Algorithms as Multilinear Tensor Equations

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Pervasive paradigms in scientific computing

What commonalities exist in simulation and data analysis applications?

- multidimensional datasets (observations, discretizations)
- higher-order relations: equations, maps, graphs, hypergraphs
- sparsity and symmetry in structure of relations
- relations lead to solution directly or as an iterative criterion
- algebraic descriptions of datasets and relations

Pervasive paradigms in scientific computing

What abstractions are needed in high performance computing?

- data abstractions reflecting native dimensionality and structure
- functions orchestrating **communication** and **synchronization**
- provably efficient building-block algorithms

Matrix computations \subset tensor computations

Tensors are convenient abstractions for multidimensional data

- one type of object for any homogeneous dataset
- enable expression of symmetries
- reveal sparsity structure of relations in multidimensional space

Tensor computations naturally extend numerical linear algebra

- = often reduce to or employ matrix algorithms
 - can leverage high performance matrix libraries
- + high-order tensors can 'act' as many matrix unfoldings
- + symmetries lower memory footprint and cost
- \bullet + tensor factorizations (CP, Tucker, tensor train, ...)

Applications of high-order tensor representations

Numerical solution to differential equations

- higher-order Taylor series expansion terms
- nonlinear terms and differential operators

Computer vision and graphics

- 2D image \otimes angle \otimes time
- compression (tensor factorizations, sparsity)

Machine learning

- sparse multi-feature discrete datasets
- reduced-order models (tensor factorizations)

Graph computations

- hypergraphs, time-dependent graphs
- clustering/partitioning/path-finding (eigenvector computations)

Divide-and-conquer algorithms representable by tensor folding

bitonic sort, FFT, scans

Applications to quantum systems

Manybody Schrödinger equation

• "curse of dimensionality" - exponential state space

Condensed matter physics

- tensor network models (e.g. DMRG), tensor per lattice site
- highly symmetric multilinear tensor representation
- $\bullet\,$ exponential state space localized $\rightarrow\,$ factorized tensor form

Quantum chemistry (electronic structure calculations)

- models of molecular structure and chemical reactions
- methods for calculating electronic correlation:
 - "Post Hartree-Fock": configuration interaction, **coupled cluster**, Møller-Plesset perturbation theory
- multi-electron states as tensors,

e.g. electron \otimes electron \otimes orbital \otimes orbital

- nonlinear equations of partially (anti)symmetric tensors
- $\bullet\,$ interactions diminish with distance $\rightarrow\,$ sparsity, low rank

- Symmetry-preserving tensor algorithms
 - contraction of order 2s symmetric tensors in $\frac{(3s)!}{(s!)^3}$ fewer multiplies
 - up to 9X speed-up for partially-symmetric contractions in coupled cluster
- Ommunication-avoiding parallel algorithms
 - novel tradeoffs: synchronization vs communication in Cholesky and stencils
 - algorithms with $p^{1/6}$ less communication on p processors for LU, QR, eigs
 - topology-aware implementations: 12X speed-up for MM, 2X for LU
- Occupies Tensor Framework (CTF)
 - first distributed-memory tensor framework supporting arbitrary contractions
 - symmetry, sparsity, multitype functions, redistributions, high-level language
- Applications to electronic structure calculations
 - codes using CTF for wavefunction methods: Aquarius, QChem, VASP, Psi4
 - $\bullet\,$ coupled cluster faster than NWChem by >10X, nearly 1 petaflop/s

Exploiting symmetry in tensors

Tensor symmetry (e.g. $A_{ij} = A_{ji}$) reduces memory and cost

- for order *d* tensor, *d*! less memory
- dot product $\sum_{i,j} A_{ij} B_{ij} = 2 \sum_{i < j} A_{ij} B_{ij} + \sum_i A_{ii} B_{ii}$
- matrix-vector multiplication¹

$$c_i = \sum_j A_{ij}b_j = \sum_j A_{ij}(b_i + b_j) - \left(\sum_j A_{ij}\right)b_i$$

rank-2 vector outer product¹

$$C_{ij} = a_i b_j + a_j b_i = (a_i + a_j)(b_i + b_j) - a_i b_i - a_j b_j$$

• squaring a symmetric matrix (or $AB + BA)^1$

$$C_{ij} = \sum_k A_{ik}A_{kj} = \sum_k (A_{ik} + A_{kj} + A_{ij})^2 - \dots$$

• for order ω contraction, $\omega!$ fewer multiplies ¹

¹S., Demmel; Technical Report, ETH Zurich, 2015.

Symmetry preserving algorithms

By exploiting symmetry, reduce multiplies (but increase adds)²

- partially symmetric contractions
 - symmetry preserving algorithm can be nested over each index group
 - reduction in multiplies \rightarrow reduction in cost, e.g. for $A_{ij}^{km} = A_{ji}^{km}$

$$c_i^{kl} = \sum_{j,m} A_{ij}^{km} b_j^{ml} = \sum_j \left(\sum_m A_{ij}^{km} (b_i^{ml} + b_j^{ml}) \right) - \sum_m \left(\sum_j A_{ij}^{km} \right) b_i^{ml}$$

- cost reductions in coupled cluster: 2X-9X for select contractions, 1.3X-2.1X for methods
- algorithms generalize to most antisymmetric tensor contractions
- for Hermitian tensors, multiplies cost 3X more than adds
 - Hermitian matrix multiplication and tridiagonal reduction (BLAS and LAPACK routines) with 25% fewer ops
- $(2/3)n^3$ bilinear rank for squaring a *nonsymmetric* matrix
- decompose symmetric contractions into smaller symmetric contractions

²S., Demmel; Technical Report, ETH Zurich, 2015.

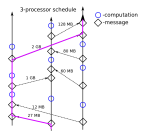
Beyond computation cost

Algorithms should minimize communication, not just computation

- data movement and synchronization cost more energy than flops
- two types of data movement:
 - vertical (intranode memory-cache)
 - horizontal (internode network transfers)
- parallel algorithm design involves tradeoffs: computation vs communication vs synchronization
- lower bounds and parameterized algorithms provide optimal solutions within a well-defined tuning space

Given a schedule of work and communication tasks on p processors, consider the following costs, accumulated along chains of tasks (as in $\alpha - \beta$, BSP, and LogGP models),

- F computation cost
- Q vertical communication cost
- W horizontal communication cost
- S synchronization cost



Communication lower bounds: previous work

Multiplication of $n \times n$ matrices

• horizontal communication lower bound³

$$W_{\rm MM} = \Omega\left(rac{n^2}{p^{2/3}}
ight)$$

• memory-dependent horizontal communication lower bound⁴

$$W_{\rm MM} = \Omega\left(rac{n^3}{p\sqrt{M}}
ight)$$

• with $M = cn^2/p$ memory, hope to obtain communication cost

$$W = O(n^2/\sqrt{cp})$$

• libraries like ScaLAPACK, Elemental optimal only for c = 1

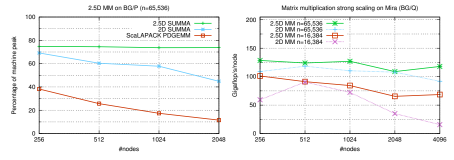
³Aggarwal, Chandra, Snir, TCS, 1990

⁺Irony, Toledo, Tiskin, JPDC, 2004

Communication-efficient matrix multiplication

Communication-avoiding algorithms for matrix multiplication have been studied extensively $^{5}\,$

They continue to be attractive on modern architectures⁶



12X speed-up, 95% reduction in comm. for n = 8K on 16K nodes of BG/P

⁵Berntsen, Par. Comp., 1989; Agarwal, Chandra, Snir, TCS, 1990; Agarwal, Balle, Gustavson, Joshi, Palkar, IBM, 1995; McColl, Tiskin, Algorithmica, 1999; ...

⁶S., Bhatele, Demmel, SC, 2011

Synchronization cost lower bounds

Unlike matrix multiplication, many algorithms in numerical linear algebra have polynomial depth (contain a long dependency path)

• matrix multiplication synchronization cost bound⁷

$$S_{\rm MM} = \Theta\left(rac{n^3}{pM^{3/2}}
ight)$$

- algorithms for Cholesky, LU, QR, SVD do not attain this bound
- low granularity computation increases synchronization cost

⁷Ballard, Demmel, Holtz, Schwartz, SIAM JMAA, 2011

Tradeoffs in the diamond DAG

Computation vs synchronization tradeoff for the $n \times n$ diamond DAG,⁸

$$F \cdot S = \Omega(n^2)$$



We generalize this idea⁹

- additionally consider horizontal communication
- allow arbitrary (polynomial or exponential) interval expansion

⁸Papadimitriou, Ullman, SIAM JC, 1987

⁹S., Carson, Knight, Demmel, SPAA 2014 (extended version, JPDC 2016)

Tradeoffs involving synchronization

We apply tradeoff lower bounds to dense linear algebra algorithms, represented via dependency hypergraphs:^{*a*}

For triangular solve with an $n \times n$ matrix,

$$F_{\text{TRSV}} \cdot S_{\text{TRSV}} = \Omega\left(n^2\right)$$

For Cholesky of an $n \times n$ matrix,

$$F_{CHOL} \cdot S_{CHOL}^2 = \Omega(n^3)$$
 $W_{CHOL} \cdot S_{CHOL} = \Omega(n^2)$

Proof employs classical Loomis-Whitney inequality:

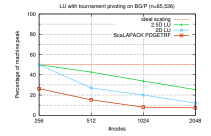
for any $R\subset\mathbb{N}\times\mathbb{N}\times\mathbb{N}$, three projections of R onto $\mathbb{N}\times\mathbb{N}$ have total size at least $|R|^{2/3}$

^aS., Carson, Knight, Demmel, SPAA 2014 (extended version, JPDC 2016)

Communication-efficient LU factorization

For any $c \in [1, p^{1/3}]$, use cn^2/p memory per processor and obtain

$$W_{\rm LU} = O(n^2/\sqrt{cp}), \qquad S_{\rm LU} = O(\sqrt{cp})$$



- LU with pairwise pivoting¹⁰ extended to tournament pivoting¹¹
- first implementation of a communication-optimal LU algorithm¹¹

¹⁰Tiskin, FGCS, 2007

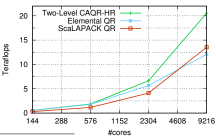
¹¹S., Demmel, Euro-Par, 2011

Communication-efficient QR factorization

- $W_{\rm QR} = O(n^2/\sqrt{cp}), S_{\rm QR} = O(\sqrt{cp})$ using Givens rotations¹²
- Householder form can be reconstructed quickly from TSQR¹³

$$Q = I - YTY^T \quad \rightarrow \quad \mathsf{LU}(I - Q) \rightarrow (Y, TY^T)$$

- enables communication-optimal Householder QR¹⁴
- Householder aggregation yields performance improvements



¹²Tiskin, FGCS, 2007

¹³Ballard, Demmel, Grigori, Jacquelin, Nguyen, S., IPDPS, 2014

¹⁴S., UCB, 2014

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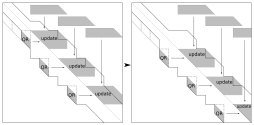
QR weak scaling on Cray XE6 (n=15K to n=131K)

Communication-efficient eigenvalue computation

For the dense symmetric matrix eigenvalue problem^a

$$W_{\mathsf{SE}} = O(n^2/\sqrt{cp}), S_{\mathsf{QR}} = O(\sqrt{cp}\log^2 p)$$

- above costs obtained by left-looking algorithm with Householder aggregation, however, with increased vertical communication
- successive band reduction minimizes both communication costs



^aS., UCB, 2014. S., Hoefler, Demmel, in preparation

Synchronization tradeoffs in stencils

Our lower bound analysis extends to sparse iterative methods:¹⁵ For computing *s* applications of a $(2m + 1)^d$ -point stencil

$$F_{\mathsf{St}} \cdot S^{d}_{\mathsf{St}} = \Omega\left(m^{2d} \cdot s^{d+1}\right) \qquad W_{\mathsf{St}} \cdot S^{d-1}_{\mathsf{St}} = \Omega\left(m^{d} \cdot s^{d}\right)$$

- proof requires generalization of Loomis-Whitney inequality to order d set and order d-1 projections
- time-blocking lowers synchronization and vertical communication costs, but raises horizontal communication
- we suggest alternative approach that minimizes vertical and horizontal communication, but not synchronization

¹⁵S., Carson, Knight, Demmel, SPAA 2014 (extended version, JPDC 2016)

Beyond the Loomis-Whitney inequalities

Loomis-Whitney inequalities are not sufficient for all computations

- symmetry preserving tensor contraction algorithms have arbitrary order projections from order d set
- bilinear algorithms¹⁶ provide a more general framework
- a bilinear algorithm is defined by matrices $F^{(A)}, F^{(B)}, F^{(C)}$,

$$c = F^{(C)}[(F^{(A)\mathsf{T}}a) \circ (F^{(B)\mathsf{T}}b)]$$

where \circ is the Hadamard (pointwise) product

$$\begin{bmatrix} \mathbf{c} \end{bmatrix} = \begin{bmatrix} \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \\ \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \end{bmatrix}} \begin{bmatrix} \left(\begin{bmatrix} \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \\ \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \\ \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \\ \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \\ \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \end{bmatrix}}^{\mathsf{T}} \begin{bmatrix} \mathbf{a} \\ \mathbf{a} \end{bmatrix} \right) \circ \begin{pmatrix} \begin{bmatrix} \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \\ \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} & \mathbf{x} \end{bmatrix}}^{\mathsf{T}} \begin{bmatrix} \mathbf{b} \\ \mathbf{b} \end{bmatrix} \end{bmatrix} \end{bmatrix}$$

communication lower bounds derived based on matrix rank¹⁷

¹⁶Pan, Springer, 1984

¹⁷S., Hoefler, Demmel, in preparation

Communication cost of symmetry preserving algorithms

For contraction of order s + v tensor with order v + t tensor¹⁸

- symmetry preserving algorithm requires $\frac{(s+v+t)!}{s!v!t!}$ fewer multiplies
- matrix-vector-like algorithms $(\min(s, v, t) = 0)$
 - vertical communication dominated by largest tensor
 - horizontal communication asymptotically greater if only unique elements are stored and $s \neq v \neq t$
- matrix-matrix-like algorithms $(\min(s, v, t) > 0)$
 - vertical and horizontal communication costs asymptotically greater for symmetry preserving algorithm when $s \neq v \neq t$

¹⁸S., Hoefler, Demmel; Technical Report, ETH Zurich, 2015.

Cyclops Tensor Framework¹⁹

- contraction/summation/functions of tensors
- distributed symmetric-packed/sparse storage via cyclic layout
- parallelization via MPI+OpenMP(+CUDA)

¹⁹S., Hammond, Demmel, UCB, 2011. S., Matthews, Hammond, Demmel, IPDPS, 2013

Cyclops Tensor Framework

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Jacobi iteration (solves Ax = b iteratively) example code snippet

```
Vector<> Jacobi(Matrix<> A, Vector<> b, int n){
    ... // split A = R + diag(1./d)
    do {
        x["i"] = d["i"]*(b["i"]-R["ij"]*x["j"]);
        r["i"] = b["i"]-A["ij"]*x["j"]; // compute residual
    } while (r.norm2() > 1.E-6); // check for convergence
    return x;
}
```

Cyclops Tensor Framework

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Jacobi iteration (solves Ax = b iteratively) example code snippet

```
Vector<> Jacobi(Matrix<> A, Vector<> b, int n){
Matrix<> R(A);
R["ii"] = 0.0;
Vector<> x(n), d(n), r(n);
Function<> inv([](double & d){ return 1./d; });
d["i"] = inv(A["ii"]); // set d to inverse of diagonal of A
do {
    x["i"] = d["i"]*(b["i"]-R["ij"]*x["j"]);
    r["i"] = b["i"]-A["ij"]*x["j"]; // compute residual
} while (r.norm2() > 1.E-6); // check for convergence
return x;
}
```

Cyclops Tensor Framework

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Møller-Plesset perturbation theory (MP3) code snippet

```
Z["abij"] += Fab["af"]*T["fbij"];
Z["abij"] -= Fij["ni"]*T["abnj"];
Z["abij"] += 0.5*Vabcd["abef"]*T["efij"];
Z["abij"] += 0.5*Vijkl["mnij"]*T["abmn"];
Z["abij"] -= Vaibj["amei"]*T["ebmj"];
```

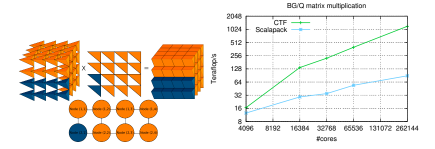
```
Betweenness centrality code snippet, for k of n nodes
void btwn_central(Matrix<int> A, Matrix<path> P, int n, int k){
  Monoid < path > mon(...,
                     [](path a, path b){
                       if (a.w<b.w) return a;</pre>
                       else if (b.w<a.w) return b;</pre>
                       else return path(a.w, a.m+b.m);
                    , \ldots);
  Matrix<path> Q(n,k,mon); // shortest path matrix
  Q["ij"] = P["ij"];
  Function<int,path> append([](int w, path p){
                         return path(w+p.w, p.m);
                       }; );
  for (int i=0; i<n; i++)</pre>
    Q["ij"] = append(A["ik"],Q["kj"]);
  . . .
```

}

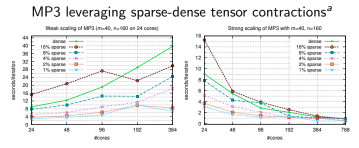
Performance of CTF for dense computations

CTF is highly tuned for massively-parallel machines

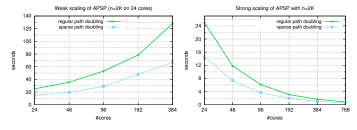
- virtualized multidimensional processor grids
- topology-aware mapping and collective communication
- performance-model-driven decomposition done at runtime
- optimized redistribution kernels for tensor transposition



Performance of CTF for sparse computations



All-pairs shortest-paths based on path doubling with sparsification^a



^aS., Hoefler, Demmel, arXiv, 2015

Coupled cluster methods

Coupled cluster provides a systematically improvable approximation to the manybody time-independent Schrödinger equation $H|\Psi\rangle=E|\Psi\rangle$

- the Hamiltonian has one- and two- electron components H = F + V
- Hartree-Fock (SCF) computes mean-field Hamiltonian: F, V
- Coupled-cluster methods (CCSD, CCSDT, CCSDTQ) consider transitions of (doubles, triples, and quadruples) of electrons to unoccupied orbitals, encoded by tensor operator T = T₁ + T₂ + T₃ + T₄
- they use an exponential ansatz for the wavefunction, $\Psi = e^T \phi$ where ϕ is a Slater determinant
- expanding 0 = $\langle \phi' | H | \Psi \rangle$ yields nonlinear equations for $\{T_i\}$ in F, V

$$0 = V_{ij}^{ab} + \mathcal{P}(a,b) \sum_{e} T_{ij}^{ae} F_{e}^{b} - \frac{1}{2} \mathcal{P}(i,j) \sum_{mnef} T_{im}^{ab} V_{ef}^{mn} T_{jn}^{ef} + \dots$$

where $\ensuremath{\mathcal{P}}$ is an antisymmetrization operator

CCSD using CTF

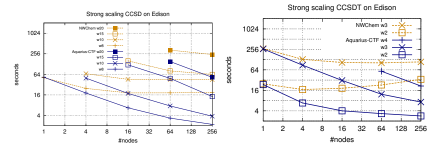
Extracted from Aquarius (Devin Matthews' code, https://github.com/devinamatthews/aquarius)

```
FMI["mi"] += 0.5*WMNEF["mnef"]*T2["efin"];
WMNIJ["mnij"] += 0.5*WMNEF["mnef"]*T2["efij"];
FAE["ae"] -= 0.5*WMNEF["mnef"]*T2["afmn"];
WAMEI["amei"] -= 0.5*WMNEF["mnef"]*T2["afmn"];
Z2["abij"] = WMNEF["ijab"];
Z2["abij"] += FAE["af"]*T2["fbij"];
Z2["abij"] -= FMI["ni"]*T2["abnj"];
Z2["abij"] += 0.5*WABEF["abef"]*T2["efij"];
Z2["abij"] += 0.5*WMNIJ["mnij"]*T2["abmn"];
Z2["abij"] -= WAMEI["amei"]*T2["ebmj"];
```

CTF-based CCSD codes exist in Aquarius, QChem, VASP, and Psi4

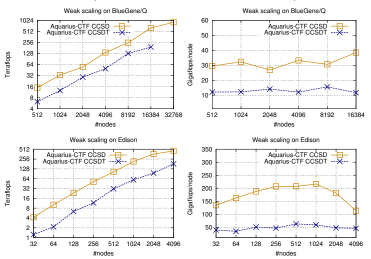
NWChem is the most commonly-used distributed-memory quantum chemistry method suite

- provides CCSD and CCSDT
- derives equations via Tensor Contraction Engine (TCE)
- generates contractions as blocked loops leveraging Global Arrays



Coupled cluster on IBM BlueGene/Q and Cray XC30

CCSD up to 55 (50) water molecules with cc-pVDZ CCSDT up to 10 water molecules with cc-pVDZ^a

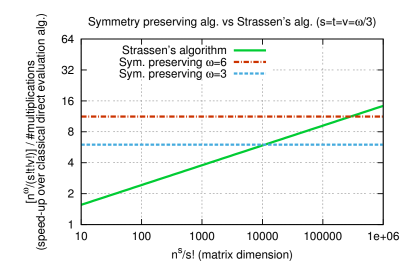


Novel results described in this talk:

- symmetry preserving algorithms
 - reduce number of multiplications in symmetric contractions by $\omega!$
 - $\bullet\,$ reduce cost of basic Hermitian matrix operations by 25%
 - reduce cost of some contractions in coupled cluster by 2X in CCSD (1.3X overall), 4X in CCSDT (2.1X overall), 9X in CCSDTQ
- communication and synchronization lower bounds
 - tradeoffs: synchronization vs computation or communication in TRSV, Cholesky, and stencils
 - rank-based lower bounds to analyze symmetric contractions
- communication avoiding dense matrix factorizations
 - new algorithms and implementations with up to $p^{1/6}$ less communication for LU, QR, symmetric eigenvalue problem
 - speed-ups of up to 2X for LU and QR over vendor-optimized libraries
- Cyclops Tensor Framework
 - first fully robust distributed-memory tensor contraction library
 - supports symmetry, sparsity, general algebraic structures
 - $\bullet\,$ coupled cluster performance more than 10X faster than state-of-the-art, reaching 1 petaflop/s performance

- symmetry in tensor computations
 - $\bullet~$ cost improvements \rightarrow fast library implementations \rightarrow application speed-ups
 - study symmetries in tensor equations and factorizations
 - consider other symmetries and relation to fast matrix multiplication
- communication-avoiding algorithms
 - existing fast implementations already used by applications (e.g. QBox)
 - find efficient methods of searching larger tuning spaces
 - algorithms for computing eigenvectors, SVD, tensor factorizations
 - study communication-efficiency of randomized algorithms
- Cyclops Tensor Framework
 - already widely-adapted in quantum chemistry, many requests for features
 - $\bullet\,$ study algorithms for tensor expressions \rightarrow factorization, scheduling, ...
 - engage new application domains (via sparsity and algebraic structures)
 - tensor networks for condensed matter-physics, particle methods
 - graph algorithms, discrete data analysis
 - graphics, computer vision, machine learning

Backup slides



Nesting of bilinear algorithms

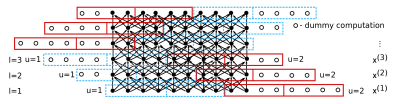
Given two bilinear algorithms:

$$\begin{split} &\Lambda_1 = &(\textbf{F}_1^{(\textbf{A})}, \textbf{F}_1^{(\textbf{B})}, \textbf{F}_1^{(\textbf{C})}) \\ &\Lambda_2 = &(\textbf{F}_2^{(\textbf{A})}, \textbf{F}_2^{(\textbf{B})}, \textbf{F}_2^{(\textbf{C})}) \end{split}$$

We can nest them by computing their tensor product

$$\begin{split} &\Lambda_1\otimes\Lambda_2\coloneqq (\textbf{F_1^{(A)}}\otimes\textbf{F_2^{(A)}},\textbf{F_1^{(B)}}\otimes\textbf{F_2^{(B)}},\textbf{F_1^{(C)}}\otimes\textbf{F_2^{(C)}})\\ &\operatorname{rank}(\Lambda_1\otimes\Lambda_2)=\operatorname{rank}(\Lambda_1)\cdot\operatorname{rank}(\Lambda_2) \end{split}$$

Block-cyclic algorithm for s-step methods

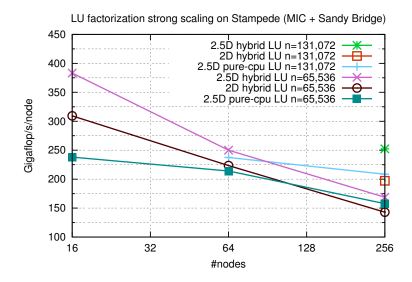


For s-steps of a $(2m+1)^d$ -point stencil with block-size of $H^{1/d}/m$,

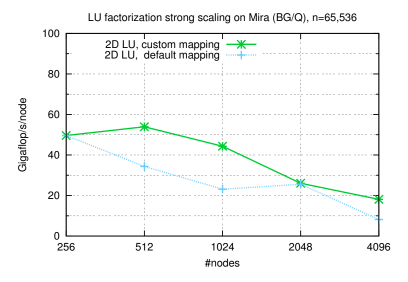
$$W_{\mathrm{Kr}} = O\left(rac{msn^d}{H^{1/d}p}
ight) \quad S_{\mathrm{Kr}} = O(sn^d/(pH)) \quad Q_{\mathrm{Kr}} = O\left(rac{msn^d}{H^{1/d}p}
ight)$$

which are good when $H = \Theta(n^d/p)$, so the algorithm is useful when the cache size is a bit smaller than n^d/p

2.5D LU on MIC

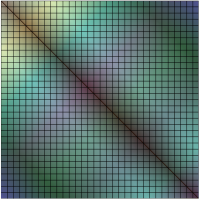


Topology-aware mapping on BG/Q

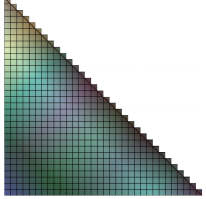


Symmetric matrix representation

Symmetric matrix



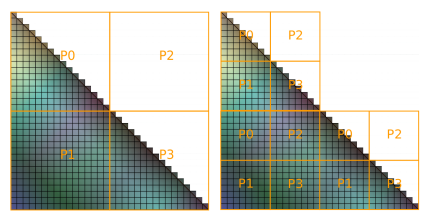
Unique part of symmetric matrix



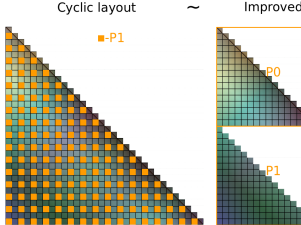
Blocked distributions of a symmetric matrix

Naive blocked layout

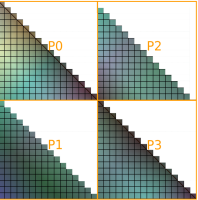
Block-cyclic layout



Cyclic distribution of a symmetric matrix



Improved blocked layout



Our CCSD factorization

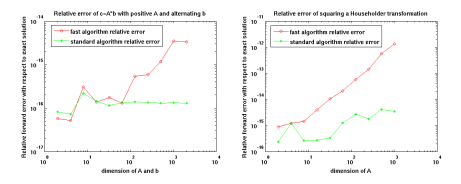
Credit to John F. Stanton and Jurgen Gauss

$$\begin{split} \tau_{ij}^{ab} &= t_{ij}^{ab} + \frac{1}{2} P_b^a P_j^i t_i^a t_j^b, \\ \tilde{F}_e^m &= f_e^m + \sum_{fn} v_{ef}^{mn} t_n^f, \\ \tilde{F}_e^a &= (1 - \delta_{ae}) f_e^a - \sum_m \tilde{F}_e^m t_m^a - \frac{1}{2} \sum_{mnf} v_{ef}^{mn} t_{mn}^{af} + \sum_{fn} v_{ef}^{an} t_n^f, \\ \tilde{F}_i^m &= (1 - \delta_{mi}) f_i^m + \sum_e \tilde{F}_e^m t_i^e + \frac{1}{2} \sum_{nef} v_{ef}^{mn} t_{in}^{ef} + \sum_{fn} v_{if}^{mn} t_n^f, \end{split}$$

Our CCSD factorization

$$\begin{split} \tilde{W}_{ei}^{mn} &= v_{ei}^{mn} + \sum_{f} v_{ef}^{mn} t_{i}^{f}, \\ \tilde{W}_{ij}^{mn} &= v_{ij}^{mn} + P_{j}^{i} \sum_{e} v_{ie}^{mn} t_{i}^{e} + \frac{1}{2} \sum_{ef} v_{ef}^{mn} \tau_{ij}^{ef}, \\ \tilde{W}_{ie}^{am} &= v_{ie}^{am} - \sum_{n} \tilde{W}_{ei}^{mn} t_{n}^{a} + \sum_{f} v_{ef}^{ma} t_{i}^{f} + \frac{1}{2} \sum_{nf} v_{ef}^{mn} t_{in}^{af}, \\ \tilde{W}_{ij}^{am} &= v_{ij}^{am} + P_{j}^{i} \sum_{e} v_{ie}^{am} t_{j}^{e} + \frac{1}{2} \sum_{ef} v_{ef}^{am} \tau_{ij}^{ef}, \\ z_{i}^{a} &= f_{i}^{a} - \sum_{m} \tilde{F}_{i}^{m} t_{m}^{a} + \sum_{e} f_{e}^{a} t_{i}^{e} + \sum_{em} v_{ei}^{ma} t_{m}^{e} + \sum_{em} v_{im}^{ae} \tilde{F}_{e}^{m} + \frac{1}{2} \sum_{efm} v_{ef}^{am} \tau_{im}^{ef} \\ &- \frac{1}{2} \sum_{emn} \tilde{W}_{ei}^{mn} t_{mn}^{ea}, \\ z_{ij}^{ab} &= v_{ij}^{ab} + P_{j}^{i} \sum_{e} v_{ie}^{ab} t_{j}^{e} + P_{b}^{a} P_{j}^{i} \sum_{me} \tilde{W}_{ie}^{am} t_{mj}^{eb} - P_{b}^{a} \sum_{m} \tilde{W}_{ij}^{am} \tau_{mn}^{ab}, \\ &+ P_{b}^{a} \sum_{e} \tilde{F}_{e}^{a} t_{ij}^{eb} - P_{j}^{i} \sum_{m} \tilde{F}_{m}^{m} t_{mj}^{ab} + \frac{1}{2} \sum_{ef} v_{ef}^{ab} \tau_{ij}^{ef} + \frac{1}{2} \sum_{mn} \tilde{W}_{ij}^{mn} \tau_{mn}^{ab}, \end{split}$$

Stability of symmetry preserving algorithms



Performance breakdown on BG/Q

Performance data for a CCSD iteration with 200 electrons and 1000 orbitals on 4096 nodes of Mira 4 processes per node, 16 threads per process Total time: 18 mins *v*-orbitals, *o*-electrons

kernel	% of time	complexity	architectural bounds
DGEMM	45%	$O(v^4 o^2 / p)$	flops/mem bandwidth
broadcasts	20%	$O(v^4 o^2 / p \sqrt{M})$	multicast bandwidth
prefix sum	10%	<i>O</i> (<i>p</i>)	allreduce bandwidth
data packing	7%	$O(v^2o^2/p)$	integer ops
all-to-all-v	7%	$O(v^2o^2/p)$	bisection bandwidth
tensor folding	4%	$O(v^2 o^2 / p)$	memory bandwidth

Tiskin's path doubling algorithm

Tiskin gives a way to do path-doubling in $F = O(n^3/p)$ operations. We can partition each \mathbf{A}^k by path size (number of edges)

$$\mathbf{A}^k = \mathbf{I} \oplus \mathbf{A}^k(1) \oplus \mathbf{A}^k(2) \oplus \ldots \oplus \mathbf{A}^k(k)$$

where each $\mathbf{A}^{k}(I)$ contains the shortest paths of up to $k \ge I$ edges, which have exactly I edges. We can see that

$$\mathbf{A}^{\prime}(l) \leq \mathbf{A}^{\prime+1}(l) \leq \ldots \leq \mathbf{A}^{n}(l) = \mathbf{A}^{*}(l),$$

in particular $\mathbf{A}^*(I)$ corresponds to a sparse subset of $\mathbf{A}^I(I)$. The algorithm works by picking $I \in [k/2, k]$ and computing

$$(\mathbf{I} \oplus \mathbf{A})^{3k/2} \leq (\mathbf{I} \oplus \mathbf{A}^k(l)) \otimes \mathbf{A}^k,$$

which finds all paths of size up to 3k/2 by taking all paths of size exactly $l \ge k/2$ followed by all paths of size up to k.