

Environmental Science Nano

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Nano impact

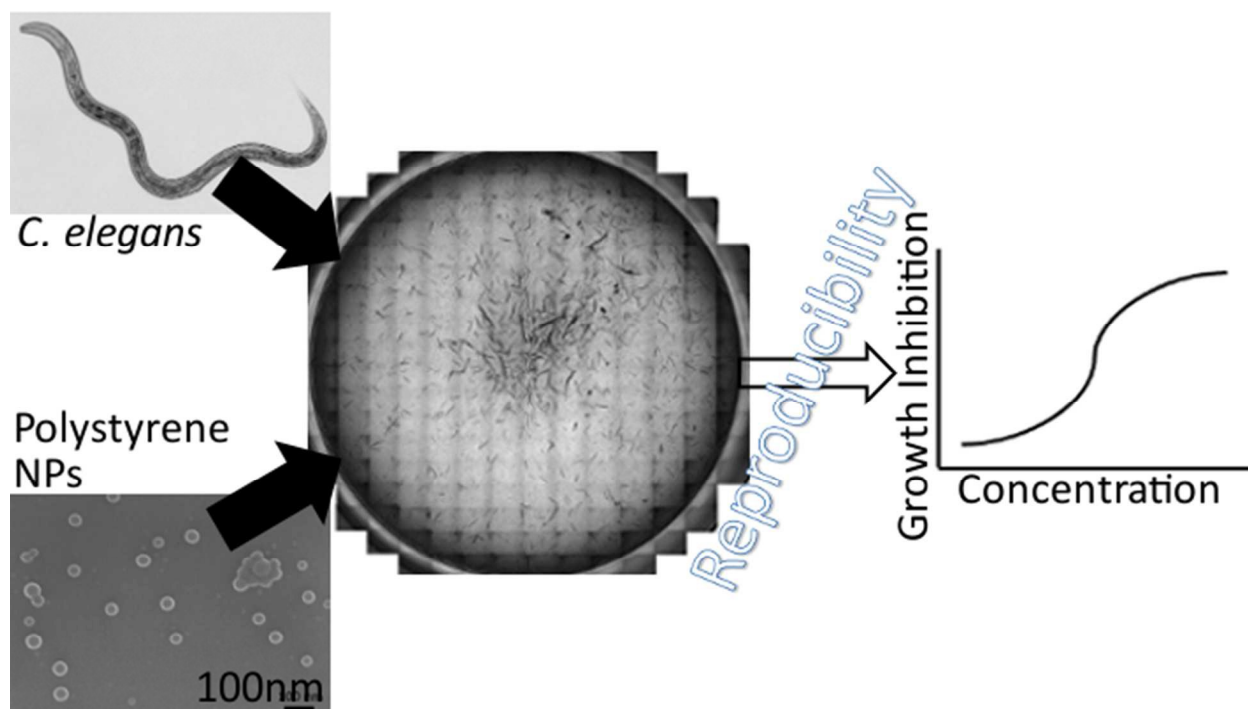
Increasing production and use of nanomaterials necessitates an understanding of their potential environmental and organismal impacts. The lack of standardized toxicity assays has hindered this understanding and led to contradictory and incomparable results. Here we evaluate the applicability of using a standard toxicity assay for *Caenorhabditis elegans* for a reference chemical control and positively charged polystyrene nanoparticles through sensitivity testing. We determined that the nanoparticles tested increase variability of our results and identified the parameters that can most strongly impact the uncertainty of the assay results.

1 Feasibility of using a standardized *Caenorhabditis elegans* toxicity test to assess nanomaterial
2 toxicity

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8 Abstract

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10 Increasing production and use of engineered nanomaterials (ENMs) has generated widespread
11 interest in measuring their environmental and human health effects. However, the lack of
12 standardized methods for these measurements has often led to contradictory results. Our goal
13 in this study was to examine the feasibility of using a standardized *Caenorhabditis elegans*
14 growth and reproduction based toxicity test designed for use with dissolved chemicals to assess
15 ENM toxicity. Sensitivity testing of seven key experimental factors identified by cause-and-
16 effect analysis revealed that bacterial feed density and plate shaking had significant effects on
17 growth inhibition by a reference toxicant, benzylcetyldimethylammonium chloride (BAC-C16).
18 Bacterial density was inversely proportional to experimental EC₅₀ values, while shaking the
19 plates during the assay caused a substantial decrease in nematode growth and reproduction in
20 control nematodes. Other factors such as bacterial viability, organism maintenance, and media

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4 21 type showed minimal effect on the test method. Using this assay with positively charged
5 22 polystyrene nanoparticles (PSNPs) revealed that the variability in the PSNP EC₅₀ values was
6 23 larger compared to those of BAC-C16. Additionally, while media type and bacterial viability did
7 24 not impact BAC-C16 toxicity, PSNP toxicity differed substantially when these parameters were
8 25 changed. PSNPs were more toxic in K⁺ medium and S-basal compared to M9 and feeding
9 26 nematodes with UV killed *E. coli* decreased toxicity of PSNPs. Test validity with ENMs and
10 27 modifications that can be made to adapt the standard *C. elegans* toxicity assay for use with
11 28 ENMs are discussed.

15 29 **Introduction**

16 30 Understanding the impact of increased production and use of engineered nanomaterials
17 31 (ENMs) on the environment and human health is essential for sustainable commercialization of
18 32 ENMs. Although researchers have been testing the hazards associated with ENMs for over a
19 33 decade,¹⁻³ the lack of standardized methods and difficulties associated with ecotoxicity testing
20 34 of ENMs has hindered these efforts and produced inconsistent results.^{4,5} For example, while
21 35 Ag is a known biocide, the impact of size of Ag ENMs on toxicity is unclear⁶ as are the
22 36 mechanisms associated with the ENM toxicity.⁷ A number of researchers attribute the toxic
23 37 effects of Ag ENMs to dissolved Ag⁸⁻¹⁰ while others show nanoparticle-specific effects.^{11,12}
24 38 Differing results among laboratories may be attributed to many factors such as differences in
25 39 protocols (i.e., ENM dispersion procedure, exposure duration), differences in ENMs of the same
26 40 elemental composition (lot to lot manufacturer variability, size, surface coating, impurities), and
27 41 improper controls leading to artifacts associated with ecotoxicity testing of the ENMs.¹³ The
28 42 choice of test organism and length of the assay also have a considerable impact on resulting
29 43 toxicity of ENMs. Very high concentrations (> 500 mg l⁻¹) of carbon nanotubes (CNTs) had little
30 44 to no effect on fish^{14,15} or amphibians,¹⁶ yet much lower concentrations (< 10 mg l⁻¹) showed
31 45 dose-dependent effects in copepods,¹⁷ *Daphnia magna*,^{18,19} and marine mussels.²⁰
32 46 Additionally, CNT concentrations causing chronic *D. magna* toxicity^{19,21} are over an order of
33 47 magnitude lower than those for short-term immobilization experiments.^{22,23} Chronic assays
34 48 that measure sub-lethal endpoints, which are more sensitive to lower, more environmentally
35 49 realistic concentrations, are needed to understand potential ecological impacts of ENMs.²⁴⁻²⁶
36 50 These chronic assays show effects at orders of magnitude lower concentrations than acute
37 51 assays.^{27,28}

38 52 The nematode *Caenorhabditis elegans* (*C. elegans*) is rapidly becoming a model for
39 53 environmental and developmental toxicological research.^{2,29-33} Nematodes are possibly the
40 54 most abundant multicellular organism on the planet and are important organisms in benthic
41 55 and soil food webs.^{34,35} Therefore, using them as model organisms in toxicological assays
42 56 would provide insight into the potential impacts of pollutants on the environment. Additionally,
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4 57 *C. elegans* is well suited to toxicity assays due to its ability to grow and reproduce in soil and
5 58 aqueous environments, ease of culture, and importance in soil food webs. In some cases,
6 59 toxicity results with *C. elegans* have even correlated well to endpoints in mouse and rat studies,
7 60^{36, 37} thus providing important data at a fraction of the effort and cost. *C. elegans* toxicity
8 61 measurements can provide a strategy for chemical or ENM screening² to facilitate tiered
9 62 testing approaches and minimize animal testing.

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13 63 The International Standards Organization (ISO) published a standard toxicity assay with *C.*
14 64 *elegans*³⁸ that allows the assessment of a potential toxin or contaminated environmental
15 65 sample on sublethal physiological parameters (growth and reproduction).³⁹ The method is
16 66 designed for use with dissolved chemicals or contaminated soils or sediments and guidance for
17 67 using this method with ENMs is not readily available. This is important because ENMs may need
18 68 to be dispersed prior to toxicity testing and may agglomerate, settle, and/or dissolve during the
19 69 test and substantially change their toxicity.^{1, 3, 40} Although the ISO method for *C. elegans*
20 70 toxicity testing is available, it appears to only have been used for testing FeOx⁴¹ and, with
21 71 modification, for testing TiO₂ ENMs.⁴² Other researchers have used a variety of non-
22 72 standardized methods that included different media, exposure time, feed density, and
23 73 endpoints.⁴³⁻⁴⁶ Unsurprisingly, the studies yielded different findings and the variable conditions
24 74 limit direct data comparison among laboratories. Even if the ISO standard method was used
25 75 more often, researchers who use the ISO method for nanoecotoxicity testing may make
26 76 different ENM-specific modifications, thus leading to additional variability in the results among
27 77 different laboratories.

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35 78 The purpose of this study was to assess the feasibility of using the *C. elegans* ISO method with
36 79 ENMs by determining the impact of test modifications, that may be required for ENMs, on the
37 80 toxicity results. In order to assess what steps of the ISO protocol may have the largest impact
38 81 on the assay results, a cause-and-effect (C&E) analysis was performed.⁴⁷ This approach, which
39 82 is a thought exercise originally developed for quality manufacturing and used for propagation of
40 83 error analysis in analytical chemistry, identifies the expected major “causes” of variability and
41 84 their “effect” on the assay result. This information was then used to conduct sensitivity testing
42 85 of the ISO method by modifying various parameters of the protocol and determining how they
43 86 impacted the toxicity results for a positive control. Our hypothesis was that this approach could
44 87 provide information on the robustness of the test procedure and reveal which assay steps have
45 88 the greatest impact on the variability of the results. These steps could then be modified to
46 89 decrease variability or control measurements could be incorporated to more carefully track
47 90 these sources of variability. After the sensitivity testing, a plate layout with multiple control
48 91 measurements was developed and used to test the toxicity of a model ENM: positively charged
49 92 polystyrene nanoparticles (PSNPs). PSNPs were selected because they could be purchased in a
50 93 highly concentrated, stable suspension and their toxicity was hypothesized to be predominantly

94 attributable to the positively charged surface coating on the ENMs and not to dissolved ions or
95 other non-specific nano effects. We then performed similar sensitivity testing with PSNPs to
96 identify possible artifacts of the assay and evaluate the impacts of changing different
97 parameters on the assay results with an ENM.

98 **Methods**

99 Overview of *C. elegans* toxicity assay

100 The duration of the ISO *C. elegans* assay is 96 h, during which time juvenile *C. elegans* mature
101 into adult hermaphrodites, become gravid, lay eggs, and the eggs hatch into new juveniles. The
102 test includes a bactericidal positive control, benzylcetyldimethylammonium chloride (BAC-C16),
103 for which preliminary EC₅₀ values for growth and reproduction have been established. The
104 nematodes are fed *Escherichia coli* (OP 50 strain) in a defined bacterial density during the assay.
105 Procedures for the toxicity assay were followed as detailed in ISO 10872³⁸ and described in the
106 supplementary information (Fig. S1). Briefly, 10 first stage juvenile (J1) *C. elegans* were added
107 to each well of a 12 well plate along with the test material and *E. coli* as a food source. The
108 plate was incubated at 20 °C for 96 h, at which point the nematodes were killed by heating and
109 stained. In order to measure nematode length and to determine the number of new young
110 nematodes, the plates were imaged using light microscopy. Quantitative assessment of growth
111 and reproduction inhibition were determined based on nematodes in control wells without
112 added toxicants.

113 Cause-and-effect analysis

114 A C&E analysis was conducted to identify which experimental factors may impact the test
115 results by examining each step of the assay and determining how alterations of the protocol
116 (that may be helpful for ENM stability) may impact results. The resulting output is a C&E
117 diagram with main branches that depict steps that are major sources of variability and side
118 branches that describe sources of variability that contribute to the main branches (Fig. 1 and
119 Table S1). Information from this analysis was then used to design experiments that
120 implemented sensitivity testing for quantifying the variability from the components of the C&E
121 diagram.

122 Nematode cultures

123 Wild type, Bristol strain N2 *C. elegans* were obtained from the Caenorhabditis Genetics Center
124 (CGC, University of Minnesota) and maintained on nematode growth medium (NGM) with *E.*
125 *coli* as feed. The nematodes were allowed to starve and become dauer larvae, which can
126 survive for several months without feed. First stage juvenile (J1) *C. elegans* were obtained by

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4 127 plating dauer larvae on a fresh *E. coli* lawn grown overnight on a NGM plate at 37°C and
5 128 allowing the plate to incubate at 20 °C for 72 h.

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7 129 Test material preparation

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10 130 Polystyrene nanoparticles (PSNPs) were purchased from Bangs Laboratories (Bangs
11 131 Laboratories Inc., Fishers, IN, USA) and were described as amine coated with a diameter of 58
12 132 nm. Primary particle size of 200 particles was measured via scanning electron microscopy using
13 133 a Zeiss NVision 40 (Zeiss International, Oberkochen, Germany) focused ion beam/scanning
14 134 electron microscope operating at 15 kV, and was determined to be 51 nm ± 9 nm (Fig. S2). To
15 135 understand the behavior of ENMs in the test media across the duration of the study, a 100 mg l⁻¹
16 136 suspension of PSNPs was prepared in 50 % S basal, K⁺ medium, and M9 (media composition
17 137 described in SI) and the particle size was measured via dynamic light scattering (ZetaPALS Zeta
18 138 Potential Analyzer, Brookhaven Instruments Corporation, Holtsville, NY) immediately and after
19 139 the suspension was placed into a 20 °C incubator after 96 h. BAC-C16 (Acros Organics (97 %
20 140 pure), Geel, Belgium) and PSNPs were prepared by diluting the samples in ultrapure water to
21 141 produce concentrations twice as much as those used in the assay. The water was vortexed
22 142 during addition of PSNPs to help maintain stability of the suspension.⁴⁸ Although the ISO
23 143 standard suggests testing only a concentration of 15 mg l⁻¹ of BAC-C16 for routine analysis, a
24 144 range of concentrations from 3.5 mg l⁻¹ to 40.5 mg l⁻¹ were tested to produce an EC₅₀ value
25 145 during each experiment. To ensure that the PSNP coating or any other dissolved component
26 146 that may have remained from synthesis was not causing a toxic effect, an 800 mg l⁻¹ PSNP
27 147 suspension was prepared in water, allowed to settle for 2 h, and then passed through a 0.02 μm
28 148 filter. The resulting solution was used as an ENM filtrate control in the toxicity assay.^{13,22} In
29 149 order to test all of the concentrations of BAC-C16 as well as PSNPs and controls associated with
30 150 the PSNPs, a novel plate design was implemented as illustrated in Fig. S3.

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33 151 *E. coli* preparation

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43 152 A suspension of *E. coli* was prepared prior to the toxicity assay by inoculating 1 l of sterilized
44 153 Luria Broth (Miller's LB broth base, Invitrogen, Thermo Fisher Scientific) with 100 μl of a frozen
45 154 culture of *E. coli*. This culture was set on a shaker incubator at 37 °C and 15.7 rad s⁻¹ (150 rpm)
46 155 for 17 h. The culture was then transferred into 250 ml polypropylene centrifuge tubes and spun
47 156 at 2000 x g (Allegra 25R Centrifuge, Beckman Coulter, Fullerton, CA, USA) and 4 °C for 20 min.
48 157 The supernatant was decanted and the bacteria pellets resuspended into 50 ml of M9. This was
49 158 repeated two more times. This suspension was then diluted in M9 and measured using a
50 159 turbidity meter (HI 88713, Hanna Instruments, Woonsocket, RI, USA), which was calibrated
51 160 using four formazin suspensions ranging from 15 formazin absorbance units (FAU) to 2000 FAU.
52 161 ⁴⁹ This calibration curve enabled the calculation of the dilution needed to obtain the specified *E.*
53 162 *coli* concentration of 1000 FAU set forth in the ISO *C. elegans* protocol.⁴⁹ A 5 mg ml⁻¹ solution of

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3 163 cholesterol (NIST SRM 911c) dissolved in 100 % ethanol was then added to the feed suspension
4 164 to achieve a 0.2 % v/v concentration of cholesterol.

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7 165 Toxicity assay

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9 166 The test wells were prepared by adding 500 μl of the test material and 500 μl of the *E. coli*
10 167 suspension to each well. Two methods were used to obtain J1 nematodes for the toxicity tests,
11 168 either a filtering method or a bleaching method. For the filtering method specified in ISO
12 169 10872, nematodes were washed from the culture plates onto a 5 μm polyester mesh filter
13 170 (Hepfinger, Munich, Germany) using 8 ml of M9. The filtrate contained mainly J1 nematodes,
14 171 however, second stage juveniles (J2) nematodes were also present. To avoid this, nematodes
15 172 were also synchronized using a standard bleaching protocol adapted from⁵⁰ in which a mixed
16 173 culture of nematodes were exposed to a sodium hypochlorite and sodium hydroxide solution
17 174 for 10 minutes, washed with sterile water three times and the eggs were allowed to hatch in
18 175 sterile water overnight. Bleached nematodes were only J1 stage as development is arrested
19 176 with no food present. Ten J1 nematodes were added to each well and the test was initiated by
20 177 placing the plates into a 20 °C incubator, in the dark, and leaving them undisturbed for 96 h. All
21 178 J1 nematodes not used in the test were stained with Rose Bengal (500 μl of a 300 mg l^{-1} stock
22 179 added to 5 ml), heated at 80 °C for 10 min to kill and straighten them, 30 individuals were
23 180 measured, as described in the SI, to determine the initial nematode length. At the end of the
24 181 test, 200 μl of a 300 mg l^{-1} stock of Rose Bengal was added to each well and the plate was
25 182 heated at 80 °C for 10 min to kill and straighten all of the nematodes. The plate was allowed to
26 183 cool for at least 1 h prior to imaging. All plates were stored at 4 °C and imaged within one week
27 184 after the experiment concluded. Imaging details can be found in the SI. After imaging, total
28 185 length of adult nematodes was measured and young were counted. Reproductive counts are
29 186 expressed as young per adult hermaphrodite.

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41 187 Sensitivity testing

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43 188 To experimentally determine which of the six parameters identified in the C&E analysis (Table
44 189 S1) had the greatest impact on the assay results, the identified parameters were adjusted and
45 190 were compared to those obtained from the conditions indicated in the ISO standard. We tested
46 191 effects of 1) the type of culture matrix from which nematodes were harvested (plate versus
47 192 liquid culture), 2) two different manufacturers of BAC-C16 (Acros Organics (97 % pure), Geel,
48 193 Belgium, was mainly used in the study, and compared to Alfa Aesar Ward Hill (95 % pure), MA,
49 194 USA), 3) different assay media (S-basal medium, K^+ medium, and M9 medium), 4) the viability of
50 195 the *E. coli* feed, 5) the size of wells (12 well and 24 well plate) 6) the feed density during the
51 196 assay, and 7) whether the plates were shaken during the assay. Details of each test can be
52 197 found in the SI.

198 Data analysis

199 Mean growth of nematodes in each well was calculated by subtracting the mean length of adult
200 hermaphrodites by the mean length of J1 nematodes measured at the start of the assay.

201 Inhibition of growth (G_I) was calculated for each nematode as follows:

$$G_I = 100 - \frac{L_F - L_I}{G} * 100$$

202 where L_F is final length of the individual nematode at the end of the assay, L_I is the mean initial
203 J1 length at the start of the assay, and G is the mean growth of the control nematodes during
204 the assay. Inhibition of reproduction (R_I) was calculated for each well as follows:

$$R_I = \frac{R_C - R_W}{R_C} * 100$$

205 where R_C is the mean reproduction per adult hermaphrodite found for the control wells and R_W
206 is the reproduction per adult hermaphrodite found in the test well. EC_{50} for growth and
207 reproduction was determined using a four parameter logistic function in GraphPad Prism (V
208 6.04, GraphPad Software, Inc).

209 Results and Discussion

210 C&E analysis

211 The C&E analysis provided a framework for identifying the factors of the assay that may cause
212 the greatest variability or uncertainty in the assay measurements. Six main categories were
213 identified: organism maintenance, the reference chemical, bacteria, the assay protocol,
214 microscopy, and ENM specific issues (Fig. 1 and Table S1). These branches include the
215 parameters outlined in the ISO protocol and potential modifications of the protocol for use with
216 ENMs. Branch 1 concerns the culturing procedure for *C. elegans*. The nematodes can be
217 cultured on agar plates containing a bacterial lawn or in liquid culture. The most common liquid
218 culture is S-basal media. However, nematodes cultured in liquid media are longer and thinner
219 than those from agar plates⁵¹ and it is unknown if the culturing procedure impacts the
220 response of juveniles that are harvested for the exposure to toxicants. Branch 2 identifies
221 sources of variability related to the reference chemical, BAC-C16. BAC-C16, which is not easily
222 quantified, is difficult to dissolve in water, and little is known about its stability over time.
223 Therefore, there may be significant variability in the chemical toxicity among the batches from
224 suppliers and how well the chemical dissolves in water, all of which may impact *C. elegans*
225 growth inhibition results. Branch 3 concerns the *E. coli* density used in the assay, which is
226 difficult to measure accurately and could increase the assay variability as it may change during
227 the assay as a result of bacterial growth and interactions with the toxin. Branch 4 identifies

228 several factors in the assay protocol described in the ISO document, as well as adaptations for
229 testing ENMs that may help keep the ENMs suspended. For example, different media
230 preparations or incorporating plate shaking during the assay may impact the ENM suspension
231 and the assay results. Branch 5 encompasses procedures for imaging nematodes for growth
232 measurements. Sources of variability in this branch are associated with microscope calibration,
233 nematode identification due to focus artifacts, interference from *E. coli* or debris, and user-to-
234 user variability. Branch 6 catalogs ENM specific concerns included producing a reproducible
235 dispersion, changes to the ENM during the assay (e.g., settling, dissolution), and toxicity of the
236 ENM to the bacteria or interactions of ENM with bacteria such as heteroagglomeration.

237 Sensitivity testing with the BAC-C16 reference toxicant

238 The sensitivity testing of the ISO protocol yielded important insights regarding the parameters
239 that impact the assay results (Fig. 2A-G and Fig. S4A-G). We found that the type of culture from
240 which nematodes were harvested, the reference chemical vendor, the media that the assay
241 was performed in, the *E. coli* viability, and the size of wells had minimal effect on the assay
242 results with BAC-C16, as shown in Fig. 2A-E for growth, and Fig S4A-E for reproduction. The lack
243 of impact on toxicity observed with BAC-C16 in different media may give researchers more
244 flexibility to select which media works best for the ENM they are using. Media flexibility allows
245 for use of more environmentally relevant media such as simulated porewater.⁵² While the
246 media we used in this study are some of the more commonly used in the literature, other
247 media have shown differences in growth of nematodes.⁵³ Therefore, measuring control
248 nematode growth in the chosen media is vital to understanding the potential impacts of a toxin.
249 Our results suggest that the protocol described in the original ISO document is robust to media
250 composition changes that were tested here. Changes in the nematode sensitivity to BAC-C16
251 was minimal when the nematodes were fed UV treated bacteria instead of live bacteria (Figure
252 2D and Fig. S4D). The EC₅₀ for growth of BAC-C16 was 17.85 mg l⁻¹ (95 % CI: 17.15 mg l⁻¹ to
253 18.57 mg l⁻¹) for nematodes fed live bacteria and 14.87 mg l⁻¹ (95 % CI: 14.17 mg l⁻¹ to 15.61 mg
254 l⁻¹) for nematodes fed UV treated bacteria. The EC50 for reproduction of BAC-C16 was 11.13 mg
255 l⁻¹ (95 % CI: 10.16 mg l⁻¹ to 12.19 mg l⁻¹) for nematodes fed live bacteria and 10.33 mg l⁻¹ (95 %
256 CI: 0.06 mg l⁻¹ to 1690 mg l⁻¹) for nematodes fed UV treated bacteria. Note the extremely high
257 variability found for reproduction, especially for the nematodes fed UV treated bacteria.
258 However, it is unclear if other researchers found impacts of bacterial viability on nematode
259 growth. While we observed minimal impact of *E. coli* inactivation on growth of *C. elegans*
260 similar to other studies,^{46, 54} several studies have found that life span of *C. elegans* is increased
261 when fed with growth inhibited or dead *E. coli*.^{55, 56}

262 Of the parameters tested, changes in the *E. coli* feed density and plate shaking had the largest
263 impact on assay results for BAC-C16. The feed density had a large impact on toxicity of BAC-

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3 264 C16. At 100 FAU feed density levels, 15 mg l⁻¹ of BAC-C16 completely inhibited nematode
4 growth, while at 1100 FAU feed density, 15 mg l⁻¹ of BAC-C16 did not affect growth (Fig. 2F).
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6 266 Similarly, nematodes did not reproduce below 500 FAU feed but reproduced as much if not
7 more than the control at ≥ 900 FAU (Fig. S4F). This result indicates that the assay positive
8 267 control is highly sensitive to feed density. Höss et. al.⁵⁷ found a similar result with Cd exposure
9 268 and suggested that binding of Cd to *E. coli* cells may impact bioavailability of the metal.
10 269 However, the method by which researchers measure bacterial density may impact the amount
11 270 of feed being administered. Bacterial density measurements were conducted in this study using
12 271 a turbidity meter as described in ISO method 10872,³⁸ but researchers use different methods
13 272 to quantify bacterial density such as photometers,^{47, 51} plate readers,⁵² or simply specify a
14 273 wavelength with no indication of instrumentation.^{35, 37, 53} The impact of using different
15 274 instruments to quantify bacteria densities is unclear, but our sensitivity testing suggests that a
16 275 50 FAU difference in bacterial concentration can change growth inhibition of BAC-C16 by as
17 276 much as 19 % (Fig. 2F). It is unclear if this is due to the nematodes having access to different
18 277 amounts of feed or if increasing amounts of *E. coli* decreases the availability of BAC-C16 to the
19 278 nematodes. Distinguishing the direct toxic effect of a chemical on growth inhibition from an
20 279 indirect effect on bacterial concentration is not possible with the current ISO method.
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22 281 Shaking plates during the assay decreased control growth by >300 μm (approximately 19 %
23 282 decrease) after 96 h and increased inhibition of growth of 15 mg l⁻¹ BAC-C16 by >36 %
24 283 compared to not shaking the plates (Fig. 2G). Similarly, shaking plates reduced reproductive
25 284 output by approximately 70% compared to not shaking (Fig. S4G). Shaking the plates greatly
26 285 increased test variability; the mean EC₅₀ (± 1 SD) for growth with plate shaking was 20.5 ± 13.1
27 286 mg l⁻¹ (n=16), yet decreased to 18.7 ± 2.6 mg l⁻¹ (n=16) without shaking. While leaving the plates
28 287 undisturbed during the assay may allow nematodes easier access to settled *E. coli*, aggregated
29 288 ENMs may also settle on the bottom of the wells, potentially increasing exposure of the
30 289 nematodes to the ENMs as well as changing exposure from ENMs to aggregates of these ENMs
31 290⁵⁸. It is important to consider these features of the test system when interpreting the results of
32 291 the assay with test ENMs. For example, choosing a media that minimizes aggregation, if
33 292 possible, would help to alleviate this issue.

34 293 Sensitivity testing with PSNPs and comparison to reference chemical results

35 294 Based on the results of the C&E and sensitivity testing with BAC-C16, we designed the layout of
36 295 12 well plates that provide five control features to assess the quality of the results (Fig. S3).
37 296 There are multiple advantages of testing the EC₅₀ value of the reference chemical and ENM on
38 297 each of three plates including that the plate-to-plate variability can be quantified and that the
39 298 EC₅₀ values for the BAC-C16 need to be within benchmark specifications for the ENM result to
40 299 be considered valid. For our laboratory, the mean EC₅₀ (± 1 SD) of BAC-C16 was 18.7 ± 2.6 mg l⁻¹

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3 300 (n = 16) and ranged from 14.4 mg l⁻¹ to 22.3 mg l⁻¹ (Fig. 3A). Mean inhibition of growth at 15 mg
4 301 l⁻¹ BAC-C16 was 34.1 ± 12.5 % and ranged from 18.1 to 58.7 % (Fig. 3B). While our growth
5 302 inhibition results were mainly within the 20 % to 80 % requirement as stated in ISO 10872,³⁸
6 303 several of our tests showed < 20 % inhibition at 15 mg l⁻¹ BAC-C16. However, an interlaboratory
7 304 study among eight laboratories showed a range of EC₅₀ values for growth from 11.9 to 18.9 mg
8 305 l⁻¹,³⁹ suggesting that our variability is similar to those in the interlaboratory study. This chemical
9 306 control and specification can be used to qualify the robustness of the measurement process.

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14 307 The results from conducting the assay with PSNPs three separate times (Fig. 4A) indicated that
15 308 the mean EC₅₀ for growth was 71.7 ± 37.2 mg l⁻¹ and ranged from 42.7 mg l⁻¹ to 113.7 mg l⁻¹. The
16 309 mean EC₅₀ for reproduction was 21.4 ± 10.5 mg l⁻¹ and ranged from 10.0 mg l⁻¹ to 30.7 mg l⁻¹.
17 310 While no published research has investigated the toxicity of PSNPs on *C. elegans*, cellular
18 311 toxicity assays indicate almost an order of magnitude lower EC₅₀ values than those observed for
19 312 *C. elegans*.⁴⁷ Several concentrations of BAC-C16 were tested in the same plates as the PSNPs to
20 313 compare the variability between the two substances (Fig. 4B). The coefficient of variations of
21 314 the growth EC₅₀ values for three independent assays were 9 % and 52 % for the BAC-16 and
22 315 PSNPs, respectively, indicating that the EC₅₀ values were substantially more variable for PSNPs.
23 316 No inhibition of growth was observed in the ENM filtrate control, suggesting that no leaching of
24 317 toxic chemicals from the ENM occurred. However, there were differences in *E. coli*
25 318 agglomeration in the presence of PSNPs and large *E. coli* agglomerates formed almost
26 319 immediately after addition (Fig. S5B). This observation suggests an additional indirect toxicity
27 320 mechanism (i.e., heteroagglomeration) that should be considered when testing ENMs. It is not
28 321 clear if the observed toxicity is due to the ENM or due to a secondary effect that results from
29 322 the ENM interaction with *E. coli* feed. For example, it is possible that the aggregates may change
30 323 the availability of feed to the nematodes. Experiments to further dissect the observed
31 324 nematode toxicity will be explored in a subsequent study.

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41 325 Unlike results for BAC-C16, media composition had a strong influence on the toxicity of PSNPs
42 326 (Fig. 5A and Fig. S6A), suggesting that care must be taken to understand the behaviors of the
43 327 ENM in the system. Growth EC₅₀ values for the three media were 23.7 mg l⁻¹ (95 % CI: 21.4 mg l⁻¹
44 328 to 26.2 mg l⁻¹), 5.9 mg l⁻¹ (95 % CI: 5.5 mg l⁻¹ to 6.3 mg l⁻¹), and 8.8 mg l⁻¹ (95 % CI: 8.1 mg l⁻¹ to
45 329 9.5 mg l⁻¹) for M9, K⁺ medium, and S-basal respectively. For reproduction, EC₅₀ values could not
46 330 be calculated for M9 due to high variability but were 2.7 mg l⁻¹ (95 % CI: 2.4 mg l⁻¹ to 3.1 mg l⁻¹),
47 331 and 2.8 mg l⁻¹ (95 % CI: 2.7 mg l⁻¹ to 3.0 mg l⁻¹) for K⁺ medium and S-basal respectively; while
48 332 these experiments were repeated at least twice, conducting the experiments with a different
49 333 concentration test range may have enabled the calculation of reproduction EC₅₀ values but was
50 334 not tested. While K⁺ medium has the lowest ionic strength and S-basal had the highest ionic
51 335 strength of the three media we tested, PSNPs were least toxic in M9. This may be due to the
52 336 fact that K⁺ medium and S-basal contain two different types of divalent cations, Ca²⁺ and Mg²⁺,

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3 337 while M9 contains only Mg^{2+} . As previously reported, the presence of divalent cations can
4 338 potentiate ENM aggregation in liquid media,^{59, 60} which may impact toxicity. Immediately after
5 339 addition, PSNPs agglomerated in S-basal (mean \pm SD: 1117.8 nm \pm 15.2 nm) and M9 (199.4 nm
6 340 \pm 4.0 nm) but not in K^+ medium (64.2 nm \pm 0.5 nm). After 96 h PSNPs increased in size in S-basal
7 341 (1966.3 nm \pm 512.3 nm) and M9 (649.6 nm \pm 14.0 nm) but remained similar in K^+ medium (58.4
8 342 nm \pm 0.5 nm). However, these measurements were run without *E. coli* present, the presence of
9 343 which may impact PSNP agglomeration. The media composition may be highly relevant for
10 344 other ENMs such as Ag ENMs which react readily with chloride;⁶¹ a media without chloride
11 345 salts may be needed to obtain the lowest EC_{50} values for Ag ENMs but such a media would have
12 346 lower environmental relevance.⁵² Similarly, bacteria viability influenced PSNP toxicity (Fig. 5B
13 347 and Fig. S6B), but did not impact the toxicity of BAC-C16. EC_{50} values for growth were 38.1 mg l⁻¹
14 348 (95% CI: 30.5 mg l⁻¹ to 47.6 mg l⁻¹) for nematodes fed live *E. coli* and 45.4 mg l⁻¹ (95% CI: 32.5
15 349 mg l⁻¹ to 63.4 mg l⁻¹) for those fed UV killed *E. coli*. EC_{50} values for reproduction could not be
16 350 calculated due to high variability. UV killed bacteria decreased PSNP toxicity, suggesting that
17 351 the interaction between PSNPs and *E. coli* may be hindered when bacteria are UV killed. This
18 352 suggests a potential assay modification to avoid this artifact. Similar to that of BAC-C16 results,
19 353 feed density greatly impacted toxicity of PSNPs (Fig. 5C and Fig. S6C). At 70 mg l⁻¹ PSNPs
20 354 nematode growth was similar to that of the control when feed was increased to 1100 FAU but
21 355 minimal to no growth was observed for feed densities between 100 and 550 FAU. Similarly, no
22 356 reproduction was observed at 70 mg l⁻¹ PSNPs until feed was increased to 900 FAU and at 1100
23 357 FAU, reproduction was similar to that of the control (Fig. S6C). Variability for all PSNP assays
24 358 was increased compared to data for BAC-C16.

359 Conclusion

360 This paper describes a process to assess the robustness and reproducibility of an ISO *C. elegans*
361 ecotoxicity assay and the utility of this assay for testing the potential effects of ENMs. Our
362 cause-and-effect analysis followed by a sensitivity testing revealed that *E. coli* concentration
363 and plate shaking have a large impact on nematode growth and toxicity of the control toxicant
364 BAC-C16. Lastly, we found that *E. coli* concentration, bacterial viability, and media composition
365 impacted PSNPs toxicity to *C. elegans*, illustrating the need to understand how ENM toxicity is
366 impacted by assay parameters. Given that studies in the literature often use a range of *E. coli*
367 concentrations and media compositions, the impact of these parameters should be better
368 understood using a broader range of conditions (bacteria concentrations, media, and types of
369 nanoparticles) to elucidate how data from multiple studies can be combined for environmental
370 risk assessment. In addition, the development of a more precise and robust method for
371 quantifying the bacteria concentration could help decrease the variability of the assay.

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3 372 While our findings with PSNPs illustrate the need to better understand the main factors
4 373 contributing to variability in assays when including ENMs, further experiments are needed to
5 374 better understand the robustness of the assay for use with varying nanoparticles (e.g., with
6 375 different surface coatings or sizes), because there may be biases or artifacts in the assay that
7 376 were not uncovered by testing only a single nanoparticle. Based on the findings of this study,
8 377 our recommendation is for this standardized method to be used more broadly in the
9 378 nanotoxicology literature. If modifications are made to the assay for which the assay is sensitive
10 379 (e.g., lower bacteria concentrations of different test media), it would be helpful to enable data
11 380 comparability among laboratories to also test the ENP using the conditions described in the ISO
12 381 assay. However, additional testing of the robustness of this assay with different ENPs may
13 382 reveal other important biases or limits to the applicability of this assay which should also be
14 383 taken into consideration.

15 384 The use of quality tools such as the cause-and-effect diagram and sensitivity testing allowed us
16 385 to systematically identify the parameters of the nematode culturing and toxicity assay that had
17 386 the greatest impact on assay results. This process can aid in reducing variability and increasing
18 387 reliability of standardized ecotoxicity tests and other key environmental measurements.

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27 394 recommendation or endorsement by the National Institute of Standards and Technology, nor
28 395 does it imply that it is necessarily the best available for the purpose.

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503 Figure Captions

504 Figure 1. Cause-and-effect analysis of ISO 10872 protocol. The six main branches indicate the
505 factors that we have identified that have the greatest potential to cause variability in assay
506 results. For detailed descriptions see Table S1.

507 Figure 2. Sensitivity testing of ISO 10872 conducted by altering test conditions (shown in Figure
508 1) and comparing the outcome to the original protocol. The test parameters altered were (A)
509 the culture from which the nematodes were harvested for the assay, (B) the manufacturer of
510 the positive control BAC C16, (C) the media that the test was performed in, (D) bacterial
511 viability, (E) the assay performed in a 24 well plate instead of 12 well (F) the amount of feed
512 used in the assay (all exposures include 15 mg l⁻¹ BAC-C16), and (G) whether the plates were
513 shaken or left undisturbed. For each plot, growth data shown are mean ± one standard
514 deviation, n=3 for each data point.

515 Figure 3. Control charting of EC₅₀ values (A) and inhibition of growth at 15 mg l⁻¹ (B) of *C.*
516 *elegans* exposed to BAC-C16 in 96 h standard toxicity assays conducted over several months
517 based on ISO 10872. Data presented as mean ± one standard deviation. The vertical bars
518 represent the date we stopped shaking plates during the assays.

519 Figure 4. Variability of the adapted toxicity assay for growth inhibition of A) PSNPs and B) BAC-
520 C16 conducted on three different days. Data are shown as mean ± one standard deviation. N =
521 3 wells, each with 10 nematodes.

522 Figure 5. Sensitivity testing of ISO 10872 containing PSNPs. The test parameters altered were
523 (A) The feed density, (B) the media that the test was performed in, and (C) the bacterial density.
524 Growth data presented as mean ± one standard deviation. For each experiment N = 3 wells,
525 each with 10 nematodes. Experiments were performed twice and data are combined.

Figure 1.

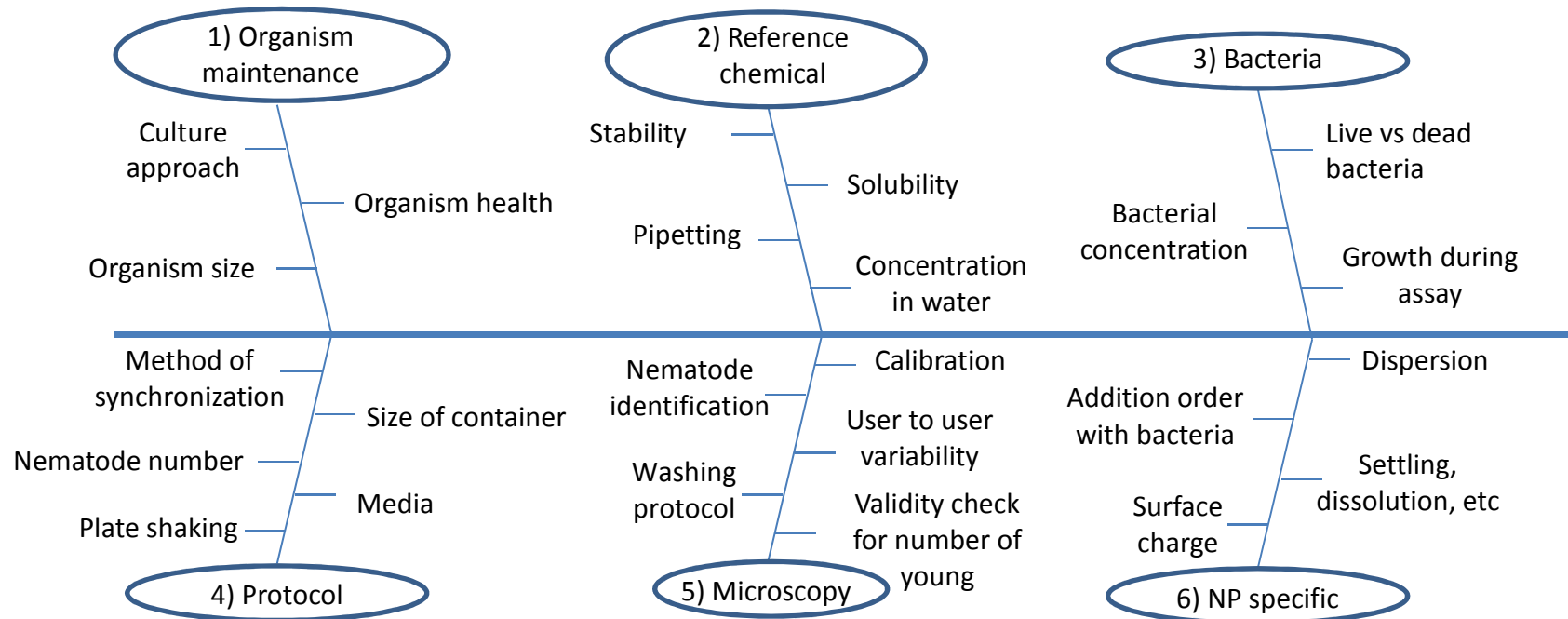


Figure 2.

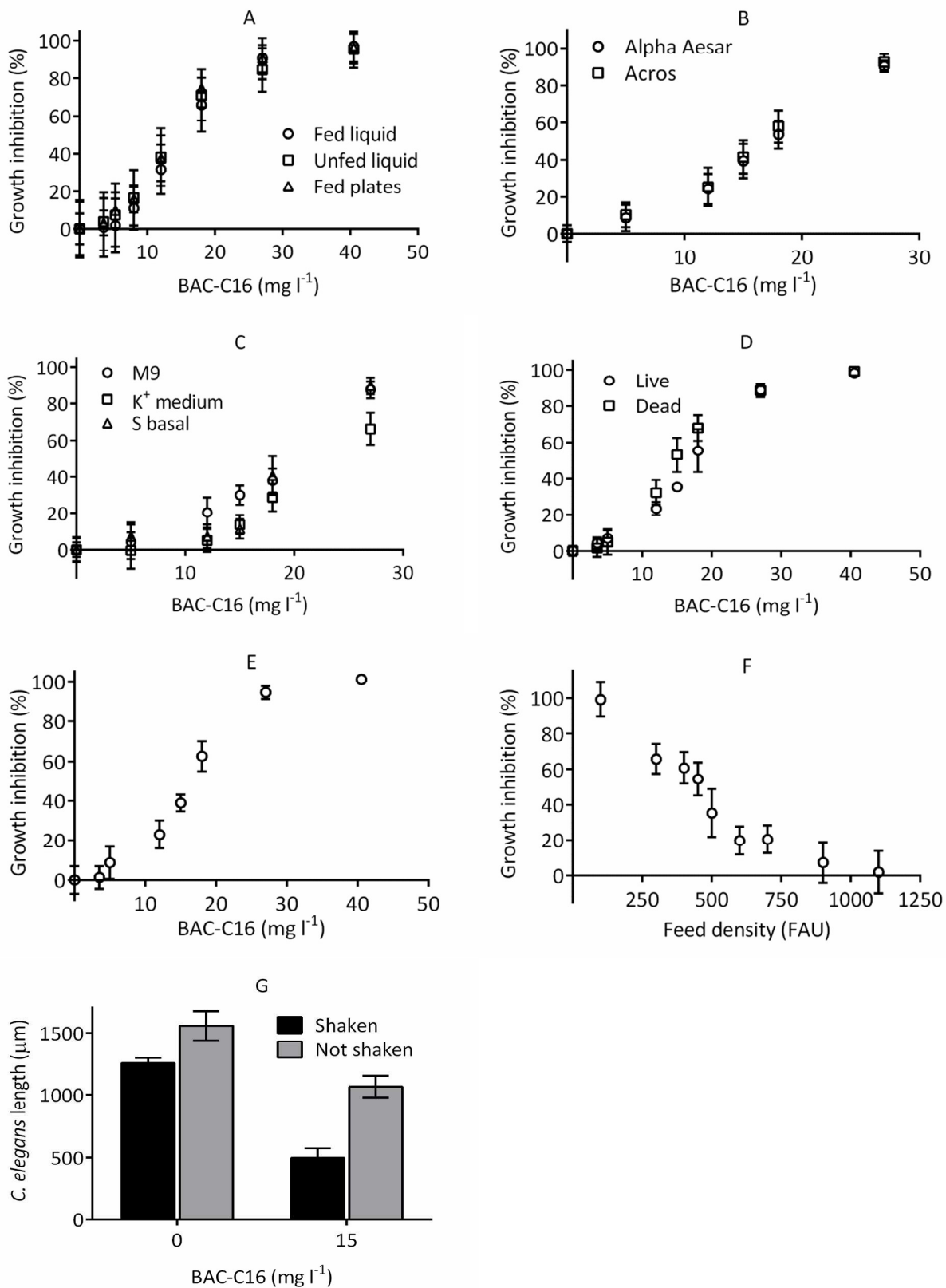


Figure 3.

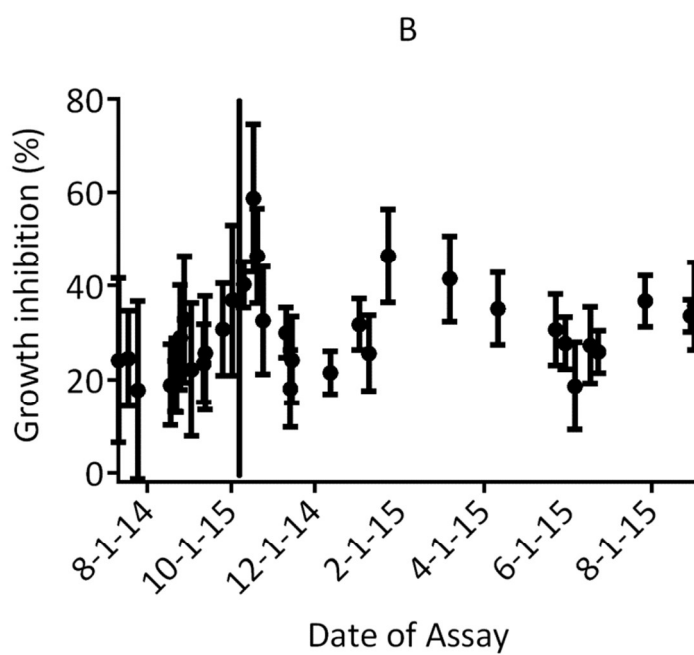
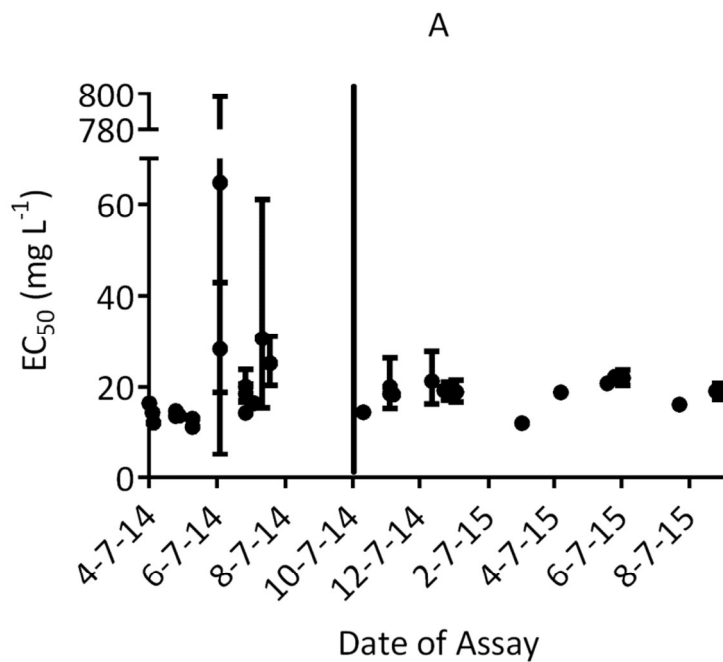


Figure 4.

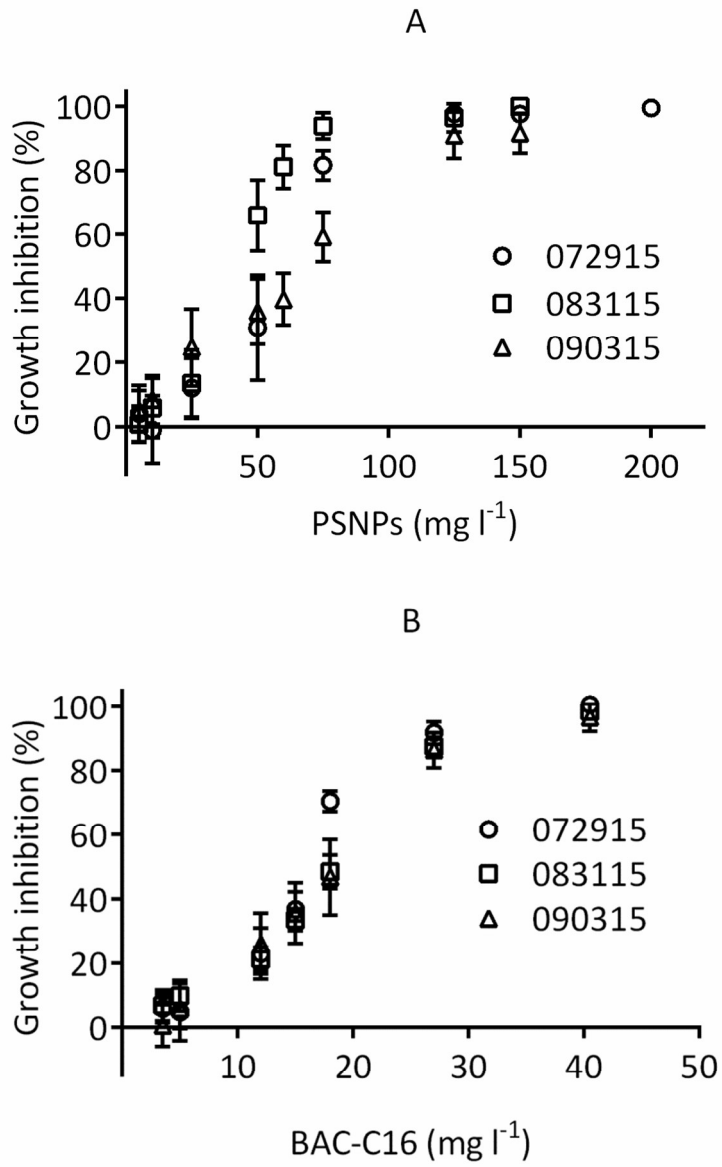


Figure 5.

