# A commercial seaweed extract structured microbial

- 2 communities associated with tomato and pepper roots
- 3 and significantly increased crop yield
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- 8 B3B 1X8
- Seaweeds have been used as a source of natural fertilizer and biostimulant in agriculture for centuries. However, their effects on soil and crop roots microbiota remain unclear. Here, we used a commercially available Ascophyllum nodosum Extract (ANE) to test its effect on bacterial and fungal communities of rhizospheric soils and roots of pepper and tomato plants in greenhouse trials. Two independent trials were conducted in a split block design to test to effect of ANE amendment. We used amplicon sequencing targeting fungal ITS and bacterial 16S rRNA gene to determine microbial community structure changes. We find that productivity parameters of root, shoot and fruit biomass were positively and significantly influenced by the amendment. In addition, a-diversity differed 17 significantly between amended and control plants, but only in some of the experimental conditions. Species composition among sites (b-diversity) differed according to the 19 amendment treatment in all four communities (fungal-root, fungal-soil, bacterial-root and 20 bacterial-soil). Finally, we identified a number of candidate taxa most strongly correlated 21 with crop yield increases. Further studies on isolation and characterization of these microbial taxa linked to the application of liquid seaweed extract may help to enhance crop yield in sustainable agro-ecosystems.
- Keywords: Stella Maris®, 16S, ITS, soil microbial diversity, Illumina MiSeq, ANE, Amplicon Sequence Variants, OTU

### 27 SIGNIFICANCE STATEMENT

While seaweeds have been used as a source of natural fertilizer and biostimulant for 28 centuries, their effects on soil microbiota remain unclear. Here, we showed that plant 29 productivity was positively and significantly influenced by the addition of a liquid Asco-30 phyllum nodosum seaweed extract. Then, we used amplicon sequencing targeting fungal 31 ITS and bacterial 16S rRNA gene to quantify Amplicon Sequence Variants and changes 32 in the microbial community structure. Both bacterial and fungal species composition differed according to the amendment treatment and we identified a number of candidate 34 taxa whose presence was correlated with plant yield increases. As such, these promising microbial candidates linked to the application of liquid seaweed extract may help to enhance crop yield in sustainable agro-ecosystems.

### 38 INTRODUCTION

Seaweeds (also known as marine macroalgae) have been used as a source of organic 39 matter and mineral nutrients for centuries, especially in coastal areas (Khan et al., 2009; 40 Craigie, 2011). Liquid seaweed extracts, developed in the 1950s in order to concentrate 41 plant growth-stimulating compounds, facilitate their usage (Milton, 1952). Today, most 42 commercially available extracts are made from the brown algae Ascophyllum nodosum, Ecklonia maxima or Laminaria spp. Unlike modern chemical fertilizers, seaweed extracts are biodegradable, non-toxic and come from a renewable resource (Dhargalkar and Pereira, 2005). Industry-funded bodies such as the European Biostimulant Industry Coalition and the United States Biostimulant Coalition have been working to accommodate biostimulants into mainstream legal architecture. These organizations extoll benefits arising from modes-of-action research, agricultural applications and positive effects on yield and quality of many commercial species (i.e. fruits, vegetables, turf, ornamentals and woody species). Legal recognition will further allow a better integration of various biostimulants, 51 including Ascophyllum nodosum Extracts (ANE) into sustainable long-term crop management programs (Craigie, 2011; Jardin, 2015).

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55 Starting in the 1990's, high quality ANE was developed and let to an increased usage by
56 farmers, in addition to an increase in cause-effect research, especially on plant diseases
57 (Jayaraj and Ali, 2015). Since then, several comprehensive reviews have described the
58 effects of seaweed extracts on agricultural plant productivity (Khan *et al.*, 2009; Craigie,
59 2010, 2011; Battacharyya *et al.*, 2015). The science points to wide-ranging effects from
60 biotic to abiotic resistance, effects on growth and development, and ultimately, to their
61 impact on plant establishment, crop yield and/or quality, and shelf life.

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At the physiological level, these extracts have been found to influence hormones levels

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that in turn, influence physiological processes even at very low concentrations (Wally *et al.*, 2013). Noted increases in the activity of superoxide dismutase, glutathione peroxidase and ascorbate peroxidase helped support the argument that ANE improve plant tolerance to oxidative stress (Ayad *et al.*, 1997; Schmidt and Zhang, 1997; Ayad, 1998; Allen *et al.*, 2001). Positive effects were also found on phytoalexin production suggesting that ANE may increase activity of these protective enzymes that target oxidizing toxins naturally emitted by disease pathogens (Lizzi *et al.*, 1998; Jayaraj *et al.*, 2008; Jayaraman *et al.*, 2010).

Improved plant stress resistance and tolerance to foliar and soil pest treatments is attributed to a cascade of various physiological reactions. ANE can impact plant-signaling mechanisms through a multitude of plant processes and cellular modifications including osmotic/oxidative stresses such as salinity, freezing and drought stress (Jithesh *et al.*, 2012). ANE can also impart drought-stress tolerance to plants by reducing stomatal conductance and cellular electrolyte leakage (Shotton and Martynenko, unpublished data; Spann and Little, 2011). These results indicate that ANE can influence cellular membrane maintenance leading to a higher tolerance for various osmotic stresses and can mitigate oxidative damage.

Although published evidence has detailed systemic effects of ANE on plant development, the effect of ANE on the biology of the rhizosphere is still largely unknown. The
soil rhizosphere harbors a large microbial biodiversity that contributes to the aggregation
of particles, enhances nutrient cycling and delivery to plants, degrades toxic substances,
allows better soil water retention and plays a role in plant disease management. It has
been suggested that the plant immune system is composed of inherent surveillance systems that perceive several general microbial elicitors, which allow plants to switch from
growth and development into a defense mode (Newman *et al.*, 2013). This may allow
the plant to avoid infection from potentially harmful microbes. An examination of sus-

tainable products that can positively influence microbial interactions between plant roots and soil biota will in turn help to further understand soil borne plant-pathogens competition dynamics. The effect of ANE on the bacterial profile suggests that ANE applications increased strawberry root and shoot growth, berry yield, and rhizosphere microbial diversity and physiological activity (Alam *et al.*, 2013). Similar results were found in sandy loam soils as Alam and colleagues (2014) showed a strong relationship between carrot growth, soil microbial populations and activity.

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The recent development of culture-independent molecular techniques and high throughput sequencing should permit to circumvent the inherent biases of culture-based approaches by targeting the ubiquitous component of life, DNA. In turn, this will help to
identify a larger proportion of the microbial diversity and lead to a better understanding of the soil microbial response to seaweed extract. DNA barcoding targeting specific
regions of the genome (e.g. ITS: fungi, 16s ribosomal genes: bacteria) are now regarded
as a prerequisite procedure to comprehensively document the diversity and ecology of
microbial organisms (Toju *et al.*, 2012; Klindworth *et al.*, 2013).

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Here, the general objective was to quantify the impact of a commercial seaweed extract on plant growth and test how the fungal and bacterial communities responded to the addition of these extracts. We also aimed to identify specific taxon positively correlated with increases in plant productivity following ANE amendments. We hypothesized that the 111 addition of liquid seaweed extracts would improve productivity and alter significantly 112 the fungal and bacterial communities. We used a commercially available ANE, Stella 113 Maris®, developed by Acadian Seaplants Ltd (NS, Canada). Stella Maris® is derived 114 from the marine algae A. nodosum, and harvested from the nutrient-laden waters of the 115 North Atlantic off the Eastern Coast of Canada. We tested the effect of ANE amendment 116 on two agricultural plants commonly grown in greenhouse conditions (tomato and pep-117

per). Several traits related to plant productivity were measured and soil and root bacterial and fungal diversity were quantified using High Throughput Illumina (San Diego, CA,

<sup>120</sup> USA) Miseq sequencing.

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## MATERIAL AND METHOD

Experimental design

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Greenhouse trials were set up in large trays (60x30x18 cm LxWxH) using two different 125 crops: tomato (Solanum lycopersicum L.) and pepper (Capsicum annuum L.). Tomato culti-126 var Totem Hybrid#A371 was planted in November 16th 2015 and pepper cultivar Ace Hy-127 brid#318 was planted in December 9th 2015. Tomato and pepper seeds were purchased 128 from William Dam Seeds Ltd (ON, Canada). These cultivars were selected for green-129 house production. Soil was collected from an agricultural field under organic regime at 130 the IRDA research station in St-Bruno (Qc, Canada, 45°32′59.6"N, 73°21′08.0"W) on Oc-13: tober 7th 2015. The soil was a loamy sand and was collected from the 15 cm top layer. Natural soil was homogenized and put into trays, filled to 15 cm in height. Soil analy-133 sis was done using a commercial service provided by EnvironeX (formerly AgriDirect, Longueuil, QC) and soil characteristics are shown in Table S1. Eight seeds per tray were planted and after germination, only four seedlings per tray were kept.

For each plant species, a randomized split block design (Table S2) was used with four trays set up per block and eight blocks for each trial. Half of the trays were amended with ANE, as described below. Half of the trays were also planted (planting effect) with four plants per tray, while the other trays were not planted. This allowed a direct comparison of fungal and bacteria soil communities with respect to the amendment and planting

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Two different amendment regimes were used according to the plant species. For tomatoes, plants were amended using multipurpose organic fertilizer (pure hen manure, 18 g per tray repeated every 4 weeks, 5-3-2) from Acti-sol (Notre-Dame-du-Bon-Conseil, QC) in addition to Stella Maris® (3.5 ml per 1L, each tray received 250 ml, repeated every

2 weeks) for the duration of the experiment. The other half were not treated, but watered with 250ml per tray instead. The physico-chemical composition of Stella Maris® is shown in Table S3. For the pepper experiment, the amendment treatment consisted solely 151 of Stella Maris® (3.5 ml per 1L, each tray received 250 ml, repeated every 2 weeks) for 152 the duration of the experiment. The other half was not amended, but watered with 250 153 ml per tray instead. Both experiments were managed under organic farming practices. 154 Thrips were controlled using Neoseiulus cucumeris (syn. Amblyseius cucumeris) (1 bag per 155 plant), Fungus gnats were also controlled using predatory mite Gaeolaelaps gillespiei (1L; 156 Natural Insect Control, ON). Plants were treated once a week with Milstop, a Potassium 157 Bicarbonate-based foliar fungicide to control the powdery mildew on both crops. 158

160 Plant productivity

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Tomato and pepper experiments were harvested on March 29th 2016. The following traits assessed plant productivity: fruit number, fruit weight, shoots fresh weight and roots fresh weight. Together, these traits are expected to represent well the plant overall productivity. Traits were measured on three plants chosen randomly per tray for each amended / control plant, crop (tomato / pepper) and block (eight blocks) for a total of 96 samples. In addition, both shoot and root samples were dried in a 70 degrees drying oven, and dry weights were quantified after 48 hours.

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Sample preparation, DNA extraction and High throughput sequencing

Soil and root samples were taken for both experiments. Soil DNA was extracted using NucleoSpin® Soil DNA extraction kit (Macherey-Nagel, BioLinx, ON) on 250 mg of soil, following the manufacturer's protocol. Roots were first washed with tap water and rinsed with sterile water. Chopped roots sub-samples (100 mg) were subjected to DNA extraction using DNeasy Plant Mini kit (Qiagen Inc - Canada, ON), following the manufacturer's recommendations. Amplicon sequencing targeting bacterial 16S rRNA gene and

fungal ITS was performed on both root and soil samples. For fungal ITS, we used the following primers with the universal CS1 and CS2 adapters: CS1\_ITS3\_KYO2 (5'-ACA CTG ACG ACA TGG TTC TAC AGA TGA AGA ACG YAG 179 YRA A-3') and CS2\_ITS4\_KYO3 (5'-TAC GGT AGC AGA GAC TTG GTC TCT BTT VCC 180 KCT TCA CTC G-3') to produce a final amplicon size of approximately 430bp including 181 adapters (Toju et al., 2012). 182 183 For bacterial 16S, we used the following primers with CS1 and CS2 universal adapters: 184 341F (5'-CCT ACG GGN GGC WGC AG-3') and 805R (5'-GAC TAC CAG GGT ATC TAA 185 TC-3') to produce a final amplicon size of approximately 460 bp and targeting specifically 186 the bacterial V3-V4 region of the 16S ribosomal gene (Klindworth et al., 2013). 187 188 DNA samples were then barcoded, pooled and sequenced (2X300bp, paired-end) using 189 an Illumina MiSeq sequencer through a commercial service provided by the Genome 190 Quebec Innovation Centre (Montreal, QC). Sequences were demultiplexed by the sequencing facility and further processed as described below. **Bioinformatics** All bioinformatics, statistical, and graphical analyses further described were performed in R 3.5.1 (R Core Team, 2018) and detailed scripts are available here (https://github. 196 com/seb951/Acadian\_Seaplants). 197 198

We used the R package DADA2 (Callahan *et al.*, 2016) to infer *Amplicon Sequence Variants*(ASV). DADA2 offers accurate sample inference from amplicon data with single-nucleotide
resolution in an open source environment. Unlike the Operational Taxonomic Unit (OTU)
approach (e.g. Schloss *et al.*, 2009; Caporaso *et al.*, 2010), ASV are not treated as cluster of

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sequences defined with an *ad hoc* sequence similarity threshold. Instead, after sequences are quality trimmed and error-corrected, DADA2 reveals the unique members of the sequenced community, thus allowing sequences and abundance counts to be comparable among studies (Callahan *et al.*, 2016).

First, sequences were trimmed following strict quality thresholds (removing primers and 208 low quality nucleotides, see parameter details in the accompanying R scripts). Follow-209 ing this, we applied the error model algorithm of DADA2, which incorporates quality in-210 formation after filtering, unlike other OTU based methods. Then dereplication, sample 211 inference, merging of paired end reads and removal of chimera were performed in or-212 der to obtain a sequence (ASV) table of abundance per sample. Taxonomy was assigned 213 through the DADA2 pipeline using the Ribosomal Database Project (RDP) Naive Bayesian 214 Classifier algorithm from Wang et al. (2007). Depending on support (minimum boot-215 strap support of 80), we assigned taxonomy from Kingdom to species. We used the 216 silva database formatted for DADA2 to infer bacterial taxa (Callahan, 2018). We used the 217 Unite (Community, 2018) fasta release (including singletons) to infer fungal taxa after formatting it to the DADA2 format using a custom R script. The pipeline was run on a multithreaded (48 CPUs) computer infrastructure provided by Westgrid (https://www. 220 westgrid.ca/support/systems/cedar) and Compute Canada (www.computecanada.ca). 221 Note that the pipeline was run separately for fungal-root, fungal-soil, bacteria-soil and bacteria-root samples given that these were sequenced separately and therefore a specific 223 error model for each dataset was calculated. 224

226 Statistical analyses - plant productivity

Each plant species (tomato and pepper) were analyzed separately. We tested for the amendment effect (tomato: hen manure + ANE, pepper: ANE) on six plant productivity measures (fruit number, average fruit weight, shoots fresh weight, roots fresh weight,

shoots dry weight, roots dry weight). We used Linear Mixed effect Models (LMM) in
the R package NLME (Pinheiro *et al.*, 2017), which are more appropriate than an Analysis
of Variance (ANOVA) given the current block design (blocks and replicates were treated
as random variables). All six plant productivity measures were either square root or log
transformed in order to help satisfy the assumption of normality and homogeneity of the
variance of the residuals in the LMM statistical framework. For the variables *fruit number*and *average fruit weight*, we also verified statistical significance using a permutation-based
2-way ANOVA (Anderson and Legendre, 1999) given that the residuals of the LMM were
not normally distributed. Results were similar according to the 2-way ANOVA.

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240 Statistical analyses - microbial and fungal diversity

For each fungal-root, fungal-soil, bacterial-root and bacterial-soil datasets, we removed samples that showed poor sequencing output and contained few ASV. In addition, for bacterial root and soil datasets, we removed ASV that were taxonomically assigned to *mitochondria* or *chloroplast* given that these were likely sequences from the plants themselves. To remove low quality samples, we first summed the abundance of all ASV for each sample ( $\sum_{i=1}^{n} ASV$ ) and eliminated samples that had fewer that a summed abundance of 1,000. In addition, we removed ASV from our dataset that were present in fewer than 5% of the samples (less than ten individuals in the soil samples or less than five in the root samples). This was done to remove very rare ASV unique to a block or replicate, but not found in the majority of samples.

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We then conducted community-based analyses looking at the amendment effect on ASV abundance in the tomato and pepper experiments separately. To visualize communities and reduce the complexity of the datasets, relative abundance of all taxa was calculated per family using the R package DPLYR (Wickham *et al.*, 2015) and barplots were drawn using GGPLOT2 (Wickham, 2016). ASV alpha (*a*)-diversity was calculated based on all ASV

(excluding rare ASV, see paragraph above) for each sample using the inverse Simpson diversity index in VEGAN (Oksanen *et al.*, 2013). The effect of the amendment and planting for soil communities were assessed using a Linear Mixed effect Model (LMM) model in the R package NLME (Pinheiro *et al.*, 2017), given the unbalanced, replicated block design. Alpha diversity was *log* transformed in order to help satisfy the assumption of normality of the residuals in the LMM statistical framework.

Using the community matrix data of ASV abundance, we performed PERmutational Multivariate ANalysis Of VAriance tests (PERMANOVA; Anderson, 2001) to identify relationships between the communities according to the experimental design. Data were analyzed separately for fungal-root, fungal-soil, bacterial-root and bacterial-soil in tomatoes and peppers. The ASV abundance matrix was Hellinger-transformed and significance was assessed using 10,000 permutations in vegan (Oksanen *et al.*, 2013). Blocks and replicates were factored as strata in the model.

We also performed redundancy analyses (RDAs) using the Hellinger-transformed ASV abundance matrix in vegan (Oksanen et al., 2013) to visually assess the grouping of sam-273 ples, ASV and their association with productivity variables (species scaling based on ASV 274 matrix). Data were analyzed separately for fungal-root, fungal-soil, bacterial-root and bacterial-soil in tomatoes and peppers, giving a total of eight RDAs. Statistical significance of the RDAs were tested using an ANOVA like permutation test (10,000 permuta-277 tions) in VEGAN. Data were constrained based on four productivity measures (fruit number, average fruits weight, shoots fresh weight, roots fresh weight). We excluded the 279 shoots & roots dry weights as constraints to simplify the model. In addition, these were 280 highly collinear with the fresh weight already included as constraints ( $r^2$ =0.98 and 0.76 281 for shoot dry/fresh weights and root dry/fresh weights, respectively). 282

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Finally, we attempted to identify candidate ASV positively associated with productivity. 284 As such, we identified the ten ASV most positively associated with the measures of fruit number, shoots fresh weight and roots fresh weight from each redundancy analysis for a 286 total of 40 fungal and 40 bacterial candidate ASV. We aligned candidate sequences from 287 these candidates ASV using the Bioconductor R package DECIPHER (Wright, 2016) and 288 build pairwise distances matrices using a JC69 substitution models of DNA sequence 289 evolution (equal base frequencies, Jukes and Cantor, 1969) in PHANGORN (Schliep, 2010). 290 Phylogenetic trees (neighbour-joining) for bacteria and fungi were plotted using APE (Par-291 adis et al., 2004). This permitted to identify if similar candidate ASV were found under 292 different experimental conditions (soil/root, pepper/tomato), thus reinforcing their role 293 in productivity increase, and increasing the probability that they are true positives. 294

### 96 RESULTS

297 Effects of the amendment treatment on productivity

The effects of the amendment treatment on tomato (hen manure + ANE) and pepper 298 (ANE) were determined by measuring six agronomic parameters (fruit number, average 299 fruit weight, shoots fresh weight, shoots dry weight, roots fresh weight, roots dry weight). 300 We observed a significant increase of almost all these agronomic parameters (LMM, p-301 value<0.005, Figure 1) for amended plants except for the average fruit fresh weight for 302 tomato that did not differ between amended and control plants (LMM,  $F_{(1,23)} = 1.81$ , p-303 value=0.19, Figure 1 and Figure S1). The amendment effect was stronger in the tomato 304 plants (fold changes between amended and control plants shown in Figure 1), likely due to the fact that these plants were fertilized with both hen manure and ANE. 306

308 Amplicon Sequencing

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A total of 2.7 million paired-end raw reads were obtained for all samples combined (976,000 for fungi-soil, 920,000 for fungi-root, 309,000 for bacteria-soil and 535,000 for bacteria-root, Table S4). On average, 47,664 paired-end reads were obtained per sample.

After quality filters were applied, including removing chimeras, and paired-end reads were merged, an average of 19,690 sequences remained per sample. From 192 soil samples for fungi and bacteria, and 96 root samples for fungi and bacteria, three fungi-soil samples, 15 fungi-root samples and one bacteria-root samples were removed because they had to few reads based on our strict quality thresholds.

The DADA2 pipeline inferred 6,112 fungal-soil, 845 fungal-root, 9,352 bacterial-soil and 2,023 bacterial-roots ASV (Table S4). In bacteria-soil, we further removed a total of 79
ASV whose taxonomy corresponded to *mitochondria* or *chloroplast* and represented 0.1% of all sequencing reads. In bacteria-root samples, we removed a total of 284 ASV that

corresponded to *mitochondria* or *chloroplast* and represented 89% of all sequencing reads.

After filtering out rare ASV, we retained 413, 106, 807 and 262 ASV respectively for fungal
soil, fungal-root, bacterial-soil and bacterial-roots. These retained ASV comprised 94%,

95%, 89% and 11% of all filtered-merged sequences assigned to ASV by the DADA2 pipeline

in the fungal-soil, fungal-root, bacterial-soil and bacterial-root samples, respectively.

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Fungal and bacterial diversity in root and soil biotopes

The microbial community structures of soil and root samples were analyzed and the relative abundance of their taxa was determined at the family level (Figures 2 & 3). Nectriaceae dominated the fungal communities, both in the root and soil samples, while the
bacterial family Bacilaceae dominated to a lesser extent the soil samples. Bacterial root
communities harbored a number of different families: Streptomycetaceae, Sphingomonadaceae, Rhizobiaceae and Pseudomonadaceae among others.

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336 Local (a-diversity)

The *a*-diversity was calculated separately for each sample, under each experimental condition (fungi-soil, fungi-root, bacteria-soil and bacteria-root for both tomato and pepper, Figure 4). Linear mixed effects models showed that the *a*-diversity (Inverse Simpson Index) was significantly higher in the soil biotope that in the roots for both fungi (mean *a*-diversity soil-fungi = 2.88 vs. mean *a*-diversity root-fungi = 27.3,  $F_{(1,239)}$ =899.5, *p*-value<0.0001) and bacteria (mean *a*-diversity soil-bacteria = 4.7 vs. mean *a*-diversity root-bacteria = 69.2,  $F_{(1,223)}$ =1198.1, *p*-value<0.0001).

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In soil samples, fungal a-diversity was not significantly different in amended versus control plants for neither tomato ( $F_{(1,66)}$ =1.6, p-value=0.21) nor pepper ( $F_{(1,69)}$ =1.2, p-value=0.05). In root samples, fungal a-diversity was significantly different in amended versus control plants for tomato ( $F_{(1,21)}$ =10.2, p-value=0.004), but not pepper ( $F_{(1,56)}$ =3.1,

*p*-value=0.10). 350 In soil samples, bacterial *a*-diversity was significantly different in amended versus control 351 plants for pepper ( $F_{(1.69)}$ =31.5, p-value<0.0001), but not tomato ( $F_{(1.69)}$ =1.9, p-value=0.17). 352 In root samples, bacterial a-diversity was significantly different in amended versus con-353 trol plants for tomato ( $F_{(1,22)}$ =39.7, p-value<0.0001), but not pepper ( $F_{(1,4)}$ =0.17, p-value=0.70). 354 355 Differences in species composition among sites 356 Using a PERMANOVA, we identified that the ANE amendment treatment had a highly 357 significant effect on both fungal and bacterial community structures (Table 1). This effect 358 was stronger in the root (9-30% of variance explained in the models) than in the soil (3-6%) 359 of variance explained in the models). Planting also had a significant effect on fungal and 360 bacterial community structures (12-24% of variance explained in the models). 361 362 Redundancy analyses (RDAs, Figures 5 for fungi and Figure 6 for bacteria) illustrated that 363 roots fresh weight, shoots fresh weight and fruit number responded similarly, while average fruit weight behaved differentially as noted previously (in fact nearly orthogonally to the other three parameters in most ordinations). In addition, it showed that fertilized samples clustered together and were positively correlated with increases in these four productivity measures. All RDA model tested were significant ( $F_{(4,10)}>1.4$ , p-value<0.03 368 for all models). 369 370 Next, we identified, for each RDA, the ten ASV most closely related to the three con-371 straints of the model (roots fresh weight, shoots fresh weight and fruit number). These 372 ASV were considered as putative candidate taxa most positively impacted by increases 373

in productivity due to the ANE amendment treatment. We further analyzed the corre-

sponding sequences for these eighty candidate ASV (ten candidates \* eight ordinations)

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in two separate alignments (one for fungi and one for bacterial ASV) and their accompanying phylogenetic trees.

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In fungi, we identified a number of different ASVs positively associated with productiv-379 ity (Figure S2). Notably, five different ASVs taxonomically assigned to the family Mi-380 croascaceae (phylum Ascomycota) in all conditions except the pepper-root were posi-381 tively associated to productivity. In addition, two ASV assigned to Mortierella spp (soil 382 saprotrophs in the phylum Mucoromycota), and a cluster of six different fungal closely 383 related ASV in tomato-soil (ASV67 & ASV132), tomato-root (ASV10, ASV1017, ASV1018, 384 ASV1019) and pepper-soil (ASV67) were positively associated to productivity in both 385 tomato and pepper roots. Given that no taxonomy was assigned to these sequences 386 through the DADA2 RDP bootstrap approach, we used a BLASTn (Altschul et al., 1997) ap-387 proach to identify the most closely related sequences against NCBI nr. The most closely 388 related fungal reference sequences were from an uncultured fungus clone (BLASTn, 86% 389 identity, e-value=9e-58, sequence ID: EU517002.1). Similarly, two unknown ASV (ASV61 390 & ASV81) also matched an uncultured fungus clone (BLASTn, 94% identity, e-value=4e-165, sequence ID: DQ900965.1). Finally, another cluster of ASVs in the pepper-root was assigned to Olpidium brassicae, a putative fungal parasite belonging to flagellate fungi (Lay 393 et al., 2018).

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In bacteria-roots, we identified a large diversity of ASV positively correlated (increased abundance of these ASV) with productivity (Figure S3), Among others we identified *Rhi-zobium*, *Sphingomonas*, *Sphingobium*, *Bradyrhizobium* in both the soil and root biotopes and tomato and pepper species.

### 100 DISCUSSION

In the current study, we investigated the effects of *Ascophyllum nodosum* extracts (ANE)
on root, shoot and fruit biomass in addition to bacterial and fungal communities in tomato
and pepper. Overall parameters related to plant growth (root, shoot and fruit weights)
significantly increased in both plant species in response to amendment treatment. These
results corroborate previous studies documenting the impact of ANE on productivity in
strawberries (Alam *et al.*, 2013) and carrots (Alam *et al.*, 2014).

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In the tomato experimental set up, the effect of the amendment treatment was especially 408 high, likely due to the fact that plants were also fertilized with hen manure in addition to ANE (Figure 1). This was not the case for the pepper plants and the increase in pro-410 ductivity was solely due to the addition of ANE. The commercial extract used in this investigation contained about 0.1% nitrogen, 0.2% phosphorus, 5% potassium, along with several micronutrients (Table S3). In the current experimental setup, ANE was diluted to 3.5 ml/L prior to application (250 ml per tray every two weeks). In fact, in the tomato plants the amounts of N, P and K supplied via the application of ANE were 200-1000 415 times less than from the hen manure itself. As such, these nutrients were given at very 416 low concentrations relative to the crop requirements and are not expected to significantly 417 impact growth relative to a regular agricultural fertility program (Bruulsema et al., 2012; 418 Alam et al., 2013). Instead, organic molecules such as betaines, polyamines, cytokinins, 419 auxins, oligosaccharides, amino acids and vitamins present in ANE have been found to 420 have overall beneficial productivity effects on plant growth (Khan et al., 2009; Craigie, 421 2010, 2011; Battacharyya *et al.*, 2015). 422

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One of primary goal of the study was to document how bacterial and fungal communities responded to the addition of ANE. We used Illumina MiSeq amplicon sequencing targeting DNA regions specific to fungi (ITS) and bacteria (16S). Then, we identified bacterial and fungal taxa present in the samples using a relatively novel bioinformatics approach developed by Callahan *et al.* (2016). The approach, based on the widely used programming language R (R Core Team, 2018), identifies unique, non-clustered sequences (ASV) that are then comparable among studies. In addition, the current analytical pipeline uses a bayesian classifier for taxonomy rather than the widely used BLAST approach, thus providing more conservative, but more accurate taxonomic identifications (Wang *et al.*, 2007).

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In the current experimental set up, most ASV identified were rare and unique to one or a 435 few sample. In fact, approximately 90% of all ASV were discarded given that they were 436 found in singletons or present in very few samples and were thus not representative of 437 a particular experimental treatment. These 'rare' ASV comprised a small minority of all 438 sequencing reads (approximately 5% of all sequences), a pattern reminiscent of the early 439 species abundance models showing that in most ecological communities, few species are 440 exceptionally abundant whereas most are rare (Fisher et al., 1943). In addition, a large fraction of the sequencing reads in the root bacterial communities likely originated from the plants themselves (identified as chloroplast or mitochondria). This may be partly explained by the fact that most of root biomass collected was from large roots (Fig. S1B), rather than fine root hair where most microbial biological activity likely takes place (Pre-445 gitzer et al., 2002) and therefore the total biomass extracted in the current study likely 446 consisted of more root cell rather than biologically active bacterial biomass. 447

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Nectriaceae, a family of fungi in the order Hypocreales and often encountered as saprotrophs on decaying organic matter comprised most of the diversity both in the soil and plant roots (between 25-70% of the total number of sequencing reads, Figure 2). With respect to bacterial communities of the soil, these were much more diverse and comprised

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many different families (Figure 3). The amendment treatment had a significant effect on both fungal and bacterial *a*-diversity (total richness of ASV) in the root biotope, except for bacteria in pepper plants. In the soil biotope, it only had a significant effect for bacteria in the tomato plants (Figure 4).

The amendment treatment significantly influenced fungal and bacterial community com-458 position (b-diversity) among root and soil biotopes. This effect was relatively small (3-33%) 459 of variance explained in the models, Table 1) but significant, implying that the addition of 460 ANE (pepper) or ANE and hen manure (tomato) is, at least partly, responsible for shaping 461 microbial communities. In addition, a significant proportion of the variance in soil com-462 munities (12-24%) was explained by the planting effect, showing how plants can alter 463 their microbiome. Finally, we also tested the effect of plant species identity on commu-464 nity structure on a combined dataset comprised of both the tomato and pepper plants. In 465 the root biotope, we find that this effect (26 and 20% of variance explained in the mod-466 els for fungal and bacterial communities, respectively, Table S5) is in line with numerous studies reporting how plants select their microbial communities (Chaparro et al., 2014; Reinhold-Hurek et al., 2015). Nevertheless, we recognize that the current experimental setup precludes any strong conclusion regarding the plant species' effect of community structure, as it does not allow to explicitly disentangle the species effect from the "addition of hen manure" effect.

Finally, we also aimed to identify candidate taxa positively correlated with increased plant productivity in response to ANE application. In fungi, one cluster of ASV taxonomically assigned to *Mortierella* (soil saprotrophs in the phylum Mucoromycota) was positively correlated to productivity in both tomato and pepper roots. Interestingly, Li et al. (2018) found that a closely related species (*M. elongata*) can improve soil health and stimulate production of plant growth hormones. In their study, Chung et al. (2007)

506

showed how increases in productivity led to greater microbial biomass and greater number of saprophytic and arbuscular mycorrhizal fungi. Perhaps, this is explained by the fact that an increase in plant productivity can lead to greater substrate availability, potentially increasing the activity of saprophytic fungi feeding on this organic matter substrate.

Several other fungal ASV in tomato (soil) and pepper (root) were linked to increases in productivity. Surprisingly, a putative plant pathogenic fungus (*Olpidium brassicaceae*, Figure S2) was positively associated with increased productivity. However, *O. brassicae* has been correlated with a decrease plant growth, but only when present in large amount (Lay *et al.*, 2018). In addition, *O. brassicae* is likely a species complex and has been shown to constitute a large proportion of the plant roots or rhizosphere fungal community in many different systems, particularly in Brassicaceae crops (Lay *et al.*, 2018).

In bacteria roots samples, a diverse number of ASV were positively impacted by the amendment treatment (Figure S3) and many of those are known to be present in the root endosphere (e.g. *Rhizobium*, *Sphingomonas*, *Sphingobium*, *Bradyrhizobium* spp, Tkacz and Poole, 2015). For example *Rhizobium*, and *Bradyrhizobium* spp. have been shown to promote plant growth, P solubilization, N fixation and overall productivity in both legume and non-legumes species such as radishes (Antoun *et al.*, 1998; Avis *et al.*, 2008).

It is now well established that seaweed extracts have a positive effect on agricultural plant productivity. Concurrently, DNA barcoding permits a more comprehensive understanding of the diversity and ecology of microbial organisms and how they interact. In fact, plants and microbes should likely be redefined as *holobionts*, an assemblage of different species that forms an ecological unit (Margulis and Fester, 1991). In this study, we showed that the addition of ANE increased plant productivity. It also increased, by a small, but significant margin, the fungal and bacterial (only in the rhizosphere) biodiversity and

changed the microbial community structure in the roots and in the rhizosphere of both

tomato and pepper plants. Finally, we identified bacterial and fungal taxa, especially saprotroph, that were positivity associated with plant productivity. Further studies, for example using inoculum of the candidate microbial species linked to increases in productivity that we identified, may help to identify a causative link between liquid seaweed extracts, microbes and productivity.

512

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Table 1: Variance explained by the terms in the PER-MANOVA models

	amendment	planting	amendment:planting
fungi-soil (tomato)	0.05***	0.24***	0.02**
fungi-root (tomato)	0.29***	NA	NA
bacteria-soil (tomato)	0.06***	0.17***	0.04**
bacteria-root (tomato)	0.33***	NA	NA
fungi-soil (pepper)	0.03**	0.2***	0.02*
fungi-root (pepper)	0.1***	NA	NA
bacteria-soil (pepper)	0.06***	0.12***	0.02*
bacteria-root (pepper)	0.19	NA	NA

 $r^2$  (percentage of variance explained by the term in the model); \*p-value<0.05, \*\*<0.005, \*\*\*<0.0005

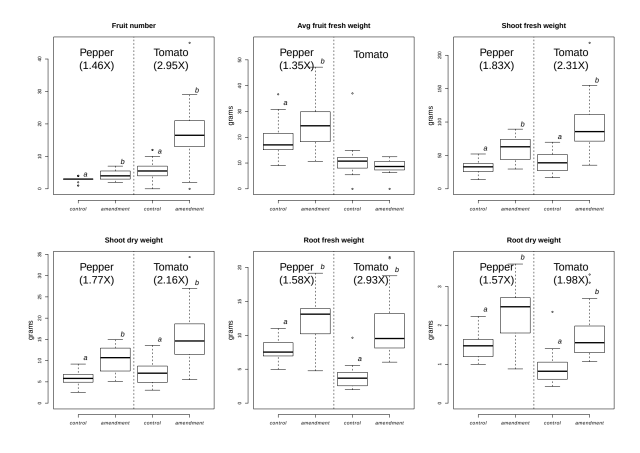


Figure 1: Measures of plant productivity. a and b subscripts above boxplots denote significant differences (p-value < 0.005) according to the amendment effect (tomato: hen manure + ANE, pepper: ANE). Fold changes between the mean of the control and amended plants were also noted for significant differences (for pepper and tomato separately).

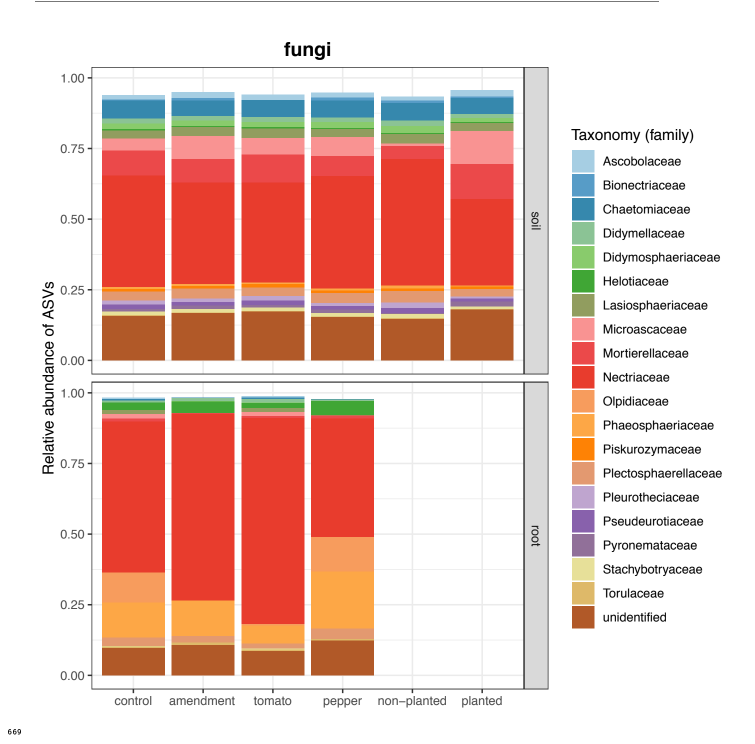


Figure 2: Barplots of the relative abundance of fungal ASV for fungi

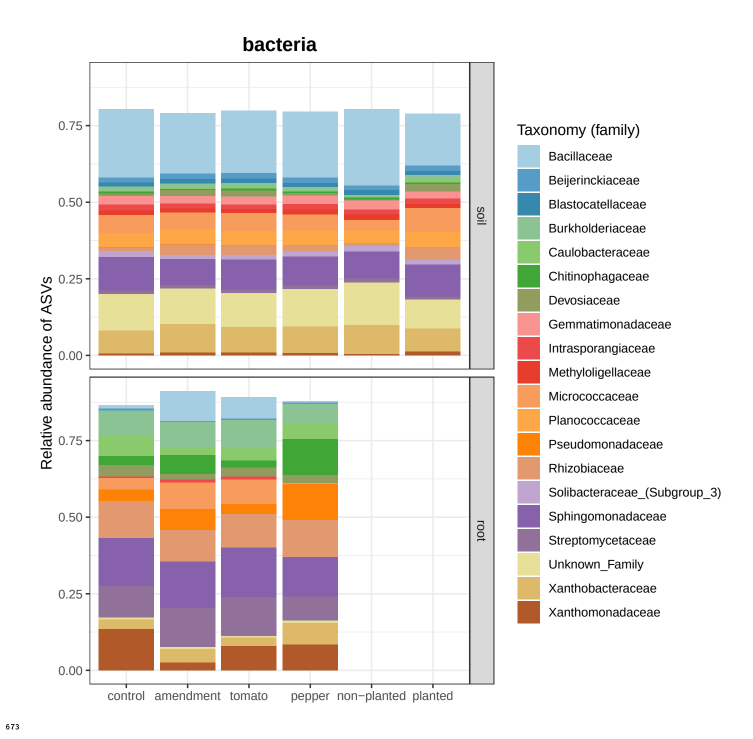


Figure 3: Barplots of the relative abundance of bacterial ASV for bacteria

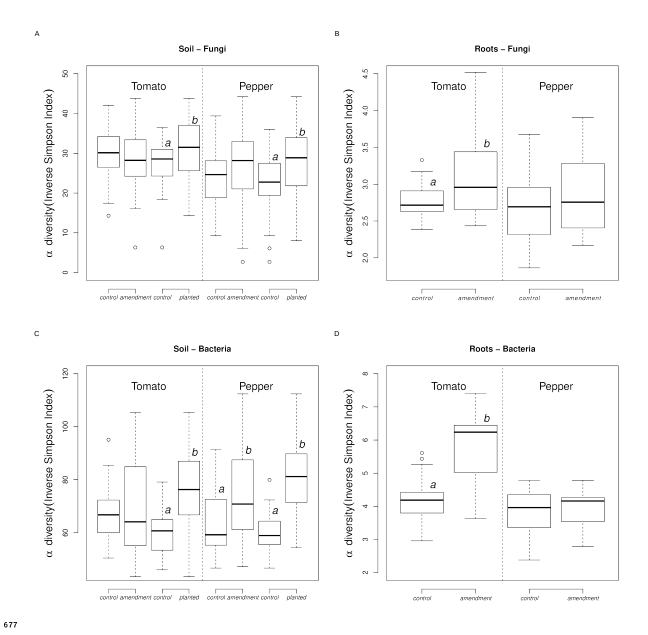


Figure 4: Boxplot of a-diversity according to the amendment and planting effect for fungal-root, fungal-soil, bacteria-soil and bacteria-root for tomato and pepper. a and b subscripts above boxplots denote significant differences (p-value < 0.05).

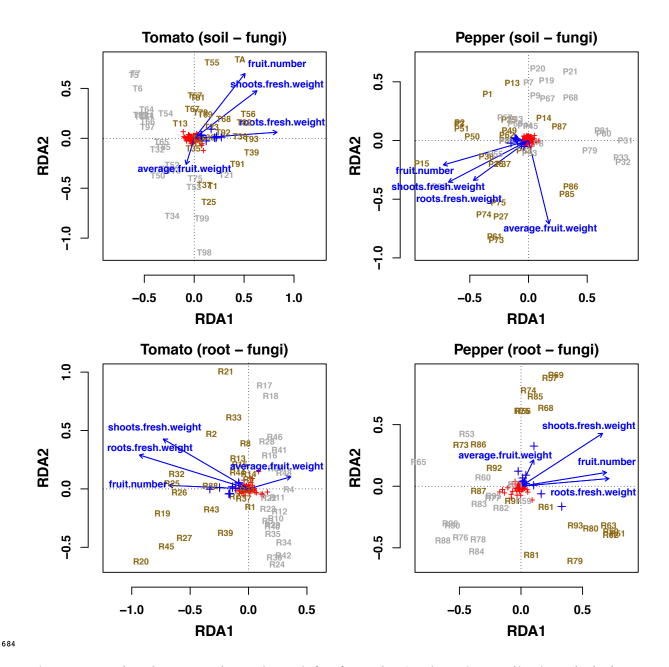


Figure 5: Redundancy analyses (RDA) for fungal ASV (species scaling). Labeled samples were colored in gray (unfertilized) or dark yellow (fertilized). Red + signs represent individual ASV, while blue + signs are the ten ASV most closely associated with the three productivity measures of root fresh weight, shoots fresh weight and fruit number. Blue arrows are the four productivity measures used as constraints in the ordinations.

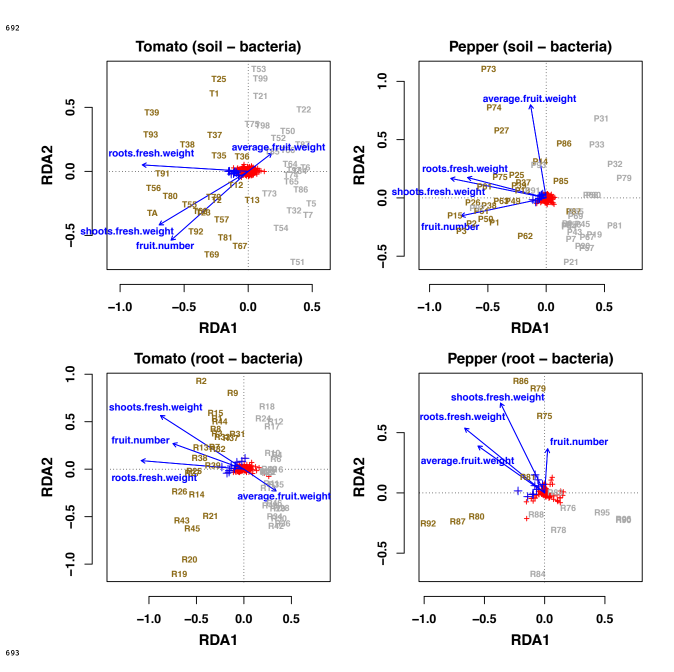


Figure 6: Redundancy analyses (RDA) for bacterial ASV (species scaling). Labeled samples were colored in gray (unfertilized) or dark yellow (fertilized). Red + signs represent individual ASV, while blue + signs represent the ten ASV most closely associated with the three productivity measures of root fresh weight, shoots fresh weight and fruit number. Blue arrows are the four productivity measures used as constraints in the ordinations.