High-Q Gold and Silicon Nitride Bilayer Nanostrings

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Low-mass, high-Q, silicon nitride nanostrings are at the cutting edge of nanomechanical devices for sensing applications. Here we show that the addition of a chemically functionalizable gold overlayer does not adversely affect the Q of the fundamental out-of-plane mode. Instead the device retains its mechanical responsiveness while gaining sensitivity to molecular bonding. Furthermore, differences in thermal expansion within the bilayer give rise to internal stresses that can be electrically controlled. In particular, an alternating current (AC) excites resonant motion of the nanostring. This AC thermoelastic actuation is simple, robust, and provides an integrated approach to sensor actuation.

Silicon nitride nanostrings1,2 are an exciting class of mechanical resonator, exceptional for their ultra-high quality factors (Qs)3 their tolerance to irregularities in the fabrication process (since their resonant frequencies depend only on the device length)4 their harmonic normal mode spectrum, and the ease with which their vibrational motion can be detected optically (by virtue of the low absorption of silicon nitride at visible wavelengths).5,10 But in order for these devices to be used efficiently in room temperature molecular sensing applications,11–13 they should have a surface that is chemically functionalizable. Silicon nitride, however, is largely inert.

The most common workaround, routinely carried out with silicon and silicon nitride atomic force microscope (AFM) cantilevers, is to coat the resonator with gold. Then a molecule of choice14 can be affixed to the gold overlayer, attached by way of a thiol intermediary.15,16 The addition of the gold has no adverse consequences for static, stress-response measurements,15 but it generally does for measurements that are dynamic in nature. For example in the cantilever case, metallization causes significant dissipation in the resonator15,16 lowering its Q and reducing its usefulness as a sensor—although this can be circumvented by depositing only in areas that do not become stressed upon actuation.17

Accordingly, our initial expectation was that application of a uniform gold layer to a silicon nitride nanoribbon would ruin the device’s ultra-high Q, which is typically on the order of 105 but can be as high as 7 × 106 (see Ref. 3). Instead, we show here that a 53 nm thick metallic layer on top of a 250 nm thick silicon nitride nanoribbon does not adversely affect the Q of the fundamental frequency, even though it nearly doubles the total mass of the string. (Our longest gold-covered device, at 210 µm, has Q ≥ 1.6 × 109 for the fundamental mode and a total mass of ≈ 840 pg.) It does, however, reduce the Qs associated with higher harmonics, so that the Q–versus–resonant-frequency behavior that was previously flat (up to even-odd effects across modes) now decays with frequency. This change is characteristic of a string system whose dominant dissipation mechanism is no longer localized at the anchor points3

Another important feature of the gold layer is that it renders the string sensitive to temperature through the bimaterial effect (a differential stress between the gold and silicon nitride layers due to different coefficients of thermal expansion).20,21 While this could prove useful in temperature sensing22 or provide an independent measurement of the resonator temperature,23 our focus lies elsewhere. In particular, we demonstrate that it is possible not only to vary the temperature of the device electrically via direct current (DC) ohmic heating but to actuate resonant motion via alternating current (AC) thermoelastic heating. Both the DC and AC effects have practical uses for sensors: the first allows for device regeneration (i.e., it provides a mechanism to desorb molecules, returning the device to a clean state), which facilitates thermal studies of affixed molecules15,24 and, as recently shown,25 can provide exceptional frequency stability in ultra-low-noise nanomechanical sensing; the second provides integrated actuation26,27 and obviates the need for a piezoelectric buzzer or other external driving mechanism.

Fabrication of the bilayer nanostrings is straightforward, since only one lithography step is necessary. We start with stoichiometric silicon nitride (250 nm) deposited onto silicon dioxide (2 μm) on a silicon handle (Rogue Valley Microdevices). This is subsequently sputtered with 10 nm of chromium, followed by 43 nm of gold. (We refer to the resulting device as bilayer nanostring, since the intermediate chromium layer is thin and serves only an adhesion function.) Standard optical lithography is then performed. This is followed by wet etching of the gold and then chromium, reactive ion etching of the silicon nitride, and device release with a buffered oxide etch of the silicon dioxide. The resulting

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strings are shown in Fig. 1. The gold is slightly over-etched, although this is not important since the string’s normal modes are insensitive to the exact dimensions. This fault tolerance during fabrication is advantageous for bulk processing.

Measurements are performed by optical interferometry using a 632.8 nm laser focused onto the nanostring, as described elsewhere[4,5] inside an optical access vacuum chamber in which the temperature of the sample stage can be controlled and accurately determined. The optical power incident on the device is 13 µW—below the onset of optical heating effects. The interferometric signal is amplified 25-fold after detection and is analyzed using a high frequency lock-in amplifier (Zurich Instruments HF2LI), which allows us to extract the peak frequency and Q by fitting to the power spectral density function (PSD) (see supplementary material of Ref. 8). This analysis is appropriate regardless of the actuation method: thermomechanical, external piezoelectric, or integrated ohmic (thermoelastic). In addition, our optical system can be scanned with respect to the sample using closed-loop piezo stages,[33] providing spatial maps of the amplitude and phase output by the Zurich lock-in. Hence, we can unambiguously distinguish between in-plane and out-of-plane motion.

We emphasize that while we have verified the following results using numerous nanostrings of varying lengths, for consistency we present data for a single 210 µm long by 2.75 µm wide bilayer nanostring with an initial resistance of 176 Ω. The Qs of this bilayer nanostring in vacuum (≈2 × 10⁻⁶ torr) are shown in Fig. 2. These Qs are compared with out-of-plane modes of uncoated strings of a very similar geometry (red squares) from Ref. 8. It is immediately clear that the Q of the fundamental out-of-plane mode is essentially unchanged (although the resonant frequency itself is reduced by about 25% because of the addition of the gold). This device is therefore an excellent candidate for a functionalizable, high-Q nanomechanical sensor.

On the other hand, the higher order modes are damped as compared to the bare string. Both bare and gold bilayer devices demonstrate the alternating even/odd behavior previously seen for silicon nitride membranes[23] and nanostrings[8] which has been reported to originate from dissipation due to phonon tunneling.[23] Ignoring this oscillation, the Q of the bare device is essentially flat with respect to frequency (or, equivalently, mode number). The Q of the gold bilayer strings, however, decays with frequency. A key difference appears to be the high thermal conductivity of the metallic layer, which allows for efficient transport of energy from within the string (dissipated as a result of internal stresses during bulk bending) to the outside.

As we have previously shown[8] the Q varies by mode according to

$$Q_n = \frac{m \omega_n}{\gamma_n} \approx \frac{m}{\gamma_{\text{visc}} \omega_n - \gamma_{\text{visc}} + \gamma_{\text{bulk}} \omega_n} \quad (1)$$

The tilde decorated quantities are damping coefficients from which the leading order frequency dependence has been factored out. The viscous damping term, $\gamma_{\text{visc}}$, can be neglected since the experimental chamber has been evacuated far past the point where the measured Q exhibits a pressure dependence. In the case of the bare string, $\gamma_{\text{anchor}}$ is the overwhelmingly dominant contribution[8] energy is lost primarily in dissipative processes in the vicinity of the anchor points, and the Q has no frequency dependence other than the even/odd mode effect. The Q of the gold bilayer string, however, shows a decaying tail that can only be fit with a non-vanishing $\gamma_{\text{bulk}}$ term, attributable to bulk dissipation. Specifically, we find from the out-of-plane data in Fig. 2 that $\omega_1 \gamma_{\text{bulk}} / \gamma_{\text{anchor}} = 0.38 \pm 0.08$. Determination of the microscopic bulk dissipation mechanism would require further detailed studies[25]. We remark that the Qs of the higher order modes are still rather impressive ($> 10^6$), suggesting that this device can be used for multimode sensing[25,26].

Beyond altering the dissipation, the gold overlayer adds functionality to the nanostring system by introducing bimaterial temperature dependence. In particular, we have observed that the resonant frequencies of the nanostring shift with heating. There are two scenarios to consider: (i) the sample stage is held in thermal equilibrium at a uniform temperature $T = T_{\text{room}} + \Delta T$; (ii) heating is highly localized in the string, and the rest of the system—including the supports and substrate—can be viewed as a heat bath at $T_{\text{room}}$. The second scenario corresponds to the Joule heating case in which a current is passed through the metallic overlayer.

In the first scenario, the frequencies have the form

$$f_n = \frac{\omega_n}{2\pi} = \frac{n}{2L} \sqrt{\frac{A_1 \sigma - \sum_{k=1}^{3} A_k E_k \alpha_{k,0} \Delta T}{\sum_{k=1}^{3} A_k p_k}}, \quad (2)$$

where $\alpha_{k,0} = \alpha_k - \alpha_0$, and the parameters $E_k$, $A_k$ are the corresponding Young’s moduli and cross-sectional areas. (See Ref. 43 and references therein.) These material parameters are defined for each layer with the indices $k = 1$ (silicon nitride), $k = 2$ (chromium), and $k = 3$ (gold).

For small temperature changes, it is sufficient to consider

![FIG. 2. Q as a function of mode number for the 210 µm long gold bilayer nanostring (orange circles are out-of-plane and blue triangles are in-plane) and a bare string of similar geometry (red squares) from Ref. 8. Here the string is actuated using the external piezoelectric. The blue and orange curves are fits to Eq. (1) with $\gamma_{\text{visc}} = 0.$](image-url)
The simple analysis we have outlined is primarily limited by the fact that we have supposed the temperature profile $\Delta T(x, t) = \Delta T(t)$ to have no meaningful variation along the length of the string. Still, it reliably captures the fact that the thermal time scale is on the order of microseconds (rather than nanoseconds, as for the device described in Ref. [3]), orders of magnitude faster than the time for the mechanical energy to be transferred to the environment, as dictated by the high $Q$ of the string: $Q/f_1 \approx 200 \, \text{ms}$.

In our AC local heating scenario, the power pumped into the string is given by $P = \dot{V}^2/R$, where $R$ is the string’s electrical resistance and $\dot{V}(t) = V_0 \cos \omega t$ is the alternating voltage applied across the string. Since the silicon nitride is insulating ($R_1 \approx \infty$), we can estimate the resistance from the geometry of the device using standard values $r_2$, $r_3$ for the resistivity of Cr and Au: $R \approx (\sum_{k=1}^{3} 1/R_k)^{-1} \approx L(A_2/r_2 + A_3/r_3)^{-1} = 49 \, \Omega$. The 176 $\Omega$ we measure experimentally is higher, mainly because it includes the resistance of the metallic supports. Accordingly, the heat content of the string (ignoring the initial transient) varies in time as

$$U(t) = \left( \frac{V_0^2 \tau_\text{th}}{2R} \right) \left[ 1 + \frac{\cos(2\omega t - \phi)}{(1 + 4\omega^2 \tau_\text{th}^2 \phi^2) \cos \phi} \right]$$

with a phase lag $\phi = \tan^{-1}(2\omega \tau_\text{th})$ that depends only on the thermal outflow time $\tau_\text{th}$. We can decompose the corresponding temperature variation into a time-invariant background $\Delta T_{\text{ave}} = V_0^2 \tau_\text{th}/2CR$ over which is superposed the weaker, alternating behaviour $\Delta T_{\text{alt}}$. Hence, the analogue energy $\dot{U}$ comes into equilibrium when $\dot{U} = P - \dot{U}/\tau_\text{th} = 0$. Here, $\tau_\text{th}$ represents the half-life for thermal energy to leak out of the string. The energy outflow can be computed as a sum of thermal currents (of the form $j = -\kappa \nabla T$) driven by a gradient $\nabla T \approx \Delta T/(L/2)$. We make use of the device’s overall heat capacity $C = L \sum_{k=1}^{3} A_k \rho_k c_k \approx 3.70 \times 10^{-10} \, \text{J/K}$ to relate the change in heat energy to a corresponding change in temperature and thus to produce an estimate of

$$\tau_\text{th} = \frac{L^2 \sum_{k=1}^{3} A_k \rho_k c_k}{2 \sum_{k'=1}^{3} A_{k'} \kappa_{k'}} \approx 750 \, \mu\text{s}.$$  \hspace{1cm} (4)

(Values of the density $\rho_k$, specific heat $c_k$, and thermal conductivity $\kappa_k$ are tabulated per layer in the supplementary material[5]) Note that $\tau_\text{th}$ scales as $L^2$, since the equilibration process is essentially diffusive; this implies that, near resonance, $\omega \tau_\text{th} \propto L$. Because the layers themselves come into local thermal equilibrium on a fast time scale of around $Ct_1/Lw_1 \kappa_1 \approx 5 \, \text{ns}$ (where $t_1$ and $w_1$ are the thickness and width of the silicon nitride respectively), we proceed as if the silicon nitride, chromium, and gold layers are at the same temperature instantaneously at each point along the string. This is a safe assumption up to a frequency scale $1/(2\pi \cdot 5 \, \text{ns}) \approx 32 \, \text{MHz}$, which is well above the oscillation frequency of any of the harmonic modes that we can measure in our device.
\[ f_n(\Delta T) - f_n(0) \approx -\frac{V_0^2 \tau_{th}^2}{4\sigma A_1 CR} \sum_{k=1}^{3} A_k E_k a_k \]
\[ = -\frac{L}{8\sigma A_1 R} \left( \sum_{k'=1}^{3} A_{k'} E_{k'} a_{k'} \right) V_0^2. \]

The relative frequency shift is quadratic in \( V_0 \) and always negative—consistent with our observations in Fig. 2(c). In contrast to the relative frequency shift described by Eq. 3, which is insensitive to the device length, the effect described in Eq. 6 is predicted to scale as \( \tau_{th}/C \sim L \) (and thus can be made arbitrary large by choosing a long enough string). The predicted coefficient, 6.82 V\(^{-2}\), is about ten times larger than the value 0.648(9) V\(^{-2}\) obtained from a fit to the data. Theory and experiment can be made to coincide if instead we take the thermal time to be \( \tau_{th} = 72 \mu s \). The discrepancy likely stems from the over-simplified, uniform temperature profile along the length of the string.

In the vicinity of the fundamental mode of our device, \( 2\omega_1 \tau_{th} = 4\pi f_1 \tau_{th} \approx 708 \gg 1 \), so that \( \phi \) has essentially achieved its maximum value, \( \pi/2 \). In this limit, \( (\Delta T)_{th} = (V_0^2/2CR\omega) \sin(2\omega t) \), and the magnitude of the alternating component of the temperature with respect to the average goes as \( 1/\omega \tau_{th} \ll 1 \). What’s astonishing is that this tiny alternating local heating is sufficient to actuate the device and to do so very efficiently. Specifically, we are able to use this local heating to actuate the bilayer nanostring by passing an AC current along the length of the device. Note that this is subtly different from the bimaterial effect in cantilevers and non-uniformly coated beams, where the gold exerts a force on the cantilever that pushes it towards the substrate\(^{29,30}\). In such a scenario, the force is directional, and hence the AC drive voltage must be applied at frequency \( f/2 \) in order to produce heating and bending at \( f \). In contrast, in our system there is no preference towards or away from the substrate as the gold expands, since it forms a continuous layer over the string. Heating and bending occur twice per string oscillation, and therefore the applied drive voltage and resulting motion have the same frequency. We are also able to actuate the device at \( f/2 \), although \( \approx 1000 \) times less efficiently—consistent with the above scenario.

In Fig. 2(b), we show frequency sweeps using the reference output of the Zurich lock-in amplifier to resonantly drive the nanostring. This integrated electronic actuation scheme is simple, compact, and robust. With root-mean-square AC voltages as small as 100 \( \mu V \) we can clearly detect resonant motion, while the maximum signal-to-noise is achieved with roughly 15 to 20 mV. At higher drives the nanostring becomes nonlinear. Yet even at quite high drives, in vacuum, we see no change in the resistance of the gold layer. Hence the local heating of the bilayer nanostring is nondestructive to the gold layer in the linear actuation regime and beyond. In addition, we note that the resonance frequency is entirely non-hysteretic at these actuation voltages—further evidence that the AC ohmic actuation scheme is robust.

Finally we note that the AC ohmic actuation technique works for both higher-order harmonics in vacuum (not shown) as well as for the fundamental modes in air. In the ambient pressure case, the nanostrings experience strong viscous damping, especially when the motion is perpendicular to the string’s substantial width. The \( Q \) of the first out-of-plane mode is only 40 while that of the in-plane mode is 160 [see Fig. 4(a)]. There is a greater cross-section for molecular collisions, and thus energy loss, in the out-of-plane direction. The voltages required to actuate the nanostring in air (\( \approx 400 \) mV) result in resistance changes and eventual failure. Improving the \( Q \) in viscous environments\(^{29,30}\) would decrease the required actuation voltages and prevent such failure.

In conclusion, gold bilayer nanostrings are an enhancement to the already exciting system of silicon nitride nanostrings. They enable chemical functionalization and molecular detection, provide a scheme for local heating to study thermal response of molecules or desorb unwanted molecules, and provide simple and integrated actuation. And unlike traditional micromechanical resonators\(^{13}\), they maintain their high quality factors even with a continuous metallic coating.

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SUPPLEMENTARY MATERIAL

Device geometry

The device is shown in cross section in Fig. [S1]. Each of the silicon nitride ($k = 1$), chromium ($k = 2$), and gold ($k = 3$) layers is of constant thickness $t_k$ and has an area $A_k$ and a bending moment $I_k$ given by

$$A_k = \int_{\zeta_{k-1}}^{\zeta_k} d\zeta w(\zeta),$$
$$I_k = \int_{\zeta_{k-1}}^{\zeta_k} d\zeta (\zeta - \bar{\zeta})^2 w(\zeta).$$

The limits of integration are the cumulative layer thicknesses

$$\zeta_k = \sum_{l=1}^{k} t_l.$$ 

The bending axis of the composite string is fixed at a height $\bar{\zeta} = 136 \text{ nm} = 0.54 t_1$ (measured from the bottom surface) that minimizes the total bending constant $\sum_{k=1}^{3} E_k I_k/(1 - v_k^2)$. Hence, the $I_k$ values themselves depend indirectly on the Young’s modulus $E_k$ and Poisson ratio $\nu_k$ of the material in each layer.

Based on our best understanding of the wet- and dry-etch steps in the fabrication process, we propose that the layered
The values \( w_1 = 2.75 \mu m \) and \( w_3 = 2.18 \mu m \) are estimated from a (top-down) scanning electron microscopy image of the device. We use a pixel-contrast histogram analysis to perform edge detection. Finally, the areas and bending moments have the following values:

\[
(A_1, A_2, A_3) = (68.8, 2.16, 9.08) \times 10^{-14} m^2,\\
(I_1, I_2, I_3) = (36.6, 3.08, 19.4) \times 10^{-28} m^4.
\]  

The device geometry and dimensions we have indicated correspond to a device of total mass 841 pg and heat capacity 3.70 \( \times 10^{-10} \) J/K.

**Normal Modes**

The layered beam is described by a Lagrangian

\[
\mathcal{L} = \int_{0}^{L} dx \left( \frac{M}{2L} (\partial_t z)^2 - U[z(x,t)] \right),
\]  

where \( M/L = \sum_{k=1}^{3} \rho_k A_k \) is the (uniform) mass per unit length and

\[
U = \frac{1}{2} \sigma A_1 (\partial_x z)^2 + \frac{1}{8} \sum_{k=1}^{3} E_k A_k (\partial_x z)^4
+ \frac{1}{2} \sum_{k=1}^{3} \frac{E_k I_k}{1 - \nu_k^2} (\partial_z^2 x)^2
\]  

is the elastic energy density (including tension, elongation, and bending terms) associated with its transverse deformation. \( U \) penalizes any deviation from the equilibrium position \( z(x,t) = 0 \). At this stage, we ignore thermal expansion effects and restrict ourselves to a model that satisfies \( U[z(x,t)] = U[z(L - x,t)] \) and \( U[z(x,t)] = U[-z(x,t)] \). In other words, we assume that the system is invariant under the transformations \( z(x) \rightarrow z(L - x) \) and \( z(x) \rightarrow -z(x) \), even though the latter symmetry can in principle be broken by the layered structure.

We follow the usual prescription

\[
\frac{d}{dt} \frac{\delta \mathcal{L}}{\delta (\partial_t z(x,t))} = \frac{\delta \mathcal{L}}{\delta z(x,t)}
\]  

(S.6)

and make use of the variational identities

\[
\frac{\delta}{\delta z(x,t)} \int dx (\partial_t z)^2 = -2 \partial_z^2 z
\]

(S.7)

\[
\frac{\delta}{\delta z(x,t)} \int dx (\partial_t z)^4 = -12 (\partial_z^2 z)^2 \partial_z^2 z
\]

(S.8)

to obtain the equation of motion

\[
M \partial_z^2 z = \tau \partial_z^2 z + \frac{3}{2} S (\partial_z^2 z)^2 \partial_z^2 z - B \partial_z^2 z.
\]

Here, \( \tau = LA_1 \sigma \) is the tension in the composite string, and \( S = L \sum_{k=1}^{3} E_k A_k \) and \( B = L \sum_{k=1}^{3} E_k I_k/(1 - \nu_k^2) \) are its stretching and bending constants. It is sufficient to treat \( (\partial_z^2 z)^2 \partial_z^2 z \), the term that is nonlinear in \( z \), at the mean field level. We do so by replacing \( (\partial_z^2 z)^2 \) by its spatially averaged value

\[
\mu = \frac{1}{L} \int_{0}^{L} dx (\partial_t z)^2.
\]  

(S.9)

Equation (S.8) differs from a pure wave equation by terms that are proportional to the Young’s moduli. The relevant situation here is when these are sufficiently small that the behavior of the system is almost perfectly string-like. When the system is driven in the \( n \)th pure string mode \( z_n(x,t) = a_n \sin(n \pi x/L) e^{i \omega_n t} \), it oscillates at an angular frequency

\[
\omega_n^2 = \frac{1}{M} \left[ \frac{n^2 \pi^2}{L^2} \left( \tau + \frac{3}{2} S \mu \right) + \frac{n^4 \pi^4}{L^4} B \right].
\]  

(S.10)

For weak nonlinearity, the mean square displacement \( \langle a_n^2 \rangle \sim \mu \) is small compared to \( L^2 \tau/S \). In that limit,

\[
f_n^2 = \frac{1}{4M} \left( \frac{n^2 \tau}{L^2} + \frac{n^4 \pi^2}{L^4} B \right).
\]  

(S.11)

Accordingly, high order modes exhibit a small \( n^3 \) correction away from a perfectly harmonic spectrum:

\[
f_n \approx \frac{n}{2L} \sqrt{\frac{\tau}{M} + \frac{n^3 \pi^2}{4L^3} \frac{B}{\sqrt{\tau M}}}.
\]  

(S.12)
TABLE S-I. Material properties of the layered nanostring and substrate. Unless otherwise specified, reference values for the bulk material are used (drawn from CRC Handbook of Chemistry and Physics, 92nd Edition).

| layer (#) | density (g/cm³) | Young’s modulus (GPa) | Poisson ratio | thermal expansion α (×10⁶ K⁻¹) | specific heat c (J/kg·K) | resistivity ρ (Ω·m) | thermal conductivity κ (W/m·K) |
|-----------|-----------------|-----------------------|---------------|---------------------------------|--------------------------|---------------------|--------------------------------|
| Au (3)    | 19.30           | 79                    | 0.44          | 14.2                            | 129.1                    | 22.14               | 318                           |
| Cr (2)    | 7.19            | 279                   | 0.21          | 4.9                             | 448.2                    | 125                 | 93.9                          |
| Si₃N₄ (1) | 3.00            | 265                   | 0.20          | 1.27a                           | 700                      | ∞                   | 30                            |
| Si (0)    |                 |                       |               |                                 |                          |                     |                                |

*a* determined by multimode calibration measurements  
*b* from Ref. 41  
*c* from Ref. 42

**Thermal Expansion**

Expansion of the layers due to heating leads to a correction that enters the elongation term of the elastic energy density:

\[
U = \frac{1}{2} A_1 \sigma (\partial_x z)^2 + \frac{1}{8} \sum_{k=1}^{3} E_k A_k \left[ (\partial_x z)^2 - 2 \alpha_k \Delta T \right]^2 + \frac{1}{2} \sum_{k=1}^{3} \frac{E_k I_k}{1 - \nu_k^{-2}} (\partial_x^2 z)^2. \tag{S.13}
\]

Note that we are not attempting to model interfacial shearing and peeling stresses. We just assume that the layers are held together by a sufficiently large through-thickness spring constant. Expanding Eq. \( \text{(S.13)} \) to give

\[
U = \text{const} + \frac{1}{2L} \left[ \left( \tau - L \sum_{k=1}^{3} E_k A_k \alpha_k \Delta T \right) (\partial_x z)^2 + B(\partial_x^2 z)^2 + \frac{1}{4} S(\partial_x z)^4 \right], \tag{S.14}
\]

we see that the effect of heating is simply to shift the tension. Applying this observation to Eq. \( \text{(S.11)} \) gives

\[
\frac{f_n(\Delta T) - f_n(0)}{f_n(0)} = -\frac{1}{8ML(f_n/n)^2} \sum_{k=1}^{3} E_k A_k \alpha_k \Delta T \\
\approx -\frac{1}{2 \sigma A_1} \sum_{k=1}^{3} E_k A_k \alpha_k \Delta T. \tag{S.15}
\]

The last line above is correct to the extent that \( f_n/n \approx f_1 \).

**Material properties**

The properties of silicon nitride thin films are quite different from those of the bulk material. For example, the density and coefficient of thermal expansion are generally lower. So we have taken care to find values that are appropriate to our particular devices.

For the purposes of calibration, we prepare a bare silicon nitride nanostring with no metallic coating. The dimensions of this device are 215 μm (length), 2.1 μm (width), and 250 nm (thickness). The frequencies (in MHz) of the first six resonant modes are determined to be 1.19508, 2.38389, 3.58222, 4.77447, 5.98263, and 7.19362. In this case, Eq. \( \text{(S.11)} \) reduces to

\[
(2L f_n)^2 = n^2 \frac{\sigma}{\rho} + n^4 \frac{E}{\rho} \frac{(t/L)^2}{12(1-\nu^2)} = n^2 \frac{\sigma}{\rho} + n^4 \frac{E}{\rho} 1.16 \times 10^{-6}. \tag{S.16}
\]

A best fit of this form to the data produces

\[
\frac{\sigma}{\rho} = 2.622(3) \times 10^5 \text{ m}^2/\text{s}^2, \\
\frac{E}{\rho} = 8.7(8) \times 10^7 \text{ m}^2/\text{s}^2, \tag{S.17}
\]

and hence \( E/\sigma = 330(30) \). We have no direct measurement that closes this set of relations, but if we assume the manufacturer’s quoted value of 0.8 GPa for the tensile stress, then \( E = 265 \) GPa and \( \rho = 3.05 \text{ g/cm}^3 \). These values are consistent with other measurements in the literature.[41]

Furthermore, we obtain the relative frequency shift for each of modes 1, 3, and 5. The measurement is performed over a dense grid of temperature values from 293 K to 320 K and produces a linear slope

\[
\frac{E(\alpha_0 - \alpha_1)}{2 \sigma} = 0.000219(1) \text{ K}^{-1} \tag{S.18}
\]

that is consistent across all three modes. Taking \( \alpha_0 = 2.59 \times 10^{-6} \text{ K}^{-1} \) as the coefficient of thermal expansion for the silicon substrate,[42] we find \( \alpha_1 = 1.27 \times 10^{-6} \text{ K}^{-1} \). This is very close to the value \( 1.23 \times 10^{-6} \text{ K}^{-1} \) reported in Ref. [43]