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! Skeleton 2-1/2D Electromagnetic MPI/OpenMP PIC code
! written by Viktor K. Decyk, UCLA
     program mpbpic2
     use mpbpush2_h
     use pplib2 h
     use omplib h
     implicit none
     integer, parameter :: indx = 9, indy =
     integer, parameter :: npx = 3072, npy =
                                                 3072
     integer, parameter :: ndim = 3
     real, parameter :: tend = 10.0, dt = 0.04, qme = -1.0
     real, parameter :: tend = 10.0, dt = 0.025, qme = -1.0
     real, parameter :: vtx = 1.0, vty = 1.0, vx0 = 0.0, vy0 = 0.0
     real, parameter :: vtz = 1.0, vz0 = 0.0
     real :: ax = .912871, ay = .912871, ci = 0.1
! idimp = dimension of phase space = 5
! relativity = (no, yes) = (0,1) = relativity is used
     integer :: idimp = 5, ipbc = 1, relativity = 1
! idps = number of partition boundaries
     integer :: idps = 2
     real :: wke = 0.0, we = 0.0, wf = 0.0, wm = 0.0, wt = 0.0
! sorting tiles, should be less than or equal to 32
     integer :: mx = 16, my = 16
! fraction of extra particles needed for particle management
     real :: xtras = 0.2
! declare scalars for standard code
     integer :: nx, ny, nxh, nyh, nxe, nye, nxeh, nnxe, nxyh, nxhy
     integer :: mx1, ntime, nloop, isign, ierr
     real :: qbme, affp, dth
     double precision :: np
! declare scalars for MPI code
     integer :: ntpose = 1
     integer :: nvp, idproc, kstrt, npmax, kxp, kyp, nypmx, nypmn
     integer :: nyp, noff, npp, nps, myp1, mxyp1
! declare scalars for OpenMP code
     integer :: nppmx, nppmx0, nbmaxp, ntmaxp, npbmx, irc
     integer :: nvpp
! declare arrays for standard code
     real, dimension(:,:), pointer :: part
     real, dimension(:,:), pointer :: qe
     real, dimension(:,:,:), pointer :: cue, fxyze, bxyze
     complex, dimension(:,:,:), pointer :: exyz, bxyz
     complex, dimension(:,:), pointer :: qt
     complex, dimension(:,:,:), pointer :: cut, fxyt, bxyt
     complex, dimension(:,:), pointer :: ffc
     integer, dimension(:), pointer :: mixup
     complex, dimension(:), pointer :: sct
     real, dimension(7) :: wtot, work
! declare arrays for MPI code
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complex, dimension(:,:,:), pointer :: bs, br
      real, dimension(:,:), pointer :: sbufl, sbufr, rbufl, rbufr
      real, dimension(:), pointer :: edges
      real, dimension(:), pointer :: scs, scr
! declare arrays for OpenMP code
     real, dimension(:,:,:), pointer :: ppart, ppbuff
      integer, dimension(:), pointer :: kpic
      integer, dimension(:,:), pointer :: ncl
      integer, dimension(:,:,:), pointer :: iholep
      integer, dimension(:,:), pointer :: ncll, nclr, mcll, mclr
! declare and initialize timing data
     real :: time
      integer, dimension(4) :: itime
     real :: tdpost = 0.0, tguard = 0.0, ttp = 0.0, tfield = 0.0
     real :: tdjpost = 0.0, tpush = 0.0, tsort = 0.0, tmov = 0.0
     real, dimension(2) :: tfft
     double precision :: dtime
!
     irc = 0
! nvpp = number of shared memory nodes (0=default)
     nvpp = 0
     write (*,*) 'enter number of nodes:'
     read (5,*) nvpp
! initialize for shared memory parallel processing
     call INIT_OMP(nvpp)
1
! initialize scalars for standard code
      np = dble(npx)*dble(npy)
     nx = 2**indx; ny = 2**indy; nxh = nx/2; nyh = ny/2
     nxe = nx + 2; nye = ny + 2; nxeh = nxe/2; nnxe = ndim*nxe
     nxyh = max(nx,ny)/2; nxhy = max(nxh,ny)
     mx1 = (nx - 1)/mx + 1
     nloop = tend/dt + .0001; ntime = 0
     qbme = qme
     affp = dble(nx)*dble(ny)/np
     dth = 0.0
! nvp = number of distributed memory nodes
! initialize for distributed memory parallel processing
     call PPINIT2(idproc, nvp)
     kstrt = idproc + 1
! check if too many processors
      if (nvp > ny) then
         if (kstrt==1) then
        write (*,*) 'Too many processors requested: ny, nvp=', ny, nvp
         endif
         go to 3000
      endif
! initialize data for MPI code
      allocate(edges(idps))
! calculate partition variables: edges, nyp, noff, nypmx
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! edges(1:2) = lower:upper boundary of particle partition
! nyp = number of primary (complete) gridpoints in particle partition
! noff = lowermost global gridpoint in particle partition
! nypmx = maximum size of particle partition, including guard cells
! nypmn = minimum value of nyp
     call PDICOMP2L(edges,nyp,noff,nypmx,nypmn,ny,kstrt,nvp,idps)
     if (nypmn < 1) then
         if (kstrt==1) then
           write (*,*) 'combination not supported nvp, ny =',nvp,ny
        go to 3000
     endif
!
! initialize additional scalars for MPI code
! kxp = number of complex grids in each field partition in x direction
     kxp = (nxh - 1)/nvp + 1
! kyp = number of complex grids in each field partition in y direction
     kyp = (ny - 1)/nvp + 1
! npmax = maximum number of electrons in each partition
     npmax = (np/nvp)*1.25
! myp1 = number of tiles in y direction
     myp1 = (nyp - 1)/my + 1; mxyp1 = mx1*myp1
! allocate and initialize data for standard code
     allocate(part(idimp,npmax))
     allocate(qe(nxe,nypmx),fxyze(ndim,nxe,nypmx))
     allocate(cue(ndim,nxe,nypmx),bxyze(ndim,nxe,nypmx))
     allocate(exyz(ndim,nye,kxp),bxyz(ndim,nye,kxp))
     allocate(qt(nye,kxp),fxyt(ndim,nye,kxp))
     allocate(cut(ndim,nye,kxp),bxyt(ndim,nye,kxp))
     allocate(ffc(nyh,kxp),mixup(nxhy),sct(nxyh))
     allocate(kpic(mxyp1))
! allocate and initialize data for MPI code
     allocate(bs(ndim,kxp,kyp),br(ndim,kxp,kyp))
     allocate(scs(ndim*nxe),scr(ndim*nxe))
! prepare fft tables
     call WPFFT2RINIT(mixup,sct,indx,indy,nxhy,nxyh)
! calculate form factors
     isign = 0
     call MPPOIS23(qt,fxyt,isign,ffc,ax,ay,affp,we,nx,ny,kstrt,nye,kxp,&
! initialize electrons
     nps = 1
     npp = 0
     call PDISTR2H(part,edges,npp,nps,vtx,vty,vtz,vx0,vy0,vz0,npx,npy, &
    &nx,ny,idimp,npmax,idps,ipbc,ierr)
! check for particle initialization error
     if (ierr /= 0) then
         if (kstrt==1) then
           write (*,*) 'particle initialization error: ierr=', ierr
        endif
         go to 3000
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endif
! initialize transverse electromagnetic fields
      exyz = cmplx(0.0,0.0)
      bxyz = cmplx(0.0,0.0)
!
! find number of particles in each of mx, my tiles: updates kpic, nppmx
      call PPDBLKP2L(part,kpic,npp,noff,nppmx,idimp,npmax,mx,my,mx1,
     &mxyp1,irc)
      if (irc /= 0) then
        write (*,*) 'PPDBLKP2L error, irc=', irc
         stop
      endif
! allocate vector particle data
      nppmx0 = (1.0 + xtras)*nppmx
      ntmaxp = xtras*nppmx
      npbmx = xtras*nppmx
      nbmaxp = 0.25*mx1*npbmx
      allocate(sbufl(idimp,nbmaxp),sbufr(idimp,nbmaxp))
      allocate(rbufl(idimp,nbmaxp),rbufr(idimp,nbmaxp))
      allocate(ppart(idimp,nppmx0,mxyp1))
      allocate(ppbuff(idimp,npbmx,mxyp1))
      allocate(ncl(8,mxyp1))
      allocate(iholep(2,ntmaxp+1,mxyp1))
      allocate(ncll(3,mx1),nclr(3,mx1),mcll(3,mx1),mclr(3,mx1))
! copy ordered particle data for OpenMP
      call PPPMOVIN2L(part,ppart,kpic,npp,noff,nppmx0,idimp,npmax,mx,my,&
     &mx1,mxyp1,irc)
      if (irc /= 0) then
        write (*,*) kstrt, 'PPPMOVIN2L overflow error, irc=', irc
         call PPABORT()
         stop
      endif
! sanity check
      call PPPCHECK2L(ppart,kpic,noff,nyp,idimp,nppmx0,nx,mx,my,mx1,myp1&
     &,irc)
      if (irc /= 0) then
        write (*,*) kstrt, 'PPPCHECK2L error: irc=', irc
        call PPABORT()
         stop
      endif
!
      if (dt > 0.45*ci) then
         if (kstrt==1) then
            write (*,*) 'Warning: Courant condition may be exceeded!'
         endif
      endif
!
! * * * start main iteration loop * * *
 500 if (nloop <= ntime) go to 2000
      if (kstrt==1) write (*,*) 'ntime = ', ntime
!
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! deposit current with standard procedure: updates part, cue, ihole
     call dtimer(dtime,itime,-1)
     cue = 0.0
      if (relativity==1) then
        call PPGRJPPOSTF2L(ppart,cue,kpic,ncl,iholep,noff,nyp,qme,dth, &
     &ci,nppmx0,idimp,nx,ny,mx,my,nxe,nypmx,mx1,mxyp1,ntmaxp,irc)
     else
         call PPGJPPOSTF2L(ppart, cue, kpic, ncl, iholep, noff, nyp, qme, dth, &
     &nppmx0,idimp,nx,ny,mx,my,nxe,nypmx,mx1,mxyp1,ntmaxp,irc)
     call dtimer(dtime, itime, 1)
     time = real(dtime)
     tdjpost = tdjpost + time
      if (irc \neq 0) then
         if (relativity==1) then
           write (*,*) kstrt, 'PPGRJPPOSTF2L error: irc=', irc
           write (*,*) kstrt, 'PPGJPPOSTF2L error: irc=', irc
         endif
        call PPABORT()
        stop
     endif
! reorder particles by tile with OpenMP
! first part of particle reorder on x and y cell with mx, my tiles:
! updates ppart, ppbuff, ncl, iholep, irc, sbufl, sbufr, ncll, nclr
     call dtimer(dtime,itime,-1)
      call PPPORDERF2LA(ppart,ppbuff,sbufl,sbufr,ncl,iholep,ncll,nclr, &
     &idimp,nppmx0,mx1,myp1,npbmx,ntmaxp,nbmaxp,irc)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tsort = tsort + time
      if (irc /= 0) then
        write (*,*) kstrt, 'PPPORDERF2LA error: ntmaxp, irc=',ntmaxp,irc
        call PPABORT()
         stop
     endif
! move particles into appropriate spatial regions:
! updates rbufr, rbufl, mcll, mclr
     call dtimer(dtime,itime,-1)
      call PPPMOVE2(sbufr,sbufl,rbufr,rbufl,ncll,nclr,mcll,mclr,kstrt, &
     &nvp,idimp,nbmaxp,mx1)
     call dtimer(dtime,itime,1)
     time = real(dtime)
      tmov = tmov + time
! second part of particle reorder on x and y cell with mx, my tiles:
! updates ppart, kpic
     call dtimer(dtime,itime,-1)
      call PPPORDER2LB(ppart,ppbuff,rbuf1,rbufr,kpic,ncl,iholep,mcll,
     &mclr,idimp,nppmx0,mx1,myp1,npbmx,ntmaxp,nbmaxp,irc)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tsort = tsort + time
     if (irc /= 0) then
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write (*,*) kstrt, 'PPPORDER2LB error: nppmx0, irc=',nppmx0,irc
         call PPABORT()
         stop
      endif
!
! deposit charge with standard procedure: updates qe
      call dtimer(dtime, itime, -1)
      qe = 0.0
      call PPGPPOST2L(ppart, qe, kpic, noff, qme, idimp, nppmx0, mx, my, nxe,
     &nypmx,mx1,mxyp1)
      call dtimer(dtime,itime,1)
      time = real(dtime)
      tdpost = tdpost + time
! add guard cells with standard procedure: updates cue, qe
      call dtimer(dtime,itime,-1)
      call PPACGUARD2XL(cue,nyp,nx,ndim,nxe,nypmx)
      call PPNACGUARD2L(cue,scr,nyp,nx,ndim,kstrt,nvp,nxe,nypmx)
      call PPAGUARD2XL(qe,nyp,nx,nxe,nypmx)
      call PPNAGUARD2L(qe,scr,nyp,nx,kstrt,nvp,nxe,nypmx)
      call dtimer(dtime, itime, 1)
      time = real(dtime)
      tquard = tquard + time
! transform charge to fourier space with standard procedure: updates qt
! modifies qe
      call dtimer(dtime,itime,-1)
      isign = -1
      call WPPFFT2RM(qe,qt,bs,br,isign,ntpose,mixup,sct,ttp,indx,indy, &
     &kstrt,nvp,nxeh,nye,kxp,kyp,nypmx,nxhy,nxyh)
      call dtimer(dtime,itime,1)
      time = real(dtime)
      tfft(1) = tfft(1) + time
      tfft(2) = tfft(2) + ttp
!
! transform current to fourier space with standard procedure: updates cut
! modifies cue
      call dtimer(dtime,itime,-1)
      isign = -1
      call WPPFFT2RM3(cue,cut,bs,br,isign,ntpose,mixup,sct,ttp,indx,indy&
     &, kstrt, nvp, nxeh, nye, kxp, kyp, nypmx, nxhy, nxyh)
      call dtimer(dtime,itime,1)
      time = real(dtime)
      tfft(1) = tfft(1) + time
      tfft(2) = tfft(2) + ttp
! take transverse part of current with standard procedure: updates cut
      call dtimer(dtime,itime,-1)
      call MPPCUPERP2(cut,nx,ny,kstrt,nye,kxp)
      call dtimer(dtime, itime, 1)
      time = real(dtime)
      tfield = tfield + time
! calculate electromagnetic fields in fourier space with standard
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! procedure: updates exyz, bxyz
     call dtimer(dtime, itime, -1)
      if (ntime==0) then
        call MIPPBPOISP23(cut,bxyz,ffc,ci,wm,nx,ny,kstrt,nye,kxp,nyh)
        wf = 0.0
        dth = 0.5*dt
     else
         call MPPMAXWEL2(exyz,bxyz,cut,ffc,affp,ci,dt,wf,wm,nx,ny,kstrt,&
     &nye,kxp,nyh)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tfield = tfield + time
! calculate force/charge in fourier space with standard procedure:
! updates fxyt
     call dtimer(dtime, itime, -1)
      isign = -1
     call MPPOIS23(qt,fxyt,isiqn,ffc,ax,ay,affp,we,nx,ny,kstrt,nye,kxp,&
     &nyh)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tfield = tfield + time
! add longitudinal and transverse electric fields with standard
! procedure: updates fxyt
     call dtimer(dtime,itime,-1)
      isign = 1
     call MPPEMFIELD2(fxyt,exyz,ffc,isign,nx,ny,kstrt,nye,kxp,nyh)
! copy magnetic field with standard procedure: updates bxyt
     isign = -1
     call MPPEMFIELD2(bxyt,bxyz,ffc,isign,nx,ny,kstrt,nye,kxp,nyh)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tfield = tfield + time
! transform force to real space with standard procedure: updates fxyze
! modifies fxyt
     call dtimer(dtime, itime, -1)
     isign = 1
     call WPPFFT2RM3(fxyze,fxyt,bs,br,isign,ntpose,mixup,sct,ttp,indx, &
     &indy,kstrt,nvp,nxeh,nye,kxp,kyp,nypmx,nxhy,nxyh)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tfft(1) = tfft(1) + time
     tfft(2) = tfft(2) + ttp
! transform magnetic field to real space with standard procedure:
! updates bxyze, modifies bxyt
     call dtimer(dtime, itime, -1)
      isign = 1
     call WPPFFT2RM3(bxyze,bxyt,bs,br,isign,ntpose,mixup,sct,ttp,indx, &
     &indy,kstrt,nvp,nxeh,nye,kxp,kyp,nypmx,nxhy,nxyh)
     call dtimer(dtime,itime,1)
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time = real(dtime)
     tfft(1) = tfft(1) + time
     tfft(2) = tfft(2) + ttp
!
! copy guard cells with standard procedure: updates fxyze, bxyze
     call dtimer(dtime, itime, -1)
     call PPNCGUARD2L(fxyze,nyp,kstrt,nvp,nnxe,nypmx)
     call PPCGUARD2XL(fxyze,nyp,nx,ndim,nxe,nypmx)
     call PPNCGUARD2L(bxyze,nyp,kstrt,nvp,nnxe,nypmx)
     call PPCGUARD2XL(bxyze,nyp,nx,ndim,nxe,nypmx)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tquard = tquard + time
! push particles: updates part, wke, and ihole
     call dtimer(dtime,itime,-1)
     wke = 0.0
     if (relativity==1) then
         call PPGRBPPUSHF23L(ppart,fxyze,bxyze,kpic,ncl,iholep,noff,nyp,&
    &qbme,dt,dth,ci,wke,idimp,nppmx0,nx,ny,mx,my,nxe,nypmx,mx1,mxyp1, &
    &ntmaxp,irc)
     else
         call PPGBPPUSHF23L(ppart,fxyze,bxyze,kpic,ncl,iholep,noff,nyp, &
    &qbme,dt,dth,wke,idimp,nppmx0,nx,ny,mx,my,nxe,nypmx,mx1,mxyp1,
    &ntmaxp,irc)
     endif
     call dtimer(dtime, itime, 1)
     time = real(dtime)
     tpush = tpush + time
      if (irc /= 0) then
         if (relativity==1) then
           write (*,*) kstrt, 'PPGRBPPUSHF23L error: irc=', irc
        else
           write (*,*) kstrt, 'PPGBPPUSHF23L error: irc=', irc
        endif
        call PPABORT()
         stop
     endif
! reorder particles by tile with OpenMP
! first part of particle reorder on x and y cell with mx, my tiles:
! updates ppart, ppbuff, ncl, iholep, irc, sbufl, sbufr, ncll, nclr
     call dtimer(dtime,itime,-1)
     call PPPORDERF2LA(ppart,ppbuff,sbufl,sbufr,ncl,iholep,ncll,nclr, &
    &idimp,nppmx0,mx1,myp1,npbmx,ntmaxp,nbmaxp,irc)
     call dtimer(dtime,itime,1)
     time = real(dtime)
     tsort = tsort + time
      if (irc /= 0) then
        write (*,*) kstrt, 'PPPORDERF2LA error: ntmaxp, irc=',ntmaxp,irc
        call PPABORT()
         stop
     endif
! move particles into appropriate spatial regions:
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! updates rbufr, rbufl, mcll, mclr
      call dtimer(dtime, itime, -1)
      call PPPMOVE2(sbufr,sbufl,rbufr,rbufl,ncll,nclr,mcll,mclr,kstrt, &
     &nvp,idimp,nbmaxp,mx1)
      call dtimer(dtime,itime,1)
      time = real(dtime)
      tmov = tmov + time
! second part of particle reorder on x and y cell with mx, my tiles:
! updates ppart, kpic
      call dtimer(dtime,itime,-1)
      call PPPORDER2LB(ppart,ppbuff,rbufl,rbufr,kpic,ncl,iholep,mcll,
     &mclr,idimp,nppmx0,mx1,myp1,npbmx,ntmaxp,nbmaxp,irc)
      call dtimer(dtime,itime,1)
     time = real(dtime)
      tsort = tsort + time
      if (irc /= 0) then
        write (*,*) kstrt, 'PPPORDER2LB error: nppmx0, irc=',nppmx0,irc
         call PPABORT()
         stop
      endif
!
! energy diagnostic
      wt = we + wf + wm
     wtot(1) = wt
     wtot(2) = wke
      wtot(3) = 0.0
      wtot(4) = wke + wt
     wtot(5) = we
     wtot(6) = wf
      wtot(7) = wm
      call PPSUM(wtot,work,7)
     wke = wtot(2)
     we = wtot(5)
     wf = wtot(6)
     wm = wtot(7)
      if (ntime==0) then
         if (kstrt.eq.1) then
            wt = we + wf + wm
            write (*,*) 'Initial Total Field, Kinetic and Total Energies&
     &:'
            write (*,'(3e14.7)') wt, wke, wke + wt
            write (*,*) 'Initial Electrostatic, Transverse Electric and &
     &Magnetic Field Energies: '
            write (*,'(3e14.7)') we, wf, wm
         endif
      endif
      ntime = ntime + 1
      go to 500
2000 continue
! * * * end main iteration loop * * *
      if (kstrt.eq.1) then
        write (*,*) 'ntime, relativity = ', ntime, relativity
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write (*,*) 'MPI nodes nvp = ', nvp
        wt = we + wf + wm
        write (*,*) 'Final Total Field, Kinetic and Total Energies:'
        write (*,'(3e14.7)') wt, wke, wke + wt
        write (*,*) 'Final Electrostatic, Transverse Electric and Magne&
     &tic Field Energies:'
        write (*,'(3e14.7)') we, wf, wm
!
        write (*,*)
        write (*,*) 'deposit time = ', tdpost
        write (*,*) 'current deposit time = ', tdjpost
        tdpost = tdpost + tdjpost
        write (*,*) 'total deposit time = ', tdpost
        write (*,*) 'guard time = ', tguard
        write (*,*) 'solver time = ', tfield
        write (*,*) 'fft and transpose time = ', tfft(1), tfft(2)
        write (*,*) 'push time = ', tpush
        write (*,*) 'particle move time = ', tmov
        write (*,*) 'sort time = ', tsort
        tfield = tfield + tguard + tfft(1)
        write (*,*) 'total solver time = ', tfield
        time = tdpost + tpush + tmov + tsort
        write (*,*) 'total particle time = ', time
        wt = time + tfield
        write (*,*) 'total time = ', wt
        write (*,*)
!
        wt = 1.0e+09/(real(nloop)*real(np))
        write (*,*) 'Push Time (nsec) = ', tpush*wt
        write (*,*) 'Deposit Time (nsec) = ', tdpost*wt
        write (*,*) 'Sort Time (nsec) = ', tsort*wt
        write (*,*) 'Total Particle Time (nsec) = ', time*wt
     endif
3000 continue
     call PPEXIT()
      stop
     end program
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