Molecular Modeling and Electron Transport in Polyethylene

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ABSTRACT

Polyethylene is commonly used as an insulator for AC power cables. However it is known to undergo chemical and physical change which can lead to dielectric breakdown. Despite almost eighty years of experimental characterization of its electrical properties, very little is known about the details of the electrical behaviour of this material at the molecular level. An understanding of the mechanisms of charge trapping and transport could help in the development of materials with better insulating properties required for the next generation of high voltage AC and DC cables. Molecular simulation techniques provide a unique tool with which to study dielectric processes at the atomic and electronic level. Here we summarise simulation methodologies which have been used to study the properties of PE at the molecular level, elucidating the role of morphology in the trapping of excess electrons. We find that polyethylene has localised states due to conformational trapping extending below the mobility edge (above which the excess electrons are delocalised), at -0.1±0.1eV with respect to the vacuum level. These trap states with localisation lengths between 0.3 and 1.2nm have energies as low as -0.4±0.1eV in the amorphous and interfacial regions of polyethylene with more positive values in lamella structures. Crystalline regions have a mobility edge at +0.46±0.1eV, so we would expect transport by electrons excited above the mobility edge to delocalised states to be predominantly through amorphous regions if they percolate the sample.

Index Terms — Molecular simulations, polyethylene, methodologies, electron trapping, electron transport, morphology.

1 INTRODUCTION

Polyethylene’s low electric conductivity, low dielectric loss, high dielectric strength and outstanding mechanical properties [1] make it a common choice as an insulator for AC power cables. However, under certain conditions, polyethylene may suffer from chemical and electrical aging or degradation after long-standing operation under voltage, which may finally lead to dielectric breakdown [2].

A major contribution to aging and breakdown is thought to be made by trapped charges, called the ‘space charge’ [2]. The accumulation of space charge distorts the local electric field, facilitating phenomena such as treeing as well as generating chemical and physical defects through the local release of trapping energies (of the order of an eV), eventually leading to the degradation of the insulation, although the exact mechanisms operating are still not fully understood. The formation and transportation mechanism of space charge has been the subject of extensive experimentation [2]. For example, by monitoring the threshold electric field for charge accumulation it has been found that charge injection is strongly affected by the material of the electrode [3-5]. Space charge formation and transport phenomena under DC stress can be observed using the Pulsed Electro-Acoustic (PEA) method [6-9]. This technique has been used to show that the interfaces either between multilayers of PE or between PE and the electrode, is prone to space charge accumulation, and thus might be considered to be weak points with respect to insulating properties [10-12].
Despite a vast literature concerned with the experimental characterization of its electrical properties, very little is known about the details of the electrical behavior of this material, especially at the molecular level. An understanding of the mechanisms of charge trapping and transport in these insulators and how such processes are affected by local physical structures and chemical impurities could help in the development of materials with better insulating properties required for the next generation of high voltage AC and DC cables. In what follows we summarize molecular simulation methodologies which have been used to study the properties of PE at the molecular level, elucidating the role of morphology as well as chemical imperfections and additives on the trapping of excess electrons.

The paper is organized as follows: Section II presents simulation methodologies which have been found useful in understanding PE, including: Molecular Dynamics (MD) for modeling morphology, Density Functional Theory (DFT) for calculating electronic properties using the generated morphologies, the Lanczos method which can be used to predict the density of states of excess electrons in a given PE structure, and the Surface Hopping method which predicts the quantum transitions between such states. Section III reviews current work focusing on the relationship between the electronic states and PE morphologies. The findings are discussed and some conclusions presented in Section IV.

2 MOLECULAR METHODOLOGIES

In many branches of science and engineering, molecular simulation techniques (including MD, DFT, etc.) are an integral part of modern research programs. They can provide a theoretical basis upon which to interpret experimental results as well as a tool for discovery in their own right. For example, computer simulations can explore the properties of ‘perfect’ materials free from chemical and morphological contamination [13]. Once the relationship between the simulated and experimental systems is established, the simulation can be used to explore the effects of systematically changing both external (pressure, temperature, density, etc.) and internal (composition, morphology, defect structure, etc.) variables. In this section, we provide a brief introduction to simulation techniques which have been found to be useful for polymeric materials.

2.1 MOLECULAR DYNAMICS METHODS

Molecular dynamics (MD) [13, 14] is widely used to simulate the trajectories of molecules or atoms in a classical many-body system by solving Newton’s equations of motion. The thermodynamic, structural and transport properties of the system can then be obtained by taking time averages over the trajectories.

This method is widely used for almost any molecular system at moderate or high temperatures (such as at normal conditions, 300K), so that quantum effects in the dynamics of atoms or molecules can be safely neglected.

The interactions between the molecules are described using a force-field, that is, a set of potential energy functions. These functions are usually empirical, consisting of intra-molecular terms representing the energy associated with for example atom-atom bonds, and orientation (valence and dihedral angles) and intermolecular terms representing the van der Waals or dispersion interactions, Coulomb interactions between charges, directional bonds such as H-bonds and, where necessary, three or many-body terms.

Given the force-field and a set of initial conditions, the trajectory of the whole system is usually integrated as a function of time using a simple time-step algorithm such as the velocity Verlet method [14, 15]. Clearly, it is important to choose a time-step small enough so that the generated trajectories are realistic. Typical time-steps for a polymeric system are of the order of a few femtoseconds and averages are taken over thousands to hundreds of thousands of configurations (time-steps) of the system.

The standard MD simulation generates a micro-canonical ensemble, with constant number of particles (N), volume (V) and energy (E). The instantaneous temperature T is obtained from the total kinetic energy as given by the equipartition theorem.

By introducing to the simulation a thermostat to fix the system temperature (popular choices are those given by Berenson, Andersen or Nose-Hoover) or a barostat to fix the pressure, MD simulations can be also carried out in a canonical ensemble (NVT) or an isothermal-isobaric ensemble (NPT) [14, 15].

When only equilibrium properties are required, Monte Carlo (MC) simulation [14] provides an alternative methodology. MC does not generate realistic trajectories but samples configuration space using Metropolis sampling to generate a set of configurations consistent with the statistical mechanical probability distribution for the given ensemble (i.e. isothermal-isobaric corresponding to constant number of atoms N, pressure P, and temperature T or grand canonical corresponding to constant chemical potential µ, volume V, and temperature T). The fictitious trajectories are generated by randomly changing the microscopic degrees of freedom (atom coordinates, etc.). The name of the method comes from the famous Monte Carlo casino, due to the extensive use of random numbers. Thermal equilibrium is usually guaranteed by implementing an ergodic sampling that also complies with detailed balance.

Thus far we have only considered classical methods where the electronic degrees of freedom are incorporated into the potential function describing the inter-atomic interaction. For explicit electronic properties, such as the density of electron states, or excess electron dynamics, a quantum mechanical simulation method is required.

2.2 DENSITY FUNCTIONAL THEORY

Density Functional Theory (DFT) (see book eg. reference [16]) is a very powerful and successful method for investigating the electronic structure of a many-body system (i.e. a multi-electron system). In many cases the results of DFT calculations for solid-state systems agree very well with experimental data. For covalent, metallic and ionic bonds DFT will usually be within 2-3% for the geometry (bond...
DFT is based on the Hohenberg–Kohn (HK) theorems [16, 18]. The first one states that “The ground-state energy from the Schrödinger equation is a unique functional of the electron density”, which implies that the ground-state electron density n(r) uniquely determines all properties, including the energy and wave function, of the ground state. The second HK theorem defines an important property of the functional: “The electron density that minimizes the energy of the overall functional is the true electron density corresponding to the full solution of the Schrödinger equation”. Therefore, by minimizing the functional of the electron density, the electronic structure of the system can be then calculated.

The Schrödinger equation of the many electron system can be rewritten as a set of single-particle equations for an auxiliary non-interacting system, the Kohn-Sham (KS) equation [19],

\[ \frac{\hbar^2}{2m} \nabla^2 + V(r) + V_H(r) + V_{XC}(r) \psi_i(r) = \epsilon_i \psi_i(r) \]  

The terms on the left-hand side of this equation are, in order, the kinetic energy; the Coulomb interactions between the electron and the nuclei; the Hartree term describing the electron-electron Coulomb repulsion; and the exchange-correlation potential which includes all the many-particle interactions. This latter term can be defined as a functional derivative of the exchange-correlation energy as,

\[ V_{XC}(r) = \frac{\partial E_{XC}(r)}{\partial n(r)} \]  

where n(r) refers to the electron density. If the exchange-correlation energy is known as a functional of the density, we have a closed set of self-consistent equations yielding a solution to the electronic structure problem.

Since the Hartree term V_H(r) and V_{XC}(R) depend on n(r), which depends on the KS wave-functions \( \psi(r) \) which in turn depend on the total effective potential \( V(r) + V_H(r) + V_{XC}(r) \), the problem of solving the Kohn–Sham equation has to be done in a self-consistent way. Starting with an initial guess for n(r), the corresponding effective potential can be calculated and the Kohn-Sham equations solved for \( \psi(r) \). From these a new density is calculated and the process starts again. This procedure is then repeated until convergence is reached.

The exact form of the exchange-correlation energy functional is only known for the free electron gas. Thus, in practice, E_{xc} is calculated using approximations, such as local density approximation (LDA) [19], or generalized gradient approximations (GGA) [17], or B3LYP [20], see also [21]. Finding the right functional can be considered the equivalent problem to finding the potential function in a classical simulation.

The Kohn-Sham wave functions are also sometimes identified with electronic states, and the KS energies are taken as estimates of single-particle energies, including that of an excess electron or a hole in the system. While this is not a rigorous procedure, this identification provides a simple method to obtain estimates in many systems that would be very difficult to obtain otherwise. Note however that the results of this uncontrolled approximation are not expected to be as accurate as the standard results (total energy, total electronic density, plus the atomic positions obtained when used for equilibration or dynamics) of DFT.

Within the framework of the Born-Oppenheimer adiabatic approximation, an ab initio Molecular Dynamics technique can be obtained when the forces exerted by the electrons on the nuclei are computed using DFT at every time-step. The nuclei are then moved using Newton’s laws as in the standard MD method. One of such approaches is the so-called Car-Parrinello method [22]. As an artefact to speed up the calculations, the Car-Parrinello method explicitly introduces the electronic degrees of freedom as fictitious variables. The resulting fictitious dynamics keeps the electrons on the electronic ground state, thus avoiding an explicit electronic minimization at each time-step. From among all ab-initio Molecular Dynamics techniques, the Car-Parrinello method is perhaps the most widely used.

Finally, let us note that DFT calculations do not provide fully ab initio solutions of the full Schrödinger equation because the exact functional is not known, however, approximate functionals can be tuned to experimental data if sufficient information is available for the system of interest. Among the shortcomings of DFT is its inability to account accurately for the weak van der Waals attractions that exist between atoms and molecules (see [23] for a summary of current progress). In addition, being strictly a ground state theory standard DFT cannot predict excited states (however Time-dependent density functional theory [24] can predict the properties of excited states of a multi-electron system) and is not capable of predicting accurate band gaps for semiconducting and insulating materials. However many variations and elaborations of density functional theory codes exist which address these problems (see for example [25, 26] for the successful prediction of band gaps), and in general the method is extremely powerful.

In the following, we conclude this section by discussing two methods which allow quantum calculations of a single electron in the system. These calculations are appropriate for the study of excess electrons in the dielectric, when the number of them is not too large, so that we can treat the excess electrons as independent particles. These methods are more amenable when compared to DFT, requiring much less
computer power and without many of the typical difficulties of the DFT techniques.

2.3 LANCZOS METHOD

The Lanczos algorithm is a numerical method for diagonalising (i.e. finding the eigenstates and eigenfunctions) for a given matrix A. Here the matrix of interest is the Hamiltonian \( H \) describing the interaction of an excess electron with PE through a pseudopotential \( V(x,y,z) \), which is an input to the calculation (for details see [27] and references therein). The Schrodinger equation is then solved for the ground and excited states of the excess electron (a single electron) for given configurations of atoms, these corresponding to different morphologies. The Lanczos method is particularly useful for the purpose of solving the Schrodinger equation for a single electron on a three dimensional grid of points in real space \((x,y,z)\), because the Hamiltonian is sparse, having many negligible terms. It is based on the generation of a set (or a set of blocks) of basis functions which are designed to allow the efficient diagonalisation of \( H \), so that the ground and low lying excited states of the excess electron are found. This calculation directly provides the information required to estimate the density of excess electron states of our system. This information can then be used to calculate transport properties such as excess electron mobilities due to extended states.

We briefly describe here the Block Lanczos method proposed by Webster, Rossky and Friesner [28, 29]. We start from a trial set of \( n \) basis functions \( \{ \psi_n \} \) comprising the vectors \( \psi_i \) (for example sine or cosine functions of position). The operator \( W = \exp(-\beta H) \), where \( \beta \) is a positive constant, is applied to this set to get a second set of eigenfunctions \( \{ \psi_n^2 \} \). In each step the Gram-Schmidt method [30] is used to guarantee that this new set is orthogonal to the first set and the basis functions in a given set are kept orthogonal between themselves. The first set \( \{ \psi_n \} \) is the first ‘block’ of our space, (called the Krylov space), with which we diagonalise \( W \). The set \( \{ \psi_n^2 \} \) is the second block, and so on. By repeated application of the operator \( W \), we can obtain \( N_n \) blocks, which constitute the space within which the operator \( W \) is diagonalised. \( W \) has the same eigenvalues and eigenfunctions as \( H \) and is used because it increases the convergence of the solution towards the ground and lower excited states of the system, which are our primary interest. The results depend on the parameter \( \beta \), which is varied until the solutions converge.

2.4 SURFACE HOPPING

The Lanczos method provides the excess electron properties for a single atomic configuration. The surface hopping technique (see reference [31]) is a method to simulate the dynamics of a quantum particle (here to be identified with an excess electron) in interaction with a larger classical system (such as PE), which allows us to go beyond the Born-Oppenheimer adiabatic approximation. We describe here the algorithm proposed by Tully [32, 33], which accounts approximately for the quantum transitions between energy levels (surfaces) and the way these transitions (hopping) affect the dynamics of atoms in the system, which are treated classically. The term ‘surface’ here refers to the single electron energy eigenstates of the system.

Let \( r=(x,y,z) \) refer to the coordinates of the quantum particle and \( R(t)=[R_i(t)] \) those of the classical molecules, then the total Hamiltonian describing the electronic motion, \( H_e(r,R) \), will be a time-dependent operator. The wave function of our quantum particle \( \psi(r,t) \) can be solved using the time-dependent Schrödinger equation,

\[
\frac{i\hbar}{\partial t} \psi = H \psi
\]  

(3)

It is helpful to use a set of adiabatic basis functions \( \Phi_n(r,R) \), which can be computed using the Lanczos algorithm described in the previous Section,

\[
\psi(r,t) = \sum_n a_n(t) \Phi_n(r,R(t))
\]  

(4)

Where the \( a_n(t) \) are expansion coefficients. Substituting (4) into (3), multiplying from the left by \( \Phi_m \) and integrating over \( r \) gives

\[
\frac{d a_m}{dt} = \sum_n d_{mn} a_n
\]  

(5)

where

\[
d_{nn} = -iE_n \delta_{nn} / \hbar - <\phi_m | \partial / \partial t | \phi_n>
\]  

(6)

\[
<\phi_m | \partial / \partial t | \phi_n> = \sum_j \frac{dR_j}{dt} \cdot D_{mn}^j
\]  

(7)

\[
D_{mn}^j = <\phi_m | \nabla_j | \phi_n>
\]  

(8)

By rewriting Eq. (5) in the equivalent density matrix notation, we define \( \rho_{mn} = a_n^* a_m \), so that the Schrödinger equation can be written as

\[
\frac{d \rho_{mn}}{dt} = -\sum_n 2 \text{Re}(d_{mn}^* d_{nm})
\]  

(9)

The diagonal elements \( \rho_{nn} \) are the electronic state population and the off-diagonal elements \( \rho_{mn} \) define the coherence. Each term in the sum can be interpreted as the number of transitions from the state \( n \) to the state \( m \) per unit time. Therefore, during a small time step \( \delta \) we can express this quantity as the probability that the electron is in state \( n \) multiplied to the probability of the transition to other state:
Once a transition to other state has been decided by using Monte Carlo transition probability [14], we must change the kinetic energy of the classical subsystem in order to conserve energy. This is usually done by scaling the velocities \( v_i \) of the classical particles in the direction of the corresponding non-adiabatic coupling vector \( D_{nm}(t) \).

In this implementation of the surface hopping method, the classical particles are moved at fixed intervals according to MD. Whether there is a quantum transition is decided during the last time step (i.e. if \( t \) is the current time step, the transition is decided in the interval from \( t - \Delta t \) to \( t \)). If the hop is accepted we scale the atom’s velocities at \( t \). The quantum force over the atom \( j \) can be derived from the Helman-Feynman theorem as

\[
F_j = - \langle \phi_n | \nabla_j H_e | \phi_n \rangle \tag{11}
\]

This implementation of surface hopping method makes it possible to study the non-adiabatic dynamics of a small quantum system in contact with a larger classical system in an approximate way. As such, it goes beyond the scope of DFT or the Car-Parrinello method, since these latter techniques assume that the whole system always remains in the ground-state.

### 3 Molecular Modelling of Polyethylene

#### 3.1 PE Morphologies and Its Conformational Disorders

Early simulation studies on bulk PE used n-alkane short chains to mimic long chain PE. Simulations of PE chains, each with 50 CH\(_2\) units, below and above the glass transition temperature, was performed to study PE physical and conformational properties such as chain conformation, glass transition or free-volume distribution [34-37]. The transition between rotational isomers in PE chains of 100 CH\(_2\) units was analyzed in [38]. All-trans PE chains of 60 CH\(_2\) units was studied [39] around the melting point. However, none of those simulations observed the chain folding phenomenon widely seen in experiments, and believed to be one of the basic motifs of the PE bulk sample.

Quantum effects on an assumed orthorhombic phase of crystalline PE were studied in reference [40] by means of path-integral Monte Carlo (PIMC) simulations. The low temperature range, from 25K to 300K, was explored in the NPT ensemble by looking at PE chains of two lengths, C\(_{12}\) and C\(_{24}\) (with a total number of atoms of 432 and 864). Structural parameters such as the lattice constants, bond lengths, and bond angles and their dependence with temperature were analyzed, showing their reduction as the temperature is decreased.

By performing MD simulations of PE chains of various lengths, using a united-atom model with torsional barriers of 2, 3 and 6 Kcal/mol, with the remaining parameters being taken directly from the Dreiding forcefield, the chain folding phenomenon was observed [41, 42]. They found that during simulations with a torsional barrier of 2kcal/mol, PE chain folding occurs when the chains have more than 150 CH\(_2\) units, long enough that the average inter-molecular van der Waals energy is low enough to compensate the increase in energy owing to torsion angle and bond angle deformation in the folds. The reported behaviour was in good agreement with the experimental observations [43].

More recently, multi-phase polyethylene morphologies including crystalline, amorphous, lamellae and interfacial regions have been studied using MD [44, 45]. As discussed in reference [2], a polyethylene sample may contain amorphous, lamellae and crystalline regions of various sizes depending on the material processing details. In reference [44], model crystalline and lamellae regions were created by geometry optimization using the all-atom COMPASS forcefield in Materials Studio 5.5 [46]. For the lamellae regions, gauche and anti-gauche defects were introduced into ten all-trans PE chains with 552 CH\(_2\) units in order to enable them to fold back and forth upon themselves, resulting in a lamellar thickness of around 5.0 nm. The structures were then imported into LAMMPS simulation package [47], and equilibrated using a united-atom force field [48] optimized for long chain n-alkanes. The amorphous and interfacial regions were prepared by melting the corresponding lamellae blocks either in NPT or N\(\mu\)T ensembles with a time-varying thermostat in LAMMPS. The resulting morphologies were in good agreement with experiment.

As observed experimentally [49], polyethylene contains a significant number of nanometre sized voids. In order to explore their influence on excess electrons, nanometre sized voids were produced in the simulated amorphous phase by expanding a test particle [44]. By calculating the Gibbs free energy to create different radius of spherical voids in amorphous PE, a surface tension of 36 mN/M was obtained, in good agreement with the experimental value of 35.7 mN/M at 20 °C for linear HDPE [50].

#### 3.2 Electronic Properties of PE

Surface trapping states of PE was studied [51] using DFT in the gradient-corrected local density approximation (LDA), supplemented with empirical long-range tails in order to properly account for van der Waals forces. Two PE surfaces were considered, representative of two classes of surfaces that differ in the orientation of the PE chains with respect to the planar surface i.e. with the chains oriented either parallel or perpendicular to the surface vector. A negative electron-affinity was found, with values of -0.17eV and -0.10 eV for surfaces with chains perpendicular and parallel to the surface, respectively. Negative electron-affinities are in agreement with experiment and other simulation results (see below), though in this DFT study the analysis is based on the identification of KS eigenvalues (section 2) with single-particle energy levels.

Dynamical DFT simulations have also been performed [52] within the framework of the Car-Parrinello (CP) technique. Calculations were carried out with an injected electron neutralized by a background charge in a crystalline cell with four PE chains (each of them with 7 CH\(_2\) units), starting from an equilibrium configuration. A similar calculation was also
performed with an injected hole. A self-trapped state for the injected electron was found, localizing near an inter-chain area which involved a pair of trans-gauche defects. The hole was observed to remain delocalized throughout the simulation cell, being of an intra-chain nature.

In reference [53], an electron-hole pair, an exciton, was injected in a similar crystalline cell, being observed to be long-lived, displaying no apparent direct channel for non-radiative recombination. In contrast, very recent DFT calculations [54] have found a direct channel for the recombination of the exciton via the breaking of a C-H bond. The discrepancy with the results in [53] might be due to the exchange-correlation functional employed or the different methods chosen to improve the deficiencies of standard DFT. However, reference [54] presents a convincing scenario in which the recombination of an electron-hole pair under electrical stress could lead to degradation of PE.

Research focused on the electron trapping density of states identifying both physical and chemical defects were also carried out. To model the physical defects [55], MD simulations of amorphous tridecane (n-C\textsubscript{13}H\textsubscript{28}) were used as a model of short sections of PE to generate localized conformational defects. The trap energy was defined as the difference between the electron affinity of the wax molecule with and without conformational defects (i.e. not all trans) as $E_t = EA(C_{13}H_{28}) - EA(n-C_{13}H_{28})\text{un-trans.}$ The electron affinity was obtained from DFT as implemented in the code DMol [56]. A similar methodology was used to estimate trapping energies for chemical defects and an approximate excess electron density of states obtained. Other workers have used similar methods [57-59]. Some results for chemical defects are given in table 1, clearly there are some very deep traps (>0.5 eV) but also some shallow traps (<0.5 eV) which overlap in energy with those caused by conformational defects.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Trap depth (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-decanone (C\textsubscript{10}H\textsubscript{10})</td>
<td>-0.453</td>
</tr>
<tr>
<td>5-vinyl (C\textsubscript{4}H\textsubscript{8})</td>
<td>-0.122</td>
</tr>
<tr>
<td>5-decanol (C\textsubscript{10}H\textsubscript{10})</td>
<td>-0.186</td>
</tr>
<tr>
<td>5-decanal (C\textsubscript{10}H\textsubscript{10})</td>
<td>-0.445</td>
</tr>
<tr>
<td>Cumylalcohol (C\textsubscript{9}H\textsubscript{12})</td>
<td>-0.28</td>
</tr>
<tr>
<td>Acetophenone (C\textsubscript{8}H\textsubscript{8})</td>
<td>-0.9</td>
</tr>
<tr>
<td>Alpha-methylstyrene (C\textsubscript{8}H\textsubscript{11})</td>
<td>-1.53</td>
</tr>
<tr>
<td>Cumene (C\textsubscript{8}H\textsubscript{12})</td>
<td>-0.04</td>
</tr>
</tbody>
</table>

A more detailed study of the physical defects in amorphous and crystalline PE and their effect on excess electrons was carried out [27, 60] using Lanczos method to compute the excess electronic states of static configurations taken from MD simulations (for ethane, methane, propane, crystalline PE, and amorphous PE using a single chain with 300 CH\textsubscript{2} units). The pseudo-potential, describing the interaction of the excess electron with the atoms, was adjusted to fit the experimental data for the threshold of conduction in fluid ethane and propane. It contained short-range repulsive potentials, and an attractive part, which accounts for the polarization interaction between the excess electron and the dielectric, based on multi-centre polarizabilities obtained by fully ab initio methods. The amorphous PE samples showed Anderson localization [61], with a mobility edge, separating localized and delocalized states. The electronic density of states (DoS) and the electron mobility at the amorphous phase was calculated using the Kubo–Greenwood equation [62]. Further work [63], using non-adiabatic simulations of an excess electron in amorphous PE, permitting the deformation of the material due to the presence of the electron, showed the spontaneous formation of localized small polaron states in which the electron is confined. Despite allowing non-adiabatic transitions by using Tully’s surface hopping algorithm [64], the simulations showed mainly adiabatic dynamics.

Very recently, the analysis of references [27, 60] has been extended to larger systems and more representative regions of PE, including lamellae and interfaces between lamellae and amorphous [44, 45]. A clear correlation between local atomic density (calculated as the number of CH\textsubscript{2} united atoms per unit volume in a cube of side 0.5nm at a given position) and electron probability has been found in all PE phases, showing that the electron is more likely to localize in lower density regions.

Simple visual inspection of the localized states showed that the electron is sitting at regions with a reduced local atomic density, as illustrated in Figure 1. In the interfaces between lamellae and amorphous phases, the lowest atomic density values are found either in the amorphous regions or near the interface [45]. Figure 2 shows a typical ground state localized in that area.

In order to make this association more quantitative, we have calculated the Pearson correlation coefficient between the probability density of the excess electron and the local PE atomic density [45]. This dimensionless coefficient is usually defined as

$$r = \frac{\sum_{i=1}^{n}(X_i - \overline{X})(Y_i - \overline{Y})}{\sqrt{\sum_{i=1}^{n}(X_i - \overline{X})^2} \sqrt{\sum_{i=1}^{n}(Y_i - \overline{Y})^2}}$$  \hspace{1cm} (12)$$

where \(X\) and \(Y\) are the variables of interest, here to be identified with the electronic probability density (the square of the wave function) in a given electronic state and the local atomic density. The index \(i\) in (12) runs through the 3D grid points in the system. The coefficient \(r\) measures the strength of the linear association between both variables: a value of zero indicates no correlation, while nonzero values show correlation or anti-correlation. In the amorphous phase, starting with the ground state, the Pearson correlation shows a significant anti-correlation value of about -0.4 in regions of the material where the electron has a probability of 90%. The
next first 10 excited states, still below the mobility edge, show a correlation of about -0.2, while for the first 10 states above the mobility edge the correlation is reduced to -0.1, which is still significant, especially taking into account that these latter states are extended.

Another way to look at this correlation is presented in reference [44]. First, the free volume or cavity number distribution in the system is computed after identifying cavities with local minima of the atomic density. A radius is assigned to each cavity based on the radial distribution function from the centre of the cavity. Then, each localized state is associated with a cavity, providing an average energy and localization length as a function of the cavity radius. The cavity number distribution can be then mapped to produce a prediction of the density of states (DoS) based on the free volume distribution. This prediction is shown [44] to accurately predict the observed DoS in the three PE bulk phases studied- amorphous, lamellae and crystalline, when a correction based on the localization length is carried out. The correction is needed because localized states with higher energy values are not restricted to single cavities, instead extending through adjacent ones.

4 DISCUSSION AND CONCLUSIONS

As we have seen, molecular modeling techniques have enabled some progress to be made in understanding the fundamentals of charge trapping and electron transport in polyethylene. Previous work has identified physical, (conformational) [52, 55] and chemical (impurities and decomposition products) [57-59] electron traps in models of polyethylene and characterized them using ab initio methods. Some of this information was used to create a preliminary distribution function representing the density of trap states (DoS) as a function of electron energy and employed in a Monte Carlo simulation to predict the current-voltage [65] characteristics of model polyethylene, showing how once the DoS is known the electrical properties can be estimated. A more complete treatment of conformational trapping using the fast Fourier transform block Lanczos diagonalization algorithm, described in section 2, has resulted in densities of states for the different morphologies found in polyethylene [44, 45]. A simple theory based on this work now links the physical traps in polyethylene to nanovoids associated with regions of low density in the material.

Our current understanding of conformational excess electron trapping based on molecular modeling [44] has localized states extending below the mobility edge (above which the electrons are delocalized), at -0.1±0.1eV with respect to the vacuum level. These trap states with localization lengths between 0.3 and 1.2nm have energies as low as -0.4±0.1eV in the amorphous and interfacial regions of polyethylene with more positive values in lamella structures. Note crystalline regions have a mobility edge at +0.46 ±0.1eV, so we would expect transport by excess electrons excited above the mobility edge to delocalized states to be predominantly through amorphous regions if they percolate the sample.
At low electron concentrations, transport above the mobility edge will be dominated by the filling of the deepest traps which are likely to be chemical traps ~ -1eV. As the deepest traps are filled the excess electron mobility will increase dramatically towards a value that corresponds to multiple trapping between more shallow conformational traps [65].

Nonadiabatic simulations (Section 2) of an excess electron in amorphous PE at room temperature showed the spontaneous formation of localized small polaron states in which the electron was confined in a spherically shaped region with a typical localization length of 0.5nm. The self-trapping energy was ~0.06 ±0.03eV, with a lifetime on the time scale of a few tens of picoseconds. The smallness of the self-trapping energy is consistent with an adiabatic hopping mechanism assisted by phonons, as observed in the simulations. The contribution to the mobility due to hopping between these self-trapped states may well be of same order of magnitude as the mobility due to excited electrons above the mobility edge.

The model studies confirm the picture of deep chemical and shallow conformational electron traps current in [2]. They also however provide the means to go further and begin to investigate the influence of trapped electrons, the space charge, on the surrounding material. For example there has been very little work on the fate of the energy given up by trapped electrons in polyethylene (but see [58] for trapped excitons) which may alter the environment leading to local damage and ageing. The very recent DFT results [54] suggesting PE degradation by exciton recombination provide an interesting scenario that needs to be confirmed. In this regard, though molecular modeling studies provide a great insight in the understanding of polymeric dielectrics, these studies need to be assisted by experiments on very pure polymeric materials (including polyethylene) in which the effects of chemical traps from additive chemicals and/or radiation damage on electrical properties can be separated from those of physical traps related to the various polymer morphologies present in real materials.

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References
