On the monophyly of chromalveolates using a six-protein phylogeny of eukaryotes

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A global phylogeny of major eukaryotic lineages is a significant and ongoing challenge to molecular phylogenetics. Currently, there are five hypothesized major lineages or 'supergroups' of eukaryotes. One of these, the chromalveolates, represents a large fraction of protist and algal diversity. The chromalveolate hypothesis was originally based on similarities between the photosynthetic organelles (plastids) found in many of its members and has been supported by analyses of plastid-related genes. However, since plastids can move between eukaryotic lineages, it is important to provide additional support from data generated from the nuclear-cytosolic host lineage. Genes coding for six different cytosolic proteins from a variety of chromalveolates (yielding 68 new gene sequences) have been characterized so that multiple gene analyses, including all six major lineages of chromalveolates, could be compared and concatenated with data representing all five hypothesized supergroups. Overall support for much of the phylogenies is decreased over previous analyses that concatenated fewer genes for fewer taxa. Nevertheless, four of the six chromalveolate lineages (apicomplexans, ciliates, dinoflagellates and heterokonts) consistently form a monophyletic assemblage, whereas the remaining two (cryptomonads and haptophytes) form a weakly supported group. Whereas these results are consistent with the monophyly of chromalveolates inferred from plastid data, testing this hypothesis is going to require a substantial increase in data from a wide variety of organisms.

INTRODUCTION

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A well-resolved global phylogeny of eukaryotes has been a major goal in understanding eukaryotic evolution and biodiversity because eukaryotic macroevolutionary events can only be interpreted accurately within the context of the relationships among these organisms. For example, plastids (the photosynthetic organelles of plants and algae) have spread between eukaryotic lineages via secondary endosymbiosis, but without a clear understanding of the relationships between plastid-bearing eukaryotes, it is very difficult to trace the history of plastid gains and losses accurately (Archibald & Keeling, 2002). Indeed, plastids illustrate this need especially well, since much of what is known about algal relationships is based on plastidencoded genes or genes for plastid-targeted proteins (Fast *et al.*, 2001; Harper & Keeling, 2003; Yoon *et al.*, 2002b), which can evolve independently of the host. This does not invalidate such inferences, but it is desirable to buttress these conclusions with supporting data from host lineages.

The first large-scale molecular phylogenies of eukaryotes, based on small-subunit rRNA (SSU rRNA), consisted of a basal ladder of protists followed by an explosive 'crown' of other eukaryotes - animals, fungi and plants (Cavalier-Smith & Chao, 1996; Kumar & Rzhetsky, 1996; Sogin & Silberman, 1998; Van de Peer & De Wachter, 1997). The first protein-coding gene phylogenies gave similar results (Brown & Doolittle, 1995; Hashimoto et al., 1994; Kamaishi et al., 1996), but also contained important incongruencies and over time, these contradictions have added up to serious questions about the 'shape' of the eukaryotic tree. Comparative phylogenetics based on several well-sampled protein-coding genes has shown that the overall laddercrown structure of the tree is not a true representation of eukaryotic evolution. Nearly all of the statistically wellsupported 'deep-branching' eukaryotes are now known to be artifacts of heterogeneous evolutionary rates or longbranch attraction (Embley & Hirt, 1998; Gribaldo & Philippe, 2002; Hirt et al., 1999; Morin, 2000; Philippe, 2000; Van de Peer et al., 2000b). All individual gene phylogenies seem to contain a few misleading branches and problematic taxa, but fortunately different trees seem

Abbreviations: EF-1 alpha, elongation factor-1 alpha; EST, expressed sequence tag; gDNA, genomic DNA; HSP, heat-shock protein; ML, maximum-likelihood; SSU rRNA, small-subunit rRNA.

The GenBank/EMBL/DDBJ accession numbers for the sequences reported in this paper are AY729814-AY729874 and AY739894-AY739895.

Individual ProML phylogenies (Figs A-E) are available as supplementary material in IJSEM Online.

to have different problems. Thus, reconstructing a global eukaryotic tree seems to remain within reach.

A new hypothesis is emerging based on several different kinds of mutually reinforcing data from many different genes. These data can take the following forms: a single phylogeny based on multiple concatenated genes (Bapteste et al., 2002; Yoon et al., 2002b); several individual phylogenies that all converge on the same relationship (Fast et al., 2002; Simpson et al., 2002b); non-phylogenetic molecular evidence such as insertions, deletions, gene duplications or gene fusions (Archibald et al., 2002; Baldauf & Palmer, 1993; Keeling & Palmer, 2001; Stechmann & Cavalier-Smith, 2002); and/or non-molecular data such as ultrastructure (Simpson et al., 2002a). The hypothesis emerging from this approach is that there are five major divisions or 'supergroups' of eukaryotes representing most or perhaps even all eukaryotic diversity (Keeling, 2004). Many of these do not have formal taxonomic names that are universally accepted, so for simplicity they can be informally referred to as the rhizaria, chromalveolates, excavates, unikonts and plants.

The chromalveolate supergroup represents a large fraction of known eukaryotic diversity, ranging from tiny microbial parasites to kelps as long as 50 m. In fact, chromalveolates account for about half of the recognized species of protists and algae (Cavalier-Smith, 2004). The chromalveolates comprise six subgroups, each of which is a large and diverse group in its own right. The alveolates contain the apicomplexans, ciliates and dinoflagellates, whereas the chromists are made up of the cryptomonads, haptophytes and heterokonts (also called stramenopiles). Various permutations including some, but not all, of these lineages have been put forward in the past (e.g. Chadefaud, 1950; Christensen, 1989), but these were largely restricted to photosynthetic members and included some groups (e.g. euglenids) that are now known to be very distantly related to other putative chromalveolates. Moreover, the current chromalveolate hypothesis is based on a very specific model of plastid evolution. Cryptomonads, haptophytes, heterokonts and dinoflagellates all have members that contain plastids derived from secondary endosymbiosis with a red alga, whereas apicomplexans contain a cryptic plastid of putative red algal origin (Blanchard & Hicks, 1999; Fast et al., 2001; Funes et al., 2002; Köhler et al., 1997; McFadden & Waller, 1997; Williamson et al., 1994; Zhang et al., 2000). Based on this, it was proposed that these plastids originated from a single endosymbiotic event in their common ancestor (Cavalier-Smith, 1999).

A single endosymbiotic event also leads to the prediction that both the host lineages and their plastids share a common evolutionary history. The only molecular evidence, in addition to other kinds of evidence, supporting this supergroup comes from plastid-targeted and plastidencoded proteins. In the first instance, plastid-targeted GAPDH has been shown to be the product of a rare gene duplication of the cytosolic homologue, which has replaced the ancestral, cyanobacterial form (Fast *et al.*, 2001; Harper

& Keeling, 2003). This replacement is a strong marker for a common origin of chromalveolate plastids. From the plastid genome itself, a phylogeny based on five concatenated genes supports the union of cryptomonads, haptophytes and heterokonts (Yoon et al., 2002a). Given that plastids can move between lineages, however, these conclusions will be significantly stronger if the host lineages can also be demonstrated to be related. Molecular data have confirmed the alveolates as a group (Fast et al., 2002; Van de Peer et al., 2000a; Van de Peer & De Wachter, 1997) and there are data from a four-gene concatenated set indicating that ciliates, apicomplexans and heterokonts form a group (Baldauf et al., 2000). However, no large-scale analysis of nuclear-cytoplasmic genes has included all major chromalveolate lineages and, moreover, none has included representation from the remaining four supergroups, which is desirable to make certain than none of these are positioned between members of the supergroup being tested.

Here, the monophyly of the chromalveolates has been examined within the context of a global phylogeny of eukaryotes using six nucleus-encoded genes for cytoplasmic proteins: actin, alpha-tubulin, beta-tubulin, elongation factor-1 alpha (EF-1 alpha), heat-shock protein 70 (HSP70) and heat-shock protein 90 (HSP90). Homologues of these genes were characterized from a variety of chromalveolates so that each gene was known from at least one representative of all six chromalveolate subgroups. In addition, genes from representatives of two other supergroups (rhizarians and excavates) have also been characterized since molecular data from these groups are sparse. Consequently, this is the first analysis that includes multiple gene data from all five supergroups.

METHODS

Strains and culture conditions. Unialgal axenic cultures of the cryptomonads *Guillardia theta* CCMP 327 and *Rhodomonas salina* CCMP 1319, the dinoflagellate *Heterocapsa triquetra* CCMP 449, the haptophytes *Isochrysis galbana* CCMP 1323, *Pavlova lutheri* CCMP 1325 and *Prymnesium parvum* CCMP 1926 and the heterokonts *Cyclotella cryptica* CCMP 333, *Mallomonas rasilis* CCMP 478 and *Phaeodactylum tricornutum* CCMP 1327 were obtained from the Provasoli-Guillard National Center for Culture of Marine Phytoplankton (McKown Point, West Boothbay Harbor, ME, USA) and grown in f/2-Si medium at 16 °C (12:12 light: dark cycle).

The heterokont *Spumella uniguttata* was the prey organism for a culture of the apicomplexan *Colpodella edax* isolated from a freshwater pond near Borok (Yaroslavskaja, Russian Federation). Genes were obtained from *Spumella uniguttata* in several attempts to sequence the *Colpodella edax* homologues (Kuvardina *et al.*, 2002).

Genomic DNAs (gDNAs) from the oomycete heterokonts *Apodachlya* brachynema CBS 557.69, Brevilegnia macrospora CBS 132.37, Phytophthora palmivora CBS 236.30, Plectospira myriandra CBS 523.87, Pythium graminicola CBS 327.62 and Thraustotheca clavata CBS 343.33 were kindly donated by A. W. DeCock (Centraalbureau voor Schimmelcultures, Utrecht, The Netherlands) and gDNA from the raphidophyte heterokont *Heterosigma akashiwo* was kindly donated by K. Ishida (Department of Biology, Kanazawa University, Kanazawa, Japan).

Species	Actin	alpha-Tubulin	beta-Tubulin	HSP90	EF-1 alpha	HSP70
Chromalveolates						
Apicomplexans						
Cryptosporidium parvum	+	+	+	+	+	+
Eimeria tenella	+	$+^{a_{\star}}$	+	+	$+^{b}$	$+^{a}$
Plasmodium falciparum	+	+	+	+	+	+
Ciliates						
Paramecium tetraurelia	+	+	+	+	+	_
Tetrahymena pyriformis	+	+	+	+	+	_
Tetrahymena thermophila	+	+	+	+	_	+
Dinoflagellates	1	I	I	I		1
Crypthecodinium cohnii		+ ^c	+	1	_	
	+			+	_ _+	+
Heterocapsa triquetra	+	+	+	+	-1	+
Oxyrrhis marina	+	+	+	+	_	-
Perkinsus marinus	+	+	+	+	_	_
Cryptomonads					4	
Guillardia theta	+	+	+	+	$+^{d}$	+
Haptophytes						
Isochrysis galbana	+	+	+	+	-†	-
Pavlova lutheri	_	+	+	+	_	-
Prymnesium parvum	+	+	+	+	_	-
Heterokonts						
Apodachlya brachynema	+	+	+	+	+	_
Brevilegnia macrospora	+	_	+	+	+	+
Heterosigma akashiwo	+	+	+	+	+	+
Mallomonas rasilis	+	+	+	+	+	_
Phaeodactylum tricornutum	+	+	+	+	+	_
Phytophthora palmivora	+	+	+	+	+	+
Plectospira myriandra	+	+	+	+	+	_
Pythium graminicola	+	+	+	+	_	_
Spumella uniguttata	+	I	+	+	+	+
Thalassiosira pseudonana		1			+	+
Thaussiosira pseudonana Thraustotheca clavata	+	+	+	+	Ŧ	Ŧ
Unikonts	+	+	+	+		
Animals						
Caenorhabditis elegans	+	+	+	+	+	+
Danio rerio	+	+	+	+	+	+
Drosophila melanogaster	+	+	+	+ "	+	+
Homo sapiens	+	+	+	+	+	+
Rattus norvegicus	+	+	+	+	+	+
Fungi						
Ajellomyces capsulatus	+	+	+	+	+	+
Candida albicans	+	+	+	+	+	+
Neurospora crassa	+	+	+	+	+	+
Saccharomyces cerevisiae	+	+	+	+	+	+
Schizosaccharomyces pombe	+	+	+	+	+	+
Mycetozoans						
, Dictyostelium discoideum	+	+	+	+	+	+
Rhizarians						
Chlorarachniophytes						
Bigelowiella natans	+	+	+	+	-+	+
Plants		1	,		I	1
Land plants						
Arabidopsis thaliana	1	1	1	1	1	. 1
	+	+	+	+	+	+
Oryza sativa	+	+	+	+	+	+

Table 1. Taxa and genes represented in analyses of the four-gene and six-gene concatenated datasets

Table 1. cont.

Species	Actin	alpha-Tubulin	beta-Tubulin	HSP90	EF-1 alpha	HSP70
Zea mays	+	+	+	+	+	+
Green algae						
Chlamydomonas reinhardtii	+	+	+	+	— †	+
Red algae						
Porphyra yezoensis	+	$+^{f}$	+ ^g	+	+ g	+
Excavates						
Diplomonads						
Giardia intestinalis	+	+	+	$+^{h}$	+	+
Parabasalia						
Trichomonas vaginalis	+	+	+	+	+	+
Euglenozoans						
Euglena gracilis	+	+	+	+	+	_
Leishmania donovani	$+^{i}$	+	$+^{j}$	+	$+^{l}$	$+^{i}$
Trypanosoma brucei	+	+	+	+	+	+
Trypanosoma cruzi	+	+	+	+	+	+
Heterolobosea						
Naegleria gruberi	+	+	+	+	$+^{m}$	_
Oxymonads						
Streblomastix strix	_	+	+	+	+	+

*Sequences obtained from species other than those listed are indicated as follows: *a*, *Eimeria acervulina*; *b*, *Eimeria bovis*; *c*, *Amphidinium herdmannii*; *d*, *Rhodomonas salina*; *e*, *Drosophila auraria*; *f*, *Prionitis lanceolata*; *g*, *Porphyra purpurea*; *h*, *Hexamita inflata*; *i*, *Leishmania major*; *j*, *Leishmania mexicana*; *l*, *Leishmania brasiliensis*; *m*, *Naegleria andersoni*.

†The sequence was determined, but was very divergent and was not used in phylogenetic analyses.

Algal cultures were harvested by centrifugation and cell pellets were lysed by grinding in a Knotes Duall 20 tissue homogenizer. gDNAs were extracted using the DNeasy Plant Mini kit (Qiagen) and total RNA was isolated using Trizol reagent (Invitrogen).

Characterization of new gene and cDNA sequences. PCR amplification was carried out using the following primers: actin, GGCCTGGAARCAYTTNCGRTGNAC and GAGAAGATGACNC-ARATH or TGGGAYGAYATGGARAARATHTGG; alpha-tubulin, CGCGGCCTCARGTNGGNAAYGCNTGYTGGGA and CGCGCC-ATNCCYTCNCCNACRTACCA; beta-tubulin, GCCTGCAGGNCA-RTGYGGNAAYCA and TCCTCGAGTRAAYTCCATYTCRTCCAT; EF-1 alpha, AACATCGTCGTGATHGGNCAYGTNGA and CTTG-ATCACNCCNACNGCNACNGT; HSP70, AAGATCATCGGNAT-HGAYYTN and CTGAACGATNCCRTTNGCATC; and HSP90, ACGTTYTAYWSNAAYAARGARAT and CGCCTTCATMATNCSY-TCCATRTTNGC. PCR products were gel-purified and then cloned into the TOPO-TA vector pCR2.1 (Invitrogen) and multiple clones of each were sequenced on both strands with ABI BigDye terminator chemistry (Applied Biosystems).

HSP90 and HSP70 cDNAs from the host and endosymbiont of the chlorarachniophyte *Bigelowiella natans* CCMP 621 were obtained from an ongoing EST (expressed sequence tag) project and each was fully sequenced. HSP70 cDNA was identified in ongoing EST projects from *Isochrysis galbana* and was fully sequenced. HSP70 from the oxymonad *Streblomastix strix* was amplified using primers AAGATCATCG-GNATHGAYYTNGGNACNAC and CTGAACGATNCRTTNGCRT-CDATRTCRAA from DNA isolated from enriched hindgut material of the Pacific dampwood termite (*Zootermopsis angusticolis*) as described previously (Keeling & Leander, 2003).

Molecular phylogeny. Newly determined sequences were added to

existing amino acid alignments. A total of eight alignments was subjected to phylogenetic analyses. Six of these were the respective alignments for the six protein-coding genes. The two remaining alignments were concatenations of different genes. One was a concatenation of actin, alpha-tubulin, beta-tubulin and HSP90 (AABH), whereas the second was a concatenation of all six genes (AABHHE). The resulting dimensions (number of taxa × number of amino acid positions) for all eight alignments were as follows: actin, 78 × 245; alpha-tubulin, 71 × 391; beta-tubulin, 73 × 380; EF-1 alpha, 48 × 422; HSP70, 45 × 505; HSP90, 61 × 516; AABH, 50 × 1554; and AABHHE, 50 × 2481. The amino acid alignments are available from the authors on request.

Distance analyses were performed on the individual datasets and the four-gene concatenated dataset only (AABH). Parsimony and maximum-likelihood (ML) analyses were performed on all datasets. For the six individual datasets and the four-gene concatenated dataset, ML distances were calculated using TREE-PUZZLE 5.0 (Strimmer & von Haeseler, 1996), using the WAG substitution matrix with the frequency of amino acid usage calculated from the data. Rate-acrosssite variation was modelled on a discrete gamma distribution with eight rate categories, estimating invariable sites and the shape parameter alpha from the data. Distance trees were constructed with weighted neighbour-joining using WEIGHBOR 1.0.1a (Bruno et al., 2000) and Fitch-Margoliash using FITCH 3.6a (Felsenstein, 1993). Fitch-Margoliash trees were inferred using the global rearrangements option and 10 input-order jumbles. Weighted neighbour-joining and Fitch-Margoliash bootstrap trees were constructed (without global rearrangements and implementing two input-order jumbles in Fitch-Margoliash) from 100 resampled datasets with gamma-corrected distances (with the rate category parameters above) using PUZZLEBOOT 1.0.3 (by M. Holder and A. Roger; http://www.tree-puzzle.de).

Protein ML analyses were performed using ProML 3.6a (Felsenstein, 1993) and PhyML (Guindon & Gascuel, 2003). ProML was performed with amino acid frequencies estimated from the data and rate-across-site variation was modelled on a gamma curve using the -r option with eight variable rate categories and an invariable sites category (rates and frequencies estimated by TREE-PUZZLE). Trees were searched with global rearrangements and a randomized input order of sequences with two jumbles. ProML bootstrap trees were constructed as above, but with no site-to-site rate variation and a single randomized input order. PhyML was performed using an input tree generated by BIONJ, the JTT model of amino acid substitution, the proportion of variable rates estimated from the data and nine categories of substitution rates (eight variable and one variable; parameters estimated by TREE-PUZZLE). PhyML bootstrap trees were constructed using the same parameters as the individual ML trees.

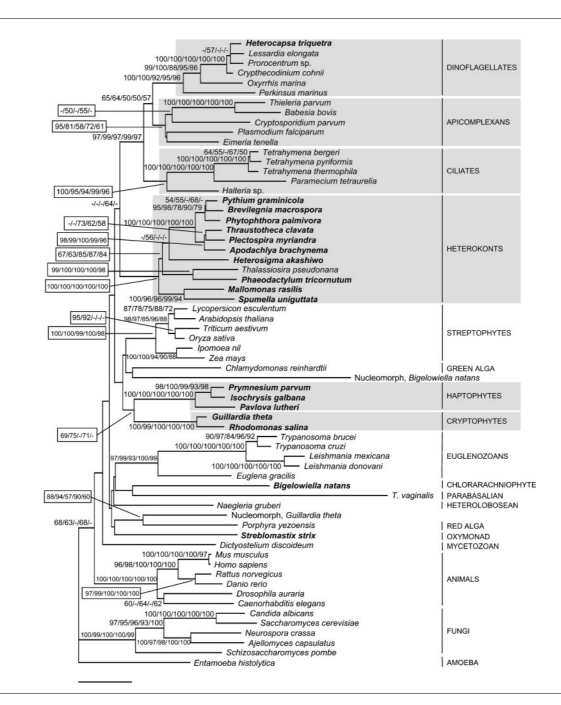


Fig. 1. Protein maximum-likelihood phylogeny (ProML) of HSP90. Bootstrap values are shown for nodes that received support over 50 % and are (left to right) weighted neighbour-joining, Fitch-Margoliash, ProML, PhyML and parsimony (dashes represent support lower than 50 %). Chromalveolate taxa are highlighted in shaded boxes and major groups are bracketed and labelled to the right. Newly determined sequence data are represented by taxon names in bold. Bar, 0.1 changes per site. Abbreviation: *T. vaginalis, Trichomonas vaginalis.*

Parsimony analysis was completed using PAUP 4.0b10 (50 random sequence additions; gaps treated as missing data) using a heuristic search with steepest descent and tree bisection-reconnection branch swapping (Swofford, 2002). Parsimony bootstraps were completed (100 replicates) using 10 random sequence additions.

Distance analyses were not performed on the six-gene concatenated dataset due to missing data for a number of taxa (see Table 1). Parsimony, ML and the respective bootstrap analyses were performed on this dataset as described above for the other gene analyses.

RESULTS AND DISCUSSION

Six genes representing five supergroups

Comparable datasets for six genes (actin, alpha-tubulin, beta-tubulin, EF-1 alpha, HSP90 and HSP70) were developed by sequencing 68 new genes or cDNAs from a variety of chromalveolates (primarily the most poorly sampled dinoflagellates, cryptomonads, haptophytes and heterokonts) and other eukaryotes (Table 1). A table of GenBank/ EMBL/DDBJ accession numbers for all taxa included in our various phylogenetic analyses is available as supplementary material in IJSEM Online. The overall aim was for all six major subgroups of chromalveolate and all five hypothesized supergroups to be represented in all six datasets and for representation to be sufficiently similar to allow genes to be concatenated. In addition, missing genes or cDNAs were also characterized from the excavate (oxymonad) Streblomastix strix and the rhizarian (the chlorarachniophyte cercozoan) Bigelowiella natans. No multiple gene phylogeny of eukaryotes has yet included a rhizarian (e.g. a cercozoan), which is an important omission when testing the monophyly of any one of the other supergroups.

Individual gene phylogenies

Of the six genes analysed here, the phylogeny of HSP90 (Fig. 1) has been investigated the least, but has shown promise. The phylogeny of alveolates has recently been examined with an expanded HSP90 dataset (Leander & Keeling, 2004). Moderate support was found for grouping alveolates and heterokonts, a result that has received varying levels of support from other molecular investigations (Baldauf et al., 2000; Ben Ali et al., 2001; Dacks et al., 2002; Stechmann & Cavalier-Smith, 2003; Van de Peer & De Wachter, 1997). Sequence data for cryptomonads, haptophytes or a diverse sample of heterokonts, however, has been lacking from these analyses. With the addition of these data, our HSP90 phylogeny shows a weak relationship between the strongly supported alveolate and heterokont clades. Interestingly, the strongly supported cryptomonad and haptophyte clades branch together with intermediate to weak support. The positions of both these larger groups, however, remain equivocal.

Phylogenies of the tubulins, actin, EF-1 alpha and HSP70 have all been analysed more extensively than HSP90 (e.g. Baldauf & Palmer, 1993; Baldauf *et al.*, 2000; Bhattacharya & Weber, 1997; Hashimoto *et al.*, 1995; Keeling, 2001;

Table 2. Summary of support for single- and multiple-gene analyses

beta-	HT,	
a-tubulin,	eterokonts;	
in, alph	HK, h	
of actir	hytes;	
tion c	aptop	
ıcatena	gae; HA, h	
I, Coi	n algae;	
AABF	green	
ctively.	gi; G,	
respec	F, fun	
hyML,	ozoa;	
and P	, euglen	
roML	ds; E,	
ash, P	omona	
1 argoli	, diple	
itch-N	ytes; D	
iing, F	ptoph	vailable
ur-joir	CR, cryp	Not av
eighbo	mists; C	S. NA,
hted n	CH, chron	ophyte
r weig	; CH,	strept
are fo	unimals	and S,
/alues	AN, a	algae;
strap v	olates;	R, red
, boot	L, alve	asalia;
Except where indicated, bootstrap values are for weighted n	and HSP90; AL, alveolates; AN, animals; CH,	P, parabasalia; R, red algae; and S, streptoph
ere in	d HSF	••
pt wh	tubulin an	rolobosea
Exce	tubu	hetei

Gene	CR	НА	НК	CH	AL	AL-HK	AL-CH	S-G	AN-F	E-HT	D-P
06dSH	100/99/100/100	00/100/100/100/001/00/100/100/100/100/00/	100/100/100/100	-/-/-/-	64/26/66/26	-/-/-/64	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-	-/-/-/-
Actın alpha-Tubulin	96/90/81/74 81/60/95/81	98/92/100/100 98/92/100/100	/0/4/0/ -/-/-/-	-/-/-/-	-/-/-/-	-/-/~C0/-	-/-/-/-	-/-/-/-	-/-/-/- 93/60/84/72	-/-/-/- -/-/-	-/-/-/-
beta-Tubulin	-/-/-	96/94/91/94	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-	-/-/-/-	85/93/83/93	-/-/-/-	99/92/69/62‡	72/81/73/57
EF-1 alpha	$NA\ddagger$	NA§	-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-	73/82/92/80	-/-/-/-	90/91/67/66
HSP70	NA‡	NA‡	53/77/74/99	-/-/-/-	-/-/-/-	-/-/-/-	-/-/-/-	88/83/65/86	-/-/-/-	-/-/-/-	-/-/-
AABH	$NA\ddagger$	100/100/100/100	99/100/100/100	-/-/-/-	98/97/98/100	66/53/-/59	-/-/-/-	100/100/100/100	-/-/-78	85/56/61/83	100/98/97/86
AABHHEII	NA‡	100/99	100/99	-/-	80/93	66/84	-/-	66/66	70/92	-/65	98/96
*Support for grou †Support for grou	uping apicomplexa uping <i>Euglena gra</i>	*Support for grouping apicomplexans, dinoflagellates and heterokonts †Support for grouping <i>Euglena gracilis</i> and <i>Naegleria gruberi</i> .	nd heterokonts. <i>ruberi</i> .								
+OILLY OLLE SEGUES	ince represented un	IS LANUL TOL TILLS BEL	IC.								

5The haptophyte sequence was unusually divergent and not included in this dataset (Keeling & Inagaki, 2004).

dataset

not completed for the six-gene

IIDistance analyses were

Keeling & Doolittle, 1996; Leander & Keeling, 2004; Saldarriaga et al., 2003) and the general characteristics of our phylogenies are similar to those given in these published reports. Bootstrap supports from these analyses are summarized in Table 2 [phylogenies (Figs A-E) are available as supplementary material in IJSEM Online]. Overall, the monophyly of each of the chromist lineages was recovered and strongly supported. As a group, the chromists were not recovered by any of the analyses of the six genes. Alveolates were resolved as monophyletic in actin, with the previously noted exception of ciliates (Keeling, 2001), and HSP70 phylogenies, but not with any appreciable bootstrap support. Not surprisingly, a chromalveolate clade was never recovered, but topologically, alveolates were positioned as a sister clade to the heterokonts in analyses of actin, EF-1 alpha, HSP70 and HSP90. For comparison, the support for four other more widely accepted groupings is also shown in Table 2 (green algae+streptophytes, animals+fungi, euglenozoans+heteroloboseans and diplomonads+parabasalia). Most of these groups are not overwhelmingly supported by individual phylogenies either and instead are supported erratically by different genes.

Six-gene and four-gene concatenated phylogenies

Few supported features of any of the individual phylogenies contradicted results from another gene and four of these genes have been concatenated in previous studies (Baldauf et al., 2000). Accordingly, datasets of four- and six-gene concatenations were analysed to compare with the results of the individual phylogenies (see Table 1 for a complete list of taxa used in these concatenations). A four-gene concatenation consisting of actin, alpha-tubulin, beta-tubulin and HSP90 (AABH) was possible for 50 taxa representing the six subgroups of chromalveolates and the five eukaryotic supergroups. In many respects, the phylogeny inferred from these data (Fig. 2) is similar to that recovered for HSP90. There were, however, some notable differences (e.g. the monophyly of the discicristates: euglenozoans+heteroloboseans) and bootstrap support for most nodes was significantly higher (Table 2). Support for the various groupings of chromalveolate taxa was also similar to the results of HSP90 analyses. Alveolates and heterokonts were grouped with moderate support, whereas a cryptomonad

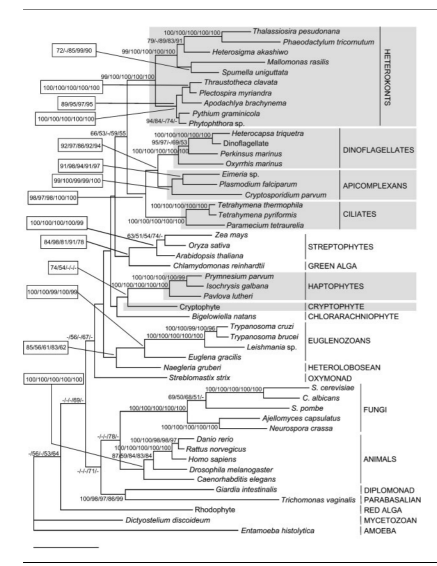


Fig. 2. Protein maximum-likelihood phylogeny (ProML) based on a concatenation of sequences for four nuclear-cytosolic proteincoding genes: actin, alpha-tubulin, betatubulin and HSP90. Bootstrap values and labelling formats are as for Fig. 1. The terms 'Cryptomonad', 'Dinoflagellate' and 'Rhodophyte' refer to concatenations of genes from more than one species representing these taxonomic groups (see Table 1). Bar, 0-1 changes per site. Abbreviations: *C. albicans, Candida albicans; S. cerevisiae, Saccharomyces cerevisiae; S. pombe, Schizosaccharomyces pombe.* and haptophyte clade received weak support, a topology also recovered by Stechmann & Cavalier-Smith (2003). The individual chromalveolate groups all received strong support, with the exception of the cryptomonads, which were represented by a single set of concatenated genes.

Adding data for EF-1 alpha and HSP70 to the concatenated dataset for analysis yielded a tree (Fig. 3) with topology that was essentially identical to that recovered by analysis of the four-gene dataset (Fig. 2). Bootstrap support was also virtually identical to the four-gene tree with the following qualitative differences: a slight decrease in support for monophyly of the alveolates, a slight increase in support for an alveolate/heterokont clade and increased support for monophyly of the opisthokonts (animals + fungi).

Conclusions: chromalveolates within a global eukaryotic phylogeny

The impetus behind the chromalveolate theory was to limit the number of endosymbiotic events in plastid evolution by proposing that the chromalveolate plastids all share a common origin (Cavalier-Smith, 1999). Indeed, molecular investigations based on plastid-encoded genes and plastidtargeted proteins have provided overlapping results that strongly support this hypothesis (Fast et al., 2001; Harper & Keeling, 2003; Yoon et al., 2002b). There is now growing support from individual and multiple gene phylogenies based on nucleus-encoded cytosolic proteins for the monophyly of an alveolate/heterokont clade. Moreover, the weak relationship observed between cryptomonads and haptophytes (never seen in nuclear gene trees before) is also consistent with the chromalveolate hypothesis. Considering these data within the context of the original cellular characteristics that prompted the recognition of the chromalyeolates suggests that these groups are all related and that, by extension, their plastids probably originated via a single secondary endosymbiotic event. Though there is presently no evidence from the phylogeny or cell biology of chromalveolates to suggest that they are not a monophyletic assemblage, it is also clear that more evidence from the nuclear-host lineage is necessary.

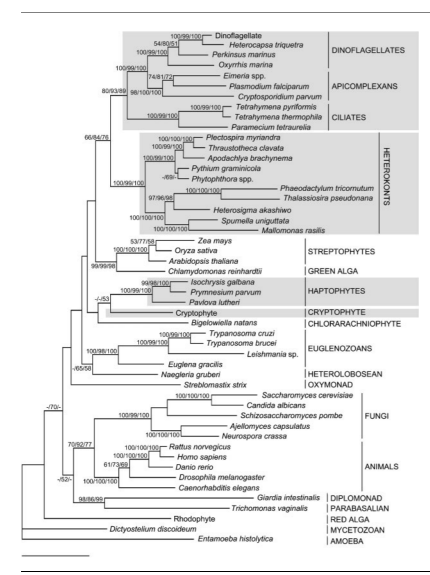


Fig. 3. Protein maximum-likelihood phylogeny (ProML) based on a concatenation of sequences for six nuclear-cytosolic proteincoding genes: actin, alpha-tubulin, betatubulin, EF-1 alpha, HSP70 and HSP90. Bootstrap values are shown for nodes that received support over 50% and are (left to right) ProML, PhyML and parsimony (dashes represent support lower than 50%). Labelling formats are as for Fig. 1. The terms 'Cryptomonad', 'Dinoflagellate' and 'Rhodophyte' refer to concatenations of genes from more than one species representing these taxonomic groups (see Table 1). Bar, 0.1 changes per site.

In general, uniting distant relatives (for, if the chromalveolates are a monophyletic group, they are an ancient group and distantly related) is a difficult problem that requires considerably more data than are currently available. A global phylogeny of eukaryotes based on nuclear-cytosolic proteins has been examined with four genes and 61 taxa (Baldauf et al., 2000), providing reasonable resolution. However, adding more taxa (six lineages that had not been previously represented were analysed here - cercozoans, cryptomonads, dinoflagellates, oxymonads, Entamoeba and haptophytes) has undermined overall support for the tree considerably. Other multiple gene analyses of eukaryotic phylogeny (e.g. Bapteste et al., 2002) have used many more genes (100) but fewer taxa (30) and were addressing important, but relatively narrow questions. A global phylogeny representing even an approximation of eukaryotic diversity remains to be achieved and will likely require a mixture of these approaches when more taxa have been sampled sufficiently to include a number of additional genes for analysis.

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