

The Unicellular Ancestry of Groucho-Mediated Repression and the Origins of Metazoan Transcription Factors

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Abstract

Groucho is a co-repressor that interacts with many transcription factors playing a crucial role in animal development. The evolutionary origins of Groucho are not clear. It is generally regarded as being a distinct animal-specific protein, although with similarities to the yeast Tup-like proteins. Here, it is shown that Groucho has true orthologs in unicellular relatives of animals. Based on their phylogenetic distribution, and an analysis of ligand-binding residues, these genes are unlikely to be orthologs of the fungal Tup-like genes. By identifying conserved candidate Groucho interaction motifs (GIMs) in nonmetazoan transcription factors, it is demonstrated that the details of molecular interactions between Groucho and transcription factors are likely to have been established prior to the origin of animals, but that the association of GIMs with many transcription factor types can be regarded as a metazoan innovation.

Key words: transcription factor evolution, transcriptional repression, animal evolution.

Introduction

Many of the genes controlling animal development encode a circumscribed set of transcription factor domains, and many of these domains have a pre-metazoan ancestry, with homologs found in nonmetazoan eukaryotes (Sebé-Pedrós et al. 2011; de Mendoza et al. 2013). Although clearly the developmental contexts of these proteins (such as patterning the nervous system) cannot exist in unicellular organisms, what is less clear is how directly analogous their molecular functions are. DNA-binding specificities of several transcription factor domains have been shown to be broadly similar between single celled eukaryotes and animals (Kwon et al. 2012; Nakagawa et al. 2013; Sebé-Pedrós et al. 2013), and this is, to a large extent, expected from the conservation of DNA binding amino acids. If there are functional differences between proteins containing these domains (i.e., they are not, within the limits of their expression patterns, completely interchangeable), they are likely to have involved the protein–protein interactions (Wagner 2007; Copley 2008; Sebé-Pedrós et al. 2013; Hudry et al. 2014).

The proteins known as Groucho in *Drosophila*, and the Transducin Like Enhancers of Split (TLEs) in vertebrates are common interaction partners of animal transcription factors.

The Groucho-like proteins act as transcriptional co-repressors and orthologs are found in all animal genomes (Copley 2005). No orthologs have been reported outside the Metazoa, but in fungi the Tup-like (TUP1 in *Saccharomyces cerevisiae*, tup11 & tup12 in *Schizosaccharomyces pombe*) proteins, that also act as transcriptional repressors, are sometimes regarded as the equivalent of Groucho (Chen and Courey 2000).

The relationship between TUP1-like and Groucho-like genes (for convenience referred to here as Tup and Groucho) has not been well defined. While noting functional similarities, Fisher and Caudy (1998) suggested that “it may be more accurate to consider TUP1 and Groucho proteins as analogous rather than truly homologous”. Flores-Saaib and Courey (2000) mentioned that the overall similarity between Groucho and Tup WD40 regions was not significantly greater than between Groucho and other WD40 repeat containing proteins without functional similarities. Based on a more detailed analysis of the sequences of corresponding repeats, they went on to suggest that the proteins were “structurally and therefore perhaps functionally, related”, and proceeding on this basis, demonstrated similarities of Groucho and TUP1 histone interactions. Pickles et al. (2002) stated that the two were “increasingly considered as functional equivalents”. Other

recent authors have more or less explicitly considered them orthologs—that is, encoded in genes related by speciation events (Matsumura et al. 2012; Asada et al. 2015).

There are, however, marked differences in the biology of Tups and Grouchos. Yeast TUP1 proteins form a functional complex with the TPR repeat containing CYC8 (*ssn6* in *Sc. pombe*) protein (Tzamarias and Struhl 1994), but there is no similar co-factor requirement for Groucho. Groucho interacts with EH1 & WRPW protein motifs from a variety of animal transcription factors (Jennings and Ish-Horowicz 2008). Pearl, Ish-Horowicz and others stated that there were no obvious WRPW motif proteins in yeast, suggesting that yeast transcription factors interact with Tup via amphipathic helices similar to the EH1 motif (Jennings et al. 2006). There are not, however, any reported yeast transcription factors with motifs matching the metazoan EH1 consensus. The metazoan EH1 motif as currently described typically begins with a phenylalanine, or less often tyrosine, with a consensus of FS[VI]xx[IL][LM] (see Copley 2005). Without the F or Y, the motif is poorly specified and large numbers of amphipathic helices would be expected to match.

The importance of Groucho-mediated repression in animal development, its inferred presence in the most recent common ancestor of the animals and absence in other groups, raises the question of its evolutionary origin. The presence of an analogous Tup system in yeasts could, however, potentially shed light on this, if its relationship with Groucho were better understood. To enquire farther into the origins of Groucho, the evolutionary history of Tup and Groucho and their likely molecular interactions were examined, with a particular focus on recently available genomic and transcriptomic data from close unicellular relatives of the animals.

Taxonomic Distribution of Tup and Groucho-like Proteins

The Groucho and Tup proteins are composed of N-terminal coiled-coil domains and a C-terminal 7 bladed β -propeller composed of WD40 repeats. WD40 repeats are widespread in animal and eukaryotic proteins, and their repeating nature makes them particularly prone to mis-alignment, making similarity scores hard to interpret. In contrast, structural superimpositions of the 3D structures of the N-terminal domains of TUP1 and TLE suggest that they are distinct from each other and unique to these proteins (fig. 1)—the superficial resemblance of the coiled coils is not reflected in any statistically significant sequence similarity. I conjecture that proteins containing a TLE_N Pfam region are orthologs of Groucho/TLE and those containing a Tup_N region, orthologs of TUP1, and that these regions can be used as proxies to determine the phylogenetic distribution of their respective genes. Later, phylogenetic analysis of the recovered sequences will show this conjecture to be correct.

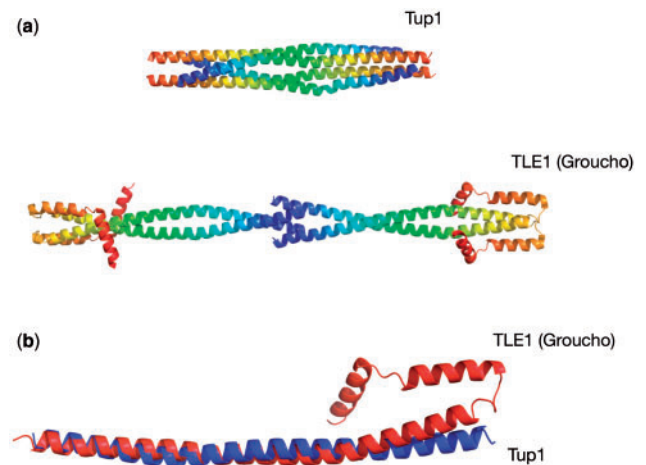


FIG. 1.—The N-terminal domains of Groucho/TLE (PDB: 4om2) and TUP1 (PDB: 3vp8) do not share statistically significant sequence similarity: (a) they adopt different quaternary structures—chains colored from blue at N-terminus to red at C-terminus; (b) both contain coiled-coil regions but with differing degrees of curvature and Groucho/TLE contains additional C-terminal helices.

Accordingly, I searched the nr protein database from the NCBI with the Pfam hidden Markov models Tup_N and TLE_N, using their associated ‘gathering’ threshold bit scores as cut-offs (TLE_N, 24.0 bits; Tup_N 22.4 bits) (Finn et al. 2016). Significant ($E < 0.001$) TLE_N hits were to animals (with the exception of *Sphaeroforma arctica*—discussed below). Among the nonbilaterians, significant TLE_N hits were found to sequences in the cnidarians, *Nematostella vectensis* and *Hydra vulgaris*; the placozoan, *Trichoplax adhaerens*; and the sponge *Amphimedon queenslandica*. Further searches of the draft contigs and scaffolds of ctenophore genomes using representative protein sequences revealed likely Groucho candidates in *Mnemiopsis leidyi* and *Pleurobrachia bachei*. These results show that, irrespective of controversies of the branching order of nonbilaterian animals (Pisani et al. 2015; Whelan et al. 2015), Groucho was present in the common ancestor of all extant animals. This is consistent with the same inference drawn from the phylogenetic distribution of EH1 motifs (Copley 2005). A single TLE_N hit was also identified to a nonmetazoan eukaryote—*Trimastix pyriformis*, represented in the NCBI TSA archive. Hits to Tup_N were primarily to Fungi. Nonfungal eukaryotes included Oomycetes, Amoebozoa, *Naegleria gruberi* (Heterolobosea), *Galdieria sulphuraria* (Rhodophyta), *Ectocarpus siliculosus* (stramenopiles), and *Guillardia theta* (Cryptophyta).

To more closely examine the separation between fungal and animal sequences, I searched both Tup_N and TLE_N against the proteins of the “Origin of Multicellularity Project” (Ruiz-Trillo et al. 2007). For TLE_N (i.e., Groucho/TLE) this resulted in significant matches to *Ameobidium parasiticum* and *Sp. arctica*, but no other species within this

project. Both of these taxa belong to the clade of Ichthyosporia. For Tup_N (i.e. Tup), I found significant matches to proteins from *Spizellomyces punctatus*, *Mortierella verticillata*, and *Allomyces macrogynus*, the three fungal taxa represented in the project. Notably, I was unable to find matches of either Tup_N or TLE_N to choanoflagellate (*Monosiga*, *Salpingoeca*) or filasterean (*Capsaspora owczarzaki*) protein sets, both of which are more closely related to the Metazoa than the ichthyosporians. I also searched Tup_N and TLE_N against protein sets generated from the data in Torruella et al. (2015) (see Methods). This resulted in further matches of TLE_N (Groucho) to Ichthyosporia and Corallochytrium taxa, and Tup_N matches to Nutomonas and Nuclearia species, essentially confirming the phylogenetic distribution of the ‘Multicellularity Project’ set.

The use of Groucho and Tup N-terminal domains conveniently avoids cross matching between different WD40-repeat containing proteins, but it is possible that some *bona fide* orthologs of these proteins diverged before the N-terminus became associated with the WD40 repeats, or subsequently lost the N-terminus. To identify possible orthologs of Tup and Groucho that may be lacking these domains in some species, I also performed searches using alignments of the complete β -propeller domain, implemented as a global-local model using the hmmer 2 software package. Two models were used, one built using representative metazoan Groucho sequences, and the other fungal Tup sequences. These models were searched against the nr database of the NCBI, and the eukaryotic sequence databases described above. As the models represent homologous sequences, there is considerable overlap between their hit lists. I conservatively defined the Groucho hit list to be those sequences scoring higher than the first Tup_N domain containing hit, and the Tup hit list to be those sequences scoring higher than the last nonfungal Tup_N domain containing hit with a positive score. There were no Tup_N hits in the Groucho hit list or TLE_N in the Tup hit list.

By combining the sequences matching the HMMs, I produced an alignment of representatives of the two groups of sequences (see [Supplementary Material](#) online) and hence a single phylogenetic tree using the LG + Gamma model of sequence evolution as implemented in the phym1 package (see Methods). The tree shows clear separation of the Tup and Groucho groups, essentially mirroring the division between Holozoa (i.e., animals and their closest single celled relatives, but excluding fungi) and nonholozoans (fig. 2). The Tup group includes all fungal sequences and the nonfungal eukaryotic sequences mentioned above, concordant with the analysis based on the presence of the Tup_N domain. It also includes additional nonmetazoan non-Tup_N containing proteins, including that from *Fonticula alba*, a member of the Fonticula that together with the Nuclearia forms the sister group to fungi. The only nonholozoans in the Groucho branch of the tree are the TLE_N containing *Trimastix* sequence mentioned above and, in addition, a further *Naegleria* sequence lacking

both TLE_N and Tup_N regions in the N-terminus. Importantly, the *Trimastix* and *Naegleria* sequences do not cluster within the bulk of the holozoan sequences, ruling out simple cross-species contamination, but to the base of the holozoan clade, as would be expected if they were *bona fide* Groucho-like sequences from nonholozoan eukaryotes. Analysis using a Bayesian tree reconstruction approach (see Methods and [Supplementary fig S1, Supplementary Material](#) online) produced similar results, with a strongly supported Groucho clade and *Naegleria* the first diverging lineage within it. In this analysis, however, the *Trimastix* sequence branched within the Teretosporea, although with weak support.

Again, no hits were identified to choanoflagellates or filastereans. This β -propeller-based search also identified orthologs of the human TLE6 gene which, although a Groucho family member, lacks the TLE_N region, and another vertebrate family lacking the TLE_N, exemplified by the human locus 102723796 and mouse Gm21964 gene, for which there does not appear to be evidence of expression.

Structural and Sequence Features Discriminating between Tup and Groucho

Aligning the WD40 repeat containing regions of proteins containing either a Tup_N or TLE_N N-terminal regions enabled the analysis of key residue differences between Grouchos and Tup. In particular, as 3D structures of Groucho (specifically, the human ortholog TLE1) in complex with EH1 and WRPW peptides are available (Jennings et al. 2006), the identity and conservation of residues mediating these interactions could be compared between Groucho and Tup.

I superimposed the structures of the C-terminal domains of TUP1, apo-Groucho, Groucho with EH1 bound, and Groucho with WRPW bound, using the STAMP package (Russell and Barton 1992). Inspection of residues within 5 Å of the EH1 peptide shows that they are found in comparable positions (fig. 3). The major visual difference lies in the orientation of the side chains of TUP1-Y580 and Groucho-F661. This appears to be a consequence of a further substitution, TUP1-L596 versus Groucho-E677. Whereas the charged Groucho side chain is extended away from the bulk of the protein fold, the nonpolar TUP1 residue is half buried within the fold, with the side chain contacting Y580 and re-orienting it towards the ‘pore’ of the β -propeller domain, relative to the orientation of Phe found in all the Groucho crystal structures. If the TUP1 residue configuration were observed in Groucho, there would be a steric clash between the Y580 equivalent and the Phe of the EH1 motif (or W of the WRPW motif) (fig. 3). From this, it appears as though the Tup fold as observed is incompatible with EH1 or WRPW binding in the configurations seen in current crystal structures. This is in accord with the result that no true EH1 or

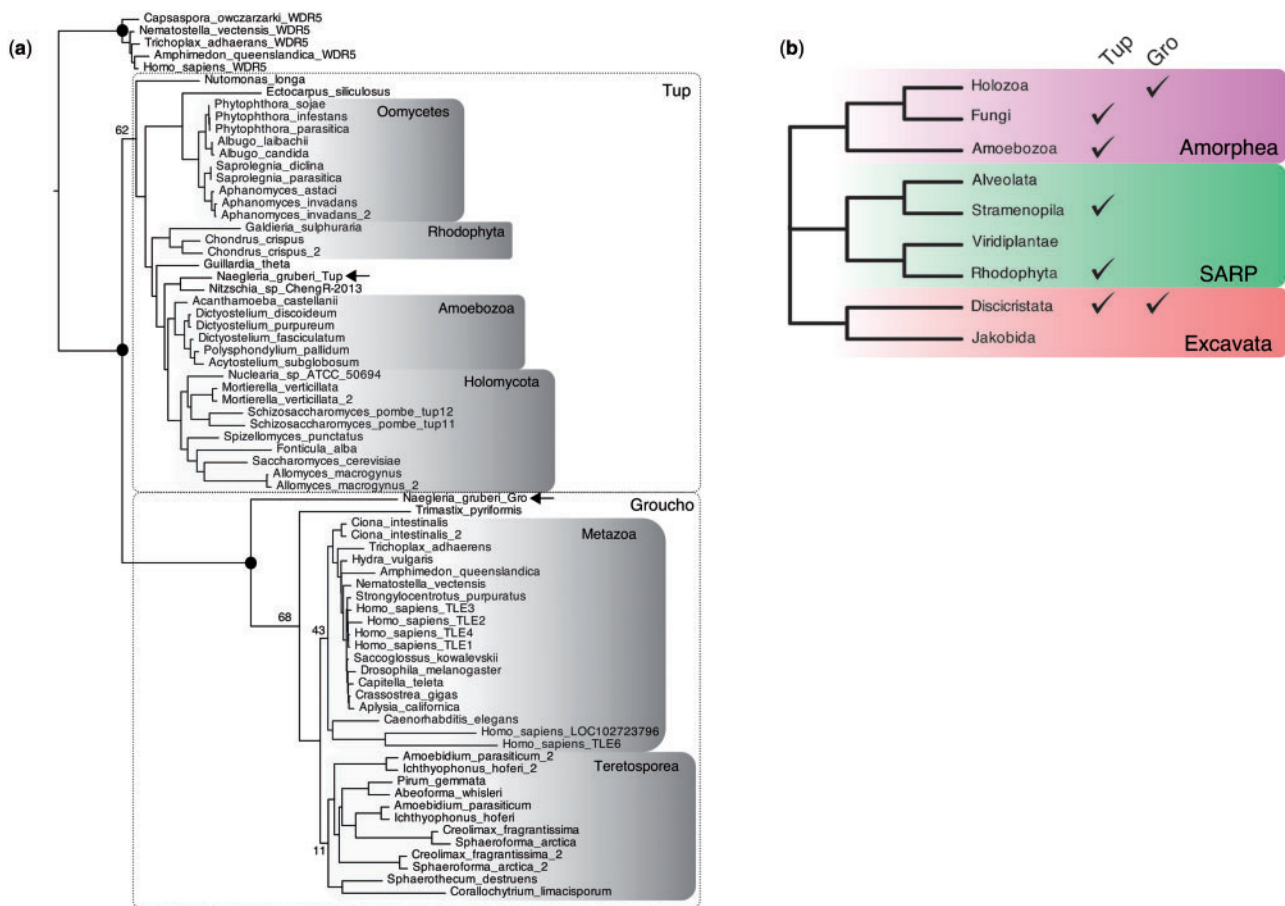


Fig. 2.—(a) Phylogenetic tree of aligned WD40 sequences from Groucho and Tup, with representative WDR5 proteins as an outgroup. The Tup and Groucho clades are boxed and labeled. Sequences within the Tup group typically include a Tup_N N-terminal motif, and those within the Groucho group a TLE_N motif. The *Naegleria* sequences are indicated with arrows. ‘Teretosporea’ is a clade of Ichthyosporia and *Corallochytrium*, defined in Torruella et al. (2015). Black circles on nodes represent complete bootstrap support, with numbers giving values for other nodes central to the discrimination of Tup and Groucho. Sequences that uniquely define the leaves are available in the [Supplementary Material](#) online. (b) The distribution of Groucho and Tup orthologs identified in this study with respect to the three major eukaryotic groups (eukaryotic tree adapted from He et al. 2014).

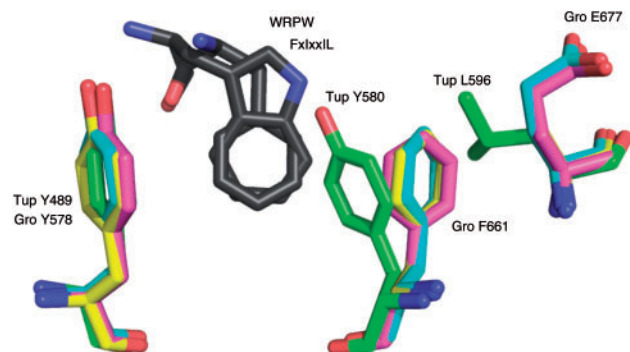


Fig. 3.—Side view of the ligand-binding pocket residues of TLE1 (three structures, with bound EH1 (PDB: 2ce8), WRPW (PDB: 2ce9), and nonbound forms (PDB: 1grx)), with equivalent TUP1 (PDB: 1erj) residues superimposed. The F and W of the TLE1 bound ligands are shown (Sprague et al. 2000; Jennings et al. 2006).

WRPW like motifs have been reported in yeast transcription factors (although see below).

This residue dichotomy (Y,L in Tup-like and F,E in Groucho-like) is conserved (fig. 4). Proteins that have a TUP_N domain have the Y,L pair, whereas those having a TLE_N have the F,E pair, with the exceptions of the most divergent single celled eukaryotic taxa and the vertebrate TLE6 orthologs. As the remainder of the protein binding pocket, interacting with the other ligand residues, appears conserved, it is possible that Tup-like proteins may be able to bind EH1 like motifs that lack the initial Phe residue.

Tup Interaction Motifs in Fungi

The *S. cerevisiae* gene MAT α 2, a homeobox containing transcription factor and regulator of mating type genes, interacts with the WD40 region of TUP1. Mutation of N-terminal

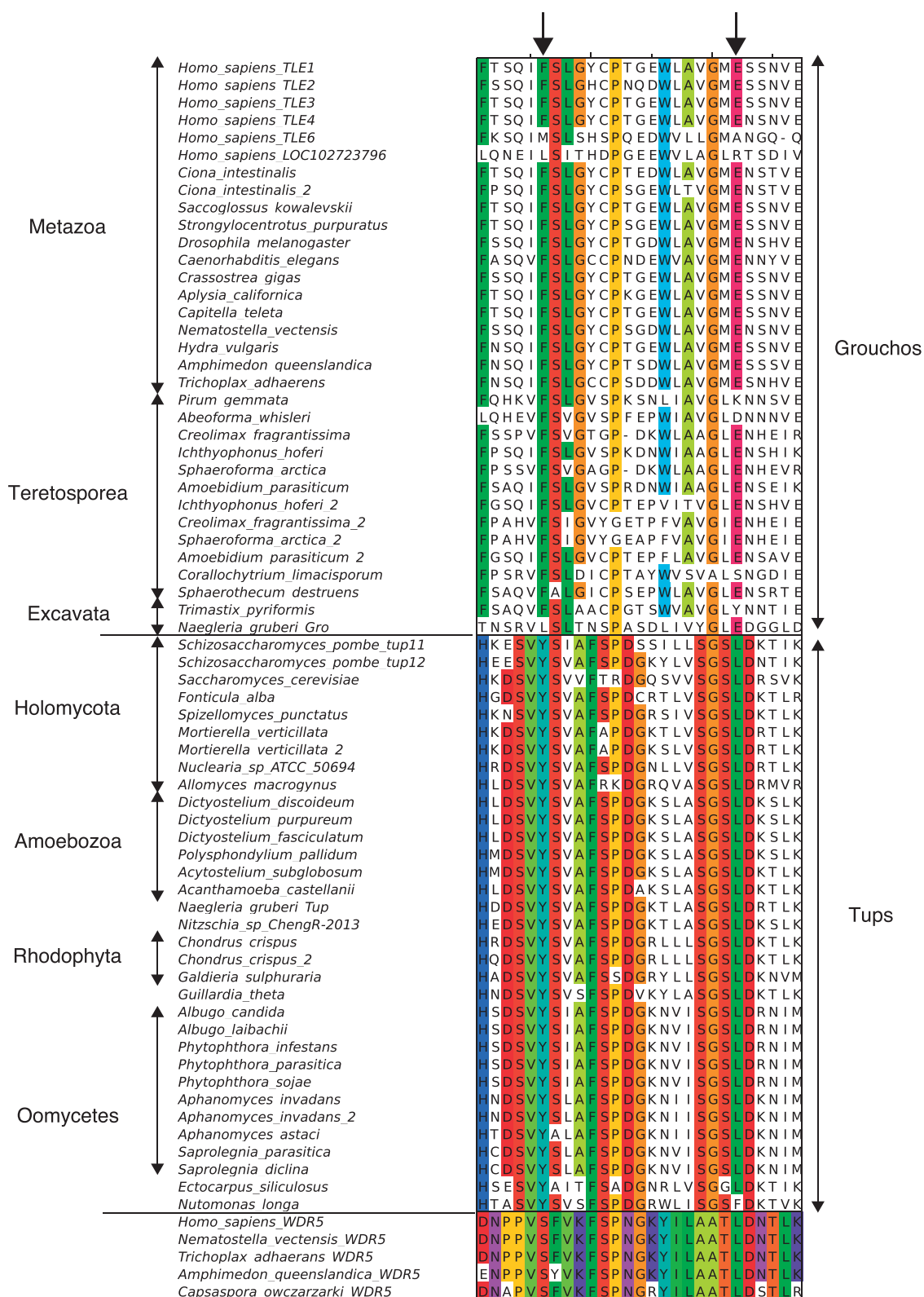


Fig. 4.—Sequence conservation of the ligand-binding pocket of the Groucho and Tup proteins (outgroup WDR5 members are also shown). The region is extracted from the full multiple sequence alignment, columns that are >80% identical within a class are colored by amino acid type (Taylor 1997). The dichotomous F,E (in Groucho) and Y,L (in Tup) residues, likely to contribute to ligand recognition, are marked with arrows.

residues Ile4, Leu9, or Leu10, (or also Gly71) disrupts the interaction with TUP1 (Komachi et al. 1994). The characteristic spacing of these residues (i.e., IxLxxLL) is obviously reminiscent of the EH1 motif (FxLxxIL). This N-terminal motif is well conserved in other yeast MAT α 2 orthologs, and has been recently interpreted as a modified version of the EH1 motif by Bürglin and Affolter (2015).

In order to screen for similar motifs in a well-characterized fungal genome, I searched protein alignments of orthologous genes from *Schizosaccharomyces* group genomes (*Sc. pombe*, *Schizosaccharomyces japonicus*, *Schizosaccharomyces octosporus*, and *Schizosaccharomyces cryophilus*) (see Methods). Instances of the motif [VI]xx[IL][LM] (essentially the highest scoring residue types of the EH1^{hox} motif without the first two residues) that matched the *Sc. pombe* sequence in nondomain regions of proteins containing transcription factors domains, and where the motif was conserved in the *Schizosaccharomyces* group alignment, were considered further.

The pombase database (McDowall et al. 2015) annotates 90 genes with a molecular function of 'RNA polymerase II core promoter proximal region sequence-specific DNA binding' (GO:0000978), of which 60 are represented in the set of aligned orthologs. Of these, 10 contain [VI]xx[IL][LM] motifs conserved in all pombe group species, and eight are recorded as interacting with tup11 or tup12 (the pombe TUP1 orthologs) in the biogrid database (Oughtred et al. 2016). Of the proteins containing the motif, three, *fep1*, *res1* and *scr1*, interact with tup11/12 according to biogrid, with an additional one (*sak1*) predicted as interacting via homology to an *S. cerevisiae* interacting pair in the STRING database (Szkłarczyk et al. 2015). These numbers suggest an enrichment of the conserved motif in transcription factors that interact with tup11/tup12 relative to those that do not ($P=0.0343$, Fisher's exact test), but it must be noted that as the total number of interactors and genes are small, the result is not especially robust. Furthermore, no studies have specifically focussed on the protein-interaction partners of tup11/tup12 in pombe (or TUP1 in *S. cerevisiae*), leading to the possibility that there are considerable numbers of interacting partners yet to be discovered.

Groucho Interaction Motifs in Ichthyosporean Transcription Factors

The analyses of the N-terminal domains and WD40 repeat regions show that orthologs of Groucho are present in the Ichthyosporeans *A. parasiticum* and *Sp. arctica*. If unicellular Groucho orthologs have the same molecular function, as indicated by the conservation of the key Y,E residue pair in the WD40 domain, we would expect to be able to identify proteins containing EH1 or WRPW motifs. By analogy with metazoan Grouchos, we might further expect these motifs to be preferentially present in transcription factor proteins.

To better understand the role of these proteins, I searched a database of proteins from taxa in the evolution of multicellularity project, with the EH1^{hox} hidden Markov model described in Copley (2005). As these motifs typically occur in nonprotein domain contexts, I first masked known Pfam domains. Retrieving hits containing known transcription factor domains (Wilson et al. 2008), eight out the top 10 were found in *Amoebidium* and *Sphaeroforma*, the only genomes in the set that encode groucho-like proteins. As these sequences are all uncharacterized experimentally, I sought evidence of function via evolutionary constraint on sequence evolution by searching for orthologs and paralogs that shared these putative EH1 motifs. In addition to the 'evolution of multicellularity project' proteins, I searched proteins generated from the assembled transcriptomes of the 'Close Relatives of Animals and Fungi' project (Torruella et al. 2015).

Two paralogs within *Amoebidium* mutually supported each other, showing conservation of EH1 motifs in the absence of conservation of surrounding sequence (supplementary fig. S2a, Supplementary Material online). A protein from *Sphaeroforma*, including an N-terminal MYND ZnF and C2H2 Zn fingers contained an EH1 motif that was conserved in an orthologous *Creolimax fragrantissima* sequence (supplementary fig. S2b, Supplementary Material online). An *Amoebidium* sequence including Ankyrin repeats and GATA ZnF had an ortholog in *Ichthyophonus hoferi*, showing conservation of the EH1 motif (supplementary fig. S2c, Supplementary Material online). An additional GATA ZnF protein from *Sp. arctica* with an N-terminal EH1 motif had readily identifiable orthologs in *Creolimax*, *Amoebidium*, *Pirum gemmata*, and *Abeoforma whisleri*. An EH1 motif was identifiable in the *Creolimax* ortholog. The *Amoebidium*, *Pirum*, and *Abeoforma* sequences did not contain EH1-like motifs, but instead, and remarkably, conserved WRPW motifs at equivalent positions, suggestive of convergent evolution of distinct binding motifs within orthologous proteins (supplementary fig. S2d, Supplementary Material online).

The Groucho ortholog in *N. gruberi* does not contain the Tyrosine of the Y,E pair, but instead Leucine. Despite the availability of predicted proteins for the complete genome sequence, I was not able to detect significant matches of the EH1 or WRPW motifs associated with *Naegleria* transcription factor domains. Similarly, no EH1 or WRPW matches were detected in the available transcripts from *T. pyriformis*, where the F and Y amino acids correspond to the Y,E pair.

Groucho Interaction Motifs in Nonbilaterian Metazoan Transcription Factors

The major metazoan associations of EH1 motifs are with the homeobox, forkhead, and T-box transcription factor domains (Copley 2005). Of these associations, all are found in sponges and ctenophores, with only the EH1 forkhead association

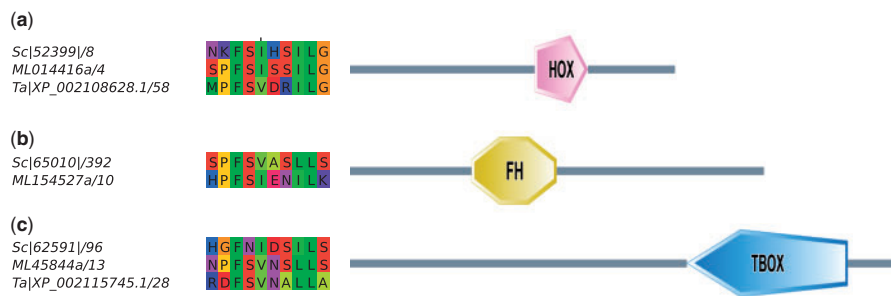


FIG. 5.—Example associations from nonbilaterian metazoans of EH1 motifs with (a) Homeobox, (b) Forkhead, and (c) T-box domains. Sc = *Sycon ciliatum*, a sponge; ML = *Mnemiopsis leidyi*, a ctenophore; Ta = *Trichoplax adhaerens*. Sequence accessions correspond to the databases described in the Methods. Domain diagrams represent the *Sycon* sequences.

being absent from *Trichoplax* (fig. 5). The [WF]RP[WY] motif is associated with HLH and Runt transcription factors. The HLH association is present in sponges, ctenophores, and *Trichoplax*. The [WF]RP[WY] Runt association is found in *Mnemiopsis* (ML03045a), but not in available sponge or *Trichoplax* sequences. Interestingly, the *Mnemiopsis* T-box protein ML45844a encodes a T-box, EH1 motif, and a C-terminal WRPW motif.

Notably, neither the EH1 nor WRPW-type motif is found associated with GATA transcription factors, as found in *Ichthyosporea*. Although there are associations between EH1 motifs and C2H2 Zinc fingers in bilateria, none are found in the nonbilaterian Metazoa investigated here. There is thus a discontinuity between single-celled eukaryotes and the Metazoa.

Discussion

The data presented here show that the origins of the metazoan transcriptional co-repressor Groucho predate the Metazoa. I have identified likely Groucho orthologs in the single celled eukaryotes of the Ichthyosporean clade, and further, identified Ichthyosporean transcription factors that contain conserved Groucho Interaction Motifs (GIMs). The Ichthyosporean transcription factors with GIMs have no obvious relationships to the typical metazoan proteins containing GIMs, suggesting that the quantitative expansion in transcription factor numbers in the animal stem lineage (de Mendoza et al. 2013) co-occurred with a re-wiring of protein-protein interactions, to make use of Groucho mediated repression.

The Tup and Groucho proteins have long been regarded as functional equivalents in fungi and animals respectively. Their dichotomous phylogenetic distribution (Tup in fungi, Groucho in animals) and shared role in transcriptional repression has been suggestive of an orthologous relationship. Increased sequence sampling of eukaryotic species has extended the range of Tup-like genes beyond the fungi, including non-opisthokont species. In the phylogenetic analysis presented here, the Groucho group clearly does not arise from within

this Tup clade, but rather has a sister group relationship with it, suggesting an equally ancient history. Furthermore, the phylogenetic distribution of Groucho and Tup orthologs revealed two excavate species, *N. gruberi* and *T. pyriformis* that appear to encode Groucho-like proteins, with *Naegleria* also encoding a Tup protein. Taken together with the broad eukaryotic distribution of Tup, this presents a *prima facie* case that both Groucho-like and Tup-like proteins were present in the eukaryotic common ancestor, although clearly distinguishing between this and alternative scenarios of horizontal gene transfer or contamination (or mis-identified species) and phylogenetic reconstruction artefacts would be made easier by the availability of more non-parasitic eukaryotic genome sequences.

Detailed comparison of the 3D protein structures of Groucho and Tup, at the level of the conservation in Tup of the binding site residues of Groucho is further suggestive of a nonorthologous relationship between the two. In particular, two amino acid substitutions play a role in restructuring the binding site of Groucho. The need for multiple substitutions and the presumed biological requirement of functional continuity is more likely to have occurred in a duplicated gene copy. Analysis of yeast transcription factors and their conservation suggests some likely genes encoding candidate interaction motifs, but not, apparently, to the extent seen in Metazoa. Two possibilities suggest themselves: first, the ability to discriminate 'F' as the first motif residue enables a greater utility, in the sense that LxxLL is more likely to occur by chance in protein sequences, making it harder to discriminate between 'functional' and nonfunctional motifs; second, that the TPR repeat containing CYC8/ssn6 co-factor plays a crucial role in transcription factor recognition in yeasts, and that specificity is not encoded solely in the WD40 β -propeller domain. Interestingly, the TPR repeats of CYC8/ssn6 appear to be orthologous to the TPR repeats encoded in the human histone demethylase KDM6A/UTY genes (they are reciprocal blast best hits, data not shown), and these latter proteins have been shown to interact with TLE1 (Grbavec et al. 1999).

Among the eukaryotes, plants encode no WD40 β -propeller domains that are obviously orthologs of Groucho or Tup. The *Arabidopsis* protein TOPLESS is frequently described as being a plant equivalent of Groucho/Tup (Liu and Karmarkar 2008), but at the level of primary sequence, contains two WD40 β -propeller domains and distinct N-terminal domains. TOPLESS binds LxLxL motifs present in many plant transcription factor proteins. The recently solved 3D structure of the N-terminal domain, however, demonstrates that the interaction of the peptide motif is with this, rather than the WD40 domain as found in Groucho/Tup, suggesting it has arisen via an independent evolutionary path (Jennings and Ish-Horowitz 2008; Ke et al. 2015).

Groucho proteins also interact with the transcription factor TCF/LEF, the effector of WNT signalling, via an interaction of their N-terminal domains (Chodaparambil et al. 2014). The fact that unicellular eukaryotes encode orthologs of Groucho, but not TCF like transcription factors, suggests that interactions with groucho via EH1 and WPRW type motifs arose before those with TCF/LEF. This inference is consistent with the fact that WNT ligands are found only within the Metazoa.

Materials and Methods

Data Sources

The NR protein database was downloaded from the NCBI (20th September 2015) <ftp://ftp.ncbi.nih.gov/blast/db/FASTA/nr.gz>.

Proteins from the 'Origins of Multicellularity' project were downloaded from https://www.broadinstitute.org/annotation/genome/multicellularity_project/MultiHome.html.

Sequence reads from species referred to in Torruella et al. (2015) were downloaded from the EBI ENA database and assembled using Trinity with open reading frames being identified using Transdecoder (Grabherr et al. 2011).

Predicted proteins from *Schizosaccharomyces* were downloaded from <https://www.broadinstitute.org/scientific-community/science/projects/fungal-genome-initiative/schizosaccharomyces-genomes-project>.

Sponge proteins were downloaded from <http://compagen.org/datasets.html>.

Mnemiopsis leidyi protein models were taken from: <http://research.nhgri.nih.gov/mnemiopsis/download/download.cgi?dl=proteome>.

Trichoplax adhaerens protein models were taken from: ftp://ftp.ncbi.nih.gov/genomes/refseq/invertebrate/Trichoplax_adhaerens/.

Phylogenetic Analysis

Representative WD40 containing regions from Tup and Gro proteins were aligned using the MAFFT program (using the L-INS-i options) (Kato and Standley 2013). WD40 sequences

from WDR5 proteins from *Capsaspora owczarzaki*, *Amphimedon queenslandica*, *T. adhaerens*, *N. vectensis*, and human were added to serve as an outgroup. Ragged N and C-termini were trimmed, but the alignment was otherwise unedited. Analysis using the proTest3 software gave LG + G as the best fitting model (Darriba et al. 2011). Accordingly, phylogenetic analysis was performed using Phyml with the LG + G model (Le and Gascuel 2008), with other parameters left as defaults (Guindon et al. 2010). 100 bootstrap replicates were performed. The data were also analyzed with phylobayes, which uses a Bayesian rather than Maximum Likelihood approach (Lartillot et al. 2009), again using the LG + G model and using two chains. Chains were run for 35,000 generations. A consensus tree was produced using bpcomp from the phylobayes package, discarding the first 20,000 generations, giving a maxdiff of 0.1 and a meandiff of 0.003.

Ortholog Identification in the *Schizosaccharomyces* Group

In order to screen for similar motifs in a well-characterized fungal genome, I inferred orthologous groups in the *Schizosaccharomyces* group genomes (*Sc. pombe*, *Sc. japonicus*, *Sc. octosporus*, and *Sc. cryophilus*), via mutually consistent groups of four reciprocal best hits in all against all searches performed with the phmmer program from the hmmer package (<http://hmmer.org>). Instances of PFAM domains within *Sc. pombe* sequences were recorded using hmmssearch from the hmmer package, and only motif matches occurring to sequence regions outside these coordinates were assessed for conservation in the remaining three species.

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Supplementary Material

Supplementary material and figures S1 and S2 are available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

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