## Quantum Espresso 6.7 with GPU support

# Webpage

https://www.quantum-espresso.org/ https://gitlab.com/QEF/q-e-gpu/-/releases

#### Version

6.7-gpu

### **Build Environment**

- PGI 20.4
- MKL 2019.0.5 (intel 2019 update 5)
- CUDA 10.1

## Files Required

- q-e-gpu-qe-gpu-6.7.tar.gz
- openmpi-4.0.2.tar.bz2
- (PBS Pro files under /local/apl/lx/pbs14)

### **Build Procedure**

```
#!/bin/sh
VERSION=6.7
FULLVER=${VERSION}
BASEDIR=/home/users/${USER}/Software/QE/${VERSION}-gpu
TARBALL=${BASEDIR}/q-e-gpu-qe-gpu-${FULLVER}.tar.gz
INSTDIR=/local/apl/lx/espresso67-gpu
# openmpi
WORKDIR=/work/users/${USER}
OMPIVER=4.0.2
OMPIROOT=${INSTDIR}/openmpi-${OMPIVER}
PBSROOT=/local/apl/lx/pbs14
PARALLEL=12
umask 0022
module purge
module load pgi/20.4
module load mkl/2019.0.5
export LANG=C
export LC_ALL=C
ulimit -s unlimited
# build openmpi first
cd ${WORKDIR}
if [ -d openmpi-${OMPIVER} ]; then
mv openmpi-${OMPIVER} openmpi-erase
rm -rf openmpi-erase &
fi
tar jxf ${OMPITARBALL}
cd openmpi-${OMPIVER}
mkdir rccs && cd rccs
```

```
CC=pgcc CXX=pgc++ FC=pgfortran \
 ../configure --prefix=${OMPIROOT} \
        --with-tm=${PBSROOT} \
        --enable-mpi-cxx \
        --enable-mpi1-compatibility \
        --with-psm2
make -j ${PARALLEL} && make install && make check
# openmpi setting
export OMPI MCA btl openib allow ib=1
export CPATH="${OMPIROOT}/include:${CPATH}"
export LIBRARY_PATH="${OMPIROOT}/lib:${LIBRARY_PATH}"
export LD_LIBRARY_PATH="${OMPIROOT}/lib:${LD_LIBRARY_PATH}"
export PATH="${OMPIROOT}/bin:${PATH}"
# qe build
cd ${INSTDIR}
if [ -d q-e-gpu-qe-gpu-${FULLVER} ]; then
mv q-e-gpu-qe-gpu-${FULLVER} q-e-gpu-qe-gpu-erase
rm -rf q-e-gpu-qe-gpu-erase &
tar zxf ${TARBALL}
cd q-e-gpu-qe-gpu-${FULLVER}
mv * .[a-zA-Z]* ../
cd ../ && rmdir q-e-gpu-qe-gpu-${FULLVER}
export MPIF90=mpif90
export MPIFC=mpif90
export MPIF77=mpif90
export MPICC=mpicc
export MPICXX=mpicxx
# --with-cuda should point cuda bundled with pgi... but i couldn't do it...
FC=pgfortran F90=pgfortran F77=pgfortran CC=pgcc CXX=pgc++\
./configure --enable-openmp \
       --enable-parallel \
        --with-cuda=/local/apl/lx/cuda-10.1 \
        --with-cuda-cc=60 \
        --with-cuda-runtime=10.1
# force to add cc70 support
sed -i -e "s/cc60/cc60,cc70/" make.inc
make -j${PARALLEL} pw cp
cd test-suite
make run-tests-pw-serial
make run-tests-cp-serial
make clean
make run-tests-pw-parallel
make run-tests-cp-parallel
cd ..
```

### Notes

- There were errors on atomic\_cmpset\* tests of OpenMPI. However, all the QE tests have passed successfully.
- Many functions of pw.x seem to be supported by GPU.
- (GPU version of cp does not seem to be very useful for now.)
  - $\circ~$  Limited parts such as fft can be performed by GPUs.
  - Conjugate gradient (cg) is not yet supported.
- OpenMP is enabled in this build.
- It works both on P100 and V100.
- On MPI parallel, single GPU would be assigned to single process.
  - Processes can share single GPU, but it may not be very effective. (Two processes on single GPU might be

advantageous in some situation.)

- Please use mpirun in /local/apl/lx/espresso67-gpu/openmpi-4.0.2 when you perform MPI parallel runs. (Please check sample job script under espresso67-gpu/samples/.)
- Some of calcul; ations might not be supported by GPU. Please check official document or the output file (timing information at the end).
- Very small calculation may not be significantly accelerated by GPU.
- Large calculation may fail due to memory allocation error (probably of GPU memory). This error may be avoided by using multiple GPUs with MPI parallel run.