2005

Experimentally Derived Resistivity for Dielectric Samples From the CRRES Internal Discharge Monitor

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Recommended Citation

Green, Nelson W.; Frederickson, A. Robb; and Dennison, JR, "Experimentally Derived Resistivity for Dielectric Samples From the CRRES Internal Discharge Monitor" (2005). All Physics Faculty Publications. Paper 1476.

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Abstract

Resistivity values were experimentally determined using charge storage methods for six samples remaining from the construction of the Internal Discharge Monitor (IDM) flown on the Combined Release and Radiation Effects Satellite (CRRES). Three tests were performed over a period of four to five weeks each in a vacuum of \(\sim 5 \times 10^{-6}\) torr with an average temperature of \(\sim 25^\circ C\) to simulate a space environment. Samples tested included FR4, PTFE, and alumina with copper electrodes attached to one or more of the sample surfaces. FR4 circuit board material was found to have a dark current resistivity of \(\sim 1 \times 10^{18}\) \(\Omega\)-cm and a moderately high polarization current. Fiber filled PTFE exhibited little polarization current and a dark current resistivity of \(\sim 3 \times 10^{20}\) \(\Omega\)-cm. Alumina had a measured dark current resistivity of \(\sim 3 \times 10^{17}\) \(\Omega\)-cm, with a very large and more rapid polarization. Experimentally determined resistivity values were two to three orders of magnitude more than found using standard ASTM test methods. The one minute wait time suggested for the standard ASTM tests is much shorter than the measured polarization current decay times for each sample indicating that the primary currents used to determine ASTM resistivity are caused by the polarization of molecules in the applied electric field rather than charge transport through the bulk of the dielectric. Testing over much longer periods of time in vacuum is required to allow this polarization current to decay away and to allow the observation of charged particles transport through a dielectric material. Application of a simple physics-based model allows separation of the polarization current and dark current components from long duration measurements of resistivity over day- to month-long time scales. Model parameters are directly related to the magnitude of charge transfer and storage and the rate of charge transport.

Introduction

Standard constant-voltage ASTM test methods of very high resistivity dielectrics [1,2] do not provide accurate resistivity values for dielectrics appropriate for use in spacecraft charging applications [3,4]. These standard methods rely on electrometer measurements of current, voltage or resistance and are typically instrumentation resolution limited to accurate measurements of resistivities of less than \(10^{12}\) to \(10^{15}\) \(\Omega\)-cm [1,4]. Inconsistencies in sample humidity, sample temperature, initial voltages and other factors from such tests cause significant variability in results [1]. Further, the duration of standard tests are short enough that the primary currents used to determine resistivity are often caused by the polarization of molecules by the applied electric field rather than by charge transport through the bulk of the dielectric [4,5,6]. Testing over much longer periods of time in a well-controlled vacuum environment is required to allow this polarization current to become small so that accurate observation of the more relevant charged particle transport through a dielectric material is possible. For space applications this is particularly important since dielectrics on the spacecraft will be exposed to space plasmas and radiation for months or years. Unless dissipated by leakage through the dielectric, charge will build up within the dielectric inducing large electric fields that can lead to dielectric breakdown and potentially harmful ESD pulses.

Selected samples remaining from the Internal Discharge Monitor (IDM) experiment on the CRRES satellite [7,8] were tested for charge storage for NASA at the Jet Propulsion Laboratory. The sample set on CRRES was chosen to cover a range of dark current resistivity values and polarization magnitudes and rates. Hence, the set provides an excellent test bed for both the charge storage method of resistivity measurements and behavior of dielectrics in the space environment. By measuring the decay of stored charge in these dielectric samples, more accurate and appropriate resistivity values for the sample materials have been determined. Preliminary measurements of resistivities measured with the charge storage method for similar samples were shown to be critical in accurate modeling of the discharge.

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Table 1. List of Samples with CRRES IDM channel reference

| Material  | Thickness (cm) | Electrode | Mount Type | IDM Channel | Material Properties (ASTM Standard) [1] | | Material (D150) | (D 257) | (D 150) | (D 149) |
|-----------|----------------|-----------|------------|-------------|------------------------------------------|-----------|----------|----------|----------|
| PTFE      | 0.229          | Dual      | Open       | 11          | 2.1 @ 1 MHz                              | 1×10^18   | 0.0003   | 20       |
| PTFE*     | 0.229          | Back      | Open       | 16          | 5.4 @ 1 kHz                              | >10^9     | 0.035    | 27       |
| FR4*      | 0.119          | Back      | Closed     | 15          | 9.6 @ 1 MHz                              | 1×10^14   | 0.001    | 9.8      |
| FR4       | 0.119          | Back      | Open       | 15          |                                          |           |          |          |
| FR4       | 0.317          | Dual      | Open       | 8           |                                          |           |          |          |
| FR4       | 0.317          | Back      | Open       | 4,12        |                                          |           |          |          |
| Alumina*  | 0.102          | Back      | Open       | 7           |                                          |           |          |          |

* Full analysis presented in this paper.

Figure 1. Diagram of vacuum chamber arrangement as used while testing the CRRES IDM samples.

Figure 2. Detail of the capacitive measurement system used to measure sample surface potential

pulsing of samples during the CRRES mission [9,10]. The new resistivity values reported here are expected to further enhance the usefulness of the knowledge gained from the IDM experiment by producing experimental resistivity values for several of the samples.

Samples tested were 5x5 cm squares with copper electrodes on one or both surfaces. Materials included fiber-filled PTFE, Micaply FR4, and alumina (Al₂O₃) [7]. Three sets of tests were performed over a period of four to five weeks each in a vacuum of ~5×10⁻⁶ torr to simulate a space environment. Details for each sample, including standard ASTM material properties and the corresponding CRRES IDM channel, are given in Table 1. Pulse histories from the CRRES IDM for each sample are documented in the references [9,11,12].

**Test Procedure**

Samples were mounted on a circular carousel (Figure 1) inserted into a vacuum chamber behind another metallic plate with a single opening into the interior allowing each sample to be charged individually. Also mounted on the shutter was an electrically isolated sensor plate used to measure each sample’s surface potential one at a time from outside of the chamber with a capacitive voltmeter [Trek, model 341] (Figure 2). Measurements represented an average surface potential over an area approximately equal to the 19 cm² surface area of the sensor plate. Connections to the electrodes on the back of each sample were brought through the chamber door for individual control or monitoring of each sample when charging. A calibration coefficient was calculated for each sample to relate measured potentials to actual sample surface potentials.
Samples were charged with electrons by one of two methods: placing a positive potential on each sample and attracting thermionically generated electrons from an energized filament near ground potential, or by floating an energized filament at a highly negative potential compared to the grounded samples. In either case, the energy of incident electrons was roughly equal to the difference between the filament and the sample potentials. For the three samples analyzed fully in this paper, the former method was used.

Three charging runs lasting for 20, 25, and 35 days respectively were performed with the CRRES IDM samples. Two charging runs were conducted successively after an initial 4 day sample conditioning in vacuum. The third run was performed on the same samples after approximately two months at atmosphere, after a 2 day sample conditioning period in vacuum. Sample temperature was not closely monitored, but an average temperature of 25 °C (laboratory room temperature) is assumed. Measurements of the surface potentials were taken initially every few minutes, but as the changes between successive measurements became smaller, the interval between measurements increased first to hours then to days.

Further details of the instrumentation and test methods are found in the references [3,5,6,10,13].

Resistivity Model

Since the actual amount of charged particles on the surface of the materials could not be measured directly, each sample’s surface potential was monitored to observe the changes in the electric field due to polarization of the material and, ultimately, dark current conduction of charge though the dielectric. A relatively rapid initial drop in the surface potential was expected for each sample due to dielectric polarization in the sample material. This initial decrease in potential was found to vary widely due to material properties. As any polar molecules in the material rotated to align with the electric field created by the charges on the surface of the sample, or migrate within the dielectric to interfaces, they created a polarization electric field in opposition to that formed by the incident electrons. Since the measured surface potential was dependent on electric field strength from the sample, the opposing field reduced the measured voltage without necessarily indicating a reduction in the number of charged particles on the surface of the sample. Simultaneously, charged particles may have been conducted through the material, but the majority of the short-term change in surface potential for high resistivity materials was thought to be through polarization of the sample material. As polarization reached saturation, further change in surface potential due to this effect became negligible and any further change was due to a reduction in the number of charged particles remaining on the surface of the charged sample. The charged particles that left the surface moved into the dielectric material filling electron traps or conducting through the material to ground. The dark current resistivity of the material was determined by the rate of charged particle transport, in the long-term asymptotic limit of charge storage measurements.

A simple model of the measured surface voltage as a function of elapsed time for the charge storage method \( V_{CS}(t) \) in terms of the initial and final surface voltages \( (V_o, V_\infty) \) and initial and final relative permittivities \( (\varepsilon_o, \varepsilon_\infty) \), where \( \varepsilon_\infty=8.854 \times 10^{-12} \) F/m is the permittivity of free space, \( \varepsilon \) is the permittivity in a dielectric medium, and \( \varepsilon_r \equiv \varepsilon/\varepsilon_\infty \) is the relative permittivity predicts [4]

\[
V_{CS}(t; V_o, V_\infty, \varepsilon_r^o, \varepsilon_r^\infty, \tau_{DC}, \tau_p) = \frac{(V_o - V_\infty) e^{-t/\tau_{DC}} + V_\infty}{(\varepsilon_r^o - \varepsilon_r^\infty) e^{-t/\tau_p} + \varepsilon_r^o}
\]

(1)

The polarization decay time, \( \tau_p \), measures the rate of the response of the medium to an applied electric field, and can be thought of as the rate at which the dipoles align within the material to the electric field \( E \). It is the time it takes for the bound surface charge to increase to \((1-1/e)\) (or 63%) of its final value. The charge storage decay time, \( \tau_{DC} \), is the time it takes for the free surface charge to drop to \( 1/e \) (or 37%) of its initial value and is directly proportional to the dark current resistivity \( \rho_{DC}=\tau_{DC}/(\varepsilon_o \varepsilon_\infty) \). Note that in this simple model, the polarization decay time, dark current decay time and resistivity are all intrinsic material properties, independent of surface area or thickness. If there is no initial polarization, \( \varepsilon_r^o=1 \). If there are no free charges trapped within the dielectric as it is transported through the material and \( t \rightarrow \infty \), then this results in a residual potential, \( V_o=0 \). In the limit of short time, with \( \tau_{DC}=\tau_p \) and \( \varepsilon_r^\infty=1 \),

\[
\lim_{t \rightarrow \tau_p} V_{CS}(t; V_o, \varepsilon_r^\infty, \tau_p) = V_o \left[ \varepsilon_r^\infty \left( 1 - e^{-t/\tau_p} \right) \right]^{-1}
\]

(2)

In the limit of long time, with \( \tau_{DC}=\tau_p \), \( \varepsilon_r^o=1 \) and \( V_o=0 \),

\[
\lim_{t \rightarrow \infty} V_{CS}(t; V_o, \tau_{DC}) = \frac{V_\infty}{e^{-t/\tau_{DC}}}
\]

(3)
Figure 3. Surface potentials functions of time for (a) PTFE, (b) FR4 and (c) alumina. Curves shows fits with three parameter fit using Equation (1) (dashdot), five parameter fit using Equation (1) (solid), early time limit model using Equation (2) (dashed) and the late time limit model with Equation (3) (dotted). Note the log-log plots of (b) and (c). For (c), there is also a modified 3-parameter fit with an additional decay mechanism. Charge as a function of elapsed time for (d) PTFE, (e) FR4 and (f) alumina. Plots are based on a three parameter fit using Equation (1). The initial and final values of the free charge from the fit are also shown.
The unusually large (~20%) residual voltage, and the defect density is high due to the composite nature of the material. The FR4 samples tested were a thermoset epoxy resin, fiberglass reinforced, Cu-clad laminate made by Micaply 3M Co. The FR4 samples displayed intermediate charge storage characteristics. FR4 showed a fairly rapid initial drop and the slow rise of the bound charge predicted in Figure 4b. The higher ratio of total charge to free charge in Figure 3b is indicative of higher polarization than in PTFE and a relative dielectric constant of >5. The polymer and glass in FR4 have permanent dipoles—unlike PTFE—and the defect density is high due to the composite nature of the material. The unusually large (~20%) residual voltage, ~2.1×10^3 (Ω·cm), a factor of ~5 less than our measured ρDC.

### Test Results

A total of seven samples were charged and monitored for each of the three runs. Analyses of the data for three of the samples are presented below representing the general results for each sample material. For each analysis presented, the surface voltage measurements were fit using a least-squares fit method for:

(i) the full data set using Eq. (1) with five fitting parameters, \( V_0, \varepsilon_r, V_\infty, \tau_P, \tau_{DC} \), and \( \rho_D \);
(ii) the full data set using Eq. (1) with three fitting parameters \( \varepsilon_r, \tau_{DC} \), and \( \rho_D \) plus \( \varepsilon_r=1 \) and \( V_0=0 \),
(iii) the initial six data points using Eq. (2) with \( \varepsilon_r \) and \( \tau_P \) as fitting parameters, and
(iv) the last six data points using Eq. (3) with \( \tau_{DC} \) as a fitting parameter.

In each case, \( V_0 \) was set to the measured initial voltage. Results for the fits are listed in Table 2.

### PTFE Charge Decay

The PTFE samples tested were a “Type 250” fiber–filled composite with a polytetrafluoroethylene matrix from the 3M Co. The decay pattern of the PTFE samples is significantly different from that of the other samples tested, and reflects the physical properties of the material. PTFE is known as a non-polar polymer, with a very low polarizability and the high affinity of fluorine for its electrons, the polymer has no permanent dipole moment and orientational polarization is not a major contributor. Thus, polarization in PTFE results rapidly from induced dipoles through electronic and atomic polarization or more slowly due to defects through interfacial polarizability. Response of the long chain polymers and modifications of defects occurs slowly for PTFE, as evidenced by the relatively long polarization decay time \( \tau_P \sim 18 \text{ hr} \) and the slow rise of the bound charge predicted in Figure 3b. PTFE has a very high dark current resistivity; this is evident in the very large value of the dark current decay constant \( \tau_{DC} \sim 1 \text{ yr} \) and in the slow decay of free charge predicted in Figure 3b. The measured \( \rho_D \) is ~300 times larger than the \( \rho_{ASTM} \) value from standard handbooks. The polarization decay constant corresponds to a resistivity of \(~7\times10^{17} \Omega\cdot\text{cm}\), which is only slightly less than the ASTM value of \(>1\times10^{17} \Omega\cdot\text{cm} \); this is consistent with the ASTM fallacy of making measurements after only 1 min of voltage application, when the polarization current still dominates.

### FR4 Charge Decay

The FR4 samples tested were a thermoset epoxy resin, fiberglass reinforced, Cu-clad laminate made by Micaply Co. FR4 is a standard designation for a broad class of composite materials typically used for printed circuit boards [15,16]. FR4 showed a fairly rapid initial drop in potential immediately after charging due to polarization. Response of the long chain polymers and modifications of defects of the FR4 composite were similar to those for PTFE, as evidenced by a similar long polarization decay time \( \tau_P \sim 25 \text{ hr} \) and the slow rise of the bound charge predicted in Figure 4b. The higher ratio of total charge to free charge in Figure 4b is indicative of higher polarization than in PTFE and a relative dielectric constant of \(>5 \). The polymer and glass in FR4 have permanent dipoles—unlike PTFE—and the defect density is high due to the composite nature of the material. The unusually large (~20%) residual voltage, \( V_\infty \), suggests that there is substantial residual charge in the FR4 sample. The FR4 has a dark current resistivity between the other two samples; this is evident in the intermediate dark current decay constant \( \tau_{DC} \sim 5 \text{ days} \) and in the modest decay of free charge predicted in Figure 4b. Comparison of the measured \( \rho_D \) to an ASTM standard value is not meaningful; the ASTM value listed [14] was not for the specific material tested but was rather from the FR4 standards [15,16] that only specifies that \( \rho_{ASTM} \) not be less than \(10^{17} \Omega\cdot\text{cm} \). Measurements with a different technique on a similar FR4 spacecraft material found a dark current resistivity of \(~2.12\times10^{17} \Omega\cdot\text{cm} \), a factor of ~5 less than our measured \( \rho_D \).

<table>
<thead>
<tr>
<th>Material</th>
<th>Thickness (cm)</th>
<th>( \varepsilon_r )</th>
<th>( \varepsilon_r' )</th>
<th>( V_0 ) (volt)</th>
<th>( V_\infty ) (volt)</th>
<th>( \tau_P ) (hr)</th>
<th>( \tau_{DC} ) (day)</th>
<th>( \rho_D ) parameter (Ω·cm)</th>
<th>( \rho_1 ) parameter (Ω·cm)</th>
<th>( \rho_5 ) parameter (Ω·cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTFE</td>
<td>0.229</td>
<td>1.01</td>
<td>1.11</td>
<td>778</td>
<td>10.2</td>
<td>17.9</td>
<td>339</td>
<td>(3.0\times10^9)</td>
<td>(2.9\times10^9)</td>
<td>(3\times10^2)</td>
</tr>
<tr>
<td>FR4</td>
<td>0.317</td>
<td>1.03</td>
<td>4.68</td>
<td>498</td>
<td>107</td>
<td>25.1</td>
<td>5.01</td>
<td>(1.1\times10^{18})</td>
<td>(1.6\times10^{18})</td>
<td>(&lt;1\times10^9)</td>
</tr>
<tr>
<td>Alumina</td>
<td>0.102</td>
<td>1.02</td>
<td>3.00</td>
<td>318</td>
<td>5.14</td>
<td>6.35</td>
<td>0.891</td>
<td>(2.9\times10^{17})</td>
<td>(3.0\times10^{17})</td>
<td>(3\times10^4)</td>
</tr>
</tbody>
</table>

* Results listed in columns 2-7 are for 5-parameter fits using Eq. (1).
Alumina Charge Decay

The alumina sample tested was a ~1 mm thick bulk alumina material, attached to a Cu substrate with silver–filled epoxy [7]. The alumina is believed to be Type II material with a Al$_2$O$_3$ content of >93% [14]; this is reflected in the values listed in Table 1. The behavior of the alumina sample is significantly different than the PTFE and FR4 polymer samples, due to its nature as a ceramic. Alumina has one of the highest dielectric constants of common ceramics, with a value of about 10. This follows mostly from the large permanent dipole moment of the Al$_2$O$_3$ unit cell that results from appreciable charge redistribution in the ionic/covalent bonds. The observation that the polarization decay constant of alumina is shorter than the polymers is too be expected as much of the polarization of alumina results from atomic polarizability, that is distortion of the atoms within the unit cell. This leads to a large initial rise in the bound charge (see Figure 5b). However, the bound charge never exceeds the initial free charge because the polarization decay constant $\tau_p$~6 hr is not too much shorter than $\tau_{DC}$. This behavior is evident in the decay of the bound charge in Figure 5b. The alumina has a much lower dark current resistivity than either polymer; this is evident in the relatively small dark current decay constant $\tau_{DC}$~20 hr and in the more rapid decay of free charge predicted in Figure 4b. The measured polarization and dark current resistivities are both approximately 3 orders of magnitude larger than the ASTM handbook value of $\sim 1 \times 10^{18}$ $\Omega$-cm [14]. The fact that $\rho_{ASTM}$ $\ll \rho_p$ may reflect the sensitivity of alumina to the nature of defects of specific samples or to the humidity.

It is interesting to note that there is evidence of a small charge (~1% of the initial free charge) that decays with a very long decay constant of >1 yr. This is apparent in the long time charge decay in Figure 5a. This term was modeled by modification of the exponential term of the numerator of Eq. (1) to include a second decay mechanism, $e^{-t/\tau_{DC}} \rightarrow e^{-t/\tau_{DC}} + \alpha_H e^{-t/\tau_H}$. A modified 3-parameter fit found $\tau_H = 2.84$, $\tau_p = 4.85$ hr, $\tau_{DC} = 19.8$ hr $\rightarrow \rho_{DC} = 2.6 \times 10^{17}$ $\Omega$-cm with $\alpha_H = 0.9\%$ and $\tau_p = 17.1$ days. We speculate that this may be related to the slow dissipation of charge trapped in deep level defect states of the alumina.

Conclusion

Laboratory testing has found that resistivity values for samples tested with the charge storage method were two to three orders of magnitude more than those given by standard ASTM test methods. The difference in measured resistivity is largely attributed to the dominance of polarization currents in the first hours after the application of an external electric field. When charge is deposited on the surface of dielectric samples held in a vacuum, the polarization current decays to an insignificant value, typically this effect is much faster than the dissipation of charge through the material. After the polarization current has been minimized, charge transport can more easily be observed and the resistivity calculated. The semi-empirical model applied in this paper has been found to accurately fit the data and to produce physically reasonable results based on the fitting parameters.

Three dielectric materials were tested and general results are listed in the analysis above. Fiber filled PTFE exhibited little polarization current and a dark current resistivity of $\sim 3 \times 10^{20}$ $\Omega$-cm. FR4 circuit board material was found to have a dark current resistivity of $\sim 1 \times 10^{19}$ $\Omega$-cm. Alumina had a measured dark current resistivity of $\sim 3 \times 10^{17}$ $\Omega$-cm, with very large and more rapid polarization.

With these measured values, and others to come, the detailed analysis of the charging history of the CRRES IDM mission begun with great success by Frederickson and Brautigam [9] can be continued for more CRRES samples. It should be noted that the values calculated here are for samples that have not been exposed to radiation and have only been exposed to small amounts of low energy electrons. The resistivity of these materials may change, and change significantly, with exposure to space radiation. These results need to be verified through further analysis of the gathered data including that for other thicknesses and additional electrode configurations.

Acknowledgements

The authors would like to give thanks and acknowledgement to Jerilyn Brunson of Utah State University and Charles Benson of the Jet Propulsion Laboratory for their help and assistance in this effort. The research described in this paper was carried out at the Jet Propulsion Laboratory, California Institute of Technology, under a contract with the National Aeronautics and Space Administration. The majority of the support for this work was from a contract for the NASA Space Environments and Effects (SEE) Program.
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