

Article

Measurement of the Relative Free Radical Content of Insulating Oils of Petroleum Origin

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Abstract: Oil/paper insulation degradation in transformers involves chemical and physical changes in the materials. Some of the chemical reactions involve very reactive intermediates called free radicals. Free radicals play a major role in a wide variety of ageing processes. The detection of these reactive species in oil may, in principle, provide useful information for monitoring oil degradation. This manuscript details a laboratory technique, which determines the relative content of free radicals in insulating oils of petroleum origin by a spectrophotometric method. Free radicals may be formed in oils under operating or test conditions. The procedure enables the determination of the relative concentration of free radicals, which can act as the precursors of decay products such as charge carriers, oxidized molecules, as well as polymerization products. The technique involves using a reactive free radical reagent, 2,2-diphenyl-1-picrylhydrazyl (DPPH), added to oil to assess free radical concentration. This method is applicable to new, reclaimed, or used oils as well as naturally or artificially oxidized oil (the cause of aging can be chemical, physical, or electrical). In this contribution, free radicals were assessed following electrical discharge application in oil.

Keywords: free radicals; insulating oil; reaction kinetics

1. Introduction

The condition of power transformer insulation is paramount since its reliability is important to society and power utilities along with the techno-economic growth of any country. For power transformers to operate normally it is important to isolate live parts from each other and the ground, current not leaking from one path to another through materials not intended to be a conducting path. Insulation includes all three forms of matter, sometimes with a simple shape but often involved a combination of forms, such as the solid/liquid or solid/gas. Heat dissipation can be achieved by circulating certain fluids (liquids/gases), which also ensure electrical insulation of energized conductors. The insulating-fluids market is therefore likely to be dominated by liquids [1]. Petroleum based oils, the so-called mineral oils, are used as the sole insulation material only in regions where, by design, electrical stresses are relatively low. The functions of oil are to strengthen the dielectric properties of solid insulation, to electrically insulate active parts from grounded ones, to dissipate heat from the core and coils to the radiators, and to quench eventual arcs [2].

The insulating oils used in large power transformers may suffer from thermal, electrical and/or environmental stresses, even under service conditions [1]. This will, in turn, cause and accelerate the deterioration of the insulation, posing a potential threat to the equipment safety. Aging is a phenomenon resulting in slow and irreversible changes in material properties [3–6]. The process implies the decomposition of vulnerable hydrocarbons resulting in a modification of the macromolecular structure of the liquid and solid materials. In general, transformer oil loses its stability gradually and becomes decomposed and oxidized [3–9].

Electrical stress together with heat and moisture, in the presence of oxygen, oxidises the oil producing free radicals, acids, and sludge that are deleterious to the transformer [6,10,11]. Free radicals may be generated by thermal decomposition, electrical discharge, electrolysis at an electrode, mechanical damage, chemical reactions, and high energy radiation and are thus implicated in the ageing process [12,13]. John at The Nineteen Symposium on Electrical Insulation underscored the paramount importance of free radicals in the physical organic chemistry of mineral insulating oils [14]. These by-products are deleterious to the transformer and catalyze further oxidation of the oil. The electric stress accelerates ageing, possibly by increasing the precipitation of acid products of oil degradation onto paper surfaces. Chemically speaking, the acid build-up will worsen the insulating paper tensile strength. Aggressive decay products being absorbed by the solid insulation attack the cellulose fibers too. Sludge produced may stick onto the large surface of transformer boards stopping heat being dissipated which results in overheating of insulation and accelerates degradation processes. The cracking process of cellulose (depolymerisation by a succession of chemical reactions) causes chain scissions, the release of gases and water into the surrounding oil and some large molecules such as furfurals [15]. In composite oil-impregnated dielectrics for high voltage insulation, oil is usually the weaker component of the system, both in dielectric strength and in reaction to environmental stress. Knowledge of the stability of insulating oils under electrical stress is of upmost importance to both electrical-equipment designers and operating engineers. Free radicals are very reactive and can adversely affect the chemical, physical, and dielectric properties of the insulating liquid [12,13]. Therefore, the detection of free radicals in oil may provide useful information on the inception of oil/paper system degradation. The method proposed allows the condition of insulating oil to be assessed through investigation of free radicals.

An attempt to develop on-line diagnostic tools and field monitoring techniques by Electron spin resonance (ESR) was explored previously [12]. From the obtained results, the authors concluded that the ESR technique was not suitable for on-line monitoring of the condition of oil/paper insulation of high voltage equipment. In this manuscript, a sensitive spectrophotometric method is used as a quality control tool for the determination of free radical scavenging activity.

2. Background on Transformer Insulation's Aging

As long as chemical bonds inside the hydrocarbon chains are not broken, the generation of decay products is impeded. The sources of energy capable of splitting of a molecular covalent bond made up of a pair of electrons are three fold [1,9,16,17]:

- The strong electro-magnetic field that triggers the free electron injection process in the liquid insulation.
- The thermal energy generated by the active parts.
- Finally, the aggressiveness of dissolved oxygen.

Copper and aluminum in windings and copper alloys and iron in other transformer components act as reaction catalysts while moisture (which is considered to be the number one enemy of the paper insulation), oxygen, heat, ageing by-products, dirt, vibration, electrical stress, *etc.*, accelerate the process.

When weak bonds split, free radicals are formed. These radicals are very unstable and react quickly with other compounds, trying to capture the electron needed to gain stability. Free radicals generally attack the nearest stable molecule, "stealing" its electron. When the "attacked-molecule" loses its electron, it becomes a free radical itself, initiating a chain reaction. Once the process is started, it can cascade, finally accelerating the degradation of the insulation.

Broken molecules and "knocked-out" hydrogen atoms are free radicals and hence paramagnetic and so will move under the influence of the local magnetic field. By secondary chemical reactions, the small fractions of the broken molecules form stable compounds, mainly gases, that dissolve in the oil without modifying the one phase system. Even though the primary decay products are short-lived free radicals, as long as normal hydrocarbon chains surround and trap them, almost all pairs recombine and no net chemical reactions can take place. This phenomenon is known as a "cage effect" and occurs when the population of free radicals is minimal and the thermal agitation of molecules is low. As the population of free radicals increases, their unpaired electron can be coupled with a free electron to become a charge carrier that tends to increase the dissipation factor of the oil [18].

Due to the fact that the oxygen molecule has in its ground state two unpaired electrons, a second effect must be considered. The unpaired electron of free radicals can be coupled with one of the two unpaired electrons of oxygen molecules. This "diradical" character of oxygen makes the attack on vulnerable hydrocarbons very successful and the result of such reactions is a hydro-peroxide, see Reaction (1), that causes the oil to change color from bright yellow to amber [11,19–21]:

$$RH + O - O \to ROOH \tag{1}$$

The redox reactions promoted by the presence of copper which is capable of producing charge carriers that increase the dissipation factor of oil, has to be emphasized as well [20,21], see Reaction (2):

 $ROOH + [cation]^+ \rightarrow [cation]^{++} + RO_{-} + -OH$ (2)

The oxidation rate is accelerated by temperature and oxygen content. Since the peroxidic bond is weak and can easily be broken even by absorbing thermal energy, the two new free radicals that arise promote the auto-oxidation chain reaction [20,21], see Reaction (3):

$$ROOH \xrightarrow{\text{Thermal energy}} RO- + -OH$$
(3)

This is the initiation of the well-known auto-oxidation reaction, which can generate not only soluble decay products which darken the color of in-service-aged/used oil, but also produce large insoluble molecules. Indeed, large free radicals may combine, leading to the formation of insoluble colloidal suspensions that affect the properties of the insulating fluid. The combination of two large free radicals leads to the formation of large colloidal compounds having a molecular weight between 500 and 600 (sludge that is a solid phase) that are no longer soluble in the oil. When two large free radicals combine they can generate a similar insoluble hydrocarbon without oxygen in the middle, the decay product formed has the generic name of x-wax. The increase in concentration of free radicals increases random chemical reactions between free radicals and soluble and insoluble oil-born decay products are the outcome (Figure 1).



Figure 1. Simplified decomposition mechanism of oil insulation [17].

3. Experimental Procedures

3.1. Ionization Procedure

Free electrons are the primary source of energy for the breakdown of vulnerable covalent bonds (approximately 4 eV \approx 386 kJ·mol⁻¹). Pioneering work by Forster [22] describes the mechanism by which high-voltage fields interact with insulating oils. His extensive research in this field reveals that electrons escape from the conduction band of the metal conductor and are emitted from its surface, especially during very short but frequent commutation voltage surges [23]. Lesser quantities of energy below this threshold simply will not do the job. The free electrons injected into the liquid insulation are accelerated by the electric field. The collision of a fast electron with a hydrocarbon molecule M may be either elastic or inelastic, leading to very different results:

(i) Elastic collision:

$$e^{-}_{fast} + M_{slow} \rightarrow e^{-}_{less\,fast} + M_{faster}$$
 (4)

(ii) Inelastic collision:

$$e_{fast} + M_{slow} \rightarrow e_{slower} + M^*$$
 (excited molecule) (5)

or

$$\rightarrow e^{-}_{slower} + M^{+} + e^{-} \tag{6}$$

Whereas stable molecules reaching their singlet excitation level usually release the absorbed energy as a quantum of harmless fluorescent light:

$$M^* \to M + h\nu \tag{7}$$

Vulnerable molecules decompose and generate a pair of free radicals:

$$R - R' + h\nu \to R \cdot + R' \cdot \tag{8}$$

As the population of free radicals increases, some gaseous or liquid fractions may capture a free electron and form an ion:

$$\mathbf{R} \cdot + \mathbf{e}^- \to \mathbf{R}^- \tag{9}$$

The accumulation of such ionized molecules increases the dissipation factor of oil-paper insulation. Alternatively, large free radicals may combine, leading to the formation of an insoluble colloidal suspension.

A Merell-based test cell type, defined in the ASTM Test Method D 6180 [24], was used (Figure 2). The free electrons process are generated by a cylindrical copper electrode 15 mm (0.6 in) in diameter and 10 mm (0.4 in) long sealed in a 500 mL round bottomed glass flask. The electrode is placed in the center of the discharge cell and suspended above the oil. An oil volume of 50 mL was used to perform the tests.



Figure 2. Discharge cell according to ASTM Designation D6180 [23].

3.2. Free Radical Assessment Technique

Test specimen sampled from a transformer generally exhibits little change in 24 h (chemical reactions between free radicals is possible only by contact). A test specimen which has undergone accelerated aging contains a large amount of free radicals. As a result, the probability of a reaction between them is higher than insitu conditions. Therefore the free radical measurement must be performed within a short period of time. Both types of specimens should be sealed or nitrogen blanketed to prevent reaction with oxygen. Therefore, test specimens taken from transformers shall be tested 24 h following sampling while test specimen that has undergone any accelerated aging process is to be tested within 24 h.

The T60U UV-VIS spectrophotometer (by P.G. Instruments Ltd., Lutterworth, UK), having a wavelength accuracy ± 1 nm was used to assess the relative concentration of free radicals in electrical insulating oils. Microsoft-based software is provided to process the data. One pathway by which a liquid dielectric degrades results from the homolytic cleavage of some vulnerable hydrocarbon bonds under the effect of a high voltage.

The reactive free radical reagent, 2,2-diphenyl-1-picrylhydrazyl (DPPH), is added to a solution whose free radical concentration is to be determined. The rate that the DPPH disappears is directly proportional to the relative free radical content at a particular instant of time. The DPPH solution has an intense blue color, which changes to yellow as unpaired electrons are coupled. This reaction is monitored spectrophotometrically. The DPPH solution has a maximum absorbance at 520 nm. The disappearance of DPPH can be determined by plotting the absorbance against time at 520 nm. The presence of free radicals in solution will increase the rate at which DPPH disappears from the background solution; the higher the free radical concentration in the test specimen, the faster DPPH disappears. The relative free radical concentration of an insulating oil test specimen is determined as follows: initially,

the absorbance of the background solution of known concentration is recorded. Subsequently the decreasing absorbance of the oil specimen added to the background solution is plotted. Finally the subtraction of the oil specimen absorbance from the background solution results in a display curve of the reaction. The absorbance at 240 s/4 min from the beginning of the reaction is reported [13].

There is a direct correlation between the disappearance of the DPPH tracer and the concentration of radicals at a certain time. The concentration of DPPH of 10^{-5} mole, even at this low level, is blue-violet coloured. The formation of a new species due to DPPH reacting with the free electron of another free radical changes the colour from blue-violet to yellow, with a corresponding decrease in absorbance of the light at blue-violet wavelengths and the increase in absorbance at yellow wavelengths [25]. The larger the quantities of radicals present in the oil, the faster the reduction of light at blue-violet wavelengths and the increase at yellow wavelengths. The scavenging activity of DPPH can be represented by the following chemical reactions, where AH symbolizes an anti-oxidant and R a free radical:

$$DPPH + AH \to DPPH-H + A \tag{10}$$

$$DPPH + R \to DPPH-R \tag{11}$$

Using the UV-VIS (ultraviolet visible) spectrophotometer to sweep from 480 to 580 nm every 60 s, the measurement of the amplitude was achieved by confirming the validity in amplitude and wavelength by using a control solution. The DPPH reference solution at a 0.01% concentration normally provides a constant absorbance of 0.500 (± 0.005) at 520 nm. The absorption information can be presented also by percent transmittance as function of wavelength or wavenumber. The inverse relationship between absorbance (Abs) and transmittance (*T*) is not linear, but logarithmic [26].

3.2.1. Materials Needed

Any automated Visible Spectrophotometer capable of handling matched liquid samples using glass cuvettes of 1 cm sample path length, along with spectrograde toluene and spectrograde stable DPPH are needed. The testing procedure is done at ambient temperature 25 °C (\pm 5 °C). Any differences due to the properties of the glass cuvettes used in the tests are compensated for using the reference solution.

The DPPH decolouration in presence of free radicals is best rationalised as resulting from an abstraction reaction between DPPH and the solvent (tri-chlorobenzene, toluene or benzene) [27]. A 0.01% DPPH standard solution in toluene from an intermediate 0.1% DPPH stock solution is prepared. 0.1 grams of DPPH is dissolved in a 100 mL volumetric flask and filled to the mark with toluene for 0.1% DPPH stock solution. 5 mL of 0.1% DPPH solution is put into a 50 mL volumetric flask and fill to the mark with toluene for 0.01% DPPH solution.

3.2.2. Preparation of the Reference Solution

2 mL of the test specimen is added to a 10 mL volumetric flask and filled to the mark with toluene. The solvent diminishes the viscosity of oil to accelerate free radicals collision to couple their unpaired electrons.

3.2.3. Measurement of the Reference Solution

The scan region is set from 480 nm to 580 nm. A glass cuvette is filled with the reference solution and the instrument adjusted to read zero absorbance.

3.2.4. Preparation of the Background Solution

2 mL of DPPH standard solution (0.01%) is added to a 10 mL volumetric flask and filled to the mark with toluene. This allows having the same concentration as the oil solution.

3.2.5. Measurement of the Background Solution

The background solution is scanned to determine the maximum absorbance and its corresponding wavelength which should be 0.5 ± 0.005 at 520 nm. If it is not, the DPPH standard solution should be adjusted by either adding more stock solution or toluene and the method standard run again.

Using the time scan mode (480 nm to 580 nm), the absorbance of background solution is recorded at 520 nm for 10 min at one minute intervals. The values should be $0.500 (\pm 0.005)$. This data is saved.

3.2.6. Preparation of the Test Specimen Solution

2 mL of DPPH (eventually adjusted) and 2 mL of the oil sample are added to a 10 mL volumetric flask.

3.2.7. Measurement of the Test Specimen

The following steps allow collection and analysis of data:

- Begin timing for 2.0 min when the first drop of oil contacts the DPPH solution. Fill to the mark with toluene and transfer the test specimen solution into a glass cuvette.
- After two minutes has elapsed from the beginning of the reaction, record and scan the spectrum at one minute intervals for at least 10 min.
- Subtract the test specimen spectrum from the background solution to obtain the display curve.
- The magnitude at 520 nm is then recorded for each measurement.

4. Results of the Investigations

The investigations were carried out on virgin/new Voltesso 35 oil, and aged/used oil collected from a power transformer that has been in service for 20 years. The measurements were done before and after exposure to electrical discharge activity in the oil for 5 h. The observed reduction in absorbance reflects the kinetic of reaction between the free radicals in the DPPH solution and in the oil test specimen. The subtraction of test specimen absorbance from background solution results in the increasing display curves shown in Figures 3 and 4, the change in absorbance at 520 nm representing the total consumption of free radicals with time. The slope of the curve at a particular point also gives an indication of the rate of the reaction.



Figure 3. Transmittance with DPPH of new oil and new oil ionized as a function of time.

A comparison of the free radical reactions in new and aged oil, with no ionization and after ionization, was carried out to determine their stability under electrical discharge. The ionization procedure was performed according to ASTM D6180 [24]. The results of the investigations are presented in Figures 3 and 4.



Figure 4. Transmittance with DPPH of used oil and used oil ionized as a function of time.

It can be seen that, over time, more radicals react with the DPPH. Comparison of the initial value for new and used oils demonstrates that the presence of free radicals is increased in aged oils, *i.e.*, within 2 min more DPPH has changed from blue-violet to yellow.

The DPPH demonstrates the presence of free radicals, and can therefore serve as free radicals chemical marker in insulating fluids. The results clearly show the increase in transmittance if the oil is aged, after 10 min the transmittance has become 40% for new oil yet has become 65% for aged oil.

In addition, the rate of change of the DPPH concentration is much faster in used ionized oil than in new ionized oil. These results demonstrate the applicability of the test method in determining the relative Free Radical Content of Insulating Oils of Petroleum Origin.

The quantitative relationship between the free radicals content in the test specimen and absorbance is given by reporting its value at 2 min [13]. The measurement of the relative free radical content of the four samples, with each sample determined six times, is reported in Table 1.

Oil Condition	Percent Transmittance Values at Two Minutes after Mixing	
New oil	9.2 ± 0.3	
New oil ionized	30.7 ± 0.4	
Used oil	17.66 ± 0.4	
Used oil ionized	50.65 ± 0.6	

Table 1. Data used to develop repeatability estimates.

Repeatability

The repeatability or test-retest reliability has been evaluated based on the analysis with the same method, using the same spectrophotometer and the same oil sample (new naphthenic-based mineral oil) repeated on different days. Note that two measurements have been performed on days 8, 9, and 15 at 6 h intervals. This data is listed in Table 2. These results suggest that the repeatability is 0.31 Absorbance units.

Day Tested —	Two Minutes after Mixing	
	Absorbance	Transmittance (%)
Day 1	1.40	3.9811
Day 8	1.46	3.4674
Day 8	1.42	3.8019
Day 9	1.44	3.6308
Day 9	1.38	4.1687
Day 15	1.47	3.3884
Day 15	1.43	3.7154
Day 28	1.47	3.3884
Day 29	1.47	3.3884
Day 33	1.51	3.0903

Table 2. Data used to develop repeatability estimates.

5. Conclusions

In this manuscript, a laboratory testing technique is proposed to assess free radical concentration in insulating mineral oil. A reactive free radical reagent: 2,2-diphenyl-1-picrylhydrazyl (DPPH), was added to a solution whose free radical concentration is to be determined using a UV-VIS spectrophotometric method. The quantitative relationship between a standard electrical stress during a given period of time and the amount of vulnerable hydrocarbons in mineral insulating oils was

addressed. All generated free radicals, large and small, are paramagnetic. In a real life conditions, these are attracted by the windings contributing to the partial discharges. Hence, these unstable molecules can be particularly harmful during a transient over voltage, when the generated gases and ionized decay products can initiate incipient electrical failure.

In addition, the theoretical premises that, incipient electrical discharges contribute to free radical generation, is experimentally confirmed under laboratory conditions.

This testing method which is applicable to new oils as well as naturally or artificially oxidized oil, may be adopted for Measurement of the relative Free Radical Content of Insulating Oils of Petroleum Origin. With this test method, the role played by free radicals in the oil decaying and gassing can be investigated.

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Author Contributions

John Sabau designed this research and gave the whole guidance; Issouf Fofana and Amidou Betie collected all the data, carried out calculations, result display and analysis and wrote the manuscript. All authors read and approved the final manuscript.

Conflicts of Interest

The authors declare no conflicts of interest.

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