

825 N. Rutledge Springfield, Illinois 62702 217.782.9780

### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### **Volatile Organic Compounds by GC/MS**

 Method:
 524.3
 Prepared:
 07/23/21 08:00

 Units:
 ug/L
 Analyzed:
 07/23/21 17:34

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,1,1-Trichloroethane	< 0.50		0.50
1,1,2-Trichloroethane	< 0.50		0.50
1,1-Dichloroethene	< 0.50		0.50
1,2,4-Trichlorobenzene	< 0.50		0.50
1,2-Dichlorobenzene	< 0.50		0.50
1,2-Dichloroethane	< 0.50		0.50
1,2-Dichloropropane	< 0.50		0.50
1,4-Dichlorobenzene	< 0.50		0.50
Benzene	< 0.50		0.50
Carbon tetrachloride	< 0.50		0.50
Chlorobenzene	< 0.50		0.50
cis-1,2-Dichloroethene	< 0.50		0.50
Ethylbenzene	< 0.50		0.50
Methyl tert-butyl ether	< 0.50		0.50
Methylene chloride	< 0.50		0.50
Styrene	< 0.50		0.50
Tetrachloroethene	< 0.50		0.50
Toluene	< 0.50		0.50
trans-1,2-Dichloroethene	< 0.50		0.50
Trichloroethene	< 0.50		0.50
Vinyl chloride	< 0.50		0.50
Xylenes, total	< 0.50		0.50



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 11:41

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,1,1,2-Tetrachloroethane	< 2.0		2.0
1,1,1-Trichloroethane	< 2.0		2.0
1,1,2,2-Tetrachloroethane	< 2.0		2.0
1,1,2-Trichloroethane	< 2.0		2.0
1,1-Dichloroethane	< 2.0		2.0
1,1-Dichloroethene	< 2.0		2.0
1,1-Dichloropropene	< 2.0		2.0
1,2,3-Trichloropropane	< 2.0		2.0
1,2-Dibromoethane	< 2.0		2.0
1,2-Dichloroethane	< 2.0		2.0
1,2-Dichloropropane	< 2.0		2.0
1,3-Dichloropropane	< 2.0		2.0
2,2-Dichloropropane	< 2.0		2.0
2-Butanone (MEK)	< 10		10
2-Hexanone (MBK)	< 5.0		5.0
4-Methyl-2-pentanone (MIBK)	< 10		10
Acetone	< 10		10
Benzene	< 2.0		2.0
Bromobenzene	< 2.0		2.0
Bromochloromethane	< 2.0		2.0
Bromodichloromethane	< 2.0		2.0
Bromoform	< 5.0		5.0
Bromomethane	< 5.0	01	5.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 11:41

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Carbon disulfide	< 2.0		2.0
Carbon tetrachloride	< 2.0		2.0
Chlorobenzene	< 2.0		2.0
Chloroethane	< 2.0		2.0
Chloroform	< 2.0		2.0
Chloromethane	< 2.0		2.0
cis-1,2-Dichloroethene	< 2.0		2.0
cis-1,3-Dichloropropene	< 2.0		2.0
Dibromochloromethane	< 5.0		5.0
Dibromomethane	< 2.0		2.0
Ethylbenzene	< 2.0		2.0
Isopropylbenzene	< 2.0		2.0
Methyl tert-butyl ether	< 2.0		2.0
Methylene chloride	< 5.0		5.0
Styrene	< 2.0		2.0
Tetrachloroethene	< 2.0		2.0
Toluene	< 2.0		2.0
trans-1,2-Dichloroethene	< 2.0		2.0
trans-1,3-Dichloropropene	< 5.0		5.0
Trichloroethene	< 2.0		2.0
Trichlorofluoromethane	< 2.0		2.0
Vinyl chloride	< 2.0		2.0
Xylenes, total	< 2.0		2.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/23/21 10:52

 Units:
 ug/L
 Analyzed:
 07/26/21 17:56

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,2,4,5-Tetrachlorobenzene	< 1.5		1.5
1,2,4-Trichlorobenzene	< 1.5		1.5
1,2-Dichlorobenzene	< 1.5		1.5
1,2-Dinitrobenzene	< 1.5		1.5
1,3-Dichlorobenzene	< 1.5		1.5
1,3-Dinitrobenzene	< 5.0		5.0
1,4-Dichlorobenzene	< 1.5		1.5
1,4-Dinitrobenzene	< 5.0		5.0
1-Chloronaphthalene	< 1.5		1.5
1-Naphthylamine	< 5.0		5.0
2,2-Oxybis(1-chloropropane)	< 1.5		1.5
2,3,4,6-Tetrachlorophenol	< 1.5		1.5
2,4,5-Trichlorophenol	< 1.5		1.5
2,4,6-Trichlorophenol	< 1.5		1.5
2,4-Dichlorophenol	< 1.5		1.5
2,4-Dimethylphenol	< 1.5		1.5
2,4-Dinitrophenol	< 7.5		7.5
2,4-Dinitrotoluene	< 5.0		5.0
2,6-Dichlorophenol	< 1.5		1.5
2,6-Dinitrotoluene	< 1.5		1.5
2-Chloronaphthalene	< 1.5		1.5
2-Chlorophenol	< 1.5		1.5
2-Methylnaphthalene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/23/21 10:52

 Units:
 ug/L
 Analyzed:
 07/26/21 17:56

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
2-Methylphenol	< 1.5		1.5
2-Naphthylamine	< 5.0		5.0
2-Nitroaniline	< 1.5		1.5
2-Nitrophenol	< 5.0		5.0
2-Picoline	< 1.5		1.5
3,3-Dichlorobenzidine	< 1.5		1.5
3-Nitroaniline	< 1.5		1.5
4,6-Dinitro-2-methylphenol	< 5.0		5.0
4-Bromophenyl phenyl ether	< 1.5		1.5
4-Chloro-3-methylphenol	< 1.5		1.5
4-Chloroaniline	< 1.5		1.5
4-Chlorophenyl phenyl ether	< 1.5		1.5
4-Methylphenol	< 1.5		1.5
4-Nitroaniline	< 1.5		1.5
4-Nitrobiphenyl	< 5.0		5.0
4-Nitrophenol	< 5.0		5.0
5-Nitroacenaphthene	< 5.0		5.0
7,12-Dimethylbenzo(a)anthracene	< 5.0		5.0
Acenaphthene	< 1.5		1.5
Acenaphthylene	< 1.5		1.5
Acetophenone	< 1.5		1.5
Anthracene	< 1.5		1.5
Azobenzene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **G210** Lab Sample ID: **21G0930-01** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/23/21 10:52

 Units:
 ug/L
 Analyzed:
 07/26/21 17:56

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Benzo(a)anthracene	< 1.5		1.5
Benzo(a)pyrene	< 1.5		1.5
Benzo(b)fluoranthene	< 1.5		1.5
Benzo(ghi)perylene	< 5.0		5.0
Benzo(k)fluoranthene	< 1.5		1.5
Bis(2-chloroethoxy)methane	< 1.5		1.5
Bis(2-chloroethyl)ether	< 1.5		1.5
Bis(2-ethylhexyl)phthalate	< 5.0		5.0
Butyl benzyl phthalate	< 5.0		5.0
Carbazole	< 1.5		1.5
Chrysene	< 1.5		1.5
Dibenzo(a,h)anthracene	< 5.0		5.0
Dibenzofuran	< 1.5		1.5
Diethylphthalate	< 1.5		1.5
Dimethylphthalate	< 1.5		1.5
Di-n-butylphthalate	< 1.5		1.5
Di-n-octylphthalate	< 5.0		5.0
Diphenylamine	< 1.5		1.5
Ethyl methanesulfonate	< 1.5		1.5
Fluoranthene	< 1.5		1.5
Fluorene	< 1.5		1.5
Hexachlorobenzene	< 1.5		1.5
Hexachlorobutadiene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **G210** Lab Sample ID: **21G0930-01** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/23/21 10:52

 Units:
 ug/L
 Analyzed:
 07/26/21 17:56

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Hexachlorocyclopentadiene	< 1.5		1.5
Hexachloroethane	< 1.5		1.5
Hexachloropropene	< 1.5		1.5
Indeno(1,2,3-cd)pyrene	< 5.0		5.0
Isodrin	< 1.5		1.5
Isophorone	< 1.5		1.5
Isosafrole	< 1.5		1.5
Mestranol	< 5.0		5.0
Methyl methanesulfonate	< 1.5		1.5
Naphthalene	< 1.5		1.5
Nitrobenzene	< 1.5		1.5
N-Nitrosodi-n-butylamine	< 1.5		1.5
N-Nitrosodi-n-propylamine	< 1.5		1.5
N-Nitrosopiperidine	< 1.5		1.5
p-Dimethylaminoazobenzene	< 1.5		1.5
Pentachlorobenzene	< 1.5		1.5
Pentachloronitrobenzene	< 1.5		1.5
Pentachlorophenol	< 5.0		5.0
Phenacetin	< 1.5		1.5
Phenanthrene	< 1.5		1.5
Phenol	< 1.5		1.5
Pronamide	< 1.5		1.5
Pyrene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name:	CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **Client Sample ID: 21G0930-01** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

#### Semivolatiles by GC/MS

Method: 8270 Prepared: 07/23/21 10:52

Units: ug/L Analyzed: 07/26/21 17:56

<u>Analyte</u> <u>Result</u> <u>Qualifier</u> <u>Reporting Limit</u>

Pyridine < 1.5 1.5
Safrole < 1.5 1.5

#### **Hexavalent Chromium**

Method: 218.6 Prepared: 07/23/21 08:45

Units: ug/L Analyzed: 07/23/21 08:45

Analyte Result Qualifier Reporting Limit

Hexavalent Chromium <50.0 50.0

#### Mercury by EPA Method 245.1

Method: 245.1 Prepared: 07/26/21 15:25

Units: ug/L Analyzed: 07/27/21 10:32

Analyte Result Qualifier Reporting Limit

Mercury < 0.06 0.06



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Metals (Digested Drinking Water) by EPA 200 Series Methods ICP

 Method:
 200.7

 Units:
 ug/L

 Analyzed:
 07/26/21 11:38

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Boron	33.7		25.0
Calcium	28000		300
Hardness	126000		1980
Iron	< 200		200
Magnesium	13700		300
Potassium	< 1400		1400
Silica	18500		2500
Sodium	151000		1000
Strontium	18.3		10.0

### Metals by EPA 200 Series Methods ICP/MS

 Method:
 200.8
 Prepared:
 07/28/21 11:20

 Units:
 ug/L
 Analyzed:
 07/28/21 16:32

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Aluminum	< 100		100
Antimony	< 2.00		2.00
Arsenic	< 1.00		1.00
Barium	< 5.00		5.00
Beryllium	< 1.00		1.00
Cadmium	< 3.00		3.00
Chromium	< 5.00		5.00



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-01

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:50

### Metals by EPA 200 Series Methods ICP/MS

 Method:
 200.8
 Prepared:
 07/28/21 11:20

 Units:
 ug/L
 Analyzed:
 07/28/21 16:32

<u>Analyte</u>	<u>Result</u>	<b>Qualifier</b>	Reporting Limit
Cobalt	< 10.0		10.0
Copper	< 100		100
Lead	< 5.00		5.00
Manganese	< 15.0		15.0
Molybdenum	< 20.0		20.0
Nickel	< 25.0		25.0
Selenium	< 2.00		2.00
Silver	< 10.0		10.0
Thallium	< 2.00		2.00
Vanadium	< 5.00		5.00
Zinc	< 100		100



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### **Volatile Organic Compounds by GC/MS**

 Method:
 524.3
 Prepared:
 07/23/21 08:00

 Units:
 ug/L
 Analyzed:
 07/23/21 17:58

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,1,1-Trichloroethane	< 0.50		0.50
1,1,2-Trichloroethane	< 0.50		0.50
1,1-Dichloroethene	< 0.50		0.50
1,2,4-Trichlorobenzene	< 0.50		0.50
1,2-Dichlorobenzene	< 0.50		0.50
1,2-Dichloroethane	< 0.50		0.50
1,2-Dichloropropane	< 0.50		0.50
1,4-Dichlorobenzene	< 0.50		0.50
Benzene	< 0.50		0.50
Carbon tetrachloride	< 0.50		0.50
Chlorobenzene	< 0.50		0.50
cis-1,2-Dichloroethene	< 0.50		0.50
Ethylbenzene	< 0.50		0.50
Methyl tert-butyl ether	< 0.50		0.50
Methylene chloride	< 0.50		0.50
Styrene	< 0.50		0.50
Tetrachloroethene	< 0.50		0.50
Toluene	< 0.50		0.50
trans-1,2-Dichloroethene	< 0.50		0.50
Trichloroethene	< 0.50		0.50
Vinyl chloride	< 0.50		0.50
Xylenes, total	< 0.50		0.50



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 12:04

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,1,1,2-Tetrachloroethane	< 2.0		2.0
1,1,1-Trichloroethane	< 2.0		2.0
1,1,2,2-Tetrachloroethane	< 2.0		2.0
1,1,2-Trichloroethane	< 2.0		2.0
1,1-Dichloroethane	< 2.0		2.0
1,1-Dichloroethene	< 2.0		2.0
1,1-Dichloropropene	< 2.0		2.0
1,2,3-Trichloropropane	< 2.0		2.0
1,2-Dibromoethane	< 2.0		2.0
1,2-Dichloroethane	< 2.0		2.0
1,2-Dichloropropane	< 2.0		2.0
1,3-Dichloropropane	< 2.0		2.0
2,2-Dichloropropane	< 2.0		2.0
2-Butanone (MEK)	< 10		10
2-Hexanone (MBK)	< 5.0		5.0
4-Methyl-2-pentanone (MIBK)	< 10		10
Acetone	< 10		10
Benzene	< 2.0		2.0
Bromobenzene	< 2.0		2.0
Bromochloromethane	< 2.0		2.0
Bromodichloromethane	< 2.0		2.0
Bromoform	< 5.0		5.0
Bromomethane	< 5.0	O1	5.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 12:04

<u>Analyte</u>	Result	<u>Qualifier</u>	Reporting Limit
Carbon disulfide	< 2.0		2.0
Carbon tetrachloride	< 2.0		2.0
Chlorobenzene	< 2.0		2.0
Chloroethane	< 2.0		2.0
Chloroform	< 2.0		2.0
Chloromethane	< 2.0		2.0
cis-1,2-Dichloroethene	< 2.0		2.0
cis-1,3-Dichloropropene	< 2.0		2.0
Dibromochloromethane	< 5.0		5.0
Dibromomethane	< 2.0		2.0
Ethylbenzene	< 2.0		2.0
Isopropylbenzene	< 2.0		2.0
Methyl tert-butyl ether	< 2.0		2.0
Methylene chloride	< 5.0		5.0
Styrene	< 2.0		2.0
Tetrachloroethene	< 2.0		2.0
Toluene	< 2.0		2.0
trans-1,2-Dichloroethene	< 2.0		2.0
trans-1,3-Dichloropropene	< 5.0		5.0
Trichloroethene	< 2.0		2.0
Trichlorofluoromethane	< 2.0		2.0
Vinyl chloride	< 2.0		2.0
Xylenes, total	< 2.0		2.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **G209** Lab Sample ID: **21G0930-02** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 13:40

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,2,4,5-Tetrachlorobenzene	< 1.6		1.6
1,2,4-Trichlorobenzene	< 1.6		1.6
1,2-Dichlorobenzene	< 1.6		1.6
1,2-Dinitrobenzene	< 1.6		1.6
1,3-Dichlorobenzene	< 1.6		1.6
1,3-Dinitrobenzene	< 5.3		5.3
1,4-Dichlorobenzene	< 1.6		1.6
1,4-Dinitrobenzene	< 5.3		5.3
1-Chloronaphthalene	< 1.6		1.6
1-Naphthylamine	< 5.3		5.3
2,2-Oxybis(1-chloropropane)	< 1.6		1.6
2,3,4,6-Tetrachlorophenol	< 1.6		1.6
2,4,5-Trichlorophenol	< 1.6		1.6
2,4,6-Trichlorophenol	< 1.6		1.6
2,4-Dichlorophenol	< 1.6		1.6
2,4-Dimethylphenol	< 1.6		1.6
2,4-Dinitrophenol	< 5.3		5.3
2,4-Dinitrotoluene	< 5.3		5.3
2,6-Dichlorophenol	< 1.6		1.6
2,6-Dinitrotoluene	< 1.6		1.6
2-Chloronaphthalene	< 1.6		1.6
2-Chlorophenol	< 1.6		1.6
2-Methylnaphthalene	< 1.6		1.6



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 13:40

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
2-Methylphenol	< 1.6		1.6
2-Naphthylamine	< 5.3		5.3
2-Nitroaniline	< 1.6		1.6
2-Nitrophenol	< 5.3		5.3
2-Picoline	< 1.6		1.6
3,3-Dichlorobenzidine	< 1.6		1.6
3-Nitroaniline	< 1.6		1.6
4,6-Dinitro-2-methylphenol	< 5.3		5.3
4-Bromophenyl phenyl ether	< 1.6		1.6
4-Chloro-3-methylphenol	< 1.6		1.6
4-Chloroaniline	< 1.6		1.6
4-Chlorophenyl phenyl ether	< 1.6		1.6
4-Methylphenol	< 1.6		1.6
4-Nitroaniline	< 1.6		1.6
4-Nitrobiphenyl	< 5.3		5.3
4-Nitrophenol	< 5.3		5.3
5-Nitroacenaphthene	< 5.3		5.3
7,12-Dimethylbenzo(a)anthracene	< 5.3		5.3
Acenaphthene	< 1.6		1.6
Acenaphthylene	< 1.6		1.6
Acetophenone	< 1.6		1.6
Anthracene	< 1.6		1.6
Azobenzene	< 1.6		1.6



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 13:40

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Benzo(a)anthracene	< 1.6		1.6
Benzo(a)pyrene	< 1.6		1.6
Benzo(b)fluoranthene	< 1.6		1.6
Benzo(ghi)perylene	< 5.3		5.3
Benzo(k)fluoranthene	< 1.6		1.6
Bis(2-chloroethoxy)methane	< 1.6		1.6
Bis(2-chloroethyl)ether	< 1.6		1.6
Bis(2-ethylhexyl)phthalate	< 5.3		5.3
Butyl benzyl phthalate	< 5.3		5.3
Carbazole	< 1.6		1.6
Chrysene	< 1.6		1.6
Dibenzo(a,h)anthracene	< 5.3		5.3
Dibenzofuran	< 1.6		1.6
Diethylphthalate	< 1.6		1.6
Dimethylphthalate	< 1.6		1.6
Di-n-butylphthalate	< 1.6		1.6
Di-n-octylphthalate	< 5.3		5.3
Diphenylamine	< 1.6		1.6
Ethyl methanesulfonate	< 1.6		1.6
Fluoranthene	< 1.6		1.6
Fluorene	< 1.6		1.6
Hexachlorobenzene	< 1.6		1.6
Hexachlorobutadiene	< 1.6		1.6



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#### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

#### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 13:40

Qualifier **Reporting Limit Analyte** Result < 1.6 1.6 Hexachlorocyclopentadiene 1.6 < 1.6 Hexachloroethane < 1.6 1.6 Hexachloropropene 5.3 < 5.3 Indeno(1,2,3-cd)pyrene < 1.6 1.6 Isodrin < 1.6 1.6 Isophorone < 1.6 1.6 Isosafrole 5.3 Mestranol < 5.3 < 1.6 1.6 Methyl methanesulfonate < 1.6 1.6 Naphthalene < 1.6 1.6 Nitrobenzene < 1.6 1.6 N-Nitrosodi-n-butylamine < 1.6 1.6 N-Nitrosodi-n-propylamine < 1.6 1.6 N-Nitrosopiperidine < 1.6 1.6 p-Dimethylaminoazobenzene < 1.6 1.6 Pentachlorobenzene < 1.6 1.6 Pentachloronitrobenzene 5.3 Pentachlorophenol < 5.3 < 1.6 1.6 Phenacetin < 1.6 Phenanthrene 1.6 Phenol < 1.6 1.6 < 1.6 1.6 Pronamide < 1.6 1.6 Pyrene



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### **LABORATORY RESULTS**

CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **G209** Lab Sample ID: **21G0930-02** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

Semivolatiles by GC/MS

Method: 8270 Prepared: 07/26/21 10:55

Units: ug/L Analyzed: 07/27/21 13:40

Analyte Result Qualifier Reporting Limit

Pyridine < 1.6 1.6 Safrole < 1.6 1.6

**Hexavalent Chromium** 

Method: 218.6 Prepared: 07/23/21 08:45

Units: ug/L Analyzed: 07/23/21 08:45

Analyte Result Qualifier Reporting Limit

Hexavalent Chromium < 50.0 50.0

**Mercury by EPA Method 245.1** 

Method: 245.1 Prepared: 07/26/21 15:25

Units: ug/L Analyzed: 07/27/21 10:34

Analyte Result Qualifier Reporting Limit

Mercury < 0.06 0.06



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-02

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Metals (Digested Drinking Water) by EPA 200 Series Methods ICP

 Method:
 200.7

 Units:
 ug/L

 Analyzed:
 07/26/21 11:42

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Boron	27.1		25.0
Calcium	78300		300
Hardness	352000		1980
Iron	< 200		200
Magnesium	37900		300
Potassium	1450		1400
Silica	18400		2500
Sodium	29800		1000
Strontium	52.7		10.0

#### Metals by EPA 200 Series Methods ICP/MS

 Method:
 200.8
 Prepared:
 07/28/21 11:20

 Units:
 ug/L
 Analyzed:
 07/28/21 16:36

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Aluminum	< 100		100
Antimony	< 2.00		2.00
Arsenic	< 1.00		1.00
Barium	8.04		5.00
Beryllium	< 1.00		1.00
Cadmium	< 3.00		3.00
Chromium	< 5.00		5.00



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **G209** Lab Sample ID: **21G0930-02** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 12:55

### Metals by EPA 200 Series Methods ICP/MS

 Method:
 200.8
 Prepared:
 07/28/21 11:20

 Units:
 ug/L
 Analyzed:
 07/28/21 16:36

<u>Analyte</u>	<u>Result</u>	<b>Qualifier</b>	Reporting Limit
Cobalt	< 10.0		10.0
Copper	< 100		100
Lead	< 5.00		5.00
Manganese	< 15.0		15.0
Molybdenum	< 20.0		20.0
Nickel	< 25.0		25.0
Selenium	< 2.00		2.00
Silver	< 10.0		10.0
Thallium	< 2.00		2.00
Vanadium	< 5.00		5.00
Zinc	< 100		100



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-03

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### **Volatile Organic Compounds by GC/MS**

 Method:
 524.3
 Prepared:
 07/23/21 08:00

 Units:
 ug/L
 Analyzed:
 07/23/21 18:21

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,1,1-Trichloroethane	< 0.50		0.50
1,1,2-Trichloroethane	< 0.50		0.50
1,1-Dichloroethene	< 0.50		0.50
1,2,4-Trichlorobenzene	< 0.50	J2	0.50
1,2-Dichlorobenzene	< 0.50	J2	0.50
1,2-Dichloroethane	< 0.50		0.50
1,2-Dichloropropane	< 0.50		0.50
1,4-Dichlorobenzene	< 0.50	J2	0.50
Benzene	< 0.50		0.50
Carbon tetrachloride	< 0.50		0.50
Chlorobenzene	< 0.50		0.50
cis-1,2-Dichloroethene	< 0.50		0.50
Ethylbenzene	< 0.50		0.50
Methyl tert-butyl ether	< 0.50		0.50
Methylene chloride	< 0.50		0.50
Styrene	< 0.50		0.50
Tetrachloroethene	< 0.50		0.50
Toluene	< 0.50		0.50
trans-1,2-Dichloroethene	< 0.50		0.50
Trichloroethene	< 0.50		0.50
Vinyl chloride	< 0.50		0.50
Xylenes, total	< 0.50		0.50



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **FB1** Lab Sample ID: **21G0930-03** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 12:26

<u>Analyte</u>	Result	Qualifier	Reporting Limit
1,1,1,2-Tetrachloroethane	< 2.0		2.0
1,1,1-Trichloroethane	< 2.0		2.0
1,1,2,2-Tetrachloroethane	< 2.0		2.0
1,1,2-Trichloroethane	< 2.0		2.0
1,1-Dichloroethane	< 2.0		2.0
1,1-Dichloroethene	< 2.0		2.0
1,1-Dichloropropene	< 2.0		2.0
1,2,3-Trichloropropane	< 2.0		2.0
1,2-Dibromoethane	< 2.0		2.0
1,2-Dichloroethane	< 2.0		2.0
1,2-Dichloropropane	< 2.0		2.0
1,3-Dichloropropane	< 2.0		2.0
2,2-Dichloropropane	< 2.0		2.0
2-Butanone (MEK)	< 10		10
2-Hexanone (MBK)	< 5.0		5.0
4-Methyl-2-pentanone (MIBK)	< 10		10
Acetone	< 10		10
Benzene	< 2.0		2.0
Bromobenzene	< 2.0		2.0
Bromochloromethane	< 2.0		2.0
Bromodichloromethane	< 2.0		2.0
Bromoform	< 5.0		5.0
Bromomethane	< 5.0	01	5.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **FB1** Lab Sample ID: **21G0930-03** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 12:26

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Carbon disulfide	< 2.0		2.0
Carbon tetrachloride	< 2.0		2.0
Chlorobenzene	< 2.0		2.0
Chloroethane	< 2.0		2.0
Chloroform	< 2.0		2.0
Chloromethane	< 2.0		2.0
cis-1,2-Dichloroethene	< 2.0		2.0
cis-1,3-Dichloropropene	< 2.0		2.0
Dibromochloromethane	< 5.0		5.0
Dibromomethane	< 2.0		2.0
Ethylbenzene	< 2.0		2.0
Isopropylbenzene	< 2.0		2.0
Methyl tert-butyl ether	< 2.0		2.0
Methylene chloride	< 5.0		5.0
Styrene	< 2.0		2.0
Tetrachloroethene	< 2.0		2.0
Toluene	< 2.0		2.0
trans-1,2-Dichloroethene	< 2.0		2.0
trans-1,3-Dichloropropene	< 5.0		5.0
Trichloroethene	< 2.0		2.0
Trichlorofluoromethane	< 2.0		2.0
Vinyl chloride	< 2.0		2.0
Xylenes, total	< 2.0		2.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-03

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 14:14

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,2,4,5-Tetrachlorobenzene	< 1.5		1.5
1,2,4-Trichlorobenzene	< 1.5		1.5
1,2-Dichlorobenzene	< 1.5		1.5
1,2-Dinitrobenzene	< 1.5		1.5
1,3-Dichlorobenzene	< 1.5		1.5
1,3-Dinitrobenzene	< 5.0		5.0
1,4-Dichlorobenzene	< 1.5		1.5
1,4-Dinitrobenzene	< 5.0		5.0
1-Chloronaphthalene	< 1.5		1.5
1-Naphthylamine	< 5.0		5.0
2,2-Oxybis(1-chloropropane)	< 1.5		1.5
2,3,4,6-Tetrachlorophenol	< 1.5		1.5
2,4,5-Trichlorophenol	< 1.5		1.5
2,4,6-Trichlorophenol	< 1.5		1.5
2,4-Dichlorophenol	< 1.5		1.5
2,4-Dimethylphenol	< 1.5		1.5
2,4-Dinitrophenol	< 5.0		5.0
2,4-Dinitrotoluene	< 5.0		5.0
2,6-Dichlorophenol	< 1.5		1.5
2,6-Dinitrotoluene	< 1.5		1.5
2-Chloronaphthalene	< 1.5		1.5
2-Chlorophenol	< 1.5		1.5
2-Methylnaphthalene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-03

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 14:14

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
2-Methylphenol	< 1.5		1.5
2-Naphthylamine	< 5.0		5.0
2-Nitroaniline	< 1.5		1.5
2-Nitrophenol	< 5.0		5.0
2-Picoline	< 1.5		1.5
3,3-Dichlorobenzidine	< 1.5		1.5
3-Nitroaniline	< 1.5		1.5
4,6-Dinitro-2-methylphenol	< 5.0		5.0
4-Bromophenyl phenyl ether	< 1.5		1.5
4-Chloro-3-methylphenol	< 1.5		1.5
4-Chloroaniline	< 1.5		1.5
4-Chlorophenyl phenyl ether	< 1.5		1.5
4-Methylphenol	< 1.5		1.5
4-Nitroaniline	< 1.5		1.5
4-Nitrobiphenyl	< 5.0		5.0
4-Nitrophenol	< 5.0		5.0
5-Nitroacenaphthene	< 5.0		5.0
7,12-Dimethylbenzo(a)anthracene	< 5.0		5.0
Acenaphthene	< 1.5		1.5
Acenaphthylene	< 1.5		1.5
Acetophenone	< 1.5		1.5
Anthracene	< 1.5		1.5
Azobenzene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-03

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 14:14

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Benzo(a)anthracene	< 1.5		1.5
Benzo(a)pyrene	< 1.5		1.5
Benzo(b)fluoranthene	< 1.5		1.5
Benzo(ghi)perylene	< 5.0		5.0
Benzo(k)fluoranthene	< 1.5		1.5
Bis(2-chloroethoxy)methane	< 1.5		1.5
Bis(2-chloroethyl)ether	< 1.5		1.5
Bis(2-ethylhexyl)phthalate	< 5.0		5.0
Butyl benzyl phthalate	< 5.0		5.0
Carbazole	< 1.5		1.5
Chrysene	< 1.5		1.5
Dibenzo(a,h)anthracene	< 5.0		5.0
Dibenzofuran	< 1.5		1.5
Diethylphthalate	< 1.5		1.5
Dimethylphthalate	< 1.5		1.5
Di-n-butylphthalate	< 1.5		1.5
Di-n-octylphthalate	< 5.0		5.0
Diphenylamine	< 1.5		1.5
Ethyl methanesulfonate	< 1.5		1.5
Fluoranthene	< 1.5		1.5
Fluorene	< 1.5		1.5
Hexachlorobenzene	< 1.5		1.5
Hexachlorobutadiene	< 1.5		1.5



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **FB1** Lab Sample ID: **21G0930-03** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Semivolatiles by GC/MS

 Method:
 8270
 Prepared:
 07/26/21 10:55

 Units:
 ug/L
 Analyzed:
 07/27/21 14:14

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Hexachlorocyclopentadiene	< 1.5		1.5
Hexachloroethane	< 1.5		1.5
Hexachloropropene	< 1.5		1.5
Indeno(1,2,3-cd)pyrene	< 5.0		5.0
Isodrin	< 1.5		1.5
Isophorone	< 1.5		1.5
Isosafrole	< 1.5		1.5
Mestranol	< 5.0		5.0
Methyl methanesulfonate	< 1.5		1.5
Naphthalene	< 1.5		1.5
Nitrobenzene	< 1.5		1.5
N-Nitrosodi-n-butylamine	< 1.5		1.5
N-Nitrosodi-n-propylamine	< 1.5		1.5
N-Nitrosopiperidine	< 1.5		1.5
p-Dimethylaminoazobenzene	< 1.5		1.5
Pentachlorobenzene	< 1.5		1.5
Pentachloronitrobenzene	< 1.5		1.5
Pentachlorophenol	< 5.0		5.0
Phenacetin	< 1.5		1.5
Phenanthrene	< 1.5		1.5
Phenol	< 1.5		1.5
Pronamide	< 1.5		1.5
Pyrene	< 1.5		1.5



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### **LABORATORY RESULTS**

V:	ame:	CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-03

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

Semivolatiles by GC/MS

Method: 8270 Prepared: 07/26/21 10:55

Units: ug/L Analyzed: 07/27/21 14:14

<u>Analyte</u> <u>Result</u> <u>Qualifier</u> <u>Reporting Limit</u>

Pyridine < 1.5 1.5
Safrole < 1.5 1.5

**Hexavalent Chromium** 

Method: 218.6 Prepared: 07/23/21 08:45

Units: ug/L Analyzed: 07/23/21 08:45

Analyte Result Qualifier Reporting Limit

Hexavalent Chromium <50.0 50.0

Mercury by EPA Method 245.1

Method: 245.1 Prepared: 07/26/21 15:25

Units: ug/L Analyzed: 07/27/21 10:37

Analyte Result Qualifier Reporting Limit

Mercury < 0.06 0.06



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21
Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: FB1 Lab Sample ID: 21G0930-03

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Metals (Digested Drinking Water) by EPA 200 Series Methods ICP

 Method:
 200.7

 Units:
 ug/L

 Analyzed:
 07/26/21 11:45

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Boron	< 25.0		25.0
Calcium	< 300		300
Hardness	< 1980		1980
Iron	< 200		200
Magnesium	< 300		300
Potassium	< 1400		1400
Silica	< 2500		2500
Sodium	< 1000		1000
Strontium	< 10.0		10.0

### Metals by EPA 200 Series Methods ICP/MS

 Method:
 200.8
 Prepared:
 07/28/21 11:20

 Units:
 ug/L
 Analyzed:
 07/28/21 16:41

<u>Analyte</u>	Result	Qualifier_	Reporting Limit
Aluminum	< 100		100
Antimony	< 2.00		2.00
Arsenic	< 1.00		1.00
Barium	< 5.00		5.00
Beryllium	< 1.00		1.00
Cadmium	< 3.00		3.00
Chromium	< 5.00		5.00



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: **FB1** Lab Sample ID: **21G0930-03** 

Matrix: Water Collected By: JW Date/Time Collected: 07/22/21 13:00

### Metals by EPA 200 Series Methods ICP/MS

 Method:
 200.8
 Prepared:
 07/28/21 11:20

 Units:
 ug/L
 Analyzed:
 07/28/21 16:41

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
Cobalt	< 10.0		10.0
Copper	< 100		100
Lead	< 5.00		5.00
Manganese	< 15.0		15.0
Molybdenum	< 20.0		20.0
Nickel	< 25.0		25.0
Selenium	< 2.00		2.00
Silver	< 10.0		10.0
Thallium	< 2.00		2.00
Vanadium	< 5.00		5.00
Zinc	< 100		100



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-04

Matrix: Water Collected By: Date/Time Collected: 07/22/21 0:00

### Volatile Organic Compounds by GC/MS

 Method:
 524.3
 Prepared:
 07/23/21 08:00

 Units:
 ug/L
 Analyzed:
 07/23/21 20:42

<u>Analyte</u>	Result	<b>Qualifier</b>	Reporting Limit
1,1,1-Trichloroethane	< 0.50		0.50
1,1,2-Trichloroethane	< 0.50		0.50
1,1-Dichloroethene	< 0.50		0.50
1,2,4-Trichlorobenzene	< 0.50		0.50
1,2-Dichlorobenzene	< 0.50		0.50
1,2-Dichloroethane	< 0.50		0.50
1,2-Dichloropropane	< 0.50		0.50
1,4-Dichlorobenzene	< 0.50		0.50
Benzene	< 0.50		0.50
Carbon tetrachloride	< 0.50		0.50
Chlorobenzene	< 0.50		0.50
cis-1,2-Dichloroethene	< 0.50		0.50
Ethylbenzene	< 0.50		0.50
Methyl tert-butyl ether	< 0.50		0.50
Methylene chloride	< 0.50		0.50
Styrene	< 0.50		0.50
Tetrachloroethene	< 0.50		0.50
Toluene	< 0.50		0.50
trans-1,2-Dichloroethene	< 0.50		0.50
Trichloroethene	< 0.50		0.50
Vinyl chloride	< 0.50		0.50
Xylenes, total	< 0.50		0.50



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-04

Matrix: Water Collected By: Date/Time Collected: 07/22/21 0:00

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 16:00

<u>Analyte</u>	Result	Qualifier	Reporting Limit
1,1,1,2-Tetrachloroethane	< 2.0		2.0
1,1,1-Trichloroethane	< 2.0		2.0
1,1,2,2-Tetrachloroethane	< 2.0		2.0
1,1,2-Trichloroethane	< 2.0		2.0
1,1-Dichloroethane	< 2.0		2.0
1,1-Dichloroethene	< 2.0		2.0
1,1-Dichloropropene	< 2.0		2.0
1,2,3-Trichloropropane	< 2.0		2.0
1,2-Dibromoethane	< 2.0		2.0
1,2-Dichloroethane	< 2.0		2.0
1,2-Dichloropropane	< 2.0		2.0
1,3-Dichloropropane	< 2.0		2.0
2,2-Dichloropropane	< 2.0		2.0
2-Butanone (MEK)	< 10		10
2-Hexanone (MBK)	< 5.0		5.0
4-Methyl-2-pentanone (MIBK)	< 10		10
Acetone	< 10		10
Benzene	< 2.0		2.0
Bromobenzene	< 2.0		2.0
Bromochloromethane	< 2.0		2.0
Bromodichloromethane	< 2.0		2.0
Bromoform	< 5.0		5.0
Bromomethane	< 5.0	01	5.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

Client Sample ID: Lab Sample ID: 21G0930-04

Matrix: Water Collected By: Date/Time Collected: 07/22/21 0:00

### Volatiles Organic Compounds by Purge and Trap GC/MS

 Method:
 8260
 Prepared:
 07/26/21 08:00

 Units:
 ug/L
 Analyzed:
 07/27/21 16:00

<u>Analyte</u>	Result	Qualifier	Reporting Limit
Carbon disulfide	< 2.0		2.0
Carbon tetrachloride	< 2.0		2.0
Chlorobenzene	< 2.0		2.0
Chloroethane	< 2.0		2.0
Chloroform	< 2.0		2.0
Chloromethane	< 2.0		2.0
cis-1,2-Dichloroethene	< 2.0		2.0
cis-1,3-Dichloropropene	< 2.0		2.0
Dibromochloromethane	< 5.0		5.0
Dibromomethane	< 2.0		2.0
Ethylbenzene	< 2.0		2.0
Isopropylbenzene	< 2.0		2.0
Methyl tert-butyl ether	< 2.0		2.0
Methylene chloride	< 5.0		5.0
Styrene	< 2.0		2.0
Tetrachloroethene	< 2.0		2.0
Toluene	< 2.0		2.0
trans-1,2-Dichloroethene	< 2.0		2.0
trans-1,3-Dichloropropene	< 5.0		5.0
Trichloroethene	< 2.0		2.0
Trichlorofluoromethane	< 2.0		2.0
Vinyl chloride	< 2.0		2.0
Xylenes, total	< 2.0		2.0



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### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00



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#### **LABORATORY RESULTS**

Name: CHEMTOOL

Project/Facility Number: 2010355004 Date Received: 07/22/21

Funding Code: CS29 B50 Temperature C: 1.00

#### **Notes and Definitions**

O1 Quality control sample failed high - possible high bias or false positive result.

J2 Internal Standard criteria were not met.

ND Analyte NOT DETECTED at or above the reporting limit

Non-NELAP accredited

Method 8270: There was insufficient amount of sample to perform a matrix spike duplicate analysis. NELAC and method requirements were not met.

Method 8270: There was insufficient amount of sample to perform a matrix spike and matrix spike duplicate analysis for 21G0930-02 and -03 as these samples had to be extracted in a separate batch. NELAC and method requirements were not met.

Drinking Water Methods 200.7 and 200.8 were assigned to this work order for Metals analysis. No samples in this work order required a digestion to be performed based on turbidity.

Report Authorized by:

Tom Weiss Laboratory Manager The results in this report apply to the samples analyzed in accordance with the chain of custody document. This analytical report must be reproduced in its entirety. Test results meet all requirements of NELAC (accredited by Florida DOH #E37645). If you have any questions about this report, please contact Tom Weiss, Laboratory Manager, at 217.782.9780.

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