A Broad Phylogenetic Survey Unveils the Diversity and Evolution of Telomeres in Eukaryotes

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Abstract

Telomeres, ubiquitous and essential structures of eukaryotic chromosomes, are known to come in a variety of forms, but knowledge about their actual diversity and evolution across the whole phylogenetic breadth of the eukaryotic life remains fragmentary. To fill this gap, we employed a complex experimental approach to probe telomeric minisatellites in various phylogenetically diverse groups of algae. Our most remarkable results include the following findings: 1) algae of the streptophyte class Klebsormidiophyceae possess the Chlamydomonas-type telomeric repeat (TTTTAGGG) or, in at least one species, a novel TTTTAGG repeat, indicating an evolutionary transition from the Arabidopsis-type repeat (TTTAGGG) ancestral for Chloroplastida; 2) the Arabidopsis-type repeat is also present in telomeres of Xanthophyceae, in contrast to the presence of the human-type repeat (TTAGGG) in other ochrophytes studied, and of the photosynthetic alveolate Chromera velia, consistent with its phylogenetic position close to apicomplexans and dinoflagellates; 3) glaucophytes and haptophytes exhibit the human-type repeat in their telomeres; and 4) ulvophytes and rhodophytes have unusual telomere structures recalcitrant to standard analysis. To obtain additional details on the distribution of different telomere types in eukaryotes, we performed in silico analyses of genomic data from major eukaryotic lineages, utilizing also genome assemblies from our on-going genome projects for representatives of three hitherto unsampled lineages (jakobids, malawimonads, and goniomonads). These analyses confirm the human-type repeat as the most common and possibly ancestral in eukaryotes, but alternative motifs replaced it along the phylogeny of diverse eukaryotic lineages, some of them several times independently.

Key words: algae, telomerase activity, Excavata, comparative genomics, Goniomonas.

Introduction

The termini of linear eukaryotic chromosomes are protected by telomeres. Their DNA part is formed typically by a long array of conserved minisatellite sequences that tend to be conserved in particular groups of organisms, for example, TTAGGG in vertebrates and fungi (named here as the human-type; Meyne et al. 1989), TTTAGGG in most plants (Arabidopsis-type; Richards and Ausubel 1988), or TTAGGG in insects (Okazaki et al. 1993; Frydrychova et al. 2004; Vítková et al. 2005). These telomeres are maintained by a special reverse transcriptase, telomerase, which elongates telomeres by addition of telomeric repeats and thus solves the so-called end replication problem (reviewed in Chan and Blackburn 2004). However, many exceptions to these rules are known, for example, telomeres maintained by retrotransposons in Drosophila melanogaster (Biessmann and Mason 2003).
diverse telomeric minisatellite sequences in yeasts (Teixeira and Gilson 2005), plants with the human-type or unknown telomeric sequences (Sýkorová et al. 2003a, 2003b, 2006), or novel telomeric sequence in Arthropoda (Vitkova et al. 2005; Mravinac et al. 2011). An interesting diversity of telomeres was recently described also in the green algal group Chlamydomonadales, where at least two evolutionary transitions from the ancestral TTTAGGG type to the TTTTAGGG (Chlamydomonas-type) occurred in the clade Chloromonadia and independently in a subclade of the Reinhardtinia clade; moreover, the human-type telomeric sequence was found in some green algal species that fall within the Dunallielia and Stephanosphaeria clades (Fulnečková et al. 2012).

Eukaryotic algae are a polyphyletic assemblage of phylogenetically diverse organisms with different life styles and strategies and are thus of interest for telomere biology, because they represent a substantial portion of the euchromatic phylogenetic diversity. Three algal lineages Glaucothyra, Rhodophyta, and Chloroplastida (the latter including also land plants) represent direct descendants of an ancestral alga with a cyanobacterium-derived plastid and are thought to form a monophyletic “supergroup” called Archaeoplastida, whereas other algal groups obtained their plastids from red or green algae through a process called secondary or tertiary endosymbiosis (Archibald 2009). Different telomere types were described in chromosomes of the nucleus and the nucleomorph (vestigial endosymbiont-derived nucleus) of cryptophytes or chlorarachniophytes, with the nucleomorph telomeres presumably descending from original telomeres of the ancestral-engulfed algal endosymbiont (Gilson and McFadden 1995; Zauner et al. 2000). Previous genome sequencing projects reported human-type telomeric sequences in the diatoms Thalassiosira pseudonana and Phaeodactylym tricornutum (Armburst et al. 2004; Bowler et al. 2008) and an unusual AATG6 sequence in the red alga Cyanidioschyzon merolae (Nozaki et al. 2007). Synthesis of Arabidopsis-type telomeric repeats by telomerase was observed in dinoflagellates (Fojtová et al. 2010; Zielke and Bodnar 2010), illustrating the diversity of telomeric sequences in Alveolata, where ciliates possess telomeres with TTGGGG or TTTTGGGG repeats (Prescott 1994) and apicomplexan taxa display several related telomere types (see Kissing and DeBarry 2011 for review).

Excavata is a potentially monophyletic “supergroup” of unicellular eukaryotes that ancestrally share a characteristic ventral feeding groove and an associated specifically organized microtubular cytoskeleton (Simpson 2003; Hampel et al. 2009). Excavates may occupy a key position in the eukaryotic phylogeny and include many important or biologically interesting species, yet they remain among the most poorly explored eukaryotic supergroups with regard to their molecular genetic and genomic features. This holds true also with regard to telomere biology, as telomeres have been characterized only in very few excavates, including the parasitic trypanosomatids (Van der Ploeg et al. 1984; Lira et al. 2007) and the diplomonad Giardia intestinalis (=G. lamblia; Le Blancq et al. 1991). Genome sequences have been additionally reported for the parabasalid Trichomonas vaginalis (Carlton et al. 2007) and the heterolobosean Naegleria gruberi (Fritz-Laylin et al. 2010), but no information is available on their telomeres. For other deep excavate lineages, such as Jakobida, Prexostyla, or Malawimonadida, no knowledge about telomeres and no representative genome sequences are available.

Here, we tested the activity of telomerase and investigated the presence of minisatellite repeats in diverse algal lineages using experimental approaches. In addition, our on-going genome sequencing projects enabled us to obtain the first data about telomeres in jakobids, malawimonadids, and gonion monads (the latter group representing the closest heterotrophic relatives of the cryptophyte algae). Finally, we surveyed diverse eukaryotic genome sequences available in public databases and inferred their telomeric sequences by in silico analyses. This combination of experimental and bioinformatic analyses allowed us to describe the diversity of telomeres across the eukaryotic phylogeny (fig. 1), to confirm a predominant occurrence of the human-type telomeric sequence in basal lineages, and to demonstrate independent acquisition of the same telomeric repeats in various phylogenetic lineages.

Materials and Methods

Algal Cultures and DNA Extraction

The algal material used in this study originated from culture collections as specified in supplementary table S1, Supplementary Material online. Algae were grown in the recommended liquid media BBM or MASM (www.ccap.ac.uk/media/pdrcipies.htm; last accessed February 21, 2013), or on nutrient agar plates. Chromera velia was grown in a modified f/2 medium (www.ccap.ac.uk/media/pdrcipies.htm; last accessed February 21, 2013) in which natural seawater was replaced by seawater prepared by dissolving 23.38 g of “Red Sea” salt (Red Sea, USA) in 1 l of distilled water. Phaeodactylum tricornutum was cultivated in MASM medium supplemented with 30 mg of Na2SiO3·9H2O per liter.

The absence/presence of eukaryotic contaminants was monitored microscopically and using algal cultures grown on BBM and bacterial LB agar plates. The identity of most of algal samples was verified by polymerase chain reaction (PCR) amplification (discussed later) and sequencing of the internal transcribed spacer and/or small subunit (SSU) ribosomal DNA (rDNA) regions, and when necessary (in some green algae), phylogenetic analyses were conducted to confirm the assignment into specific algal classes (data not shown). Genomic DNA for PCR amplification was isolated using the “modified IRRI” method (Collard et al. 2007). Genomic DNA from a control alga Chlorella vulgaris (TEL01, supplementary table S1, Supplementary Material online) was isolated...
according to protocol described by Saghai-Maroof et al. (1984). Isolation of DNA from other algal samples was performed according to a previously described protocol (Fulněcková et al. 2012), which involves the use of proteinase K during the lysis step. The DNA sample concentrations were estimated from agarose gels.

**Dot-Blot Hybridization**

Genomic DNA samples (~1 µg per sample) were dot-blotted onto Amersham Hybond-XL nylon membrane (GE Healthcare) and hybridized with radioactively end-labelled oligonucleotide probes (ATSB, CHSB, HUSB, TTCAGGG-SB, TTTAGGG-SB, T4AG2-SB, T3G3-SB, T2CG3-SB, Red alga-SB, supplementary table S5, Supplementary Material online).
endonucleases a s e f o r 1 5 a n d 4 5 m i n ( o r 6 0 m i n ) , a n d t h e n b y t h e r e s t r i c t i o n to Neplechova et al. (2005). Briefly, membranes were hybridized at 55 °C for 16 h and washed at 55 °C under low stringency conditions (2x saline sodium citrate [SSC] and 0.1x sodium dodecylsulphate [SDS]); the final wash for the ATSB and HUSB oligonucleotides was done using a high-stringency washing buffer (0.6x SSC and 0.1x SDS) to avoid cross-hybridization. Membranes for rehybridization with another probe were gently washed three times in 0.5% SDS at 80 °C. A control probe of mixed SSU and large subunit (LSU) rDNA fragments was prepared by mixing an equal amount of PCR products from several phylogenetically diverse algae (TEL213 Rhodella maculata, TEL97 Klebsormidium subtilissimum, TEL211 Tetraselminus chui, TEL207 Euglena geniculata, TEL01 Chl. vulgaris) obtained using amplification by a combination of gene-specific primers [18S rDNA-F, 18S rDNA-R (18S rDNA-F, 18S rDNA-R (Katana et al. 2001) for TEL207, TEL211, TEL97, and TEL01; p4 and p23 [Van der Auwera et al. 1994] for TEL97 and TEL211; and ITS-A [Blattner 1999] for TEL207, TEL211, TEL97, and TEL01). The mixture was labelled by DecaLabel DNA Labeling Kit (Fermentas, Thermo Scientific). The probe was used for final rehybridization overnight at 62 °C and low stringency conditions (2x SSC, 0.1x SDS) or at 65 °C and high stringency conditions (0.2x SSC, 0.1% SDS). Membranes were exposed to autoradiography screens and signals were visualized using a phosphoimager FLA5000 (FujiFilm) and evaluated by the Multigauge software (FujiFilm).

Restriction Digestion, Pulsed-Field Gel Electrophoresis, and Southern Hybridization

Genomic DNA samples (1–5 μg) were digested with restriction endonucleases Rsal, Alul, or TaqI (NEB) and run on an 0.9% agarose gel in Tris–acetate–EDTA (TAE) buffer; the DNA fragments were alkali blotted and hybridized using the same hybridization and washing conditions as for dot-blots (for exceptions see supplementary material, Supplementary Material online). Agarose plugs with high-molecular-mass DNA samples for pulsed-field electrophoresis were prepared from lyophilized algal samples, and BAL-31 and restriction enzyme digestion was performed as described in Šýkorová et al. (2005). Briefly, agarose plugs with high-molecular-weight DNA (TEL206 E. stellata, TEL207 E. geniculata, TEL133 Eustigmatus polyphem, TEL201 Vischeria punctata, TEL103 K. nitens, TEL97 K. subtilissimum, and TEL131 Porphyridium purpurreum) were digested with BAL31 nuclease for 15 and 45 min (or 60 min), and then by the restriction endonucleases SmaI (TEL97 and TEL131) or HindIII (TEL206, 207, 133, 201, 103) (all enzymes from NEB). The DNA was then analyzed by pulsed-field gel electrophoresis using a CHEF Mapper (BioRad) under the following conditions: 1% agarose (BioRad) gel in 0.5x TBE buffer, 6 V/cm, pulses 0.5–35 s for 20 h at 13 °C. Gels were alkali blotted and hybridized with the telomere probes.

Telomere Repeat Amplification Protocol Assay

Telomerase activity was investigated using a protocol originally developed for plant telomerases (Fitzgerald et al. 1996; Šýkorová et al. 2003b) and applied with modifications to dinoflagellates (Fojtová et al. 2010) and green algae (Fulněcková et al. 2012) (supplementary fig. S2A, Supplementary Material online).

**Table S2, Supplementary Material** online as described in Šýkorová et al. (2003b) with minor modifications according to Neplechová et al. (2005). Briefly, membranes were hybridized at 55 °C for 16 h and washed at 55 °C under low stringency conditions (2x saline sodium citrate [SSC] and 0.1x sodium dodecylsulphate [SDS]); the final wash for the ATSB and HUSB oligonucleotides was done using a high-stringency washing buffer (0.6x SSC and 0.1x SDS) to avoid cross-hybridization. Membranes for rehybridization with another probe were gently washed three times in 0.5% SDS at 80 °C. A control probe of mixed SSU and large subunit (LSU) rDNA fragments was prepared by mixing an equal amount of PCR products from several phylogenetically diverse algae (TEL213 Rhodella maculata, TEL97 Klebsormidium subtilissimum, TEL211 Tetraselminus chui, TEL207 Euglena geniculata, TEL01 Chl. vulgaris) obtained using amplification by a combination of gene-specific primers (18S rDNA-F and 18S rDNA-R (Katana et al. 2001) for TEL207, TEL211, TEL97, and TEL01; p4 and p23 [Van der Auwera et al. 1994] for TEL97 and TEL211; and ITS-A [Blattner 1999] for TEL207, TEL211, TEL97, and TEL01). The mixture was labelled by DecaLabel DNA Labeling Kit (Fermentas, Thermo Scientific). The probe was used for final rehybridization overnight at 62 °C and low stringency conditions (2x SSC, 0.1x SDS) or at 65 °C and high stringency conditions (0.2x SSC, 0.1% SDS). Membranes were exposed to autoradiography screens and signals were visualized using a phosphoimager FLA5000 (FujiFilm) and evaluated by the Multigauge software (FujiFilm).

Gathering Genomic Data for Jakobids, Malawimonads, and Goniomonads

Complemented draft genome sequences and their systematic analyses of the jakobid Andalucia godoyi, the malawimonad Malawimonas californiana, and the goniomonad Goniomonas avonlea will be published elsewhere together with details on DNA isolation, sequencing, and assembly protocols. Briefly, *And. godoyi* (ATCC PRA-185) and *M. californiana* (ATCC 50740) were sequenced by GS FLX Titanium platform (454 Life Sciences/Roche) employing both shotgun and pair-end libraries. Draft assemblies were generated using Newbler 2.6. Genome sequence data of the recently described species *G. avonlea* (Kim and Archibald 2013) were generated using
the Illumina sequencing platform from multiple libraries, including two standard short insert libraries (300 bp) and mate pair libraries (2 and 6 kbp) (Beijing Genomics Institute; McGill University and Genome Quebec Innovation Centre). Errors in the raw illumina data were corrected using ALLPATHS-LG (Gnerre et al. 2011), and the corrected reads were assembled using the ABYSS de novo assembler (Simpson et al. 2009).

Bioinformatic Analyses of Telomeric Sequences
Candidate telomeric sequences were searched in our genome assemblies (discussed earlier) and in sequenced genomes available in various databases, including GenBank (www.ncbi.nlm.nih.gov; last accessed February 21, 2013), TraceArchive and Sequence Read Archive (SRA; http://www.ncbi.nlm.nih.gov/Traces/home; last accessed February 21, 2013), Joint Genome Institute (www.jgi.doe.gov; last accessed February 21, 2013), Broad Institute (http://www.broad.mit.edu/; last accessed February 21, 2013), University of Tokyo (http://merolae.biol.s.u-tokyo.ac.jp/; last accessed February 21, 2013), and EMBL (http://ct.bork.embl.de; last accessed February 21, 2013). Data sets of whole-genome sequence assemblies were downloaded in the FASTA format and searched using the BioEdit software and a string search for TTAGGG, TTTAGGG, TTTGGGG, and TTAGG types of a telomere-like repetitive minisatellites. The position and distribution of candidate telomeric sequences (terminal and/or internal) in the genome assembly was subsequently assessed by an automatic search for the respective strings. In addition, unassembled genomic reads from Phaeocystis antarctica and Porphyra umbilicalis (Sanger reads in the TraceArchive and Illumina data in the SRA archive, respectively) were searched using as a query a trimer of candidate sequences or one repeat of published telomere sequence of Cya. merolae (Nozaki et al. 2007).

Results
Sample Collection and Analyses of Algal Telomeres
To cover a phylogenetically wide sample of algae, we cultivated 48 algal strains from culture collections (supplementary table S1, Supplementary Material online), including members of Chloroplastida (Chlorophyta and Streptophyta), Rhodophyta, Glaucophyta, Haptophyta, Alveolata, Ochrophyta (Bacillariophyceae, Xanthophyceae, and Eustigmatophyceae), and Euglenozoa (supplementary table S1, Supplementary Material online). We examined all algal strains for telomerase activity by the TRAP assay (supplementary fig. S2A, Supplementary Material online) and cloned the TRAP products from 31 strains to determine what DNA sequence forms the ends of chromosomes (i.e., what sequence is synthesized by telomerase) (table 1 and figs. 2 and 3; supplementary fig. S2, Supplementary Material online). Algal strains used in this study came both from groups where the telomeric sequence could be presumed from published data and from groups where the telomeric sequence has not been described yet. In the latter cases, a set of alternative reverse primer sequences were used in combination with three different substrate primers to avoid false-negative results. In a subset of algal strains (32 in total), the occurrence of variant minisatellite telomeric repeats was examined by Southern hybridization (dot-blot hybridization and/or terminal restriction fragment [TRF] analysis; fig. 4 and supplementary fig. S3, Supplementary Material online) using telomeric oligonucleotide probes (supplementary table S2, Supplementary Material online). A terminal position of candidate telomeric sequences was tested by BAL31 nuclease digestion and Southern hybridization in eight algal species (representative samples shown in fig. 5, supplementary figs. S3 and S4, Supplementary Material online).

Telomerase Activity Screening in Archaeplastida Using TRAP Assay
We investigated 23 and 8 algal strains from Chlorophyta and Streptophyta, respectively, for the presence of a telomerase activity using the reverse primer TELPR30-3A with the Arabidopsis-type telomeric sequence. The Arabidopsis-type sequence was presumed as an ancestral telomere type for this group based on our previous results (Fulněcková et al. 2012). Algal strains from the chlorophyte classes Chlorophyceae, Trebouxiophyceae, and Chlororodendrophyceae showed positive telomerase activity with products of a 7-nucleotide (nt) periodicity (fig. 2) and cloned TRAP products confirmed synthesis of the Arabidopsis-type telomeric sequence (table 1). A comparison of telomerase activity in the “crude” telomerase extracts, the PEG-purified extracts, and the PEG-norpurified protein fraction revealed that the telomerase activity is present also in the PEG-norpurified fraction in all these telomerase-positive algal strains (supplementary table S3, Supplementary Material online). However, algal strains from the class Ulvophyceae failed to show a reproducible telomerase activity. Testing of reverse primers with sequences corresponding to alternative telomere types or minisatellite variants and/or using different substrate primers to cope with possible telomerase substrate preference did not produce positive results (supplementary table S4, Supplementary Material online). A control experiment excluded the presence of telomerase inhibitors in algal extracts (supplementary fig. S1, Supplementary Material online; discussed later). In two cases, we experienced a very weak ladder of TRAP products using human-type reverse primer; however, we identified fungal contaminants in the respective two algal cultures by PCR (see Materials and Methods), which might be responsible for this residual activity in the samples tested.

Three algal strains representing different branches of the streptophyte class Zygnematophyceae (TEL181 Zygnema
**FIG. 2.**—Telomerase activity in Archaeplastida investigated by TRAP assay. Telomerase activity in representative algal strains of Glaucophyta (A, TEL195 Gl. nostochinearum), Rhodophyta (B, TEL213 R. maculata), Chlorophyta (C, TEL211 T. chui; D, TEL121 Dictyochloropsis irregularis; E, TEL94 Pseudendocloniopsis botryoides and TEL124 Pseudendoclonium basiliense), and Streptophyta (F, TEL198 Mesotaenium endlicherianum; G, TEL97 Klebsormidium subtilissimum, TEL100 K. dissectum, TEL101 K. flaccidum, TEL103 K. nitens, TEL187 K. crenulatum) grouped according to their phylogenetic provenance (indicated above panels); the activity is shown using combinations of substrate and reverse primers—GG(21) and HUTC (human-type primer) (A), 47F and TELPR30-3A (Arabidopsis-type) (B, C, F), or pSSyF and TELPR30-3A (D, E). Synthesis of telomeric repeats corresponding to the human-type and the Arabidopsis-type sequence (compare with table 1) was observed in Glaucophyta (A) and three green algal classes (C, Chlorodendrophyceae; D, Trebouxiophyceae; E, Zygnematophyceae), respectively. Negative results were obtained in Rhodophyta (B) and Ulvophyceae (E). The Klebsormidiophyceae samples (G) showed synthesis of two different telomere types; alternative combinations of the substrate primer TS21 and the Chlamydomonas-type repeat (T4AG2-C), or of the substrate primer pSSyF and the TTTAGG-type repeat reverse primer (CHTRAPRev1), or of the sample control (0.3 µg) were used as a pattern control of an 8-, a 7-, and a 6-nt periodicity ladder, respectively; negative control (−), no extract.

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Diversity and Evolution of Telomeres in Eukaryotes

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**FIG. 2.**—Telomerase activity in Archaeplastida investigated by TRAP assay. Telomerase activity in representative algal strains of Glaucophyta (A, TEL195 Gl. nostochinearum), Rhodophyta (B, TEL213 R. maculata), Chlorophyta (C, TEL211 T. chui; D, TEL121 Dictyochloropsis irregularis; E, TEL94 Pseudendocloniopsis botryoides and TEL124 Pseudendoclonium basiliense), and Streptophyta (F, TEL198 Mesotaenium endlicherianum; G, TEL97 Klebsormidium subtilissimum, TEL100 K. dissectum, TEL101 K. flaccidum, TEL103 K. nitens, TEL187 K. crenulatum) grouped according to their phylogenetic provenance (indicated above panels); the activity is shown using combinations of substrate and reverse primers—GG(21) and HUTC (human-type primer) (A), 47F and TELPR30-3A (Arabidopsis-type) (B, C, F), or pSSyF and TELPR30-3A (D, E). Synthesis of telomeric repeats corresponding to the human-type and the Arabidopsis-type sequence (compare with table 1) was observed in Glaucophyta (A) and three green algal classes (C, Chlorodendrophyceae; D, Trebouxiophyceae; E, Zygnematophyceae), respectively. Negative results were obtained in Rhodophyta (B) and Ulvophyceae (E). The Klebsormidiophyceae samples (G) showed synthesis of two different telomere types; alternative combinations of the substrate primer TS21 and the Chlamydomonas-type repeat reverse primer (CHTRAPRev1), or of the substrate primer pSSyF and the TTTAGG-type repeat reverse primer (T4AG2-C), displayed synthesis of a 7- or an 8-nt periodicity of TRAP products (arrows) by telomerase of K. subtilissimum (TEL97) or other Klebsormidium spp., respectively (table 1). Differences in efficiency of telomerase purification during preparation from protein extracts are documented in samples shown on C, D, F, and G (summarized in supplementary table S3, Supplementary Material online). Triangles indicate different amounts of total protein (0.1 and 1 µg) in protein extract without PEG precipitation (crude, cr.), in fractions nonprecipitated (supernatant, sup) and precipitated by PEG (telomerase extract, ex), except TEL94 (E: 0.1, 0.2 µg), TEL124 (E: 0.1, 0.5 µg), TEL97 (G: 0.1, 0.8 µg), TEL100 (G: 0.1, 0.3 µg), and TEL101 (G: 0.1, 0.4 µg). When one sample is indicated, 1 µg or a higher amount of total protein mentioned earlier was used, except TEL121 (D: all 0.5 µg), TEL211 (C: sup 0.5 µg), TEL94 (E: cr. 0.1, sup 0.3 µg), and TEL101 (G: sup 0.4 µg). Telomerase-enriched extracts (50 ng of total protein) from Chlamydomonas hydra (TTTGGG), Arabidopsis thaliana seedlings (TTTAGG), and Euglena stellata (TTAGGG) were used as a pattern control of an 8-, a 7-, and a 6-nt periodicity ladder, respectively; negative control (−), no extract.
Three algal strains covering different branches of rhodophytes failed to show telomerase activity when investigated using six variants of a reverse oligonucleotide primer derived from the telomeric sequence known in *Chlamydomonas* (supplementary tables S2 and S4, Supplementary Material online) or four alternative reverse primers (supplementary table S4, Supplementary Material online) including the *Arabidopsis*-type (fig. 2 and table 1). To check whether the failure of the TRAP assay could be caused by the presence of inhibitors in algal telomerase extracts, we performed a control experiment, in which the negative red algal telomerase extracts was added to a positive control extract from *A. thaliana* in ratio 1:1 or 3:1. Neither of the red algal extracts posed a clear inhibitory effect (supplementary fig. S1, Supplementary Material online). Further analyses of telomerase activity using three different substrate primers with the human-type reverse primer showed a positive result in all three red algal samples, but only using the substrate primer 47F (supplementary fig. S5, Supplementary Material online). To verify the identity of the amplified minisatellite repeat, we performed TRAP reactions also with an alternative reverse primer (T3AG2-C) that was able to amplify successfully the TRAP products containing human-type repeats from other algal species (discussed in supplementary fig. 3).
| Phylum/Class | TEL | Species | Telomere Type | No. of Clones | No. of Repeats | Activity (Primers) | Minisatellite Variant | T/G-Slippage | Mismatches | Other Combinations |
|-------------|-----|---------|---------------|---------------|---------------|-------------------|----------------------|--------------|-------------|-------------------|
| **Glaucophyta** | | | | | | | | | | |
| 195 | Glaucostis nostochinearum | TTAGGG | 2 | 39 | +++ (a, f) | 38 | 1 | 0 | 0 | 0 | +++ (b, f) |
| Rhodophyta | 131 | Porphyridium purpureum | Unknown | n.a. | /C0 | | | | | | |
| 213 | Rhodella maculata | Unknown | n.a. | | | | | | | |
| 214 | Rhodosorus marinus | Unknown | n.a. | | | | | | | |
| **Chlorophyta/Chlorophyceae** | | | | | | | | | | |
| 87 | Scenedesmus vacuolatus | TTAGGG | 2 | 17 | ++ (e, g) | 0 | 17 | 0 | 0 | 0 | |
| 89 | Muriella decor | TTAGGG | 4 | 78 | +++ (c, f) | 0 | 77 | 1 | 0 | 0 | |
| 91 | Mychonastes homosphaera | TTAGGG | 3 | 33 | ++ (e, g) | 0 | 33 | 0 | 0 | 0 | |
| 98 | Chromochloris zofingiensis | TTAGGG | 6 | 95 | +++ (a, f) | 0 | 50 | 1 | 0 | 4 | |
| 108 | Neochloris conjuncta | TTAGGG | 3 | 24 | +++ (b, f) | 0 | 24 | 0 | 0 | 0 | |
| 123 | Chlorochloris hypnoroporum | TTAGGG | 5 | 38 | +++ (b, f) | 1 | 38 | 0 | 0 | 1 | 1 | +++ (c, f) |
| 138 | Pseudomonilia aurantica | TTAGGG | 2 | 18 | +++ (b, f) | 2 | 13 | 1 | 0 | 0 | |
| 140 | Follicularia paradoxicus | TTAGGG | 3 | 21 | +++ (b, f) | 1 | 20 | 0 | 0 | 0 | +++ (c, f) |
| 188 | Bracteacoccus cohaerens | 7-nt (tttaggg) | n.a. | | | | | | | | |
| **Chlorophyta/Trebouxiophyceae** | | | | | | | | | | |
| 84 | Jaagella alpicola | TTAGGG | 2 | 30 | +++ (e, g) | 3 | 24 | 2 | 0 | 1 | 0 | +++ (c, f) |
| 85 | Asterochloris phycobiontica | TTAGGG | 5 | 67 | +++ (c, f) | 0 | 64 | 1 | 0 | 2 | |
| 88 | Heterochlorella luteoviridis | TTAGGG | 3 | 32 | +++ (a, f) | 0 | 31 | 0 | 0 | 0 | 1 | +++ (c, f) |
| 90 | Auxenochlorella protothecoides | TTAGGG | 4 | 30 | +++ (d) | 0 | 30 | 0 | 0 | 0 | 0 | +++ (c, f) |
| 121 | Dictyochloropsis irregularis | TTAGGG | 4 | 51 | +++ (a, f) | 0 | 53 | 0 | 0 | 0 | |
| 134 | Heterochlorella gracilis* | TTAGGG | 7-rt (tttaggg) | n.a. | +++ (c, f) | | | | | | |
| **Chlorophyta/Ulvophyceae** | | | | | | | | | | |
| 211 | Tetraselmis chui | TTAGGG | 4 | 38 | +++ (b, f) | 0 | 22 | 9 | 0 | 5 | 3 | |
| 212 | T. striata | TTAGGG | 4 | 18 | +++ (b, f) | 0 | 9 | 7 | 0 | 2 | 0 | |
| **Chlorophyta/Olvophyceae** | | | | | | | | | | |
| 86 | Planophila laeviriris | Unknown | n.a. | | | | | | | | |
| 94 | Pseudendocloniopsis bartyioides | Unknown | n.a. | | | | | | | | |
| 111 | Pseudendoclonium printii | Unknown | n.a. | | | | | | | | |
| 124 | Pseudendoclonium basiliense | Unknown | n.a. | | | | | | | | |
| 137 | Desmochloris halophila | Unknown | n.a. | | | | | | | | |
| 139 | Pirula salina | Unknown | n.a. | | | | | | | | |

(continued)
| Phylum/Class | TEL | Species | Telomere Type | No. of Clones | No. of Repeats | Activity (Primers) | Minisatellite Variant | T/G-Slippage | Mismatches | Other Combinations |
|-------------|-----|---------|---------------|---------------|---------------|-------------------|----------------------|--------------|------------|-------------------|
| Streptophyta/ Klebsormidiophyceae | 97  | *Klebsormidium subtilissimum* | TTTAGGG | 3 | 53 | ++ (a, i) | 0 | 0 | 0 | 53 | 0 | 0 | ++/ (c, i); ++/ (d, i); ++/ (d, j); − (d, k); ++/ (b, f); − (d, h) |
| | 100 | *K. dissectum* | 8-nt (ttttaggg) | n.a. | + (d, k) | 0 | 0 | 0 | 53 | 0 | 1 | 1 | ++/ (d, β) +/ (d, h) |
| | 103 | *K. nitens* | TTTAGGG | 2 | 25 | + (a, i) | 0 | 0 | 0 | 23 | 0 | 1 | 1 | ++/ (d, β) +/ (d, h) |
| | 187 | *K. crenulatum* | 8-nt (ttttaggg) | n.a. | ++ (d, k) | 0 | 0 | 0 | 33 | 0 | 4 | 0 | − (b, f); − (a, f); + (c, i) + (c, l); + (c, k) + (c, l); + (c, i) |
| Streptophyta/ Zygnematophyceae | 181 | *Zygnema circumcarinatum* | TTTAGGG | 5 | 38 | + (d, f) | 4 | 29 | 0 | 0 | 1 | 1 | + (b, f) |
| | 196 | *Mesotaenium endlichianum* | TTTAGGG | 4 | 38 | +++ (d, f) | 0 | 35 | 0 | 0 | 1 | 1 | |
| | 198 | *Mesotaenium endlichianum* | TTTAGGG | 4 | 18 | +++ (d, f) | 0 | 15 | 0 | 0 | 3 | 0 | +++ (d, f) |
| Ochrophyta/ Xanthophyceae | 95  | *Xanthomonas cf. hormidioides* | TTTAGGG | 1 | 13 | + (b, f) | 2 | 6 | 2 | 0 | 2 | 0 | ++/ (a, f); +/ (c, h) |
| | 202 | *Pseudochloris meiringensis* | TTTAGGG | 4 | 39 | +++ (b, f) | 0 | 39 | 0 | 0 | 0 | 0 | +++ (a, f); +/ (c, h) |
| | 203 | *X. hormidioides* | TTTAGGG | 3 | 35 | +++ (a, f) | 7 | 10 | 0 | 1 | 4 | 0 | |
| | 204 | *Heterococcus protonematoide* | TTTAGGG | 5 | 85 | +++ (b, f) | 0 | 83 | 0 | 0 | 1 | 1 | − (a, f) |
| | 205 | *Batrydiopsis intercedens* | TTTAGGG | 3 | 53 | +++ (a, f) | 6 | 42 | 2 | 0 | 3 | 0 | +++ (b, f); +/ (c, h) +/ (c, h) |
| Ochrophyta/ Eustigmatophyceae | 133 | *Eustigmatos polyphek* | Unknown | n.a. | | | | | | | | | |
| | 201 | *Vischeria punctata* | Unknown | n.a. | | | | | | | | | |
| Ochrophyta/ Bacillariophyceae | 231 | *Paeodactylum tricornutum* | Unknown | n.a. | | | | | | | | | |
| Haptophyta | 210 | *Pavlova lutheri* | TTTAGGG | 7 | 216 | +++ (c, h) | 215 | 0 | 0 | 0 | 0 | 1 | |
| Alveolata | 233 | *Chromera vela* | TTTAGGG | 4 | 90 | ++ (b, f) | 0 | 79 | 0 | 0 | 0 | 11 | 0 |
| Euglenophyceae | 185 | *Euglena anabaena* | TTTAGGG | 2 | 39 | + (b, f) | 32 | 6 | 0 | 0 | 0 | 0 | |
| | 206 | *E. stellata* | 6-nt (ttagg) | n.a. | +++ (c, h) | 35 | 2 | 0 | 0 | 0 | 0 | 0 | |
| | 207 | *E. geniculata* | TTTAGGG | 3 | 49 | ++ (c, h) | 49 | 0 | 0 | 0 | 0 | 0 | − (b, f) |

**Note.**—Telomere type was determined from the nucleotide periodicity of the TRAP products (lower case letters) or from cloned TRAP products (upper case letters). Primers: a, pSSyF; b, 47F; c, GG(21); d, T521; e, CAMV; f, TELPR30-3A; g, TELPR; h, HUTC; i, T4AG2-C; j, T4AG2-PR; k, CHTRTTRAPRev1; l, TTTAGG3-C; n.a., not analyzed. Minisatellite variants refer to number of indicated telomere motifs detected among telomerase products, the major telomere types are indicated in bold. Errors classified as T- or G-slippage are additional T or G nucleotides incorporated into the reiterated unit. Other errors, mostly nucleotide deletion or A/G substitution, are considered nucleotide misincorporations (mismatches).
FIG. 4.—Dot-blot hybridization of genomic DNA with telomere and telomere-like minisatellite probes. Genomic DNA samples (1–2 μg per dot) from algal strains and control samples (listed according to phylogeny position on left) were blotted and hybridized with radioactively labeled oligonucleotide probes representing different telomere types and derived sequences (indicated above dot columns). Samples in which the telomere type was revealed in a telomerase analysis (figs. 2 and 3, table 1) hybridized with the corresponding oligonucleotide, but occurrence of other minisatellites was also indicated. The T4AG2 and CHSB probes cross-hybridized and could not be distinguished by Southern hybridization (for details see supplementary material, Supplementary Material online). The T2CG3 minisatellite showed a signal across algal samples, but not in a terminal position (see supplementary material, Supplementary Material online); a similar situation was described in *Allium* (*Sykorova et al.* 2006). Control DNA samples represent the *Arabidopsis*-type (*Chlorella vulgaris*) and the human-type telomeres (*human* and *Ipheion uniflorum*, Alliaceae); note that the plant DNA contains also a portion of the ancestral *Arabidopsis*-type minisatellite. Control rehybridization of membranes was done with a mixed rDNA probe (see Materials and Methods for details); n.a., not analyzed.
Telomerase Activity in Other Algal Groups

We investigated telomerase activity also in representatives of other algal groups. According to published data, diatoms and euglenophytes should possess telomeres formed by the human-type telomere repeats (Dooijes et al. 2000; Armbrust et al. 2004), but we were not able to detect any telomerase activity (fig. 3) in the diatom TEL231 *P. tricornutum*. In contrast, the three *Euglena* species tested (Euglenophyceae) and the haptophyte TEL210 *Pavlova lutheri* displayed high telomerase activity and synthesis of the human-type telomeric repeats (table 1). Control experiments designed to investigate possible preferences in substrate primer sequence or presence of inhibitors in diatom telomerase extract excluded these technical reasons of the TRAP assay failure (supplementary table S4 and fig. S1, Supplementary Material online; refer the earlier discussion). An observed 7-nt periodicity and cloning of the TRAP products confirmed synthesis of the expected Arabidopsis-telomeric type in C. veila (fig. 3 and table 1), which is in agreement with its phylogenetic position within Alveolata close to dinoflagellates known from previous experiments to possess telomeres formed by the Arabidopsis-telomeric type sequence (Fojtová et al. 2010). Samples from the classes Xanthophyceae and Eustigmatophyceae showed very different results, despite the fact that they both belong to the algal phylum Ochrophyta within Stramenopiles. Although all five of xanthophytes (TEL95 *Xanthonema* cf. hormidoides, TEL202 *Pleurochlois* meiningensis, TEL203 *Xanthonema* debile, TEL204 *Heterococcus* proconenatoideae, and TEL205 *Botrydiopsis* intercedens) showed telomerase synthesizing the Arabidopsis-telomeric type sequence, the two eustigmatophyte strains investigated (TEL133 *Eus. polyphem* and TEL201 *V. punctata*) did not reveal any reproducible telomerase activity (fig. 3). Similar to diatoms, control experiments using different combinations of substrate and reverse primers showed negative result (supplementary table S4, Supplementary Material online), whereas a presence of inhibitors was excluded (discussed earlier, supplementary fig. S1, Supplementary Material online). A PEG purification step was successful in telomerase enrichment of protein extracts from algal strains of Euglenophyceae, Haptophyta, Xanthophyceae, and C. veila (supplementary table S3, Supplementary Material online).

Dot-Blot Hybridization Screening and Testing for Telomeric Localization of Minisatellite Repeats Using BAL 31 Digestion

We screened samples of algal genomic DNA by Southern hybridization using radioactively labeled oligonucleotides as probes (fig. 4) to unveil a possible occurrence of other telomere-like minisatellites in the respective genomes and to possibly identify candidate telomeric sequences in samples with no detected telomerase activity. We experienced difficulties in DNA extraction from several algal strains, mainly from Zygningmatoxyceae and rhodophytes, which showed the presence of colored substances and a low DNA yield; moreover, genomic DNA extraction was not successful for TEL196 *Micrasterias crux-melitensis* and TEL198 *Mesotaenium endlicherianum*. For the remaining samples from Chlorophyta, Streptophyta, Xanthophyceae, Euglenophyceae, Haptophyta, Glauco cytphyta, and C. veila, the dot-blot hybridization confirmed
the presence of telomeric minisatellites identified as “true”
telomeric types synthesized by telomerase in the respective
algal strains. However, dot-blot hybridization of genomic DNA
from telomerase-negative strains did not suggest any
other candidate telomeric sequence and in general, dot-blot
hybridization signals were much weaker than we experienced
in our previous study (Fulnečková et al. 2012). A weak signal of
control hybridization with a mixed probe consisting of a mix-
ture of LSU and SSU rDNA sequences (see Material and
Methods) may be caused by a wide phylogenetic span of our
algal collection and a limited similarity among rDNA sequences
or by a low quality of genomic DNA prepared by the proteinase
K-based method, because we observed difficulties in PCR am-
plification of control rDNA sequences and other Southern
hybridization experiments (supplementary material, Supple-
mentary Material online). The terminal position of a candidate
human-type telomeric sequence in euglenophytes (TEL206
and TEL207) and the terminal position of the *Chlamydomonas*
type or the TTTTAGG-type of a telomeric sequence in
Klebsormidiophyceae (TEL103, TEL187, and TEL97) were veri-
fied using BAL 31 nuclease digestion (fig. 5; supplementary
figs. S3 and S4, Supplementary Material online). Subsequent
dehybridization of BAL31-digested samples of TEL97 (*K. sub-
tilissium*) with the *Chlamydomonas*-type sequence probe
confirmed the presence of both sequence types in TRFs (sup-
plementary fig. S3B, the bottom panel, Supplementary
Material online). Investigation of the TRF lengths showed
that the TTTTAGG-type sequences hybridize with 0.7–1.5 kb
long fragments (supplementary fig. S3A, right panel,
Supplementary Material online), suggesting short telomers
similar to Chlamydomonadales. Correspondingly, digestion
with Small digestion produced longer restriction fragments
and the signal of both TTTTAGG- and *Chlamydomonas*-type
probes was distributed among multiple BAL31-sensitive frag-
ments of 2.5–23 kb length (supplementary fig. S3B, the bottom panel, Supplementary
Material online). Besides these, short BAL31-resistant fragments (1.3–2.3 kb) representing in-
terstitial telomeric sequences could also be seen in the hybrid-
ization patterns of both probes. The presence of internal
telomere repeats is also apparent in *K. crenulatum* (supple-
mentary fig. S4A, Supplementary Material online). Although high-
molecular-weight restriction fragments hybridizing with
*Chlamydomonas*-type probe shortened upon BAL31 treatment
(supplementary fig. S4A, the left panel, Supplementary Mate-
rial online), the low-molecular-weight fragments were resistant
to BAL31, which reflects their internal (nontelomeric) positions
(supplementary fig. S4A, the right panel, Supplementary
Material online). We also performed BAL31 digestion on
both strains of Eustigmatophyceae to check whether the qual-
ity of the genomic DNA could be the reason for the failure of
dot-blot hybridization. Probing with the *Arabidopsis*-type or
the human-type telomeric sequence, which are expected as
candidate telomere types due to the phylogenetic position of
Eustigmatophyceae in Ochrophyta, did not produce any
specific signal (supplementary fig. S4, Supplementary
Material online), thus confirming the negative results of the
TRAP assay and dot-blot hybridization. BAL 31 nuclease diges-
tion was performed also in TEL31 *Porphyridium purpureum*
samples, but both investigated probes (Cyanidioschyzon-type
and human-type) failed to show any specific signal.

Identification of Candidate Telomere Sequences in
Genome Sequences of Phylogenetically Diverse
Eukaryotes

To cover a wider spectrum of phylogenetic lineages across
the tree of eukaryotic life, we coupled our experimental in-
vestigations with in silico searches for candidate telomeric
sequences in published or publicly available genome se-
quences, focusing on groups that have been ignored or
poorly studied with regard to their telomeres. In addition,
we take advantage of the genome data yielded by our
on-going genome sequencing projects for three phylogeneti-
ically unique organisms, the jakobid *And. godoyi*, the mala-
wimonad *M. californiana*, and the goniomonad *Gon.
avonlea*. We also used available genomic sequences to
verify the presence of telomeric sequences that have been
described previously for the respective organisms by meth-
ods in telomere biology. We searched the genome assem-
blies for stretches consisting of repeated units of the major
known types of telomeric sequences (T,AnGn) and assessed
them as candidate telomeric minisatellites by taking into
account their position and orientation with respect to adjacent
sequences. Our simple database search could not uncover
degenerated telomere types, like those known from yeasts,
and experimental tests are also required to confirm the ter-
minal position of the candidate sequences. We searched 143
genomes (including 32 from species where the telomeric
sequence has been published before) and 80 of them
showed a convincing pattern of telomeric sequence (supple-
mentary table S5, Supplementary Material online). A majority
of the genomes that showed a dominant presence of the
candidate sequence in terminal regions also exhibited intern-
al telomeric repeats occurring in short stretches or in large
blocks (>100 bp of uninterrupted minisatellite). The gen-
omes where we found only short or occasional repeats
positioned terminally and/or in large internal blocks were
considered inconclusive and ignored for the summary of the
phyletic distribution of telomeric sequences in eukaryotes
shown in figure 1 (except species with previously published
telomeric sequences; supplementary table S5, Supple-
mentary Material online). In several cases, we identified
unexpected candidate telomeric repeats in genomes repre-
senting hitherto unstudied key phylogenetic groups, for
example, TTTCGGG in the parasitic relative of dinoflagellates
*Perkinsus marinus*, TTTCGGG in the heterolobosean
*N. gruberi*, TTAGG in the labyrinthulid *Aurantiochytrium limacina*, and the highly unusual 10–11 nucleotide
repeat unit TTTATT(T)AGGG in the rhodophyte *Galdieria sulphuraria* (fig. 1 and supplementary table S5, Supplementary Material online). In addition, minisatellites that differed from the types “canonical” for the respective organismal groups were found in fungi and stramenopiles, indicating the evolutionary flexibility of the telomeric sequence at various phylogenetic scales. Our database searches corroborated the experimental results from Haptophyta, Glauco phyta, and Chlorophyta, whereas no genome assemblies from Xanthophyceae, Chromera, Ulvophyceae, Euglenophyceae, or dinoflagellates were available for analysis. The genome assemblies of two Nannochloropsis species (Eustigmatophyceae) displayed the presence of large internal blocks of TTAGGG-type repeats in addition to several terminally positioned stretches; and without experimental evidence, these repeats should be taken as a candidate telomere sequence. Searches of unassembled genomic reads available for the red alga Por. umbilicalis did not identify the telomere types described in *Cya. merolae* genome or predicted in the *Galdieria sulphuraria* genome (discussed earlier), but revealed a large number of reads containing a TTAGGG-type minisatellite. In several cases, these repeats could be assessed as internal, but whether any of the other sequences represent the true telomere cannot be verified without a full genome assembly or an experiment. A similar result was achieved when we searched unassembled Sanger reads from an on-going genome project for the haptophyte *Pha. antarctica*.

**Discussion**

People have long been fascinated by the question how mechanisms of linear chromosome maintenance might have originated. It is believed that recombination-based pathways, which today serve mostly as a backup mechanism (Fajkus et al. 2005), were original and were subsequently replaced by a more successful, steady, and efficient synthesis of telomeres consisting of minisatellite sequences by telomerase. Telomeres of most investigated organismal groups are generally conserved within the groups and conform to a limited number of minisatellite types. The repeat units of these minisatellites are mainly variants of the $T_mA_nG_2$ sequence and their evolutionary success presumably depends on their properties stemming from the G-rich sequence and their capacity to form alternative DNA structures like the G-quartet or the T-loop typical for telomere function (de Lange 2005). The telomere sequence of the red alga *Cya. merolae* is somewhat atypical, but it is a G-rich sequence and genes for telomerase subunits were predicted in the genomic sequence (Nozaki et al. 2007). It is generally assumed that telomeres are formed by T/G-rich minisatellites and maintained by telomerase until alternative telomere and maintenance mechanisms are shown for a given organism. The results of our new experimental and in silico analyses have substantially expanded the sampling of telomere structures across the eukaryotic phylogeny. In addition, telomerase activity has also now been tested against a number of evolutionarily distant groups. The data enable us to paint a picture of telomere evolution across eukaryotes with an unprecedented level of detail (fig. 1), although many aspects of the scheme remain to be clarified.

**Characterization of Telomeres in Many Algal Groups Remains Technically Challenging**

In algae, telomerase activity has been previously experimentally proven in photosynthetic and nonphotosynthetic dinoflagellates (Fojtova et al. 2010; Zielke and Bodnar 2010) and in chlorophytes (Fulnecˇkova et al. 2012). Despite a published completed genome sequence including predicted telomere sequences and a telomerase gene (Bowler et al. 2008), our TRAP assay was unsuccessful in the diatom *P. tricornutum*. This experimental failure could be for some unknown technical reason or for the telomerase being active only in specific developmental stages (e.g., during auxospore formation) that were not presented in our samples at a detectable level. Another possible explanation comes from experiments in the silkworm *Bombbyx mori*, where telomeres are maintained by telomerase with a very low processivity and thus difficult to be detected (Sasaki and Fujiwara 2000). Active telomerase synthesizing the Arabidopsis-type telomeric sequences was demonstrated by our experiments in *C. velia* (Alveolata), Xanthophyceae, chlorophytes, and streptophyte algae. In contrast, we demonstrated the human-type telomeric sequence synthesized by telomerase in euglenophytes, haptophytes (Pav. lutheri), and glaucophytes (Gla. nostochinearum).

Unexpectedly, we did not detect telomerase activity in the classes Eustigmatophyceae (Ochrophyta) and Ulvophyceae (Chlorophyta), and also Southern hybridization of genomic DNA with other minisatellite telomeric probes failed to resolve telomeric sequences, as was observed also in rhodophytes. A low signal intensity by Southern hybridization could be caused by impurities in the genomic DNA, which in our case had to be isolated by the protease K-based method (see supplementary material, Supplementary Material online). Control TRAP experiments excluded the presence of telomerase inhibitors in protein extracts of all telomerase-negative algal strains (a diatom, Eustigmatophyceae, Ulvophyceae, and Rhodophyta). Our use of a different substrate primers should accommodate variable telomerase preferences for substrate sequence. Despite these controls, it remains possible that there are technical reasons that prevent the telomere motifs from being resolved. Remarkably, we found a similar behavior of telomerases during PEG purification from protein extracts in various algal groups, suggesting their similar biochemical properties. In most cases, telomerase was enriched in a purified fraction, with the exception of protein extracts from chlorophytes and a glaucophyte alga, which showed telomerase activity also in the nonprecipitated fraction (supplementary table S3, Supplementary Material online). However, PEG
purification led to removal of compounds inhibiting TRAP assay from these extracts (Ševčíková et al. 2013).

We did not detect telomerase activity in three red algal representatives investigated for synthesis of Cyanidioschyzon-type telomeric sequences. This may be due to specific problems with PCR-based TRAP assay (a difficult G-rich template of a candidate sequence) or to occurrence of different telomeric sequences, because the investigated red algal species are only distantly related to Cya. merolae. The behavior of the three rhodophyte strains investigated here does not seem to be a peculiarity of one particular lineage, because they represent three different deeply diverged rhodophyte classes (fig. 1 and supplementary table S1, Supplementary Material online). Searches for telomeric motifs in databases suggest that some rhodophytes might have the human-type telomeric sequence, because a great portion of genomic reads from Por. umbilicalis (representing an additional class Bangiophyceae) contain this sequence type. However, many of them could be scored as internal sequences, and other telomere-like minisatellites also occur in the genomic reads, making it difficult to distinguish the "true" telomeric type without an experiment. Moreover, the assembled genome sequence of Gal. sulphuraria (a relative of Cya. merolae also belonging to Cyanidiophyceae) showed the presence of an unusual minisatellite repeat (TTTATTAGGG or TTTATTTAGGG) predominantly at the ends of scaffolds, suggesting a putative telomeric position. The sequence of this repeat unit seems to be an AT-rich variant derived from the typical TTAGGG telomeric minisatellite. Neither the human-type nor other investigated telomere types (including the Cyanidioschyzon-type) were found in the Galdieria genome assembly. The path of the telomere evolution in rhodophytes thus remains unclear. Assuming that Archaeplastida are monophyletic (but see, e.g., Burki et al. 2012) and that the human-type telomeric repeat occurs in the presumably basal archaeplastid lineage Glaucophyta as well as in most other eukaryotic lineages (discussed later), it is possible that the last common ancestor of Archaeplastida had the TTAGGG telomeric sequence. This would then mean that this telomeric type could be retained by some red algal lineages, but further testing of archaeplastid monophyly, and more evidence for the presence of the human-type telomere in rhodophytes, are needed to corroborate this scenario.

Novel Telomere Forms Seem to Have Evolved in Some Algal Groups

In plants and algae, two categories of evolutionary changes of telomere sequences have been described previously: 1) change of the telomeric sequence synthesized by telomerase to a related minisatellite sequence, that is, from the Arabidopsis-type to the human type in Asparagales (Sýkorová et al. 2003b) and from the Arabidopsis-type to the Chlamydomonas type or the human type in Chlamydomonadales (Fulnečková et al. 2012); and 2) loss of a typical minisatellite and of telomerase activity reported in the genus Allium (Asparagales) (Sýkorová et al. 2006) and three genera of Solanaceae (Sýkorová et al. 2003a). Our results bring evidence for further examples following the first and possibly also the second category.

The first is exemplified by switches to variant minisatellite telomeric sequences in Klebsormidio phyceae, specifically a 1-nt addition resulting in a change from the ancestral TTTAGGG (Arabidopsis type) to TTTTAGGG (Chlamydomonas type) early in the evolution of Klebsormidio phyceae, followed by a 1-nt deletion in the lineage leading to TEL97 K. subtilissimum, which resulted in the unusual TTTTAGG sequence (this interpretation is based on the nested phylogenetic position of the latter K. subtilissimum within the genus Klebsormidium; Rindi et al. 2011). The TEL97 strain still possesses a large amount of ancestral Chlamydomonas-type sequence in its genome, which is similar to the situation in Hyacinthaceae (Asparagales) that have telomerase synthesizing the human-type sequence, but the ancestral telomere type still occurs in the genome at a level detectable by fluorescent in situ hybridization (Adams et al. 2001). The negative result of the telomerase assay using typical telomeric minisatellites as reverse primers in Eustigmatophyceae and Ulvophyceae suggests a change of the second type, but a more detailed study is needed to pinpoint the evolutionary transition in the telomere sequence and/or maintenance mechanism. Similarities between land plants and algae are also seen in different substrate primer usage (see supplementary material, Supplementary Material online) that was demonstrated in Asparagales, and in that case its phyletic pattern did not show a simple correspondence to the phylogeny of this group (Sýkorová, Leitch, Fajkus 2006). It seems that the substrate primer usage (supplementary fig. S2, Supplementary Material online) is linked to enzyme properties of telomerases in different species, which generally show a varying accuracy in minisatellite repeat synthesis (Fitzgerald et al. 2001; Sýkorová et al. 2003b).

Telomere Evolution in the Context of the Phylogeny of Eukaryotes

Although the telomere structure remains unknown or enigmatic for many deeply branching eukaryotic lineages, including those with reference genome sequences available (e.g., Parabasalia, Archaeomobae, or Apusomonadida), a combination of previously published data with the results of our experiment and in silico searches allows to paint at least the major outlines of telomere evolution in eukaryotes (fig. 1). Metazoa (Traut et al. 2007; Gomes et al. 2010) and Fungi (Teixeira and Gilson 2005) show the TTAGGG-type as ancestral to both groups and all changes in their telomeric sequences seem to have happened later in their evolution. The same evolutionary path, that is, from the ancestral TTAGGG type to secondarily derived alternative telomeric repeats, can
now be deduced for Amoebozoa, Excavata, Stramenopiles, and probably also Archaeoplastida (fig. 1). The TTAGGG motif remains the only telomeric repeat known in Choanoflagellata, Haptophyta, and Rhizaria (excluding nucleomorph genomes); hence, it is likely that it is also ancestral for these groups, but a much better sampling (especially for Rhizaria) is needed to confirm this. On the other hand, the plant type of telomeric repeats (TTTAGGG) may be ancestral in cryptomonads, which include cryptophyte algae and their heterotrophic sister lineage goniomonads. Inference on the type of the telomeric repeat is difficult for the last common ancestor of Alveolata, because the two principal lineages, that is, Myzozoa (including dinoflagellates, Perkinsids, apicomplexans, and chromerids) and ciliates, have different telomeres.

Considering the wide occurrence of the TTAGGG telomeric repeat and its inferred ancestral presence in most major eukaryotic groups, it is tempting to suggest that the human-type telomeric repeat was ancestral for eukaryotes as a whole. Deducing the ancestral state of any character for any taxon requires knowledge on the position of the taxon’s root (the deepest branching point). The question about the root of the eukaryotic phylogeny has not yet settled, and at least four contradictory hypotheses, supported by different sources of evidence, have been suggested recently (fig. 1). Interestingly, all those root positions are compatible with the idea of the TTAGGG being the ancestral telomeric repeat for eukaryotes.

Another important aspect to consider is the frequency of homoplasies in the evolution in telomeric sequences. Indeed, assuming the TTAGGG motif as ancestral, different phylogenetic lineages evolved independently on each other to the same alternative motifs. A most notable case concerns the TTAGGG being the ancestral telomeric repeat for Myzozoa (including dinoflagellates, Perkinsids, apicomplexans, and chromerids) and ciliates, with different telomeres.

In summary, our experiments and analyses substantially extend the sampling of telomere diversity in eukaryotes and strengthen the view of telomeres as evolutionarily flexible structures of eukaryotic genomes. The much more comprehensive picture of the phyletic distribution of various telomere types should facilitate future studies on mechanistic causes of the evolutionary changes in telomeres and on underlying driving forces.

Supplementary Material
Supplementary material, figures S1–S5, and tables S1–S5 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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