Degradation of Leakage Currents and Reliability Prediction for Tantalum Capacitors

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Key Words: acceleration factors, degradation, failures, leakage current, tantalum capacitors

SUMMARY & CONCLUSIONS
Two types of failures in solid tantalum capacitors, catastrophic and parametric, and their mechanisms are described. Analysis of voltage and temperature reliability acceleration factors reported in literature shows a wide spread of results and requires more investigation.

In this work, leakage currents in two types of chip tantalum capacitors were monitored during highly accelerated life testing (HALT) at different temperatures and voltages.

Distributions of degradation rates were approximated using a general log-linear Weibull model and yielded voltage acceleration constants \( B = 9.8 \pm 0.5 \) and \( 5.5 \). The activation energies were \( E_a = 1.65 \) eV and \( 1.42 \) eV. The model allows for conservative estimations of times to failure and was validated by long-term life test data.

Parametric degradation and failures are reversible and can be annealed at high temperatures. The process is attributed to migration of charged oxygen vacancies that reduce the barrier height at the MnO2/Ta2O5 interface and increase injection of electrons from the MnO2 cathode. Analysis showed that the activation energy of the vacancies’ migration is \( \sim 1.1 \) eV.

I INTRODUCTION
Two types of failures are known for solid chip tantalum capacitors with manganese oxide cathodes operating at steady state conditions. Type I appears as a sharp rise of direct current leakage (DCL) indicating breakdown of the dielectric, and type II as a gradual increase of currents with time resulting in parametric failures.

 Failures in tantalum capacitors are often explained by field-induced crystallization of the amorphous anodic tantalum pentoxide dielectric [1, 2]. These crystals gradually grow with time of operation under the amorphous Ta2O5 layer and eventually disrupt the dielectric.

Type I failures can be explained based on the time dependent dielectric breakdown (TDDDB) model [3]. According to the thermochemical model, the breakdown occurs when the local electrical field weakens polar molecular bonds with time to the level at which thermal energy is sufficient to cause the breakage.

Type II failures appear as a gradual degradation of DCL with time that might eventually exceed the specified limit. Experiments showed that in some capacitors currents might increase with time by several orders of magnitude and result eventually in a thermal runaway and catastrophic failures. The process was explained by migration of positively charged oxygen vacancies in the Ta2O5 dielectric [4, 5].

The voltage acceleration factor for reliability calculations that is determined as a ratio of failure rates at the test and rated (VR) voltages, can be written in the form:

\[
AF_V = \exp \left[ B \times \left( \frac{V_{\text{test}}}{V_R} - 1 \right) \right],
\]

where \( B \) is the voltage acceleration constant.

The temperature acceleration factor is determined as a ratio of failure rates measured at temperatures \( T_1 \) and \( T_2 \) and is described by the Arrhenius equation:

\[
AF_T = \exp \left[ \frac{E_a}{k} \times \left( \frac{1}{T_1} - \frac{1}{T_2} \right) \right],
\]

where \( k = 8.617 \times 10^{-5} \) eV/K is the Boltzmann constant, and \( E_a \) is the activation energy.

Only scarce information was found in the literature about reliability acceleration factors for tantalum capacitors, and most of the existing data are related to catastrophic, type I, failures. Table 1 shows values of \( B \) and \( E_a \) for type I and type II failures estimated based on technical publications. Constant \( B \) for type I failures varies in a wide range, from 6 to 20.6 and activation energies are in the range from 0.7 eV to 2.1 eV. At these ranges, a 20% change of operating voltage, would cause variation of \( AF_V \) by more than an order of magnitude, from 3.3 to 61.5. Temperature increase from a typical operating conditions of 55 °C, to test conditions at 125 °C, would vary \( AF_T \) from 78 to 4.7x10\(^5\).

Assessment of failure rates in military grade tantalum capacitors is based on the assumption that the constant \( B \) is the same for all part types, \( B = 18.77 \). However, analysis [6] showed that this assumption can cause errors in FR up to three orders of magnitude because actual values of \( B \) can vary substantially from lot to lot. This requires development of express methods to assess acceleration factors for different types of capacitors. One such method that has been suggested in [3] is based on measurements of breakdown voltages and the TDDDB model. Another express method is based on analysis of times to failures (TTF) observed during life step stress testing [7].

These methods were used for type I, or catastrophic failures that are typically detected by blown fuses connected in series with each capacitor under test during burning-in or so called...
Weibull grading testing. Type II failures are often observed during qualification life tests that are carried out at 125 °C and require periodic measurements of leakage currents during the test.

This work was undertaken to evaluate acceleration factors for reliability testing based on the rate of degradation of leakage currents.

<table>
<thead>
<tr>
<th>Type</th>
<th>B</th>
<th>$E_a$, eV</th>
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<th>Ref.</th>
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<td>9.3</td>
<td>1.2</td>
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<td>[8]</td>
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<td>Rate of annealing and TSD</td>
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Table 1. Literature data for voltage acceleration constants and activation energies for type I and type II failures.

2 TECHNIQUE

Two lots of molded case chip 6.8 μF tantalum capacitors rated to 25 V and 35 V were selected for this study based on preliminary tests showing that capacitors from these lots have substantial degradation of leakage currents at high temperatures. Twenty capacitors of each type were used for the HALT with monitoring of leakage currents at different temperatures and voltages. Each test consisted of 30 hours stress under bias followed by depolarization ($V = 0$) at the same temperature for 3 hours. The testing was carried out at room temperature, 85 °C, 105 °C, 125 °C and 145 °C. For capacitors rated to 25 V, stress voltages increased from 15 V to 25 V, and finally to 35 V. The currents were monitored by scanning voltages across 100 kohm resistors connected in series with each capacitor.

Typical $I-t$ characteristics observed during the testing are shown in Figure 1. A general trend for DCL variations is an initial decrease followed by increasing current degradation. The initial decrease is due to absorption currents that follow the empirical Curie - von Schweidler power law, $I \sim t^n$, with the exponent $n$ close to 1. These currents might be substantially larger than the intrinsic leakage currents even at 125 °C.

A portion of the $I-t$ curve with increasing currents was approximated with a linear function:

$$I(t) = I_0 + \alpha \times t$$

(1)

where $I_0$ is the initial current, and $\alpha$ is the rate of degradation.

Figure 1. Variations of leakage currents with time at 105°C, 25V. Insert shows the same chart in double logarithmic coordinates.

Rates of current degradation were calculated for each part at each test condition and distributions of $\alpha$ were approximated with Weibull law:

$$F(\alpha) = 1 - \exp \left[ -\left( \frac{\alpha}{\eta} \right) ^{\beta} \right] .$$

(4)

where $\eta > 0$ is the characteristic degradation rate (scale parameter), and $\beta > 0$ is the shape parameter (slope).

The values of $\beta$ and $\eta$ were determined using a general log-linear model at a reciprocal transformation of absolute temperature. ALTA-7 software available from ReliaSoft was used to calculate parameters of the model based on the maximum likelihood estimation (MLE) method. According to this method, all distributions are assumed to have the same slopes of Weibull distributions, and the scale parameter depends on temperature and voltage as:

$$\eta(T, V) = a_0 \times \exp \left( \frac{a_1}{T} \right) \times \exp \left( a_2 \times V \frac{T}{T_a} \right) ,$$

(5)

where $a_i$ are parameters of the model, and $T$ is the absolute temperature.

Based on the model’s parameters, the voltage acceleration constant $B$ and the activation energy of degradation, $E_a$, can be calculated: $B = a_2 \times VR$ and $E_a = -a_1 / k$.

3 TEST RESULTS

An example of experimental data obtained during leakage current monitoring for a capacitor rated to 35 V at 85 °C and different voltages and their linear approximations are shown in Figure 2.

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The rate of degradation increased more than three orders of magnitude with voltage, from ~7E-12 A/sec at 56 V to more than 1E-8 A/sec at 77 V. Eventually, the part failed after ~30 hours of testing at 77 V. The rate of degradation increased exponentially with voltage, and calculations for this sample yielded voltage acceleration constant \( B = 12.4 \).

Degradation of currents with time at 125 °C and different voltages for 6.8 µF, 25 V capacitors is shown in Figure 3. Currents increased linearly with time during the whole period of testing at 25 V, but showed a trend for saturation at 35 V. A similar trend was also observed at 145 °C and 25 V.

Values of \( \alpha \) increase with \( I_{min} \) according to a power law with the exponent ~1.4.

After testing of capacitors rated to 25 V at 125 °C and 35 V, leakage currents at 85 °C and 25 V were monitored for 10 hours twice: immediately after the testing and then after baking at 175 °C for 20 hours. No parametric failures were observed during the initial 85 °C testing, but all parts failed DCL requirements (1.7 µA) when the measurements were made after HALT. Baking resulted in a substantial reduction of currents, and ~90% of capacitors were within the specified limits after the bake. The baking cycle that included measurements of currents at 85 °C 25 V right after HALT, annealing, and post-bake measurements was repeated after the last 30-hours test at 25 V and 145 °C. To evaluate the effect of annealing duration, the 85 °C measurements were carried out after 10, 30, and 60 hours of bake at 175 °C. Variations of currents measured at 25 V and 85 °C through the testing are shown in Figure 5 and indicate reproducible variations of leakage currents in the HALT-bake cycles. Leakage currents decrease with the time of baking and a 60-hour bake at 175 °C can completely restore the values of DCL observed before the testing even for samples that had currents degraded by almost three orders of magnitude.

Degradation rates calculated for 25 V capacitors at different temperatures are plotted against the minimal values of DCL, \( I_{min} \), in Figure 4. The results indicate a good correlation between these parameters and the possibility of selecting the most stable capacitors based on the level of initial currents. The

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high rates is of the most interest. To simplify calculations in the following analysis, we consider samples in the slow-degrading subgroup (below 35% of the distribution) and high-degrading subgroup (above 35% of the distribution) separately.

Distributions of $\alpha$ for 6.8 $\mu$F 35 V capacitors were unimodal. These parts were tested similar to 6.8 $\mu$F 25 V capacitors, but at voltages from 49 V to 77 V with 7 V increments at 85 °C, and at 125 °C the measurements were made at 42 V only. Experimental data and results of modeling of HALT are shown in Weibull coordinates in Figure 7.

Similar distributions were obtained for 6.8 $\mu$F 25 V capacitors separately, for the fast-degrading and slow-degrading subgroups. Results of analysis using the log-linear model, Equation (5), as well as voltage acceleration constants and activation energies are displayed in Table 2. Constants $B$ and $E_a$ are similar for the slow-degrading subgroup of capacitors rated to 25 V and to 35 V capacitors ($B_{av} = 9.8 \pm 0.5$ and $E_{aav} = 1.65 \pm 0.1 \text{ eV}$). However, both constants are smaller (~45% for $B$ and ~20% for $E_a$) for the fast degrading subgroup.

Results of life testing for six samples of 6.8 $\mu$F 25 V capacitors tested at 125 °C and 16.6 V for 2000 hours are presented at Figure 8. Two out of six samples failed marginally, one after 1000 hrs and another after 2000 hrs. Apparently, the currents are stabilizing after ~1500 hrs, so it is quite possible that other parts in this group would not fail if testing continued.

![Degradation rate, 6.8uF 25V at 25V](image)

Figure 6. Distributions of degradation rates in 6.8 $\mu$F 25 V capacitors at 25 V and temperatures from 85 °C to 145 °C.

Distributions of $\alpha$ for 6.8 $\mu$F 35 V capacitors were unimodal. These parts were tested similar to 6.8 $\mu$F 25 V capacitors, but at voltages from 49 V to 77 V with 7 V increments at 85 °C, and at 125 °C the measurements were made at 42 V only. Experimental data and results of modeling of HALT are shown in Weibull coordinates in Figure 7.

![Degradation rates for 6.8uF 35V capacitors](image)

Figure 7. Distributions of degradation rates during HALT for 6.8 $\mu$F, 25 V capacitors at different temperatures and voltages. Marks correspond to experimental data and lines are approximation with the log-linear model. A dashed line corresponds to the “use life”, 55 °C, 10 V, conditions.

To validate the model, degradation rates at the life test conditions (125 °C, 16.6 V) were calculated for the slow-degrading and fast degrading subgroups for 6.8 $\mu$F 25 V capacitors, and times to failure were determined for each capacitor based on the linear approximation:

$$ TTF = \frac{I_{crit} - I_0}{\alpha} $$

where $I_{crit} = 40.8 \mu$A is the critical current at life test conditions.

![Degradation rate, 6.8uF 25V at 25V](image)

Figure 8. Variations of current with time for 6 samples of 6.8 $\mu$F 25 V capacitors during life testing.

A comparison of times to failure calculated based on the model and experimental data obtained during life testing is shown in Fig.9. The calculated and experimental TTF data are close thus validating the model. However, calculations predict smaller times to failure, likely due to the saturation of currents and decreasing rate of degradation with time that was neglected in the model. The slope of the TTF distribution for the fast-degrading subgroup of capacitors is 1.96, which indicates wear-out type of failures.

The model was also used to assess TTF at the use conditions, 55 °C and 10 V. The relevant distribution is shown in Figure 9 and indicates that the probability of parametric failures is less than 1% after ~800 years of operation. The results show a substantial, more than 5 orders of magnitude increase in the life expectancy as temperature and voltage decrease. The acceleration factor of life testing compared to the operating conditions was calculated based on Table 1 data: $AF = A_{F_1} \times A_{F_T} = 36 \times 6800 = 2.4 \times 10^5$. For these two conditions

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<thead>
<tr>
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<th>High-rate</th>
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<td>0.99</td>
<td>1.95</td>
</tr>
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<td>12.59</td>
<td>12.53</td>
<td>12.29</td>
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<td>-1.9300</td>
<td>-1.6400</td>
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</tr>
<tr>
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<td>0.218</td>
<td>0.294</td>
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</tr>
<tr>
<td>$E_a$, eV</td>
<td>1.66</td>
<td>1.42</td>
<td>1.65</td>
</tr>
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</table>

Table 2. Parameters of log-linear model.
the acceleration caused by temperature is almost two orders of magnitude greater than by voltage.

It is possible to show that in this approximation, the degradation rate depends on the initial intrinsic current, $I_0(T)$, and mobility of oxygen vacancies:

$$\alpha(T) = I_0(T) \times \frac{\beta_s}{kT} \times \left( \frac{V_{\text{test}}}{d} \right)^{0.5} \times q \times \mu(T) \times N_v \times \frac{d}{2\alpha^2_0} , \quad (9)$$

Both parameters in this equation, $I_0$ and $\mu$, are increasing with temperature according to the Arrhenius law, hence the activation energy of the degradation rate is a sum of activation energies of leakage currents, $E_d$ and $V_o^{++}$ migration, $E_v$:

$$E_d = E_i + E_v \quad , \quad (10)$$

Results of this work indicate $E_d \sim 1.65$ eV, and based on the temperature dependence of $I_{\text{min}}$ for both types of capacitors is in the range from 0.5 to 0.6 eV, so $E_v$ is $\sim 1.1$ eV. The latter value is close to the range of values reported previously. Based on the rate of annealing and measurements of the thermally stimulated depolarization (TSD), the activation energy for mobility of $V_o^{++}$ was 1.1 to 1.2 eV [5, 17]. Analysis of local atomic structure before and after annealing in thin (~100 nm) amorphous Ta2O5 layers showed that activation energy of oxygen diffusion via vacancies is in the range from 1 to 1.3 eV [19]. Interestingly, $E_v$ for diffusion of oxygen ions in ceramic capacitors has similar values, from 0.9 to 1.1 eV [20].

The range of activation energies observed in this work for type II failures is overlapped by values reported for type I failures. Although more data are necessary to reveal possible differences, both types of failures are associated with ionic processes, and thus might have similar activation energies and acceleration factors.

The suggested mechanism can explain a correlation observed between the minimal, or initial, leakage currents and the rate of degradation (Figure 4). Capacitors with higher concentration of $V_o^{++}$ would have a greater degradation rate and higher levels of initial leakage currents because the stresses that parts experienced during manufacturing testing (e.g. burning-in) could cause a certain level of degradation before HALT. It is also possible that oxygen vacancies being donors for electrons would increase conductivity of the dielectric and greater DCL.

The presence of low- and fast-degrading subgroups in the population of 6.8 µF 25 V capacitors might be due to a poor reproducibility of the anodic oxidation process that causes variations in conditions at the surface of the tantalum pellet after sintering. Similar bimodal distributions of TTF for metal-insulator-metal (MIM) capacitors with deposited Ta2O5 dielectric was observed in [8, 21]. The results were explained by the presence of two failure mechanisms: thermal runaway for capacitors having low TTF, and intrinsic wear-out for capacitor failing at times that are more than two orders of magnitude greater.

Our data show that the fast degrading capacitors failed parametrically, without thermal runaway. Due to saturation of leakage currents with time, there might be no catastrophic failures if testing continued. More than that, samples in the slow degrading subgroup might not fail even parametrically. However, this does not exclude the possibility that another

Figure 9. Distributions of the calculated and experimental times to failure. Experimental data correspond to results in Figure 8, and “calculation” is based on the results of HALT and log-linear model.

5 DISCUSSION

The reversibility of degradation of leakage currents that was also observed earlier [17] precludes the possibility of parametric failures caused by the field-induced crystallization of the Ta2O5 dielectric. The most probable cause of degradation is migration of positively charged oxygen vacancies, $V_o^{++}$. Under applied bias, oxygen vacancies move to the cathode, reduce the barrier at the MnO2/Ta2O5 interface, and increase injection of electrons into the dielectric by the Schottky mechanism. High-temperature annealing causes redistribution of $V_o^{++}$, thus increasing the barrier height, and decreasing leakage currents.

A variation of the normalized leakage currents with time during the degradation process depends on the effective surface charge density $Q_s(t)$ at the dielectric/cathode interface and can be written as [5]:

$$I(t) = I_0 \times \exp \left[ \frac{\beta_s}{kT} \left( \frac{V_{\text{test}}}{d} + \frac{Q_s(t)}{\alpha_0} \right)^{0.5} - \left( \frac{V_{\text{test}}}{d} \right)^{0.5} \right] , \quad (7)$$

where $\beta_s = 2.71 \times 10^{14}$ Cs/mV is the Schottky constant, $\varepsilon = 5$ is the high frequency dielectric constant, $\alpha_0 = 8.85 \times 10^{-12}$ F/m is the dielectric constant of vacuum, and $d$ is the thickness of Ta2O5 dielectric that for 25 V capacitors is $\sim 150$ nm.

Considering that during HALT at 125 °C and 25 V the ratio $I_{\text{max}}/I_0$ is $\sim 10^3$, calculations per Eq.(7) yielded $Q_s \sim 0.01$ C/m². Assuming that this charge was initially distributed evenly through the thickness of oxide, the concentration of $V_o^{++}$ should be $\sim 4.2 \times 10^{17}$ cm⁻³ minimum. The actual concentration of $V_o^{++}$, $N_v$, is likely at least an order of magnitude greater, which corresponds to literature data [18].

In a first approximation, we can assume that at the beginning of the degradation process, the surface charge increases with time linearly:

$$Q_s(t) = q \times \mu \times E \times N_v \times t \quad , \quad (8)$$

where $q = 1.6 \times 10^{-19}$ C is the electron charge, $\mu$ is the mobility of oxygen vacancies, and $E = V_{\text{test}}/d$ is the electric field.
mechanism, e.g. TDDB or field induced crystallization, would cause failures if duration of testing is increased substantially. A reason for decreasing of the degradation rate with time might be a limited concentration of $V_{o}^{++}$ in the oxide, or compensation of the positive charge by electrons or oxygen atoms injected from the MnO$_2$ cathode layer [22].

REFERENCES


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