

Substrate interactions and promiscuity in a viral DNA packaging motor

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The ASCE (additional strand, conserved E) superfamily of proteins consists of structurally similar ATPases associated with diverse cellular activities involving metabolism and transport of proteins and nucleic acids in all forms of life¹. A subset of these enzymes consists of multimeric ringed pumps responsible for DNA transport in processes including genome packaging in adenoviruses, herpesviruses, poxviruses and tailed bacteriophages². Although their mechanism of mechanochemical conversion is beginning to be understood³, little is known about how these motors engage their nucleic acid substrates. Questions remain as to whether the motors contact a single DNA element, such as a phosphate or a base, or whether contacts are distributed over several parts of the DNA. Furthermore, the role of these contacts in the mechanochemical cycle is unknown. Here we use the genome packaging motor of the *Bacillus subtilis* bacteriophage ϕ 29 (ref. 4) to address these questions. The full mechanochemical cycle of the motor, in which the ATPase is a pentameric-ring⁵ of gene product 16 (gp16), involves two phases—an ATP-loading dwell followed by a translocation burst of four 2.5-base-pair (bp) steps⁶ triggered by hydrolysis product release⁷. By challenging the motor with a variety of modified DNA substrates, we show that during the dwell phase important contacts are made with adjacent phosphates every 10-bp on the 5′–3′ strand in the direction of packaging. As well as providing stable, long-lived contacts, these phosphate interactions also regulate the chemical cycle. In contrast, during the burst phase, we find that DNA translocation is driven against large forces by extensive contacts, some of which are not specific to the chemical moieties of DNA. Such promiscuous, nonspecific contacts may reflect common translocase–substrate interactions for both the nucleic acid and protein translocases of the ASCE superfamily¹.

To test the role of the phosphate backbone charge in motor–DNA interactions we inserted a 10-bp region of double-stranded methylphosphonate DNA (dsMeP) into the middle of an ~8-kilobase-pair (kb) native DNA molecule (Fig. 1a), and followed the packaging of this molecule by a single ϕ 29 prohead–motor complex using optical tweezers. In MeP the charged oxygen on DNA is replaced with an uncharged isosteric methyl group while conserving the B-form structure of DNA^{8,9} (Fig. 1b, inset). Thus, it is possible to determine the role of this chemical modification in a native geometric context. Figure 1b shows sample packaging traces under 5 pN of constant tension and saturating [ATP] (1 mM). Packaging proceeds normally until the motor encounters the inserted modification where it pauses, and then either successfully traverses the insert or completely

dissociates. In stark contrast to related helicases, in which disruption of a single charge interaction^{10–14} completely abolishes translocation, the packaging motor traverses 10 bp of neutral DNA with a probability of ~80% under a tension of 5 pN (Fig. 1c).

To rule out the possibility that the motor crosses the neutral insert by diffusive fluctuations as opposed to making direct contact with uncharged moieties, we took advantage of the strong force dependence of diffusive traversal times¹⁵. We found that there is only a twofold increase in pause duration with a 15 pN increase in force (Fig. 1c)—much less than the 10⁵-fold increase predicted for diffusion across a 10-bp distance (Supplementary Discussion). Furthermore, lowering [ATP] increases the pause duration and decreases the traversal probability, providing further support for an active, ATP-dependent crossing mechanism. Thus, the motor actively traverses the insert by making contacts with elements other than the phosphate charge, albeit with reduced efficiency, indicating that native packaging involves both charge and non-charge contacts.

To determine whether phosphate charges from both strands are equally important, we created hybrid inserts in which only one strand contains the MeP backbone. We used a 30-bp insert to accentuate the effect of the uncharged section because the traversal probability of a 30-bp dsMeP insert at 5 pN is ~4%. At this force the traversal probability of the hybrid insert with MeP on the strand packaged from 3′–5′ is almost 90%, whereas that of the hybrid insert with MeP in the 5′–3′ strand is reduced to 10% (Fig. 1d). This result clearly indicates that the most important phosphate interactions are made with the 5′–3′ strand in the direction of packaging. Such preferential interaction with a single DNA strand has been shown for the monomeric dsDNA translocase EcoR124I (ref. 16), although it is more surprising in this case of a ring-ATPase in which several subunits of the ring are simultaneously in close proximity to both strands.

Next, we addressed whether a critical length is involved in the interaction of the motor and its DNA substrate. Figure 1e shows that as we increase the length of the double-stranded neutral insert from 5 to 10 bp there is no statistically significant change in traversal probability, but the 1-bp increase from 10 to 11 bp results in a twofold reduction. Further increasing the length to 15 bp does not produce a similar change. The location of this discrete change in traversal probability at first seems to be inconsistent with the 10 bp of DNA packaged by the motor each full mechanochemical cycle⁶, but these results are easily reconciled if the motor makes contact with two adjacent phosphates, with either contact being sufficient for packaging (Fig. 1f). The co-crystal structure of the related BPV helicase E1 with its single-stranded DNA (ssDNA) substrate reveals simultaneous

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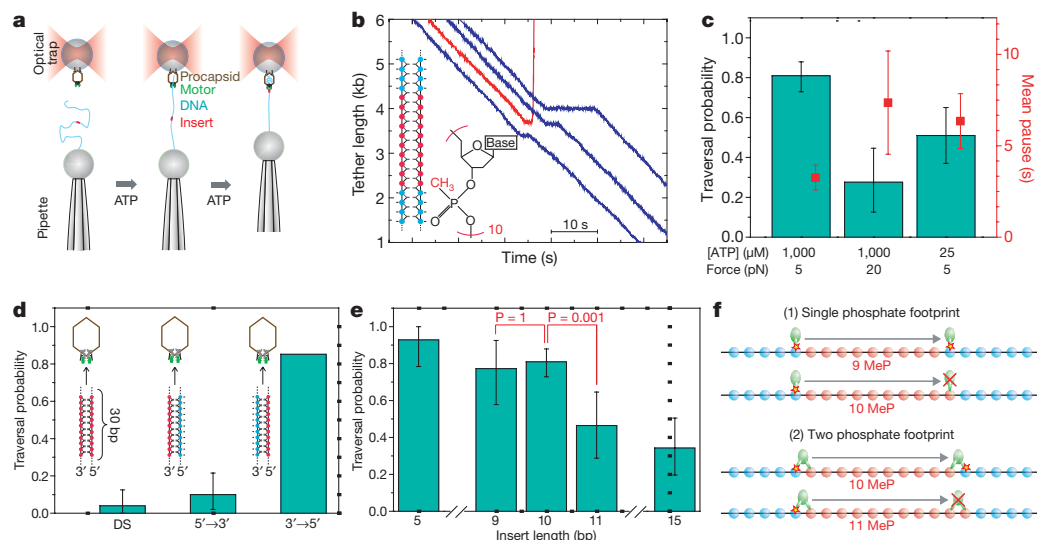


Figure 1 | Packaging of neutral DNA analogues. **a**, A prohead–motor complex bound to a microsphere is held in an optical trap while a micropipette holds a second microsphere bound to DNA containing a modified insert. A tether is formed and packaging is initiated when the beads are briefly brought into close proximity in the presence of ATP. **b**, Representative packaging traces of DNA containing 10-bp dsMeP at a constant load of 5 pN. Blue traces show traversal after a pause, and the red trace shows a terminal dissociation event after a pause. Inset is a schematic of the insert, with MeP nucleotides in red and unmodified nucleotides in blue, and the chemical structure of a MeP nucleotide. **c**, Force and ATP dependence of traversal probability and pause duration of 10-bp dsMeP inserts. **d**, Traversal probability of 30-bp dsMeP and DNA–MeP hybrid

inserts at 5 pN. **e**, Traversal probability of dsMeP inserts at 5 pN force as a function of insert length. P values (two-tailed Fisher exact test) between 9 and 10 bp, and between 10 and 11 bp, are indicated. **f**, Translocation cycle length and footprint size limits from MeP length dependence. This scheme shows the position of a subunit that contacts the DNA before and after a full mechanochemical cycle, that is, 10 bp. Contact with a single phosphate would produce a drop in traversal probability between 9- and 10-bp dsMeP, whereas contact with two phosphates would produce the observed drop between 10- and 11-bp dsMeP. In **c–e** the traversal probability is plotted using the Laplace estimator²⁶, with 95% confidence intervals from the adjusted Wald method²⁷, and error bars of pause durations denote the s.e.m.

contact with adjacent phosphate charges¹⁷, lending support to this interpretation.

We next investigated the specific role of these phosphates in the mechanochemical cycle by probing the base-pair-scale dynamics of the motor at an uncharged insert. Phosphate–motor interactions may have two possible roles in the mechanochemical cycle: they may provide the long-lived contacts that are required to keep the enzyme attached to the DNA, or they may have a sensory role, accelerating a chemical rate, such as ATP hydrolysis, upon detecting that the DNA is bound and properly oriented. These two roles of the phosphate charge, although not mutually exclusive, can be revealed by characteristic dynamics of the motor as it traverses the modified insert. If the phosphate provides load-bearing contacts, its absence will increase the dissociation rate of the motor, and the insert-induced pause will consist of a series of attempts to package followed by small slips. Alternatively, if the role of the phosphate is sensory, we expect the time between packaging steps to be lengthened, owing to the decreased rate of catalytic turnover.

To determine the dynamics of the motor as it crosses a neutral insert (10-bp dsMeP), we followed packaging using dual-trap optical tweezers with higher spatial and temporal resolution⁶. The pauses observed at low resolution are actually remarkably dynamic events, containing two types of sub-pauses that occur at distinct locations along the modified DNA insert (Fig. 2a). The first type of sub-pause, which we term ‘upstream pauses’ because it occurs at longer DNA tether lengths, is followed by either brief disengagement of the motor (slips) or packaging attempts. These attempts are themselves followed by a second class of sub-pause, which we term ‘downstream pauses’. After slips from either the upstream or the downstream pauses, the motor typically recovers and repackages the DNA to the position of the upstream pauses. Occasionally the motor does not recover from a slip, resulting in a terminal slip. The branching probabilities of these events are shown in Fig. 2b.

The upstream sub-pauses occur in a uniform position on a given tether, ± 1 bp (s.d.), and have longer average durations, 1.00 ± 0.08 s

(s.e.m.; Fig. 2c, e), whereas the downstream sub-pauses occur at the end of attempts of different sizes, ± 3 bp (s.d.), and have shorter durations, 80 ± 10 ms (s.e.m., Fig. 2d, f). The upstream- and downstream-pause time distributions are both well-described by single exponential

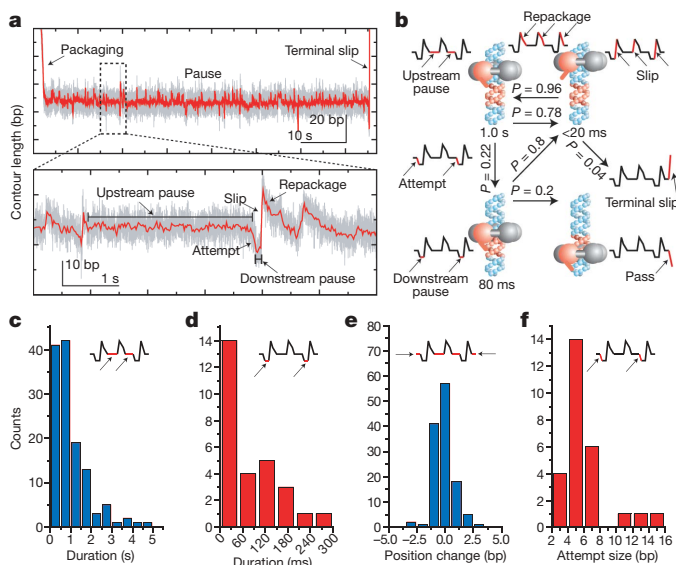


Figure 2 | High-resolution dynamics at a neutral-DNA insert. **a**, Base-pair scale dynamics at 10 bp of dsMeP consist of two classes of sub-pauses, upstream and downstream pauses, separated by attempts and punctuated by slips and repackaging events. **b**, A cartoon model of the dynamics of these pauses with average lifetimes and inter-conversion probabilities. The full statistics of these states are in Supplementary Table 3. **c**, **d**, Histograms of upstream (**c**) and downstream (**d**) pause durations. The distributions have n_{\min} values—the ratio of the mean squared to the variance—of 1.1 ± 0.1 (s.d.) and 1.3 ± 0.4 (s.d.), respectively, and are thus well-described by single exponential decays. **e**, Histogram of position changes in the upstream pause. **f**, Histogram of the distance between upstream and downstream pauses.

decays (Fig. 2c, d), suggesting that each class of sub-pause corresponds to an individual kinetic process, as opposed to a weighted sum of different kinetic processes, and that each of them is governed by a single rate-limiting kinetic event. In combination with their distinct mean durations, these observations suggest that each class of sub-pause corresponds to one of the mechanochemical phases of the motor: ATP-loading dwell or stepping burst.

To identify the kinetic state of the motor associated with each class of sub-pause, we measured the dynamics of motors traversing 5 bp of neutral DNA, an insert shorter than the 10-bp span of the full mechanochemical cycle⁶. Whereas contacts during the burst phase occur every 2.5 bp, contacts during the dwell phase should only occur every 10 bp. Thus, all motors must make burst phase contacts with the neutral DNA, while a fraction of the motors will never make dwell phase contacts with the neutral DNA. If the upstream pause occurs during the dwell phase, then only a fraction of the motors will show such a pause. Only ~50% of the motors packaging 5 bp of dsMeP display a long pause (Supplementary Fig. 1), strongly suggesting that the upstream pause corresponds to the dwell phase, and hence the downstream pause to the burst phase. This conclusion is further supported by the average duration of the downstream pauses and the exponential nature of their distribution, which are both similar to those of the micro-dwells observed during the burst phase on charged DNA under similar conditions⁶. Finally, the highly uniform positions of the upstream pauses in a given tether indicate that this single-base-pair position is unique; suggesting that after slipping and repositioning, the motor resides at the boundary of the charged and neutral DNA during its dwell phase.

The multiple slip/attempt phenotype observed shows that removing the phosphate charge reduces the processivity of the motor (Fig. 2a). Because the motor slips from both upstream and downstream pauses, load-bearing phosphate contacts are made during both the dwell and the burst phases of the cycle. Notably, however, upstream pauses are on average tenfold longer than dwells on normal DNA under the same experimental conditions (~100 ms)⁶; thus, the absence of the phosphate charge slows the dwell phase, indicating a sensory role for this contact. The fact that a single kinetic event dominates the upstream pause duration, in sharp contrast to the ~four kinetic events that are known to be rate-limiting for the dwell phase on charged DNA⁶, indicates that the mechanochemical cycle of the packaging motor contains a single kinetic checkpoint—a process that halts the chemical cycle until the DNA is correctly positioned. In contrast, the average duration of the contacts during the burst phase is not significantly modified on neutral DNA, and hence these contacts probably do not involve a significant sensory role.

These dynamics suggest a mechanism for the motor to cross neutral inserts. Successful traversal is a kinetic competition between the increased off-rate of the motor on neutral DNA and the time it takes to successfully complete the mechanochemical events necessary to traverse this modified DNA. There is a small probability of loading the necessary ATP molecules and starting the burst, out-competing the increased slipping rate on the neutral DNA; however, as illustrated by the probabilities in Fig. 2b, the large probability of recovering from a slip allows the motor to attempt to package the insert many times, amplifying this small probability to the large traversal probabilities observed in Fig. 1. This kinetic competition also explains the marked decrease in the ability of the motor to package 30 bp of neutral DNA because successful traversal would require several consecutive complete mechanochemical cycles to out-compete slipping. Finally, the branching probabilities observed (Fig. 2b) predict traversal probabilities consistent with the values in Fig. 1, providing further support for this model (Supplementary Discussion).

The fact that the motor resides for a finite time in the neutral DNA (Fig. 2) provides further evidence for additional, non-charge contacts. To test the role of sugars and bases in these interactions, we created an insert with these elements removed (Fig. 3a). These chemical modifications also disrupt the helical geometry of the DNA; thus, to isolate

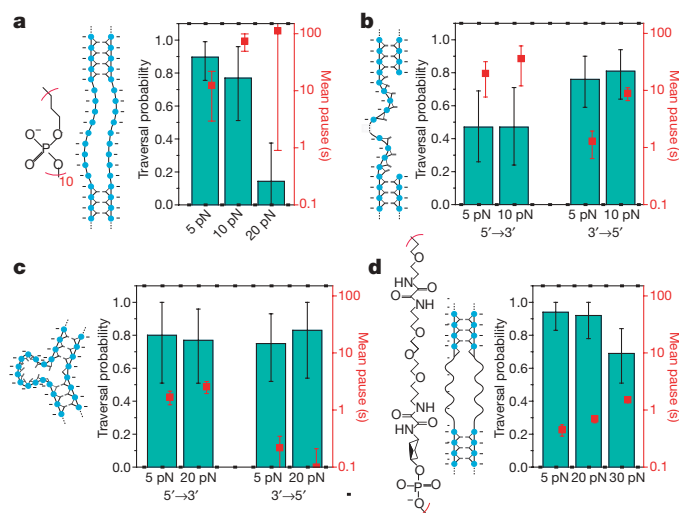


Figure 3 | Substrate promiscuity. **a–d**, The force dependence of traversal probabilities and pause durations of different modified DNA substrates. **a**, 10-nucleotide double-stranded abasic phosphate backbone. **b**, 20-nucleotide ssDNA (poly-AC) on 5'–3' and 3'–5' strands. **c**, 10-bp bulge (poly-AC) on 5'–3' and 3'–5' strands. **d**, Double-stranded linker. The traversal probabilities are plotted using the Laplace estimator²⁶, with 95% confidence intervals from the adjusted Wald method²⁷. The error bars of the pause durations are the s.e.m. (The results for other sizes of bulges and gaps are listed in Supplementary Table 2.)

the role of DNA geometry, we created inserts with no chemical modification yet with large disruptions to the helix; namely, single-stranded gaps (Fig. 3b) and unpaired bulges (Fig. 3c). Remarkably, the ϕ 29 motor is capable of traversing all of these modifications (Fig. 3a–c), and the force dependence of the mean pause durations again rules out a purely diffusive model (Supplementary Table 4). Notably, the motor shows lower traversal probability and higher pause durations for gaps on the 5'–3' strand than those on the 3'–5' strand, with a similar trend seen in pause durations of the bulges, consistent with the important 5'–3' contacts discussed above. Our findings are consistent with the successful packaging of short unpaired bulges in phage lambda¹⁸, although they differ from reported results in ϕ 29 (ref. 19) and T4 (ref. 20), in which single-stranded gaps were not packaged. However, disparate experimental methods probably account for these apparent differences (Supplementary Discussion).

Many of the modifications we have probed (Figs 1 and 3) change many features of the DNA simultaneously; thus, to extract the relative importance of the different chemical moieties while controlling for other changes, we performed a set of multivariate logistic regressions²¹ on different subsets of our data. This analysis provides a quantitative ranking of the importance of the different contacts and their force dependence (Fig. 4 and Supplementary Fig. 2). These regressions confirm the importance of the phosphates every 10 bp on the 5'–3' strand and show that these contacts are only important under the application of force. Regression analysis also shows less important contacts with phosphates on both strands on a smaller distance scale, which, remarkably, remain important when extrapolated to zero applied load. This residual importance may arise from the force the motor generates on the DNA, as opposed to the external load applied optically, supporting our conclusion that these contacts occur during the burst phase.

The regression analysis also provides information on the nature of the non-charge contacts, revealing that important, but minor, contacts are made with bases or sugars (Supplementary Fig. 2a). However, the analysis also predicts that removing all of the characteristic features of nucleic acid—phosphates, bases, sugars and native double-helix structure—will result in a reduced but finite traversal probability (Supplementary Discussion). Thus, a component of these additional, non-charge interactions is not specific to DNA. We tested this

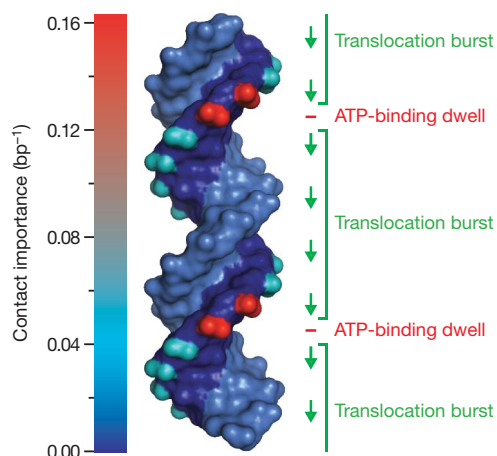


Figure 4 | The motor–DNA contacts. The relative importance of the motor–DNA contacts are marked using the program PyMOL with a quantitative colour scale with magnitudes inferred from the traversal probabilities at 5 pN for the measured modifications (Figs 1 and 3, Supplementary Table 2, Supplementary Discussion and Supplementary Fig. 2). The units of ‘contact importance’ correspond to the inverse of the distance over which the removal of the specific moiety would reduce the traversal probability to 50% (Supplementary Discussion). The mechanochemical phase of the motor as it moves along the DNA is indicated to the right.

surprising result by challenging the motor with a polymer lacking any resemblance to nucleic acid. Notably, the motor packages this insert as well (Fig. 3d), revealing a nonspecific component to the motor–DNA interaction. Such interactions may correspond to chemical contacts, that is, hydrogen bonds or hydrophobic interactions, which do not require a specific nucleic acid moiety. Alternatively, it is possible that these interactions are steric in nature. A prediction of such a steric drive mechanism is that amino acids essential for DNA translocation can be replaced by any residue with a bulky side chain, a behaviour suggested by mutational analysis of the DNA binding loop of the related hexameric helicase SV40 (ref. 22).

Taken together, our data indicate that the motor–DNA interaction involves a wide variety of contacts with the full complement of nucleic acid moieties, as well as contacts not specific to DNA. Furthermore, our data suggest that the type of motor–DNA interaction changes during the course of the mechanochemical cycle. During the dwell phase, when ATP is loaded, the motor maintains strong, load-bearing contact with the DNA via interactions with adjacent charges on the 5′–3′ strand every 10 bp. These contacts have a sensory role, coupling the mechanical and chemical cycles. In the burst phase, when the DNA is translocated, the motor makes contact with a variety of chemical moieties, including charges on both strands, bases or sugars, and additional non-DNA-specific contacts. Given the relatively long duration of the contacts during the dwell phase, specific, strong, ionic interactions may be preferable. In contrast, contacts during active translocation in the burst phase must be made and broken more quickly; thus, it would be advantageous to make weaker, more promiscuous contacts. The important motor–DNA interactions and the specific point in the mechanochemical cycle at which these contacts are made are summarized in Fig. 4.

Our findings have broad implications for the mechanism of the packaging motor. In particular, nonspecific contacts can have an important role in generating a step size that is a non-integer repeat of the chemical periodicity of DNA⁶. Moreover, the observation that important contacts are made below distances of ~10 bp provides evidence against an open-ring model of translocation in which only two subunits contact the DNA, thereby limiting the mechanism by which a non-integer step size can be generated⁶. In parallel, our findings have broader implications for the family of ASCE ring-ATPases. The use of different contacts during distinct portions of

the mechanochemical cycle, as well as the use of sensory contacts, could act as a general mechanism by which multimeric motors synchronize and coordinate the hydrolysis cycles of their individual subunits with the position of their substrate. The use of nonspecific, perhaps steric, contacts by nucleic acid and polypeptide translocases²³ may reflect the existence of a conserved translocation mechanism shared by members of the ASCE superfamily; such nonspecific contacts may have facilitated the evolution of peptide translocases from nucleic acid translocases²⁴.

METHODS SUMMARY

Substrates were prepared by ligating custom oligonucleotide inserts with unmodified DNA. Single-molecule packaging assays were performed as described^{16,7,25}. Insert-induced pauses were identified according to their location in the packaging trace.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

Received 3 June; accepted 20 August 2009.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

Acknowledgements We thank C. L. Hetherington, M. Kopaczynska, A. Spakowitz and J. M. Berger for critical discussions, and D. Reid, M. T. Couvillon and N. L. S. Chavez for preliminary work leading to this publication. K.A. acknowledges the PMMB fellowship through the Burroughs Wellcome Fund, A.T.P. the NIH Molecular Biophysics Training Grant, A.K. the Human Frontier Science Program Cross-Disciplinary Fellowship, J.R.M. the NSF Graduate Research Fellowship, and Y.R.C. the Burroughs Wellcome Fund Career Award at the Scientific Interface for funding. This research was supported in part by the National Institutes of Health (NIH) grants GM-071552, DE-003606 and GM-059604. The content is solely the

responsibility of the authors and does not necessarily represent the official views of the NIH.

Author Contributions K.A., A.T.P., A.K. and J.R.M. conducted the experiments; K.A., A.T.P., A.K., J.R.M. and Y.R.C. performed the analysis; S.G., P.J.J. and D.L.A. prepared and provided experimental materials; and K.A., A.T.P., A.K., J.R.M., Y.R.C., S.G., P.J.J. and C.B. wrote the paper. K.A., A.T.P., A.K. and J.R.M. contributed equally to this work.

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METHODS

DNA substrates. For each construct, two (or in the case of the gaps, three) modified or unmodified DNA oligonucleotides (listed in Supplementary Table 1) were purchased from Fidelity Systems or IDT. The oligonucleotides hybridize to form a DNA segment with 3-nucleotide overhangs. Using PCR, restriction digest and agarose gel purification, 4,187- and 4,008-bp DNA fragments with distinct non-palindromic 3-nucleotide overhangs were prepared. The primer for the 4,008-bp PCR fragment contains a terminal biotin. These fragments were ligated to the insert oligonucleotides and the 8.2-kb product was gel purified.

Single-molecule packaging. Single-molecule packaging assays were performed as previously described^{4,7,25,28} in packaging buffer containing 50 mM Tris-HCl, pH 7.8, 50 mM NaCl, 5 mM MgCl₂ and 1 mM ATP, unless mentioned otherwise. The biotinylated DNA is bound to a 2.1- μ m streptavidin-coated polystyrene bead held by a micropipette and the prohead-gp16-ATPase complex is attached to a second bead held in an optical trap. Packaging is initiated *in situ* by bringing the beads together^{25,28}. The instrument²⁹ is run in force-feedback mode, in which the separation between the beads is adjusted to maintain constant tension in the DNA molecule. High-resolution packaging measurements were conducted on a dual trap instrument described previously³⁰ with materials prepared in the same fashion as described above, except 860-nm beads were used.

Analysis. The modified DNA insert is expected to reach the motor after packaging 4.2 kb. However, variability in attachment geometry of the DNA to the beads and other systematic errors introduce uncertainty in the measurement of tether length, so the insert was considered to be between the 2.5- and 4.5-kb positions. The longest pause in this interval was scored as the insert-induced pause. In traces

in which a pause could not be identified, a pause of 0.1 s, the temporal resolution, was assigned. The rates of pausing (0.017 kb⁻¹) and slipping (0.02 kb⁻¹) during normal packaging⁷ are small enough that these events do not significantly bias our measurements. All probabilities were calculated with the Laplace estimator²⁶, which is given by $P_{\text{Laplace}} = (x + 1)/(n + 2)$, in which x is the number of successes and n is the number of trials. This is a better estimator than the maximum-likelihood x/n , especially when n is small or the probabilities being estimated are near zero or one²⁶. n_{min} values are a measure of the degree to which a distribution is exponential and provide a strict lower limit on the number of rate-limiting kinetic events⁶. These values were calculated by taking the ratio of the mean dwell time squared over the variance in the dwell times. n_{min} error bars are standard deviations and were calculated using a bootstrap method.

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