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## Formin homology 2 domains occur in multiple contexts in angiosperms

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### Abstract

**Background:** Involvement of conservative molecular modules and cellular mechanisms in the widely diversified processes of eukaryotic cell morphogenesis leads to the intriguing question: how do similar proteins contribute to dissimilar morphogenetic outputs. Formins (FH2 proteins) play a central part in the control of actin organization and dynamics, providing a good example of evolutionarily versatile use of a conserved protein domain in the context of a variety of lineage-specific structural and signalling interactions.

**Results:** In order to identify possible plant-specific sequence features within the FH2 protein family, we performed a detailed analysis of angiosperm formin-related sequences available in public databases, with particular focus on the complete *Arabidopsis* genome and the nearly finished rice genome sequence. This has led to revision of the current annotation of half of the 22 *Arabidopsis* formin-related genes. Comparative analysis of the two plant genomes revealed a good conservation of the previously described two subfamilies of plant formins (Class I and Class II), as well as several subfamilies within them that appear to predate the separation of monocot and dicot plants. Moreover, a number of plant Class II formins share an additional conserved domain, related to the protein phosphatase/tensin/auxilin fold. However, considerable inter-species variability sets limits to generalization of any functional conclusions reached on a single species such as *Arabidopsis*.

**Conclusions:** The plant-specific domain context of the conserved FH2 domain, as well as plant-specific features of the domain itself, may reflect distinct functional requirements in plant cells. The variability of formin structures found in plants far exceeds that known from both fungi and metazoans, suggesting a possible contribution of FH2 proteins in the evolution of the plant type of multicellularity.

### Background

Proteins of the formin family (FH2 proteins) have an important role in the organization of the actin cytoskeleton in organisms as diverse as fungi, slime molds, meta-

zoa and plants (reviewed in [1-3]). Formins have been implicated in processes such as budding of yeast cells, cytokinesis in *Drosophila* and *Caenorhabditis*, and formation of fruiting bodies in *Dictyostelium* (see e.g. [4-9]).

Known mutations affecting formin function in vertebrates cause limb deformity and deafness [10,11], again suggesting a role in morphogenetic processes.

Formins are defined by the presence of a hallmark domain, FH2, accompanied by a proline-rich FH1 domain and often also by other conserved sequence motifs shared only by a subset of FH2 proteins, such as the FH3 domain, a GTPase-binding domain (GBD), or coiled-coil regions [4,12,13]. The FH2 domain, whose structure has been recently determined [14,15], acts as a dimer, nucleating new actin filaments by a novel Arp2/3 independent mechanism, which has been well documented in both yeast and metazoans [16-19]. This provides a mechanistic basis for the observed morphogenetic role of formins. The proline-rich FH1 motif binds profilin and contributes to the actin-nucleating activity and its regulation [20,21]. Domains outside FH1 and FH2 provide a variety of "interfaces" for integration of the actin nucleating module into cellular regulatory networks (reviewed in [1,2,22]). For instance, a subfamily of Diaphanous-related formins may be mediating the effects of Rho class small GTPases on actin assembly and dynamics via a specific conserved domain [23,24]. Other formins communicate with universal "adaptor" domains such as SH3 or WW (see e.g. [25-28], and at least indirectly even with the microtubule cytoskeleton [29-31].

Members of the formin family have been found also in higher plants, both experimentally [32] and by a bioinformatic approach [2,33]. Formin-related sequences encoded by the complete *Arabidopsis* genome can be divided into two distinct subfamilies. One of them (Class I) contains mostly proteins with putative membrane insertion signals, and often with extensin-like proline-rich stretches in the predicted extracytoplasmic domain. This suggests a possible plant-specific mechanism of cytoskeleton-membrane connection, or even transmembrane anchorage of the cytoskeleton to the cell wall in case of plasmalemma-localized formins [33], which is now supported also by experimental data [32,34]. No such motifs – and no conserved domains whatsoever besides FH1 and FH2 – were described in Class II formins so far. However, detailed analysis of the N-termini of *Arabidopsis* formins could not have been performed on the basis of a genome annotation where the majority of Class II formin genes appeared to be N-terminally truncated [2].

Here we report the results of a detailed structural and phylogenetic analysis of a collection of angiosperm formin-related sequences currently available in public databases, including the nearly complete rice genome. Using a comparative approach, we were able to refine the current annotation of the *Arabidopsis* formin-related genes, and to identify a novel N-terminal conserved domain shared by

the majority of plant Class II formins. Moreover, the structure of this domain, which is related to the conserved and well-characterized protein phosphatase/tensin/auxilin fold, suggests a second possible plant-specific mechanism for integrating the formin-associated actin nucleation complexes into the cellular context.

## Results and discussion

### **An inventory of *Arabidopsis* formins**

An exhaustive *in silico* search of the *Arabidopsis* genome has identified 22 occurrences of the FH2 domain in total of 21 annotated loci, described previously as AtFH1 to AtFH21 [2] (see Table 1); the AtFH15 locus appears to encode two FH2 domains. Most of the formin-related genes reside at positions interspersed throughout the *Arabidopsis* genome. However, 5 loci (AtFH15, AtFH16, AtFH19, AtFH20 and AtFH21) form a tight cluster on chromosome 5.

At least partial cDNA sequences are available for 17 of the 21 loci. Expression data from the NASC microarray collection [35] suggest that the remaining genes are significantly expressed at least under some circumstances, although for AtFH12 and AtFH21 only very low transcript levels have been detected. Complete cDNAs have been sequenced only for AtFH1, AtFH5, AtFH9 and AtFH10, while the current genome annotation of the remaining genes is based mainly on automated splicing prediction. Such predictions are known to be error-prone, and inclusion of cDNA data can improve the annotation considerably [36,37]. Homology among members of a large gene family can be used as an additional guide for identification of mispredicted intron-exon boundaries (see e.g. [38]). Taking into account both cDNA data and homology, we have found that the current annotation of 10 of the remaining 17 loci appears to be incorrect, and suggested modifications, although we still could not reliably identify the N-terminal exons of AtFH3 and AtFH12 (see Table 1 and Additional file 1).

Most of the suggested modifications represent extension of exons, or inclusion of extra exons that restore missing portions of the conserved FH2 domain, or of N-terminal regions of homology shared by multiple family members and revealed in TBLASTN searches (see Methods). This is the case of AtFH12, AtFH13, AtFH14, AtFH17, AtFH18 and AtFH20. In the case of AtFH20, the N-terminal portion of FH2, which was missing in the original prediction, was found in a neighboring gene (At5g07750), which therefore probably represents a part of the AtFH20 locus erroneously annotated as a separate gene.

In three loci (AtFH3, AtFH4 and AtFH16) attempts to restore a missing exon within FH2 revealed probable frameshift errors in the genome sequence. For AtFH3, we

**Table 1: FH2 proteins encoded by the *Arabidopsis thaliana* genome**

Gene	AGI locus	cDNA (complete) <sup>a</sup>	cDNA (partial) <sup>a,b</sup>	No. of coding exons	Protein/ORF sequence <sup>a</sup>	Class	Domain structure <sup>c</sup>	Synonyms	References and notes
<b>AtFH1</b>	At3g25500	AF174427.1	-	4	AAF14548.1	Ia	A	AFH1, AtFORMIN8	[2,32,33]
<b>AtFH2</b>	At2g43800	NA	AV545883.1 BU635310.1	4	AAB64026.1	Ia	A	AtFORMIN2, AtORF1	[2,13,33]
<b>AtFH3</b>	At4g15190 At4g15200	NA	AV557654.1	6	BK004092 <sup>d</sup>	Ic	A?	AtFORMIN3	[2,33]; 5' truncated, presumed genomic sequence error
<b>AtFH4</b>	At1g24150	NA	AI998115.1 BE526568.1	2	BK004101 <sup>d</sup>	Ie	A	AtFORMIN4	[2,33]; presumed genomic sequence error
<b>AtFH5</b>	At5g54650	AY042801.1	-	6	AAK68741.1	Ic	A	AtFORMIN5	[2,33]
<b>AtFH6</b>	At5g67470	NA	F19772.1 BX829650.1	4	BAB08455.1	Ib	A	AtFORMIN6	[2,33]
<b>AtFH7</b>	At1g59910	NA	AV542102.1 BX817601.1	2	AAD39332.1	Ie	B	AtFORMIN7	[2,33]
<b>AtFH8</b>	At1g70140	NA	AY050956.2	2	AAB61101.1	Ie	A	AtFORMIN1, AtORF2	[2,13,33]
<b>AtFH9</b>	At5g48360	AK118458.1	-	4	BAC43066.1	I	A	-	[2]
<b>AtFH10</b>	At3g07540	AY050396.1	-	3	AAK91412.1	I	A	-	[2]
<b>AtFH11</b>	At3g05470	NA	NA	4	AAF64546.1	Id	A	-	[2]
<b>AtFH12</b>	At1g42980	NA	NA	11	BK004100 <sup>d</sup>	II	C	-	[2]; 5' truncated
<b>AtFH13</b>	At5g58160	NA	N65121.1 AA394985.1	14	BK004099 <sup>d</sup>	II	D	-	[2]; alternative splicing
<b>AtFH14</b>	At1g31810	NA	AV528978.1 AV548682.1	17	BK004098 <sup>d</sup>	II	D	-	[2]
<b>AtFH15<sup>a</sup></b>	At5g07650	NA	AV543211.1	13	BK004097 <sup>d</sup>	II	E	-	[2]; alternative splicing or two genes
<b>AtFH15<sup>b</sup></b>	At5g07650	NA	NA	12	BK004096 <sup>d</sup>	II	C	-	
<b>AtFH16</b>	At5g07770	NA	AV526999.1 AV527418.1 AV520899.1 AV529184.1 AV520610.1	16	BK004095 <sup>d</sup>	II	B	-	[2]; presumed genomic sequence error
<b>AtFH17</b>	At3g32400	NA	NA	16	BK004094 <sup>d</sup>	II	C	-	[2]
<b>AtFH18</b>	At2g25050	NA	AV558611.1	16	BK004093 <sup>d</sup>	II	D	-	[2]
<b>AtFH19</b>	At5g07780	NA	AI998622.1	14	BAB09942.1	II	E	-	[2]
<b>AtFH20</b>	At5g07740 At5g07750	NA	AV558046.1 BE525429.1 AV554850.1	15	BK004102 <sup>d</sup>	II	D	-	[2]; alternative splicing
<b>AtFH21</b>	At5g07760	NA	NA	24	BAB11455.1	II	F	-	[2]

Notes: <sup>a</sup>GenBank/EMBL/DDBJ accession numbers; <sup>b</sup>selected cDNAs/ESTs providing maximal coverage of the locus, given only if complete cDNA not available; <sup>c</sup>see Figure 3; <sup>d</sup>deposited in the Third Party Annotation section of GenBank as a part of this study, see also Additional file 1; NA – not available.

have included an internal exon exhibiting homology to the very closely related sequenced AtFH5 cDNA and a 3' extension of the ORF that was suggested by an alternative GenScan prediction (see Methods) as a new exon preceded by an unusually short (11 bp) intron. Since the smallest (protozoan) introns reported so far are 13 bp short and the majority of short introns in plants exceed the length of 30 bp [39], we suspect that the presumed

"intron", which would contain a stop codon, may in fact be a part of a contiguous exon disrupted by omission of 1 base. Also in the internal exon, an extra base must be introduced in order to maintain the reading frame. Since both suspect areas are extremely GC-rich (and therefore notoriously difficult to sequence), we believe that an error in the genomic data is a likely explanation in both cases. For AtFH4, the original annotation predicts an intron dis-

rupted by an in-frame stop codon at the position of the conserved G-N-X-M-N motif. However, this intron is poorly supported by WebGene and GenScan predictions and apparently not spliced in a sequenced cDNA, which contains an extra base in this area and restores the reading frame within a highly conserved portion of FH2. For AtFH16, the splicing pattern could have been inferred with a reasonable confidence, since most of the locus is covered by cDNAs. However, a contiguous reading frame throughout the conserved FH2 domain can be maintained only by inserting an extra base in a GC-rich area not covered by cDNA, again suggesting a sequencing error.

In case of AtFH15, the locus with two FH2 domains, a sequenced cDNA ends by a stretch corresponding to a presumed intron, which contains multiple stop-codons. We believe that the locus either represents two related neighboring genes (further referred to as AtFH15a and AtFH15b), or produces multiple gene products by alternative splicing. Evidence for cDNA-supported alternative splicing, documented in metazoan formins [40], has been found also for AtFH13 and AtFH20, as well as for two rice formin homologues (see below and Additional file 2). In the following structural analysis only the longest predicted proteins have been taken into account.

Nine of the eleven mispredicted loci code for formins previously classified as Class II, while most Class I formins appear to be predicted correctly. The complex structure of Class II formin genes, which possess substantially more intron-exon boundaries than their Class I relatives, may be sufficient to explain the difference [36]. Moreover, both mispredicted Class I loci appear to contain sequencing errors, and one of them, AtFH3, is expressed almost exclusively in pollen according to the results of a recent microarray analysis [35,41]. This narrow tissue specificity might be associated with a modification of the "house-keeping" splicing apparatus, whose function has been so far characterized mainly on the basis of data from vegetative tissues. We therefore believe that the difficulties in predicting AtFH3 structure (including the lack of a reliable N-terminus) may partly reflect the particular expression pattern of this gene.

#### **Phylogeny of the plant FH2 proteins**

Previous phylogenetic analyses of plant formins [2,33] included only *Arabidopsis* data. We have used the re-annotated *Arabidopsis* formin sequences as a query for identifying genes encoding FH2 proteins from available angiosperm sequences in the public databases, including the nearly complete rice genome and a recently published large collection of rice cDNAs [42]. At least partial cDNA or genomic sequences corresponding to 79 putative formin-related genes from cotton, soybean, barley, tomato, trefoil, alfalfa, tobacco, rice, pea, sorghum,

potato, wheat, grapevine and maize have been found (Additional files 2 to 4).

Complete FH2 domain sequence could have been reconstructed for 29 of the non-*Arabidopsis* sequences. This subset, together with 22 *Arabidopsis* FH2 domains and a selection of fungal, slime mold and metazoan formins, has been used to construct an unrooted phylogenetic tree (Fig. 1) using the NJ method [43]. For the remaining sequences, closest neighbors have been determined using the BLAST algorithm (Table 2). All of the plant FH2 domains studied so far, including the incomplete ones, can be unequivocally assigned to one of the two previously proposed classes [2]; Table 2). Also the overall domain composition and domain order of available complete plant formins – i.e. sequences outside FH2 – reflects rather well the dichotomy between Class I and Class II formins (see below).

The presence of two classes of formins appears to be a general feature of plants. However, although several subclasses containing representatives of more than one species can be distinguished within the two classes (see Fig. 1 – branches Ia to Ie, IIa and IIb, Table 1, Table 2), only occasionally true orthology between genes from different species was established. A similar pattern has been previously observed for another large plant gene family encoding the plethora of phospholipase D isoforms [38]. Very closely related proteins that might represent true orthologues (Table 2, sequences in bold) were found mostly within limited taxonomical groups such as the grasses (barley, rice, sorghum, wheat), the legumes (soybean, alfalfa), or *Solanaceae* (tobacco, potato and tomato). The observed pattern of paralog distribution may suggest that a number of gene duplications or polyploidization events occurred relatively recently compared to the separation of the angiosperm lineages included in the analysis. An extreme example of such a recent gene multiplication is presented by the well-defined subgroup of Class II genes corresponding to the clustered loci on the *Arabidopsis* chromosome V (marked by asterisk in Fig. 1).

#### **Structural diversity of the plant FH2 domains**

Surprisingly, major differences between Class I and Class II formins have been found within the relatively well-conserved FH2 domain. Hallmark features for both classes of plant formins can be identified already at the level of amino acid sequence in the C-terminal portion of the FH2 domain, which is less conserved than the central area around the G-N-X-M-N motif (see Additional file 5). While Class I formins contain a consensus V/I-R-D-F-L motif about 170–190 aa from the conserved core, Class II proteins possess a signature M-H-Y-L/Y-C-K, located usually 31 aa downstream of the central motif.

**Table 2: Phylogenetic relationships of non-arabidopsis plant FH2 proteins**

Gene	Organism	Class	Closest relatives	Gene	Organism	Class	Closest relatives
<b>BvFH1</b>	<i>Beta vulgaris</i>	I	AtFH5 107/202 (52%)	<b>NbFH6</b>	<i>N. benthamiana</i>	lc	<b>NbFH1</b> 47/48 (97%); <b>AtFH5</b> 41/48 (85%)
<b>GaFH1</b>	<i>Gossypium arboreum</i>	lc	AtFH5 131/172 (76%)	<b>NbFH7</b>	<i>N. benthamiana</i>	la	<b>NtFH2</b> 154/181 (85%); <b>AtFH1</b> 60/188 (31%)
<b>GaFH2</b>	<i>G. arboreum</i>	la	MtFH1 116/171 (67%); <b>AtFH1</b> 105/169 (62%)	<b>NtFH1</b>	<i>Nicotiana tabacum</i>	la*	<b>StFH4</b> 177/218 (81%); <b>AtFH1</b> 216/293 (73%)
<b>GhFH1</b>	<i>Gossypium hirsutum</i>	I	<b>SpFH1</b> 89/109 (81%); <b>AtFH1</b> 144/197 (73%)	<b>NtFH2</b>	<i>N. tabacum</i>	la*	<b>AtFH1</b> 313/474 (66%)
<b>GhFH2</b>	<i>G. hirsutum</i>	lb	LeFH1 165/227 (72%); <b>AtFH6</b> 158/221 (71%)	<b>NtFH3</b>	<i>N. tabacum</i>	lc	<b>AtFH5</b> 109/156 (69%)
<b>GmFH1</b>	<i>Glycine max</i>	II*	<b>MtFH5</b> 113/133 (84%); <b>AtFH13</b> 182/278 (65%)	<b>NtFH4</b>	<i>N. tabacum</i>	II	<b>GmFH1</b> 105/165 (63%); <b>AtFH13</b> 97/195 (49%)
<b>GmFH2</b>	<i>G. max</i>	IIa	<b>OsFH3</b> 136/161 (84%); <b>AtFH14</b> 132/161 (81%)	<b>NtFH5</b>	<i>N. tabacum</i>	lc	<b>StFH5</b> 58/87 (66%); <b>AtFH5</b> 36/50 (72%)
<b>GmFH3</b>	<i>G. max</i>	II	<b>AtFH18</b> 125/153 (81%)	<b>NtFH6</b>	<i>N. tabacum</i>	I	<b>StFH5</b> 71/150 (47%); <b>AtFH3</b> 56/153 (36%)
<b>GmFH4</b>	<i>G. max</i>	la	<b>MtFH1</b> 98/107 (91%); <b>AtFH1</b> 88/105 (83%)	<b>OsFH1</b>	<i>Oryza sativa</i>	I*	<b>SpFH1</b> 180/205 (87%); <b>AtFH1</b> 299/442 (67%)
<b>GmFH5</b>	<i>G. max</i>	I	<b>StFH4</b> 77/125 (61%); <b>AtFH1</b> 75/112 (66%)	<b>OsFH2</b>	<i>O. sativa</i>	Id*	<b>AtFH1</b> 227/409 (55%)
<b>GmFH6</b>	<i>G. max</i>	lc	LeFH3 103/141 (73%); <b>AtFH5</b> 97/142 (68%)	<b>OsFH3</b>	<i>O. sativa</i>	IIa*	<b>GmFH2</b> 136/160 (85%); <b>AtFH14</b> 303/461 (65%)
<b>HvFH1</b>	<i>Hordeum vulgare</i>	II	<b>OsFH5</b> 190/217 (87%); <b>AtFH18</b> 146/217 (67%)	<b>OsFH4</b>	<i>O. sativa</i>	lb*	<b>OsFH8</b> 373/495 (75%); <b>AtFH6</b> 292/486 (60%)
<b>HvFH2</b>	<i>H. vulgare</i>	I	<b>OsFH13</b> 144/168 (85%); <b>AtFH6</b> 114/164 (69%)	<b>OsFH5</b>	<i>O. sativa</i>	II*	<b>HvFH1</b> 190/217 (87%); <b>AtFH20</b> 259/445 (58%)
<b>HvFH3</b>	<i>H. vulgare</i>	I	<b>OsFH13</b> 177/237 (74%); <b>AtFH6</b> 121/227 (53%)	<b>OsFH6</b>	<i>O. sativa</i>	II*	<b>AtFH18</b> 265/382 (69%)
<b>HvFH4</b>	<i>H. vulgare</i>	I*	<b>OsFH1</b> 256/316 (81%); <b>AtFH1</b> 198/289 (68%)	<b>OsFH7</b>	<i>O. sativa</i>	IIb*	<b>HvFH5</b> 139/163 (85%); <b>AtFH18</b> 264/428 (61%)
<b>HvFH5</b>	<i>H. vulgare</i>	IIb	<b>OsFH7</b> 132/153 (86%); <b>AtFH18</b> 103/149 (69%)	<b>OsFH8</b>	<i>O. sativa</i>	lb*	<b>OsFH4</b> 356/486 (73%); <b>AtFH6</b> 287/433 (66%)
<b>HvFH6</b>	<i>H. vulgare</i>	I	<b>OsFH1</b> 138/154 (89%); <b>AtFH1</b> 91/151 (60%)	<b>OsFH9</b>	<i>O. sativa</i>	Id*	<b>AtFH11</b> 220/404 (54%)
<b>HvFH7</b>	<i>H. vulgare</i>	lc*	<b>OsFH11</b> 156/225 (69%); <b>AtFH5</b> 147/226 (65%)	<b>OsFH10</b>	<i>O. sativa</i>	lc*	<b>OsFH11</b> 235/456 (51%); <b>AtFH5</b> 243/463 (52%)
<b>HvFH8</b>	<i>H. vulgare</i>	I	<b>SpFH2</b> 98/120 (81%); <b>AtFH1</b> 94/186 (50%)	<b>OsFH11</b>	<i>O. sativa</i>	lc*	<b>AtFH5</b> 265/481 (55%)
<b>HvFH9</b>	<i>H. vulgare</i>	I	<b>OsFH11</b> 70/129 (54%); <b>AtFH5</b> 47/110 (42%)	<b>OsFH12</b>	<i>O. sativa</i>	II*	<b>AtFH20</b> 209/395 (52%)
<b>HvFH10</b>	<i>H. vulgare</i>	I	<b>OsFH14</b> 145/196 (73%); <b>AtFH1</b> 86/193 (44%)	<b>OsFH13</b>	<i>O. sativa</i>	I	<b>HvFH2</b> 144/168 (85%); <b>AtFH6</b> 224/443 (50%)
<b>LeFH1</b>	<i>Lycopersicon esculentum</i>	lb*	<b>StFH1</b> 211/235 (89%); <b>AtFH6</b> 318/455 (69%)	<b>OsFH14</b>	<i>O. sativa</i>	I*	<b>AtFH1</b> 187/381 (49%)
<b>LeFH2</b>	<i>L. esculentum</i>	IIa*	<b>StFH6</b> 140/155 (90%); <b>AtFH14</b> 268/349 (76%)	<b>OsFH15</b>	<i>O. sativa</i>	I*	<b>OsFH1</b> 243/425 (57%); <b>AtFH1</b> 228/416 (54%)
<b>LeFH3</b>	<i>L. esculentum</i>	lc*	<b>StFH5</b> 189/196 (96%); <b>AtFH5</b> 292/406 (71%)	<b>OsFH16</b>	<i>O. sativa</i>	I*	<b>SbFH2</b> 148/167 (88%); <b>AtFH4</b> 234/441 (53%)
<b>LeFH4</b>	<i>L. esculentum</i>	I	<b>NtFH1</b> 164/258 (63%); <b>AtFH1</b> 131/246 (53%)	<b>PsFH1</b>	<i>Pisum sativum</i>	lc*	<b>AtFH5</b> 275/439 (62%)
<b>LeFH5</b>	<i>L. esculentum</i>	la	<b>AtFH1</b> 155/222 (69%)	<b>SbFH1</b>	<i>Sorghum bicolor</i>	II	<b>HvFH1</b> 105/126 (83%); <b>AtFH18</b> 93/147 (63%)
<b>LeFH6</b>	<i>L. esculentum</i>	II	<b>OsFH5</b> 95/144 (65%); <b>AtFH20</b> 99/178 (55%)	<b>SbFH2</b>	<i>S. bicolor</i>	Ie	<b>OsFH16</b> 137/167 (82%); <b>AtFH7</b> 107/167 (64%)
<b>LeFH7</b>	<i>L. esculentum</i>	I	<b>NbFH1</b> 66/98 (67%); <b>AtFH5</b> 62/108 (57%)	<b>SpFH1</b>	<i>Sorghum propinquum</i>	I	<b>OsFH1</b> 180/205 (87%); <b>AtFH1</b> 146/205 (71%)
<b>LjFH1</b>	<i>Lotus japonicus</i>	Id	<b>AtFH11</b> 111/140 (79%)	<b>SpFH2</b>	<i>S. propinquum</i>	I	<b>OsFH15</b> 172/204 (84%); <b>AtFH1</b> 128/205 (62%)
<b>MtFH1</b>	<i>Medicago truncatula</i>	la	<b>GmFH4</b> 98/106 (92%); <b>AtFH1</b> 215/299 (71%)	<b>SpFH3</b>	<i>S. propinquum</i>	I	<b>OsFH1</b> 93/111 (83%); <b>AtFH1</b> 72/91 (79%)
<b>MtFH2</b>	<i>M. truncatula</i>	la	<b>AtFH1</b> 143/182 (78%)	<b>StFH1</b>	<i>Solanum tuberosum</i>	lb*	<b>LeFH1</b> 211/235 (89%); <b>AtFH6</b> 186/248 (75%)

**Table 2: Phylogenetic relationships of non-arabidopsis plant FH2 proteins (Continued)**

<b>MtFH3</b>	<i>M. truncatula</i>	lc	NbFH1 135/189 (71%); AtFH5 128/188 (68%)	<b>StFH2</b>	<i>S. tuberosum</i>	la	<b>GhFH1</b> 127/148 (85%); AtFH1 143/208 (68%)
<b>MtFH4</b>	<i>M. truncatula</i>	ll	GmFH3 129/181 (71%); AtFH13 126/191 (65%)	<b>StFH3</b>	<i>S. tuberosum</i>	ll	OsFH6 156/216 (72%); AtFH18 150/216 (69%)
<b>MtFH5</b>	<i>M. truncatula</i>	ll	GmFH1 102/133 (76%); AtFH18 83/166 (50%)	<b>StFH4</b>	<i>S. tuberosum</i>	la	<b>NtFH1</b> 186/217 (85%); AtFH1 149/211 (70%)
<b>MtFH6</b>	<i>M. truncatula</i>	ll	AtFH20 76/124 (61%)	<b>StFH5</b>	<i>S. tuberosum</i>	lc	<b>LeFH3</b> 189/196 (96%); AtFH5 152/229 (66%)
<b>NbFH1</b>	<i>Nicotiana benthamiana</i>	lc	<b>LeFH3</b> 225/241 (93%); AtFH5 189/240 (78%)	<b>StFH6</b>	<i>S. tuberosum</i>	lla	<b>LeFH2</b> 153/155 (98%); AtFH14 123/154 (79%)
<b>NbFH2</b>	<i>N. benthamiana</i>	lb	<b>LeFH1</b> 138/150 (92%); AtFH6 100/162 (61%)	<b>TaFHI</b>	<i>Triticum aestivum</i>	l	<b>HvFH4</b> 178/196 (90%); AtFH1 145/192 (75%)
<b>NbFH3</b>	<i>N. benthamiana</i>	lc	<b>StFH5</b> 100/124 (80%); AtFH5 91/143 (63%)	<b>VvFHI</b>	<i>Vitis vinifera</i>	l	OsFH1 114/185 (61%); AtFH1 104/192 (54%)
<b>NbFH4</b>	<i>N. benthamiana</i>	l	AtFH6 115/278 (41%)	<b>ZmFHI</b>	<i>Zea mays</i>	l	OsFH1 119/154 (77%); AtFH1 81/154 (52%)
<b>NbFH5</b>	<i>N. benthamiana</i>	l	NtFH2 101/146 (69%); AtFH1 77/140 (55%)				

As "closest relatives", sequences with best match altogether and best *Arabidopsis* match are shown (defined as identity at least 80 % across at least 100 amino acids, if available, or best BLAST score). Numbers denote fraction of identical amino acids throughout the length of sequence analysed (putative orthologues with more than 80 % identity in bold). Sequences marked by an asterisk are included in Fig. 1. The partial sequences MtFH4, MtFH5 and MtFH6 might correspond to different parts of the same gene. For database references and protein sequence predictions see Additional files 2 to 4.

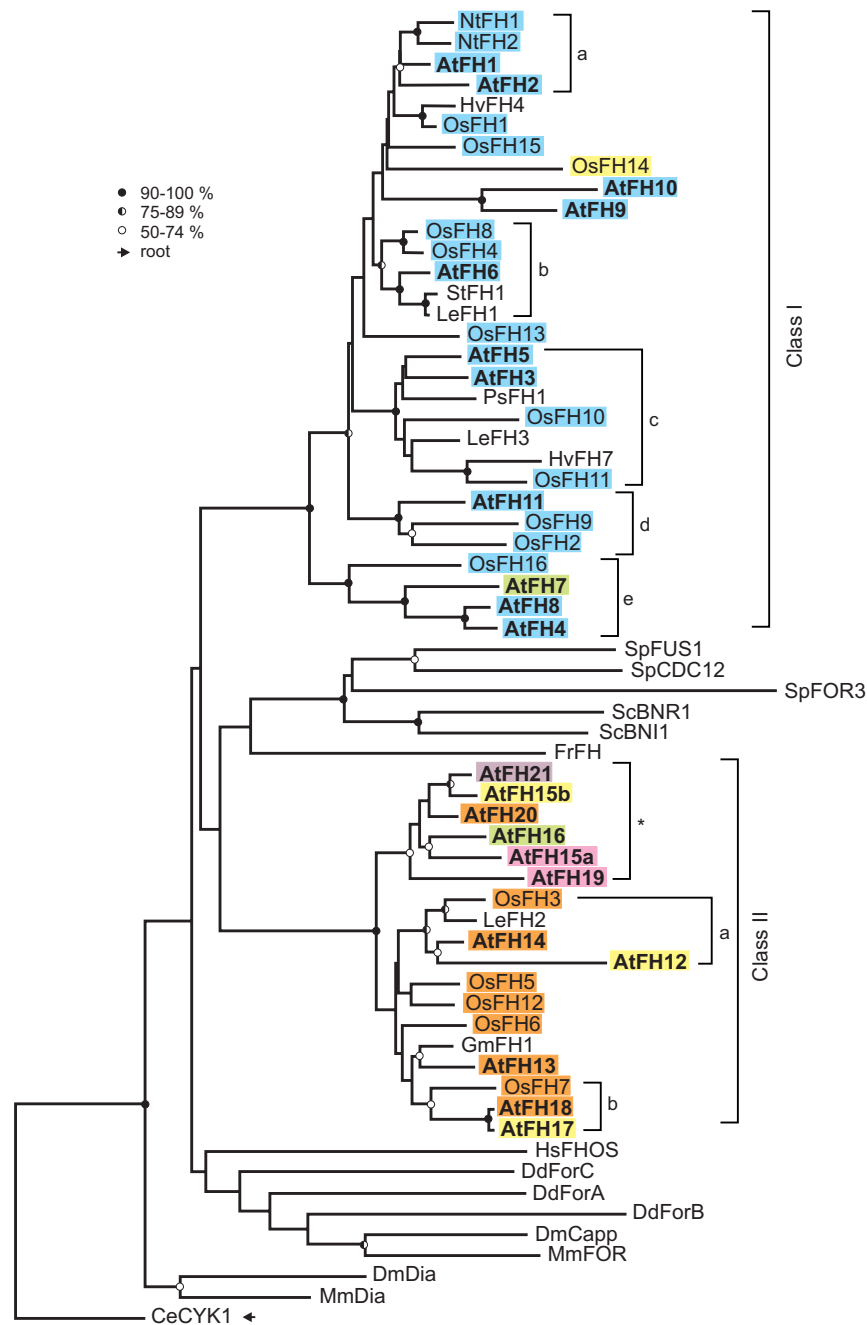
Data on 3-dimensional structure of the FH2 domain have been published recently for the yeast Bni1p formin [14], see also Fig. 2). Its FH2 domain folds into a structure consisting from N-terminal "lasso", connected by a predominantly helical linker to the globular "knob" region, which is followed by a coiled-coil assembly of three  $\alpha$ -helices and a "post" domain, again predominantly  $\alpha$ -helical. The protein can dimerize through interaction of the lasso and post domains, producing flexible ring-like "head to tail" dimers with putative actin-binding sites on the part of the inner surface of the ring, provided by the knob. Residues directly participating in the actin-binding site have been identified by site-directed mutagenesis of selected positions conserved among Bni1p and metazoan formins of the Diaphanous type. Mutants unable to form dimers do not nucleate actin in vitro, and proteolytically cleaved "hemidimers" have been shown to block barbed end elongation, acting as capping proteins rather than nucleators [14,15]. Apparently, dimerization can add another dimension to the diversity of plant formins, since, in theory, the 22 *Arabidopsis* FH2 domains could produce up to 484 different homo- and heterodimers. However, the actual number will be lower, since formins do exhibit tissue-specific expression patterns, as documented by analysis of available expression data [35]. Moreover, structural differences may prevent heterodimerization of some protein pairs.

Comparison of *Arabidopsis* FH2 domain sequences with the sequence of Bni1p (see Additional file 5) revealed surprising plant-specific features in Class I formins (Fig. 2). All Class I formins except AtFH9 and AtFH10 have a small or non-polar amino acid at the position corresponding to

K1639 of Bni1p, a conserved Lys residue contributing to the actin binding site and required for efficient nucleation [14]. However, all Class I proteins have a relatively large insertion (12–54 aa) in the vicinity (position 1620 of Bni1p), suggesting an alternative construction of the actin-binding site. It is worth noting that the shortest insertions were found in AtFH9 and AtFH10, i. e. the only Class I formins that have kept – or, more likely, restored – the consensus Lys residue. On the other hand, AtFH9 and AtFH10, which are mutually closely related (see Fig. 1), exhibit deviations from the Bni1p/Diaphanous consensus in portions of the molecule that are involved in dimerization (a deletion in the lasso of AtFH9, altered structure of the post in AtFH10). It is tempting to speculate that these alterations might result in a restriction of (hetero)dimerizing capability of AtFH9 and AtFH10, although we cannot, at present, predict which dimers will be preferred or excluded.

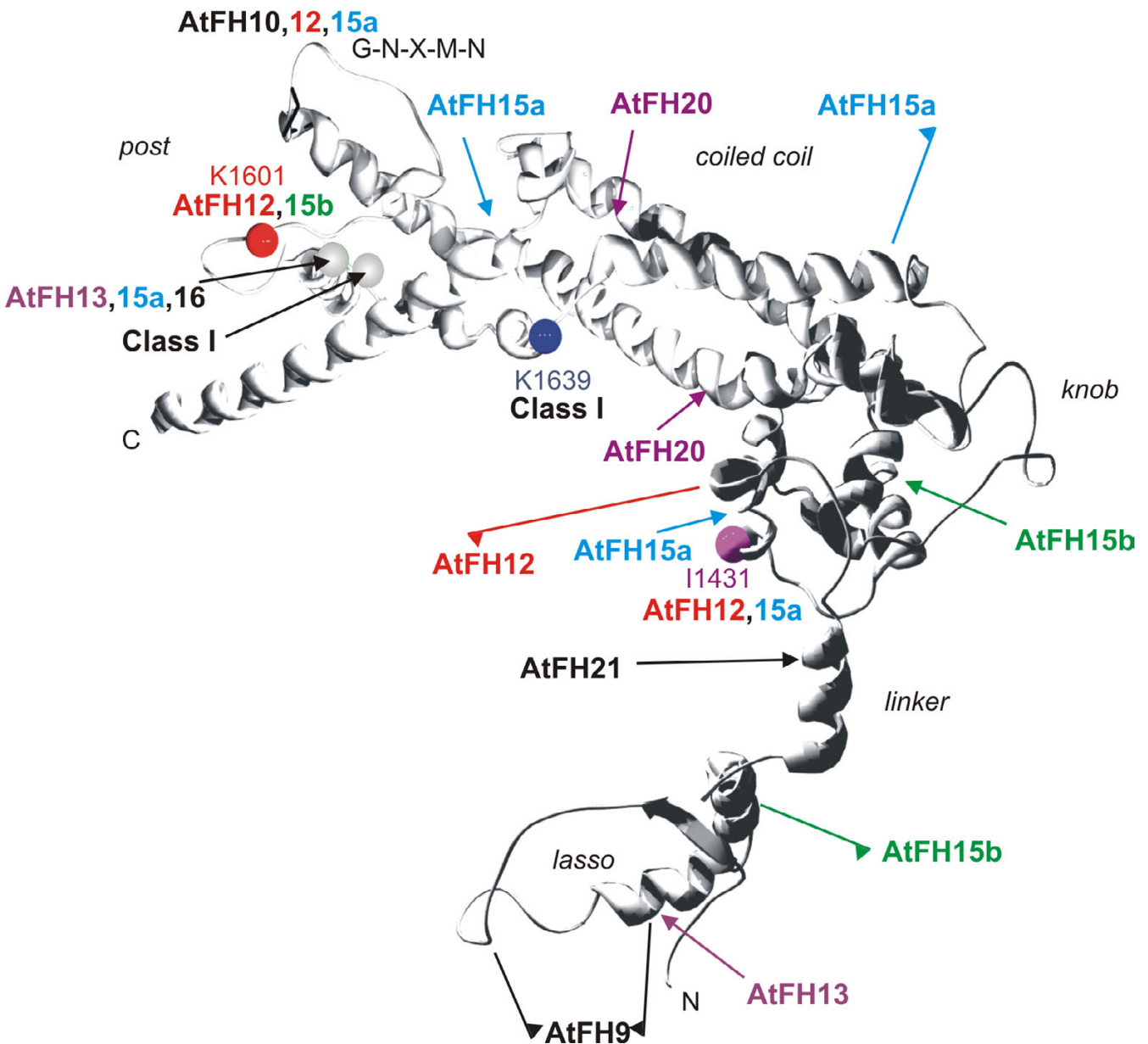
On the other hand, the overall structure of most Class II formins basically corresponds to the Bni1p/Diaphanous consensus, with several notable exceptions. AtFH20 contains two insertions in different strands of the coiled-coil part of the molecule. Insertions of 15–43 aa have been found also in the post region of AtFH13, AtFH15a and AtFH16, close to the site of the common insertion in Class I formins. AtFH13 has also an insertion in the lasso region, with possible effect on dimerization.

The case of AtFH15a and its neighbour or splicing variant AtFH15b is rather enigmatic, since both proteins miss substantial portions of the FH2 domain (part of post and coiled-coil in AtFH15a, lasso in AtFH15b) and, moreover,



**Figure 1**

**An unrooted phylogenetic tree of the plant FH2 domains.** For description of the plant genes see Table 1 and Additional file 2. Selected fungal and metazoan sequences are included: fission yeast Cdc12 (Sp CDC12, CAA92232.1), Fus1 (Sp FUS1, T43296) and For3 (Sp FOR3, CAA22841.1), budding yeast Bni1 (Sc BNII, P41832) and Bnr1 (Sc BNRI, P40450), *Dictyostelium* ForA (Dd ForA, BAC16796.1), ForB (Dd ForB, BAC16797.1) and ForC (Dd ForC, BAC16798.1), *Caenorhabditis* Cyk-1 (Ce CYK1, AAM15566.1), *Drosophila* Diaphanous (Dm Dia, P48608) and Cappucino (Dm Capp, 2123320A), mouse Formin (Mm FOR, Q05860) and Diaphanous (Mm Dia, AAC53280.1), fugu Formin (Fr FH, AAC34395.1), human FHOS (Hs FHOS, AAD39906.1). Symbols at nodes denote percentual bootstrap values (out of 500 replicates); no symbol means less than 50 % node stability, the sequence used as forced root for tree construction is marked by an arrow. For complete or nearly complete plant genes, sequences are color-coded according to their overall domain structure (see Fig. 3). Proteins encoded by the *Arabidopsis* chromosome V cluster are denoted by an asterisk.



**Figure 2**  
**Summary of structural variation in plant FH2 domains.** Structure of the yeast Bni1p FH2 domain (PDB 1UX5), with marked positions of major insertions (arrows), deletions (flags pointing towards the missing portion of sequence) and conserved site mutations (colored balls) found in *Arabidopsis* formins. Grey balls denote positions of insertions found in multiple proteins, numbers correspond to conserved amino acid positions in Bni1p.

each of them lacks one of two residues important for actin nucleation – an essential isoleucine (I1431 of Bni1) in AtFH15a and a lysine (K1601 of Bni1) in AtFH15b. We suspect that AtFH15a and AtFH15b may present two out of many possible splicing alternatives of the complex AtFH15 locus, which may be encoding multiple proteins,

including perhaps both a "complete" active formin and regulatory variants without nucleation and/or dimerization activity.

Two more proteins have mutations in the conserved positions required for actin nucleation. In AtFH21, the K1601



mutation is accompanied by a large insertion in the flexible linker. This appears to be a result of a partial gene duplication involving also the lasso-linker area of FH2, which has been subsequently lost from the posterior copy. Most dramatic deviation from the conserved structure has been found in AtFH12, which lacks both I1431 and K1601 and the whole lasso-linker assembly, while the presumed dimerization region of the post is altered. Such a protein may perhaps present a naturally occurring "hemidimer" variant, acting as a barbed end cap rather than a nucleating centre.

#### **Plant formins exhibit variable domain composition**

Following the phylogenetic analysis, we have examined the overall domain composition of plant FH2-containing proteins, searching for known sequence or structure motifs. Several patterns of conserved domain order can be distinguished in the complete plant formin sequences (Fig. 3). We further refer to these patterns as structural types A through F. Besides of the conserved domains, some formins contain long stretches of sequence (55–870 amino acids) lacking any conserved motifs, located either between FH1 and FH2 (AtFH6, AtFH21) or C-terminally (AtFH16, OsFH14).

Most plant formins contain proline-rich sequences, often called FH1 in the formin context. However, neither the FH3 domain nor additional motifs common in FH proteins, such as the DBD motif shared by diaphanous-related formins, or the coiled coil, were found in plant FH proteins (a single coiled-coil domain, located between FH1 and FH2, has been found in AtFH21).

It has been noted previously [33] that most *Arabidopsis* Class I formins contain putative secretion or membrane insertion signals and transmembrane segments, indicating that they may be integral membrane proteins. A second proline-rich domain, reminiscent of some cell wall proteins such as the extensins, is often located in the presumed extracytoplasmic portion of the protein (Fig. 3, structure A). Indeed, association with insoluble cellular fractions has been reported for one of the presumed transmembrane formins [32], providing support for the hypothesis that this type of formins may mediate anchorage of actin nucleation sites to the cell wall across the plasmalemma. We found similar sequences also in the majority of complete non-*Arabidopsis* Class I sequences (see color coding in Fig. 1), although transmembrane segments appeared to be on the edge of significance for the rice sequences OsFH10 and OsFH11. We believe that also AtFH3 may be a type A (transmembrane) formin, since its current predicted sequence appears to be 5'-truncated, and its closest relative, AtFH5, exhibits type A structure. The only non-membrane Class I formin in *Arabidopsis* is AtFH7, which resembles "standard" animal formins with

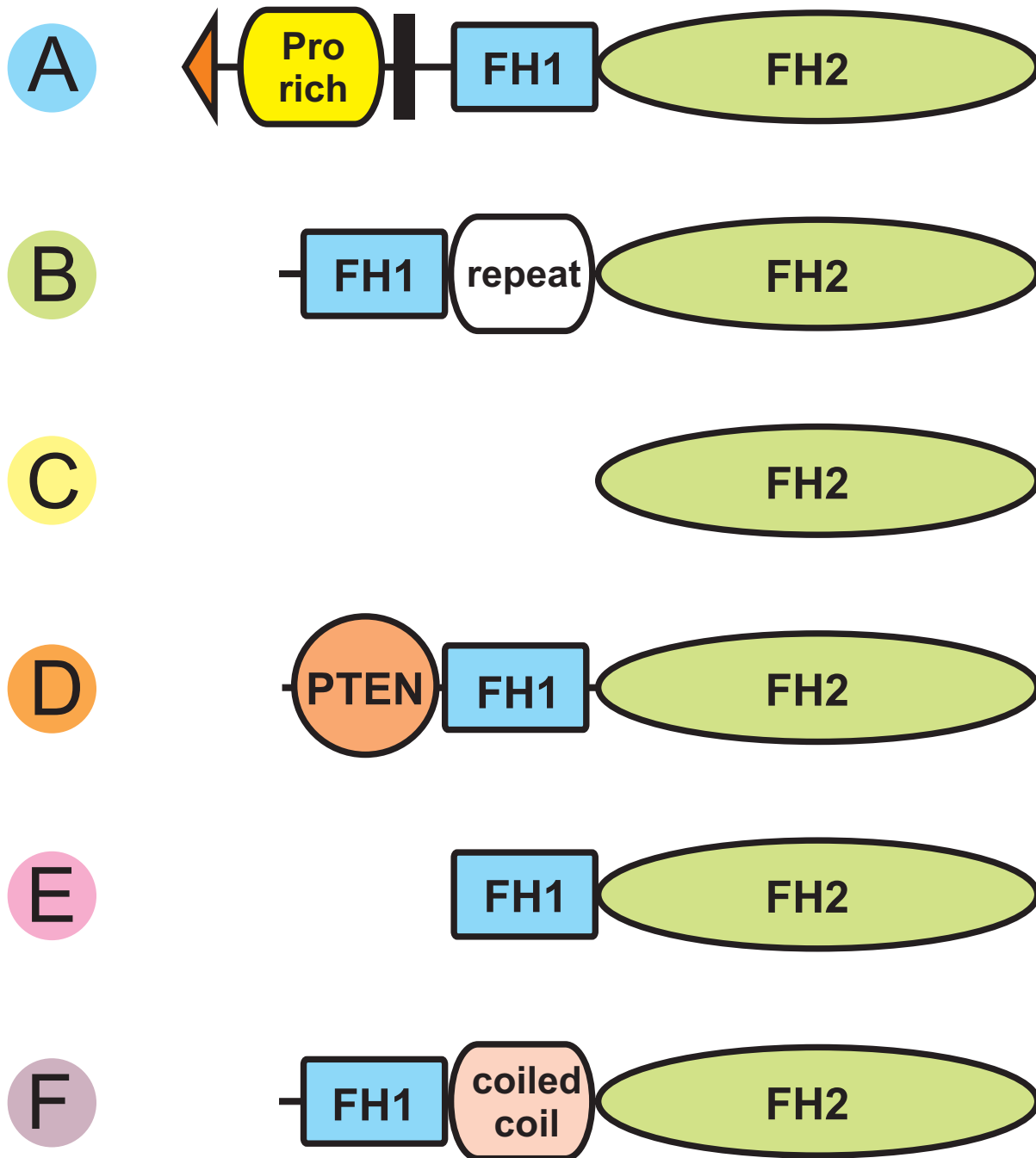
FH1 and FH2 motifs but possesses a unique repetitive structure in its N-terminal half. No other Class I proteins of this structure have been found so far, and it remains to be clarified whether this is a representative of a common, though less abundant, Class I formin type, or a relatively recent modification, present perhaps only in one or a handful of species. The same can be said about the remaining non-membrane Class I formin from rice, OsFH14, which has an extremely long C-terminal extension unparalleled elsewhere. However, until complete cDNA sequence becomes available, we cannot exclude the possibility of artifacts resulting from wrong splicing prediction in the weakly conserved parts of these loci.

While examining the predicted protein structures of Class II formins, we have noticed an area of mutual similarity in the N-terminal half of a subset of these proteins. This area exhibits also considerable similarity to the structurally well-characterized PTEN domain known from metazoans (see below; Fig. 4). Subsequently, we have found this motif, located N-terminally from the conventional FH1 and FH2 domains, in a majority of Class II formins (type D). The only exceptions are the rather diverged AtFH12 protein, AtFH17 and a group of mutually related formins encoded by genes of the cluster on the *Arabidopsis* chromosome V that apparently arose by a relatively recent series of gene duplication events. A clear distinction between Class I and Class II formins is therefore not restricted to the structural features of the FH2 domain itself, but extends also to features outside FH1 and FH2; presence of the PTEN-related domain can be considered a hallmark feature of a subset of Class II plant formins.

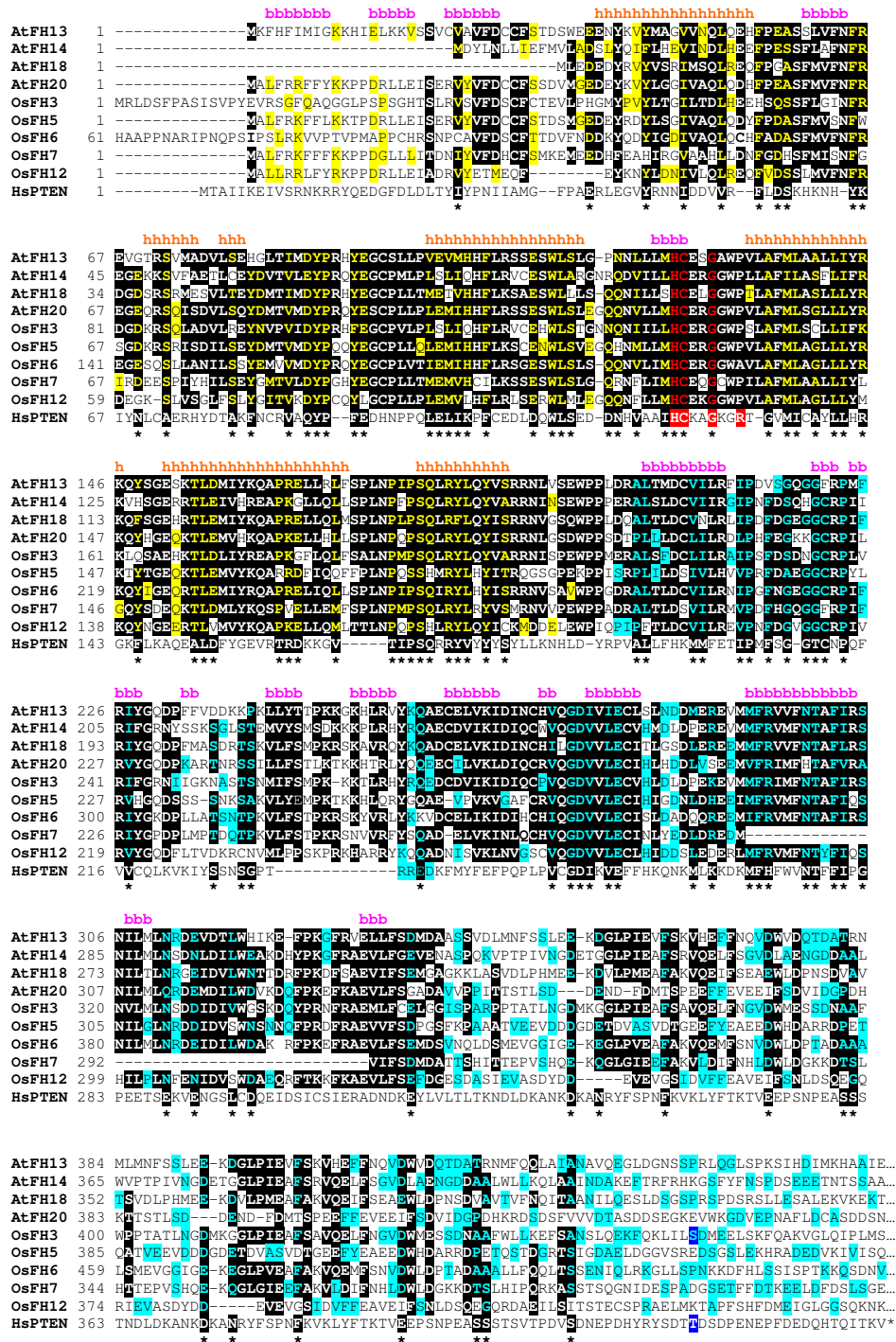
Aberrations from the characteristic domain composition of Class I and Class II formins (types A and D, respectively) appear to be mostly *Arabidopsis*-specific, and often associated with relatively recent gene duplications (a partial internal duplication involving a segment of the FH2 domain has been identified in AtFH21). Although an *Arabidopsis*-specific tendency to duplicate formin-related sequences cannot be excluded a priori, we believe that a more likely explanation is that duplicated sequences are notoriously difficult to analyse, and therefore tend to be the last ones to make their way into genome databases.

#### **A phosphatase/tensin/PTEN-related domain in most Class II formins**

While screening for known domains in plant formins using the SMART package [44], we found a significant match to the undefined specificity protein phosphatase domain (PTPc\_DSPc, SM0012, BLAST E = 9. 10<sup>-6</sup>) in the N-terminal part of AtFH18. Position-specific iterated BLAST [45] revealed a conserved domain (KOG2283 – clathrin coat dissociation kinase GAK/PTEN/auxilin) in the same region. The conserved domain falls into an area



**Figure 3**  
**Domain composition of plant FH2 proteins.** Schematic representation of the domain composition and order encountered in plant FH2 proteins (domains of variable size, such as FH1, and unique sequences not to scale). Note that only structures E and F correspond to those found outside the plant kingdom.



**Figure 4**  
**The PTEN domain of selected plant formins.** For terminology of the plant proteins see Tables 1 and 2; the remaining sequence in the alignment is human PTEN (HsPTEN, AAD13528.1). Amino acids conserved between at least one of the plant sequences and PTEN are shown in yellow for the protein phosphatase-related domain and in light blue for the C2 domain; residues conserved between at least six plant formins are inverted, and marked by asterisks if found also in PTEN. The lipid/protein phosphatase signature is in red, the putative regulatory phosphorylation site (T383) in dark blue. Note that only OsFH3 can be phosphorylated at the corresponding position. Secondary structure prediction for AtFH13 is shown above the alignment (a –  $\alpha$ -helix, b –  $\beta$ -sheet); results for other *Arabidopsis* formins were analogous.

exhibiting high degree of conservation between AtFH18 and three other Arabidopsis family members (AtFH13, 14 and 20), and a related motif has been found also in several non-Arabidopsis Class II plant formins (OsFH3, 5, 6, 7 and 12, MtFH6; see Fig. 4).

Search of the 3D-PSSM protein fold library [46] with the multiple alignment of AtFH13, 14, 18 and 20 as a probe predicted significant structural similarity to a well-defined three-dimensional structure element, the c1d5ra\_ fold. The prototype of this fold, the PTEN tumor suppressor (PDB 1d5r), is a member of a wider superfamily that includes, among others, also protein phosphatases, tensin and the tensin/auxilin domain of the cyclin G associated protein kinase (GAK) – i.e. sequences that have defined the KOG2283 domain. The presence of a PTEN-related structural motif has been independently confirmed by another sequence-structure threading method – FUGUE [47], while a third algorithm – SAM-T99 [48] – failed to recognize any conserved elements in this area. However, a secondary structure similar to that established for PTEN has been independently predicted from the sequences of AtFH13, 14 and 18 by PSIPRED [49]. Taken together, these results indicate the presence of a conserved sequence and structure motif in multiple Class II plant formins. We will further refer to this sequence/structure element as the PTEN domain.

The prototype of the PTEN domain is the human PTEN (phosphatase and tensin-related) antioncogene, whose mutation results in rapid development of multi-organ tumors in humans [50,51]. The conserved part of human PTEN protein consists of two structural units. First unit (corresponding to positions 7–185 of the human protein) has the structural similarity to dual specificity protein phosphatases, contains the active site signature of protein phosphatases, HCXXGXXR, and possesses both a lipid phosphatase activity with a strong affinity to PtdIns(3,4,5)P3 and a weak protein tyrosine phosphatase activity [52,53]. The second structural unit, related to a class of domains collectively referred to as C2, will be discussed in more detail below.

A portion of the PTEN molecule, including the phosphatase-like domain, exhibits significant similarity to the N-terminal domain of tensin, a multifunctional component of integrin-mediated focal adhesions known to participate in cell motility and cell adhesion [50,54]. A related domain was found also in some metazoan auxilins (proteins involved in uncoating of clathrin-coated vesicles), and in the cyclin G-associated protein kinase (GAK), which has an auxilin-like domain [50,55].

Some of the metazoan proteins sharing the PTEN domain are involved in the structural organization of cell regions

exhibiting a complex cytoskeletal pattern. Tensin is associated with actin in focal adhesions, acting both as a barbed-end cap and a cross-linking protein [56]. Overexpression of PTEN results in alteration of the structure of the actin cytoskeleton [57]. Moreover, although PTEN does not exhibit a specific intracellular localization, it binds to proteins localized to tight junctions in epithelia, and appears to be essential for embryonic development in mice. Together with its apparent participation in the control of cell growth, adhesion, migration, invasion and apoptosis, this suggests a possible role in the building of the cell surface and in cellular processes that involve a substantial contribution of the actin cytoskeleton [53]. In plants, homologous proteins may be therefore expected to participate in the construction of the cytoplasmic portion of the cell wall – membrane – cytoskeleton continuum.

#### ***The PTEN-related domain may have a structural rather than catalytic role***

To our surprise, PTEN domains in plant formin sequences bear mutations that make both protein and lipid phosphatase activity very unlikely (Fig. 4). The last arginine residue in the phosphatase active site, which has been shown to be crucial for catalysis [58,59], is replaced by hydrophobic or small polar residues in the plant proteins. Another residue essential for catalysis, Asp 92, that act as a general acid to facilitate protonation of the phenolic oxygen atom of the tyrosyl group [60] was in the plant formins substituted by glycine. We therefore believe that the function of plant PTEN domains is rather structural than catalytic.

Such a structural role could perhaps be attributed mainly to the second portion of the conserved PTEN motif. This second conserved unit (corresponding to positions 186–351 of human PTEN) is similar to the C2 domain that is known from a variety of proteins with multiple functions including membrane fusion, vesicular transport, GTPase regulation, protein phosphorylation, and protein degradation. The C2 domain mediates protein-membrane or protein-protein interactions, often dependent on the presence of Ca<sup>2+</sup> ions. However, not all C2 domains exhibit the same membrane-association mechanism. Some of them, such as those of synaptotagmin or phospholipase C $\beta$ , bind to membrane surface, while others, such as the C2 domain of phospholipase A2, invade the membrane by insertion of variable C2 domain loops [61]. The C2 domain of PTEN cannot bind calcium ions. Instead, two stretches of basic residues mediate peripheral binding to the membrane by electrostatic interaction [62]. The PTEN C2 domain therefore defines a specific class of calcium-independent C2 domains [61].

We were unable to detect C2 domains in plant formins by any of the software used to find PTEN similarity (see

above and Methods). This can be due to a huge sequence divergence within the C2 domains family, reflecting the diversity of functions they participate in. However, upon visual inspection of sequence alignment, we found obvious sequence similarity to the C2 domain of human PTEN protein, including the regions that make PTEN  $\text{Ca}^{2+}$  independent (Fig. 4).

Using the human PTEN C2 domain as a template, we were able to produce 3D models of three representative plant formin C2 domains, AtFH13, AtFH14 and AtFH15, further strengthening the notion that plant Class II formins do possess a C2 domain. Surface charge distributions of all three formin C2 domains resemble those of phospholipase A2 (PLA2) rather than PTEN (Fig. 5). Membrane association of the PLA2 C2 domain is  $\text{Ca}^{2+}$ -dependent and occurs via hydrophobic interactions with zwitterionic phospholipids and insertion of C2 loops into the membrane. However, since the formin C2 domains appear to be unable to bind  $\text{Ca}^{2+}$  (at least not in the same manner as PLA2), although they perhaps could be neutralised by other means, it remains to be decided experimentally whether plant Class II formins interact with membranes via their C2 domains.

Phosphorylation of a threonine residue downstream from the C2 domain (at position 383 of human PTEN) was recently found to modulate the ability of PTEN to modify cell migration in culture [63]. However, the phosphorylated motif, including the crucial threonine, is not conserved in plant sequences, suggesting that this regulation may be specific for the metazoan lineage (Fig. 4).

It is therefore tempting to speculate that the widespread occurrence of the PTEN domain among plant Class II formins may suggest a function analogous to that proposed for the transmembrane segments in their Class I counterparts. A variant PTEN domain that has lost its catalytic activity by mutation but retained its intracellular localisation may act as an anchor positioning the actin-nucleating sites (FH2) to some intracellular (and possibly cell surface/plasmalemma-associated) structures.

## Conclusions

One of the most intriguing questions of contemporary molecular biology is how can vastly dissimilar organisms develop utilizing a limited repertoire of basically similar molecules derived from a relatively small set of conserved protein domains? Processes of eukaryotic cell morphogenesis, such as shaping of the actin cytoskeleton, provide a good example of such a versatile usage of conserved molecular mechanisms. Well-conserved protein domains, such as the FH1/FH2 motifs shared by formins and participating in actin nucleation, are being used for a broad range of cellular tasks in various eukaryotic lineages. It is

plausible to assume that the diversity of cellular functions may to a large extent stem from the context those domains assume within the framework of larger, multidomain protein molecules.

We have examined the phylogeny and molecular context of the FH2 domain in available angiosperm formin homologues. Besides of confirming the existence of two classes of plant formins, suggested previously solely on the basis of Arabidopsis data, we could identify a novel domain shared by a significant portion of Class II formins and possibly involved in the association between formin-based actin nucleation modules and other cellular structures.

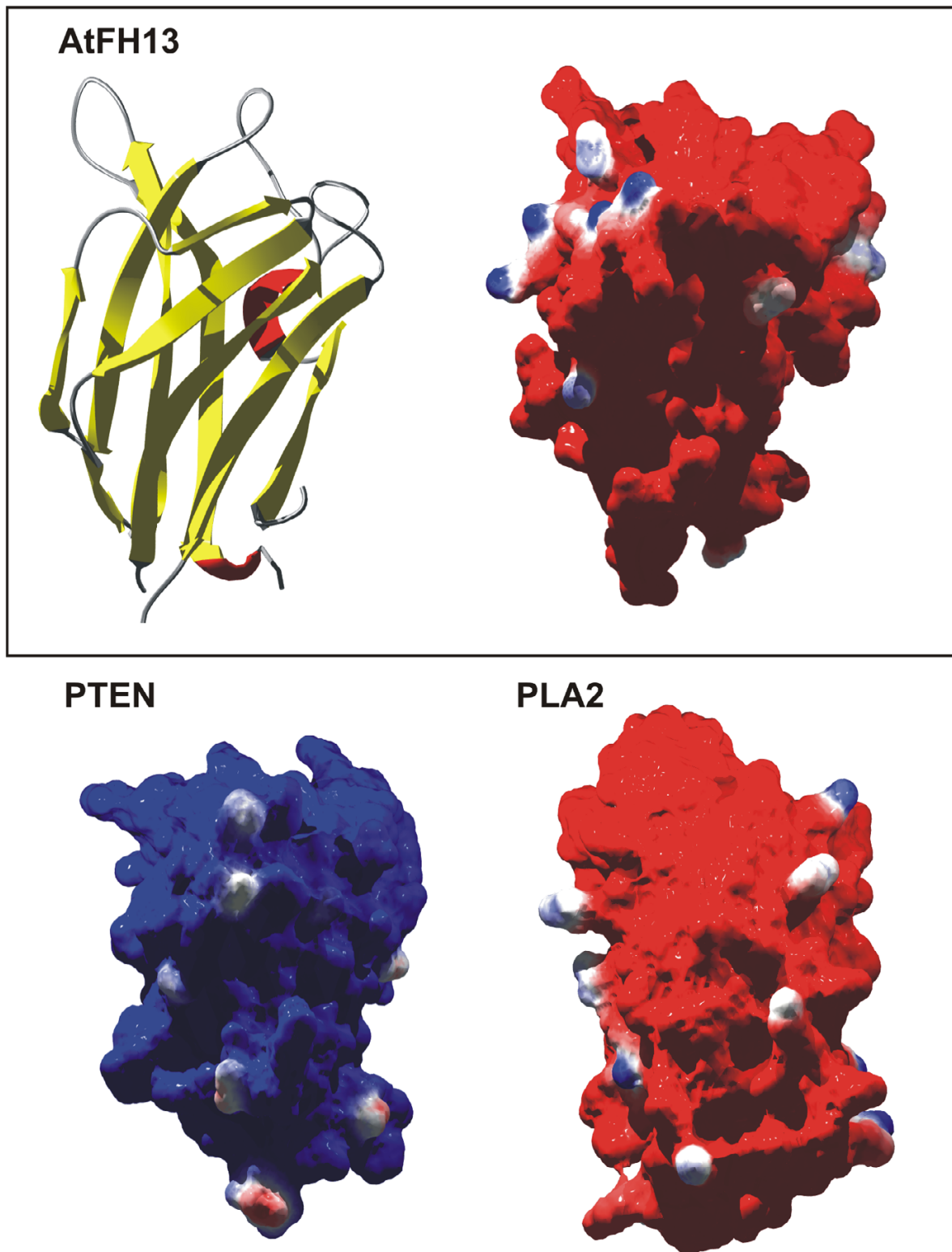
Another interesting aspect is the extremely dynamic evolution of the angiosperm formin family, with ample evidence for multiple gene duplication events (sometimes accompanied by major domain rearrangements) not seen in other species studied so far. At the moment, we can only speculate whether this is a feature specific for a small selection of plant taxa, or a general characteristic of plant formins. However, we hope that publication of more complete plant genomes can help to resolve this question in a not so distant future.

The enormous variety of formins within a single plant organism far exceeds that found in fungi and metazoans, where only a handful of formin genes exist, although their diversity can be enhanced by alternative splicing [13] and heterodimerization [14]. However, even the relatively small *Arabidopsis* genome contains no less than 21 FH2 domain-encoding loci, while possibilities for alternative splicing and dimerization remain open, resulting in literally hundreds of possible species of the functional formin dimer. It is tempting to speculate that this diversity may be related to the demands of actin-nucleation site positioning with respect to precisely defined cellular surfaces in the context of a multicellular body consisting of cells endowed with rather rigid surface structures. Consequently, a major increase in formin diversity could be expected at the transition between unicellular green algae and multicellular plants. However, testing of this hypothesis would have to wait until more complete data from algal and gymnosperm genome projects become available.

## Methods

### Identification of FH2-containing plant genes

*Arabidopsis* loci encoding putative formin homologues have been identified by BLASTP and TBLASTN searches [64] of both predicted protein and complete genome sequence databases (at TAIR and NCBI) using previously characterized members of the family [33] as the query. The same loci have been found by both approaches.

**Figure 5**

**Model of the C2 domain of AtFH13.** 3D model of the AtFH13 C2 domain and its predicted surface potential compared to that of human PTEN (PDB 1d5r) and calcium-free human PLA2 (PDB 1bci) C2 domains (red – negative, blue – positive). All models are oriented membrane side upwards. Analogous results have been obtained also for AtFH14 and AtFH18.

Analogous searches have been performed with the most diverged sequences from the first round as the query, until no new significant matches appeared. Presence of FH2 domains in predicted candidate open reading frames has been confirmed by a SMART search [44] for all genes. Sequences from other plant species coding for related proteins have been identified in analogous searches of Entrez *nr*, *est* and *htgs* databases.

#### **Detection of gene expression**

For *Arabidopsis* genes with no available cDNA sequence, microarray slides with highest transcript levels within the NASCArrays data set [35] have been detected using the Spot History tool. Visual inspection of the Detection parameter in full slide data has been used to confirm that detected expression levels were significantly above zero.

#### **Gene structure predictions**

For every *Arabidopsis* formin-related gene, exhaustive TBLASTN searches of the Entrez *nr* and *est* databases have been performed in order to identify all cDNAs derived from the chromosomal locus. Resulting cDNA sequences were aligned to the genomic and predicted ORF sequences with the aid of the MACAW program [65]. Conflicts between the genomic and cDNA sequence have been resolved in favor of the genomic version unless strong evidence suggested genomic sequence errors (see Results and Discussion). In cases of either a complex splicing pattern without experimental support or apparent deletions in the FH2 portion of the sequence, an alternative splicing prediction has been obtained using GenScan [66,67]. GenScan usually agreed reasonably well with the original genome annotation, although it did miss exons occasionally. WebGene [68] or MZEF [69] have been used as well in some cases, however these programs appeared to be inferior with respect to both performance (agreement with cDNA or the FH2 consensus) and output readability. The original splicing predictions have been modified for some genes, taking into account alternative predictions and the structure of closest homologues (see Results, Table 1 and Additional file 1). Programs of the Sequence manipulation suite [70] version 2 [71] have been used for general sequence manipulation, assembly and translation tasks.

For non-*Arabidopsis* genomic sequences, gene models have been produced in an analogous manner, based on combination of existing genome annotation (if available), GenScan and WebGene predictions and alignment to the closest *Arabidopsis* relatives.

#### **Protein sequence alignments and phylogeny reconstruction**

To produce an alignment of FH2 domain sequences, selected representative members of divergent branches of

the family have been aligned using ClustalW [72] configured to run as a helper application under the BioEdit package [73], using the BLOSUM matrix series. The resulting core alignment has been modified manually using BioEdit, taking into account independently produced alignments of selected subsets of sequences, made with the aid of the MACAW software [65] as described previously [33]. Sequences closely related to those already present in the core alignment have been merged to the alignment manually in the BioEdit environment. Alignments of the PTEN domain have been produced in an analogous manner, taking into account 3D structure data (see below).

All positions containing gaps in at least one sequence have been removed prior to the construction of the phylogenetic tree. An unrooted phylogenetic tree based on the resulting alignment has been produced with the aid of the Treecon package [74] using the neighbor-joining (NJ) algorithm [43] with Poisson correction for distance estimation.

#### **Protein domain recognition**

The SMART program package [44,75] version 4 has been used for identification of known domains, secretory signals (by the SignalP method – [76]), transmembrane segments (by the TMHMM algorithm – [77]) and repetitive sequence motifs within predicted formin sequences. Unless stated otherwise, only signals above the default significance threshold have been taken into account.

Additional series of domain searches has been performed using reverse position-specific BLAST [45] against the CDD database (version 1.65), which contains a larger selection of domains than the SMART collection (including FH3 and DBD), with the same results.

#### **3D structure searches and alignments**

The 3D-PSSM threading algorithm [46] has been used to find possible known protein folds in the shared N-terminal domain of AtFH13, 14, 18 and 20, using a multiple alignment of the four sequences as a probe. The result appeared to be highly significant (over 95 % confidence, PSSM E = 0.00179 for the 1d5ra\_ fold). For independent confirmation, a FUGUE version v2.s.07 [47,78] search against the HOMSTRAD fold library has been performed for two selected sequences (AtFH13 and AtFH14), resulting in identification of the same fold in both cases as "certain". A third algorithm, SAM-T99 [48,79] produced no significant results. However, this program is known to have a very powerful filter against false negatives that loses some positives, especially those with predominantly helical structures (see manuals at the program website).

**Secondary structure predictions and homology modelling**

Secondary structure of type II formins was predicted using a PSIPRED server [49,80].

The structure of the human Phosphoinositide phosphatase PTEN (1d5r) was used as a template for modeling the C2 domains of AtFH13, AtFH14 and AtFH18. The template and the target structures were aligned with ClustalW [72] and the resulting alignment was manually edited with help of the secondary structure prediction outputs.

The WHAT IF program [81] was used for modeling as described [82]. The sequence alignments and coordinates of the models are available as Additional files 6 to 9.

SwissProt Deep View [83,84] has been used to calculate electrostatic potentials of the 3-D models of formins and to generate their figures, which have been graphically visualized using the PovRay 3.5 raytracing software [85].

**Authors' contributions**

FC conceived of the study, participated in database searches and sequence analyses, carried out the phylogenetic analysis and drafted the manuscript. MN carried out the 3D structure analyses and model building. DP participated in database searching and re-annotation of *Arabidopsis* genes. VŽ participated in the microarray data analysis and substantially contributed to the design of the study and writing of the manuscript. All authors read and approved the final manuscript.

**Additional material****Additional File 1**

*Nucleotide sequence alignments used to generate revised ORF predictions for selected Arabidopsis formins*

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**Additional File 2**

*Non-arabidopsis plant FH2 proteins included in the analysis GenBank/EMBL/DDBJ accession numbers are shown where available; AF3, AF4 – see Additional files 3 or 4; NA – not available.*

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**Additional File 3**

*Predicted protein sequences of rice formins*

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**Additional File 4**

*Translations of non-arabidopsis, non-rice ESTs or EST assemblies encoding (partial) FH2 proteins.*

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**Additional File 5**

*Alignment of the FH2 domain sequences used for dendrogram construction*

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**Additional File 6**

*Alignment used for construction of plant formins C2 models.*

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**Additional File 7**

*Coordinates of the model of the AtFH13 C2 domain.*

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**Additional File 8**

*Coordinates of the model of the AtFH14 C2 domain.*

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**Additional File 9**

*Coordinates of the model of the AtFH18 C2 domain.*

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