Study of Phase Transitions
by Means of Nuclear Magnetic Resonance Phenomena

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STUDY OF PHASE TRANSITIONS BY MEANS OF NUCLEAR MAGNETIC RESONANCE PHENOMENA

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Abstract

Although the nuclear magnetic resonance condition generally depends only on the gyromagnetic ratio of the resonating nuclei, the width and amplitude of the resonance depend critically on the substance containing the resonating nuclei. A basic factor affecting the width is the characteristic flipping time for the substance - i.e., the average time it takes a molecule to change its orientation appreciably. This fact has been applied in a study of a group of molecular solids which exhibit two or more phase in order to investigate the mechanisms of these transitions. Of the two mechanisms proposed, rotational and order-disorder, the former was eliminated in favor of the latter for HCl, HB,, HI, H,S, and H,Se, while free molecular rotation is shown to be plausible in CH, and CH,D.

This report is a condensation of a thesis with the title "Study of the Solid State by Means of Nuclear Magnetic Resonance Phenomena" submitted by the author in partial fulfillment of the requirements for the degree of Doctor of Philosophy in Physics at the Massachusetts Institute of Technology, 1948.
1. Introduction

The first successful detection of radio-frequency transitions between Zeeman levels of nuclei in liquids and solids was announced by Purcell, Torrey, and Pound and by Bloch, Hansen, and Packard in 1946. It became apparent early in these investigations that the characteristics of a nuclear magnetic resonance depend directly on the material in which the resonating nuclei reside. This, coupled with the fact that the ratio of resonant frequency to magnetic field is a linear function of the nuclear g-factor, indicated three lines of research:

(1) The accurate measurement of nuclear g-factors.
(2) An investigation of the interactions which affect the resonance characteristics - i.e. nuclear relaxation processes.
(3) Study of internal properties of solids and liquids by means of nuclear magnetic resonance phenomena. It is one phase of the last of these characteristic experiments with which this investigation is chiefly concerned.

If a substance containing magnetic nuclei of magnetic moment $\mu$ and angular momentum $\hbar$ is placed in a z-directed magnetic field $H_0$, there will result a nuclear paramagnetism which may be found from Curie's law:

$$M_0 = \frac{N(g_n \mu_n)^2 I(I+1)}{3kT} H_0$$  \hspace{1cm} (1)$$

where $M_0$ is the static magnetization, $N$ the number of nuclei/cc., $g$ is the nuclear g-factor, $\mu_n$ the nuclear magneton, $I$ the nuclear angular momentum in...
units of $h/2\pi$, $k$ the Boltzmann factor, and $T$ the absolute temperature. Now if a radio-frequency field is introduced in the $x$-$y$ plane with the Larmor frequency

$$\nu_o = \frac{\mu H_0}{\hbar} \text{ or } \omega_o = \gamma H_0$$

(2)

where $\gamma = \mu/\hbar$, a resonant absorption and dispersion will occur. For diamagnetic materials the ratio of $\nu_o$ to $H_0$ generally depends only on the magnetic moment and angular momentum of the nuclei concerned, and the peak absorption will be found at the same resonance condition for the same nuclei, regardless of the material in which the nuclei reside. The width and magnitude of the nuclear magnetic resonance, however, are found to depend critically on the substance containing the resonating nuclei. More specifically, these properties are found to depend on two modes of interaction affecting the nuclei: (1) The interaction between the nuclei and the lattice containing them, expressed in terms of the spin-lattice or thermal relaxation time $T_1$. It is the resultant interchange of energy which enables the nuclear system to attain thermal equilibrium at a temperature $T$ and to attempt to maintain this equilibrium upon exposure to resonance radiation, (2) The magnetic interaction between the nuclei, which is expressed in terms of a spin-spin relaxation time $T_2$. This effect manifests itself primarily in a broadening of the resonance line width. The experiments described below are chiefly concerned with this latter mode of interaction.

If a resonating nucleus is surrounded by magnetic nuclei, there will be superimposed on $H_0$ a local magnetic field $H_{loc}$ which may be of the order of several gauss. The resonant condition is then $\omega_0 = \gamma(H_0 + H_{loc})$. Since $H_{loc}$ depends on the orientations of all neighbors, the result is a dispersion of the magnetic fields at the various nuclei about the applied $H_0$, the average field at the nuclei still being $H_0$. From the resultant broadening in the energy levels, $\Delta E = \hbar \gamma H_{loc}$, we may define a time $T_2$ on the basis of the uncertainty principle of quantum mechanics: $T_2 = \hbar/\Delta E = 1/\gamma H_{loc}$. It should be noted that $H_{loc}$ represents a deviation of the magnetic field from an average $H_0$. Therefore, gradients in the magnetic field due to the inhomogeneity of the magnet serve also to broaden the line.

Since the internuclear fields are expected to be of the order of a few gauss or more in magnitude, most solids fit well with this extremely elementary picture, giving resonance line widths of a few gauss or more. That this picture on the basis of a rigid lattice is inadequate becomes particularly obvious in the case of most liquids and gases and a few solids,  

1. There are other factors which may contribute to the line width but which do not enter into the cases under study. A more complete discussion of this matter may be found in the reference given in footnote 4 on p. 1.
where line widths of a small fraction of a gauss are found. The key to these lines may be seen from the fact that no mode of spin-lattice relaxation is possible in the rigid lattice so far considered. A generally useful and satisfactory theory of both spin-lattice and spin-spin interactions has been developed by B.P.P. On the basis of general lattice motion they derive fairly accurate values for $T_1$ and explain details of line narrowing.

Qualitatively the resonance line width is reduced by an averaging out of the internuclear fields due to the lattice motion. This is particularly obvious in the extreme case of a nucleus residing in a freely rotating molecule, since rotational periods are much smaller than times associated with the nuclear resonance. In most cases of interest all orientations of the molecule are equally probable. As a consequence, the internuclear fields essentially average out to zero so that an extremely narrow line results. Actually in the case of water a line width of $10^{-4}$ gauss is predicted. In practice it is impossible to observe this "true" line width since it would require, at a working field of 7000 gauss, a homogeneity of almost one part in $10^8$ over the sample. Thus the line width in most liquids is determined entirely by the homogeneity of the magnet.

The theory referred to above approximates its description of the lattice motion by means of a single parameter $\tau_0$. This so-called correlation time (or "flipping" time) is essentially the average time it takes for a molecule to change its position or orientation appreciably. This is very closely related, through a factor of the order of one, to the "characteristic relaxation time" introduced by Debye in his theory of polar molecules. The effect of $\tau_0$ on $T_2$ and on the line width may be seen as follows:

Let us denote by $T_2^*$ the spin-spin relaxation time in the limiting case of the rigid lattice. Thus, for this case, $T_2^*$ represents the average lifetime of a nucleus in a particular energy state (or orientation). Now if $\tau_0 \gg T_2^*$, the slow variation of the internuclear fields during the "lifetime" of a nuclear orientation will have a negligible effect on the spin-spin interaction. Therefore we have

\begin{equation}
\text{Case I: } \tau_0 \gg T_2^*. \quad T_2 = T_2^* \quad \Delta H = \frac{1}{T_2^*}
\end{equation}

1. Bloembergen, Purcell, Pound, loc. cit.
If on the contrary \( T_0 \ll T_2 \), considerable averaging out of the internuclear fields occurs during a normal nuclear level "lifetime". As a result the spin-spin interaction is greatly reduced and the relaxation time increased accordingly. In this case it may be shown that, in general, the relaxation time varies inversely as the correlation time.

**Case II:** \( T_0 \ll T_2 \) 

\[
T_2 \approx \frac{1}{T_0} 
\]

(3b)

\[
\Delta H \propto T_0. 
\]

Thus it becomes apparent that the line width is a sensitive function of the correlation time.

On the basis of a simplified calculation, by taking into account the effect of only the nearest neighbor of a resonating nucleus, B.P.P. find for \( T_2 \) the relation:

\[
(1/T_2)^2 = K \tan^{-1}(2 T_0/T_2) 
\]

(4)

where \( K \) is a constant which includes the internuclear distance and other constants which are independent of frequency and of temperature. This has the properties discussed above; namely, for \( T_0 \gg T_2 \), \( T_2 = (2/\pi K)^{1/2} \), which may be shown to be equal to \( T_2 \), while for \( T_0 \ll T_2 \) we find \( T_2 = 1/2K T_0 \). The form of a plot of \( \log T_2 \) vs. \( \log T_0 \) is immediately apparent. This is made up of two predominant portions, a horizontal line for \( T_0 \ll T_2 \) and a line whose slope is \(-1\) for \( T_0 \ll T_2 \). If one reasons that the correlation time loses its "control" of \( T_2 \) when it reaches a value equal to \( T_2 \), one finds that the break between the horizontal and sloping part of this graph occurs near \( T_0 = T_2^{1/2} \). Very little accuracy is lost if we approximate this plot by two straight lines, one horizontal and the other with slope \(-1\), intersecting at \( T_0 = T_2^{1/2} \).

Another factor which affects \( T_2 \) is the spin-lattice relaxation time. It may be seen that if the motion of the lattice transporting the nuclei has frequency components near \( \nu_0 \), this motion is capable of inducing transitions between Zeeman levels. Although this picture is oversimplified, it indicates essentially the means of energy transfer between the lattice and the nuclei. \( T_1 \) is found to be \( 1/2W \), where \( W \) is the probability of an induced transition. Now if \( T_1 \) is comparable to \( T_2 \), the former will also serve to limit the lifetime of a nucleus in a particular state. Thus \( T_1 \) may also contribute to the line width. However, under the conditions of the experiments, one of two situations generally exists. Either \( T_1 \gg T_2 \), which is usually the case for broad lines, or \( T_1 \ll T_2 \) for the sample, and both are in the range of .01 sec.
to several seconds. In the former cases $T_1$ has no effect on the line width, while in the latter case the line width is determined by the homogeneity of the magnetic field, and $T_1$ does not influence the experimentally observed line widths for small r-f fields.

2. Phase Transitions in Solids

Many solids undergo drastic changes in their physical properties at critical temperatures. Such transitions may be placed in one of two categories. The so-called first-order transitions occur at one particular temperature and are accompanied by a latent heat, usually due to a change in crystal symmetry. In contrast, second-order transitions are identified by $\lambda$-points with an anomalously high specific heat over the transition temperature range, generally of the order of a few degrees Kelvin. One class of solids showing second-order transitions are alloys. The order-disorder phenomenon responsible for the transitions in these substances is well understood. However, there is a large class of substances, including many molecular solids, which show second-order transitions whose mechanisms are not completely understood. These transitions may possibly be explained in terms of a change in molecular activity. It is this factor which points out the potentialities of studying this group of solids by the nuclear magnetic resonance experiment, since it has been shown that internal activity has a profound effect on the properties of the resonance. For the sake of completeness it should be noted that a few first-order transitions - e.g., in HCl - should also be of interest in any such investigation.

There are two conflicting theories to explain the transitions in question. One theory, put forward by Pauling\(^1\) and Fowler\(^2,3\) associates the transition temperature with a change from rotational oscillation of the molecules or ionic groups below the critical temperature to a phase in which most of the molecules are freely rotating. The existence of $\lambda$-point transitions may be derived by statistical mechanics by assuming the rotation to be a cooperative phenomenon. That is, one considers that the rotation of a molecule is not independent of the motions of its neighbors and that the potential against rotation is a function of the degree to which its neighbors are rotating.

The second theory, developed by Frenkel\(^4\) and Landau\(^5\) assumes that the transition is from an ordered to a disordered state. In this theory

there is assumed a preferred orientation, occupied by a majority of the molecules, below the transition, while above the transition the molecules occupy with equal probability one of two, or more, equilibrium orientations. It should be noted that both below and above the transition the molecules flip between equilibrium orientations. The distinguishing factor between the ordered and disordered phases is that the molecules predominate in a preferred orientation in the ordered phase; however, in the disordered phase, a preferred orientation ceases to exist.

It is quite possible that either of these mechanisms may be responsible in different cases.

Since the line width of the nuclear resonance depends on the average reorientation time of the molecule through $T_2^*$, we see that it is potentially possible to gain information about these transitions and their associate mechanisms by means of a nuclear resonance in the substance in question. Let us now investigate the order of magnitude of the quantities involved, so as to determine what effect these mechanisms may be expected to have on the line width.

In general we should expect the line width in the rigid lattice to be three gauss or more. Since the gyromagnetic ratio $\gamma$ is about $2.7 \times 10^4$ for protons, this indicates that $T_2^*$ should be less than $12 \mu$sec. Now, as indicated in Sec. 1, the line width is controlled by the molecular flipping time when $\gamma_0$ is less than $T_2^*/2$. Thus the line width is generally sensitive to $\gamma_0$ for values of $\gamma_0$ smaller than $8 \mu$sec. At 7000 gauss the limit of homogeneity of the magnet used during these experiments was roughly 0.2 gauss. Therefore the lowest values of $\gamma_0$ which can be derived from line-width measurements would be in the range .05 to .5 sec.

We may obtain an idea of expected values of $\gamma_0$ from Debye's "Polar Molecules".\(^1\) For liquid normal propyl alcohol he lists values from $26 \times 10^{-10}$ second at $-60^\circ$C to $0.9 \times 10^{-10}$ second at $+20^\circ$C for the characteristic reorientation time defined in his theory. Also we should expect the reorientation time for the case of a free rotation in a solid to be of the order of a rotational period: $10^{-13}$ second or less. Therefore, it would be expected that, for rotating molecules in a solid, characteristic times of at most $10^{-10}$ second would be found.

Ice may be considered as an example of a solid with nonrotating molecules. A fairly rapid motion occurs at temperatures just below the freezing point, but the activity decreases quite rapidly as the temperature is decreased. From experiments on the dielectric constant of ice at radio frequencies Debye gives values of $2.7 \mu$sec at $-5^\circ$C and $18 \mu$sec at $-22^\circ$C for his

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reorientation time. Undoubtedly a considerable variation in these values would occur for different substances, but the above would lead us to expect values somewhere in the microseconds range. It should be noted that this range is precisely that in which the line width is effective in measuring $\gamma_c$.

The marked difference between the values of $\tau_c$ for rotation and non rotation, a factor of at least $10^4$, should be carefully noted. In addition, it is important that in the case of non rotation $\gamma_c$ may possibly fall in a range in which it directly affects the observed line width, while for rotation $\gamma_c$ is so small that the observed line width must, in all practical cases, be only as narrow as the homogeneity of the magnet permits.

The expected effect of phase transitions on line widths may now be discussed on the basis of the expected values of $T_2^*$ and $\gamma_c$. For the case of a rotational transformation, the line should be broad, typical of most solids, below the transition; upon reaching the transition, where free rotation becomes predominant, one should expect a rapid decrease, over the range of the transition, to a narrow line limited in width only by magnet inhomogeneity.

The picture in the case of an order-disorder transition is not so clear cut, since there may be little or no change in $\gamma_c$ at such a transition. Even in this case there may be some sharp change in the line width at the transition temperature, since the critical temperature is frequently accompanied by a change in lattice constants or in crystal structure. The crystal structure generally becomes more symmetrical in the highest temperature phase. This would result in a change in $T_2^*$ even if $\gamma_c$ does not decrease to the range in which it affects $T_2$. A discontinuous decrease of $\gamma_c$ at the transition may, of course, contribute to a change in line width. In any case one might expect a decrease of $\gamma_c$ with increasing temperature, which may or may not reveal itself in the line width measurements. As Frenkel points out, $\gamma_c$ should be a function of the type

$$\gamma_c = \gamma_c^0 e^{U/kT}$$  \hspace{1cm} (5)

where $\gamma_c^0$ is the period of the rotational oscillations which would occur at absolute zero and $U$ is the activation energy required to reorient a small group of molecules. Of course $U$ itself may be a complicated function of the temperature $T$.

To summarize, if a moderate change, or zero change, in line width occurs at the transition, this must be identified with the order-disorder mechanism. On the contrary, if a rapid change to a very narrow line occurs

1. Frenkel, loc. cit.
at the transition, the rotational transformation mechanism may be indicated. However, the order-disorder mechanism is not completely ruled out on the basis of this evidence alone.

A third mechanism which applies to a few cases, for example Rochelle salt, is the so-called displacive transition. In this case the unit cell of the crystal suffers a small distortion during which a new symmetry element appears or disappears. Although this type of transition has not yet been studied experimentally by the method proposed here, it is expected that this would show very little effect on the nuclear resonance line width.

Essentially complete lists of substances which exhibit phase transitions may be found in review articles by Eucken and by Smyth. As one would expect on the basis of the mechanisms introduced to explain the transitions, these transitions are found primarily in substances composed of light molecules or of internal groups which would have low moment of inertia.

Pauling proves that for the hydrogen molecule and the methane molecule the eigenfunctions and energy levels, even in the lowest quantum state, closely approximate those for a freely rotating molecule. The case of hydrogen has recently been studied by Rollin and collaborators by the nuclear resonance method. Other substances which show transitions include: the hydrogen halides, except for the lightest member of the group, HF; the \( \text{H}_2\text{X} \) group, with the exception of the lightest member, \( \text{H}_2\text{O} \); the \( \text{XH}_3 \) group, also except for the lightest member, \( \text{NH}_3 \); a number of ammonium salts, including the ammonium halides; and a multitude of organic molecules.

3. **Apparatus and Cryogeny**

The circuit used is essentially the one described by B.P.P. The water-cooled electromagnet used for most of these experiments was designed by F. Bitter and constructed in the Research Laboratory of Electronics Shop. The magnet is fitted with detachable pole pieces eight inches in diameter and a gap of 2 1/4 inches. The pole faces are shimmed to minimize field gradients at the center of the gap. To satisfy the demands of current constancy - one part in \( 10^5 \) or better - the magnet was generally run from a 50-volt bank of submarine storage batteries. With the coils in parallel, the total current required for an induction of 7000 gauss (the resonant field for 30 Mc) is about 40 amperes. This current is controlled by a series of advance strips.
and, for less precise work, by a heavy-duty, field-control rheostat. The d-c field is generally modulated at 60 or 30 cycles by means of separately wound inner layers of the magnet coils. These supplementary coils are fed either from a variac connected to the 110-volt power line or from a current amplifier excited by a Hewlett-Packard 200-C audio oscillator. The current was monitored by means of a shunt and a Rubicon potentiometer.

The signal generator is either a General Radio 805-C or 605-B, the former having a higher voltage output and a greater frequency range. However, the 605 has the advantage that its power supply can conveniently be replaced by a well-regulated and filtered d-c power supply. This minimizes unwanted frequency and amplitude modulation on the r-f carrier.

Most of the experiments were carried out at 30 Mc (a working field of about 7000 gauss). For work at this frequency the output of the r-f bridge feeds into a low noise-figure preamplifier. Further amplification is then obtained by means of a Hallicrafters SX-28 receiver. The method of detecting the output depends on the type of signal under observation. If the resonance is fairly narrow and intense, it is most convenient to observe the receiver output on an oscilloscope whose base line sweep is the same frequency as the modulating field. The amplitude of this modulating field is adjusted to be at least a few times the resonance line width for this case. Under these conditions the line may be measured directly from the resulting oscilloscope trace, if one calibrates the modulation amplitude.

In the case of less intense resonances of a gauss or more in width the output of the receiver is fed instead to a 30-cycle twin-T narrow-band amplifier with a lock-in feature. In this case the modulation amplitude is reduced to a small fraction of the line width and the output, read on a 0-1 ma meter, is proportional to the slope of the resonance curve. The damping of the meter, which determines the bandwidth, also limits how rapidly one can traverse a resonance curve. In this case the absorption component of the resonance is more generally used and the width measured is the distance between the inflection points, i.e., between the peak deflections of the output meter. If one assumes a line shape resembling a universal resonance curve, as results from Bloch's development for example, the above definition of line width is $1/3\frac{2}{3}$ times the distance between half-value points, which is normally defined as the line width. The distance between peaks was measured in terms of the

3. Bloch makes certain approximations which lead to the assumed line shape. This line shape is only an approximation, however. Since there is no completely adequate theory for line shape, any definition of line width is arbitrary to a certain degree.
deflection of the potentiometer galvanometer, the potentiometer setting meanwhile remaining fixed.

The temperatures which must be obtained in order to study practically all of the phase transitions fall in the range from about 20°K to room temperature. Preliminary low-temperature experiments were carried on in the 9 1/2-inch gap between the unshimmed poles of the M.I.T. cyclotron magnet. These experiments were carried on in glass dewars with appropriate liquid baths as the low-temperature reservoir: helium (in one experiment), hydrogen, nitrogen, oxygen, methane, ethylene, and a dry ice-carbon bisulfide solution. In many cases the liquid had to be discarded just before making observations for one of two reasons: (1) Proton-containing liquids around the r-f coil gave appreciable signals. (2) A boiling effect, due to the lack of complete rigidity in the r-f coil, caused a large increase in background noise. Also, by removing the liquid and allowing the temperature to drift upward, it was possible to cover any temperature range desired. However, these methods were not generally suitable for quantitative measurements because: (1) The bridge balance was very sensitive to small changes in temperature. (2) The continual drift in temperature left the temperature and thermal equilibrium of the sample very much in doubt, particularly in regions of high specific heats in the vicinity of phase transitions.

The above work indicated the requirements on the cryostat used for most of the quantitative work. This was designed to fit in a glass dewar specially made for use in the small electromagnet. This dewar is about 1 1/2 inches I.D. over the bottom 5 1/2 inches, with a 2 1/8 O.D. to fit conveniently in the gap. For the remaining 8 inches of its length the dewar widens out to 3 1/8 inches I.D. to provide a greater volume for a liquid reservoir.

The cryostat operates on the principle of a variable heat leak between a thick-walled copper pot containing the sample at the bottom and a reservoir containing a suitable liquid in the wide part of the dewar. The heat leak is controlled by means of a heater placed just above the copper pot. The pot and the reservoir are connected by brass tubing with a .031-inch wall thickness, which represents a compromise between too slow a heat leak on the one hand and a too rapid loss in liquid on the other. A 3/4-inch copper tube was placed from the top of the reservoir with a heater wound over almost its entire length. This was used to keep the entire gas column above the pot at a temperature at least as high as that of the pot, in order to prevent convection currents. The space between the copper tube and the reservoir was filled with an insulating material. The reservoir was made vacuum-tight to permit pumping on the liquid and to enable convenient removal of explosive and noxious vapors. In addition, the construction of the cover was such that the space in the dewar, but outside the reservoir, could be evacuated, so that the cryostat could be made adaptable for use with liquid hydrogen.
Resistance thermometers were used to determine temperature, with one wound about midway along the copper pot and the other in the gas column several inches above the pot. Resistance was measured by means of a Leeds and Northrup Wheatstone Bridge (Type "S" Testing Set). This was found to be quite adequate for the purpose of the experiments above 80 K. Since the lower-resistance thermometer reads the average temperature of the pot rather than the temperature of the sample, the time-dependent relationship between these was checked by placing a thermocouple in the position of the sample. As a result, a waiting period of the order of 20 minutes was adopted after changes in temperature of the order of 5 to 10 K. Generally, two or more readings of line width were taken over a time of 10 or more minutes after the waiting period, and data were not accepted until these could be repeated to within the estimated accuracy of the readings. To check this procedure, line width data were usually taken in the direction of increasing as well as decreasing temperature. Practically no cases were found of hysteresis due to a lack of thermal equilibrium. The greatest trouble occurred in the vicinity of transition temperatures, because of the high specific heats (or latent heats). Actually some of these effects may be inherent in these substances, since hysteresis effects have been reported by investigators of phase transitions by other methods.

Those samples which are gases at room temperature were condensed through a capillary opening into thin-walled glass ampules dipped in liquid nitrogen. After the sample was collected, the ampule was sealed by applying a small hot flame to the capillary. The seal was then worked around into a hook so that the sample could be suspended from a string for ease in storing the samples and transferring them to the cryostat. The ampules were stored in liquid nitrogen. An outline of the chemical preparation of the samples is given in the Appendix.

4. Results and Discussion

Those substances which have been studied in some detail include (1) normal and heavy methane, CH₄ and CH₃D; (2) the hydrogen halides HCl, HBr, and HI; (3) the H₂X group, H₂S and H₂Se. In addition, some preliminary results will be reported for ammonium chloride and for natural and synthetic rubber.

4.1. The Methanes. It was the exploratory work with normal methane which gave the first indication of the possible application of this method to the study of phase transitions. Since the construction of the cryostat, no work has been done at hydrogen temperatures, the range of interest for these substances. Therefore, the results for these solids were obtained entirely during the exploratory experiments.
Normal methane shows a single λ-point transition at 20.5°K, occurring over a range of 3K. Its melting point is at 90.6°K. The methane crystal is cubic both above and below the λ-point, and the transition shows some hysteresis.1

Methane is one of the first solids in which a nuclear resonance has been observed whose line width is appreciably less than one gauss. This line width was observed must about 77.4°K and just above 20.4°K. In both cases observations were made within a few minutes after pouring out the low-temperature liquid, nitrogen and hydrogen, respectively. Since even the lower-temperature observation was believed to be above the λ-point, one experiment was tried in liquid helium in a dewar of half a liter capacity, in order to reach temperatures definitely below the λ-point. No explicit temperature measurements were made, however. That temperatures below the λ-point were reached was verified by the observation of a line width of about 10 gauss on the lock-in amplifier, as well as of a broad hump of considerable amplitude on the oscilloscope. There was no marked change in the appearance of resonance for approximately 30 minutes. Then, within a period of less than one minute, during which the bridge was being rebalanced, the resonance became narrow, as had been observed above hydrogen temperature. We therefore conclude that there was a transition in the line width from about 10 gauss to roughly 0.2 gauss in a range of the order of less than 1K. The corresponding transition in $\gamma$ is therefore from greater than 2 µsec to 0.03 µsec or less. As was pointed out in Sec. 2, this behavior is what would be expected in the case of a rotational transformation. However, the possibility that this is an order-disorder transition accompanied by a decrease in the flipping time to less than 0.03 µsec is not excluded.

In the case of CH₃D two λ-points were found, at 15.5°K with a range of 2°, and at 22.6°K with a range of 3°. The crystal structure of this solid has not been investigated, to the knowledge of the author. The melting point of deuterated methane is also at 90.6°K.

A line-width behavior very similar to that observed in normal methane was found at the higher-temperature λ-point. No investigation was carried on below 20.4°K. In this case the liquid hydrogen in the dewar was removed, except for a small amount at the bottom of the dewar. Upon returning the dewar to the magnet and balancing the bridge, a resonance of the order of 3 gauss in width was observed. No apparent change occurred in this resonance for about ten minutes. Then, within a period of less than ten seconds the line narrowed to a width of less than 0.3 gauss. This series of observations was repeated. In this case the change in line width represents

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1. Data concerning transition temperatures, etc., are from references given in footnotes 1 and 2, p. 8.
a change in $\gamma_c$ from greater than 5 μsec below the transition to less than 0.3 μsec above the transition.

As is the conclusion for CH$_4$, the behavior of the nuclear resonance line width in CH$_3$D in the vicinity of its upper $\lambda$-point is what one would expect for a rotational transformation. However, the possibility of its being an order-disorder transition is not absolutely eliminated. Although observations were not carried down to the lower $\lambda$-point, by process of elimination one would expect this to be an order-disorder transition.

4.2. The Hydrogen Halides. HCl shows a first-order transition at 98.36°K which is accompanied by a change in crystal structure from cubic in the higher-temperature modification to a less symmetrical structure at lower temperatures. HCl melts at 158.9°K.

The resultant curve of line width versus temperature, shown in Fig. 1, divides into three sections. In the comparatively short temperature range investigated below the transition the line width appears to be decreasing slowly with increasing temperature. This may be interpreted in one of two ways: (1) This line width is a measure of the rigid-lattice line width, and the temperature variation is due to slight changes in the

![Fig. 1. Line width vs. temperature for hydrogen chloride.](image-url)
lattice constants with temperature. (2) $\tau_c$ is decreasing slowly with temperature and is in the range where it affects the line width. It would be difficult to prove either one of these assumptions until an accurate rigid-lattice line width can be computed for the lower-temperature modification.

An apparently sharp drop in line width at the transition temperature constitutes the boundary between the first and second portions of the graph. In the temperature range from 98.4°K to about 135°K the graph indicates almost a perfectly linear decrease in line width with the temperature. In this case we are almost certainly in the range where $\Delta H$ is proportional to $\tau_c$, which implies a linear decrease in $\tau_c$ from about 3 usec at 100°K to about 0.08 usec at 135°K. However, there is some ambiguity in the interpretation of the sharp drop at the transition. It is likely that this is due at least in part to a change in the rigid-lattice line width accompanying the change in crystal structure. There is also some possibility that this may be associated with a sharp change in the characteristic flipping time, particularly if the first alternative above were the case and $\tau_c$ had not affected the line width below the transition.

Beyond 135°K the observed line width is determined entirely by the homogeneity of the magnetic field. This limiting line width is roughly 0.2 gauss at a resonant field of 7000 gauss. No change in the amplitude of the resonance, as observed on the oscilloscope, was noted upon melting the sample. There is little to be gained by speculating as to the variation of $\tau_c$ above 135°K, although it should be noted that the extreme possibility of free molecular rotation setting in below the melting point is not ruled out by these data. However, this does not imply a rotational transformation, for which case free molecular rotation sets in at the transition. Therefore, there is no doubt that this line-width variation fits in with what would be expected for the order-disorder transition mechanism. The possibility of a rotational transformation is completely eliminated.

Hydrogen bromide has $\lambda$-points at 89°K, with a 3° range, and at 113° and 117°K, with a 2.5° and 1.5° range, respectively. A change from a less symmetrical structure to a cubic structure occurs at the lowest transition. Hysteresis is found at all three transitions. HBr melts at 186.2°K.

Referring to the line-width versus temperature graph in Fig. 2, we see that there may be a slight change in line width at the lowest transition. However, in the temperature range between 90° and about 140°K, the line width curve is characterized by a very slow decrease with increasing temperature. It is believed that in this range we are measuring the rigid-lattice line width, although it is not impossible that the line width is affected by the characteristic flipping time, which may therefore be changing very slowly with temperature. If the former interpretation is assumed, it is believed that $\tau_c$ has become sufficiently small to take control of the line.
width in the vicinity of 140°K. Above this temperature the line width responds similarly to what was found in the higher-temperature modification of HCl. Between 140° and about 160°K this line width drops very rapidly from about 3 gauss to the observable limit of about 0.2 gauss, the drop representing, on the basis of our interpretation, a decrease in $\gamma_0$ from roughly 5 μsec to about 0.3 μsec. Again there is a range of somewhat over 20°K in the solid states in which the observed line width depends only on the homogeneity of the magnetic field. As in the case of HCl the possibility of rotational transformation at any of the $\lambda$-points is clearly eliminated, while the data satisfy the implication of the order-disorder mechanism.

The remaining member of the hydrogen halide series, hydrogen iodide, has two $\lambda$-transitions at 70° and 125°K, each with a range of 5°. HI is tetragonal (face-centered) above 70°K, while no data seem to be available for the lowest-temperature modification.

The line-width-vs.-temperature data shown in Fig. 3 are almost identical in character to those found for HBr. In this case the investigation has not been carried to the lowest-temperature modification. For HI
the line width is almost constant, with possibly a very slow decrease, from 80° to about 150°K. There is then a comparatively rapid decrease in line width from about 2.5 gauss to the observable limit of about 0.2 gauss, between 150° and about 200°K, representing an assumed decrease in $\gamma_C$ from about 6 μsec to roughly 0.5 μsec. Again there is a temperature range of somewhat over 20K° in the solid state in which the observed line width is determined entirely by the homogeneity of the magnetic field. As is the case for HCl and HBr, the data clearly deny the Pauling-Fowler interpretation of the $\lambda$-points in favor of the Frenkel-Landau interpretation.

4.3. Hydrogen Sulfide and Hydrogen Selenide. The lighter substance of the two, H$_2$S, shows phase transitions at 103.6°K and at 126.2°K. The former is a $\lambda$-transition with a range of 0.9K° and some hysteresis, while the latter is apparently a first-order transition, sharp and showing no hysteresis. The intermediate-temperature modification is known to be cubic. H$_2$S melts at 187.6°K.
The graph of the line-width-vs.-temperature data for $\text{H}_2\text{S}$, as shown in Fig. 4, may be interpreted in much the same way as the results for the hydrogen halides. In the lowest-temperature modification line width is apparently constant and is probably the rigid-lattice line width for this phase. A sharp drop in line width occurs in the vicinity of 103.6 K, the lower-transition temperature, and the line width is constant in the range of the intermediate-temperature modification, leading us again to interpret this as the rigid-lattice line width for this phase. At the upper-transition temperature another small discontinuity in line width apparently occurs followed by a sharp break in the slope at about 135 K. If we interpret this as the temperature at which $\tau_c$ becomes effective in narrowing the line, this indicates an approximately linear variation in $\tau_c$ from 3 μsec to 0.1 μsec between 135° and 165 K. Again there is a range of about 20° in the solid state in which the observed line width is limited only by the homogeneity of the magnetic field. As is the case with the hydrogen halides, the above behavior eliminates the possibility of a rotational transformation and indicates the order-disorder mechanism as the source of the transitions.

Fig. 4. Line width vs. temperature for hydrogen sulfide.
Hydrogen selenide shows two phase transitions in the solid state. The one at 82.3°K is a λ-point with a range of 7° and showing some hysteresis, while the upper transition at 172.5°K is sharp with no hysteresis, and may be a first-order transition, similar to H₂S. Hydrogen selenide is known to be cubic in its two higher-temperature modifications. The melting point is at 207.4°K.

The nuclear resonance data for protons in H₂Se, shown in Fig. 5, have somewhat different characteristics from those described for the previous cases. There is some decrease in line width at the lower transition. However, the drastic increase in line width between 110° and about 140°K and the subsequent decrease between 140° and 172°K have no precedence. If one assumes that the line width is proportional to \( \gamma \) in this range, then the indicated variation in \( \gamma \) is very startling. If it is recalled from Sec. 2 that \( \gamma \) should be proportional to \( \exp(U/kT) \), where \( U \) is the activation energy required to reorient a small group of molecules, this variation indicates that the barrier against reorientation would have to increase drastically between 110° and 140°K, so as to more than compensate for the inverse tempera-
ture dependence in the exponential. It is difficult to see how this could happen.

Although there is a considerable dispersion in the data in the vicinity of the upper transition, there seems to be a discontinuous decrease in the line width at this temperature to the comparatively narrow width of about 1.4 gauss, followed by a decrease to the observable limit of 0.2 gauss at 200°C, in this case only about 8°C below the melting point.

One factor which may have contributed to the strange results in the case of H₂Se is the possibility of the presence of some HCl as impurity.

4.4. Preliminary Results for Ammonium Chloride and for Rubberlike Materials. In both cases data were taken only during the exploratory experiments.

In the case of NH₄Cl, which has a λ-point at 242.8°C, a line width of about 5.5 gauss was found at room temperature and somewhere in the vicinity of the λ-point. At nitrogen temperature a line width of 18 gauss was observed. As in the other cases this definitely indicates that this is an order-disorder type of transition,¹ and not a rotational transformation, as has frequently been assumed.²

In natural, unvulcanized rubber a narrow line was observed, limited only by the homogeneity in the magnetic field. Therefore, the possibility of extremely short flipping times for the methyl side groups or for the hydrogen bonds, for example, is quite plausible. However, there is an apparent decrease in the number participating in this motion as the temperature is decreased, since the amplitude of the resonance decreases drastically while there is no change in the other characteristics of the resonance. A small, narrow pip is still visible at dry-ice temperature, about -78°C. Vulcanization had no effect at room temperature, while carbon loading did seem to broaden the resonance slightly. As was expected, a sample of GR-S synthetic rubber gave only a broad resonance.

4.5. Summary of the Results. We have indicated and applied a method capable of distinguishing the mechanisms of those λ-points, as well as a few first-order transformations, for which the transitions may be attributed to a change in molecular reorientation time. In the experimental studies it was shown in most cases that the transitions cannot possibly be associated with the onset of molecular rotation but satisfy, instead, the conditions of an order-disorder transformation. Those substances which fall into this classification are HCl, HBr, HI, H₂S, H₂Se, and NH₄Cl. In a few cases the results indicate the plausibility of a rotational transformation, although the order-disorder mechanism is not absolutely eliminated for these transitions. These exceptional solids are CH₄ and CH₃D at the upper transition, as well as natural rubber, for

2. Smyth, loc. cit.
which internal rotation is shown to be reasonable but is not proved unquestionably.

The experimental data for the methanes, in particular, should be made more quantitative under better conditions of temperature control. In addition, measurements of the spin-lattice relaxation time $T_1$ may possibly eliminate the ambiguity in these rare cases. This follows from the fact that $T_1$ is also a function of the characteristic flipping time $T$, with the advantage that no limitations will be imposed in the direction of short flipping times by the homogeneity of the magnetic field. However, the measurement of $T_1$ is much more difficult than the measurement of line width, or $T_2$, which has been shown to be adequate in most cases.

Appendix

Following is a brief outline of the methods by which the commercial unavailable samples were produced. In all cases at least one stage of fractionation was carried on before condensation into the ampules.

1. Hydrogen Chloride. Hydrogen chloride was prepared by dropping concentrated sulphuric acid on a mixture of hydrochloric acid and sodium chloride. The gas was dried by passing through four tubes of $P_2O_5$. It was collected in a 100-ml flask in a liquid nitrogen trap.

2. Hydrogen Bromide. Forty-eight percent hydrobromic acid was dropped slowly onto a mixture of freshly ignited sand and phosphoric pentoxide. The resulting gas was passed through two tubes of red phosphorus to remove free bromine, and through three tubes of $P_2O_5$ to remove water vapor. The gas was first collected in a 100-ml flask (drawn down in the neck) submerged in a liquid nitrogen bath.

3. Hydrogen Iodide. Hydrogen iodide was prepared by the substitution reaction $NaI + H_3PO_4 \rightarrow HI + NaH_2PO_4$. Moderate heat was applied. The apparatus was completely covered with aluminum foil to prevent decomposition of the hydrogen iodide by light. Free iodine was removed by freezing out in a 3/4-inch pyrex coil in a $-30^\circ$C trap. The temperature of the trap was maintained by regulating a dry air flow through a glass-wool-packed U-tube immersed in liquid nitrogen. This temperature was easily kept to within $\pm 1^\circ$C. Further traces of iodine were removed by passing the gas through two tubes of red phosphorus. Moisture was removed by passing the gas through three tubes of $P_2O_5$. The gas was collected in two U-tubes in liquid nitrogen baths. The HI first condensed into the U-tubes as a pure white solid. It was observed then

that after a few seconds' exposure to light a progressive discoloration of the solid HI took place, probably due to decomposition of the HI. Although considerable effort was made to prevent exposure to light during the fractionation, some decomposition did take place. The finished frozen gas therefore shows some discoloration, presumably due to free I₂.

4. Hydrogen Selenide. Hydrogen selenide was made by dropping dilute HCl slowly onto freshly made aluminum selenide: \( \text{Al}_2\text{Se}_3 + (3+x)\text{H}_2\text{O} \rightarrow 3\text{H}_2\text{Se} + + \text{Al}_2\text{O}_3 \cdot x\text{H}_2\text{O} \). The aluminum selenide was prepared by mixing 50 grams of finely divided precipitated selenium with 30 grams of aluminum powder that had been washed with acetone and ether: \( 2\text{Al} + 3\text{Se} \rightarrow \text{Al}_2\text{Se}_3 \). Five grams of the mixture were placed in a fire-clay crucible and ignited with glowing magnesium ribbon. Every few seconds three grams of the mixture were added to the glowing mass in the crucible. The aluminum selenide was stored in a vacuum dessicator until used. Dilute hydrochloric acid was slowly dropped from the dropping funnel on to the freshly made aluminum selenide. The evolved hydrogen selenide passed through a drying train of two tubes of CaCl₂ and three tubes of phosphorus pentoxide. It was collected in a 100-ml flask in a liquid nitrogen bath. Metaphosphoric acid was used to lubricate the ground-glass joints in the glass system.

5. Deuterated Methane. The methyl deuteride was prepared by the Grignard reaction: \( \text{CH}_3\text{I} + \text{M} \rightarrow \text{CH}_3\text{MgI} ; \text{CH}_3\text{MgI} + \text{D}_2\text{O} \rightarrow \text{CH}_3\text{D} + \text{MgI(OD)} \), carried on in dibutyl ether as solvent. The dibutyl ether and methyl iodide were first carefully dried and redistilled. The heavy water was dropped slowly onto the methyl magnesium iodide. The gas resulting from the reaction passed through concentrated sulfuric acid to remove the dibutyl ether, and then through three tubes of P₂O₅ for drying. In this case the gas was condensed directly into the ampules.

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