Nano-La$_2$O$_3$ Induces Honeybee (Apis mellifera) Death and Enriches for Pathogens in Honeybee Gut Bacterial Communities

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Honeybees (Apis mellifera) can be exposed via numerous potential pathways to ambient nanoparticles (NPs), including rare earth oxide (REO) NPs that are increasingly used and released into the environment. Gut microorganisms are pivotal in mediating honeybee health, but how REO NPs may affect honeybee health and gut microbiota remains poorly understood. To address this knowledge gap, honeybees were fed pollen and sucrose syrup containing 0, 1, 10, 100, and 1000 mg kg$^{-1}$ of nano-La$_2$O$_3$ for 12 days. Nano-La$_2$O$_3$ exerted detrimental effects on honeybee physiology, as reflected by dose-dependent adverse effects of nano-La$_2$O$_3$ on survival, pollen consumption, and body weight ($p < 0.05$). Nano-La$_2$O$_3$ caused the dysbiosis of honeybee gut bacterial communities, as evidenced by the change of gut bacterial community composition, the enrichment of pathogenic *Serratia* and *Frischella*, and the alteration of digestion-related taxa *Bombella* ($p < 0.05$). There were significant correlations between honeybee physiological parameters and the relative abundances of pathogenic *Serratia* and *Frischella* ($p < 0.05$), underscoring linkages between honeybee health and gut bacterial communities. Taken together, this study demonstrates that nano-La$_2$O$_3$ can cause detrimental effects on honeybee health, potentially by disordered gut bacterial communities. This study thus reveals a previously overlooked effect of nano-La$_2$O$_3$ on the ecologically and economically important honeybee species *Apis mellifera*.

Keywords: *Apis mellifera*, rare earth oxide nanoparticles, honeybee gut microbiota, nano-La$_2$O$_3$, honeybee health
INTRODUCTION

Honeybees (e.g., *Apis mellifera*) provide essential pollination services for agricultural ecosystems and valuable apiary products for human nutritional needs (Klein et al., 2007). Due to their extensive social activities within 14 km² foraging areas, honeybees are exposed to conventional contaminants, for example, pesticides, antibiotics, and respirable suspended particulate matters, that may lead to the decline of honeybee colonies or the deterioration of honeybee health status (Bargànska et al., 2016; Thimmegowda et al., 2020; Bondarenko et al., 2021; Kapoor et al., 2021). Among the emerging contaminants, rare earth oxide (REO) nanoparticles (NPs), characterized by their unique chemical and physical properties, have been one of most widely used materials in various industries and biotechnology applications (Mastronardi et al., 2015; Gao et al., 2017; Kos et al., 2017). For example, the fertilizers, pesticides, and germination stimulants containing or engineered with REO NPs have been widely used to enhance the efficiency and sustainability of agricultural practices, and nano-La₂O₃ is reported to account for approximately 30% of REO NPs additives and have higher cytotoxicity compared with other REO NPs (De la Torre Roche et al., 2015; Mastronardi et al., 2015; Servin et al., 2015; Gao et al., 2017). This make honeybees highly susceptible to the exposure and toxicity of REO NPs, through contacting with and ingesting these particles directly or indirectly from the surrounding environments, especially the agricultural ones, such as plant and flower surfaces, pollen and nectar, and soil and dust (Ma et al., 2011; De la Torre Roche et al., 2015; Kos et al., 2017; Radziwill-Bienkowska et al., 2018). Once soil was contaminated by nano-La₂O₃, plants can serve as a potential intermediary pathway that could bioaccumulate and transport them to primary consumers, for example, *Acheta domestica*, *Tenebrioidea*, and honeybees (Ma et al., 2011; De la Torre Roche et al., 2015). Related studies have also demonstrated that honeybees come in contact with metal oxide NPs (e.g., CeO₂, CdO, and PbO) through surface particle adhesion, dust inhalation, foraging on contaminated food, or water (Kos et al., 2017; Al Naggar et al., 2020). Therefore, honeybees may suffer the environmental exposure of terrestrial REO NPs and serve as sensitive indicators of environmental quality. Despite the fact that honeybee gut microorganisms take important roles in maintaining host immunity, metabolism, and health (Kwong and Moran, 2016), it is unknown whether and how REO NPs exposure will cause deterioration of honeybee health and dysbiosis of honeybee gut microbiota and whether the gut dysbiosis will further mediate the toxic effect of environmental contaminants on honeybees health.

Although there are few studies regarding effects of REO NPs on honeybee health, related research implies that metal oxide NPs may adversely affect honeybee health, through mechanisms relating to, for example, signals blocking, reactive oxygen species (ROS), and cell damage (Nel et al., 2006; Benelli, 2018). For example, the mortality rate of *Apis mellifera* increased with exposure concentrations of REO NPs (nano-TiO₂, nano-ZnO-TiO₂, and nano-Ag-TiO₂), implying a dose-dependent toxic effect of metal oxide NPs on honeybee (Dabour et al., 2019). Also, exposure to nano-CeO₂ causes undesirable neurological effects on honeybee *Apis mellifera*, by inhibiting the activity of membrane acetylcholinesterase (AChE) which further influences cholinergic function of the nervous system (Kos et al., 2017). Also, nano-CdO and nano-PbO can enhance ROS production and thus cause free radical-induced oxidative damage to honeybee *Apis millefera*, accompanied by anti-oxidative responses, for example, increased catalase production (Al Naggar et al., 2020).

REO NPs may also disturb honeybee gut microbiota. Previous studies have shown the undesirable effects of various types of NPs on the soil microbiomes and the gut microbiota of animals and insects (Han et al., 2014; Ge et al., 2016; Zhu et al., 2018; Chen et al., 2021). For instance, nano-ZnO and carbonaceous nanoparticles disturb the soil bacterial community structure and change functionally important microbial groups associated with C, N, and S cycling (Ge et al., 2018; Xu et al., 2018a; Wu et al., 2019). Silver NPs alter the gut bacterial communities of *Drosophila* and *Collembola* (Han et al., 2014; Zhu et al., 2018). But, it is still unclear whether, and how, REO NPs affect gut microbiota of honeybees, specifically the model species *Apis mellifera*.
Honeybee gut microbiota could be influenced by various factors, including pathogens, antibiotics, pesticides, diet, and host attributes and environmental conditions (Cox-Foster et al., 2007; Raymann et al., 2017; Liu et al., 2019; Ge et al., 2021). Previous study has demonstrated that the shifts of gut bacterial communities in bumblebees may serve as a characteristic of disease states, featuring as low abundance of core species and high incidence of opportunistic environmental bacteria (Cariveau et al., 2014). Given the significant roles of honeybee gut microbiota in maintaining host health and fitness (Kwong and Moran, 2016; Zhang et al., 2020) and the likelihood of honeybee exposure to REOs via ingestion, it is imperative to understand the ecological effects of REO NPs on honeybee gut microbiota to guide the safe design and application of REO NPs.

In this study, the aims were to (1) investigate the toxicity of REO NPs on honeybee health and gut bacterial communities and (2) explore the relationship between REO NP exposure, gut bacterial communities, and host responses. The working hypotheses were that REO NPs would, in a dose-dependent fashion, disturb honeybee gut microbiota and also directly impact honeybee physiology. A further hypothesis was that the overall effects of REO NPs across physiology and gut microbial effects would be additive. To test these hypotheses, honeybees were fed food amended with different concentrations of nano-La$_2$O$_3$. Here, nano-La$_2$O$_3$ was used as a representative REO NP because of its multifunctionality and high cytotoxicity (De la Torre Roche et al., 2015; Gao et al., 2017). Honeybee survival, pollen consumption, and body weight were quantified, and the composition of the honeybee gut bacterial community was investigated. This is the first study to analyze the effects of nano-La$_2$O$_3$ on honeybees and their gut microbiota, and the results contribute new knowledge regarding the environmental risks of REO NPs.

**MATERIALS AND METHODS**

**The Model Honeybee**

Honeybees (*Apis mellifera*) used for nano-La$_2$O$_3$ exposure experiment were incubated at the Institute of Apicultural Research apiary, Chinese Academy of Agricultural Science following standard protocols (Liu et al., 2019). Briefly, brood frames of a single hive with capped honeybee pupae were placed in an RXZ-380C climate-controlled incubator (Ningbo, China; 34 ± 1°C, 60 ± 10% relative humidity, in darkness) for up to 12 h to obtain honeybee specimens. The newly emerged honeybees (less than 12 h old) were randomly divided into five rearing cages with 120 honeybees per cage and further incubated for 1 week (30 ± 1°C, 45 ± 5% relative humidity, in darkness) by feeding fresh pollen, sterile sucrose solution (50% wt/wt), and deionized water *ad libitum* to initiate microbial colonization in the gut (Ellegaard and Engel, 2019). After one-week pre-incubation, the adult honeybees were exposed to nano-La$_2$O$_3$ NPs (Day 0).

**La$_2$O$_3$ NPs**

Nano-La$_2$O$_3$ (25 ± 5 nm) was obtained from the University of California Center for the Environmental Implications of Nanotechnology and characterized in a previous study (Qi et al., 2019). Briefly, the nano-La$_2$O$_3$ studied was of >99.99% purity and composed of spherical particles that aggregated in deionized water (pH = 6.8) to 589 ± 16 nm. The zeta potential of this nano-La$_2$O$_3$ was previously determined to be 9 ± 1 mV in deionized water (pH = 6.8), and the dissolution extent was determined to be 14% after incubating (24 h at 37°C) in an acidic aqueous solution (HCl, 50 μg mL$^{-1}$, pH = 4.5; Li et al., 2014).

**Experimental Design**

Fresh pollen grains were collected from *Camellia sinensis*, freeze-dried under vacuum in a lyophilizer (Songyuan Huaxing, Beijing, China), and ground into a fine powder using a mortar and pestle. To obtain a homogeneous NP distribution, nano-La$_2$O$_3$ powder was thoroughly mixed with the ground pollen with a handheld kitchen mixer for 10 min, diluted to a series of concentrations (2.5, 25, 250, and 2500 mg kg$^{-1}$ pollen) using a 10-fold dilution method (Ge et al., 2018), and then stored separately at −20°C for daily use. Before daily dietary exposure, the mixture of pollen and nano-La$_2$O$_3$ was dispersed (1:1.5 weight ratio) into an aqueous sucrose solution (50% wt/wt in sterile water) to promote ingestion by honeybees (Jack et al., 2016). Therefore, the final target exposure doses of nano-La$_2$O$_3$ in the mixed pollen and sucrose syrup were 1, 10, 100, and 1,000 mg kg$^{-1}$. Negative exposure control was also conducted by treating honeybees with the mixture of pollen (without nano-La$_2$O$_3$) and sucrose solution (1:1.5 weight ratio).

The exposure doses were chosen to represent several scenarios of dietary exposure: possible environmental concentrations (low or medium concentrations), predicted REO NP environmental hotspots (high concentrations), and potential scenarios (the highest concentrations) based on previous studies (Wen et al., 2001; Tyler, 2004; Gottschalk et al., 2009) and some assumptions. Previous studies reported the concentrations of La in plants (0.004–40 mg kg$^{-1}$; Wen et al., 2001; Tyler, 2004), surface soils (5.5–44 mg kg$^{-1}$; Tyler, 2004), and sediments (5–321 mg kg$^{-1}$; Tyler, 2004; Xu et al., 2018b); and the proportion of oxidation state of La was 35–70% (Wen et al., 2001). Also, La compounds tend to be colloidal or nanoclusters (< 200 nm) in environmental matrices (Ma et al., 2011; Kulaksz and Bau, 2013), and approximate 1–30% nanoparticles can be isolated from bulk soil (Theng and Yuan, 2008). We thus assumed that 1–30% of the La$_2$O$_3$ in environmental matrices was nano-La$_2$O$_3$. Based on this assumption, the estimated concentrations of nano-La$_2$O$_3$ were calculated as 0.00002–10 mg kg$^{-1}$ in plants, 0.02–11 mg kg$^{-1}$ in surface soils, and 0.02–79 mg kg$^{-1}$ in sediments. Therefore, the low and medium concentrations of nano-La$_2$O$_3$ (1 and 10 mg kg$^{-1}$) used in this study were comparable to the estimated concentrations of nano-La$_2$O$_3$ in environmental matrices. Considering that NP distributions in terrestrial environments may be highly heterogeneous such that very high concentrations may occur in localized areas (Gottschalk et al., 2009). The
highest dose was also regarded as simulating an extreme endmember concentration within ranges of reported simulated or measured metal oxide NP concentrations in terrestrial environments (Holden et al., 2014). Using the highest concentration here of 1,000 mg kg⁻¹ also allows for examining future potential scenarios of NP environmental buildup, as a situation being previously considered (Priester et al., 2013; De la Torre Roche et al., 2015).

To examine the impacts of nano-La₂O₃ ingestion, adult honeybees in five rearing cages in climate-controlled incubator (Ningbo Jiangnan, Ningbo, China; 30 ± 1°C with 45 ± 5% relative humidity, in darkness) were orally exposed to different concentrations of nano-La₂O₃ (0, 1, 10, 100, and 1,000 mg kg⁻¹) for up to 12 days (Liu et al., 2019). Each cage contained three sterile Petri dish feeders: one feeder containing 4 g of a freshly prepared pollen and sterile sucrose syrup mixture containing 0, 1, 10, 100, and 1,000 mg kg⁻¹ nano-La₂O₃ for dietary exposure; another feeder containing 20 g sucrose solution (50% wt/wt in sterile water) to minimize the indirect effect of insufficient feeding; and the third feeder containing 30 ml deionized water (Di Pasquale et al., 2013). The amounts provided were more than sufficient for dietary and water needs and were replaced daily with equal amounts during the exposure experiment. A control cage that contained dietary provisions and water, but no honeybees, was also conducted in the incubator simultaneously to measure water evaporation.

**La Residue in Honeybees**

To assess the naturally environmental exposure of honeybees to La, 3 wild honeybees were randomly collected using a sweep net in the field within Beijing Botanical Garden where the Institute of Apicultural Research is located. To measure the La residue in honeybees under controlled laboratory conditions, 3 honeybees of each treatment were sacrificed after 12 days exposure, with each honeybee serving as a biological replicate. Each honeybee was washed and frozen in liquid nitrogen. For each honeybee, the whole body was added with 5 ml concentrated nitric acid (HNO₃) and 1 ml hydrogen peroxide (H₂O₂, 30% v/v) in a 56 ml digestion vessel (Zarić et al., 2016). The digestion was conducted using a microwave digestion system (Anton Paar GmbH, Graz, Austria) based on the following scheme: 15 min from room temperature to 200°C, 15 min at 200°C, and cooling down to the room temperature (Zarić et al., 2016). The digestion solution was diluted to 10 ml with deionized water in volumetric flasks and analyzed with an inductively coupled plasma optical emission spectrometer (PerkinElmer, Palo Alto, CA, United States).

**Honeybee Physiological Parameters**

A live census was taken daily, by counting the number of mobile honeybees in each cage, to assess survivorship; any dead honeybees were removed after the census daily. The total mixture consumption of nano-La₂O₃ was calculated by subtracting the mass of the remaining mixture and water evaporation from the initial 4 g mixture supplied daily. The 40% mass of total mixture consumption was divided by the number of surviving honeybees at each 24 h interval in each cage to calculate the pollen consumption per honeybee per day. Further, to determine the mass of the whole honeybee body, 12 live honeybees were randomly sampled from each cage and randomly separated into three groups of 4 honeybees, transferring into individual 50-ml sterile centrifuge tubes and weighted by the subtraction method with a digital balance (Mettler-Toledo, Shanghai, China; ± 0.001 g).

**Gut Sample Collection**

The entire exposure test lasted for 12 days. After 6 and 12 days, five live honeybees as individual replicates were sampled randomly from each cage and cold-anesthetized (−20°C, 1 min). All immobilized honeybees were dissected on ice to collect the entire gut with flame-sterilized forceps under aseptic conditions. Each gut sample was placed into a 2-ml sterile centrifuge tube and immediately frozen in liquid nitrogen for subsequent DNA extraction and gut bacterial community analysis (Liu et al., 2019).

**DNA Extraction, PCR, and High-Throughput Sequencing**

Gut DNA was extracted from the honeybee gut samples using the FastDNA SPIN Kit (MP Biomedicals, Santa Ana, CA, United States) according to manufacturer’s protocol. Extraction blanks, containing all the components except gut samples, were used as quality controls. The DNA from the whole gut of one bee was dissolved in 100 μl TE buffer, quantified with NanoDrop 2000 (Thermo Scientific, Wilmington, DE, United States), and stored at −80°C until use.

The V3-V4 hypervariable regions of the bacterial 16S rRNA gene were amplified in triplicates with the primer set 338F (5′-ACTCCTACGGGAGGCAGCAG-3′) and 806R (5′-GGACTACHVGGGTWTCTAAT-3′). The oligonucleotides of six-base barcodes were incorporated with the forward and the reverse primers to distinguish sequencing samples. Polymerase Chain Reaction (PCR) was conducted in a 20 μl reaction mixture, containing 4 μl of 5× FastPfu Buffer, 2 μl of 2.5 mM dNTPs, 0.8 μl of each primer (5 μm), 0.4 μl of FastPfu Polymerase (TransGen Biotech, Beijing, China), and 10 ng of template DNA. Each reaction was performed under the following procedures: denaturation at 95°C for 3 min, annealing for 25 cycles (95°C for 30 s, 55°C for 30 s, and 72°C for 45 s), and a final extension at 72°C for 10 min. Reaction mixtures without DNA templates served as negative controls to test for contamination. The size and quality of PCR products were checked by TapeStation (Agilent Technologies, Santa Clara, CA, United States). The triplicate PCR products of each sample were pooled and purified using the AxyPrep DNA Gel Extraction Kit (Axygen Biosciences, Union City, CA, United States) and then quantified using the QuantiFluor-ST (Promega, Madison, WI, United States). After normalization in equimolar amounts, the purified amplicons were sequenced on an Illumina MiSeq platform (Illumina, San Diego, CA, United States) according to standard protocols.
Bioinformatic Analysis

The raw sequencing data were processed using the Quantitative Insights Into Microbial Ecology. First, sequences were filtered according to Liu et al. (2020). Then, the high-quality sequences were merged using FLASH.2 Finally, the merged sequences were clustered into operational taxonomic units (OTUs) at 97% identity threshold using UPPARSE,3 while chimeric sequences were removed using UCHIME. The OTUs were assigned to a taxonomic unit by RDP Classifier4 against the SILVA 16S rRNA database (Release 128)5 using a confidence threshold of 70%. The sequences were subsampled to the minimum depth (30,707) prior to analysis.

Statistical Analysis

The “survival” of honeybees was estimated by the Kaplan-Meier (KM) method taking into account the numbers of survivors and dead honeybees, and the samples sacrificed during the study. The KM method is considered efficient and sufficiently general for estimating survival curve (Motta et al., 2018; Al Naggar et al., 2020). Statistical differences following various treatments were assessed by a log-rank paired test, and the p values were adjusted using a Bonferroni procedure. Cumulative pollen consumption indicated the total mass of pollen consumption from the first day to each time point, and the potential maximum honeybee pollen intake was estimated by fitting the cumulative pollen consumption to a first-order kinetic equation. Honeybee weight loss indicated the loss of body weight at each time point relative to the initial weight (Day 0), and the apparently zero-order rate of weight loss was obtained from a linear regression based on untransformed data. Comparisons of equation coefficients between the control and different treatments were achieved by bootstrapping (1000 times) followed by pairwise t test (Zhou et al., 2008).

One-way ANOVA with a post hoc least significant difference test was performed to test the differences among treatments. The contents of La were log-transformed for normality prior to ANOVA. The rarefied sample-OTU matrix was log-transformed to reduce the influence of highly abundant species (Anderson et al., 2006). Principal coordinates analysis (PCoA; using function “pcoa”), permutational multivariate analysis of variance (PERMANOVA; “adonis”), and the distance-based multivariate dispersion test (“betadisper”) were performed using VEGEN package in R6 to assess the influence of nano-La2O3 on the gut bacterial community composition.

Due to the simple composition and significant inter-individual variation in the gut samples (Kwong and Moran, 2016), the genera that occurred in more than half of 55 samples were defined as the common ones. Spearman correlation was used to identify affected taxa whose relative abundance was significantly correlated (p < 0.05) with nano-La2O3 exposure dose at day 6 and 12. Linear or exponential regression analysis was further conducted to examine the relationship between affected taxa and host physiological parameters (survival, cumulative pollen consumption, and weight loss) and nano-La2O3 exposure dose. Analyses were executed using R7,8 SPSS (SPSS, Chicago, IL, United States), or SigmaPlot (Systat Software, Chicago, IL, United States).

RESULTS AND DISCUSSION

La Content in Honeybees

Wild honeybees in the field were assayed to evaluate the environmental background of La in honeybees. Incubated honeybees with nano-La2O3 exposure were also assayed to assess treatment effects on uptake of dietary exposure of nano-La2O3. The background La residue (0.20 μg bee−1) in wild honeybees was significantly higher than the lab-reared ones with no (exposure, 0 mg kg−1; residual, 0.06 μg bee−1) or low-dosage (1 mg kg−1; 0.04 μg bee−1) exposure of nano-La2O3 (p < 0.05; Figure 1), suggesting that honeybees did suffer La exposure under field condition. For incubated honeybees, the La residues increased significantly with La2O3 exposure doses (Spearman’s R = 0.86, p < 0.001; Figure 1). Notably, La residue in the honeybees treated with medium dosage (10 mg kg−1) nano-La2O3 exposure were comparable to the background La residue in wild honeybees (p > 0.05; Figure 1). Therefore, the medium dosage of nano-La2O3 used in this study could be taken into account to predict the natural La exposure to honeybees, although it remains a challenge to characterize the complex forms of La during translocation in environmental matrices (Ma et al., 2011; Li et al., 2014). La residues in high (100 mg kg−1) and the highest dosage (1,000 mg kg−1) treatments were approximately 7 and 55 times that of wild honeybees (p < 0.05; Figure 1). It is reported that high concentrations of La could

1http://cbcb.jhu.edu/software/FLASH/
2http://drive5.com/uparse/
3http://rdp.cme.msu.edu/
4http://www.arb-silva.de/
5https://www.r-project.org/
6http://www.r-project.org/
result in irreversibly adverse impacts to plants and *Daphnia magna*, despite of its neutral effects on *Chlorella* sp. (Balusamy et al., 2015; Yue et al., 2017). In addition, the measured La residue was lower than the corresponding cumulative consumption (Supplementary Figure 1), indicating that nano-La$_2$O$_3$ was released to the intestinal environment and partially excreted through the gut. Therefore, we further conducted honeybee toxicity assays to explore the potential effects of nano-La$_2$O$_3$ on honeybee health and gut microbiota.

**Nano-La$_2$O$_3$ Deteriorates Honeybee Health**

The Kaplan-Meier survival curves of honeybees showed that survival significantly decreased in the high or highest NP
concentration treatments compared to the control (p < 0.001), while there were negligible decreases in low and medium treatments (p = 1 and p = 0.07, respectively; Figure 2; Supplementary Table 1). This suggests a dose-dependent toxic effect of nano-La$_2$O$_3$ on honeybee's survival, which is in line with a previous study (Dabour et al., 2019). Previous studies indicated that nano-La$_2$O$_3$ is chronically toxic to the lung that can strip membrane phosphate groups in acidifying lysosomes and induce cellular and pulmonary damage (Li et al., 2014). Pollen provides most of the nutrients (e.g., proteins, amino acids, and lipids) for honeybee physiological development (Di Pasquale et al., 2013). The results revealed major differences across the time course of cumulative pollen consumption in different treatments (Figure 3A,B; Supplementary Table 2), indicating dose-dependent toxic effects of nano-La$_2$O$_3$ on honeybee nutrition intake (Glavan et al., 2017). In addition, there were positive relationships between weight loss and exposure days, such that the rate of the weight losses for the two highest dose treatments was significantly greater than that of the control (p < 0.05, Figures 3C,D; Supplementary Table 3). Thus, exposure to sufficient doses of nano-La$_2$O$_3$ can decrease the cumulative body weight of honeybees. Body weight is a sensitive indicator of nutritional and energetic effects, which are tightly linked with gut microbiota (Zheng et al., 2017).

Nano-La$_2$O$_3$ Causes Dysbiosis of Honeybee Gut Bacterial Communities

To access whether honeybee gut bacterial community will respond to nano-La$_2$O$_3$ exposure, we extracted gut DNA (Supplementary Figure 2) and conducted bacterial 16S rRNA gene sequencing. The results of PCoA, PERMANOVA, and distance-based multivariate dispersion showed that nano-La$_2$O$_3$ exposure can cause significant shift of gut bacterial community composition (p < 0.05; Figures 4A–F), with or without the consideration of the effect of exposure days (day 6 and 12; p < 0.05; Figures 4A,D). These results suggest that Nano-La$_2$O$_3$ exposure can cause significant honeybee gut bacterial compositional dysbiosis in a relative short term of within 6-day exposure. When the dose of nano-La$_2$O$_3$ exposure was assessed, a gradual shift of gut bacterial community composition with increasing exposure dose of nano-La$_2$O$_3$ was observed at both day 6 (linear regression, p < 0.05) and day 12 (p = 0.07), while the significant effects were only observed for the highest dose (1,000 mg kg$^{-1}$) at both days 6 and 12 as evidenced by the pairwise comparison of gut community compositional differences (p < 0.05; multivariate dispersion test; Figures 4E,F). These...
Nano-La$_2$O$_3$ Exposure Affects Honeybee Health by Causing Dysbiosis of Its Gut Bacterial Communities

Gut microbiota play crucial roles in host health (Engel et al., 2016; Zhang et al., 2020). The relationships between the abundances of main bacterial taxa and the physiological parameters of honeybees (survival, cumulative pollen consumption, and weight loss) were assessed with linear or exponential regression (Supplementary Table 4). The result showed that the abundances of pathogenic *Serratia* and *Frischella* were significantly related to honeybee physiological parameters at either day 6 or day 12 ($p<0.05$; Figure 6).

**Figure 5** | Heatmap illustrating the relative sequence abundances (log2-transformed; 30,707 depth basis) of common bacterial genera in the guts of honeybees of different treatments. Asterisk indicate significant relationship (Spearman’s correlation) between the relative abundances of core genera and exposure dose of nano-La$_2$O$_3$ (*$p<0.05$, **$p<0.01$, ***$p<0.001$).
Our result showed a significant decrease of honeybee survival with the increasing abundances of the pathogenic genera *Serratia* and *Frischella* at day 12 (Figures 6A,B). This is in line with previous studies, showing that the enrichment of pathogens, for example, the genera *Serratia* and *Frischella*, deteriorates host development and increases host mortality (Doublet et al., 2015; Maes et al., 2016; Raymann et al., 2018). Our results, as well as the previous evidence, imply that the response of the gut microbiota and the specific functional taxa, including the pathogenic ones, may act important roles in mediating the effects of contaminants, such as nano-La$_2$O$_3$, assessed in this study, on host health (Raymann et al., 2017; Motta et al., 2018). Notably, these correlations did not provide conclusive evidence that the enrichment of the pathogenic *Serratia* and *Frischella* directly cause honeybee death, but they are suggestive. Further, perhaps in vivo, infection experiments are needed to establish the virulence of pathogens from various contaminant exposures and to assess whether pathogenesis derives directly from contaminant-promoted thrive of pathogen or indirectly from contaminant-induced inhibition of beneficial taxa.

Gut dysbiosis induces dramatic effects on honeybee health (Hamdi et al., 2011). Gut microbiota affect host weight by mediating host nutritional physiology (e.g., vitellogenin level; Zheng et al., 2017). Hence, gut dysbiosis can cause metabolic disorders and impair host development through altering hormone production (Cryan and Dinan, 2012). In this study, the abundance of *Serratia* was significantly related to cumulative pollen consumption and weight loss at day 12 (Figures 6C,E). The abundance of *Frischella* exponentially increased, while the body weight decreased under nano-La$_2$O$_3$ exposure (Figures 6D,F). According to these results, an inference is that the markers of...
gut dysbiosis, pathogenic *Serratia* and *Frischella*, may affect hormonal signaling-related gene expression and subsequently induce pathophysiological responses (e.g., impaired nutrition intake and body weight; Maes et al., 2016; Raymann et al., 2018). However, testing this hypothesis requires studying the relationship between hormone gene expression and gut pathogen abundance. Generally, it may be difficult to disentangle the effects of nano-La$_2$O$_3$ on gut bacterial communities independent of the direct effects on the host, which may, in turn, alter the tolerance to pathogens (Raymann et al., 2017). For example, nano-La$_2$O$_3$ exposure inhibited honeybee pollen intake and thus affected the resistance threshold of honeybees to pathogen stress (Di Pasquale et al., 2013). Although several studies have explored how contaminant-induced shifts in gut microbiota affect host physiology (Raymann et al., 2017; Motta et al., 2018), evidence of direct effects is currently lacking. In the future, it would be important to investigate how changes in honeybee health following environment disturbances may trigger changes to gut microbiota.

**CONCLUSION**

In summary, this study examined the effects of nano-La$_2$O$_3$ on honeybee health and gut bacterial communities. Our results provide evidence that nano-La$_2$O$_3$ exerted dose-dependent detrimental effects on honeybee physiology as reflected by the decrease in honeybee survival, pollen consumption, and body weight. Further, the exposures of 0 to 100 mg kg$^{-1}$ nano-La$_2$O$_3$ had no significant effects on gut bacterial community, while the exposure dose of 1,000 mg kg$^{-1}$ caused a significant community compositional shift. Besides, the specific genera within the community, including the pathogenic *Serratia* and *Frischella*, and the digestion-related bacteria *Bombella*, also responded significantly to nano-La$_2$O$_3$ exposure. Moreover, honeybee physiological impairments were significantly related to the enrichment of *Serratia* and *Frischella*. Collectively, these findings suggest that pathogen enrichment and gut dysbiosis may be at least partially responsible for adverse effects of nano-La$_2$O$_3$ exposure to honeybee health, thus extending our knowledge regarding the effects of nano-La$_2$O$_3$ on honeybee (*Apis mellifera*).

**DATA AVAILABILITY STATEMENT**

The 16S rRNA datasets generated in this study can be found in the SRA archive in GenBank under the BioProject PRJNA764692.

**ETHICS STATEMENT**

This study was reviewed and approved by the ethics committee of Institute of Apicultural Research, Chinese Academy of Agricultural Sciences (IAR, CAAS).

**AUTHOR CONTRIBUTIONS**

YG and Y-JL proposed the project and designed the experiments. Q-YD, ZJ, Y-YW, and Y-JL performed the exposure experiment and collected honeybee physiological data. X-TB, JW, and Y-JL collected the sequencing data. TX characterized the nanoparticles. X-TB and ZJ analyzed and visualized the data. X-TB and YG wrote the original draft. QZ, TX, BX, ZJ, and PH reviewed and edited the manuscript. All authors contributed to the data interpretation and paper writing.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.780943/full#supplementary-material

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