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Temperature controlled high-throughput magnetic tweezers show striking difference in activation energies of replicating viral RNA-dependent RNA polymerases

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ABSTRACT

RNA virus survival depends on efficient viral genome replication, which is performed by the viral RNA dependent RNA polymerase (RdRp). The recent development of high throughput magnetic tweezers has enabled the simultaneous observation of dozens of viral RdRp elongation traces on kilobases long templates, and this has shown that RdRp nucleotide addition kinetics is stochastically interrupted by rare pauses of 1–1000 s duration, of which the short-lived ones (1–10 s) are the temporal signature of a low fidelity catalytic pathway. We present a simple and precise temperature controlled system for magnetic tweezers to characterize the replication kinetics temperature dependence between 25°C and 45°C of RdRps from three RNA viruses, i.e. the double-stranded RNA bacteriophage $\Phi 6$, and the positive-sense single-stranded RNA poliovirus (PV) and human rhinovirus C (HRV-C). We found that $\Phi 6$ RdRp is largely temperature insensitive, while PV and HRV-C RdRps replication kinetics are activated by temperature. Furthermore, the activation energies we measured for PV RdRp catalytic state corroborate previous estimations from ensemble pre-steady state kinetic studies, further confirming the catalytic origin of the short pauses and their link to temperature independent RdRp fidelity. This work will enable future

temperature controlled study of biomolecular complex at the single molecule level.

INTRODUCTION

Genome replication is essential to any organism. To achieve this task, every RNA virus encodes an RNA dependent RNA polymerase (RdRp) that synthesizes all the viral RNA, either to form messenger RNA for viral protein translation, or to produce new viral genomes that are enclosed into viral particles as they mature into infectious virions. Furthermore, the viral RdRp also evolves the viral genome either by incorporating mutations (1,2) while replicating the viral genome, or by assisting RNA recombination (3). The viral RdRp is a key player in successful infection, and is therefore the target of many antiviral drugs (4,5). Of special interest is the mechanochemical cycle of nucleotide incorporation occurring at the RdRp catalytic site (6). This site is conserved among RNA viruses (7), and resembles the one of the DNA and RNA polymerases in the A family with their typical cupped right hand shape (7–9). PV and $\Phi 6$ RdRps are model for the RdRps of positive-sense single-stranded (ss) and double-stranded (ds) viruses, respectively. Pre-steady-state kinetics, structural biology, molecular dynamics and next generation sequencing studies on PV RdRp and its genome have largely shaped our understanding on nucleotide selection and incorporation kinetics by viral RdRps (10), while structural and biochemical analyses on the $\Phi 6$ RdRp have provided insights into the de novo initiation mechanism (11–13). However,

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a complete description of the kinetic cycle of viral RdRp elongation is still lacking. Indeed, the ensemble techniques are unable to access RdRp kinetics on genome-long templates, i.e. kilobases (kb), and cannot interrogate rare asynchronous events, such as nucleotide misincorporation or antiviral nucleotide analogue incorporation in the presence of cognate canonical NTPs. Our recent single molecule studies on $\Phi 6$ and PV RdRps elongation kinetics have partly filled this gap, shedding light on the kinetic of elongation pauses of various biochemical origins (14–16). This work relied on the concomitant development of high throughput magnetic tweezers, a powerful single molecule force and torque spectroscopy technique enabling the characterization of protein-nucleic acids interactions at both high throughput (15,17–20) and high-resolution (21–23), and of a new analysis approach based on First-Passage statistics (24,25). However, the home-built magnetic tweezers instrument used for these studies did not include a temperature control system, and the experiments were therefore performed at room temperature. Ensemble kinetic assays have demonstrated the temperature dependence of RdRps activity, either in initiation for dengue and Zika virus RdRps (26), or in elongation for PV RdRp (27). Furthermore, temperature controlled experiments performed on *Escherichia coli* RNA polymerase (RNAP) at the single molecule level using optical tweezers have further informed on the mechanochemical cycle of nucleotide addition and on the nature of RNAP pauses, showing that off-pathway pauses have no enthalpic contribution, i.e. pause exit rate demonstrating no temperature dependence, whereas nucleotide addition rate is dominated by enthalpy, i.e. having strong temperature dependence (28,29). A temperature dependent study of viral RdRps elongation kinetics would thus significantly complement our current understanding of their mechanochemical cycle, and warrants the development of temperature controlled high throughput magnetic tweezers.

Recently, several studies have reported on the development of custom temperature controlled magnetic tweezers assays, i.e. relying either on a totally custom approach (home-built proportional-integral-derivative (PID) controller, heating elements and thermistors disposed at several locations on the microscope, and a custom graphic user interfaces (GUI)), or a dedicated commercial device (30–34). In the present study, we take advantage of a simple and robust commercial device from Thorlabs (originally designed to control the temperature on 1 inch diameter optical tubes and provided with its own GUI) to precisely control the temperature on an oil immersion microscope objective. Doing so, we are able to maintain the temperature within $\sim 0.1^\circ$ from room temperature up to 60°C in the whole field of view, i.e. with dimensions of (0.5×0.4) mm, of a home-built high throughput magnetic tweezers.

We applied this temperature controlled high throughput magnetic tweezers to study the temperature dependence of the replication kinetics of three viral RdRps, i.e. $\Phi 6$, PV and HRV-C RdRps. On the one hand, we expect a different response to temperature for $\Phi 6$ and PV RdRps, as $\Phi 6$ and PV are respectively a plant pathogenic bacteria bacteriophage and a human virus, with respective host optimal temperature of 28°C and 37°C . On the other hand, we expect a similar response to temperature for

PV and HRV-C RdRps, as both are human enteroviruses. Indeed, we show that PV and HRV-C RdRps present a steep and similar increase in nucleotide addition rate and in the exit rates for the short pauses (0.3–10 s), while the same kinetic states are lesser affected for $\Phi 6$ RdRp. Furthermore, we noticed that the long pauses (>20 s) probability is largely unaffected for all RdRp. For the three RNA viruses we studied, we show that RNA virus replication rate is optimum near the optimum growth temperature of the host cell. Moreover, we believe that the easy implementation and low-cost of the temperature control system we have characterized makes it very attractive and will therefore be of large interest to the single molecule community using instruments that include an oil immersion objective, as well as for fluorescence microscopy and optical tweezers.

MATERIALS AND METHODS

High throughput magnetic tweezers apparatus

The high throughput magnetic tweezers apparatus has been previously described (15,18,35). Shortly, it is a custom inverted microscope with a $50\times$ oil immersion objective (CFI Plan Achrom 50 XH, NA 0.9, Nikon, Germany), on top of which a flow cell is mounted. The assembly, surface preparation of the flow cell and nucleic acid tethering of the magnetic beads is described in the paragraph below. To apply an attractive force to the magnetic beads and stretch the nucleic acid tether, a pair of vertically aligned permanent magnets (5 mm cubes, SuperMagne, Switzerland) separated by a 1 mm gap (36) are positioned above the objective; the vertical position and rotation of the beads are controlled by the M-126-PD1 and C-150 motors (Physik Instrumente PI, GmbH & Co. KG, Karlsruhe, Germany), respectively (Figure 1A). The field of view is illuminated through the magnets gap by a collimated LED-light source located above it, and is imaged onto a large chip CMOS camera (Dalsa Falcon2 FA-80-12M1H, Stemmer Imaging, Germany). The temperature control system is made of a flexible resistive foil heater with an integrated $10\text{ M}\Omega$ thermistor (HT10K, Thorlabs) wrapped around an oil immersion objective (CFI Plan Achrom 50 XH, NA 0.9, Nikon, Germany) and further insulated by several layers of kapton tape (KAP22-075, Thorlabs). The heating foil is connected to a PID temperature controller (TC200 PID controller, Thorlabs) to adjust the temperature within $\sim 0.1^\circ\text{C}$. We used the Thorlabs GUI to control the heating system via USB on the data acquisition computer.

Flow cell fabrication

The flow cell assembly has been described previously (35). The flow cell was mounted on the magnetic tweezers setup and rinsed with 1 ml of $1\times$ phosphate buffered saline (PBS). $100\ \mu\text{l}$ of a 1:1000 dilution of $3\ \mu\text{m}$ polystyrene beads (LB30, Sigma Aldrich, Germany, stock concentration 1.828×10^{11} particles per milliliter) were added and after a 3-minute incubation, the flow cell was rinsed with 1 ml PBS. $40\ \mu\text{l}$ of anti-digoxigenin Fab fragments (1 mg/ml) were added and the excess rinsed away with 1 ml PBS after 30 min incubation. The flow cell was then treated

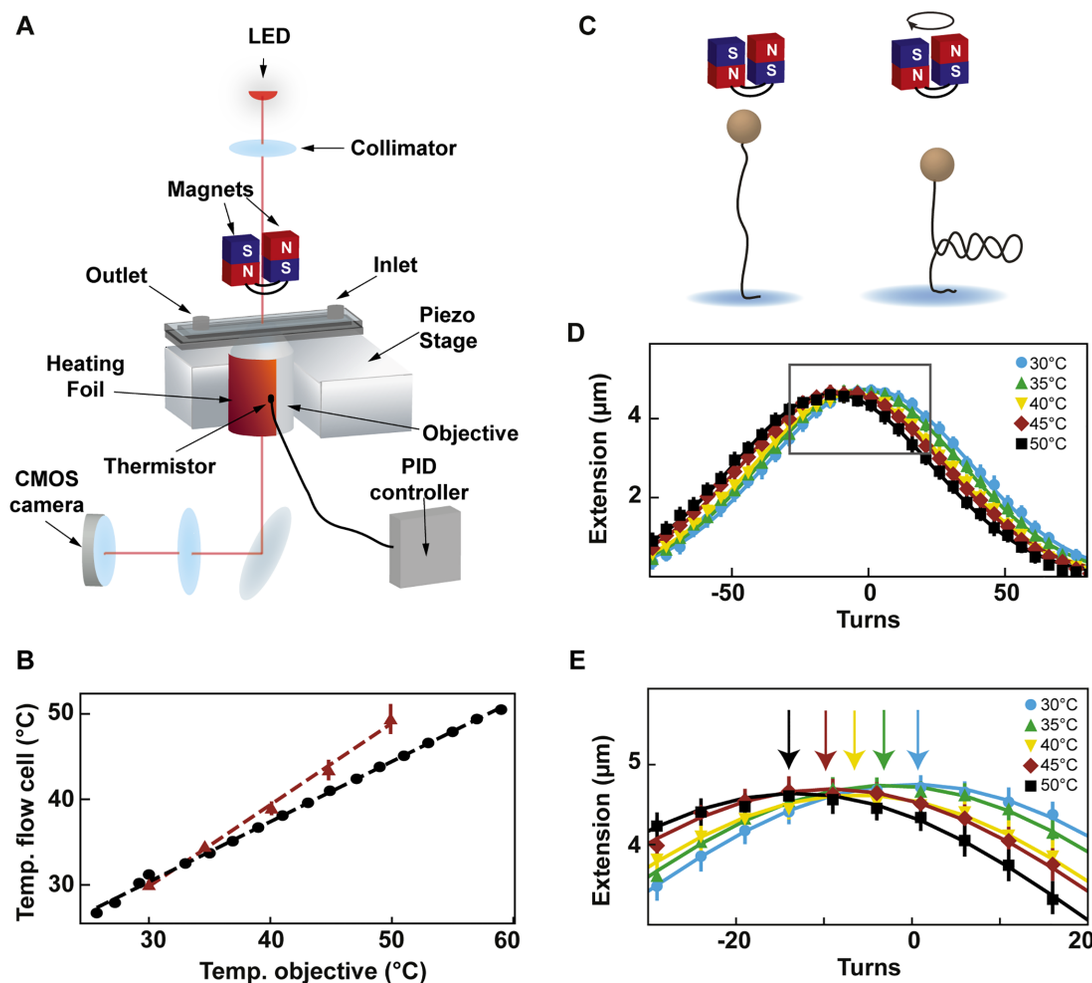


Figure 1. Description and calibration of the temperature control device installed in the high throughput magnetic tweezers assay. (A) Schematic of the magnetic tweezers assay. (B) Temperature measured at the surface of the flow cell as a function of the temperature measured at the objective by the thermistor of the heating foil. Red triangles: temperature at the surface of the flow cell extracted from the extension-rotation experiments (Material and methods; error bars are the standard deviation for seven tethers). Black circles: temperature at the surface of the flow cell measured using a macroscopic thermistor (Supplementary Figure S1B). The dashed lines represent the linear fits. (C) Schematic of supercoiling of linear dsDNA. (D) The median rotation–extension of seven different MyOne magnetic beads tethered by 20.6 kb dsDNA at either 30°C (light blue), 35°C (green), 40°C (yellow), 45°C (red) or 50°C (black), experiencing a 0.3 pN force. Each rotation–extension experiment for a given temperature is fitted by a Gaussian. (E) A magnification of the shift of the maximum extension position. Arrows indicate maxima.

with bovine serum albumin (BSA, New England Biolabs) (100 mg/ml) for 10 min and rinsed with 1 ml PBS. To remove BSA excess from the surface, high salt buffer (10 mM Tris, 1 mM EDTA pH 8.0, supplemented with 750 mM NaCl and 2 mM sodium azide) was flushed through the flow cell and rinsed 10 min later with TE buffer (TE supplemented with 150 mM NaCl and 2 mM sodium azide). To establish the temperature dependent rotation-extension experiments, we carefully mixed 10 μ l of streptavidin coated MyOne magnetic beads (ThermoFisher, Germany) and with \sim 0.25 ng of 20.6 kb long coilable double-stranded DNA tether, diluted it to 40 μ l, and flushed the solution into the flow cell (35). For the RdRp experiment, we mixed 5 μ l of dsRNA at \sim 0.1 ng/ μ l with 20 μ l of streptavidin coated M270 magnetic beads (Invitrogen), diluted it to 40 μ l and flushed the solution into the flow cell. For both experiments, the flow cell was rinsed to remove unattached magnetic beads after few minutes of incubation.

Coilable DNA construct for temperature calibration

The 20.6 kb coilable DNA construct fabrication was described in (35).

dsRNA construct

The construct employed here, which has been previously described in detail (15), a 4 kb long single-stranded splint to which four ssRNAs are annealed: a biotin-labeled strand to attach to the streptavidin-coated magnetic bead, a spacer, \sim 2.9 kb template, and a digoxigenin-labeled strand to attach the magnetic bead to the surface glass surface. The template strand ends in 3' with either a ssRNA flap made of 3 C residues followed by 15 U residues—used for Φ 6 RdRp to catalyze *de novo* initiation, or a small hairpin with the sequence ACGCUUUCGCGT followed by 15 U residues to initiate PV and HRV-C RdRp catalyzed RNA synthesis via primer extension (27,37) (Figure 2A).

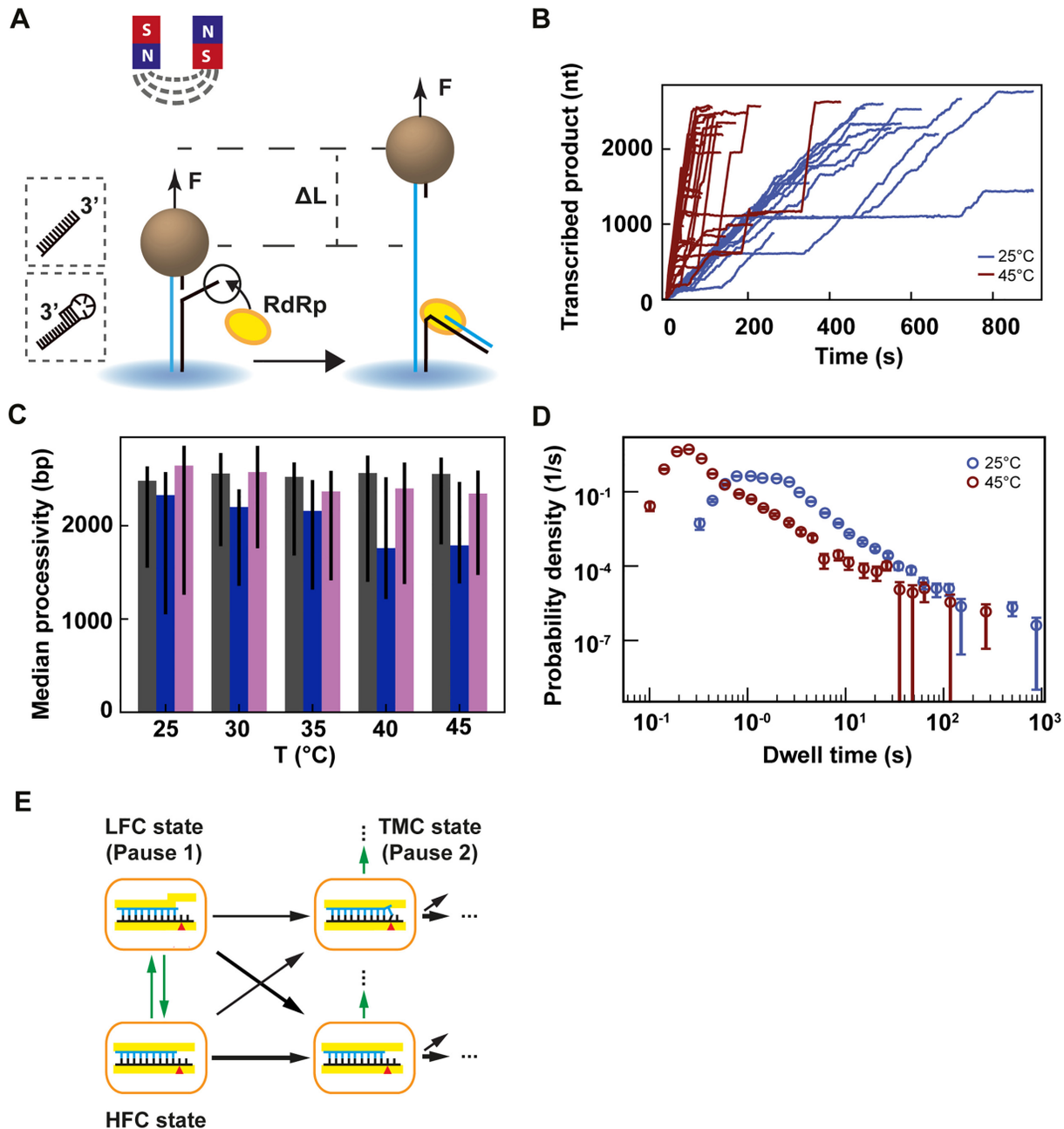


Figure 2. Temperature dependent magnetic tweezers assay show an increased replication rate for human rhinovirus C RdRp at increasing temperatures. (A) A schematic overview of the experimental assay (not to scale). A M-270 magnetic bead is tethered by a dsRNA that experiences a constant force F . The RdRp initiates at the 3'-end of the template strand, which ends either with a single-stranded 3' overhang or a short hairpin (see insert on the left). In the presence of NTPs, the RdRp replicate and unwinds the template strand, creating a ssRNA tether with a difference in extension length ΔL in comparison to the dsRNA tether at a force F . (B) Low pass filtered (0.5 Hz) HRV-C RdRp activity traces at 25°C (blue) ($N = 18$) and 45°C (red) ($N = 42$), while applying a 30 pN constant force. (C) Median processivity of $\Phi 6$ (gray), PV (blue) and HRV-C (pink) RdRps. The error bars denote the 68% confidence interval. (D) The dwell times probability density distribution extracted from at least 35 traces of HRV-C RdRp at either 25°C (blue) or 45°C (red) acquired as in (B). The error bars represent the 36% confidence interval obtained from 1000 bootstrapped datasets. (E) Kinetic model of nucleotide incorporation and mismatch by RdRps (adapted from Ref. (15)). HFC: high fidelity catalytic state; LFC: low fidelity catalytic state; TMC: terminal mismatch catalytic state.

Purification of $\Phi 6$ P2 RdRp

Recombinant N-terminally histidine-tagged $\Phi 6$ RdRp was expressed from plasmid pAA5 in *Escherichia coli* BL21 (DE3) (38) and purified using Ni-NTA affinity column (Qiagen), HiTrap™ Heparin HP column and HiTrap™ Q HP column (GE Healthcare) as previously described (39). The purified protein was stored in 50% glycerol, 140 mM NaCl, 50 mM Tris-HCl pH 8.0, 0.1 mM EDTA, 0.1% Triton-X 100 at -20°C .

Construction of HRV-C and PV RdRp bacterial expression plasmids

The HRV-C RdRp (3D gene) was cloned into the pSUMO bacterial expression plasmid using a similar procedure as described for PV RdRp (3D gene) (40). This system allows for the production of SUMO fusion proteins containing an amino-terminal hexahistidine tag fused to SUMO that can be purified by Ni-NTA chromatography and subsequently processed by the SUMO protease, Ulp1. Briefly, the HRV-C and PV RdRps coding region was amplified using respectively the HRV-C15 cDNA (Accession# GU219984.1) and the PV type 1 cDNA (Accession# V01149.1) as template, as described in (41). The PCR product of HRV-C RdRp was gel purified and cloned into the pSUMO plasmid using BsaI and Sall sites.

Expression and purification of HRV-C and PV RdRps

HRV-C RdRp was expressed and purified using the same procedure reported for PV RdRp (40,42). Briefly, expression was performed at 25°C by auto-induction, cells harvested, lysed by French Press, subjected to PEI precipitation followed by AMS04 precipitation, Ni-NTA chromatography, cleavage by Ulp1, phosphocellulose chromatography, gel filtration and the protein concentrated using Vivaspin concentrators. *Reaction buffer for $\Phi 6$ P2*. The P2 reaction buffer is composed of 50 mM HEPES pH 7.9, 20 mM ammonium acetate, 3% (w/v) polyethylene glycol 4000, 0.1 mM EDTA (pH 8.0), 5 mM MgCl_2 , 2 mM MnCl_2 , 0.01% Triton X-100, 5% Superase RNase inhibitor (Life Technologies), 20 $\mu\text{g}/\text{ml}$ BSA and 9 nM P2 RdRp.

Reaction buffer for PV and HRV-C RdRps

The PV/HRV-C RdRp reaction buffer (coined PV reaction buffer) is composed of 50 mM HEPES pH 7.9, 5 mM MgCl_2 , 0.01% Triton X-100, 5% Superase RNase inhibitor (Life Technologies). To stall PV and HRV-C RdRps on the template, 500 nM of RdRp and 1 mM of ATP, CTP and GTP in PV reaction buffer were incubated in the flow chamber for ~ 10 min, and the flow cell was subsequently rinsed.

Single-molecule RdRp replication activity experiments

We select the dsRNA tether in the flow cell in reaction buffer as previously described (15). To prevent reinitiation on the template-product duplex when PV/HRV-C RdRp falls off, we first stalled the RdRp and rinsed the flow chamber with 400 μl of reaction buffer. Subsequently, we flushed in the reaction buffer, and grabbed the data for 30 min at 58 Hz

camera acquisition rate, at a defined temperature and 30 pN force. The force is defined by the magnet distance to the flow chamber top surface, as described in (35) with a relative error of $\sim 7\%$ (standard deviation). The reaction buffer contains either 1 mM concentration of all NTPs for PV and HRV-C RdRps or 1 mM ATP/GTP and 0.2 mM CTP/UTP for $\Phi 6$ RdRp. The images are analyzed in real-time using custom-written routines in Labview 2017 and CUDA nVidia to extract the (x, y, z) position of up to ~ 800 magnetic beads simultaneously (18). The micrometric change in tether extension upon RdRp activity (Figure 2A) is converted into numbers of transcribed nucleotides using the force-extension relationships for dsRNA and ssRNA constructs (15). RdRp activity traces are low-pass filtered at 0.5 Hz using a Kaiser-Bessel window, from which the dwell time were extracted as previously described (14,15,24).

Processivity

The processivity was measured from activity traces from which the tether did not detach within 5 min after the apparent end of the trace. We represented in Figure 2C the median processivity of all the traces for a given temperature and RdRp.

Stochastic-pausing model

There are many kinetic models that are consistent with the empirical dwell-time distributions we observe, and we here work under the assumption that the probability of pausing is low enough that there is only one rate-limiting pause in each dwell-time window. This assumption washes out most details of the kinetic scheme that connects pauses and nucleotide addition, but allows us to determine the general form of the dwell-time distribution without specifying how the pauses are connected to the nucleotide addition pathway

$$p_{dw}(t) \propto p_{na} \Gamma\left(t; N_{dw}, \frac{1}{k_{na}}\right) + Q(t) \left(\sum_{n=1}^{N_{sp}} p_n k_n e^{-k_n t} + \frac{a_{bt}}{2(1+t/1s)^{3/2}} \right). \quad (1)$$

In the above expression, the gamma function in the first term contributes the portion p_{na} of dwell-times that originate in the RdRp crossing the dwell-time window of size N_{dw} base pairs without pausing; the second term is a sum of contributions originating in pause-dominated transitions, each contributing a fraction p_n of dwell-times; the third term captures the asymptotic power-law decay (amplitude a_{bt}) of the probability of dwell-times dominated by a backtrack. The backtracked asymptotic term needs to be regularized for times shorter than the diffusive backtrack step. We have introduced a regularization at 1 s, but the precise timescale does not matter, as long as it is set within the region where the exponential pauses dominate over the backtrack. From left to right, each term of (Equation 1) is dominating the distribution for successively longer dwell-times.

A cut-off factor $Q(t)$ for short times is introduced to account for the fact that the dwell-time window includes N_{dw}

nucleotide-addition steps,

$$Q(t) = \frac{(tk_{na}/N_{dw})^{N_{dw}-1}}{1 + (tk_{na}/N_{dw})^{N_{dw}-1}}.$$

The fit results dependence on these cut-offs is negligible as long as they are introduced in regions where the corresponding term is sub-dominant. Here the cut is placed under the center of the elongation peak, guaranteeing that it is placed where pausing is sub-dominant.

Maximum likelihood estimation

The normalized version of (Equation 1) is the dwell-time distribution fit to the experimentally collected dwell-times t_i by minimizing the likelihood function (43):

$$L = - \sum_i \ln p_{dw}(t_i)$$

with respect to rates and probabilistic weights. We calculate the errors in our parameter estimates by bootstrapping (44) the system 100 times, and report the one-sigma confidence intervals among the bootstrapped data sets.

Dominating in a dwell-time window versus dominating in one step

The fractions p_n represent the probability that a particular rate k_n dominates the dwell-time. We want to relate this to the probability P_n that a specific exit rate dominates within a 1-nt transcription window. Assuming we have labelled the pauses so that $k_{n-1} > k_n$, we can relate the probability of having rate n dominating in N_{dw} steps to the probability of having it dominate in one step through

$$p_n = \left(\sum_{m=0}^n P_m \right)^{N_{dw}} - \left(\sum_{m=0}^{n-1} P_m \right)^{N_{dw}},$$

$$p_0 = p_{na} = P_{na}^{N_{dw}} = P_0^{N_{dw}} \quad (2)$$

The first term in (Equation 2) represents the probability of having no pauses longer than the n th pause in the dwell-time window, and the second term represents the probability of having no pauses longer than the $(n-1)$ th pause. The difference between the two terms is the probability that the n th pause will dominate. This can be inverted to yield a relation between the single-step probabilities (P_n) and the dwell-time window probabilities (p_n)

$$P_n = \left(\sum_{m=0}^n p_m \right)^{1/N_{dw}} - \left(\sum_{m=0}^{n-1} p_m \right)^{1/N_{dw}},$$

$$P_0 = p_0^{1/N_{dw}}.$$

This relationship has been used throughout the manuscript to relate our fits over a dwell-time window to the single-step probabilities.

RESULTS

Establishing temperature control of high throughput magnetic tweezers

To perform a temperature-controlled experiment in a magnetic tweezers assay, one needs to maintain the temperature setpoint constant in the field of view. To fulfill this condition using a magnetic tweezers instrument, it is sufficient to only control the temperature at the objective (given that an oil immersion objective is used). Indeed, the field of view is separated by at least one centimeter from the base plate (Supplementary Figure S1A) and as glass is a poor thermal conductor, the field of view remains insulated from heat transferred from the base plate (30). One may suggest that not maintaining the temperature on the top coverslip of the flow chamber may result in convection flows that would perturb the measurement. The Rayleigh number (Ra) compares convection over conduction (45). Natural convection occurs for $Ra \sim 2000$, which is much larger than our estimation of Ra in our experimental conditions (for a flow cell inner thickness ~ 0.23 mm), i.e. $Ra \sim 4$, and therefore convection does not occur in our conditions. We have mounted a temperature control device derived from Thorlabs parts on the objective, meaning a flexible resistive foil heater including a thermistor that is strapped around the oil immersion objective (Supplementary Figure S1B) and a PID controller to adjust the temperature setpoint (Figure 1A, Materials and Methods). This system has several advantages in comparison with custom ones: it is commercially available, low price, and it comes with a graphic user interface (GUI) to adjust the temperature from the data acquisition computer. We first calibrated the temperature measured on the objective side versus the temperature measured in the flow cell. To this end, we used a digital portable thermometer (Gochange, Amazon.de, $\pm 1.5\%$ error), whose ~ 1 mm³ temperature probe was inserted in the flow cell and immersed in ultrapure water (Supplementary Figure S1A). In absence of heating, i.e. at room temperature, the two sensors reported the same temperature. We repeated this measurement several times at different temperature setpoints assigned on the GUI to the PID controller, from 25°C to 60°C, i.e. the maximum temperature tolerated by the objective. Between each measurement, we waited at least 30 min for the objective temperature to equilibrate, as the focal plane of the objective shifts until equilibrium is reached (Supplementary Figure S1C). We observed a linear relationship between the temperature at the objective and the temperature in the flow chamber, i.e. $T_{\text{flow cell}}(T_{\text{obj}}) = (0.7 * T_{\text{obj}} + 9.23) \text{ } ^\circ\text{C}$ (Figure 1B, Supplementary Table S1). Because the probe integrates the temperature over a cubic millimeter volume, the thermometer measurement may not be representative of the temperature within few microns above the bottom coverslip surface of the flow cell. To perform an in situ calibration directly at the surface, we took advantage of the well-characterized temperature dependence of the DNA helical twist, which decreases by $\sim 11^\circ/\text{C}/\text{kb}$ (46–48). Magnetic tweezers are an ideal technique to measure the twist of a coilable molecule, such as fully double-stranded DNA, as they enable a precise control of the torque applied on such molecule (Figure 1C). We performed several DNA

rotation-extension experiments at 0.3 pN and at different temperatures using a coilable DNA molecule (30,49) (Figure 1D, Materials and Methods). After waiting at least 30 min for the objective temperature to equilibrate to the set-point value (Supplementary Figure S1C) (30), we evaluated the spatiotemporal resolution of the set up. To this end, we monitored the z -axis position of a surface attached 3 μm diameter bead, to which another reference bead position has been subtracted, at either 25°C (Supplementary Figure S1D) or 35°C (Supplementary Figure S1E). The trace at each temperature shows little drift, which is further supported by their respective Allan deviation (35,50) (Supplementary Figure S1F). The extension of DNA molecules at different supercoiling density shows a typical bell-like shape distribution, with the maximum extension centered at zero turn (Figure 1D). This shape is well fitted by a Gaussian function, which enables the evaluation of the zero turn maximum extension (Figure 1E). If DNA helical twist decreases, the zero turn position shifts towards the negative turns. If DNA helical twist increases, the zero turn position now shifts towards the positive turns. Changing the temperature by steps of 5°C from 30°C to 50°C, and centering at zero turn the maximum extension obtained at 30°C, we observed that the rotation-extension traces shift towards the negative turns upon temperature increases, as expected from a reduction in DNA helical twist (Figure 1D, E). Converting the shift of the peak of the traces into temperature variation using the above described relationship between DNA helical twist and temperature, we extracted almost a one-to-one linear relationship between the temperature measured at the surface of the flow chamber and on the side of the objective, i.e. $T_{\text{flow cell}}(T_{\text{obj}}) = (0.95 * T_{\text{obj}} + 1.18) \text{ } ^\circ\text{C}$ (Figure 1B, Supplementary Table S1). If not indicated otherwise, we will use this equation for temperature conversion in the subsequent sections of the study. As our magnetic tweezers assay has a very large field of view of $\sim(0.5 \times 0.4) \text{ mm}$, we evaluated the homogeneity of the temperature over the whole field of view. To this end, we performed a rotation extension experiment at both at 25°C and 45°C with the same DNA tether at five different locations in the field of view by moving the flow cell with the micrometric screws of the stage, and we did not measure any difference of the rotation-extension traces for a given temperature. This result confirms that the temperature is constant over the whole field of view (Supplementary Figure S1E) (32). We have now shown that using a simple and commercially available temperature control system, we are able to adjust and maintain the temperature in the entire field of view of a high throughput magnetic tweezers instrument. This assay is thus suitable to investigate the temperature dependence of viral RdRps elongation kinetics.

Setting up high throughput magnetic tweezers to study viral replication

We have recently developed a high throughput magnetic tweezers assay to investigate viral RdRp replication kinetics (14–16). As RdRps use an RNA template, we made a non-coilable dsRNA construct to tether the magnetic beads to the glass coverslip surface of the flow cell as previously described (15) (Figure 2A, Materials and Methods). The 3'-end of the template RNA is terminated by either a blunt

ssRNA to initiate $\Phi 6$ RdRp (51), or by a small hairpin to initiate PV or HRV-C RdRps (14,37) (Figure 2A). Following successful initiation, the RdRp displaces the template strand from the tethering strand, converting the tether from dsRNA to ssRNA, which increases linearly the end-to-end extension of the tether by ΔL at constant force F and is subsequently converted from microns to nucleotides (15) (Figure 2A). Of note, the force is not directly applied to the RdRp, but on the tethering RNA strand, which lowers the energy barrier to the RdRp forward translocation formed by the downstream double-stranded RNA fork. Using high throughput magnetic tweezers, dozens of RdRp activity traces are simultaneously acquired (Figure 2B, Supplementary Figure S2A, C). In this work, we study the influence of temperature on the replication kinetics of three model RdRps. The direct observation of HRV-C RdRp traces show that an increase in temperature from 25°C to 45°C dramatically increases its average replication rate (Figure 2B). This is also observable for PV RdRp, and, to a smaller extent, for $\Phi 6$ RdRp (Supplementary Figure S2A, C). Looking at the processivity of the three RdRps for the probed temperature range, we evaluated a constant processivity for $\Phi 6$ and HRV-C RdRps and a slight decay for PV RdRp above 40°C (Figure 2C). To further quantitate the change in replication kinetics, we performed a dwell time analysis of the 0.5 Hz low pass filtered traces. Specifically, the traces were scanned with non-overlapping windows of 10 nt to evaluate the duration of ten successive nucleotide incorporations, after which the collected dwell times are assembled into a probability density distribution (Figure 2D, Supplementary Figures S2B, D, S3) (14,15,24). The dwell time analysis is a direct measure of the event(s) that dominated in time over ten successive nucleotide incorporations, i.e. either the incorporation of the ten nucleotides only or a pause that interrupted the ten nucleotides addition (Supplementary Figure S1F). Looking at the dwell times distribution extracted from the HRV-C traces acquired at 45°C (Figure 2B) represented in a log-binned histogram, the bins describe a bell-like shape until $\sim 0.4 \text{ s}$, then a shoulder from $\sim 0.4 \text{ s}$ till $\sim 3 \text{ s}$, and finally a straight line for the longer dwell times (Figure 2D, Supplementary Figure S1F). Each region of the distribution is the temporal signature of a specific biochemical event in elongating RdRps. To describe these events, we previously introduced four probability distribution functions: to fit the bell-like shape region, a gamma distribution that describes ten successive nucleotide additions cycles without pause; to fit the shoulders, two exponentially distributed pauses (Pause 1 and Pause 2); and finally to fit the dwell times dominated by the longest pauses, a power law distribution (15,16,52) (Supplementary Figure S1F, Materials and Methods). We have previously showed (14,15) that RdRp incorporates cognate NTPs fast through the high fidelity catalytic (HFC) state (Figure 2E, gamma distribution), and rarely interconverts into another conformation, i.e. the low fidelity catalytic (LFC) state (Figure 2E), from which the RdRp slowly incorporates either a cognate NTP appearing as Pause 1 (diagonal arrow in Figure 2E) or a non-cognate NTP. In the latter, the nucleotide addition following the mismatch addition is done even slower than Pause 1 through a state we coined the terminal mismatch catalytic (TMC) state (Figure 2E) and appears in

the traces as Pause 2. Furthermore, we also showed that the power law distributed pause relates to RdRp backtracking (16,52). Using a maximum likelihood estimation (MLE) procedure, we extracted the parameters for each distribution, i.e. exit rates and probabilities (Material and Methods, Supplementary Table S2, Figure S3) (24). Having acquired and analyzed viral replication traces at several temperatures, we now describe how the parameters extracted from the MLE procedure vary for $\Phi 6$, PV and HRV-C RdRps, at constant force, i.e. 30 pN, and saturating NTPs concentration, i.e. 1 mM.

The nucleotide addition rates of RdRps from an environmental phage and from two human viruses show different responses to temperature increase

As noted above, HRV-C and PV RdRps show a dramatic increase in average elongation rate when increasing the temperature (Figure 2B, D, Supplementary Figure S2CD), while only a mild effect is observable for $\Phi 6$ RdRp (Supplementary Figure S2A, B). We first look into the pause-free nucleotide addition rate described by the bell-like shape gamma distribution (Figure 3A, Supplementary Figures S1F, S3). For temperature varying from 25°C to 45°C, the nucleotide addition rate of PV RdRp increases from (17.8 \pm 0.3) nt/s to (52.7 \pm 0.5) nt/s, i.e. almost a 3-fold increase, and in agreement with previous bulk measurements (27). Similarly, HRV-C RdRp nucleotide addition rate strongly increases, i.e. from (9.6 \pm 0.1) nt/s to (42.3 \pm 0.3) nt/s, i.e. a 4-fold increase. On the other hand, $\Phi 6$ RdRp nucleotide addition rate only varies from (16.0 \pm 0.1) nt/s to (24.6 \pm 0.2) nt/s for the same temperature span, in line with a previous ensemble measurement (53) (Figure 3A, Supplementary Table S2). Investigating the temperature response of a reaction, one is able to determine whether the reaction is exothermic or endothermic from the Van't Hoff equation by determining whether the sign of the reaction rate derivative by the temperature is negative or positive, respectively. Furthermore, it enables the measurement of the activation energy E_A of the reaction by fitting the reaction rate k , i.e. nucleotide addition or pause exit, as a function of the temperature with the Arrhenius equation:

$$k(T) = Pe^{-\frac{E_A}{RT}} \quad (3)$$

where P is the preexponential factor, T is the absolute temperature, and R is the universal gas constant (54) (Figure 3B). As the nucleotide addition rate increases with temperature, the reaction is endothermic. Evaluating the activation energy (\pm standard deviation) of the nucleotide addition rate for the three RdRps using (Equation 3), we found $E_{A,\Phi 6} = (4.1 \pm 0.4)$ kcal/mol, $E_{A,PV} = (10.7 \pm 1.0)$ kcal/mol and $E_{A,HRV-C} = (14.3 \pm 1.2)$ kcal/mol ((Equation 1)). $\Phi 6$ and PV RdRp nucleotide addition rates are not affected by increasing the mechanical tension on the dsRNA construct (14,15). Therefore, the nucleotide addition rate is not limited by the rate of the polymerase translocating to the $n + 1$ position, and the activation energy measured here relates to another rate limiting process, such as e.g. the phosphoryl transfer (6). Conclusively, our results confirm a previous pre-steady state kinetic estimation of the free

energy landscape of nucleotide addition for PV RdRp (6). Our results demonstrate the large impact of temperature on the nucleotide addition rate for human viruses, i.e. PV and HRV-C RdRp, and the mild impact on the environmental virus, i.e. the RdRp of *Pseudomonas* bacteriophage $\Phi 6$.

Human virus and environmental bacteriophage RdRps elongation competent pauses are differently affected by temperature

We then investigated how Pause 1 and Pause 2 kinetics and probabilities (Figure 4A and E) are affected by temperature. We previously showed that Pause 1 and Pause 2 are the signature of slow nucleotide addition cycles that appear as short pauses in the RdRp elongation activity traces (14,15). For PV and HRV-C RdRps, Pause 1 exit rate increases dramatically, i.e. by 3- and 6-fold, respectively (Supplementary Table S2, Figure 4BF), which is further highlighted by their respective large Pause 1 activation energy, i.e. $E_{A,PV} = (11.0 \pm 1.1)$ kcal/mol and $E_{A,HRV-C} = (15.2 \pm 4.9)$ kcal/mol (Supplementary Table S2, Figure 4B, C), similar to the nucleotide addition activation energy (Figure 3C). Pause 2 exit rate also experiences a steep increase, i.e. by 4- and 10-fold for PV and HRV-C RdRps, respectively, and eventually saturates at 35°C (Supplementary Table S2, Figure 4F). We decided to fit only the values of Pause 2 exit rate below 35°C for PV and HRV-C RdRps to fit Equation (3), as this equation does not describe the plateau observed at higher temperatures. Of note, to achieve reasonable fits to the HRV-C dwell time distribution collected at 40°C, we were forced to restrict Pause 2 exit rate $k_2 > 0.2$ s⁻¹ (Supplementary Figure S3). The large increase of Pause 2 exit rates is further supported by an activation energy larger than for Pause 1, i.e. $E_{A,PV} = (23.2 \pm 0.5)$ kcal/mol and $E_{A,HRV-C} = (34.3 \pm 1.8)$ kcal/mol (Equation 3, Figure 4G). Using the temperature calibration from the macroscopic thermistor (Figure 1B), we observed a significant variation in activation energies for PV RdRp (Supplementary Figure S4), which further supports the utilization of a calibration in the immediate vicinity of the surface, such as DNA rotation extension experiments (Figure 1B). Interestingly, Pause 1 and Pause 2 exit rates for PV and $\Phi 6$ RdRps increase with assisting tension (14,15), which suggests that forward translocation is the rate limiting step of slow nucleotide addition through this parallel kinetic pathway (Figure 2E). The large activation energy of Pause 2 is therefore consistent with the barrier to RdRp translocation induced by a nucleotide mismatch and is in agreement with previous ensemble measurements (6). The saturation of Pause 2 exit rates $> 35^\circ\text{C}$ suggests that Pause 2 is made of (at least) two successive kinetics states, one being temperature dependent and rate limiting up to 35°C, e.g. the translocation over a nucleotide mismatch, and a second one being not temperature dependent and rate limiting above 35°C, which is not determined here. Looking now at $\Phi 6$ RdRp, we observed a very different trend. Indeed, Pause 1 and Pause 2 exit rates are relatively stable, i.e. $k_1 = (1.7 \pm 0.4)$ s⁻¹ and $k_2 = (0.26 \pm 0.09)$ s⁻¹ respectively (mean \pm standard deviation, Supplementary Table S2, Figure 4BF). Therefore, the Arrhenius equation fits poorly to $\Phi 6$ RdRp pause exit rates

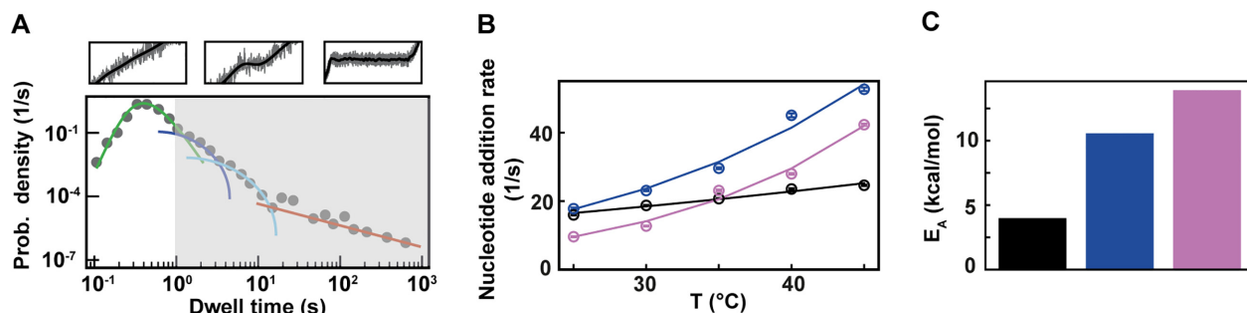


Figure 3. $\Phi 6$, PV and HRV-C RdRps show different nucleotide addition rates response to temperature increase. The corresponding data for each RdRp is represented in black, blue and pink, respectively in (B) and (C). (A) Section (not shaded) of the dwell time distribution that contributes to the MLE fit (green solid line) of the nucleotide addition (Supplementary Figure S1D). (B) The nucleotide addition rates (circles) as obtained from the maximum likelihood estimation (MLE) fits. The solid lines represent the respective fit of the Arrhenius equation. The error bars denote the standard deviation from 100 bootstraps of the MLE procedure (Materials and Methods). (C) Activation energy E_A extracted from the fits in (B).

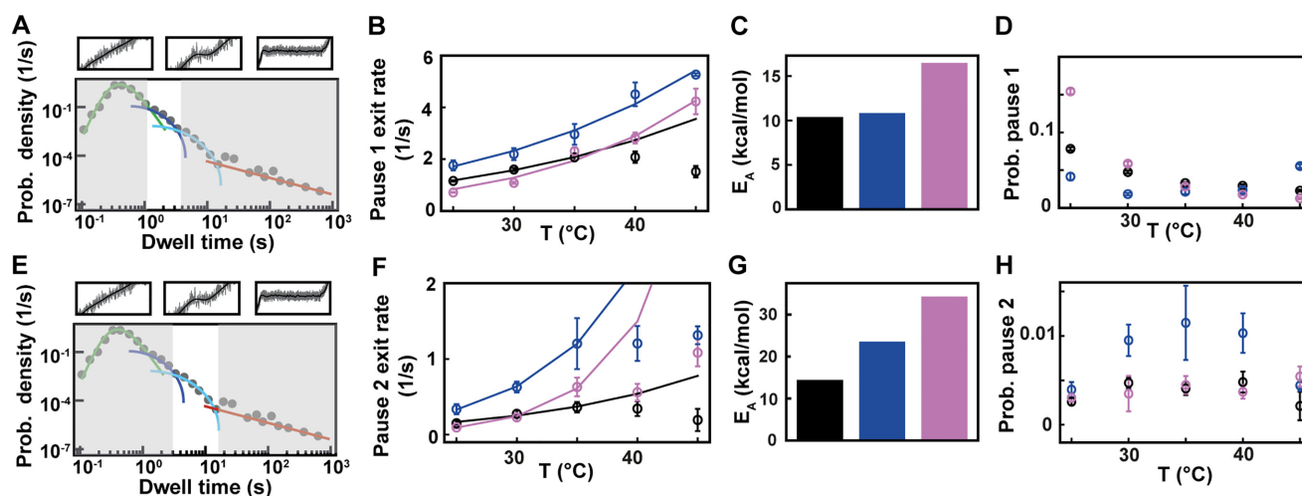


Figure 4. Pause 1 and Pause 2 exit rates and probabilities of $\Phi 6$, PV and HRV-C RdRps demonstrate different activation by temperature. The data for each RdRp is represented in black, blue and pink, respectively. (A) Section (not shaded) of the dwell time distribution that contributes to the MLE fit (dark blue solid line) of Pause 1. (B) The pause 1 exit rates (circles) extracted from the MLE fits as a function of the temperature. The lines represent the fitted Arrhenius equation. For $\Phi 6$ RdRp, only the data at 25°C, 30°C and 35°C were considered for the fit. (C) Activation energy for Pause 1 exit rate. (D) The probabilities to be in Pause 1 state as a function of the temperature. (E) Section (not shaded) of the dwell time distribution that contributes to the MLE fit (light blue solid line) of Pause 2. (F) The Pause 2 exit rates (circles) extracted from MLE fits. The lines represent the fitted Arrhenius equation. Only the data at 25°C, 30°C and 35°C were considered for the fit. (G) Activation energy for Pause 2 exit rate. (H) The probabilities to be in Pause 2 state as a function of the temperature. Error bars in B, D F and H denote the standard deviation extracted from 100 bootstraps of the MLE procedure.

evolution with temperature (Figure 4BF), leading to a poor estimation of the activation energies of Pause 1 and Pause 2 (Figure 4CG, Supplementary Table S2). Off-pathway pause exit rates have been shown to be temperature insensitive in *E. coli* RNAP (28,29), and could offer an attractive explanation for the relative insensitivity of Pause 1 and Pause 2 from $\Phi 6$ RdRp. However, $\Phi 6$ RdRp elongation rate also increases little with temperature, and therefore we suspect that the thermal activation of $\Phi 6$ RdRp elongation kinetics occurs at lower temperature than the range we explored. Pause 1 probabilities are largely unaffected by temperature for $\Phi 6$ and PV RdRps, while it decreases when temperature increases for HRV-C RdRp (Figure 4D). However, the nucleotide addition rate and Pause 1 regions of the HRV-C RdRp dwell time distributions have a strong overlap at low temperature (Figure 3B, Supplementary Figure S3), which may bias the probability we measured. Pause 2 probability

is largely constant for all RdRps (Figure 4H). As for nucleotide addition rate, we observe a very different response to temperature for the human RNA virus RdRps and the environmental dsRNA bacteriophage, which could be an evolutionary advantage for a bacteriophage to not respond too strongly on natural environmental temperature variation in the room temperature range.

Backtrack related long pause probability mildly decreases with temperature increase for $\Phi 6$ RdRp, while remaining constant for PV and HRV-C RdRps

We next look at the last part of the dwell time distribution, which describes the pauses longer larger than 20 s (Figure 5A) and are related to polymerase backtrack, i.e. a backward diffusion of the polymerase on its template leading to an RNA product 3'-end out of register (Supplementary

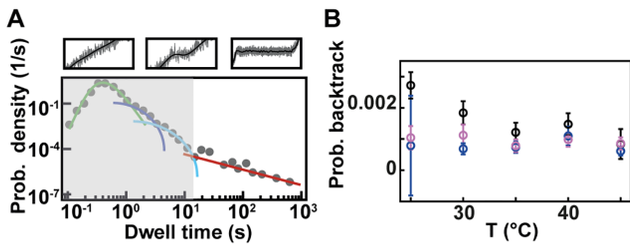


Figure 5. Temperature has little effect on backtrack probabilities for $\Phi 6$, PV and HRV-C RdRps. (A) Section (not shaded) of the dwell time distribution that contributes to the fit (red solid line) of the backtrack pause. (B) The probabilities for backtrack (circles) as obtained from MLE fits. The error bars denote the standard deviation extracted from 100 bootstraps of the MLE procedure. The data for $\Phi 6$, PV and HRV-C RdRps is represented in black, blue and pink, respectively.

Figure S5). The polymerase eventually returns in register by hopping from base to base with a forward and backward rate. This is modeled by a one dimensional diffusion process through a periodical energy landscape, and is described in the dwell time distribution by a power law distribution with two parameters, i.e. a $-3/2$ exponent and a characteristic hopping rate (28,52,55) (Materials and Methods). While we have previously observed $\Phi 6$ RdRp backtracking (16), this behavior had remained elusive for PV RdRp in absence of the nucleotide analogue T-1106 (14). We suspected that the absence of long backtrack pauses in our previous study likely originated from an RNase contamination of the PV RdRp stock, which consequently decreased the tether lifetime and prevented the observation of these rare and long pauses. Having improved PV and HRV-C RdRps purification protocols (Material and Methods), we are now able to show exemplary traces of deep backtracks (Supplementary Figure S5) (shallow ones cannot be resolved due to the limited spatiotemporal resolution of the assay) and characterize the backtrack-related pauses for these two enterovirus RdRps (Figure 5, Supplementary Figure S3). The average hopping rate is hidden in the dwell time distribution behind the shoulder of Pause 1 and Pause 2, as they statistically dominate, and we have therefore decided to fix this parameter at 1 s^{-1} , a value that has been directly measured for *E. coli* RNAP (52,56). Using this fixed hopping rate, we evaluated the probability for the polymerase to enter a backtrack pause as a function of the temperature. For temperatures varying from 25°C to 45°C , $\Phi 6$ RdRp probability to enter a backtrack pause mildly decreases, i.e. from 0.0027 to 0.0008, while for PV and HRV-C RdRps, this probability is largely constant, i.e. (0.0008 ± 0.0002) and (0.0009 ± 0.0002) (mean \pm standard deviation), respectively (Figure 5B, Supplementary Table S2). Though we should be cautious with the extracted probabilities as we do not know the average hopping rate of a backtracking RdRp, we observe a different backtracking behavior between an environmental bacteriophage RdRp and human virus RdRps.

DISCUSSION

Single molecule biophysics has revolutionized our view of molecular biology by enabling the observation of the kinetic

processes of single enzymes, increasing significantly our understanding of these complex molecular motors (57). In particular, single molecule biophysics has further complemented the existing wealth of knowledge collected in ensemble measurements with transient, rare and asynchronous kinetic events to assemble complete mechanochemical pathways. However, while bulk studies are easily performed at physiological temperature, i.e. 37°C for human and *E. coli* enzymes, many single molecule studies are performed at room temperature ($\sim 21^{\circ}\text{C}$) because of the technical difficulties of inserting a temperature control system within home-built high-end microscopes. Here, we present a simple temperature control system with several advantages: low cost, commercially available (Thorlabs), and easily mounted on an oil immersion objective. We calibrated the temperature in the field of view of a home-built magnetic tweezers instrument by performing rotation-extension experiments on a supercoiled DNA at several temperatures, using the well-known relationship describing the decrease in DNA helical twist as a function of temperature and achieving better performance than using a macroscopic thermometer probe.

We have previously investigated the kinetics of PV RdRp at room temperature, extracting a slow nucleotide addition rate of ~ 10 nt/s, even at saturating concentration of NTPs and high assisting force (14). This result was consistent with previous bulk measurements performed at the same temperature on the same enzyme and using ssRNA templates (27). Having established a temperature controlled magnetic tweezers assay, we have presented in this study how temperature affects the replication kinetics of RdRps from several different viruses, i.e. $\Phi 6$ (a bacteriophage infecting a plant pathogenic bacteria), PV and HRV-C (two human enteroviruses). These RdRps responded very differently to temperature change. $\Phi 6$ RdRp is only mildly affected in the temperature range we probed here, which is in line with previous work (53). *Pseudomonas syringae*, $\Phi 6$ natural host, grows at wide range of environmental temperatures (from 0 up to 36°C) with an wide optimum around $\sim 28^{\circ}\text{C}$ (58). However, the reproduction cycle of $\Phi 6$ is restricted at temperatures above 30°C (59). Consequently, we suggest that $\Phi 6$ RdRp has evolved to support viral genome replication under different environmental temperatures, and therefore is not thermally activated $>25^{\circ}\text{C}$. On the other hand, the kinetics of PV and HRV-C RdRps are strongly affected by the increase in temperature. The data we present here definitely supports that the temperature is the predominant factor behind the slow replication we previously observed, as we have now measured nucleotide addition rates similar to the one measured in bulk at the same temperatures, using ssRNA templates (27). PV and HRV-C RdRps showed a parallel trend, with PV nucleotide addition rate saturating at a lower temperature. Interestingly, the activation energy we measured for the nucleotide addition rate and Pause 1, and the one for Pause 2 are both consistent with the energy barrier for cognate and non-cognate nucleotide addition, respectively, estimated by pre-steady state kinetic analysis (6). This result further supports our model where Pause 1 is a slow nucleotide addition event and Pause 2 is the signature of mismatch nucleotide addition (14,15) (Figure 2E), and suggests that temperature does not affect nucleotide misincorporation rate (Pause 2 probability is largely constant).

Finally, we also show here that the replication traces of non-de novo initiating RdRps, i.e. PV and HRV-C, present long pauses related to polymerase backtrack, similarly to *de novo* initiating $\Phi 6$ RdRp (Figure 5B) (16). We anticipate that this behavior is ubiquitous in all RNA virus RdRps and may be an important feature for viral replication. On a technical note, we show here that our analysis framework is particularly suitable to detect even very short lived pauses, i.e. <1 s, while standard pause-removal approaches can only efficiently detect pauses longer than 1 s (60). Therefore, we are confident that our approach enables a good and less biased estimation of the nucleotide addition rate.

In conclusion, we have established a simple and easy to set up objective-based temperature control system for magnetic tweezers. This approach is easily applicable to any microscope using an immersion objective to ensure a good heat transfer. We believe that the simplicity and availability of our assay makes it very attractive, and will therefore be widely applied in the field. In addition, the RdRps replication kinetics study brings new evidences in support of our model of nucleotide mismatch addition.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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REFERENCES

- Lauring, A.S., Frydman, J. and Andino, R. (2013) The role of mutational robustness in RNA virus evolution. *Nat. Rev. Microbiol.*, **11**, 327–336.
- Peck, K.M. and Lauring, A.S. (2018) Complexities of viral mutation rates. *J. Virol.*, **92**, e01031–17.
- Simon-Loriere, E. and Holmes, E.C. (2011) Why do RNA viruses recombine? *Nat. Rev. Microbiol.*, **9**, 617–626.
- Bekerman, E. and Einav, S. (2015) Infectious disease. Combating emerging viral threats. *Science (New York)*, **348**, 282–283.
- Crotty, S., Maag, D., Arnold, J.J., Zhong, W., Lau, J.Y., Hong, Z., Andino, R. and Cameron, C.E. (2000) The broad-spectrum antiviral ribonucleoside ribavirin is an RNA virus mutagen. *Nat. Med.*, **6**, 1375–1379.
- Arnold, J.J. and Cameron, C.E. (2004) Poliovirus RNA-dependent RNA polymerase (3DPOL): pre-steady-state kinetic analysis of ribonucleotide incorporation in the presence of Mg^{2+} . *Biochemistry*, **43**, 5126–5137.
- te Velthuis, A.J. (2014) Common and unique features of viral RNA-dependent polymerases. *Cell. Mol. Life Sci. CMLS*, **71**, 4403–4420.
- Monttinen, H.A., Ravantti, J.J., Stuart, D.I. and Poranen, M.M. (2014) Automated structural comparisons clarify the phylogeny of the right-hand-shaped polymerases. *Mol. Biol. Evol.*, **31**, 2741–2752.
- Steitz, T.A. (1999) DNA polymerases: structural diversity and common mechanisms. *J. Biol. Chem.*, **274**, 17395–17398.
- Cameron, C.E., Moustafa, I.M. and Arnold, J.J. (2016) Fidelity of nucleotide incorporation by the RNA-dependent RNA polymerase from poliovirus. *Enzymes*, **39**, 293–323.
- Wright, S., Poranen, M.M., Bamford, D.H., Stuart, D.I. and Grimes, J.M. (2012) Noncatalytic ions direct the RNA-dependent RNA polymerase of bacterial double-stranded RNA virus varphi6 from *de novo* initiation to elongation. *J. Virol.*, **86**, 2837–2849.
- Butcher, S.J., Grimes, J.M., Makeyev, E.V., Bamford, D.H. and Stuart, D.I. (2001) A mechanism for initiating RNA-dependent RNA polymerization. *Nature*, **410**, 235–240.
- Ren, Z. and Ghose, R. (2011) Slow conformational dynamics in the cystoviral RNA-directed RNA polymerase P2: influence of substrate nucleotides and template RNA. *Biochemistry*, **50**, 1875–1884.
- Dulin, D., Arnold, J.J., van Laar, T., Oh, H.S., Lee, C., Perkins, A.L., Harki, D.A., Depken, M., Cameron, C.E. and Dekker, N.H. (2017) Signatures of nucleotide analog incorporation by an RNA-dependent RNA polymerase revealed using high-throughput magnetic tweezers. *Cell Rep.*, **21**, 1063–1076.
- Dulin, D., Vilfan, I.D., Berghuis, B.A., Hage, S., Bamford, D.H., Poranen, M.M., Depken, M. and Dekker, N.H. (2015) Elongation-competent pauses govern the fidelity of a viral RNA-dependent RNA polymerase. *Cell Rep.*, **10**, 983–992.
- Dulin, D., Vilfan, I.D., Berghuis, B.A., Poranen, M.M., Depken, M. and Dekker, N.H. (2015) Backtracking behavior in viral RNA-dependent RNA polymerase provides the basis for a second initiation site. *Nucleic Acids Res.*, **43**, 10421–10429.
- Berghuis, B.A., Dulin, D., Xu, Z.Q., van Laar, T., Cross, B., Janissen, R., Jergic, S., Dixon, N.E., Depken, M. and Dekker, N.H. (2015) Strand separation establishes a sustained lock at the Tus-Ter replication fork barrier. *Nat. Chem. Biol.*, **11**, 579–585.
- Cnossen, J.P., Dulin, D. and Dekker, N.H. (2014) An optimized software framework for real-time, high-throughput tracking of spherical beads. *Rev. Sci. Instrum.*, **85**, 103712.
- Ribeck, N. and Saleh, O.A. (2008) Multiplexed single-molecule measurements with magnetic tweezers. *Rev. Sci. Instrum.*, **79**, 094301.
- De Vlamincq, I., Henighan, T., van Loenhout, M.T., Pfeiffer, I., Huijts, J., Kerssemakers, J.W., Katan, A.J., van Langen-Suurling, A., van der Drift, E., Wyman, C. *et al.* (2011) Highly parallel magnetic tweezers by targeted DNA tethering. *Nano Lett.*, **11**, 5489–5493.
- Dulin, D., Cui, T.J., Cnossen, J., Docter, M.W., Lipfert, J. and Dekker, N.H. (2015) High spatiotemporal-resolution magnetic tweezers: calibration and applications for DNA dynamics. *Biophys. J.*, **109**, 2113–2125.
- Dulin, D., Barland, S., Hachair, X. and Pedaci, F. (2014) Efficient illumination for microsecond tracking microscopy. *PLoS One*, **9**, e107335.
- Huhle, A., Klaue, D., Brutzer, H., Daldrop, P., Joo, S., Otto, O., Keyser, U.F. and Seidel, R. (2015) Camera-based three-dimensional real-time particle tracking at kHz rates and Angstrom accuracy. *Nat. Commun.*, **6**, 5885.
- Dulin, D., Berghuis, B.A., Depken, M. and Dekker, N.H. (2015) Untangling reaction pathways through modern approaches to high-throughput single-molecule force-spectroscopy experiments. *Curr. Opin. Struct. Biol.*, **34**, 116–122.
- Redner, S. (2007) In: *A Guide to First-Passage Processes*. Cambridge University Press.

26. Potisopon,S., Ferron,F., Fattorini,V., Selisko,B. and Canard,B. (2017) Substrate selectivity of dengue and Zika virus NS5 polymerase towards 2'-modified nucleotide analogues. *Antiviral Res.*, **140**, 25–36.
27. Gong,P., Campagnola,G. and Peersen,O.B. (2009) A quantitative stopped-flow fluorescence assay for measuring polymerase elongation rates. *Anal. Biochem.*, **391**, 45–55.
28. Mejia,Y.X., Mao,H., Forde,N.R. and Bustamante,C. (2008) Thermal probing of E. coli RNA polymerase off-pathway mechanisms. *J. Mol. Biol.*, **382**, 628–637.
29. Abbondanzieri,E.A., Shaevitz,J.W. and Block,S.M. (2005) Picocalorimetry of transcription by RNA polymerase. *Biophys. J.*, **89**, L61–L63.
30. Kriegel,F., Matek,C., Drsata,T., Kulenkampff,K., Tschirpke,S., Zacharias,M., Lankas,F. and Lipfert,J. (2018) The temperature dependence of the helical twist of DNA. *Nucleic Acids Res.*, **46**, 7998–8009.
31. Gollnick,B., Carrasco,C., Zuttion,F., Gilhooly,N.S., Dillingham,M.S. and Moreno-Herrero,F. (2015) Probing DNA helicase kinetics with temperature-controlled magnetic tweezers. *Small*, **11**, 1273–1284.
32. Galburt,E.A., Tomko,E.J., Stump,W.T. and Ruiz Manzano,A. (2014) Force-dependent melting of supercoiled DNA at thermophilic temperatures. *Biophys. Chem.*, **187–188**, 23–28.
33. Seidel,R., Bloom,J.G., Dekker,C. and Szczelkun,M.D. (2008) Motor step size and ATP coupling efficiency of the dsDNA translocase EcoR124I. *EMBO J.*, **27**, 1388–1398.
34. Park,J.S., Lee,K.J., Hong,S.C. and Hyon,J.Y. (2008) Temperature dependence of DNA elasticity and cisplatin activity studied with a temperature-controlled magnetic tweezers system. *J. Korean Phys. Soc.*, **52**, 1927–1931.
35. Ostrofet,E., Papini,F.S. and Dulin,D. (2018) Correction-free force calibration for magnetic tweezers experiments. *Sci. Rep.*, **8**, 15920.
36. Lipfert,J., Hao,X. and Dekker,N.H. (2009) Quantitative modeling and optimization of magnetic tweezers. *Biophys. J.*, **96**, 5040–5049.
37. Mestas,S.P., Sholders,A.J. and Peersen,O.B. (2007) A fluorescence polarization-based screening assay for nucleic acid polymerase elongation activity. *Anal. Biochem.*, **365**, 194–200.
38. Sukhodolets,V.V. (2006) [Unequal crossing-over in *Escherichia coli*]. *Genetika*, **42**, 1526–1535.
39. Makeyev,E.V. and Bamford,D.H. (2000) Replicase activity of purified recombinant protein P2 of double-stranded RNA bacteriophage phi6. *EMBO J.*, **19**, 124–133.
40. Arnold,J.J., Bernal,A., Uche,U., Sterner,D.E., Butt,T.R., Cameron,C.E. and Mattern,M.R. (2006) Small ubiquitin-like modifying protein isopeptidase assay based on poliovirus RNA polymerase activity. *Anal. Biochem.*, **350**, 214–221.
41. Gizzi,A.S., Grove,T.L., Arnold,J.J., Jose,J., Jangra,R.K., Garforth,S.J., Du,Q., Cahill,S.M., Dulyaninova,N.G., Love,J.D. *et al.* (2018) A naturally occurring antiviral ribonucleotide encoded by the human genome. *Nature*, **558**, 610–614.
42. Gohara,D.W., Ha,C.S., Kumar,S., Ghosh,B., Arnold,J.J., Wisniewski,T.J. and Cameron,C.E. (1999) Production of “authentic” poliovirus RNA-dependent RNA polymerase (3D(pol)) by ubiquitin-protease-mediated cleavage in *Escherichia coli*. *Protein Expr. Purif.*, **17**, 128–138.
43. Cowan,G. (1998) In: *Statistical Data Analysis*. Oxford University Press.
44. Press,W.H., Flannery,B.P., Teukolsky,S.A. and Vetterling,W.T. (1992) In: *Numerical Recipes in C: The Art of Scientific Computing*. Cambridge University Press.
45. Squires,T.M. and Quake,S.R. (2005) Microfluidics: Fluid physics at the nanoliter scale. *Rev. Mod. Phys.*, **7**, 977–1026.
46. Charbonnier,F., Erauso,G., Barbeyron,T., Prieur,D. and Forterre,P. (1992) Evidence that a plasmid from a hyperthermophilic archaeobacterium is relaxed at physiological temperatures. *J. Bacteriol.*, **174**, 6103–6108.
47. Depew,D.E. and Wang,J.C. (1975) Conformational fluctuations of DNA helix. *PNAS*, **72**, 4275–4279.
48. Duguet,M. (1993) The helical repeat of DNA at high temperature. *Nucleic Acids Res.*, **21**, 463–468.
49. Strick,T.R., Croquette,V. and Bensimon,D. (1998) Homologous pairing in stretched supercoiled DNA. *PNAS*, **95**, 10579–10583.
50. Lansdorp,B.M. and Saleh,O.A. (2012) Power spectrum and Allan variance methods for calibrating single-molecule video-tracking instruments. *Rev. Sci. Instrum.*, **83**, 025115.
51. van Dijk,A.A., Makeyev,E.V. and Bamford,D.H. (2004) Initiation of viral RNA-dependent RNA polymerization. *J. Gen. Virol.*, **85**, 1077–1093.
52. Depken,M., Galburt,E.A. and Grill,S.W. (2009) The origin of short transcriptional pauses. *Biophys. J.*, **96**, 2189–2193.
53. Sarin,L.P., Poranen,M.M., Lehti,N.M., Ravantti,J.J., Koivunen,M.R., Aalto,A.P., van Dijk,A.A., Stuart,D.I., Grimes,J.M. and Bamford,D.H. (2009) Insights into the pre-initiation events of bacteriophage phi6 RNA-dependent RNA polymerase: towards the assembly of a productive binary complex. *Nucleic Acids Res.*, **37**, 1182–1192.
54. Winzor,D.J. and Jackson,C.M. (2006) Interpretation of the temperature dependence of equilibrium and rate constants. *J. Mol. Recognit.*, **19**, 389–407.
55. Galburt,E.A., Grill,S.W., Wiedmann,A., Lubkowska,L., Choy,J., Nogales,E., Kashlev,M. and Bustamante,C. (2007) Backtracking determines the force sensitivity of RNAP II in a factor-dependent manner. *Nature*, **446**, 820–823.
56. Shaevitz,J.W., Abbondanzieri,E.A., Landick,R. and Block,S.M. (2003) Backtracking by single RNA polymerase molecules observed at near-base-pair resolution. *Nature*, **426**, 684–687.
57. Ostrofet,E., Papini,F.S., Malinen,A.M. and Dulin,D. (2019) In: Joo,C and Rueda,D (eds). *Biophysics of RNA-protein interactions*. Springer, NY., pp. 109–141.
58. Young,J.M., Luketina,R.C. and Marshall,A.M. (1977) The effects on temperature on growth in vitro of *Pseudomonas syringae* and *Xanthomonas pruni*. *J. Appl. Bacteriol.*, **42**, 345–354.
59. Sands,J.A., Cupp,J., Keith,A. and Snipes,W. (1974) Temperature sensitivity of the assembly process of the enveloped bacteriophage phi6. *Biochim. Biophys. Acta*, **373**, 277–285.
60. Malik,O., Khamis,H., Rudnizky,S., Marx,A. and Kaplan,A. (2017) Pausing kinetics dominates strand-displacement polymerization by reverse transcriptase. *Nucleic Acids Res.*, **45**, 10190–10205.