Bad News: Hardness Results

Good News: Tractability Results

Summary 0

On the Complexity of SNP Block Partitioning Under the Perfect Phylogeny Model

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Introduction

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Outline

1 Introduction

- The Model and the Problem
- The Integrated Approach

2 Bad News: Hardness Results

- Hardness of PP-Partitioning of Haplotype Matrices
- Hardness of PP-Partitioning of Genotype Matrices

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- Perfect Path Phylogenies
- Tractability of PPP-Partitioning of Genotype Matrices

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The Model and the Problem

What is haplotyping and why is it important?

You hopefully know this after the previous three talks...

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The Model and the Problem

General formalization of haplotyping.

Inputs

- A genotype matrix G.
- The rows of the matrix are taxa / individuals.
- The columns of the matrix are SNP sites / characters.

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Outputs

- A haplotype matrix H.
- Pairs of rows in H explain the rows of G.
- The haplotypes in *H* are biologically plausible.

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Summary

The Model and the Problem

Our formalization of haplotyping.

Inputs

- A genotype matrix G.
- The rows of the matrix are individuals / taxa.
- The columns of the matrix are SNP sites / characters.
- The problem is directed: one haplotype is known.
- The input is biallelic: there are only two homozygous states (0 and 1) and one heterozygous state (2).

Outputs

- A haplotype matrix H.
- Pairs of rows in H explain the rows of G.
- The haplotypes in *H* form a perfect phylogeny.

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The Model and the Problem

We can do perfect phylogeny haplotyping efficiently, but ...

Data may be missing.

- This makes the problem NP-complete
- ... even for very restricted cases.

Solutions:

- Additional assumption like the rich data hypothesis.
- In a perfect phylogeny is possible.
 - This can be caused by chromosomal crossing-over effects.
 - This can be caused by incorrect data.
 - This can be caused by multiple mutations at the same sites.

Solutions:

- Look for phylogenetic networks.
- Correct data.
- Find blocks where a perfect phylogeny is possible.

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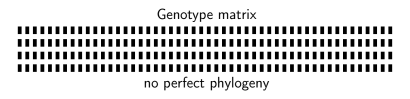
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The Integrated Approach

How blocks help in perfect phylogeny haplotyping.

- **1** Partition the site set into overlapping contiguous blocks.
- Compute a perfect phylogeny for each block and combine them.
- **③** Use dynamic programming for finding the partition.



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Genotype matrix

perfect phylogeny

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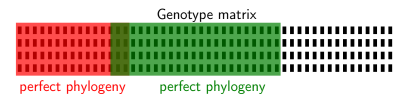
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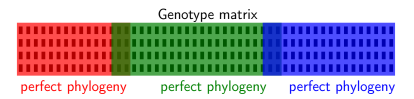
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The Integrated Approach

Objective of the integrated approach.

- Partition the site set into noncontiguous blocks.
- Compute a perfect phylogeny for each block and combine them.
- Occupies Compute partition while computing perfect phylogenies.

Genotype matrix no perfect phylogeny

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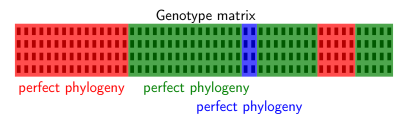
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The Integrated Approach

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The Integrated Approach

The formal computational problem.

We are interested in the computational complexity of the function χ_{PP} :

- It gets genotype matrices as input.
- It maps them to a number k.
- This number is minimal such that the sites can be covered by k sets, each admitting a perfect phylogeny.
 (We call this a pp-partition.)

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Hardness of PP-Partitioning of Haplotype Matrices

Finding pp-partitions of haplotype matrices.

We start with a special case:

- The inputs *M* are already haplotype matrices.
- The inputs *M* do not allow a perfect phylogeny.
- What is $\chi_{PP}(M)$?

Exar	nple	9			
	0	0	0	1	
	0	1	0	0	
	1	0	0	0	
	0	1	0	0	No perfect phylogeny is possible.
<i>M</i> :	1	0	0	0	
	0	1	0	1	
	1	1	0	0	
	0	0	1	0	
	1	0	1	0	

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Exan	nple	9			
М:	0 0 1 0 1 0 1 0 1	0 1 0 1 1 1 0 0 0	0 0 0 0 0 0 1 1	1 0 0 1 0 0 0 0	Perfect phylogeny Perfect phylogeny $\chi_{PP}(M) = 2.$

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Summary

Hardness of PP-Partitioning of Haplotype Matrices

Bad news about pp-partitions of haplotype matrices.

Theorem

Finding optimal pp-partition of haplotype matrices is equivalent to finding optimal graph colorings.

Proof sketch for first direction.

- Let G be a graph.
- 2 Build a matrix with a column for each vertex of G.
- For each edge of G add four rows inducing the submatrix $\begin{pmatrix} 0 & 0 \\ 1 & 1 \\ 1 & 1 \end{pmatrix}$.
- The submatrix enforces that the columns lie in different perfect phylogenies.

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Hardness of PP-Partitioning of Haplotype Matrices

Implications for pp-partitions of haplotype matrices.

Corollary

If $\chi_{PP}(M) = 2$ for a haplotype matrix M, we can find an optimal pp-partition in polynomial time.

Corollary

Computing χ_{PP} for haplotype matrices is

- NP-hard,
- not fixed-parameter tractable, unless P = NP,
- very hard to approximate.

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Summary

Hardness of PP-Partitioning of Genotype Matrices

Finding pp-partitions of genotype matrices.

Now comes the general case:

- The inputs *M* are genotype matrices.
- The inputs *M* do not allow a perfect phylogeny.
- What is $\chi_{PP}(M)$?

Exar	nple	9			
	2	2	2	2	
	1	0	0	0	No perfect phylogeny is possible.
<i>M</i> :	0	0 0	0	1	
<i>IVI</i> .	0	0	1	0	
	0	2	2	0	
	1	1	0	0	

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Summary

Hardness of PP-Partitioning of Genotype Matrices

Finding pp-partitions of genotype matrices.

Now comes the general case:

- The inputs *M* are genotype matrices.
- The inputs *M* do not allow a perfect phylogeny.
- What is $\chi_{PP}(M)$?

Example										
M :	2 1 0 0 0 1	2 0 0 2 1	2 0 1 2 0	2 0 1 0 0	Perfect phylogeny Perfect phylogeny $\chi_{PP}(M) = 2.$					

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Summary

Hardness of PP-Partitioning of Genotype Matrices

Bad news about pp-partitions of haplotype matrices.

Theorem

Finding optimal pp-partition of genotype matrices is at least as hard as finding optimal colorings of 3-uniform hypergraphs.

Proof sketch.

- Let G be a 3-uniform hypergraph.
- 2 Build a matrix with a column for each vertex of G.
- For each hyperedge of G add four rows inducing the submatrix $\begin{pmatrix} 2 & 2 & 2 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}$.
- The submatrix enforces that the three columns do not all lie in the same perfect phylogeny.

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Summary

Hardness of PP-Partitioning of Genotype Matrices

Implications for pp-partitions of genotype matrices.

Corollary

Even if we know $\chi_{PP}(M) = 2$ for a genotype matrix M, finding a pp-partition of any fixed size is still

- NP-hard,
- not fixed-parameter tractable, unless P = NP,
- very hard to approximate.

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Perfect Path Phylogenies

Automatic optimal pp-partitioning is hopeless, but...

- The hardness results are worst-case results for highly artificial inputs.
- Real biological data might have special properties that make the problem tractable.
- One such property is that perfect phylogenies are often perfect path phylogenies:
 In HapMap data, in 70% of the blocks where a perfect phylogeny is possible a perfect path phylogeny is also possible.

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Perfect Path Phylogenies

Example of a perfect path phylogeny.

Genotype matrix					Haplotype matrix				Perfect path phylogeny		
	А	В	С			А	В	С			
	2	2	2	.		1	0	0	A / \ B		
<i>G</i> :	0	2	0			0	1	1			
	2	0	0			0	0	0	i i i		
	0	2	2		<i>H</i> :	0	1	0			
_			_			0	0	0	Ľ		
						1	0	0	_ Ğ		
						0	0	0			
						0	1	1			

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Perfect Path Phylogenies

The modified formal computational problem.

We are interested in the computational complexity of the function χ_{PPP} :

- It gets genotype matrices as input.
- It maps them to a number k.
- This number is minimal such that the sites can be covered by k sets, each admitting a perfect path phylogeny.
 (We call this a ppp-partition.)

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Tractability of PPP-Partitioning of Genotype Matrices

Good news about ppp-partitions of genotype matrices.

Theorem

Optimal ppp-partitions of genotype matrices can be computed in polynomial time.

Algorithm

- Build the following partial order:
 - Can one column be above the other in a phylogeny?
 - Can the columns be the two children of the root of a perfect path phylogeny?
- Over the partial order with as few compatible chain pairs as possible.

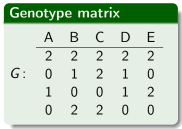
For this, a maximal matching in a special graph needs to be computed.

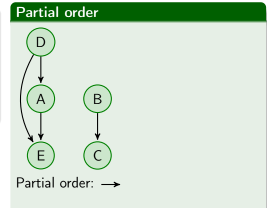


Summary

- Finding optimal pp-partitions is intractable.
- It is even intractable to find a pp-partition when just two noncontiguous blocks are known to suffice.
- For perfect path phylogenies, optimal partitions can be computed in polynomial time.

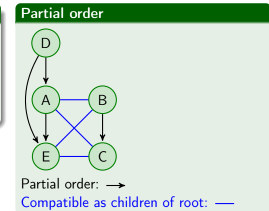
The algorithm in action. Computation of the partial order.



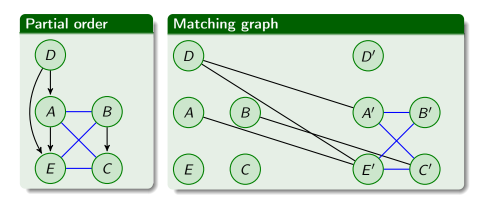


The algorithm in action. Computation of the partial order.

Genotype matrix									
	А	В	С	D	Е				
	2	2	2	2	2				
<i>G</i> :	0	1	2	1	0				
	1	0	0	1	2				
	0	2	2	0	0				

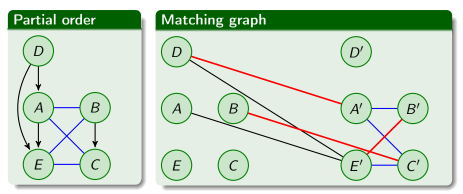


The algorithm in action. The matching in the special graph.





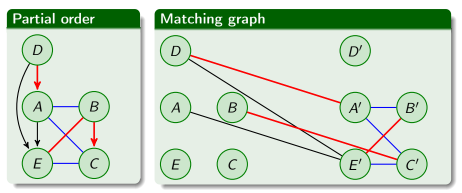
The algorithm in action. The matching in the special graph.



A maximal matching in the matching graph



The algorithm in action. The matching in the special graph.



A maximal matching in the matching graph induces perfect path phylogenies.

