CHAPTER 5

Radionuclides in Nature

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Practically all natural materials contain radioactive nuclides, though usually in such low concentrations that they can be detected only by very sensitive analysis. This is true for water (rain water, rivers, lakes, sea), rocks and soil, and all living matter, as well as for

TABLE 5.1.a. Long-lived cosmogenic radionuclides appearing in meteorites and rain water

Nuclide	Half-life (years)	Decay Mode & particle energy (MeV)	Atmospheric production rate (atoms m ⁻² s ⁻¹)	
3 _H	12.32	β 0.0186	2500	
¹⁰ Be	1.52×10^{6}	β^{-} 0.555	300	
¹⁴ C	5715	β^{-} 0.1565	17 000-25 000	
²² Na	2.605	β^{+} 0.545	0.5	
²⁶ Al	7.1×10^5	β^+ 1.16	1.2	
³² Si	160	β^{-} 0.213	1.6	
³⁵ S	0.239 (87.2 d	$\beta^{-} 0.167$	14	
³⁶ Cl	3.01×10^{5}	β^{-} 0.709	60	
³⁹ Ar	268	β^{-} 0.565	56	
⁵³ Mn	3.7×10^{6}	EC (0.596)		
⁸¹ Kr	2.2×10^{5}	EC (0.28)		

Values within parenthesis after EC are decay energies.

structures based on natural raw materials except where great care has resulted in use of a non-radioactive material. The radionuclides in our environment can be divided into (i) those formed from cosmic radiation, (ii) those with lifetimes comparable to the age of the earth, (iii) those that are part of the natural decay chains beginning with thorium and uranium, and (iv) those introduced in nature by modern techniques. Sources can be categorized as: (i) cosmogenic, (ii) and (iii) primordial, and (iv) anthropogenic.

5.1. Cosmogenic radionuclides

5.1.1. *Survey*

Cosmic irradiation of the atmosphere produces neutrons and protons (Ch. 10) which react with N_2 , O_2 , Ar, etc. resulting in the production of radioactive nuclides, some of which are listed in Table 5.1 a and b. These nuclides are produced at constant rates and brought

TABLE 5.1.b. Short-lived cosmogenic radionuclides appearing in rain water

Nuclide	Half-life	Decay mode and particle energy (MeV)
⁷ Be	53.28 d	EC (0.862)
²⁴ Na	14.96 h	β^{-} 1.389
²⁸ Mg	21.0 h	β^{-} 0.459
³² P	14.28 d	β^{-} 1.710
³³ P	25.3 d	β^{-} 0.249
³⁹ Cl	55.6 min	β^- 1.91

Value within parenthesis is decay energy Production rates (atoms m⁻² s⁻¹): ⁷Be 81, ³⁹Cl 16 to the earth surface by rain water. Though they are formed in extremely low concentrations, the global inventory is by no means small (§§5.1.2 and 5.1.3). Equilibrium is assumed to be established between the production rate and the mean residence time of these radionuclides in terrestrial reservoirs (the atmosphere, the sea, lakes, soil, plants, etc) leading to constant specific radioactivities of the elements in each reservoir. If a reservoir is closed from the environment, its specific radioactivity decreases. This can be used to determine exposure times of meteorites to cosmic radiation (and the constancy of the cosmic radiation field, using ⁸¹Kr), dating marine sediments (using ¹⁰Be, ²⁶Al), groundwater (³⁶Cl), glacial ice (¹⁰Be), dead biological materials (¹⁴C), etc. The shorter-lived cosmogenic radionuclides have been used as natural tracers for atmospheric mixing and precipitation processes (e.g. ³⁹Cl or ³⁸S). Only T and ¹⁴C are of sufficient importance to deserve further discussion.

5.1.2. Tritium

Satellite measurements have shown that the earth receives some of the tritium ejected from the sun. Much larger amounts are formed in the atmosphere through nuclear reactions; e.g., between fast neutrons and nitrogen atoms

$$n(fast) + {}^{14}N \rightarrow {}^{12}C + {}^{3}H$$
 (5.1)

The yield for this reaction is about 2500 atoms tritium per second per square meter of the earth's surface; the global inventory is therefore about 1.3×10^{18} Bq. Tritium has a half-life of 12.33 y, decaying by weak β emission to 3 He. It is rapidly incorporated in water, entering the global hydrological cycle. The average residence time in the atmosphere is about 2 y which is a small fraction of the half-life, as once the tritiated water reaches the lower troposphere, it rains out in 5 - 20 days. If we define 1 TU (Tritium Unit) as 1 tritium atom per 10^{18} hydrogen atoms, 1 TU corresponds to 118 Bq/m³. Before the advent of nuclear energy, surface waters contained 2 - 8 TU (an average value of 3.5 TU is commonly used). The tritium content in water now commonly is of the order 20 - 40 TU. Rainwater contains between 4 and 25 TU, lower at the equatorial zone and increasing with latitude.

Tritium is also a product in the nuclear energy cycle, some of which is released to the atmosphere and some to the hydro sphere. The emissions differ between reactor types (usually in the order HWR > PWR > BWR, see Ch. 19) and is a function of the energy production. Assuming the annual releases to be 40 TBq/GW_e (Giga Watt electricity) from an average power plant and 600 TBq/GW_e from a typical reprocessing plant, the annual global injection of tritium in the environment is estimated to ~10 PBq in 1992. Though this is a small fraction of the natural production, it causes local increases.

The hydrogen bomb tests conducted in the atmosphere during the decade of the 1950's and early 1960's injected large amounts of tritium into the geosphere; 2.6×10^{20} Bq up to the end of the tests in 1962. This considerably exceeds the natural production inventory.

Before 1952 (first hydrogen bomb tests) the tritium content could be used to date water (i.e. determine when it became isolated from contact with the atmosphere). This was very

useful e.g. for determining ice ages. However, due to the much larger content of anthropogenic tritium presently, this is no longer a useful technique for such dating.

Tritium in concentrations as low as 1 TU can be measured in low background proportional counters, and, after isotope enrichment (e.g. by electrolysis of alkaline water, by which tritium is enriched in the remainder), down to 0.01 TU. For very low concentrations mass spectrometry is preferred.

5.1.3. 14-Carbon

¹⁴C is produced in the atmosphere by a variety of reactions, the most important being between thermalized neutrons from cosmic radiation and nitrogen atoms:

$$n(slow) + {}^{14}N \rightarrow {}^{14}C + {}^{1}H$$
 (5.2)

This reaction occurs with a yield of approximately 22 000 atoms 14 C formed per s and m² of the earth's surface; the global annual production rate is ~ 1 PBq, and global inventory ~ 8500 PBq (corresponding to ~ 75 tons). Of this amount ~ 140 PBq remain in the atmosphere while the rest is incorporated in terrestrial material. All living material (incl. body tissue) has a 14 C concentration of ~ 227 Bq/kg. The half-life of 14 C is 5715 y; it decays by soft β^- emission (E_{max} 158 keV).

 14 C is also formed by reaction (5.2) in nuclear tests. From these 220 PBq is assumed to have been injected into the atmosphere up to 1990. This 14 C comes to equilibrium with other atmospheric carbon (CO₂) in 1 - 2 years. Some 14 C, about 18 TBq/GW_e per year, is also released from nuclear power plants (mainly from HWR, Ch. 19 and 22). The global atmospheric value is < 300 TBq/y.

The combustion of fossil fuel adds CO_2 , which is almost free of ¹⁴C, to the atmosphere, thus reducing the specific activity (the dilution was about 3% for the period 1900 - 1970). Taking all anthropogenic sources into account, a global average specific activity of modern carbon is now 13.56 \pm 0.07 dpm/g C. In §5.8.1 we discuss dating by the "C14-method", for which such a figure is important.

5.2. Primordial radionuclides

5.2.1. Very long-lived nuclides lighter than lead

As the detection technique for radioactivity has been refined, a number of long-lived radionuclides have been discovered in nature. The lightest have been mentioned in §5.1. The heavier ones, not belonging to the natural radioactive decay series of uranium and thorium, are listed in Table 5.2. ⁵⁰V is the nuclide of lowest elemental specific activity (~0.0001 Bq/g) while the highest are ⁸⁷Rb and ¹⁸⁷Re (each ~900 Bq/g). As our ability to make reliable measurements of low activities increases, the number of elements between potassium and lead with radioactive isotopes in nature can be expected to increase.

Nuclide	Isotopic abundance %	Decay mode and particle energy MeV	Half-life (years)	
40 _K	0.0117	β EC 1.31	1.26×10 ⁹	
50 _V	0.250	β^{-} EC (0.601)	$>1.4\times10^{17}$	
⁸⁷ Rb	27.83	β^{-} 0.273	4.88×10^{10}	
115 _{In}	95.72	β^- 1.0	4.4×10^{14}	
¹²³ Te	0.905	EC (0.052)	1.3×10^{13}	
¹³⁸ La	0.092	β EC	1.06×10^{11}	
¹⁴⁴ Nd	23.80	α	2.1×10^{15}	
¹⁴⁷ Sm	15.0	α 2.23	1.06×10^{11}	
¹⁴⁸ Sm	11.3	α 1.96	7×10^{15}	
¹⁷⁶ Lu	2.59	β^{-} (1.188)	3.8×10^{10}	
¹⁷⁴ Hf	0.162	α	2×10^{15}	
¹⁸⁷ Re	62.60	β^{-} 0.0025	4.2×10^{10}	
190Pt	0.012	α	6.5×10^{11}	

TABLE 5.2. Primordial radionuclides for Z < 82 (Pb)

Values within parenthesis refer to decay energies.

Because of the long half-lives of these nuclides they must have been formed at the time of (or possibly even before) the formation of the solar system and of the earth. When the earth's crust solidified, these radionuclides became trapped in rocks. As they decayed, decay products accumulated in the closed rock environment. By measuring the amount of parent and daughter nuclides, it is possible with the half-life to calculate how long this environment (e.g. a rock formation) has existed. This is the bases for nuclear dating (also called "radioactive clocks"), and almost all of the nuclides in Table 5.2 can be used for this purpose. In § 5.8 we discuss dating methods for the K-Ar and Rb-Sr systems.

A careful look at these naturally occurring long-lived nuclei reveals that some of them appear in *short* decay series, e.g. $^{152}\text{Gd} \rightarrow ^{148}\text{Sm} \rightarrow ^{144}\text{Nd} \rightarrow ^{140}\text{Ce}$ and $^{190}\text{Pt} \rightarrow ^{186}\text{Os} \rightarrow ^{182}\text{W}$. The heavy element series beginning with U and Th isotopes are therefore referred to as *long* decay series.

5.2.2. Elements in the natural radioactive decay series

In Chapter 1 we briefly discussed the existence of four long series of genetically related radioactive nuclides beginning with Th, U or Np and ending with Pb or Bi. In Figure 5.1 we present all known isotopes of elements $_{81}$ Tl to $_{92}$ U. Some of these nuclides occur naturally in the long decay series shown in Figure 2.1\frac{1}{2}. Others are produced through nuclear reactions according to the schemes in Figure 4.8 and principles described in Ch. 12-15. The first series in Figure 1.2 is known as the thorium decay series, and consists of a group of radionuclides related through decay in which all the mass numbers are evenly divisible by four (the 4n series). It has its natural origin in 232 Th which occurs with 100% isotopic abundance. Natural thorium has a specific activity S of 4.06 MBq/kg, as its

¹ The student can easily trace the decay series in Fig. 5.1 with a transparent marker.

half-life through α -decay is 1.41×10^{10} y. The terminal nuclide in this decay series is the stable species ^{208}Pb (also known as ThD). The transformation from the original parent to the final product requires 6 α - and 4 β -decays. The longest-lived intermediate is 5.76 y ^{228}Ra .

The uranium decay series consist of a group of nuclides that, when their mass number is divided by 4, have a remainder of 2 (the 4n + 2 series). The parent of this series is 238 U with a natural abundance of 99.3%; it undergoes α -decay with a half-life of 4.46×10^9 y. The stable end product of the uranium series is 206 Pb, which is reached after 8 α - and 6 β -decay steps.

The specific activity of ²³⁸U is 12.44 MBq/kg ²³⁸U. However, because *natural uranium* consists of 3 isotopes, ²³⁸U, ²³⁵U and ²³⁴U, whose isotopic abundances are 99.2745%, 0.7200% and 0.0055%, respectively, the specific activity of natural uranium is 25.4 MBq/kg.

The uranium decay series provides the most important isotopes of elements radium, radon, and polonium, which can be isolated in the processing of uranium minerals. Each ton of uranium is associated with 0.340 g of 226 Ra. Freshly isolated 226 Ra reaches radioactive equilibrium with its decay products to 210 Pb in about two weeks (see Fig. 1.2). Many of these products emit energetic γ -rays, which resulted in the use of Ra as a γ -source in medical treatment of cancer (radiation therapy). However, the medical importance of radium has diminished greatly since the introduction of other radiation sources, and presently the largest use of radium is as small neutron sources (see Table 12.2).

Although the chemistry of radium is relatively simple (like barium), the fact that it produces a radioactive gas (radon) complicates its handling. The decay of radon produces "airborne" radioactive atoms of At, Po, Bi, and Pb. Since uranium is a common element in rocks (see §5.4) it is also a common constituent of building materials. Such material emits Rn, as discussed further in §5.6. Work with radium compounds should be carried out within enclosures to avoid exposure to Rn and its daughters.

The actinium decay series consists of a group of nuclides whose mass number divided by 4 leaves a remainder of 3 (the 4n + 3 series). This series begins with the uranium isotope 235 U, which has a half-life of 7.04×10^8 y and a specific activity of 8×10^4 MBq/kg. The stable end product of the series is 207 Pb, which is formed after 7 α - and 4 β -decays. The actinium series includes the most important isotopes of the elements protactinium, actinium, francium, and astatine. Inasmuch as 235 U is a component of natural uranium, these elements can be isolated in the processing of uranium minerals. The longest-lived protactinium isotope, 231 Pa ($t_{1/2}$ 3.28 \times 10⁴ y) has been isolated on the 100 g scale, and is the main isotope for the study of protactinium chemistry. 227 Ac ($t_{1/2}$ 21.8 y) is the longest-lived actinium isotope.

5.3. Transuranic elements in nature and the Np decay series

A fourth long radioactive decay series, the *neptunium series* (Fig. 1.2), is composed of nuclides having mass numbers which divided by 4 have a remainder of 1 (the 4n + 1 series). The name comes from the longest lived A = 4n + 1 nuclide heavier than Bi, 237 Np, which is considered as the parent species; it has a half-life of 2.14×10^6 y. Inasmuch as this half-life is considerably shorter than the age of the earth, primordial 237 Np no longer exists on earth, and, therefore, the neptunium series is not found as a natural

Part								
256.0254 Part Part	Ra 202 0.7 ms a 7.86	Fr 201 48 ms a 7.388/100	Rn 200 1.0 s a 6.902/~96	At 199 7.2 s a 6.64/90	50	Bi 197 4.9m 9.3m a 5.77 e/100 /~0.11 m.g g c y 866	Pb 196 36.4 m e/100 g y 283/?	TI 195 3.6s 1.13h 17/100 4; \$ 7.384 1.8/3 g /91 7.564
Second	88Ra 226.0254	Fr 200 19 ms a 7.488	Rn 199 0.32s 0.62s a 7.06 a 7.00 m 9	At 198 1.0s 4.2s a 6.86 a 6.76 /84 m /90 g	Po 197 26s 56s a 6 39 a 6 28 /84 /44 c/16 c/56	196 196 196 196 196 196 196 196 196 196	Pb 195 15.0m ~ 15m 6.5i m e/100 7100 9 7384 7384	E m
Back		87FR		At 197 3.7 s 0.35 s a 6.71 a 6.96		Bi 195 1.5m 3.0m a 6 11 a 5.42 /4 g /<0.2	4 -	
Sept			197 66ms a7.260 9		Po 195 193 463 0670 0651 7-90 775 9:0 m;0	1.6m a 5.65	193 4.0m 4/100 9 7365	192 9.6m v: p
Sabra Sabr			96 8		Po 194 0.39 s 0.846/~ 100			- 191
Sapeta S			Constitution of the last of th	At 194 0.18 s			191 1.4m 0 5.29 7~0.6;	
S4Po Po 190 Po 191 Po					N 10	Bi 191 115s 12s 3 16.88 a 6.31 c 550g 760 c	0	
84Po Po 190 2.4 ms 208.9804 2.86.4 ps 208.6 p			,		91 sm		68 83	T1 188
84Po 1089-904 208-9					90 sr	189 s 0.88s a 6.67		
Ph 180								86 28.1s 28.1s 3.5.66 7.7.0
Pb 180				,			98	T1185
Pb 180						86 15 ms 2 7 18	22	**
83Bi 208.9804 Deb 180 Pb 181 Pb 182 Pb 183 4 ms 45 ms 55 ms 6.30 s 4 ms 7 21/2						35	8 ° °	
Pb 180 Pb 181 Pb 182 2							~	
Pb 180 Pb 181 45 ms 45 ms 45 ms 1.5 s 83 7.21 0 645 6.28							Pb 182 55 ms a 6.921/?	T1 181
Pb 180 4 ms 5 4 ms 1							18 sr	0
2 2 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1							0	
								83

FIG. 5.1. Chart of all known isotopes of elements 81Tl to 92U. The legend to the chart is given in Fig. 4.8.

U 224 0.7 ms	Pa 223 6.5 ms a/100 8.006/55	Th 222 2.2 ms a/100 7 982	Ac 221 52 ms a/100 7.848/70	Ra 220 23 ms 2100 7 455/99	Fr 219 21 ms av100 7.312/99	Rn 218 35 ms 0x/100 /.133 /90.8	At 217 32.3 ms a/~ 100 7.069 /96.6 p /0.01	Po 216 150 ms a.100 8.779/~100	8i 215 7.6 m 7100 2847	Pb 214 26.8 m \$ 0.7/100 y 352/37	
U 223 U 18 µ8 C C C C C C C C C C C C C C C C C C	Pa 222 P. 4.3 ms 6 4.100 8.21 ar/10	Th 221 Th 1.68 ms 24/100 8 146/56 avic	Ac 220 A 28 ms	Ra 219 R 10 ms a/100 7.878/96 a/10 17.316	Fr 218 F 22ms 1.0ms a 2.10	Rn 217 R 0.54 ms av100 7.740/100 /393.	At 216 A 3 3 3 3 4 2 100 0 100	Po 215 P 1.780 ms 1 a/~100 2.386/100 a.10	Bi 214 B	Pb 213 P 210.2 m 2 6.100 proy	
U 222	Pa 221 5.9 μs σ 9.06	Th 220 9.7 μs α 8.790/100	Ac 219 11.8 µs α 8.664/100	Ra 218 25.6 µs a 8.350/100	Fr 217	Rn 216 45 µs a 8.050/100	At 215 0.1 ms 8.026/~100	Po 214 163.7 µs a/100 7.687;-100	Bi 213 45.59 m a/25.869/2 § 1.4/88	Pb 212 10.64 h 6 0.3/100 7 239/44	
	Pa 220 0.78 µs	Th 219 1.05 µs a 9.340/100	Ac 218 1.1 µs α 9.206/100	Ra 217 1.6 µs a 8.992/100	Fr 216 0.70 µs a 9.006/100	Rn 215 2.3 µs α 8.6/4/100	At 214 760 270 560 11	Po 213 4.2 µs a/100 8.376/100	Bi 212 25m 1.008h a.93 a.36 6.34 6.05 753 /26	Pb 211 36.1 m 8-1.4100 y 405/4, 832/4	T1 210 1.30 m \$ 1.8/100 y 798/99
	Pa 219 53 ns	Th 218 109 ns u 9.865/100	Ac 217 0.74µ 69ns a a 10.54 9.65	Ra 216 2.0ns 0.18µs 0.55 9.36	Fr 215 0.09 µs 0.360/100	Rn 214 6.6 0.7 0.2 ne ne 7 a a ne 10. 10 a	At 213 0.11 µs	Po 212 45s 298ns a/100 a 11.7 8.784 /87 y /100	Bi 211 2.17 m a.89.76623 .84 9	Pb 210 22.3 γ α/2×10° 3.720; p/100	71 209 2.18 m \$ 1.8/100 y 1568/98
U 219 ~42 µs	Pa 218 0.12 ms	Th 217 252 µs 4 9.250/100	Ac 216 0.33 -0.33 ms ms au100 a/100 9.03 9.07	Ra 215 1.6 ms artoo 8.700/98	Fr 214 3.35ms 5.0 ms ar100? ar100 8.476 8.427 761 193	Rn 213 25.0 ms a/100 8.087/99;	At 212 119ms 314ms ar100 ar100 7.84 7.88 765 784	Po 211 25.5a 516ma c/100 c/100 7.27 7.45 /81 /99:	Bi 210 3.0 × 5.013d 10 ⁴ y α 4.68 α/100 / ~0; β 4.86 /~100	Pb 209 3.263 h 8.0.6/100	TI 208 3.053 m 8 1.8/100 v 2614/99.8
U 218	Pa 217 1.6ms 4.9ms 10.16 8.33	Th 216 0.18ms 28ms 0.9m 2.92	Ac 215 0.17 s a 7.604:99.91	Ra 214 2.46 s a 7 136/39.94 c,0.06	Fr 213 34.6 s a 6.776:98.4	Rn 212 24 m 0:100 6:250/-100	At 211 7.22 h 0.5867/42 e/58.9	Po 210 138.376 d α 5.304/100 γ 803/0 001	Bi 209 100 % u0011+0023	Pb 208 52.4 % 9 0.000487	TI 207 1.33s 4.77m 17.100 pr.100 y 1000 y 897 187 /0.2
92U 238.0289 07.59.278 0 _{6,7} 3.4 0 _{6,6} 2.2	Pa 216 0.2 s a/~80 7 87	Th 215 1.2 s w/100 7 395/52	Ac 214 8.2 s or>96 7.214/45	Ra 213 2.1ms 2.7m o/~1 o/80 8.47 6.62 // /39:c	Fr 212 20.0 m 0/43 6.262/16 к/57	Rn 211 14.6 h av26 5.74; y 674	At 210 8.1 h a.o.2 5.524/0.05	Po 209 102 y ¤ 4.880/99 c/0.3	BI 208 3.68 × 10 ⁵ y c/100 y 2614/100	Pb 207 22.1 % 0 0.709	TI 206 TI 207 3.76m 4.20m 1.33 4.77m II/100 61.5 IT/100
	Pa 215 14 ms « 8.09	Th 214 0.10 s a 7 677/100	Ac 213 0.80 s a 7.364/100	Ra 212 13 s a 8 901/7	Fr 211 3.10 m a 6.534/> 90 e/< 20	Rn 210 2.4 h 2.96 6.040/96 20/47/458/1.6	At 209 5.41 h α 5.641/4 «/96	Po 208 2.898 y a 5.116/100	Bi 207 31.55 y c/~100 g*/0.01	Pb 206 24.1 % o 0.0305	TI 205 70.476 %
	Pa 214 17 ms a 8.12	Th 213 0.14 s	Ac 212 0.93 s a 7.379/100	Ra 211 13 s a 6.911,?	Fr 210 3.18 m a 6.543/00	Rn 209 28.5 m 0.17 6.039.17 ; 0.79; § 1/4	At 208 1.63 h axx,6 6.6410.6 k/90:p*/3	Po 207 2.8s 5.84h IT/100 a 5.12 y 814 /0.02 /99 e/99 g	Bi 206 6.24 d c.p' /100 y 803/88	Pb 205 1.53×10 ⁷ y 100 y	TI 204 3.78 y 6/2.6 8 0.8/97.5
	Pa 213 5.3 ms a 8.24	Th 212 30 ms	Ac 211 0.25 s 07.481//	Ra 210 3.7 s a 7.019/100	Fr 209 50.0 s 6.646/100	Rn 208 24.4 m α/80 8.138/80 ; ε/40; γ 427	At 207 1.8 h α 5.759/10 ε:β** /90	Po 206 8.8 d a 5.223/5 c/96 g	Bi 205 15,31 d 0/~100 8*/0:1	Pb 204 1.4 % 1.12 h	T1 203 29.524 %
	Pa 212 5.1 ms α 8.27	Th 211 37 ms	Ac 210 0.35 s c 7.462/100	Ra 209 4.6 s e 7.010/100	Fr 208 58.6 s e.638/100	Rn 207 9.3 m α/23 θ.126/23 : c;β'/77 g	At 206 29.4 m ¤ 5.70311; c:82 µ-3.117	Po 205 1.86 h a 5.22/0 1 g c/98: 5 ⁻ /2	Bi 204 11.22 h c/100 v 899/98	Pb 203 6.3s 2.169d IT/100 e/100 y 826 y 279 /72 /80	TI 202 12.23 d c/100 y 440/91
	91P8 231.0359	90Th 232.0381 07.40.85	Ac 209 90 ms a 7.59/100	Ra 208 1.3 s 0.7 133:-95	Fr 207 14.8 s a 6.766/93	Rn 206 5.67 m a 6.260:68 c/32 y 498:7	At 205 26.2 m a 5.902:10 k:p* /90	Po 204 3.53 h a 5.3770.7 c.99.3	Bi 203 11.76 h c/~100 g.m p^~ 1.40.2	Pb 202 3.62h 8.3× ε/10 10 ⁴ /γ 17/90 ε/100 1,961 no γ	TI 201 3.05 d 6/100 y 167/9
			Ac 208 25ms 96ms a 7.76 a 7.57 m2 9	Ra 207 55ms 1.3s a 7.32 a 7.13 /15 m /80 g	Fr 206	Rn 205 2.83 m a/23 6.263/23 7.265/2	At 204 9.2 m α 5.961.4 ε/68:β ⁻ /29:	Po 203 45s 36m IT 05.384 U A.1 641/51 c/93;	Bi 202 1.72 h c, p'; g v 579/7	Pb 201 1.02m 9.33h IT/100 c/-10 1,629 0 /54 p+	TI 200 1.088 d e/-100 b+/0.4
			Ac 207 22 ms	Ra 206 0.24 s a 7.270100	Fr 205	Rn 204 1.24 m a 6.417/68	At 203 7.4 m a 6.088/31 g	Po 202 44.7 m 44.7 m c.98	Bi 201 59.1m 1.8h α 5.24 ε/100 /~0; ε; γ 629 9 IT: /?	Pb 200 21.5 h «/100 y 148/38	TI 199 7.42 h 0/100 g y 455/12.
			89Ac 227.0278	Ra 205 0.17s 0.21s a 7.37 a 7.34 m	Fr 204	Rn 203 28s 45s a 6.55 a 6.50 /100 /66	At 202	Po 201 8.9 m 15.3 m a 5.75 a 5.88 /3x/57 /2 g	Bi 200 31m 38.4m 2,0 e/89 100 pf:11	Pb 199 12.2m 1.6h 17.83 e.89 e.7 p ¹ .71 1,424 y 387	TI 198 1.87h 6.3h 6.54 6.1577 1746 9412 9412 /83
				Ra 204 59 ms	Fr 203 0.55 s ¤ 7.132/100	Rn 202 9.85 s 0.5.641/85 c/~15.9	At 201 1.5 m 8.34471	Po 200 11.5 m a5.883/15	Bi 199 24.7m 27m a c/100 5.484 y 424/i i < 2.8 m; g	Pb 198 2.4 h c/100 g y 290/36	71 197 2.84 h e: \$7/2 g y 426 f13
				Ra 203 33ms - 1mg a 7.62 a 7.58 m	Fr 202	Rn 201 3.8 s 7.0 s a 6.77 a 6.72 i-90 i-80 em e g	At 200	Po 199 4.2m 5.2m a 6.06 a 5.95 /39 m /12 g c/81 g c/88 m	Bi 198 7.7 11. 10. 8 8 3 If m m I, 6; 6;	Pb 197 43m ~8m c/81 c.p ⁺ 17/19 /100 y 386 y 386	T1 196 1.41h 1.84h 6/96 e; IT/5 \$ ⁴ /14 y 426 y 426

FIG. 5.1. Continued from previous page.

U 242 16.8 m \$ 1000 Y 58.79									
U 240 14.1 h \$ 04100 744/2			1						
U 239 23.5 m p 12/100 y 75/52	Pa 238 2.3 m p 1.7/100 y 1015	Th 237 5.0 m 8 /100							
U 238 99.2745 % 4.47 × 10 ⁹ y g	Pa 237 8.7 m β 1.4/100 γ 854/34	Th 236 37.5 m g 1.0/100 v 111/4		Rs 234 30 s 87100					
0.237 6.75 d 8.02 ./100 7.60/33	Pa 236 9.1 m 8 20./100 y 642/29	Th 235 7.1 m p 1.4/100	Ac 234 44 s p /100 v 1847	Ra 233 30 s 67100	Fr 232 5 s 67100 7126				
U 236 2.34 × 10 ⁷ y 0/100 4.494/74 y 49/0.08	Pa 235 24.2 m p 14/100 y 128	Th 234 24.10 d p 0.2100 y 63/4	Ac 233 2.42 m p.100 y 523	Ra 232 4.2 m p.//oo y 471	Fr 231 17.5 s 67,100 7,433				
0.0055 % 0.7200 % 2.46 × 105 y 7.04 × 108 y 0.100 4.73573 ar × 100 4.388	Pa 234 1.17m 6.70h 17/0.13 po.6 \$2.3/100 /99.97 y131	Th 233 22.3 m g 1.2/100 v 28/2.6	Ac 232 1.98 m b /100 y 885	Ra 231 1.72 m p-7100 y 410	Fr 230 19.1 s p7100 y711				
	Pa 233 27.0 d p 03 /100 y 312/36	Th 232 1.41×10 ¹⁰ y α/100 4.010/77y	Ac 231 7.5 m y 283	Ra 230 1.55 h p. 0.3/100 y 72	Fr 229 50.2 s p-7100 y 310	Rn 228 1.08 m g /100 y 125			
U 233 1.59 × 10 ⁵ y α:'-100 4 824 /84.y 42/0.06	Pa 232 1.31 d 1.31 d p 0.3	Th 231 1.063 d 8 0.3/100 7 26/15	Ac 230 2.03 m p 2.7/100 v 455/9	Ra 229 4.0 m β-1.8/100	Fr 228 39 s p7100 y 474	Rn 227 22.5 s p.100 y.162			
U 232 68.9 y a/~100 5 320 768 .: Y58/0.2	Pa 231 3.28 × 10 ⁴ y a/100 5.014/25	Th 230 7.54 × 10 ⁴ y α:1004.88778 : y 68:0.4	Ac 229 1.045 h p 1.1/100 y 165	Ra 228 5.75 y \$10.04/100 y 14	Fr 227 2.47 m 8° 1.8/100 y 90	Rn 226 7.4 m p.1100			
U 231 4.2 d α 5.4560.006 ρ/99.996 γ 26/13	Pa 230 17.4 d 0/0.003 5.348 /0.0007; c/9 p0.5/10; y	Th 229 7880 y 7.54 × 10 ⁴ y a/100.4.845/56 a/100.4.887/78: y 134/5 y 68/0.4	Ac 228 6.13 h a4.270:6×10° p 1.2/~100	Ra 227 42.2 m F 1.3/100 y 27/17	Fr 226 48 s p 3.2/100 y 254	Rn 225 4.5 m \$ 729			
U 230 20.8 d a/100 5.888/67	Pa 229 1.50 d α.0.25 5.580 λ0.09 c/99.75 γ	Th 228 1.913 y 0.100 5.423/73	Ac 227 21.773 y a/1.4 4.853/0.85	Ra 226 1600 y a/100 4.784/94 , y 186/3	Fr 225 4.0 m p-1.6/100 y 182	Rn 224 1.78 h p./100 y.281.	At 223 50 s \$7100		
U 229 58 m a/-20 6.362/64 c-80, y 123	Pa 228 22 h α/~2 6.078 c/~96, β*/0.2 γ 310	Th 226 Th 227 31 m 18.72 d add 6.038/26 add	Ac 225 Ac 226 10.0 d 1.2 d 1.2 d 1.00 d 2.3990.008	Ra 225 14.8 d p. 03/100 y 40/29	Fr 224 3.3 m p. 2.6/100 y 216	Rn 223 23.2 m g / 100 y 593	At 222 54 s		
U 228 9.1 m a/>96 6.681/70; c/ < 5, y 95/2	Pa 227 38.3 m a/~86 6.466/43 , c/~16	31 m 21 m 21 m 21 00 6.338/78	Ac 225 10.0 d a/100 5.830/52	Re 224 3.66 d or - 100 5.886 /96y241/4	Fr 223 21.8 m or 6.340/0.006 § 1.1/99.994	Rn 222 3.825 d a/100 5.490 /99.9:	AT 221 2.3 m gr100		
U 227 1.1 m a/100 6.86	Pe 225 Pe 226 Pe 38.3 1.8 s 1.8 m 38.3 0x:1007.246/70 0774 6.863/39 0x-86 0x;00.726 6.46645	Th 225 8.72 m av~90 6.479 ; e/~10	Ac 224 2.9 h av~10 6.138 ; e/~90	Re 223 11.43 d ar~100 6.716 /63;7269/14	Fr 222 14.2 m av<1 \$1.8/>99	25 m 25 m a/22 6 037/18 p0.8/78	At 220 3.71 m a 5.493/8 p.92		
U 226 0.2 s a 7.57	Pa 225 1.8 s ¤/100 7.246/70	Th 223 Th 224 Th 225 m 2	Ac 223 2.10 m ays 8.847/44 : e/1	Ra 222 38.0 s av~100 6.666 /97y324/3	4.9 m 4.9 m 4.100 6.341/83	Rn 220 55.6 s a/100 6.288/99.93	At 219 54 s a e 275/97 p/3	Po 218 3.05 m a/~100 8.002/100	
U 225	Ps 224 0.95 s α/100 7.856	Th 223 0.66 \$ @/1007.287/80 y 140	Ac 222 1.05m 5.0s a/>89 a/100 6.81 7.01 124 /94	Ra 221 28 s 02100 6.813/38 7 90/15	Fr 220 27.4 s ar99.6 5.686 § 70.4, y 45/2	Rn 219 3.96 s a/100 6.819/81	At 218 -2 s -2 s 6.696/90	Po 217 < 10 s a 6.539/100	Bi 216 3.6 m y 550?

FIG. 5.1. Continued from previous page.

occurrence. However, Np has been discovered in the spectrum of some stars. All known Np-isotopes are presented in Figure 16.1.

Very small amounts of 237 Np, as well as of 239 Pu, have been discovered on earth; the half-lives of 239 Pu (in the 4n+3 series) is 2.411×10^4 y. Both isotopes are too short-lived to have survived the 4 eons since the solar system was formed. However, they are always found in minerals containing uranium and thorium and it is believed that the neutrons produced in these minerals through (α,n) and (γ,n) reactions with U and Th as well as by spontaneous fission of 238 U form the neptunium and plutonium through n-capture and β -decay processes. The n-production rate in the uranium mineral pitchblende (containing $\sim 50\%$ U) is about 50 n/kg s. The typical value for the 239 Pu/ 238 U ratio in minerals is 3×10^{-12} .

The end product of the neptunium series is 209 Bi, which is the only stable isotope of bismuth. Seven α - and four β -decays are required in the sequence from the parent 237 Np to 209 Bi. An important nuclide in the neptunium decay series is the uranium isotope 233 U, which has a half-life of 1.59×10^5 y (the most stable intermediate) and, like 235 U, is fissionable by slow neutrons.

The long-lived plutonium isotope 244 Pu (belonging to the 4n series; see also Fig. 16.1), which decays through α -emission and spontaneous fission (0.13%) with a total half-life of 8.26×10^7 y, was discovered in rare earth minerals in 1971. If this is a survival of primeval 244 Pu, only 10^{-15} % of the original can remain. An alternate possibility is that this 244 Pu is a contaminant from cosmic dust (e.g. from a supernova explosion in more recent times than the age of the solar system).

5.4. Thorium

5.4.1. Isotopes

Natural thorium consists 100% of the isotope 232 Th which is the parent nuclide of the thorium decay series. The specific radioactivity for thorium is lower than that of uranium, and it is normally treated as a non-radioactive element. For radioactive tracer studies the nuclide 234 Th ($t_{1/2}$ 24.1 d) is used after separation from natural uranium.

5.4.2. Occurrence and production

Thorium is somewhat more common in nature than uranium, with an average content in the earth's crust of 10 ppm (by comparison the average abundance of lead is about 16 ppm in the earth's crust). In minerals it occurs only as oxide. The content of thorium in sea water is $< 0.5 \times 10^{-3} \text{ g/m}^3$, which is lower than that of uranium because of the lower solubility of Th⁴⁺ compounds (the most stable valency state of Th).

The most common thorium mineral is monazite, a golden brown rare earth phosphate containing 1 - 15% ThO₂ and usually 0.1 - 1% U₃O₈. It is also found in small amounts in granite and gneiss. The largest deposits of monazite are found in India, Egypt, South Africa, the USA, and Canada, with 200 - 400 kton ThO₂ in each country. The size of natural resources are defined in terms of ore reserves which can be economically processed. Thus, the total reserves at commercial price in 1991 was estimated to > 2 Mt ThO₂.

Because thorium often occurs with other valuable metals (in addition to the lanthanides) such as niobium, uranium and zirconium, it can be produced as a byproduct.

The following procedure is used for producing thorium from monazite sand. The sand is digested with hot concentrated alkali which converts the oxide to hydroxide. The filtered hydroxide is dissolved in hydroxhloric acid and the pH adjusted between 5 and 6, which precipitates the thorium hydroxide but not the main fraction of lanthanide elements. The thorium hydroxide is dissolved in nitric acid and selectively extracted with methyl isobutyl ketone or tributyl phosphate in kerosene. This gives a rather pure organic solution of $Th(NO_3)_4$. The thorium is stripped from the organic phase by washing with alkali solution.

5.4.3. Uses

Thorium metal is used as electrode material in gas discharge lamps, and as getter for absorption of rest gases in high vacuum technique. ThO₂ (melting point 3300 °C) is highly refractory and used for high temperature furnace linings. Thorium salts are of little practical use. Because Th⁴⁺ is a stable tetravalent ion with properties very similar to the tetravalent actinides, Th⁴⁺ is often used as an analog for the An(IV) ions; most common to use are 228 Th (1.91 y) or 230 Th (7.54 × 10⁴ y) and 234 Th (14 2 24.5 d), which can be isolated from old 232 Th or 238 U, see Fig. 1.2. Thorium may become important to the nuclear energy industry as a fuel in high temperature gas-cooled reactors and may be used in the future in thorium-breeder reactors (Ch. 20).

5.5. Uranium

5.5.1. Isotopes

Natural uranium consists of 3 isotopes, ²³⁴U, ²³⁵U and ²³⁸U, members of the natural decay series discussed in §5.2.2. Uranium is an important raw material for nuclear energy production (see Ch. 19 and 21).

The specific radioactivity of natural uranium makes it a weak radiological hazard (see Ch. 18). It is also chemically toxic and precautions should be taken against inhaling uranium dust for which the threshold limit is 0.20 mg/m³ air (about the same as for lead).

5.5.2. Occurrence, resources and production capacity

Uranium appears in a large number of minerals (at least 60 are known). The earth's crust contains 3 - 4 ppm U, which makes it about as abundant as arsenic or boron. Uranium is found at this relative concentration in the large granitic rock bodies formed by slow cooling of the magma about 1.7 - 2.5 eons ago (1 eon = 10^9 y = 1 billion years). It is also found in younger rocks at higher concentrations ("ore bodies").

Geochemists now begin to understand how these ore bodies were formed. They are usually located downstream from mountain ranges. As the atmosphere became oxidizing about 1.7 eons ago, rain penetrated into rock fractures and pores, bringing the uranium to

its hexavalent state and dissolving it as an anionic complex (likely as a carbonate, $UO_2(CO_3)_n^{2-2n}$, or as a sulphate complex, $UO_2(SO_4)_n^{2-2n}$), often at elevated temperatures. As the water and the dissolved uranium migrated downstream, regions of reducing material were encountered, either inorganic (e.g. pyrite) or organic (e.g. humic) matter, which caused reduction to U(IV). Since most U(IV) compounds are insoluble, the uranium precipitated, possibly as the sulphide or, more likely, as the hydroxide. Many of these original uranium precipitates were later covered by sedimentary material.

In most minerals uranium is in the tetravalent state. The most important one is uraninite $(UO_{2+x}, x = 0.01 \text{ to } 0.25)$, in which the uranium concentration is 50 - 90%; it is found in Western Europe, Central Africa (e.g. Katanga, Gabon) and Canada (e.g. Cigar Lake) and Australia (e.g. Koongara). In the USA and Russia carnotite (a K + U vanadate) is the most important mineral and contains 54% uranium. In the high grade ores the mineral is mixed with other minerals so the average uranium concentration in the crushed ore is much less: e.g. $\leq 0.5\%$ on the Colorado Plateau. Uranium is often found in lower concentration, of the order of 0.01 - 0.03%, in association with other valuable minerals such as apatite, shale, peat, etc.

At the 1999 price of \sim \$30 per kg U, the known (total of *measured*, *indicated* and *inferred*) world reserves of uranium which could be recovered economically was about 3.3 Mton U_3O_8 . This corresponds to \sim 40 years consumption assuming LWR:s and present nuclear capacity to increase by \sim 1% a year. As energy cost increases, it is estimated that about 20 Mton can be recovered at a higher cost. Sea water contains some 4500 Mtons U but it is uncertain if U can be recovered economically from this huge resource.

The uranium production in 1999 was ~ 37 kton of U_3O_8 . The ten largest producers, in decreasing order, were Canada, Australia, Niger, Namibia, Uzbekistan, Russia, USA, Kazakhstan, South Africa and the Czech Republic. The Cigar Lake mine in Canada is interesting for its location (430 m below a lake), size (~ 130 kton U), and high U-content (up to 19%) which poses a difficult working environment (Rn plus quite high levels of γ -radiation); the mine is expected to open in 2005. The past over capacity in U-production has caused large U stockpiles and a small interest in prospecting and opening of new mines.

5.5.3. Production techniques

Uranium ores differ widely in composition, containing a variety of other elements which must be removed. As a result the production methods differ considerably depending on the particular ore to be processed although in every case very selective processes must be used. The following is a common scheme.

The ore is mined in open pits or underground. The ore is crushed and concentrated through flotation. If the uranium is in the tetravalent state it is oxidized in piles by air, sometimes with the aid of bacteria. The material is subsequently treated with sulfuric acid which dissolves the uranium as the sulphate complex, $UO_2(SO_4)_2^{2-}$ (the feed). This complex can be selectively removed from the aqueous solution by means of anion exchange resins or, more commonly, by extraction into an organic solvent (solvent extraction). In the latter case an extractant (i.e. an organic compound with ability to form a U-organic complex soluble in organic solvents) dissolved in kerosene is used; depending on the aqueous feed composition, various extractants have been applied. A typical flow-sheet is illustrated in Figure 5.2, and the basic chemical principles in Table 5.3. The final product is commonly

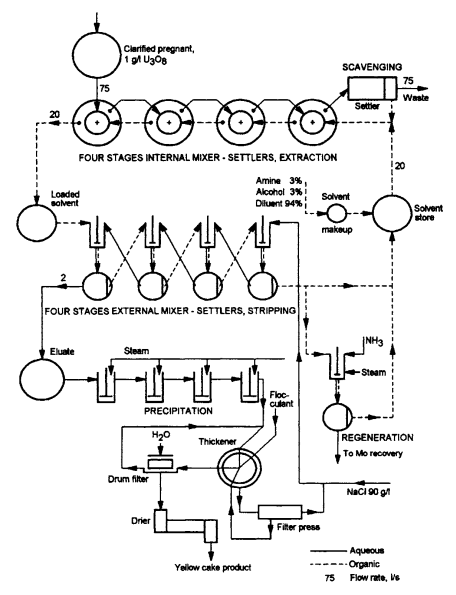


FIG. 5.2. Amine extraction circuit, Kerr-McGee Corp., Grants Mill, New Mexico. Each S-X battery consists of 4 mixer-settlers. Solid lines aqueous, dashed lines organic flow. Numbers are flow rates (l/s).

ammonium diuranate, which is referred to as yellow cake; it contains 65-70% U. It is so free of radioactive uranium daughters that it can be handled safely in drums without any radiation protection. Yellow cake is further purified in order to obtain a final pure product of U₃O₈, usually better than 99.98% pure; its content of neutron poisons (nuclides which have high capture cross-sections for neutrons such as B, Cd, Dy; see Ch. 19) is less than 0.00002%.

Heating yellow cake in hydrogen gas produces UO₂ which in sintered form is used as nuclear reactor fuel. If the UO₂ is exposed to F₂, UF₄ is obtained ("green salt"), which in a thermite process with calcium metal yields metallic uranium. The metal is slowly oxidized in air at room temperature; the metal powder is very reactive at elevated temperatures and can then be used to remove practically all impurities from rare gases.

TABLE 5.3. Chemical bases for the Ames (amine) and Dapex (HDEHP) processes for uranium recovery from sulfuric acid leach liquors. (From Musicas and Schulz.)

	AMEX	DAPEX
Extraction	$UO_2^{2+} + SO_4^{2-} + 2(R_3NH)_2SO_4 \rightleftharpoons$ $(R_3NH)_4UO_2(SO_4)_3$ $R_3N = $ Alamine 336 or Adogen 364 (trialkylamines with alkyl groups with $8 - 10$ carbons)	UO ₂ ²⁺ + 2(HDEHP) ₂ ≠ UO ₂ (HDEHP.DEHP) ₂ + 2H ⁺
Stripping	Acidic stripping $(R_3NH)_4UO_2(SO_4)_3 + 4HX \rightleftarrows 4R_3NHX + UO_2^{2+} + 3HSO_4^{-} + H^+$ $HX = HCl \text{ or } HNO_3$ Neutral stripping $(R_3NH)_4UO_2(SO_4)_3 + (NH_4)_2SO_4 + 4NH_3 \rightleftarrows 4R_3N + UO_2(SO_4)_3^{4-} + 6NH_4^+ + SO_4^{2-}$ Alkaline stripping $(R_3NH)_4UO_2(SO_4)_3 + 7Na_2CO_3 \rightleftarrows 4R_3N + UO_2(CO_3)_3^{4-} + 4HCO_3^{-} + 3SO_4^{2-} + 14Na^+$	Alkaline stripping UO ₂ (HDEHP.DEHP) ₂ + 4Na ₂ CO ₃ ₹ UO ₂ (CO ₃) ₃ ⁴⁻ + 4NaDEHP + H ₂ O + CO ₂ † + 4Na ⁺ (TBP must be added to avoid the third phase formation owing to low organic phase solubility of NaDEHP.) Acidic stripping UO ₂ (HDEHP.DEHP) ₂ + 2H ⁺ ₹ 2(HDEHP) ₂ + UO ₂ ²⁺

The chemistry of aqueous uranium is discussed in §16.3 together with the chemistry of the other actinides. Production of reactor fuel and reprocessing is described in Chapter 21.

5.5.4. Production wastes

The milling operation produces tailings consisting of fine-coarse particles in a water slurry containing most of the radioactive uranium decay products, of which radium is the most hazardous. As most tailings are not highly radioactive (e.g. from mining of low grade ores) they are dumped on outside the plant; leach water from the dumps will then, combined with mine water, enter local streams. In dry areas, dusts from the tailings may spread by winds.

This is the main picture and explains the high dose commitment values in Table 22.1. However, many health authorities now require conditioning of the tailings to reduce harmful effects to the environment: recycling of waste water, precipitation and removal of radium from solution, and neutralization to precipitate heavy metals. The slurry is usually transported to an impoundment basin where the solid particles settle out, and the effluent is treated for removal of activities before discharge into a settling pond. Treatment ponds may contain low permeability liners where appropriate to control seepage. Barium chloride may be added to the tailings to precipitate dissolved ²²⁶Ra as Ra-Ba sulfate. Lime and limestone may be added to the tailings to raise pH.

The final dry waste is stored either on surface or in shallow basins, though underground storage also has been practiced (abandoned mines). In the former case, the tailings are covered by up to 3 meters of earth fill to restrict erosion, and some water tight material to protect it against rain. In some cases (e.g. Sweden) the land on top of mine waste and tailings have been reclaimed for farming.

5.6. Radium and radon in the environment

In uranium ore radioactive equilibrium is established between the mother ²³⁸U and daughters in the decay chain (see heavy arrow in Fig. 5.1, 4n + 2 series) at the rate of the daughter half-life and at the level of the mother decay rate. The chain passes ²²⁶Ra and ²²²Rn and daughters down to ²⁰⁶Pb at a rate corresponding to the original amount of ²³⁸U, as long as the host material is undisturbed. The equilibrium between ²²⁶Ra and ²¹⁰Pb is established within some weeks. Since ²²²Rn is a gaseous intermediate, its daughters are likely to be formed in air ("radon daughters"). Radon diffuses out of thorium and uranium minerals, and adds radioactivity to the ground water and to the atmosphere both by its own presence and that of its daughters. Since Ra and Rn are among the most radio-toxic substances existing, causing bone and lung cancer at relatively low concentrations (the risk levels are discussed in Ch. 18), special attention must be devoted to their appearance in nature.

Common Rn-concentrations in ground water are 5 - 300 kBq/m³, but in areas of U-rich granite values ≥ 1 MBq ²²²Rn/m³ occur. Tap water usually contains of the order of 1 kBq/m³. In many places water from hot mineral wells is considered beneficial to health both for bathing and for drinking ("spas" or hot springs). The water may be warm due to radiogenic heating at the source (minerals rich in U or Th) and have a high content of dissolved radium and radon. Thus, in famous spas in Europe the ²²²Rn concentration for "therapeutic inhalation" may be 1 MBq/m³ air (Baden-Baden). From the Joachimsthal U-mine in Bohemia, which contains a number of hot wells (29°C), water containing 10 - 15 MBq ²²²Rn/m³ is pumped to spas, where it is used as medical treatment against rheumatism (a 30 min bath a day).

The average exhalation rate of radon from the ground is $5 - 50 \text{ mBq/m}^2\text{s}$, leading to a near ground level radon concentration of $1 - 10 \text{ Bq/m}^3$, but varies widely with ground conditions. In soil over Swedish uranium bearing shale ($\approx 300 \text{ ppm U}$) the ^{222}Rn concentration can exceed 1 MBq/m^3 , though on the average the concentration in Scandinavian air (the main geology consists of granitic rocks) is only 3 Bq/m^3 . Representative values are for the US $0.1 - 10 \text{ Bq/m}^3$, UK and Germany ~ 3 , and 10 (average) for France. The concentration in air above ground depends also on temperature and wind conditions.

Many ores contain small amounts of uranium. During processing, uranium and/or its daughters may enter the product, causing a radioactive contamination problem. For example, when apatite is used to produce phosphoric acid, the gypsum by-product contains all the radium originally present, producing a γ -ray and inhalation hazard from Rn-daughters, making it unsuitable for building material.

Radon concentrations in indoor air may be quite high, depending on site and building material. The 226 Ra content e.g. in German building materials varies from > 500 (gypsum) to 60 (brick) Bq/kg; slag used in Poland contains < 800, Italian tuff \approx 280, concrete in Hungary \approx 13, and white bricks in the UK only \approx 4 Bq/kg. The indoor concentration of radon also depends on the way in which the house is built and used (poorly ventilated, etc). In the US it varies between < 1 to > 1000 Bq/m³. Authorities no longer recommend very tight houses, as suggested in the 1970's to reduce heating costs. At levels < 70 Bq/m³ the Rn-hazard is considered negligible. In Sweden (pop. 8 M people) 50% of the houses have

70 - 200 Bq/m³, while 40 000 houses have been classified as "Rn-houses", i.e. their indoor concentration of Rn plus daughters exceed 400 Bq/m³. If the ground is the main source of radon, ventilation of the basement may be sufficient to declassify a "Rn-house".

Even coal contains small amounts of uranium, 4 to 300 kBq/ton, a typical value being 20 kBq/ton. When coal is burnt, the more volatile U-daughter products are released into the atmosphere, while the less volatile ones are deposited in the dust filters. A 1 GW_e coal-fired power plant typically releases 60 GBq ²²²Rn and 5 GBq ²¹⁰Pb+Po annually, while ~3 MBq daughter products are obtained per ton of fly ash.

5.7. Disequilibrium

Uranium ores at surface level are usually identified from the penetrating γ -ray emissions of the daughter products. However, it has been observed that some peat, which absorb uranium from local drainage, contain very little of the daughter products because they have been formed so recently that radioactive equilibrium has not been established.

The decay products of uranium passes over 10 elements (Fig. 5.1), all with very different chemical properties. These elements are transported by groundwater (*migrate*), the solute composition of which varies with the surrounding rock/soil minerals. The different elements migrate at different rates due to their different chemistries, dissolving in some areas and precipitating in others. If the mother and daughter in a radioactive chain migrate at different velocities during a time which is short in relation to the daughter half-life, the radioactive equilibrium is disturbed; this is referred to as *disequilibrium*. Such disequilibrium can be used to measure the age of the sample.

Let us consider the essential steps of the uranium decay series:

In this decay series the nuclide pairs which are suitable for determination of ages are associated with the time periods given between the double arrows of the two connected isotopes. For example, the 238 U-decay to 234 U passes over the short-lived intermediates 234 Th ($t_{1/2}$ 24.1 d) and 234 Pa ($t_{1/2}$ 1.17 m). The Th-isotope is long-lived enough to follow its own chemistry in a dynamic system. In strongly acidic solutions it forms Th^{4+} ions, while U forms UO_2^{2+} ions; the behavior of these two ions with regard to complex formation (e.g. by carbonates, hydroxyl or humic acid) and solubility is drastically different in neutral waters, leading to different migration rates for the two elements. For example, 238 U may migrate away, while 234 Th is precipitated or sorbed. As a result, when 234 Th decays (via the rapid equilibrium with 234 Pa) to 234 U, the latter is free from 238 U. From the deviation from the original activity ratio of 238 U/ 234 U = 1.0, (238 U refers to the concentration of that isotope, see §2.3.2) the time since 238 U and 234 Th separated (i.e. the age of the sample) is deduced.

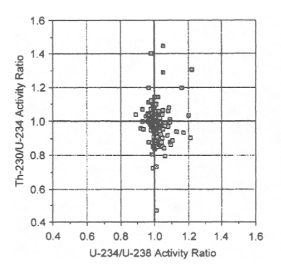
An example of such a system is the sedimentary uranium deposit in Tono, Japan, studied by Nohara et al. The ground water moves through the area with a velocity of 0.001 to 1

m/y. The specific activity ratios, measured by α - and $\beta\gamma$ -spectrometry, of $^{234}U/^{238}U$ is plotted against $^{230}Th/^{234}U$ (a), and $^{226}Ra/^{230}Th$ against $^{210}Pb/^{226}Ra$ (b) for a large number of rock samples in Figure 5.3. If radioactive equilibrium existed all ratios should be 1.0. The observation of deviations from 0.5 to 1.5 in (a) and up to 5 in (b) indicate that U, Th, Ra and Pb have migrated at different velocities in recent times. A detailed analysis yields the age of the U deposit and the migration rates of the daughter elements: the U and Th has not migrated during the past several hundred thousand years (this is probably also the age of the U deposit); Ra has migrated a few meters in the last 10000 y.

5.8. Age determination from radioactive decay

Prior to the discovery of radioactivity, geologists could obtain only poor estimates of the time scale of the evolution of the earth. The oldest geologic materials were assumed to be some 10 million years old, and it was believed that this represented the age of the earth. However, with the discovery of radioactivity early in this century, geologists developed more objective methods for such age determination ("nuclear clocks"). In 1907 B. B. Boltwood obtained a value of 2.2×10^9 y for the age uranium and thorium minerals, assuming that all U and Th ultimately decayed to lead. Considering how few isotopes in the chains had been discovered at the time, the calculation was surprisingly good. Nuclear clocks have provided primary data on the age and evolution of the earth (nuclear geochronology) as well as the formation of elements and of the universe (cosmochronology).

The cosmogenic radionuclides with relatively short half-lives can be used to date materials of more recent origin; e.g., ³H for water movements in the geosphere, and ¹⁴C for organic material of archaeologic interest. Practically all of the primordial radionuclides (Table 5.2) can be used for dating geologic materials: ⁴⁰K/⁴⁰Ar for igneous (plutonic) rocks (i.e. rocks which have solidified from a rather homogenous melt); ⁸⁷Rb/⁸⁷Sr for metamorphic and sedimentary rocks; ¹⁴⁷Sm/¹⁴³Nd for rock-forming silicate, phosphate and carbonate minerals; ¹⁸⁷Re/¹⁸⁷Os for sulfides and metallic material like iron meteorites, etc.



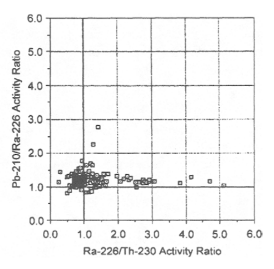


FIG. 5.3. Activity ratios ²³⁴U/²³⁸U vs ²³⁰Th/²³⁴U, and ²²⁶Ra/²³⁰Th vs ²¹⁰Pb/²²⁶Ra for rock samples from the Tono uranium deposit. (From Nohara et al.)

5.8.1. Dating by the ¹⁴C method

It is reasonable to assume that the production of ¹⁴C in the atmosphere has been constant for at least a million years, which means that equilibrium exists between the rates of formation and decay of the ¹⁴C in the atmosphere. Moreover, the half-life of ¹⁴C is sufficient to allow equilibrium between the ¹⁴C in the atmosphere, the oceans (including precipitations to ocean bottoms), and exchangeable carbon in natural materials. Thus from measurement of the specific radioactivity of carbon, it should be possible to determine when the sample became isolated from its natural environmental compartment.

The discovery that all living organic material has a certain specific radioactivity due to ¹⁴C led W. Libby to a new method for determination of the age of biological material. This method, which has been of great importance in dating archeological, geological, etc materials, is based on the assumption (i) that cosmogenic ¹⁴C has been produced at a constant rate, (ii) that the amount of anthropologic ¹⁴C is negligible compared to the cosmogenic, (iii) that after the organism incorporating the biological material died, no exchange occurs between the carbon atoms of the material and those of the surroundings. In such material the number of ¹⁴C atoms decreases with time according to the half-life of ¹⁴C. The equation is (for notation, see § 4.13)

$$^{14}C \text{ (Bq/g)} = ^{14}C_0 e^{-0.693t/5568}$$
 (5.3a)

or

$$t(y) = (\log^{14}C_0 - \log^{14}C) 5568/0.301$$
 (5.3b)

where $^{14}C_0$ is the initial 14 C activity of a standard ($^{14}C_0 \approx 14$ dpm/g, c.f. § 5.1.3) at time of death of the plant, etc (t=0). The reference time for 14 C ages is AD 1950, which is indicated with the letters bp or BP (for "before present"). The half-life of 5568 y is a standard reference value introduced by Libby. For example, if the specific activity of a sample is measured to be 0.1 dpm/g, then (5.3b) gives a value of 39 700 years as the time since the material ceased to exchange its carbon. Only with extreme care and very sophisticated equipment can a specimen this old be determined with reliability, but shorter times can be measured more accurately since the specific activities are larger. 14 C ages are used for dating specimen of 300 to 50 000 y, with an uncertainty of 10 - 100 y. 14 C-determinations were originally made by transferring the carbon into carbon dioxide, which was measured in an internal GM-counter (Ch. 8). Later, transformation into methane was preferred, and the CH₄ counted in an internal proportional counter. The most sensitive technique today is to introduce the sample into the ion source of a tandem-van-der-Graaff accelerator and "count" the amount of 14 C-ions relative to 12 C-ions by mass-spectrometry.

There are many cautions that must be observed in the use of carbon dating. In addition to those mentioned above, there is the possibility of isotopic effects in metabolic processes. These could cause 14 C to be slightly depleted relative to 12 C due to chemical reactions in the biological material. To take this into account to get the correct "solar age" instead of a slightly misleading " 14 C age", the isotopic depletion can be determined from deviations in the $^{12}C/^{13}C$ ratio. A " $\delta^{13}C$ correction" is introduced according to

$$^{14}C_{\rm corr} = ^{14}C \left\{ 1 - 2(\delta^{13}C + 25)/1000 \right\}$$
 (5.4)

The value of $\delta^{13}C$ differs for various substances; e.g. -35 to -20 for terrestrial organic matter, -8 to -7 for atmospheric CO₂, etc. Figure 5.4 illustrates the use of various $\delta^{13}C$ values for biological matter. These corrections lead to considerable adjustments of conventional ¹⁴C ages; e.g. for the period 7 000 to 2 000 years ago, corrections increase ages by up to 1 000 y.

Another necessary correction has to be made for changes in the cosmic ray irradiation, which, a priori was assumed to be constant, but is not over long geological periods. By counting the number of annular rings on old trees such as the Sequoia gigantea, which can be almost 4000 years old, and determining the ¹⁴C content of each ring, the variation of cosmic radiation through thousands of years can be studied accurately, see Figure 5.5. Obviously there is a periodic cycle of some 10000 y, which coincides with the variation in warm periods and small ice ages.

A large number of important age determinations have been made with the ¹⁴C dating method. It was believed that a large ice cap had covered parts of the North American continent until about 35 000 years ago. Dating of wood and peat by the ¹⁴C method has shown that the ice must have lasted until about 11 000 years ago. Moreover, several hundred pairs of sandals found in a cave in Oregon have been shown to be about 9 000 years old, indicating that tribes with significant cultures had developed soon after the withdrawal of the ice cap from North America. By analyzing inorganic carbon in bone apatite, it has been determined that hunters in Arizona killed mammoths there 11 300 years ago. Another example of the use of ¹⁴C dating which has attracted widespread attention involves the Dead Sea scrolls. There was considerable controversy about their authenticity until ¹⁴C dating showed their age to be slightly more than 1 900 years.

Some researchers claim that 14 C-ages lag behind with as much as 3 500 years for times close to 20 000 y, when compared to U-Th dating (§5.8.4). Such adjustments play a large role in the debate about cultural migrations. It is interesting to note that adjusted 14 C-dates now have led to the conclusion that e.g. the stone monuments in Carnac, Brittany, are believed to be $> 6\,000$ years old, i.e. older than the cultures in Egypt and Babylon. Similarly, grave mounds near the ancient edge of the inland ice in southern Norway and in Newfoundland have been found to be $> 7\,000$ years old.

5.8.2. Dating by K-Ar method

Potassium is the eighth most abundant element in the earth's crust and occurs in many important rock-forming minerals. The radioactive isotope ⁴⁰K is present only to 0.0117% in natural potassium. It has a branched decay as follows:

EC, 10.7%

----->
40
Ar (99.6% of natural Ar)

 40 K(t_{1/2} 1.28 × 10⁹ y) \langle

-----> 40 Ca (96.8% of natural Ca)

 β , 89.3%

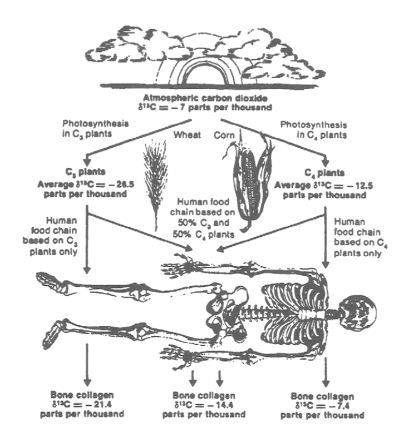


FIG. 5.4. Ratio of 13 C/ 12 C in human bone depends on diet and affects the δ^{13} C correction factor. (From American Scientist, 70 (1982) 602.)

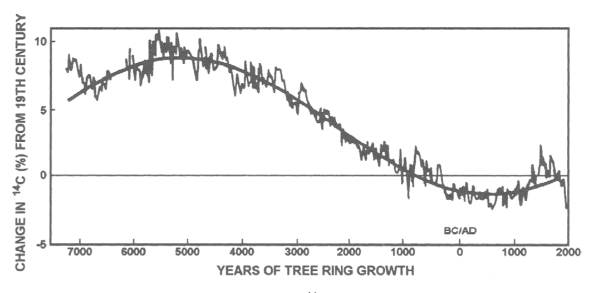


FIG. 5.5. Long term variation in atmospheric ¹⁴C activity as determined from tree rings. (From H. E. Suess, La Jolla Radiocarbon Lab.)

 $\lambda_{EC}=0.578\times 10^{10}~{\rm y}^{-1},~\lambda_{\beta}=4.837\times 10^{-10}~{\rm y}^{-1},~E_{\rm max}~1.32$ MeV. Because the $^{40}{\rm K}$ half-life is of the same magnitude as the age of the earth, measurement of the $^{40}{\rm K}/{\rm Ar}$ ratio can be used to determine ages of the oldest K-containing minerals (for notation, see §2.3.2). From the decay scheme one can derive the equation

$$t = \lambda^{-1} \ln[\{{}^{40}Ar/(0.107 {}^{40}K)\} + 1]$$
 (5.6)

where λ is the total decay constant (= λ_{EC} + λ_{β} ; see §4.12), ^{40}Ar represents the concentration of ^{40}Ar -atoms in the sample (i.e. number of radiogenic atoms of ^{40}Ar per unit sample weight), and ^{40}K is the present atomic abundance of ^{40}K -atoms. This equation assumes that all ^{40}Ar in the mineral is radiogenic. Any amount of non-radiogenic ^{40}Ar present initially (e.g. dissolved in the magma from which the mineral formed) is denoted as $^{40}Ar_i$. Thus the present amount of ^{40}Ar is the sum of the radiogenic amount and that originally present:

$$^{40}Ar = ^{40}Ar_{i} + \lambda_{EC} \lambda^{-1} ^{40}K (e^{\lambda t} - 1)$$
 (5.7)

The problem is that both t and $^{40}Ar_i$ are unknown. This can be solved by dividing equation (5.7) by the number of ^{36}Ar atoms in the sample, yielding

$$^{40}Ar/^{36}Ar = (^{40}Ar/^{36}Ar)_{i} + (\lambda_{EC}/\lambda)(^{40}K/^{36}Ar)\{e^{\lambda t} - 1\}$$
 (5.8)

Because 36 Ar is a stable non-radiogenic isotope, its amount in the sample should not change by time. Thus the measured $^{40}Ar/^{36}Ar$ ratio is the sum of the original and the radiogenic contributions from 40 Ar. Mineral samples of the same origin (so-called cogenetic) should have the same initial $^{40}Ar/^{36}Ar$ ratio and be of the same age (same t), though the $^{40}K/^{36}Ar$ may vary. Thus for cogenetic samples a plot of $^{40}Ar/^{36}Ar$ versus $^{40}K/^{36}Ar$ should yield a straight line, from the slope of which t is calculated. This line is referred to as an isochron. From the isochron for the rock sample from Tanzania in Figure 5.6, its age is calculated to be $(2.04 \pm 0.02) \times 10^6$ y, which is an interesting result as remains of early humans has been found in the same tuff matrix.

5.8.3. Dating by Rb-Sr method

Some uncertainty is associated with the use of the $^{40}K/^{40}Ar$ ratio because of the possibility of the loss of gaseous argon from minerals. An alternative method is based on 87 Rb/Sr system:

The measurement of the ${}^{87}Rb/{}^{87}Sr$ ratio by isotope dilution and mass spectrometry is one of the most reliable methods for geologic age determinations. Meteorite values as high as 4.7×10^9 y have been obtained, indicating that this is the age of the solar system. For the

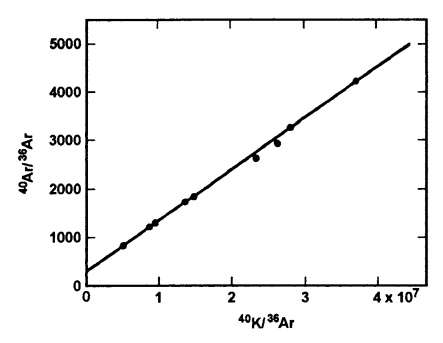


FIG. 5.6. K-Ar isochron of tuff from Olduvai Gorge, Tanzania, yielding a slope corresponding to an age of 2.04 ± 0.02 My. (From Faure.)

Rb-Sr clock a relation of type (5.9) is valid although corrections must be made for any non-radiogenic 87 Sr present; it is believed that the primordial 87 Sr/ 86 Sr ratio was 0.70.

5.8.4. Dating based on ²³⁸U decay

In the uranium decay series 8 α -particles are emitted in the decay from 238 U to 206 Pb. Thus for every 8 helium atoms found in a uranium mineral, one atom of 238 U must have decayed to 206 Pb. Designating the number of original uranium atoms in the sample at time 0 as $^{238}U_{\rm o}$, the number which has decayed with time t would be $^{238}U_{\rm o} - ^{238}U$, where the last number refers to the uranium atoms present now. Then $^{238}U_{\rm o} - ^{238}U = He/8$, where He is the number of helium atoms made. From this it relatively easy to derive an equation from which t, i.e. the age of the mineral, can be calculated once the values of ^{238}U and He are known. For accurate calculations, it is necessary to correct for the formation of helium from the decay of the 235 U and 232 Th also present in the mineral (see Fig. 5.1). Further, if the mineral has lost any helium through diffusion or other processes during its existence, the helium content would be abnormally low, leading to erroneously small values of t. This method, therefore, can only give lower limits of the ages of minerals.

Another common method of dating U-minerals is by considering its content of lead isotopes. Lead has four stable isotopes of which three are end products of radioactive decay series. The fourth lead isotope, ²⁰⁴Pb, is found in lead minerals in about 1.4% isotopic abundance and has no radio-genetic origin. At the time of formation of the earth, all the ²⁰⁴Pb in nature must have been mixed with unknown amounts of the other lead isotopes. If a lead-containing mineral lacks ²⁰⁴Pb, it can be assumed that presence of the other lead isotopes together with uranium and/or thorium must be due to their formation in the decay

series. In such ²⁰⁴Pb-free minerals if it is possible to determine the amount of the parent nuclide ²³⁸U and of the end product ²⁰⁶Pb, the age of the mineral can be obtained from the general equation

$$t = \lambda^{-1} \ln(1 + N_{\rm d}/N_{\rm p})$$
 (5.10)

where $N_{\rm p}$ is the number of parent atoms (e.g. 238 U) and $N_{\rm d}$ the number of radiogenic daughter atoms (e.g. 206 Pb), and λ is the decay constant of the parent. The atomic ratios between a number of isotopic pairs as a function of time is shown in Figure 5.7.

This method is more reliable than the helium method since there is very little possibility of any of the lead that has been formed by radioactive decay having diffused or been leached from the mineral during its geologic age. Mineral samples from the earth have yielded values as great as 3×10^9 y by the lead content method. Unfortunately, there is a problem with this method also. The decay series all pass through isotopes of the inert gas radon, and if some of the radon is lost from the mineral the 206 Pb content leads to an erroneously low age. However, if relation (5.13) is used for different decay series (i.e. the 238 U and the 235 U series) and the same t is obtained, the data are said to be concordant. Such data increases the confidence in the measured age.

Such data increases the confidence in the measured age.

Since the lifetimes of the uranium isotopes ²³⁸U and ²³⁵U are different, the isotopic ratio between their end products ²⁰⁶Pb and ²⁰⁷Pb can also be used for age determination. One can derive the relationship

$${}^{207}Pb/{}^{206}Pb = (1/138)(e^{\lambda_{235}t} - 1)/(e^{\lambda_{238}t} - 1)$$
 (5.11)

where the factor 1/138 is the present isotopic abundance ratio of the uranium isotopes. This method has given values of 2.6×10^9 y for uranium and thorium minerals. When applied to stony meteorites, a somewhat longer age of $(4.55 \pm 0.07) \times 10^9$ y is obtained.

When lead is extracted from the ore it is in secular equilibrium with its precursors radium and uranium. The radium and most of its descendants are removed during processing while the ²¹⁰Pb accompanies the other lead isotopes. The separation of radium from lead is not always complete. Because of the long half-life of ²²⁶Ra (1600 y) as compared to the short-lived ²¹⁰Pb (22.3 y), the determination of the excess ²¹⁰Pb over the equilibrium amount received from ²²⁶Ra decay provides a scale for the time since the manufacture of the lead. This was used to verify forgeries of paintings claimed to be made by the Dutch artist Vermeer van Delft (1632 - 1675). The paintings were in fact made by H van Meegeren (1889 - 1948), in the 1940's and were so excellent in style that every authority accepted them as authentic. When threatened by the death penalty for selling such national treasures to the Germans during World War II, van Meegeren confessed the forgery, and his story was verified by dating the lead in the "lead white", PbSO₄, in the fake "Vermeers".

5.9. Natural radioactivity of the oceans

The total amounts of 238 U and 232 Th in the ocean are 4.3×10^{12} kg (53 EBq) and 6.9×10^{10} kg (0.3 EBq), respectively. The oceans contain a much higher concentration of uranium than would be expected from its abundance in rocks. It also occurs at much larger

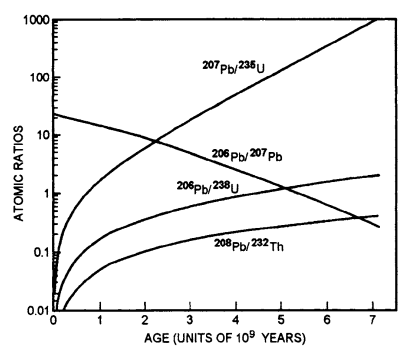


FIG. 5.7. Atomic ratios at the time of measurement as a function of the age of the minerals. (From E. K. Hyde.)

concentration than thorium, even though the latter element is more abundant in rocks. This is explained by the absence of easily soluble thorium complexes, in contrast to the strong $UO_2(CO_3)_3^{4-}$ complex. ²³⁸U decays to ²³⁴Th ($t_{1/2}$ 24.1 d), which precipitates out or absorbs to colloids before it decays to ²³⁴U, causing a slight disequilibrium between the two U-isotopes. This is used to determine the mean residence time of Th in the oceans: ~200 y; by contrast, the residence time of U in the ocean is 5×10^5 y. The decay passes the long-lived ²³⁰Th ($t_{1/2}$ 75 400 y), which deposits in sediments before radioactive equilibrium is reached with ²²⁶Ra ($t_{1/2}$ 1600 y); consequently the specific activity of ²²⁶Ra is less than expected from the U-content (39 Bq/m³), only 1 - 10 Bq/m³. It should be noted that the ²²²Rn concentration in surface waters is in disequilibrium with ²²⁶Ra because Rn is vented by heat, turbulence, etc.

As the total content of 40 K in the oceans is 74×10^{12} kg, corresponding to 19.4×10^3 EBq, 40 K is the largest source of radioactivity in the oceans. Minor activities come from 3 H as HTO and from 14 C as dissolved CO₂ or HCO $_3^-$.

5.10. Anthropogenic radioactivity in nature

In the analysis of a sample for its content of natural radioactivity it is necessart today to consider the possibility that the sample has become contaminated by "non-natural" radioactivities, i.e. radionuclides added to by human activities (so-called anthropogenic sources). Nuclear weapons tests, nuclear satellites burnt-up in the atmosphere, and nuclear power accidents may release large amounts of activities, see Table 5.4. The nuclear power industry is permitted by health authorities to continually release small, controlled amounts of specified radionuclides into the atmosphere and into open waters, Tables 19.7 and 21.10.

Effluents may also come from large radiotracer users (especially hospitals). In most cases these releases are well known and the area concerned ("contaminated") identified. From a global standpoint, compared to natural radioactivity, these releases are minor, but they do add to the our "natural" radiation background, as discussed in Chapters 18 and 22.

Usually one distinguishes between "near field" and "far field" effects of radioactivity releases. Near field effects are observed close to the release source, as for example the nuclear power plant or nuclear waste storage facility. The dissolution of nuclear waste by rain or ground water is a typical near field problem. As the source is known, it can be controlled and its environment monitored. If the radioactivity exceeds permitted levels, access to the contaminated area can be restricted. Far field effects involve the behavior of radionuclides which have spread out of such a restricted area, caused either by nuclear power accidents and weapons tests or by leakage from nuclear power plants.

In this paragraph we briefly describe some of the largest anthropogenic sources causing far field effects, i.e. nuclear weapons tests and nuclear power plant accidents. The cause of the releases is discussed in Chapter 19. Chapter 22 discusses both near and far field effects in further detail, particularly with regard to chemical properties: liquid releases from nuclear power plants, dissolution of solidified nuclear waste and of fall-out particles, migration in the environment, and possible consequences.

5.10.1. Nuclear weapons

Nuclear weapons were tested in the atmosphere up to 1990, with total releases up to 2×10^{20} Bq fission products, as well as some lesser amounts of Pu isotopes; cf. Table 22.2. Most of the debris injected into the troposphere had a mean residence time of ~ 30 d, causing fall-out mostly in the neighborhood of the test area. Some of the debris passed through the tropopause and entered the stratosphere where it was carried by winds around the globe at approximately the latitude of the release. The residence time in the stratosphere

TABLE 5.4. Events leading to large injections of radionuclides into the atmosphere

Source	Country	Time	Radioactivity Bq	Important nuclides
Hiroshima & Nagasaki	Japan	1945	4×10 ¹⁶	Fiss.prod. Actinides
Atmospheric weapons tests	USA USSR	-1963	2×10 ²⁰	Fiss. prod. Actinides
Windscale	UK	1957	1×10^{15}	¹³¹ I
Chelyabinsk (Kysthym)	USSR	1957	8×10 ¹⁶	Fiss. prod. ⁹⁰ Sr, ¹³⁷ Cs
Harrisburg	USA	1979	1×10 ¹²	Noble gases, ¹³¹ I
Chernobyl	USSR	1986	2×10 ¹⁸	¹³⁷ Cs

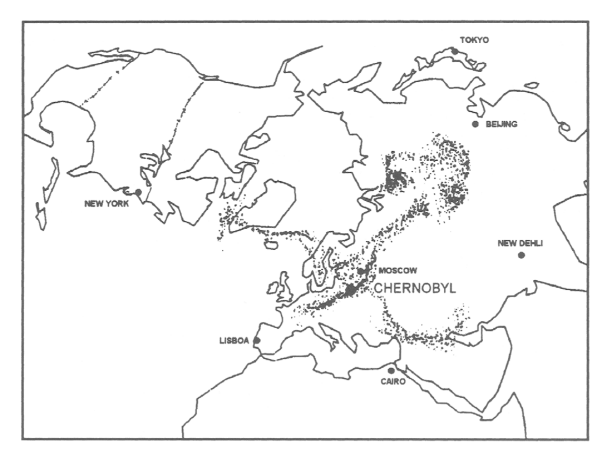


FIG. 5.8. The distribution of the airborne radioactivity from Chernobyl.

is 3 - 24 months depending on latitude (greatest at the equator). The average surface deposition is 2500 Bq/m² at 40 - 50°N, decreasing towards north an south (< 800 at > 70°N and at < 20°N). The global integrated deposition density for ²⁴¹Pu is ~440 Bq/m² and the air concentration is ~0.8 mBq/m³.

Because the atmospheric tests stopped in 1990, the fission products have decayed for > 10 y, leaving almost only 90 Sr, 137 Cs and Pu "active". Comparing with global natural radioactivity the contributions of these isotopes to our environment is negligible today, except for T and 14 C.

5.10.2. Nuclear power plant accidents

In 1957 a fire developed in one of the gas cooled graphite moderated rectors at Windscale, UK. The main radionuclides released were (in TBq) 131 I 700, 137 Cs 20, 89 Sr 3 and 90 Sr 0.3. The largest air concentration close to the plant was 20 kBq/m³. The Cs activity deposited was ≤ 4 kBq/m². The ground activity is now back to normal.

In 1979 a partial core melt-down occurred in a reactor at Three Mile Island power station (TMI), *Harrisburg*, Pennsylvania. Although the fission products released from the core were almost completely contained within the building, the Xe and Kr and some iodine (~ 1 TBq 131 I) were released. No deposition occurred on ground outside the building.

The explosion and fire in one of the power reactors at *Chernobyl*, Soviet Union, in 1986, was a more severe accident (§§20.1.2 and 22.3). For several days large amounts of fission products and actinides were ejected and spread over large areas of the former USSR and Europe, see Figure 5.8. Almost 20% of the fission products came down over Scandinavia, causing a deposition > 120 kBq 137 Cs/m² over the city Gävle in Sweden (170 km north of Stockholm). The plume spread down over Central Europe, causing very uneven deposition due to wind pattern and local rains: e.g., Munich, Germany, received \leq 25 kBq 137 Cs and 0.2 kBq 90 Sr per m², while Mainz (400 km away) received 180 Bq 137 Cs and \sim 0.001 Bq 90 Sr per m², see also §22.3.

In 1957 an explosion occurred in a nuclear waste storage facility (probably due to reactions between organic matter and nitrate) at Kyshtym south of Sverdlovsk, USSR, leading to the contamination of approximately 1600 km² of land by 8×10^{16} Bq fission products, causing local contaminations exceeding 10^{10} Bq/m² (2×10^{8} 90 Sr and 137 Cs). The area is still uninhabitable, see also §21.10.

5.10.3. Releases from nuclear power plants

All nuclear power plants are carefully controlled by the national radiation protection boards, and strict limits for releases are set. Usually it is quite easy for the power plants to meet these requirements. These releases are small compared to the natural radioactivities and mostly negligible in the far field, see §§20.4 and 21.8.

5.10.4. Other anthropogenic additions

In 1975 the London Dumping Convention set limits to dumping of nuclear waste in the Oceans to places far of shipping trades and fishing areas and at depths > 4000 m. Earlier dumping had been common practice by the nuclear powers even in narrow (the British Channel) and shallow (up to 50 m) waters to get rid of large volumes of low level long-lived waste. Officially reported dumpings were made as early as in 1946 and continued in the Pacific and the North Atlantic until 1982 and in the Barents and Kara seas even later. East of the British Islands the U.K. has dumped 665 TBq, France 134 TBq and other countries together some 35 GBq. In total some 45 PBq have been dumped at 46 different sites, mostly at depths of 1400 - 6500 m. Measurements reveal that in some cases radioactive nuclides are leaking out from the containers, causing bottom contamination at the site; however, the radionuclides are rapidly diluted as they are carried away by bottom currents.

Recently it was disclosed that the former USSR made huge dumps of nuclear waste in the waters east of Novaya Semlya: some 10 000 containers and 13 nuclear reactors (8 still containing fuel), mainly from nuclear submarines. In some places the waste is at very shallow waters (some 30 m). The total activity is estimated to > 60 PBq. The waste will ultimately leak out and spread with westerly currents towards the Barents Sea and the North Atlantic. However, the releases are expected to be rapidly diluted to harmless concentrations.

5.11. Exercises

- 5.1. Cosmic-ray irradiation of the atmosphere yields 0.036^{10} Be atoms cm⁻² s⁻¹. If this ¹⁰Be is rapidly carried down into sea water, which is assumed to have a volume of 1.4×10^{18} m³, what will the equilibrium radioactivity of ¹⁰Be in 1 m³ sea water be? The earth's surface area is 510×10^6 km².
- 5.2. In Greenland ice the ¹⁰Be radioactivity has been measured to be 0.0184 dpm m⁻³. How old is this ice if it was formed out of water in equilibrium with cosmic-ray ¹⁰Be (see previous question)?
- 5.3. The CO₂ in the atmosphere is in exchange with carbon in living organisms, humus, dissolved organic compounds, and carbonate in the sea water, the latter being the main reservoir (88%) of all exchangeable carbon. The amount of such exchangeable carbon is estimated to be 7.9 g cm⁻² of the earth. When cosmic-ray produced ¹⁴C is mixed into this exchangeable carbon, what will the specific activity become?
- 5.4. On the label of a bottle of cognac bought in 1976 it is stated that the cognac is over 20 years old. An analysis showed a tritium content of 80 TU. Discuss the trustworthiness of the statement.
- 5.5. 0.11 cm³ helium gas at NTP was isolated from 100 g of uranium mineral containing 5 ppm uranium. How old is the mineral?
- 5.6. A mineral was found to contain 39.1 g K and 87.2×10^{-6} liter Ar at NTP. How old is the mineral?
- 5.7. A uranium mineral was found to contain the lead isotopes ²⁰⁴Pb, ²⁰⁶Pb, and ²⁰⁷Pb in the ratio 1:1087:388, as determined with a mass spectrometer. Estimate the age of the mineral.
- 5.8. The heat flow from the earth's crust is 0.060 W m⁻². The mean thickness of the crust is 17 km and the earth's radius is 6371 km. The average concentration of uranium, thorium, and potassium in granite is estimated to be 4 ppm (by weight) 18 ppm, and 3.6%, respectively. Assuming that 7% by volume of the crust is made up of granite (feldspar + quartz, density 2.6 g cm⁻³), what will the heat flow at the earth's surface be from each of these elements? Assume β -heat as 1/3 E_{max} ; for α -decay assume $E_{\alpha} = Q_{\alpha}$. Each ⁴⁰K decay by the EC-branch emits one 1.46 MeV γ . Discuss the results.
- 5.9. A 1 GW_e nuclear power station uses annually about 30 t uranium enriched to 3% ²³⁵U. (a) How much natural uranium has been produced to keep it running? Assume waste stream from isotope separation plant to contain 0.3% ²³⁵U. (b) How much low grade ore (assume 0.06% uranium) must be mined, if the uranium recovery efficiency in the process is 70%?
- 5.10. The assumed uranium resources in Japan are 4 kt, in Argentina 12 kt and in France 48 kt U₃O₈. How many 1 GW_e reactor years can these uranium amounts sustain in each country at the uranium consumption rate (a) of the previous exercise?

5.12. Literature

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