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November 16, 2018

Mr. Christopher Bittner Standards Coordinator Utah Dept. of Environmental Quality 195 N 1950 W Salt Lake City, UT 84116 Dr. Gary Belovsky Environ. Res. Center & Dept. Biol Sci. University of Notre Dame Notre Dame, IN 46556

RE: Results of Acute Lead Toxicity to Brine Shrimp

Mr. Bittner / Dr. Belovsky:

Enclosed is a copy of the final report entitled *Acute Toxicity of Lead to* Artemia franciscana *under Static-Renewal Test Conditions.*

We greatly appreciate this opportunity to provide our services to you. Please do not hesitate to contact us if you have any questions.

Sincerely.

Rami B. Naddy, Ph.D. V Manager / Senior Toxicologist naddyrb.tre@gmail.com

David A. Pillard, Ph.D. Manager / Senior Toxicologist pillardda.tre@gmail.com

Enclosure:

17001-474-018

University of Notre Dame Notre Dame, IN

Acute Toxicity of Lead to *Artemia franciscana* Under Static-Renewal Test Conditions

Prepared by:





TRE Environmental Strategies, LLC 100 Racquette Drive, Unit A Fort Collins, CO 80524

Document No. 17001-474-018

November 2018

Study Title

Acute Toxicity of Lead to Artemia franciscana in Laboratory Reconstituted Salt Water Under Static-Renewal Test Conditions

Study Period

Start: February 5, 2018 @ 15:30 End: February 9, 2018 @ 16:05

Performing Laboratory



TRE Environmental Strategies, LLC 100 Racquette Drive, Unit A Fort Collins, CO 80524

Telephone: (970) 416-0916 FAX: (970) 490-2963

Laboratory Project ID

17001-474-018

SUMMARY

Study Directors	Rami B. Naddy, Ph.D. David A. Pillard, Ph.D.
Test Facility	TRE Environmental Strategies, LLC 100 Racquette Drive, Unit A Fort Collins, Colorado 80524 (970) 416-0916
Location of Data	TRE Environmental Strategies, LLC 100 Racquette Drive, Unit A Fort Collins, Colorado 80524 (or an offsite storage location)
Test Substance	Lead Nitrate (Pb(NO ₃) ₂ ; Fisher Scientific Lot #164061; TRE #C17-020)
Subject	Static-Renewal Toxicity Test
Test Dates	Initiated: February 05, 2018 @ 1530 Terminated: February 09, 2018 @ 1605
Length of Study	96 Hours
Test Species	Artemia franciscana
Source of Organisms	Brine Shrimp Direct (Ogden, UT)
Age of Test Organisms	48 Hours
Test Concentrations	Nominal Lead Concentrations: 0, 48,000, 80,000, 134,000, 222,000, 371,000 µg Pb/L
Dilution Water	Laboratory Saltwater Reconstituted Water (rGSL; RW #13175) Target: Salinity ~ 120 ppt
Results	96-Hour LC ₅₀ Based on Measured Lead Concentrations: 130,000 μg total recoverable Pb/L; 95% C.I. (121,300 – 139,300) μg dissolved Pb/L; 95% C.I. ¹

¹ Dissolved median lethal concentration was not calculated because of the low and variable dissolved lead concentrations compared to total recoverable lead concentrations. See analytical results.

Sponsor and Laboratory Information

	Environmental Research Center
0	University of Notre Dame
Sponsor	97 Galvin Life Sciences Center
	Notre Dame, IN 46556
Project Officer	Gary E. Belovsky, Ph.D. (574) 631-0172
	TRE Environmental Strategies, LLC
Testing Feeility	100 Racquette Drive, Unit A
Testing Facility	Fort Collins, Colorado 80524
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Study Directore	Rami B. Naddy, Ph.D. (970) 416-0916 email: naddyrb.tre@gmail.com
Study Directors	David A. Pillard, Ph.D. (970) 416-0916 email: pillardda.tre@gmail.com

Introduction

This report presents the results of a study conducted to determine the acute toxicity of lead (as lead nitrate) to *Artemia franciscana* in a laboratory reconstituted salt water under static-renewal test conditions. The ultimate goal is to use these data to aid in the development of ambient water quality criteria for metals for the protection of species in the Great Salt Lake.

Methods followed the Work Plan for Great Salt Lake Toxicity Test, Version 8, October 23, 2016; Dr. Belovsky, University of Notre Dame, except where noted. The target water used in these studies was prepared to mimic Gilbert Bay water (see work plan; Appendix A), with a salinity of 120 ppt.

All toxicity tests were conducted at TRE (Fort Collins, CO). Chemical confirmation of lead was carried out at ALS Environmental (Kelso, WA; primary analytical laboratory) and Brooks Applied Labs (Bothell, WA; secondary analytical laboratory).

METHODS Test Media

The artificial reconstituted Great Salt Lake (rGSL) water was used in holding and testing conditions for the brine shrimp (but not in hatching). It was prepared as follows with ASTM Type I (Milli-Q[®]) water (ASTM 2012):

-) Crystal Sea marine mix: 50.95 g/L
- J Potassium chloride (KCl): 2.99 g/L
- / Magnesium sulfate (MgSO₄): 6.19 g/L
- J Sodium chloride (NaCl): 65.77 g/L

Salts were added to Milli-Q[®] water in a 20 L carboy and stirred to mix salts. Analytical results for this water are reported in the results section. The laboratory reconstituted salt water had an initial salinity of ~120 ppt.

Test Organisms

Test organisms were *Artemia franciscana* nauplii obtained as cysts from Brine Shrimp Direct (Ogden, UT). Brine shrimp were hatched in a 1.0-L separatory funnel containing artificial reconstituted seawater (29 ppt at 29 °C) made using Crystal Sea Marine Mix according to TRE SOP #5104. Hatched nauplii were kept in the pre-test holding conditions for 48 hours before test initiation.

The food source for the *A. franciscana* was the salt water alga *Dunaliella viridis*. *D. viridis* was cultured at ~10°C with 16:8 h light: dark photoperiod under constant aeration. The media used to culture *D. viridis* consisted of the following added to Milli-Q[®] water:

- Morton's Water Softener Salt (80 ml/L)
-) Crystal Sea marine mix (53.3 ml/L)

The media was mixed well and filtered (Buchner funnel) through a 110-mm Whatman[®] #4 filter paper and sterilized using a pressure cooker. Nutrients (1-3 ml) were added to the *D. viridis* cultures 1-2X per week. The nutrient solution consisted of Milli-Q[®] water (80 ml), blue solution (10 ml), and P/N solution (10 ml). The Blue solution consisted of 41.7 g/L Hydrosol, 27.5 g/L calcium nitrate, and 22.1 g/L ammonium nitrate dissolved in Milli-Q[®] water. The P/N (phosphorus/nitrogen) solution consists of 8.79 g/L monopotassium phosphate and 20.0 g/L ammonium nitrate dissolved in Milli-Q[®] water.

Pre-Test Conditions

In the pre-test holding conditions 200 < 24-h old brine shrimp nauplii were placed in 50 ml of 120 ppt rGSL seawater in a 80-ml Pyrex beaker and fed *D. viridis* at a concentration of 100 µg/L Chl*a* (chlorophyll a) at the beginning of the holding period (no additional food was added during the pre-test holding period). The solutions were gently aerated during the holding period. Five batches of brine shrimp (e.g., 200 organisms each) were prepared and held in this manner to ensure a sufficient number of organisms for the toxicity test. After 48-h, the organisms were transferred to the test chambers using a block design (e.g., organisms from the first chamber were used for all 'A' replicates, etc.).

Test Conditions

The chemical used in testing was lead nitrate $(Pb(NO_3)_2;$ Fisher Scientific, Lot #164061, TRE # C17-020). Individual test solutions were prepared by addition of the appropriate volume of the lead stock solution (3,900 mg/L as Pb) to rGSL water. Algae (*D. viridis*) were added at a concentration of 145 µg/L Chla along with the metal stock solution and rGSL to a total volume of 1.0 L. This volume was sufficient for biological and analytical sampling. The solutions were allowed to equilibrate for ~3 hours prior to use in testing.

After the equilibration period, the solutions were shaken to re-suspend any settled algae / precipitate to ensure homogenous distribution to test chambers for testing and for analytical sampling (there was a visible metal precipitate observed in the higher lead treatments). Approximately 150 ml of solution was poured into each test chamber (n = 5). Test chambers consisted of 12 ounce Pro-Kal[®] polypropylene cups². Twenty 48-h old brine shrimp nauplii were placed in each test chamber, and test chambers were placed in a randomized design in a temperature-controlled water bath (20°C). The photoperiod was 16 h light:8 h dark using fluorescent lighting.

The study consisted of a 96-h exposure period in which *A. franciscana* were exposed to different concentrations of lead. The test solutions were renewed after 48 h³ with test solutions that had been equilibrated with *D. viridis* (same feeding rate as at test initiation) for ~ 3-h. Surviving organisms were verified at 48 h (test solution renewal) and 96 h (organisms were not handled on days 1 and 3 so surviving organisms on these days were estimates). While organism growth was included in the work plan it was determined not to be required for these studies and therefore, only survival was measured.

² Polypropylene test chambers were used instead of HDPE test chambers as mentioned in the work plan ³ The work plan mentioned renewals every 24-h although preliminary analytical work indicated that 48-h renewals were sufficient for analytical and biological needs; therefore the latter renewal cycle was adopted.

Further detail is provided in Table 1 below.

Туре	96-h Static-Renewal Acute (renewal at 48-h; see deviation to work plan)
Test Endpoints	Mortality (no response to stimulus); see deviation to work plan regarding growth endpoint
Test Concentrations (nominal)	0 (control), 48000, 80000, 134000, 222000, and 371000 Îg Pb/L
Quality Criterion	90% control survival
Analytical Confirmation	Test initiation (new): Dissolved and total recoverable samples for each treatment Test renewal (new and old): Dissolved and total recoverable samples for each treatment Test termination (old): Dissolved and total recoverable samples for each treatment
Lead Analyses	ICPMS (EPA Method 200.7) ALS Environmental – Primary Analytical Laboratory ICP-QQQ-MS (EPA Method 1638) Brooks Applied Labs – Secondary Analytical Laboratory
Statistical Analyses	96-h median lethal concentrations were determined using CETIS 2014

Table 1. Additional Test Conditions in the Toxicity Test

Reference Toxicant

Three reference toxicant studies were performed with *A. franciscana* to determine the relative sensitivity of the organism. A summary of the results of the reference toxicant tests is presented in the results section below.

RESULTS

The initial characteristics of the rGSL water used in this experiment are reported in Table 2.

Table 2. Initial Dilution/Control Water Characterization

Batch No.	рН	Hard. (mg/L) ^a	Alk. (mg/L)ª	Spec. Cond. (~S/cm)	Salinity (ppt)
13175	8.3	12,900	187	195,400	122

^a As CaCO₃

^b Total residual chlorine

The batch of rGSL water was also analyzed for dissolved and total recoverable metals (Table 3).

Total _____ Total Recoverable

Table 3. Characterization of rGSL Water used in Toxicity Testing

Total Recoverable (μg/L)	Dissolved (µg/L)	Total Recoverable (μg/L) with food added
1.0 U	1.0 U	1.0 U
0.50 U	0.50 U	0.50 U
0.020 U	0.020 U	0.020 U
0.020 U	0.020 U	0.026
0.56	0.25	37.2
0.41	0.28	4.93
0.077	0.032	0.898
0.20 U	0.20 U	
22.0	22.0	23.4
1.0 U	1.0 U	
0.044	0.020 U	0.208
0.102	0.022	0.465
9.68	6.21	52.6
	Recoverable (μg/L) 1.0 U 0.50 U 0.020 U 0.020 U 0.56 0.41 0.077 0.20 U 22.0 1.0 U 0.044 0.102	Recoverable (μg/L)Dissolved (μg/L)1.0 U1.0 U0.50 U0.50 U0.50 U0.50 U0.020 U0.020 U0.020 U0.020 U0.020 U0.020 U0.560.250.410.280.0770.0320.20 U0.20 U22.022.01.0 U1.0 U0.0440.020 U0.1020.022

Analyzed by ALS Environmental Laboratory (Appendix B)

U = Analyte was not detected at or above the MRL/MDL, which is reported.

Food (algae) was not present in these samples unless noted. There was insufficient volume to analyze the rGSL water with food added for Hg and Se.

Dissolved organic carbon (DOC) and total organic carbon was also measured in the rGSL control water with food added. The DOC and TOC was <1.0 mg/L.

The range of water quality parameters measured during the toxicity test is provided in the table below (Table 4). Overall, the pH, dissolved oxygen, conductivity, and temperature were similar among treatments.

Treatment (Nominal Conc., μg Pb/L)	р	н	Disso Oxygen			ictivity /cm)	-	erature C)
	Low	High	Low	High	Low	High	Low	High
0 (Control)	8.0	8.3	5.1	5.4	122.3	195.4	20	21
48,000	7.9	8.1	5.1	5.4	123.8	196.3	20	21
80,000	7.8	8.0	5.0	5.5	123.1	193.1	20	21
134,000	7.6	7.9	4.9	5.4	123.6	190.5	20	21
222,000	7.3	7.8	4.9	5.4	124.7	192.0	20	21
371,000	7.0	7.4	4.7	5.2	128.8	186.5	20	21

Table 4. Physical and Chemical Data Measured during the Toxicity Test

Analytical Confirmation

Samples were collected for total and dissolved lead analyses from new and old test solutions as outlined in the Methods section. Average concentrations for all treatments are presented in Table 5.

Table 5. Measu	Ired Lead C	Concentrations
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Nominal					
Conc.	Avg Total Rec.	% of	Avg. Dissolved	% of	Diss. / Tot.
(µg Pb/L)	(µg Pb /L)	Nominal	(µg Pb/L)	Nominal	(%) ^a
0 (control)	53 U		53 U		
48,000	38,925	81	4,398	9	7
80,000	65,700	82	7,075	9	11
134,000	109,500	82	9,563	7	10
222,000	199,250	90	25,150	11	18
371,000	288,500	78	65,650	18	35

^a Initial samples only

Note: Refer to Appendix B for a summary of analytical results

U = less than the MRL / MDL (which is reported)

In comparing total recoverable versus dissolved lead concentrations, dissolved lead concentrations were 7-35% of the total recoverable lead concentrations and 7-18% of nominal

concentrations. Dissolved lead concentrations do not appear to be representative of actual lead concentrations, and therefore, endpoints were not calculated using dissolved lead values.

Quality assurance checks included both duplicate and split sample analyses. Duplicate samples (an aliquot of the same test concentration split and sent to the primary laboratory in two separate bottles) and split samples (an aliquot of the same test concentration split and sent to the primary and secondary analytical laboratories). In addition, a blank sample (Milli-Q[®] water) was sent to the primary laboratory and analyzed for total recoverable and dissolved lead. Results are presented in Table 6 below. A summary of all the samples analyzed is included in Appendix B.

Nominal Conc. (μg Pb/L)	Sample Type	Total Rec. (μg Pb /L)	Dissolved (µg Pb/L)	Relative Standard Deviation
48,000	Original		6,370	1.48
40,000	Duplicate		6,490	1.40
	Original		6,260	1.87
80,000	Duplicate		6,080	1.07
80,000	Original	65,900		7.45
	Split	71,000		7.45
	Original		11,100	0.00
134,000	Duplicate		11,100	0.00
134,000	Original	107,000		15.52
	Split	125,000		15.52
	Original	201,000		2.92
222,000	Duplicate	204,000	11,100 11,100 	2.92
222,000	Original	193,000		3.56
	Split	200,000		3.50
371,000	Original		18,300	20.59
371,000	Split		22,500	20.09
Control	Original	0.077		
Control	Split	<1.02 U		
Milli-Q water	Blank	53 U	53 U	

Table 6. Measured Lead Concentrations – QA Samples

Organism Response

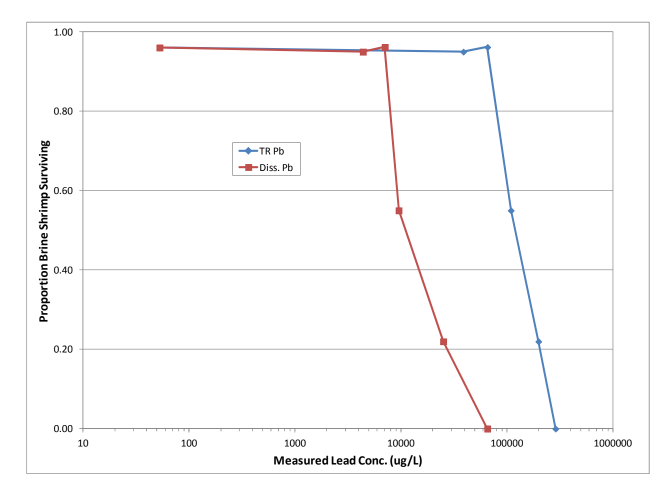
The definitive test was initiated February 5, 2018 at 1530 hours and was terminated on February 9, 2018 at 1605 hours. *A. franciscana* survival at 48 h and 96 h is shown in Table 7.

Nominal Conc.	% Survival		
(~g Pb/L)	0 h	48 h	96 h
0 (control)	100	97.0	96.0
48,000	100	97.0	95.0
80,000	100	99.0	96.2
134,000	100	95.0	55.0
222,000	100	86.0	22.0
371,000	100	0.0	0.0

Note: See Appendix C for a copy of raw data

A graphical representation of the 96-hour survival data, based on measured lead concentrations (log scale) is shown in Figure 1.

Figure 1. Proportion of *Artemia franciscana* Surviving at 96 hours versus Measured Lead Concentrations.



Dissolved lead concentrations were included in Figure 1 for comparison, even though dissolved lead concentrations do not provide a good estimate of nominal lead concentrations.

The calculated 96 hour LC_{50} values for *A. franciscana* are provided below for total recoverable lead (Table 7). The median lethal concentration was not calculated for dissolved lead because of the low and variable dissolved lead concentrations (see Table 5).

Table 7. 96 hour Media	n Lethal Lead	I Concentrations (~ g/L)
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Endpoint	Total Rec. (µg Pb/L)	Dissolved (µg Pb/L)
LC ₅₀	130,000	
95% C.I.	121,300 – 139,300	
Method	Trimmed Spearman-Karber	

Note: median lethal dissolved lead value was not calculated because dissolved lead was not found to be a good surrogate of nominal lead concentrations. See analytical results.

Reference Toxicant Studies

Three additional studies were initiated with *A. franciscana* using arsenic for the purpose of reference toxicant tests. The average results of these reference toxicant studies (mean ± 2 standard deviations) are presented below (Table 8).

Table 8. Reference Toxicant Test Results for A. franciscana

96-h LC₅₀ (mg As/L)	TRE Historical 95% Control Limits (mg As/L)			
	Low	High		
70.57	53.97	87.16		

Note: Values are expressed as mg/L of nominal arsenic.

References

- ASTM. 2012. Standard Guide for Conducting Acute Toxicity Tests on Test Materials with Fishes, Macroinvertebrates, and Amphibians. E 729-96. Annual Book of ASTM Standards, Volume 11.05, Section 11, Water and Environmental Technology.
- Belovsky, G. Work Plan for Great Salt Lake Toxicity Tests, Version 8. October 23, 2016.
 Environmental Research Center and Department of Biological Sciences. University of Notre Dame.
- CETIS. 2014. Comprehensive Environmental Toxicity Information System. User Guide (version 1.8.7). Tidepool Scientific, LLC. McKinleyville, CA.

STATEMENT OF PROCEDURAL COMPLIANCE

I certify that this document and all attachments were prepared under my direction or supervision in accordance with a system designed to assure that qualified personnel properly gather and evaluate the information submitted. Based on my inquiry of the person or persons who manage the system, or those persons directly responsible for gathering the information, the information submitted is, to the best of my knowledge, accurate and complete.

Rami B. Naddy, Ph.D. Study Director

)ovenber 13 2018

Date

STATEMENT OF QUALITY ASSURANCE

The test data were reviewed by the Quality Assurance Unit to assure that the study was performed in accordance with the protocol and standard operating procedures. This report is an accurate reflection of the raw data generated at TRE.

Quality Assurance Unit

lovember 16, 2018

Date

APPENDIX A

WORK PLAN

Work Plan For Great Salt Lake Toxicity Tests, Version 8 October 23, 2016

Gary Belovsky Environmental Research Center & Department of Biological Sciences University of Notre Dame

Introduction:

Great Salt Lake (GSL) is a unique ecosystem, the fourth largest (largest in the western hemisphere) hypersaline lake in the world¹⁶. Invertebrate life in the GSL is relatively species poor due to the high salinity of the lake and is dominated by brine shrimp (*Artemia franciscana*) and brine fly larvae. Two or more species of brine flies occur in the GSL with *E. cinerea* the most abundant by several orders of magnitude. Thus, *Artemia franciscana* and *Ephydra cinerea* are the dominant grazers in the GSL food web in Gilbert Bay (South Arm) of Great Salt Lake, and are the focus of this study. These invertebrates are very abundant and are the main source of food for many resident and migrating water birds, which have important ecological and conservation value. Some of these water bird species are threatened or endangered or have other legal protections.

The purpose of this project is to determine toxic levels of copper, arsenic and lead, to brine shrimp and brine fly larvae so that water quality criteria (WQC) can be developed for GSL as described in the State of Utah Division of Water Quality (UDWQ) Great Salt Lake Strategy²². These potential pollutants were identified as the highest priorities by UDWQ after public feedback and comments.

We will conduct acute toxicity tests of the above substances for brine shrimp and brine fly larvae (*E. cinerea*) and develop a plan of work to assess chronic toxicity of these trace elements. *E. cinerea* has been chosen as a test species because of its local abundance end ecological dominance and it has been successfully cultured in the laboratory. American Society for Testing and Materials (ASTM) has not sanctioned standard methods for toxicity testing with either of these species. However, other ASTM-approved methods for invertebrates are established and will be used as a guide for the conduct of the toxicity testing proposed here.

In this work plan we present the protocols that lead to uniform rearing of the brine shrimp and brine fly larvae, as well as production of control and test (pollutant) solutions. We then focus on range-finding and acute testing, which must be completed before proceeding with the ecologically more relevant chronic testing (survival and reproduction with life-time exposure to the pollutant), which will be detailed in a future work plan. <u>Range-finding</u> establishes the range of concentrations for each pollutant that produces short term (96 hour) toxicity (assessed by mortality and growth). Once the range for each pollutant has been established, <u>acute tests</u> will be conducted to establish the concentration of each pollutant that will produce 50% mortality over 96 hours of exposure (LC50). For this project, the Belovsky laboratory has primary responsibility for developing the culturing methods for the brine flies and brine shrimp and will provide brine flies to the bioassay laboratory. The bioassay laboratory has primary responsibility for conducting the toxicity tests for both species –*Artemia franciscana* cultured at the bioassay laboratory per protocols provided by the Belovsky lab, and *Ephydra cinerea* which will be provide by the Belovsky lab.

Source animals and rearing conditions:

Given the uniqueness of Gilbert Bay, Great Salt Lake compared to other aquatic environments for which ecotoxicology studies have been developed, it is critical that our toxicology studies provide results that are applicable to the lake's environment. The environmental conditions found in the Great Salt Lake in April – October (the time when brine shrimp are present) over a 20 year period $(1994 - 2013^{16, 17})$ are summarized in the table below.

Consistency of *Artemia* will be ensured by using a single batch of brine shrimp cysts (resting eggs) that have been commercially harvested from GSL. Brine fly larvae will be obtained from a colony maintained at UND, where GSL was the source of individuals starting the colony and the colony has been in existence for approximately two years (6 - 8 generations). The 3rd instar were selected because this is the longest and final larval stage when the most growth and development occurs²⁰. This stage can be easily collected without damaging them from the rugose surface of bioherms on which they are raised in the laboratory colonies. Furthermore, this life stage is long enough to support shipping the larvae and robust enough to experience <10% mortality in transit.

Both species will be reared in environmental chambers that maintain temperature (\pm 1°C) and a light:dark cycle (16:8, ~summer day using full spectrum lighting) using the following protocols:

Brine shrimp will be hatched in 10 gallon aquaria at a salinity of 45 ppt, the optimum for hatching and hatchling survival. Nauplii will be used in the bioassays as it is thought that they are most susceptible due to their small size and less developed exoskeleton. Twenty-four hours after hatching, the nauplii will be transferred to artificial GSL water (see below). Over the initial 2 days post hatch, individuals are fed ad libidum a high quality phytoplankton (*Dunaliella* sp.: 40 µg Chl_a/L/2 days) maintained in culture.

Brine fly larvae will be reared in plastic containers (60 cm X 60 cm X 25 cm) that contain 12 cm of water that is maintained at average lake conditions specified above at the University of Notre Dame. Gravel and GSL bioherm (approximately 30 cm X 15 cm X 15 cm) serve as a substrate for larvae and pupae, and an above water platform is provided to emerging adults for resting and mating. Larvae will be fed ad libidum (pupae and adults do not feed) a high quality food (*Dunaliella* sp.: 40 µg chl_a/L/2 days) from a colony established from the GSL. Notre Dame personnel ship late 2nd or 3rd instar larvae based on size (FedEx overnight) with a resin fiber pad (3M TM Scotch Brite #86) for attachment and food (*Dunaliella* sp.) in a plastic bag with head space and bags in a cooler with ice. Larvae will then be acclimated in artificial GSL water (see below) for a minimum of four days prior to the beginning of each test.

Dilution Water:

Artificial reconstituted GSL water (rGSL) (Table 1, salinity = 120 ppt mass/volume) will be made to duplicate Gilbert Bay water as close as practical. This specific artificial reconstituted water was selected based on considerations of the data quality objectives (see Appendix 1).

The rGSL attempted to match the average concentrations of salts that are shown in Table 2 based on Utah Geological Survey measurements from Gilbert Bay¹⁸ (Table 2). Initially, no attempt will be made to mimic the dissolved organic content (DOC) of Gilbert Bay. DOC in Gilbert Bay water has been reported to reduce the toxicity of copper and other metals compared to artificial Great Salt Lake water³ (Brix et al 2006). Using Gilbert Bay water instead of artificial water would duplicate the DOC concentrations at the time the water was collected but the representativeness over time is unknown. For instance, reported DOC concentrations include 7 mg/L to 42 mg/L^{3,2}. Amending the rGSL with DOC was considered but was primarily rejected by UDWQ because of the lack of data to support determining an appropriate target DOC concentration.

For the acute tests, rGSL will be made using reagent grade Crystal SeaTM Bioassay Laboratory Formula Marinemix® (Marine Enterprises, Baltimore, Maryland). Marinemix is approved by EPA for toxicity testing¹⁹ and American Chemical Society (ACS) grade salts to deionized water in 20L Nalgene[®] carboys. Facility Deionized filtered water will be used. pH will be maintained at 7.9 ± 0.2 via the addition of 0.1N nitric acid or 0.1N sodium hydroxide as necessary. Filtered triplicate samples will be collected from each batch to verify that the salt concentrations and pH are within acceptable ranges.

Recipe	g/L
MarineMix	50.95
KCl	2.99
MgSO ₄	6.19
NaCl	65.77

Table 1. Reconstituted	l Great Salt Lake water	used for acute toxicity testing.
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Constituent	Great Salt Lake	rGSL	%
	Average		match
Na	42.0	41.53	98.54
Mg	4.4	3.19	100.77
Cl	74.0	69.33	100.48
К	2.8	2.14	100.40
Ca	0.3	0.62	99.73
SO_4	9.3	8.85	99.95
HCO ₃	0.4	0.22	100.14
CO3	No data	0.02	No
			data

Table 2. Reconstituted Great Salt Lake (rGSL) water nominal concentrations compared to Utah Geological Survey average Great Salt Lake (Gilbert Bay) Concentrations from 1966-2013¹⁸.

Prior to initiating testing using organisms, the rGSL will be characterized for trace metals and metalloids and the analytical results from the copper range finding test (Appendix 2) will be confirmed. To characterize the rGSL, the concentrations of antimony, arsenic, beryllium, cadmium, chromium, copper, lead, mercury, nickel, selenium, silver, thallium and zinc will be measured. To confirm the analytical results from the copper range finding test, a sample of the

rGSL will be spiked with $450 \pm 50 \mu g/l$ of copper as CuCl₂. The salinity of the rGSL may need adjustment to achieve the target salinity of 120 ppt after spiking with CuCl₂. The copper-spiked rGSL will be stirred to mix and both filtered (0.45 μ m) and unfiltered samples collected and analyzed. The measured concentrations in the filtered samples should be at least 70% of nominal. If measured concentrations are less than 70% of nominal, additional experiments will be pursued to determine the fate of the copper spikes.

Test solutions (treatments):

The pollutants chosen for these studies were selected by UDWQ after soliciting public comment. American Chemical Society (ACS) reagent grade chemicals will be employed. To maximize comparability of test results with published ecotoxicology studies²⁻¹⁴ that have been conducted with other aquatic invertebrates, the following chemicals will be used to produce test solutions:

- As (arsenic) as sodium arsenate;
- Cu (copper) as copper chloride;
- Pb (lead) as lead nitrate.

Stock solutions of each trace metal will be made fresh for each of three rounds of acute toxicity testing as recommended by US EPA. Solutions will be made in Teflon sample bottles and solution concentrations will be verified at the beginning and end of each exposure 24-hour day of the bioassay (see details below).

Range finding tests:

For each pollutant to be tested, a preliminary range finding test will be conducted to establish concentrations to be used in the acute tests so that well resolved concentration-response (mortality or immobility) curves result. Initial range finding will be conducted with 10-fold increases in dissolved concentrations (e.g., 0 (controls), 10, 100, 1,000, 10,000, 100,000 µg of the pollutant/L) to ensure that we can develop well-resolved concentration response curves. Exposure to each concentration will be tested in triplicate for each test species over a 96 hour period on a static renewal basis (100% water changes occurring at 24, 48 and 72 hours), as follows:

Brine shrimp – 48 hour old nauplii will be used for all acute tests (including preliminary range finding tests). Test chambers will be 250 mL HDPE beakers containing 150 mL of test solution. Nauplii will be hatched in Marine Mix formula at 45ppt and then transferred to MarineMix at 120 ppt for tests. The shrimp nauplii will be fed *Dunalliela* from culture at a rate of 190 µg Chl_a/L/day. These feeding rations were determined to be the minimum rations required to achieve \geq 90% control survivorship in feeding trials were conducted at UND. The feeding trials were conducted using 5 replicate treatments using the same rGSL, temperature, number of nauplii and container size as the acute tests. The initial survival was <90% and the food was increased by 50% until >90% survival was observed. Food was decreased from this rate to the midpoint of the next lowest feeding rate and the trials repeated until the minimum amount of food resulting in \geq 90% survival was determined (190 µg Chl_a/L/day).

For the acute tests, feeding rates will be measured by taking at least three (3) Chl_a measurements of the feeding solution with the fluorometer (equivalent FSUs, fluorescent signal units, for the

necessary μ g Chl_a feeding level) will be made after mixing vigorously between measurements. The test conditions are summarized in Table 3.

Brine fly larvae – 3^{rd} instar larvae will used in all acute and range finding tests. Ten individuals per replicate will be held in a 250 ml HDPE beaker containing 150 ml of test solution. The larvae will be fed *Dunalliela* from culture at the minimum necessary to achieve rate $\geq 90\%$ control survivorship as determined at UND using the same feeding trial methods as for nauplii. This food level is attained by vacuum filtering at ~1 atmosphere 150 mL of 490 µg Chl_a/L through a glass microfibre filter (Whatman Catalog # 1825 024) 50 mL at a time. The FSU of the filtrate will be measured and recorded to verify that the phytoplankton were captured on the filter. The filter paper will be placed phytoplankton-side up in the bottom of the empty beaker. A piece of a resin fiber pad (3M TM Scotch Brite #86) weighted with a paperclip will cover the top of the filter and provide a substrate for the larvae. The larvae are then added to the beaker.

For brine shrimp and brine fly larvae, beakers will be acid washed (5% HNO₃), rinsed several times with deionized water and dried under a laminar flow hood prior to use. Each beaker will be covered with parafilm® to reduce evaporative water loss and associated changes in salinity and pollutant concentration. The resulting headspace in each beaker will provide ample gas exchange (O_2 and CO_2) on a 24 hour basis for the animals between daily treatment solution changes. Individuals will be randomly assigned to treatment groups, and the placement of beakers in the incubators will be randomized as well. pH, DO, and conductivity in each beaker will be checked daily, as well as in a "monitoring replicate" (no test organisms present) for each concentration.

Dead and immobile individuals will be recorded daily. Dead individuals are those that are immobile and unresponsive to stimuli (touched with pipette), while immobile individuals are not observed to move until touched with the pipette. Dead individuals are removed by pipette daily at the time of treatment solution change. After the sampling and methods for measuring concentrations in the test solutions are verified to be accurate, analytical chemistry of the treatment solutions will not be conducted in the range testing work, because it is only necessary to establish which treatment solutions produce mortality or immobility and to reduce analytical costs.

Acute Toxicity Assays:

Concentration ranges for acute assay treatment solutions – Based on range finding tests, we will run appropriate dilution series such that we can make robust statistical estimates of LC50 concentrations for each species. Six test concentrations (including controls) will be used for each species. If it is determined that the concentrations required to elicit mortality exceed the solubility limit of the metal under our test conditions, acute testing will be halted and the pollutant will proceed to chronic testing.

Acute assay protocols – Acute assay protocols are summarized in Table 5. Assays will be executed as described in range finding tests (see above) with the exception that 5 replicates will be used per test concentration (as opposed to 3 for the range finding tests). Fresh test solutions will be made daily and verified (see below). Tests will be run on a static renewal basis

with a 100% water change and the removal of dead test organisms occurring on a daily basis. At the termination of each test, surviving organisms will be counted, and dried on pre-weighed filter paper for analysis of growth differences between treatments. Mortality data will be analyzed via probit analysis.

<u>Analytical chemistry.</u> Test exposure concentrations will be verified daily at the beginning and end of each 24 hour exposure period to ensure that we are accurately characterizing exposure conditions. When possible and supported by the range finding results, brine flies and brine shrimp will be tested simultaneously to reduce analytical costs. Table 4 illustrates the number of analytical samples (88) needed assuming that both brine shrimp and brine flies are tested simultaneously with 5 exposure concentrations plus the control. Three of these exposure concentrations are assumed to be identical for both brine flies and brine shrimp and 2 of the exposure concentrations are unique to both brine flies and brine shrimp.

Note that the addition of *Dunaliella* as a food source for *Artemia* nauplii and brine fly larvae will require that post exposure water samples are filtered to remove particulates (*Dunaliella*). Because the addition of live cells (*Dunaliella*) as a food source is likely to decrease the dissolved concentrations of the test chemicals, all statistical analyses will be based on the geometric mean of the initial (pre-exposure) and final (post-exposure) dissolved concentrations. Treatment solution samples will be filtered through acid washed (5% HNO₃) 0.45 µm syringe filters that have had 3 volumes of sample water passed through them prior to retaining the sample in the appropriate acid washed (5% HNO₃) sample tubes (500 ml). Comparisons of pre-exposure test solutions will be made between filtered and unfiltered samples to describe the relationship between total and dissolved metals under our test conditions. Samples will be preserved and kept at 4°C in the dark in 15mL conical tubes for As, Cu and Pb samples which will be stabilized with Omnitrace nitric acid or as instructed by the analytical laboratory

Acute assay data analyses – Two toxic endpoints will be recorded - mortality and immobility. Records of daily deaths and immobility will be recorded, but analyses will be conducted on overall mortality and immobility over the entire 96 hour assay period for a given pollutant concentration. Measures of mortality and immobility for a given pollutant concentration will be presented relative to the respective values observed in the simultaneous controls (no pollutant). For an assay to be considered successful, \geq 90% of individuals in the control must survive.

With the above measures, the concentration-response (mortality or immobility) curves will be developed for a given pollutant and organism. These curves will be calculated via standard analytical procedures with diagnostic checks for homogeneity of variances using standard statistical packages . With the concentration-response curves, a number of toxicity effects for a pollutant can be estimated:

1) <u>LC50 and EC50</u> is computed as the concentration eliciting 50% mortality (LC50) and 50% immobility (EC50) relative to the organism's control values.

2) <u>Lowest concentration (LOEC)</u> affecting mortality and immobility is defined as the first test concentration to produce a statistically significant increase in mortality or immobility relative to control values.

3) <u>No effect concentrations (NOEC)</u> is the next lowest concentration tested relative to the LOEC.

Data archiving -- all water chemistry, QA/QC data, and toxicity (mortality and immobility) data will be provided to UDWQ and made available to any interested parties

	Brine shrimp	Brine Flies
Test Type	Static renewal	Static renewal
Test Duration	96-hr	96-hr
Temperature	20 +/- 1 °C	20 +/- 1 °C
рН	7.9 +/-0.1; adjusted with 0.1N nitric acid and/or 0.1N sodium hydroxide as needed	7.9 +/-0.1; adjusted with 0.1N nitric acid and/or 0.1N sodium hydroxide as needed
Salinity	120 ppt	120 ppt
Photoperiod	16 hr light/8 hr dark	16 hr light/8 hr dark
Test Concentration or Dilution Series	5 exposures + control (40% dilution series TBD by range finder test)	5 exposures + control (40% dilution series TBD by range finder test)
Test Chamber Size	250 ml	250 ml
Test Solution Volume	150 ml	150 ml
Renewal of Test Solution	daily	daily
Age of Test Organism	48 hr nauplii	3rd instar
Number of Organisms per Test Chamber	20	10
Number of Replicates per Concentration: range finder (acute test)	3(5)	3(5)
Number of Organisms per Concentration: range finder (acute test)	60(100)	30(50)
Feeding	Daily (see text)	Daily (see text)
Endpoint	mortality (LC50) and immobility (EC50)	mortality (LC50) and immobility (EC50)
Test Acceptability	≥ 90% control survivorship	≥ 90% control survivorship

Table 3. Summary of acute testing methods.

Time (hours)	0	24		4	48		72	
Exposure Solutions	initial	initial	final	initial	final	initial	final	final
Control	1	1	2	1	2	1	2	2
Brine Shrimp	1	1	1	1	1	1	1	1
Brine Shrimp	1	1	1	1	1	1	1	1
Both	1	1	2	1	2	1	2	2
Both	1	1	2	1	2	1	2	2
Both	1	1	2	1	2	1	2	2
Brine Flies	1	1	1	1	1	1	1	1
Brine Flies	1	1	1	1	1	1	1	1
Duplicate	1	1	1	1	1	1	1	1

Table 4. Minimum number of analytical samples for each toxicant tested (see text for details

Chronic Toxicity Test Plan of Work:

While conducting the acute testing portion of this project, we will develop a plan of work for chronic testing of the priority pollutants described above.

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Appendix 1

Bioassay Water For Great Salt Lake Bioassays



State of Utah

GARY R. HERBERT

Department of Environmental Quality

Alan Matheson

Executive Director

Governor

MEMORANDUM

TO: Great Salt Lake Bioassay Team

FROM: Chris Bittner

DATE: April 23, 2015 (edited for inclusion in Work Plan 5/4/16)

PROJECT: Great Salt Lake Toxicity Bioassays for Brine Shrimp and Brine Flies

SUBJECT: Bioassay water for Great Salt Lake Toxicity Bioassays

Conclusions

After evaluating how well each media met the data quality objectives, none of the media can currently be concluded to be superior. The DWQ Round Robin medium was scored the highest but has not been tested with the test organisms and concerns remain regarding the long-term stability.

Data Quality Objectives

- 1. USEPA approval is required because the test results are intended to ultimately support the development and promulgation of numeric criteria. USEPA approval is not independent of the other data quality objectives (DQOs) because if the other DQOs are met, USEPA would likely approve the results for criteria development. However, the salinity of Great Salt Lake (GSL) is not specifically addressed by USEPA Guidance or Rules and unavoidable deviations from the existing guidance and rules are anticipated. These deviations must ultimately be acceptable to USEPA for criteria development.
- 2. The test medium must support the test organisms which at minimum are anticipated to include brine shrimp, brine flies, and green algae. If the test medium doesn't support the test organisms as defined by acceptable survival, growth, and reproduction in the negative controls, the results won't support the development of numeric criteria.
- 3. The test medium must have minimal potential confounders that either increase or decrease toxicity. The goal of having no potentially confounding issues is the ideal and meeting this

DQO is anticipated to limited to accepting the least amount of potential confounders. Confounders include factors such as ion balance, pH, presence of pollutants to be tested, dissolved organic carbon and hardness. Some of these factors are discussed in the context of DQO 5 because ideally, the medium mimics GSL's concentrations of these modifiers of toxicity.

- 4. The medium must be able to be replicated over time. Considerations for this DQO include that the source of the materials used for the medium should be stable over time and have documented quality control to ensure that any deviations can be identified and addressed if necessary. Based on the resources that are anticipated to be available for conducting bioassays and the number of existing pollutants, numeric criteria development for GSL is a long term project (e.g., 20+ years). Tests conducted 20 years should give the same results.
- 5. The DQO that the medium should be representative of Great Salt Lake is related to DQO 3 for minimal confounders affecting toxicity. The representativeness DQO is specifically specified because in a regulatory context, if bias is present, bias that overestimates toxicity is much more acceptable than bias that may underestimate toxicity. USEPA recommendations for toxicity testing media are intended to avoid underestimating toxicity for waters across the nation at the expense of potentially overestimating toxicity. To address this potential overestimation, effluent limits or potentially criteria can be modified on a site-specific basis using the Water Effects Ratio (WER). A WER is the ratio of toxicity between conducting bioassays using USEPA standard laboratory bioassay water and site water for dilution. The results may be the national criteria are either over- or under protective for the specific site. The GSL criteria are intended to be site-specific, so the bioassay medium should accurately reflect the toxicity, or lack thereof, of GSL waters obviating the need for WERs.
- 6. The medium should be stable over time. At minimum, the media must be stable for the test duration and ideally, the media would be stable over longer time frames. The stability of the medium may also affect the reproducibility of the toxicity testing.
- 7. The medium should be able to be replicated by any qualified laboratory. This DQO is similar to the other precision-related DQOs such as replication over time. The media composition must be sufficiently documented and the materials readily available to any qualified laboratory to meet this DQO.
- 8. Cost and convenience are the least important consideration but resource constraints are still an influential factor. When the scores for each DQO were summed, the cost and convenience score was not included.

Evaluation of Media

Several potential media were investigated:

- GSL Water (Brix et al., 2006)
- Barnes and Wurtsbaugh (2015)
- Belovsky
- DWQ Marinemix
- DWQ Round Robin

The advantages and disadvantages of each approach relative to the data quality objectives (DQOs) are summarized in the following text. Media that met, or were anticipated to potentially meet the minimum requirements are qualitatively scored on a scale of 1 to 10 for each DQO. Scores are summarized in Table 5.

Brix et al. 2006

Brix et al. (2006) used GSL water, artificial seawater, and the Bagshaw et al. (1986) media for conducting their bioassays. Although not documented in the paper, Bill Adams' (coauthor with Brix) recollection was that the GSL water was not filtered. When contacted, Mr. Adams opined that because of the quantities of water needed, filtering through a 0.45 μ m filter wasn't a practical option. While a larger filter may not clog as fast, decanting may be just as efficient as filtering. Brix et al. (2006) diluted the GSL water with deionized water to the salinity of seawater. Brix et al. (2006) concluded that the toxicity of copper in GSL water was much less compared to artificial seawater or the media of Bagshaw et al. (1986). Neither the artificial seawater medium nor the Bagshaw medium were considered further because they do not appear representative of Gilbert Bay water with regards to toxicity.

The use of GSL water might be approvable by USEPA. At minimum, this will require addressing the existing contaminant concentrations assuming they are low enough to not significantly affect the test results. A score of 7 is assigned for USEPA approval because using GSL water would require a deviation from existing USEPA protocols.

GSL water is anticipated to support the test organisms under current lake conditions and is scored a 10 for this DQO.

GSL water has many potential confounders that could affect the toxicity results. Existing pollutant concentrations (further discussed in the Marinemix section) for some metals are known but data for the majority of organic priority pollutants are unavailable. GSL water may also have unidentified pathogens or introduce other undesirable organisms into the bioassays. The concentration of dissolved organic carbon, sulfate, and hardness are expected to decrease the toxicity of metals. However, these affects are reflective of actual site conditions and are not considered undesirable for criteria development specific and limited to GSL. GSL water is scored a 5 for this DQO because of existing pollutant concentrations.

The consistency of GSL water over time is uncertain. Lake salinity is known to fluctuate over time (see Figure 1) and a protocol to address these potential fluctuations would need to be developed if salinity is determined to significantly affect toxicity. For instance, the test protocol may require salinities of 11 to 13%. If GSL water was higher, deionized water could be used to lower the salinity to the target range. UGS reports that the major ion ratios have remained similar (Gwynn, 1998). However, future changes are possible because of for instance, changes by the mineral extraction industries that selectively remove some of the salts and are required to return the remainder to the Lake. At GSL salinities lower than 11%, additional salts would have to be added. Depending on the quantity of salt required, the ion ratios of GSL water may not be achievable because maintaining the ion ratios may result in precipitation. This is judged to have a small potential because much higher salinities exist in GSL. Pollutant concentrations in GSL may increase over time resulting in the water being unacceptable for criteria development at some time

in the future. Other factors affecting toxicity such as pH, dissolved organic carbon, and sulfate would have to be tracked and their impacts on toxicity accounted for. GSL water is scored a 5 because of the identified uncertainties.

GSL water is representative of current conditions in GSL and is scored a 10. GSL water has several physiochemical parameters that are known to decrease toxicity and accurately accounting for these effects is important. GSL water is scored a 10 because the potential changes over time were addressed by the preceding DQO.

GSL water is anticipated to be stable relative to the concentrations of ions. The potential for biologically mediated changes are unknown. Based on the difficulty in obtaining and shipping the water, GSL water would have to be stable over time to allow large quantities to be collected and stored. Protocols would need to be established that verify the stability of GSL water over time. GSL water is scored an 8 for this DQO.

GSL water can be replicated by any laboratory provided that protocols are developed for collecting and storing the water. The protocols should also establish the tolerance ranges for key parameters. GSL water is scored a 10 for this DQO.

GSL water is anticipated to be costly because of the logistical issues of collection presumably by boat from the more remote areas of the Lake and the cost of shipping large quantities of water. Storage of the water may be inconvenient and the collection of additional water may be restricted by factors such as weather and/or season. GSL water is scored a 4 for this DQO.

Barnes and Wurtsbaugh, 2015

Barnes and Wurtsbaugh (2015) prepared a medium with salinity concentrations ranging from 10 g/l to 275 g/l. The media were prepared using equal parts deionized and GSL water and an inorganic salt mix consisting of 84% Instant Ocean, 14% NaCl, and 3% K₂SO₄. Final salinities and major ion concentrations were measured.

The Barnes and Wurtsbaugh (2015) medium would require similar analyses as the GSL water because it includes GSL water to meet USEPA approval. In addition, Instant Ocean is not a currently approved artificial salt mix for conducting bioassays. Presumably, USEPA could approve the use of this mix provided that protocols were developed to document the contents of the final solutions. This media is scored a 7.

The Barnes and Wurtsbaugh (2015) medium was used successfully to conduct experiments on brine shrimp. The suitability of this media for culturing algae or brine flies is unknown resulting in a score of 8.

Because the Barnes and Wurtsbaugh (2015) medium contains GSL water, the same concerns regarding confounders applies but to a lesser degree because only ½ of the liquid portion of the media is GSL water. The Instant Ocean has trace concentrations of the same inorganic substances that will be tested for the bioassays. Figures 2 through 8 compare the concentrations of metals in Instant Ocean at 120 ppt versus GSL water when data were available for both media. Note that none of the media considered is either 100% Instant Ocean or 100% Marinemix. With the exception of the DWQ Marinemix medium, this assumption is not anticipated to significantly

overestimate the metals concentrations because metals were not measured in all materials by Barnes and Wurtsbaugh (2015) or Belovsky.

The Instant Ocean concentrations were scaled up from the data of Hovanec and Coshland (2002). Atkinson and Bingman (1999), the only source for Instant Ocean trace metals that was peer reviewed, reported much higher trace metal concentrations than Hovanec and Coshland (2002) whose concentrations are corroborated by the data reported by Marulla and O'Toole (2005). As noted by Hovanec and Coshland (2002), the ICP/MS analytical method that they employed is considered more reliable than the ICP used by Atkinson and Bingman (1999). The comparisons were subsequently based on the data from Hovanec and Coshland (2002). The GSL data were reported by Adams et al., (2015) or were based on 2 years of recent sampling by DWQ. The GSL data were not normalized to 120 ppt salinity. The figures also include the same data for Crystal Sea Bioassay Laboratory Formula Marinemix which was provided by the manufacturer.

The following observations are based on a qualitative analysis of the comparisons shown on Figures 2 through 8. Arsenic concentrations are higher in the lake than in the Marinemix (Figure 2). No data were found for the Instant Ocean arsenic concentrations. Cadmium concentrations in Instant Ocean are higher than Marinemix or GSL (Figure 3). Copper concentrations are similar between Marinemix, Instant Ocean (not detected), and GSL (Figure 4). Lead (Figure 5) and zinc (Figure 9) concentrations in Marinemix are higher than Instant Ocean or GSL. Mercury concentrations are similar between Marinemix and GSL but no data were available for Instant Ocean (Figure 6). Nickel concentrations in Instant Ocean are higher than Marinemix and GSL (Figure 7). Selenium concentrations are higher in Marinemix than GSL but no data were available for Instant Ocean (Figure 8). The media mixtures using Instant Ocean and Marinemix were was assigned a score of 7.

The Barnes and Wurtsbaugh (2015) media can be replicated over time. Although the medium would have some of the same potential deficiencies as were discussed for GSL water, the salinity can be increased with the salt mixture which would negate the impacts of varying salinity in the lake. However if other parameters in the GSL water changed significantly, the media may not easily replicated over time. The media was assigned a score of 7.

The Barnes and Wurtsbaugh (2015) media is reasonably similar to GSL water with respect to major ion concentrations (Table 2). However, the media has about half Ca and Mg as GSL water and higher concentrations of Cl and SO₄. The dissolved organic carbon concentration is not reported but is likely lower than the 7 to 42 mg/l reported by Brix et al. (2006) and Wurtsbaugh and Jones (2012), respectively. This media was scored 6 primarily due to the uncertainties regarding organic carbon.

The Barnes and Wurtsbaugh (2015) medium is presumed to be stable based on the duration of the microcosm experiments conducted. The medium was assigned a score of 10. The medium could also be replicated by a qualified laboratory and was assigned a score of 10. For cost and convenience, the medium was assigned a score of 6 because compared to 100% GSL water, half as much water is needed.

Table 2 excerpted from Barnes and Wurtsbaugh (2015)

Table 1. Ionic weight proportions of the Gunnison Bay (328 g L⁻¹) and Gilbert Bay (110 g L⁻¹) of the Great Salt Lake (from Sturm 1980) and measured ionic composition of water from six of the salinity treatments in the microcosm experiment.

	Great Salt La	Microcosm Experiment (Nominal Salinities)						
Major Ions	Gunnison Bay (328 g L ⁻¹)	Gilbert Bay (110 g L ⁻¹)	10 g L ⁻¹	50 g L ⁻¹	100 g L^{-1}	150 g L^{-1}	200 g L^{-1}	250 g L^{-1}
Na ⁺	.320	.313	.350	.330	.307	.314	.313	.320
K^{+}	.026	.027	.017	.025	.024	.025	.024	.025
Mg ⁺²	.032	.035	.037	.020	.017	.016	.016	.016
Ca^{+2}	.001	.002	.004	.001	.001	.001	.001	.001
Cl	.554	.551	.502	.538	.571	.562	.563	.562
$\mathrm{SO_4}^{-2}$.067	.073	.090	.086	.080	.082	.083	.076

Belovsky

Belovsky has successfully used a 60:40 mix by volume of Morton Solar Salt and Instant Ocean to lab to culture brine shrimp and brine flies for the proposed bioassays. The Morton Solar Salt is commercial water softening salt extracted from GSL and the ion concentrations are as reported by the manufacturer. Both Belovsky and Wurtsbaugh use Instant Ocean in their media combined with either GSL water or salt extracted from GSL water and the following includes a specific discussion only where the scores or rationale differ.

Belovsky's medium has been successfully used to culture GSL algae, brine shrimp and brine flies and was assigned a score of 10. Belovsky's medium was assigned a score of 6 for potential confounders. No data for trace metals were available for the Morton Solar Salt and the score could increase or decrease if this data were obtained.

A comparison of the major ions in Belovsky's medium to GSL water is shown in Figure 10 for assessing the representativeness compared to GSL water. Belovsky's media has more NaCl and less K, SO₄, Ca, and Mg than GSL water. These concentrations were estimated assuming that 60:40 ratio by volume was equivalent to 60:40 by weight and the scores could change based on actual analytical or more refined nominal estimates.

Belovsky's medium is the least expensive medium considered and uses salts that are easily obtained and was scored 10 for cost and convenience.

DWQ Marinemix

A DWQ proposed medium consisting of Marinemix, NaCl and KCl in ratios of 84:13:2, respectively was attempted. This medium may have matched GSL water better than Barnes and Wurtsbaugh (2015) or Belovksy but when the medium was attempted, a precipitate formed and preliminary testing indicated lower survival of brine flies and brine shrimp, potentially in part due to reduced algal production. Jim from EPA unsuccessfully attempted to permanently dissolve the precipitate by reducing the pH. After consulting with the Marinemix manufacturer, Belvosky

reported that at salinities above approximately 5%, calcium would precipitate. This medium was abandoned because of the precipitate. The precipitation also suggests that the Marinemix (USPEPA approved) could not simply be substituted for Instant Ocean in Barnes and Wurtsbaugh's (2015) or Belovsky's media recipes.

DWQ Round Robin

DWQ initiated a laboratory round robin study to verify the analytical methods commonly used for analyzing the GSL samples. The initial matrix was formulated using reagent grade salts only and precipitates formed at salinities well below 12%. Different chemical forms of the salts were attempted and the medium shown in Table 3 did not exhibit a precipitate when anhydrous salts were used. Precipitates did form at salinities above 13% and the SO₄ was reduced by 50% as shown in Table 4 to prevent precipitation up to a salinity of 20%. The medium described in Table 4 was not considered further because the major ion concentrations deviate substantially from GSL. With salts of sufficient quality (e.g., laboratory grade), the medium in Table 3 would be approvable by USEPA. The primary unknown with this media is if the test organisms would tolerate it and if potential nutrient deficiencies exist. Because this is unknown, a score of 3 was assigned which could change if the organism were tested.

No potential confounders are identified and the medium was scored a 10. The medium would be replicable over time. The media is representative of GSL for the major ions but is lacking dissolved organic carbon, so was scored a 7. The potential addition of organic carbon could potentially change this score. The medium was stable over a couple of weeks but longer-term stability was not tested, so a score of 9 was assigned. The media can be replicated by any qualified laboratory and was scored a 10.

The salts are available from laboratory suppliers but the costs will be higher than e.g., Instant Ocean. Large term storage may require conditions to preserve the anhydrous condition of the salts resulting in a score of 6.

Page 8

					Table 3					
			DWQ Ro	und Robin	Media Matr	rix at 13% S	alinity			
Salt	Added (g/l)	[Na+]	[Mg++]	[K+]	[Ca++]	[H+]	[CI-]	[SO4=]	[CO3=]	[OH-]
CaCl2+2H2O	1.1				0.299944		0.530661			
MgCl2 (anhy)	8.1		2.06775				6.032335			
NaCl	106.56	41.91779					64.64221			
KCI	5.38			2.821532			2.558468			
CaSO4										
MgSO4 (anhy)	11.6		2.342456					9.258026		
Na2SO4										
K2SO4										
NaHSO4										
CaCO3										
MgCO3										
Na2CO3										
K2CO3										
NaHCO3	0.65	0.177883				0.007737			0.464379	
Ca(OH)2										
Mg(OH)2										
NaOH										
КОН										
Mass		0.317328	0.033245	0.021269	0.002261		0.556049	0.069789		
Fraction										
Target Conc.	133.39	42	4.4	2.8	0.3	1E-08	74	9.3	0.43	0.000017
(g/l)										
% of Target		100.2	100.	100.8	100.0		99.7	99.5	108	

`

Table 5Qualitative Scores for Bioassay Media Considered								
Data Quality Objective	GSL Water	Barnes and Wurtsbaugh (2015)	Belovsky	DWQ Round Robin				
USEPA Approval	7	7	7	10				
Media supports test organisms	10	8	10	3				
Media has minimal potential confounders affecting the toxicity	5	7	6	10				
Media can be replicated over time	5	7	7	10				
Media is representative of Great Salt Lake	10	6	6	7				
Media is stable over test duration	8	10	10	9				
Media can be replicated at any laboratory	10	10	10	10				
Low Cost and Convenient	4	6	8	6				
SUM without cost score	55	55	56	59				

References

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Marulla, M. and T. O'Toole. 2005. "The Inland Reef Aquaria Salt Study, Part II." Advanced

Wurtsbaugh, W.A. and E.F. Jones. 2012. The Great Salt Lake's Deep Brine Layer and Its Importance for Mercury Bioaccumulation in Brine Shrimp (Artemia franciscana). Final Report to the Utah Division of Forestry, Fire and State Lands. May 22. http://digitalcommons.usu.edu/cgi/viewcontent.cgi?article=1547&context=wats_facpub

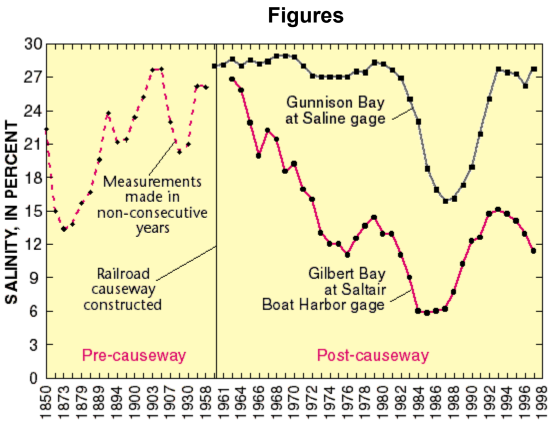


Figure 1. Great Salt Lake Salinity over time from USGS (http://ut.water.usgs.gov/greatsaltlake/salinity/)

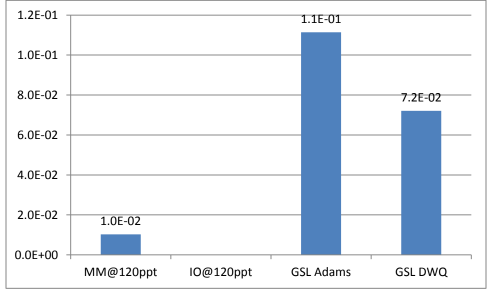


Figure 2. Comparison of mean arsenic concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality. No data available for Instant Ocean (IO).

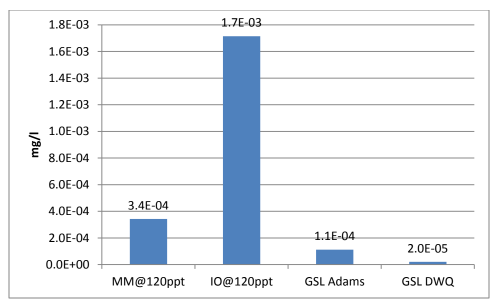


Figure 3. Comparison of mean cadmium concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality

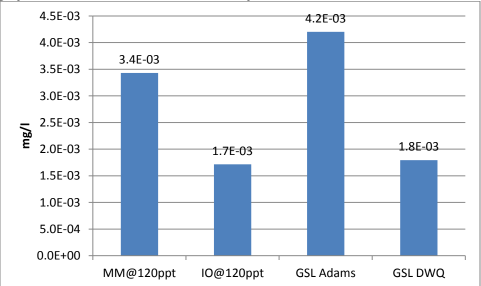


Figure 4. Comparison of mean copper concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality. Copper was nondetect for Instant Ocean.

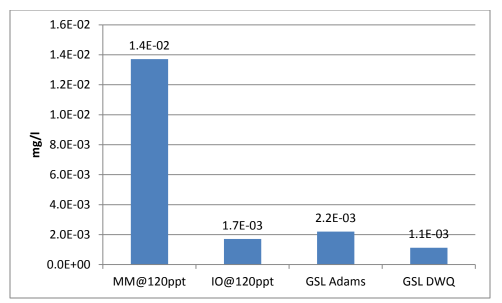


Figure 5. Comparison of mean lead concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality. Lead was nondetect for Instant Ocean.

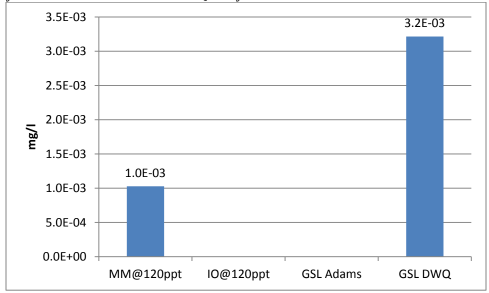


Figure 6. Comparison of mean lead concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality. No data were available for Instant Ocean or GSL Adams.

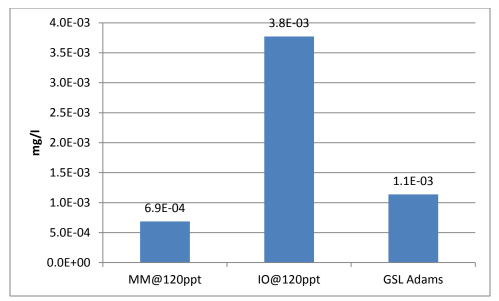


Figure 7. Comparison of mean nickel concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality

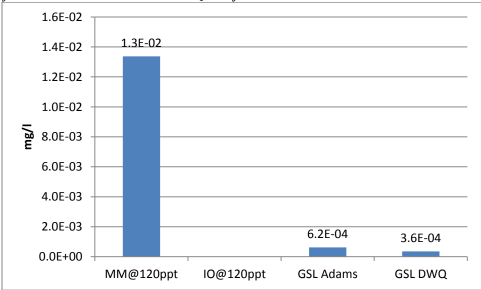


Figure 8. Comparison of mean selenium concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality. No data were available for Instant Ocean.

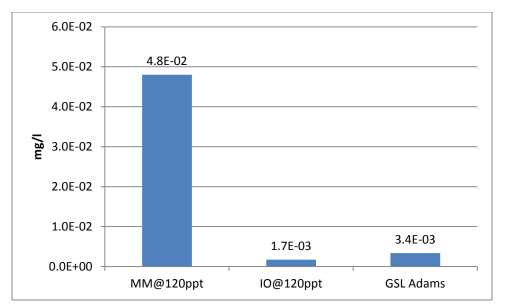


Figure 9. Comparison of mean zinc concentrations. MM=Marinemix, IO=Instant Ocean, GSL Adams=Great Salt Lake Adams et al.,2015, GSL DWQ= Great Salt Lake based on 2 years of sampling by the Utah Division of Water Quality. Zinc was nondetect for Instant Ocean.

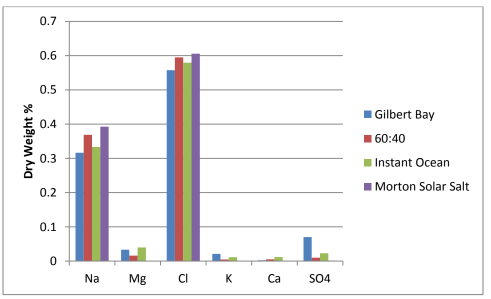


Figure 10. Comparison of major ions between Great Salt Lake (Gilbert Bay), a 60:40 Morton Solar Salt : Instant Ocean assuming volume is accurate surrogate for mass, Instant Ocean and Morton Solar Salt

APPENDIX 2

Results from Initial Range Finding for Copper and Brine Shrimp

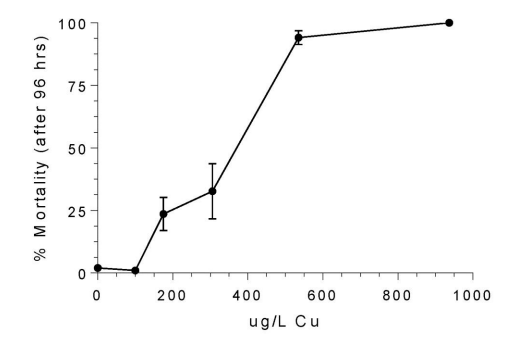
The pre-test water samples were collected after the water had been renewed, but prior to the addition of algae and brine shrimp each day as to hopefully ensure the correct copper concentrations were in solution.

The post-test water samples were collected each following day, prior to water renewal, as why those samples start at 24 hours. These samples were filtered in order to remove the algae.

The blanks were one replicate that was treated the same as all other replicates (water renewal each day and algae present), but it did not have brine shrimp in solution. Water samples were taken from these replicates after each 24 hours (post-test) to monitor the copper concentrations without brine shrimp in solution.

Ultra-pure nitric acid prior to shipment to the lab and were clear at the time of shipping.

96-Hr Acute Copper Assay Results



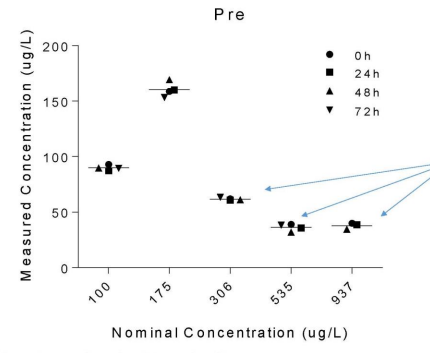
Observations:

• Mortality increased with increased nominal concentrations of copper

 However, measured copper concentrations are not in agreement with nominal values.....see slides 2-5

1

Measured Copper Concentrations <u>Pre-test</u> for Each Day



Observations:

• Background copper in control solutions varied from 5.3-9.6 ug/L

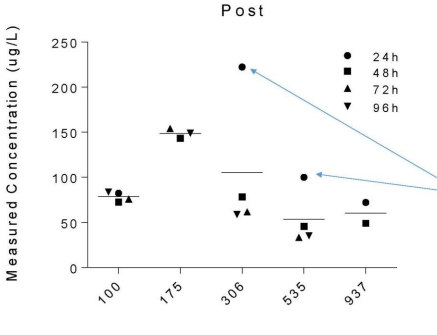
• Measured concentrations of copper at first 2 concentrations were acceptable

• The three highest concentrations had lower measured concentrations than expected <u>Possible Explanation:</u> Precipitation and removal via filtration

2

*All samples were filtered via 0.2um syringe filters

Measured Copper Concentrations <u>Post-test</u> for Each Day



Nominal Concentration (ug/L)

Observations:

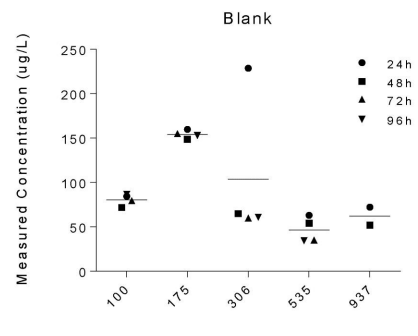
• Similar trends in measured copper across the range of concentrations

• Reduction from pre-test concentrations, especially for 175 ug/L

Possible explanation: Accumulation in algae (see slide 5)

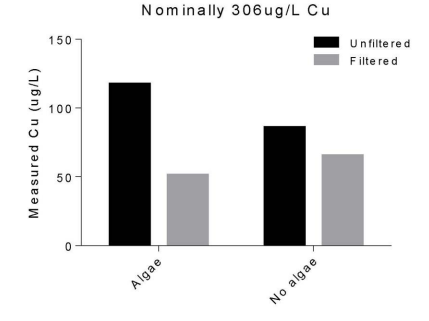
• Higher concentration than pre test for certain measurements

Measured Copper Concentrations **Post-test (Blanks)** for Each Day



Nominal Concentration (ug/L)

Filtered vs. Unfiltered Samples



Observations:

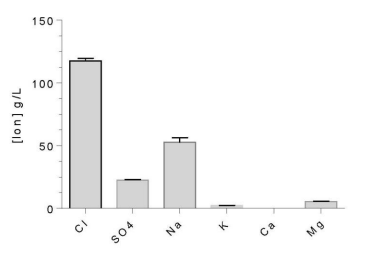
• Filtering the sample via .2um has an effect on measured concentrations

• Copper appears to be associated with algae, either apically bound or accumulated

• Measured copper is lower than target concentration

lon	Average in lake (g/L)	Average measured (g/L)	Difference
Cl	74	116.5	42.5
SO4	9.3	22.8	13.5
Na	42	55.8	13.8
К	2.8	2.3	-0.5
Ca	0.3	0.34	0.04
Mg	4.4	5.3	0.9

Major Ion Concentrations



A run of background metals will be important moving forward. These initial samples were with a lower grade NaCl than what we are using now

6

APPENDIX B

ANALYTICAL CHEMISTRY INFORMATION

-ead (as lead nitrate)	
issL - Definitive Brine Shrimp study with L	lename: analytical_Pb_definitive2.xlsx

Page / or 3

		Dissolved	10 / 01	100%	104%		%26		68%		35%		16%	2
		Dis		+		-	_		_	-			-	Ļ
			E E	-	7%/		11%		10%	2	18%		35%	
		Diss	%		%6		%6		7%		11%		18%	2
		Total	% of nominal		81%		82%		82%		%06		78%	2
			AVG Diss	53.0	4 398		7.075		9.563		25.150		65 650	
			AVG Total	53.0	38.925		65.700		109.500		199.250		288.500	
8	PIO	54	Dav 4 Diss - old	53 U	5.600	,	6,260	6,080	7,150		10.200			53 U
020918	DID	T4	Dav 4 Total - old Dav 4 Diss - old	53 U	38,300		65,600		107,000		200,000			53 U
	New	D3	2 Total - new Day 2 Diss - new	53 U	6,370	6,490	7,920		12,400		42,400			
020718	New	13	Dav		39,500		65,900		107,000		203,000			
18	PIO	D2	Day 2 Total - old Day 2 Diss - old	53 U	2,870		6,780		7,600		12,500		18,300	
020718	PIO	12	Day 2 Total - old	53 U	39,400		67,400		113,000		201,000	204,000	252,000	
8	New	D1	Initial Diss	53 U	2,750		7,340		11,100	11,100	35,500		113,000	
020518	New	11	Initial Total	53 U	38,500		63,900		111,000		193,000		325,000	
		Nominal	conc. (ug/L)	-	48,000		80,000		134,000		222,000		371,000	MQ Blank
			Sample ID	SP1	SP2	SP2 dup	SP3	SP3 dup	SP4	SP4 dup	SP5	SP5 dup	SP6	ΤQ

Note: Calculations (e.g., % nominal) may appear slightly different due to rounding differences. U = < MRL / MDL

RSD 1.48%			
RSD 0.00%			
	1.0 U		1.0 U
	Control		Control
DOC	SP1	TOC	SP1

RSD 1.87%

RSD 2.92%

ł

TAN MAN 3/29/18

Long list of metals for arsenic definitive study filename: analytical_Pb_definitive2.xlsx

No Food	SP1.0hN-T/D(NF))	Food	SP1.0hN-T/D(F)
	ug/L	ug/L		ug/L
	<u>TR</u>	<u>Diss.</u>		TR
Antimony	1.0 U	1.0 U	Antimony	1.0 U
Arsenic	0.50 U	0.50 U	Arsenic	0.50 U
Beryllium	0.020 U	0.020 U	Beryllium	0.020 U
Cadmium	0.020 U	0.020 U	Cadmium	0.026
Chromium	0.56	0.25	Chromium	a 37.2
Copper	0.41	0.28	Copper	4.93
Lead	0.077	0.032	Lead	0.898
Mercury	0.20 U	0.20 U	Mercury	×
Nickel	22.0	22.0	Nickel	23.4
Selenium	1.0 U	1.0 U	Selenium	₿⁄
Silver	0.044	0.020 U	Silver	0.208
Thallium	0.102	0.022	Thallium	0.465
Zinc	9.68	6.21	Zinc	52.6

U = < MRL / MDL

ARLIMOL & insufficient volume to analyze these two analytes

GSL - Definitive Brine Shrimp study with Lead (as lead nitrate) filename: analytical_Pb_definitive2.xlsx comparison of orginal vs split analytical samples

Page $\frac{2}{3}$ of $\frac{3}{2}$

vilats wan to

			% Precision (RSD)				7.45%		15.52%		3.56%		20.59%
	Old	T4	av 4 Total - old					107.000	125,000				
020918	New (T4	Day 2 Total - new Day 2 Diss - old Day 4 Total - new Day 4 Total - old % Precision (RSD)	0.077	<1.02 U								
8	Old	D3	Day 2 Diss - old									18,300	22,500
020718	New	13	Day 2 Total - new			65,900	71,000						
020518	New	Τ1	conc. (ug/L) Day 1 Total - new							193,000	200,000		
		Nominal	conc. (ug/L)	0	0	80,000	80,000	134,000	134,000	222,000	222,000	371,000	371,000
				original	split	original	split	original	split	original	split	original	split
			Sample ID	Control	Control	SP3	SP3	SP4	SP4	SP5	SP5	SP6	SP6

Note: Calculations (e.g., % nominal) may appear slightly different due to rounding differences. U = < MRL / MDL

split samples analyzed at Brooks Applied Labs

original samples analyzed by ALS Labs

nb: control waters analyzed without the addition of algae

APPENDIX C

BIOLOGICAL TEST DATA

Page 1 of _____ QA Form No. 051 Revision 5 Effective 02/14

TOXICITY DATA PACKAGE COVER SHEET

an: m3/30/18

Test Type:	Acute	Project Number:	17001-474-018
Test Substance:	Pb(NO3)2		ia franciscana
Dilution Water Type:	120 ppt rGSL	Organism Lot or Batc	×
Concurrent Control Water Typ		Age: 27.24 (<72 h	
Date and Time Test Began:	2/8/18 @1530-1710	Date and Time Test E	
Protocol Number:	A	Investigator(s):	1000 4111 @ 1605
Background Information		11001igator(s). <u>1-9-</u>	who have and
Background Information			
Type of Test:	Static - Renewal	pH control?: <u>Yes</u> If yes, give % CO ₂ :	<u>No</u> N/A
Test Temperature:	20 ± 1 °C	Env. Chmbr/Bath #: _1	
Photoperiod:	16 h light : 8 h dark	Light Intensity: <u>50 -100</u>	
Test Solution Vol.:	150 ml	Number of Replicates	
Length of Test:	96 h	Number of Organisms	
Type of Food and Quantity per	Chamber: see food sheet	Feeding Frequency:	
Test Substance Characterization	on Parameters and Frequency:		<u> </u>
Dead the second		itiation TRC: In	Notion-
••	Conductivity: <u>Daily* & Termination</u>		
Test Concentrations (Mass:Volu		, 222, and 371 mg/L Pb (n	
Agency Summary Sheet(s)?:	NA	-222, and 37 1 mg/2 PD (n	ominal)
	ztetie -	7.21	
	Test Dates: to	128/19	LC50: 77.9 maddl
Hist. 95% Control Limits: 5	<u>8.97 to 87.16</u> Method	for Determining Ref. Tox.	Value:Probit
Special Procedures and Cons	iderations:		
* Measure chemistries at test ter	minotion exclanation		
	mination or when 100% mortality is c	observed in a treatment.	
Study Director Initials:			
Vo		7	
(DA3N 2/5/18CF		

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TEST SUBSTANCE USAGE LOG

Project Number:

Project Number:17001	-474-018		C	A: DAP 10/29
	Sample 1	Sample 2	Sample 3	Sample 4
Test Substance Number	017-020	, 2		•
Test Substance Stock Solution Preparation Date	5	spile shell		
Sample Type	Propared in L			
Date Test Substance Received	N/A			
Dilution Water Number RW# or TRE#, circle one	13175			
Concurrent Control Water RW#	NA			
Date(s) Used	2/5/18 2/1)18			

Preparation of Test Solutions

Test	Test	Dilution	Total				Test	Dilution	Total
Substance	Substance	Water	Volume				Substance	Water	Volume
Conc. (mg/L	Volume	Volume	(ml)				Volume	Volume	(ml)
As Nominal)	(ml)	(ml)			ρ		(ml)	(ml)	
0					see As spiking sheet	·			
48					() "				
80					11				
134					11				
222					H				
371					17				
								_	
Total	0	0	0						
Initials / Date	WZN	Brines	ring act	fe	2/5/18				ť
Initials / Date			/						
Initials / Date	from "	2/118						· · · · · · · · · · · · · · · · · · ·	
Initials / Date						Ī			
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Initials / Date								······	

OUTN 2/5/18E

Page <u>3</u> of <u>19</u> QA Form No. 052 Revision 3 Effective 02/14

ACUTE BIOLOGICAL DATA am An 3/30/18 Project Number: 17001-474-018 Test Species: Artemia franciscana Number of Surviving Organisms (mg/L)Test 0 24 48 72 96 Conc. Replicate Hours Hours 48 5 Hours Hours Hours 964 Remarks 0 20 А 5 97 7 M Ð, . 85 0.85 20 В \mathbb{Z} 2 20 0 1 20 С え \mathcal{O} 20 ,9790 1 96% D 20 2 O. 20 1 mont Е 20 1 0.952 48 20 А 19 Ø .95 .95 20210 20210 В 7 20 С 18 5 97% 758 .90 .85 20 D 10 2 0 20 Е 20 Q .95 20 80 20 А 9 .95 20 В 70 0 1 20 С 99% 70 96.29 O1 20 D Q Ģ .95 22 72 Е 20 .91 -20 134 А XINF 19 3 \bigcirc 10 В 20 L $\overline{\mathcal{V}}$ С Х XINF 18 95% 55% 20 う D $\boldsymbol{\chi}$ XINF 19 W La × Е XINF r 20 Х 222 Z А 8 XIND N В 0 Ĺ 20 С 0 2 86% 22% 0 20 S D W Е 6 371 20 А 6 0 -> 180 В 20 \bigcirc 0 % С 0 20 0 0% 20 D \bigcirc 0 20 Е 6 \mathcal{O} 25/19 2/6/18 24/13 2/9/18 Date: 1555 1420 1530-1710 Time: 1605

2 MW 2/1/18 E Obm 2/2/18 WP

o~/mm

hm

ΜŴ

REN/OAR

Initials:

Page 4 of QA Form No. 126a Revision 1 Effective 02/14

ACUTE CHEMICAL DATA

Project Number: 17001-474-018

QA: PSV 10/28/15

Test Species: Artemia franciscana

(mg/L)	<u> </u>		IEW				DLD		Meter # (All Conc.)					
	Day 0	Day 1	Day 2	Day 3	Day 1	Day 2	Day 3	Day 4	Day 0	Day	Day	Day	Da	
Conc.: 0	KUT	5/317	15	<u> </u>	Rep.	Rep.	Bep.	Rep.		1	2	3	4	
<u>(</u> эН	83	820	81		8.0	8.0				r				
D.O. (mg/L) 🕻	5.1	0.00	5.2		5.5	8.0	<u>8.1</u> 54	8.1 5.2	PM24	FM210			HMZ	
Гетр. (°С)	20		20		21	21	197	20,2	L-13	LDO		400		
Cond. (µS/cm)	195400		122300		<u> </u>			13190		15	L-30 15	Los	40	
Hard. (mg/L)	12900						1	Horro	<u> </u>				15	
Alk. (mg/L)	187										Titration Titration			
RC (mg/L)	- NA								#22		Titration			
IH3 (mg/L)	NA				. 9 9				HAPI					
alinity (ppt)	122								#1					
Conc.: 48										<u></u>	<u></u>			
H .O. (mg/L) 🕺			80		7.9	80	8.0	80						
	5.1		5.2		5.4	5.1	5.4	5.7						
emp. (°C)			20		21	20	-70	20						
ond. (µS/cm)	196320	 	23800					12820	2					
ard. (mg/L) lk. (mg/L)														
RC (mg/L)														
$H_3 (mg/L)$														
Conc.: 80				r										
	8.0		7,8		7.91	791	170	0 D						
	5.2		51				a il	60						
	20		26		<u>55</u> 21		<u>5.4</u> 20	\mathcal{D}						
	93100		2300		<u> </u>			20						
Conc.: 134				╘╼═╾┥			[/	302						
I	7.8		7.6		7.8	7.8	78	19		<u> </u>				
O. (mg/L) 🖌 .	5.2		5.1				5.42							
	20		20			$\frac{1}{20}$	25	$\frac{5}{21}$						
nd. (µS/cm)	90500		13600					2980						
Date: 2		Ž.	118		2/2/18/0	hhar	AleR	AI	<u>·</u> ∔					
Time: 1			30		A	lordi	• • • •							
Initials:	SU		om			mi								

Note: Hardness, alkalinity, TRC, and NH3 data appearing on this page have been transcribed from the wet chemistry log, QA Form No. 084

Page 5 of $\int Q$ QA Form No. 126b **Revision 1** Effective 02/14

QA: DAP 10/28/18

ACUTE CHEMICAL DATA

Project Number: 17001-474-018

Test Species: Artemia franciscana

		-	IEW			(DLD			Mete	r # (All C	Conc.)	
(mg/L)	Day 0	Day 1	Day 2	Day 3	Day 1	Day 2	Day 3	Day 4	Day 0	Day 1	Day 2	Day 3	Day
Conc.:					Rep.	Rep.	Rep.	Rep.		<u> </u>	2		4
рН													
D.O. (mg/L)							<u> </u>						
Temp. (°C)													
Cond. (µS/cm)													
Conc.:													
рН													
D.O. (mg/L)													
Temp. (°C)													
Cond. (µS/cm)													
Conc.:							·	<u>ار ، ، ، ، ، ، ، ، ، ، ، ، ، ، ، ، ، ، ،</u>	<u> </u>				
pН													
D.O. (mg/L)													
Temp. (°C)													
Cond. (µS/cm)													
Conc.: 222													
pH	7.3		7.7		7.5	77	76	7.8					
D.O. (mg/L) 🗡	52		5.0		5.3	47	<u>Ş</u> Ÿ	üq					
Temp. (°C)	20		20		21	20	21	76					
Cond. (µS/cm)	1920	Ø	24700		4	74		337	,				
Conc.: 371													
рН	7.0				7.2	74							
D.O. (mg/L) 🔻	5.2				52	<u><u>u</u><u>r</u></u>							
Temp. (°C)	20		820		21	20							
Cond. (µS/cm)	18650	Ø			14000	51288	Ċ,						
lard. (mg/L)													
Alk. (mg/L)													
FRC (mg/L)													
NH3 (mg/L)													
Salinity (ppt)													
Date:	2/5/18	2	7/18		ZKel18	mat	48/18	2018					
Time:			احتما		1550 MW -	645	120	1610					
	NZN		em		~~~	<u> ~ ~ ~ </u>	200	D~					

Note: Hardness, alkalinity, TRC, and NH3 data appearing on this page have been transcribed from the wet chemistry log, QA Form No. 084

Page 6 of $\underline{12}$ QA Form No. 055 Revision 3 Effective 02/14

DAILY TOXICITY TEST LOG

		a	A: MR 3/301
Project Num		<u>~</u>	n: M2 51501
Test Species	S: Artemia franciscana		
General	Measured salinity of rGSL water: <u>122</u> ppt	Feeding	Initials/Date
Comments			initials/Date
Test Day 0	Random ChartMulconThermometer #:M-/STest Solution Mixed at: $7200 - 1220$ $1200 - 1220$ $1200 - 1220$ Food Added at: $530 - 120$ $1200 - 1220$ $1200 - 1220$ Test Organisms Added at: $1530 - 120$ $1200 - 120$ TakeDO $1200 - 120$ $1200 - 1220$ Real Time Temp.= $19 ^{\circ}C$ Range = $19 - 200 ^{\circ}C$	yes cea food prop sheet	1250) 2/5/18
		none	MW 21(d18
Test Day 2	Real Time Temp.= 19°C Range = 19-20 °C Take D2 New & Bld analytical USed New Plostocic Real Time Temp.= 19°C Range = 19-20 °C	yes See food prop sheet NOne	em 21/113
est Day 3		Nove	2/3/10
est Day 4	Real Time Temp.= 1 °C Range = 1 °C °C	None	24113
est Day 5			
est Day 6			
est Day 7			
est Day 8			

(24: An 3/30/18/ Page 2 of 19 Situa 1220 (white) Situa 1220 (white) Start Spihing @ 1120 -1150 @ 217118 Smm stat ale 1220 100 TR 50 Diss $\frac{(\text{measured, nominal was 4,000 mg/L})}{3900.00} \frac{(C12 - 020)}{\text{mg/L Pb}} \frac{(C12 - 020)}{(C12 - 020)} \frac{(C12$ 81/2/2 dilution series Jumi of stade laft Take 50 ml for QC dup - D2 0.0 Definitive TEST 9.5% % spike of vol 18 Volume per treatment (L) 1.0 Total Vol(L) 1.000 1.000 1.000 1.000 1.000 0.0 0.00 ~ 219.33 95.13 57.08 34.25 20.55 12.33 Stock (m) **Brine Shrimp Acute Studies** Conc. ug/L 371,000 2222,600 133,560 80,136 80,136 48,082 0 Primary stock @ Feb 2018 0 u 4 u u Total Trtmnt ~

Survival of brine shrimp in definitive Lead study 17001-474-018

Page 7 of 19 122 3/29/18 QA: 040 10/28/18

8-h Survival	4			6-h Survival	9		
ave		avg					
<u>% survival</u> % surviva	<u># @ 48-h</u>	<u>% survival</u>	<u>% survival</u>	<u>#@96-h</u>	<u>initial #</u>	<u>rep</u>	<u>Pb trtmnt</u>
85.0%	17		85.0%	17	20	а	0
100.0%	20		100%	20	20	b	
100.0%	20		100%	20	20	С	
100.0%	20		100%	20	20	d	
100.0% 97.00%	21	96.05%	95.2%	20	21	e	
95.0%	19		95%	19	20	а	48
100.0%	21		100%	21	21	b	
90.0%	18		85%	17	20	С	
100.0%	20		100%	20	20	d	
100.0% 97.0%	20	95.0%	95%	19	20	е	<u> </u>
100.0%	20		95%	19	20	а	80
100.0%	20		100%	20	20	b	
100.0%	20		100%	20	20	С	
95.0%	19		95%	19	20	d	
100.0% 99.0%	22	96.2%	91%	20	22	e	
95.0%	19		50%	10	20	а	134
100.0%	20		70%	14	20	b	
90.0%	18		55%	11	20	С	
95.0%	19		65%	13	20	d	
95.0% 95.0%	19	55.0%	35%	7	20	е	
90.0%	18		25%	5	20	а	222
80.0%	16		20%	4	20	b	
80.0%	16		15%	3	20	С	
90.0%	18		45%	9	20	d	
90.0% 86.0%	18	22.0%	5%	1	20	е	
0.0%	0	······································	0%	0	20	а	371
0.0%	0		0%	0	20	b	
0.0%	0		0%	0	20	С	
0.0%	0		0%	0	20	d	
0.0% 0.0%	0	0.0%	0%	0	20	е	

19001-474-018

	0.040609	P1-4 •
	0.052594	P3-1 ●
	0.058357	P2-5
	0.064117	P5-1
5	0.111139	P4-2 •
	0.155116	P2-2 •
	0.180092	P6-4∙
	0.214803	P4-5●
	0.241688	P1-1•
10	0.258089	P6-5 •
	0.260786	P3-5 •
	0.316961	P1-5 •
	0.331968	P3-2 •
	0.348368	P1-2 •
77	0.375865	P5-5 •
•	0.385366	P4-1
	0.387737	P6-2 🖕
	0.399265	P2-4 •
	0.462034	P2-3 •
71	0.46881	P2-1
	0.474399	P5-3 •
	0.490044	P5-4 /
	0.505853	P6-1 ●
2	0.658912	P6-3 •
	0.669352	P3-4 •
	0.690513	P4-3 •
	0.894144	P4-4 •
	0.90844	P1-3 🖡
	0.953544	P5-2 *

Random Chart for Pb definitive study

0.975395 P3-3

-018 Rep. 9 Jg 2/5/18 N3N QA: Der 10/29/18

Page <u>Lb</u> of <u>9</u> QA Form No.159 Revision: 0 Effective: 03/2015

Activities Log

DUNEN 14/2/18

Date	Time	Initials	Notes
2/3/18	1400	-DAP	Obtained a 1000 brine shamp haupling -200
			hauplii in each of 5 bockors. Each beaker containing a 2.72 ml of algal culture of 47.28 ml of 1656.
			rGSL Batch: RW 13168
			Algal culture = 070617 Nauplii were collocted from 2/3/19 noon batche Algae was collocted by
			Ashley Romano on 2/2/17 & held @ 10% in Chambor #1
\checkmark	\checkmark	V	in Chambon #1 Boatons with brine shrimp & algoe placod in isolated agranium in Bath #1, with goatle arrazion.

			Rige I of R	L R	
Predicting Chl a from Absorbance at 450 nm	Models Update on:	5/22/2017	GA: DOV	CA: Day is/29/18	ι.
⁵ redict. Chl a (ug/L) from abs at 450 nm (2.5 cm cuvette):	Chl a = 1092.5x - 96.878		Slope Y-inte 1092.5	Y-intercept -96.878	
Predict. Chl a (ug/L) from abs at 450 nm (1.0 cm cuvette):	Chl a = 1983.6x - 179.98		1983.6	-179.98	
Calc. Absorb. Conc	Calc. Chl a Concn. (ug/L)				
	-96.878				
Enter Absorbance (1.0 cm) here: 1.016	1835.3576				
From MoUN Collure					

Re-feeding (before test) calculation of Volume of D. viridis Culture Media Required for Feeding

21/22/01 000 : YOS Papele of 12

Intermediate Calculations

0.1

1.835358

Final Values

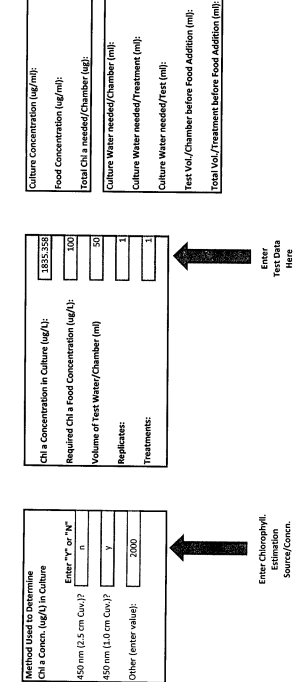
2.724265

47.27574

2.724265

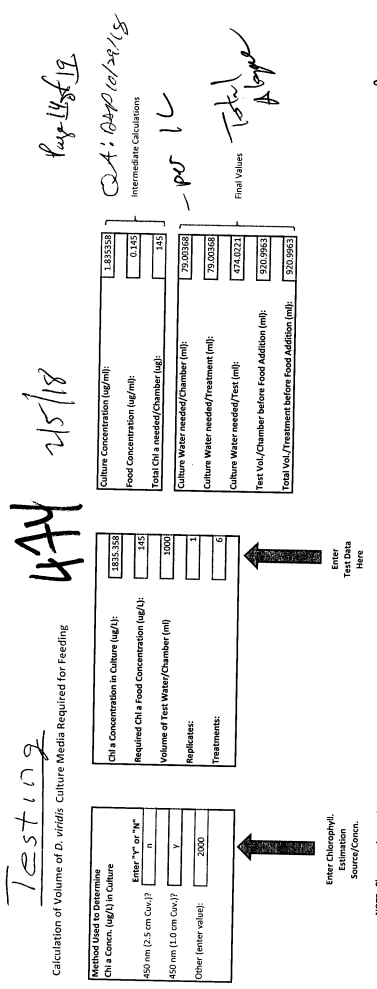
2.724265

47.27574



NOTE: Change values only in cells outlined in GREEN; do not change cells outlined in RED; these are equations

Predicting Chl a from Abcorboon at 200	-	2/5/18		Purp.	fup 13 . f19
	nuance at 450 hm	Models Update on:	5/22/2017	04:00	21/22/01 dro : 20
Predict. Chl a (ug/L) from abs at 450 nm (2.5 cm cuvette):	0 nm (2.5 cm cuvette):	Chl a = 1092.5x - 96.878		Slope Y-intercept 1092.5 -96.8	ept -96.878
Predict. Chl a (ug/L) from abs at 450 nm (1.0 cm cuvette):	0 nm (1.0 cm cuvette):	Chl a = 1983.6x - 179.98		1983.6	-179.98
Enter Absorbance (2.5 cm) here: Enter Absorbance (1.0 cm) here:	Absorb. 0 1.016	Calc. Chl a Concn. (ug/L) -96.878 1835.3576]



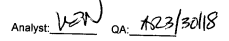
NOTE: Change values only in cells outlined in GREEN; do not change cells outlined in RED; these are equations

79 ml is earle

DAP Reeds 2 40 ml algre

	mia fran	<u>ciscar</u>	NR					port Date: st Code:	28		16:16 (p 1 of 2 3 01-2655-760
Mysidopsis	-96-h Acute Su	rvival Test	t			· · · · · · · · · · · · · · · · · · ·			TRE Env		ntal Strategies
Analysis ID Analyzed:	10-2908-59 29 Mar-18		-	96h Surviva Trimmed Sp		rber		TIS Versio	on: CETISv		
Batch ID: Start Date: Ending Date Duration:	12-9169-670 05 Feb-18 1 e: 09 Feb-18 1 4d 1h	5:30 6:05	Test Type: Protocol: Species: J	Survival (96) EPA/821/R-(h) 02-012 (200 Dahia //~}c		An	alyst: L uent: S ne: C	ab Tech Seawater Crystal Sea		
	16-6160-874 e: 05 Feb-18 1 te: 05 Feb-18 1 e: 4h	1:00 I 4:00 S		530A2324 Ambient San	nple		Clie	ent: G	iSL-ND pecial Studies		
	earman-Kärbe						$\overline{\bigcirc}$			1	314
Control Three		Threshold		Mu	Sigma		LC50 3	95% LC	L 95% UCL		
		0.0396	0.50%	2.114	0.015		130	121.3	139.3		
96h Survival Total PG	I Rate Summar Control Type	у			Cal	culated Varia	ate(A/B)				
	Control Type Dilution Water		Mean	Min	Max	Std Err	Std Dev	CV%	%Effect	Α	в
38.92	Dilution water	5	0.9605	0.85	1	0.02912	0.06511	6.78%	0.0%	97	101
65.7		5 5	0.95	0.85	1	0.02739	0.06124	6.45%	1.09%	96	101
109.5		5 5	0.9618	0.9091	1	0.01728	0.03865	4.02%	-0.14%	98	102
199.2		5 5	0.55	0.35	0.7	0.06124	0.1369	24.9%	42.7%	55	100
288.5		5 5	0.22	0.05	0.45	0.06633	0.1483	67.4%	77.1%	22	100
96h Survival	Rate Detail		0	0	0	0	0	······	100.0%	0	100
	Control Type	Rep 1	Rep 2	Rep 3	Ban 4	D			Proti		ta.
	Dilution Water	0.85	1	1 1	Rep 4	Rep 5					NA
8.92		0.95	1		1	0.9524			~ Aliv) one	
5.7		0.95	1	0.85 1	1	0.95			P	all' '	N Contraction of the second se
09.5		0.5		1	0.95	0.9091			X	1	
99.2		0.5	0.7	0.55	0.65	0.35			pe		
88.5		0.25	0.2 0	0.15	0.45	0.05					
	Rate Binomials			0	0	0					
-mg/L	Control Type	, Rep 1	Rep 2	Rep 3	Rep 4	Den C					
.053	Dilution Water	17/20	20/20	20/20	20/20	Rep 5					
		19/20	21/21	17/20		20/21					
8.92			~ 1/ ~	17720	20/20	19/20					
		19/20	20/20	20/20	10/00	00/0-					
5.7		19/20 10/20	20/20	20/20	19/20	20/22					
8.92 5.7 09.5 99.2		19/20 10/20 5/20	20/20 14/20 4/20	20/20 11/20 3/20	19/20 13/20 9/20	20/22 7/20 1/20					

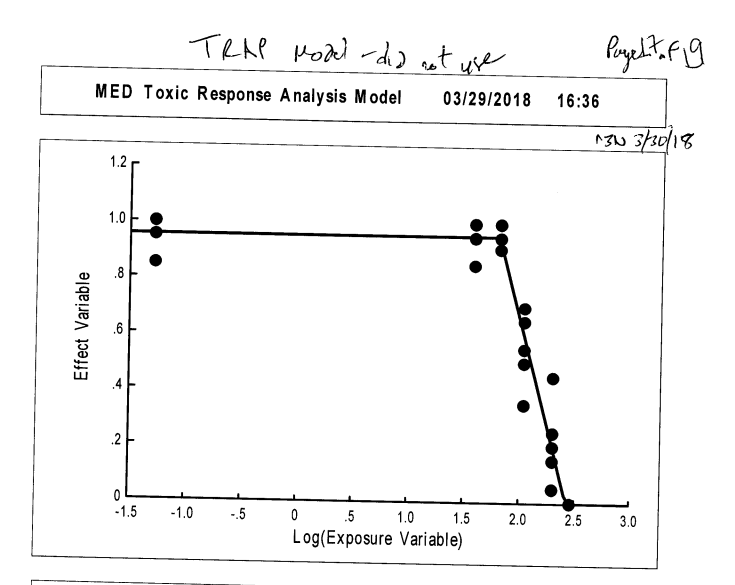
Ome for RISIN 3/30/18 9



CETIS Ana	llytical Report			Report Date: Test Code:	29 Mar-18 16:16 (p 2 of 2)
Mysidopsis 9	6-h Acute Survival T	est		Test Code;	474-018 01-2655-7609
Analysis ID: Analyzed:	10-2908-5999 29 Mar-18 16:16	Endpoint: Analysis:	96h Survival Rate Trimmed Spearman-Kärber	CETIS Version:	TRE Environmental Strategies CETISv1.8.7
Graphics				Official Results:	Yes
0.9 0.8 0.7 0.6 0.6 0.3 0.2 0.1 0.0 0.1 0.0 0.1 0.0 0.1 0.0 0.1 0.0 0.1 0.0 0.1 0.0 0.1 0.1	50 100	150 200	250 300		

C-mg/L

QA: Are 3/30/18 Ner Analyst:



	Parameter Summar	y (Piecewise L	inear Regress.	ion Analysis)	
Parameter	Guess	FinalEst	StdError	95%LCL	95%UCL
LogX 50	2.096	2.117	0.018	2.079	2.155
S	1.3257	1.6041	0.1321	1.3330	1.8752
Y 0	0.9575	0.9552	0.0284	0.8970	1.0134

Effect Concentration Summary								
%Effect	Xp Est	95%LCL	95%UCL					
50.0	130.98	120.06	142.90					
20.0	85.15	76.33	95.00					
10.0	73.77	65.06	83.64					
5.0	68.66	60.01	78.56					
0.0	63.90	55.32	73.81					

MED Toxic Response Analysis Model, Version 1.00

Ray 18 5 F 19 1310 3130/18

16:36

MED Toxic	Response	Analysis	Model	03/29/2018

Regression Analysis of Variance							
Source	df	SS	MS	F	Alpha		
Total	29	4.651	0.16038				
Regression	2	4.433	2.21671	275.	0.0000		
Error	27	0.218	0.00806				

		Data Summary		
Exposure	Obs Effects	Pred Effects	Residual	Weight
-1.276	0.850	0.955	0.105	1.
-1.276	0.952	0.955	0.003	1.
-1.276	1.000	0.955	-0.045	1.
-1.276	1.000	0.955	-0.045	1.
-1.276	1.000	0.955	-0.045	1.
1.590	0.850	0.955	0.105	1
1.590	0.950	0.955	0.005	1
1.590	0.950	0.955	0.005	1
1.590	1.000	0.955	-0.045	1
1.590	1.000	0.955	-0.045	1
1.818	0.910	0.937	0.027	1
1.818	0.950	0.937	-0.013	1
1.818	0.950	0.937	-0.013	1
1.818	1.000	0.937	-0.063	1
1.818	1.000	0.937	-0.063	1
2.039	0.350	0.597	0.247	1
2.039	0.500	0.597	0.097	1
2.039	0.550	0.597	0.047	1
2.039	0.650	0.597	-0.053	1
2.039	0.700	0.597	-0.103	1
2.299	0.050	0.198	0.148	1
2.299	0.150	0.198	0.048	1
2.299	0.200	0.198	-0.002	1.
2.299	0.250	0.198	-0.052	1.
2.299	0.450	0.198	-0.252	1.
2.460	0.000	0.000	0.000	1.

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MED Toxic Response Analysis Model, Version 1.00

MED Toxic Response Analysis Model 03/29/2018 16:36

		Data Summary		
Exposure	Obs Effects	Pred Effects	Residual	Weigh
2.460	0.000	0.000	0.000	1
2.460	0.000	0.000	0.000	1
2.460	0.000	0.000	0.000	1
2.460	0.000	0.000	0.000	1

Error Summary
 No Errors

Puge 19 of 19 N312/3018