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Copper Oxide Nanomaterial Fate in Plant Tissue: Nanoscale Impacts on Reproductive Tissues

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Copper Oxide nanomaterial fate in plant tissue: Nanoscale impacts on reproductive tissues

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1 Copper Oxide nanomaterial fate in plant tissue: Nanoscale impacts on

2 reproductive tissues

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Abstract

- 21 A thorough understanding of the implications of chronic low dose exposure to Engineered
- Nanomaterials (ENMs) through the food chain is lacking. The present study aimed to characterize
- such response in *Cucurbita pepo* L. (zucchini) upon exposure to a potential nanoscale fertilizer:
- copper oxide (CuO) nanoparticles. Zucchini was grown in soil amended with nano-CuO, bulk CuO
- 25 (100 mg Kg⁻¹) and CuSO₄ (320 mg Kg⁻¹) from germination to flowering (60 days). Nano-CuO
- treatment had no impact on plant morphology or growth, nor pollen formation and viability. The
- 27 uptake of Cu was comparable in the plant tissues under all treatments. RNA-seq analyses on

vegetative and reproductive tissues highlighted common and nanoscale-specific component of the
response. Mitochondrial and chloroplast functions were uniquely modulated in response to
nanomaterial exposure as compared with conventional bulk and salt forms. X-ray Absorption
Spectroscopy (XAS) showed that Cu local structure changed upon nano-CuO internalization,
suggesting potential nanoparticles biotransformation within the plant tissues. These findings
demonstrate the physiological, cellular, and molecular consequences related to nano-CuO application
as a plant fertilizer, highlighting the differential mechanisms involved in the exposure to nano-CuO,
bulk or salt and the pathways of plant response to minimize environmental and health risk, through
sustainable nano-enabled agricultural strategies.

Keywords: nanomaterials, nanofertilization, RNA-seq, pollen, biotransformation, Cucurbita pepo.

Synopsis

Fertilization with nanoscale CuO affected zucchini at the physiological and molecular levels, from

roots to flowers, with significant internalization and particle biotransformation being evident.

Introduction

In recent years, interest in the utilization of nanotechnology to produce nano-enabled materials and delivery platforms to address the progressive inefficiency of mineral fertilization has been rapidly growing. 1.2 However, these nanomaterials are by their very nature more reactive and bioavailable than traditional forms utilized in agriculture, 1-3 and as such, have raised concerns over sustainability and safety with regard to human and environmental health. For example, direct utilization in agriculture raises the clear possibility of food chain contamination from staple plants to humans through the direct consumption of contaminated plant products as could happen with rice, maize and peanuts. 3.4 In addition, the potential widespread use of engineered nanomaterials (ENMs) as part of "nanoagricultural chemistry" has created concerns over damage to non-target organisms and to potential trophic transfer through terrestrial food chains, through vegetables grazing by simple insects and then spreading out to higher insects and predators. 5-9 As such, safety assessment and sustainability evaluation must be a core component in novel materials formulations for agri-food production purposes (e.g. nanopesticides, nanofertilizers). 1,10-13

A number of recent studies have demonstrated the unique potential of Cu-based nanomaterials in agriculture, ^{11,14,15} including field studies demonstrating how materials such as Cu₃(PO4)₂ based-nanosheets and commercial CuO NPs can be foliar applied to young seedlings so as to increase plant growth and suppress *Fusarium* spp. infections of tomato (*Solanum lycopersicum* L.) and watermelon (*Citrullus lanatus* L.) in full life cycle studies. Furthermore, Ma *et al.* (2020)¹³ studied how nanomaterial chemistry could be tuned to optimize the effects of pathogen suppression and nutrient release from Cu-based ENMs on sudden death syndrome (SDS) in soybean (*Glycine max* L.), developing a thermodynamic model to describe how morphology and matrix effects are implicated in Cu release and plant response. In addition, Cu-based nanoformulations are known to interact with organic acids in plant root exudates. These interactions significantly influence ENM stability, biotransformation and bioavailability, ¹⁶ as well as induce modifications in the plant metabolome. ¹⁷

The role of Cu nanomaterial bioavailability has also been investigated through trophic transfer experiments, including an assessment of how initial chemical form is impacted by relevant weathering conditions and subsequent material transformation.¹⁸ These findings highlight the importance of controlling ENMs physico-chemical properties (e.g. morphology, composition and dissolution) so as to develop safer and more sustainable nanoscale formulations for agriculture, simultaneously enhancing the targeting and delivery efficiency to optimize utilization of resources while minimizing negative impacts on the environment.¹

The current study investigated the potential effects of CuO NPs on zucchini (*Cucurbita pepo* L.) from a morphological, physiological, molecular, and atomic perspective, with a particular focus on gametogenesis and pollen development. Conventional CuO bulk material and CuSO₄ salts were used as controls to clarify nanoscale-specific effects of CuO NPs with regard to its fate within the plant tissues. Particular attention was focused on the function and regulation of chloroplast and mitochondrion activity, which play a critical role in pollen development. The coupling of a global transcriptomic approach (by RNA-seq) with synchrotron-based analyses, such as μ-X-ray Fluorescence (μ-XRF) mapping and Extended X-ray absorption fine structure (EXAFS) spectroscopy of Cu states, enabled a thorough understanding of the connection between the observed biological response and the physico-chemical condition of CuO NPs at a molecular and sub-molecular level in different plant tissues.

Methods

Nanomaterial characterization

Copper oxide nanoparticles (CuO NPs) (99% purity; 40 nm average sized) were purchased from U.S. Research Nanomaterials, Inc. (Houston, TX). Cu represents 79.8% of the total molecular weight of the molecule. CuO NPs were characterized by electron microscopy (TEM, Talos F200S G2, SEM FEG Thermo Fisher Scientific, Waltham, MA, USA) as reported in Figure S1. The average particle size (dh) and zeta (ζ) potential in ddH₂O were 533.9 \pm 47.2 nm and of -24.7 \pm 1.4 mV as

determined by Zetasizer Nano Series ZS90 (Malvern Instruments, Malvern, UK), after sonication by a Fisher Scientific Model 505 Sonic Dismembrator (Fisher Scientific, Waltham, MA) at 40% amplitude for 60s. CuO bulk material and CuSO₄·5H₂O were purchased from Sigma Aldrich (St. Louis, MO, US).

For particle dissolution analysis, CuO NPs and CuO bulk solutions (1000 mg L⁻¹) were prepared in ddH₂O, avoiding shaking and light, and portions were collected after 1, 2, 3, 7, and 14d. Aliquots of 1 ml for each sample were precipitated by ultracentrifugation at 30000 rpm, for 10 min, at 20°C (Optima Max-XP Ultracentrifuge, Beckman-Coulter Inc., Brea, CA, USA). The liquid phase was collected and digested in 4 mL of 1M HNO₃ (purity: 67% w/w) for 40 min at 200°C using a VELP DK20 digester (VELP Scientifica, Usmate, Italy). The digests were analysed by flame atomic absorption spectroscopy (FA-AAS; AA240FS, Agilent Technologies, Santa Clara, CA, USA) for the presence of Cu (lamp current: 4 mA; fuel: acetylene; support: air; wavelength 324.7 nm; slit with: 0.5 nm; linearity of calibration, R²: 0.9982), in three replicates. The average dissolution for CuO bulk and CuO NPs were between 0.1% and 0.15%, respectively, considering the theoretical value of 100% dissolution of ionic copper in CuSO₄.

Plant exposure

Cucurbita pepo L. (cv. Costata Romanesco) seeds were pre-germinated in vermiculite amended with Hoagland's Solution (10%) for 10 days prior to transplanting to soil. The Cu concentration administered in pre-germination was about 2 μg Kg⁻¹. Zucchini seeds were purchased from Johnny's Selected Seeds (Albion, ME, USA). The experimental soil was collected from the Connecticut Agricultural Experiment Station (CAES) Lockwood Farm in Hamden, CT, USA. Individual solutions of CuO NPs and CuO bulk material in water (30% water capacity of soil/vermiculite mixture) were probe sonicated by a Fisher Scientific Model 505 Sonic Dismembrator (Fisher Scientific, Waltham, MA) at 40% amplitude for 60–120s to maximize dispersion. Solutions

of CuO NPs, CuO (bulk) or (copper sulfate) CuSO₄ were slowly added to pots containing 500g of soil and manually mixed. The final concentration of NPs and bulk CuO in pots was 100 mg kg⁻¹ while for CuSO₄·5H₂O (copper sulfate pentahydrate), the amount was 320 mg kg⁻¹. Considering the molecular weight of the single molecules taken into account, this represented a total concentration of approximately 80 mg kg⁻¹ for all the treatments. Additional data related to the Cu release in treated soils are reported in Supporting Information (SI). The concentrations utilized were chosen to be below the limit considered as potential Cu contamination in soil, and yet still able to provide information on chronic plant exposure. Furthermore, the low dose utilized and the long growth period (60 days) are indicative of a chronic exposure scenario that is not common in the literature. Zucchini seedlings were planted (one each pot) and grown indoor under supplemental fluorescent lighting (60 μE m² sec) under a photoperiod of 16h light at approximately 22–28 °C until flowering. Plants were top watered during a 60-d growth period. For all the conditions, ten biological replicates were included.

Pollen morphology and pollen viability

Alexander's staining protocol was used to test pollen viability. Free anthers were collected when pollen was mature but anthers were still non-dehiscent (stage 12-13), and were fixed in Carnoy's fixative (6 ethanol: 3 chloroform: 1 acetic acid) for 2h. Mature pollen was collected and stained as described in Peterson *et al.* (2010).²² After staining, all aborted and non-aborted pollen grains were counted using a Zeiss Apotome 2 microscope at 20x magnification (Zeiss, Oberkochen, Germany). Pollen grains were analysed fresh with no fixation or staining; they were collected from mature flowers and positioned on 2 cm diameter stainless-steel sample holder (stub) covered with adhesive carbon tape. An environmental scanning electron microscope (ESEM) FEG2500 FEI (FEI Europe, Eindhoven, The Netherlands) operating in low-vacuum (60 Pa) with LFD (Large Field Detector) was used to enable optimal Secondary Electron (SE) imaging. The cone PLA (Pressure Limiting Aperture) of 500 µm improved the signal available to the Bruker X-ray detector,

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QUANTAX XFlash. SE imaging was performed at 10 KeV with a beam size of 2.5 μ m, EDX analysis at 20 KeV acceleration voltage, final lens aperture of 40 μ m, and beam size of 4 μ m. SE images and EDX spectra were collected from samples treated with CuO NPs, CuO bulk and untreated controls.

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Metal uptake measurement

Flowers harvested for elemental analyses were sampled and thoroughly rinsed with tap water, MilliQ water (resistivity: 18.2 M Ω cm) and 2% HNO₃ (0.01 M) to remove soil and surface-attached NPs. To determine Cu content in the tissues, fresh samples were dried at 100 °C for 72 h and digested in HNO₃ (purity: 67% w/w) at 115 °C, 25 min. After 30 min, 1 mL of H₂O₂ (purity: 30 % w/w) was added to each digestion tube and the samples treated for an additional 30 min prior dilution to 50 mL with ddH₂O. The digested samples from the 10 biological replicates per treatment were analysed by inductively coupled plasma mass spectrometry (ICP-MS) Agilent 7500ce (Agilent Technologies, Santa Clara, CA) for Cu presence (63 amu). The digests Cu content were quantified against a fivepoint calibration curve based on certified reference material (SPEX CertiPrep, Metuchen, NJ, USA), and that had been previously evaluated for linearity (R2: 0.9999) and accuracy. Analytical blanks, matrix blanks, and calibration verification samples were included in each sequence. Roots, stems, leaves and flower biomass samples were collected after 60-d for elemental analysis. Samples (10 replicates for each tissue) were digested as in the case of ICP-MS analyses and analysed by Atomic Absorption Spectroscopy (AAS) (AA240FS device, Agilent Technologies, Santa Clara, CA; lamp current: 4 mA; fuel: acetylene; support: air; wavelength 324.7 nm; slit with: 0.5 nm; linearity of calibration, R²: 0.9993). AAS analyses were conducted with a four-point calibration curve based on standard reference material (SPEX CertiPrep, Metuchen, NJ). Biomass and Cu content in all tissues were evaluated by a one-way ANOVA with a pairwise Tukey's multiple comparison test (IBM SPSS v. 26.0).

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RNA extraction and whole transcriptome analysis

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RNA samples were extracted from 0.1g (fresh weight) of pollen, leaves or roots samples from unamended control, CuO NP, CuO bulk and CuSO₄ treatments. Total RNA was extracted from 0.1 g of fresh plant material using a Sigma-Aldrich Spectrum Plant Total RNA Kit (Sigma-Aldrich, St. Louis, MO). Three biological replicates per treatment were used. Total RNA quality was assessed by gel electrophoresis and RNA quantity was determined using a Thermo Scientific Nanodrop Lite Spectrophotometer (Thermo Fisher Scientific, Wilmington, DE). Samples were sent to IGA Technologies Srl (Udine, IT) for RNA sequencing service. TruSeq Stranded mRNA kit (Illumina, San Diego, CA) was used for library preparation following the manufacturer's instructions. RNA samples were quantified and quality tested by Agilent 2100 Bioanalyzer RNA assay (Agilent Technologies, Santa Clara, CA). Final libraries were checked Agilent Bioanalyzer DNA assay (Agilent Technologies, Santa Clara, CA). Libraries were prepared for sequencing and sequenced on single-end 75 bp mode on NextSeq 500 (Illumina, San Diego, CA). Alignment of reads to the reference transcriptome available on Cucurbitgenomics database (http://cucurbitgenomics.org/)²³ was performed using STAR software with default parameters. The resulting raw data have been normalized and the differentially expressed genes were identified using a 2.3 threshold of FPKM data (in log2). Data have been deposited in the NCBI GEO database (accession number GSE173716). A student t test was applied for analysis of homogeneity of variance, statistical analysis for scatter plots, box and whiskers graphs. A principal component analysis (PCA) was performed with R statistical software (www.r-project.org). Venny bioinformatics tool (http://bioinfogp.cnb.csic.es/tools/venny/) was used for the generation of Venn diagrams. Gene Ontology (GO) analysis and A. thaliana ortholog gene identification was performed by the Cucurbitgenomics database. The GO term enrichment analysis was conducted using a cut-off p-value

of 0.05 for cellular components and biological processes, and 0.03 for relevant pathways,

respectively. Network analysis was performed using the GeneMANIA data service (http://www.genemania.org/) using *A. thaliana* orthologues genes.

Samples preparation for synchrotron-based analyses

μ-XRF and XANES analyses were performed at ELETTRA, Sincrotrone Trieste, in order to analyse Cu presence and distribution in tissues; root, leaf and flower samples (0.1 g, fresh weight) were cut and submerged in glutaraldehyde triphosphate in Eppendorf tubes for fixation. After three days the samples were dehydrated in gradients of alcohol (from 25 to 100%) and fixed with epoxy resin following Kurth et al. (2009).²⁴ To analyse potential variations in local structures within the tissues, bulk X-ray Absorption Spectroscopy (XAS) analyses at BM08 "LISA" beamline at ESRF have been performed. The protocols described in Marmiroli et al. (2020)²⁵ were applied. Briefly, samples were mixed with pure cellulose powder (Sigma Aldrich, St. Louis, MO, USA) and pressed into 1.3 cm diameter pellets using an amount of material sufficient to keep the total absorption (μ) ≤1.5 above the edge.

Low Energy μ-XRF (LE μ-XRF)

μ-XRF analyses in the soft X-ray regime were performed at the TwinMic beamline at ELETTRA, Sincrotrone Trieste, Italy.²⁶ For the present experiment, the TwinMic microscope was operated in scanning transmission mode (SXM), the beam was focused on the sample through a zone plate (600 μm in diameter with a 50 nm outermost zone width), and a micrometric or sub-micrometric probe size was delivered. While the sample was raster-scanned perpendicularly to the incoming monochromatic beam, a fast readout CCD camera collected the transmitted X-rays and an 8-silicon drift detector-based XRF system acquired the emitted fluorescence photons.²⁷ The obtained absorption and phase contrast images outline the morphological features of the sample at sub-

micrometer length scales, whereas the simultaneous detection of the low energy μ-XRF correlates the elemental distribution to the morphology. The elemental distribution was then obtained by deconvolving and fitting the XRF spectra with PyMCA software.²⁸ A photon energy of 1.26 keV was used to excite and obtain optimal emission conditions for the elements of major interest (Cu, Na, Ni and Fe) with a spot size of 1.45 μm and a dwell time of 8 s per pixel for XRF mapping and a CCD dwell time of 50 ms per SXM imaging. Each map lasted approximately 5-7 h, depending on the dimensions of the scanned area.

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XRF and X-ray absorption near edge structure (XANES) mapping

Zucchini root and flower thin section samples were also investigated by means of XRF mapping and XAS at the XRF beamline, ELETTRA Sincrotrone Trieste (Italy)²⁹ which covers a different energy range compared to TwinMic and allows detection of heavier elements and K lines of transition metals. The experiment was conducted using a Si(111) monochromator and standard 45°/45° geometry for fluorescence mode measurements, using an XFlash 5030 SDD (Bruker, Berlin, Germany). Higher order harmonics contamination was suppressed by a pair of parallel plane mirrors intercepting the beam in grazing incidence. Thin sections of samples embedded in resin were sealed between two Mylar foils and fixed on the Al sample holder using a Delrin interlocking ring. This design was necessary to secure the samples and to have a system compatible with the working conditions of the Ultra High Vacuum Chamber (UHVC, 10⁻⁷ mbar) available at the XRF beamline. XRF maps were collected with an incident beam energy of 10 keV and a beam size at the exit slits of 200x100 µm² (HxV) to inspect the elemental distribution. Based on the XRF maps acquired on the samples (data not shown), the areas with higher content of Fe, Cu and Zn were selected to collect XANES spectra at the relative K-edges. The Si(111) monochromator was calibrated before the measurements using reference metal foils. All spectra were collected using 5 seconds per step and a variable energy step as a function of the energy: Large step (5 eV) in the first 200 eV of the spectrum,

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smaller step (0.2 eV) in the near-edge region and a k-constant step of 0.07 Å⁻¹ further above the absorption edge. Multiple spectra were collected and merged in order to increase the signal to noise ratio. The oxidation state was determined using least-squares Linear Combination Fitting (LCF) based on reference spectra collected on compounds of known oxidation state. Background removal, normalization of XANES spectra and LCF analyses were performed using the ATHENA software package.³⁰

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Extended X-ray absorption fine structure (EXAFS)

Extended X-ray absorption fine structure (EXAFS) measurements at the Cu K-edge (8978.9 eV) were performed at the LISA CRG beamline (BM08)³¹ at the European Synchrotron Radiation Facility (ESRF, Grenoble, France) using plant samples and three model compounds: CuO (bulk), CuO NPs and CuSO₄·5H₂O. The main optical features of the beamline were a fixed exit monochromator with a pair of Si(111) crystals (energy resolution $\Delta E/E \approx 1.33 \cdot 10^{-4}$); Si mirrors were used for harmonics rejection (E cutoff ≈ 15 KeV). Energy was calibrated with a Cu reference foil (8978.9 eV). Spectra of plant samples were acquired at 80 K, in order to minimize beam-induced damage, with a constant k step of 0.05 Å⁻¹ up to a maximum k value of 12.5 Å⁻¹; model compounds were measured at room temperature with a k step of 0.03 Å⁻¹ up to k=18 Å⁻¹. Plant samples were measured in fluorescence mode with a 12-element HP-Ge detector, 32 while model compounds were measured in transmission mode. Multiple spectra were collected and merged in order to increase the signal to noise ratio. ATHENA software³⁰ was used to calibrate the energy and to average multiple spectra. Standard procedures were followed to extract the structural extended EXAFS signals $(k \cdot \chi(k))$, including pre-edge background removal, spline modelling of bare atomic background, edge step normalization, and energy calibration.³³ Model atomic clusters centered on the absorber atom were obtained by ATOMS;³⁴ theoretical amplitude and phase functions were generated using the FEFF8 code.³⁵ EXAFS spectra were fitted through the ARTEMIS software in the Fourier-Transform (FT) space.³⁰

Results and Discussion

Pollen morphology and viability

Pollen grain morphology was analysed by ESEM of transverse sections of developing mature anthers; no overt differences were observed across treatments (Figure S2a). Pollen viability was also evaluated to determine the male gametophyte developmental stage and the preservation of plant reproductive fitness. Similar to morphology, there were no differences across CuO NP, CuO bulk and CuSO₄ treatments as compared to the untreated control, with a pollen viability approximately 100% in all cases (Figure S2b).

Previous studies have demonstrated that copper can be toxic to seed and pollen germination, pollen viability and pollen tube growth; Sharafi (2014)³⁶ showed that high concentrations (250 mg Kg⁻¹) of copper cause an almost complete inhibition of pollen germination and pollen tube lengthening in almond (*Prunus dulcis* (Mill.) D.A. Webb cultivars). Similar results were observed in *Pisum sativum*, where copper (35-700 mg Kg⁻¹) was highly toxic to pollen germination.³⁷ It is unclear if zucchini exhibits a unique tolerance to copper; importantly, few studies have investigated the potential effects of copper nanomaterials on pollen formation and maturation. Kumbhakar *et al.* (2016)³⁸ showed that both copper and cadmium-based NPs reduced pollen fertility in black cumin (*Nigella sativa* L.), both during pollen formation and in developmental maturation process. Similarly, in *Coriandrum sativum* L. CdS NPs and CuO NPs induced physiological alterations and cytological aberrations in meiotic cells, and decreased viability of pollen.³⁹ The alteration types and frequencies in meiotic cells of *C. sativum* following NPs treatments (0.25-1 mg L⁻¹) were less severe than those reported in *Nigella sativa*.³⁸ Notably, in the present study the Cu concentration used was specifically selected to be below the limit considered for Cu contamination in soil.¹⁹⁻²⁰

Plant biomass and metal content

After flowering, plants were harvested, and the fresh mass of roots and shoots was measured (Table S1-S2). Treatment with CuO, CuO NPs and CuSO₄ had no impact on zucchini biomass (fresh weight) compared to untreated control. These results align with much of the present literature, showing that exposure to CuO NPs did not negatively impact the biological parameters in agricultural crops. Tamez et al. (2019)⁴⁰ reported no significant changes in zucchini root and leaf biomass upon exposure to comparable concentrations of CuO NPs. Pagano et al. (2016)²¹ demonstrated that CuO NPs had no effect on *C. pepo* biomass at a higher concentration (500 mg Kg⁻¹) and with an experimental design that provided greater direct interaction between NPs and tissues (vermiculite growth media). Alternatively, studies conducted with the model plant *Arabidopsis thaliana* grown in hydroponic conditions showed a strong reduction in root length after exposure to CuO NPs (10-20 mg L⁻¹).⁴¹ These contrasting results demonstrate the importance of CuO NPs dose to biological response, and also highlight the influence of growth medium, plant species, and the exposure time to observed effects.

The Cu content in soil and in the different tissues of zucchini plants was determined by Atomic Absorption Spectroscopy (AAS), as shown in Table S3. The Cu content in soil was determined (14.18 mg Kg⁻¹) in order to justify the concentrations utilized for the experiment, highlighting a different metal release percentage within the soil, starting from the same relative concentration of Cu potentially available of 80 mg Kg⁻¹ (salt > NP > bulk). Although there was a trend for increased Cu content of tissues with all Cu treatments, only plant roots from the CuSO₄ exposure were significantly increased. To validate the AAS results on flowers, analysis of the Cu content was performed also by Inductively Coupled Plasma Mass Spectrometry (ICP-MS). Here, results show that the Cu content from the CuO NPs and bulk material treatment was 43 and 30% (significant at p<0.05) greater than the untreated control, respectively (Table S4). This finding demonstrates that CuO NPs addition to the soil does result in Cu accumulation in the reproductive tissues, although there is no difference

based on particle size. However, these results were impacted by instrument limits of detection (ICP-MS: 0.04 µg L⁻¹ vs AAS: 0.05 mg L⁻¹) and quantitation (ICP-MS: 0.1 µg mL⁻¹ vs AAS: 0.1 mg L⁻¹). A previous study from our group focused on zucchini exposed to nanoscale Cu evaluated a broad set of physiological assays, including chlorophyll content, mitochondrial functionality, and metal content in different plant tissues (roots, stems, leaves), and also demonstrated significant Cu translocation from roots to both stems and leaves.²¹

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RNA-seq data analysis: critical aspects related to the CuO NP molecular response

Given the evidence of an active translocation of Cu into reproductive tissues, the plant transcriptomic response of different tissues and organs to exposure was evaluated using the highquality assembly of the C. pepo genome (NCBI BioProject PRJNA386743, sequences length 263 Mbps; 34240 ORFs) published on Cucurbitgenomics database.²³ Statistical analysis of RNA-seq datasets showed high homogeneity between treatments in the different tissues, with comparable averages and dispersions (Figures S3-S6). After normalization to the untreated control, comparison across CuO NPs, CuO bulk and CuSO₄ exposure in roots showed 4420, 6540, and 4747 differentially expressed genes in the three treatments, respectively. In leaves, the CuO NPs treatment showed a lower number of differentially expressed genes compared to the other treatments: 3122 genes were up- or down-regulated with CuO exposure, whereas the values for CuO bulk and CuSO₄ were 9924 and 9103, respectively. The number of differentially expressed genes in CuO NPs exposed pollen was also markedly lower in comparison to the treatments in the other tissues: 1829, 2112 and 2163 respectively for CuO NPs, CuO bulk and CuSO₄ treatments. This large quantitative difference in gene expression was certainly related with a larger number of different biological processes performed in roots and leaves, as compared to pollen, but could be also due to the lower amount of the Cu (in different forms) translocated to pollen.

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Venn diagrams of up- and down-regulated genes in roots (Figure 1) and the relative GO enrichment (Supplementary Information, SI2) data show that the transcripts in common among all the tested conditions were only 4.3% and 16.3% of total genes up- and down-regulated, respectively. In the roots, the specific molecular responses to the three different treatments were largely independent of each other, as shown by the low percentage of gene functions common among all the three conditions tested and by the two-by-two common classes. Metabolic processes and ribosome translation were the most highly represented groups in biological processes related to CuO NPs and CuO bulk (Figure 2; details in Supplementary Information, SI2), together with mitochondrial activity. In the CuSO₄ treatment, unlike the NPs and bulk exposure, a nuclear component was represented, and this can be related to greater Cu ion toxicity. 42 The percentage of genes commonly up- or downregulated in leaves was similar to that observed in roots: 4.7% and 16.4% total shared genes, respectively (Figure 1). The percentage of genes in common between CuO bulk and CuSO₄ increased dramatically, both for up- and down-regulated transcripts, to 40.1% and 46.6%, respectively. This observation may correlate with the Cu ion release from the CuO bulk material within the plant tissues, which seems to be higher than for CuO NPs, in spite of the similar dissolution rate in ddH₂O. Genes involved in metabolic and energetic processes are among the more enriched genes; in addition, GO terms related to chloroplast genes are well represented, as are genes for abiotic stimuli response (Figure 2; details in Supplementary Information, SI3). Previous studies with A. thaliana highlighted the role of chloroplast as a potential target of ENMs exposure. 43 Wang et al. (2016)44 showed that CuO NPs block electron transport between the two photosystems which can cause an excessive ROS accumulation and oxidative stress, damaging biological molecules and disrupting of cellular metabolism. Furthermore, CuO NPs strongly up-regulate ZAT12, a transcription factor implicated in abiotic stress response, with a key role in ROS signalling pathway and co-expressed with ORF31, a chloroplastic electron carrier involved in photosynthesis that has been identified as a potential biomarker of ENM exposure. 21 In pollen, the percentage of genes up- or down-regulated common to all treatments increased as compared to the leaves and roots: 25.5% and 33%, respectively (Figure

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1). The percentage of genes up- and down-regulated specifically related to CuO NPs response is significant (21.1% and 12.5%), when compared to the other two treatments. This data strengthens the idea that CuO NPs were not only translocated (intact or modified) into the floral parts of the plant, but once there, they trigger a "nanoscale-specific" response which is different from the response observed in roots and leaves. These results likely reflect a multifaceted response, including partial dissolution of CuO NPs and CuO bulk giving rise to a "non-specific Cu response", along with a non-dissolved component exerting a nanoscale-specific response. It is also reasonable to suppose that amount of Cu²⁺ derived from the three treatments increased as a consequence of the interaction with plant organs and tissues.

Pollen has a significantly lower number of expressed genes as compared to the vegetative tissues, but data highlight some pollen-specific functions and other components which have been described as unique to sporophyte tissues.⁴⁵ The difference between the leaves was related to the low level of expression of genes involved in energy metabolism, especially photosynthesis, since pollen is not photosynthetically active. Another difference between treated and untreated plants was in pollen with higher expression level of genes with functions in ion transport, cell-wall and starch metabolism, and cytoskeleton dynamics (Figure 2; details in Supplementary Information, SI4). Previous studies showed that polarized internal gradients and/or external fluxes of protons, potassium, and chloride had a role in pollen tube function, 46 but that ion channel and transporter involvement in ion fluxes across the plasma membrane in pollen tubes is still largely unknown. Starch biosynthesis during the final phases of pollen maturation is fundamental because starch is a reserve source of energy for pollen survival and it may also act as a metabolic checkpoint for pollen maturity. This pathway is prematurely aborted whenever starch levels remain below a critical amount, strongly linking pollen viability to starch deficiency.⁴⁷ A key aspect of pollen tube tip growth is the constant construction of new cell wall and plasma membrane at the tube apex. Vesicles delivering this material are mediated by the actin cytoskeleton.⁴⁸

The whole transcriptome analysis of *C. pepo* treated with CuO NPs showed interesting insights from a functional point of view. Chloroplast and mitochondrial function were critical in regulating the response to CuO NPs and the energy metabolism in all plant tissues, which becomes primarily mitochondrial functionality in pollen formation and development (Figure 2; details in Supplementary Information, SI5). A network analysis produced for chloroplast genes in leaves, and for mitochondrial genes in roots, leaves and pollen, respectively, shows the physical interactions between the reported gene targets (Figures S7-S10). Genes highlighted in heatmaps and Venn diagrams (Figure 3) showed a certain specificity to CuO NPs response, in particular in roots and leaves. In the case of pollen, the percentage of common regulated genes among CuO NPs, CuO bulk and CuSO₄ treatment is increased (Figure 3), suggesting that during translocation from roots to shoots there was an increase in ionic Cu presence. Additional information about potential sensitive targets in pollen development, derived from the study of orthologue genes in the yeast *S. cerevisiae*, were investigated, highlighting a certain level of commonality in the response with the RNA-seq analyses. Results are reported and described in Supplementary information (Table S5, and Figure S11).

CuO NPs biotransformation

μ-XRF analyses performed at the TwinMic beamline, ²⁶ an example of which is depicted in Figure 4, showed that in the root sections, Cu in general was mainly detectable on cell walls and more visible in the treatments with CuSO₄, followed by nanoparticle and bulk forms where Cu content was very close to TwinMic detection limits; the higher presence of Cu in the treatments with CuSO₄ salt is due to the salt dissociation in the soil and followed by ready Cu accumulation in the roots. Notably, the roots were thoroughly washed before the resin embedding procedures to avoid external contamination. Fe was highly present in all root samples, including the controls, likely because it was abundant in the soil. The roots maps for Cu (Figure 4a) are consistent with those obtained by Servin *et al.* (2017).¹⁸ In the flower samples (Figure 4b), it is possible to observe the pollen sacs and the

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completely developed pollen grain; one exception is for the CuSO₄ treatment, where the pollen sacs were noticeably smaller and possessed fewer pollen grains. Interestingly, Cu was present in the roots, in particular in the cell wall, where other elements such as Ca, which is an important cofactor in building of the cell wall, are known to be present. 49 In CuO bulk treatment Cu was almost not detectable in roots. Although the resolution of the maps does not allow nanoparticle visualization, the EXAFS analyses (Table S6, Figure 5) confirm that CuO NPs in the plants were biotransformed. This suggests that cellular and molecular activities remodel and biotransform the nanoparticles. Cu was present in the flowers treated with all the three types of Cu-based materials, but there were minimal differences in the signal intensity and in the localization of the element. The treatment with CuO NPs did not hinder formation of the flower or pollen and did not result in overt phytotoxicity, but there were nanoscale-specific molecular effects at the transcriptomic levels as described by RNA-seq analysis. The same was true for the bulk Cu treatment, although treatment with CuSO₄ did appear to negatively affect gamete formation. The idea of a biotransformation of CuO NPs once within the plant tissues has been reported in the literature; Servin et al. (2017) reported that after treatment, transformed CuO NPs products were detected in roots as Cu₂O, Cu₂S and Cu-acetate. ¹⁸ These biotransformation processes significantly influence NPs bioavailability and effects in plants, including broad main metabolic and physiological processes, as well as gametogenesis. 18,50

Figure 5 shows normalized XANES and EXAFS spectra of the plant samples, together with those of the model compounds CuO, CuO NPs and CuSO₄·5H₂O and the EXAFS multiparameter fits. Both XANES (results shown in Figure S12) and EXAFS features show that the CuO NPs structure is closely related to the CuO bulk structure. EXAFS multiparameter fits (Table S6) were performed on both CuO samples based on the tenorite structure,⁵¹ yielding the same results in terms of interatomic distances and path degeneracies, with both refined parameters closely agreeing with the theoretical ones. EXAFS quantitative results on plant samples (Figure 5, Table S6) clearly indicate that the CuO structure is not fully preserved within the plants tissues, in particular in the roots, and that after uptake, the particles are biotransformed over time, leading to Cu²⁺ release starting in roots and increasing up

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to the flower. Specifically, the prominent signal at R > 4.5 Å in the Fourier Transform (FT) spectra in both CuO bulk and NPs is not visible any longer in plant samples. Moreover, the peaks at R< 4.5 Å are markedly weakened. In addition, no overt differences are observable among plant samples treated with two different CuO types. First, shell distances in both roots and flowers are typical of Cu²⁺ in square planar coordination with O and thus compatible with a remnant structure of CuO. However, the Cu local environment in the higher shells shows small differences between roots and flowers. Indeed, a second Cu-O shell path is systematically present in both flowers and roots at significantly different distances: ≈2.7 Å in roots and 2.8 Å in flowers. Moreover, a Cu-Cu path at ≈3.55 Å is always present in roots while the same path could not be fitted in flowers, the only exception being plants treated with CuSO₄, where the path distance is much longer (3.79 Å). In shoots of the Cu hyperaccumulator plant Crassula helmsii, Kupper at al. (2009)⁵² reported Cu²⁺ O ligands at 2.001 Å, indicating Cu bonds with small organic acids. Mijovilovich et al. (2009)⁵³ studied the leaves of Noccea caerulescences (ecotype Ganges), an hyperaccumulator plant of Cd and Zn but sensitive to Cu, reported ligands for Cu²⁺ at 1.9 Å sulfur atoms, indicating ligands with S rich molecules, and at 4.5 Å, a double ligand Cu-Cu that they attributed to Cu biomineralization. These findings do not align with our results and this is likely due to plant species differences; hyperaccumulator plants having a unique and specific metabolic profile that is different from that of non-hyperaccumulator species such as Cucurbita pepo.

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Environmental implications

Given the essential role of Cu to the plant life cycle and its biotic response to disease, there has been significant interest in its use as a potential nanofertilizer. However, in certain plants and under certain concentrations, phytotoxicity has been observed. In the present study, the effect of three types of Cu (CuO NPs, CuO bulk, and CuSO₄) was compared in *C. pepo* using a broad range of physiological and molecular endpoints, with a focus on the process of male gametogenesis and pollen

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production, which are essential to reproduction and from fruit formation and ultimately, to plant yield. In a dioic species such as zucchini, gamete fertilization depends primarily on pollen quality and vitality of the parental plant, which then mediates fruit production. From the morphological and physiological perspective, as inferred by ESEM and XRF analyses, there were few differences between the three forms of Cu (CuO NPs, CuO bulk and CuSO₄) in the roots, but as the Cu was translocated to the flower, the CuSO₄ treatment exerted a more marked negative effect on pollen viability. This increasing toxic response likely was a function of the complete dissolution to Cu²⁺ ions in this medium and the increased reactivity of Cu in this form. Conversely, the NP form exerted almost no effects on pollen and exhibited a reduced stimulation of Cu uptake, possibly being a function of Cu complexed to organic ligands within the plant tissues that mitigated chemical dissolution. The CuO bulk material results for the molecular and physiological endpoints were more similar to CuSO₄, in spite of the CuO bulk and CuO NPs dissolution behavior in ddH₂O being quite similar. Interesting, some nanoscale materials release ions at a greater rate than bulk materials, due to increased surface area and volume. 14 However, coatings, complexation and corona formation could modulate dissolution. The transcriptomic analysis of the different tissues and flowers showed that metabolic processes and ribosome translation were highly represented among the most responsive pathways. Chloroplast and especially mitochondrial functions were particularly affected in response to CuO NPs, which agrees with previous data and aligns with the organelle's role in energy metabolism in all plant tissues, 44 and specifically in pollen formation and development. In addition, the EXAFS features demonstrate the occurrence of CuO NPs biotransformation, highlighting a similar Cu local environment from roots to flowers. The similarity of the Cu environment after different treatments seemed to depend more on the plant tissue than on the type of treatment, suggesting that the biotransformed Cu environment is reached after substantial dissolution of Cu ions. followed by stabilization of Cu in complexes whose nature is more dependent on the plant characteristics than on the type of treatment. Indeed, the transcriptomic data showed that at the molecular level, the response was partially nanoscale-specific, including in the pollen. Similar

phenomena have been reported for other nanomaterials such as CeO₂ NPs and CdS QDs.^{25,54} The evidence for the formation of Cu ions when CuO NPs are accumulated by the plants and the fact that still there is a certain level of nanoscale-specific response suggests that *in planta* biotransformation processes are significant and critical to overall plant response. Overall, this suggests nanoscale CuO NPs as nanofertilizers likely presents minimal concerns to general plant health. A thorough and mechanistic understanding of these processes such as that provided by this study will be necessary to ensure the safe and sustainable application of Cu-based and other nanoscale materials in nano-enabled agricultural strategies.

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Authors contribution

MM, LP, NM and JCW coordinated the study and designed the experiments. RR, LP and RD performed individual experiments and analysed the physiological and molecular data in with collaboration of MM and RRu. AG, GA, performed the synchrotron analyses in Trieste with collaboration of SP, GG and VB. AP, FA and GOL (remotely) performed the XAS measurements at

531	LISA BM08 beamline at ESRF; GOL performed EXAFS data analysis in collaboration with FA. All
532	authors contributed to manuscript revision and approved the final version.
533	
534	Conflict of interest
535	The authors declare no conflict of interest.
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538	Supplementary Information (SI) description:
539	Supplementary information included (SI1):
540	Method section and results for qPCR of genes involved in gametogenesis
541	Results of LCF analysis of XANES spectra
542	Figure S1: CuO nanoparticles visualization by TEM
543	Figure S2: ESEM micrographs pollen grains and pollen viability assay
544	Figure S3: Statistics of the genes datasets from roots samples
545	Figure S4: Statistics of the genes datasets from leaves samples
546	Figure S5: Statistics of the genes datasets from pollen samples
547	Figure S6: PCA of all data profiles
548	Figure S7: Gene network of chloroplast targets observed in leaves treated with CuO NPs
549	Figure S8: Gene network of mitochondrial targets observed in roots treated with CuO NPs
550	Figure S9: Gene network of mitochondrial targets observed in leaves treated with CuO NPs
551	Figure S10: Gene network of mitochondrial targets observed in pollen treated with CuO NPs
552	Figure S11: Heatmap transciprotmics genes involved in meiosis and gametogenesis
553	Figure S12: XANES fits and relative K-edge data
554	Table S1: Biomass of roots and shoots
555	Table S2: Flower biomass
556	Table S3: Copper concentration measured in soil, roots, shoots and flowers by AAS

- Table S4: Copper concentration measured in flowers by ICP-MS
- Table S5: Genes' information and primer sequences utilized in Real time PCR assay
- 559 Table S6: EXAFS multiparameter fit details for studied samples and
- 560 reference compounds

- *Supplementary information reported in excel format:*
- Supplementary Information 2 (SI2): GO analysis of up- and down-regulated genes exposed to CuO
- NPs, CuO bulk and CuSO₄ in roots.
- Supplementary Information 3 (SI3): GO analysis of up- and down-regulated genes exposed to CuO
- NPs, CuO bulk and CuSO₄ in leaves.
- Supplementary Information 4 (SI4): GO analysis of up- and down-regulated genes exposed to CuO
- NPs, CuO bulk and CuSO₄ in pollen.
- Supplementary Information 5 (SI5): A. thaliana ortholog genes analysis of relevant chloroplast and
- mitochondrial targets isolated from *C. pepo* exposed to CuO NPs, CuO bulk and CuSO₄ in roots,
- leaves and pollen.

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Figure captions:

Figure 1. Comparison of high-throughput transcriptional datasets related to the molecular response of *C. pepo* in condition of treatment with CuO NPs, CuO bulk and CuSO₄, in the roots (a), leaves (b), pollen (c), represented with Venn diagrams. Up-regulated and down-regulated genes are reported on left and right side, respectively. Percentage of identity between CuO NPs, CuO bulk and CuSO₄ is also reported. Data were normalized on the untreated controls, with a 2.3 threshold of raw data (in log2). Data highlighted an increase in the percentage of common genes involved in the response to the three different Cu-based forms from roots to pollen, suggesting an increased bioavailability of free Cu in the plant shoots.

Figure 2. GO biological processes expressed in percentage (%) of gene cluster enriched, related to roots (a), leaves (b), and pollen (c) related to the treatment with 100 mg kg⁻¹ of CuO NPs. Upregulated and down-regulated genes are reported as blue and orange bars, respectively. Additional details related to GO analyses in the different tissues are available in Supplementary Information, SI2-SI4.

Figure 3. Heatmaps and Venn diagrams comparison of the genes involved in chloroplast functions in response to CuO NPs, CuO bulk and CuSO₄, in leaves (a), and in mitochondrial functions identified in roots (b), leaves (c) and pollen (d) tissues. Data related to the specific genes are reported in Supplementary Information, SI5. Data confirmed the increase in percentage of common modulated genes from roots to pollen in response to the three different Cu-based forms utilized for the treatments.

Figure 4. μ-XRF maps of (a) roots and (b) flowers from plants treated with CuO NPs. Names of the mapped elements are on top of each figure. The maps are related to the black and white square on top left (Abs) which is the 20x image of the cells in the root tissue and pollen sac tissues treated with CuO NPs. Cu map is always the last in the second row for (a) and (b).

Figure 5. XANES spectra of the measured samples and model compounds (a). Cu K-edge k²-weighted EXAFS region (b) and Fourier transforms (c) of plant tissues and model compounds. Solid lines are data, red lines are fits. Energy was calibrated with a Cu reference foil (8978.9 eV). In order to minimize beam-induced damage, spectra of samples were acquired at 80 K with a constant k step of 0.05 Å-1 up to a maximum k value of 12.5 Å-1 for plant tissues while model compounds were measured at room temperature with a k step of 0.03 Å-1 up to k=18 Å-1. Plant samples were measured in the fluorescence mode with a 12-element HP-Ge detector.

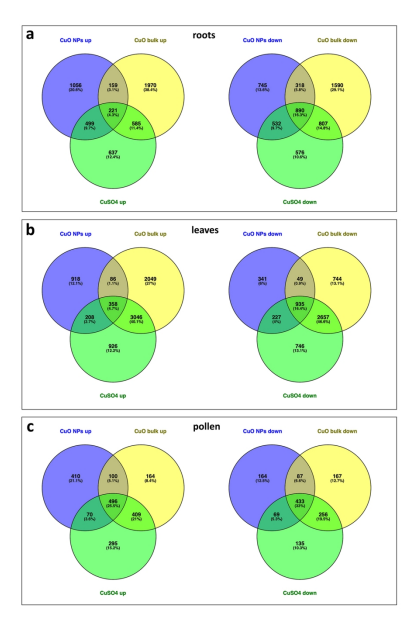


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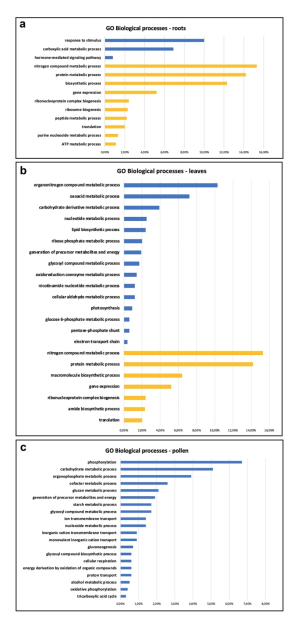


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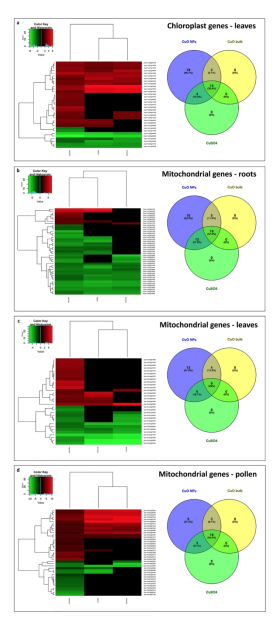


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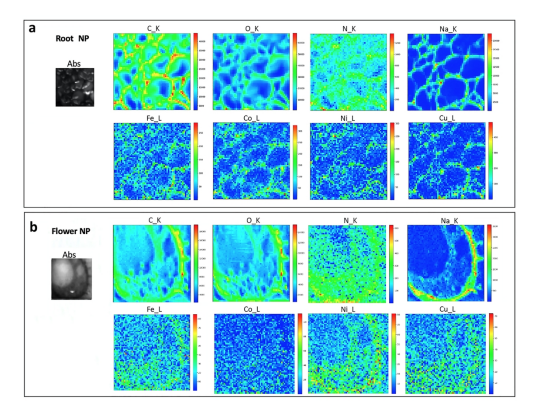


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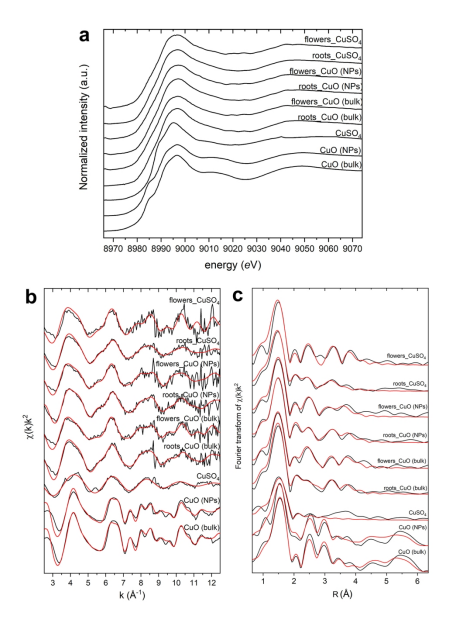


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