MODELING OF THERMAL JOINT RESISTANCE OF POLYMER-METAL ROUGH INTERFACES

M. Bahrami¹, M. M. Yovanovich², and E. E. Marotta ³

Abstract

A compact analytical model is developed for predicting thermal joint resistance of rough polymer-metal interfaces in a vacuum. The joint resistance includes two components: i) bulk resistance of the polymer and ii) micro, constriction/spreading resistance of the microcontacts at the interface. Performing a deformation analysis, it is shown that the deformation mode of the polymer asperities is plastic. The required input parameters of the model can be measured in the laboratory and/or found in the open literature. It is observed that the thermophysical properties of the polymer control the thermal joint resistance and the metallic body properties have a second order effect on the thermal joint resistance. A non-dimensional parameter, i.e., ratio of microcontacts over bulk thermal resistances, is proposed as a criterion to specify the relative importance of the microcontacts thermal resistance. The present model is compared with more than 140 experimental data points collected for a selected number of polymers and showed good agreement.

Nomenclature

\[ A = \text{area, } m^2 \]
\[ a_s = \text{radius of microcontacts, } m \]
\[ b_L = \text{specimen radius, } m \]
\[ E = \text{Young’s modulus, } Pa \]
\[ E' = \text{effective elastic modulus, } Pa \]
\[ F = \text{applied load, } N \]
\[ H_{mic} = \text{microhardness, } Pa \]
\[ H_e = \text{elastic microhardness, } Pa \]
\[ h = \text{thermal conductance, } W/m^2K \]
\[ k = \text{thermal conductivity, } W/mK \]
\[ k^* = \text{non-dimensional thermal conductivity, } \equiv k_p/k_s \]
\[ m = \text{combined mean absolute surface slope, } [-] \]
\[ n_s = \text{number of microcontacts} \]
\[ P = \text{apparent contact pressure, } Pa \]
\[ P^* = \text{non-dimensional pressure, } \equiv P/H_{mic} \]
\[ Q = \text{heat flow rate, } W \]
\[ R = \text{thermal resistance, } K/W \]
\[ T = \text{temperature, } K \]
\[ t = \text{thickness of polymer specimen, } m \]
\[ TCR = \text{thermal contact resistance} \]
\[ TIM = \text{thermal interface material} \]
\[ Y = \text{mean surface plane separation, } m \]
\[ \Theta = \text{non-dimensional parameter } \equiv \frac{R_s}{R_b} \]
\[ \nu = \text{Poisson’s ratio} \]

\[ \gamma = \text{plasticity index } \equiv \frac{H_{mic}}{E'm} \]
\[ \lambda = \text{non-dimensional separation } \equiv \frac{Y}{\sqrt{2}\sigma} \]
\[ \sigma = \text{combined RMS surface roughness, } m \]

Subscripts

\[ 0 = \text{reference value} \]
\[ 1, 2 = \text{solid 1, 2} \]
\[ a = \text{apparent} \]
\[ b = \text{bulk} \]
\[ c = \text{contact} \]
\[ e = \text{elastic} \]
\[ FM = \text{Fuller Marotta} \]
\[ g = \text{glass temperature} \]
\[ j = \text{joint} \]
\[ mic = \text{micro} \]
\[ p = \text{plastic} \]
\[ r = \text{real} \]
\[ s = \text{solid, micro} \]

¹Ph.D. Candidate. Department of Mechanical Engineering, University of Waterloo, Waterloo, ON, Canada N2L 3G1.
²Distinguished Professor Emeritus. Department of Mechanical Engineering, University of Waterloo, Waterloo, ON, Canada N2L 3G1. Fellow ASME.
³TEES Associate Research Professor. Department of Mechanical Engineering, Texas A & M University, College Station, Texas 77813-3123, USA.
1 INTRODUCTION

The continued growth in performance and functionality of microelectronic and avionic systems has resulted in a significant increase in heat dissipation rates and presents a great challenge to thermal engineers. A number of failure mechanisms in electronic devices, such as inter-metallic growth, metal migration, and void formation are related to thermal effects. Following the Arrhenius law, the rate of these failures is approximately doubled with every 10 °C increase above 80 °C in the operating temperature of the device [1]. The heat generated must pass through a complex network of thermal resistances to dissipate from the junction to the surroundings. The most significant resistance is the thermal contact resistance (TCR) at the interface between the package and its heat sink or heat pipe. TCR may be reduced by two methods: 1) increasing the real contact area, accomplished by a) increasing contact pressure, or b) reducing the roughness and out-of-flatness of the contacting surfaces before the interface is formed; and 2) using a thermal interface material (TIM) of higher thermal conductivity that can conform to the imperfect surface features of the mating surfaces. Load constraints on electronic components make it unfeasible to use high contact pressure. Also, manufacturing highly finished surfaces is not practical due to cost constraints. Therefore, the practical alternative is to use a TIM applied at a moderate contact pressure. Most TIMs are polymeric materials filled with thermally conductive particles. Therefore, TCR of the metal-polymer interface is an important issue in microelectronics and chip cooling.

In addition to microelectronics, the use of polymers in everyday items is rapidly increasing in a wide variety of applications. Polyethylene is commercially used for packing films and wire insulation. Polyvinyl chloride (PVC) is used in automobile roofs, pipes, valves, and fittings. Polypropylene is incorporated into military hardware and communication equipment. Teflon is used as a seal, valve, nonstick coating, and within electronic equipment. Acrylonitrile-butadiene-styrene (ABS) has commercial use for electronic housings, nylon within electrical and electronic equipment. Delrin has a potential to replace metal in mechanical and structural applications. Polycarbonate is used for machine parts and propellers and phenolic is incorporated in many types of equipment [2].

2 PROBLEM STATEMENT

Figure 1 illustrates the geometry of a polymer-metal joint in a vacuum which has been used in the existing experimental investigations. A polymer specimen of thickness \( t \) is sandwiched between two cylindrical rough nominally flat metal specimens of radius \( b_L \). Thermocouples are mounted in metal flux meters so the heat transfer rate \( Q \) can be determined. Also, metal interface temperatures are estimated by extrapolating the temperature profiles in solids 1 and 2. Three thermal resistances exist in the described joint, i.e., the contact resistance at interface 1, the bulk resistance of the polymer, and the contact resistance at interface 2. To reduce the overall mean temperature and also considering that the TCR is identical at both interfaces 1 and 2, thermal paste is applied at one of the interfaces, therefore the TCR of that interface becomes very small and can be neglected.

Since the contact is in a vacuum, heat is assumed to be transferred, at the metal-polymer interface, only by conduction through microcontacts. The conduction/spreading resistances at microcontacts \( R_s \) is defined as

\[
R_s = \frac{\Delta T}{Q}
\]

where \( \Delta T \) is the temperature drop at the polymer metal interface and \( Q \) is the heat transferred. Heat transfer via radiation across the joint can be neglected as long as the mean temperature of contacting surfaces are not too high, i.e., less than 700 K [3]. In addition, heat flow must overcome the bulk resistance of the polymer specimen \( R_b \) that can be calculated as

\[
R_b = \frac{t}{A_b k_p}
\]

where \( t \), \( A_a = \pi b_L^2 \), and \( k_p \) are the thickness of the polymer at the applied load, the apparent contact area, and thermal conductivity of the polymer, respectively. As shown in Fig. 1, the contact resistance \( R_c \) and the bulk resistance \( R_b \) are in series, therefore the joint resistance can be determined from

\[
R_j = R_c + R_b
\]
thermal conductivities and the thermal joint resistance of several thermoplastic and thermoset polymers over a range of temperatures and contact pressures. Thermal conductivity of the polymers reported in [2] showed small variation as the temperature was changed over the range of 10 to 100°C. Marotta and Fletcher [2] compared their measured thermal joint resistance data $R_j$ with the microcontacts resistance $R_s$ calculated from the Mikic elastic model [6] and the Cooper et al. plastic model [7]. It was shown that both models failed to predict the trend of the data [2]. The comparisons of Marotta and Fletcher [2] were not appropriate because their measured joint resistance, which include the microcontacts and the bulk resistances ($R_s + R_b$) were compared only against the microcontacts resistance $R_s$ predicted by the models. Also, they did not offer any model to predict the joint resistance.

Fuller and Marotta [8] conducted experiments in a vacuum with several polymers and developed an elastic contact model for predicting the polymer-metal joint resistance. Their model accounted for the bulk resistance and the TCR of the joint, i.e., Eq. (3). The TCR component of Fuller and Marotta model was based on the assumption that the deformation of asperities was elastic. They showed good agreement with their experimental data. However, assuming elastic deformation of microcontacts may lead to physically impossible effective elastic microhardness for polymers, discussed in section 4.3.

The thermal contact resistance theory is based on the following premises: 1) the heat flow passes through the contact plane, thus the heat flow direction is perpendicular to the contact plane and 2) the equivalent contact simplification, see section 4.2 and Fig. 2. The contact plane or the apparent area is the projection of contacting surfaces on the plane normal to the direction of the applied load, thus the real contact area is always less than, or at its limit, equal to the apparent area. Fuller and Marotta [8] and Parihar and Wright [5] stated that “since polymers have comparatively lower modulus of elasticity, the calculated real contact area can be greater than the apparent area.” This premise is not correct.

The preceding shows the need for developing model(s) that enables one to predict the TCR of a polymer-metal interfaces. The objective of this study is to develop a compact analytical model for the thermal joint resistance of rough polymer-metal contacts in a vacuum.

4 THEORETICAL BACKGROUND

Real, or engineering surfaces, have roughness. If the asperities of a surface are isotropic and randomly distributed over the surface, the surface is called Gaussian. Due to the random nature of roughness, the microcontacts are distributed randomly in the apparent contact area. The real contact area $A_r$, the summation of these microcontacts, forms a small portion of the nominal contact area, typically a few percent of the nominal contact area [9]. The
The combined roughness perfectly smooth surface, Fig. 2, for more detail see [10]. The combined surface characteristics of the two surfaces with a contact between a single Gaussian surface that has the contact between two Gaussian rough surfaces is modeled by assuming hemispherical asperities whose height and surface slope have Gaussian distributions. The CMY model was equivalent surface approximation, derived relationships for (CMY) [7], based on the level-crossing theory and using the equivalent surface. The plastic deformation of the polymer and the applied load exists. As a result, Hooke’s law can be used to determine the bulk elastic deformation of the polymer specimen shown in Fig. 1 as

\[ \varepsilon = \frac{\Delta t}{t_0} = \frac{P}{E_p} \]  

(5)

where \( \Delta t = t_0 - t \), \( E_p \), and \( P \) are the bulk deformation, elastic modulus, and the nominal contact pressure, respectively.

### 4.1 Polymer Microhardness

Microhardness of polymers can be measured by static penetration of the specimen with a standard indenter at a known pressure similar to metals [11]. The strain boundaries for plastic deformation below the indenter are critically dependent on microstructural factors e.g. crystal size and perfection, degree of crystallinity, etc. Indentation during a microhardness test permanently deforms only a small volume element, in the order of \( 10^9 \) to \( 10^{11} \) nm\(^3\) for a non-destructive test [11]. Thus the contact stress between the indenter and the specimen is much larger than the compressive yield stress of the specimen. The material under the indenter consists of a zone of severe plastic deformation, about 4 to 5 times the penetration depth of the indenter into the specimen, surrounded by a larger zone of elastic deformation. Together these zones generate stresses that supports the force exerted by the indenter.

### 4.2 Thermal Resistance of Microcontacts

For temperatures less than the glassy temperature, a polymer’s mechanical response, both macro and micro, is similar to metals. Therefore, one can apply existing TCR models, originally developed for metals, to determine the joint resistance of polymers. TCR models can be categorized into two main groups: plastic and elastic. The fundamental assumptions of the TCR theory, which are common in both groups can be summarized as

- contacting surfaces are rough, isotropic, with a Gaussian asperity distribution
- behavior of a given microcontact is independent of all other microcontacts
- interfacial force on any microcontact spot acts normally (no frictional or tangential forces)
- the deformation mechanics, i.e., the stress and displacement fields are uniquely determined by the shape of the equivalent surface.

With the concept of equivalent roughness, the plastic model assumes that the asperities are flattened or equivalently penetrate into the smooth surface without any change in the shape of the part of surfaces not yet in contact. As mentioned in the previous section, the pressure at microcontacts is sufficiently large, i.e., larger than the strength of the materials in contact. Tabor [9] suggested that these contact pressures were equal to the flow pressure of the softer of the two contacting materials. Therefore, in plastic models the pressure at microcontacts is effectively independent of load and the contact geometry. The real area of contact is then proportional to the load, \( A_c/A_a = P/H_{mic} \), where \( P \) is the apparent contact pressure. Cooper et al. (CMY) [7], based on the level-crossing theory and using the equivalent surface approximation, derived relationships for mean microcontact size and number of microcontacts by assuming hemispherical asperities whose height and surface slopes have Gaussian distributions. The CMY model was...
essentially based on the assumption that each microcontact consists of two hemispherical asperities in symmetric plastic contact. Later Yovanovich [12] summarized the CMY [7] model and reported relationships for calculating the contact parameters; and also proposed a compact expression for calculating the TCR of conforming rough joints as

$$h_p = 1.25 k_s (m/\sigma) (P/H\text{mic})^{0.95} \quad (6)$$

where \(k_s\) is the harmonic mean of the thermal conductivities

$$k_s = \frac{2k_1 k_p}{k_1 + k_p}$$

Considering the fact that the plastic deformation is irreversible and cannot be repeated on subsequent loadings, Arhart [13] stated that the normal contact of rough surfaces could be plastic at first several contacts but for moving machine parts that meet millions during their life the contact must reach an “elastic” state. Arhart [13] showed that any elastic model based on simple Hertzian theory in which the number of contacts remains constant, as the load increases, will give \(A_e \sim F^{2/3}\), which does not satisfy the observed proportionality \(A_e \sim F\) reported by Tabor [9]. But, if the average contact size remains constant, and the number of microcontact increases, the area would be proportional to the load. This approach led to an “effective elastic hardness” \(H_e\).

Mikic [6] based on the CMY model proposed an elastic model. He assumed that the elastic real contact area is half of the plastic contact area, i.e., \(A_{elastic}/A_{plastic} = 1/2\). Mikic’s model satisfied the linear proportionality between the applied load and the real contact area; he also proposed an effective elastic microhardness as [6]

$$H_e = \frac{E'm}{\sqrt{2}} \quad (7)$$

with,

$$\frac{1}{E'} = \frac{1 - \nu_1^2}{E_1} + \frac{1 - \nu_2^2}{E_2}$$

where \(E, \nu\) are the elastic modulus and Poisson’s ratio, respectively. Mikic proposed an expression for calculating the TCR of conforming rough joints assuming elastic deformation [6]

$$h_e = 1.55 k_s (m/\sigma) (P/H_e)^{0.94} \quad (8)$$

It can be seen from Eq. (8) that \(h_e\) is a very weak function of \(m\). Another correlation was suggested by Mikic assuming a typical value of \(m = 0.1\) as, \(h_e = 1.55 (k_s/\sigma) (P/H_e)^{0.94}\).

Table 1 summarizes the relationships of the CMY [7] plastic model and the Mikic [6] elastic model.

<table>
<thead>
<tr>
<th>Model</th>
<th>Relation</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\kappa = 1) elastic</td>
<td>(A_e = \frac{\kappa}{4} \text{erfc}(\lambda))</td>
</tr>
<tr>
<td>(\kappa = 2) plastic</td>
<td>(n_s = \frac{1}{16} \left(\frac{m}{\sigma}\right)^2 \text{exp} \left(-2\lambda^2\right) A_a)</td>
</tr>
<tr>
<td>(\lambda = Y/\sqrt{2}\sigma)</td>
<td>(A_a = \pi b^2 \left(\frac{\sigma}{m}\right) \text{exp} \left(\lambda^2\right) \text{erfc}(\lambda))</td>
</tr>
<tr>
<td>Plastic [7]</td>
<td>(\lambda = \text{erfc}^{-1}(2P/H\text{mic}))</td>
</tr>
<tr>
<td>Elastic [6]</td>
<td>(\lambda = \text{erfc}^{-1}(4P/H_e))</td>
</tr>
</tbody>
</table>

### 4.3 Deformation Mode of Asperities

A priori assumption of deformation mode of asperities could lead to wrong conclusions; the effective elastic microhardness \(H_e\), calculated from Eq. (7), could result in unrealistic values larger than the material microhardness, i.e., \(H_e > H\text{mic}\). This is physically impossible. To avoid this, Mikic [6] performed an analysis to determine the mode of deformation and proposed the plasticity index:

$$\gamma = \frac{H\text{mic}}{E'm} \quad (9)$$

According to [6] for surfaces with \(\gamma \geq 3\), 90% of the actual area will have the elastic contact pressure, therefore, the contact will be predominantly elastic; and for \(\gamma \leq 0.33\), 90% of the actual area will have the plastic contact pressure, therefore, the contact will be predominantly plastic.

$$\begin{cases} 
\gamma \leq 0.33 & \text{asperities deform plastically} \\
0.33 \leq \gamma \leq 3.0 & \text{transition} \\
\gamma \leq 3.0 & \text{asperities deform elastically}
\end{cases} \quad (10)$$

Mikic [6] concluded that for most engineering surfaces the asperity deformation mode is plastic and the average asperity pressure is the microhardness. It can be seen that the deformation mode of asperities depends on material properties \((E'\text{ and } H\text{mic})\) and the shape of asperities \(m\). Mikic also reported that the mode of deformation, as stated by Greenwood and Williamson [14], is not sensitive to the pressure level.

Fuller and Marotta [8] developed an elastic model for calculating the TCR of polymer-metal interface and showed good agreement with the experimental data. They reported an effective elastic microhardness as

$$H_{e,FM} = \frac{E_p m}{2.3} \quad (11)$$

In developing Eq. (11), only the polymer Young’s modulus \(E_p\) was considered instead of the effective elastic modulus.
In other words, Eq. (11) neglects the lateral strain, i.e., the effect of Poisson’s ratio.

Fuller and Marotta [8] did not implement the deformation analysis proposed by Mikic. Applying the deformation analysis, one observes that the deformation mode of asperities is plastic for most polymers studied. Assuming elastic deformation mode of asperities (when $\gamma < 3$) may result in unrealistic situations where the effective elastic microhardness becomes larger than the actual (measured) hardness value, i.e., $H_e > H_{mic}$.

The deformation was assumed elastic in [8] based on the surface measurements performed by Parihar and Wright [5]. Parihar and Wright [5] measured the surface roughness and asperities slope of a stainless steel flux meter and a silicone rubber specimen before and after loading. They concluded that the contact was elastic since the roughness measurements were identical. This conclusion is incorrect. The mean RMS roughness $\sigma$ and the mean absolute surface slope $m$, based on their definition, are statistical measures of rough surfaces. These values will not be influenced by the deformation due to contact since the real contact area is a small fraction of the apparent area. Clausing and Chao [15] measured roughness of metallic specimens before and after loading and reported identical values. They also measured the joint resistance as the applied load was varied; then removed the load and broke the joint and rotated the upper specimen by 90° and repeated the test with the same pair. No changes were observed in thermal joint resistance measurements [16] which was consistent with their roughness measurements before and after loading.

In conclusion, as stated by Mikic [6] and Greenwood and Williamson [14], the deformation mode of asperities is not an arbitrary choice and should be determined using the plasticity index.

5 PRESENT MODEL

The present model is based on the premise that the load-displacement behavior of polymers is similar to metals in the temperature range of interest, i.e., temperatures below the glassy temperature. A successful thermal joint resistance model accounts for both the bulk and the thermal contact resistance of the polymer metal joint. The bulk resistance of the polymer can be calculated combining Eqs. (2) and (5) as

$$R_b = \frac{t_0 (1 - P/E_p)}{A_a k_p}$$

Equation (12) was first used by Fuller and Marotta [8] to calculate the bulk conductance. A deformation analysis is performed and the plasticity indices, Eq. (9), are listed for the polymers used in this study in Table 3. Since the plasticity index $\gamma$ approaches 0.33 and for most polymers is less than 3, the deformation mode of asperities is assumed to be plastic in this study.

The microcontacts are often assumed to be isothermal [10]. Thermal constriction/spreading resistance of microcontacts can be modeled using a flux tube geometry [7] or if microcontacts are considered to be located far enough from each other, the isothermal heat source on a half-space solution [3] can be used. Bahrami et al. [17] compared these solutions and showed that the microcontacts can be considered as heat sources on a half-space for most engineering applications. Bahrami et al. [17] assumed plastically deformed asperities and used scale analysis techniques and developed a compact model to predict thermal constriction/spreading resistance through the microcontacts, $R_s$

$$R_s = \frac{0.565H_{mic} (\sigma/m)}{k_s F}$$

where $F$ is the applied load. Since Eq. (13) yields close values to the Yovanovich [12] expression, Eq. (6), thus both relationships can be used. The joint resistance then can be calculated from Eq. (3) as

$$R_j = \frac{0.565H_{mic} (\sigma/m)}{k_s P A_a} + \frac{t_0 (1 - P/E_p)}{A_a k_p}$$

Since the thermal conductivity of polymers is relatively small compared to metals, thus the harmonic mean of thermal conductivities of a metal-polymer interface is controlled by the polymer thermal conductivity, i.e., $k_s \approx k_p$. Also, values of the effective elastic modulus $E'$ and the joint microhardness $H_{mic}$ are controlled by the polymer values. Therefore, the joint resistance is dominated by the polymer parameters.

The asperity slope values reported by Fuller and Marotta [8] are relatively large compared to previously measured slope values published by Marotta and Fletcher [2]. Marotta and Fletcher [2] slope values are used to develop a

<table>
<thead>
<tr>
<th>Test</th>
<th>$\sigma$</th>
<th>$m$</th>
<th>$E_p$</th>
<th>$H_{mic}$</th>
<th>$k_p$</th>
<th>$t_0$</th>
<th>$\gamma$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delrin1</td>
<td>2.19</td>
<td>0.26</td>
<td>3.59</td>
<td>0.37</td>
<td>0.38</td>
<td>1.27</td>
<td>0.37</td>
</tr>
<tr>
<td>Delrin2</td>
<td>2.29</td>
<td>0.26</td>
<td>3.59</td>
<td>0.37</td>
<td>0.38</td>
<td>1.32</td>
<td>0.36</td>
</tr>
<tr>
<td>Polyethylene</td>
<td>0.81</td>
<td>0.15</td>
<td>2.38</td>
<td>0.15</td>
<td>0.22</td>
<td>1.44</td>
<td>0.38</td>
</tr>
<tr>
<td>PVC</td>
<td>0.74</td>
<td>0.15</td>
<td>4.14</td>
<td>0.15</td>
<td>0.17</td>
<td>1.31</td>
<td>0.23</td>
</tr>
<tr>
<td>Al 6061</td>
<td>0.51</td>
<td>0.05</td>
<td>72.1</td>
<td>-</td>
<td>183</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
correlation to estimate the surface asperity slope as a function of roughness as follows:

$$m = 0.19 \left( \frac{\sigma}{\sigma_0} \right)^{0.52}$$  \hspace{1cm} (15)

where $\sigma$ is in micron and $\sigma_0 = 1 \mu m$. The slope values listed in Tables 1 and 2 are the estimated values using Eq. (15).

Figures 3 to 5 illustrate comparisons between the present model, Eq. (14), and the experimental data sets Delrin1, Delrin2, and Polyethylene collected by Fuller and Marotta [8], respectively. Their experimental set up is described in section 2 and schematically shown in Fig. 1 with $b_L = 12.7 \ mm$. The mean temperature of the polymer specimens were maintained at $40 \pm 1^\circ C$ which was lower than the glassy temperature for the polymers tested. The combined roughness $\sigma$ and surface asperities slope $m$ (corrected values using Eq. (15)), thermal conductivity $k_p$, microhardness $H_{mic}$, and elastic modulus $E_p$ for polymer specimens are summarized in Table 2.

The comparisons show the two components of the joint resistance i) constriction/spreading resistance of microcontacts $R_s$, i.e., the TCR resistance at the metal-polymer interface and ii) the polymer bulk resistance $R_b$. As shown, the bulk resistances of polymers remained almost constant as the applied load was increased during the tests. This was because of the relatively small changes in the polymer thickness $t$. It can also be seen that at relatively light loads the microcontacts resistance controls the joint resistance. As the applied load increases, the contact resistance decreases linearly, Eq. (13) and the joint resistance approaches the bulk resistance at relatively high loads. These trends can be observed in all experimental data.

The present model is also compared with the experimental data collected by Marotta and Fletcher [2]. The experimental equipment was similar to that used in [8] with $b_L = 12.7 \ mm$. The mean temperature of the polymer specimens were maintained at $40^\circ C$. They measured the joint resistance of several polymer specimens. The combined roughness $\sigma$ and surface asperities slope $m$ (corrected values using Eq. (15)), thermal conductivity $k_p$, microhardness $H_{mic}$, and elastic modulus $E_p$ for the polymer specimens tested by [2] are summarized in Table 3.

Figures 6 and 7 show the comparison between the
Table 3. POLYMERS CHARACTERISTIC DATA, MAROTTA AND FLETCHER [2]

<table>
<thead>
<tr>
<th>Test</th>
<th>σ</th>
<th>m</th>
<th>E</th>
<th>$H_{mic}$</th>
<th>$k_p$</th>
<th>$t_0$</th>
<th>γ</th>
</tr>
</thead>
<tbody>
<tr>
<td>µm</td>
<td>GPa</td>
<td>GPa</td>
<td>W/mK</td>
<td>mm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ABS</td>
<td>0.93</td>
<td>0.17</td>
<td>2.90</td>
<td>0.17</td>
<td>0.18</td>
<td>1.59</td>
<td>0.30</td>
</tr>
<tr>
<td>Delrin</td>
<td>1.42</td>
<td>0.21</td>
<td>3.59</td>
<td>0.37</td>
<td>0.38</td>
<td>1.62</td>
<td>0.46</td>
</tr>
<tr>
<td>Nylon</td>
<td>1.23</td>
<td>0.20</td>
<td>2.11</td>
<td>0.41</td>
<td>0.31</td>
<td>1.65</td>
<td>0.90</td>
</tr>
<tr>
<td>Phenolic</td>
<td>1.13</td>
<td>0.19</td>
<td>6.80</td>
<td>0.36</td>
<td>0.65</td>
<td>1.56</td>
<td>0.26</td>
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<tr>
<td>Poly1</td>
<td>0.82</td>
<td>0.16</td>
<td>2.39</td>
<td>0.14</td>
<td>0.22</td>
<td>1.62</td>
<td>0.32</td>
</tr>
<tr>
<td>Poly2</td>
<td>1.92</td>
<td>0.24</td>
<td>3.00</td>
<td>0.13</td>
<td>0.45</td>
<td>1.66</td>
<td>0.17</td>
</tr>
<tr>
<td>Poly3</td>
<td>1.33</td>
<td>0.20</td>
<td>1.90</td>
<td>0.41</td>
<td>0.31</td>
<td>1.64</td>
<td>0.97</td>
</tr>
<tr>
<td>PVC</td>
<td>0.56</td>
<td>0.14</td>
<td>2.50</td>
<td>0.15</td>
<td>0.17</td>
<td>1.62</td>
<td>0.37</td>
</tr>
<tr>
<td>Teflon</td>
<td>1.52</td>
<td>0.22</td>
<td>0.46</td>
<td>0.20</td>
<td>0.25</td>
<td>1.68</td>
<td>1.78</td>
</tr>
<tr>
<td>Al 6061</td>
<td>0.51</td>
<td>0.05</td>
<td>72.1</td>
<td>−</td>
<td>208</td>
<td>−</td>
<td>−</td>
</tr>
</tbody>
</table>

Poly1: Polycarbonate, Poly2: Polyethylene, Poly3: Polypropylene

Figure 6. COMPARISON BETWEEN PRESENT MODEL AND ABS, MAROTTA AND FLETCHER [2]

The proposed model, Eq. (14), and the experimental data sets ABS and PVC collected by Marotta and Fletcher [8], respectively. As can be observed, the range of applied load in the experiments was relatively high. As a result, the bulk thermal resistance is the controlling component and the contact resistance is relatively small. As shown, the present model predicts the trend of the data very well.

The proposed model can be non-dimensionalized with respect to the bulk thermal resistance as follows:

$$R_j^* = \frac{R_j}{R_b} = 1 + \Theta$$

where the non-dimensional parameter $\Theta$ is

$$\Theta = \frac{R_s}{R_b} = \frac{k^*}{P^*} \left( \frac{\sigma}{m} \right) \left( \frac{t_0}{E_p} \right) (1 - P/E_p)$$

where $k^* = k_p/k_s$ and $P^* = P/H_{mic}$. The non-dimensional parameter $\Theta$ includes all the joint parameters: contact pressure, macro and micro geometrical parameters, i.e., $\sigma$, $m$, and $t_0$ as well as thermal conductivities $k_p$, $k_s$ and elastic and plastic mechanical properties of the joint $E_p$ and $H_{mic}$. Based on this non-dimensional parameter, we define a criterion to specify the relative importance of the thermal joint resistance components as a function of the joint input parameters

$$\Theta \ll 1 \quad R_b \text{ controls } R_j$$

$$\Theta \sim 1 \quad \text{both } R_s \text{ and } R_b \text{ are important}$$

$$\Theta \gg 1 \quad R_s \text{ controls } R_j$$

Figure 8 presents a non-dimensional comparison between the data and the model, Eq. (16), where the non-dimensional pressure $P^* = P/H_{mic}$ is varied over a wide range. All experimental data collected by [2] and [8], more than 120 data points in 13 data sets, are included in the comparison. Two asymptotes defined in Eq. (16) are also shown in the plot. As the non-dimensional pressure varies from relatively small to large values, the non-dimensional
joint resistance predicted by the model moves from the TCR to the bulk resistance asymptote. The experimental data covers a relatively wide range of the non-dimensional parameter \( \Theta \), \( 0.03 \leq \Theta \leq 15.72 \) and follows the trend of the model. The RMS relative difference between the proposed model and all experimental data is approximately 12.7%. The total average uncertainty of both experimental data sets were reported to be approximately 17% by [2] and [8].

6 SUMMARY

Thermal joint resistance of rough polymer-metal interfaces in a vacuum is studied and a compact analytical model is developed for temperatures less than the glassy temperature. Performing a critical review, we show that the existing model in the literature is based on incorrect assumptions that results in unrealistic values for microhardness of the polymers.

The present model assumes that the mechanical behaviors of polymers are similar to metals for temperatures below the glassy temperature. The existing plastic and elastic TCR models are reviewed and the deformation modes of surface asperities are discussed. It is shown that the deformation mode of asperities, for most of the tested polymers, is close to plastic. The proposed joint resistance model includes two components: i) bulk resistance of the polymer and ii) TCR of the microcontacts at the interface.

It is observed that the thermophysical properties of the polymer controls the thermal joint resistance and the metallic body properties have a second order effect on the thermal joint resistance.

A new non-dimensional parameter is introduced which represents the ratio of the TCR over the bulk thermal resistances. Based on this non-dimensional parameter, a criterion is proposed for specifying the relative importance of TCR of the joint.

The present model is compared with 13 polymer-metal data sets, 127 experimental data points that cover a variety of polymers, collected by [2] and [8]; and shows good agreement. The RMS difference between the model and data is approximately 12.7% over the entire range of the comparison.

Further experimental investigation is highly recommended for lighter loads, where the thermal microcontacts resistance \( R_s \) controls the joint resistance \( R_j \). The non-dimensional parameter \( \Theta \) can be used to identify the light load range. Also, additional microhardness and surface measurements (\( \sigma, m \)) are required at different roughness levels to confirm the deformation mode of asperities for polymers.

ACKNOWLEDGMENT

The authors (M.B. and M. M. Y.) gratefully acknowledge the financial support of the Centre for Microelectronics Assembly and Packaging, CMAP and the Natural Sciences and Engineering Research Council of Canada, NSERC.

REFERENCES

Figure 8. COMPARISON BETWEEN NON-DIMENSIONAL JOINT RESISTANCE PREDICTED BY PRESENT MODEL AND EXPERIMENTAL OF FULLER AND MAROTTA [8] AND MAROTTA AND FLETCHER [2]


