PARALLEL COMPUTING FOR QCD ON A PENTIUM CLUSTER

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Motivated by the computational demands of our research and budgetary constraints which are common to many research institutions, we built a “poor man’s supercomputer”, a cluster of PC nodes which together can perform parallel calculations at a fraction of the price of a commercial supercomputer. We describe the construction, cost, and performance of our cluster.

1 Introduction

The lattice field theory group at the Zhongshan University physics department is in a period of rapid development. The group’s interests involve such topics as finite density QCD, Hamiltonian Monte Carlo lattice field theory, lattice supersymmetry, and lattice quantum gravity. All of these topics can be studied through Monte Carlo simulation, but can be quite costly in terms of computing power. In order to do large scale numerical investigations of these topics, we required a corresponding development of our local computing resources. We decided building a PC cluster capable of parallel computing was the most economical way to build computing power. This type of cluster is often termed a “Beowulf Cluster” and was pioneered by the United States’ National Aeronautics and Space Administration. The Beowulf Project’s website also proved to be a valuable resource in the construction of our cluster.

2 Construction

2.1 Hardware

One big advantage of a PC cluster over other types of supercomputers is the low cost and easy availability of the hardware components. All the hardware in our cluster is available at retail computer suppliers. This gives us great flexibility in both building the cluster and in any future upgrades or expansions we may choose to make.

At the present our cluster consists of ten PCs, each one has two 500MHz Pentium III processors. Additionally, each has 128MB of RAM, an 8GB EIDE hard disk, a 100Mbit/sec Ethernet card, a CDROM, a floppy drive and a basic graphics card. In practice the CDROM, the floppy drive, and even the graphics card could be considered extraneous, as all interaction with the nodes could be done through the network. However, with these components, all of which are relatively cheap in comparison to the total cost, the operating system installation and occasional maintenance is significantly easier. One node has a larger hard disk (20GB) and a SCSI adaptor, for communication with a tape drive for disk backups. The entire cluster shares one monitor, mouse and keyboard.

A 100 Mbit/s fast Ethernet switch handles the inter-node communication. The switch has 24 ports so the cluster is expandable to a total of 48 processors using the current scheme. Of course it is possible to link multiple switches or use nodes with more than two processors, so the possibilities for a larger cluster are nearly boundless.

2.2 Software

The cluster runs on the Linux operating system. The reasons for choosing Linux are manifold. It is cheap. It can easily support multiple users. It easily supports network file systems (shared hard disks) and allows user accounts to be shared across the cluster. Furthermore, compilers for C, C++, and Fortran are free.

To operate the cluster as a parallel computer, the programmer must design the algorithm so that it appropriately divides the task among the individual processors. He or she must then include appropriate message passing functions in the code which allow information to be sent and received by the various processors. We use MPI (Message Passing Interface), one of the most popular message passing standards. By adopting such a widely used standard we are able to share C, C++, and Fortran programs with other colleagues who may be using any of a large variety of computing platforms.

3 Performance and Cost

3.1 General Cost Comparison

The LINPACK benchmarking test on a single processor shows that a single 500MHz Pentium processor has a peak speed of about 100 Mflops (100 million floating point operations per second). The LINPACK benchmark, a widely used standard, is used to compare the performance of different computer systems. A high LINPACK score indicates that the computer is able to perform more floating point operations per second than other computers.

With the LINPACK benchmark, the cluster is capable of performing parallel calculations at a fraction of the price of a commercial supercomputer. The total cost of the cluster is significantly lower than that of a commercial supercomputer. The operating system installation and occasional maintenance is significantly easier. One node has a larger hard disk (20GB) and a SCSI adaptor, for communication with a tape drive for disk backups. The entire cluster shares one monitor, mouse and keyboard.

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point operations per second). The peak speed for a cluster of 20 such processors therefore approaches 2 Gflops.

The cost of our cluster was a little under US$14000 in 1999. This gives an approximate cost of US$7 per MFlop. For comparison we can examine the cost of a commercially produced supercomputer. The Cray company offers a starring model of its T3E-1200E supercomputer cluster for US$63000. This includes six processors, each capable of processing at a peak speed of 1200 Mflops. The cost per MFlop on the T3E-1200E machine is therefore US$87.50. We have constructed a machine that is an order of magnitude cheaper per MFlop.

We should note here that six of our two-processor nodes are not equivalent to a single Cray node. One of the slowest parts of a parallel computation is the communication. Fewer faster nodes always require less communication and the communication channels are inherently faster on the Cray machine. However, some problems are by their nature easier to divide into nearly independent parts. These problems require less communication and the penalty for a system with slower communication or more nodes is smaller. It is for just these types of problems that a PC cluster such as ours is particularly well suited. One can take full advantage of the processing power. For most easily parallelizable problems, a PC cluster seems to be the most economical solution for way of providing computing power.

3.2 QCD Benchmarking

In particular, we are interested in using for Monte Carlo simulations of lattice quantum chromodynamics (QCD). Lattice QCD simulations are well suited for parallelization as they involve mostly local calculations on a lattice. The algorithm can conveniently divide the lattice and assign the sections to different processors. The communication between the nodes therefore is not extremely large.

We have tested the performance of our cluster in actual lattice QCD simulations. Hioki and Nakamura provide comparison performance data on SX-4 (NEC), SR2201 (Hitachi), Cenju-3 (NEC) and Paragon (Intel) machines. Specifically, we compare the computing time per link update in microseconds per link and the inter-node communication speed in MB/sec. The link update is a fundamental computational task within the QCD simulation and is therefore a useful standard. The test was a simulation of improved pure gluage lattice action (1 x 1 plaquet and 1 x 2 rectangle terms) on a 16^4 lattice. In each case the simulation was run on 16 processors. We used β = 6.0

We used the QCDimMPI Fortran code. Table shows the results of this testing.

| Machine          | µ-sec/link | MB/sec |
|------------------|------------|--------|
| SX-4             | 4.50       | 45     |
| SR2201           | 31.4       | 28     |
| Cenju-3          | 57.42      | 8.1    |
| Paragon          | 149        | 9.0    |
| ZSU’s Pentium cluster | 7.3      | 11.5   |

4 Conclusions

We have reported on our efforts to build a parallel computing facility that fits the demands and budget of a developing lattice field theory group. We feel that a PC cluster can provide a very flexible and extremely economical computing solution that is able to run parallel programs written using the most popular message passing standard, MPI. Furthermore we believe that this may in fact be the first such cluster at an academic physics institution in the People’s Republic of China.

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