Developing Multilayer Thin Film Strain Sensors With High Thermal Stability

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Abstract
A multilayer thin film strain sensor for large temperature range use is under development using a reactively-sputtered process. The sensor is capable of being fabricated in fine line widths utilizing the sacrificial-layer lift-off process that is used for microfabricated noble-metal sensors. Tantalum nitride films were optimized using reactive sputtering with an unbalanced magnetron source. A first approximation model of multilayer resistance and temperature coefficient of resistance was used to set the film thicknesses in the multilayer film sensor. Two multifunctional sensors were fabricated using multilayered films of tantalum nitride and palladium chromium, and tested for low temperature resistivity, TCR and strain response. The low temperature coefficient of resistance of the films will result in improved stability in thin film sensors for low to high temperature use.

Nomenclature
CTE  coefficient of thermal expansion (ppm/°C)
d  film thickness (Å)
ε  strain; change per unit length (με)
εa  apparent strain due to temperature rather than applied strain (με)
δεa/δT  apparent strain sensitivity to temperature changes (με/°C)
γ  gauge factor of strain gauge
δl/l  length change per unit length (ε)
με  unit of microstrain; typically defined as 10⁻⁶ inch change per inch length (μin/in)
PdCr  palladium chromium alloy
ρ  electrical resistivity (μΩ-cm)
σ  electrical conductivity (Ω-cm)⁻¹
TaN  tantalum nitride (no specific phase)
TCR  temperature coefficient of resistance (ppm/°C)

I. Introduction
A. Challenge of Sensors for Propulsion Systems
To advance knowledge in fundamental aeronautics and develop technologies for safer, lighter, quieter, and more fuel efficient aircraft, instrumentation technologies are being developed by the National Aeronautics and Space Administration (NASA) in support of its mission to pioneer the future in space exploration, scientific discovery, and aeronautics research. These technologies also enable the capabilities for long duration, more distant human and robotic missions for the Vision for Space Exploration.

The Sensors and Electronics Branch of NASA Glenn Research Center (GRC) has an in-house effort to develop thin film sensors for surface measurement in propulsion system research. The sensors include those for strain, temperature, heat flux and surface flow which will enable critical vehicle health monitoring and characterization of components of future space and air vehicles.

The use of sensors made of thin films has several advantages over wire or foil sensors. Thin film sensors do not require special machining of the components on which they are mounted, and, with thicknesses less than 10 μm,
they are considerably thinner than wire or foils. Thin film sensors are thus much less disturbing to the operating environment, and have a minimal impact on the physical characteristics of the supporting components.

The need to consider ceramic sensing elements is brought about by the temperature limits of metal thin film sensors in propulsion system applications. Longer-term stability of thin film sensors made of noble metals has been demonstrated at 1100 °C for 25 hr (ref. 1) The capability for thin film sensors to operate in 1500 °C environments for 25 hr or more is considered critical for ceramic turbine engine development (refs. 2 and 3). For future space transportation vehicles, temperatures of propulsion system components of at least 1650 to 3000 °C are expected (ref. 4).

B. Limits of Metal Film Sensors

A limitation of thin films used as sensors to measure strain is that their resistance changes as the temperature changes. This apparent strain (εa) can be falsely interpreted as actual strain on the component being monitored. For static strain applications for use on gas turbine engines, the current required accuracy is ±200 μin/in (με), approximately ±10 percent of full scale, with the goal of ±1 percent accuracy (ref. 5). The thin film palladium-chromium (PdCr) alloy strain gauge, developed at NASA GRC for high temperature strain measurement application, is stable to 1100 °C, but has a temperature coefficient of resistance (TCR) of 135 ppm/°C and an apparent strain sensitivity (δεa/δT) of 85 με/°C, requiring temperature compensation for high temperature static strain measurements (refs. 1 and 5). Currently, this compensation is in the form of setting a “ballast” potentiometer in a bridge to perform first order elimination of the apparent strain at a particular temperature, but deviations from this matched temperature results in measured apparent strain (ref. 6).

A thin film multifunctional sensor developed at NASA GRC that can measure directional strain, flow, heat flux, and temperature utilizes this PdCr alloy (ref. 7), but does not incorporate a compensation bridge in its design and is limited to dynamic strain measurements at high temperature. A thin film strain sensor with thermal stability over a wide range of temperatures would allow high temperature static measurements with the multifunctional sensor as well as a more passive method of eliminating apparent strain without the need for a compensation bridge. NASA GRC has thus begun an in-house effort to develop thin film strain sensors with high thermal stability. Ultimately, the goal is to be able to achieve the desired ±20 με accuracy of measured applied static strain being no less than 0.1 percent of a total strain measurement (= applied + apparent + drift strain), or ±20,000 με. The total apparent strain of 20,000 με limits the desired apparent strain sensitivity to temperature to be less than ±20 με/°C over the current temperature range. As this goal is approached, the drift strain (creep) will also be considered as part of the total strain measurement.

II. Ceramic-Based Film Sensor Development

A. Background

Since 1991, there have been many investigations into the application of ceramic thin films for use as high temperature thin film strain gauges. A summary of notable high temperature thin film strain gauges is given in table 1. Thin film resistors for regulating electronics based on doped nickel-chromium alloy films with a TCR of ±5 ppm/°C and greater are common in the electronics industry, but they are generally restricted to a temperature range between –55 to 125 °C (ref. 8), and thus not suitable for our applications.

<table>
<thead>
<tr>
<th>Gauge material</th>
<th>TCR (ppm/°C)</th>
<th>Gauge factor (γ) (δR/Rε)</th>
<th>Apparent strain sensitivity (με/°C)</th>
<th>Maximum use temperature (if reported)</th>
<th>Fabrication notes</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni-20%Cr</td>
<td>290</td>
<td>2.5</td>
<td>116</td>
<td>700 °C</td>
<td>COTS standard</td>
<td>14,15</td>
</tr>
<tr>
<td>Pd-13%Cr</td>
<td>135</td>
<td>2 to 1.4</td>
<td>85</td>
<td>1100 °C</td>
<td>NASA standard</td>
<td>1</td>
</tr>
<tr>
<td>AlN</td>
<td>–1281 to 109</td>
<td>3.7 to 15</td>
<td>–344 to 29</td>
<td>&gt;1100 °C</td>
<td>Al reacted with N</td>
<td>16</td>
</tr>
<tr>
<td>ITO</td>
<td>–469 to 230</td>
<td>–6.5 to 11.4</td>
<td>–35 to 72</td>
<td>&gt;1100 °C</td>
<td>Oxygen doping</td>
<td>17</td>
</tr>
<tr>
<td>Al:ITO</td>
<td>–1200</td>
<td>8</td>
<td>–150</td>
<td>1280 °C</td>
<td>Aluminum doping</td>
<td>18</td>
</tr>
<tr>
<td>TaB2</td>
<td>–50</td>
<td>1.4</td>
<td>–36</td>
<td>&lt;3225 °C</td>
<td>Nitrogen doping</td>
<td>19</td>
</tr>
<tr>
<td>Cu:TaN</td>
<td>–800 to 200</td>
<td>2.3 to 5.1</td>
<td>–348 to 87</td>
<td>(not reported)</td>
<td>Ta reacted w/N; Cu doping</td>
<td>20</td>
</tr>
<tr>
<td>TaN</td>
<td>–80</td>
<td>3.5</td>
<td>–23</td>
<td>(not reported)</td>
<td>Ta reacted with N</td>
<td>21</td>
</tr>
<tr>
<td>TaON</td>
<td>–290</td>
<td>3.5</td>
<td>–83</td>
<td>(not reported)</td>
<td>Ta reacted with N; 1 percent Ox</td>
<td>22</td>
</tr>
</tbody>
</table>

TABLE 1.—A REVIEW OF HIGH TEMPERATURE THIN FILM STRAIN GAUGE APPLICATIONS
The gauge factor (γ) of the strain gauge relates the sensitivity of the gauge to strain (ε = δl/l), as shown in equation (1).

$$\frac{\delta R}{R} = \gamma \frac{\delta l}{l} = \gamma \varepsilon$$  (1)

The apparent strain sensitivity to temperature ($\delta \varepsilon_a/\delta T$) is the TCR divided by the gauge factor plus the difference in the substrate and the gauge material’s coefficient of thermal expansion (CTE), as shown in equation (2). The difference in the CTE’s is expected to be less than 5 ppm/°C based on the materials that we are exploring, and this will be left as an uncertainty in our apparent strain calculation.

$$\frac{\delta \varepsilon_a}{\delta T} = \frac{\text{TCR}}{\gamma} + \Delta \text{CTE}$$  (2)

Based on the reported gauges in table 1, the maximum use temperature of tantalum nitride (TaN) may be up to 2000 °C, the most attractive for high temperature applications. As a thin film, TaN is known as a stable high temperature resistor with TCR between 200 and –200 ppm/°C, depending on the fabrication process, nitride phase produced (e.g., Ta2N, TaN, Ta5N6, etc.), incorporation of oxide on the TaN grains and the degree of amorphous structure (refs. 9 to 12). The report of studies (refs. 13) of resistors using NiCr and TaNiCr interlayers with Ta2N to achieve TCR between 5 and –5 ppm/°C began our investigation to develop a tantalum nitride film for use with the PdCr strain gauge to achieve the passive elimination of apparent strain sensitivity.

### B. Tantalum Nitride Film Fabrication

The first step to developing an interlayered or multilayered TaN/PdCr film was to develop a process of reactively sputtering the tantalum nitride consistently. The first depositions were on alumina substrates with the film resistivity measured using a four-point probe (ref. 23) with spacing of 2.54 mm. The resistivity of Ta2N films can vary between 200 and 264 μΩ-cm, again due to the incorporation of oxide and degree of amorphous structure (refs. 9 to 12). The results of these first runs are given in table 2. The resistivities given in the table are considered accurate to ±1.8 percent.

<table>
<thead>
<tr>
<th>Sample</th>
<th>RF power (W)</th>
<th>Process pressure (mTorr)</th>
<th>Argon flow (sccm)</th>
<th>Nitrogen flow (sccm)</th>
<th>Run time (sec)</th>
<th>Film thickness (μm)</th>
<th>Resistivity ρ (μΩ-cm)</th>
<th>TCR (ppm/°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JG40901</td>
<td>250</td>
<td>8</td>
<td>20</td>
<td>0</td>
<td>3600</td>
<td>1.8</td>
<td>74</td>
<td>----</td>
</tr>
<tr>
<td>JG40903</td>
<td>250</td>
<td>8</td>
<td>20</td>
<td>2</td>
<td>7200</td>
<td>2.2</td>
<td>383</td>
<td>----</td>
</tr>
<tr>
<td>JG40908</td>
<td>250</td>
<td>8</td>
<td>38</td>
<td>2</td>
<td>7200</td>
<td>5.0</td>
<td>265</td>
<td>----</td>
</tr>
<tr>
<td>JG40909</td>
<td>250</td>
<td>8</td>
<td>38</td>
<td>2</td>
<td>7200</td>
<td>4.8</td>
<td>269</td>
<td>----</td>
</tr>
<tr>
<td>JG40922</td>
<td>250</td>
<td>8</td>
<td>38</td>
<td>2</td>
<td>7200</td>
<td>3.9</td>
<td>259</td>
<td>–93</td>
</tr>
<tr>
<td>JG41123</td>
<td>250</td>
<td>4</td>
<td>38</td>
<td>2</td>
<td>7200</td>
<td>3.7</td>
<td>232</td>
<td>–400</td>
</tr>
<tr>
<td>JG41124</td>
<td>250</td>
<td>3</td>
<td>38</td>
<td>2</td>
<td>7200</td>
<td>3.7</td>
<td>269</td>
<td>–100</td>
</tr>
<tr>
<td>JG50114</td>
<td>250</td>
<td>3</td>
<td>38</td>
<td>2</td>
<td>3600</td>
<td>3.7</td>
<td>440</td>
<td>–110</td>
</tr>
<tr>
<td>JG50115</td>
<td>125</td>
<td>2</td>
<td>38</td>
<td>2</td>
<td>3600</td>
<td>1.0</td>
<td>155</td>
<td>–200</td>
</tr>
</tbody>
</table>

The confidence gained with the results of the last two runs resulted in the fabrication of a multifunctional sensor (run JG40922) with a length to width ratio (l/w) of 290 of tantalum nitride consistently. The first depositions were on alumina substrates with the film resistivity measured using a four-point probe (ref. 23) with spacing of 2.54 mm. The resistivity of Ta2N films can vary between 200 and 264 μΩ-cm, again due to the incorporation of oxide and degree of amorphous structure (refs. 9 to 12). The results of these first runs are given in table 2. The resistivities given in the table are considered accurate to ±1.8 percent.

The confidence gained with the results of the last two runs resulted in the fabrication of a multifunctional sensor (run JG40922) with a length to width ratio (l/w) of 290 of tantalum nitride using the run parameters of runs JG40908 and JG40909. The sensor was patterned using the sacrificial-layer lift-off process that is used for microfabricated noble-metal sensors of fine line widths (ref. 24). The completed sensor is shown in figure 1. The TCR was measured to be –93 ppm/°C, and the resistivity 259 μΩ-cm. The strain sensitivity was measured using the algorithm developed for the sensor (ref. 7), and the gauge factor was found to be 3.9±0.1, and the angle resolution was determined to be less than ±0.2°. This translates to an apparent strain sensitivity of –24 με/°C, similar to what was reported by Ayerd, et al. (ref. 21) and close to our goal of ≤±20 με/°C. A graph showing the output of the gauge factor vs. applied strain versus the applied strain is shown in figure 2.
Target arcing and degradation, however, was becoming an issue, with the thickness of the sensor of run JG40922 about 20 percent less than the initial runs JG40908 and JG40909. For the next set of runs, the system pressure and power were lowered to prevent the arcing and degradation of the tantalum target. The films were patterned using a shadow mask with $l/w = 21.88$, allowing the resistivity to be measured as well as the TCR using a four wire method with a data acquisition system. The resulting film of run JG50115 had a TCR of $-200$ ppm/°C and a resistivity of 155 $\mu\Omega$-cm.
III. Multilayered Film Fabrication

A. Multilayer Film Approach

Once the TaN film deposition parameters were finalized, the multilayered TaN/PdCr films were attempted. A cross-section schematic of the multilayered film is shown in figure 3. As an estimate for the relative thicknesses, the assumption is made that because the film thicknesses (d) are greater than the grain size, and thus electron mean free path, the films can be treated as three independent layers in parallel (to first approximation) (ref. 25).

The conductivity (σ) is the inverse of the resistivity (ρ), and because the first and last layers are the same material (TaN) and the second is PdCr, the conductivities add as in equation (3).

\[
\sigma_{\text{Total}} = \sigma_{\text{TaN}} \frac{d_{\text{TaN}}}{d_{\text{Total}}} + \sigma_{\text{PdCr}} \frac{d_{\text{PdCr}}}{d_{\text{Total}}} \tag{3}
\]

Similarly, the TCR (α) for multiple layers is derived as in equation (4).

\[
a_{\text{Total}} = \frac{d_{\text{TaN}} \sigma_{\text{TaN}} \alpha_{\text{TaN}} + d_{\text{PdCr}} \sigma_{\text{PdCr}} \alpha_{\text{PdCr}}}{d_{\text{TaN}} \sigma_{\text{TaN}} + d_{\text{PdCr}} \sigma_{\text{PdCr}} } \tag{4}
\]

To cancel out the TCR, the numerator is set to zero, as shown in equation (5).

\[
d_{\text{TaN}} \sigma_{\text{TaN}} \alpha_{\text{TaN}} = -d_{\text{PdCr}} \sigma_{\text{PdCr}} \alpha_{\text{PdCr}} \tag{5}
\]

The ratio of PdCr film thickness to TaN film thickness then can be derived in equation (6).

\[
\frac{d_{\text{PdCr}}}{d_{\text{TaN}}} = -\frac{\sigma_{\text{TaN}} \alpha_{\text{TaN}}}{\sigma_{\text{PdCr}} \alpha_{\text{PdCr}}} = -\frac{\rho_{\text{PdCr}}}{\rho_{\text{TaN}}} \frac{\alpha_{\text{TaN}}}{\alpha_{\text{PdCr}}} \tag{6}
\]

So, to a first approximation, the TCR of the multilayer can be minimized with the knowledge of the resistivity and TCR of the component films. Using the values of TaN from JG50115, and the resistivity and TCR of PdCr as 110 μΩ-cm and 170 ppm/°C respectively, the ratio of PdCr to TaN thickness to cancel the multilayer film TCR is found to be 0.835.

![Cross-section schematic of the multilayer thin film sensor.](Figure 3.)
B. Multilayered Film Samples

Three sets of TaN/PdCr multilayer films were fabricated with different PdCr to TaN thicknesses, with each set having two samples using identical machine parameters. The films were patterned using a shadow mask with $l/w = 21.88$ as shown in figure 4, allowing the resistivity to be measured as well as the TCR using a four wire method with a data acquisition system. After the two samples in a set were completed, they were annealed at 600 °C in $<10^{-6}$ Torr vacuum for 8 hr to stabilize the multilayer films. The results are shown in table 3, and the conductivity and TCR are shown in graphs in figures 5 and 6 respectively with the theoretical plot based on values of TaN from JG50115. Note that the top 560Å of the samples are being forced to be TaO, which is used to control the passivating oxide that would normally form on TaN films. Examining each of these sets, the conductivity and TCR of JG50510 does appear to be inconsistent with the rest of the films, so much as to call into question the reaction of the formation of tantalum nitride in the fabrication process.

As TaN films seem to vary more than annealed metallic films from bulk values in literature (refs. 9 to 12 and 26), the determination of the resistivity and TCR of the TaN films of the multilayer films will shed light on the variation of the parameters for TaN for these runs. From equations (3) and (4), the TCR and resistivity of the TaN portion of the films formed in the multilayer can be determined.

Since the thicknesses of the films individually are measured, as is the total conductivity and TCR (as shown in table 3), the TaN conductivity can be estimated from equation (3):

$$\sigma_{TaN} = \frac{\sigma_{Total} d_{Total} - \sigma_{PdCr} d_{PdCr}}{d_{TaN}}$$

(7)
Likewise, the TCR can be estimated from equation (4):

$$\alpha_{\text{TaN}} = \frac{d_{\text{Total}}\sigma_{\text{Total}}\alpha_{\text{Total}} - d_{\text{PdCr}}\sigma_{\text{PdCr}}\alpha_{\text{PdCr}}}{\sigma_{\text{TaN}}d_{\text{TaN}}}$$  \hspace{1cm} (8)

Table 4 gives the results of these calculations for the six samples. The TCR and resistivity for the TaN films for JG50510 is observed to be more similar to Ta+Ta₂N mixed films than to pure Ta₂N films (ref. 9). If the TaN for JG50510 is included, an average resistivity and TCR for the TaN for the samples are 189±31 μΩ-cm and –93±94 ppm/°C respectively. Without JG50510, the resistivity and TCR are 199±24 μΩ-cm and –130±30 ppm/°C. The latter values are used in the “best fit” curves in figures 5 and 6.

**TABLE 4.—TaN PROPERTIES FROM TaN/PdCr DEPOSITIONS**

<table>
<thead>
<tr>
<th>Sample</th>
<th>TaN conductivity (Ω-cm)⁻¹</th>
<th>TaN resistivity (μΩ-cm)</th>
<th>TaN TCR (ppm/°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JG50331</td>
<td>4820</td>
<td>207</td>
<td>–170</td>
</tr>
<tr>
<td>JG50401</td>
<td>5360</td>
<td>186</td>
<td>–135</td>
</tr>
<tr>
<td>JG50509</td>
<td>5920</td>
<td>169</td>
<td>–110</td>
</tr>
<tr>
<td>JG50510</td>
<td>6630</td>
<td>151</td>
<td>91</td>
</tr>
<tr>
<td>JG60427</td>
<td>4655</td>
<td>215</td>
<td>–93</td>
</tr>
<tr>
<td>JG60428</td>
<td>4400</td>
<td>227</td>
<td>–139</td>
</tr>
</tbody>
</table>

![Figure 5.—Conductivity of TaN/PdCr multilayer at room temperature versus PdCr/TaN thickness ratio.](image)
C. Multilayered Thin Film Multifunctional Sensor Application

To test the TaN/PdCr multilayer as multifunctional sensors, several multifunctional sensors were fabricated using the pattern shown in figure 7 with l/w = 447. The properties of the sensors are summarized in table 5, which also contains the TaN multifunctional sensor (JG40922) as a reference. The first multilayer sensor (JG51003) was fabricated using the ratio of PdCr to TaN of 0.32, based on the average calculated properties of TaN in table 4. The strain output is graphed in figure 8. The resistivity and TCR of the first sensor were considerably different than what was expected. The second run (JG60522) has a PdCr to TaN ratio of 0.86, which reflects the suspected incorporation of oxygen into the multilayer film through our sacrificial copper/nitric acid etch lithographic process that is used to pattern the sensors. The resistivity and gauge factor are lower, as expected with the increased PdCr, but the TCR was very erratic, varying between 0 and 30 ppm/°C over a temperature range of 20 to 120 °C. The erratic TCR was due to a failure of the thermocouple leads that forced the use of an external thermocouple in contact with the sample in the oven. The leads will need to be repaired to determine a more precise TCR value. If the average TCR of +15 ppm/°C is repeatable, the apparent strain sensitivity to temperature of +12 με/°C meets our goal of less than ±20 με/°C for static strain measurements, at least on this first level of testing. Further testing will need to be conducted to 700 and then to 1300 °C to validate the concept, similar to validation tests done on conventional high temperature flame-sprayed instrumentation.

<table>
<thead>
<tr>
<th>Sample</th>
<th>TaN thickness (Å) (±250Å)</th>
<th>PdCr thickness (Å) (±250Å)</th>
<th>TaN+TaO thickness (Å) (±250Å)</th>
<th>Sample resistivity (µΩ-cm) (±5 percent)</th>
<th>Sample TCR (ppm/°C) (±5 ppm/°C) (±5 percent)</th>
<th>Gauge factor (δR/R/ε) (±5 percent)</th>
<th>Apparent strain sensitivity (µε/°C) (±10 percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JG40922</td>
<td>39,000</td>
<td>----</td>
<td>----</td>
<td>259</td>
<td>-93</td>
<td>3.9</td>
<td>-24</td>
</tr>
<tr>
<td>JG51003</td>
<td>5095</td>
<td>3256</td>
<td>5554</td>
<td>198</td>
<td>-121</td>
<td>1.89</td>
<td>-64</td>
</tr>
<tr>
<td>JG60522</td>
<td>2900</td>
<td>4600</td>
<td>3450</td>
<td>146</td>
<td>15±15</td>
<td>1.23</td>
<td>12±12</td>
</tr>
</tbody>
</table>
IV. Conclusions

In order to have a more passive method of negating changes of resistance due to temperature, a process was developed to create a multilayer ceramic/metal thin film sensor of TaN and PdCr. The process was developed to use existing capabilities of the microfabrication cleanroom facility at NASA GRC, and to allow the fabrication of embedded sensors directly on components. The new sensor was tested on alumina substrates, since alumina is used as the typical insulator for thin film sensors on components. The sensor is capable of being fabricated in fine line widths utilizing the sacrificial-layer lift-off process that is used for microfabricated noble-metal sensors.

The TaN films were optimized for Ta$_2$N properties using reactive sputtering with an unbalanced magnetron source. A first approximation model of multilayer resistance and TCR was used to set the film thicknesses of PdCr and TaN in the multilayer film sensor. Two multifunctional sensors were fabricated using TaN/PdCr multilayered films, and tested for low temperature resistivity, TCR and strain response.
Though there were initial difficulties translating the large area multilayered film samples to a fine lined sensor, the sensors tested improve existing apparent strain sensitivity by at least 30 percent. The low thermal coefficient of resistance compared to existing high temperature strain gauges allows for the multifunctional sensor to be used in static measurements with little or no temperature compensation. The sensor is not attacked by acids as readily as metals, making it survivable in non-oxidizing harsh environments. The reactively-sputtered fabrication technique also allows for fairly quick fabrication times compared to other ceramic-based sensors. Further tests are required to validate the sensitivity, stability, repeatability, interference and durability of the concept.

References


### Developing Multilayer Thin Film Strain Sensors With High Thermal Stability

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#### Abstract
A multilayer thin film strain sensor for large temperature range use is under development using a reactively-sputtered process. The sensor is capable of being fabricated in fine line widths utilizing the sacrificial-layer lift-off process that is used for microfabricated noble-metal sensors. Tantalum nitride films were optimized using reactive sputtering with an unbalanced magnetron source. A first approximation model of multilayer resistance and temperature coefficient of resistance was used to set the film thicknesses in the multilayer film sensor. Two multifunctional sensors were fabricated using multilayered films of tantalum nitride and palladium chromium, and tested for low temperature resistivity, TCR and strain response. The low temperature coefficient of resistance of the films will result in improved stability in thin film sensors for low to high temperature use.