



REFERENCE ARCHITECTURES OF DELL EMC READY BUNDLE FOR HPC LIFE SCIENCES

Refresh with 14th Generation servers

ABSTRACT

Dell EMC's flexible HPC architecture for Life Sciences has been through a dramatic improvement with new Intel® Xeon® Scalable Processors. Dell EMC Ready Bundle for HPC Life Sciences equipped with better 14G servers, faster CPUs, and more memory bring a much higher performance in terms of throughput compared to the previous generation especially in genomic data processing.

March 2018

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EXECUTIVE SUMMARY

Since Dell EMC announced Dell EMC HPC solution for Life Science in September 2016, the current Dell EMC Ready Bundle for HPC Life Sciences can process 485 genomes per dayⁱ with 64x C6420s and Dell EMC Isilon F800 in our benchmarking. This is roughly a two-fold improvement from Dell EMC HPC System for Life Science v.1.1 due to the introduction of Dell EMC's 14th generation servers which include the latest Intel® Xeon® Scalable processors (code name: Skylake), updated server portfolio, improved memory, and storage subsystem performance (1).

This whitepaper describes the architectural changes and updates to the follow-on of Dell EMC HPC System for Life Science v1.1. It explains new features, demonstrates the benefits, and shows the improved performance.

AUDIENCE

This document is intended for organizations interested in accelerating genomic research with advanced computing and data management solutions. System administrators, solution architects, and others within those organizations constitute the target audience.

INTRODUCTION

Although the successful completion of the Human Genome Project was announced on April 14, 2003 after a 13-year-long endeavor and numerous exciting breakthroughs in technology and medicine, there's still a lot of work ahead for understanding and using the human genome. As an iterative advance in sequencing technology has accumulated, these cutting-edge technologies allow us to look at many different perspectives or levels of genetic organization such as whole genome sequencing, copy number variations, chromosomal structural variations, chromosomal methylation, global gene expression profiling, differentially expressed genes, and so on. However, despite the scores of data we generated, we still face the challenge of understanding the basic biology behind the human genome and the mechanisms of human diseases. There will not be a better time than now to evaluate if we have overlooked the current approach, "analyze large numbers of diverse samples with the highest resolution possible" because analyzing a large number of sequencing data alone has been unsuccessful to identify key genes/variants in many common diseases where majority of genetic variations seem to be involved randomly. This is the main reason that data integration becomes more important than before. We have no doubt that genome research could help fight human disease; however, combining proteomics and clinical data with genomics data will increase the chances of winning the fight. As the life science community is gradually moving toward to data integration, **Extract, Transform, Load** (ETL) process will be a new burden to an IT department (2).

Dell EMC Ready Bundle for HPC Life Sciences has been evolving to cover various needs for Life Science data analysis from a system only for **Next Generation Sequencing** (NGS) data processing to a system that can be used for Molecular Dynamics Simulations (MDS), Cryo-EM data analysis and *De Novo* assembly in addition to DNA sequencing data processing. Further, we are planning to cover more applications from other areas of Life Sciences and re-design the system suitable for data integration. In this project, we tested an additional two Dell EMC Isilon storages as the importance of finding suitable storages for variable purposes grows. This is an additional step to build a HPC system as a tool to integrate all the cellular data, biochemistry, genomics, proteomics and biophysics into a single frame of work and to be ready for the high-demanding era of ETL process.

SOLUTION OVERVIEW

Especially, HPC in Life Sciences requires a flexible architecture to accommodate various system requirements. Dell EMC Ready Bundle for HPC Life Sciences was created to meet this need. It is a pre-integrated, tested, tuned, and leverages the most relevant of Dell EMC's high-performance computing line of products and best-in-class partner products (3). It encompasses all of the hardware resources required for various life sciences data analysis, while providing an optimal balance of compute density, energy efficiency and performance.

ARCHITECTURE

Dell EMC Ready Bundle for HPC Life Sciences provides high flexibility. The platform is available in three variants which are determined by the cluster interconnect selected for the storages. In the current version, the following options are available:

- Dell EMC PowerEdge™ C6420 compute subsystem with Intel® Omni-Path (OPA) fabric or Mellanox InfiniBand® (IB) EDR fabric
 - o Storage choice:
 - Dell EMC Ready Bundle for HPC Lustre Storage as a performance scratch space
 - Dell EMC Ready Bundle for HPC NFS Storage as a home directory space
- PowerEdge C6420 compute subsystem with 10/40GbE fabric
 - o Storage choice:
 - Either Dell EMC Isilon F800 or Dell EMC Isilon H600 as a performance scratch space
 - Dell EMC Ready Bundle for HPC NFS Storage as a home directory space
- Add-on compute nodes:
 - o Dell EMC PowerEdge R940
 - This server covers large memory applications such as *De Novo* Assembly
 - o Dell EMC PowerEdge C4130
 - A server for accelerators like NVIDIA GPUs for Molecular Dynamics Simulations

In addition to the compute, network, and storage options, there are several other components that perform different functions in the Dell EMC Ready Bundle for HPC Life Sciences. These include CIFS gateway, fat node, acceleration node and other management components. Each of these components is described in detail in the subsequent section.

The solutions are nearly identical for Intel® OPA and IB EDR versions except for a few changes in the switching infrastructure and network adapter. The solution ships in a deep and wide 48U rack enclosure, which helps to make PDU mounting and cable management easier. Figure 1 shows the components of two fully loaded racks using 64x Dell EMC PowerEdge C6420 rack server chassis as a compute subsystem, Dell EMC PowerEdge R940 as a fat node, Dell EMC PowerEdge C4130 as an accelerator node, Dell EMC Ready Bundle for HPC NFS Storage, Dell EMC Ready Bundle for HPC Lustre Storage and Intel® OPA as the cluster's high-speed interconnect.

Compute and Management Components

There are several considerations when selecting the servers for master node, login node, compute node, fat node and accelerator node. For master, 1U form factor Dell EMC PowerEdge R440 is recommended. The master node is responsible for managing the compute nodes and optimizing the overall compute capacity. The login node (Dell EMC PowerEdge R640 is recommended) is used for user access, compilations and job submissions. Usually, master and login nodes are the only nodes that communicate with the outside world, and they act as a middle point between the actual cluster and the outside network. For this reason, high availability can be provided for master and login nodes. An example solution is illustrated in Figure 1, and the high-speed interconnect is configured to 2:1 blocking fat tree topology with Intel® OPA fabric.

Ideally, the compute nodes in a cluster should be as identical as possible since the performance of parallel computation is bounded by the slowest component in the cluster. Heterogeneous clusters do work, but it requires careful execution to achieve the best performance; however, for Life Sciences applications, heterogeneous clusters make perfect sense to handle completely independent workloads such as DNA-Seq, *De Novo* assembly or Molecular Dynamics Simulations. These workloads require quite different hardware components. Hence, we recommend Dell EMC PowerEdge C6420 as a compute node to handle NGS data processing due to its density, a wide choice of CPUs, and high maximum memory capacity. Dell EMC PowerEdge R940 is an optional node with 6TB of RDIMM /LRDIMM memory and is recommended for customers who need to run applications requiring large memory such as *De Novo* assembly. Accelerators are used to speed up computationally intensive applications, such as molecular dynamics simulation applications. We tested configurations G and K for this solution.

The compute and management infrastructure consists of the following components.

- Compute
 - o Dell EMC PowerEdge C6400 enclosure with 4x C6420 servers
 - CPU: 2x Intel® Xeon® Gold 6148 20c 2.4GHz (Skylake)
 - Memory: 12 x 32GB @2666 MHz
 - Disk: 6x 480GB 12Gbps SAS Mixed Used SSDs
 - Interconnect: Intel® Omni-Path, IB EDR or/and 10/40GbE
 - BIOS System Profile: Performance Optimized
 - Logical Processor: Disabledⁱⁱ
 - Virtualization Technology: Disabled
 - OS: RHEL 7.4
 - o Dell EMC PowerEdge C4130 with up to four NVIDIA® Tesla™ P100 or V100 GPUs (K configuration recommended, Figure 9)
 - CPU: 2x Intel® Xeon® E5-2690 v4 14c 2.6GHz (Broadwell)
 - Memory: 16x 16GB @2400 MHz
 - Disk: 9TB HDD
 - BIOS System Profile: Performance Optimized
 - Logical Processor: Disabled
 - Virtualization Technology: Disabled
 - OS: RHEL 7.4
 - o Dell EMC PowerEdge R940
 - CPU: 4x Intel® Xeon® Platinum 8168, 24c 2.7GHz (Skylake)
 - Memory: 48x 32GB @2666 MHz
 - Disk: 12x 1.2TB 10K RPM SAS 12Gbps 512n 2.5in Hot-plug Hard Drive in RAID 0
 - BIOS System Profile: Performance Optimized
 - Logical Processor: Disabled
 - Virtualization Technology: Disabled
 - OS: RHEL 7.4

- Management
 - o Dell EMC PowerEdge R440 for master node
 - CPU: Intel® Xeon® Gold 5118, 12c 2.3GHz (Skylake)
 - Memory: 12x 8GB @2666 MHz
 - Disk: 10x 1.8TB 10K RPM SAS 12Gbps 512e 2.5in Hot-plug Hard Drive in RAID 10
 - BIOS System Profile: Performance Optimized
 - Logical Processor: Enabled
 - Virtualization Technology: Disabled
 - OS: RHEL 7.4
 - o Dell EMC PowerEdge R640 for login node and CIFS gateway (optional)
 - CPU: Intel Xeon Gold 6132, 14c 2.6GHz (Skylake)
 - Memory: 12x 16GB @2666 MHz
 - Disk: 4x 1.2TB 10K RPM SAS 12Gbps 512n 2.5in Hot-plug Hard Drive
 - BIOS System Profile: Performance Optimized
 - Logical Processor: Enabled
 - Virtualization Technology: Disabled
 - OS: RHEL 7.4

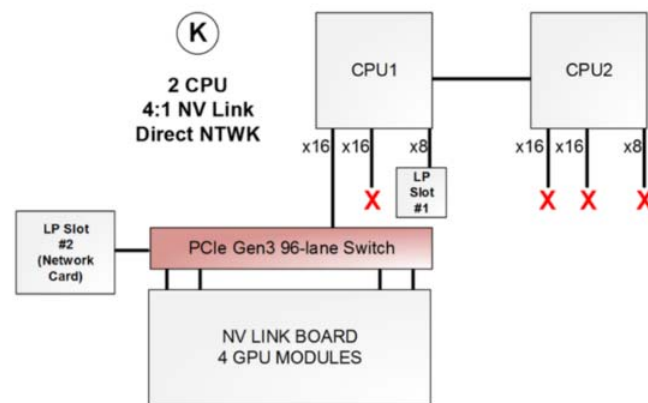


Figure 1 K configuration

Storage Components

The storage infrastructure consists of the following components and the details of the configurations can be found from the references:

- Dell EMC Ready Bundle for HPC NFS Storage (NSS7.0-HA) (4)
- Dell EMC Ready Bundle for HPC Lustre Storage (4)
- Dell EMC Isilon Scale-out NAS Product Family; Dell EMC Isilon F800 All-flash (5)
or Dell EMC Isilon Hybrid Scale-out NAS H600 (5)

Table 1 Tested Dell EMC Isilon Configurations

	Dell EMC Isilon F800	Dell EMC Isilon H600
NUMBER OF NODES	4	4
CPU PER NODE	Intel® Xeon™ CPU E5-2697A v4 @2.60 GHz	Intel® Xeon™ CPU E5-2680 v4 @2.40GHz
MEMORY PER NODE	256GB	256GB
STORAGE CAPACITY	Total usable space: 166.8 TB, 41.7 TB per node	Total usable space: 126.8 TB, 31.7 TB per node
SSD L3 CACHE	N/A	2.9 TB per node
NETWORK	Front end network: 40GbE Back end network: 40GbE	Front end network: 40GbE Back end network: IB QDR
OS	Isilon OneFS v8.1.0 DEV.0	Isilon OneFS v8.1.0.0 B_8_1_0_011

Network Components

Dell Networking H1048-OPF

Intel® Omni-Path Architecture (OPA) is an evolution of the Intel® True Scale Fabric Cray Aries interconnect and internal Intel® IP [9]. In contrast to Intel® True Scale Fabric edge switches that support 36 ports of InfiniBand QDR-40Gbps performance, the new Intel® Omni-Path fabric edge switches support 48 ports of 100Gbps performance. The switching latency for True Scale edge switches is 165ns-175ns. The switching latency for the 48-port Omni-Path edge switch has been reduced to around 100ns-110ns. The Omni-Path host fabric interface (HFI) MPI messaging rate is expected to be around 160 million messages per second (MMPS) with a link bandwidth of 100Gbps.



Figure 2 Dell Networking H1048-OPF

Mellanox SB7700

This 36-port Non-blocking Managed InfiniBand EDR 100Gb/s Switch System provides the highest performing fabric solution in a 1U form factor by delivering up to 7.2Tb/s of non-blocking bandwidth with 90ns port-to-port latency.



Figure 3 Mellanox SB7700

Dell Networking Z9100-ON

The Dell EMC Networking Z9100-ON is a 10/25/40/50/100GbE fixed switch purpose-built for applications in high-performance data center and computing environments. 1RU high-density 10/25/40/50/100GbE fixed switch with choice of up to 32 ports of 100GbE (QSFP28), 64 ports of 50GbE (QSFP+), 32 ports of 40GbE (QSFP+), 128 ports of 25GbE (QSFP+) or 128+2 ports of 10GbE (using breakout cable).



Figure 4 Dell Networking Z9100-ON

Dell Networking S3048-ON

Management traffic typically communicates with the Baseboard Management Controller (BMC) on the compute nodes using IPMI. The management network is used to push images or packages to the compute nodes from the master nodes and for reporting data from client to the master node. Dell EMC Networking S3048-ON is recommended for management network.



Figure 5 Dell Networking S3048-ON

Reference Architecture Case 1

This reference architecture is configured with Intel® OPA fabric, Dell EMC Ready Bundle for HPC NFS Storage and Dell EMC Ready Bundle for HPC Lustre Storage. Dell EMC Ready Bundle for HPC NFS Storage, referred to as Dell NFS Storage Solution-High Availability configuration (NSS7.0-HA) is configured for general usage such as a home directory while Dell EMC Ready Bundle for HPC Lustre Storage serves as a high-performance scratch storage.

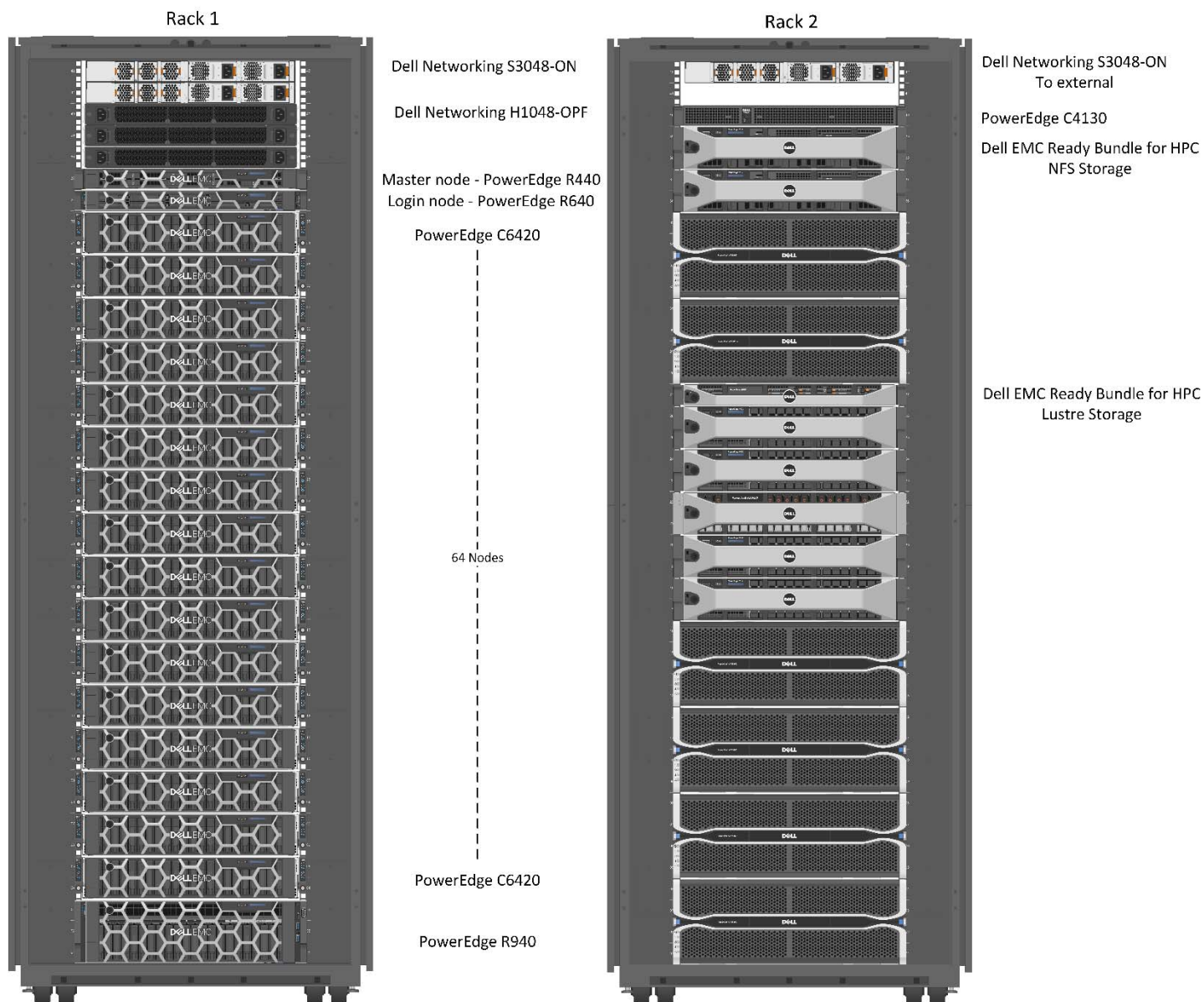


Figure 6 Dell EMC Ready Bundle for HPC Life Sciences with Intel® OPA fabric

Figure 2 illustrates the details of management network. It requires three Dell Networking S3048-ON switches. Two switches in Rack 1 (Figure 1) make up the internal management network for deploying, provisioning, managing, and monitoring the cluster. One of the switches in Rack 2 has 48 ports that are split into multiple untagged virtual LANs to accommodate multiple internal networks.

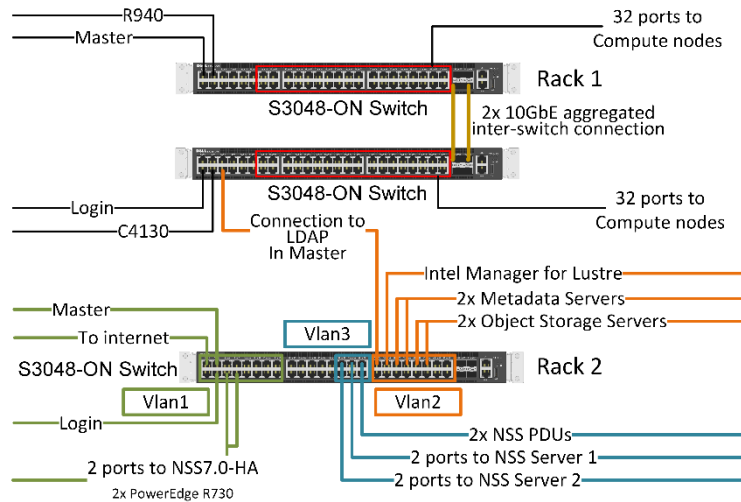


Figure 7 Management Network

Figure 3 shows high-speed interconnect with Intel® OPA. The network topology is 2:1 blocking fat tree which requires three Dell Network H1048-OPF 48 port switches.

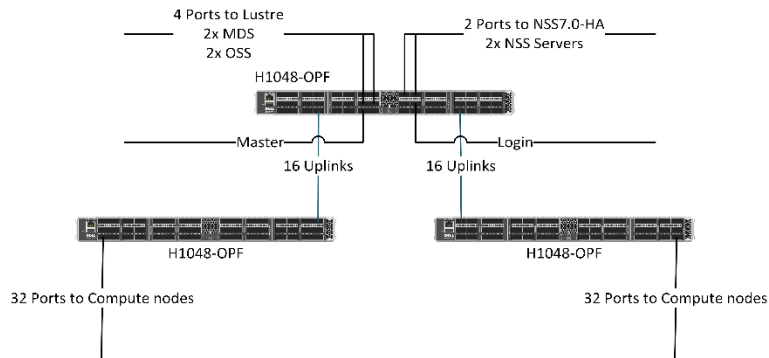


Figure 8 Interconnection with Intel® OPA; 2:1 blocking fat tree topology

Reference Architecture Case 2

Dell EMC Isilon storage, F800 All-flash is configured into Dell EMC Ready Bundle for HPC Life Sciences. In this reference architecture, the interconnect for the F800 has 8 ports of 40GbE connections. Since two 40GbE external connections in each node of the F800 can be aggregated, the Z9100-ON switch is configured to support two-port aggregation on each node. Hence, as shown in Figure 5, two 40GbE ports are aggregated to one port channel which connects to a node in the F800.

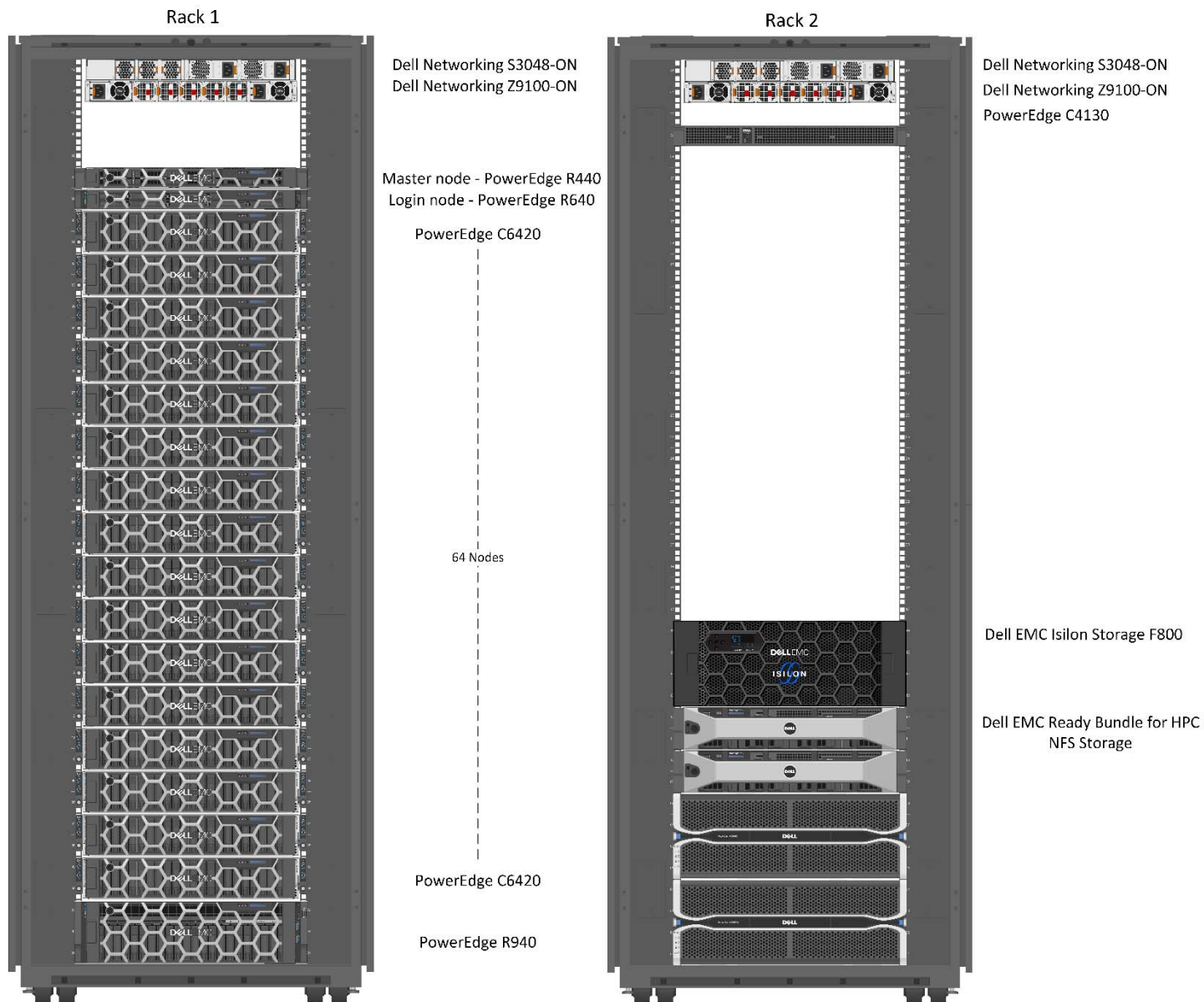


Figure 9 Dell EMC Ready Bundle for HPC Life Sciences with 10GbE fabric and F800

Eight ports on each Z9100-ON switch in both racks are used to connect between the switches. All eight ports are aggregated to a single port-channel.

The Z9100-ON switches are configured with multiple untagged virtual LANs to accommodate the backend network of the F800. Vlan2s in the both switches are private to the backend networks of the F800. A total of eight 40GbE ports are assigned to this private network.

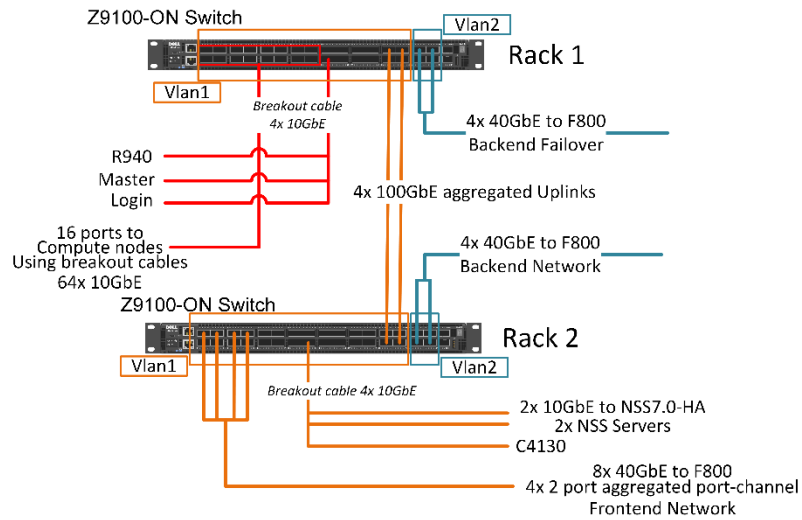


Figure 10 Interconnection with 10GbE (1:1 Non-blocking)

The management network is identical to case 1 except for the connections to the Lustre storage (Figure 6).

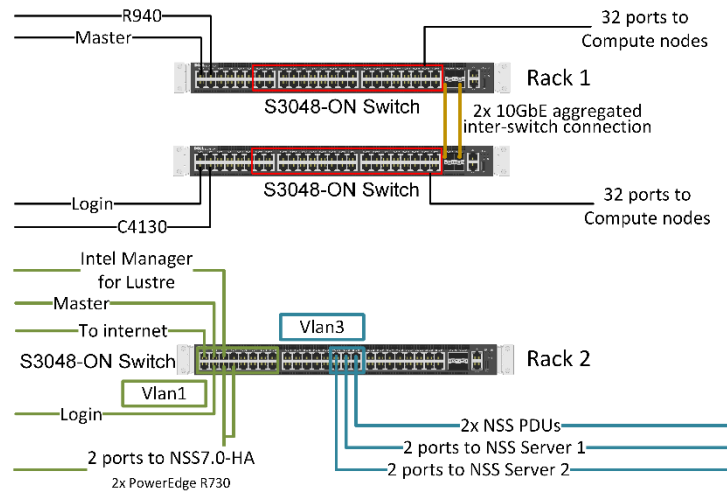


Figure 11 Management Network

Reference Architecture Case 3

This example shows Dell EMC Ready Bundle for HPC Life Sciences with Dell EMC Isilon Storage H600. The H600 is one of Dell EMC Isilon Hybrid Scale-out NAS storages. The details of the storage description can be found in the next sections.

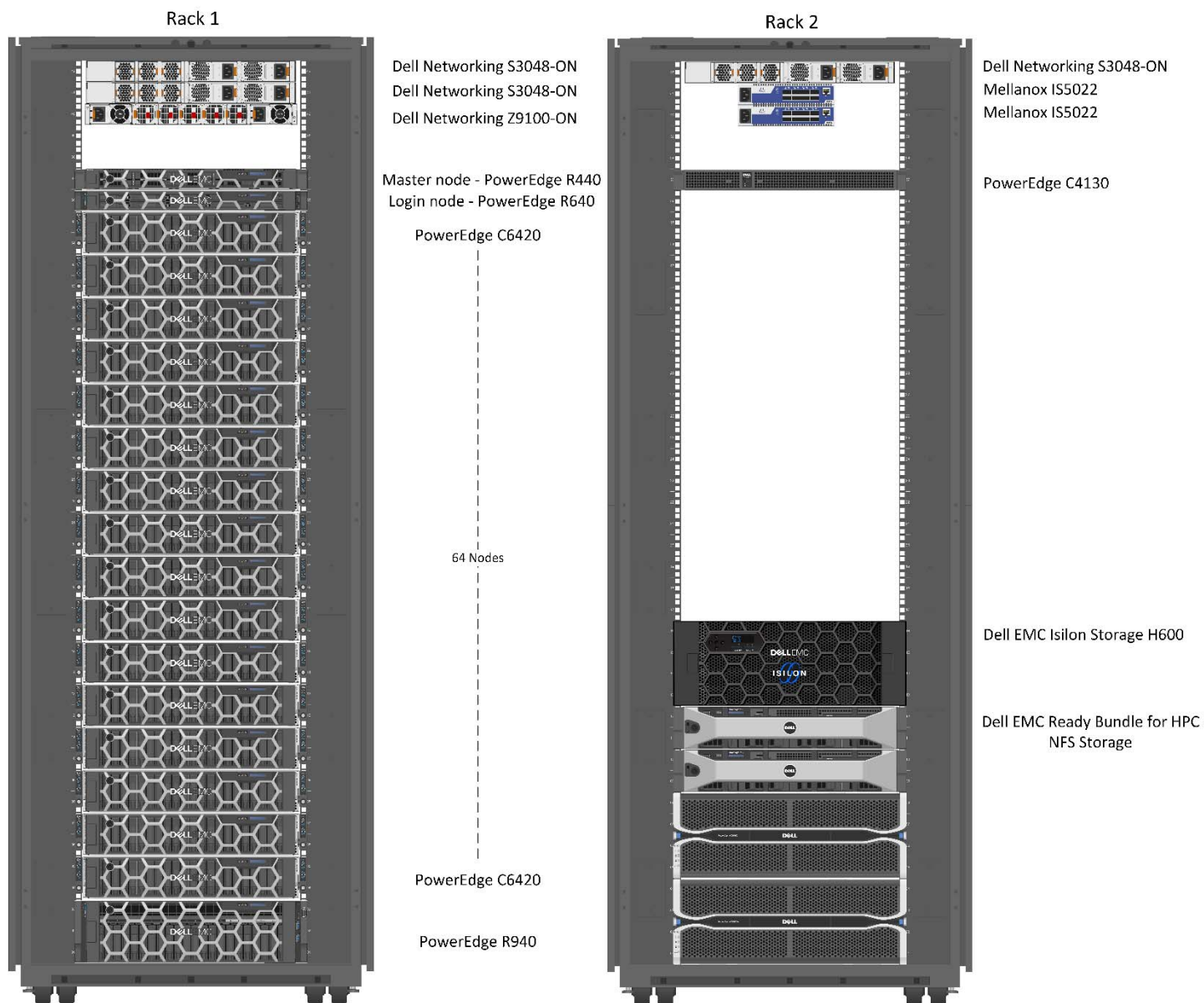


Figure 12 Dell EMC Ready Bundle for HPC Life Sciences with 10GbE fabric and H600

The management network for this reference architecture is identical to the one shown in Figure 6. Also, 10/40GbE and IB QDR interconnect networks are illustrated in Figure 8. The H600 uses IB QDR as the backend network. Two 8 ports IS 5022 switches are used to connect interconnect four storage nodes in the H600. Like the F800's frontend network, the H600 supports 2x 40GbE aggregate external link per node.

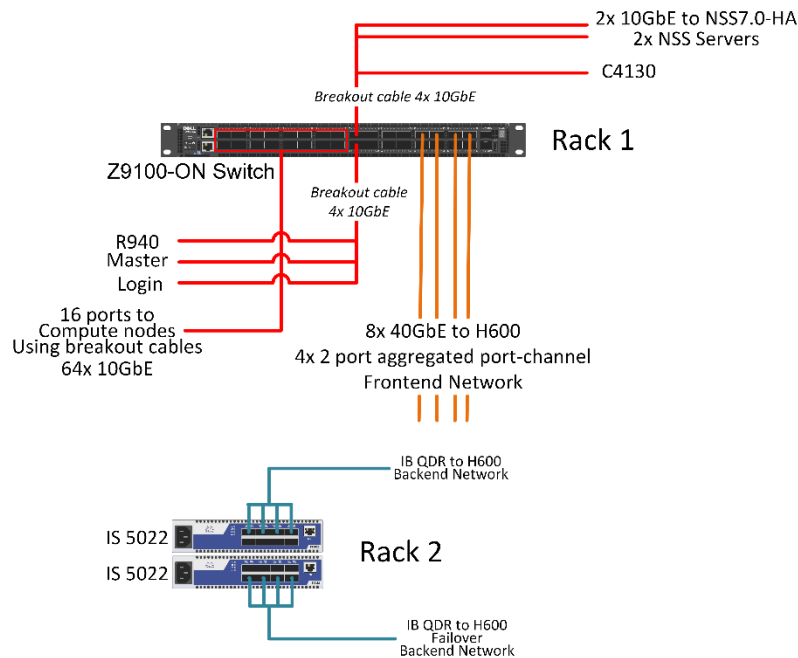


Figure 13 Interconnection with 10GbE (1:1 Non-blocking) and IB QDR

Software Components

Along with the hardware components, the solution includes the following software components:

- Bright Cluster Manager®
- BioBuilds

Bright Cluster Manager

Bright Computing is a commercial software that provides comprehensive software solutions for deploying and managing HPC clusters, big data clusters and OpenStack in the data center and in the cloud (6). Bright cluster Manager can be used to deploy complete clusters over bare metal and manage them effectively. Once the cluster is up and running, the graphical user interface monitors every single node and reports if it detects any software or hardware events.

BioBuilds

BioBuilds is a well maintained, versioned, and continuously growing collection of open-source bio-informatics tools from Lab7 (7). They are prebuilt and optimized for a variety of platforms and environments. BioBuilds solves most software challenges faced by the life sciences domain.

- Imagine a newer version of a tool being released. Updating it may not be straight forward and would probably involve updating all of the dependencies the software has as well. BioBuilds includes the software and its supporting dependencies for ease of deployment.
- Using BioBuilds among all of the collaborators can ensure reproducibility since everyone is running the same version of the software. In short, it is a turnkey application package.

PERFORMANCE EVALUATION AND ANALYSIS

GENOMICS/NGS DATA ANALYSIS PERFORMANCE

A typical variant calling pipeline consists of three major steps 1) aligning sequence reads to a reference genome sequence; 2) identifying regions containing SNPs/InDels; and 3) performing preliminary downstream analysis. In the tested pipeline, BWA 0.7.2-r1039 is used for the alignment step and Genome Analysis Tool Kit (GATK) is selected for the variant calling step. These are considered standard tools for aligning and variant calling in whole genome or exome sequencing data analysis. The version of GATK for the tests is 3.6, and the actual workflow tested was obtained from the workshop, 'GATK Best Practices and Beyond'. In this workshop, they introduce a new workflow with three phases.

- Best Practices Phase 1: Pre-processing
- Best Practices Phase 2A: Calling germline variants
- Best Practices Phase 2B: Calling somatic variants
- Best Practices Phase 3: Preliminary analyses

Here we tested phase 1, phase 2A and phase 3 for a germline variant calling pipeline. The details of commands used in the benchmark are in APPENDIX A. GRCh37 (Genome Reference Consortium Human build 37) was used as a reference genome sequence, and 30x whole human genome sequencing data from the Illumina platinum genomes project, named ERR091571_1.fastq.gz and ERR091571_2.fastq.gz were used for a baseline test (11).

It is ideal to use non-identical sequence data for each run. However, it is extremely difficult to collect non-identical sequence data having more than 30x depth of coverage from the public domain. Hence, we used a single sequence data set for multiple simultaneous runs. A clear drawback of this practice is that the running time of Phase 2, Step 2 might not reflect the true running time as researchers tend to analyze multiple samples together. Also, this step is known to be less scalable. The running time of this step increases as the number of samples increases. A subtle pitfall is a storage cache effect. Since all of the simultaneous runs will read/write roughly at the same time, the run time would be shorter than real cases. Despite these built-in inaccuracies, this variant analysis performance test can provide valuable insights to estimating how much resources are required for an identical or even similar analysis pipeline with a defined workload.

The throughput of Dell HPC Solution for Life Sciences

Total run time is the elapsed wall time from the earliest start of Phase 1, Step 1 to the latest completion of Phase 3, Step 2. Time measurement for each step is from the latest completion time of the previous step to the latest completion time of the current step as illustrated in Figure 16.

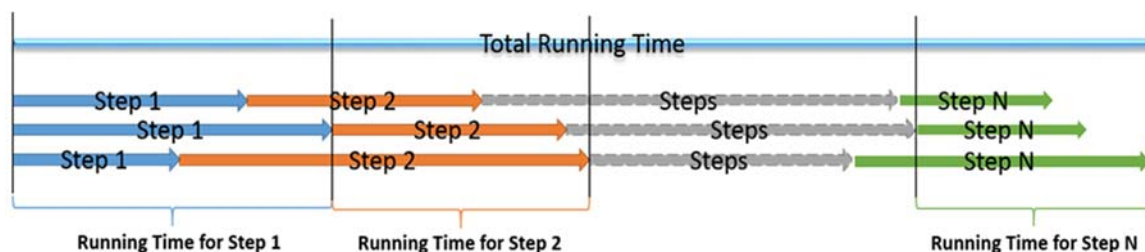


Figure 14 Running time measurement method

Feeding multiple samples into an analytical pipeline is the simplest way to increase parallelism, and this practice will improve the throughput of a system if a system is well designed to accommodate the sample load. In Figure 17, the throughputs in total number of genomes per day for all tests with various numbers of 30x whole genome sequencing data are summarized. The tests performed here are designed to demonstrate performance at the server level, not for comparisons on individual components. At the same time, the tests were also designed to estimate the sizing information of Dell EMC Isilon F800/H600 and Dell EMC Lustre Storage. The data points in Figure 17 are calculated based on the total number of samples (X axis in the figure) that were processed concurrently. The number of genomes per day metric is obtained from total running time taken to process the total number of samples in a test. The smoothed curves are generated by using a polynomial spline with the piecewise polynomial degree of 3 generating B-spline basis matrix. The details of BWA-GATK pipeline information can be obtained from the Broad Institute web site (10).

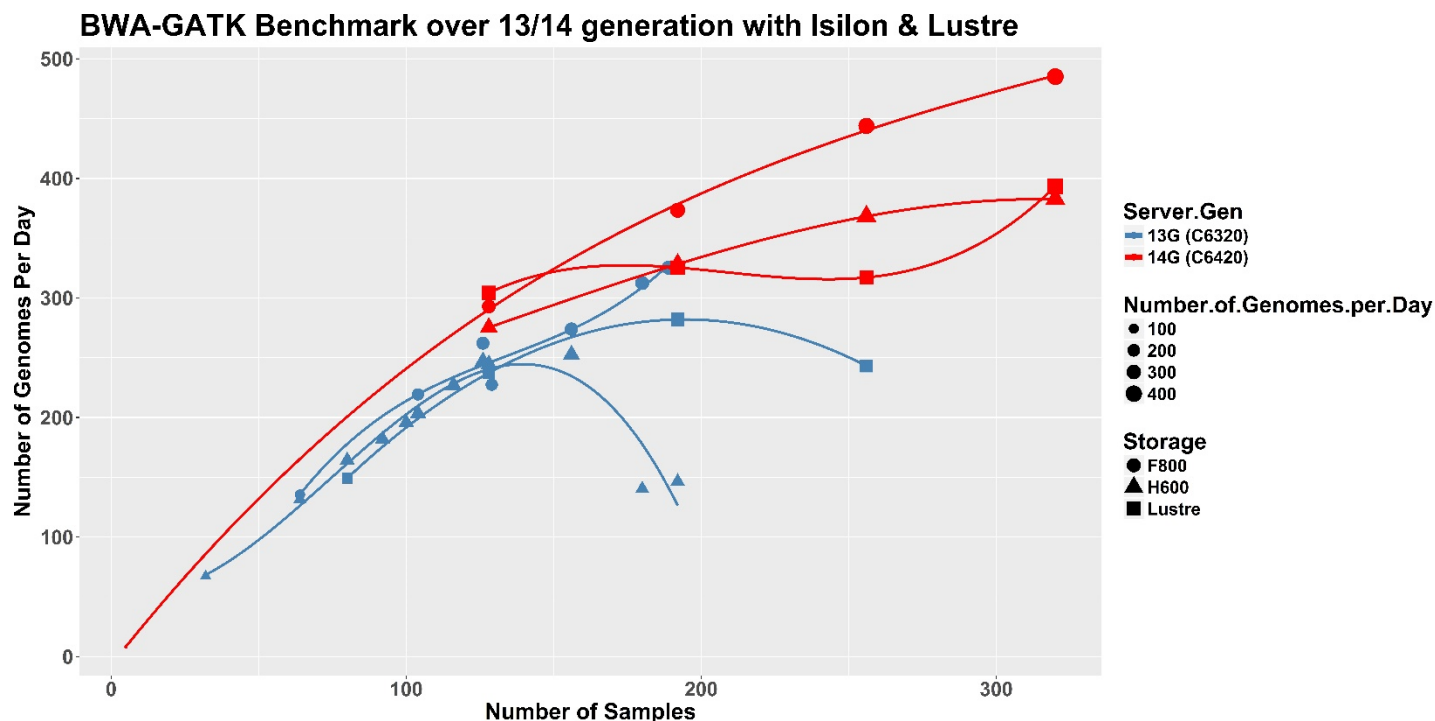


Figure 15 Performances in 13/14 generation servers with Isilon and Lustre

The number of compute nodes used for the tests are 64x C6420s and 63x C6320s (64x C6320s for testing H600). The number of samples per node was increased to get the desired total number of samples processed concurrently. For C6320 (13G), 3 samples per node was the maximum number of samples each node can process. 64, 104, and 126 test results for 13G system (blue) were with 2 samples per node while 129, 156, 180, 189 and 192 sample test results were obtained from 3 samples per node. For C6420 (14G), the tests were performed with maximum 5 samples per node. The plot for 14G was generated by processing 1, 2, 3, 4, and 5 samples per node. The number of samples per node is limited by the amount of memory in a system. 128 GB and 192 GB of RAM were used in 13G and 14G system, respectively. C6420s show a better scaling behavior than C6320s. 13G server with Broadwell CPUs seems to be more sensitive to the number of samples loaded onto system as shown from the results of 126 vs 129 sample tests on all the storages tested in this study.

Dell EMC PowerEdge C6420 has at least a 12% performance gain compared to the previous generation. Each C6420 compute node with 192 GB RAM can process about seven 30x whole human genomes per day. This number could be increased if the C6420 compute node is configured with more memory. In addition to the improvement on the 14G server side, four Isilon F800 nodes in a 4U chassis can support 64x C6420s and 320 30x whole human genomes concurrently.

MOLECULAR DYNAMICS SIMULATION SOFTWARE PERFORMANCE

Over the past decade, GPUs have become popular in scientific computing because of their great ability to exploit a high degree of parallelism. NVIDIA has a handful of life sciences applications optimized and to run on their general-purpose GPUs. Unfortunately, these GPUs can only be programmed with CUDA, OpenACC and the OpenCL framework. Most of the life sciences community is not familiar with these frameworks, and so few biologists or bioinformaticians can make efficient use of GPU architectures. However, GPUs have been making inroads into the molecular dynamics and electron microscopy fields. These fields require heavy computational work to simulate biomolecular structures or their interactions and reconstruct 3D images from millions of 2D images generated from an electron microscope.

We selected two different molecular dynamics applications to run tests on a PowerEdge C4130 with P100 and V100. The applications are Amber and LAMMPS (12) (13).

Molecular dynamics application test configuration

The Dell EMC PowerEdge C4130 with Intel® Xeon® Dual E5-2690 v4 with 256 GB DDR4 2400MHz and P100 and V100 in G and K configurations (14) (15). Unfortunately, we were not able to complete integration tests with Dell EMC Ready Bundle for HPC Life Sciences with various interconnect settings. However, we believe the integration tests will not pose any problem since molecular dynamics applications are not bounded by either storage I/O or inter-communication bandwidth.

The NVIDIA® Tesla® V100 accelerator is one of the most advanced accelerators available in the market right now and was launched within one year of the P100 release. In fact, Dell EMC is the first in the industry to integrate Tesla V100 and bring it to market. As was the case with the P100, V100 supports two form factors: V100-PCIe and the mezzanine version V100-SXM2. The Dell EMC PowerEdge C4130 server supports both types of V100 and P100 GPU cards.

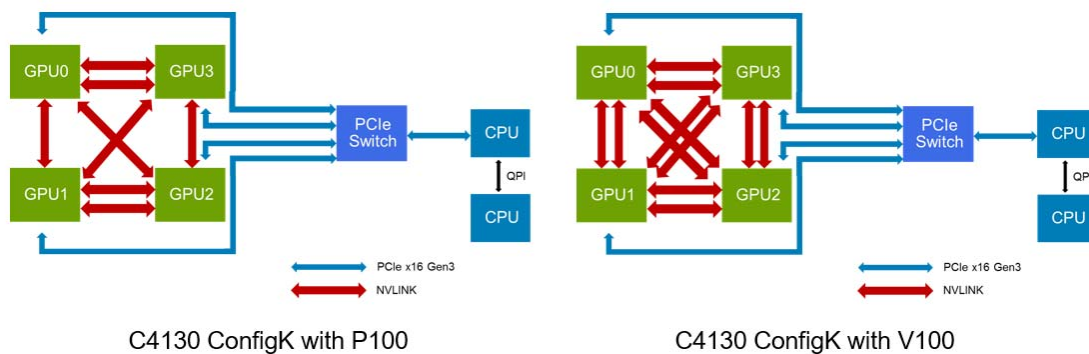


Figure 16 V100 and P100 Topologies on C4130 configuration K

Amber benchmark suite

This suite includes the Joint Amber-Charmm (JAC) benchmark considering dihydrofolate reductase (DHFR) in an explicit water bath with cubic periodic boundary conditions. The major assumptions are that the DHFR molecule presents in water without surface effect and its movement assumed to follow microcanonical (NVE) ensemble which assumes constant amount of substance (N), volume (V), and energy (E). Hence, the sum of kinetic (KE) and potential energy (PE) is conserved, in other words, Temperature (T) and Pressure (P) are unregulated. JAC benchmark repeats simulations with Isothermal-isobaric (NPT) ensemble that assumes N, P and T are conserved. It corresponds most closely to laboratory conditions with a flask open to ambient temperature and pressure. Beside these settings, Particle mesh Ewald (PME) is the choice of algorithm to calculate electrostatic forces in molecular dynamics simulations. Other biomolecules simulated in this benchmark suite are Factor IX (one of the serine proteases of the coagulation system), cellulose and Satellite Tobacco Mosaic Virus (STMV). Here, we report the results from DHFR and STMV data.

Figure 19

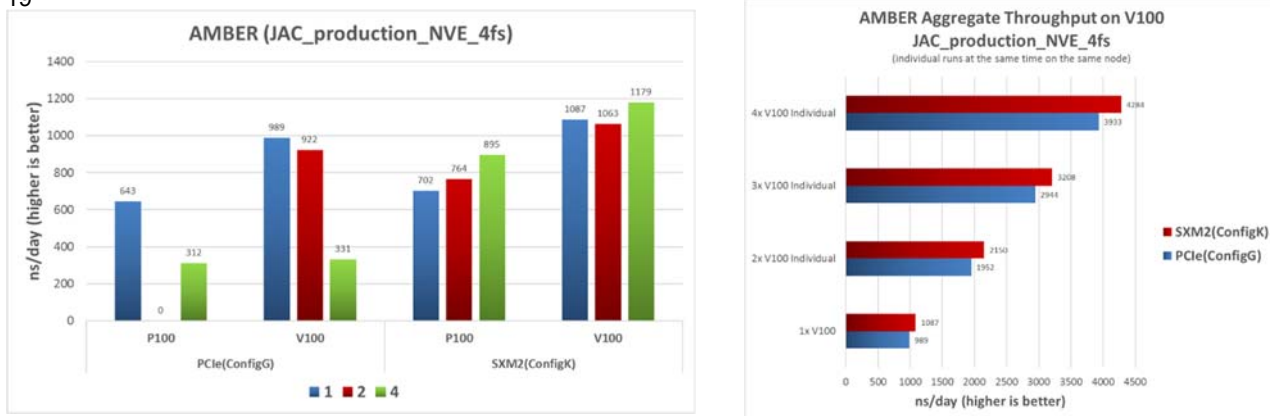


Figure 17 AMBER JAC Benchmark

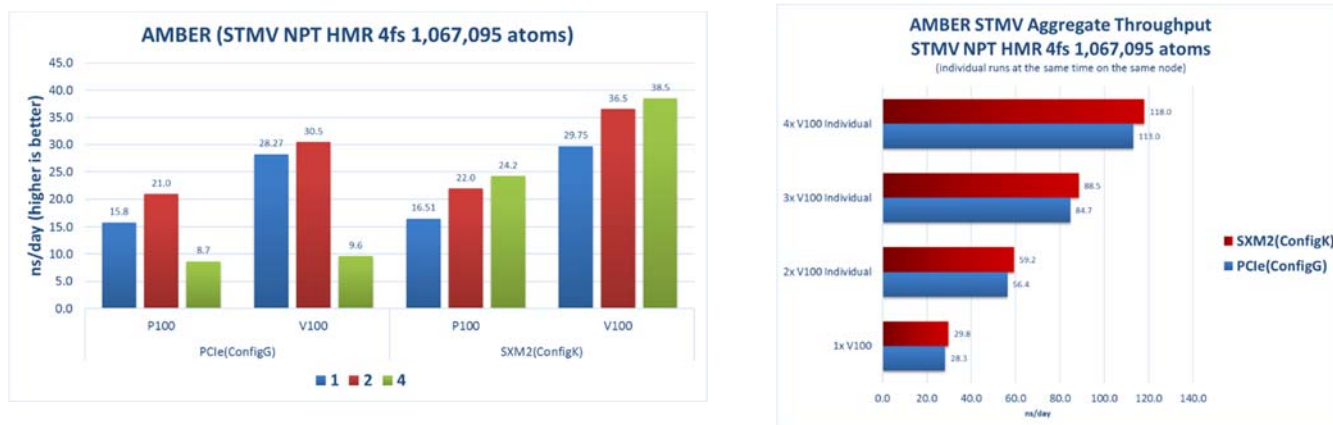


Figure 18 AMBER STMV benchmark

Figure 19 and Figure 20 illustrate AMBER's results with DHFR and STMV dataset. On SXM2 system (Config K), AMBER scales weakly with 2 and 4 GPUs. Even though the scaling is not strong, V100 has noticeable improvement than P100, giving ~78% increase in single card runs, and 1x V100 is actually 23% faster than 4x P100. On the PCIe (Config G) side, one and two cards perform similar to SXM2; however, four cards' results dropped sharply. This is because PCIe (Config G) only supports Peer-to-Peer access between GPU0/1 and GPU2/3 and not among all four GPUs. Since AMBER has redesigned the way data transfers among GPUs to address the PCIe bottleneck, it relies heavily on Peer-to-Peer access for performance with multiple GPU cards. Hence a fast, direct interconnect like NVLink between all GPUs in SXM2 (Config K) is vital for AMBER multiple GPU performance. To compensate for a single job's weak scaling on multiple GPUs, there is another use case promoted by AMBER developers, which is running multiple jobs in the same node

concurrently but where each job uses only 1 or 2 GPUs. Figure 5 shows the results of 1-4 individual jobs on one C4130 with V100s and the numbers indicate that those individual jobs have little impact on each other. This is because AMBER is designed to run pretty much entirely on the GPUs and has very low dependency on the CPU. The aggregate throughput of multiple individual jobs scales linearly in this case. Without any card to card communication, the 5% better performance on SXM2 is contributed by its higher clock speed.

LAMMPS

Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) is a classical molecular dynamics code and has potentials for solid-state materials (metals, semiconductors) and soft matter (biomolecules, polymers) and coarse-grained or mesoscopic systems. It can be used to model atoms or, more generically, as a parallel particle simulator at the atomic, meso, or continuum scale. It runs on single processors or in parallel using message-passing techniques and a spatial-decomposition of the simulation domain. Many of its models have versions that provide accelerated performance on CPUs, GPUs, and Intel Xeon Phis. The code is designed to be easy to modify or extend with new functionality.

We performed a 3D Lennard-Jones melt simulation package which comes with LAMMPS. It simulates the movement of molecules confined in a squared box. LAMMPS has built-in functions for placing molecules inside the box and moving them according to Newton's laws. This is arguably one of the simplest models capable of reproducing the complete thermodynamic behavior of classical fluids. The details of the simulation set-up can be found in Appendix B.

Figure 21 shows LAMMPS performance on both configurations G and K. The testing dataset is Lennard-Jones liquid dataset, which contains 512,000 atoms, and LAMMPS compiled with the kokkos package. V100 is 71% and 81% faster on Config G and Config K respectively. Comparing V100-SXM2 (Config K) and V100-PCIe (Config G), the former is 5% faster due to NVLINK and higher CUDA core frequency.

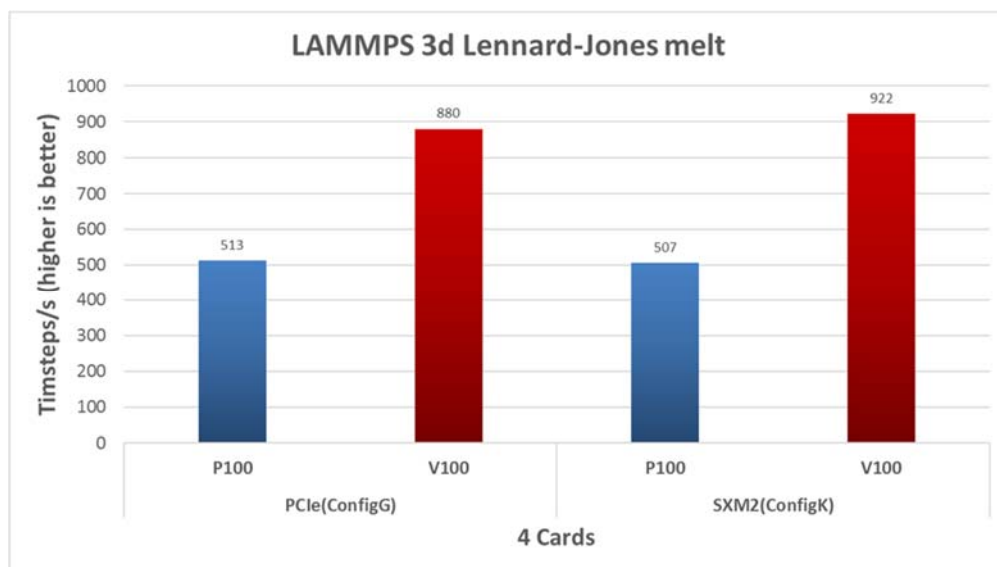


Figure 19 LAMMPS 3D Lennard-Jones melt simulation

The C4130 server with NVIDIA® Tesla® V100™ GPUs demonstrates exceptional performance for HPC applications that require faster computational speed and highest data throughput. Applications like AMBER and LAMMPS were boosted with C4130 configuration K, owing to P2P access, higher bandwidth of NVLink and higher CUDA core clock speed. Overall, a PowerEdge C4130 with Tesla V100 GPUs performs 1.54x to 1.8x faster than a C4130 with P100 for AMBER and LAMMPS.

CRYO-EM PERFORMANCE

The purpose of this study was to validate the optimized Relion (for REgularised Likelihood Optimization) on Dell EMC PowerEdge C6420s with Skylake CPUs. Relion was developed from the Scheres lab at MRC Laboratory of Molecular Biology. It uses an empirical Bayesian approach to refining multiple 3D images or 2D class averages for the data generated from CryoElectron Microscopy (Cryo-EM). The impressive performance gain from Intel®'s efforts in the collaboration of Relion development team reduced the performance gap between CPUs and GPUs. The CPU/GPU performance comparison results are not shown here; however, the performance gap becomes single digit fold between Skylake CPU systems and Broadwell CPU/Tesla P100 GPU systems.

Essentially, Cryo-EM is a type of transmission electron microscopy (TEM) for imaging frozen-hydrated specimens at cryogenic temperatures. Specimens remain in their native state without the need for dyes or fixatives, allowing the study of fine cellular structures, viruses and protein complexes at molecular resolution. A rapid vitrification at cryogenic temperature is the key step to avoid water molecule crystallization and forming amorphous solid that does almost no damage to the sample structure. Regular electron microscopy requires samples to be prepared in complex ways, and the sample preparations make hard to retaining the original molecular structures. Cryo-EM is not perfect like X-ray crystallography; however, it has quickly gained the popularity in the research community due to the simple sample preparation steps and flexibility of the sample size, complexity, and non-rigid structure. As the resolution revolution in Cryo-EM progresses due to the 40+ years of dedicated work from the structural biology community, we now can yield accurate, detailed 3D models of intricate biological structures at the sub-cellular and molecular scales.

The tests were performed on 8 nodes of Dell PowerEdge C6420s which is a part of Dell EMC Ready Bundle for HPC Life Sciences.

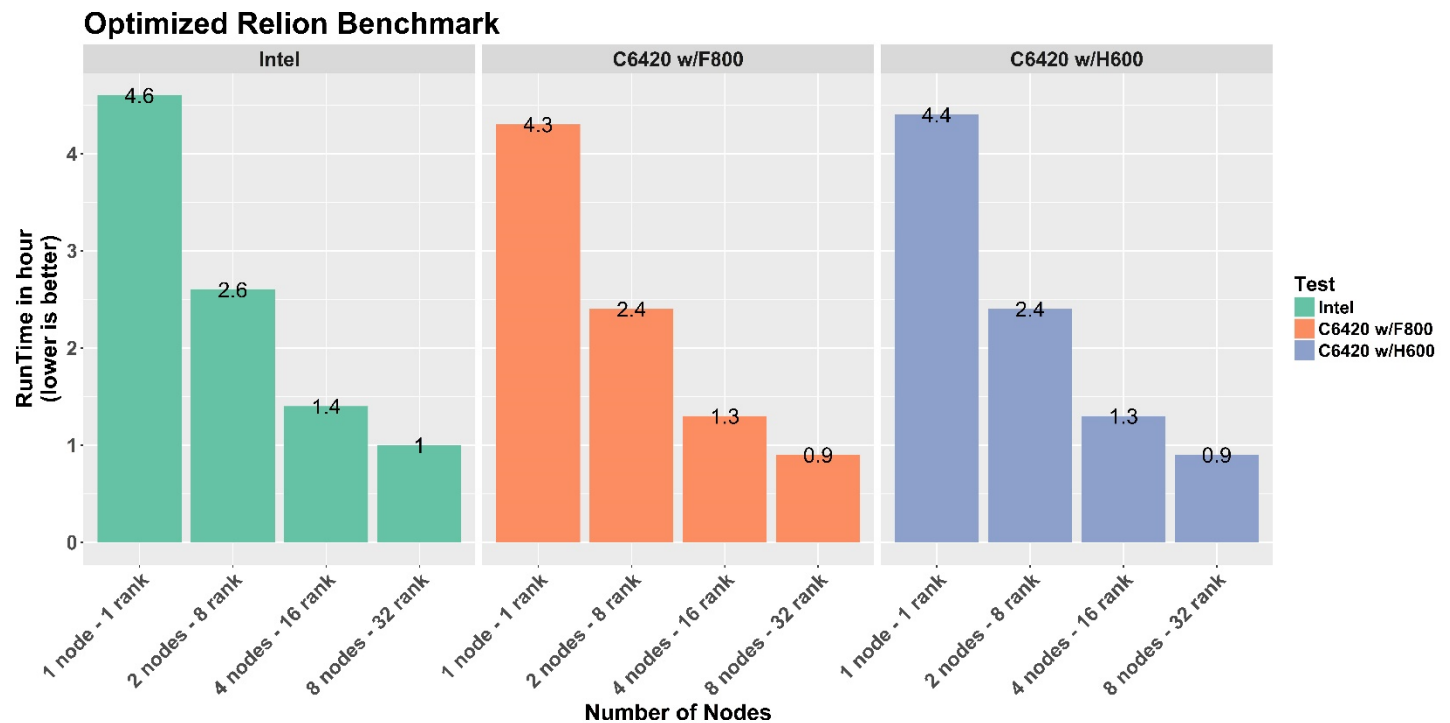


Figure 20 Optimized Relion Benchmark

Dell EMC PowerEdge C6420 shows that it is an ideal compute platform for the Optimized Relion. It scales well over various number of compute nodes with Plasmodium ribosome data. In the future study, we plan to use a larger protein data and more compute nodes to accomplish more comprehensive scaling tests.

CONCLUSION

Overall, 14th generation servers with Skylake and larger/faster memory size (due to higher number of memory channels compare with Broadwell) show a better throughput on BWA-GATK pipeline. The throughput for this type of work improved from four 30x genomes per day per C6320 to seven 30x genomes per day per C6420. Also, we verified that Dell EMC Isilon storage, F800 and H600, can be used for high-performance scratch storage while they provide all the conveniences of easy maintenance, scaling, and various supported file systems. In addition to that, we observed a better performance on Cryo-EM data process with Intel®'s optimized Relion codes. Unfortunately, we could not test Dell EMC PowerEdge C4140 with NVIDIA® Tesla™ V100; however, the V100 with PowerEdge C4130 still shows impressive performance compared to P100. We believe the current version of Dell EMC Ready Bundle for HPC Life Sciences is ready for data centric high-performance computing.

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ⁱ The tested sequencing data's depth of coverage is 30x when we assume that the sequence reads are generated from germ cells which are haploids.

ⁱⁱ For the best performance for the Optimized Relion application, Logical Processor should be enabled.