Graphical Models for Genomic Selection

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Background
A Bayesian network (BN) [6, 7] is a combination of:

- a directed acyclic graph $\mathcal{G} = (\mathbf{V}, \mathcal{A})$, in which each node $v_i \in \mathbf{V}$ corresponds to a random variable $X_i$ (a gene, a trait, an environmental factor, etc.);
- a global probability distribution over $\mathbf{X} = \{X_i\}$, which can be split into simpler local probability distributions according to the arcs $a_{ij} \in \mathcal{A}$ present in the graph.

This combination allows a compact representation of the joint distribution of high-dimensional problems, and simplifies inference using the graphical properties of $\mathcal{G}$. 
The two main properties of Bayesian networks

The defining characteristic of BNs is that graphical separation implies (conditional) probabilistic independence. As a result, the global distribution factorises into local distributions: each one is associated with a node $X_i$ and depends only on its parents $\Pi_{X_i}$,

$$ P(X) = \prod_{i=1}^{p} P(X_i \mid \Pi_{X_i}). $$

In addition, we can visually identify the Markov blanket of each node $X_i$ (the set of nodes that completely separates $X_i$ from the rest of the graph, and thus includes all the knowledge needed to do inference on $X_i$).
From the definition, if we have a set of traits and markers for each variety, all we need for GS and GWAS are the Markov blankets of the traits [11]. Using common sense, we can make some additional assumptions:

- traits can depend on markers, but not vice versa;
- traits that are measured after the variety is harvested can depend on traits that are measured while the variety is still in the field (and obviously on the markers as well), but not vice versa.

Most markers are discarded when the Markov blankets are learned. Only those that are parents of one or more traits are retained; all other markers’ effects are indirect and redundant once the Markov blankets have been learned. Assumptions on the direction of the dependencies allow to reduce Markov blankets learning to learning the parents of each trait, which is a much simpler task.
Learning
Learning the Bayesian network

1. **Feature Selection.**
   
   1.1 For each trait, use the SI-HITON-PC algorithm [1, 10] to learn the parents and the children of the trait; children can only be other traits, parents are mostly markers, spouses can be either. Dependencies are assessed with Student's $t$-test for Pearson’s correlation [5] and $\alpha = 0.01$.
   
   1.2 Drop all the markers which are not parents of any trait.

2. **Structure Learning.** Learn the structure of the BN from the nodes selected in the previous step, setting the directions of the arcs according to the assumptions in the previous slide. The optimal structure can be identified with a suitable goodness-of-fit criterion such as BIC [9]. This follows the spirit of other hybrid approaches [3, 12], that have shown to be well-performing in literature.

3. **Parameter Learning.** Learn the parameters of the BN as a Gaussian BN [6]: each local distribution in a linear regression and the global distribution is a hierarchical linear model.
The local distribution of each trait $X_i$ is a linear model

$$X_i = \mu + \Pi_{X_i} \beta + \varepsilon$$

$$= \mu + X_j \beta_j + \ldots + X_k \beta_k + X_l \beta_l + \ldots + X_m \beta_m + \varepsilon$$

which can be estimated any frequentist or Bayesian approach in which the nodes in $\Pi_{X_i}$ are treated as fixed effects (e.g. ridge regression [4], elastic net [13], etc.).

For each marker $X_i$, the nodes in $\Pi_{X_i}$ are other markers in LD with $X_i$ since $\text{COR}(X_i, X_j|\Pi_{X_i}) \neq 0 \iff \beta_j \neq 0$. This is also intuitively true for markers that are children of $X_i$, as LD is symmetric.
A caveat about causal interpretations

Even though “good” BNs have a structure that mirrors cause-effect relationships [8], and even though there is ample literature on how to learn causal BNs from observational data, **inferring causal effects from a BN requires great care** even with completely independent data (i.e. with no family structure).
The MAGIC data (Multiparent Advanced Generation Inter-Cross) include 721 varieties, 16K markers and the following phenotypes:

- flowering time (FT);
- height (HT);
- yield (YLD);
- yellow rust, as measured in the glasshouse (YR.GLASS);
- yellow rust, as measured in the field (YR.FIELD);
- mildew (MIL) and
- fusarium (FUS).

Varieties with missing phenotypes or family information and markers with > 20% missing data were dropped. The phenotypes were adjusted for family structure via BLUP and the markers screened for MAF > 0.01 and COR < 0.99.
51 nodes (7 traits, 44 markers), 86 arcs, 137 parameters for 600 obs.
Phenotypic traits in MAGIC

- FT
- MIL
- HT
- YR.GLASS
- YLD
- YR.FIELD
- FUS
Friedman et al. [2] proposed an approach to assess the strength of each arc based on bootstrap resampling and model averaging:

1. For $b = 1, 2, \ldots, m$:
   1.1 sample a new data set $X^*_b$ from the original data $X$ using either parametric or nonparametric bootstrap;
   1.2 learn the structure of the graphical model $G_b = (V, E_b)$ from $X^*_b$.

2. Estimate the confidence that each possible arc $a_i$ is present in the true network structure $G_0 = (V, A_0)$ as

$$\hat{p}_i = \hat{P}(a_i) = \frac{1}{m} \sum_{b=1}^{m} 1_{\{a_i \in A_b\}},$$

where $1_{\{e_i \in E_b\}}$ is equal to 1 if $a_i \in A_b$ and 0 otherwise.
Averaged Bayesian network from MAGIC

81 out of 86 arcs from the original BN are significant.
Phenotypic traits in MAGIC

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<th>from</th>
<th>to</th>
<th>strength</th>
<th>direction</th>
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Arcs in the BN are highlighted in red in the table.

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Inference
Inference in Bayesian networks

Inference for BNs usually takes two forms:

- **conditional probability queries**, in which the distribution of one or more nodes of interest is investigated conditional on a second set of nodes (which are either completely or partially fixed);
- **maximum a posteriori queries**, in which the most likely outcome of a certain event (involving one or more nodes) conditional on evidence on a set of nodes (which are often completely fixed for computational reasons).

In practice this amounts to answering “what if?” questions (hence the name queries) about what could happen in observed or unobserved scenarios using posterior probabilities or density functions.
Flowering time: what if we fix directly related alleles?

Fixing 6 genes that are parents of FT in the BN to be homozygotes for early flowering (EARLY) or for late flowering (LATE).
Flowering time: which nodes we used...
Fixing 8 genes that are parents of YR.FIELD, then another 7 that are parents of YR.GLASS, either to be homozygotes for yellow rust susceptibility or for yellow rust resistance.
Yellow rust: nodes farther away can help...
If we have two varieties for which we scored low levels of fusarium (0 to 2), and are among the top 25% yielding, but one is tall (top 25%) and one is short (bottom 25%), which is the most probable allele for gene G3140?
G3140: information travels backwards...
Conclusions
Bayesian networks provide an **intuitive representation** of the relationships linking sets of phenotypes and markers, both within and between each other.

Given a few reasonable assumptions, we can learn a Bayesian network for multiple trait GWAS and GS efficiently and reusing state-of-the-art general-purpose algorithms.

Once learned, Bayesian networks provide a **flexible tool for inference** on both the markers and the phenotypes.

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