The effect of seaweed extract on tomato plant growth, productivity and soil

Hashmath Inayath Hussain1 · Naga Kasinadhuni2 · Tony Arioli1,3

Received: 28 October 2020 / Revised and accepted: 6 January 2021 / Published online: 1 February 2021 © The Author(s) 2021

Abstract

This study investigated the effects of seaweed extract (SWE) made from the brown algae Durvillaea potatorum and Ascophyllum nodosum on plants and soil. The application of SWE to soil growing tomato plants showed dual effects. SWE comprehensively improved tomato plant growth (flower clusters, flower number, fruit number, root length, root and shoot dry weight, SPAD) and increased plant productivity (yield and quality). Similarly, SWE application effected soil biology at the soil root zone by increasing total bacterial count and available soil nitrogen and impacting bacterial community diversity with an increase in certain bacterial families linked to soil health. A broader understanding of the effects of SWE on the plant-soil ecosystem may offer breakthrough approaches for sustainable food production.

Keywords Microbes · Productivity · Quality · Seaweed extract · Rhizosphere · Soil · Tomatoes

Introduction

The role of agricultural biostimulants in food production worldwide is expanding, especially those made from seaweed extracts (SWE). Biostimulants made from seaweeds have properties that improve plant growth, yield and quality and increase tolerance to abiotic and biotic stresses, as well as improving nutrient use (Arioli et al. 2015; Shukla et al. 2019). For example, field studies have demonstrated that SWE, applied at relatively low rates as foliar sprays or soil fertigation, increase yield among diverse crops such as wine grapes, sugarcane, strawberries and vegetables (Khan et al. 2009; Arioli et al. 2015; Shukla et al. 2019). Studies using Arabidopsis, soybean and tomato plants have demonstrated that the application of SWE can improve plant tolerance to abiotic stresses such as drought, freezing and salt (Nair et al. 2012; Martynenko et al. 2016; Santaniello et al. 2017; Goni et al. 2018; Jithesh et al. 2019). At a molecular level, laboratory studies have demonstrated that certain SWE have an intriguing capacity to activate plant defence response systems including the Systemic Acquired Resistance mechanism (Goni et al. 2016; Islam et al. 2020; Omidbakhshfard et al. 2020). In addition, soil-applied SWE provide a wide range of nutrients for soil microbiota utilisation.

The microbiota at the interface between plant roots and soil (rhizosphere) have been linked to improved plant growth and health (Vives-Peris et al. 2020). For example, microbiota at the rhizosphere can protect plants against pathogens (Mendes et al. 2011; Rolfe et al. 2019), help withstand stress to adverse environmental conditions (Rolli et al. 2015; Hartman and Tringe 2019) and alter plant development, for example, by influencing the timing of plant flowering (Lu et al. 2018; Olanrewaju et al. 2019). The rhizosphere environment is highly interactive because plants manipulate the composition of their root exudates (Olanrewaju et al. 2019). Consequently, a broader understanding of the effect of SWE on plant and soil biology is needed to advance agriculture.

We hypothesise that the soil application of a SWE made from two brown seaweeds (Ascophyllum nodosum and Durvillaea potatorum) would enhance plant growth and soil biology and would influence the plant-soil ecosystem. We chose an extract made from two seaweeds because transcriptomics research has shown an expanded set of plant gene responses compared with the extracts made from single-origin seaweeds (Islam et al. 2020). For new insights, we investigated tomato plant growth and soil biology parameters concurrently.
We designed a greenhouse experiment using tomato plants growing in pots and receiving soil applications of SWE at regular intervals and scientific assessments focused on flowering and fruit harvest and quality. Soil biology was investigated using metagenomics sequencing at the commencement of flowering and phenotypic assessments followed for flowering, fruit number, root growth, leaf chlorophyll and fruit yield and quality. Flowering was chosen because the transition from vegetative to propagative growth is a critical plant development stage for field crops.

The aim of the study was to determine the effect of SWE concurrently on (i) plant growth and productivity and (ii) the rhizosphere microbiology.

Method

Seedling establishment

Tomato seeds (Lycopersicon esculentum Var. Grosse Lisse) were used in this study (from Seeds and Select Organic, Queensland, Australia). Commercial seedling trays were filled with commercially purchased seed raising mix (Debco Seed & Cutting Mix, Victoria, Australia). The tomato seeds were planted into the seedling trays at a rate of one seed per cell. The trays were placed at 21 °C for 4 weeks in a temperature-controlled greenhouse for germination and the soil mix was kept moist as required.

Experimental plant trial

A greenhouse potting trial was conducted using a commercially available seaweed extract manufactured by alkaline hydrolysis from Durvillaea potatorum and Ascophyllum nodosum and fortified with calcium at 1% (w/v) (Seasol Plus Calcium, Melbourne, Australia). Detailed information about the seaweed extract such as seaweed composition, mineral content and manufacture has been published previously (Arioli et al. 2015). A seaweed extract with calcium was selected because tomato fruits are susceptible to calcium disorders such as blossom end rot and yellow shoulder disorder. Four-week-old tomato seedlings were transplanted to pots (400 mm) containing Premium Potting Mix containing controlled-release fertiliser granules (N16%, P3%, K13%; Grow Better Garden Products, Victoria, Australia). Twenty pots containing the tomato seedlings were transferred into the greenhouse (8m × 8m × 6m, L × W × H) exposed to local ambient spring and summer conditions and not temperature controlled. The plants were climate hardened for a 1 week before the first treatment application. For application, the SWE was applied at 1:400 dilution and water was used for the untreated control. The SWE dilution rate was selected because of field and greenhouse studies demonstrating the efficacy at this dilution (Mattner et al. 2013, 2018; Arioli et al. 2020) and because of root growth testing (Arioli et al. 2015). The SWE treatment was applied to the soil in each pot using a piped irrigation system fitted with (i) a dosing unit (Dosatron, Australia) set to deliver the required dilution and (ii) a frequency controller set to deliver twice daily doses of 250 mL (to each pot). The trial was conducted as a randomised complete block design with ten replicates for each treatment, with 20 plants in total. Three plant developmental time points were used for the different assessments: flower initiation, flowering and harvest, which occurred at 29, 64 and 95 days after potting (DAP). The flower initiation day (29 DAP) was determined by monitoring the plants daily for the appearance of the first open flower.

Plant growth measurements

Plant growth was investigated at flowering and harvest using five replicates (plants) per treatment (ten plants in total) by assessing flower cluster and fruit number (Hussain et al. 2017), root length (Doerner 2008), shoot dry weight and root dry weight (Huang et al. 2017), and tomato leaf chlorophyll was assessed using a soil plant analysis development meter SPAD (SPAD-502, Konica Minolta, Japan) (Yao et al. 2020). Shoot and root samples were dried at 70 °C in a drying oven and dry weights were measured using analytical scales (Westlabs Model 663-543) after 72 h.

Soluble solids concentration

Tomato fruits harvested at 95 DAP were sent for fruit quality analysis (Freshhort, Victoria, Australia). Tomato fruit soluble solid concentration (SSC) in degree Brix (°Brix) were measured by cutting a ripe section from the cheek of each fruit within a replicate, placing all sections in a plastic bag, hand squeezing the sections and pouring 10 mL of extracted juice through a fine-mesh sieve into a 15-mL Eppendorf tube. Approximately 0.5 mL of juice was then poured from the tube onto the lens of a temperature-compensated digital refractometer (ATAGO PAL-1) with a measurement accuracy of ± 0.2 °Brix. The refractometer was calibrated with distilled water prior to SSC measurements of each replicate. The measurement was repeated, and the mean SSC in °Brix was calculated from the two measurements per replicate.

For each replicate, the remaining juice was frozen at −20 °C. All juice samples were thawed at 20 °C, and 3 mL of juice from each sample was diluted in 5 mL of distilled water once the juice temperature in all samples was above 15 °C. The titratable acidity of each sample was then measured via endpoint titration to pH 8.2 with 0.1 M NaOH using an automatic titrator (Steroglass Titre X) and AS23 Micro autosampler. The mean titratable acidity for each replicate was calculated as grams of citric acid equivalent per litre of juice using the NaOH titre volume.
Soil testing

At flower initiation (29 DAP), the soil growing the potted tomato plants were assessed for microbiology using metagenomics sequencing, available soil nitrogen and the number of bacterial cells. The soil samples (approximately 400 g) were carefully collected from individual pots by selecting soil in contact with the central tomato root zone (rhizosphere) at a depth of 10 cm and avoiding the bulk soil zone (non-rhizosphere) that was not in contact with the roots. For metagenomics testing, ten individual soil samples were collected in total, five individual soil replicates for each treatment. The soil in contact with the roots (root zone soil) was placed into separate zip lock bags using a clean spatula. The soil samples, 50 g for each replicate, was sent to the Australian Genomic Research Foundation (AGRF) for metagenomics shotgun sequencing (NGS) and analysis. DNA was extracted from the soil using DNeasy PowerSoil Pro Kit (Qiagen, Germany). DNA libraries were prepared for each replicate using the Illumina TruSeq Nano DNA Library Prep Kit (Illumina, USA) with a 350 bp insert library and sequenced using the Illumina HiSeq 2500 platform (Illumina, USA) and 100 base pair (bp) paired read chemistry. The root zone soil samples, containing 200 g soil, five individual replicates per treatment, were sent for available soil nitrogen testing (SWEP Laboratories Pty Ltd, Melbourne, Australia). Available soil nitrogen refers to the nitrogen that is available for plants to absorb and measured as nitrate-nitrogen (NO₃-N) only. The method involves a calcium chloride extraction and UV-visible colorimetric assay at 550 nm (Industry Method 7C2, Rayment and Lyons 2011). The root zone soil samples, containing 50 g soil, ten individual replicates per treatment, were sent for a standardised bacterial plate count test (SWEP Laboratories Pty Ltd, Melbourne, Australia). The test measures the number of bacterial colony forming units per gram of soil (cfu g⁻¹).

Bioinformatics analysis

To build the taxonomical profiles from whole metagenomics sequencing dataset, the fastq files containing the raw reads were imported into CLC Genomics Workbench v20.0 (QIAGEN, Aarhus C Denmark, http://www.clcbio.com), using the following options: Illumina import, paired-reads, paired-end (forward-reverse) and minimum distance of 1 and a maximum distance of 1000 (default). Microbial Genomics Module (MGM) available as a plugin in the CLC Genomics Workbench was used to perform quality control (QC) and taxonomical profiling of the raw reads. The trimmed sequences were aligned to a CLC Microbial Reference Database (Version June 2019) from Genbank (250,000 basepairs). The permutational multivariate analysis of variance (PERMANOVA) test was conducted using CLC Genomics Workbench version v20.0 between two treatments based on Bray–Curtis distance among all taxonomic levels of each community. Beta diversity using principal coordinate analysis (PCoA) plots were generated among samples using CLC Microbial Genomics Module.

Statistical analysis

For the plant assessments the treatment effect on plants was analysed using a randomised block design experiment setup and Agronomic Research Management software (ARM, South Dakota, USA). GraphPad Prism statistical software (GraphPad, USA) was used to perform the statistical analysis. A t-test between the two treatments was used to show the statistical significance of alpha 0.05. Means followed by the same letter do not significantly differ. For the fruit quality (SSC) assessment, data for the treatment effect on fruit quality were analysed as a completely randomised design experiment using two-way ANOVA in GenStat 17 (VSN International Ltd., UK). Violations of the ANOVA assumption of normality in the data, such as non-normality (Skewness, Kurtosis) or heterogeneity of treatment variance, were assessed using residual error plots, skewness and kurtosis tests of normality and Bartlett’s test of homogeneity of variance. Where necessary, the appropriate data correction transformation was applied to data prior to ANOVA based on optimal values of lambda calculated from Box-Cox analysis in Genstat.

For the soil microbial analysis, at the family level, differential abundance analysis was done using the CLC Genomics Workbench v20.0. Post-QC and taxonomical profiling of the individual fastq files from each treatment, the CLC MGM tool was used to perform the differential abundance analysis between the two treatments. This CLC MGM tool performed a generalised linear model differential abundance test on the samples defined by metadata. The tool modelled each feature based on a separate Generalised Linear Model, where it assumed that the abundances followed a negative binomial distribution. The Wald test was used to determine significance between the treatments.

Results

Effects of SWE treatment on plant growth and productivity

At the flowering (64 DAP) and harvest (95 DAP) stages, plant growth was assessed by determining (i) the number of flower clusters, (ii) fruit number, (iii) shoot dry weight and (iv) root dry weight. A statistical analysis found a significant increase in all the plant growth parameters in the SWE treatment relative to the untreated control (P < 0.05, Fig. 1; P = 0.0049 for day 64 DAP number of flower clusters, P = 0.0463 for day 95...
DAP number of flower clusters, \( P = 0.0014 \) for day 95 DAP number of fruit, \( P = 0.0289 \) for day 64 DAP shoot dry weight, \( P = 0.0005 \) for 95 DAP shoot dry weight, \( P = 0.0149 \) for day 64 DAP root dry weight, \( P = 0.0207 \) for 95 DAP root dry weight) except for the average fruit number at the flowering stage. At flowering, the fruit number was found to be larger in the SWE-treated plants but not significantly.

Additional plant assessments were undertaken at the flowering stage: (i) root length, (ii) flower number per plant and (iii) chlorophyll content. Statistical analysis found a significant increase in all the plant growth parameters due to the SWE treatment relative to the untreated control (\( P < 0.05 \), Fig. 2; \( P = 0.0475 \) for day 64 DAP root length, \( P = 0.0008 \) for day 64 DAP fruit number per plant, \( P = 0.0220 \) for chlorophyll content).

Plant productivity was assessed by measuring fruit yield per plant. Statistical analysis found a significant increase in fruit weight per plant due to the SWE treatment relative to the untreated control (\( P = value < 0.05 \), Fig. 3; \( P = 0.0075 \) for day 95 DAP fruit weight per plant). Fruit quality was assessed by measuring SSC. Fruit harvested from SWE-treated plants had a mean SSC of 5.6 °Brix, and fruit from untreated plants had a mean SSC of 5.2 °Brix. Statistical analysis found that the increase (8%) in SSC due to the SWE treatment relative to the untreated control was significant at \( P < 0.1 \). The increases in the overall plant growth phenotypes are in alignment with productivity and quality increases.

**Soil microbiology and available soil nitrogen**

At the flower initiation stage, soil collected at the tomato root zone were assessed for bacterial numbers and available soil nitrogen. Statistical analysis found a significant increase in the number of soil bacteria (\( P \)-value < 0.05, Fig. 4; \( P = 0.0097 \)) and the available nitrogen in the soil (\( P \)-value < 0.05, Fig. 5; \( P = 0.0492 \)) due to the SWE treatment relative to the untreated control. Also, we noted that the SWE treatment increased the total active fungi cfu g\(^{-1}\) (8.58 × 10\(^6\) vs 5.97 × 10\(^6\) ), Total active yeast cfu g\(^{-1}\) (1.02 × 10\(^5\) vs 7.20 × 10\(^4\) ) and total plant growth-promoting bacteria cfu g\(^{-1}\) by 2.2-, 1.4-, 1.4-fold, respectively, but these results were not statistically significant (at \( P < 0.05 \)).

**Soil microbial composition**

The root zone soils collected at the flower initiation stage were assessed for microbial composition. Taxonomic profiling was used to investigate the abundance profiles at the domain and microbial levels. For absolute abundance at the domain level,
the average number of reads for both the treatments aligned to Bacteria (97.78%) followed by Eukaryota (2.03%) and Archaea (0.19%). At the microbial level, the stacked bar chart (Fig. 6) shows the relative abundance profiles after the alignment of the raw reads to the microbial reference database for the treatments (untreated control and SWE). The taxonomical profiling of the potting mix soil revealed 83 bacterial families between the treatments. Nakamurellaceae was the most abundant (38%) in the potting mix treated with SWE, and it was absent in the untreated control samples while Mycobacteriaceae, Bradyrhizobiaceae, Thermonosporaceae inter alia were common between the treatments. Differences in microbial diversity and abundance between the treatments were confirmed by additional statistical analysis.

**Beta analysis—microbial communities cluster distinctly between treatments at flower initiation**

The Bray-Curtis distance metric was used to determine multivariate sample distances, which were visualised through the ordination of a principal coordinate analysis (PCoA). The PCoA shows the control samples clustered relative to SWE treatment samples (S1). PCo1 was found to account for the majority of the variation (74%).

**Differential abundance analysis between the treatments**

Analysis using permutational multivariate analysis of variance (PERMANOVA) confirmed substantial differences in the structures of the bacterial communities between the SWE and control
potting mix soils ($P \leq 0.05$; Table 1). Among the bacterial taxonomic families detected, 38 complete bacterial taxonomic were significantly different between the SWE- and control treated potting mixes. The SWE treatment enriched 14 bacterial families that showed a log fold change greater than or equal to 2 ($P \leq 0.05$; S2) and depleted 14 bacterial families that showed a log fold change lesser than or equal to $-2$ ($P \leq 0.05$; S3).

Nakamurellaceae, Cytophagaceae Sterolibacteriaceae, Archangiaceae, Moraxellaceae, Brevibacteriaceae, Chlorobiaceae and Dermatophilaceae were the top 8 bacterial families significantly enriched in the potting mix soil treated with SWE (S2).

**Discussion**

**The effect of SWE on plants**

This study found that the effect of SWE on plants significantly increased numerous plant growth parameters: number of flower clusters and flowers, fruit number, dry weight of foliage and roots and root length. SWE had a positive effect on tomato plant yield and the total soluble solid concentration in tomato fruit. SWE application was found to significantly increase the chlorophyll levels in the tomato leaves at the flowering stage,
as determined by SPAD testing, an assessment used to assess crop nitrogen status (Yuan et al. 2016; Gholizadeh et al. 2017). Collectively, the effects due to the SWE are agronomically significant as prerequisites for increasing tomato productivity and fruit quality.

In our study, it is unlikely that the plant responses are due to a fertiliser effect of the SWE since the experiment included fertiliser in the potting soil (both in the control and SWE treatment). The applied fertiliser was substantially higher than the nutrients in the diluted SWE that was used. Furthermore, field research using the same SWE used in this study found that repeated soil applications did not change the soil macronutrient content (Mattner et al. 2013).

Our results, using an extract made from dual seaweeds, are consistent with published research demonstrating that SWE can improve plant growth and productivity (Khan et al. 2009; Shukla et al. 2019). Tomato studies using SWE made from Ascophyllum nodosum or Sargassum horneri demonstrated these SWE can enhance different aspects of plant growth. For example, Ali et al. (2016) reported that an SWE made from A. nodosum increased the number of flower clusters and fruit yield in tomato greenhouse and field trials. Renaut et al. (2019) found an SWE made from A. nodosum increased fruit number, shoot dry weight and root dry weight. Yao et al. (2020) found that an SWE made from S. horneri increased tomato yield and enhanced SPAD, chlorophyll content and photosynthesis capacity. More broadly, the soil application of SWE made from dual seaweeds (A. nodosum and Durvillaea potatorum) has been reported to increase growth and yield parameters in broccoli, sugarcane and strawberry field trials (Mattner et al. 2013, 2018; Arioli et al. 2020). Santaniello et al. (2017) used another SWE to report the effects on photosynthetic performance in Arabidopsis. Islam et al. (2020) reported SWE increased root growth in Arabidopsis. Given the effects of SWE reported across many different farming and growth environments using different crops, this implies their mode of action is conserved with an inherent plasticity when used as a foliar spray or soil application.

The effect of SWE on Soil

We investigated the effect of SWE, as a soil application, on the soil microbiology at the onset of flowering and focused on the characterisation of the soil bacteria located at the rhizosphere rather than the bulk soil. Flowering is a critical transition development point where plants reallocate metabolism and require nutrition, especially potassium and calcium, for plant and fruit growth (Lee et al. 2017). In addition, Lu et al. (2018) reported the influence of rhizosphere microorganisms on the timing of plant flowering. Therefore, we utilised the flowering stage as an important juncture linking soil biology and plant productivity.

At flower initiation we found evidence for changes in the soil microbiology before the plant phenotype assessments were undertaken. Firstly, we found that the recurring applications of SWE increased the total number of bacteria in the soil. From an ecosystem perspective, the increase in soil bacteria is significant because the additional bacteria provide cellular matter to decompose and supply the soil carbon and nitrogen cycles. The SWE also increased the available nitrogen content in the soil, which highlights an uncharacterized influence on the bacterial communities that mineralize soil nitrogen. The differences in the soil bacterial communities are supported by a Bray-Curtis analysis confirming that the SWE treatment created diverse and distinct soil populations. Collectively, these soil changes are relevant to agriculture because nitrogen is an essential nutrient for plant growth, and the cellular matter is vital to the soil organic matter reservoir in depleted farming soils.

Our metagenome analysis revealed a plethora of microbial communities at the root-soil rhizosphere. Taxonomical profiling revealed three major microbial domains: Bacteria, Archaea and Eukaryota. The Bacteria domain dominated in relative abundance, compared with the Archaea and Eukaryota domains. Soils rich in nitrogen fertiliser have been found to harbour dominate bacterial populations (Jia and Conrad 2009; Monreal et al. 2018).

Differential abundance analysis found that the SWE treatment enhanced 14 bacterial families and reduced 14 bacterial families. Our results are in general agreement with reports that showed an increased number of bacterial families in soils treated with SWE (Khan et al. 2012, 2013; Shukla et al. 2019). Also, SWE has been shown to increase fungal families in soils amended with organic matter (Renaut et al. 2019). Upon review of the 14 bacterial families that were significantly enriched by SWE treatment, we found certain members to have functions associated with enhanced soil properties and plant growth (Muhammad and Shahida 2006; Sgroy et al. 2009; Miransari 2013; Rolli et al. 2015; Shan et al. 2018; Rana et al. 2019). For example, some of the species that belong to the family Nakamurellaceae have been associated with symbiotic endophytes (Whitman 2015; Singh and Dubey 2018; Nouioui et al. 2019). The Cytophagaceae and Archangiaceae families have members known to digest polysaccharides for growth and for cycling carbon (Dawid 2000; McBride et al. 2014). The Sterolibacteriaceae family have members associated with enhancing plant tolerance to heavy metals (Chain et al. 2003). The families that were found to be reduced in abundance tended to be associated with marine habits or had no ascribable relevance except for the Beijerinkiaceae family due to their ability to grow in habitats scarce in nitrogen. Although metagenomic analysis is a powerful technique, additional analysis is needed to undercover the functionalities of the communities and their fit within the agricultural ecosystem.
Effects of SWE at the ecosystem level

Our results demonstrate the application of SWE to soil-influenced plant growth and soil characteristics. However, plants and soils are part of a broader plant-growing ecosystem involving a multitude of interactions, natural nutrient cycles and an interrelated food web.

Plants growing in soils develop intimate associations with the soil microbial community, particularly at the rhizosphere (Hartman and Tringe 2019). These relationships can take multiple forms leading to beneficial, deadly or commensal interactions. Plants use their carbon-rich root exudates to influence their interactions in the ecosystem (Thakur and Geisen 2019). Plants can sequester microbial communities that facilitate plant nutrient uptake, the suppression of biotic stress, the activation plant defence responses and enhanced plant tolerance to abiotic pathogens (Olanrewaju et al. 2019; Rolfe et al. 2019; Vives-Peris et al. 2020).

Soil biogeochemistry includes an important role for microbes (Neal et al. 2020). Organic matter residing in soil is processed by microbes and acts to bind mineral and colloids particles. Microbes associated with these surfaces have distinct gene assemblages (Hartman and Tringe 2019; Neal et al. 2020). This type of research has led to the proposal that soil-microbe systems have self-organising capabilities with carbon flux being the critical factor in soil systems (Neal et al. 2020).

An important insight from our study is that the application of SWE to the soil will resonate at the ecosystem level when plants are growing. We hypothesize that plants with improved growth and root systems, due to SWE application, are supplying relatively more exudates into the soil, increasing microbial biomass, thereby benefiting the overall soil biogeochemical cycles. This proposal is consistent with research demonstrating that plant species with nutrient acquisitive properties regulate a plant-soil carbon-nitrogen feedback system operating at the soil root interface (Henneron et al. 2020; Wei et al. 2020). Despite the complexity, further research from the agroecosystem perspective is required to elucidate the collective benefits of using SWE for sustainable agriculture and food production.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s10811-021-02387-2.

Acknowledgements We recognise and thank Jeffery Lang for his excellent technical assistance growing the experimental plants and conducting the greenhouse assessments.

Authors’ contributions All co-authors contributed to the final version of the work and approved the manuscript for publication. HH: leading the experimental research, interpretation of data and writing the manuscript; TA: interpretation of data and writing the manuscript; NK: bioinformatics and review of the manuscript.

Funding Funding for the experimental research was provided by Seasol International Pty Ltd (Australia). No funding was received to assist with the preparation or submission of this manuscript.

Compliance with ethical standards

Conflict of interest Seasol International (SI) is the manufacturer of seaweed extract (Seasol) in Australia. TA and HH are scientific researchers at Seasol International. TA is an adjunct Associate Professor at Deakin University. The authors declare that the research was conducted in the absence of any commercial or financial relationship that could be construed as a potential conflict of interest.

Consent for publication All the authors provide consent to publish.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

Ali N, Farrell A, Ramsubhag A, Jayaraman J (2016) The effect of Ascophyllum nodosum extract on the growth, yield and fruit quality of tomato grown under tropical conditions. J Appl Phycol 28:1353–1362
Arioli T, Mattner S, Winberg P (2015) Applications of seaweed extracts in Australian agriculture: past, present and future. J Appl Phycol 27:2007–2015
Arioli T, Hepworth G, Farnsworth B (2020) Effect of seaweed extract application on sugarcane production. Proc Aust Soc Sugar Cane Technol 42:393–396
Chain P, Lamerdin J, Larimer F, Regala W, Lao V, Land M, Hauser L, Hooper A, Klotz M, Norton J, Sayavedra-Soto L, Arciero D, Hommes N, Whittaker M, Arp D (2003) Complete genome sequence of the ammonia-oxidizing bacterium and obligate chemolithoautotroph Nitrosomonas europaea. J Bacteriol 185:2759–2773
Dawid W (2000) Biology and global distribution of myxobacteria in soils. FEMS Microbiol Rev 24:403–427
Doerner P (2008) Phenotypic analysis of Arabidopsis mutants: quantitative analysis of root growth. Cold Spring Harb Protoc 2008:prot4960
Gholizadeh A, Saberioon M, Boruvka L, Wayayok A, Soom M (2017) Leaf chlorophyll and nitrogen dynamics and their relationship to lowland rice yield for site-specific paddy management. Int Proc Agric 4:259–268
Goni O, Fort A, Quille P, Mckewen P, Spillane C, O’Connell S (2016) Comparative transcriptome analysis of two Ascophyllum nodosum extract biostimulants: same seaweed but different. J Agric Food Chem 64:2980–2989
Goni O, Quille P, O’Connell S (2018) Ascophyllum nodosum extract biostimulants and their role in enhancing tolerance to drought stress in tomato plants. Plant Physiol Biochem 126:63–73
Hartman K, Tringe S (2019) Interactions between plants and soil shaping the root microbiome under abiotic stress. Biochem J 476:2705–2724
**Publisher's note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.