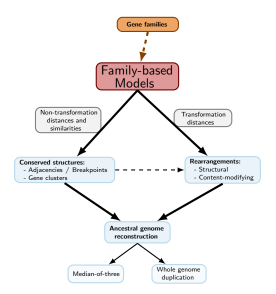
Topics of today:

- 1. Family-based \times Family-free setting
- 2. Family-free DCJ distance
- 3. Family-free DCJ-indel distance

Family-based setting



Are family assignments accurate?



Are family assignments accurate?

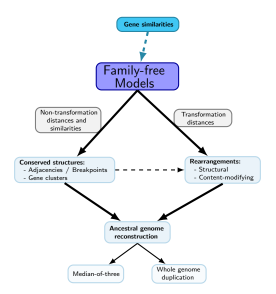


Family assignments are most of the time made automatically

• Even in the absence of errors, there may be ambiguities:

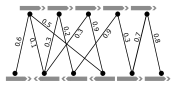


Alternative: family-free setting



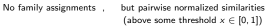
Family-free DCJ distance

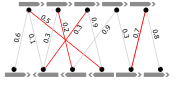
No family assignments , but pairwise normalized similarities (above some threshold $x \in [0, 1]$)



x = 0.1

Family-free DCJ distance

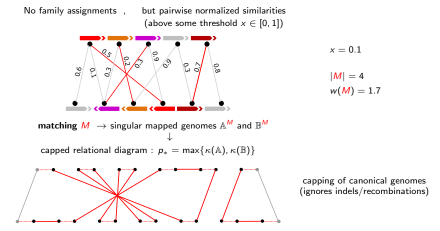




x = 0.1|M| = 4w(M) = 1.7

matching M

Family-free DCJ distance



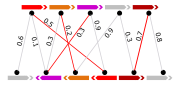
Taking the weights into consideration

Weighted DCJ distance of ma	appe	d genomes
$wd_{ ext{DCJ}}(\mathbb{A}^M,\mathbb{B}^M)=d_{ ext{DCJ}}(\mathbb{A}^M,\mathbb{B}^M)$	+	M - w(M)
$= p_* + M - \mathcal{C} $	+	M - w(M)

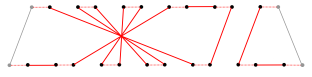
|M| - w(M) : penalizes edges of M with similarity < 1

genes that are not covered by M are simply ignored

 $|\mathbf{M}| = 4$ is maximal, $w(\mathbf{M}) = 1.7$

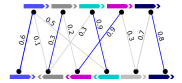


M	M	$d_{\rm DCJ}$	M - w(M)	$wd_{\rm DCJ}$
Μ	4	1	2.3	3.3

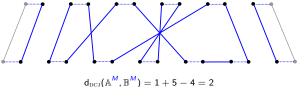


$$\begin{split} & \mathsf{d}_{\mathrm{DCJ}}(\mathbb{A}^{M},\mathbb{B}^{M}) = 1 + 4 - 4 = 1 \\ & \mathsf{wd}_{\mathrm{DCJ}}(\mathbb{A}^{M},\mathbb{B}^{M}) = 1 + 2.3 = 3.3 \end{split}$$

|M| = 5 is maximal, w(M) = 3.5

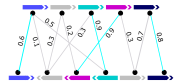


Μ	M	$d_{\rm DCJ}$	M - w(M)	$wd_{\rm DCJ}$
Μ	4	1	2.3	3.3
M	5	2	1.5	3.5
			1	'

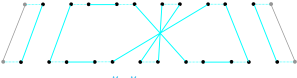


 $wd_{DCJ}(\mathbb{A}^{M}, \mathbb{B}^{M}) = 2 + 1.5 = 3.5$

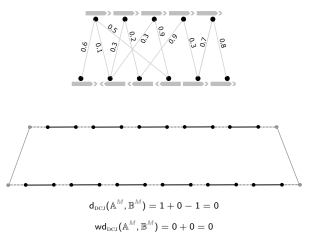
|M| = 4 is non-maximal, w(M) = 3.2



Μ	M	d_{DCJ}	M - w(M)	wd_{DCJ}
Μ	4	1	2.3	3.3
M	5	2	1.5	3.5
M	4	1	0.8	1.8
				1

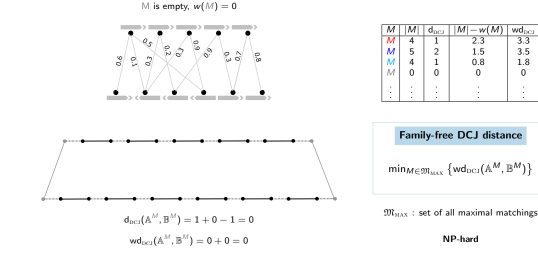


$$\begin{split} &\mathsf{d}_{\text{DCJ}}(\mathbb{A}^{\mathcal{M}},\mathbb{B}^{\mathcal{M}})=1+4-4=1\\ &\mathsf{wd}_{\text{DCJ}}(\mathbb{A}^{\mathcal{M}},\mathbb{B}^{\mathcal{M}})=1+0.8=1.8 \end{split}$$



M is empty, w(M) = 0

M	M	$d_{\rm DCJ}$	M - w(M)	wd_{DCJ}
Μ	4	1	2.3	3.3
M	5	2	1.5	3.5
Μ	4	1	0.8	1.8
Μ	0	0	0	0



3.3

3.5

1.8

0

Objective

We have a capped multi-relational graph, but here each gene can be potentially ignored:

 \Rightarrow each gene has an indel edge and ignoring a gene is done by selecting its indel edge

Besides variables and contraints from Shao-Lin-Moret, an extra constraint assures a maximal matching:

at least one of the vertices connected by an edge in the similarity graph must be chosen at most one of the corresponding indel edges in the capped MRG can be selected:

$$\begin{array}{l} x_a + x_b \leq 1 \\ x_a + x_b \leq 1 \end{array} \qquad \begin{cases} \forall \ a \in E_{\mathrm{D}}^{\mathrm{a}}, \ b \in E_{\mathrm{D}}^{\mathrm{b}}, \\ \text{siblings } d, \ e \in E_{\xi} \text{ and} \\ a \cap (d \cup e) \neq \emptyset \text{ and } b \cap (d \cup e) \neq \emptyset \end{array}$$

Weighted DCJ distance formula

 $wd_{DCJ}(\mathbb{A}^{M}, \mathbb{B}^{M}) = p_{*} + |M| - |C| + |M| - w(M) = p_{*} + 2|M| - |C| - w(M)$

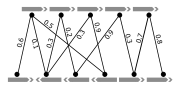
function:

$$\min \quad p_* + \sum_{e \in E_\xi} x_e - \sum_{1 \le i \le |V|} c_i - \frac{1}{2} \sum_{e \in E_\xi} w_e x_e$$

(one edge of the matching corresponds to a pair of edges in E_{ξ})

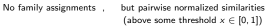
Family-free DCJ-indel distance

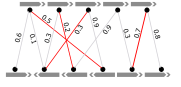
No family assignments , but pairwise normalized similarities (above some threshold $x \in [0, 1]$)



x = 0.1

Family-free DCJ-indel distance

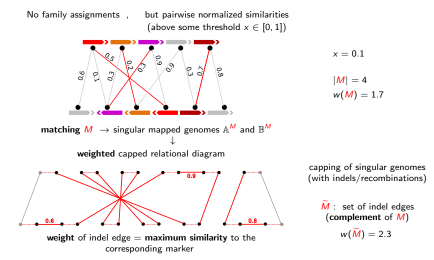




x = 0.1|M| = 4w(M) = 1.7

matching M

Family-free DCJ-indel distance



Taking the weights into consideration

Weighted DCJ-indel distance of mapped genomes

$$wd_{DCJ}^{ID}(\mathbb{A}^{M}, \mathbb{B}^{M}) = d_{DCJ}^{ID}(\mathbb{A}^{M}, \mathbb{B}^{M}) + [|M| - w(M)] + [w(\widetilde{M})]$$

$$= p_{*} + |M| - |\mathcal{C}| + \sum_{c \in \mathcal{C} \cup S} \lambda(c) + |M| - w(M) + w(\widetilde{M})$$

$$= p_{*} + |M| - |\mathcal{C}| + |\mathcal{C}^{r}| + |S| + \frac{\aleph}{2} + |M| - w(M) + w(\widetilde{M})$$

$$= p_{*} + |M| - |\mathcal{C}^{\tilde{r}}| + |S| + \frac{\aleph}{2} + |M| - w(M) + w(\widetilde{M})$$

|M| - w(M) : penalizes edges of M with similarity < 1

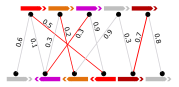
 $w(\widetilde{M})$: penalizes markers of \widetilde{M} with some similarity > 0

 C^r : set of indel-enclosing cycles

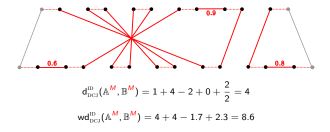
 $\mathcal{C}^{\tilde{r}}$: set of indel-free cycles

S : set of circular singletons

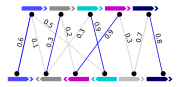
 $|\mathbf{M}| = 4$ is maximal, $w(\mathbf{M}) = 1.7$, $w(\widetilde{\mathbf{M}}) = 2.3$



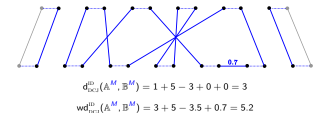
М	M	$d_{\rm DCJ}^{\rm ID}$	M - w(M)	$w(\widetilde{M})$	$wd_{\rm DCJ}^{\rm ID}$
М	4	4	2.3	2.3	8.6
			I		



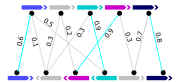
|M| = 5 is maximal, w(M) = 3.5, $w(\widetilde{M}) = 0.7$



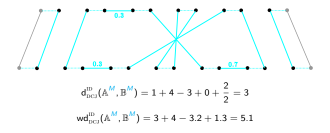
Μ	M	$d_{\rm DCJ}^{\rm ID}$	M - w(M)	$w(\widetilde{M})$	$wd_{\rm DCJ}^{\rm ID}$
М	4	4	2.3	2.3	8.6 5.2
M	5	3	1.5	0.7	5.2
	0	Ű	1.0	0.1	0.2



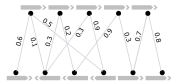
|M| = 4 is non-maximal, w(M) = 3.2, $w(\widetilde{M}) = 1.3$



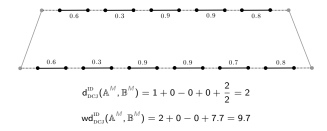
М	M	$d_{\rm DCJ}^{\rm ID}$	M - w(M)	$w(\widetilde{M})$	wd_{DCJ}^{ID}
Μ	4	4	2.3	2.3	8.6
M	5	3	1.5	0.7	5.2
M	4	3	0.8	1.3	5.1



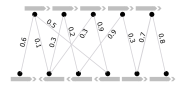
M is empty, w(M) = 0, $w(\widetilde{M}) = 7.7$



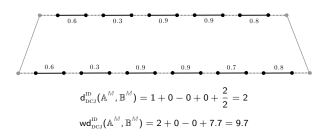
M	M	$d_{\rm DCJ}^{\rm ID}$	M - w(M)	$w(\widetilde{M})$	$wd_{\rm DCJ}^{\rm ID}$
Μ	4	4	2.3	2.3	8.6
M	5	3	1.5	0.7	5.2
M	4	3	0.8	1.3	5.1
M	0	2	0	7.7	9.7



M is empty, w(M) = 0, $w(\widetilde{M}) = 7.7$



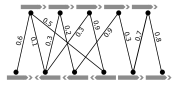
М	M	$d_{\rm DCJ}^{\rm ID}$	M - w(M)	$w(\widetilde{M})$	$wd_{\rm DCJ}^{\rm ID}$
Μ	4	4	2.3	2.3	8.6
Μ	5	3	1.5	0.7	5.2
Μ	4	3	0.8	1.3	5.1
Μ	0	2	0	7.7	9.7
			:	:	
	:	:	:	:	:



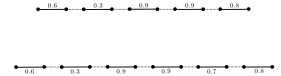
Family-free DCJ-indel distance
$\min_{M\in\mathfrak{M}}\left\{wd_{\scriptscriptstyle\mathrm{DCJ}}^{\scriptscriptstyle\mathrm{ID}}(\mathbb{A}^M,\mathbb{B}^M) ight\}$



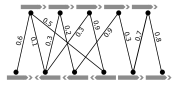
NP-hard



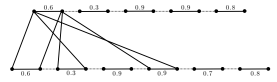
Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$



each marker has its weighted indel edge (weight = max. similarity)

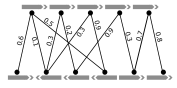


Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$

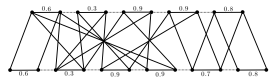


sibling weights are omitted

each marker has its weighted indel edge (weight = max. similarity)

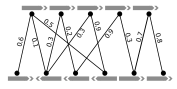


Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$

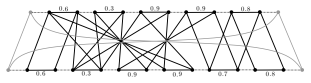


sibling weights are omitted

each marker has its weighted indel edge (weight = max. similarity)



Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$

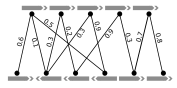


sibling weights are omitted

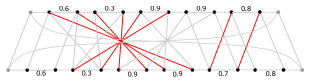
each marker has its weighted indel edge (weight = max. similarity)

add 2p* vertices (cap extremities)

link each cap extremity in genome $\mathbb A$ to each cap extremity in genome $\mathbb B$



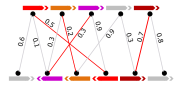
Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$



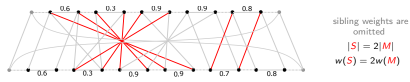
sibling weights are omitted

matching of extremity edges:

sibling set S (pairs of siblings)

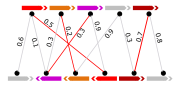


Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$

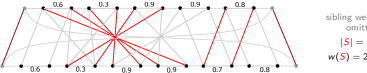


matching of extremity edges:

sibling set $S \rightarrow M$ (pairs of siblings)



Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$



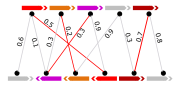


$$|\mathbf{S}| = 2|\mathbf{M}|$$
$$w(\mathbf{S}) = 2w(\mathbf{M})$$

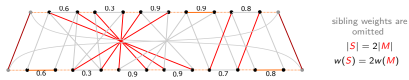
matching of extremity edges:

sibling set $S \rightarrow M$ (pairs of siblings)

maximal capping set P (covers all cap extremities)



Capped family-free relational diagram : $p_* = \max{\kappa(\mathbb{A}), \kappa(\mathbb{B})}$



matching of extremity edges:

sibling set $S \rightarrow M$ (pairs of siblings)

maximal capping set P (covers all cap extremities) $\begin{array}{l} \text{capped consistent decomposition} \\ Q[S, P] \end{array} \begin{cases} S \cup P \\ \text{all adjacency edges} \\ \text{complement} \quad \widetilde{S} \equiv \widetilde{M} \end{array}$

Optimization formula

DCJ-indel distance of a capped consistent decomposition

$$\mathsf{d}^{\text{\tiny ID}}_{\text{\tiny DCJ}}(\boldsymbol{\mathcal{Q}[S,P]}) = p_* + \boxed{\frac{|S|}{2}} - |\mathcal{C}^{\tilde{r}}| + |S| + \frac{\aleph}{2}$$

$$|S| = 2|M|$$

Optimization formula

DCJ-indel distance of a capped consistent decomposition

$$\mathsf{d}^{\text{ID}}_{\text{DCJ}}(\boldsymbol{Q}[\boldsymbol{S},\boldsymbol{P}]) = \boldsymbol{p}_* + \boxed{\frac{|\boldsymbol{S}|}{2}} - |\mathcal{C}^{\tilde{r}}| + |\boldsymbol{S}| + \frac{\aleph}{2}$$

$$|S| = 2|M|$$

w(S) = 2w(M)

Weighted DCJ-indel distance of a capped consistent decomposition

$$wd_{DCJ}^{ID}(Q[S, P]) = d_{DCJ}^{ID}(Q[S, P]) + \boxed{\frac{|S|}{2} - \frac{w(S)}{2}} + w(\widetilde{S})$$

$$= p_* + \frac{|S|}{2} - |\mathcal{C}^{\widetilde{r}}| + |S| + \frac{\aleph}{2} + \frac{|S|}{2} - \frac{w(S)}{2} + w(\widetilde{S})$$

$$= p_* + |S| - |\mathcal{C}^{\widetilde{r}}| + |S| + \frac{\aleph}{2} - \frac{w(S)}{2} + w(\widetilde{S})$$

Optimization formula

DCJ-indel distance of a capped consistent decomposition

$$\mathsf{d}^{\text{ID}}_{\text{DCJ}}(\boldsymbol{Q}[\boldsymbol{S},\boldsymbol{P}]) = \boldsymbol{p}_* + \boxed{\frac{|\boldsymbol{S}|}{2}} - |\mathcal{C}^{\tilde{r}}| + |\boldsymbol{S}| + \frac{\aleph}{2}$$

$$|S| = 2|M|$$

Weighted DCJ-indel distance of a capped consistent decomposition

$$wd_{DCJ}^{ID}(Q[S, P]) = d_{DCJ}^{ID}(Q[S, P]) + \boxed{\frac{|S|}{2} - \frac{w(S)}{2}} + w(\widetilde{S})$$

$$= p_* + \frac{|S|}{2} - |\mathcal{C}^{\widetilde{r}}| + |S| + \frac{\aleph}{2} + \frac{|S|}{2} - \frac{w(S)}{2} + w(\widetilde{S})$$

$$= p_* + |S| - |\mathcal{C}^{\widetilde{r}}| + |S| + \frac{\aleph}{2} - \frac{w(S)}{2} + w(\widetilde{S})$$

$$w(S)=2w(M)$$

Family-free DCJ-indel distance $\min_{S \in \mathfrak{S}, P \in \mathfrak{P}_{MAX}} \left\{ wd_{DCJ}^{ID}(Q[S, P]) \right\}$

 $\mathfrak{S}_{}$: set of sibling sets $\mathfrak{P}_{_{MAX}}$: set of maximal capping sets

Previous formulations:

DCJ distance of balanced genomes (Shao et al., 2014)

DCJ-indel distance of natural genomes (Bohnenkämper et al., 2020)

Selecting a consistent decomposition: (Shao *et al.*, 2014) $x_{a} = 1 \quad \forall \ a \in E_{\alpha}^{\mathbb{A}} \cup E_{\alpha}^{\mathbb{B}}$ $\sum_{uv \in E} x_{uv} = 2 \quad \forall \ u \in V$ $x_{e} = x_{d} \quad \forall \ e, d \in E_{\xi}, \ e, d \text{ are siblings}$

Previous formulations:

DCJ distance of balanced genomes (Shao et al., 2014)

DCJ-indel distance of natural genomes (Bohnenkämper et al., 2020)

Previous formulations:

- DCJ distance of balanced genomes (Shao et al., 2014)
- DCJ-indel distance of natural genomes (Bohnenkämper et al., 2020)

Counting singletons: (Bohnenkämper *et al.*, 2020) $\sum_{e \in E_{\text{ID}}^{k}} x_{e} - |k| \leq s_{k} \quad \forall \ k \in K$

Previous formulations:

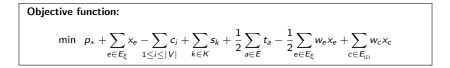
DCJ distance of balanced genomes (Shao et al., 2014)

DCJ-indel distance of natural genomes (Bohnenkämper et al., 2020)

Counting transition					
(Bohnenkämper <i>et al.</i>	, 202	0)			
	r _v r _{v'}	≤ ≥		$ \forall \ uv \in E_{\text{ID}}^{\mathbb{A}} \\ \forall \ u'v' \in E_{\text{ID}}^{\mathbb{B}} $	
	t _{uv} t _{uv}	≥ >	$\left. \begin{array}{c} r_{v} - r_{u} - (1 - x_{uv}) \\ r_{u} - r_{v} - (1 - x_{uv}) \end{array} \right\}$	$\forall uv \in E$	(C.08)
$\sum x_d$ -				$\forall \ \mathbf{a} \in \mathbf{E}^{\mathbb{A}}_{lpha}$	(C.09)
$d\!\in\!E_{ ext{ID}}^{\mathbb{A}}\ d\!\cap\!e\! e\!\!\neq\!\!\emptyset$					
	te	=	0	$\forall \ \mathbf{e} \in \mathbf{E} \setminus \mathbf{E}^{\mathbb{A}}_{\alpha}$	(C.10)

Weighted DCJ-indel distance formula

$$wd_{\text{DCJ}}^{\text{ID}}(Q[S,P]) = p_* + |S| - |\mathcal{C}^{\tilde{r}}| + |S| + \frac{\aleph}{2} - \frac{w(S)}{2} + w(\widetilde{S})$$



Available at https://gitlab.ub.uni-bielefeld.de/gi/gen-diff

Running times (or gap in %) for CPLEX with max. CPU time of 3h

Pairwise comparisons of Drosophila genomes

 \sim 13,000 genes per genome, distributed in 5-6 chromosomes

gene similarities obtained using FFGC (Doerr et al., 2018):

considering all similarities that are strictly greater than x = 0, the pairwise similarity graphs have an average of 11.2 and at most 95 connections per gene.

DIFF on similarity graphs of x = 0.3, with an average of 1.92 and at most 31 connections per gene:

pseudoobscura	sechellia	simulans	yakuba	busckii
0.76%	4,431.78s	109.60s	201.49s	540.19s
	163.12s	764.24s	5,782.73s	290.12s
		103.33s	146.88s	415.23s
			216.77s	115.54s
				153.36s
	P	0.76% 4,431.78s	0.76% 4,431.78s 109.60s 163.12s 764.24s	0.76% 4,431.78s 109.60s 201.49s 163.12s 764.24s 5,782.73s 103.33s 146.88s

(3h=10,800s):

 DING on OMA families with an average of 1.04 and at most 23 occurrences:

All comparisons finished very fast, ranging from 2 to 32 seconds.

Comparing DIFF and DING on CPLEX with max. CPU time of 3h

Balancing the number of multiple connections in both models:

Extending the connected components of similarity graphs to cliques

DING on families derived from similarity graphs extended cliques: All but one comparisons reached the time limit of 3h.

DIFF on similarity graphs with extended cliques (new edges received weight=0.3): Only one comparison reached the time limit of 3h, the others took 380 seconds on average.

Probable explanation:

There is a larger number of co-optimal solutions in the DCJ-indel distance of natural genomes. In the family-free DCJ-indel model the co-optimality is reduced by weights, allowing DIFF to converge faster.

 \Rightarrow Indeed, in a simulation in which the weights of all edges of the similarity graphs were set to 1, the running times of DIFF were much slower than those of DING for instances with the same number of multiple connections.

Gene homologies established by DIFF compared to Flybase

Flybase (flybase.org) established gene families (homolog gene sets) tor the three species

D. melanogaster D. simulans D. yakuba

Classification of pairs of homologous genes inferred for these three species with DIFF (for x = 0.3):

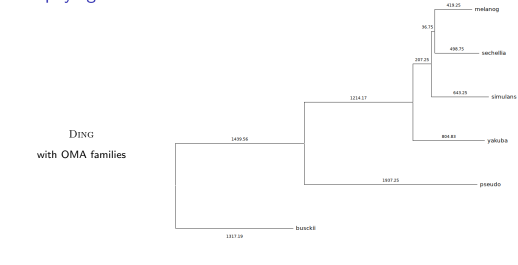
Match: (97.3%) both genes are in the same Flybase family;

New: (1.4%) both genes are not part of any Flybase family;

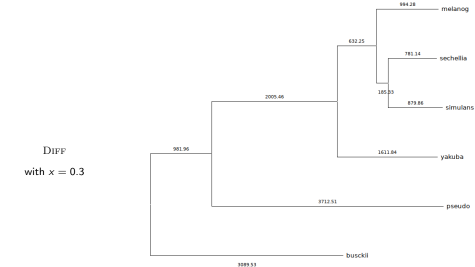
Extension: (1.1%) one of the two genes is not part of any Flybase family;

Mismatch: (0.2%) each gene is in a different Flybase family.

Inferred phylogenies

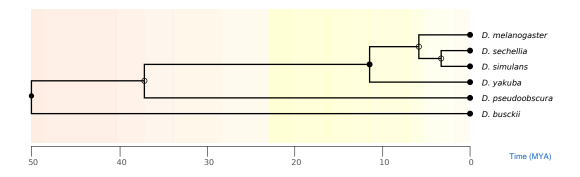


Inferred phylogenies



Reference phylogeny

TimeTree



References

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