



# Vanadium

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# V

# 2 Vanadium

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#### Element Data

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- 7 Atomic Symbol: V
- 8 Atomic Number: 23
- 9 Atomic Weight: 50.9415 g/mol
- 10 Isotopes and Abundances: <sup>50</sup>V 0.25 %, <sup>51</sup>V 99.75 %
- 11 *I Atm Melting Point: 1910 °C*
- 12 1 Atm Boiling Point: 3407 °C
- 13 Common Valences: 5+, 4+, 3+, 2+
- 14 *Ionic Radii: 64 pm (in 3+, 6-fold coordination)*
- 15 Pauling Electronegativity: 1.63
- 16 First Ionization Energy: 6.74 eV
- 17 *Chondritic (CI) Abundance: 54.6 ppm<sup>a</sup>*
- 18 Silicate Earth Abundance: 86 ppm<sup>a</sup>
- 19 Crustal Abundance:  $138 \text{ ppm}^{\overline{b}}$
- 20 Seawater Abundance:  $\sim 30-37 \text{ nmol/kg}^{c}$
- 21 Core Abundance:  $\sim 150 \text{ ppm}^d$
- <sup>22</sup> <sup>*a*</sup>*Palme et al.* (2014)
- $^{b}$ Rudnick and Gao (2014)
- <sup>24</sup> <sup>c</sup>Bruland et al. (2014)
- $^{25}$  <sup>d</sup>McDonough (2014)

### 26 **Properties**

Vanadium (chemical symbol, V) is a d-block transition metal,
silver in color, appearing in the first long period of the periodic table between titanium and chromium. Vanadium has
two stable isotopes: <sup>50</sup>V and <sup>51</sup>V, with atomic abundance of

0.25 % and 99.75 %, respectively. Vanadium has several 31 oxidation forms (between 2+ and 5+). In the lithosphere, 32 V occurs as reducing V(III) form, whereas in oxidizing con- 33 ditions V prevails under V(IV) form. Vanadium(II) is partic- 34 ularly unstable in the environment. Vanadium(III) is more 35 stable than V(II), but it is also gradually oxidized by the air 36 or dissolved oxygen. Vanadium(V) is expected to be the 37 prevailing form in waters exposed to atmospheric oxygen, 38 whereas V(IV) may be present in reducing environments. 39 Depending upon geometry and environment, V ionic radii 40 vary between 36 pm and 79 pm. Vanadium has a high melting 41 point of 1910 °C and is a mildly incompatible, refractory, 42 lithophile (siderophile in the iron core and chondrites) ele- 43 ment. Vanadium has an electronegativity of 1.63 on the Pau- 44 ling scale and displays a first ionization potential of 45 6.74 eV. More details can be found in Richards (2006) and 46 Haynes (2015). 47

## **History and Use**

Vanadium was first discovered by del Rio in 1801 and was 49 incorrectly considered as impure chromium. Vanadium was 50 rediscovered and named by Sefström in 1830 after the Scan- 51 dinavian goddess Vanadis. In 1867, Roscoe isolated V in 52 nearly pure form by reducing V chloride with hydrogen. 53 Vanadium of high purity (99.3–99.8 %) was not produced 54 until 1927. The V redox chemistry and its specific application 55 to renewable energy technologies (e.g., vanadium redox bat- 56 tery; a type of rechargeable flow battery using V in different 57 oxidation states to store chemical potential energy) have 58 already add to the demand for this element and will definitely 59 continue to increase in the near future. Vanadium demand for 60 construction materials is the greatest consumer of mined V - 61 (Moskalyk and Alfantazi, 2003). Indeed, vanadium is the 62 most widely used alloying element for strengthening steels 63 employed in buildings and bridges. Vanadium volume that is 64 cycled through terrestrial and aquatic reservoirs can thus be 65

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expected to further growth. It results in uncontrolled
V releases in the environment, and environmental issues
appear in developing and developed countries (Imtiaz et al.,

69 **2015**).

#### 70 Geochemical Behavior

Widely and sparsely distributed, V is not found as the free
metal in nature. It originates from primary sources such as
ores (i.e., iron oxides deposits), metallurgical slags (i.e.,
processing of U and Ti ores), and petroleum residues. Other
sources include vanadiferous sandstones, bauxite, coals, or oil
shales.

Vanadium is widely distributed in igneous and sedimen-77 tary rocks and minerals. Among more than fifty minerals, 78 carnotite  $[K_2(UO_2)_2(VO_4)_2.3H_2O]$ , roscoelite  $[KV^{3+}_2(Si_3A])$ 79 O<sub>10</sub>(OH)<sub>2</sub>], vanadinite [Pb<sub>5</sub>(VO<sub>4</sub>)<sub>3</sub>Cl], mottramite [PbCu 80  $(VO_4)(OH)$ ], and patrónite  $(VS_4)$  are the most important 81 V-carriers. It is also present in some crude oils as organic 82 complexes. Its average content in the Earth's crust is approx-83 imately 138 ppm and is similar to that of Zn and Ni. It is, 84 however, more dispersed in the crust than either of those 85 elements, and concentrated mineral deposits are consequently 86 rare. The average V contents of CI chondrites range from 87 54.6 ppm to 56.5 ppm. The silicate Earth has a V content of 88 86 ppm, the bulk Earth of 105 ppm, whereas its content in the 89 metallic core is slightly higher (150 ppm). Vanadium contents 90 in the bulk continental crust vary significantly from 96 ppm to 91 230 ppm. These aspects have been further reviewed by 92 McDonough (2014), Palme et al. (2014), and Rudnick and 93 Gao (2014). 94

Historically, geochemists' interest in vanadium has 95 derived from the occurrence of variable redox states in Earth 96 surface environments, +3, +4, +5, and as a consequence its 97 application as a redox indicator (Huang et al., 2015). Indeed, 98 V chemical speciation and solubility are strong functions of 99 pH and Eh conditions. Incorporation of reduced V species 100 (III, IV) in clays and other secondary minerals provides a 101 useful redox indicator for past anoxic conditions. Moreover, 102 V behaves as an incompatible element because V(III) is 103 preferentially incorporated into crystals: V-bearing minerals 104 which crystallized in equilibrium with a low fO2 magma have 105 a higher V content than crystals from the same magma with 106 high  $fO_2$  (Mallmann and O'Neill, 2009). It results in an 107 increasing interest in V partitioning during high 108 temperature - high pressure magma differentiation processes. 109 Studies on V partitioning among Earth's crustal reservoirs 110 have provided critical insights into the interaction and evolu-111 tion of these reservoirs. Vanadium depletion in the silicate 112 Earth is explained by it preferential partitioning into the 113 metallic core during "deep magma ocean" process (at high 114 pressure and low  $fO_2$ ; Wood et al., 2006). The core could 115

contain half of the total V budget of the bulk Earth 116 (McDonough, 2014). Perspectives to exploit V stable isotope 117 chemistry are considered: Wu et al. (2015) proposed firstprinciples calculations that predict V isotopes fractionation 119 among V species with different valences in aqueous systems 120 and during sorption of V(V) to goethite. 121

Vanadium is also a naturally occurring element in air, soil, 122 plants, and water. The concentration of dissolved V in natural 123 freshwater and seawater is generally several mg/L or lower 124 (Bruland et al., 2014; Gaillardet et al., 2014). Vanadium 125 concentrations in marine and nonmarine sediments are gen- 126 erally below 100 ppm, reflecting a net accumulation from the 127 aqueous phase. Vanadium in trace amounts represents an 128 essential element for normal cell growth, but it may cause 129 adverse effects when its concentration is much greater than a 130 few tenths of  $\mu g/L$  (Chatteriee, 2009). Most data on the 131 release of V into the environment have been related to indus- 132 trial activities, especially from oil refineries and power plants 133 using V-rich fuel oil, and coal crude oil is enriched in V with 134 respect to many other trace elements, with concentrations 135 occasionally exceeding 1 mg/L. Thus, the fraction of 136 dissolved V in surface waters might be an environmental 137 indicator of oil combustion or pollution (Hope, 2008). Such 138 pollution sources may be responsible for appreciable amounts 139 of V into the environment, well above the natural background 140 levels associated with rock weathering and sediment 141 leaching. Fluvial dissolved V concentrations might also be 142 indicative of the types of rocks being weathered or of the 143 nature of the involved weathering processes. Weathering rate 144 and source rock type, rather than solution chemistry or anthro- 145 pogenic influences, appeared to be the important controlling 146 factors on fluvial dissolved V concentrations (Hope, 2008). 147

The oxidation rate of V(IV) to V(V) and the equilibrium 148 between these two species in aqueous solution will regulate 149 V prevalence in water (Pourret et al., 2012). It further depends 150 on several factors, such as pH, V concentration, redox poten- 151 tial, ionic strength of the aqueous system, and biological 152 activity. In water, V(IV) is commonly present as a vanadyl 153 cation  $[VO^{2^+}, VO(OH)^+]$ , whereas V(V) exists as a vanadate 154 oxyanion (H<sub>2</sub>VO<sub>4</sub><sup>-</sup>, HVO<sub>4</sub><sup>2-</sup>). VO<sup>2+</sup> is strongly adsorbed 155 onto solid phases, including organic and oxyhydroxide 156 phases. Adsorption of anionic V (H<sub>2</sub>VO<sub>4</sub><sup>-</sup>, HVO<sub>4</sub><sup>2-</sup>) is 157 much lower than the cations; however, VO<sup>2+</sup> solubility may 158 be greatly increased through complexation with organic mat- 159 ter. Whereas V(IV) is not thermodynamically stable at 160 pH > 7, complexation by various organic and inorganic spe-161 cies may considerably increase its stability. Eventually, V 162 (V) oxidation state ion is more toxic than V(IV) ion one. 163

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## 164 Biological Utilization and Toxicity

Whereas low concentrations of V are required for plants to 165 positively influence chlorophyll synthesis, K consumption, or 166 N assimilation, higher V concentrations are toxic as they 167 cause chlorosis and limit growth. At trace amounts V is essen-168 tial for normal cell growth, whereas possibly toxic at high 169 concentrations, especially when occurring as pentoxid 170 Chronic exposure may result in inflammation of bronchi and 171 trachea, eyes and skin irritation, pulmonary edema, and sys-172

173 temic poisonii

#### 174 Summary

Vanadium's occurrence receives more attention as the global 175 demand for V increases. However, its prevalence in environ-176 ment becomes a new critical issue. Vanadium has been 177 suggested to be a potentially dangerous pollutant, and the 178 United States Environmental Protection Agency classifies 179 V in the priority list of environmental risk elements. Fixing 180 the environmental issue of V is constrained by the limited 181 understanding of V biogeochemistry relative to other d-block 182 transition metals. А better understanding 183 of V biogeochemical behavior may support to assess the risk to 184 the environment and to human health and to assist in devel-185 oping new remediation tools. 186

#### 187 Cross-References

- 188 

  Complexation
- 189 Lithophile Elements
- 190 **•** Oxidation-Reduction
- 191 **•** Transition Elements

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