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## PCOS

- (Report results for microbe-outcome network). From the microbe-outcome network (Figure 2), the following nine genera are causal to PCOS ('group' node): *Alistipes*, *Blautia*, *Burkholderia*, *Desulfovibrio*, *Holdemanella*, *Knoellia*, *Prevotellaceae* NK3B31 group, *Ruminococcus*, and *Ruminococcus gnavus* group. We find their individual causal effects with do-calculus (Table 2).



(Report results for microbe-microbe network). From the microbe-outcome network (Figure 1), the following five genera are causal to T2D ('IRIS' node): *Butyrivibrio*, *Clostridium* XIVb, *Odoribacter*, *unclassified Bacteria*, and *unclassified Firmicutes*. To further investigate their individual effects, we implement do-calculus through logistic regression models on T2D given the neighbors of the genus of interest (Table 1). We implement three models: a simple logistic regression model regressed on the five genera causal to T2D, a logistic regression model regressed on a microbe and its neighbors, and a logistic regression model regressed on a microbe and its mediators. Their significance is denoted with a .

Table 1. Do-Calculus Results for T2D.

(Insert VAE results).

Genus	Model 1	Model 2	Literature Agreement
<i>Alistipes</i>	0.153458	4.68e-05	Unknown
<i>Blautia</i>	0	0	Unknown
<i>Burkholderia</i>	0	0	Unknown
<i>Desulfovibrio</i>	0	0	Unknown
<i>Holdemanella</i>	0	0	Unknown
<i>Knoellia</i>	0	0	Unknown
<i>Prevotellaceae</i> NK3B31 group	0	0	Unknown
<i>Ruminococcus</i>	0	0	Unknown
<i>Ruminococcus gnavus</i> group	0	0	Unknown

Table 2. Do-Calculus Results for PCOS.

(Insert VAE results).

Answer questions 1, 2, and 3. Explain BIRDMAN. Point to website for more results. Fix references to et al for many authors. We would like to thank our mentors, and Dr. Sam Degregori for guidance throughout this project.

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Causal discovery attempts to recover the true causal structure of a system given observed data. One way to model this causal structure is through a directed graphical model. A widely-used general-purpose causal discovery algorithm is the Peter-Clark (PC) algorithm [?]. It follows these key steps:

1. Start with a **complete undirected graph** (each node connected to all other nodes).
2. **Remove edges** based on statistical independence and conditional independence tests.
3. **Identify v-structures** (patterns like  $X \rightarrow Y \leftarrow Z$ ) to infer causal directions.
4. **Apply Meek's rules** to orient additional edges while preserving v-structures.

The result is a **CPDAG (Completed Partially Directed Acyclic Graph)**, which represents a set of causal structures consistent with the observed data, also known as the Markov Equivalence Class (MEC).

## Why Use PC?

- Works for different data types (as long as independence tests match the data distribution).
- Efficient for large datasets.
- Assumes the **causal Markov condition**, the **faithfulness** assumption, and **no hidden confounders**.