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      attributes(global) subroutine gpuppush2l(ppart, fxy, kplic, qbm, dt, ek, &
      &idimp, nppmx, nx, ny, mx, my, nxv, nyv, mx1, mxy1, ipbc)
! for 2d code, this subroutine updates particle co-ordinates and
! velocities using leap-frog scheme in time and first-order linear
! interpolation in space, with various boundary conditions.
! threaded version using guard cells
! data read in tiles
! particles stored segmented array
! 44 flops/particle, 12 loads, 4 stores
! input: all, output: ppart, ek
! equations used are:
!  $vx(t+dt/2) = vx(t-dt/2) + (q/m)*fx(x(t),y(t))*dt,$ 
!  $vy(t+dt/2) = vy(t-dt/2) + (q/m)*fy(x(t),y(t))*dt,$ 
! where  $q/m$  is charge/mass, and
!  $x(t+dt) = x(t) + vx(t+dt/2)*dt,$   $y(t+dt) = y(t) + vy(t+dt/2)*dt$ 
!  $fx(x(t),y(t))$  and  $fy(x(t),y(t))$  are approximated by interpolation from
! the nearest grid points:
!  $fx(x,y) = (1-dy)*((1-dx)*fx(n,m)+dx*fx(n+1,m)) + dy*((1-dx)*fx(n,m+1)$ 
!  $+ dx*fx(n+1,m+1))$ 
!  $fy(x,y) = (1-dy)*((1-dx)*fy(n,m)+dx*fy(n+1,m)) + dy*((1-dx)*fy(n,m+1)$ 
!  $+ dx*fy(n+1,m+1))$ 
! where  $n,m$  = leftmost grid points and  $dx = x-n,$   $dy = y-m$ 
! ppart(n,1,m) = position x of particle n in tile m
! ppart(n,2,m) = position y of particle n in tile m
! ppart(n,3,m) = velocity vx of particle n in tile m
! ppart(n,4,m) = velocity vy of particle n in tile m
! fxy(1,j,k) = x component of force/charge at grid (j,k)
! fxy(2,j,k) = y component of force/charge at grid (j,k)
! that is, convolution of electric field over particle shape
! kpic = number of particles per tile
! qbm = particle charge/mass
! dt = time interval between successive calculations
! kinetic energy/mass at time t is also calculated, using
!  $ek = .125*sum((vx(t+dt/2)+vx(t-dt/2))^2+(vy(t+dt/2)+vy(t-dt/2))^2)$ 
! idimp = size of phase space = 4
! nppmx = maximum number of particles in tile
! nx/ny = system length in x/y direction
! mx/my = number of grids in sorting cell in x/y
! nxv = first dimension of field arrays, must be  $\geq nx+1$ 
! nyv = second dimension of field arrays, must be  $\geq ny+1$ 
! mx1 = (system length in x direction - 1)/mx + 1
! mxy1 = mx1*my1, where my1 = (system length in y direction - 1)/my + 1
! ipbc = particle boundary condition = (0,1,2,3) =
! (none,2d periodic,2d reflecting,mixed reflecting/periodic)
      implicit none
      integer, value :: idimp, nppmx, nx, ny, mx, my, nxv, nyv
      integer, value :: mx1, mxy1, ipbc
      real, value :: qbm, dt
      real, dimension(nppmx,idimp,mxy1) :: ppart
      real, dimension(2,nxv,nyv) :: fxy
      integer, dimension(mxy1) :: kplic
      real, dimension(mxy1) :: ek
! local data

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integer :: noff, moff, npp, mxv
integer :: i, j, k, ii, nn, mm
real :: qtm, edgelx, edgely, edgerx, edgerly
real :: dxp, dyp, amx, amy, x, y, dx, dy, vx, vy
! The sizes of the shared memory arrays are as follows:
! float sfx(2*(mx+1)*(my+1)), sek(blockDim%x)
! to conserve memory, sek overlaps with sfx
! and the name sfx is used instead of sek
  real, shared, dimension(*) :: sfx
  double precision :: sum1
  qtm = qbm*dt
  sum1 = 0.0d0
! set boundary values
  edgelx = 0.0
  edgely = 0.0
  edgerx = real(nx)
  edgerly = real(ny)
  if (ipbc==2) then
    edgelx = 1.0
    edgely = 1.0
    edgerx = real(nx-1)
    edgerly = real(ny-1)
  else if (ipbc==3) then
    edgelx = 1.0
    edgerx = real(nx-1)
  endif
  mxv = mx + 1
! k = tile number
  k = blockIdx%x + blockDim*x*(blockIdx%y - 1)
! loop over tiles
  if (k <= mxy1) then
    noff = (k - 1)/mx1
    moff = my*noff
    noff = mx*(k - mx1*noff - 1)
    npp = kp1c(k)
! load local fields from global array
    nn = min(mx,nx-noff) + 1
    mm = min(my,ny-moff) + 1
    ii = threadIdx%x
    do while (ii <= mxv*(my+1))
      j = (ii - 1)/mxv
      i = ii - mxv*j
      j = j + 1
      if ((i <= nn) .and. (j <= mm)) then
        sfx(2*ii-1) = fxy(1,i+noff,j+moff)
        sfx(2*ii) = fxy(2,i+noff,j+moff)
      endif
      ii = ii + blockDim%x
    enddo
! synchronize threads
    call syncthreads()
! loop over particles in tile
    j = threadIdx%x
    do while (j <= npp)

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! find interpolation weights
    x = ppart(j,1,k)
    nn = x
    y = ppart(j,2,k)
    mm = y
    dxp = x - real(nn)
    dyp = y - real(mm)
    nn = 2*(nn - noff) + 2*mxv*(mm - moff) + 1
    amx = 1.0 - dxp
    amy = 1.0 - dyp
! find acceleration
    dx = amx*sfxxy(nn)
    dy = amx*sfxxy(nn+1)
    dx = amy*(dxp*sfxxy(nn+2) + dx)
    dy = amy*(dyp*sfxxy(nn+3) + dy)
    nn = nn + 2*mxv
    vx = amx*sfxxy(nn)
    vy = amx*sfxxy(nn+1)
    dx = dx + dyp*(dxp*sfxxy(nn+2) + vx)
    dy = dy + dyp*(dyp*sfxxy(nn+3) + vy)
! new velocity
    vx = ppart(j,3,k)
    vy = ppart(j,4,k)
    dx = vx + qtm*dx
    dy = vy + qtm*dy
! average kinetic energy
    vx = vx + dx
    vy = vy + dy
    sum1 = sum1 + dble(vx*vx + vy*vy)
    ppart(j,3,k) = dx
    ppart(j,4,k) = dy
! new position
    dx = x + dx*dt
    dy = y + dy*dt
! reflecting boundary conditions
    if (ipbc==2) then
        if ((dx < edgelx).or.(dx >= edgerx)) then
            dx = ppart(j,1,k)
            ppart(j,3,k) = -ppart(j,3,k)
        endif
        if ((dy < edgely).or.(dy >= edgery)) then
            dy = ppart(j,2,k)
            ppart(j,4,k) = -ppart(j,4,k)
        endif
    endif
! mixed reflecting/periodic boundary conditions
    else if (ipbc==3) then
        if ((dx < edgelx).or.(dx >= edgerx)) then
            dx = ppart(j,1,k)
            ppart(j,3,k) = -ppart(j,3,k)
        endif
    endif
! set new position
    ppart(j,1,k) = dx
    ppart(j,2,k) = dy

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        j = j + blockDim%x
    enddo
! synchronize threads
    call syncthreads()
! add kinetic energies in tile
    sfxxy(threadIdx%x) = real(sum1)
! synchronize threads
    call syncthreads()
    call lsum2(sfxxy,blockDim%x)
! normalize kinetic energy of tile
    if (threadIdx%x==1) ek(k) = 0.125*sfxxy(1)
endif
end subroutine
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!-----
      attributes(global) subroutine gpu2ppost2l(ppart,q,kpic,qm,nppmx, &
      &idimp,mx,my,nxv,nyv,mx1,mxy1)
! for 2d code, this subroutine calculates particle charge density
! using first-order linear interpolation, periodic boundaries
! threaded version using guard cells
! data deposited in tiles
! particles stored segmented array
! 17 flops/particle, 6 loads, 4 stores
! input: all, output: q
! charge density is approximated by values at the nearest grid points
!  $q(n,m)=qm*(1.-dx)*(1.-dy)$ 
!  $q(n+1,m)=qm*dx*(1.-dy)$ 
!  $q(n,m+1)=qm*(1.-dx)*dy$ 
!  $q(n+1,m+1)=qm*dx*dy$ 
! where n,m = leftmost grid points and  $dx = x-n$ ,  $dy = y-m$ 
! ppart(n,1,m) = position x of particle n in tile m
! ppart(n,2,m) = position y of particle n in tile m
!  $q(j,k)$  = charge density at grid point j,k
! kpic = number of particles per tile
! qm = charge on particle, in units of e
! nppmx = maximum number of particles in tile
! idimp = size of phase space = 4
! mx/my = number of grids in sorting cell in x/y
! nxv = first dimension of charge array, must be  $\geq nx+1$ 
! nyv = second dimension of charge array, must be  $\geq ny+1$ 
! mx1 = (system length in x direction - 1)/mx + 1
! mxy1 = mx1*my1, where my1 = (system length in y direction - 1)/my + 1
      implicit none
      integer, value :: nppmx, idimp, mx, my, nxv, nyv, mx1, mxy1
      real, value :: qm
      real, dimension(nppmx,idimp,mxy1) :: ppart
      real, dimension(nxv,nyv) :: q
      integer, dimension(mxy1) :: kpic
! local data
      integer :: noff, moff, npp, mxv
      integer :: i, j, k, ii, nn, mm, np, mp
      real :: dxp, dyp, amx, amy, old
! The size of the shared memory array is as follows:
! real sq((mx+1)*(my+1))
      real, shared, dimension((mx+1)*(my+1)) :: sq
      mxv = mx + 1
! k = tile number
      k = blockIdx%x + gridDim%x*(blockIdx%y - 1)
! loop over tiles
      if (k <= mxy1) then
         noff = (k - 1)/mx1
         moff = my*noff
         noff = mx*(k - mx1*noff - 1)
         npp = kpic(k)
! zero out local accumulator
         i = threadIdx%x
         do while (i <= mxv*(my+1))
            sq(i) = 0.0

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        i = i + blockDim%x
    enddo
! synchronize threads
    call syncthreads()
! loop over particles in tile
    j = threadIdx%x
    do while (j <= npp)
! find interpolation weights
        dxp = ppart(j,1,k)
        nn = dxp
        dyp = ppart(j,2,k)
        mm = dyp
        dxp = qm*(dxp - real(nn))
        dyp = dyp - real(mm)
        nn = nn - noff + 1
        mm = mxv*(mm - moff)
        amx = qm - dxp
        mp = mm + mxv
        amy = 1.0 - dyp
        np = nn + 1
! deposit charge within tile to local accumulator
! original deposit charge, has data hazard on GPU
!         sq(np+mp) = sq(np+mp) + dxp*dyp
!         sq(nn+mp) = sq(nn+mp) + amx*dyp
!         sq(np+mm) = sq(np+mm) + dxp*amy
!         sq(nn+mm) = sq(nn+mm) + amx*amy
! for devices with compute capability 2.x
        old = atomicAdd(sq(np+mp),dxp*dyp)
        old = atomicAdd(sq(nn+mp),amx*dyp)
        old = atomicAdd(sq(np+mm),dxp*amy)
        old = atomicAdd(sq(nn+mm),amx*amy)
        j = j + blockDim%x
    enddo
! synchronize threads
    call syncthreads()
! deposit charge to global array
    nn = min(mxv,nxv-noff)
    mm = min(my+1,nyv-moff)
    ii = threadIdx%x
    do while (ii <= mxv*(my+1))
        j = (ii - 1)/mxv
        i = ii - mxv*j
        j = j + 1
        if ((i <= nn) .and. (j <= mm)) then
! original deposit charge, has data hazard on GPU
!         q(i+noff,j+moff) = q(i+noff,j+moff) + sq(ii)
! for devices with compute capability 2.x
            old = atomicAdd(q(i+noff,j+moff),sq(ii))
        endif
        ii = ii + blockDim%x
    enddo
endif
end subroutine

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