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/* Skeleton 2D Electrostatic MPI/OpenMP PIC code */
/* written by Viktor K. Decyk, UCLA */
#include <stdlib.h>
#include <stdio.h>
#include <complex.h>
#include <sys/time.h>
#include "mppush2.h"
#include "pplib2.h"
#include "omplib.h"
void dtimer(double *time, struct timeval *itime, int icntrl);
int main(int argc, char *argv[]) {
   int indx = 9, indy = 9;
   int npx = 3072, npy = 3072;
   int ndim = 2;
   float tend = 10.0, dt = 0.1, qme = -1.0;
   float vtx = 1.0, vty = 1.0, vx0 = 0.0, vy0 = 0.0;
   float ax = .912871, ay = .912871;
/* idimp = dimension of phase space = 4 */
   int idimp = 4, ipbc = 1;
/* idps = number of partition boundaries */
   int idps = 2;
   float wke = 0.0, we = 0.0, wt = 0.0;
/* sorting tiles, should be less than or equal to 32 */
   int mx = 16, my = 16;
/* fraction of extra particles needed for particle management */
   float xtras = 0.2;
/* declare scalars for standard code */
   int nx, ny, nxh, nyh, nxe, nye, nxeh, nnxe, nxyh, nxhy;
   int mx1, ntime, nloop, isign, ierr;
   float qbme, affp;
   double np;
/* declare scalars for MPI code */
   int ntpose = 1;
   int nvp, idproc, kstrt, npmax, kxp, kyp, nypmx, nypmn;
   int nyp, noff, npp, nps, myp1, mxyp1;
/* declare scalars for OpenMP code */
   int nppmx, nppmx0, nbmaxp, ntmaxp, npbmx, irc;
   int nvpp;
/* declare arrays for standard code */
   float *part = NULL;
   float *qe = NULL;
   float *fxye = NULL;
   float complex *qt = NULL;
   float complex *fxyt = NULL;
   float complex *ffc = NULL;
   int *mixup = NULL;
   float complex *sct = NULL;
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float wtot[4], work[4];
/* declare arrays for MPI code */
   float complex *bs = NULL, *br = NULL;
   float *sbufl = NULL, *sbufr = NULL, *rbufl = NULL, *rbufr = NULL;
   float *edges = NULL;
   float *scs = NULL, *scr = NULL;
/* declare arrays for OpenMP code */
   float *ppart = NULL, *ppbuff = NULL;
   int *kpic = NULL;
   int *ncl = NULL, *iholep = NULL;
   int *ncll = NULL, *nclr = NULL, *mcll = NULL, *mclr = NULL;
/* declare and initialize timing data */
   float time;
   struct timeval itime;
   float tdpost = 0.0, tguard = 0.0, ttp = 0.0, tfield = 0.0;
   float tpush = 0.0, tsort = 0.0, tmov = 0.0;
   float tfft[2] = \{0.0,0.0\};
   double dtime;
   irc = 0;
/* nvpp = number of shared memory nodes (0=default) */
  nvpp = 0;
/* printf("enter number of nodes:\n"); */
/* scanf("%i",&nvpp);
/* initialize for shared memory parallel processing */
   cinit_omp(nvpp);
/* initialize scalars for standard code */
   np = (double) npx*(double) npy;
   nx = 1L << indx; ny = 1L << indy; nxh = nx/2; nyh = ny/2;
   nxe = nx + 2; nye = ny + 2; nxeh = nxe/2; nnxe = ndim*nxe;
  nxyh = (nx > ny ? nx : ny)/2; nxhy = nxh > ny ? nxh : ny;
  mx1 = (nx - 1)/mx + 1;
  nloop = tend/dt + .0001; ntime = 0;
  qbme = qme;
   affp = (double) nx*(double) ny/np;
/* nvp = number of distributed memory nodes */
/* initialize for distributed memory parallel processing */
   cppinit2(&idproc,&nvp,argc,argv);
   kstrt = idproc + 1;
/* check if too many processors */
   if (nvp > ny) {
      if (kstrt==1) {
         printf("Too many processors requested: ny, nvp=%d,%d\n",ny,nvp);
      goto L3000;
   }
/* initialize data for MPI code */
   edges = (float *) malloc(idps*sizeof(float));
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*/
/* calculate partition variables: edges, nyp, noff, nypmx
/* edges[0:1] = 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
                                                                        */
   cpdicomp21(edges,&nyp,&noff,&nypmx,&nypmn,ny,kstrt,nvp,idps);
  if (nypmn < 1) {
      if (kstrt==1) {
        printf("combination not supported nvp, ny =%d,%d\n",ny,nvp);
     goto L3000;
   }
/* 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 */
   part = (float *) malloc(idimp*npmax*sizeof(float));
  qe = (float *) malloc(nxe*nypmx*sizeof(float));
  fxye = (float *) malloc(ndim*nxe*nypmx*sizeof(float));
  qt = (float complex *) malloc(nye*kxp*sizeof(float complex));
   fxyt = (float complex *) malloc(ndim*nye*kxp*sizeof(float complex));
   ffc = (float complex *) malloc(nyh*kxp*sizeof(float complex));
  mixup = (int *) malloc(nxhy*sizeof(int));
   sct = (float complex *) malloc(nxyh*sizeof(float complex));
  kpic = (int *) malloc(mxyp1*sizeof(int));
/* allocate and initialize data for MPI code */
   bs = (float complex *) malloc(ndim*kxp*kyp*sizeof(float complex));
   br = (float complex *) malloc(ndim*kxp*kyp*sizeof(float complex));
   scs = (float *) malloc(nxe*2*sizeof(float));
   scr = (float *) malloc(nxe*2*sizeof(float));
/* prepare fft tables */
   cwpfft2rinit(mixup,sct,indx,indy,nxhy,nxyh);
/* calculate form factors */
   isign = 0;
   cmppois22(qt,fxyt,isign,ffc,ax,ay,affp,&we,nx,ny,kstrt,nye,kxp,nyh);
/* initialize electrons */
  nps = 1;
   npp = 0;
   cpdistr2(part,edges,&npp,nps,vtx,vty,vx0,vy0,npx,npy,nx,ny,idimp,
           npmax,idps,ipbc,&ierr);
/* check for particle initialization error */
   if (ierr != 0) {
      if (kstrt==1) {
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printf("particle initialization error: ierr=%d\n",ierr);
      }
      goto L3000;
   }
/* find number of particles in each of mx, my tiles: updates kpic, nppmx */
   cppdblkp21(part,kpic,npp,noff,&nppmx,idimp,npmax,mx,my,mx1,mxyp1,
   if (irc != 0) {
     printf("%d,cppdblkp21 error, irc=%d\n",kstrt,irc);
      cppabort();
      exit(1);
/* allocate vector particle data */
   nppmx0 = (1.0 + xtras)*nppmx;
   ntmaxp = xtras*nppmx;
   npbmx = xtras*nppmx;
   nbmaxp = 0.25*mx1*npbmx;
   sbufl = (float *) malloc(idimp*nbmaxp*sizeof(float));
   sbufr = (float *) malloc(idimp*nbmaxp*sizeof(float));
   rbufl = (float *) malloc(idimp*nbmaxp*sizeof(float));
   rbufr = (float *) malloc(idimp*nbmaxp*sizeof(float));
   ppart = (float *) malloc(idimp*nppmx0*mxyp1*sizeof(float));
   ppbuff = (float *) malloc(idimp*npbmx*mxyp1*sizeof(float));
   ncl = (int *) malloc(8*mxyp1*sizeof(int));
   iholep = (int *) malloc(2*(ntmaxp+1)*mxyp1*sizeof(int));
   ncll = (int *) malloc(3*mxyp1*sizeof(int));
   nclr = (int *) malloc(3*mxyp1*sizeof(int));
   mcll = (int *) malloc(3*mxyp1*sizeof(int));
   mclr = (int *) malloc(3*mxyp1*sizeof(int));
/* copy ordered particle data for OpenMP */
   cpppmovin2l(part,ppart,kpic,npp,noff,nppmx0,idimp,npmax,mx,my,mx1,
                  mxyp1,&irc);
   if (irc != 0) {
      printf("%d,cpppmovin2l overflow error, irc=%d\n",kstrt,irc);
      cppabort();
      exit(1);
   }
/* sanity check */
   cpppcheck2l(ppart,kpic,noff,nyp,idimp,nppmx0,nx,mx,my,mx1,myp1,&irc);
   if (irc != 0) {
      printf("%d,cpppcheck2l error: irc=%d\n",kstrt,irc);
      cppabort();
      exit(1);
   }
/* * * * start main iteration loop * * * */
L500: if (nloop <= ntime)
         goto L2000;
/*
      if (kstrt==1) printf("ntime = %i\n",ntime); */
/* deposit charge with standard procedure: updates qe */
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dtimer(&dtime,&itime,-1);
      for (j = 0; j < nxe*nypmx; j++) {
         qe[j] = 0.0;
      }
      cppgppost21(ppart,qe,kpic,noff,qme,idimp,nppmx0,mx,my,nxe,nypmx,
                  mx1,mxyp1);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tdpost += time;
/* add quard cells with standard procedure: updates qe */
      dtimer(&dtime,&itime,-1);
      cppaguard2xl(qe,nyp,nx,nxe,nypmx);
      cppnaguard21(qe,scr,nyp,nx,kstrt,nvp,nxe,nypmx);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tquard += time;
/* transform charge to fourier space with standard procedure: updates qt */
/* modifies qe */
      dtimer(&dtime, &itime, -1);
      isign = -1;
      cwppfft2rm((float complex *)qe,qt,bs,br,isiqn,ntpose,mixup,sct,
                 &ttp,indx,indy,kstrt,nvp,nxeh,nye,kxp,kyp,nypmx,nxhy,
                 nxyh);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tfft[0] += time;
      tfft[1] += ttp;
/* calculate force/charge in fourier space with standard procedure: */
/* updates fxyt */
      dtimer(&dtime, &itime, -1);
      isign = -1;
      cmppois22(qt,fxyt,isign,ffc,ax,ay,affp,&we,nx,ny,kstrt,nye,kxp,
                nyh);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tfield += time;
/* transform force to real space with standard procedure: updates fxye */
/* modifies fxyt */
      dtimer(&dtime,&itime,-1);
      isign = 1;
      cwppfft2rm2((float complex *)fxye,fxyt,bs,br,isiqn,ntpose,mixup,
                  sct, &ttp, indx, indy, kstrt, nvp, nxeh, nye, kxp, kyp, nypmx,
                  nxhy, nxyh);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tfft[0] += time;
      tfft[1] += ttp;
/* copy guard cells with standard procedure: updates fxye */
      dtimer(&dtime, &itime, -1);
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cppncquard21(fxye,nyp,kstrt,nvp,nnxe,nypmx);
      cppcguard2xl(fxye,nyp,nx,ndim,nxe,nypmx);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tguard += time;
/* push particles: updates part, wke, and ihole */
      dtimer(&dtime,&itime,-1);
      wke = 0.0;
      cppgppushf21(ppart,fxye,kpic,ncl,iholep,noff,nyp,qbme,dt,&wke,nx,
                   ny, mx, my, idimp, nppmx0, nxe, nypmx, mx1, mxyp1, ntmaxp,
                   &irc);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tpush += time;
      if (irc != 0) {
         printf("%d,cppgppushf2l error: irc=%d\n",kstrt,irc);
         cppabort();
         exit(1);
      }
/* 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 */
      dtimer(&dtime, &itime, -1);
      cppporderf2la(ppart,ppbuff,sbufl,sbufr,ncl,iholep,ncll,nclr,
                    idimp,nppmx0,mx1,myp1,npbmx,ntmaxp,nbmaxp,&irc);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tsort += time;
      if (irc != 0) {
         printf("%d,cppporderf2la error:ntmaxp,irc=%d,%d\n",kstrt,
                ntmaxp,irc);
         cppabort();
         exit(1);
/* move particles into appropriate spatial regions: */
/* updates rbufr, rbufl, mcll, mclr */
      dtimer(&dtime,&itime,-1);
      cpppmove2(sbufr,sbufl,rbufr,rbufl,ncll,nclr,mcll,mclr,kstrt,nvp,
                idimp,nbmaxp,mx1);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tmov += time;
/* second part of particle reorder on x and y cell with mx, my tiles: */
/* updates ppart, kpic */
      dtimer(&dtime,&itime,-1);
      cppporder2lb(ppart,ppbuff,rbufl,rbufr,kpic,ncl,iholep,mcll,mclr,
                   idimp,nppmx0,mx1,myp1,npbmx,ntmaxp,nbmaxp,&irc);
      dtimer(&dtime, &itime, 1);
      time = (float) dtime;
      tsort += time;
      if (irc != 0) {
         printf("%d,cppporder2lb error:nppmx0,irc=%d,%d\n",kstrt,nppmx0,
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irc);
         cppabort();
         exit(1);
      }
/* energy diagnostic */
      wtot[0] = we;
      wtot[1] = wke;
      wtot[2] = 0.0;
      wtot[3] = we + wke;
      cppsum(wtot,work,4);
      we = wtot[0];
      wke = wtot[1];
      if (ntime==0) {
         if (kstrt==1) {
            printf("Initial Field, Kinetic and Total Energies:\n");
            printf("%e %e %e\n",we,wke,wke+we);
         }
      }
      ntime += 1;
      goto L500;
L2000:
/* * * * end main iteration loop * * * */
   if (kstrt==1) {
      printf("ntime = %i\n",ntime);
      printf("MPI nodes nvp = %i\n",nvp);
      printf("Final Field, Kinetic and Total Energies:\n");
      printf("%e %e %e\n",we,wke,wke+we);
      printf("\n");
      printf("deposit time = %f\n",tdpost);
      printf("guard time = %f\n",tguard);
      printf("solver time = %f\n",tfield);
      printf("fft and transpose time = %f,%f\n",tfft[0],tfft[1]);
      printf("push time = %f\n",tpush);
      printf("particle move time = %f\n",tmov);
      printf("sort time = %f\n",tsort);
      tfield += tguard + tfft[0];
      printf("total solver time = %f\n",tfield);
      time = tdpost + tpush + tmov + tsort;
      printf("total particle time = %f\n",time);
      wt = time + tfield;
      printf("total time = %f\n",wt);
      printf("\n");
      wt = 1.0e+09/(((float) nloop)*((float) np));
      printf("Push Time (nsec) = %f\n",tpush*wt);
      printf("Deposit Time (nsec) = %f\n",tdpost*wt);
      printf("Sort Time (nsec) = %f\n",tsort*wt);
      printf("Total Particle Time (nsec) = %f\n",time*wt);
   }
```

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L3000:
    cppexit();
    return 0;
}
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