Phylogenetic and comparative gene expression analysis of barley (*Hordeum vulgare*)

WRKY transcription factor family reveals putatively retained functions between monocots and dicots

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Abstract

**Background:** WRKY proteins belong to the WRKY-GCM1 superfamily of zinc finger transcription factors that have been subject to a large plant-specific diversification. For the cereal crop barley (*Hordeum vulgare*), three different WRKY proteins have been characterized so far, as regulators in sucrose signaling, in pathogen defense, and in response to cold and drought, respectively. However, their phylogenetic relationship remained unresolved.

**Results:** In this study, we used the available sequence information to identify a minimum number of 45 barley WRKY transcription factor (HvWRKY) genes. According to their structural features the HvWRKY factors were classified into the previously defined polyphyletic WRKY subgroups 1 to 3. Furthermore, we could assign putative orthologs of the HvWRKY proteins in Arabidopsis and rice. While in most cases clades of orthologous proteins were formed within each group or subgroup, other clades were composed of paralogous proteins for the grasses and Arabidopsis only, which is indicative of specific gene radiation events. To gain insight into their putative functions, we examined expression profiles of *WRKY* genes from publicly available microarray data resources and found group specific expression patterns. While putative orthologs of the HvWRKY transcription factors have been inferred from phylogenetic sequence analysis, we performed a comparative expression analysis of WRKY genes in Arabidopsis and barley. Indeed, highly correlative expression profiles were found between some of the putative orthologs.

**Conclusions:** *HvWRKY* genes have not only undergone radiation in monocot or dicot species, but exhibit evolutionary traits specific to grasses. HvWRKY proteins exhibited
not only sequence similarities between orthologs with Arabidopsis, but also relatedness in their expression patterns. This correlative expression is indicative for a putative conserved function of related WRKY proteins in mono- and dicot species.
Introduction

In plants, WRKY proteins constitute a large family of transcription factors. Since their first identification in sweet potato and wild oats [1, 2], the WRKY proteins have been described as transcriptional regulators in multiple plant species and in various processes such as pathogen defense [3-9], trichome [10] and seed development [11], as well as leaf senescence [12, 13]. The common feature of WRKY proteins is the existence of either one or two WRKY domains. The WRKY domain is an approximately 60 amino acids long spanning DNA binding domain, which contains a highly conserved amino acid motif, WRKYGQK, at its N-terminus and a metal chelating zinc finger signature at the C-terminus. The crystal structure of one WRKY domain has been resolved recently [14] and provides an insight into how the conserved residues of the WRKY domain facilitate the binding to their cognate DNA-element, the W-box. While most of the functional studies of WRKY proteins indicate a binding to the conserved nucleotide consensus sequence TGAC(C/T) of the W-box, the binding to another element called SURE has also been described [15].

Based on the number of WRKY domains present and on the type of zinc finger WRKY proteins have been classified into groups 1, 2a to 2e and 3 [16]. WRKY transcription factors seem to have their evolutionary origin in ancient eukaryotes with the most basal WRKY genes identified in the unicellular eukaryote Giardia lamblia and the slime mold Dictyostelium discoideum [17, 18]. While the WRKY-GCM1 superfamily has been found in all organism kingdoms, the conserved WRKY-type signatures of these genes have not yet been detected in the lineage of true fungi and animals [19]. In contrast to that, phylogenetic analysis revealed that the WRKY transcription factor family has
expanded to great extent in higher plants [20]. For example, the WRKY gene family consists of 72 members in Arabidopsis [16] and at least 81 members in rice [6, 18, 21, 22].

The cereal crop barley is of high importance for food, feed and brewing. After maize, rice and wheat, barley ranked as number four in cereal production in 2005[23]. Due to its wide ecological potential and adaptability it is cultivated world wide and under diverse environmental conditions. For example, barley is grown on soils and in altitudes that are unsuitable for wheat and oats [24]. The genome of barley is estimated to be of approximately 5000Mb in size and has not been fully sequenced yet. However, the GenBank EST dataset (http://www.ncbi.nlm.nih.gov/) for barley contains almost 440,000 ESTs and provides a valuable resource for bioinformatics analyses. In addition, the Affymetrix 22K Barley1 GeneChip microarray, representing probe sets for roughly 22,000 genes [25], was released on the market in 2004. Nowadays, a number of barley microarray experiments are accessible via different public sources such as BarleyBase [26] or GEO [27] and allow for in silico expression analyses of barley genes.

Several studies have shown an interest in barley WRKY proteins (HvWRKY) and their function. HvWRKY1, also named HvWRKY38, has been reported to be involved in several regulatory processes, such as the response to cold and drought [28], and the repression of the alpha-amylase gene *Amy32b* during seed germination [29]. Recently, the same WRKY protein and its paralog HvWRKY2 have been described as repressors of the PAMP-triggered basal defense upon infection with *Blumeria graminis* [8]. Apart from HvWRKY proteins functioning in response to biotic and abiotic stresses, the group 1
WRKY protein SUSIBA2 has been shown to mediate sucrose signaling in barley cells during regulation of starch synthesis [15, 30].

In this study we identified 45 members of the WRKY gene family in barley based on publicly available sequence information. We describe their phylogenetic relationship and try to assign putative orthologs from barley, Arabidopsis and rice. Expression analysis using microarray data suggests the involvement of some of the HvWRKY proteins in response to powdery mildew and in plant development. Moreover, correlated expression of orthologous barley and Arabidopsis WRKY genes imply that their function is retained in monocot and dicot species.

Results

Identification of 45 WRKY protein coding sequences in barley

To estimate the minimum number of WRKY genes in barley, publicly available sequence data was searched with the BLAST program [31] at the NCBI website (http://www.ncbi.nlm.nih.gov/). The protein sequence of WRKY domains from representative members for all subgroups from Arabidopsis (AtWRKY1, -18, -6, -8, -7, -14 and -30) were used as input. The primary search resulted in 142 non-redundant hits, of which 20 were removed as they did not contain the conserved amino acid sequence of the WRKY domain. The remaining 122 sequences were screened for partial overlaps by pairwise comparison. This analysis identified 45 WRKY open reading frames (ORFs) unique for barley. A keyword search on NCBI returned eight previously annotated HvWRKY protein sequences. As indicated by corresponding EST sequences, all eight ORFs were expressed. In order to create a uniform nomenclature with consecutive
numbering, we renamed sequences annotated as putative WRKY2, 3, 4, 5 and 6 to putative HvWRKY2, 3, 4, 5 and 6, respectively. Sequences of HvWRKY3 and 4 could be extended based on available EST information. The sequence of putative WRKY1 protein, CAD60651, is identical with the sequence AAS48544 which has been published as HvWRKY-38 [28]. We designated both sequences as HvWRKY1 and did not annotate any sequence as HvWRKY38 to prevent confusion. SUSIBA2 received the formal name HvWRKY46. The remaining sequences were annotated accordingly as HvWRKY7 to HvWRKY37 and HvWRKY39 to HvWRKY45 (Table 1). Seven of the HvWRKY sequences included start and stop codons and, as judged by protein sequence alignments with WRKY proteins from rice (data not shown), constitute complete coding sequences. Further 39 HvWRKY sequences represent partial coding sequences.

In addition to the sequence searches in public databases, we used PCR techniques to extend sequence information for HvWRKY6 and HvWRKY9. This approach allowed for the identification of a second WRKY domain in the sequence of HvWRKY6. In case of HvWRKY9, a characteristic domain with the highly conserved amino acid sequence HARF [32] could be detected at the beginning of the coding sequence.

While Eulgem et al. [16] defined WRKY proteins by the conserved heptapeptide WRKYGQK, slight variations of this sequence have been described for a number of rice WRKY proteins [20, 21]. Similarly, a number of HvWRKY sequences have amino acid sequence substitutions in the conserved WRKY signature. In HvWRKY18, 19 and 20 the WRKY domain displays the sequence WRKYGKK, in HvWRKY33 and 34 WRKYGEK, in HvWRKY36 WRKYGQN and in the C-terminal domain of HvWRKY24 WSKYGQM (Figure 1). Moreover, a unique zinc-finger motif (C-X_4,5-C-
X_{22-23}-H-X-H) was described as a second characteristic feature of WRKY proteins [16]. In group 3 WRKY proteins, this motif is slightly altered to C-X_{7}-C-X_{23}-H-X-C. In barley, three of the so far characterized group 3 WRKY proteins, namely HvWRKY33 and 34 showed an extension of the zinc finger motif (C-X_{7}-C-X_{24}-H-X-C) compared to the patterns described by Eulgem et al. [16] (Figure 1). Again, similar modifications of the zinc-finger patterns were detected for rice WRKY proteins and have been discussed to be a monocot-specific feature [18, 20, 21].

**Phylogenetic analysis of barley WRKY proteins**

While the phylogenetic analysis showed that WRKY proteins are of monophyletic nature [19], the assignment to defined groups is polyphyletic and either based on the number of WRKY domains or on the type of zinc finger motif. To study the phylogenetic relationship of the 45 HvWRKY proteins we performed a multiple sequence alignment of the about 60 amino acid spanning the WRKY domain of all HvWRKYs. To obtain a better separation of the different groups and subgroups, WRKY domains from members of different Arabidopsis and rice WRKY proteins were included for the analysis. For each of the groups 1, 2a to 2e and 3 one representative was chosen randomly. These were AtWRKY20, -40, -72, -50, -74, -65, -54 for Arabidopsis and OsWRKY82, -71, -27, -25, -68, -2, -46 for rice (according to Xie et al. [21]). Both C-terminal and N-terminal WRKY domains of group 1 WRKY proteins have been treated as independent sequences, so that a total number of 51 barley, 8 Arabidopsis and 8 rice WRKY domains were analyzed.

As shown in Figure 2, for all subgroups of WRKY proteins we could identify at least one representative in the barley genome. For example, five HvWRKY sequences code for
proteins with two WRKY domains and clearly group with the group 1 members AtWRKY2 and OsWRKY82. The single WRKY domain identified for HvWRKY43 clusters within the C-terminal group 1 domains, indicating that a second N-terminal domain remains likely to be identified. Interestingly, the 24 identified group 2 WRKY members of barley are distributed very unevenly among the five subgroups. Whereas 11 HvWRKYs form a distinct subclade with the characteristic members of subgroup 2c, only a single barley WRKY, HvWRKY37, belongs to group 2b. Group 3 is represented by 13 single-WRKY domain barley proteins. Surprisingly, both WRKY domains of HvWRKY24 cluster with these proteins. Again, this phenomenon has been observed for at least four rice WRKY proteins [21], namely OsWRKY41, -61, -63 and -81.

The total number of barley WRKY genes will remain unknown until the sequencing of the barley genome is completed. However, we constructed an alignment and a phylogenetic tree of 72 Arabidopsis, 45 barley and 81 rice WRKY proteins to be able to estimate this number (Additional Files 1 and 2). As described earlier by Zhang and Wang [20], group 2 and 3 domains were closer related to the C-terminal domains of group 1 whereas the N-terminal WRKY domains of group 1 members form a separate monophyletic cluster. On the other hand, separate subclades of the WRKY domains of subgroup 2c underline the paraphyletic nature of these subgroups (Additional File 2).

The grouping of HvWRKY36, HvWRKY6 and OsWRKY82 based solely on the phylogenetic analysis of their WRKY domain sequences can not be performed unambiguously. However, the presence of two WRKY domains in HvWRKY6 and OsWRKY82 is indicative for group 1 members. Both C-terminal and N-terminal WRKY domains of HvWRKY6 and OsWRKY82 support basal monophyletic subclades of group
1 WRKY proteins (Figure 1 and Additional File 2), which might be specific for grasses [21].

The number of WRKY proteins within each plant assigned to the different paraphyletic subgroups 1 to 3 was calculated and is summarized in Table 2. A simple comparison of the total number of Arabidopsis and rice WRKY genes with the number of HvWRKYS indicates that approximately 50% of the barley WRKY genes have been identified. Also, the distribution of WRKY proteins showed that HvWRKYS are underestimated in several subgroups. Nevertheless, similar numbers of barley and rice WRKY proteins in subgroups 2a and 2d may indicate that all HvWRKY proteins of this subgroup have been identified.

**Monocot-specific radiation and identification of orthologous WRKY genes**

Based on the WRKY domains of all proteins from barley, Arabidopsis and rice, we calculated a large phylogenetic tree to gain a closer insight into the evolutionary relationships between WRKY proteins (Additional File 2). From that, we wanted to assign putative WRKY orthologs between these sequences. Additionally, nine WRKY domains of the moss *Physcomitrella patens*, two each from *Giardia lamblia* and *Dictyostelium discoideum* as well as additional zinc finger domains of other WRKY-GCM1-like proteins were included as basal outgroups.

In almost all WRKY subgroups clades exist that solely consist of rice WRKY domains or a combination of rice and Arabidopsis WRKY domains. These groups might comprise orthologs of yet unidentified HvWRKY proteins. In contrast, rice and barley WRKY domains form a number of monocot-specific clades in group 2c and group 3.
These clades reveal clusters of orthologous WRKY proteins for rice and barley. For example, in subgroup 2c, AtWRKY50 forms a basal accession to a cluster of five barley and six rice WRKY domains. Similarly, in a subgroup of group 3 ten barley and nine rice WRKY proteins constitute putative monocotyledonous orthologs to a group of three paralogous Arabidopsis WRKY proteins (Additional File 2). The barley and rice WRKY proteins themselves form paralogous or orthologous groups and hence, would indicate monocot-specific diversification of WRKY genes. This assumption was further supported by BLAST searches in dbEST, which did not return orthologous WRKY domains of any other dicotyledonous species for the monocot-specific subclades (data not shown). Thus, our results support the idea of radiation events in groups 2c and 3 WRKY proteins that are specific for monocot species. This diversification has most likely occurred after the divergence of the ancestors of dicotyledonous and monocotyledonous plants.

The assignment of orthologs is crucial in order to transfer the knowledge on WRKY protein functions from the model plants rice and Arabidopsis to barley. Based on the presented phylogenetic tree (Additional File 2), we assigned orthologs for most of the WRKY genes described (Table 1). In a few cases, orthologs could not be defined unambiguously. However, as gene orthology implies similarity in gene function, the assignment presented provides a solid base for functional analyses.

**Powdery mildew infection affects expression of *HvWRKY* genes**

The analysis of temporal and spatial expression patterns is an initial step towards the functional characterization of genes. The expression of numerous WRKY genes from various plant species is affected by biotic and abiotic stresses ([6] and references therein).
and implies the regulatory role of WRKY proteins in stress responses. In order to investigate if this is also true for \textit{HvWRKY} genes, we applied publicly available microarray data of powdery mildew- (\textit{Blumeria graminis}) infected barley leaves. Totally 23 probesets on the Barley1 GeneChip could be assigned to 20 different \textit{HvWRKY} genes (Table 3). Datasets of two experiments named BB4 and BB7 were downloaded from BarleyBase [26, 33]. In Figure 3, the expression profiles of 20 \textit{HvWRKY} genes are displayed for two genotypes (Mlo and mlo5) under infection versus non-infection conditions (BB7) as well as during a time course experiment using the host genotype Mla1 and the Bgh isolate 5874 (BB4, see publication for details). Even though the setup of the BB7 and BB4 experiments differs, some general trends can be seen in both experiments. Roughly 50\% of all \textit{HvWRKY} genes show altered expression levels upon infection with \textit{Blumeria graminis}. \textit{HvWRKY}1 and \textit{HvWRKY}2 have previously been shown to be involved in the regulation of defense mechanisms [8]. \textit{HvWRKY}1 expression appears to be drastically increased under the one experimental condition and lower in the second, while \textit{HvWRKY}2 shows increased expression levels in both experiments. In addition to these regulators, a number of other putatively pathogen responses related to \textit{HvWRKY} genes can be assigned based on their expression profiles. For example, the gene for \textit{HvWRKY}3, which is closely related to the well-studied pathogen response regulators \textit{AtWRKY}18, -40 and -60, shows an increase in expression after 20h of pathogen treatment. Also, candidates for pathogen response regulators can be found in subgroups 2c and 3. Interestingly, these are WRKY groups that have undergone monocot-specific radiation events, which might thus have been related to an increased biotic stress during evolution. In addition to upregulated genes, there are \textit{HvWRKY} genes with a decrease in
transcript levels, such as most of the *HvWRKY* genes in group 1. However, several *HvWRKY* genes are not affected in their expression upon powdery mildew infection and, therefore, might function in regulatory processes other than pathogen response such as abiotic stress or development.

**Orthologous barley and Arabidopsis *WRKY* genes exhibit correlative expression signatures during plant development**

A number of WRKY proteins have been reported to be involved in plant developmental processes [10-12]. To investigate if *HvWRKY* transcript levels vary during plant development, we used publicly microarray data of different developmental stages from barley [34]. In addition, we investigated whether phylogenetically related *WRKY* genes of barley and Arabidopsis exhibited similarities in their expression patterns. As not all *WRKY* genes of both barley and Arabidopsis are represented on the respective microarrays, we compared the expression patterns of 16 *HvWRKY* genes and their putative Arabidopsis orthologs. We extracted expression data for ten organs in barley at different developmental stages, coleoptile, mesocotyl, radicule, leaf, root, inflorescence, anthers, and caryopsis 5, 10 and 16 days post anthesis (d.p.a.), as was described by Druka et al. [34].

The expression profiles reveal spatial and temporal variations in expression of *HvWRKY* genes in different barley organs (Additional File 3). Some genes, like *HvWRKY2*, -4 and -46, have similar expression levels in all investigated tissues. In contrast, genes such as *HvWRKY5*, -6, -9, -39 and -41 reveal more than 3-fold difference in expression levels in different organs. Furthermore, the transcript levels of barley *WRKY* genes in the same
organ show temporal variations. For example, \textit{HvWRKY32} is expressed in young caryopses at 5 d.p.a. but is clearly reduced in later stages of caryopsis development (Figure 4). Similarly, the expression levels of \textit{HvWRKY6} are low in radicules but significantly higher in older roots. In addition, paralogous barley \textit{WRKY} genes such as \textit{HvWRKY5} and -20 reveal similar expression patterns, indicating redundant function.

Next, we performed a comparative expression analysis for both barley and Arabidopsis \textit{WRKY} genes on the extracted subsets of the microarray expression datasets of homologous organs by Druka et al. [34] and Schmid et al. [35].

The Pearson correlation of the normalized signal intensities was calculated for the potentially orthologous \textit{WRKY} genes. The resulting correlation coefficients ranged from -0.49 to +0.91 (Table 4). The average correlation of all the putative orthologous \textit{WRKY} genes was +0.24 and differed significantly ($p$-value 0.009) from the average expression correlation of a control dataset composed of 101 randomly chosen gene pairs (+0.01; Additional File 3). Selection of only the best correlated pairs of \textit{HvWRKY} and \textit{AtWRKY} orthologs even increased the average correlation coefficient to +0.37. The observed high correlation is mirrored by the overall expression pattern of the compared \textit{WRKY} genes (Figure 4). For example, the expression levels of the orthologous pair in group 1, \textit{AtWRKY20} and \textit{HvWRKY46}, hardly differ in any of the ten organs and are clearly correlated. In contrast, the expression profiles of \textit{HvWRKY9} and \textit{AtWRKY7} are observably anticorrelated. Genes for the group 2e WRKY protein \textit{HvWRKY39} and its closely related Arabidopsis orthologs \textit{AtWRKY65} and -69 show a consistent increase in mRNA level in radicules and roots with low expression levels in leaves. According to our phylogenetic analysis, \textit{HvWRKY6} has a single ortholog in rice, \textit{OsWRKY82}, and two
potential orthologs in Arabidopsis, namely AtWRKY1 and AtWRKY32. The \textit{HvWRKY6} expression profile resembles that of \textit{AtWRKY1} (+0.41) in vegetative tissues but is more similar to the profile of \textit{AtWRKY32} (+0.45) in reproductive organs such as the inflorescence and the anthers. This might indicate that the Arabidopsis \textit{WRKY} genes have undergone functional changes after a duplication event.

Next, we performed RT-PCR experiments on cDNA from three different organs (roots, leaves and infructescences). To extend our studies and support our findings, we also included samples from rice, \textit{Oryza sativa japonica}, for the analysis. The signal intensities of the bands were quantified and normalized to the \textit{Actin} controls to calculate the normalized fold intensities for each organ and organism, respectively (Additional File 4). First of all, the expression trajectories of all tested barley and Arabidopsis genes reproduced in general the data of the microarrays experiments (Figure 5). Transcripts of the three orthologs \textit{HvWRKY46}, \textit{OsWRKY78} and \textit{AtWRKY20} were the most abundant and present to the same amount in all organs. Although \textit{HvWRKY39} and \textit{OsWRKY13} constitute close relatives, their normalized intensities differ especially in leaves, where \textit{HvWRKY39} transcripts can not be detected (Figure 5).

Intriguingly, the expression of the closely related group 2c \textit{WRKYs} \textit{HvWRKY13}, \textit{OsWRKY23}, \textit{AtWRKY43} and \textit{AtWRKY56} is highly correlated and strongly increased in seed stages compared to leaves (Figure 4 and 5). From this we can conclude that these four \textit{WRKY} genes are of high general importance in all of the three plants for seed development as the compared organs, siliques and caryopses or infructescences, differ greatly in their structure and composition.
Discussion

In this study we present the annotation of 45 members of the *WRKY* gene family in barley and classify them according to the WRKY groups 1 to 3 (Figures 1 and 2, Table 1). The comparison to the total number of *WRKYs* in Arabidopsis and rice (Table 2) allowed us to estimate that we identified about 50% of all barley *WRKY* genes. This is in accordance with a study by Ryu and colleagues [6] who found EST sequences for 44% of all *OsWRKY* genes.

In order to transfer the knowledge of WRKY studies in Arabidopsis and rice to barley, we created a phylogenetic tree including the WRKY protein sequences of all three species. This tree allows for functional predictions based on earlier studies. For example, *AtWRKY7* (group 2d) has been shown to bind to calmodulin in a Ca$^{2+}$-dependent manner by a calmodulin-binding domain (CaMBD), which has been found in all other group 2d WRKY proteins of Arabidopsis [32]. According to our alignment tree, *HvWRKY9* is the ortholog of *AtWRKY7* (Additional File 2) and might therefore have similar calmodulin-binding capacity. Even though the sequence of *HvWRKY9* does not span the CaMBD, we could identify such a CaMBD for *HvWRKY7* and *HvWRKY10*. Also, a sequence alignment of all group 2d *OsWRKY* proteins revealed CaMBD in four of six *OsWRKY* proteins (data not shown), pointing to the conserved function of this domain in angiosperms. Interestingly, none of the group 2d *HvWRKY* genes displayed significant changes in expression levels upon powdery mildew infection (Figure 3). This underlines the uniformity of group 2d *HvWRKY* proteins, and also leads to the speculation whether these *HvWRKY* proteins are impaired in the regulation of pathogen response in a calmodulin-binding dependent manner.
For rice and Arabidopsis members of WRKY group 2a an involvement in the response to fungal pathogens [6, 36] has been demonstrated. Recently, HvWRKY1 and HvWRKY2 have been shown to play an important role in the regulation of the response to the fungal pathogen *Blumeria graminis* [8]. Our alignment tree indicates the existence of two further group 2a WRKY proteins in barley, HvWRKY3 and HvWRKY23, which therefore are also potential regulators of pathogen response. In fact, the expression levels of all group 2a *HvWRKY* genes alter upon powdery mildew infection (Figure 3).

Interestingly, three out of six rice members of the monocot-specific subclade of group 2c having AtWRKY50 as a basal member show increased expression levels upon infection with the fungus *Magnaporthe grisea* [6]. The same is true for at least two barley members, namely *HvWRKY5* and HvWRKY20, which show changes in expression levels in *Blumeria graminis* challenged leaves (Figure 3). This could indicate that the diversification of this particular subclade of WRKY transcription factors might have occurred in response to pathogen stress. As all so far identified pathogen-responsive OsWRKYs (OsWRKY7, -10, -67) have corresponding orthologs in barley, these proteins are also potential regulators of pathogen response.

Xie and co-workers [37] showed synergistic interactions of OsWRKY71 (group 2a) and OsWRKY51 (group 2d) in regulation of *Amy32b* gene expression in rice aleurone cells. Parts of this work have been used in a comparative analysis in barley, where HvWRKY1 (HvWRKY38) was shown to be involved in regulating *Amy32b* expression [29]. In fact, based on our alignment, OsWRKY71 and HvWRKY1 were shown to be putative orthologs. However, an interaction partner of HvWRKY1 with functional similarities to OsWRKY51 has so far not been described [29], although one may
postulate that the ortholog to OsWRKY51, HvWRKY10, constitutes a good candidate as an interaction partner to HvWRKY1. In summary, the sequence comparison of WRKY proteins of different species aids in assigning functions to different WRKY proteins and allows for target-oriented hypothesis testing.

Microarray-based studies have become a highly valuable source for functional genomics [38] and huge publicly available datasets allow scientists to monitor the expression of their genes of interest. In this study, we performed a cross-species expression analysis using the microarray data of developmental studies in barley and Arabidopsis [34, 35]. We considered similarities in expression profiles as indicator for evolutionary conserved functional similarities of orthologous genes. However, our study was limited by the fact that only 20 of the HvWRKY genes and 61 of the AtWRKY genes are represented by probesets on the respective microarrays. Also, there was an obvious risk of not comparing true orthologs as only about 50% of all HvWRKY genes could be identified (Table 2). In addition, growth conditions for barley and Arabidopsis plants differed and some of the organs we considered homologous harbor plant lineage-specific functions (see [34, 35]). Nonetheless, we observed an over-average Pearson correlation of +0.24 in the gene expression of putative WRKY orthologs (Table 4, Figure 4) between these two species. This is consistent with the results of a genome-wide expression comparisons of five developmental stages of Arabidopsis and rice, which returned correlation coefficients in the same range [39]. Also, a comparative study of the expression of members of the basic helix-loop-helix transcription factors in Arabidopsis and rice indicates conserved expression patterns of orthologs in four corresponding plant organs [40]. When only considering the best pairing WRKY orthologs, the average
Pearson correlation for gene expression increases to +0.37. This is mostly due to a reduction of putatively orthologous gene-pairs exhibiting non-correlating and more random expression patterns (Additional File 5). Several reasons might explain why some of these putative orthologs of barley and Arabidopsis show more random or negatively correlated expression profiles. First, the function of the orthologous WRKY genes might have changed during evolution, mirrored by changes in expression patterns. Moreover, the expression of these WRKY genes might not be related to any developmental processes but e.g. being activated upon abiotic and biotic stresses. As we only present a minimum number of HvWRKYs, we might as well not have assigned the true barley ortholog to the Arabidopsis WRKY genes investigated, leading to non-correlated expression of non-orthologous genes.

In this study we have shown the potential value of comparative expression analysis between sequenced model species and non-sequenced species of high economic impact. Nowadays, microarrays for several non-sequenced crop species are commercially available. The reported combination of phylogenetic and expressional analysis may offer a valuable tool for future studies and may benefit the understanding of gene functions in these crops as well.

**Conclusions**

The basis for deriving orthology between related proteins of different plant species is their high degree in similarity. Here, we could provide that this orthology is not restricted to sequence similarities alone but reaches out to orthology in expression patterns. We speculate that this observation is not restricted to the WRKY proteins. Nonetheless, the
combination of phylogenetic characterization of HvWRKY genes and the subsequent comparative expression analysis of orthologous genes in Arabidopsis provides a solid basis for further comparative functional genomics studies of this gene family in the cereal crop barley.

Materials and methods

Database search and sequence annotation

To find ESTs coding for WRKY proteins in barley (*Hordeum vulgare*), we performed a tBLASTn search [31] on the NCBI dbEST dataset for barley (http://www.ncbi.nlm.nih.gov/BLAST). The WRKY domain sequences of seven Arabidopsis (*Arabidopsis thaliana*) WRKY family members (AtWRKY1,18, 6, 8, 7 14 and 30), each representing one of the WRKY subgroups described by Eulgem et al. [16], were used as query sequences. Default settings were used except for a deselected low complexity filter. 142 best hits were further processed. To confirm the obtained cDNA sequences, the nucleotide sequences were translated into amino acid sequences, which were then examined for the existence of the heptapeptide WRKYGEK and its observed variations [20]. 20 sequences were removed as they did not contain any WRKY sequence. The remaining ESTs were aligned using ClustalW [41] and manually screened for redundant sequences, resulting in 45 unique WRKY protein coding sequences. The 45 sequences were used as query sequences in a BLASTn search against PlantGDB’s EST assemblies, so called PUT in order to obtain maximum sequence length of overlapping ESTs (http://www.plantgdb.org/) [42].
A keyword search on NCBI returned 7 mRNA and one genomic sequence of the species *Hordeum vulgare* [GenBank:AJ536667, GenBank:AJ853838 GenBank:AJ853839, GenBank:AJ853840, GenBank:AJ853841, GenBank: AJ853842, GenBank:AY323206, GenBank:AY541586]. Multiple sequence alignment revealed that we had identified EST representing all of them. To create a uniform nomenclature with constitutive numbering, the sequences annotated as “Hordeum vulgare mRNA for putative WRKY protein” 2 [GenBank:AJ853838] and 5 [GenBank:AJ853841] were named HvWRKY2 and 5, respectively. The GenBank sequence AY541586 (“Hordeum vulgare WRKY transcription factor (wrky38) gene”) was published as HvWRKY-38 [28] and is identical with the “Hordeum vulgare mRNA for putative WRKY1 protein” [GenBank:AJ536667]. We named both sequences HvWRKY1 and did not annotate any sequence as “HvWRKY38” to prevent confusion. Sequences “Hordeum vulgare partial mRNA for putative WRKY3 protein” [GenBank:AJ853839] and “Hordeum vulgare partial mRNA for putative WRKY4 protein”[GenBank:AJ853840] showed 100% nucleotide sequence similarity to longer EST sequences which were consequently annotated as HvWRKY3 [GenBank:EF488104] and HvWRKY4 [GenBank:EF488105]. The sequence “Hordeum vulgare partial mRNA for putative WRKY6 protein” [GenBank:AJ853842] had been annotated with a single C-terminal WRKY domain. We extended the sequence to the N-terminal WRKY domain by sequencing and annotated the sequence as HvWRKY6 [GenBank:EF488106]. The sequence “Hordeum vulgare SUSIBA2 (susiba2) mRNA” [GenBank:AY323206] [15] was named HvWRKY46. All remaining identified barley WRKY sequences were annotated on NCBI as HvWRKY7 to -37 and HvWRKY39 to -45 (see Table 1 for accession numbers).
Phylogenetic analysis

To gain insights into the evolutionary relationships of barley WRKY proteins and WRKY proteins of Arabidopsis and rice (*Oryza sativa*), we performed multiple alignments of the species WRKY proteins. Pairwise alignment using NCBI blastP (http://www.ncbi.nlm.nih.gov/blast/Blast.cgi) was used for paraphyletic grouping of HvWRKY36, HvWRKY6 and OsWRKY82 into their respective WRKY groups. Sequence information for 72 Arabidopsis WRKY proteins (AtWRKY) and 81 rice WRKY proteins (OsWRKY, as annotated by [21]) was retrieved from NCBI. WRKY proteins of *Physcomitrella patens* [GenBank:AAL78681, GenBank:ABI64128 GenBank: ABI64129, GenBank: ABI64130, GenBank: ABI64131, GenBank: ABI64132, GenBank: ABI64133, GenBank: ABI64134, GenBank ABI64135, GenBank:ABI64136], *Giardia lamblia* [GenBank:XM_765980] and *Dictyostelium discoideum* [GenBank:XM_638694] as well as zinc-finger domains of three additional members of the WRKY-GCM1superfamily, namely Arabidopsis Mutator-like transposase [GenBank:AAC97215], *Homo sapiens* CRA a FLYWCH-type zinc finger 1 [GenBank:EAW85450] and *Mus musculus* FLYWCH-type zinc finger 1 [GenBank;AAH38031] were included to form basal outgroups. The 60 amino acid spanning WRKY core domains described by Eulgem and co-workers [16] of all proteins including all barley WRKY proteins (HvWRKY) was used to create multiple protein sequence alignments using ClustalW (http://www.ebi.ac.uk/clustalw/) [41]. Default settings were applied for the alignment in Figure 1. For Additional Figure1, default settings were used except for using a PAM matrix, a gap opening penalty of 2 and a gap
extension penalty of 0.05. The phylogenetic trees in Figure 2 and Additional File 2 were strict consensus trees of up to 200000 trees calculated with three different programs, ClustalW (http://www.ebi.ac.uk/clustalw/), Phylip (http://evolution.genetics.washington.edu/phylip.html) and PAUP (http://paup.csit.fsu.edu/), which all resulted in identical tree topologies. Bootstrap values have been calculated from 1000 iterations using Phylip (http://evolution.genetics.washington.edu/phylip.html). Phylogenetic trees were drawn using TreeView software version 1.6.6 (http://taxonomy.zoology.gla.ac.uk/rod/treeview.html). The tree in Additional Figure 2 was rooted with the zinc finger domain of Arabidopsis mutator-like transposase (AtMutTrans) [GenBank:AAC97215] as outgroup.

**Microarray based expression analysis**

For expression analysis, publicly available microarray data of the Affymetrix Barley1 GeneChip were used. The annotations of the Barley1 GeneChip were updated by the HarvEST database (http://harvest.ucr.edu/; version 1.51). In order to identify probe sets for the barley WRKY genes described in this study, BLAST searches against the Barley1 GeneChip assemblies were performed on the Affymetrix Netaffx website (http://www.affymetrix.com/analysis/index.affx). For 20 of the characterized WRKY genes, probesets could be assigned unambiguously. HvWRKY2 and HvWRKY46 are represented by three and two probesets, respectively, so that a total number of 23 probesets was used to examine the expression of barley WRKY genes. Additional probesets which - according to the HarvEST annotations - have rice or Arabidopsis
WRKY genes as best matches were not considered further as they could not be related to any WRKY domain spanning barley EST. The Affymetrix CEL files of powdery mildew treatment experiments (accession numbers BB4 and BB7) as well as the developmental baseline (accession number BB3; [34]) were downloaded from BarleyBase (http://www.plexdb.org/) [26]. Affymetrix CEL files of the developmental baseline of Arabidopsis [35] were retrieved from TAIR (http://www.arabidopsis.org; accession number ME00319). Analysis of microarray data was performed using GeneSpring software version 7 (Agilent, http://www.agilent.com).

A per-Chip normalization to the median was applied to obtain comparability. The arrays were adjusted for background optical noise using sequence dependent robust multi-array averaging (GC-RMA) software [43] and normalized using quantile normalization. From the resulting signal intensities, fold change values were calculated.

Expression data were extracted for homologous plant organs (1 - 10) for barley and Arabidopsis, respectively: 1 (coleoptile/cotyledon), 2 (mesocotyl/hypocotyl), 3 (radicule/seedling roots), 4 (leaf/leaf), 5 (root/root), 6 (inflorescence/inflorescence), 7 (anthers/stamen), 8 (caryopsis 5dpa/seed stage 6), 9 (caryopsis 10dpa/seed stage 8) and 10 (caryopsis 15dpa/seed stage 10).

The correlation for gene expression levels between corresponding organs from barley and Arabidopsis were analyzed by calculating the Pearson correlation of the normalized signal intensities with spreadsheet analysis of Microsoft Excel. As a control, the signal intensities of 100 randomly chosen probesets of the Barley1 GeneChip were compared to 100 randomly chosen Arabidopsis genes represented by the ATH1 GeneChip. For both
comparisons, the average correlation was calculated. Students t-test was used to test the statistical significance of the difference in correlation of the two datasets.

**Plant material**

Barley and Arabidopsis (Catania-0) plants were grown in 10 cm pots in soil in the green house (long day; ~24°C; 40% rel. humidity). Rice, Oryza sativa japonica cv. Nipponbare, was grown in 60 cm pots in water logged soil in the green house (long day; ~27°C; ~70% rel. humidity).

**RNA extraction**

Leaf and root plant material from four week old barley and rice plants was harvested at the middle of the day. Infructescences were harvested at the age of eleven days post anthesis. Tissues of Arabidopsis were harvested from three week old Catania-0 plants. Plant material was pooled from several plants. Total RNA was isolated according to the RNAqueous system (Ambion; barley) or Trizol (Invitrogen; Arabidopsis and rice) following the manufacturer’s recommendations. Plant RNA isolation Aid solution (Ambion) was added to all samples. After extraction, the samples were DNase treated using the DNA-free kit (Ambion). RNA concentration was measured using a NanoDrop ND-1000 (Fisher Scientific).

**RT-PCR**

A two-step semi-quantitative RT-PCR method was used to measure gene expression. First strand cDNA synthesis was performed using SuperScript III Reverse Transcriptase
(Invitrogen) on 2 - 5 µg total mRNA and oligo dT primers (Invitrogen). The relative amount of gene expression for different WRKY genes was determined within linear amplification ranges. One or two microliters cDNA template (equivalent to 50 and 100ng RNA) were used for each PCR. Number of cycles used and primer sequences for WRKY gene and Actin controls are listed in Additional File 6.

Signal intensities of each of the RT-PCR amplicons have been quantified using KODAK Molecular Imaging Systems and Software. Normalized signal intensities have been calculated by dividing the intensity for each WRKY gene by the intensity of its respective Actin control signal. Normalized fold intensities were calculated for reasons of comparison from the normalized signal intensities divided by the average of the normalized signal intensities of the three organs (mean expression intensity).

**Sequencing of two Barley WRKY genes**

Multiple sequence alignments indicated, that EST sequences BU990739 and BM370186 might possibly represent parts of the C-terminal and the N-terminal WRKY domains of HvWRKY6. To obtain the interjacent sequence, a PCR was performed with HvWRKY6 forward primer 5’-GCATATTCAGAAGGGCTGCCGAG-3’ and HvWRKY6 reverse primer 5’-CTGCCCATACTTACGCCATCTG-3’.

Sequence information of the WRKY domain for HvWRKY9 was available (CB879962) and multiple alignments indicated that HvWRKY9 belonged to the subgroup of HARF-domain containing WRKY proteins (for HARF sequence information, see [16]). To extend the sequence further to the N-terminus towards the HARF domain, a PCR was performed with HvWRKY9 forward primer: 5’-
AATGGCAGACAAGGCCATGCTAGG-3’ and HvWRKY9 reverse primer 5’-ATACTTCCGCCACGAGAATTCA-3’. In a sequencing approach, DNA of barley leaves was isolated using a protocol by Edwards et al. and treated with RNase H [44, 45]. PCR was performed with Phusion-Polymerase according to the manufacturer’s recommendations (Finnzymes). The PCR products were gel-purified and cloned into a pCR 2.1-TOPO vector (Invitrogen) for sequencing at 4BaseLab (Reutlingen/Germany).

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**Figure legends**

**Figure 1.** Multiple alignment of the identified barley (HvWRKY) and selected Arabidopsis (AtWRKY) and rice (OsWRKY) WRKY domain sequences.

N-terminal and C-terminal WRKY domains of group 1 WRKY proteins are indicated as _NTD and _CTD, respectively. The conserved WRKY amino acid signature is highlighted in bold letters, the amino acids forming the zinc-finger motif are displayed in grey, gaps are marked with dashes.

**Figure 2.** Unrooted phylogenetic tree of identified barley WRKY proteins. The tree was calculated on the basis of WRKY domain sequences of the barley (HvWRKY) and selected Arabidopsis (AtWRKY) and rice (OsWRKY) protein sequences as representatives for the different groups. WRKY groups and subgroups 1 to 3 are highlighted in different colors. Color coded grouping of HvWRKY36, HvWRKY6 and OsWRKY82 is based on analysis with sequences extending over the WRKY domain consensus.

**Figure 3.** *HvWRKY* gene expression upon infection with *Blumeria graminis* (Bgh).

Normalized signal intensities of 23 probesets representing HvWRKY genes are displayed for microarray experiments BB4 and BB7 (according to BarleyBase [26]). Experimental samples and timepoints are indicated on the x-axis. Mlo, mlo5 and Mla1 represent different barley genotypes, Bgh_5874 represents a particular strain of powdery mildew (see BarleyBase [26] for experimental details). Fold changes compared to the control (BB7) or timepoint zero (BB4) are color coded as indicated. *HvWRKY* gene probesets are
arranged according to WRKY groups 1 to 3. Several probesets representing the same
gene are named _a to _c.

**Figure 4.** Selected expression profiles barley *WRKY* genes (*HvWRKY*) and their designate orthologs in Arabidopsis (*AtWRKY*).
The signal intensities of the developmental baseline experiments by Druka *et al.* [34] and Schmid *et al.* [35] are normalized to the mean expression values and are plotted in log-scale for different homologous plant organs (1-10). The respective organs of barley and Arabidopsis are: 1 (coleoptile/ cotyledon), 2 (mesocotyl/ hypocotyl), 3 (radicule/ seedling roots), 4 (leaf/ leaf), 5 (root/ root), 6 (inflorescence/ inflorescence), 7 (anthers/ stamen), 8 (caryopsis 5dpa/ seed stage 6), 9 (caryopsis 10dpa/ seed stage 8) and 10 (caryopsis 15dpa/ seed stage 10).

**Figure 5.** Expression of selected barley *WRKY* genes (*HvWRKY*) and their designate orthologs in Arabidopsis (*AtWRKY*) or rice (*OsWRKY*).
The normalized fold difference (upper diagram) of the signal intensities of each amplicon has been calculated from the RT-PCR experiments (lower diagram) to estimate the expression trajectories of *WRKY* genes relative to its *Actin* controls over three different organs in three plant species. RT-PCR analysis (lower panels) with mRNA of barley, Arabidopsis and rice isolated from homologous organs (roots, left; leaves, middle; infructescence, right).
### Table 1. Identified WRKY proteins in barley and their putative orthologs in Arabidopsis and rice based on phylogenetic studies of their respective WRKY domain sequences.

| HvWRKY       | Accession number | WRKY group | AtWRKY ortholog | OsWRKY ortholog |
|--------------|------------------|------------|-----------------|-----------------|
| HvWRKY1/38   | AJ536667         | IIa        | 18,40,60        | 28,71           |
| HvWRKY2      | AJ853838         | IIa        | 18,40,60        | 28,71           |
| HvWRKY3      | EF488104         | IIa        | 18,40,60        | 76              |
| HvWRKY4      | EF488105         | III        | 54,70           | 47              |
| HvWRKY5      | AJ853841         | IIc        | 50              | 77              |
| HvWRKY6      | EF488106         | I          | 1               | 82              |
| HvWRKY7      | DQ840406         | IID        | 11,17           | 68              |
| HvWRKY8      | DQ840407         | IID        | 39,74           | *               |
| HvWRKY9      | DQ840408         | IID        | 7               | *               |
| HvWRKY10     | DQ840409         | IID        | *               | 51              |
| HvWRKY11     | DQ840410         | IID        | *               | 6               |
| HvWRKY12     | DQ840411         | IIc        | 75              | 72              |
| HvWRKY13     | DQ840412         | IIc        | 56              | 23              |
| HvWRKY14     | DQ840413         | IIc        | 13              | 79              |
| HvWRKY15     | DQ840414         | IIc        | 57              | 3,29            |
| HvWRKY16     | DQ840415         | IIc        | 50              | 67              |
| HvWRKY17     | DQ840416         | IIc        | 50              | 10              |
| HvWRKY18     | DQ840417         | IIc        | 50              | 26,59           |
| HvWRKY19     | DQ840418         | IIc        | 50              | 7               |
| HvWRKY20     | DQ840419         | IIc        | 50              | 7               |
| HvWRKY21     | DQ863105         | III        | 30,41,53        | 15              |
| HvWRKY22     | DQ863106         | III        | 30,41,53        | 74              |
| HvWRKY23     | DQ863131         | IIa        | 18,40,60        | 28,71           |
| HvWRKY24     | DQ863108         | I          | 30,41,53        | 63              |
| HvWRKY25     | DQ863109         | III        | 30,41,53        | 81              |
| HvWRKY26     | DQ863110         | III        | 30,41,53        | 15              |
| HvWRKY27     | DQ863111         | III        | 30,41,53        | 19              |
| HvWRKY28     | DQ863112         | III        | 30,41,53        | 74              |
| HvWRKY29     | DQ863113         | III        | 30,41,53        | 15              |
| HvWRKY30     | DQ863114         | III        | 30,41,53        | 69              |
| HvWRKY31     | DQ863115         | III        | 54,70           | 48              |
| HvWRKY32     | DQ863116         | III        | 54,70           | 45              |
| HvWRKY33     | DQ863117         | III        | *               | 46              |
| HvWRKY34     | DQ863118         | III        | *               | 46              |
| HvWRKY35     | DQ863119         | I          | 19              | *               |
| HvWRKY36     | DQ863130         | IIc        | 49              | 17              |
| HvWRKY37     | DQ863121         | IIb        | 72              | 73              |
| HvWRKY39     | DQ863122         | IIe        | 65              | 13              |
| HvWRKY40  | DQ863123 | I   | 44  | 38  |
|-----------|----------|-----|-----|-----|
| HvWRKY41  | DQ863124 | I   | *   | *   |
| HvWRKY42  | DQ863125 | I   | 2   | 80  |
| HvWRKY43  | DQ863126 | I   | 33  | 24  |
| HvWRKY44  | DQ863129 | Ile | 22  | 39  |
| HvWRKY45  | DQ863128 | Ile | 69  | 14  |
| HvWRKY46  | AY323206 | I   | 20  | 78  |
| (Susiba2) |          |     |     |     |

* ortholog could not be assigned with significance
Table 2. Number of WRKY proteins of Arabidopsis (AtWRKY), rice (OsWRKY) and barley (HvWRKY) in the WRKY subgroups 1 to 3*.

| WRKY group | AtWRKY | OsWRKY** | HvWRKY |
|------------|--------|----------|--------|
| 1          | 15     | 16       | 8      |
| 2a         | 3      | 4        | 4      |
| 2b         | 8      | 7        | 1      |
| 2c         | 17     | 20       | 11     |
| 2d         | 7      | 6        | 5      |
| 2e         | 8      | 8        | 3      |
| 3          | 14     | 20       | 13     |
| Total      | 72     | 81       | 45     |

* according to Eulgem et al. [16]
** according to Xie et al. [21]
Table 3. Probesets of *HvWRKY* genes on the Barley1GeneChip.

| *HvWRKY* | WRKY group | Barley1 Chip probeset |
|-----------|------------|-----------------------|
| *HvWRKY6* | I          | CONTIG14308_AT        |
| *HvWRKY41* | I          | CONTIG12033_AT        |
| *HvWRKY42* | I          | CONTIG15657_AT        |
| *HvWRKY46* | I          | CONTIG7243_AT; RBAAL15J13_S_AT |
| (Susiba2) *HvWRKY1* | IIa | CONTIG4386_AT        |
| (HvWRKY-38) *HvWRKY2* | IIa | CONTIG4387_AT; HB25K10R_S_AT; HB25K10R_AT |
| *HvWRKY3* | IIa        | EBRO02_SQ004_H10_AT   |
| *HvWRKY23* | IIa        | CONTIG21110_AT        |
| *HvWRKY5* | IIc        | CONTIG18462           |
| *HvWRKY13* | IIc        | CONTIG13268_AT        |
| *HvWRKY20* | IIc        | CONTIG10168_AT        |
| *HvWRKY7* | IIId       | CONTIG7798_AT         |
| *HvWRKY8* | IIId       | CONTIG23011_AT        |
| *HvWRKY9* | IIId       | CONTIG22226_AT        |
| *HvWRKY10* | IIId       | CONTIG16040_AT        |
| *HvWRKY39* | IIe        | CONTIG13375_AT        |
| *HvWRKY4* | III        | CONTIG20450_AT        |
| *HvWRKY30* | III        | CONTIG12005_AT        |
| *HvWRKY32* | III        | S0001000055P18F1_S_AT |
| *HvWRKY34* | III        | CONTIG10471_AT        |
Table 4. Pearson correlation coefficients of the expression profiles of *HvWRKY* genes and their designate *AtWRKY* orthologs.

| *HvWRKY* | *AtWRKY* | Correlation coefficient |
|----------|----------|------------------------|
| *HvWRKY1* | *AtWRKY18* | 0.06 |
|          | *AtWRKY40* | -0.11 |
|          | *AtWRKY60* | -0.18 |
| *HvWRKY2* | *AtWRKY18* | -0.05 |
|          | *AtWRKY40* | -0.10 |
|          | *AtWRKY60* | -0.31 |
| *HvWRKY3* | *AtWRKY18* | 0.21 |
|          | *AtWRKY40* | -0.24 |
|          | *AtWRKY60* | -0.11 |
| *HvWRKY4* | *AtWRKY54* | 0.52 |
|          | *AtWRKY70* | 0.23 |
| *HvWRKY6* | *AtWRKY1* | 0.41 |
|          | *AtWRKY32* | 0.45 |
| *HvWRKY7* | *AtWRKY11* | 0.38 |
|          | *AtWRKY17* | 0.66 |
| *HvWRKY8* | *AtWRKY39* | 0.20 |
|          | *AtWRKY74* | 0.30 |
| *HvWRKY9* | *AtWRKY7* | -0.49 |
|          | *AtWRKY15* | 0.12 |
| *HvWRKY13* | *AtWRKY43* | 0.82 |
|          | *AtWRKY56* | 0.54 |
| *HvWRKY23* | *AtWRKY30* | 0.33 |
|          | *AtWRKY53* | 0.39 |
| *HvWRKY28* | *AtWRKY30* | -0.12 |
|          | *AtWRKY53* | -0.07 |
| *HvWRKY32* | *AtWRKY54* | 0.70 |
|          | *AtWRKY70* | 0.42 |
| *HvWRKY34* | *AtWRKY55* | -0.37 |
| *HvWRKY39* | *AtWRKY65* | 0.88 |
|          | *AtWRKY69* | 0.91 |
| *HvWRKY42* | *AtWRKY2* | 0.78 |
| *HvWRKY46* | *AtWRKY20* | 0.50 |

Average correlation all WRKY gene pairs: 0.24
Average correlation best correlated WRKY gene pairs*: 0.37
Average correlation random genes 0.01

* For each *HvWRKY* gene, the *AtWRKY* gene with the highest correlation value was chosen to calculate the average correlation of best pairs.

** Correlation of 101 random genes of barley and Arabidopsis. Probesets on the Barley1 GeneChip and ATH1 GeneChip were randomly selected and the average of the Pearson correlation of their expression profiles was calculated.
Description of additional datafiles

File name: Additional File 1

File format: .doc

Title of data: Multiple alignment of WRKY_GCM1-like domains

Description of data: Multiple alignment of WRKY_GCM1-like domain amino acid sequences from Arabidopsis (AtWRKY), barley (HvWRKY), rice (OsWRKY), Physcomitrella (PpWRKY), Dictyostelium discoideum (DdWRKY), Giardia lamblia (GlWRKY), Homo sapiens (HsFLYWCH) and Mus musculus (MmFLYWCH). N-terminal and C-terminal WRKY domains of group 1 WRKY proteins are indicated as _N and _C, respectively. The conserved WRKY signature is highlighted in bold letters, the amino acids forming the zinc-finger motif are shaded with grey, gaps in the alignment are indicated by dashes.

File name: Additional File 2

File format: .pdf

Title of data: Phylogenetic tree based on all WRKY_GCM1-like domains considered in the Alignment of Additional File 1.

Description of data: Phylogenetic tree based on the WRKY domain amino acid sequences given in Additional Figure 1. Sequences of Arabidopsis (AtWRKY), barley (HvWRKY), Physcomitrella (PpWRKY), rice (OsWRKY), Dictyostelium discoideum (DdWRKY), Giardia lamblia (GlWRKY) Homo sapiens (HsFLYWCH) and Mus musculus (MmFLYWCH) are arranged in clusters with the zinc finger domain of the
Arabidopsis mutator-like transposase (AtMutTrans) as outgroup. Tree topology has been confirmed by using three different programs. Bootstrap values from 1000 iterations and bigger than 500 are included in the tree. HvWRKY sequences are highlighted in bold letters. Groups and subgoups of WRKY1 to 3 are indicated by bars on the right side.

**File name:** Additional File 3

**File format:** .xls

**Title of data:** Normalized signal intensities of *HvWRKY* genes, *AtWRKY* genes and a set of 101 randomly chosen gene pairs

**Description of data:** The table contains all essential data for the comparative gene expression analysis for barley and Arabidopsis *WRKY* genes as well as control gene pairs on the extracted subsets of the microarray expression datasets of homologous organs by Druka et al. [34] and Schmid et al. [35].

**File name:** Additional File 4

**File format:** .xls

**Title of data:** Signal intensities of RT-PCR experiment with barley, Arabidopsis and rice *WRKY* genes

**Description of data:** Tables contain raw signal intensity values of bands from RT-PCR experiments shown in Figure 5.

**File name:** Additional File 5

**File format:** .pdf
**Title of data:** Distribution of the relative correlation coefficient in randomly chosen and \( HvWRKY-AtWRKY \) gene pairs.

**Description of data:** The relative correlation coefficient was calculated from the normalized signal intensities of the gene pairs and plotted according to its occurrence. Random gene pairs exhibited an average correlation coefficient of 0.01. WRKY gene pairs exhibited a divided curve with two maxima. When only the best pairing orthologs were considered, the first peak of more random distribution is lost.

**File name:** Additional File 6

**File format:** .xls

**Title of data:** Primer sequences used in RT-PCR experiment with barley, Arabidopsis and rice \( WRKY \) genes

**Description of data:** Table containing the sequences of primers, numbers of cycles and expected fragment size of RT-PCR experiments depicted in Figure 5.
The figure presents the normalized fold intensity of various WRKY genes across different plant tissues and developmental stages. The X-axis represents the tissue types: Roots, Leaves, and Inflorescence. The Y-axis represents the normalized fold intensity.

For Barley (Hv), the genes HvWRKY6, HvWRKY9, HvWRKY13, HvWRKY39, HvWRKY46, and HvActin are analyzed.

For Arabidopsis (At), the genes AtWRKY32, AtWRKY7, AtWRKY56, AtWRKY20, and AtActin2 are analyzed.

For Rice (Os), the genes OsWRKY23, OsWRKY13, OsWRKY78, and OsActin1 are analyzed.

The graphs illustrate differential expression patterns for each gene across the different tissues and developmental stages.