

## AMBER20 update 13

### ウェブページ

<http://ambermd.org/>

### バージョン

Amber20 update 13, AmberTools 21 update 12

### ビルド環境

- GCC 9.2.1 (gcc-toolset-9)
- MKL 2022.2.1
- CUDA 11.2
- OpenMPI 4.1.4 (HPC-X 2.11)
  - 実際のビルド時には OpenMPI 4.1.5 (HPC-X 2.13.1 を使用)していたが、問題が発生したため、runtime だけを HPC-X 2.11 に切り替え
  - 以下の手順では HPC-X 2.11 で記述。

### ビルドに必要なファイル

- Amber20.tar.bz2
- AmberTools20.tar.bz2
  - (Amber20 update.1-13 & AmberTools20 update.1-15 & AmberTools21 update.1-12; スクリプト内で取得)
- patch-nmrat-gpu.cpp

```
--- src/pmemd/src/cuda/gpu.cpp.org 2022-01-06 16:02:15.915217989 +0900
+++ src/pmemd/src/cuda/gpu.cpp 2022-01-06 16:02:26.857121731 +0900
@@ -2849,7 +2849,7 @@
 }
 // torsions, resttype = 3
 else if (resttype[i] == 3) {
- if (nmrat[i][0] >= 0 && nmrat[i][1] >= 0 && nmrat[i][2] >= 0 && nmrat[3] >= 0) {
+ if (nmrat[i][0] >= 0 && nmrat[i][1] >= 0 && nmrat[i][2] >= 0 && nmrat[i][3] >= 0) {
   torsions++;
 }
 else {
```

- patch-cpptraj-configure

```
--- AmberTools/src/cpptraj/configure.org 2022-01-13 22:56:46.000000000 +0900
+++ AmberTools/src/cpptraj/configure 2022-01-13 22:57:10.000000000 +0900
@@ -1771,8 +1771,6 @@
 if [ "${LIB_STAT[$LPARANC]}" != 'off' -a $USE_MPI -eq 0 ]; then
   WrmMsg "Parallel NetCDF enabled but MPI not specified. Assuming '-mpi'."
   USE_MPI=1
- elif [ $USE_MPI -ne 0 -a "${LIB_STAT[$LPARANC]}" = 'off' ]; then
-   LIB_STAT[$LPARANC]='enabled'
fi
# If we are using the bundled ARPACK then we will need C++/Fortran linking.
if [ "${LIB_STAT[$LARPAC]}" = 'bundled' ]; then
```

- patch-configure\_python (miniconda のかわりに miniforge を利用)

```
--- AmberTools/src/configure_python.org 2022-01-12 15:46:09.042775250 +0900
+++ AmberTools/src/configure_python 2022-01-12 15:48:09.177986821 +0900
@@ -107,8 +107,7 @@

echo "Downloading the latest Miniconda distribution"
if [ $mac -eq 1 ]; then
- curl -L -# https://repo.continuum.io/miniconda/Miniconda${version}-${MINICONDA_VERSION}-MacOSX-x86_64.sh > \
-   miniconda.sh
+ :
```

```

else
  # Try to figure out if the machine builds 32- or 64-bit binaries,
  # respectively.
@@ -145,23 +144,23 @@
    ;;
  esac
  if [ $bit -eq 32 ]; then
-   wget https://repo.continuum.io/miniconda/Miniconda${version}-${MINICONDA_VERSION}-Linux-x86.sh \
-     -O miniconda.sh
+   exit 0
  else
-   wget https://repo.continuum.io/miniconda/Miniconda${version}-${MINICONDA_VERSION}-Linux-x86_64.sh \
-     -O miniconda.sh
+   wget https://github.com/conda-forge/miniforge/releases/download/${MINICONDA_VERSION}/Miniforge${version}-${MINICONDA_VERSION}-
Linux-x86_64.sh \
+     -O miniforge.sh
  fi
fi

-if [ -d "$prefix/miniconda" ]; then
-  echo "Deleting existing miniconda at $prefix/miniconda"
-  /bin/rm -fr "$prefix/miniconda"
+if [ -d "$prefix/miniforge" ]; then
+  echo "Deleting existing miniforge at $prefix/miniforge"
+  /bin/rm -fr "$prefix/miniforge"
fi

echo "Installing Miniconda Python."
-bash miniconda.sh -b -p "$prefix/miniconda" || error "Installing miniconda failed"
+bash miniforge.sh -b -p "$prefix/miniforge" || error "Installing miniconda failed"
+ln -s ./miniforge ./miniconda

-export PATH="$prefix/miniconda/bin":$PATH"
+export PATH="$prefix/miniforge/bin":$PATH"
echo "Updating and installing required and optional packages..."

$prefix/miniconda/bin/python -m pip install pip --upgrade
@@ -172,7 +171,7 @@
# Use pip to install matplotlib so we don't have to pull in the entire Qt
# dependency. And cache inside the Miniconda directory, since we don't want to
# be writing outside $AMBERHOME unless specifically requested to
-$prefix/miniconda/bin/python -m pip --cache-dir=$prefix/miniconda/pkgs \
+$prefix/miniconda/bin/python -m pip --cache-dir=$prefix/miniforge/pkgs \
  install matplotlib || install_matplotlib='yes'

if [ ! -z "$install_matplotlib" ]; then
@@ -183,22 +182,22 @@
mkdir -p $prefix/lib
pwd=`pwd`
cd "$prefix/bin"
-ln -sf ../miniconda/bin/python amber.python || error "Linking Amber's Miniconda Python"
-ln -sf ../miniconda/bin/conda amber.conda || error "Linking Amber's Miniconda conda"
-ln -sf ../miniconda/bin/ipython amber.ipython || error "Linking Amber's Miniconda ipython"
-ln -sf ../miniconda/bin/jupyter amber.jupyter || error "Linking Amber's Miniconda jupyter"
-ln -sf ../miniconda/bin/pip amber.pip || error "Linking Amber's Miniconda pip"
+ln -sf ../miniforge/bin/python amber.python || error "Linking Amber's Miniconda Python"
+ln -sf ../miniforge/bin/conda amber.conda || error "Linking Amber's Miniconda conda"
+ln -sf ../miniforge/bin/ipython amber.ipython || error "Linking Amber's Miniconda ipython"
+ln -sf ../miniforge/bin/jupyter amber.jupyter || error "Linking Amber's Miniconda jupyter"
+ln -sf ../miniforge/bin/pip amber.pip || error "Linking Amber's Miniconda pip"
cd "$prefix/lib"
-for dir in ../miniconda/lib/tcl*; do
+for dir in ../miniforge/lib/tcl*; do
  ln -sf "$dir" || error "Linking TCL library folder $dir"
done

```

```

-for dir in ../miniconda/lib/tk*; do
+for dir in ../miniforge/lib/tk*; do
    ln -sf "$dir" || error "Linking TK library folder $dir"
done
cd $cwd
echo ""
-echo "Done. Miniconda installed in $prefix/miniconda"
+echo "Done. Miniforge installed in $prefix/miniforge"

-/bin/rm -f miniconda.sh
+/bin/rm -f miniforge.sh

-$prefix/miniconda/bin/conda clean --all --yes
+$prefix/miniforge/bin/conda clean --all --yes

```

## ビルド手順

```

#!/bin/sh

VERSION=20
TOOLSVERSION=20 # will be upgraded to 21

MINIFORGE_VERSION="4.11.0-4" # ad hoc custom version

INSTALL_DIR="/apl/amber/20u13"
TARBALL_DIR="/home/users/${USER}/Software/AMBER/20"

PATCH0=${TARBALL_DIR}/patch-nmrat-gpu.cpp
PATCH1=${TARBALL_DIR}/patch-cpptraj-configure
PATCHX=${TARBALL_DIR}/patch-configure_python

PARALLEL=12

#-----
module purge
module load gcc-toolset/9
module load mkl/2022.2.1
module load cuda/11.2
module load openmpi/4.1.4-hpcx/gcc9

export AMBERHOME=${INSTALL_DIR}
export CUDA_HOME="/apl/cuda/11.2"

export LANG=C
export LC_ALL=C

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

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

ulimit -s unlimited

# prep files
cd ${AMBERHOME}
bunzip2 -c ${TARBALL_DIR}/Amber${VERSION}.tar.bz2 | tar xf -
bunzip2 -c ${TARBALL_DIR}/AmberTools${TOOLSVERSION}.tar.bz2 | tar xf -

```

```

mv amber${VERSION}_src/* .
rmdir amber${VERSION}_src

patch -p0 < $PATCHX
sed -i -e "s/=latest/=${MINIFORGE_VERSION}/" AmberTools/src/configure_python

# install python first. otherwise, update_amber failed to connect ambermd.org
./AmberTools/src/configure_python -v 3
AMBER_PYTHON=$AMBERHOME/bin/amber.python

# cheat /usr/bin/env python...
cd bin && ln -s amber.python ./python && cd ..
OLDPATH=${PATH}
export PATH="${AMBERHOME}/bin:${PATH}"

# apply patches and update AmberTools
echo y | $AMBER_PYTHON ./update_amber --upgrade
$AMBER_PYTHON ./update_amber --update

# remove evidence
rm -f ${AMBERHOME}/bin/python
export PATH="${OLDPATH}"

# patch
# see http://archive.ambermd.org/202110/0206.html
patch -p0 < $PATCH0
# ad hoc something for pnetcdf
patch -p0 < $PATCH1

# ad hoc fix for cuda 11.2
sed -i -e "s/11\./11.2/" AmberTools/src/configure2

# reaxff-puremd is openmp only (tentatively)
# quick is tentatively not enabled; libstdc++ related issue?
echo "[GPU serial edition (two versions)]"
LANG=C ./configure --no-updates -cuda gnu
make -j${PARALLEL} install && make clean

echo "[GPU parallel edition (two versions)]"
LANG=C ./configure --no-updates -mpi -cuda gnu
make -j${PARALLEL} install && make clean

echo "[CPU serial edition]"
LANG=C ./configure --no-updates gnu
make -j${PARALLEL} install && make clean

echo "[CPU openmp edition]"
LANG=C ./configure --no-updates -mkl -reaxff-puremd-openmp -openmp gnu
make -j${PARALLEL} install && make clean

echo "[CPU parallel edition]"
LANG=C ./configure --no-updates -mkl -mpi gnu
make -j${PARALLEL} install && make clean

# run tests
. ${AMBERHOME}/amber.sh
cd ${AMBERHOME}

# ad hoc work-around
# https://github.com/Amber-MD/pdb4amber/issues/85#issuecomment-672812778
cd ${AMBERHOME}/lib/python*/site-packages
if [ -d ParmEd*/parmed ]; then
  ln -s ParmEd*/parmed ./parmed
fi
if [ -d pdb4amber*/pdb4amber ]; then

```

```

In -s pdb4amber*/pdb4amber ./pdb4amber
fi
if [ -d pytraj*/pytraj ]; then
  In -s pytraj*/pytraj ./pytraj
fi

# parallel tests first
cd ${AMBERHOME}
export DO_PARALLEL="mpirun -np 2"
make test.parallel && make clean.test

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

cd ${AMBERHOME}
chmod 700 src

```

## テスト

- GPU のテストについてはフロントエンドノードで実行できないため、以下のスクリプトで別途実行

```

#!/bin/sh

module purge
module load gcc-toolset/9
module load mkl/2022.2.1
module load cuda/11.2
module load openmpi/4.1.4-hpcx/gcc9

export AMBERHOME="/apl/amber/20u13"
export CUDA_HOME="/apl/cuda/11.2"

export LANG=C
export LC_ALL=C

ulimit -s unlimited

# parallel tests first
cd ${AMBERHOME}
. ${AMBERHOME}/amber.sh

## gpu tests (elsewhere)
export DO_PARALLEL="mpirun -np 2"
make test.cuda_parallel && make clean.test # DPFP
cd test; ./test_amber_cuda_parallel.sh SPFP; make clean; cd ../

unset DO_PARALLEL
make test.cuda_serial && make clean.test # DPFP
cd test; ./test_amber_cuda_serial.sh SPFP; make clean; cd ../

```

- テスト結果は /apl/amber/20u13/logs/ に有り。基本的には軽微な数値エラー。

## メモ

- 前システムでのビルド時と同様に configure を利用。cmake については検討せず。
- CUDA 11.2 は configure では正式に対応されていなかったため、少し無理矢理に対応。
- HPC-X 2.13.1 では GPU での多ノード並列時にエラーが発生して計算できず。runtime を HPC-X 2.11 に切り替えることで解消。

- MMPBSA.py のパス周りに問題があり、実行に失敗。以下を MMPBSA.py, MMPBSA.py.MPI に以下の 2 行を強制的に加えることで解消
  - 前回のシステムではこのような問題は発生していない。

```
import sys
sys.path.append( "/apl/amber/20u13/AmberTools/src/mmpbsa_py/MMPBSA_mods" )
```