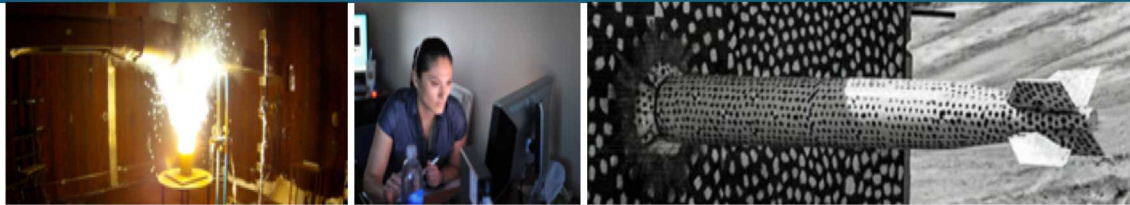


SAND2019-XXXXX



SAND2019-2243C

Performance portable parallel sparse CP-APR tensor decompositions



SIAM CSE19, 03/01/2019

PRESENTED BY

Keita Teranishi

**Christopher Forster (NVIDIA), Richard Barrett,
Daniel Dunlavy, and Tamara Kolda**



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Develop production quality library software to perform CP factorization for **Poisson Regression Problems** for HPC platforms

Tensor Tool Box (<http://www.tensortoolbox.org>)

- Matlab only!

Support several HPC platforms

- Node parallelism (Multicore, Manycore and GPUs)

Major Questions

- Software Design
- Performance Tuning

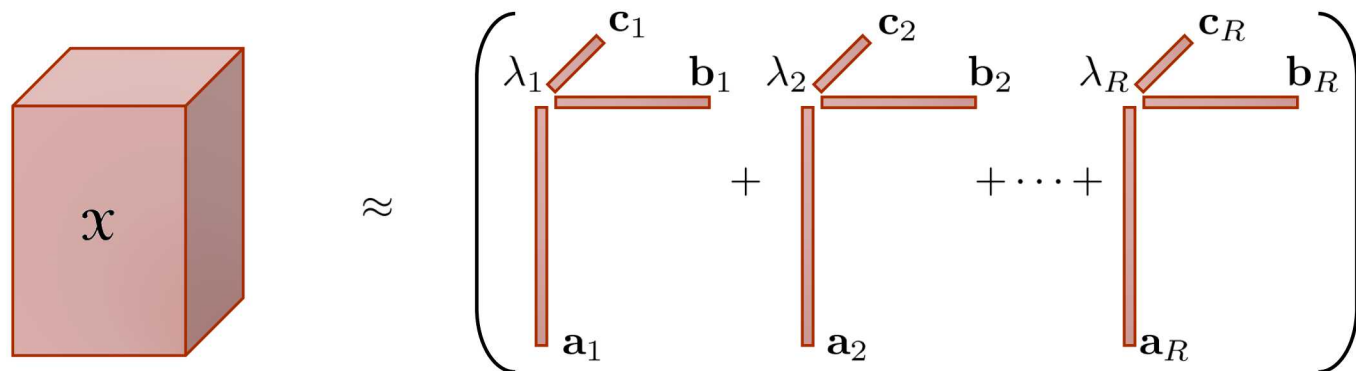
This talk

- We are interested in two major variants
 - **Multiplicative Updates**
 - Projected Damped Newton for Row-subproblems



CANDECOMP/PARAFAC (CP) Model

Express the important feature of data using a small number of vector outer products



$$\mathcal{X} \approx \left(\lambda_1 \mathbf{a}_1 \circ \mathbf{b}_1 \circ \mathbf{c}_1 + \lambda_2 \mathbf{a}_2 \circ \mathbf{b}_2 \circ \mathbf{c}_2 + \dots + \lambda_R \mathbf{a}_R \circ \mathbf{b}_R \circ \mathbf{c}_R \right)$$

Model: $\mathcal{M} = \sum_r \lambda_r \mathbf{a}_r \circ \mathbf{b}_r \circ \mathbf{c}_r$

$$x_{ijk} \approx m_{ijk} = \sum_r \lambda_r a_{ir} b_{jr} c_{kr}$$

Key references: Hitchcock (1927), Harshman (1970), Carroll and Chang (1970)

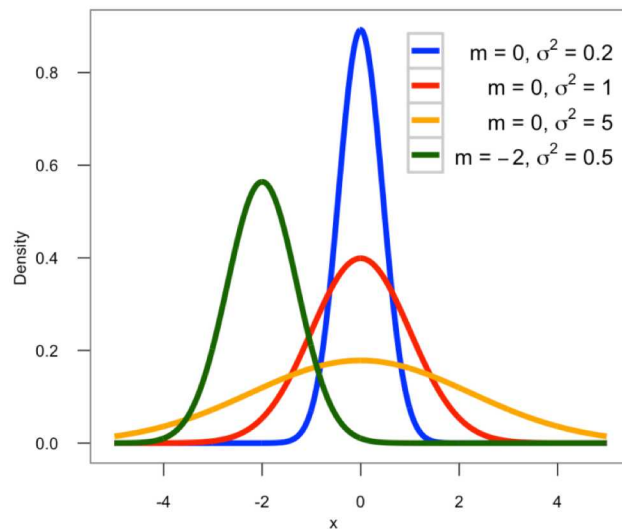


Gaussian (typical)

The random variable x is a continuous real-valued number.

$$x \sim N(m, \sigma^2)$$

$$P(X = x) = \frac{\exp(-\frac{(x-m)^2}{2\sigma^2})}{\sqrt{2\pi\sigma^2}}$$

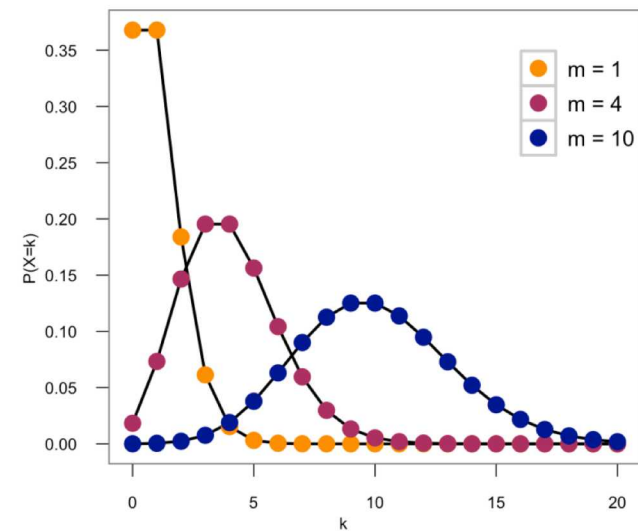


Poisson

The random variable x is a discrete nonnegative integer.

$$x \sim \text{Poisson}(m)$$

$$P(X = x) = \frac{\exp(-m)m^x}{x!}$$



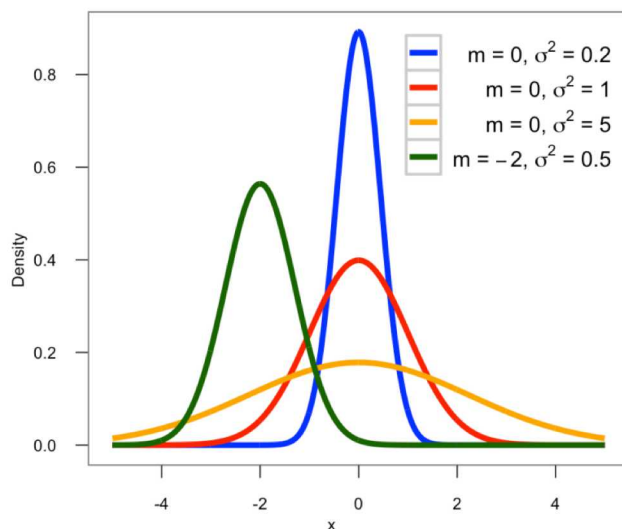


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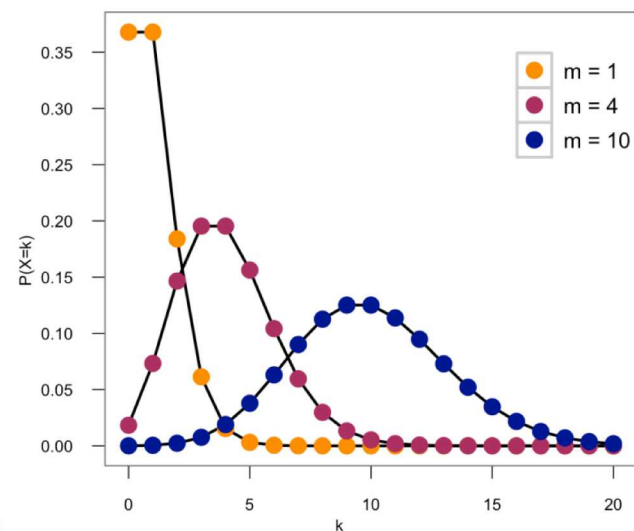


Poisson

The random variable x is a discrete nonnegative integer.

$$x \sim \text{Poisson}(m)$$

$$P(X = x) = \frac{\exp(-m)m^x}{x!}$$





$$\mathcal{X} \sim \text{Poisson} \left(\lambda_1 \begin{matrix} \text{---} \mathbf{c}_1 \\ | \quad \text{---} \mathbf{b}_1 \\ | \quad \text{---} \mathbf{a}_1 \end{matrix} + \lambda_2 \begin{matrix} \text{---} \mathbf{c}_2 \\ | \quad \text{---} \mathbf{b}_2 \\ | \quad \text{---} \mathbf{a}_2 \end{matrix} + \dots + \lambda_R \begin{matrix} \text{---} \mathbf{c}_R \\ | \quad \text{---} \mathbf{b}_R \\ | \quad \text{---} \mathbf{a}_R \end{matrix} \right)$$

Model: Poisson distribution (nonnegative factorization)

$$x_{ijk} \sim \text{Poisson}(m_{ijk}) \text{ where } m_{ijk} = \sum_r \lambda_r a_{ir} b_{jr} c_{kr}$$

- Nonconvex problem!
 - Assume R is given
- Minimization problem with constraint
 - The decomposed vectors must be non-negative
- Alternating Poisson Regression (Chi and Kolda, 2011)
 - Assume $(d-1)$ factor matrices are known and solve for the remaining one

Alternating Poisson Regression (CP-APR)



Repeat until converged...

1. $\bar{\mathbf{A}} \leftarrow \arg \min_{\bar{\mathbf{A}} \geq 0} \sum_{ijk} m_{ijk} - x_{ijk} \log m_{ijk} \text{ s.t. } \mathcal{M} = \sum_r \bar{\mathbf{a}}_r \circ \mathbf{b}_r \circ \mathbf{c}_r$
 2. $\lambda \leftarrow \mathbf{e}^\top \bar{\mathbf{A}}; \mathbf{A} \leftarrow \bar{\mathbf{A}} \cdot \text{diag}(1/\lambda)$
 3. $\bar{\mathbf{B}} \leftarrow \arg \min_{\bar{\mathbf{B}} \geq 0} \sum_{ijk} m_{ijk} - x_{ijk} \log m_{ijk} \text{ s.t. } \mathcal{M} = \sum_r \mathbf{a}_r \circ \bar{\mathbf{b}}_r \circ \mathbf{c}_r$
 4. $\lambda \leftarrow \mathbf{e}^\top \bar{\mathbf{B}}; \mathbf{B} \leftarrow \bar{\mathbf{B}} \cdot \text{diag}(1/\lambda)$
 5. $\bar{\mathbf{C}} \leftarrow \arg \min_{\bar{\mathbf{C}} \geq 0} \sum_{ijk} m_{ijk} - x_{ijk} \log m_{ijk} \text{ s.t. } \mathcal{M} = \sum_r \mathbf{a}_r \circ \mathbf{b}_r \circ \bar{\mathbf{c}}_r$
 6. $\lambda \leftarrow \mathbf{e}^\top \bar{\mathbf{C}}; \mathbf{C} \leftarrow \bar{\mathbf{C}} \cdot \text{diag}(1/\lambda)$
- Fix $\mathbf{B}, \mathbf{C}$; solve for $\mathbf{A}$
- Fix $\mathbf{A}, \mathbf{C}$; solve for $\mathbf{B}$
- Fix $\mathbf{A}, \mathbf{B}$; solve for $\mathbf{C}$

Convergence
Theory

Theorem: The CP-APR algorithm will **converge to a constrained stationary point** if the subproblems are strictly convex and solved exactly at each iteration. (Chi and Kolda, 2011)

8 Accuracy is High For Very Sparse Data



Data: 1000 x 800 x 600 Tensor with R=10 Components

CP-APR: Max Iterations = 200, Max Inner Iterations = 30 (10 per mode), Tol = $1e-4$ (KKT)

CP-ALS: Max Iterations = 200, Tol = $1e-8$ (change in fit)

Nonzeros	Poisson Regression FMS	Gaussian Regression FMS
480,000 (.100%)	0.99	0.57
240,000 (.050%)	0.81	0.49
48,000 (.010%)	0.77	0.47
24,000 (.005%)	0.74	0.46



Algorithm 1: CPAPR, Alternating Block Framework

```

1 CPAPR ( $\mathcal{X}, \mathcal{M}$ );
   Input : Sparse  $N$ -mode Tensor  $\mathcal{X}$  of size  $I_1 \times I_2 \times \dots \times I_N$  and the
           number of components  $R$ 
   Output: Kruskal Tensor  $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$ 
2 Initialize
3 repeat
4   for  $n = 1, \dots, N$  do
5     Let  $\Pi^{(n)} = (A^{(N)} \odot \dots \odot A^{(n+1)} \odot A^{(n-1)} \odot \dots \odot A^{(1)})^T$ 
6     Compute  $\bar{A}^{(n)}$  that minimize  $f(\bar{A}^{(n)})$  s.t.  $\bar{A}^{(n)} \geq 0$ 
7      $A^{(n)} \leftarrow \bar{A}^{(n)}$ 
8   end
9 until all mode subproblems converged;

```

Minimization problem is expressed as:

$$\min_{\bar{A}^{(n)} \geq 0} f(\bar{A}^{(n)}) = e^T [\bar{A}^{(n)} \Pi^{(n)} - X_{(n)} * \log(\bar{A}^{(n)} \Pi^{(n)})] e$$



$\odot$ is called Khatori-Rao product
(Column wise Kronecker product)

$$C = [C_1 | C_2 | C_3]$$

$$D = [D_1 | D_2 | D_3]$$

$$C \odot D = [C_1 \otimes D_1 | C_2 \otimes D_2 | C_3 \otimes D_3]$$

Algorithm 1: CPAPR

1 CPAPR ($\mathcal{X}, \mathcal{M}$);

Input : Sparse N -mode Tensor $\mathcal{X}$ of size $I_1 \times I_2 \times \dots \times I_N$ and the number of components R

Output: Kruskal Tensor $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$

2 **Initialize**

3 **repeat**

4 **for** $n = 1, \dots, N$ **do**

5 Let $\Pi^{(n)} = (A^{(N)} \odot \dots \odot A^{(n+1)} \odot A^{(n-1)} \odot \dots \odot A^{(1)})^T$

6 Compute $\bar{A}^{(n)}$ that minimize $f(\bar{A}^{(n)})$ s.t. $\bar{A}^{(n)} \geq 0$

7 $A^{(n)} \leftarrow \bar{A}^{(n)}$

8 **end**

9 **until** *all mode subproblems converged*;

Minimization problem is expressed as:

$$\min_{\bar{A}^{(n)} \geq 0} f(\bar{A}^{(n)}) = e^T [\bar{A}^{(n)} \Pi^{(n)} - X_{(n)} * \log(\bar{A}^{(n)} \Pi^{(n)})] e$$



$\odot$ is called Khatori-Rao product
(Column wise Kronecker product)

$$C = [C_1 | C_2 | C_3]$$

$$D = [D_1 | D_2 | D_3]$$

$$C \odot D = [C_1 \otimes D_1 | C_2 \otimes D_2 | C_3 \otimes D_3]$$

Algorithm 1: CPAPR

1 CPAPR ($\mathcal{X}, \mathcal{M}$);

Input : Sparse N -m

 number of components n

Output: Kruskal Tensor $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$

2 **Initialize**

3 **repeat**

4 **for** $n = 1, \dots, N$ **do**

5 Let $\Pi^{(n)} = (A^{(N)} \odot \dots \odot A^{(n+1)} \odot A^{(n-1)} \odot \dots \odot A^{(1)})^T$

6 Compute $\bar{A}^{(n)}$ that minimize $f(\bar{A}^{(n)})$ s.t. $\bar{A}^{(n)} \geq 0$

7 $A^{(n)} \leftarrow \bar{A}^{(n)}$

8 **end**

9 **until** *all mode subproblems converged*;

Π is expressed in sparse matrix (indices and values).

Minimization problem is expressed as:

$$\min_{\bar{A}^{(n)} \geq 0} f(\bar{A}^{(n)}) = e^T [\bar{A}^{(n)} \Pi^{(n)} - X_{(n)} * \log(\bar{A}^{(n)} \Pi^{(n)})] e$$



$\odot$ is called Khatori-Rao product
(Column wise Kronecker product)

$$C = [C_1 | C_2 | C_3]$$

$$D = [D_1 | D_2 | D_3]$$

$$C \odot D = [C_1 \otimes D_1 | C_2 \otimes D_2 | C_3 \otimes D_3]$$

Algorithm 1: CPAPR

1 CPAPR ($\mathcal{X}, \mathcal{M}$);

Input : Sparse N -m
number of components n

Π is expressed in sparse matrix (indices and values).

Output

2 **Initial**

3 **repeat**

4 **for**

5

6

7

8 **end**

9 **until**

Minimize

1

- 2 major approaches

- **Multiplicative Updates** like Lee & Seung (2000) for matrices, but extended by E. C. Chi and T. G. Kolda. *On Tensors, Sparsity, and Nonnegative Factorizations*, SIAM Journal on Matrix Analysis and Applications 33(4):1272-1299, December 2012.
- **Newton and Quasi-Newton method for Row-subblems** by S. Hansen, T. Plantenga and T. G. Kolda. *Newton-Based Optimization for Kullback-Leibler Nonnegative Tensor Factorizations*, to appear in Optimization Methods and Software, 2015.



Multiplicative Update (MU)

Projected Damped Newton for Row-subproblems (PDNR)

Key computations

- Khatri-Rao Product $\Pi^{(n)}$
- Multiplicative Update Modifier (10+ iterations)

Key computations

- Khatri-Rao Product $\Pi^{(n)}$
- Constrained Non-linear Newton-based optimization for each row

Key features

- Factor matrix is updated all at once
- Exploits the convexity of row subproblems for global convergence

Key features

- Factor matrix can be updated by rows
- Exploits the convexity of row-subproblems



Algorithm 1: CP-APR-MU, Multiplicative Update

```

1 CP-APR-MU ( $\mathcal{X}, \mathcal{M}$ );
   Input : Sparse  $N$ -mode Tensor  $\mathcal{X}$  of size  $I_1 \times I_2 \times \dots \times I_N$  and the
           number of components  $R$ 
   Output: Kruskal Tensor  $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$ 
2 Initialize
3 repeat
4   for  $n = 1, \dots, N$  do
5      $B \leftarrow (A^{(n)} + S)\Lambda$  ( $S$  is used to remove inadmissible zeros)
6     Let  $\Pi^{(n)} = (A^{(N)} \odot \dots \odot A^{(n+1)} \odot A^{(n-1)} \odot \dots \odot A^{(1)})^T$ 
7     for  $i = 1, \dots, 10$  do
8        $\Phi^{(n)} \leftarrow (X_{(n)} \oslash \max(B\Pi^{(n)}, \epsilon))(\Pi^{(n)})^T$ 
9        $B \leftarrow B * \Phi^{(n)}$ 
10    end
11     $\lambda = e^T B$ 
12     $A^{(n)} \leftarrow B\Lambda^{-1}$ , where  $\Lambda = \text{diag}(\lambda)$ 
13  end
14 until all mode subproblems converged;
  
```

Key Computations



Algorithm 1: CPAPR-PDNR algorithm

```

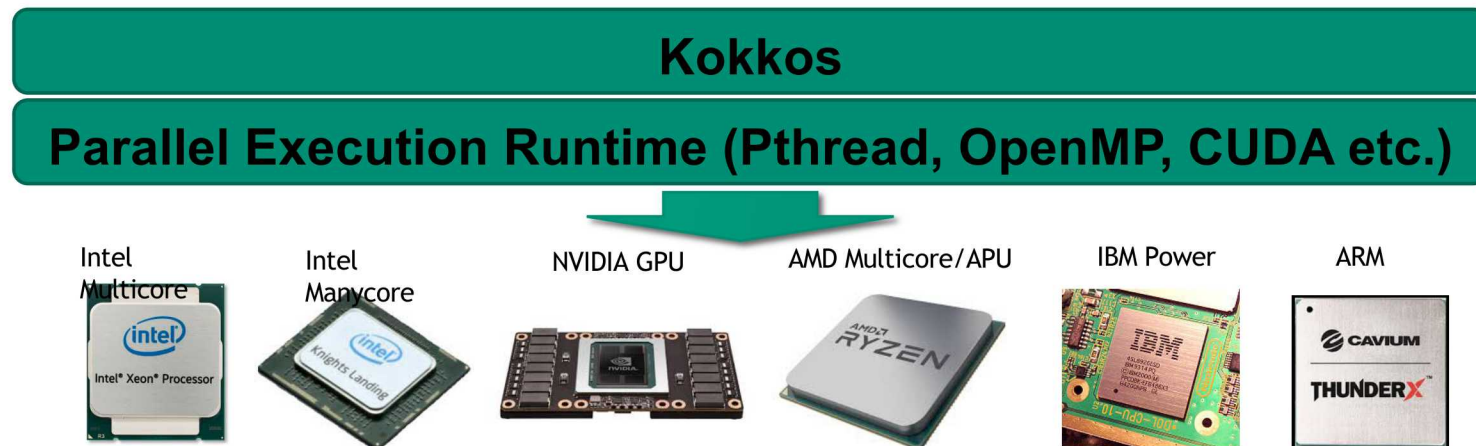
1 CPAPR_PDNR ( $\mathcal{X}, \mathcal{M}$ );
   Input : Sparse  $N$ -mode Tensor  $\mathcal{X}$  of size  $I_1 \times I_2 \times \dots \times I_N$  and the
           number of components  $R$ 
   Output: Kruskal Tensor  $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$ 
2 Initialize
3 repeat
4   for  $n = 1, \dots, N$  do
5     Let  $\Pi^{(n)} = (A^{(N)} \odot \dots \odot A^{(n+1)} \odot A^{(n-1)} \odot \dots \odot A^{(1)})^T$ 
6     for  $i = 1, \dots, I_n$  do
7       Find  $b_i^{(n)}$  s.t.  $\min_{b_i^{(n)} \geq 0} f_{\text{row}}(b_i^{(n)}, x_i^{(n)}, \Pi^{(n)})$ 
8     end
9      $\lambda = e^T B^{(n)}$  where  $B^{(n)} = [b_1^{(n)} \dots b_{I_n}^{(n)}]^T$ 
10     $A^{(n)} \leftarrow B^{(n)} \Lambda^{-1}$ , where  $\Lambda = \text{diag}(\lambda)$ 
11  end
12 until all mode subproblems converged;
  
```

Key Computations



Focus on on-node parallelism for multiple architectures

- Multiple choices for programming
 - OpenMP, OpenACC, CUDA, Pthread ...
 - Manage different low-level hardware features (cache, device memory, NUMA...)
- Our Solution: **Use Kokkos for productivity and performance portability**
 - Abstraction of parallel loops
 - Abstraction Data layout (row-major, column major, programmable memory)
 - Same code to support multiple architectures





Templated C++ Library by Sandia National Labs (Edwards, et al)

- Serve as substrate layer of sparse matrix and vector kernels
- Support any machine precisions
 - Float, Double, etc

Kokkos::View() accommodates performance-aware multidimensional array data objects

- Light-weight C++ class to accommodate abstractions for platform specific features (host, device, GPU's shared memory, data access pattern, etc.)

Parallelizing loops using C++ language standard

- **Lambda**
- Functors

Extensive support of atomics



Serial

```
for (size_t i = 0; i < N; ++i)
{
    /* loop body */
}
```

OpenMP

```
#pragma omp parallel for
for (size_t i = 0; i < N; ++i)
{
    /* loop body */
}
```

Kokkos

```
parallel_for (( N, [=], (const size_t i)
{
    /* loop body */
});
```

Kokkos information courtesy of Carter Edwards

Provide parallel loop operations using C++ language features

Conceptually, the usage is no more difficult than OpenMP. The annotations just go in different places.

Support for task parallel computing is ongoing (Task Parallel Kokkos and UINTHA)



Algorithm 1: CP-APR-MU in source

```

1 CP-APR-MU  $X, M, R$ ;
   Input : Sparse  $N$ -mode Tensor  $X$  of size  $I_1 \times I_2 \times \dots \times I_N$  and the
           number of components  $R$ 
   Output: Kruskal Tensor  $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$ 
2 initializeBuffer( $X, R$ )
3  $\mathcal{E} \leftarrow \text{computeIndexMap}(X)$ 
4 repeat
5   for  $n = 1, \dots, N$  do
6      $M \leftarrow \text{offset}(M, n)$  (Remove inadmissible zeros)
7      $M \leftarrow \text{distribute}(M, n)$  (Scale the elements of  $A^n$  by  $\lambda$ )
8      $\Pi^{(n)} \leftarrow \text{computePi}(M, \mathcal{E}^{(n)})$ 
9     for  $i = 1, \dots, 10$  do
10       $\Phi_i^{(n)} \leftarrow \text{computePhi}(A_i^{(n)}, \Pi^{(n)}, \mathcal{E}^{(n)})$ 
11       $A_{i+1}^{(n)} \leftarrow A_i^{(n)} \Phi_i^{(n)}$ 
12    end
13     $M \leftarrow \text{normalize}(M, A, n)$ 
14  end
15 until all mode subproblems converged;

```

Data Parallel



Algorithm 1: CP-APR-PDNR in source

```

1 CP-APR-PDNR  $X, M, R$ ;
   Input : Sparse  $N$ -mode Tensor  $X$  of size  $I_1 \times I_2 \times \dots \times I_N$  and the
           number of components  $R$ 
   Output: Kruskal Tensor  $\mathcal{M} = [\lambda; A^{(1)} \dots A^{(N)}]$ 
2 initializeBuffer( $X, R$ )
3  $\mathcal{E} \leftarrow \text{computeIndexMap}(X)$ 
4 repeat
5   for  $n = 1, \dots, N$  do
6      $M \leftarrow \text{distribute}(M, n)$  (Scale the elements of  $A^n$  by  $\lambda$ )
7      $\Pi^{(n)} \leftarrow \text{computePi}(A, \mathcal{E}^{(n)})$ 
8     parallel_for  $i = 1, \dots, I_n$  do
9        $a_i^n \leftarrow \text{rowSolvePDNR}(a_i^n, X^n, \Pi^n, \mathcal{E}_i^{(n)})$ 
10    end
11     $M \leftarrow \text{normalize}(M, A, n)$ 
12  end
13 until all mode subproblems converged;
  
```

Data Parallel

Task Parallel



Use Kokkos::View for all data structures

Sparse Tensor

- Similar to the Coordinate (COO) Format in Sparse Matrix representation

Atomics

- Expensive for CPUs and Manycore
 - Data Rows of Factor Matrices
- Efficient for the latest GPUs

CP-APR-PDNR

- **Nested Parallelism**
 - Top Level: Individual Newton Solve
 - Second Level: Vectorized operations

Mode-1	1	1	1	1	2	2	4	1	3	3	5	4
Mode-2	3	4	1	3	5	2	4	2	3	5	1	4
Mode-3	5	1	2	2	2	3	3	1	5	4	1	4
INDICES (_indices)												
Nonzero Entries (_data)	0.2	0.5	0.1	0.7	0.9	2.1	0.4	0.5	0.1	0.4	0.6	0.1



Modifier Computation is the major part of CP-APR-MU.

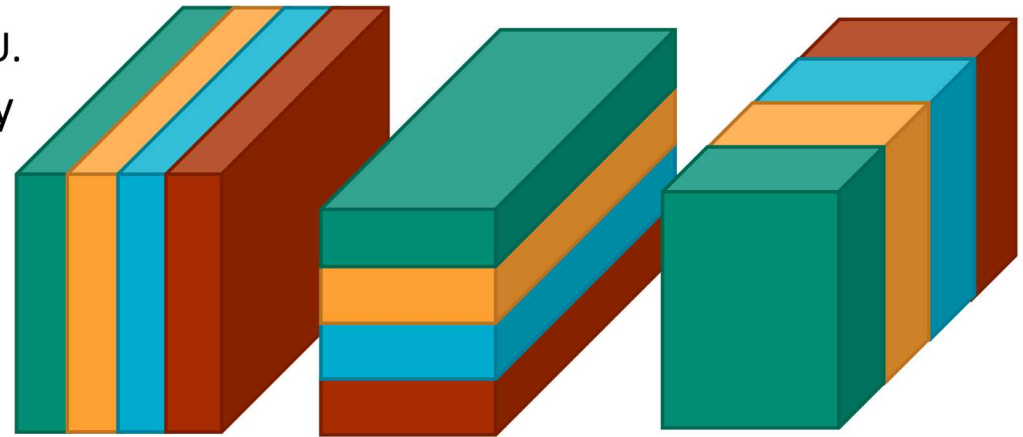
- **Two ways** to parallelize, which affects the way to access the output factor matrices

1. Partition with respect to the mode

- **No atomics** to access the output vectors by partition
- **Extra indexing** is required to access nonzero entries by partition (**reordering**)

2. Partition COO sparse tensor storage format

- **No extra indexing** is required
 - Need efficient hardware supported **atomics**
 - The output vector elements are updated by multiple threads concurrently
 - Large outermost loop irrespective of the mode sizes
- Recent work by Smith and Karypis, and Li and Vuduc suggest more efficient data format than COO



Mode-1	1	1	1	1	2	2	4	1	3	3	5	4
Mode-2	3	4	1	3	5	2	4	2	3	5	1	4
Mode-3	5	1	2	2	2	3	3	1	5	4	1	4
INDICES (_indices)												
Nonzero Entries (_data)	0.2	0.5	0.1	0.7	0.9	2.1	0.4	0.5	0.1	0.4	0.6	0.1



Strong Scalability

- Problem size is fixed

Random Tensor

- 3K x 4K x 5K, 10M nonzero entries
- **100 outer iterations**

Realistic Problems

- Count Data (Non-negative)
- Available at <http://frostdt.io/>
- **10 outer iterations**

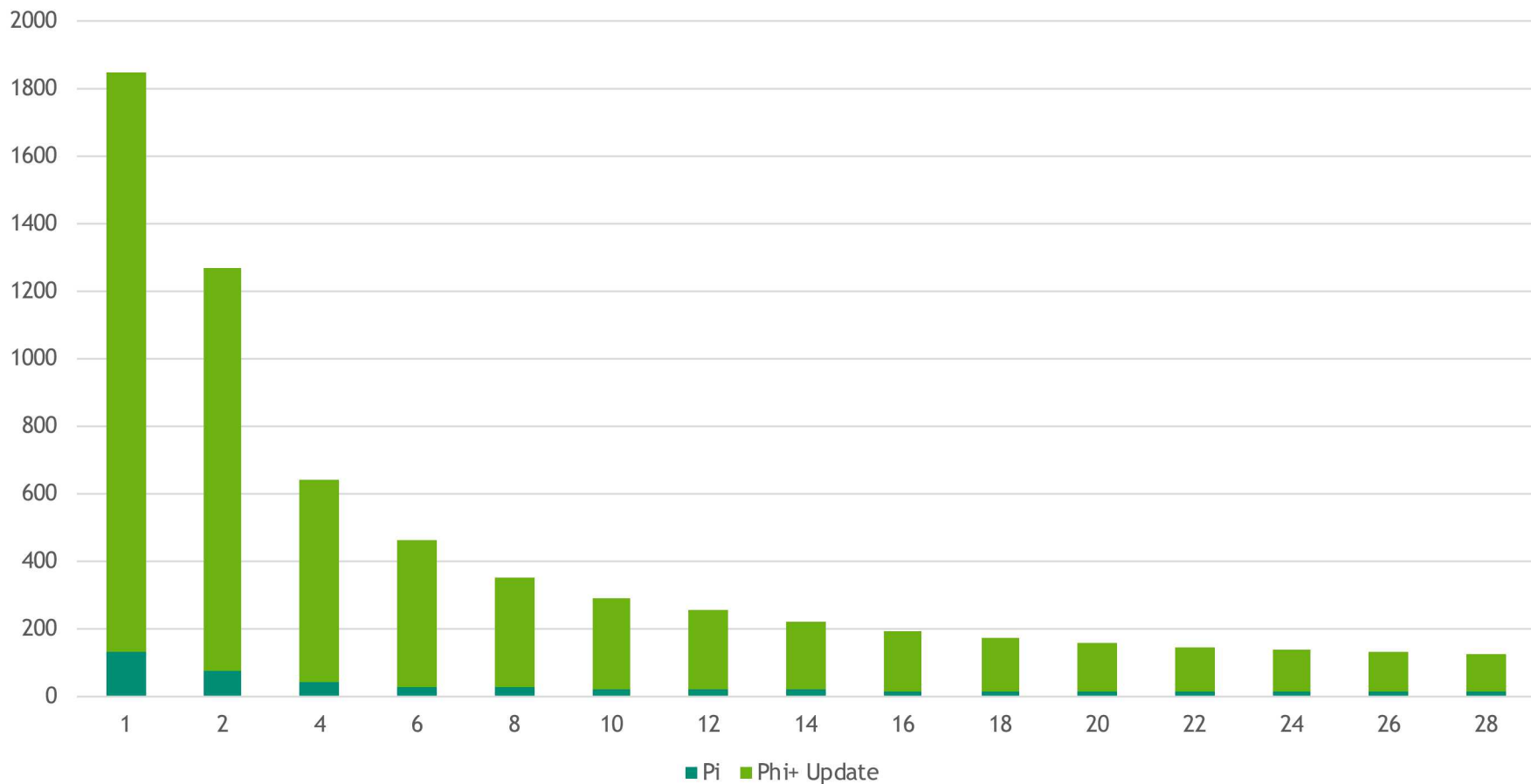
Data	Dimensions	Nonzeros	Rank (*)
LBNL	2K x 4K x 2K x 4K x 866K	1.7M	10
NELL-2	12K x 9K x 29K	77M	10
NELL-1	3M x 2M x 25M	144M	10
Delicious	500K x 17M x 3M x 1K	140M	10

(*) if not indicated.

Scalability of CPAPR-MU on CPU (Random)



CP-APR-MU method, 100 outer-iterations, (3000 x 4000 x 5000, 10M nonzero entries), $R=100$, 2 Haswell (14 core) CPUs per node, HyperThreading disabled



CP-APR-MU: Performance on GPUs (10 inner, 10 outer iterations, 10 components)

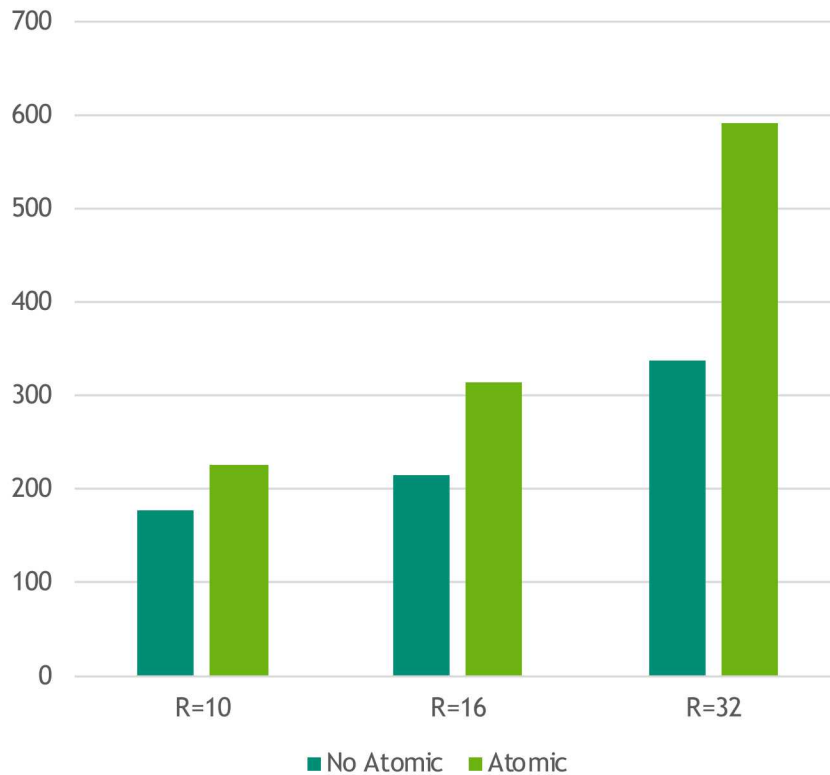


Data	Haswell CPU 1-core		2 Haswell CPUs 14-cores		2 Haswell CPUs 28-cores		Intel KNL (Cache Mode) 68-core CPU		NVIDIA P100 GPU		NVIDIA V100 GPU	
	Time(s)	Speedup	Time(s)	Speedup	Time(s)	Speedup	Time(s)	Speedup	Time(s)	Speedup	Time(s)	Speedup
Random	185	1	22	8.4	13	14.11	8.4	22.01	4.47	41.31	3.01	61.53
LBNL	39	1	19	2.05	13	3.0	33	1.18	2.99	13.04	2.09	18.66
NELL-2	1157	1	137	8.44	87	13.29	100	11.02	47.17	24.52	28.80	40.17
NELL-1	3365	1	397	16.62	258	20.9	257	10.86	OOM		OOM	
Delicious	4170	1	2183	1.91	1872	2.23	3463	1.41	OOM		OOM	

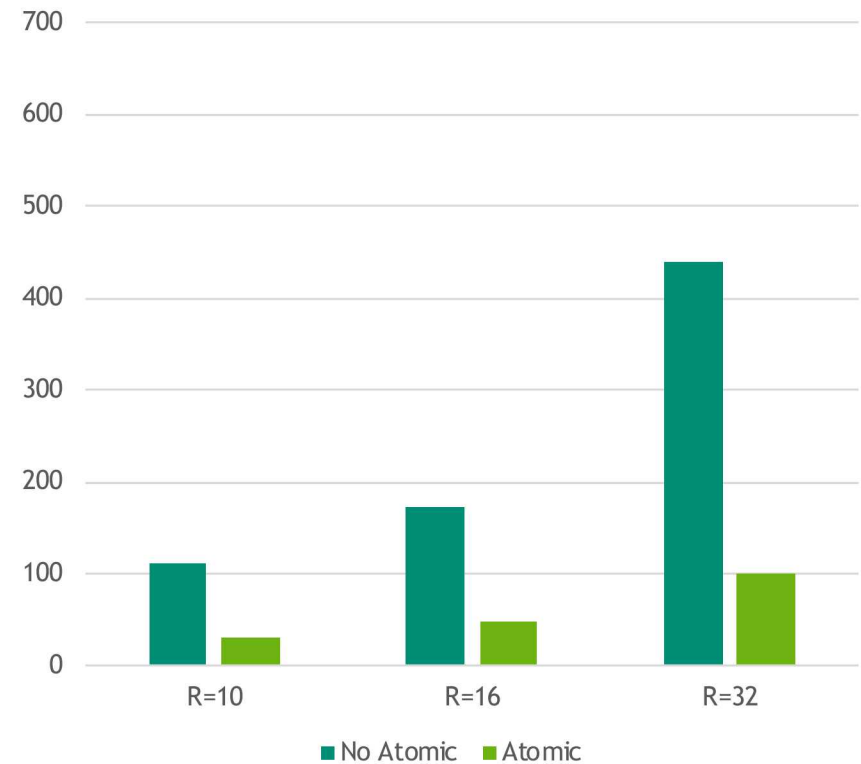
Performance Comparison: Atomic vs Non-Atomic



Performance of CP-APR-MU on
Haswell CPUs
3Kx4Kx5K Random Sparse Tensor



Performance of CP-APR-MU on
V100
3Kx4Kx5K Random Sparse Tensor

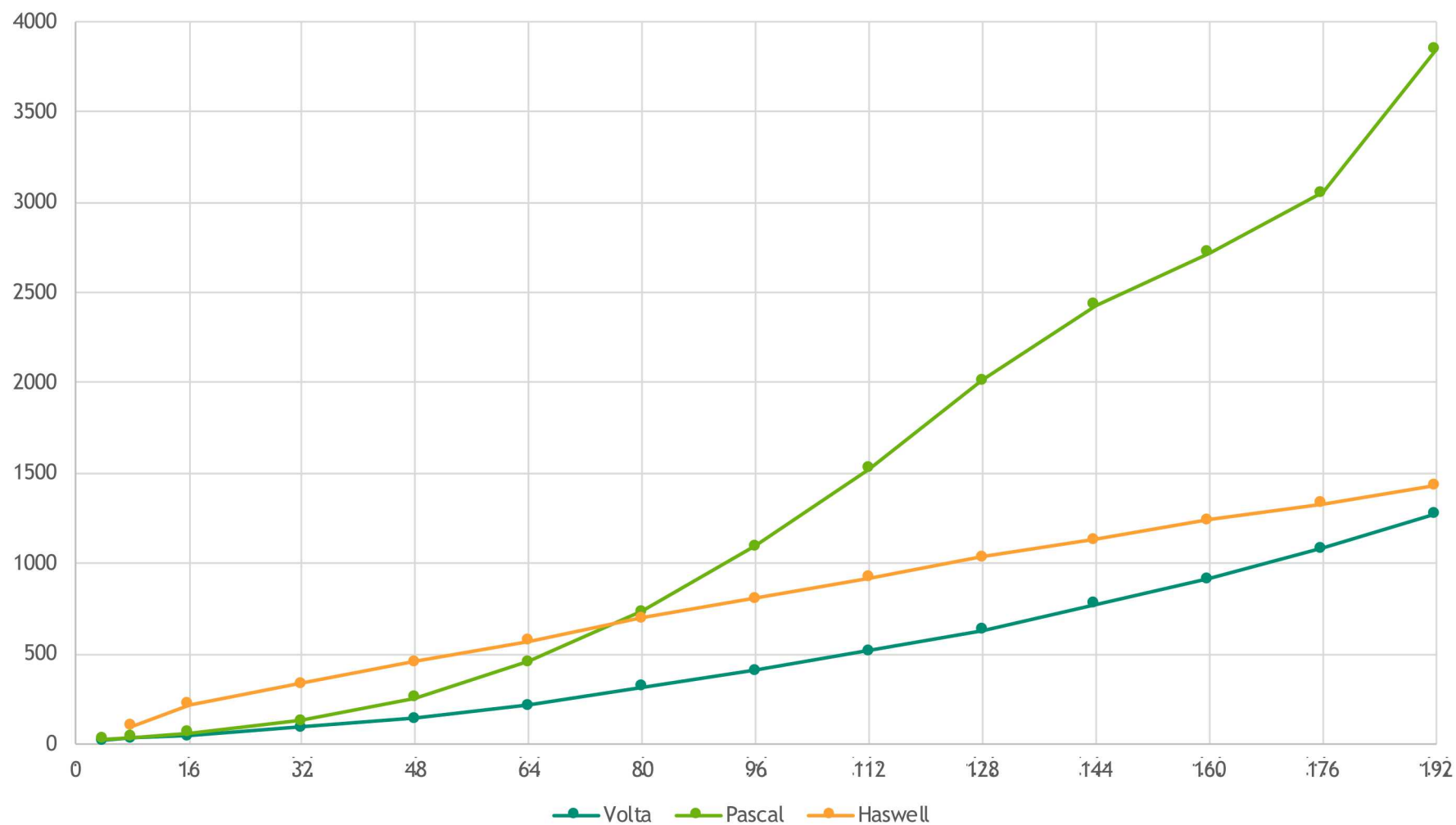


Intel CPUs: Software-based atomic operations

NVIDIA GPUs: Hardware-based atomic operations



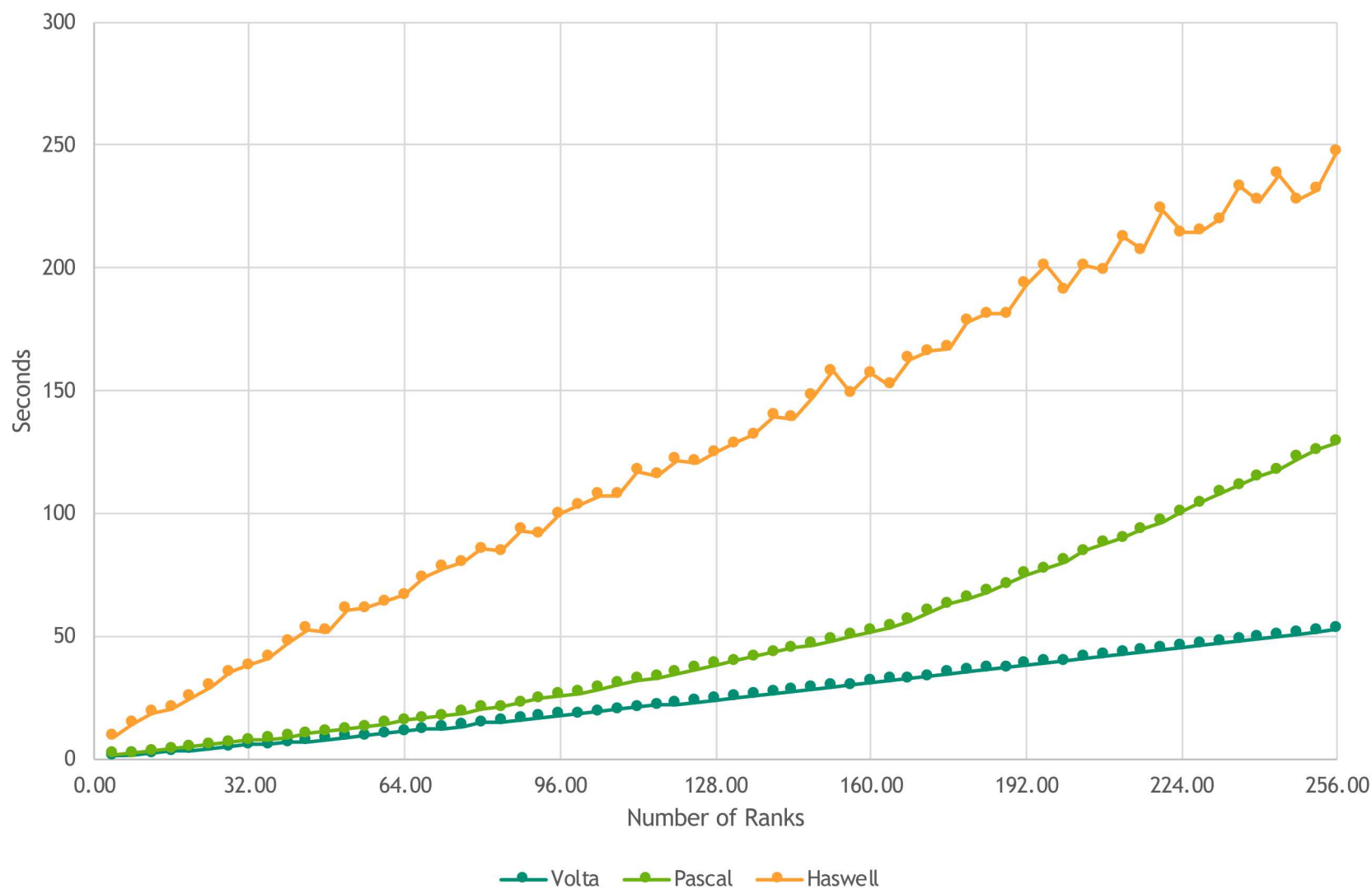
CP-APR-MU (Random tensor 3Kx4Kx5K, 100 outer iterations)



Performance of CP-APR-MU (LBNL-Network) with respect to different rank sizes



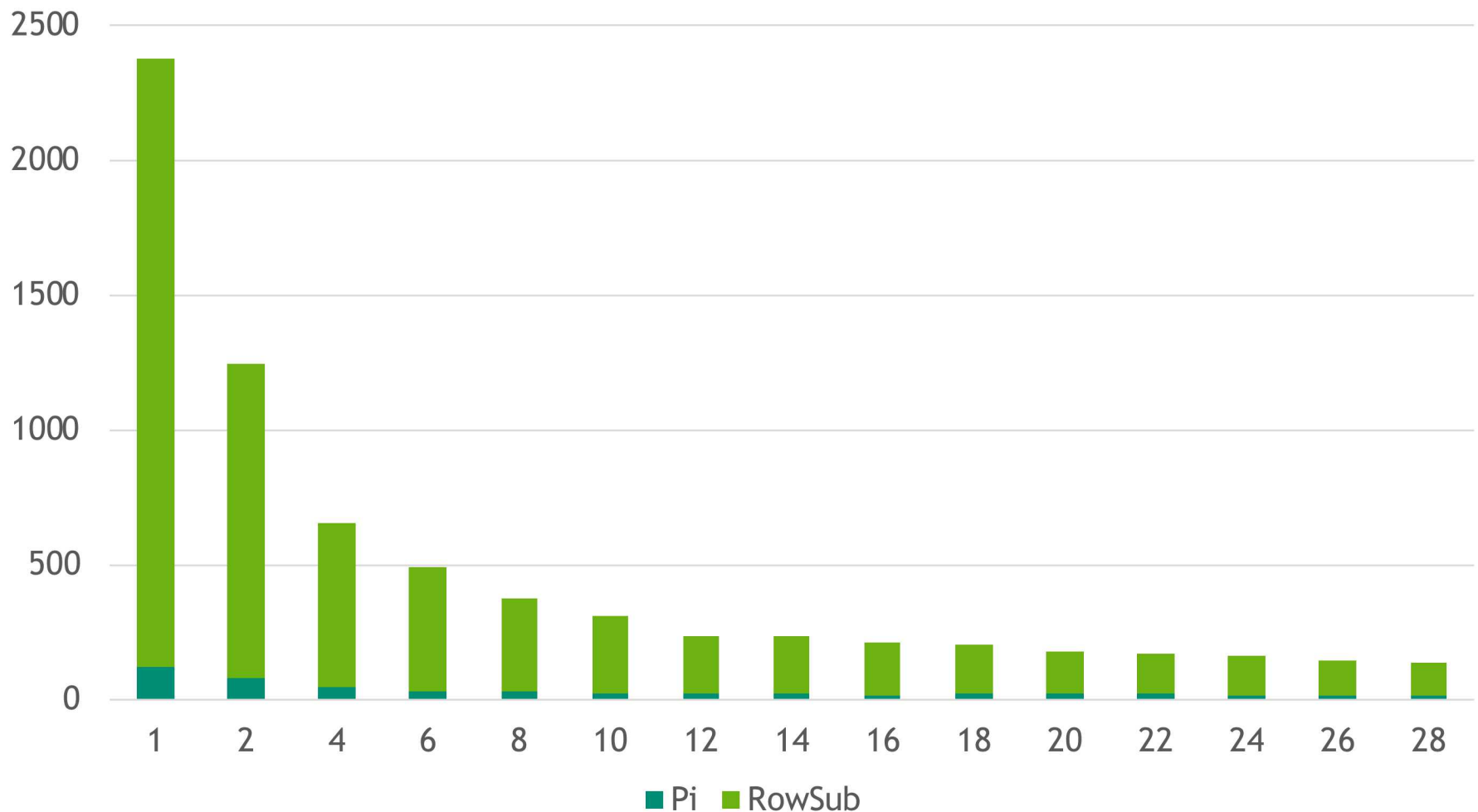
CP-APR-MU (LBNL-NETWORK, 10 outer iterations)



CPAPR-PDNR on CPU(Random)



CpAPR-PDNR method, 100 outer-iterations, 1831221 inner iterations total, (3000 x 4000 x 5000, 10M nonzero entries), R=10, 2 Haswell (14 core) CPUs per node, HyperThreading disabled





Data	Haswell CPU 1 core		2 Haswell CPUs 14 cores		2 Haswell CPUs 28 cores		KNL (Cache Mode) 68-core CPU	
	Time(s)	Speedup	Time(s)	Speedup	Time(s)	Speedup	Time(s)	Speedup
Random	238	1	23.7	10.03	14.6	16.28	17.8	13.36
LBNL	1049	1	187	2.35	191	2.30	761	1.38
NELL-2	2154	1	326	6.63	319	6.77	655	3.29
NELL-1	17212	1	4241	4.05	3974	4.33	☹	
Delicious	28053	1	3684	5.15	3138	6.05	☹	



Our implementation exhibits very good scalability with the random tensor.

- Similar mode sizes
- Regular distribution of nonzero entries
 - Some cache effects
 - Kokkos is NUMA-aware for contiguous memory access (first-touch)

Some scalability issues with the realistic tensor problems.

- Irregular nonzero distribution and disparity in mode sizes
- PDNR needs some improvement to handle row subproblems
- Preprocessing could improve the locality
 - Explicit Data partitioning (Smith and Karypis)



Development of Portable on-node Parallel CP-APR Solvers

- Data parallelism for MU method
- Mixed Data/Task parallelism for PDNR method
- Multiple Architecture Support using Kokkos
- Performance on CPU, Manycore and GPUs

Scalable Performance for random sparse tensor

Future Work

- Projected Quasi-Newton for Row-subproblems (PQNR)
- GPU and Manycore support for PDNR and PQNR
- Performance tuning to handle irregular nonzero distributions and disparity in mode sizes

Difference between P100 and V100 (300W)



GPU PERFORMANCE COMPARISON

	P100	V100	Ratio
DL Training	10 TFLOPS	120 TFLOPS	12x
DL Inferencing	21 TFLOPS	120 TFLOPS	6x
FP64/FP32	5/10 TFLOPS	7.5/15 TFLOPS	1.5x
HBM2 Bandwidth	720 GB/s	900 GB/s	1.2x
STREAM Triad Perf	557 GB/s	855 GB/s	1.5x
NVLink Bandwidth	160 GB/s	300 GB/s	1.9x
L2 Cache	4 MB	6 MB	1.5x
L1 Caches	1.3 MB	10 MB	7.7x

Courtesy: NVIDIA