

# The Multithreaded version of FORM

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## Abstract

We present TFORM, the version of the symbolic manipulation system FORM that can make simultaneous use of several processors in a shared memory architecture. The implementation uses Posix threads, also called pthreads, and is therefore easily portable between various operating systems. Most existing FORM programs will be able to take advantage of the increased processing power, without the need for modifications. In some cases some minor additions may be needed. For a computer with two processors a typical improvement factor in the running time is 1.7 when compared to the traditional version of FORM. In the case of computers with 4 processors a typical improvement factor in the execution time is slightly above 3.

# 1 Introduction

The symbolic manipulation system FORM [1] has been available for more than 17 years. It has been used for many calculations that involve large intermediate expressions, many of them in quantum field theory. It is estimated that the existence of FORM has caused the advance of calculations in field theory by one order in perturbation theory. None of the more spectacular calculations of refs [2–13] would have been possible with other available systems.

The advantages of FORM over other systems are its capability to handle very large expressions and its speed in processing them. Part of the advances in the handling of these large expressions can of course be attributed to the steady increase in the power of computers. Because the calculations that are undertaken require more and more computer power it has been noticed that the use of more than one processor at the same time can lead to great benefits in a system like FORM. The first attempt took place in 1991 with the special computer that had been constructed at FNAL. This experiment was however not continued due to problems with access to this computer from outside. It led however to a constant awareness of what was needed inside FORM to keep the possibility of parallelization and hence its internal structure was kept more or less compatible with these needs.

In the late nineties a joint project was started with the University of Karlsruhe for adapting FORM to run on special computers with several processors. In the beginning this led mostly to a big struggle with the hardware and the system software, but when the hardware and the system libraries became more stable good results became available. This has led to the program ParFORM which uses most of the FORM sources with some extra code in addition. It has been described in the literature [14–18] and several calculations have greatly benefited from it. Its main drawback at the moment is that it runs most effectively on machines that are rather expensive. Most people however may have access to simpler computers with more than a single processor. It is nowadays rather common to have computers with two or four processors using a shared memory model. It becomes even more relevant because most leading vendors switch to dual- and multicore processors. Hence it was judged important to create a version of FORM that can make efficient use of such systems. Whereas ParFORM uses a message passing protocol named MPI<sup>1</sup> which is quite good for systems that have processors with separated memory or consist of networks of computers, computers with a shared memory can work with a more efficient communication. The best method here seems the use of Posix<sup>2</sup> threads or pthreads. A thread is a semi-process, that has its own stack, and executes a given piece of code. Unlike a real process, the thread normally shares its memory with other threads. Pthreads adhere to strict standards and the libraries for them are widely available. These threads can also make more efficient use of the available resources as the number of threads is not restricted to the number of available processors. Hence if some threads have to wait for data, the processors that become available can give attention to other threads. The above considerations have led to the creation of TFORM, the multithreaded version of FORM,

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<sup>1</sup>More about it can be found at <http://www.mpi-forum.org/>

<sup>2</sup>Posix, "a Portable Operating System Interface for uniX" is the collective name for a family of related standards.

which is described below.

When a symbolic system has to operate in a parallel mode one has to worry about a number of things. The most important is whether the components of an expression can be treated individually. In the case of FORM this can be done, because FORM allows only so-called local operations (operations that act on only a single term at a time). Another problem might be a central variable administration. It should be independent of the order in which the terms are processed.

Once the above has been established one has to design strategies for those parts of the processing in which information needs to be exchanged between the processors. In general there are moments in which expressions need to be brought to a unique form. This means that when the contents of an expression are spread out over the processors, these processors will have to send pieces of the expression to each other. To avoid total chaos there should be a single processor that determines which processors will continue processing which parts of the expression. Normally this will mean that temporarily each expression will be under the complete control of a single processor. This constitutes a bottleneck in the processing. In the case of FORM this bottleneck consists of two phases: The first is the reading of the expression from the disk and distributing the terms over the 'workers'. The second bottleneck is the final stage of the processing of the terms of an expression in which the terms are sorted and put together. This sorting is done by merging. In its very last stage it is a single processor that will have to do the final merge and write the resulting terms in order to the disk. Much attention has to be given to the efficiency of the code for these bottlenecks as it puts a limit on the improvement of the combined running time.

There are some variables in FORM that are an intermediate between private and common variables. Common variables are variables of which there is only a single value for every process to use. Private variables are variables of which each process has its own copy, or of which each term generates its own value. The variables we refer to are called dollar-variables because their names start with the dollar character. They can take values and their values can be used both during the common phases of a program like preprocessing and compilation when only the master process is active, and during the execution phases of the program when things are done on a term by term basis. In the latter case, when we have a central/common administration, their value may depend on the order in which the terms are treated. If each processor has its own administration for these variables in the end each processor may have a different value, and the question arises as to which value we need for the common value. If this problem cannot be resolved the corresponding part of the program cannot be parallelized. New statements have been implemented to help FORM with such conflicts and to minimize the number of cases in which it has to decide to run a given module in 'sequential' mode. Internally this is done differently in ParFORM (each processor has its own administration and the resolution of the conflict is at the end of the processing of an expression) than in TFORM (one central administration with the resolution of conflicts at the moment a variable obtains a value). At the user level however this difference shouldn't be noticeable and both will act identically.

Special attention has to be given to simultaneous file access. When several processors need part of an expression which has been stored on disk, they may have to wait for each

other. Hence it is best that each processor has its own caching system. This will minimize waiting time. Another problem occurs in certain systems with the simultaneous use of the disk by the processors in the later stages of the sorting of a large expression. We have encountered cases in which the ensuing traffic jams actually made a program on 4 processors slower than when it was run on a single processor.

It is rather important that programs need as few modifications as possible to be able to use the benefits of running on several processors simultaneously. We think we have been able to do this with FORM. This is illustrated with a number of test programs that were originally developed for running on a single processor (and optimized for it). We try to run them unchanged and then measure the improvement in running time (wall clock time on a computer that isn't engaged in any other major tasks). The actual improvement depends of course on the programming style and particularly on what is the ratio of the time spent by the algebraic operations of the statements inside the modules versus the sorting at the end of the module. The intensive use of external files may influence it also. Finally, work done during the compilation and the printing stages is done only inside a single processor and hence lowers the efficiency as well.

In the second and third sections we will discuss the issues that needed addressing inside the C-code of FORM in order to allow it to run several threads simultaneously. In the second section we give attention to the modifications of the existing code that were needed before the actual parallelization could be attempted. Then in the third section we pass on to the FORM specific pieces of code that had to be designed to make the multi-threaded running efficient. In the fourth section we test the resulting program called TFORM with a number of existing programs that are available in the FORM distribution.

## 2 Cleaning up the internals of the FORM sources

The first problem in designing multi-threaded programs is making sure that all routines are reentrant or thread-safe. This means that several instances of the same routine should be capable of running at the same time without unintentionally influencing each other. Hence C-code like

```
static int ScratchArray[100];
int Multiply(int *term1,int *term2)
{
/*
    use ScratchArray to temporarily store pieces of the terms
*/
}
```

will usually cause disasters as all instances of the routine Multiply will write into the same array at the same time. Therefore we need a very strict separation in FORM of which variables are common variables and which variables are private variables, with which we mean variables of which each thread needs its own copy. Usually this is done by storing all private variables in the stack, but in the case of FORM this would involve some very big arrays and not all operating systems may be prepared to provide that much

stack space. In addition it would require much restructuring inside FORM. Fortunately the organization of the variables in FORM was already such that there was another easy mechanism to make this separation and the code didn't have to be modified very much.

All common variables in the sequential version of FORM are part of substructures of a single common data structure named A. This was done originally to allow compilers to use offsets to the contents of a single address register to refer to all common data. At the time it made FORM about 10% faster. These substructures were ordered by type of use, like the substructure P containing variables that are used/set by the preprocessor, the substructure C containing the variables that concern the compiler, the substructure N containing scratch variables used by the various routines to communicate with each other during running, etc. Rather than referring to these structures as A.P or A.N there are macro's and A.N is referred to by the macro AN. All that had to be done was to define a new type of data structure and move all substructures (these were the N, R and T substructures; all others remained inside the A structure) of which the different threads need their own copy to this structure. Each thread will allocate its own copy of this structure and we define a new data type B which is a pointer to this private structure. After this we redefine the macro's AN, AR and AT into B->N, B->R and B->T and once each routine knows from which thread it is running and hence knows the value it needs for this variable B, the rest of the routine needs no further changes. Because all parameter fields to the routines are already run by macro's (originally this was to facilitate the differences between ANSI-C and non-ANSI-C), it was also easy to redefine some macro's and automatically pass the pointer B to routines that need it.

The above changes affect nearly all the code and make the program slightly slower on average although we have met rare cases in which TFORM on a single processor is a little bit faster than the sequential version with its original definitions of the macro's.

The next changes concern the reading of files. In the sequential version it was possible to position a file in one routine and read it in another. One could also assume that a file was still at the position at which it was left after the previous access. In a parallel environment this is no longer possible. Hence one needs to use locks (for which one can use so-called mutex variables). But because one wants to lock the access to a file for as short a time as possible, it is important to have the positioning of the file and the access to it as close together as possible. This required a number of modifications and was a rather annoying source of errors. The caching system had to be adapted as well to allow each processor to have its own cache, even though the file is opened only once. The way the system works now is that if an expression fits completely inside memory and hence lies completely inside the cache<sup>3</sup> of the master processor, all worker processors read from the cache of the master, but they have their own set of pointers to from where in the cache they are reading. If the expression is too big for the cache of the master and is actually residing on disk, all workers will use their own cache of which the size is  $1/N$  times the size of the cache of the master with  $N$  the number of worker threads. These worker caches are in addition to the cache of the master and hence in total we need twice the size of the cache of the master. It is hard to improve on this as the master also needs the cache for reading the input and distributing the terms over the workers. Trying to improve on this would

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<sup>3</sup>The size of such a cache is determined by the setup variable ScratchSize.

add much complexity to the code.

The next file problem comes with the stored expressions. It is somewhat related to the relocation problem when the executable code of a program is being loaded into the memory, just before running. In the sequential version expressions are stored with their symbol table. This may tell that for instance the variable  $x$  is represented by the symbol number 5 inside the stored expression. When the stored expression is used, its symbol table is compared with the table of active objects and the variable  $x$  may either not be in it yet or may have a different number. If necessary, the variable  $x$  will be entered in the symbol tables. In any case, a renumbering table has to be made because the symbol number of  $x$  may now be different, for instance 12 rather than the 5 it has in the stored version. Originally this would be done during the execution of the program at the moment an expression would be needed. In a parallel environment however different terms may use different expressions. This means that the variables may not be added in the same order when several processors are trying to do this simultaneously. Hence it should be clear that when several threads want to do this at the same time we need either a sophisticated system of locks around the common variable administration, or we can make errors. The solution selected was to let the compiler (which runs on the single master thread) read the symbol tables of the expressions that *might* be used. This avoids conflicts in the numbering without having to resort to locks. On the other hand, if an expression is not used because during running the control flow avoids the statement in which it would be used, its symbols are read nevertheless. This could potentially result in wasteful use of space in the name tables. It was however considered the lesser evil.

For stored expressions multiple caching is also very important. A new caching system was implemented of which the sequential version of FORM also benefits. Therefore one may notice that, also in the sequential version of FORM, programs that make intensive use of stored expressions will execute faster now.

The final cleanup of the code concerned the pattern matching where use was made of some common variables in a very intransparent way. These had to be transferred to the private data space.

The above cleanup concerns all versions of FORM. Hence the moving of the positioning statements of files have their effect on the sequential version as well. By working this way it is much easier to maintain FORM as basically there is only a single version of the source code. The setting of just one parameter controls whether the compiler will produce the sequential version or the multithreaded version.

### 3 The parallelization

For the parallelization we use the conventions of the Posix threads, shortly called pthreads. These are implemented by means of the pthread library, which is available on all modern UNIX systems. For the definition of the concepts and their use we used ref. [19]. The model we use consists of the startup of one master thread and  $N$  worker threads in a so-called thread pool. This seems most efficient on a computer with  $N$  processors. Often when the master has to do work, at least one of the workers is idle and conversely when all workers are occupied, often the master has very little to do. We will show in the tests

that indeed this works well.

The workers are started only once at the startup of TFORM. They allocate their own private memory inside a single data structure of which the address (called B as explained above) is stored in a common array from which the address can be recovered when the number of the worker is known. On computers on which the threads are moved between the processors in a semi-random fashion, the individual allocations, rather than a single common allocation that is parcelled up afterwards, leads to no benefits, but it allows to create a coherency between workers and memory on those computers for which it does make a difference. Currently nothing is done with this.

Next all private arrays for caching and sorting are allocated by the workers and the master. After this the workers go to sleep. The master waits till all workers are sleeping and then starts the execution of the FORM program in which the preprocessor and the compiler treat each module before executing it. The execution phase of each module is the part that is eligible for parallelization.

In FORM each expression is executed in sequence which means one after the other. The same happens with the terms inside the expressions. The order is determined by the final sorting in the previous module. In TFORM (as well as in ParFORM) the master process reads the terms of one expression and distributes them over the workers. The terms are usually bunched together in something called buckets. This is because signals<sup>4</sup> have to be sent to the worker that should receive the terms and these signals turn out to be rather costly. For many programs the optimal bucket size lies between 100 and 1000 terms. Currently the default has been set to 500 terms. To keep the workers waiting as short as possible, the total number of buckets is twice the number of workers. This way the master process can prepare spare buckets. Once a worker becomes available all the master has to do is to copy some necessary ‘environment’ variables to the private data structure of the worker and the pointer to the bucket. After that a wakeup signal can be sent.

The next critical point is when the last terms in the input expression have been sent. After this workers will become available, but there are no further terms for them. This can become inefficient, if there is a single worker that has one or more terms that require much CPU time. Often such difficult terms tend to be grouped together and hence they could be inside the same bucket. For this a load balancing system has been designed in which the master processor will look inside the buckets of the workers and possibly steal some terms back to give to idle workers. This works well, but it fails in the case that there is a single term that uses most of the CPU time. Such is the case in the following FORM code:

```
Symbols x1, ..., x10;
Local F = (x1+...+x10)^10;
id x10 = 1-x1-...-x9;
.end
```

A new expressions always starts as a single term and inside the module it will be expanded. Hence the above example cannot use the parallelization, because its single term

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<sup>4</sup>Technically they aren't signals. Calls to `pthread_cond_signal()` are part of the Pthreads mechanisms for synchronizing processes. For the sake of simplicity we will refer to them as signals.

will end up inside a single worker.

In principle FORM is designed in such a way that also this load can be balanced. It is possible to interfere in the expansion tree and organize the delegation of subtasks to idle co-workers. It does however need a number of strategic decisions about when and where in the tree this interference should take place. This will be considered in a later version. For now one should remember that if one would like to make efficient use of the parallelization, one should avoid having the early modules doing too much work and keep the work for the modules in which there are already a large number of terms at the input. In the case of the above example one could benefit from parallelization if the program would read

```
Symbols x1, ..., x10;
Local F = (x1+...+x10)^10;
.sort
id x10 = 1-x1-...-x9;
.end
```

because the id-statement which does most of the work, is now executed at a moment that the master can distribute terms over the workers. At the same time the overhead of the extra sort is still negligible compared to the total amount of work to be done.

Once the input terms have all been processed, it is time to do the final sorting. Here we are faced with the second bottleneck, because now the completed expression should be put together under the control of the single master processor. The idea is to have the workers do as much of the sorting as possible. The master will then merge the sorted results of the workers.

We tried several methods for this sorting. In the first version of TFORM each worker did a regular sort as is done in the sequential version of FORM. This means that each might use a sort file and each would write its output to an output scratch file. Then the master would merge these files in the same way that the patches in a single sort file would be merged. The only difference was that rather than reading from different sorted patches inside a single file, the patches would come from different files. The results were disastrous. Apparently the LINUX file system that was used (ReiserFS) becomes very inefficient when several files are accessed at the same time by the same program. The test program running on four processors made the program actually take more than four times as much real time as when it was running on just a single processor, and the whole computer became very slow. The exact cause is still not clear and the issue remains under investigation.

The solution that has been selected cuts out the output scratch files for the workers. Instead the workers write their output directly into the sort buffers of the master. The master will then sort them simultaneously, provided this is possible. A system of data blocks has been set up to allow this simultaneous work. Each 'stream' from the workers inside the sort buffer of the master is divided into 10 blocks that are arranged in a circular fashion. Locks prevent the master from accessing blocks that have not been filled by the workers and workers from writing into blocks that have not yet been treated by the master. The result is that now each worker needs only one file, its sort file. Performance is much better, but still there are problems with the LINUX file system we used. The same



program that needed four times as much real time in the first approach, now needs a bit more than half the amount of real time than the sequential version. And during some of the sorting the computer is virtually unusable from the terminal. This depends very much on the way the file activity is organized. The size of the sorting buffers and the cache are of great influence. It also depends on which computer is used. We illustrate this with the first example in the next section (case N=15). In all other examples however this effect doesn't occur.

The next problem to address is the treatment of the dollar variables. These variables are in principle common but if one isn't careful each processor may obtain a private value and hence they are a potential source of conflicts. Let us have a look at some examples:

```

#$m1 = 0;
#$mc = 0;
if ( count(x1,1) > $m1 ) $m1 = count_(x1,1);
if ( count(y,1) > 0 ) $mc = $mc+1;
if ( match(f(x?$mx)) );
    id g($mx,n?) = n+$mx;
endif;
.sort

```

We have here three uses of dollar variables that can lead to problems. In the case of the variable \$m1 two workers could make the compare at the same time, after which the eventual value might not become the maximum of those two. In the case of \$mc one could have that worker 1 picks up the value, let us say 6, then worker 2 also picks up 6, then worker 1 writes 7, after which worker 2 writes 7. And similar accidents can happen with \$mx.

In the case of these accidents the placing of locks is counterproductive. One would have to place the lock around most of the substitution tree and hence the workers would obstruct each other rather thoroughly. There is however a solution. If one can tell TFORM what is meant with the dollar variable, TFORM might be able to deal with it without much overhead. This is done with the ModuleOption statement at the end of the module. In this case we would add the statement

```
ModuleOption,maximum,$m1,sum,$mc,local,$mx;
```

before the .sort instruction. This way TFORM knows that we collect a maximum in \$m1 and hence makes a compare, if needed locks the variable, makes a new compare (!) and if needed stores the new value. Then it releases the lock on the variable. This implementation has a side effect. The following code would produce the same result:

```

#$m1 = 0;
$m1 = count_(x1,1);
ModuleOption,maximum,$m1;
.sort

```

The maximum declaration in the ModuleOption statement makes the original if statement superfluous. It would however be very unwise to program this way because the sequential

version of FORM just ignores this part of the ModuleOption statement and hence would not produce the proper result. Also the different working of ParFORM would not give the correct result.

The case of the sum is more complicated. We have to place a lock immediately when the statement starts its evaluation and we can release the lock only after the new assignment. This is rather expensive and therefore should be used with great care. Of course this sum option can be abused. One could apply it to other assignments that are not sums. One is strongly advised against this, because in the future the inner workings may change and only a proper sum will work correctly<sup>5</sup>. In the case of a local use of a dollar variable (like  $\$m_x$ ), each thread will have its own copy. If there was a common value at the start of the module, this private/local variable will be initialized by it. At the end of the module the private copy is deleted. The common copy will still have the value that it had at the start of the module.

In ParFORM the treatment of the above dollar variables is somewhat different. Each processor has its own copy of the variable administration. Hence  $\$m_l$  will become a local/private maximum and  $\$m_c$  a sum over all terms treated in that processor. Only at the end of the module the private values are combined into a common value, using the information in the ModuleOption statement.

Of course there can be uses of dollar variables that cannot be parallelized. If TFORM is not helped in its use of dollar variables it will automatically switch to sequential mode for the module(s) in which this dollar variable is given a value during execution. Hence old programs will continue to run. They may however not benefit from the parallelization in such modules. The sequential version of FORM accepts the ModuleOption statement, but it just ignores most of its options. When writing programs that use dollar variables, it is best to plan the ModuleOption statements with it. This way the program will run optimally both in the sequential and in the parallel mode.

The only other instance of the use of local/private definitions of common variables concerns the preprocessor variables as redefined by the redefine statement. At first one might think that here we run into the same problems as with the dollar variables, but this is not the case. The value of a preprocessor variable can only be used by the preprocessor. Therefore during execution it is a write-only variable. If we remember which term was responsible for the last redefinition, we have to make the redefinition only when the number of the current input term is greater or equal to the number of the input term that caused the last redefinition. Hence this problem has been solved inside FORM and needs no special attention from the user.

## 4 The performance

The above modifications allow the parallel running of all existing FORM programs. The only exception are the modules in which dollar variables are assigned during execution time and that are not aided by ModuleOption statements. These will be run in sequential

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<sup>5</sup>One could think of substituting zero for the occurrence of  $\$m_c$  in the right hand side and adding the obtained value for the complete right hand side into the existing value of  $\$m_c$ . This would reduce the time for locking the variable by a considerable amount.

mode. The running itself is rather simple. One calls TFORM in the same way FORM used to be called but an extra argument in the command tail is needed to specify the number of worker threads as in:

```
tform -w4 -l calcdia
```

The `-w` argument specifies the number of workers. If this number is zero or one, there will only be a master thread. Invocation of a single worker would only cost overhead for communication and wouldn't improve performance. Omission of the `-w` argument is the same as `-w0`.

The first testcase we use is a simple program for computing so-called chromatic polynomials. The program is described in a set of example files in the FORM distribution. We use the program named `p15.frm` for a variety of lattice sizes. The main part of the code is a loop given by

```
Multiply F'i';
repeat id d(?a,k?,?b)*d(?c,k?,?d) = d(?a,?c,k,?b,?d);
Symmetrize d;
repeat id d(?a,k?,k?,?b) = d(?a,k,?b);
#do d = 1, 'D'
  id d(k'i',?a,k1?kk0[x],?b,k2?kk'd'[x],?c) = 0;
#enddo
id,ifmatch->1,d(k'i',k?) = 1;
id,ifmatch->1,d(k'i',?b) = d(?b);
  Multiply acc([q-1]+1);
Label 1;
id d = 1;
.sort:'i';
```

The loop contains a few `id`-statements with non-trivial pattern matching and some polynomial arithmetic inside the 'polyfun' named `acc`. We ran this program on a variety of computers.

The first computer, designated P, has two Pentium4 processors at 1.7 GHz. The second computer, designated N, contains 4 Opteron processors at 2.6 GHz. The third computer is a SGI computer with 32 Itanium processors at 1.3 GHz. Although the last computer uses shared memory there is a hierarchy. Also one has to specify how many and which processors are assigned to the job. We will designate the SGI computer S# in which # is the number of processors reserved for the task. These processors are usually selected to have the best memory latency for the selected number. In one case we did a test in which we selected the memory in such a way that the situation was the worst possible. This made an almost two percent difference in execution time. Hence the effect doesn't seem very relevant. In the case that we specify zero workers we used the sequential version of FORM. In the case we specify one worker we used TFORM with only the master thread.

Lattice	Computer	Workers	Time(sec)	Improvement
10	N	0	12.45	
10	N	1	11.92	1.0445
10	N	2	6.64	1.8750
10	N	3	4.83	2.5776
10	N	4	4.30	2.8953
10	N	5	4.37	2.8490
10	N	6	4.35	2.8621
10	N	7	4.30	2.8953
10	N	8	4.36	2.8555
10	N	16	4.50	2.7667
10	N	32	4.95	2.5152
10	P	0	29.96	
10	P	1	30.67	0.9769
10	P	2	17.45	1.7169

Table 1: Performance for chromatic polynomials on simple lattices.

We notice here a number of things. One is that on the Opteron this program is a rare example in which TFORM in single thread mode is faster than the sequential version of FORM. Next we see that there is a saturation effect, because the computer N has four processors. With more than 4 workers the improvement factor starts going down again. At first the effect is not very strong. In the case of 32 workers though the master has to start doing more work in the sorting as it has to merge 32 streams which gives on average 5 compares per term. This effect becomes noticeable. The optimum is at 4 workers, as the master seems to be able to get enough time on the various processors. In total the master processor used 1.16s of CPU time in that run. Finally we make a remark about running a smaller number of threads than there are processors. In the case of two workers on a machine with four processors the master can run simultaneously with the workers. This can make a difference during the final stages of the sorting and during the filling of the input buckets. Hence the efficiency will be slightly higher than it would be on a similar machine with two processors. We see this effect here when we compare the efficiencies for two workers on the machines N (4 processors) and P (2 processors), although it is not completely clear whether this is the only reason for the difference.

For bigger lattices we obtain:

Lattice	Computer	Workers	Time(sec)	Improvement
14	N	0	2809.57	
14	N	1	2695.64	1.0423
14	N	2	1443.07	1.9469
14	N	3	1059.07	2.6529
14	N	4	923.86	3.0411
14	P	0	15786.37	
14	P	2	9163.65	1.7227
15	N	0	12541.39	
15	N	1	12234.77	1.0251
15	N	2	7601.00	1.6500
15	N	3	5332.38	2.3519
15	N	4	3892.80	3.2217
15	N	6	4046.58	3.0993
15	N	8	4100.53	3.0585
15	P	0	77349.12	
15	P	2	45996.63	1.6816
15	S0	0	21897.79	
15	S2	2	13853.82	1.5806
15	S4	4	7867.53	2.7833
15	S8	8	6622.27	3.3067
15	S16	16	5543.48	3.9502
15	S32	32	5664.99	3.8655

Table 2: Performance for chromatic polynomials on bigger lattices.

In the case of the 14x14 lattice on the computer N the expression could still fit inside the allocated buffers. We see a similar pattern as before. For the 15x15 case however the sorting stage needed heavy use of disk files. We used a large sorting buffer of 2 Gigabytes. This is still not large enough to avoid the use of a sort file for each of the workers. One can see that the behaviour is much better on the Opteron computer (N) than on the SGI machine. To compare we also ran with smaller buffers (400 Mbytes). This causes many more and smaller disk operations. It gives the following results:

Lattice	Computer	Workers	Time(sec)	Improvement
15	N	0	14807.98	
15	N	2	9834.92	1.5057
15	N	4	7833.86	1.8903
15	S0	0	25312.04	
15	S2	2	15208.59	1.6643
15	S4	4	12879.12	1.9654
15	S8	8	11832.35	2.1392
15	S16	16	10490.21	2.4129
15	S32	32	10200.72	2.4814

Table 3: As in table 2 but with smaller buffers.

Clearly lots of (nearly) simultaneous disk operations have a bad effect on the performance. It is guessed that this effect is due to the particular type of file system used (ReiserFS) as another file system doesn't show this effect<sup>6</sup> The slower computer P, which has less memory and hence smaller buffers anyway, has already problems with the disk when running in the sequential mode and hence the effect seems to be less pronounced. The different ratio in the speed between the computers as compared to the  $N = 10$  lattice is due to several factors. First the bigger coefficients in the bigger lattices make the 64 bits Opteron more efficient. And second the Opteron computer has much more memory. For the determination of the improvement factor this is however irrelevant.

The second example concerns a number of Feynman diagrams as computed for past publications. The program used for these computations is called mincer and it has been heavily optimized for use with version 2 of FORM. Later some extra optimizations were added for the use with version 3. Not a single line was changed for the runs with TFORM. We label the diagrams d1c, d10c and d11c as they come from a set computed for the nonsinglet sector of the form factor F2 in deep inelastic scattering. They are three loop diagrams of the non-planar type and we calculate Mellin moments.

Diagram	Moment	Computer	Workers	Time(sec)	Improvement
d1c	10	N	0	32.70	
d1c	10	N	1	33.41	0.9799
d1c	10	N	2	18.29	1.7900
d1c	10	N	3	12.85	2.5479
d1c	10	N	4	10.39	3.1511
d1c	16	N	0	694.02	
d1c	16	N	1	727.52	0.9540
d1c	16	N	2	387.74	1.7899
d1c	16	N	3	266.23	2.6068
d1c	16	N	4	208.74	3.3248
d1c	10	P	0	108.55	
d1c	10	P	2	60.32	1.7996
d1c	16	P	0	2759.26	
d1c	16	P	2	1463.70	1.8851
d10c	10	N	0	880.66	
d10c	10	N	1	894.69	0.9843
d10c	10	N	2	474.23	1.8570
d10c	10	N	3	328.68	2.6794
d10c	10	N	4	263.33	3.3443
d11c	10	N	0	3208.02	
d11c	10	N	1	3271.74	0.9805
d11c	10	N	2	1719.05	1.8662
d11c	10	N	3	1221.12	2.6271
d11c	10	N	4	971.87	3.3009

Table 4: Runs with Mincer for three different diagrams.

<sup>6</sup>This was tested with heavy file copy operations rather than with TFORM.

The diagram d11c was the ‘most difficult’ diagram in the set of all diagrams that had to be computed for this reaction. Because more work is done inside the modules the efficiency here is higher than in the previous example. One has to consider that in the case of multithreaded runs there is an overhead of 5% - 10% as compared to the sequential version. This means that the theoretical limit is an improvement factor of about 3.6. on a computer with 4 processors. One has to add that there is also a part of the time in which only the master thread is active. Hence an improvement factor of 3.3 on a machine with four processors is quite high.

Diagram	Moment	Computer	Workers	Time(sec)	Master	Improvement
d11c	10	S1	0	7042.83		
d11c	10	S2	2	3834.41	196.21	1.8367
d11c	10	S3	3	2625.51	273.05	2.6825
d11c	10	S4	4	2117.88	291.37	3.3254
d11c	10	S6	6	1644.43	392.37	4.2828
d11c	10	S8	8	1335.42	418.30	5.2739
d11c	10	S12	12	1093.35	483.02	6.4415
d11c	10	S16	16	1002.00	522.96	7.0288
d11c	10	S24	24	953.70	606.45	7.3847
d11c	10	S32	32	957.79	658.27	7.3532

Table 5: Runs with Mincer on the SGI computer using TFORM.

On the sgi machine we notice a clear example of saturation. Till 4 threads the improvement is quite nice. From 4 to 16 threads the improvement isn’t very spectacular but might still be worth it. Above that one can hardly see any further improvement and one is just wasting resources. For 32 processors the improvement even worsens. The dominant reason is clearly the time spent by the master process. This increases when more threads are involved. The two effects one can think of immediately are the extra work that has to be done by the master in the final sorting and the fact that more threads have to be provided with data, running the risk that threads have to wait. This last effect seems to be present, but even if it wouldn’t be present, the efficiency would be no more than about 8 for 32 processors. If we ignore this last effect (which can be controled somewhat with the setting of the variable ThreadBucketSize) the real time would roughly follow a formula like

$$T = c_1 + c_2/N + c_3^2 \log N \quad (1)$$

in which N is the number of workers and  $c_1, c_2, c_3$  are constants. One can read from the table that each extra compare that has to be done by the master thread (5 in the case of 32 processors) adds more than 100 sec to its CPU time and hence  $c_3 \approx 115$ . Therefore the sorting seems to be the main culprit. Clearly much of the saturation can be alleviated by a future change in the workload of the master during the final stages of the sorting. As the values of  $c_1, c_2, c_3$  depend on the problem the exact value of the saturation in the improvement factor is hard to predict.

We managed to compare this last example with the current version of ParFORM. This gives similar results as can be seen from the next table:

Diagram	Moment	Computer	Workers	Time(sec)	Improvement
d11c	10	S1	0	6998.00	
d11c	10	S3	2	3704.21	1.8892
d11c	10	S4	3	3353.51	2.0868
d11c	10	S5	4	2117.88	3.3042
d11c	10	S8	7	1642.14	4.2615
d11c	10	S9	8	1374.69	5.0906
d11c	10	S13	12	1221.45	5.7293
d11c	10	S16	15	1163.73	6.0134
d11c	10	S17	16	1088.27	6.4304
d11c	10	S24	23	1074.26	6.5143
d11c	10	S25	24	986.38	7.0946
d11c	10	S32	31	1010.73	6.9237

Table 6: Runs with Mincer on the SGI computer using ParFORM.

We see here the same saturation. One should also realize that in ParFORM one processor is reserved for the master process exclusively. This explains the rather bumpy transition when the number of processors is around a power of two. On the whole, the slower MPI communication costs a marginal amount of efficiency. This situation is better when the size of the various buffers is tuned to ParFORM and the specific computer. In that case S32 reaches an efficiency of 7.4247. In the case of an experimental version of ParFORM in data is transferred via shared memory the maximum efficiency becomes 7.7132 for S32.

The final example concerns the solution of a large system of equations. The equations in question form all relations that are known between multiple zeta values of the same weight. This system was described in ref [20] and the programs can be found in the FORM distribution under ‘summer’. In ref [20] the equations were only worked out and solved till weight 9, but because the computers have become faster and we have parallel processing we decided to try weight 10 as well. It turned out that most of the CPU time for the weight 10 calculations is used for the calculation of GCD’s of large integers (like 300 decimal digits). Considering that originally FORM was optimized for relatively short integers and that hence some improvements might be possible, the GCD algorithms were studied carefully and indeed a significant improvement for large integers was found. This made the runs of the weight 10 programs faster by a factor 6.95. For weight 9 the improvement was a factor 2.76 as there the numbers were not quite as large. We will give the timings with the new version<sup>7</sup>. It should be noted that a part of the program in which a number of equations are prepared for a Gaussian elimination is better done inside a single processor. This does spoil the parallelization a bit for the lower weights. For the higher weights most work is in the rational arithmetic, even with the improved routines, and hence this gives a much better efficiency. But it does give a relatively large value for the CPU time used by the master process.

<sup>7</sup>Because the calculations of the GCD’s is mostly local/private during term generation and normalization, the efficiency of the parallelization is somewhat better for the old version. We have reached there a record factor of 3.49 for the weight 10 calculation on 4 processors, but the total execution time was of course much larger than with the new version.



Weight	Computer	Workers	Time(sec)	CPU master(sec)	Improvement
7	N	0	8.43		
7	N	4	8.74	4.91	0.9645
8	N	0	78.71		
8	N	4	62.73	36.92	1.2547
9	N	0	2026.15		
9	N	2	1330.41	372.65	1.5230
9	N	4	909.47	373.40	2.2278
10	N	0	130860.59		
10	N	2	71479.05	5510.02	1.8308
10	N	4	39151.96	5590.78	3.3424
10	S1	0	233797.28		
10	S2	2	125072.98	10013.77	1.8693
10	S4	4	69075.57	10645.14	3.3847
10	S8	8	41987.38	11693.31	5.5683
10	S12	12	32910.15	12194.61	7.1041
10	S16	16	28317.13	12564.38	8.2564
10	S24	24	24108.43	13189.96	9.6977
10	S32	32	22143.32	13633.52	10.5584

Table 7: Solving dependencies between multiple zeta values.

Beyond weight 9 the test couldn't be done on 32-bits architectures, because the size of the tables would be too large (FORM has a limitation on the number of elements in a single table which depends on the word size). We see on the Silicon Graphics computer that the CPU time of the master process becomes the limiting factor in the efficiency when we increase the number of processors. The eventual value of the improvement factor is of course a function of how much work can be done completely locally. As the rational arithmetic is very local, the factor here is somewhat better than in the previous examples. The waiting of workers for input terms and the signals involved in sending them were quite a factor here. The runs we show here were with a ThreadBucketSize of 1000. With a value of 100 the improvement factor for 32 processors went down to 7.75.

It is also clear that when the equations are relatively simple the solution selected doesn't benefit much from parallelization. This is in the nature of the problem, and only because of the vast amount of rational arithmetic the more complicated cases give good improvements.

## 5 Conclusions

TFORM is working and can handle in principle all existing programs that would run on version 3.1 of FORM<sup>8</sup>. On computers with a limited number of processors the improvement in running time is quite good. The more complicated the program the better the improvement.

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<sup>8</sup>It is of course not excluded that the extensive changes have caused the inclusion of some new bugs that still have not been caught. If encountered, please report them to the authors.

We have compared the performance of TFORM with that of ParFORM. As is to be expected, when the two can run on the same computers they give more or less the same increase in efficiency. Each have their own strong points. ParFORM can run on a larger number of multi processor systems. The internal organization allows TFORM to run more existing programs in parallel. Also for a small number of processors TFORM can run more efficiently as it doesn't have to reserve a whole processor for the master.

For computers with a larger number of processors there is a strong saturation effect due to the increase in tasks that have to be done by the master thread. The way to improve TFORM in the future has to be sought in the lightening of the load of the master process. One way is to remove the signals that are sent around between the master and the workers when the terms are distributed over the workers. The most important however seems to be a change in the final stages of the sorting. In principle it is possible to set this up as a binary tree in which all but the last step are done by workers, and maybe it is even possible to let a worker do the last step, making the master only responsible for writing the final result. This will be tried in a future version of TFORM. There will however always be a program dependent limit as a sorting tree will always imply a bottleneck in which one compare per term is to be done inside a single thread<sup>9</sup>. In the ideal case the execution time will follow a rule like

$$T = c_1 + c_2/N \quad (2)$$

in which the values of  $c_1$  and  $c_2$  are problem dependent. One can try to optimize programs in such a way that for a given number of workers the execution time is optimal. We have not yet experimented with this but we expect that some users will do this in the future.

Considering the bottlenecks it should be clear that at the moment only for special problems the use of a very large number of processors can be beneficial. For a small number of processors however the current version of TFORM is more than adequate.

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<sup>9</sup>One could imagine a sorting system in which two processes are working simultaneously, one from the smallest term upward and one from the largest term downward. This creates an enormous complexity while the benefits are relatively limited.

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