

# PHYLOGENETIC ANALYSIS OF PHOTOSYSTEMS I AND II



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Photosystem I (PS I) and Photosystem II (PS II) reaction centers include heterodimers of the gene products of *psaA* (PSI-A) and *psaB* (PSI-B) and of *psbA* (D1) and *psbD* (D2), respectively. Structural and sequence similarity exists among the heterodimer pair PSI-A and PSI-B, and D1 and D2 (Svensson et al., 1990; Golbeck and Bryant, 1991; Nitschke and Rutherford, 1991; Blankenship, 1992; Ruffle et al., 1992). Many sequences are now available for reaction center proteins of PS I and PS II. Golbeck and Bryant (1991) presented alignments of PSI-A and PSI-B for 7 taxa. Svensson et al. (1992) compiled 38 sequences of D1 (27 taxa) and 15 sequences of D2 (12 taxa) and showed that the sequences were strongly conserved. Ruffle et al. (1992) have shown relationships among the D1 and D2 proteins. In this paper, we present the first phylogenetic analysis of PSI-A and PSI-B, and extend the existing reaction center alignments of D1 and D2 to include several additional taxa.

Selected published protein sequences were retrieved from SWISS-PROT, PIR, and GenPept data bases using BLAST (Altschul et al., 1990). Additional sequences were obtained for *Porphyra purpurea* (M. Reith and J. Munholland, unpublished), *Cyanophora paradoxa* (V. L. Stirewelt and D. A. Bryant, unpublished), and *Zea mays* (Sopory et al., 1993, from Larrinua and McLaughlin, 1987). In total, 39 D1 and 11 D2 sequences of different genera, and 14 PSI-A and 14 PSI-B sequences, representing 13 genera each, were aligned. Alignments were done as described by Svensson et al. (1992), and Golbeck and Bryant (1991). The resulting data matrices were analyzed using PAUP (Phylogenetic Analysis Using Parsimony) version 3.1.1 (Swofford, 1993). All heuristics searches were replicated 100 times with random addition sequence and tree bisection-reconnection branch swapping. The options mulpars, steepest descent, collapse, and acctran optimization were selected. Gaps were treated as missing data. The amount of phylogenetic information in the parsimony analyses was estimated using the consistency, CI, (excluding uninformative characters) and retention, RI, indices. The trees were rooted by positioning the root along the branch connecting *Synechococcus* to the rest of the network.

Multiple alignments of D1, D2, PSI-A, and PSI-B sequences resulted in matrices of 360, 353, 804, and 743 positions, respectively; consensus sequences for each group of reaction center proteins are presented below (Complete data matrices are available upon request.).

## D1 (Photosystem II)

PROKARYOTES	MHTTLERRES	ASLWERFCSW	ITSTDERLYV	GWFGLMIPT	LLAATICFII	AFIAAPPVDI	DGIREPVAGS	LLYGNIIISG	AVVPSSNAIG	LHFYPIWEAA
ALGAE	AI	A E			T SV		S	T	I T	
MONOCOTS	AI	T G N			T SV		S		II T A	
DICOTS	AI	E G N			T SV		S		I T A	
PROKARYOTES	SLDENLYNGG	PYOLVVFHFL	IGICCYMGRE	WELSYRLGMR	PNICVAYSAP	VAATAVFLI	YPIGQGSFSD	GMP LGISGTF	NFMLVFOEH	NILMHPFHM
ALGAE		I F V			A F	S V			I	
MONOCOTS	V	E I L VA			A				I	
DICOTS	V	E I L VA			A	F			I	
PROKARYOTES	GVAGVFGGSL	FSAMHGSVLT	SSLVRETTEN	ESQNYGKFG	QEEETYNIVA	AHGYPGRLLF	OYASFNNRSR	LHFFLAAMPV	VGIWFTALGV	STMAFNLNGF
ALGAE			I	A N						
MONOCOTS			I	A E R						I
DICOTS			I	A E R						I
PROKARYOTES	NFNQSVVDSQ	GRVINTWADI	INRANLQMEV	MHERNAHNF	LDLASVESAP	VALTAPSING				
ALGAE					A	---	---			
MONOCOTS					A	---	---			
DICOTS					A	---	---			

D2 (Photosystem II)

PROKARYOTES	MTIAVGRAAA	VERGWFDVLD	DWLKRRDRFVF	IGWSGLLLEP	CAPMALGGWL	TGTTFTVTSWY	THGLASSYLE	GANFLTAVVT	SPADAFGHSL	LFLWGPEAQN
ALGAE	I TTYK EK	DA			SYF		T	C A S T	NSMA	V
MONOCOTS	L VPK E NDL	IM	R		YF			C A S T	NSLA	L
DICOTS	L FTK D NDL	IM	R		YF V			C A T	NSLA	L
PROKARYOTES	NLTRWFQGG	LMFFVALHGA	FGLIGFMLRQ	FEIARLVGIR	PYNAIAFSGP	IAVFVSVFLM	YPLGQSSWFF	APSEFGVAGIF	RFILFLQGFH	NWTLNPFPHM
ALGAE	DF C I	A		N	A	I		A		
MONOCOTS	DF C	T	A	L Q	S	I		A		
DICOTS	DF C	T		L Q		I		A		H
PROKARYOTES	GVAGILGGAL	LCAIHGATVE	NTLFEDGEDS	NTFRAFEP	AZETYSNVTA	NRFWSQIFGI	AFSNKRWLHF	FMLFVPTVGL	WNSSVGIVGL	ALNLRAYDFV
ALGAE			DGA	N	S	V			AI	
MONOCOTS			DGA	N		V			AI V	
DICOTS			DGA	N		V			AL V	
PROKARYOTES	SQELRAAEDP	EFETFYTKNI	LLNEGIRAWM	APQDOPHENF	IFPEEVLP	RG NAL				
ALGAE	I			A	V					
MONOCOTS	I			A	L					
DICOTS	I			A	L					

PSI-A (Photosystem I)

PROKARYOTES	MTISPPPEREA	-KARVVVDND	PVPTSFEKWA	KPGHFDRTLA	RGPKTTTMIW	NLHALAHDFD	THTSDLEDIS	RKIPSAHFQ	LAVVFIWLSG	MYPHGAKFSN
ALGAE	T	K VKIA K	E	S S K N		D	S	V Q	GII	
MONOCOTS	M RS P--	-EVKI R	K E R	S L K D		D	S E	V Q	SII	
DICOTS	I RS P--	-EVKIL R	K E R	S I K D		D	S E	V Q	SII L	
PROKARYOTES	YEAMLADPTG	IKPSAQVWVP	IVGOGILNGD	VGGGFHGIQI	TSCILFQLMRA	SGITNSFQLY	CTAIGGLVMA	GLMLFAGWFH	YHRAPKLEW	FQNVESMLNH
ALGAE	S		E	Q	L	SEL	T	A F	A	
MONOCOTS	S	G	E	R	S F I	SEL		A IF S	A A	SHD
DICOTS	S	G	E	R	F I	SEL		A F A	A A	D
PROKARYOTES	HLAGLLCLGS	LAWAGHQIHV	SLPINKLLDA	GVAARDIPLP	HEFILNPSLY	PELDPGFASG	LTPFFTLNWG	AYSDFLTFKG	GLNPVTGGLW	LSDTAHRHLA
ALGAE	G		S	DPQE		RD M AQ Y S SK	A	S E		
MONOCOTS		S	QF	DP E		RD L AQ Y S ER A		K AE S R	I	I
DICOTS		S	QF N	DP-E		RD L AQ Y S E A		K A R		T
PROKARYOTES	IAVLFIAGH	MYRTNWGIGH	SLREILEAHK	GPFTGEGHKG	LYEVLTSWH	AQLAINLAML	GSLTIIVAOH	MYAMPYPYL	ATDPTQLSL	FTHHMWIGF
ALGAE	V LV		M R		I		F S H	S		
MONOCOTS	I L		G D	QR	I	SL	T V H			
DICOTS	I L		G D	Q	I	L	V H			
PROKARYOTES	LIVGGAHGA	IFMVRDYDPA	MNQNNVLDLV	LRRDRAIISH	LNWVCIFLGF	HSFGLYIHD	TMRALGRPOD	MFSDTAIQLQ	PVFAQWVQNL	HTLAPGGTAP
ALGAE	C AG A		N Y L	I			S		I T F L	
MONOCOTS	A A		T TRY DL				S		AG RL	
DICOTS	A A		T TRY DL		A		S		I T A SA	
PROKARYOTES	NALA-TASYA	FGGDVVAVGG	KVAMMPIVLG	TADFMVHHIH	AFTIHVTVLI	LLKGVLFARS	SRLIPDKANL	GFRFPDGGP	RGGTCQVSGM	DHVFLGLFWM
ALGAE	- M G- ELG H		S	S						
MONOCOTS	G TTS SLTW	G EL	LL P	L						
DICOTS	G TAS SLTW	G S L	LL P	L						
PROKARYOTES	YNSLSIVIFH	FSWKMSQDVW	GTVAPDGTVS	HITCGNFAQS	AITINGWLRD	FLWAQASQVI	NSYGSALSAY	GLMFLGAHFV	WAFSLMFLFS	GRGYWQELIE
ALGAE	V		TAS G			S	Q	I		
MONOCOTS	I V		ISDQ V T		S		Q S	F		
DICOTS	I V		SISDQ V T		S		Q S	F		
PROKARYOTES	SIWVAHNKLG	VAPAIQPRAL	SIIQGRAVGV	AHYLLGGIAT	TWAFFLARIL	SVG				
ALGAE			T		S	I A				
MONOCOTS		T		T		I A				
DICOTS		T	V	T		I A				

PSI-B (Photosystem I)

PROKARYOTES	MATKFPKFSQ	GIAQDPTTRR	IMYGIATAHD	PETHDGMTEE	NLYQKIFASH	FGHLAIIFLW	TSGNLFHVAW	QGNFQZQVKD	PVNTRPIAHA	IWDPHFGKAA
ALGAE		D		S		Q		IT	LHV	P
MONOCOTS	ELR R	G	F	S DI	R N	Q		S IQ		P
DICOTS	LR R	G	F	S DI	R N	Q		Q		P
PROKARYOTES	VEAFTQAGAS	NPVNIAYSGV	YHWMYTIGMR	TNGELYQGAI	FLILASLAL	FAGWLHLOPK	FRPSLSWFKN	AESRLNHHLA	GLFGVSSLAW	AGHLVHVVAIP
ALGAE	RG G		Q	QD T SV	VSA F			S		T
MONOCOTS	RG A G		Q	L ED T L	F ST S	IG	WK	S		T
DICOTS	RG L		Q	L ED T L	AI S	I	WK V	S		T
PROKARYOTES	ESRGQHVQWD	NFLSTMHPA	GLAFFPTGNW	GVYAQNPDTA	SHVFGTSQGA	GTAILTFLGG	FHPQTESLWL	TDIAHHHLAI	AVLFIVAGHM	YRTNFGIGHS
ALGAE	R	VL Q	S	AA		E S Q		M		R
MONOCOTS	A EY R N	DVL Q	G LL Q	NL	SS N L T				FI LI	
DICOTS	A EY R N	DVL	G L Q	NL	SS L A				FI L	
PROKARYOTES	IKELNAHTK	VGGPFGAGHQ	GLYDTYNSL	HFQLGLALAS	LGVITSLVAQ	HMYSLPPYAF	IAQDHTTQAA	LYTHHQYIAG	FLMVGAFAHG	AIFLVRDYDP
ALGAE	L A D KP PS NL	K	I				I F		I C	FI
MONOCOTS	DL E TP P RL R K		I				S	F	I T	FI
DICOTS	M DL D IP P RL R K		I				A	F		I
PROKARYOTES	AQNKGVLAR	VLDHKEAIIIS	HLSWVSLFLG	FHTLGLYVHN	DVVVAFGTPE	KOILIEPVFA	OPIQAAGHCKL	LYGFDVLLSN	PDSIASTAMP	NYGNVWLPW
ALGAE	E	I	T		MQ		A	S	STVS	GO S---
MONOCOTS	E ED	M	A		ML	I	S T T	I	S TSGP	FN GR T---
DICOTS	E ED	M E	A		ML	I	S T S	I	S TSGP	FW GR S---
PROKARYOTES	LDAINSQTNS	LFLTICPGDF	LVHHAIALGL	HTTLLILVKG	ALDARGSKLM	PDKKDFGYAF	PCDGPGRGGT	CDISANDAFY	LAMFWMLNTI	GWVTFYHWHK
ALGAE	NNQ									
MONOCOTS	N V ENS									
DICOTS	N V ENS									
PROKARYOTES	HLGVWEGNVA	QFNESSTYLM	GWFRDYLWLN	SSQLINGYNP	FGTNNLSVMA	WMFLFGHLVW	ATGFMFLISW	RGYMQELIET	LVWAHERTPL	ANLVRNKKDP
ALGAE	TL				M S	F			K	
MONOCOTS	ITL QG S				M S				A	I
DICOTS	TL QG S				M S				A	I
PROKARYOTES	VALSIVQARL	VGLAHFSVGY	ILTYAAFLIA	STAGKFG						
ALGAE			F	S						
MONOCOTS			F	S						
DICOTS			F	S						

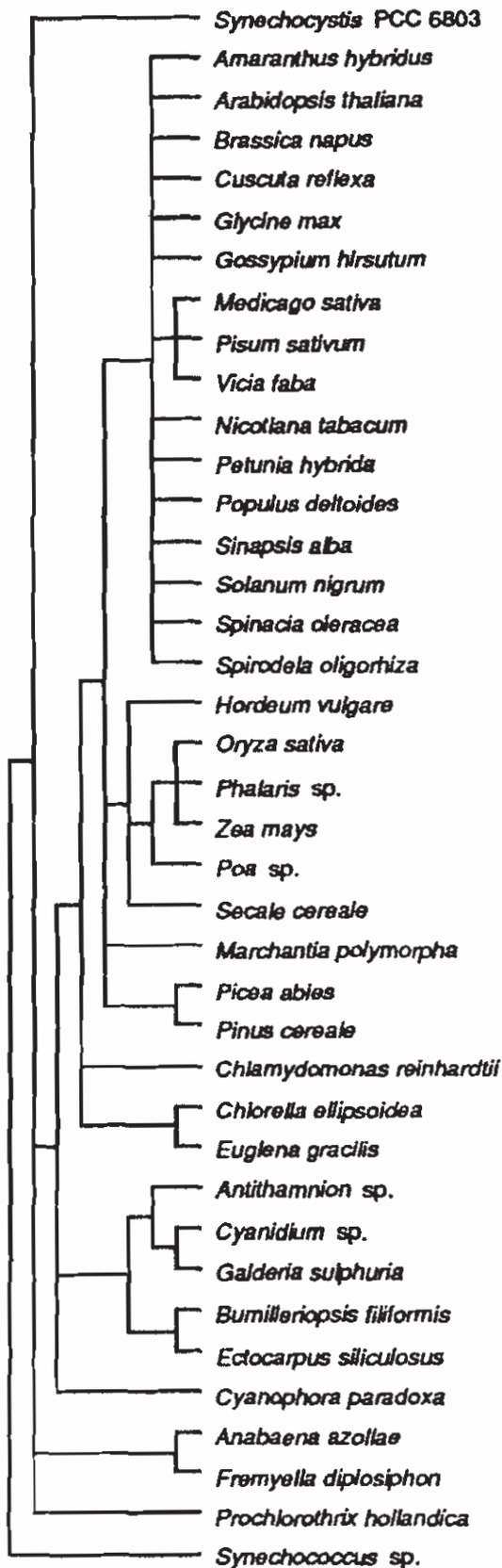


Fig. 1. Strict consensus of 1,660 minimal length 400-step trees derived from parsimony analysis of selected *psbA* (D1) sequences. CI=0.62, RI=0.69.

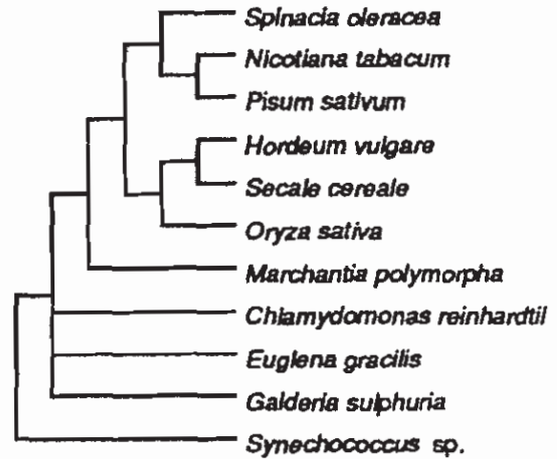


Fig. 2. Strict consensus of 4 minimal length 144-step trees derived from parsimony analysis of selected *psbD* (D2) sequences. CI=0.73, RI=0.62.

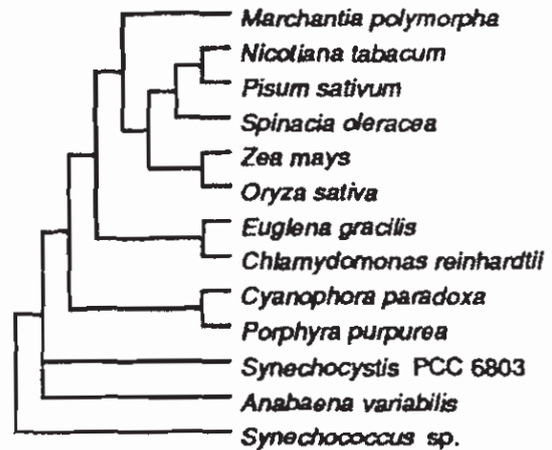


Fig. 3. Strict consensus of 2 minimal length 659-step trees derived from parsimony analysis of selected PSI-A sequences. CI=0.69, RI=0.69.

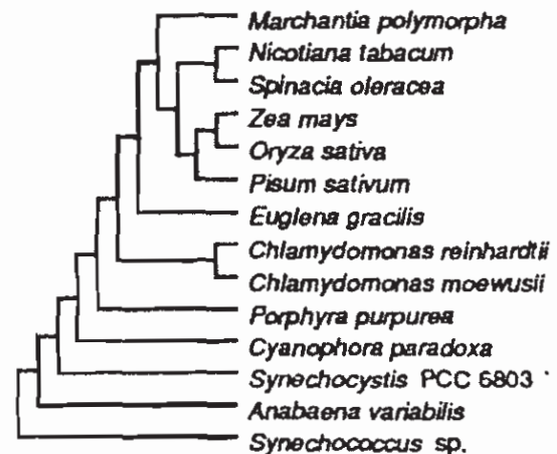


Fig. 4. Single maximally parsimonious 696-step tree derived from the analysis of selected PSI-B sequences. CI=0.72, RI=0.74.

The multiple alignment of each of the reaction center subunits was facilitated by the introduction of none (i.e., D2) or very few gaps (D1, one gap; PSI-B, three gaps; PSI-A, 16 gaps). Analyses of the D1, D2, and PSI-A data matrices resulted in 1,660, 4, and 2 maximally parsimonious topologies, whose strict consensus trees, with accompanying tree lengths, consistency and retention indices, are shown in Figs. 1, 2, and 3, respectively. A single maximally parsimonious tree obtained from the analysis of PSI-B sequences, is presented in Fig. 4. PS I and II reaction center proteins are highly conserved among the taxa examined. The inclusion of the *Pisum sativum* A2 PSI-A sequence (Lehmbeck et al., 1986) resulted in a single maximally-parsimonious tree of 1033 steps, 374 steps greater than the trees constructed without this sequence (CI excluding uninformative characters = 0.70; RI = 0.64). In this single PSI-A tree (not shown), this *Pisum* sequence arises as sister taxon to *Euglena* and away from the other *Pisum* and dicot sequences. In most cases, the evolutionary relationships inferred on the basis of PS I and II reaction center protein sequences and maximum parsimony are in agreement with traditional concepts of relationship. Generally the dicots and monocots each comprise a clade and are sistergroups.

**Remarks:** We have presented here the current consensus sequence of core PS II reaction center proteins (D1 and D2), and of core PS I reaction center proteins (PSI-A and PSI-B) based on published as well as unpublished sequences. Further, we have presented here the first phylogenetic analysis of PSI-A and PSI-B proteins and an extended analysis of D1 and D2 proteins. The phylogenetic trees, thus obtained, provide a hypothesis of the evolutionary relationship of the reaction centers of prokaryotes, cyanelle, rhodophytes, chlorophytes and other eukaryotes (gymnosperms, monocots and dicots).

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