Quad-Double computation package Copyright (C) 2003-2012

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To build this library, follow the steps below. Some system specific notes are at the end of this file.

Build Instructions ______

1. Run the configure script by typing

./configure

The script will attempt to automatically detect various system-dependent variables used during compilation (such as the C++/fortran compiler, compiler flags, and linker flags).

If you want to specify a particular C++ / F90 compiler and their flags, you can set them as environmental variables. For example:

FC=ifc FCFLAGS="-02 -FR" ./configure

Important variables are

C++ compiler to use CXXFLAGS C++ compiler flags to use C compiler to use (for C demo program) CFLAGS C compiler flags to use (for C demo program) FC

Fortran 90 compiler

FCFLAGS Fortran 90 compiler flags to use

FCLIBS Fortran 90 libraries needed to to link with C++ code.

See ./configure --help to see other options.

- 3. The configure script should also have created the files 'config.h' and 'include/qd/qd config.h', which will contain the compile time defines. Examine these and edit them if necessary. In most cases no edits are necessary, since the options are detected when configure was run.
- 4. Type "make". This will build the library, and necessary Fortran wrappers.
- 5. Optionally, one can build and run some simple test programs. To do this, type "make check". Some programs run during this phase is a good demonstration of how to use the qd library in C++.
- 6. You can now install the QD library by issuing "make install".
- 7. If you want to build some sample programs written in C++ you can type "make cpp-demo".

- 8. If you want to build some sample programs written in Fortran 90, you can type "make fortran-demo".
- 9. If you want to compile the Experimental Mathematician's Toolkit, type "make toolkit". This will compile the Fortran-90 codes in the toolkit cirectory, including the "mathinit" and "mathtool" Read the "README" file in the toolkit directory for additional details.

System-Specific Notes

Linux x86 / Itanium

You can use g++ to compile the C++ code. The Fortran 90 codes can be compiled using Intel Fortran 95 compiler

http://www.intel.com/software/products/compilers/flin/

available freely for non-commercial uses. There is also a C++ compiler available (for non-commercial use) at

http://www.intel.com/software/products/compilers/clin/

which can be used to compile the C++ portion. By default the configure script will use the Intel compiler if found.

Apple (OS X)

For Apple OS X Intel-badsed systems, it is recommended that you use the g++-4.0 (or higher) compiler and the gfortran compiler. The g++-4.0 (or higher) compiler and related command-line tools are now available via this URL (see command-line tools): https://developer.apple.com/downloads/index.action

The gfortran compiler can be downloaded from: http://www.macresearch.org/files/gfortran/gfortran-4.3-Nov.mpkg.zip

After installing these compilers, in the main gd directory type

./configure CXX=q++ FC=qfortran FCFLAGS=-m64

then type "make" to construct the library. See the "README" file on how to construct a compile-link script for your own codes.

IBM (Power)

With IBM's xlC/xlf90 compilers, you may want to experiment with --enable-fma option which uses a faster code but relies on the compiler to generate a fused multiply-accumulate instruction. WARNING: since the compiler is not required to produce such instructions, this is not guaranteed to work. Please test before using.