Experimental signatures of the quantum–classical transition in a nanomechanical oscillator modeled as a damped-driven double-well problem

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Abstract
We demonstrate robust and reliable signatures for the transition from quantum to classical behavior in the position probability distribution of a damped double-well system using the quantum state diffusion approach to open quantum systems. We argue that these signatures are within experimental reach, for example in a doubly clamped nanomechanical beam.

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(Some figures may appear in color only in the online journal)

1. Introduction

The difference between quantum and classical behavior is of fundamental interest with many experimental consequences in contexts such as superconducting quantum interference devices, cold atom systems and significantly in nanoelectromechanical systems (NEMS). NEMS are small, oscillate at high frequencies and are typically maintained at low temperatures. Most of their parameters are adjustable, making them an ideal setting for exploring the transition from quantum to classical behavior. Recent experimental results [1, 2] indicate that such systems can now be prepared and observed in their ground state, thus enabling the study of genuinely nonclassical behavior in the dynamics. The exciting prospect of being able to observe the quantum–classical transition in detail has contributed to interest in quantum-mechanical NEMS (occasionally referred to as QEMS in the literature), whose study is an active and rapidly expanding field of research [3].

A seminal theoretical study of the quantum–classical transition in NEMS was carried out by Katz et al [4] and Carr et al [5] using a nonlinear resonator, which is a paradigmatic model for several common types of NEMS that yields quantum behavior distinct from the classical. They considered an isolated resonator as well as one modeled as an open system (coupled to the environment) in search of experimental signatures of such quantum behavior. Specifically, they examined the dynamics of the classical phase space distribution in such a system compared to that of the quantum-mechanical Wigner function, and looked at a zero-temperature and a finite-temperature environment. They were able to find differences in experimentally accessible signatures: in particular, the quantum-mechanical version of the resonator has nonzero probability of being found in a position where the classical resonator has zero probability of being found. However, these differences are relatively small and blur away quickly when experimental noise or finite-temperature effects through thermal noise are considered.

In this paper, by considering a slightly different but equally accessible nonlinear oscillator, we are able to obtain more pronounced experimental signatures of the quantum–classical transition; these are arguably more visible at finite temperatures and in the presence of experimental noise. Further, we show that rather than comparing just quantum and classical behavior, there is insight to be gained from studying the continuous transition where the classical behavior emerges as the limiting case of the quantum
dynamics. That is, although systems are fundamentally described by quantum mechanics, their behavior changes as we increase the size of the system (affecting the characteristic action scaled by \( \hbar \)), the effect of the environment (decoherence) or other parameters of the system. For our system, we indeed recover the classical behavior as the limiting case. The details of the emergence of the classical limit are illuminating, and in particular, the signatures do not always change monotonically, thus establishing that comparing a single quantum solution with a classical solution leads to an incomplete picture.

Katz et al considered the system with the Hamiltonian

\[
H_{\text{sys}} = \frac{1}{2} p^2 + \frac{1}{2} x^2 + \frac{1}{2} x^4 - x F \cos \omega t. \tag{1}
\]

This quadratic nonlinear oscillator is within experimental reach in the near-quantum regime, and is an excellent system to be studied. However, even more interesting effects such as quantum tunneling are obtained if we change the sign so that the \( x^2 \) term is negative [6]. This yields the so-called double-well Duffing oscillator, which has the Hamiltonian \( H_{\text{sys}} = \frac{1}{2} p^2 - \frac{1}{2} x^2 + \frac{1}{2} x^4 - x F \cos \omega t \) and allows for clearer signatures of the classical–quantum transition as we demonstrate below.

This paper is organized as follows. We start by motivating the modeling of a specific NEMS as a doubly clamped beam to show the connection between experimental parameters and model parameters. We then sketch the quantum state diffusion (QSD) approach, where the Lindblad operators act within the stochastic Schrödinger equations to incorporate the effect of the environment, utilized to model the behavior of such an oscillator understood as an open quantum system. These two allow us to establish that changing the sign of the quadratic term is within current experimental capabilities and that such experiments are in or close to the realm where the quantum–classical transition can be explored. We then present the results and conclude with our analysis.

2. NEMS modeled as a doubly clamped beam

Many nanoelectromechanical devices can be modeled as doubly clamped beams driven near resonance, and continuum mechanics continues to serve as an adequate model even at the submicron scale [7–9]. Usually, magnetomotive [10] or optical [7] actuation is used to study the resonance behaviors of NEMS such as dynamically induced bistability, hysteresis and effects of parameter attenuation on energy dissipation. The resonant behavior of the doubly clamped NEMS structure has been experimentally identified with the fundamental bending mode [11] of the problem. We sketch here the theory allowing us to relate the resonance frequency to characteristic parameters of NEMS and to determine the scale of the system with respect to \( \hbar \), establishing the regime of the transition to classical behavior in these systems.

Starting with the assumption of an ideal beam of rectangular cross-sectional area, we can obtain the potential energy \( V \) of a doubly clamped beam under tension along the longitudinal direction. This consists of two parts [12]: elastic and bending potential energy,

\[
V_{\text{elastic}} = T \left[ \int_0^{l_0} \sqrt{1 + (y')^2} \, dx - l_0 \right] \tag{2}
\]

and

\[
V_{\text{bending}} = \frac{EI}{2} \int_0^{l_0} (y'')^2 \, dx. \tag{3}
\]

\( T \) is the external force on the beam in the form of a tension in the longitudinal direction along the beam, \( E \) is the elastic modulus, \( I \) is the moment of inertia of the cross-section and \( l_0 \) is the separation of two ends that clamp the beam. Note that \( l_0 \) need not be the length of the beam when it is neither stretched nor compressed because a static doubly clamped beam may already be buckled due to inherent tension along the beam. \( V_{\text{elastic}} \) is due to tension built up in the beam, which causes the elastic beam to deform and therefore gain energy. This is usually referred to as strain energy [13], while \( V_{\text{bending}} \) is purely due to bending. The quantity \( E I \), termed ‘flexural rigidity’, denotes the force necessary for bending a beam by a unit of curvature.

The tension \( T \) along the beam has two parts [14]: firstly, an inherent tension \( T_0 \) (either from compression at both ends of the beam or the manufacturing process) and an additional tension \( \Delta T \) (due to transverse motion that stretches the beam), whence \( T = T_0 + \Delta T \). When the beam is stretched, its length changes and therefore the additional tension \( \Delta T \) is given by a fractional change in length multiplied by the elastic modulus and the cross-section area \( A \) [14]:

\[
\Delta T = \frac{T_0 - l_0}{l_0} E A, \tag{4}
\]

where \( l_0 \) refers to the actual length of the buckled beam. We can rewrite this length as

\[
l = l_0 + \Delta l = \int_0^{l_0} \sqrt{1 + (y')^2} \approx \int_0^{l_0} \left(1 + \frac{1}{2}(y')^2 \right) \, dx,
\]

\[
= l_0 + \frac{1}{2} \int_0^{l_0} (y')^2 \, dx. \tag{4}
\]

This yields that the change in tension \( \Delta T = \frac{E A}{2l_0} \int_0^{l_0} (y')^2 \, dx \), whence the total tension

\[
T = T_0 + \frac{E A}{2l_0} \int_0^{l_0} (y')^2 \, dx. \tag{5}
\]

Substituting these expressions for \( \Delta l \) and \( T \) in the expression for elastic potential energy above, we obtain

\[
V_{\text{elastic}} = \frac{T_0}{2} \int_0^{l_0} (y')^2 \, dx + \frac{EA}{4l_0} \left[ \int_0^{l_0} (y')^2 \, dx \right]^2. \tag{6}
\]

Thus, the total potential energy is

\[
V = V_{\text{elastic}} + V_{\text{bending}}
\]

\[
= \frac{T_0}{2} \int_0^{l_0} (y')^2 \, dx + \frac{EA}{4l_0} \left[ \int_0^{l_0} (y')^2 \, dx \right]^2 + \frac{EI}{2} \int_0^{l_0} (y'')^2 \, dx. \tag{7}
\]

At low temperatures, as is often true for NEMS experiments, it is sufficient to consider only the fundamental mode. Mode amplitude solutions for this fundamental mode \( y(x) \) of the form [15]

\[
y(x) = \frac{1}{2} Y \left[ 1 - \cos(2\pi \frac{x}{l_0}) \right], \tag{8}
\]

where \( Y \) is the transverse displacement of the center point of the beam. When we make this substitution, the fundamental mode amplitude \( Y \)—that is, the location of the vibrating central point—is
effectively the position variable of a single particle evolving in a potential \( V \), which as a function of \( Y \) is given by
\[
V(Y) = \frac{EI\pi^4}{l_0^4} Y^2 + \frac{\pi^2 T_0}{4l_0^2} Y^2 + \frac{EA\pi^4}{16l_0^4} Y^4
= Y^2 \left( \frac{EI\pi^4}{l_0^4} + \frac{\pi^2 T_0}{4l_0^2} \right) + \frac{EA\pi^4}{16l_0^4} Y^4.
\]
(8)

Here \( I \), the moment of inertia of the rectangular cross-section, is \( \frac{4}{15} \bar{a} \), where \( a \) is the thickness of the beam and \( b \) its width. \( I \) varies with the geometry of the beam for specific experimental setups. Further, depending on whether the inherent tension \( T_0 \) compresses or stretches the beam, the sign of \( T_0 \) can be negative or positive. Compressive inherent tension along the beam can arise from manufacturing processes, and can vary from 200 MPa to 1 GPa [15]. In general, the inherent tension is not known \( a \) priori in an experimental setup. Instead, by considering the continuum mechanics model and fitting the experimentally measurable resonance frequency, the inherent tension can be calculated [11]. For a sufficiently large inherent compressive tension, \( T_0 < -\frac{2\pi^2 \bar{E} \bar{I}}{l_0^4} \), we obtain a double-well potential and the beam is in a state of ‘Euler instability’ [9, 17], meaning that it is buckled.

Thus, a single-well potential or a double-well potential can be experimentally achieved in doubly clamped NEMS structures and these systems can then be studied for their dynamical behavior. The dynamics of a single-well NEMS, actuated near the fundamental bending mode frequency has been extensively studied [18, 19] but that of the double-well case has not yet been fully explored. The results of [11] suggest that it is within experimental reach and the above clarifies that, in general, tuning the characteristic parameters should enable switching from a single-well to a double-well potential.

3. Open quantum systems

We now turn to the quantum version of the model for the problem and connect characteristic parameters with the degree of ‘quantumness’ of the system. An open quantum system with weak system–environment coupling and a Markovian environment is modeled by a master equation
\[
\dot{\rho}(t) = \frac{-i}{\hbar} \left[ \hat{H}, \rho \right] - \frac{1}{2} \sum_j (\hat{L}_j \rho \hat{L}_j^\dagger + \rho \hat{L}_j^\dagger \hat{L}_j) + \sum_j \hat{L}_j \rho \hat{L}_j^\dagger,
\]
(9)

where \( \hat{L}_j \) is the Lindblad operator representing the system–environment interaction and \( \hat{\rho} \) is the reduced density operator. The QSD approach corresponds to solving quantum trajectories which are unravellings of the master equation. This allows for numerical efficiencies, and physical insights not available via the master equation. Specifically, we can use the QSD numerical library [20] to solve a stochastic version of the Schrödinger equation
\[
d|\psi(t)\rangle = \frac{i}{\hbar} \hat{H} |\psi(t)\rangle dt
+ \sum_j \left( \hat{L}_j^\dagger \hat{L}_j \frac{1}{2} - \frac{1}{2} \hat{L}_j^\dagger \hat{L}_j \langle \hat{L}_j \rangle \right) |\psi(t)\rangle dt
+ \sum_j \left( \hat{L}_j - \langle \hat{L}_j \rangle \right) |\psi(t)\rangle d\xi_j.
\]
(10)

The solution \( |\psi(t)\rangle \) to equation (10) is a ‘quantum trajectory’, and we obtain \( \hat{\rho} = \frac{1}{M} \sum_i |\psi_i\rangle \langle \psi_i| \), as the mean over an ensemble of \( M \) normalized pure state projection operators [20]. This \( \hat{\rho} \) satisfies the Lindblad master equation (9). That is, starting with an ensemble of identical pure states, we obtain the time evolution of multiple trajectories that evolve into an ensemble of different pure states due to interaction with the environment; this is the density matrix.

For the open quantum double-well Duffing oscillator, \( \hat{H} \) and \( \hat{L}_j \) in equation (10) are chosen as [4, 5, 21–23]
\[
\hat{H} = \hat{H}_D + \hat{H}_R + \hat{H}_ex,
\]
(11)
\[
\hat{H}_D = \frac{1}{2m} \hat{p}^2 + \frac{m\omega_0^2}{4l_0^2} \hat{x}^2 - \frac{m\omega_0^2}{2} \hat{x}^2,
\]
(12)
\[
\hat{H}_R = \frac{\gamma}{2} \left( \hat{x} \hat{p} + \hat{p} \hat{x} \right),
\]
(13)
\[
\hat{H}_ex = -gml\omega_0^2 \hat{x} \cos(\omega t),
\]
(14)
\[
\hat{L}_1 = \sqrt{\gamma} \left( 1 + \bar{n} \right) \left( \sqrt{\frac{m\omega_0}{\hbar}} \hat{x} + i \sqrt{\frac{\gamma}{m\omega_0\hbar}} \hat{p} \right),
\]
(15)
\[
\hat{L}_2 = \sqrt{\gamma} \left( 1 - \bar{n} \right) \left( \sqrt{\frac{m\omega_0}{\hbar}} \hat{x} - i \sqrt{\frac{\gamma}{m\omega_0\hbar}} \hat{p} \right),
\]
(16)
where \( \gamma \) is the strength of coupling between the oscillator and the thermal bath environment. We now define \( \hat{x} \) and \( \hat{p} \) in unitless forms \( \hat{Q} = \sqrt{m\omega_0/\hbar} \hat{x} \) and \( \hat{P} = \sqrt{m\omega_0\hbar} \hat{p} \), respectively [22, 23], yielding the dimensionless set of equations
\[
\dot{\hat{H}}_D = \frac{1}{2} \hat{P}^2 + \frac{\bar{q}^2}{4} \hat{Q}^4 - \frac{1}{2} \hat{Q}^2,
\]
(17)
\[
\dot{\hat{H}}_R = \frac{\Gamma}{2} \left( \hat{Q} \hat{P} + \hat{P} \hat{Q} \right),
\]
(18)
\[
\dot{\hat{H}}_ex = -\frac{g}{\bar{p}} \hat{Q} \cos(\Omega t),
\]
(19)
\[
\dot{\hat{L}}_1 = \sqrt{1 + \bar{n}} \left( \hat{Q} + i \hat{P} \right),
\]
(20)
\[
\dot{\hat{L}}_2 = \sqrt{1 - \bar{n}} \left( \hat{Q} - i \hat{P} \right),
\]
(21)
where the time is measured in units of \( \omega_0^{-1} \) so that \( \Gamma = \gamma/\omega_0, \Omega = \omega/\omega_0 \). In equations (15), (16), (20) and (21), \( \bar{n} = (e^{\hbar\Omega/ksT} - 1)^{-1} \) is the Bose–Einstein distribution of thermal photons representing the effect of the environment at finite \( T \) and is evaluated at the natural frequency of the oscillator [24].
This dimensionless formulation isolates the scaling factor \( \beta = \sqrt{\frac{S}{\bar{S}}} \) as the single length scale of the problem, where \( S \) is the classical action, such that \( \beta^2 = \frac{\hbar}{m \bar{S} \omega_{\text{osc}}} \). For the classical equation of motion for a damped Duffing oscillator, which is \( \ddot{x} + 2 \Gamma \dot{x} + \beta^2 x^3 - x = \frac{\hbar}{\bar{S}} \cos(\Omega t) \), solutions to the classical equation of motion are unchanged with respect to different values of \( \beta \) except for the length scale of the phase space. However, changing \( \beta \) changes the quantum system considerably.

Comparing the quartic potential equation (8) for the motion of the fundamental mode of the nanomechanical resonator with the potential term in the Duffing Hamiltonian equation (12), it is straightforward to see that for the quantum nanoelectromechanical resonator

\[
\beta^2 = \frac{\hbar}{S} = \frac{h_0}{8\pi^2} \sqrt{\frac{A}{2E\rho l^3}} \left[ (1 + \frac{v_0 l}{\pi^2 E l}) \right]^3, \tag{22}
\]

where, as noted in section 2, we restrict our consideration to resonators of the double-well shape with \( T_0 < -\frac{\pi^2 \hbar}{2E \rho l^2} \). The scaling parameter \( \beta^2 \) determines the way in which the action of the nanomechanical system scales relative to \( \hbar \).

As such, studying the system as \( \beta \) is varied allows us to explore the quantum–classical transition. Previous theoretical work [21–23] for the Duffing problem with a focus on the behavior of the quantum trajectories has confirmed that \( \beta \to 0 \) indeed recovers the classical limiting behavior, including the presence of ‘strange attractors’ of chaos in the phase space, while \( \beta = 1 \) shows behavior unique to the quantum regime. There is interesting physics in the intermediate regime [21], and in particular, the details of the transition as \( \beta \) changes are informative (as below). As shown in table 1, doubly clamped nanoelectromechanical beams can be made of different materials including Si [6, 15], single-walled carbon nanotubes (SWNT) or multi-walled nanotubes (MWNT) [7] and even metals such as gold [25]. The above analysis for the Duffing oscillator also works for a doubly clamped Pt nanowire [26], which has a high fundamental frequency (\( >1 \) GHz). The characteristic parameters given in table 1, for example, show that current experimental setups can be tuned within the range of \( \beta \) considered in this paper. Specifically, for the SWNT experiment of Witkamp et al [11] (see table 1), they used a device of radius \( r = 1.6 \) nm and length \( l_0 = 1.15 \) \( \mu \)m. This yields, using their other device parameters and \( I = \frac{\pi^4 \rho}{16} \), that

\[
\beta^2 = 5.34 \times 10^{-6} \times \left( \frac{\lambda}{4\pi^2} - 1 \right)^{-3/2}, \tag{23}
\]

where \( \lambda = \frac{v_0 l}{\pi^2 E l} \) is greater than \( 4\pi^2 \) to yield the double-well shape. If \( \lambda \) is in the range (39.5–60), we obtain \( \beta \) in the range (0.65–0.04), which as we see below is precisely the range needed for mapping the transition. Witkamp et al report \( \lambda = 26 \), so this is clearly well within the reach of current experiments.

There are other parameters of interest related to the system–environment interactions. In particular, the experimentally measurable quality factor \( Q \) quantifies the energy lost in comparison to the total energy of the resonator during one complete driving period, which can be related to the damping \( \Gamma \). A comprehensive account of the mechanisms of the quality factor, \( Q \), in NEMS has not been established [25], but for the case of doubly clamped beam, \( Q \), can arise from clamping, thermal elastic damping as well as defects of the crystalline structure of the material [8]. Usually, \( Q \) can be measured experimentally using magnetomotive techniques [10]. For low temperatures (20 K and below), \( Q \) can be as large as of the order of \( 10^5 \) [18]. Lowering the temperature further to the millikelvin range depresses energy dissipation further, and recent experiments [25, 27] have achieved quality factors of the order of \( 10^7 \). Although these experiments have pushed toward high \( Q \) and correspondingly low \( \Gamma \), it is clearly possible to make systems more dissipative, thus allowing exploration of a large range of \( \Gamma \).

### 4. Results and discussions

The solutions to equation (10) yield ‘quantum trajectories’, which have been previously discussed in detail in the context of the recovery of classical behavior from quantum mechanics [21–23]. What we focus our attention on below is the ensemble-and-time-averaged asymptotic behavior of the system, defined through the probability distributions

\[
P_{\text{avg}}(x) = \sum_{i=1}^{M} \sum_{k=1}^{N} \frac{1}{MN} |\psi^i(t)|^2 \delta(x - x^i(t)), \tag{24}
\]

with \( t_{\text{transient}} + 4\pi t > t > t_{\text{transient}} \). In doing this, we are averaging over multiple \( M \) trajectories generated in QSD simulations to yield the behavior of density matrices accessible in the laboratory. We also eliminate phase-dependent idiosyncracies of these distributions and correlate better with experimental capabilities regarding the measurement of the dynamics of the quadratures by time-averaging trajectories \( |\psi^i(t)| \) with \( N \) samples of each trajectory over two driving periods, taken after a suitably long transient time \( t_{\text{transient}} \).

The time averaging is not necessary but is useful because (a) in exploring the quantum–classical transition, the properties that signal change most clearly as the parameters are varied are the asymptotic, global characteristics of the system rather than unrepresentative ‘snapshots’ at specific times. Also, (b) the time averaging eliminates experimental difficulties in determining the phase of the driving relative to the observation, in addition to allowing one to quickly build statistics.

We have explored \( P_{\text{avg}}(x) \) at both zero temperature (that is, \( \bar{n} = 0 \)) and finite temperature for several \( \beta \); we

| Type            | Length, \( l_0 \) (nm) | Diameter (nm) | Density \( \rho \) (kg m\(^{-3}\)) | Elastic modulus \( E \) (TPa) |
|-----------------|------------------------|---------------|-------------------------------|-----------------------------|
| Si [6]          | 50–10\(^4\)           | 5 \times 10–10 \times 20 | 2330                          | 0.137                       |
| SWNT [7, 11]    | 50–10\(^6\)           | 1             | 1930                          | 1.25                        |
| MWNT [7]        | 50–10\(^4\)           | 20            | 1930                          | 1.25                        |
| Pt nanowire [26]| 1300                   | 43            | 168                           | 21 090                      |
report here $\beta = 0.01$, 0.3 and 1.0. For each of these values, when we change the coupling strength $\Gamma$, we also looked for signatures of the quantum-classical transition through the changes in $P_{\text{avg}}(x)$. Figure 1 shows $P_{\text{avg}}(x)$ at zero temperature. With high dissipation ($\Gamma = 0.3$) we see what we term nonmonotonicity in the behavior of the probability distribution as a function of $\beta$. Specifically, the probability distribution changes as follows: (a) at $\beta = 0.01$, it is an asymmetric object centered in a single well; (b) at $\beta = 0.3$, it is an essentially symmetric double-peak structure across both wells; and (c) at $\beta = 0.1$, it is a symmetric single-peak straddling the central potential barrier.

We understand these results as follows: for $\beta = 0.01$, the most classical case, the dissipation results in the particle being unable to overcome the central potential barrier even with driving and therefore being confined in a single well. The probability peaks at both ends of the probability distribution correspond to the classical turning points of the oscillator orbit in phase space. That is, the dynamics in this situation are identical to those that would be generated by a classical trajectory, and unsurprisingly, this means that the probability distribution is also entirely explained by classical considerations.

When we increase the quantumness of the system (decrease the scale of the system) to $\beta = 0.3$, the particle acquires a nonzero probability of being in both wells. This is classically impossible given that it is a zero-temperature problem with high damping. The interwell transitions could indeed occur because, as the system gets more quantum mechanical with $\beta$ increasing, the scale of the Lindblad term also effectively increases. It is therefore possible that the particle continues to act essentially classically but the increased effect of the noise leads to a classical noise-activated hopping over the barrier. However, our studies show that the particle is behaving entirely quantum mechanically during the interwell transition, whence this transition is a quantum tunneling effect. Specifically, we have seen that the Wigner function for the particle shows classically forbidden negative regions throughout its dynamics. We have also seen that the energy expectation value remains negative, while the expectation value of the position transits past the origin [21]. Finally, we found the instances in time when a wavepacket has significant probability on either side of the barrier, which is incommensurate with it being a localized wavepacket behaving like a classical particle hopping over the barrier. Incidentally, the last two demonstrate an advantage of using the QSD wavefunction-based approach, since such information is not available in the master equation approach. The symmetric double-peak structure of $P_{\text{avg}}(x)$ at this value of $\beta$ indicates that the ensemble average over the wave-packet behavior leads to an asymptotically equal probability in either well.

For $\beta = 1$, the most quantum case, the probability is peaked at the origin. There are multiple ways of understanding this. One way is that the zero-point energy is such that the lowest states of the potential are above the barrier. As such, the probability is peaked in the middle of the quartic larger well, ignoring the quadratic bump at the bottom. Another way of understanding this situation is to realize that when $\beta$ is increased above 0.3, the scale of the system changes such that the two peaks of the symmetric distribution from $\beta = 0.3$ come closer together until they merge.

These three figures show not only that there is a clear difference between the quantum and the classical probability behavior in the double-well Duffing oscillator compared to the single-well version, but also there are interesting things to be learned from studying the intermediate regime.

As an alternative case, we also present the results for a slightly lower dissipation, with $\Gamma = 0.125$. In this case, even at the classical limit, the dissipation is not strong enough to confine the particle in one well. Specifically, the particle trajectory is that of a strange attractor across both wells [21–23], which yields the unusual shape for the probability distribution. The difference in classical behavior does not persist, and the distributions at higher $\beta$ for this $\Gamma$ value follow the same pattern as those for $\Gamma = 0.3$. As such, comparing the two sets of figures, the results at $\Gamma = 0.3$ show a more dramatic transition from the classical to the quantum regime.

At even lower values of $\Gamma$, the vanishing dissipation means that the wavefunctions do not localize in a basis as happens for greater dissipation [20]. This leads to the standard semiclassical convergence problem for the calculations of $P_{\text{avg}}(x)$. However, we are able to compute $P_{\text{avg}}(x)$ at a finite temperature for each set of $\beta$ and $\Gamma$ and these are presented in figure 2. We see immediately that the results are similar to those at zero temperature in figure 1 although we can see that
temperatures corresponding to \( \approx 4 \) thermal photons begin to eliminate the structure of the distribution(s) at \( \beta = 0.3 \); this is roughly where quantum effects are equal to thermal effects. As one can see, for large dissipation (\( \Gamma = 0.3 \)) and the most classical case, the particle is still confined in one well; that is, thermal activation is not high enough to induce switching between wells. The probability distributions are, of course, broadened because of thermal effects, but the signatures of the transition when \( \beta \) is increased are robust. The last row of figure 2 reassuringly shows the existence of a smooth transition away from zero-temperature behavior.

In conclusion, we have shown that in an open double-well quantum oscillator, clear signatures of quantum behavior and, in particular, the transition away from classical behavior can be found in the changing shapes of an asymptotically obtained probability distribution \( P_{av}(x) \). At the right parameters, this transition from classical to quantum-mechanical behavior can be nonmonotonic and therefore should be clearly visible. Further, these transitions are robust at finite temperatures. The experimentally tunable parameters in NEMS allow for flexibility in exploring the parameter landscape of the transition and are excellent candidates to study this transition. We believe that such experiments are viable in doubly clamped NEMS (see table 1).

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**Figure 2.** \( P_{av}(x) \) plots for various parameters at a finite temperature. For the first three rows, the number of thermal photons \( \bar{n} = 4 \) and \( \beta \) increases from left to right as 0.01, 0.3 and 1.0. \( \Gamma \) increases down the grid as 0.03, 0.125 and 0.3. The last row has \( \Gamma = 0.3 \) and \( \bar{n} = 0.5 \) and shows a smooth transition from the zero-temperature case in figure 1.
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