Molecular sled is an eleven-amino acid vehicle facilitating biochemical interactions via sliding components along DNA

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Recently, we showed the adenovirus proteinase interacts productively with its protein substrates in vitro and in vivo in nascent virus particles via one-dimensional diffusion along the viral DNA. The mechanism by which this occurs has heretofore been unknown. We show sliding of these proteins along DNA occurs on a new vehicle in molecular biology, a ‘molecular sled’ named pVIc. This 11-amino acid viral peptide binds to DNA independent of sequence. pVIc slides on DNA, exhibiting the fastest one-dimensional diffusion constant, $26 \pm 1.8 \times 10^6$ (bp)$^2 s^{-1}$. pVIc is a ‘molecular sled,’ because it can slide heterologous cargos along DNA, for example, a streptavidin tetramer. Similar peptides, for example, from the C terminus of β-actin or NLSIII of the p53 protein, slide along DNA. Characteristics of the ‘molecular sled’ in its milieu (virion, nucleus) have implications for how proteins in the nucleus of cells interact and imply a new form of biochemistry, one-dimensional biochemistry.
The adenovirus proteinase (AVP), a 23 kDa cysteine proteinase, is activated inside a nascent virion, so it can cleave multiple copies of virion precursor proteins used in virion assembly. Only then is the virus particle infectious. AVP is activated by two cofactors, pVlc, the 11-amino acid peptide (GVQSLKRRRCF) from the C terminus of the precursor to adenovirus protein VI, pVI, and the double-stranded linear viral DNA genome of about 35,000 bp (refs 5,6). The cofactors increase the $k_{cat}/K_m$ for substrate hydrolysis. In the presence of DNA, the activity of AVP increases 11-fold; in the presence of pVlc it increases 118-fold. In the presence of both cofactors, the activity of AVP increases synergistically, 34,000-fold. Inside nascent virus particles, the activated AVP and its protein substrate pVI slide on DNA over hundreds to thousands of base pairs, depending in part on the ionic strength, via one-dimensional diffusion to facilitate productive interactions with DNA-bound precursor protein substrates. Facilitation is needed, because the high concentration of viral DNA prevents these sequence-independent DNA-binding proteins from diffusing effectively in a three-dimensional space as they are essentially irreversibly bound to the viral DNA.

AVP, which is synthesized in an inactive form, is partially activated by pVI by forming AVP–pVlc complexes according to the following mechanism: AVP binds randomly to DNA and does not slide. pVI also binds randomly to DNA, but it slides along DNA with a significant one-dimensional diffusion constant, $1.45 \times 10^6$ (bp)$^2$ s$^{-1}$ (ref 12). pVI slides into AVP. AVP, partially activated by being bound to the viral DNA, cleaves pVI first at its N terminus and then at its C terminus. pVIc, released by cleavage of pVI at its C terminus, binds to the AVP molecule that cut it out. A disulfide bond is formed between Cys104 of pVlc and Cys104 of AVP resulting in the very active AVP–pVIc complex, an irreversibly activated form of the enzyme. The processing of the virion proteins, cleavage at >2,000 sites per virion by ~50 AVP–pVIc complexes per virion, occurs by the following mechanism: covalent, active AVP–pVIc complexes bound to DNA, the most active form of the enzyme, slide along the viral DNA with a one-dimensional diffusion constant of $2.1 \times 10^6$ (bp)$^2$ s$^{-1}$ to encounter and cleave precursor proteins, which are also nonspecifically bound to the viral DNA.

Here we ask by what mechanism do pVI and AVP–pVlc complexes slide along DNA to render virus particles infectious? We show that the 11-amino acid peptide pVlc is necessary and sufficient for sliding. pVlc alone binds to DNA independent of DNA sequence and slides along DNA. Furthermore, pVlc is a ‘molecular sled,’ because it not only slides pVI and protein VI along DNA, it can also slide heterologous cargos. For example, streptavidin tetramers do not bind to DNA. Streptavidin tetramers attached to pVlc not only bind to DNA but also slide along DNA. Fragments of human proteins homologous to pVlc, the 8 amino acids at the C terminus of β-actin and a 13-amino acid peptide containing nuclear location signal III, NLSIII, of the p53 protein, exhibit similar DNA binding and sliding activities. This suggests the DNA contour may be a previously unrecognized venue for productive protein–protein interactions via one-dimensional biochemistry.

### Results

#### A role for pVlc in sliding along DNA

By what physical mechanism do pVI and AVP–pVlc complexes slide along DNA to render nascent adenovirus particles infectious (Supplementary Discussion)? Both proteins have in common pVlc, whether as the C terminus of pVI or as the 11-amino acid peptide cofactor covalently attached to AVP in the AVP–pVlc complex. Whereas AVP–pVlc complexes slide on DNA, AVP alone does not slide, although it binds to DNA. This raises the possibility that pVlc is a modular component able to confer sliding ability on proteins to which it is attached, for example, AVP and protein VI. If so, does its binding induce conformational changes that lead to the appearance of a sliding interface able to interact with DNA or does pVlc itself present such a sliding interface?

#### pVlc binds to DNA

If pVlc is a modular component able to confer sliding ability on proteins, then, it may be able to bind to DNA by itself and slide on DNA by itself (Supplementary Note 1). pVlc binds to DNA (Fig. 1a) with a $K_d$ of 26 ± 25 nM (Table 1). Six to seven base pairs of DNA are occluded (Supplementary Note 2) by the binding of one pVlc molecule to DNA (Fig. 1b,c and Table 1). Because the number of pVlc molecules that can bind to DNA is proportional to the length of the DNA, we conclude that binding is independent of DNA sequence (Fig. 1b,c). The amino-acid sequence of pVlc is conserved, Fig. 1d. The crystal structure of pVlc in the AVP–pVlc complex is highlighted in Supplementary Fig. 1; it shows that the pVlc moiety in the complex is exposed to solvent and, therefore, capable of interacting with DNA.

#### pVlc slides on DNA

To see if pVlc slides on DNA, we performed a facilitated diffusion assay (Supplementary Discussion) using total internal reflection fluorescence microscopy with flow-stretched dsDNA at pH 6.5 in 2 mM NaCl and pVlc labelled with a single fluorescent dye molecule. Individual pVlc molecules were observed not only binding to random locations on the DNA as predicted, but, surprisingly, they were observed diffusing along the DNA as well. Indeed, pVlc exhibited robust sliding activity, with many molecules remaining bound to the DNA for longer than a second, some up to 6 s, while sliding over tens of thousands of base pairs (more than half the length of the adenovirus genome). For example, the molecule whose motion is shown in the raw image data in Fig. 2a travelled over ~30,000 base pairs during a 3 second binding event. Despite the optical resolution limit imposed by diffraction, the centroid position (see Methods section) of each signal in each frame was determined with sub-diffraction-limited resolution, <50 nm (red line in Fig. 2a).

The one-dimensional diffusion constant of pVlc was the highest average value yet reported for diffusion along DNA. The trajectories of 35 pVlc molecules sliding randomly in both directions on DNA are plotted in Fig. 2b. The mean square displacement (MSD) for each molecule shown in Fig. 2b is plotted in Fig. 2c. The MSD for each molecule was approximately linear with diffusion time, indicating transport dominated by Brownian motion. From the MSD initial slopes, one-dimensional diffusion constants ($D_0$) were estimated according to $D_0 = \langle \Delta x^2 \rangle / (2 \Delta t)$ and are displayed in the histogram in Fig. 2d. The mean one-dimensional diffusion constant was $26 \pm 1.8 \times 10^6$ (bp)$^2$ s$^{-1}$ with an s.d. of $11 \times 10^6$ (bp)$^2$ s$^{-1}$ (Table 1).

Given this large one-dimensional diffusion constant, we considered the possibility that processes other than sliding along DNA might be occurring (Supplementary Note 3). Sliding consists of one-dimensional translocation along the DNA with the polypeptide in continuous contact with the DNA. Hopping can be mistaken for sliding in tracking experiments. Hopping occurs when the polypeptide repeatedly dissociates from the DNA and rebinds at nearby locations on the DNA, in a microdiffusion process. If hopping were occurring, the diffusion constant should increase as the salt concentration is increased, since the residence time on DNA decreases. Experiments at a 10-fold-higher salt concentration, 20 mM, showed the one-dimensional diffusion constant to be slightly lower, 17.9 ± 3.5 × 10^6 (bp)$^2$ s$^{-1}$ with an s.d. of 10.7 × 10^6 (bp)$^2$ s$^{-1}$.
Figure 1 | Binding of pVIc to DNA. (a) The equilibrium dissociation constant, $K_{\text{d(app)}}$, for the binding of pVIc to DNA was determined by incubating 10 nM Cy3B-pVIc with increasing amounts of 12-mer dsDNA and measuring the change in anisotropy. The experiment was repeated five times. The line through the closed circles represents the nonlinear regression fit of the experimental data to a 1:1 ligand-receptor model. The $K_{\text{d(app)}}$ was 264 ± 25 nM (ref. 14). (b) The stoichiometry of binding of pVIc to DNA was determined by incubating 10 nM Cy3B-pVIc and 10 μM pVIc with increasing amounts of 36-mer dsDNA and measuring the change in anisotropy. The two straight lines were drawn using the data in the filled-in circles. The point with the open circle was not included in the fits for the lines. The intersection point of the two lines is the minimal concentration of DNA required to saturate 10 bp dsDNA and measuring the change in anisotropy. The two straight lines were drawn using the data in the filled-in circles. The point with the open circle was not included in the fits for the lines. The intersection point of the two lines is the minimal concentration of DNA required to saturate 10 bp dsDNA; it indicates six molecules of pVIc saturate one molecule of 36-mer dsDNA. (c) The number of base pairs of DNA occluded by the binding of one pVIc to DNA. (d) The ratio of (pVIc) to (DNA) as a function of DNA length (bp) for the binding of pVIc to DNA was determined by incubating 10 nM Cy3B-pVIc and 10 μM pVIc with increasing amounts of 36-mer dsDNA and measuring the change in anisotropy. The two straight lines were drawn using the data in the filled-in circles. The point with the open circle was not included in the fits for the lines. The intersection point of the two lines is the minimal concentration of DNA required to saturate 10 μM pVIc; it indicates six molecules of pVIc saturate one molecule of 36-mer dsDNA.

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Model for pVIc and AVP–pVIc complexes binding to DNA. A model of pVIc bound to DNA that reflects the simplest interpretation of a body of disparate data, especially the ionic strength dependence of the binding of pVI12 and of AVP–pVIc complexes to DNA14, was generated by moving the crystal structure of the AVP–pVIc complex onto the crystal structure of DNA, Fig. 3a. The only direct contacts are made by the four basic residues of pVIc, which form ion pairs23 with as many as four DNA phosphate groups, Fig. 3b,c. In the crystal structure of the AVP–pVIc complex, the four basic residues of pVIc are part of a β-strand; they are exposed to solvent2. Although we have placed pVIc in the major groove, we cannot rule out binding in the minor groove. If instead of placing the structure of pVIc taken from the atomic coordinates of the AVP–pVIc structure on to the structure of DNA, the structure of the entire AVP–pVIc complex is manually moved to the DNA, Fig. 3d, no additional contacts between the protein and DNA are made. Furthermore, the active site of AVP, coloured pink, shows that it is accessible to being able to bind substrate while the AVP–pVIc complex is bound to DNA. In this model, AVP–pVIc complexes bind to DNA solely via the pVIc moiety, more
To test whether pVIc has general translocase activity, we attached a vehicle capable of carrying any cargo attached to it along DNA.

Figure 2 | One-dimensional diffusion of pVIc along DNA. (a) Rapid motion of a pVIc molecule along flow-stretched dsDNA recorded at 50 frames per second. The kymograph was generated from raw images showing motion along DNA (each vertical strip of pixels is taken from one movie frame) as a function of time (horizontal axis). The red line represents the position estimate in each frame determined using Gaussian centroid determination. (b) Diffusion of pVIc along flow-stretched dsDNA. The 11-amino acid peptide pVIc diffuses rapidly along DNA (x(t), left axis, 35 trajectories). (c) MSD of the trajectories shown in b along the DNA (Δx(t)^2, left axis). (d) Histogram of the diffusion constants for pVIc diffusing along dsDNA. The initial slopes of the MSD of each of the 35 trajectories from pVIc molecules sliding on DNA plotted in c were used to calculate one-dimensional diffusion constants (D_x) according to D_x = <Δx(t)^2>/2Δt. The results are displayed in the histogram; the mean was equal to 26.0 ± 10^6 (bp^2 s^-1) (ref. 14). In b and c motion transverse to the DNA (γ(t) and <Δγ(t)^2>), respectively, right axis) is represented on the same scale, as a control.

pVIc is a molecular sled. This model and extensive biochemical data suggest that pVIc may be a universal 'molecular sled,' a vehicle capable of carrying any cargo attached to it along DNA. To test whether pVIc has general translocase activity, we attached a large, heterologous cargo, the streptavidin tetramer, to pVIc and assayed the complex for DNA binding and sliding activity. In a control experiment, streptavidin did not bind to DNA (Fig. 4a).

However, pVIc–streptavidin complexes not only bound to DNA (Fig. 4a and Table 1), they diffused efficiently along DNA with a one-dimensional diffusion constant of 2.21 ± 0.21 × 10^6 (bp^2 s^-1) (Fig. 4b and Table 1); the lower diffusion constant is consistent specifically via the four basic amino acids. These four amino acids are conserved (Fig. 1d).

| Protein (bp DNA) | M_W (aa) | D_x ± s.e. (bp^2 s^-1) x 10^-6 | s.d. D_x (bp^2 s^-1) x 10^-6 | K_{d(app.)} nM (pH) | DNA footprint (bp) | Notes |
|------------------|----------|-------------------------------|-----------------------------|----------------------|-------------------|-------|
| pVIc (12)        | 1,350 (11) | ND                            | 693 ± 84 (pH 8)              | 7                    | 1, 2, 3           |
| pVIc(low salt)   | 26.0 ± 1.8 | 0.1                          | 264 ± 25 (pH 7)              | 7                    | 1, 4, 5           |
| pVIc(high salt)  | 17.9 ± 3.5 | 0.1                          | ND                           | ND                   | 1, 4              |
| AVP (12)         | 2,087 (204) | (0.02 ± 0.07)                | 63.1 ± 5.8 (pH 7)            | ND                   | 1, 3, 6, 7        |
| AVP-pVIc (36)    | 24,435 (215) | 1.9                          | 4.6 ± 2.2 (pH 7)             | 6                    | 1, 3, 5, 8        |
| AVP-pVIc (high salt) | 17.1 ± 3.5 | 1.6                          | ND                           | ND                   | 1, 4, 5           |
| (pVIc-biotin: streptavidin) (18) | 2.21 ± 0.21 | 0.2                          | 35 ± 5.0 (pH 7.5)            | ND                   | 1, 8              |
| pVI (33)         | 27,014 (250) | 1.45 ± 0.13                  | 46 ± 1.6 (pH 8)              | 8                    | 1, 2, 7, 9        |
| Protein VI (33)  | 22,118 (206) | 1.61                         | 307 ± 38 (pH 8)              | 8                    | 1, 2, 9           |
| 8-Actin-C (12)   | 988 (8)    | 5.45                          | 5.0 ± 0.8 (pH 7)             | ND                   | 1                |
| 11-Actin-C (12)  | 1,230 (11) | 6.40                          | 3.29                         | ND                   | 1                |
| 13-p53-C (30)    | 1,593 (13) | 11.2 ± 0.8                    | 780 ± 96 (pH 7.4)            | ND                   | 1, 10             |

AVP, Adenosine proteinase; ND, not determined; pVIc, peptide derived from C terminus of pVI; pVII, the precursor to protein VI; pVI, Protein VI; Adenosine protein VI. Notes: (1) to convert from bp to nm: 10^6 (bp^2 s^-1) = 102-400 (nm^2 s^-1). For K_{d(app.)} determinations, at pH 7.5 or 8, the dye was fluorescein, and the label was on the DNA; at pH 7 the dye was Cy3B and the label was on the protein. For pVIc-biotin: streptavidin experiments at pH 7.5, the dye used was Alexa Fluor 546, and there were two dye molecules per streptavidin. pVIc was labelled with Cy3B at Cys50; AVP and AVP-pVIc were labelled with Cy3B at Cys199. The actin C-terminal peptides were labelled on their cysteine residue with Cy3B. pVI was labelled at Cys249 with Cy3B. The p53 CTD segment (aa368-388) was labelled at its N terminus with tetramethyl rhodamine. (2) assay buffer was 20 mM Tris-HCl, pH 8, 0.025% DDM, 0.1 mM DTT; (3) ref. 10; (4) binding assay buffer was 20 mM HEPES, pH 7, 0.025% DDM, 10 mM NaCl and 0.1 mM DTT. The NaCl concentration in the sliding assay buffer was 2–6 mM NaCl; in the high salt-sliding assay buffer, 20–25 mM NaCl was present (see Supplementary Text); (5) ref. 14; (6) whole population-mean D_x calculated from one population (99–96% of the molecules bound to DNA) having a D_x of zero and another population (7–4% of the molecules bound to DNA) having a D_x of 1.7 × 10^6 (bp^2 s^-1), with s.d. of 1.9 × 10^6 (bp^2 s^-1); (7) ref. 11; (8) assay buffer was 20 mM Na phosphate, pH 7.5, 0.05% DDM; (9) ref. 12, (10) Assay in 10 mM sodium phosphate buffered at pH 6.5 or 7.4.
are coloured in light blue and dark blue, respectively. (residues of pVIc, the one lysine residue and the three arginine residues, complex (PDB ID: 1NLN) is shown docked to the DNA. The four basic residues of pVIc, the one lysine residue and the three arginine residues, are coloured in light blue and dark blue, respectively. (The DNA-pVIc complex was rotated ~90° on its x axis to show the contacts of the peptide with the major groove. (d) The AVP-pVIc structure is displayed, showing that DNA binding is dominated by the pVIc moiety. The majority of the binding enthalpy between the protein-peptide complex and DNA is likely to originate through electrostatic interactions between the four, contiguous, basic residues of pVIc (KRRR, in blue) in the AVP-pVIc complex and phosphate groups (in red) in the backbone of the dsDNA. The active site of AVP is coloured pink, showing that the proteinase active site is sterically unhindered by DNA binding, consistent with the proteinase being able to bind to and cleave substrates while bound to and sliding on DNA. The figure was rendered using PyMol (The PyMOL Molecular Graphics System, Version 1.2r.3pre, Schrödinger, LLC).

with the larger size of streptavidin tetramers compared with AVP, in their corresponding complexes with pVIc. Thus, pVIc is a ‘molecular sled’ that can transport heterologous cargos along DNA. This is also the most convincing evidence that pVIc contributes the entirety of the sliding contacts between protein and DNA in all the contexts where we observe sliding activity in adenovirus proteins.

Figure 3 | Model for the binding of pVIc and AVP-pVIc complexes to DNA. (a) The structure of a B-form DNA dodecamer (PDB ID: 1HQ7) is displayed with four of its phosphate groups coloured in red. (b) The structure of pVIc obtained from the crystal structure of the AVP-pVIc complex (PDB ID: 1NLN) is shown docked to the DNA. The four basic residues of pVIc, the one lysine residue and the three arginine residues, are coloured in light blue and dark blue, respectively. (c) The DNA-pVIc complex was rotated ~90° on its x axis to show the contacts of the peptide with the major groove. (d) The AVP-pVIc structure is displayed, showing that DNA binding is dominated by the pVIc moiety. The majority of the binding enthalpy between the protein-peptide complex and DNA is likely to originate through electrostatic interactions between the four, contiguous, basic residues of pVIc (KRRR, in blue) in the AVP-pVIc complex and phosphate groups (in red) in the backbone of the dsDNA. The active site of AVP is coloured pink, showing that the proteinase active site is sterically unhindered by DNA binding, consistent with the proteinase being able to bind to and cleave substrates while bound to and sliding on DNA. The figure was rendered using PyMol (The PyMOL Molecular Graphics System, Version 1.2r.3pre, Schrödinger, LLC).

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Conservation and ubiquity of the pVIc amino-acid sequence. The four, contiguous basic amino acids of pVIc (KRRR) function not only in binding to and sliding on DNA but also function as a nuclear location signal (NLS) in its pVIc moiety. The amino-acid sequence of pVIc is conserved among adenoviruses, Fig. 1d. Across human adenovirus serotypes, as well as among simian, porcine and murine adenoviruses, there is strict conservation of the KRRR motif within pVIc. Are there cellular proteins with sequences homologous to pVIc that slide on DNA? Since the NLS sequences of many nuclear-targeted proteins contain contiguous stretches of basic amino acids like those in the NLS in pVIc, is it possible that proteins that enter the nucleus of a cell via an NLS bind to and slide on DNA to interact with other proteins?

C-terminal peptides from β-actin slide on DNA. Are there cellular proteins with sequences homologous to pVIc that slide on DNA? The C terminus of the major human protein β-actin, like pVIc, can stimulate AVP activity. Of the last eight amino acids of β-actin (SIVHRKCF), four are identical and three are homologous to the last eight amino acids of pVIc (SLKRRRRCF), Fig. 1d. The three contiguous basic residues at the C terminus of β-actin constitute the single longest stretch of basic residues in this acidic protein. Peptides comprised of the last 8 amino acids of β-actin, 8-actin-C and the last 11 amino acids of β-actin, 11-actin-C (SGPSIVHRKCF), bound tightly to DNA (Table 1) and diffused rapidly along it (Fig. 4c and Supplementary Fig. 3a,c). The MSD for each molecule was approximately linear (Supplementary Fig. 3b,d) yielding mean one-dimensional diffusion constants of 5.4–6.4 × 10^6 (bp)^2 s^-1, Table 1. β-actin has been found in virus particles and in the nucleus of cells and has been shown to be involved in chromatin remodelling and gene activation.

Peptide with an NLS from the p53 protein slides on DNA. Are there proteins that bind to and slide along DNA using an NLS as a molecular sled? The extremely important tumour-suppressor protein p53 has been shown to slide on DNA, and sliding activity has been localized to its C terminus. p53 contains three functional NLSs, including two in the C-terminal domain (CTD; aa 363–393). A 13-amino acid peptide fragment of the p53 CTD (STSRHKKLMFKTE, aa 376–388) containing NLSIII (RHKKLM) binds to DNA (Table 1) and exhibits robust sliding activity with a measured mean diffusion constant of 11.2 ± 0.8 × 10^6 (bp)^2 s^-1 (Fig. 4d, Table 1 and Supplementary Fig. 4a,b). Thus a peptide with an NLS from a cellular protein can bind to and slide along DNA.

Discussion
Here we asked the question: by what mechanism do two adenovirus proteins, a substrate and an enzyme, slide along DNA to form productive collisions, collisions that lead to binding? We showed they slide via pVIc, an 11-amino acid peptide. pVIc itself binds to DNA with a nM equilibrium dissociation constant; binding is independent of DNA sequence; once bound, pVIc covers 6–7 base pairs, and its diffusion constants of 5.4–6.4 × 10^6 (bp)^2 s^-1 with an s.d. of 11 × 10^6 (bp)^2 s^-1 (ref. 14). Is pVIc a ‘molecular sled’ capable of carrying cargos such as AVP or protein VI and possibly heterologous proteins or does it induce sliding by the adenovirus proteins by inducing a conformational change that enables them to slide? Streptavidin does not bind to DNA, yet a streptavidin-pVIc complex not only binds to DNA but also slides along DNA. Thus pVIc is a ‘molecular sled’ that can slide not only adenovirus proteins along DNA but also heterologous cargos along DNA. A peptide similar in sequence to pVIc, the C terminus of β-actin, was shown to slide on DNA. A model is presented that depicts pVIc binding to DNA via iron pairs formed between its four basic amino-acid residues and four phosphate groups on DNA. These four basic amino acids of pVIc constitute a functional NLS raising the possibility that proteins with similar NLSs slide on DNA. We showed that a peptide containing NLSIII from the p53 protein slides on DNA.
Figure 4 | Sliding of heterologous proteins and peptides. (a) The equilibrium dissociation constant for the binding of (pVlc–biotin)–streptavidin to 18-mer dsDNA was determined by fluorescence resonance energy transfer. The quenching of the fluorescence intensity of the donor molecule, fluorescein-labelled 18-bp dsDNA, as a function of the concentration of the acceptor molecule (pVlc–biotin)–streptavidin Alexa Fluor 546 is shown by the closed circles. The relative fluorescence intensity is the fluorescence intensity at a specific concentration of acceptor divided by the initial fluorescence intensity of the donor in the absence of acceptor. The line through the closed circles represents the nonlinear regression fit of the experimental data to a 1:1 ligand–receptor model. The open circles represent data from the titration of 10 nM fluorescein-labelled 18-mer dsDNA with streptavidin Alexa Fluor 546. These data indicated that streptavidin Alexa Fluor 546 did not bind to DNA. (b) (pVlc–biotin)–streptavidin complexes diffuse rapidly along DNA (x(t), left axis, 106 trajectories). Motion transverse to the DNA (y(t), right axis, is represented on the same scale, as a control. (c) A peptide (SVHHRKCF) with the last 8 amino acids of β-actin diffuses rapidly (69 trajectories) along DNA x(t). (d) The 13-amino acid peptide of NLSIII of the p53 protein (STSRHKLMFKTE) diffuses rapidly along DNA at pH 6.5 (45 trajectories).

That pVlc slides on DNA by itself was a surprise; the 11-amino acid peptide lacks the large, structured DNA-binding interface present in proteins previously shown to slide on DNA. Consistent with this is the observation that in the crystal structure of the AVP–pVlc complex, pVlc is tightly bound to AVP. There are 24 non-β-strand hydrogen bonds, 6 β-strand hydrogen bonds and one-covalent bond. A disulfide bond is formed between Cys104 of AVP and Cys107 of pVlc. Structured DNA-binding interfaces have been presumed necessary to achieve the low free-energy barriers to sliding (comparable to k_BT) that are needed for effective searching along DNA. This is understood to be achieved in sequence-non-specific protein–DNA complexes by indirect or frustrated protein–DNA contacts that ‘lubricate’ the binding interface. Such ‘buffered’ protein–DNA contacts are thought to support rapid sliding by minimizing differences in free energy as the protein translates along the DNA and have been observed in protein–DNA complexes (interfacial water, long-range interactions, for example, between helix dipoles and the DNA). One-dimensional diffusion constants for some proteins are known to be sensitive to point mutations and the presence of individual charged residues. Together, these observations established the expectation that precision-tuned, three-dimensional structures are required for high-speed sliding. However, we observed both robust and persistent sliding on DNA by pVlc, a peptide predicted by several secondary structure prediction programs to have no stable secondary structure. The mean one-dimensional diffusion constant of pVlc was the highest yet reported for any object sliding along DNA, 26 ± 1.8 × 10^6 (bp)^2 s^-1 (Table 1).
parameters of the hOgg1–streptavidin complex very similar to those of the pVIc–streptavidin complex. The diffusion constant of the hOgg1–streptavidin conjugate was measured as $2.2 \times 10^6$ (bp)$^2$ s$^{-1}$. This value is consistent with the prediction from the physical model as were the one-dimensional diffusion constants of the pVIc–streptavidin complex and the AVP–pVIc complex (Table 1).14

How do AVP–pVIc complexes and pVIc bind to DNA? Experiments on the ionic strength dependence of the sliding of AVP–pVIc complexes and of pVI on DNA are consistent with the pVIc moiety mediating the binding to DNA. The model in Fig. 3 depicts the sliding interface as consisting of the four contiguous basic amino acids of pVIc forming ion pairs with up to four phosphate groups on the DNA. The model is consistent with a binding footprint on DNA of 7 bp for pVIc, the AVP–pVIc complex and pVI (Table 1). The major factor driving the non-specific interaction between AVP–pVIc complexes6,10,12 and pVI and DNA comes from the entropic contribution on the release of counter ions. The sequence-non-specific interactions between AVP–pVIc complexes and pVI and DNA exhibit a substantial dependence on the monovalent sodium ion concentration. This dependence reflects the electrostatic component of the binding reaction47. The electrostatic component originates from the formation of ion pairs between positively charged groups on the AVP–pVIc complex or on pVI and negatively charged phosphate groups on DNA. After binding occurs, there is a concomitant release of counter ions from the DNA and, possibly, from AVP–pVIc complexes or pVI. From an analysis of the equilibrium association constants for the binding of AVP–pVIc complexes or pVI to 12-mer dsDNA as a function of the Na$^+$ concentration, an accurate estimate of the number of ion pairs involved in the interaction was obtained. For AVP–pVIc complexes and for pVI, 2.2 (ref. 10) and 2.9 (ref. 12) ion pairs, respectively, were involved in complex formation with 12-mer dsDNA.

How does pVIc slide on DNA? Dynamic formation and breakage of the ionic contacts may explain the observed motility and processivity of AVP–pVIc complexes and pVIc on DNA.46 During movement of the sled along DNA, one or more of the four potential ion pairs could break and re-form elsewhere; concomitantly, the other ion pairs could remain intact, holding the moving peptide to at least one part of the DNA at all times. This is consistent with the average of 2–3 ion pairs observed in pVIc-mediated protein–DNA complexes.10,12 Multiple binding sites used simultaneously could explain the high processivity observed in these single-molecule translocation experiments. We had also previously shown that sliding of AVP–pVIc complexes occurs along a helical path (Supplementary Note 4) with the protein complex rotating as it moves so that its DNA-binding face maintains contact with the DNA.15

The model for the binding of pVIc and AVP–pVIc complexes to DNA explains a long-standing enigma about the crystal structure of the AVP–pVIc complex: Why is pVIc which exerts powerful control on the rate of catalysis, bound quite far from the active-site residues involved in catalysis? A secondary structure model of the AVP–pVIc complex with the atoms of the pVIc residue colour coded is shown in Supplementary Fig. 1. The pVIc cysteine residue, which forms a disulphide bond with Cys104 of the main chain, is 32 Å away from the active-site nucleophile Cys122. The residue of pVIc closest to the active site is Val$^2$, whose side chain is 14.5 Å from Cys122. Heretofore, it had been thought that the only function of pVIc was to act as a cofactor by increasing the catalytic power of the active site. However, knowing a second function of pVIc, as described here, namely sliding of AVP along DNA, it became clear why the pVIc-binding site must be far away from the active site: if the active site of AVP were close to pVIc bound to DNA, that configuration may physically interfere with the ability of the active site to recognize and cleave a precursor protein also bound to the same DNA molecule. Separation of the active site from pVIc enables the enzyme active site to interact readily with its seven, different substrates.

Current views on one-dimensional diffusion of proteins along DNA are mostly concerned with proteins that locate and bind to a specific site on DNA, proteins involved in DNA metabolism. Ionic strength is thought to be a major factor in determining the search dynamics. For example, in low salt in vitro, the lac repressor binds to its operator with a second-order rate constant of $7 \times 10^9$ M$^{-1}$ s$^{-1}$ (ref. 49). However, in the presence of 100 mM KCl, the rate falls 100-fold50, to the diffusion-limited rate$^{41,52}$. The reason given for this extremely fast interaction at low ionic strength is that there is an electrostatic attraction between a positively charged site on the repressor and the negatively charged phosphate groups on the DNA. The decrease in the association rate at physiological ionic strength is due to the solvent counter ions screening the electrostatic attraction. From this, it was concluded that at physiological ionic strength the rate-determining step in binding to a specific DNA site is the diffusion of the protein to the DNA molecule, not the transfer of the protein molecule from a non-specific site to a specific site.53

Here we are concerned with how and why some adenovirus proteins inside the virion at physiological ionic strength locate and productively bind to other proteins via one-dimensional diffusion on the viral DNA. What is the rate-determining step in this environment? Our sliding assays (Supplementary Discussion) were done under non-physiological conditions, at low ionic strength. Raising the ionic strength to physiological conditions (0.15 M NaCl) will decrease the association rate of a protein binding to DNA; however, the enormously high concentration of DNA in the virion54 will increase the association rate of a protein binding to DNA. The second-order rate constant for binding of a protein to DNA is proportional to the DNA binding site concentration, which is huge, 0.5 M in the virion. Also, increasing the ionic strength to physiological conditions will have a minimal effect on the one-dimensional diffusion constant for sliding along DNA as that process is insensitive to ionic strength.

So, what are the factors that determine the three-dimensional rate of diffusion of pVI and AVP–pVIc complexes in the virion? Certainty the rate of diffusion in three-dimensional space is much faster, by orders of magnitude$^{48,53}$, than it is in the one-dimensional space on the DNA contour. But, is that effectively true for some adenovirus proteins inside the adenovirus virion? It is not. Inside the virion, two factors act together to reduce the effective three-dimensional bimolecular association rate between the proteinase and its substrates. The high concentration of DNA drives these sequence-independent DNA-binding proteins (the proteinase and its seven different substrates)$^{4,5}$ onto the DNA by mass action. For example, in an adenovirus virion, the concentration of DNA-binding sites for AVP–pVIc complexes is ~500 mM (ref. 54), and the $K_d$ for the binding of AVP–pVIc complexes to DNA at 0.15 M NaCl (ref. 10) is 10 mM. Therefore at equilibrium, the ratio of protein free to protein bound in the virus particle, $(K_d(DNA))^{-1}$, is $2 \times 10^{-3}$. This means that at any one time, for each AVP–pVIc molecule not bound to DNA, there will be 50,000 AVP–pVIc molecules bound to and presumably sliding along DNA. In a virion with 50 molecules of AVP only 0.001 molecules of AVP will be free; 99.998% of the 50 AVP–pVIc complexes will be bound. Thus, most of the time, almost none of the AVP–pVIc complexes is free in solution in a given virion. And this would diminish, by lowering the duty cycle for three-dimensional diffusion, the ‘effective’ three-dimensional diffusion constant by a similar factor, 50,000-fold. A similar situation occurs with pVI. At a NaCl concentration of 0.001 M,
the estimated $K_d$ for the binding of pV1 to DNA is 0.027 nM (ref. 12); this increases $>350,000$-fold, to 9.5 μM, at physiological ionic strength. However, at 0.15 M NaCl, the ratio of protein free to protein bound, $(K_d)/(DNA)^{-1}$ in the virus particle, is $1.9\times 10^{-5}$.

Even though at high ionic strength the residence time per event will be decreased, the net amount of time a protein is sliding on DNA will be quite large. This is because there will be numerous short sliding events, over hundreds of base pairs, as opposed to fewer but longer sliding events. An added benefit of high ionic strength and DNA concentration is the amount of unique DNA scanned per unit time by sliding; this will be increased since, presumably, hopping or jumping will occur after each dissociation event thereby reducing the redundancy of the one-dimensional search. This balance between one-dimensional searching and dissociation/rebinding would have the additional benefit of enabling efficient bypass of obstacles on the DNA.

The second factor that reduces the effective three-dimensional bimolecular association rate between the proteinate and its substrates is the mesh size of the DNA matrix. We calculate the mesh size of the DNA matrix inside the virion to be $<1$ nm (ref. 48). Three-dimensional diffusion was shown to be reduced by as much as an order of magnitude when the mesh size of a dynamic polymer network equals the size of the diffusing object. The AVP–pV1c complex is ovoid with dimensions of $41\times 44\times 55 \text{Å}$ (refs 1,2). Since the diameter of the AVP–pV1c complex is much larger than the mesh size, and the DNA network has restricted mobility by being packed in the small volume of the virion, a further factor of 10 in the reduction of the diffusion constant is a conservative estimate as to the effect of the DNA matrix on the three-dimensional diffusion of AVP–pV1c complexes.

And, there is no other way of compensating for these reductions in the effective diffusion rate. For example, the DNA genome cannot move to promote protein–protein interactions as it is jammed inside the virion because of being packaged under high pressure. The pressure exerted by the DNA on the virion walls, possibly in excess of 100 atm (ref. 56), results in a large friction between the DNA and the inner surface of the virion, freezing the DNA in place and rendering DNA-bound proteins likewise immobile.

Together, the enormous DNA concentration in the virion, which diminishes the three-dimensional diffusion constant by 50,000-fold and the mesh size of the DNA, which diminishes the three-dimensional diffusion constant by a factor of 10, indicate the ‘effective’ three-dimensional diffusion constant would be reduced by a factor of over 50,000. Thus, the reason these adenovirus proteins form productive collisions via one-dimensional diffusion along DNA is that they have no choice; doing so by three-dimensional diffusion is impossible. In this case, the rate-determining step is not the diffusion of a protein molecule to a DNA molecule but the transfer of a molecule along the DNA to its target protein.

A similar situation probably exists in the nucleus of mammalian cells and in the nucleoids of bacterial cells, where the concentration of DNA binding sites is also high. A discussion of initial diffusion-limited reaction rates among protein molecules and approximate lower limits of time to react half the molecules in a population of molecules under different DNA-binding and one-dimensional diffusion scenarios is presented in Supplementary Table 1 and in Supplementary Discussion.

The data shown here that elucidated the mechanism of sliding of two adenovirus proteins have several interesting and potentially important implications. The discovery that short, basic peptides can confer a DNA binding and sliding activity on a protein vastly increases the possible number and types of proteins that may bind to DNA nonspecifically and exhibit sliding activity.

Tight DNA binding is not necessary for effective sliding activity. Since the concentration of DNA binding sites in the nucleus of mammalian cells and the nucleoid of bacterial cells is high (similar to that in adenovirus virions), even proteins with short basic sequences that have weak, non-specific DNA-binding affinity will be bound most of the time to DNA, being driven there by the DNA concentration. For example, that pV1c with its NLS sequence (KRRR) binds to DNA and slides AVP–pV1c complexes and pV1 on DNA and that NLSII from the p53 protein binds to and slides along DNA imply that possibly all proteins that enter the nucleus of a cell with similar, basic NLSs may bind to and slide along DNA. If so, this will have a significant effect on protein–protein association rates in these environments. Presumably all NLSs must be on the surface of a protein to be able to be recognized by the nuclear import apparatus. That the activity of ‘molecular sleds’ seems to be encoded purely in their primary sequences raises the tantalizing possibility that DNA binding and sliding activity may be predicted from genomic data. This is how we discovered the sliding activity at the C terminus of β-actin and NLSIII of the p53 protein.

To date, one-dimensional diffusion along DNA has been found almost exclusively within the purview of proteins that search for a specific locus on the DNA genome. We had previously observed that the one-dimensional compartment along DNA can also support, promote and regulate transactions between proteins. For example, sliding on DNA is required for the activation of AVP by pV1. If only a simple bimolecular interaction between AVP and pV1 were needed for activation, then mixing purified AVP7 with purified pV1 should result in the cleavage of pV1 to yield pVc followed by the formation of active AVP–pV1c complexes. However, when this was done, no enzymatic activity was detected, even at μM concentrations of AVP and pV1. DNA is required for the activation of AVP by pV1. Both AVP and pV1 bind to DNA with apparent equilibrium dissociation constants of 63 nM (ref. 10) and 46 nM (ref. 12), respectively. When we repeated the experiment but in the presence of dsDNA, by 1 h, 100% of the pV1 was cleaved and used to form active AVP–pV1c complexes. Thus, activation of AVP to AVP–pV1c complexes by pV1 required the presence of DNA. Furthermore, AVP and pV1 both must be on the same DNA molecule for activation to occur as activation can only occur in cis, not trans11. Thus, sliding along DNA is required for the activation of AVP by pV1 to AVP–pV1c complexes.

That protein–protein interactions can be regulated by DNA binding and one-dimensional diffusion greatly increases the scope of biochemistry mediated by facilitated diffusion on DNA: it also has implications for biotechnology. Targeting protein–protein interactions to the DNA genome localizes their interaction in the cell. And DNA might be more than just a ‘highway’ accelerating reactions. It could have a dramatic effect on the specificity of protein–protein interactions by introducing new mechanisms. On the DNA contour, bimolecular interactions may not only be regulated by one-dimensional diffusion constants, but also by the relative orientations of the DNA-bound proteins. If, for example, an enzyme and its substrate become optimally orientated for a productive collision when both are bound to the same DNA helix, the fraction of productive enzyme-substrate collisions mediated by sliding may be orders of magnitude greater, possibly as high as 1/1, far higher than in the case of the random orientation collisions that occur in three-dimensional space.

In conclusion, we have identified and characterized the physical mechanism by which two adenovirus proteins slide on DNA. They use a ‘molecular sled,’ a new vehicle in molecular biology, an 11-amino acid peptide that slides on DNA by itself or,
when attached to a heterologous cargo, can carry the cargo along DNA. Sliding along DNA is the only way sequence-independent DNA-binding proteins can interact with each other in certain milieu, for example, virion or nucleus, have implications for DNA-binding proteins can interact with each other in certain milieu, for example, virion or nucleus, have implications for DNA. Sliding along DNA is the only way sequence-independent when attached to a heterologous cargo, can carry the cargo along DNA. Characteristics of the ‘molecular sled’ in its milieu, for example, virion or nucleus, have implications for how proteins can interact and imply that a new form of biochemistry, one-dimensional biochemistry, is operative.

**Methods**

**Materials.** pVlc (GQVSLKRRRCF), streptavidin Alexa Fluor 546 conjugate, 5’-fluorescein-labelled 12-mer ssDNA (5’-GAGACAGTACGAT-3’), 5’-fluorescein-labeled 18-mer ssDNA (5’-CAGGAAAACAGTCTGAC-3’), and 5’-fluorescein-labeled 36-mer ssDNA (5’-GATTTGACATGAGTGTGCTGGATGTGATAG TGA-3’) were purchased from Invitrogen (Carlsbad, CA). Labeled ssDNAs were annealed to their complimentary strands according to standard protocols. Octylglucoside was purchased from Fisher Scientific (Faden, NJ). The concentrations of pVIc, 8-Actin-C and 11-Actin-C were determined by high-pressure liquid chromatography Research Genetics (Huntsville, AL, USA). Cy3B-maleimide was purchased from GE Healthcare (Piscataway, NJ). Biotin-PEG-3K maleimide was purchased from Nektar. Octylglycoside was purchased from Fisher Scientific (Faden, NJ). N-decyl-β-D-maltopyranoside (DDM) was purchased from Anatrace (Maumee, OH). The concentrations of pVlc, 8-Actin-C and 11-Actin-C were determined by titration of the cysteine residue with Ellman’s reagent using an extinction coefficient of 14,150 M⁻¹ cm⁻¹ at 412 nm for released thionitrobenzoate. The N-terminal TMR-labelled human p53 CTG segment (AA 376-388) was purchased from the Biopolymer & Proteomics Laboratory at MIT (> 85% purity).

**Binding of pVlc to DNA.** The equilibrium dissociation constant, **Kd(lipp)**, for the binding of pVlc to DNA was determined by incubating 10 nM Cy3B-pVlc in 20 mM HEPES (pH 7), 10 mM NaCl, 0.025% DDM, and 0.1 mM dithiothreitol (DTT) with increasing amounts of 12-mer dsDNA and measuring the change in anisotropy. The line through the closed circles in Fig. 1a represents the nonlinear regression fit of the experimental data to a 1:1 ligand–receptor model. The **Kd(lipp)** was 264 ± 25 nM (ref. 14). The stoichiometry of binding of pVlc to DNA was determined by incubating 10 nM Cy3B-pVlc and 10 μM pVlc with increasing amounts of 36-mer dsDNA in 10 mM Tris-HCl (pH 8) and 2 mM β-octylglucoside and measuring the change in anisotropy, Fig. 1b. The two straight lines were drawn using the data in the filled-in circles. The point with the open circle was not included in the fits to the lines. The intersection point of the two lines is the minimal concentration of DNA required to saturate 10 molecules of pVlc; it indicates six molecules of pVlc saturate one molecule of 36-mer dsDNA. The number of base pairs of DNA occluded by the binding of one pVlc to DNA: the experiment in Fig. 1b was repeated but with 12-mer and with 18-mer dsDNA, and the stoichiometries of binding plotted versus the DNA length in base pairs, Fig. 1c. The lines represent the number of molecules of pVlc occupying a 18-mer dsDNA. For determining the **Kd** of the 13-amino acid peptide containing NLSIII from the p53 protein, the fluorophore-labelled peptide concentrations were kept constant at 10 nM, and the 30 bp dsDNA concentrations ranged from 0 to 3 μM.

**Sliding assay.** The sliding assay, based on total internal reflection fluorescence microscopy, used phage λ DNA molecules (48,302 bp) attached at one end to a glass cover slip surface by a biotin linkage9,10. A flow cell was constructed on top of the coverslip such that when a laminar flow of buffer is applied, the DNA strand slides in the direction of flow and orient itself parallel with and in close proximity to the surface of the cover slip. Evanescent waves (532 nm) from a LASER reflecting off the coverslip-buffer interface are used to illuminate a very small volume (~ 1 pl) within a few hundred nanometres of the glass surface. The interaction of a single, fluorescently labeled protein with DNA was visualized with low fluorescence background by wide-field microscopy using a sensitive CCD camera (Photometrics Cascade 512:B) or scientific CMOS (Hamamatsu Orca Flash 4.0). Despite the optical resolution limit imposed by diffraction, the centroid position of each signal in each frame is determined with sub-diffraction-limited resolution, typically 10-30 nm for bright fluorescent conjugates. Labeled peptides or proteins were infused at concentration of 1-2 nM at rates of 20-50 μl h⁻¹. High flow rates were chosen to drive the longitudinal DNA fluctuation faster than the imaging frame rate16. The assay buffer consisted of 10 mM MES (pH 6.5), 2-25 mM NaCl, 50 μM EDTA, 20 mM ethanol, 5% (v/v) glycerol, 0.1 M DTT or β-mercaptoethanol. ‘High salt’ measurements were conducted with 2-6 mM NaCl, ‘higher salt’ measurements were conducted with 20-25 mM NaCl. **Fluorescent labelling of peptides.** The cysteine side chains of pVlc, 8-Actin-C, 11-Actin-C were labelled at a concentration of 200 μM in 25 mM Hepes (pH 7.0), 25 mM NaCl, and 20 mM ethylene by the addition of Cy3B-maleimide to 600 μM. Labelling reactions were incubated at 21 °C in the dark for 2.5 h. Dye-conjugated peptides were purified from unreacted dye and peptide on a 15 cm × 4.6 mm Discovery C18, 5 μm column. Peptides were eluted via a linear acetonitrile gradient from 0% (v/v) acetonitrile to 100% (v/v) acetonitrile in 40% (v/v) acetonitrile. The concentration of the dye-conjugated peptide was determined by measuring the dye concentration spectrophotometrically using εCy3B-fluor = 130,000 M⁻¹ cm⁻¹. The conjugated peptide was aliquoted, dried and stored at -20 °C.

**Alexa fluor 546-streptavidin-biotinylated pVlc conjugates.** Alexa Fluor 546-streptavidin-biotinylated pVlc conjugates were formed as follows: pVlc was biotinylated by incubating 3.4 mM pVlc in 100 mM sodium phosphate (pH 7.0) and 20 mM ethylenediamine tetraacetic acid (pH 8.5) with Ellman’s reagent for 30 min. The biotinylated pVlc was purified on a Discovery C18 column equilibrated in 0.1% (v/v) trifluoroacetic acid using a linear gradient of acetonitrile. Fractions off the column that stimulated AVP activity and were unreactive to Ellman’s reagent were pooled. Next, ~850 pmol of the biotin-PGEG-3K were serially diluted 1:2 with acetonitrile and each dilution lyophilized to dryness using a Labconco CentriVap. Then, a constant volume of 16.7 μM Streptavidin Alexa Fluor 546 was added resulting in molar ratios of (pVlc-biotin) to streptavidin Alexa Fluor 546 of 1:1, 2:1, 4:1, 8:1 and 16:1. After incubation for 1 h, the conjugates were stored at 4 °C.

**Fluorescence anisotropy.** Steady-state fluorescence anisotropy measurements were performed using an ISS model PC-1 photon counting spectrophluorometer (ISS, Champaign, IL) equipped with a 495 nm excitation wavelength. The parallel and vertical emission components for the Cy3B dye, the excitation wavelength was 564 nm, with 8 nm slits placed before and after a monochromator. The parallel and vertical emission components were measured through a 580 nm bandpass filter with a FWHM of 10 nm. For fluorescein dye, the excitation wavelength was 495 nm, with 8 nm slits placed before and after a monochromator. The parallel and vertical emission components were measured through a 530 nm long-pass filter. For measurements on the M5 instrument, samples were prepared in a 96-well plate, and the excitation/emission/cutoff wavelengths were set at 544/590/570 nm (alternatively 590 nm cutoff; both cutoff settings generating substantially identical **Kd** values from fits).

**Calculation of **Kd** values.** The **Kd** was calculated by fitting the fluorescence anisotropy data to a one-to-one stoichiometry binding model according to the equation:

\[ r_{\text{obs}} = r - \left( r_{1} - r_{f} \right) \left( \frac{[P_{1}]}{[P_{1}] + [D_{1}] + K_{d}} \right) \left( \frac{[P_{1}]}{[P_{1}] + [D_{1}] + K_{d}} \right) \left( \frac{[P_{T}]}{[P_{T}] + [D_{T}] + K_{d}} \right) \]

where \( r_{\text{obs}} \) is the observed anisotropy; \( r_{1} \) the anisotropy of free 12-mer dsDNA; \( r_{f} \) the anisotropy of protein bound DNA; \([P_{1}]\) the total protein concentration; \([D_{1}]\) the total DNA concentration (10 nM); and \( K_{d} \) is the equilibrium dissociation constant. The parameters in the nonlinear regression analysis were \( K_{d} \) and \( K_{d} \). **Fluorescence resonance energy transfer.** Steady-state fluorescence measurements were measured with an ISS PC-1 spectrophluorometer (ISS, Champaign, IL) with a 300-W xenon arc lamp and 19 amp lamp current. The excitation and emission wavelengths were 490 and 520 nm, respectively, using 8 nm excitation and emission slits.
emission slits. A 1 ml solution of 10 nM fluorescein-labelled 18-mer dsDNA in 20 mM sodium phosphate, pH 7.5, 0.05% DDM was placed inside a 1 cm standard quartz cuvette and was titrated with increasing amounts of streptavidin Alexa Fluor 546 or the complex (pVlc-biotin)-streptavidin Alexa Fluor 546. The solution was mixed and allowed to equilibrate for 2 min before measuring the fluorescence intensity for a maximum of 10 s (an average of 11 measurements). To correct for the decrease in fluorescence intensity due to inner filter effects, a 10-nM solution of streptavidin Alexa Fluor 546 was titrated with streptavidin Alexa Fluor 546.

**Centroid determination and analysis of trajectories.** Commercial particle-tracking software (Diatrack 3.0, Semasoph, Switzerland) and custom particle-tracking code developed in house were used to identify individual DNA-bound protein molecules and track their trajectories. Molecules exhibiting transverse displacements consistent with flow-stretched DNA were included in the analyses, while molecules exhibiting trajectories characteristic of surface adsorption were excluded. Molecular trajectories were analysed by methods similar to those previously published.[18]

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Author contributions
W.F.M. and P.C.B. designed the experiments and supervised the work. W.J.M., V.G., K.X. and P.C.B. carried out the experiments. W.F.M., W.J.M., K.X., V.G. and P.C.B. processed the data. W.F.M., W.J.M. and P.C.B. wrote the manuscript.

Additional information
Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: W.F.M., W.J.M., V.G. and P.C.B. are inventors on a patent application describing uses of a ‘molecular sled’. The remaining author declares no competing financial interest.

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