INTRODUCTION
The Institute of Agricultural and Fisheries Research (ILVO) has established a general Genomics platform, joining forces in the genomics research of the four ILVO Units. One part of the project focuses on the study of microbial communities of four different environments: soil, seawater, rumen (cows) and the intestine (pigs). This poster presents the research of two of these environmental niches: the rhizosphere of the plant (a soil environment) and (micro)plastics (a marine environment).

Soil environment – Rhizosphere

The rhizosphere of the plant is defined as the narrow zone around the roots and can contain more than 10^11 microbial species per gram of soil. The effect of different soil treatments on the rhizosphere microbiome of lettuce and strawberry, and the influence on plant health are investigated. Initially, biochar, the solid coproduct of biomass pyrolysis (a technique used for biofuel production), is used as a soil additive.

Approach
The microbial rhizosphere of lettuce and strawberry

Amplicon sequencing
16S rDNA (V3-V4 region)

Metatranscriptomics

Composition and dynamics of the microbial community

Preliminary Results
Initially, the 16S rDNA V3 region was sequenced of three lettuce rhizosphere samples: potting soil (a), field soil (b) and field soil with 1% biochar (c) (HiSeq 2500, 91bp paired end sequencing).

As expected, the bacterial taxa of potting soil and field soil showed a major difference. Potting soil was dominated by Acidobacteria, whereas field soil contained especially Actinobacteria. In both soil types, Proteobacteria were dominantly present.

The addition of biochar seemed to induce a microbial shift. This result confirms previous observations of DGGE and PLFA analysis.

Future experiments
Amplicon sequencing of the V3-V4 region of the 16S rDNA will be used primarily to investigate the influence of biochar and other soil treatments on the rhizosphere microbiome of lettuce and strawberry. The activity of the rhizosphere microbiome will be studied using metatranscriptomics.

Marine environment – Microplastics

Plastic particles with a diameter less than 5 mm are defined as microplastics. They are produced by the industry or are derived of larger plastics due to the influence of UV light and mechanical/physical factors. These microplastics are colonized by various bacteria. The microbial load of these microplastics is investigated in function of biodegradation.

Approach
The microbial load of microplastics

Amplicon sequencing
16S rDNA (V3-V4 region)

Whole genome shotgun sequencing

Analysis of the genes expressed in the microbial community

Preliminary Results
Initially, a pilot study was performed. The 16S rDNA V3 region was sequenced of two samples: a beach pellet and a blue synthetic rope of the marine environment (HiSeq 2500, 91bp paired end sequencing).

The bacterial community of both kind of plastics was different. The beach pellet mainly showed a colonization of Psychrobacter, whereas the Pseudoalteromonadaceae dominated the synthetic rope.

These results confirmed previous tests with DGGE analysis, cultivation of the bacteria and cloning.

Future experiments
The bacterial species present on the microplastics will be identified using V3-V4 16S rDNA amplicon sequencing. To identify plastic degrading bacteria, the functions present in the microbial environment will be studied by using whole genome shotgun sequencing.

CONCLUSION
Metagenomic research of microplastics and the rhizosphere of lettuce gives information about the microbial composition and functions of the species present in their environment. Knowledge about the microbial load on microplastics and the functions these bacteria perform could help in the detection of marine plastic-degrading bacteria.

Understanding the influence of biochar on the rhizosphere microbiome and the effect on soil-borne pathogens, could lead to the development of novel ways to restore soil quality and enhance plant health.