The Effects of Crosslinking Byproducts on the Electrical Properties of Low Density Polyethylene

by

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ABSTRACT

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THE EFFECT OF CROSSLINKING BYPRODUCTS ON THE ELECTRICAL PROPERTIES OF LOW DENSITY POLYETHYLENE

By Nuriziani Hussin

Crosslinked polyethylene (XLPE) is widely used for high voltage insulation in power transmission systems. However, it has been found that, after crosslinking with Dicumyl Peroxide (DCP), the crosslinking byproducts such as acetophenone, α-methylstyrene and cumyl alcohol have a significant influence on electrical properties of XLPE power cables. This thesis distinguished the individual contribution of the crosslinking byproducts on space charge formation, dielectric properties, dc conductivity as well as the ac breakdown strength. Percentage weight increases as well as the Fourier Transform Infrared (FTIR) spectrum were used to monitor the chemical level in the soaked samples. Despite high concentration of byproducts in the LDPE film compared to practical, the measurement results have successfully reveal the contribution of each byproduct on the electrical properties. It should be noted that some consideration should be taken when taking the quantitative value from the result obtained.

Space charge accumulation was measured using the pulsed electroacoustic (PEA) technique. Homocharges are observed in acetophenone and α-methylstyrene soaked LDPE. Meanwhile heterocharge formed in cumyl alcohol soaked LDPE. From the charge decay profile in dc condition, these chemicals are observed to assist the transportation of the charges in the sample bulk due to shallow traps from the byproducts. These shallow traps assist the trapping process into deep traps when ac field is applied to the byproduct soaked LDPE. As a result, more charges trapped in deep traps were found in soaked LDPE compared to clean LDPE. In addition, from the space charge measurement in ac condition,
it is proved that the amount of charge trapped in deep traps also depends on the population of shallow traps in the polymer which is contradicted to the literature where the byproducts are normally associated to the deep traps.

Permittivity values of acetophenone, α-methylstyrene soaked LDPE and cumyl alcohol are slightly higher than permittivity value of the clean untreated LDPE. Cumyl alcohol soaked LDPE has higher dielectric loss at lower frequency due to Maxwell-Wagner-Sillars polarisation as well as space charge polarisation effect. In contrast, acetophenone does not change the dielectric loss value and α-methylstyrene gives very little effect. These byproducts have very high dc conductivity values. It is also proposed that the chemicals provide shallow traps that aid the charge movement and this is consistent with the mobility values that calculated from the conduction current result. The ac breakdown results however show no significant difference from the breakdown strength of clean LDPE. Based on ac space charge results and ac breakdown test results, it is concluded that the byproducts have little effects in ac condition.
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Authors Declaration

I, Nuriziani Hussin, declare that the thesis entitled “The Effects of Crosslinking Byproducts on The Electrical Properties of Low Density Polyethylene”, and the work presented in the thesis are both my own and have been generated by me as a result of my own original research. I confirm that:

1. This work was done wholly in candidature for a research degree at the University of Southampton;
2. Where any part of this thesis has previously been submitted for a degree or any other qualification at the University of Southampton or any other institution, that this has been clearly attributed;
3. Where I have consulted the published work of others, this is always clearly attributed;
4. Where the work of others has been quoted the source is always given, and with the exception of such quotations this thesis is entirely my own work;
5. I have acknowledged all main sources of help;
6. Where the thesis is based on work done by myself jointly with others, I have made clear exactly what contributions were my own; Some of this work have been published as:


• Hussin, N and Chen, G (2009) *Analysis of Space Charge Formation in LDPE in the Presence of Crosslinking Byproducts.* Submitted to IEEE Journal of Transactions on Dielectrics and Electrical Insulation (Accepted)


Signed: ___________________________ Date: ____________________

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Definitions and Abbreviations Used

Definitions

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<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
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<tr>
<td>E</td>
<td>Electric field (V/m)</td>
</tr>
<tr>
<td>V</td>
<td>Voltage/potential (V)</td>
</tr>
<tr>
<td>i(t)</td>
<td>Current (A)</td>
</tr>
<tr>
<td>u</td>
<td>Velocity field</td>
</tr>
<tr>
<td>t</td>
<td>Time</td>
</tr>
<tr>
<td>e</td>
<td>Electron charge</td>
</tr>
<tr>
<td>eV</td>
<td>Electron volt</td>
</tr>
<tr>
<td>(\varepsilon_o)</td>
<td>Permittivity of vacuum</td>
</tr>
<tr>
<td>(\varepsilon_r)</td>
<td>Relative permittivity</td>
</tr>
<tr>
<td>(\varepsilon')</td>
<td>Dielectric constant / real permittivity</td>
</tr>
<tr>
<td>(\varepsilon'')</td>
<td>Imaginary part of permittivity</td>
</tr>
<tr>
<td>(\varepsilon^{*})</td>
<td>Complex permittivity</td>
</tr>
<tr>
<td>(\varepsilon'_0)</td>
<td>Real permittivity at zero frequency</td>
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<tr>
<td>(\varepsilon'_\infty)</td>
<td>Real permittivity at limiting high frequency</td>
</tr>
<tr>
<td>(\tan \delta)</td>
<td>Dielectric loss tangent</td>
</tr>
<tr>
<td>(\rho)</td>
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<tr>
<td>(\phi)</td>
<td>Thermal activation energy (eV)</td>
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<tr>
<td>(\Phi)</td>
<td>Potential barrier</td>
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<tr>
<td>(\mu)</td>
<td>Mobility (m(^2)/Vs)</td>
</tr>
<tr>
<td>(\sigma)</td>
<td>Conductivity (S/m)</td>
</tr>
<tr>
<td>T</td>
<td>Temperature (°K)</td>
</tr>
<tr>
<td>(c_p(t))</td>
<td>Pulse voltage (V)</td>
</tr>
<tr>
<td>P(x,t)</td>
<td>Pressure pulse</td>
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ρ(x,t) Total charge density (C/m$^3$)
q(x,t) Total charge (C)
k Boltzmann constant (8.62 x 10$^{-5}$ eV/K)
h Plank’s constant
f Frequency
ω Angular frequency
J Current density
m Mass
d Thickness
Dn Fick’s diffusion coefficient
n Number density of charge
$f_{\text{pulse}}$ Pulse generator frequency
$f_{\text{ac}}$ Alternating current frequency
$E_s/V_s$ Stressing Field / Voltage in PEA measurement
$V_p$ Pulse voltage in PEA measurement
$C^*$ Complex capacitance
$C_0$ Capacitance in air gap
Z Impedance
R Resistance
α Weibull scale parameter
β Weibull shape parameter

Abbreviations

PE Polyethylene
LDPE Low density polyethylene
XLPE Cross-linked polyethylene
DCP Dicumyl peroxide
HDPE High density polyethylene
MDPE Medium density polyethylene
TR-XLPE Tree retarded crosslinked polyethylene
C Carbon
H  Hydrogen
O  Oxygen
Si  Silane
Al  Aluminium
DC or d.c  Direct current
AC or a.c  Alternating current
HV  High voltage
HVAC  High voltage alternating current
HVDC  High voltage direct current
LIMM  Laser intensity modulation method
LIPP  Laser induced pressure pulse
LLDPE  Linear low density polyethylene
PEA  Pulsed electro-acoustic
TSM  Thermal Step method
PMMA  Polyethyl methacrylate
PVDF  Polyvinylidene fluoride
SCLC  Space charge limited current
V_{TFL}  Trap-filled-limit
FFT  Fast Fourier transforms
IFFT  Inverse Fast Fourier transforms
GPIB  General Purpose interface bus
TTL  Transistor-transistor logic
GPIB  General Purpose interface bus
STP  Shielded twisted pair
BNC  Bayonet Neill concelma
FTIR  Fourier transforms infrared
OS  Open system
CS  Close system
Volt off  Measurement during short circuit
Volt on  Measurement during close circuit
a+α  acetophenone plus α-methylstyrene in LDPE
a+c  acetophenone plus cumyl alcohol in LDPE
α+c  α-methylstyrene plus cumyl alcohol in LDPE
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<tr>
<th>$E_{\text{RMS}}$</th>
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<tr>
<td>MWS</td>
<td>Maxwell-Wagner-Sillars</td>
</tr>
<tr>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>PDF</td>
<td>Probability density function</td>
</tr>
<tr>
<td>ML</td>
<td>Maximum likelihood</td>
</tr>
<tr>
<td>$E_b$</td>
<td>Breakdown strength</td>
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Chapter 1  Introduction

1.1 Overview of Research

The development of underground cable started with the telegraph lines. The earliest use of continuously insulated conductor was recorded in 1812 by B. Schilling when an electrical pulse was sent through a cable insulated with strips of India rubber. It was then expanded into the electrical lighting system where a steady flow of a considerable amount of energy was needed. T. A. Edison who planned his first installation for New York City decided to have an underground system distribution. During his time, copper rods insulated with wrapping of jute were used. This design gave a pleasing performance for low voltage service. This power distribution is still limited to the area capable of being supplied from one source if the regulation did not exceed the maximum bound.

In 1882, ac distribution was employed by L. Gaulard and J. D. Gibbs and during the early installation, the cable operated at 2,000 volt, the overhead was used due to their satisfactory performance and almost universally employed. In Washington and Chicago, overhead wires were prohibited, hence a number of underground lines were emplaced. Many insulation and method of installation were tried but the successfulness was little. However, in 1890, Ferranti had successfully installed the first line insulated with paper between Deptford to London for single-phase operation at 10,000 volt and some were still in use at the original voltage after more than 50 years. The conductors were insulated with wide strips of paper applied helically around the conductor before saturated with rosin-based oil.
Impregnated paper became most universal form of insulation cable for underground transmission due to its low dielectric loss, low dissipation factor and high dielectric strength. Impregnated paper insulation consists of several layers of paper tapes, wrapped helically around the conductor. The total wall of papers is subsequently heated, vacuum dried and impregnated with an insulating fluid. The impregnating fluid used changed from time to time. Circa 1925, impregnating compound changed from rosin-based compound to mineral oil or oil blended to obtain higher viscosity. Around 1983, polybutene replaced oil as the insulating fluid. The reliability of paper insulated as the insulating material was excellent. However, it required a high degree skill for appropriate splicing and terminating.

The insulation of cable system is then shifted towards polymer, particularly polyethylene (PE), as more development and research done in that area. Low Density Polyethylene (LDPE) was mainly used in the cable insulation system for its low cost, moisture and chemicals resistance, low temperature flexibility and excellent electrical properties. However, power cables typically operate at high temperatures and hence the insulation surrounding the conductor is subjected to elevated temperatures and a temperature gradient. With a melting point approximately at 110ºC, LDPE could not operate in the temperature range for paper-oil insulated cable. During the last decade, the voltage rating for underground cable has become increased due to higher electrical usage. In the future, the electrical stress will become increase and hence, the need for a better polymer that could withstand higher electrical stress has become vital.

1.1.1 High Voltage Cable

It was stated by the International Electrotechnical Commission (IEC 502) that the standard for HV level is the range of 35 kV <V<230 kV [1]. A full detail on the range of different voltage level is presented in Table 1-1. Different type and thickness of insulation material is used for each operating voltage due to the thermal and mechanical properties of the material. For example, based on IEC 502 specification, at rated voltage of 20kV, the insulation thickness for XLPE cable is 5.5mm meanwhile for PVC cable is 6.4mm. However not all countries follow the IEC standard. In country like Japan, the insulation thickness specified by the national standard for XLPE insulated cables is greater than the
IEC but for export purpose, the IEC value is followed. In USA, which have different voltage rating, the IEC standard for insulation thickness is not applicable [2].

Table 1-1: Classification of the operating voltage levels [1].

<table>
<thead>
<tr>
<th>System</th>
<th>Nominal voltage ($V_r$) kV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Voltage (LV)</td>
<td>$V_r &lt; 1$</td>
</tr>
<tr>
<td>Medium Voltage (MV)</td>
<td>$1 &lt; V_r &lt; 35$</td>
</tr>
<tr>
<td>High Voltage (HV)</td>
<td>$35 &lt; V_r &lt; 230$</td>
</tr>
<tr>
<td>Extra High Voltage (EHV)</td>
<td>$230 &lt; V_r &lt; 800$</td>
</tr>
</tbody>
</table>

In 1955, Crosslinked polyethylene (XLPE) was first patented and with the crosslinking technique, the thermal stability of PE is improved [3]. XLPE has now been used widely in high voltage insulation system. Compared to LDPE, XLPE has slightly higher dielectric loss, lasts longer and shows less moisture sensitive [4]. Cables with XLPE insulation for operating voltage up to 132 kV have now has been established widely. The use of metallic barrier which usually an extruded sheath has been introduced to avoid the incidence of water trees in the cable. The incentive of using XLPE has allow the cable to operate at 90°C compared to 70°C for LDPE and 80°C for HDPE [2].

Despite of its ability to withstand high temperature, it is undeniable that XLPE has some other weaknesses as a cable insulator. Easy accumulation of space charge in XLPE for instance, become the main anxiety in employing this material in insulation system [5]. It was considered that the crosslinking byproducts of well used crosslinking agent, Dicumyl Peroxide (DCP), antioxidant or the crosslink polymer morphology could generate space charge in the insulation material [6]. Through research N. Hirai reported that the crosslinked structure has no distinct trapping property [7]. Sulphur-containing antioxidant in the combination effect of acetophenone creates heterocharges [8]. The antioxidant by itself cannot trap any charges in the bulk. The trapping of the free radical from the crosslinking process is sometime inevitable, due to viscosity effect. As a result, the crosslinking is not completed and the residual peroxide as well as the byproducts could become a major factor to degradation process.
DCP which is widely used as the crosslinking agent creates volatile crosslinking by-products such as acetophenone, cumyl alcohol, α-methylstyrene and cumene which also affect the insulation properties. These byproducts are said to cause charges trapped in the XLPE. Other than causing space charge formation in the sample, the byproducts also have an effect on other electrical properties of XLPE like increasing the tan δ value, temporary increase in breakdown strength, and so forth [3].

In practice, degassing is introduced after the crosslinking process to remove the crosslinking byproducts from the insulation by subjecting the XLPE cable to moderately high temperature for a certain period. Ideally it is possible to remove all byproducts in all sample. However the times involved are rather long even for small cable. For example, it took 500 hours for acetophenone to be degassed to 10% of its’ original concentration in an average MV cable (approximately 5.5mm thick) [3]. Although degassing can occur freely from the outer surface of the insulation, the process is constrained from the inner surface. Hence the byproducts are more likely to stay in the insulation nearer to the conductor. Degassing process may take longer for a thicker sample and even much slower as the outer insulation shield (semiconductor shield) is added to the cable. Temperature could be increased to fasten the degassing process. However, without extra care, very high temperature may result to damage to the core through thermal expansion and softening the insulator. In addition, it was reported that degassing may also change the morphology of the dielectric [9].

1.2 Research Objective and Aims

It is clear that the main advantages of XLPE are its good electrical, mechanical and thermal properties. However, due to the existence of the impurities in XLPE including the crosslinking byproducts, researches have now looking for other material that could match the excellent properties of XLPE. However, the knowledge that we have on each of the impurities and how they actually cause problem is very little. It is believed that by understanding on how these individual impurities, in this case the crosslinking byproducts affect the electrical properties of XLPE, some improvement could be done on the conventional XLPE cable. Research on the effect of water molecules on the formation of water treeing in XLPE has lead to the production tree-retarded crosslinked polyethylene
(TR-XLPE). It is hope that by doing this research, comparable improvement could be made on XLPE cable.

Each and every byproduct in XLPE affect differently to the polymer’s electrical properties and thus, the experiments will be conducted so that the effect of each of the byproducts can be distinguished. XLPE film consisted of a few parts which are the crosslinked chains, uncrosslinked chains, peroxides, volatile crosslinking byproducts, antioxidant as well as stabilizer. Each of the XLPE part may affect the electrical properties differently. On the basis of understanding the individual contribution of the crosslinking byproducts, it is important to have very consistent polymer base for the byproducts to reside in order to have a fair comparison between the byproducts. Due to that, instead of using XLPE as the base polymer, LDPE is chosen.

In this research, a few objectives have been outlined;

1. Crosslinking byproducts are known to be one of the major sources of space charge accumulation. It is hoped that by investigating the space charge profile of crosslinking byproducts in LDPE will give some understanding on the mechanism of charge formation in the polymer. Through space charge measurement, the trapping characteristics of polymer as well as the crosslinking byproducts could be revealed. We are hopping that by studying space charge profile during dc and ac stressing will give more understanding on how the electric field affects the trapping behaviour.

2. For any material to become a good insulator, the dielectric constant along with dielectric loss value should be low. Investigation on the dielectric spectroscopy of the crosslinking byproducts will tell us how this additives affecting these values. Also, from this measurement, the polarisation process that happens in the byproduct soaked sample could be studied.

3. An excellent insulation will have very small conductivity and most of the current is due to capacitive charging current that will decay by time. In this thesis, we aim to
study the changes in the conductivity value of LDPE in the presence of the crosslinking byproducts. From the result, we are aiming to calculate the mobility of the carriers in each byproducts soaked sample.

4. It is common to associate the space charge formation with breakdown strength of the polymer. After all, this is one of the reasons why space charge accumulation is unfavourable in cable insulation. Here, we are intended to compare the breakdown strength of the dielectrics with and without the byproducts. Hopefully we could see here to what extend these byproducts affecting the breakdown strength value.

5. From the results that obtained from all measurements stated above, we are aiming to compare the characteristics of each of the byproducts and how they individually influence the electrical properties of the LDPE.

1.3 Thesis Outline

This thesis consisted of a few chapters that are arranged as follows.

**Chapter 1** outlines the overview of the research as well as the objectives of the work.

**Chapter 2** presents the details about the material that will be used in this research, which is polyethylene. This chapter includes background knowledge on the crosslinking process and introduction to the crosslinking byproducts.

Since the main problem caused by the crosslinking byproducts is the space charge formation, **Chapter 3** describes the space charge phenomenon in the polymer including the conduction mechanism in polymer.

In **Chapter 4** the techniques used to measure space charge are explained and the fundamental concepts of the measurements are also presented.
In Chapter 5, the sample preparation which explains about the soaking process is reported. The similar procedures are applied to all samples before any measurement is conducted. In other words, this chapter will be referred by other chapters to explain on the samples that were used in the measurement.

Chapter 6 describes the space charge characteristics of the clean and soaked samples at dc and ac fields. Charge decay profiles of the samples are used to understand the trapping characteristics of the byproducts soaked LDPE.

Chapter 7 shows the dielectric spectroscopy of the liquid byproducts, clean LDPE as well as soaked LDPE. The results are compared and the polarisation effect is observed from the dielectric spectrum.

Conduction current measurement is presented in Chapter 8. In this chapter, the conductivity result is being related to the results obtained in space charge measurement. Also, the mobility of the carriers is calculated and compared.

Chapter 9 describes the breakdown measurement of the byproducts soaked LDPE. Results in this chapter will also be linked to the space charge results from Chapter 6.

Last but not least, Chapter 10 presents the conclusions and future work for this research.
Chapter 2  Polyethylene

2.1 Introduction

Polyethylene is produced since 1936 and until now, it becomes the most popular material used for insulation. It is produced at pressure up to thousand atmospheres, which gives a polymer of single chain type but with short branches along the main chain. Polyethylene is created from polymerization of ethylene where in this process the double bond in ethylene between carbon atoms is opened up and forms a long carbon chain, as shown in Figure 2-1.

\[ \sim \text{CH}_2\text{=CH}_2 \quad \xrightarrow{\text{polymerization}} \quad \sim \text{CH}_2\text{-CH}_2\text{-CH}_2\text{-CH}_2\text{-CH}_2 \sim \]

**Ethylene (gas)**  **Polyethylene (solid)**

Figure 2-1: The formation of polyethylene from ethylene.

As a thermoplastic, polyethylene melts to a liquid when heated and freezes to a brittle, very glassy state when cooled sufficiently. Polyethylene is categorized into groups by its density and branching. The density of PE is actually a measure of crystallinity. In term of manufacture, PE is classified into ‘high density’, ‘medium density’, ‘low density’ or ‘linear low density’. Method of polyethylene manufacture controls the exact chemical structure, which in turn controls the properties of PE. The polymer chain is not always linear. In polymerization, the process always leads to side chain attached to the long main chain. This is called chain branching and it contributes to the increase of molecular weight. The length and the distribution on the branches affect physical properties as well as the ability of the PE to crystallize.
In reality, chains in PE are not ‘straight’ as in Figure 2-2 above. The polymer has high tendency to coil or in other words, has a random configuration which is influenced by the branching. These entanglements will contribute to some resistance to movement when the chains are pulled apart. Hence, the entanglements influence the mechanical properties of the polymer. The ability of the polymer chain to coil as well as tendency to align themselves relative to each other, make PE grouped into the semicrystalline polymer. The chain portions that are aligned is called ‘crystalline’ and the chain portions that coil are described as ‘amorphous’ as illustrated in Figure 2-3.
The crystalline regions give PE many good properties for cable insulation, such as toughness, high modulus and moisture and gas permeation resistance. The ‘tighter’ chain packing in the aligned portions also increases the density of the polymer. Hence, increased crystallinity also depicts higher density. On the other hand, amorphous regions increase the flexibility, ductility and facilitate processing.

High Density Polyethylene (HDPE) for instance, has a density range of greater or equal to 0.941 g/cm³. With low degree of branching, HDPE has a stronger intermolecular forces and tensile strength. It is used in products and packaging such as containers, folding chairs, storage sheds and water pipes. Medium Density Polyethylene (MDPE) with a density of 0.926-0.940 g/cm³ is normally used in gas pipes and fittings, sacks, shrink film, packaging film, carrier bags and screw closures [11]. Due to their stiffness which makes them difficult to handle, HDPE and MDPE are not suitable for cable insulation.

Low Density Polyethylene (LDPE) is one of the favourite materials used for insulation. LDPE has a density range of 0.910 - 0.925 g/cm³ and is generally 50-60% crystalline [12]. Compared to HDPE, LDPE has more branching and thus, it has weaker intermolecular forces, lower tensile strength and higher ductility. Its excellent properties such as low dielectric loss, high dielectric strength, chemical inertness, low moisture up-take and ease of extrusion, make it easy to use in many kinds of telecom and power applications.

Table 2-1: General properties of LDPE [13].

<table>
<thead>
<tr>
<th>Properties</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molecular Weight</td>
<td>28.0 g/mol</td>
</tr>
<tr>
<td>Dielectric constant at 1MHz</td>
<td>2.2-2.35</td>
</tr>
<tr>
<td>Dielectric strength</td>
<td>27 kV mm⁻¹</td>
</tr>
<tr>
<td>Coefficient of thermal expansion</td>
<td>(100-200 )x10⁶ K⁻¹</td>
</tr>
<tr>
<td>Thermal conductivity at 23°C</td>
<td>0.33 W m⁻¹ K⁻¹</td>
</tr>
</tbody>
</table>

Table 2-1 shows some general properties of LDPE. The discrepancy between LDPE and Linear Low Density Polyethylene (LLDPE) is the absence of long chain branches in
LLDPE due to different manufacturing process in both LDPE and LLDPE. Although LLDPE has higher tensile strength and puncture resistance than LDPE, LLDPE is more difficult to process compared to LDPE. One disadvantage of LDPE is its low melting point which makes it unsuitable for high voltage cable insulation. However, with research developments, a polymer that has almost the same electrical properties as LDPE and can withstand higher temperature was produced.

2.2 The Crosslinking Polyethylene

Crosslinking process is a process of joining different polyethylene chain together by chemical reaction. The crosslinking of LDPE with Dicumyl Peroxide (DCP) to form XLPE was first accomplished by Gilbert and Precopio in 1955. XLPE can be considered to a branched PE with the end of the branches are connected to a different PE chain. Figure 2-4 shows the fundamental crosslinking process in polymer.

![Figure 2-4: Simplified description of crosslink network](image)

Crosslinking process retains the electrical properties of LDPE and at the same time improves certain properties of PE. Such properties are resistance to deformation and stress-cracking as well as improve tensile strength and modulus. Usually, antioxidant is added in the crosslinking process to prevent oxidative degradation to happen. If the oxidative degradation is significant, it may be harmful as it can lead to chemical changes within the insulation that could introduce more polar material which in turn, may change the electrical properties that prone to failure during ageing [4]. There are a few crosslinking mechanisms that are used in manufacture which are the high energy radiation-induced crosslinking, silane-induced crosslinking, and peroxide-induced crosslinking.
2.2.1 **High Energy Radiation-Induced Crosslinking**

A beam of electrons emanating from special equipment can interact with the polymer chains and form free radicals. These reactive radicals interact with other chains and hence induce crosslinking. In this radiation crosslinking process, the high energy electrons interact with the polymer chain and cleave the C-H or C-C bond. When C-H bond is broken, one hydrogen atom is released and \(~C\cdot\) radical tries to stabilize itself by combining with other radicals hence provides the crosslinking. The hydrogen atom combines with another hydrogen atom to form hydrogen gas. The radiation process is performed at room temperature which means polymer structure is in both crystalline and amorphous state [4]. Radiation-induced crosslinking is employed primarily for low voltage cable where the speed of crosslinking is a key issue. One weakness found in radiation-induced crosslinking is the non-uniformity of energy absorbed by thick specimen. The total energy absorbed is depending upon the electron beam energy. Therefore the degree of crosslinking by electron beam technology is not uniform within the component thickness and depends on the geometry. However, radiation-induced crosslinking do not produce any byproduct that stay in the polyethylene.

2.2.2 **Silane-Induced Crosslinking**

![Chemical reactions](image)

Figure 2-5: Silane-induced crosslinking process [14]. (1) The crosslinking reactions of silane-grafted polyethylene through the hydrolysis of silyl trimethoxy groups with water. (2) Condensation of the formed silanol groups.
Silane-induced crosslinking is a method that employs “moisture curing”. This type of crosslinking utilises the grafting of silane onto the PE and followed by the condensation reaction of silane graft by moisture [15]. The most common silane used in the manufacture is vinyl trimethoxysilane. The silane is commenced into PE by melt grafting using peroxide. Silane and peroxide are mixed together, compounded in an extruder at high temperature and immersed into water. Water induces chemical reaction leading to crosslinking [4,14]. The process is explained in Figure 2-5.

Silane-induced crosslinking has various advantages, such as easy processing, low cost and capital investment, and favourable properties in the processed materials. It also has less homocharge when stressed, lower conduction current, as well as has higher ac breakdown strength than peroxide crosslinked polyethylene [16]. However, this technique has been commonly employed for low voltage cables, as thicker cable insulation will take longer time for it to be crosslinked.

2.2.3 Peroxide-induced Crosslinking

The most common way in crosslinking polyethylene is through the use of peroxide agent. This organic peroxide agent is stable in room temperature but decompose at elevated temperature. For example, Dicumyl peroxide (DCP), which is well used peroxide for medium and high voltage cable [17], decomposed at 120°C [18]. The crosslinking agent is normally incorporated into PE pallets by the material supplier. After extrusion process (conversion of pallets into cable insulation), the PE is now surrounding the conductor and placed into a long curing tube where high temperature and pressure are applied. The PE is then melted and peroxide decomposes and induces the crosslinking process.

When DCP decomposes during curing process, the peroxide splits into active oxygen containing components and attract hydrogen atom from the polymer. As a result, it forms a free radical that is unstable. Polymer chain becomes active and unstable. Thus two such chains naturally combine to crosslink to stabilize the system and forms the XLPE [4]. During this process, several byproducts are formed from the peroxide and the major ones are methane, cumyl alcohol (dimethyl benzyl alcohol), acetophenone and α-methylstyrene.
When the free radical is generated, it will undergo 2 types of reaction to reach stabilisation. These ‘roots’ can be observed in the Figure 2-6. Firstly, the radical could rearrange itself and doffs a methyl radical to form acetophenone. The reactive methyl radical may also pick off a hydrogen atom from the polymer and forming methane gas. Secondly, the radical could also ‘grab’ the hydrogen atom from the PE chain and composes the relatively stable cumyl alcohol. Water may also form if the dicumyloxy radical expels a hydroxyl and hydrogen radical, to form water. As water is formed, the dicumyloxy radical converted to α-methylstyrene.
Cumyl Alcohol, acetophenone and methane are always found in greatest quantities in XLPE. Acetophenone is a liquid at ambient temperature due to its low melting point (approximately 20º C). It has somewhat sweet odour and not soluble in water and extent depending on the temperature. The methane gas must be allowed to readily protrude out of the freshly manufactured XLPE cable. α-methylstyrene could be found in smaller concentration. And sometimes, from further reaction of α-methylstyrene, another byproduct called cumene could be produced.

To ensure that the peroxide is uniformly dispersed which then means the cable is uniformly crosslinked, the temperature and pressure need to be properly controlled throughout the curing tube. This crosslinking process must happen in amorphous region, since the crystalline region cannot hold the peroxide preceding to the extrusion process. As the curing tube is heated, the whole polymer is amorphous and peroxide is considered uniformly dispersed. As the cable is cooled down, recrystallisation takes place and the byproducts are ‘forced’ into the new amorphous region.

2.2.4 Tree Retarded Crosslinked Polyethylene (TR-XLPE)

Numerous researches have been done to develop a better insulator. XLPE has been improved to be water treeing resistant. The earliest TR-XLPE cable made available in the early 1980s. A few approaches are proposed to achieve this property.

1. Replacing the polyethylene homo-polymer with a more polar copolymer.
2. Using an additive or placing additives into the homo-polymer (or co-polymer)
3. Using both of the options above

It was reported that there are high possibility for several additives present in the polymer. This additive serves as the agent to bind water molecules and provides the resistance to water tree. Acetophenone and dodecyl alcohol were found to facilitate resistance to treeing. Fillers may also trap water. Silane groups as we discussed before react with water. The overall aim here is to stop water from damaging the insulation by holding or trapping the water.
One issue appears in regarding TR-XLPE, on the definition of TR-XLPE or the degree of water tree retardancy. Although in this material, water tree growth has been removed along with the increment of AC breakdown strength compared to XLPE, one unclear issue is to what extend does the material has be improved in order to be considered ‘tree-retardant’ material. Table 2-2 compares the components that presence in XLPE and TR-XLPE

Table 2-2: Comparison between XLPE and TR-XLPE Insulated Cable

<table>
<thead>
<tr>
<th>Components</th>
<th>XLPE</th>
<th>TR-XLPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tree Retardant Additives</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Residual of DCP</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Crosslinking byproducts</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acetophenone</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Cumyl alcohol</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>α-methylstyrene</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Antioxidant + antioxidant byproducts</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

2.2.5 Antioxidant

During manufacturing process, it is usual for the insulation cable to undergo oxidative degradation. This process happens as the insulation temperature increases up to the level in which oxidation happens. If this happens, the molecular structure may change due to chemical reaction in the insulator hence introducing more polar molecules in the insulator. Antioxidant also changes the morphology of the polymer as it appears to have acted as a nucleating agent [19].

Antioxidant is incorporated into the polymer pallets to prevent the degradation mechanism process from happening. Antioxidant will naturally decompose in the extruder due to high thermal environment. At the beginning of curing process, the antioxidant will reside in the amorphous region. As the temperature increases, the crystalline melted and the antioxidant can migrate over the whole part of the molten polymer. During cooling process, the remaining un-reacted antioxidant as well as the non-volatile antioxidant degradation byproducts will inhabit in the amorphous region.
Many researches have been reported on the influence of antioxidant on electrical properties of XLPE. Goshowaki et. al in [20] reported that antioxidant containing sulphur has a great influence on the conductivity of XLPE. This measurement however was conducted on XLPE in which the conductivity value may be affected by the byproducts remaining in the XLPE. This hypothesis is in agreement with Sekii et. al in [21] who observed the formation of heterocharge in XLPE containing sulphur-base antioxidant. It is presumed that the heterocharge is created in the combine effect of the antioxidant and acetophenone. However reports in [22,23] stated that LDPE with antioxidant may generate deep traps or increase the deep trap density.

2.2.6 Peroxide Crosslinking Byproducts

Several properties of acetophenone, cumyl alcohol and α-methylstyrene are presented in Table 2-3.

Table 2-3: The physical properties of the chemical byproducts.

<table>
<thead>
<tr>
<th>Chemical Byproducts</th>
<th>Chemical Structure</th>
<th>Acetophenone</th>
<th>Cumyl Alcohol</th>
<th>α-methylstyrene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical Structure</td>
<td><img src="image" alt="Chemical Structure" /></td>
<td><img src="image" alt="Chemical Structure" /></td>
<td><img src="image" alt="Chemical Structure" /></td>
<td></td>
</tr>
<tr>
<td>%wt in cable [3]</td>
<td>0.6%</td>
<td>1.2%</td>
<td>0.02%</td>
<td></td>
</tr>
<tr>
<td>Melting point</td>
<td>19 - 20 °C</td>
<td>32-34 °C</td>
<td>-24°C</td>
<td></td>
</tr>
<tr>
<td>Boiling Point</td>
<td>198 - 204 °C</td>
<td>88-90°C</td>
<td>164-168 °C</td>
<td></td>
</tr>
<tr>
<td>Dielectric constant</td>
<td>17</td>
<td>9.7</td>
<td>3.8</td>
<td></td>
</tr>
<tr>
<td>Conductivity</td>
<td>$2.4 \times 10^{-4}$ S/m</td>
<td>$1.4 \times 10^{-7}$ S/m</td>
<td>$3.6 \times 10^{-9}$ S/m</td>
<td></td>
</tr>
<tr>
<td>Appearance</td>
<td>-clear to light yellow liquid -slightly soluble in water -Sweet odour</td>
<td>-Clear slightly yellow liquid -Insoluble in water</td>
<td>-Colourless clear liquid</td>
<td></td>
</tr>
</tbody>
</table>
A lot of researches have been done to investigate the properties of the byproducts in the performance of XLPE cable. There were various papers reporting on the contribution of crosslinking byproducts on the performance of XLPE and quite a number of them detailed the research on each of the byproducts. And these researches relate the decomposition of DCP with various electrical properties. For instance, Aida et al compared how each of the byproducts influence tan δ value [24]. At low electrical stress, tan δ value does not alter by adding the additives. However, this value is increased at high electrical stress region [25]. Limited sources have discussed on the effect of these byproducts on the breakdown. As in [3,26,27] the authors showed that short term AC breakdown strength of XLPE is improved with the presence of these chemicals. However, there are also some authors reported no significant effect of AC breakdown strength due to their existence in polymer [28].

Nevertheless, the effects of crosslinking byproducts on the accumulation of space charge were the most reported of all. Heterocharges were said to be observed in the bulk of the untreated XLPE [29-32]. The accumulation of heterocharges is explained by the mechanism of ion-pair separation from the byproducts [30]. However, the space charge accumulation was also associated with the production of water in the decomposition of cumyl alcohol [3,33]. The hole injection gets enhanced due to the increase of water content. This is considered to be the presence of water ions (H$_3$O$^+$) [34]. However, the main cause for heterocharges formation is the crosslinking byproducts in the XLPE.

T. Ohara et. al in [35] that focuses on the acetonophenone suggested that this byproduct increases the conductivity of the soaked LDPE. By using the coated sample, they proved that holes are moving towards the uncoated side. Acetonophenone changes the $\sigma$/$\varepsilon$ value which causing charge accumulation at the interface of soaked and un-soaked sample. Same result was observed by Y. Ohki et. al, and from calculation, acetonophenone soaked LDPE was said to have higher conductivity than $\alpha$-methylstyrene soaked LDPE which are $10^{13}$ and $10^6$ times of un-soaked sample, respectively [5]. And due to the difference in ($\sigma$/$\varepsilon$) at the interface of soaked and un-soaked sample, charges are generated at the interface. In the same paper, the authors concluded that positive charge migration in the LDPE soaked in $\alpha$-methylstyrene is faster than the one soaked in acetonophenone due to the permeation velocity in LDPE.
In most of the reports, acetophenone and α-methylstyrene give no effect on the space charge formation. It was said that the chemicals accelerate the charges transportation in LDPE thus leave no charge in LDPE [7,36,37]. It was also said that acetophenone reduces the injection barriers and enhance the charge injection [38]. However, heterocharges were observed when water molecule and acetophenone present in LDPE. Hirai et. al in [36] reported that the conduction current for acetophenone or α-methylstyrene decreases with time, unlike the conduction current for cumyl alcohol that initially increases, and then decreases with time. However, despite of its high conduction current, cumyl alcohol was observed to cause deep traps in soaked LDPE sample. This observation is also reported in other papers. The formation of homocharge can be observed as deep traps from cumyl alcohol trap the charges injected from the electrodes [39].

2.3 Chapter Summary

In this chapter the fundamental theory of polymer particularly PE is presented. PE is graded based on its’ density with each density of PE is useful for different usage. Later, when crosslinking process is introduced, XLPE has become an alternative to the oil impregnated paper for cable insulation for higher voltage cable. Table 2-4 presents the comparison between the types of crosslinking available.

Table 2-4: Comparison of the crosslinking technique.

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Crosslinking type</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>High Energy Radiation</td>
</tr>
<tr>
<td>Operating temperature</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Byproducts</td>
<td>-</td>
</tr>
<tr>
<td>Advantages</td>
<td>-No byproduct</td>
</tr>
<tr>
<td></td>
<td>Acetophenone, methane, α-methylstyrene, cumyl alcohol, cumene</td>
</tr>
<tr>
<td></td>
<td>-uniform crosslinked</td>
</tr>
</tbody>
</table>
Peroxide induced crosslinking has been the most popular crosslinking technique despite the formation of byproducts in the crosslinked structure. Peroxide induced crosslinking could offer a uniform crosslinking process in thicker sample which suitable for high voltage operation. In chapter 6, this type of crosslinking will be performed to produce XLPE sample for space charge measurement.
Chapter 3  Space Charge in Polymer

3.1 Introduction

From the context of dielectric material, the term ‘space charge’ corresponds to the carriers (electrons, holes and ions) that move around in the dielectric material by the applied field and become trapped in the material. Low charge carrier mobility and charge trapping within the polymer give rise to the space charge, leading to localised electric stress enhancement. The formation of space charge may distort the electrical field and distribution throughout the cable insulation thickness which may lead to the premature failure of the cable insulation at stresses below the anticipated value [30,32]. This condition becomes more severe during dc application. The strong correlation between the space charge and breakdown in cable insulation is reported in [40] where the evidence linking space charge build up, tree growth and breakdown in XLPE is presented. The presence of space charge is also linked to aging acceleration mediated by lowering of degradation process activation energy [41,42].

There are a few mechanisms of space charge formation in the polymer. Basically, space charge may result from;

i) Charge injection and extraction from electrodes;

ii) Electric field assisted ionisation of impurities in the polymeric material or from an inhomogeneous polarization;

iii) Application of a mechanical/thermal stress.
For the purpose of this research, we will only focus on the effect of impurities in the polymer as the other variables such as type of electrodes, and mechanical/thermal stress will remain constant. Homocharges and heterocharges are terms that are often used to describe the polarities of the space charges. Homocharges refer to the charges with the same polarity to the adjacent electrode while heterocharges refer to the opposite polarity. When voltage is applied across the polymer, small molecules may become ionised and drift towards the electrode with opposite polarity which eventually produce heterocharges. This type of space charges will increase the stress at the electrode interface but reduce the electric stress in the bulk. Meanwhile, homocharges are usually formed from the charge injection or charge extraction near the electrode. Homocharges reduce the stress in the vicinity of the electrode interface but enhance the bulk stress [43].

3.2 Trapping and Detrapping

As voltage is applied across the polymer, all carriers experience the same force but they move with different velocities. These are what we called as fast and slow moving charges. The charge movement depends on the ‘roots’ at which it takes towards the electrode.

This phenomenon is possible with the existence of traps in the dielectric bulk. Localised states that results from assorted physical attributes such as broken bonds within the crystalline structure, chain branches, polarised states, dislocations foreign particles, crosslinking impurities and additives will act as traps to the charge carriers at energy states below the conduction band of the polymer. These traps for electrons are known as the acceptor and the traps for holes are known as donors. However, there are also traps that act as the recombination centres by trapping both electrons and holes [44,45].

The best way to describe traps is by the energy level diagram of polymer, which consists of a conductive band which is separated from the valence band by a huge forbidden gap (more than 7eV) [46]. It can be pictured as a potential well that exists in the energy diagram where it effectively acts as a potential barrier that obstructs the carrier movement in the conduction or valance band [47]. Generally, traps are divided into two types; deep
traps and shallow traps, depending on their depth. The former energetically located 0.8 to 1.4 eV below the conduction band and the latter is at 0.1 to 0.3 eV below the conduction band. Trapped charges normally reside longer in the deep traps and inevitably introduce space charge. Under the influence of the electric field the trapped charges may begin to detrap and move. Figure 3-1 shows the representation of the traps in electron energy diagram.

![Energy Band Diagram](image)

**Figure 3-1: Schematic representation of shallow and deep traps in energy band diagram.**

Basically, the slow moving charges are the charges which are trapped into the deep traps, while the fast moving charges are the free moving charges or charges that are trapped in the shallow traps.

When discussing about the charge transportation in insulating polymer, there are two parts need to be considered; the charge injection from the electrode and the high field conduction process in the bulk.
3.3 Electrode Process

3.3.1 Schottky Injection

Injection due to Schottky theory is based on the potential barrier height of the metal-insulator interface. This value depends on the type of interface. In the case of metal-vacuum boundary, barrier height equal to the work function of the metal. However for metal-dielectric interface, the barrier height depends on the metal-dielectric work function difference as well as the local condition of the electric polarisation. [43].

In the simple band structure, the barrier between metal and insulation is assumed to be precipitous as shown in Figure 3-3 (a). However in Schottky injection theory, the barrier height is reduced by the electrostatic attraction between the electron and metal. In this case, the metal is positively charge as it loss an electron to the dielectric. As a result, the barrier height is constantly changed due to the potential energy of the electron.
Figure 3-3: Modification of the potential barrier between metal and polymer by an applied field. (a) total barrier height, (b) shape of barrier including effect of coulombic image force, (c) potential energy due to the applied field, and (d) total barrier shape and (e) illustration of the classic law of image.

To calculate the attraction between the metal and electron, consider the electrode to be replaced by an equal and opposite polarity charged point at equal distance from behind the
electrode surface. This is known as image charge theorem which is illustrated in Figure 3-3(e). If electron is in x-distance from the interface, then the distance separating electron and image is 2x, and the force of attraction due to Coulomb’s Law is:

\[
F_i = \frac{e^2}{4\pi\varepsilon_0 \varepsilon_r (2x)^2} = \frac{e^2}{16\pi\varepsilon_0 \varepsilon_r x^2}
\]

Equation 3-1

Also the potential barrier due to the image field (Figure 3-3(b)) is given by:

\[
\Phi_i = \int_x^\infty F_i(x) dx = \frac{e^2}{16\pi\varepsilon_0 \varepsilon_r x}
\]

Equation 3-2

Potential due to applied field, \(\Phi_a\) is represented in a straight declining line illustrated in Figure 3-3(c) given by

\[
\Phi_a = -e(Ex)
\]

Equation 3-3

Hence, the total resulting potential, given by \(\Phi(x) = \Phi_i + \Phi_a\) is

\[
\Phi(x) = \frac{e^2}{16\pi\varepsilon_0 \varepsilon_r x} - e(Ex)
\]

Equation 3-4

The maximum potential is given by

\[
\frac{d\Phi}{dx} = -\frac{e^2}{16\pi\varepsilon_0 \varepsilon_r x} - eE = 0 \quad , \quad x = x_m
\]

Equation 3-5

Considering magnitude of electric field \(E = |E|\), we could obtain maximum stationary point \(x_m\)

\[
x_m = \sqrt{\frac{e}{16\pi\varepsilon_0 \varepsilon_r E}}
\]

Equation 3-6

And the barrier height is now reduced by \(\Phi_m\) (Figure 3-3(d)) ;
\[ \Phi_m = -eV = \frac{e^2E}{\sqrt{4\pi\varepsilon_0\varepsilon_r}} \]  
Equation 3-7

So that the effective barrier height is

\[ \Phi_{eff} = \Phi - \frac{eE}{\sqrt{4\pi\varepsilon_0\varepsilon_r}} \]  
Equation 3-8

Employing Richardson-Dushman equation, \( J_T = AT^2 \exp\left(-\frac{\Phi}{kT}\right) \), to Schottky injection effect:

\[ J(E) = AT^2 \exp\left(-\frac{\Phi_{eff}}{kT}\right) \]  
Equation 3-9

\[ J(E) = AT^2 \exp\left(-\frac{\Phi}{kT}\right) \exp\left(\frac{1}{kT} \sqrt{\frac{e^2E}{4\pi\varepsilon_0\varepsilon_r}}\right) \]  
Equation 3-10

where \( A \) is the Richardson-Dushman constant and \( T \) is the absolute temperature. Giving the current density:

\[ J(E) = J_T \exp\left(\frac{1}{2kT} \sqrt{\frac{e^2E}{\pi\varepsilon_0\varepsilon_r}}\right) \]  
Equation 3-11

3.3.2 **Fowler-Nordheim Injection**

Fowler-Nordheim effect is also known as the tunnelling effect from bulk metal to other bulk crystalline solids. It is found that electron may tunnel through a potential barrier although it has less energy to overcome the barrier’s height. This is due to the particle-wave duality characteristic of electron as shown in Figure 3-4(a). As the electron wave travel through the barrier, the wave will attenuate reducing its’ amplitude but leaving the energy unchanged.

Figure 3-4 (b) shows the schematic diagram of tunnelling injection through a barrier. If very high field is applied, the height and width of the potential barrier will be reduced.
This tunnelling process is width dependent and in fact, in this charge injection theory, the width of the potential barrier is much more important than the barrier height. The probability of tunnelling decreases for taller and wider barriers. This emission process will occur at which the potential barrier is the thinnest, marked as $x_0$ in Figure 3-4 (b).

![Diagram of Fowler-Norheim tunnelling injection](image)

Figure 3-4: Illustration of Fowler Norheim tunnelling injection; a) electron wave travel through the barrier, and b) potential barrier reduction due to applied field.

The conduction current from Fowler-Norheim injection can be described as;

$$J = \frac{e^3 E^2}{8\pi h \Phi_{\text{eff}}} \exp\left(-\frac{8\pi \sqrt{2m}}{3e h E} \Phi_{\text{eff}}^2\right)$$

Equation 3-12
Here $e$ is the electron charge, $h$ is the Plank constant, $m$ is the mass, $\Phi_{\text{eff}}$ as the effective potential barrier height and $E$ the electric field. The hole transport, which only happen via electron vacancies, requires inter-chain hole transfer, a reverse resonant tunnelling process of electron [48] and therefore, hole transfer normally happen in crystal region. The tunnelling process only happens at very high field where the barrier width is very thin.

3.4 Bulk Process

3.4.1 Space-Charge Limited Conduction

Space-charge limited current is found to be incredibly dependent on the thickness in which the current is the result from the movement of charges that injected into the polymer. Polymer with high dielectric constant will lead to high charge built up in the material.

(a) Trap-free SCLC

In this model, an ideal dielectric is considered, which has [43];

- No trap
- No thermally generated charge, all charges come from injected charge
- Good injection due to Ohmic contact
- Negative-charged carrier only (i.e. electrons)

The dielectric with $d$ thickness is placed in between two parallel electrodes. The $x$-direction is perpendicular to the electrodes, straight through the dielectric. The conduction current which have three components; drift, diffusion and displacement current is given by:

$$J = ne\mu E - eD_n \frac{dn}{dx} + \varepsilon_0 \varepsilon_r \frac{dE}{dt}$$

Equation 3-13

where $D_n$ is the Fick’s diffusion coefficient. Two assumptions are made here which are; steady state condition, $dE/dt=0$, and space charge concentration is constant throughout the dielectric (although in real life space charge build up near electrodes will experience the greatest diffusion). In the latter, the diffusion term will be neglected. From Poisson equation;
\[
\frac{dE}{dx} = \frac{ne}{\varepsilon_0 \varepsilon_r} \quad \text{Equation 3-14}
\]

So that:

\[
J \approx \varepsilon_0 \varepsilon_r \mu E \frac{dE}{dx} \quad \text{Equation 3-15}
\]

By rearranging this equation and integrating both sides gives;

\[
E = \sqrt{\frac{2J}{\varepsilon_0 \varepsilon_r \mu}} (x + x_0) \quad \text{Equation 3-16}
\]

where \(x_0\) is a constant of integration. Integrate once again from \(x=0\) to \(x=d\) with \(V(0)=V\) and \(V(d) = 0\), it gives

\[
V = \int_0^d E \, dx
\]

\[
= \int_0^d \sqrt{\frac{2J}{\varepsilon_0 \varepsilon_r \mu}} (x + x_0) \, dx
\]

\[
= \frac{2}{3} \sqrt{\frac{2J}{\varepsilon_0 \varepsilon_r \mu}} \left( \left(d + x_0 \right)^{3} - x_0^{3} \right) \quad \text{Equation 3-17}
\]

Assuming \(x_0 \ll d\) so that the current density is given by:

\[
J = \frac{9\varepsilon_0 \varepsilon_r \mu V^2}{8d^3} \quad \text{Equation 3-18}
\]

This is also known as Mott & Gurney square law in which the current is proportional to square of the voltage. By rearranging equation 3-18, mobility could be obtained from;

\[
\mu = \frac{8jd^3}{9V^2 \varepsilon_0 \varepsilon_r} \quad \text{Equation 3-19}
\]
(b) **Trap-limited SCLC**

Unlike the trap-free dielectric with all carriers in conduction band, dielectric with traps will have some carriers being trapped in the material. Although the injected charges contribute to the space charge, but only a portion of it contribute to the current. Hence, a fraction of \( \theta = \frac{n_c}{n_t} \), where \( n_c \) and \( n_t \) attribute to the number density of conduction band electrons and that of occupied trap states respectively, is necessary to be added into the current equation:

\[
J = \theta \frac{9 \varepsilon_0 \varepsilon_r \mu V^2}{8 s^3}
\]

Equation 3-20

Figure 3-5: Log-log plot of current-voltage characteristics for one carrier, SCLC injection into an insulator.

SCLC current-voltage characteristic is commonly illustrated in log-log plot due to their power law relationship. This characteristic is demonstrated in figure 3-5. At low voltage, SCLC effect is not noticeable thus Ohmic conduction dominates the current-voltage
characteristic. As the voltage becomes higher, the SCLC effect starts to appear as it entering the shallow trapping region. As the voltage increases further, more electrons injected and ultimately fill in all the traps giving the current rises vertically at $V_{TFL}$ (Trap-Filled- Limit). This steep increase the current-voltage characteristic will follow the trap-free SCLC relationship having similar slope with that in the trapping region. The concentration of the trapping state could be found by knowing $V_{TFL}$ [49].

3.4.2 Hopping Conduction

![Diagram of hopping conduction mechanism](image)

Figure 3-6: Hopping conduction mechanism; before and after electric field application.

When no electric field applied to the polymer, electron may move in any direction. The probability of this electron to move in any direction is $1/6$ given by the positive and negative direction of three coordinates that perpendicular to each other [50]. With electric stress application, the barrier height is reduced and this reduction will help promote charge tunnelling effect. With higher electric fields, the potential barrier in the forward direction will continue to reduce and make conduction easier.

During the application of electric field, the barrier height will be changed. The new barriers’ height along or against the direction of electric field become $W + \frac{1}{2} \delta E e$ and $W - \frac{1}{2} \delta E e$ respectively. Thus, the probability of electron hopping in these two directions is;
where \( v \) is the escape frequency of electrons.

### 3.4.3 Poole-Frenkel Effect

Poole and Frankel mechanism is the bulk limited version of the Schottky effect at electrode interface. In contrast to the Schottky effect, Poole and Frankel mechanism reduce the trap barrier localising the carries in the dielectric itself. For this mechanism to occur, the dielectric must have a wide band gap with donor or acceptor resides in it. With the presence of electric field across the dielectric material, reduction on the potential barrier will occur and the schematic diagram of this effect can be seen in Figure 3-7.

![Figure 3-7: Schematic diagram of the Poole-Frenkel effect.](image)

The potential energy, \( V(r) \) related to the Coulombic interaction between the electron and ionised donor is given by:

\[
V(r) = -\frac{e}{4\varepsilon_0\varepsilon_r r}
\]

Equation 3-23
where \( r \) is the distance of separation. With the application of electric field of \( E \), the potential energy becomes:

\[
V(r) = -\frac{e}{4\varepsilon_0\varepsilon_r r} - Er
\]

Equation 3-24

Applying the procedure in 3.3.1, the position of the maximum potential surface, \( r_m \) is given by

\[
r_m = \sqrt[4]{\frac{e}{4\pi\varepsilon_0\varepsilon_r E}}
\]

Equation 3-25

And thus giving \( \Phi_m \):

\[
\Phi_m = \sqrt[4]{\frac{e^3E}{\pi\varepsilon_0\varepsilon_r}}
\]

Equation 3-26

By reducing the new potential barrier into the equation of Schottky injection, the current density obtained is:

\[
J(E) = J_T \exp \left( \frac{1}{kT} \sqrt[4]{\frac{e^3E}{\pi\varepsilon_0\varepsilon_r}} \right)
\]

Equation 3-27

Equation 3-27 could be rewrite as:

\[
\ln J(E) = \frac{1}{kT} \sqrt[4]{\frac{e^3}{\pi\varepsilon_0\varepsilon_r}} \sqrt{E} + \ln J_T
\]

Equation 3-28

Which clearly illustrate a straight line with slope \( \frac{1}{kT} \sqrt[4]{\frac{e^3}{\pi\varepsilon_0\varepsilon_r}} \) when \( \ln J(E) \) is plotted against \( \sqrt{E} \).
3.5 Charge Transfer and Polymer Morphology

Polymer chains in PE always arrange itself to form crystalline lamella ribbons. The lamellas normally grow from a nucleating site to form spherulites which may have diameter of more than 10µm. This array of lamella crystal is separated by less dense amorphous polymer. Figure 3-8 illustrates the spherulites in PE. J.P. Jones et al. in his work [48], mentioned about the effects of polymer morphology to charge transfer.

Figure 3-8: Morphology of semi-crystalline PE showing spherulites array of lamella crystallite.[48]

Figure 3-9: Change of energy level in electron transfer due to the polymer morphology. [48]
Electron for example, will move in inter-lamellar-phase particularly in chains that align with the applied electric field, to avoid PE chains in which has higher energy barrier. Localised polaronic energy state, such as at T in Figure 3-9 is found due to the neighbouring electrons, acts as the restricting barrier to electron transport. Therefore, electron transport depends on;

1. Local polarisation
2. Chain conformation

In the same figure, as electron travels from region 1 to 3, higher energy required to overcome the barrier at region 2. The closer the electron path to the polymer chain, the higher energy is required to attain them. Thus, this path is less preferred. The numerous hills of electron affinity are due to the chain configuration in PE and region with lowered density tends to become macro (deep) traps. Hence, the polymer creates an energy landscape in which the electron travels.

Figure 3-10: Positive hole transfer between two chain via valance band of polymer chain and inter-chain tunnelling which is difficult at a-a and easy at b-b. [48]

Holes on the other hand, only travel via electron vacancies in polymer chains. For a hole to transfer in a long range, it will depend on two transfer mechanisms;

1. Inter-chain hole transfer, and
2. Reverse resonant tunnelling process of electron between chains that are close to each other.

Thus, in amorphous region, holes transfer becomes more difficult due to the morphology of the region which encouraging the hole trapping.

3.6 Chapter Summary

This chapter is important as later in this thesis we will discuss on the charge build up in the tested samples and the influence of crosslinking byproducts on charge injection at electrodes, charge movement in the bulk as well as changing the trapping landscape of the dielectric. Thus it is important to grab the idea of deep and shallow traps from the beginning so that the result obtained from the tests could easily be analysed. The effect of hopping will also be seen in the tested sample during the charge decay process in space charge measurement. No voltage is applied across the sample and charge in the tested sample decay gradually depending on the type of sample.

The theories on conduction process that presented in this chapter will also be useful as we discuss the effect of byproducts on dc conductivity. Some of the conduction models will be referred to calculate the mobility of the charges in samples. Last but not least, the effect of dielectric morphology on charge conduction demonstrates the importance of local condition as well as the polarisation effect of the neighbouring molecules towards the trapping properties of the dielectric.
Chapter 4  Space Charge Measurement Techniques

Since the term space charge is introduced, there have been a few devices developed to enable the space charge in polymer to be determined. Three main non-destructive techniques have been adopted which are the thermal step or pulse method, laser-induced pressure pulse (LIPP) method and also last but not least the pulse electroacoustic (PEA) technique [51]. These methods use space charge displacement technique in order to measure the charges that exist in the specimen. The difference among them is the technique used to introduce the small displacement to the charges and this matter will be discussed in here.

4.1 Thermal Step Method

Collins, in mid-1970s, was the first person who introduced diffusion of heat pulses in the sample to measure space charge. Later, in 1988 the thermal step method (TSM) was proposed by Toureille which measures the external current in response to the thermal expansion in the sample [52].
Figure 4-1: Diagram of thermal step principle.

To have a better understanding on the principle of the thermal step method, let’s consider a dielectric slab with a thickness \( L \), that contains \( Q \) charges at a distance \( x \). Before the experiment, the sample is at low temperature. The sample is placed in between two electrodes and at \( t=0 \), the hot source is placed at one electrode at position \( x=0 \), meanwhile the cold sink is at \( x=L \). The thermal expansion can be created by a flash light or illumination of a laser. During short circuit, the electrodes are connected to the ground resulting to no potential difference. When the thermal step is introduced to the system, the material layers shifted resulting to the charges \( Q \) in the sample to be displaced. Consequently, the induced image charges \( Q_1 \) and \( Q_2 \) at the electrodes are also modified to fulfil the electrostatic equilibrium:

\[
Q_i + Q_1 + Q_2 = 0 \quad \text{Equation 4-1}
\]

As a result, small current is produced in the external circuit and it is given by [53,54] :

\[
I(t) = -AC \int_0^L E(x) \frac{\partial \Delta T(x,t)}{\partial t} \, dx \quad \text{Equation 4-2}
\]

42
where \( A \) is the parameter related to the variation in thickness and permittivity with temperature, \( C \) is the material capacitance and \(-AC= \delta C/\delta T\) the variation of \( C \) with temperature and \( E(x) \) is the local electric field in the sample.

Application of a suitable deconvolution procedure using Fourier analysis allows extraction of the space charge distribution from the current flow data. However, this technique is particularly suitable for thicker samples, between 2 and 20mm since there is essentially an unlimited amount of heat available and useful data can be collected until the temperature distribution across the sample has equilibrated. Due to this low resolution, some other new methods have been introduced so that the space charge in thinner sample could be measured.

4.2 Laser-Induced Pressure Pulse

![Laser Induced Pressure Pulse System](image)

Laser Induced Pressure Pulse (LIPP) initiates the charge displacement from the pressure wave that is produced when the laser beam hits the target. LIPP system is illustrated in figure 4-2. In this case, the target used is a semiconductor electrode which has the same acoustic impedance with the dielectric material. As the pressure wave travels through the sample thickness, it induces an external signal (current or voltage) which tells the electric field profile of the sample. In the case of external current, the induced signal is given by \([55,56]\);
where $A$ is a constant that depends on the sample properties (permittivity, compressibility and electrical capacity), $d$ is its thickness, $E(x)$ the local field resulting from both the applied voltage and from space charge, and $P(x, t)$ the pressure pulse. By deconvolution technique, the charge distribution of the sample could be obtained. Compared to TSM, LIPP allows any space charge measurement to be conducted on thin samples by controlling the laser intensity to obtain a high signal-to-noise ratio in order to have a good data. However, there are a few technical problems encountered when applying this method [55].

During the formation of the pressure pulse, the optical energy is converted into mechanical energy. Consequently, some of the high optical energy may be converted into heat which increases the local temperature and change the target surface into gas. This gas could be ionised and forms ‘plasma’ which radiates electromagnetic energy that can disturb the signal and increase the noise in the system. The formation of plasma can be observed in Figure 4-3. In addition, due to the heating effect, thermal step diffusion may occur in the measured sample. If this happens, the induced signal may result from both thermal and mechanical stresses and the signal cannot be treated.

The impact of laser beam to the target is more severe when it burns away the target electrode with each laser pulse. With the damaged target surface, the induced signal becomes out of phase. From equation 4-3, the current produced is dependent to the internal field distribution $E(x)$ and its pressure shape $P(x, t)$. The pressure shape is related to the laser beam energy which normally have 5% variation (for commercial laser beam). This non-uniform laser beam energy may lead to variation of $i(t)$ signal which consequently affects the charge profile. Due to these limitations, the use of piezoelectric is more preferable compared to a laser in generating the pressure pulse. This brings us to the final method, which uses a better technique to measure space charge.
4.3 Pulsed Electroacoustic Technique

Due to simplicity, low cost, non-destructive and ease of implementation, PEA method is now more commonly used, rather than the other two. Figure 4-4 shows the experimental setup of the pulse electroacoustic (PEA) technique. It consists of two aluminium electrodes...
with semiconductor attached to the top electrode to match with the acoustic impedance of the dielectric sample. PEA introduces the charge displacement by applying short voltage pulse $e_p(t)$ with duration $\Delta T$ across the sample. Any charges $\rho(x)$ in the sample will experience perturbation force that causes the charge displacement. This displacement launches an acoustic wave which travels through the sample and bottom electrode, towards the transducer. The generated acoustic wave has amplitude proportional to the local charge density in the sample. Piezoelectric transducer will then transform the acoustic wave into electric signal.

Piezoelectric transducer which is made of Polyvinylidene Difluoride (PVDF) is connected to oscilloscope via amplifier to enlarge the voltage signal. The oscilloscope will then record the signal as a function of time [51,57,58]. The spatial resolution for acoustic technique depends on the product of the pressure pulse duration and the speed of sound in the sample. In the PEA method, pressure pulse duration is equivalent to the voltage pulse duration. However, the thickness of the piezoelectric transducer must also be considered since the practical resolution improves as the transducer thickness decreases. The thickness of the device is normally 4 to 110µm. An absorber is attached to the transducer to delay and restrain the reflection of the acoustic wave from going back to the transducer. Other than acting as the bottom electrode, aluminium also plays an important role of delaying the arrival of the acoustic pulse until the instability due to the pulse source disappears before allowing the acoustic wave to reach the transducer. A full description on the acoustic method in measuring electric charge distribution is reported in [59].
Figure 4-5 illustrates the schematic diagram of the PEA system. As shown in the figure, $\sigma_1$ and $\sigma_2$ appears at the electrode, as the surface image charges. These values are dependent on the electric field $E_{dc}$ as well as the magnitude and distribution of the total charge density $\rho(x)$. The magnitude of $\sigma_1$ (at $x=0$) and $\sigma_2$ (at $x=d$) are described by equations 4-4 and 4-5 respectively [60].

Equation 4-4

$$\sigma_1 = -\varepsilon E_{dc} \int_0^d \frac{d-x}{d} \rho(x) \, dx$$

Equation 4-5

$$\sigma_2 = \varepsilon E_{dc} \int_0^d \frac{x}{d} \rho(x) \, dx$$

where $\varepsilon = \varepsilon_0 \varepsilon_r$ and $d$ is the sample thickness. The electric field will generate surface charges equal in magnitude but opposite sign, meanwhile the total charge in the bulk will generate surface charges in different magnitude but the same sign. Equation 4-6 shows the corresponding voltage proportional to the space charge density $\rho(x)$ [60]:

$$V_s(t) = G_0 \left[ K_1 \sigma_1 \delta \left( t - \frac{B}{v_{Al}} \right) + K_2 K_3 \sigma_2 \delta \left( t - \frac{B}{v_{Al}} - \frac{d}{v_{sa}} \right) \right] e_p \Delta T + 0.5 K_3 v_{sa} \rho(x = v_{sa} t)$$

Equation 4-6
Vs is the electric signal obtained in the time domain which represents the charge distribution, where $\sigma_1$ and $\sigma_2$ are the surface charges at the electrodes at time $t$. $B$ and $d$ are the electrode and sample thickness respectively. $v_{Al}$ and $v_{sa}$ are the sound velocity through the aluminium electrode and sample thickness respectively. $\Delta T$ is the width of the pulse, $\rho$ is the bulk charge and $e_p$ is the amplitude of the pulse voltage. $G_0$ is a constant meanwhile $K_1$, $K_2$ and $K_3$ are the transmission coefficients associated to the impedance of the materials, and are given by equation 4-7 to 4-9;

$$K_1 = \frac{Z_{Al}}{Z_{Al}+Z_{sa}} \quad \text{Equation 4-7}$$

$$K_2 = \frac{Z_{sa}}{Z_{Al}+Z_{sa}} \quad \text{Equation 4-8}$$

$$K_3 = \frac{2Z_{Al}}{Z_{Al}+Z_{sa}} \quad \text{Equation 4-9}$$

where $Z_{Al}$ and $Z_{sa}$ are the acoustic impedances of the aluminium electrode and sample respectively. In equation 4-6, the constant $G_0$ needs to be determined and this is typically done by applying a small voltage across the sample to generate a known charge density on the two surfaces at the electrodes. If there is no bulk charge in the material, the third term in equation 4-6 will become zero and thus,

$$V_{\sigma_1}(t) = G_0 K_1 \sigma_1 e_p \delta \left( t - \frac{B}{v_{Al}} \right) \Delta T \quad \text{Equation 4-10}$$

$$V_{\sigma_2}(t) = G_0 K_2 K_3 \sigma_2 e_p \delta \left( t - \frac{B}{v_{Al}} - \frac{d}{v_{sa}} \right) \Delta T \quad \text{Equation 4-11}$$

Where $K_1=0.5 \ K_3$ (from equation 4-7 and 4-9) and $V_{\sigma_1}(t)$ and $V_{\sigma_2}(t)$ are the output voltage associated to the surface charges $\sigma_1$ and $\sigma_2$. In practice, Equation 4-10 is used since the bottom electrode is nearer to the transducer and thus less loss due to attenuation and dispersion of the signal. The propagated signal will gradually decrease in amplitude and broaden its shape due to scattering, absorption and dissipation of energy [61]. By integrating equation 4-10 within $\Delta T$ region, as well as applying equation $\sigma_1 = -\varepsilon E_{dc}$, we obtain [60];
Substituting equation 4-13 to the third term of equation 4-6, the space charge density profile \( \rho(x) \) which is proportional to the output voltage \( V_s(t) \) is obtained;

\[
V_s(t) = (G_0 K_1 v_{sa} \Delta T) \rho(x = v_{sa} t) e_p
\]

Equation 4-14

In the PEA software, \( V_s \) value is calculated from electric field and charge density of the sample. This value will match the applied voltage across the tested sample from the volts on measurement signal. In some cases, the output signal can sometimes be misinterpreted as space charge, where it actually is a result from the frequency response of the transducer and amplifier. The transducer with a high capacitance (typically 0.1-5nF) as well as low noise and high frequency amplifiers (500 MHz) which often have low input impedance (50Ω), acts as a high pass-filter. If \( G(f) \) is the transfer function of the system, the output signal in frequency domain, \( V(f) \) can be described as;

\[
V(f) = G(f) \rho(f)
\]

Equation 4-15

where \( \rho(f) \) is the impulse response of the system, including the sensor and amplifier in the frequency domain. If \( G(f) \) can be found, the actual space charge distribution can be computed using Fast Fourier Transform (FFT), division, software filtering and Inverse Fast Fourier Transform (IFFT). The full mathematical principle behind deconvolution is well documented in [62].

4.3.1 Calibration of Pulsed Electroacoustic Signal

Ideally in a clean sample, no bulk charge can be found. However, due to the system respond and noise subsequent to the applied pulse voltage, the signal from PEA at bulk of sample is not always zero. The surface charge \( \sigma_1 \) and \( \sigma_2 \) are produced by the applied pulse voltage and the actual pulse voltage on sample is related to the capacitance ratio of
decoupling capacitor and sample. Therefore, the calibration of the PEA signal is a must to eliminate the effect.

By applying a dc voltage in the opposite direction to the pulse voltage, these induced charge at the top and bottom electrodes cancel each other. Due to the equivalent applied voltage to the output signal from the pulse voltage, it is possible to cancel the effect of the latter. The relationship between the pulse voltage and its equivalent dc voltage can be used to improve the charge analysis. For example, 600V pulse voltage can be used to calibrate sample stressed at 2kV. Due to the noise from surrounding and from pulse generator, signal from pulse voltage only could have low signal-to-noise ratio (SNR). Thus, calibration process requires additional voltage to perform a better calibration. It is also important to confirm that no bulk charge is produced by the applied dc voltage. From previous studies [63,64], in a 200μm LDPE sample, homocharge is initiated when a 3kV dc is applied. At the same time, the applied voltage must be high enough to yield a signal of acceptable SNR.

Hence, a suitable signal of calibration is the signal measured at 2kV. This value however will vary with the thickness. A higher voltage could be used for a thicker sample. For sample with a trapped charge, the same batch of samples without charge is used for calibration purpose. The measurement is carried out on the sample without involving sample loading and unloading in the PEA system to avoid the slightest difference in the sample assembling which will result in a variation in the output signal [63]. It should also be noted here that calibration process must be conducted on every sample before any further measurement is done on the sample.

In Tony Davies High Voltage Laboratory in University Of Southampton, this technique has been used for many years due to its advantages as stated above. PEA measurement for cable insulation is also available to measure the space charge accumulation in plaque or coaxial-cables. Improvement has been made on the conventional cable PEA system by Fu and Chen [65], to allow measurement on the cable at the same radius. A flat electrode system was introduced and employed in the system and the measuring assembly is illustrated in Figure 4-6 below:
Figure 4-6: Illustration of a modified pulsed electro-acoustic (PEA) system for cable [65].

By using this equipment, the space charge profile in XLPE cable with temperature gradient was successfully measured [66].

4.3.2 Pulsed Electroacoustic Technique In AC Condition

The accumulation of space charge under ac condition has not been fully investigated due to a few factors. Firstly it may due to less interest on this study because of smaller space charge accumulation in ac condition making it less problematic than that at dc. Secondly, ac space charge measurement needs to have faster data acquisition and good synchronisation with the ac supply to measure the phase related charge [67]. Previously, authors like Wang et. al and Chong et. al have used the point on wave technique to measure the ac space charge profile [64,68]. This method is shown in Figures 4-7 and 4-8.
Figure 4-7: Illustration of point on wave method[69]

Figure 4-8: Demonstration of pulse train and applied voltage in point on wave method [69]

Due to low switching frequency of the pulse generator as well as the slow data transferring between the oscilloscope and GPIB card, only one set of data could be recorded per cycle. ‘m’ in figure 4-8 is defined as the number of point to be measured in one cycle meanwhile ‘N’ is the average number of pulses required on each point to obtain a good signal to noise ratio. The number of m is limited by the frequency of the pulse generator such that $f_{\text{pulse}} < m f_{\text{ac}}$. If a sine waveform of 50Hz applied with the 500Hz pulse generator, maximum
number of m will be 10. To have 500 pulses for averaging on these 10 data points, the required time for this process is 20ms \times 10 \times 500 = 100s. This long measurement time may cause inaccurate result as the charges profile especially at high voltage may change during this period.

To overcome these problems, a new system has been introduced [70,71]. In Tony Davies High Voltage Laboratory, this modification and improvement was done by Xu [72]. A specially built pulse generator that could produce 4kV narrow pulse of at most 5ns width as well as Eclipse signal averager console are used to give high sampling rate, high speed pulse and high speed data acquisition. Synchronisation of the pulse and ac waveform is provided by the function generator which sends trigger signal to both pulse generator and HV trek amplifier. Figure 4-9 shows the system set up that was used for ac measurement.

![Diagram of Eclipse PEA system setup for AC measurement](image-url)

Figure 4-9: The Eclipse PEA system setup for AC measurement [70].
The HV Pulse apply to PEA cell, Trigged by the Eclipse

Figure 4-10: The average set of pulses in one cycle for the new PEA system.

In the new system, the relation between the pulse generator frequency and ac signal frequency is described as $f_{pulse} = N \times m \times f_{ac}$. With sampling interval of 0.5, 1 or 2ns, the system could record up to 262,000 data points. The eclipse system has its own interlacing requirements for example 0.5ns interval and 10 records could give 40 averages. For an applied sine wave of 10 Hz and 2 kHz HV pulse, if 20 points are defined, 10 averages can be accomplished. To obtain 200 averages, 20 cycles is required and the system only take 2s to finish the measurement compared to the old system that took 400s (200x0.1x20=400s) for similar measurement [70].

4.4 Chapter Summary

In Tony Davies High Voltage Laboratory, space charge measurements used to be conducted using LIPP. However, due to its’ complex operation, this technique has no longer been employed. Some issues related to this equipment includes high frequency noise during ac stress, offset on the point on wave voltage, effect on the semicon thickness to measurement resolution, degradation of the target material as well as complicated signal
correction that performed after the measurement [73]. Since then, LIPP has been replaced by PEA technique due to simplicity, low cost, non-destructive and ease of implementation. This technique has shown its reliability to produce good data for space charge measurement.
Chapter 5   Sample preparation

5.1 Preparation of XLPE film

The polyethylene in pallet form, containing almost 5% of DCP was heated at 150°C in a pre-shaped mould as shown in Figure 5-1. Mould release liquid was applied on the mould surface to ease the removal process of the sample after it has been crosslinked. When the pallet melt, a pressure of 2 ton (equivalent to 0.2 Mpa) was applied for 30 seconds to push any bubbles out of the mould. The sample was then left to crosslinked at 200°C for 5 minutes. After that the mould was quenched cooled using tap water. The fresh XLPE was peeled out from the mould and cut into circular sample with a diameter of ≈19mm. The thickness of the samples made from this mould varies from 180μm to 210μm.

Figure 5-1: Mould tool [74]
To investigate the effect of byproducts in XLPE, the fresh XLPE was degassed in the vacuum oven at 80°C for 4 days. It was reported in [75] that degassing process may change the quality of the crystal forming more perfects crystals depending on the temperature, degassing period as well as the sample thickness. However, the amount of byproducts is more significant compared to the change of morphology in term of space charge formation since it was found that samples with comparable amount of byproducts but different morphology structure have almost similar charge profile. In order to confirm the existence of any byproduct in the fresh and degassed XLPE samples, the Fourier transform infrared (FTIR) spectrum has been used and this spectrum will be discussed later in this chapter.

Gulmine et. al reported that LDPE pallets with 1.5% of DCP that crosslinked at 170°C for 15 minutes produce 84% of gel content[76]. Thus, with the XLPE sample produced here, one can estimate that the gel content in the sample is as high as 80-90%.

5.2 Preparation Of Soaked LDPE Sample

All samples that were used in this part are supplied by GoodFellow [77] (supplier), with the density of 0.92gcm\(^{-3}\). The samples with a thickness of 180µm were cut into circular films. The diameter of the circular films varies depending on the electrode size of the measurement equipment. In this research, LDPE was chosen over XLPE to eliminate any possibility of having more than one byproduct in the sample. In addition, the possibility of having byproduct-free XLPE via degassing cannot be guaranteed [39]. Thus, LDPE is chosen over XLPE.

In the soaking process, the LDPE film acts like a sponge which means the byproducts will be absorbed by the film and settle in the space in between the crystalline structure as well as in the amorphous region. This condition is similar to the real XLPE cable where during the crystallisation process, the byproducts are displaced into the amorphous region. Hence, the tests could be conducted in a condition similar to the real cable.
5.2.1 *The Soaking Process*

All samples were soaked at room temperature except for those soaked into cumyl alcohol which were treated at 80ºC since the melting temperature for cumyl alcohol is 23ºC. The soaking process of the samples took 2 hours. To obtain optimum absorption, a small weight was used to keep the sample immersed in the chemicals. The absorption rate can be observed from the percentage weight increase and these values are plotted in the graph of Figure 5-2. The percentage weight increases rapidly during the first 20 minutes before it becomes steadier until the soaking process finish. After one hour, the percentage of α-methylstyrene is about 4 times larger than acetophenone due to low permeation velocity of the latter into LDPE. By having almost similar amount of acetophenone and α-methylstyrene in soaked sample, we could check if any difference on electrical properties of test sample is due to the different amount of byproducts in the sample. This result shows that by soaking LDPE sample into the chemicals for 2 hours, the amount of them in the sample is sufficient enough for electrical test to be conducted on it. These values are similar to the values that are reported in [37].

It is also worth to address here that this percentage weight of byproducts per sample film is not relative to the actual percentage of the byproducts in commercial XLPE which is much lower. These samples were soaked so that the distribution of byproducts was uniform across the sample thickness. The small increment of the byproducts in the samples after 20 minutes of soaking indicates that the byproducts were uniformly distributed in the sample. Hence, the big amount of byproduct is inevitable. In the soaked samples, the amount of acetophenone, cumyl alcohol and α-methylstyrene are 3, 8 and 500 times larger that in practice, respectively. This discrepancy should not a big problem in the research since the focus here is to identify the individual effects of the byproducts towards the electrical performance of LDPE. However, extra consideration on the percentage weight differences must be taken when the quantitative value from the measurement is to be used in real cable. Later in chapter 8, the percentage weights of the byproducts are considered in calculation of the conductivity increment by the byproducts.

It is also important to see the decay of the chemicals in the soaked sample so that we could see how long the chemicals could reside in the sample due to its volatility. The decay of
these chemicals in soaked LDPE can be observed in Figure 5-3. From the result, we learnt that the test should be conducted within one hour after the 2 hours of soaking to guarantee sufficient amount of chemicals in the tested sample so that the objective to see effect of the chemicals on the electrical properties of LDPE could be fully achieved. However, the chemicals could retain longer in the sample if it is in a closed environment. As in PEA method, the sample is placed in between 2 electrodes where there are only small room for the chemicals to evaporate. As in the graph below, it can be considered that the amount of chemicals is constant throughout the measurement.

Figure 5-2: The Soaking rate of acetophenone, \(\alpha\)-methylstyrene and cumyl alcohol in LDPE.
In dielectric spectroscopy and conductivity measurement, all samples were coated with gold to obtain good contact between the electrode and insulator. It is also observed in these measurements that gold coated samples produce better result (high signal-to-noise ratio) compared to the uncoated samples. The coating process was carried out before soaking to prevent any removal of the byproducts from samples due to the employment of vacuum during the gold coating process. The percentage weight of the byproducts in each samples are almost similar and these values are presented in Table 5-1. The specifications of samples used in each measurement are also presented in Table 5-2.

Table 5-1: Comparison of percentage weight of byproducts in coated and uncoated samples.

<table>
<thead>
<tr>
<th>Byproducts</th>
<th>Acetophenone</th>
<th>α-methylstyrene</th>
<th>Cumyl alcohol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncoated LDPE</td>
<td>2.09%</td>
<td>10.07%</td>
<td>11.36%</td>
</tr>
<tr>
<td>Coated LDPE</td>
<td>2.26%</td>
<td>9.33%</td>
<td>10.09%</td>
</tr>
</tbody>
</table>
Table 5-2: Samples specification for each measurement.

<table>
<thead>
<tr>
<th>Test/Measurement</th>
<th>Sample Thickness</th>
<th>Sample diameter</th>
<th>Gold coated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Space charge measurement</td>
<td>180 µm</td>
<td>36mm</td>
<td>No</td>
</tr>
<tr>
<td>Dielectric Spectroscopy</td>
<td>180 µm</td>
<td>25mm</td>
<td>Yes</td>
</tr>
<tr>
<td>measurement</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DC Conductivity test</td>
<td>50 µm (Clean LDPE )</td>
<td>50mm</td>
<td>Yes</td>
</tr>
<tr>
<td>AC Breakdown Test</td>
<td>50 µm</td>
<td>-</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>180 µm (soaked samples)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

5.3 Fourier Transform Infrared

In this research, FTIR was also used to study the byproducts in the sample. The infrared spectrum was obtained by applying the infrared radiation through the sample. Some of the radiation will be absorbed by the sample and some of it is transmitted. Depending on the responses from the vibrational spectroscopy, the molecule structure could be determined. Some vibration modes of the molecules are symmetrical stretching, anti-symmetrical stretching, scissoring, rocking and twisting. Figure 5-4 illustrates the vibration mode of CH₂ group. The resulting spectrum represents a fingerprint of a sample with absorption peak corresponding to the frequencies of vibration between the bonds constructing the material.
Figure 5-5 illustrates the principle of FTIR. FTIR consists of two mirrors located at a right angle to each other, oriented perpendicularly to a beam splitter at a 45° angle relative to the two mirrors. Radiation incident on the beam splitter is then divided into two parts, each of which propagates down one of the two arms and is reflected off one of the mirrors. The two beams are then recombined and transmitted out to the detector. When the position of one mirror is continuously varied along the axis of the corresponding arm, an interference pattern is swept out as the two phase-shifted beams interfere with each other [78].
According to quantum mechanics, these frequencies correspond to the difference in the energy between the ground state ($E_0$, lowest frequency) and several excited states ($E_n$, higher frequencies). The transition between these two states will involve the absorption of the light energy (determined by the wavelength) equal to the difference in energy between the two states [79]. FTIR is usually used to identify unknown materials, determine the quality or consistency of a sample, as well as determining the amount of components in a mixture. This technique has successfully adapted before [80].

FTIR spectrum can be used to detect the existence of the crosslinking byproducts in the sample. For instance, C=O for acetophenone, O-H or C-O for cumyl alcohol and C=C for $\alpha$-methylstyrene. To represent the main peaks and associated interactions for the entire wavenumber range, Table 5-3 has been formed.
Table 5-3: Wavenumbers and related vibrations compiled from [81], [82]* and [83]**

<table>
<thead>
<tr>
<th>Functional group</th>
<th>Description of Vibration</th>
<th>Wave number (cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>O-H</td>
<td>stretch, H-bonded</td>
<td>3200-3600</td>
</tr>
<tr>
<td>O-H</td>
<td>stretch, free</td>
<td>3500-3700</td>
</tr>
<tr>
<td>C-H</td>
<td>stretch</td>
<td>2850-3000</td>
</tr>
<tr>
<td>C=O</td>
<td>stretch</td>
<td>1670-1820</td>
</tr>
<tr>
<td>C=C</td>
<td>stretch</td>
<td>1400-1600</td>
</tr>
<tr>
<td>-C-H</td>
<td>bending</td>
<td>1350-1480</td>
</tr>
<tr>
<td>C=C</td>
<td>-</td>
<td>1375, 1263**</td>
</tr>
<tr>
<td>C-O</td>
<td>stretch</td>
<td>1050-1150</td>
</tr>
<tr>
<td>C-H</td>
<td>bending</td>
<td>936*</td>
</tr>
<tr>
<td>C=CH₂</td>
<td>twisting</td>
<td>578*</td>
</tr>
</tbody>
</table>

Figure 5-6 illustrates the FTIR absorbance spectrum of the fresh and degassed XLPE in comparison to LDPE spectrum. The spectrum confirms that 4 days of degassing at 80°C could not remove the whole byproducts as shown in the red line in the figure. Other than FTIR, the effect of degassing could also be observed by weight reduction throughout the degassing period such as reported in [9]. However, this technique will not divulge the exact byproducts left in the sample. More importantly, there is no accurate measure to tell that the sample is free from any byproducts. The small peak seen in the degassed XLPE spectrum from 2800-3000 cm⁻¹ may due to the C-H peak that was detected during the background scanning. This peak is considered as the measurement error.

The amount of byproducts in the sample could also be quantified by using the absorbance spectrum. Figure 5-7 shows the relation of the absorbance intensity with the soaking time. Absorbance level depends on the absorptivity of the species at that particular wavenumber, the path length as well as the concentration of the species. The absorbance level that presented in Figure 5-7 is the height of the peak corresponding to the molecular bond that exist in the byproduct. For acetophenone, peak of C=O was observed at 1694cm⁻¹ and peak of C=C in α-methylstyrene is observed at 1630cm⁻¹. Meanwhile the peak of O-H in cumyl alcohol appears to be at 3370cm⁻¹. The level of acetophenone and cumyl alcohol are
almost similar to what have been reported in [5] and [36]. However the absorbance level of \(\alpha\)-methylstyrene shows some deviation.

![FTIR absorbance spectrum of fresh XLPE, degassed XLPE, and LDPE sample.](image)

Figure 5-6: FTIR absorbance spectrum of fresh XLPE, degassed XLPE, and LDPE sample.

![Absorbance level of the crosslinking byproducts in the samples during soaking process.](image)

Figure 5-7: The Absorption level of the crosslinking byproducts in the samples during soaking process.
5.4 Chapter Summary

The importance to ensure only one byproduct exist in each sample is emphasised here. Therefore LDPE is chosen as the base material over XLPE film. The soaking process took 2 hours and the amount of the byproducts in each sample is measured using percentage weight as well as the FTIR absorbance level.

The percentage weight of byproducts in the test samples is higher than in practice. This limitation is unavoidable in order to obtain uniform byproduct distribution across the sample thickness. However, since the focus of this research is to gain knowledge on the influence of these byproducts on electrical properties, this big amount of byproducts will not affect the research aim. Furthermore, some calculation could be made to estimate the quantitative increment or decrement of the measured value by each percent of the byproducts.
Chapter 6  Space charge Measurement

6.1 The Measurement Procedures.

In the space charge measurement, a pulse voltage of 600 V with a duration of 5 ns was applied to the sample to generate an acoustic signal wave. Two types of base materials were used in this measurement; XLPE and LDPE. For the former material, fresh and degassed XLPE (degassed for 4 days at 80°C) were stressed at 8kV for 2 hours followed by an hour of decay.

For LDPE based sample, all samples were soaked into the chemicals for 2 hours and subsequently applied to 3 different positive dc voltages which were 5kV, 8kV and 10kV. For space charge measurement, PEA technique was employed. Readings were taken using the easy data software in Lab View environment for every 10 minutes. In this measurement, semiconductor (top electrode, anode) and aluminium (bottom electrode, cathode) were applied directly to the specimens.

Before the sample was stressed at the specified voltage, 2kV dc voltage was applied across the sample for calibration purpose which has been discussed in Chapter 3 before. The charge density pattern at 2kV was then used in the PEA software as the reference signal in software written in Lab View TM environment.
During charging process, two measurements were conducted which is volts on and volts off measurements. For volt on, the measurement was taken while the power was still ‘on’, whiles for volt off measurement, the reading was taken 10 seconds after the dc voltage was removed (short circuit condition). As soon as the reading was taken, the sample was recharged again until the next reading time, for 1 hour of total charging period. Consequently, the power supply was permanently removed and the sample is discharged. This time, the decay of the accumulated charge in specimen was measured for another hour. For a better understanding on the volt on and volt off measurement procedure, Figure 6-1 could be referred.

![Figure 6-1: The volt on and volt off measurement](image)

6.2 Measurements With And Without Voltage Applied

Volt on measurement will include both fast and slow moving carriers. Meanwhile, for volt off measurement, the results will only show the slow moving carries. These fast and slow moving carries signify the path in which the carriers will take towards the opposite polarity electrode. The fast moving carriers are the carriers that are free from trap or those which
escape from the traps very shortly after the dc voltage is removed. On the other hand, the slow moving carriers are the carriers that are trapped in the deep traps and thus their movement are very limited depending on the energy that is required to overcome the barrier. As the power supply is disconnected, the carriers with energy less than the barrier will stay in the traps.

6.2.1 Fresh XLPE

![Graph](image)

Figure 6-2: The space charge profile of fresh XLPE during a) volt on and b) volt off.

The volt on result for fresh XLPE sample shows that the formation of homocharge in the vicinity of the electrodes. As the sample is stressed for longer time, more negative charges
are seen penetrated into the sample bulk. When the voltage is removed from the sample, it is observed that heterocharges are accumulated very near to the electrodes. The heterocharge peaks are increasing with the stressing time and becomes constant after 45 minutes. In term of the bulk charge, electrons or negative charge carrier dominate this region. The charge profile of the fresh sample is presented in Figure 6-2.

Figure 6-3: The space charge profile of degassed XLPE during a) volt on and b) volt off.

The volt on charge profile for the degassed XLPE sample (Figure 6-3(a)) shows the formation of positive heterocharge near the cathode. The positive peak near electrode, as well as the spread of positive charge in the sample bulk increase by time. Volt off result in Figure 6-3 (b) reveals that there is a small amount of negative heterocharge accumulated near the anode. However these negative hetero charges get smaller by time. From both
results, it is seen that the XLPE samples causes the formation of heterocharge in the sample. The fresh XLPE sample has more charge injection in the sample hence out numbering the heterocharge that actually accumulated near the electrodes. The amount of heterocharge is also less in the degassed sample due to less byproduct left in the sample. These charges may be due to the remaining byproducts that still reside in the XLPE sample. Equivalent results were reported in [84]. After degassing XLPE sample for 3 days at 90°C, Fu et. al also observed charges in degassed sample although instead of heterocharges, homocharges was observed near electrode.

Figure 6-4: The decay charge profile of a) fresh XLPE and b) degassed XLPE.

The charge decay in fresh XLPE is seen to be very fast in the first 30 minutes and become slower after that. The charge profile after decaying for 60 minutes is very similar to that of 30 minutes. The charge decay characteristic of the degassed sample is not straight forward.
The negative charge near the cathode may be drifted into the electrode revealing the positive charges that reside in the region. As a result, the positive peak appears in the vicinity of the cathode. The space charge profile of fresh and degassed XLPE do not give much understanding on the characteristics of the byproducts in term of space charge accumulation. Hence, separate investigation on each byproduct is crucial.

6.2.2 Clean LDPE

Before we could analyse the accumulation of space charge in the soaked sample, it is important to study the charge behaviours in the clean LDPE so that the analysis of the space charge accumulation will not be deluded. The results of clean untreated LDPE will be used as the reference to the results obtained for soaked samples. The charge build up in the clean LDPE can be seen in Figure 6-5.

a) Clean LDPE stressed at 5kV,
b) Clean LDPE stressed at 8kV,

c) Clean LDPE stressed at 10kV,

Figure 6-5: Charge Density of 180µm clean LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt ON condition

This observation is consistent in the volt off measurement results and in fact, the charge build up could be seen even clearer. These results are presented in Figure 6-6. For the sample stressed at 5 kV, the increasing stress time results in the charges of both polarities gradually increasing in the sample. Negative charges are trapped in the region near to the cathode which stops them to move towards the anode. It is also observed that the injected positive charges tend to move towards the cathode due to different polarity attraction. At 8kV and 10kV, a greater volume of negative charges injected from the cathode and slowly
Figure 6-6: Charge Density of 180µm clean LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt OFF condition.
the charges drifted in the sample bulk. It is clear that the meeting point of positive and negative charges moves towards the middle of the sample with increasing applied voltage. The gradual increase of positive and negative charges is similar to the result in [30].

6.2.3 Acetophenone Soaked LDPE

For acetophenone soaked LDPE sample, the volt off results at the three voltages are illustrated in Figure 6-7. This time, the domination of negative charges is obvious. In the vicinity of the positive electrode, the amount of positive charge decreases with time indicating either the positive charges migrated towards the nearby electrode, or the amount of negative charges is greater than that of positive charges at that specific area. It is observed that the amount of negative charges that move into the bulk of the sample, towards to anode, is proportional to the stressing time. This observation can be seen at all three stressing voltages and more noticeable at higher voltage. As the positive charges from the anode are suppressed, one observes distinguishable differences in the charge profile of LDPE soaked in acetophenone compared to an untreated sample.

![Graph](image)

(a) Acetophenone soaked LDPE stressed at 5kV,
Figure 6-7: Charge Density of 180µm acetophenone soaked LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt OFF condition.

The negative charge build up is also seen in the volt on measurement result at 8 kV and 10 kV. This result is presented in Figure 6-8. In sample stressed at 5kV, any charge build up could hardly been seen.
Figure 6-8: Charge Density of 180µm acetophenone soaked LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt ON condition.
6.2.4 \( \alpha \)-methylstyrene Soaked LDPE

Volt on measurement results for \( \alpha \)-methylstyrene soaked LDPE is presented in Figure 6-9 meanwhile charge density during volt off, is illustrated in Figure 6-10. The charge profiles for \( \alpha \)-methylstyrene soaked LDPE and untreated clean sample are almost similar except that the positive charges in clean untreated LDPE increases gradually compared to the easy domination of positive charge in the \( \alpha \)-methylstyrene soaked sample. Due to that, we could say that \( \alpha \)-methylstyrene assists the movement of positive charges into the sample bulk.

\[\text{\( \alpha \)-methylstyrene soaked LDPE at 5kV}\]

\[\text{\( \alpha \)-methylstyrene soaked LDPE at 8kV}\]

80
c) α-methylstyrene soaked LDPE at 10kV

Figure 6-9: Charge Density of 180µm α-methylstyrene soaked LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt ON condition.

In term of magnitude, samples stressed at 5kV and 8kV shows almost the same amount of charge accumulation in sample. At 10kV (Figure 6-10(c)) positive charges have already accumulated in the bulk of the sample as early as the first 10 minutes of stressing. Similarly to Figure 6-10, authors have reported [5] a rapid migration of positive charges in a α-methylstyrene soaked sample, it is concluded that α-methylstyrene has a greater effect on the positive charge migration than acetophenone as it can permeate faster into the polymer structure.
Figure 6-10: Charge Density of 180µm α-methylstyrene soaked LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt OFF condition.
6.2.5  *Cumyl Alcohol Soaked LDPE*

The space charge characteristic of cumyl alcohol soaked LDPE is different from the two former chemicals discussed. The charge density of the sample during volt on and volt off are shown in Figure 6-11 and Figure 6-12 respectively.

Cumyl alcohol causes charge accumulation of both charge polarities with heterocharges appeared when the sample is stressed at 5kV and 8kV. Cumyl alcohol introduces more charge injection into the sample compared to the other byproducts. Both positive and negative charges were introduced into the sample. In Figure 6-11 (a) and (b), after 10 minutes charging, the negative charges migrated into the bulk, reducing the height of the positive peak near cathode. After 20 minutes, the positive peak is totally disappears and negative charges are now accumulated in the bulk of sample. Similar observation is seen for the negative peak near the anode. However, as the voltage is increased, heterocharges begin to diminish gradually due to charge injection from the electrodes. This phenomenon is reported in [85]. Heterocharges are also observed in cumyl alcohol soaked sample that reported in [86] but not discussed.

![Diagram of Charge Density vs Thickness](image)

(a) cumyl alcohol soaked LDPE stressed at 5kV
b) cumyl alcohol soaked LDPE stressed at 8kV

c) cumyl alcohol soaked LDPE stressed at 10kV

Figure 6-11: Charge Density of 180µm cumyl alcohol soaked LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt ON condition.
Figure 6-12: Charge Density of 180µm cumyl alcohol soaked LDPE; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, during Volt OFF condition.
6.3 Charge Decay Results

6.3.1 Clean LDPE

By monitoring charge decay after the applied voltage is removed, one can see the influence of these byproduct chemicals on the charge transportation in the insulator. Figure 6-13 shows the changes in charge profiles in clean LDPE after the removal of the applied voltage.

Charge decay in clean LDPE after stressed at 5kV,

b) Charge decay in clean LDPE after stressed at 8kV
c) Charge decay in clean LDPE after stressed at 10kV

Figure 6-13: Charge Decay in 180um clean LDPE after; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, for 1 hour

The clean LDPE shows a very slow charge decay rate as more charges are left in the bulk of sample even after 60 minutes. The negative charges near the top electrode shows an increment in density as more positive charges possibly leak into the anode, leaving the negative charges in the vicinity of electrode.

6.3.2 Byproducts Soaked LDPE

Figure 6-14 to Figure 6-16 show the changes in charge profiles in the soaked LDPE samples after the removal of the applied voltage started by acetophenone followed by α-methylstyrene and cumyl alcohol soaked LDPE.
Figure 6-14: Charge Decay in 180µm acetophenone soaked LDPE after; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, for 1 hour
Figure 6-15: Charge Decay in 180µm α-methylstyrene soaked LDPE after; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, for 1 hour.
Figure 6-16: Charge Decay in 180µm cumyl alcohol soaked LDPE after; a) stressed at 5kV, b) stressed at 8kV, c) stressed at 10kV, for 1 hour
For acetophenone, α-methylstyrene and cumyl alcohol soaked samples, the decays are extremely fast especially in the first 10 minutes. As the decay results are referred, the byproducts assist the movement of the carriers in LDPE and reduce the number of charges trapped in the existing traps of the polymer. This argument is supported by the results from volt off measurement, where more charges could drift into the bulk instead of trapped at the adjacent of the electrodes.

By integrating the space charge density over the insulation thickness at a certain time \( t \), \( \rho(x, t; E) \) the amount of charge in the sample, \( q(t; E) \) is obtained [87].

\[
q(t; E) = \int_0^L \rho(x, t; E) | \cdot S \cdot dx
\]  
Equation 6-1

where 0 and L denote the position of the electrode excluding the charges at the electrodes, E is the electric field and S is the electrode area. The charge profile of the sample could be analysed as the total charge value during charging and decay is plotted over time. The results of this calculation are presented in Figure 6-17.

The effect of the chemicals on the total charge is the utmost at elevated voltage and no big difference is observed at lower voltage. This is shown in the left graphs in Figure 6-17. In samples charged at 5kV and 8kV, cumyl alcohol causes more charge accumulation as compared to that of the clean LDPE. Acetophenone and α-methylstyrene reduce the amount of total charge accumulated in the LDPE. At 10kV, acetophenone and cumyl alcohol introduce more charge accumulation to the clean LDPE.

After comparing all soaked samples and the clean LDPE, α-methylstyrene appears to cause the least effect on the charge accumulation in the insulator. In all three stressing voltages, the smallest amount of charges is found in α-methylstyrene soaked LDPE and the value is smaller than that of clean LDPE sample. This result is similar to the result reported in [88] where it was concluded that α-methylstyrene has no contribution to the space charge formation. On the other hand, cumyl alcohol and acetophenone show a clear contribution to the charge built up in the insulator, especially at high voltage.
Figure 6-17: The total charge in the samples that stressed at 5kV, 8kV and 10kV during charging (LEFT) and decay (RIGHT) process.
6.3.3 Rate of Charge Decay

The rate of charge decay could give an indication of charge mobility and thus the trap depth. Despite the increase of charge injection and accumulation in the insulator, the byproducts speed up the decay process particularly in the first 15 minutes. From this point, the charge decay is comparable to the untreated LDPE. This decay could be seen in Figure 6-17. From the charge decay results, LDPE soaked in α-methylstyrene has the greatest rate of decay followed by acetophenone and cumyl alcohol. α-methylstyrene results in 80% of the original charges within a sample to decay. As the rate of decay is commonly associated with shallow and deep traps, where rapid decay is observed in the former [89], one can conclude that the chemical byproducts reduce the number of charges trapped in the deep traps. This is demonstrated by a slower rate of decay in a smaller number of charges, as seen in the decay curves. Based on the results one assumes that the charges will initially fill the shallow traps prior to the deep traps. This hypothesis has been verified recently [90]. It is reasonable to suggest that the byproduct chemicals modify the trapping characteristics of LDPE in two possible ways which are,

1) by increasing the population of the shallow traps, and

2) by reducing the number of deep traps in the LDPE, replacing them with the shallow traps.

In untreated LDPE, deep electron traps are often found in regions of reduced density, such as sub-microvoids with surrounded polymer chains [48]. These voids and naturally occurring spaces between the crystalline structures will fill with chemicals in the soaking process and result in a change in the charge trapping and de-trapping characteristics associated with the region. This is dependent on the type of bonding the chemical byproducts induce within the system such as benzene ring, carbonyl group, double bond, and hydroxyl groups [37].

It should be noted that the evaluation made here is contradict to the report in [91] which associates the crosslinking byproducts to deep traps. This report is based on the calculations of electron affinity and trap depth. This trap depth values however have been obtained without considering the polarisation that neighbouring molecules induced on the model molecule which is non negligible in general cases [92].
6.4 Two Chemicals In One Sample

The analysis made above is based on an assumption that the chemical function of the byproducts molecule act separately from each other. At this stage, it is difficult to achieve a similar mixture ratio to that in practice via soaking process. This is because of the difficulty to control the amount of byproducts in the soaked sample and at the same time have a uniform distribution across the film thickness. The main concern here is to examine potential interactions amongst different byproducts. As it is difficult to see these effects with a mixture of three byproducts, only two byproducts are mixed at one time. The results are compared to those obtained in the previous section.

Similar procedures as before, including the soaking, charging and decaying processes were applied to the samples. However in this part, the samples were stressed at one stressing voltage which is at 8kV. With 3 different chemicals, we have 3 different combinations of the chemicals, which have been simplified in Table 6-1 below. The volt off measurement results are presented in Figures 6-18 to 6-20.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Chemical</th>
</tr>
</thead>
<tbody>
<tr>
<td>a+α</td>
<td>acetophenone plus α-methylstylene in LDPE</td>
</tr>
<tr>
<td>a+c</td>
<td>acetophenone plus cumyl alcohol in LDPE</td>
</tr>
<tr>
<td>α+c</td>
<td>α-methylstylene plus cumyl alcohol in LDPE</td>
</tr>
</tbody>
</table>
Figure 6-18: Volt off measurement for Sample a+α

Figure 6-19: Volt off measurement for Sample a+c
Figure 6-20: Volt off measurement for Sample α+c

For Sample a+α (Figure 6-18), the Volt profile illustrates how negative charges dominate the sample bulk. This profile is similar to data obtained from acetophenone soaked LDPE, which suggests that acetophenone has a greater influence than α-methylstyrene in terms of controlling the charge build up. The former byproduct suppresses positive charge injection from the anode. Similar characteristic is observed in Figure 6-7(b).

In Figure 6-19, the domination of acetophenone over the other byproduct is also shown in Sample a+c. Although negative peak is only observed near to the cathode, the amount of positive charges that reduce with charging time and the positive peak moves towards the anode showed that the charges are suppressed. This observation is believed to be related to the existence of acetophenone in the sample.

No heterocharges appeared in Sample a+c and Sample α+c (Figure 6-20), although cumyl alcohol presents in the samples. In Sample α+c, positive charges are trapped in the vicinity of the anode and at the same time, the negative charges migrate from the cathode to the sample bulk. As a result, more negative charge is seen in the sample. It is believed that some positive charges are trapped in the vicinity of the anode, this is consistent with the space charge profile in α-methylstyrene and cumyl alcohol soaked LDPE. However, due to low resolution, these charges appear in only one positive peak. As the charge dynamics do
not show any distinctive pattern associated with any of the individual byproducts, a conclusion cannot be drawn as to which byproduct is the most dominant.

After one hour charging, the samples were left to decay. Again, the amount of charges in samples is calculated and the results are shown in Figure 6-21. The results for sample soaked into single chemical that stressed at 8kV are also included for comparison purpose.

![Figure 6-21: Total charge accumulated in sample during decay in samples.](image)

After being stressed for an hour, the total charge accumulated in Sample a+α is the highest followed by Sample α+c and Sample a+c. Once again, we could observe the similar slow decay after about 20 minutes open circuit, which is due to the deep traps in the polymer. Despite having the least total charge, the rate of charge decay in Sample a+c is low relative to the other two samples. It is believed that the presence of α-methylstyrene in samples a-α and α+c cause a faster rate of decay. The result indicates that in terms of charge accumulation, it is rather difficult to see which byproduct causes the most charge injection into the insulator. However, α-methylstyrene maintains its characteristic as the charge decay accelerator while in the existence of other byproducts in the insulator.
6.5 AC Space Charge Profile

Similar to DC measurement semicon and aluminium were used as the top and bottom electrodes respectively. Samples were stressed at 4kV peak voltage which equivalent to 28kV\textsubscript{rms}/mm, for 4 hours at the frequency of 50Hz. Space charge profile of the sample was collected in every hour. This measurement is followed by 20 minutes of charge decay. 800V electric pulse at 2 kHz was applied to produce the acoustic signal. Due to very small data signal that will be obtained in this test especially for short circuit measurement, data need to be denoised before further analysis could be done.

Volt on results show no significant difference in the charge profile of the clean LDPE and the soaked samples. These results are included in the Appendix. Hence, the Volt off measurement results are referred to study their differences. Figure 6-22 shows the charge profile of all samples during volt off condition.

![Graph showing charge density over position]

\textbf{a) Clean LDPE,}
b) Acetophenone soaked LDPE,

c) α-methylstyrene soaked LDPE

d) Cumyl alcohol soaked LDPE

Figure 6-22: Charge Density of a) Clean LDPE, b) acetophenone soaked LDPE, c) α-methylstyrene soaked LDPE and d) cumyl alcohol soaked LDPE during Volt off condition after 28 kV/mm ac.

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In the clean LDPE sample, positive and negative charges are observed in the bulk. The positive peak is near to the aluminium electrode and negative peak is near to the semicon electrode. However the positive peak is wider than the negative peak. Both of these peaks amplitude reduce with stressing time. This reduction may due to more charges recombined during the second half cycle compared to the charge injected into the film. Meanwhile in acetophenone soaked LDPE, more negative charges reside in the bulk of the sample. The negative charge decreases with time within the first 3 hours of stressing. The charges suddenly get bigger after 4 hours. Positive and negative peak is also observed in the vicinity of semicon and aluminium electrodes respectively.

In α-methylstyrene soaked LDPE negative charge is also detected in the vicinity of aluminium electrode and penetrates into the sample bulk. Positive peak is observed near the semicon electrode. In the presence of α-methylstyrene, the negative charge in the bulk as well as the positive peak near the top electrode increase by time. On the other hand, negative charges near aluminium electrode become lesser by time. Cumyl alcohol soaked sample is the only sample that shows a total domination of negative charge in the sample. The negative peak near the semicon electrode gets higher as the charging time gets longer.

It is difficult to have a direct comparison between the samples that were stressed at ac and dc in term of charge density profile. Although both tests used similar electrode system, the difference on the material for top and bottom electrode making the analysis on the charge injection in samples during ac stress becomes very complicated. Hence, total charge in every sample was calculated by integrating the charge density over the sample thickness. Figure 6-23 shows the total charge decay in samples stressed at ac.
The charge decay profiles for ac stressed samples were recorded up to 20 minutes (1200 s). The fluctuation that observed in acetophenone soaked LDPE and cumyl alcohol soaked LDPE is most likely due to noises that appear in the data. The total charge accumulated during ac stress is much lower compared to the total charge under dc condition. This is due to the recombination of the majority of electrons and holes alternately injected each half cycle [93]. Despite this small amount of charge, the accumulation of space charge under ac condition should not be neglected particularly at lower frequency where the formation of space charge is more severe [94].

Charge decay profile of ac stressed samples is not as direct as that of dc stressed samples. Overall, the results suggested that after a few hours stressed, the charges movement in the samples become very static. The rate of charge decay is very small. This observation proves that the charges left in the sample during decay process are the charges that were trapped in the deep traps.

**During DC Stress**

From dc space charge measurement, it was observed that the byproducts have contributed to very fast initial charge decay followed by slower decay. It is suggested that the byproducts either (a) increase the number of shallow traps in the samples, or (b) reduce the density of deep traps in the sample. With continuous current in one direction, the charges
will initially fill into the shallow traps and conduct through the dielectric to the opposite polarity electrode. Accordingly, the conduction current is higher for the byproduct soaked samples compared to the clean LDPE.

**During AC Stress**

In ac, charges will move back and forward due to the polarity change of the electrodes. It is in agreement with the results obtained in dc test that the byproducts introduce more shallow traps in the sample causing higher traps density in the sample.

During the first half cycle, the charges will be injected into the sample. Since higher shallow trap density found in byproducts soaked LDPE, there is higher probability of the charges to be trapped either in shallow or deep traps. However due to the polarity reverse, charges trapped in shallow traps can be easily extracted into the electrode. The remaining charges will shift further into the bulk, moving back and forward due to the polarity reverse and eventually trapped into the deep traps.

With more shallow traps, the crosslinking byproducts ease the movement of the charges further into the bulk and have higher probability to be trapped into the deep traps. As a result, samples with crosslinking byproduct posses more charges compared to the clean sample. The dependency of charges trapping dynamics on the total number of shallow and deep traps is mentioned by Chen et. al in [90] analysing the trapping and detrapping characteristics of polymer by using PEA technique. The analysis however does not consider the trapping process from shallow to deep traps. It is clear now, that this process should be taken into account when analysing the trapping and detrapping process especially during ac stress.

6.6 Chapter Summary

It was observed from comparison between the fresh XLPE and degassed XLPE that both samples have heterocharge accumulation near the electrodes. With more byproducts in the sample, fresh XLPE sample has more injected charge in the sample.
The homocharges are found in acetophenone and α-methylstyrene soaked LDPE. However, acetophenone causes negative charge to accumulate in the sample bulk, meanwhile α-methylstyrene allows the positive charge to dominate the sample bulk. Cumyl alcohol is the only byproduct that introduces heterocharges in the LDPE sample. These heterocharges become less after more charges injected into the sample and over cover the heterocharges. In term of total charge, α-methylstyrene soaked LDPE has shown the least charges accumulated in the sample.

With the existence of 2 byproducts in one sample, acetophenone is dominant in determining the profile of the charge accumulation. Any sample with α-methylstyrene has the fastest charge decay. Charge decay profiles from dc space charge measurement reveal that the crosslinking byproducts introduce shallow trap into the LDPE structure. During dc stress, these shallow traps assist the movement of charges across the polymer thickness causing fewer charges trapped in deep traps.

During ac stress, the shallow traps in crosslinking byproduct cause more charges to be trapped in the polymer. With more charges trapped and move further into the bulk, hence there is higher probability for the charges to be trapped in the deep traps due to the polarity reverse that happen throughout the stressing period.
Chapter 7  Dielectric Spectroscopy Measurement

7.1  Introduction

The electrical properties are determined by the degree of polarity of a material. A dielectric like polyethylene which only consists of carbon and hydrogen or methylene chains is non-polar in nature. Such material has low conductivity value. If any polar component exists in the chain, the polymer will then become more polar and its characteristics are no longer as explained before. When a polymer is subjected to an electrical field, the polymer will become polarised where the polar parts of the chain will respond to the electric field applied. As a result, the polymer will be pulled into two directions due to two charge polarities in the chain. Although ideally polyethylene is a non-polar material, in real world compositions there are always small amount of polar components such as impurities, which present in the polymer structure. Under electric field application, these impurities will also experience polarisation [95].

In the case of ac, we will see that the polar chain will be shifting back and forth according to the polarity change. The rate of changes of the alignment of this chain is controlled by the frequency. As the frequency increases, it occurs at one point that the chain cannot ‘keep up’ with the changing frequency. With various functional groups in the polymer, different groups will be sensitive to different frequencies. As the frequency increases, no changes will be seen in real permittivity value ($\varepsilon'$) provided that the dipole can respond. When the dipole could no longer rotate as fast as the changing field, charges cannot be held and the dielectric constant value changes with frequency. This phenomenon not only
occurs in a polar structure, but also occurs in other electrical polarisation such as electronic, atomic and orientational polarisation. Figure 7-1 and Figure 7-2 illustrate the frequency response of different dielectric mechanisms across the measured frequency.

**Figure 7-1**: Three main polarisations from lower to higher frequency [96].
Permittivity ($\varepsilon'$) represents the ability of the material to polarise by an electric field. Dissipation factor or dielectric loss ($\tan \delta$) is a measure of the amount of energy loss as heat rather than transferred as electrical energy. For a good insulator, the value of $\varepsilon'$ and $\tan \delta$ should be low. To understand the concept of permittivity, it is easier to first discuss a closely related property, capacitance ($C$). Consider a dielectric sandwiched between two parallel metal plates. The capacitance and conductance ($G$) for the dielectric are [97]:

\[
C = \frac{\varepsilon \varepsilon_0 A}{d} \quad \text{Equation 7-1}
\]

\[
G = \frac{\sigma A}{d} = \frac{1}{R} \quad \text{Equation 7-2}
\]

where $A$ in $\text{m}^2$ is the area of electrodes, $d$ in $\text{m}$ is the electrode separation, $\sigma$ in $\text{W}^{-1}\text{m}^{-1}$ and $R$ in $\Omega$ is the resistance. However, in ac voltage waveform, the ‘formulae’ is not as simple as that in dc voltage. It is more convenient to use a vector diagram to describe these two properties with real and imaginary part as in Figure 7-3.
Figure 7-3: Vector diagram of the electrical response of a dielectric [97].

The phase angle $\theta$ represents the angle by which the current waveform leads the voltage waveform. The impedance $Z$ of a dielectric is given by:

$$\frac{1}{Z} = \frac{1}{R} + j\omega C$$  \hspace{1cm} \text{Equation 7-3}

The complex dielectric function $\varepsilon^\ast(\omega)$ is determined from:

$$\varepsilon^\ast(\omega) = \varepsilon' - j\varepsilon''$$  \hspace{1cm} \text{Equation 7-4}

where $\omega=2\pi f$ is the angular frequency. One could also express the complex capacitance $C^\ast$ of a dielectric as:

$$C^\ast = \varepsilon^\ast C_0$$  \hspace{1cm} \text{Equation 7-5}

Where $C_0$ is the capacitance of the cell with air given by $C_0=\varepsilon_0 A/d$ for a parallel plate capacitor and the impedance $Z$ is given by:

$$Z = \frac{1}{j\omega C^\ast} = \frac{1}{j\omega C_0 (\varepsilon' - j\varepsilon'')} = \frac{R}{1 + j\omega CR}$$  \hspace{1cm} \text{Equation 7-6}

From equation 7-6, equating the real and imaginary parts, we obtain the real permittivity as expected,
And

\[ \epsilon'' = \frac{1}{\omega \epsilon_0 R} = \frac{\sigma}{\omega \epsilon_0} \]  

Equation 7-8

And thus

\[ \tan \delta = \frac{\epsilon''}{\epsilon'} = \frac{1}{\omega RC} = \frac{\sigma}{\omega \epsilon' \epsilon_0} \]  

Equation 7-9

Cole-Cole plots has been introduced by Kenneth and Robert in 1941 [98]. Cole-Cole plot is produced by plotting the imaginary permittivity against real permittivity in log-log plot which produce a curve that normally in a semicircular shape (Figure 7-4) for a fully Debye relaxation process. This plots is used to determine the zero-frequency dielectric constant \( \epsilon'_0 \) and the limiting high frequency value \( \epsilon''_\infty \) [99]. Equation that defines the Cole-Cole plots is given by;

\[ \epsilon = \epsilon_\infty + \frac{\epsilon_s + \epsilon_\infty}{1 + (i\omega \tau)^\alpha} \]  

Equation 7-10

Here \( \epsilon_s \) is the static permittivity (at 0Hz), \( \epsilon_\infty \) is the permittivity at maximum frequency, \( \tau \) is the relaxation time and \( \alpha \) is the Cole exponent, which have a specific value for different material. Good background on Cole-Cole plots is described in [100].

Figure 7-4: Semicircular Cole-Cole plot.
7.2 Measurement procedures

Dielectric measurement was performed on the liquid byproduct and the byproducts soaked LDPE. Hence different cells are required for the solid sample and liquid sample. Figure 7-5 shows the cells used for solid and liquid samples in this measurement. In both measurements, a Solartron 1296 Dielectric Interface and Schlumberger SI 1260 Impedance/phase gain analyser were used. Dielectric spectroscopy of the samples was measured at room temperature. During measurement, the dielectric cell is placed in a metal box to avoid any unwanted noise to interfere with the measurement. To have a good contact between the sample and electrode, the film samples are initially coated with gold prior to the soaking process. By applying 25mA current for 4 minutes, the samples are coated by 32nm thick of gold on each side. The parameters used in this test are shown in table 7-1.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value/Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency</td>
<td>10mHz -10MHz</td>
</tr>
<tr>
<td>AC Bias</td>
<td>1V</td>
</tr>
<tr>
<td>DC Bias</td>
<td>0V</td>
</tr>
<tr>
<td>Integration</td>
<td>10 cycle</td>
</tr>
<tr>
<td>Sample Area</td>
<td>Circular, 25mm diameter (S)</td>
</tr>
<tr>
<td></td>
<td>Circular, 30mm diameter (L)</td>
</tr>
<tr>
<td>Sample Thickness</td>
<td>0.18mm (S)</td>
</tr>
</tbody>
</table>
Figure 7-5: (a) The solid and liquid cells used in experiment and (b) metal box to avoid signal from noise.

7.3 Dielectric Measurement Results For Liquid Samples

The real and permittivity values for the liquid byproducts are calculated from equation 7-7 and the results as well as the imaginary permittivity values are shown in log-log plot in
Figure 7-6. At low frequency, the permittivity value of acetophenone is the highest ($≈10^6$) followed by cumyl alcohol ($≈10^5$) and last but not least $\alpha$-methylstyrene ($≈1$). As the frequency increases, the real permittivity value is reduced rapidly. From the literature [101], the static permittivity values of acetophenone, $\alpha$-methylstyrene and cumyl alcohol are 17.44, 2.28 and 5.61 respectively. It is clear from Figure 7-6 that the measured values are too high that are not acceptable at some frequencies. This enormous values are contributed by the combination effect of DC conductivity and the anomalous conductance due to the polarisation process [96]. As a result, the equipment that measured the impedance value which was largely contributed by the resistance had misinterpreted it as the capacitive value hence giving high dielectric constant. The dielectric constant values that are acceptable for acetophenone, cumyl alcohol and $\alpha$-methylstyrene liquids from Figure 7-6 are from 50 Hz, 0.3 Hz and 0.01Hz onwards respectively. In these range of frequencies, the dielectric constant values are more acceptable and closer to theoretical values.

Figure 7-7 shows the tan $\delta$ values for the byproducts liquids. For the tan $\delta$ curve, after comparing the values to that of the empty cell, the relaxation peak for $\alpha$-methylstyrene, cumyl alcohol and acetophenone are observed at frequency range of $10^{-2}$ to $10^{-1}$ Hz, $10^{-1}$ to 1 Hz and $10^{2}$ to $10^{3}$ Hz respectively.

Figure 7-6: The real part of $\varepsilon'$ of acetophenone, $\alpha$-methylstyrene and cumyl alcohol.
7.4 Dielectric Measurement Results For Solid Samples

The relative permittivity value obtained for the clean LDPE is \( \approx 2.31 \). Similar measurement was conducted on uncoated LDPE sample. As a result, permittivity value that obtained is \( \approx 2.1 \) which is quite low compare to the theoretical value which typically lies in the range of \( 2.25 < \varepsilon_r < 2.3 \). Therefore, it is quite essential to gold coat the sample film in order to obtain a more accurate film.

The additional byproducts in the material cause small increment on the permittivity value. Among the three byproducts, acetophenone gives the highest permittivity value, followed by \( \alpha \)-methylstyrene and cumyl alcohol. Despite this small effect, the \( \tan \delta \) curves show some changes when the byproducts exist in the LDPE film. The value of \( \varepsilon' \) and \( \tan \delta \) values are presented in Figure 7-8 and Figure 7-9 respectively.
It is observed that acetophenone have very similar tan δ values of that of the clean LDPE. α-methylstyrene increases the tan δ value in the range of 0.1Hz-10Hz. The increment is however very little. On the other hand, cumyl alcohol increases the ε” and tan δ values especially at lower frequency. This is obviously not the actual values corresponding to the
bulk dielectric permittivity characteristic of cumyl alcohol soaked LDPE but rather are due to the charge build up at interfaces between the bulk of the sample forming spurious boundary layers (interfacial or Maxwell-Wagner-Sillars (MWS) polarisation [96]) and between the sample-electrode interface (Space charge polarisation [102]). At lower frequency, there is more time for the charges to move in and build up at the sample-electrode interface within the half-cycle of the applied AC, giving rise to the measured capacitance and hence the effective dielectric permittivity values. On contrary, this will not happen at high frequency as very short time is available for the charge movement in one half-cycle, the charges could not keep up with the change of electric field and thus only bulk polarisation mechanism contribute to the electric polarisation. The Cole-Cole plots of the clean sample as well as the byproducts soaked sample is presented in Figure 7-10. The Cole-Cole plots do not form a semicircular shape indicating the samples do not undergo Debye relaxation process. However it is shown here that cumyl alcohol has a large increase in conduction compared to the other samples.

![Cole-Cole Plots of the clean LDPE and the byproducts soaked LDPE.](image)

Figure 7-10: Cole-Cole Plots of the clean LDPE and the byproducts soaked LDPE.

With the results obtained in permittivity and dielectric loss spectrum, cumyl alcohol is proved to introduce either more mobile carriers into the insulator, or increase the charge mobility in the insulator [103]. The frequency-dependent real part of alternating current conductivity \( \sigma_{ac} \) of the dielectric is obtained from the relation [104]:

\[
\varepsilon'' = \frac{1}{\varepsilon' - j \omega \varepsilon_0}
\]
From this equation, the conductivity values are calculated and presented in Figure 7-11 below. Acetophenone and α-methylstyrene give almost no effect on the conductivity values. As expected cumyl alcohol increases the conductivity of LDPE film. This result however, shows the conductivity value when AC current is applied to the sample which may not be directly equal to the result obtained from DC conduction current measurement. Polarisation that might occur under the influence of AC current may delude the actual conductivity value of the samples. A DC characteristic may be obtained from dielectric spectroscopy at a very low frequency where the polarisation effect is minimal.

![Figure 7-11: The conductivity of the clean LDPE and the soaked LDPE.](image-url)
7.5 Chapter Summary

The dielectric spectroscopy measurements have been carried out on the liquid byproducts as well as the byproduct soaked LDPE. The value of permittivity and dielectric loss of the byproducts soaked LDPE films do not really resemble the measured values of the parent byproducts. It may possible to relate the small percentage weight of acetophenone in LDPE to its small effect on the dielectric constant and loss of LDPE. However, this relation is not true for α-methylstyrene sample. Thus, it is proposed that these differences may due to different polarisation effect of the byproducts in LDPE. In the future, it is worth to check the effects of several byproducts in one LDPE samples on dielectric spectroscopy.

Acetophenone soaked LDPE has the highest permittivity value followed by α-methylstyrene and cumyl alcohol soaked LDPE. Among the samples, only cumyl alcohol soaked LDPE exhibit the MWS polarisation and space charge polarisation effect at low frequencies. As a result, the dielectric loss and conductivity value of this sample is much higher than the rest of the measured samples.

On the other hand, the acetophenone soaked LDPE and α-methylstyrene soaked LDPE have very similar dielectric loss values to the clean LDPE. Last but not least, the Cole-Cole plots show that the soaked samples do not exhibit the Debye relaxation process.
Chapter 8  Conduction Current Measurement

8.1 Introduction

Generally, polymer has very low conductivity. Despite its high insulating properties, polymer could not stop some tiny flow of charges injected from the electrode at high voltage. Conductivity measurement has been a commercial interest because of the relation of cable aging or failure due to the local damage induced by charge flow and/or recombination. Due to the amount of traps in polymer, it is said that the polymer has a trap-controlled conductivity [105].

Conduction current consist of three components namely the leakage current, charging current and absorption current. The later two currents rise dramatically during the initial charging period but gradually reduce to a smaller value and eventually approaching zero. The charging current or also known as capacitive current becomes very high due to the electron rush into the negative electrode and be drawn from the positive electrode. Absorption current is due to the gradual change of polarisation of polymer (orientation of the polar side of the polymer chain by the electric field) [106] and accumulated charges under the influence of the electric field stress. This transient current is followed by a steady current which is the leakage.
8.2 Procedures

Samples for conduction current measurement have been coated with gold on both sides before they undergo the soaking process. 2 cm mask diameter was used to match the electrode size. Samples were stressed at 3, 5 and 8 kV which are equivalent to 16.7kV/mm, 27.8kV/mm and 44.4kV/mm respectively. The measurement took 3600 seconds (60 minutes) to be completed and readings were taken for every 20 seconds. Keithley 6487 picoammeter has been used to measure the current. The electrodes were placed in an oven to maintain the temperature throughout the experiment. However, the temperature is limited to room temperature due to the nature of the byproducts that will evaporate at high temperature. It is crucial to maintain the same amount of byproducts in the sample throughout the experiment. The circuit connection is illustrated in Figure 8-1.

180µm samples were used for the soaked sample measurement. The clean LDPE measurement however requires a thinner sample and larger electrode due to very small current flow in the sample. Here, 50µm LDPE film and 5cm electrode diameter are used. In addition, higher electric fields need to be applied for the measurement to obtain a higher signal to noise ratio.

![Circuit connection for conduction current measurement.](image)

Figure 8-1: Circuit connection for conduction current measurement.
8.3 Conduction Current Result

![Figure 8-2](image)

**Figure 8-2**: The current density-time characteristic of clean LDPE at 60kV/mm and 120kV/mm at room temperature.

To obtain the conduction current value without being deceived by the charging and absorption current, it is necessary to measure the current after a specified period of time. In this research, the conduction current reading is taken at 3500 seconds. Figure 8-2 shows the current density profile of the clean LDPE sample during charging. Although the field applied is very big, the current flow in the sample is very small. After 1 hour charging at 60kV/mm, the current density becomes as low as \(\approx 1 \times 10^{-3} \text{pA/m}^2\) and \(\approx 2.8 \times 10^{-3} \text{pA/m}^2\) at 120kV/mm. One can imagine that the current in clean LDPE at lower electric field is so much lower.

Comparing the current density of the soaked sample to that of the clean LDPE, it is quite obvious that the byproducts enhance the conduction current value in at least one order of magnitude. These values are presented in Figure 8-3 to Figure 8-5. From the results, cumyl alcohol causes the biggest jump to the current value followed by \(\alpha\)-methylstyrene and then acetophenone.
Charged at 3kV, the current density in acetophenone soaked LDPE initially started at $\approx 5 \times 10^{-8}$ A/m$^2$ and this decreased to $\approx 3.5 \times 10^{-8}$ A/m$^2$ after 60 minutes charging. The reduction of the current is very little. The current density of acetophenone soaked LDPE after charged at 5kV and 8kV are $8 \times 10^{-8}$ A/m$^2$ and $2.5 \times 10^{-7}$ A/m$^2$ respectively. At higher charging voltage, the current reduction over the charging period becomes bigger.

Figure 8-3: The current-time characteristic of acetophenone soaked LDPE at 3, 5 and 8kV at room temperature.

Figure 8-4: The current-time characteristic of $\alpha$-methylstyrene soaked LDPE at 3, 5 and 8kV at room temperature.
Figure 8-5: The current-time characteristic of cumyl alcohol soaked LDPE at 3, 5 and 8kV at room temperature.

For $\alpha$-methylstyrene soaked LDPE, similar current decay is seen in the samples charged at different voltages. Cumyl alcohol soaked LDPE however shows a different result. Similar decay is seen in the current from samples charged at 3 and 5kV. Meanwhile sample that charged at 8kV has faster current decay. Table 8-1 shows the current density value of all samples at 3500s.

Table 8-1: Current density of samples at 3500s

<table>
<thead>
<tr>
<th>Sample</th>
<th>Charging Voltage</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3kV</td>
</tr>
<tr>
<td>Acetophenone soaked LDPE</td>
<td>3.5 x 10^{-8} A/m^2</td>
</tr>
<tr>
<td>$\alpha$-methylstyrene</td>
<td>1.8 x 10^{-7} A/m^2</td>
</tr>
<tr>
<td>soaked LDPE</td>
<td></td>
</tr>
<tr>
<td>Cumyl Alcohol</td>
<td>3.5 x 10^{-6} A/m^2</td>
</tr>
<tr>
<td>soaked LDPE</td>
<td></td>
</tr>
</tbody>
</table>
8.4 The Correlation of DC Conductivity with space charge

This observation may be explained by the heterocharges that formed in cumyl alcohol soaked sample. Heterocharges enhance the electric field at the metal-insulator interface causing more charges to drift into the dielectric. Cumyl alcohol allows more charges pass through the LDPE leading to the increment of the current. Figure 6-12 in Chapter 6, demonstrates the fast growth of positive and negative charges both in bulk and near the electrodes.

The decaying transient current is proved to be bulk origin [105]. The accumulation of heterocharges reduce the electric field acting in the bulk and hence it explain the large current decay after the high initial current in cumyl alcohol soaked LDPE as seen in Figure 8-5.

Despite having the highest conductivity value, acetophenone (1.8x 10^-6 S/m) becomes less conductive compared to cumyl alcohol (1.4 x10^-7 S/m) as it presence in LDPE [101]. This discrepancy might due to the small amount of acetophenone diffused in soaked LDPE compared to the other byproducts. Comparing a clean LDPE to the acetophenone as well as α-methylstyrene soaked LDPE, more charges injected from the electrode forming homocharge in the samples, and hence the diffusion of charges through the bulk may also increase. This is why bigger current was found in the soaked LDPE.

Conductivity (σ) values of the samples are calculated from the result of conduction current via equation:

\[ \sigma = J/E \]  \hspace{1cm} \text{Equation 8-1}

and the values are plotted against electric field in Figure 8-6. The conductivity values for acetophenone and α-methylstyrene soaked LDPE increase linearly with the electric field. Meanwhile conductivity of cumyl alcohol soaked LDPE shows a different pattern in the relationship with electric field. Initially, the conductivity value increase dramatically but as the electric field increases, the slope of the graph start to decrease. It can be observed that cumyl alcohol amplifies the current that can pass through the dielectric with only small change of electric field.
As presented in Table 8-2, each byproduct-soaked-LDPE has different amount of byproducts. This is due to the different diffusion rate of the byproducts into LDPE. R. Minami et.al reported in [107] that the conductivity increases with the soaking time, i.e., more byproducts chemical in the sample film. Hence, it is only justified to measure the influence of each byproduct on conductivity value relatively to the amount of these byproducts in the samples. Table 8-2 presented the conductivity gain by each percent of byproducts in LDPE. These values indicate the conductivity increment (in percent) caused by each percent of byproducts in the soaked sample and they are dependent on the electric field. These values show that with only small amount of crosslinking byproducts left in the insulator, particularly cumyl alcohol, the conductivity of the insulator could be magnified to a very big value that may damage the insulation.

Table 8-2: Conductivity Gain by each percent of byproduct in LDPE

<table>
<thead>
<tr>
<th>Sample</th>
<th>Electric Field (kV/mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>16.70</td>
</tr>
<tr>
<td>Acetophenone soaked LDPE</td>
<td>28.5%</td>
</tr>
<tr>
<td>α-methylstyrene soaked LDPE</td>
<td>93.8%</td>
</tr>
<tr>
<td>Cumyl Alcohol soaked LDPE</td>
<td>4957.26%</td>
</tr>
</tbody>
</table>
The effects of polar groups on conduction in polyethylene have been reported before [108]. Although the work done was on acetoxy group, similar effect could be observed from the polar groups that exist in the crosslinking byproducts. With the formation of heterocharges in cumyl alcohol soaked LDPE as well as a massive impact on the conductivity value, cumyl alcohol has a special feature that does not exist in acetophenone and α-methylstyrene. It is necessary to associate the unique characteristic with the existence of the hydroxyl group that exists in cumyl alcohol.

In addition, it was found that the conductivity values are thickness dependent. Extra measurements were conducted on 50μm byproducts soaked LDPE samples. Figure 8-7 plots the conductivity values for the samples at different electric filed. Result shown in figure 8-6 also included for comparison. Thinner sample has lower conductivity. Notice the 50μm cumyl alcohol soaked LDPE at field around 20 and 40kV/mm. The conductivity values are two orders smaller than that of 180μm samples around the same electric field.

Comparing the conductivity values for the 50μm samples, cumyl alcohol still acquires the highest conductivity value. Acetophenone and α-methylstyrene soaked LDPE have almost similar conductivity value, two order lower than that of cumyl alcohol soaked LDPE. Last but not least is the clean LDPE. The conductivity value of LDPE is as low as $10^{-17}$. It is seen from Chapter 6 before, although the total charge that injected into clean LDPE is comparable with the total charge in byproducts, most of the charges are trapped in deep traps. As a result, very little charge contributes to conduction in clean LDPE.
Figure 8-7: The conductivity of the 50μm and 180μm clean and soaked samples at different electric field.

8.5 Carriers Mobility

The power law dependency of current density to electric field is often found in many research of electrical conduction of dielectric polymer. The power law is given by [109]:

\[ J = aE^n \]  \hspace{1cm} \text{Equation 8-2}

And can be written as

\[ \ln J = \ln a + n \ln E \]  \hspace{1cm} \text{Equation 8-3}

This gives a straight line with slope n of ln J vs. ln E plot. Different n value indicates different conduction mechanism. In the case of ohmic conduction, where n = 1, it is usually as the result of electric field independency of carrier concentration and mobility [110]. In the case of space-charge-limited-conduction (SCLC), Equation 3-18 is referred, which give the value of n ≈ 2. For a combination effect of non-ohmic contact, e.g. Schottky barrier with the effect of space charge, the values of lie in the range of 2 to 4 [111]. Montanari in his work [110,112] uses this relation to distinguish the electrical degradation threshold of PE.
J-E characteristic of the soaked samples are presented in Figure 8-8. The J-E curves of three byproducts-soaked samples are almost similar. The slope values for acetophenone, α-methylstyrene and cumyl alcohol soaked LDPE are 1.68, 1.81, and 2.19 respectively. According to [113] current distributions with such $n$ values could be described by the SCLC theory. It may however be expressed by the Schottky laws as well as the ionic conduction. However SCLC plot can effectively display the space charge accumulation process on the basis of charging current.

Mobility could be obtained from SCLC theory from equation 3-19, which is $\mu = \frac{8Jd^3}{9\eta^2\varepsilon_0\varepsilon_r}$. The relative permittivity of the samples is taken from the measured value presented in Chapter 7. Since permittivity value for cumyl alcohol soaked sample increase at lower field due to the ionic conduction, the permittivity value used for the calculation is taken at 10Hz before the ionic conduction just started to affect the permittivity value. Figure 8-8 plots the charge mobility values against electric field.
It must be pointed out, regarded Figure 8-9, the mobility values calculated by the derivation from SCLC theory refer properly to the mobility of free charge carrier where $\theta = \frac{n_c}{n_t}$ is assumed to be equal to one. Hence, in a dielectric with traps such as the samples measured, the mobility values should be expected to be lower than the plotted values.

### 8.6 Chapter Summary

The conductivity of the clean LDPE is very low and hence the conduction current measurement can only be conducted at a very high electric field. The conductivity of the byproducts soaked LDPE is so much higher in at least 8 times higher than the conductivity of the clean LDPE.

This finding is different from the total accumulated charges calculated from the space charge measurement. From Figure 6-16, it is clear that the charge in LDPE is considered in the equivalent range of that of the byproducts soaked sample. Thus, we can say that the ratio of the free charge carrier to the trapped charges ($n_c/n_t$) is very small and hence very little charges contribute to conduction.
Cumyl alcohol soaked LDPE has the highest conductivity which is 2 times bigger than α-methylstyrene soaked LDPE and 3 times bigger than the acetophenone soaked LDPE. This observation could be linked to the hydroxyl group that appear in cumyl alcohol as well as the heterocharge formation in the cumyl alcohol soaked LDPE.
Chapter 9  AC Breakdown

9.1 Introduction

Breakdown in dielectric insulation occurs when the insulation can no longer insulate and forced to conduct electricity. Dielectric breakdown in polymer is very complicated due to its complex structure. Breakdown mechanism is always associated with the internal field. In term of mechanical breakdown theory, internal field causes electromechanical force to be sufficient enough to break the material [114]. Meanwhile for thermal theory, internal field is related to electrical conductivity which increases the temperature up to the melting point of the material [43]. Last but not least, in the electronics breakdown theory, breakdown happens when the internal field reaches its critical value. The summary of breakdown process that occurs in solid dielectric is presented in Figure 9-1.

**Electronic breakdown Theories**
- intrinsic breakdown
- Electronic thermal breakdown
- Electron avalanche breakdown field emission breakdown
- Space charge-enhance critical field breakdown

**Thermal breakdown Theories**
- Steady state thermal breakdown
- Impulse thermal breakdown

**Mechanical Breakdown Theories**
- Electromechanical breakdown

Figure 9-1: Electrical breakdown theories of solid dielectrics.
Electronics breakdown theory can be classified into three mechanisms which are intrinsic, avalanche and space charge-enhanced critical field. During the application of very high electric field, some electrons from the valance band cross the band gap into the conduction band. As more and more electrons become available for conduction, conduction current across the insulation becomes huge and this phenomenon is called intrinsic breakdown. Avalanche theory was initially developed for a dielectric that is considered as a polar crystal. Avalanche happens when an electron from cathode with sufficient energy collide with another atom/molecule producing more electron and hole. If these electrons produce further two electrons, an avalanche of $n$ electron will be produced in $n$ generation. The process will be repeated and propagate across the insulation thickness.

![Avalanche Process Diagram](image)

**Figure 9-2: Avalanche process**

Electrons injection in voids is controlled by the space charge that accumulated on the surface of void. Electrons (injected from the electrode or from the polymer) get trapped on the void surface as they travel to the opposite electrode. Negative electron affinity of polyethylene promotes surface state of the electron under the bottom of conduction band. The electrons that travel in polymer will get trapped on the surface of void and the process continues until the Coulombic repulsion between them is high enough to increase their energy level to de-trap by Schottky effect or thermal activation. In the void, the energy
barrier is much lower than in the PE. Electrons injected into the void are accelerated by applied electric field and if the field is high enough, it could cause avalanche in the void.

It is agreed that breakdown strength is defined as the voltage at which beyond this value, the dielectric can no longer maintain its ability to insulate. This value however is not an absolute number. This value varies depending on several factors that may influence the breakdown mechanism. These factors need to be reviewed so the limitation of dielectric strength value could be understood better. Some of the factors that have a big influence to breakdown strength values are;

- Dielectric thickness
- Temperature
- Electrode shape and size
- Rate of voltage increase

Other than the obvious reason that thinner sample will have less chance for defects to exist, the dependency of breakdown strength on thickness could also be explained by Maxwell stress [115,116]. Breakdown strength of dielectrics decreases with its thickness. Temperature control is important as the dielectric strength is related to the temperature of the specimen at the time of breakdown. The temperature dependent of dc breakdown field is usually decreased with temperature however this relation is non-linear. The electrode shape and size will determine the electric field distribution on the test sample. This factor is very important for small sample testing. Rogowski type electrode and ball bearing electrode could provide a uniform stress gradient which is important in order to obtain a reliable result. However, needle tests are also performed to investigate the breakdown phenomenon under divergent field. Rate of voltage increase may also affect the breakdown strength value. As the breakdown is performed at a constant ramp-rate (in which the electric field increases from zero at a predetermined rate), the breakdown value obtained will normally be greater than the value that obtained in step-rise test procedure (field increase at every set rise time). The impulse breakdown strength is reported to increase as the ramp rate is increase[117]. Hence, it is important to state the conditions together with the breakdown strength value since different condition may give different breakdown value.
9.2 Weibull Distribution Of AC Breakdown

Two-parameter Weibull Distribution is most commonly used for characterizing the time of failure for solid insulation. This distribution has been accepted as IEEE 930/IEC 62539 standard. The cumulative probability of failure for two parameter Weibull distribution can be expressed by:

\[ P_F(t) = 1 - \exp \left( -\frac{t}{\alpha} \right)^\beta \]  \hspace{1cm} \text{Equation 9-1}

where \( t \) is the measured variable (\( t \) or \( E \)), \( \alpha \) (eta) is the scale parameter, and \( \beta \) (beta) represent the shape parameter. The higher the value of \( \beta \), the narrower the spread of times (or fields) to failure. From another perspective, the value of \( \beta \) shows the reliability of such material to insulate the HV cable before the material breakdown. The probability density function (PDF) of equation 9-1 is given by:

\[ g(t) = \frac{dP_F(t)}{dt} = \beta \alpha^{-\beta} t^{\beta-1} \exp \left\{ \left( \frac{t}{\alpha} \right)^\beta \right\} \] \hspace{1cm} \text{Equation 9-2}

Figure 9-3(a) and (b) show the plot of cumulative probability of failure and probability density function respectively for the cases of \( \beta=0.5 \), 1.0, 1.5, 2.0 and \( \beta=3.0 \). At a constant value of \( \alpha \), which in this case \( \alpha=1 \), various value of \( \beta \) will result to different slope.

In order to calculate the cumulative probability of failure, it is necessary first to rank the breakdowns in order of time (or field) and then estimate the true probability of failure for each point from \( i=1 \) to \( i=n \) for \( n \) samples. This probability of failure is best obtained at the median, in which the estimation proposed by Bernard and Bos-Levenbach [118] given by:

\[ F(i,n) = \frac{i-0.3}{n+0.4} \] \hspace{1cm} \text{Equation 9-3}
Figure 9-3: (a) The cumulative probability of failure and (b) probability density function for the cases of $\beta=0.5, 1.0, 1.5, 2.0$ and $\beta=3.0$. [43]

Obtaining Weibull parameters is possible by rearranging equation 9-1 as a linear function;

$$\log_{10}(-\log_e[1-P_F(t)]) = \beta \log_{10}(t) - \beta \log_{10}(\alpha)$$

Equation 9-4
Plotting $\log_{10}\left(-\log_e[1 - P_F(t)]\right)$ versus $\log_{10}(\alpha)$ will result in a straight line of slope $\beta$. This linear regression however will be bias at the extreme of distribution. Therefore, maximum likelihood (ML) estimation is used to approximate the values of $\alpha$ and $\beta$ which is the product of PDF at each data point.

$$L(p) = g(x_1\alpha, \beta) \times g(x_2\alpha, \beta) \times \ldots \times g(x_n\alpha, \beta) \quad \text{Equation 9-5}$$

In practice, $\log L(p)$ is used rather than $L(p)$ (likelihood function) and the relevant equation of two-parameter Weibull distribution becomes;

$$\frac{1}{\beta_{ML}} = \frac{\sum_{i=1}^{n} \left(x_i^{\beta_{ML}}\right) \ln x_i}{\sum_{i=1}^{n} x_i^{\beta_{ML}}} - \frac{1}{n} \sum_{i=1}^{n} \ln x_i \quad \text{Equation 9-6}$$

$$\alpha_{ML} = \left[\frac{1}{n} \sum_{i=1}^{n} x_i^{\beta_{ML}}\right]^{\frac{1}{\beta_{ML}}} \quad \text{Equation 9-7}$$

where $\beta_{ML}$ and $\alpha_{ML}$ are the shape and scale parameters obtained from maximum likelihood estimation. Nevertheless, this estimation is known to be biased to the value of $\beta$ for small $n$ and the value $\alpha$ become more bias for small $\beta$.

In this study, Weibull software supplied by ReliaSoft® was used to perform the Weibull distribution.

### 9.1 Experiment Procedures

LDPE films of 50µm thick were soaked into acetophenone and $\alpha$-methylstyrene at room temperature meanwhile into cumyl alcohol at 80°C. The samples were used to perform the breakdown strength ($E_b$) of the LDPE with the byproducts chemical reside in the sample. The AC breakdown measurements were conducted at a ramp rate of 50V/s at room temperature. Ball-bearing electrodes were used. Weibull distribution is used to analyse the AC breakdown result. Figure 9-4 shows the equipment setting in the breakdown rig.
9.2 AC breakdown test result

Figure 9-5 shows the plot of probability of failure, \( F(i, n) \) versus the breakdown strength for all samples. The Weibull parameters, shape and scale parameters are estimated from these plots by using special software. The dotted lines drawn in the figures are the estimates value of 90% confidence intervals for each measurement.

Looking at the \( \alpha \) value, which is the scale parameter of the breakdown strength, these values shows that some of the byproducts increase the AC breakdown strength. Acetophenone and cumyl alcohol increase the breakdown strength for about 10kV/mm. This result is in agreement with result reported in [120,121]. Meanwhile it is shown here that \( \alpha \)-methylstyrene reduces the scale parameter value for about 10kV/mm lower than the clean LDPE. These values are presented in Table 9-1. The results obtained here is in agreement with [121] due to resistance of the byproducts impregnated samples to partial discharge. Nevertheless, the breakdown increment in the samples reported in the paper is much higher than the values obtained in our experiment. Overall, the breakdown strengths
Figure 9-5: The Weibull Breakdown probability plot of a) Clean LDPE, b) acetophenone soaked LDPE, c) α-methylstyrene soaked LDPE and d) cumyl alcohol soaked LDPE
of all samples fall in the same range of values. Thus the byproducts have no effect on the ac breakdown strength of LDPE and similar observation seen in [28].

The exact breakdown mechanisms of the samples are difficult to be confirmed since it may involve more than one mechanism in the breakdown process. However it is worth to mention here that author like Fukuma reported breakdown measurement in conjunction with the space charge measurement [122]. Here, the role of space charge is observed as the space charge dynamics in the sample is captured just before breakdown happens. Nevertheless, most of the works that link space charge and breakdown strength are mainly related to direct field where the effect of space charge is much more severe. The results in Chapter 6 have shown the different impact of ac and dc field on the accumulation of space charge. Hence, it is believed that the effect of byproducts on dc breakdown strength will be more severe. With the breakdown process governed by the maximum electric field in the sample, space charge effect on dc field will play the utmost role to cause breakdown to happen.

Table 9-1: Weibull analysis of breakdown strength test result

<table>
<thead>
<tr>
<th>Sample</th>
<th>% added crosslinking byproducts (%)</th>
<th>$\eta\alpha$ (kV/mm)</th>
<th>Beta, $\beta$</th>
<th>Range of $E_b$ (kV/mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDPE</td>
<td>0</td>
<td>192.2 ± 6.2</td>
<td>11.9</td>
<td>148-226</td>
</tr>
<tr>
<td>LDPE + Acetophenone</td>
<td>2.42</td>
<td>198.8 ± 5.2</td>
<td>14.6</td>
<td>165-220</td>
</tr>
<tr>
<td>LDPE + $\alpha$-Methylstyrene</td>
<td>13.20</td>
<td>183.5 ± 5.6</td>
<td>11.3</td>
<td>127-216</td>
</tr>
<tr>
<td>LDPE + Cumyl Alcohol</td>
<td>14.36</td>
<td>195.1 ± 6.8</td>
<td>11.0</td>
<td>171-231</td>
</tr>
</tbody>
</table>
9.3 Chapter Summary

AC breakdown test has been performed on the clean LDPE and crosslinking byproducts soaked LDPE. Weibull distribution is used to give the statistical features of the breakdown phenomena of the byproducts soaked LDPE.

By referring to the α values from the distribution, acetophenone and cumyl alcohol slightly increase the ac breakdown strength of LDPE. On contrary, α-methylstyrene slightly reduces the breakdown strength value.

In term of β, this value is very similar between the soaked sample and the clean LDPE, except for the acetophenone soaked LDPE which shows a higher β value. The overall ac breakdown characteristic of the soaked samples are very much similar to that of the clean LDPE. The breakdown values for each sample fall in the same range. Thus the byproducts have no effect on the ac breakdown strength of LDPE.

The small effect of crosslinking byproducts on ac breakdown is also observed in ac space charge measurement. In the latter, although the byproducts increase the total charge accumulated in the LDPE sample, the increment is very small compared to dc case. Hence, it is believe that the crosslinking byproducts may also have greater effect on dc breakdown strength due to high charge mobility in the soaked sample in dc condition as seen in Chapter 6 and Chapter 8. This hypothesis is also based on the report on the influence of the byproducts towards impulse breakdown [123].
Chapter 10  Conclusions and Future Works

10.1 Conclusions

This thesis provides a report on the research done on the crosslinking byproducts that remain in XLPE after the crosslinking process. Three main byproducts were investigated here which are acetophenone, α-methylstyrene and cumyl alcohol. To obtain a single product in each measured sample, LDPE was used as the base material for the soaking process. Through this research, consequences of crosslinking byproducts towards space charge accumulation across the polymeric insulation and their effect towards the charge conduction, permittivity and ac breakdown strength are realised. Some conclusions can be made.

1. Space Charge Accumulation in byproducts soaked LDPE

The presence of byproducts in XLPE cable is mostly associated with space charge formation in the cable insulator. Here, the individual effects of the byproducts on the formation of space charge were investigated. The presence of the byproducts changes the space charge pattern in the samples. Homocharges are seen in the vicinity of the electrodes for acetophenone and α-methylstyrene soaked samples. Although the clean sample also possesses similar homocharge, the charge pattern is different. Acetophenone ease the movement of negative charges in LDPE causing more negative charge accumulation in the sample bulk. On contrary, α-methylstyrene causes more positive charge build up in the sample bulk. Dissimilar to the former two byproducts, cumyl alcohol introduces
heterocharges near both electrodes. The formation of heterocharges is believed due to the hydroxyl group that presence in cumyl alcohol.

We were also able to calculate the total charge in each sample and compared with the total charge in clean LDPE. Cumyl alcohol has the highest total charge followed by clean LDPE, acetophenone and α-methylstyrene. However at higher field, acetophenone introduces more charges compare to clean LDPE. α-methylstyrene on the other hand consistently reduces the amount of total charge introduced into the insulator. This characteristic is also observed in samples with 2 byproducts, where any samples with α-methylstyrene have low total charge.

The charge decay process reveals the ability of the byproducts to speed up the charge reduction in samples. 2 different charge decay rates leads us to 2 different traps depth in polymer. Hence, the influence of the byproducts on each population of traps could be determined. It was found that the byproducts increase the number of shallow traps in the samples and reduce the number of charges that are trapped in deep traps. α-methylstyrene soaked LDPE have the fastest charge decay followed by acetophenone and then cumyl alcohol.

An improved PEA system enables us to measure the space charge property of the samples in ac condition. It was found that the byproducts once again increase the total charge in the sample although the amount is so much smaller than that in dc. This time however, the charges that trapped in the samples are identified as the charges in deep traps due to their very slow charge decay. These findings together with the results obtained in dc condition lead us to another conclusion. The shallow traps that exist in crosslinking byproducts increase the number of charges trapped in the sample. As they move further into the bulk, the probability of the charges to be trapped into deep traps is higher and hence we see more charges in deep traps in the byproducts soaked sample. This conclusion is significant in determining the trapping probability of deep traps in any model. From this research, it is found that this value depends on the concentration of shallow traps in the dielectrics.
2. Dielectric Spectroscopy Measurement

In applying a material as the cable insulation, the dielectric constants (permittivity) as well as the dielectric loss tangents are two important parameters that need to be considered. As we measure the byproducts soaked LDPE, it was found that acetophenone give a small increase of the dielectric constant value of LDPE. The other byproducts do not affect much on this value.

Cumyl alcohol soaked LDPE shows a gradual increase of the permittivity value as the spectrum shifted to lower frequency. The dielectric loss value for this sample is also very high particularly at lower frequency. This result may be due to the MWS polarisation effect and electrode polarisation effect as well as the ionic conduction in the sample. This phenomenon however was not observed in acetophenone and α-methylstyrene soaked LDPE. Hence it is believe that the hydroxyl group in cumyl alcohol plays the main role in causing such behaviour.

The conductivity values were calculated using the imaginary part of permittivity. As expected, cumyl alcohol has the highest conductivity value.

3. DC Conductivity measurement

Conductivity values of the sample are measured at room temperature. DC conductivity of clean LDPE is very low. The influence of electric field on conductivity is observed. Cumyl alcohol has the biggest effect on the conductivity of the soaked LDPE followed by α-methylstyrene and acetophenone. This result may be explained by the formation of heterocharges in the vicinity of electrode which observed in space charge measurement, enhancing the electric field near the metal-dielectric interface and lead to the increment of the current value. Cumyl alcohol is proven to be more conductive in LDPE compared to the other byproducts even at low electric field.

Through the experiment, we were able to calculate the mobility values of the carriers in the samples by using SCLC theory.
4. Dielectric breakdown on AC field

AC breakdown test was conducted using ball-bearing electrodes which were placed in silicon oil. The result reveals that the breakdown strength of the byproduct soaked LDPE is almost similar to that of the clean LDPE. Although acetophenone and cumyl alcohol slightly increase the α value, also α-methylstyrene slightly reduces the α value, the breakdown strength for each samples fall in the same range.

In other words, the crosslinking byproducts have little effect on ac breakdown strength. By relating the ac breakdown strength with ac space charge measurement result, one could agree that the little difference on space charge formed in byproducts soaked LDPE and LDPE samples that were stressed in ac field may lead to small difference on ac breakdown strength among the samples.

Since the effect of the crosslinking byproducts on space charge formation at dc field is more severe, we believe that the byproducts will also influence the dc breakdown strength value of LDPE.

5. Byproducts

After comparing the influence of acetophenone, α-methylstyrene and cumyl alcohol on the electrical properties that discussed above, it could be concluded that the total amount of byproducts in each sample is not one of the factors that contribute to the different observation on each tested sample. We believe that these differences are related to the polar group that exist in each byproduct molecule. The hydroxyl group in cumyl alcohol may cause the heterocharge in the stressed sample due to the ionisation process in the sample. The O-H bond may also causes ionic conduction as well as MWS and electrode polarisation during the dielectric spectroscopy measurement. As a result, cumyl alcohol has the biggest dielectric loss compared to the other samples.

The carbonyl group (C=O) in acetophenone and conjugate double bond (C=C) in α-methylstyrene have lead to the formation of homocharges in stressed samples. They however are observed to assist the movement of different type of charge carriers. Both of
these byproducts do not change the dielectric loss of LDPE and their charge mobility values are only one order of magnitude differ.

The three crosslinking byproducts have very small effect on electrical properties of LDPE during ac condition. This is observed from the ac space charge measurement as well as the ac breakdown measurement. In term of charge decay, we could conclude that the byproducts particularly α-methylstyrene leave a positive effect on the process. With the fastest charge decay, α-methylstyrene could remove all of the charges in the sample much faster than other samples. The byproducts however have affected the total amount and distribution of space charge as well as the charge dynamics of the sample. The effects of the byproducts become much severe at higher electric field. With very high DC conductivity, the byproducts have proved to be the drawback to the insulation material. Cumyl alcohol has shown its ability to introduce more charges into the LDPE with the formation of heterocharge. It also has the highest dielectric loss and dc conductivity which leads to the highest charge mobility value. Based on these results, we could conclude that cumyl alcohol is the most dangerous crosslinking byproduct compared to the others.

10.2 Future Work

Several areas of this investigation have the potential to be continued by further research and experimentation, some of which is detailed below.

1. Much knowledge could be gained by studying the space charge profile of the polymer. One big advantage that could contribute to this research is by measuring the conduction current simultaneously with the space charge profile. By obtaining the current values, the carrier mobility could be calculated without applying any ambiguous assumption to simplify the equation. With this, the carriers’ mobility value could be attained accurately.

2. In this thesis, we were trying to link the space charge profile to the breakdown result that obtained from a different system. This comparison however will be much more accurate if the breakdown test could be conducted on the PEA system.
By having this system, we will be able to monitor the space charge and electric field in the sample just before breakdown happens. DC breakdown measurement of the byproduct soaked LDPE could also be done to verify the contribution of the byproduct.

3. It is also an interest of this research to relate the electrical properties that are measured to the chemical structure of the byproducts such as the molecule size, since they are both related to each other. The trapping characteristic might be influenced by the chemical groups in the byproducts. One thing that could be done is to use various chemicals with similar group to confirm their effects on electrical properties.

4. The influence of space charge on the electrical performance of a material depends on several factors such as the amount and distribution of space charge and charge dynamics. These are affected by the magnitude and duration of the applied electric field, temperature and electrode material. The works that reported in this thesis present the short-term effects of the byproducts. These results may be different from the samples that are stressed for a longer duration. Thus, a long-term effect of the byproducts should also be studied to reveal the hazardousness of these byproducts on the electrical properties. By doing this, along with selecting the right experiment variables, the results obtained will be closer to the actual condition in the real insulation cable.
References


W. Choo, G. Chen. *Electric field determination in DC polymeric power cable in the presence of space charge and temperature gradient under dc conditions*, in 2007, Beijing, China, pp. 321-324.


Appendix: AC Volt On result

Charge Density of 180µm clean LDPE during Volt on condition after 4 hours stress at 28 kV/mm ac.
Charge Density of 180µm acetophenone soaked LDPE during Volt on condition after 4 hours stress at 28 kV/mm ac.
Charge Density of 180µm α-methylstyrene soaked LDPE duringVolt on condition after 4 hours stress at 28 kV/mm ac.
Charge Density of 180µm cumyl alcohol soaked LDPE during Volt on condition after 4 hours stress at 28 kV/mm ac.