Origin and Diversification of Basic-Helix-Loop-Helix Proteins in Plants

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Abstract

Basic helix-loop-helix (bHLH) proteins are a class of transcription factors found throughout eukaryotic organisms. Classification of the complete sets of bHLH proteins in the sequenced genomes of *Arabidopsis thaliana* and *Oryza sativa* (rice) has defined the diversity of these proteins among flowering plants. However, the evolutionary relationships of different plant bHLH groups and the diversity of bHLH proteins in more ancestral groups of plants are currently unknown. In this study, we use whole-genome sequences from nine species of land plants and algae to define the relationships between these proteins in plants. We show that few (less than 5) bHLH proteins are encoded in the genomes of chlorophytes and red algae. In contrast, many bHLH proteins (100–170) are encoded in the genomes of land plants (embryophytes). Phylogenetic analyses suggest that plant bHLH proteins are monophyletic and constitute 26 subfamilies. Twenty of these subfamilies existed in the common ancestors of extant mosses and vascular plants, whereas six further subfamilies evolved among the vascular plants. In addition to the conserved bHLH domains, most subfamilies are characterized by the presence of highly conserved short amino acid motifs. We conclude that much of the diversity of plant bHLH proteins was established in early land plants, over 440 million years ago.

Key words: bHLH, transcription factor, plants, algae, evolution, phylogeny.

Introduction

The basic helix-loop-helix (bHLH) domain is a highly conserved amino acid motif that defines a group of transcription factors. It was originally described in animals (Murre et al. 1989) and soon discovered in all the major eukaryotic lineages. Proteins that contain a bHLH domain (referred to as bHLH proteins) are involved in a myriad of regulatory processes. Their functions include regulating neurogenesis, myogenesis, and heart development in animals (Massari and Murre 2000; Jones 2004); controlling phosphate uptake and glycolysis in yeast (Robinson and Lopes 2000); or modulating secondary metabolism pathways, epidermal differentiation, and responses to environmental factors in plants (Ramsay and Glover 2005; Castillon et al. 2007).

The bHLH domain consists of 50–60 amino acids that form two distinct segments: a stretch of 10–15 predominantly basic amino acids (the basic region) and a section of roughly 40 amino acids predicted to form two amphipathic α -helices separated by a loop of variable length (the helix-loop-helix region). Structural analyses of mammalian and yeast bHLH proteins showed that the basic region forms the main interface where contact with DNA occurs, whereas the two helices promote the formation of homoor heterodimers between bHLH proteins, a prerequisite for DNA binding to occur (Jones 2004).

Phylogenetic analyses have classified the diversity of bHLH proteins into a number of distinct groups. Over 50 bHLH proteins are encoded in the genomes of most an-

imals (metazoans) and are typically classified into six major groups (A-F), based on their ability to bind DNA (Atchley and Fitch 1997; Ledent and Vervoort 2001; Jones 2004). Detailed analyses using whole-genome sequences showed that animal bHLH could be further classified in several smaller subfamilies that are highly conserved across major metazoan lineages (Ledent and Vervoort 2001; Simionato et al. 2007). Phylogenetic analyses indicate that 44 of these subfamilies were present in the common ancestor of all bilaterians, which is thought to have existed sometime before 600 million years ago (Ma) (Simionato et al. 2007). The genomes of Arabidopsis thaliana and Oryza sativa (rice) encode even more bHLH sequences than animals. Different phylogenetic studies proposed the classification of plant bHLH into 15-25 subgroups (Buck and Atchley 2003; Heim et al. 2003; Toledo-Ortiz et al. 2003; Li et al. 2006b). However, the origin and evolutionary history of these groups cannot be understood using A. thaliana and O. sativa sequences alone. The characterization of the evolution of plant bHLH diversity requires the phylogenetic analysis of bHLH proteins from a more diverse selection of plants, including algae, bryophytes, and different lineages of vascular plants.

In this study, we characterized the evolution of bHLH proteins in plants, defined here as the organisms that are likely to have been derived from the primary endosymbiotic event that gave rise to the red algae, chlorophytes, and land plants (Rodríguez-Ezpeleta et al. 2005). We show that the plant bHLH family is monophyletic and underwent

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a major radiation before the evolution of the mosses. The bHLH groups established in the early land plants over 400 Ma were conserved during subsequent plant evolution, although there were many gene duplications and losses within these groups. Our analysis defines 26 subfamilies that represent deep evolutionary relationships between plant bHLH proteins.

Materials and Methods

Sequence Retrieval

The A. thaliana bHLH reported by Bailey et al. (2003), Heim et al. (2003) and Toledo-Ortiz et al. (2003) were retrieved from The Arabidopsis Information Resource (http://www. arabidopsis.org/). A clear bHLH domain was not found in At1g31050 (AtbHLH111) and At1g22380 (AtbHLH152), so they were not further used in this study; we could not find At2g20095 (AtbHLH133) and At4g38071 (AtbHLH131) in any database. A data set of predicted O. sativa L. ssp. japonica bHLH proteins was retrieved from the Plant TFDB (Guo et al. 2008) and combined with the bHLH protein sequences reported by Li et al. (2006b), retrieved from the Rice Genome Annotation Project (http://rice.plantbiology.msu. edu/). Eleven new proteins were numbered following the nomenclature style of Li et al. (2006b), whereas a clear bHLH was not found in Os01g65080 (OsbHLH033), Os04g35000 (OsbHLH145), Os11g02054 (OsbHLH160), and Os12g02020 (OsbHLH161). A data set of predicted Physcomitrella patens bHLH was retrieved from the Plant TFDB (Guo et al. 2008). A direct search of genes annotated as bHLH was performed on the genome assembly of Selaginella moellendorffii v1.0 (http://www.jgi.doe.gov/). HMMsearch (Eddy 1998) was used to screen the genome assemblies of Cyanidioschyzon merolae (Matsuzaki et al. 2004), Chlamydomonas reinhardtii v3.0 (Merchant et al. 2007), Ostreoscoccus tauri v2.0 (Palenik et al. 2007), Thalassiosira pseudonana v3.0 (Armbrust et al. 2004), and the draft assemblies of Chlorella vulgaris C-169 and Volvox carteri (http://www.jgi.doe.gov/) with the PFAM profile hidden Markov model (pHMM) HLH_ls.hmm (http://pfam. sanger.ac.uk/).

Five Homo sapiens and four Amphimedon queenslandica (demosponge) representative sequences of the major metazoan groups of bHLH proteins (based on Jones 2004; Simionato et al. 2007) were retrieved from GenBank; group F proteins are not clearly alignable to other bHLH (Ledent et al. 2002) and so they were not used in this study. The Saccharomyces cerevisiae bHLH proteins reported by Robinson and Lopes (2000) were retrieved from http://www.yeastgenome.org/.

For simplicity, all sequences were renamed according to the supplementary table S1 (Supplementary Material online). The complete amino acid sequence of all proteins can be found in supplementary data 1 (Supplementary Material online).

Alignment and Phylogenetic Analysis

Protein sequences were prealigned using HMMalign (Eddy 1998) and the pHMM HLH_ls.hmm from PFAM (http://

pfam.sanger.ac.uk/). The bHLH region was then extensively manually aligned in BioEdit (http://www.mbio.ncsu.edu/ BioEdit/BioEdit.html). Unambiguous aligned positions were used for the subsequent phylogenetic analyses (supplementary fig. \$1, Supplementary Material online). The Jones, Taylor, and Thorton (JTT) model was selected as the best-fitting amino acid substitution model with the Akaike information criterion implemented in ProtTest (Abascal et al. 2005). The maximum likelihood (ML) analyses were done with the program PhyML version 3.0.1 (Guindon and Gascuel 2003) using the JTT model of amino acid substitution, an estimated gamma distribution parameter and an Shimodaira-Hasegawa-like approximate likelihood ratio test. The PHYLIP package version 3.67 (Felsenstein 1989) was used to perform 100 bootstrap replicas of a neighbor joining (NJ) tree based on a JTT distance matrix. PAUP* version 4.0b10 (Swofford 2003) was used to perform 100 bootstrap replicas of a maximum parsimony (MP) tree. The Bayesian analysis was performed with MrBayes version 3.1.2 (http://mrbayes.csit.fsu.edu/): two independent runs were computed for 10 million generations, at which point the standard deviation of split frequencies was less than 0.01; one tree was saved every 100 generations, and 75,000 trees from each run were summarized to give rise to the final cladogram. All trees were visualized using the program Figtree (http://tree.bio.ed.ac.uk/software/ figtree/).

Alignments of the bHLH domain of related sequences were used to build pHMMs with HMMbuild (Eddy 1998). The pHMMs were used to classify proteins not used in the phylogenetic analyses in the plant bHLH subfamilies. The pHMMs were visualized with HMM Logo (Schuster-Bockler et al. 2004).

Detection of Conserved Motifs

The MEME and FIMO software (Bailey and Elkan 1994) were used to discover patterns in the complete amino acid sequences of plant bHLH proteins. Each motif was individually checked so that incorrect or insignificant matches were discarded. The complete plant amino acid sequences were also screened against the PFAM 23.0 database (http://pfam.sanger.ac.uk/).

Results

All the Major Groups of Land Plants Have Large Numbers of bHLH Proteins

Previous phylogenetic analyses of plant bHLH proteins were based on the genome sequences of *A. thaliana* and *O. sativa* (Buck and Atchley 2003; Heim et al. 2003; Toledo-Ortiz et al. 2003; Li et al. 2006b). This provided a useful, but limited, phylogenetic framework for the classification of bHLH proteins in flowering plants (angiosperms). Nevertheless, it provided no insight into the diversity of this family in the earlier diverging groups of land plants. To determine if these subfamilies were angiosperm specific or if they arose earlier in plant evolution and to understand the deeper evolutionary history of this family in plants, we

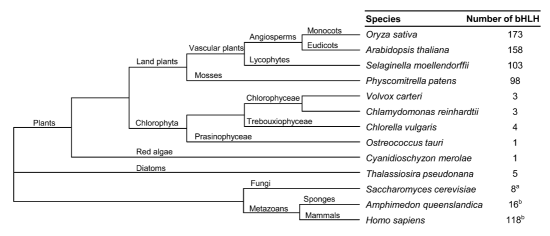


Fig. 1. Phylogenetic relationships of the species used in this study. The total number of bHLH proteins found in the genome of each species is indicated. The cladogram is based on the current view of plant and eukaryotic phylogeny (Baldauf 2003; Lewis and McCourt 2004; Rodríguez-Ezpeleta et al. 2005); ^aRobinson and Lopes (2000); ^bSimionato et al. (2007).

searched for bHLH protein coding sequences in the complete genome of the lycophyte S. moellendorffii, the moss P. patens, the chlorophytes V. carteri, C. reinhardtii, C. vulgaris, and O. tauri, and the red alga C. merolae. These sequences were combined with the previously reported A. thaliana and O. sativa sequences to generate a primary data set consisting of 544 bHLH sequences representing the major evolutionary lineages of plants (fig. 1). We then extended this data set to include proteins from selected eukaryotic groups: the full set of bHLH proteins encoded in the genomes of the diatom T. pseudonana and the fungi S. cerevisiae, plus representative bHLH sequences from the sponge A. queenslandica and H. sapiens (fig. 1).

There are large numbers of bHLH proteins in all species of land plants (embryophytes) sequenced to date. A. thaliana and O. sativa have over 150 bHLH sequences in their genomes, making it the second largest family of transcription factors in angiosperms (Xiong et al. 2005). Approximately 100 bHLH proteins are encoded in the genomes of the lycophyte S. moellendorffii and the moss P. patens (fig. 1). In contrast, we found less than five bHLH-encoding sequences in the genome of each chlorophyte and red alga examined. Other unicellular eukaryotic organisms such as the diatom T. pseudonana and S. cerevisiae also have small numbers (less than 10) of bHLH proteins (fig. 1, Robinson and Lopes 2000). In animals, the sponge A. queenslandica has 16 bHLH-encoding genes, whereas most bilaterians have over 50 genes (Simionato et al. 2007).

Animals and land plants have considerably more bHLH sequences than other eukaryotic organisms. This suggests that the increase in the number of bHLH proteins occurred independently during the evolution of plants and animals.

Key Amino Acid Residues Are Highly Conserved Between Plant and Metazoan bHLH Proteins

To characterize the molecular evolution of plant bHLH proteins, we aligned the retrieved amino acid sequences in the conserved bHLH region (fig. 2, supplementary fig. S1, Supplementary Material online). The first 10–15 amino acids correspond to the basic region, where most interactions

with the DNA are made (Ferré-D'Amaré et al. 1993). Most animal bHLH proteins bind to hexanucleotide sequences (5'-CANNTG-3') known as E-boxes. All E-box-binding bHLH proteins have a glutamic acid (E) residue at position 9 that directly contacts the DNA at the CA nucleotides of the hexanucleotide sequence (Ferré-D'Amaré et al. 1993; Atchley et al. 1999). In plants, the critical E_9 residue is present in 74% of the proteins analyzed (supplementary fig. S1, Supplementary Material online). Other positions of the basic region allow a better discrimination of the target DNA sequences and are easily distinguishable in the major animal bHLH groups (Atchley and Fitch 1997; Ledent and Vervoort 2001; Jones 2004; Atchley and Zhao 2007). Animal group A proteins bind the CAGCTG (or CACCTG) Ebox configuration and have a diagnostic arginine (R) at position 8. Animal group B proteins have a lysine (K) or histidine (H) residue at position 5 and an R at position 13 and bind the CACGTG (or CATGTTG) E-box configuration. In plants, 53% of the bHLH proteins have the characteristic animal group B configuration H₅-E₉-R₁₃ and only 8% have the typical R₈-E₉ found in animal group A. This suggests that most plant bHLH proteins also bind to E-boxes. Indeed, a number of plant bHLH proteins have been shown to bind the CACGTG sequence (e.g., Martínez-García et al. 2000; Toledo-Ortiz et al. 2003; Qian et al. 2007), which is classically known in plants as a G-box motif (Giuliano et al. 1988). Group E animal proteins, that bind N-boxes (CACGCG or CACGAG), have the same H₅-E₉-R₁₃ configuration as group B and a proline (P) at position 6. This configuration is absent in all the 544 plant bHLH proteins analyzed. The remaining animal bHLH groups C and F proteins contain extra PAS and COE domains, not found in plant bHLH proteins, whereas group D proteins are atypical bHLH without a basic domain; 11% of the plant proteins have a conserved Q_5 - A_9 - R_{13} motif (supplementary fig. \$1, Supplementary Material online), not present in animals. This raises the interesting possibility that these proteins bind to a novel target DNA sequence. Other frequent basic amino acids in animal bHLH, such as R in positions 10 and 12, are also highly conserved in plants (73% and 90%, respectively).

Hs Max	12345-67890123 KRAHH-NALERKRR	11111122222-222 45678901234-5678 DHIKDSFHSLR-DSVP	2333-34-44 9123-47-89	555555555566666 012345678901234 SRAQILDKATEYIQYM
AtbHLH067	-RINH-IAVERNRR	RQMNEHINSLR-ALLP	PSYI-QR-GI	QASIVGGAINYVKVL
PpbHLH048	-RENH-IWSERQRR	KGMNYLFSTLR-SLLP	HPT-SK-TI	KSTVVGEIIKYIESL
PpbHLH092	RGVNH-FATERQRR	EYLNEKYQTLR-SLVP	NP-TK-AI	RASIVADAIEYVKEL
AtbHLH017	EPLNH-VEAERQRR	EKLNQRFYALR-SVVP	NI-SK-MI	KASLLGDAISYIKEL
AtbHLH116	PAKNLMAERRRR	KKLNDRLYMLR-SVVP	KI-SK-MI	RASILGDAIDYLKEL
OsbHLH012	SIKNH-VMSERRRW	EKLNEMFLTLK-SLVP	SI-DK-VI	KASSLAETIAYLKEL
AtbHLH032	ASKSH-SEAERRRR	ERINTHLAKLR-SILP	NT-TK-TI	KASLLAEVIQHMKEL
PpbHLH046	PRSKH-SATEQRRR	SKINDRFQMLR-DLVP	HSD-QK-RI	KASFLLEVIEYIQVL
OsbHLH025	NQLQH-MISERKRR	EKLNDSFLALK-AVLP	PG-SK-KI	KTSILIRAREYVKSL
AtbHLH125	KKMKH-RDIERQRR	QEVSSLFKRLR-TLLP	FQYIQ-GK-RS	TSDHIVQAVNYIKDL
AtbHLH015	AAEVH-NLSERKRR	DRINERMKALQ-ELIP	RC-NK-SI	KASMLDEAIEYMKSL
PpbHLH022	-QD-H-IMAERKRR	EKLSQRFIALS-AIVP	GL-KK-MI	KASVLGDAIKYVKTL
PpbHLH042	VRKVHKADREKLRR	DRLNEQFGELAGVLDP	DR-PK-NI	KATILGDSVQVVKDL
PpbHLH014	GGAGNKACREKMRR	DRLNDRFLELGAILEP	GRPPK-TI	KATILSDAVRILTQL
SmbHLH001	ASKNLVSERKRR	KKLNERLYSLR-AIVP	KI-SK-MI	KASIVADAIDYVQEL
SmbHLH042	ATDPQ-SIYARQRR	ERINERLRALQ-GLVP	NG-AK-VI	IVTMLEEAINYVKFL
AtbHLH049	ATNSH-SLAERVRR	EKISERMKFLQ-DLVP	GC-NKVT	KAVMLDEIINYVQSL
AtbHLH082	ATDPH-SIAERLRR	ERIAERMKSLQ-ELVP	NT-NK-TI	KASMLDEIIEYVRFL
PpbHLH027	SKDPQ-SVAARHRR	ERISDRIRVLQ-RLVP	GG-TK-MI	TASMLDEAIHYVKFL
PpbHLH074	SVEPQ-SVAARHRR	KKISERIRVLE-KLIP	GG-NK-MI	TATMLDEAIEYVKFL
SmbHLH103	ANDPQ-SIAARQRR	ERISQRLKILQ-DLVP	NG-SK-VI	LVTMLEKAINYVKFM
AtbHLH144	GSASS-SNNDGKGR	KKMKKMMGVLR-RIVP	GG-EQ-MN	TACVLDEAVQYLKSL
PpbHLH094	ATHPR-SIAERVRR	GKISERMKKLQ-ELVP	NS-DRQTN	TADMLDDAVEYVKQL
AtbHLH134	RSRQA-SSSSRISD	DQITDLISKLR-QSIP	EIRQNRRS-NT-VS	SASKVLQETCNYIRNL
OsbHLH066	AATTT-NKRPRVRR	ERLGERIIALQ-QLVS	PF-GK-SI	TASVLHEALGYIRFL
AtbHLH155	-PGES-SRPRPRDR	QLIQDRIKELR-ELVP	NG-SK-CS	IDSLLERTIKHMLFL

Fig. 2. Alignment of the bHLH domain of representative plant proteins. A representative of each of the 26 subfamilies of plant bHLH is shown, together with the human protein Max, a well-characterized bHLH protein. The shaded boxes indicate the position of the DNA-binding basic region, the two α-helixes, and the variable loop region (Ferré-D'Amaré et al. 1993). The numbering of the amino acids follows (Atchley and Fitch 1997). This is a subset of the full alignment with all the proteins used in this study (supplementary fig. 1, Supplementary Material online).

The α -helices promote the formation of homo- or heterodimeric complexes between bHLH proteins. The structure of a dimer is stabilized by the hydrophobic amino acids isoleucine (I), leucine (L), and valine (V) in conserved positions in the bHLH domain (Ferré-D'Amaré et al. 1993). These positions are highly conserved in animals (Atchley et al. 1999) and in plants (fig. 2). An L residue is present in sites 23 and 64 in 99% and 96% of the plant proteins and in 98% and 80% of the animal proteins, respectively. Sites 54 and 61 have an I, L, or V in 99% and 93% of the plant proteins and in 98% and 93% of the animal proteins, respectively. A conserved P breaks the first helix and starts a loop of variable length (usually six to nine residues in plants). Some loop residues are also conserved: site 47 is K or R in 88% of the plant proteins (supplementary fig. \$1, Supplementary Material online) and 82% of the animal proteins (Atchley et al. 1999).

The high degree of sequence similarity between the bHLH domain of plant and animal proteins, particularly in key DNA-interacting basic amino acids and in helix-stabilizing hydrophobic amino acids, indicates that the molecular structure and transcription factor activity of bHLH proteins are conserved between animals and plants.

Twenty bHLH Subfamilies Found in Flowering Plants Were also Present in Early Land Plants

To understand the evolutionary relationships between plant bHLH proteins, we used conserved regions of the alignment shown in supplementary figure \$1 (Supplementary Material online) to compute phylogenetic trees. An ML analysis shows that proteins from different species cluster together in compact clades with high support values (fig. 3, supplementary fig. S2, Supplementary Material online). MP and NJ analyses support the existence of most of these clades (supplementary fig. S2, Supplementary Material online). Based on the topology of the trees, clade support values, branch lengths, and visual inspection of the bHLH amino acid sequences, we defined 26 subfamilies of bHLH proteins (fig. 3, supplementary fig. S2, Supplementary Material online). These subfamilies are mostly consistent with the groups proposed by previous phylogenetic analyses of plant bHLH using A. thaliana and O. sativa sequences alone (Buck and Atchley 2003; Heim et al. 2003; Toledo-Ortiz et al. 2003; Li et al. 2006b). We adopted the A. thaliana bHLH group nomenclature proposed by Heim et al. (2003) to label these subfamilies, with some modifications, for example, Ib was divided in Ib(1) and Ib(2), and IIIa and IIIc were combined into III(a + c). We also defined three new groups (XIII, XIV, and XV) that include 28 A. thaliana sequences not present in the analysis by Heim et al. Of the 544 proteins analyzed, 10% do not clearly fall in any of the 26 subfamilies and were classified as "orphans" (supplementary fig. S2, Supplementary Material online). These proteins often have a high degree of sequence divergence from other bHLH: This may be due to lineage-specific specializations or, alternatively, they may correspond to pseudogene sequences. One of the A. thaliana groups proposed by Heim and colleagues (group VI,

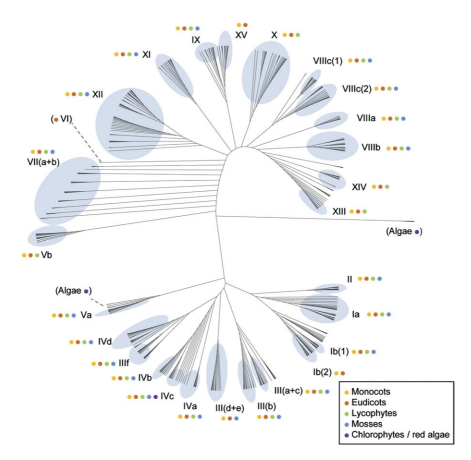


Fig. 3. Twenty subfamilies of bHLH were already established in the common ancestral of vascular plants and mosses. Maximum likelihood analysis of 544 plant bHLH, shown as an unrooted cladogram. The blue balloons delineate the 26 subfamilies of plant bHLH proteins. Colored dots symbolize the species to which the proteins in each group belong (yellow: Oryza sativa [monocot]; red: Arabidopsis thaliana [eudicot]; green: Selaginella moellendorffii [lycophyte]; blue: Physcomitrella patens [moss]; purple: Volvox carteri, Chlamydomonas reinhardtii, Chlorella vulgaris, Ostreococcus tauri, and Cyanidioschyzon merolae (chlorophytes and red algae). A full tree with protein names, proportional branch lengths, and clade support values is given in supplementary fig. S2 (Supplementary Material online).

consisting of only two proteins) falls in this "orphan" category.

Of the 26 plant bHLH subfamilies, 3 include only angiosperm proteins and 23 include angiosperm and lycophyte proteins (fig. 3). Because the last common ancestor of angiosperms and lycophytes lived sometime in the Upper Silurian period before 415 Ma (Kenrick and Crane 1997), this implies that these 23 bHLH subfamilies are at least 415 million years (My) old. Interestingly, 20 of these subfamilies include not only vascular plants but also moss proteins. Given that the oldest evidence for the existence of vascular plants is trilete spores in Upper Ordovician sediments (Steemans et al. 2009), it suggests that these subfamilies are more than 443 My old. A bHLH protein from the chlorophyte algae *O. tauri* is a member of subfamily IVc (fig. 3). This suggests that this subfamily may be over 1 billion years old (Heckman et al. 2001).

A clade composed of *V. carteri, C. reinhardtii,* and *C. vulgaris* bHLH proteins is sister to the proteins in subfamily Va. However, we did not include these chlorophyte proteins into the Va subfamily as this relationship is not strongly supported. Nevertheless, this relationship suggests that subfamily Va is phylogenetically closer to chlorophyte pro-

teins than to any other land plant proteins. Another group of *V. carteri, C. reinhardtii,* and *C. vulgaris* proteins forms a clade that is clearly distinct from other plant proteins (fig. 3); this probably represents a group that evolved among the chlorophytes or, alternatively, was present in the common ancestors of the chlorophytes and land plants but maintained among the chlorophytes and lost in the ancestors of land plants. The only bHLH-encoding gene found in the genome of the red algae *C. merolae* could not be allocated to any chlorophyte or land plant bHLH clade.

In summary, the phylogenetic analysis shows that plant bHLH proteins form 26 distinct subfamilies or evolutionary lineages; 20 of these subfamilies were already present in early land plants 443 Ma, by which time the mosses had diverged from the vascular plants. Despite several rounds of gene duplications and losses in different plant lineages, these subfamilies have been highly conserved throughout plant evolution.

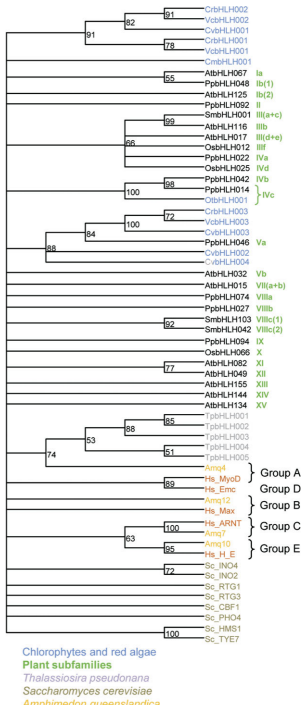
Plant bHLH Proteins Are Monophyletic

The phylogenetic information contained in the 50-60 amino acids of the bHLH allows delimitation of major

evolutionary lineages of proteins in plants but does not allow good resolution of deeper nodes that represent the phylogenetic relationships between different bHLH subfamilies; these basal nodes often have low support values (supplementary fig. S2, Supplementary Material online) and vary when using NJ or MP analyses (data not shown). Similar poor resolution was observed in previous classifications of bHLH proteins in other groups of organisms (Atchley and Fitch 1997; Ledent and Vervoort 2001; Buck and Atchley 2003; Toledo-Ortiz et al. 2003; Li et al. 2006b). Thus, the inter-subfamily relationships shown in figure 3 should be interpreted cautiously. We initially tried to incorporate non-plant bHLH sequences in the ML analysis. However, the large number of proteins and the great evolutionary distances (and consequent high degree of sequence divergence) caused the non-plant proteins to form very long branches, nested within plant clades with no obvious sequence similarity (data not shown). To circumvent this problem, we opted to perform a phylogenetic analysis on a simplified alignment (supplementary fig. S1, Supplementary Material online) that includes chlorophytes, red algae, diatom, and yeast proteins plus representatives of the 26 plant subfamilies and of the higher order metazoan groups (Atchley and Fitch 1997; Simionato et al. 2007).

The deep evolutionary relationships between many proteins were still not resolved: most branches in the Bayesian phylogenetic tree had low support values (fig. 4). However, some close relationships between different plant bHLH subfamilies (fig. 3) were supported by this analysis. For example, subfamilies IVc and Va were probably established in the common ancestors of chlorophyte algae and land plants; subfamily IVb possibly evolved later among land plants from subfamily IVc proteins. Pairs of subfamilies such as VIIIc(1)/VIIIc(2), XI/XII, and III(a + c)/IIIb seem to form monophyletic lineages. Interestingly, the five diatom sequences and a sponge group A protein form a wellsupported clade. Closer examination of the amino acid sequence of the five diatom bHLH proteins reveals that each of these proteins have an arginine in position 8 of the bHLH domain, a defining characteristic of group A proteins (Atchley and Fitch 1997). Although beyond the scope of this study, this suggests that group A might predate the origin of opisthokontes, the eukaryotic lineage that includes fungi and animals.

No clustering of plant proteins with proteins from other eukaryotic organisms is found on the Bayesian tree (fig. 4). The small number of bHLH proteins found in the genomes of different chlorophytes and red algae (fig. 1) suggests that the first plants had one or a few bHLH proteins, from which all modern plant bHLH descended and radiated. This view is consistent with previous analyses that highlighted the distant relationship of angiosperm and animal bHLH proteins (Ledent and Vervoort 2001; Buck and Atchley 2003; Toledo-Ortiz et al. 2003). The lack of discernible phylogenetic relationships between bHLH subfamilies in plants and other eukaryotic organisms supports the hypothesis that plant bHLH proteins are monophyletic.



Amphimedon queenslandica Homo sapiens

Fig. 4. Plant bHLH do not group with other eukaryote bHLH. A Bayesian analysis was performed on an alignment of the bHLH sequence of one representative of each of the 26 subfamilies of plant bHLH, all the chlorophyte and red algae proteins, 5 proteins found in diatom Thalassiosira pseudonana, 8 Saccharomyces cerevisiae proteins, and representatives of 5 major groups of metazoan bHLH in the sponge Amphimedon queenslandica and Homo sapiens. The tree is unrooted. The numbers in the clades are posterior probability values; clades with less than 50% support were collapsed.

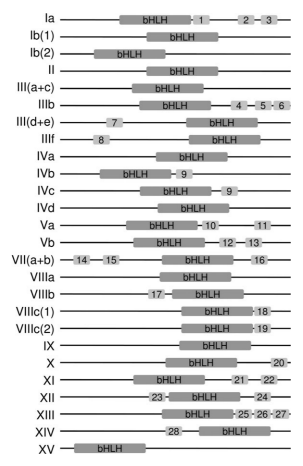


Fig. 5. Non-bHLH amino acid motifs are highly conserved in each bHLH subfamily. An idealized representation of a typical member of each bHLH subfamily is shown, with the bHLH domain and other conserved motifs drawn as shaded boxes. The diagrams are not drawn to scale. The sequences of each motif in individual proteins are given in supplementary table S2 (Supplementary Material online).

Conserved Non-bHLH Motifs Are Present in Most Plant bHLH Subfamilies

The amino acids sequences outside the bHLH region are generally divergent, even in closely related proteins from the same species. Nevertheless, it has been reported that short conserved amino acid motifs are often present in related plant bHLH proteins (Heim et al. 2003; Li et al. 2006b). If our plant bHLH classification were correct, then we expected that such motifs should be conserved within subfamilies. To determine if non-bHLH motifs were conserved throughout plant evolution, we searched for amino acid patterns in our data set of plant bHLH proteins. We found 28 motifs that are represented in both angiosperm and non-angiosperm proteins (supplementary table S2, Supplementary Material online). The relative position of each of these motifs is conserved (fig. 5): most are located C-terminal to the bHLH domain, which itself is generally located toward the C-terminal half of plant proteins. Each of these motifs is only found in members of the same subfamily, apart from motif 9, which is found in both IVb and IVc proteins (fig. 5). None of the 28 conserved motifs corresponds to known domains in the PFAM database. Motifs

14 and 15, present in several proteins of subfamily VII(a + b), overlap with the active phytochrome binding (APB) motif, shown to mediate the binding of several A. thaliana bHLH proteins to phytochrome B (Khanna et al. 2004). Motif 9 (present in IVb and IVc proteins) has a typical leucine zipper (LZ) conformation. The LZ is a dimerization domain that occurs in several regulatory proteins and consists of a periodic repetition of leucine followed by six other residues (Bornberg-Bauer et al. 1998). Several animal bHLH proteins also have an LZ immediately C-terminal to the second helix (Atchley and Fitch 1997). However, its presence in unrelated bHLH proteins suggested a multiple origin of the LZ domain in animal bHLH proteins (Atchley and Fitch 1997; Morgenstern and Atchley 1999). We could not find similarities between the bHLH sequences of IVb/IVc proteins and animal bHLH-LZ proteins. Therefore, it is likely that the acquisition of an LZ motif in bHLH proteins occurred independently in plant and animals. The occurrence of conserved domains outside the bHLH domain strongly supports the classification made on the basis of alignments of the bHLH sequence.

We also queried the PFAM database of protein domains with the 544 plant bHLH proteins and found significant matches to an ACT domain in several unrelated proteins (OsbHLH036, VcbHLH001, CrbHLH002, OsbHLH170, VcbHLH002, and PpbHLH097). The ACT is a regulatory ligand-binding domain found in a diverse group of proteins, mostly metabolic enzymes (Chipman and Shaanan 2001). The occurrence of the ACT domain in plant bHLH proteins was previously reported (Anantharaman et al. 2001), and an ACT-like domain was found to mediate homodimerization of the maize R protein (Feller et al. 2006). Feller et al. also found ACT-like domains in over 30 A. thaliana proteins using low-stringency structure-based searches, but we could not confirm this using our stringent motif-based search methods. The occurrence of the ACT domain in a few proteins from different bHLH subfamilies suggests that the ACT-bHLH association occurred multiple times, possibly through domain-shuffling processes. Such mechanisms have been proposed to play an important role in the evolution of several metazoan bHLH proteins (Morgenstern and Atchley 1999; Ledent and Vervoort 2001).

The presence of highly conserved motifs among proteins of the same subfamily supports the phylogenetic relationships inferred from the bHLH domain sequence alone. The conservation of these extra domains during plant evolution suggests that they are essential for the function of the bHLH proteins in the respective subfamilies. Nevertheless, the presence of the ACT domain in a few unrelated proteins also indicates that domain-shuffling processes may have played a small role in plant bHLH evolution.

Discussion

Our analysis shows that most of the major subfamilies of plant bHLH transcription factors were already present in early land plants, before the divergence of mosses and vascular plants. The recent advent of large-scale sequencing projects has shown that many of the gene families that control angiosperm development were present in early land plants (Floyd and Bowman 2007). However, unlike the bHLH family, many of these families (such as the MIKC MADS-box and TCP transcription factors) diversified after the divergence of lycophytes from the other vascular plants (Floyd and Bowman 2007). We envisage two major alternative hypotheses that would explain the early radiation of the bHLH proteins in plants. The first is that the radiation occurred in parallel with the evolution of multicellularity, before the transition of plants to terrestrial environments. The increase in the number of cell types and morphological complexity brought about by multicellularity would have been programmed by increasingly elaborate gene regulatory networks. bHLH proteins, with their ability to heterodimerize and differentially control gene expression, might have become an ideal tool to assemble such complex regulatory pathways. Consistent with this view is the observation that the first large radiation of the bHLH family in metazoans may have accompanied the evolution of multicellularity (Simionato et al. 2007). A second hypothesis is that the diversification of plant bHLH proteins accompanied the colonization of the land. The challenges faced by plants in a dry terrestrial environment led to the evolution of many novel structures and physiological mechanisms, orchestrated by versatile gene regulatory networks. Distinguishing between these alternatives will require knowledge of the number of bHLH proteins encoded in the genomes of multicellular algae. The sequence of a charophycean (multicellular aquatic algae, sister group to land plants) genome would allow the testing of these hypotheses, but unfortunately only a handful of expressed sequence tags are currently available.

All sequenced genomes of chlorophytes and red algae encode few bHLH proteins (fig. 1). We detected three distinct evolutionary lineages in chlorophytes (fig. 3). One lineage includes both chlorophytes and land plants (subfamilies IVc and IVb), implying that it predates the divergence of chlorophytes from the ancestors of land plants, over 1 billion years ago (Heckman et al. 2001). Interestingly, a characteristic of these two subfamilies is the presence of an LZ motif associated with the bHLH domain. This association has also occurred, independently, in animals. A second lineage of chlorophyte proteins is more similar to subfamily Va than to any other bHLH subfamily, although support for monophyly is poor. A third lineage is distinct from all other plant bHLH proteins and possibly evolved only in chlorophytes. The only bHLH protein found in red algae could not be clearly allocated to any clade. This suggests that none of the 26 subfamilies of plant bHLH proteins was established at the time of divergence of red algae from other plants, 1.5 billion years ago (Yoon et al. 2004). Alternatively, these protein lineages were lost in a C. merolae ancestor but are still present in other red algae; the availability of additional whole-genome sequences from red algae will help to clarify this. However, the small number of bHLH found in all the chlorophytes and red algae examined (fig. 1) and the lack of clear phylogenetic relationships with other eukaryotic bHLH proteins (fig. 4) allows us to confidently deduce that all bHLH proteins found in plants evolved after the primary endosymbiotic event that led to the evolution of plastids and are not represented in other eukaryotic groups.

Plant transcription factor families usually have high expansion rates compared with metazoan families, caused by elevated rates of retention of duplicated genes (Shiu et al. 2005). Accordingly, there are usually many (1-12) proteins per species in each of the 26 plant bHLH subfamilies (supplementary fig. S2, Supplementary Material online), in contrast with the small number (1-4) of genes found in each of the 44 metazoan subfamilies (Ledent and Vervoort 2001; Simionato et al. 2007). Members of the same plant bHLH subfamily are frequently involved in the same biological process (table 1). Usually the functions of these proteins overlap, causing them to be partially or totally redundant (e.g., HEC or BEE proteins). A striking exception comes from three A. thaliana subfamily la proteins, MUTE, SPEECH-LESS, and FAMA: they play nonoverlapping roles in controlling sequential cell fate specification during stomatal differentiation, in a pathway surprisingly similar to metazoan bHLH proteins controlling muscle and neural development (Nadeau 2009; Serna 2009). Interestingly, the function of these proteins seems to be mostly conserved in rice and maize homologs, despite these species having considerably different stomata morphology and differentiation patterns (Liu et al. 2009). Other examples of members of the same bHLH subfamily regulating similar processes in different species are currently known (table 1). A new challenge will be to understand how the function of bHLH proteins has changed during plant evolution. An interesting glimpse comes from subfamily VIIIc(1), where the P. patens proteins PpRSL1 and PpRSL2—the only moss bHLH proteins that have been characterized so far-were shown to be required for the development of rhizoids (Menand et al. 2007). Rhizoids were lost during vascular plant evolution but the two representatives of subfamily VIIIc(1) in A. thaliana (AtRHD6 and AtRSL1) are required for the formation of root hairs, analogous structures to rhizoids with a similar rooting function (Menand et al. 2007). This suggests that these proteins were independently recruited to fulfil similar functions during land plant evolution.

The presence of highly conserved motifs (such as the APB motif in PIF proteins) in the different plant bHLH subfamilies (fig. 5) indicates that the partners of molecular interactions are also conserved. This is particularly exciting because it suggests that protein interactions that are at the base of gene regulatory networks are highly conserved across plants. Several plant bHLH proteins are known to form transcription complexes with MYB proteins (Ramsay and Glover 2005). Although the early evolution of MYB proteins in plants has not been characterized, we found over 30 MYB sequences in the genome of *C. reinhardtii* and more than 150 sequences in *P. patens* (data not shown). Given the large number of both bHLH and MYB proteins in mosses, it is appealing to hypothesize that the bHLH–MYB complex had evolved early in land plant

Table 1. Functionally Characterized bHLH Proteins from Different Plant Species.

Name	bHLH Number	Function	Reference
ubfamily Ia			
AtMUTE	AtbHLH045	Control sequential cell fate	Nadeau (2009); Serna (2009)
AtFAMA	AtbHLH097	specification during stomatal differentiation	
AtSPCH	AtbHLH098		
OsMUTE	OsbHLH055	Control stomata development	Liu et al. (2009)
OsFAMA	OsbHLH051	·	
OsSPCH2	OsbHLH053		
ubfamily Ib(1)	000.121.1033		
RGE1/ZHOUPI	AtbHLH095	Regulates embryonic development	Kondou et al. (2008);
	Attrictions	and endosperm breakdown	Yang et al. (2008)
ubfamily Ib(2)			
OsIRO2	OsbHLH056	Regulates genes involved in Fe uptake under Fe-deficiency conditions	Yuko et al. (2007)
ubfamily III(a + c)			
FIT ,	AtbHLH029	Required for the up-regulation of responses to iron deficiency in Arabidopsis roots	Bauer et al. (2007)
RERJ1	OsbHLH006	Involved in the rice shoot growth inhibition	Kiribuchi et al. (2004)
	Osbriciooo	caused by jasmonic acid	Killbucili et al. (2004)
ubfamily IIIb			
ICE/SCRM	AtbHLH116	Control stomatal development; implicated in	Chinnusamy et al. (2003);
ICE2/SCRM2	AtbHLH033	the cold acclimation response and freezing	Kanaoka et al. (2008);
		tolerance	Fursova et al. (2009)
TalCE41	Wheat ^a	Potential activators of the cold-responsive genes	Badawi et al. (2008)
TalCE87			
Subfamily III(d + e)			
MYC2/JAI1/JIN1	AtbHLH006	Involved in abscisic acid, jasmonic acid and light	Abe et al. (2003);
MTC2/JATT/JINT	ALDITLITOOO	• •	• •
		signalling pathways	Lorenzo et al. (2004);
			Yadav et al. (2005)
AIB	AtbHLH017	Involved in abscisic acid signalling	Li et al. (2007)
PsGBF	Pea ^a	Regulates phenylpropanoid biosynthetic pathway	Qian et al. (2007)
Subfamily IIIf			
TT8	AtbHLH042	Partially redundantly regulate anthocyanin	Nesi et al. (2000);
GL3	AtbHLH001	biosynthesis, trichome and root hair development	Payne et al. (2000);
EGL3	AtbHLH002		Bernhardt et al. (2003); Zhang et al. (2003)
Ra/OSB1	0-611111012	December the such assessed bis such asia mathematic	Ludwig et al. (1989);
	OsbHLH013	Regulate the anthocyanin biosynthetic pathway	
Rb	OsbHLH165		Burr et al. (1996);
Rc	OsbHLH017		Hu et al. (2000);
OSB2	OsbHLH016		Spelt et al. (2000);
Lc	Maize ^a		Sakamoto
IN1	Maize ^a		et al. (2001); Sweeney
An1	Petunia ^a		et al. (2006)
Subfamily IVa			()
NAI1	AtbHLH020	Required for the formation of an ER-derived	Matsushima et al. (2004)
		structure, the ER body	
Subfamily IVc			
ILR3	AtbHLH105	Modulate metal homeostasis and auxin-conjugate metabolism	Rampey et al. (2006)
Subfamily Va			
BIM1	AtbHLH046	Implicated in brassinosteroid signaling	Yin et al. (2005)
BIM2	AtbHLH102		• •
BIM3	AtbHLH141		
Subfamily VII(a + b)			
PIF1/PIL5	Athul Unic	Bind to activated phytochromes and mediate	Castillon et al. (2007);
•	AtbHLH015	· ·	
PIF3	AtbHLH008	light and gibberellin signaling responses; PIF4	de Lucas et al. (2008);
PIF4	AtbHLH009	was recently shown to also mediate plant architecture	Leivar et al. (2008);
PIF5/PIL6	AtbHLH065	responses to high temperatures	Koini et al. (2009)
PIF7	AtbHLH072		
HFR1	AtbHLH026	Mediate both phytochrome and cryptochrome signaling	Duek and Fankhauser (2003)
SPATULA	AtbHLH024	Regulator of carpel margin development; mediator of Heisler et al. (2001);	
ALCATRAZ	AtbHLH073	germination responses to light and temperature Penfield et al. (2005) Required for the formation of a cell layer necessary for Rajani and Sundaresan (2001)	
		fruit dehiscence	
UNE10	AtbHLH016	Involved in the fertilization process	Pagnussat et al. (2005)

Table 1. Continued

Name	bHLH Number	Function	Reference
BP-5	OsbHLH102	Involved in the regulation of amylose synthesis in the rice endosperm	Zhu et al. (2003)
Subfamily VIIIb		·	
HEC1	AtbHLH088	Redundantly control the	Gremski et al. (2007)
HEC2	AtbHLH037	development of the transmitting tract	
HEC3	AtbHLH043	and stigma; each of these proteins can form heterodimers with SPATULA	
LAX	OsbHLH123	Regulator of axillary meristem generation in rice	Komatsu et al. (2003)
INDEHISCENT	AtbHLH040	Required for the differentiation, in the Arabidopsis fruit, of three cell types involved in seed dispersal	Liljegren et al. (2004)
Subfamily VIIIc(1)			
AtRHD6	AtbHLH083	Required for the formation of root hairs	Menand et al. (2007)
AtRSL1	AtbHLH086		
PpRSL1	PpbHLH043	Redundantly required for the	Menand et al. (2007)
PpRSL2	PpbHLH033	development of rhizoids and caulonemata	
Subfamily VIIIc(2)			14 (aaaa)
RSL2	AtbHLH085	Partially redundant and involved in root hair	Yi (2008)
RSL3	AtbHLH084	development	
RSL4	AtbHLH054		
RSL5	AtbHLH139		
Subfamily XI	A+LII 11050	locational to the featilization access	D
UNE12	AtbHLH059 OsbHLH096	Involved in the fertilization process Involved in the responses to phosphate	Pagnussat et al. (2005) Yi et al. (2005)
PTF1	OSDITLINOS	deficiency stress	11 et al. (2005)
Subfamily XII	A - L - L L L L L L L L L	Control of the control of the	5 (2005)
ZCW32/BPE	AtbHLH031	Controls petal size	Szecsi et al. (2006)
BEE1 BEE2	AtbHLH044	Redundant positive regulators of brassinosteroid	Friedrichsen et al. (2002)
BEE3	AtbHLH058	signalling	
CIB1	AtbHLH050	Shown to interact with the blue-light receptor	Liu et al. (2008)
CIB5	AtbHLH063 AtbHLH076	CRY2 and promote floral initiation	Liu et al. (2008)
Subfamily XIII	Attitition	CK12 and promote notal initiation	
LHW	AtbHLH156	Regulates the size of the vascular initial population in the root meristem	Ohashi-Ito and Bergmann (2007)
Subfamily XIV		micial population in the root mension	
SAC51	AtbHLH142	Involved in a spermidine synthase mediated stem elongation process	lmai et al. (2006)
Subfamily XV			
PRE1	AtbHLH136	Proposed to act as positive regulators	Lee et al. (2006)
PRE2	AtbHLH134	of gibberellin signalling	
PRE3	AtbHLH135		
PRE4	AtbHLH161		
PRE5	At3g28857 ^a		
PRE6	At1g26945 ^a		
KIDARI	At1g26945 ^a	Represses light signal transduction; interacts and negatively regulates HFR1	Hyun and Lee (2006)
Orphans			
AMS	AtbHLH021	Required for correct anther development,	Sorensen et al. (2003)
DYT1	AtbHLH022	particularly tapetum development	Zhang et al. (2006)
TDR	OsbHLH005		Li et al. (2006a)
Udt1 MEE8	OsbHLH164	Paguired for early ambruo development	Jung et al. (2005)
MEE8 Fer	AtbHLH108 Tomato ^a	Required for early embryo development Controls iron-uptake responses in roots	Pagnussat et al. (2005) Ling et al. (2002)
Gmyc1	Gerbera ^a	Regulates the expression of an anthocyanin	Elomaa et al. (1998)
delila	Antirrhinum ^a	pathway enzyme Regulates the pattern of anthocyanin pigmentation	Goodrich et al. (1992)
JAF13	Petunia ^a	Regulates the anthocyanin biosynthetic pathway	Quatrochio et al. (1998)
PAR1	At2g42870 ^a	Negatively control growth and metabolic shade	Roig-Villanova et al. (2007)
PAR2	At3g58850 ^a	avoidance responses	

Note.—ER, endoplasmic reticulum.

^a These proteins were not included in our phylogenetic analysis; their classification was based on pHMM scores to subfamily-specific pHMMs.

evolution. The picture that emerges from this and other studies is that much of the complex regulatory machinery that we are currently dissecting in "higher" plants was actually invented by very "simple" ones, early in land plant evolution. The recent reappraisal of algae, bryophytes, and lycophytes as experimental organisms will be an excellent tool to clarify the molecular and biological foundations of many of these processes.

Supplementary Material

Supplementary tables S1 and S2, figures S1 and S2, and data 1 are available at *Molecular Biology and Evolution* online (http://www.mbe.oxfordjournals.org/).

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