acceleration
  dx = amx*sfx(1,nn,mm)
  dy = amx*sfx(2,nn,mm)
  dx = amy*(dxp*sfx(1,nn+1,mm) + dx)
  dy = amy*(dyp*sfx(2,nn+1,mm) + dy)
  vx = amx*sfx(1,nn,mm+1)
  vy = amx*sfx(2,nn,mm+1)
  dx = dx + dyp*(dxp*sfx(1,nn+1,mm+1) + vx)
  dy = dy + dyp*(dyp*sfx(2,nn+1,mm+1) + vy)
c new velocity
  vx = ppart(3,j,k)
  vy = ppart(4,j,k)
  dx = vx + qtm*dx
  dy = vy + qtm*dy
c average kinetic energy
  vx = vx + dx
  vy = vy + dy
  sum1 = sum1 + (vx*vx + vy*vy)
  ppart(3,j,k) = dx
  ppart(4,j,k) = dy
c new position
  dx = x + dx*dt
  dy = y + dy*dt
c find particles going out of bounds
  mm = 0
c count how many particles are going in each direction in ncl
c save their address and destination in ihole
c use periodic boundary conditions and check for roundoff error
c mm = direction particle is going
  if (dx.ge.edgerx) then
    if (dx.ge.anx) dx = dx - anx
    mm = 2
  else if (dx.lt.edgelx) then
    if (dx.lt.0.0) then
      dx = dx + anx
      if (dx.lt.anx) then
        mm = 1
      else
        dx = 0.0
      endif
    else

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        mm = 1
    endif
endif
if (dy.ge.edgery) then
    if (dy.ge.any) dy = dy - any
    mm = mm + 6
else if (dy.lt.edgely) then
    if (dy.lt.0.0) then
        dy = dy + any
        if (dy.lt.any) then
            mm = mm + 3
        else
            dy = 0.0
        endif
    else
        mm = mm + 3
    endif
endif
c set new position
ppart(1,j,k) = dx
ppart(2,j,k) = dy
c increment counters
if (mm.gt.0) then
    ncl(mm,k) = ncl(mm,k) + 1
    ih = ih + 1
    if (ih.le.ntmax) then
        ihole(1,ih+1,k) = j
        ihole(2,ih+1,k) = mm
    else
        nh = 1
    endif
endif
sum2 = sum2 + sum1
40 continue
c set error and end of file flag
c ihole overflow
if (nh.gt.0) then
    irc = ih
    ih = -ih
endif
ihole(1,1,k) = ih
50 continue
!$OMP END PARALLEL DO
c normalize kinetic energy
ek = ek + 0.125*sum2
return
end
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c-----
      subroutine GPPOST2L(ppart,q,kpic,qm,nppmx,idimp,mx,my,nxv,nyv,mx1,
     1mxy1)
c for 2d code, this subroutine calculates particle charge density
c using first-order linear interpolation, periodic boundaries
c OpenM version using guard cells
c data deposited in tiles
c particles stored segmented array
c 17 flops/particle, 6 loads, 4 stores
c input: all, output: q
c charge density is approximated by values at the nearest grid points
c q(n,m)=qm*(1.-dx)*(1.-dy)
c q(n+1,m)=qm*dx*(1.-dy)
c q(n,m+1)=qm*(1.-dx)*dy
c q(n+1,m+1)=qm*dx*dy
c where n,m = leftmost grid points and dx = x-n, dy = y-m
c ppart(1,n,m) = position x of particle n in tile m
c ppart(2,n,m) = position y of particle n in tile m
c q(j,k) = charge density at grid point j,k
c kpic = number of particles per tile
c qm = charge on particle, in units of e
c nppmx = maximum number of particles in tile
c idimp = size of phase space = 4
c mx/my = number of grids in sorting cell in x/y
c nxv = first dimension of charge array, must be >= nx+1
c nyv = second dimension of charge array, must be >= ny+1
c mx1 = (system length in x direction - 1)/mx + 1
c mxy1 = mx1*my1, where my1 = (system length in y direction - 1)/my + 1
      implicit none
      integer nppmx, idimp, mx, my, nxv, nyv, mx1, mxy1
      real qm
      real ppart, q
      integer kpic
      dimension ppart(idimp,nppmx,mxy1), q(nxv,nyv)
      dimension kpic(mxy1)
c local data
      integer MXV, MYV
      parameter(MXV=33,MYV=33)
      integer noff, moff, npp
      integer i, j, k, nn, mm
      real x, y, dxp, dyp, amx, amy
      real sq
c      dimension sq(MXV,MYV)
      dimension sq(mx+1,my+1)
c error if local array is too small
c      if ((mx.ge.MXV).or.(my.ge.MYV)) return
c loop over tiles
!$OMP PARALLEL DO
!$OMP& PRIVATE(i,j,k,noff,moff,npp,nn,mm,x,y,dxp,dyp,amx,amy,sq)
      do 80 k = 1, mxy1
      noff = (k - 1)/mx1
      moff = my*noff
      noff = mx*(k - mx1*noff - 1)
      npp = kpic(k)

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c zero out local accumulator
do 20 j = 1, my+1
do 10 i = 1, mx+1
sq(i,j) = 0.0
10 continue
20 continue
c loop over particles in tile
do 30 j = 1, npp
c find interpolation weights
x = ppart(1,j,k)
y = ppart(2,j,k)
nn = x
mm = y
dxp = qm*(x - real(nn))
dyp = y - real(mm)
nn = nn - noff + 1
mm = mm - moff + 1
amx = qm - dxp
amy = 1.0 - dyp
c deposit charge within tile to local accumulator
x = sq(nn,mm) + amx*amy
y = sq(nn+1,mm) + dxp*amy
sq(nn,mm) = x
sq(nn+1,mm) = y
x = sq(nn,mm+1) + amx*dyp
y = sq(nn+1,mm+1) + dxp*dyp
sq(nn,mm+1) = x
sq(nn+1,mm+1) = y
30 continue
c deposit charge to interior points in global array
nn = min(mx,nxv-noff)
mm = min(my,nyv-moff)
do 50 j = 2, mm
do 40 i = 2, nn
q(i+noff,j+moff) = q(i+noff,j+moff) + sq(i,j)
40 continue
50 continue
c deposit charge to edge points in global array
mm = min(my+1,nyv-moff)
do 60 i = 2, nn
!$OMP ATOMIC
q(i+noff,1+moff) = q(i+noff,1+moff) + sq(i,1)
if (mm > my) then
!$OMP ATOMIC
q(i+noff,mm+moff) = q(i+noff,mm+moff) + sq(i,mm)
endif
60 continue
nn = min(mx+1,nxv-noff)
do 70 j = 1, mm
!$OMP ATOMIC
q(1+noff,j+moff) = q(1+noff,j+moff) + sq(1,j)
if (nn > mx) then
!$OMP ATOMIC
q(nn+noff,j+moff) = q(nn+noff,j+moff) + sq(nn,j)

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    endif
70 continue
80 continue
!$OMP END PARALLEL DO
      return
end
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