

Implementation of elastic-plastic model in PDLAMMPS

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1 Introduction

In this documentation the discussion is focused on implementation of elastic plastic peridynamics model in PDLAMMPS [PLPS08],[Pli95]. The peridynamic elastic-plastic formulation (and its time integration) used in this work was developed by John Mitchell at Sandia National Lab [Mit11]. In the PDLAMMPS a new pair-`peri`-style is added in order to incorporate the elastic-plastic formulation. The new source codes `pair-peri-eps.cpp` and `pair-peri-eps.h` are introduced. Besides, `fix-peri-neigh.cpp` and `fix-peri-neigh.h` are modified for introducing elastic-plastic solid model (EPS).

2 Algorithm and implementation

The total extension state can be decomposed into two parts [Mit11] ¹:

¹For details, please see documentation of state based peridynamics plasticity model [Mit11].

$$\text{Total extension: } e(Y) = e^i(Y) + e^d(Y) \quad (1)$$

$$\text{Volumetric extension: } e^i(Y) = \frac{\theta(Y)|X|}{3} \quad (2)$$

$$\text{Deviatoric extension: } e^d(Y) = |Y| - |X| - \frac{\theta(Y)|X|}{3} \quad (3)$$

Here, $|Y|$, $|X|$ and θ are the reference state, deformation state and dilation state, respectively. The deviatoric extension can be written as:

$$e^d(Y) = e^{de}(Y) + e^{dp(i)}(Y) \quad (4)$$

Here, $e^{de}(Y)$ and $e^{dp(i)}$ are the elastic and plastic parts of the deviatoric extension. The elastic-plastic force scalar state can be written as:

$$t = t^i + t^d = -\frac{3p}{m}\omega\underline{x} + \alpha\omega(e^d - e^{dp}) \quad (5)$$

p , k , α , t^i and t^d are hydrostatic pressure, bulk modulus, elastic properties volumetric and deviatoric scalar force states, respectively. $\alpha = \frac{15\mu}{m}$; where, μ , m are the shear modulus and weighted volume, respectively. The influence function: $\omega(\xi) = \frac{1}{\|\xi\|}$, $\|\xi\|$ is the scalar reference state. At the current or $(n+1)^{th}$ timestep the scalar trial force state t_{trial}^d and its norm $\|t_{trial}^d\|$ are calculated.

$$t_{trial}^d = \frac{15\mu}{m}\omega(e_{n+1}^d - e_{n+1}^{dp}). \quad (6)$$

Equation. 6 is implemented in the function in `pair_peri_eps.cpp`:

```
double PairPeriEPS::compute_DeviatoricForceStateNorm(int i)
{
    int j,jj,jnum,itype,jtype;
    double xtmp,ytmp,ztmp,dex,dely,delz;
    double xtmp0,ytmp0,ztmp0,dex0,dely0,delz0;
    double rsq,r,dr;
    double delta;
    double tdtrial;
    double norm = 0.0;
    double **x = atom->x;
```

```

int *type = atom->type;
double **x0 = atom->x0;
double *s0 = atom->s0;
int nlocal = atom->nlocal;
double *vfrac = atom->vfrac;
double vfrac_scale = 1.0;
double lc = domain->lattice->xlattice;
double half_lc = 0.5*lc;
double **r0 =
    ((FixPeriNeigh *) modify->fix[ifix_peri])->r0;
int **partner =
    ((FixPeriNeigh *) modify->fix[ifix_peri])->partner;
int *npartner =
    ((FixPeriNeigh *) modify->fix[ifix_peri])->npartner;
double *wvolume =
    ((FixPeriNeigh *) modify->fix[ifix_peri])->wvolume;
double **deviatorPlasticextension =
    ((FixPeriNeigh *) modify->fix[ifix_peri])->deviatorPlasticextension;
int periodic =
    domain->xperiodic || domain->yperiodic || domain->zperiodic;

// compute the dilatation theta

xtmp = x[i][0];
ytmp = x[i][1];
ztmp = x[i][2];
xtmp0 = x0[i][0];
ytmp0 = x0[i][1];
ztmp0 = x0[i][2];
jnum = npartner[i];
itype = type[i];

for (jj = 0; jj < jnum; jj++) {
    if (partner[i][jj] == 0) continue;
    j = atom->map(partner[i][jj]);
    // check if lost a partner without first breaking bond
    if (j < 0) {
        partner[i][jj] = 0;
    }
}

```

```

        continue;
    }
    delx = xtmp - x[j][0];
    dely = ytmp - x[j][1];
    delz = ztmp - x[j][2];
    if (periodic) domain->minimum_image(delx,dely,delz);
    rsq = delx*delx + dely*dely + delz*delz;
    delx0 = xtmp0 - x0[j][0];
    dely0 = ytmp0 - x0[j][1];
    delz0 = ztmp0 - x0[j][2];
    if (periodic) domain->minimum_image(delx0,dely0,delz0);
    r = sqrt(rsq);
    dr = r - r0[i][jj];
    if (fabs(dr) < 2.2204e-016) dr = 0.0;

        // scale vfrac[j] if particle j near the horizon
    double vfrac_scale;

    jtype = type[j];
    double delta = cut[itype][jtype];

    // scale vfrac[j] if particle j near the horizon

    if ((fabs(r0[i][jj] - delta)) <= half_lc)
        vfrac_scale = (-1.0/(2*half_lc))*(r0[i][jj]) +
            (1.0 + ((delta - half_lc)/(2*half_lc) ) );
    else vfrac_scale = 1.0;

    double ed = dr - (theta[i] * r0[i][jj])/3;
    double edPNP1 = deviatorPlasticextension[i][jj];

    jtype = type[j];
    delta = cut[itype][jtype];

    double omega_plus =
        influence_function(-1.0*delx0,-1.0*dely0,-1.0*delz0);
    double omega_minus =
        influence_function(delx0,dely0,delz0);

```

```

double stretch = dr / r0[i][jj];

tdtrial = ( 15 * shearmodulus[itype][itype] ) *
  ((omega_plus * theta[i] / wvolume[i]) +
  ( omega_minus * theta[j] / wvolume[j] ) ) * (ed - edPNP1);

norm += tdtrial * tdtrial * vfrac[j] * vfrac_scale;
}
return sqrt(norm);
}

```

The yield function $f(t_{trial}^d)$ is written based on $\|t_{trial}^d\|$ and yield stress σ_Y (Eq. 6).

$$f(t_{trial}^d) = \frac{\|t_{trial}^d\|^2}{2} - \frac{25\sigma_Y^2}{8\pi\delta^5} \quad (7)$$

Here, δ is the horizon. If $f(t_{trial}^d) < 0$ the step is elastic. Otherwise, it is plastic. If the step is elastic:

$$t_{n+1}^d = t_{trial}^d, \quad (8)$$

$$e_{n+1}^{dp} = e_n^{dp}. \quad (9)$$

For plastic step:

$$\Delta\lambda = \frac{1}{\alpha} \left[\frac{\|t_{trial}^d\|}{\sqrt{2\psi_0}} - 1 \right], \quad (10)$$

$$t_{n+1}^d = \sqrt{2\psi_0} \frac{t_{trial}^d}{\|t_{trial}^d\|}, \quad (11)$$

$$e_{n+1}^{dp} = e_n^{dp} + \Delta\lambda t_{n+1}^d. \quad (12)$$

The return algorithm is implemented in `PairPeriEPS::compute(int, int)`.

3 LAMMPS command for PD EPS

There is no significant change in the LAMMPS input script for elastic-plastic model. For PD EPS the LAMMPS commands are:

```
pair_style peri/eps
pair_coeff i j Bulk_modulus Shear_modulus s00 alpha Yield_Stress
```

4 Conclusion

The LAMMPS implementation of peridynamic elastic-plastic model is still in beta phase. Any bug or issue can be informed to the authors through `rezwanur.rahman@utsa.edu`.

5 Acknowledgment

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References

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6 Appendix

In the current updated version of PDLAMMPS there are two new compute commands are included. One is **compute ID group-ID plasticity/atom**, and **compute ID group-ID dilatation/atom**. Compute style plasticity/atom is applicable for pair style peri/eps and compute style dilatation/atom is applicable for pair styles peri/lps, peri/ves and peri/eps. In compute style plasticity, λ for each peridynamic node is calculated.

The dilatation at each peridynamics node θ [SEW⁺07] is computed by compute dilatation/atom command. The equation to calculate theta is

$$\theta(\mathbf{x}) = \frac{3}{m(\mathbf{x})} \int_{\mathcal{H}_x} \omega\langle\xi\rangle \underline{x}\langle\xi\rangle e\langle\xi\rangle dV_\xi \quad (13)$$

$$m(\mathbf{x}) = \int_{\mathcal{H}_x} \omega\langle\xi\rangle \underline{x}\langle\xi\rangle \underline{x}\langle\xi\rangle dV_\xi \quad (14)$$

Here, \mathcal{H}_x , $e\langle\xi\rangle$, $\underline{x}\langle\xi\rangle = \|\xi\|$, dV_ξ are the horizon (i.e. neighborhood) of a peridynamic node, bond extension state, reference position scalar state and volume of a peridynamic node or particle. $m(\mathbf{x})$ is the weighted volume. The numerical implementation of Eq. 13 and 14 is explained in the PDLAMMPS documentation [PSP⁺10]. In the **compute ID group-ID dilatation/atom** the calculated values of θ at each peridynamic node or atom is stored in a vector.