Improved performance of QCD code on ALiCE

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We present results for the performance of QCD code on ALiCE, the Alpha-Linux Cluster Engine at Wuppertal. We describe the techniques employed to optimise the code, including the metaprogramming of assembler kernels, the effects of data layout and an investigation into the overheads incurred by the communication.

1. Introduction

In a typical lattice QCD project the total runtime of code on a supercomputing platform is often measured in months or even years. This means that even a modest improvement in the performance of the code can yield very tangible benefits. There are two aspects to the optimisation of code for parallel machines: single-node optimisation and the minimisation of the overhead incurred by inter-node communications.

The former requires that the code be written to take full advantage of the high performance available from today's advanced hardware. The latter is of particular importance on cluster machines, like ALiCE, where the scalability of code can be a serious problem.

2. Single node optimisation

Experience tells us that the dominant part of a typical lattice QCD code is that implementing the multiplication of a vector by the fermion matrix so it is here that the effort should be made. Secondly, the use of hand-coded optimised assembler routines can dramatically improve performance since the programmer can use information about the code which is unavailable to the compiler.

The disadvantage with assembler routines is that they are difficult to develop and harder to maintain, in addition to the obvious lack of portability. We address these problems by adopting a metacoding approach; writing a C++ program to write the assembler code for us. We have developed special software tools to enable this.

![Figure 1. Wilson matrix multiplication in single (above) and double (below) precision. The vertical lines indicate the volumes at which the data fills the level 1 (L1) and level 2 (L2) caches.](image-url)
2.1. Metacode software toolkit

The first stage in creating the assembler routine is to reduce the computational task to elementary assembler-level abstract instructions, e.g. load a datum from memory into registers, perform arithmetic on the data, cache management, etc.

In order to write the metacode we have developed a system of C++ classes and routines which automatically schedule the instructions to hide the instruction latencies as much as possible and automatically manage the register usage.

When the metacode written using these routines is compiled and run, the abstract instructions with their arguments are translated into an actual assembly language and written to a file.

By basing the toolkit design on an abstract RISC ISA it should be possible to produce assembler code for any RISC machine by changing the architecture-dependent parameters. Here we show the results on ALiCE, a cluster of Compaq DS10 servers which have a 616 Mhz, 4-way superscalar Alpha 21264 processor with a 64Kb 2-way set-associative level 1 (on-chip) data cache and a 2Mb level 2 (off-chip) cache.

2.2. QCD kernels

An advantage of the metacode toolkit as that it permits a large degree of flexibility in writing various assembler kernels; different approaches can be tried and compared, and the kernels can be rewritten to adapt to changes in the action or algorithm.

Figure 1 shows the improvement, over a wide range of lattice volumes, in the performance of the Wilson matrix multiplication routine when written with assembler kernels over that of the original implementation in C. To demonstrate the effect in a more realistic environment, the inversion of the Wilson matrix using BiCGStab is shown in figure 2.

3. Cluster performance

ALiCE is clustered using ParaStation 3 over 64bit/33MHz Myrinet. Our code uses MPICH 1.2.3 to do the communications.

We test the multinode performance of the BiCGStab solver on a $16^d$ lattice running on $n = 1, 2, 4, 8$ and 16 nodes arranged in a 1-dimensional $(1 \times n)$ grid and a 2-dimensional (square) grid. We use a standard metric of parallel performance:

$$\text{speedup} = \frac{\text{speed on } n \text{ nodes}}{\text{speed on 1 node}}$$

Our original implementation used a conventional array ordering for all the fields, where each lattice site with coordinates $(x_0, x_1, x_2, x_3)$ is numbered $n = x_3 + N_3 x_2 + N_2 x_1 + N_1 x_0$ where $N_\mu$ is the size of the local lattice in direction $\mu$. This is illustrated in figure 3 (left) which shows that while the data along the boundary in one direction is contiguous, in the second direction it is strided. Investigations into the performance of our MPI communications suggest that the communication of strided data introduces an overhead of at least 20% compared to contiguous data. This explains the poor scaling of the solver on a 2-dimensional grid shown in figure 4. The scaling on the 1-dimensional grid suffers from the increasingly unfavourable surface-to-volume ratio of the local lattice.

The solution to these problems appears to be to rearrange the data layout so that the sites on
the lattice boundaries are ordered in a contiguous fashion, illustrated in figure 4 (right).

Separating the boundary and interior sites in this way has the additional advantage that computation can proceed on the interior sites while the boundary sites are waiting for a non-blocking communication to finish. Figure 4 shows that using this new data layout greatly improves the speedup of the solver. The new data layout does not adversely affect single node performance.

4. Summary

We have introduced a flexible software toolkit [1] which can successfully generate optimised assembler routines for performance-critical parts of our lattice QCD code. On a single node we see a 100–150% improvement in the Wilson matrix solver performance at single precision and 50–100% at double precision.

We demonstrate that good scaling performance can be achieved on ALiCE if the data layout and communication strategy is carefully adapted to suit the communication needs.

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References

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