

Title: Air pollution alters *Caenorhabditis elegans* development and lifespan: responses to traffic-related nanoparticulate matter (nPM)

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Abstract

Air pollution is a heterogeneous environmental toxicant that impacts humans throughout their life. We introduce *Caenorhabditis elegans* as a valuable air pollution model with its short lifespan, medium-throughput capabilities, and highly conserved biological pathways that impact healthspan. We exposed developmental and adult life stages of *C. elegans* to airborne nano-sized particulate matter (nPM) produced by traffic emissions and measured biological and molecular endpoints that changed in response. Acute nPM did not cause lethality in *C. elegans*, but short-term exposure during larval stage 1 caused delayed development. Gene expression responses to nPM exposure overlapped with responses of mouse and cell culture models of nPM exposure in previous studies. We showed further that the *skn-1*/Nrf2 antioxidant response has a role in the development and hormetic effects of nPM. This study introduces the worm as a new resource and complementary model for mouse and cultured cell systems to study air pollution toxicity across the lifespan.

Keywords: *Caenorhabditis elegans*, air pollution, nPM, *skn-1*, development

Introduction

Epidemiological studies show that air pollution is associated with multiple chronic health hazards of older age including Alzheimer's disease (AD), ischemic heart disease and stroke, lung cancer, and chronic obstructive pulmonary diseases - all of which decrease life expectancy (1-5). Understanding the underlying mechanisms between air pollution and these diseases requires modeling both air pollution and the resulting biological responses.

Traffic related air pollution (TRAP) particles are a complex environmental toxicant consisting of a variety of inflammogens and toxicants derived from vast heterogeneous sources. It is estimated that ambient particulate matter (PM <2.5 micron diameter) contributes to 3-4 million deaths (around 7.6% of total global death) and 103.1 million (4.2%) global disability-adjusted life year annually (1). The mortality number is apart from 3.5 million deaths attributed to indoor air pollution such as solid fuels for cooking and heating (6). It is essential that gerontological research understand the effects of environmental toxicants over organismal lifespan, the association with common aging phenotypes, and also identify the biomarkers that can predict these effects (reviewed in (7)).

The nanosized subfraction of TRAP (nPM) has consistent toxic effects in rodent and cell models (8). Biochemical and cell assays of air pollution toxicity, while widely used, are not good predictors of *in vivo* responses for multicellular organisms (e.g. Dithiothreitol [DTT], ascorbic acid [AA]-glutathione [GSH], and MTT (9). Exposure of mouse models to reaerosolized nPM results diverse systemic and organ-specific localized effects that involve different biological networks such as oxidative stress and antioxidant responses, innate immunity, and the nervous system (10-13). These responses were dependent on the dosage of PM samples and the developmental stage of the exposure. Mouse models are a valuable tool to study air pollution toxicity, but they are limited as a biological model; low

reproduction yield, long lifespan, expense, and ethical considerations in these animals can reduce the feasibility of this model and also the statistical power of any experiments.

Caenorhabditis elegans is a valuable model for TRAP toxicology with a potential for much higher throughput than rodents (14). Humans and worms share several basic physiological and stress response processes with homologues in most human genes (60-80%), including multiple signal transduction pathways (15). Easy maintenance, large scale production, small size, body transparency, full genomic characterization, complete cell lineage map, and mutant libraries make the worm an ideal model for gene network and environment interactions (16). While there are limitations such as lack of specific organ equivalents, a smaller immune response system, and large differences in overall lifespan, *C. elegans* allows medium-throughput whole organism-level assays with multiple end points (e.g. development, reproduction, feeding, lifespan, locomotion) (15). Worms have been used to assess the toxicity of terrestrial environmental samples (e.g. soils, sludge, river sediments, (17, 18), pesticides (e.g. Glyphosate, Paraquat, Endosulfan and Dichlorvos for neurotoxicity, DNA damage, sterility and embryonic lethality) (19), metal toxicity (e.g. Ag, Cd, Pb, Fe), lifespan, fertility, growth (20), nanoparticles (21), drugs (22), toxins (e.g. nicotine) (23-25), as well as other bioreactive molecules including NaAsO₂, NaF, caffeine and DMSO (26). While airborne bacteria have been tested in *C. elegans* (17), its responses to TRAP have not been studied. Considering the usefulness of *C. elegans* in diverse toxicology models, this study introduces *C. elegans* as a multicellular model organism for air pollution toxicity.

Our prior mouse and cell culture studies show that nPM can induce oxidative stress, systemic inflammation, and selective neuroinflammation in different brain regions as well as exacerbate amyloidogenesis processes in AD mouse models (5, 10-13). We also noticed that there is an interaction between age and air pollution mediated toxicity (10). Here we examined different nPM dosages, as well as developmental and lifespan effects in *C. elegans* to evaluate the similarities of this model to higher

organisms in several cell survival pathways and Alzheimer A β -related genes. In contemplating the response of cytoprotective transcription factor SKN-1/Nrf2 in *C. elegans* to nPM, we also investigated the association of SKN-1 in an nPM-mediated developmental delay phenotype as well as changes in lifespan.

Method

C. elegans strain maintenance

C. elegans were maintained at 20°C unless otherwise noted. Strains used were Bristol N2 (wild type), LG335 *skn-1(zu135)/nT1[qIs51(myo-2::GFP;pes-10::GFP;F22B7.9::GFP)]*, and CL2166 *dvIs19 [(pAF15)gst-4p::GFP::NLS]* III. Some strains were provided by the CGC, funded by NIH Office of Research Infrastructure Programs (P40 OD010440). LG335 was a gift from the Leonard Guarente laboratory. *E. coli* strain OP50 was used for all non-RNAi experiments and for general *C. elegans* maintenance. For synchronization prior to plating, gravid adults were treated with a solution of bleach and hypochlorite to harvest eggs; then, eggs were washed and rocked overnight in M9 solution, allowing the hatching and L1 arrest/synchronization (27).

Airborne nano-sized particle collection:

Ambient nano-sized particles (diameter<0.18 μ m) were collected on 8x10 inch commercially available Zeflour PTFE filters (Pall Life Sciences, Ann Arbor, MI) using a High-Volume Ultrafine Particle (HVUP) Sampler (28) operating at a sampling flow rate of 400 liters/min flow rate at the Particle Instrumentation Unit (PIU) of University of Southern California located within 150m downwind of a major freeway (I-

110). Gravimetric mass (nPM mass concentration) was determined from pre- and post-weighing the filters under controlled temperature (22–24 °C) and relative humidity (40–50%) conditions. The filter-deposited nPM was eluted by sonication into ultrapure deionized (milli-Q) water (11) providing the concentrated slurry suspension used for these exposures. A portion of the aqueous suspension was chemically characterized. After acid digestion, samples were analyzed by high resolution inductively coupled plasma sector field mass spectrometry (SF-ICPMS). Another portion was analyzed using a Sievers 900 Total Organic Carbon Analyzer to determine total organic carbon (TOC) content (Fig. S1). This characterized suspension was used in all the experiments of this study.

RNA interference

The *E. coli* strain HT115 (DE3), harboring either the empty L4440 plasmid ("Control RNAi") or the *skn-1* RNAi plasmid (dsRNA production of *skn-1* sequence - Ahringer Library), was grown 16-18hrs at 37°C overnight. Cultures were seeded onto RNAi plates (normal NGM plates with 5mM isopropyl-β-D-thiogalactoside (IPTG) and 50 µg/ml carbenicillin) and left overnight to generate dsRNA for experiments (maintained at 20°C during and after dsRNA generation). To optimize RNAi of *skn-1* in offspring, P0 worms were plated on bacteria expressing *skn-1* RNAi for 12, 18, 24, or 48 hrs (Fig. S2); this was done to further reduce maternally-deposited *skn-1* mRNA transcripts as well as to deposit *skn-1* RNAi in the F1 generation prior to hatching. To control developmental and RNAi timing, these adults were placed in 15°C for the duration of *skn-1* RNAi exposure. Adult exposure for 24 or 48 hr caused 90-100% of dead F1 eggs. 18 hrs was chosen as an acceptable 20-50% egg death while decreasing *skn-1* transcripts in living animals to ~70% (Fig. S2).

Optimization of air pollution exposure model: Two routes of nPM exposure (liquid or chronic exposure on growth medium plates [chronic exposure data not shown]) and duration (1, 2, 4, 8, 24 hours) were tested in larval stage 1 or 4 (L1 or L4) *C. elegans*, as well as solvents (M9 buffer or K medium). 1 hr exposure in diluted nPM at different dosages (1-200 μg nPM/ml) with M9 was chosen for further experiments as it was the fastest exposure in inducing phenotypes without compromising developmental timing. For treatments, worms were washed into an Eppendorf tube, brought to a known volume, and nPM was added to each tube to achieve the listed concentration. Worms were incubated at 20°C for all experiments, unless otherwise indicated. Worms were gently rocked 1 hr for even distribution of nPM. After exposure, worms were washed once before plating at time 0 in post exposure time. Worms treated as "L1s" were treated immediately at the synchronized developmental stage (prior to feeding).

Size analysis:

Following exposure of L1 stage for 1 hr to nPM and re-plating, worms were incubated at 20 °C for 72 hrs. Body size was analyzed by area using ImageJ (average of width \times length).

Pharyngeal Pumping:

Worms were treated with 200 $\mu\text{g}/\text{ml}$ nPM for 1 hr at L1 stage, then allowed to recover for 24hrs on regular agar food plates described above (~100-150 worms/plate). These plates were then directly placed under a 10x objective, allowed to settle for 1 minute to prevent changes in pumping due to plate movement/vibrations, and individual worms were then followed and recorded using the Movie Recorder in the ZEN 2 software at 6-8ms exposure (Zeiss Axio Imager) for 10-15 seconds (making sure to

keep the pharynx in frame). Slowed-down (~4x) movies were later analyzed for pharyngeal pumping rate. Worms without pharyngeal pumping (dead, lethargus) during recording were excluded (estimated <5% of total worms observed).

Fluorescent analysis of *gst-4p::GFP* labeled *C. elegans*:

gst-4p::GFP animals were treated with the listed nPM concentration (0-200 $\mu\text{g/ml}$) for 1 hr and then imaged 4 or 24 hrs after exposure. Recovered worms were then plated into 10-12 μl droplets of M9 solution directly onto slides and 10mM sodium azide was added. Once animals were paralyzed, coverslips were slowly placed at a diagonal angle on top of the M9-immersed animals to prevent bursting/flattening. Slides were then immediately taken to image whole worms using an oil-based 40x objective (Numerical aperture 1.4) with DIC and GFP filters (Zeiss Axio Imager, an epifluorescence microscope). Animals were imaged to have as much of the pharynx, germline, and intestine in focus as possible. As prolonged exposure to sodium azide can eventually induce *gst-4::GFP* (29), each plate was imaged within 20-30 minutes, and a new plate was made after this time if more worms were needed - both done to avoid inflation of the GFP signal. Later, fluorescence was measured (the "measure" command) on the imaged worms via ImageJ using the polygon selection tool surrounding both the whole animal and a section containing no worm(s) next to the animal (within the same image) as background (Whole worm autofluorescence background was not considered in the analysis). Corrected total cell fluorescence (CTCF) was then determined using Microsoft Excel. $\text{CTCF} = \text{Integrated Density} - (\text{Area of selected cell} \times \text{Mean fluorescence of background readings})$.

Lifespan

C. elegans were treated with 50 µg/ml nPM at either L1 stage or at L1 and Day 2 adult (~96 hrs). Synchronized L1 animals (after nPM exposure) were plated on control RNAi or *skn-1* RNAi. Worms were re-plated each day of reproduction (Day 1-5) to avoid overcrowding from progeny. Individuals were checked each day for survival by prodding with a platinum wire to verify touch responsiveness. Individual worms were censored from survival analysis for gross morbidity (bursting, vulval protrusion, crawling off plate, etc.).

Statistics: Size and *gst-4p::GFP* analysis results were compared using ANOVA followed by Tukey post hoc with calculating multiplicity adjusted P values by Graphpad Prism 7. In qPCR time points and the experiments with only two groups, pairwise comparison was done with T distribution tests in Graphpad Prism 7. Lifespan was analyzed by log-rank and Wilcoxon tests in JMP software. Heatmap and dendrogram analysis was done in R by the CompexHeatmap package of Bioconductor using Euclidean distance matrix and complete linkage for clustering of the genes. Factorial models were made from size, *gst-4p::GFP*, and *skn-1* RNAi data to analyze the significance of interaction terms using R.

Quantitative real-time PCR:

L1 or L4 worms were exposed to 50 µg/ml nPM for 1 hr, and then worms were collected and washed in M9 solution at 0 (immediately after 1hr exposure), 1, 2, 4, or 8 hrs after exposure. "Populations" in figure legends refer to biological replicates of 500-1000 worms. Worms were pelleted at collection, 500 µl of TriZol reagent (Zymo) was added. RNA was extracted using TriZol method (11) with an additional step of treatment with DNase, lysis by T&C lysis and protein precipitation by MPC reagent (Epicenter, USA) for purification. RNA concentration and purity were assessed by O.D. 230, 260, and 280 by

spectrophotometry before qRT-PCR with gene specific primers in Table S1. The data was normalized to *ama-1* as housekeeping gene. For Figure 2, *skn-1* RNAi was administered to P0 adults, and F1 progeny were analyzed (as "RNA interference" above).

3. Results:

3.1. Acute nPM exposure drives a reduction in *C. elegans* body size

Different exposure methods were tested to determine the acute lethal dosage 50 (LD50) of nPM samples. Larval stage 1 and 4 (L1 and L4, chosen to represent near the beginning and end of development) wild type N2-Bristol worms (~100 per replicate; at least 3 replicates/group) were exposed to a range of nPM concentrations (0.5-200 µg/ml) for 1, 4, or 24 hrs in liquid culture or on growth medium plates. *C. elegans* did not show any acute mortality at the highest dose in any of tested conditions during 3 days after exposure.

However, nPM exposure did affect development in a dose-dependent manner (Fig. 1A). Exposure of L1 stage animals to nPM in M9 buffer (10, 50 and 200 µg/ml) for only 1 hr showed dose dependent size reduction (13-26% of the worm's area) in day 1. This size difference was not due to lack of food intake because pharyngeal pumping rates were not impaired (Fig. 1B). Since nPM at 50 µg/ml caused the same size reduction as 200 µg/ml, the lower dose was used for further experiments.

3.2 *skn-1* is activated in response to nPM exposure

Given the size impairment under nPM exposure, we were interested in the underpinning biological response - including any defense mechanisms that were activated. TRAP nPM is a heterogeneous environmental toxicant consisting of multiple elements and water-soluble organic carbons with high oxidant activity (Fig. S1). We chose to examine SKN-1/Nrf2, a cytoprotective

transcription factor, due to its established anti-oxidant defense response (30) and role in development (31); specifically, we used the reporter *gst-4p::GFP* (GFP linked to Glutathione S-transferase, a target of SKN-1) as an estimate of activation of SKN-1 in animals treated with nPM. In the *gst-4p::GFP* strain, both L1 and L4 had dose dependent increases in response as early as 4 hr post-exposure to nPM. By 18 hr post-exposure of L1 stage and 24 hr post-exposure of L4 stage, we observed reduction of GFP signal in most of the nPM exposed animals (all groups except for 10 µg/ml nPM exposed L1s) (Fig. 2A, Fig. S3) suggesting a rapid return of reversal of the gene expression changes (perhaps compensatorily) post-SKN-1 activation.

To determine the role of SKN-1 activation in response to nPM stress, we down-regulated SKN-1 by RNAi before exposure to nPM. First, we tested the effects of short term nPM exposure on lifespan at different life stages to try to understand if certain exposure windows were more important for any lifespan effects. Contrary to the negative survival effects observed in cell lines (9), dual nPM exposure (50 µg/ml) during development (L1 and L3), or triple nPM exposure (50 µg/ml) during reproduction stage (D1, D3 and D5) had a modest increase of the lifespan. Increase of exposure intervals by including of both developmental and reproduction stages removed this lifespan difference. We then examined if dual nPM exposure (50 µg/ml) would have the same outcome in *skn-1* knockdown worms. While worms fed with control RNAi had a modest increase in mean lifespan (1.1-day, $p=0.015$ Log rank test, analysis in 90% of total lifespan), reducing *skn-1* expression by RNAi ablated this increase in lifespan (Fig 2B, Table S2). In general, *skn-1* knockdown lead to decrease of total lifespan regardless of nPM exposure (Fig S4). While our confidence in the observed lifespan change is limited by the number of animals used ($n=200-300$ per replicate), we are confident in our finding that the high nPM concentrations utilized do not negatively impact *C. elegans* lifespan.

We investigated *skn-1* knockout (*skn-1(zu135)*) for its response to nPM. While L1 exposure of wild type worms to 50 µg/ml nPM reduced day 1 adult size by 20%, *skn-1(zu135)* worms did not show this decrease in size (Fig. 2C). In both conditions, *skn-1(zu135)* worms were smaller overall compared to wild type.

3.3 Gene expression changes induced by nPM exposure

To understand molecular responses to nPM, selected mRNAs were measured by qPCR to analyze a panel of genes involved in *C. elegans* stress responses, development, vitellogenesis, innate immunity, amyloid processing, and TGF-β signaling pathway in L1 or L4 (Fig. 3, S5-S8). In general, animals exposed at L1 had larger nPM mediated mRNA changes, particularly in the first hours of exposure compared to L4 animals; moreover, the genes with largest nPM responses in L1 stage did not change in L4 stage exposed animals (e.g. *gst-4*, *daf-2*, *apl-1*, *sel-12*).

In L1 stage animals, 1 hr exposure to nPM immediately (0 hr post-exposure) changed the expression of several genes (Fig. 3, S6-8). Some of the observed changes at this time included heat shock responses (e.g. *hsp-4* [expression/control=0.5 fold]), hormone receptor and development (e.g. *daf-2* [0.58], *daf-12* [0.61]), vitellogenin (e.g. *vit-6* [0.64]) and Alzheimer amyloid processing genes (e.g. *apl-1* [0.64], *lrp-1* [0.68] and *sel-12* [0.55]) and TGF-β signaling pathway (e.g. *daf-7* [0.53]). At 1 hr post-exposure, the expression of most genes returned to baseline (pre-exposure) levels (exceptions were *hsf-1* [0.6], *hsp-4* [0.47] and *daf-2* [0.82]). This shift continued and lead to upregulation of several genes at 2 hr post-exposure of L1s in response to nPM. These responses consisted of *skn-1* antioxidant target genes (e.g. *gst-4* [3.5]), metal response genes (e.g. *cdr-1* [5.7]), innate immune response (e.g. *abf-2* [2.8]) and amyloid processing genes (e.g. *sel-12* [2.9]). At 8 hr post-exposure, these changes returned to base line or were decreased compared to controls. Metal-sensing genes (e.g. *aip-1* [0.54]) and heat shock

responses (e.g. *hsf-1* [0.46]) were among the genes with lower mRNA levels compared to controls at 8 hr post-exposure (Fig. 3).

In L4 stage animals, nPM exposure did not significantly alter the expression of most genes examined. However, the gene expression followed the trend observed in L1 exposed animals with increased expression at 1 hr post-exposure and return to baseline at 4 hr post-exposure (Fig. 3). nPM caused decrease of *tol-1* mRNA [-0.86] at this time. These data suggest that transcriptional responses to nPM exposure are dependent on developmental stage.

3.4 The role of *skn-1* transcriptional activity in response to nPM exposure.

Given the responsiveness of *gst-4* to nPM, we further studied the role of SKN-1 activity in nPM mediated toxicity. The expression of *skn-1* and its downstream genes was targeted by *skn-1* RNAi at L1, which blocked *gst-4* response in first 2 hours after nPM exposure (Fig. 4); *gst-4* was the only gene with significant nPM and the associated interaction with *skn-1* or time based on multivariate regression analysis. At 0 hr post-exposure, all SKN-1 targets were lower (*skn-1* [0.7], *gst-4* [0.58], *gcs-1* [0.47] and *ugt-11* [0.25]) vs negative control RNAi. Control animals had significant *gst-4* [1.56] mRNA increase at 0 hr post-exposure to nPM, which was not observed for *skn-1* RNAi. Contrary to *skn-1*-related genes after RNAi, nPM still induced some innate immune responses, e.g. *abf-2* [0.7 nPM/control at 0 hr post-exposure]. Moreover *skn-1* RNAi slightly increased *abf-2* response (+30%) at 2 hr post-exposure regardless of nPM. For the amyloid processing genes, *skn-1* RNAi decreased *sel-12* (-40%) with no further response to nPM at 0 hr post-exposure. At 2 hr post-exposure, nPM still decreased *sel-12* mRNA in worms fed Control RNAi (-20%), but not when fed *skn-1* RNAi. Taken together with the *gst-4::GFP* expression data, it appears that SKN-1 may have a role in mediating the transcriptional responses to nPM exposure.

Discussion

This study introduces *C. elegans* as a valuable short-lived model with potential for medium-throughput for genetic and toxicological studies of air pollution toxicity in humans. Most notable are the transcriptomic responses to nPM, a toxic subfraction of the ultrafine air pollution particulate matter. We identified several conserved genes that show developmental sensitivity to nPM. While nPM does not cause lethality in adult *C. elegans* up to 200 µg/ml, developmental exposure to nPM does reduce worm size and alter mRNA levels, dependent on SKN-1. Despite the observed reduction in worm size under nPM exposure, the time from which an animal hatches from an egg to laying its first egg is roughly the same (egg to egg time) (Fig. S3). As such, development is not markedly perturbed. Genes responsive to nPM include *skn-1*, the *C. elegans* homologue of the mammalian *NRF2*, which has fundamental roles in air pollution toxicity (9). In addition, 50 µg/ml of nPM given twice early in the *C. elegans* life cycle caused a mild increase in lifespan [1.2 days in mean lifespan], again dependent on SKN-1.

Although *C. elegans* lacks a respiratory system, exposing animals to cigarette smoke in smoking chambers for 3hrs impaired intestinal bacterial clearance and gene expression changes (32); we observe similar gene expression changes following nPM exposure (e.g. *tir-1*: -3.24, and *hso-16.2*: 2.14 fold change). In the insect *Drosophila*, chronic exposure to air pollution decreased lifespan by 50% (33), and a similar exposure chamber was also used for exposing *C. elegans* to cigarette smoke (32). Future studies using this methodology will allow comparisons between liquid and aerosolized delivery of nPM. It is possible that suspended particles could limit the toxic effects of nPM, and exposure to aerosolized particles should be considered for future experiments. Using lifelong liquid cultures to maintain worms could also be used to increase exposure of nPM.

While most collected nPM has a cell cytotoxicity 50 (CC50) of 10-20 µg/ml (9), our data show that there is no acute lethality to nPM even at concentrations as high as 200 µg/ml, which have no

obvious counterpart in human exposure. This lack of toxicity at high levels of nPM suggests that *C. elegans* may be highly resistant against nPM, through a currently unknown mechanism. It is possible that molecular defense pathways, such as those we observe, are important for this resistance, or perhaps its outer cuticle simply provides a strong barrier against the toxicity. While 200µg/ml is an extremely high dose compared to normal human exposure, it may be prudent in the future to test "mega-doses" (e.g. >500µg/ml) to arrive at a true LD50 in *C. elegans*. However, our collection protocol hits a technical limitation around 200µg/ml, and even with those concentrations, the volume becomes limiting in that it is difficult to do many follow-up experiments using the same nPM batch (as we have done here), requiring constant re-characterization of different nPM batches. We are interested in attempting higher concentrations to find a true LD50 if we find a way to make it technically feasible in the future.

Our study examined the role of *skn-1*/Nrf2 in the observed physiological and molecular changes of nPM. SKN-1/Nrf2 is a transcription factor well-known for affecting both development (31) and cytoprotection/detoxification (30), among other genes. Exposure of young adult mice (3 mo), but not older adult mice (18 mo), to nPM induced Nrf2-dependent phase II detoxifying enzymes such as GCLC and GCLM in lung, liver and brain samples (34). These age differences further show the different susceptibility of life stages to air pollution toxicity. Down-regulation of *skn-1*/Nrf2 in *C. elegans* did not cause lethality in response to nPM. However, *skn-1* knockout animals no longer had a reduction in size under nPM exposure (albeit were smaller overall). It is possible this developmental effect may involve *sel-12*, as *sel-12* mRNA was decreased by *skn-1* RNAi - similar to size reduction in these animals. In view of SKN-1 targets, nPM specifically activated *gst-4* with minimal change in *gcs-1* expression. Genome wide screening of the genes associated with *skn-1*-mediated detoxifying responses showed involvement of alternative pathways (e.g. *apb-2* and *csn-2*) in activation of *gcs-1*, regardless of *skn-1* activation (35). Thus *gcs-1* may be regulated independently of other phase 2 genes. It is worth noting that while animals

lacking *skn-1* mRNA transcripts experience embryonic lethality, it is unclear whether progeny that do survive the transgenerational RNAi exposure are "sicker" than wild type, or if their survival indicates that enough *skn-1* mRNA was around to reach a "survival threshold" where they are healthy enough (observationally, they do not seem "sick"). In addition, while we focused on the effects of *skn-1*, it is likely that other defense-associated transcription factors could be implicated in these gene expression changes or otherwise. While each individual experiment on *skn-1*-nPM interactions yielded modest, albeit significant changes, in sum, our data strongly suggests a role for *skn-1* under nPM exposure.

We must consider if *C. elegans* is an appropriate model for human air pollution toxicity, because humans have more complex circulatory and immune systems and 100-fold longer lifespans. Our previous studies of air pollution toxicity in mouse and cell culture models nonetheless show overlap in oxidative stress, inflammation and amyloidogenesis pathways with these *C. elegans* findings (5). This study investigates our previous findings of air pollution toxicity based on prior mouse and cell culture models. Specifically, we used TRAP ultrafine particulate matter (<0.2 μm dia.) in this study. UltrafinePM may be more toxic compared to PM_{2.5} (36, 37), but is not currently regulated or monitored by the United States Environmental Protection Agency.

Air pollution and oxidative stress are strongly associated (38-40). Previously, we showed only 5 hr exposure to air pollution sufficed for oxidative damage of membrane lipids in olfactory epithelium of exposed mice, assayed as 4-HNE (13). Oxidative stress (e.g. *gst-4*) was a prominent response to nPM. The other consistent air pollution response in cell cultures, mice, and humans is inflammation. Microarray analysis of the primary mixed glial culture responses to nPM showed MyD88 dependent activation of TLR4, suggesting this pathway as an important upstream sensors of air pollution leading to inflammation (41). Exposed mice had higher levels of brain TNF α depending on the duration of nPM exposure (8, 11). Humans exposed to diesel exhaust also show rapid systemic inflammatory responses

(42). Notably, TLR-associated gene homologues (*tol-1*: TLR4, *abf-2*: downstream of *tol-1*) were among the earliest responses of *C. elegans* to nPM.

Air pollution is recently recognized as an environmental risk factor for Alzheimer's disease (AD) and accelerated cognitive decline (5, 43). In mouse and cell models, exposure to nPM increased production of the A β peptide (5). *C. elegans* also shows similar effects of air pollution in the responsiveness of its homologous amyloid processing genes to nPM. Several amyloid-related genes are associated with *C. elegans* development; for example, inactivation of *apl-1/APP* results in penetrant lethality during the L1 to L2 transition due to molting defects (44). On the other hand, overexpression of *apl-1/APP* results in penetrant L1 lethality, shortened body length and morphological, locomotive, and reproductive effects (44, 45). *sel-12/PSEN* is one of the genes regulating *apl-1/APP* cleavage and trafficking (44). The expression of both *apl-1/APP* [0 hr: 0.64] and *sel-12/PSEN* [0 hr: 0.55, 2 hrs: 2.88] is significantly changed in L1 stage following nPM exposure, suggesting the importance of *apl-1* and *sel-12* expression for the developmental responses to nPM. In addition, air pollution is linked with developmental changes in humans; for example, childhood obesity is associated with maternal exposure to ambient air polycyclic aromatic hydrocarbons during pregnancy (46). Moreover, we showed that prenatal exposure of mice to nPM alters neuronal differentiation and depression-like responses (12). Lack of *sel-12* lead to elevated endoplasmic reticulum (ER) mitochondrial Ca²⁺ signaling and oxidative stress due to mitochondrial superoxide production (47). Thus, *sel-12/PSEN* might be a part of antioxidant response to nPM.

We initially hypothesized exposure to nPM might be fatal for *C. elegans*, especially during development. Instead, we found that short term exposure to air pollution could cause a hormetic increase of lifespan. This effect is apparently dose dependent, since exposing the worms for multiple times during development and reproduction ablated this effect of air pollution. Hormesis, wherein small

doses of a toxin or stress can induce increases in health and lifespan, is well established in eukaryotes (48), including *C. elegans* (49). While modest, our data suggest the hormetic effect of air pollution might be dependent on a functional *skn-1* response, indicating its importance on later lifespan in addition to its importance in physical development. If exposure to nPM is administered more chronically, perhaps by air exposure chambers, longer or more constant doses may eventually cause decreases in lifespan.

Future studies will further identify signaling pathways in responses to air pollution and hopefully help to identify drug and/or diet intervention strategies to counteract these ambient toxins. Systematic study of the life stages of *C. elegans* will identify critical periods of vulnerability to acute and chronic air pollution exposure. *C. elegans* could productively complement rodent models for studies of air pollution toxicity throughout the life cycle.

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Author contributions

Conceptualization, S.P.C., C.E.F., T.E.M.; Methodology, A.H., H.M.D. and S.P.C.; nPM collection and characterization: F.S., C.S.; Investigation, A.H., H.M.D., N.S. and S.P.C.; Writing, A.H., H.M.D. and S.P.C.; Supervision, Project Administration, and Funding Acquisition, S.P.C. and C.E.F.

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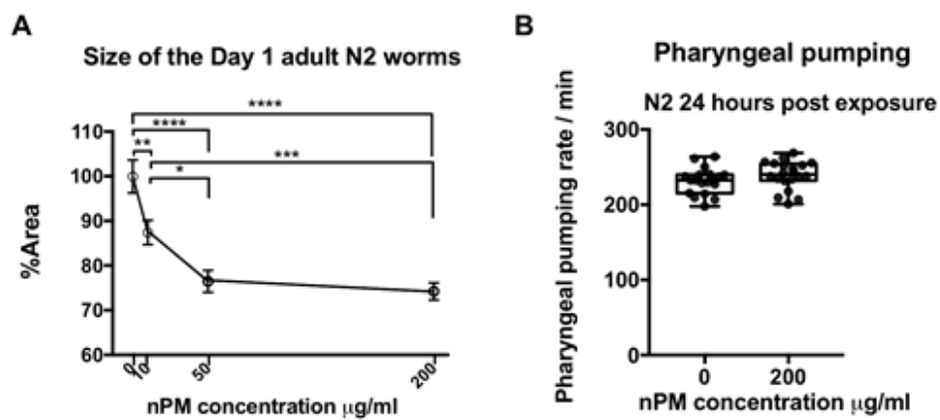
Figure 1. Acute nPM exposure of L1 worms results in adult worm size reduction. A) Dose dependent size changes (mean±SEM) of Day 1 adult animal body area following 1 hr exposure to nPM in L1 stage (n=3*15 to 22 /group). The average area of the control group was 0.06 mm² with average length of 0.97 mm. B) Pharyngeal pumping rate of animals (n=3*~7/group) analyzed at 24 hrs post-exposure of the L1 stage wildtype animals to 200 µg/ml nPM. Statistical tests include ANOVA followed by Tukey post hoc with correction for multiple statistical hypothesis testing for size and t test for pharyngeal pumping. Adjusted p-values: <0.05 (*), <0.01(**), <0.001(***), <0.0001(****).

Figure 2. nPM mediated SKN-1 response underlies developmental and lifespan effects of TRAP particulate matter. A) Dose response *gst-4* protein responses to nPM in L1 and L4 *gst-4p::GFP* strain (n=3×3 to 7/group). Representative images of L1 animals are presented in front of the graph. B) Examining the interaction of different life stages on nPM (50 µg/ml) mediated hormesis effects (n=3×52-101/group) and the role of *skn-1* response in this process until 90% survival in wild type non-exposed animals (n=2-3×47-53/group) (Summary statistics in Table S2). L: Larval stage, D: Day adult. Multivariate model: GST-4 ~ nPM×Life stage×Time post exposure, F=21.36, adjusted R²=0.42, p<0.0001****. (nPM):(Time) interaction, p=0.003**, β=-0.0002 C) Analysis of adult *skn-1(zu135)* mutants and wild type worms exposed to nPM during development (n=3×13 to 27/group). Multivariate model: size ~ *skn-1*×nPM, F=37.72, adjusted R²=0.35, p<0.0001****, *skn-1*:nPM interaction: p=0.03*, β=0.24. Statistical tests: Survival data was analyzed by Log-Rank and Wilcoxon tests. Size: Univariate ANOVA followed by Tukey post hoc with correction for multiple statistical hypothesis testing and also Multivariate ANOVA for interaction was used for size and *gst-4p::GFP* results. Adjusted p-values: <0.05 (*), <0.01(**), <0.001(***), <0.0001(****). Please refer to Table S3 for summary statistics of multivariate models.

Figure 3. mRNA response to nPM varies with developmental stage. A) Heatmap showing the mRNA changes of the selected genes in L1 and L4 stage animals exposed to 50 $\mu\text{g}/\text{ml}$ nPM for 1 hr (n=4-5 replicates, ~500 animals/replicate). RNA levels were followed for 8 hr post-exposure in L1 stage animals and 4 hr post-exposure in L4 stage animals. Responses are clustered based on Euclidian distance using complete linkage method. The heatmap is annotated with the time point, treatment, life stage, average of mRNA changes and the function of target genes. Significant changes are shown in the heatmap. B) mRNA changes of Alzheimer homologue genes in *C. elegans* to nPM. The t-test was used to compare nPM and controls at each time. p-values: <0.05 (*), <0.01(**), <0.001(***), <0.0001(****).

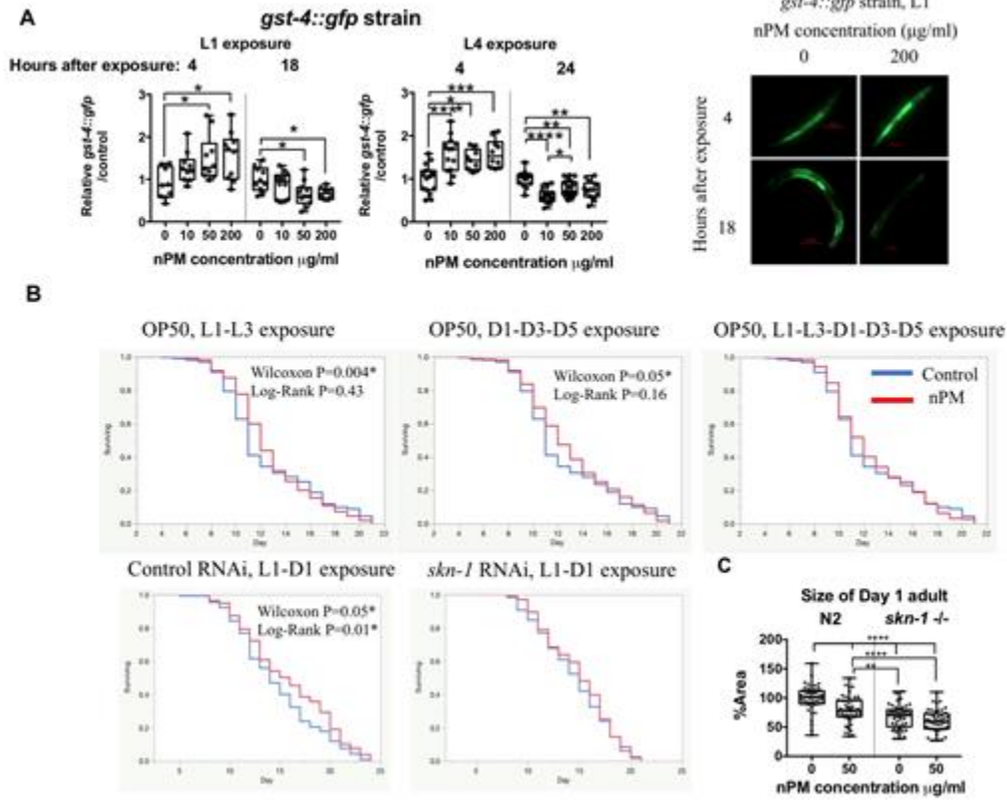
Figure 4. SKN-1 mediates some transcriptional responses of nPM. RNA responses to 1 hr nPM (50 $\mu\text{g}/\text{ml}$ nPM) after *skn-1* RNAi (n=3-5 replicates, ~500 animals/replicate). Pairwise t-test was used to compare Control RNAi vehicle controls with *skn-1* RNAi or nPM vs controls at each time. Multivariate coefficients of the *gst-4* expression regressed on nPM, *skn-1* RNAi, time and the associated interactions: *gst-4* was the only gene with significant nPM and nPM:*skn-1* interaction. p-values: <0.05 (*), <0.01(**), <0.001(***), <0.0001(****).

Figure 1



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Figure 2



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Figure 3

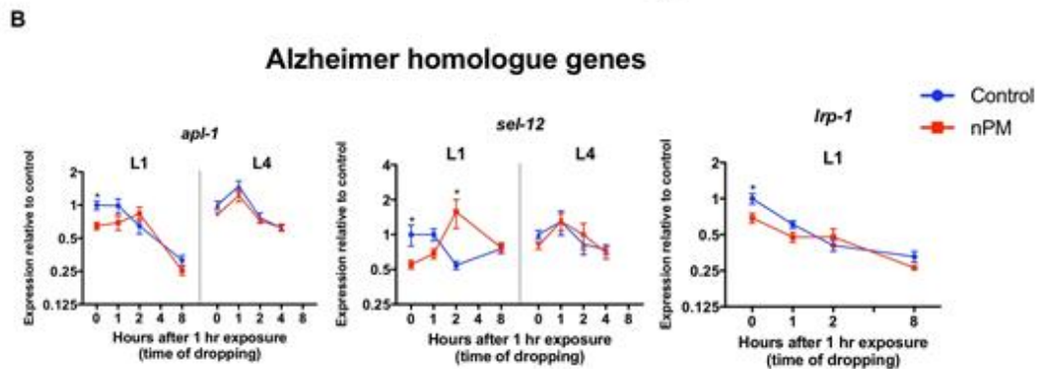
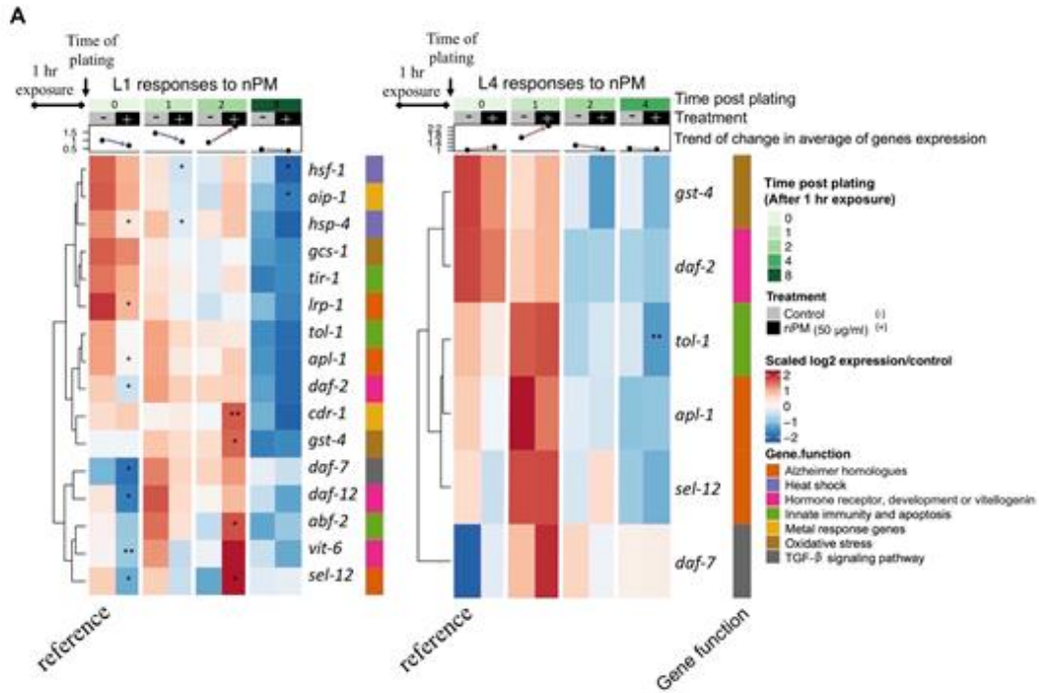
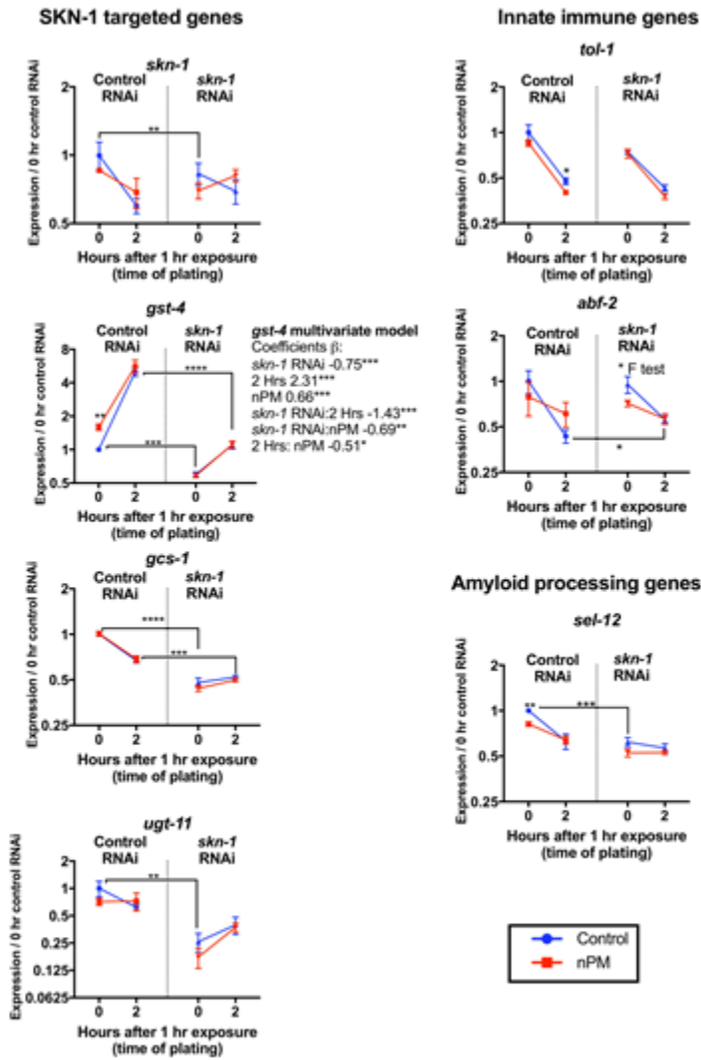


Figure 4



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