Topics of today:

- 1. Short introduction to / review of ILP
- 2. DCJ distance of balanced genomes
- 3. ILP for computing the DCJ distance of balanced genomes

Optimization problem - simple example: optimal meal problem

	Vitamin				
Let the following table give the nutritional values of a portion of each of two types of food		А	С	D	Calories
	Food P	225	100	200	600
	Food B	600	100	75	300
	Min. intake	1800	550	600	

A solution to this optimization problem is a meal composed of portions of Food P and Food B

The variables are $\begin{cases} x_p: \ \# \text{ of portions of Food P} \\ x_b: \ \# \text{ of portions of Food B} \end{cases}$

The number of portions cannot be negative, therefore the respective **domains** are

 $x_p \geq 0$ and $x_b \geq 0$

For achieving the minimum intake of each vitamin, any meal has to respect the **constraints** on the amount of vitamins:

 $\begin{array}{l} 225 x_{p} + 600 x_{b} \geq 1800 \mbox{ (Vitamin A)} \\ 100 x_{p} + 100 x_{b} \geq 550 \mbox{ (Vitamin C)} \\ 200 x_{p} + 75 x_{b} \geq 600 \mbox{ (Vitamin D)} \end{array}$

The **objective** is the minimization of calories in the meal: minimize $600x_p + 300x_b$.

(As not eating violates the vitamin constraints, the empty meal is an infeasible solution.)

Linear Programming (LP)

Given a set of decision variables

(each variable has a **domain**, stating its valid values)

Given a set of linear constraints in the given variables where ...

infeasible solution: tuple of valid values that violates at least one constraint

feasible solution: tuple of valid values that satisfies all constraints the set of all feasible solutions is the solution space (if no solution is feasible, the solution space is empty and the optimization problem itself is infeasible)

The problem is finding a best feasible solution by minimizing or maximizing the linear $\boldsymbol{objective}\ \boldsymbol{function}$ in the given variables (the objetive function defines the quality of the feasible solutions)

Linear Optimization Problem

LP - simple example: optimal meal problem

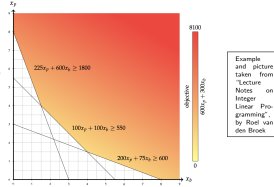
Graphical illustration:

The lines represent the constraints

The **colored area** corresponds to the **solution space** (set of feasible solutions), and the **gradient** indicates the value of the **objective function** in the solution space

The optimal meal consists of:

 $\begin{cases} x_p = 1.5 \ (\# \text{ of portions of Food P}) \\ x_b = 4 \quad (\# \text{ of portions of Food B}) \\ \text{for a total of 2100 calories} \end{cases}$



The computation time of an LP is polynomial in the number of variables and constraints.

Integer Linear Programming (ILP)

Variation of LP, where each variable is restricted to integer values

Powerful tool in combinatorial optimization:

many problems feature discrete decisions that can be modeled in an ILP

In contrast to linear programs, which are all solvable to optimality in polynomial time (in the number of variables and constraints), often there is no known polynomial bound on the computation time of integer linear programs

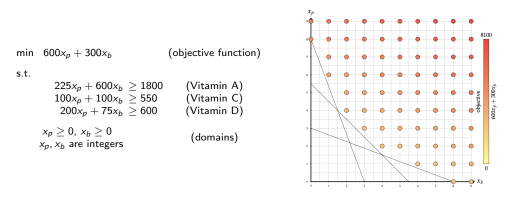
Each ILP has a corresponding LP relaxation, by allowing the variables to assume non-integers values:

- Each feasible solution in the ILP is also feasible in the LP relaxation, but not vice versa.
- The optimal solution of the LP relaxation is a lower/upper bound for the optimal solution of the ILP.

ILP - simple example: optimal meal problem

Consider the LP for solving the optimal meal problem, and suppose that we want to avoid meals that consist of partial portions of the two types of food.

For this goal we only need to set the domains of the variables x_p and x_b to be non-negative integers:



The ILP formulation has three optimal solutions that result in 2400 calories

$$\begin{cases} x_p = 2, x_b = 4 \\ x_p = 1, x_b = 6 \\ x_p = 0, x_b = 8 \end{cases}$$

ILP - NP-hard problem with simple formulation

MAXIMUM INDEPENDENT SET PROBLEM of a graph G = (V, E)

ILP formulation:

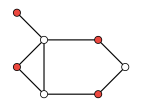
Let $V = \{v_1, v_2, ..., v_n\}$

Associate a **binary variable** to each vertex v_i : $x_i = \begin{cases} 1 & \text{if } v_i \text{ is in the independent set,} \\ 0 & \text{otherwise} \end{cases}$

Constraint: at most one vertex of each edge $(v_i v_j) \in E$ can be included in the independent set: $x_i + x_j \leq 1$

Objective: maximize the sum of all n binary x_k variables.

 $\begin{array}{ll} \max \sum_{k=1}^{n} x_k & (\text{objective function}) \\ \text{s.t.} & \\ x_i + x_j \leq 1 & \forall (v_i v_j) \in E \\ x_k \in \{0, 1\} & \forall v_k \in V & (\text{non-adjacent}) \\ \end{array}$



(MAXIMUM INDEPENDENT SET PROBLEM is NP-hard, but modeling it as an ILP is straightforward)

Quiz 1

1 Which of the following statements are true?

A linear program can always be solved in polynomial time in the number of variables and constraints.

An integer linear program cannot be solved in polynomial time in the number of variables and constraints.

X NP-hard problems can be solved by linear programming.

D Each feasible solution in the ILP is also feasible in its LP relaxation, but not vice versa.

The optimal solution of an ILP is a lower/upper bound for the optimal solution of its LP relaxation.



Matched genomes derived from balanced genomes

Balanced genomes
$$\mathbb{A}$$
 and \mathbb{B}

$$\begin{cases}
\mathcal{F}_{\star} = \mathcal{F}(\mathbb{A}) = \mathcal{F}(\mathbb{B}) \\
\mathcal{G}_{\star} = \mathcal{G}(\mathbb{A}) = \mathcal{G}(\mathbb{B}) \\
\text{for each family } f \in \mathcal{F}_{\star}, \ \Phi(f, \mathbb{A}) = \Phi(f, \mathbb{B})
\end{cases}$$

Transforming \mathbb{A} and \mathbb{B} into **matched** canonical genomes \mathbb{A}^{\ddagger} and \mathbb{B}^{\ddagger} :

for each family $f \in \mathcal{F}_{\star}$, determine which occurrence of f in \mathbb{A} matches each occurrence of f in \mathbb{B}

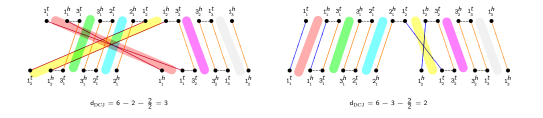
The number of common genes between any pair of matched genomes \mathbb{A}^{\ddagger} and \mathbb{B}^{\ddagger} is $n_* = |\mathcal{G}_*|$

Matched genomes derived from balanced genomes

Matched occurrences receive the same index in \mathbb{A}^\ddagger and in \mathbb{B}^\ddagger

The number of common genes between any pair of matched genomes \mathbb{A}^{\ddagger} and \mathbb{B}^{\ddagger} is $n_* = |\mathcal{G}_{\star}|$

Example: $\mathbb{A} = [132131]$ and $\mathbb{B} = [132]$ [131] with $n_* = 6$



DCJ distance of balanced genomes

 \mathfrak{M} : set of all possible pairs of matched canonical genomes obtained from balanced genomes $\mathbb A$ and $\mathbb B$

DCJ distance of $\mathbb A$ and $\mathbb B {:}$

$$\mathsf{d}_{\mathrm{DCJ}}(\mathbb{A},\mathbb{B}) = \min_{(\mathbb{A}^{\ddagger},\mathbb{B}^{\ddagger})\in\mathfrak{M}} \{\mathsf{d}_{\mathrm{DCJ}}(\mathbb{A}^{\ddagger},\mathbb{B}^{\ddagger})\}$$

NP-hard

Multi-relational graph of balanced genomes

Given two balanced genomes \mathbb{A} and \mathbb{B} , their multi-relational graph $MRG(\mathbb{A}, \mathbb{B}) = (V, E)$ is described as follows:

 V = V(ξ(A)) ∪ V(ξ(B)) : there is a vertex for each extremity of each gene in A and a vertex for each extremity of each gene in B

Each vertex v has a label $\ell(v)$, that corresponds to the gene extremity it represents.

2. $E = E_{\Gamma}(\mathbb{A}) \cup E_{\Gamma}(\mathbb{B}) \cup E_{\xi}$, where:

► Adjacency edges:
$$\begin{cases} E_{\Gamma}(\mathbb{A}) = \{uv : u, v \in V(\xi(\mathbb{A})) \text{ and } \ell(u)\ell(v) \in \Gamma(\mathbb{A})\} \\ E_{\Gamma}(\mathbb{B}) = \{uv : u, v \in V(\xi(\mathbb{B})) \text{ and } \ell(u)\ell(v) \in \Gamma(\mathbb{B})\} \end{cases}$$

• Extremity edges:
$$E_{\xi} = \{uv : u \in V(\xi(\mathbb{A})) \text{ and } v \in V(\xi(\mathbb{B})) \text{ and } \ell(u) = \ell(v)\}$$

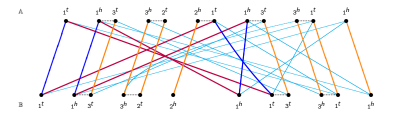
Vertices can have degree greater than two in the multi-relational graph:

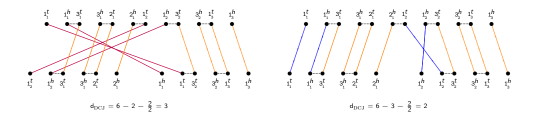
For each family $f \in \mathcal{F}_{\star}$, let $m_f = \phi(f, \mathbb{A}, \mathbb{B})$.

The number of extremity edges inciding in each vertex representing an extremity of an occurrence of f is m_f .

Multi-relational graph of balanced genomes

Example: $\mathbb{A} = [132131]$ and $\mathbb{B} = [132]$ [131]





Sibling-sets of $MRG(\mathbb{A}, \mathbb{B})$

 $f_{\mathbb{A}:i}$ and $f_{\mathbb{B}:k} \Rightarrow$ genes (occurrences of the same family $f \in \mathcal{F}_{\star}$) in genomes \mathbb{A} and \mathbb{B} .

Siblings: pair of extremity edges that connect $\begin{cases} f_{\mathbb{A}:i}^{h} \text{ to } f_{\mathbb{B}:k}^{h} \\ f_{\mathbb{A}:i}^{t} \text{ to } f_{\mathbb{B}:k}^{h} \end{cases}$

Sibling-set $S \subseteq E_{\xi}$ {is composed of pairs of siblings does not contain any pair of incident edges is a marketing

There is a bijection between pairs of (partially) matched genomes and sibling-sets of $MRG(\mathbb{A}, \mathbb{B})$:

- ▶ Denote by $\mathbb{A}^{\ddagger s}$ and $\mathbb{B}^{\ddagger s}$ the matched genomes corresponding to the sibling-set *S*
- ▶ If S is maximal, \mathbb{A}^{\ddagger_S} and \mathbb{B}^{\ddagger_S} are canonical (completely matched) genomes

Consistent decompositions of $MRG(\mathbb{A}, \mathbb{B})$

$$\mathbf{Consistent \ decomposition \ } D[S] \begin{cases} -\text{ is induced by a maximal sibling-set } S \\ -\text{ is the union of } S \text{ with all adjacency edges} \\ -\text{ covers all vertices of } MRG(\mathbb{A}, \mathbb{B}) \\ -\text{ is composed of cycles and paths:} \\ \mathbf{weight \ of \ } D[S]: \ w(D[S]) = |\mathcal{C}^D| + \frac{|\mathcal{P}_{\mathbb{AB}}^D|}{2} \\ \mathbf{d}_{\mathrm{DCJ}}(D[S]) = n_* - w(D[S]) \end{cases}$$

The DCJ distance of balanced \mathbb{A} and \mathbb{B} can then be computed by the following equation:

$$d_{DCJ}(\mathbb{A}, \mathbb{B}) = \min_{S \in \mathfrak{S}_{MAX}} \{ d_{DCJ}(D[S]) \} = n_* - \max_{S \in \mathfrak{S}_{MAX}} \{ w(D[S]) \},$$

where
$$\begin{cases} \mathfrak{S}_{MAX} \text{ is the set of all maximal sibling-sets of } MRG(\mathbb{A}, \mathbb{B}) \\ n_* \text{ is constant for any consistent decomposition} \end{cases}$$

If $d_{DCJ}(D[S]) = d_{DCJ}(\mathbb{A}, \mathbb{B})$, the consistent decomposition D[S] is said to be **optimal**.

Capped multi-relational graph $CMRG(\mathbb{A}, \mathbb{B})$

 $\mathsf{Example:} \ \mathbb{A} = \texttt{[132131]} \ \mathsf{and} \ \mathbb{B} = \texttt{[132]} \ \ \texttt{[\overline{1}31]} \ , \ \ p_* = \max\{\kappa(\mathbb{A}), \kappa(\mathbb{B})\} = 2, \ \ a_* = |\kappa(\mathbb{A}) - \kappa(\mathbb{B})| = 1$

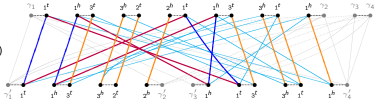
Add $2p_*$ cap extremities to each genome

Add the new adjacencies (including the a_* adj. between caps) to $E_{\Gamma}(\mathbb{A})$ and to $E_{\Gamma}(\mathbb{B})$

 $E_{\xi'}$: set of edges connecting cap extremities

 $E = E_{\Gamma}(\mathbb{A}) \cup E_{\Gamma}(\mathbb{B}) \cup E_{\xi} \cup E_{\xi'}$

Non-optimal capping

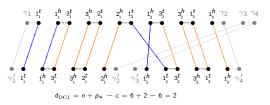


 $CMRG(\mathbb{A}, \mathbb{B})$ includes all possible cappings of each maximal sibling-set

Two distinct cappings of the maximal sibling-set composed of blue + orange edges:

 $\gamma_{1} \mathbf{1}_{1}^{t} \mathbf{1}_{1}^{h} \mathbf{3}_{1}^{t} \mathbf{3}_{1}^{h} \mathbf{2}_{1}^{t} \mathbf{2}_{1}^{h} \mathbf{1}_{2}^{t} \mathbf{1}_{2}^{h} \mathbf{3}_{2}^{t} \mathbf{3}_{2}^{h} \mathbf{1}_{3}^{t} \mathbf{1}_{3}^{h} \mathbf{1}_{3}^{h} \gamma_{2} \gamma_{3} \gamma_{4}$ $\gamma_{1}^{t} \mathbf{1}_{1}^{t} \mathbf{1}_{1}^{h} \mathbf{3}_{1}^{t} \mathbf{3}_{1}^{t} \mathbf{2}_{1}^{t} \mathbf{2}_{2}^{h} \gamma_{2}^{\prime} \gamma_{3}^{\prime} \mathbf{1}_{3}^{h} \mathbf{1}_{1}^{t} \mathbf{1}_{2}^{t} \mathbf{3}_{2}^{t} \mathbf{3}_{2}^{h} \mathbf{1}_{3}^{t} \mathbf{1}_{3}^{h} \gamma_{4}^{\prime}$ $d_{DC1} = n + p_{*} - c = 6 + 2 - 5 = 3$

Optimal capping



Capped consistent decompositions of $CMRG(\mathbb{A}, \mathbb{B})$

Capping-set $P \subseteq E_{\xi'}$: does not contain any pair of incident edges

Capped consistent decomposition Q[S, P]	(- is induced by a maximal sibling-set S and a maximal caping-set P
	- is the union of S with P and with all adjacency edges
	$-$ covers all vertices of $CMRG(\mathbb{A},\mathbb{B})$
) – is composed of cycles only:
	weight of $Q[S, P]$: $w(Q[S, P]) = C^Q $
	$d_{\mathrm{DCJ}}(Q[S,P]) = n_* + p_* - w(Q[S,P])$

 $\text{For each maximal sibling set } S: \qquad \mathsf{d}_{\text{DCJ}}(D[S]) = \min_{P \in \mathfrak{P}_{\text{MAX}}} \{ \mathsf{d}_{\text{DCJ}}(Q[S,P]) \}$

The DCJ distance of balanced genomes $\mathbb A$ and $\mathbb B$ can be computed by the following equation:

$$d_{\rm DCJ}(\mathbb{A},\mathbb{B}) = \min_{S \in \mathfrak{S}_{\rm MAX}, P \in \mathfrak{P}_{\rm MAX}} \{ d_{\rm DCJ}(Q[S,P]) \} = n_* + p_* - \max_{S \in \mathfrak{S}_{\rm MAX}, P \in \mathfrak{P}_{\rm MAX}} \{ w(Q[S,P]) \},$$
where
$$\begin{cases} \mathfrak{S}_{\rm MAX} \text{ is the set of all maximal sibling-sets of } CMRG(\mathbb{A},\mathbb{B}) \\ \mathfrak{P}_{\rm MAX} \text{ is the set of all maximal capping-sets of } CMRG(\mathbb{A},\mathbb{B}) \\ n_* \text{ and } p_* \text{ are constant for any capped consistent decomposition} \end{cases}$$

If $d_{DCJ}(Q[S, P]) = d_{DCJ}(\mathbb{A}, \mathbb{B})$, the capped consistent decomposition Q[S, P] is said to be **optimal**.

Quiz 2

1 Which of the following statements are true?

X The multi-relational graph is a collection of paths and cycles.

B A consistent decomposition of the multi-relational graph is a collection of paths and cycles.

C There is a bijection between consistent decompositions of $MRG(\mathbb{A}, \mathbb{B})$ and pairs of matched canonical genomes. $5 \rightarrow t$

There is a bijection between capped consistent decompositions of $CMRG(\mathbb{A}, \mathbb{B})$ and pairs of matched canonical genomes. 5, ? $\downarrow \rightarrow \ddagger$

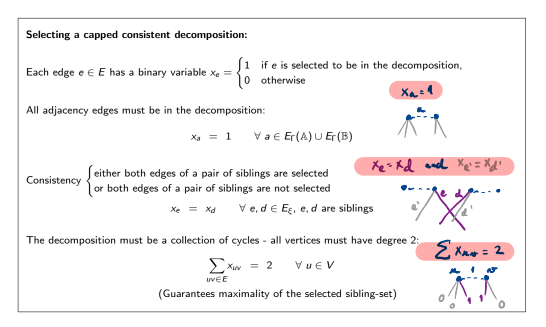
- 2 Given that Φ(f, A, B) is the number of occurrences of each family f in A and in B, the number of pairs of matched canonical genomes derived from balanced genomes A and B is...
- 3 The number of distinct caping sets is

A 2p*

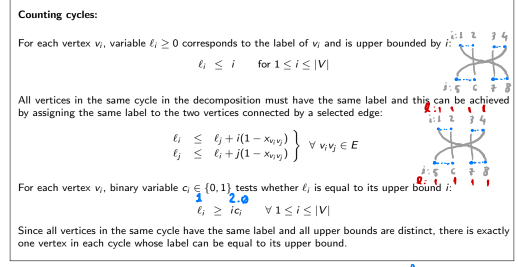
$$\bigwedge_{f \in \mathcal{F}_{\star}} \Phi(f, \mathbb{A}, \mathbb{B})!$$

$$\mathsf{B} \ 2\sum_{f\in\mathcal{F}_{\star}} \Phi(f,\mathbb{A},\mathbb{B})!$$

ILP to find an optimal capped consistent decomposition of $CMRG(\mathbb{A}, \mathbb{B})$



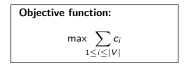
ILP to find an optimal capped consistent decomposition of $CMRG(\mathbb{A}, \mathbb{B})$



ILP to find an optimal capped consistent decomposition of $CMRG(\mathbb{A}, \mathbb{B})$

DCJ distance of balanced genomes

$$d_{\text{DCJ}}(\mathbb{A}, \mathbb{B}) = \min_{S \in \mathfrak{S}_{\text{MAX}}, P \in \mathfrak{P}_{\text{MAX}}} \{ d_{\text{DCJ}}(Q[S, P]) \}$$
$$= n_* + p_* - \max_{S \in \mathfrak{S}_{\text{MAX}}, P \in \mathfrak{P}_{\text{MAX}}} \{ w(Q[S, P]) \}$$
$$= n_* + p_* - \max_{S \in \mathfrak{S}_{\text{MAX}}, P \in \mathfrak{P}_{\text{MAX}}} \{ |\mathcal{C}^{Q[S, P]}| \}$$



Quiz 3

Computing the DCJ distance of balanced genomes:

Match the functions to their corresponding parts of the ILP!

Each adjacency edge is in the
decomposition(1)
$$(A)$$
 $\ell_i \leq \ell_j + i(1 - x_{\{v_i, v_j\}})$ $\forall \{v_i, v_j\} \in E$ Cycle labels of adjacent ver-
tices are the same(2)
 (B) (B)
 $\{u,v\} \in E$ $\sum_{\{u,v\} \in E} x_{\{u,v\}} = 2$ $\forall u \in V$ Sibling edges are only selected
together(3)
 (C) (C) $i \cdot c_i \leq \ell_i$ $\forall 1 \leq i \leq |V|$ A decomposition consists only
of simple cycles(4)
 (D) $x_a = 1$ $\forall a \in E_{\Gamma}(\mathbb{A}) \cup E_{\Gamma}(\mathbb{B})$ A cycle is only counted at the
vertex with the smallest label(5)
 (E) $x_e = x_d$ $\forall e, d \in E_{\xi}$ such that
 e and d are siblings

References

An Exact Algorithm to Compute the Double-Cut-and-Join Distance for Genomes with Duplicate Genes

(Mingfu Shao, Yu Lin, and Bernard M. E. Moret)

JCB, vol. 22, no. 5, pp 425-435 (2015)