Iterative sparse matrix partitioning

Supervisor: Prof. dr. Rob H. Bisseling

Davide Taviani

October 7th, 2013



Universiteit Utrecht

Parallel sparse matrix-vector multiplication

At the core of many iterative solvers (e.g. conjugate gradient method) lies a simple operation: sparse matrix-vector multiplication.

Given:

- ▶ $m \times n$ sparse matrix A (N nonzeros, $N \ll mn$)
- $\rightarrow n \times 1 \text{ vector } \vec{v}$

we want to compute

$$\vec{u} = A\vec{v}$$



Parallel sparse matrix-vector multiplication

Usually A is fairly large and a lot of computations are required:

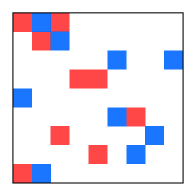
- \triangleright $\mathcal{O}(mn)$ following the definition of matrix-vector multiplication;
- \triangleright $\mathcal{O}(N)$ only considering the nonzero elements.

We split the computations among p processors to improve speed.

We make a **partition** of the set of the nonzeros of A, obtaining p disjoint sets A_0, \ldots, A_{p-1} .

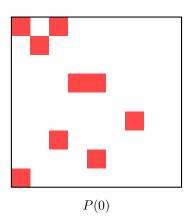
Furthermore, also the input vector \vec{v} and the final output \vec{u} can be divided among those p processors (their distribution might not necessarily be the same).

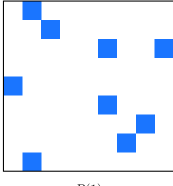
Example of a partition of a 9×9 matrix with 18 nonzeros, with p=2.





Local view of the matrix for every processor:





P(1)

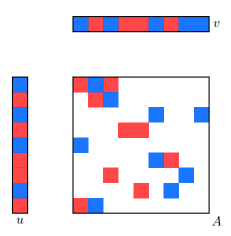


Parallel sparse matrix-vector multiplication is made (essentially) by 3 phases:

- I) fan-out
- II) local multiplication
- III) fan-in

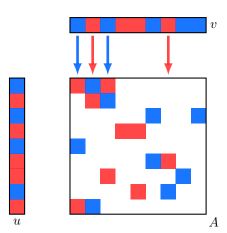


A is partitioned along with u and v





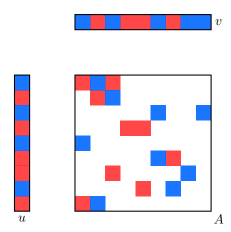
Fan-out: each processor receives the required elements of \vec{v} from the others (according to its distribution)



Universiteit Utrecht



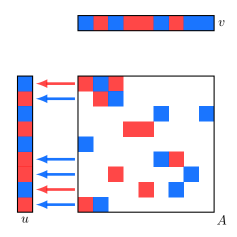
Local multiplication: where the actual computation is performed







Fan-in: where each processor sends his contributions to the other processors according to the distribution of \vec{u}







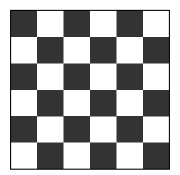
To optimize this process:

- ▶ I and III involve communication: it has to be **minimized**
- ► II is a computation step: we need **balance** in the size of the partitions

Optimization problem: partition the nonzeros such that the balance constraint is satisfied and the communication volume is minimized.

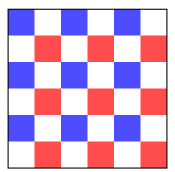


As a last example, a 6×6 "checkerboard" matrix:

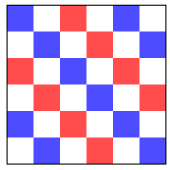




Two different partitionings result in extremely different communication volumes.



(a) Rows and columns are not split, therefore there is no need for communication



(b) Every row and column is split and causes communication during fan-in and fan-out.

Universiteit Utrecht

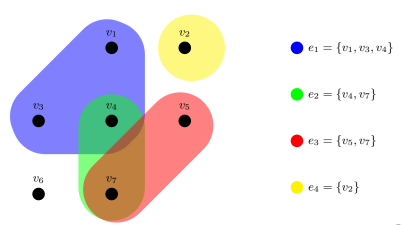


Exact modeling of the matrix partitioning problem through hypergraph partitioning.

- ▶ A partition of a hypergraph is simply the partition of the set of vertices V into V_0, \ldots, V_{p-1} .
- A hyperedge $e = \{v_1, \dots, v_k\}$ is **cut** if two of its vertices belong to different sets of the partition.



Hypergraph: a graph in which a hyperedge can connect more than two vertices (i.e. a subset of the vertex set V)





There are several models to translate the matrix partitioning to hypergraph partitioning:

▶ 1-dimensional

- \diamond **row-net**: each column of A is a vertex in the hypergraph, each row a hyperedge. If $a_{ij} \neq 0$, then column A_i is placed in the hyperedge j.
- column-net: identical to the previous one, with the roles of columns and rows exchanged

As hypergraph partitioning consists in assignment of the vertices, columns/rows are uncut. Advantage of eliminating completely one source of communication, but being 1-dimensional is often a too strong restriction.



▶ 2-dimensional

 \diamond fine grain: nonzeros of A are vertices, rows and columns are hyperedges. The nonzero a_{ij} is placed in the hyperedges i and j

A lot of freedom in partitioning (each nonzero can be assigned individually), but the size of the hypergraph (N vertices) is often too large.

 medium grain: middle ground between 1-dimensional models and fine-grain

Good compromise between the size of the hypergraph and freedom during the partitioning.

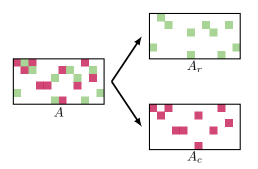


(Daniel M. Pelt and Rob Bisseling, 2013, to appear)





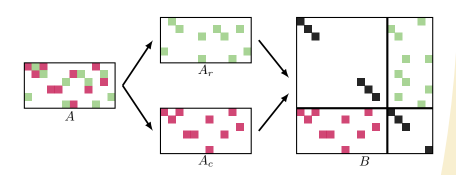
(Daniel M. Pelt and Rob Bisseling, 2013, to appear)



▶ Initial split of A into A_c and A_r



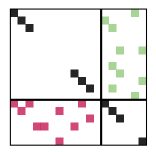
(Daniel M. Pelt and Rob Bisseling, 2013, to appear)



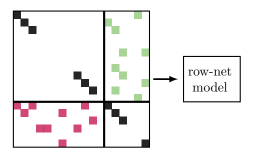
- ▶ Initial split of A into A_c and A_r
- ▶ Construction of the $(m+n) \times (m+n)$ matrix B (with dummy diagonal elements)





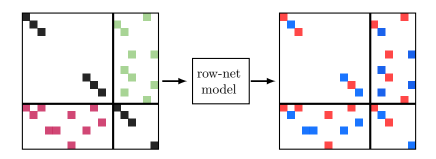






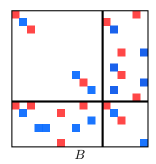
ightharpoonup Partitioning of B with the row-net model (columns are kept together)



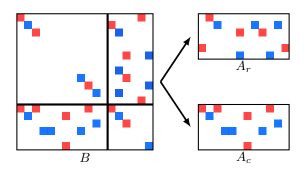


ightharpoonup Partitioning of B with the row-net model (columns are kept together)





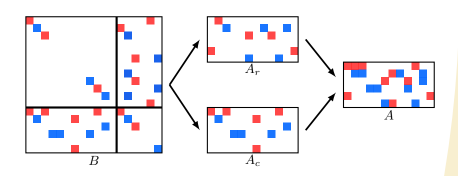




ightharpoonup Retrieval of A_r and A_c with the new partitioning



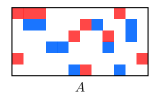




- ightharpoonup Retrieval of A_r and A_c with the new partitioning
- ightharpoonup Reassembling of A







Clusters of nonzeros are grouped together:

- in A_r we kept together elements of the same row;
- ightharpoonup in A_c elements of the same column.



Research directions

Two research directions:

- ightharpoonup Improving the initial partitioning of A
- ▶ Development of a fully iterative scheme: lowering the communication value by using information on the previous partitioning

These directions can be combined: we can try to find efficient ways of splitting A into A_r and A_c , distinguishing between:

- ▶ partition-oblivious heuristics: no prior information is required
- ightharpoonup partition-aware heuristics: requirement of A already partitioned



General remarks

A few general principles to guide us in the construction of the heuristics:

- ▶ short rows/columns (w.r.t. the number of nonzeros) are more likely to be uncut in a good partitioning
- ▶ if a row/column is uncut, the partitioner decided at the previous iteration that it was convenient to do so.
 We shall try to keep, as much as possible, those rows/columns uncut again.



A simple heuristic is the extension of the original algorithm used in medium-grain.

Partition-oblivious version:

```
for all a_{ij} \in A do

if nz_r(i) < nz_c(j) then

assign a_{ij} to A_r

else if nz_c(j) < nz_r(i) then

assign a_{ij} to A_c

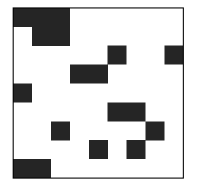
else

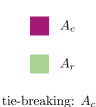
assign a_{ij} to according to tie-breaker

end if

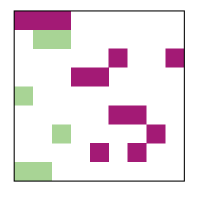
end for
```

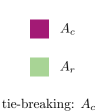












Universiteit Utrecht

Partition-aware version:

```
for all a_{ij} \in A do

if row i is uncut and column j is cut then

assign a_{ij} to A_r

else if row i is cut and column j is uncut then

assign a_{ij} to A_c

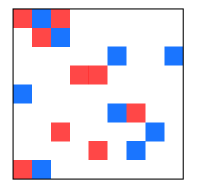
else

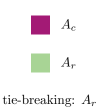
assign a_{ij} as in the partition-oblivious variant

end if

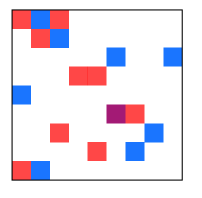
end for
```

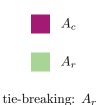




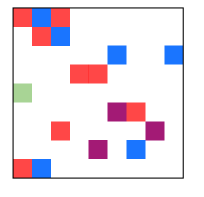


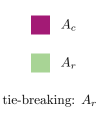






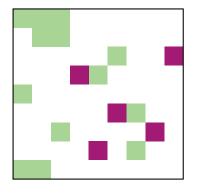








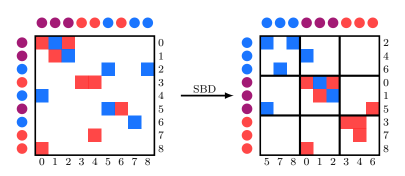
Individual assignment of nonzeros



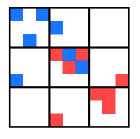




Separated Block Diagonal (SBD) form of a partitioned matrix: we separate uncut and cut rows and columns.





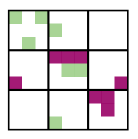


The SBD form is a 3×3 block matrix

$$\begin{bmatrix} \dot{A}_{00} & \dot{A}_{01} \\ \dot{A}_{10} & \dot{A}_{11} & \dot{A}_{12} \\ & \dot{A}_{21} & \dot{A}_{22} \end{bmatrix}$$

 \dot{A}_{01} , \dot{A}_{10} , \dot{A}_{12} , \dot{A}_{21} can be easily assigned in our framework.





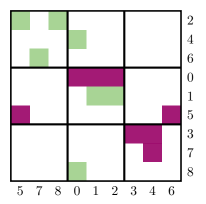
$$\begin{bmatrix} A_r/A_c & A_r \\ A_c & M & A_c \\ & A_r & A_r/A_c \end{bmatrix}$$

$$\blacksquare A_c$$

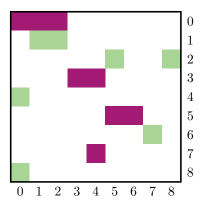
 $\dot{A}_{01}, \dot{A}_{10}, \dot{A}_{12}, \dot{A}_{21}$ can be easily assigned in our framework.

- ▶ A_r/A_c means that the size of the block determines whether it is assigned to A_r or A_c ;
- ▶ the nonzeros in the middle block are assigned individually (*M* stands for "mixed" assignment)







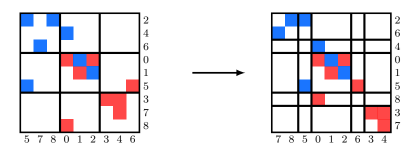


▶ We reverse the permutations of rows and columns, obtaining *A* back, with new assignment.

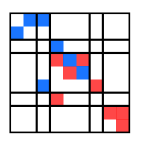




Separated Block Diagonal form of order 2 (SBD2) of a matrix: we split the top, bottom, left and right blocks, separating the empty and nonempty parts.





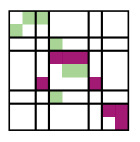


The SBD2 form of a matrix is the following 5×5 block matrix:

$$\begin{bmatrix} \ddot{A}_{00} & \ddot{A}_{01} \\ \ddot{A}_{10} & \ddot{A}_{11} & \ddot{A}_{12} \\ & \ddot{A}_{21} & \ddot{A}_{22} & \ddot{A}_{23} \\ & & \ddot{A}_{32} & \ddot{A}_{33} & \ddot{A}_{34} \\ & & & \ddot{A}_{43} & \ddot{A}_{44} \end{bmatrix}$$

In this form, other than having information on nonzeros (rows/columns cut/uncut), we also have information on their neighbors (nonzeros in the same row and column).



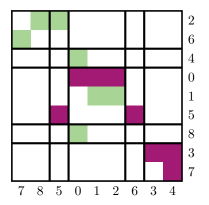


$$\begin{bmatrix} A_r & A_r \\ A_c & A_r/A_c & A_r \\ & A_c & M & A_c \\ & & A_r & A_r/A_c & A_c \\ & & & A_r & A_c \end{bmatrix}$$

In this form, other than having information on nonzeros (rows/columns cut/uncut), we also have information on their neighbors (nonzeros in the same row and column).

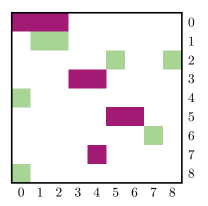


Individual assignment of blocks of nonzeros





Individual assignment of blocks of nonzeros



 \blacktriangleright We reverse the permutations of rows and columns, obtaining A back, with new assignment.





Partial assignment of rows and columns

- ▶ Main idea: Every time we assign a nonzero to either A_r or A_c , all the other nonzeros in the same row/column should be assigned to it as well, to prevent communication.
- ▶ Main issue: Hard to assign complete rows/column: a nonzero cannot be assigned to both A_r and A_c .

We need to reason in terms of **partial assignment**:

- computation of a **priority vector**: a permutation of the indices $\{0, \ldots, m+n-1\}$ (decreasing priority)
 - $\diamond \{0,\ldots,m-1\}$ correspond to rows;
 - $\{m, \ldots, m+n-1\}$ to columns.
- overpainting algorithm.

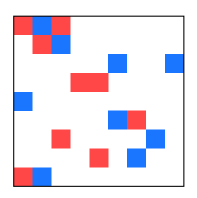


```
Require: Priority vector v, matrix A
Ensure: A_r, A_c
A_r := A_c := \emptyset
for i = m + n - 1, \dots, 0 do
if v_i < m then
Add the nonzeros of row i to A_r
else
Add the nonzeros of column i - m to A_c
end if
end for
```

- In this formulation of the algorithm, every nonzero is assigned twice:
- ▶ the algorithm is **completely deterministic**: A_r and A_c depend entirely on the priority vector v.



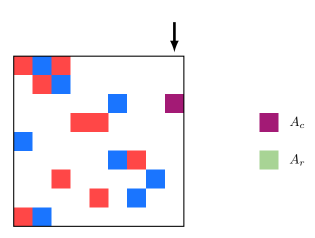




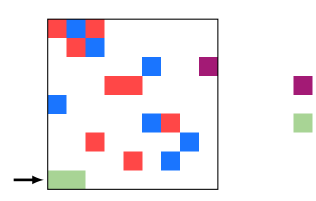




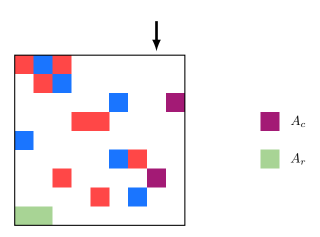




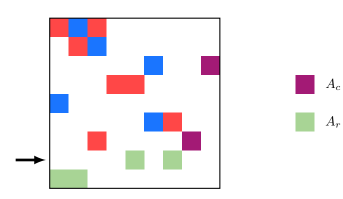




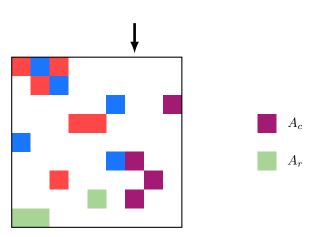




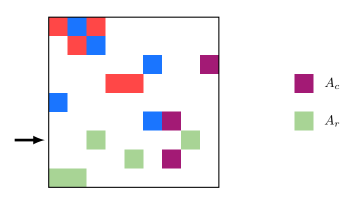




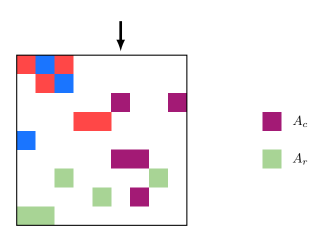




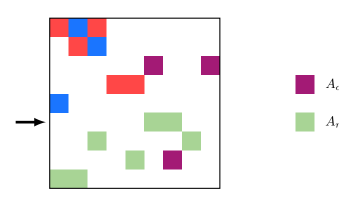




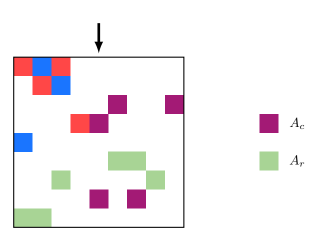




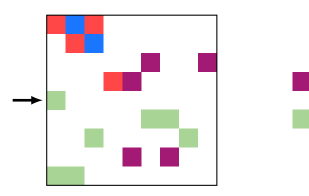




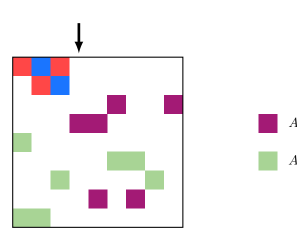




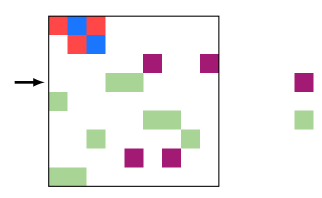




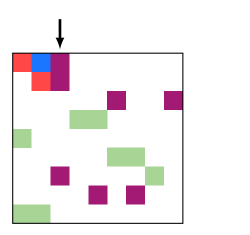








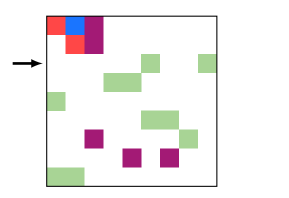








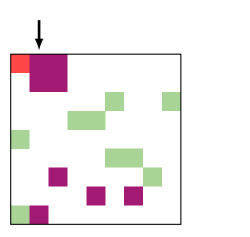








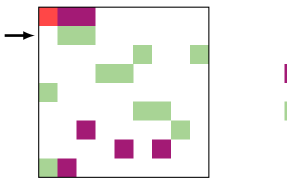




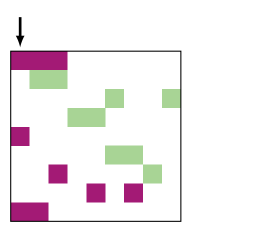








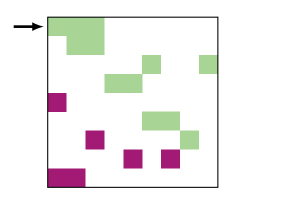


















We used a structured approach for the construction of v: 30 different heuristics.

Generating schemes with three steps:

- 1. Usage of previous partitioning
- 2. Sorting (w.r.t the number of nonzeros, in ascending order)
- 3. Internal order of indices



We used a structured approach for the construction of v: 30 different heuristics.

Generating schemes with three steps:

- 1. Usage of previous partitioning
 - partition-oblivious
 - partition-aware
- 2. Sorting (w.r.t the number of nonzeros, in ascending order)
- 3. Internal order of indices



We used a structured approach for the construction of v: 30 different heuristics.

Generating schemes with three steps:

1. Usage of previous partitioning

- 2. Sorting (w.r.t the number of nonzeros, in ascending order)
 - sorted (with or without refinement)
 - unsorted
- 3. Internal order of indices



We used a structured approach for the construction of v: 30 different heuristics.

Generating schemes with three steps:

1. Usage of previous partitioning

2. Sorting (w.r.t the number of nonzeros, in ascending order)

- 3. Internal order of indices
 - concatenation
 - mixing (either alternation or spread)
 - random (only when not sorting)
 - simple (only when sorting)



Independent set formulation

Partial assignment of rows and columns seems an interesting idea, but we want to reduce, as much as possible, the number of cut rows/columns.

Goal: Find the biggest subset of $\{0, \ldots, m+n-1\}$ which can be assigned completely (i.e. full rows and full columns) without causing communication.

Graph theory approach: translating the sparsity pattern of A in a particular way, we are looking for a **maximum independent set**.



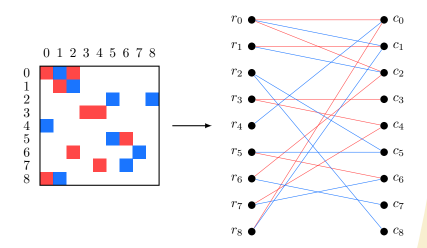
Construction of the graph

We construct the bipartite graph $G = (L \cup R, E)$ as follows:

- ▶ Rows and columns are vertices
 - $L = \{r_0, \dots, r_{m-1}\}$
 - $\land R = \{c_0, \dots, c_{n-1}\}$
- Edges correspond to nonzeros: $e = (r_i, c_j) \iff a_{ij} \neq 0$



Construction of the graph





Maximum independent set

Definition

An independent set is a subset $V' \subseteq V$ such that $\forall u, v \in V'$, $(u, v) \notin E$. A maximum independent set is an independent set of G with maximum cardinality.

- ▶ Our desired object is the maximum independent set
- ► The complement of a (maximum) independent set is a (minimum) vertex cover



Maximum independent set

In general, computing a maximum independent set is as hard as partitioning the matrix (both NP-hard problems).

But, luckily, our graph is bipartite:

- ► Kőnig's Theorem: on bipartite graphs, maximum matchings and minimum vertex covers have the same size;
- ▶ Hopcroft-Karp algorithm: $\mathcal{O}\left(N\sqrt{m+n}\right)$ algorithm to compute a maximum matching on a bipartite graph

In our case it is not very demanding to compute a maximum independent set.



Maximum independent set

Given the set of indices I, let S_I denote the maximum independent set computed on the matrix A(I).

One partition-oblivious heuristic to compute v:

1. let
$$I = \{0, ..., m + n - 1\}$$
, then $v := (S_I, I \setminus S_I)$

For partition-aware heuristics, let U be the set of uncut indices, C be the set of cut indices; we have three possibilities:

- 1. we compute S_U and have $v := (S_U, U \setminus S_U, C)$;
- 2. we compute S_U , S_C and have $v := (S_U, U \setminus S_U, S_C, C \setminus S_C)$;
- 3. we compute S_U , then we define $U' := U \setminus S_U$ and compute $S_{C \cup U'}$, having $v := (S_U, S_{C \cup U'}, (C \cup U') \setminus S_{C \cup U'})$.



General framework for experiments

Require: Sparse matrix A

Ensure: Partitioning for the matrix A

Partition A with Mondriaan using the medium-grain method

for $i = 1, \ldots, iter_{max}$ do

Use any of the heuristics described previously to compute A_r and A_c

Construct B from A_r and A_c

Partition B with Mondriaan using the row-net model

Re-construct A with the new partitioning

end for

Unique framework for both partition-oblivious and partition-aware types of heuristics.



Implementation

All of the heuristics have been implemented following these steps:

- 1. MATLAB prototyping
- 2. Core C implementation (MATLAB compatibility through MEX files)
- 3. Full C implementation

The Hopcroft-Karp algorithm for the maximum independent set computation was implemented in the Python programming language.



Implementation

Randomness involved during the computation of A_r and A_c and during the actual partitioning. To obtain meaningful results:

- ▶ 20 independent initial partitionings
- for each, 5 independent runs of the heuristic and subsequent partitioning ($iter_{max} = 1$)

18 matrices used for tests:

- ► rectangular vs. square
- ▶ 10th Dimacs Implementation Challenge



Preliminary selection

Wide selections of heuristics, preliminary selection is necessary.

5 matrices used:

- ▶ dfl001;
- ▶ tbdlinux;
- ▶ nug30;
- rgg_n_2_18_s0;
- bcsstk30.



Preliminary selection

Partition-oblivious heuristics (17 different algorithms):

- ▶ In general, results are much worse than medium-grain method.
- ▶ Mixing rows and columns in partial assignment is a bad idea
- ► Individual assignment of nonzero (po_localview) is the best strategy (7% worse than medium-grain)
- ▶ Maximum independent set computation (po_is) yields interesting results (16% lower communication volume in one matrix, but in general 12% worse than medium-grain)

2 heuristics selected for deeper investigation.



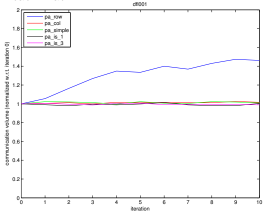
Preliminary selection

Partition-aware heuristics (21 different algorithms):

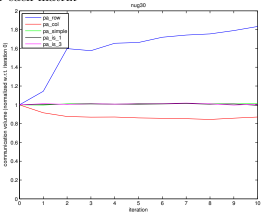
- ▶ Results closer to medium-grain efficiency
- ► SBD and SBD2 forms are not worthwile, nor individual assignment of nonzeros
- ▶ Refinement in sorting does not yield a substantial difference
- ► Unsorted concatenation of rows and columns (pa_row and pa_col) produces good results with rectangular matrices: they can be combined into a po_localbest heuristic (which tries both and picks the best)
- ► Maximum independent set strategies (pa_is_1 and pa_is_3) are very close to medium-grain, even better in a few matrices
- 5 heuristics selected for deeper investigation.



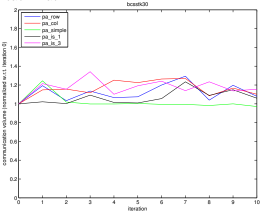
We are developing a fully iterative scheme: how many iterations do we have to execute?



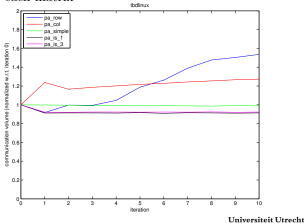
We are developing a fully iterative scheme: how many iterations do we have to execute?



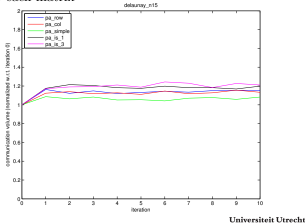
We are developing a fully iterative scheme: how many iterations do we have to execute?



We are developing a fully iterative scheme: how many iterations do we have to execute?



We are developing a fully iterative scheme: how many iterations do we have to execute?



We are developing a fully iterative scheme: how many iterations do we have to execute?

- ▶ Usually 1 iteration is enough to show improvements, if any.
- ▶ More iterations can worsen the communication volume.



Analysis of the performance of the best heuristics

Partition-oblivious heuristics:

- ▶ No devised heuristic was able to improve medium-grain
- ▶ The preliminary results confirmed:
 - ♦ Individual assignment of nonzeros 7% worse than medium-grain
 - \diamond Computing the maximum independent set 22% worse than medium-grain



Analysis of the performance of the best heuristics

Partition-aware heuristics:

- ► Concatenation interesting strategy:
 - ♦ pa_row and pa_col 8% worse than medium-grain
 - ⋄ localbest method takes best of both: only 4% worse than medium-grain
- ► Similar good results for the other strategies (between 4% and 8% higher communication volume than medium-grain)
- ▶ No algorithm was able to beat medium-grain
- ► Considering only rectangular matrices, our methods work better: they improve medium-grain, even if only by little (1-2%)



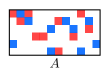
Is there something we can do to improve the results?

Medium-grain employs a procedure of **iterative refinement**:

- 1. A is partitioned into two sets $(A_0 \text{ and } A_1)$
- 2. we create again the matrix B of the medium-grain method (example: $A_r = A_0$ and $A_c = A_1$)
- 3. we retain communication volume: the first n columns of B are assigned to a single processor, and similarly for the other m
- 4. we create the hypergraph from this B and a single run of Kernighan-Lin is performed
- 5. we repeat steps 1-4 until no improvement is found, then we swap the roles of A_0 and A_1 for the creation of A_r and A_c
- 6. we repeat step 5 until no other improvement is found

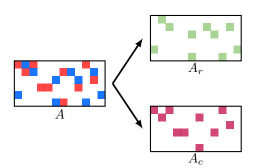


Kernighan-Lin method is **monotonically non-increasing**: during iterative refinement, the communication volume is either lowered or remains at the same value.





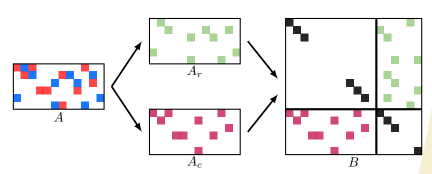
Kernighan-Lin method is **monotonically non-increasing**: during iterative refinement, the communication volume is either lowered or remains at the same value.





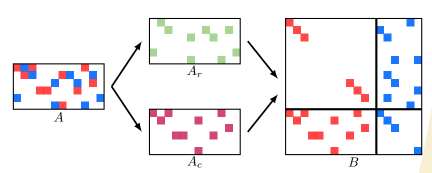


Kernighan-Lin method is **monotonically non-increasing**: during iterative refinement, the communication volume is either lowered or remains at the same value.





Kernighan-Lin method is **monotonically non-increasing**: during iterative refinement, the communication volume is either lowered or remains at the same value.







With iterative refinements, results are in general better:

- ▶ partition-oblivious algorithms:
 - ♦ po_localview still 7% worse than medium-grain
 - ♦ po_is now 6% worse than medium-grain (down from 22%)
- ▶ partition-aware algorithms:
 - pa_row and pa_col now 2% and 1% worse than medium-grain (down from 8%)
 - \diamond localbest now 1% better than medium-grain (down from 4% worse)
 - ♦ pa_simple now 2% worse than medium-grain (down from 4%)
 - pa_is_1 and pa_is_3 now 1% worse than medium-grain (down from 5% and 8%)

Now, with rectangular matrices, computing the independent set produces an average communication volume 4% lower than medium-grain.

Universiteit Utrecht

Conclusions

We originally had two research directions:

▶ Improving the quality of the initial partitioning

▶ Developing a fully-iterative scheme



Conclusions

We originally had two research directions:

- ▶ Improving the quality of the initial partitioning
 - ♦ We were not able to outperform medium-grain
- ▶ Developing a fully-iterative scheme



Conclusions

We originally had two research directions:

▶ Improving the quality of the initial partitioning

- ▶ Developing a fully-iterative scheme
 - \diamond We were able to outperform medium-grain only by a small margin
 - ♦ Computing the independent set is worthwile. Also results about concatenation of rows and columns can be explained with it.
 - ♦ Our approach works well with rectangular matrices



Further research

A number of possibilities for further development:

- ► Keep testing other strategies to gain more confidence on medium-grain
- Maximum weighted independent set to maximize the number of nonzeros completely assigned
- ► If our approach is confirmed to work consistently well with rectangular matrices, it could be added to Mondriaan:
 - the program detects that the matrix is strongly rectangular and asks user for input;
 - 2. the user decides whether he wants to sacrifice computation time for a better partitioning and execute our approach

