The GNU libstdc++ parallel mode: Software Engineering Considerations

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ABSTRACT
The C++ Standard Library implementation provided with the free GNU C++ compiler, libstdc++, provides a “parallel mode” as of version 4.3. Using this mode enables existing serial code to take advantage of many parallelized STL algorithms, an approach to making use of multi-core processors which are now or will soon will be ubiquitous.

This paper describes the goals, the design choices and the software engineering issues discovered during design and implementation, and the results of user testing, presenting possible solutions to outstanding issues. We address standards compliance, influencing algorithm and parameter choices at compile and run time, and the interplay between execution speed, the executable size and the compilation time.

Categories and Subject Descriptors
D.2.2 [Software Engineering]: Design Tools and Techniques—Software libraries; D.1.3 [Programming Techniques]: Concurrent Programming—Parallel Programming

General Terms
Algorithms, Design, Languages, Standardization

Keywords
Parallel Algorithm Libraries

1. INTRODUCTION
Nowadays, nearly all personal computers ship with more than one processor core on a chip. However, the increased computing performance from the underlying hardware is often rendered useless due to insufficiently parallelized software. Thus, the high performance possibilities inherent in multi-CPU hardware is at present largely theoretical.

Parallelizing software is difficult. There exist a wide variety of approaches to extracting parallelism: new languages, language extensions, specialized libraries, specialized compilers. The tools for parallel software engineering are quite primitive as compared to existing tools for traditional serial software development.

In addition, writing properly parallelized programs is hard. Software developers not only have to think about the implementation of threading, but also about how to actually parallelize the algorithms, take into account hardware variance, and then do testing and debugging. This is beyond the abilities of many programmers, and thus costly and error-prone when attempted.

C++ is a very well-established programming language with a rich syntax, supporting both object-oriented and generic programming. It comes with no inherent performance penalty, as do languages requiring a virtual machine, e.g. Java or C#. This makes it a ideal candidate for applications executing compute-intensive algorithms. The standardization of C++ [3] has set a common ground for language and library implementations, which are widely available and often of high quality. The C++ standard is also evolving, as can be seen by the proposed upcoming standard, termed C++0x [2]. The Standard Template Library [13] is part of the existing library specification, and provides many useful generic data structures and algorithms to its users. Since this interface is long-established, well-understood, and widely used, it is a very good way to bring easy-to-use parallel algorithms to the programmer. By utilizing parallelized STL algorithms as black boxes, the programmer can completely abstract from parallel programming, but still benefit from multi-core power. If activated, a usual call to an STL algorithm will cause its parallel execution. So this is a very good example for the concept of encapsulation, too. It even works for existing code, and the parallelization can optionally happen in an incremental way.

The implementation of the parallel mode was begun under the name Multi-Core Standard Template Library (MCSTL) [14] in 2006, followed by its integration into the STL implementation of the free GNU C++ compiler (g++) in 2007. g++ is available for a large number of platforms, including x86, x86_64, IA-64, and Sparc. It provides state-of-the-art optimization.

The parallel mode includes functionality like load balancing embarrassingly parallel work, sorting, merging two or more sorted sequences, random shuffling, partition of a sequence by a pivot, finding single elements or substrings in sequences, prefix sums, and many more. Thus, it enables the programmer to use parallelized standard algorithms as

\[ 1 \text{Both free as beer and free as speech. The compiler is open-source and licensed under the GNU Public License, the libstdc++ under the GPL with runtime exception. That means that even commercial programs may use the library without any having to publish the program’s source code.} \]

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This paper states the goals of libstdc++ parallel mode and shows up the software engineering issues we encountered while developing and integrating it. For many of them, we present resolutions, which might include trade-offs between desirable properties. Some of them cannot be resolved satisfactorily in the current setting. In this case, the paper intends to foster research for development of the C++ language and its standard library. The solutions presented here are propositions, may not be integrated into the current parallel mode code (yet), and are subject to change.

The algorithmic aspects have been considered in [14]. The general performance and scalability of the algorithms is comparable to the ones in the MCSTL, and has been evaluated in that context before. Still, we will discuss performance implications stemming from the fact that a library must be general, thus making specific optimizations difficult.

### 2. SOFTWARE ENGINEERING ASPECTS

Major software-engineering-related goals are:

1. Transparent integration of parallel algorithms, i.e. compile in sequential mode and parallel mode, without changes to the user code. Pragmatic balance between adherence to the C++ standard and parallelization/software engineering benefits.
2. Possibility to choose and tune algorithms, and to adjust the degree of parallelism, in order to achieve best performance.
3. Maintainability of the library code, i.e. little code duplication, much code reuse.
4. Limited increase in compilation time and executable size, with respect to the user application.

#### 2.1 Threading

We chose the OpenMP compiler extension and API in version 2.5 [12] for managing the threads. It has many advantages over strictly library-based approaches like pthread.

- **OpenMP** provides so-called “fork-join” parallelism, which is right what we need for parallelizing single algorithms. The work is divided, the threads start, they re-join, the function call returns.
- **Low-level issues** like thread pooling, and getting information about the environment, e.g. the number of cores, can be left to the OpenMP runtime. The user can configure the runtime environment using standardized procedures.
- **Augmented code also compiles and runs correctly** (though not in parallel) with OpenMP stub support, which can be added to every compiler, easily by the user.
- **OpenMP is platform-independent.** GCC supports OpenMP 2.5 from version 4.2. It is also supported by all major commercial C++ compilers.

In addition to OpenMP, the parallel mode uses atomic operations like **fetch-and-add**, as provided by the GCC’s lower-level libraries. The abstraction layers are visualized in Figure 1.

#### 2.1.1 OpenMP Compliance

A program incorporating algorithms from the libstdc++ parallel mode tries to behave like any other OpenMP-parallelized program. This includes obedience to the environment variables defined in the OpenMP standard. Most important are `OMP_NUM_THREADS` and `OMP_DYNAMIC`, which influence the number of threads. By default, `OMP_NUM_THREADS` is usually set to the number of cores available in the system, `OMP_DYNAMIC` is switched off. Although an OpenMP `parallel` clause takes an argument on the desired number of threads, this value poses an upper limit only. If the dynamic mode is switched on, the runtime can decide to schedule a smaller number of threads to the parallel region. In the end, this means that the number of threads is clear only after the `parallel` statement, i.e. in the parallel region. The code must not make any assumptions about the number of threads beforehand. In many algorithms, this leads to `single` block right after the beginning of the parallel region, in order to distribute the work appropriately. For best performance, the compiler should identify this case, and start the threads only after the `single` block, having fixed the number of threads in advance, and thus returning the right value from `omp_get_num_threads`.

The environment variable `OMP_SCHEDULER` is less important, since it only influences OpenMP `parallel for` loops, which are not used by the parallel mode at the moment.

Nested parallelism is used by algorithms, namely the parallel quicksort variants. It is a classic example for parallel divide-and-conquer. Conceptually, the threads partition the sequence into two parts, in parallel. After that, the threads are split into two groups corresponding to the sizes of the sub-sequences. When only one thread remains in a group, it solves the problem on its own, possibly helping out slower threads via work-stealing.

In the actual implementation, the threads are not split, but newly started, relying on the OpenMP runtime’s thread pool in terms of efficiency. The desired effect is achieved by having a `parallel` region with two `sections`. In each of this sections, an appropriate number of worker threads are spawned for the parallel partitioning. After that, the sorting step is called recursively, which leads to nested parallelism.
There is a termination condition that stops the spawning of new threads such that the total number of threads never exceeds a certain maximum number, determined by calling `omp_get_max_threads` at the start of the algorithm. The runtime deciding to execute two sections one after the other does not threaten correctness, either.

If the user switches off nested parallelism using the `OMP_NESTED` environment variable, this scheme is destroyed. The algorithm will still run correctly, but its speedup will be limited to 2.

Of course, an equivalent algorithm could be implemented using a single parallel region, by scheduling all arising work “manually”. However, this would imply the following disadvantages:

- The parallel partitioning step could not be done by the already existing parallelized `partition` call, because the threads are already in existence. A less elegant solution, minimizing code duplication, would be to tear apart the partition algorithm, and to double-use its parts in both the quicksort and the `partition` implementation.

- Threads of a group have to barrier synchronize once in a while, which can imply (OS-based) waiting. However, the OpenMP `barrier` statement takes all threads in a `team` into account, not only the ones in the algorithm-specified and data-dependent group. Solving this case inside the OpenMP specification could only be done by using `omp_set_lock` and `omp_get_lock`, which is beyond their intended purpose. OpenMP lock functionality does not support the monitor concept where multiple threads can be notified at once. Thus, each thread would have to wait for its predecessor thread to finish, with respect to a predetermined order. This adds up the inefficiencies.

A future elegant resolution without the problems noted here, using OpenMP 3.0, is sketched in Section 2.11.

### 2.2 Activation of the Parallel Mode

The use of the parallel mode STL should be optional, i.e. by default, it is deactivated by the compiler. If the programmer wants to use the parallel mode, there should be two options:

1. Activate the parallel mode by default, i.e. all STL algorithms that have parallel implementations available, will have parallel code included into the program.

2. Activate the parallel mode on request, i.e. specific algorithm calls can be annotated by the programmer to call the parallel version of the algorithm.

By declaring the nested namespace using a `inline namespace _parallel;` declaration\(^3\), the contents of namespace `_parallel` are transparently imported into the surrounding one, superseding the already existing symbols there. This procedure is called `namespace association` [11]. The parallel implementations from namespace `_gnu_parallel` are imported into the unaltered namespace `std`, the parallel calls thereby hiding the sequential implementations\(^3\). The sequential versions are not completely inaccessible in this case, it is just that the imported functions are chosen by default. This is just what we want for the first case. If the programmer wants to force sequential execution at a certain spot, he just adds `_gnu_parallel::sequential_tag()` behind all other call arguments.

In the second case, the programmer just has to prepend the namespace qualifier `_gnu_parallel::` to his call, to switch to the parallel version explicitly.

### 2.3 Sequence Access through Iterators

Most STL algorithms take one or more sequences as their main argument(s). To abstract from the underlying representation, the iterator concept is used. A sequence is usually specified by passing an iterator referencing the first element `begin()`, and an iterator referencing a virtual element behind the last element `end()`. This allows a consistent treatment of arrays, regarding pointers as iterators. Also, different iterator types or instantiations can access a single sequence in different ways, e.g. selecting a range, advancing with a stride, going backwards. On the one hand side, this adds a lot of flexibility.

On the other hand side, information gets lost when passing iterators. The best example for that is the length of a linked list. A linked list container might implement the size query efficiently, just by book keeping the number of inserted and deleted elements. When an algorithm is passed the corresponding `begin()` and `end()` iterators, however, this information is lost, since the iterators do not include a reference to the underlying container. Since the iterators are not random-access iterators, the distance between them cannot be computed efficiently. The only way to find out the length of the sequence is to traverse the whole sequence, taking sequential linear time, and impossible to parallelize.

Since all parallel mode algorithms are `data-parallel`, the input must be split into parts, the splitting positions being scattered along the whole sequence. This can only be done in sublinear time, if random access to the elements is granted. Hence, parallel execution seems to be limited to random-accessible sequences. We have given thought on how to workaround this problem with using as little sequential computation as possible [7], but this is a secondary choice.

The decision on whether random access is possible happens at compile time, using partial template specialization. For each used iterator, an appropriate specialization of `std::iterator_traits` must be available, as requested for all iterators by the standard [3, Section 24.3.3].

### 2.4 Code Reuse

Parallel algorithms, in general, are far more complicated than sequential ones. For example, the one-liner describing `for_each` explodes into a parallel version of many hundred lines, implementing complex load balancing mechanisms like work-stealing.

On the one hand, this fact makes the usage of a library even more valuable, since you do not have to write that code yourself. On the other hand, the library itself is threatened by hard-to-maintain complexity. For this reason,

\(^3\)A similar construction has already existed in the libstdc++, namely for the `debug mode`.\n
the library programmers must make efforts to keep the code as maintainable as possible, without diminishing benefits. Two important aspects in that are fostering code reuse, and avoiding code duplication.

Code duplication is particularly tempting for the STL, since it contains many algorithm calls that act similarly. for_each and (unary) transform, for example, only differ in the fact that the functor passed to transform is allowed (even supposed) to modify its input in-place. There are several other similar calls, basically all of them being "embarrassingly parallel", traversing one or two input sequences. One could write the code for for_each, copy and paste it several times, and change some little details. This is feasible for the sequential versions, but unmaintainable for the parallel ones. A bug fix or an algorithmic improvement would have to be applied in many places, making the process extremely error-prone.

In particular the algorithmic portions, must be reused already by the library itself. For a family of algorithms, a most general version of the parallel algorithm is factored out, and subsequently called by all members of the family. The surplus arguments must be filled with dummies, e.g. noop-functors and dummy iterators. Similar action is taken for default functors. For example, the sequential implementation provides two completely independent implementations for both versions of accumulate, one having explicitly specified the addition functor, the other taking the default operator+ for the respective type. For the complex parallel version, the implicit version must be redirected to the explicit one by filling in an equivalent functor as soon as possible (see Figure 3), to avoid code duplication. We have to rely on the compiler’s ability to inline simple operators, and to optimize away empty functions. Otherwise, the performance penalty will be considerable. Compiling without optimizations will therefore impede the parallel implementation to a greater extent than the sequential one.

There are some algorithms that are a sensible extensions of the standard library. One of them is the generalization of merge to accept multiple sequences, instead of only two. This is internally used for the mergesort implementation, therefore called multiway mergesort. Of course, multiple sequences can be merged together one after the other, but this is less efficient, not in asymptotic running time, but due to additional accesses to memory. So we pass on this functionality to the programmer under the name multiway_merge.

Other functions are helpers for exploiting parallelism in higher levels. This includes the simple, but error-prone function for splitting a range into parts of about the same size (plus/minus 1). Much more complex is the routine for selecting the element of a certain global rank, from several sorted sequences. Both routines are used intensively by the parallel mode internals: it is only natural to make them accessible to the programmer. One approach would be to document the calls in <gnu_parallel>, and attempt to then standardize most-used functionality in namespace std for some future version of C++.

2.6 Sequential Performance

Since parallelization is about performance, the question arises whether the existing sequential algorithms are sufficiently optimized. An asymptotic penalty in running time is accepted for integer sorting, since the STL provides only comparison-based sorting, taking \( O(n \log n) \) time. Specialized integer sorting algorithms take only \( O(n) \) time for this task. We could specialize the sort routine for all kinds of integer elements, to support at least ascending and descending order. However, the library user usually wants to sort a more complex object including an integer key. Thus, the interface would have to be changed quite radically. Not a comparison functor would have to be passed, but a functor extracting the key from the element. This could lead to a syntax similar to the upcoming hash maps.

2.5 Extensions to the Standard Library

2.7 Algorithm Variants
#include <vector>
#include <algorithm>

int main() {
    std::vector<int> vi(100000000);
    std::sort(vi.begin(), vi.end() TAG); return 0;
}

sort(begin, end, comp=std::less<int>())
parallel_sort(begin, end, comp, stable=false)
1. parallel_sort_mwms(..., stable);
2. parallel_sort_qs(...);
3. parallel_sort_qsb(...);
    respective OpenMP-parallelized code

Figure 3: Minimal sorting example, and call stack for “runtime variant choice”, i.e. TAG="" with the three possible runtime outcomes.

For some functions, the parallel mode offers multiple algorithmic variants, in order to allow best execution speed in a variety of circumstances. Particularly sophisticated are the three variants for sorting. This may sound like a lot, but note that the sequential implementation of stable_sort also has two completely independent versions.

The parallel mode features two kinds of parallel quicksort, unbalanced and balanced, as well as a the multiway mergesort. The quicksort variants work completely in place, and are quite small in executable size (see Table 1). On the downside, they do not support stable sort, and they require nested parallel regions.

Multivay mergesort provides stability, and does not use nested parallelism. However, it requires a temporary copy of the data. Experiments show that it scales better down in terms of running time, i.e. for small inputs.

While mergesort can give stronger theoretical guarantees on the running time, the quicksort variants may not scale at all in the unlikely theoretical worst case. In practice, the balanced variant can better handle the case of cores being unexpectedly unavailable because of other processes.

### 2.8 Code and Binary Size, Compilation Time

Generic programming trades off executable size for optimal adaptation to and integration of the specified data type, resulting in best performance. Instantiating an algorithm for each data type separately allows high-level programming without any run-time overhead, which would be implied by the use of virtual functions or similar run-time decision-making. The increase in executable size that comes with generic programming is considered an acceptable trade off. If code size is the most important factor for a particular use-case, you can still derive all used data types from a base class with virtualized functions and instantiate the algorithm for the base class.

For the parallel mode implementation in particular, the algorithms are more complex and multiplied by variants, and thus the problem gets more important. As a result, the code size for the algorithms can grow up to a factor of 4 compared to the sequential code. This is shown in Table 1, for the program in Figure 3:

For TAG, the corresponding tags are inserted, in order to choose the algorithm variant. The code was compiled using the options `-D3 -g0 -fopenmp -D_GLIBCXX_PARALLEL -DNDEBUG` with a prerelease version of GCC 4.3, running on a x86_64 machine.

The compilation time also differs, it approximately scales with the size of the resulting executable. This is mostly due to the optimization passes, not the parsing, as can be seen by comparing the timings for the debug mode.

As a conclusion from these measurements, it can be stated that including all variants of an algorithm for a run-time choice is best avoided. Instead, the programmer should be able to choose a specific algorithm variant at compile time, and the default should include only the parallel algorithm forming the best choice.

### 2.9 C++ Standard Compliance

Strict 100% compliance to the C++ standard makes parallelization virtually impossible. This is because the standard was designed for serial use, without having parallel processing in mind at all.

The algorithm descriptions often do not specify only a input/output relation, but instead provides code examples for which the operation is “equivalent.” This implies the order in which the elements are processed (say in `for_each`), and thus ultimately forbids parallelization.

### 2.9.1 Functors

The library calls take many kinds of encapsulated user code as arguments.

- Explicit functors to process/transform an element, making up the main functionality of the algorithm. The functor object might contain “state” information. Examples: functors for `for_each`
- Implicit methods and functions, inherent to the value type. Examples: Overloaded (converting) constructors and assignment operators
- Implicit methods and functions, inherent to the iterator type. Examples: Increment operator (`operator++`), dereference operator (`operator*`), difference operator (`operator-`), default constructor
- Stateful helper functions. Example: random number generator for `random_shuffle`.
- (Hopefully) stateless helper functions. Example: Comparison functors, e.g. used by `sort`.

All of this user code might be able to interact with the environment in an non-thread-safe way through side effects.

<table>
<thead>
<tr>
<th>Algorithm Variant(s)</th>
<th>Size</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequential</td>
<td>15479</td>
<td>0.74</td>
</tr>
<tr>
<td>Quicksort</td>
<td>22387</td>
<td>1.49</td>
</tr>
<tr>
<td>Balanced Quicksort</td>
<td>26989</td>
<td>1.84</td>
</tr>
<tr>
<td>Multiway-Mergesort Sampling</td>
<td>36002</td>
<td>3.49</td>
</tr>
<tr>
<td>Default</td>
<td>41229</td>
<td>4.68</td>
</tr>
<tr>
<td>(Multiway-Mergesort Exact)</td>
<td>41237</td>
<td>4.78</td>
</tr>
<tr>
<td>Multiway-Mergesort Exact</td>
<td>46003</td>
<td>5.48</td>
</tr>
<tr>
<td>All Parallel Variants (run-time choice)</td>
<td>61543</td>
<td>6.50</td>
</tr>
</tbody>
</table>

Table 1: Executable sizes (in bytes) and compilation time (in seconds) for different algorithms variants.
These are most legitimate for the first class of functors. However, for some routines, the standard states that they “shall not have any side effects”, namely transform [3, Section 25.2.3] and the numerical algorithms accumulate, inner_product, partial_sum, and adjacent_difference.

For the other classes, they usually imply a bad programming style, or an abuse of functionality.

The extensive code re-use, justified in Section 2.4, yields another detail problem. Functors have to be passed to subroutines (analogous to the call stack in Figure 3), and the question arises whether this should by done by value or by reference. The common prototypes of the algorithm calls take the functors by value. This prohibits returning any “state” to the caller, which is good news. To avoid copying the same thing multiple times, one should pass on references to subroutines.

Another hitch stemming from the more complex parallel algorithms is the use of user data types with unfortunate properties. For example, many algorithms have to copy elements from the input to some temporary storage. Unfortunately, the library may not assume that the given data type is default-constructable, so the temporary storage cannot be easily allocated. The algorithms never produce “new” elements, so we can always circumvent this problem. One option is to use pointers to the existing elements, the other is to reserve raw memory and to construct the element using placement new, calling the copy constructor. In the latter case, you have to be extremely careful in order not to construct multiple objects in the same place without calling the destructor for the former ones.

Determining the return type of a functor is a persistent, but minor problem. The most elegant way for the programmer to declare a functor, is to (indirectly) derive from a standard base class like unary_function or binary_function. These base classes take the argument type and the return type as template parameters, so they can be read out from the type members (first)_argument_type, second_argument_type, and result_type. However, it is more realistic not to rely on the types being defined appropriately. For the return type, there is an alternative, namely applying the typeof keyword to a “fake” function call. This function call is in fact never executed, but just serves to find out the return type: typedef typeof(functor(input)) return_type;

The return type is needed for inner_product, where the type of the intermediate values (after multiplying) need not coincide with the return type. Example:

```c++
struct A;

bool operator*(const A& a1, const A& a2)
{ /*... */ }

void foo()
{
  A* S1 = new A[100], * S2 = new A[100];
  double c =
  std::inner_product(a, a+100, b, 0);
}
```

The return type bool of operator* is needed in order to pass on a regular functor to the “embarrassingly parallel” infrastructure, instead of just using the + and * operators.

### 2.9.2 Exception Safety

Two things that avoid each other like the plague are parallelism and exceptions. There is no theoretical reason for this, rather a lack of well-specified behavior in existing C++ implementations.

The problem surfaces when a routine in a thread throws an exception which is not caught inside the thread and propagates up the stack.

When the parallelization is done explicitly, this situation can be avoided by carefully specifying throw and catch regions within each thread of parallel execution such that an exception never leaves the originating thread. The consequences to program execution can then be handled by checking for error conditions, e.g. forcing the other threads to terminate, or somehow sensibly resuming execution.

For a library which takes functors, such a specific solution is not applicable, and the search for a more robust solution is an area of active research. Some general restrictions should apply, as follows. First, the algorithms themselves should not throw any exceptions. This is easy to achieve since “wrong” input is not possible, in most cases. Although some algorithms rely on the given input sequences being sorted. However, even the sequential versions don not check for this precondition, and as a result may deliver wrong results, not terminate, or crash the whole program. For all algorithms taking a comparison functor, inconsistent return values (failing to be a strict weak ordering) can lead to undefined behavior, just as in the sequential case.

Second, exceptions have to be expected only from user-defined code, as listed in Section 2.9.1. But left as an open issue is the problem of a sensible reaction to an exception being thrown. Rethrowing the exception after the clean termination of the algorithm is a reasonable choice, as is stopping all involved threads as soon as possible. If more than one thread throws before the execution can be stopped, one of the exceptions would have to be resolved before the other. But which? Unfortunately, the current C++ standard does not allow this kind of reaction as two active exceptions leads to a call of std::terminate. A catch clause can either specify an exception type to catch, or catch all exceptions, as marked by an ellipsis (...). In the first case, the type of the exception is known, so it could be rethrown. In the second case, the exception object is unknown and one can only attempt to deduce it via nested try blocks containing rethrows. Sadly, the set of all exceptions to expect are not known to the library. Throwing some general exception in place of the original one would be an work-around.

The best way to handle this would be to extend OpenMP with definite exception handling semantics. As it stands, throwing exceptions over parallel region boundaries is explicitly specified as forbidden.

Little has been published about solving this problem. One paper [8] only considers terminating the program in a standard-compliant way, not giving any thoughts to resuming it sensibly. To our knowledge, the only proposal for handling exceptions in the context of OpenMP is a paper

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4This is an area that is not detailed in the current C++ standard, and still under development in C++0x: some C++ implementations allow a “per-thread” exception handler, and others do not. Lack of portability has hampered experimentation in this area.

5This is particularly likely to happen for out-of-memory-related exceptions.

6A proposal for the upcoming standard could come to the aid of all this, see Section 2.10
proposing integration of OpenMP into Java [9, Section 4.3]. Their suggestion is that after the occurrence of an exception, the threads should terminate as soon as possible. To achieve this, the catching thread sets a cancellation flag, which is read out by the other threads at certain cancellation points, namely barrier, critical, parallel, and all OpenMP work-sharing directives. A similar approach for C++ could lead to more effective exception handling in the parallel mode.

2.9.3 Tag Parameters
In order to allow compile time algorithm choices such as forcing sequential execution, the function prototypes must be augmented by additional “tag” parameters. This makes some of the standard compliance tests fail because of the augmented method signature. For user code, this should not make any difference, since all standard-compliant calls will still succeed.

2.9.4 Conclusion on Standard Compliance
So far, the libstdc++ parallel mode works with “reasonably” programmed code. This includes assumptions as side-effect-safe and exception-free functors passed to the parallelized algorithms.

Others are inherent to parallel computing, like side-effect issues and exceptions. Those could be mitigated by providing thread-safe containers, as provided by the Intel Threading Building Blocks [1].

2.10 Benefits from Upcoming C++ Standard
The upcoming next version of the C++ standard will provide functionality that comes in particularly handy for the parallel mode.

The easy availability and standardization of thread-local storage could simplify the parallel mode code. The introduction of powerful atomic operations will reduce the amount of compiler-specific calls. A proposal [5] on how to carry exceptions between threads has been posted recently. By calling rethrow exception, an exception originally thrown in another thread, can be “repeated” in the current thread. This would allow exception handling in the parallel mode in an explicit way, i.e. the algorithms would catch all user-induced exceptions, terminate properly, and rethrow the exception after the parallel region.

Concepts and static assertions simplify checking for preconditions, e.g. an iterator being random-accessible. Lambda functions allow for more elegant definition of functors, thereby making the usage of STL algorithms easier and more intuitive. The auto feature will provide an elegant, minimal, and complete solution for the return type issue.

So-called rvalue references allow to avoid unnecessary copying of (potentially complex) objects. Instead, they can be “moved” explicitly, implying that the data must not be accessed in the original location any longer. By providing a specific move constructor, the programmer can thereby make moving much more efficient than the combination of copy construction/assignment and deletion. This helps the STL algorithms in general, because many of them actually only move data, e.g. sort and random_shuffle. The parallel mode benefits because copying data loads the memory connection, a common bottleneck for parallel speedup.

2.11 Benefits from Upcoming OpenMP

Although the OpenMP standard version 3.0 is not yet final, some of the proposed innovations by the draft look promising.

The standard draft specifies the parallel for loop to work on C++ random access iterators in addition to integer loop variables. This supplements the for_each functionality. One drawback is that no scheduling is required that provides a worst-case guarantee on running time.

A major addition is the task construct, which allows (recursive) asynchronous processing of code blocks, scheduled by the OpenMP runtime. This could greatly simplify the code for recursive routines like the quicksort variants, and avoid the nested parallelism drawback. The use of the platform-specific yield call could also be made obsolete by this construct. The yield call proposes to the OS to give control to some other thread, and is usually called when out of work, in order to avoid other threads to wait unnecessarily long. It is mostly needed when there are more threads scheduled than cores (currently) available.

A publication actually proposed to include an equivalent directive into OpenMP [16]. In the OpenMP 3.0 draft, nothing new is said about exception handling.

2.12 Testing
Rigorous testing of the libstdc++ parallel mode would cover many aspects in an in-depth manner: rigorous correctness, conformance to C++ standards, performance of executables, compilation time and binary size to name a few.

Testing procedures to date have concentrated on conformance and performance.

For conformance to ISO C++ specifications, the test suite accompanying the libstdc++ is used. This test suite is comprised of over three thousand individual test cases, and is run twice: once for sequential code and again for parallel code. Over 99% of the tests succeed in parallel mode, with failures explained by Section 2.9.3.

Coverage could be improved. The functional and conformance checks are often limited to very simple, hand-written examples covering only specific inputs. This is not sufficient for the parallel implementation, since many of its subroutines have fallbacks for very small inputs, e.g. a sequence length smaller than the desired thread number. Larger inputs could be generated on the fly, and the results compared to the sequential runs to assess correctness.

The confidence in the correctness of the algorithms are based on the test program available from the original MCSTL, so far. Integrating this program as a whole into the libstdc++ test suite is difficult because of the quite different paradigms used.

2.13 Performance Aspects
To what extent should an implementation of a general purpose standard library should aim for best execution speed? On the one hand, such an implementation may be used by very many people, which makes optimization worthwhile. On the other hand, such a library must be very robust and general, which impedes finely-tuned optimization. By definition, hand-written code can always be as least as good as a library because nobody keeps you from doing the same thing as the library does. However, the genericity is gone, and thus it will be too much of an effort to optimize all algorithm instantiations of a program.
to this extent. Such solutions tend to be fragile, and require constant maintenance.

The STL implementations are mostly asymptotically efficient, losing only a constant factor to an optimized version. Things are different for parallelization. Under the reasonable assumption that the number of cores in a computer will continue to increase, a sequential implementation of the STL will lose a growing factor compared to the parallelized one. This is what makes parallelized solutions so essential.

In terms of a widely-available highly-used algorithm and data structures library, there is actually not much competition to the STL. LEDA [10] covers much more functionality, but lags behind the development of C++, and as a result does not fully exploit its generic programming capabilities. In addition, an excessive usage of virtual functions leads to bad performance for many cases. A very nice example for a library that pulls out all the stops of C++, is the Computational Geometry Algorithms Library (CGAL) [6]. However, it is limited to geometry and internally re-uses much of the STL. The Boost libraries try to extend the standard library with additional functionality, and many are prospected for inclusion into upcoming versions of the standard.

2.14 Heuristics, Tuning

Even if parallel code is included into the executable, this does not mean that the algorithm is always executed in parallel. Heuristic considerations might cause a fallback to the sequential version at runtime, the most obvious case being that there is only one processor core available. In addition, the input size and the data type are taken into account when deciding whether a function call should be executed in parallel. Many of the heuristics and thresholds can be configured at run-time.

Also, in several cases, tuning parameters allow to trade off granularity of load-balancing against synchronization overheads. Load-balancing work in a fine-grained way shows worst-case performance, but can add too much synchronization overhead to be worthwhile.

2.15 Interaction with Other Libraries

We have integrated the libstdc++ parallel mode with another library akin, namely the STXXL [4]. The STXXL provides functionality alike the STL with respect to external memory, i.e. data on hard disks. Since the library can load several parallel disks very efficiently, it is often actually compute-bound. Using the parallel mode for the internal memory operations results in significant speedups for various applications.

Another natural candidate for using the parallel mode would be CGAL.

3. CONCLUSION AND FUTURE WORK

In this paper, we have presented the transparent integration of parallel algorithm implementations into the libstdc++. We have proposed solutions for most of the arisen issues, many of them already implemented or in the process of being implemented. For the others, we show starting points.

One of the hardest remaining problems is the decision of whether to switch to parallel execution or not. This is crucial for the resulting program performance in the worst case, since the parallelization overhead cancels out speedups for “to easy” problems, i.e. algorithm calls that have to little or too local data for communication of the data being worthwhile. By testing and profiling, the application programmer can discover break-even points and tuning parameters for a specific algorithm on a specific data type, a specific functors, and so on. However, this is tedious and does not lend to deployment on several platforms at the same time with the same code. As a result, developing methods for auto-tuning the algorithms are underway. Even with the noted deficiencies, the parallel mode user can already work at a higher level of abstraction and performance.

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4. REFERENCES