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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

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Getting Help and Support

Intel provides a support web site that contains a rich repository of self help information, including getting started tips, known product issues, product errata, license information, user forums, and more. Visit the Intel® MKL support website at http://www.intel.com/software/products/support/.

You can get context-sensitive help when editing your code in the Microsoft Visual Studio® integrated development environment (IDE). See Getting Assistance for Programming in the Microsoft Visual Studio® IDE for details.
Introducing the Intel® Math Kernel Library

Intel® Math Kernel Library (Intel® MKL) is a computing math library of highly optimized, extensively threaded routines for applications that require maximum performance. The library provides Fortran and C programming language interfaces. Intel® MKL C language interfaces can be called from applications written in either C or C++, as well as in any other language that can reference a C interface.

Intel® MKL provides comprehensive functionality support in these major areas of computation:

• BLAS (level 1, 2, and 3) and LAPACK linear algebra routines, offering vector, vector-matrix, and matrix-matrix operations.
• ScalAPACK distributed processing linear algebra routines, as well as the Basic Linear Algebra Communications Subprograms (BLACS) and the Parallel Basic Linear Algebra Subprograms (PBLAS).
• Intel® MKL PARDISO (a direct sparse solver based on Parallel Direct Sparse Solver PARDISO*), an iterative sparse solver, and supporting sparse BLAS (level 1, 2, and 3) routines for solving sparse systems of equations, as well as a distributed version of Intel® MKL PARDISO solver provided for use on clusters.
• Fast Fourier transform (FFT) functions in one, two, or three dimensions with support for mixed radices (not limited to sizes that are powers of 2), as well as distributed versions of these functions provided for use on clusters.
• Vector Mathematics (VM) routines for optimized mathematical operations on vectors.
• Vector Statistics (VS) routines, which offer high-performance vectorized random number generators (RNG) for several probability distributions, convolution and correlation routines, and summary statistics functions.
• Data Fitting Library, which provides capabilities for spline-based approximation of functions, derivatives and integrals of functions, and search.
• Extended Eigensolver, a shared memory programming (SMP) version of an eigensolver based on the Feast Eigenvalue Solver.

For details see the Intel® MKL Developer Reference.

Intel® Math Kernel Library (Intel® MKL) is optimized for performance on Intel processors. Intel® MKL also runs on non-Intel x86-compatible processors.

For Windows* and Linux* systems based on Intel® 64 Architecture, Intel® MKL also includes support for the Intel® Many Integrated Core Architecture (Intel® MIC Architecture) and provides libraries to help you port your applications to Intel MIC Architecture.

NOTE

Intel® MKL provides limited input validation to minimize the performance overheads. It is your responsibility when using Intel® MKL to ensure that input data has the required format and does not contain invalid characters. These can cause unexpected behavior of the library. Examples of the inputs that may result in unexpected behavior:

• Not-a-number (NaN) and other special floating point values
• Large inputs may lead to accumulator overflow

As the Intel® MKL API accepts raw pointers, it is your application's responsibility to validate the buffer sizes before passing them to the library. The library requires subroutine and function parameters to be valid before being passed. While some Intel® MKL routines do limited checking of parameter errors, your application should check for NULL pointers, for example.
Optimization Notice

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.
What's New

This Developer Guide documents Intel® Math Kernel Library 2020 Update 1. The Developer Guide has been updated to fix inaccuracies in the document.
Notational Conventions

The following term is used in reference to the operating system.

Windows* This term refers to information that is valid on all supported Windows* operating systems.

The following notations are used to refer to Intel® MKL directories.

<parent directory> The installation directory that includes Intel® MKL directory; for example, the directory for Intel® Parallel Studio XE Composer Edition.

<mkl directory> The main directory where Intel® MKL is installed:

<mkl directory> = <parent directory>

Replace this placeholder with the specific pathname in the configuring, linking, and building instructions.

The following font conventions are used in this document.

Italic Italic is used for emphasis and also indicates document names in body text, for example:

see Intel® MKL Developer Reference.

Monospace lowercase mixed with uppercase Indicates:

• Commands and command-line options, for example,

  ifort myprog.f mkl_blas95.lib mkl_c.lib libiomp5md.lib

• Filenames, directory names, and pathnames, for example,

  C:\Program Files\Java\jdk1.5.0_09

• C/C++ code fragments, for example,

  a = new double [SIZE*SIZE];

UPPERCASE MONOSPACE Indicates system variables, for example, $MKLPATH.

Monospace italic Indicates a parameter in discussions, for example, lda.

When enclosed in angle brackets, indicates a placeholder for an identifier, an expression, a string, a symbol, or a value, for example, <mkl directory>. Substitute one of these items for the placeholder.

[ items ] Square brackets indicate that the items enclosed in brackets are optional.

{ item | item } Braces indicate that only one of the items listed between braces should be selected. A vertical bar ( | ) separates the items.
Related Information

To reference how to use the library in your application, use this guide in conjunction with the following documents:

- The *Intel® Math Kernel Library Developer Reference*, which provides reference information on routine functionalities, parameter descriptions, interfaces, calling syntaxes, and return values.
- The *Intel® Math Kernel Library for Windows® OS Release Notes*. 
Getting Started

Optimization Notice

Intel’s compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

Checking Your Installation

After installing the Intel® Math Kernel Library (Intel® MKL), verify that the library is properly installed and configured:

1. Intel® MKL installs in the <parent directory> directory.
   
   Check that the subdirectory of <parent directory> referred to as <mkl directory> was created.
   
   Check that subdirectories for Intel® MKL redistributable DLLs redist\ia32_win\mkl and redist\intel64_win\mkl were created in the <parent directory> directory (See redist.txt in the Intel® MKL documentation directory for a list of files that can be redistributed.)

2. If you want to keep multiple versions of Intel® MKL installed on your system, update your build scripts to point to the correct Intel® MKL version.

3. Check that the mklvars.bat file appears in the <mkl directory>\bin directory.
   
   Use this file to assign Intel® MKL-specific values to several environment variables, as explained in Scripts to Set Environment Variables Setting Environment Variables.

4. To understand how the Intel® MKL directories are structured, see Structure of the Intel® Math Kernel Library.

5. To make sure that Intel® MKL runs on your system, launch an Intel® MKL example, as explained in Using Code Examples.

See Also
Notational Conventions

Setting Environment Variables

When the installation of Intel® MKL for Windows® is complete, set the PATH, LIB, and INCLUDE environment variables in the command shell using the mklvars.bat script in the bin subdirectory of the Intel® MKL installation directory.

The script accepts the parameters, explained in the following table:

<table>
<thead>
<tr>
<th>Setting Specified</th>
<th>Required (Yes/No)</th>
<th>Possible Values</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Architecture</td>
<td>Yes</td>
<td>intel64</td>
<td></td>
</tr>
<tr>
<td>Setting Specified</td>
<td>Required (Yes/No)</td>
<td>Possible Values</td>
<td>Comment</td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------</td>
<td>-----------------</td>
<td>---------</td>
</tr>
<tr>
<td>Use of Intel® MKL Fortran modules precompiled with the Intel® Visual Fortran compiler</td>
<td>No</td>
<td>mod</td>
<td>Supply this parameter only if you are using this compiler.</td>
</tr>
<tr>
<td>Programming interface (LP64 or ILP64)</td>
<td>No</td>
<td>lp64, default ilp64</td>
<td></td>
</tr>
</tbody>
</table>

For example:

- The command
  
mklvars intel64
  
sets the environment for Intel® MKL to use the Intel 64 architecture.

- The command
  
mklvars intel64 mod ilp64
  
sets the environment for Intel® MKL to use the Intel 64 architecture, ILP64 programming interface, and Fortran modules.

- The command
  
mklvars intel64 mod
  
sets the environment for Intel® MKL to use the Intel 64 architecture, LP64 interface, and Fortran modules.

**NOTE**
Supply the parameter specifying the architecture first, if it is needed. Values of the other two parameters can be listed in any order.

See Also
High-level Directory Structure
Intel® MKL Interface Libraries and Modules
Fortran 95 Interfaces to LAPACK and BLAS
Setting the Number of Threads Using an OpenMP* Environment Variable

## Compiler Support

Intel® MKL supports compilers identified in the *Release Notes*. However, the library has been successfully used with other compilers as well.

Although Compaq no longer supports the Compaq Visual Fortran* (CVF) compiler, Intel® MKL still preserves the CVF interface in the IA-32 architecture implementation. You can use this interface with the Intel® Fortran Compiler. Intel® MKL provides both stdcall (default CVF interface) and cdecl (default interface of the Microsoft Visual C* application) interfaces for the IA-32 architecture.

When building Intel® MKL code examples, you can select a compiler:

- For Fortran examples: Intel® or PGI* compiler
- For C examples: Intel, Microsoft Visual C++*, or PGI compiler

Intel® MKL provides a set of include files to simplify program development by specifying enumerated values and prototypes for the respective functions. Calling Intel® MKL functions from your application without an appropriate include file may lead to incorrect behavior of the functions.


**See Also**
Compiling an Application that Calls Intel® MKL and Uses the CVF Calling Conventions
Using the cdecl and stdcall Interfaces
Intel® MKL Include Files

## Using Code Examples

The Intel® MKL package includes code examples, located in the `examples` subdirectory of the installation directory. Use the examples to determine:

- Whether Intel® MKL is working on your system
- How you should call the library
- How to link the library

If an Intel® MKL component that you selected during installation includes code examples, these examples are provided in a separate archive. Extract the examples from the archives before use.

For each component, the examples are grouped in subdirectories mainly by Intel® MKL function domains and programming languages. For instance, the `blas` subdirectory (extracted from the `examples_core` archive) contains a makefile to build the BLAS examples and the `vmlc` subdirectory contains the makefile to build the C examples for Vector Mathematics functions. Source code for the examples is in the next-level `sources` subdirectory.

**See Also**
High-level Directory Structure

## What You Need to Know Before You Begin Using the Intel® Math Kernel Library

<table>
<thead>
<tr>
<th>Target platform</th>
<th>Identify the architecture of your target machine:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• IA-32 or compatible</td>
</tr>
<tr>
<td></td>
<td>• Intel® 64 or compatible</td>
</tr>
</tbody>
</table>

**Reason:** Because Intel® MKL libraries are located in directories corresponding to your particular architecture (see Architecture Support), you should provide proper paths on your link lines (see Linking Examples). To configure your development environment for the use with Intel® MKL, set your environment variables using the script corresponding to your architecture (see Scripts to Set Environment Variables Setting Environment Variables for details).

<table>
<thead>
<tr>
<th>Mathematical problem</th>
<th>Identify all Intel® MKL function domains that you require:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• BLAS</td>
</tr>
<tr>
<td></td>
<td>• Sparse BLAS</td>
</tr>
<tr>
<td></td>
<td>• LAPACK</td>
</tr>
<tr>
<td></td>
<td>• PBLAS</td>
</tr>
<tr>
<td></td>
<td>• ScaLAPACK</td>
</tr>
<tr>
<td></td>
<td>• Sparse Solver routines</td>
</tr>
<tr>
<td></td>
<td>• Parallel Direct Sparse Solvers for Clusters</td>
</tr>
<tr>
<td></td>
<td>• Vector Mathematics functions (VM)</td>
</tr>
<tr>
<td></td>
<td>• Vector Statistics functions (VS)</td>
</tr>
<tr>
<td></td>
<td>• Fourier Transform functions (FFT)</td>
</tr>
<tr>
<td></td>
<td>• Cluster FFT</td>
</tr>
<tr>
<td></td>
<td>• Trigonometric Transform routines</td>
</tr>
<tr>
<td></td>
<td>• Poisson, Laplace, and Helmholtz Solver routines</td>
</tr>
<tr>
<td></td>
<td>• Optimization (Trust-Region) Solver routines</td>
</tr>
</tbody>
</table>
• Data Fitting Functions
• Extended Eigensolver Functions

**Reason:** The function domain you intend to use narrows the search in the *Intel® MKL Developer Reference* for specific routines you need. Additionally, if you are using the Intel® MKL cluster software, your link line is function-domain specific (see *Working with the Intel® Math Kernel Library Cluster Software*). Coding tips may also depend on the function domain (see *Other Tips and Techniques to Improve Performance*).

**Programming language**

Intel® MKL provides support for both Fortran and C/C++ programming. Identify the language interfaces that your function domains support (see *Appendix A: Intel® Math Kernel Library Language Interfaces Support*).

**Reason:** Intel® MKL provides language-specific include files for each function domain to simplify program development (see *Language Interfaces Support _by Function Domain*).

For a list of language-specific interface libraries and modules and an example how to generate them, see also *Using Language-Specific Interfaces with Intel® Math Kernel Library*.

**Range of integer data**

If your system is based on the Intel 64 architecture, identify whether your application performs calculations with large data arrays (of more than $2^{31}-1$ elements).

**Reason:** To operate on large data arrays, you need to select the ILP64 interface, where integers are 64-bit; otherwise, use the default, LP64, interface, where integers are 32-bit (see *Using the ILP64 Interface vs*).

**Threading model**

Identify whether and how your application is threaded:

• Threaded with the Intel compiler
• Threaded with a third-party compiler
• Not threaded

**Reason:** The compiler you use to thread your application determines which threading library you should link with your application. For applications threaded with a third-party compiler you may need to use Intel® MKL in the sequential mode (for more information, see *Linking with Threading Libraries*).

**Number of threads**

If your application uses an OpenMP* threading run-time library, determine the number of threads you want Intel® MKL to use.

**Reason:** By default, the OpenMP* run-time library sets the number of threads for Intel® MKL. If you need a different number, you have to set it yourself using one of the available mechanisms. For more information, see *Improving Performance with Threading*.

**Linking model**

Decide which linking model is appropriate for linking your application with Intel® MKL libraries:

• Static
• Dynamic

**Reason:** The link libraries for static and dynamic linking are different. For the list of link libraries for static and dynamic models, linking examples, and other relevant topics, like how to save disk space by creating a custom dynamic library, see *Linking Your Application with the Intel® Math Kernel Library*.

**MPI used**

Decide what MPI you will use with the Intel® MKL cluster software. You are strongly encouraged to use the latest available version of Intel® MPI.

**Reason:** To link your application with ScaLAPACK and/or Cluster FFT, the libraries corresponding to your particular MPI should be listed on the link line (see *Working with the Intel® Math Kernel Library Cluster Software*).
Structure of the Intel® Math Kernel Library

Optimization Notice

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

Architecture Support

Intel® Math Kernel Library (Intel® MKL) for Windows® provides architecture-specific implementations for supported platforms. The following table lists the supported architectures and directories where each architecture-specific implementation is located.

<table>
<thead>
<tr>
<th>Architecture</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>IA-32 or compatible</td>
<td><code>&lt;mkl directory&gt;</code>\lib\ia32_win</td>
</tr>
<tr>
<td></td>
<td><code>&lt;mkl directory&gt;</code>\redist\ia32_win\mkl(DLLs)</td>
</tr>
<tr>
<td>Intel® 64 or compatible</td>
<td><code>&lt;mkl directory&gt;</code>\lib\intel64_win</td>
</tr>
<tr>
<td></td>
<td><code>&lt;mkl directory&gt;</code>\redist\intel64_win\mkl(DLLs)</td>
</tr>
</tbody>
</table>

See Also
High-level Directory Structure
Notational Conventions
Detailed Structure of the IA-32 Architecture Directories
Detailed Structure of the Intel® 64 Architecture Directories

High-level Directory Structure

<table>
<thead>
<tr>
<th>Directory</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>&lt;mkl directory&gt;</code></td>
<td>Installation directory of the Intel® Math Kernel Library (Intel® MKL)</td>
</tr>
</tbody>
</table>

Subdirectories of `<mkl directory>`

<table>
<thead>
<tr>
<th>Subdirectory</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bin</td>
<td>Batch files to set environmental variables in the user shell</td>
</tr>
<tr>
<td>\mkl\bin\pkgconfig</td>
<td>Files for the most popular configurations of the Intel® MKL</td>
</tr>
<tr>
<td>Directory</td>
<td>Contents</td>
</tr>
<tr>
<td>--------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>benchmarks\linpack</td>
<td>Shared-Memory (SMP) version of the LINPACK benchmark</td>
</tr>
<tr>
<td>benchmarks\mp\linpack</td>
<td>Message-passing interface (MPI) version of the LINPACK benchmark</td>
</tr>
<tr>
<td>lib\ia32\win</td>
<td>Static libraries and static interfaces to DLLs for the IA-32 architecture</td>
</tr>
<tr>
<td>lib\intel64\win</td>
<td>Static libraries and static interfaces to DLLs for the Intel® 64 architecture</td>
</tr>
<tr>
<td>examples</td>
<td>Source and data files for Intel® MKL examples. Provided in archives corresponding to Intel® MKL components selected during installation.</td>
</tr>
<tr>
<td>include</td>
<td>Include files for the library routines and examples</td>
</tr>
<tr>
<td>include\ia32</td>
<td>Fortran 95 .mod files for the IA-32 architecture and Intel Fortran compiler</td>
</tr>
<tr>
<td>include\intel64\lp64</td>
<td>Fortran 95 .mod files for the Intel® 64 architecture, Intel® Fortran compiler, and LP64 interface</td>
</tr>
<tr>
<td>include\intel64\ilp64</td>
<td>Fortran 95 .mod files for the Intel® 64 architecture, Intel Fortran compiler, and ILP64 interface</td>
</tr>
<tr>
<td>include\fftw</td>
<td>Header files for the FFTW2 and FFTW3 interfaces</td>
</tr>
<tr>
<td>interfaces\blas95</td>
<td>Fortran 95 interfaces to BLAS and a makefile to build the library</td>
</tr>
<tr>
<td>interfaces\fftw2x_cdft</td>
<td>MPI FFTW 2.x interfaces to Intel® MKL Cluster FFT</td>
</tr>
<tr>
<td>interfaces\fftw3x_cdft</td>
<td>MPI FFTW 3.x interfaces to Intel® MKL Cluster FFT</td>
</tr>
<tr>
<td>interfaces\fftw2xc</td>
<td>FFTW 2.x interfaces to the Intel® MKL FFT (C interface)</td>
</tr>
<tr>
<td>interfaces\fftw2xf</td>
<td>FFTW 2.x interfaces to the Intel® MKL FFT (Fortran interface)</td>
</tr>
<tr>
<td>interfaces\fftw3xc</td>
<td>FFTW 3.x interfaces to the Intel® MKL FFT (C interface)</td>
</tr>
<tr>
<td>interfaces\fftw3xf</td>
<td>FFTW 3.x interfaces to the Intel® MKL FFT (Fortran interface)</td>
</tr>
<tr>
<td>interfaces\lapack95</td>
<td>Fortran 95 interfaces to LAPACK and a makefile to build the library</td>
</tr>
<tr>
<td>tools</td>
<td>Command-line link tool and tools for creating custom dynamically linkable libraries</td>
</tr>
<tr>
<td>tools\builder</td>
<td>Tools for creating custom dynamically linkable libraries</td>
</tr>
</tbody>
</table>

**Subdirectories of** `<parent directory>`

<table>
<thead>
<tr>
<th>Directory</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>redist\ia32\win\mkl</td>
<td>DLLs for applications running on processors with the IA-32 architecture</td>
</tr>
<tr>
<td>redist\intel64\win\mkl</td>
<td>DLLs for applications running on processors with Intel® 64 architecture</td>
</tr>
</tbody>
</table>

**See Also**

Notational Conventions
Using Code Examples
Layered Model Concept

Intel® MKL is structured to support multiple compilers and interfaces, both serial and multi-threaded modes, different implementations of threading run-time libraries, and a wide range of processors. Conceptually Intel® MKL can be divided into distinct parts to support different interfaces, threading models, and core computations:

1. Interface Layer
2. Threading Layer
3. Computational Layer

You can combine Intel® MKL libraries to meet your needs by linking with one library in each part layer-by-layer.

To support threading with different compilers, you also need to use an appropriate threading run-time library (RTL). These libraries are provided by compilers and are not included in Intel® MKL.

The following table provides more details of each layer.

<table>
<thead>
<tr>
<th>Layer</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interface Layer</td>
<td>This layer matches compiled code of your application with the threading and/or computational parts of the library. This layer provides:</td>
</tr>
<tr>
<td></td>
<td>• cdecl and CVF default interfaces.</td>
</tr>
<tr>
<td></td>
<td>• LP64 and ILP64 interfaces.</td>
</tr>
<tr>
<td></td>
<td>• Compatibility with compilers that return function values differently.</td>
</tr>
<tr>
<td>Threading Layer</td>
<td>This layer:</td>
</tr>
<tr>
<td></td>
<td>• Provides a way to link threaded Intel® MKL with supported compilers.</td>
</tr>
<tr>
<td></td>
<td>• Enables you to link with a threaded or sequential mode of the library.</td>
</tr>
<tr>
<td></td>
<td>This layer is compiled for different environments (threaded or sequential) and compilers (from Intel and PGI*).</td>
</tr>
<tr>
<td>Computational Layer</td>
<td>This layer accommodates multiple architectures through identification of architecture features and chooses the appropriate binary code at run time.</td>
</tr>
</tbody>
</table>

See Also
Using the ILP64 Interface vs. LP64 Interface
Linking Your Application with the Intel® Math Kernel Library
Linking with Threading Libraries
Linking Your Application with the Intel® Math Kernel Library

**Optimization Notice**

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

**Linking Quick Start**

Intel® Math Kernel Library (Intel® MKL) provides several options for quick linking of your application. The simplest options depend on your development environment:

- **Intel® Parallel Studio XE Composer Edition compiler**
  - see Using the /Qmkl Compiler Option.

- **Microsoft Visual Studio® Integrated Development Environment (IDE)**
  - see Automatically Linking a Project in the Visual Studio® Integrated Development Environment with Intel® MKL.

Other options are independent of your development environment, but depend on the way you link:

- **Explicit dynamic linking**
  - see Using the Single Dynamic Library for how to simplify your link line.

- **Explicitly listing libraries on your link line**
  - see Selecting Libraries to Link with for a summary of the libraries.

- **Using pkg-config tool to get compilation and link lines**
  - see Using pkg-config metadata files for a summary on how to use Intel® MKL pkg-config metadata files.

- **Using an interactive interface**
  - see Using the Link-line Advisor to determine libraries and options to specify on your link or compilation line.

- **Using an internally provided tool**
  - see Using the Command-line Link Tool to determine libraries, options, and environment variables or even compile and build your application.

**Using the /Qmkl Compiler Option**

The Intel® Parallel Studio XE Composer Edition compiler supports the following variants of the /Qmkl compiler option:

- `/Qmkl`
- `/Qmkl:parallel`

To link with a certain Intel® MKL threading layer depending on the threading option provided:
For -qopenmp the OpenMP threading layer for Intel compilers
For -tbb the Intel® Threading Building Blocks (Intel® TBB) threading layer

/Qmkl:sequential
to link with sequential version of Intel® MKL.
/Qmkl:cluster
to link with Intel® MKL cluster components (sequential) that use Intel MPI.

NOTE
The -qopenmp option has higher priority than -tbb in choosing the Intel® MKL threading layer for linking.

For more information on the /Qmkl compiler option, see the Intel Compiler User and Reference Guides.
For each variant of the /Qmkl option, the compiler links your application using the following conventions:
• cdecl for the IA-32 architecture
• LP64 for the Intel® 64 architecture

If you specify any variant of the /Qmkl compiler option, the compiler automatically includes the Intel® MKL libraries. In cases not covered by the option, use the Link-line Advisor or see Linking in Detail.

See Also
Using the ILP64 Interface vs. LP64 Interface
Using the Link-line Advisor
Intel® Software Documentation Library for Intel® compiler documentation

Automatically Linking a Project in the Visual Studio® Integrated Development Environment with Intel® MKL

After a default installation of the Intel® Math Kernel Library (Intel® MKL) or Intel® Parallel Studio XE Composer Edition, you can easily configure your project to automatically link with Intel® MKL.

Automatically Linking Your Microsoft Visual C/C++® Project with Intel® MKL

Configure your Microsoft Visual C/C++® project for automatic linking with Intel® MKL as follows:
2. Change the Use MKL property setting by selecting Parallel, Sequential, or Cluster as appropriate.

Specific Intel® MKL libraries that link with your application may depend on more project settings. For details, see the documentation for Intel® Parallel Studio XE Composer Edition for C++.

See Also
Intel® Software Documentation Library for the documentation for Intel® Parallel Studio XE Composer Edition

Automatically Linking Your Intel® Visual Fortran Project with Intel® MKL

Configure your Intel® Visual Fortran project for automatic linking with Intel® MKL as follows:
Go to Project > Properties > Libraries > Use Intel Math Kernel Library and select Parallel, Sequential, or Cluster as appropriate.

Specific Intel® MKL libraries that link with your application may depend on more project settings. For details see the documentation for Intel® Parallel Studio XE Composer Edition for Fortran.
Using the Single Dynamic Library

You can simplify your link line through the use of the Intel® MKL Single Dynamic Library (SDL).

To use SDL, place mkl_rt.lib on your link line. For example:

icl.exe application.c mkl_rt.lib

mkl_rt.lib is the import library for mkl_rt.dll.

SDL enables you to select the interface and threading library for Intel® MKL at run time. By default, linking with SDL provides:

- Intel LP64 interface on systems based on the Intel® 64 architecture
- Intel threading

To use other interfaces or change threading preferences, including use of the sequential version of Intel® MKL, you need to specify your choices using functions or environment variables as explained in section Dynamically Selecting the Interface and Threading Layer.

Selecting Libraries to Link with

To link with Intel® MKL:

- Choose one library from the Interface layer and one library from the Threading layer
- Add the only library from the Computational layer and run-time libraries (RTL)

The following table lists Intel® MKL libraries to link with your application.

<table>
<thead>
<tr>
<th>Interface layer</th>
<th>Threading layer</th>
<th>Computational layer</th>
<th>RTL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intel® 64 architecture, static linking</strong></td>
<td>mkl_intel_l64.lib</td>
<td>mkl_intel_thread.l64.lib</td>
<td>mkl_core.lib</td>
</tr>
<tr>
<td><strong>Intel® 64 architecture, dynamic linking</strong></td>
<td>mkl_intel_l64_dll.lib</td>
<td>mkl_intel_thread_dll.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
<tr>
<td><strong>Intel® Many Integrated Core Architecture (Intel® MIC Architecture), static linking</strong></td>
<td>libmkl_intel_l64.a</td>
<td>libmkl_intel_thread.a</td>
<td>libmkl_core.a</td>
</tr>
<tr>
<td><strong>Intel MIC Architecture, dynamic linking</strong></td>
<td>libmkl_intel_l64.so</td>
<td>libmkl_intel_thread.so</td>
<td>libmkl_core.so</td>
</tr>
</tbody>
</table>

The Single Dynamic Library (SDL) automatically links interface, threading, and computational libraries and thus simplifies linking. The following table lists Intel® MKL libraries for dynamic linking using SDL.

See Dynamically Selecting the Interface and Threading Layer for how to set the interface and threading layers at run time through function calls or environment settings.
Linking with libiomp5md.lib is not required.

For exceptions and alternatives to the libraries listed above, see Linking in Detail.

See Also
Layered Model Concept
Using the Link-line Advisor
Using the /Qmkl Compiler Option
Working with the Cluster Software

Using the Link-line Advisor

Use the Intel® MKL Link-line Advisor to determine the libraries and options to specify on your link or compilation line.

The latest version of the tool is available at https://software.intel.com/en-us/articles/intel-mkl-link-line-advisor. The tool is also available in the documentation directory of the product.

The Advisor requests information about your system and on how you intend to use Intel® MKL (link dynamically or statically, use threaded or sequential mode, and so on). The tool automatically generates the appropriate link line for your application.

See Also
High-level Directory Structure

Using the Command-line Link Tool

Use the command-line Link tool provided by Intel® MKL to simplify building your application with Intel® MKL.

The tool not only provides the options, libraries, and environment variables to use, but also performs compilation and building of your application.

The tool mkl_link_tool.exe is installed in the <mkl_directory>\bin directory, and supports the modes described in the following table.

Intel MKL Command-line Link Tool Modes

<table>
<thead>
<tr>
<th>Mode</th>
<th>Description</th>
<th>Usage</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inquiry</td>
<td>The tool returns the compiler options, libraries, or environment variables</td>
<td>Get Intel® MKL libraries</td>
<td>mkl_link_tool -libs [Intel MKL Link Tool options]</td>
</tr>
<tr>
<td></td>
<td>necessary to build and execute the application.</td>
<td>Get compilation options</td>
<td>mkl_link_tool -opts [Intel MKL Link Tool options]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Get environment variables for application executable</td>
<td>mkl_link_tool -env [Intel MKL Link Tool options]</td>
</tr>
<tr>
<td>Compilation</td>
<td>The Intel® MKL Link Tool builds the application.</td>
<td>—</td>
<td>mkl_link_tool [options] &lt;compiler&gt; [options2] file1 [file2 ...]]</td>
</tr>
<tr>
<td>Mode</td>
<td>Description</td>
<td>Usage</td>
<td>Example</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>Interactive</td>
<td>Allows you to go through all possible Intel® MKL Link Tool supported options. The output provides libraries, options, or environment variables as in the inquiry mode, or a built application as in the compilation mode (depending on what you specify).</td>
<td></td>
<td>mkl_link_tool - interactive</td>
</tr>
</tbody>
</table>

where:

- **options** represents any number of Link Tool options
- **compiler** represents the compiler name: ifort, icc (icpc, icl), cl, gcc (g++), gfortran, pgcc (pgcp), pgf77 (pgf90, pgf95, pgfortran), mpiic, mpiifort, mpic (mpic++), mpif77 (mpif90, mpif95)
- **options2** represents any number of compiler options

Use the **-help** option for full help with the Intel® MKL Link Tool and to show the defaults for the current system.

## Linking Examples

### See Also

* Using the Link-line Advisor
* Examples for Linking with ScaLAPACK and Cluster FFT

### Linking on IA-32 Architecture Systems

The following examples illustrate linking that uses Intel(R) compilers. Most examples use the `.f` Fortran source file. C/C++ users should instead specify a `.cpp` (C++) or `.c` (C) file and replace **ifort** with **icl**:

- **Static linking of myprog.f and OpenMP* threaded Intel® MKL supporting the cdecl interface:**
  ```
  ifort myprog.f mkl_intel_c.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib
  ```
- **Dynamic linking of myprog.f and OpenMP* threaded Intel® MKL supporting the cdecl interface:**
  ```
  ifort myprog.f mkl_intel_c_dll.lib mkl_intel_thread_dll.lib mkl_core_dll.lib
  ```
- **Static linking of myprog.f and sequential version of Intel® MKL supporting the cdecl interface:**
  ```
  ifort myprog.f mkl_intel_c.lib mkl_sequential.lib mkl_core.lib
  ```
Dynamic linking of myprog.f and sequential version of Intel® MKL supporting the cdecl interface:

```fortran
ifort myprog.f mkl_intel_c_dll.lib mkl_sequential_dll.lib mkl_core_dll.lib
```

Static linking of user code myprog.f and OpenMP* threadedIntel® MKL supporting the stdcall interface:

```fortran
ifort myprog.f mkl_intel_s.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib
```

Dynamic linking of user code myprog.f and OpenMP* threadedIntel® MKL supporting the stdcall interface:

```fortran
ifort myprog.f mkl_intel_s_dll.lib mkl_intel_thread_dll.lib mkl_core_dll.lib libiomp5md.lib
```

Dynamic linking of myprog.f and OpenMP* threaded or sequential Intel® MKL supporting the cdecl or stdcall interface (Call the mkl_set_threading_layer function or set value of the MKL_THREADING_LAYER environment variable to choose threaded or sequential mode):

```fortran
ifort myprog.f mkl_rt.lib
```

Static linking of myprog.f, Fortran 95 LAPACK interface, and OpenMP* threaded Intel® MKL supporting the cdecl interface:

```fortran
ifort myprog.f mkl_lapack95.lib mkl_intel_c.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib
```

Static linking of myprog.f, Fortran 95 BLAS interface, and OpenMP* threaded Intel® MKL supporting the cdecl interface:

```fortran
ifort myprog.f mkl_blas95.lib mkl_intel_c.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib
```

Static linking of myprog.c and Intel® MKL threaded with Intel® Threading Building Blocks (Intel® TBB), provided that the LIB environment variable contains the path to Intel TBB library:

```fortran
icl myprog.c /link /libpath:$MKLPATH -I$MKLINCLUDE mkl_intel.lib mkl_tbb_thread.lib mkl_core.lib tbb.lib
```

Dynamic linking of myprog.c and Intel® MKL threaded with Intel TBB, provided that the LIB environment variable contains the path to Intel TBB library:

```fortran
icl myprog.c /link /libpath:$MKLPATH -I$MKLINCLUDE mkl_intel_dll.lib mkl_tbb_thread_dll.lib mkl_core_dll.lib tbb.lib
```

See Also
Fortran 95 Interfaces to LAPACK and BLAS
Examples for linking a C application using cluster components
Examples for linking a Fortran application using cluster components
Using the Single Dynamic Library

**Linking on Intel(R) 64 Architecture Systems**

The following examples illustrate linking that uses Intel(R) compilers.

Most examples use the .f Fortran source file. C/C++ users should instead specify a .cpp (C++) or .c (C) file and replace ifort with icl:

- Static linking of myprog.f and OpenMP* threadedIntel® MKL supporting the LP64 interface:

```fortran
ifort myprog.f mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib
```

- Dynamic linking of myprog.f and OpenMP* threadedIntel® MKL supporting the LP64 interface:

```fortran
ifort myprog.f mkl_intel_lp64_dll.lib mkl_intel_thread_dll.lib mkl_core_dll.lib libiomp5md.lib
```

- Static linking of myprog.f and sequential version of Intel® MKL supporting the LP64 interface:

```fortran
ifort myprog.f mkl_intel_lp64.lib mkl_sequential.lib mkl_core.lib
```

- Dynamic linking of myprog.f and sequential version of Intel® MKL supporting the LP64 interface:

```fortran
ifort myprog.f mkl_intel_lp64_dll.lib mkl_sequential_dll.lib mkl_core_dll.lib
```
• Static linking of myprog.f and OpenMP* threaded Intel® MKL supporting the ILP64 interface:
  ifort myprog.f mkl_intel_ilp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib

• Dynamic linking of myprog.f and OpenMP* threaded Intel® MKL supporting the ILP64 interface:
  ifort myprog.f mkl_intel_ilp64_dll.lib mkl_intel_thread_dll.lib mkl_core_dll.lib
  libiomp5md.lib

• Dynamic linking of user code myprog.f and OpenMP* threaded or sequential Intel® MKL supporting the LP64 or ILP64 interface (Call appropriate functions or set environment variables to choose threaded or sequential mode and to set the interface):
  ifort myprog.f mkl_rt.lib

• Static linking of myprog.f, Fortran 95 LAPACK interface, and OpenMP* threaded Intel® MKL supporting the LP64 interface:
  ifort myprog.f mkl_lapack95_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib
  mkl_core.lib libiomp5md.lib

• Static linking of myprog.f, Fortran 95 BLAS interface, and OpenMP* threaded Intel® MKL supporting the LP64 interface:
  ifort myprog.f mkl_blas95_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib
  mkl_core.lib libiomp5md.lib

• Static linking of myprog.c and Intel® MKL threaded with Intel® Threading Building Blocks (Intel® TBB), provided that the LIB environment variable contains the path to Intel TBB library:
  icl myprog.c /link /libpath:%MKLPATH% -I%MKLINCLUDE% mkl_intel_lp64.lib
  mkl_tbb_thread.lib mkl_core.lib tbb.lib

• Dynamic linking of myprog.c and Intel® MKL threaded with Intel TBB, provided that the LIB environment variable contains the path to Intel TBB library:
  icl myprog.c /link /libpath:%MKLPATH% -I%MKLINCLUDE% mkl_intel_lp64_dll.lib
  mkl_tbb_thread_dll.lib mkl_core_dll.lib tbb.lib

See Also
Fortran 95 Interfaces to LAPACK and BLAS
Examples for linking a C application using cluster components
Examples for linking a Fortran application using cluster components
Using the Single Dynamic Library

Linking in Detail
This section recommends which libraries to link with depending on your Intel® MKL usage scenario and provides details of the linking.

Dynamically Selecting the Interface and Threading Layer
The Single Dynamic Library (SDL) enables you to dynamically select the interface and threading layer for Intel® MKL.

Setting the Interface Layer
To set the interface layer at run time, use the mkl_set_interface_layer function or the MKL_INTERFACE_LAYER environment variable.

Available interface layers depend on the architecture of your system.

On systems based on the Intel® 64 architecture, LP64 and ILP64 interfaces are available. The following table provides values to be used to set each interface layer.
Specifying the Interface Layer

<table>
<thead>
<tr>
<th>Interface Layer</th>
<th>Value of MKL_INTERFACE_LAYER</th>
<th>Value of the Parameter of mkl_set_interface_layer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel LP64, default</td>
<td>LP64</td>
<td>MKL_INTERFACE_LP64</td>
</tr>
<tr>
<td>Intel ILP64</td>
<td>ILP64</td>
<td>MKL_INTERFACE_ILP64</td>
</tr>
</tbody>
</table>

If the mkl_set_interface_layer function is called, the environment variable MKL_INTERFACE_LAYER is ignored.

See the Intel® MKL Developer Reference for details of the mkl_set_interface_layer function.

On systems based on the IA-32 architecture, the cdecl and stdcall interfaces are available. These interfaces have different function naming conventions, and SDL selects between cdecl and stdcall at link time according to the function names.

Setting the Threading Layer

To set the threading layer at run time, use the mkl_set_threading_layer function or the MKL_THREADING_LAYER environment variable. The following table lists available threading layers along with the values to be used to set each layer.

Specifying the Threading Layer

<table>
<thead>
<tr>
<th>Threading Layer</th>
<th>Value of MKL_THREADING_LAYER</th>
<th>Value of the Parameter of mkl_set_threading_layer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel threading, default</td>
<td>INTEL</td>
<td>MKL_THREADING_INTEL</td>
</tr>
<tr>
<td>Sequential mode of Intel® MKL</td>
<td>SEQUENTIAL</td>
<td>MKL_THREADING_SEQUENTIAL</td>
</tr>
<tr>
<td>PGI threading†</td>
<td>PGI</td>
<td>MKL_THREADING_PGI</td>
</tr>
<tr>
<td>Intel TBB threading</td>
<td>TBB</td>
<td>MKL_THREADING_TBB</td>
</tr>
</tbody>
</table>

† Not supported by the SDL for Intel® Many Integrated Core Architecture.

If the mkl_set_threading_layer function is called, the environment variable MKL_THREADING_LAYER is ignored.

See the Intel® MKL Developer Reference for details of the mkl_set_threading_layer function.

Replacing Error Handling and Progress Information Routines

You can replace the Intel® MKL error handling routine xerbla or progress information routine mkl_progress with your own function. If you are using SDL, to replace xerbla or mkl_progress, call the mkl_set_xerbla and mkl_set_progress function, respectively. See the Intel® MKL Developer Reference for details.

NOTE
If you are using SDL, you cannot perform the replacement by linking the object file with your implementation of xerbla or mkl_progress.

See Also
Using the Single Dynamic Library
Layered Model Concept
Using the cdecl and stdcall Interfaces
Directory Structure in Detail

**Linking with Interface Libraries**

**Using the cdecl and stdcall Interfaces**

cdecl and stdcall calling conventions differ in the way how the stack is restored after a function call. Intel® MKL supports both conventions in its IA-32 architecture implementation through the `mkl_intel_c[.dll].lib` and `mkl_intel_s[.dll].lib` interface libraries. These libraries assume the defaults of different compilers, which also differ in the position of the string lengths in the lists of parameters passed to the calling program, as explained in the following table:

<table>
<thead>
<tr>
<th>Library for Static Linking</th>
<th>Library for Dynamic Linking</th>
<th>Calling Convention</th>
<th>Position of String Lengths in Parameter Lists</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_intel_c.lib</td>
<td>mkl_intel_c_dll.lib</td>
<td>cdecl</td>
<td>At the end</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>The defaults of Intel® C++ and Intel® Fortran compilers</td>
</tr>
<tr>
<td>mkl_intel_s.lib</td>
<td>mkl_intel_s_dll.lib</td>
<td>stdcall</td>
<td>Immediately after the string address</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>The defaults of Compaq Visual Fortran* (CVF) compiler</td>
</tr>
</tbody>
</table>

**Important**

To avoid errors, ensure that the calling and called programs use the same calling convention.

To use the cdecl or stdcall calling convention, use appropriate calling syntax in C applications and appropriate compiler options for Fortran applications.

If you are using a C compiler, to link with the cdecl or stdcall interface library, call Intel® MKL routines in your code as explained in the table below:

<table>
<thead>
<tr>
<th>Interface Library</th>
<th>Calling Intel® MKL Routines</th>
</tr>
</thead>
</table>
| mkl_intel_c[_dll].lib | Use the following declaration:  
  ```
  <type> name( <prototype variable1>, <prototype variable2>, .. );
  ``` |
| mkl_intel_s[_dll].lib | Call a routine with the following statement:  
  ```
  extern __stdcall name( <prototype variable1>, <prototype variable2>, .. );
  ``` |

If you are using a Fortran compiler, to link with the cdecl or stdcall interface library, provide compiler options as explained in the table below:

<table>
<thead>
<tr>
<th>Interface Library</th>
<th>Compiler Options</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_intel_c[_dll].lib</td>
<td>/Gm</td>
<td>/Gm and /iface:cvf options enable compatibility of the CVF and Powerstation calling conventions</td>
</tr>
<tr>
<td>mkl_intel_s[_dll].lib</td>
<td>/Gm or /iface:cvf</td>
<td></td>
</tr>
</tbody>
</table>
### Interface Library

<table>
<thead>
<tr>
<th>Interface Library</th>
<th>Compiler Options</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_intel_s[_dll].lib</td>
<td>Default</td>
<td>CVF compiler</td>
</tr>
<tr>
<td>mkl_intel_c[_dll].lib</td>
<td>/iface=(cref,</td>
<td></td>
</tr>
<tr>
<td></td>
<td>nomixed_str_len_arg)</td>
<td></td>
</tr>
</tbody>
</table>

#### See Also

- Using the stdcall Calling Convention in C/C++
- Compiling an Application that Calls Intel® MKL and Uses the CVF Calling Conventions

### Using the ILP64 Interface vs. LP64 Interface

The Intel® MKL ILP64 libraries use the 64-bit integer type (necessary for indexing large arrays, with more than \(2^{31}-1\) elements), whereas the LP64 libraries index arrays with the 32-bit integer type.

The LP64 and ILP64 interfaces are implemented in the Interface layer. Link with the following interface libraries for the LP64 or ILP64 interface, respectively:

- `mkl_intel_lp64.lib` or `mkl_intel_ilp64.lib` for static linking
- `mkl_intel_lp64_dll.lib` or `mkl_intel_ilp64_dll.lib` for dynamic linking

The ILP64 interface provides for the following:

- Support large data arrays (with more than \(2^{31}-1\) elements)
- Enable compiling your Fortran code with the `/4I8` compiler option

The LP64 interface provides compatibility with the previous Intel® MKL versions because “LP64” is just a new name for the only interface that the Intel® MKL versions lower than 9.1 provided. Choose the ILP64 interface if your application uses Intel® MKL for calculations with large data arrays or the library may be used so in future.

Intel® MKL provides the same include directory for the ILP64 and LP64 interfaces.

### Compiling for LP64/ILP64

The table below shows how to compile for the ILP64 and LP64 interfaces:

#### Fortran

| Compiling for ILP64 | ifort /4I8 /I<mkl directory>
include ... |
|---------------------|--------------------------------------------------|
| Compiling for LP64  | ifort /I<mkl directory>
include ... |

#### C or C++

| Compiling for ILP64 | icl /DMKL_ILP64 /I<mkl directory>
include ... |
|---------------------|--------------------------------------------------|
| Compiling for LP64  | icl /I<mkl directory>
include ... |

#### Caution

Linking of an application compiled with the `/4I8` or `/DMKL_ILP64` option to the LP64 libraries may result in unpredictable consequences and erroneous output.
Coding for ILP64

You do not need to change existing code if you are not using the ILP64 interface.

To migrate to ILP64 or write new code for ILP64, use appropriate types for parameters of the Intel® MKL functions and subroutines:

<table>
<thead>
<tr>
<th>Integer Types</th>
<th>Fortran</th>
<th>C or C++</th>
</tr>
</thead>
<tbody>
<tr>
<td>32-bit integers</td>
<td>INTEGER*4 or INTEGER(KIND=4)</td>
<td>int</td>
</tr>
<tr>
<td>Universal integers for ILP64/</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LP64:</td>
<td>INTEGER (without specifying KIND)</td>
<td>MKL_INT</td>
</tr>
<tr>
<td>• 64-bit for ILP64</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• 32-bit otherwise</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Universal integers for ILP64/</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LP64:</td>
<td>INTEGER*8 or INTEGER(KIND=8)</td>
<td>MKL_INT64</td>
</tr>
<tr>
<td>• 64-bit integers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FFT interface integers for ILP64/</td>
<td></td>
<td>MKL_LONG</td>
</tr>
<tr>
<td>LP64</td>
<td>INTEGER (without specifying KIND)</td>
<td></td>
</tr>
</tbody>
</table>

To determine the type of an integer parameter of a function, use appropriate include files. For functions that support only a Fortran interface, use the C/C++ include files *.h.

The above table explains which integer parameters of functions become 64-bit and which remain 32-bit for ILP64. The table applies to most Intel® MKL functions except some Vector Mathematics and Vector Statistics functions, which require integer parameters to be 64-bit or 32-bit regardless of the interface:

• **Vector Mathematics**: The `mode` parameter of the functions is 64-bit.
• **Random Number Generators (RNG)**:
  - All discrete RNG except `viRngUniformBits64` are 32-bit.
  - The `viRngUniformBits64` generator function and `vslSkipAheadStream` service function are 64-bit.
• **Summary Statistics**: The `estimate` parameter of the `vslsSSCompute/vsldSSCompute` function is 64-bit.

Refer to the *Intel® MKL Developer Reference* for more information.

To better understand ILP64 interface details, see also examples.

Limitations

All Intel® MKL function domains support ILP64 programming but FFTW interfaces to Intel® MKL:

• FFTW 2.x wrappers do not support ILP64.
• FFTW 3.x wrappers support ILP64 by a dedicated set of functions `plan_guru64`.

See Also

- High-level Directory Structure
- Intel® MKL Include Files
- Language Interfaces Support, by Function Domain
- Layered Model Concept
- Directory Structure in Detail
Linking with Fortran 95 Interface Libraries

The mkl_blas95*.lib and mkl_lapack95*.lib libraries contain Fortran 95 interfaces for BLAS and LAPACK, respectively, which are compiler-dependent. In the Intel® MKL package, they are prebuilt for the Intel® Fortran compiler. If you are using a different compiler, build these libraries before using the interface.

See Also
Fortran 95 Interfaces to LAPACK and BLAS
Compiler-dependent Functions and Fortran 90 Modules

Linking with Threading Libraries

Intel® MKL threading layer defines how Intel® MKL functions utilize multiple computing cores of the system that the application runs on. You must link your application with one appropriate Intel® MKL library in this layer, as explained below. Depending on whether this is a threading or a sequential library, Intel® MKL runs in a parallel or sequential mode, respectively.

In the parallel mode, Intel® MKL utilizes multiple processor cores available on your system, uses the OpenMP* or Intel TBB threading technology, and requires a proper threading run-time library (RTL) to be linked with your application. Independently of use of Intel® MKL, the application may also require a threading RTL. You should link not more than one threading RTL to your application. Threading RTLs are provided by your compiler. Intel® MKL provides several threading libraries, each dependent on the threading RTL of a certain compiler, and your choice of the Intel® MKL threading library must be consistent with the threading RTL that you use in your application.

The OpenMP RTL of the Intel® compiler is the libiomp5md.lib library, located under <parent directory>\compiler\lib. You can find additional information about the Intel OpenMP RTL at https://www.openmp.org.

The Intel TBB RTL of the Intel® compiler is the tbb.lib library, located under <parent directory>\tbb\lib. You can find additional information about the Intel TBB RTL at https://www.threadingbuildingblocks.org.

In the sequential mode, Intel® MKL runs unthreaded code, does not require a threading RTL, and does not respond to environment variables and functions controlling the number of threads. Avoid using the library in the sequential mode unless you have a particular reason for that, such as the following:

- Your application needs a threading RTL that none of Intel® MKL threading libraries is compatible with
- Your application is already threaded at a top level, and using parallel Intel® MKL only degrades the application performance by interfering with that threading
- Your application is intended to be run on a single thread, like a message-passing Interface (MPI) application

It is critical to link the application with the proper RTL. The table below explains what library in the Intel® MKL threading layer and what threading RTL you should choose under different scenarios:

<table>
<thead>
<tr>
<th>Application</th>
<th>Intel® MKL</th>
<th>RTL Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uses OpenMP</td>
<td>Compiled with</td>
<td>Execution Mode</td>
</tr>
<tr>
<td>no</td>
<td>any compiler</td>
<td>parallel</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>no</td>
<td>any compiler</td>
<td>parallel</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Linking Your Application with the Intel® Math Kernel Library

<table>
<thead>
<tr>
<th>Application</th>
<th>Intel® MKL</th>
<th>RTL Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uses OpenMP</td>
<td>Compiled with</td>
<td>Execution Mode</td>
</tr>
<tr>
<td>no</td>
<td>any compiler</td>
<td>sequential</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>yes</td>
<td>Intel compiler</td>
<td>parallel</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>yes</td>
<td>PGI* compiler</td>
<td>parallel</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>yes</td>
<td>any other compiler</td>
<td>parallel</td>
</tr>
</tbody>
</table>

See Also
Layered Model Concept
Notational Conventions

Linking with Computational Libraries
If you are not using the Intel® MKL ScaLAPACK and Cluster Fast Fourier Transforms (FFT), you need to link your application with only one computational library, depending on the linking method:

<table>
<thead>
<tr>
<th>Static Linking</th>
<th>Dynamic Linking</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_core.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
</tbody>
</table>

Computational Libraries for Applications that Use ScaLAPACK or Cluster FFT
ScaLAPACK and Cluster FFT require more computational libraries, which may depend on your architecture.

The following table lists computational libraries for IA-32 architecture applications that use ScaLAPACK or Cluster FFT.
**Computational Libraries for IA-32 Architecture**

<table>
<thead>
<tr>
<th>Function domain</th>
<th>Static Linking</th>
<th>Dynamic Linking</th>
</tr>
</thead>
<tbody>
<tr>
<td>ScaLAPACK †</td>
<td>mkl_scalapack_core.lib</td>
<td>mkl_scalapack_core_dll.lib</td>
</tr>
<tr>
<td></td>
<td>mkl_core.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
<tr>
<td>Cluster Fourier Transform Functions ‡</td>
<td>mkl_cdft_core.lib</td>
<td>mkl_cdft_core_dll.lib</td>
</tr>
<tr>
<td></td>
<td>mkl_core.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
</tbody>
</table>

† Also add the library with BLACS routines corresponding to the MPI used.

The following table lists computational libraries for Intel® 64 or Intel® Many Integrated Core Architecture applications that use ScaLAPACK or Cluster FFT.

**Computational Libraries for the Intel® 64 or Intel® Many Integrated Core Architecture**

<table>
<thead>
<tr>
<th>Function domain</th>
<th>Static Linking</th>
<th>Dynamic Linking</th>
</tr>
</thead>
<tbody>
<tr>
<td>ScaLAPACK, LP64 interface ‡</td>
<td>mkl_scalapack_lp64.lib</td>
<td>mkl_scalapack_lp64_dll.lib</td>
</tr>
<tr>
<td></td>
<td>mkl_core.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
<tr>
<td>ScaLAPACK, ILP64 interface ‡</td>
<td>mkl_scalapack_ilp64.lib</td>
<td>mkl_scalapack_ilp64_dll.lib</td>
</tr>
<tr>
<td></td>
<td>mkl_core.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
<tr>
<td>Cluster Fourier Transform Functions ‡</td>
<td>mkl_cdft_core.lib</td>
<td>mkl_cdft_core_dll.lib</td>
</tr>
<tr>
<td></td>
<td>mkl_core.lib</td>
<td>mkl_core_dll.lib</td>
</tr>
</tbody>
</table>

‡ Also add the library with BLACS routines corresponding to the MPI used.

**See Also**
- Linking with ScaLAPACK and Cluster FFT
- Using the Link-line Advisor
- Using the ILP64 Interface vs. LP64 Interface

**Linking with Compiler Run-time Libraries**

Dynamically link libiomp5 or tbb library even if you link other libraries statically.

Linking to the libiomp5 statically can be problematic because the more complex your operating environment or application, the more likely redundant copies of the library are included. This may result in performance issues (oversubscription of threads) and even incorrect results.

To link libiomp5 or tbb dynamically, be sure the PATH environment variable is defined correctly.

Sometimes you may improve performance of your application with threaded Intel® MKL by using the /MT compiler option. The compiler driver will pass the option to the linker and the latter will load multi-thread (MT) static run-time libraries.

However, to link a Vector Mathematics (VM) application that uses the errno variable for error reporting, compile and link your code using the option that depends on the linking model:
- /MT for linking with static Intel® MKL libraries
- /MD for linking with dynamic Intel® MKL libraries

**See Also**
- Setting Environment Variables
- Layered Model Concept
Linking with System Libraries

If your system is based on the Intel® 64architecture, be aware that Microsoft SDK builds 1289 or higher provide the `bufferoverflowu.lib` library to resolve the `__security_cookie` external references. Makefiles for examples and tests include this library by using the `buf_lib=bufferoverflowu.lib` macro. If you are using older SDKs, leave this macro empty on your command line as follows: `buf_lib= `. 

See Also
Linking Examples

Building Custom Dynamic-link Libraries

Custom dynamic-link libraries (DLL) reduce the collection of functions available in Intel® MKL libraries to those required to solve your particular problems, which helps to save disk space and build your own dynamic libraries for distribution.

The Intel® MKL custom DLL builder enables you to create a dynamic library containing the selected functions and located in the `tools\builder` directory. The builder contains a makefile and a definition file with the list of functions.

Using the Custom Dynamic-link Library Builder in the Command-line Mode

To build a custom DLL, use the following command:

```
nmake target [<options>]
```

The following table lists possible values of `target` and explains what the command does for each value:

<table>
<thead>
<tr>
<th>Value</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>libia32</td>
<td>The builder uses static Intel® MKL interface, threading, and core libraries to build a customDLL for the IA-32 architecture.</td>
</tr>
<tr>
<td>libintel64</td>
<td>The builder uses static Intel® MKL interface, threading, and core libraries to build a customDLL for the Intel® 64 architecture.</td>
</tr>
<tr>
<td>dllia32</td>
<td>The builder uses the single dynamic library <code>libmkl_rt.dll</code> to build a custom DLL for the IA-32 architecture.</td>
</tr>
<tr>
<td>dllintel64</td>
<td>The builder uses the single dynamic library <code>libmkl_rt.dll</code> to build a custom DLL for the Intel® 64 architecture.</td>
</tr>
<tr>
<td>help</td>
<td>The command prints Help on the custom DLL builder</td>
</tr>
</tbody>
</table>

The `<options>` placeholder stands for the list of parameters that define macros to be used by the makefile. The following table describes these parameters:

<table>
<thead>
<tr>
<th>Parameter [Values]</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>interface</td>
<td>Defines which programming interface to use. Possible values:</td>
</tr>
<tr>
<td></td>
<td>• For the IA-32 architecture, {cdecl</td>
</tr>
<tr>
<td></td>
<td>• For the Intel 64 architecture, {lp64</td>
</tr>
<tr>
<td>threading =</td>
<td>Defines whether to use the Intel® MKL in the threaded or sequential mode. The default value is sequential.</td>
</tr>
</tbody>
</table>
## Parameter [Values]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>cluster = {yes</td>
<td>no}</td>
</tr>
<tr>
<td>blacs_mpi = {intelmpi</td>
<td>mpich2</td>
</tr>
<tr>
<td>blacs_name = &lt;lib name&gt;</td>
<td>Specifies the name (without extension) of a custom Intel® MKL BLACS library to use. Ignored if 'cluster=no'. If 'blacs_mpi' is specified, 'blacs_name' is ignored. The default value is mkl_blacs_&lt;blacs_mpi&gt;_&lt;interface&gt;.</td>
</tr>
<tr>
<td>mpi = &lt;lib name&gt;</td>
<td>Specifies the name (without extension) of the MPI library used to build the custom DLL. Ignored if 'cluster=no'. The default value is mpi.</td>
</tr>
<tr>
<td>export = &lt;file name&gt;</td>
<td>Specifies the full name of the file that contains the list of entry-point functions to be included in the DLL. The default name is user_example_list (no extension).</td>
</tr>
<tr>
<td>name = &lt;dll name&gt;</td>
<td>Specifies the name of the dll and interface library to be created. By default, the names of the created libraries are mkl_custom.dll and mkl_custom.lib.</td>
</tr>
<tr>
<td>xerbla = &lt;error handler&gt;</td>
<td>Specifies the name of the object file &lt;user_xerbla&gt;.obj that contains the error handler of the user. The makefile adds this error handler to the library for use instead of the default Intel® MKL error handler xerbla. If you omit this parameter, the native Intel® MKL xerbla is used. See the description of the xerbla function in the Intel® MKL Developer Reference to develop your own error handler. For the IA-32 architecture, the object file should be in the interface defined by the interface macro (cdecl or stdcall).</td>
</tr>
<tr>
<td>MKLROOT = &lt;mkl directory&gt;</td>
<td>Specifies the location of Intel® MKL libraries used to build the custom DLL. By default, the builder uses the Intel® MKL installation directory.</td>
</tr>
<tr>
<td>uwd_compat = {yes</td>
<td>no}</td>
</tr>
<tr>
<td>buf_lib</td>
<td>Manages resolution of the __security_cookie external references in the custom DLL on systems based on the Intel® 64 architecture. By default, the makefile uses the bufferoverflowu.lib library of Microsoft SDK builds 1289 or higher. This library resolves the __security_cookie external references. To avoid using this library, set the empty value of this parameter. Therefore, if you are using an older SDK, set buf_lib=.</td>
</tr>
<tr>
<td>Caution</td>
<td>Use the buf_lib parameter only with the empty value. Incorrect value of the parameter causes builder errors.</td>
</tr>
<tr>
<td>crt = &lt;c runtime library&gt;</td>
<td>Specifies the name of the Microsoft C run-time library to be used to build the custom DLL. By default, the builder uses msvcr100.dll.</td>
</tr>
<tr>
<td>manifest = {yes</td>
<td>no</td>
</tr>
<tr>
<td>Parameter [Values]</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| • If *manifest=yes*, the manifest file with the name defined by the *name* parameter above and the *manifest* extension is created.  
• If *manifest=no*, the manifest file is not be created.  
• If *manifest=embed*, the manifest is embedded into the DLL. |

By default, the builder does not use the *manifest* parameter.

All of the above parameters are optional. However, you must make the system and c-runtime (crt) libraries and link.exe available by setting the *PATH* and *LIB* environment variables appropriately. You can do this in the following ways:

- Manually
- If you are using Microsoft Visual Studio (VS), call the `vcvarsall.bat` script with the appropriate 32-bit (x86) or 64-bit (x64 or amd-64) architecture flag.
- If you are using the Intel compiler, use the `compilervars.bat` script with the appropriate 32-bit (x86) or 64-bit (x64 or amd-64) architecture flag.

In the simplest case, the command line is:

```bash
# source Visual Studio environment variables  
call vcvarsall.bat x86  
# run custom dll builder script  
nmake ia32
```

and the missing options have default values. This command creates the `mkl_custom.dll` and `mkl_custom.lib` libraries with the cdecl interface for processors using the IA-32 architecture. The command takes the list of functions from the `functions_list` file and uses the native Intel® MKL error handler `xerbla`.

Here is an example of a more complex case:

```bash
# source Visual Studio environment variables  
call vcvarsall.bat x86  
# run custom dll builder script  
nmake ia32 interface=stdcall export=my_func_list.txt name=mkl_small xerbla=my_xerbla.obj
```

In this case, the command creates the `mkl_small.dll` and `mkl_small.lib` libraries with the stdcall interface for processors using the IA-32 architecture. The command takes the list of functions from the `my_func_list.txt` file and uses the error handler of the user `my_xerbla.obj`.

To build a UWD-compatible custom dll, use the `uwd_compat=yes` option. For this purpose, you must make a different set of universal system (`OneCore.lib`) and universal c-runtime (`ucrt.lib`) libraries available. You can get these libraries by downloading Windows 10 SDK 10.0.17134.0 (version 1803) or newer. Make sure to source the Visual Studio environment with the appropriate native architecture to add the libraries to your path.

This example shows how to create a 64-bit architecture library, `mkl_uwd_compat.dll`, that is UWD-compatible with the `lp64` interface using `my_function_list.txt` for specific functionality:

```bash
# source Visual Studio environment variables, LIB should have paths to desired OneCore.lib and universal crt libraries  
call vcvarsall.bat x64  
# run custom dll builder script  
nmake intel64 interface=lp64 export=my_func_list.txt uwd_compat=yes name=mkl_uwd_compat
```
See Also
Linking with System Libraries

Composing a List of Functions

To compose a list of functions for a minimal custom DLL needed for your application, you can use the following procedure:

1. Link your application with installed Intel® MKL libraries to make sure the application builds.
2. Remove all Intel® MKL libraries from the link line and start linking.
   Unresolved symbols indicate Intel® MKL functions that your application uses.
3. Include these functions in the list.

   **Important**
   Each time your application starts using more Intel® MKL functions, update the list to include the new functions.

See Also
Specifying Function Names

Specifying Function Names

In the file with the list of functions for your custom DLL, adjust function names to the required interface. For example, you can list the cdecl entry points as follows:

DGEMM
DTRSM
DDOT
DGETRF
DGETRS
cblas_dgemm
cblas_ddot

You can list the stdcall entry points as follows:

 DGEMM@60
__DDOT@20
__DGETRF@24

For more examples, see domain-specific lists of function names in the `<mkl directory>\tools\builder` folder. This folder contains lists of function names for both cdecl or stdcall interfaces.

**NOTE**
The lists of function names are provided in the `<mkl directory>\tools\builder` folder merely as examples. See Composing a List of Functions for how to compose lists of functions for your custom DLL.
Tip
Names of Fortran-style routines (BLAS, LAPACK, etc.) can be both upper-case or lower-case, with or without the trailing underscore. For example, these names are equivalent:
BLAS: dgemm, DGEMM, dgemm_, DGEMM
LAPACK: dgetrf, DGETRF, dgetrf_, DGETRF_.

Properly capitalize names of C support functions in the function list. To do this, follow the guidelines below:

1. In the mkl_service.h include file, look up a #define directive for your function (mkl_service.h is included in the mkl.h header file).
2. Take the function name from the replacement part of that directive.

For example, the #define directive for the mkl_disable_fast_mm function is
#define mkl_disable_fast_mm MKL_Disable_Fast_MM.

Capitalize the name of this function in the list like this: MKL_Disable_Fast_MM.
For the names of the Fortran support functions, see the tip.

Building a Custom Dynamic-link Library in the Visual Studio® Development System

You can build a custom dynamic-link library (DLL) in the Microsoft Visual Studio® Development System (VS*). To do this, use projects available in the tools\builder\MSVS_Projects subdirectory of the Intel® MKL directory. The directory contains subdirectories with projects for the respective versions of the Visual Studio Development System, for example, VS2012. For each version of VS two solutions are available:

- libia32.sln builds a custom DLL for the IA-32 architecture.
- libintel64.sln builds a custom DLL for the Intel® 64 architecture.

The builder uses the following default settings for the custom DLL:

- Interface: cdecl for the IA-32 architecture and LP64 for the Intel 64 architecture
- Error handler: Native Intel® MKL xerbla
- Create Microsoft manifest: yes
- List of functions: in the project's source file examples.def

To build a custom DLL:

1. Set the MKLROOT environment variable with the installation directory of the Intel® MKL version you are going to use.
2. Open the libia32.sln or libintel64.sln solution depending on the architecture of your system.
   The solution includes the following projects:
   - i_malloc_dll
   - vml_dll_core
   - cdecl_parallel (in libia32.sln) or lp64_parallel (in libintel64.sln)
   - cdecl_sequential (in libia32.sln) or lp64_sequential (in libintel64.sln)
3. [Optional] To change any of the default settings, select the project depending on whether the DLL will use Intel® MKL functions in the sequential or multi-threaded mode:
   - In the libia32 solution, select the cdecl_sequential or cdecl_parallel project.
   - In the libintel64 solution, select the lp64_sequential or lp64_parallel project.
4. [Optional] To build the DLL that uses the stdcall interface for the IA-32 architecture or the ILP64 interface for the Intel 64 architecture:
   b. In the libia32 solution, change mkl_intel_c.lib to mkl_intel_s.lib.
      In the libintel64 solution, change mkl_intel_lp64.lib to mkl_intel_ilp64.lib.
5. [Optional] To include your own error handler in the DLL:
   a. Select Project>Properties>Configuration Properties>Linker>Input.
   b. Add <user_xerbla>.obj
5. [Optional] To turn off creation of the manifest:
   b. Select: no.
6. [Optional] To change the list of functions to be included in the DLL:
   a. Select Source Files.
   b. Edit the examples.def file. Refer to Specifying Function Names for how to specify entry points.
8. To build the library, select Build>Build Solution.

See Also
Using the Custom Dynamic-link Library Builder in the Command-line Mode

Distributing Your Custom Dynamic-link Library
To enable use of your custom DLL in a threaded mode, distribute libiomp5md.dll along with the custom DLL.

Building a Universal Windows Driver
Intel® MKL is compatible with Universal Windows Drivers (UWDs) whose binaries are Win10 SDK version 1803 (also called Redstone 4, RS4 or Win 10 SDK build 10.0.17134.0) or higher. The Windows system calls used in the statically linked libraries are defined in:

- OneCore.Lib for headless applications like Nano Server.
- OneCoreUAP.Lib for a UWD.

A Windows application or driver that links to either of these libraries can use functionality from statically linked Intel® MKL to accelerate computations. For more information on design criteria for a UWD, see the Microsoft website.

This table summarizes important information about compatibility between Intel® MKL and UWDs.

### Compatibility between Intel MKL and Universal Windows Driver (UWD)

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Compatibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>SDK</td>
<td>10.0.17134.0 (also called Redstone 4, RS4, version 1803) or higher</td>
</tr>
<tr>
<td>Architecture</td>
<td>• ia32</td>
</tr>
<tr>
<td></td>
<td>• intel64</td>
</tr>
<tr>
<td>Universal API sets</td>
<td>• OneCore.Lib for headless applications like Nano Server</td>
</tr>
<tr>
<td></td>
<td>• OneCoreUAP.Lib for a UWD</td>
</tr>
<tr>
<td>Threading layers</td>
<td>• Sequential</td>
</tr>
<tr>
<td></td>
<td>• Intel® Threading Building Blocks (Intel® TBB). See Using Intel® TBB in UWP Applications.</td>
</tr>
<tr>
<td>UWD Compliance</td>
<td>• Intel® MKL with static linking libraries</td>
</tr>
<tr>
<td></td>
<td>• For dynamic linking libraries, use the custom DLL builder with uwd_compat=yes to construct a custom DLL that is compatible with UWD. See Using the Custom DLL Builder.</td>
</tr>
</tbody>
</table>
NOTE The UWD links to Intel® MKL using the same link line settings as drivers or executables that are not universal. See Link Line Advisor for recommended linking models.

### UWD Compliance for 32-bit Architecture (ia32)
These tables describe compliance between Intel® MKL and UWD on 32-bit (ia32) architecture for static and dynamic linked libraries.

#### Static Linking

<table>
<thead>
<tr>
<th>Layer</th>
<th>Library</th>
<th>Compatibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fortran interface</td>
<td>mkl_blas95.lib</td>
<td>Not applicable¹</td>
</tr>
<tr>
<td></td>
<td>mkl_lapack95.lib</td>
<td></td>
</tr>
<tr>
<td>Interface layer</td>
<td>mkl_intel_c.lib (cdecl)</td>
<td>✓</td>
</tr>
<tr>
<td></td>
<td>mkl_intel_s.lib (stdcall)</td>
<td>✓</td>
</tr>
<tr>
<td>Threading layer</td>
<td>mkl_sequential.lib</td>
<td>✓</td>
</tr>
<tr>
<td></td>
<td>mkl_tbb_thread.lib (if using UWD Intel® TBB ²)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_intel_thread.lib (OpenMP)</td>
<td>×</td>
</tr>
<tr>
<td>Computational layer</td>
<td>mkl_core.lib</td>
<td>✓</td>
</tr>
</tbody>
</table>

¹ These are Fortran 95 wrappers for BLAS (BLAS95) and LAPACK (LAPACK95). Windows drivers are typically written in C, C++ or ASM. Fortran wrappers are UWD-compliant but not useful for building UWDs.

² See Using Intel® TBB in UWP Applications. Starting with the Intel® TBB 2018 release, you can find a prebuilt UWD-compliant Intel® TBB library in `<tbb_distribution_root>\lib\<target_architecture>\vc14_uwd`.

#### Dynamic Linking

<table>
<thead>
<tr>
<th>Layer</th>
<th>Library</th>
<th>Compatibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fortran interface</td>
<td>mkl_blas95_dll.lib</td>
<td>Not applicable¹</td>
</tr>
<tr>
<td></td>
<td>mkl_lapack95_dll.lib</td>
<td></td>
</tr>
<tr>
<td>Interface layer</td>
<td>mkl_intel_c_dll.lib (cdecl)</td>
<td>See UWD Compliance in the compatibility table above.</td>
</tr>
<tr>
<td></td>
<td>mkl_intel_s_dll.lib (stdcall)</td>
<td></td>
</tr>
<tr>
<td>Threading layer</td>
<td>mkl_sequential_dll.lib</td>
<td>See UWD Compliance in the compatibility table above.</td>
</tr>
<tr>
<td></td>
<td>mkl_tbb_thread_dll.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_intel_thread_dll.lib (OpenMP)</td>
<td></td>
</tr>
<tr>
<td>Computational layer</td>
<td>mkl_core_dll.lib</td>
<td>See UWD Compliance in the compatibility table above.</td>
</tr>
</tbody>
</table>

¹ These are Fortran 95 wrappers for BLAS (BLAS95) and LAPACK (LAPACK95). Windows drivers are typically written in C, C++ or ASM. Fortran wrappers are UWD-compliant but not useful for building UWDs.
**UWD Compliance for 64-bit Architecture (intel64)**

These tables describe compliance between Intel® MKL and UWD on 64-bit (intel64) architecture for static and dynamic linked libraries.

### Static Linking

<table>
<thead>
<tr>
<th>Layer</th>
<th>Library</th>
<th>Compatibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fortran interface layer</td>
<td>mkl_blas95_lp64.lib</td>
<td>Not applicable$^1$</td>
</tr>
<tr>
<td></td>
<td>mkl_blas95_ilp64.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_lapack95_lp64.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_lapack95_ilp64.lib</td>
<td></td>
</tr>
<tr>
<td>Interface layer</td>
<td>mkl_intel_lp64.lib</td>
<td>✓</td>
</tr>
<tr>
<td></td>
<td>mkl_intel_ilp64.lib</td>
<td>✓</td>
</tr>
<tr>
<td>Threading layer</td>
<td>mkl_sequential.lib</td>
<td>✓</td>
</tr>
<tr>
<td></td>
<td>mkl_tbb_thread.lib</td>
<td>✓ (if using UWD Intel® TBB$^2$)</td>
</tr>
<tr>
<td></td>
<td>mkl_intel_thread.lib (OpenMP)</td>
<td>×</td>
</tr>
<tr>
<td></td>
<td>mkl_pgi_thread.lib (PGI OpenMP)</td>
<td>×</td>
</tr>
<tr>
<td>Computational layer</td>
<td>mkl_core.lib</td>
<td>✓</td>
</tr>
<tr>
<td>MPI layer</td>
<td>mkl_blacs_lp64.lib</td>
<td>×</td>
</tr>
<tr>
<td></td>
<td>mkl_blacs_ilp64.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_cdft_core.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_scalapack_lp64.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_scalapack_ilp64.lib</td>
<td></td>
</tr>
</tbody>
</table>

$^1$ These are Fortran 95 wrappers for BLAS (BLAS95) and LAPACK (LAPACK95). Windows drivers are typically written in C, C++ or ASM. Fortran wrappers are UWD-compliant but not useful for building UWDs.


### Dynamic Linking

<table>
<thead>
<tr>
<th>Layer</th>
<th>Library</th>
<th>Compatibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fortran interface layer</td>
<td>mkl_blas95_lp64.lib</td>
<td>Not applicable$^1$</td>
</tr>
<tr>
<td></td>
<td>mkl_blas95_ilp64.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_lapack95_lp64.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_lapack95_ilp64.lib</td>
<td></td>
</tr>
<tr>
<td>Interface layer</td>
<td>mkl_intel_lp64_dll.lib</td>
<td>See UWD Compliance in the compatibility table above.</td>
</tr>
<tr>
<td></td>
<td>mkl_intel_ilp64_dll.lib</td>
<td></td>
</tr>
<tr>
<td>Threading layer</td>
<td>mkl_sequential_dll.lib</td>
<td></td>
</tr>
<tr>
<td>Layer</td>
<td>Library</td>
<td>Compatibility</td>
</tr>
<tr>
<td>---------------------</td>
<td>----------------------------------------------</td>
<td>---------------------------------------------------</td>
</tr>
<tr>
<td>Computational layer</td>
<td>mkl_tbb_thread_dll.lib</td>
<td>See <a href="#">UWD Compliance</a> in the compatibility table above.</td>
</tr>
<tr>
<td></td>
<td>mkl_intel_thread_dll.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(OpenMP)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_pgi_thread_dll.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(PGI OpenMP)</td>
<td></td>
</tr>
<tr>
<td>Computational layer</td>
<td>mkl_core_dll.lib</td>
<td>See <a href="#">UWD Compliance</a> in the compatibility table above.</td>
</tr>
<tr>
<td>MPI layer</td>
<td>mkl_blacs_lp64_dll.lib</td>
<td>See <a href="#">UWD Compliance</a> in the compatibility table above.</td>
</tr>
<tr>
<td></td>
<td>mkl_blacs_ilp64_dll.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_cdft_core_dll.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_scalapack_lp64_dll.lib</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mkl_scalapack_ilp64_dll.lib</td>
<td></td>
</tr>
</tbody>
</table>

1 These are Fortran 95 wrappers for BLAS (BLAS95) and LAPACK (LAPACK95). Windows drivers are typically written in C, C++ or ASM. Fortran wrappers are UWD-compliant but not useful for building UWDs.
Managing Performance and Memory

### Optimization Notice

Intel’s compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

### Improving Performance with Threading

Intel® Math Kernel Library (Intel® MKL) is extensively parallelized. See OpenMP* Threaded Functions and Problems and Functions Threaded with Intel® Threading Building Blocks for lists of threaded functions and problems that can be threaded.

Intel® MKL isthread-safe, which means that all Intel® MKL functions (except the LAPACK deprecated routine `lacx`) work correctly during simultaneous execution by multiple threads. In particular, any chunk of threaded Intel® MKL code provides access for multiple threads to the same shared data, while permitting only one thread at any given time to access a shared piece of data. Therefore, you can call Intel® MKL from multiple threads and not worry about the function instances interfering with each other.

If you are using OpenMP* threading technology, you can use the environment variable `OMP_NUM_THREADS` to specify the number of threads or the equivalent OpenMP run-time function calls. Intel® MKL also offers variables that are independent of OpenMP, such as `MKL_NUM_THREADS`, and equivalent Intel® MKL functions for thread management. The Intel® MKL variables are always inspected first, then the OpenMP variables are examined, and if neither is used, the OpenMP software chooses the default number of threads.

By default, Intel® MKL uses the number of OpenMP threads equal to the number of physical cores on the system.

If you are using the Intel TBB threading technology, the OpenMP threading controls, such as the `OMP_NUM_THREADS` environment variable or `MKL_NUM_THREADS` function, have no effect. Use the Intel TBB application programming interface to control the number of threads.

To achieve higher performance, set the number of threads to the number of processors or physical cores, as summarized in Techniques to Set the Number of Threads.

**See Also**

Managing Multi-core Performance

### OpenMP* Threaded Functions and Problems

The following Intel® MKL function domains are threaded with the OpenMP* technology:

- Direct sparse solver.
- LAPACK.
  
  For a list of threaded routines, see LAPACK Routines.
- Level1 and Level2 BLAS.
  
  For a list of threaded routines, see BLAS Level1 and Level2 Routines.
• All Level 3 BLAS and all Sparse BLAS routines except Level 2 Sparse Triangular solvers.
• All Vector Mathematics functions (except service functions).
• FFT.

For a list of FFT transforms that can be threaded, see Threaded FFT Problems.

LAPACK Routines
In this section, ? stands for a precision prefix of each flavor of the respective routine and may have the value of s, d, c, or z.

The following LAPACK routines are threaded with OpenMP*:

• Linear equations, computational routines:
• Orthogonal factorization, computational routines:
• Singular Value Decomposition, computational routines:
  ?gebrd, ?bdsqr
• Symmetric Eigenvalue Problems, computational routines:
• Generalized Nonsymmetric Eigenvalue Problems, computational routines:
  ?chgeqz, ?zhgeqz.

A number of other LAPACK routines, which are based on threaded LAPACK or BLAS routines, make effective use of OpenMP* parallelism:


Threaded BLAS Level1 and Level2 Routines
In the following list, ? stands for a precision prefix of each flavor of the respective routine and may have the value of s, d, c, or z.

The following routines are threaded with OpenMP* for Intel® Core™2 Duo and Intel® Core™ i7 processors:

• Level1 BLAS:
  ?axpy, ?copy, ?swap, ddot/sdot, ddotc, drot/srot
• Level2 BLAS:
  ?gemv, ?trsv, ?trmv, dsysy, dsysr, dsys2/ssyr, dsymv/ssym

Threaded FFT Problems
The following characteristics of a specific problem determine whether your FFT computation may be threaded with OpenMP*:

• rank
• domain
• size/length
• precision (single or double)
• placement (in-place or out-of-place)
• strides
• number of transforms
• layout (for example, interleaved or split layout of complex data)
Most FFT problems are threaded. In particular, computation of multiple transforms in one call (number of transforms > 1) is threaded. Details of which transforms are threaded follow.

**One-dimensional (1D) transforms**

1D transforms are threaded in many cases.

1D complex-to-complex (c2c) transforms of size \( N \) using interleaved complex data layout are threaded under the following conditions depending on the architecture:

<table>
<thead>
<tr>
<th>Architecture</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel® 64</td>
<td>( N ) is a power of 2, ( \log_2(N) &gt; 9 ), the transform is double-precision out-of-place, and input/output strides equal 1.</td>
</tr>
<tr>
<td>IA-32</td>
<td>( N ) is a power of 2, ( \log_2(N) &gt; 13 ), and the transform is single-precision.</td>
</tr>
<tr>
<td></td>
<td>( N ) is a power of 2, ( \log_2(N) &gt; 14 ), and the transform is double-precision.</td>
</tr>
<tr>
<td>Any</td>
<td>( N ) is composite, ( \log_2(N) &gt; 16 ), and input/output strides equal 1.</td>
</tr>
</tbody>
</table>

1D complex-to-complex transforms using split-complex layout are not threaded.

**Multidimensional transforms**

All multidimensional transforms on large-volume data are threaded.

**Functions Threaded with Intel® Threading Building Blocks**

In this section, ? stands for a precision prefix or suffix of the routine name and may have the value of s, d, c, or z.

The following Intel® MKL function domains are threaded with Intel® Threading Building Blocks (Intel® TBB):

- LAPACK.
  - For a list of threaded routines, see LAPACK Routines.
- Entire Level3 BLAS.
- Fast Poisson, Laplace, and Helmholtz Solver (Poisson Library).
- All Vector Mathematics functions (except service functions).
- Intel® MKL PARDISO, a direct sparse solver based on Parallel Direct Sparse Solver (PARDISO*).
  - For details, see Intel® MKL PARDISO Steps.
- Sparse BLAS.
  - For a list of threaded routines, see Sparse BLAS Routines.

**LAPACK Routines**

The following LAPACK routines are threaded with Intel TBB:


A number of other LAPACK routines, which are based on threaded LAPACK or BLAS routines, make effective use of Intel TBB threading:


**Intel® MKL PARDISO Steps**

Intel® MKL PARDISO is threaded with Intel TBB in the reordering and factorization steps. However, routines performing the solving step are still called sequentially when using Intel TBB.
Sparse BLAS Routines

The Sparse BLAS inspector-executor application programming interface routines `mkl_sparse_?_mv` are threaded with Intel TBB for the general compressed sparse row (CSR) and block sparse row (BSR) formats. The following Sparse BLAS inspector-executor application programming routines are threaded with Intel TBB:

- `mkl_sparse_?_mv` using the general compressed sparse row (CSR) and block sparse row (BSR) matrix formats.
- `mkl_sparse_?_mm` using the general CSR sparse matrix format and both row and column major storage formats for the dense matrix.

Avoiding Conflicts in the Execution Environment

Certain situations can cause conflicts in the execution environment that make the use of threads in Intel® MKL problematic. This section briefly discusses why these problems exist and how to avoid them.

If your program is parallelized by other means than Intel® OpenMP* run-time library (RTL) and Intel TBB RTL, several calls to Intel® MKL may operate in a multithreaded mode at the same time and result in slow performance due to overuse of machine resources.

The following table considers several cases where the conflicts may arise and provides recommendations depending on your threading model:

<table>
<thead>
<tr>
<th>Threading model</th>
<th>Discussion</th>
</tr>
</thead>
<tbody>
<tr>
<td>You parallelize the program using the technology other than Intel OpenMP and Intel TBB (for example: Win32* threads on Windows*).</td>
<td>If more than one thread calls Intel® MKL, and the function being called is threaded, it may be important that you turn off Intel® MKL threading. Set the number of threads to one by any of the available means (see Techniques to Set the Number of Threads).</td>
</tr>
<tr>
<td>You parallelize the program using OpenMP directives and/or pragmas and compile the program using a non-Intel compiler.</td>
<td>To avoid simultaneous activities of multiple threading RTLs, link the program against the Intel® MKL threading library that matches the compiler you use (see Linking Examples on how to do this). If this is not possible, use Intel® MKL in the sequential mode. To do this, you should link with the appropriate threading library: <code>mkl_sequential.lib</code> or <code>mkl_sequential.dll</code> (see Appendix C: Directory Structure in Detail).</td>
</tr>
<tr>
<td>You thread the program using Intel TBB threading technology and compile the program using a non-Intel compiler.</td>
<td>To avoid simultaneous activities of multiple threading RTLs, link the program against the Intel® MKL Intel TBB threading library and Intel TBB RTL if it matches the compiler you use. If this is not possible, use Intel® MKL in the sequential mode. To do this, link with the appropriate threading library: <code>mkl_sequential.lib</code> or <code>mkl_sequential_dll.lib</code> (see Appendix C: Directory Structure in Detail).</td>
</tr>
<tr>
<td>You run multiple programs calling Intel® MKL on a multiprocessor system, for example, a program parallelized using a message-passing interface (MPI).</td>
<td>The threading RTLs from different programs you run may place a large number of threads on the same processor on the system and therefore overuse the machine resources. In this case, one of the solutions is to set the number of threads to one by any of the available means (see Techniques to Set the Number of Threads). The Intel® Distribution for LINPACK* Benchmark section discusses another solution for a Hybrid (OpenMP* + MPI) mode.</td>
</tr>
</tbody>
</table>

Using the `mkl_set_num_threads` and `mkl_domain_set_num_threads` functions to control parallelism of Intel® MKL from parallel user threads may result in a race condition that impacts the performance of the application because these functions operate on internal control variables that are global, that is, apply to all threads. For example, if parallel user threads call these functions to set different numbers of threads for the same function domain, the number of threads actually set is unpredictable. To avoid this kind of data races, use the `mkl_set_num_threads_local` function (see the “Support Functions” chapter in the Intel® MKL Developer Reference for the function description).
Techniques to Set the Number of Threads

Use the following techniques to specify the number of OpenMP threads to use in Intel® MKL:

- Set one of the OpenMP or Intel® MKL environment variables:
  - OMP_NUM_THREADS
  - MKL_NUM_THREADS
  - MKL_DOMAIN_NUM_THREADS
- Call one of the OpenMP or Intel® MKL functions:
  - omp_set_num_threads()
  - mkl_set_num_threads()
  - mkl_domain_set_num_threads()
  - mkl_set_num_threads_local()

**NOTE**
A call to the mkl_set_num_threads or mkl_domain_set_num_threads function changes the number of OpenMP threads available to all in-progress calls (in concurrent threads) and future calls to Intel® MKL and may result in slow Intel® MKL performance and/or race conditions reported by run-time tools, such as Intel® Inspector.

To avoid such situations, use the mkl_set_num_threads_local function (see the "Support Functions" section in the Intel® MKL Developer Reference for the function description).

When choosing the appropriate technique, take into account the following rules:

- The Intel® MKL threading controls take precedence over the OpenMP controls because they are inspected first.
- A function call takes precedence over any environment settings. The exception, which is a consequence of the previous rule, is that a call to the OpenMP subroutine omp_set_num_threads() does not have precedence over the settings of Intel® MKL environment variables such as MKL_NUM_THREADS. See Using Additional Threading Control for more details.
- You cannot change run-time behavior in the course of the run using the environment variables because they are read only once at the first call to Intel® MKL.

If you use the Intel TBB threading technology, read the documentation for the tbb::task_scheduler_init class at https://www.threadingbuildingblocks.org/documentation to find out how to specify the number of threads.

Setting the Number of Threads Using an OpenMP* Environment Variable

You can set the number of threads using the environment variable OMP_NUM_THREADS. To change the number of OpenMP threads, in the command shell in which the program is going to run, enter:

```plaintext
set OMP_NUM_THREADS=<number of threads to use>.
```

Some shells require the variable and its value to be exported:

```plaintext
export OMP_NUM_THREADS=<number of threads to use>.
```

You can alternatively assign value to the environment variable using Microsoft Windows*OS Control Panel. Note that you will not benefit from setting this variable on Microsoft Windows* 98 or Windows* ME because multiprocessing is not supported.
See Also
Using Additional Threading Control

Changing the Number of OpenMP* Threads at Run Time

You cannot change the number of OpenMP threads at run time using environment variables. However, you can call OpenMP routines to do this. Specifically, the following sample code shows how to change the number of threads during run time using the `omp_set_num_threads()` routine. For more options, see also Techniques to Set the Number of Threads.

The example is provided for both C and Fortran languages. To run the example in C, use the `omp.h` header file from the Intel(R) compiler package. If you do not have the Intel compiler but wish to explore the functionality in the example, use Fortran API for `omp_set_num_threads()` rather than the C version. For example, `omp_set_num_threads_()`.

```c
// ******** C language ********
#include "omp.h"
#include "mkl.h"
#include <stdio.h>
#define SIZE 1000
int main(int args, char *argv[]){
    double *a, *b, *c;
    a = (double*)malloc(sizeof(double)*SIZE*SIZE);
    b = (double*)malloc(sizeof(double)*SIZE*SIZE);
    c = (double*)malloc(sizeof(double)*SIZE*SIZE);
    double alpha=1, beta=1;
    int m=SIZE, n=SIZE, k=SIZE, lda=SIZE, ldb=SIZE, ldc=SIZE, i=0, j=0;
    char transa='n', transb='n';
    for( i=0; i<SIZE; i++)
    {
        printf("row	a	\n");
        printf("\n%d:	%f	%f\n", i, a[i*SIZE], c[i*SIZE]);
    }
   omp_set_num_threads(1);
    for( i=0; i<SIZE; i++)
    {
        printf("row	a	\n");
        printf("\n%d:	%f	%f\n", i, a[i*SIZE], c[i*SIZE]);
    }
}
```

```fortran
// ******* C language *******
#include "mp.h"
#include "mk1.h"
#include <stdio.h>
#define SIZE 1000
integer function main(int args, character *argv[]){
    integer a, b, c;
    a = (double*)malloc(sizeof(double)*SIZE*SIZE);
    b = (double*)malloc(sizeof(double)*SIZE*SIZE);
    c = (double*)malloc(sizeof(double)*SIZE*SIZE);
    double alpha=1, beta=1;
    integer m=SIZE, n=SIZE, k=SIZE, lda=SIZE, ldb=SIZE, ldc=SIZE, i=0, j=0;
    character transa='n', transb='n';
    do i=0, i<SIZE, i++
    {
        print("row\ta\t\n");
        print("\n%d:	%f	%f\n", i, a(i*SIZE), c(i*SIZE));
    }
    call omp_set_num_threads(1);
    do i=0, i<SIZE, i++
    {
        print("row\ta\t\n");
        print("\n%d:	%f	%f\n", i, a(i*SIZE), c(i*SIZE));
    }
}```
for ( i=0; i<10; i++)
{
    printf("%d: \t%f \t%f\n", i, a[i*SIZE], c[i*SIZE]);
}
omp_set_num_threads(2);
for( i=0; i<SIZE; i++)
{
    for( j=0; j<SIZE; j++)
    {
        a[i*SIZE+j]= (double)(i+j);
        b[i*SIZE+j]= (double)(i*j);
        c[i*SIZE+j]= (double)0;
    }
}
cblas_dgemm(CblasRowMajor, CblasNoTrans, CblasNoTrans,
        m, n, k, alpha, a, lda, b, ldb, beta, c, ldc);
printf("row\ta\tc\n");
for ( i=0;i<10;i++)
{
    printf("%d: \t%f \t%f\n", i, a[i*SIZE],
        c[i*SIZE]);
}
free (a);
free (b);
free (c);
return 0;

// ******** Fortran language ********
PROGRAM DGEMM_DIFF_THREADS
INTEGER N, I, J
PARAMETER (N=100)
REAL*8 A(N,N),B(N,N),C(N,N)
REAL*8 ALPHA, BETA
ALPHA = 1.1
BETA = -1.2
DO I=1,N
    DO J=1,N
        A(I,J) = I+J
        B(I,J) = I*j
        C(I,J) = 0.0
    END DO
END DO
CALL DGEMM('N','N',N,N,N,ALPHA,A,N,B,N,BETA,C,N)
print *, 'Row A C'
DO i=1,10
    write(*,'(I4,F20.8,F20.8)') I, A(1,I),C(1,I)
END DO
CALL OMP_SET_NUM_THREADS(1);
DO I=1,N
    DO J=1,N
        A(I,J) = I+J
        B(I,J) = I*j
        C(I,J) = 0.0
    END DO
Using Additional Threading Control

Intel® MKL-specific Environment Variables for OpenMP Threading Control

Intel® MKL provides environment variables and support functions to control Intel® MKL threading independently of OpenMP. The Intel® MKL-specific threading controls take precedence over their OpenMP equivalents. Use the Intel® MKL-specific threading controls to distribute OpenMP threads between Intel® MKL and the rest of your program.

**NOTE**
Some Intel® MKL routines may use fewer OpenMP threads than suggested by the threading controls if either the underlying algorithms do not support the suggested number of OpenMP threads or the routines perform better with fewer OpenMP threads because of lower OpenMP overhead and/or better data locality. Set the `MKL_DYNAMIC` environment variable to `FALSE` or call `mkl_set_dynamic(0)` to use the suggested number of OpenMP threads whenever the algorithms permit and regardless of OpenMP overhead and data locality.

Section "Number of User Threads" in the "Fourier Transform Functions" chapter of the *Intel® MKL Developer Reference* shows how the Intel® MKL threading controls help to set the number of threads for the FFT computation.

The table below lists the Intel® MKL environment variables for threading control, their equivalent functions, and OMP counterparts:

<table>
<thead>
<tr>
<th>Environment Variable</th>
<th>Support Function</th>
<th>Comment</th>
<th>Equivalent OpenMP* Environment Variable</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>MKL_NUM_THREADS</code></td>
<td><code>mkl_set_num_threads</code></td>
<td>Suggests the number of OpenMP threads to use.</td>
<td><code>OMP_NUM_THREADS</code></td>
</tr>
</tbody>
</table>
### Environment Variable | Support Function | Comment | Equivalent OpenMP* Environment Variable
--- | --- | --- | ---
MKL_DOMAIN_NUM_THREADS | mkl_domain_set_num_threads | Suggests the number of OpenMP threads for a particular function domain. |  
MKL_DYNAMIC | mkl_set_dynamic | Enables Intel® MKL to dynamically change the number of OpenMP threads. | OMP_DYNAMIC

**NOTE**

Call `mkl_set_num_threads()` to force Intel® MKL to use a given number of OpenMP threads and prevent it from reacting to the environment variables `MKL_NUM_THREADS`, `MKL_DOMAIN_NUM_THREADS`, and `OMP_NUM_THREADS`.

The example below shows how to force Intel® MKL to use one thread:

```c
// ******* C language *******
#include <mkl.h>
...
mkl_set_num_threads( 1 );
```

```fortran
// ******* Fortran language *******
...
call mkl_set_num_threads( 1 )
```

See the *Intel® MKL Developer Reference* for the detailed description of the threading control functions, their parameters, calling syntax, and more code examples.

**MKL_DYNAMIC**

The `MKL_DYNAMIC` environment variable enables Intel® MKL to dynamically change the number of threads.

The default value of `MKL_DYNAMIC` is **TRUE**, regardless of `OMP_DYNAMIC`, whose default value may be **FALSE**. When `MKL_DYNAMIC` is **TRUE**, Intel® MKL may use fewer OpenMP threads than the maximum number you specify.

For example, `MKL_DYNAMIC` set to **TRUE** enables optimal choice of the number of threads in the following cases:

- If the requested number of threads exceeds the number of physical cores (perhaps because of using the Intel® Hyper-Threading Technology), Intel® MKL scales down the number of OpenMP threads to the number of physical cores.
- If you are able to detect the presence of a message-passing interface (MPI), but cannot determine whether it has been called in a thread-safe mode, Intel® MKL runs one OpenMP thread.

When `MKL_DYNAMIC` is **FALSE**, Intel® MKL uses the suggested number of OpenMP threads whenever the underlying algorithms permit. For example, if you attempt to do a size one matrix-matrix multiply across eight threads, the library may instead choose to use only one thread because it is impractical to use eight threads in this event.
If Intel® MKL is called from an OpenMP parallel region in your program, Intel® MKL uses only one thread by default. If you want Intel® MKL to go parallel in such a call, link your program against an OpenMP threading RTL supported by Intel® MKL and set the environment variables:

- **OMP_NESTED** to **TRUE**
- **OMP_DYNAMIC** and **MKL_DYNAMIC** to **FALSE**
- **MKL_NUM_THREADS** to some reasonable value

With these settings, Intel® MKL uses **MKL_NUM_THREADS** threads when it is called from the OpenMP parallel region in your program.

In general, set **MKL_DYNAMIC** to **FALSE** only under circumstances that Intel® MKL is unable to detect, for example, to use nested parallelism where the library is already called from a parallel section.

### MKL_DOMAIN_NUM_THREADS

The **MKL_DOMAIN_NUM_THREADS** environment variable suggests the number of OpenMP threads for a particular function domain.

**MKL_DOMAIN_NUM_THREADS** accepts a string value `<MKL-env-string>`, which must have the following format:

```
<MKL-env-string> ::= <MKL-domain-env-string> { <delimiter><MKL-domain-env-string> }
<delimiter> ::= [ <space-symbol>* ] ( <space-symbol> | <comma-symbol> | <semicolon-symbol> | <colon-symbol> ) [ <space-symbol>* ]
<MKL-domain-env-string> ::= <MKL-domain-env-name><uses><number-of-threads>
<MKL-domain-env-name> ::= MKL_DOMAIN_ALL | MKL_DOMAIN_BLAS | MKL_DOMAIN_FFT | MKL_DOMAIN_VML | MKL_DOMAIN_PARDISO
<uses> ::= [ <space-symbol>* ] ( <space-symbol> | <equality-sign> | <comma-symbol>) [ <space-symbol>* ]
<number-of-threads> ::= <positive-number>
<positive-number> ::= <decimal-positive-number> | <octal-number> | <hexadecimal-number>
```

In the syntax above, values of `<MKL-domain-env-name>` indicate function domains as follows:

- **MKL_DOMAIN_ALL** — All function domains
- **MKL_DOMAIN_BLAS** — BLAS Routines
- **MKL_DOMAIN_FFT** — non-cluster Fourier Transform Functions
- **MKL_DOMAIN_LAPACK** — LAPACK Routines
- **MKL_DOMAIN_VML** — Vector Mathematics (VM)
- **MKL_DOMAIN_PARDISO** — Intel® MKL PARDISO, a direct sparse solver based on Parallel Direct Sparse Solver (PARDISO*)

For example, you could set the **MKL_DOMAIN_NUM_THREADS** environment variable to any of the following string variants, in this case, defining three specific domain variables internal to Intel® MKL:

```
MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL=2, MKL_DOMAIN_BLAS=1, MKL_DOMAIN_FFT=4"
MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL 2 : MKL_DOMAIN_BLAS 1 : MKL_DOMAIN_FFT 4"
MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL=2 : MKL_DOMAIN_BLAS=1 : MKL_DOMAIN_FFT=4"
MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL=2; MKL_DOMAIN_BLAS=1; MKL_DOMAIN_FFT=4"
MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL=2 MKL_DOMAIN_BLAS 1, MKL_DOMAIN_FFT 4"
MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL,2: MKL_DOMAIN_BLAS 1, MKL_DOMAIN_FFT,4"
```
NOTE Prepend the appropriate set/export/setenv command for your command shell and operating system. Refer to Setting the Environment Variables for Threading Control for more details.

The global variables MKL_DOMAIN_ALL, MKL_DOMAIN_BLAS, MKL_DOMAIN_FFT, MKL_DOMAIN_VML, and MKL_DOMAIN_PARDISO, as well as the interface for the Intel® MKL threading control functions, can be found in the mkl.h header file.

NOTE You can retrieve the values of the specific domain variables that you have set in your code with a call to the mkl_get_domain_max_threads(domain_name) function per the Fortran and C interface with the desired domain variable name.

This table illustrates how values of MKL_DOMAIN_NUM_THREADS are interpreted.

<table>
<thead>
<tr>
<th>Value of MKL_DOMAIN_NUM_THREADS</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>MKL_DOMAIN_ALL=4</td>
<td>All parts of Intel® MKL should try four OpenMP threads. The actual number of threads may be still different because of the MKL_DYNAMIC setting or system resource issues. The setting is equivalent to MKL_NUM_THREADS = 4.</td>
</tr>
<tr>
<td>MKL_DOMAIN_ALL=1, MKL_DOMAIN_BLAS=4</td>
<td>All parts of Intel® MKL should try one OpenMP thread, except for BLAS, which is suggested to try four threads.</td>
</tr>
<tr>
<td>MKL_DOMAIN_VML=2</td>
<td>VM should try two OpenMP threads. The setting affects no other part of Intel® MKL.</td>
</tr>
</tbody>
</table>

Be aware that the domain-specific settings take precedence over the overall ones. For example, the "MKL_DOMAIN_BLAS=4" value of MKL_DOMAIN_NUM_THREADS suggests trying four OpenMP threads for BLAS, regardless of later setting MKL_NUM_THREADS, and a function call "mkl_domain_set_num_threads ( 4, MKL_DOMAIN_BLAS );" suggests the same, regardless of later calls to mkl_set_num_threads(). However, a function call with input "MKL_DOMAIN_ALL", such as "mkl_domain_set_num_threads (4, MKL_DOMAIN_ALL);" is equivalent to "mkl_set_num_threads(4)", and thus it will be overwritten by later calls to mkl_set_num_threads. Similarly, the environment setting of MKL_DOMAIN_NUM_THREADS with "MKL_DOMAIN_ALL=4" will be overwritten with MKL_NUM_THREADS = 2.

Whereas the MKL_DOMAIN_NUM_THREADS environment variable enables you set several variables at once, for example, "MKL_DOMAIN_BLAS=4, MKL_DOMAIN_FFT=2", the corresponding function does not take string syntax. So, to do the same with the function calls, you may need to make several calls, which in this example are as follows:

mkl_domain_set_num_threads ( 4, MKL_DOMAIN_BLAS );
mkl_domain_set_num_threads ( 2, MKL_DOMAIN_FFT );

MKL_NUM_STRIPES

The MKL_NUM_STRIPES environment variable controls the Intel® MKL threading algorithm for ?gemm functions. When MKL_NUM_STRIPES is set to a positive integer value nstripes, Intel® MKL tries to use a number of partitions equal tonstripes along the leading dimension of the output matrix.

The following table explains how the value nstripes of MKL_NUM_STRIPES defines the partitioning algorithm used by Intel® MKL for ?gemm output matrix; max_threads_for_mkl denotes the maximum number of OpenMP threads for Intel® MKL:
Value of MKL_NUM_STRIPES | Partitioning Algorithm
--- | ---
$1 < nstripes < \frac{\text{max\_threads\_for\_mkl}}{2}$ | 2D partitioning with the number of partitions equal to $nstripes$:
- Horizontal, for column-major ordering.
- Vertical, for row-major ordering.

$nstripes = 1$ | 1D partitioning algorithm along the opposite direction of the leading dimension.

$nstripes \geq \frac{\text{max\_threads\_for\_mkl}}{2}$ | 1D partitioning algorithm along the leading dimension.

$nstripes < 0$ | The default Intel® MKL threading algorithm.

The following figure shows the partitioning of an output matrix for $nstripes = 4$ and a total number of 8 OpenMP threads for column-major and row-major orderings:

You can use support functions mkl_set_num_stripes and mkl_get_num_stripes to set and query the number of stripes, respectively.

**Setting the Environment Variables for Threading Control**

To set the environment variables used for threading control, in the command shell in which the program is going to run, enter:

```bash
set <VARIABLE NAME>=<value>
```

For example:

```bash
set MKL_NUM_THREADS=4
set MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL=1, MKL_DOMAIN_BLAS=4"
set MKL_DYNAMIC=FALSE
set MKL_NUM_STRIPES=4
```

Some shells require the variable and its value to be exported:

```bash
export <VARIABLE NAME>=<value>
```
For example:

```plaintext
export MKL_NUM_THREADS=4
export MKL_DOMAIN_NUM_THREADS="MKL_DOMAIN_ALL=1, MKL_DOMAIN_BLAS=4"
export MKL_DYNAMIC=FALSE
export MKL_NUM_STRIPES=4
```

You can alternatively assign values to the environment variables using Microsoft Windows* OS Control Panel.

### Calling Intel® MKL Functions from Multi-threaded Applications

This section summarizes typical usage models and available options for calling Intel® MKL functions from multi-threaded applications. These recommendations apply to any multi-threading environments: OpenMP*, Intel® Threading Building Blocks, Windows* threads, and others.

#### Usage model: disable Intel® MKL internal threading for the whole application

**When used:** Intel® MKL internal threading interferes with application's own threading or may slow down the application.

**Example:** the application is threaded at top level, or the application runs concurrently with other applications.

**Options:**

- Link statically or dynamically with the sequential library
- Link with the Single Dynamic Library `mkl_rt.lib` and select the sequential library using an environment variable or a function call:
  - Set `MKL_THREADING_LAYER=sequential`
  - Call `mkl_set_threading_layer(MKL_THREADING_SEQUENTIAL)`‡

#### Usage model: partition system resources among application threads

**When used:** application threads are specialized for a particular computation.

**Example:** one thread solves equations on all cores but one, while another thread running on a single core updates a database.

**Linking Options:**

- Link statically or dynamically with a threading library
- Link with the Single Dynamic Library `mkl_rt.lib` and select a threading library using an environment variable or a function call:
  - `set MKL_THREADING_LAYER=intel` or `MKL_THREADING_LAYER=tbb`
  - `call mkl_set_threading_layer(MKL_THREADING_INTEL)` or `mkl_set_threading_layer(MKL_THREADING_TBB)`

**Other Options for OpenMP Threading:**

- Set the `MKL_NUM_THREADS` environment variable to a desired number of OpenMP threads for Intel® MKL.
- Set the `MKL_DOMAIN_NUM_THREADS` environment variable to a desired number of OpenMP threads for Intel® MKL for a particular function domain.

Use if the application threads work with different Intel® MKL function domains.

- Call `mkl_set_num_threads()`

Use to globally set a desired number of OpenMP threads for Intel® MKL at run time.

- Call `mkl_domain_set_num_threads()`.

Use if at some point application threads start working with different Intel® MKL function domains.
• Call `mkl_set_num_threads_local()`.

  Use to set the number of OpenMP threads for Intel® MKL called from a particular thread.

  **NOTE**
  If your application uses OpenMP* threading, you may need to provide additional settings:

  • Set the environment variable `OMP_NESTED=TRUE`, or alternatively call `omp_set_nested(1)`, to enable OpenMP nested parallelism.
  • Set the environment variable `MKL_DYNAMIC=FALSE`, or alternatively call `mkl_set_dynamic(0)`, to prevent Intel® MKL from dynamically reducing the number of OpenMP threads in nested parallel regions.

  For details of the mentioned functions, see the Support Functions section of the *Intel® MKL Developer Reference*, available in the Intel Software Documentation Library.

<table>
<thead>
<tr>
<th>Optimization Notice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.</td>
</tr>
<tr>
<td>Notice revision #20110804</td>
</tr>
</tbody>
</table>

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

**See Also**

- Linking with Threading Libraries
- Dynamically Selecting the Interface and Threading Layer
- Intel® MKL-specific Environment Variables for OpenMP Threading Control
- `MKL_DOMAIN_NUM_THREADS`
- Avoiding Conflicts in the Execution Environment
- Intel Software Documentation Library

**Using Intel® Hyper-Threading Technology**

Intel® Hyper-Threading Technology (Intel® HT Technology) is especially effective when each thread performs different types of operations and when there are under-utilized resources on the processor. However, Intel® MKL fits neither of these criteria because the threaded portions of the library execute at high efficiencies using most of the available resources and perform identical operations on each thread. You may obtain higher performance by disabling Intel HT Technology.

If you run with Intel HT Technology enabled, performance may be especially impacted if you run on fewer threads than physical cores. Moreover, if, for example, there are two threads to every physical core, the thread scheduler may assign two threads to some cores and ignore the other cores altogether. If you are using the OpenMP* library of the Intel Compiler, read the respective User Guide on how to best set the thread affinity interface to avoid this situation. For Intel® MKL, apply the following setting:

```bash
export KMP_AFFINITY=granularity=fine,compact,1,0
```

If you are using the Intel TBB threading technology, read the documentation on the `tbb::affinity_partitioner` class at [https://www.threadingbuildingblocks.org/documentation](https://www.threadingbuildingblocks.org/documentation) to find out how to affinitize Intel TBB threads.
Managing Multi-core Performance

You can obtain best performance on systems with multi-core processors by requiring that threads do not migrate from core to core. To do this, bind threads to the CPU cores by setting an affinity mask to threads. Use one of the following options:

- OpenMP facilities (if available), for example, the `KMP_AFFINITY` environment variable using the Intel OpenMP library
- A system function, as explained below
- Intel TBB facilities (if available), for example, the `tbb::affinity_partitioner` class (for details, see https://www.threadingbuildingblocks.org/documentation)

Consider the following performance issue:

- The system has two sockets with two cores each, for a total of four cores (CPUs).
- The application sets the number of OpenMP threads to four and calls an Intel® MKL LAPACK routine. This call takes considerably different amounts of time from run to run.

To resolve this issue, before calling Intel® MKL, set an affinity mask for each OpenMP thread using the `KMP_AFFINITY` environment variable or the `SetThreadAffinityMask` system function. The following code example shows how to resolve the issue by setting an affinity mask by operating system means using the Intel compiler. The code calls the function `SetThreadAffinityMask` to bind the threads to appropriate cores, preventing migration of the threads. Then the Intel® MKL LAPACK routine is called:

```c
// Set affinity mask
#include <windows.h>
#include <omp.h>
int main(void) {
#pragma omp parallel default(shared)
{
  int tid = omp_get_thread_num();
  // 2 packages x 2 cores/pkg x 1 threads/core (4 total cores)
  DWORD_PTR mask = (1 << (tid == 0 ? 0 : 2));
  SetThreadAffinityMask( GetCurrentThread(), mask );
}
// Call Intel MKL LAPACK routine
return 0;
}
```

Compile the application with the Intel compiler using the following command:

```bash
icl /Qopenmp test_application.c
```

where `test_application.c` is the filename for the application.

Build the application. Run it in four threads, for example, by using the environment variable to set the number of threads:

```bash
set OMP_NUM_THREADS=4
test_application.exe
```

See Windows API documentation at msdn.microsoft.com/ for the restrictions on the usage of Windows API routines and particulars of the `SetThreadAffinityMask` function used in the above example.

See also a similar example at en.wikipedia.org/wiki/Affinity_mask.
Improving Performance for Small Size Problems

The overhead of calling an Intel® MKL function for small problem sizes can be significant when the function has a large number of parameters or internally checks parameter errors. To reduce the performance overhead for these small size problems, the Intel® MKL direct call feature works in conjunction with the compiler to preprocess the calling parameters to supported Intel® MKL functions and directly call or inline special optimized small-matrix kernels that bypass error checking. For a list of functions supporting direct call, see Limitations of the Direct Call.

To activate the feature, do the following:

• Compile your C or Fortran code with the preprocessor macro depending on whether a threaded or sequential mode of Intel® MKL is required by supplying the compiler option as explained below:

<table>
<thead>
<tr>
<th>Intel® MKL Mode</th>
<th>Macro</th>
<th>Compiler Option</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threaded</td>
<td>MKL_DIRECT_CALL</td>
<td>/DMKL_DIRECT_CALL</td>
</tr>
<tr>
<td>Sequential</td>
<td>MKL_DIRECT_CALL_SEQ</td>
<td>/DMKL_DIRECT_CALL_SEQ</td>
</tr>
</tbody>
</table>

• For Fortran applications:
  • Enable preprocessor by using the /fpp option for Intel® Fortran Compiler and -Mpreprocess option for PGI* compilers.
  • Include the Intel® MKL Fortran include file mkl_direct_call.fi.

Intel® MKL skips error checking and intermediate function calls if the problem size is small enough (for example: a call to a function that supports direct call, such as dgemm, with matrix ranks smaller than 50).

Using MKL_DIRECT_CALL in C Applications

The following examples of code and link lines show how to activate direct calls to Intel® MKL kernels in C applications:

• Include the mkl.h header file:

```c
#include "mkl.h"
int main(void) {

    // Call Intel MKL Dgemm

    return 0;
}
```

• For multi-threaded Intel® MKL, compile with MKL_DIRECT_CALL preprocessor macro:

```
icl /DMKL_DIRECT_CALL /Qstd=c99 your_application.c mkl_intel_lp64.lib mkl_core.lib
mkl_intel_thread.lib /Qopenmp -I%MKLROOT%/include
```

• To use Intel® MKL in the sequential mode, compile with MKL_DIRECT_CALL_SEQ preprocessor macro:

```
icl /DMKL_DIRECT_CALL_SEQ /Qstd=c99 your_application.c mkl_intel_lp64.lib mkl_core.lib
```
Using MKL_DIRECT_CALL in Fortran Applications

The following examples of code and link lines show how to activate direct calls to Intel® MKL kernels in Fortran applications:

- Include mkl_direct_call.fi, to be preprocessed by the Fortran compiler preprocessor

```fortran
include "mkl_direct_call.fi"
program DGEMM_MAIN
    ...
    * Call Intel MKL DGEMM
    ...
    call sub1()
    stop 1
end

* A subroutine that calls DGEMM
subroutine sub1
    *
    * Call Intel MKL DGEMM
end
```

- For multi-threaded Intel® MKL, compile with /fpp option for Intel Fortran compiler (or with -Mpreprocess for PGI compilers) and with MKL_DIRECT_CALL preprocessor macro:

```
ifort /DMKL_DIRECT_CALL /fpp your_application.f mkl_intel_lp64.lib mkl_core.lib
    mkl_intel_thread.lib
    /Qopenmp -I%MKLROOT%/include
```

- To use Intel® MKL in the sequential mode, compile with /fpp option for Intel Fortran compiler (or with -Mpreprocess for PGI compilers) and with MKL_DIRECT_CALL_SEQ preprocessor macro:

```
ifort /DMKL_DIRECT_CALL_SEQ /fpp your_application.f mkl_intel_lp64.lib mkl_core.lib
    mkl_sequential.lib
    -I%MKLROOT%/include
```

Limitations of the Direct Call

Directly calling the Intel® MKL kernels has the following limitations:

- If the MKL_DIRECT_CALL or MKL_DIRECT_CALL_SEQ macro is used, Intel® MKL may skip error checking.

  **Important**
  
  With a limited error checking, you are responsible for checking the correctness of function parameters to avoid unsafe and incorrect code execution.

- The feature is only available for the following functions:
  - Intel® MKL Verbose mode, Conditional Numerical Reproducibility, and BLAS95 interfaces are not supported.
  - GNU* Fortran compilers are not supported.
  - For C applications, you must enable mixing declarations and user code by providing the /Qstd=c99 option for Intel® compilers.
  - In a fixed format Fortran source code compiled with PGI compilers, the lines containing Intel® MKL functions must end at least seven columns before the line ending column, usually, in a column with the index not greater than 72 – 7 = 65.
NOTE
The direct call feature substitutes the names of Intel® MKL functions with longer counterparts, which can cause the lines to exceed the column limit for a fixed format Fortran source code compiled with PGI compilers. Because the compilers ignore any part of the line that exceeds the limit, the behavior of the program can be unpredictable.

Other Tips and Techniques to Improve Performance

See Also
Managing Performance of the Cluster Fourier Transform Functions

Coding Techniques
This section discusses coding techniques to improve performance on processors based on supported architectures.

To improve performance, properly align arrays in your code. Additional conditions can improve performance for specific function domains.

Data Alignment and Leading Dimensions
To improve performance of your application that calls Intel® MKL, align your arrays on 64-byte boundaries and ensure that the leading dimensions of the arrays are divisible by 64/element_size, where element_size is the number of bytes for the matrix elements (4 for single-precision real, 8 for double-precision real and single-precision complex, and 16 for double-precision complex). For more details, see Example of Data Alignment.

For Intel® Xeon Phi™ processor x200 product family, codenamed Knights Landing, align your matrices on 4096-byte boundaries and set the leading dimension to the following integer expression:

\[
(((n \times \text{element\_size} + 511) / 512) \times 512 + 64) / \text{element\_size},
\]

where \(n\) is the matrix dimension along the leading dimension.

LAPACK Packed Routines
The routines with the names that contain the letters HP, OP, PP, SP, TP, UP in the matrix type and storage position (the second and third letters respectively) operate on the matrices in the packed format (see LAPACK "Routine Naming Conventions" sections in the Intel® MKL Developer Reference). Their functionality is strictly equivalent to the functionality of the unpacked routines with the names containing the letters HE, OR, PO, SY, TR, UN in the same positions, but the performance is significantly lower.

If the memory restriction is not too tight, use an unpacked routine for better performance. In this case, you need to allocate \(N^2/2\) more memory than the memory required by a respective packed routine, where \(N\) is the problem size (the number of equations).

For example, to speed up solving a symmetric eigenproblem with an expert driver, use the unpacked routine:

```fortran
  call dsyevx(jobz, range, uplo, n, a, lda, vl, vu, il, iu, abstol, m, w, z, ldz, work, lwork, iwork, ifail, info)
```

where \(a\) is the dimension \(lda\)-by-\(n\), which is at least \(N^2\) elements,
instead of the packed routine:

```fortran
    call dspevx(jobz, range, uplo, n, ap, vl, vu, il, iu, abstol, m, w, z, ldz, work, iwork, ifail, info)
```

where `ap` is the dimension $N^*(N+1)/2$.

**See Also**
Managing Performance of the Cluster Fourier Transform Functions

### Improving Intel(R) MKL Performance on Specific Processors

#### Dual-Core Intel® Xeon® Processor 5100 Series

To get the best performance with Intel® MKL on Dual-Core Intel® Xeon® processor 5100 series systems, enable the Hardware DPL (streaming data) Prefetcher functionality of this processor. To configure this functionality, use the appropriate BIOS settings, as described in your BIOS documentation.

### Operating on Denormals

The IEEE 754-2008 standard, "An IEEE Standard for Binary Floating-Point Arithmetic", defines denormal (or subnormal) numbers as non-zero numbers smaller than the smallest possible normalized numbers for a specific floating-point format. Floating-point operations on denormals are slower than on normalized operands because denormal operands and results are usually handled through a software assist mechanism rather than directly in hardware. This software processing causes Intel® MKL functions that consume denormals to run slower than with normalized floating-point numbers.

You can mitigate this performance issue by setting the appropriate bit fields in the MXCSR floating-point control register to flush denormals to zero (FTZ) or to replace any denormals loaded from memory with zero (DAZ). Check your compiler documentation to determine whether it has options to control FTZ and DAZ. Note that these compiler options may slightly affect accuracy.

### Using Memory Functions

#### Avoiding Memory Leaks in Intel® MKL

When running, Intel® MKL allocates and deallocates internal buffers to facilitate better performance. However, in some cases this behavior may result in memory leaks. To avoid memory leaks, you can do either of the following:

- Set the `MKL_DISABLE_FAST_MM` environment variable to 1 or call the `mkl_disable_fast_mm()` function. Be aware that this change may negatively impact performance of some Intel® MKL functions, especially for small problem sizes.
- Call the `mkl_free_buffers()` function or the `mkl_thread_free_buffers()` function in the current thread.

For the descriptions of the memory functions, see the Intel® MKL Developer Reference, available in the Intel Software Documentation Library.

**See Also**
Intel Software Documentation Library
Redefining Memory Functions

In C/C++ programs, you can replace Intel® MKL memory functions that the library uses by default with your own functions. To do this, use the **memory renaming** feature.

Memory Renaming

Intel® MKL memory management by default uses standard C run-time memory functions to allocate or free memory. These functions can be replaced using memory renaming.

Intel® MKL accesses the memory functions by pointers: `i_malloc`, `i_free`, `i_calloc`, and `i_realloc`, which are visible at the application level. These pointers initially hold addresses of the standard C run-time memory functions `malloc`, `free`, `calloc`, and `realloc`, respectively. You can programmatically redefine values of these pointers to the addresses of your application’s memory management functions.

Redirecting the pointers is the only correct way to use your own set of memory management functions. If you call your own memory functions without redirecting the pointers, the memory will get managed by two independent memory management packages, which may cause unexpected memory issues.

How to Redefine Memory Functions

To redefine memory functions, use the following procedure:

If you are using the statically linked Intel® MKL,

1. Include the `i_malloc.h` header file in your code.
   This header file contains all declarations required for replacing the memory allocation functions. The header file also describes how memory allocation can be replaced in those Intel libraries that support this feature.
2. Redefine values of pointers `i_malloc`, `i_free`, `i_calloc`, and `i_realloc` prior to the first call to Intel® MKL functions, as shown in the following example:

   ```c
   #include "i_malloc.h"
   ...
   i_malloc = my_malloc;
   i_calloc = my_calloc;
   i_realloc = my_realloc;
   i_free = my_free;
   ...
   // Now you may call Intel MKL functions
   ```

If you are using the dynamically linked Intel® MKL,

1. Include the `i_malloc.h` header file in your code.
2. Redefine values of pointers `i_malloc_dll`, `i_free_dll`, `i_calloc_dll`, and `i_realloc_dll` prior to the first call to Intel® MKL functions, as shown in the following example:

   ```c
   #include "i_malloc.h"
   ...
   i_malloc_dll = my_malloc;
   i_calloc_dll = my_calloc;
   i_realloc_dll = my_realloc;
   i_free_dll = my_free;
   ...
   // Now you may call Intel MKL functions
   ```
The Intel® Math Kernel Library (Intel® MKL) provides broad support for Fortran and C/C++ programming. However, not all functions support both Fortran and C interfaces. For example, some LAPACK functions have no C interface. You can call such functions from C using mixed-language programming.

If you want to use LAPACK or BLAS functions that support Fortran 77 in the Fortran 95 environment, additional effort may be initially required to build compiler-specific interface libraries and modules from the source code provided with Intel® MKL.

### Optimization Notice

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

### See Also

Language Interfaces Support, by Function Domain

### Using Language-Specific Interfaces with Intel® Math Kernel Library

This section discusses mixed-language programming and the use of language-specific interfaces with Intel® MKL.

See also the "FFTW Interface to Intel® Math Kernel Library" Appendix in the Intel® MKL Developer Reference for details of the FFTW interfaces to Intel® MKL.

### Interface Libraries and Modules

You can create the following interface libraries and modules using the respective makefiles located in the interfaces directory.

<table>
<thead>
<tr>
<th>File name</th>
<th>Contains</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_blas95.lib</td>
<td>Fortran 95 wrappers for BLAS (BLAS95) for IA-32 architecture.</td>
</tr>
<tr>
<td>mkl_blas95_ilp64.lib</td>
<td>Fortran 95 wrappers for BLAS (BLAS95) supporting LP64 interface.</td>
</tr>
<tr>
<td>mkl_blas95_lp64.lib</td>
<td>Fortran 95 wrappers for BLAS (BLAS95) supporting ILP64 interface.</td>
</tr>
</tbody>
</table>
### File name | Contains
---|---
mkl_lapack95.lib<sup>1</sup> | Fortran 95 wrappers for LAPACK (LAPACK95) for IA-32 architecture.
mkl_lapack95_lp64.lib<sup>1</sup> | Fortran 95 wrappers for LAPACK (LAPACK95) supporting LP64 interface.
mkl_lapack95_ilp64.lib<sup>1</sup> | Fortran 95 wrappers for LAPACK (LAPACK95) supporting ILP64 interface.
fft2xc_intel.lib<sup>1</sup> | Interfaces for FFTW version 2.x (C interface for Intel compilers) to call Intel® MKL FFT.
fft2xc_ms.lib | Contains interfaces for FFTW version 2.x (C interface for Microsoft compilers) to call Intel® MKL FFT.
fft2xf_intel.lib | Interfaces for FFTW version 2.x (Fortran interface for Intel compilers) to call Intel® MKL FFT.
fft3xc_intel.lib<sup>2</sup> | Interfaces for FFTW version 3.x (C interface for Intel compiler) to call Intel® MKL FFT.
fft3xc_ms.lib | Interfaces for FFTW version 3.x (C interface for Microsoft compilers) to call Intel® MKL FFT.
fft3xf_intel.lib<sup>2</sup> | Interfaces for FFTW version 3.x (Fortran interface for Intel compilers) to call Intel® MKL FFT.
fft2x_cdft_SINGLE.lib | Single-precision interfaces for MPI FFTW version 2.x (C interface) to call Intel® MKL cluster FFT.
fft2x_cdft_DOUBLE.lib | Double-precision interfaces for MPI FFTW version 2.x (C interface) to call Intel® MKL cluster FFT.
fft3x_cdft.lib | Interfaces for MPI FFTW version 3.x (C interface) to call Intel® MKL cluster FFT.
fft3x_cdft_ilp64.lib | Interfaces for MPI FFTW version 3.x (C interface) to call Intel® MKL cluster FFT supporting the ILP64 interface.

### Modules, in architecture- and interface-specific subdirectories of the Intel® MKL include directory

- **blas95.mod<sup>1</sup>** | Fortran 95 interface module for BLAS (BLAS95).
- **lapack95.mod<sup>1</sup>** | Fortran 95 interface module for LAPACK (LAPACK95).
- **f95_precision.mod<sup>1</sup>** | Fortran 95 definition of precision parameters for BLAS95 and LAPACK95.
- **mkl_service.mod<sup>1</sup>** | Fortran 95 interface module for Intel® MKL support functions.

<sup>1</sup> Prebuilt for the Intel® Fortran compiler

<sup>2</sup> FFTW3 interfaces are integrated with Intel® MKL. Look into `<mkl directory>\interfaces\fftw3x*.makefile` for options defining how to build and where to place the standalone library with the wrappers.

**See Also**

Fortran 95 Interfaces to LAPACK and BLAS
Fortran 95 Interfaces to LAPACK and BLAS

Fortran 95 interfaces are compiler-dependent. Intel® MKL provides the interface libraries and modules precompiled with the Intel® Fortran compiler. Additionally, the Fortran 95 interfaces and wrappers are delivered as sources. (For more information, see Compiler-dependent Functions and Fortran 90 Modules). If you are using a different compiler, build the appropriate library and modules with your compiler and link the library as a user’s library:

1. Go to the respective directory `{mkl directory}`/interfaces/blas95 or `{mkl directory}`/interfaces/lapack95
2. Type:
   - For the IA 32 architecture, make libia32 install_dir =<user_dir>
   - nmake libintel64 [interface=lp64|ilp64] install_dir=<user dir>

   **Important**
   The parameter install_dir is required.

As a result, the required library is built and installed in the `<user dir>`/lib directory, and the .mod files are built and installed in the `<user dir>`/include/arch/[{lp64|ilp64}] directory, where arch is `{ia32, intel64}`.

By default, the ifort compiler is assumed. You may change the compiler with an additional parameter of nmake:

FC=<compiler>.

For example, the command

nmake libintel64 FC=f95 install_dir=<userf95 dir> interface=lp64

builds the required library and .mod files and installs them in subdirectories of `<userf95 dir>`.

To delete the library from the building directory, type:

- For the IA-32 architecture, make cleania32 INSTALL_DIR=<user_dir>
- nmake cleanintel64 [interface=lp64|ilp64] install_dir=<user dir>
- make clean INSTALL_DIR=<user_dir>

   **Caution**
   Even if you have administrative rights, avoid setting install_dir=..\.. or install_dir={mkl directory} in a build or clean command above because these settings replace or delete the Intel® MKL prebuilt Fortran 95 library and modules.

Compiler-dependent Functions and Fortran 90 Modules

Compiler-dependent functions occur whenever the compiler inserts into the object code function calls that are resolved in its run-time library (RTL). Linking of such code without the appropriate RTL will result in undefined symbols. Intel® MKL has been designed to minimize RTL dependencies.

In cases where RTL dependencies might arise, the functions are delivered as source code and you need to compile the code with whatever compiler you are using for your application.

In particular, Fortran 90 modules result in the compiler-specific code generation requiring RTL support. Therefore, Intel® MKL delivers these modules compiled with the Intel compiler, along with source code, to be used with different compilers.
Using the stdcall Calling Convention in C/C++

Intel® MKL supports stdcall calling convention for the following function domains:

- BLAS, except CBLAS, compact API, and JIT API
- Sparse BLAS
- LAPACK
- Vector Mathematics
- Vector Statistics (VS)
- Intel® MKL PARDISO, a direct sparse solver based on Parallel Direct Sparse Solver (PARDISO*)
- Direct Sparse Solvers
- RCI Iterative Solvers for Linear Problems (Iterative Sparse Solvers based on Reverse Communication Interface)
- Support Functions

To use the stdcall calling convention in C/C++, follow the guidelines below:

- In your function calls, pass lengths of character strings to the functions. For example, compare the following calls to the VS function vslLoadStreamF:

  cdecl:
  ```c
  errstatus = vslLoadStreamF(&stream, "streamfile.bin");
  ```

  stdcall:
  ```c
  errstatus = vslLoadStreamF(&stream, "streamfile.bin", 14);
  ```

- Define the MKL_STDCALL macro using either of the following techniques:
  - Define the macro in your source code before including Intel® MKL header files:
    ```c
    #define MKL_STDCALL
    #include "mkl.h"
    ```
  - Pass the macro to the compiler. For example:
    ```c
    icl -DMKL_STDCALL foo.c
    ```

- Link your application with the following library:
  - mkl_intel_s.lib for static linking
  - mkl_intel_s_dll.lib for dynamic linking

**Caution** Avoid linking with the Single Dynamic Library mkl_rt.dll because its support of the stdcall calling convention is limited.

See Also

Using the cdecl and stdcall Interfaces
Compiling an Application that Calls Intel® MKL and Uses the CVF Calling Conventions
Intel® MKL Include Files

Compiling an Application that Calls the Intel® Math Kernel Library and Uses the CVF Calling Conventions

The IA-32 architecture implementation of Intel® MKL supports the Compaq Visual Fortran* (CVF) calling convention by providing the stdcall interface.

Although the Intel® MKL does not provide the CVF interface in its Intel® 64 architecture implementation, you can use the Intel® Visual Fortran Compiler to compile your Intel® 64 architecture application that calls Intel® MKL and uses the CVF calling convention. To do this:

- Provide the following compiler options to enable compatibility with the CVF calling convention:
  ```c
  /Gm or /iface:cvf
  ```
Additionally provide the following options to enable calling Intel® MKL from your application:
    /iface:nomixed_str_len_arg

See Also
Using the cdecl and stdcall Interfaces
Compiler Support

Mixed-language Programming with the Intel Math Kernel Library

Appendix A Intel® Math Kernel Library Language Interfaces Support lists the programming languages supported for each Intel® MKL function domain. However, you can call Intel® MKL routines from different language environments.

See also these Knowledge Base articles:


Calling LAPACK, BLAS, and CBLAS Routines from C/C++ Language Environments

Not all Intel® MKL function domains support both C and Fortran environments. To use Intel® MKL Fortran-style functions in C/C++ environments, you should observe certain conventions, which are discussed for LAPACK and BLAS in the subsections below.

Caution
Avoid calling BLAS 95/LAPACK 95 from C/C++. Such calls require skills in manipulating the descriptor of a deferred-shape array, which is the Fortran 90 type. Moreover, BLAS95/LAPACK95 routines contain links to a Fortran RTL.

LAPACK and BLAS

Because LAPACK and BLAS routines are Fortran-style, when calling them from C-language programs, follow the Fortran-style calling conventions:

• Pass variables by address, not by value.
  Function calls in Example "Calling a Complex BLAS Level 1 Function from C++" and Example "Using CBLAS Interface Instead of Calling BLAS Directly from C" illustrate this.
• Store your data in Fortran style, that is, column-major rather than row-major order.

With row-major order, adopted in C, the last array index changes most quickly and the first one changes most slowly when traversing the memory segment where the array is stored. With Fortran-style column-major order, the last index changes most slowly whereas the first index changes most quickly (as illustrated by the figure below for a two-dimensional array).
For example, if a two-dimensional matrix $A$ of size $m \times n$ is stored densely in a one-dimensional array $B$, you can access a matrix element like this:

$$A[i][j] = B[i*n+j] \text{ in C} \quad (i=0, \ldots , m-1, j=0, \ldots , n-1)$$

$$A(i,j) = B((j-1)*m+i) \text{ in Fortran} \quad (i=1, \ldots , m, j=1, \ldots , n).$$

When calling LAPACK or BLAS routines from C, be aware that because the Fortran language is case-insensitive, the routine names can be both upper-case or lower-case, with or without the trailing underscore. For example, the following names are equivalent:

- LAPACK: dgetrf, DGETRF, dgetrf_, and DGETRF_
- BLAS: dgemm, DGEMM, dgemm_, and DGEMM_

See Example "Calling a Complex BLAS Level 1 Function from C++" on how to call BLAS routines from C. See also the Intel® MKL Developer Reference for a description of the C interface to LAPACK functions.

**CBLAS**

Instead of calling BLAS routines from a C-language program, you can use the CBLAS interface.

CBLAS is a C-style interface to the BLAS routines. You can call CBLAS routines using regular C-style calls. Use the mkl.h header file with the CBLAS interface. The header file specifies enumerated values and prototypes of all the functions. It also determines whether the program is being compiled with a C++ compiler, and if it is, the included file will be correct for use with C++ compilation. Example "Using CBLAS Interface Instead of Calling BLAS Directly from C" illustrates the use of the CBLAS interface.

**C Interface to LAPACK**

Instead of calling LAPACK routines from a C-language program, you can use the C interface to LAPACK provided by Intel® MKL.

The C interface to LAPACK is a C-style interface to the LAPACK routines. This interface supports matrices in row-major and column-major order, which you can define in the first function argument matrix_order. Use the mkl.h header file with the C interface to LAPACK. mkl.h includes the mkl_lapacke.h header file, which specifies constants and prototypes of all the functions. It also determines whether the program is being compiled with a C++ compiler, and if it is, the included file will be correct for use with C++ compilation. You can find examples of the C interface to LAPACK in the examples\lapacke subdirectory in the Intel® MKL installation directory.

**Using Complex Types in C/C++**

As described in the documentation for the Intel® Visual Fortran Compiler, C/C++ does not directly implement the Fortran types COMPLEX(4) and COMPLEX(8). However, you can write equivalent structures. The type COMPLEX(4) consists of two 4-byte floating-point numbers. The first of them is the real-number component, and the second one is the imaginary-number component. The type COMPLEX(8) is similar to COMPLEX(4) except that it contains two 8-byte floating-point numbers.
Intel® MKL provides complex types MKL_Complex8 and MKL_Complex16, which are structures equivalent to the Fortran complex types COMPLEX(4) and COMPLEX(8), respectively. The MKL_Complex8 and MKL_Complex16 types are defined in the mkl_types.h header file. You can use these types to define complex data. You can also redefine the types with your own types before including the mkl_types.h header file. The only requirement is that the types must be compatible with the Fortran complex layout, that is, the complex type must be a pair of real numbers for the values of real and imaginary parts.

For example, you can use the following definitions in your C++ code:

```c++
#define MKL_Complex8 std::complex<float>

and

#define MKL_Complex16 std::complex<double>
```

See Example "Calling a Complex BLAS Level 1 Function from C++" for details. You can also define these types in the command line:

```bash
-DMKL_Complex8="std::complex<float>"
```

See Also

Intel® Software Documentation Library for the Intel® Visual Fortran Compiler documentation

for the Intel® Visual Fortran Compiler documentation

Calling BLAS Functions that Return the Complex Values in C/C++ Code

Complex values that functions return are handled differently in C and Fortran. Because BLAS is Fortran-style, you need to be careful when handling a call from C to a BLAS function that returns complex values. However, in addition to normal function calls, Fortran enables calling functions as though they were subroutines, which provides a mechanism for returning the complex value correctly when the function is called from a C program. When a Fortran function is called as a subroutine, the return value is the first parameter in the calling sequence. You can use this feature to call a BLAS function from C.

The following example shows how a call to a Fortran function as a subroutine converts to a call from C and the hidden parameter result gets exposed:

Normal Fortran function call:

```fortran
result = cdotc( n, x, 1, y, 1 )
```

A call to the function as a subroutine:

```fortran
call cdotc( result, n, x, 1, y, 1)
```

A call to the function from C:

```c
cdotc( &result, &n, x, &one, y, &one )
```

NOTE

Intel® MKL has both upper-case and lower-case entry points in the Fortran-style (case-insensitive) BLAS, with or without the trailing underscore. So, all these names are equivalent and acceptable: cdotc, CDOTC, cdotc_, and CDOTC_.

The above example shows one of the ways to call several level 1 BLAS functions that return complex values from your C and C++ applications. An easier way is to use the CBLAS interface. For instance, you can call the same function using the CBLAS interface as follows:

```c
clasm_cdotc( n, x, 1, y, 1, &result )
```

NOTE

The complex value comes last on the argument list in this case.

The following examples show use of the Fortran-style BLAS interface from C and C++, as well as the CBLAS (C language) interface:
• Example "Calling a Complex BLAS Level 1 Function from C"
• Example "Calling a Complex BLAS Level 1 Function from C++"
• Example "Using CBLAS Interface Instead of Calling BLAS Directly from C"

Example "Calling a Complex BLAS Level 1 Function from C"

The example below illustrates a call from a C program to the complex BLAS Level 1 function `zdotc()`. This function computes the dot product of two double-precision complex vectors.

In this example, the complex dot product is returned in the structure `c`.

```c
#include "mkl.h"
define N 5
int main()
{
    int n = N, inca = 1, incb = 1, i;
    MKL_Complex16 a[N], b[N], c;
    for( i = 0; i < n; i++ )
    {
        a[i].real = (double)i; a[i].imag = (double)i * 2.0;
        b[i].real = (double)(n - i); b[i].imag = (double)i * 2.0;
    }
    zdotc( &c, &n, a, &inca, b, &incb );
    printf( "The complex dot product is: ( %6.2f, %6.2f)\n", c.real, c.imag );
    return 0;
}
```

Example "Calling a Complex BLAS Level 1 Function from C++"

Below is the C++ implementation:

```c++
#include <complex>
#include <iostream>
define MKL_Complex16 std::complex<double>
#include "mkl.h"
define N 5
int main()
{
    int n, inca = 1, incb = 1, i;
    std::complex<double> a[N], b[N], c;
    n = N;
    for( i = 0; i < n; i++ )
    {
        a[i] = std::complex<double>(i,i*2.0);
        b[i] = std::complex<double>(n-i,i*2.0);
    }
    zdotc(&c, &n, a, &inca, b, &incb );
    std::cout << "The complex dot product is: " << c << std::endl;
    return 0;
}
```
Example "Using CBLAS Interface Instead of Calling BLAS Directly from C"

This example uses CBLAS:

```c
#include <stdio.h>
#include "mkl.h"

typedef struct{ double re; double im; } complex16;
#define N 5

int main()
{
    int n, inca = 1, incb = 1, i;
    complex16 a[N], b[N], c;
    n = N;
    for( i = 0; i < n; i++ )
    {
        a[i].re = (double)i; a[i].im = (double)i * 2.0;
        b[i].re = (double)(n - i); b[i].im = (double)i * 2.0;
    }
    cblas_zdotc_sub(n, a, inca, b, incb, &c);
    printf( "The complex dot product is: ( %6.2f, %6.2f)\n", c.re, c.im );
    return 0;
}
```
Intel® Math Kernel Library (Intel® MKL) offers functions and environment variables that help you obtain Conditional Numerical Reproducibility (CNR) of floating-point results when calling the library functions from your application. These new controls enable Intel® MKL to run in a special mode, when functions return bitwise reproducible floating-point results from run to run under the following conditions:

- Calls to Intel® MKL occur in a single executable
- The number of computational threads used by the library does not change in the run

For a limited set of routines, you can eliminate the second condition by using Intel® MKL in strict CNR mode.

It is well known that for general single and double precision IEEE floating-point numbers, the associative property does not always hold, meaning \((a+b)+c\) may not equal \(a+(b+c)\). Let’s consider a specific example. In infinite precision arithmetic \(2^{-63} + 1 + -1 = 2^{-63}\). If this same computation is done on a computer using double precision floating-point numbers, a rounding error is introduced, and the order of operations becomes important:

\begin{align*}
(2^{-63} + 1) + (-1) &\approx 1 + (-1) = 0 \\
2^{-63} + (1 + (-1)) &\approx 2^{-63} + 0 = 2^{-63}
\end{align*}

This inconsistency in results due to order of operations is precisely what the new functionality addresses.

The application related factors that affect the order of floating-point operations within a single executable program include selection of a code path based on run-time processor dispatching, alignment of data arrays, variation in number of threads, threaded algorithms and internal floating-point control settings. You can control most of these factors by controlling the number of threads and floating-point settings and by taking steps to align memory when it is allocated (see the Getting Reproducible Results with Intel® MKL knowledge base article for details). However, run-time dispatching and certain threaded algorithms do not allow users to make changes that can ensure the same order of operations from run to run.

Intel® MKL does run-time processor dispatching in order to identify the appropriate internal code paths to traverse for the Intel® MKL functions called by the application. The code paths chosen may differ across a wide range of Intel processors and Intel architecture compatible processors and may provide differing levels of performance. For example, an Intel® MKL function running on an Intel® Pentium® 4 processor may run one code path, while on the latest Intel® Xeon® processor it will run another code path. This happens because each unique code path has been optimized to match the features available on the underlying processor. One key way that the new features of a processor are exposed to the programmer is through the instruction set architecture (ISA). Because of this, code branches in Intel® MKL are designated by the latest ISA they use for optimizations: from the Intel® Streaming SIMD Extensions 2 (Intel® SSE2) to the Intel® Advanced Vector Extensions2 (Intel® AVX2). The feature-based approach introduces a challenge: if any of the internal floating-point operations are done in a different order or are re-associated, the computed results may differ.

Dispatching optimized code paths based on the capabilities of the processor on which the code is running is central to the optimization approach used by Intel® MKL. So it is natural that consistent results require some performance trade-offs. If limited to a particular code path, performance of Intel® MKL can in some circumstances degrade by more than a half. To understand this, note that matrix-multiply performance nearly doubled with the introduction of new processors supporting Intel AVX2 instructions. Even if the code branch is not restricted, performance can degrade by 10-20% because the new functionality restricts algorithms to maintain the order of operations.

**Optimization Notice**

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-
Optimization Notice

dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

Getting Started with Conditional Numerical Reproducibility

Intel® MKL offers functions and environment variables to help you get reproducible results. You can configure Intel® MKL using functions or environment variables, but the functions provide more flexibility.

The following specific examples introduce you to the conditional numerical reproducibility.

While these examples recommend aligning input and output data, you can supply unaligned data to Intel® MKL functions running in the CNR mode, but refer to Reproducibility Conditions for details related to data alignment.

Intel CPUs supporting Intel AVX2

To ensure Intel® MKL calls return the same results on every Intel CPU supporting Intel AVX2 instructions:

1. Make sure that your application uses a fixed number of threads
2. (Recommended) Properly align input and output arrays in Intel® MKL function calls
3. Do either of the following:
   - Call
     `mkl_cbwr_set(MKL_CBWR_AVX2)`
   - Set the environment variable:
     `set MKL_CBWR = AVX2`

NOTE
On non-Intel CPUs and on Intel CPUs that do not support Intel AVX2, this environment setting may cause results to differ because the AUTO branch is used instead, while the above function call returns an error and does not enable the CNR mode.

Intel CPUs supporting Intel SSE2

To ensure Intel® MKL calls return the same results on every Intel CPU supporting Intel SSE2 instructions:

1. Make sure that your application uses a fixed number of threads
2. (Recommended) Properly align input and output arrays in Intel® MKL function calls
3. Do either of the following:
   - Call
     `mkl_cbwr_set(MKL_CBWR_SSE2)`
   - Set the environment variable:
     `set MKL_CBWR = SSE2`
NOTE
On non-Intel CPUs, this environment setting may cause results to differ because the AUTO branch is used instead, while the above function call returns an error and does not enable the CNR mode.

Intel or Intel compatible CPUs supporting Intel SSE2
On non-Intel CPUs, only the MKL_CBWR_AUTO and MKL_CBWR_COMPATIBLE options are supported for function calls and only AUTO and COMPATIBLE options for environment settings.

To ensure Intel® MKL calls return the same results on all Intel or Intel compatible CPUs supporting Intel SSE2 instructions:
1. Make sure that your application uses a fixed number of threads
2. (Recommended) Properly align input and output arrays in Intel® MKL function calls
3. Do either of the following:
   - Call
     
     mkl_cbwr_set(MKL_CBWR_COMPATIBLE)
   - Set the environment variable:
     
     set MKL_CBWR = COMPATIBLE

NOTE
The special MKL_CBWR_COMPATIBLE/COMPATIBLE option is provided because Intel and Intel compatible CPUs have a few instructions, such as approximation instructions rcpps/rsqrtps, that may return different results. This option ensures that Intel® MKL does not use these instructions and forces a single Intel SSE2 only code path to be executed.

Next steps
See Specifying the Code Branches for details of specifying the branch using environment variables.

See the following sections in the Intel® MKL Developer Reference:
Support Functions for Conditional Numerical Reproducibility for how to configure the CNR mode of Intel® MKL using functions.
Intel® MKL PARDISO - Parallel Direct Sparse Solver Interface for how to configure the CNR mode for PARDISO.

See Also
Code Examples

Specifying Code Branches
Intel® MKL provides a conditional numerical reproducibility (CNR) functionality that enables you to obtain reproducible results from MKL routines. When enabling CNR, you choose a specific code branch of Intel® MKL that corresponds to the instruction set architecture (ISA) that you target. You can specify the code branch and other CNR options using the MKL_CBWR environment variable.

- MKL_CBWR="<branch>,[STRICT]"
- MKL_CBWR="BRANCH=<branch>,[STRICT]"

Use the STRICT flag to enable strict CNR mode. For more information, see Reproducibility Conditions.
The `<branch>` placeholder specifies the CNR branch with one of the following values:

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUTO</td>
<td>CNR mode uses the standard ISA-based dispatching model while ensuring fixed cache sizes, deterministic reductions, and static scheduling</td>
</tr>
<tr>
<td>COMPATIBLE</td>
<td>Intel® Streaming SIMD Extensions 2 (Intel® SSE2) without rcpps/rsqrtps instructions</td>
</tr>
<tr>
<td>SSE2</td>
<td>Intel SSE2</td>
</tr>
<tr>
<td>SSE3</td>
<td>DEPRECATED. Intel® Streaming SIMD Extensions 3 (Intel® SSE3). This setting is kept for backward compatibility and is equivalent to SSE2.</td>
</tr>
<tr>
<td>SSSE3</td>
<td>Supplemental Streaming SIMD Extensions 3 (SSSE3)</td>
</tr>
<tr>
<td>SSE4_2</td>
<td>Intel® Streaming SIMD Extensions 4.2 (Intel® SSE4.2)</td>
</tr>
<tr>
<td>AVX</td>
<td>Intel® Advanced Vector Extensions (Intel® AVX)</td>
</tr>
<tr>
<td>AVX512</td>
<td>Intel AVX-512 on Intel® Xeon® processors</td>
</tr>
<tr>
<td>AVX512_E1</td>
<td>Intel® Advanced Vector Extensions 512 (Intel® AVX-512) with support for Vector Neural Network Instructions</td>
</tr>
<tr>
<td>AVX512_MIC</td>
<td>Intel® Advanced Vector Extensions 512 (Intel® AVX-512) on Intel® Xeon Phi™ processors</td>
</tr>
<tr>
<td>AVX512_MIC_E1</td>
<td>Intel® Advanced Vector Extensions 512 (Intel® AVX-512) with support for Vector Neural Network Instructions on Intel® Xeon Phi™ processors</td>
</tr>
</tbody>
</table>

When specifying the CNR branch, be aware of the following:

- Reproducible results are provided under Reproducibility Conditions.
- Settings other than AUTO or COMPATIBLE are available only for Intel processors.
- To get the CNR branch optimized for the processor where your program is currently running, choose the value of AUTO or call the mkl_cbwr_get_auto_branch function.
- Strict CNR mode is only supported for AVX2, AVX512, AVX512_E1, AVX512_MIC, and AVX512_MIC_E1 branches. You can also use strict CNR mode with the AUTO branch when running on Intel processors that support one of these instruction set architectures (ISAs).

Setting the MKL_CBWR environment variable or a call to an equivalent mkl_cbwr_set function fixes the code branch and sets the reproducibility mode.

**NOTE**

- If the value of the branch is incorrect or your processor or operating system does not support the specified ISA, CNR ignores this value and uses the AUTO branch without providing any warning messages.
- Calls to functions that define the behavior of CNR must precede any of the math library functions that they control.
- Settings specified by the functions take precedence over the settings specified by the environment variable.

See the Intel® MKL Developer Reference for how to specify the branches using functions.

**See Also**

Getting Started with Conditional Numerical Reproducibility
Reproducibility Conditions

To get reproducible results from run to run, ensure that the number of threads is fixed and constant. Specifically:

- If you are running your program with OpenMP* parallelization on different processors, explicitly specify the number of threads.
- To ensure that your application has deterministic behavior with OpenMP* parallelization and does not adjust the number of threads dynamically at run time, set MKL_DYNAMIC and OMP_DYNAMIC to FALSE. This is especially needed if you are running your program on different systems.
- If you are running your program with the Intel® Threading Building Blocks parallelization, numerical reproducibility is not guaranteed.

Strict CNR Mode

In strict CNR mode, Intel® MKL provides bitwise reproducible results for a limited set of functions and code branches even when the number of threads changes. These routines and branches support strict CNR mode (64-bit libraries only):

- Intel® Advanced Vector Extensions 2 (Intel® AVX2) or Intel® Advanced Vector Extensions 512 (Intel® AVX-512).

When using other routines or CNR branches, Intel® MKL operates in standard (non-strict) CNR mode, subject to the restrictions described above. Enabling strict CNR mode can reduce performance.

**NOTE**

- As usual, you should align your data, even in CNR mode, to obtain the best possible performance. While CNR mode also fully supports unaligned input and output data, the use of it might reduce the performance of some Intel® MKL functions on earlier Intel processors. Refer to coding techniques that improve performance for more details.
- Conditional Numerical Reproducibility does not ensure that bitwise-identical NaN values are generated when the input data contains NaN values.
- If dynamic memory allocation fails on one run but succeeds on another run, you may fail to get reproducible results between these two runs.

See Also
MKL_DYNAMIC
Coding Techniques

Setting the Environment Variable for Conditional Numerical Reproducibility

The following examples illustrate the use of the MKL_CBWR environment variable. The first command sets Intel® MKL to run in the CNR mode based on the default dispatching for your platform. The other two commands are equivalent and set the CNR branch to Intel AVX:

- `set MKL_CBWR=AUTO`
- `set MKL_CBWR=AVX`
- `set MKL_CBWR=BRANCH=AVX`

See Also
Specifying Code Branches
Code Examples

The following simple programs show how to obtain reproducible results from run to run of Intel® MKL functions. See the Intel® MKL Developer Reference for more examples.

C Example of CNR

```c
#include <mkl.h>
int main(void) {
    int my_cbwr_branch;
    /* Align all input/output data on 64-byte boundaries */
    /* "for best performance of Intel® MKL "*/
    void *darray;
    int darray_size=1000;
    /* Set alignment value in bytes */
    int alignment=64;
    /* Allocate aligned array */
    darray = mkl_malloc (sizeof(double)*darray_size, alignment);
    /* Find the available MKL_CBWR_BRANCH automatically */
    my_cbwr_branch = mkl_cbwr_get_auto_branch();
    /* User code without Intel MKL calls */
    /* Piece of the code where CNR of Intel MKL is needed */
    /* The performance of Intel MKL functions might be reduced for CNR mode */
    /* If the "IF" statement below is commented out, Intel® MKL will run in a regular mode, */
    /* and data alignment will allow you to get best performance */
    if (mkl_cbwr_set(my_cbwr_branch)) {
        printf("Error in setting MKL_CBWR_BRANCH! Aborting\n");
        return;
    }
    /* CNR calls to Intel MKL + any other code */
    /* Free the allocated aligned array */
    mkl_free(darray);
}
```

Fortran Example of CNR

```fortran
PROGRAM MAIN
    INCLUDE 'mkl.fi'
    INTEGER*4 MY_CBWR_BRANCH
    ! Align all input/output data on 64-byte boundaries
    ! "for best performance of Intel® MKL
    ! Declare Intel MKL memory allocation routine
    ifndef _IA32
        INTEGER MKL_MALLOC
    else
        INTEGER*8 MKL_MALLOC
    endif
    EXTERNAL MKL_MALLOC, MKL_FREE
    DOUBLE PRECISION DARRAY
    POINTER (P_DARRAY,DARRAY(1))
    INTEGER DARRAY_SIZE
    PARAMETER (DARRAY_SIZE=1000)
    ! Set alignment value in bytes
    INTEGER ALIGNMENT
    PARAMETER (ALIGNMENT=64)
    ! Allocate aligned array
    P_DARRAY = MKL_MALLOC (%VAL(8*DARRAY_SIZE), %VAL(ALIGNMENT));
    ! Find the available MKL_CBWR_BRANCH automatically
```
MY_CBWR_BRANCH = MKL_CBWR_GET_AUTO_BRANCH()
! User code without Intel MKL calls
! Piece of the code where CNR of Intel MKL is needed
! The performance of Intel MKL functions may be reduced for CNR mode
! If the "IF" statement below is commented out, Intel® MKL will run in a regular mode,
! and data alignment will allow you to get best performance
   IF (MKL_CBWR_SET (MY_CBWR_BRANCH) .NE. MKL_CBWR_SUCCESS) THEN
      PRINT *, 'Error in setting MKL_CBWR_BRANCH! Aborting..
      RETURN
   ENDIF
! CNR calls to Intel MKL + any other code
! Free the allocated aligned array
   CALL MKL_FREE(P_DARRAY)
END

Use of CNR with Unaligned Data in C

#include <mkl.h>
int main(void) {
   int my_cbwr_branch;
   /* If it is not possible to align all input/output data on 64-byte boundaries */
   /* to achieve performance, use unaligned IO data with possible performance */
   /* penalty */
   /* Using unaligned IO data */
   double *darray;
   int darray_size=1000;
   /* Allocate array, malloc aligns data on 8/16-byte boundary only */
   darray = (double *)malloc (sizeof(double)*darray_size);
   /* Find the available MKL_CBWR_BRANCH automatically */
   my_cbwr_branch = mkl_cbwr_get_auto_branch();
   /* User code without Intel MKL calls */
   /* Piece of the code where CNR of Intel MKL is needed */
   /* The performance of Intel MKL functions might be reduced for CNR mode */
   /* If the "IF" statement below is commented out, Intel MKL will run in a regular mode, */
   /* and you will NOT get best performance without data alignment */
   if (mkl_cbwr_set(my_cbwr_branch)) {
      printf("Error in setting MKL_CBWR_BRANCH! Aborting..
      return;
   }
   /* CNR calls to Intel MKL + any other code */
   /* Free the allocated array */
   free(darray);
}

Use of CNR with Unaligned Data in Fortran

PROGRAM MAIN
   INCLUDE 'mkl.fi'
   INTEGER*4 MY_CBWR_BRANCH
   ! If it is not possible to align all input/output data on 64-byte boundaries
   ! to achieve performance, use unaligned IO data with possible performance
   ! penalty
   DOUBLE PRECISION, DIMENSION(:), ALLOCATABLE :: DARRAY
   INTEGER DARRAY_SIZE, STATUS
   PARAMETER (DARRAY_SIZE=1000)
   ! Allocate array with undefined alignment
   ALLOCATE(DARRAY(DARRAY_SIZE));
   ! Find the available MKL_CBWR_BRANCH automatically
   ""
MY_CBWR_BRANCH = MKL_CBWR_GET_AUTO_BRANCH()
! User code without Intel MKL calls
! Piece of the code where CNR of Intel MKL is needed
! The performance of Intel MKL functions might be reduced for CNR mode
! If the "IF" statement below is commented out, Intel MKL will run in a regular mode,
! and you will NOT get best performance without data alignment
   IF (MKL_CBWR_SET(MY_CBWR_BRANCH) .NE. MKL_CBWR_SUCCESS) THEN
      PRINT *, 'Error in setting MKL_CBWR_BRANCH! Aborting…'
      RETURN
   ENDIF
! CNR calls to Intel MKL + any other code
! Free the allocated array
   DEALLOCATE(DARRAY)
END
Coding Tips

This section provides coding tips for managing data alignment and version-specific compilation.

See Also
Mixed-language Programming with the Intel® Math Kernel Library  Tips on language-specific programming
Managing Performance and Memory  Coding tips related to performance improvement and use of memory functions
Obtaining Numerically Reproducible Results  Tips for obtaining numerically reproducible results of computations

Example of Data Alignment

Needs for best performance with Intel® MKL or for reproducible results from run to run of Intel® MKL functions require alignment of data arrays. The following example shows how to align an array on 64-byte boundaries. To do this, use `mkl_malloc()` in place of system provided memory allocators, as shown in the code example below.

Aligning Addresses on 64-byte Boundaries

```c
// ******* C language *******
...  
#include <stdlib.h>
#include <mkl.h>
...  
void *darray;
int workspace;
// Set value of alignment
int alignment=64;
...  
// Allocate aligned workspace
darray = mkl_malloc( sizeof(double)*workspace, alignment );
...
// call the program using Intel MKL
mkl_app( darray );
...  
// Free workspace
mkl_free( darray );
```
integer*8 mkl_malloc
endif
external mkl_malloc, mkl_free, mkl_app
...
double precision darray
pointer (p_wrk,darray(l))
integer workspace
...
! Allocate aligned workspace
p_wrk = mkl_malloc( %val(8*workspace), %val(alignment) )
...
! call the program using Intel MKL
call mkl_app( darray )
...
! Free workspace
call mkl_free(p_wrk)

Using Predefined Preprocessor Symbols for Intel® MKL Version-Dependent Compilation

Preprocessor symbols (macros) substitute values in a program before it is compiled. The substitution is performed in the preprocessing phase.

The following preprocessor symbols are available:

<table>
<thead>
<tr>
<th>Predefined Preprocessor Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INTEL_MKL</strong></td>
<td>Intel® MKL major version</td>
</tr>
<tr>
<td><strong>INTEL_MKL_MINOR</strong></td>
<td>Intel® MKL minor version</td>
</tr>
<tr>
<td><strong>INTEL_MKL_UPDATE</strong></td>
<td>Intel® MKL update number</td>
</tr>
<tr>
<td>INTEL_MKL_VERSION</td>
<td>Intel® MKL full version in the following format:</td>
</tr>
</tbody>
</table>

```
INTEL_MKL_VERSION =
(_INTEL_MKL__*100+__INTEL_MKL_MINOR__)*100+__INTEL_MKL_UPDATE__
```

These symbols enable conditional compilation of code that uses new features introduced in a particular version of the library.

To perform conditional compilation:

1. Depending on your compiler, include in your code the file where the macros are defined:
   - C/C++ compiler: `mkl_version.h`, or `mkl.h`, which includes `mkl_version.h`
   - Intel® Fortran compiler: `mkl.fi`
   - Any Fortran compiler with enabled preprocessing: `mkl_version.h`
   Read the documentation for your compiler for the option that enables preprocessing.

2. [Optionally] Use the following preprocessor directives to check whether the macro is defined:
   - `#ifdef, #endif` for C/C++
   - `!DEC$IF DEFINED, !DEC$ENDIF` for Fortran

3. Use preprocessor directives for conditional inclusion of code:
• #if, #endif for C/C++
• !DEC$IF, !DEC$ENDIF for Fortran

Example
This example shows how to compile a code segment conditionally for a specific version of Intel® MKL. In this case, the version is 11.2 Update 4:

Intel® Fortran Compiler:

```fortran
#include "mkl.fi"
!DEC$IF DEFINED INTEL_MKL_VERSION
!DEC$IF INTEL_MKL_VERSION .EQ. 110204
*     Code to be conditionally compiled
!DEC$ENDIF
!DEC$ENDIF
```

C/C++ Compiler. Fortran Compiler with Enabled Preprocessing:

```c
#include "mkl.h"
#ifdef INTEL_MKL_VERSION
#if INTEL_MKL_VERSION == 110204
...     Code to be conditionally compiled
#else
...     Code to be conditionally compiled
#endif
```

```c
#endif
```
Managing Output

Using Intel® MKL Verbose Mode

When building applications that call Intel® MKL functions, it may be useful to determine:

- which computational functions are called,
- what parameters are passed to them, and
- how much time is spent to execute the functions.

You can get an application to print this information to a standard output device by enabling Intel® MKL Verbose. Functions that can print this information are referred to as verbose-enabled functions.

When Verbose mode is active in an Intel® MKL domain, every call of a verbose-enabled function finishes with printing a human-readable line describing the call. However, if your application gets terminated for some reason during the function call, no information for that function will be printed. The first call to a verbose-enabled function also prints a version information line.

To enable the Intel® MKL Verbose mode for an application, do one of the following:

- set the environment variable MKL_VERBOSE to 1, or
- call the support function mkl_verbose(1).

To disable the Intel® MKL Verbose mode, call the mkl_verbose(0) function. Both enabling and disabling of the Verbose mode using the function call takes precedence over the environment setting. For a full description of the mkl_verbose function, see either the Intel® MKL Developer Reference for C or the Intel® MKL Developer Reference for Fortran. Both references are available in the Intel® Software Documentation Library.

You can enable Intel® MKL Verbose mode in these domains:

- BLAS
- LAPACK
- ScaLAPACK (selected functionality)
- FFT

Intel® MKL Verbose mode is not a thread-local but a global state. In other words, if an application changes the mode from multiple threads, the result is undefined.

**WARNING**

The performance of an application may degrade with the Verbose mode enabled, especially when the number of calls to verbose-enabled functions is large, because every call to a verbose-enabled function requires an output operation.

See Also

Intel Software Documentation Library

Version Information Line

In the Intel® MKL Verbose mode, the first call to a verbose-enabled function prints a version information line. The line begins with the MKL_VERBOSE character string and uses spaces as delimiters. The format of the rest of the line may change in a future release.

The following table lists information contained in a version information line and provides available links for more information:
<table>
<thead>
<tr>
<th>Information</th>
<th>Description</th>
<th>Related Links</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel® MKL version.</td>
<td>This information is separated by a comma from the rest of the line.</td>
<td></td>
</tr>
<tr>
<td>Operating system.</td>
<td>Possible values:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Ln​x for Linux® OS</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Win for Windows® OS</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• OSX for macOS*</td>
<td></td>
</tr>
<tr>
<td>The host CPU frequency.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intel® MKL interface layer used by the application.</td>
<td>Possible values:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• stdcall or cdecl</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• lp64 or ilp64 on systems based on the Intel® 64 architecture.</td>
<td></td>
</tr>
<tr>
<td>Intel® MKL threading layer used by the application.</td>
<td>Possible values:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>intel_thread, tbb_thread, pgi_thread, or sequential.</td>
<td></td>
</tr>
<tr>
<td>The following is an example of a version information line:</td>
<td>MKL_VERBOSE Intel(R) MKL 11.2 Beta build 20131126 for Intel(R) 64 architecture Intel(R) Advanced Vector Extensions (Intel(R) AVX) Enabled Processor, Win 3.10GHz lp64 intel_thread</td>
<td></td>
</tr>
<tr>
<td><strong>Call Description Line</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>In Intel® MKL Verbose mode, each verbose-enabled function called from your application prints a call description line. The line begins with the MKL_VERBOSE character string and uses spaces as delimiters. The format of the rest of the line may change in a future release.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>The following table lists information contained in a call description line and provides available links for more information:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Information</td>
<td>Description</td>
<td>Related Links</td>
</tr>
<tr>
<td>The name of the function.</td>
<td>Although the name printed may differ from the name used in the source code of the application (for example, the cblas_ prefix of CBLAS functions is not printed), you can easily recognize the function by the printed name.</td>
<td></td>
</tr>
<tr>
<td>Values of the arguments.</td>
<td>• The values are listed in the order of the formal argument list.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>The list directly follows the function name, it is parenthesized and comma-separated.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Arrays are printed as addresses (to see the alignment of the data).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Integer scalar parameters passed by reference are printed by value.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Zero values are printed for NULL references.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Character values are printed without quotes.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>For all parameters passed by reference, the values printed are the values returned by the function. For example, the printed value of the info parameter of a LAPACK function is its value after the function execution.</td>
<td></td>
</tr>
<tr>
<td>Information</td>
<td>Description</td>
<td>Related Links</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Time taken by the function.</td>
<td>• The time is printed in convenient units (seconds, milliseconds, and so on), which are explicitly indicated. &lt;br&gt;• The time may fluctuate from run to run. &lt;br&gt;• The time printed may occasionally be larger than the time actually taken by the function call, especially for small problem sizes and multi-socket machines. To reduce this effect, bind threads that call Intel® MKL to CPU cores by setting an affinity mask.</td>
<td>Managing Multi-core Performance for options to set an affinity mask.</td>
</tr>
<tr>
<td>Value of the MKL_CBWR environment variable.</td>
<td>The value printed is prefixed with CNR:</td>
<td>Getting Started with Conditional Numerical Reproducibility</td>
</tr>
<tr>
<td>Value of the MKL_DYNAMIC environment variable.</td>
<td>The value printed is prefixed with Dyn:</td>
<td>MKL_DYNAMIC</td>
</tr>
<tr>
<td>Status of the Intel® MKL memory manager.</td>
<td>The value printed is prefixed with FastMM:</td>
<td>Avoiding Memory Leaks in Intel® MKL for a description of the Intel® MKL memory manager</td>
</tr>
<tr>
<td>OpenMP* thread number of the calling thread.</td>
<td>The value printed is prefixed with TID:</td>
<td></td>
</tr>
<tr>
<td>Values of Intel® MKL environment variables defining the general and domain-specific numbers of threads, separated by a comma.</td>
<td>The first value printed is prefixed with NThr:</td>
<td>Intel® MKL-specific Environment Variables for Threading Control</td>
</tr>
</tbody>
</table>

The following is an example of a call description line:

```plaintext
MKL_VERBOSE DGEMM(n,n, 1000,1000,240,0x7ff708bb30,0x7ff28e92b000,240,0x7ff708bb38,0x7ff28e08d000,1000) 1.66ms CNR:OFF Dyn:1 FastMM:1 TID:0 NThr:16,FFT:2
```

The following information is not printed because of limitations of Intel® MKL Verbose mode:

- Input values of parameters passed by reference if the values were changed by the function.
  - For example, if a LAPACK function is called with a workspace query, that is, the value of the `lwork` parameter equals `-1` on input, the call description line prints the result of the query and not `-1`.
- Return values of functions.
  - For example, the value returned by the function `ilaenv` is not printed.
- Floating-point scalars passed by reference.
Working with the Intel® Math Kernel Library Cluster Software

Intel® Math Kernel Library (Intel® MKL) includes distributed memory function domains for use on clusters:

- ScaLAPACK
- Cluster Fourier Transform Functions (Cluster FFT)
- Parallel Direct Sparse Solvers for Clusters (Cluster Sparse Solver)

ScaLAPACK, Cluster FFT, and Cluster Sparse Solver are only provided for the Intel® 64 and Intel® Many Integrated Core architectures.

**Important**

ScaLAPACK, Cluster FFT, and Cluster Sparse Solver function domains are not installed by default. To use them, explicitly select the appropriate component during installation.

**Optimization Notice**

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

**See Also**

- Intel® Math Kernel Library Structure
- Managing Performance of the Cluster Fourier Transform Functions
- Intel® Distribution for LINPACK® Benchmark

**Message-Passing Interface Support**


To link applications with ScaLAPACK, Cluster FFT, or Cluster Sparse Solver, you need to configure your system depending on your message-passing interface (MPI) implementation as explained below.

If you are using MPICH2, do the following:

1. Add mpich2\include to the include path (assuming the default MPICH2 installation).
2. Add mpich2\lib to the library path.
3. Add mpi.lib to your link command.
4. Add fmpich2.lib to your Fortran link command.
5. Add cxx.lib to your Release target link command and cxxd.lib to your Debug target link command for C++ programs.

If you are using the Microsoft MPI, do the following:

1. Add Microsoft Compute Cluster Pack\include to the include path (assuming the default installation of the Microsoft MPI).
2. Add Microsoft Compute Cluster Pack\Lib\AMD64 to the library path.
3. Add msmpi.lib to your link command.

If you are using the Intel® MPI, do the following:

1. Add the following string to the include path: %ProgramFiles%\Intel\MPI\<ver>\intel64\include, where <ver> is the directory for a particular MPI version, for example, %ProgramFiles%\Intel\MPI \5.1\intel64\include.
2. Add the following string to the library path: %ProgramFiles%\Intel\MPI\<ver>\intel64\lib, for example, %ProgramFiles%\Intel\MPI\5.1\intel64\lib.
3. Add impi.lib and impicxx.lib to your link command.

Check the documentation that comes with your MPI implementation for implementation-specific details of linking.

Linking with Intel® MKL Cluster Software

The Intel® MKL ScaLAPACK, Cluster FFT, and Cluster Sparse Solver support MPI implementations identified in the Intel® MKL Release Notes.

To link with ScaLAPACK, Cluster FFT, and/or Cluster Sparse Solver, use the following commands:

```bash
set lib =<path to MKL libraries>;<path to MPI libraries>;%lib%
<linker> <files to link> [<MKL cluster library>] <BLACS><MKL core libraries><MPI libraries>
```

where the placeholders stand for paths and libraries as explained in the following table:

<table>
<thead>
<tr>
<th>&lt;path to MKL libraries&gt;</th>
<th>&lt;mkl directory&gt;\lib\intel64_win. If you performed the Scripts to Set Environment Variables Setting Environment Variables step of the Getting Started process, you do not need to add this directory to the lib environment variable.</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;path to MPI libraries&gt;</td>
<td>Typically the lib subdirectory in the MPI installation directory.</td>
</tr>
<tr>
<td>&lt;linker&gt;</td>
<td>One of icl, ifort, xilink.</td>
</tr>
<tr>
<td>&lt;MKL cluster library&gt;</td>
<td>One of libraries for ScaLAPACK or Cluster FFT listed in Appendix C: Directory Structure in Detail. For example, for the LP64 interface, it is mkl_scalapack_lp64.lib or mkl_cdft_core.lib. Cluster Sparse Solver does not require an additional computation library.</td>
</tr>
<tr>
<td>&lt;BLACS&gt;</td>
<td>The BLACS library corresponding to your , programming interface (LP64 or ILP64), and MPI version. These libraries are listed in Appendix C: Directory Structure in Detail. For example, for the LP64 interface, choose one of mkl_blacs_intelmpi_lp64.lib, mkl_blacs_mpich2_lp64.lib, or mkl_blacs_msmpi_lp64.lib in the case of static linking and mkl_blacs_lp64_dll.lib in the case of dynamic linking.</td>
</tr>
<tr>
<td>&lt;MKL core libraries&gt;</td>
<td>Intel® MKL libraries other than libraries with ScaLAPACK, Cluster FFT, or Cluster Sparse Solver.</td>
</tr>
</tbody>
</table>
Tip
Use the Using the Link-line Advisor to quickly choose the appropriate set of *<MKL cluster Library>, <BLACS>, and <MKL core libraries>.*

Intel MPI provides prepackaged scripts for its linkers to help you link using the respective linker. Therefore, if you are using Intel MPI, the best way to link is to use the following commands:

```bash
<path to Intel MPI binaries>\mpivars.bat
set lib = <path to MKL libraries>%lib%
<mpilinker><files to link> [<MKL cluster Library>]<BLACS><MKL core libraries>
```

where the placeholders that are not yet defined are explained in the following table:

<table>
<thead>
<tr>
<th>Placeholder</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;path to MPI binaries&gt;</td>
<td>By default, the bin subdirectory in the MPI installation directory.</td>
</tr>
<tr>
<td>&lt;MPI linker&gt;</td>
<td>mpiclon mpiifort</td>
</tr>
</tbody>
</table>

See Also
Linking Your Application with the Intel(R) Math Kernel Library
Examples of Linking for Clusters

Determining the Number of OpenMP* Threads

The OpenMP* run-time library responds to the environment variable OMP_NUM_THREADS. Intel® MKL also has other mechanisms to set the number of OpenMP threads, such as the MKL_NUM_THREADS or MKL_DOMAIN_NUM_THREADS environment variables (see Using Additional Threading Control).

Make sure that the relevant environment variables have the same and correct values on all the nodes. Intel® MKL does not set the default number of OpenMP threads to one, but depends on the OpenMP libraries used with the compiler to set the default number. For the threading layer based on the Intel compiler (mkl_intel_thread.lib), this value is the number of CPUs according to the OS.

Caution
Avoid over-prescribing the number of OpenMP threads, which may occur, for instance, when the number of MPI ranks per node and the number of OpenMP threads per node are both greater than one. The number of MPI ranks per node multiplied by the number of OpenMP threads per node should not exceed the number of hardware threads per node.

The OMP_NUM_THREADS environment variable is assumed in the discussion below.

Set OMP_NUM_THREADS so that the product of its value and the number of MPI ranks per node equals the number of real processors or cores of a node. If the Intel® Hyper-Threading Technology is enabled on the node, use only half number of the processors that are visible on Windows OS.

Important
For Cluster Sparse Solver, set the number of OpenMP threads to a number greater than one because the implementation of the solver only supports a multithreaded algorithm.

See Also
Setting Environment Variables on a Cluster
Using DLLs

All the needed DLLs must be visible on all the nodes at run time, and you should install Intel® Math Kernel Library (Intel® MKL) on each node of the cluster. You can use Remote Installation Services (RIS) provided by Microsoft to remotely install the library on each of the nodes that are part of your cluster. The best way to make the DLLs visible is to point to these libraries in the \PATH\ environment variable. See Setting Environment Variables on a Cluster on how to set the value of the PATH environment variable.

The ScaLAPACK DLLs in the <mkl directory>\redist\intel64_win\mkl directory use the MPI dispatching mechanism. MPI dispatching is based on the MKL_BLACS_MPI environment variable. The BLACS DLL uses MKL_BLACS_MPI for choosing the needed MPI libraries. The table below lists possible values of the variable.

<table>
<thead>
<tr>
<th>Value</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPICH2</td>
<td>Default value. MPICH2 for Windows* OS is used for message passing</td>
</tr>
<tr>
<td>INTELMPI</td>
<td>Intel MPI is used for message passing</td>
</tr>
<tr>
<td>MSMPI</td>
<td>Microsoft MPI is used for message passing</td>
</tr>
<tr>
<td>CUSTOM</td>
<td>Intel® MKL MPI wrappers built with a custom MPI are used for message passing</td>
</tr>
</tbody>
</table>

If you are using a non-default MPI, assign the same appropriate value to MKL_BLACS_MPI on all nodes.

See Also
Setting Environment Variables on a Cluster
Notational Conventions

Setting Environment Variables on a Cluster

By default, when you call the MPI launch command mpiexec, the entire launching node environment is passed to the MPI processes. However, if there are undefined variables or variables that are different from what is stored in your environment, you can use -env or -genv options with mpiexec. Each of these options take two arguments- the name and the value of the environment variable to be passed.

- genv NAME1 VALUE1 -genv NAME2 VALUE2
  - env NAME VALUE -genv

See these MPICH2 examples on how to set the value of OMP_NUM_THREADS explicitly:

mpiexec -genv OMP_NUM_THREADS 2 ....

mpiexec -n 1 -host first -env OMP_NUM_THREADS 2 test.exe : -n 2 -host second -env OMP_NUM_THREADS 3 test.exe ....

See these Intel MPI examples on how to set the value of MKL_BLACS_MPI explicitly:

mpiexec -genv MKL_BLACS_MPI INTELMPI ....

mpiexec -n 1 -host first -env MKL_BLACS_MPI INTELMPI test.exe : -n 1 -host second -env MKL_BLACS_MPI INTELMPI test.exe.

If you want to pass all environment variables by default and avoid passing these values explicitly, modify the user or system environment variables on each Windows node. From the Start menu, select Settings > Control Panel > System > Advanced > Environment Variables.

If you are using Microsoft MPI, the ways of setting environment variables described above are also applicable if the Microsoft Single Program Multiple Data (SPMD) process managers are running in a debug mode on all nodes of the cluster. However, the best way to set environment variables is by using the Job Scheduler with the Microsoft Management Console (MMC) and/or the Command Line Interface (CLI) to submit a job and pass environment variables. For more information about MMC and CLI, see the Microsoft Help and Support page at the Microsoft Web site (http://www.microsoft.com/).
Interaction with the Message-passing Interface

To improve performance of cluster applications, it is critical for Intel® MKL to use the optimal number of threads, as well as the correct thread affinity. Usually, the optimal number is the number of available cores per node divided by the number of MPI processes per node. You can set the number of threads using one of the available methods, described in Techniques to Set the Number of Threads.

If the number of threads is not set, Intel® MKL checks whether it runs under MPI provided by the Intel® MPI Library. If this is true, the following environment variables define Intel® MKL threading behavior:

- $I_MPI_THREAD_LEVEL$
- MKL_MPI_PPN
- $I_MPI_NUMBER_OF_MPI_PROCESSES_PER_NODE$
- $I_MPI_PIN_MAPPING$
-OMPI_COMM_WORLD_LOCAL_SIZE
- MPI_LOCALNRANKS

The threading behavior depends on the value of $I_MPI_THREAD_LEVEL$ as follows:

- 0 or undefined.
  Intel® MKL considers that thread support level of Intel MPI Library is $MPI_THREAD_SINGLE$ and defaults to sequential execution.
- 1, 2, or 3.
  This value determines Intel® MKL conclusion of the thread support level:
  - 1 - $MPI_THREAD_FUNNELED$
  - 2 - $MPI_THREAD_SERIALIZED$
  - 3 - $MPI_THREAD_MULTIPLE$

In all these cases, Intel® MKL determines the number of MPI processes per node using the other environment variables listed and defaults to the number of threads equal to the number of available cores per node divided by the number of MPI processes per node.

**Important**
Instead of relying on the discussed implicit settings, explicitly set the number of threads for Intel® MKL.

---

**Optimization Notice**

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

**See Also**
Managing Multi-core Performance

[Intel® Software Documentation Library](https://software.intel.com)  for more information on Intel MPI Library
Using a Custom Message-Passing Interface

While different message-passing interface (MPI) libraries are compatible at the application programming interface (API) level, they are often incompatible at the application binary interface (ABI) level. Therefore, Intel® MKL provides a set of prebuilt BLACS libraries that support certain MPI libraries, but this, however, does not enable use of Intel® MKL with other MPI libraries. To fill this gap, Intel® MKL also includes the MKL MPI wrapper, which provides an MPI-independent ABI to Intel® MKL. The adaptor is provided as source code. To use Intel® MKL with an MPI library that is not supported by default, you can use the adapter to build custom static or dynamic BLACS libraries and use them similarly to the prebuilt libraries.

Building a Custom BLACS Library

The MKL MPI wrapper is located in the `<mkl directory>`\interfaces\mklmpi directory.

To build a custom BLACS library, from the above directory run the `nmake` command.

For example: the command

```
nmake libintel64
```

builds a static custom BLACS library `mkl_blacs_custom_lp64.lib` using the default MPI compiler on your system. Look into the `<mkl directory>`\interfaces\mklmpi\makefile for targets and variables that define how to build the custom library. In particular, you can specify the compiler through the `MPICC` variable.

For more control over the building process, refer to the documentation available through the command `nmake help`

Using a Custom BLACS Library

In the case of static linking, use custom BLACS libraries exactly the same way as you use the prebuilt BLACS libraries, but pass the custom library to the linker. For example, instead of passing the `mkl_blacs_intelmpi_lp64.lib` static library, pass `mkl_blacs_custom_lp64.lib`.

To use a dynamic custom BLACS library:

1. Link your application the same way as when you use the prebuilt BLACS library.
2. Call the `mkl_set_mpi` support function or set the `MKL_BLACS_MPI` environment variable to one of the following values:
   - `CUSTOM`
     to load a custom library with the default name `mkl_blacs_custom_lp64.dll` or `mkl_blacs_custom_ilp64.dll`, depending on whether the BLACS interface linked against your application is LP64 or ILP64.
   - `<dll_name>`
     to load the specified BLACS DLL.

**NOTE**

Intel® MKL looks for the specified DLL either in the directory with Intel® MKL dynamic libraries or in the directory with the application executable.

For a description of the `mkl_set_mpi` function, see the *Intel® MKL Developer Reference*.

See Also

Linking with Intel® MKL Cluster Software
Examples of Linking for Clusters

This section provides examples of linking with ScALAPACK, Cluster FFT, and Cluster Sparse Solver.

Note that a binary linked with the Intel® MKL cluster function domains runs the same way as any other MPI application (refer to the documentation that comes with your MPI implementation).

For further linking examples, see the support website for Intel products at https://www.intel.com/software/products/support/.

See Also
Directory Structure in Detail

Examples for Linking a C Application

These examples illustrate linking of an application under the following conditions:

- Main module is in C.
- MPICH2 is installed in c:\mpich2x64.
- You are using the Intel® C++ Compiler.
- Intel® MKL functions use LP64 interfaces.

To link with ScALAPACK for a cluster of Intel® 64 architecture based systems, set the environment variable and use the link line as follows:

```bash
set lib=c:\mpich2x64\lib;\<mkl directory>\lib\intel64_win;%lib%
icl <user files to link> mkl_scalapack_lp64.lib mkl_blacs_mpich2_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib mpi.lib cxx.lib bufferoverflowu.lib
```

To link with Cluster FFT for a cluster of Intel® 64 architecture based systems, set the environment variable and use the link line as follows:

```bash
set lib=c:\mpich2x64\lib;\<mkl directory>\lib\intel64_win;%lib%
icl <user files to link> mkl_cdft_core.lib mkl_blacs_mpich2_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib mpi.lib cxx.lib bufferoverflowu.lib
```

To link with Cluster Sparse Solver for a cluster of Intel® 64 architecture based systems, set the environment variable and use the link line as follows:

```bash
set lib=c:\mpich2x64\lib;\<mkl directory>\lib\intel64_win;%lib%
icl <user files to link> mkl_blacs_mpich2_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib mpi.lib cxx.lib bufferoverflowu.lib
```

See Also
Linking with Intel® MKL Cluster Software
Using the Link-line Advisor
Linking with System Libraries

Examples for Linking a Fortran Application

These examples illustrate linking of an application under the following conditions:

- Main module is in Fortran.
- Microsoft Windows Compute Cluster Pack SDK is installed in c:\MS CCP SDK.
- You are using the Intel® Fortran Compiler.
• Intel® MKL functions use LP64 interfaces.

To link with ScaLAPACK for a cluster of Intel® 64 architecture based systems, set the environment variable and use the link line as follows:

```bash
set lib="c:\MS CCP SDK\Lib\AMD64";<mkl directory>\lib\intel64_win;%lib%
ifort <user files to link> mkl_scalapack_lp64.lib mkl_blacs_mpich2_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib msmpi.lib bufferoverflowu.lib
```

To link with Cluster FFT for a cluster of Intel® 64 architecture based systems, set the environment variable and use the link line as follows:

```bash
set lib="c:\MS CCP SDK\Lib\AMD64";<mkl directory>\lib\intel64_win;%lib%
ifort <user files to link> mkl_cdft_core.lib mkl_blacs_mpich2_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib msmpi.lib bufferoverflowu.lib
```

To link with Cluster Sparse Solver for a cluster of Intel® 64 architecture based systems, set the environment variable and use the link line as follows:

```bash
set lib="c:\MS CCP SDK\Lib\AMD64";<mkl directory>\lib\intel64_win;%lib%
ifort <user files to link> mkl_blacs_mpich2_lp64.lib mkl_intel_lp64.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib msmpi.lib bufferoverflowu.lib
```

**See Also**
- Linking with Intel® MKL Cluster Software
- Using the Link-line Advisor
- Linking with System Libraries
Managing Behavior of the Intel(R) Math Kernel Library with Environment Variables

See Also
Intel® MKL-specific Environment Variables for Threading Control

Specifying the Code Branches
for how to use an environment variable to specify the code branch for Conditional Numerical Reproducibility

Using Intel® MKL Verbose Mode
for how to use an environment variable to set the verbose mode

Managing Behavior of Function Domains with Environment Variables

Setting the Default Mode of Vector Math with an Environment Variable
Intel® Math Kernel Library (Intel® MKL) enables overriding the default setting of the Vector Mathematics (VM) global mode using the MKL_VML_MODE environment variable.

Because the mode is set or can be changed in different ways, their precedence determines the actual mode used. The settings and function calls that set or change the VM mode are listed below, with the precedence growing from lowest to highest:

1. The default setting
2. The MKL_VML_MODE environment variable
3. A call vmlSetMode function
4. A call to any VM function other than a service function

For more details, see the Vector Mathematical Functions section in the Intel® MKL Developer Reference and the description of the vmlSetMode function in particular.

To set the MKL_VML_MODE environment variable, use the following command:

```
set MKL_VML_MODE=<mode-string>
```

In this command, <mode-string> controls error handling behavior and computation accuracy, consists of one or several comma-separated values of the mode parameter listed in the table below, and meets these requirements:

- Not more than one accuracy control value is permitted
- Any combination of error control values except VML_ERRMODE_DEFAULT is permitted
- No denormalized numbers control values are permitted

<table>
<thead>
<tr>
<th>Value of mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy Control</td>
<td>high accuracy versions of VM functions</td>
</tr>
<tr>
<td>VML_HA</td>
<td>low accuracy versions of VM functions</td>
</tr>
<tr>
<td>VML_LA</td>
<td>enhanced performance accuracy versions of VM functions</td>
</tr>
<tr>
<td>Denormalized Numbers Handling Control</td>
<td></td>
</tr>
</tbody>
</table>

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### Managing Performance of the Cluster Fourier Transform Functions

Performance of Intel® MKL Cluster FFT (CFFT) in different applications mainly depends on the cluster configuration, performance of message-passing interface (MPI) communications, and configuration of the run. Note that MPI communications usually take approximately 70% of the overall CFFT compute time. For more flexibility of control over time-consuming aspects of CFFT algorithms, Intel® MKL provides the `MKL_CDFT` environment variable to set special values that affect CFFT performance. To improve performance of your application that intensively calls CFFT, you can use the environment variable to set optimal values for your cluster, application, MPI, and so on.

The `MKL_CDFT` environment variable has the following syntax, explained in the table below:

\[
\text{MKL_CDFT}=\text{option1}[=\text{value1}],\text{option2}[=\text{value2}],...,\text{optionN}[=\text{valueN}]
\]

### Important

While this table explains the settings that usually improve performance under certain conditions, the actual performance highly depends on the configuration of your cluster. Therefore, experiment with the listed values to speed up your computations.

<table>
<thead>
<tr>
<th>Option</th>
<th>Possible Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>alltoallv</td>
<td>0 (default)</td>
<td>Configures CFFT to use the standard MPI_Alltoallv function to perform global transpositions.</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>Configures CFFT to use a series of calls to MPI_Isend and MPI_Irecv instead of the MPI_Alltoallv function.</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>Configures CFFT to merge global transposition with data movements in the local memory. CFFT performs global transpositions by calling MPI_Isend and MPI_Irecv in this case.</td>
</tr>
</tbody>
</table>

### Managing Behavior of the Intel® Math Kernel Library with Environment Variables

This command provides an example of valid settings for the `MKL_VML_MODE` environment variable:

set MKL_VML_MODE=VML_LA,VML_ERRMODE_ERRNO,VML_ERRMODE_STDERR

**NOTE**

VM ignores the `MKL_VML_MODE` environment variable in the case of incorrect or misspelled settings of `mode`.

<table>
<thead>
<tr>
<th>Value of mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>VML_FTZDAZ_ON</td>
<td>Faster processing of denormalized inputs is enabled.</td>
</tr>
<tr>
<td>VML_FTZDAZ_OFF</td>
<td>Faster processing of denormalized inputs is disabled.</td>
</tr>
<tr>
<td>Error Mode Control</td>
<td></td>
</tr>
<tr>
<td>VML_ERRMODE_IGNORE</td>
<td>On computation error, VM Error status is updated, but otherwise no action is set. Cannot be combined with other VML_ERRMODE settings.</td>
</tr>
<tr>
<td>VML_ERRMODE_NOERR</td>
<td>On computation error, VM Error status is not updated and no action is set. Cannot be combined with other VML_ERRMODE settings.</td>
</tr>
<tr>
<td>VML_ERRMODE_STDERR</td>
<td>On error, the error text information is written to stderr.</td>
</tr>
<tr>
<td>VML_ERRMODE_EXCEPT</td>
<td>On error, an exception is raised.</td>
</tr>
<tr>
<td>VML_ERRMODE_CALLBACK</td>
<td>On error, an additional error handler function is called.</td>
</tr>
<tr>
<td>VML_ERRMODE_DEFAULT</td>
<td>On error, an exception is raised and an additional error handler function is called.</td>
</tr>
<tr>
<td>Option</td>
<td>Possible Values</td>
</tr>
<tr>
<td>--------------</td>
<td>----------------</td>
</tr>
<tr>
<td>wo_omatcopy</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>-1 (default)</td>
</tr>
<tr>
<td>enable_soi</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

The following example illustrates usage of the environment variable:

```
set MKL_CDFT=wo_omatcopy=1,alltoallv=4,enable_soi
mpirun -ppn 2 -n 16 mkl_cdft_app.exe
```

**Optimization Notice**

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

**Managing Invalid Input Checking in LAPACKE Functions**

The high-level interface includes an optional, on by default, NaN check on all matrix inputs before calling any LAPACK routine. This option affects all routines. If an input matrix contains any NaNs, the input parameter corresponding to this matrix is flagged with a return value error. For example, if the fifth parameter is found to contain a NaN, the routine returns the value, -5. The middle-level interface does not contain the NaN check.

NaN checking on matrix input can be expensive. By default, NaN checking is turned on. LAPACKE provides a way to set it through the environment variable:
• Setting environment variable `LAPACKE_NANCHECK` to 0 turns OFF NaN-checking
• Setting environment variable `LAPACKE_NANCHECK` to 1 turns ON NaN-checking

The other way is the call the `LAPACKE_set_nancheck` function; see the Developer Reference for C's LAPACK Auxiliary Routines section for more information.

Note that the NaN-checking flag value set by the call to `LAPACKE_set_nancheck` always has higher priority than the environment variable, `LAPACKE_NANCHECK`.

### Instruction Set Specific Dispatching on Intel® Architectures

Intel® MKL automatically queries and then dispatches the code path supported on your Intel® processor to the optimal instruction set architecture (ISA) by default. The `MKL_ENABLE_INSTRUCTIONS` environment variable or the `mkl_enable_instructions` support function enables you to dispatch to an ISA-specific code path of your choice. For example, you can run the Intel® Advanced Vector Extensions (Intel® AVX) code path on an Intel processor based on Intel® Advanced Vector Extensions 2 (Intel® AVX2), or you can run the Intel® Streaming SIMD Extensions 4.2 (Intel® SSE4.2) code path on an Intel AVX-enabled Intel processor. This feature is not available on non-Intel processors.

In some cases Intel® MKL also provides support for upcoming architectures ahead of hardware availability, but the library does not automatically dispatch the code path specific to an upcoming ISA by default. If for your exploratory work you need to enable an ISA for an Intel processor that is not yet released or if you are working in a simulated environment, you can use the `MKL_ENABLE_INSTRUCTIONS` environment variable or `mkl_enable_instructions` support function.

The following table lists possible values of `MKL_ENABLE_INSTRUCTIONS` alongside the corresponding ISA supported by a given processor. `MKL_ENABLE_INSTRUCTIONS` dispatches to the default ISA if the ISA requested is not supported on the particular Intel processor. For example, if you request to run the Intel AVX512 code path on a processor based on Intel AVX2, Intel® MKL runs the Intel AVX2 code path. The table also explains whether the ISA is dispatched by default on the processor that supports this ISA.

<table>
<thead>
<tr>
<th>Value of <code>MKL_ENABLE_INSTRUCTIONS</code></th>
<th>ISA</th>
<th>Dispatched by Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVX512</td>
<td>Intel® Advanced Vector Extensions (Intel® AVX-512) for systems based on Intel® Xeon® processors</td>
<td>Yes</td>
</tr>
<tr>
<td>AVX512_E1</td>
<td>Intel® Advanced Vector Extensions (Intel® AVX-512) with support for Vector Neural Network Instructions.</td>
<td>Yes</td>
</tr>
<tr>
<td>AVX512_E2</td>
<td>ICX: Intel® Advanced Vector Extensions (Intel® AVX-512) enabled processors.</td>
<td>Yes</td>
</tr>
<tr>
<td>AVX512_MIC</td>
<td>Intel® Advanced Vector Extensions (Intel® AVX-512) for systems based on Intel® Xeon Phi™ processors</td>
<td>Yes</td>
</tr>
<tr>
<td>AVX512_MIC_E1</td>
<td>Intel® Advanced Vector Extensions 512 (Intel® AVX-512) for Intel® Many Integrated Core Architecture (Intel® MIC Architecture) with support for AVX512_4FMAPS and AVX512_4VNNIW instruction groups enabled processors</td>
<td>Yes</td>
</tr>
<tr>
<td>AVX2</td>
<td>Intel® AVX2</td>
<td>Yes</td>
</tr>
<tr>
<td>AVX</td>
<td>Intel® AVX</td>
<td>Yes</td>
</tr>
<tr>
<td>SSE4_2</td>
<td>Intel® SSE4.2</td>
<td>Yes</td>
</tr>
</tbody>
</table>
For more details about the `mkl_enable_instructions` function, including the argument values, see the *Intel® MKL Developer Reference*.

For example:

- To turn on automatic CPU-based dispatching of Intel AVX-512 with support of AVX512_4FMAPS and AVX512_4VNNI instruction groups on systems based on Intel Xeon Phi processors, do one of the following:
  - Call
    ```
    mkl_enable_instructions(MKL_ENABLE_AVX512_MIC_E1)
    ```
  - Set the environment variable:
    ```
    set MKL_ENABLE_INSTRUCTIONS=AVX512_MIC_E1
    ```
  - To configure the library not to dispatch more recent architectures than Intel AVX2, do one of the following:
    - Call
      ```
      mkl_enable_instructions(MKL_ENABLE_AVX2)
      ```
    - Set the environment variable:
      ```
      set MKL_ENABLE_INSTRUCTIONS=AVX2
      ```

  **NOTE**
  Settings specified by the `mkl_enable_instructions` function take precedence over the settings specified by the `MKL_ENABLE_INSTRUCTIONS` environment variable.

---

**Optimization Notice**

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.
Programming with Intel® Math Kernel Library in Integrated Development Environments (IDE)

Configuring Your Integrated Development Environment to Link with Intel® Math Kernel Library

See these Knowledge Base articles for how to configure your Integrated Development Environment for linking with Intel® MKL:


Configuring the Microsoft Visual C/C++* Development System to Link with Intel® MKL

Steps for configuring Microsoft Visual C/C++* development system for linking with Intel® Math Kernel Library (Intel® MKL) depend on whether you installed the C++ Integration(s) in Microsoft Visual Studio® component of the Intel® Parallel Studio XE Composer Edition:

- If you installed the integration component, see Automatically Linking Your Microsoft Visual C/C++* Project with Intel® MKL.
- If you did not install the integration component or need more control over Intel® MKL libraries to link, you can configure the Microsoft Visual C++* development system by performing the following steps. Though some versions of the Visual C++* development system may vary slightly in the menu items mentioned below, the fundamental configuring steps are applicable to all these versions.

1. In Solution Explorer, right-click your project and click Properties
2. Select Configuration Properties > VC++ Directories
3. Select Include Directories. Add the directory for the Intel® MKL include files, that is, `<mkl directory>`\include
4. Select Library Directories. Add architecture-specific directories for Intel® MKL and OpenMP* libraries, for example: `<mkl directory>`\lib\ia32 and `<compiler directory>`\windows\compiler\lib \ia32_win
5. Select Executable Directories. Add architecture-specific directories with dynamic-link libraries:
   - For OpenMP* support, for example: `<compiler directory>`\windows\redist\ia32_win\compiler
   - For Intel® MKL (only if you link dynamically), for example: `<mkl directory>`\redist\ia32
6. Select Configuration Properties > Custom Build Setup > Additional Dependencies. Add the libraries required, for example, mkl_intel_c.lib mkl_intel_thread.lib mkl_core.lib libiomp5md.lib

See Also

Intel® Software Documentation Library for the documentation for Intel Parallel Studio XE Composer Edition
for the documentation for Intel Parallel Studio XE Composer Edition
Notational Conventions

Getting Assistance for Programming in the Microsoft Visual Studio* IDE

Using Context-Sensitive Help

You can get context-sensitive help when typing your code in the Visual Studio* IDE Code Editor. To open the help topic describing an Intel® MKL function called in your code, select the function name and press F1. The topic with the function description opens in the Microsoft Help Viewer or your Web browser depending on the Visual Studio IDE Help settings.

Using the IntelliSense* Capability

IntelliSense is a set of native Visual Studio*(VS) IDE features that make language references easily accessible.

The user programming with Intel® MKL in the VS Code Editor can employ two IntelliSense features: Parameter Info and Complete Word.

Both features use header files. Therefore, to benefit from IntelliSense, make sure the path to the include files is specified in the VS or solution settings. For example, see Configuring the Microsoft Visual C/C++* Development System to Link with Intel® MKL on how to do this.
Parameter Info
The Parameter Info feature displays the parameter list for a function to give information on the number and types of parameters. This feature requires adding the `include` statement with the appropriate Intel® MKL header file to your code.

To get the list of parameters of a function specified in the header file,
1. Type the function name.
2. Type the opening parenthesis.

This brings up the tooltip with the list of the function parameters:

![Parameter Info Example](image)

Complete Word
For a software library, the Complete Word feature types or prompts for the rest of the name defined in the header file once you type the first few characters of the name in your code. This feature requires adding the `include` statement with the appropriate Intel® MKL header file to your code.

To complete the name of the function or named constant specified in the header file,
1. Type the first few characters of the name.
2. Press Alt+RIGHT ARROW or Ctrl+SPACEBAR.
   - If you have typed enough characters to disambiguate the name, the rest of the name is typed automatically. Otherwise, a pop-up list appears with the names specified in the header file
3. Select the name from the list, if needed.
Intel® Math Kernel Library Benchmarks

Optimization Notice

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

Intel® Optimized LINPACK Benchmark for Windows*

Intel® Optimized LINPACK Benchmark for Windows* is a generalization of the LINPACK 1000 benchmark. It solves a dense (real*8) system of linear equations (Ax=b), measures the amount of time it takes to factor and solve the system, converts that time into a performance rate, and tests the results for accuracy. The generalization is in the number of equations (N) it can solve, which is not limited to 1000. It uses partial pivoting to assure the accuracy of the results.

Do not use this benchmark to report LINPACK 100 performance because that is a compiled-code only benchmark. This is a shared-memory (SMP) implementation which runs on a single platform. Do not confuse this benchmark with:

• Intel® Distribution for LINPACK* Benchmark, which is a distributed memory version of the same benchmark.
• LINPACK, the library, which has been expanded upon by the LAPACK library.

Intel provides optimized versions of the LINPACK benchmarks to help you obtain high LINPACK benchmark results on your genuine Intel processor systems more easily than with the High Performance Linpack (HPL) benchmark.

Additional information on this software, as well as on other Intel® software performance products, is available at https://www.intel.com/software/products/.

Acknowledgement

This product includes software developed at the University of Tennessee, Knoxville, Innovative Computing Laboratories.

Contents of the Intel® Optimized LINPACK Benchmark

The Intel Optimized LINPACK Benchmark for Windows* contains the following files, located in the benchmarks\linpack\ subdirectory of the Intel® Math Kernel Library (Intel® MKL) directory:

<table>
<thead>
<tr>
<th>File in benchmarks\linpack\</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>linpack_xeon32.exe</td>
<td>The 32-bit program executable for a system based on Intel® Xeon® processor or Intel® Xeon® processor MP with or without Intel® Streaming SIMD Extensions 3 (SSE3).</td>
</tr>
<tr>
<td>File in benchmarks\linpack\</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>linpack_xeon64.exe</td>
<td>The 64-bit program executable for a system with Intel Xeon processor using Intel® 64 architecture.</td>
</tr>
<tr>
<td>runme_xeon32.bat</td>
<td>A sample shell script for executing a pre-determined problem set for linpack_xeon32.exe.</td>
</tr>
<tr>
<td>runme_xeon64.bat</td>
<td>A sample shell script for executing a pre-determined problem set for linpack_xeon64.exe.</td>
</tr>
<tr>
<td>lininput_xeon32</td>
<td>Input file for a pre-determined problem for the runme_xeon32 script.</td>
</tr>
<tr>
<td>lininput_xeon64</td>
<td>Input file for a pre-determined problem for the runme_xeon64 script.</td>
</tr>
<tr>
<td>help.lpk</td>
<td>Simple help file.</td>
</tr>
<tr>
<td>xhelp.lpk</td>
<td>Extended help file.</td>
</tr>
</tbody>
</table>

These files are not available immediately after installation and appear as a result of execution of an appropriate runme script.

| win_xeon32.txt              | Result of the runme_xeon32 script execution. |
| win_xeon64.txt              | Result of the runme_xeon64 script execution. |

See Also

High-level Directory Structure

Running the Software

To obtain results for the pre-determined sample problem sizes on a given system, type:

`runme_xeon32.bat`

`runme_xeon64.bat`

To run the software for other problem sizes, see the extended help included with the program. You can view extended help by running the program executable with the -e option:

`linpack_xeon32.exe -e`

`linpack_xeon64.exe -e`

The pre-defined data input files `lininput_xeon32` and `lininput_xeon64` are examples. Different systems have different numbers of processors or amounts of memory and therefore require new input files. The extended help can give insight into proper ways to change the sample input files.

Each input file requires the following minimum amount of memory:

- `lininput_xeon32`: 2 GB
- `lininput_xeon64`: 16 GB

If the system has less memory than the above sample data input requires, you may need to edit or create your own data input files, as explained in the extended help.

The Intel Optimized LINPACK Benchmark determines the optimal number of OpenMP threads to use. To run a different number, you can set the `OMP_NUM_THREADS` or `MKL_NUM_THREADS` environment variable inside a sample script. If you run the Intel Optimized LINPACK Benchmark without setting the number of threads, it defaults to the number of physical cores.
Optimization Notice

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

Known Limitations of the Intel® Optimized LINPACK Benchmark

The following limitations are known for the Intel Optimized LINPACK Benchmark for Windows*:

- Intel Optimized LINPACK Benchmark supports only OpenMP threading
- Intel Optimized LINPACK Benchmark is threaded to effectively use multiple processors. So, in multi-processor systems, best performance will be obtained with the Intel® Hyper-Threading Technology turned off, which ensures that the operating system assigns threads to physical processors only.
- If an incomplete data input file is given, the binaries may either hang or fault. See the sample data input files and/or the extended help for insight into creating a correct data input file.

Intel® Distribution for LINPACK® Benchmark

Overview of the Intel Distribution for LINPACK Benchmark

The Intel® Distribution for LINPACK® Benchmark is based on modifications and additions to High-Performance LINPACK (HPL) (http://www.netlib.org/benchmark/hpl/) from Innovative Computing Laboratories (ICL) at the University of Tennessee, Knoxville. The Intel Distribution for LINPACK Benchmark can be used for TOP500 runs (see http://www.top500.org) and for benchmarking your cluster. To use the benchmark you need to be familiar with HPL usage. The Intel Distribution for LINPACK Benchmark provides some enhancements designed to make the HPL usage more convenient and to use Intel® Message-Passing Interface (MPI) settings to improve performance.

The Intel Distribution for LINPACK Benchmark measures the amount of time it takes to factor and solve a random dense system of linear equations (Ax=b) in real*8 precision, converts that time into a performance rate, and tests the results for accuracy. The benchmark uses random number generation and full row pivoting to ensure the accuracy of the results.

Intel provides optimized versions of the LINPACK benchmarks to help you obtain high LINPACK benchmark results on your systems based on genuine Intel processors more easily than with the standard HPL benchmark. The prebuilt binaries require Intel® MPI library be installed on the cluster. The run-time version of Intel MPI library is free and can be downloaded from https://www.intel.com/software/products/.

The Intel package includes software developed at the University of Tennessee, Knoxville, ICL, and neither the University nor ICL endorse or promote this product. Although HPL is redistributable under certain conditions, this particular package is subject to the Intel® Math Kernel Library (Intel® MKL) license.

Intel® MKL provides prebuilt binaries that are linked against Intel MPI libraries either statically or dynamically. In addition, binaries linked with a customized MPI implementation can be created using the Intel® MKL MPI wrappers.
Performance of statically and dynamically linked prebuilt binaries may be different. The performance of both depends on the version of Intel MPI you are using. You can build binaries statically or dynamically linked against a particular version of Intel MPI by yourself.

HPL code is homogeneous by nature: it requires that each MPI process runs in an environment with similar CPU and memory constraints. The Intel Distribution for LINPACK Benchmark supports heterogeneity, meaning that the data distribution can be balanced to the performance requirements of each node, provided that there is enough memory on that node to support additional work. For information on how to configure Intel® MKL to use the internode heterogeneity, see Heterogeneous Support in the Intel Distribution for LINPACK Benchmark.

Contents of the Intel Distribution for LINPACK Benchmark

The Intel Distribution for LINPACK Benchmark includes prebuilt binaries linked with Intel® MPI library. For a customized MPI implementation, tools are also included to build a binary using Intel® MKL MPI wrappers. All the files are located in the .\benchmarks\mp_linpack\ subdirectory of the Intel® MKL directory.

<table>
<thead>
<tr>
<th>File in (\langle mkl\ directory\rangle\backslash benchmarks \backslash mp_linpack)</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>COPYRIGHT</td>
<td>Original Netlib HPL copyright document.</td>
</tr>
<tr>
<td>readme.txt</td>
<td>Information about the files provided.</td>
</tr>
</tbody>
</table>

Prebuilt executables for performance testing

| xhpl_intel64_dynamic.exe          | Prebuilt binary for the Intel® 64 architecture dynamically linked against Intel MPI library. |

Run scripts and an input file example

| runme_intel64_dynamic.bat         | Sample run script for the Intel® 64 architecture and binary dynamically linked against Intel MPI library. |
| runme_intel64_prv.bat             | Script that sets HPL environment variables. It is called by runme_intel64_dynamic.bat. |
| HPL.dat                           | Example of an HPL configuration file. |

Prebuilt libraries and utilities for building with a customized MPI implementation

| libhpl_intel64.lib                | Library file required to build Intel Distribution for LINPACK Benchmark for the Intel® 64 architecture with a customized MPI implementation. |
| HPL_main.c                        | Source code required to build Intel Distribution for LINPACK Benchmark for the Intel® 64 architecture with a customized MPI implementation. |
| build.bat                         | Build script for creating Intel Distribution for LINPACK Benchmark for the Intel® 64 architecture with a customized MPI implementation. |

Utilities for building Netlib HPL from source code

| Make.Windows_Intel64              | Makefile for building Netlib HPL from source code. |

*For a list of supported versions of the Intel MPI Library, see system requirements in the Intel® MKL Release Notes.*
Building the Intel Distribution for LINPACK Benchmark for a Customized MPI Implementation

The Intel Distribution for LINPACK Benchmark contains a sample build script `build.bat`. If you are using a customized MPI implementation, this script builds a binary using Intel® MKL MPI wrappers. To build the binary, follow these steps:

1. Specify the location of Intel® MKL to be used (`MKLROOT`)
2. Set up your MPI environment
3. Run the script `build.bat`

See Also
High-level Directory Structure

Building the Netlib HPL from Source Code

The source code for Intel Distribution for LINPACK Benchmark is not provided. However, you can download reference Netlib HPL source code from http://www.netlib.org/benchmark/hpl/. To build the HPL:

1. Download and extract the source code.
2. Copy the makefile `<mkl directory>\benchmarks\mp_linpack\Make.Windows_Intel64` to your HPL directory
3. Edit `Make.Windows_Intel64` as appropriate
4. Build the HPL binary:
   ```
   $> nmake -f Make.Windows_Intel64
   ```
5. Check that the built binary is available in the current directory.

**NOTE**
The Intel Distribution for LINPACK Benchmark may contain additional optimizations compared to the reference Netlib HPL implementation.

Optimization Notice

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Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

See Also
Contents of the Intel Distribution for LINPACK Benchmark

Configuring Parameters

The most significant parameters in `HPL.dat` are `P`, `Q`, `NB`, and `N`. Specify them as follows:
- $P$ and $Q$ - the number of rows and columns in the process grid, respectively. $P \times Q$ must be the number of MPI processes that HPL is using. Choose $P \leq Q$.
- $NB$ - the block size of the data distribution.

The table below shows recommended values of $NB$ for different Intel® processors:

<table>
<thead>
<tr>
<th>Processor</th>
<th>$NB$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intel® Xeon® Processor X56*/E56*/E7-<em>/<em>E7</em>/X7</em> (codenamed Nehalem or Westmere)</td>
<td>256</td>
</tr>
<tr>
<td>Intel Xeon Processor E26*/E26* v2 (codenamed Sandy Bridge or Ivy Bridge)</td>
<td>256</td>
</tr>
<tr>
<td>Intel Xeon Processor E26* v3/E26* v4 (codenamed Haswell or Broadwell)</td>
<td>192</td>
</tr>
<tr>
<td>Intel® Core® i3/i5/i7-6* Processor (codenamed Skylake Client)</td>
<td>192</td>
</tr>
<tr>
<td>Intel® Xeon Phi™ Processor 72* (codenamed Knights Landing)</td>
<td>336</td>
</tr>
<tr>
<td>Intel Xeon Processor supporting Intel® Advanced Vector Extensions 512 (Intel® AVX-512) instructions (codenamed Skylake Server)</td>
<td>384</td>
</tr>
</tbody>
</table>

- $N$ - the problem size:
  - For homogeneous runs, choose $N$ divisible by $NB \times LCM(P, Q)$, where $LCM$ is the least common multiple of the two numbers.
  - For heterogeneous runs, see Heterogeneous Support in the Intel Distribution for LINPACK Benchmark for how to choose $N$.

**NOTE**
Increasing $N$ usually increases performance, but the size of $N$ is bounded by memory. In general, you can compute the memory required to store the matrix (which does not count internal buffers) as $8 \times N \times N / (P \times Q)$ bytes, where $N$ is the problem size and $P$ and $Q$ are the process grids in HPL.dat. A general rule of thumb is to choose a problem size that fills 80% of memory.

### Ease-of-use Command-line Parameters

The Intel Distribution for LINPACK Benchmark supports command-line parameters for HPL that help you to avoid making small changes in the HPL.dat input file every time you do a new run.

Placeholders in this command line illustrate these parameters:

```plaintext
xhpl.exe -n <problem size> -m <memory size in Mbytes> -b <block size> -p <grid row dimn> -q <grid column dimn>
```

You can also use command-line parameters with the sample runme script. For example:

```plaintext
runme_intel64_dynamic.bat -m <memory size in Mbytes> -b <block size> -p <grid row dimn> -q <grid column dimn>
```

For more command-line parameters, see Heterogeneous Support in the Intel Distribution for LINPACK Benchmark.

If you want to run for $N=10000$ on a 1x3 grid, execute this command, provided that the other parameters in HPL.dat and the script are correct:

```plaintext
runme_intel64_dynamic.bat -n 10000 -p 1 -q 3
```

By using the $m$ parameter you can scale by the memory size instead of the problem size. The $m$ parameter only refers to the size of the matrix storage. Therefore, to use matrices that fit in 50000 Mbytes with $NB=256$ on 16 nodes, adjust the script to set the total number of MPI processes to 16 and execute this command:

```plaintext
runme_intel64_dynamic.bat -m 50000 -b 256 -p 4 -q 4
```
Running the Intel Distribution for LINPACK Benchmark

To run the Intel Distribution for LINPACK Benchmark on multiple nodes or on one node with multiple MPI processes, you need to use MPI and either modify HPL.dat or use Ease-of-use Command-line Parameters. The following example describes how to run the dynamically-linked prebuilt Intel Distribution for LINPACK Benchmark binary using the script provided. To run other binaries, adjust the steps accordingly.

1. Load the necessary environment variables for the Intel MPI Library and Intel® compiler:
   ```
   <parent directory>\bin\compilervars.batintel64
   <mpi directory>\bin64\mpivars.bat
   ```

2. In HPL.dat, set the problem size N to 10000. Because this setting is for a test run, the problem size should be small.

3. For better performance, enable non-uniform memory access (NUMA) on your system and configure to run an MPI process for each NUMA socket as explained below.

   **NOTE**
   High-bandwidth Multi-Channel Dynamic Random Access Memory (MCDRAM) on the second-generation Intel Xeon Phi processors may appear to be a NUMA node. However, because there are no CPUs on this node, do not run an MPI process for it.

   - Refer to your BIOS settings to enable NUMA on your system.
   - Set the following variables at the top of the runme_intel64_dynamic.bat script according to your cluster configuration:
     ```
     MPI_PROC_NUM The total number of MPI processes.
     MPI_PER_NODE The number of MPI processes per each cluster node.
     ```
   - In the HPL.dat file, set the parameters Ps and Qs so that Ps * Qs equals the number of MPI processes. For example, for 2 processes, set Ps to 1 and Qs to 2. Alternatively, leave the HPL.dat file as is and launch with -p and -q command-line parameters.

4. Execute runme_intel64_dynamic.bat script:
   ```
   runme_intel64_dynamic.bat
   ```

5. Rerun the test increasing the size of the problem until the matrix size uses about 80% of the available memory. To do this, either modify Ns in line 6 of HPL.dat or use the -n command-line parameter:
   ```
   • For 16 GB: 40000 Ns
   • For 32 GB: 56000 Ns
   • For 64 GB: 83000 Ns
   ```

See Also

Notational Conventions
Building the Intel Distribution for LINPACK Benchmark for a Customized MPI Implementation
Building the Netlib HPL from Source Code

Heterogeneous Support in the Intel Distribution for LINPACK Benchmark

Intel Distribution for LINPACK Benchmark achieves heterogeneous support by distributing the matrix data unequally between the nodes. The heterogeneous factor command-line parameter f controls the amount of work to be assigned to the more powerful nodes, while the command-line parameter c controls the number of process columns for the faster nodes:

```bash
xhpl.exe -n <problem size> -b <block size> -p <grid row dimn> -q <grid column dimn> -f <heterogeneous factor> -c <number of faster processor columns>
```
If the heterogeneous factor is 2.5, roughly 2.5 times the work will be put on the more powerful nodes. The more work you put on the more powerful nodes, the more memory you might be wasting on the other nodes if all nodes have equal amount of memory. If your cluster includes many different types of nodes, you may need multiple heterogeneous factors.

Let $P$ be the number of rows and $Q$ the number of columns in your processor grid ($P \times Q$). The work must be \textit{homogeneous} within each processor column because vertical operations, such as pivoting or panel factorization, are synchronizing operations. When there are two different types of nodes, use MPI to process all the faster nodes first and make sure the "PMAP process mapping" (line 9) of HPL.dat is set to 1 for \textit{Column-major} mapping. Because all the nodes must be the same within a process column, the number of faster nodes must always be a multiple of $P$, and you can specify the faster nodes by setting the number of process columns $C$ for the faster nodes with the $c$ command-line parameter. The $-f 1.0 -c 0$ setting corresponds to the default homogeneous behavior.

To understand how to choose the problem size $N$ for a heterogeneous run, first consider a homogeneous system, where you might choose $N$ as follows:

$$N \approx \sqrt{\text{Memory Utilization} \times P \times Q \times \text{Memory Size in Bytes} / 8}$$

\text{Memory Utilization} is usually around 0.8 for homogeneous Intel Xeon processor systems. On a heterogeneous system, you may apply a different formula for $N$ for each set of nodes that are the same and then choose the minimum $N$ over all sets. Suppose you have a cluster with only one heterogeneous factor $F$ and the number of processor columns (out of the total $Q$) in the group with that heterogeneous factor equal to $C$. That group contains $P \times C$ nodes. First compute the sum of the parts: $S = F \times P \times C + P \times (Q-C)$. Note that on a homogeneous system $S = P \times Q, F = 1, \text{and } C = Q$. Take $N$ as

$$N \approx \sqrt{\text{Memory Utilization} \times P \times Q 	imes ((F \times P \times C) / S) \times \text{Memory Size in Bytes} / 8}$$

or simply scale down the value of $N$ for the homogeneous system by $\sqrt{F \times P \times C / S}$.

\textbf{Example}

Suppose the cluster has 100 nodes each having 64 GB of memory, and 20 of the nodes are 2.7 times as powerful as the other 80. Run one MPI process per node for a total of 100 MPI processes. Assume a square processor grid $P=Q=10$, which conveniently divides up the faster nodes evenly. Normally, the HPL documentation recommends choosing a matrix size that consumes 80 percent of available memory. If $N$ is the size of the matrix, the matrix consumes $8N^2/(P \times Q)$ bytes. So a homogeneous run might look like:

```
xhpl.exe -n 820000 -b 256 -p 10 -q 10
```

If you redistribute the matrix and run the heterogeneous Intel Distribution for LINPACK Benchmark, you can take advantage of the faster nodes. But because some of the nodes will contain 2.7 times as much data as the other nodes, you must shrink the problem size (unless the faster nodes also happen to have 2.7 times as much memory). Instead of $0.8 \times 64 \times 10^9$ total memory size, we have only $0.8 \times 64 \times 20 + 0.8 \times 64 \times 2.7 \times 80$ total memory size, which is less than half the original space. So the problem size in this case would be $526000$. Because $P=10$ and there are 20 faster nodes, two processor columns are faster. If you arrange MPI to send these nodes first to the application, the command line looks like:

```
xhpl.exe -n 526000 -b 1024 -p 10 -q 10 -f 2.7 -c 2
```

The $m$ parameter may be misleading for heterogeneous calculations because it calculates the problem size assuming all the nodes have the same amount of data.

\textbf{Warning}

The number of faster nodes must be $C \times P$. If the number of faster nodes is not divisible by $P$, you might not be able to take advantage of the extra performance potential by giving the faster nodes extra work.
While it suffices to simply provide \( f \) and \( c \) command-line parameters if you need only one heterogeneous factor, you must add lines to the \texttt{HPL.dat} input to support multiple heterogeneous factors. For the above example (two processor columns have nodes that are 2.7 times faster), instead of passing \( f \) and \( c \) command-line parameters you can modify the \texttt{HPL.dat} input file by adding these two lines to the end:

\begin{verbatim}
1         number of heterogeneous factors
0 1 2.7   [start_column, stop_column, heterogeneous factor for that range]
\end{verbatim}

\textbf{NOTE}
Numbering of processor columns starts at 0. The start and stopping numbers must be between 0 and \( Q-1 \) (inclusive).

If instead there are three different types of nodes in a cluster and you need at least two heterogeneous factors, change the number in the first row above from 1 to 2 and follow that line with two lines specifying the start column, stopping column, and heterogeneous factor.

When choosing parameters for heterogeneous support in \texttt{HPL.dat}, primarily focus on the most powerful nodes. The larger the heterogeneous factor, the more balanced the cluster may be from a performance viewpoint, but the more imbalanced from a memory viewpoint. At some point, further performance balancing might affect the memory too much. If this is the case, try to reduce any changes done for the faster nodes (such as in block sizes). Experiment with values in \texttt{HPL.dat} carefully because wrong values may greatly hinder performance.

When tuning on a heterogeneous cluster, do not immediately attempt a heterogeneous run, but do the following:

1. Break the cluster down into multiple homogeneous clusters.
2. Make heterogeneous adjustments for performance balancing. For instance, if you have two different sets of nodes where one is three times as powerful as the other, it must do three times the work.
3. Figure out the approximate size of the problem (per node) that you can run on each piece.
4. Do some homogeneous runs with those problem sizes per node and the final block size needed for the heterogeneous run and find the best parameters.
5. Use these parameters for an initial heterogeneous run.

\section*{Environment Variables}

The table below lists Intel\textsuperscript{®} MKL environment variables to control runs of the Intel Distribution for LINPACK Benchmark.

\begin{table}[h]
\centering
\begin{tabular}{lll}
\hline
\textbf{Environment Variable} & \textbf{Description} & \textbf{Value} \\
\hline
\texttt{HPL\_LARGEPAGE} & Defines the memory mapping to be used for the Intel Xeon processor. & \begin{itemize}
\item 0 or 1:
\begin{itemize}
\item 0 - normal memory mapping, default.
\item 1 - memory mapping with large pages (2 MB per page mapping). It may increase performance.
\end{itemize}
\end{itemize} \\
\hline
\texttt{HPL\_LOG} & Controls the level of detail for the HPL output. & \begin{itemize}
\item An integer ranging from 0 to 2:
\begin{itemize}
\item 0 - no log is displayed.
\item 1 - only one root node displays a log, exactly the same as the \texttt{ASYOUGO} option provides.
\end{itemize}
\end{itemize} \\
\hline
\end{tabular}
\end{table}
### Environment Variable

<table>
<thead>
<tr>
<th>Environment Variable</th>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPL_HOST_CORE, HPL_HOST_NODE</td>
<td>Specifies cores or Non-Uniform Memory Access (NUMA) nodes to be used. <strong>HPL_HOST_NODE requires NUMA mode to be enabled. You can check whether it is enabled by the <code>numactl --hardware</code> command.</strong> The default behavior is auto-detection of the core or NUMA node.</td>
<td>A list of integers ranging from 0 to the largest number of a core or NUMA node in the cluster and separated as explained in example 3.</td>
</tr>
<tr>
<td>HPL_SWAPWIDTH</td>
<td>Specifies width for each swap operation.</td>
<td>16 or 24. The default is 24.</td>
</tr>
</tbody>
</table>

You can set Intel Distribution for LINPACK Benchmark environment variables using the `PMI_RANK` and `PMI_SIZE` environment variables of the Intel MPI library, and you can create a shell script to automate the process.

### Examples of Environment Settings

<table>
<thead>
<tr>
<th>#</th>
<th>Settings</th>
<th>Behavior of the Intel Distribution for LINPACK Benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Nothing specified</td>
<td>All Intel Xeon processors in the cluster are used.</td>
</tr>
<tr>
<td>2</td>
<td>HPL_MIC_DEVICE=0,2,3,8,9,10</td>
<td>Intel Xeon processor cores 1,2,3,8,9, and 10 are used.</td>
</tr>
<tr>
<td>3</td>
<td>HPL_HOST_NODE=1</td>
<td>Only Intel Xeon processor cores on NUMA node 1 are used.</td>
</tr>
</tbody>
</table>

### Improving Performance of Your Cluster

To improve cluster performance, follow these steps, provided all required software is installed on each node:

1. Reboot all nodes.
2. Ensure all nodes are in identical conditions and no zombie processes are left running from prior HPL runs. To do this, run single-node Stream and Intel Distribution for LINPACK Benchmark on every node. Ensure results are within 10% of each other (problem size must be large enough depending on memory size and CPU speed). Investigate nodes with low performance for hardware/software problems.
3. Check that your cluster interconnects are working. Run a test over the complete cluster using an MPI test for bandwidth and latency, such as one found in the Intel® MPI Benchmarks package.
4. Run an Intel Distribution for LINPACK Benchmark on pairs of two or four nodes and ensure results are within 10% of each other. The problem size must be large enough depending on the memory size and CPU speed.
5. Run a small problem size over the complete cluster to ensure correctness.
6. Increase the problem size and run the real test load.
7. In case of problems go back to step 2.

Before making a heterogeneous run, always run its homogeneous equivalent first.
See Also
Mixed-language Programming with Intel® MKL

Language Interfaces Support, by Function Domain

The following table shows language interfaces that Intel® Math Kernel Library (Intel® MKL) provides for each function domain. However, Intel® MKL routines can be called from other languages using mixed-language programming. See Mixed-language Programming with the Intel Math Kernel Library for an example of how to call Fortran routines from C/C++.

<table>
<thead>
<tr>
<th>Function Domain</th>
<th>Fortran interface</th>
<th>C/C++ interface</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basic Linear Algebra Subprograms (BLAS)</td>
<td>Yes</td>
<td>through CBLAS</td>
</tr>
<tr>
<td>BLAS-like extension transposition routines</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Sparse BLAS Level 1</td>
<td>Yes</td>
<td>through CBLAS</td>
</tr>
<tr>
<td>Sparse BLAS Level 2 and 3</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>LAPACK routines for solving systems of linear equations</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>LAPACK routines for solving least-squares problems, eigenvalue and singular</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>value problems, and Sylvester's equations</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Auxiliary and utility LAPACK routines</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Parallel Basic Linear Algebra Subprograms (PBLAS)</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>ScALAPACK</td>
<td>Yes</td>
<td>†</td>
</tr>
<tr>
<td>Direct Sparse Solvers/</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Intel® MKL PARDISO, a direct sparse solver based on Parallel Direct Sparse</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solver (PARDISO*)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parallel Direct Sparse Solvers for Clusters</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Other Direct and Iterative Sparse Solver routines</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Vector Mathematics (VM)</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Vector Statistics (VS)</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Fast Fourier Transforms (FFT)</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Cluster FFT</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Trigonometric Transforms</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Fast Poisson, Laplace, and Helmholtz Solver (Poisson Library)</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Optimization (Trust-Region) Solver</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Function Domain</td>
<td>Fortran interface</td>
<td>C/C++ interface</td>
</tr>
<tr>
<td>---------------------------------------</td>
<td>-------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>Data Fitting</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Extended Eigensolver</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Support functions (including memory allocation)</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

† Supported using a mixed language programming call. See Include Files for the respective header file.

## Include Files

The table below lists Intel® MKL include files.

<table>
<thead>
<tr>
<th>Function Domain/ Purpose</th>
<th>Fortran Include Files</th>
<th>C/C++ Include Files</th>
</tr>
</thead>
<tbody>
<tr>
<td>All function domains</td>
<td>mkl.fi</td>
<td>mkl.h</td>
</tr>
<tr>
<td>BLACS</td>
<td>mkl_blacs.h‡‡</td>
<td></td>
</tr>
<tr>
<td>BLAS</td>
<td>blas.f90</td>
<td>mkl_blas.h†</td>
</tr>
<tr>
<td></td>
<td>mkl_blas.fi†</td>
<td></td>
</tr>
<tr>
<td>BLAS-like Extension Transposition Routines</td>
<td>mkl_trans.fi†</td>
<td>mkl_trans.h‡</td>
</tr>
<tr>
<td>CBLAS Interface to BLAS</td>
<td></td>
<td>mkl_cblas.h‡</td>
</tr>
<tr>
<td>Sparse BLAS</td>
<td>mkl_spblas.fi†</td>
<td>mkl_spblas.h‡</td>
</tr>
<tr>
<td>LAPACK</td>
<td>lapack.f90</td>
<td>mkl_lapack.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_lapack.fi†</td>
<td></td>
</tr>
<tr>
<td>C Interface to LAPACK</td>
<td></td>
<td>mkl_lapacke.h†</td>
</tr>
<tr>
<td>PBLAS</td>
<td>mkl_pblas.h‡‡</td>
<td></td>
</tr>
<tr>
<td>ScaLAPACK</td>
<td>mkl_scalapack.h‡‡</td>
<td></td>
</tr>
<tr>
<td>Intel® MKL PARDISO</td>
<td>mkl_pardiso.f90</td>
<td>mkl_pardiso.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_pardiso.fi†</td>
<td></td>
</tr>
<tr>
<td>Parallel Direct Sparse Solvers for Clusters</td>
<td>mkl_cluster_sparse_solver.f90</td>
<td>mkl_cluster_sparse_solver.h‡</td>
</tr>
<tr>
<td>Direct Sparse Solver (DSS)</td>
<td>mkl_dss.f90</td>
<td>mkl_dss.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_dss.fi†</td>
<td></td>
</tr>
<tr>
<td>RCI Iterative Solvers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ILU Factorization</td>
<td>mkl_rci.f90</td>
<td>mkl_rci.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_rci.fi†</td>
<td></td>
</tr>
<tr>
<td>Optimization Solver</td>
<td>mkl_rci.f90</td>
<td>mkl_rci.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_rci.fi†</td>
<td></td>
</tr>
<tr>
<td>Vector Mathematics</td>
<td>mkl_vml.90</td>
<td>mkl_vml.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_vml.fi†</td>
<td></td>
</tr>
<tr>
<td>Vector Statistics</td>
<td>mkl_vsl.f90</td>
<td>mkl_vsl.h‡</td>
</tr>
<tr>
<td></td>
<td>mkl_vsl.fi†</td>
<td></td>
</tr>
<tr>
<td>Function Domain/ Purpose</td>
<td>Fortran Include Files</td>
<td>C/C++ Include Files</td>
</tr>
<tr>
<td>------------------------------------------</td>
<td>----------------------------------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td>Fast Fourier Transforms</td>
<td>mkl_dfti.f90</td>
<td>mkl_dfti.h†</td>
</tr>
<tr>
<td>Cluster Fast Fourier Transforms</td>
<td>mkl_cdft.f90</td>
<td>mkl_cdft.h‡‡</td>
</tr>
<tr>
<td>Partial Differential Equations Support</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trigonometric Transforms</td>
<td>mkl_trig_transforms.f90</td>
<td>mkl_trig_transform.h†</td>
</tr>
<tr>
<td>Poisson Solvers</td>
<td>mkl_poisson.f90</td>
<td>mkl_poisson.h†</td>
</tr>
<tr>
<td>Data Fitting</td>
<td>mkl_df.f90</td>
<td>mkl_df.h§</td>
</tr>
<tr>
<td>Extended Eigensolver</td>
<td>mkl_solvers_ee.fi†</td>
<td>mkl_solvers_ee.h§</td>
</tr>
<tr>
<td>Support functions</td>
<td>mkl_service.f90</td>
<td>mkl_service.h†</td>
</tr>
<tr>
<td></td>
<td>mkl_service.fi†</td>
<td></td>
</tr>
<tr>
<td>Declarations for replacing memory allocation functions. See Redefining Memory Functions for details.</td>
<td></td>
<td>i_malloc.h</td>
</tr>
<tr>
<td>Auxiliary macros to determine the version of Intel® MKL at compile time.</td>
<td>mkl_version</td>
<td>mkl_version‡</td>
</tr>
</tbody>
</table>

† You can use the mkl.fi include file in your code instead.
‡ You can include the mkl.h header file in your code instead.
‡‡ Also include the mkl.h header file in your code.

See Also
Language Interfaces Support, by Function Domain
Support for Third-Party Interfaces

FFTW Interface Support

Intel® Math Kernel Library (Intel® MKL) offers two collections of wrappers for the FFTW interface (www.fftw.org). The wrappers are the superstructure of FFTW to be used for calling the Intel® MKL Fourier transform functions. These collections correspond to the FFTW versions 2.x and 3.x and the Intel® MKL versions 7.0 and later.

These wrappers enable using Intel® MKL Fourier transforms to improve the performance of programs that use FFTW without changing the program source code. See the "FFTW Interface to Intel® Math Kernel Library" appendix in the Intel® MKL Developer Reference for details on the use of the wrappers.

Important
For ease of use, the FFTW3 interface is also integrated in Intel® MKL.

Caution
The FFTW2 and FFTW3 interfaces are not compatible with each other. Avoid linking to both of them. If you must do so, first modify the wrapper source code for FFTW2:

1. Change every instance of `fftw_destroy_plan` in the `fftw2xc` interface to `fftw2_destroy_plan`.
2. Change all the corresponding file names accordingly.
3. Rebuild the pertinent libraries.
Tables in this section show contents of the Intel® Math Kernel Library (Intel® MKL) architecture-specific directories.

**Optimization Notice**

Intel's compilers may or may not optimize to the same degree for non-Intel microprocessors for optimizations that are not unique to Intel microprocessors. These optimizations include SSE2, SSE3, and SSSE3 instruction sets and other optimizations. Intel does not guarantee the availability, functionality, or effectiveness of any optimization on microprocessors not manufactured by Intel. Microprocessor-dependent optimizations in this product are intended for use with Intel microprocessors. Certain optimizations not specific to Intel microarchitecture are reserved for Intel microprocessors. Please refer to the applicable product User and Reference Guides for more information regarding the specific instruction sets covered by this notice.

Notice revision #20110804

This notice covers the following instruction sets: SSE2, SSE4.2, AVX2, AVX-512.

**See Also**

- High-level Directory Structure
- Using Language-Specific Interfaces with Intel® MKL

**Intel® Math Kernel Library Benchmarks**

**Detailed Structure of the IA-32 Architecture Directories**

**Static Libraries in the lib\ia32_win Directory**

Some of the libraries in this directory are optional. However, some optional libraries are installed by default, while the rest are not. To get those libraries that are not installed by default, explicitly select the specified optional component during installation.

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Name</td>
</tr>
<tr>
<td><strong>Interface Layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_intel_c.lib</td>
<td>cdecl interface library</td>
<td></td>
</tr>
<tr>
<td>mkl_intel_s.lib</td>
<td>CVF default interface library</td>
<td></td>
</tr>
<tr>
<td>mkl_blas95.lib</td>
<td>Fortran 95 interface library for BLAS. Supports the Intel® Fortran compiler</td>
<td>Fortran 95 interfaces for BLAS and LAPACK</td>
</tr>
</tbody>
</table>
### File Contents

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_lapack95.lib</td>
<td>Fortran 95 interface library for LAPACK. Supports the Intel® Fortran compiler</td>
<td>Fortran 95 interfaces for BLAS and LAPACK</td>
</tr>
</tbody>
</table>

**Threading Layer**

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_intel_thread.lib</td>
<td>OpenMP threading library for the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_tbb_thread.lib</td>
<td>Intel® Threading Building Blocks (Intel® TBB) threading library for the Intel compilers</td>
<td>Intel TBB threading support</td>
</tr>
<tr>
<td>mkl_sequential.lib</td>
<td>Sequential library</td>
<td></td>
</tr>
</tbody>
</table>

**Computational Layer**

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_core.lib</td>
<td>Kernel library for IA-32 architecture</td>
<td></td>
</tr>
</tbody>
</table>

### Dynamic Libraries in the *lib\ia32_win* Directory

Some of the libraries in this directory are optional. However, some optional libraries are installed by default, while the rest are not. To get those libraries that are not installed by default, explicitly select the specified optional component during installation.

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_rt.lib</td>
<td>Single Dynamic Library to be used for linking</td>
<td></td>
</tr>
</tbody>
</table>

**Interface Layer**

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_intel_c_dll.lib</td>
<td>cdecl interface library for dynamic linking</td>
<td></td>
</tr>
<tr>
<td>mkl_intel_s_dll.lib</td>
<td>CVF default interface library for dynamic linking</td>
<td></td>
</tr>
</tbody>
</table>

**Threading Layer**

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_intel_thread_dll.lib</td>
<td>OpenMP threading library for dynamic linking with the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_tbb_thread_dll.lib</td>
<td>Intel TBB threading library for the Intel compilers</td>
<td>Intel TBB threading support</td>
</tr>
</tbody>
</table>
### Contents of the redist\ia32_win\mkl Directory

Some of the libraries in this directory are optional. However, some optional libraries are installed by default, while the rest are not. To get those libraries that are not installed by default, explicitly select the specified optional component during installation.

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_sequential_dll.lib</td>
<td>Sequential library for dynamic linking</td>
<td></td>
</tr>
<tr>
<td><strong>Computational Layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_core_dll.lib</td>
<td>Core library for dynamic linking</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_rt.dll</td>
<td>Single Dynamic Library</td>
<td></td>
</tr>
<tr>
<td><strong>Threading Layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_intel_thread.dll</td>
<td>Dynamic OpenMP threading library for the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_tbb_thread.dll</td>
<td>Dynamic Intel TBB threading library for the Intel compilers</td>
<td>Intel TBB threading support</td>
</tr>
<tr>
<td>mkl_sequential.dll</td>
<td>Dynamic sequential library</td>
<td></td>
</tr>
<tr>
<td><strong>Computational Layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_core.dll</td>
<td>Core library containing processor-independent code and a dispatcher for dynamic loading of processor-specific code</td>
<td></td>
</tr>
<tr>
<td>mkl_p4.dll</td>
<td>Pentium® 4 processor kernel</td>
<td></td>
</tr>
<tr>
<td>mkl_p4m.dll</td>
<td>Kernel library for Intel® Supplemental Streaming SIMD Extensions 3 (Intel® SSSE3) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_p4m3.dll</td>
<td>Kernel library for Intel® Streaming SIMD Extensions 4.2 (Intel® SSE4.2) enabled processors</td>
<td></td>
</tr>
<tr>
<td>File</td>
<td>Contents</td>
<td>Optional Component</td>
</tr>
<tr>
<td>------------------</td>
<td>--------------------------------------------------------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>mkl_avx.dll</td>
<td>Kernel library for Intel® Advanced Vector Extensions (Intel® AVX) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_avx2.dll</td>
<td>Kernel library for Intel® Advanced Vector Extensions 2 (Intel® AVX2) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_avx512.dll</td>
<td>Kernel library for Intel® Advanced Vector Extensions 512 (Intel® AVX-512) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_p4.dll</td>
<td>Vector Mathematics (VM)/Vector Statistics (VS)/Data Fitting (DF) part of Pentium® 4 processor kernel</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_p4m.dll</td>
<td>VM/VS/DF for Intel® SSSE3 enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_p4m2.dll</td>
<td>VM/VS/DF for 45nm Hi-k Intel® Core™ 2 and Intel Xeon® processor families</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_p4m3.dll</td>
<td>VM/VS/DF for Intel® SSE4.2 enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_avx.dll</td>
<td>VM/VS/DF optimized for Intel® AVX enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_avx2.dll</td>
<td>VM/VS/DF optimized for Intel® AVX2 enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_avx512.dll</td>
<td>VM/VS/DF optimized for Intel® AVX-512 enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_ia.dll</td>
<td>VM/VS/DF default kernel for newer Intel® architecture processors</td>
<td></td>
</tr>
<tr>
<td>libmkl_vml_cmpt.dll</td>
<td>VM/VS/DF library for conditional numerical reproducibility</td>
<td></td>
</tr>
</tbody>
</table>
## Detailed Structure of the Intel® 64 Architecture Directories

### Static Libraries in the lib\intel64_win Directory

Some of the libraries in this directory are optional. However, some optional libraries are installed by default, while the rest are not. To get those libraries that are not installed by default, explicitly select the specified optional component during installation.

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>File</td>
<td>Contents</td>
<td>Optional Component</td>
</tr>
<tr>
<td></td>
<td>Name</td>
<td>Installed by Default</td>
</tr>
<tr>
<td></td>
<td>Installed by Default</td>
<td></td>
</tr>
<tr>
<td><strong>Interface Layer</strong></td>
<td><strong>Name</strong></td>
<td><strong>Installed by Default</strong></td>
</tr>
<tr>
<td>mkl_intel_lp64.lib</td>
<td>LP64 interface library for the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_intel_ilp64.lib</td>
<td>ILP64 interface library for the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_blas95_lp64.lib</td>
<td>Fortran 95 interface library for BLAS. Supports the Intel® Fortran compiler and LP64 interface</td>
<td>Yes</td>
</tr>
<tr>
<td>mkl_blas95_ilp64.lib</td>
<td>Fortran 95 interface library for BLAS. Supports the Intel® Fortran compiler and ILP64 interface</td>
<td>Yes</td>
</tr>
<tr>
<td>File</td>
<td>Contents</td>
<td>Optional Component</td>
</tr>
<tr>
<td>-------------------------</td>
<td>--------------------------------------------------------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>mkl_lapack95_lib</td>
<td>Fortran 95 interface library for LAPACK. Supports the Intel® Fortran compiler and LP64 interface</td>
<td>Fortran 95 interfaces for BLAS and LAPACK Yes</td>
</tr>
<tr>
<td>mkl_lapack95_ilp64.lib</td>
<td>Fortran 95 interface library for LAPACK. Supports the Intel® Fortran compiler and ILP64 interface</td>
<td>Fortran 95 interfaces for BLAS and LAPACK Yes</td>
</tr>
<tr>
<td>Threading Layer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_intel_thread.lib</td>
<td>OpenMP threading library for the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_tbb_thread.lib</td>
<td>Intel® Threading Building Blocks (Intel® TBB) threading library for the Intel compilers</td>
<td>Intel TBB threading support Yes</td>
</tr>
<tr>
<td>mkl_pgi_thread.lib</td>
<td>OpenMP threading library for the PGI* compiler</td>
<td>PGI* Compiler support</td>
</tr>
<tr>
<td>mkl_sequential.lib</td>
<td>Sequential library</td>
<td></td>
</tr>
<tr>
<td>Computational Layer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_core.lib</td>
<td>Kernel library for the Intel® 64 architecture</td>
<td></td>
</tr>
<tr>
<td>Cluster Libraries</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mkl_scalapack_lib</td>
<td>ScaLAPACK routine library supporting the LP64 interface</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_scalapack_ilp64.lib</td>
<td>ScaLAPACK routine library supporting the ILP64 interface</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_cdft_core.lib</td>
<td>Cluster version of FFTs</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_intelmpi_lib</td>
<td>LP64 version of BLACS routines supporting Intel® MPI Library</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_intelmpi_ilp64.lib</td>
<td>ILP64 version of BLACS routines supporting Intel MPI Library</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_mpich2_lib</td>
<td>LP64 version of BLACS routines supporting MPICH2 or higher</td>
<td>Cluster support</td>
</tr>
</tbody>
</table>
### Dynamic Libraries in the `lib\intel64_win` Directory

Some of the libraries in this directory are optional. However, some optional libraries are installed by default, while the rest are not. To get those libraries that are not installed by default, explicitly select the specified optional component during installation.

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_blacs_mpich2_ilp64.lib</td>
<td>ILP64 version of BLACS routines supporting MPICH2 or higher</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_msmipi_lp64.lib</td>
<td>LP64 version of BLACS routines supporting Microsoft* MPI</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_msmipi_ilp64.lib</td>
<td>ILP64 version of BLACS routines supporting Microsoft* MPI</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_rt.lib</td>
<td>Single Dynamic Library to be used for linking</td>
<td></td>
</tr>
<tr>
<td>mkl_intel_lp64_dll.lib</td>
<td>LP64 interface library for dynamic linking with the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_intel_ilp64_dll.lib</td>
<td>ILP64 interface library for dynamic linking with the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_intel_thread_dll.lib</td>
<td>OpenMP threading library for dynamic linking with the Intel compilers</td>
<td></td>
</tr>
<tr>
<td>mkl_tbb_thread_dll.lib</td>
<td>Intel TBB threading library for the Intel compilers</td>
<td>Intel TBB threading support</td>
</tr>
<tr>
<td>mkl_pgi_thread_dll.lib</td>
<td>OpenMP threading library for dynamic linking with the PGI* compiler</td>
<td>PGI* Compiler support</td>
</tr>
<tr>
<td>mkl_sequential_dll.lib</td>
<td>Sequential library for dynamic linking</td>
<td></td>
</tr>
</tbody>
</table>

### Directory Structure in Detail

- **Interface Layer**
- **Threading Layer**
- **Computational Layer**
### Contents of the `redist\intel64_win\mkl` Directory

Some of the libraries in this directory are optional. However, some optional libraries are installed by default, while the rest are not. To get those libraries that are not installed by default, explicitly select the specified optional component during installation.

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>mkl_core_dll.lib</code></td>
<td>Core library for dynamic linking</td>
<td></td>
</tr>
<tr>
<td><strong>Cluster Libraries</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>mkl_scalapack_lp64_dll.lib</code></td>
<td>ScaLAPACK routine library for dynamic linking supporting the LP64 interface</td>
<td>Cluster support</td>
</tr>
<tr>
<td><code>mkl_scalapack_ilp64_dll.lib</code></td>
<td>ScaLAPACK routine library for dynamic linking supporting the ILP64 interface</td>
<td>Cluster support</td>
</tr>
<tr>
<td><code>mkl_cdft_core_dll.lib</code></td>
<td>Cluster FFT library for dynamic linking</td>
<td>Cluster support</td>
</tr>
<tr>
<td><code>mkl_blacs_lp64_dll.lib</code></td>
<td>LP64 version of BLACS interface library for dynamic linking</td>
<td>Cluster support</td>
</tr>
<tr>
<td><code>mkl_blacs_ilp64_dll.lib</code></td>
<td>ILP64 version of BLACS interface library for dynamic linking</td>
<td>Cluster support</td>
</tr>
<tr>
<td><code>mkl_rt.dll</code></td>
<td>Single Dynamic Library</td>
<td></td>
</tr>
<tr>
<td><strong>Threading layer</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>mkl_intel_thread.dll</code></td>
<td>Dynamic OpenMP threading library for the Intel compilers</td>
<td></td>
</tr>
<tr>
<td><code>mkl_tbb_thread.dll</code></td>
<td>Dynamic Intel TBB threading library for the Intel compilers</td>
<td>Intel TBB threading support</td>
</tr>
<tr>
<td><code>mkl_pgi_thread.dll</code></td>
<td>Dynamic OpenMP threading library for the PGI* compiler</td>
<td>PGI* compiler support</td>
</tr>
<tr>
<td><code>mkl_sequential.dll</code></td>
<td>Dynamic sequential library</td>
<td></td>
</tr>
<tr>
<td>File</td>
<td>Contents</td>
<td>Optional Component</td>
</tr>
<tr>
<td>----------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>mkl_core.dll</td>
<td>Core library containing processor-independent code and a dispatcher for dynamic loading of processor-specific code</td>
<td></td>
</tr>
<tr>
<td>mkl_def.dll</td>
<td>Default kernel for the Intel® 64 architecture</td>
<td></td>
</tr>
<tr>
<td>mkl_mc.dll</td>
<td>Kernel library for Intel® Supplemental Streaming SIMD Extensions 3 (Intel® SSSE3) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_mc3.dll</td>
<td>Kernel library for Intel® Streaming SIMD Extensions 4.2 (Intel® SSE4.2) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_avx.dll</td>
<td>Kernel library optimized for Intel® Advanced Vector Extensions (Intel® AVX) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_avx2.dll</td>
<td>Kernel library optimized for Intel® Advanced Vector Extensions 2 (Intel® AVX2) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_avx512.dll</td>
<td>Kernel library optimized for Intel® Advanced Vector Extensions 512 (Intel® AVX-512) enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_def.dll</td>
<td>Vector Mathematics (VM)/Vector Statistics (VS)/Data Fitting (DF) part of default kernel</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_mc.dll</td>
<td>VM/VS/DF for Intel® SSSE3 enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_mc2.dll</td>
<td>VM/VS/DF for 45nm Hi-k Intel® Core®-2 and Intel Xeon® processor families</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_mc3.dll</td>
<td>VM/VS/DF for Intel® SSE4.2 enabled processors</td>
<td></td>
</tr>
<tr>
<td>File</td>
<td>Contents</td>
<td>Optional Component</td>
</tr>
<tr>
<td>------------------------------</td>
<td>--------------------------------------------------------------------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>mkl_vml_avx.dll</td>
<td>VM/VS/DF optimized for Intel® AVX enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_avx2.dll</td>
<td>VM/VS/DF optimized for Intel® AVX2 enabled processors</td>
<td></td>
</tr>
<tr>
<td>mkl_vml_avx512.dll</td>
<td>VM/VS/DF optimized for Intel® AVX-512 enabled processors</td>
<td></td>
</tr>
<tr>
<td>libmkl_vml_cmpt.dll</td>
<td>VM/VS/DF library for conditional numerical reproducibility</td>
<td></td>
</tr>
<tr>
<td>libimalloc.dll</td>
<td>Dynamic library to support renaming of memory functions</td>
<td></td>
</tr>
</tbody>
</table>

**Cluster Libraries**

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
<th>Optional Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>mkl_scalapack_lp64.dll</td>
<td>ScaLAPACK routine library supporting the LP64 interface</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_scalapack_ilp64.dll</td>
<td>ScaLAPACK routine library supporting the ILP64 interface</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_cdft_core.dll</td>
<td>Cluster FFT dynamic library</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_lp64.dll</td>
<td>LP64 version of BLACS routines</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_ilp64.dll</td>
<td>ILP64 version of BLACS routines</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_intelmpi_lp64.dll</td>
<td>LP64 version of BLACS routines for Intel® MPI Library</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_intelmpi_ilp64.dll</td>
<td>ILP64 version of BLACS routines for Intel MPI Library</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_mpich2_lp64.dll</td>
<td>LP64 version of BLACS routines for MPICH2 or higher</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_mpich2_ilp64.dll</td>
<td>ILP64 version of BLACS routines for MPICH2 or higher</td>
<td>Cluster support</td>
</tr>
<tr>
<td>mkl_blacs_msmpi_lp64.dll</td>
<td>LP64 version of BLACS routines for Microsoft* MPI</td>
<td>Cluster support</td>
</tr>
<tr>
<td>File</td>
<td>Contents</td>
<td>Optional Component</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>--------------------------------------------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>mkl_blacs_msmpi_ilp64.dll</td>
<td>ILP64 version of BLACS routines for Microsoft* MPI</td>
<td>Cluster support</td>
</tr>
<tr>
<td><strong>Message Catalogs</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1033\mkl_msg.dll</td>
<td>Catalog of Intel® Math Kernel Library (Intel® MKL) messages in English</td>
<td></td>
</tr>
<tr>
<td>1041\mkl_msg.dll</td>
<td>Catalog of Intel® MKL messages in Japanese. Available only if Intel® MKL provides Japanese localization. Please see the Release Notes for this information.</td>
<td></td>
</tr>
</tbody>
</table>
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