XCONFIGURE

XCONFIGURE is a collection of configure wrapper scripts for various HPC applications. The purpose of the scripts is to configure the application in question to make use of Intel's software development tools (Intel Compiler, Intel MPI, Intel MKL). This may sound cumbersome, but it actually helps to rely on a "build recipe", which is known to expose the highest performance or to reliably complete the build process.

Each application (or library) is hosted in a separate subdirectory. In order to configure (and ultimately build) an application, one may clone or download the entire collection.

```
git clone https://github.com/hfp/xconfigure.git
```

Alternatively, one can rely on a single script which then downloads a specific wrapper into the current working directory (of the desired application).

```
wget https://github.com/hfp/xconfigure/raw/master/configure-get.sh
chmod +x configure-get.sh
./configure-get.sh qe hsw
```

To configure an application, please follow into one of the aforementioned subfolders and read the build recipe of this application e.g., qe in case of Quantum Espresso.

Similar Prjoects

- Spack Package Manager: http://computation.llnl.gov/projects/spack-hpc-package-manager
- EasyBuild / EasyConfig (University of Gent): https://github.com/hpcugent

CP2K

Build and Run Instructions

The build and run instructions for CP2K using Intel Tools are exercised at https://github.com/hfp/libxsmm/tree/master/documentation/cp2k.md (pdf).

Please note, in terms of functionality it is beneficial to rely on LIBINT and LIBXC, whereas ELPA eventually provides an improved performance.

Sanity Check

There is nothing that can replace the full regression test suite to be clear. However in order to quickly check whether a build is sane or not, one can run for instance tests/QS/benchmark/H2O-64.inp and check if the SCF iteration prints like the following:

Step		Update	method	Time	Convergence	Total energy	Change
1	ОТ	DIIS	0.15E+00	0.5	0.01337191	-1059.6804814927	-1.06E+03
2	OT	DIIS	0.15E+00	0.3	0.00866338	-1073.3635678409	-1.37E+01
3	OT	DIIS	0.15E+00	0.3	0.00615351	-1082.2282197787	-8.86E+00
4	OT	DIIS	0.15E+00	0.3	0.00431587	-1088.6720379505	-6.44E+00
5	OT	DIIS	0.15E+00	0.3	0.00329037	-1092.3459788564	-3.67E+00
6	OT	DIIS	0.15E+00	0.3	0.00250764	-1095.1407783214	-2.79E+00
7	OT	DIIS	0.15E+00	0.3	0.00187043	-1097.2047924571	-2.06E+00
8	OT	DIIS	0.15E+00	0.3	0.00144439	-1098.4309205383	-1.23E+00
9	OT	DIIS	0.15E+00	0.3	0.00112474	-1099.2105625375	-7.80E-01
10	OT	DIIS	0.15E+00	0.3	0.00101434	-1099.5709299131	-3.60E-01
Г	٦						

The column called "Convergence" has to monotonically converge towards zero.

Eigenvalue SoLvers for Petaflop-Applications (ELPA)

Build Instructions

Download and unpack ELPA, and make the configure wrapper scripts available in ELPA's root folder. It is recommended to package the state (Tarball or similar), which is achieved after downloading the wrapper scripts. It appears that ELPA's make clean (or similar Makefile target) is cleaning up the entire directory including all "non-ELPA content" (the directory still remains unclean enough to make subsequent builds unsuccessful).

```
wget http://elpa.mpcdf.mpg.de/html/Releases/2016.11.001.pre/elpa-2016.11.001.pre.tar.gz
tar xvf elpa-2016.11.001.pre.tar.gz
cd elpa-2016.11.001.pre
wget https://github.com/hfp/xconfigure/raw/master/configure-get.sh
chmod +x configure-get.sh
./configure-get.sh elpa
```

Please make the Intel Compiler available on the command line. This actually depends on the environment. For instance, many HPC centers rely on module load.

```
source /opt/intel/compilers_and_libraries_2017.0.098/linux/bin/compilervars.sh intel64
```

For example, to configure and make for an Intel Xeon E5v4 processor (formerly codenamed "Broadwell"):

```
./configure-elpa-hsw-omp.sh
make -j; make install
```

For different targets (instruction set extensions) or different versions of the Intel Compiler, the configure scripts support an additional argument ("default" is the default tagname):

```
./configure-elpa-hsw-omp.sh tagname
```

As shown above, an arbitrary "tagname" can be given (without editing the script). This might be used to build multiple variants of the ELPA library.

References

https://software.intel.com/en-us/articles/quantum-espresso-for-the-intel-xeon-phi-processor

LIBINT

Build Instructions

Download and unpack LIBINT, and make the configure wrapper scripts available in LIBINT's root folder.

```
wget https://github.com/evaleev/libint/archive/release-1-1-6.tar.gz
tar xvf release-1-1-6.tar.gz
cd libint-release-1-1-6
wget https://github.com/hfp/xconfigure/raw/master/configure-get.sh
chmod +x configure-get.sh
./configure-get.sh libint
```

Please make the Intel Compiler available on the command line. This actually depends on the environment. For instance, many HPC centers rely on module load.

For example, to configure and make for an Intel Xeon E5v4 processor (formerly codenamed "Broadwell"):

```
./configure-libint-hsw.sh
make -j ; make install
```

The version 1.x line of LIBINT does not support to cross-compile for an architecture (a future version of the wrapper scripts may patch this ability into LIBINT 1.x). Therefore, one might rely on the Intel Software Development Emulator (Intel SDE) to compile LIBINT for targets, which cannot execute on the compile-host.

```
/software/intel/sde/sde -knl -- make
```

To speed-up compilation, "make" might be carried out in phases: after "printing the code" (c-files), the make execution continues with building the object-file where no SDE needed. The latter phase can be sped up by interrupting "make", and executing it without SDE. The root cause of the entire problem is that the driver printing the c-code is (needlessly) compiled using the architecture-flags that are not supported on the host.

Further, for different targets (instruction set extensions) or different versions of the Intel Compiler, the configure scripts support an additional argument ("default" is the default tagname):

```
./configure-libint-hsw.sh tagname
```

As shown above, an arbitrary "tagname" can be given (without editing the script). This might be used to build multiple variants of the LIBINT library.

References

TBD

LIBXC

Build Instructions

TBD

http://www.tddft.org/programs/octopus/wiki/index.php/Libxc:download http://www.tddft.org/programs/octopus/down.php?file=libxc/libxc-3.0.0.tar.gz

References

TBD

LIBXSMM

LIBXSMM is a library targeting Intel Architecture (x86) for small, dense or sparse matrix multiplications, and small convolutions. The build instructions can be found at https://github.com/hfp/libxsmm (pdf).

Quantum Espresso (QE)

Build Instructions

Download and unpack Quantum Espresso, and make the configure wrapper scripts available in QE's root folder. Please note that the configure wrapper scripts support QE 6.0 and QE 6.1 (prior support for 5.x is dropped). Before building QE, one needs to complete the ELPA build recipe!

```
wget http://www.qe-forge.org/gf/download/frsrelease/240/1075/qe-6.1.tar.gz
tar xvf qe-6.1.tar.gz
cd qe-6.1
wget https://github.com/hfp/xconfigure/raw/master/configure-get.sh
chmod +x configure-get.sh
./configure-get.sh qe
```

Please make the Intel Compiler available on the command line, which may vary with the computing environment. For instance, many HPC centers rely on module load.

```
source /opt/intel/compilers_and_libraries_2017.0.098/linux/bin/compilervars.sh intel64
```

For example, configure for an Intel Xeon E5v4 processor (formerly codenamed "Broadwell"), and build the desired application(s) e.g., "pw", "cp", or "all".

```
./configure-qe-hsw-omp.sh
make pw -j
```

Building "all" (or make without target argument) requires to repeat make all until no compilation error occurs. This is because of some incorrect build dependencies (build order issue which might have been introduced by the configure wrapper scripts). In case of starting over, one can run make distclean, reconfigure the application, and build it again. For different targets (instruction set extensions) or different versions of the Intel Compiler, the configure scripts support an additional argument ("default" is the default tagname):

```
./configure-qe-hsw-omp.sh tagname
```

As shown above, an arbitrary "tagname" can be given (without editing the script). This might be used to build multiple variants of QE. Please note: this tagname also selects the corresponding ELPA library (or should match the tagname used to build ELPA). Make sure to save your current QE build before building an additional variant!

Run Instructions

To run Quantum Espresso in an optimal fashion depends on the workload and on the "parallelization levels", which can be exploited by the workload in question. These parallelization levels apply to execution phases (or major algorithms) rather than staying in a hierarchical relationship (levels). It is recommended to read some of the primary references explaining these parallelization levels (a number of them can be found in the Internet including some presentation slides). Time to solution may vary by factors depending on whether these levels are orchestrated or not. To specify these levels, one uses command line arguments along with the QE executable(s):

- -npool: try to maximize the number of pools. The number depends on the workload e.g., if the number of k-points can be distributed among independent pools. Indeed, trial and error is a rather quick to check if workload fails to pass the initialization phase. One may use prime numbers: 2, 3, 5, etc. (default is 1). For example, when npool=2 worked it might be worth trying npool=4. On the other hand, increasing the number pools duplicates the memory consumption accordingly (larger numbers are increasingly unlikely to work).
- -ndiag: this number determines the number of ranks per pool used for dense linear algebra operations (DGEMM and ZGEMM). For example, if 64 ranks are used in total per node and npool=2, then put ndiag=32 (QE selects the next square number which is less-equal than the given number e.g., ndiag=25 in the previous example).
- -ntg: specifies the number of tasks groups per pool being used for e.g., FFTs. One can start with NTG=\$((NUMNODES*NRANKS/(NPOOL*2))). If NTG becomes zero, NTG=\${NRANKS} should be used (number of ranks per node). Please note the given formula is only a rule of thumb, and the number of task groups also depends on the number of ranks as the workload is scaled out.

To run QE, below command line can be a starting point ("numbers" are presented as Shell variables to better understand the inner mechanics). Important for hybrid builds (MPI and OpenMP together) are the given environment variables. The kmp_affinity assumes Hyperthreaded is enabled (granularity=fine), and the "scatter" policy allows to easily run less than the maximum number of Hyperthreads per core. As a rule of thumb, OpenMP adds only little overhead (often not worth a pure MPI application) but allows to scale further out when compared to pure MPI builds.

```
mpirun -bootstrap ssh -genvall \
  -np $((NRANKS_PER_NODE*NUMNODES)) -perhost ${NRANKS} \
  -genv I_MPI_PIN_DOMAIN=auto \
  -genv KMP_AFFINITY=scatter,granularity=fine,1 \
  -genv OMP_NUM_THREADS=${NTHREADS_PER_RANK} \
  /path/to/pw.x \<command-line-arguments\>
```

References

 $https://software.intel.com/en-us/articles/quantum-espresso-for-the-intel-xeon-phi-processor \\ http://www.quantum-espresso.org/wp-content/uploads/Doc/user_guide/node18.html$