Introduction to large-scale computing 大規模計算序論

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Contents

- Necessity of large-scale computing
- Methods of the large-scale computing(Mainly parallelization)
- Today's supercomputers
- Future direction of large-scale computing

Necessity of large-scale computing

- Systems we want to study have many atoms.
- The computational cost increases N^3 , N^6 , or exponentially.(N is the number of atoms, electrons,...)
- Some physical quantities needs a huge computational cost in spite of a small system(high accuracy).
- There is scientific and/or industrial significance obtained by large-scale computing.

Problems raised by large-scale computing

- Because of huge date must be handled, huge HDD and fast IO is needed. (~30PB in K computer, ~150PB in Fugaku(post-K))
- The problem of data analysis including visualization.
- In some cases, data analysis costs more time than computational time.

• ...

When we perform large-scale computing,...

- It costs huge computational time.
- In some cases, huge physical memory of CPU is needed.
- In order to solve these problems, we have to improve software codes to reduce the computational time and memory.

Parallel computing is definitely necessary.

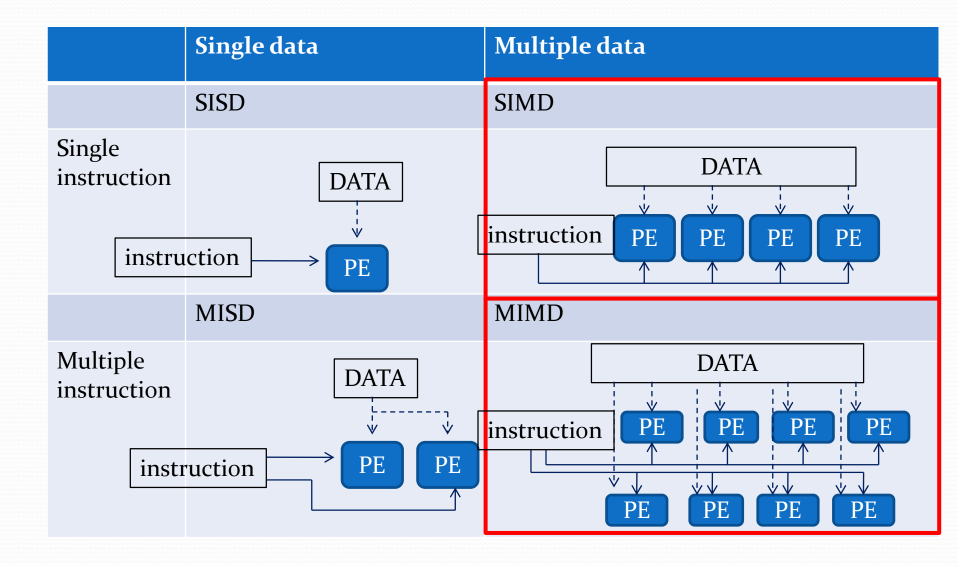
Methods for large-scale computing

- Speed-up techniques
 - Vectorization→SIMD(many cores)
 - Parallelization
 - Order-*N* method(reduce the cost $o(N^p) \rightarrow o(N)(p=1)$)
 - Use of specialized machine or accelerator(GPU).
- These are used in a code depending on one's necessity.

The easiest way to speed up is to be able to realize it by choosing proper compile options(vectorization, Automatic parallelization, optimization,...).

Since it is not possible to speed up the code automatically, we have to tune the code significantly.

Concept of Flynn's taxonomy



Vectorization

- A vector processor, or array processor, is a CPU that implements an instruction set containing instructions that operate on one-dimensional arrays of data called vectors(row, column, or diagonal elements, etc...).
- Vectorization is the method that we program a code to construct regularly-arrayed data, and process the data at a time by the vector CPU.
- Vectorization realizes the reduction of a CPU time.
- Today, this concept is included in SIMD.

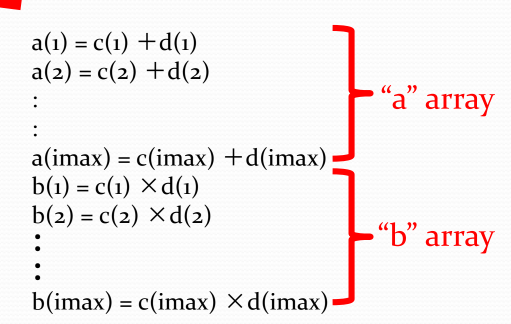
Vector processing

do i=1,imax a(i) = c(i) + d(i) $b(i) = c(i) \times d(i)$ enddo

Scalar processing

We tune to increase the part being processed at the deepest do-loop in multi do-loop.

Vector processing



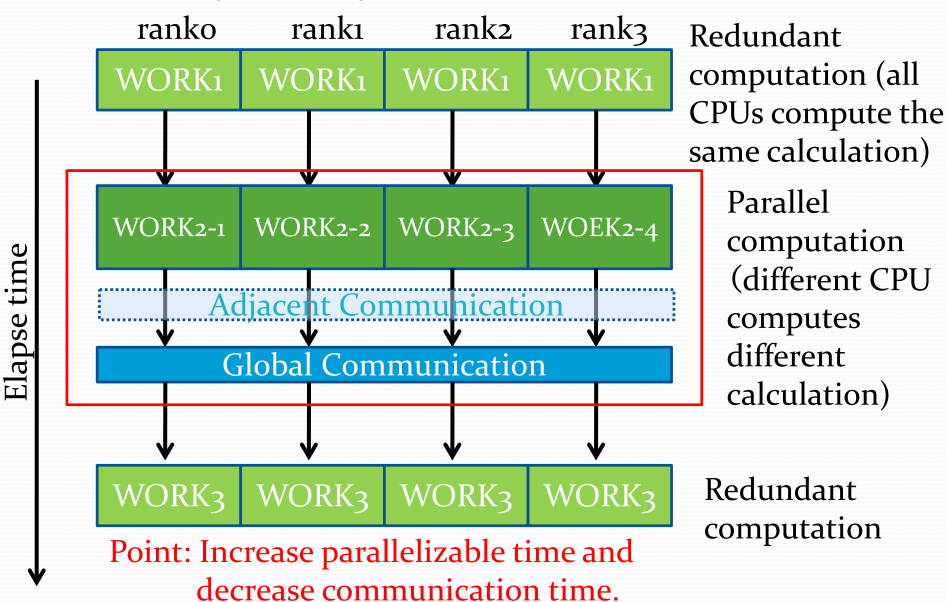
Order-N

- Realized by algorithm
 - →No approximation Ex.: Screened-KKR
- Realized by approximation of Hamiltonian
 - →Approximation by localization of interaction. This type of methods were actively studied in 90's.
 - Roughly speaking, matrices are block-diagonalized or including similar method.
 - $N \times N \rightarrow N \times N_{local}$ (fixed). If N increases, N_{local} not increase. (like a tight-binding approximation)
 - The accuracy of the approximation can be adjustable depending on a required accuracy.
 - One of the famous method is a Divide and conquer method (分割統治法). This is used in "OpenMX(Ozaki)", "DC(Kobayashi and Nakai)", CONQUEST (Bowler, Miyazaki, Gillan)...

Parallelization

- This is the method to shorten the elapse time by sharing task with more than one CPU without changing the processing time of the task.
- This is used to share big memory(array).
- SIMD(Single instruction, multiple data),
 MIMD(Multiple instruction, multiple data)

Concept of parallelization



Problems in parallelization

- The ratio of parallelizable part in an elapse time on 1CPU sequential job is conclusive (Amdahl' law).
- When we parallelize a software code, the data communication between nodes must occur. If the number of nodes increases, the communication time between nodes costs more than the computational time.(In the case of classical MD simulation, this is a crucial problem.)

Amdahl's Law

Speed up ratio
$$\alpha_P \equiv \frac{t_1}{t_{N_{\rm core}}}$$
 Elapse time of one job on 1 core that $t_{N_{\rm core}}$ Elapse time of one job on N core

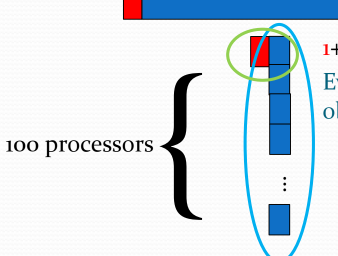
$$(\alpha_P)_{\text{max}} = t_1 \times \left(\frac{1}{t_{N_{\text{core}}}}\right)_{\text{max}} = \frac{1}{(1-P) + \frac{P}{N_{\text{core}}}}$$

P: parallelizable time in a sequential job

Upper bound of parallelizable ratio

For examples,

Not parallelizable (1 sec.) parallelizable (99 sec.)



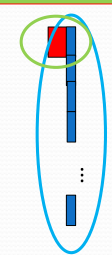
1+0.99=1.99 sec.(almost 50 times faster)

Even w obtain We have to increase the parallelizable part as possible.

We have to decrease the non parallelizable part as possible.

1000 processors

We feel that the merit of parallelization does not increase even if the number of processors increases more and more.



1+0.099=1.099 sec. (almost 91 times faster)
Although we uses 1000

Although we uses 1000 processors, the efficiency does not become 100 times even!!!

Gustafson-Barsis' law

Speed-up ratio $\leq n + (1-n)s$

n: the number of cores

s: the ratio of sequential computation(this part cannot parallelize.)

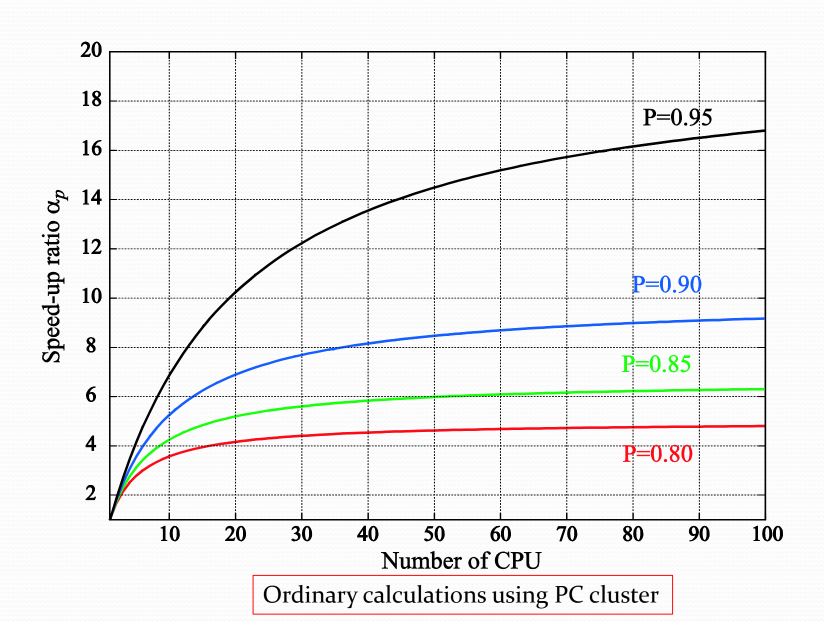
In this law, the communication time among nodes does not considered.

Indices of parallelization efficiency

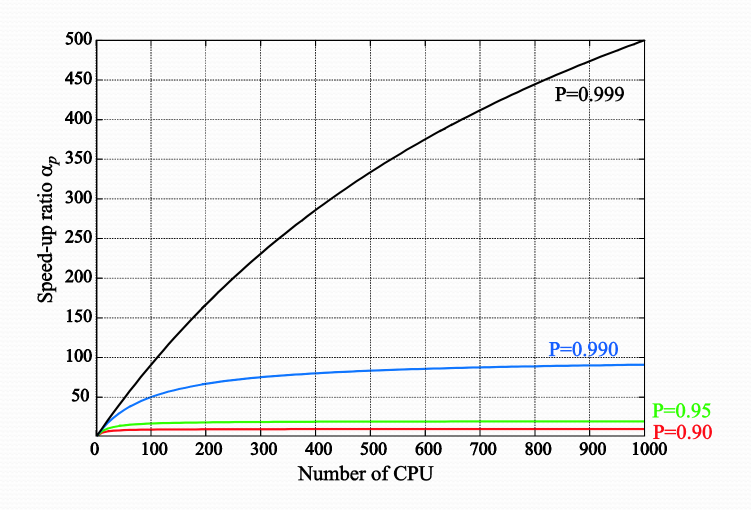
- Strong scaling—The overall problem size (number of atoms or electrons etc...) is fixed and the number of processors is increased(based on Amdahl's law).
- Weak scaling The problem size per processor is fixed and the overall problem size is increased with increasing processors(based on Gustafson's law). The processing of one processor is not changed. The ideal situation is the computation time should be constant even if the increase of the number of processors. If the computation time increases, we can see the non parallel part remains, and if the adjacent communication time increases, the method of the communication has a problem.

Reference: http://www.r-ccs.riken.jp/library/event/tokuronB_180406.html

Example of Amdahl's law(1)

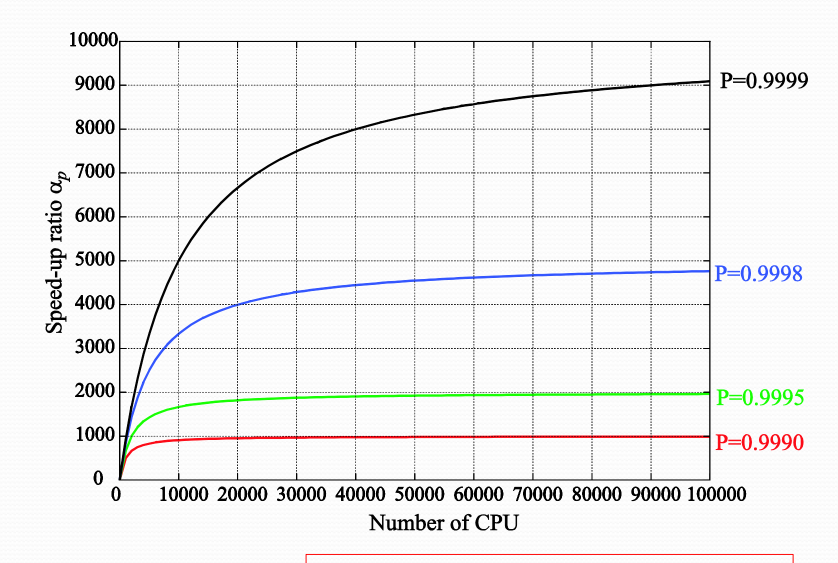


Example of Amdahl's law(2)



Calculations using high-end PC cluster or supercomputers

Example of Amdahl's law(3)

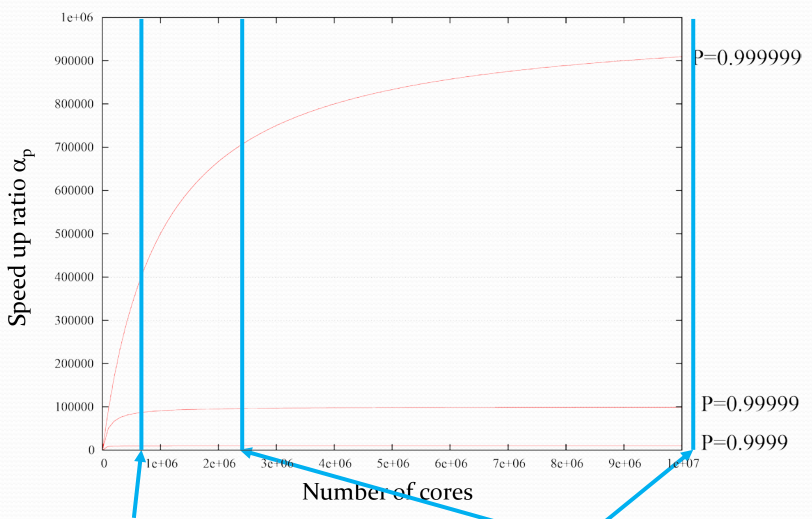


K computer: 705,024 cores

Calculations using high-end supercomputers

Example of Amdahl's law(4)

Calculations using top class supercomputers in the world



K computer: 705,024 cores

Summit. 2,397,824 cores

Sunway TaihuLight: 10,649,600 cores

Even if you are just a user,

- you need to be aware of the parallelization efficiency of the software you use because you must effectively use machine time(machine point, or cost(usage fee)) you can use.
- you need to understand the characteristics of the computer and make efforts to increase the calculation efficiency.
- you need to choose the best compile option to make executable file.
- you need to choose the best parameters for the best performance of the software to obtain the best scientific results.

Parallel programming languages

- MPI(message passing interface)
 - The most versatile method.
 - Both core-to-core and node-to-node communication can be handled.
 - The programming may not be easy.
- OpenMP
 - Only core-to-core communication.
 - The programming may be easier than that of MPI.

PVM

• HPF(High Performance Fortran)

(within non-complex case)

Now, these are seldom used.

Example of parallelization with MPI

```
program main
include (mpif.h)
call MPI_INIT(IERR)
call MPI_COMM_SIZE(MPI_COMM_WORLD,NPROCS,IERR)
call MPI_COMM_RANK(MPI_COMM_WORLD,MYRANK,IERR)
                                                        ista, iend are the first array
call para_range(1, n, nprocs, myrank, ista, iend)+
                                                        numbers allocated to each
do i=ista, iend
                                                        processor.
                                         Each processor has only allocated elements of
  a(i)=a(i)+value
                                         the array a().
enddo
call MPI_ALLREDUCE(a,a1,n_element,
                                                     Sum up all elements of the array a()
   & MPI_DOUBLE_PRECISION,
                                                     for all processors and distributed it
                                                     for all processors (all processors have
   & MPI_SUM, MPI_COMM_WORLD, IERR)
                                                     the same value of a()).
call MPI_FINALIZE(IERR)
                              It is necessary to rewrite so as to calculate only the part
                              allocated to each node and to communicate the data.
end
```

Execution: mpirun (or mpiexec) –np 4(# of parallel) ./a.out

Example of parallelization with OpenMP

```
program main
implicit none
integer omp_get_thread_num,I
double precision z(100), a, x(100), y
do i= 1, 100
  z(i) = 0.0
  x(i) = 2.0
 end do
                                            code.
a = 4.0
y = 1.0
print *, "Welcome to the parallel world"
!somp parallel
print *, "Excuted on", omp_get_thread_num()
call daxpy(z, a, x, y)
!somp end parallel
end program main
subroutine daxpy(z, a, x, y)
integer I
double precision z(100), a, x(100), y
!$omp parallel do
                                            Do parallel of just below do loop
do i = 1, 100
  z(i) = a * x (i) + y
enddo
return
```

end

For simple parallelization, we only need to insert directives, so if we compile without parallelization options the directives are commented out, so basically we do not need to change the original

Directive of the start of parallel processing

Directive of the end of parallel processing

Execution: We do not use a special command. Set environmental value "OMP NUM THREADS=4" (# of parallel)

The difference between MPI and OpenMP

- OpenMP can be parallelizable only for cores which share the memory. We cannot make a large-scale parallelization code by using OpenMP only.
- The typical software codes which correspond to the large-scale parallelization are tuned by using MPI for its main part.
- The role of OpenMP is to assist the speed up of the code which is parallelized by using MPI.
- Even if you do not develop the software code, you have to know the details of the code in order to get maximum performance.

The other environments to develop parallel computing codes using GPU

CUFFT, CUBLAS,...

easy

- Only call libraries to accelerate specific parts. The other parts do not be accelerated.
- OpenACC
 - a programming standard for parallel computing developed by Cray, CAPS, NVIDIA and PGI, designed to simplify parallel programming of heterogenious CPU/GPU system.
 - Very similar with OpenMP. In near future, OpenMP and OpenACC may be merged.
 - Fortran and C are supported.
- CUDA

hard

- a parallel computing platform and programming model created by NVIDIA and implemented by the graphics processing units (GPUs) that they produce. (only PGI Fortran can used.))
- OpenCL
 - a framework for writing programs that execute across heterogeneous platforms consisting of central processing units (CPUs), graphics processing units (GPUs), DSPs and other processors.

OpenACC(Open Accelerator)

```
program picalc
   implicit none
   integer, parameter :: n=1000000
   integer :: i
   real(kind=8) :: t, pi
   pi = 0.0
 !$acc parallel loop
   do i=0, n-1
    t = (i+0.5)/n
    pi = pi + 4.0/(1.0 + t*t)
   end do
 !$acc end parallel loop
   print *, 'pi=', pi/n
 end program picalc
```

As the development environment, the best results can be obtained by improving according to the message issued by the compiler.

http://www.nvidia.co.jp/object/openacc-gpu-directives-jp.html http://www.cms-initiative.jp/ja/events/2014-haishin http://www.r-ccs.riken.jp/library/event/tokuronB_180406.html (The 14th lecture given by Dr. A. Naruse is useful.)

The cutting-edge techniques

- Avoid memory wall problem(cache control)
 - Computing power>data transfer from memory to computing unit(CPU)→reuse the data on caches
- Pipeline processing
- Continuous access in do loop
- loop unrolling
- Divide data into blocks
- Use the highly-optimized libraries
- ...

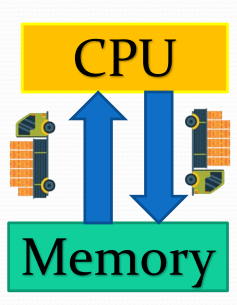
We have to know details of hardware to make highly optimized software codes.

Please see these sites; (only Japanese)

http://www.r-ccs.riken.jp/library/event/tokurona_170406.html http://www.r-ccs.riken.jp/library/event/tokuronB_180406.html

Memory wall problem

Old computer

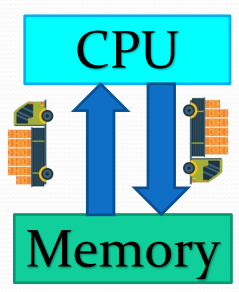


Computation performance (number of clocks) was slow (~10MHz)

The memory can not be accessed until a certain time passes

Sending data from memory to the CPU was slow, and the computation speed was also slow, so it was not the problem at "data transfer speed ≈ calculated speed".

Today's computer



The computation performance becomes extremely fast (~a few GHz)

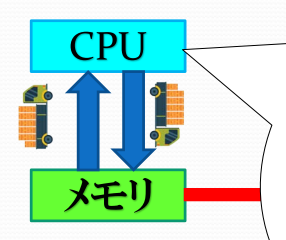
The fact that memory cannot be accessed only for a fixed time does not change much.

Sending data from the memory to the CPU is slow, but since the computing speed has become extremely fast, the CPU cannot wait "the transfer speed << compute speed" (supplement with cache etc...).

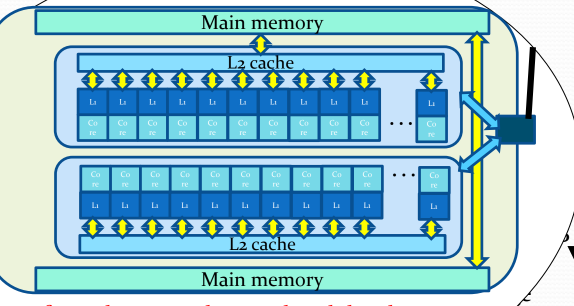
Memory wall problem

Impotant!: Data transfer is the most expensive at all levels.

Today's computer



Once the data is sent from the CPU, the code is program. it as much as possible and reduc number of calls to the memory as ma possible. Data transfer between the memory and the register or cache is fast, but the capacity is small.



If you do not understand and develop a complex CPU configuration, the performance will not be achieved transfer from

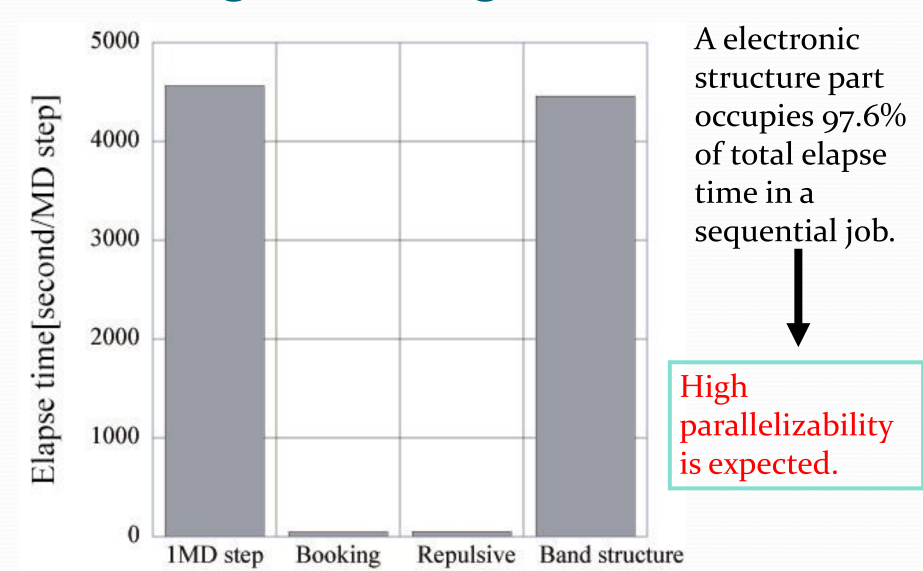
> the mam memory to the CPU is 100 times longer than access to the data on the register.

pacity

High performance Python

- In place of Fortran and C / C ++, which have been used in scientific computing in the past, HPC software development has been advanced in the USA by using Python.
- Python has overwhelming code readability compared to Fortran and C / C ++.
- →It is more likely that the maintenance of the code will be continued without reading by experts.
- We do not need write everything in Python. If we know that another language is faster than Python for a certain part, we should use the part in the original. We combine Python and the other language and improve the ease of maintenance. (It is said that Python is a "glue language" because it is a language that "sticks" programs in multiple languages.)

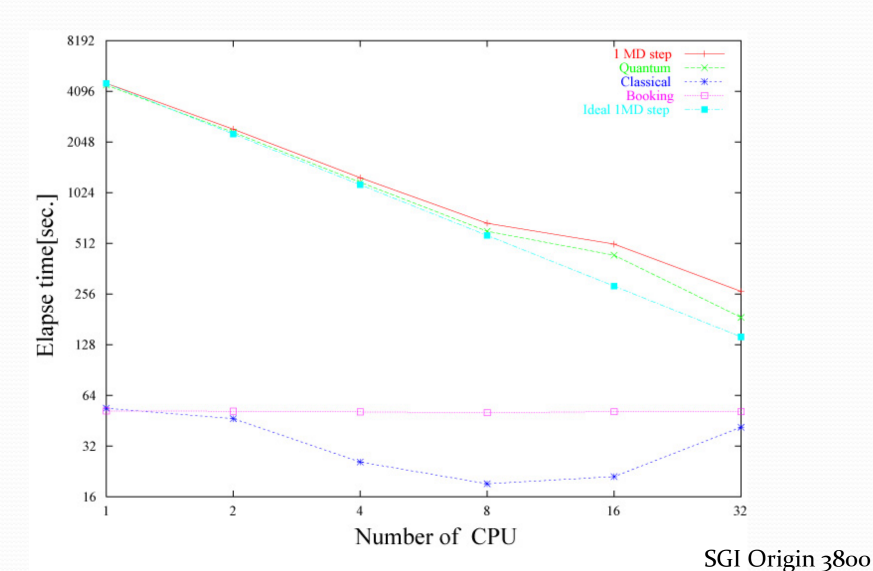
Analysis of the computational time in order-N tight-binding method



Parallelization by means of MPI

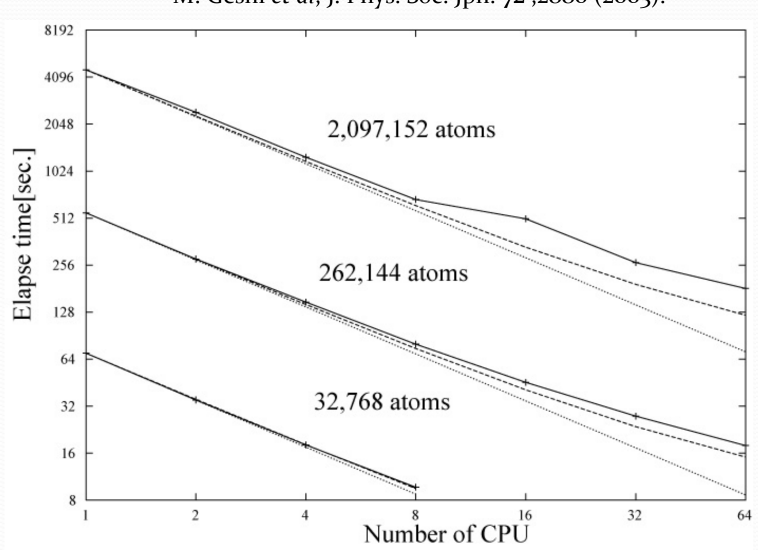
- We used MPI_ALLREDUCE and MPI_BCAST to communicate data. (Although this is quite simple and includes useless data communication, the speed-up ratio was good.)
- We divided do-loop as outer as possible.
- In this method, we can make all matrix elements as long as we determine the atomic positions. These data determined from 1 MD step are stored in CPU memory of all node. Although almost all data are useless, we do not need to communicate the data among nodes and there is no loss of data communication.

Elapse time of each part as a function of the number of CPU



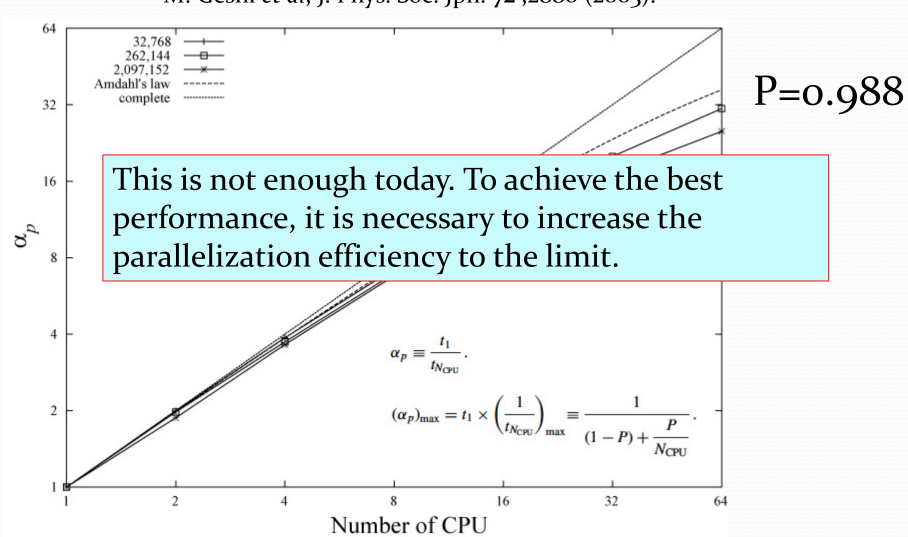
Elapse time as a function of the number of CPU

M. Geshi et al, J. Phys. Soc. Jpn. 72,2880 (2003).



Parallelization ratio

M. Geshi et al, J. Phys. Soc. Jpn. 72,2880 (2003).



Today's supercomputers

Classes of parallel computers

- Multicore computing
 - a processor that includes multiple execution units ("cores") on the same chip.
- Symmetric multiprocessing
 - a computer system with multiple identical processors that share memory and connect via a bus.
- Distributed computing
 - a distributed memory computer system in which the processing elements are connected by a network. (Cluster computing, Massive parallel processing, Grid computing)
- Specialized parallel computers
 - Within parallel computing, there are specialized parallel devices that remain niche areas of interest. (GPGPU,Application-specific integrated circuits, Vector processors)

The latest top 500 supercomputer

list (2018.11)

IBM



http://www.top5oo.org/

GPU

GPU

Designed for data and AI

Intel Xeon(many-cores)

GPU

Intel Xeon(many-cores

GPU

By the way, K-computer is 18, Oakforest PACS is 14.

GPU



The latest top 500 supercomputer list (2019.6)(Latest)





IBM製





United States



http://www.top5oo.org/

GPU

GPU

By the way, Oakforest-PACS is 16, and K-computer is 20. (The usage of the K-computer stopped at August 16, and it was shutdown at the end of August.)

GPU

Intel XeonPhi
(This architecture was gone.)
GPU

GPU

Linpack

- A software library for performing numerical linear algebra on digital computers.
- It is used for the performance assessment of supercomputers. In practice, High-Performance Linpack(HPL) is used.
- It makes use of the BLAS libraries for performing basic vector and matrix operations.
- The benchmark used in the LINPACK Benchmark is to solve a dense system of linear equations.

This benchmark shows nothing more than one side of performance of supercomputers.

HPC Challenge

The HPC Challenge benchmark consists of basically 7 tests:

- HPL the Linpack TPP benchmark which measures the floating point rate of execution for solving a linear system of equations.
- DGEMM measures the floating point rate of execution of double precision real matrix-matrix multiplication.
- STREAM a simple synthetic benchmark program that measures sustainable memory bandwidth (in GB/s) and the corresponding computation rate for simple vector kernel.
- PTRANS (parallel matrix transpose) exercises the communications where pairs of processors communicate with each other simultaneously. It is a useful test of the total communications capacity of the network.
- RandomAccess measures the rate of integer random updates of memory (GUPS)
- FFT measures the floating point rate of execution of double precision complex one-dimensional Discrete Fourier Transform (DFT).
- Communication bandwidth and latency a set of tests to measure latency and bandwidth of a number of simultaneous communication patterns; based on b_eff (effective bandwidth benchmark).

http://www.hpcchallenge.org/

HPCG Benchmark(From 2014)

The High Performance Conjugate Gradients (HPCG) Benchmark project is an effort to create a new metric for ranking HPC systems. HPCG is intended as a complement to the High Performance LINPACK (HPL) benchmark, currently used to rank the TOP500 computing systems. The computational and data access patterns of HPL are still representative of some important scalable applications, but not all. HPCG is designed to exercise computational and data access patterns that more closely match a different and broad set of important applications, and to give incentive to computer system designers to invest in capabilities that will have impact on the collective performance of these applications.

HPCG is a complete, stand-alone code that measures the performance of basic operations in a unified code:

- •Sparse matrix-vector multiplication.
- Vector updates.
- •Global dot products.
- Local symmetric Gauss-Seidel smoother.
- •Sparse triangular solve (as part of the Gauss-Seidel smoother).
- •Driven by multigrid preconditioned conjugate gradient algorithm that exercises the key kernels on a nested set of coarse grids.
- •Reference implementation is written in C++ with MPI and OpenMP support.

http://www.hpcg-benchmark.org/index.html

HPCG ranking(2019.6)(Latest)

| HPITE | Lict tor | June 2019 |
|-------|----------|-----------|
| | | |

| Rank | TOP500 Rank | System | Cores | Rmax (TFlop/s) | HPCG (TFlop/s) |
|------|----------------|--|------------|-------------------|-------------------|
| 1 | 1 | Summit - IBM Power System AC922, IBM POWER9 22C 3.076Hz, NVIDIA Volta GV100, Dual-rail Mellanox EDR Infiniband (/system/179397), IBM DOE/SC/Oak Ridge National Laboratory (/site/48553) United States | 2,414,592 | 148,600.0 | 2925.75 |
| 2 | 2 | Sierra - IBM Power System S922LC, IBM POWER9 22C 3.1GHz, NVIDIA Volta GV100, Dual-rail Mellanox EDR Infiniband (/system/179398), IBM / NVIDIA / Mellanox DDE/NNSA/LLNL (/site/49763) United States | 1,572,480 | 94,640.0 | 1795.67 |
| 3 | 20 | K computer, SPARC64 VIIIfx 2.0GHz, Tofu interconnect (/system/177232), Fujitsu RIKEN Advanced Institute for Computational Science (AICS) (/site/50313) Japan | 705,024 | 10,510.0 | 602.74 |
| 4 | 7 | Trinity - Cray XC40, Xeon E5-2698v3 16C 2.39Hz, Intel Xeon Phi 7250 68C 1.4GHz, Aries interconnect I/system/178610], Cray Inc. DOE/NNSA/LANL/SNL (/site/50334) United States | 979,072 | 20,158.7 | 546.12 |
| 5 | 8 | Al Bridging Cloud Infrastructure (ABCI) - PRIMERGY CX2570 MA, Xeon Gold 6148 20C 2.46Hz, NVIDIA Tesla V100 SXM2, Infiniband EDR I/system/179393], Fujitsu National Institute of Advanced Industrial Science and Technology (AIST) (/site/50762) | 391,680 | 19,880.0 | 508.85 |
| 6 | 6 | Piz Daint - Cray XC50, Xeon E5-2690v3 12C 2.6GHz, Aries interconnect , NVIDIA Tesla P100 (/system/177824), Cray Inc. Swiss National Supercomputing Centre (CSCS) (/site/50422) Switzerland | 387,872 | 21,230.0 | 496.98 |
| 7 | 3 | Sunway TaihuLight - Sunway MPP, Sunway SW26010 260C 1.450Hz, Sunway (/system/178764), NRCPC National Supercomputing Center in Wuxi (/site/50623) | 10,649,600 | 93,014.6 | 480.85 |
| 8 | 15 | Nurion - Cray CS500, Intel Xeon Phi 7250 68C 1.4GHz, Intel Omni-Path (/system/179421), Cray Inc. Korea Institute of Science and Technology Information (/site/50770) Korea South | 570,020 | 13,929.3 | 391.45 |
| 9 | 16 | Oakforest-PACS - PRIMERGY CX1640 M1, Intel Xeon Phi 7250 68C 1.4GHz, Intel Omni-Path (/system/178932), Fujitsu Joint Center for Advanced High Performance Computing (/site/50673) Japan | 556,104 | 13,554.6 | 385.48 |
| 10 | 14 | Cori - Cray XC40, Intel Xeon Phi 7250 68C 1.4GHz, Aries interconnect l/system/178924), Cray Inc. DOE/SC/LBNL/NERSC l/site/48429) United States | 622,336 | 14,014.7 | 355.44 |

USA

USA

Japan

USA

Japan

China

Japan

USA

Green500(省エネスパコンリスト)

• The purpose of the Green500 is to provide a ranking of the most energy-efficient supercomputers in the world. For decades, the notion of "performance" has been synonymous with "speed" (as measured in FLOPS). This particular focus has led to the emergence of supercomputers that consume egregious amounts of electrical power and produce so much heat that extravagant cooling facilities must be constructed to ensure proper operation. In addition, the emphasis on speed as the ultimate metric has caused other metrics such as reliability, availability, and usability to be largely ignored. As a result, there has been an extraordinary increase in the total cost of ownership (TCO) of a supercomputer.

http://www.green500.org/

Green 500(2019.6)(Latest)

| | Rank | TOP500 Rank | System | Cores | Rmax (TFlop/s) | | Power Efficiency (GFlops/watts) |
|-----|------|----------------|---|-----------|-------------------|------------------------|---------------------------------------|
| | 1 | 472 | Shoubu system B - ZettaScaler-2.2, Xeon D-1571 16C 1.3GHz, Infiniband EDR, PEZY-SC2 [/system/179165], PEZY Computing / Exascaler Inc. Advanced Center for Computing and Communication, RIKEN (/site/50577) Japan | 953,280 | 1,063.3 | ⁶⁰ Ja∶ | pan |
| GPU | 2 | 470 | DGX SaturnV Volta - NVIDIA DGX-1 Volta36, Xeon E5-2698v4 200-2 26Hz, Infiniband EDR, NVIDIA Tesla V100 (/system/1791osi, Nvidia NVIDIA Corporation (/site/48448) United States | 22,440 | 1,070.0 | 97 U | 15.113 S A |
| GPU | 3 | 1 | Summit row System AC922, IBM POWER9 22C 3. /GHz, NVIDIA VOIA 6 \ \ 100, Dual-rail Mellanox EDR Intoliband (/system/179/7), IBM DOE/90; 32 Bit - (Vational Laboratory (/site/48553) United States | 2,414,592 | 148,600.0 | 10,096 | 14.719 ISA |
| GPU | 4 | 8 | Al Bridging Cloud Infrastructure (AD69: - F Nm6SQY CX2570 M4, Xeon Gold 6148 200 2.46Hz, NVIDIA Testav100 SXM2, Infiniband EDR (Jsystem, Y9393), Fujitsu National Institute of Advanced Industrial Science and Technology (AIST) (Jsite/S0762) Japan | 391,680 | 19,880.0 | ^{1,649} Ja | ıpan |
| GPU | 5 | 394 | MareNostrum P9 CTE - IBM Power System AC922, IBM CONTENT 2023, IGHz, Dual-rail Mellanox EDR Infiniband, NVIDIA Tesla V1-0 [/system/179442], IBM Barcelona Supplicomputing Center [/site/49748] | 18,360 | 1,145.0 | 81 | ^{14.131} J S A |
| GPU | 6 | 25 | TSUBAME3.0 - SGI ICE YX IP139-SXM2, Xe n E5-2680v4 140 2.4GHz, Intel Omn(-Path, NVIDIA Tesla P III0 SXM2 (/system/179093), HPE GSIC Center, Tokyo Institute of Technology (/site/47912) Japan | 135,828 | 8,125.0 | ⁷⁹² Ja | 13.704 1 pan |
| GPU | 7 | 11 | PANGEA III - IBM Power System AC922 JoM POWER9 180 3.456Hz, Dual-rail Mellanox EDR Infini band, NVIDIA Volta GV100 [/system/179689], IBM Total Exploration Production [/site/49546] France | 291,024 | 17,860.0 | 1,367 | 13.065 |
| GPU | 8 | 2 | Sierra - IBM Power System S922L9, IBM SOWER9 22C 3.16Hz, NVIDIA Volta GV107, Dual-rail Mellanox DR Infiniband [/system/17939 l], IBM / NVIDIA / Mellanox DOE/NNSA/LLNL [/site/49783] United States | 1,572,480 | 94,640.0 | 7,438 | ^{12.723} J S A |
| | 9 | 43 | Advanced Computing System[PreE] - Sugon TC8600, Hygon Dhyana 32C 26Hz, Deep Computing Processor, 2006b 6D- Torus (/system/179593), Sugon Sugon (/site/S0797) China | 163,840 | 4,325.0 | 380 | 11.382 |
| GPU | 10 | 23 | Taiwania 2 - QCT QuantaGrid D546-4U/LC, Xeon Gold 6154 18C 3GHz, Mellanox InfiniBand EDR, NVIDIA Tesla V100 SXM2 (System/179590), Quanta Somputer / Taiwan Fixed Network / ASUS Cloud National Center for High Performance Computing (/site/48468) Taiwan | 170,352 | 9,000.0 | 798 | 11.285 |

This list fluctuates violently every time. That is why emphasis is placed on the development of this technology.

These machines balance energy saving and computing performance.

https://www.top5oo.org/green5oo/

Graph500 (graph analysis for big data)

Brief Introduction

Data intensive supercomputer applications are increasingly important for HPC workloads, but are illsuited for platforms designed for 3D physics simulations. Current benchmarks and performance metrics
do not provide useful information on the suitability of supercomputing systems for data intensive
applications. A new set of benchmarks is needed in order to guide the design of hardware architectures
and software systems intended to support such applications and to help procurements. Graph algorithms
are a core part of many analytics workloads.

Backed by a <u>steering committee</u> of over 50 international HPC experts from academia, industry, and national laboratories, Graph 500 will establish a set of large-scale benchmarks for these applications. The Graph 500 steering committee is in the process of developing comprehensive benchmarks to address three application kernels: concurrent search, optimization (single source shortest path), and edge-oriented (maximal independent set). Further, we are in the process of addressing five graph-related business areas: Cybersecurity, Medical Informatics, Data Enrichment, Social Networks, and Symbolic Networks.

This is the first serious approach to complement the Top 500 with data intensive applications. Additionally, we are working with the SPEC committee to include our benchmark in their CPU benchmark suite. We also maintain a list of the most energy efficient systems for data intensive computing. The Green Graph 500 will be released together with the Graph 500 list. We anticipate the list will rotate between ISC and SC in future years.

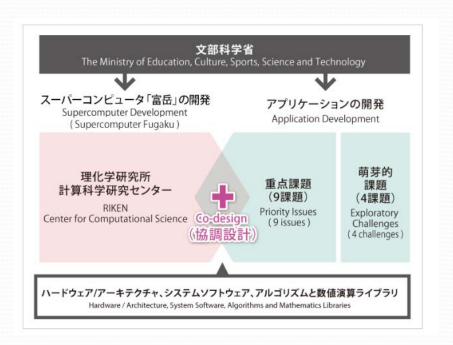
Graph500 (2019 June)(Latest)

Performance evaluation of "graph analysis" used for analysis of big data

| JUNE | 2019 BFS | | | | | | | | | | | | | | | | | | |
|------|------------------|---|---|---|---------------------|--|---------------------|----------------------|------|--|--|--------|----------|----------------------|-------------------------|------------|---------|----------------------|-----------------|
| Show | 100 ▼ entri | ies | Search: | | | | | | | | | NUMBER | NUMBER | | | | | | |
| RANK | PREVIOUS RANK | \$ MACHINE | VENDOR \$ | TYPE 4 | NETWORK ¢ | INSTALLATION SITE | \$ LOCATION \$ | COUNTRY | YEAR | APPLICATION | ‡ USAGE : | | OF ; | | IMPLEMENTATION | \$ SCALE\$ | GTEPS⊕ | CONSTRUCTION TIME | POWE |
| 1 | 1 | K computer | Fujitsu | Custom | Tofu | RIKEN Advanced Institute for Computational Science (AICS) | Kobe Hyogo | Japan | 2011 | Various scientific and instudria fields | al Academic and industry | 82944 | 663552 | 1327100 gigabytes | Custom | 40 | 31302.4 | 609.395 seconds | 0 Watts |
| 2 | 2 | Sunway TaihuLight | NRCPC | Sunway MPP | Sunway | National Supercomputing Center in Wuxi | Wuxi | China | 2015 | research | research | 40768 | 10599680 | 1304580 gigabytes | Custom | 40 | 23755.7 | 961.455 seconds | |
| 3 | 3 | DOE/NNSA/LLNL Sequoia | IBM | BlueGene/Q Power BQC 16C 1.60 GHz | Custom | Lawrence Livermore National Laboratory | Livermore CA | USA | 2012 | Scientific Research | Government | 98304 | 1572864 | 1572860 gigabytes | Custom | 41 | 23751 | 368.7 seconds | 638976 Watts |
| 4 | 4 | DOE/SC/Argonne National Laboratory Mira | IBM | BlueGene/Q Power BQC 16C 1.60 GHz | Custom | Argonne National Laboratory | Chicago IL | USA | 2012 | Scientific Research | Research | 49152 | 786432 | 786432 gigabytes | Custom | 40 | 14982 | 259.8 seconds | 0 Watts |
| 5 | 5 | SuperMUC-NG | Lenovo | ThinkSystem SD530 Xeon Platinum 8174 24C 3.1GHz Intel Omni-Path | | Leibniz Rechenzentrum | Garching | Germany | 2018 | Academic | Research | 4096 | 196608 | 393216 | custom-aml-heavy-diropt | 39 | 6279.47 | 1547.10 | |
| 6 | 6 | JUQUEEN | IBM | BlueGene/Q Power BQC 16C 1.60 GHz | Custom | Forschungszentrum Juelich (FZJ) | Juelich | Germany | 2012 | Scientific Research | Research | 16384 | 262144 | 262144 gigabytes | Custom | 38 | 5848 | 565 seconds | 108134 Watts |
| 7 | 8 | Fermi | IBM | BlueGene/Q Power BQC 16C 1.60 GHz | Custom | CINECA | Casalecchio Di Reno | Italy | 2012 | Scientific Research | Academic | 8192 | 131072 | 131072 gigabytes | Custom | 37 | 2567 | 400 seconds | 0 Watts |
| 8 | 9 | NERSC Cori - 1024 haswell partition | Cray | XC40 | Aries | NERSC/LBNL | DOE/SC/LBNL/NERSC | United States | 2017 | Government | basic science and simulation | 1024 | 32768 | 133376 | custom-aml-heavy-diropt | 37 | 2562.16 | 1369.29 | |
| 9 | 11 | Tianhe-2 (MilkyWay-2) | National University of Defense Technology | MPP | TH Express-2 | Changsha China | Changsha China | China | 2013 | Science research Big data | Research | 8192 | 196608 | 524288 gigabytes | custom | 36 | 2061.48 | 456.071 seconds | 0 Watts |
| 10 | new | Nurion | Cray | CS500 Intel Xeon Phi 7250 68C 1.4GHz Intel Omni-Path | Intel Omni- Path | Korea Institute of Science and Technology Information | Daejeon | Korea Republic Of | 2018 | Research | Various scientific and instudrial fields | 1024 | 65536 | 98304 | custom-aml-heavy-diropt | 37 | 1456.46 | 3519 | |

http://www.graph500.org/

post-K "Fugaku" project





Published specifications of post-K

ノード Nodes

| Armv8.2-A SVE 512bit 富士通鉱領:ハードウェアパリア、セクタキャッシュ、プリフェッチ | | | | |
|---|--|--|--|--|
| 48 + 2アシスタントコア 4 CMG (Core Memory Group, NUMA nodeのこと) | | | | |
| DP: 2.7+ TF, SP: 5.4+ TF, HP: 10.8 TF | | | | |
| L1D/core: 64 KiB, 4way, 230+ GB/s (load), 115+ GB/s (store) | | | | |
| L2/CMG: 8 MiB, 16way L2/node: 3.6+ TB/s L2/core: 115+ GB/s (load), 57+ GB/s (store) | | | | |
| HBM2 32 GiB, 1024 GB/s | | | | |
| Tofu Interconnect D (28 Gbps x 2 lane x 10 port) | | | | |
| PCIe Gen3 x16 | | | | |
| 7nm FinFET | | | | |
| | | | | |

プログラミング環境 Programming Environment

| Fortran2008 & Fortran2018サブセット C11 & GNU拡張仕様・Clang拡張仕様 C++14 & C++17サブセット & GNU拡張仕様・Clang拡張仕様 OpenMP 4.5 & OpenMP 5.0サブセット Java XcalableMP FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II 科学技術計算用ライブラリ SSL II (Fujitsu) | | | | | | | |
|---|---------------------------------------|--|--|--|--|--|--|
| コンパイラ C++14 & C++17サブセット & GNU拡張仕様・Clang拡張仕様 OpenMP 4.5 & OpenMP 5.0サブセット Java XcalableMP FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II | | Fortran2008 & Fortran2018サブセット | | | | | |
| OpenMP 4.5 & OpenMP 5.0サブセット Java XcalableMP FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II | | C11 & GNU拡張仕様・Clang拡張仕様 | | | | | |
| Java XcalableMP FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II | コンパイラ | C++14 & C++17サブセット & GNU拡張仕様・Clang拡張仕様 | | | | | |
| XcalableMP FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II | | OpenMP 4.5 & OpenMP 5.0サブセット | | | | | |
| 並列プログラミング FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II | | Java | | | | | |
| FPDPS スクリプト Python + Numpy + Scipy BLAS, LAPACK, ScaLAPACK, SSL II | ************************************* | XcalableMP | | | | | |
| BLAS, LAPACK, ScaLAPACK, SSL II | 型列ノログフミング | FPDPS | | | | | |
| | スクリプト | Python + Numpy + Scipy | | | | | |
| 科学技術計算用ライブラリ SSL II (Fujitsu) | | BLAS, LAPACK, ScaLAPACK, SSL II | | | | | |
| | 科学技術計算用ライブラリ | SSL II (Fujitsu) | | | | | |
| EigenEXA, Batched BLAS | | EigenEXA, Batched BLAS | | | | | |

システムソフトウェア System software

| os | Linux |
|---------|--|
| 03 | McKernel |
| MPI | Fujitsu MPI (Based on OpenMPI), MPICH |
| File IO | LLIO |
| The 10 | Application-oriented file IO libraries |

ストレージ Storages

_{第一階層} 1st level

- グローバルファイルシステムのキャッシュ
- テンポラリーファイルシステム
 - 計算ノードのローカルファイルシステム
 - ジョブの共有ファイルシステム

第二階層 2nd level

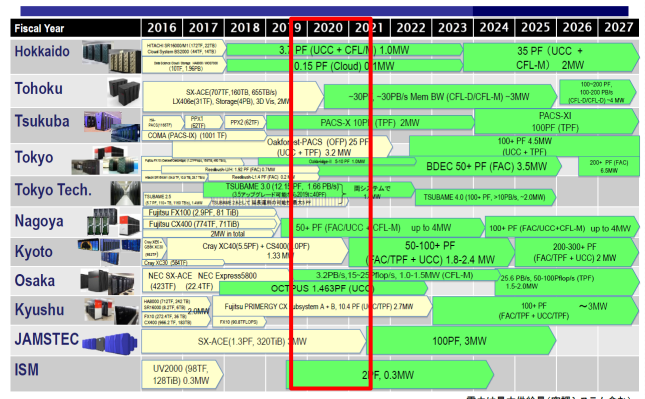
• Lustreベースのファイルシステム

3rd level

アーカイブ用ストレージ

HPCI supercomputer situation

HPCI第2階層システム 運用&整備計画 (2017年10月時点)



電力は最大供給量(空調システム含む)

- ✓ The highest use hope is the name FX 100. The highest use hope is the name FX 100. At present, it is scheduled to suspend operation in FY 2018, but extension for one year is being studied.
- ✓ Since there are supercomputers besides these (supercomputer of ISSP etc...), there is still much computing resources.

K computer is working on 24h/365days

- K computer was active for public use on 24hours/365 days for general users.
 - Different from K-computer, almost all the top class machines of the top500 ranking are for closed users and are used for limited purposes.
- The several rankings showed that K computer is almighty.
- Users may optimize their code for K computer easier than for the other ones(in the sense that it does not use a special architecture like GPU.).

Fugaku will be operated in the same way as K-computer.

Future direction of large-scale computing

From the above rankings

- From now on, both performance and energy-saving are important (Summit makes energy saving and computing performance highly compatible).
- The trend of many-core system with not so high frequency CPU is still going on. →Fugaku project is proceeding in this direction.
- Or many supercomputers have GPU and realize energy saving.
- As GPU is prominent in the field of AI and machine learning, this architecture will continue to be adopted in the future.
- However, I do not know whether supercomputers made up of GPU are effective in all fields or not.
- Probably, many supercomputers designed for data analysis and AI will appear after now.

However,...

- Effective performances measured by LINPACK or HPCG benchmarks are not necessarily accordance with the performance of our software codes. Usually, we spend so much effort to get better performance on supercomputers than typical PC clusters.
 - →The post-K machine, Fugaku, will be constructed for getting better performance for scientific application than benchmark programs (Co-design). (However, it may be the national mission to get the first prize of top 500 for taxpayers.)
- The effort goes beyond knowledge and techniques of physicists, chemist, biologists,... We need those of computer scientists and numerical mathematicians and need to collaborate them.
- Fugaku will be also a computer for general purpose. It may be inferior to a dedicated machine.(cf. Anton (supercomputer for performing classical MD) is 100~1000 faster than normal supercomputers.)

Future direction1

- The tendency of architecture is multi cores or many cores (including SIMD,GPU, ...).
- It is not easy to use all cores effectively.

(As personal opinion, I think that I do not have to worry about using all the cores and it is enough to get the best efficiency to obtain the calculation result.)

- Is it reasonable and proper to devote many researchers' effort to develop parallel efficiency? (very serious problem!!!)
- For DFT calculations, it may be difficult to speed-up the software by using GPU without breakthrough.
- We have to perform feasibility studies continuously to follow a leading-edge architecture.

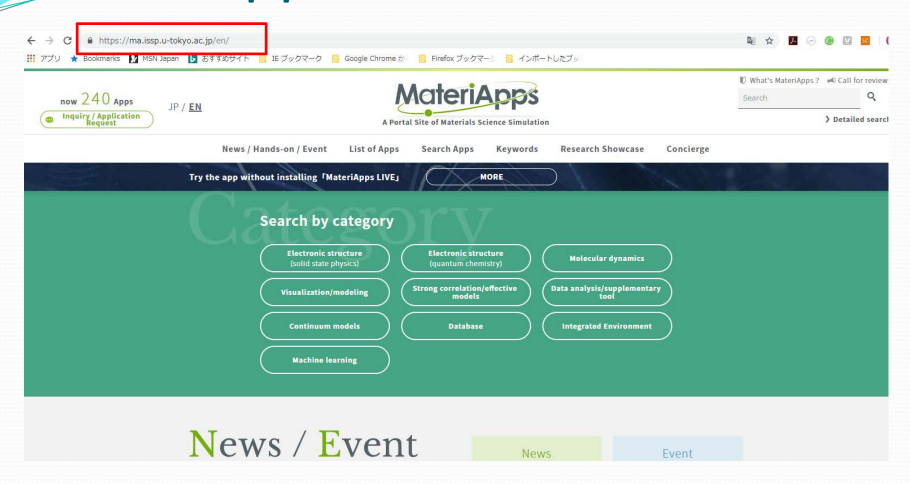
Future direction2

- The United States develops both architectures and programming language frameworks. We should contribute those.
- It is too much hard-pressed for physicists and chemists to cover computer science or numerical analysis.
 - →it is needed to establish a system to promote to work together with computational scientists and computer scientists or experts of numerical analysis.
- Still now, the guarantee of accuracy of calculation is inadequacy. There is no standard for summation of data from cores (threads) on MPI or OpenMP,... We must treat it carefully, otherwise our calculation may become meaningless.

Future direction3

- Today's amassed skills through K computer can be used during the next 10 years. (In Japan, Fugaku project, and the next project)
- However, the development of supercomputer must be stop in near future. (Prof. Hiraki said it was 2029. The reason is the necessity of huge electric power, failure of Moors' law, etc...)
- We should consider a new idea to get high performance of our own calculations for both software and hardware as soon as possible. We may undergo a paradigm shift before 2029.
- For the time being, data transfer will continue to be the most costly at all levels.

MateriApps



This portal site provides the information of applications that are used in materials science. You can find your desired application. "MateriApps LIVE" gives you an environment to perform open-source software codes without any installation process.

Finally,

- We should advance the development of human resources to be able to use supercomputers and/or HPC machines effectively.
- The relation between the ranking of top500 and the software code we use is not necessarily clear. We need to have the knowledge of the architecture of the supercomputers and/or the HPC machines if we'd like to maximize the performance of our software codes.
- We should cultivate people who can develop the software code.
- We should cultivate people who can effectively use scientific software codes to progress the researches. (CMD-WS, distance learning etc...)

Related texts



- These books are for developers.
- Integrates technologies for development on massively parallel computers such as supercomputers.
- Lecture videos and texts are available on the following site(Japanese only).

https://www.r-ccs.riken.jp/library/event/tokuronA_2019.html https://www.r-ccs.riken.jp/library/event/tokuronB_180406.html