Facilitating reproducibility in scientific computing:  
Principles and practice*

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Abstract
The foundation of scientific research is theory and experiment, carefully documented in open publications, in part so that other researchers can reproduce and validate the claimed findings. Unfortunately, the field of scientific and mathematical computing has evolved in ways that often do not meet these high standards. In published computational work, frequently there is no record of the workflow process that produced the published computational results, and in some cases, even the code is missing or has been changed significantly since the study was completed. In other cases, the computation is subject to statistical errors or numerical variability that makes it difficult for other researchers to reconstruct results. Thus confusion often reigns.

That tide may be changing, though, in the wake of recent efforts that recognize both the need for explicit and widely implemented standards, and also the opportunity to do computational research work more effectively. This chapter discusses the roots of the reproducibility problem in scientific computing and summarizes some possible solutions that have been suggested in the community.

1 Introduction
By many measures, the record of the field of modern high-performance scientific and mathematical computing is one of remarkable success. Accelerated by relentless advances of Moore’s law, this technology has enabled researchers in many fields to perform computations that would have been unthinkable in earlier times. Indeed, computing is rapidly

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becoming central to both scientific theory and scientific experiment. Computation is already essential in data analysis, visualization, interpretation and inference.

The progress in performance over the past few decades is truly remarkable, arguably without peer in the history of modern science and technology. For example, in the July 2014 edition of the Top 500 list of the world’s most powerful supercomputers (see Figure 1), the best system performs at over 30 Pflop/s (i.e., 30 “petaflops” or 30 quadrillion floating-point operations per second), a level that exceeds the sum of the top 500 performance figures approximately ten years earlier [4]. Note also that a 2014-era Apple MacPro workstation, which features approximately 7 Tflop/s (i.e., 7 “teraflops” or 7 trillion floating-point operations per second) peak performance, is roughly on a par with the #1 system of the Top 500 list from 15 years earlier (assuming that the MacPro’s Linpack performance is at least 15% of its peak performance).

Just as importantly, advances in algorithms and parallel implementation techniques have, in many cases, outstripped the advance from raw hardware advances alone. To mention but a single well-known example, the fast Fourier transform (“FFT”) algorithm reduces the number of operations required to evaluate the “discrete Fourier transform,” a very important and very widely employed computation (used, for example, to process

Figure 1: Performance of the Top 500 computers: Red = #1 system; orange = #500 system; blue = sum of #1 through #500.
signals in cell phones), from $8n^2$ arithmetic operations to just $5n\log_2 n$, where $n$ is the total size of the dataset. For large $n$, the savings are enormous. For example, when $n$ is one billion, the FFT algorithm is more than six million times more efficient.

Yet there are problems and challenges that must be faced in computational science, one of which is how to deal with a growing problem of non-reproducibility in computational results.

A December 2012 workshop on reproducibility in computing, held at the Institute for Computational and Experimental Research in Mathematics (ICERM) at Brown University, USA, noted that

Science is built upon the foundations of theory and experiment validated and improved through open, transparent communication. With the increasingly central role of computation in scientific discovery, this means communicating all details of the computations needed for others to replicate the experiment. ... The “reproducible research” movement recognizes that traditional scientific research and publication practices now fall short of this ideal, and encourages all those involved in the production of computational science ... to facilitate and practice really reproducible research. [50]

From the ICERM workshop and other recent discussions, four key issues have emerged:

1. The need to institute a culture of reproducibility.
2. The danger of statistical overfitting and other errors in data analysis.
3. The need for greater rigor and forthrightness in performance reporting.
4. The growing concern over numerical reproducibility.

In the remainder of the chapter, we shall discuss each of these items and will describe solutions that have been proposed within the community.

2 A culture of reproducibility

As mentioned above, the huge increases in performance due to hardware and algorithm improvement have permitted enormously more complex and higher fidelity simulations to be performed. But these same advances also carry an increased risk of generating results that either do not stand up to rigorous analysis, or cannot reliably be reproduced by independent researchers. As noted in [50], the culture that has arisen surrounding scientific computing has evolved in ways that often make it difficult to maintain reproducibility — to verify findings, to efficiently build on past research, or even to apply the basic tenets of the scientific method to computational procedures.
Researchers in subjects like experimental biology and physics are taught to keep careful lab notebooks documenting all aspects of their procedure. In contrast, computational work is often done in a much more cavalier manner, with no clear record of the exact “workflow” leading to a published paper or result. As a result, it is not an exaggeration to say that in most published results in the area of scientific computing, reproducing the exact results by an outside team of researchers (or even, after the fact, by the same team) is often impossible. This raises the disquieting possibility that numerous published results may be unreliable or even downright invalid [26, 37].

The present authors have personally witnessed these attitudes in the field. Colleagues have told us that they can’t provide full details of their computational techniques, since such details are their “competitive edge.” Others have dismissed the importance of keeping track of the source code used for published results. “Source code? Our student wrote that, but he is gone now.”

The ICERM report [50] summarized this problem as follows:

Unfortunately the scientific culture surrounding computational work has evolved in ways that often make it difficult to verify findings, efficiently build on past research, or even to apply the basic tenets of the scientific method to computational procedures. Bench scientists are taught to keep careful lab notebooks documenting all aspects of the materials and methods they use including their negative as well as positive results, but computational work is often done in a much less careful, transparent, or well-documented manner. Often there is no record of the workflow process or the code actually used to obtain the published results, let alone a record of the false starts. This ultimately has a detrimental effect on researchers’ own productivity, their ability to build on past results or participate in community efforts, and the credibility of the research among other scientists and the public.

2.1 Documenting the workflow

The first and foremost concern raised in the ICERM report [50] was the need to carefully document the experimental environment and workflow of a computation. Even if all of this information is not included in the published account of the work, a record should be kept while the study is underway, and the information should be preserved. Specific recommendations include the following [50]:

1. A precise statement of assertions to be made in the paper.
2. The computational approach, and why it constitutes a rigorous test.
3. Complete statements of, or references to, every algorithm employed.
4. Auxiliary software (both research and commercial software).
5. Test environment, including hardware, software and number of processors.
6. Data reduction and statistical analysis methods.
7. Adequacy of precision level and grid resolution.
8. Full statement or summary of experimental results.
9. Verification and validation tests performed.
10. Availability of computer code, input data and output data.
11. Curation: where are code and data available?
12. Instructions for repeating computational experiments.
13. Terms of use and licensing. Ideally code and data “default to open.”
14. Avenues explored and negative findings.
15. Proper citation of all code and data used.

2.2 Tools to aid in documenting workflow and managing data

A number of the presentations at the ICERM workshop, as summarized in the report [50] described emerging tools to assist researchers in documenting their workflow. Here is a brief summary of some of these tools, condensed and adapted from [50]:

- **Literate programming, authoring, and publishing tools.** These tools enable users to write and publish documents that integrate the text and figures seen in reports with code and data used to generate both text and graphical results. This process is typically not interactive, and requires a separate compilation step. Some examples here are WEB, Sweave, and knitr, as well as programming-language-independent tools such as Dexy, Lepton, and noweb. Other authoring environments include SHARE, Doxygen, Sphinx, CWEB, and the Collage Authoring Environment.

- **Tools that define and execute structured computation and track provenance.** Provenance refers to the tracking of chronology and origin of research objects, such as data, source code, figures, and results. Tools that record provenance of computations include VisTrails, Kepler, Taverna, Sumatra, Pegasus, Galaxy, Workflow4ever, and Madagascar.

- **Integrated tools for version control and collaboration.** Tools that track and manage work as it evolves facilitate reproducibility among a group of collaborators. With the advent of version control systems (e.g., Git, Mercurial, SVN, CVS), it has become
easier to track the investigation of new ideas, and collaborative version control sites like Github, Google Code, BitBucket, and Sourceforge enable such ideas to be more easily shared. Furthermore, these web-based systems ease tasks like code review and feature integration, and encourage collaboration.

- **Tools that express computations as notebooks.** These tools represent sequences of commands and calculations as an interactive worksheet. Examples include both closed-source tools such as MATLAB (through the publish and app features), Maple, and Mathematica, as well as open-source tools such as IPython, Sage, RStudio (with knitr), and TeXmacs.

- **Tools that capture and preserve a software environment.** A major challenge in reproducing computations is installing the prerequisite software environment. New tools make it possible to exactly capture the computational environment. For instance, Docker, VirtualBox, VMWare, or Vagrant can be used to construct a virtual machine image containing the environment. Blueprint analyzes the configuration of a machine and outputs its text description. ReproZip captures all the dependencies, files and binaries of the experiment, and also creates a workflow specification for the VisTrails system in order to make the execution and exploration process easier. Application virtualization tools, such as CDE (Code, Data, and Environment), attach themselves to the computational process in order to find and capture software dependencies. Some computational environments can also be constructed and made available in the cloud, and others feature full workflow tracking. Examples include Synapse/clearScience and HUBzero including nanoHUB.

- **Interactive theorem proving systems for verifying mathematics and computation.** “Interactive theorem proving”, a method of formal verification, uses computational proof assistants to construct formal axiomatic proofs of mathematical claims. Examples include coq, Mizar, HOL4, HOL Light, ProofPowerHOL, Isabelle, ACL2, Nuprl, Veritas, and PVS. Notable theorems such as the Four Color Theorem and the Prime Number Theorem have been verified in this way, and Thomas Hales’s Flyspeck project, using HOL Light and Isabelle, aimed to obtain a formal proof of the Kepler conjecture. Each one of these projects produces machine-readable and exchangeable code that can be integrated in to other programs. For instance, each formula in the web version of NIST’s authoritative Digital Library of Mathematical Functions may be downloaded in TeX or MathML (or indeed as a PNG image) and the fragment directly embedded in an article or other code. This dramatically reduces chances of transcription error and other infelicities being introduced.

- **Post-publication tools persistently connecting data, code, workflows, and articles** Although digital scholarly objects may be made available, it is quite possible each object may be hosted by different services and may reside at different locations on the
web (such as authors’ websites, journal supplemental materials documents, and various data and code repositories). Services such as http://ResearchCompendia.org, http://RunMyCode.org and http://ipol.im attempt to co-locate these objects on the web to enable both reproducibility and the persistent connection of these objects [52, 51].

The development of software tools enabling reproducible research is a new and rapidly growing area of research. We believe that the difficulty of working reproducibly will be significantly reduced as these and other tools continue to be adopted and improved — and their use simplified. But prodding the scientific computing community, including researchers, funding agencies, journal editorial boards, lab mangers and promotion committee members, to broadly adopt and encourage such tools remains a challenge [41, 51].

2.3 Other cultural changes

While the above recommendations and tools will help, there is also a need to fundamentally rethink the “reward system” in the scientific computing field (and indeed, of all scientific research that employs computing). Journal editors need to acknowledge importance of computational details, and to encourage full documentation of the computational techniques, perhaps on an auxiliary persistent website if not in the journal itself. Reviewers, research institutions and, especially, funding agencies need to recognize the importance of computing and computing professionals, and to allocate funding for after-the-grant support and repositories. Also, researchers, journals and research institutions need to encourage publication of negative results—other researchers can often learn from these experiences. Furthermore, as we have learned from other fields, only publishing “positive” results inadvertently introduces a bias into the field.

3 Statistical overfitting

As mentioned above, the explosion of computing power and algorithms over the past few decades has placed computational tools of enormous power in the hands of practicing scientists. But one unfortunate side effect of this progress is that it has also greatly magnified the potential for serious errors, such as statistical overfitting.

Statistical overfitting, in this context, means either proposing a model for an input dataset that inherently possesses a higher level of complexity than that of the input dataset being used to generate or test it, or else trying many variations of a model on an input dataset and then only presenting results from the one model variation that appears to best fit the data. In many such cases, the model fits the data well only by fluke, since it is really fitting only the idiosyncrasies of the specific dataset in question, and has little or no descriptive or predictive power beyond the particular dataset used in the analysis.
Statistical overfitting can be thought of as an instance of “selection bias,” wherein one presents the results of only those tests that support well one’s hypothesis [34].

The problem of statistical overfitting in computational science is perhaps best illustrated in the field of mathematical finance. A frequently-encountered difficulty is that a proposed investment strategy or a fund based on this strategy looks great on paper, based on its “backtest” performance, but fall flat when actually fielded in practice.

In the field of finance, a backtest is the usage of a historical market dataset to test how a proposed investment strategy would have performed had it been in effect over the period in question. The trouble is that in our present era, when a computer program can easily analyze thousands, millions or even billions of variations of a given investment strategy, it is almost certain that the optimal strategy, measured by its backtest performance, will be statistically overfit and thus of dubious predictive value.

In two 2014 papers, two of the present authors (Bailey and Borwein, with two other collaborators) derived formulas for (a) relating the number of trials to the minimum backtest length, and (b) computing the probability of backtest overfitting. They also demonstrated that under the assumption of memory in markets, overfit strategies are actually somewhat prone to lose money [13, 12].

How easy is it to overfit backtest data? Very! If only 2 years of daily backtest data are available, then no more than 7 strategy variations should be tried. If only 5 years of daily backtest data are available, then no more than 45 strategy variations should be tried (see Figure 2). In general, if a financial analyst or researcher does not report the number of trials $N$ explored when developing an investment strategy, then it is impossible for a third party to properly “reconstruct” the results, in the sense of correctly ascertaining the true level of reward and risk. Indeed, it is shown in [13, 12] that for given any desired performance goal, a financial analyst just needs to keep running her computer program, trying alternative parameters for her strategy, and eventually she will find a variation that achieves the desired performance goal, yet the resulting strategy will have no statistically valid predictive efficacy whatsoever.

Along this line, backtest overfitting can also be seen as an instance of the “filedrawer problem,” i.e., trying many variations of a given algorithm but only reporting the “winner” [38, 48].

### 3.1 A hands-on demonstration of backtest overfitting

To better understand the phenomenon of backtest overfitting, two of the present authors, together with researchers from Boston College and the Lawrence Berkeley National Laboratory, developed an online simulator of backtest overfitting (see http://datagrid.lbl.gov/backtest). It first generates a pseudorandom time series representing, say, several years of daily market prices, and then adjusts parameters of a simple investment strategy (monthly entry date, holding period, stop loss, etc.) to produce an “optimal” strategy. However, when this “optimal” strategy is then applied to a second dataset, it typically
fails miserably, evidence of severe overfitting. Figure 3 shows the performance of a typical “optimal” strategy, based on an input pseudorandom time series dataset, and Figure 4 shows the performance of this “optimal” strategy on a new dataset. Note that the Sharpe ratio (“SR”), which is a standard performance measure used in finance, is 1.32 for Figure 3, indicating fairly good performance, while the same statistic for Figure 4 is −0.03, indicating a completely ineffective strategy. See [15] for additional details.

In short, the field of mathematical/computational finance, like numerous other fields that employ “big data,” must recognize the need to pay careful attention to the rules of statistical inference when drawing conclusions. Sloppy data analysis no longer suffices.

3.2 Why the silence?

Among other things, the material highlighted above raises the question of why so many in computational science have remained largely silent with the regards to colleagues who, knowingly or not, employ questionable or non-reproducible practices in their work. In the computational finance field, this includes those who fail to disclose the number of models used in developing an investment strategy, make vague market predictions that do not permit rigorous testing, misuse probability theory and statistics, or employ pseudoscientific charts, graphs or technical jargon (e.g., “Fibonacci ratios,” “cycles,” “waves,” “pivot points,” etc.). As two of us wrote in [13], “Our silence is consent, making us accomplices.
Figure 3: Final optimized strategy applied to input dataset. Note that the Sharpe ratio is 1.32, indicating a fairly effective strategy on this dataset.

in these abuses.” And ignorance is a poor excuse.

Similarly, in several instances when we have observed instances of reproducibility problems and sloppy data analysis within other arenas of scientific computing (see the next few sections), a surprisingly typical reaction is to acknowledge that this is a significant problem in the field, but that it is either futile, unnecessary or inappropriate to make much fuss about it. Given recent experiences, it is clear that this approach is no longer acceptable.

4 Performance reporting in high-performance computing

One aspect of reproducibility in scientific computing that unfortunately is coming to the fore once again is the need for researchers to more carefully analyze and report data on the performance of their codes.
Figure 4: Final optimized strategy applied to new dataset. Note that the Sharpe ratio is -0.03, indicating a completely ineffective strategy on this dataset.

4.1 A 1992 perspective

Some background is in order here. One of the present authors (Bailey) was privileged to participate in some of the earliest implementations and analysis of highly parallel scientific computing, back in the late 1980s and early 1990s. In this time period, many new parallel systems had been introduced; each vendor claimed theirs was “best.” What’s more, many scientific researchers were almost as excited about the potential of highly parallel systems as were the computer vendors themselves. Few standard benchmarks and testing methodologies had been established, so it was hard to reproduce published performance results. Thus much confusion reigned. Overall, the level of rigor and peer review in the field was relatively low.

In response, Bailey published a humorous essay entitled “Twelve ways to fool the masses,” which was intended to gently poke some fun at some of these less-than-rigorous practices, hoping that this bit of humor would prod many in the community to tighten their standards and be more careful when analyzing and reporting performance.

The “Twelve ways” article was published in Supercomputing Review, which subse-
quently went defunct. Here is a brief reprise, taken from [6]:

1. Quote 32-bit performance results, not 64-bit results, but don’t mention this in paper.

2. Present performance figures for an inner kernel, then represent these figures as the performance of the entire application.

3. Quietly employ assembly code and other low-level language constructs.

4. Scale up the problem size with the number of processors, but omit any mention of this.

5. Quote performance results projected to a full system.

6. Compare your results against scalar, unoptimized code on conventional systems.

7. When run times are compared, compare with an old code on an obsolete system.


9. Quote performance as processor utilization, parallel speedups or Mflop/s per dollar.

10. Mutilate the algorithm used in the parallel implementation to match the architecture.

11. Measure parallel run times on a dedicated system, but measure conventional run times in a busy environment.

12. If all else fails, show pretty pictures and animated videos, and don’t discuss performance.

Since abuses continued, Bailey subsequently published a paper in which he explicitly called out a number of these practices, citing actual examples from peer-reviewed papers (although the actual authors and titles of these papers were not presented in the paper, out of courtesy) [6]. Here is a brief summary of some of the examples:

- **Scaling performance results to full-sized system.** In some published papers and conference presentations, performance results on small-sized parallel systems were linearly scaled to full-sized systems, without even clearly disclosing this fact. For example, in several cases 8,192-CPU performance results were linearly scaled to 65,536-CPU results, simply by multiplying by eight. Sometimes this fact came to light only in the question-answer period of a technical presentation. A typical rationale was, “We can’t afford a full-sized system.”
• Using inefficient algorithms on highly parallel systems. In other cases, inefficient algorithms were employed for the highly parallel implementation, requiring many more operations, thus producing artificially high performance rates. For instance, some researchers cited partial differential equation simulation performance based explicit schemes, for applications where implicit schemes were known to be much more efficient. Another paper cited performance for computing a 3-D discrete Fourier transform by direct evaluation of the defining formula, which requires $8n^2$ operations, rather than by using a fast Fourier transform (FFT), which requires $5n \log_2 n$ operations. Obviously, for sufficiently large problems, FFT-based computations can produce desired results with vastly fewer operations.

This is not to say that alternate algorithms should not be employed, but only that when computing performance rates, one should base the operation count on the best practical serial algorithm, rather than on the actual number of operations performed (or cite both sets of figures).

• Not actually performing a claimed computation. One practice that was particularly troubling was that of not actually performing computations that are mentioned in the study. For example, in one paper the authors wrote in the Abstract that their implementation of a certain application on a CM-2 system (a parallel computer available in the 1992 time frame) runs at “300-800 Mflop/s on a full [64K] CM-2, or at the speed of a single processor of a Cray-2 on 1/4 of a CM-2.” However, in the actual text of the paper, one reads that “This computation requires 568 iterations (taking 272 seconds) on a 16K Connection Machine.” Note that it was actually run on 16,384 processors, not 65,536 processors — the claimed performance figures in the Abstract evidently were merely the 16,384-CPU performance figures multiplied by four. One then reads that “In contrast, a Convex C210 requires 909 seconds to compute this example. Experience indicates that for a wide range of problems, a C210 is about 1/4 the speed of a single processor Cray-2.” In other words, the authors did not actually perform the computation on a Cray-2, as clearly suggested in the Abstract; instead they ran the computation on a Convex C210 and employed a rather questionable conversion factor to obtain the Cray-2 figure.

• Questionable performance plots. The graphical representation of performance data also left much to be desired in many cases. Figure 5 shows a plot from one study that compares run times of various sizes of a certain computation on a parallel systems, in the lower curve, versus the same computations on a vector system (a widely used scientific computer system at the time), in the upper curve. The raw data for this graph is as follows:
In the text of the paper where this plot appears, one reads that in the last entry, the 3:11:50 figure is an “estimate” — this problem size was not actually run on the vector system. The authors also acknowledged that the code for the vector system was not optimized for that system. Note however, by examining the raw data, that the parallel system is actually slower than the vector system for all cases, except for the last (estimated) entry. Also, except for the last entry, all real data in the graph is in the lower left corner. In other words, the design of the plot leaves much to be desired; a log-log plot should have been used for such data.

Figure 5: Performance plot [run time, parallel (lower) vs vector (upper)].
4.2 Fast forward to 2014: New ways to fool the masses

Perhaps given that a full generation has passed since the earlier era mentioned above, present-day researchers are not as fully aware of the potential pitfalls of performance reporting. In any event, various high-performance computing researchers have noted a “resurrection” of some of these questionable practices.

Even more importantly, the advent of many new computer architectures and designs has resulted in a number of “new” performance reporting practices uniquely suited to these systems. Here are some that are of particular concern:

- Citing performance rates for a run with only one processor core active in a shared-memory multi-core node, producing artificially inflated performance (since there is no shared memory interference) and wasting resources (since most cores are idle). For example, some studies have cited performance on “1024 cores” of a highly parallel computer system, even though the code was run on 1024 multicore nodes (each node containing 16 cores), using only one core per node, and with 15 out of 16 cores idle on each node. Note that since this implementation wastes 15/16 of the parallel system being used, it would never be tolerated in day-to-day usage.

- Claiming that since one is using a graphics processing unit (GPU) system, that efficient parallel algorithms must be discarded in favor of “more appropriate” algorithms. Again, while some algorithms may indeed need to be changed for GPU systems, it is important that when reporting performance, one base the operation count on the best practical serial algorithm. See “Using inefficient algorithms on highly parallel systems” in the previous subsection.

- Citing performance rates only for a core algorithm (such as FFT or linear system solution), even though full-scale applications have been run on the system (most likely because performance figures for the full-scale applications are not very good).

- Running a code numerous times, but only publishing the best performance figure in the paper (recall the problems of the pharmaceutical industry from only publishing the results of successful trials).

- Basing published performance results on runs using special hardware, operating system or compiler settings that are not appropriate for real-world production usage.

In each of these cases, note that in addition to issues of professional ethics and rigor, these practices result make it essentially impossible for other researchers to reconstruct claimed performance results, or to build on past research experiences to move the field forward. After all, if one set of researchers report that a given algorithm is the best known approach for performing a given application on a certain computer system, other researchers may also employ this approach, but then wonder why their performance results seem significantly less impressive than expected.
In other words, reproducible practices are more than just the “right” thing to do; they are essential for making real progress in the field.

5 Numerical reproducibility

The ICERM report also emphasized rapidly growing challenge of numerical reproducibility:

Numerical round-off error and numerical differences are greatly magnified as computational simulations are scaled up to run on highly parallel systems. As a result, it is increasingly difficult to determine whether a code has been correctly ported to a new system, because computational results quickly diverge from standard benchmark cases. And it is doubly difficult for other researchers, using independently written codes and distinct computer systems, to reproduce published results [50].

5.1 Floating-point arithmetic

Difficulties with numerical reproducibility have their roots in the inescapable realities of floating-point arithmetic. “Floating-point arithmetic” means the common way of representing non-integer quantities on the computer, by means of a binary sign and exponent, followed by a binary mantissa — the binary equivalent of scientific notation, e.g., \(3.14159 \times 10^{13}\). For most combinations of operands, computer arithmetic operations involving floating-point numbers produce approximate results. Thus roundoff error is unavoidable in floating-point computation.

For many scientific and engineering computations, particularly those involving empirical data, IEEE 32-bit floating-point arithmetic (roughly 7-digit accuracy) is sufficiently accurate to produce reliable, reproducible results, as required by the nature of the computation. For more demanding applications, IEEE 64-bit arithmetic (roughly 15-digit accuracy) is required. Both formats are supported on almost all scientific computer systems.

Unfortunately, with the greatly expanded scale of computation now possible on the enormously powerful, highly parallel computer systems now being deployed, even IEEE 64-bit arithmetic is sometimes insufficient. This is because numerical difficulties and sensitivities that are minor and inconsequential on a small-scale calculation may be major and highly consequential once the computation is scaled up to petascale or exascale size. In such computations, numerical roundoff error may accumulate to the point that the course of the computation is altered (i.e., by taking the wrong branch in a conditional branch instruction), or else the final results are no longer reproducible across different computer systems and implementations, thus rendering the computations either useless or misleading for the science under investigation.
5.2 Numerical reproducibility problems in real applications

As a single example, the ATLAS (acronym for “A Toroidal LHC Apparatus”) experiment at the Large Hadron Collider must track charged particles to within 10-micron accuracy over a 10-meter run, and, with very high reliability, correctly identify them. The software portion of the experiment consists of over five million lines code (C++ and Python), which has been developed over many years by literally thousands of researchers.

Recently, some ATLAS researchers reported to us that in an attempt to improve performance of the code, they changed the underlying math library. When this was done, they found some collisions were missed and others were misidentified. That such a minor code alteration (which should only affect the lowest-order bits produced by transcendental function evaluations) had such a large high-level effect suggests that their code has significant numerical sensitivities, and results may even be invalid in certain cases. How can they possibly track down where these sensitivities occur, much less correct them?

As another example of this sort, researchers working with an atmospheric simulation code had been annoyed by the difficulty of reproducing benchmark results. Even when their code was merely ported from one system to another, or run with a different number of processors, the computed data typically diverged from a benchmark run after just a few days of simulated time. As a result, it was very difficult even for the developers of this code to ensure that bugs were not introduced into the code when changes are made, and even more problematic for other researchers to reproduce their results. Some divergence is to be expected for computations of this sort, since atmospheric simulations are well-known to be fundamentally “chaotic,” but did it really need to be so bad?

After an in-depth analysis of this code, researchers He and Ding found that merely by employing double-double arithmetic (roughly 31-digit arithmetic) in two key summation loops, almost all of this numerical variability disappeared. With this minor change, benchmark results could be accurately reproduced for a significantly longer time, with very little increase in run time [35].

Researchers working with some large computational physics applications reported similar difficulties with reproducibility. As with the atmospheric model code, they were able to dramatically increase the reproducibility of their results by employing a form of custom high-precision arithmetic in several critical summation loops [47].

5.3 High-precision arithmetic and numerical reproducibility

As noted above, using high-precision arithmetic (i.e., higher than the standard IEEE 64-bit arithmetic ) is often quite useful in ameliorating numerical difficulties and enhancing reproducibility.

By far the most common form of extra-precision arithmetic is known as “double-double” arithmetic (approximately 31-digit accuracy). This datatype consists of a pair of 64-bit IEEE floats \((s, t)\), where \(s\) is the 64-bit floating-point value closest to the desired value, and
is the difference (positive or negative) between the true value and \( s \). “Quad-double” arithmetic operates on strings of four IEEE 64-bit floats, providing roughly 62-digit accuracy. Software to support these datatypes is widely available, e.g., the QD package [36]. Extensions of such schemes can be used to achieve any desired level of precision. Software to perform such computations has been available for quite some time, for example in the commercial packages *Mathematica* and *Maple*. However, until 10 or 15 years ago those with applications written in more conventional languages, such as C++ or Fortran-90, often found it necessary to rewrite their codes. Nowadays there are several freely available high-precision software packages, together with accompanying high-level language interfaces, utilizing operator overloading, that make code conversions relatively painless. A current list of such software packages is given in [8].

Obviously there is an extra cost for performing high-precision arithmetic, but in many cases high-precision arithmetic is often only needed for a small portion of code, so that the total run time may increase only moderately.

Recently a team led by James Demmel of the University of California, Berkeley have begun developing software facilities to find and ameliorate numerical anomalies in large-scale computations. These include facilities to: test the level of numerical accuracy required for an application; delimit the portions of code that are inaccurate; search the space of possible code modifications; repair numerical difficulties, including usage of high-precision arithmetic; and even navigate through a hierarchy of precision levels (32-bit, 64-bit or higher) as needed. The current version of this tool is known as “Precimonious.” Details are presented in [49]. Some related research on numerical reproducibility is given in [44, 43].

5.4 Computations that require extra precision

While it may come as a surprise to some, there is a growing body of important scientific computations that actually *requires* high-precision arithmetic to obtain numerically reliable, reproducible results. Typically such computations involve highly ill-conditioned linear systems, large summations, long-time simulations, highly parallel simulations, high-resolution computations, or experimental mathematics computations.

The following example, condensed from [7], shows how the need for high-precision arithmetic may arise even in very innocent-looking settings. Suppose one suspects that the data \((1, 32771, 262217, 885493, 2101313, 4111751, 7124761)\) are given by an integer polynomial for integer arguments \((0, 1, \ldots, 6)\). Most scientists and engineers will employ a familiar least-squares scheme to recover this polynomial [46, pg. 44].

Performing these computations with IEEE 64-bit arithmetic and the LINPACK routines [1] for LU decomposition, with final results rounded to the nearest integer, finds the correct coefficients \((1, 0, 0, 32769, 0, 0, 1)\) (i.e., \( f(x) = 1 + (2^{15} + 1)x^3 + x^6 \)), but it fails when given the 9-long sequence generated by the function \( f(x) = 1 + (2^{20} + 1)x^4 + x^8 \). On a MacPro system, for example, the resulting rounded coefficients are \((1, 6, -16, 14, 1048570, 2, 0, 0, 1)\), which differ badly from the correct coefficients \((1, 0, 0, 0, 1048577, 0, 0, 0, 1)\).
From a numerical analysis point of view, a better approach is to employ a Lagrange interpolation scheme or the more sophisticated Demmel-Koev algorithm [25]. An implementation of either scheme with 64-bit IEEE-754 arithmetic finds the correct polynomial in the degree-8 case, but even these schemes both fail in the degree-12 case. By contrast, merely by modifying a simple LINPACK program to employ double-double arithmetic, using the QD software [36], all three problems (degrees 6, 8 and 12) are correctly solved without incident. See [7] for additional details.

There are numerous full-fledged scientific applications that also require higher precision to obtain numerically reliable, reproducible results. Here is a very brief summary of some applications that are known to the present authors. See [7] and [8] for additional details:

1. A computer-assisted solution of Smale’s 14th problem (18-digit arithmetic) [53].
2. Obtaining accurate eigenvalues for a class of anharmonic oscillators (18-digit arithmetic) [42].
4. Studies of n-body Coulomb atomic systems (120-digit arithmetic) [30].
5. Computing solutions to the Schrödinger equation (from quantum theory) for the lithium atom [56] and, using some of the same machinery, to compute a more accurate numerical value of the fine structure constant of physics (approx. $7.2973525698 \times 10^{-3}$) (31-digit arithmetic) [57].
6. Computing scattering amplitudes of collisions at the Large Hadron Collider (31-digit arithmetic) [28, 19, 45, 24].
7. Studies of dynamical systems using the Taylor method (up to 500-digit arithmetic) [16, 17, 18].
8. Studies of periodic orbits in dynamical systems using the Lindstedt-Poincaré technique of perturbation theory, together with Newton’s method for solving nonlinear systems and Fourier interpolation (1000-digit arithmetic) [54, 55, 5].

6 High-precision arithmetic in experimental mathematics and mathematical physics

Very high-precision floating-point arithmetic is essential to obtain reproducible results in experimental mathematics and in related mathematical physics applications [20, 21]. Indeed, high-precision computation is now considered one of two staples of the field, along with symbolic computing (see the next section).
Many of these computations involve variants of Ferguson’s PSLQ integer relation detection algorithm [14]. Given an \( n \)-long vector \((x_i)\) of floating-point numbers, the PSLQ algorithm finds the integer coefficients \((a_i)\), not all zero, such that \(a_1x_1 + a_2x_2 + \cdots + a_nx_n = 0\) (to available precision), or else determines that there is no such relation within a certain bound on the size of the coefficients. Integer relation detection typically requires very high precision, both in the input data and in the operation of the algorithm, to obtain numerically reproducible results.

6.1 The BBP formula for \( \pi \)

One of the earliest applications of PSLQ and high-precision arithmetic was to numerically discover what is now known as the “BBP” formula for \( \pi \):

\[
\pi = \sum_{k=0}^{\infty} \frac{1}{16^k} \left( \frac{4}{8k+1} - \frac{2}{8k+4} - \frac{1}{8k+5} - \frac{1}{8k+6} \right).
\]

This formula had not been known previously in the mathematical literature. It was discovered using computations with 200-digit arithmetic.

What is particularly remarkable about this formula is that after a simple manipulation, it can be used to calculate a string of binary or base-16 digits of \( \pi \) beginning at the \( n \)-th digit, without needing to calculate any of the first \( n-1 \) digits [20, pp. 135–143].

Since the discovery of the BBP formula in 1996, numerous other formulas of this type have been found for other well-known mathematical constants, again by employing the PSLQ algorithm and very high-precision arithmetic. For example, two formulas of this type were subsequently discovered for \( \pi^2 \). One permits arbitrary-position digits to be calculated in base 2 (binary), while the other permits arbitrary-position digits to be calculated in base 3 (ternary).

6.2 Ising integrals

Very high-precision computations, combined with variants of the PSLQ algorithm, have been remarkably effective in mathematical physics settings. We summarize here just one or two examples of this methodology in action, following [7] and some related references as shown below.

In one study, high-precision software was employed to study the following classes of integrals [10]. The \( C_n \) are connected to quantum field theory, the \( D_n \) integrals arise in the
Ising theory of mathematical physics, while the $E_n$ integrands are derived from $D_n$:

$$C_n = \frac{4}{n!} \int_0^\infty \cdots \int_0^\infty \frac{1}{(\sum_{j=1}^n (u_j + 1/u_j))^2} u_1 \cdots u_n \, du_1 \cdots du_n$$

$$D_n = \frac{4}{n!} \int_0^\infty \cdots \int_0^\infty \prod_{i<j}^n \left(\frac{u_i - u_j}{u_i + u_j}\right)^2 \frac{1}{(\sum_{j=1}^n (u_j + 1/u_j))^2} u_1 \cdots u_n \, du_1 \cdots du_n$$

$$E_n = 2 \int_0^1 \cdots \int_0^1 \left(\prod_{1 \leq j < k \leq n} \frac{u_k - u_j}{u_k + u_j}\right)^2 dt_2 \cdots dt_n.$$

In the last line $u_k = \prod_{i=1}^k t_i$.

In general, it is very difficult to compute high-precision numerical values of $n$-dimensional integrals such as these. But the $C_n$ integrals can be written as one-dimensional integrals as follows:

$$C_n = \frac{2^n}{n!} \int_0^\infty p K_0^n(p) \, dp,$$

where $K_0$ is the modified Bessel function [2], and such integrals can be evaluated to high precision. For large $n$, these numerical values approach the limit

$$\lim_{n \to \infty} C_n = 0.63047350337438679612204192710\ldots.$$

This numerical value was quickly identified, using the Inverse Symbolic Calculator 2.0 (available at http://carma-lx1.newcastle.edu.au:8087), as

$$\lim_{n \to \infty} C_n = 2e^{-2\gamma},$$

where $\gamma = 0.5772156649\ldots$ is Euler’s constant. This identity was then proven [10]. Some results were also obtained for the $D_n$ and $E_n$ integrals, although the numerical calculations involved there were much more expensive, requiring highly parallel computation [10, 9].

### 6.3 Lattice sums arising from the Poisson equation

One recent application of very high-precision arithmetic (thousands of digits) arose out of attempts to solve the Poisson equation, which, as our colleague Richard Crandall (deceased December 2012) has shown, arises in various contexts such as the analysis of crystal structures and even the sharpening of photographic images. The following lattice sums arise as values of basis functions in the Poisson solutions [11]:

$$\phi_n(r_1, \ldots, r_n) = \frac{1}{\pi^2} \sum_{m_1, \ldots, m_n \text{ odd}} e^{i\pi(m_1r_1 + \cdots + m_nr_n)} \frac{e^{i\pi(m_1^2 + \cdots + m_n^2)}}{m_1^2 + \cdots + m_n^2}.$$
By noting some striking connections with Jacobi theta-function values, Crandall, Zucker, and two of the present authors were able to develop new closed forms for certain values of the arguments \( r_k \) \cite{11}. In particular, after extensive high-precision numerical experimentation, they discovered, then were able to prove, the remarkable fact that for rational numbers \( x \) and \( y \),

\[
\phi_2(x, y) = \frac{1}{\pi} \log A,
\]

where \( A \) is an algebraic number, namely the root of an algebraic equation with integer coefficients.

In this case they computed \( \alpha = A^8 = \exp(8\pi\phi_2(x, y)) \) (as it turned out, the ‘8’ substantially reduces the degree of polynomials and so computational cost), and then generated the vector \( (1, \alpha, \alpha^2, \cdots, \alpha^d) \), which, for various conjectured values of \( d \), was input to a multipair PSLQ program. When successful, the PSLQ program returned the vector of integer coefficients \( (a_0, a_1, a_2, \cdots, a_d) \) of a polynomial satisfied by \( \alpha \) as output. With some experimentation on the degree \( d \), and after symbolic verification using Mathematica, they were able to ensure that the resulting polynomial is in fact the minimal polynomial satisfied by \( \alpha \). Table 1 shows some examples of the minimal polynomials discovered by this process. Using this data, they were able to conjecture a formula that gives the degree \( d \) as a function of \( k \) \cite{11}.

These computations required prodigiously high precision to produce reproducible re-

<table>
<thead>
<tr>
<th>( k )</th>
<th>Minimal polynomial for ( \exp(8\pi\phi_2(1/k, 1/k)) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>( 1 + 52\alpha - 26\alpha^2 - 12\alpha^3 + \alpha^4 )</td>
</tr>
<tr>
<td>6</td>
<td>( 1 - 28\alpha + 6\alpha^2 - 28\alpha^3 + \alpha^4 )</td>
</tr>
<tr>
<td>7</td>
<td>( -1 - 196\alpha + 1302\alpha^2 - 14756\alpha^3 + 15673\alpha^4 + 42168\alpha^5 - 111916\alpha^6 + 82264\alpha^7 - 35231\alpha^8 + 19852\alpha^9 - 2954\alpha^{10} + 7\alpha^{11} )</td>
</tr>
<tr>
<td>8</td>
<td>( 1 - 88\alpha + 92\alpha^2 - 872\alpha^3 + 1990\alpha^4 - 872\alpha^5 + 92\alpha^6 - 88\alpha^7 + \alpha^8 )</td>
</tr>
<tr>
<td>9</td>
<td>( -1 - 534\alpha + 10923\alpha^2 - 342864\alpha^3 + 2304684\alpha^4 - 7820712\alpha^5 + 13729068\alpha^6 - 22321584\alpha^7 + 39775986\alpha^8 - 44431044\alpha^9 + 19899882\alpha^{10} + 3546576\alpha^{11} - 8458020\alpha^{12} + 4009176\alpha^{13} - 273348\alpha^{14} + 121392\alpha^{15} - 11385\alpha^{16} - 342\alpha^{17} + 3\alpha^{18} )</td>
</tr>
<tr>
<td>10</td>
<td>( 1 - 216\alpha + 860\alpha^2 - 744\alpha^3 + 454\alpha^4 - 744\alpha^5 + 860\alpha^6 - 216\alpha^7 + \alpha^8 )</td>
</tr>
</tbody>
</table>

Table 1: Minimal polynomials discovered by very high precision PSLQ computations.
results: up to 20,000-digit floating-point arithmetic in some cases, such as in finding the degree-128 polynomial satisfied by $\alpha = \exp(8\pi \phi_2(1/32, 1/32))$. Related computations in a follow-up study required up to 50,000-digit arithmetic.

In each of these examples, the usage of very high-precision arithmetic is not optional, but absolutely essential to the computation — the relations in question could not possibly be recovered in a reliable, reproducible fashion except by using these exalted levels of precision. Future research in this field will focus not only on how to perform such computations in the most efficient way possible, but also how to explore and then certify that the precision level used in such computations is sufficient to obtain reproducible results.

7 Reproducibility in symbolic computing

Closely related to the high-precision computations described in the past two sections is the usage of symbolic computing in experimental mathematics and mathematical physics. At the present time, the commercial packages Maple and Mathematica are most commonly used for symbolic computations, although other packages, such as Sage and Magma are also used by some.

But like all other classes of computer software, symbolic manipulation software has bugs and limitations. In some cases, the software itself detects that it has encountered difficulties, and outputs messages acknowledging that its results might not be reliable. But not always.

For example, consider the integral

$$W_2 = \int_0^1 \int_0^1 |e^{2\pi ix} + e^{2\pi iy}| \, dx \, dy,$$

which arose in research into the properties of $n$-step random walks on the plane [23, 39]. The latest editions of Maple (version 18) and Mathematica (version 9) both declare that $W_2 = 0$, in spite of the obvious fact that this integral is positive and nonzero. Indeed $W_2 = 4/\pi$ is the expected distance traveled by a uniform random walk in two steps. It is worth pointing out that (2) can easily be rewritten as

$$W_2 = \int_0^1 |e^{2\pi ix}| \left( \int_0^1 |1 + e^{2\pi i(y-x)}| \, dy \right) \, dx = \int_0^1 |1 + e^{2\pi iy}| \, dy,$$

which is in a form that both Maple and Mathematica can evaluate, but this “human” observation is not exploited by the commercial packages. Whether this behavior is a “bug” or a “feature” is matter of some dispute, although it is fair to expect the package to notify the user that the integrand function is problematic.

As another example, three researchers in Spain encountered difficulties with Mathematica while attempting to check a number theory conjecture. They ultimately found errors that they could exhibit even in fairly modest-sized examples. For example, they presented
an example involving a $14 \times 14$ matrix of pseudorandom integers between $-99$ and $99$, which was then multiplied by a certain diagonal matrix with large entries, and then added to another matrix of pseudorandom integers between $-999$ and $999$. When they then attempted to compute the determinant of the resulting fixed matrix using Mathematica version 9, they found that the results were not consistent — they often obtained different answers for the same problem [27]. This problem now seems to have been resolved in Mathematica version 10, according to some tests by the present authors.

Symbolic computing applications often are quite expensive, with jobs running for many hours, days or months. For example, recently three researchers attempted to update an earlier effort to explore Giuga’s 1950 conjecture, which is that an integer $n \geq 2$ is prime if and only if

$$
\sum_{k=1}^{n-1} k^{n-1} \equiv -1 \pmod{n}.
$$

They computationally verified this conjecture for all composite $n$ with up to 4771 prime factors, which means that any counterexample must have at least 19,907 digits. This multithreaded computation required 95 hours. Increasing this bound significantly would currently take decades or centuries [22].

8 Why should we trust the results of computation?

These examples raise the question of why should anyone trust the results of any computation, numeric or symbolic. After all, there are numerous potential sources of error, including user programming bugs, symbolic software bugs, compiler bugs, hardware errors, and I/O errors. None of these can be categorically ruled out. However, substantial confidence can be gained in a result by reproducing it with different software systems, hardware systems, programming systems, or by repeating the computation with detailed checks of intermediate results.

For example, programmers of large computations of digits of $\pi$ have for many years certified their results by repeating the computation using a completely different algorithm. If all but a few trailing digits agree, then this is impressive evidence that both computations are almost certainly correct. In a similar vein, programmers of large, highly parallel numerical simulations, after being warned by systems administrators that occasional system hardware or software errors are inevitable, are inserting internal checks into their codes. For example, in problems where the computation carries forward both a large matrix and its inverse, programmers have inserted periodic checks that the product of these two matrices is indeed the identity matrix, to within acceptable numerical error.

In other cases, researchers are resorting to formal methods, wherein results are verified by software that performs a computer verification of every logical inference in the proof, back to the fundamental axioms of mathematics. For example, University of Pittsburgh
researcher Thomas Hales recently completed a computer-aided proof of the Kepler conjecture, namely the assertion that the most efficient way to pack equal-sized spheres is the same scheme used by grocers to stack oranges [31]. Some objected to this approach, so Hales has attempted to re-do the proof using formal methods [32]. This project is now complete [33].

8.1 Is “Free” Software Better?

We conclude by briefly commenting on open-source versus commercial software (a full treatment would require discussion of complexities such as licensing, user adoption, and numerous other topics). Some open-source software for mathematical computing is quite popular. For example, GeoGebra, which is based on Cabri, is now very popular in schools as replacement for Sketchpad. But the question is whether such software will be preserved when the core group of founders and developers lose interest or move on to other projects or employment. This is also an issue with large-scale commercial products, but it is more pronounced with open-source project.

One advantage of Maple over Mathematica is that most of the Maple source code is accessible, while Mathematica is entirely sealed. As a result, it is often difficult to track down or rectify problems that arise. Similarly, Cinderella is very robust, unlike GeoGebra, and mathematically sophisticated—using Riemann surfaces to ensure that complicated constructions do not crash. That said, it is the product of two talented and committed mathematicians but only two, and it is only slightly commercial. In general, software vendors and open source producers do not provide the teacher support that has is assumed in the textbook market.

One other word of warning is given by the experience of the Heartbleed bug [3], which many cybersecurity observers termed “catastrophic.” This is a bug in the OpenSSL cryptography library, which is incorporated in many other software packages, commercial and open-source. A fixed version of the OpenSSL was quickly released, but it will take time to identify all instances. In this case, although almost everyone uses the flawed software, no one really “owns” it, making ongoing maintenance problematic.

9 Conclusions

The advent of high-performance computer systems, all the way from laptops and workstations to systems with thousands or even millions of processing elements, has opened new vistas to the field of scientific computing and computer modeling. It has revolutionized numerous fields of scientific research, from climate modeling and materials science to protein biology, nuclear physics and even research mathematics (as we have seen in the examples above). In the coming years, this technology will be exploited in numerous industrial and engineering applications as well, including “virtual” product testing and market analysis.
However, like numerous other fields that are embracing “big data,” scientific computing must confront issues of reliability and reproducibility, both in computed results and also in auxiliary statistical analysis, visualization and performance. For one thing, these computations are becoming very expensive, both in terms of the acquisition and maintenance costs of the supercomputers being used, but even more so in terms of the human time to develop and update the requisite application programs.

Thus it is increasingly essential that reliability and reproducibility be foremost in these activities: designing the workflow process right from the start to ensure reliability and reproducibility — carefully documenting each step, including code preparation, data preparation, execution, post-execution analysis and visualization, through to the eventual publication of the results. A little effort spent early on will be rewarded with much less confusion and much more productivity at the end.

References


152, 2005.

research in computational harmonic analysis. *Computing in Science and Engineering*,
11, January 2009.

[27] A. J. Duran, M. Perez, and J. L. Varona. Misfortunes of a mathematicians’ trio using


symplectic integrators for the solar system. *Celestial Mechanics and Dynamical


2005.

ThomasHales.pdf.

flyspeck/wiki/AnnouncingCompletion.


[35] Y. He and C. Ding. Using accurate arithmetics to improve numerical reproducibility


[38] J. P. A. Ioannidis. Why most published research findings are false. *PLOS Medicine*,
August 2005. DOI: 10.1371/journal.pmed.0020124.


