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Predicting the phytotoxic mechanism of action of LiCoO₂ nanomaterials using a novel multiplexed algal cytological imaging (MACI) assay and machine learning†

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Currently, there is a lack of knowledge of how complex metal oxide nanomaterials, like LiCoO₂ (LCO) nanosheets, interact with eukaryotic green algae. Previously, LCO was reported to cause a number of physiological impacts to *Raphidocelis subcapitata* including endpoints related to growth, reproduction, pigment & lipid biosynthesis, and carbon biomass assimilation. Furthermore, LCO was proven to physically enter the cells, thus indicating the possibility for it to directly interact with key subcellular components. However, the mechanisms through which LCO interacts with these key subcellular components is still unknown. This study assesses the interactions of LCO at the biointerface of *R. subcapitata* using a novel multiplexed algal cytological imaging (MACI) assay and machine learning in order to predict its phytotoxic mechanism of action (MoA). Algal cells were exposed to varying concentrations of LCO, and their phenotypic profiles were compared to that of cells treated with reference chemicals which had already established MoAs. Hierarchical clustering and machine learning analyses indicated photosynthetic electron transport to be the most probable phytotoxic MoA of LCO. Additionally, single-cell chlorophyll fluorescence results demonstrated an increase in irreversibly oxidized photosystem II proteins. Lastly, LCO-treated cells were observed to have less nuclei/cell and less DNA content/nucleus when compared to non-treated cell controls.

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Environmental significance

Complex metal oxide nanomaterials, like LiCoO₂, are among some of the most widely produced nanomaterials in commerce, yet there is still little information on how they may interact with algae and other plant-type organisms. Furthermore, in general, there is a need for more non-targeted, high-throughput profiling assays that can quickly and effectively characterize nanomaterial mechanisms of action in environmentally relevant organisms, like algae. This study applies a novel phenotypic profiling approach for predicting the mechanisms through which nanomaterials, like LiCoO₂, interact at the biointerface of plant-type organisms in a way that is quick, efficient, and cost-effective. Using this approach, it was found that the most probable mechanism of action of LiCoO₂ in algae is that of photosynthetic electron transport inhibition.

Introduction

As the number of engineered nanomaterials found in the environment and commerce expands, understanding the breadth of their environmental consequences is a challenge we currently face. In particular, a class of nanomaterials we

need more toxicological data on are complex metal oxide nanomaterials, like LiCoO₂ (LCO) nanosheets due to their high volume of production and presence in commerce. LCO is one of the most commonly used cathode materials in rechargeable Li-ion batteries and can be found in a multitude of consumer electronics from computers and smart phones to high-end electric vehicles. As such, the annual production rate for LCO has skyrocketed to levels of environmental significance in recent years. What makes this matter particularly concerning, however, is that there is little to no infrastructure for recycling or for properly disposing of LIB's, nor is there any economic incentive to do so as it is cheaper to simply manufacture new battery materials. For

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example, it is currently estimated that less than 5% of all LIB's are being recycled, with the rest ending up in landfills or being disposed of in other mean of un-sustainable storage.3 As a result, this means that there is a high probability for LCO to be unintendedly released into the environment, thus posing an ecological risk. And what exacerbates this risk is that LCO contains high-valence metals with unique catalytic properties, high reactivity, and known inherent toxicity. 4,5 Thus, understanding the impacts of LCO to a wide diversity of environmentally relevant organisms across multiple trophic levels is of paramount importance. Current research has been done to assess LCO's toxicological impacts and mechanism of action (MoA) in higher animaltype organisms. For example, Curtis et al., 2022⁶ has reported LCO to cause a differential expression in electron transport and energy metabolism related genes across different fish and invertebrate species. Furthermore, Niemuth et al., 20207 has reported LCO to participate in redox reactions and alter the redox state of and Fe-S proteins, which are important for electron transport. However, our understanding of LCO's impacts to plant-type organisms is still limited.

In a previous study, LCO was reported to cause a number of physiological impacts to Raphidocelis subcapitata, such as reduced growth, altered pigment biosynthesis, and impaired photosynthetic productivity.8 Furthermore, LCO nanoparticles were proven to physically enter the algal cells, thus implying that they undergo direct interactions with key subcellular compartments.8 However, our understanding of the mechanisms governing these consequences is still unclear. Given that photosynthesis was impacted in algae and its reverse reaction, respiration, has also been reportedly impacted in higher animal species,6 it's likely that they may experience similar mechanistic disturbances. For example, key photosynthetic proteins like photosystem II (PSII), which facilitate electron flow in chloroplasts, could be impacted.

In order to assess the interactions of LCO at the biointerface of algal cells, a multiplexed algal cytological imaging (MACI) assay in combination with data mining and machine learning techniques can be used. MACI, as described in a previous study by Ostovich & Klaper, 2023,9 is a type of high-throughput phenotypic profiling assay which involves the use of fluorescence cytochemistry to visualize multiple subcellular structures within the cell, and automated high-content fluorescence imaging to take hundreds of snapshots of their morphology in a consistent manner. In general, high-content phenotypic profiling works off the principal that the morphology of a cell is very sensitive to environmental cues¹⁰ and that subtle, yet reproducible, alterations to subsets of morphological features can be used as a framework to characterize compound-specific interactions and predict MoAs of novel compounds when comparing their phenotypic profiles to that of compounds with established MoAs. 11 Additionally, image-based profiling assays provide the advantage of speed and cost effectiveness when compared to transcriptomic- and proteomic-based profiling assays. 12,13 However, MACI, in the

context of eco- and phytotoxicological research, provides an additional benefit compared to other mainstream highthroughput phenotypic profiling assays, like Cell Painting, 14 in that it relies on the fluorescent labeling of subcellular structures that are unique to the architecture of algal cells, as opposed to human/animal cells. This is beneficial in terms of phytotoxicological research as it would allow researchers to accurately characterize the MoAs that are unique to plant-type organisms, especially MoAs that target the chloroplast (not present in animal cells). Additionally, MACI could be particularly useful in terms of environmental relevance as planttype organisms, like microalgae, on average, constitute the largest amount of biomass in the environment and drive environmental processes that affect ecosystem dynamics. 15,16 Therefore data obtained from MACI could potentially be used to make predictions that are tied to larger ecosystem level consequences.

In this study, algal cells were exposed to LCO for 24 hours, and MACI was used to characterize subcellular changes in their phenotype with the goal of understanding LCO-algal interactions. R. subcapitata, was chosen as the model organism in this study as this particular algal species is a US EPA established model for environmental toxicology, 17 as well as an important bioindicator species for monitoring water quality.¹⁸ In addition, it maintains its unicellular nature under stress, which is ideal for downstream segmentation in bioimage analyses. In comparison, other common types of microalgae, like Chlamydomonas spp. or Scenedesmus spp., could potentially form colonies or coenobia in response to environmental stressors, 19,20 which would make it harder to distinguish individual cells from one another. Lastly, R. subcapitata was also chosen in order to make direct connections to previous algalbased LCO studies in the literature, which also used this algal species as a toxicological model organism.8 The phenotypic profiles of LCO-treated cells were compared to several reference compounds with established MoAs to predict the phytotoxic MoA of LCO. These reference compounds were chosen to represent MoAs that have been reported for other nanomaterials like membrane disruption, 21 DNA damage, 22 and more. The similarities between LCO and reference compound profiles were evaluated using hierarchical clustering based on Euclidean distance. An additional deep learning convolutional neural network (CNN) approach was also used to characterize the MoA experienced in individual cells as a means to predict LCO's phytotoxic MoA.

Materials and methods

LCO synthesis and characterization

Li_xCoO₂ nanosheets were synthesized using techniques described in previous studies. 8,23,24 18.2 MΩ cm⁻¹ water was used for each step during the synthesis. A (Co(OH)2) precursor was prepared using a precipitation reaction between LiOH and Co(NO₃)₂·6H₂O. 1 M Co(NO₃)₂·6H₂O was added to a 0.1 M solution of LiOH, drop-by-drop. The precipitate was isolated and washed with 3 cycles of centrifugation for 5 min at 4696 g

in order to isolate a pellet of particles, and then resuspended in water. Next, the supernatant was removed after washing and the solid product was dried in a vacuum oven at 30 °C overnight. The Co(OH)₂ precursor was then lithiated to form Li_xCoO₂ by adding 0.20 g Co(OH)2 particles to a molten salt flux of 6:4 molar ratio of LiNO3: LiOH at 200 °C in a PTFE container equipped with magnetic stirring in a silicone oil bath. The particles were heated and stirred in this molten salt flux for 30 min and the reaction was quenched with water. The precipitate was isolated and washed by 3 cycles of centrifugation for 5 min at 4696 g to isolate a pellet of particles, and were then resuspended in water. Then the product was isolated from the supernatant and dried in a vacuum oven at 30 °C overnight. The particles, which were digested in aqua regia, were analyzed using inductively coupled plasma - optical emission spectroscopy (ICP-OES) to yield a Li: Co ratio of 0.92:1. Surface area measurements, determined by nitrogen physisorption, yielded a surface area value of 125 m² g⁻¹. Individual LCO particles were imaged and sized using a FEI Tecnai T12 transition electron microscope (TEM). Only particles completely captured in each TEM micrograph were measured. Thickness was measured on particles that appeared very dark as this means they were viewed edge-on. Length of particles were measured if clearly defined endpoints were visible and if it could reasonably be assumed that it was a single particle as opposed to an aggregate. Dynamic light scattering (DLS) and zeta potential measurements of LCO suspensions in OECD 201 media were obtained with a Zetasizer Nano ZS Size Analyzer

Algal cell culture

from Malvern PANalytical.

A stock culture of R. subcapitata, inoculated at 1×10^5 cells per mL, was grown in a 1 L Erlenmeyer flask and cultured in OECD 201 media.²⁵ Cells were illuminated continuously in an incubator with a full spectrum T8 light bulb at a photon flux of 70 $\mu E\ m^{-2}\ s^{-1}$. The stock culture was mixed with an orbital shaker at a speed of 111 rpm.

Exposure setup

This exposure was done to assess predict the phytotoxic MoA of LCO on R. subcapitata by comparing changes in the complex phenotype of LCO-treated cells to that of reference chemicals with established MoAs after 24 hours of exposure. Each reference chemical and their associated MoA are reported in Table 1, below.

An additional LCO exposure was done at 48 hours to better evaluate physiological endpoints such as nucleation state. Algae were exposed to one of four LCO concentrations or untreated control (0 μg mL⁻¹, 0.01 μg mL⁻¹, 0.1 μg mL⁻¹, 1 μg mL⁻¹, & 10 μg mL⁻¹ LCO), an ion control that contained the concentration of lithium and cobalt ions that would be present in the algae media containing 10 μg mL⁻¹ of LCO after 24 or 48 hours, depending on the exposure duration. In each treatment, 900 µL aliquots of algal stock culture were seeded into individual 1.5 mL microcentrifuge tubes after

Table 1 Reference chemicals with known MoAs

Chemical	Mechanism of action	Abbreviation	Ref.
Aclonifen	Carotenoid biosynthesis inhibition	CBI	26
Carfentrazone	Membrane disruption	MD	27
DCMU	PSII photochemistry inhibition	PPI	28
Glufosinate	N ₂ metabolism inhibition	NMI	29
H_2O_2	Oxidative stress	OS	30
Metolachlor	Very-long-chain fatty acid synthesis inhibition	VLCFASI	31
MSMA	OP uncoupler/e ⁻ transport inhibition	OPU/e ⁻ TI	32
Zeocin	DNA damage	DD	33

cells were growing exponentially. A stock suspension of LCO was constituted at 100 μg mL⁻¹ in OECD 201 media. For this study, the OECD 201 media was made to be deficient in EDTA in order to prevent the mitigation of any metal-induced stress. This suspension was sonicated for 25 minutes before the addition to respective samples to break up any aggregated nanoparticles. Additionally, an ion solution made from LiOH and CoCl2·H2O, also constituted the OECD 201 media, was made at 10× the concentration of ion dissolution of 10 µg mL⁻¹ of LCO after 24 and 48 hours, depending on the exposure duration. For each treatment, OECD 201 media, LCO suspension, or 10× ion solution was added to each 900 μL cell suspension at a final volume of 1 mL. The samples were then placed under full spectrum illumination, with tube lids open, at a photon flux of 70 $\mu E \text{ m}^{-2} \text{ s}^{-1}$ for 24 hours. The 24 hour exposure duration was chosen for MoA prediction as this timepoint has been shown to better delineate initial phenotypic impacts,34 while the 48 hour exposure duration was chosen to examine physiological endpoints.

Single-cell chlorophyll fluorescence

At the conclusion of these exposures, a 50 µL aliquot from each sample was plated in to a well of a glass bottom 384 well plate (Cellvis, P384W-1.5H-N) and spun gently at 600 RPM for 1 minute to concentrate cells at the bottom of the well. Cells were not stained nor fixed with glutaraldehyde for this assay as to prevent interference with raw fluorescence. Cells were then dark adapted for 30 min to ensure PSII centers were in an open, or oxidized, state before imaging with an ImageXpress Micro XLS High-Content Screening System. For image acquisition, cells were visualized using a Cy5 filter (ex/em: 628/692) in a manner similar to that of a PAM Fluorometer, such that cells are exposed to a low intensity light beam followed by a saturating light beam to measure the minimal (F_0) and maximal (F_m) chlorophyll fluorescence, respectively (Fig. 1) using the fluorescence cytochemistry parameters in Table 2. After acquiring images, bioimage analysis was used to measure the respective fluorescence intensities. Using the "mean_integrated_intensity" values for F_0 and F_m , the variable fluorescence (F_v) and quantum efficiency of PSII (F_v/F_m) of individual cells were calculated using the following equation:

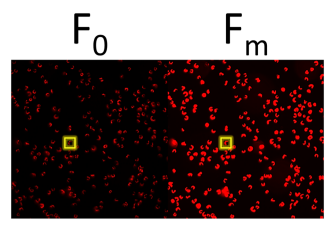


Fig. 1 Representative fluorescence micrograph of single cells at an F_0 and $F_{\rm m}$ state, respectively.

$$\frac{F_{\rm v}}{F_{\rm m}} = \frac{F_{\rm m} - F_0}{F_{\rm m}} \tag{1}$$

Multiplexed algal cytological imaging (MACI) assay

The MACI assay was carried out as using methods described by Ostovich & Klaper, 2023. At the end of the exposure, 905 µL aliquots from each sample were transferred to sterile 1.5 mL microcentrifuge tubes. Commercially available fluorescent probes were used to stain nuclei and lipid droplets using NucBlue (Thermo Fisher, R37605) and BODIPY 505/515 (Thermo Fisher, D3921), respectively, and glutaraldehyde was used to fix the algal cells. After adding reagents, all reactions were incubated overnight, at 4 °C to minimize enzymatic degradation and maintain the integrity of the subcellular structures. After incubating reactions, cells were centrifuged at 4000×g for 5 min, washed 2× with 1× phosphate buffered saline (PBS), and resuspended in PBS. Cells from each sample were loaded into a well of a glass bottom 384 well plate (Cellvis, P384W-1.5H-N) at a seeding density of $\sim 2 \times 10^3$ cells per mm² for optimal distribution of cells across the well surface. After loading cells, the well plate was then spun gently at 600 RPM for 1 minute to concentrate cells at the bottom of the well. Images were acquired at 9 sites per well with an ImageXpress Micro XLS High-Content Screening System with a 60× Plan Fluor 0.85 NA air immersion objective (Molecular Devices, 1-6300-0414), using the Cy5, GFP, and DAPI fluorescent channels to visualize the chloroplast, lipid droplets, and nuclei, respectively. To enhance image contrast and resolution, the digital confocal feature was used during image acquisition. For representative cell images with higher resolution, some images were also acquired with a 100× CFI L PLAN EPI CC 0.85 NA air immersion objective (Molecular Devices, 1-6300-0419).

Bioimage analysis

After acquiring images CellProfiler, 35 was used for image preprocessing, object segmentation, and morphological feature extraction at the resolution of individual cells. Morphological features related to area, shape, intensity, and granularity of each subcellular structure were extracted, in addition to cytoplasmic intensity features to add more measurements for comparing the phenotypic profiles of LCO and reference chemical treatments, which were run on separate plates. These data were exported to a local SQLite database file and were then extracted using the RSOLite package in R.³⁶

Phenotypic profiling - fingerprint analysis

Phenotypic response data was analyzed using the methods described by Ostovich & Klaper, 2023.9 The data was firstly processed by aggregating single-cell morphological feature measurements to per-image and then per-well values, which was done by taking the cell and image means, respectively. Secondly, well data from each compound and dose were then normalized to the non-treated cell control by computing a Z-score. In order to verify whether LCO elicited a change to the entire phenotypic profile of treated cells, a partial least squares-discriminant analysis (PLS-DA) was performed in R using the mixOmics package.³⁷ Before feeding phenotypic response data into the PLS-DA models, an ANOVA was performed across all features for each reference chemical to remove any non-informative features with little variance (p-values > 0.05). Lastly, factor analysis was used to further reduce the dimensionality of phenotypic data vectors, and the fingerprints were subsequently compared to one another using hierarchical clustering based on Euclidean distance in R.

Phenotypic profiling - convolutional neural networks

In addition to fingerprint analysis, a CNN was also trained on a small subset of reference compound treated cells (~9.3%) using the classifier module on CellProfiler Analyst (Ver 3.0).38 A separate bin was created for each reference chemical and the non-treated cell control in the classifier module, where around 1000 randomly fetched cells from each treatment were placed in each respective bin. For the non-treated cell control, cells form the reference chemical exposure and LCO exposure were both used to account for plate-to-plate and run-to-run variations. After training the CNN, it was used to score the entire experiment by classifying

Table 2 Single-cell chlorophyll fluorescence cytochemistry parameters

Intensity	Channel	Excitation (nm)	Emission (nm)	Exposure time (nm)
Minimal (F_0)	Cy5	628/40	692/40	35 ms
Maximal (F_m)	Cy5	628/40	692/40	270 ms

individual cells into predicted mechanistic classes, and computing enrichment scores for each sample as the logit area under the receiver operating characteristic curve. An ANOVA and a Tukey post hoc test was used to evaluate the significance of predicted mechanistic class enrichments for each treatment.

Statistical analysis

All statistical analyses were performed using R Studio. 39,40 A Shapiro-Wilk test was used to verify normal distribution and a One-Way ANOVA was used to compare variance among group means, while a Tukey post hoc test was used for multiple comparisons. In each analysis, significant differences were determined with a 95% confidence interval.

Results and discussion

LCO characterization

The sizing of single LCO particles, using TEM micrographs (found in ESI†) suggested an average thickness and length of 5.54 ± 2.01 nm and 39.63 ± 17.35 nm, respectively. The zeta potential values for LCO suspended in OECD 201 media (found in ESI†) indicate that this nanomaterial is negatively charged between -17 and -20 mV in higher concentrations, and around -8 mV in the lowest concentration (0.01 µg mL⁻¹). However, at all concentrations, the lower magnitude of the zeta potential (<|30| mV) suggests that LCO is not highly stable in suspension with OECD 201 media. Furthermore, the hydrodynamic diameter of LCO also generally increases with increasing concentration, thus also suggesting increasing aggregation of LCO nanoparticles with increasing concentration. These results generally coincide with LCO's behavior in other aqueous media, like moderately hard reconstituted water⁴¹ and Bold's modified freshwater solution,8 as well.

LCO significantly alters algal cell morphology after 24 hours

After exposing R. subcapitata with LCO for 24 hours, the cells were stained and imaged using the MACI protocol (Fig. 2a). In the representative fluorescence micrograph, the LCO-treated cells appear to be larger compared to the control cell, and also appear to have distinct levels of chlorophyll, TAG, and DNA content, which visually support the claim that LCO has an impact on cell morphology. A CellProfiler pipeline was used to convert the high-content image data into quantitative data where 570 unique, unbiased, morphological features of individual cells were used to generate phenotypic fingerprints of molecular interaction. In order to quantitatively verify if LCO treatment significantly alters algal cell morphology, a PLS-DA was used to analyze subtle changes in the complex phenotypic profile of LCO-treated cells with increasing concentration. Based on the PLS-DA response plot (Fig. 2b), LCO-treated cells display a significant separation between response groups, thereby indicating that LCO does elicit a significant, and measurable, change to cell morphology after 24 hours. Based on X-variate data, a significant separation from the control group can be seen at starting at the 0.1 µg mL⁻¹ response group. Here, the ion control response group also exhibited a significant separation from the control group. However, the ion control response group, which represents the amount of Li⁺ and Co²⁺ ions released from 10 µg mL-1 of LCO after 24 hours, was also significantly different form the 10 µg mL⁻¹ response group. Interestingly, this data suggests that while the ions do have somewhat of an impact on cell morphology, they are separate from nano-specific impacts.

Phenotypic profiles of LCO-treated cells compared to reference chemical-treated cells

In order to predict the phytotoxic MoA of LCO, the phenotypic profiles of LCO-treated cells were compared to reference

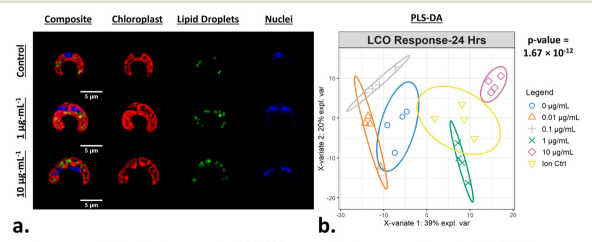


Fig. 2 Phenotypic responses of algal cells when exposed to LCO. (a) Representative fluorescence micrograph of MACI labeling patterns in treated and non-treated algal cells. (b) A PLS-DA response plot graphically describes the change across complex morphological feature data with increasing concentration of LCO; ellipses represent 95% confidence intervals and p-values represent ANOVA statistics across the 1st latent variable between response groups.

chemical-treated cells with known MoAs. For this purpose, the 10 μM reference chemical data was compared to the 1 μg mL⁻¹ LCO data, which is around 10 µM of LCO, to compare responses of similar concentrations.

After constructing the phenotypic fingerprints, an ANOVA was used to identify individual features that carry little information, which were removed from the analysis given a p-value > 0.05. Additionally, factor analysis was used to further reduce the dimensionality of the phenotypic data vectors down to 7 eigen features/factors in order to minimize redundant measurements while preserving variance. The phenotypic fingerprints across all replicates for each reference chemical and LCO treatment were hierarchically clustered based on Euclidean distance (Fig. 3). Euclidean distance is a data mining technique that measures how far apart two points are in a high-dimensional feature space, 42 and can be calculated as the square root of the sum of squares between two vectors.43

The hierarchical clustering analysis was able to identify four main clusters. LCO-treated samples were clustered with DMCU and three of the MSMA-treated samples, thus indicating that their phenotypic profiles are most similar to one another. The MoAs of DMCU and MSMA are both related to electron transport inhibition in the chloroplast and mitochondrion, respectively. 28,32 Based on the results of the hierarchical clustering analysis, we can deduce that the probable MoA of LCO is also likely related to electron transport inhibition. This would makes sense as LCO tends to target proteins and pathways that are involved in transport of electrons.7,6

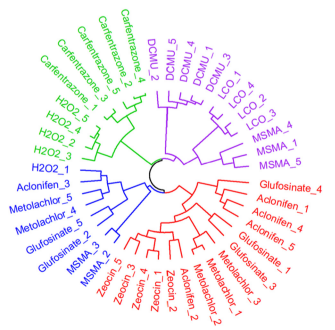


Fig. 3 Dendrogram of hierarchically clustered LCO treated samples compared to reference chemical threated samples. Branches represent relative Euclidean distances between samples and nearest neighboring samples indicate the most similar phenotypic profiles to one-another.

Predicting the MoA of individual LCO-treated cells using convolutional neural networks

In addition to the hierarchical clustering analysis, a convolutional neural network was also used to classify individual LCO-treated cells into mechanistic classes. In CellProfiler Analyst, a CNN was trained on a small subset of randomly fetched cells from each treatment (~9.3% of cells from the combined LCO and reference chemical exposures) using 50 × 50 neurons per layer. For the healthy cell class, the CNN was trained on cells form the non-treated cell controls of both the LCO and reference chemical exposures as a means to account for variations due to separate plates and separate runs. Based on the confusion matrix (Fig. 4), the CNN model was able to predict the correct mechanistic class across training cells with a moderately good classification accuracy of 73.82%. Once trained, the CNN model was used to score each cell in the LCO exposure, based on its individual phenotype, with a predicted mechanistic class, and then calculate enrichment scores for each sample.

The average enrichment scores for the LCO exposure are visualized in a heatmap (Fig. 5) where values with asterisks (*) represent the mechanistic classes which are significantly enriched in each treatment when compared to the control. Based on these results, the CNN model predicted PSII photochemistry inhibition, or photosynthetic electron transport, as the most probable MoA of LCO with the highest and most significantly enriched scores. This was most notable in the 1 µg mL⁻¹ and ion control treatments. These treatments also obtained high enrichment scores in the oxidative phosphorylation uncoupler/mitochondrial electron

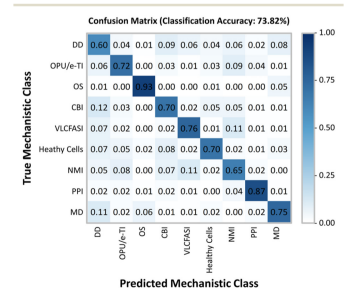


Fig. 4 Convolutional neural network construction. A small subset of randomly fetched cells in each treatment from the reference chemical and LCO exposures were fed into a convolutional neural network model. Based on the training data, this CNN model yields a classification accuracy of 73.82% at correctly classifying cells by their true mechanistic class.

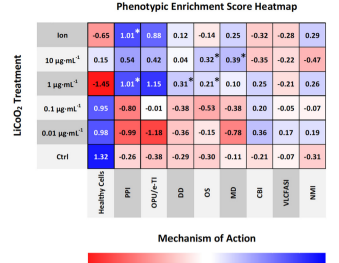


Fig. 5 Phenotypic enrichment score heatmap. The CNN model classifies each cell across the entire LCO exposure with a predicted mechanistic class based on its phenotype. Enrichment scores for each mechanistic class are calculated in each sample. Heatmap values represent average treatment enrichment scores; values with asterisks (*) represent significantly enriched mechanistic classes compared to the control

transport inhibition mechanistic class; however, the scores were not significantly different from the control due to larger deviations in the enrichment data. This data coincides with the hierarchical clustering data (Fig. 3) in that electron transport inhibition in general appears to be the most probable phytotoxic MoA of LCO, but the CNN data suggests that this disturbance is more prevalent in the photosynthetic pathway.

Part of the advantage of using a CNN classifier model for phenotypic profiling is the fact that it can identify rare objects,³⁸ or smaller subpopulations of cells, in an otherwise heterogeneous cell population, that have distinct mechanistic classes that may go undetected with other phenotypic profiling methods that average across all cells. Interestingly, based off the enrichment data, there were also three other significantly enriched mechanistic classes found in much smaller subpopulations of cells. In the 1 µg mL⁻¹ treatment, the DNA damage and oxidative stress mechanistic classes were also significantly enriched. Additionally, in the 10 μg mL⁻¹ treatment, the oxidative stress and membrane disruption mechanistic classes were significantly enriched. These results may make sense as these mechanistic classes were only enriched in the highest concentrations of LCO, which are inherently more likely to have more cells undergoing a cytotoxic shock in which disturbances such as DNA damage, membrane disruption, and high levels of oxidative stress are experienced. However, it is important to reiterate here that the main MoA experienced by these cells is still largely photosynthetic electron transport inhibition as the other mechanistic classed were only indicated in a small percentage of cells.

Chlorophyll fluorescence markers indicate that LCO alters the oxidation state of PSII proteins

Chlorophyll fluorescence is a non-invasive way to determine relative amounts of chlorophyll content in plant-type organisms and to assess the overall efficiency of photochemistry. 44,45 Several methods currently exist to make these sorts of measurements: pulse amplitude modulation fluorometers, for example, are typically used to make these measurements in leaf tissue and algal cells. PAM's work by, first, exposing the sample to a pulse of low intensity light to get the minimal fluorescence, as most of the energy will enter photochemistry and only a small portion will be re-emitted as fluorescence signal. This is then followed by a pulse of saturating light to get the maximal fluorescence, which at this point, the PSII reaction centers will already be reduced and thus most of the excited energy will be re-emitted as fluorescence signal. In general, the minimal fluorescence is comparable to the levels of chlorophyll present in leaf and algal samples.⁴⁴ Furthermore, by taking the ratio of minimal and maximal fluorescence, the quantum efficiency of PSII can be calculated, 46 which is a good indicator of photochemistry efficiency. 47 Here, this same concept was applied using high-content fluorescence microscopy for algal cells, and with this tool, the relative chlorophyll levels and quantum efficiencies of PSII evaluated for individual cells.

Fig. 6a displays measurements of single-cell chlorophyll fluorescence after being exposed to LCO for 48 hours; each red dot represents an individual cell. LCO treated cells had exhibited significant increases in chlorophyll content compared to that of the control (Fig. 6b). A quantum efficiency of PSII around 0.7 is considered healthy and/or normal for eukaryotic algal cells, 48,49 so based on the data in Fig. 6c, even LCO treated cells are still within a normal range. Interesting, however, LCO treated cells exhibited a significant decrease in the quantum efficiency of PSII, compared to the control, in somewhat of a dose-dependent manner. In the context of the PSII biochemistry, this data would suggest that, in LCO treated cells, there is a higher fraction of damaged, or inactive, PSII reaction centers.⁵⁰ This type of disturbance is typically due to the irreversible oxidation of the D1 and D2 proteins in the PSII complex, usually following the production of O2. and OH radicals.⁵¹ These results, coincide with the hierarchical clustering (Fig. 3) and CNN (Fig. 5) analyses which predicted photosynthetic electron transport inhibition as the main phytotoxic MoA of LCO. Additionally, these results could help explain the reductions in the net production of carbon biomass reported in previous studies.8

LCO significantly impacts the nucleation state and DNA content after 48 hours

In eukaryotic cells, the nucleus in an important subcellular compartment which houses the genetic material of the organism and is responsible for regulating gene expression and facilitating cellular division. Most green algae contain a unique

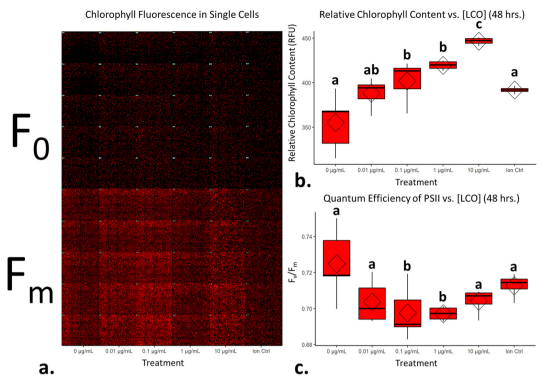


Fig. 6 LCO significantly alters markers of chlorophyll fluorescence. (a) Representative montage micrograph of algal cells across all treatments at a state of minimal and maximal fluorescence after 48 hours of exposure to LCO; (b) relative chlorophyll content after 48 hours of exposure to LCO; (c) quantum efficiency of PSII. Diamonds on boxplots represent treatment means and significant differences were determined using a one-way ANOVA with a Tukey post hoc test for multiple comparisons; columns with different letters differ significantly (p < 0.05).

multiple fission reproductive pattern in which one mother cell can divide into several daughter cells, depending on the environmental cues. 19 R. subcapitata, in particular, has the ability to divide into eight daughter cells,18 and measuring the number of nuclei per cell, as well as the relative amount of DNA content per nucleus, can be used to describe instances of cell cycle disruption/arrest. 31,19,52 After exposing cells to LCO for 48 hours, cells were stained with NucBlue, in order to fluorescently label the DNA content within the algae. A CellProfiler pipeline was used the count the number of nuclei/cell, as well as measure the integrated intensity of each nuclei, which was taken as relative DNA content. On average, LCO-treated sample contained a higher percentage of cells with only one nucleus when compared to the non-treated samples (Fig. 7a). This affect increased in a dose-dependent manner, with significant differences from the control observed at 10 µg mL-1 LCO and the ion control. The opposite trend was observed for cells with more than one nucleus (i.e. 2 & 4 nuclei), which also changed in a dose-dependent manner, with significant differences from the control observed at 10 µg mL-1 LCO and the ion control. Looking only at the cells with one nucleus, LCO-treated cells had, on average, a lesser relative amount of DNA content when compared to the non-treated control cell (Fig. 7b). This was also observed in a dose-dependent manner, but with significant differences from the control at 0.1 μg mL⁻¹ LCO, 10 μg mL⁻¹ LCO, and the ion control. Interestingly, when assessing the

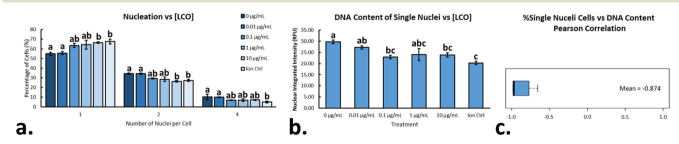


Fig. 7 Nucleation state and relative DNA content in LCO-treated cells. (a) Percentage of cells with 1, 2, or 4 nuclei after 48 hours of exposure to LCO; (b) relative DNA content of single nucleated cells; (c) Pearson correlation between the percentage of single nucleated cells and their relative DNA content. Significant differences were determined using a one-way ANOVA with a Tukey post hoc for multiple comparisons; columns with different letters differ significantly (p < 0.05). Error bars represent SEM.

Pearson correlation between the two trends, a strong negative correlation was observed (Fig. 7c). Together, these results indicate that, on average, LCO-treated cells exhibit an increased instance in the delay/arrest of cell cycle progression, specifically in the earlier stages of the cell cycle (stage 1/2) before first nuclear division.⁵² Based on these results, it may explain the instances of increased growth inhibition and biovolume of LCO-treated algal cells reported in previous studies.8 In this case, it may be possible that LCO had an impact on nuclear/ cellular division through the impairment of photosynthetic electron transport.⁵³ These results are similar to that of DCMU treated cells, which, on average, had been reported to have significantly less nuclei/cell when compared to non-treated control cells, and had appeared to have inhibited nuclear division.9 This would also be consistent with other photosynthetic organisms, like Euglena gracilis, in which similar impacts have been reported under impaired photosynthetic electron transport.53

Conclusion

In this study, MACI and machine learning techniques were used to assess the interactions of LCO at the biointerface of R. subcapitata cells and to predict the phytotoxic MoA of LCO. Algal cells were exposed to varying concentrations of LCO, and their phenotypic profiles were compared to that of cells treated with reference chemicals with established MoAs. The described analyses predicted photosynthetic electron transport to be the most probable phytotoxic MoA of LCO, and single-cell chlorophyll fluorescence demonstrated an increase in irreversibly oxidized photosystem II proteins, thus fortifying the MACI assay prediction and coinciding with the impaired carbon biomass assimilation reported in previous work. Lastly, LCOtreated cells were observed to have less nuclei/cell and less DNA content/nucleus when compared to non-treated cell controls, which suggests an interference with cell cycle progression, also complementing the growth inhibition and biovolume data reported in previous work.

Conflicts of interest

There are no conflicts to declare.

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