

Gromacs 5.1.5 for LX with GPU support (intel compiler)

ウェブページ

<http://www.gromacs.org/>

バージョン

5.1.5

ビルド環境

- ▶ Intel Compiler 2015.1.133
- ▶ Intel MPI 5.0.2
- ▶ Intel MKL 11.2.1
- ▶ CUDA 8.0.61

ビルドに必要なファイル

- ▶ gromacs 5.1.5.tar.gz
- ▶ (regressiontests-5.1.5.tar.gz; テストセット)

ビルド手順

```
#!/bin/sh
VERSION=5.1.5
GROMACS_TARBALL=/home/users/${USER}/gromacs-${VERSION}.tar.gz
#REGRESSION=/home/users/${USER}/regressiontests-${VERSION} # unpacked
INSTALL_PREFIX=/local/apl/lx/gromacs515-CUDA
WORKDIR=/work/users/${USER}
#-----
umask 0022
cd ${WORKDIR}
if [ -d gromacs-${VERSION} ]; then
  mv gromacs-${VERSION} gromacs_erase
  rm -rf gromacs_erase &
fi
tar xzf ${GROMACS_TARBALL}
cd gromacs-${VERSION}
# intel 15 or 16 is required for cuda-8.0
. /local/apl/lx/intel2015update1/bin/compilervars.sh intel64
# compiler setting for single node versions
export CC=icc
export CXX=icpc
export F77=ifort
export F90=ifort
export FC=ifort
# single precision, no MPI
mkdir rccs-cuda
cd rccs-cuda
cmake .. \
  -DCMAKE_INSTALL_PREFIX=${INSTALL_PREFIX} \
  -DCMAKE_VERBOSE_MAKEFILE=ON \
  -DGMX_MPI=OFF \
  -DGMX_GPU=ON \
  -DGMX_DOUBLE=OFF \
  -DGMX_THREAD_MPI=ON \
  -DGMX_FFT_LIBRARY=mkl \
  -DREGRESSIONTEST_DOWNLOAD=OFF
make -j12 && make install
cd ..
# corresponding version of intel MPI
. /local/apl/lx/intel2015update1/mpi/5.0.2.044/intel64/bin/mpivars.sh
# compiler setting for MPI versions
export CC=mpiicc
export CXX=mpiicpc
export F77=mpiifort
export F90=mpiifort
export FC=mpiifort
# single precision, with MPI
mkdir rccs-cuda-mpi
cd rccs-cuda-mpi
cmake .. \
```

```
-DCMAKE_INSTALL_PREFIX=${INSTALL_PREFIX} \  
-DCMAKE_VERBOSE_MAKEFILE=ON \  
-DGMX_MPI=ON \  
-DGMX_GPU=ON \  
-DGMX_DOUBLE=OFF \  
-DGMX_THREAD_MPI=OFF \  
-DGMX_FFT_LIBRARY=mkl \  
-DREGRESSIONTEST_DOWNLOAD=OFF  
make -j12 && make install  
cd ..
```
