Research Paper

Isopentenyltransferase-1 (IPT1) knockout in Physcomitrella together with phylogenetic analyses of IPTs provide insights into evolution of plant cytokinin biosynthesis

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Abstract

The moss Physcomitrella patens is part of an early divergent clade of land plants utilizing the plant hormone cytokinin for growth control. The rate-limiting step of cytokinin biosynthesis is mediated by isopentenyltransferases (IPTs), found in land plants either as adenylate-IPTs or as tRNA-IPTs. Although a dominant part of cytokinins in flowering plants are synthesized by adenylate-IPTs, the Physcomitrella genome only encodes homologues of tRNA-IPTs. This study therefore looked into the question of whether cytokinins in moss derive from tRNA exclusively. Targeted gene knockout of ipt1 (d|ipt1) along with localization studies revealed that the chloroplast-bound IPT1 was almost exclusively responsible for the A37 prenylation of tRNA in Physcomitrella. Ultra-performance liquid chromatography–tandem mass spectrometry (UPLC-MS/MS)-based cytokinin profiling demonstrated that the total amount of all free cytokinins in tissue was almost unaffected. However, the knockout plants showed increased levels of the N6-isopentenyladenine (iP)- and trans-zeatin (tZ)-type cytokinins, considered to provide active forms, while cis-zeatin (cZ)-type cytokinins were reduced. The data provide evidence for an additional and unexpected tRNA-independent cytokinin biosynthetic pathway in moss. Comprehensive phylogenetic analysis indicates a diversification of tRNA-IPT-like genes in bryophytes probably related to additional functions.

Key words: Bryophyte, cytokinin, isopentenyladenosine, isopentenyltransferases, moss, tRNA.

Introduction

Cytokinins (Cks) are N⁶-substituted adenine derivatives acting as phytohormones in various developmental processes in plants. Natural Cks represent a large group of phytohormones which can be divided into an aromatic and an isoprenoid group depending on the side chain coupled to the N⁶ of the adenine (Strnad, 1997; Mok and Mok, 2001). The isoprenoid...
Cks occur as four types, the 6-isopentenyladenine (iP), cis-zeatin (cZ), trans-zeatin (tZ), and dihydrozeatin (DHZ) types, varying in abundance as well as stereochemistry of the side chain hydroxyl group and/or in the saturation of the side chain in the case of DHZ types. Furthermore, Cks can be vastly modified, either at the N7, N9, or N5 position of the purine ring or at the terminal side chain hydroxyl group. Major modifications are glycosylation, aminoacylation, and phosphorylation (Mok and Mok, 2001). The different types and conjugates of Cks can differ greatly in their biological activity and abundance (Sakakibara, 2006).

Cytokinin biosynthesis

The rate-limiting step in biosynthesis of isoprenoid Cks in plants is catalysed by isopentenyltransferases (IPTs), adding an isoprenoid side chain to the N6- amino group of an adenine nucleotide. Depending on the nature of the nucleotide substrate, two different forms of IPTs are known in plants (Fig. 1) (Kamada-Nobusada and Sakakibara, 2009). Adenylate-IPTs use either ATP, ADP, or AMP as a substrate and are encoded in multiple copies in all flowering plants (Kakimoto, 2001; Takei et al., 2001). The adenylyl-IPTs of Arabidopsis are mainly responsible for the synthesis of iP- and tZ-type Cks (Miyawaki et al., 2006). Interestingly, the first adenylyl-IPT to be described was cloned from the plant pathogen Agrobacterium tumefaciens and was shown to be the factor driving Agrobacterium-induced tumorigenesis in plant root tissues (Barry et al., 1984; Lichtenstein et al., 1984), and proved to be functional in the moss Physcomitrella patens (Reutter et al., 1998).

In budding bioassays, the free bases iP, tZ, and benzyladenine (BA) are the most effective, while their corresponding ribosides are less active. Strikingly, in the same assays, none of

Arabidopsis, tRNA-IPTs are exclusively responsible for the synthesis of cZ-type Cks (Miyawaki et al., 2006).

Cytokinins in Physcomitrella

The last common ancestor (LCA) of mosses and flowering plants lived ~500 million years ago (MYA) (Lang et al., 2010). Consequently, development of the moss Physcomitrella is mainly controlled by the evolutionarily old phytohormones auxin, Ck, and abscisic acid (for reviews, see Decker et al., 2006; Cho et al., 2009; von Schwartzzenberg, 2009). Physcomitrella has been established as a model system in evo-devo research and plant molecular genetics due to the availability of a fully sequenced genome as well as exceptionally high rates of homologous recombination enabling reverse genetics (Schaefer and Zryd, 1997; Kamisugi et al., 2006; Lang et al., 2008; Rensing et al., 2008; Prigge and Bezanilla, 2010). Thus the evolution of phytohormone signalling and metabolism can be studied in comparative analyses utilizing moss (Vandenbussche et al., 2007; Paponov et al., 2009; Kopecka et al., 2013).

Physcomitrella is the first bryophyte with a reported Ck profile (von Schwartzzenberg et al., 2007). In protonema from liquid cultures, the cZ-type cytokinins are dominant, with cZROG (cis-zeatin riboside O-glucoside) accounting for 80% of the overall Ck pool. The other Ck types, iP, tZ, and DHZ, were also detectable at intracellular concentrations in the picomolar range. In the culture medium, the iP conjugates were found to be the most abundant type, with 6-isopentenyladenosine-5′-monophosphate (iPR5′MP) as a main constituent.

Cks are of particular importance for the progression of the life cycle of mosses as they induce the production of three-faceted apical cells (buds), which later develop into the leafy shoots (Cove and Ashton, 1984). An external application of Cks results in a strong overproduction of buds (Reski and Abel, 1985), and internal Ck accumulation precedes bud formation (Schulz et al., 2000). However, not all Cks exert the same hormonal activity when applied to protonema cultures. In budding bioassays, the free bases iP, tZ, and benzyladenine (BA) are the most effective, while their corresponding ribosides are less active. Strikingly, in the same assays, none of

![Fig. 1. Simplified scheme of the two alternative cytokinin (Ck) biosynthetic pathways in plants. (a) Prenylation of UNN-decoding tRNAs at adenine (A37) by tRNA-IPTs and subsequent release of cytokinin nucleotides by tRNA degradation. (b) Direct synthesis by adenylyl-IPTs. DMAPP, dimethylallyl diphosphate; HMBDP, hydroxymethylbutenyl diphosphate. Data taken from Sakakibara (2006). (This figure is available in colour at JXB online.)](https://example.com/fig1.png)
the tested cZ conjugates had a bud-inducing effect, leading to the conclusion that cZ-type Cks are not contributing to bud induction (von Schwartzenberg et al., 2007). This apparent lack of hormonal activity stands in contrast to the dominant abundance of cZ-type Cks in Physcomitrella tissue.

The full genomic sequence of Physcomitrella (Rensing et al., 2008; Zimmer et al., 2013) reveals seven ipt genes, which are at first sight all to be considered as tRNA-IPTs based on their sequence homology to other plant tRNA-IPTs (Sakakibara, 2006; Frebort et al., 2011). IPT1 has been studied by Yevdakova and von Schwartzenberg (2007). A yeast complementation assay confirmed the IPT1 protein to be a functional tRNA-IPT. Findings such as the dominant occurrence of cZ-type Cks in tRNA and whole tissue extracts of Physcomitrella, together with the sequence homology of all IPTs to known, functional tRNA-IPTs, led to the working hypothesis that the tRNA biosynthetic pathway is of great importance in Physcomitrella (Yevdakova et al., 2008; von Schwartzenberg, 2009). The remaining six IPTs have only rarely been studied, but Patil and Nicander (2013) have shown recently that IPT4 and IPT5 also possess tRNA-IPT function.

In this study, insights are provided into the origins of Cks in Physcomitrella focusing on IPT1 by the generation of targeted gene knockout plants. The results of the phenotypic analysis, detailed analysis of free and tRNA-bound Cks, and a phylogeny of IPT proteins reveal unexpected, multiple origins of Cks and indicate that the in vivo function of annotated tRNA-IPTs has to be re-assessed in Physcomitrella and possibly other organisms.

Materials and methods

Plant material and growth conditions

The Physcomitrella patens (Hedw.) Bruch & Schimp wild type used in this study was originally collected from Grandsen Wood, Huntingdonshire, UK (1968) and is the clone that was sequenced. Under standard growth conditions, strains were kept at 25 °C in white light (Phillips TML, Hamburg, Germany) at 100 μmol m⁻² s⁻¹ under a light:dark cycle of 16:8 h. Tissue of Physcomitrella grown in Petri dishes on ABCTV agar medium according to Wang et al. (1968) was grown in Huntingdonshire, UK (1968) and is the clone that was sequenced. The gene models of the moss IPT family were manually curated based on cDNA and expressed sequence tag (EST) evidence using the genome browser and annotation service of the moss model organism database (http://www.cosmos.org; Lang et al., 2005). The complete open reading frame of PpIPT1 (GenBank accession no. EF512463.1; curated cosmos.org gene model Pp1s96_115V6_lindner.1; excluding the stop codon) was amplified from pFL61_PpIPT1 (Yevdakova and von Schwartzenberg, 2007) with the primers ipt1_for(AvrII), CGCGCCCTAGGATGTTGATTTGCGATTTGAG; and ipt1_rev(BamHI), ATGGGGATCCTGAGAAGTCGCTCAAGGATG, digested with AvrII and BamHI, and cloned into the pLNU vector (http://www.dna-cloning.com/vectors/Vectors_with_markers/plNU-GFP.gb) 5’ of the green fluorescent protein (GFP)-encoding sequence (pLNU-GFP). PpIPT1 vector card is given in Supplementary Fig. S1 available at JXB online). The plasmid was then transfected into Physcomitrella protoplasts by polyethylene glycol (PEG)-mediated transfer according to Schaefer et al. (1991). GFP fluorescence in transiently transformed protoplasts was monitored 5 d after transfection using a TCS SPE confocal laser scanning microscope, and images were acquired with the LAS AF lite software (both Leica, Wetzlar, Germany).

Generation and characterization of ipt1 mutants

IPT1-deficient mutants were generated by gene targeting. The gene-disrupting vector pBNR_PpIPT1, based on pBNR (Schaefer et al., 2010), carries two ~1000 bp 5’ and 3’ genomic fragments of the native ipt1 locus (deletion of the Pp1 assembly region scaffold_96:726668...728288), flanking a 3SS:nptII resistance-mediating cassette. Recombinant transformants were obtained in three cycles of selection on ABCNTV agar medium containing G418 and were characterized by PCR and reverse transcription–PCR (RT–PCR) for ipt1 gene disruption as well as loss of transcript (Supplementary Fig. S3 at JXB online). The selected mutant moss lines are named d|ipt1#49, dipt1#10, and dipt1#13, and have been deposited at the International Moss Stock Center (http://www.moss-stock-center.org/) with the respective IMSC accession numbers 40697, 40698, and 40699.

Cytokinin measurements in tissue, medium, and tRNA samples

Physcomitrella wild type and three independent d|ipt1 mutants were grown in liquid cultures under standard conditions as previously described in von Schwartzenberg et al. (2007). Prior to preparation of cultures, the tissue used for inoculation was washed with an excess of fresh medium. For sampling, 200 ml aliquots were taken after 22 d of culture. Tissue and medium from the same sample were separated by filtration. The tissue was immediately frozen in liquid nitrogen, and medium and tissue were stored at −20 °C for a maximum of 2 weeks prior to freeze-drying.

For analysis of endogenous Cks, extraction and purification were performed according to the method described by Novák et al. (2003). The samples were purified using a combined cation (SCX-cartridge), anion (DEAE-Sephadex-C18-cartridge) exchanger and immunofinity chromatography (IAC) based on a wide range of specific monoclonal antibodies against Cks. The levels of Cks were quantified by ultra-performance liquid chromatography–electrospray tandem mass spectrometry (UPLC-MS/MS) using stable isotope-labelled internal standards as a reference (Novák et al., 2008).

In order to obtain sufficient quantities of tissue for Ck determination in tRNA, cultures independent from those used for free Ck profiling were grown under the same conditions for 22 d.

Extraction and purification of tRNA was performed according to a protocol described by Maass and Klämbt (1981), including modifications described by Stirk et al. (2011). Aliquots of 60 g of tissue were hydrolysed with 2 M KOH overnight and dephosphorylated by alkaline phosphatase. Internal standards were added and the samples were purified by IAC based on a wide range of specific monoclonal antibodies against Cks (Faiss et al., 1997). Cks in these samples were quantified by UPLC-MS/MS as described above.

Computational and phylogenetic analysis of the IPT family

Members of the IPT family in the genomes of 12 selected sequenced green plants and algae (Arabidopsis thaliana, Chlamydomonas reinhardtii, Chlorella variabilis, Cyanidioschyzon merolae, Micromonas pusilla, Micromonas sp., Oryza sativa, Ostreococcus lucimarinus,
Ostreococcus tauri, Selaginella moellendorffii, Volvox carteri, and Zea mays) were identified using protein clustering as described previously (Bartels et al., 2010; Lang et al., 2010). The resulting candidate list was collated with the clusters in the Phytozome plant gene family database (Goodstein et al., 2012) and previous descriptions of the IPT family in Arabidopsis, rice, and maize. To provide further taxonomic coverage, additional members of the IPT family were added based on their InterPro protein domain annotation (Hunter et al., 2012). Using the InterPro database web interface, proteins containing the IPT protein domain (PF01745; IPR002627) from selected cyanobacteria, Alveolates, Euglenozoa, Stramenopiles, and Agrobacterium tumefaciens were included into the candidate set. Sequence data are available as a supplementary file at JXB online. To derive a high-quality alignment, the manually curated alignment of the TRIT1 tRNA isopentenyltransferase family from the TreeFam database (TF315069) was used as a starting point for profile alignment using T-Coffee. 8.14 (Notredame et al., 2000). Based on the resulting full alignment (5517 amino acid columns), a Neighbor–Joining (NJ) guide tree was calculated using the Sorensid matrix (Sonnhammer and Hollich, 2005) implemented in a modified version of QuickTree and MrBayes (Ronquist and Huelsenbeck, 2003). MrBayes was run with parameter settings lset nst=6 enies using QuickTree and MrBayes (2002) applying bootstrap resampling with 1000 replicates. The resulting tree was used to reorder the alignment and manually curate and clip it using Jalview (Clamp et al., 2004) by removing phylogenetically uninformative sites and reducing the full alignment to a clipped and reduced alignment spanning 203 residues and comprising 97 proteins. A full list of all IPT family members in this curated alignment is provided in Supplementary Table S1 at JXB online. The resulting clipped protein alignment was used to infer the IPT gene phylogenies using QuickTree and MrBayes (Ronquist and Huelsenbeck, 2003). MrBayes was run with parameter settings lset nst=6 rates=invgamma Ngammaracat=4; prset aamodelpr=mixed; mcmodel cpmodel=cata; mcmctol=0.01 stoprule=yes nnruns=2 nchains=128 samplefreq=1000 temp=0.2 starttree=random redburnin=yes burninfrac=0.25 savebrels=yes.

Results

Phylogenetic analyses reveal a complex evolutionary history of IPTs

In order to resolve the evolutionary past of the moss IPTs and to test their tRNA-IPT ancestry, a comprehensive phylogenetic study was performed considering multiple databases and genomes that comprises protein sequences from 22 bacterial (Actinobacteria, Cyanobacteria, and Proteobacteria) and 23 eukaryotic species (Amoebozoa, Fungi, Metazoa, Rhodophyta, Stramenopiles, and Viridiplantae). The resulting taxonomic distribution of IPT functionality across the different kingdoms of life (Fig. 2) raises the question of their evolutionary history.

As previously reported (e.g. Frebort et al., 2011), tRNA-IPTs are found in all major clades of bacteria and eukaryotes, but are absent from Archaea. In contrast, adenylate-IPTs follow a more fragmented distribution. The functionally confirmed adenylate-IPTs are found in flowering plants (Kakimoto, 2001; Miyawaki et al., 2006; Sakamoto et al., 2006; Brugiere et al., 2008), the slime mould Dictyostelium discoideum (Taya et al., 1978), and several bacterial plant pathogens or symbionts (e.g. Akiyoshi et al., 1985). The present phylogenetic analysis suggests a complex evolutionary history of IPTs. Figure 2 depicts the relationships of IPTs as a midpoint-rooted phylogram with branch support values and colour-coding of branches based on proposed functional IPT classes. Besides the tRNA-IPTs, whose function has been initially identified as prenylating certain tRNAs in order to increase translational precision (Konevega et al., 2006), the flowering plant adenylate-IPTs are thought to use ADP or ATP as substrate, while the slime mould and bacterial forms seem to prefer AMP. Accordingly, Kakimoto (2003) classified IPTs into functional groups of DMAPP:tRNA-IPTs (here tRNA-IPT, Fig. 2; blue branches), DMAPP:AMP-IPTs (here AMP-IPTs, red branches), and DMAPP:ADP/ATP-IPTs (here ADT/ATP-IPTs, green branches).

The absence of adenylate-IPT homologues in Physcomitrella gave rise to the hypothesis that in early divergent land plants CK biosynthesis is mediated exclusively by tRNA prenylation and subsequent degradation. A functional characterization of Physcomitrella IPT1 was performed in order to test this hypothesis.

IPT1 is localized in chloroplasts

Based on sequence analyses, the Physcomitrella IPT1 protein was predicted to be targeted to chloroplasts (Yevdakova and von Schwartzenberg, 2007). For experimental validation of the chloroplast localization of IPT1, a IPT1::GFP fusion protein was transiently expressed in Physcomitrella protoplasts. The subsequent CLSM-based imaging is consistent with a subcellular localization of IPT1 in plastids (Fig. 3), whereas the control transfection leading to an expression of GFP alone showed fluorescence signals in the cytoplasm. It was observed that the IPT1::GFP-mediated fluorescence was not evenly distributed within the chloroplast. The confinement of the GFP signal to the chloroplast has been confirmed by spinning disc CLSM for individual chloroplasts (Supplementary Fig. S2 at JXB online).

Characterization of d|ipt1 knockout mutants

In order to draw conclusions on the function of IPT1 and its relevance for CK biosynthesis, targeted knockout mutants, dipt1 plants, were generated taking advantage of the high frequency of homologous recombination in Physcomitrella. After selection of stable recombinant plants, the haploid status of the selected strains (#9, #10, and #13) was confirmed by flow cytometry (data not shown) in order to exclude that protoplast fusion had occurred (Schween et al., 2003, 2005). PCR on genomic DNA from the dipt1 plants and subsequent sequencing showed that the expected gene replacement, namely the partial ipt1 deletion and full integration of the 35S::nptII resistance cassette, had occurred at the ipt1 locus (Supplementary Fig. S3a at JXB online). Further, RT–PCR-based analyses confirmed the absence of ipt1 transcript in the selected mutants. In order to strengthen the results and to minimize unlikely effects of eventual additional random ectopic integrations, analyses were performed with three independent dipt1 plants; the results for one of the mutants (dipt1#9) is presented in the following. The results for the two additional independent mutants confirming the findings are given in the supplementary data available at JXB online.
Fig. 2. IPT phylogeny. Branch lengths and support in the form of posterior probabilities (presented at internal nodes and branch thickness) were inferred using MrBayes (Ronquist and Huelsenbeck, 2003) based on a manually curated T-Coffee (Notredame et al., 2000) protein alignment comprising 97 representative members of the IPT family. Confirmed IPT functions are designated with yellow dots: DMAPP:tRNA-IPT references, MiaA_Ecoli (Caillet and Droogmans, 1988), TRIT1_HUMAN (Golovko et al., 2000), MOD5_YEAST (Dihanich et al., 1987), AtIPT2 (Golovko et al., 2002), AtIPT9 (Miyawaki et al., 2006), PpIPT1 (Yevdakova and von Schwartzenberg, 2007), PpIPT3,5 (Patil and Nicander, 2013). DMAPP:AMP-IPT references, IPT_DCDI (Taya et al., 1978), IPTZ_AGRT7 (Akiyoshi et al., 1985), IPTZ_AGTT5 (Beaty et al., 1986); DMAPP:ADT/ATP-IPT references, AtIPT1-4 (Kakimoto, 2001), AtIPT3,5,6,7,8 (Miyawaki et al., 2006; Sakakibara, 2006), OsIPT1,3 (Sakamoto et al., 2006), ZmIPT2 (Smijers et al., 2008), Proposed (homology-based) functionality for the clades is given by the colouring of the branches. Gene identifier and species names are available in Supplementary Table S1 at JXB online.
Phenotype and cytokinin bioassays

The d|ipt1 plants expressed a morphological phenotype at the level of protonema. Mutant chloronema and caulonema cells were smaller compared with the wild type and showed a higher degree of branching (Fig. 4a, b). Compared with the wild type, d|ipt1 colonies revealed a reduced diameter (Fig. 4e, f). d|ipt1 mutant lines were not affected in bud formation; that is, mutant cultures grown on solid, as well as in liquid medium still formed buds at approximately the same time point and with frequencies comparable with the wild type (Fig. 4c, d). Gametophore development of the d|ipt1 mutant strains was also not significantly altered (Fig. 4f).

In order to check whether the phenotype observed for d|ipt1 mutant protonema could be reversed by externally applied Cks, mutants were treated with different concentrations of iP and BA (20–500 nM). During 2 weeks of microscopic observation, the phenotype of reduced colony growth persisted and budding was induced on the mutant protonema to the same extent as on the wild type (Supplementary Fig. S4 at JXB online).

Levels of cytokinins in d|ipt1 mutants

In order to assess the impact of the deletion of ipt1 on levels of free and tRNA-bound Cks, they were quantified in media and tissue of liquid cultures after 22 d of cultivation by UPLC-MS/MS.

tRNA-bound cytokinins

Assuming that in Physcomitrella tRNA is a major source of Cks, the amount of Ck ribosides in dephosphorylated tRNA hydrolysates was determined. cZR had already been reported as the dominant Ck type in tRNA of the wild type (Yevdakova et al., 2008), correlating well with the dominance of free cZ-type Cks in tissue. tRNA extracts of d|ipt1 mutants revealed strongly reduced contents of all analysed Ck ribosides (Fig. 5a). tZR and DHZR dropped to levels below the limit of detection. The more abundant cZR and iPR were reduced to <1% compared with the wild type, thus revealing the essential importance of IPT1 for tRNA A37 prenylation.

Free cytokinins in tissue

In order to uncover the contribution of tRNA-bound Cks to the overall Ck production in Physcomitrella, free Cks were determined in tissues of the wild type and d|ipt1 mutants. As the methanolic extraction and the applied purification and quantification protocol are specific for low molecular weight Cks, interference by tRNA-bound Cks is unlikely. UPLC-MS/MS analysis revealed that levels of cZ-type Cks were reduced from 537 pmol g–1 dry weight (DW) in the wild type to 132 pmol g–1 DW in d|ipt1 mutants. Furthermore, levels of the less abundant DHZ-type Cks (15 pmol g–1 DW in the wild type) are reduced to only 5 pmol g–1 DW in d|ipt1 plants.

Surprisingly, a different situation was monitored for the levels of iP- and tZ-type Cks. In contrast to cZ- and DHZ-type Cks, the levels of free iP- and tZ-type Cks were strongly increased in the tissues of d|ipt1 plants (Fig. 5b). For example, iPR5’MP, which at 56 pmol g–1 DW was already the most abundant iP metabolite in the wild type, was increased...
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~4-fold to levels up to 233 pmol g⁻¹ DW in dipt1 mutants (Supplementary Table S2 at *JXB* online).

Extracellular cytokinins

Previously, the iP-type Cks were found to be highly abundant in the culture media of *Physcomitrella* (*Reutter et al.*, 1998; *Schulz et al.*, 2001; *von Schwartzzenberg et al.*, 2007). In the dipt1 plants, the levels of extracellular iP and iPR₅MP were determined to be at least 15-fold higher than in the wild type. In the mutants, the concentration of iPR₅MP in the medium during 22 d of cultivation accumulated to a concentration of 3697 pM compared with 242 pM in the wild type (Supplementary Table S3 at *JXB* online).

In summary, the results of Ck analyses clearly demonstrated that the knockout of *ipt1* does not result in a general decrease of all Cks, but leads to a drastic differential alteration of the levels of the different Ck types.

**Discussion**

Apart from its general utility as a plant evo-devo model (*Lang et al.*, 2008; *Prigge and Bezanilla*, 2010), two striking aspects are of particular interest in studying Ck biosynthesis in the early divergent land plant *P. patens*: (i) it is the most basal member of the green lineage with a sequenced genome that encodes major elements of Ck biosynthesis, metabolism, degradation, and signalling (*Rensing et al.*, 2008; *Pils and Heyl*, 2009); and (ii) the moss genes encoding the first step in Ck biosynthesis exclusively represent homologues to tRNA-IPTs, a type of IPTs that in flowering plants is considered to be only of minor importance to biosynthesis of active free Cks (*Miyawaki et al.*, 2006). The origin of these proteins and especially the function of IPT1 are the key points of this study.

*Physcomitrella has a unique position in the IPT phylogeny*

Given the topology of the tree and the taxonomic distribution generated here (Fig. 2), it can be concluded that the origin of the IPT gene family lies within the ancestor of extant bacteria. Although the longest internal branch leads to AMP-IPTs, this clade is unlikely to represent the ancestral form of IPTs, because the AMP-IPT clade is formed entirely by proteins from either slime moulds or cyanobacterial and proteobacterial plant symbionts or pathogens (red clade in Fig. 2). These bacterial genes, encoded by symbiosis or pathogenicity islands (PAIs) within the main genome or plasmids, being important for host interactions, are known to have been transferred laterally across taxa and consequently have deviating evolutionary rates (*Boyd et al.*, 2009). This holds true also for the putative cyanobacterial adenylate-IPTs as they are found only in species that are also observed as symbionts of specific fungi, liverworts, horwnorts, ferns, gymnosperms, and angiosperms (*Yamada et al.*, 2012, and references within).

Lastly, the *Dictyostelium* adenylate-IPT has been identified as a xenologue which was probably acquired by lateral transfer from bacteria (*Eichinger et al.*, 2005), which is consistent with its position in the tree generated here. The true origin of bacterial AMP-IPT genes is speculative and could either be xenologous or date back to a duplication event in the first bacterial lineage that chose plants as hosts and was subsequently transferred horizontally as part of PAIs.

Consequently, the origin of IPTs lies within bacterial tRNA-IPTs. Based on the tree topology and the taxonomic composition of the two plant tRNA-IPT clades (Fig. 2 class I and II) following the outgroup of AMP-IPTs, the root of eukaryotic IPTs most probably traces back to ancestral α-proteobacteria and to the initial endosymbiotic event leading to extant mitochondria.

Subsequently, this ancestral mitochondrial protein acquired additional domains enabling its function to be extended to
the cytoplasm. While Metazoa and Fungi have kept a single copy, Plantae, namely the lineage formed by the second endosymbiotic event, either transferred the cyanobacterial copy to the nuclear genome or duplicated the mitochondrial tRNA-IPT. The topology of the tree favours the latter scenario. The ancestral eukaryotic IPT was duplicated and resulted in two classes of tRNA-IPT (class I and II). While prasinophytes algae [here: *Micromonas* (Miep) and *Ostreococcus* (Ostlu and Ostta)] as well as seed plants (as exemplified by the well characterized *Arabidopsis*, rice, and maize IPTs) retained both classes of tRNA-IPTs, the distinct lineages leading to the studied Stramenopiles, the red alga, chlorophyte algae, the bryophyte, and the lycophyte, retained only class I IPTs. After an additional duplication event possibly dating back to the LCA of flowering plants, one additional copy of class II tRNA-IPTs subsequently lost the capability to bind tRNAs and diversified into the extant ADP/ATP-IPTs found in flowering plants (green clade in Fig. 2).

As previously reported (Sakakibara, 2006; Frebort et al., 2011), the seven *Physcomitrella* IPTs cluster within the clade of class I tRNA-IPTs. All other species covered in this class code for a single copy tRNA-IPT, which underlines the remarkable evolutionary position of *Physcomitrella* (and probably other mosses) regarding IPTs.

Evidence for a plastidic origin of tRNA-bound Cks

The localization of IPT1:GFP (Fig. 3) corroborated previous sequence-based predictions which also suggested plastidic targeting of IPT1 (Yedvakova and von Schwartzzenberg, 2007). Nevertheless, the uneven distribution of the plastidic fluorescence signal could also point to a specific suborganellar localization of the IPT1 protein in the stroma, for example in close proximity to the plastid ribosomes (Newell et al., 2012).

Although the obtained merged pictures and 3D reconstructions of CLSM stacks favour the assumption that IPT1 is present in the stroma of chloroplasts (Supplementary Fig. S2 at *JXB* online), it is not possible at present state to resolve its exact subplastidic localization. The observed aggregation of IPT1::GFP protein could also be interpreted as an overexpression artefact of the maize ubiquitin promoter resulting in an excess of GFP fusion protein which might have precipitated unevenly in the chloroplast.

IPT1 localized to the chloroplast taken together with IPT1 as the exclusive mechanism of tRNA prenylation does not explain how the tRNA pools in the mitochondria and cytosol are prenylated. The present data do not rule out a function for IPT1 in the cytosol and mitochondria, although no GFP signal was detectable in these compartments. An inaccurate folding of the fusion protein, for example, could have masked targeting sites. For the tRNA-IPT MOD5 from *Saccharomyces cerevisiae*, it has been shown that according to different translational starts, the protein is either transported into mitochondria or stays in the cytosol (Gillman et al., 1991). Further, it has been shown for AtIPT3 in *Arabidopsis* that farnesylation can also determine targeting to different subcellular compartments (Galichet et al., 2008). Due to the scarce knowledge of tRNA trafficking in plant cells, it can only be speculated on as to whether A37 modified tRNAs can be translocated from the chloroplast to other cellular compartments.

Under the assumption that the targeting of IPT1:GFP was not biased by artefacts, the present results point to the chloroplast providing a large amount of *Physcomitrella* prenylated tRNA. This is a contrast to AtIPT2, a tRNA-IPT from *Arabidopsis*, which has been shown to be localized in the cytosol where it is considered to be a significant source of cZ-type Cks using DMAPP from the mevalonate pathway as a side chain donor (Kasahara et al., 2004). For the other tRNA-IPT of *Arabidopsis*, AtIPT9, to which PpIPT1 is an orthologue, no information on localization is available.

**IPT1 is the main enzyme catalysing tRNA prenylation**

The deletion of ipt1 as one of seven ipt genes had a severe effect on the prenylation of A37 in tRNA. The amount of tRNA-bound cZr, iP, tZr, and DHZR in the dipt1 mutants was strongly reduced (Fig. 5) and thus the contribution of the remaining six IPTs to tRNA prenylation can be considered as insignificant in moss protonema. It is therefore concluded that plastidic IPT1 is almost solely responsible for this A37 modification in *Physcomitrella* tRNA. The obtained results are comparable with the report of Dihanich et al. (1987) who found in yeast that the deletion of the single copy tRNA-IPT gene Mod5 led to residual levels of <1.5% tRNA-bound iP compared with the wild type.

If the tRNA pathway was the only pathway for Ck biosynthesis, as suggested by the apparent absence of adenylate-IPTs, a Ck-deficient tRNA as obtained after IPT1 knockout would have led to an overall Ck-deficient plant. These plants should express strongly impaired bud formation as shown for Ck deficiency after overexpression of cytokinin oxidase (von Schwartzzenberg et al., 2007).

Indeed the IPT1 knockout plants showed a strong phenotype which was mainly characterized by altered colony growth, cell shape, and branching of protonema (Fig. 4a, b). This phenotype could not be reversed by addition of exogenous Cks (Fig. 4; Supplementary Fig. S4 at *JXB* online). However, the bud formation as a sensitive marker of Ck response in *Physcomitrella* was neither delayed nor reduced in comparison with the wild type. The reduced growth in colony area of the mutants with its concomitantly enhanced differentiation (Fig. 4e, f) can be mimicked in wild-type agar cultures by exogenous application of iP (Fig. 4g, h) (Thelander et al., 2005; Richter et al., 2012). This morphological feature of dipt1 mutants indicates an overproduction rather than the expected limitation in biosynthesis of bud-inducing Cks.

**Profiling of free Cks indicates a tRNA-independent Ck biosynthesis**

A fully satisfactory explanation of the dipt1 phenotype is only possible after a detailed analysis of the profile of free Cks in the mutant and wild type, which uncovered a differential situation with respect to individual Ck types. While
the levels of tRNA-bound Cks (cZR, iP, tZR, and DHZJR) were strongly reduced, the level of the free Cks revealed a significant reduction only for cZ- and DHZ-type Cks to about a quarter of the respective amount in the wild type. Since IPT1 is localized in chloroplasts, it can be deduced that the majority of free cZ- and DHZ-type Cks is of plastidic tRNA origin.

In contrast to the reduction of free cZ- and DHZ-type Cks, surprisingly a 4-fold increase for the iP- and tZ-type Cks was found. The fact that there is no overall loss of free Cks in the mutants but an increase of tZ and iP conjugates coupled to an almost total loss of tRNA-bound Cks, is strong evidence for an unexpected TRNA-independent pathway for Ck biosynthesis in Physcomitrella.

This finding is in obvious disagreement with the predicted ipt gene complement, which seems to be comprised exclusively of homologues of tRNA-IPTs. Thus, the biosynthetic origin of tZ- and iP-type Cks needs to be re-assessed as tRNA is very unlikely to be their only source.

In Arabidopsis it was described that different pathways are responsible for the formation of distinct types of Cks. The tRNA-IPT-deficient double mutants of Atipt2,9 were shown, like the d|ipt1 mutant, to contain no Cks in their tRNA fraction (Miyawaki et al., 2006). However, in contrast to the Physcomitrella mutants, with 25% residual free cZ-type Cks, the AtIPT2,9-deficient plants did not produce any cZ-type Cks.

In contrast to Arabidopsis where tZ- and iP-type Cks are dominant, the cZ type are the most abundant cytokinins in Physcomitrella. Despite this difference in the Ck profiles, in both plants iP and tZ represent the physiologically most active forms (Sakakibara, 2006; von Schwartzenberg et al., 2007).

Physcomitrella and Arabidopsis differ greatly in the levels of free iP- and tZ-type Cks in the absence of one (PpIPT1) or two tRNA-IPTs (AtIPT2 and -9). While in Arabidopsis the level of free iP- and tZ-type Cks is not greatly affected in the tRNA-IPT double mutants, the d|ipt1 mutants show a vastly increased level of those Cks, thus hinting at differences in Ck biosynthesis, and its regulation, between the bryophyte Physcomitrella and flowering plants. In Physcomitrella, the deficiency of IPT1 apparently causes an up-regulation of the tRNA-independent formation of iP and tZ.

The knockout of ipt1 seems to interfere mainly with the biosynthesis but not with the interconversion between different conjugates of free Cks, thus leading to either generally increased or reduced overall concentrations of the different Ck types (Fig. 5b; Supplementary Table S2 at JXB online). The relative distribution of the different Ck forms within one type (e.g. the ratio of bases to ribosides or nucleotides) is only slightly affected (Supplementary Fig. S5, Table S2 available at JXB online).

Concerning the profile of free Cks, the deletion of IPT1 leads to a shift from cZ-type dominance in the wild type to an iP-type dominance of free cytokinins in d|ipt1 tissue (Fig. 5b). These strong changes in the internal Ck profile also affect the concentration of Cks in the culture medium, with the concentration of iP-type Cks being greatly enhanced in the mutants.

The increase of iP-type Cks in the culture medium is likely to explain certain aspects of the phenotype of d|ipt1 plants. Both a reduction in colony growth area and an increased differentiation might be a direct consequence of the increased external levels of iP and iP, mirrored by the growth habitus of wild-type plants grown on iP-containing media. The fact that the frequency of bud induction is comparable in the mutant and wild type (Fig. 2; Supplementary Fig. S4b at JXB online) shows that the d|ipt1 plants are probably not affected in Ck signalling.

The results of the analyses of d|ipt1 mutants suggest that in Physcomitrella tRNA-bound cZR is most probably the main source for free cZ-type Cks. Considering the loss of tRNA-bound Cks and the concomitant existence of substantial amounts of free iP- and tZ-type Cks, the IPT1-deficient plants strongly indicate the existence of a second, tRNA-independent pathway in the moss. This pathway seems to be important in the production of the active Cks in Physcomitrella.

Given the Ck profile of the d|ipt1 mutant and the hence deduced existence of a tRNA-independent Ck biosynthetic pathway, the clustering of moss IPTs separate from adenylate-IPTs leads to three possible hypotheses. (i) There is an alternative pathway mediating the tRNA-independent production of Cks by an as yet unknown enzyme. (ii) One or several of the other six moss IPTs belong to an as yet uncovered family of (maybe bryophyte-specific) IPTs which have convergently lost the capacity to bind tRNA, evolving adenylate-IPT functionality in parallel to flowering plant ADP/ATP-IPTs. (iii) One or several of the moss IPTs indeed belongs to the same ADP/ATP-IPT family as those of flowering plants, but the phylogenetic signal connecting them has been obscured by an as yet undetermined mechanism. In the last case, the origin of plant adenylate-IPTs would trace back to the ancestor of land plants. Based on the available data, especially the topology and branch lengths of the PpIPT subtree, alternative (ii) seems to be the most probable evolutionary scenario.

The functionally confirmed tRNA-IPT PpIPT1 clusters basal to all other moss IPTs. Some of these six additional IPTs deviate substantially in their branch lengths, indicating changes in the rate of evolution. In order to see whether the increased amounts of iP- and tZ-type Cks are due to an increased expression of one of the remaining IPTs in the d|ipt1 mutant, real-time analyses have been performed. ipt2.1 and ipt2.2 did not show expression either in the wild type or in the mutant. The remaining ipt genes (ipt3–ipt6) showed no strong or consistent changes (Supplementary Fig. S6 at JXB online). Which of the proposed evolutionary hypotheses ultimately can be verified clearly requires additional experimental work. Therefore, studies have been initiated to characterize functionally the in planta function of the remaining members of the Physcomitrella IPT family in order to clarify the identity of the unexpected tRNA-independent Ck biosynthetic pathway observed in Physcomitrella.

**Supplementary data**

Supplementary data are available at JXB online.

**Figure S1.** Vector card for pLNU-GFP_Ppipt1.
Figure S2. Localization of IPT1 within Physcomitrella chloroplasts.
Figure S3. Generation and characterization of dipt1 mutants.
Figure S4. Ck response of dipt1 mutants and the wild type.
Figure S5. Relative level of cytokinins in 22-day-old liquid cultures.
Figure S6. Relative expression of the ipt gene family in dipt1 mutants.
Table S1. Gene identifiers for the phylogenetic analyses performed.
Table S2. Average levels of intracellular free isoprene-type Cks in tissue.
Table S3. Average levels of extracellular free isoprene-type Cks in media.
Table S4. Average levels of tRNA-bound isoprene-Ck ribosides in tissue.

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References

Akiyoshi DE, Regier DA, Jen G, Gordon MP. 1985. Cloning and nucleotide sequence of the tzs gene from Agrobacterium tumefaciens strain T37. Nucleic Acids Research 13, 2773–2788.

Barry GF, Rogers SG, Fraley RT, Brand L. 1984. Identification of a cloned cytokinin biosynthetic gene. Proceedings of the National Academy of Sciences, USA 81, 4776–4780.

Bartels S, Gonzalez Besteiro MA, Lang D, Ulm R. 2010. Emerging functions for plant MAP kinase phosphatases. Trends in Plant Science 15, 322–329.

Beaty JS, Powell GK, Lica L, Regier DA, Macdonald EMS, Hommes NG, Morris RO. 1986. Tzs, a nopaline T plasmid gene from Agrobacterium tumefaciens associated with trans-zeatin biosynthesis. Molecular and General Genetics 203, 274–280.

Boyd EF, Almagro-Moreno S, Parent MA. 2009. Genomic islands are dynamic, ancient integrative elements in bacterial evolution. Trends in Microbiology 17, 47–53.

Brugiere N, Humbert S, Rizzo N, Bohn J, Habben JE. 2009. A member of the maize isopentenyl transferase gene family, Zea mays isopentenyl transferase 2 (ZmIPT2), encodes a cytokinin biosynthetic enzyme expressed during kernel development. Plant Molecular Biology 67, 215–229.

Caillet J, Droogmans L. 1988. Molecular cloning of the Escherichia coli miaA gene involved in the formation of δ2-isopentenyl adenosine in tRNA. Journal of Bacteriology 170, 4147–4152.

Cho SH, von Schwartzzenberg K, Quattraro R. 2009. The role of abscisic acid in stress tolerance. Annual Plant Reviews Volume 36: the moss Physcomitrella patens. Oxford: Wiley-Blackwell, 282–297.

Clamp M, Cuff J, Searle SM, Barton GJ. 2004. The Jalview Java alignment editor. Bioinformatics 20, 426–427.

Cove D, Ashton N. 1984. The hormonal regulation of gametophytic development in bryophytes. In: Dyer AF, Duckett JG, eds. The experimental biology of bryophytes. London: Academic Press, 177–201.

Decker EL, Frank W, Sarnighaen E, Reski R. 2006. Moss systems biology en route: phytohormones in Physcomitrella development. Plant Biology 8, 397–405.

Dihanich ME, Najarian D, Clark R, Gillman EC, Martin NC, Hopper AK. 1987. Isolation and characterization of MOD5, a gene required for isopenstenylation of cytoplasmic and mitochondrial tRNAs of Saccharomyces cerevisiae. Molecular and Cellular Biology 7, 177–184.

Eichinger L, Pachetat JA, Gluckner G, et al. 2005. The genome of the social amoeba Dictostelium discoideum. Nature 435, 43–57.

Fais M, Zalubilova J, Strnad M, Schmulling T. 1997. Conditional transgenic expression of the ipt gene indicates a function for cytokinins in paracrine signaling in whole tobacco plants. The Plant Journal 12, 401–415.

Freibrot I, Kowalska M, Hlusa T, Frebortova J, Galuszka P. 2011. Evolution of cytokinin biosynthesis and degradation. Journal of Experimental Botany 62, 2451–2452.

Galichet A, Hoyerova K, Kaminek M, Gruissem W. 2008. Farnesyltransferase directs AITP3 subcellular localization and modulates cytokinin biosynthesis in Arabidopsis. Plant Physiology 146, 1155–1164.

Gillman EC, Slusher LB, Martin NC, Hopper AK. 1991. MOD5 translation initiation sites determine N6-isopenstenyladenosine modification of mitochondrial and cytoplasmic tRNA. Molecular and Cellular Biology 11, 2382–2390.

Golovko A, Hjalm G, Sitton F, Nicander B. 2000. Cloning of a human tRNA isopentenyl transferase. Gene 255, 55–93.

Golovko A, Sitton F, Tillberg E, Nicander B. 2002. Identification of a tRNA isopenstenytransferase gene from Arabidopsis thaliana. Plant Molecular Biology 49, 161–169.

Goodstein DM, Shu S, Howson R, et al. 2012. Phytozome: a comparative platform for green plant genomics. Nucleic Acids Research 40, D1178–D1186.

Howe K, Bateman A, Durbin R. 2002. QuickTree: building huge Neighbor-Joining trees of protein sequences. Bioinformatics 18, 1546–1547.

Hunter S, Jones P, Mitchell A, et al. 2012. InterPro in 2011: new developments in the family and domain prediction database. Nucleic Acids Research 40, D306–D312.

Kakimoto T. 2001. Identification of plant cytokinin biosynthetic enzymes as dimethylallyl diphosphate:ATP/ADP isopenstenytransferases. Plant and Cell Physiology 42, 677–685.

Kakimoto T. 2003. Biosynthesis of cytokinins. Journal of Plant Research 116, 233–239.

Kamada-Nobusada T, Sakakibara H. 2009. Molecular basis for cytokinins biosynthesis. Phytochemistry 70, 444–449.

Kamisugi Y, Schlink K, Rensing SA, Schwechke D, von Stackelberg M, Cuming AC, Reski R, Coke DJ. 2006. The mechanism of gene targeting in Physcomitrella patens: homologous recombination, concatenation and multiple integration. Nucleic Acids Research 34, 6205–6214.

Kasahara H, Takei K, Ueda N, Hishiyama S, Yamaya T, Kamei Y, Yamaguchi S, Sakakibara H. 2004. Distinct isopensterynol origins of cis and trans-zeatin biosynthesis in Arabidopsis. Journal of Biological Chemistry 279, 14049–14054.

Knight CD, Coke D, Boyd PJ, Ashton N. 1988. The isolation of biochemical and developmental mutants in Physcomitrella patens. In: Gilme JM, ed. Methods in Bryology: Proceedings of the Bryology Methods Workshop, Mainz. Harfot Botanical Laboratory, Nichinan, Japan, 47–58.

Konevga AL, Soboleva NG, Makhno VI, Peshekhonov AV, Katunin VI. 2006. The effect of modification of tRNA nucleotide-37 on the tRNA interaction with the P- and A-site of the 70S ribosome Escherichia coli. Molecular Biology 40, 669–683.

Kopcma N, Blaschke H, Kopecny D, Vigouroux A, Koncikova R, Novak O, Kotland O, Strnad M, Morera S, von Schwartzzenberg K. 2013. Structure and function of nucleoside hydrolases from Physcomitrella and maize catalyzing hydrolysis of purine, pyrimidine and cytokinin ribosides. Plant Physiology 163, 1568–1583.

Lang D, Eisinger J, Reski R, Rensing SA. 2005. Representation and high-quality annotation of the Physcomitrella patens transcriptome.
demonstrates a high proportion of proteins involved in metabolism in mosses. Plant Biology 7, 238–250.

Lang D, Weibe B, Timmerhaus G, Richardt S, Riano-Pachon DM, Correa LG, Reski R, Mueller-Roeder B, Rensing SA. 2010. Genome-wide phylogenetic comparative analysis of plant transcriptional regulation: a timeline of loss, gain, expansion, and correlation with complexity. Genome Biology and Evolution 2, 488–503.

Lang D, Zimmer AD, Rensing SA, Reski R. 2008. Exploring plant biodiversity: the Physcomitrella genome and beyond. Trends in Plant Science 13, 542–549.

Lichtenstein C, Klee H, Montoya A, Garfinkel D, Fuller S, Flores C, Nester E, Gordon M. 1984. Nucleotide sequence and transcript mapping of the tnr gene of the PiTAgNC octopine Ti-plasmid: a bacterial gene involved in plant tumorigenesis. Journal of Molecular and Applied Genetics 2, 354–362.

Maass H, Klämbt D. 1981. On the biogenesis of cytokinins in roots of Phaseolus vulgaris. Planta 151, 353–358.

Miyawaki K, Tarkowski P, Matsumoto-Kitano M, Kato T, Sato S, Tarkowska D, Tabata S, Sandberg G, Kakimoto T. 2006. Roles of Arabidopsis ATP/ADP isopentenyltransferases and tRNA isopentenyltransferases in cytokinin biosynthesis. Proceedings of the National Academy of Sciences, USA 103, 16058–16063.

Mok DWS, Mok MC. 2001. Cytokinin metabolism and action. Annual Review of Plant Physiology and Plant Molecular Biology 52, 89–118.

Murai N. 1994. Cytokinin biosynthesis in tRNA and cytokinin incorporation into plant RNA. In: Mok DWS, Mok MC, eds Cytokinins: chemistry, activity, and function. Boca Raton, FL: CRC Press, 87–99.

Newell CA, Natesan SK, Sullivan JA, Jouhet J, Kavanagh TA, Gray JC. 2012. Exclusion of plastid nucleoids and ribosomes from stromules in tobacco and Arabidopsis. The Plant Journal 69, 399–410.

Notroff C, Higgins DG, Heringa J. 2000. T-Coffee: a novel method for fast and accurate multiple sequence alignment. Journal of Molecular Biology 302, 205–217.

Novák O, Hauserova E, Amakorova P, Doležal K, Strnad M, 2008. Cytokinin profiling in plant tissues using ultra-performance liquid chromatography-electrospray tandem mass spectrometry. Phytochemistry 69, 2214–2224.

Novák O, Tarkowski P, Tarkowská D, Doležal K, Lenobel R, Strnad M. 2003. Quantitative analysis of cytokinins in plants by liquid chromatography–single-quadrupole mass spectrometry. Analytica Chimica Acta 490, 207–218.

Papounov P, Teale W, Lang D, Paponov M, Reski R, Rensing SA, Palme K. 2009. The evolution of nuclear auxin signalling. BMC Evolutionary Biology 9, 126.

Patil G, Nicander B. 2013. Identification of two additional members of the tRNA isopentenyltransferase family in Physcomitrella patens. Plant Molecular Biology 82, 417–426.

Pils B, Heyl A. 2009. Unraveling the evolution of cytokinin signaling. Plant Physiolology 151, 782–791.

Prigge MJ, Bezanilla M. 2010. Evolutionary crossroads in developmental biology. Physcomitrella patens. Development 137, 3535–3543.

Rensing SA, Lang D, Zimmer AD, et al. 2008. The Physcomitrella genome reveals evolutionary insights into the conquest of land by plants. Science 319, 64–69.

Reski R, Abel WO. 1985. Induction of budding on chloronemata and caulonemata of the moss, Physcomitrella patens, using isopentenyladenine. Planta 165, 354–358.

Reutter K, Atzorn R, Hadeler B, Schmulling T, Reski R. 1998. Expression of the bacterial ipt gene in Physcomitrella patens rescues mutations in budding and in plastid division. Planta 206, 196–203.

Richter H, Lieberei R, Strnad M, Novak O, Gruz J, Rensing SA, von Schwartzzenberg K. 2012. Polyphenol oxidases in Physcomitrella: functional rpo1 knockout modulates cytokinin-dependent development in the moss Physcomitrella patens. Journal of Experimental Botany 63, 5121–5135.

Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19, 1572–1574.

Sakakibara H. 2006. Cytokinins: activity, biosynthesis, and translocation. Annual Review of Plant Biology 57, 431–449.

Sacramento H, Sakakibara H, Kojima M, Yamamoto Y, Nagasaki H, Inukai Y, Sato Y, Matsuoka M. 2006. Ectopic expression of KNOTTED1-like homeobox protein induces expression of cytokinin biosynthesis genes in rice. Plant Physiology 142, 54–62.

Schafer D, Zryd JP, Knight CD, Cove DJ. 1991. Stable transformation of the moss Physcomitrella patens. Molecular and General Genetics 226, 419–424.

Schafer DG, Delacote F, Charlot F, Vrielynck N, Guyon-Debast A, Le Guin S, Neuhaus JM, Doutriaux MP, Nogue F. 2010. RAD51 loss of function abolishes gene targeting and de-represses illegitimate integration in the moss Physcomitrella patens. DNA Repair 9, 526–533.

Schafer DG, Zryd JP. 1997. Efficient gene targeting in the moss Physcomitrella patens. The Plant Journal 11, 1195–1206.

Schulz P, Reski R, Maldiney R, Laloue M, von Schwartzzenberg K. 2000. Kinetics of cytokinin production and bud formation in Physcomitrella: analysis of wild type, a developmental mutant and two of its ipt transgensics. Journal of Plant Physiology 156, 768–774.

Schulz PA, Hofmann AH, Russo VEA, Hartmann E, Laloue M, von Schwartzzenberg K. 2001. Cytokinin overproducing ove mutants of Physcomitrella patens show increased riboside to base conversion. Plant Physiology 126, 1224–1231.

Schween G, Gorr G, Hohe A, Reski R. 2003. Unique tissue-specific cell cycle in Physcomitrella. Plant Biology 5, 50–58.

Schween G, Schulte J, Reski R, Hohe A. 2005. Effect of ploidy level on growth, differentiation, and morphology in Physcomitrella patens. Bryology 108, 27–35.

Sonnhhammer ELL, Hollich V. 2005. Scoredist: a simple and robust protein sequence distance estimator. BMC Bioinformatics 6, 108.

Stirk WA, Balint P, Sipoo G, Dobrev PI, Novak O, Dolezal K, Strnad M, van Staden J, Ordog V. 2011. Endogenous cytokinins in synchronized Chlorella cultures in relation to light and the cell cycle. European Journal of Physiology 46, 73–73.

Strnad M. 1997. The aromatic cytokinins. Physiologia Plantarum 101, 674–688.

Takel K, Sakakibara H, Sugiyama T. 2001. Identification of genes encoding adenylyl isopentenyltransferase, a cytokinin biosynthesis enzyme, in Arabidopsis thaliana. Journal of Biological Chemistry 276, 26405–26410.

Taya Y, Tanaka Y, Nishimura S. 1978. 5′-Amp is a direct precursor of cytokinin in Dicotyolum discoidum. Nature 271, 545–547.

Thelander M, Olsson T, Ronne H. 2005. Effect of the energy supply on filamentous growth and development in Physcomitrella patens. Journal of Experimental Botany 56, 653–662.

Vandenbussche F, Fierro AC, Wiedemann G, Reski R, Van Der Straeten D. 2007. Evolutionary conservation of plant gibberellin signalling pathway components. BMC Plant Biology 7, 65.

von Schwartzzenberg K. 2009. Hormonal regulation of development by auxin and cytokinin in moss. Annual Plant Reviews Volume 36: the moss Physcomitrella patens. Oxford: Wiley-Blackwell, 246–281.

von Schwartzzenberg K, Nunez MF, Blaschke H, Dobrev PI, Novak O, Motyka V, Strnad M. 2007. Cytokinins in the bryophyte Physcomitrella patens: analyses of activity, distribution, and cytokinin oxidase/dehydrogenase overexpression reveal the role of extracellular cytokinins. Plant Physiology 145, 786–800.

Wang TL, Beutelmann P, Cove DJ. 1981. Cytokinin biosynthesis in mutants of the moss Physcomitrella patens. Plant Physiology 68, 739–744.

Yamada S, Ohkubo S, Miyashita H, Setoguchi H. 2012. Genetic diversity of symbiotic cyanobacteria in Cymas revoluta (Cyanodaceae). FEMS Microbiology Ecology 81, 696–706.

Yevdakova NA, Motyka V, Malbeck J, Travnickova A, Novak O, Strnad M, von Schwartzzenberg K. 2008. Evidence for importance of IRN-dependent cytokinin biosynthetic pathway in the moss Physcomitrella patens. Journal of Plant Growth Regulation 27, 271–281.

Yevdakova NA, von Schwartzzenberg K. 2007. Characterisation of a prokaryote-type tRNA-isopentenyltransferase gene from the moss Physcomitrella patens. Planta 226, 683–695.

Zimmer AD, Lang D, Buchta K, Rombauts S, Nishiyama T, Hasebe M, Van de Peer Y, Rensing SA, Reski R. 2013. Reannotation and extended community resources for the genome of the non-seed plant Physcomitrella patens provide insights into the evolution of plant gene structures and functions. BMC Genomics 14, 498.