



TAS3 miR390-dependent loci in non-vascular land plants: towards a comprehensive reconstruction of the gene evolutionary history

Sergey Y. Morozov¹, Irina A. Milyutina¹, Tatiana N. Erokhina²,
Liudmila V. Ozerova³, Alexey V. Troitsky¹ and Andrey G. Solovyev^{1,4}

¹ Belozersky Institute of Physico-Chemical Biology, Moscow State University, Moscow, Russia

² Shemyakin-Ovchinnikov Institute of Bioorganic Chemistry, Russian Academy of Science, Moscow, Russia

³ Tsitsin Main Botanical Garden, Russian Academy of Science, Moscow, Russia

⁴ Institute of Molecular Medicine, Sechenov First Moscow State Medical University, Moscow, Russia

ABSTRACT

Trans-acting small interfering RNAs (ta-siRNAs) are transcribed from protein non-coding genomic TAS loci and belong to a plant-specific class of endogenous small RNAs. These siRNAs have been found to regulate gene expression in most taxa including seed plants, gymnosperms, ferns and mosses. In this study, bioinformatic and experimental PCR-based approaches were used as tools to analyze TAS3 and TAS6 loci in transcriptomes and genomic DNAs from representatives of evolutionary distant non-vascular plant taxa such as Bryophyta, Marchantiophyta and Anthocerotophyta. We revealed previously undiscovered TAS3 loci in plant classes Sphagnopsida and Anthocerotopsida, as well as TAS6 loci in Bryophyta classes Tetrarchidiopsida, Polytrichopsida, Andreaeopsida and Takakiopsida. These data further unveil the evolutionary pathway of the miR390-dependent TAS3 loci in land plants. We also identified charophyte alga sequences coding for SUPPRESSOR OF GENE SILENCING 3 (SGS3), which is required for generation of ta-siRNAs in plants, and hypothesized that the appearance of TAS3-related sequences could take place at a very early step in evolutionary transition from charophyte algae to an earliest common ancestor of land plants.

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Corresponding author

Sergey Y. Morozov,
morozov@genebee.msu.su

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page 19

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INTRODUCTION

Plant chromosomal loci of trans-acting small interfering RNAs (ta-siRNAs) and microRNAs (miRNAs) encode non-protein-coding and protein-coding precursor transcripts, which are synthesized by RNA polymerase II and include cap-structures and poly-(A) tails. In plants, primary miRNA transcripts forming internal imperfect hairpins are processed by a protein complex including Dicer-like protein 1 (DCL1), HYL1 and SERRATE to give RNA duplexes with 2-nucleotide 3'-overhangs, which are then terminally methylated by specific RNA methylase HEN1. One strand of such duplexes, being typically of 21 nucleotides in

length and representing a mature miRNA, is selectively recruited by Argonaut (AGO) family protein to an effector complex targeting a specific RNA for AGO-mediated endonucleolytic cleavage or translational repression (Rogers & Chen, 2013; Axtell, 2013; Bologna & Voinnet, 2014; Borges & Martienssen, 2015; Chorostecki et al., 2017).

Some specific microRNAs are able to initiate production of ta-siRNAs (and other secondary phased RNAs—phasiRNAs) by a step-by-step processing of long double-stranded RNA by DCL4 from a start point defined by miRNA-directed cleavage of a single-stranded RNA precursor in a “phased” pattern. These PHAS loci include non-coding TAS genes and genes encoding penta-tricopeptide repeat-containing proteins (PPRs), nucleotide-binding and leucine-rich repeat-containing proteins (NB-LRRs), or MYB transcription factors (Allen & Howell, 2010; Zhai et al., 2011; Xia et al., 2013; Fei, Xia & Meyers, 2013; Axtell, 2013; Yoshikawa, 2013; Zheng et al., 2015; Komiya, 2017; Liu et al., 2018; Deng et al., 2018). Biogenesis of ta-siRNAs includes initial AGO-dependent miRNA binding at single or dual sites of the precursor transcripts and their subsequent cleavage. The further process is dependent on plant RNA-dependent RNA polymerase 6 (RDR6) and SGS3 proteins participating in the formation of dsRNA, which is then cleaved in a sequential and phased manner by DCL4 with assistance of DRB4 (dsRNA binding protein). The resulting ta-siRNAs (mostly of 21 bp in length), similar to miRNAs, are methylated by HEN1 protein (Allen & Howell, 2010; Axtell, 2013; Fei, Xia & Meyers, 2013; Yoshikawa, 2013; Bologna & Voinnet, 2014; Komiya, 2017; Deng et al., 2018).

Arabidopsis TAS3a transcript, first identified by Allen et al. (2005), gives rise to two near-identical 21-nucleotide tasiARFs targeting the mRNAs of some auxin-responsive transcription factors (ARF2, ARF3/ETT and ARF4). Most angiosperm TAS3 primary transcripts are recognized by miR390 and cleaved by AGO7 at the 3' target site, whereas the 5' miRNA target site is non-cleavable. However, the number of miR390 cleavage sites, organization of tasiARF sequence blocks and phasing registers may vary among different TAS3 genes of vascular plants (Allen & Howell, 2010; Axtell, 2013; Fei, Xia & Meyers, 2013; Zheng et al., 2015; Xia et al., 2013; Xia, Xu & Meyers, 2017; De Felippes et al., 2017; Komiya, 2017; Deng et al., 2018). Moreover, miR390 may additionally target and inhibit protein-coding gene transcripts, such as StCDPK1 related to auxin-responsive pathway (Santin et al., 2017).

Previously, we described a new method for identification of plant TAS3 loci based on PCR with a pair of oligodeoxyribonucleotide primers mimicking miR390. The method was found to be efficient for dicotyledonous plants, cycads, conifers, and mosses (Krasnikova et al., 2009; Krasnikova et al., 2011; Krasnikova et al., 2013; Ozerova et al., 2013). Importantly, at that time the structural and functional information on bryophyte TAS3 loci was available only for the model plant *Physcomitrella patens* (Arif, Frank & Khraiwesh, 2013), and we used our PCR-based approach as a phylogenetic profiling tool to identify relatives of *P. patens* TAS3 loci in 26 additional moss species of class Bryopsida and several mosses of classes Polytrichopsida, Tetraphidopsida and Andreaeopsida. Moreover, we found a putative pre-miR390 genomic sequence for an additional moss class, Oedipodipsida (Krasnikova et al., 2013). Our studies revealed that a representative of Marchantiophyta (liverwort *Marchantia polymorpha*, class Marchantiopsida) could also encode a candidate miR390 gene

and a potential TAS3-like locus (Krasnikova *et al.*, 2013). This finding extended the known evolutionary history of TAS3 loci to the proposed most basal land plant lineage (Ruhfel *et al.*, 2014; Bowman *et al.*, 2017). In addition, we sequenced putative pre-miR390 genomic locus for *Harpanthus flotovianus* (Marchantiophyta, class Jungermanniopsida) (Krasnikova *et al.*, 2013). Later, our findings of TAS3-like and miR390 loci were experimentally confirmed in the studies of the transcriptomes of Marchantiophyta plants *M. polymorpha* (Lin *et al.*, 2016; Tsuzuki *et al.*, 2016) and *Pellia endiviifolia* (class Jungermanniopsida) (Alaba *et al.*, 2015).

New genomic and transcriptomic sequence data for basal Viridiplantae appeared in NCBI (<http://ncbi.nlm.nih.gov/sra>) and Phytozome (<http://www.phytozome.net>) databases prompted us to perform new experimental and *in silico* analyses of TAS3 loci in basal taxons of Viridiplantae. In this paper, we identified previously unrecognized TAS3 loci in classes Sphagnopsida and Anthocerotopsida, as well as composite TAS6/TAS3 loci in classes Tetrarchidiopsida, Polytrichopsida, Andreaeopsida and Takakiopsida. Additionally, we revealed SGS3-coding sequences in charophytes and analyzed their evolutionary links.

MATERIALS AND METHODS

Dried material for *Sphagnum angustifolium* and *S. girgensohnii* were taken from herbarium at Department of Biology, Moscow State University. Total DNA was extracted from dry plants using the Nucleospin Plant Extraction Kit (Macherey-Nagel, Düren, Germany) according to the protocol of the manufacturer. For PCR amplification, the following degenerate primers were used: a forward primer Spha-TASP (5'-GGCGRTAWCCYACTGAGCTA-3') and reverse primer Spha-TASM (5'-TAGCTCAGGAGRGATAMMBMRA-3'). For PCR, 30 cycles were used with a melting temperature of 94 °C –3', and the next steps are as follows: an annealing temperature 94 °C –20", 65 °C –20", 58 °C –30", and an extending temperature of 72 °C followed by a final extension at 72 °C for 5'. PCR products were separated by electrophoresis of samples in a 1.5% agarose gel and purified using the Gel Extraction Kit (Qiagen, Hilden, Germany). For cloning, the PCR-amplified DNA bands isolated from gel were ligated into pGEM-T (Promega, Madison, WI, USA). The resulting clones were screened by length in 1,5% agarose gel. The plasmids were used as templates in sequencing reactions with an automated sequencer (Applied Biosystems, Foster City, CA, USA) 3730 DNA Analyzer with facilities of "Genom" (Moscow, Russia).

Sequences for comparative analysis were retrieved from NCBI (<http://www.ncbi.nlm.nih.gov/>), Phytozome (<http://www.phytozome.net>) and 1,000 Plant Transcriptome Project ("1KP") (<http://1kp-project.com/blast.html>). Sequence similarities were analysed by NCBI Blast at <http://blast.ncbi.nlm.nih.gov/BlastAlign.cgi>. The presence of open reading frames within retrieved sequences was analysed at <http://web.expasy.org/translate/>. The nucleic acid sequences and deduced amino acid sequences were analyzed and assembled using the NCBI. Conserved domains in the amino acid sequences SGS3 were identified using the CD-Search of the NCBI.

The sequences of SGS3 protein and TAS3 nucleotide sequences were aligned by MAFFT version 7 software (Katoh & Standley, 2014). The phylogenetic tree was constructed by the

Neighbor-Joining method with 1,000 bootstrap replications in MEGA7 (Kumar, Stecher & Tamura, 2016). The evolutionary distances were computed using the JTT matrix-based method and are in the units of the number of amino acid substitutions per site. The rate variation among sites was modeled with a gamma distribution (shape parameter = 1).

RESULTS

TAS3 loci in Bryophyta (classes Sphagnopsida and Takakiopsida)

It is commonly accepted that mosses of classes Sphagnopsida and Takakiopsida represent most basal lineages in Bryophyta (Shaw et al., 2010; Shaw, Szövényi & Shaw, 2011; Rosato et al., 2016). Previously, using primers, which have allowed us to detect pre-miR390 and TAS3 loci in Bryopsida and some other moss classes, we failed to identify pre-miR390 and TAS3 genes in genus *Sphagnum* (Krasnikova et al., 2013). However, a predicted sequence of pri-miR390 from *Sphagnum fallax* was recently reported (Xia, Xu & Meyers, 2017). This finding prompted us to re-evaluate the occurrence of TAS3-like loci in Sphagnopsida. To this end, we designed a new pair of degenerated PCR primers Spha-TASP and Spha-TASM, which differed from those used previously (Krasnikova et al., 2011; Krasnikova et al., 2013). As a positive control, we used plasmid DNA carrying cloned TAS3 gene of *Andreaea rupestris*, a representative of basal Bryophyta (Krasnikova et al., 2013). Like the positive control, two total DNA probes from *Sphagnum angustifolium* and *S. girgensohnii* gave a single main PCR product of the expected size (Fig. 1). Cloning and sequencing of these PCR fragments revealed two TAS3-like primary structures having 285 (*S. angustifolium*) and 292 (*S. girgensohnii*) bases in length and exhibiting 96% identity (e -value = $2e^{-131}$). We named these loci as Sphan-285 and Sphgi-292, (Fig. 2, Fig. S1 and Table 1). Despite that the degenerate miR390-mimicking primers were used for amplification of these loci, miR390 recognition sites in TAS3 species well corresponded to genomic loci of TAS3 extracted from genomic and transcriptomic data of other plants from genus *Sphagnum* (Fig. 2).

Peatmosses *S. angustifolium* and *S. girgensohnii* belong to subgenera *Cuspidata* and *Acutifolia*, respectively (Shaw et al., 2010; Shaw et al., 2016). To extend search for TAS3-like loci inside genus *Sphagnum* we performed bioinformatics analysis of the nucleotide sequences in databases available at NCBI (Sequence Read Archive) and Phytozome (version 12.1). Phytozome has recently released genome assembly of bog moss *S. fallax* (version 0.5). Bog moss belongs to subgenus *Cuspidata* and represents the most closely related moss to *S. angustifolium* (Shaw et al., 2016). BLASTN search at Phytozome allowed us to reveal a TAS3-like locus (supercontig super_37), which has 100% identity to the TAS3 locus of *S. angustifolium* sequenced in this study (Fig. S1 and Table 1). Unexpectedly, we found an additional TAS3-like locus in *S. fallax* (transcript Sphfalx0293s0011, supercontig super_293). This TAS3 locus in bog moss has 277 nucleotides in length and showed only a distant relation to the *S. angustifolium* TAS3 (Fig. 2, Fig. S1 and Table 1).

To further analyze Sphagnopsida TAS3-related loci, we used BLAST analysis of Sequence Read Archive (SRA), which is the NCBI database collecting sequence data obtained by the use of next generation sequence (NGS) technology. Assembly of sequence reads of *S. recurvum* (subgenus *Cuspidata*) retrieved by BLAST search using *S. fallax* sequences as

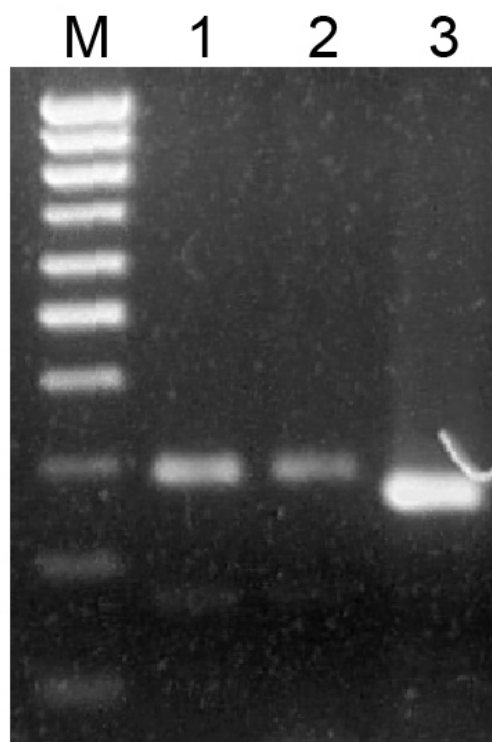


Figure 1 Analysis of PCR products in 1.5% agarose gel. Amplification of genomic DNA sequences flanked by miR390 and miR390* sites. PCR products were obtained on genomic DNAs with degenerate primers. *Sphagnum angustifolium* (1), *Sphagnum girgensohnii* (2), *Andreaea rupestris* (3). (M), DNA size markers including bands ranging from 100 bp to 1,000 bp with 100 bp step (Sibenzyme).

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queries revealed two TAS3 loci (Table 1). The first locus (Sphre-283) is 283 nucleotides in length and has 98% identity to Sphan-285. The second locus (Sphre-277) shows 98% identity to Sphfalx0293s0011 (Table 1, Fig. S1). These findings indicate that two distant TAS3 loci in species of a particular subgenus of genus *Sphagnum* are extremely similar.

We also analyzed the SRA database of subgenus *Sphagnum* (Shaw et al., 2010; Shaw et al., 2016). It was found that *S. magellanicum* belonging to this subgenus also encode two TAS3 loci called Sphma-285 (285 nt size) and Sphma-286 (286 nt size) (Fig. S1 and Table 1). Unlike *S. fallax* and *S. recurvum*, in *S. magellanicum* TAS3 loci are more similar, showing 86% identity (Fig. 2). Both Sphma-285 and Sphma-286 had 85% identity to Sphan-285 (Fig. 2). It was found that TAS3-like locus (Sphpa) from one more representative of subgenus *Sphagnum* (*S. palustre*) exhibited 98% identity to Sphma-285 (Fig. S1 and Table 1). The SRA database also contained sequence reads of two representatives from subgenus *Subsecunda* (Shaw et al., 2010; Shaw et al., 2016). Our BLAST analysis and subsequent assembly of retrieved reads revealed a single TAS3 locus in *S. cribrosum* (Sphcri, 291 nt size) showing 95% identity to Sphan-285 and 81% identity to Sphma-286 (Fig. 2, Fig. S1 and Table 1) and a partial TAS3-like sequence in *S. lescurii* (Fig. S1 and Table 1).

Analysis of the SRA database of *Takakia lepidozoides* (class Takakiopsida) allowed us to reveal only one TAS3-like sequence (Take-207) (Fig. S1 and Table 1). The same sequence

(A)

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Sphan-285          GCGGTAACCCCTCTGAGC TAAGTTAAAC-GGATAGGG-TTGTTGTTTGAAGTAGATT-----TGTGTGTTTTTAATGCTTTTT
S.fallax super 37 GCGGTAACCCCTCTGAGC TAAGTTAAAC-GGATAGGG-TTGTTGTTTGAAGTAGATT-----TGTGTGTTTTTAATGCTTTTT
Sphfalx0293s001  GCGGTAACCCCTCTGAGC TAAGTTAAAC-GGATAGGG-TTGTTGTTTGAAGTAGATT-----TGTGTGTTTTTAATGCTTTTT
Sphma-285         GCGGTAACCCCTCTGAGC TAAGTTGAGC-GGATAGGG-TTGTTGTTTGAAGTAGATT-----TGTGTGTTTTTAATGCTTTTT
Sphre-283        GCGGTAACCCCTCTGAGC TAAGTTGAGC-GGATAGGG-TTGTTGTTTGAAGTAGATT-----TGTGTGTTTTTAATGCTTTTT
Sphpa            GCGGTAACCCCTCTGAGC TAAGTTGAGC-GGATAGGG-TTGTTGTTTGAAGTAGATT-----TGTGTGTTTTTAATGCTTTTT
Sphgi-292        GCGGTAACCCCTCTGAGC TAAGTTAAACAGATAGGG-TTGTTGTTTGAAGTAGATTG--TGTGTGTTTTTAATGCTTTT
Sphcri           GCGGTAACCCCTCTGAGC TAAGTTAAACAGATAGGG-TTGTTGTTTGAAGTAGATTG--TGTGTGTTTTTAATGCTTTT
Sphma-286        GCGGTAACCCCTCTGAGC TAAATTTGAGGGAATAGGG-TTGAGCCTTGAAGTAGAAT-----TGTGAAATTTTAATGATTTT
Sphre-277        GCGGTAACCCCTCTGAGC TAA--TTTGGGATTAAGG-TTGATTTGCAACATTA-----TGTGATGTTTTT
Takakia          GCGGTAACCCCTCTGAGC TAA-----GCCAGTAG-----TGTGATGTTTTT
Folioceros       GCGGTAACCCCTCTGAGC TGA-----GAAAGAG-----TGTGATGTTTTT
*****

Sphan-285          AGTAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
S.fallax super 37 AGTAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
Sphfalx0293s001  AGTAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
Sphma-285         AGTAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
Sphre-283        AGTAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
Sphpa            AGTAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
Sphgi-292        AGAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAG-TATAAGCCCTTGT---TTCAGATATGAATT
Sphcri           AGAAGGAAGGAAAGCTGAATGTTAGGGTTAACATAATTA-----TTATGTTTTTAGAGAGATAAACCCTTGT---TTCAGATATGAATT
Sphma-286        AGTAAGGAAGGAAACATGAAAGTTACGGTTATCATAATTA-----TTATGTTTTTAG-TAAATGCCCTTGT---TTGAAATATGACTT
Sphre-277        AGTTAGTATGGGTTATGAATTTAGTATCTTTGTTTT-----TAAATCAACAT-TATTGATCATGTC---AAGAACATAATGT
Takakia          -----AGGTTGGTTGAGGG-----GGCACTAGG--ACACTCCCGGC-----CTGTGCGGAT
Folioceros       -GCAAGGTTGGGGTGGCTGGCGGGCGGCCCTTGTAAACGGGTTAAGCACCAAC--GGACGCCCTGGCAGCTCAGACGCCACCC
.....

Sphan-285          CTATAGCTTGAAGACATGACAAAC---ATGTTGTTTCGTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
S.fallax super 37 CTATAGCTTGAAGACATGACAAAC---ATGTTGTTTCGTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
Sphfalx0293s001  CTATAGCTTGAAGACATGACAAAC---ATGTTGTTTCGTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
Sphma-285         CTATAGCTTGAAGACATGATAAAC---ATGTTATTCCTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGTCAT
Sphre-283        CTATAGCTTGAAGACATGACAAAC---ATGTTGTTTCGTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
Sphpa            CTATAGCTTGAAGACATGATAAAC---ATGTTATTCCTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
Sphgi-292        CTATAGCTTGAAGACATGACAAAC---ATGTTGTTTCGTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
Sphcri           CTATAGCTTGAAGACATGACAAAC---ATGTTGTTTCCTCATCTCATGATCA-CCTGCAGACCTACCCCTTGAGACAAAATGTTTGACAT
Sphma-286        ATATAGCTTGAAGACATAATAAAA---AAGAAATTCATCATCTTACGACCT-CCTGCAGACCTACCCCTTGAGATAAATGTTTGACAT
Sphre-277        TGATCGAATTCAAAAGTCATTCTACTATGTTTGTACCCATCACCTCATCTCTCTCTGAGACCTACCCCTTGAGACAAAATGTTTGACAT
Takakia          ATGGTGGCC TAGGGTGTGATGAGT---GCTTTA--CCAGCACCTCACATTGGCCCA GCGGCTACCCCTTGAGTACAAGGGACTGCAACT
Folioceros       ACGGCTCCG TAGGGTGTGATGAGT---GCTTTA CCTAGCGCTCAGCCCTGGCGAG GCGGCTACCCCTTGAGTACACGGGCTGGGACAT
.....

Sphan-285          TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
S.fallax super 37 TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphfalx0293s001  TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphma-285         TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphre-283        TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphpa            TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphgi-292        TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphcri           TATTGCAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphma-286        TATTGAAACAT-CTGTCAAT TTAGTTATCACTCCTGAGCTA
Sphre-277        TATTGCAACAC-CTGTCAAT TCGATATCACTCCTGAGCTA
Takakia          CTTTGGCCATCTCTGTAAT TTGTTTATCACTCCTGAGCTA
Folioceros       CCCTGCACGGCCCTGTCGG-TTACGTATCACTCCTGAGCTA
.....

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(B)

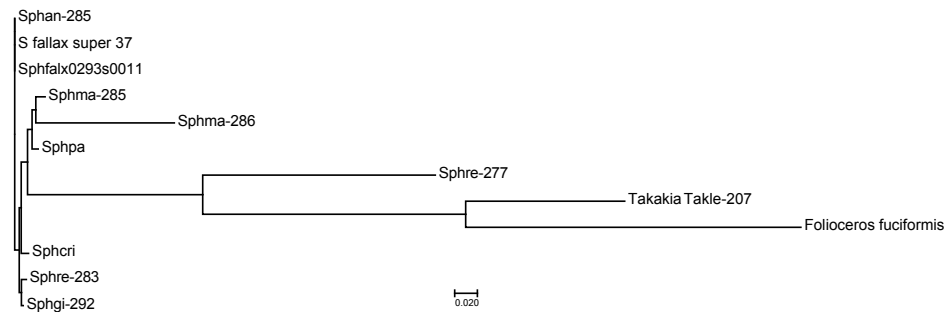


Figure 2 Analysis of TAS3 loci in genus *Sphagnum*. (A) Multiple sequence alignment of available nucleotide sequences of TAS3-like loci from mosses of genus *Sphagnum* along with TAS3 loci of *Takakia lepidozoides* and *Folioceros fuciformis*. Alignment was generated at MAFFT6 program. The miR390 target sites are in yellow, and putative tasiARF-a2 site is in green; tasiAP2 site is in blue. (B) The minimal evolution phylogenetic tree based on analysis of the aligned TAS3 genes from mosses of genus *Sphagnum*. This tree was generated according MAFFT6 program. For full plant names and accession numbers see Table 1.

Full-size DOI: 10.7717/peerj.4636/fig-2

Table 1 List of the putative TAS3 loci in Sphagnopsida and Takakiopsida.

Plant species	Locus name	Subgenus	Length	Sequence source
<i>Sphagnum angustifolium</i>	Sphan-285	<i>Cuspidata</i>	285 nts	MF682529
<i>S. girgensohnii</i>	Sphgi-292	<i>Acutifolia</i>	292 nts	MF682530
<i>S. fallax</i>	contig super_37	<i>Cuspidata</i>	285 nts	SRX2120232
<i>S. fallax</i>	Sphfalx0293s0011	<i>Cuspidata</i>	277 nts	Sphfalx0293s0011 ^a
<i>S. recurvum</i>	Sphre-283	<i>Cuspidata</i>	283 nts	SRX1513231
<i>S. recurvum</i>	Sphre-277	<i>Cuspidata</i>	277 nts	SRX1513231
<i>S. magellanicum</i>	Sphma-285	<i>Sphagnum</i>	285 nts	SRX2330962
<i>S. magellanicum</i>	Sphma-286	<i>Sphagnum</i>	286 nts	SRX2330962
<i>S. palustre</i>	Sphpa	<i>Sphagnum</i>	partial	SRX1516347
<i>S. cribrosum</i>	Sphcri	<i>Subsecunda</i>	291 nts	ERX443237
<i>S. lescurii</i>	Sphle	<i>Subsecunda</i>	partial	ERX337183
<i>Takakia lepidozoides</i>	Takle-207	Not applicable	207 nts	ERX2100030 SKQD-2076588 ^b

Notes.^aPHYTOZOME accession.^b1 KP accession (Xia, Xu & Meyers, 2017).

was revealed in a longer assembly which was found recently upon search of 1KP database (Xia, Xu & Meyers, 2017).

Since Takakiopsida and Sphagnopsida are most basal sister lines to all other Bryophyta (Shaw et al., 2010; Shaw, Szövényi & Shaw, 2011; Rosato et al., 2016; Puttick et al., 2018), it was very interesting to compare the structural organization of Takakiopsida and Sphagnopsida TAS3 loci with other classes of Bryophyta. Our previous detailed analysis of approximately 40 TAS3 loci in Bryophyta (Krasnikova et al., 2011; Krasnikova et al., 2013) showed that the general structure of moss TAS3 is similar in all taxa and fits the structural organization of *Physcomitrella patens* genes, comprising dual miR390 target sites on the 5' and 3' borders and internal monomeric tasiAP2 sequence followed by tasiARF sequence positioned in 20–30 bases. We revealed that phylogenetic tree of TAS3-like loci in Bryophyta showed clear subdivision of their sequences into two main clades (see Fig 5 in Krasnikova et al., 2013). The first group was formed by a cluster of sequences close to *P. patens* TAS3 species PpTAS3a, PpTAS3d, and PpTAS3f, and the second one—by those close to PpTAS3b, PpTAS3c, and PpTAS3e. The recent paper on the structure of TAS3 loci in lower land plants (Xia, Xu & Meyers, 2017) has shown the structure-functional basis for this phylogenetic subdivision. TAS3 species of the first group (PpTAS3a/PpTAS3d/PpTAS3f cluster) were shown to form class III of TAS3-like loci and contain, in addition to the previously reported tasiAP2 and tasiARF-a2 sequences, newly discovered tasiARF-a3 sequence positioned 5' according to tasiAP2 (Fig. 3). Among TAS3 species of basal Bryophyta, *Andreaea rupestris* locus 13-Aru (Krasnikova et al., 2013) belongs to class III (Fig. 3).

BLAST comparison of *T. lepidozoides* TAS3 with known Bryopsida loci showed that Takle-207 belongs to class II of TAS3 with typical positioning of tasiAP2 and tasiARF-a2 sequences (Fig. 2 and Fig. S1). On the other hand, none of Sphagnopsida TAS3-like sequences (Table 1) showed conventional internal structural organization

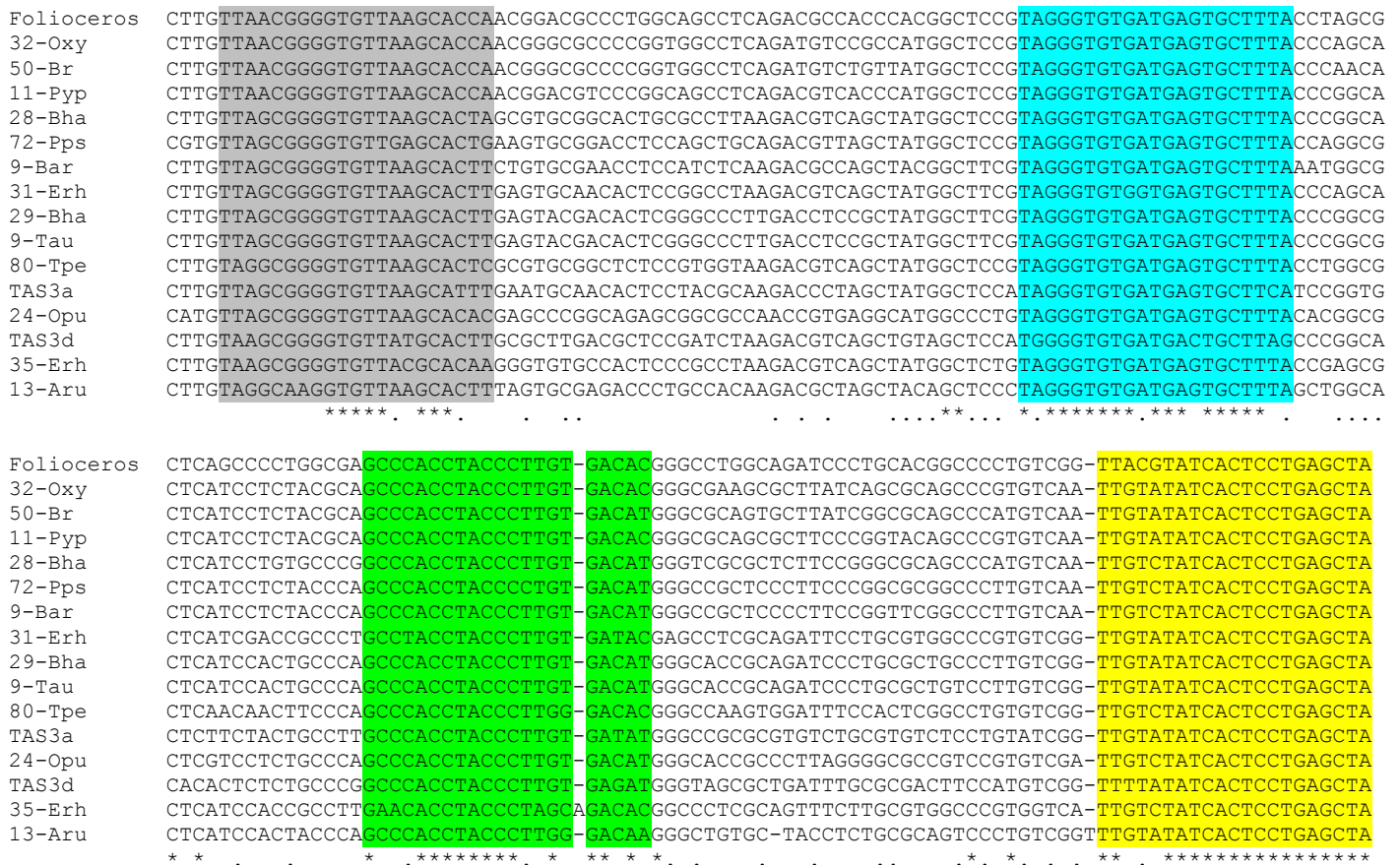


Figure 3 Multiple sequence alignment of selected available nucleotide sequences of class III TAS3-like loci from Bryopsida mosses along with TAS3 locus of *Folioceros fuciformis*. Alignment was generated at MAFFT6 program. The miR390 target sites are in yellow; putative tasiARF-a2 site is in green; tasiAP2 is in blue, and tasiARF-a3 is shaded. Note that all sequences are cut in the 5'-terminal TAS3 area to exclude non-aligned regions.

Full-size [DOI: 10.7717/peerj.4636/fig-3](https://doi.org/10.7717/peerj.4636/fig-3)

of the most moss TAS3 species. The only recognizable conserved site, except miR390-targeting regions, was identified as tasiARF-a2 sequence, which was found to be conserved between two very distant TAS3 loci in *S. fallax* and *S. recurvum* (Fig. 2). The mentioned above tasiARF sequences, tasiARF-a2 and tasiARF-a3, showed no sequence similarity suggesting their independent origins. These tasiRNAs were found to be formed from different strands of the TAS3 dsRNA intermediate and target different regions of ARF genes (Xia, Xu & Meyers, 2017). Inhibition of production of both tasiARF RNAs in *P. patens* resulted in obvious developmental defects exhibited, in particular, as alterations in gametophore initiation, protonemal branch determinacy and caulonemal differentiation (Plavskin et al., 2016).

Comparison of nucleotide sequences between TAS3 species of several moss classes revealed in many plants obvious similarity of nucleotide sequence blocks including tasiAP2 site and immediate upstream 21 bp block occurring in the same 21-bp-phase (Fig. 4A). We hypothesized that this sequence block may correspond to novel previously unrecognized

(A)

Andreaea1	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	43
Andreaea2	AAGACGCTAGCTACAGCTCC	TAGGGTGTGATGAGTGCTTTA	190
Tetraphis1	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	137
Tetraphis2	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	182
Tetraphis3	AAGACGCCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	440
Plagiomnium	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	315
Leucobryum	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	1006
Racomitrium	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	304
Philonotis	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	314
Dicranum	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	58
Encalypta	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	1009
Ceratodon	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	192
Niphotrichum	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	455
Funaria	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	250
Schwetschkeop	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	1632
Aulacomnium	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	278
Syntrichia	AGGACGTCAGCTATGGTCC	TAGGGTGTGATGAGTGCTTTA	154
Diphyscium1	AAGACGTCAGCTATGCTTC	TAGGGTGTGATGAGTGCTTTA	047
Diphyscium2	AAGACGTCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTAA	486
Pohlia	AAGACGCCAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTTA	38
Bryum	AAGACGCCAGCTACGGCTCC	TAGGGTGTGATGAGTGCTTTA	240
PHYSO TAS3A	AAGACCTAGCTATGGCTCC	TAGGGTGTGATGAGTGCTTCA	158

(B)

3' - UUCUGCaGUCGAUaCCGAGGc
caagacagttacggatcatctggacgtaccggaggtggtctagggc aagatgatagctacggctcat
Q D S Y G S S G R T G G G L G Q D D S Y G S - 28

3' - UUCUGCaGUCGAUaCCGAGGc
tcctctggctcgcaactggaggtggcctcggcc aagacgacagctatggttcctctggccgc
S S G R T G G G L G Q D D S Y G S S G R - 56

3' - UUCUGCaGUCGAUaCCGAGGc
ggtggcggcctcggta aagatgatagctacggctcctcaggacgtaccggggaaataca
G G G L G K D D S Y G S S G R T G G N T - 96

3' - UUCUGCaGUCGAUaCCGAGGc
cttgac aagatgattcgtatggatcctctggacaaactggcggtaacagcggatacggc
L G Q D D S Y G S S G Q T G G N S G Y G - 136

Figure 4 Novel potential ta-siRNA in Bryopsida plants. (A) Multiple sequence alignments of nucleotide sequence blocks including tasiAP2 site and preceding 21 bp site of putative ta-siRNA of *Andreaea rupestris* TAS3-like locus WOGB_2010369 with the corresponding transcript sequences of moss TAS3 loci. BLASTN was used at 1 KP blast site. For the complete TAS3 transcript sequences see [Xia, Xu & Meyers, 2017](#). The putative tasiAP2 site is in blue, and preceding putative ta-siRNA site is in violet. *Andreaea1* - *Andreaea rupestris* WOGB_2010369; *Andreaea2* - *Andreaea rupestris* WOGB_2002765; *Tetraphis1* - *Tetraphis pellucida* HVBQ_2019753; *Tetraphis2* - *Tetraphis pellucida* HVBQ_2011866; *Tetraphis3* - *Tetraphis pellucida* HVBQ_2005644; *Plagiomnium* - *Plagiomnium insigne* BGXB_2010105; *Leucobryum* - *Leucobryum glaucum* RGKI_2062694; *Racomitrium* - *Racomitrium varium* RDOO_2117129; *Philonotis* - *Philonotis fontana* ORKS_2058791; *Dicranum* - *Dicranum scoparium* NGTD_2078536; *Encalypta* - *Encalypta streptocarpa* KEFD_2058811; (continued on next page...)

Full-size  DOI: 10.7717/peerj.4636/fig-4

Figure 4 (...continued)

Ceratodon - *Ceratodon_purpureus* FFPD_2044193; *Niphotrichum* - *Niphotrichum_elongatum* ABCD_2000143; *Funaria* - *Funaria* sp. XWHK_2042016; *Schwetschkeop* - *Schwetschkeopsis_fabronia* IGUH_2166854; *Aulacomnium* - *Aulacomnium_heterostichum* WNGH_2088134; *Syntrichia* - *Syntrichia_princeps* GRKU_2074985; *Diphyscium1* - *Diphyscium_foliosum* AWOI_2069791; *Diphyscium2* - *Diphyscium_foliosum* AWOI_2006305; *Pohlia* - *Pohlia_nutans* GACA01023180; *Bryum* *argenteum* - Unigene33538 GCZP01053768; PHYSCO TAS3A - *Physcomitrella_patens* TAS3a BK005825. (B) The example target transcript sequence (lower case letters) from *Leucobryum_albidum* (VMXJ_2127900) is presented alongside with the predicted novel ta-siRNA shown in blue and above the transcript. Lower case letters in ta-siRNA indicate non-Watson-Crick pairing positions. Complementary mRNA sequences are in violet; conserved amino acid sequence signatures are in yellow. Numbers indicate codon positions of the target gene.

ta-siRNA in many moss species. This putative siRNA in its single-stranded form really presents in *P. patens* transcriptome as minus-sense 21-nucleotide ta-siRNA (see NCBI SRA accessions [SRX903096](#)–[SRX903105](#)) like tasiARF RNA ([Arif et al., 2012](#)). Thus, we speculated that novel hypothetical ta-siRNA might be produced from TAS3, and its minus-strand is complementary to uncharacterized well-conserved, protein-coding moss mRNAs which have homologs also in conifers and angiosperms ([Fig. 4B](#); [Fig. S2](#)).

TAS3 loci in Anthocerotophyta

Taking into account the finding of TAS3-like loci in classes Sphagnopsida and Takakiopsida and previously published data ([Krasnikova et al., 2013](#); [Xia, Xu & Meyers, 2017](#)), one can conclude that the only remaining blind-spot in land plants with respect to TAS3 is represented by phylum Anthocerotophyta. Relationships between liverworts, mosses and hornworts are still obscure. Moreover, the question remains which bryophyte phylum is a sister line to all other land plants ([Qiu, 2008](#); [Shaw, Szövényi & Shaw, 2011](#); [Harrison, 2017](#); [Puttick et al., 2018](#)). Recent molecular phylogenetic analysis, in which three bryophyte lineages were resolved, revealed that a clade with mosses and liverworts could form a sister group to the tracheophytes, whereas the hornworts is sister line to all other land plants ([Wickett et al., 2014](#)). However, analyses of the plastid genome sequences suggested another branching order of the phylogenetic tree, with hornworts rather than moss/liverwort clade being a sister group to tracheophytes ([Lewis, Mishler & Vilgalys, 1997](#); [Samigullin et al., 2002](#); [Ruhfel et al., 2014](#); [Lemieux, Otis & Turmel, 2016](#)). Moreover, some very recent nuclear gene comparisons also suggested that hornworts could be a sister clade to tracheophytes, and liverworts plus mosses might be closer to a common ancestor of land plants ([Rosato et al., 2016](#); [Bowman et al., 2017](#)). However, this ancestor could have more tracheophyte-like characteristics than Setaphyta (liverworts/mosses) due to secondary simplification in liverworts ([Puttick et al., 2018](#)).

Analysis of the SRA database of Anthocerotophyta revealed a putative TAS3-like sequence in *Folioceros_fuciformis* (family *Anthocerotaceae*). Unexpectedly, the discovered TAS3-like sequence (Folfu) was found to be 244 nucleotides in length and obviously similar to Bryophyta class III TAS3 species ([Fig. 3](#), [Fig. S3](#) and [Table 2](#)). The identity of Folfu to some moss TAS3 sequences exceeds 80% being therefore even higher than between some related Bryopsida species ([Fig. 3](#)). Thus these data clearly indicate a close relation of TAS3 in Anthocerotophyta to Bryophyta TAS3 (excepting Sphagnopsida).

Table 2 List of the putative TAS3 loci in Anthocerotophyta and Marchantiophyta.

Plant species	Class/subclass	Order	Length	Sequence source
<i>Folioceros fuciformis</i>	Anthocerotopsida/Anthocerotidae	Anthocerotales	244 nts	SRS2162762
<i>Marchantia polymorpha</i> 1-Mpo	Marchantiopsida/Marchantiidae	Marchantiales	256 nts	KC812742
<i>Marchantia emarginata</i>	Marchantiopsida/Marchantiidae	Marchantiales	262 nts	SRX1952816
<i>Conocephalum japonicum</i>	Marchantiopsida/Marchantiidae	Marchantiales	252 nts	SRX1952810
<i>Ricciocarpos natans</i>	Marchantiopsida/Marchantiidae	Marchantiales	235 nts	ERX337127
<i>Dumortiera hirsuta</i>	Marchantiopsida/Marchantiidae	Marchantiales	243 nts	SRX1126014
<i>Plagiochasma appendiculatum</i>	Marchantiopsida/Marchantiidae	Marchantiales	247 nts	SRX1741567
<i>Conocephalum conicum</i>	Marchantiopsida/Marchantiidae	Marchantiales	248 nts	ILBQ_2006554 ^a
<i>Lunularia cruciata</i>	Marchantiopsida/Marchantiidae	Lunulariales	220 nts	TXVB_2071521 ^a
<i>Marchantia paleacea</i>	Marchantiopsida/Marchantiidae	Marchantiales	257 nts	HMHL_2051051 ^a
<i>Metzgeria crassipilis</i>	Jungermannopsida/Metzgeriidae	Metzgeriales	226 nts	ERX337128
<i>Pellia endiviifolia</i>	Jungermannopsida/Pelliidae	Pelliales	192 nts	SRX726500

Notes.

^a1 KP accession (Xia, Xu & Meyers, 2017).

TAS3 loci in Marchantiophyta

Some of the recent molecular phylogenetic reconstructions suggested that Marchantiophyta species could represent a sister clade to all other land plants (see above). Therefore, finding and comparative analyses of TAS3 loci in this taxon represented a significant interest for understanding early events in TAS3 evolution. In contrast to class Marchantiopsida, where putative TAS3 and pre-miR390 loci were previously identified (Krasnikova *et al.*, 2013; Lin *et al.*, 2016; Tsuzuki *et al.*, 2016), for class Jungermannopsida only potential pre-miR390 loci were found in *Pellia endiviifolia* and *Harpanthus flotovianus* (Krasnikova *et al.*, 2013; Alaba *et al.*, 2015). Assuming that miR390 was found to be among eight most conserved miRNA species in land plants (Xia *et al.*, 2013; You *et al.*, 2017; Liu *et al.*, 2018), Jungermannopsida could be expected to encode TAS3 loci.

To detect new potential TAS3 loci, we performed BLAST analysis of the SRA database for species of class Jungermannopsida using *Marchantia polymorpha* TAS3 sequence (1-Mpo) as a query. Using this approach we revealed a set of reads and assembled a single TAS3-like locus (Pelen-192) for *Pellia endiviifolia* (192 nt size). In addition, TAS3 locus of 226 nucleotides in length was found in *Metzgeria crassipilis* (Metcr-226) (Fig. 5, Table 2, Fig. S3). The latter locus was also recently revealed in a search of 1KP database (Xia, Xu & Meyers, 2017).

TAS3 1-Mpo sequence was further used for BLAST analysis of other Marchantiopsida sequences available at the NCBI SRA database. As a result, we retrieved sequence reads and assembled five full-length TAS3-like sequences in *Plagiochasma appendiculatum* (Plaap-247), *Dumortiera hirsuta* (Dumhi-243), *Marchantia emarginata* (Marem-262), *Ricciocarpos natans* (Ricna-235) and *Conocephalum japonicum* (Conja-252) (Fig. 5, Table 2, Fig. S3). Recent bioinformatics analysis of 1KP database revealed three additional full-length TAS3-like sequences in *Conocephalum conicum*, *Lunularia cruciata* and *Marchantia paleacea*

(A)

```

Takakia      GCGGCTAACCTTCCTGAGCTAGCCAGTAGAG-----GGTGGGTTGAGGGGGGCAC----
Folioceros  GCGGTTATCCTTCCTGAGCTGAGAAAGAAGGCAAG-----GTTGGGGTGGCGTGGCGGGCGGCGCTTGT
1-Mpo       GACGGTATCCTTCCTGAGCTAAAAG-----ATGTAGC-----TTCTGTACATCTCACACGACATCTCATTTGAA
Marchantia  GACGGTATCCTTCCTGAGCTAGGAAGAAGGAG-----ATGTAGC-----TTCTGTACATCTCACACGACATCTCGTTGCA
Conocephal. Jap GACGGTATCCTTCCTGAGCTAGGAGGAAGAG-----ATGTAGCAGCGGGGACCTTACCTGTACATCTCACACGACATCTCGTTGAA
Conocephal. Con GACGGTATCCTTCCTGAGCTAGGAGGAAGAG-----ATGTATCAAGCGGTGCCTTACCTGTACATCTCACACGACATCTCGTTGAA
Dumortiera  GACGGTATCCTTCCTGAGCTAGGAGGAAGAG-----ATGTAGC-----TGTTCCTGTACATCTCACACGACATCTCATTTGTA
Plagiochasma GACGGTATCCTTCCTGAGCTAGGAGGAAGAG-----ATGTAGC-----TAGTGTACATCTCACACGACATCTCTCGTTTGTG
Ricciocarpos GACGGTATCCTTCCTGAGCTAGGAGGAAGAG-----AAAT-----CTGTACATCTCACACGACATCTTCTTTGGTA
Lunularia   GACGTTATCCTTCCTGAGCTAGGAAGAAGGAG-----ATGTAGC-----TTCTGTACATCTCACACGACATGTTTACTAGAA
Metzgeria   GACGTTATCCTTCCTGAGCTAGACAGACGTGACAGGATGTATC-----TGGCTACTGTGCATCTCACACTATGGGAGGGCTGGCGA
Pellia      GACGTTATCCGTCCTGAGCTAGATGATGTGG-----TTCTGTGCATCTCACACTACATGCGGCCT----
*..** ** * ..***** ..

Takakia      -----TAGGACACTTCCCGC-----CTTGTGCCGATATGGTGG-----C
Folioceros  TAACGGGGTGTAAAGCAC-----CAACGGAGCCCTGGC-----AGCCTCAGACGCCACCCACGGCTCC-----G
1-Mpo       TGTTCAAATCTTTAGTGACTGAATCGAATCTAAAGTTAATTTGACTTCAATAGAGACTAGTTTCCGGGAGAAACTGTCCAGTTAGCAG
Marchantia  TGTTCAAATCTCTCGGATGTAAGTCACATACAGAAGTTAATTTGACGGCAAGCGAGACACATGTCCGGGACGGACACCCCTGGTTAGCAT
Conocephalum TGTGCATGT-----CACAATTTAATTTGGATGATCAACGAGATAAATGTGTGGATGGACTCTCTGGTAGCAT
Conocephalum TGTGCATGT-----CACAATTTAATTTGGATGATCAACGAGATAAATGTGTGGATGGACTCTCTGGTAGCAT
Dumortiera  TGTTCAAAT-----CAAAATTAATTTGACACACACAGAGACAGAGCTGAGGGACAGAGACTTTTGTAGCAT
Plagiochasma TGTTCAAAT-----CAGAAGTCAATTTGATGACAAAAGAGACAGAGCTGACAGCGGACCGCTTTGGCCAGCAT
Ricciocarpos TGTTCAAAT-----CAGAATATGATTCGAAGATACACGAGATAAATTTGTGATACGGACACCTCTGCCTAAAAT
Lunularia   AATCTAG-----TAGTAAACAATTTGTGGGACAGGCTGCTGGTAAGCAC
Metzgeria   GCTGGAGCTGTT-----CTCAGCCTTTATGGTAGATACCCTTGGTCAGCTT
Pellia      -----CTTTGCTGGCAATGACAACATAGGTAGCAT

Takakia      TAGGGTGTGATGAGTGCTTTACCAG-CACCTCACATTTGCCCAAGCCGTCCTACCCTTGGTACAGGGGACTGCAACTCTTTGCGCCATCC
Folioceros  TAGGGTGTGATGAGTGCTTTACTAGCGCTCAGCCCTGGCGAGCCACCTACCTTGTGACAGGGCCTGGCAGATCCCTGCACGGCCC
1-Mpo       GAGGGTGTGATGAGTGCTTTACT-----GTTCCAGGATCCCCACCCCTCCT-----CCACTGCCTATTTCTAGGCTCGCG
Marchantia  GAGGGTGTGATGAGTGCTTTACCC-----GGCTCGGGTCCCGTCCCTTCC-----CCACTGCCTATGCTAGGCTCGCC
Conocephalum GAGGGTGTGATGAGTGCTTTACCA-----TACAGGGGTTCCTGTCCTTTCTTC-----CCGTGCATATGCTAGGTTCCGCC
Conocephalum TAGGGTGTGATGAGTGCTTTACCA-----TACAGGGGTTCCTGTCCTTTCTTC-----CCGTGCATATGCTAGGTTCCGCC
Dumortiera  GAGGGTGTGATGAGTGCTTTACCA-----GGCAAGGGTTCACGTCCTTTCTTC-----CCATGCTATGCTAGGTTCCGCC
Plagiochasma GAGGGTGTGATGAGTGCTTTACCA-----GGCCAGGGTTCCTGTCCTTTCTTC-----CCATGCTATGCTAGGTTCCGCC
Ricciocarpos GAGGGTGTGATGAGTGCTTTACTA-----GGCAGGGTTCACGTCCTTTCTTC-----CCACTGCATATGCTAGGTTCCGCC
Lunularia   TAGAGTGTGATAGTGCTTTACCA-----GACGGGGTTCCTGTCCTTTCTTC-----CCACTGCCTATGCTAGGTTCCGCC
Metzgeria   TAGGGTGTGATATGCTTTGACGT-----GGCAGTCTACCCCGCCCTGGG-----CAAATGCCTGCATCTTCTATCTG
Pellia      TAGGGTGTGATATGCTTTGACGT-----GACTTTCCCTCCTGCCCTGATTT-----GTTTTCCTGTGCTAGGCTAGTC
..***** ** ..* ..* ..*

Takakia      TTGTAATTTGTTTATCACTCCTGAGCTA
Folioceros  CTGTCCG--TTACGTATCACTCCTGAGCTA
1-Mpo       -TTAC--CTGCCTATCCCTCTTGAGCTA
Marchantia  -TGAC--CTGCCTATCCCTCTTGAGCTA
Conocephalum -TGAC--CTGCCTATCCCTCTTGAGCTA
Conocephalum -TGAC--CTGCCTATCCCTCTTGAGCTA
Dumortiera  -TGAC--CTGCCTATCCCTCTTGAGCTA
Plagiochasma -TGAC--CTGCCTATCCCTCTTGAGCTA
Ricciocarpos -TGAC--CTGCCTATCCCTCTTGAGCTA
Lunularia   -TGAC--CTGCCTATCCCTCTTGAGCTA
Metzgeria   -TTAA--TTTCCTATCCCTCTTGAGCTA
Pellia      TTGTA--TCTCCTATCCCTCTTGAGCTA
* ..* ..* ..* ..* ..*

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(B)

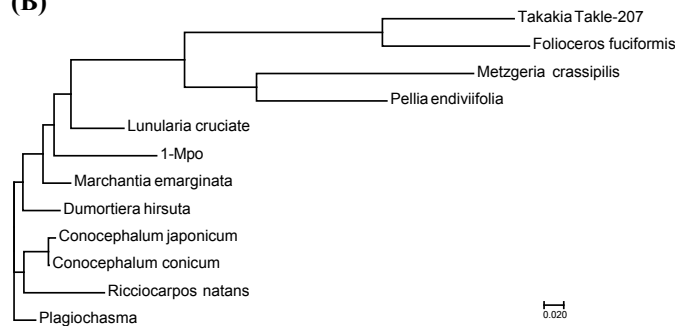


Figure 5 Analysis of TAS3 loci in Marchantiophyta plants. (A) Multiple sequence alignment of available nucleotide sequences of TAS3-like loci from Marchantiophyta plants along with TAS3 loci of *Takakia lepidozoides* and *Folioceros fuciformis*. Alignment was generated at MAFFT6 program. The miR390 target sites are in yellow, and putative tasiARF-a2 site is in green; tasiAP2 site is in blue. (B) The minimal evolution phylogenetic tree based on analysis of the aligned TAS3 genes from Marchantiophyta plants. This tree was generated according MAFFT6 program. For full plant names and accession numbers see Table 2.

Full-size DOI: 10.7717/peerj.4636/fig-5

(Xia, Xu & Meyers, 2017) (Table 2). Thus, totally 11 TAS3-like loci have been found in Marchantiophyta.

TAS6 loci in Bryophyta

Previous studies of *P. patens* revealed three novel non-coding PHAS loci (TAS6) which were located in rather close genomic proximity to PpTAS3 loci (PpTAS3a, PpTAS3d, and PpTAS3f) and expressed as common RNA precursors with these TAS3 species (Cho, Coruh & Axtell, 2012; Arif et al., 2012; Arif, Frank & Khraiweh, 2013). Moreover, miR529 and miR156 were suggested to influence accumulation of ta-siRNAs specific not only for TAS6, but also for PpTAS3a (Cho, Coruh & Axtell, 2012). We have found that localization of TAS6 loci close to TAS3 genes in common transcripts was not unique for *P. patens* (subclass Funariidae), since these loci were also found to be encoded by three other mosses of subclasses Bryidae and Dicranidae (Krasnikova et al., 2013).

For further search of the combined TAS6/TAS3 loci, we performed bioinformatics analysis of 1KP database. Although nucleotide sequences of miR156 and related miR529, as well as their recognition sites in RNA transcripts, are highly conserved among land plants (Morea et al., 2016; Axtell & Meyers, 2018), the internal sequences between dual miR156/miR529 recognition sites show little or no similarity even between different TAS6 loci of *P. patens* (Arif et al., 2012). So we used, as queries for BLAST search, the individual full-length TAS6/TAS3 loci including most characterized locus encoding PpTAS3a (Fig. 6), as well as those for PpTAS3d and PpTAS3f. First, it was found that in addition to four previously found Bryopsida species, encoding TAS6/TAS3 loci, these loci could be revealed in basal subclasses Timmiidae (*Timmia austriaca*) and Diphysciidae (*Diphyscium foliosum*) (Shaw, Szövényi & Shaw, 2011; Table 3, Fig. S4). List of TAS6/TAS3 loci in other moss subclasses was also significantly extended: we found 18 new loci in Bryidae, seven loci in Dicranidae and four loci in Funariidae (Table 3, Fig. S4). These novel loci showed recognizable but varying sequence similarities to the PpTAS3a-containing locus (Fig. 6). Second, most importantly, putative TAS6/TAS3 loci were revealed in four basal classes of Bryophyta, namely, Tetraphidiopsida, Polytrichopsida, Andreaeopsida and Takakiopsida (Table 3, Fig. S4). These novel loci had a similar organization to Bryopsida TAS6/TAS3 species (Fig. 6). However, no TAS6-specific sequence signatures were found in the vicinity of genomic *S. fallax* and *M. polymorpha* TAS3 loci upon analysis of the corresponding Phytozome genome contigs.

Phylogeny of SGS3 as a characteristic molecular component of TAS3 pathway

It was shown that some species of green algae could encode ancient types of dicer-like proteins, RDRs, and AGOs. On the other hand, no encoded SGS3 proteins were revealed for these algae (Zheng et al., 2015). Since SGS3 was found to be essential for production of tasiARF RNAs in moss *P. patens* (Plavskin et al., 2016), we performed sequence analyses to identify possible SGS3 genes in charophytes. For identification of SGS3 protein orthologs among land nonvascular plants and charophytes, we used as a query the most conserved region of *P. patens* SGS3 including short zinc binding zf-XS domain and RNA recognition

Table 3 List of the putative TAS6/TAS3 loci of Bryophyta in transcribed sequences found in 1 KP database.

Plant species	Class/subclass	Order	Length ^a and type	Sequence source
<i>Timmia austriaca</i>	Bryopsida/Timmiidae	Timmiales	TAS6/TAS3 (874 nts)	ZQRI-2061439 ZQRI-2063082
<i>Thuidium delicatulum</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (837 nts)	EEMJ-2003175
<i>Hypnum subimponens</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (823 nts)	LNSF-2068452
<i>Pseudotaxiphyllum elegans</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (1,590 nts)	QKQO-2009669
<i>Anomodon attenuatus</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (843 nts)	QMWB-2059873
<i>Anomodon rostratus</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (829 nts)	VBMM-2003482
<i>Schwetschkeopsis fabronia</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (854 nts)	IGUH-2166854
<i>Leucodon sciuroides</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (852 nts)	ZACW-2016434
<i>Fontinalis antipyretica</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (1,410 nts)	DHWX-2007057
<i>Rhytidiadelphus loreus</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (830 nts)	WSPM-2009782
<i>Rhynchostegium serrulatum</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (853 nts)	JADL-2047695
<i>Climacium dendroides</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (809 nts)	MIRS-2012325
<i>Calliergon cordifolium</i>	Bryopsida/Bryidae	Hypnales	TAS6 (95 nts)	TAVP-2006322
<i>Neckera douglasii</i>	Bryopsida/Bryidae	Hypnales	TAS6/TAS3 (839 nts)	TMAJ-2023603
<i>Plagiomnium insigne</i>	Bryopsida/Bryidae	Bryales	TAS6/TAS3 (914 nts)	BGXB-2010105
<i>Orthotrichum lyellii</i>	Bryopsida/Bryidae	Orthotrichales	TAS6 (192 nts)	CMEQ-2080784
<i>Hedwigia ciliata</i>	Bryopsida/Bryidae	Hedwigiales	TAS6/TAS3 (877 nts)	YWNF-2050742
<i>Philonotis fontana</i>	Bryopsida/Bryidae	Bartramiales	TAS6/TAS3 (893 nts)	ORKS-2058791
<i>Aulacomnium heterostichum</i>	Bryopsida/Bryidae	Rhizogoniales	TAS6/TAS3 (863 nts)	WNGH-2088134
<i>Scouleria aquatic</i>	Bryopsida/Dicranidae	Scouleriales	TAS6/TAS3 (partial)	BPSG-2088977
<i>Syntrichia princeps</i>	Bryopsida/Dicranidae	Pottiales	TAS6/TAS3 (partial)	GRKU-2074985
<i>Leucobryum glaucum</i>	Bryopsida/Dicranidae	Dicranales	TAS6/TAS3 (763 nts)	RGKI-2062694
<i>Leucobryum albidum</i>	Bryopsida/Dicranidae	Dicranales	TAS6/TAS3 (763 nts)	VMXJ-2128109
<i>Dicranum scoparium</i>	Bryopsida/Dicranidae	Dicranales	TAS6 (105 nts)	NGTD-2092412
<i>Ceratodon purpureus</i>	Bryopsida/Dicranidae	Pseudoditrichales	TAS6/TAS3 (1,121 nts)	FFPD-2005850 SRX2065999
<i>Racomitrium varium</i>	Bryopsida/Dicranidae	Grimmiales	TAS6/TAS3 (724 nts)	RDOO-2117129
<i>Physcomitrium_sp.</i>	Bryopsida/Funariidae	Funariales	TAS6 (partial)	YEPO-2071108
<i>Physcomitrium_sp.</i>	Bryopsida/Funariidae	Funariales	TAS6 (178 nts)	YEPO-2000016
<i>Physcomitrium_sp.</i>	Bryopsida/Funariidae	Funariales	TAS6/TAS3 (821 nts)	YEPO-2016361
<i>Encalypta streptocarpa</i>	Bryopsida/Funariidae	Encalyptales	TAS6/TAS3 (883 nts)	KEFD-2058811
<i>Diphyscium foliosum</i>	Bryopsida/Diphysciidae	Diphyscales	TAS6/TAS3 (832 nts)	AWOI-2069791
<i>Tetraphis pellucida</i>	Tetraphidopsida	Tetraphidales	TAS6 (partial)	HVBQ-2112923
<i>Atrichum angustatum</i>	Polytrichopsida	Polytrichales	TAS6/TAS3 (810 nts)	ZTHV-2082998
<i>Andreaea rupestris</i>	Andreaeopsida	Andreaeales	TAS6/TAS3 (869 nts)	WOGB-2010369
<i>Takakia lepidozoioides</i>	Takakiopsida	Takakiales	TAS6/TAS3 (1,040 nts)	SKQD-2076588

Notes.

^aThe length indicates total size of TAS6-TAS3 complex element (from the 5' miR529 target site in TAS6 to 3' miR390 target site in TAS3) or isolated TAS6 (between miR529 and miR156 target sites).

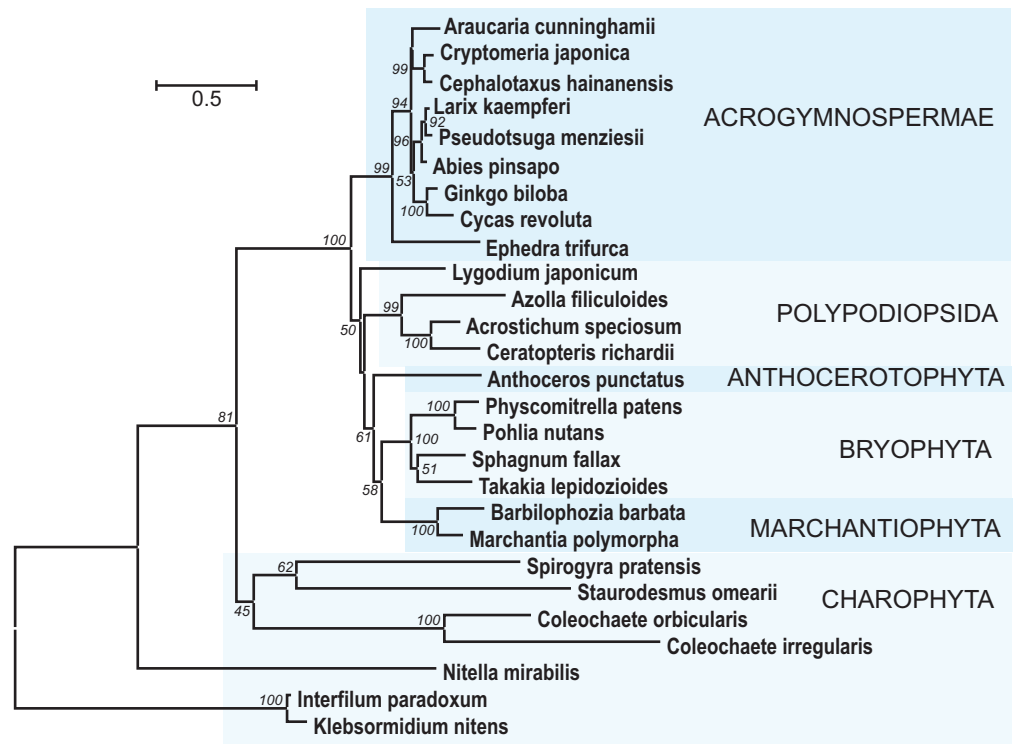


Figure 7 The phylogenetic tree constructed from conserved regions of SGS3 protein sequences from 27 selected streptophytes by the Neighbor-Joining method with 1000 bootstrap replications. There were a total of 419 positions in the final dataset. Evolutionary analyses were conducted in MEGA7 (Kumar, Stecher & Tamura, 2016). Bootstrap support values $\geq 50\%$ are shown. The evolutionary distances were computed using the JTT matrix-based method and are in the units of the number of amino acid substitutions per site. The rate variation among sites was modeled with a gamma distribution (shape parameter = 1). The tree was rooted at two Klebsormidiaceae charophytes. Accession numbers from NCBI or PHYTO-ZOME data banks see in Fig. S5.

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XS domain (Bateman, 2002; Zhang & Trudeau, 2008). Importantly, the short N-terminal zf-XS domain is characteristic for functional SGS3 proteins, since the XS domain-containing protein of *Selaginella moellendofii* lacking TAS-generating machinery (Banks et al., 2011) possesses no zf-XS domain upstream of XS domain and instead contains the C-terminal RING zf region (see NCBI accession XP_002979112). However, it should be noted that the lack of TAS3 pathway and SGS3 is not universal for lycophytes (Xia, Xu & Meyers, 2017).

In addition to class Bryopsida, SGS3 protein sequences were revealed for members of classes Marchantiopsida, Jungermanniopsida, Anthocerotopsida, Takakiopsida and Sphagnopsida (Fig. 7 and Fig. S5). Most importantly, search for the SGS3 coding sequences in transcriptomes of four charophyte classes (Klebsormidiophyceae) also revealed the SGS3-like proteins in representatives of all these taxa (Fig. 7, Figs. S5, S6). This observation was in agreement with the fact that SGS3-like coding sequence was found in the fully sequenced and annotated genome of *Klebsormidium nitens* (NCBI accession GAQ92898) (Hori et al.,

2014). Moreover, the characteristic motifs of land plant SGS3 proteins (Bateman, 2002) were revealed in the protein sequences from charophyte algae (Figs. S5, S6).

Importantly, in the dendrogram based on comparisons of 27 aligned SGS3 protein sequences, the position of charophytes (Fig. 7) nearly corresponded to the commonly accepted Viridiplantae phylogenetic tree (Shaw, Szövényi & Shaw, 2011; Delwiche & Cooper, 2015; Harrison, 2017), where class Zygnemophyceae (*Spirogyra pratensis*) was a sister group for all land plants. It has become clear that evolving the SGS3-like genes was not directly connected to the appearance of TAS loci in Viridiplantae, since Chlorophyta species, lacking SGS3, encode not only critical enzyme machinery including DCLs, RDRs, and AGOs (You et al., 2017), but also PHAS loci (Zheng et al., 2015). Despite our extensive searches, no SGS3 genes could be identified also in brown and red algae, and this is in agreement with previously published data on green algae (Zheng et al., 2015).

DISCUSSION

Our current analyses revealed previously undiscovered TAS3 loci in bryophytes from classes Takakiopsida, Sphagnopsida and Anthocerotopsida. In *Folioceros fuciformis* (family *Anthocerotaceae*) we found a TAS3-like sequence which is obviously similar to Bryopsida class III TAS3 species (Fig. 3), whereas Takakiopsida TAS3 locus showed relatedness to class II of TAS3 with typical positioning of tasiAP2 and tasiARF-a2 sequences and no tasiARF-a3 sites (Xia, Xu & Meyers, 2017). Unexpectedly, all predicted Sphagnopsida TAS3 loci showed no conventional internal structural organization of the moss class II and class III TAS3 species. Excepting miR390-targeting regions, the only recognizable conserved site was tasiARF-a2 sequence (Fig. 2).

It was shown that structural organization of Marchantiopsida and Jungermanniopsida TAS3 loci were quite similar, whereas Marchantiophyta TAS3 species were obviously different from those of Bryophyta. These loci were proposed to belong to TAS3 class I with species containing two conserved sequence blocks which presumably represent functional ta-siRNAs (Xia, Xu & Meyers, 2017). One of these blocks was found in the vicinity of the 3'-terminal miR390 binding site and corresponded to Bryopsida tasi-AP2 sequence (Krasnikova et al., 2013), whereas another one (tasiARF-a1), unique among lower land plants, was located closer to the 5'-terminal miR390 binding site in Marchantiopsida and Jungermanniopsida TAS3 (Tszuzuki et al., 2016; Xia, Xu & Meyers, 2017) (Fig. 5, Fig. S3).

Assuming occurrence of tasiARFs as potential products of TAS3 in all main lineages of land plants (Marchantiophyta, Anthocerotophyta and Bryophyta), recent paper proposed that the earliest function of TAS3 could contribute to the production of ta-siRNAs targeting ARF genes, and, since green algae encode no ARF genes, TAS3 likely appeared first in land plants (Xia, Xu & Meyers, 2017). However, extensive comparative sequence analysis showed that charophyte algae representing the sister group to all land plants (colonized terrestrial environments approximately 480 million years ago, see for references Fischer (2018) and Puttick et al. (2018)) could also encode ARF-like proteins including all sequence domains typical for bryophyte and angiosperm ARFs (Mutte et al., 2017). Moreover, our current data showed that TAS3-like loci are encoded by the representatives of all main

taxa among non-vascular plants. These observations suggest that the TAS3 evolution started in a common ancestor of land plants, likely belonging to a still unknown lineage of charophytes. Identification of the canonical motifs of land plant SGS3 in charophyte proteins (see above) indirectly supports this speculation. However, it should be kept in mind that evolving the SGS3-like genes could not be connected solely to the appearance of PHAS loci in Viridiplantae, since green algae and brown algae species were found to encode not only essential silencing machinery enzymes including DCLs, RDRs and AGOs, but also PHAS loci (Billoud *et al.*, 2014; Zheng *et al.*, 2015; Singh *et al.*, 2015; Dueck *et al.*, 2016; You *et al.*, 2017; Cock *et al.*, 2017). Finally, it can be proposed that the failure to identify charophyte TAS3 loci may be related to (i) the incompleteness of the available sequence data; (ii) evolving by charophytes the one-hit TAS3 genes (De Felippes *et al.*, 2017); or (iii) the use of miRNA species with sequences other than land plant miR390 for TAS precursor processing.

CONCLUSIONS

The current data on the structural organization of TAS3-like loci in all main classes of land non-vascular plants reveal three types of TAS3 loci, namely, (i) Bryopsida-like TAS3 (classes II and III, Xia, Xu & Meyers, 2017) found in Bryophyta plants (excepting Sphagnopsida) and Anthocerotopsida, (ii) Marchantiophyta-like TAS3 (class I, Xia, Xu & Meyers, 2017) and (iii) Sphagnopsida-like TAS3 (this paper). Clearly recognizable common ta-siRNAs is represented in these TAS3 types by tasiARF sequences. Occurrence of primitive SGS3 and ARF genes in charophytes (Mutte *et al.*, 2017 and this paper) supports the idea that TAS3-like genes first appeared in the hypothetical common precursor of land plants (Xia, Xu & Meyers, 2017) to regulate auxin signaling (Leyser, 2018).

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- Sergey Y. Morozov conceived and designed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Irina A. Milyutina conceived and designed the experiments, performed the experiments, contributed reagents/materials/analysis tools, approved the final draft.
- Tatiana N. Erokhina and Liudmila V. Ozerova performed the experiments, contributed reagents/materials/analysis tools, prepared figures and/or tables, approved the final draft.
- Alexey V. Troitsky and Andrey G. Solovyev conceived and designed the experiments, analyzed the data, authored or reviewed drafts of the paper, approved the final draft.

DNA Deposition

The following information was supplied regarding the deposition of DNA sequences:

New sequences described here are accessible via GenBank accession numbers [MF682529](#) and [MF682530](#).

Supplemental Information

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