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Catch bonding in the forced dissociation of a polymer endpoint

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Applying a force to certain supramolecular bonds may initially stabilize them, manifested by a lower dissociation rate. We show that this behavior, known as catch bonding and by now broadly reported in numerous biophysics bonds, is generically expected when either or both the trapping potential and the force applied to the bond possess some degree of nonlinearity. We enumerate possible scenarios and for each identify the possibility and, if applicable, the criterion for catch bonding to occur. The effect is robustly predicted by Kramers theory and Mean First Passage Time theory and confirmed in direct molecular dynamics simulation. Among the catch scenarios, one plays out essentially any time the force on the bond originates in a polymeric object, implying that some degree of catch bond behavior is to be expected in many settings relevant to polymer network mechanics or optical tweezer experiments.

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I. INTRODUCTION

How long does it take a fluctuating particle to escape the trap of a confining potential well? The question is one of the staples of statistical mechanics and, in its simplest incarnation, gives rise to Kramer’s well-known expressions for the rate at which a particle crosses a potential energy barrier—the rate exponentially decaying with increasing barrier height [1]. This escape problem features in a wide range of problems in statistical mechanics and has important applications and consequences in materials science, soft matter, and biological physics for its capacity to predict, under general external conditions, the dissociation rate of bound states and, thereby, the mean lifetime of bonds between, for instance, receptor-ligand pairs [2–4].

Escape kinetics change when forces are taken into consideration. Generally, an applied force aligning with the escape path will hasten dissociation; the naive Kramers prediction is, again, an exponential decrease in lifetime with increasing force [1]. As Ref. [5] showed, forced escape scenarios become considerably richer when the energy landscape is multidimensional—in particular, they note the curious possibility of the lifetime of the bound state initially increasing with the applied force, a phenomenon they termed rollover. This counterintuitive behavior—a bond is strengthened by applying a force to it—is no longer a theoretical curiosity but has, in recent years, been experimentally demonstrated in a range of noncovalent biophysical bonds where it has become widely known as a catch bond [4,6]. For protein-protein bonds, such behavior is generally ascribed to specific conformational properties of the molecules involved [7–11]; we will show that catch bond behavior is generic for protein complexes in which multiple pathways for dissociation exist. In this paper, we will define a catch bond to mean a bound state whose average lifetime possesses an initial regime of increase with force:

$$\frac{d}{df} \langle \tau \rangle(f) \bigg|_{f=0} > 0.$$  

We will show that this behavior is generic when the applied force is nonlinear (i.e., depends asymmetrically on the position of the particle in the trap), even in (but not limited to) one-dimensional energy landscapes, and moreover even in symmetric one-dimensional potentials. The only geometrical requirement for the trap to show catch bond behavior is that there are multiple escape pathways, i.e., two separate escape pathways in a situation where escape occurs along a one-dimensional reaction coordinate, or a continuum of escape paths in higher dimensional dissociation landscapes. Our findings demonstrate the broader generality, without the need for further assumptions, of the implication of entropic elasticity in catch bonding [12].

Our paper is organized as follows: First, we outline the general framework of a forced one-dimensional escape process in the Kramers sense (with rates depending only on barrier height) and summarize the various trapping and forcing scenarios that give rise to catch bonding. Next, we specify the case of a particle that is pulled out of a confining trap by a polymer under tension and study the full differential equation determining the mean first passage time (MFPT) for a general asymmetric potential. Our explicit analytical solution proves that a nonlinear force-extension relation for the polymer is a necessary condition for catch bonding to occur. We conclude our paper with a three-dimensional molecular dynamics (MD) simulation of the escape of a tensed semiflexible polymer with one end trapped in a symmetric, harmonic potential well showing significant (approximately 10% increase in unbinding time) catch bonding, while also highlighting some of the more subtle features of polymer unbinding.

II. KRAMERS APPROACH

An intuitive model to rationalize equilibrium catch bond behavior is the so-called Two-Pathway Model (TPM), developed by Pereverzev et al. [13]. A particle is confined to a local minimum of the trap’s potential energy $U_f(x)$, where $x$ is the position of the particle in the trap. This local minimum $U_f = 0$...
(the bound state) is flanked by two different energy barriers. In any trap configuration, there are two escape paths: one left (L) and one right (R). By applying a force $f$ that is orthogonal to the pathway with the lowest barrier, however, the effective barrier height in the secondary direction is lowered and escape along the second unbinding pathway becomes increasingly likely and frequent. Provided the dissociation lengths of the two pathways are suitably chosen, the same applied force may simultaneously increase the barrier height of the original pathway, making it less likely. The combined escape problem over the two barriers determines the overall escape time, and the differential response to the applied force may result in an increase in the average unbinding time as a force is applied, that is, catch bond behavior. As such, catch bonding in the TPM is a result of an asymmetric trap, in combination with a linear forcing.

We generalize the one-dimensional TPM by making no assumptions on the energy barriers, but assuming only that there are two pathways by which the bond can unbind. We will continue to call those pathways left (L) and right (R), respectively. The bare escape rates corresponding to the pathways are called $k^0_L$ and $k^0_R$ and in the Kramers picture are determined by the barrier heights presented by the trap potential only

$$k^0_i = \nu \exp(-\beta U(b_i)) \quad (i = L, R), \tag{2}$$

with $\nu$ the attempt frequency and $\beta = (k_B T)^{-1}$. For two otherwise independent processes, we thus expect the combined rate of escape to be $k^0 = k^0_L + k^0_R$. These rates define the survival probabilities $S_L(t)$ and $S_R(t)$—the probabilities that unbinding did not occur along the L or R pathway prior to time $t$—as

$$S_L(t) = e^{-k^0_LT}, \quad S_R(t) = e^{-k^0_RT}, \tag{3}$$

and permit us to compute $\pi_{L}(t) dt$ and $\pi_{R}(t) dt$, the likelihood that the first unbinding event occurs along L (R) between $t$ and $t + dt$, as the joint probability of unbinding along L (R) between those times, and having survived unbinding along R (L) up until $t$:

$$\pi_{L}(t) = -\frac{dS_L(t)}{dt}, \quad \pi_{R}(t) = -\frac{dS_R(t)}{dt}. \tag{4}$$

The asymptotic probabilities $P_L$ and $P_R$ of unbinding along either direction are then given by

$$P_L = \int_0^\infty \pi_{L}(t) dt = \frac{k^0_L}{k^0_L + k^0_R}, \tag{5}$$

$$P_R = \int_0^\infty \pi_{R}(t) dt = \frac{k^0_R}{k^0_L + k^0_R}. \tag{6}$$

This makes intuitive sense; if one of the two rates is very high compared to the other, fast unbinding preempts dissociation along the slow pathway. It is, however, also instructive to consider the average unbinding times, separately for each of the two directions. The mean unbinding times along L and R are computed as

$$\langle \tau^0_L \rangle = \int_0^\infty \frac{\pi_{L}(t)}{P_L} dt = \frac{1}{k^0_L + k^0_R}, \tag{7}$$

$$\langle \tau^0_R \rangle = \int_0^\infty \frac{\pi_{R}(t)}{P_R} dt = \frac{1}{k^0_L + k^0_R}. \tag{8}$$

where the factors $P_L$ and $P_R$ are included to normalize the L and R unbinding distributions; $\langle \tau_L \rangle$ and $\langle \tau_R \rangle$ are thus the mean unbinding times of each path, averaged over only the unbinding events in that particular direction. This reveals two interesting properties: the presence of a second unbinding path always decreases the average unbinding lifetime along a primary path (the unbinding is, of course, as least as fast as it was originally), but the average time it takes to escape along either direction becomes the same. Counterintuitively, perhaps, the escape along a very slow direction is greatly sped up by the presence of a secondary, fast unbinding route; this is because only the rare, very quick event along the slow path is not precluded by the abundant, fast rebinding. The overall number of such events may become very small, though, as borne out by $P_L$ and $P_R$. Averaging the escape time over all events, both L and R, gives the expected result mean lifetime $\langle \tau \rangle$ of the bond:

$$\langle \tau^0 \rangle = P_L \langle \tau^0_L \rangle + P_R \langle \tau^0_R \rangle = \frac{1}{k^0_L + k^0_R} = \frac{1}{k^0}. \tag{9}$$

Thus, we find that due to the presence of the other pathway, each path acquires the same escape time. This escape time, moreover, is equal to the overall lifetime of the bound state

$$\langle \tau^0 \rangle = \langle \tau^0_L \rangle = \langle \tau^0_R \rangle. \tag{10}$$

How do external manipulations, such as the application of a force, affect $\langle \tau \rangle$? Let us consider now the case in which the unbinding process takes place in the presence of an additional, external potential $U_{ext}(x, \varepsilon)$ which defines a (possibly position-dependent) force $f_{ext}(x, \varepsilon) = -\partial U_{ext}/\partial x$. The parameter $\varepsilon$ quantifies some manner of tuning this external potential; in the simplest case it is the force itself $[U_{ext}(x, \varepsilon) = -\varepsilon x]$, but it may also represent some more general way of altering the externally applied potential. Assuming that the rate at which the particle crosses the boundaries $b_L$ or $b_R$ of the trap still depends on the barrier height only (we will examine the validity of this assumption in the latter part of this article), $U_{ext}(x, \varepsilon)$ shifts the transition rates according to

$$k_i = k^0_i \exp(-\beta U_{ext}(b_i, \varepsilon)) \quad (i = L, R). \tag{11}$$

To assess the possibility of catch bonding, we compute the change in the overall lifetime $\langle \tau \rangle$:

$$\langle \tau \rangle(\varepsilon) = \langle \tau \rangle(\varepsilon = 0) + \langle \Delta \tau \rangle(\varepsilon), \tag{12}$$

due to a small change in $\varepsilon$. This is done by substituting Eq. (11) into Eq. (9) and expanding to lowest (first) order in $\varepsilon$, to yield

$$\langle \Delta \tau \rangle(\varepsilon) = \beta \varepsilon \left[ \frac{\exp(-\beta U_{ext}(b_L, 0))}{k^0_L} + \frac{\exp(-\beta U_{ext}(b_R, 0))}{k^0_R} \right] k^0 \int_0^\infty \frac{\partial U_{ext}}{\partial \varepsilon}(b_i, \varepsilon) \frac{\partial U_{ext}}{\partial \varepsilon}(b_i, \varepsilon) \right] + O(\varepsilon^2). \tag{13}$$

The essential information in this cumbersome expression may be condensed somewhat. The first factor, between brackets, is strictly positive and does not affect the sign of $\langle \Delta \tau \rangle(\varepsilon)$. For $\langle \Delta \tau \rangle$ to increase with increasing $\varepsilon$, the following "catch
the energy landscapes with corresponding catch bond criteria.

FIG. 1. (a) Illustration of the bond; the bead (red) is trapped in a potential well while an external potential acts on it. (b) Scenarios for the energy landscapes with corresponding catch bond criteria.

calling the dissociation length \(-b_L = b_R = b\), the criterion for increased lifetime reduces to

\[
(iia) \quad k_L^0 \kappa^0 > 1.
\]

This is the TPM catch bonding scenario: if an external force opposing escape along the initially favored (most frequent) pathway is applied, the overall escape rate increases. However, also in case \((iib)\) catch bonding may occur; in this case, the criterion reduces to

\[
(ii) \quad 1 > \frac{b_R}{b_L}.
\]

Thus, even for identical barrier heights applying a pulling force in the direction where the dissociation length is shortest will increase the overall lifetime of the bond. Obviously, suitably chosen combinations of trap asymmetries \((iia)\) and \((iib)\) also produce catch bonding.

C. Symmetric trap, harmonic external potential

For symmetric \(U_1(x)\), and thus again \(-b_L = b_R = b\) and \(k_L^0 = k_R^0\), the left-hand side of Eq. (14) is equal to one. Suppose now that the external potential is supplied by a harmonic spring of rest length \(\ell_0\) and spring constant \(\kappa^{(2)}\), whose left end point is attached to the particle in the trap, and whose right end is at \(x = \ell_0(1 + \epsilon)\). \(\epsilon\) thus measures the extension of the spring and relates to the tension in the spring. In that case, which is scenario (iii) of Fig. 1, the external potential (with \(x = 0\) the center of the trap) is given by

\[
U_{\text{ext}}(x,\epsilon) = \frac{\kappa^{(2)}}{2}(\epsilon\ell_0 - x)^2.
\]

Straightforward substitution into Eq. (14) yields that in this case, too, the catch criterion cannot be met, as both the left- and right-hand sides of Eq. (14) are 1. Moreover, expanding the change in lifetime to second order in \(\epsilon\) reveals that \((\Delta \tau)(\epsilon) \sim -\epsilon^2\), and thus suggests that the lifetime at zero displacement (i.e., zero force applied to the spring) is a maximum.

D. Symmetric trap, anharmonic external potential

We may also consider the same trap conditions as case (iii), but with an anharmonic, nonlinear external potential (we will use a third power here, but the argument is valid with the inclusion of arbitrary odd powers):

\[
U_{\text{ext}}(x,\epsilon) = \frac{\kappa^{(2)}}{2}(\epsilon\ell_0 - x)^2 + \frac{\kappa^{(3)}}{3}(\epsilon\ell_0 - x)^3.
\]

In this case, scenario (iv) of Fig. 1, the catch criterion may be cast as

\[
\left(\frac{\kappa^{(2)}}{2} + \frac{\kappa^{(3)}}{3}\right)e^{\pm ib\alpha} < 1,
\]

demonstrating that for suitably chosen nonlinearity \(\kappa^{(3)}\) catch bonding is possible. In particular, one recognizes the limits \(\kappa^{(3)} \to 0\) [recovering case (iii)] to preclude catch bonding, whereas \(\kappa^{(2)} \to 0\) will always give catch bonding, for all positive values of \(\kappa^{(3)}\).

Summarizing, a range of configurations of external forcing and trap properties give rise to an increase in lifetime with applied force (or with the extension of a springlike tether...
to which the escaping particle is attached). In particular, we show here that such a regime is generic for nonlinearly elastic tethers. Examples of such are all polymers, flexible and semiflexible. Most proteins also display highly nonlinear force-extension relationships, and our result suggests a route towards catch binding that is not due to any specific allosteric or conformational mechanism, but rather is encoded within the universal departures from nonlinearity in protein mechanics. To explore this mechanism in more detail, we focus on cases (iii) and (iv) in the following.

The Kramers analysis assumes $L$ and $R$ transition rates to depend only on barrier height and ignores the shape of the potential along the different unbinding pathways. In order to account for those, and check the robustness of the Kramers predictions, we now compute the lifetimes via the full mean first passage time formalism.

### III. MEAN FIRST PASSAGE TIME

The average time for the particle to escape the trap, and thus for dissociation of the bond, is the mean first passage time (MFPT) for the particle to pass the boundaries of the trap. The MFPT $\langle \tau(x) \rangle$, with $x$ the point of departure inside the trap, obeys the differential equation [14]

$$ f_{tot}(x) \frac{d}{dx} \langle \tau(x) \rangle + \frac{d^2}{dx^2} \langle \tau(x) \rangle = -\frac{1}{D}, $$

supplemented with the boundary condition that $\langle \tau(x) \rangle = 0$ at the boundaries $b_L$ and $b_R$ of the trap. Here is $D$ the diffusion constant for the fluctuating particle, and $f_{tot}(x)$ is the total force acting on the bead, defined as $-\frac{dU}{dx}$ with $U_{tot}(x) = U_f(x) + U_{ext}(x)$.

We will focus on potential catch bonding for symmetric traps, in combination with harmonic or anharmonic external forcing to mimic the situation where the escaping particle is actually one end of a polymeric object or protein. Thus, we choose as our trapping potential

$$ U_f(x) = \begin{cases} \frac{1}{2} \kappa^{(i)} (x^2 - b^2) & |x| \leq b \\ 0 & x > b \end{cases}, $$

i.e., harmonic with trap spring constant $\kappa^{(i)}$ in the region $[-b,b]$, and zero elsewhere (this choice, as well as the discontinuity at the boundaries, is immaterial; the potential outside of the trap region is irrelevant to the escape problem). Escape is defined as first passage across the left or right boundary.

#### A. Symmetric trap, harmonic external potential

When the particle is attached to a harmonic spring with an energy of the form of Eq. (17), resembling scenario (iii) of Fig. 1, the total force on the particle is given by

$$ f_{tot}(x,\epsilon) = -\kappa^{(i)} x + \kappa^{(2)} (\epsilon \ell_0 - x). $$

When we set $\epsilon = 0$, the end point of the spring is fixed a distance $\ell_0$ away from the center of the trap and the total force becomes simply $f_{tot}(x,0) = -\kappa^{(i)} x + \kappa^{(2)} x$. In this case, Eq. (20) with boundary conditions $\langle \tau_{0}(\pm b) \rangle = 0$ may be solved analytically:

$$ \langle \tau_{0}(x_0) \rangle = \frac{b^2}{2D} F_2 \left( \begin{smallmatrix} 1,1; 1/2, 1/2 \end{smallmatrix} ; \frac{b^2 (\epsilon^{(i)} + \kappa^{(2)}/2 \ell_0)}{2D} \right) - \frac{\kappa^{(1)} b^2}{2D} F_2 \left( \begin{smallmatrix} 1,1; 3/2, 3/2 \end{smallmatrix} ; \frac{\kappa^{(2)} b^2 (\epsilon^{(i)} + \kappa^{(2)}/2 \ell_0)}{2D} \right), $$

where $F_2(a;b;z)$ is the generalized hypergeometric function, and $x_0$ is the position of the bead at $t = 0$. ($\tau(x_0)$) is plotted in Fig. 2(a) for various values of $\kappa = \kappa^{(i)} + \kappa^{(2)}$. As expected, the escape time is maximal when the particle departs from the center of the trap; we shall take this value $\langle \tau(0) \rangle$ at $\epsilon = 0$ as our reference time.

To study the effect of pulling the spring under tension, we now increase $\epsilon$ away from zero. This results in an extended tether in the center of the trap, and $f_{tot}$ is now given by Eq. (22). The effect of this small change may be studied perturbatively. When we expand the perturbed solution to second order in $\epsilon$ as

$$ \langle \tau_{\epsilon}(x_0) \rangle = \langle \tau_{0}(x_0) \rangle + \epsilon \langle \tau_{1}(x_0) \rangle + \epsilon^2 \langle \tau_{2}(x_0) \rangle + O(\epsilon^3), $$

with $\langle \tau_{0}(x_0) \rangle$ is the exact solution for $\epsilon = 0$, $\langle \tau_{1}(x_0) \rangle$ and $\langle \tau_{2}(x_0) \rangle$ (and, indeed, the higher order corrections) may be obtained from an order-by-order set of recursive differential equations,

$$ -x_0 (\kappa^{(i)} + \kappa^{(2)}) \frac{d}{dx_0} \langle \tau_{\epsilon} \rangle + \frac{\kappa^{(2)} \ell_0}{D} \frac{d}{dx_0} \langle \tau_{\epsilon-1} \rangle + \frac{d^2}{dx_0^2} \langle \tau_{\epsilon} \rangle = 0, $$

FIG. 2. (a) The average unbinding time $\langle \tau(x_0) \rangle$ obtained by Eq. (23) for a cross-link which released at $x_0$ at $t = 0$. The sum of $\kappa_i$ and $\kappa_s$ is 2 (black line), 4 (blue line), and 6 (red line). (b) The first three terms of the perturbed system for $\kappa = 2$. 042405-5
with boundary conditions \(\langle \tau_0(\pm b) \rangle = 0\). These equations, too, may be analytically solved for \(\langle \tau_1(x_0) \rangle\) and \(\langle \tau_2(x_0) \rangle\). Figure 2(b) graphs these first two corrections to the \(\varepsilon = 0\) result and confirms what was already suggested by the Kramers analysis: \(\langle \tau_1(0) \rangle = 0\), meaning that there is no effect, to first order in \(\varepsilon\), on the lifetime of the bond (i.e., no catch bond).

Moreover, we also see that indeed the second order correction \(\langle \tau_2(0) \rangle = 0\) is negative, proving that the zero-\(\varepsilon\) lifetime is indeed a maximum. Attaching the escaping particle to a linear spring cannot increase the lifetime in a symmetric potential.

B. Symmetric trap, anharmonic external potential

The Kramers theory predicts catch bonding in the case of an anharmonic external potential. To connect this to a more realistic setting, we will specify to the case of a Worm-Like Chain tether; instead of a Hookean spring the escaping particle is now connected to a semiflexible polymer with a contour length \(\ell_c\) and a persistence length \(\ell_p\), what resembles scenario (iv) of Fig. 1. The force-extension relation for such a polymer is strongly nonlinear and given by [15,16]

\[
\ell(f) = \ell_c - \frac{k_B T}{2f} \left[ \frac{f \ell_c^2}{k_B T \ell_p} \coth \left( \frac{f \ell_c^2}{k_B T \ell_p} \right) - 1 \right].
\]

The equilibrium length for such a polymer is \(\ell_0 = \ell_c^2/\ell_p\). Expanding \(\ell(f) - \ell_0\) to second order in the force \(f\) and inverting the relation yields the force-extension relation to first anharmonic order for the semiflexible WLC:

\[
f_{\text{WLC}}(x, \varepsilon) = 90k_B T \left( \frac{\ell_p}{\ell_c^2} \right)^2 (\varepsilon \ell_0 - x) + \frac{5400}{7} k_B T \left( \frac{\ell_p}{\ell_c^2} \right)^3 (\varepsilon \ell_0 - x)^2.
\]

So, with the identification

\[
\kappa^{(2)} = 90k_B T \left( \frac{\ell_p}{\ell_c^2} \right)^2,
\]

\[
\kappa^{(3)} = \frac{5400}{7} k_B T \left( \frac{\ell_p}{\ell_c^2} \right)^3,
\]

we may use the external potential of Eq. (18) to capture the lowest relevant nonlinear order of tethering by a semiflexible WLC. Obviously, for larger extensions (and thus forces) we may need to go to higher orders in \((\varepsilon \ell_0 - x)\).

To verify the catch bonding effect predicted by Kramers for this anharmonic force extension relation on the mean unbinding time, we numerically solve Eq. (20) for a semiflexible polymer with \(\ell_c = \ell_p = 20\), confined to a trap with a spring constant of \(\kappa^{(2)} = 10\) and a dissociation length \(b = 1\). While we can no longer treat this case analytically, we do have access to the full range of \(\varepsilon\) allowing us to compute also the total extent of the catch effect. Comparing the full WLC force-extension relation with its harmonic and first anharmonic approximations [potentials graphed in Fig. 3(a)] confirms the Kramers predictions; to harmonic order, there is no catch bond effect, but the inclusion of the first anharmonic term creates a regime of increasing lifetime with rising \(\varepsilon\). Thus, indeed, a nonlinear anharmonicity [third order terms or higher, odd powers, in \(U_{\text{ext}}(x, \varepsilon)\)] is a prerequisite for this type of catch bond effect. The full WLC curve shows that the rise in lifetime continues over an extended range of \(\varepsilon\), and that the total induced lifetime increase can attain values of over 10% [see Fig. 3(b)].

So both the Kramers analysis and the MFPT computations show a catch bonding effect for a particle in a symmetric, one-dimensional trap with multiple escape pathways, attached to a polymer or, in fact, any tether possessing some anharmonic response. In the final part of this paper, we assess the real-life validity of the effect by a molecular dynamics (MD) simulation in three dimensions.

IV. MOLECULAR DYNAMICS

We use LAMMPS MD [17] to perform a Langevin dynamics simulation, implicitly taking into account the solvent, of a fluctuating semiflexible bead-spring chain consisting of \(N\) beads with diameter \(d\), connected by identical Hookean springs with a spring constant \(K_{sp}\) and rest length equal to \(d\) and including a bending contribution to the chain energy, quantified by a bend stiffness \(K_b\). The resulting chain energy is

\[
U_{\text{MD}} = \sum_{i=1}^{N} K_{sp}(\ell_i - d)^2 + \sum_{i=1}^{N-1} K_{sp} \theta_i^2,
\]

where \(\ell_i\) the length of spring \(i\), and \(\theta_i\) the angle between spring \(i\) and spring \(i + 1\). One of the end beads is trapped in a harmonic potential with spherical symmetry, the other end is fixed at a distance \(\ell = \ell_0(1 + \varepsilon)\) from the center of the trap. In the simulation a polymer consisting of \(N = 40\) springs that
that even in symmetric three-dimensional trapping potentials, tether anharmonicity suffices to generate a catch bond effect. There is, however, no satisfactory quantitative agreement between the numerical solution of the Eq. (20) and the MD simulations, despite the identical parameters. We believe this to be due to the anomalous diffusion of the polymer end point within the trap. As we show in Fig. 4(b), the mean-squared displacement \( \langle \delta x^2 \rangle \) of the end bead scales as \( \langle \delta x^2 \rangle \sim t^{3/4} \), that is, slower than the expected power of 1. This subdiffusive motion of a polymer end point is well known [18] but invalidates the derivation that produces Eq. (20). To fully capture the escape problem of a polymer end point, we suggest that a version of Eq. (20) with fractional derivatives might be required, but this is beyond the scope of our current presentation.

V. SUMMARY AND CONCLUSION

Our results prove, that catch bonding—bound states whose lifetime increases with applied force—is a generic feature of bonds where either the trap itself, or the external structures that a ligand is connected to, or both, display some degree of nonlinear response. These effects are robustly predicted by Kramers theory and MFPT theory and are confirmed in direct MD simulation.

The mechanism we describe suggests that catch bonding may be more prevalent than previously assumed, because it does not require finely tuned structural or conformational properties of bond participants—the minimal requirements being that (1) multiple escape paths exist, in one or more dimensions (a criterion not universally met in protein-protein binding) and that (2) the tether is nonlinearly elastic (a criterion readily met in proteins). The same effects are predicted to occur in reversible (noncovalent) bonding in, for instance, polymeric materials; these reversible links should, under very general conditions, also show a regime of some strengthening when loaded. It will be instructive, in future analysis, to assess the theoretical limits of this effect to inspire the design of novel, responsive catch bond materials.