Distributions of random fields in solids: contribution of the nearest defect

A M Stoneham
Theoretical Physics Division, Building 424.4, AERE, Harwell, Oxon OX11 0RA, UK

Received 26 August 1982

Abstract. This paper discusses the statistical distribution of internal fields due to random defects in solids. It extends previous discussions by separating explicitly the contribution of the nearest defect from that due to all others. Possible applications include those to preferential defect pairing, to hopping conduction in randomly doped crystals, and to questions of the existence of phase transitions in random dipole systems. As a by-product, we note that the random strains in solids have a statistically negligible probability of localising a muon (assumed not self-trapped) except at the sites of defects themselves.

1. Introduction

There are many cases in which observable effects arise from the random fields of statistically distributed defects in solids. The fields themselves may be strain fields, electric fields, or it may be the electrostatic potential or electrical field gradient which varies from site to site. This subject was reviewed by Stoneham (1969), who surveyed the methods of calculating the distribution $I(\varepsilon)$ of an internal field $\varepsilon$.

Calculations of the distribution involve two main pieces of information. First, one needs to know the contribution $\varepsilon(z)$ to $\varepsilon$ of a single defect whose relative position and other defining parameters are given by $z$. Secondly, one needs to know something about the statistical distribution. This is usually described by a straightforward generalisation $p(z)$ of the pair distribution function of the defects causing the random field relative to the probe species which monitors $I(\varepsilon)$. Given expressions for $p(z)$, $\varepsilon(z)$ and the density of defects one can calculate the distribution $I(\varepsilon)$. One can also obtain the moments of the distribution $M_N$, where

$$M_N = \int d\varepsilon I(\varepsilon)\varepsilon^N,$$

and some special features like asymptotic expressions. The moments, however, are rarely useful, since they do not always correspond to what is measured, and often diverge, though they are simple to calculate formally.

In this note we look at $I(\varepsilon)$ in some simple cases, and we project out the contribution of the closest defect. This has several advantages. One is that, if the circumstances of direct interest are dominated by the nearest defect, calculations can be made much simpler. A second is that both $p(z)$ and $\varepsilon(z)$ are often described by some simple form at large separations, with corrections at short distances only. In many such cases these
corrections will be important only in the case of the closest defect, and can be handled more easily in the separated form. But the main advantage is conceptual: one can identify directly the contribution of the many distant defects and separate this from the specific contribution of the closest defect. The main complication is, of course, the fact that the distance from a specific site to the nearest defect is not unique, but is itself a distributed quantity. One specific application we have in mind occurs in muon spin rotation. If a muon moves through a lattice containing point defects, and if it localises where the strain field (e.g. the dilatation) is a minimum, will the sites of localisation be at isolated sites (Yaouanc 1982) or only significantly adjacent to the defects themselves? Other potential applications occur in conductivity and in phase transitions (if indeed these occur) in systems containing random or statistically distributed defects.

2. Theory

2.1. The statistical method

Here I simply quote the results derived by Stoneham (1969). The distribution of internal fields given by the statistical method is \( I(\epsilon) \), where:

\[
I(\epsilon) = \frac{1}{2\pi} \int_{-\infty}^{\infty} dx \exp(i\epsilon x) \exp[-\rho J(x)]
\]

and where the density of defects is \( \rho = N/\int dz p(z) \) where \( N \) is the number of defects in the chosen volume of integration. For simplicity we shall usually choose symmetric cases where \( I(\epsilon) \) and \( I(-\epsilon) \) are equal. The expressions then simplify:

\[
I(\epsilon) = \frac{1}{\pi} \int_0^{\infty} dx \cos(x\epsilon) \exp[-\rho J(x)]
\]

\[
J(x) = \int dz p(z) \{1 - \exp[-i\epsilon z(z)]\}
\]

where \( J(x) \) is now real.

In the very simplest and most important cases, \( \epsilon \) depends only on the relative position of the perturber. If the perturbing defects are distributed completely at random, then \( p(z) \) \( dz \) takes the following forms:

\[
p(z) \ dz \rightarrow dr \quad (1 \text{ dimension})
\]

\[
\rightarrow r \ dr \ d\theta \quad (2 \text{ dimensions})
\]

\[
\rightarrow r^2 \ dr \ d\Omega \quad (3 \text{ dimensions}).
\]

Here \( \int d\theta = 2\pi \) and \( \int d\Omega = 4\pi \) when integrated over angles. If the distribution is not random, \( p(z) \) is multiplied by a weighting factor \( f(r) \). With the definitions (5) the density of defects, \( \rho \), is the number per unit length (one dimension), per unit area (two dimensions) or per unit volume (three dimensions).

2.2. The nearest defect

Here we follow the early arguments of Hertz (1909) and Chandrasekhar (1943). The probability \( P_N(r) \) that the nearest defect lies between \( r \) and \( r + dr \) of a chosen site is the
product of two probabilities: $\hat{p}(r)$, which is the probability that there is a defect in this range, and $q(r)$, the probability that there is no defect closer than $r$:

$$P_N(r)\, dr = \hat{p}(r)q(r)\, dr.$$  

(6)

The two component probabilities are related, for $q(r)$ is clearly the product of all the component probabilities $[1 - \hat{p}(x)\, dx]$ that there is no defect between $x$ and $x + dx$ for $0 \leq x \leq r$. Hence (either directly or following the discussions of Hertz and of Chandrasekhar) one can show:

$$P_N(r)\, dr = \exp\left(-\int_0^r \hat{p}(x)\, dx\right)\hat{p}(r)\, dr.$$  

(7)

In general $\hat{p}$ is a function of $z$, and inspection shows that $\hat{p}(z)$ of the present section is related to $p(z)$ by the simple factor $\rho$, the density of defects, plus a second factor to take into account internal degrees of freedom, if any. An integral over angles is implied too if we are only concerned with distance, irrespective of direction. In the important case described by (5) we find, for example,

$$\hat{p}(x)\, dx = \rho 4\pi\, dx\, x^2$$  

(8)

in three dimensions, whence the Hertz-Chandrasekhar result:

$$P_N(r)\, dr = \rho \exp(-\frac{4\pi r^3}{3})4\pi r^2\, dr.$$  

(9)

It is useful to give results for one, two and three dimensions. Whilst Hertz quotes most probable, mean and median nearest-neighbour distances in each case, the distributions are implicit only. Table 1 quotes the results, apparently given here for the first time, and introduces simplified notation in terms of $L$, the average nearest-neighbour distance.

<table>
<thead>
<tr>
<th>Form of $p(z), dz$ integrated over angles</th>
<th>One dimension</th>
<th>Two dimensions</th>
<th>Three dimensions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density of defects $\rho$</td>
<td>$\rho_1$ = number per unit length</td>
<td>$\rho_2$ = number per unit area</td>
<td>$\rho_3$ = number per unit volume</td>
</tr>
<tr>
<td>$L, \rho_1 = 1$</td>
<td>$2\pi r, dr$</td>
<td>$\frac{4\pi r^2}{3}, dr$</td>
<td></td>
</tr>
<tr>
<td>Form of $P_N(r), dr$</td>
<td>$\rho_1 \exp(-\rho_1 r)$</td>
<td>$\rho_2 \exp(-\rho_2 r)^2$</td>
<td>$\rho_3 \exp(-\frac{4\pi r^3}{3})$</td>
</tr>
<tr>
<td>Form of $P_N(y), dy$ with $y = r/L$</td>
<td>$\rho_1 \exp(-y)$</td>
<td>$\rho_2 \exp(-y^2)$</td>
<td>$\rho_3 \exp(-\frac{4\pi r^3}{3})$</td>
</tr>
</tbody>
</table>

It may happen that the defects are not distributed completely at random. Two simple examples arise when there are attractive interactions, giving preferential pairing, and site exclusion, since two defects cannot occupy the same lattice site. If the effect of the interaction is to modify the pair distribution function by $f(r, \Omega)$, where $f \to 1$ at large distances to ensure the average density $\rho$ is defined consistently, then the main change is the replacement

$$\hat{p}(r) \to \hat{p}(r) \int d\Omega f(r, \Omega).$$  

(10)

This leads to more complexity, but does not cause any serious problems.
2.3. Contribution of the nearest defect

The main idea exploited in the next section is that the contribution of the nearest defect to the distribution of $\varepsilon$ is $I_N(\varepsilon)$ where

$$I_N(\varepsilon) \, d\varepsilon = \text{probability that the nearest defect lies between the contours on which it produces perturbations } \varepsilon \text{ and } \varepsilon + d\varepsilon.$$  \hspace{1cm} (11)

Thus, if $\varepsilon(z)$ is a function of $r$ only, and if $dR$ is defined by $dR = d\varepsilon(\varepsilon(r)/dr)$ and $R$ by $\varepsilon = \varepsilon(R)$, then we find

$$I_N(\varepsilon) \, d\varepsilon = P_N(R) \, dR.$$  \hspace{1cm} (12)

The only technical problem is the inversion of $\varepsilon = \varepsilon(r)$ to give the contours $R = R(\varepsilon)$ over which the defect would give the same perturbation $\varepsilon$. We note that $I_N(\varepsilon)$ itself is not necessarily normalised; it is $P_N(R)$ which is normalised.

For reference, it is useful to define the several distributions needed here: $I(\varepsilon) = \text{probability that the internal field from all defects lies between } \varepsilon \text{ and } \varepsilon + d\varepsilon$; $I_N(\varepsilon) = \text{probability that the nearest defect contributes an internal field between } \varepsilon \text{ and } \varepsilon + d\varepsilon$; $I_0(\varepsilon) = \text{probability that the internal field from all defects except the nearest lies between } \varepsilon \text{ and } \varepsilon + d\varepsilon$.

2.4. Contribution of all defects but the nearest

The full distribution $I(\varepsilon)$ is simply the convolution of $I_N(\varepsilon)$ and $I_0(\varepsilon)$, a second distribution which comes entirely from the defects other than the nearest. The simplest way to obtain $I_0(\varepsilon)$ is from the convolution theorem. Expressing the Fourier transforms of $I$, $I_N$ and $I_0$ as $\hat{I}$, $\hat{I}_N$ and $\hat{I}_0$ respectively, $\hat{I} = \hat{I}_N \hat{I}_0$. The desired distribution $I_0(\varepsilon)$ can be found from the back-transform of $\hat{I}/\hat{I}_N$. In practice, this is complicated. However, it is clear that $I_N(\varepsilon)$ dominates at the large values of $\varepsilon$, whilst $I_0(\varepsilon)$ is the major contributor at small $\varepsilon$, falling rapidly to zero as $\varepsilon$ rises.

3. Examples of $I_N(\varepsilon)$

We now calculate the distributions of random fields due to the nearest defect only, beginning with simpler cases and generalising systematically.

Case I. Consider the simple inverse power-law potential given by

$$\varepsilon(z) = A/r^m.$$  \hspace{1cm} (13)

The inversion of (13) gives these expressions:

$$r = A^{1/m}e^{-1/m} \hspace{1cm} (14)$$
$$dr = -(1/m)A^{1/m}e^{-(1-1/m)} \, d\varepsilon. \hspace{1cm} (15)$$

Equation (12) now gives us

$$I_N(\varepsilon) \, d\varepsilon = P_N\left[\frac{A}{\varepsilon}\right]^{1/m} \frac{A^{1/m}}{m} e^{-(1-1/m)} \, d\varepsilon$$

where the irrelevant negative sign has been dropped. This equation is general, applying to all distributions of defects, and so to all forms of $P_N(r)$. We illustrate the result using
the simplest case of all, namely a random distribution. The results in one, two and three dimensions are most conveniently expressed in terms of $\varepsilon_L$:

$$\varepsilon_L = A/L^m$$  \hspace{1cm} (17)

the perturbation due to a defect at the average spacing, $L$. Using table 1, we obtain:

$$I_N(\varepsilon) \, d\varepsilon = \begin{cases} 
(1/m)(\varepsilon_L/e)^{1/m-1} \left[ \exp(\varepsilon_L/e)^{1/m} \right] d(\varepsilon/e_L) \\
(2/m)(\varepsilon_L/e)^{2/m-1} \exp \left[ - (\varepsilon_L/e)^{2/m} \right] d(\varepsilon/e_L) \\
(3/m)(\varepsilon_L/e)^{3/m-1} \exp \left[ - (\varepsilon_L/e)^{3/m} \right] d(\varepsilon/e_L)
\end{cases}$$

\hspace{1cm} (18a, 18b, 18c)

The results at large perturbations (large $\varepsilon$) reduce to the asymptotic solutions given in § 4.5 of Stoneham (1969). At the small-perturbation limit (small $\varepsilon$), $I_N(\varepsilon)$ tends to zero very rapidly. This corresponds to the obvious point that it is rare for the nearest neighbour to be very far away.

**Case II.** In case I, the perturbation always had the same sign: $\varepsilon(z)$ of (13) is always positive. We now consider a case in which there are two types of defect which produce perturbations of opposite signs. One, with density $\rho_+$, gives a perturbation $\varepsilon_+ = Ar^{-m}$; the other, with density $\rho_-$, gives a perturbation $\varepsilon_- = -Ar^{-m}$. If the defects are again randomly distributed, then the resulting distribution is simply the weighted mean of the two components:

$$I_N(\varepsilon) \, d\varepsilon = d\varepsilon \left( \rho_- I_+(\varepsilon) + \rho_- I_-(\varepsilon) \right) / (\rho_- + \rho_-).$$

\hspace{1cm} (19)

The two components are quite distinct here, giving opposite sign contributions to $\varepsilon$.

When $\rho_-$ and $\rho_-$ are equal, $I_N(\varepsilon)$ becomes symmetric, i.e. $I_N(\varepsilon)$ and $I_N(-\varepsilon)$ are identical. In this limit we may usefully compare $I_N(\varepsilon)$ with the full distribution $I(\varepsilon)$ from all defects present. In the same notation we have for the general case:

$$\rho I(x) = 6 \int dy \, y^2 \left[ 1 - \cos(x \varepsilon_L y^{-m}) \right] + 6ix \int dy \, y^2 \sin(x \varepsilon_L y^{-m})$$

\hspace{1cm} (20)

where $\varepsilon_L = AL^{-m}$ and $4\pi L^3 \rho = 1$, with $\rho = \frac{1}{2} (\rho_+ + \rho_-)$ and $\alpha = (\rho_- - \rho_+)/ (\rho_- + \rho_-)$. For the specific case $\rho_+ = \rho_-$ we find:

$$I(\varepsilon) = \frac{1}{\pi} \int_0^\infty dx \, \cos(x \varepsilon) \exp(-x^{3m} \varepsilon_L^{3m} \tilde{J}_m)$$

\hspace{1cm} (21)

$$\tilde{J}_m = \frac{6}{m} \int_{u_1}^{u_2} du \, \frac{1 - \cos u}{u^{1+3/m}}.$$  \hspace{1cm} (22)

In practice the lower limit (strictly $u_1 = x \varepsilon_{\text{min}}$, where $\varepsilon_{\text{min}}$ is the perturbation of the most distant allowed defect) can be taken as zero. Likewise, the upper limit (strictly $x \varepsilon_{\text{max}}$, where $\varepsilon_{\text{max}}$ is the perturbation of the closest permitted defect) may normally be taken as infinite. The dependence of $\tilde{J}_m$ on $x$ is thus modest. Provided this dependence can be ignored, $\tilde{J}_m$ is just a number. So long as $m$ is finite and bigger than $3/2$, the integral converges satisfactorily. In the special case $m = 3$, $\tilde{J}_m$ is simply $\pi$. The expressions take a rather simple form:

$$I(\varepsilon) \, d\varepsilon \rightarrow \frac{\varepsilon_L^2}{\varepsilon_L^2 + \varepsilon} \, d(\varepsilon/\varepsilon_L)$$

\hspace{1cm} (23)

$$I_N(\varepsilon) \, d\varepsilon \rightarrow (\varepsilon_L/e)^2 \exp(-\varepsilon_L/e) \, d(\varepsilon/e_L).$$  \hspace{1cm} (24)
This confirms the point already made that $I(\varepsilon)$ tends to $I_N(\varepsilon)$ at large $y$. Close inspection shows a less obvious feature: $I_N$ can exceed $I$, though the difference falls to zero as $\varepsilon$ increases. If one writes $|\varepsilon| = \pi \varepsilon_L y$, for example,

$$I_N(\varepsilon)/I(\varepsilon) = (1 + y^{-2}) \exp(-1/\pi y),$$

then $I_N(\varepsilon)$ is 45% bigger than $I$ at $y = 1$, the point of half maximum. The underlying reason is that $I_N(\varepsilon)$ singles out the nearest defect which gives a specific sign of perturbation, whereas $I(\varepsilon)$ is a compromise between the contributions of the two species with opposite signs.

In the case described by (23) and (24) we can also calculate the distribution $I_0(\varepsilon)$ from defects other than the nearest. The cosine transforms of (23) and (24) yield:

$$I(x) = \frac{1}{2} \exp(-\pi x)$$

$$I_N(x) = -2\sqrt{x} \ker'(2\sqrt{x})$$

with $\ker'(y)$ as defined in Abramowitz and Stegun (1965), § 9.9. The reverse transform must be carried out numerically, and gives the results shown in figure 1.

**Figure 1.** The three distributions shown are for a $1/r^3$ interaction with defects of both signs. $I(\varepsilon)$ is the full distribution, $I_N(\varepsilon)$ that from the nearest defect, and $I_0(\varepsilon)$ from all defects but the nearest. The numerical inversion for $I$ is difficult technically, so that, whilst the resulting curve is basically correct, it may be inaccurate in detail.

**Case III.** Assume (13) is replaced by some other monotonic power law $\varepsilon = \varphi(r)$, so that $\varphi^{-1}(\varepsilon)$ gives a single value $r$. Define $\psi(\varepsilon) = d\varphi^{-1}(\varepsilon)/d\varepsilon$. Then we have as the analogue of (16):

$$I_N(\varepsilon) \, d\varepsilon = P[\varphi^{-1}(\varepsilon)] \psi(\varepsilon) \, d\varepsilon.$$  

(28)
Distributions of random fields in solids

Thus, if we were to assume
\[ \epsilon(r) = A \exp(-\beta r) \]  
and to define \( \epsilon_L = \epsilon(L) \) again, we should find:
\[ I_N(\epsilon) = (3/\beta \epsilon) \left[ \ln(\epsilon_{ij}/\epsilon) \right]^3 \exp - \left[ \ln(\epsilon_{ij}/\epsilon) \right]^3. \]

4. Discussion

4.1. Possible applications

The discussion so far has aimed mainly at clarifying contributions to random internal strain fields. In fact, such a division into components is useful in several applications. These include hopping conduction in randomly doped crystals, and certain ordering transitions for random dipolar impurities.

Calculation of hopping conduction in randomly doped insulators and semiconductors involves two configuration-dependent factors. One concerns the overlap of states associated with those nearby pairs of the dopants between which hops occur. This leads to a transition matrix element \( M_{ij} \), falling off roughly exponentially with separation \( R_{ij} \). The other factor involves the energy difference \( \Delta E_{ij} \) between the initial and final states. This term varies exponentially with inverse temperature. The usual analyses (Mott and Davis 1971, § 2.9) then give the limiting cases of variable-range hopping, of Miller and Abrahams (1960) and so on.

Even for a given spacing \( R_{ij} \), both \( M_{ij} \) and \( \Delta E_{ij} \) are distributed quantities. For amorphous systems, the structural disorder is important. In such cases an empirical distribution, constant near \( E_F \), is usually assumed. For randomly doped crystalline systems, the dopant–dopant interactions may be the major factor in the spread of \( M_{ij} \) and \( \Delta E_{ij} \). If so, the distributions of \( M_{ij} \) and of \( \Delta E_{ij} \) can be obtained using the arguments of § 2.4. In principle, one can predict both the absolute value of the density of states and the qualitative shape of the distribution; this may permit a quantitative prediction of the hopping conductivity which improves on some current models.

Cooperative phenomena involving randomly distributed dipoles have been observed in many cases. With elastic dipoles one sees martensitic transitions (see e.g. Nishiyama 1978); for electric, or electric and elastic, dipoles one finds systems like alkali halides doped with \( \text{OH}^- \) or \( \text{CN}^- \) (see e.g. Potter and Anderson 1981, or Garland et al 1982). One feature is that a fairly clear phase transition is observed only above a critical concentration. This can be understood in several ways. Stoneham and Bullough (1970) noted that the mean field driving alignment was much less than the fluctuations in the aligning field. This means, of course, that below the critical concentration, many dipoles are ‘frozen out’ in small clusters, each with their own high ordering temperature. Since the closest defects interact most strongly, and since pairs are the smallest clusters (and possibly the only ones of importance if the critical concentration is low) one may use generalisations of our present results to estimate what the critical concentration is. This involves the self-consistent solution of two equations. One relates the mean field causing alignment \( \bar{\epsilon} \) to the fractional concentration of dipoles not frozen out. If \( c \) is the total concentration and \( c_\ast \) is that concentration frozen out, \( \bar{\epsilon} \) is roughly proportional to \((c - c_\ast)\). The second equation defines \( c_\ast \) as the concentration fraction for which the local field (e.g. that of the nearest defect) exceeds the mean field \( \bar{\epsilon} \) by some specified amount. Thus \( \bar{\epsilon} = \alpha(c - c_\ast) \) and \( c_\ast = \int d\epsilon P(\epsilon) \theta(\epsilon - \bar{\epsilon}) \) in which \( \theta \) might be the Heav-
inside function. In principle, these might be solved to deduce a concentration below which no phase transition is expected. In practice some generalisation is needed, since site exclusion becomes important at about the concentrations of interest.

4.2. Related distributions

The distributions we have calculated are those of a scalar \( \varepsilon \). Frequently, the variable of interest is a vector (like an electric field) or a tensor (e.g. a strain). There are then two types of distribution: one is the projection of the vector onto a specific direction, i.e. \( \varepsilon_p = \sum a_i \nu_i \), with \( \sum |a_i|^2 = 1 \), the other the magnitude of the vector irrespective of direction, i.e. \( \varepsilon_m = (\sum |\nu_i|^2)^{1/2} \). Here \( \varepsilon \) is a unit vector; there are obvious generalisations for tensor quantities. Chandrasekhar (1943, §IV.2) shows that the two distributions are related by:

\[
I_m(\varepsilon) = 4\pi\varepsilon^2 I_p(\varepsilon)
\]

in three dimensions. This has one obvious consequence: unless the distribution of the projection \( \varepsilon_p \) is singular (\( I_p(\varepsilon) \sim \varepsilon^{-2} \) at least) then \( I_m(\varepsilon_m) \) tends to zero with \( \varepsilon_m \): there are essentially no sites at which the magnitude \( \varepsilon_m \) of the vector is zero. There are, of course, the actual defect sites themselves, though usually these are excluded from the counting because the expressions for \( \varepsilon(z) \) are singular when extrapolated to defect sites.

4.2. Are there minima in \( \varepsilon \) other than at defects?

In muon spin rotation one needs to know whether there are sites remote from defects where the muon will find an energy minimum. Suppose the energy of interaction with a defect is \( U(r) \). Then we may seek minima in \( \sum U_i \) by looking at the distribution of forces \( \nabla U_i = F \). The minima correspond to zeros in \( |F| \). We can see immediately from (31) and from the examples of §3 (and those given by Chandrasekhar (1943) or by Stoneham (1969)) that absolute minima in \( U \) (zeros in \( |F| \)) are statistically negligible in common random systems. This excludes (a) minima at defects, (b) the statistically negligible special cases where \( |F| \) vanishes for fortuitous reasons, and (c) special potentials \( U(r) \) which are constant over finite volumes. We can also see from our earlier calculations here that it is the nearest defect which dominates in the force field. One may assume that the muon will move (normally by diffusion) rapidly to a nearby defect; indeed, our calculations suggest it will be the nearest defect with high probability. Trapping in isolation, other than self-trapping, is unlikely unless the strain field of an individual defect (e.g. the nearest) has large volumes of constant strain; surfaces of constant strain do not suffice.

Acknowledgments

I am most grateful to Dr J Rae for the transform giving equation (27) and to Dr J A G Temple for figure 1 and the numerical analysis on which it is based.

References

Abramowitz M and Stegun I A 1965 *Handbook of Mathematical Functions* (New York: Dover)
Distributions of random fields in solids

Chandrasekhar S 1943 Rev. Mod. Phys. 15 1
Miller A and Abrahams E 1960 Phys. Rev. 120 745
Stoneham A M 1969 Rev. Mod. Phys. 41 82
Yaouanc A 1982 Phys. Lett. 87A 423