Electrical characterization of organic (amorphous) materials

Peter Stallinga
Universidade do Algarve / Center for Electronics Optoelectronics and Telecommunications

E-MRS Warszawa
19/IX/2013 9:00-9:30
Symp. F 12.1
Abstract (for finding this doc on Google)

Electrical characterization of organic (amorphous) electronic materials

Organic electronic materials, or amorphous electronic materials in general, have relatively low conductivity and this limits their application to the low-frequency electronics market. To describe electronic conduction in these materials it is common to use Percolation or (Variable Range) Hopping Theory (the two being equivalent). This is an inheritance from the earlier organic materials that were invariably insulators, where conduction was a perturbation -- movement of charge was a rare event. It will be argued here that for electronic materials, instead, it is better to revert to classical semiconductor theories, like Band Theory[1]. If we include a large density of traps in the energy system, all observed phenomena are easily explained. This includes 1) Strong temperature dependent charge-carrier mobility, 2) Field-dependent mobility, 3) Anomalous transient behavior. Moreover, it is consistent with observations in many types of devices, ranging from two-terminal devices such as diodes to three-terminal devices such as thin-film transistors[2].

Introduction to organic electronics

The conduction model

Amorphous materials

Theories:
- (Variable Range) Hopping
- Percolation Theory
- (Good old) Band Theory

Devices/measurements
- Thin Film Transistors
- (Anomalous) Transients (DLTS, etc.)
- Temperature and bias dependent mobility
Conjugated organics have paths with alternating single and double bonds.

**Conjugation:**
*Delocalization* and opening of energy gap makes them semiconductors.

**Oligomers** (small molecules)
Good for FETs
(High mob. because of better crystallinity)

**Polymers**
Good for LEDs
(Because of fewer impurities)
The solid-state **physics** tripartition

- Making excellent samples
  - Float zone, Czochralski, etc.
  - Bragg
- Developing excellent theories
  - Drude, Hartree-Fock, Bloch, Landau

**Growth**

**Theory**

**Experiment**

- Doing excellent measurements
  - DLTS, Raman, FTIR, PL, AFM, STM, ....

Keywords: predictability & reproducibility
The solid-state physics tripartition

Initially 1/2-cons were ignored. “Shitty materials”

Later: Exactly this dependence on impurities makes them extremely powerful

Semiconductor industry. Brought to you by physics!
Semiconductor (device) theory

(for electronic **device physics**)
The entire band structure can **effectively** be summarized as

- NV full levels at EV
- NC levels at EC
- Effective mass (and $\mu$)
- Donor and acceptor levels

Sze: 800 pages of approximation good enough to describe all electronic ½-con devices
The chemistry tripartition

Most effort spent here
**Synthesis & Growth**
(Very complicated!
Much worse than non-organics)
**Nano**-structures!

Chemistry dominates organic “plastic” material research
Plastics traditionally non-conductive

Using theories of non-conductive materials (e.g. Hopping)

Doping is not relevant (yet)

**Ex.: Sub-threshold swing**
(which is not a parameter in TFTs!)

Experimental equipment is black boxes with buttons

= Use of non-conduction models (also for conductive materials!)
Conduction Mechanism Theories

Amorphous ('disordered') materials are mostly modeled by **Variable Range Hopping / Percolation Theory** (Evros, Shklovskii, Mott, 1950's)

Crystalline materials are invariably modeled by **Band Theory**

Why? Are these materials so different?

Crystalline

Poly-crystalline

Amorphous

Band Theory

Percolation Theory / (Variable Range) Hopping
Both crystalline and amorphous materials have covalent bonds. In crystals they give rise to band structure ....

Crystals were only used because they are easier to produce reproducibly. **Crystal theories apply to non-crystals too!** (Ioffe, Regel, and Gubanov)
Crystal schmystal

By far the most materials in nature are amorphous

Crystalline materials: Scientists not describing nature, but scientists creating a new universe to study
Get out of the sandbox and start studying nature!

**Crystalline materials are not going to save the planet**
(from the Global Warming threat [if it is not a scam*])

Poly-crystalline silicon solar panel:
30 years to return energy invested (no electronics can stand 30 years outside in the weather!)
There has not yet been renewable energies (except hydro) that saved oil!

*: Visit www.stallinga.org/Climate where the scam is exposed
Non crystals

“The fact that devices were invented and theories developed for crystals, does not mean that the theories are valid **only** for crystals”

- (P.S. 2011)

“A periodic electric field of the lattice is not essential for the occurrence of typical semiconducting properties and the band model may be applied also in the case in which there is a loss of periodicity of the lattice”

- Ioffe, Regel and Gubanov [1]

Both crystalline and amorphous materials have covalent bonds. In crystals they give rise to **band structure** .... so ... also in **amorphous materials**!

Band Theory for Amorphous materials

Crystal:

```
Crystal:

E (eV)

<table>
<thead>
<tr>
<th>k</th>
</tr>
</thead>
<tbody>
<tr>
<td>L</td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td>E (eV)</td>
</tr>
<tr>
<td>0</td>
</tr>
<tr>
<td>-10</td>
</tr>
</tbody>
</table>

'clean' electronic levels
```

Amorphous:

```
Amorphous:

E (eV)

<table>
<thead>
<tr>
<th>k</th>
</tr>
</thead>
<tbody>
<tr>
<td>L</td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td>E (eV)</td>
</tr>
<tr>
<td>0</td>
</tr>
<tr>
<td>-10</td>
</tr>
</tbody>
</table>

'messy' electronic levels
```
Amorphous Band Diagram

Energy diagram consists of
- conductive (delocalized) band states
- (localized) trap states

Successfully used to describe amorphous silicon thin-film transistors [1]

Applied to organic thin-film transistors “The Algarve TFT Model” [1]

1) $\rho = \rho_T + n$
2) $\rho = -V_g C_{ox}$
3) $I = -n \mu dV(x)/dx$

Si: $\gamma = 1$
Organics: can be 7 (not a small perturbation!)

There are **no contact effects** [1]
People often imagine 'Schottky barriers' at the metal – ½-con interface
  This does not make sense
People often imagine resistors at the contacts
  This does not make sense
All observed 'anomalies' are caused by traps

Transistor (accumulation TFT) can be made of **any material**. Not limited to ½ conductors. Can be a potato.
We made a **metal** transistor [2].

Trap 'depth' depends on electric field [1]:

\[ \mu = \mu_0 \exp \left( -\frac{E_A}{kT} \right) \]

\[ E_A = E_T - \sqrt{\frac{qE}{\pi \varepsilon}} \]

FIG. 1. Potential energy as a function of distance from the positive ion. Full line, without an external field, dotted line in the presence of the field.

full line represents the normal potential energy of the electron as a function of the distance from the positive ion while the dotted line represents the same quantity in the presence of the field. The height of the potential barrier is lowered in the field by the amount

\[ \Delta U = eE_r_0 + e^2/\varepsilon r_0, \]

where \( r_0 \), the distance to the maximum from the ion, is given by \( e^2/\varepsilon r_0^2 = eE \). Thus \( r_0 = (e/eF)^{1/2} \) and

\[ \Delta U = 2eE r_0 = 2e(eF/e)^{1/2}. \]

Crystalline Silicon:
- Traps scarce
- Traps discrete
- Filling immediate
- Emptying thermally activated

\[ \tau = \exp \left( - \frac{E_A}{kT} \right) \]

\[ C(t) = \Delta C \exp \left( - \frac{t}{\tau} \right) + C_0 \]

DLTS: measure \( \tau \) as a function of \( T \) and you get \( E_A \)
Transients

Amorphous materials:
- Traps abundant
- Traps distributed in energy
- Filling thermally activated
- Emptying thermally activated

\[ \tau = \exp \left( -\frac{E_A}{kT} \right) \]

\[ C(t) = \Delta C \exp \left( -\frac{t}{\tau} \right) + C_0 \]

Abundant level: DLTS: \( C(t) \) is not exponential. \( C^2 \) is!

Distributed \( E_A \) and \( \tau \): transient is not exponential. DLTS will be a mess. LT-DLTS will be a mess.

These scientists from the crystalline era had an easy life indeed!
Transients

Distributed \( E_A \) and \( \tau \): transient is not exponential. DLTS will be a mess. LT-DLTS will be a mess. Transient will be convolution of transients [1] ... ... resulting theoretically in a power law, for example

\[
I_{ds}(t) = t^{-\alpha}
\]

In practice, empirically, transients are more stretched-exponential [2]. Without, after nearly two centuries, no theoretical justification:

\[
I_{ds}(t) = \exp\left[-\left(\frac{t}{\tau}\right)^\beta\right]
\]

"In any finite critical system, it is well known that the power-law description must give way to another regime" [1]

Any (finite, 'alpha-stable') distribution of relaxation times will result in a stretched exponential [2]

\[ I_{ds}(t) = t^{-\alpha} \]  

\[ I_{ds}(t) = \exp\left[-\left(\frac{t}{\tau}\right)^\beta\right] \]

Effects of abundant traps (amorphous)

a) transfer (non-lin)

\[ I_d \sim V_g^{\alpha+1} \]

b) output (non-lin)

\[ I_d \sim V_{ds} \]

P.F.

c) Meyer-Neldel

\[ \mu_{FET} \sim 1/T \]

V_{gn}

V_{g1}

d) ToF (non rect.)

\[ \log(I) \sim \log(t) \]

\[ \log(I) \sim \log(t) \]

Kohlrausch (1847)

\[ \log(I) \sim t \]

\[ I_d \sim V_g \]

--- Trap-free  --- single trap  --- with traps
Summary

Organic (amorphous) electronic do have future outside sandbox

Percolation / Variable Range Hopping compared with Band Theory
→ Band Theory wins (with traps)

Organic Electronic Devices Characterization
- Thin Film Transistor
- Transients
Very complex! Not a black box plug-and-play measurement

Percolation theory or hopping theory, used to describe the electronic behavior of devices, is very popular in the chemistry-dominated research area of organic electronics. The reason is that chemists think in units (“molecules”) such as the phenyl ring in a polymer, with specific energy levels (i.e., the lowest unoccupied molecular orbital (LUMO) and highest occupied molecular orbital (HOMO)), and conduction of charge is considered a perturbation to the molecular electronic levels. A typical organic semiconductor consists of a one-dimensional chain — the conjugation backbone — along which current is visualized. Thinking in terms of conduction paths assumes very natural and three-dimensional conduction in terms of “hopping” to neighboring conjugated segments means an obvious approach. Charges reside on specific sites and only occasionally make a jump to neighboring units. This way of thinking has successfully described the low-conductivity end of measurement sets, but also because we will get bogged down in equations. Yet, it is useful to introduce the models globally, the function of models that is. That is the scope of this Essay. The equations will be kept here in a minimum.

The variable range hopping (VRH) theory is based on the idea that charges are localized but can every now and then jump (“hop”) to another localized state. The probability of hopping between two states of spatial separation $s$ and energy separation $W$ is given by

$$P(E,W) = \exp \left( -\frac{W}{kT} \right)$$

with $k$ Boltzmann constant, and $a$ a constant. The total conductivity can then be found as the integral over all energy states and distances, which can be worked out into the previously mentioned Mott equation (Mott 1949 and Fillmore 1968).

Comparison of theories in Adv. Mat. 23, 3356 (2011)

Special thanks to Henrique Gomes and the University of The Algarve and Portugal in general

10Q4ur@+jon