Density of States and Conductivity of Granular Metal or Array of Quantum Dots

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The conductivity of a granular metal or an array of quantum dots usually has the temperature dependence associated with variable range hopping within the soft Coulomb gap of density of states. This is difficult to explain because neutral dots have a hard charging gap at the Fermi level. We show that uncontrolled or intentional doping of the insulator around dots by donors leads to random charging of dots and finite bare density of states at the Fermi level. Then Coulomb interactions between electrons of distant dots results in the a soft Coulomb gap. We show that in a sparse array of dots the bare density of states oscillates as a function of concentration of donors and causes periodic changes in the temperature dependence of conductivity. In a dense array of dots the bare density of states is totally smeared if there are several donors per dot in the insulator.

I. INTRODUCTION

Conduction of samples where metallic granules are surrounded by some kind of insulator (granular metals) have been studied intensively for decades\(^1\)–\(^9\). If volume fraction of the metal \(x\) is large, metallic granules touch each other and conductivity is metallic. When \(x\) decreases and crosses percolation threshold \(x_c\), granules become isolated from each other and granular metal goes through metal-insulator transition. It is generally observed\(^1\)–\(^3\) that at the insulator side of transition the temperature dependence of conductivity obeys

\[
\sigma = \sigma_0 \exp \left[ - \left( \frac{T_0}{T} \right)^{1/2} \right].
\]

Recently, similar temperature dependence was observed in doped systems of self-assembled germanium quantum dots on the silicon surface\(^10\) and CdSe nanocrystal thin films\(^11\). Below we talk about both granular metal and semiconductor dot structures universally using the word dot for brevity.

Efros-Shklovskii (ES) law. In a \(n\)-type lightly doped compensated semiconductor all acceptors are negatively charged and randomly situated, an equal number of donors are charged positively. Together all random charges create a random potential shifting donor levels up and down. This results in finite bare DOS \(g_0(\mu)\) at the Fermi level \(\mu\). Long range Coulomb interaction of localized electrons then produces the parabolic Coulomb gap in DOS

\[
g(\varepsilon) = \frac{3}{\pi} \kappa^3 \varepsilon^2 / e^6
\]

at the Fermi level (Fig. 1) leading to ES law with \(T_0 = C \cdot e^2 / \kappa \xi\), where \(C\) is a constant factor, \(\kappa\) is dielectric constant and \(\xi\) is the localization length.

FIG. 1. The shape of Coulomb gap in the vicinity of the Fermi level. Bare density of states in the absence of long range Coulomb interaction is shown by the dashed line. Occupied states are shaded.

In contrary to a doped semiconductor, in an array of neutral dots the bare DOS at the Fermi level \(g_0(\mu) = 0\) and there is no justification for ES law, which requires a non-zero \(g_0(\mu)\) to begin with. Let us consider a sparse array of dots with the same radius \(R\) shown on Fig. 2. Charging energy levels of such array are shown in Fig. 3. Here we assume that dots are large enough, so that spacing between charging levels \(e^2 / \kappa R\) is much greater than spacing between quantum levels of the dot with a given charge. Empty peaks in Fig. 3 correspond to energies necessary to charge a neutral dot by the first, second and so on electrons transferring them from the macroscopic piece of the same metal. Shaded (filled) peaks correspond
to, taken with the sign minus, energies necessary to extract first, second and so on electrons from the dot, or in other words, this is density of states of holes. The Fermi level of the array is at zero between two peaks and coincides with the Fermi level of macroscopic piece of the same metal.

\[ \mathcal{E} \]

\[ \begin{array}{c|c}
3 e^2 & \text{occupation}
\end{array} \]

\[ \begin{array}{c|c}
2 k R & \text{energy}
\end{array} \]

\[ \begin{array}{c|c}
e^2 & \text{probability}
\end{array} \]

\[ \begin{array}{c|c}
\mu & \text{charge}
\end{array} \]

\[ \begin{array}{c|c}
- e^2 & \text{occupation}
\end{array} \]

\[ \begin{array}{c|c}
- 3 e^2 & \text{energy}
\end{array} \]

\[ \begin{array}{c|c}
2 k R & \text{occupation}
\end{array} \]

\[ \begin{array}{c|c}
g_0(\mathcal{E}) & \text{probability}
\end{array} \]

FIG. 3. The bare density of ground states (BDOGS) of a clean system of identical dots consists of equidistant peaks. Occupied states are shaded.

We want to emphasize that in each dot we are dealing with the ground state at a given number of electrons and exclude excited states (higher quantum levels) because the ground states of dots determine the low temperature hopping transport. Indeed, in the Miller-Abrahams network\textsuperscript{13} of resistances connecting all dots, the exponentially large activation factor of each resistance depends on probabilities of occupation of a dot by a given number of electrons. Exponential temperature dependencies of these probabilities, as well as the partition functions of dots are determined by ground state energies\textsuperscript{14}. Thus, the density of states of the array we need can be called bare density of ground states (BDOGS).

Note that in a lightly doped semiconductor with several equivalent conduction band minima each donor has excited states close to the ground state, i.e. situation is similar to large dots. Still exponential temperature dependence of hopping conductivity depends only on BDOGS\textsuperscript{14,13}.

Let us illustrate the role of BDOGS in a clean sparse array (Fig. 3). In this case, conductivity requires activation of electron-hole pairs or in other words, transfer of an electron between two originally neutral dots. Obviously concentrations of positive and negative dots and the hopping conductivity obeys

\[ \sigma = \sigma_0 \exp[-E_c/2k_BT] \quad (3) \]

independently of excited states, (Same result can be obtained in Miller-Abrahams resistor network approach.) where \( E_c = e^2/\kappa R \) is the charging energy.

How then can we explain observation of ES law? Apparently BDOGS shown in Fig. 3 should be smeared in the vicinity of the Fermi level due to some kind of disorder.

A simplest source of disorder is distribution of sizes or capacitances of dots. Indeed, charging energies of larger dots are smaller and this can result in the low energy tail of the first empty peak of BDOGS and symmetric high energy tail of the first occupied peak. Still in a neutral system these tails do not overlap and \( g_0(\mu) \) is zero, so that this kind of disorder does not lead to ES law. Sheng et al.\textsuperscript{3} assumed that in a reasonably dense system of neutral granules there is a special distribution of distances between granules or their mutual capacitances, which can lead to ES law. This assumption was found incompatible with other experiments\textsuperscript{5}. But more importantly it was noticed\textsuperscript{6} that in a system of large granules made of the same metal all granules remain neutral in ground state at any distribution of mutual and individual capacitances. Therefore, inter-granular excitation of electron-hole pairs has a gap.

This leads to an important conclusion that granular metal may have finite BDOGS \( g_0(\mu) \) and show ES conductivity only if in the ground state of the system granules are charged\textsuperscript{4–9}. Formally one can imagine that this happens if granules have different work functions\textsuperscript{4,6}. Several possible mechanisms of such fluctuations were discussed in literature. Cuevas et al.\textsuperscript{8} claimed that even in large dots there are large fluctuations of the Fermi level \( \delta E_F \) due to random positions of neutral impurities in different dots. To our mind, this possibility can be rejected using for \( \delta E_F \) a simple estimate of typical fluctuation of the average potential in the metallic dot due to fluctuation of number of impurities there. For three dimensional case one easily gets \( \delta E_F \sim E_F/c^{1/2}/(k_F R)^{3/2}, \) where \( c \) is relative concentration of impurities in the dot, \( k_F \) is Fermi wave-vector, and we assume that size of an impurity is of the order of \( k_F^{-1} \) and its potential \( U \sim E_F. \) At large \( R \) this energy is apparently smaller than charging energy \( e^2/\kappa R \) even at \( c \sim 1, \) so that dots remain neutral. Baskin and Entin\textsuperscript{9} considered fluctuations of surface part of the dot energy as a reason of ionization of dots. Again if we assume that these fluctuations are result of random distribution of point-like impurities located on the dot surface we come to conclusion that corresponding fluctuation of energy decreases like \( 1/R^2 \) and can not compete with the charging energy. Thus, simplest internal mechanisms of fluctuations of the work function mentioned above are too weak to charge array of three-dimensional dots.

In this paper we study models of arrays of dots affected by external disorder, where the origin of charging and BDOGS \( g_0 \) is transparent and in some cases controllable. We assume that the insulator between clean dots has a concentration of donors \( N_D \) with electron energy \( E_D \) higher than Fermi energy \( \mu \) in dots, so that at low temperatures all donor donate an electron to dots and charge them. For a given concentration of dots \( N \)
(Fig. 2) we can introduce the average number of extra electrons per dot \( \nu \equiv N_D/N \), which by analogy with quantum Hall effect can be called the filling factor. Maximum number of electrons which can be added to a dot is \( n_{\text{max}} \approx (E_D - \mu)/(e^2/\kappa R) \). We assume that \( \nu < n_{\text{max}} \) so that all donors lose their electrons. In the semiconductor language we are dealing with compensated p-type semiconductor where dots play the role of multi-charged acceptors.

In Sec. II we summarize our main results for BDOGS at the Fermi level \( g_0(\mu) \) as function of the filling factor \( \nu \). In Sec. III the sparse 3D array of dots (Fig. 2) is discussed. In Sec. IV we extend this discussion to 2D array. In Sec. V we study in detail the super-dense 3D array shown in Fig. 4, which is the other extreme, and then comment on the realistic moderately dense array (Fig. 14). We also see how the properties of sparse 3D arrays cross over to dense arrays. After studying different arrays one can see that external doping brings about non-zero \( g_0(\mu) \) and leads to ES law.

**II. SUMMARY OF RESULTS**

We start from sparse three dimensional arrays of dots of the same radius \( R \), which are randomly situated in space with concentration \( N \ll (4\pi R^3/3)^{-1} \). We assume that the dots are big enough so that Coulomb effect overweights quantum level spacing. For sparse arrays of dots, doping introduces two types of charges: positive, empty donors and negatively charged dots. Both are randomly situated and create random potentials growing with \( \nu \). These charges result in two effects (Fig. 5). Firstly, the \( \delta \)-shaped peaks in BDOGS \( g_0(\varepsilon) \) become somewhat smeared, since the energy it takes to bring an electron to a dot is affected by the random potential (the effect is similar to that of gedanken random gate potential). As a result, each peak gets tails. Secondly, electrons coming from donors fill some dot states and hence move the position of the Fermi level up.

As a result \( g_0(\mu) \) may oscillate with \( \nu \). For example, at \( \nu = 1/2 \) the Fermi level \( \mu \) is in the middle of the BDOGS peak and for that reason \( g_0(\mu) \) can be rather large. On the other hand, at \( \nu = 1 \) the Fermi level \( \mu \) is in a tail between two BDOGS peaks, and \( g_0(\mu) \) tends to be much lower. Actually the total dependence of \( g_0(\mu) \) on \( \nu \) is somewhat more complicated and consists of three parts, the growing part, the oscillating part and the saturation part (Fig. 6). The first (growing) part takes place at small \( \nu \), where the situation is similar to \( p \)-type semiconductors at low degree of compensation\(^1\). We get BDOGS growing linearly with \( \nu \)

\[
g_0(\mu) \sim \kappa \nu N^{2/3}/e^2 \quad (\nu << 1). \tag{4}\]

In the second (oscillating) part, \( \mu \) typically dwells in BDOGS peaks,

\[
g_{\text{max}}(\mu) \sim \frac{N^{2/3} \kappa}{e^2 \nu^{1/3}} \quad (1 < \nu < \nu_s). \tag{5}\]

When \( \nu \) is very close to integers, the Fermi level drop into minima of \( g_0(\varepsilon) \) where

\[
g_{\text{min}}(\mu) \sim \frac{\nu^{8/3} N^{5/3} R^2}{e^2/(\kappa R)} \quad (1 < \nu < \nu_s). \tag{6}\]

Here \( \nu_s \sim 1/(NR^3)^{1/4} >> 1 \) is the filling factor at which oscillations become relatively small and the third (saturation) part starts over. This happens because fluctuations of the Coulomb potential are so big that BDOGS is almost uniform everywhere

\[
g_0 \sim \frac{N}{e^2/(\kappa R)} \quad (\nu > \nu_s). \tag{7}\]

The number of large oscillations \( \nu_s \) is big only when dots are far from each other. Here and below we often omit numerical coefficients.
For less ideal arrays where the dots have slightly different sizes, the distribution of sizes can wash away the oscillations to a certain extent, making the line of maxima lower and the line of minima higher. As a result the two lines approach each other faster ($\nu_s$ is smaller) than the situation of identical sizes.

Oscillations of $g_0(\mu)$ lead to periodic transitions between the ES law and the Mott’s law at a given low temperature as shown in Fig. 7. This happens because in the very vicinity of $\mu$ the long range Coulomb interaction creates the parabolic Coulomb gap (not shown in Fig. 5). The width of Coulomb gap as follows from Eq. (2) and Fig. 1 depends on $g_0$

$$\Delta \sim \sqrt{g_0(\mu)e^6/\kappa^3}. \quad (8)$$

At large $g_0$ the Coulomb gap is wide ($\Delta >> k_BT$) and the conductivity obeys the ES law (Eq. (1)), which does not depend on $g_0$. At a very small $g_0$ the width of the Coulomb gap is smaller than $k_BT$ and becomes irrelevant for conductivity. In this case we arrive at the Mott’s law region

$$\sigma = \sigma_0 \cdot \exp \left[ -\left( \frac{\beta}{g_0(\mu) a^3 T} \right)^{1/4} \right], \quad (9)$$

with strong dependence on $g_0(\mu)$. Here $\beta$ is a numerical coefficient$^4$. Oscillations of BDOGS $g_0(\mu)$ with $\nu$ lead to oscillations of other measurable quantities. It was shown$^{10,18}$ that in systems with the Coulomb gap BDOGS $g_0(\mu)$ plays the role of the thermodynamic density of states $dn/d\mu$, where $n$ is concentration of electrons, so that it determines screening radius of the system $r_s = (4\pi g_0 e^2/\kappa)^{-1/2}$ used below for self-consistent calculations of BDOGS. One can also measure BDOGS at the Fermi level with help of extremely low temperature (lower than typical quantum level spacing) specific heat and microwave absorption$^4, 17$

FIG. 6. BDOGS at the Fermi level $g_0(\mu)$ of a sparse 3D array as a function of filling factor (solid line). The reference to the equation appropriate for a part of the curve is shown next to it. Dashed lines describe locations of minima and maxima of the oscillating part.

FIG. 7. Ranges of ES and Mott’s laws alternating with growth of filling factor $\nu$ at a given low temperature.

Until now we talked about 3D arrays of dots. In Sec. IV we consider arrays of dots located in a 2D plane. In a 2D array dots can be charged by donors located in parallel to the plane (δ-layer)$^{10,18}$. In 2D case, however, there is a more practical way to charge dots using a metallic gate parallel to the plane of dots. At $\nu >> 1$ results for $g_0(\mu)$ are almost independent on the way of charging, because most of the random potential is created by dot charges. Dependence of $g_0(\mu)$ on $\nu$ in 2D qualitatively looks similar to the 3D case discussed above, but quantitatively it is somewhat different (see Sec. IV).

Dependence of $g_0(\mu)$ on $\nu$ in 2D should lead to oscillations of conductivity. Some oscillation were observed in experiments$^{10,18}$. But there are additional ways to measure this dependence in a gated in 2D structure. First$^{10}$, one can study change of conductivity where the gate voltage charges dots. Second$^{19}$, one can study inverse small signal AC capacitance of unit area $1/C$, which is related to the two-dimensional screening radius of the dot system $r_s = \kappa/(2\pi g_0 e^2)$ by equation

$$1/C = 4\pi \kappa (d_0 + r_s), \quad (10)$$

where $d_0$ is distance from gate to the plane of dots. This method is similar to magneto-capacitance measurements in quantum Hall effect$^{20}$.

Until now we talked about sparse arrays of dots. For a sparse array of dots the variable range hopping conductivity is measurable only when the tunneling length in insulator is large enough. This probably can be achieved in some cases in narrow gap semiconductors. Otherwise conductivity is so small that it can not be measured. In this case, thermodynamic measurements of oscillating quantities may be more convenient.

It is much easier to measure conductivity in a dense array of dots. Actually most of cited experiments are done at $NR^3 \sim 1$. In Sec. V we study clean metallic cubes separated by thin layers of an insulator doped by donors in the “super-dense” limit $d << R$ (Fig. 4). It is assumed that the tunneling conductance between two cubes $G << e^2/h$. We show that in the dense array BDOGS as a function of the average number of donors per dot $\nu$ behaves as shown in Fig. 8. At $\nu << 1$ BDOGS is very small and grows super-linearly with $\nu$ (like $\nu^4$), and at $\nu \sim 1$ it saturates at the value $\tilde{g}_s \sim \kappa/(e^2 R D d)$. 
III. BDOGS IN SPARSE THREE-DIMENSIONAL ARRAYS.

1. Saturation of BDOGS

With enough impurities the BDOGS is strongly smeared and it approaches an averaged value given by Eq. (7). This happens when the fluctuation of potential $\gamma$ is large enough, namely

$$\gamma > e^2/\kappa R .$$  \hspace{1cm} (11)

In order to find the critical $\nu_s$ of the saturation BDOGS, let us study the fluctuation of electric potential. For a given quantum dot, all the charges within a distance of the screening radius $r_s$ contribute into a collective Coulomb potential. There are two kinds of charges, positive donors and negative dots. Let us compare the charge fluctuations made by them. The average number of impurities within $r_s$ is $N\nu r_s^3$, with a typical charge fluctuation $\nu e/\sqrt{N\nu r_s^3}$. For the charged dots, the typical fluctuation of dot number in $r_s$ is $\sqrt{N\nu r_s^3}$, and each dot carries a charge of $\nu e$ on average, so the typical charge fluctuation is $\nu e \sqrt{N\nu r_s^3}$. At $\nu \gg 1$ the charge fluctuations produced by charged dots are larger than that of donors, so that the fluctuation of the potential energy of an electron is mostly related to charge fluctuations of dots

$$\gamma \sim \frac{\nu e^2}{\kappa r_s} \sqrt{N\nu r_s^3} .$$  \hspace{1cm} (12)

On the other hand, the BDOGS at the Fermi level $g_0(\mu)$ determines the linear screening radius due to redistribution of electrons between dots

$$r_s = \frac{1}{\sqrt{4\pi g_0(\mu) e^2/\kappa}} \approx \frac{1}{\sqrt{N R}} .$$  \hspace{1cm} (13)

With the help of Eqs. (12) and (13), the requirement (11) becomes

$$\gamma \sim e^2 \nu (N/R)^{1/4} > e^2/\kappa R .$$  \hspace{1cm} (14)

Therefore only at $\nu > \nu_s \equiv 1/(NR^3)^{1/4}$ can we have big enough potential fluctuation. At $\nu > \nu_s$ BDOGS is smeared so strongly that at a given $\nu$ BDOGS $g(\varepsilon) \sim g_s$ at all energies, and $g_0(\mu)$ does not depend on the position of $\mu$ any longer.

2. Line of maxima of oscillating BDOGS.

When $\nu$ is close to a half-integer $M - 1/2$ ($M$ is an integer) the Fermi level is in the middle of the $M$th peak in the BDOGS of dots (Fig. 9).

![FIG. 9. The origin of g_max. The solid horizontal line is the Fermi level, the dashed line is the fluctuating potential energy. Two peaks of BDOGS of the array are shown, as well as two corresponding levels of 4 dots.

The single dot charging levels vary with the electric potential energy. The typical fluctuation of the potential energy is determined by Eqs. (12), which broadens the BDOGS peak (Fig. 9). The BDOGS at the peak is

$$g_{\text{max}}(\mu) \sim g_{\text{peak}}(\varepsilon) \sim \frac{N}{\gamma} .$$  \hspace{1cm} (15)

In such a situation the linear screening radius is

$$r_s \approx \frac{1}{\sqrt{4\pi g_{\text{max}}(\mu) e^2/\kappa}} .$$  \hspace{1cm} (16)

At $1 < \nu < \nu_s$ Eqs. (12), (15) and (16) self-consistently determine all three unknowns: the BDOGS $g_{\text{max}}(\mu)$, the screening radius and the typical fluctuation of the potential energy. We arrive at Eq. (5) and

$$r_s \approx \frac{\nu^{2/3} N^{1/3}}{N^{-1/3}} ;$$  \hspace{1cm} (17)

$$\gamma \sim e^2 \nu^{4/3} N^{-1/3};$$  \hspace{1cm} (18)

It is clear from Eq. (5) that $g_{\text{max}}$ arrives at the value $g_s \sim N/((e^2/\kappa R)$ when $\nu \sim \nu_s \equiv 1/(NR^3)^{1/4} > 1$. Note that the theory of this subsection is similar to Ref. 21.

3. Line of minima of oscillating BDOGS.

![FIG. 8. BDOGS at the Fermi level $g_0(\mu)$ of a “super-dense” array as a function of filling factor $\nu$.

The distribution of electrons between dots is mostly related to charge fluctuations of donors, so that the fluctuation of the potential energy is large enough. For a given quantum dot, all the charges within a distance $r_s$ contribute into a collective Coulomb potential. There are two kinds of charges, positive donors and negative dots. Let us compare the charge fluctuations made by them. The average number of impurities within $r_s$ is $N\nu r_s^3$, with a typical charge fluctuation $e/\sqrt{N\nu r_s^3}$. For the charged dots, the typical fluctuation of dot number in $r_s$ is $\sqrt{N\nu r_s^3}$, and each dot carries a charge of $\nu e$ on average, so the typical charge fluctuation is $\nu e \sqrt{N\nu r_s^3}$. At $\nu \gg 1$ the charge fluctuations produced by charged dots are larger than that of donors, so that the fluctuation of the potential energy of an electron is mostly related to charge fluctuations of dots

$$\gamma \sim \frac{\nu e^2}{\kappa r_s} \sqrt{N\nu r_s^3} .$$  \hspace{1cm} (12)

On the other hand, the BDOGS at the Fermi level $g_0(\mu)$ determines the linear screening radius due to redistribution of electrons between dots

$$r_s = \frac{1}{\sqrt{4\pi g_0(\mu) e^2/\kappa}} \approx \frac{1}{\sqrt{N R}} .$$  \hspace{1cm} (13)

With the help of Eqs. (12) and (13), the requirement (11) becomes

$$\gamma \sim e^2 \nu (N/R)^{1/4} > e^2/\kappa R .$$  \hspace{1cm} (14)

Therefore only at $\nu > \nu_s \equiv 1/(NR^3)^{1/4}$ can we have big enough potential fluctuation. At $\nu > \nu_s$ BDOGS is smeared so strongly that at a given $\nu$ BDOGS $g(\varepsilon) \sim g_s$ at all energies, and $g_0(\mu)$ does not depend on the position of $\mu$ any longer.

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The single dot charging levels vary with the electric potential energy. The typical fluctuation of the potential energy is determined by Eqs. (12), which broadens the BDOGS peak (Fig. 9). The BDOGS at the peak is

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At $1 < \nu < \nu_s$ Eqs. (12), (15) and (16) self-consistently determine all three unknowns: the BDOGS $g_{\text{max}}(\mu)$, the screening radius and the typical fluctuation of the potential energy. We arrive at Eq. (5) and

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It is clear from Eq. (5) that $g_{\text{max}}$ arrives at the value $g_s \sim N/((e^2/\kappa R)$ when $\nu \sim \nu_s \equiv 1/(NR^3)^{1/4} > 1$. Note that the theory of this subsection is similar to Ref. 21.

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\[ r_s \sim \frac{1}{N^{1/2}R^2}. \tag{20} \]

The potential fluctuation forms wells and humps (Fig. 10). Electrons fall into wells and holes fall into humps. We will discuss the wells and electrons, although our discussion is equally applicable to humps and holes. Inside a well of size \( r_s \) there are other humps and wells of shorter range. Electrons inside wells find themselves in wells of smaller scale and so on until they reach the shortest scale \( l \).

The shortest range \( l \) can be determined by the fact that each dot accepts only one electron. The excess number of charges inside such a well is of the order \( \nu \sqrt{Nl^2} \). The maximum number of electrons in the well is of the same order, otherwise the well turns into a hump. On the other hand the maximum number of electrons is \( Nl^3 \), so that \( Nl^3 \approx \nu \sqrt{Nl^2} \). This gives

\[ l \sim (\nu^2/N)^{1/3} \tag{21} \]

for the characteristic size of the shortest scale well, and

\[ \gamma(l) \sim \frac{\nu e^2}{\kappa l} \sqrt{Nl^2} \sim \nu^{4/3} N^{1/3} v^2 / \kappa. \tag{22} \]

for the characteristic depth. It can be checked that \( l << r_s \) as long as \( \nu << \nu_s \), in accordance with our assumption.

As we said, in a cube of size \( r_s \), the fluctuation of extra charge is \( q \sim \nu e \sqrt{N r_s^3} \), which is balanced by the screening charge \( \Delta N r_s^3 e \) (\( \Delta N \) is concentration of dots participating in screening):

\[ \Delta N r_s^3 e \sim \nu e \sqrt{N r_s^3}, \tag{23} \]

therefore

\[ \Delta N \sim \nu^4 N^2 R^3. \tag{24} \]

With the help of Eqs. (22) and (24), one can estimate the BDOGS at the Fermi level as \( g_{\text{min}}(\mu) \sim \Delta N / \gamma(l) \), and arrive at Eq. (6). For peaks of \( g(\varepsilon) \) at \( \varepsilon = \pm e^2/2\kappa R \) we have \( g(\varepsilon) \approx g_s > g_{\text{min}}(\mu) \) (see Fig. 10).

At \( \nu \sim \nu_s \) the shortest scale of wells and humps \( l \) approaches the longest scale of wells and humps \( r_s \), and each dot can accept more than one electrons or holes because \( \gamma > e^2/\kappa R \). Therefore the non-linear screening crosses over to linear screening at \( \nu \sim \nu_s \). At the same time the fluctuation of potential energy \( \gamma \) is big enough to smear the BDOGS. It can be checked that \( g_{\text{min}} \) and \( g_{\text{max}} \) merge to \( g_s \) at \( \nu = \nu_s \).

4. Growing part of BDOGS.

When concentration of impurities is rather low, \( \nu << 1 \), an electron released by a donor would like to stay on a dot close to the donor. The common situation is a donor accompanied by one charged dot (1-complex), while some donors stay alone (0-complex), and some donors are accompanied by two charged dots (2-complex). Charge conservation requires \( N_0 = N_2 \), where \( N_0 \) and \( N_2 \) are concentrations of 0- and 2-complexes.

At \( \nu << 1 \) nearly all donors are independent, and \( N_0, N_1 \) and \( N_2 \) are proportional to \( \nu \). Therefore the Fermi level remains a constant and the BDOGS grows linearly with \( \nu \) at \( \nu << 1 \).

The details of this problem is the same as in a weakly compensated doped semiconductors. It has been studied quantitatively\(^{15} \), counting the numbers of 0- and 2-complexes. The quantitative results are
\[ N_0 = N\nu \exp \left( -\frac{4\pi e^2 N}{3\kappa|\mu|^3} \right), \]  
\[ N_2 = 7.14 \times 10^{-4} \cdot \left( \frac{4\pi}{3} \right)^2 \cdot N\nu \left( \frac{e^2 N^{1/3}}{\kappa \mu} \right)^6, \]  
\[ \mu = -0.99e^2 N^{1/3}/\kappa. \]  
As the concentration of impurities \( N\nu = N_0 + N_1 + N_2 \) is not changed, the concentration of electrons on dots is controlled by the Fermi level \( \mu \), therefore

\[ g_<(\mu) = \frac{dN_2}{d\mu} - \frac{dN_0}{d\mu} = 0.2\kappa\nu N^{2/3}/e^2. \]  
We kept all coefficients in this subsection because they are known.

Thus \( g_<(\mu) \) is proportional to \( \nu \) and follows Eq. (4). This picture works well at \( \nu << 1 \). At \( \nu \geq 1 \) each dot is typically affected by more than one impurities. It is straightforward to check that Eq. (4) matches Eq. (5) at \( \nu \sim 1 \).

5. Periodic array of dots.

Above we have discussed sparse arrays in which the dots are situated randomly. Now we turn to a question, what if the dots are arranged periodically in space?

The qualitative picture in Fig. 6 is still mostly correct. At \( \nu >> 1 \) there are still the oscillations of \( g_0(\mu) \) with \( \nu \). One should notice that the strongly charged periodic dots do not contribute to the fluctuations of electric potential and only donors produce the fluctuations. As we discussed above these fluctuations are weaker than in the random sparse array. Therefore more donors are needed to smear BDOGS to the same extent. In other words, the lines of maxima and minima in this case converge slower and the value of \( \nu_s \) is larger.

The linear growth of BDOGS at \( \nu << 1 \) shown in Fig. 6 is not valid in a periodic array, since 0- and 2-complexes originate from the random distribution of dots in space. In periodic array a typical donor donates an electron to the nearest dots and forms an electric dipole. It takes additional energy to take an electron out of a dipole and put it near another dipole. Therefore a hard gap is formed at the vicinity of the Fermi level. Only the rare cases where several donors are close to each other can produce non-zero BDOGS at the Fermi level. As a result \( g_0(\mu) \) does not grow linearly with \( \nu \), but with higher power. The similar situation will be studied more thoroughly in the dense array of Sec. V. Therefore, here we gave only a brief description of the result.

IV. BDOGS IN A SPARSE TWO-DIMENSIONAL ARRAY

The \( g_0(\mu) \) as a function of \( \nu \) in 2D looks similar to 3D (Fig. 6). It also has growing part, oscillating part and saturation part. However, in 2D oscillations survive longer than in 3D, because the collective potential in 2D is much weaker. Indeed, the fluctuation of potential energy in 3D grows with \( r_s \) as a power law

\[ \gamma_{3D} \sim \frac{\nu e^2}{\kappa r_s} \sqrt{N r_s^2} \sim r_s^{1/2}, \]  
while in 2D the long-range fluctuation of potential energy grows slower than logarithmically with \( r_s \)

\[ \gamma_{2D} \sim \left( \int_{\sqrt{N}}^{r_s} \frac{\nu e^2}{\kappa R} N^2 R^2 \right)^{1/2} \propto \ln(\sqrt{N r_s}). \]  
Therefore the line of minima of oscillating part in 2D is quite different from that of 3D. The tail in the \( g(\varepsilon) \) vs. \( \varepsilon \) curve is produced here by Coulomb interaction of nearest neighbor dots, and the minimum value of \( g \) appears to be (at \( 1 << \nu < r_s = 1/\sqrt{N R^2} \))

\[ g_{\min} \sim N^2(\nu e^2/\kappa)^2/\varepsilon \]  
\[ g_{\min} \sim \nu^2 N^2 R^3 \kappa/e^2. \]  
The 2D linear screening radius produced by \( g_{\min} \) is

\[ r_s = \frac{\kappa}{2\pi g_{\min} e^2} \sim \frac{1}{\nu^2 N^2 R^3}. \]  
One can verify using Eq. (30) that \( \gamma_{2D} << e^2/\kappa R \) for such \( r_s \) and therefore long-range contribution can be neglected.

The growing part at \( \nu < 1/2 \) can be studied in the similar way as in 3D. That is, one can use the idea about 0-complex, 1-complex and 2-complex to find the Fermi level

\[ \mu \sim -e^2 N^{1/2}/\kappa \]  
and BDOGS at the Fermi level (at \( \nu << 1 \))

\[ g_<(\mu) \sim \kappa N^{1/2}/e^2. \]  
For the maximum line of oscillating part, the collective typical fluctuation of potential energy is given by Eq. (30) and BDOGS at the Fermi level is

\[ g_{\max}(\mu) \sim \frac{N}{\gamma} \sim \frac{N^{1/2} R}{e^2 \nu} \]  
\[ 1 << \nu < \nu_s = 1/\sqrt{N R^2}. \]  
Here we dropped the logarithmic factor and therefore \( r_s \) disappeared from the equations. In this approximation unlike 3D situation, we do not need the self-consistent method. As a result \( g_{\max}(\mu) \) decreases with \( \nu \) much slower than in 3D.
Both Eqs. (35) and (31) reach \( g_\nu (\mu) \sim N/(e^2/\kappa R) \) at \( \nu \sim \nu_s = 1/\sqrt{N R^2} \), where the distinction between peaks and valleys disappears.

Up to now we discussed charging of two-dimensional array of dots by donors or a gate situated outside of dots. In some two-dimensional systems electrons may be provided to a dot by donors located close to the dots. This can be done, for example, by creating a two-dimensional gas with help of close layer of donors parallel to its plane and then by etching outside dots both the gas and the donor layer. If such dots are essentially two-dimensional and heavily doped, internally induced fluctuations can cause their charging, in contrary to what we said in introduction about three-dimensional dots. Indeed, in this case, fluctuations of number of electrons in the dot and corresponding of the Fermi level decrease inverse proportionally to the dot radius. This decay is similar to the charging energy, so that ratio of these energies is just proportional to the ratio of kinetic and potential energies of electron in the dot.

V. DENSE THREE-DIMENSIONAL ARRAYS OF DOTS

Until now we dealt with sparse arrays of dots where \( NR^3 \ll 1 \) as shown in Fig. 2. Let us discuss what happens when the small parameter \( NR^3 \) grows. The saturation filling factor \( \nu_s = (NR^3)^{-1/4} \) decreases, which means the number of oscillations decreases as well. The oscillations disappear at \( \nu_s \sim 1 \) when \( NR^3 \sim 1 \), i.e. when the typical separation between dots is of the order of the dot size.

Let us now concentrate on the other extreme (Fig. 4) where say cubic dots are closely packed in a cubic lattice with the period \( R \) and the width \( d \) of an insulator between dots is small \( d << R \). The insulator is uniformly doped by donors. We show below that at a large enough average number of donors per dot, \( \nu > 1 \), the BDOGS is smeared. On the other hand at \( \nu << 1 \) typical donors do not contribute to BDOGS at the Fermi level but rare clusters of donors result in a small BDOGS.

For the system of metallic cubic dots \( \{i\} \) separated by a clean insulator, potentials \( \phi_i \) and charges \( Q_i \) of dots are related by linear equations

\[
\phi_i = \sum_j P_{ij} Q_j .
\]  

and

\[
Q_i = \sum_j C_{ij} \phi_j ,
\]

where \( P_{ij} \) is the reciprocal matrix of the capacitance matrix \( C_{ij} \). The coefficients satisfy \( P_{ij} = P_{ji}, 0 \leq P_{ij} \leq P_{ii} \) and depend on the relative positions of \( i \) and \( j \) only. Without donors the Hamiltonian can be written as

\[
H = \frac{1}{2} \sum_{ij} P_{ij} Q_i Q_j = \frac{1}{2} \sum_i P_{ii} Q_i^2 + \frac{1}{2} \sum_{i \neq j} P_{ij} Q_i Q_j .
\]

Of course, it has minimum when all \( Q_i = 0 \).

In order to understand the role of ionized donors let us study an isolated donor between the dots A and B (Fig. 11). The positive donor is completely screened by negative charges \( q_A \) and \( q_B \) appearing of the surface of A and B, i.e. \( q_A + q_B + e = 0 \). These charges leave compensating positive charges \( -q_A \) and \( -q_B \) in the dots A and B. If \( x_A \) and \( x_B \) are distances from the donor to the dots A and B respectively then

\[
q_A x_B = q_B x_A .
\]

One can easily arrive at Eq. (39) imagining that the donor for a moment is replaced by the uniformly charged plane parallel to the faces of cube A and B (with the same \( x_A \) and \( x_B \)), and minimizing the sum of Coulomb energies of two emerging plane capacitors with distances between planes \( x_A \) and \( x_B \) as a function of \( q_A \). Note that at \( d << R \) all points of the charged plane are in the same conditions and the plane fields are just a superposition of fields of these charges. This means Eq. (39) should also hold for a point charge of this plane or just for a donor. Eq. (39) means that potentials of dot A and B are equal. In other words, the group made of the positive donor, \( q_A \) and \( q_B \) produces an electric field which is confined in the vicinity of the donor. The group does not affect the electric potential of any dot. Therefore the group can be totally forgotten when we study the charges and potentials of the dots, or in other words it can be totally ignored in the Hamiltonian.
On the other hand, the compensating positive charges $-q_A$ and $-q_B$ cannot be ignored. One can say that the donor’s charge is split into two fractional parts $-q_A$ and $-q_B$. Charges $-q_i$ are uniformly distributed between 0 and $e$.

When there are more than one donor around a dot A we can simply add them up $Q_{AD} = -q_{A1} - q_{A2} - \cdots$ to get the total positive charge induced by donors on the dot A.

Each donor also donates an electron to the array of dots. We assume that $d >> a$, where $a$ is the tunneling length of electrons in the insulator so that the conductance $G$ between two dots is much smaller than $e^2/h$. In this case each dot can get only integer number of extra electrons $n_i$. The neutrality of the whole system requires $-e \sum n_i = \sum Q_{iD}$. For a given fractional set \{n_i\} the integer set \{n_i\} minimizes the Hamiltonian

$$H = \sum_i \frac{P_{ii}}{2} (Q_{iD} - n_i e)^2 + \sum_{i \neq j} \frac{P_{ij}}{2} (Q_{iD} - n_i e)(Q_{jD} - n_j e).$$  

(40)

This energy is calculated from the reference point of the sum of energies of neutral groups around each donor, which does not depend on $n_i$. If each donated electrons were shared by dots this fractional charges would neutralize $Q_{iD}$ and make $H = 0$ and all $\phi_i = 0$. This is equivalent to what would happen if all grain are grounded. Discreteness of $n_i$ makes finding the ground state $n_i$ non-trivial. It is clear that in the globally neutral system the chemical potential $\mu$ lies in the same place as for a system of neutral dots, i.e., $\mu = 0$. In the ground state we can introduce “one-dot” energies of an empty dot state above the Fermi level assuming that charges of other dots are fixed

$$\varepsilon_i^{(e)} = H(n_i + 1) - H(n_i) = -e \phi_i + \frac{1}{2} P_{ii} e^2, \quad (41)$$

where

$$\phi_i = \sum_j P_{ij} (Q_{j} - n_j e)$$  \quad (42)

is the electrostatic potential of the dot $i$. For an occupied state in the similar way

$$\varepsilon_i^{(o)} = H(n_i) - H(n_i - 1) = -e \phi_i - \frac{1}{2} P_{ii} e^2. \quad (43)$$

In the absence of donors when all dots are neutral, i.e., $\phi_i = 0$, we get $\varepsilon_i^{(e)} = +\frac{1}{2} P_{ii} e^2$ and $\varepsilon_i^{(o)} = -\frac{1}{2} P_{ii} e^2$. The single electron BDOGS for neutral dots looks the same as in Fig. 3 except the separation of peaks or the level spacing is smaller now $e^2 P_{ii} \sim e^2/\kappa_{eff} R \approx e^2 d^2 \rho$ (the calculation of $P_{ii}$ is in Appendix). Here

$$\kappa_{eff} \approx \kappa R/d \quad (44)$$

is the effective dielectric constant of the array. For charged dots stability of the global ground state requires $\varepsilon_i^{(e)} > \mu = 0$ and $\varepsilon_i^{(o)} < \mu = 0$.

Let us study the BDOGS in the case when there are more than one donors per dot ($\nu \gg 1$). In order to find BDOGS we neglect the interaction terms in Eq. (40) and consider only diagonal terms. As a result the ground state of the system is realized by such a set of \{ni\} that $-e/2 \leq Q_{iD} - n_i e \leq e/2$. So effective charges on different dots in the ground state of the array are uniformly distributed from $-e/2$ to $e/2$. This gives the energy independent BDOGS

$$\tilde{g}_s(\varepsilon) \sim \frac{1}{e^2 P_{ii} R^2}. \quad (45)$$

Consider now the role of non-diagonal terms. At a big distance $r_{ij} >> R$ the coefficients $P_{ij}$ have the Coulomb form $P_{ij} \approx 1/\kappa_{eff} r_{ij}$. They, therefore, result in a density of ground states (DOGS) with the Coulomb gap at the Fermi level, which in turn leads to ES law.

The finite BDOGS at the Fermi level and ES law at $\nu \gg 1$, originates from random distribution of charges $Q_i$, which in turn is related to the fact that several donors contribute into each $Q_{iD}$. Even for nearest neighbor dots $Q_{iD}$ and $Q_{jD}$ are independent since they include contributions of different donors.

We turn now to the case of a small density of donors $\nu << 1$ and first deal with typical donors isolated from each other. As discussed above for such an isolated donor between the dots A and B, the charges induced by the donor are related by $Q_{AD} + Q_{BD} = +e$. If the donor is closer to the dot A, $Q_{AD} > e/2 > Q_{BD} > 0$. If we neglect interaction between the dots A and B in Eqs. (41) and (43) we arrive at finite BDOGS of isolated donors at the Fermi level. States at the Fermi level are provided by dots where $Q_{BD} = 1/2 - \delta$ with $\delta << 1$. Then the dot B has a state just above the Fermi level and A has a state just below it. The fact that these states are so closely correlated in space leads to their elimination by the interaction between the dots A and B. Indeed, in the ground state the electron donated by the donor is on the dot A. The total charges on A and B is $Q_A = Q_{AD} - e = -Q_{BD}$ and $Q_B = Q_{BD}$. The electric potential on B is $\phi_B = P_{BB} Q_B + P_{BA} Q_A$ and according to Eq. (41) all the levels on B are moved down by $e \phi_B$. Using the fact $0 \leq Q_{BD} \leq e/2$, we have

$$0 \leq e \phi_B \leq (1 - \alpha) P_{BB} e^2/2, \quad (46)$$

where $\alpha \equiv P_{BA}/P_{AA} = 0.34 \pm 0.01$ represents the interaction between nearest neighbor dots (see the calculation of $\alpha$ in Appendix). According to Eq. (41) the lowest vacant level of dot B is at least $\alpha P_{BB} e^2/2$ above the Fermi level. Similarly, according to Eq. (43) the highest filled level on the dot A is moved up by $-e \phi_A$ with $0 \leq -e \phi_A \leq (1 - \alpha) P_{AA} e^2/2$ and it is at least $\alpha P_{AA} e^2/2$ below the Fermi level (Fig. 12).
Thus, we showed that for $\nu \ll 1$ typical isolated donors have a substantial hard gap of width $\alpha P_\nu e^2$ in their density of states. This gap can not be destroyed by long range interactions of such donors because they form neutral complexes and weakly interact with each other at long distances.

One can also think of a donor in the neutral complex with electron of the neighboring dot A as a hydrogen-like donor in a semiconductor. Then the hard gap obtained above is similar to the Hubbard gap. The electric dipole made of a donor and an electron in principle can accept another electron with a weak binding energy, but the energy difference or Hubbard gap between the two electronic levels makes sure that only the lower level is filled in the ground state.

Up to now we talked about typical isolated donors at $\nu \ll 1$. In fact there are rare configurations that make a non-zero BDOGS at the Fermi level. Let us show how this happens.

Consider, for example, a cluster of 4 donors around one dot (Fig. 13). The charges of donors are shared by 5 dots in such a way that charge of each donor is effectively split into $4e/5$ going to the central dot and $e/5$ going to its nearest neighbor. The middle dot has effective charge $Q_{ij} = 16e/5$ and each of the four neighbor dots has $Q_{ij} = e/5$. Let us show that in the ground state of the cluster $n_i = 3$ for the central dot and $n_i = 0$ for all others, i.e. there are only three electrons on the cluster of four donors so that the cluster has a net charge $+e$. It is effectively shared by the 5 dots, each getting $+e/5$.

According to Eq. (41) addition of the fourth electron to this cluster costs

$$\varepsilon_i^{(e)} = \left[ \frac{1}{5} (4\alpha + 1) + \frac{1}{2} \right] P_{ii} e^2 \approx 0.03 P_{ii} e^2$$  \hspace{1cm}(47)$$

where $\alpha = P_{12}/P_{11}$, and as shown in Appendix $\alpha = 0.3405$. This means that $\varepsilon_i^{(e)} > \mu = 0$ and the positively charged cluster of Fig. 13 is stable with respect of bringing the fourth neutralizing electron. The physical reason for such stability is uniform smearing of the net charge over 5 dots which diminishes potential of the charged cluster at the Fermi level.

We have shown above an example of a positive “donor-like” cluster. Moving each of the four donors away from the central dot in the direction of the neighbor dot one can construct an “acceptor-like” cluster in which charge $-e$ is evenly shared by 5 dots. Such a cluster accepts an electron released by the cluster in Fig. 13. Thus, some clusters charge themselves by “self-compensation” similarly to amphoteric impurities in semiconductors. Of course, there are also neutral clusters and tuning positions of donors between dots one can continuously go from a neutral cluster to the charged one. This means that BDOGS at the Fermi level created by clusters is finite.

Using coefficients $P_{ij}$ calculated in Appendix, it is easy to find that the smallest charged cluster in the ground state is the cluster of 4 dots with 3 donors between them. Therefore, at $\nu \ll 1$ the BDOGS at the Fermi level grows as $\gamma_0(\mu) \propto \nu^3$ and it begins to be significant at $\nu \sim 1$ (see Fig. 8).

The structure of self charging clusters discussed above is similar to the mechanism which provides a non-zero bare density of states in a system of quasi-one-dimensional electron crystals at low impurity concentration. There is also similarity to self-compensation of clusters of several donors in uncompensated semiconductor near the insulator-metal transition.

As we saw above in the “super-dense” array of dots with $d \ll R$, a large enough concentration of donors can easily smear BDOGS. This is a natural explanation of the experiments, where the ES law is observed. Let us consider parameter $T_0$ of the ES law (Eq. (1)) for a dense array of dots at $\nu \gg 1$. According to Ref. 4 $T_0 \sim e^2/(\kappa_{eff} \xi)$, where $\xi$ is the localization length for tunneling to distances much larger than $R$. When an electron tunnels through insulator it accumulates dimensionless action $d/a$, where $a$ is the tunneling decay.
length in the insulator. On the distance $x$ electron accumulates $x/R$ such actions. Thus, its wave function decays as $\exp(-x d/R a) = \exp(-x \xi/a)$, where the localization length $\xi = a R/d$. This enhancement of localization length is similar to the one derived for disordered granular system near percolation threshold\textsuperscript{25}. It leads to

$$T_0 \sim \frac{e^2 d}{\kappa_{\text{eff}} a R} = \frac{e^2 d^2}{\kappa R^2 a}.$$ \textsuperscript{(48)}

We see that with decreasing $d$ the temperature $T_0$ decreases as $d^2$. This continues until the conductance of the insulator layer $G = (e^2/h)(Rk_F)^2 e^{-2d/a}$ reaches the quantum limit $G \sim e^2/\hbar$, i.e. while $d > d_c \equiv a \ln(Rk_F)$. At $d << d_c$ the characteristic ES temperature $T_0(d)$ should vanish when $d \to 0$, but the way how this happens is still unknown (see also recent publications\textsuperscript{26,27} concerned with the case $G >> 1$).

In the above calculation of the localization length we assumed that tunneling through a metallic dot does not accumulate more action. Actually a dirty dot provides logarithmic contribution to the tunneling action\textsuperscript{28}. We assume that it is much smaller than $2d/a$ resulting from crossing the insulator.

Until now we discussed only the “super-dense” array shown in Fig. 4. Let us consider what happens in the system of almost densely packed metallic spheres (Fig. 14), where insulator occupies a larger fraction of space than in Fig. 4.

![Fig. 14. An almost densely packed array of metallic spheres ($d << R$). A donor is shown by + and electron donated by it is shown by -.
](image)

One can show that in such a system BDOGS as a function of number of donors per dot $\nu$ behaves qualitatively similar to Fig. 8. At smaller $\nu$ only a cluster of donors around a dot can create a finite BDOGS, while at $\nu >> 1$ donors easily smear BDOGS. In this case, characteristic temperature of ES law is $T_0 \approx e^2 d/a R$. It loses one power of $d/R$ because the dielectric constant of such systems does not diverge at $d \to 0$. When distance between dots $d$ grows and exceeds $R$, we get sparse periodic array and recover oscillations of BDOGS at $\nu >> 1$. At $\nu << 1$ there is a smooth cross-over between two super-linear growths of BDOGS related to rare donor clusters in the vicinity of some dots.

Although we arrived at $\nu >> 1$ as the universal criterion of substantial smearing of BDOGS for both dense arrays (Figs. 4 and 14) and sparse arrays (Figs. 2), we should understand that the required concentration of donors $N_D$ in the insulator grows while volume fraction of insulator decreases from the arrays on Figs. 2 and to the one on Fig. 14 and then to the one on Fig. 4.

In some cases metallic granules are first coated by the layer of a doped insulator (for example, the metal’s oxide) and then compressed into the bulk array with a clean insulator of different kind of filling or just air filling empty space (Fig. 15). For simplicity we assume the dielectric constant of the coating insulator is close to that of the filling space between dots.

![Fig. 15. An array of coated dots. The width of coating insulator is $w$. A donor is shown by + and electron donated by it is shown by -.
](image)

We would like to consider BDOGS for such a model of granular metal in order to understand cross-over between the studied above case of donors residing outside dots and those on the dot surface or inside dots. In the latter case, donors are screened and act as short range neutral impurities. According to the introduction such impurities can not charge a neutral array of dots. Thus, effect of donors in the layer of the width $w << R$ should decrease as $w$ vanishes.

Indeed, one can calculate, fluctuation of average the potential of donor layer of the width $w$ which plays the role of fluctuation of work function of dots. It can be estimated as a potential created by the layer of the width $w$ by fluctuating concentration $(N_D w R^2)^{1/2}/w R^2$. One can imagine that, correction to the work function is created by effective random double layer. Such correction to the work function is of order $e^2((N_D w^3)^{1/2}/\kappa R$. It becomes larger than the charging energy only at $N_D w^3 >> 1$\textsuperscript{29}. This condition of BDOGS smearing is obviously much stronger than “one donor per dot” condition valid for a uniform insulator. When $w$ approaches the lattice constant the role of donors is so much weakened that in order to smear BDOGS one needs the relative concentration of donors to be of the order of 50%. At the surface, donors and their images are still forming dipoles with fluctuating dipole moments, which create fluctuating double layer potential. On the other hand, when donors enter inside the dot, each donor becomes exponentially screened from all sides, the dipoles and the random double layer disappear. This diminishes the role of donors further so that as we said in introduction even large concentration of impurities inside the whole three dimensional dot can not charge the dot.
VI. CONCLUSION

This paper addresses the problem of explanation of the temperature dependence of the conductivity observed in granular metals and arrays of quantum dots. We show that ES law is the most natural explanation for this dependence (in spite of recently expressed doubts30). For this purpose we present a simple model with random charging of clean metallic dots in both sparse and dense arrays of dots. In both cases we assume that the insulator separating dots is uniformly doped by donors, which donate their electrons to dots. Random positions of dots in the sparse case (Fig. 1) and random positions of donors in the dense periodic model (Fig. 5) lead to random charging of dots in the global ground state and to filling of the gap of the bare density of individual dots at the Fermi level. The long range Coulomb interaction creates a soft Coulomb gap on the background of the finite bare density of ground states in the vicinity of the Fermi level. At low enough temperature this leads to the ES variable range hopping conductivity, too. It was shown that at low temperatures a strong electric field replaces temperature dependence (in spite of recently expressed doubts30). As a result non-ohmic ES law reads

\[ \frac{1}{6} \sum_{\vec{r}'} \phi(\vec{r}) = \frac{Q_0 d}{6 \kappa R^2} \delta_{\vec{r},\vec{r}'} , \tag{49} \]

where \( \vec{r} \) are 6 nearest neighbor of the site \( \vec{r} \), and \( \phi(\vec{r}) \) is defined only on the discrete sites \( \vec{r} = (n_x R, n_y R, n_z R) \). Eq. (49) can be solved using discrete Fourier transform.

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APPENDIX: MATRIX ELEMENTS \( P_{ij} \) FOR NEIGHBORING DOTS.

Here we study the coefficients \( P_{ij} \) in Eq. (36). We know the diagonal term \( P_{ii} \sim 1/\kappa_{eff} R \), and for distant neighbors \( r_{ij} \gg R \) one can show that \( P_{ij} \sim 1/\kappa_{eff} r_{ij} \). But the coefficients between close neighbors require further consideration.

The dense array of periodic dots is equivalent to the lattice of identical capacitors connecting adjacent dots. Such a system can be studied in the language of an equivalent resistor network. Imagine a cubic lattice network with an identical resistance between every nearest neighbor sites. Suppose that current \( I \) goes into a lattice site \( A \) and travels out through the network to infinity. Label the potential of site \( A \) to be \( V \), the potential of nearby dots \( B, C, D, \ldots \) to be \( \alpha_{AB} V, \alpha_{AC} V, \alpha_{AD} V, \ldots \). What are the values of the \( \alpha_{AB}, \alpha_{AC}, \alpha_{AD}, \) and so on? Here we are using the dimensionless coefficients such as \( \alpha_{AB} = P_{AB}/P_{AA} \), and we know all of them are between 0 and 1.

A standard numerical calculation uses the fact that the current going into a site equals the current going out of it (Kirchhoff law), which means the potential of a site equals the average potential of its nearest neighbors. Using lattices of \( N \times N \times N \) sites and keeping the potential at central cite to be 1 and that of boundaries to 0, one adjusts the potential values of the other sites so that the potential of each site is very close to the average potential of its nearest neighbors. Using lattices with \( N \) up to 97 we find \( \alpha \equiv \alpha_{01} = 0.34 \pm 0.005, \alpha_{02} = 0.166 \pm 0.005, \alpha_{03} = 0.106 \pm 0.005, \alpha_{11} = 0.216 \pm 0.005, \alpha_{111} = 0.17 \pm 0.005 \) (here 111 represents the vector between the two dots measured in the units of the lattice constant).

There is also an analytic way to find the coefficients32. Imagine the situation when there is charge \( Q_0 \) on the central dot while all the other dots remain neutral, one can use the equation similar to Kirchhoff law mentioned above

\[ \phi(\vec{r}) - \frac{1}{6} \sum_{\vec{r}'} \phi(\vec{r}') = \frac{Q_0 d}{6 \kappa R^2} \delta_{\vec{r},\vec{r}'} \]

where \( \vec{r} \) are 6 nearest neighbor of the site \( \vec{r} \), and \( \phi(\vec{r}) \) is defined only on the discrete sites \( \vec{r} = (n_x R, n_y R, n_z R) \).
\[ \phi_\vec{k} - \frac{1}{6} \phi_\vec{k} \cdot (\cos k_x R + \cos k_y R + \cos k_z R) = \frac{Q_0 d}{6 \kappa R^2}. \]  

(50)

One can find \( \phi_\vec{k} \) and

\[ \phi(\vec{r}) = \frac{dQ_0}{2\kappa R^2} \int_{BZ} \frac{d^3k}{(2\pi)^3} \frac{R^3 \cos(\vec{k} \cdot \vec{r})}{3 - \cos k_x R - \cos k_y R - \cos k_z R} \]  

(51)

where the integration is over the Brillouin zone \( k_x, k_y, k_z \in (-\pi/R, \pi/R) \). One can use numerical integration to find

\[ P_{00} \equiv \frac{\phi(\vec{0})}{Q_0} = \frac{0.2527d}{\kappa R^2}, \quad \alpha \equiv \alpha_{01} = \frac{P_{01}}{P_{00}} = 0.3405, \]

\[ \alpha_{02} = 0.1697, \quad \alpha_{00} = 0.1089, \quad \alpha_{11} = 0.2183 \]  

(52)

and so on. These results agree with the results of the first method.

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