Insulation Resistance Degradation in Ni–BaTiO₃ Multilayer Ceramic Capacitors

Donhang (David) Liu

Abstract—Insulation resistance (IR) degradation in Ni–BaTiO₃ multilayer ceramic capacitors has been characterized by the measurement of both time to failure (TTF) and direct current leakage as a function of stress time under highly accelerated life test conditions. The measured leakage current-time dependence data fit well to an exponential form, and a characteristic growth time \( \tau \) can be determined. A greater value of \( \tau \) represents a slower IR degradation process. Oxygen vacancy migration and localization at the grain boundary region results in the formation of Schottky barrier height and has been found to be the main reason for IR degradation in Ni–BaTiO₃ capacitors. The reduction of barrier height as a function of time follows an exponential relation of \( \phi(t) = \phi(0)e^{-2kt} \), where the degradation rate constant \( K = k_0e^{(-E_d/kT)} \) is inversely proportional to the mean TTF (MTTF) and can be determined using an Arrhenius plot. For oxygen vacancy electromigration, a lower barrier height \( \phi(0) \) will favor a slow IR degradation process, but a lower \( \phi(0) \) will also promote electronic carrier conduction across the barrier and decrease the IR. As a result, a moderate barrier height \( \phi(0) \) (and therefore a moderate IR value) with a longer MTTF (smaller degradation rate constant \( K \)) will result in a minimized IR degradation process and the most improved reliability in Ni–BaTiO₃ multilayer ceramic capacitors.

Index Terms—Barium titanate, ceramic capacitors, dielectric degradation, insulation resistance (IR), reliability.

I. INTRODUCTION

Insulation resistance (IR) degradation related to oxygen vacancy migration has been considered to be the primary cause of reliability degradation of multilayer ceramic capacitors (MLCCs) with base-metal electrodes (BMEs). The behavior is characterized by a slow increase in the leakage current under an applied direct current (dc) field stress. To reveal IR degradation in a timely manner, MLCCs are often degraded under highly accelerated life test (HALT) conditions with different temperatures and applied voltages. Previous studies have shown that there are three possible factors related to the IR degradation of BaTiO₃-based BME MLCCs: the dielectric layer, the BaTiO₃ grain boundaries, and the Ni–BaTiO₃ internal electrode interfaces [1]–[4].

Unlike traditional BaTiO₃-based MLCCs with precious-metal electrodes (PMEs), BME MLCCs are co-fired in a reducing atmosphere to avoid oxidation of the electrodes. Despite the reoxidation process, there is still a significant amount of oxygen vacancies that are accommodated in the BaTiO₃ dielectric layers. The failure mechanism of BME MLCCs is thought to be dominated by the electromigration of oxygen vacancies through the grain boundaries in the dielectric layers [5]–[8]. Wasser [9], Wasser et al. [10], [11], and Baiatu et al. [12] studied the IR degradation in ambient-fired SrTiO₃ ceramic and acceptor-doped single-crystal SrTiO₃. The results showed that IR degradation begins with oxygen vacancy electromigration toward the cathode with respect to time, field, and temperature. Segregation of defects and dopants is found at the grain boundaries during the sintering process and results in the formation of space-charge layers at the grain boundaries.

The formation of double Schottky depletion layers at the grain boundaries of ceramic BaTiO₃ and their impact on the properties of BaTiO₃ ceramics was first proposed in [13] to explain the unique positive temperature coefficient of resistance (PTCR) behavior around the Curie temperature, which only existed in donor-doped semiconducting BaTiO₃ ceramics. This model indicates that the depletion barriers are formed because of the electron trapping by acceptor states at grain boundaries. Jonker [14], [15] later extended the Heywang model, considering the influence of ferroelectric polarization on resistivity below the Curie temperature.

In BaTiO₃-based MLCCs, the depletion layers are believed not only to be depleted of electron carriers and therefore to be highly resistive but also to act as electrical barriers against oxygen vacancy electromigration and, thus, to slow down the degradation process [2], [10], [16]. Although high-resistance depletion layers both at grain boundaries and at electrode interfaces limit electronic conduction and the transport of oxygen vacancies across dielectric layers, oxygen vacancy migration is either trapped near the grain boundary depletion layers or blocked by electrode interfaces, and these charged oxygen vacancies are neutralized by the reduction of Ti⁺⁺ near the cathode and the rare-earth (RE) doping in the BaTiO₃ dielectric. This agrees with a number of recently published works about first-principles calculations of RE element doping, the local atomic configuration, and the solution energy of oxygen vacancies [17]–[19].

In this paper, the IR degradation in Ni–BaTiO₃ MLCCs was investigated for three commercial BME capacitors qualified to the same reliability level but made by different manufacturers. This paper provides insight into the time-dependent correlations among oxygen vacancy migration, vacancy trapping,
TABLE I

<table>
<thead>
<tr>
<th>Part ID</th>
<th>Stress Level</th>
<th>E (kV/mm)</th>
<th>V/Grain</th>
</tr>
</thead>
<tbody>
<tr>
<td>AA46450</td>
<td>250V 175C</td>
<td>39.1236</td>
<td>14.75</td>
</tr>
<tr>
<td>0.47μF, 50V, 0805</td>
<td>225V 165C</td>
<td>35.2113</td>
<td>13.27</td>
</tr>
<tr>
<td>Manufacturer A</td>
<td>250V 165C</td>
<td>39.1236</td>
<td>14.75</td>
</tr>
<tr>
<td>98 layers</td>
<td>250V 155C</td>
<td>39.1236</td>
<td>14.75</td>
</tr>
<tr>
<td>BaTiO3 thickness= 6.39 μm</td>
<td>315V 155C</td>
<td>49.2958</td>
<td>18.59</td>
</tr>
<tr>
<td>Ave. grain size= 0.38 μm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AB47450</td>
<td>250V 175C</td>
<td>43.0886</td>
<td>14.13</td>
</tr>
<tr>
<td>0.47μF, 50V, 0805</td>
<td>225V 165C</td>
<td>38.7797</td>
<td>12.72</td>
</tr>
<tr>
<td>Manufacturer B</td>
<td>250V 165C</td>
<td>43.0886</td>
<td>14.13</td>
</tr>
<tr>
<td>100 layers</td>
<td>250V 155C</td>
<td>43.0886</td>
<td>14.13</td>
</tr>
<tr>
<td>BaTiO3 thickness= 5.80 μm</td>
<td>315V 155C</td>
<td>54.2916</td>
<td>17.81</td>
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<tr>
<td>Ave. grain size= 0.33 μm</td>
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<td></td>
<td></td>
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<tr>
<td>AC47450</td>
<td>250V 175C</td>
<td>30.8642</td>
<td>12.45</td>
</tr>
<tr>
<td>0.47μF, 50V, 0805</td>
<td>225V 165C</td>
<td>27.7778</td>
<td>11.20</td>
</tr>
<tr>
<td>Manufacturer C</td>
<td>250V 165C</td>
<td>30.8642</td>
<td>12.45</td>
</tr>
<tr>
<td>103 layers</td>
<td>250V 155C</td>
<td>30.8642</td>
<td>12.45</td>
</tr>
<tr>
<td>BaTiO3 thickness= 8.10 μm</td>
<td>315V 155C</td>
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<td>15.68</td>
</tr>
<tr>
<td>Ave. grain size= 0.40 μm</td>
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</table>

and the depletion layer height. A theoretical model based on the oxygen vacancy migration and entrapment at grain boundary is proposed to show the relationship between the reliability, characterized by mean time to failure (MTTF), and the depletion layer height reduction during the IR degradation in the BaTiO3 dielectric materials of BME capacitors.

II. EXPERIMENTAL PROCEDURE

Three automotive-grade commercial BME capacitors, AA47450, AB47450, and AC47450, with the same chip size, capacitance, and rated voltage, but made by different manufacturers, were selected for this paper. These BME capacitors were qualified per AEC-Q200, a specification document developed by the Automotive Electronics Council of United States for ceramic capacitors and passive components used in harsh automotive environments. The microstructures of the three BME capacitor lots were investigated by cross section processing of five units per capacitor type and were examined using a scanning electron microscope to reveal the number of dielectric layers, average grain size, and average dielectric layer thickness.

Table I summarizes the specifications, number of dielectric layers, corresponding electrical field (V/thickness), and volts per grain at a given stress condition used to degrade the capacitors to reveal their failure modes. The electric field can be calculated by dividing the measured dielectric thickness by the applied voltage. The volts per grain (the voltage applied to each individual BaTiO3 grain) can be calculated using the dielectric thickness and measured average grain size data from the microstructure analysis.

The results in Table I show that at a given applied voltage, the values of volts per grain are very similar among the three BME capacitor lots, indicating that these capacitors are not only manufactured and qualified to the same reliability level but they will also have a similar applied electrical strength during degradation. The three BME capacitor lots were degraded together under the same HALT temperature and applied voltage conditions, as shown in Table I. Twenty BME capacitors were tested for each stress condition and for each capacitor type.

The traditional HALT measures only the time-to-failure (TTF) data of each capacitor at a given stress condition, and then uses a statistical model to determine the MTTF data at this stress condition [19]–[22]. This approach is based on a single failure mode assumption and is adequate for most ceramic capacitors with precious-metal electrodes (PMEs). However, as has been previously shown [26]–[29], many BME capacitors, when degraded under HALT conditions, reveal a more complicated failure mechanism with two distinct failure modes. A recently performed failure analysis of these degraded BME capacitors also confirms the existence of two distinct failure modes [37]. For this reason, the measurement of TTF data alone will not result in an accurate prediction of the reliability life of these BME capacitors. To characterize the multiple failure modes in BME capacitors, the leakage current was also monitored in situ and was measured every 1–3 s for each capacitor unit. The reliability life of a capacitor is, thus, not only determined by the TTF data but also by the measured leakage current data used to distinguish the failure modes. To differentiate this new approach from the traditional single-failure mode HALT that measures TTF data only, this modified method that has been used with multiple failure modes has been dubbed highly accelerated life stress testing (HALST) [29]. The term HALST is used through the remainder of this paper as appropriate.

When IR degradation is accelerated under HALST conditions, a certain value of IR or of leakage current must be used as the failure criterion at which the TTF can be determined. Per MIL-PRF-123, BaTiO3-based MLCCs built for high-reliability applications were constructed with a minimum dielectric thickness of 25 μm for rated voltages greater than 50 V. In this case, the leakage current level was very low, even under HALST conditions. The MLCC was considered a failure when IR at a given stress condition was degraded three to four orders below its initial value [20], [21]. Waser et al. [10] have also defined characteristic lifetime \( t_{ch} \) at which the leakage
current has risen one decade above its minimum value to evaluate the degradation rate. With the progress in Ni–BaTiO₃ MLCC development, dielectric thicknesses have been significantly reduced, even below 2 μm [22]. As a result, leakage current has increased significantly, and Ni–BaTiO₃ MLCCs are considered to have failed at a fairly low IR value. Some reports define the TTF as the time at which IR is degraded to a value of 500 kΩ [22], [23], 1000 kΩ [24], or 10% of its initial value [4], while others set a constant leakage current limit at 100 μA for all of the stress conditions being used [25]–[29]. After a review of most of the failure criteria that have been reported for determining the TTF of MLCCs with BaTiO₃ dielectrics, the author decided to use a single current limit of 100 μA to determine the TTF for all stress levels and for all MLCC samples, based on the following considerations.

During HALST, each MLCC unit was connected in series to a current-limiting resistor with a dc power supply. The voltage drop across the series resistor was measured and used to calculate the leakage current of the capacitor. Since the voltage drop on the current-limiting resistor is dependent on the current value, a leakage current higher than 100 μA can result in a voltage drop on the current-limiting resistor of more than 5% of the overall applied voltage. This is true for a number of BME capacitors with thin dielectric layers that can be degraded under fairly low applied voltages. Although this voltage drop can be reduced if a smaller value resistor is used, the voltage value reading on a small resistor may be too low to ensure the accuracy of the IR measurement. For example, when an MLCC with a 100 kΩ resistor is accelerated at 150 V, the voltage drop will be 100 kΩ · 100 μA = 10 V, which is 6.7% of the total applied voltage. For a 1 kΩ resistor, the voltage drop is only 0.1 V across the resistor, which may be too low to show the details of the leakage current changing with the stress time. In this paper, the values of the resistor were chosen such that the maximum voltage drop at the current-limiting resistor will be no more than 3% of the applied voltage under all HALST conditions.

III. RESULTS AND DISCUSSION
A. Characterization of Time-Dependent Leakage Current and MTTF Data

1) Leakage Current and MTTF Data: Fig. 1 shows the leakage current against stress time for the three BME capacitor lots at a given stress condition: 155 °C, 250 V (5 × rated voltage).

The plots in Fig. 1 have the same scale on the x-axis. It can be seen that these capacitor lots, when degraded at the same stress condition, revealed significantly different TTF data. Further details about the characterization of the failure patterns and about the method to determine the acceleration factors have been discussed in [29].

A two-parameter Weibull plot can be made when TTF data at a given stress level are available. The MTTF, a statistical parameter that measures reliability, can be determined as

\[ \text{MTTF} = \eta \Gamma(1 + \beta^{-1}) \]  \hspace{1cm} (1)

where slope $\beta$ is the dimensionless shape parameter whose value is often characteristic of the particular failure mode, $\eta$ is the scale parameter that represents the time at which 63.2% of the population has failed, and $\Gamma(x)$ is the gamma function of $x$.

Table II summarizes the calculated MTTF data using (1) when all TTF data at a given stress level were used to make a two-parameter Weibull plot. The reliability life, as characterized by MTTF, was more than one magnitude of difference among the capacitor lots under the same stress condition.

2) Characterization of Leakage Current Data for a Slow Degradation Failure Mode: As shown in Table I, all three BME capacitor lots that were made for the same reliability level revealed almost identical construction and microstructure parameters and were also degraded under very similar stress conditions. However, the MTTF data contained in Table II show significant differences in reliability life for these BME capacitors. To understand the mechanism that determines the reliability life, the leakage current data shown in Fig. 1 were replotted with a different scale in the x-axis to reveal...
Fig. 2. Leakage data are replotted from Fig. 1 with a different scale in the x-axis to reveal details in the failure modes among the three BME capacitor lots.

details of the differences in their failure modes. As shown in Fig. 2, the TTF data appear to be highly dependent on the failure mode exhibited during the HALST regimen.

As discussed in [27]–[29], two failure modes can be identified in these BME capacitor lots: catastrophic and slow degradation. A catastrophic failure is characterized by a time-accelerating increase in leakage current that is mainly due to existing processing defects (voids, cracks, delaminations, and so on) or to extrinsic defects. A slow degradation failure is characterized by a near-linear increase in leakage current against stress time; this is caused by the electromigration of oxygen vacancies (intrinsic defects). The TTF data shown in Fig. 2 clearly indicate that BME capacitors with slow degradation failures exhibit the largest MTTF values (AA47450), and those with the most catastrophic failures showed the smallest MTTF values (AC47450). Capacitor lot AB47450 shows failures with both failure modes and with MTTF values in between those of the other two capacitor lots.

As shown in Fig. 2, for a certain period of stress time, the leakage current follows a slow degradation failure mode, i.e., a gradual increase in leakage current against stress time. With a further increase of stress time, some capacitors will fail catastrophically, while some will retain the slow degradation failure mode until the failure criterion is reached.

The leakage data shown in Fig. 2 have been curved-fitted with a number of different mathematical functions (power law, exponential, linear, logarithmic, and so on). Although the leakage data shown in Fig. 2 appear to be linear against most of the stress time measured, the curve-fitting results have shown that the exponential form of

$$I = I(t_0) e^{\left(\frac{t - t_0}{\tau_{SD}}\right)}$$  \hspace{1cm} (2)

fits the leakage data better than a linear form. In (2), $I$ is the measured leakage current, $I(t_0)$ is the leakage value at $t = t_0$, and $\tau_{SD}$ is a characteristic exponential growth time.

Fig. 3 shows an example of curve fitting using (2) for two capacitor samples with different failure patterns. Both appear to fit well to the exponential form of (2) for most of the stress time.

The meaning of $\tau_{SD}$ can be illustrated by the following example: let $I_1$ and $I_2$ be the leakages at $t_1$ and $t_2$, respectively, for a slow degradation failure. If one assumes $I_2/I_1 = 2$, then (2) can be rewritten as

$$\frac{I_2}{I_1} = e^{\left(\frac{t_2 - t_1}{\tau_{SD}}\right)} = (\frac{I_2}{I_1})^{\frac{1}{\tau_{SD}}} = 2$$

and

$$\tau_{SD} = \frac{\Delta t}{\ln(2)} \approx 1.4427 \cdot \Delta t$$  \hspace{1cm} (3)

where $\Delta t$ is the time at which the leakage current doubles in value. The greater the value of $\tau_{SD}$, the longer the timespan of a degradation failure, indicating a slower IR degradation process.

At a given stress condition, when the $\tau_{SD}$ of all 20 BME capacitors being tested was obtained using the curve-fitting method described above, a two-parameter Weibull plot could be made, and an average value of $\tau_{SD}$, $\langle \tau_{SD} \rangle$, was defined as the scale parameter $\eta$ of the Weibull plot.

Table III summarizes the values of $\langle \tau_{SD} \rangle$ determined for various stress conditions. The table can be used to compare the corresponding MTTF data. In most cases, the value
the PTCR effect in donor-doped semiconducting BaTiO$_3$ boundary, as shown in Fig. 4, was initially proposed to explain the formation of a double Schottky barrier layer at a grain surface acceptor states whose number is $N_d$. IR Degradation Model for BME Capacitors

the initial leakage current value for the capacitors. then the relationship depends on the value chosen and on the value of leakage current is selected as the failure criterion, Ni–BaTiO$_3$ MLCCs [2], [4], [9]–[11]. The typical barrier has also been suggested to explain the IR degradation in ceramics [13]–[15], the same barrier depletion layer model of $\phi$. Although oxygen vacancies migrate under an applied dc field and the weakly bonded electrons can be trapped by the surface acceptor states, the value of depletion layer thickness $d$ in (4) is always in the submicrometer range, indicating that $N_d \gg n_s$. Therefore, one can assume that $N_d(t) \approx N_d(0)$, and one can also assume time independence. Therefore, the time-dependent barrier height $\phi$ can be expressed as

$$\frac{d\phi(t)}{dt} = \frac{d}{dt} \left( \frac{e^2n_s^2}{8\varepsilon_0\varepsilon_r N_d} \right) = \frac{e^2}{4\varepsilon_0\varepsilon_r N_d} \frac{dn_s(t)}{dt}. $$

To determine $dn_s(t)/dt$, the following facts were considered.

1) $n_s(t)$ is trapped electrons at surface acceptor states in the grain boundary regions. The negative space charge due to trapped electrons is compensated for by the formation of a positive space charge region near the grain boundary, which behaves like a depletion barrier layer to electron conduction.

2) The computational analysis on the trapping of oxygen vacancies at grain boundaries with respect to local atomic configuration and energy shows that grain boundaries attract oxygen vacancies and trap them at specific sites at which local cation density is lower than that in the grain interior [17].

3) Since oxygen vacancies behave like donors, they possess positive space charges when ionized. The same positive space charge in a barrier layer at a grain boundary will thus act as a resistance for positively charged oxygen vacancy diffusion in a polycrystalline BaTiO$_3$ dielectric. As a result, when an ionized oxygen vacancy migrates under a dc field and reaches the barrier layer, it has a tendency to become trapped there. The electroneutrality condition requires that the weakly bonded two electrons that are moving in a conduction band now have to be localized to make the trapped oxygen vacancy neutralize and to become part of the crystalline structure.

When Kröger and Vink [31] symbols are used, the process can be described by

$$V_O = V_O^{**} + 2e'. $$

As reported in [4], the localized electrons that compensate the $V_O^{**}$ localization can be trapped with the reduction of

<table>
<thead>
<tr>
<th>Test Conditions</th>
<th>MTTF (Minutes)</th>
<th>$&lt;\tau_{SD}&gt;$ (Minutes)</th>
</tr>
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<tbody>
<tr>
<td>250V, 175°C</td>
<td>1466</td>
<td>3333</td>
</tr>
<tr>
<td>250V, 165°C</td>
<td>9869</td>
<td>11111</td>
</tr>
<tr>
<td>250V, 155°C</td>
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<tr>
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<tr>
<td>250V, 175°C</td>
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</tr>
<tr>
<td>250V, 165°C</td>
<td>1140</td>
<td>1714</td>
</tr>
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<td>250V, 155°C</td>
<td>3659</td>
<td>3333</td>
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<tr>
<td>Test Conditions</td>
<td></td>
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<tr>
<td>250V, 175°C</td>
<td>319</td>
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</tr>
<tr>
<td>250V, 165°C</td>
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<td>250V, 155°C</td>
<td>1479</td>
<td>1667</td>
</tr>
<tr>
<td>Test Conditions</td>
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</tr>
</tbody>
</table>

where $N_d$ is the donor concentration, $d$ is the depletion layer thickness, $e$ is the electron charge, and $\varepsilon_0\varepsilon_r$ is the dielectric constant. The electroneutrality condition in the depletion layer satisfies the following requirement [13]:

$$d = \frac{n_s}{2N_d} $$

where $n_s$ is the concentration of trapped electrons at grain boundary surface acceptor states (the number is $N_s$) (cm$^{-2}$). The barrier height $\phi$ can be rewritten as

$$\phi = \frac{e^2n_s^2}{8\varepsilon_0\varepsilon_r N_d}. $$

Equation (5) has often been used to estimate the grain boundary barrier height in semiconducting BaTiO$_3$ ceramics [34], [38]. In Ni–BaTiO$_3$ MLCCs, $N_d$ is mainly determined by the bulk concentration of ionized oxygen vacancies. Fig. 4. Schematic of the formation of a double Schottky barrier around the grain boundary of a BaTiO$_3$ ceramic capacitor, where $\phi$ is the depletion barrier height; $n_s$ is the concentration of trapped electrons at the grain boundary surface acceptor states whose number is $N_s$ (cm$^{-2}$); and $E_s$ is the energy level of the surface states from the conduction level of the bulk.

of $<\tau_{SD}>$ was greater than that of MTTF, but it was smaller in a few lower-stress levels where the catastrophic failure mode is dominant. Certainly, the relationship between TTF and $\tau_{SD}$ depends upon the failure criterion, and if a fixed absolute value of leakage current is selected as the failure criterion, then the relationship depends on the value chosen and on the initial leakage current value for the capacitors. B. IR Degradation Model for BME Capacitors

1) Time-Dependent Depletion Layer Height $\phi(t)$: Although the formation of a double Schottky barrier layer at a grain boundary, as shown in Fig. 4, was initially proposed to explain the PTCR effect in donor-doped semiconducting BaTiO$_3$ ceramics [13]–[15], the same barrier depletion layer model has also been suggested to explain the IR degradation in Ni–BaTiO$_3$ MLCCs [2], [4], [9]–[11]. The typical barrier height can be expressed as

$$\phi = \frac{e^2N_d d^2}{2\varepsilon_0\varepsilon_r} $$

where $N_d$ is the donor concentration, $d$ is the depletion layer thickness, $e$ is the electron charge, and $\varepsilon_0\varepsilon_r$ is the dielectric constant. The electroneutrality condition in the depletion layer satisfies the following requirement [13]:

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$$\frac{d\phi(t)}{dt} = \frac{d}{dt} \left( \frac{e^2n_s^2}{8\varepsilon_0\varepsilon_r N_d} \right) = \frac{e^2}{4\varepsilon_0\varepsilon_r N_d} \frac{dn_s(t)}{dt}. $$

To determine $dn_s(t)/dt$, the following facts were considered.

1) $n_s(t)$ is trapped electrons at surface acceptor states in the grain boundary regions. The negative space charge due to trapped electrons is compensated for by the formation of a positive space charge region near the grain boundary, which behaves like a depletion barrier layer to electron conduction.

2) The computational analysis on the trapping of oxygen vacancies at grain boundaries with respect to local atomic configuration and energy shows that grain boundaries attract oxygen vacancies and trap them at specific sites at which local cation density is lower than that in the grain interior [17].

3) Since oxygen vacancies behave like donors, they possess positive space charges when ionized. The same positive space charge in a barrier layer at a grain boundary will thus act as a resistance for positively charged oxygen vacancy diffusion in a polycrystalline BaTiO$_3$ dielectric. As a result, when an ionized oxygen vacancy migrates under a dc field and reaches the barrier layer, it has a tendency to become trapped there. The electroneutrality condition requires that the weakly bonded two electrons that are moving in a conduction band now have to be localized to make the trapped oxygen vacancy neutralize and to become part of the crystalline structure.

When Kröger and Vink [31] symbols are used, the process can be described by

$$V_O = V_O^{**} + 2e'. $$

As reported in [4], the localized electrons that compensate the $V_O^{**}$ localization can be trapped with the reduction of
the remaining trapped electrons in acceptor states can be localized to make the trapped states balanced by the trapped electrons in surface acceptor layer and reduce the barrier height. Since the barrier height is balanced by the trapped electrons in surface acceptor states \( n_s(t) \), the reduction in barrier height will lower the Fermi level at grain boundary, and then will reduce the \( n_s(t) \). If we assume that \( \Delta n_O \) is the electron concentration that has been localized to make the trapped \( V_{O}^{\bullet\bullet} \) neutral, \( \Delta n_O \) should meet the following conditions: at \( t = 0 \), \( \Delta n_O(0) = 0 \), and at \( t \to \infty \), \( \Delta n_O = n_s(0) \), i.e., all trapped electrons at \( t = 0 \) in the surface acceptor states \( n_s(0) \) will eventually be fully compensated by the localized electrons that neutralize the trapped \( V_{O}^{\bullet\bullet} \). However, with a further increase of \( \Delta n_O \) as more \( V_{O}^{\bullet\bullet} \) are trapped and neutralized, the electrically negative feature of \( n_s(t) \) will further retard the localization of electrons and reduce the localization rate of \( \Delta n_O \). Therefore, the change of \( \Delta n_O \) as a function of \( t \) can be expressed by a first-order reaction according to the reaction rate theory [30]

\[
\frac{d\Delta n_O(t)}{dt} = K(t)[n_s(0) - \Delta n_O(t)]
\]

and

\[
\int^{\Delta n_O(t)}_{\Delta n_O(0)} \frac{d\Delta n_O(t)}{\Delta n_O(t) - n_s(0)} = \int^{t}_0 -K(t)dt
\]

where \( K(t) \) is the degradation rate constant and \( n_s(0) - \Delta n_O(t) = n_s(t) \) is the trapped electron concentration at surface acceptor states at time \( t \). If \( \Delta n_O(t) \) is only balanced by \( n_s(t) \) near the Fermi level, \( K(t) = K = K_0 e^{-\frac{E_k}{kT}} \), can be simplified as a time-independent constant in which \( E_k \) is the activation energy that is required for \( V_{O}^{\bullet\bullet} \) in electromigration to be neutralized at a grain boundary region per (6) and \( k \) is the Boltzmann constant. Since \( \Delta n_O(0) = 0 \), (7) finally yields

\[
n_s(0) - \Delta n_O(t) = n_s(0)(1 - e^{-Kt})
\]

and

\[
\Delta n_O(t) = n_s(0)(1 - e^{-Kt})
\]

the remaining trapped electrons in acceptor states can be simply expressed according to (8) as

\[
n_s(0) - \Delta n_O(t) = n_s(0) - n_s(0)(1 - e^{-Kt}) = n_s(0)e^{-Kt}.
\]

Combining (5) and (8) yields a time-dependent barrier height

\[
\phi(t) = \frac{e^2[n_s(0) - \Delta n_O(t)]^2}{8\varepsilon_0\varepsilon_rN_d} = \phi(0)e^{-2Kt}.
\]

This relationship indicates that the barrier height will exponentially decrease with time due to the oxygen vacancy entrapment at grain boundaries.

2) Determination of Degradation Rate Constant \( K \): The measurement of \( I-V \) characteristics of ceramic BaTiO\(_3\) inside the grain interior and at the grain boundary has shown that under an applied field of 100 kV/cm, the current density inside the grain and at the grain boundary can differ by several orders of magnitude. The difference increases significantly as temperature increases [33]. It is the grain boundary that holds the high resistivity of the ceramic BaTiO\(_3\). If all grain boundaries inside a dielectric layer are assumed to have a uniform barrier height \( \phi(t) \), the time-dependent resistivity \( \rho(t) \) of an MLCC can be written as [13]

\[
\rho(t) = \rho_0 e^{\left(\frac{\phi(0)}{kT}\right)}
\]

where \( \rho_0 \) is the resistivity of the grain interior. According to (2), the time-dependent current density of an MLCC \( j(t) \) for a slow degradation failure mode can be expressed as

\[
j(t) = j(t_0)e^{-\frac{t-t_0}{\tau_{SD}}} = \frac{E}{\rho(t)}
\]

or

\[
\rho(t) = \frac{E}{j(t_0)}e^{-\frac{t-t_0}{\tau_{SD}}}
\]

where \( j(t_0) \) is the current density at \( t = t_0 \) and \( E \) is the applied field. Combining (10) and (11) results in

\[
\rho(t) = \rho_0 \exp\left(\frac{\phi(t)}{kT}\right) = \rho_0 \exp\left(\frac{\phi(0)e^{-2Kt}}{kT}\right)
\]

\[
= \frac{E}{j(0)} \exp\left(-\frac{t-t_0}{\tau_{SD}}\right).
\]

At a given stress level, \( E \) is a constant, so

\[
\frac{t-t_0}{\tau_{SD}} \approx -\frac{\phi(0)}{kT}e^{-2Kt}.
\]

Using \( \tau_{SD} \), the average of \( \tau_{SD} \), to replace \( \tau_{SD} \), and \( e^{-x} \approx 1 – x \) when \( x \) is small, the integration of (12) results in

\[
\int^{MTTF}_{0} \frac{t-t_0}{\tau_{SD}} dt = -\int^{MTTF}_{0} \frac{\phi(0)}{kT} \cdot e^{-2Kt} dt
\]

\[
\approx -\int^{MTTF}_{0} \frac{\phi(0)}{kT} (1-2Kt) dt
\]

and

\[
\frac{1}{2\tau_{SD}} \approx \frac{\phi(0)}{kT} \left( K - \frac{1}{MTTF} \right).
\]

This gives rise to

\[
\frac{1}{MTTF} = K - \frac{kT}{2\phi(0)\tau_{SD}} \approx K_0 e^{-\frac{E_k}{kT}}.
\]

Equation (13) is the well-known Prokopowicz–Vaskas equation for MLCCs, when applied voltage is a constant [35]. The degradation rate constant \( K \) can now be determined by an Arrhenius plot using the MTTF data obtained at various temperatures and at a constant voltage.

Using the MTTF data at different temperatures and at a given voltage (250 V) for the three BME capacitor lots, a corresponding Arrhenius plot according to (13) can be plotted, as shown in Fig. 5. Both activation energy \( E_k \) and degradation rate constant \( K_0 \) can now be determined.

Table IV lists the activation energy \( E_k \) and constant \( K_0 \) at three different temperatures for the three BME capacitor lots. The calculated \( K \) values shown in Table IV were used to estimate the MTTF data per (13). A comparison between the measured MTTF data and the calculated data shows fairly good agreement. All calculated MTTF values are smaller than the measured ones. For lot AA47450, the MTTF data
measured at 175 °C were excluded for the estimation of the degradation rate constant $K$ because its value is much smaller when combined with other MTTF data points to give rise to a good linear fitting. Since the BME capacitor lot under degradation was so leaky at this temperature, the results of self-heating due to the leakage current could result in a significant amount of temperature increase, and thus an acceleration in the failure of the BME capacitors.

C. IR Degradation Mechanism Due to Oxygen Vacancy Electromigration

1) Reliability and Oxygen Vacancy Migration: According to (9)

$$\phi(t) = \phi(0)e^{-2Kt} = \phi(0)e^{-2\frac{t}{\text{MTTF}}}$$

and

$$\frac{1}{\text{MTTF}} = \approx K_0e^{-\frac{E_k}{kT}}$$

where $K = K_0e^{-E_k/kT}$ is the degradation rate constant for oxygen vacancy entrapment at grain boundaries. Although $K$ and MTTF can be related to each other per the Prokopowicz–Vaskas equation, the physical meaning of $K$ was clearly defined in the proposed model. To make Ni–BaTiO$_3$ BME capacitors with improved reliability, the value of $K$ must be minimized. This requires a smaller value of the degradation constant $K_0$, which is the total number of oxygen vacancies that are migratable; and a large value of $E_k$, which is the activation energy required for $V_O^{**}$ to be neutralized and trapped near the depletion layer at grain boundaries. This has been proven from the calculated $K$ values shown in Table IV, where AA47450 and AB47450 are shown to have similar values of $E_k$ but a much smaller value for $K_0$, which gives rise to a significantly improved reliability life (larger MTTF) for capacitor AA47450.

On the other hand, since a typical barrier height value of $\phi(0) \approx 1.30$ eV has been reported in [4], a large value of $E_k > \phi(0)$ is necessary to slow down the $V_O^{**}$ entrapment, because a large value of $E_k$ also means that the entrapment of $V_O^{**}$ at grain boundaries may not be an energetically favorable process unless the barrier height $\phi(t)$ is high enough to be comparable to $E_k$. This is the case for AC47450, where $E_k = 1.11$ eV is smaller than that of $\phi(0)$. Therefore, the oxygen vacancies are more energetically favorable for local entrapment than for migrating across the depletion barrier, and they result in a very small reliability life when compared with the other two BME capacitor lots.

Finally, since $N_d \gg n_e$ per (4), only a tiny fraction of oxygen vacancies can be trapped at the grain boundaries during the electromigration across the dielectric layer to cause IR degradation. This indicates that the effort to simply reduce the level of $V_O^{**}$ in the dielectric material would not significantly slow down the IR degradation. Indeed, a small value of $K_0$ indicates that the number of electromigrateable $V_O^{**}$ must be minimized. The recent first-principles calculations based on the density function theory have shown that some kinds of dopants in BaTiO$_3$ dielectrics, such as RE and alkaline-earth (AE) elements, were effective in improving the reliability life of Ni–BaTiO$_3$ capacitors [17]–[19]. According to the IR degradation model proposed here, both RE and AE doping would benefit from the frozen migrateable $V_O^{**}$, and thus reduce the value of $K_0$. On the other hand, some RE doping elements, such as Dy and Er, due to their amphoteric characteristics, will also benefit from the formation of deep surface acceptable states $N_S$ to result in a large value of $E_k$.

The correlations between the formulation and the value of the degradation rate constant for oxygen vacancy entrapment $K$ as defined in this paper needs to be further investigated.

2) Insulation Resistance and Reliability Life: According to (12)

$$\frac{L}{\text{MTTF}} = \approx e^{-\frac{\phi(0) - \phi(t)}{kT}}e^{-2Kt}$$

which indicates that a slower IR degradation process, characterized by a larger value of $r_{SD}$, would give rise to a smaller value of $\phi(0)$. This makes sense since a smaller $\phi(0)$ will energetically be favorable to the continuous electromigration of $V_O^{**}$ without being trapped at a grain boundary to cause an IR degradation. However, this is only one part of the equation; $\phi(0)$ also presents the barrier height for the conduction band electronic carriers. A lower $\phi(0)$ will facilitate electronic conduction and will also deteriorate the IR. As a result, when electron conduction and oxygen vacancy electromigration are both considered, a moderate initial barrier height $\phi(0)$ and a smaller $K$ are the keys for slowing the IR degradation in Ni–BaTiO$_3$-based ceramic capacitors, and thus improving their reliability life.

As an important conclusion of this model development, higher IR values may not always result in a larger MTTF, but a slower IR degradation rate (smaller $K$) will always have this effect. This conclusion can be verified from the measured leakage data shown in Fig. 3, where sample AC47450 was shown to have the highest initial IR values and smallest MTTF and AA47450 had the lowest initial IR values but the largest MTTF among the group. A higher $\phi(0)$ generally means a higher resistance, and therefore a higher electrical strength when a dc voltage is applied. A higher dc electrical strength makes it more likely...
that a thermal-related electrical breakdown will be experienced due to the localization of the electrical strength [9]–[11].

3) Oxygen Vacancy Migration and Compensation: Since only a very small portion of $V_{O}^{**}$ may be trapped at the grain boundaries, the majority of $V_{O}^{**}$ will continually migrate and will eventually reach the dielectric layer and internal Ni electrode interface, as has been shown by previously reported electron energy loss spectroscopy and high-resolution transmission electron microscope observations [4]. Since there is no evidence to show that $V_{O}^{**}$ can be transferred across the cathode electrode layer [4], [10] most $V_{O}^{**}$ capable of migration will now pile up along the Ni-electrode dielectric interface. To neutralize these vacancies, a significant number of electrons is required, which can only be obtained from the cathode electron injection. The energy required for cathode electron injection at the dielectric-electrode interface is $\sim 1.25$ eV [36]. If $E_k$ is less than this, most of the oxygen vacancies will be energetically favorable for localization, and a quick IR degradation will occur. This is exactly the case for lot AC47450.

The high concentration of localized electrons due to the compensation of the pile-up of oxygen vacancies will not only dramatically change the local stoichiometry of the BaTiO$_3$ dielectric but it will also lead to a leakage current increase during IR degradation. This will cause a local temperature increase and will eventually lead to the breakdown at the Ni–BaTiO$_3$ interface. The initial failure site of the dielectric-electrode interface was clearly revealed in a previously published failure analysis work regarding Ni–BaTiO$_3$ ceramic capacitors [37].

A model based on the existence of depletion layers at grain boundaries and on the entrapment of electromigrateable oxygen vacancies was proposed to explain the MTTF difference among these BME capacitors. The MTTF has been found to be directly related to the degradation rate constant $K$ of oxygen vacancy entrapment at grain boundaries. The MTTF and $K$ were found to follow the traditional Prokopowicz–Vaskas equation at a constant applied voltage. A lower depletion layer height $\phi(0)$ is energetically favorable for a slower degradation rate and a longer reliability life. However, when both oxygen vacancy migration and electronic conduction are considered, a $\phi(0)$ with a moderate height would give rise to the best reliability performance.

It is the conclusion of this paper that reliability will not be improved simply by increasing the IR. Indeed, Ni–BaTiO$_3$ BME capacitors with a smaller IR degradation rate constant $K$, or a smaller amount of entrapped $V_{O}^{**}$ at grain boundaries, will always give rise to a longer reliability life.

Both RE and AE doping may have profound impacts on the values of degradation rate constant $K$, and the impacts needs to be further investigated.

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REFERENCES


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