

Amber22

Webpage

<http://ambermd.org/>

Version

Amber22, AmberTools 22

Build Environment

- Intel Parallel Studio 2018 Update4 (MPI only)
- GCC 9.3.1 (devtoolset-9)
- CUDA 11.1 Update 1

Files Required

- Amber22.tar.bz2
- AmberTools22.tar.bz2
- patch-cmake-python
 - use miniforge instead of miniconda (to avoid license issue of anaconda repository)

```
-- cmake/UseMiniconda.cmake.org      2022-05-27 09:43:57.000000000 +0900
+++ cmake/UseMiniconda.cmake    2022-05-27 09:56:28.000000000 +0900
@@ -84,11 +84,14 @@
        endif()
    endif()

-   set(MINICONDA_INSTALLER_FILENAME "Miniconda${PYTHON_MAJOR_RELEASE}-${MINICONDA_VERSION}-${CONTINUUM_SYSTEM_NAME}-
${CONTINUUM_BITS}.${INSTALLER_SUFFIX}")
+   #set(MINICONDA_INSTALLER_FILENAME "Miniconda${PYTHON_MAJOR_RELEASE}-${MINICONDA_VERSION}-${CONTINUUM_SYSTEM_NAME}-
${CONTINUUM_BITS}.${INSTALLER_SUFFIX}")
+   ## mkamiya: assume x86_64 Linux...
+   set(MINICONDA_INSTALLER_FILENAME "Miniforge${PYTHON_MAJOR_RELEASE}-Linux-x86_64.sh")

    # location to download the installer to
    set(MINICONDA_INSTALLER ${MINICONDA_DOWNLOAD_DIR}/${MINICONDA_INSTALLER_FILENAME})
-   set(INSTALLER_URL "http://repo.continuum.io/miniconda/${MINICONDA_INSTALLER_FILENAME}")
+   #set(INSTALLER_URL "http://repo.continuum.io/miniconda/${MINICONDA_INSTALLER_FILENAME}")
+   set(INSTALLER_URL "https://github.com/conda-
forge/miniforge/releases/${MINICONDA_VERSION}/download/${MINICONDA_INSTALLER_FILENAME}")

    # If we've already downloaded the installer, use it.
    if(EXISTS "${MINICONDA_INSTALLER}")
```

- amber.csh.patch (this file was modified after the installation; not mentioned in the procedure below)
 - The trick to get script directory does not always work. Old-style way (hard-coded) is employed this time.

```
-- amber.csh.org     2022-06-06 16:53:47.000000000 +0900
+++ amber.csh 2022-06-06 16:57:01.000000000 +0900
@@ -16,11 +16,13 @@
        echo "      Your shell does not appear to be a C shell: $0"
    endif

-# Get path used for this source file (credit scott brozell).
-set invocationpath = `echo $_ | cut -d' ' -f2- | sed "s@${myname}.*@@"`
-if ( "$invocationpath" == "" ) then
-    set invocationpath = '.'
-endif
+## RCCS: this does not always work. Use oldstyle setting...
+set invocationpath = "/local/apl/lx/amber22-up0"
+## Get path used for this source file (credit scott brozell).
+#set invocationpath = `echo $_ | cut -d' ' -f2- | sed "s@${myname}.*@@"`
```

```
+#if ( "$invocationpath" == " " ) then
+#
+    set invocationpath = '!
+#
+endif

setenv AMBERHOME `cd "$invocationpath" >&! /dev/null; pwd` 
setenv PATH "$AMBERHOME/bin:$PATH"
```

Build Procedure

```
#!/bin/sh

VERSION=22
TOOLSVERSION=22

# amber22 + AmberTools22
INSTALL_DIR="/local/apl/lx/amber22-up0"
WORKDIR="/work/users/${USER}/work-amber"
TARBALL_DIR="/home/users/${USER}/Software/AMBER/22"

PATCHX=${TARBALL_DIR}/patch-cmake-python

PARALLEL=12

#-----
module purge
module load mpi/intelmpi/2018.4.274
module load scl/devtoolset-9
module load cuda/11.1
module load cmake/3.16.3

export CUDA_HOME="/local/apl/lx/cuda-11.1"

export LANG=C
export LC_ALL=C
ulimit -s unlimited

# install directory has to be prepared before running this script
if [ ! -d $WORKDIR ]; then
    echo "Create $WORKDIR before running this script."
    exit 1
fi

# build directory must be empty
if [ "$(ls -A $WORKDIR)" ]; then
    echo "Target directory $WORKDIR not empty"
    exit 2
fi

# install directory must be empty
if [ "$(ls -A $INSTALL_DIR)" ]; then
    echo "Target directory $INSTALL_DIR not empty"
    exit 2
fi

# prep files
cd $WORKDIR
if [ -d amber${VERSION}_src ]; then
    mv -f amber${VERSION}_src amber_erase
    rm -rf amber_erase &
fi

bunzip2 -c ${TARBALL_DIR}/Amber${VERSION}.tar.bz2 | tar xf -
bunzip2 -c ${TARBALL_DIR}/AmberTools${TOOLSVERSION}.tar.bz2 | tar xf -

# prep python and update
cd amber${VERSION}_src
```

```

export AMBERHOME=${WORKDIR}/amber${VERSION}_src

patch -p0 < $PATCHX

# CPU serial with installation of tests
echo "[CPU serial edition]"
mkdir build_cpu_serial && cd build_cpu_serial
cmake .. \
-DCMAKE_INSTALL_PREFIX=${INSTALL_DIR} \
-DCOMPILER=GNU \
-DMPI=FALSE \
-DCUDA=FALSE \
-DINSTALL_TESTS=TRUE \
-DDOWNLOAD_MINICONDA=TRUE \
-DFORCE_INTERNAL_LIBS="arpack" \
-DBUILD_QUICK=TRUE \
-DCHECK_UPDATES=TRUE

make -j${PARALLEL} install && make clean
cd ../ && rm -rf build_cpu_serial

# mark its origin at installation directory
cd ${INSTALL_DIR}
ln -s ./miniconda ./miniforge

# ad hoc fix shebang of amber.conda
cd miniconda/bin
# ad hoc ad hoc ad hoc ad hoc
perm=$(stat -c "%a" conda)
head -n 1 ipython >> conda.new
sed -e "1d" conda >> conda.new
mv -f conda.new conda
chmod $perm conda
# ad hoc ad hoc ad hoc ad hoc

cd ${WORKDIR}/amber${VERSION}_src

# reuse installed python
AMBER_PYTHON=${INSTALL_DIR}/bin/amber.python

# CUDA, serial, gcc
echo "[GPU serial edition]"
mkdir build_gpu_serial && cd build_gpu_serial
cmake .. \
-DCMAKE_INSTALL_PREFIX=${INSTALL_DIR} \
-DCOMPILER=GNU \
-DMPI=FALSE \
-DCUDA=TRUE \
-DINSTALL_TESTS=FALSE \
-DDOWNLOAD_MINICONDA=FALSE \
-DPYTHON_EXECUTABLE=${AMBER_PYTHON} \
-DCUDA_TOOLKIT_ROOT_DIR=${CUDA_HOME} \
-DFORCE_INTERNAL_LIBS="arpack" \
-DBUILD_QUICK=TRUE \
-DCHECK_UPDATES=FALSE

make -j${PARALLEL} install && make clean
cd ../ && rm -rf build_gpu_serial

# GPU parallel
echo "[GPU parallel edition]"
mkdir build_gpu_parallel && cd build_gpu_parallel
cmake .. \
-DCMAKE_INSTALL_PREFIX=${INSTALL_DIR} \
-DCOMPILER=GNU \

```

```

-DMPI=TRUE \
-DCUDA=TRUE \
-DINSTALL_TESTS=FALSE \
-DDOWNLOAD_MINICONDA=FALSE \
-DPYTHON_EXECUTABLE=${AMBER_PYTHON} \
-DCUDA_TOOLKIT_ROOT_DIR=${CUDA_HOME} \
-DFORCE_INTERNAL_LIBS="arpack" \
-DBUILD_QUICK=TRUE \
-DCHECK_UPDATES=FALSE

make -j${PARALLEL} install && make clean
cd .. && rm -rf build_gpu_parallel

# CPU openmp
echo "[CPU openmp edition]"
mkdir build_cpu_openmp && cd build_cpu_openmp
cmake .. \
-DCMAKE_INSTALL_PREFIX=${INSTALL_DIR} \
-DCOMPILER=GNU \
-DMPI=FALSE \
-DOPENMP=TRUE \
-DCUDA=FALSE \
-DINSTALL_TESTS=FALSE \
-DDOWNLOAD_MINICONDA=FALSE \
-DPYTHON_EXECUTABLE=${AMBER_PYTHON} \
-DFORCE_INTERNAL_LIBS="arpack" \
-DBUILD_REAXFF_PUREMD=TRUE \
-DBUILD_QUICK=TRUE \
-DCHECK_UPDATES=FALSE

make -j${PARALLEL} install && make clean
cd .. && rm -rf build_cpu_openmp

# CPU mpi (don't build mpi+openmp version)
echo "[CPU parallel edition]"
mkdir build_cpu_parallel && cd build_cpu_parallel
cmake .. \
-DCMAKE_INSTALL_PREFIX=${INSTALL_DIR} \
-DCOMPILER=GNU \
-DMPI=TRUE \
-DOPENMP=FALSE \
-DCUDA=FALSE \
-DINSTALL_TESTS=FALSE \
-DDOWNLOAD_MINICONDA=FALSE \
-DPYTHON_EXECUTABLE=${AMBER_PYTHON} \
-DFORCE_INTERNAL_LIBS="arpack" \
-DBUILD_QUICK=TRUE \
-DCHECK_UPDATES=FALSE

make -j${PARALLEL} install && make clean
cd .. && rm -rf build_cpu_parallel

# ad hoc fix for shebang
cd ${INSTALL_DIR}/bin
for f in *; do
grep -d skip "#!.*python" $f > /dev/null
if [ $? -eq 0 ]; then
perm=$(stat -c "%a" $f)
head -n 1 amber.condal >> ${f}.new
sed -e "1d" ${f} >> ${f}.new
mv -f ${f}.new ${f}
chmod $perm ${f}
fi
done

```

```

# run tests
cd ${INSTALL_DIR}
. ${INSTALL_DIR}/amber.sh
# now, $AMBERHOME should be $INSTALL_DIR

# parallel tests first
export DO_PARALLEL="mpirun -np 2"

make test.parallel && make clean.test
make test.cuda.parallel && make clean.test # DPFP
cd test; ./test_amber_cuda_parallel.sh SPFP; make clean; cd ../

export DO_PARALLEL="mpirun -np 4"
cd test; make test.parallel.4proc; make clean; cd ../

unset DO_PARALLEL

# openmp tests
make test.openmp && make clean.test

# serial tests
make test.serial && make clean.test
make test.cuda.serial && make clean.test # DPFP
cd test; ./test_amber_cuda_serial.sh SPFP; make clean && cd ../

```

Tests

- Test results are available under /local/apl/lx/amber22-up0/logs.
- All the tests excluding pbsa_cuda_cg ones (see below) seem to be passed successfully.

メモ

- Performance on P100/V100 GPUs seem to be worse than amber20 by ~5%. (tested on JAC system)
 - amber18 is the fastest for P100, amber20 is the fastest for V100
 - In the current RCCS system, you might want to use amber18/20 if you don't need new functions of amber22. (There could be some updates later, though.)
- Gcc10 (devtoolset-10) failed to build pmemd.cuda. Gcc7 or 8 (devtoolset-7 or devtoolset-8) is OK.
- If cuda-11.6.1 is used, there seems to be a slight performance loss on V100 (tested on JAC system). We thus employ cuda 11.1.
 - Official benchmark results for P100/V100 are not yet available now (June 6, 2022).
- arpack is installed in our frontend nodes but not in computation nodes (ccca*). That's why we added -DFORCE_INTERNAL_LIBS="arpack" flag.
 - Generally, this flag may not be necessary. This is very specific issue to the current RCCS system.
- quick is enabled for all the versions, reaxff_puremd is enabled only for OpenMP version.
- MKL is disabled for all the versions. This is because it is difficult to use MKL only for some of versions like [the amber20-up12 case](#).
 - (Mixing MKL and non-MKL versions may be the problem. There may be some trick to overcome this...)
- pbsa_cuda_cg tests failed just alike amber20-cmake (ambertools20/21) case.
 - (reported to official ML)
 - If pbsa.cuda is built using configure script (need some modifications, though), this bug does not occur.
- Environment setting script "amber.csh" failed to locate script directory in some case. The script is manually modified and the path is hard-coded. (see file diff above)