Structural Studies on the Rare Earth Carboxylates

14. A Structural Study of the Orthorhombic Trishydroxyacetates of

Lanthanum(III) and Gadolinium(III)

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The structures of the orthorhombic tris-glycolates of lanthanum and gadolinium have been determined from three-dimensional X-ray intensity data, obtained by a linear single crystal diffractometer. The structures of both compounds have been described with a model using anisotropic thermal parameters on the heavy atom only. This model was refined to a final weighted R-value, equal to 0.043 and 0.035, in the lanthanum and gadolinium structures, respectively. A comparison of the corresponding interatomic bond distances indicates that the average metal—oxygen distances decrease approximately as the decrease in the ionic radius of the central ion. The change in size of the central ion also affects the oxygen—oxygen contact distances in the coordination polyhedron and the conformation of the ligands.

The metal-ligand bonds in lanthanoid(III) compounds involve little or no interaction between the metal $4\,f$ orbitals and the ligand orbitals. There are, as a result of this, no pronounced directional bonds of the type found among the d-transition elements in rare earth complexes. Phenomenologically, one may often regard the structures of lanthanoid compounds as a result of the compromise between the achievement of spherical symmetry for the central ion and the minimization of ligand—ligand repulsions in the coordination sphere. For polydentate ligands one has also to take into account the geometrical constraints imposed by the geometry of the ligand and its possibility to form hydrogen or metal bonded networks.

The difference in energy between various coordination geometries in compounds of high coordination numbers is often quite small. Hence, even small changes in any of the factors mentioned above are sufficient for the formation of a new phase. Examples of polymorphism are also frequent, e.g. the A, B, and C types of the rare earth sesquioxides, the orthorhombic 3 and monoclinic 4 rare earth tris-glycolates and the various rare earth dipicolinates, 5 described in previous parts of this series.

The present investigation has been made, mainly in order to investigate how a change in the size of the central ion within an isostructural series of compounds affects the coordination geometry and the conformation of the ligands. The main points of interest may be summarized as:

- a. How are the packing of the oxygen donors in the coordination sphere and the oxygen—oxygen contact distances changed by a change of size of the central ion?
- b. Are there any significant differences between the corresponding bond angles and distances within glycolate ligands in different crystallographic surroundings? How large is the variation in the conformation angle in the ligand? This angle is defined as the angle (the dihedral angle) between the projection of the C-O and the C-O(H) vectors viewed along the C-C vector. A discussion pertinent to these problems is found in part 8 of this series.
- c. Do possible conformation and packing changes among the ligands result in a change in the system of hydrogen bonds?

The compounds chosen for the study are the orthorhombic tris-glycolates of lanthanum and gadolinium, in the following abbreviated LAGLYC and GDGLYC, respectively. The crystal and molecular structure of the latter compound has been determined in part 3 of this series.³

The compounds studied contain approx. 41 % of a heavy atom, the ratios $Z^2_{\text{heavy}}/\sum Z^2_{\text{light}}$, being equal to 4 and 5, respectively. Hence, accurate determinations of the parameters for the light atoms are difficult. In order to obtain data of high accuracy, a single crystal diffractometer has been used in recording the various spectra for the X-ray structure determinations.

EXPERIMENTAL

Single crystal work. LAGLYC and GDGLYC were prepared as described before. The habits and sizes of the crystals used are given in Table I. Intensity data for the two compounds were collected at 25.0 \pm 0.2°C with an automatic linear single crystal diffractometer of type PAILRED. One LAGLYC crystal was used in recording a total of 1819 reflexions in the layers 0kl-11kl and h0l-h1l. Two GDGLYC crystals were used in recording the layers hk0-hk7 (1527 reflexions) and h0l-h10l (1441 reflexions), respectively. In both structures the region $\sin\theta \leq 0.80$ was investigated. Mo $K\alpha$ radiation was used. It was made monochromatic by reflexion in the (002) plane of a graphite crystal. The take-off angle was 6°. The pulse height discrimination levels were set for approximately a 95 % window, centered at the Mo $K\alpha$ peak. Coincidence losses were negligible, except for the four strongest reflexions in LAGLYC, where they amounted to at most 5%. The electronic stability was checked by measuring the intensity of four standard reflexions for each layer. Only random variations, usually less than 3%, were observed over a 24 h period. The counter aperture was 1.5° and the integrated intensities were determined by using the ω -scan technique. The scan speed was 1°/min and the scan ranges varied from 3° to 5°. The larger scan range was used in the vicinity of the ω -axis. Stationary background counts, B_1 and B_2 , were measured for 10 sec in LAGLYC and 20 sec in GDGLYC, at both ends of each scan interval.

The corrected integrated peak counts, I, were calculated from the eqn.

$$I = C - t_{\rm c} \frac{B_1 + B_2}{2 \ t_{\rm B}}$$

where C is the total integrated peak count obtained in a time $t_{\rm c}$ and $t_{\rm B}$ is the time for each of the background counts. The variances of the corrected intensities were estimated, using the expression

$$\sigma^2(I) = C + t_c^2(B_1 + B_2)/4t_B^2 + (0.025I)^2$$

Table 1. Unit cell parameters and volumes with their corresponding standard deviations for the various isostructural orthorhombic tris(hydroxyacetato) lanthanoid(III) compounds, $M(HOCH_2COO)_3$.

Central ion M	a/Å	b/Å	c/Å	V/ų
La	10.384(5)	11.564(4)	7.975(2)	957.5(4)
Ce	10.380(4)	11.541(3)	7.902(2)	946.7(4)
\mathbf{Pr}	10.401(6)	11.527(4)	7.905(2)	947.7(4)
Nd	10.362(5)	11.411(3)	7.772(1)	918.9(3)
Sm	10.306(6)	11.335(4)	7.724(2)	902.4(4)
Eu	10.303(4)	11.306(5)	7.698(4)	896.7(5)
Gd	10.298(2)	11.275(2)	7.649(1)	888.1(5)
$\mathbf{T}\mathbf{b}$	10.284(4)	11.247(4)	7.636(4)	883.2(5)

Table 2. Powder data for the orthorhombic rare earth tris-glycolates, $MC_6H_9O_9$. The observed values of $\sin^2\theta\times 10^4$ are denoted obs, while the corresponding quantities calculated from the least squares refined lattice parameters in Table 1 are denoted calc.

hk 1				Nd			0d	Intensit	-	
	obs calc	obs calc	obs calc	obs calc	Obs calc	obs calc	obs calc	QQ.	obs	calc
101	148.5 148.6	149.4 150.3	150.8 150.0	153.5 153.7 180.3 182.6	156.0 155.6	155.7 156.3	157.9 157.6	VW	157.6	158.1
0 2 0	176.9 177.8	178.2 178.5	180.3 178.9		185.6 185.0 203.1 201.8	185.9 186.0 203.3 202.8	187.4 187.0 204.8 204.4	n: VS	187.7	187.9 205.1
111	193.7 193.0 220.3 220.5	194.7 194.9 219.5 220.6	196.7 194.8 221.3 219.8	200.5 199.4 220.7 221.4	224.8 223.8	223.2 223.9	224.9 224.2	m+	223.7	224.7
210	264.6 264.9	263.7 265.3	265.3 264.5	268.6 267.0	270.2 270.1	269.1 270.4	271.9 270.9	m	271.8	271.7
5 5 0	398.5 398.2	399.8 399.1	401.6 398.7	405.3 404.0	409.5 408.8	409.5 409.9	410.8 411.2	w	413.1	412.7
221	498.3 [491.7	499.6 494.3	490.9 493.8	498.1 502.3	505.5 508.4 513.5 515.9	508.8 510.2 521.8 518.7	509.6 512.7 521.5 522.3	WW.	515.4 528.1	514.6 524.8
0 5 1	550.6 548.5	490.7	- 497.7	512.1 509.2 565.8 564.5	- 571.9	521.8 518.7 - 574.7	- 578.4	EH.	581,1	581.0
022	554.1 551.6	552.4 551.9 558.6 559.1	559.0 \(\begin{pmatrix} 559.6 \\ 559.3 \end{pmatrix}	565.8 564.5 - 576.1	586.3 583.4	588.6 587.1	592.5 593.3	8	596.6	595.6
301	590.9 589.5	591.9 591.6	reso e	600.0 596.5	602.6 603.1	606.3 604.2	605.3 605.9	m+	608.2	607.6
505	597.4 594.3	600.4 601.3	292. [1600.1	615.2 _615.0	623.5 622.2	626.7 625.1	630.0 630.4	m+	634.0	632.4
122	608.9 606.7	613.8 614.3	607.5 614.2	631.7 631.5	640.2 639.4	643.1 642.4	651.5 -644.9	m.	653.9	651.8
230	- 620.4 634.5 634.0	621.3 622.2 634.3 636.2	621.9 622.3 628.3 634.2	643.9 642.2	649.3 649.4	650.7 650.7	652.7	m-	055.9	654.6
311	640,6 638.7	645.6 645.9	639.8 644.9	659.3 660.6	670,4 668.5	672.4 671.6	676.5 677.2	m+	679.7	679.4
0 4 0	F211 i	F713 R	715.7	728,3 730.2	740.1	743.9	747.3 7748.0	m+	749.0	751.7
231	711.0 713.9	714.7 717.3	724.3 717.4	1750.0	1/39.1	L/42.0	[140.5			749.5
5 5 5	776.4 772.0	778.6 779.8	775.2 779.0	796.8 797.5	806.1 807.2 891.2 895.2	812.8 811.1 894.6 895.8	817.6 817.4 897.1 896.7	a m+	822.8 899.3	820.3
4 0 0 0 1 3	882.6 \[\frac{881.9}{865.5} \]	881.4 882.6 902.8 901.1	884.5 879.0 - 900.6	890.3 885.6	Folio 7	953.4 949.1	060.8	VVW	099.3	964.3
410	925.9 926.4	928.3 927.2	- 923.7	935.9 931.3	938.3 941.4	941.3 942.3	944.2 943.4	W	943.3	946.0
240	932.4 931.5	935.5 934.5	932.5 935.4	951.5 951.7	962.8 963.9	966.2 967.8	961.9 960.8	VVW	974.5	976.5
1 1 3	945.9 940.6	957.4 956.3	- 955.5	988.4 [986.5	993.1 1998.6	1010 1005	1018 1017	5 W		1020
401	976.0 975.4	975.9 977.8	971.6 974.1	1984.0	1019 1019	993.3 996.1 1023 1023	999.2 998.2 1028 1027	w	999.0 1025	1001
331 232	988.2 989.5 997.5 994.3	994.5 993.2 1002 1003	982.7 992.1 1002 1003	1010 1007 1027 1026	1042 1039	1025 1025	· - 1051	-	1025	1055
322	- 1048	1051 1056	- 1054	f 102k	1084 1087	1092 1091	1099 1098	vvw	1098	1101
420	- 1060	- 1061	1066 1058	1075 1068	1079 1080	1080 1082	1083 1084	VVW	1086	1087
042	1091 1085	1098 1094	- 1096	1124 1124	1142 1139	1147 1145	- 1154		1160	1159
051	1206 1204	1214 1211	- 1213	1239 1239	1256 1256 F1312	1260 1263	- 1270 [1326		1276	1277 J1333
15 1 33 2	1264 {1260	1271 {1266	- 1268	1295 1302	1314 1318	1318 1323	1326 1331	sr\	1333	1336
430	1284 1282	1284 1284	1283 1282	1296	L1312	1314	1318 1317	ra	1321	1322
412	1301 1300	1308 1308	- 1304	1330 1325	1345 1340	1344 1343	1352 1350	VS	1359	1354
242	[1305	1321 1315	1319 1316	1346 1345	1366 1362 1374 1380	1370 1369	1379 1378	m+ 6	1385	1384
25 0	1335 1332	1340 1336	1332 1338	- 1362 1382 1384	1374 1380 1404 1400	1381 1386 1410 1406	1393 1393 1419 1418	m-	1423	1399 1423
303	1359 1382	1354 1353 1397 1398	1349 1350 1392 1395	1382 1384 1430 1429	1452 1446	1457 1453	1465 1465	n+	1470	1470
4 2 2	- 1433	1444 1442	- 1438	1464 1462	- 1479	- 1483	- 1490		-	1495
501	1471 1471	1 hosk 1474	- 1469	1486 1482	1500 1498	1499 1500	1504 1503	71	1504	1507
233	- 1461	L1479	1480 _1478	1518	- 1537 1603 1594	1549 1545 - 1604	- 1559 1625 1625		1566	1565 1631
00 4 32 3	- 1495 1509 1515	- 1523 - 1531	1526 {1522	1568 1574	1588 1585	1595 1592	1606 1605	m+	1612	1611
114	- 1595	1600 1606	1610 1610	1645 1643	1663 1665	- 1707	1689 1683	10+	1728	1734
060	F1600	1630 1625	1629 1626	F1671	- 1692	1674 1674	T1718		-	1591
1 4 3	1607	1622	- 1621	1671 11675	- 1696	1702	1722 - 1728	m-	-	1725
432	1650 1656	1665 1665	1665 1662	1693 1690	1710 1710 1783 1779	1716 1715 - 1787	- 1724 - 1799		-	1730 1810
252 161	1701 1705 1746 1748	1719 1717 1759 1756	1726 1718 - 1760	1753 1756 1795 1797	- 1821	- 1830	1850 1841	W	1847	1850
2 4 3	1774 1773	1795 1791	- 1791	- 1837	- 1860	- 1870	- 1886		-	1894
224	- 1893	1914 1922	1920	1976 1978	2007 2002	- 2014	2034 2036	m+	-	2043
600	1979 1984	1980 1986	- 1978	- 1993	2058 2060	2025 2016	- 2018 2063 2064	m÷	2070	2023
610	2031 2029 2184 2189	2032 2030	2015 2022	2031 2038	2267 2268	2059 2062 2281 2278	- 2289	ure	2010	2070
36 1 26 2	2184 2189 2200 2194	2191 2198 2203 2207	2207 {2200	- 2240 - 2258	Engg.	- 2299	- 2313			J2324
503	2222 2219	2244 2236	2226 2229	- 2269	2288 1 2297	- 2302	- 2515		2323	2322
61 2	2406 2403	2418 2411	alim J2403	2423 2432	2458 2459	2463 2463	2467 2470	w	2482	2477
404	2372 2377	- 2405	L2401	2459 2460	2480 2489	2493 2500 - 2563	- 2522		-	2530 2604
105	2388 2391	- 2434	2432 2432	- 2515	2539 - 2546	- 2503	- 2595		-	

All intensities were corrected with the Lorentz, polarization, and absorption factors. The linear absorption coefficients and the range of the transmission factors are given in Table 3.

Powder work Powder photographs were taken at room temperature with a Guinier-Hägg camera, using $CuK\alpha$ radiation. Lead nitrate (cubic, $\alpha=7.8568$ Å) was used as an internal standard. The least-squares refined lattice parameters for the isomorphous orthorhombic $Mglyc_3$ phases (M=La-Tb) are given in Table 1 and the observed and calculated values of $\sin^2\theta$ in Table 2.

REFINEMENT OF THE STRUCTURES

LAGLYC and GDGLYC are isostructural. The space group is $Pn2_1a$ with the general fourfold positions $(x.y,z; \ \bar{x},\frac{1}{2}+y,\bar{z}; \ \frac{1}{2}-x,\frac{1}{2}+y,\frac{1}{2}+z; \ \frac{1}{2}+x,y,\frac{1}{2}-z)$. Preliminary values of the atomic parameters for the refinement of the 16 non-hydrogen atoms were obtained from Ref. 3. The atomic scattering factors used were taken from International Tables ⁸ (oxygen and carbon) and from Cromer et al. ⁷ (lanthanum and gadolinium). The function minimized was $\sum w(|F_0|-|F_c|)^2$ where the weights w were equal to $1/\sigma^2$ ($|F_0|$) for reflexions obeying the conditions $I>3\sigma(I)$ and $0.80 \le |F_0|/|F_c| \le 1.25$; other reflexions were

Table 3. Data on the crystals used for the structure determination of LAGLYC and GDGLYC and the course of the refinement of the two structures. The values of R and wR are defined as $R = \sum ||F_0| - |F_c||/\sum |F_0|$ and $wR = [\sum w(|F_0| - |F_c|)^2/\sum w|F_0|^2]^{1/2}$, respectively.

	LAGLYC a-axis	GDGLYC b-axis	GDGLYC c-axis	GDGLYC Both settings averaged
Habit of the crystal Approximate dimensions	Approx. spherical Radius 0.15 mm	Tabular (100) 0.09 × 0.09 ×	Tabular (100) 0.15×0.15×	-
1		$\times 0.03$ mm	0.035 mm ³	_
μ/cm^{-1}	45.7	82.3	82.3	
Range of transmission	0 55 0 61	0.00 0.55	0.17 - 0.50	
factors Total number of independent reflexions recorded for	0.57 - 0.61	0.33 - 0.55	0.17-0.50	-
$\sin \theta < 0.80$	1819	1441	1527	1785
No. of reflexions				
with $I \leq 3\sigma(I)$	239	316	$\bf 372$	570
No. of reflexions with $I \leq 0$ Isotropic refinement, in- dividual scale factors	35	106	118	89
R (all reflexions)	0.0463	0.0664	0.0610	_
ωR	0.0440	0.0518	0.0469	
No, of reflexions with $\omega \neq 0$ Refinement with aniso- tropic thermal parameters on the metal atom and an	1516	978	1035	_
over-all scale factor				0.0800
R_{p}	0.0460	0.0617	0.0573	0.0522
ωR	0.0432	0.0464	0.0402	0.0352
No. of reflexions with $\omega \neq 0$ Anisotropic refinement of all atoms	1512	982	977	1190
R	0.0408	_	-	
ωR	0.0383			_
No. of reflexions with $\omega \neq 0$	1530	_	-	

Table 4. Coordinates and thermal parameters for the various atoms in the orthorhombic LAGLYC and GDGLYC structures. The space group is $Pna2_1$ and the anisotropic thermal parameters for lanthanum and gadolinium have been calculated from the expression: $\exp[-(h^2\beta_{11} + hk\beta_{12} + \dots)]$.

Atom	Group	$x \times$	104	y >	< 10⁴	$z \times 1$	104	В	$^2/{ m \AA}^2$
		La	Gd	La.	Gd	La	Gd	La	Gd
O(1)	COO-	1997(6)	2030(11)	1364(6)	1244(11)	2750(8)	2554(17)	1.27(10)	1.81(22)
O(2)	COO-	2092(7)	2024(10)	2996(7)	2995(10)	1271(10)	1275(14)	1.94(12)	1.42(17)
O(3)	$-\mathrm{OH}$	4540(6)	4509(9)	1314(6)	1304(8)	3428(7)	3514(12)	1.48(10)	1.54(17)
C(1)	COO_	2722(10)	2622(16)	2183(9)	2218(14)	2083(12)	2152(25)	1.43(16)	0.91(29)
C(2)	\mathbf{coh}	4044(7)	4040(11)	2321(6)	2244(11)	2593(8)	2401(16)	0.91(9)	0.88(18)
O(4)	coo_{-}	4784(3)	4804(4)	72(8)	86(11)	6950(4)	6858(6)	1.21(6)	1.15(10)
O(5)	COO-	5599(4)	5672(5)	135(5)	-129(8)	9538(5)	9523(7)	1.54(7)	1.12(10)
O(6)	- OH	2412(4)	2453(5)	158(5)	149(8)	8107(5)	8156(7)	1.49(7)	1.37(10)
C(3)	coo-	4672(5)	4736(6)	56(11)	54(20)	8527(5)	8478(9)	1.27(7)	1.17(12)
C(4)	COH	3376(6)	3436(8)	244(5)	$-25\dot{5}(8)$	9277(8)	9360(12)	1.65(10)	1.41(15)
O(7)	COO-	2041(6)	1968(9)	-1273(6)	-1327(9)	2504(8)	2803(13)	1.17(9)	0.86(15)
O(8)	COO-	2180(6)	2180(11)	-2895(7)	-2959(12)	996(9)	1071(16)	1.65(10)	2.24(24)
O(9)	-oH	4487(6)	4454(7)	-1222(5)	-1182(7)	3148(7)	3205(11)	1.16(9)	0.67(12)
C(5)	COO-	2560(10)	2546(18)	-2194(9)	-2156(15)	2028(11)	2046(29)	1.27(15)	1.48(36)
C(6)	COH	4046(10)	4035(13)	-2239(9)	-2376(11)	2272(12)	2696(18)	2.39(17)	1.46(25)
<u>M</u>		2912(0.2)	2922(0.3)	0(0)	0(0)	4391(0.2)	4973(0.6)		
	β_{11}	< 10⁴	$\beta_{22} \times 10^4$	β_{33}	104	$\beta_{12}\!\times10^4$	β_1	$_3 \times 10^4$	$\beta_{23} \times 10^4$
La	16.	7(3)	13.3(2)	30.	1(3)	-3.5(1.2)	0.	2(3)	4.6(1.1)
Gd	12.	1(4)	14.4(3)	27.	L(4)	-1.0(1.0)	0.	5(3)	3.8(1.1)

given zero weight. The convergence was followed by the usual discrepancy indices R and wR, defined in Table 3, where details of the various models used in the refinements are also given. The refined atomic parameters with their estimated standard deviations are given in Table 4. Tables of observed and calculated structure factors may be obtained from the Department of Physical Chemistry at the University of Lund. A final difference synthesis, calculated by using the refined parameters given in Table 4, showed the presence

Table 5. Analysis of the weighting schemes for the LAGLYC and the GDGLYC (b-axis data) structures. The averages of $(|F_o| - |F_c|)^2 = w \Delta^2$ are normalized and the weights are estimated from the Poisson statistics of the counting data (p. 1481).

$ F_{ m o} $ interval LAGLYC	Number of reflexions	$\overline{w \Delta^2}$	$ F_{ m o} $ interval ${ m GDGLYC}$	Number of reflexions	$\overline{w \Delta^2}$
0.0 - 9.3	90	0.29	0.0 - 14.4	80	0.26
9.3 - 13.1	128	0.60	14.4 - 20.3	93	0.38
13.1 - 16.9	141	0.87	20.3 - 29.3	99	0.93
16.9 - 22.3	143	1.16	29.3 - 42.2	97	1.28
22.3 - 28.6	148	1.51	42.2 - 55.8	98	1.10
28.6 - 36.3	144	1.57	55.8 - 67.8	98	0.74
36.3 - 46.5	148	0.93	67.8 - 81.6	101	1.19
46.5 - 59.7	174	1.02	81.6 - 94.6	104	1.09
59.7 - 80.3	194	0.62	94.6 - 117.6	104	0.75
80.3 - 234.8	195	1.44	117.6 - 234.4	103	2.27

of a peak equal to $3e/Å^3$ at the central ion sites in both structures. The other parts of the electron density maps showed only small spurious peaks, at most $0.5~e/Å^3$, above a smooth background.

An examination of the values of $\overline{ud^2}$ between different $|F_0|$ -intervals (Table 5) indicates that the weighting procedure used is not entirely satisfactory. Hence, the estimated standard deviations given in Table 4 may be too small. The weight analysis indicates the occurrence of systematic errors in the measurements. In the Single Crystal Intensity Measurement Project Report 9,10 of the International Union of Crystallography it has been shown, that serious instrumental malfunctions may occur with linear diffractometers, causing a systematic error in the structure factors with increasing angle of inclination (see Ref. 9, p. 13). However, no error of this type seems to be present in our diffractometer, as judged by the fact, that intensity data, obtained from two different settings, only showed small random variations, of the order 2-3%, between the $|F_0|$ -values of the same reflexions.

The scale factors for the various layers were improved in the first stage of the refinement. The relative values thus obtained agreed within 2-3% with those calculated from data in the second setting. The covariance between the scale factors and the anisotropic thermal parameters is usually large. The anisotropic thermal parameters are also more sensitive towards systematic errors than the positional parameters. In order to investigate the effect of the first factor, two series of anisotropic refinements of all atoms and an overall scale factor were tried on the LAGLYC data. The relative scale-factors for the various layers, obtained from the isotropic refinement, were varied at random within 2-3% between the two refinements. Examination of the two sets of parameters showed that the coordinates for all atoms and the anisotropic thermal parameters for lanthanum, in general, had changed less than one standard deviation with the change of scale. On the other hand, the β_{ij} -values (especially β_{11}) of some of the light atoms had changed with 50%. Hence, it was concluded, that no physically meaningful model of the structure with anisotropic thermal parameters on the light atoms could be used.

A fairly large number of the reflexions recorded have zero weight (Table 3). Most of these have h+l=2n+1 and $I\leq 3\sigma(I)$. This is due to the value of z for the heavy atom, which is close to $\frac{1}{2}$. Hence, its contribution to the structure factor is small for the re-

flexions, obeying the above condition.

The experimental data were not corrected for extinction as there was no significant difference between the $|F_0|$ and $|F_c|$ values for strong reflexions at low values of sin θ .

Programmes used. All computing work was carried out on the UNIVAC 1108 Computer at Lund, Sweden, using the programmes DRF, DATAP2, LALS, DISTAN, PLANES, ORTEP, and CELSIUS.¹¹

DISCUSSION

Selected interatomic distances and angles for the LAGLYC and GDGLYC structures are given in Table 6. A stereoscopic view of the two structures is given in Fig. 1. Some symmetry-related sites have been given superscripts of the following significance

$$\begin{array}{lll} \text{(i)}\ \frac{1}{2}-x,-\frac{1}{2}+y,\ \frac{1}{2}+z & \text{(iv)}\ \frac{1}{2}+x,\ y,\ \frac{1}{2}-z \\ \text{(ii)}\ -\frac{1}{2}+x,\ y,\ \frac{1}{2}-z & \text{(v)}\ \frac{1}{2}+x,\ 1+y,\ \frac{1}{2}-z \\ \text{(iii)}\ \frac{1}{2}-x,\ \frac{1}{2}+y,\ \frac{1}{2}+z & \end{array}$$

where x, y, z are coordinates of the crystal-chemical unit, given in Table 4. A description of the GDGLYC structure, based on photographic intensity data, has been given in part 3.³ The parameters, obtained by photographic and diffractometer methods in general, agree within $3 \times \sqrt{\sigma_1^2 + \sigma_2^2}$, where σ_1 and σ_2 are the estimated standard deviations, obtained by the two methods. One exception is the x-coordinate of O(9). However, the diffractometer data give a bond length to gadolinium for this atom, which is in better accordance with the other Gd – O distances than was the case with the photographic data.

Table 6. Selected bond distances (in Å) and angles with their corresponding standard deviations in LAGLYC and GDGLYC.

	dev	lations in La	GLIC and GDGLIC.		
	La V	Vithin the co Gd	ordination polyhedra	La	Gd
$\begin{array}{c} M-O(1)\\ M-O(2^i)\\ M-O(2^i)\\ M-O(3)\\ M-O(4)\\ M-O(5^{ii})\\ M-O(6)\\ M-O(7)\\ M-O(8^{iii})\\ M-O(9)\\ O(1)-O(5^{iii})\\ O(5^{ii})-O(8^{iii})\\ O(2^i)-O(4)\\ O(2^i)-O(9) \end{array}$	2.533(7) 2.552(8) 2.569(7) 2.525(4) 2.444(4) 2.592(4) 2.594(7) 2.580(8) 2.587(6) 2.967(8) 2.858(9) 3.271(9) 3.139(11) 3.116(10)		$\begin{array}{lll} 2) & O(1) - O(9) \\ O(4) - O(8^{iii}) \\ O(3) - O(4) \\ O(3) - O(8^{iii}) \\ O(3) - O(8^{iii}) \\ O(3) - O(9) \\ O(0) - O(2^{i}) \\ O(0) - O(5^{ii}) \\ O(6) - O(5^{ii}) \\ O(6) - O(8^{iii}) \\ O(7) - O(1) \\ 8) & O(7) - O(2^{i}) \\ 5) & \angle O(1) - M - O(3) \\ \end{array}$	3.505(9) 3.966(9) 3.203(10) 3.164(8) 2.865(9) 2.941(9) 2.828(6) 2.843(9) 3.056(10) 3.248(10) 3.248(8) 63.8(2) 61.9(1)	3.234(13) 3.732(14) 3.061(16) 2.915(12) 2.744(15) 2.807(12) 2.868(14) 2.766(7) 2.686(16) 2.899(15) 2.949(15) 2.788(12) 64.8(3) 64.5(2)
O(4) - O(9)	3.395(7)	3.156(1	$1) \angle O(7) - M - O(9)$	60.0(2)	63.6(3)
		With	in ligand 1		
	La	Gd		La	Gd
$\begin{array}{c} C(1) - O(1) \\ C(1) - O(2) \\ C(1) - C(2) \\ C(1) - C(2) \\ C(2) - O(3) \\ O(1) - O(3) \end{array}$	1.322(12) 1.315(13) 1.441(13) 1.437(10) 2.696(10)	1.262(20) 1.291(20) 1.473(20) 1.440(15) 2.657(15)		115.3(9) 120.6(8) 122.3(9) 112.5(7)	118.9(15) 117.0(13) 122.6(14) 113.2(11)
		With	nin ligand 2		
	\mathbf{La}	Gd	g -	\mathbf{La}	Gd
$\begin{array}{c} C(3) - O(4) \\ C(3) - O(5) \\ C(3) - C(4) \\ C(4) - O(6) \\ O(4) - O(6) \end{array}$	1.263(5) 1.259(6) 1.489(8) 1.372(8) 2.632(6)	1.240(9) 1.269(9) 1.539(12) 1.441(11) 2.618(7)	$ \angle O(4) - C(3) - O(5) \angle O(4) - C(3) - C(4) \angle O(5) - C(3) - C(4) \angle O(6) - C(4) - C(3) $	124.5(5) 118.8(5) 115.0(5) 112.0(5)	126.2(7) 119.5(7) 110.5(8) 105.1(7)
		With	in ligand 3		
	La	Gd		La	Gd
C(5) - O(7) $C(5) - O(8)$ $C(5) - C(6)$ $C(6) - O(9)$ $O(7) - O(9)$	1.252(12) 1.222(12) 1.556(15) 1.443(12) 2.591(9)	1.249(21) 1.230(23) 1.630(23) 1.464(15) 2.584(12)		129.0(10) 114.7(9) 112.6(9) 110.3(8)	133.0(18) 114.9(15) 111.2(14) 102.7(10)
		Possible	hydrogen bonds		
	La	Gd	Ljurogon bonus	La	Gd
$O(3) - O(1^{iv})$	2.719(9)	2.723(15)	$\angle C(2) - O(3) - O(1^{iv})$ $\angle C(1^{iv}) - O(1^{iv}) - O(3)$	104.0(2) 114.3(6)	99.3(7) 111.0(10)
$O(6) - O(4^{ii})$	2.731(6)	2.729(7)	$\angle C(4) - O(6) - O(4^{ii})$	137.9(4)	134.2(5)
$O(9) - O(7^{v})$	2.704(9)	2.706(12)	$ \angle C(3^{ii}) - O(4^{ii}) - O(6) \angle C(6) - O(9) - O(7^{v}) \angle C(5^{v}) - O(7^{v}) - O(9) $	$\begin{array}{c} 96.3(3) \\ 112.5(6) \\ 101.6(6) \end{array}$	$\begin{array}{c} 93.5(4) \\ 98.7(7) \\ 111.7(11) \end{array}$
$O(6) - O(4^{ii})$	2.731(6)	2.729(7)	$\begin{array}{l} $	114.3(6) 137.9(4) 96.3(3) 112.5(6)	111.0(1 134.2(8 93.5(4 98.7(1

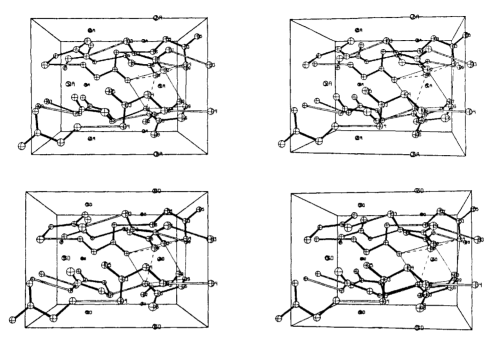


Fig. 1. Stereoscopic view of the LAGLYC and GDGLYC structures. The stick bonds between the ligand atoms are filled and those between possible hydrogen bonded atoms are unfilled. The picture has been drawn by using the program ORTEP and the atoms are represented by thermal spheres (or ellipsoids for the heavy atoms), formally scaled to include 50 % of the probability distribution.

The average metal-oxygen bond distances decrease from 2.55₂ Å in LAGLYC to 2.44. Å in GDGLYC. This change is approximately equal to the decrease in the ionic radius of the two central ions (0.12 Å). The average oxygen - oxygen contact distances in the coordination polyhedron also decrease with decreasing size of the central ion, the averages in the two compounds being 3.05, Å and 2.92, Å, respectively. The decrease is considerably larger for the distances above 3.1 Å than for those below, the averages being 0.22 Å and 0.03 Å, respectively. Some of the shorter oxygen – oxygen contact distances actually increase from LAGLYC to GDGLYC. This finding is a satisfactory piece of experimental evidence for the importance of oxygen - oxygen repulsions in deciding the geometry of solid rare earth complexes, a point, which has been discussed before by Albertsson 12 and by the present author. 13 The change in size of the central ion does not result in any significant changes in the distances between the two oxygen donors in the chelate ring (the ligand bite). The conformation of the ligand is instead changed in order to fit into the different crystallographic surroundings; a change wich is most noticeable in the conformation angle. The differences in conformation are most easily seen in Fig. 1 and in Table 7. The various conformations are a result both of differences between corresponding bond angles in the various ligands and of

Table 7. The deviation in Å of the central ion and ligand atoms from the least-squares planes, formed by the equatorial oxygens and the C-COO groups, respectively. The conformation angle in the various ligands has also been included in the last line of the table.

Atom	Equatori	al plane	Atom	Liga	nd I	Atom	Liga	nd 2	Atom	Liga	nd 3
· - · · · -	La	Gd		La	Gd		La	Gd		La	Gd
M	- 0.025	- 0.012	M	-0.412	- 0.826	M	- 0.078	0.215	M	- 0.554	0.117
O(3)	0	0	O(1)	-0.027	0.023	O(4)	0.027	0.044	O(7)	0.041	-0.020
O(7)	0	0	O(2)	-0.028	0.025	O(5)	0.027	0.039	O(8)	0.042	-0.020
O(9)	0	0	C(1)	0.082	-0.070	C(3)	-0.075	-0.113	C(5)	-0.110	0.053
` ,			C(2)	-0.027	0.021	C(4)	0.021	0.030	$C(\theta)$	0.027	-0.012
			O(3)	0.154	-0.178	O(6)	-0.083	-0.353	O(9)	0.187	0.603
Confo	rmation a	ngle O(1)	- C(1) -	-C(2)-O(3)	O(4)-	C(3) - C(4)	-O(6)	O(7)—	C(5) — C(6) -	-O(9)
				17.7	2.8		-13.6	29.0		5.1	30.9

"inversion" of the atoms in the least-squares plane, formed by the C-COO group.

Some of the corresponding interatomic bond distances in the ligands, mainly the C-C distances, are significantly different from one another as judged by the t-test. These differences might well be artefacts, brought about by the presence of systematic errors in the data, as mentioned on p. 1484. Most of the bond distances within the ligands show, on the whole, a satisfactory agreement between one another, i.e. a change in size of the central ion does not seem to result in any significant changes in the intra-ligand bond distances.

Hydrogen bonding and unit cell dimensions. The hydrogen bond scheme is outlined in Table 6 and in Fig. 1. There are only small differences between the various hydrogen bonded oxygen—oxygen distances in the two structures and the change in packing and conformation of the ligands do not result in any significant changes in the hydrogen bonds. One might even guess, that the preservation of an energetically favourable network of hydrogen bonds is as important for the conformation of the ligand as the size of the central ion (cf. part 8, p. 3358).

The symmetry related ligands are connected by hydrogen bonds as shown in Fig. 1. All ligands are aligned along the a axis, inclined approximately 30°, 90°, and 150° to the ab plane. A decrease in the metal—oxygen bond distances moves the ligands closer to the central ion and results in a larger decrease in b and c than in a for decreasing radius of the central ion (cf. Table 1). The smaller decrease in a is due to the alignment of the ligands along this axis in connection with the near constancy of the ligand dimensions and the hydrogen bonded oxygen—oxygen distances.

COMPARISON BETWEEN DIFFERENT GLYCOLATE STRUCTURES

The ligands in solid complexes may often be approximated as rigid bodies with some rotational freedom around the single bonds, i.e. the carbon – carbon bond in the glycolate ion. The experimental values of bond distances and an-

Table 8. Ranges and means of the various interatomic distances and angles within the glycolate ion, obtained by X-ray structure determinations of various solid glycolate compounds.

	C-C Bond in Å	C-O Bond in Å	C-O(H) Bond in Å	Ligand bite in Å	Sum of angles around the carboxylate carbon
Potassium ¹⁴ and					
rubidium 15 hydrogen	1.511 - 1.519	1.241 - 1.264	1.418 - 1.426	2.687 - 2.733	359.9
bisglycolates	1.51_{5}	1.24_{8}	1.42_{2}	2.71_{0}	359.9
Copper(II) glycolate ¹	- -	1.24 - 1.27		_ `	_
	1.53	1.25_{5}	1.43	2.57	359.9
Europium tris-		-			
glycolate ⁸	1.48 - 1.58	1.27 - 1.36	1.41 - 1.46	2.66 - 2.72	360
	1.52	1.32	1.43	2.69	360
Hydroxyacetato-oxy-	1.49 - 1.54	1.20 - 1.26		_	359 - 360
acetato-aquo-	1.51	1.24	1.44	2.54	360
erbium(III) hydrate ¹⁷					
Erbium tris-glycolate	1.48 - 1.54	1.24 - 1.32	1.36 - 1.50	2.54 - 2.57	359 - 360
dihydrate 18	1.52	1.27	1.44	2.56	360
Lanthanum and gado-	1.441 - 1.630	1.222 - 1.322	1.372 - 1.464	2.584 - 2.696	356.2 - 358.5
linium tris-glycolates	1.52_{1}	1.26_{\star}	1.43_{s}	2.63_{0}	358.3

gles, available in the literature on this ligand, are given in Table 8. These data indicate that the rigid body approximation is a fairly good model for the description of glycolate ions in crystallographic different surroundings.

Planar ligands are found in the Cu, K, Rb, and Eu (two ligands) compounds. The data from the alkali compounds, where the metal-ligand interactions are expected to be weaker than those in the other compounds, might indicate, that the planar form is the energetically favoured one for the "free" glycolate ion. The observed deviations from planarity are mainly caused by a twist of the hydroxy—oxygen from the C—COO plane, approximately 4° in ERGLYC¹⁷ and at most 20° in GDGLYC. The conformation differences among the C—COO groups in LAGLYC and GDGLYC have been mentioned previously.

The ligand bite in the rare earth glycolate complexes is significantly shorter in eight- than in nine-coordinated complexes, 2.54 Å and 2.65 Å, respectively. This difference may be due to a larger average attraction between the metal and the donor atoms in the former compounds. Another result of this increased attraction is a shortening of the average metal—oxygen bond distance in the eight-coordinated complexes.

The orthorhombic rare earth tris-glycolate phases are thermodynamically more stable than the corresponding monoclinic phases (at 25°C and 1 atm) for the elements La—Sm, while the opposite is true for Eu—Tb. It is tempting to speculate over the possible structural causes of these differences in relative stability. The average oxygen—oxygen contact distances are approximately the same in the two structures as exemplified by the monoclinic EUGLYC and the orthorhombic GDGLYC. Hence, differences in the oxygen—oxygen repulsions do not seem to be the deciding factor for the relative stability of

the two phases. This may instead be determined by differences in the planarity of the ligands and in the hydrogen bonded oxygen – oxygen distances between the two structure types, e.g., the average hydrogen bonded oxygen - oxygen distance is 0.10 Å shorter in GDGLYC than in EUGLYC.

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