RESEARCH ARTICLE



Open Access

Molecular evolution of *psbA* gene in ferns: unraveling selective pressure and co-evolutionary pattern

Lin Sen^{1,2}, Mario A Fares^{3,4}, Ying-Juan Su⁵ and Ting Wang^{2*}

Abstract

Background: The photosynthetic oxygen-evolving photo system II (PS II) produces almost the entire oxygen in the atmosphere. This unique biochemical system comprises a functional core complex that is encoded by *psbA* and other genes. Unraveling the evolutionary dynamics of this gene is of particular interest owing to its direct role in oxygen production. *psbA* underwent gene duplication in leptosporangiates, in which both copies have been preserved since. Because gene duplication is often followed by the non-fictionalization of one of the copies and its subsequent erosion, preservation of both *psbA* copies pinpoint functional or regulatory specialization events. The aim of this study was to investigate the molecular evolution of *psbA* among fern lineages.

Results: We sequenced *psbA*, which encodes D1 protein in the core complex of PSII, in 20 species representing 8 orders of extant ferns; then we searched for selection and convolution signatures in *psbA* across the 11 fern orders. Collectively, our results indicate that: (1) selective constraints among D1 protein relaxed after the duplication in 4 leptosporangiate orders; (2) a handful positively selected codons were detected within species of single copy *psbA*, but none in duplicated ones; (3) a few sites among D1 protein were involved in co-evolution process which may intimate significant functional/structural communications between them.

Conclusions: The strong competition between ferns and angiosperms for light may have been the main cause for a continuous fixation of adaptive amino acid changes in *psbA*, in particular after its duplication. Alternatively, a single *psbA* copy may have undergone bursts of adaptive changes at the molecular level to overcome angiosperms competition. The strong signature of positive Darwinian selection in a major part of D1 protein is testament to this. At the same time, species own two *psbA* copies hardly have positive selection signals among the D1 protein coding sequences. In this study, eleven co-evolving sites have been detected via different molecules, which may be more important than others.

Background

The photosynthetic oxygen-evolving photo system II is a unique biochemical system that is capable of oxidizing water molecules [1] and is responsible for producing the almost totality of oxygen on earth [2,3]. *psbA* gene, along with three other chloroplast (cp) genes, namely *psbB*, *psbC* and *psbD*, encodes the core proteins complex in the chlo-roplasts of ferns [4-7]. Precisely, *psbA* gene encodes D1 protein (also known as PsbA protein) in

²CAS Key Laboratory of Plant Germplasm Enhancement and Specialty Agriculture, Wuhan Botanical Garden, Chinese Academy of Sciences, Wuhan, China

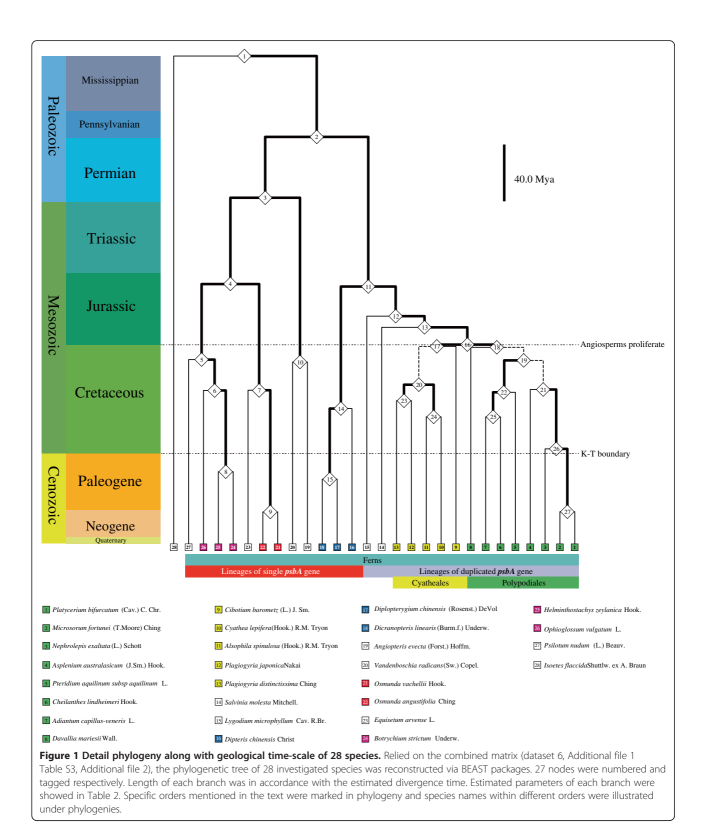
Full list of author information is available at the end of the article

ferns. Physical mapping and pastime sequencing unraveled a set of genome rearrangements around *psbA* gene in "higher" fern lineages [8-10]. Based on completely and partially sequenced fern pastimes [11-13], the *psbA* gene often locates either to inverted repeat regions (IRs) or to the large single copy region (LSC). Moreover, gene order around it is only found in one state in IRs (*rps7-psbA-trnH*), but in two possible arrangements in LSC (*trnK-psbA-trnH* or *matK-psbA-trnH*). Importantly, *psbA* gene duplicated when its location shifted from LSC to IRs [13]. This shift mainly affected four fern orders: Schizaeales, Salviniales, Cyatheales and Polypodiales, accounting for over 90% of extant fern diversity. Because both gene copies were preserved



© 2012 Sen et al.; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

^{*} Correspondence: tingwang@wbgcas.cn



since the duplication, we sought to investigate the selective pressures that may drive the evolution of this interesting gene.

Evolution after gene duplication has been a fundamental issue in evolutionary biology chiefly because of its direct link to the generation of novel functions and adaptations [14-16]. The opportunity for generating novel functions is, however, often balanced with the effects such duplications have on gene dosage [17-20]. The photosynthesis environment changed mainly as a result of the emergence of flowering plants [21], and as the open sunny ground transformed into the closed shadowy canopy. For this reason, the biodiversity of other vascular plants largely decreased, and some lineages underwent extinction [22-24]. Remarkably, rather than the result of being adept at holding on in the face of angiosperm domination, the leptosporangiate ferns may have the ability to capitalize upon it [25]. The underlying molecular functioning of this ability remains a mystery for evolutionary biologists, and the present study sheds some light on it. The colonization by these species of canopies angiosperm-dominated light environments sparks the idea that such extraordinary diversification may have been fuelled by functional innovation following *psbA* duplication. To disclose this obscurity, we performed a comprehensive analyze of the action of natural selection following the duplication in psbA.

Coevolving sites are non-independent amino acid sites during the course of protein evolution [26,27]. Among the different sites with physical or functional relationships in one protein, one mutation is likely to trigger the corresponding mutations at related sites [28]. Natural selection theory and nearly neutral theory have different explanations for the mechanism of these dynamic changes [29-31]. Despite the debates on the underlying mechanism, the compensatory mutations among the interdependent amino acid sites provide an important approach to understand protein structure and function [32]. The identification of coevolving amino acid sites will not only play a prominent role in the annotations of the function of D1 protein, but will also reflect the evolutionary pattern of the particular protein.

To achieve a better understanding on the evolutionary biology of *psbA* genes in ferns, we focused on three aims: 1) identifying gene order states around *psbA* genes in 11 fern orders; 2) detecting the selective constraints in consequence of *psbA* gene duplication; and 3) unraveling the co-evolution pattern of D1 protein among fern lineages.

Results

Sampling and sequencing

Mainly based on Smith's system [33] and Lehtonen's recent study [34], we sampled more species in Order Polypodiales (5 species) and Cyatheales (4 species), which have significantly high current fern diversity on earth, and we selected at least one species from the 11 orders (Figures 1, 2, and Table 1). To sum up, these 27 *psbA* sequences represented 11 orders and 14 families (100% of fern orders and 40% of families) providing reasonable coverage of the most taxon-rich lineage. All the sequenced 20 coding regions have 1059nt, coding 353 amino acid residues. Along with their stop code, 20 currently determined *psbA* sequences were uploaded to GenBank (Accession: JQ684679 - JQ684698, Table 1).

Since the analysis of molecular adaptive evolution strictly rejects the termination codon (i.e. TAA, TGA, TAG) in a sequence [35], coding sequences in dataset 1, 2 and 3 showed in Additional file 1 Table S3 were applied in the investigation of selective pressures.

Phylogenetic analysis

The reconstruction of phylogenetic trees is non-sensitive to the stop codes [36-38]; moreover, intergenic sequence of *psbA-trnH* was widely accepted as good indicator for barcode of land plant [39,40]. Coding sequences along with intergenic regions of *psbA-trnH* in dataset 4, 5 and 6 (Additional file 2)were utilized for the reconstruction of the phylogenetic structures for better resolution (Table 2).

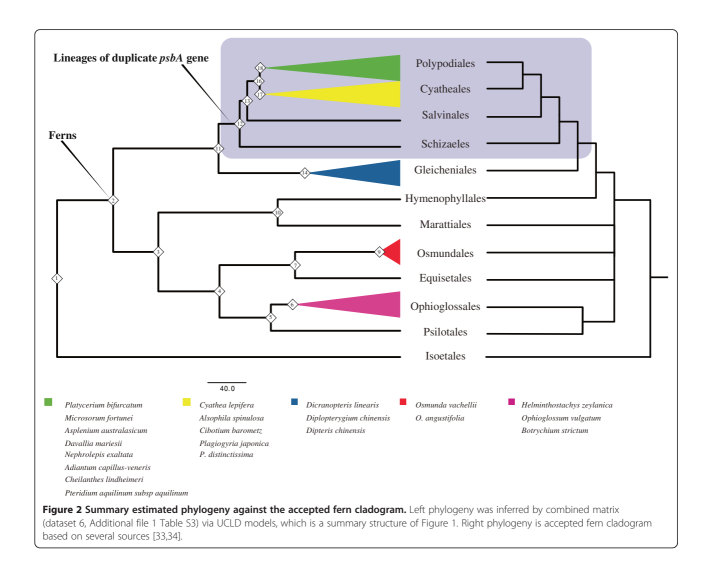
In accordance with the phylogenetic structure in previous documents and present analyses (Figure 2), we found that the combined dataset 6 might not be an accurate indicator for the reconstruction of fern phylogenetic structure under geological timescale. Even though the currently gained tree partially coincided with other relevant results from Smith's system [33] and Lehtonen's conclusion [34], we could found several disagreements.

Firstly, species from the same Orders were in the same clades (Figure 2): 1) Order Ophioglossales in node 6; 2) Order Osmundales in node 9; 3) Order Gleicheniales in node 14; 4) Order Cyatheales in node 17 and 5) Order Polypodiales in node 18. Secondly, species from the same Families were in the same sub-clade (Figure 1): 1) Family Ophioglossaceae in node 6; 2) Family Osmundaceae in node 9; 3) Family Gleicheniaceae in node 14; 4) Family Plagiogyriaceae in node 23; 5) Family Polypodiaceae in node 27. Thirdly, species from the same genera were together: 1) Genus *Osmunda* in node 9; 2) Genus *Plagiogyria* in node 23. Fourth, most esti-mated node ages had significant posterior probabilities (Table 3).

The current phylogenetic tree along with timescale was not accurate according to other published literature [25]. i) Partial structure was not the same with the known phylogeny (Figure 2): the Order Hymenophyllales was near to Marattiales in our study rather than Gleicheniales in well-known tree. ii) Several best estimated node ages were not precise, such as node 21 and 22. iii) Four nodes had low posterior possibilities (e.g. node 19, 20, 21 in Table 4).

Gene order around *psbA* gene in ferns

Results of the PCR amplification indicate that gene order around *psbA* gene in fern species could be mainly classified into three types: *trnK-psbA-trnH*,



matK-psbA-trnH and rps7-psbA-trnH. When universal primers (Additional file 1 Table S1) from conserved sequence of trnK and psbA were applied in PCR systems, five species (Botrychium strictum, Ophioglossum Helminthostachys zeylanica, vulgatum, Osmunda angustifolia and Osmunda vachellii) have positive reaction with the right sizes of the fragments. Four species (Vandenboschia radicans, Dipteris chinensis, Diplopterygium chinensis and Dicranopteris linearis) showed positive results when the primers were from conserved sequence of matK and psbA (Additional file 1 Table S2). Other eleven species (Lygodium microphyllum, Salvinia molesta, Plagiogyria distinctissima, Plagiogyria japonica, Cibotium barometz, Cyathea lepifera, Nephrolepis exaltata, Davallia mariesii, Asplenium australasicum, Microsorum fortune and Platycerium bifurcatum) showed positive results when the primers were from conserved sequence of rps7 and psbA (Additional file 1 Table S2).

Selective pressure in different psbA genes

As showed in Figures 1 and 2, we tested the action of natural selection in D1 protein using three datasets (Dataset 1, 2 and 3 in Additional file 1 Table S3) comprising both presently determined sequences (20 species in Table 1) and sequences retrieved from GenBank (7 species in Additional file 1 Table S4).

We can found subtle positive selection signals in *psbA* genes from fern species (Figure 3, Table 4 and Additional file 1 Table S6). However, when the species were divided into two groups as single and duplicated *psbA* genes, only those from single copy harbored alike signals (Table 4 and Additional file 1 Table S6). As showed in Table 4 and Additional file 1 Table S6, three codons (site 4, 91 and 155) were found as positively selected sites from dataset 1 and none from dataset 2 throughout different mathematics models.

For the detection of positive selection we used nested maximum likelihood models allowing for variation in

Order	Family	Genus	Species	Location*	GenBank Accession
Polypodiales	Polypodiaceae	Platycerium	P. bifurcatum (Cav.) C. Chr.	WBGCAS	JQ684679
		Microsorum	M. fortunei (T. Moore) Ching	WBGCAS	JQ684680
	Aspleniaceae	Asplenium	A. australasicum (J.Sm.) Hook.	WBGCAS	JQ684681
	Davalliaceae	Davallia	D. mariesii Wall.	WBGCAS	JQ684682
	Lomariopsidaceae	Nephrolepis	<i>N. exaltata</i> (L.) Schott	WBGCAS	JQ684683
Cyatheales	Cyatheaceae	Cyathea	C. lepifera (Hook.) R.M. Tryon	WBGCAS	JQ684684
	Cibotiaceae	Cibotium	C. barometz (L.) J. Sm.	WBGCAS	JQ684685
	Plagiogyriaceae	Plagiogyria	P. japonica Nakai	FLBG	JQ684686
		Plagiogyria	P. distinctissima Ching	WBGCAS	JQ684687
Salviniales	Salviniaceae	Salvinia	S. molesta Mitchell.	FLBG	JQ684688
Schizaeales	Lygodiaceae	Lygodium	L. microphyllum Cav. R.Br.	FLBG	JQ684689
Gleicheniales	Gleicheniaceae	Dicranopteris	D. linearis (Burm.f.) Underw.	FLBG	JQ684690
		Diplopterygium	D. chinensis (Rosenst.) DeVol	FLBG	JQ684691
		Dipteris	D. chinensis Christ	WBGCAS	JQ684692
Hymenophyllales	Hymenophyllaceae	Vandenboschia	V. radicans (Sw.) Copel.	WBGCAS	JQ684693
Osmundales	Osmundaceae	Osmunda	<i>O. vachellii</i> Hook.	SCBGCAS	JQ684694
		Osmunda	O. angustifolia Ching	SCBGCAS	JQ684695
Ophioglossales	Ophioglossaceae	Helminthostachys	H. zeylanica Hook.	SCBGCAS	JQ684696
		Ophioglossum	O. vulgatum L.	FLBG	JQ684697
		Botrychium	B. strictum Underw.	WBGCAS	JQ684698

Table 1 Plant materials and sampling locations in this study

*WBGCAS is short for Wuhan Botanical Garden, Chinese Academy of Sciences, Wuhan, Hubei, China. FLBG stands for Shenzhen Fairy Lake Botanical Garden, Shenzhen, Guangdong, China. SCBGCAS means South China Botanical Garden, Chinese Academy of Sciences, Guangzhou, Guangdong, China.

the ratio of non-synonymous to synonymous substitutions rates (d_N/d_S) across codons, as implemented in PAML and Selecton [41,42], and three models in Datamonkey as well [43,44]. For each dataset, one Likelihood Ratio Test (LRT) was performed for d_N/d_S heterogeneity across codons (M0/M3 test). The Discrete model (M3) fitted the data significantly better than the one-ratio model (M0), which suggested the existence of significant variation in selective constraints among codons (Table 4, Additional file 1 Table S5). Further, four LRTs were performed for the positively selected codons: M1a/M2a, M7/M8, M8a/M8 and M8a/Mechanistic empirical model (MEC). In order to take the differences between amino-acid replacement rates into account, MEC model employed a cpREV matrix to expand a 20 by 20 amino acid replacement rate matrix into a 61 by 61 sensecodon rate matrix [45]. By this means, a position with radical replacements will obtain a higher Ka value than a position with more moderate replacements (Figure 3). Log-likelihood values and estimates of parameters under various models were given in Table 4, and the likelihood ratio tests in Additional file 1 Table S5. In dataset two, alternative models (M2a and M8), permitting $\omega > 1$, failed to detect positively selected codons and showed non-significance against their null tests (M1a and M7). In M2a, though the estimated value ($\omega_2 = 34.2681$) is

Table 2 Selective pressures among <i>psbA</i> gene sequences	examined via SLAC and FEL models
--	----------------------------------

Dataset**	Model			Numbers and p	Numbers and positions of positively/negatively selected site*			
		p = 0.01	p = 0.05	<i>p</i> = 0.1	p=0.15	p=0.2	p=0.25	
One	SLAC	0/22	0/78	0/103	0/137	0/141	1(4)/144	
	FEL	0/69	1(4)/126	1(4)/150	1(4)/171	1(4)/198	1(4)/210	
Two	SLAC	0/4	0/24	0/42	0/71	0/75	0/79	
	FEL	0/22	1(155)/56	2(155, 352)/81	3(4, 155, 352)/93	4(4, 155, 350, 352)/120	5(4, 53, 155, 350, 352)/139	

*The positions of positively selected codons were showed in brackets. *p* value stands for the statistical significant levels of the models, which is preset before the data manipulation.

** The contexts of the dataset were introduced in Additional file 1 Table S3.

Table 3 The estimated parameters among phylogenetic tree via BEAST software

Node	Estimated	Posterior	Geological timescale		
No.*	Diverge Time**	probability***	Era	Period	
1	359.53	100	Paleozoic	Mississippian	
2	299.9	100	Paleozoic	Permian	
3	255.03	98.33	Paleozoic	Permian	
4	191.39	100	Mesozoic	Jurassic	
5	135.7	100	Mesozoic	Cretaceous	
6	112.95	100	Mesozoic	Cretaceous	
7	113.11	99.05	Mesozoic	Cretaceous	
8	53.02	100	Cenozoic	Paleogene	
9	23.38	100	Cenozoic	Neogene	
10	133.94	99.81	Mesozoic	Cretaceous	
11	189.57	93.15	Mesozoic	Jurassic	
12	167.73	100	Mesozoic	Jurassic	
13	159.5	99.91	Mesozoic	Jurassic	
14	99.6	100	Mesozoic	Cretaceous	
15	47.54	100	Cenozoic	Paleogene	
16	146.76	100	Mesozoic	Cretaceous	
17	145.45	100	Mesozoic	Cretaceous	
18	144.76	100	Mesozoic	Cretaceous	
19	135.12	83.86	Mesozoic	Cretaceous	
20	117.2	47.62	Mesozoic	Cretaceous	
21	113.65	68.39	Mesozoic	Cretaceous	
22	111.6	100	Mesozoic	Cretaceous	
23	105.41	100	Mesozoic	Cretaceous	
24	93.51	100	Mesozoic	Cretaceous	
25	93.48	100	Mesozoic	Cretaceous	
26	69.86	100	Cenozoic	Paleogene	
27	23.55	100	Cenozoic	Neogene	

* As showed in Figure 1 and Figure 2, all the nodes in the phylogenetic tree were numbered and marked.

** Diverge time of each branch was estimated in BEAST software and listed accordingly. The geological time scale (era and period) were inferred from the estimated diverge time of each nodes.

**** Posterior probabilities (PP) of every node were inferred via BEAST. Four of them were lower than 95%: 1) node 11; 2) node 19; 3) node 20; and 4) node 21.

greater than one, no codon ($p_2 = 0$) belongs to this kind of ω ; and the estimated value for ω was not greater than 1 in M8. Meanwhile, M8 significantly outperformed M7 in both datasets; M2a outperformed M1a as well (Additional file 1 Table S5). Collectively, we can conclude three main points under nested maximum likelihood models (Table 4): 1) subtle positive selection signal was detected in D1 protein from fern species; 2) codons within D1 protein encoded by duplicated *psbA* genes were mainly under negative selection; 3) several codons encoded by single *psbA* gene might have undergone adaptive evolution.

The random-site models, SLAC, FEL, REL and MEC methods were employed to examine the adaptive evolution of the D1 protein in eleven fern orders [41-44]. For dataset one (Table 2 and Additional file 1 Table S6), one positively selected codon (91 L) has been detected via maximum likelihood methods (Table 4), none via SLAC model, one (4 T) via FEL model (p = 0.05), two (4 T and 155 T) via REL model (PP > 95%). Our results indicate that 96.99% codons are highly conserved, 2.69% evolve neutrally, and a few (0.31%) are positively selected with $\omega > 1$ (Table 4). In the evolution of singleton *psbA* gene, positive selection at a handful of codons has played an important role in the evolutionary dynamics. On the contrary, the majority of codons are under negative selection in dataset two, while a small part is under neutral selection. No positively selected codon was found in the four fern lineages (Order Schizaeales, Salviniales, Cyatheales and Polypodiales). Our results showed that most of the sites are under purifying selection, while a small part is under neutral selection. No sites in the duplicate *psbA* gene were found under positive selection, while 4 have statistical significance in the singleton.

The selective relaxation in duplicated *psbA* genes could be observed in the adaptive selection analyses based on single likelihood ancestor counting (SLAC) and fixed effects likelihood (FEL) methods. Results in Table 2 showed a decrease of selective pressure in duplicated *psbA* genes (Table 2). For instance, 22 sites in dataset 1 were under selection while only 4 in dataset 2 via SLAC (p = 0.01). In accordance with FEL, 69 sites in dataset 1 were under selection while only 22 in dataset 2 (p = 0.01). Less codons from dataset 2 against dataset 1 were under either positive or negative selection (p = 0.01). On the other hand, no codon was identified as positive selection in dataset 2 under REL (Additional file 1 Table S6), while two amino acid sites (4 T and 155 T) were identified in dataset 1.

Co-evolutionary pattern among D1 protein

Multiple amino acid sites were involved in the coevolutionary network within D1 protein. CAPS indicated that one co-evolution pair was located between the N-terminal and α helix (Site 4 and 71) [46]. In Datamonkey [43,44], two kinds of methods for detecting coevolution (one parent and two parents) indicated that other pair has undergone co-evolution (Site 19 and 350). Moreover, the results from InterMap3D [47] indicated that ten pairs have undergone coevolution processes during their evolution dynamics (Table 5).

Discussion

Phylogenetic marker

It is a great challenge to develop DNA barcodes for land plants [40,48]. Kress and Erickson (2007) recommended

Models	Ts/Tv	np	e	Estimated value of parameters	Positively selected codons**
Dataset one*					
M0: One ratio	4.38	22	-3906.05	$\omega = 0.0184$	None
M1a: Near neutral	4.55	23	-3858.84	$p_0 = 97.14\%$, $\omega_0 = 0.00618$; $p_1 = 2.86\%$, $\omega_1 = 1$	Not allowed
M2a: Positive	4.55	25	-3858.83	$p_0 = 97.14\%$, $\omega_0 = 0.00618$; $p_1 = 2.86\%$, $\omega_1 = 1$; $p_2 = 0\%$, $\omega_2 = 79.26$	91 L (97.2%)
M3: Discrete	4.60	26	-3854.64	$p_0 = 96.31\%, \ \omega_0 = 0.00488;$ $p_1 = 3.39\%, \ \omega_1 = 0.5646;$ $p_2 = 0.3\%, \ \omega_2 = 3.9847$	91 L (99.8%)
Μ7: β	4.48	23	-3865.36	<i>p</i> =0.0267, <i>q</i> =0.614	Not allowed
Μ8: β &ω > 1	4.53	25	-3857.78	$p_0 = 97.84\%, p = 0.01575, q = 0.46508;$ $p_1 = 2.15\%, \omega = 1.08788$	91 L (99.7%)
Dataset two*					
M0: One ratio	5.97	29	-3093.72	$\omega = 0.02403$	None
M1a: Near neutral	6.07	30	-3061.96	$p_0 = 96.57\%$, $\omega_0 = 0.00127$; $p_1 = 3.43\%$, $\omega_1 = 1$	Not allowed
M2a: Positive	5.80	32	-3061.96	$p_0 = 96.57\%$, $\omega_0 = 0.00127$; $p_1 = 3.43\%$, $\omega_1 = 1$; $p_2 = 0$, $\omega_2 = 34.2681$	347E (58.3%)
M3: Discrete	5.71	33	-3060.47	$p_0 = 46.14\%$, $\omega_0 = 0$; $p_1 = 49.52\%$, $\omega_1 = 0$; $p_2 = 4.31\%$, $\omega_2 = 0.61912$	None
Μ7: β	5.75	30	-3064.99	p=0.00981, q=0.19983	Not allowed
Μ8: β &ω > 1	5.80	32	-3061.93	$p_0 = 96.61\%$, $p = 0.01744$, $q = 2.58197$; $p_1 = 3.38\%$, $\omega = 1$	347E (81.6%), 349P (61.5%), 353 G (54.4%)
Dataset three*					
M0: One ratio	4.66	47	-5742.02	$\omega = 0.01934$	None
M1a: Near neutral	4.87	48	-5654.74	$p_0 = 96.39\%$, $\omega_0 = 0.00538$; $p_1 = 3.6\%$, $\omega_1 = 1$	Not allowed
M2a: Positive	4.18	50	-5656.81	$p_0 = 96.47\%$, $\omega_0 = 0.0054$; $p_1 = 3.55\%$, $\omega_1 = 1$; $p_2 = 0$, $\omega_2 = 4.46$	71 L (72.4%), 91 L (78.0%)
M3: Discrete	4.90	51	-5637.73	$p_0 = 90.54\%, \ \omega_0 = 0; \ p_1 = 8.51\%, \ \omega_1 = 0.5308;$ $p_2 = 0.943\%, \ \omega_2 = 1.71$	71 L (96.1%), 91 L (97.1%) , 351 V (81.3%)
M7: β	4.78	48	-5648.93	<i>p</i> = 0.03078, <i>q</i> = 0.63009	Not allowed
Μ8: β &ω > 1	5.09	50	-5641.45	$p_0 = 98.21\%$, $p = 0.01555$, $q = 0.36594$; $p_1 = 1.78\%$, $\omega = 1.19067$	71 L (96.4%) , 91 L(97.2%) , 351 V (84.9%)

Table 4 Parameter estimates and log-likelihood values under models of variable ω ratios among co	odons

*The contexts of the dataset were introduced in Additional file 1 Table S3. Codons with PP > 95% were in bold. The phylogeny trees applied in the estimation were reconstructed throughout dataset introduced in Additional file 1 Table S3 accordingly.

**Positions and posterior probability of the positively selected codons were showed in this column. The posterior probabilities were given via Bayes Empirical Bayes (BEB) method.

rbcL gene and *psbA-trnH* spacer region as universal markers [49]. Our results indicated that the *psbA* and *psbA-trnH* sequences have high resolution at order and family level, but low resolution at several genera level (Figures 1 and 2, Table 5). Further investigation with extra locus will have a deeper reflection on this task.

Genome rearrangement and psbA duplication

The PCR amplification results coincide with the previous conclusion based on the completely and partially sequenced plastomes [10-13]: three kinds of gene order (*trnK-psbA-trnH*, *matK-psbA-trnH* and *rps7-psbA-trnH*) were found in the 20 species from 8 fern orders (Figure 1). The data of the extant complete fern cp genomes showed that the *trnK*- *psbA-trnH* and *matK-psbA-trnH* fragments are located in the LSC and the *rps7-psbA-trnH* fragment is in the IRs [10,13,50]. Since the large-scale inversion events involving the duplication of *psbA* gene are rare during the evolution history of fern species [8,9], one copy of *psbA* exists in those species of *trnK-psbA-trnH* and *matK-psbA-trnH* types and two in *rps7-psbA-trnH*. Their full-length *psbA* encoding sequences have not been determined in previous studies.

Evolutionary trajectory and survival strategy

The strong competition between ferns and angiosperms for light may have been the main cause for the evolutionary trajectory of *psbA* gene. Nevertheless, the occurrence of *psbA* duplication provided an alternative



Software or Website*	Co-evolutionary Pair**		
	Site1	Site2	
CAPS	4	71	
Datamonkey***	19	350	
InterMap3D	19	350	
	53	351	
	4	351	
	71	92	
	4	352	
	71	72	
	72	281	
	4	53	
	72	351	
	351	353	

*Three packages were applied in the current researches [46,47,73].

**Site1, Site 2: Coordinates of each of the interacting residues.

***Two kinds of molecules (two parents and one parent) were applied and gave the same pair.

survival strategy versus the molecular adaptive evolution at the codons of D1 protein. Several positively selected sites were found in single *psbA* gene copy and none in duplicated *psbA* (Table 2). These results indicated that single *psbA* gene might have functional adaptation via the replacement at certain positions among the D1 protein while the duplicated one might have not. Conversely to the case of single gene copies, the leptosporangiate ferns with duplicate *psbA* gene may have acquired further fitness gain through the existence of a new transcription locus for the synthesis of D1 protein, which might directly increase the efficiency of photosynthesis by protein dosage effects [20,51,52].

The structure and function of the D1 protein are conserved among cyanobacteria, red algae and plants [53,54]. The purifying selection of *psbA* in Lejeuneaceae has been noticed in a recent study [55]. Regretfully, not like the elegant investigation in *rbcL* gene [56], the detail function of the codons in D1 protein remains obscure. Mutation experiments of D1 protein will unveil the functional importance of the positively selected positions. Moreover, the further research on the relationships between positively selected mutations among *psbA* gene from fern species and the biodiversity will have great impacts on the understanding of fern evolutionary biology.

Complicated intra-molecular evolution under selective pressure

Eleven co-evolving sites (Table 5) have been detected via different molecules, and they (site 4, 19, 53, 71, 72, 92, 281, 350, 351, 352 and 353) may be more important than others in D1 protein. Point mutation experiments may have distinctive outcomes, some could cause severe functional consequences and others could result in completely undetectable change. This fact indicates that a protein is a network of interacting residues, and the core nodes in this network determine the function of the protein [57]. Future point mutation experiments aiming at the eleven sites may have butterfly effect on the protein structure and function.

psbA gene, along with three other chloroplast genes, encodes the core proteins complex in the chloroplasts of ferns. However, as we have concluded before the functional adaptation of this complex might be caused by the inter-coevolution among different proteins [32]. The current intra-molecular evolution analysis may have shortcomings on predicting the complicated coevolutionary networks among the entire functional core complex. Further conclusion could not be declared without overall analysis based on the four related genes: *psbA*, *psbB*, *psbC* and *psbD*.

Conclusion

In the current research we present evidence that point to a complex adaptive process mediating the functional innovation of the D1 protein. This process involves a multiply checking of the structural and functional consequences of the fixation of functionally novel mutations and the amelioration of the effects by such mutations may have through compensatory replacement events. A serial amino acid sites are identified as co-evolution positions while significant positive selection signals are detected in the single copy *psbA* gene from the fern species. One hypothetical scenario is put forward: i) single copy psbA gene fern species may adapt to the newly formed living circumstance by the modification of amino acids in D1 protein; ii) by the meantime, the dosage effects of D1 protein are the possible strategy against the rising of angiosperm in psbA duplicated ones. The selective relaxation in duplicated psbA genes could be observed throughout different models. However, no evidence stands for the functional divergence in the duplicated psbA genes. Future investigation will shed new lights on this question. Although this research covers only a little of the diversity of species in fern, our sampling included all 11 orders. Our research however opens exciting new avenues that will hopefully lead to a more complete understanding of the functional novelties and dosage effects in the D1 protein among ferns.

Methods

Sampling of plant materials

Plant materials of 20 fern species for the present investigation were collected from Wuhan Botanical Garden, Fairy Lake Botanical Garden and South China Botanical Garden, respectively (Table 1).

Isolation of total genomic DNA

For each species, three pieces of fresh leaves were collected to isolate genomic DNA with the modified CTAB protocols [58]. Each sample was dissolved in 50 μ l TE buffer. Roughly, the quality was determined by 1% agarose/TAE gel electrophoresis and the quantity was estimated via DNA Ladder (Takara). The bright sample under UV-light with right size was used as the template in PCR reactions. The absorption at 260 and 280 nm of qualified template DNA was measured using a 752 spectrophotometer. The purity and concentration was resolved and calculated by the A₂₆₀/A₂₈₀ ratio and A₂₆₀ absorption value.

PCR amplification and DNA sequencing

PCR amplification was carried out in 100 µl volumes containing 50 mM KCl, 10 mM Tris-HCl (pH 8.0), 0.1% Triton X-100, 1.5 mM MgCl₂, 0.2 mM each deoxynucleoside triphosphate, 2 U Taq DNA polymerase, 0.3 µM primer, 30 ng genomic DNA and DNA-free water. The 3-step and 2-step PCR protocols employed speciesspecific and universal primers, respectively. Individually, the annealing temperature in 3-step PCR reaction (Additional file 1 Table S1) of species-specific primers was calculated via Primer 3 (http://frodo.wi.mit.edu/primer3/). The thermo-cycling program was set as: 5 min at 95°C (1 cycle); 45 s at 94°C, 60s at Ta°C, 90s at 72°C (34 cycles); 10 min at 72°C (1 cycle). However, the annealing temperature was ignored in 2-step PCR reaction (Additional file 1 Table S2). The thermo-cycling program was set as: 5 min at 95°C (1 cycle); 60s at 94°C, 150 s at 60°C (35 cycles); 20 min at 72°C (1 cycle). Except as normal reactions, the genomic DNA was excluded from the reaction mix for negative control. Then the molecular weight of PCR products was verified in 1% agarose/TAE gels.

Each qualified DNA fragment amplified by the above steps was recovered and purified with a quick PCR Purification Kit (Promega), and then cloned into PMD19-T vectors (Takara). The plasmids, composed of the vectors and the DNA fragments, were transformed to *Escherichia coli* strain DH5 α . Plasmids within positive clones were extracted and sequenced with an ABI PRISM 3730 DNA analyzer. Three clones were sequenced for each amplicon to control *Taq* polymerase errors. The overlapping sequences from various amplification steps were assembled as a single contig. To ascertain the contigs' locations among cp genomes, the sequencing results were submitted to DOGMA website [59].

Multiple sequence alignments and best-fit nucleotide substitution model

Six multiple sequence alignments (hereafter MSAs) were established for the present investigation (Additional file 1 Table S3). Nucleotide sequences obtained experimentally (Table 1) plus those retrieved from the public databases (Additional file 1 Table S4) were aligned using the MUSCLE software [60]. Partially fern *psbA* sequences from GenBank were excluded from the present study.

The best-fit nucleotide substitution model for each MSA was selected via jModeltest package [61]. And it was also estimated via the automatic model selection tool at the Datamonkey website (http://www.datamonkey.org) for the coding regions [43,44].

Reconstruction of phylogeny along with time-scale

Isoetes flaccida (GenBank Accession GU191333) was selected as outgroup in the phylogenetic analyses. Two nodes were chosen to constrain for a rate consistent with the known fossils: 1) Since *Osmunda* fossils have been described from the Upper Triassic [62], the Osmundaceae clade was constrained to 199.6 million years ago (Mya, Node 4); 2) According to the fossil *Gleichenia*, node 14 were dated to have been originated 99.6 Mya [63]. Moreover, following previous estimates, another node 21 and 22 were separately dated to 110.5 Mya and 42.5 Mya at the beginning of the running: 3) Pteridaceae clade [64] and 4) Polypodiaceae clade [25].

According to authors' suggestion [37], to avoid the misspecification of dating and taxon sampling, the empty alignment was run before the real MSAs. Then BEAST 1.7 was allowed to infer topology, branch lengths, and dates for combined datasets [38]. A uniform distribution is applied over the estimating of the absolute ages via the MCMC process. For each MSA, BEAST runs 6×10^7 generations, saving data every 1,000 generations, producing 60,000 estimates of dates under a Yule speciation prior under the uncorrelated lognormal distributed relaxed clock model. Convergence statistics was analyzed in Tracer v 1.5, resulting in 54,000 post-burn-in trees. Before the consensus tree was graphically illustrated by Figtree v.1.3.1, TreeAnnotator v.1.6.1 was utilized to produce maximum clade credibility trees from the postburn-in trees and to determine the 95% probability density of ages for all nodes in the phylogenetic tree.

Detection of positively and negatively selected codons

Since identification of positive/negative (non-neutral) evolution is fundamental to the understanding of the process of diversifying/purifying selection, this subject has been the focus of several decades of mathematical and computational efforts. Different scientists have developed numerous analysis models and methods, and each has its own advantages [42-44,65]. However, the general consensus of them is that non-synonymous nucleotide substitutions (d_N), whose alternatives leading to a change in the codon and its corresponding amino acid, can be time-scaled by the number of synonymous replacements (d_S), which are nucleotide changes that only change the codon but not the amino acid and are consequently neutrally fixed and proportional to the divergence time between the sequences.

The random-site models (M0 vs. M3, M1a vs. M2a, M5 vs. null test, M7 vs. M8, and MEC vs. null test), contained in PAML package version 4.1 and Selecton version 2.2, allow the ω ratio ($\omega = d_N/d_S$) to shift among codons within the MSA and this parameter is estimated by maximum-likelihood value via Bayesian inference approach [41,42]. Additionally, the results from Selecton version 2.2 are visualized with seven-color scale for representing the different types of selection. To identify the statistical significant levels of the results, the LRT was conducted to compare the nested models [66].

Besides, another three models for detecting codons under selection are implemented on Datamonkey website [43]: SLAC, FEL and REL. SLAC originated from the Suzuki-Gojobori counting approach [67] and is quite efficient on detecting non-neutral evolution in large MSAs. Less conservative than SLAC, FEL fits a site-specific $d_{\rm N}$ and $d_{\rm S}$ in the context of codon substitution models and tests whether $d_{\rm N} = d_{\rm S}$, outperforming other two models in MSAs of intermediate size. As the most powerful model of the three, REL is improved based on the Nielsen-Yang approach [68]. Before the analysis of Natural selection, the best-fit nucleotide substitution models in the MSAs were calculated via the model-selection molecule online. Different from the posterior LRTs in the nested models, the parameter for statistical significant level (p value or Bayesian factor) was pre-set prior to the estimating processes [69].

Analysis of inter-dependent evolutionary sites

To understand the broad implications of the amino acid replacements in D1 proteins from fern species, the analysis of the evolutionary dependencies among sites to identify functional/structural dependencies among residues were conducted via five.

Based on a tree-ignorant strategy, CAPS outperforms the tree-aware strategy methods as reported by previous published work [70], which compares the correlated variance of the evolutionary rates at 2 sites corrected by the time since the divergence of the 2 sequences [46]. The significance of the results was evaluated by randomization of pairs in the alignment, calculation of their correlation values, and comparison of the real values with the distribution of 10,000 randomly sampled values. The stepdown permutation procedure was employed to correct for multiple tests and non-independence of data [71]. An alpha value of 0.001 was applied to minimize Type I errors. The correlated variability between amino acid sites was weighted by the level of substitutions per synonymous site in order to normalize parameters by the time of sequence divergence [72].

The mathematical models for detecting the coevolving residues in protein from InterMap3D and Datamonkey websites are based on a tree-aware strategy [47,73]. Currently, three different models, namely Row and Column Weighing of Mutual Information (RCW-MI), Dependency and Mutual Information/Entropy (MI/E), were implemented at the former website [47], and one (Spidermonkey) at the later [43,73]. RCW-MI and Dependency extract both the entropy dependency and the phylogenetic signal [74,75], while the MI/E extracts the entropy dependency from the signal by dividing mutual information by the joint site's entropy [76,77]. Spidermonkey molecule reconstructs the substitution history of the MSAs by ML-based methods, and analyses the joint distribution of substitution events by Bayesian graphical models [73].

Additional files

Additional file 1: Table S1. Universal primers for fern psbA gene. S2 Species-specific primers for *psbA* gene from eight fern species. S3 Six datasets of different species and fragments. S4 Species and accession number of the retrieved data. S5 LRTs of the random-site models in PAML version 4.1*. S6 Positively selected codons determined via REL model.

Additional file 2: Currently determined 20 sequences. The sequences started at the ATG codon of *psbA*, ended before the ATG codon of *tmH*. They included the stop codon of *psbA* and the intergenic region between *psbA* and *trnH*. They were applied to reconstruct the phylogenetic trees.

Abbreviations

cp: Chloroplast; FEL: Fixed effects likelihood; IR(s): Inverted repeat region(s); LSC: Large single copy region; LRT: Likelihood ratio test; Mya: Million (1 × 10⁶) years ago; MI/E: Mutual information/entropy; MSA(s): Multiple sequence alignment(s); PS II: Photo system II; REL: Random effects likelihood; RCW-MI: Row and column weighing of mutual information; SLAC: Single likelihood ancestor counting; Ta: Annealing temperature.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

LS carried out the experimental works, participated in the sequence alignment, molecular evolution analysis and drafted the manuscript. MF contributed to the analysis tools and drafted the manuscript. TW and YJS conceived of the study, and participated in its design and coordination. All authors read and approved the final manuscript.

Acknowledgments

We thank Dr. Yefu Wang at State Key Laboratory of Virology, College of Life Sciences, Wuhan University, China, for the guidance for wording, and we also thank Bo Wang and Dr. Lei Gao at the Wuhan Botanical Garden, Chinese Academy of Sciences, China, for the experimental assistance. We thank Dr. Jianqiang Li at the Wuhan Botanical Garden, Chinese Academy of Sciences, China, for the advices on species sampling. We appreciate two anonymous reviewers and other editors for their helpful suggestions. The present work was financially supported by the National Nature Science Foundation of China (No. 30970290, 31070594), and by the Knowledge Innovation Program of the Chinese Academy of Sciences (No. KSCX2-EW-J-20, KSCX2-YW-Z-0940). Dr. Mario A. Fares was supported by Spanish Ministerio de Ciencia e Inovación (No. BFU2009-12022) and Research Frontiers Program (No. 10/RFP/ GEN2685) from Science Foundation Ireland.

Author details

¹State Key Laboratory of Virology, College of Life Sciences, Wuhan University, Wuhan, China. ²CAS Key Laboratory of Plant Germplasm Enhancement and Specialty Agriculture, Wuhan Botanical Garden, Chinese Academy of Sciences, Wuhan, China. ³Integrative Systems Biology Group, Instituto de Biología Molecular y Celular de Plantas (CSIC-Universidad Politécnica de Valencia (UPV)), Valencia, Spain. ⁴Evolutionary Genetics and Bioinformatics Laboratory, Department of Genetics, University of Dublin, Trinity College, Dublin 2, Ireland. ⁵State Key Laboratory of Biocontrol, School of Life Sciences, Sun Yat-Sen University, Guangzhou, China.

Received: 12 March 2012 Accepted: 8 August 2012 Published: 16 August 2012

References

- Shi L-X, Hall M, Funk C, Schröder WP: Photosystem II, a growing complex: updates on newly discovered components and low molecular mass proteins. *Biochim Biophys Acta-Bioenerg* 2012, 1817(1):13–25.
- Pospisil P: Molecular mechanisms of production and scavenging of reactive oxygen species by photosystem II. *Biochim Biophys Acta-Bioenerg* 2012, 1817(1):218–231.
- 3. Hohmann-Marriott MF, Blankenship RE: Evolution of photosynthesis. Annu Rev Plant Biol 2011, 62:515–548.
- Barber J, Nield J, Morris E, Zheleva D, Hankamer B: The structure, function and dynamics of photosystem two. *Physiol Plant* 1997, 100(4):817–827.
- Minai L, Wostrikoff K, Wollman FA, Choquet Y: Chloroplast biogenesis of photosystem II cores involves a series of assembly-controlled steps that regulate translation. *Plant Cell* 2006, 18(1):159–175.
- Guskov A, Kern J, Gabdulkhakov A, Broser M, Zouni A, Saenger W: Cyanobacterial photosystem II at 2.9-Å resolution and the role of quinones, lipids, channels and chloride. *Nat Struct Mol Biol* 2009, 16(3):334–342.
- Pospisil P: Production of reactive oxygen species by photosystem II. Biochim Biophys Acta-Bioenerg 2009, 1787(10):1151–1160.
- Stein DB, Conant DS, Ahearn ME, Jordan ET, Kirch SA, Hasebe M, Iwatsuki K, Tan MK, Thomson JA: Structural rearrangements of the chloroplast genome provide an important phylogenetic link in ferns. *Proc Natl Acad Sci U S A* 1992, 89(5):1856–1860.
- 9. Raubeson LA, Stein DB: Insights into fern evolution from mapping chloroplast genomes. *Am Fern J* 1995, **85**(4):193–204.
- Gao L, Yi X, Yang YX, Su YJ, Wang T: Complete chloroplast genome sequence of a tree fern *Alsophila spinulosa*: insights into evolutionary changes in fern chloroplast genomes. *BMC Evol Biol* 2009, 9:130.
- Wolf PG, Rowe CA, Sinclair RB, Hasebe M: Complete nucleotide sequence of the chloroplast genome from a leptosporangiate fern, Adiantum capillus-veneris L. DNA Res 2003, 10(2):59–65.
- Roper JM, Hansen SK, Wolf PG, Karol KG, Mandoli DF, Everett KDE, Kuehl J, Boore JL: The complete plastid genome sequence of Angiopteris evecta (G. Forst.) Hoffin. (Marattiaceae). Am Fern J 2007, 97(2):95–106.
- Wolf PG, Roper JM, Duffy AM: The evolution of chloroplast genome structure in ferns. *Genome* 2010, 53(9):731–738.
- Lynch M, Conery JS: The evolutionary fate and consequences of duplicate genes. Science 2000, 290(5494):1151–1155.
- Zhang JZ: Evolution by gene duplication: an update. Trends Ecol Evol 2003, 18(6):292–298.
- Tautz D, Domazet-Loso T: The evolutionary origin of orphan genes. Nat Rev Genet 2011, 12(10):692–702.
- Gravemann S, Schnipper N, Meyer H, Vaya A, Nowaczyk MJ, Rajab A, Hofmann WK, Salewsky B, Tonnies H, Neitzel H, et al: Dosage effect of zero to three functional LBR-genes in vivo and in vitro. Nucleus 2010, 1(2):179–189.

- Innan H, Kondrashov F: The evolution of gene duplications: classifying and distinguishing between models. Nat Rev Genet 2010, 11(2):97–108.
- Colbourne JK, Pfrender ME, Gilbert D, Thomas WK, Tucker A, Oakley TH, Tokishita S, Aerts A, Arnold GJ, Basu MK, *et al*: The ecoresponsive genome of daphnia pulex. *Science* 2011, 331(6017):555–561.
- 20. Kondrashov FA, Rogozin IB, Wolf YI, Koonin EV: Selection in the evolution of gene duplications. *Genome Biol* 2002, **3**(2):8.1-8.9.
- 21. Lidgard S, Crane PR: Quantitative analyses of the early angiosperm radiation. *Nature* 1988, **331**(6154):344–346.
- 22. Niklas KJ, Tiffney BH, Knoll AH: Patterns in vascular land plant diversification. *Nature* 1983, **303**(5918):614–616.
- 23. Lupia R, Lidgard S, Crane PR: Comparing palynological abundance and diversity: implications for biotic replacement during the Cretaceous angiosperm radiation. *Paleobiology* 1999, **25**(3):305–340.
- Crisp MD, Cook LG: Cenozoic extinctions account for the low diversity of extant gymnosperms compared with angiosperms. *New Phytol* 2011, 192(4):997–1009.
- Schuettpelz E, Pryer KM: Evidence for a Cenozoic radiation of ferns in an angiosperm-dominated canopy. Proc Natl Acad Sci U S A 2009, 106(27):11200–11205.
- 26. Wang GZ, Lercher MJ: The effects of network neighbours on protein evolution. *PLoS One* 2011, 6(4):e18288.
- 27. Liang Z, Xu M, Teng MK, Niu LW, Wu JR: Coevolution is a short-distance force at the protein interaction level and correlates with the modular organization of protein networks. *FEBS Lett* 2010, **584**(19):4237–4240.
- Lovell SC, Robertson DL: An integrated view of molecular coevolution in protein-protein interactions. Mol Biol Evol 2010, 27(11):2567–2575.
- Poon A, Chao L: The rate of compensatory mutation in the DNA bacteriophage phiX174. *Genetics* 2005, 170(3):989–999.
- Mateo R, Mateu MG: Deterministic, compensatory mutational events in the capsid of foot-and-mouth disease virus in response to the introduction of mutations found in viruses from persistent infections. *J Virol* 2007, 81(4):1879–1887.
- 31. Davis BH, Poon AF, Whitlock MC: Compensatory mutations are repeatable and clustered within proteins. *Proc Biol Sci* 2009, **276**(1663):1823–1827.
- 32. Sen L, Fares MA, Liang B, Gao L, Wang B, Wang T, Su YJ: Molecular evolution of *rbcL* in three gymnosperm families: identifying adaptive and coevolutionary patterns. *Biol Direct* 2011, **6**:29.
- Smith AR, Pryer KM, Schuettpelz E, Korall P, Schneider H, Wolf PG: A classification for extant ferns. *Taxon* 2006, 55(3):705–731.
- 34. Lehtonen S: Towards resolving the complete fern tree of life. *PLoS One* 2011, 6(10):e24851.
- Yang Z: Computational molecular evolution. USA: Oxford University Press; 2006.
- 36. Maddison DR, Maddison WP: *MacClade 4: Analysis of phylogeny and character evolution*. Version 4.08a http://macclade.org edn; 2005.
- 37. Drummond AJ, Rambaut A: BEAST: Bayesian evolutionary analysis by sampling trees. *BMC Evol Biol* 2007, **7**:214.
- Drummond AJ, Suchard MA, Xie D, Rambaut A: Bayesian phylogenetics with BEAUti and the BEAST 1.7. Mol Biol Evol 2012.
- 39. Fazekas AJ, Kuzmina ML, Newmaster SG, Hollingsworth PM: DNA barcoding methods for land plants. *Methods Mol Biol* 2012, **858**:223–252.
- Chase MW, Salamin N, Wilkinson M, Dunwell JM, Kesanakurthi RP, Haidar N, Savolainen V: Land plants and DNA barcodes: short-term and long-term goals. *Philos Trans R Soc Lond B Biol Sci* 2005, 360(1462):1889–1895.
- 41. Yang Z: PAML 4: phylogenetic analysis by maximum likelihood. *Mol Biol Evol* 2007, 24(8):1586–1591.
- Stern A, Doron-Faigenboim A, Erez E, Martz E, Bacharach E, Pupko T: Selecton 2007: advanced models for detecting positive and purifying selection using a Bayesian inference approach. *Nucleic Acids Res* 2007, 35:W506–W511.
- Delport W, Poon AF, Frost SD, Pond SLK: Datamonkey 2010: a suite of phylogenetic analysis tools for evolutionary biology. *Bioinformatics* 2010, 26(19):2455–2457.
- Pond SLK, Frost SD: Datamonkey: rapid detection of selective pressure on individual sites of codon alignments. *Bioinformatics* 2005, 21(10):2531–2533.
- Doron-Faigenboim A, Pupko T: A combined empirical and mechanistic codon model. Mol Biol Evol 2007, 24(2):388–397.
- Fares MA, McNally D: CAPS: coevolution analysis using protein sequences. Bioinformatics 2006, 22(22):2821–2822.

- Gouveia-Oliveira R, Roque FS, Wernersson R, Sicheritz-Ponten T, Sackett PW, Molgaard A, Pedersen AG: InterMap3D: predicting and visualizing co-evolving protein residues. *Bioinformatics* 2009, 25(15):1963–1965.
- Fazekas AJ, Burgess KS, Kesanakurti PR, Graham SW, Newmaster SG, Husband BC, Percy DM, Hajibabaei M, Barrett SC: Multiple multilocus DNA barcodes from the plastid genome discriminate plant species equally well. *PLoS One* 2008, 3(7):e2802.
- Kress WJ, Erickson DL: A two-locus global DNA barcode for land plants: the coding rbcL gene complements the non-coding trnH-psbA spacer region. *PLoS One* 2007, 2(6):e508.
- Karol KG, Arumuganathan K, Boore JL, Duffy AM, Everett KD, Hall JD, Hansen SK, Kuehl JV, Mandoli DF, Mishler BD, et al: Complete plastome sequences of Equisetum arvense and Isoetes flaccida: implications for phylogeny and plastid genome evolution of early land plant lineages. BMC Evol Biol 2010, 10:321.
- Brown CJ, Todd KM, Rosenzweig RF: Multiple duplications of yeast hexose transport genes in response to selection in a glucose-limited environment. *Mol Biol Evol* 1998, 15(8):931–942.
- 52. Bekaert M, Conant GC Copy number alterations among mammalian enzymes cluster in the metabolic network. *Mol Biol Evol* 2011, **28**(2):1111–1121.
- Müh F, Renger T, Zouni A: Crystal structure of cyanobacterial photosystem II at 3.0 Å resolution: a closer look at the antenna system and the small membrane-intrinsic subunits. *Plant Physiol Biochem* 2008, 46(3):238–264.
- Guskov A, Gabdulkhakov A, Broser M, Glockner C, Hellmich J, Kern J, Frank J, Muh F, Saenger W, Zouni A: Recent progress in the crystallographic studies of photosystem II. Chemphyschem 2010, 11(6):1160–1171.
- Chen X, Su Y, Wang T: Adaptive evolution analysis of the *psbA* gene in Lejeuneaceae. Acta Botanica Boreali-Occidentalia Sinica (in Chinese) 2010, 30(1):1534–1544.
- Kreel N, Tabita F: Substitutions at Methionine 295 of Archaeoglobus fulgidus Ribulose-1, 5-bisphosphate Carboxylase/Oxygenase Affect Oxygen Binding and CO₂/O₂ Specificity. J Biol Chem 2007, 282(2):1341–1351.
- Lee BC, Park K, Kim D: Analysis of the residue-residue coevolution network and the functionally important residues in proteins. *Proteins* 2008, 72(3):863–872.
- Sambrook J, Fritsch EF, Maniatis T: Molecular cloning: a laboratory manual. Cold Spring Harbor. USA: Cold Spring Harbor Laboratory Press; 2001.
- Wyman SK, Jansen RK, Boore JL: Automatic annotation of organellar genomes with DOGMA. *Bioinformatics* 2004, 20(17):3252–3255.
- 60. Edgar RC: MUSCLE: a multiple sequence alignment method with reduced time and space complexity. *BMC Bioinforma* 2004, 5:113.
- Posada D: jModelTest: phylogenetic model averaging. Mol Biol Evol 2008, 25(7):1253–1256.
- Phipps C, Taylor T, Taylor E, Cuneo R, Boucher L, Yao X: Osmunda (Osmundaceae) from the Triassic of Antarctica: an example of evolutionary stasis. Am J Bot 1998, 85(6):888–895.
- Herendeen PS, Skog JE: Gleichenia chaloneri-a new fossil fern from the lower Cretaceous (Albian) of England. Int J Plant Sci 1998, 159(5):870–879.
- Sen L, Su Y, Zhang B, Wang T: Adaptive evolution of the *rbcL* gene in Pteridaceous ferns. Journal of Tropical and Subtropical Botany (in Chinese) 2010, 18(1):1–8.
- 65. Yang Z: PAML: a program package for phylogenetic analysis by maximum likelihood. *Comput Appl Biosci* 1997, **13**(5):555–556.
- Anisimova M, Bielawski JP, Yang Z: Accuracy and power of the likelihood ratio test in detecting adaptive molecular evolution. *Mol Biol Evol* 2001, 18 (8):1585–1592.
- Suzuki Y, Gojobori T: A method for detecting positive selection at single amino acid sites. Mol Biol Evol 1999, 16(10):1315–1328.
- Nielsen R, Yang Z: Likelihood models for detecting positively selected amino acid sites and applications to the HIV-1 envelope gene. *Genetics* 1998, 148(3):929–936.
- Pond SLK, Frost SD: Not so different after all: a comparison of methods for detecting amino acid sites under selection. Mol Biol Evol 2005, 22(5):1208–1222.
- Caporaso JG, Smit S, Easton BC, Hunter L, Huttley GA, Knight R: Detecting coevolution without phylogenetic trees? Tree-ignorant metrics of coevolution perform as well as tree-aware metrics. *BMC Evol Biol* 2008, 8:327.
- Taberner A, Castanera P, Silvestre E, Dopazo J: Estimation of the intrinsic rate of natural increase and its error by both algebraic and resampling approaches. *Comput Appl Biosci* 1993, 9(5):535–540.

- 72. Li WH: Unbiased estimation of the rates of synonymous and nonsynonymous substitution. *J Mol Evol* 1993, **36**(1):96–99.
- Poon AF, Lewis FI, Frost SD, Pond SLK: Spidermonkey: rapid detection of co-evolving sites using Bayesian graphical models. *Bioinformatics* 2008, 24(17):1949–1950.
- Tillier ER, Lui TW: Using multiple interdependency to separate functional from phylogenetic correlations in protein alignments. *Bioinformatics* 2003, 19(6):750–755.
- Gouveia-Oliveira R, Pedersen AG: Finding coevolving amino acid residues using row and column weighting of mutual information and multidimensional amino acid representation. *Algorithms Mol Biol* 2007, 2:12.
- Martin LC, Gloor GB, Dunn SD, Wahl LM: Using information theory to search for co-evolving residues in proteins. *Bioinformatics* 2005, 21(22):4116–4124.
- Makarova KS, Wolf YI, van der Oost J, Koonin EV: Prokaryotic homologs of Argonaute proteins are predicted to function as key components of a novel system of defense against mobile genetic elements. *Biol Direct* 2009, 4:29.

doi:10.1186/1471-2148-12-145

Cite this article as: Sen *et al.*: Molecular evolution of *psbA* gene in ferns: unraveling selective pressure and co-evolutionary pattern. *BMC Evolutionary Biology* 2012 **12**:145.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at www.biomedcentral.com/submit

BioMed Central