2L is a software package that solves a random, dense linear system of equations with N unknowns, in double-precision arithmetic on distributed-memory computers. It then prints the estimated performance of the stributed-memory computer as the ratio between the number of effective floating point operations divided by the amount of time in seconds it took to perform these operations.

How do I run the HPL test on a Bright Cluster?

What is HPL?

Slightly rewording from http://www.netlib.org/benchmark/hpl/:

It can thus be regarded as a portable as well as freely available implementation of the High Performance Computing LINPACK Benchmark.

The algorithm

- Two-dimensional block-cyclic data distribution
- Right-looking variant of the LU factorization with row partial pivoting featuring multiple look-ahead depths
- Recursive panel factorization with pivot search and column broadcast combined
- Various virtual panel broadcast topologies
  - bandwidth reducing swap-broadcast algorithm
  - backward substitution with look-ahead of depth 1

#### The HPL algorithm is changeable and can be tuned via 17 parameters:

N	Problem size	Pmap	Process mapping
NB	Blocking factor	threshold	for matrix validity test
2	Rows in process grid	Ndiv	Panels in recursion
Q	Columns in process grid	Nbmin	Recursion stopping criteria
Depth	Lookahead depth	[Swap]	Swap algorithm
Bcasts	Panel broadcasting method	L1, U	to store triangle of panel
Pfacts	Panel factorization method	Align	Memory alignment
Rfacts	Recursive factorization method	Equilibration	

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The parameters in the red cells are the most important tuning parameters, while the ones in the green cells are of somewhat lesser importance. The parameter space is huge and exploring the full parameter space is inefficient as there are more than a billion different combinations. If one considers only combination of the most important parameters the parameter space reduces to 50.000 combinations.

#### Third Party Software: How do I run the HPL test on a Bright **Cluster?** for tuning, but they are beyond the scope of this article. However, it is worth mentioning that one does not have to use the maximal problem to that tuning algorithms perform well and seem to be robust suggest that for a given architecture the amount of important parameters might

ns that doubling N requires eight times more computations. Values of the unstat command. A useful formula to estimate the problem size

n a load balancing point of view, but too small values might limit performance. A value in the

in tuning. Early te

The tuning parameters need to be stored in a file named HPL.dat.

Setting the values of HPL.dat parameters

N = sqrt (0.75 \* Number of Nodes \* Minimum memory of any node / 8)

The problem size N should be set to a large value but not too large. The amount of computations scales as the cube of that number wi order of 1000-10.000 are reasonable for modern systems. Make sure that there is no paging occurring while executing the problem by

IPL uses the block size NB for the data distribution as well as for the computational granularity. Small value ange [32,...,256] is good for modern systems with large values being more efficient for larger problem sizes.

nce the panel factorization has been

Algorithm 2: Increasing-ring (modified) is the usually the best choice

There are also a couple of web-based HPL.dat generators like the following:

• Algorithm 4: Increasing-2-ring (modified) is usually a second choice.

· Algorithm 5: Long (bandwidth reducing) - this can be a good choice for systems with very fast nodes, but a very slow network.

The Rfact and Pfact values need experimentation, but for very large clusters both parameters can be set to the right-looking part of the LU decomposition.

A final recommendation is to ensure that HPL is not going to use more than 80% of the memory available to that system

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http://www.advancedclustering.com/fag/how-do-i-tune-my-hpldat-file.html

A sample HPL.dat file

HPLinpack benchmark input file Innovative Computing Laboratory, University of Tennessee			
HPL.out	output file name (if any)		
6	device out (6=stdout,7=stderr,file)		
1	# of problems sizes (N)		
58464	Ns		
1	# of NBs		
168	NBs		
0	PMAP process mapping (0=Row-,1=Column-major)		
1	# of process grids (P x Q)		
4	Ps		
8	Qs		
16.0	threshold		
1	# of panel fact		
2	PFACTs (0=left, 1=Crout, 2=Right)		
1	# of recursive stopping criterium		
4	NBMINs (>= 1)		
1	# of panels in recursion		
2	NDIVs		
1	# of recursive panel fact.		
1	RFACTs (0=left, 1=Crout, 2=Right)		
1	# of broadcast		
1	BCASTs (0=1rg,1=1rM,2=2rg,3=2rM,4=Lng,5=LnM)		
1	# of lookahead depth		
1	DEPTHs (>=0)		
2	SWAP (0=bin-exch,1=long,2=mix)		
64	swapping threshold		
0	L1 in (0=transposed,1=no-transposed) form		
0	U in (0=transposed,1=no-transposed) form		
1	Equilibration (0=no,1=yes)		

8 memory alignment in double (> 0)

HPL requires a distributed-memory system with a tuned implementation of the BLAS libr

Prerequisites

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on of the MPI standard. The *Bright Cluster Manager* provides several modules of the OpenBLAS atect the relevant architecture and use routines optimized for that particular system.

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The relevant modules can be loaded with the command below:

[user@myHeadNode ~]\$ module load shared openmpi/gcc openblas/dynamic hpl

The other available tuned versions of OpenBLAS are:

- openblas/bulldozer

- openblas/istanbul

- openblas/nehalem

- openblas/sandybridge

Note: OpenBLAS is available in Bright 6.1. Users of Bright 5.2 and 6.0 can choose between gotoBLAS and BLAS. In the latter case the HPL binary is called xhpl.blas.

Running HPL on a Bright cluster

To run HPL after loading the modules, the user needs to issue the command:

mpiexec -machinefile nodes \_ xhpl | tee HPL.out

If your cluster has InfiniBand, and you want to see the p

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InfiniBand and Ethernet, you can do a run over Ethernet by issuing the following command:

Sample output:

-----

HPLinpack 2.1 -- High-Performance Linpack benchmark -- October 26, 2012

Written by A. Petitet and R. Clint Whaley, Innovative Computing Laboratory, UTK

Modified by Piotr Luszczek, Innovative Computing Laboratory, UTK

Modified by Julien Langou, University of Colorado Denver

-----

An explanation of the input/output parameters follows:

T/V : Wall time / encoded variant.

N : The order of the coefficient matrix A.

NB : The partitioning blocking factor.

P : The number of process rows.

Q : The number of process columns.

Time : Time in seconds to solve the linear system.

Gflops : Rate of execution for solving the linear system.

The following parameter values will be used:

N : 79232

NB : 128

PMAP : Row-major process mapping

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p : 4
Q ÷ 4
PFACT : Right
NBMIN : 4
NDIV : 2
RFACT : Crout
BCAST : lringM
DEPTH : 1
SWAP : Mix (threshold = 64)
Ll : transposed form
U : transposed form
EQUIL : yes
ALIGN : 8 double precision words
- The matrix A is randomly generated for each test.
- The following scaled residual check will be computed:
Ax-b  _oo / ( eps * (    x   _oo *    A   _oo +    b   _oo ) * N )
- The relative machine precision (eps) is taken to be 1.110223e-16
- Computational tests pass if scaled residuals are less than 16.0
T/V N NB P Q Time Gflops

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HPL\_pdgesv() start time Wed Jan 2 21:53:09 2013

HPL\_pdgesv() end time Wed Jan 2 22:31:40 2013

||Ax-b||\_00/(eps\*(||A||\_00\*||x||\_00+||b||\_00)\*N)= 0.0034949 ...... PASSED

-----

Finished 1 tests with the following results:

1 tests completed and passed residual checks,

0 tests completed and failed residual checks,

0 tests skipped because of illegal input values.

End of Tests.

Unique solution ID: #1124 Author: Panos Labropoulos Last update: 2015-07-23 09:56

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