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Hyperfine Structure in the 2084-keV State of Ce¹⁴⁰

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Differential perturbed angular-correlation measurements were made on the 2084-keV state of Ce¹⁴⁰ in aqueous solution at room temperature, and in lanthanum metal both at room temperature and at 77°K. The derived g factor of $\pm 1.014 \pm 0.038$ is compared with Rho's theory and with other experimental values. The advantages of working with metals and at low temperatures are discussed. An upper limit of approximately 200°K is set on the quantity ΔE , where $\Delta E/k$ is the energy spacing between Γ_7 and Γ_8 for Ce³⁺ in La metal. An approximate value of 4.1 ± 0.4 atomic units is derived for $\langle r^{-3} \rangle$ of the cerium 4f electron, using the above upper limit for ΔE .

I. INTRODUCTION

 $S_{\rm kev}^{\rm TUDIES}$ of the hyperfine interactions in the 2084-kev state of Ce¹⁴⁰ can yield important information about nuclear and atomic structure in this region of the periodic table. This state's mean life of 5 nsec¹ makes it inaccessible to all methods except perturbed angular correlations. In this paper we report the results of studies to date using this technique.

The nuclear properties of 5882Ce140 are especially interesting because they are accessible to theoretical treatment. The 82-neutron and 50-proton core can be neglected in calculating the properties of the low-lying states, and Bardeen-Cooper-Schrieffer (BCS) theory can be applied to the remaining 8 protons, which would just fill the $g_{7/2}$ subshell in a single-particle model. Rho has carried out such a BCS calculation.² Furthermore the calculated g factor of the 2084-keV state is quite different on the BCS theory from any of the singleparticle possibilities. The BCS wave function for this state involves a rich mixture of shell-model (quasiparticle) basis functions, and a determination of the gfactor constitutes a severe test of this theory.

The atomic properties of Ce are interesting because it has stable 3+ and 4+ oxidation states, with the configurations $4f^1$ and $4f^0$. Comparison of the effective fields for these two states affords the possibility of determining the crucial $\langle r^{-3} \rangle$ hyperfine-structure parameter for the 4f electron. We have obtained an approximate value for this parameter as well as an upper limit for the crystal-field splitting $\Delta E(\Gamma_7 - \Gamma_8)$ for Ce³⁺ in metallic lanthanum.

Sample preparation and descriptions of the measurements are given in Sec. II. In Sec. III the g factor is compared with theory. Crystal-field parameters are discussed in Sec. IV.

II. EXPERIMENTAL

Both the integral and the differential angular correlation methods were used to study the g factor of the 2084 keV state in Ce¹⁴⁰. These methods are described thoroughly in the literature.³ We give below only the particulars of our apparatus and the important details of our data-gathering procedures. For all the work reported here the radioactive parent was La¹⁴⁰, and the 329-487-keV cascade (Fig. 1) was used throughout.

A. Sample Preparation

Measurements were made on Ce140 in solution and in La metal. The La¹⁴⁰ was prepared by the reaction La¹³⁹ (n,γ) , using naturally abundant La (99.911%) La¹³⁹). Since only the trivalent state of La is stable in aqueous solution the liquid sources were prepared by simply dissolving irradiated La₂O₃ in 3M HNO₃. For the metallic sources, 20-30-mg pieces of La metal were irradiated in an argon atmosphere for 8 min at a flux of 10^{12} neutrons/cm² sec.

¹W. M. Currie, Nucl. Phys. 32, 574 (1962). ² Mannque Rho, Ph.D. thesis, Lawrence Radiation Laboratory Report, UCRL-11080, 1963 (unpublished report); Nucl. Phys. 65, 497 (1965).

³ R. M. Steffen and H. Frauenfelder, in Perturbed Angular Correlations, edited by E. Karlsson, E. Matthias, and K. Siegbahn (North-Holland Publishing Company, Amsterdam, 1964), p. 1.



B. Integral Angular Correlations

In this method a magnetic field is applied to the source perpendicular to the correlation plane and Larmor precession in the intermediate state is manifest as a phase shift in the correlation pattern. We used this method in early experiments, employing a fast-slow coincidence circuit with a resolving time of 18 nsec. Unfortunately the γ -ray spectrum of La¹⁴⁰ is quite complicated and there is a considerable amount of coincident radiation at the energies of the two peaks in question. These "background" coincidences, which arise largely from Compton-scattered 1597-keV photons, give an angular correlation pattern that is not rotated. In a preliminary report we gave a g factor of $+1.08\pm0.10$ for the 2084-keV state.⁴ Less than half of this probable error interval arose directly from counting statistics; the rest arose from uncertainties in the background correction. In a recent independent re-evaluation of the background correction we obtained a value of +0.99for the g factor. We are thus led to the conclusion that, even after a substantial experimental effort had been made (over 3×10^5 coincidences were collected), we still had measured the g factor to only about 10%. Longer counting periods would not materially help the situation, because the largest source of error was our inability to correct accurately for background. The apparatus was therefore modified to permit differential measurements.

C. Differential Angular Correlations

In these experiments the apparatus was similar to a conventional fast-slow coincidence circuit, except that a time-to-height converter operating on the overlap principle was substituted for the fast coincidence unit. The $1 \times 1\frac{1}{2}$ in. NaI(Tl) gamma-ray detectors were mounted on 30-cm quartz light guides which were in turn secured to Amperex 56 AVP photomultiplier tubes. Energy-sensitive pulses were taken from the fifth dy-

TABLE I. Half-life of the 2084-keV state of Ce¹⁴⁰.

Cascade studies	$t_{1/2}(nsec)$	Reference
329–487 keV	3.44(6)	1
329–487 keV	3.41(4)	5
392–487 keV	3.52(10)	this work

node and passed through a transistorized shaper and into a cosmic multiple coincidence unit which performed a slow coincidence. Time-sensitive pulses from the anode were limited and passed into the time-toheight converter. The pulses from this unit were sent into a pulse-height analyzer that was gated by the slow coincidence unit. The resultant display was an exponential decay curve with the half-life of the intermediate state. Such a curve is shown in Fig. 2. This curve gave a half-life of 3.52 ± 0.10 nsec for the 2084 keV state, in good agreement with the values in the literature (Table I).^{1,5}

An iron core "C" type electromagnet with a $\frac{1}{2}$ in. pole gap provided a maximum field of 42.0 kG on the sample. For low-temperature work pole tips with axial holes were used, allowing a Dewar to be inserted without destroying the axial symmetry. The maximum field with these pole tips was only 30 kG. The photomultipliers were shielded from the stray field by two concentric Mu-metal cylinders and a cold-rolled iron tube. The gain shifted less than 1% for maximum field. A time resolution of 5 nsec for the 329–487 keV settings was measured using annihilation radiation from Na²².

For differential g-factor measurements the counters were set at 135° to one another and coincidence time spectra were taken with the magnetic field directed alternately up and down relative to the correlation plane. The resultant time spectra, denoted by $W_+(t)$ and $W_-(t)$, are exponential curves modulated slightly by the Larmor precession of the intermediate state.



FIG. 2. Prompt resolution curve and decay curve for the 329-487 keV cascade in Ce¹⁴⁰.

⁶ H. J. Körner, E. Gerdau, C. Günther, K. Averbach, G. Mielken, G. Strube, and E. Bodenstedt, Z. Physik 173, 203 (1963).

⁴ R. M. Levy and D. A. Shirley, Phys. Letters **3**, 46 (1962), Erratum, Phys. Letters **3**, 256 (1963).



FIG. 3. Plot of R(t) versus delay time for the 2084-keV state of Ce¹⁴⁰ in aqueous solution, in an external field of 42.1 kG.

These spectra were combined to form the ratio R,

$$R = 2 [W_{+}(t) - W_{-}(t) / W_{+}(t) + W_{-}(t)] = (\frac{3}{2}) A_{2} \sin 2\omega t, \quad (1)$$

where A_2 is the coefficient of $P_2(\cos\theta)$ in the unperturbed angular correlation,⁶ and ω is the Larmor precession frequency. Equation (1) is applicable to cascades for which the angular correlation is given by $W(\theta) = 1 + A_2 P_2(\cos\theta)$.

Three independent runs were made on liquid sources, yielding the results given in Table II. For one of the better runs R is shown in Fig. 3. The final value for the g factor of the 2084 keV state from these data is

$g = +1.014 \pm 0.038$.

In deriving this value we have assumed that the paramagnetic correction for these experiments on liquid samples is zero. This point is discussed below. Lanthanum metal sources were studied by differential

TABLE II. Summary of results for 2084-keV state of Ce¹⁴⁰.

Source	Field (kG)	Temper- ature (deg K)	gβ	β
La ³⁺ solution La ³⁺ solution La ³⁺ solution Weighted average	29.7 29.7 42.1	298 298 298	$\begin{array}{c} 1.115(95) \\ 0.994(59) \\ 0.998(42) \\ 1.014(38) \end{array}$	(1.00) (1.00) (1.00) (1.00)
La metal La metal Weighted average (see text)	30.0 33.95	298 298	1.325(113) 1.376(75) 1.361(62)	1.31 1.36 1.35(6)
La metal La metal Weighted average	30.0 29.8	77 77	2.07(37) 2.53(23) 2.40(19)	2.04 2.49 2.37(20)

⁶ For this case A_2 is approximately -0.10. The exact value is unimportant here because there is substantial attenuation arising from the finite resolving time of the counters, etc. Furtunately, the differential method relies primarily on a frequency measurement and is relatively insensitive to amplitude. The value of A_2 is discussed in Refs. 4 and 5.

TABLE III. Measured g factors for the 2084-keV state of Ce¹⁴⁰.

Method	Result	Reference
Differential+integral Differential, less than 1 cycle Differential Differential+integral	$\begin{array}{r} +1.11 \ \pm 0.04 \\ +1.15 \ \pm 0.08 \\ +0.95 \ \pm 0.10 \\ +1.014 \pm 0.038 \end{array}$	5 7 8 this work

angular correlation at 295 and at 77°K. Derived quantities are included in Table II.

III. THE g FACTOR

Three other time-differential measurements of the g factor of the 2084-keV state of Ce¹⁴⁰ have been reported.^{5,7,8} These are summarized in Table III. The agreement among these four values is fairly good, considering the difficulty of the measurement, although none of the error limits overlap the average value of 1.06.

Cerium-140 has 58 protons and 82 neutrons. Thus it is natural to attribute most of its low-lying excited states to the protons alone, assuming that the neutrons form a closed shell. In fact the proton configuration may be regarded as a closed shell at 50, plus 8 protons. These would be expected, in a single-particle model, to fill the $g_{7/2}$ subshell in the ground state, with the nextlowest $d_{5/2}$ shell empty. On this model the configurations $g_{7/2}^2$, $d_{5/2}^2$, and $g_{7/2}d_{5/2}$ (each coupled to J=4) might be candidates for the 2084-keV state. The calculated g factors for these configurations were taken from the experimental values of $+2.7881^9$ and $+4.26^9$ respectively, for the magnetic moments of La¹³⁹ and Pr¹⁴¹ to account empirically for configuration mixing. They are, respectively, +0.80, +1.71, and +1.092. Evidently, if this model were valid, the 2084-keV state would have to be regarded as primarily of $d_{5/2} g_{7/2}$ character. This simple shell-model picture does not predict such other properties as energy-level positions and transition probabilities correctly, however,² and we must look elsewhere for wave functions that describe this state.

In the light of the success of the BCS model in accounting for nuclear structure in recent years it seems much more realistic to regard the 2084-keV level as a quasiparticle state. Rho has done so and has made a detailed calculation, using Gaussian forces, for nuclei in the Ce¹⁴⁰ region.² His calculations give g=+0.92,¹⁰ slightly below the experimental values, for the quasiparticle state that he considers best, given by

$|\Psi\rangle_{4+} \cong 0.30 |d_{5/2}^2\rangle + 0.91 |g_{7/2}^2\rangle - 0.26 |d_{5/2}g_{7/2}\rangle.$

⁷ N. Kaplan, S. Ofer, and B. Rosner, Phys. Letters 3, 291 (1963).
S. G. Cohen, N. Kaplan, and S. Ofer, in *Perturbed Angular Correlations*, edited by E. Karlsson, E. Matthias, and K. Siegbahn (North-Holland Publishing Company, Amsterdam, 1964), p. 313.
⁸ M. Schmorak, H. Wilson, P. Gatti, and L. Grodzins, Phys. Rev. 134, B718 (1964).

⁸ M. Schmorak, H. Wilson, P. Gatti, and L. Grodzins, Phys. Rev. 134, B718 (1964).
⁹ I. Lindgren, in *Perturbed Angular Correlations*, edited by E. Karlsson, E. Matthias, and K. Siegbahn (North-Holland Publishing Company, Amsterdam, 1964). Appendix 1.

lishing Company, Amsterdam, 1964), Appendix 1. ¹⁰ With the single-particle g factors given here, Rho's wave functions would give g = +0.90.

Rho's g factor was also calculated using the experimental moments of La¹³⁹ and Pr¹⁴¹. Although Rho's calculation was quite involved it nevertheless included several approximations that are necessary to make such calculations tractable. Thus we feel that the theoretical value of 0.92 represents fairly good agreement with experiment, and that the model is therefore essentially correct. At the same time a discrepancy of about 10%remains. It would take us too far afield to discuss the possibilities for adjusting parameters in the theory in order to produce a g factor larger than +0.92. Rho has discussed this complex problem. It seems doubtful that exact agreement can be obtained without improving the theory somewhat, as opposed to simply adjusting parameters.

IV. HYPERFINE INTERACTIONS

One object of these experiments was to evaluate the "paramagnetic correction" that must often be invoked in angular-correlation measurements to account for magnetic-hyperfine-structure effects arising from interaction of the nucleus with orbital electrons. These effects are usually treated by defining an effective magnetic field \mathbf{H}_{e} :

$\mathbf{H}_e = \mathbf{H}_0 + \mathbf{H}_i$.

Here \mathbf{H}_0 is the external field and \mathbf{H}_i is a hyperfine field induced by H_0 . For the general case one cannot define a time-independent vector \mathbf{H}_i having the properties of a magnetic field, because magnetic hfs can be regarded as the interaction of a nuclear moment with a magnetic field only for cases in which certain symmetry conditions are fulfilled. If we regard Ce³⁺ essentially as a free ion, we may take the quantization (z) axis along \mathbf{H}_{0} and write

$$H_i = H_N \langle J_z \rangle / J, \qquad (2)$$

where H_N is the maximum possible hyperfine field, given by

$$H_N = 2\langle r^{-3} \rangle \mu_B N \langle J \rangle. \tag{3}$$

Here $\langle r^{-3} \rangle$ is the famous radial integral for the 4f electron, μ_B is the Bohr magneton, and N is a standard factor in hyperfine-structure theory.¹¹ The parameter $\langle J_z \rangle$ is given by

$$\langle J_z \rangle \cong - [g_J \mu_B J (J+1)] / [3kT] H_0 \tag{4}$$

in high-temperature approximation. Finally then, we may write $H_e = \beta H_0$, with¹²

$$\beta = 1 + (2\mu_B^2/3kT) \langle r^{-3} \rangle Ng_J J (J+1).$$
 (5)

For Ce⁴⁺, which has the electron configuration of the xenon core, $\beta = 1$. For most nommagnetic metals β would be approximately 1, because the electrons that participate in hyperfine interactions are also in the conduction band, where their properties as fermions lead to an attenuation of their susceptibility by a factor of approximately kT/E_F , where E_F is the Fermi energy. This phenomenon, termed "Pauli paramagnetism," is familiar in metal physics.

In the metallic sources of Ce¹⁴⁰ in La we expect that the above arguments will not apply because the 4felectron is expected to be associated with the Ce ion core rather than in a conduction band. There is abundant evidence from susceptibility measurements that this is the case for pure rare-earth metals. Thus, if we may neglect crystal field effects, the Ce¹⁴⁰ atom in La metal may be treated as a free trivalent ion. Substituting N=48/35, $g_J=6/7$, $\langle r^{-3} \rangle = 3.64$ to 4.72 atomic units (a.u.)¹³⁻¹⁶ for Ce³⁺(4f¹²f_{5/2}) into Eq. 5, we find 1.35 $\leq \beta \leq 1.46$ at T = 298 °K. The experimental value (Table II) is 1.35 ± 0.06 . This quantitative agreement provides very good confirmation of the model (Ce⁴⁺ ion in solution, Ce³⁺ in La). Of course this is hardly surprising, because the cerium atom is expected to be highly ionized after β^- decay of La¹⁴⁰, and Ce⁴⁺ is the highest stable oxidation state of Ce. In aqueous solution it would certainly not be reduced in a few nsec. Lanthanum metal, on the other hand, with its conduction electrons, is about as "fast" a reducing agent as one can imagine. Nevertheless, the agreement is very encouraging, because a thorough understanding of the relevant chemistry must precede the determination of a magnetic moment by angular correlation. Cohen, Kaplan, and Ofer⁷ have made similar measurements on Ce¹⁴⁰; their results for β are in good agreement with ours.

Taking the view now that our expectations about the chemistry of our samples are confirmed, we may proceed to apply the data in Table II to the evaluation of parameters describing Ce3+ in La. The method of angular correlation rotation is especially applicable in this case, for which optical spectroscopic techniques are not applicable. Our measurements lead to only rather approximate values of these parameters, but they do demonstrate the applicability of this technique to crystal-field studies.

Lanthanum metal has a structure in which the atoms are stratified alternately in sites of cubic and hexagonal symmetry. The hexagonal sites can be simulated by the elongation of a cube along the $\langle 111 \rangle$ axis. Bleaney has shown¹³ that for praseodymium metal, with the same structure, the contributions of the hexagonal terms to Stark splitting is small compared with that of the cubic terms. We adopt the approximation that all the cerium ions in the La lattice experience a crystalline field of cubic symmetry.¹⁴ Under a cubic field the ${}^{2}f_{5/2}$ level breaks up into a quartet and a doublet. The effective

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¹¹ R. J. Elliott and K. W. H. Stevens, Proc. Roy. Soc. (London) A218, 553 (1953). ¹² C. Günther and I. Lindgren, in *Perturbed Angular Correlations*, edited by E. Karlsson, E. Matthias, and K. Siegbahn (North-Holland Publishing Company, Amsterdam, 1964), p. 357.

¹³ B. Bleaney (private communication).

¹⁴ This approximation is permissible for calculating certain hightemperature properties; it is by no means generally valid. make it here because consideration of the hexagonal term would require introducing an unknown parameter, B_6/B_4 .

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hyperfine field H_e is related to the free-ion field by $H_e = f(x)H_N$, where, for the doublet lower in energy f(x) is given by¹⁵

$$f(x) = 5 + 26e^{-x} + (32/x)(1 - e^{-x})/21 + 42e^{-x}, \quad (6)$$

where $x = \Delta E/kT$. Here ΔE is the energy spacing between the two Stark states.

It is not possible to calculate ΔE accurately, but we may estimate it by analogy with values obtained for other lattices. White and co-workers found a susceptibility anomaly for Ce in the LaAl₂ lattice¹⁵ that indicated an energy gap $\Delta E/k = 200^{\circ}$ K. Measurements

FIG. 5. Plot of $\langle r^{-3} \rangle$ versus atomic number for rare earths, after Bleaney. Three experimental points are shown and an empirical curve is drawn through them.



¹⁵ J. A. White, H. J. Williams, J. H. Wernick, and R. C. Sherwood, Phys. Rev. 131, 1039 (1963).

on Ce metal yield a splitting of 200°K.¹⁶ The limiting value of $f(\infty)$ indicates that the doublet is the lower state. Because La has a larger lattice constant than Ce we might expect that $\Delta E/k$ would be slightly lower than 200°K. In Fig. 4 we have plotted β determined for Ce in La metal at 298°K and at 77.3°K, from the data in Table II, against 1/T. The two curves A and B correspond respectively to $\Delta E/k=0$ and 200°K. The "best fit" to the data is provided by curve A, but the limits of error are such that we can only set an upper limit of $\sim 200^{\circ}$ K on $\Delta E/k$. It would be interesting to determine $\Delta E/k$ accurately by extending these measurements to lower temperatures.

Even with our very limited knowledge of the crystalfield splitting we can use the data of Table II, together with Eqs. (2)-(6), to make a tentative determination of $\langle r^{-3} \rangle$ for the 4f electron of Ce in La metal. Assuming $\Delta E/k = 200$ we obtain $\langle r^{-3} \rangle = 4.1 \pm 0.4$ a.u. For higher values of ΔE the value of $\langle r^{-3} \rangle$ would be somewhat higher.

Theoretical estimates of $\langle r^{-3} \rangle$ for the rare earths have been made by Judd and Lindgren¹⁷ and by Lindgren¹⁸ from spin-orbit coupling constants and by Freeman and Watson from unrestricted Hartree-Fock calculations.¹⁹ Bleaney has compared these with experimental determinations for several rare earths^{13,18} and suggested an empirical curve for $\langle r^{-3} \rangle$ versus atomic number. All of this information is presented graphically, along with our result for Ce³⁺, in Fig. 5. Our result is not sufficiently accurate to be definitive in extending the empirical curve, but, like the other experimental results, it lies between the two theoretical curves. It is thus a reasonable value. This lends support to our interpretation of the measurements on Ce¹⁴⁰ reported here and demonstrates the applicability of angular correlation rotation methods to the determination of such parameters.

¹⁶ T. Murao and T. Matsubara, Progr. Theoret. Phys. (Kyoto) 18, 215 (1957). ¹⁷ B. R. Judd and I. Lindgren, Phys. Rev. 122, 1802 (1961);

I. Lindgren, Nucl. Phys. 32, 157 (1962). ¹⁸ I. Lindgren, in *Perturbed Angular Correlations*, edited by

L. Linugten, in *Perturbed Angular Correlations*, edited by E. Karlsson, E. Matthias, K. Siegbahn (North-Holland Pub-lishing Company, Amsterdam, 1964), p. 322. ¹⁹ A. J. Freeman and R. E. Watson, Phys. Rev. 127, 2058 (1962).