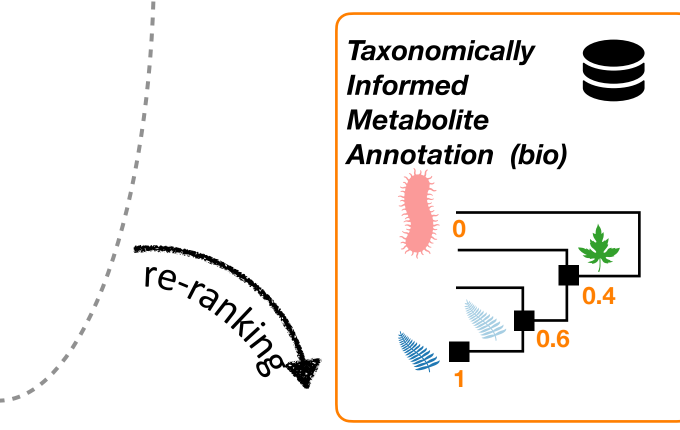


Feature ID	Spectrum	Biological source	Candidate structures	Score S <sub>1</sub>	Initial rank	
1	1 1	V.	INCHIKEY-1	0.55	1	
			INCHIKEY-2	0.53	2	
			INCHIKEY-3	0.50	3	
		•	INCHIKEY-4	0.45	4	
	•••					
1			INCHIKEY-9	0.00	9	



1			INCHIKEY-9	0.00	9	<b>*</b>	•	1	
Feature ID	Spectrum	Biological source	Candidate structures	Score S <sub>1</sub>		Candidate biological source		Combined score (S <sub>1</sub> + S <sub>2</sub> ) /2	Final rank
1			INCHIKEY-4	0.45	4		1.00	0.78	1
			INCHIKEY-1	0.55	1		0.60	0.58	2
			INCHIKEY-2	0.53	2	*	0.40	0.47	3
			INCHIKEY-9	0.00	9		0.60	0.30	4
•••	• • •	• • •	INCHIKEY-3	0.50	3	Southern Williams	0.00	0.25	5



Taxonomically Informed Metabolite Annotation (chemo) **Polyketides** 0.3 Terpenoids 0.6 Triterpenoids 1 Quassinoids

Feature ID	Spectrum	Biological source	Candidate structures	Score S <sub>1</sub>		Candidate biological source	Score S <sub>2</sub>	Combined score (S <sub>1</sub> + S <sub>2</sub> ) /2		Attributed chemical class	Candidate chemical class	Score S <sub>3</sub>	Combined score $(S_1 + S_2 + S_3)/3$	Final rank
1			INCHIKEY-4	0.45	4		1.00	0.78	1		Quassinoid	1.00	0.82	1
		INCHIKEY-1	0.55	1		0.60	0.58	2	OOH	Triterpenoids	0.60	0.58	2	
		INCHIKEY-2	0.53	2	*	0.40	0.47	3		Terpenoids	0.30	0.41	3	
			INCHIKEY-9	0.00	9		0.60	0.30	4	Quassinoid	Terpenoids	0.30	0.30	4
•••		•••	INCHIKEY-3	0.50	3	ATTION (U) AND	0.00	0.25	5		Polyketides	0.00	0.17	5