

LAMMPS 23Jun22 with GPU support

Webpage

<https://www.lammps.org>

Version

23Jun22

Build Environment

- Intel oneAPI Compiler Classic 2022.2.1
- Intel MKL 2022.2.1
- HPC-X 2.11 (Open MPI 4.1.4)
 - HPC-X 2.13.1 (Open MPI 4.1.5) was used on actual build. Runtime library was switched to HPC-X 2.11 to avoid the communication error.
 - In the following procedure, we assume HPC-X 2.11 was used.
- CUDA 11.6

Files Required

- lammps-stable.tar.gz
- (some of files will be downloaded during the installation)

Build Procedure

conda environment

Conda (miniforge) environment prepared for [lammps 29Sep21 CPU version](#) is used.

lammps

```
#!/bin/sh

VERSION=23Jun22
NAME=lammps-23Jun2022
INSTALL_PREFIX=/apl/lammps/2022-Jun23-CUDA

BASEDIR=/home/users/${USER}/Software/LAMMPS/${VERSION}
LAMMPS_TARBALL=${BASEDIR}/lammps-stable.tar.gz

WORKDIR=/gwork/users/${USER}/lammps-23Sep2022-cuda
LAMMPS_WORKDIR=${WORKDIR}/${NAME}

GPU_ARCH=sm_80
VMD_MOLFILE_INC=/home/users/${USER}/Software/VMD/1.9.4/vmd-1.9.4a57/plugins/include

PARALLEL=12

#-----
umask 0022
export LANG=C
ulimit -s unlimited

module -s purge

# load intel compiler
. ~/intel/oneapi/compiler/2022.2.1/env/vars.sh

. /apl/lammps/2022-Jun23-CUDA/conda_init.sh

module -s load mkl/2022.2.1
module -s load cuda/11.6
```

```

module -s load openmpi/4.1.4-hpcx/intel2022.2.1

export CC=mpicc
export CXX=mpicxx
export FC=mpif90
export MPICC=mpicc
export MPICXX=mpicxx
export MPIFC=mpif90

cd ${WORKDIR}
if [ -d ${NAME} ]; then
  mv ${NAME} lammmps_erase
  rm -rf lammmps_erase &
fi

tar zxf ${LAMMPS_TARBALL}

cd ${NAME}
sed -i -e "s/xHost/march=core-avx2/" cmake/CMakeLists.txt
mkdir build && cd build

# Disabled PKGs:
# FFMPEG, ADIOS, MDI, VTK: noavail
# MSCG: gsl too old
# MESSAGE: ZeroMQ support not enabled
# QUIP: failed to build
# ML-HDNNP: failed to build
# KIM: CDDL is incompatible with GPL
# LATTE: technical problem of cmake? (LAPACK and BLAS)
# NETCDF: to avoid EVP_KDF_ctrl error
# MPIIO: not maintained?

cmake ../cmake \
-DLAMMPS_MACHINE=rccs-cuda \
-DENABLE_TESTING=on \
-DCMAKE_INSTALL_PREFIX=${INSTALL_PREFIX} \
-DCMAKE_C_COMPILER=mpicc \
-DCMAKE_CXX_COMPILER=mpicxx \
-DCMAKE_Fortran_COMPILER=mpif90 \
-DCMAKE_MPI_C_COMPILER=mpicc \
-DCMAKE_MPI_CXX_COMPILER=mpicxx \
-DCMAKE_MPI_Fortran_COMPILER=mpif90 \
-DCMAKE_CXX_FLAGS_DEBUG="-Wall -Wextra -g" \
-DCMAKE_CXX_FLAGS_RELWITHDEBINFO="-Wall -Wextra -g -O2 -DNDEBUG" \
-DCMAKE_CXX_FLAGS_RELEASE="-O3 -DNDEBUG" \
-DCMAKE_Fortran_FLAGS_DEBUG="-Wall -Wextra -g" \
-DCMAKE_Fortran_FLAGS_RELWITHDEBINFO="-Wall -Wextra -g -O2 -DNDEBUG" \
-DCMAKE_Fortran_FLAGS_RELEASE="-O3 -DNDEBUG" \
-DCMAKE_C_FLAGS_DEBUG="-Wall -Wextra -g" \
-DCMAKE_C_FLAGS_RELWITHDEBINFO="-Wall -Wextra -g -O2 -DNDEBUG" \
-DCMAKE_C_FLAGS_RELEASE="-O3 -DNDEBUG" \
-DLAMMPS_EXCEPTIONS=on \
-DBUILD_SHARED_LIBS=on \
-DBUILD_TOOLS=on \
-DBUILD_MPI=on \
-DBUILD_OMP=on \
-DFFT=MKL \
-DFFT_SINGLE=on \
-DFFT_MKL_THREADS=on \
-DWITH_JPEG=yes \
-DWITH_PNG=yes \
-DWITH_GZIP=yes \
-DPKG_ASPHERE=on \
-DPKG_ATC=on \
-DPKG_AWPMO=on \

```

-DPKG_BOCS=on \
-DPKG_BODY=on \
-DPKG_BROWNIAN=on \
-DPKG_CG-DNA=on \
-DPKG_CG-SDK=on \
-DPKG_CLASS2=on \
-DPKG_COLLOID=on \
-DPKG_COLVARS=on \
-DPKG_COMPRESS=on \
-DPKG_CORESHELL=on \
-DPKG_DIELECTRIC=on \
-DPKG_DIFFRACTION=on \
-DPKG_DIPOLE=on \
-DPKG_DPD-BASIC=on \
-DPKG_DPD-MESO=on \
-DPKG_DPD-REACT=on \
-DPKG_DPD-SMOOTH=on \
-DPKG_DRUDE=on \
-DPKG_EFF=on \
-DPKG_EXTRA-COMPUTE=on \
-DPKG_EXTRA-DUMP=on \
-DPKG_EXTRA-FIX=on \
-DPKG_EXTRA-MOLECULE=on \
-DPKG_EXTRA-PAIR=on \
-DPKG_FEP=on \
-DPKG_GPU=on \
-DGPU_API=cuda \
-DGPU_ARCH=\${GPU_ARCH} \
-DPKG_GRANULAR=on \
-DPKG_H5MD=on \
-DPKG_INTEL=on \
-DPKG_INTERLAYER=on \
-DPKG_KIM=off \
-DDOWNLOAD_KIM=no \
-DPKG_KOKKOS=off \
-DPKG_KSPACE=on \
-DPKG_LATBOLTZ=on \
-DPKG_MACHDYN=on \
-DDOWNLOAD_EIGEN3=on \
-DPKG_MANIFOLD=on \
-DPKG_MANYBODY=on \
-DPKG_MC=on \
-DPKG_MDI=off \
-DPKG_MEAM=on \
-DPKG_MESONT=on \
-DPKG_MESSAGE=on \
-DPKG_MGPT=on \
-DPKG_MISC=on \
-DPKG_ML-HDNNP=off \
-DDOWNLOAD_N2P2=no \
-DPKG_ML-IAP=on \
-DPKG_ML-PACE=on \
-DPKG_ML-QUIP=off \
-DDOWNLOAD_QUIP=no \
-DPKG_ML-RANN=on \
-DPKG_ML-SNAP=on \
-DPKG_MOFFF=on \
-DPKG_MOLECULE=on \
-DPKG_MOLFILE=on \
-DMOLFILE_INCLUDE_DIR=\${VMD_MOLFILE_INC} \
-DPKG_MPIIO=off \
-DPKG_MSCG=off \
-DPKG_NETCDF=off \
-DPKG_OPENMP=on \
-DPKG_OPT=on \

```

-DPKG_ORIENT=on \
-DPKG_PERI=on \
-DPKG_PHONON=on \
-DPKG_PLUGIN=on \
-DPKG_PLUMED=on \
-DDOWNLOAD_PLUMED=yes \
-DPKG_POEMS=on \
-DPKG_PTM=on \
-DPKG_PYTHON=on \
-DPKG_QEQ=on \
-DPKG_QMMM=on \
-DPKG_QTB=on \
-DPKG_REACTION=on \
-DPKG_REAXFF=on \
-DPKG_REPLICA=on \
-DPKG_RIGID=on \
-DPKG_SCAFACOS=on \
-DDOWNLOAD_SCAFACOS=yes \
-DPKG_SHOCK=on \
-DPKG_SMTBQ=on \
-DPKG_SPH=on \
-DPKG_SPIN=on \
-DPKG_SRD=on \
-DPKG_TALLY=on \
-DPKG_UEF=on \
-DPKG_VORONOI=on \
-DDOWNLOAD_VORO=yes \
-DPKG_VTK=off \
-DPKG_YAFF=on \
-DBLAS_LIBRARIES="-qmkl" \
-DCMAKE_BUILD_TYPE=Release

```

```
make VERBOSE=1 -j ${PARALLEL}
```

```
export OMP_NUM_THREADS=2
```

```
#make test # will put error...
```

```
make install
```

```
cp -a ../examples ${INSTALL_PREFIX}
```

```
cd ${INSTALL_PREFIX}
```

```
for f in etc/profile.d/*; do
```

```
ln -s $f .
```

```
done
```

```
cd lib64
```

```
if [ -f liblammps_rccs-cuda.so ]; then
```

```
ln -s liblammps_rccs-cuda.so liblammps.so
```

```
fi
```

```
if [ -f liblammps_rccs-cuda.so.0 ]; then
```

```
ln -s liblammps_rccs-cuda.so.0 liblammps.so.0
```

```
fi
```

Enabled Packages

```

ASPHERE ATC AWPMD BOCS BODY BROWNIAN CG-DNA CG-SDK CLASS2 COLLOID COLVARS
COMPRESS CORESHELL DIELECTRIC DIFFRACTION DIPOLE DPD-BASIC DPD-MESO
DPD-REACT DPD-SMOOTH DRUDE EFF EXTRA-COMPUTE EXTRA-DUMP EXTRA-FIX EXTRA-MOLECULE
EXTRA-PAIR FEP GPU GRANULAR H5MD INTEL INTERLAYER KSPACE LATBOLTZ MACHDYN
MANIFOLD MANYBODY MC MEAM MESONT MGPT MISC ML-IAP ML-PACE ML-RANN ML-SNAP
MOFFF MOLECULE MOLFILE OPENMP OPT ORIENT PERI PHONON PLUGIN PLUMED POEMS
PTM PYTHON QEQ QMMM QTB REACTION REAXFF REPLICA RIGID SCAFACOS SHOCK
SMTBQ SPH SPIN SRD TALLY UEF VORONOI YAFF

```

Tests

("make test" was done on one of GPU computation nodes.)

Copy of test log file is available at </apl/lammps/2022-Jun23-CUDA/Testing/>.

The following tests FAILED:

- 11 - AtomStyle (Failed)
- 42 - ComputeGlobal (Failed)
- 94 - MolPairStyle:coul_diel (Failed)
- 100 - MolPairStyle:coul_shield (Failed)
- 102 - MolPairStyle:coul_slater_long (Failed)
- 137 - MolPairStyle:lj_class2_soft (Failed)
- 152 - MolPairStyle:lj_cut_soft (Failed)
- 158 - MolPairStyle:lj_expand_coul_long (Failed)
- 171 - MolPairStyle:lj_sdk_coul_long (Failed)
- 172 - MolPairStyle:lj_sdk_coul_table (Failed)
- 176 - MolPairStyle:lj_switch3_coulgauss_long (Failed)
- 200 - MolPairStyle:tip4p_long_soft (Failed)
- 203 - MolPairStyle:wf_cut (Failed)
- 211 - AtomicPairStyle:buck_coul_cut_qeq_point (Failed)
- 212 - AtomicPairStyle:buck_coul_cut_qeq_shielded (Failed)
- 229 - AtomicPairStyle:edip (Failed)
- 236 - AtomicPairStyle:meam (Failed)
- 237 - AtomicPairStyle:meam_spline (Failed)
- 238 - AtomicPairStyle:meam_sw_spline (Failed)
- 241 - AtomicPairStyle:reaxff-acks2 (Failed)
- 242 - AtomicPairStyle:reaxff-acks2_efield (Failed)
- 243 - AtomicPairStyle:reaxff (Failed)
- 244 - AtomicPairStyle:reaxff_lgvdw (Failed)
- 245 - AtomicPairStyle:reaxff_noqeq (Failed)
- 246 - AtomicPairStyle:reaxff_tabulate (Failed)
- 247 - AtomicPairStyle:reaxff_tabulate_flag (Failed)
- 264 - ManybodyPairStyle:comb (Failed)
- 272 - ManybodyPairStyle:ilp-graphene-hbn (Failed)
- 273 - ManybodyPairStyle:ilp-graphene-hbn_notaper (Failed)
- 277 - ManybodyPairStyle:lcbop (Failed)
- 286 - ManybodyPairStyle:pace_product (Failed)
- 287 - ManybodyPairStyle:pace_recursive (Failed)
- 299 - ManybodyPairStyle:tersoff (Failed)
- 304 - ManybodyPairStyle:tersoff_shift (Failed)
- 314 - BondStyle:gaussian (Failed)
- 357 - KSpaceStyle:ewald_tri (Failed)
- 359 - KSpaceStyle:pppm_ad (Failed)
- 360 - KSpaceStyle:pppm_cg (Failed)
- 362 - KSpaceStyle:pppm_cg_tiled (Failed)
- 369 - KSpaceStyle:pppm_disp_tip4p (Failed)
- 377 - KSpaceStyle:pppm_tip4p (Failed)
- 382 - KSpaceStyle:scafacos_direct (Failed)
- 383 - KSpaceStyle:scafacos_ewald (Failed)
- 384 - KSpaceStyle:scafacos_fmm (Failed)
- 385 - KSpaceStyle:scafacos_fmm_tuned (Failed)
- 386 - KSpaceStyle:scafacos_p2nfft (Failed)
- 387 - FixTimestep:adapt_coul (Failed)
- 390 - FixTimestep:addforce_const (Failed)
- 392 - FixTimestep:addtorque_const (Failed)
- 411 - FixTimestep:nph (Failed)
- 412 - FixTimestep:nph_sphere (Failed)
- 414 - FixTimestep:npt_iso (Failed)
- 415 - FixTimestep:npt_sphere_aniso (Failed)
- 416 - FixTimestep:npt_sphere_iso (Failed)
- 440 - FixTimestep:rigid_npt_small (Failed)
- 454 - FixTimestep:smd_couple (Failed)
- 462 - FixTimestep:temp_csld (Failed)
- 483 - DihedralStyle:table_cut_linear (Failed)
- 485 - DihedralStyle:table_linear (Failed)

486 - DihedralStyle:table_spline (Failed)

496 - ImproperStyle:inversion_harmonic (Failed)

- Most errors are minor numerical ones or "lattice" command problem when intel compiler is employed. There may not be serious problems.

Notes

- Built on ccfep, tested on one of GPU computation nodes.
- Parallel runs failed if HPC-X 2.13.1 runtime library was used. Switching to HPC-X 2.11 solved the problem.
- If NETCDF is set to on, the build failed. We thus disabled NETCDF this time. This may be due to the problem of system library.
- The compilation errors could be avoided if python 3.6 of the system was employed... We will try it in the next time.