Theory of rapid force spectroscopy

Jakob T. Bullerjahn¹, Sebastian Sturm¹ & Klaus Kroy¹

In dynamic force spectroscopy, single (bio-)molecular bonds are actively broken to assess their range and strength. At low loading rates, the experimentally measured statistical distributions of rupture forces can be analysed using Kramers’ theory of spontaneous unbinding. The essentially deterministic unbinding events induced by the extreme forces employed to speed up full-scale molecular simulations have been interpreted in mechanical terms, instead. Here we start from a rigorous probabilistic model of bond dynamics to develop a unified systematic theory that provides exact closed-form expressions for the rupture force distributions and mean unbinding forces, for slow and fast loading protocols. Comparing them with Brownian dynamics simulations, we find them to work well also at intermediate pulling forces. This renders them an ideal companion to Bayesian methods of data analysis, yielding an accurate tool for analysing and comparing force spectroscopy data from a wide range of experiments and simulations.

¹Universität Leipzig, Institut für theoretische Physik, 04103 Leipzig, Germany. Correspondence and requests for materials should be addressed to K.K. (email: klaus.kroy@uni-leipzig.de).
In the realm of soft matter, structural stability often hinges on delicate intermolecular bonds that are easily broken apart through external forces. The resulting maleability is characteristic of biological matter on practically all scales from single proteins over mesoscale cellular scaffold structures and individual cells to whole tissues. Thanks to the development of a variety of nanomanipulation methods, intermolecular interactions can nowadays be investigated on a single-molecule level using a number of techniques commonly referred to as ‘dynamic force spectroscopy’7–12, allowing the experimentalist to isolate single binding sites and probe their strength by quickly and reliably inducing hundreds or thousands of unbinding events. The wealth of stochastic unbinding trajectories thus obtained is routinely analysed through a schematic but effective description of molecular bonds in terms of the attraction range \( x_0 \) and the activation energy \( \delta \) of the binding potential. To extract quantitatively useful predictions from this simple model, any theory of unbinding kinetics needs to provide a reasonable approximation to the underlying molecular dynamics. At the relatively low loading rates that were conventionally realized in experiments, any transient effects arising from the finite relaxation time of the bond itself can be neglected. This has allowed for the development of a range of analytical theories of forced bond breaking that greatly facilitate the analysis and interpretation of dynamic force spectroscopy data13–18. In contrast, detailed molecular dynamics simulations2 usually operate in the opposite limit of very high loading rates, where bond breaking becomes essentially deterministic19. Recently, also single-molecule assays are increasingly resolving the hitherto elusive rapid dynamics of bond breaking and accompanying macromolecular conformational changes, for example, in protein unfolding upon rapid loading20, or taut DNA recoil21 and supercoiling22. This provides a strong incentive to also push the existing theories to higher loading rates. Moreover, in spite of the experimental progress and improved computational abilities23, matching the loading rates used in experimental and simulation studies remains challenging20, making the development of a unified analytical theory of dynamic force spectroscopy, covering both fast and slow loading rates, all the more desirable.

To this end, we derive in the following a probabilistic theory of dynamic force spectroscopy in the spirit of Bell24, Evans & Ritchie25 and Dudko, Hummer & Szabo14 that becomes exact at high external forces (or, equivalently, high loading rates) and reduces to established results at low loading rates. We provide explicit, closed-form analytical results for the most common experimental loading protocols. They agree with exact numerical simulations of the microscopic bond model for all loading rates, save for a narrow region at the crossover from diffusion-dominated to deterministic dynamics. This makes them an ideal companion to Bayesian methods25 and a natural choice for the analysis of spectroscopy experiments and simulations alike. Moreover, we show that these results constitute the lowest-order approximation to a rigorous mathematical formulation of escape kinetics, which opens the way of their systematic extension to higher precision.

Results

Theory. Molecular binding and unbinding transitions lie at the heart of every chemical reaction and have thus been thoroughly investigated long before the advent of single-molecule manipulation techniques. In 1940, Kramers laid down a comprehensive, analytically tractable theory of chemical reaction rates26 that has since then become synonymous with reaction rate theory and can still be considered state of the art for most applications. The basic idea is that a molecular bond corresponds to a local free energy minimum and thus remains stable under weak perturbations. Its thermal fluctuations can be represented, within an effective picture, by those of a Brownian particle trapped in some one-dimensional binding potential \( U(x) \), with the ‘reaction coordinate’ \( x \) typically corresponding to the distance between two binding partners. The function \( U(x) \) should have a stable minimum at \( x = 0 \), surrounded by a region of attraction extending to some finite coordinate \( x_0 > 0 \) beyond which \( U(x) \) either vanishes or turns repulsive. As soon as the particle has left the basin of attraction, \( x(t) > x_0 \), the bond can be considered broken. Excluding the possibility of quantum-mechanical tunnelling, which is negligible on a macromolecular scale, the dissociation process is a tug of war between thermal and potential forces. The thermal forces drive the bond coordinate \( x \) diffusively out of the bound state and the deterministic force \( F(x) = -\frac{\partial U}{\partial x} \) drags it back to the origin. On a probabilistic level, the dissociation rate of an ensemble of initially bound particles can then be deduced from an associated Fokker–Planck equation27, a partial differential equation for the time-dependent distribution of particles that is mathematically exact but cannot be solved in closed form without the help of further simplifying assumptions. In most cases of practical interest, the activation energy \( \delta \) is large compared with the thermal energy scale, \( \delta \gg k_BT \). Dissociation then becomes a rare event, leaving sufficient time for any transients caused by the fast microscopic bond dynamics to die out. In the steady-state situation that ensues, an accurate analytical solution of the Fokker–Planck equation is possible26, yielding a time-independent unbinding rate \( k \) that scales exponentially in the activation energy, \( k \propto \exp\left( -\frac{\delta}{k_BT} \right) \).

This provides a natural starting point for the analysis of force-induced bond rupture. An external pulling force \( F \) acting along the reaction coordinate \( x \) ‘tilts’ the internal binding potential \( U \), which lowers the effective free energy barrier and increases the unbinding rate. Within the Bell model24, the reaction distance \( x_0 \) is assumed to remain fixed under the external force, such that the force-dependent unbinding rate immediately follows as \( k(F) = k(0)\exp(\frac{F x_0}{k_BT}) \). However, reaction rates are not measured directly in single-molecule experiments, where actual unbinding times may follow a broad statistical distribution. Also, external forces are usually not constant in time. In a typical experiment, molecular bonds are pulled apart using retracting actuators such as AFM cantilevers that exhibit Hookean stretching behaviour and therefore exert forces growing linearly with their retraction distance. Assuming an external force \( F(t) \) that steadily increases in time, Evans & Ritchie13 derived from the force-dependent unbinding rate \( k(F) \) a general expression for the experimentally measured distribution \( p(F) \) of rupture forces,

\[
p(F) = \frac{k(F)}{F} \exp\left( -\int_0^F \frac{k(f)}{F} df \right).
\]  

Here, the force dependence of the loading rate \( F \) is optional, but can be used to take into account nonlinear loading protocols or nonlinearly elastic force transducers such as polymer linkers28,29. For a power-law binding potential and a cusp-shaped binding potential of finite range, Evans & Ritchie were furthermore able to improve upon the phenomenological Bell model by accounting for force-induced shifts in the reaction distance \( x_0 \). Focusing on a linearly increasing external force, Dudko, Hummer & Szabo later derived analytic results for cusp-shaped19 and linear-cubic30 binding potentials. These could finally be condensed into a uniform expression for the rupture force distribution and its first two moments that contained a new fit parameter to smoothly interpolate between different potential shapes14,15. A recent crop of theories further fleshed out the free energy landscape by taking into account the force fluctuations.
pertaining to stiff actuators\textsuperscript{16–18} (that is, actuators with a spring constant $\kappa \gtrsim \delta/x_0^3$). These theories nowadays provide the standard tools used to convert rupture force histograms obtained from force spectroscopy experiments into estimates of the underlying binding potentials.

From a dynamical point of view, these approaches still rely on a strong separation between the short timescale of internal bond dynamics and the long timescale of bond dissociation. Although this approximation is well justified in the original Kramers theory\textsuperscript{28} of a force-free bond, it is bound to break down as soon as the external loading becomes rapid enough to effectively flatten out the energy barrier before rupture occurs. In that case, unbinding becomes essentially determinist as the bond is swept away ballistically by the external force. As shown by Hummer & Szabo\textsuperscript{19} for a harmonic cusp potential, one can accurately predict the mean rupture force at high loading rates by neglecting stochastic fluctuations altogether and afterwards obtain a global approximation to the mean rupture force by manually interpolating between the high-force result and a conventional theory that accounts for the low-rate regime.

In the following, we derive a probabilistic theory of forced unbinding that includes fluctuations even at high pulling speeds and becomes exact both in the limits of high and low loading rates. Already the leading order results of this theory surpass previous work; moreover, it provides a systematic route for further improvement. We start with the same cusp-shaped binding potential as Hummer & Szabo,

$$U(x) = \begin{cases} \delta'(x/x_0^3), & x < x_0 \\ -\infty, & \text{otherwise} \end{cases} 
$$

(2)

Formally, the potential becomes negatively infinite at $x = x_0$, corresponding to an absorbing boundary condition (thus ignoring rebinding events, which quickly become negligible under external load\textsuperscript{3}). Although it can be argued\textsuperscript{14} that a linear-cubic potential constitutes a more faithful approximation to many real binding potentials, we will show in the following that the practical difference between both models is often insignificant; what we lose in generality is more than made up for by what we gain in terms of analytical tractability.

To set the stage, we first summarize some basic definitions. The time-dependent probability $W(x,t)dx$ to find any particle in the ensemble between $x$ and $x+dx$ follows from the Fokker–Planck equation\textsuperscript{27}

$$\partial_t W(x,t) = \left[ D \delta'' - \partial_x A(x,t) \right] W(x,t), \quad W(x_0,t) \equiv 0, 
$$

(3a)

$$A(x,t) \equiv -\frac{D}{k_B T} \partial_x V(x,t) + 2 \frac{\delta}{x_0^3} x. 
$$

(3b)

$$A(x,t) \equiv -\frac{D}{k_B T} \partial_x V(x,t) + 2 \frac{\delta}{x_0^3} x. 
$$

(3c)

Here, $D$ denotes the diffusion coefficient and $V(x,t)$ an external pulling force potential that may either represent some prescribed force protocol $F(t) \equiv -\partial_x V(x,t)$ or a moving external spring, if $V(x,t) \equiv \kappa [x - \gamma(t)]^2/2$. The spatially integrated distribution

$$S(t) = \int_{-\infty}^{x_0} W(x,t)dx 
$$

is the proportion of bound particles or ‘survival function’ at any given time $t$. A constant survival function $S(t) = \text{const}$ corresponds to zero escape events and thus a vanishing unbinding rate $k = 0$. Likewise, a positive escape rate $k > 0$ amounts to an exponentially decaying survival function $S(t) \equiv S(0) \text{exp}(-kt)$. In the general case, $S(t)$ may be neither constant nor exponential, but an arbitrarily complex function of time. However, it is always monotonously decaying and nonnegative and thus defines a (potentially time-dependent) generalized unbinding rate $k(t)$,

$$k(t) = -\frac{S(t)}{S(t)/t}. 
$$

(5)

In principle, we could now solve equation (3a) (or an equivalent integral formulation\textsuperscript{31,32}) numerically\textsuperscript{33} to obtain the time-dependent probability distribution $W(x,t)$, calculate the survival probability $S(t)$ via equation (4) and feed the result into equation (5) to precisely compute the unbinding rate $k(t)$ for a given set of model parameters ($\delta$, $x_0$, $D$). This procedure is, however, too computationally intensive for the inverse problem of obtaining $\delta$, $x_0$, and $D$ from experimental data. To arrive at an analytical approximation to the unbinding rate that is quickly evaluated on a computer, we write down a first approximation to $W$ by ignoring the absorbing boundary,

$$\partial_t W_G(x,t) = \left[ D \delta'' - \partial_x A(x,t) \right] W_G(x,t), 
$$

(6a)

$$W_G(x \to \pm \infty, t) = 0. 
$$

(6b)

With the external driving potential $V(x,t)$ in equation (3c), which is at most parabolic, $W_G$ is strictly Gaussian for any localized initial distribution $W_G(x,t = 0) \equiv \delta(x-x_0)$. It is easily evaluated analytically,

$$W_G(x,t|x_0) = \exp\left(-\frac{\delta^2 [x - \delta(x - x_0)]^2}{k_B T x_0^3 [1 - C^2(t)]/\gamma^3} \right) \sqrt{\frac{\pi k_B Tx_0^3}{1 - C^2(t)}}/\gamma^3, 
$$

(7)

where $\delta(t)$ follows from a purely deterministic equation of motion,

$$\ddot{x}(t) = -\frac{k_B T}{x_0^3} \dot{x}(t) + 2 \frac{\delta}{x_0^3} x - \partial_x V(x,t) |_{x=x}, 
$$

(8)

$C(t)$ denotes the positional autocorrelation function,

$$C(t) = e^{-2\gamma(t)k_BT/x_0^3}, 
$$

(9)

and $\gamma$ the relative change in the system’s overall spring constant introduced by the external actuator; that is, $\gamma = 1$ for external fields or soft springs ($k \ll \delta/x_0^3$), $\gamma = 1 + \kappa x_0^3/2\delta$ otherwise.

Going back to the time-dependent escape rate equation (5), we note that the computation of $S(t)$ via equation (4) would be needlessly complicated. Since escape events can only take place at the absorbing boundary $x_0$, the outflow of bound particles into the unbound state is equivalent to the probability flux $j(x_0,t) \equiv -\dot{S}(t)$ across $x_0$. It only depends on local properties of the probability distribution $W$, via the continuity equation

$$j(x_0,t) = A(x_0,t) W(x_0,t) - D \delta'' W(x_0,t) |_{x=x_0}. 
$$

(10)

Replacing $W$ by $W_G$ in equation (10) defines the corresponding approximate flux $j_G$, survival function $S_G = 1 - \int_{x_0}^{x} j_G(x,t)dx'$, and escape rate $k_G(t) = j_G(x_0,t)/S_G(t)$.

Note that the Gaussian expressions become asymptotically exact for pulling forces greater than the critical force needed to flatten out the energy barrier, $F >> F_c = 2\delta/x_0$ (corresponding to high loading rates $F >> DF_c/x_0^3$ in a constant-rate experiment), due to an essentially ballistic decay of the bound state. Intuitively, the probability density $W_G$ does not have time to ‘sense’ (and react to) the presence of the absorbing boundary before it is pulled across $x_0$ by the deterministic drift term $A(x,t)/W_G(x,t)$. For the escape rate, we thus find the exact asymptotic limit

$$k(t) \rightarrow k_G(t) = j_G(x_0,t)/S_G(t). 
$$

(11)

The situation is slightly more complicated for slow loading, where deterministic driving and stochastic dynamics interfere. Consider first the limiting case of a freely diffusing particle, that is, if the binding potential, the external forcing and the absorbing boundary condition are neglected. There is a purely diffusive net flux $j_{\text{diff}} = -D W'_G(x_0,t)$ at the position $x_0$, half of which is
randomly backscattered (see Fig. 1). It is suggestive that inserting a perfect absorber at $x = x_b$ would suppress the backscattering and therefore double the diffusive flux across $x_b$. Adding the deterministic drift back in, we get

$$j(x_b, t) \overset{F = 0}{\sim} j^*(x_b, t)$$  \hspace{1cm} (12a)

$$j^*(x_b, t) \equiv A(x_b, t)W_G(x_b, t) - 2D\partial_x W_G(x, t) \big|_{x=x_b}.$$  \hspace{1cm} (12b)

This expression for the flux can indeed be understood as the first-order approximation to an exact theory, as discussed in the Methods section. There we also show that $j^*(x_b, t)$ provides an asymptotically exact expression for the escape rate at low loading rates,

$$k(t) \overset{F = 0}{\sim} j^*(x_b, t)/1.$$  \hspace{1cm} (13)

We now observe that the additional factor of two in front of the diffusive component in $j^*$ compared with $j_0$ is irrelevant for the limit of large pulling rates, which is dominated by the deterministic drift term $A(x_b, t)W_G(x_b, t)$, suggesting that equation (11) still holds with $j_0$ replaced by $j^*$. Moreover, note that for high energy barriers $\delta \gg k_bT$ and low pulling rates $F \ll DF_b/x_b^2$, the Gaussian survival probability $S_G(t)$ remains close to 1 since $W_G$ extends only negligibly beyond $x_b$ for all relevant times $t$, that is,

$$1 - S_G(t) = \int_{x_b}^\infty W_G(x, t)dx = e^{-\frac{\delta - F x_b t}{k_bT}} \ll 1.$$  \hspace{1cm} (14)

Our two asymptotic results can therefore be combined into the unified approximation

$$k(t) \approx \frac{j^*(x_b, t)}{S_G(t)},$$  \hspace{1cm} (15)

which is exact both in the limits of low and high loading rates. Together with equation (1), this yields the rupture force distribution

$$p(F) \approx \frac{1}{F^2} \frac{d}{dt} \exp \left( - \int_0^t \frac{j^*(x_b, t')}{S_G(t')} dt' \right).$$  \hspace{1cm} (16)

Evaluating $p(F)$ in its general form equation (16) still requires some computational effort, since it depends in a potentially complicated way on experimental details such as the driving protocol, the initial distribution of particles within the bound state and the stiffness of the force actuator. However, for the common case of a linearly increasing pulling force $F = F_0 t$ (exerted by an external field or a soft spring with stiffness $k \ll \delta/x_b^2$) we obtain the closed analytical result

$$p(F) \approx \frac{k_0}{F} \left( 1 + \frac{F}{F_c} - \frac{2\bar{x}(F)}{x_b} \right) e^{\frac{\delta - (1 - \bar{x}(F)/x_b)}{k_bT}} \times \exp \left( - \frac{k_b T k_0}{F x_b} \left[ e^{\frac{\delta - (1 - F/F_c)}{k_bT}} - 1 \right] \right).$$  \hspace{1cm} (17)

where $k_0$ is the associated force-free Kramers rate for the cusp potential,

$$k_0 = \frac{2D\delta}{k_b T x_b^2} \left( \frac{\delta}{\pi k_b T} \right)^{k_bT},$$  \hspace{1cm} (18)

and $\bar{x}(F)$ is the force-dependent coordinate of the maximum of $W_G(x, t)$,

$$\bar{x}(F) = x_b + \frac{F}{F_c} - \frac{k_b T F x_b}{DF_c} \left( 1 - e^{-2DF_b/k_bT F_c^2} \right).$$  \hspace{1cm} (19)

In the limit of small loading rates we obtain $\bar{x}(F) \sim F x_b/F_c$ and equation (17) reduces to the expression derived by Dudko, Hummer & Szabo (DHS) in ref. 14 for $v = 1/2$. In the Methods section, we provide a compilation of analogous results for stiff force transducers, arbitrary driving protocols or arbitrary initial conditions $W(x, t = 0)$ (Table 1), as well as a closed expression for the mean rupture force $\langle F \rangle$.

Some remarks about the comparison of data and theory may be useful, here. First, note that, in any case, a comparison of data and theory for the distribution $p(F)$ is substantially more informative than merely fitting $\langle F \rangle$. Yet, every analytical approximation to the true rupture force distribution depends on at least three different parameters (binding energy $\delta$, attraction range $x_0$ and diffusivity $D$). Direct fits of experimentally obtained rupture force histograms are therefore prone to get trapped in some local optimum in parameter space, thus often missing the best possible solution. One way to obtain more reliable results is to first aggregate several data sets obtained under different loading rates and subsequently perform a ‘global’ fit using a single set of parameters only. Conventionally, this is done by either performing a least-squares fit of several histograms at once or by fitting the mean rupture force as a function of loading rate $^{13-15,19}$.

Both approaches can be improved upon systematically using a maximum-likelihood approach$^{25}$ that, instead of optimizing for some arbitrary measure of fit quality (such as the squared residual), employs Bayesian analysis to select those model parameters most likely to underlie the given rupture force histograms. Compared with conventional fitting methods, the maximum-likelihood approach straightforwardly extends to heterogeneous data sets encompassing different linker stiffnesses or loading protocols. Moreover, it has proven significantly more robust with respect to small ensemble sizes$^{25}$, a potentially crucial advantage in the analysis of real-world data sets.

**Figure 1 | Flux into an absorber at $x_b$.** (a) Given some statistical distribution $W(x, t)$ of particles, these particles can be driven across the position $x_b$ either ballistically by external forces ($\dot{j}_{bal}$) or diffusively through random thermal noise ($\dot{j}_{diff}$). (b) Inserting an absorber at $x_b$ effectively eliminates diffusive backscattering, thus doubling the diffusive contribution to the resultant probability flux $\dot{j}^*$ into $x_b$. 

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experiments. In general, the larger the range of applicability of a given theory of force spectroscopy, the better the results obtained using the maximum-likelihood approach. This fact should work to our advantage, as our model covers a much wider range of loading rates than the predominant Bell-Evans 'standard model' and its recent extensions by DHS and others. In the remainder and in the Supplementary Note 1, we illustrate the data fitting procedure with some examples and provide some practical tools and protocols for optimized data analysis.

**Application to simulation data.** Although our theory becomes exact in the limits of high and low external loading rates, the quality of our approximation at finite loading rates is *a priori* unclear. To assess the practical applicability of our method, we have thus generated synthetic rupture force distributions using Brownian Dynamics simulations of the underlying microscopic model (see Methods). As Fig. 2 exemplifies, our theory is as good as the approach by Hummer & Szabo in capturing the mean rupture force \( \langle F \rangle \) as a function of loading rate. Moreover, whereas the Hummer–Szabo model builds on an athermal treatment of the underlying microscopic equation of motion, we fully account for thermal fluctuations. These can be significant even within the 'ballistic' regime \( \dot{F} > D_F/x_b^2 \), as Fig. 3 and the experimental data by Rico et al. show. Hence, our approach allows us to perform a complete and systematic Bayesian analysis (see Methods and ref. 25) of rupture force histograms (as opposed to mean rupture forces), thus making full use of the available experimental data. As apparent from Fig. 3, our approximation still breaks down at intermediate loading rates, because in our formalism, we do not actually absorb particles as they cross the absorbing boundary at \( x_b \) but merely count them. After leaving the potential well, they build up a 'phantom population' that may later produce an unphysical backflow into the bound state. In principle, this deficiency could be rectified, at least for the cusp potential, by extending the systematic integral equation approach outlined in the Methods to higher orders. Yet, the range of applicability of equation (16) already covers the vast majority of unbinding events. By simply truncating \( p(F) \) after its first zero crossing, we obtain an analytical expression for the rupture force distribution that works well at all loading rates, except for a narrow range close to a critical value \( \dot{F}_c = D_F/x_b^2 \).

Although we find the derivation of an analytical, global approximation to the rupture force distribution \( p(F) \) satisfying in itself, its practical utility lies first and foremost in the accurate and efficient analysis of experimental unbinding data. To estimate how well our model stacks up against the prevailing methods of data analysis, we have thus analysed numerically obtained rupture force histograms, see Supplementary Fig. 1. We considered both the cusp-shaped potential and a linear-cubic binding potential (Supplementary Figs 2 and 3), using local and global (maximum-likelihood) fits both of equation (16) and of the rupture force distributions derived by DHS and Maitra & Arya (as well as a 'cusp-optimized' counterpart to their results, detailed in Supplementary Note 2) for soft and stiff linkers, respectively. Furthermore, we emulated conventional experimental procedures by fitting the mean rupture force \( \langle F \rangle \), using both the analytical expressions derived by Friddle, associated to the DHS-model, and Maitra & Arya, as well as the numerical interpolation formula derived by Hummer & Szabo. In the Supplementary Note 1, the interested reader will find a comprehensive discussion of all these approaches, the gist of which is as follows. At high loading rates, unsurprisingly, our theory proves superior to existing probabilistic theories. More importantly, however, the combination of the maximum-likelihood analysis and our globally valid approximation is powerful enough to compete with the best conventional models even for a linear-cubic binding potential while requiring neither prior knowledge of the potential shape nor additional fit parameters. The only other global model to date, the numerical interpolation formula obtained by Hummer & Szabo, yields similarly good results as our own approach if supplied with sufficient data, although the increased computational cost of numerical integration leads to longer fitting times. Using a smaller range of loading rates (that is, 3–6 decades in \( \dot{F} \) instead of 12), the higher information content of our full-histogram description can produce significantly better results. Moreover, as our approach inherits the greater generality and robustness of the Bayesian maximum-likelihood analysis, this gap is bound to widen with small ensemble sizes or heterogeneous data sets. Since the maximum-likelihood method is somewhat more difficult to implement in practice than traditional methods, we wrote (and provide for free use) a ready-made Mathematica notebook that covers some of the most common force spectroscopy setups, see Supplementary Data 1.

**Discussion**

Our theory provides an analytically tractable generic model of dynamic force spectroscopy that reproduces previously known, accurate results at low loading rates and improves upon the results derived by Hummer & Szabo for high loading rates by including stochastic fluctuations. It provides excellent results also at intermediate loading rates, thus constituting an ideal companion to Bayesian methods of data analysis. Apart from serving as a systematic and convenient replacement for current theories of force spectroscopy, our approach may prove crucial to...
the analysis of future high-speed force spectroscopy setups and full-scale MD simulations. Finally, we note that at low enough loading rates, the effect of a polymeric force transducer onto the resultant rupture force distributions can be approximated by choosing \( F(t) \) and \( \tilde{F}(t) \) accordingly\(^{20,22,23}\). At higher loading rates, the dynamics of force propagation within the polymer\(^{21,34,35}\) may become relevant, which could be an interesting subject for future theoretical developments based on our theory.

**Methods**

**Alternative derivation of equation (12b).** The (non-Gaussian) probability distribution function \( W(x,t) \) is, in the presence of an absorbing boundary at \( x = x_b \), in general not analytically tractable. However, it is related to its Gaussian counterpart \( W_G(x,t) \) in a simple manner\(^{22,33}\):

\[
W(x,t) = W_G(x,t) - \int_0^t W_G(x,t|x',\tau)(j(x',\tau)\text{d}\tau),
\]

(20a)

where \( j(x,\tau) \) is the associated probability flux, satisfying the continuity equation

\[
\frac{\partial W(x,t)}{\partial t} = -\frac{\partial j(x,t)}{\partial x} + \sigma(x,t),
\]

(21a)

with the boundary conditions \( W(x,t)|_{x=x_b-\rightarrow} = W(x_b,t) = 0 \). The Gaussian distribution \( W_G(x,t) \) and flux \( j_G(x,t) \) also satisfy the continuity equation (21a) for \( W_G(x,t)|_{x=x_b-\rightarrow} = 0 \). Alternatively\(^{36}\), the absorbing boundary condition in equation (20b) can be interpreted as a sink term \( \sigma(x,t) < 0 \) ∀\( t \), resulting in the modified equation

\[
\frac{\partial W(x,t)}{\partial t} = -\frac{\partial j(x,t)}{\partial x} + \sigma(x,t),
\]

(21b)

where \( W(x,t)|_{x=x_b-\rightarrow} = 0 \). Using the fact that the flux vanishes at infinity, \( j(x \rightarrow -\infty,\tau = 0) \), we integrate equation (21a) to obtain

\[
\frac{\partial}{\partial t} \int_{-\infty}^{x_b} W(y,t)\text{d}y = -j(x_b,t),
\]

(22)

or equivalently, \( \dot{S}(t) = -j(x_b,t) \).

Taking the time derivative of equation (20a) gives

\[
\frac{\partial W(x,t)}{\partial t} \mid_{x=x_b-\rightarrow} = \frac{d}{dt} \int_0^t W_G(x,t|x',\tau)(j(x',\tau)\text{d}\tau)
\]

\[
\equiv -\frac{\partial j_G(x,t)}{\partial x} + \int_0^t \frac{\partial j_G(x,t|x',\tau)}{\partial x}(j(x',\tau)\text{d}\tau)
\]

\[
- W_G(x,t|x_b)(j(x_b,\tau)\text{d}\tau).
\]

(23)

which can be compared with equation (21a), leading to

\[
\frac{\partial j(x,t)}{\partial x} = \frac{\partial j_G(x,t)}{\partial x} - \int_0^t \frac{\partial j_G(x,t|x',\tau)}{\partial x}(j(x',\tau)\text{d}\tau) + \delta(x - x_b)j(x_b,\tau). \]

(24)

Inserting this expression into equation (21b) and then integrating over the interval \((-\infty, x_b) \) results in

\[
j(x_b,t) = \frac{1}{2} \frac{\partial j_G(x_b,t)}{\partial x} - \int_0^\infty \frac{\partial j_G(x_b,t|x',\tau)}{\partial x}(j(x',\tau)\text{d}\tau)
\]

\[
+ \frac{1}{2} j(x_b,t) - \int_{-\infty}^{x_b} \sigma(y,t)\text{d}y.
\]

(25)

Equation (27) was originally derived by Buonocore et al. in ref. 37, where it was noted that the function \( \sigma(x,t) \) can be uniquely determined using

\[
\lim_{t \rightarrow -\infty} \psi(x_b,t|x',\tau) = 0.
\]

(28)

a criterion that follows from the fact that for all \( t \),

\[
W_C(x,t|x_b) = \delta(x-x_b).
\]

(29)

Solving equation (28), we obtain \( g(t) = -A(x_b,t)/2 \) (ref. 38) and thus

\[
\psi(x_b,t) = \frac{1}{2} A(x_b,t) W_C(x,t) - D \frac{\partial W_C(x,t)}{\partial x} |_{x=x_b}.
\]

(30)

Formally, equation (27) can be solved iteratively,

\[
j(x_b,t) = 2\psi(x_b,t) - 4 \int_0^t \psi(x_b,t|x',\tau_1)\psi(x_b,\tau_1)\text{d}\tau_1 + \ldots \]

\[
- ( -2)^{n+1} \int_0^t \psi(x_b,t|x',\tau_1) \int_0^{\tau_1} \psi(x_b,\tau_2|x_2,\tau_2) \text{d}\tau_2 \ldots \text{d}\tau_n,
\]

(31)

where \( 0 < \tau_n < \tau_{n-1} < \cdots < \tau_1 < \tau \) and \( j(x_b,\tau) \) vanishes as \( n \rightarrow \infty \). Truncating the above result after the first term yields a first-order approximation \( j(x_b,t) \) to the flux that coincides with equation (12b),

\[
j(x_b,t) \equiv j(x_b,t) \equiv 2\psi(x_b,t) = A(x_b,t) W_C(x,t) - 2D \frac{\partial W_C(x,t)}{\partial x} |_{x=x_b} = j(x_b,t).
\]

(32)

**Asymptotic exactness at low loading rates.** Our escape rate \( k(F) \) and rupture force distribution \( \rho(F) \) both reduce to the results of DHS (equations 3 and 4 in ref. 14 for \( v = 1/2 \)), in the limit \( \dot{F} \rightarrow 0 \). The flux \( j(x_b,F) \) in equation (12b) is a sum

\[
\sum_{n=0}^{\infty} \int_0^\infty \psi(x_b,t|x',\tau_1) \psi(x_b,\tau_1) \text{d}\tau_1 + \cdots + \int_0^\infty \psi(x_b,\tau_n|x_n,\tau_n) \text{d}\tau_n \ldots \text{d}\tau_1,
\]

which reduces to \( 2\psi(x_b,t) = A(x_b,t) W_C(x,t) - 2D \frac{\partial W_C(x,t)}{\partial x} |_{x=x_b} \), as \( \dot{F} \rightarrow 0 \).
Figure 4 | Measuring rupture forces with a stiff transducer. (a) As long as the bond remains intact, the combined bond-transducer system can be seen as two harmonic springs connected in series. At low pulling speeds, positional fluctuations within the bond state translate into force fluctuations that can be smoothed out via a time-moving average (see inset). (b) As long as the effective free energy barrier is still large compared with $k_BT$, the probability distribution $W(x, t)$ closely approximates a Gaussian centered within the bond state and $\langle x \rangle_{\text{slow}}(t)$ virtually coincides with $\langle x \rangle_{\text{slow}}(t)$. (c) At high pulling forces, the static force-balance argument fails, as it yields an equilibrium position $\langle x \rangle_{\text{slow}}(t)$ beyond $x_b$. The improved approximation $\langle x \rangle_{\text{slow}}(t)$ instead is always bounded above by $x_b$.

of the Gaussian flux $j_G(x_b, F) = \langle A(x, F) - D\partial_x W(x, F) \rangle_{x=x_b}$ and the diffusive flux $j_{\text{diff}}(x_b, F) = -D\partial_x W(x, F) |_{x=x_b}$. Due to their asymptotic behaviour,

$$j_G(x_b, F) \sim \frac{c \langle \mathcal{E} \rangle_{x_b} / DF_b \mathcal{E}_b}{\mathcal{E}(1 - F/F_b)}, \quad \mathcal{E} < DF_b,$$

and computed from $p'(F)$ the mean rupture force via

$$\langle F \rangle = \int_0^\infty F \, p'(F) \, dF.$$  \hspace{1cm} (36)

Alternatively, we can (for constant $F$ and $\mu = 1$) provide a rough analytical estimate that is exact in the limits of small and large loading rates, respectively,

$$\langle F \rangle = \int_0^\infty F \, p(F) \, dF \approx \langle F \rangle |_{\text{slow}} + \langle F \rangle |_{\text{fast}}.$$  \hspace{1cm} (37)

For large loading rates the mean rupture force evaluates to

$$\langle F \rangle |_{\text{fast}} \approx \int_0^\infty \frac{\partial_x \mathcal{E}_b}{2DF_b} e^{-y^2/2} \left( \left[ \frac{1}{2} + \frac{1}{2F} \right] I_{1/4} \left( \frac{\sqrt{F} \mathcal{E}_b}{2} \right) + \frac{1}{2} I_{1/4} \left( \frac{\sqrt{F} \mathcal{E}_b}{2} \right) \right) \, dF \approx \frac{\sqrt{\mathcal{E}_b}}{DF_b} + e \left( \left( \frac{\mathcal{E}_b}{2} \right)^{-1} \right),$$

where $I_{1/4}(z)$ is the modified Bessel function of the first kind. This result coincides with the deterministic mean rupture force originally derived in ref. 19.

For small loading rates $F < F_b$, we arrive at a result previously obtained from the Bell–Evans model by Gergely et al.\(^{19}\),

$$\langle F \rangle |_{\text{slow}} \approx \frac{e^{\mathcal{E}_b} \mathcal{E}_b}{\mathcal{E}_b}.$$  \hspace{1cm} (38b)

Here, $q = \exp(\mathcal{E}_b (1 - x)/\mathcal{E}_b)$, $x = \chi^1 k_\text{eff}/k_b x_b$, and $E_c(z)$ denotes the exponential integral,

$$E_c(z) = \int_0^z \frac{e^{-t}}{t} \, dt.$$  \hspace{1cm} (39)

Inserting equation (38a) and (38b) into equation (37), we thus end up with

$$\langle F \rangle \approx \frac{e^{\mathcal{E}_b} \mathcal{E}_b}{\mathcal{E}_b} + \frac{\sqrt{\mathcal{E}_b}}{DF_b} + e \left( \left( \frac{\mathcal{E}_b}{2} \right)^{-1} \right),$$

It should be noted that we provide $\langle F \rangle$ only for the sake of completeness. In fact, it is wasteful to discard experimentally measured force fluctuations instead of directly fitting the full histograms $p(F)$ and we thus advise against the use of equation (40) (as much as any other available predictions for $\langle F \rangle$) for analysing the experimental data.

**Simulations.** To generate rupture force histograms across a large range of external field strengths $F$, we have used a stochastic Euler scheme to directly integrate the equivalent Langevin equation for the Fokker–Planck equation (3a),

$$\gamma \dot{z}(t) = -U'(z) - D, \gamma \dot{z}(x, t) + \zeta(t),$$  \hspace{1cm} (41)

where $\zeta(t)$ denotes Gaussian, white noise,

$$\langle \zeta(t) \zeta(t') \rangle = 0, \quad \langle \zeta(t) \rangle = 2k_B T \gamma \delta(t - t'^{\prime})$$  \hspace{1cm} (42)

and $\gamma$ is the bond friction coefficient, $\gamma = k_b T / D$. External force is applied either through an external field $V(x, t) = -x F$ (corresponding to a deterministic force) or through a moving external spring centred at $y(t) = y_0$. Initial particle positions are drawn from a Boltzmann distribution; once the particle position $x(t)$ crosses $x_b$, we consider the bond broken and record the corresponding rupture force. Although one might argue that prior knowledge of the barrier position $x_b$ allowed us an unrealistic advantage over actual experiments, we note that even beyond the critical pulling force, strong variations in the intramolecular binding potential $U$ around $x_b$ manifest themselves in experimentally detectable force signatures, see Supplementary Note 3 and the accompanying Supplementary Fig. 4.

**Maximum-likelihood method.** Following ref. 25, the likelihood $P$ of a given set of parameters $\delta, x_b, D$ is computed as follows,

$$p(F) = \prod_{i=1}^M p(F_i | (\delta, x_b, D), \hat{F}),$$

where $p(F)$ denotes the totality of rupture forces taken into consideration

$$p(F_i | (\delta, x_b, D), \hat{F})$$

and the rupture force distribution $p$ given by equation (16), evaluated at the measured rupture forces $F_i = \hat{F}_i$ with model parameters $(\delta, x_b, D)$ and $\hat{F}$. The logarithm of the right-hand side is then maximized using the 'NMaximize' method provided by Wolfram Mathematica. Since the maximum-likelihood method is somewhat more involved than conventional fitting procedures, we provide a ready-made Mathematica notebook that allows the user to analyse her data through a simple graphical interface (Supplementary Data 1).

**Nonconstant loading rates and arbitrary initial conditions.** In deriving equation (17), we have started from a localized initial condition $W(x, t=0) = \delta(x - x_b)$, calculating the corresponding distribution of first passage times and in
Table 1 | Analytical approximations to the rupture force distribution \( p(F) \).

| General definitions |
|---------------------|
| \( W_0(x, F|x') = \frac{\exp \left( -\frac{(x' - x - \nu y)^2}{2k_B T} \right)}{\sqrt{2\pi k_B T}} \), \( \frac{W_0(x, F|x')^2}{\nu} \geq x_k \exp \left( -\frac{(x' - x - \nu y)^2}{2k_B T} \right) \) |
| \( C(F) = \exp \left( -2d_0^2 \frac{F}{k_B T} \right) \), \( \bar{x}(F) = \frac{1}{F} \left[ F(0) - C(F) - \frac{1}{F} \int_0^F \frac{C(F')}{F'} dF' \right] \) |
| \( F_c = \frac{2\nu}{x_k} \), \( M(F) = 1 + \frac{2}{\nu} \left[ \frac{1}{F} - F_c \right] \), \( F' = \{ F(t), \ F(0) \} \) |

| External field setup |
|----------------------|
| \( V(x, t) = -F(t)x, \chi = 1 \) |
| \( p(F) \approx -\frac{F(0)}{V_F} \exp \left( -\int_{F(0)}^F \frac{W_0(y|\nu \bar{x}(F) + 2d_0)/W_0(x|\nu \bar{x}(F)) dy}{F(0)} \right) \) |
| \( \int_{F(0)}^F \frac{W_0(y|\nu \bar{x}(F) + 2d_0)/W_0(x|\nu \bar{x}(F)) dy}{F(0)} \) arbitrary initial conditions \( W_0(x) \) |
| \( p(F) \approx -\frac{F(0)}{V_F} \exp \left( -\int_{F(0)}^F \frac{A(x, \nu \bar{x}(F)) + 2d_0/\nu \bar{x}(F) M(F)}{F(0) M(F)} \right) \) |
| \( \int_{F(0)}^F \frac{A(x, \nu \bar{x}(F)) + 2d_0/\nu \bar{x}(F) M(F)}{F(0) M(F)} \) equilibrium initial conditions \( W_0(x) \propto \exp(-\mu x U(x)/k_B T) \) |

| Linear force ramp \( F(t) = \hat{F} t^{\mu=1} \) equation (17) |
| \( p(F) \approx \frac{1}{F} \left[ \frac{\left( \frac{x_k}{k_B T} \right)^{\nu}}{C(x_k/k_B T)} \right] \frac{W_0(x_k)}{x_k} \) |
| \( C(F) = \exp \left( -2d_0^2 \frac{F}{k_B T} \right) \), \( \bar{x}(F) = x_k \left[ \frac{F}{k_B T} - \frac{k_B T}{F} \right] \) |
| \( \int_{F(0)}^F \frac{A(x, \nu \bar{x}(F)) + 2d_0/\nu \bar{x}(F) M(F)}{F(0) M(F)} \) equilibrium initial conditions \( W_0(x) \propto \exp(-\mu x U(x)/k_B T) \) |

We provide here closed formulas for various classical experimental situations, as well as generic expressions for nonlinear loading protocols and arbitrary initial conditions. All scenarios assume a monotonously increasing (or decreasing) external load, which allows us to unambiguously convert rupture forces into the time domain via the functional inverse \( F(t) \) of the time-dependent rupture force \( F(t) \) (that is, for a linearly increasing force ramp \( F(t) = \hat{F} t \) follows as \( F(t) = \hat{F} t \)). Notice that equation (17) is a special case of the bottom entry of this table for \( \mu = 1 \).

Stiff force transducers. For stiff transducers, that is, external spring constants \( k \gg \frac{d_0}{x_k} \), the relation between rupture time and rupture force depends to a certain degree on the experimental time resolution. Instead of a deterministic time-dependent pulling force \( F(t) \), one measures then a force within the transducer that fluctuates with the bond coordinate \( x(t) \). These fast fluctuations are typically smoothed out using a time-moving average, which in a conventional low-speed experiment corresponds to an equilibrium average, with respect to the time-dependent combined potential \( U(x) + V(x, t) \) (molecular bond + transducer), yielding thus as the measured rupture force \( F = \kappa [y(t) - x(t)] \).

As long as the effective free energy barrier is still large compared with \( k_B T \) (which can safely be assumed in the limit of low loading rates \( \hat{F} \ll d_0 x_k^2 \)), the actual distribution of bond particles is essentially a Gaussian centred within the bound state. Hence, the anharmonicity of the molecular binding potential beyond \( x_k \) can be neglected and the equilibrium position \( \langle x \rangle \) of bound particles follows from a simple force-balance argument\(^{17} \) (see Fig. 4a).

\[
\kappa_{\text{mol}} \langle x \rangle_{\text{down}}(t) = \frac{\kappa_{\text{pol}}(t)}{\kappa_{\text{torn}} + \kappa_b} \langle y \rangle_{\text{down}}(t),
\]

\[
\langle x \rangle_{\text{down}}(t) = \frac{\langle y \rangle_{\text{down}}(t)}{\kappa_{\text{mol}}(t)},
\]

where \( \kappa_{\text{mol}} = \frac{U''(x)}{2} = \frac{d_0}{x_k^2} \) for the cusp potential. Setting \( \chi = 1 + \kappa x_k^2 / 2d_0 \),

the end averaged over all possible starting points \( x_0 < x_k \) using an equilibrium Boltzmann distribution. This choice of initial condition should be appropriate to most real-world scenarios, but since our theory readily extends to nonequilibrium initial conditions (corresponding, for example, to a particle held fixed by an auxiliary trapping potential that is turned off at \( t = 0 \)), Table 1 provides a generic rupture force distribution for arbitrary choices of \( W(x, t = 0) \equiv W_0(x) \). Importantly, for a thermally stable, smoothly varying binding potential \( U \) the initial distribution is generally Gaussian around the energy minimum at \( x = 0 \),

\[
W_0(x, 0) \approx \frac{1}{2k_B T} \frac{\exp \left( -\frac{U''(0) + V''(0, t = 0)}{2k_B T} x_k^2 \right)}{\sqrt{2\pi k_B T}},
\]

but its width may deviate from the value obtained from our cusp-shaped model potential. To better account for other types of binding potentials (such as the often used linear-cubic potential), we can thus treat

\[
\frac{U''(0) + V''(0, t = 0)}{2} = \frac{U''(0) + V''(0, t = 0)}{2d_0^2 / x_k^2 + V''(0, t = 0)} \equiv \mu
\]

as an additional fit parameter, in analogy to the parameter \( \nu \) used by Dudko, Hummer & Szabo\(^{14} \) (although in contrast to \( \nu \), our fit parameter \( \mu \) is relevant only at high loading rates \( \hat{F} \gg d_0 / x_k^2 \) since at low loading rates the memory of the initial condition fades away long before rupture occurs). In Table 1, we also provide a generic expression for arbitrary force protocols \( F(t) \) that do not necessarily scale linearly in time.
this simplifies to
\[ F_{\text{low}}(t) = \kappa \langle y(t) - x(t, \text{slow}) \rangle = \frac{\kappa y(t)}{1 + \kappa / k_{\text{mol}}} \equiv \kappa y(t) \quad (49) \]

Large external forces, on the other hand, may shift \( \langle x(t, \text{slow}) \rangle \) beyond \( x_0 \) and render the static force-balance argument obviously inadequate. We obtain a reasonable generalization of equation (48b) by instead truncating the time-dependent approximate probability distribution \( W_c(x, t|x_0) \) at \( x = x_0 \) and using that to compute \( \langle x(t) \rangle \),
\[ \langle x(t) \rangle \approx \langle x(t) \rangle_c(t) \]
\[ \langle x(t) \rangle_c(t) = \frac{\int_{-\infty}^{x_0} W_c(x, t|x_0) dx}{\int_{-\infty}^{\infty} W_c(x, t|x_0) dx} \]
\[ = \frac{x(t) - \langle x(t) \rangle}{\text{erf} \left( \frac{\langle x(t) \rangle - x_0}{\sqrt{2/W(t)}} \right) + 1} \quad (50) \]

with mean \( \langle x(t) \rangle \), variance \( \sigma^2(t) \) and the autocorrelation function \( C(t) \) defined as usual (note that the particle position \( x_0 \) at \( t = 0 \) is irrelevant in the low-force limit and is eliminated in the high-force limit by averaging over the initial distribution of particle positions \( W(x_0, t = 0) \)).
\[ \tilde{x}(t) = -\kappa x_0 t^2 \int_0^t C(t') \langle y(t') \rangle dt' \quad (51a) \]
\[ \sigma^2(t) = \kappa x_0^2 \left( 1 - C^2(t) \right) / 2 \quad (51b) \]
\[ C(t) = \exp \left[ -2 \frac{dy(t)}{\kappa x_0^2} \right] \quad (51c) \]

In the limit of low pulling forces, equation (50) reduces to the established result equation (48b); \( \langle x(t) \rangle_c(t) \approx \langle x(t) \rangle \). In the high-force limit, however, an equilibrium average is no longer warranted as a force spectroscopy setup with sufficient time resolution should instead measure the transducer force at the very moment of rupture, \( F_{\text{rupture}}(t) = \kappa y(t) - x_0 \).
\[ F_{\text{rupture}}(t) = \frac{\kappa y(t)}{1 + \kappa / k_{\text{mol}}} \quad (52) \]

This equation coincides with the operational definition of ‘rupture force’ used by Hummer & Szabo \(^{19}\). Since in the limit of high loading rates, \( x_0 \) is generally negligible compared with the transducer position \( y(t) \) at the moment of rupture and since \( \langle x(t) \rangle_c(t) \) is bound above by \( x_0 \), we can use \( \kappa y(t) - \langle x(t) \rangle_c(t) \) as a global approximation to the force rupture; the disadvantage of this approach is that \( F(t) \) becomes difficult to solve for. As intermediate pulling speeds, it may be reasonable to perform the functional inversion numerically; in practice, however, we propose to simply replace \( F(t) \) by its respective asymptotic limits,
\[ F(t) \sim \begin{cases} \kappa y(t) / \chi, & \kappa y(t) \ll \Delta E / k_B T \chi^2 \\ \kappa y(t), & \text{otherwise} \end{cases} \quad (53) \]

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Author contributions
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