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# Impact of CeO<sub>2</sub> nanoparticles on the microbiota of the *S. flos-cuculi* L. (Caryophyllaceae) rhizosphere†

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The aim of this study was to correlate the influence of cerium oxide nanoparticles (nCeO<sub>2</sub>) resulting from pollution sources, on the root bacterial composition and the associated substrate (root zone soil) of *Silene flos-cuculi* (L.) using a metabarcoding technique. Currently, limited information is available regarding the direct effects of nCeO<sub>2</sub> on plants and the rhizosphere microbiota, where changes in turn could positively or negatively influence plant performance. To assess the distribution of the main bacterial phyla in the culture substrates, analyses were conducted considering both intracellular DNA (iDNA) contained within intact and live bacterial cells, and extracellular DNA (eDNA) from lysed cells. The impact of various nCeO<sub>2</sub> dosages on phyla, families, and genera was then investigated with a detailed examination of all detected members at the family and genus levels to differentiate the nCeO<sub>2</sub> treatment effects. The results revealed that 25 out of 641 identified bacterial genera, primarily anaerobic and strictly anaerobic, exhibited reduced presence in nCeO<sub>2</sub> treated samples compared to the controls. This decrease was particularly evident in species belonging to the phylum *Firmicutes*. Metabolic function analysis performed using FAPROTAX indicated a decline in fermentative, nitrogen fixation, and iron/nitrate respiration metabolisms in nCeO<sub>2</sub> treated samples, especially at higher concentrations. Conversely, there was an increase in chemoheterotrophy and aerobic chemoheterotrophic-related functions in these samples.

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## Environmental significance

Nanoparticles are even more present in the environment, for many reasons. However, little is known about their impact on the different environmental microbiota and also in relation to humans, animals, and plants. For this reason, the present work aims to evaluate the effect of nCeO<sub>2</sub> particles on the *Silene flos-cuculi* (L.) root bacterial composition and its associated substrate (root zone soil).

## 1 Introduction

Thanks to the advent of nanotechnology, new nanomaterials useful in numerous applications have been engineered.<sup>1</sup> However, the diffusion of these new materials in the environment must be studied to evaluate their possible impact on ecosystems.<sup>2</sup> In this regard, cerium (Ce)

nanoparticles are one of the most important, already widespread into the environment because of their use as an anti-corrosion component in aluminium alloys to replace hexavalent chromium as well as an additive in painting systems.<sup>3,4</sup> Ce has been also studied for decades as a catalyst and material for electrolytes and electrodes in fuel-cell systems, and nanostructured cerium oxide (NS-CeO<sub>2</sub>) materials have been developed to meet the high energy and environmental demands because of their superior redox and catalytic properties compared to bulk materials.<sup>5,6</sup> Ce nanoparticles (CeNPs) in water suspensions also found application as polishing supports, anticorrosive systems, oxidants for electrode sensors, and redox agents for medical treatment. In the last case, although Ce itself has no biological role in mammalian physiology, Ce<sup>3+</sup> soluble salts (nitrate, acetate, and chloride) have traditionally been used for biomedical purposes due to their antiemetic,

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bacteriostatic and bactericidal properties.<sup>7</sup> CeNPs have been also reported to beneficially modulate excess reactive oxygen species (ROS) in cells; however, their toxicological effects associated with their structure and stability should be considered.<sup>8</sup>

The effects of CeNPs in complex environments are influenced by several variables, including the manufacturing method, nanocrystal shape and size,<sup>9,10</sup> and redox potential.<sup>8,11,12</sup> Biological studies involving CeNPs revealed significant variability and conflicting results.<sup>13,14</sup> *In vivo* studies have shown that CeNPs are bio-persistent for long periods after inhalation<sup>15,16</sup> and oral administration,<sup>17</sup> and can be internalized by macrophages and epithelial cells.<sup>18</sup> Furthermore, given the irreversibility of their accumulation as well as the potential long-term effects and interactions of Ce oxide nanoparticles (nCeO<sub>2</sub>) with abiotic and biotic entities, it is critical to conduct research on the consequences of the large-scale use of these materials.<sup>13</sup>

For this reason, given their already present diffusion and their high persistence in soil, one of the most important aspects to evaluate are the effects of these nanoparticles on soil structure and plants, especially those intended for human and animal consumption, also in relation to the microbial population that lives symbiotically on the roots.<sup>19,20</sup> Synchrotron spectroscopic techniques have revealed that plants can improve CeNP reduction in soil and hydroponic systems, and Ce absorption and translocation differed among rice, wheat, and barley grown in soils enriched with CeNPs. However, while Ce does not accumulate in the grains of wheat, it accumulates in the tissues and grains of rice and barley.<sup>21</sup>

There are conflicting reports regarding the growth and health of plants exposed to this nanomaterial. Some studies report an improvement in factors, such as growth and increases in biomass, others testify to the induction of physiological and morphological anomalies.<sup>22–28</sup>

Some evidence suggests a possible toxic effect of this element on the microbial flora of the soil<sup>29</sup> and Ce microbial accumulation in microorganisms isolated from rare earth-rich environments.<sup>30,31</sup> For this reason, given the fundamental role of bacteria in the health and growth of plants,<sup>32,33</sup> it is important to expand knowledge on the possible effects of an accumulation of this compound in relation to the effects induced on the soil microbiota. In fact, it is possible to hypothesize that some of the effects previously observed on plants could be correlated to modifications of the symbiotic microflora, which, given the high importance and interconnection, would be reflected in the physiology of the plant.

In this regard, starting from a previous study, which reported that nCeO<sub>2</sub> is absorbed by plant roots and translocated toward aerial plant fractions, inducing differences in the development of the treated plants,<sup>34</sup> this work focused on the modification induced by different concentrations of nCeO<sub>2</sub> soil amendments on substrates on the rhizospheric microbiota of *S. flos-cuculi* L. (Caryophyllaceae). For this purpose, a next-generation sequencing technology (NGS) approach was applied.

## 2 Materials and methods

### 2.1 Experimental design and sampling

The characteristics of nCeO<sub>2</sub>, plant materials and the experimental design were set up as described in a previous study<sup>34</sup> and are summarized in Fig. 1. Briefly, *Silene flos-cuculi* (L.) was grown in an organic substrate (“Compo Sana”, pH = 6.8–7.2), with four replicates for each trial, in growing pots filled with 500 g of substrate. Control samples (CPR) were sown in the substrate without any addition of nCeO<sub>2</sub> and were irrigated with pure tap water. In contrast, the potting substrate of samples 20DR and 40DR was irrigated with water suspensions of nCeO<sub>2</sub> with a dimension of 25 nm to achieve an nCeO<sub>2</sub> substrate concentration of 200 mg kg<sup>-1</sup>. Then, growing pots were seeded with *S. flos-cuculi* (L.) and placed in full sunlight at 18–27 °C (night/day) with a relative humidity of around 60%, irrigated every three days, and seedlings were thinned to two for each pot after two weeks. In trial 20DR, a further nCeO<sub>2</sub> application of 200 mg kg<sup>-1</sup> was performed 20 days after seedling emergence (DSE), whereas for trial 40DR, two applications of 200 mg kg<sup>-1</sup> nCeO<sub>2</sub> were performed after 20 and 40 days from DSE. At 60 days from DSE, all plants were harvested. The nCeO<sub>2</sub> doses were chosen on the basis of previous results,<sup>34</sup> which demonstrated that 200 and 400 mg kg<sup>-1</sup> were found to be the

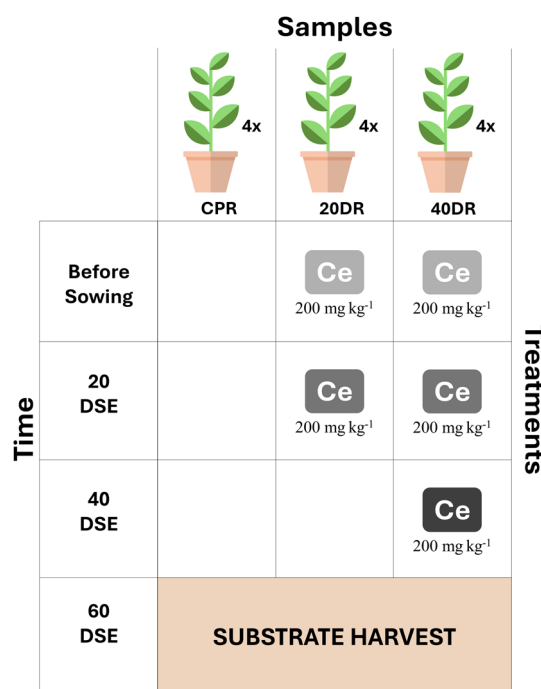


Fig. 1 Experimental setup showing the combination of treatments: control trials (CPR) were grown in the untreated growing substrate, while the substrate of Ce-treated trials was added with 200 mg kg<sup>-1</sup> nCeO<sub>2</sub> before sowing. Trial 20DR was further added with another equal dose of nCeO<sub>2</sub> at 20 days from seedling emergence (DSE), and trial 40DR with the other two nCeO<sub>2</sub> additions (200 mg kg<sup>-1</sup> + 200 mg kg<sup>-1</sup>) at 20 and 40 days from DSE, respectively. At 60 days from DSE, all the samples were harvested.



most effective in affecting plant growth parameters, with a shortage in biomass of roots and stems, and in nCeO<sub>2</sub> translocation towards aerial plant fractions. The purpose of this protocol was to simulate conditions closer to reality, where plants are probabilistically exposed to CeNPs over a more extended period, at relatively lower concentrations, but through repeated pulses.<sup>35</sup>

Before being separated from the stems and leaves, the roots were gently shaken in an empty sterile glass beaker to collect the substrate in four replicates. A sterilized cleaned trowel was used for each sample to collect the substrate for microbial community analyses. All samples were transported to the laboratory on ice, and the substrates for molecular characterization were immediately stored at -80 °C until analysis.

## 2.2 Extraction and purification of intra- and extracellular DNA

For each trial, DNA was directly extracted from equal aliquots (5 g) of substrate obtained from the four replicates performed for each trial using the following protocol:<sup>36</sup> extracellular DNA (eDNA) was extracted by gentle substrate washing with 5 mL of 0.12 M Na<sub>2</sub>HPO<sub>4</sub> (Sigma-Aldrich, Italy) at pH 8 in 50 mL Falcon tubes horizontally shaken for 30 min (80 rpm). Tubes were then centrifuged (4 °C, 30 min, 7500 × g), and the 5 mL supernatant was collected. The same procedure was repeated twice, and the resulting supernatants were pooled to obtain a final volume of 15 mL of unpurified eDNA. Then, the eDNA solution was purified using a commercial extraction kit (DNeasy® PowerMax® Soil kit, Qiagen, USA) following the manufacturer's instructions but avoiding incubation in cell lysis buffer. Intracellular DNA (iDNA) was extracted from the residual substrate pellets resulting from the alkaline washing. The pellet was collected in a 50 mL Falcon tube and processed using the extraction kit according to the manufacturer's instructions, including incubation in cell lysis buffer. At the end of purification, all DNA samples were suspended in 5 mL of 10 mM Tris solution (Sigma-Aldrich, Italy).

To remove undesired RNA residues, eDNA and iDNA solutions were incubated with RNase A (25 µL, 1 ng µL<sup>-1</sup>, Sigma-Aldrich, Italy) and shaken at 37 °C (1 h, 300 rpm); then sodium acetate (3 M, 10% v/v), NaCl (4 M, 30% v/v), and isopropyl alcohol (66% v/v) were added prior to overnight incubation at -20 °C and centrifugation (4 °C, 30 min, 7500×g). After removing the supernatant, the residual pellet was washed twice with 2 mL ethanol (80%). Finally, the eDNA and iDNA pellets were dried in a stove (10–15 min at 37 °C) and resuspended in 1 mL of *DNase free* sterile water. Purified DNA samples were quantified using a Qubit 3.0 (Life Technology, Carlsbad, California, USA), and quality was assessed using a NanoDrop spectrophotometer (Thermo Fisher Scientific, Waltham, Massachusetts, USA). The fragment length distribution was assessed using 0.8% agarose gel electrophoresis.

## 2.3 Microbiota identification by 16S rRNA gene – amplification, sequencing, and data analysis

The 16S rRNA gene sequences were amplified from the extracted DNA, according to a previous reported protocol.<sup>37</sup> Briefly, for the amplification of the V3–V4 region, the primer pair Probio\_Uni (forward primer 5'-CCTACGGGRSGCAGCAG-3') and Probio\_Rev (reverse primer 5'-ATTACCGCGGCTGCT-3') was used, verifying the integrity of amplicons by electrophoresis with an Experion workstation (BioRad, Italy). The amplicons were purified using electrophoretic separation on a 1.5% agarose gel of a Wizard SV Gen PCR Clean-Up System (Promega, Italy), with a subsequent purification step to remove primer dimers made with Agencourt AMPure XP DNA purification beads (Beckman Coulter Genomics GmbH, Germany). The 16S rRNA gene sequencing was performed using the MiSeq Platform (Illumina) at the GenProbio s.r.l. DNA sequencing facility (<http://www.genprobio.com>). The obtained raw reads were processed using the QIIME2 software suite.<sup>38</sup> The complete Probio\_Uni/Probio\_Rev amplicons were reconstructed from the assembled paired-end reads. Sequences with a length between 140 and 400 bp and a mean sequence quality score >20 were retained, whereas sequences with homopolymers >7 bp and mismatched primers were omitted. 16S rRNA amplicon sequence variants (ASVs) were identified with DADA2, which defines variants based on unique sequences,<sup>39</sup> and singletons were removed. All reads were classified to the lowest possible taxonomic rank using QIIME2 (ref. 38 and 40) using SILVA as the database.<sup>41</sup> Statistical analyses were conducted using R (v 4.2.1). Observed ASVs and Good's coverage were calculated using *ggrare* (ranacapa package 0.1.0) and represented as rarefaction curves for species richness. After rarefaction of the samples implemented through the *phyloseq* package,<sup>42</sup> the mean species diversity for each treatment (Alpha-diversity) was assessed using the Shannon index. Principal coordinate analysis (PCoA) was used to analyze the patterns of prokaryotic communities using Bray–Curtis and Jaccard dissimilarity matrixes with the *vegan* R package (v 2.5–6).<sup>32</sup> Likewise, to evaluate the similarities among samples from a qualitative point of view, principal component analysis (PCA) was applied to the bacterial group distributions at the genus granularity level, using the *scikit-learn* Python package (v. 0.23.2). To better understand the behavior of the microbiological groups in the different samples, deviations in their distributions compared to the control samples at the phylum, family, and genus granularity levels were evaluated. Finally, the ranking trends of various bacterial groups in the different samples were studied. To such an extent, other than presenting a series of qualitative results concerning the changes in the bacterial rankings in the different samples, the Spearman correlation was calculated pairwise between the samples to evaluate the degree of concordance (between -1 and 1) of the ranks. The ranking, which sorts the bacterial groups from the most to the least present in a sample, provides an overview of the bacterial group distribution.



**Table 1** BioSample and sequence read archive (SRA) accession numbers

Sample	Biosample	SRA
CPR_iDNA	SAMN31207732	SRR21875482
CPR_eDNA	SAMN31207733	SRR21875481
20DR_iDNA	SAMN31207734	SRR21875480
20DR_eDNA	SAMN31207735	SRR21875479
40DR_iDNA	SAMN31207736	SRR21875478
40DR_eDNA	SAMN31207737	SRR21875477

Given two rankings pertaining to two different samples, their similarity through the Spearman correlation was compared. In short, the Spearman correlation measures the strength and direction of the monotonic relationship between the two rankings. Instead of comparing raw data values, it assesses how well the relationship between the rankings of one sample predict the rankings of the other sample. When both samples have perfectly similar rankings, the Spearman correlation is 1 (thus the two bacterial group distributions are similar); when the rankings are completely opposite, it is  $-1$ ; and when the rankings are unrelated, it is close to 0. Furthermore, the metabolic functions of the microbiota in different samples were predicted using the FAPROTAX database.<sup>43</sup>

## 3 Results

### 3.1 Sequencing results

The total DNA that could be extracted from soil could be distinguished in the iDNA, found within cell membranes, and eDNA, representing DNA located outside cell membranes.<sup>36,44</sup> In this study, iDNA and eDNA were differentiated as considering the whole DNA could have led to bias as considering not only the present microbial community but also paleome,<sup>45</sup> overestimating the actual diversity.<sup>46,47</sup>

The obtained sequences were published in National Centre for Biotechnology Information (NCBI) under the

BioProject PRJNA888088 and are available at the BioSample accession numbers shown in Table 1.

### 3.2 Quality control report of the high-throughput sequencing

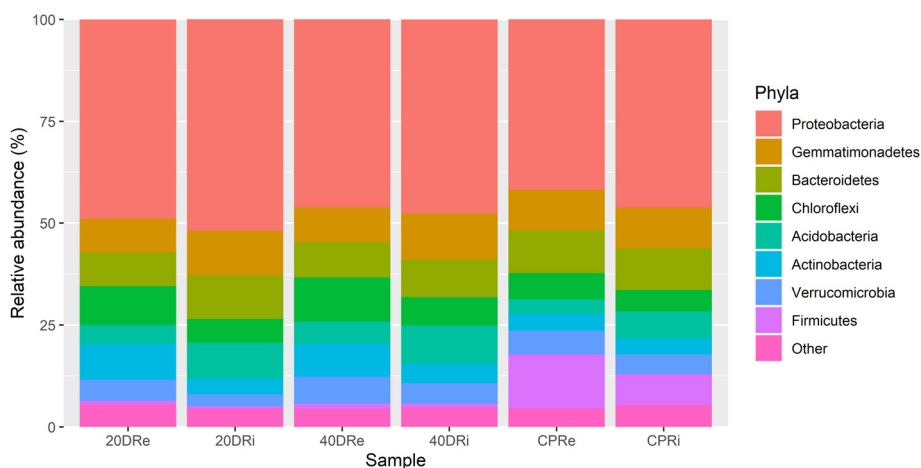
The validity and reproducibility of the sequencing data were verified using rarefaction curves. As shown in Fig. S1,<sup>†</sup> each curve reached plateau, indicating that the number of reads obtained was sufficient to effectively represent the entire bacterial population.

### 3.3 Microbial composition by high-throughput sequencing

Looking at the composition of the microbiota at a phylum level, whose percentages are reported in Table S1,<sup>†</sup> *Proteobacteria* was the phylum present with the highest percentage, with an average relative abundance of 47.11%, with a minimum percentage of 41.83% in the CPRi sample and a maximum percentage of 51.92% in the 20DRi sample, thus constituting alone almost half of the phyla present (Fig. 2). The others more abundant phyla (threshold above 1%) resulted in *Gemmatimonadetes* (9.82%), *Bacteroidetes* (9.60%), *Chloroflexi* (7.50%), *Acidobacteria* (6.46%), *Actinobacteria* (5.59%), *Verrucomicrobia* (5.02%), *Firmicutes* (4.05%) and *Patescibacteria* (1.35%).

### 3.4 Prokaryotic community diversity

Analysing the mean species diversity induced by treatments in the soil ( $\alpha$ -diversity), the Shannon–Wiener  $H$  index highlighted biodiversity differences among the samples. From the values reported in Table 2, albeit all  $H$  values were close to the average value of 5.053, CPR samples showed higher average biodiversity values than the nCeO<sub>2</sub> treated samples (CP = 5119, 20DR = 5023, 40DR = 5035), however, this difference was not statistically significant ( $p > 0.05$ ). Maximum  $H$  values (5228) were observed for CPR\_iDNA, while lower values for Ce-treated samples, with a minimum value of 4.941 for sample 20DR\_iDNA. eDNA samples showed a higher average biodiversity than those obtained from iDNA



**Fig. 2** Phylum level report of the high-throughput sequencing. 16S rDNA sequences of bacterial rhizosphere communities (stacked bar plot).



**Table 2** Shannon–Wiener H index and analysis metadata

Sample	Shannon index	Treatment	DNA origin
CPR_iDNA	5.228	CP	iDNA
CPR_eDNA	5.009	CP	eDNA
20DR_iDNA	4.941	20DR	iDNA
20DR_eDNA	5.105	20DR	eDNA
40DR_iDNA	5.001	40DR	iDNA
40DR_eDNA	5.069	40DR	eDNA

(iDNA = 5057, eDNA = 5061); however, also in this case the difference was not significant ( $p > 0.05$ ).

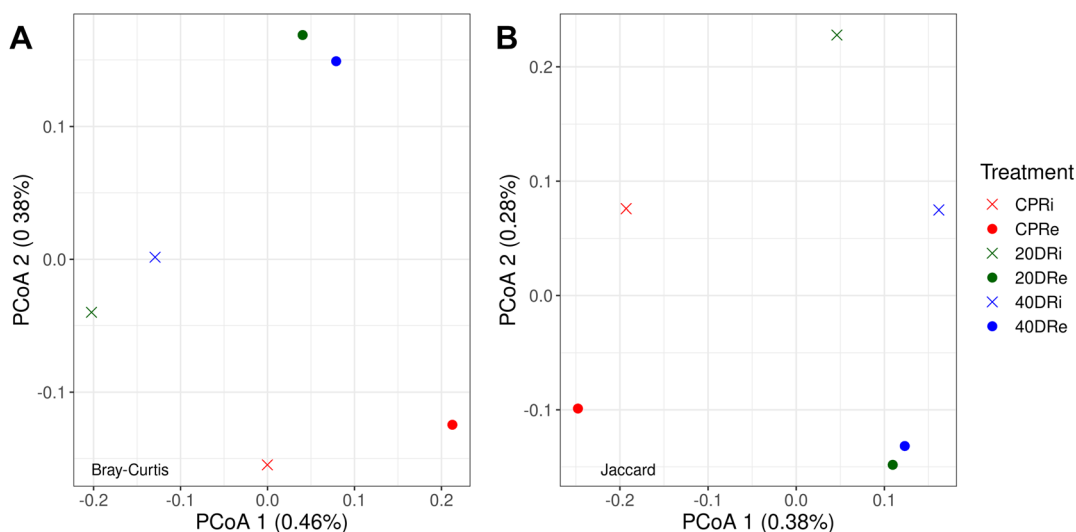
Through PCoA, it was also possible to observe how the different iDNA and eDNA samples were arranged according to the treatment. In Fig. 3A, the Bray–Curtis dissimilarity, which considers both the number of species and the frequency with which they occur, showed that nCeO<sub>2</sub> treated samples clearly separated from CPR control samples, and grouped according to the origin of the DNA (iDNA or eDNA), but independently from the nCeO<sub>2</sub> dose (20DR and 40DR). The same results could be also observed from the Jaccard dissimilarity PCoA (Fig. 3B), which, unlike the Bray–Curtis index, does not consider the frequency with which different species occur. However, the percentages of variance explained by these PCoAs were minimal, indicating that although separated, the samples had very low diversity values.

Following a similar approach, PCA was performed on the bacterial distribution of the samples at the genus granularity level, retaining the first two components. As can be seen in Fig. 4, also in this case, the treated samples grouped according to the origin of the DNA, while the two control samples clearly separated from the treated ones as well as from each other. Overall, the explained variance of the two components was 87.39% (52.91% for the first component and 34.48% for the second component).

### 3.5 Differences between detected ASVs

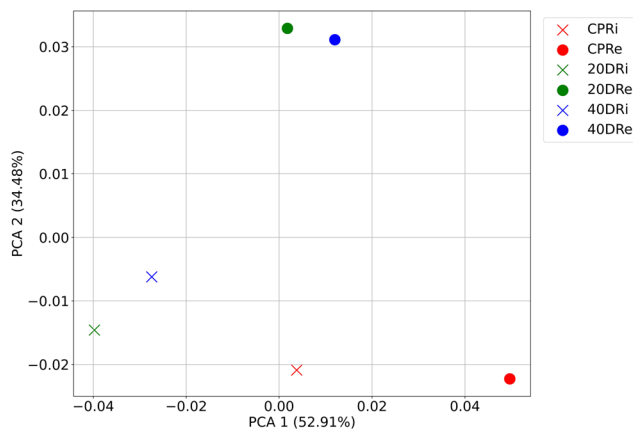
The differences induced on bacterial populations by the application of nCeO<sub>2</sub> were then evaluated by comparing the increase or decrease in the relative percentages at phylum, family and genus level of the treated samples to the control.

Of the 31 identified phyla, 13 were found to be present at >0.5% (iDNA Fig. 5A, eDNA Fig. 5B). One of the most significant differences observed was a consistent decrease of *Firmicutes* in substrates treated with nCeO<sub>2</sub> at both dosages and both in iDNA and eDNA in comparison to the corresponding control samples. In fact, from an average relative percentage of 10.35% in CPR samples, mean percentages of 0.74% and 1.07% were observed in 20DR and 40DR samples. Another decrease in the percentage of the population was observed for *Bacteroidetes*, with an average percentage of 10.33% in CPR against an average percentage of 9.23% in 20DR and 40DR samples. Conversely, in nCeO<sub>2</sub>-treated samples, *Proteobacteria* increased from an average concentration of 44.04% in CPR samples to an average concentration of 48.64% in 20DR and 40DR, *Acidobacteria* moved from 5.14% to 7.12%, as well as *Chloroflexi* and *Actinobacteria* rose from 5.88% to 8.31% and 4.06% to 6.35%, respectively. As for the differences between iDNA and eDNA, a greater proportion of eDNA was found in *Chloroflexi* (6.03% vs. 8.96%), *Actinobacteria* (4.24% vs. 5.80%), *Firmicutes* (3.00% vs. 5.11%), *Verrucomicrobia* (4.24% vs. 5.80%), and *Patescibacteria* (1.12% vs. 1.58%). In summary, considering these 13 main phyla, in comparison to control samples, in the case of the 20DRi sample, 7 phyla decreased while 6 increased, while for 40DRi 6 decreased and 7 increased. For eDNA samples, in both 20DRi and 40DRi, compared to CPRi, 5 phyla decreased and 8 increased. Comparing the values of each phylum for the iDNA and eDNA of the controls, 6 couples behaved differently in samples 20DRi and



**Fig. 3** Bray–Curtis (A) and Jaccard (B) principal coordinates analysis (PCoA) arrangement of samples according to the treatments (CPR, 20DR, and 40DR) and DNA origin (iDNA or eDNA).





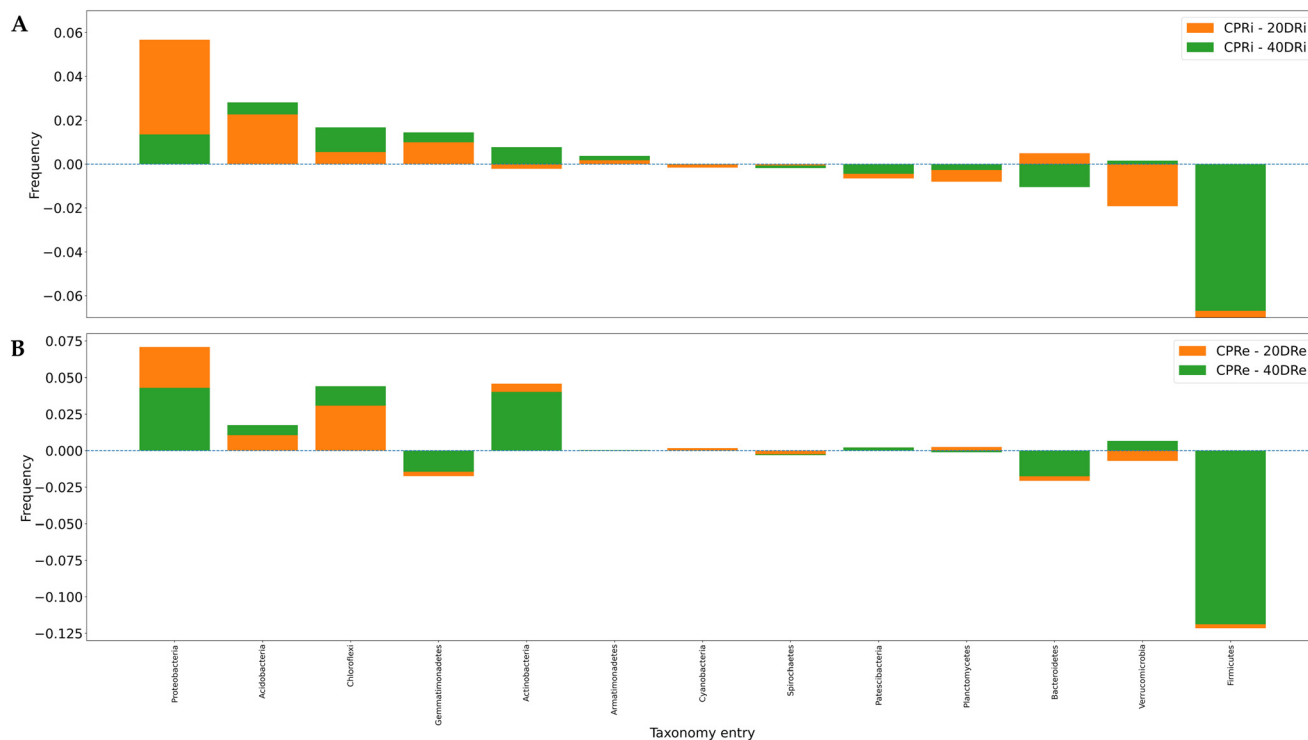
**Fig. 4** PCA (first two components) of bacterial distribution at the genus granularity level according to the treatments (CPR, 20DR, and 40DR) and DNA origin (iDNA or eDNA).

20Dre, whereas 7 couples performed equally. In samples 40DRi and 40DRe, 3 couples behaved differently and 10 had the same trend.

Moving to the family level (iDNA Fig. 6A, eDNA Fig. 6B), 88 out of 336 identified families were present at values  $>0.5\%$  and considered for this analysis. These results highlighted that two nCeO<sub>2</sub> dosages have similar effects to the triple dosage. Also, it was observed that comparing 20DRi to CPRe, 41 families decreased, 6 remained unchanged, and 41 increased, while

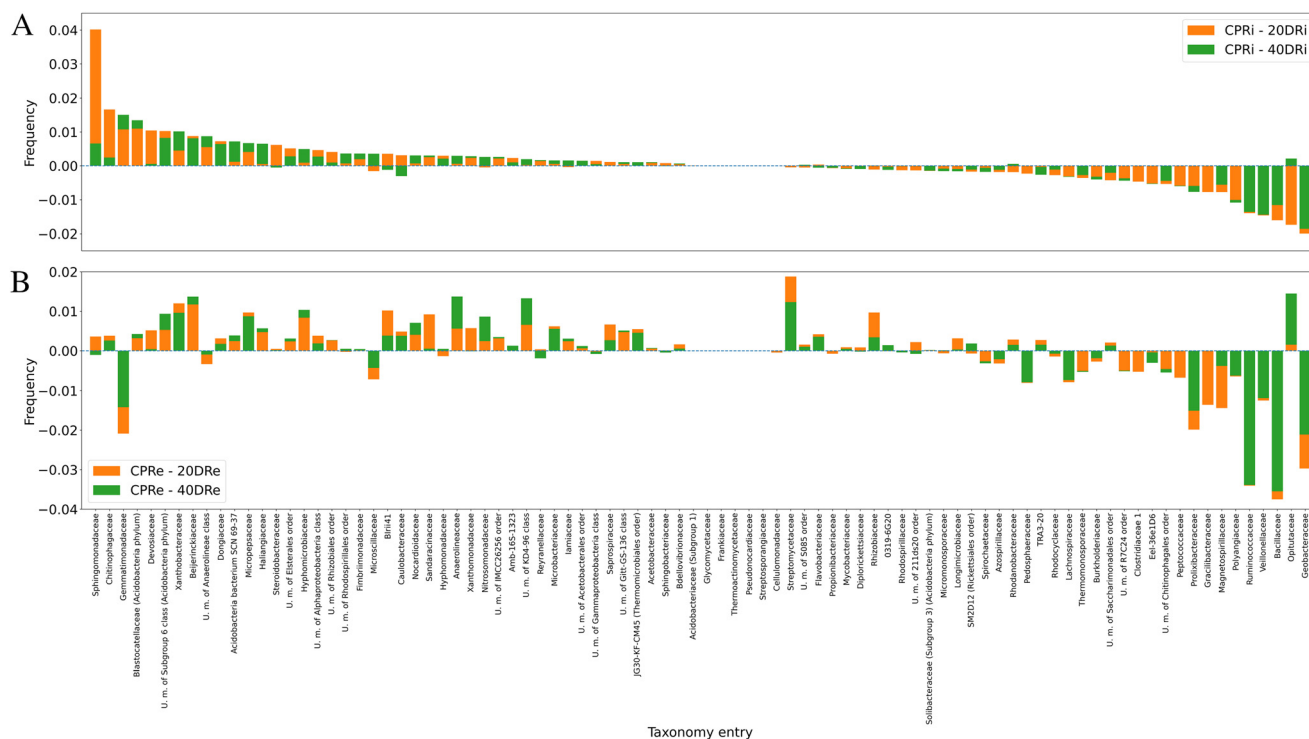
comparing 40DRi, 36 families decreased, 7 remained unchanged, and 45 increased. Similarly, for eDNA samples, comparing 20DRe to CPRe, 31 families decreased, 6 remained unchanged, and 51 increased, whereas for 40DRe, 33 families decreased, 6 remained unchanged, and 49 increased. Furthermore, comparing 20DRi and 20DRe with respect to the corresponding controls, 23 couples behaved differently, and 65 couples showed the same behaviour, whereas in the case of the 40DRi and 40DRe samples, 21 couples behaved differently, and 67 couples behaved in the same way.

Fig. 7A and B depict the deviations compared with the control samples at the genus level for iDNA and eDNA, respectively. Out of 641 detected genera, 112, whose presence was  $>0.5\%$ , were considered for the analysis. The comparison of the treatments with the controls revealed that in the case of 20DRi *versus* CPRe, 42 genera decreased, 13 did not change, and 57 increased, whereas in the case of 40DRi, 39 genera decreased, 12 remained unchanged, and 61 increased. Similarly, considering the eDNA, in 20DRe *versus* CPRe, 31 genera decreased, 12 remained unchanged, and 69 increased, whereas in 40DRe, 34 genera increased, 11 were stable, and 67 increased. Furthermore, comparing 20DRi *versus* 20DRe with reference to the corresponding controls, 31 couples behaved in the opposite way, and 81 couples reported the same trend. For 40DRi *versus* 40DRe, 23 couples behaved in the opposite way, while 89 couples had the same behaviour.

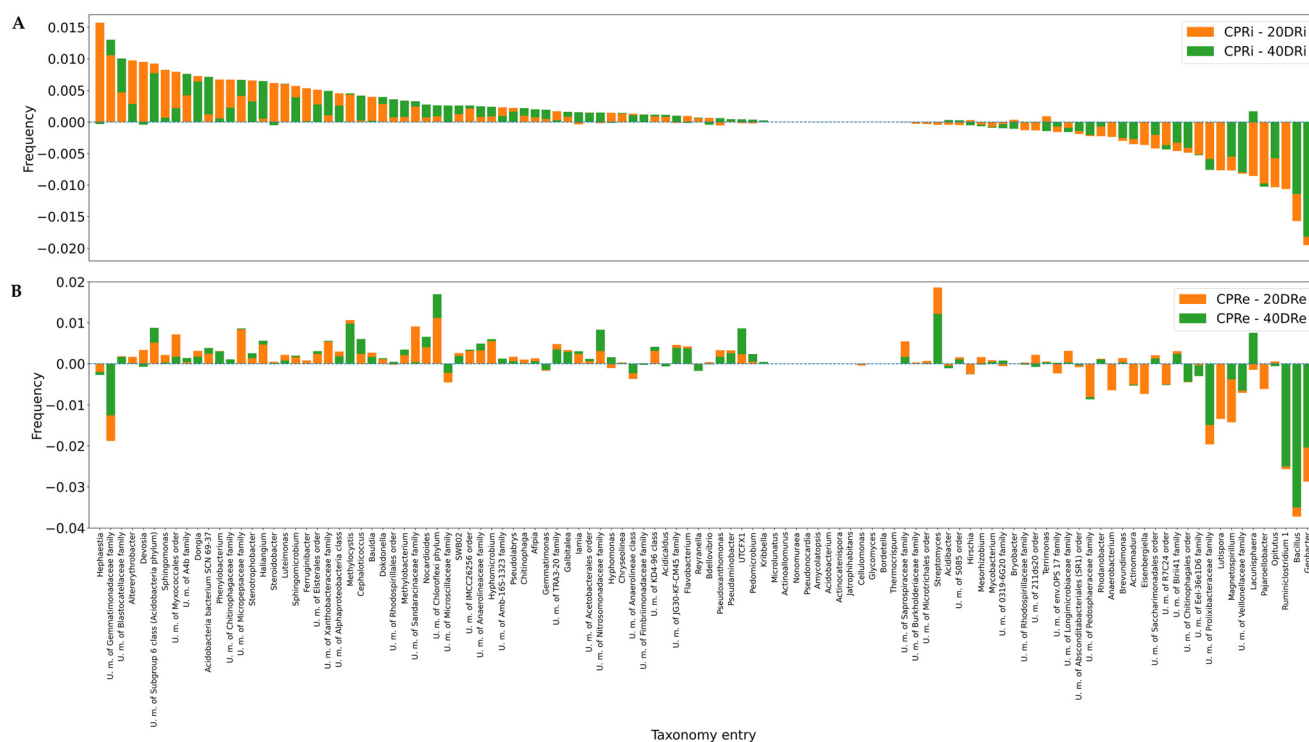


**Fig. 5** The differences in the distribution of microbiological groups for iDNA and eDNA for the different samples in comparison to the control were shown in panel (A) and (B) respectively. Bars represent different taxonomy entries and are not stacked, but rather superimposed, with each bar starting from a 0.00 baseline and the lower value presented in the forefront. Sorting of bars is determined by the frequencies of CPRe. Granularity is at the phylum level.





**Fig. 6** Deviations in the distributions of the main microbiological groups in the different samples compared to the control, granularity at the family level (6A, iDNA; 6B, and eDNA). Note that bars are not stacked but superimposed: for each taxonomy entry, both bars start from the value of 0.00, and the lowest is in the foreground with respect to the other. Bars in both graphs are sorted according to the frequencies of CPRe.



**Fig. 7** Deviations in the distributions of the main microbiological groups in the different samples compared to the control, granularity at the genus level (7A, iDNA; 7B, and eDNA). Note that bars are not stacked but superimposed: for each taxonomy entry, both bars start from the value of 0.00, and the lowest is in the foreground with respect to the other. Bars in both graphs are sorted according to the frequencies of CPRe.



**Table 3** Spearman correlation between samples

Granularity	Samples	Corr	<i>p</i> -Value	Granularity	Samples	Corr	<i>p</i> -Value
Phylum	CPRi/20DRi	0.86	$1.50 \times 10^{-4}$	Family	CPre/20DRe	0.6	$6.30 \times 10^{-10}$
Phylum	CPRi/40DRi	0.85	2.20	Family	CPre/40DRe	0.7	$3.90 \times 10^{-14}$
Phylum	20DRi/40DRi	0.99	$1.70 \times 10^{-10}$	Family	20DRe/40DRe	0.93	$2.40 \times 10^{-38}$
Phylum	CPre/20DRe	0.73	$5.00 \times 10^{-3}$	Genus	CPRi/20DRi	0.72	$7.50 \times 10^{-19}$
Phylum	CPre/40DRe	0.82	$6.20 \times 10^{-4}$	Genus	CPRi/40DRi	0.75	$4.00 \times 10^{-21}$
Phylum	20DRe/40DRe	0.97	$2.60 \times 10^{-8}$	Genus	20DRi/40DRi	0.93	$1.00 \times 10^{-49}$
Family	CPRi/20DRi	0.75	$2.50 \times 10^{-17}$	Genus	CPre/20DRe	0.6	$4.20 \times 10^{-12}$
Family	CPRi/40DRi	0.77	$1.40 \times 10^{-18}$	Genus	CPre/40DRe	0.7	$1.50 \times 10^{-17}$
Family	20DRi/40DRi	0.95	$4.10 \times 10^{-45}$	Genus	20DRe/40DRe	0.88	$2.70 \times 10^{-38}$

### 3.6 Ranking analysis: Spearman correlation

The ranking of various bacterial groups in the different samples was considered. To such an extent, the Spearman correlation was calculated pairwise between the samples at different granularity levels to evaluate the degree of concordance (between  $-1$  and  $1$ ) of the ranks (Table 3).

Generally, among the samples analysed using the same technique and treated with different quantities of nCeO<sub>2</sub>, the concordance between rankings was sufficiently high (higher than 0.92, except for one case with 0.88). In contrast, the agreement was significantly lower among the samples treated with nCeO<sub>2</sub> and the related control samples (always with the same analysis). This confirmed the qualitative results previously provided by the PCoA and PCA. Moreover, from Table 3, it can be observed that for the same pairs of samples, passing from the phylum to families and genera levels, the 20DRi/40DRi correlation was more homogeneous and similar, with values between 0.93 and 0.99. At the same time, the CPre/20DRe showed the greatest diversity with values between 0.60 and 0.73.

Fig. 8 and 9 highlight how the rankings of the different bacteria changed among the samples, reported the position in the rank for each bacterial group at the phylum granularity level, considering iDNA and eDNA. In both cases, in the treated samples a marked reduction in the presence of *Firmicutes* with respect to the untreated samples was observed. In contrast, there was an increase in both *Chloroflexi* and *Actinobacteria* (more pronounced in the iDNA samples than in the eDNA cases). *Acidobacteria* and

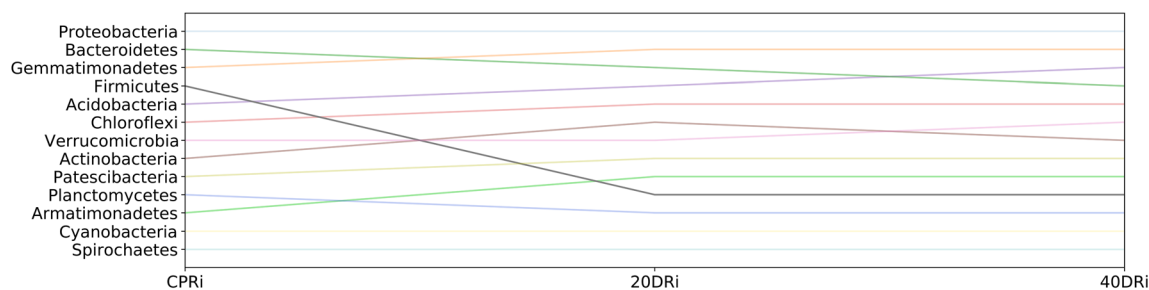
*Patescibacteria* showed similar increases in both cases. Interestingly, *Armatimonadetes* decreased when eDNA was considered and increased when iDNA was considered.

### 3.7 Metabolisms

The analysis of the metabolic functions, reconstructed using the FAPROTAX database, revealed differences induced by nCeO<sub>2</sub> to the reconstructed metabolic functions. As observed from the heatmap shown in Fig. 10, metabolic functions related to fermentation, nitrogen fixation, and iron, nitrate, and nitrogen respiration decreased in the samples treated with cerium (20DR and 40DR) compared to those in the control samples (CPR), in association with an increase in functions related to chemo-heterotrophy.

## 4 Discussion

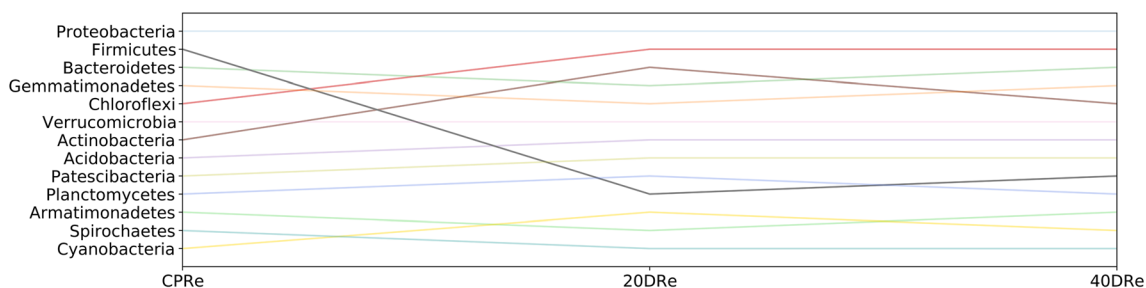
The aim of this study was to deepen the knowledge of the influence of CeNPs on the soil microbiota, and more specifically on the rhizosphere microbiota during the vegetative growth. For this purpose, iDNA and eDNA obtained from the rhizospheric soil of *Silene flos-cuculi* amended with different nCeO<sub>2</sub> concentrations in a controlled environment were considered. To date, studies have focused mainly on the relationships among soil, plant and CeNPs, but not on the relationship between CeNPs and microbiota.<sup>48,49</sup> Soil microbial communities are the poorly understood driver of many processes that directly or indirectly govern the fluxes of carbon, nitrogen, and other elements through ecosystems.<sup>47,50–52</sup> In particular, rhizospheric microbiota and



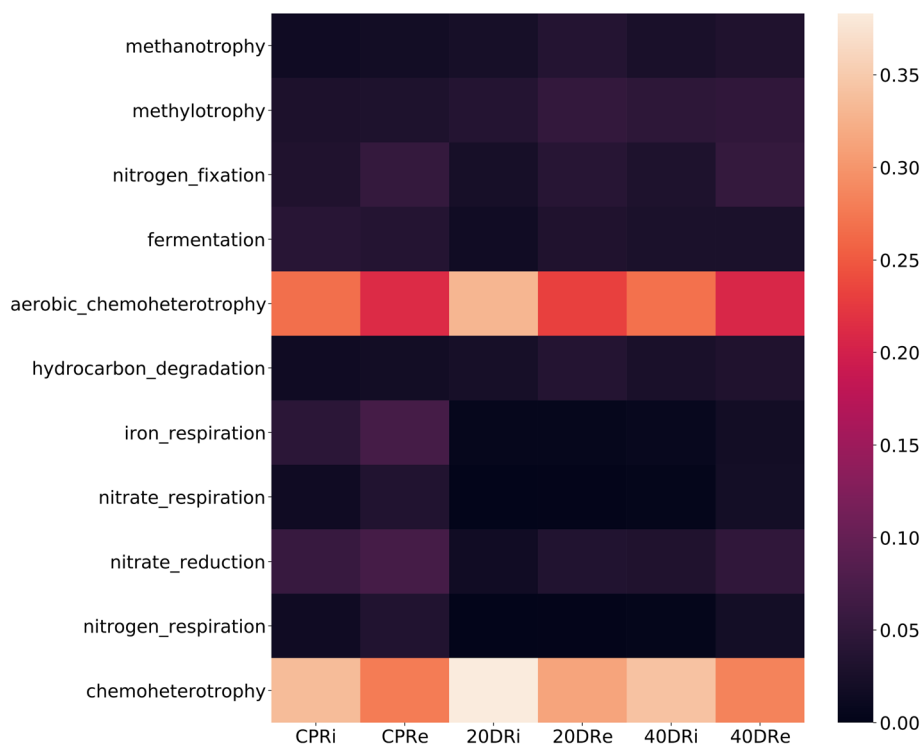
**Fig. 8** Ranking of the main microbiological groups and their changes in the different samples, iDNA case, and granularity at the phylum level. Lines of softer colors correspond to smaller changes in rankings. A higher height on the ordinate axis corresponds to a higher position in the ranking (more marked presence in the sample).







**Fig. 9** Ranking of the main microbiological groups and their changes in the different samples, eDNA case, and granularity at the phylum level. Lines of softer colors correspond to smaller changes in rankings. A higher height on the ordinate axis corresponds to a higher position in the ranking (more marked presence in the sample).



**Fig. 10** Heatmap of metabolic functions reconstructed using the FAPROTAX database. Each column corresponds to a sample and each row corresponds to a functional group. The colors shown represent the occurrence of the metabolic function (number of ASVs capable for each function, from 0 to 1).

the symbiotic processes triggered by plant growth-promoting rhizobacteria (PGPR) play a fundamental role in improving productivity by reducing fertilization and pesticide utilization.<sup>53</sup> In this regard, nCeO<sub>2</sub> can impact this mutualistic relationship in different ways, both directly and indirectly. Due to their physicochemical characteristics, CeNPs are translocated in plants through the root system to leaves, determining a specific plant modifiable adaptation response that also influences the microbiota.<sup>29</sup> Furthermore, the retention or release of oxygen from the nCeO<sub>2</sub> structure<sup>8,9,54</sup> can modify the biogeochemical conditions, inhibiting gas flow across the sediment/water interface profile<sup>55–57</sup> and therefore modifying the oxygen gradient, thus directly reducing the aerobic and facultative bacterial microflora.<sup>58</sup>

Several studies have also proved the direct antimicrobial effect of CeNPs, and their possible application for medical use<sup>59,60</sup> against both Gram-positive and Gram-negative bacteria.<sup>61,62</sup> At low pH values, nanomaterials (positively charged) adhere to microorganisms (negatively charged), acquiring an electron, which converts Ce<sup>4+</sup> to Ce<sup>3+</sup>, deranging the cell wall structure. Gram negatives and fungi are more susceptible to this effect, while Gram positives, thanks to a thick peptidoglycan layer, can modulate this effect.<sup>9,16,62,63</sup>

Therefore, biological spheres are subject to complex interactions related to the dosage and physical properties of nanoparticles present in the environment.<sup>8,10,64</sup> Lizzi *et al.* (2020)<sup>27</sup> demonstrated that CeNPs with a nominal dimension of 25 nm and 99.95% purity showed an average dimension of



126.7 ± 1.0 nm but CeNPs translocated to roots and leaves were respectively 33 ± 2 nm and 31 ± 1.5 nm. This underlined the greater catalytic action of the smaller particles,<sup>65</sup> and demonstrated that the reduction of CeNPs to Ce<sup>3+</sup> is necessary to facilitate the absorption of cerium at the root level.<sup>21,66</sup>

Given the complexity of these biotic and environmental factors, to evaluate the differences induced in this study, a NGS approach was chosen for its capable of evaluating the composition of the entire population with a high resolution degree and also allowing detection of minimal differences. In this regard, the evaluation of alpha diversity analysis results underlined a lower Shannon–Viener *H* index value in nCeO<sub>2</sub> treated samples. The same evidence of an effect on microbial populations emerged from the results of PCoA. In fact, both in Bray–Curtis and Jaccard dissimilarity matrices, the treated samples clustered separately from the control samples. However, these differences, although present, were minimal, indicating a limited effect of the treatments on the differences in the biodiversity level of the samples. It should also be underlined that the differences induced by the presence of nCeO<sub>2</sub> were poorly correlated with its concentration, as suggested by the similarity of the results obtained between samples 20DR and 40DR. From these results we can therefore observe a non-significant effect on the overall biodiversity level induced by the amendment with nCeO<sub>2</sub>. Although there was an effect on the proportion of some species present, no impoverishment of the soil was therefore observed at a biological level. In contrast, it was possible to observe a clear separation between the iDNA and eDNA samples, testifying to the importance of distinguishing these two components of the total DNA to obtain an effective analysis, capable of distinguishing the still vital microbial population from the paleome. Subsequently, focusing the analysis on the present phyla, a drastic decrease in *Firmicutes* was observed. This difference has certainly induced significant effects on a biological level, in fact *Firmicutes* are a phylum of bacteria that exerts important metabolic functions in the soil, contributing to terrestrial ecosystems in different ways. Several of these bacteria are involved in the decomposition of organic matter and nitrogen fixation<sup>67,68</sup> producing beneficial chemicals and competing with plant pathogens. Therefore their disruption and imbalance in soil could result in the emergence of several plant diseases.<sup>69</sup>

These results were in contrast with other studies, in which this difference was not reported<sup>70,71</sup> or even reported an increase of this phylum.<sup>72</sup> This evidence, however, was obtained under completely different conditions from those applied in this study. For instance, some experiments were conducted on activated sludge, in a liquid environment maintained under aerobic conditions, and the increase in *Firmicutes* was observed due to an increment of *Bacilli*. In contrast, in this study the reduction of *Firmicutes* was attributed to a decrease in *Clostridia*, reflecting disparate environmental and microbial dynamics.

Going into further detail to the genus level, of the 112 genera identified at a percentage >0.5% (out of 641 total), 25 of these were present in the controls but totally absent in the 20D and 40D treated samples (for both iDNA and eDNA) (Table S2†). Of these, 10 were Gram-negative, 12 were Gram-positive, while *Lutispora*, *Eisenbergiella*, and *Anaerovorax*, despite the resulting Gram-negative bacteria from Gram staining possessing a cell wall typical of Gram-positive bacteria. In light of these results, it was therefore not possible to associate the different antimicrobial effects of nCeO<sub>2</sub> based on the different cell wall structure as reported in the literature.

On the other hand, considering the possible differences induced by the modification of oxygen exchange due to the presence of nCeO<sub>2</sub>, the oxygen demand of the 25 genera was analyzed: 4 were aerobic, 1 facultatively anaerobic and the remaining 19 genera corresponded to anaerobic and strictly anaerobic bacteria. Currently, few studies focus on the redox cycle between the Ce<sup>3+</sup> and Ce<sup>4+</sup> states on the surface of nCeO<sub>2</sub> in the proximity of the rhizosphere, and its effects on oxygen uptake/release,<sup>73</sup> and further research is required to understand the modifications of the aerobic/facultative/anaerobic microorganism ratio from the nCeO<sub>2</sub> induced ROS.

Focusing on the effects of nCeO<sub>2</sub> on PGPR, important bacteria for soil and plant health,<sup>53,74–82</sup> the majority of which with their relative functions are summarized in Table S3,† it was possible to identify the presence of 13 PGPR genera (*Allorhizobium*, *Azospirillum*, *Pseudomonas*, *Bacillus*, *Bradyrhizobium*, *Chryseobacterium*, *Enterobacter*, *Mesorhizobium*, *Paenibacillus*, *Pantoea*, *Pseudomonas*, *Rhizobium*, and *Stenotrophomonas*). From the comparison of their presence in the different samples (Table S4†), it was possible to observe lower percentages in iDNA samples compared to eDNA. Additionally, it was possible to observe an increase in 5 genera and a decrease in 8 genera in nCeO<sub>2</sub> treated samples for iDNA.

Overall, these induced differences in the PGPR population could be directly responsible for the differences in plant growth, and given the discordant results between different studies, could indicate a specific effect between certain PGPRs and different plant species. Other studies report effects on plant growth, demonstrating both beneficial and detrimental outcomes depending on the doses employed. At high concentrations of nCeO<sub>2</sub>, an increase in *Azotobacter*, *Clostridium*, *Rhodospirillaceae*, and *Ensifer* was observed in the soybean-cultivated soil, while the populations of *Sphingomonas* and *Rhizobium* significantly decreased. Furthermore, the impact of nCeO<sub>2</sub> was notable only in the soil cultivated with soybean and not to the unplanted soil, where bacterial communities remained unaffected.<sup>83</sup> However, with the currently available knowledge, which often consists of experiments conducted on a laboratory scale, with results currently at a preliminary level, which are often contradictory, it is not yet possible to associate a specific effect on plant and soil health with the variation of each individual microbial group given the complexity of the interactions, and the functional redundancy in soil microbial



community.<sup>84,85</sup> The importance of further in-depth studies on nanomaterials to comprehend their actual effects on the environment, soil, and plants has also been emphasized by the Organization for Economic Co-operation and Development (OECD).<sup>86</sup>

To analyse the impact on all metabolic functions induced by diversity changes, the FAPROTAX database was used. Through this useful tool, it is in fact possible to investigate and assign in a more in-depth and effective way the ecological functions of soil bacteria.<sup>87</sup> This analysis showed that nCeO<sub>2</sub> treatment in 20DR and 40DR samples decreased fermentation, nitrogen fixation, iron, nitrate, and the nitrogen respiration function compared to the controls. The predominant function in all samples was chemo-heterotrophy, followed by aerobic chemo-heterotrophy, that was found to be comparable among CPRi and CPRe and 40DRi and 40DRre treated samples. Instead, in the 20DRi sample, both functionalities increased compared to those in 20DRre and the respective controls. This positive selection of aerobic heterotrophs was in accordance with the previously discussed reduction of anaerobic and strictly anaerobic genera in treated samples. Previous studies also reported that exposure to nCeO<sub>2</sub> leads to the degradation of the denitrifying process. This observation aligns with the inhibition of enzyme activities associated with nitrite oxidoreductase and nitrate reductase. Concurrently, a reduction in microbial diversity, a decrease in the network complexity of the bacterial community, and enhanced competitive dynamics among bacterial populations were observed. Specifically, there was a notable decrease in *Flexibacter* and *Acinetobacter*.<sup>88</sup> However, different results were observed for the sediment/water interface.<sup>58</sup> In this scenario, with the presence of halogens in the rhizosphere (e.g., micromolar concentrations of bromides, chlorides, or reactive oxygen species), CeNPs may have exhibited a mimetic action similar to peroxidase enzymes. This action involves the production of hypo-halogen acids (HOCl and HOBr), which interfere with quorum sensing molecules, thereby influencing the metabolism of both Gram-positive and Gram-negative bacteria.<sup>89</sup> Under these conditions, CeNPs present in soils with low chloride concentrations and reduced water content can therefore interact with the rhizosphere, inhibiting microbial quorum sensing molecules and altering the density of bacterial populations, the production of extracellular enzymes, and nitrogen and phosphorus cycles. The actions of mycorrhizae and PGPR can mitigate these effects by intervening in the ROS compounds produced in the root system, thereby reducing the mimetic and catalytic peroxidase-like activity of CeNPs.<sup>90–96</sup> Considering climate change, these conditions characterized by reduced water and humidity content in soils are expected to become increasingly prevalent. In such scenarios, in addition to the abovementioned effects, different studies reported a direct beneficial effect of nCeO<sub>2</sub>. Under water stress conditions, nCeO<sub>2</sub> affected the activity of Rubisco, thereby enhancing photosynthetic activity, mitigating oxidative stress induced by drought-related ROS, and consequently improving plant growth under these challenging conditions.<sup>97,98</sup> However, in addition to potential applications in agronomy, the direct

effects on humans resulting from the consumption of products grown in the presence of nCeO<sub>2</sub> must be thoroughly assessed. From the initial studies conducted, these nanoparticles appear to influence the gut microbiota, potentially having negative implications and thereby compromising their use as agronomical adjuvants.<sup>99–101</sup>

## 5 Conclusions

This study stems from a previous study whose results showed that nCeO<sub>2</sub> may have caused a slowdown in the vegetative growth rate of *S. flos-cuculi* (L.). The possible causes of this phenomenon were therefore sought in a modification induced by this compound in the rhizosphere microbiota.

The most abundant bacterial phylum found in the rhizosphere of the *S. flos-cuculi* substrate was *Proteobacteria* followed by *Gemmatimonadetes*, *Bacteroidetes*, *Chloroflexi*, *Acidobacteria*, *Actinobacteria*, and *Verrucomicrobia*. In contrast, *Firmicutes* which constituted 7.58% and 13.12% in the CPRi and CPRe untreated samples were greatly reduced in soil treated with nCeO<sub>2</sub> amendments, reaching thresholds lower than 1% in nCeO<sub>2</sub> treated soils.

By extending the comparison also to the phyla present in a lower percentage, the inhibition of some anaerobic and strictly anaerobic genera was observed.

Considering the families affected by the same effects both in iDNA and eDNA samples, approximately 50% of them showed inhibition compared with the control, particularly for *Prolixibacteraceae*, *Bacillaceae*, *Peptococcaceae*, *Ruminococcaceae*, *Veillonellaceae*, *Geobacteraceae*, *Thermomonosporaceae*, *Clostridiaceae*, *Azospirillaceae*, *Magnetospirillaceae*, *Gracilibacteraceae*, *Lachnospiraceae*, *Polyangiaceae* and *Burkholderiaceae*.

From these results, and in consideration of the existing studies, nCeO<sub>2</sub> particles have an influence on soil microbiota, which impacts plant health. However, the definition of a general trend of impact is not possible, due to so many variables that can affect the complex soil–root–microbiota system and the few studies present in the literature on the topic. For this reason, we strongly encourage investigations on the topic to improve the knowledge on the effects on different cultivations, which is extremely important in the case of crop rotations given the nCeO<sub>2</sub> soil accumulation, but also in consideration of better understanding the plant translocation mechanisms and its build-up in plant edible parts that could affect human and animal health.

## Data availability

The data are available, and the hyperlink to the published data is present in the manuscript.

## Author contributions

MC, LI, LM: conceptualization; MC, LI: resources; AB, MC, AC, LI, NS: methodology; MC, LI: supervision; AB, AC, AF, NS: formal analysis; AB, AC, NS: data curation; AB, AC, NS:



visualization; AB, MC, AC, NS: writing – original draft; AC, LI: writing – draft revision; MC, AC, LI: writing – review and editing; MC, LI, LM: funding acquisition.

## Conflicts of interest

The authors declare no conflict of interest.

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