## **GROUNDWATER INFORMATION SHEET**

# **Hexavalent Chromium**

The purpose of this groundwater information sheet is to provide general information regarding a specific constituent of concern (COC). The information provided herein relates to groundwater sources used for public drinking water, not water served at the tap.

	GENERAL INFORMATION		
Constituent of Concern	Hexavalent Chromium		
Synonyms	Chromium VI, Chromium Six, Chrome 6, Cr <sup>6</sup>		
Chemical Formula	Cr <sup>6</sup>		
CAS No.	18540-29-9		
Storet No.	01032		
Summary	In 2014, the California State Water Resources Control Board (SWRCB) established a Maximum Contaminant Level (MCL) for hexavalent chromium at 10 micrograms per liter (µg/L) that was withdrawn by the SWRCB in August 2017. Until a revised MCL is adopted by the SWRCB, the total chromium MCL (50 µg/L) will be used as the drinking water standard. In order to show a spatial distribution of hexavalent chromium in public wells the Detection Limit for Purposes of Reporting (DLR) of 1 µg/L is used for this fact sheet. Based on SWRCB data from 2007 to 2017, 3,778 active and standby public water supply wells (of 8,765 wells sampled) had at least one detection of hexavalent chromium above the DLR. Most wells with detections have occurred in Los Angeles (514), San Bernardino (429) and Fresno (323) counties.		

REGULATORY AND WATER QUALITY LEVELS <sup>1</sup> HEXAVALENT CHROMIUM				
Туре	Agency	Concentration		
Federal MCL <sup>2</sup>	US EPA <sup>3</sup>	None established		
State MCL	SWRCB <sup>4</sup>	In progress		
Detection Limit for Purposes of Reporting (DLR) Former California State Notification Level (NL)	SWRCB	1 μg/L for Cr6		
Public Health Goal (PHG)	OEHHA <sup>5</sup>	0.02 μg/L		
Others: HBSL <sup>6</sup> (non-cancer health effect) Cal/EPA Cancer Potency Factor as a drinking water level	USGS <sup>7</sup> Cal/EPA	20 μg/L 0.07 μg/L		

<sup>1</sup> These levels generally relate to drinking water. Other water quality levels may exist. For further information see A Compilation of Water Quality Goals, 17<sup>th</sup> Edition (SWRCB, 2016).

<sup>2</sup> MCL - Maximum Contaminant Level

<sup>3</sup> US EPA - United States Environmental Protection Agency

<sup>4</sup> SWRCB - State Water Resources Control Board

<sup>5</sup>OEHHA - Office of Environmental Health and Human Hazard Assessment

<sup>6</sup> HBSL - Health Based Screening Level

<sup>7</sup>USGS - United State Geological Survey

HEXAVALENT CHROMIUM DETECTIONS IN PUBLIC WATER WELLS <sup>8</sup>				
Detection Type	Number of Groundwater Sources			
Number of active and standby public water wells	3,778 of 8,765 wells tested.			
with Cr <sup>6</sup> concentrations above 1 $\mu$ g/L <sup>9</sup> .				
Top 3 counties with active and standby public	Los Angeles (514), San Bernardino			
water wells with Cr <sup>6</sup> concentrations above 1	(429), Fresno (323)			
μg/L.				

<sup>8</sup>Based on 2007-2017 public standby and active well (groundwater sources) data collected by the SWRCB. <sup>9</sup>Data from private domestic wells and wells with less than 15 service connections are not available.

ANALYTICAL INFORMATION				
Method	Detection Limit	Note		
US EPA 218.7	0.01 µg/L	By ion chromatography with post-column derivatization and UV-visible spectroscopic detection (SWRCB approved for drinking water)		
US EPA 218.6	0.3 µg/L	By ion chromatography (SWRCB approved for drinking water)		
USGS by GFAAS	0.05 µg/L	Cr <sup>6</sup> separation in the field, not time sensitive		
Known Limitations to Analytical Methods	Water sample pH must be adjusted to 9.0-9.5, stored at 4°C and analyzed within 24 hours.			
Public Drinking Water Testing Requirements	In January 2001, hexavalent chromium was identified as an unregulated chemical requiring monitoring. As a result, public water systems began to test for hexavalent chromium in their drinking water supplies to the DLR of 1 $\mu$ g/L. In 2014, MCL for hexavalent chromium was established at 10 $\mu$ g/L. On August 1, 2017 the MCL was repealed. Compliance with the MCL for total chromium continues to be required.			

OCCURRENCE		
Anthropogenic Sources	Chromium is a metallic chemical that originates as a contaminant in the environment from the discharges of dye and paint pigments, wood preservatives, chrome-plating liquid wastes, and leaching from hazardous waste sites. The greatest use of chromium is in metal alloys such as stainless steel; protective coatings on metal; magnetic tapes; and pigments for paints, cement, paper, rubber, composition floor covering, etc. The two largest sources of chromium emission in the atmosphere are from the chemical manufacturing and combustion of natural gas, oil and coal.	

	GAMA Program
Natural Sources	Chromium is a metal found in natural deposits of ores containing other elements, mostly as chrome-iron ore. It is also widely present in soil and plants. Under most conditions, natural chromium in the environment occurs as Cr <sup>3</sup> . Under oxidizing conditions, alkaline pH range, presence of MnO <sub>2</sub> , and minerals containing chromium, part of it may occur as hexavalent chromium dissolved in groundwater. Recent sampling of drinking water sources throughout California suggests that hexavalent chromium may occur naturally in groundwater at many locations. Naturally occurring hexavalent chromium may be associated with serpentinite-containing rock or chromium containing geologic formations.
History of Occurrence	Hexavalent chromium has been detected in groundwater at several industrial sites where wood treatment or metal plating solutions were used. Between 1952 and 1966, hexavalent chromium was used to reduce corrosion in natural gas compressor plant in Hinkley, near Barstow. Hexavalent chromium contaminated groundwater was suspected of causing cancer and tumors in local residents beginning in the mid 1980's. Since then, elevated levels of hexavalent chromium has been detected in groundwater at several other locations including: Glendale, Topock, and Kettleman City. Hexavalent chromium also occurs naturally in groundwater at the Presidio of San Francisco and Lawrence Livermore National Laboratory.
Transport Characteristics	Hexavalent chromium is readily soluble in water. Under high Eh (oxidizing) and alkaline (pH above 7) conditions, hexavalent chromium can be predominant in groundwater. However, in the presence of organic matter, ferrous iron (Fe II) and sulfide, hexavalent chromium can be readily reduced to Cr <sup>3</sup> and immobilized. Adsorption of hexavalent chromium by clayey soil and natural aquifer materials is low to moderate under near-neutral pH ranges commonly encountered in groundwater.

## **REMEDIATION & TREATMENT TECHNOLOGIES**

#### In-situ Treatment:

In several laboratory and field pilot tests, and full-scale remediation systems, hexavalent chromium has been removed using a permeable reactive barrier filled with zero-valent iron granules or surfactant-modified zeolite. Engineered chemical reduction technologies involve the addition or in-situ injection of an electron donor such as hydrogen sulfite, sodium dithionite, sodium metabisulfite, calcium metabisulfite calcium polysulfite or tin(II) chloride. Other methods include geochemical fixation, soil flushing and extraction, bioremediation and electrokinetics.

#### **Above-Ground Treatment**

Drinking water can be treated by different pump and treat remediation systems. Cr<sup>3</sup> and Cr<sup>6</sup> can be removed by reverse osmosis or ion exchange resin. The ion exchange method should be used with caution, as presence of other metals may interact with the process and decrease system effectiveness. Removal of Cr<sup>6</sup> by seaweed biosorbent and bacteria (Bacillus sp.) within packed bed reactors has also been used.

#### Natural Attenuation

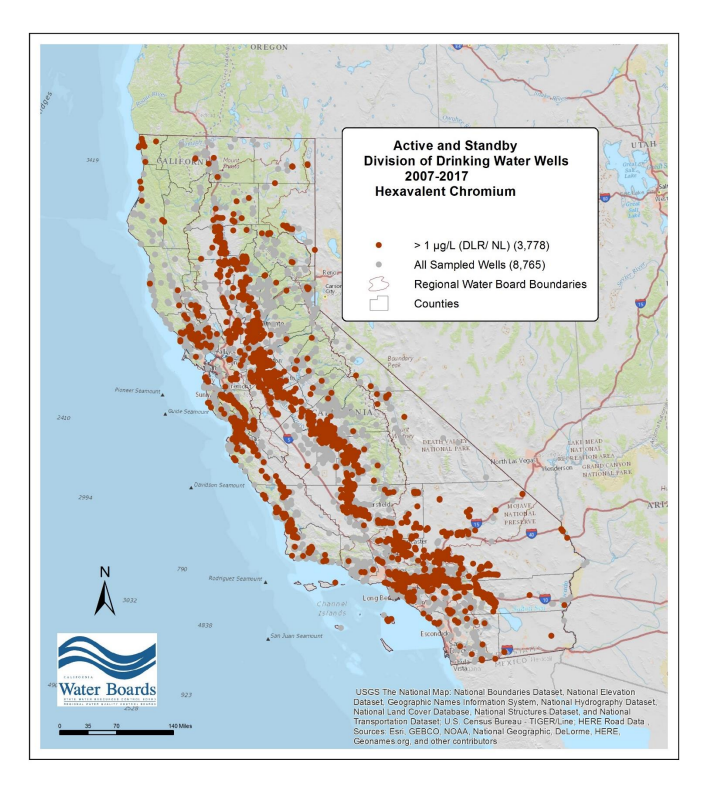
Natural attenuation of hexavalent chromium may occur in the subsurface environment through reduction by organic matter, iron hydroxides or sulfides. Prior to selection of natural attenuation as an option for remediation, the following conditions need to be demonstrated: 1) there are natural reducers present within the aquifer, 2) the amount of hexavalent chromium and other reactive constituents do not exceed the capacity of the aquifer to reduce them, 3) the rate of hexavalent chromium reduction is greater than the rate of transport of the aqueous hexavalent chromium off the impacted site, 4) the hexavalent chromium remains immobile, and 5) there is no net oxidation of  $Cr^3$  to  $Cr^6$ .

### HEALTH EFFECT INFORMATION

Hexavalent chromium is known to cause cancer in humans when inhaled. It can also damage the lining of the nose and throat and irritate the lungs. A number of scientific studies have found elevated rates of lung cancer in workers with occupational exposure to hexavalent chromium by inhalation. A few studies of workers exposed by inhalation have shown an increase in cancers of the gastrointestinal tract. When swallowed, hexavalent chromium can upset the gastrointestinal tract and damage the liver and kidneys. In recent scientific studies of laboratory animals, hexavalent chromium has been linked to cancer when ingested, although it is rapidly converted to Cr<sup>3</sup> after entering the stomach and coming into contact with organic matter.

### **KEY REFERENCES**

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Active and standby drinking water wells that had at least one detection of chromium-6 above the DLR<sup>1</sup>, 2007-2017, 3,778 wells. (Source: Public well data in GeoTracker GAMA).

<sup>1</sup>The DLR is equal to the former CA NL.

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