## SET COVERING MACHINE FOR SNPs DISCOVERY

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March, 23<sup>rd</sup> 2020 CBIO Meeting  $1. \mathrm{Short} \mathrm{reminder} \mathrm{on} \mathrm{GWAS}$ 

#### Goal of Genome-wise association studies (GWAS)

- Goal: Discover gene mutations *linked* to a disease.
- GWAS will not provide: causality relations or biological understanding of a disease.
- Applications to common diseases:
  - Inflammatory Bowel Diseases
  - Auto-immune diseases
  - ▶ Metabolic diseases (T2 diabetes, obesity, BMI)
  - Multiple sclerosis
  - ► Cancer

### The CD/CV hypothesis

- Common diseases are partly caused by common variants
- Consequence: each mutation can only have a *small effect*



#### Allele Frequency

Source: Bush et al (2012)

 $Linkage\ disequilibrium\ (LD):$  correlation between close-by alleles on the genome



Idea of using single nucleotide polymorphisms (SNPs):

- SNP: Single nucleotide polymorphism
- There are many high-LD blocks on the genome
- We can use SNPs as markers of an LD block

#### GWAS

- Gather  $n \sim 10^3$  individuals
- Observe the phenotype:
  - ▶ quantitative (BMI, cholesterol, height)
  - ▶ or qualitative (case-control for common disease).
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Source: Ikram et al (2010)

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- The genetic mutations are in interaction
- Need to consider SNPs *jointly*

• 
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Very small statistical power



## 2. Set covering machines for SNPs discovery

Each SNP has value  $\in$  {aa, aA, AA}.

Genotype	Dominant	Recessive	Allelic dosage	One-hot
aa	0	0	0	100
aA	1	0	1	010
AA	1	1	2	001

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We will use one-hot encoding  $\rightarrow$  binary features

#### Setting of the set covering machine (SCM)

- $y_i \in \{0, 1\}$  (case/control GWAS)
- $x_{i,j} \in \{0,1\}$  (one-hot encoding)
- $p \gg n$  with true model assumed to be *very sparse*

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• SCM learns a boolean function of the features:

$$f(\mathbf{x}) = \bigwedge_{j \in \mathcal{R}} h_j(\mathbf{x}),$$

where  $\mathcal{R} \in$ is the set of rules to learn.

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- SCM only learns a conjunction of SNPs to explain the phenotype.

Haussler algorithm:

- Assume there is a combination of features that perfectly classifies the dataset:  $\mathbf{y} = \bigwedge_{j \in \mathcal{R}} h_j$
- How to find the sparsest possible combination of features?
- Only consider rules that correctly classify *all* positive examples  $(y_i = 1)$ .

• Example: what conjunction of  $h_j$ s equals **y**?

	У	$h_1$	$h_2$	$h_3$	$h_4$
	0	0	0	1	1
$\mathcal{N}$	0	1	0	0	1
(negative examples)	0	1	1	0	0
* /	0	1	1	1	0
$\mathcal{P}$	1	1	1	1	1
(positive examples)	1	1	1	1	1

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	0	0	0	1	1	0
٨٢	0	1	0	0	1	0
JV	0	1	1	0	0	0
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$\mathcal{D}$	1	1	1	1	1	1
P	1	1	1	1	1	1

• Smallest number of sets  $\{1\}, \{1, 2\}, \{2, 3\}, \{3, 4\}$  whose union is  $\{1, 2, 3, 4\}$ .

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- This is the set cover problem (NP hard)
- We use a greedy approach

We choose the rule with maximum usefulness:

$$U_h = |\mathcal{A}_h| - q|\mathcal{B}_h|,$$

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- We allow errors on positive examples
- q controls this error

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	$ \mathcal{A}_h $	1	2	2	2
	$ \mathcal{B}_h $	0	2	1	0
	$\mathcal{U}_h$	1	0	1	2

Example (with q = 1):

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•  $\mathcal{R} \leftarrow \{h_4\}$ •  $\mathcal{N} \leftarrow \mathcal{N} \setminus \mathcal{A}_{h_4}$ 

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- $\mathcal{R} \leftarrow \{h_4\}$
- $\mathcal{N} \leftarrow \mathcal{N} \setminus \mathcal{A}_{h_4}$
- $\mathcal{P} \leftarrow \mathcal{P} \setminus \mathcal{B}_{h_4}$

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- We have finished the job:  $\mathcal{N} = \emptyset$
- Early stopping:  $|\mathcal{R}| \ge s$  with parameter  $s \ge 1$
- There remains only useless rules:  $|\mathcal{A}_h| = |\mathcal{B}_h| = 0$

- Each greedy step is fast to compute:
  - Let  $\mathcal{I}_{\mathcal{N}}$  be the (current) indices of the negative examples.
  - ►  $|\mathcal{A}_h| = |\mathcal{I}_N| \sum_{i \in \mathcal{I}_N} x_{i,j}$  if *h* is the presence rule of feature *j*
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- Overall complexity  $\mathcal{O}\left(|\mathcal{R}|ns\right)$
- Limited memory usage

#### Upper bound on the risk (1/2)

SCM is a sample compression algorithm

- Given a model f learnt by an SCM, there exist
  - a set of individuals  $\mathcal{Z} \in \{1, \cdots, n\}$

• a message string  $\sigma$  containing additional information, such that h can be reconstructed from  $\mathcal{Z}$ .

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• Marchand and Sokolova (2006) established that:

$$\mathbb{P}\left(\forall S \sim D, \forall h, R(h) \leq \varepsilon(h, S, \delta)\right) \geq 1 - \delta$$

- $\varepsilon$  depends on  $\mathcal{Z}$  and the of classif. errors made on  $S \setminus \mathcal{Z}$ .
- $\varepsilon$  does not depend on p.

Consequences:

- The bound does not depend on *p*: theoretical performance guarantee
- It can be used for hyperparameter selection (Marchand et Shawe-Taylor, 2002).

#### Conclusion

- Set covering machine
  - ▶ learns a boolean conjunction of SNPs
  - runs fast
  - does not suffer from  $p \gg n$
- However there are other issues:
  - Many SNPs can have same  $\mathcal{U}_h$ : which one to choose?
  - Only conjunctions of SNPs

# THANK YOU

- SCM: Marchand, M. and Shawe-Taylor, J., *The set covering machine*, JMLR, 2002
- Risk bound for SCM: Marchand, M and Sokolova, M., *Learning with Decision Lists of Data-Dependent Features*, JMLR, 2005
- GWAS: Bush, W, Moore, J., Lewitter, F., and Kann, M., *Chapter* 11: Genome-Wide Association Studies, Plos Comp. Biol., 2012
- Epistasis: Lehner, B., Molecular mechanisms of epistasis within and between genes, Trends in Gen., 2011

$$\varepsilon(h, S, \delta) = 1 - \exp\left(\frac{-1}{n - |\mathcal{Z}| - r} \left[\log\binom{m}{|\mathcal{Z}|} + \log\binom{m - |\mathcal{Z}|}{r} + |h| \log(2\mathcal{N}(\mathcal{Z})) + \log\Omega\right]\right)$$