THE NUCLEAR QUADRUPOLE RESONANCE AND CRYSTAL STRUCTURE OF SOLID IODINE

By

K.W.H. Stevens

April 20, 1954

Technical Report No. 197

Cruft Laboratory
Harvard University
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The Nuclear Quadrupole Resonance and Crystal Structure
of Solid Iodine

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K. W. H. Stevens

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Abstract

Previous attempts to explain the anisotropy of the quadrupole resonance in solid iodine are critically reviewed and an attempt is made to set up a valence bond description of the iodine molecule which contains the possibility that additional weak bonds are formed in crystalline iodine. This is primarily accomplished by introducing d-hybridization. The observed bond angles are then used to estimate some of the parameters in the orbits. Use is made of knowledge about nuclear quadrupole interaction in solid $I_2$ and in the iodine atom to determine further parameters, and in this way a picture of the iodine molecule and its behavior in a crystal is obtained. An interesting possibility which then arises is that the lobes of the wave-function of different iodine molecules do not overlap very strongly because a maximum of one molecule seems to be directed towards a minimum of another. This does not correspond to the usual ideas about bonds, but as the theory passes smoothly from the notion of maximum overlap to this new situation, no discussion is given about how it should be described. It is also seen that the observation of the quadrupole resonance in single crystals of iodine (if this becomes possible) would provide very useful information and checks on the theory.
The Nuclear Quadrupole Resonance and Crystal Structure of Solid Iodine

K. W. H. Stevens*

Cruft Laboratory, Harvard University
Cambridge 38, Massachusetts

Introduction

The observation of pure nuclear quadrupole resonances in solid iodine has been reported by a number of authors [1], who find that there are considerable departures from axial symmetry in the charge distributions near the iodine nuclei. At first sight this is rather surprising, for iodine forms a molecular crystal of $I_2$ molecules, and one would expect that the charge distributions would show cylindrical symmetry about the I-I bond, as is certainly true for a free $I_2$ molecule. An explanation of the origin of this asymmetry has been put forward by Townes and Dailey [2], who point out that the I-I bond length in the solid is greater than in the free molecule, a fact which can be understood if additional weak bonds are formed in the crystal, thus weakening the main bond. A study of the crystal structure, see Figs. 1 and 2, shows that the iodine molecules lie in planes which are fairly well spaced, and that in the planes there are two directions for the I-I bond, the different sorts of molecules lying in chains. In Fig. 2 we show an enlarged picture of an iodine molecule with the positions of a pair of nearest neighboring iodines, and if, following Townes and Dailey, we assume that the additional bonds are formed in these directions, it seems fairly clear that the charge distribution near an iodine nucleus will not have an axis of symmetry. The rest of the treatment of this problem by Townes and Dailey is more difficult to understand. They assume that the charge distribution around each of the three bonds made by any iodine has cylindrical symmetry about the bond, so that only one parameter is needed to describe the quadrupole interaction with the charge in each bond (if $z$ is a bond direction $\partial^2 V/\partial x^2 = \partial^2 V/\partial y^2 = -\frac{1}{2} \partial^2 V/\partial z^2$), and then, because the two weak bonds are of equal length, they weight them

*On leave from Nottingham University, England
each with a total charge $a$ and the main bond with $1-2a$. In this way the total quadrupole interaction of an iodine nucleus is described by two parameters, $a$ and $\frac{\partial^2 V}{\partial z^2}$; $a$ is found from the anisotropy and $\frac{\partial^2 V}{\partial z^2}$ from the magnitude of the interaction. The final step is then to show that $\frac{\partial^2 V}{\partial z^2}$ found in this way agrees with the field gradient expected for a free iodine molecule. Our main criticism of this procedure is that there is no reason to suppose that because the two additional bonds are equal in length they must have equal weight, nor is there any a priori reason to assume that, apart from the different charges, $\frac{\partial^2 V}{\partial z^2}$ is the same for the weak and strong bonds. This latter assumption becomes plausible if one assumes that the bonds are made of hybridized $s$ and $p$ orbitals, but there is then the difficulty that three bonds in a plane cannot be made from $s$ and $p$ alone. The first criticism can be illustrated very simply (see Fig. 3) where the dotted lines now represent the charge densities in the additional bonds in a schematic way. The bonds will clearly have equal lengths but they do not set up equal field gradients at a given iodine: we believe that this sort of behavior is more likely to occur than equal gradients.

With these difficulties about the validity of Townes and Dailey's model we decided to try to set up a more specific model, based on the Pauling ideas of directed valence. What is needed is a reasonable description of a free iodine molecule, then a study of how such molecules might come together to form a crystal and finally a calculation of the nuclear quadrupole interaction. This ambitious programme proves to be too difficult to carry through completely, but quite a lot of progress can be made and a number of useful results obtained.

Since this work was completed another theory about the origin of the anisotropy has been presented by Robinson, Dehmelt and Gordy [3]. This differs significantly from the previous theory and from the one we propose to give. While we do not know which theory is nearest the true solution, we feel that ours has at least a more fundamental approach than the other two and that something of this sort is required, rather than further semi-empirical speculations.
FIG. 1 THE ARRANGEMENT OF IODINE MOLECULES IN A PLANE

FIG. 2 RELATIVE POSITION OF IODINE ATOMS IN SOLID I₂
FIG. 3  AN EXAMPLE WHICH SHOWS SCHEMATICALLY HOW BOND DISTANCES CAN BE EQUAL BUT GIVE RISE TO DIFFERENT CHARGE DISTRIBUTION AT A GIVEN NUCLEUS
The Free Molecule

Each iodine atom has five 5p electrons outside closed shells and we might expect that in the molecule four of these pair off in $p_\pi$ orbitals and the remaining one goes into a $p_\sigma$ orbit and pairs up with a similar electron from the other iodine. This is too simple because hybridization is expected and to allow for this we shall generalize the above orbits by expressing them in a series of spherical harmonics. Thus, if the positive direction of $z$ is taken towards the other iodine, we have:

- 2 electrons in
  $$\frac{1}{N_1} \left\{ \mid y\rangle + a \mid yz\rangle \right\}$$

- 2 electrons in
  $$\frac{1}{N_1} \left\{ \mid x\rangle + a \mid xz\rangle \right\}$$

- 1 electron in
  $$\frac{1}{N_2} \left\{ \cos \gamma \left\{ \cos \alpha \mid z\rangle + \sin \alpha \left\{ 3z^2 - r^2 \mid 3z^2 - r^2\rangle \right\} + \sin \gamma \mid s\rangle \right\}$$

where allowance is made for $s$, $p$, $d$ hybridization, $\mid yz\rangle$ etc., denotes the angular variation of a normalized wave-function, $N_1$ and $N_2$ are normalizing factors and $a$, $a$ and $\gamma$ describe the relative admixtures of orbitals of different symmetries. The normalization is such that $N_2$ will probably be near unity (not exactly because this orbit overlaps strongly with the similar one on the other iodine), while $N_1$ will depart from unity according as $a$ is large or small.

The bonding orbit (i.e., the orbit with only one electron) is expected to have a strong maximum in the positive direction of $z$, but this is not the only direction of maximum charge concentration, particularly with $d$-hybridization. In fact, if $\theta$ is measured from the positive direction of $z$ the angular variation of the amplitude of the wave-function at some radius $r$ is proportional to:

$$u \cos a \sqrt{3} \cos \theta + v \sin a \left\{ \frac{\sqrt{15}}{2} \left( 3 \cos^2 \theta - 1 \right) \right\}$$

where $u$ and $v$ denote the values of the radial parts of the $p$ and $d$ orbitals respectively ($u$ and $v$ are usually taken to be equal). This function has turning values at $\theta = 0, 180^\circ$ and $\theta_o = \cos^{-1} \left\{ \frac{1}{\sqrt{15}} \frac{u}{v} \cot \alpha \right\}$. To make $\theta = 0$, $a$
strong maximum cot $a$ must be positive, so that $\theta_o$ lies between $90^\circ$ and $180^\circ$. Thus as a result of using $d$-hybridization, the bonding orbit has subsidiary turning values on the cone of directions corresponding to $\theta_o$ as well as at $\theta = 180^\circ$. It is convenient to have a notation for the directions of these turning values, so we shall refer to them by the angles they make, i.e., $0^\circ$, $180^\circ$, and $\theta_o$ directions.

**Crystalline Iodine**

Suppose two iodine molecules are coming together; then we may ask how they will arrange themselves. If we neglect for the moment the disturbances of their charge clouds by their mutual interactions, it seems not unlikely that they either have their $\theta_o$ directions pointing towards one another, or that the $\theta_o$ of one will point towards a $180^\circ$ direction of the other. Referring to the crystal structure (Figs. 1 and 2) it is seen that one of the subsidiary bonds is only $10^\circ$ away from the $180^\circ$ direction and that the other lies between $90^\circ$ and $180^\circ$. Thus a semi-empirical understanding of why iodine has the structure it does is obtained if we assume that the molecules arrange themselves so that the $180^\circ$ direction of one is directed towards the $\theta_o$ direction of another. This hypothesis will be further strengthened if we can show that the mutual interactions of the molecules reduce the strength of the main I-I bond and move the $180^\circ$ direction in the right direction. To consider this we suppose that the interactions cause mixing of our original three orbits, so that in the crystal the electrons are distributed in orbits as follows:

$$2 \text{ in } \cos \frac{\beta}{N1} \left[ |y\rangle + a |yz\rangle \right]$$

$$2 \text{ in } \cos \frac{\beta}{N1} \left[ |x\rangle + a |xz\rangle \right] + \sin \frac{\beta}{N2} \left[ \cos \gamma (\cos a |z\rangle + \sin a |3z^2 - r^2\rangle) + \sin \gamma |s\rangle \right]$$

$$1 \text{ in } \cos \frac{\beta}{N2} \left[ \cos \gamma (\cos a |z\rangle + \sin a |3z^2 - r^2\rangle) + \sin \gamma |s\rangle \right]$$

where the plane of the iodine molecules is taken as $xz$, so the orbit which is primarily perpendicular to this is unaltered. (We are, for the present,
neglecting the 3-dimensional nature of the problem and considering only a plane of iodine molecules). If we assume that \( \beta \) is small, the turning points of the modified bonding orbital, in the \( xz \) plane, are along \( \theta = \phi, \pi + \psi, \pm \theta_o + \chi \) where

\[
\phi = -\frac{\tan \beta}{\cos \gamma} \frac{N_2}{N_1} \frac{(u + a v \sqrt{5})}{(u \cos a + \sqrt{15} v \sin a)}
\]

\[
\psi = -\frac{\tan \beta}{\cos \gamma} \frac{N_2}{N_1} \frac{(u - a v \sqrt{5})}{(u \cos a - \sqrt{15} v \sin a)}
\]

and \( \chi = -\frac{\tan \beta}{\cos \gamma} \frac{N_2}{N_1} \frac{\{3u \cos \theta_o + a \sqrt{15} \cos 2 \theta_o\}}{\{\sqrt{3}u \cos \theta_o \cos \theta_o + 3\sqrt{5} \sin \theta_o \cos 2 \theta_o\}} \)

With some rearrangement, and using the definition of \( \theta_o \), these become:

\[
\phi = -\frac{\tan \beta}{X} \frac{\cos \theta_o (1+A)}{(\cos \theta_o - 1)}
\]

\[
\psi = -\frac{\tan \beta}{X} \frac{\cos \theta_o (1-A)}{(\cos \theta_o + 1)}
\]

\[
\chi = -\frac{\tan \beta}{X} \frac{\cos \theta_o \{\cos \theta_o + A \cos 2 \theta_o\}}{1 - \cos^2 \theta_o} = \frac{\psi \{\cos \theta_o + A \cos 2 \theta_o\}}{(1-\cos \theta_o)(1-A)}
\]

where

\[
X = \cos a \cos \gamma \frac{N_1}{N_2} \quad \text{and} \quad A = a \sqrt{5} \frac{v}{u}
\]

\( \beta \) is positive if the bond along \( \theta_o \) is strengthened, but whether the 180° bond is bent in the required direction depends on whether \( A \) is less than, or greater than, unity. Alternatively we can identify \( \psi \) with its experimental value, 10°53' and \( \theta_o + \chi \) similarly with 105°25' and use the above equations to determine \( \theta_o \) and \( \tan \beta/X \) as functions of \( A \). The results of this are given in Table 1.
Table 1

Values of parameters needed to fit the bond angles

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<td>θ₀</td>
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<td>110°</td>
<td>113.7°</td>
<td>120°</td>
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<tr>
<td>tan β/κ</td>
<td>.425</td>
<td>.457</td>
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The fact that φ is different from zero means that the main iodine bond is not directed straight towards the other iodine, which suggests that the main I-I bond is weakened in the crystal, as required. Further, to have a stable arrangement, the main bonds from the two iodines in a molecule must be deflected on opposite sides of the I-I axis, which means that the midpoint of a molecule is a center of symmetry.

One point which has been glossed over and which must now be considered is the use of the turning values of the angular part of the wave-function to determine bond directions. In so far as the Pauling-directed-bond method is valid, it would seem reasonable to associate bond directions with maxima, but in the above reasoning we have not shown that our various directions, 0, θ₀, and 180° are really directions in which the electron distribution is large. In particular it seems not unlikely that the actual orbits will be such that the charge in the 180° direction is very small, for presumably the bulk of the charge is located in θ = 0. If, then, we assume that the charge concentration is a minimum in θ = 180° we may picture a free iodine molecule as having a lot of charge along the main axis and with subsidiary cones of charge at about 105° with this axis. The crystal is then built up by having a point on the cone of one molecule on the axis of the cone of another. In this way the molecules pack fairly closely without their charges overlapping too strongly, i.e., without the formation of extra bonds. It also seems not unreasonable to suppose that this would lead to planes of fairly closely interlocked iodine molecules, with weaker forces between the planes.

The above ideas are based on physical arguments about charge densities, and it is seen that the introduction of d-hybridization is mainly responsible for the results obtained. It is clearly going to be a formidable task to try and substantiate these ideas by actual calculation of orbits, so we propose instead to
investigate the results of the nuclear quadrupole resonance to see what information this can give.

The Nuclear Quadrupole Interaction

If \((x,y,z)\) is the position of an electron with respect to a nucleus at the origin, the contribution which it makes to the electric quadrupole interaction is:

\[
\frac{e^2 Q}{2I(2I-1)} \left[ \frac{(3x^2 - r^2)}{r^5} I_x^2 + \frac{(3y^2 - r^2)}{r^5} I_y^2 + \frac{(3z^2 - r^2)}{r^5} I_z^2 + \frac{3xy}{r^5} (I_x I_y + I_y I_x) 
\right. \\
+ \left. \frac{3y^2}{r^5} (I_x I_y + I_y I_x) + \frac{3zx}{r^5} (I_x I_z + I_z I_x) \right]
\]

To obtain the full interaction this must be summed over all electrons and averaged over their orbits. Using

\[
\langle z | \frac{3z^2 - r^2}{r^5} | z \rangle = \frac{4}{5} \left( \frac{1}{r} \right)_p \\
\langle z | \frac{zx}{r^5} | x \rangle = \frac{1}{5} \left( \frac{1}{r^3} \right)_p, \quad \langle z^2 - r^2 | \frac{3z^2 - r^2}{r^5} | z^2 - r^2 \rangle = \frac{4}{7} \left( \frac{1}{r} \right)_d \\
\langle xz | \frac{3z^2 - r^2}{r^5} | xz \rangle = \frac{2}{7} \left( \frac{1}{r} \right)_d, \quad \langle xz | \frac{zx}{r^5} | z^2 - r^2 \rangle = \frac{1}{7\sqrt{3}} \left( \frac{1}{r} \right)_d
\]

and other quantities derived from them, the interaction in the present case becomes:

\[
\frac{e^2 Q}{2I(2I-1)} \left[ a_{11} I_x^2 + a_{22} I_y^2 + a_{33} I_z^2 + a_{13} (I_x I_z + I_z I_x) \right]
\]

where

\[
a_{11} = \frac{1}{r^3} \left[ \begin{array}{c}
\frac{4}{N_1} (+ \frac{1}{5} - \frac{1}{7} \lambda a^2) - \frac{2}{N_2} \cos^2 \gamma (\frac{1}{5} \cos^2 a + \frac{1}{7} \lambda \sin^2 a) \\
\sin^2 \gamma \left[ \frac{2}{N_1} \left( \frac{5}{7} \lambda a^2 \right) + \frac{2}{N_2} \cos^2 \gamma \left( \frac{1}{5} \cos^2 a - \frac{1}{7} \lambda \sin^2 a \right) \right]
\end{array} \right]
\]
These formulae are too complicated to be handled easily, so as \( \frac{1}{r^3} \) probably varies as \( t^{-3} \) for electrons with the same principal quantum number, it seems reasonable to approximate by taking \( \lambda \) as zero. Considerable simplification follows if ratios are then used, giving:

\[
\begin{align*}
\mathbf{a}_{11} &= \frac{1 - \frac{1}{2} X^2 - \frac{1}{2} \sin^2 \beta (2 + X^2)}{-2 + X^2 + \frac{1}{2} \sin^2 \beta (1 + 2X^2)} \\
\mathbf{a}_{22} &= \frac{1 - \frac{1}{2} X^2 + \frac{1}{2} \sin^2 \beta (1 - X^2)}{-2 + X^2 + \frac{1}{2} \sin^2 \beta (1 + 2X^2)} \\
\mathbf{a}_{13} &= \frac{3 \cos \beta \sin \beta X}{2 \left[-2 + X^2 + \frac{1}{2} \sin^2 \beta (1 + 2X^2)\right]} \\
\mathbf{a}_{33} &= \frac{1}{\left(-\frac{1}{3}\right)_{p}} \\
\end{align*}
\]
These relations are not independent since \( a_{11} + a_{22} + a_{33} = 0 \), but a full investigation of the quadrupole interaction would determine \( \beta \) and \( \lambda \), and then (see Table 1) \( A \) and \( \theta_0 \), which are properties of the free molecule. There are, however, a number of experimental difficulties in this, so that present information extends only to the magnitude of \( A_{33} \) and the anisotropy \( \left| \frac{A_{11} - A_{22}}{A_{33}} \right| \), where capital \( A \)'s mean that the quadrupole interaction is referred to its principal axes (i.e., \( A_{13} = 0 \)). If we assume that \( a_{13} \) is small so that the original axes are close to the principal axes, the anisotropy is approximately \( \left| a_{11} - a_{22} \right| / a_{33} \)

\[
\frac{3}{2} \sin^2 \beta = \frac{-2 + X^2 + 1/2 \sin^2 \beta (1 + 2X^2)}{1 - 2X^2 + 1/2 \sin^2 \beta (1 + 2X^2)}
\]

To get the bond angles right, we see (Table 1) that \( \tan \beta / X \) is probably of order .45, so taking \( \tan^2 \beta = .18X^2 \) the observed value of .15 for the anisotropy gives \( X^2 \sim .75 \) and \( \tan \beta \sim .37 \). Thus, although the theory contains a good many parameters, some numerical values can be obtained with some precision. Continuing, with \( X^2 = .75 \), \( N_1N_2 \cos \theta \cos \gamma = .866 \), and using \( \cos \theta_0 = -1/\sqrt{15} \), \( u/v \cot a \) and setting \( u = v \), \( \theta_0 = 110^\circ \), we obtain \( a \sim 37^\circ \) and \( N_1N_2 \cos \gamma \sim 1.1 \). If the nonbonding orbits contain no d-wave functions, \( N_1 \) is probably less than \( N_2 \) and this last relation will not be satisfied. It therefore seems that, with our model, it is necessary to have enough d-character in the nonbonding orbits to make \( N_1 > N_2 \), and also we may expect \( \gamma \sim 0 \). With \( a \sim 37^\circ \), \( \gamma \sim 0 \) the bonding orbit contains very little s-character and approximately equal amounts of p and d, which means that \( \theta = 180^\circ \) is a direction in which there is very little charge. In other words, the p and d admixtures are such that the p approximately cancels the d in the \( \theta = 180^\circ \) direction. This situation was anticipated in the discussion earlier in the paper. A more surprising result is that the \( \pi \)-orbitals, i.e., those which contain pairs of electrons moving perpendicular to the axis, have \( \sim 10\% \) d-character (amplitude of admixture, from \( N_1 \)), but on examination this is seen to make the electrons concentrate towards the middle of the molecule, which does not seem unreasonable.

The above numerical estimates were obtained by taking \( \lambda = 0 \) and neglecting \( a_{13} \). If we now use the above numerical values and do not make
these assumptions it is found that the theoretical anisotropy might be reduced to .14, depending on the value of λ. The observed value can be restored by increasing X slightly, a change which will not affect the main conclusions.

Finally, it is of interest to consider the actual magnitude of the quadrupole interaction. Our equations give:

\[
\frac{(a_{33})_{\text{crystal}}}{(a_{33})_{\text{free}} \text{ mol}} \sim \frac{-2 + X^2 + \frac{1}{2} \sin^2 \beta (1+2X^2)}{-2 + X^2} \sim 1 - \frac{1}{2} \sin^2 \beta \left( \frac{1+2X^2}{2-X^2} \right)
\]

\[
\sim .876
\]

The measured value of the quadrupole coupling in the solid is 2140 Mc/s so this ratio predicts that the free iodine value should be 2440 Mc/s.

The other thing which can be done is to compare \((a_{33})_{\text{crystal}}\) with the corresponding value for a free iodine atom, since a free atom will be described by letting \(a = \beta = \gamma = a = 0\) and \(N_1 = N_2 = 1\).

Then

\[
\frac{(a_{33})_{\text{crystal}}}{(a_{33})_{\text{free atom}}} \sim \frac{-2 + X^2 + \frac{1}{2} \sin^2 \beta (1+2X^2)}{-2N_1^2} = \frac{1.23}{N_1^2}
\]

\((a_{33})_{\text{free atom}}\) can be obtained from the analysis of the hyperfine structure of free iodine by Jaccarino [4], whence \(N_1^2 = 1.23 \frac{2292}{2140} = 1.32\). The departure of \(N_1\) from unity gives a measure of the amount of d-character in the nonbonding orbit, and this agrees well with our earlier estimate of this.

**Conclusion**

Starting with elementary ideas about the nature of chemical bonds, it has proved possible to set up a plausible model of a free iodine molecule which enables one to understand how iodine molecules may come together to form a solid having the observed structure. The theory is then tested by examining the predictions which it makes about the nuclear quadrupole interaction. A number of approximations are made in this, but it is found that the anisotropy and magnitude of the quadrupole interaction can be
correlated in a satisfactory way with the crystal structure and wave functions, thus helping to confirm the initial assumptions. The final picture of the origin of the anisotropic quadrupole interactions then differs significantly from the models of Townes and Dailey and of Robinson, Dehmelt and Gordy.

Acknowledgements

I would like to thank Professor R. V. Pound for a number of stimulating discussions and pure quadrupole resonance and O. N. R. for financial support.

References

4. V. Jaccarino, Quarterly Progress Report, Research Laboratory of Electronics, Massachusetts Institute of Technology, July 1953, p. 31.
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