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Phytosulfokine peptide library: chemical synthesis and biological evaluation on protoplast regeneration†

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Phytosulfokine (PSK) is a highly conserved plant peptide hormone (PPH) containing two sulfated tyrosine residues. PSK acts as a growth factor in *Arabidopsis thaliana* plant species and significantly improves the efficiency of cellular proliferation in plant cell culture, a process known as regeneration. However, many important plant species are recalcitrant under exposure to PSK and require e.g., artificially generated PPHs to boost regeneration. Here, we report the design, chemical synthesis, and biological evaluation of the regeneration capacity of a PSK-like peptide library. Our set of thirteen PSK analogues included N-terminal modifications, amino acid substitutions, incorporation of diastereomers, and backbone N-methylation. Insertion of the latter required solution-phase synthesis of the proper building block prior to solid-phase peptide synthesis. The biological activity of the PSK analogues was assessed in the regeneration capacity of *Brassica oleracea* cells (protoplasts). Most PSK analogues did not affect the regeneration capacity as compared to PSK. For example, when an alanine substitution was introduced and replaced either 2-Ile or 4-Thr, the bio-activity was fully abolished. However, 2-*allo*-Ile PSK and 4-*Nme*-Thr PSK showed increased activity in protoplast regeneration when compared with that of the native peptide. These findings indicated that introducing small chemical modifications in the PPH PSK provides opportunities for inducing protoplast regeneration in recalcitrant species.

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Introduction

Environmental influences significantly impact plant development and growth. Stress conditions can change plant growth, reduce crop productivity, alter gene expression, or affect cellular metabolism.^{1,2} In an adaptive response, plant tissue can withstand stress by re-initiation of cell division, or preservation of totipotent cells.³ Analogous to this process, responsive plant tissues or cells can resume cell division when exposed to media enriched with the phytohormone auxin and cytokinin, giving rise to new organs and the potential to grow a full plant that can complete its lifecycle.⁴ In the field of plant research, farming practices and plant breeding are investigated as methods to reduce plant stress.^{5–7} Unfortunately, many plant species that are important from an agricultural point of view display poor

tolerance to stress conditions, and furthermore, are challenging to improve by efficient breeding and genetic engineering methods due to their recalcitrance in regeneration. Plant hormones are key in the control of totipotency and regeneration capacity and represent signaling molecules that are involved in signal reception, signal transduction, and cellular response. To transmit information intra- and intercellularly between plant cells and their external environment, chemical signaling is used for down- and upregulation of gene expression.⁸ Various plant hormones have been identified as important regulatory molecules in plant cells and their involvement with plant signaling and molecular trafficking has been established. For example, abscisic acid, auxin, cytokinin, ethylene, and the family of gibberellins have been identified as major plant hormones.⁹ These hormones play an important role in initiating plant dormancy, cell differentiation, cell division, fruit ripening, and stem elongation, respectively. In addition to these phytohormones, the family of plant peptide hormones (PPHs) was also identified as important in plant signaling.^{10–13}

Phytosulfokine (PSK) is a secreted PPH that is bioactive even at nanomolar concentrations.¹⁴ It was first isolated from an *Asparagus officinalis* L. mesophyll cell culture medium and

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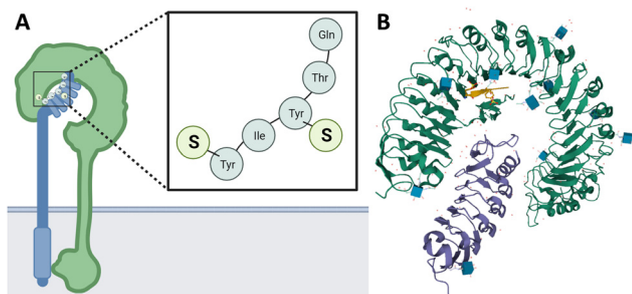


Fig. 1 (A) Illustration of phytosulfokine (PSK) being perceived at the plasma membrane by the PSK receptor (PSKR1) and co-receptor SERK1. S represents the sulfate modification on the tyrosine residue. (B) Crystal structure of the plant peptide hormone receptor-coreceptor complex including ligand (PSK-PSKR1-SERK1) in *Arabidopsis thaliana* (PDB-code: 4z64; receptor in green, co-receptor in purple, PSK in orange).

found to promote cell division of asparagus mesophyll cells cultured at low cell density.¹⁵ The PPH is a disulfated pentapeptide H-Tyr(SO₃H)-Ile-Tyr(SO₃H)-Thr-Gln-OH, which is known as PSK- α . In this peptide, both tyrosine residues contain a post-translationally installed sulfate group that is introduced on a PSK precursor by tyrosylprotein sulfotransferase (TPST) using 3'-phosphoadenosine-5'-phosphosulfate (PAPS) as a substrate.^{16,17} Analysis of PSK analogues revealed the importance of the sulfated residues for its biological activity (*i.e.*, nonsulfated PSK is inactive below 100 nM in stimulating cell proliferation at low density),¹⁸ and identified Tyr(SO₃H)-Ile-Tyr(SO₃H) as the core requirement of PSK to display activity (*i.e.*, the pharmacophore). In the cell, PSK is synthesized from 80–120 amino acid long prepropeptides that are encoded by small gene families and unlike these precursors, the mature PSK peptide is strongly conserved.^{19,20}

The formed mature peptide is recognized by the PSK receptor (PSKR1, Fig. 1A), a transmembrane receptor belonging to the leucine-rich repeat receptor-like kinase (LRR-RLK) family with more than 600 members in *Arabidopsis* and over 1100 members in rice.^{21–23} The PSKR1 contains an extracellular domain, located in the cell membrane by a single transmembrane helix, and a C-terminal intracellular kinase domain.²⁴ Binding sites of PSK to the PSKR1 have been found in the plasma membrane of rice suspension cells, asparagus, carrot, maize, tobacco, and tomato plants with photoaffinity labeling and radiolabeling experiments.^{25–27} Moreover, the crystal structure of the PSKR1 (Fig. 1B) has been resolved. In addition, the formation of the PSK-receptor complex, and allosteric activation by PSK has been discovered, leading to detailed insights in peptide-receptor interaction and receptor conformation in both bound and unbound state.^{28,29}

PSK is universally found in both monocotyledon and dicotyledon plant species and strongly promotes proliferation of plant cells in culture.¹⁹ In addition to cell proliferation, PSK also promotes plant cell differentiation, organogenesis, and somatic embryogenesis.^{30,31} Previously, PSK was tested for its ability to enhance regeneration in cauliflower varieties, showing its variety-dependent potential in shoot regeneration.^{32,33} In addition to cauliflower, PSK was also tested in sugar beet

protoplast regeneration, in which some recalcitrant cultures could be induced to cell division upon application of exogenous PSK peptide.³⁴ So far, characteristics and functions of PSK have been reviewed quite extensively.^{35–38} PSK have been categorized as a growth factor in *Arabidopsis thaliana*.^{39,40} As we now understand, it has many more activities besides growth regulation, such as responses to biotic and abiotic stress and reproductive processes.^{41,42} In recalcitrant plant species there are still several bottlenecks that need to be solved before PSK can be successfully used as a regenerator peptide. Based on literature, the PSK peptide has potential, but needs optimization for efficient plant regeneration in a broad range of plant varieties, for example *Brassica oleracea* (red cabbage). Therefore, we used PSK as starting point in search for modified PSK peptides with improved performance in protoplast regeneration of recalcitrant plant species.

Here, we report the chemical synthesis and analysis of the bio-activity of a set of PSK analogues that are not yet present in literature.^{43–45} To evaluate the PSK-like library of mutated PSK peptides, screening for any induction or enhancement of regeneration as compared to the original PSK peptide was assessed by determining their effect on the regeneration of *Brassica oleracea* protoplasts. Ultimately, this will lead to new insights into structural features of PSK-like peptides, and the effects of these peptides on cell growth, that improve the regeneration capacity and affects the stress levels of the plant.

Results and discussion

PSK was first discovered by phenotypic analysis and biochemical purification in *Asparagus officinalis* L. culture medium by Matsubayashi *et al.*¹⁵ The same group was also the first to report the chemical synthesis of PSK- α in 1996.⁴⁵ In addition to PSK- α , various analogues were reported and structure–activity relationships were established according to mitogenic activities measured by a highly sensitive bioassay method. Results of these studies indicated that the N-terminal tripeptide of PSK acts as the active core.

Chemical synthesis of phytosulfokine (PSK)

PSK (1, Fig. 2) was synthesized following standard Fmoc/*t*Bu-based solid-phase peptide synthesis (SPPS). Starting with a Wang resin preloaded with Fmoc-Gln(Trt) (loading: 0.4–0.9 mmol g⁻¹), the peptide chain was elongated with Fmoc-Thr(*t*Bu)-OH using the coupling reagents 2-(1*H*-benzotriazole-1-yl)-1,1,3,3-tetramethyl-uronium hexafluorophosphate (HBTU) and 1-hydroxy-benzotriazole (HOBT) in the presence of DIPEA. Subsequently, Fmoc-Tyr(SO₂ONp)-OH, Fmoc-Ile-OH, and Fmoc-Tyr(SO₂ONp)-OH were coupled using similar chemistry. We used commercially available Fmoc-Tyr(SO₂ONp)-OH, which contains a neopentyl (Np) protecting group for the sulfated tyrosine residue, as a stable and easy cleavable protecting group to incorporate the sulfated tyrosine residues.⁴⁶ The Np moiety was removed either by conditions involving unhindered strong nucleophiles in polar aprotic solvent with long reaction times and elevated temperatures,⁴⁷ or by treatment with an aqueous



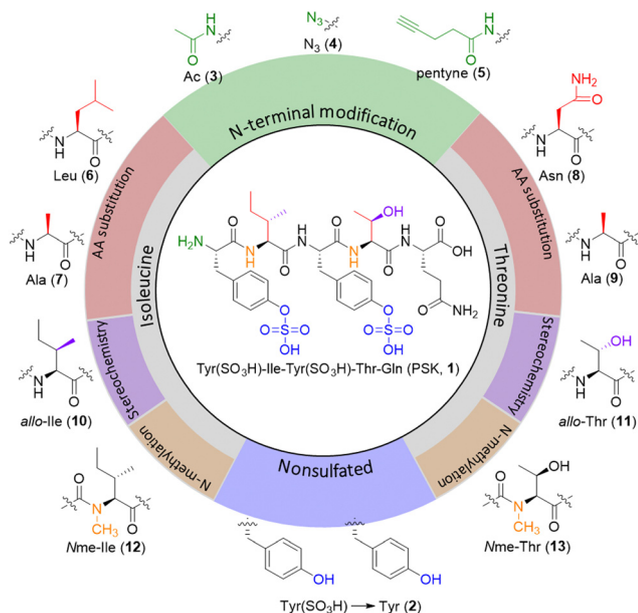


Fig. 2 Chemical structure of PSK (1) and its analogues. Nonsulfated PSK (2); N-terminal modified PSK (3–5); amino acid substituted PSK (6–9); diastereomer PSK (10–11); N-methylated PSK (12–13).

solution of ammonium acetate (NH_4OAc) at 37 °C for at least 12 h,⁴⁸ resulting in the sulfate monoester.

After completion of PSK synthesis on solid support, the resin-bound peptide H-Tyr(SO_2ONp)-Ile-Tyr(SO_2ONp)-Thr(*t*Bu)-Gln(Trt)-OH was cleaved from the solid support and concomitant removal of *tert*-butyl and trityl protecting groups with an acidic TFA cocktail (95% TFA: 2.5% triisopropylsilane (TIS): 2.5% H_2O , reaction time: 1.5–2 h). After acidic cleavage, the crude peptide was precipitated in ice-cold ether and the residue was dissolved in $\text{H}_2\text{O}:\text{MeCN}$ 6:4 from which it was lyophilized. The lyophilized product was first purified by reversed-phase (RP) HPLC using acidic buffers. After purification and lyophilization of the PSK-containing fractions, the obtained powder was dissolved in a 1–2 M NH_4OAc solution and a minimum amount of acetonitrile as co-solvent, and subsequently heated to 40–45 °C for 40 h. RP-HPLC analysis of the mixture at different time-points showed complete Np-removal after 40 h (Fig. S1, ESI[†]). The resulting sulfate monoester was purified by preparative RP-HPLC using 10 mM NH_4OAc buffers. Purified PSK was analyzed with RP-HPLC coupled to a mass spectrometer (MS), and with nuclear magnetic resonance (NMR; peak assignment was done with standard 2D NMR techniques, Fig. S2, ESI[†]). In MS, sulfate fragmentation might occur due to small traces of acid in the system (see ESI[†] for analytical spectra and data). Non-sulfated PSK (PSK-2, Fig. 2) was synthesized analogously, although added steps as result of the sulfated Tyr could be omitted. Compared to the ¹H NMR of PSK-1, non-sulfated PSK-2 only shows notable differences in ppm peaks for the protons of the tyrosine residue (Fig. S1, ESI[†]).

Design of PSK analogues

Over 50 PSK analogues have been reported in different studies.^{43–45} These analogues mainly focused on changes in

the tyrosine residues, for example by incorporation of non-proteinogenic aromatic amino acid residues or halogenated phenol amino acids, and showed that the presence of a sulfate group at Tyr-1 or Tyr-3 of the PSK peptide plays an important role in the bio-activity of the PSK peptide. Here we select analogues with different types of small modifications while maintaining the sulfated tyrosine residues intact (Fig. 2).

We first installed N-terminal modifications such as an acetyl (Ac-PSK 3), or an azide (N_3 -PSK 4) or pentynoic amide (pentynoic-PSK 5) for further bioconjugation strategies.⁴⁹ These modifications can be conveniently introduced as final step in SPPS or, for the azide, after peptide purification using a transfer reagent.⁵⁰ For mutations of the amino acids, Matsubayashi *et al.*⁴⁵ reported >95% reduction in activity of a PSK analogue which lacks the δ - CH_3 -group on position 2 (resulting from the Ile2Val substitution), but also for mutations on position 4 with for example serine. We report four additional mutations at these two sites, namely 2-Leu PSK and 2-Ala PSK (analogues 6 and 7, respectively), and 4-Asn PSK and 4-Ala PSK (analogues 8 and 9, respectively). Next, we selected analogues containing diastereomers of both isoleucine on position 2 and threonine on position 4 by using the *allo* amino acids. We incorporated 2-*allo*-Ile PSK 10 and 4-*allo*-Thr PSK 11, resulting in small changes in spatial orientation of a CH_3 -group or OH-group at the C β atom. These diastereomers can easily be introduced by SPPS using Fmoc-*allo*-Ile-OH (Bachem) or Fmoc-*allo*-Thr(*t*Bu)-OH (Bachem). Finally, we selected N-methylated modifications on positions 2 or 4, resulting in 2-Nme-Ile PSK 12 and 4-Nme-Thr PSK 13 in order to prevent hydrogen-bond formation between PSK and the receptor *via* these peptide bonds. The N-methylated peptides can be synthesized with SPPS, with in solution-phase synthesized building blocks.

Synthesis of mutated PSK peptides

N-terminal modifications. Synthesis of N-terminally modified PSK analogues started with the general procedure of PSK synthesis described above. Acetyl-PSK-3 was obtained after treatment of the H_2N -Tyr(SO_2ONp)-Ile-Tyr(SO_2ONp)-Thr(*t*Bu)-Gln(Trt)-Wang resin with acetyl capping solution. Azide was installed on PSK with imidazole-1-sulfonyl azide hydrochloride (S1, ESI[†]) as azidotransfer reagent.⁵¹ Previous studies showed the use of this reagent for simple and efficient solid-phase preparation of azido-peptides.⁵² However, on-resin the N-terminal amine was not transformed into an azide moiety. Instead, we used a procedure in solution-phase involving high pH and the presence of Cu(II) .⁵⁰ After 40 h of reaction time and lyophilization, N_3 -PSK, PSK-4, was obtained. The final N-terminal modified peptide was prepared similarly to acetyl-PSK. For the alkyne-functionalized PSK H-Tyr(SO_2ONp)-Ile-Tyr(SO_2ONp)-Thr(*t*Bu)-Gln(Trt)-Wang resin was treated with 4-pentynoic acid, HBTU, HOBt, and DIPEA, to yield pure peptide pentynoic-PSK, PSK-5, after preparative-HPLC purification.

Amino acid substitution. PSK analogues containing Leu or Ala instead of 2-Ile (analogues PSK-6 and PSK-7) or analogues that contain Asn or Ala instead of 4-Thr (analogues PSK-8 or PSK-9) were obtained by standard Fmoc/*t*Bu-based SPPS.

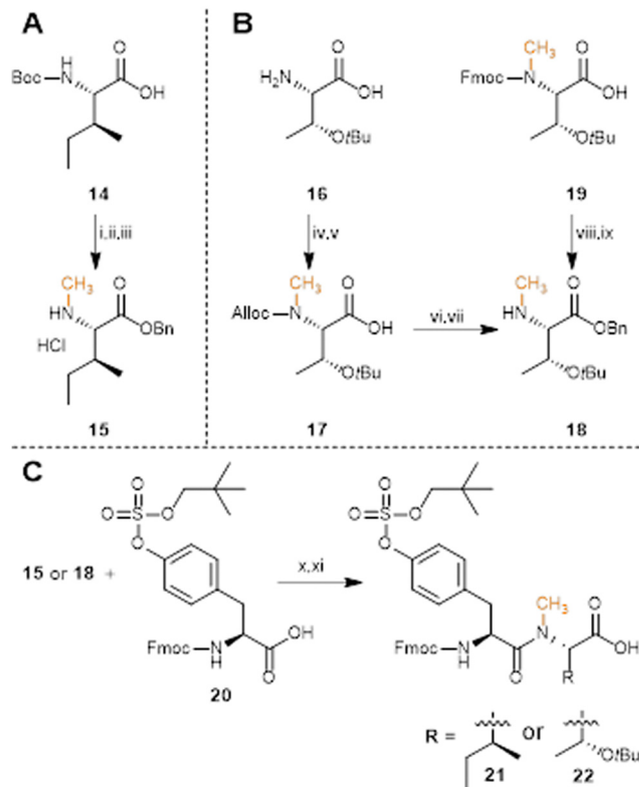


All synthesized peptides were obtained in analytically pure form and their identity was confirmed with LC-MS and NMR comparison (see ESI†, Fig. S4). Remarkably, 2-Ala PSK (PSK-7) displayed a retention time (t_R) of only 5.5 min on LC-MS, while all other single amino acid substituted PSK analogues have a t_R range between 11.2–11.5 min (on a C18 column) (see ESI†).

Diastereomers. 2-*allo*-Ile containing PSK-10 was synthesized using Fmoc-*allo*-Ile-OH, whereas 4-*allo*-Thr PSK, PSK-11, was prepared with Fmoc-*allo*-Thr(*t*Bu)-OH. Both peptides PSK-10 and PSK-11 showed a similar t_R of 11.3 min on LC-MS, which is slightly longer than that of native peptide PSK-1 (t_R = 10.4 min).

N-methylation. Despite the commercial availability of Fmoc-Nme-Ile-OH or Fmoc-Nme-Thr(*t*Bu)-OH, we were not able to couple Fmoc-Tyr(SO₂ONp)-OH in SPPS to the *N*-methylated amino acid. Different carboxyl activators such as HATU, HBTU were applied without success, and even an *in situ* prepared acid chloride⁵³ did not provide the expected coupling in SPPS. We concluded that coupling of Fmoc-Tyr(SO₂ONp)-OH to these two resin-bound *N*-alkylated β -substituted amino acids is severely hampered by steric factors. Therefore, we prepared the respective protected dipeptide building blocks in solution (Scheme 1), and coupled these to the appropriate resin-bound precursor (Scheme 2). To synthesize the isoleucine building block (Scheme 1A), we first executed a *N*-alkylation of Boc-Ile-OH 14 with methyl iodide and sodium hydride.^{54,55} The product was subsequently benzylated at the carboxylic acid using Cs₂CO₃ and benzyl bromide. The Boc protecting group was then removed with 4 M HCl in 1,4-dioxane to obtain Nme-Ile-OBn hydrochloride 15 in 74% yield over 3 steps.

For preparation of the threonine building block, two synthetic routes were explored (Scheme 1B). First, H₂N-Thr(*t*Bu)-OH 16 required an orthogonal protecting group strategy to maintain the threonine side chain protected in the final building block.⁵⁶ Therefore, the *N*-terminus was first protected with an allyloxycarbonyl (Alloc) protecting group by reacting 16 with allyl chloroformate in the presence of sodium bicarbonate. Afterwards, the carbamate amine was methylated using methyl iodide and sodium hydride to afford Alloc-Nme-Thr(*t*Bu)-OH 17, and the C-terminus was protected with a benzyl group. Finally, the Alloc protecting group was removed from the secondary amine using palladium-catalyzed allyl transfer to thiosalicylic acid as the accepting nucleophile,^{57,58} and yielded Nme-Thr(*t*Bu)-OBn 18 in 51% yield over 4 steps. We found that the same threonine building block could also be synthesized starting from commercially available Fmoc-Nme-Thr(*t*Bu)-OH 19. In two steps that involved benzylation of the carboxyl moiety and Fmoc-removal with piperidine, Nme-Thr(*t*Bu)-OBn 18 was obtained in 81% yield over 2 steps. Clearly, this last route is preferred, although the former might be more favorable for large amounts of *N*-methylated amino acid 18. Using building blocks 15 and 18, solution-phase dipeptide synthesis was performed by coupling Fmoc-Tyr(SO₂ONp)-OH using HATU and DIPEA in DMF (Scheme 1C). The formed products were hydrogenated with Pd/C and H₂ to remove the benzyl protecting group and Fmoc-Tyr(SO₂ONp)-Nme-Ile-OH 21 was obtained in



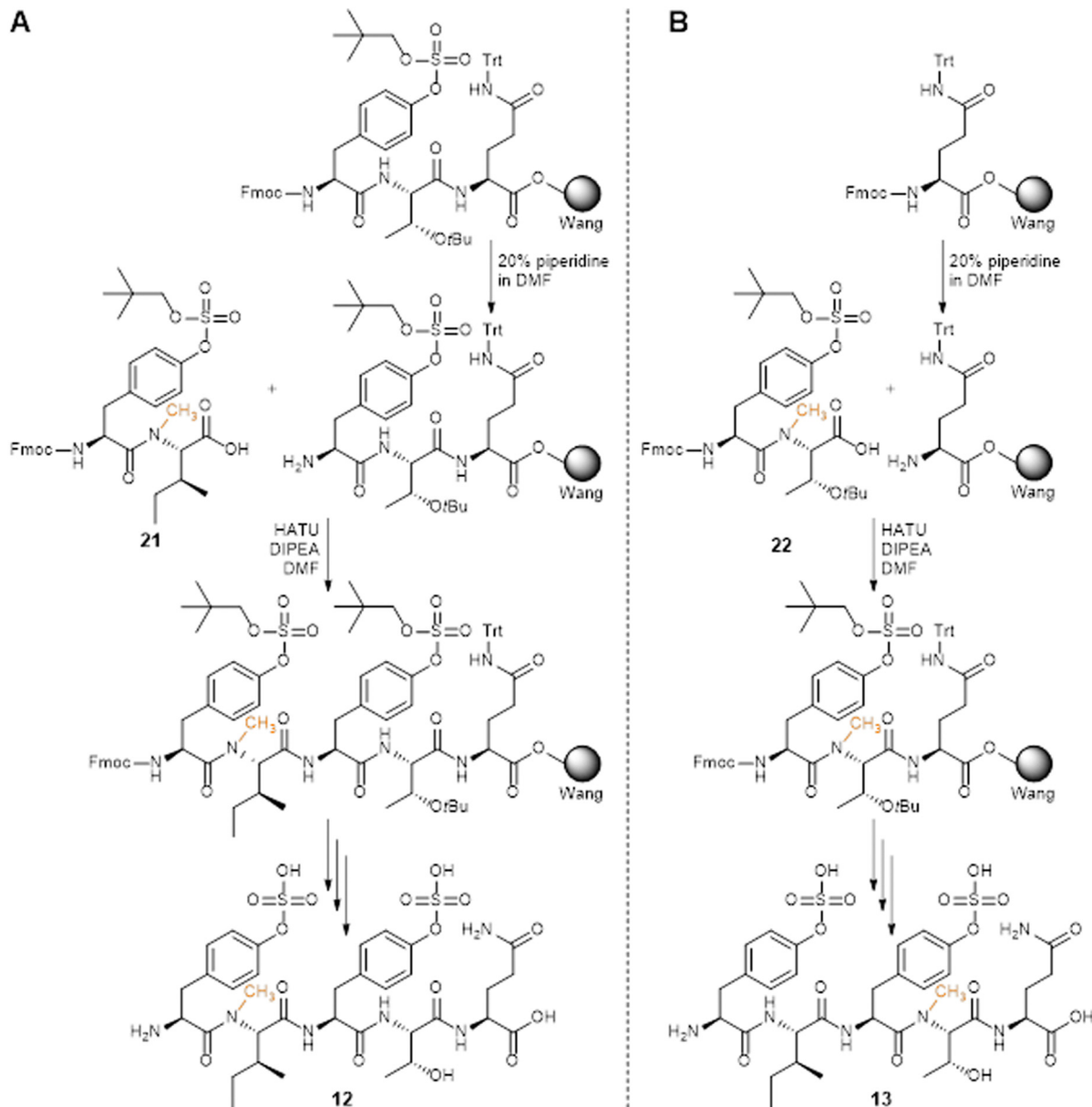
Scheme 1 (A) Synthesis of *N*-methylisoleucine 15, (B) *N*-methylthreonine 18, and (C) dipeptide building blocks containing *N*-methylisoleucine 21 and *N*-methylthreonine 22. Reagents and conditions: (i) MeI, NaH, THF, 89%; (ii) Cs₂CO₃, MeOH, then BnBr, MeCN, 84%; (iii) HCl, 1,4-dioxane, 99%; (iv) Alloc-Cl, sat. NaHCO₃, THF, 93%; (v) MeI, NaH, THF, 83%; (vi) Cs₂CO₃, MeOH, then BnBr, MeCN, 80%; (vii) 10% Pd(PPh₃)₄, thiosalicylic acid, DCM, 82%; (viii) Cs₂CO₃, MeOH, then BnBr, MeCN, 98%; (ix) piperidine, DCM, 83%; (x) HATU, DIPEA, THF, quantitative; (xi) Pd/C, H₂, EtOH, CHCl₃, 94% (21), 62% (22).

94% yield over 2 steps, whereas Fmoc-Tyr(SO₂ONp)-Nme-Thr(*t*Bu)-OH 22 was obtained in 62% yield over 2 steps.

The obtained dipeptide building blocks 21 and 22 were used in SPPS to synthesize 2-*N*-me-Ile PSK-12 and 4-*N*-me-Thr PSK-13, respectively (Scheme 2), with HATU as coupling reagent. For 2-*N*-me-Ile PSK (Scheme 2A), dipeptide 21 was coupled to H₂N-Tyr(SO₂ONp)-Thr(*t*Bu)-Gln(Trt)-Wang resin to afford the desired pentapeptide. For 4-*N*-me-Thr PSK (Scheme 2B), dipeptide 22 was coupled to H-Gln(Trt)-Wang resin and was subsequently further elongated with Fmoc-Ile-OH and Fmoc-Tyr(SO₂ONp)-OH to afford the desired pentapeptide. Sequential Fmoc-deprotection, acidic cleavage and partial deprotection, initial purification, SO₂ONp-deprotection, final purification, and lyophilization of the products afforded pure PSK analogues containing 2-*N*-me-Ile, in PSK-12, and 4-*N*-me-Thr, in PSK-13.

Protoplast proliferation assay with the synthesized PSK analogues. Chemically synthesized PSK-1 and its analogues PSK-2–13 were applied in a protoplast proliferation assay in order to investigate their ability to induce cell division of *Brassica oleracea* protoplasts. For this, protoplasts were obtained from leaves of *in vitro* grown *Brassica oleracea* plants, after which they





Scheme 2 (A) SPPS of 2-Nme-Ile PSK (PSK-**12**) using Fmoc-Tyr(SO₂ONp)-Nme-Ile-OH dipeptide **21**, synthesized in solution-phase, coupled to H-Tyr(SO₂ONp)-Thr(tBu)-Gln(Trt)-Wang resin using HATU, DIPEA, and DMF. Acidic cleavage, with TFA cocktail (95% TFA; 2.5% TIS; 2.5% H₂O, 1.5–2 h), followed by Np deprotection and purification yields pure PSK-**12**. (B) SPPS of 4-Nme-Thr PSK (PSK-**13**) using Fmoc-Tyr(SO₂ONp)-Nme-Thr(tBu)-OH **22**, synthesized in solution-phase, coupled to H-Gln(Trt)-Wang resin using HATU, DIPEA, and DMF. Respective coupling of Fmoc-Ile-OH and Fmoc-Tyr(SO₂ONp)-OH, acidic cleavage, with TFA cocktail, and subsequent Np deprotection and purification yields pure PSK-**13**.

were embedded in alginate and subsequently treated with culture medium that was supplemented with 100 nM of one of the synthetic constructs PSK-1–13, or no peptide (NP). While few calli were formed in the absence of a peptide (NP), treatment with PSK (PSK-1) resulted in significantly increased callus formation of *Brassica oleracea* red cabbage (Fig. 3A).

For the PSK analogues, the number of calli formed were counted and compared to the number of calli formed when exposed to the native peptide PSK in the same experiment. We note that PSK-5, PSK-11, and PSK-12 were also tested on red cabbage, but those results were not sufficiently reliable to

include in this graph. We found that exposing protoplasts derived from red cabbage to 2-*allo*-Ile PSK (PSK-10) and 4-Nme-Thr PSK (PSK-13) lead to calli numbers that were significantly higher than those counted for PSK ($P < 0.01$) (Fig. 3B). Interestingly, treatment of the protoplasts with Ac-PSK (PSK-3) or N₃-PSK (PSK-4) resulted in similar calli formation as compared to the native peptide (Fig. 3B), suggesting that the N-terminus tolerates these modifications. Notably, when 2-Ile is substituted for 2-Leu PSK (PSK-6) or 2-Ala (PSK-7), or when 4-Thr is substituted for 4-Ala (PSK-9), calli formation was similar to that in the absence of hormone (Fig. 3B). Higher



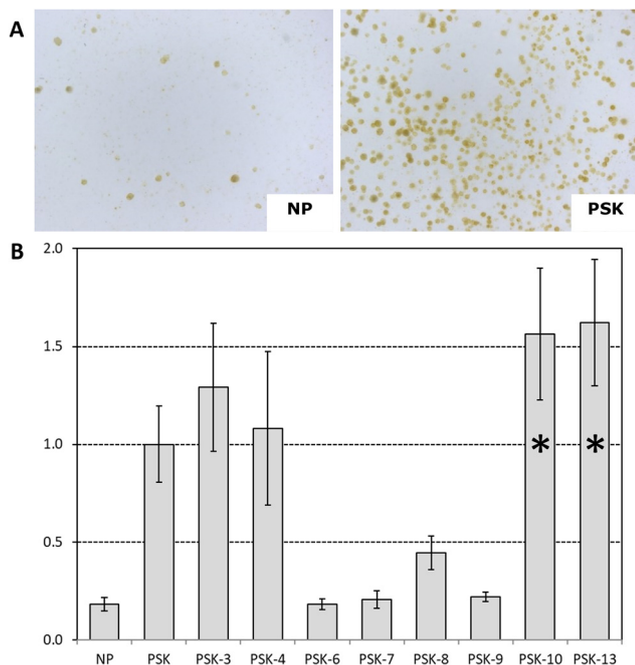


Fig. 3 (A) Representation of formed calli from *Brassica oleracea* (red cabbage) protoplasts treated with no peptide (NP, left) or 100 nM phyto-sulfokine (PSK, right). (B) Testing of N-terminal modified PSK and enantiomers of PSK for bioactivity. Different PSK analogues were tested at 100 nM and compared relative to the activity of native PSK for their ability to induce cell division, resulting in callus, in protoplasts from red cabbage. The number of calli formed was counted and compared relative to the number of calli formed with native PSK (set at 1.0). Only PSK-10 and PSK-13 outperformed the activity of PSK ($P < 0.01$). Error bars indicate the standard deviation over the number of calli in three biological replicates. NP is no peptide.

than background numbers of calli were counted when protoplasts were exposed to Asn-PSK (PSK-8), although not as much as with native PSK. In *Brassica oleracea* white cabbage protoplasts, similar observations for amino acid substituted analogues PSK-6–9 were found (Fig. S19, ESI[†]). For 2-*N*-methyl-Ile PSK (PSK-12), no increase in formed calli is observed when compared to the absence of peptide (Fig. S19, ESI[†]), while 4-*allo*-Thr PSK (PSK-11) resulted in an increased number of calli, although less than with native PSK.

Evaluation of the structures of the synthetic mutants and with respect to the observed protoplast proliferation, structure-activity relationships (SAR) could elucidate essential features for the bio-activity of the PSK peptide. First, the presence of 2-Ile and 4-Thr seems essential for the biological activity. While 4-Asn PSK (PSK-8) is still capable of inducing callus formation, the relative activity is only 30% compared to PSK (PSK-1). *N*-Terminal modifications Ac-PSK (PSK-3) and N_3 -PSK (PSK-4) resulted in similar outcomes compared to PSK (PSK-1). This indicates that the pentapeptide pharmacophore still remains bio-active while containing an extended *N*-terminal modification, which could be useful for peptide labeling. Indeed, the crystal structure of the PSKR-PSK complex indicates the presence of a cavity at the *N*-terminus, which can be occupied by the respective PSK analogues. Changes in stereochemistry

resulted in weaker bio-activity for 4-*allo*-Thr PSK (PSK-11) but higher activity for 2-*allo*-Ile PSK (PSK-10) when compared to the parent PPH PSK, with 0.7 and 1.3 relative activity, respectively. Considering the presence of a *N*-methylated amide backbone, we found that methylation of the Ile amide (PSK-12) does not lead to bio-active compounds, whereas shifting the methyl to the Thr residue, as in 4-*N*-methyl-Thr PSK (PSK-13), leads to the most active PSK analogue in our library (> 1.5 relative activity compared to PSK).

The resolved crystal structure of the PSK-PSKR1^{LRR} indicates hydrogen bonding and van der Waals packing interactions, where the most prominent contributions arise from the two sulfate moieties.²⁸ Additionally, residues from the inner surface of the PSKR1^{LRR} forms hydrogen bonds with the main chain of PSK. Polar interactions are observed in the amide moiety between Tyr-1 and Ile-2 of PSK with Thr³⁹⁸, between the Thr-4-OH of PSK and Ser³⁷⁰, and between a PSK backbone NH of Thr-4 with Ser³⁷². As the 4-*N*-methyl-Thr analogue (PSK-13) does not enable the hydrogen bond between Ser³⁷² and the PSK backbone, the higher biological activity with respect to plant protoplast regeneration shows that this interaction between the PSK analogue and the PSKR is not crucial to elicit a biological response. Changing the stereochemistry of C β in 2-Ile (as in PSK-10) resulted in increased bio-activity, which could indicate an increase in polar interaction with Thr³⁹⁸, resulting in stronger receptor activation. In contrast, the reduced bio-activity found for 4-*allo*-Thr PSK (as in PSK-11) could be directed to weaker Thr-4-OH with Ser³⁷⁰ interactions and thus less receptor activation and weaker bio-activity of the peptide.

Conclusions

We report the synthesis of *N*-methylated amino acids, their incorporation in PSK analogues, and a set of other substituted PSK-like peptides. We selected a set of peptides including *N*-terminal modifications, amino acid substitutions, diastereomers and *N*-methylation. Solution-phase synthesis was combined with solid-phase peptide synthesis and resulted in a set of 13 peptides. For the *N*-methylated analogues PSK-12 and PSK-13, building blocks 15 and 18 were synthesized with an *N*-alkylation procedure using methyl iodide. The building blocks were coupled in solution to Fmoc-Tyr(SO₂ONp)-OH using HATU as activator to obtain dipeptides 21 and 22. The chemically synthesized PSK analogues were tested in *Brassica oleracea* protoplast proliferation assays to compare relative activity to the native peptide. Bio-activity of most analogues was reduced or even abolished. Only 2-*allo*-Ile PSK (PSK-10) and 4-*N*-methyl-Thr PSK (PSK-13) displayed a significant increase in bio-activity. These findings indicate that small chemical modifications in PPHs affects bio-activity which arises from weaker or stronger polar interactions in the binding pocket of PSKR1. Potentially, making combinations of PSK-10 and PSK-13 might lead to an even more potent artificial analogue. With these assessments, a bio-active PPH could be designed, synthesized and biologically evaluated in order to improve bio-activity in different recalcitrant crops.



Author contributions

BA, RGHI and MF conceived the project, JvdS and DS prepared the chemical compounds, MF performed the biological activity studies and data curation. JvdS and BA wrote the manuscript, DS, RGHI, and MF reviewed and edited the manuscript, MF and RGHI obtained funding for this project.

Conflicts of interest

There are no conflicts to declare.

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