Zeitschrift:	Archives des sciences [1948-1980]
Herausgeber:	Société de Physique et d'Histoire Naturelle de Genève
Band:	27 (1974)
Heft:	2-3: EPR of magnetic ions in metals
Artikel:	Nuclear magnetic resonance and relaxation in metals containing magnetic impurities
Autor:	Walstedt, R.E.
DOI:	https://doi.org/10.5169/seals-739318

Nutzungsbedingungen

Die ETH-Bibliothek ist die Anbieterin der digitalisierten Zeitschriften auf E-Periodica. Sie besitzt keine Urheberrechte an den Zeitschriften und ist nicht verantwortlich für deren Inhalte. Die Rechte liegen in der Regel bei den Herausgebern beziehungsweise den externen Rechteinhabern. Das Veröffentlichen von Bildern in Print- und Online-Publikationen sowie auf Social Media-Kanälen oder Webseiten ist nur mit vorheriger Genehmigung der Rechteinhaber erlaubt. <u>Mehr erfahren</u>

Conditions d'utilisation

L'ETH Library est le fournisseur des revues numérisées. Elle ne détient aucun droit d'auteur sur les revues et n'est pas responsable de leur contenu. En règle générale, les droits sont détenus par les éditeurs ou les détenteurs de droits externes. La reproduction d'images dans des publications imprimées ou en ligne ainsi que sur des canaux de médias sociaux ou des sites web n'est autorisée qu'avec l'accord préalable des détenteurs des droits. <u>En savoir plus</u>

Terms of use

The ETH Library is the provider of the digitised journals. It does not own any copyrights to the journals and is not responsible for their content. The rights usually lie with the publishers or the external rights holders. Publishing images in print and online publications, as well as on social media channels or websites, is only permitted with the prior consent of the rights holders. <u>Find out more</u>

Download PDF: 15.07.2025

ETH-Bibliothek Zürich, E-Periodica, https://www.e-periodica.ch

IV. RELATED SUBJECTS

NUCLEAR MAGNETIC RESONANCE AND RELAXATION IN METALS CONTAINING MAGNETIC IMPURITIES

BY

R. E. WALSTEDT

Bell Laboratories, Murray Hill, New Jersey 07974

I. INTRODUCTION

Over the past fifteen years or so, nuclear magnetic resonance (NMR) has proved to be a valuable tool for the study of both the static and dynamic disturbances caused by dilute impurities in metals. To facilitate our discussion let us arbitrarily divide dilute alloy systems into categories of "local-moment" and "nonmoment" impurities, where in the spirit of this conference we take the former classification to include only impurities sufficiently well-localized to allow study by *EPR* techniques.

In the loosely-defined "nonmoment" category we may find systems very near to the moment-formation condition in the Friedel-Anderson sense [1, 2]. Also, many nearly-magnetic systems are said to represent local moments which have "condensed" into a weakly-magnetic "Kondo" ground state [3]. Such "nonmoment" impurities have been found to have magnetic properties of great interest. Dilute alloy systems of this sort have been quite intensively studied by *NMR* techniques [4].

In the present instance, however, we focus attention on the "magnetic", i.e. local-moment dilute alloys, and more specifically on certain aspects of NMR studies of such alloys to be outlined presently. One may subdivide this topic into studies of (1) host and (2) impurity nuclei and the studies themselves into (a) static (i.e. shift and linewidth) and (b) dynamic (i.e. spin-lattice relaxation) component parts. Let us consider these various aspects of the subject in turn.

(1a) Static host NMR effects. The earliest resonance work on local-moment alloys [5, 6] revealed that the host NMR line is essentially unshifted by magnetic impurities, but is considerably broadened by the oscillatory (RKKY) spin-density disturbance [7] such impurities produce. An up-to-date review of host linewidth studies is also found in Reference 4, except for the fact that there is now a closed-

form theory for such impurity lineshapes [8]. In metals where the host nuclei possess quadrupole moments, there is also the static effect known as quadrupolar "wipeout" due to the Friedel charge oscillations, wherein certain nuclei close to an impurity site simply do not appear in the NMR line. This effect is, of course, not specific to local-moment impurities as it is primarily a function of the charge screened by the conduction electrons.

(1b) Dynamic host NMR effects. In the past decade or so there have been a number of studies of host nuclear relaxation effects due to magnetic impurities [9]. It is only with the recent work of Alloul and Bernier [10, 11], however, that a correct understanding of systems such as \underline{Cu} Mn has begun to emerge. This has come about as a result of their having correctly taken account of spin-diffusion barrier phenomena in these systems for the first time. Although a final resolution of the \underline{Cu} Mn case has not yet been reached, we describe briefly the host relaxation processes and summarize the progress to date in understanding this system in part II of the paper. We also discuss the beautiful results of the McHenry and Silbernagel [12] on host relaxation when spin-diffusion is absent.

(2a) and (2b) NMR of local moment nuclei. Apart from the anomalous cases of \underline{W} Mo and \underline{W} Co [13], where the d-spin hyperfine fields of the impurities are extraordinarily small, there had until recently been virtually no results available on the NMR of local moment impurity nuclei themselves. Clearly, the properties of such resonances would give the most direct contact with the dynamics of the localized spins. The recent work of Walstedt and Warren [14] on $Al_{1-x} Cu_x:Mn$ has shown such studies to be feasible and profitable. In part III we describe this work and offer further explanation (beyond Ref. 14) for the results observed. The properties discussed include ${}^{55}Mn$ NMR shift, relaxation time, d-spin hyperfine field, susceptibility, and s-d exchange coupling.

II. HOST NUCLEAR RELAXATION BY LOCAL-MOMENT IMPURITIES

We first describe briefly the various processes by which local moment fluctuations can cause nuclear relaxation. Basically, the same processes apply to both the host and impurity nuclei as was recently demonstrated by Walstedt and Narath [15]. At low temperatures one of the most important processes, known for the host nuclei as the Benoit-de Gennes-Silhouette (*BGS*) process [16], may be regarded as a second-order perturbation process [15], where, e.g., a mutual spinflip between the nucleus and local moment spin is followed by another flip between the local moment spin and a conduction electron spin. Such a process is coherent with the direct Korringa process, and the cross term between these two is known for host nuclei) as the Giovannini-Heeger [17] process. This process is important only when the impurity effects are small, so we do not consider it further. A third mechanism which may be important is the dipolar process, which is generated by the fluctuations of the local-moment spin operator S_z . These are transmitted via the coupling terms BS_zI_{\pm} .

The effects of the BGS and dipolar terms may be represented in terms of their frequency spectra as is shown in Figure 1. Since the BGS term embodies the fluctuations of the S_+ operator, its spectrum is centred at $\omega_{eo} = \gamma_e H_o$, i.e. well above the nuclear Larmor frequency $\omega_{no} = \gamma_n H_o$. For the conditions shown in the figure,



FIG. 1. — Frequency spectra associated with dipolar and BGS relaxation mechanisms as described in the text, for conditions such that $\omega_{oe}T_{1,2e} < 1$.

 $\omega_{eo}T_{2e} \rangle \rangle$ 1, the dipolar process predominates. Here T_{2e}^{-1} is the local moment *EPR* linewidth. In the opposite extreme, $\omega_{eo}T_{2e} < 1$, the spectral densities of the two processes at ω_{on} would be comparable, and their relative strengths would depend on the coupling matrix elements.

A. Case of No Spin Diffusion

If the nuclei do not communicate with one another via spin diffusion, then nuclear relaxation dynamics become particularly simple. Such a case, namely, ${}^{27}Al$ in $LaAl_2:Gd$, has been studied by McHenry and Silbernagel [12]. Here, spin diffusion among the ${}^{27}Al$, already reduced by the large internal quadrupole splitting, appears to be almost totally inhibited by the disordering effect of the Gd impurities.

The analysis of spin-lattice relaxation in this case rests on the assumption that the relevant coupling varies, on the average, as $1/r^3$, where r is the distance from the impurity to the nucleus in question. This would apply, for example, for dipolar or *RKKY* coupling. Any single impurity then relaxes exponentially with

$$1/T_1(r) = C/r^6. (1)$$

In Eq. (1) C contains the coupling coefficients and spectral density function to be specified later. It was first shown by Tse and Hartmann [18] that after a brief initial period the system magnetization then relaxes as

$$1 - M(t) / M_0 \propto exp \left[-(t/\tau_1)^{1/2} \right]$$
(2)

with

$$1/\tau_1 = (4N_0c/3)^2 \pi^3 C, \tag{3}$$

where N_oc is the volume density of magnetic sites. The striking feature of Eq. (2) and (3) is that the parameters—except C, which is to be determined—are readily available to the experimenter. This relaxation process is therefore subject to simple and direct interpretation.

Relaxation data [12] for ${}^{27}Al$ in $La_{1-c}Gd_cAl_2$ is shown in Figure 2, where we see that the behavior of Eq. (2) is dramatically realized. Values of τ_1 were extracted as a function of field *H*, temperature *T*, and concentration *c* from a series of such



FIG. 2. — Relaxation curves ln p(t) vs. $t^{1/2} (p(t) = 1 - M(t)/M_o)$ for ²⁷ Al in $La_{1-c}Gd_cAl_2$, illustrating behavior predicted by Eq. (2). Taken from Ref. 12.

experiments. The results are displayed, in part, in Figure 3, where several features are to be noted. First, the rate τ_1^{-1} appears to vary much more slowly with c than $\propto c^2$ [Eq. (3)]. Further, from the observed variation of $\tau_1^{-1} \propto \delta B_s(x) / \delta x$, where $B_s(x)$ is the Brillouin function and $x = g\mu_B H / kT$, we conclude that the dipolar mechanism is predominant. The BGS mechanism varies, in contrast, as $B_s(x) / x$



(dashed line, Fig. 3), which is in decidedly poorer agreement. For dipolar fluctuations $C \propto \tau_m (1 + \omega_{on}^2 \tau_m^2)^{-1}$, where τ_m is the correlation time for *Gd* moment fluctuations. Since $\omega_{on} \tau_m \langle \langle 1 \rangle$ (see Ref. [12]) one concludes that $\tau_m^{-1} \propto c$ and is independent of *H* and *T* over this range. The motion of the *Gd* impurity moments is therefore dominated by spin-spin couplings. Analysis in detail [12] suggests that these originate from the *RKKY* interaction.

In conclusion, then, when nuclear spin diffusion is absent, the analysis of impurity-induced relaxation becomes straightforward and yields a considerable amount of useful information. For example, the Gd EPR line is probably exchange narrowed here, so that the numbers yielded for τ_m are obtainable in no other way.

B. Diffusion Barrier Phenomena

When some diffusion among host nuclei is possible on the time scale of τ_1 [Eq. (3)], the picture of part II-A above breaks down and the situation becomes

more difficult to analyze. In general the full interpretive apparatus developed for paramagnetic impurities in dielectric crystals [19] must be invoked.

We focus our attention on the particular case of <u>Cu</u> Mn, which is being analyzed in a series of experiments by Alloul and Bernier [10, 11]. These authors have found for the purposes of analysis that the alloy may be divided into two regions spatially as shown in Figure 4. In an approximately spherical region surrounding each impurity the conduction-electron spin-density (and therefore local NMR shift) is varying so rapidly that neighboring nuclei cannot communicate via spin-diffusion. These nuclei therefore relax according to the analysis of part II-A above. The remaining "bulk" nuclei are in good enough contact to maintain a common spin temperature and relax exponentially. The radius of the spherical region is the diffusion barrier radius b. With the electron spin density varying as $\sim r^{-3}$ it is easy to show that $n_b \propto \langle S_z \rangle$, where $n_b = 4\pi N_o b^3 / 3$ is the number of spins "isolated" within the diffusion barrier. The relaxation time of the bulk spins is simply obtained by volume integration of Eq. (1) from b to ∞ . This yields

$$1/T_1(\text{bulk}) = \frac{4\pi N_0 c C}{3b^3} = \frac{16\pi^2 N_0^2 c C}{9n_b}.$$
(4)

Equation (4) leads to the important observation, overlooked in previous work, that motion of the diffusion barrier as embodied in the factor n_b plays a major role in the *H* and *T* dependence of T_1 (bulk).

The experimental results [10] bear out the picture presented here. We summarize the most important findings briefly. The observed relaxation curves $[ln(1-M(t)/M_o) vs. t]$ are found to consist of a superposition of an $exp(-at^{1/2})$ term and an exponential (bulk) term which predominates at large t. By extrapolation these terms can be separated, yielding data for both relaxation processes as well as for n_b from the fraction which relaxes as $exp\{-at^{1/2}\}$. In this way n_b was indeed found to vary as $\langle S_z \rangle$. Taking account of this in Eq. (4), it was then found that $C \propto T^{-1}$ and independent of H. The latter result was found to hold in the region where H/T is small enough so that $\langle S_z \rangle \propto H/T$. Further, the impurity relaxation was found to be linear in c and analysis of the $exp\{-at^{1/2}\}$ portion of the relaxation gave values of C consistent with the bulk behavior.

The principal finding, i.e. that $C \propto T^{-1}$ and independent of H, suggests two possibilities for the underlying relaxation mechanism. Referring to Figure 1, the peak spectral densities for the dipolar and BGS mechanisms ($\propto T_{1e}$ and T_{2e} , respectively) both vary as T^{-1} as found. Thus one has either $\omega_{oe}T_{2e} \rangle \rangle$ 1 and therefore a dominant dipolar term, or $\omega_{oe}T_{2e} \langle \langle 1 \rangle$ with both terms contributing but BGS (by estimate) predominant. The former possibility is rendered unlikely by the fact that no evidence for a BGS term appears at even the lowest fields used ($\sim 1 kG$), making the BGS term unreasonably small. The latter option yields, however, a value for J_{sd} of several eV, in clear disagreement with the value obtained in NMR studies of ${}^{55}Mn$ in liquid \underline{Cu} Mn (see part III) as well as negative magneto-resistance data [20].

While questions remain, then, the \underline{Cu} Mn host NMR work has made important advances in understanding the nature of magnetic impurity relaxation in metals, with the clear demonstration of diffusion barrier phenomena being especially significant. For example, all previous work on this effect [21] must now be reexamined in the light of these developments. It remains an interesting challenge to unify the results of various experiments on the \underline{Cu} Mn system, with particular regard to the conjecture [11] that Kondo condensation [3] may play a role here.

III. NMR OF ⁵⁵Mn IN LIQUID $Al_{1-x}Cu_x:Mn$

We turn now to the subject of direct NMR of the local moment nucleus itself in dilute magnetic impurity systems. It is apparent that such experiments pose formidable technical difficulties, with broad lines, weak signals and a distinct possibility of unmanageably short relaxation times. Recently, however, such a study has been carried out at high temperatures on dilute Mn in $Al_{1-x}Cu_x$ alloys [14]. In this final section of the paper we discuss the background for this experiment, give a brief summary of the results and some further interpretive remarks.

The possibility of this kind of experiment was recently discussed in a paper [15] examining the relaxation processes expected for nuclei on local moment sites, with particular reference to the case of ${}^{55}Mn$ in Cu Mn. The nuclear relaxation appears to be dominated by a process which involves the localized spin as an intermediary, and which may be calculated in second-order perturbation theory. This process has been identified [15] with the BGS process for host nuclei. When the single-impurity EPR linewidth becomes comparable with the applied field H, the perturbation picture breaks down, giving at higher T/H a process which may be simply regarded as due to a fluctuating local-moment hyperfine field with a correlation time $\sim T_{1e}$:

$$1/T_{1n} = 2A_d^2 S(S+1) T_{1e}/3\hbar^2,$$
(5)

where T_{1e} is assumed to be given by a Korringa-type expression in terms of the exchange parameter J_{sd} [22]. Calculated rates T_{1n}^{-1} as a function of T for several values of H are shown in Figure 5 for Cu Mn, assuming $J_{sd} = 1 eV$. Taking $T_{1n}^{-1} \sim 10^5 \text{ sec}^{-1}$ as a rough upper limit of feasibility, we see that one must either work at He temperatures in a large field or in the neighborhood of the melting point of $Cu (\sim 1350 K)$. The latter possibility formed the basis for the work considered here.

The $Al_{1-x}Cu_x$ host alloy system was chosen as a host for Mn impurities for the reasons that (a) extensive studies of the impurity susceptibility and host NMR $({}^{27}Al$ and ${}^{63}Cu$) were available and (b) study of the ${}^{55}Mn$ NMR across this alloy series from pure Al to pure Cu permits an examination of the Mn moment-formation



FIG. 5. — Calculated contribution to spin-lattice relaxation rate of local-moment nucleus from fluctuating hyperfine coupling with moment as a function of temperature for four values of applied field. Parameters assumed here: S = 5/2, g = 2, $J_{sd} = 1 \ eV$, $A_d = 44.10^{-4} \ cm^{-1}$, (host density of states) $\rho_s = 0.16 \ eV^{-1} \ atom^{-1}$ for each spin orientation. From Ref. 15.

process. The high-temperature Mn impurity susceptibilities across this alloy series, as taken from the literature [23-25], are shown in Figure 6. As we traverse the series $Al \rightarrow Cu$, χ_{Mn} increases by a factor ~ 3 and goes from a slowly increasing function of T to the Curie law of pure <u>Cu Mn</u>. There is no sharp "transition" from nonmagnetic to magnetic behavior. One might say from χ_{Mn} alone that local-moment behavior appears for $x \ge 0.60$. The NMR results serve to sharpen this rather broad conclusion and raise the "threshold" value of x to 0.80 or more.

A description of the experiments and discussion of the results has already been given [14]. Here we summarize the main results and discuss further the moment formation process as $x \rightarrow 1$ in the host alloy series. The results are as follows:

(1) By plotting the observed ${}^{55}Mn$ shift K_{Mn} against χ_{Mn} with temperature the implicit parameter, one may partition the underlying structure of these quantities into orbital and localized *d*-spin components, yielding the variation of the orbital shift K_{orb} and the *d*-spin hyperfine coefficient α_d with composition as shown in Figures 7 and 8, respectively. α_d is proportional to the core-polarization hyperfine field per Bohr magneton H_{hf}^d . We return to a detailed discussion of these data after the summary.



FIG. 6. — Manganese impurity susceptibility χ_{Mn} in a series of liquid $Al_{1-x}Cu_x$ alloys (Refs. 23-25). The discontinuity seen in the Cu and $Al_{.20}Cu_{.80}$ data takes place at the melting point.

(2) The spin-lattice relaxation rate (corrected for variations of T and α_d), assumed to be dominated by the enhanced *d*-spin Korringa mechanism [26] at the *Al*-rich end, makes a rather sudden transition to a value larger by a factor ~ 5 as pure *Cu* is approached. This is interpreted as the onset of the mechanism of Eq. (5). A detailed analysis leads to deduced values of $J_{sd} = 0.82 \pm 0.10 \, eV$ and $1.22 \pm 0.15 \, eV$ for the liquid and solid phases of *Cu Mn*, respectively. This may be compared with the low-temperature magnetoresistance value of $0.8 \, eV$ [20].

(3) The origin of the observed "jump" in J_{sd} across the melting point is not understood. It does, however, offer an explanation for the corresponding jump of $\sim 6\%$ in χ_{Mn} at the melting point [23] in terms of the $(1 + \rho_s J_{sd})$ correction factor for the free-spin Curie susceptibility found on both the Anderson [27] and J_{sd} [3, 7]



FIG. 7. — Experimental variation of K_{orb} for $Al_{1-x}Cu_x$:⁵⁵Mn with composition (x), showing model theory curves calculated as described in the text. K_{orb} vanishes for pure Cu as expected for an S-like moment.



FIG. 8. — Experimental variation of α_d for $Al_{1-x}Cu_x$:⁵⁵Mn with composition (x), showing model theory curves calculated as described in the text.

models of the impurity state. Here ρ_s is the host density of states. The correction comes out to be of the right magnitude and sign.

For the remainder of the discussion we direct our attention to the behavior of the impurity state over the composition range where the moment "formation" process takes place as revealed by the above-mentioned data. In other words, we presume (contrary to the suggestions [24, 28] that $\underline{Al} Mn$ represents an impurity system in the Kondo-condensation state) that $\underline{Al} Mn$ does not meet the Anderson [2] criteria for moment formation because of the large virtual bound state width

$$\Delta = \pi \langle V_{kd}^2 \rangle \rho_s, \tag{6}$$

that $\underline{Cu} Mn$ (obviously) does, and that a transition from one to the other takes place as ρ_s and Δ diminish across the $Al \rightarrow Cu$ series.

As we see from Figures 7 and 8, this transition results in some rather curious behavior in K_{orb} and α_d . Both parameters are seen to increase gradually at first, then drop sharply at the *Cu*-rich end. Nothing in the variation of χ_{Mn} (Fig. 6), for example, would presage this behavior. The close correspondence between the K_{orb} and α_d curves suggests a common mechanism, and we would like to propose that something at least closely related to the Jaccarino-Walker [29] (J-W) momentformation phenomenon is operating here. It is recalled that these authors explained the occurrence of *Fe* impurity moments in *Nb-Mo* alloys in terms of near-neighbor configurations computed on the basis of statistical probabilities. The conclusion was simply that two or more *Nb* near-neighbors sufficed to quench the moment at a given *Fe* site.

Let us see how such a model might apply to the present situation, starting with the behavior of $K_{orb}(x)$ (Fig. 7). We presume the observed variation of $K_{orb}(x)$ as $x \to 1$ to be the result of a statistical "transition" from the orbital shift of the nonmoment state to the (zero-valued) K_{orb} of the fully developed moment. The initial increase of (nonmoment) K_{orb} with x is attributed to narrowing of the virtual bound state as ρ_s decreases [see Eq. (6)]. This follows because $K_{orb} = \beta \chi_{orb}$, $\beta = (2/A)$ $\langle r^{-3} \rangle_{av}$ (A = Avogadro's number), and χ_{orb} varies as Δ^{-1} or faster, if there is enhancement.

We make a quantitative estimate of $K_{orb}(x)$ as follows. At any given instant the system consists of a mixture of magnetized and unmagnetized impurities. Because of the rapid liquid motion compared with nuclear Larmor periods, a given ⁵⁵Mn nucleus will average the properties of all possible impurity states. Adopting the J-W criterion for moment formation, the experimental result is then expected to be

$$K_{orb}(x) = P_n(x) K_{orb-nm}(x), \tag{7}$$

where $P_n(x)$ is the probability of *n* or more *Al* atoms in the nearest-neighbor "shell" of a given *Mn*, and $K_{orb-nm}(x)$ is the orbital shift of the *Mn* nonmoment state at *Cu* concentration *x*. We can only guess the behavior of the latter quantity; a hypothetical $K_{orb-nm}(x)$ is plotted in Figure 7 as a dashed line. Corresponding $K_{orb}(x)$ curves for n = 1, 2, and 3 are also plotted, giving reasonable accord with the data for n = 2. Although these results are at best semiquantitative, the data are shown to be at least consistent with such a model in which two or more *Al* neighbors quench the *Mn* moment.

Application of these ideas to α_d is shown in Figure 8. Here the variation of the "nonmoment" α_d is purely hypothetical as the behavior of hyperfine fields in metals is only poorly understood [30]. We simply extrapolate the trend in α_d established

339

for $x \le 0.60$. For the "moment" α_d we adopt the value found for *Cu Mn* and assume it to be constant with x. The resulting statistical averages are again shown for n = 1, 2, or 3 *Al* neighbors required to quench the moment. The data appear to fall in the range of $n = 2 \rightarrow 3$ and again show consistency with a model of this sort.

One may also attempt to apply these ideas to the powerful local-moment relaxation mechanism, which is seen [14] to change even more abruptly than K_{orb} or α_d as $x \to 1$. It is not clear, however, that the relaxation mechanism itself is not influenced by the rapid fluctuations of environment in an $Al_{1-x}Cu_x$ alloy. Thus, one may not be able to assume it is independent of composition for the "moment" impurities, and no firm conclusion can be drawn.

The *J*-*W* picture gives a qualitatively reasonable, if not rigorous, explanation for the variation of K_{orb} and α_d with composition. The specific assumptions of this model may not be necessary here, but it offers at least a conceptual framework with which to take account of the inhomogeneous nature of these alloys, which clearly may play an important role.

The obvious question remaining is whether the J-W model can account for the occurrence of moments in these alloys in terms of the susceptibilities, as in the original work. The data of Gruber and Gardner [24] (Fig. 6) do not follow any well-defined temperature law; thus one is stopped by lack of knowledge about the unmagnetized susceptibility behavior. One can, however, compare with the Curie-Weiss behavior found for these alloys by Myers and Westin [31] in the neighborhood of room temperature. Surprisingly, these authors find little or no change or effective moment over the range $0.82 \le x \le 1$ where the most dramatic changes in our parameters take place. Thus we are forced to conclude either (1) that the behavior we observe is limited to the liquid state, (2) that the Mn moment becomes less stable at high temperatures, or (3) that the solid-phase alloys are not random.

There appears to be little theoretical basis for possibilities (1) or (2) above. For example, the measured J_{sd} values [14] suggest via the Schrieffer-Wolff transformation [32] $(J_{sd} \propto \langle V_{kd}^2 \rangle)$ that the virtual bound state is narrower in the liquid [Eq. (6)], rendering the moment *more* stable according to the Anderson [2] criterion.

Finally, we note the apparent contradiction between the impurity susceptibility results of References [24] and [31]. In particular, the $Al_{.20}Cu_{.80}$: Mn liquid-state results [24] appear to exhibit rough Curie-Weiss behavior with $\theta \sim 350 K$, as opposed to $\theta \sim 0 K$ at lower temperatures [31]. Experiments are underway to resolve these inconsistencies.

ACKNOWLEDGMENTS

The author would like to thank Y. Yafet, M. A. Butler and W. W. Warren, Jr. for several useful discussions of this material.

REFERENCES

- J. FRIEDEL, Suppl. Nuovo Cimento VII, 287 (1958); A. BLANDIN and J. FRIEDEL, J. Phys. Radium 19, 573 (1958).
- [2] P. W. ANDERSON, Phys. Rev. 124, 41 (1961).
- [3] See, e.g., J. KONDO, in *Solid State Physics*, Vol. 23, F. Seitz, D. Turnbull, and H. Ehrenreich, Eds., Academic Press, New York, 1969, p. 183; A. J. HEEGER, *idem*, p. 283.
- [4] A thorough review of NMR studies of such systems has been given by A. NARATH, CRC Critical Reviews in Solid State Sciences 3, 1 (1972).
- [5] J. OWEN, M. E. BROWNE, W. O. KNIGHT and C. KITTEL, Phys. Rev. 102, 1501 (1956).
- [6] J. OWEN, M. E. BROWNE, V. ARP and A. F. KIP, J. Phys. Chem. Solids 2, 85 (1957).
- [7] K. YOSIDA, Phys. Rev. 106, 893 (1957).
- [8] R. E. WALSTEDT and L. R. WALKER, Phys. B9, 4851 (1974).
- [9] A review of all but the most recent work is given by B. GIOVANNINI, P. PINCUS, G. GLADSTONE and A. J. HEEGER, J. de Physique 32, Cl-163 (1971).
- [10] H. ALLOUL and P. BERNIER, LT 13, Boulder, Colorado (1972); P. BERNIER and H. ALLOUL, J. Phys. F 3, 869 (1973).
- [11] H. ALLOUL and P. BERNIER, to be published.
- [12] M. MCHENRY and B. SILBERNAGEL, Phys. Rev. Lett. 27, 426 (1971); Phys. Rev. B 5, 2958 (1972).
- [13] K. C. BROG and W. H. JONES, Phys. Rev. Lett. 24, 58 (1970); A. NARATH, K. C. BROG and W. H. JONES, Phys. Rev. B 2, 2618 (1970).
- [14] R. E. WALSTEDT and W. W. WARREN, Jr., Phys. Rev. Lett. 31, 365 (1973).
- [15] R. E. WALSTEDT and A. NARATH, Phys. Rev. B 6, 4118 (1972).
- [16] H. BENOIT, P. G. DE GENNES and D. SILHOUETTE, Comptes Rendus 256, 3841 (1963).
- [17] B. GIOVANNINI and H. HEEGER, Solid State Commun. 7, 287 (1969).
- [18] D. TSE and S. R. HARTMANN, Phys. Rev. Lett. 21, 511 (1968).
- [19] G. R. KHUTSISHUILI, Proc. Inst. Phys., Academy of Science of Georgia (USSR) 4, 3 (1956);
 P. G. DE GENNES, J. Phhys. Chem. Solids 7, 345 (1958).
- [20] P. MONOD, Phys. Rev. Lett. 19, 1113 (1967).
- [21] See Ref. 9 and the papers in Ref. (10) and (11) for an extensive listing of previous work.
- [22] J. KORRINGA, Phys. Rev. 16, 601 (1950); H. HASEGAWA, Progr. Theor. Phys. (Kyoto) 21, 483 (1959).
- [23] J. A. GARDNER and C. P. FLYNN, Phil. Mag. 15, 1233 (1967).
- [24] O. F. GRUBER and J. A. GARDNER, Phys. Rev. B 4, 3994 (1971).
- [25] E. W. COLLINGS, J. Phys. (Paris), Colloq. 32, Cl-516 (1971).
- [26] P. LEDERER and D. L. MILLS, Solid State Commun. 5, 131 (1967).
- [27] D. J. SCALAPINO, Phys. Rev. Lett. 16, 937 (1966).
- [28] G. GRUNER and A. ZAWADOWSKI, Solid State Commun. 11, 663 (1972).
- [29] V. JACCARINO and L. R. WALKER, Phys. Rev. Lett. 15, 258 (1965).
- [30] One can make a rough qualitative accounting for the variation of α_d (nonmoment) in terms of the corepolarization mechanism (D. A. GOODINGS and V. HEINE, Phys. Rev. Lett. 5, 370 (1960)) through the variation of *d*-orbital size with *x*. For example as *x* increases, the number of *d*-electrons should also increase. This makes the *d*-orbitals larger, and, in turn, the core-polarization field more negative, etc.
- [31] H. P. MYERS and R. WESTIN, Phil. Mag. 8, 1969 (1963).
- [32] J. R. SCHRIEFFER and P. A. WOLFF, Phys. Rev. 149, 491 (1966).

DISCUSSION

RIVIER: How do you count the nearest neighbours in a liquid?

WALSTEDT: Well, in a liquid it turns out that people have done neutron scattering experiments that indicate that there really is a nearest neighbour shell which has essentially the same configuration as in the solid. I admit this is rather a murky point.

ALLOUL: I would just like to make some statements about what can be done by the techniques we used for the measurement of the nuclear relaxation of the host nuclei at low temperatures. You have explained how we obtained the information we are interested in, that is the fluctuations of the impurity moment. The fact is that we can measure a relaxation time of the impurity which is extremely different from the relaxation time measured by EPR in these systems which are bottlenecked. In fact we measure something that is unbottlenecked, which means that we measure directly the so-called T_{ds} which is very hard to get from EPR. In the case of CuMn and CuCr we find avery nice Korringa type of behaviour and the J value which can be drawn is much bigger than the EPR figure, at least a factor of two bigger. This is rather interesting because it could be related to the calculation that Götze did this morning.

DUPREE: I would like to say that it would be very nice if the statistical model you put forward is true, but there is some data around which you might have difficulty in explaining with this model, and these are the susceptibility measurements. These have been done on a much finer scale varying the electron densities in AlSn and AlGa for example. These show a bump analogous to the bump in the K orbital that you have there. I think a bump in the susceptibility, if true, is really difficult to explain on a statistical model.

WALSTEDT: I have been scrupulously careful to avoid saying anything about any other system !