A Viral T7 RNA Polymerase Ratcheting Along DNA With Fidelity Control

Chunhong Long a, E. Chao a, Lin-Tai Da b, Jin Yu a,⁎

a Beijing Computational Science Research Center, Beijing, 100193, China
b Shanghai Center for Systems Biomedicine, Shanghai JiaoTong University, Shanghai 200240, China

ABSTRACT

RNA polymerase (RNAP) from bacteriophage T7 is a representative single-subunit viral RNAP that can transcribe with high promoter activities without assistances from transcription factors. We accordingly studied this small transcription machine computationally as a model system to understand underlying mechanisms of mechano-chemical coupling and fidelity control in the RNAP transcription elongation. Here we summarize our computational work from several recent publications to demonstrate first how T7 RNAP translocates via Brownian alike motions along DNA right after the catalytic product release. Then we show how the backward translocation motions are prevented at post-translocation upon successful nucleotide incorporation, which is also subject to step-wise nucleotide selection and acts as a pawl for “selective ratcheting”. The structural dynamics and energetics features revealed from our atomistic molecular dynamics (MD) simulations and related analyses on the single-subunit T7 RNAP thus provided detailed and quantitative characterizations on the Brownian-ratchet working scenario of a prototypical transcription machine with sophisticated nucleotide selectivity for fidelity control. The presented mechanisms can be more or less general for structurally similar viral or mitochondrial RNAPs and some of DNA polymerases, or even for the RNAP engine of the more complicated transcription machinery in higher organisms.

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1. T7 RNA Polymerase as a Minimal Transcription Machine Model System

The RNA polymerase (RNAP) from bacteriophage T7 is regarded as one of the smallest transcription machines [1–3]. In bacteria and eukaryote species, RNAP II, the core engine of the transcription machinery, works with a variety of transcription factors to support gene expression [4–8]. RNAP II itself consists of multiple polypeptides, i.e., maintaining a complex molecular architecture. In comparison, T7 RNAP is a single-subunit enzyme with a simple hand-like structure [9–12], and it is capable of transcribing with high promoter activity or processivity, self-sufficiently, without assistances from transcription factors, from initiation to elongation and to termination. Indeed, T7 RNAP structurally resembles a wide class of DNA polymerases (DNAPs), along with some other viral and mitochondrial RNAP species [10,13]. Hence, T7 RNAP makes a minimal model system to study transcription.

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The high-resolution crystal structures of T7 RNAP had been initially determined by Sousa et al. [14] and then by Thomas A Steitz lab coworkers on its transcription initiation to elongation complexes since the late last century [15–18]. In particular, several states of T7 RNAP elongation complex have been obtained, from nucleotide insertion or substrate state to catalytic product state, and to post-translocation state, together with an additional pre-insertion complex then resolved by Temiakov et al. [19]. Meanwhile, extensive bio-chemical studies [20–23] along with single-molecule measurements on T7 RNAP elongation [24–27] provide substantial quantitative features of the enzyme kinetics, from initiation to elongation. Accordingly, physical modeling and molecular dynamics (MD) simulation on this smallest transcription machine became feasible, so that to reveal underlying molecular mechanisms and essential structural dynamics details.

We have recently studied transcription elongation of T7 RNAP by combining physical modeling and all-atom MD simulations, addressing both mechano-chemical coupling and fidelity control mechanisms during elongation [28–36]. The mechanochemistry concerns about how the protein machine utilizes chemical free energy to generate mechanical or directional motions, referring to how the chemical synthesis of RNA couples with the polymerase translocation along DNA in the RNAP system. T7 RNAP had been suggested to work via a ‘power stroke’ (PS) mechanism [17,37,38], in which product release directly drives the RNAP translocation via simultaneous protein subdomain opening. Meanwhile, T7 RNAP along with RNAPs from bacteria and eukaryotic species had also been proposed to function in a ‘Brownian ratchet’ (BR) scenario [21,25,39–42]. Below, we elaborate on how our studies actually support the BR working scenario of T7 RNAP, in which the translocation proceeds in Brownian motions after the product release, while the ratcheting part is fulfilled largely by cognate nucleotide incorporation to the growing end of the synthesizing RNA. Since non-cognate nucleotides unlikely support successful nucleotide incorporation or ratcheting, one regards that nucleotide selection plays a crucial role for the BR scenario such that an RNAP actually conducts ‘selective ratcheting’ along DNA. Accordingly, we illustrate then how T7 RNAP achieves the nucleotide selection for the transcription fidelity control. Indeed, the mechanisms can be representative or apply in general to related enzymes on the catalyzed polymerization processes, in the presence of molecular template, though specific structural elements do vary for different polymerization machines.

2. PPI Product Release Unlikely Drives the Translocation of T7 RNAP

In previous structural studies of T7 RNAP, suggestions had been made on the PS mechanism such that the pyrophosphate (PPI) product release after catalysis directly drives the translocation via rotational opening of the fingers subdomain [17]. On the other hand, early [39] and immediately later single-molecule force measurements on T7 RNAP suggested alternatively the BR scenario [25]. Accordingly, we investigated the mechano-chemical coupling by studying the PPI release first, using atomistic MD simulation [31].

Indeed, the PPI release step along with the translocation of RNAP on DNA turns out to be too fast to be monitored directly by experiments. For example, the single molecule measurements had shown that forces implemented to hinder the RNAP movements on the DNA hardly slowly down the overall elongation, suggesting that the translocation is not a rate-limiting step during an elongation cycle [25,26]. The elongation cycle of T7 RNAP, however, lasts tens of milliseconds or longer [22]. One can accordingly estimate that the fast steps of the product release and translocation happen from microseconds to sub-milliseconds [43–45], which are nevertheless too long for straightforward computational samplings by the atomistic MD simulations.

The all-atom simulation systems of T7 RNAP-DNA-RNA complex with explicit water solvent include over 100 K atoms. For systems of such a size, one can routinely simulate up to several microseconds under current high-performance computing technologies; yet it is still computationally prohibiting to further approach over tens of microseconds to millisecond time scale. Fortunately, by launching extensive sub-microseconds equilibrium simulations that spread around a wide range of conformation space for the relevant process, we were able to construct the Markov-state model (MSM) for the PPI release, and later for the translocation of T7 RNAP, which are estimated to happen at tens of microsecond time scale [31,34]. The strength and technical issues in building the MSM using MD can be found in abundant literature elsewhere [46–50].

The MSM we constructed (two hundreds micro-states according to structural root-mean-square deviations or RMSDs, and three macro-states for visualization) shows a jump–from-cavity PPI release mechanism (see Fig. 1A), in which the PPI-bound product state (S1a) and a pre-activation intermediate state (S1b) dominate the overall population (90%), while the PPI released state (S2) is achieved by thermally activating transitions S1a→S1b→S2 [31]. Inside the ‘cavity’ around the active site, PPI is held by two aspartate residues (Asp527 and Asp812) that are crucial for the catalysis. Then PPI can shift away and associate more closely with positively charged residues aligning the product release channel (e.g. Lys631 and Arg627), particularly with Lys472 that is key to assist PPI to ‘jump’ out of the cavity via the lysine side-chain swing or fluctuations. Interestingly, there is always a lysine or a homologous arginine located at the exit of the product release channel in other polymerase species (including yeast RNAP II, bacterial RNAP, human mitochondrial RNAP, and several species of DNAPs), which appears to assist the PPI release in general [31]. Note that the jump–from-cavity happens comparatively slowly at the S1b→S2 transition, which is estimated above several microseconds at least.

Essentially, the fingers subdomain does not show substantial conformational changes during the many short equilibrium simulation processes for the PPI release. Even in comparatively long simulations, the fingers subdomain or the O-helix on the subdomain shows no substantial opening, either in the PPI initially bound state, or upon PPI removal or its charge neutralization for control, up to a microsecond time scale. Anyhow, the rotational fluctuations of the O-helix increase in the control simulations, once having PPI or its charge removed. Hence, it seems that the PPI release does not necessarily couple with a progressive rotational opening of the O-helix or the fingers subdomain, which is essential to complete the RNAP translocation. Besides, the thermal activation of the PPI release also suggests that energetically it is unlikely for the release process to directly power or drive the translocation. Instead, the PPI release likely only enhances the rotational flexibilities of the fingers subdomain, which then facilitates the RNAP translocation thereafter.

3. Translocation Proceeds in Brownian Motions and is Facilitated by the O-helix Fluctuation Opening at Pre-translocation that may also Prevent Backtracking

Then we employed even more extensive MD simulations in aggregation to ~10 µs to construct the MSM of the T7 RNAP translocation on DNA, by clustering a large amount of simulation snapshots into 500 micro-states, according to time-structure independent component analysis (tICA) [51,52]. The resulted model is further simplified for visualization into a six-state macro-state representation (see Fig. 1C) [34].

In the six-state translocation network model representation, both the O-helix and Y-helix on the fingers subdomain play significant roles, and they show rotational opening motions in non-synchronized manner. Importantly, in the initial pre-translocation state (S1), both the O-helix and Y-helix show a ‘semi-open’ conformation on average (i.e., the rotation angle peaked ~15°), with significant wide fluctuations spanning from the closed conformations to open ones (34°). Microsecond transition into a less-populated pre-translocation configuration (S2) allows base un-stacking of the transition nucleotide (TN) from F644 (on the Y-helix) yet somehow quenches the O/Y-helix opening to the close status. The Y-helix opens first in the transition state (S3),
after that the O-helix opens (S4 to S6). Essentially, Y639 (on the O-helix) pushes onto the 3′-end of the RNA to allow it to move ahead of the template DNA (S3). Hence, the O-helix opening seems to well couple with the DNA forward translocation. Overall, the free energy profile of translocation appears comparatively flat so that Brownian motions dominate. The slowest step of the translocation takes place in the transition to the key intermediate state (S3), which is estimated to last over tens of microseconds at least.

Interestingly, the O-helix along with the Y-helix (or say the fingers subdomain) seems to be able to open by enhanced rotational oscillations or fluctuations (after the PPi release) in the pre-translocation state. Note that it is NOT progressive opening yet until toward the post-translocation state. Previous studies had also concerned about rotational flexibilities of the thumb subdomain [53,54]. We also noticed substantial rotational movements (~25°; non-published results) of the thumb subdomain from the pre- to the post-translocation. Backtracking turns out to be an efficient way of coordination for RNAP to deal with errors of nucleotide incorporation, i.e., by proofreading or editing; or it supports necessary pauses during the transcription elongation, e.g., to coordinate with translation by ribosome [55–57]. Although backtracking had been identified in multi-subunit RNAPs or even the single subunit mitochondrial RNAP (mtrRNAP) [58,59], it has not been detected for T7 RNAP. We then hypothesized that the very mechanism to facilitate the translocation, i.e., the O-helix fluctuation to opening in the pre-translocation state, may also prevent T7 RNAP backtracking. To test the hypothesis, we computationally designed a
mutant T7 RNAP with several residues replaced on the O-helix to mimic the mtRNAP that is structurally similar to T7 RNAP [60]. Our simulation results showed that the mutant T7 RNAP would have the O-helix closed upon the 3′-end of the RNA being pulled to initiate the backtracking from the pre-translocation, while the O-helix opens for such a response in the wild-type T7 RNAP [34]. Hence, we considered that the mutant T7 RNAP we made might be able to backtrack to some extent. Preliminary experimental studies supported the hypothesis such that the mutant T7 RNAP maintained transcription activities, but lower than the wild-type system [34]. Further experimental investigation is still needed, e.g., at a single molecule level, to confirm on the mutant and wild-type T7RNAP capabilities of backtracking.

4. Selective Ratcheting Starts From the Nucleotide Pre-insertion Checkpoint

In an RNAP elongation cycle, the incoming nucleoside triphosphate (NTP) is recruited into the RNAP enzyme active site according to the Watson-Crick (WC) base pairing with the template DNA nucleotide. The RNAP translocation allows the incorporated nucleotide at the 3′-end of the synthesizing RNA to move upstream to vacant the active site. According to the BR scenario, prior to the NTP association, RNAP can keep moving forward and backward on DNA, due to the nearly flat free energy surface of the translocation. Once the incoming NTP binds and occupies the active site, the backward movement of RNAP can be prevented, so that forward translocation is finally biased upon the full nucleotide incorporation. That says, the NTP association and incorporation act as a ‘pawl’ in a ratchet device to achieve the BR process (see Figs. 2A and 1B) [61]. Nevertheless, non-cognate nucleotide species may also bind but likely dissociate prematurely before chemical synthesis, due to the nucleotide selectivity conducted by RNAP for the purpose of fidelity control. Consequently, only those successfully incorporated nucleotides, or in principle, the cognate nucleotide species, contribute to the ratcheting of RNAP along DNA.

A prominent feature of the nucleotide addition cycle (NAC) of T7 RNAP and related single-subunit polymerase species is that a nucleotide pre-insertion state exists [12,19,62,63], and our studies confirm that the pre-insertion state serves well as an initial kinetic checkpoint to screen non-cognate nucleotide species out of the active site [30,64,36]. Indeed, the pre-insertion complex of T7 RNAP in a ‘semi-open’ conformation had been crystalized with a cognate nucleotide bound to a pre-insertion site, slightly away from the active site [19]. Although the WC base pairing had not been well captured between the pre-insertion nucleotide and the template counterpart in the crystal structure, our MD simulation on the pre-insertion complex revealed the WC base pairing formation after ~ 50 ns equilibrium simulation [30]. Besides, when we replaced the cognate nucleotide (rATP) by the non-cognate species (rGTP, dATP etc.) from the pre-insertion crystal structure complex and conducted equilibrium simulations accordingly, we found that the non-cognate nucleotide would be ‘grabbed’ by Y639, which actually blocks the insertion site of the DNA template nucleotide (i.e., the transition nucleotide or TN) in the pre-insertion complex. In such an equilibrated rGTP pre-insertion complex (see the rGTP off-path pre-insertion structure in Fig. 2B upper right, or Fig. 2D config 1 in the middle...
row), as the Y639 side chain grabs rGTP, the template TN has its base rotated away from rGTP, as well as away from its post-translocation configuration [30,64,36]. In the dATP case (see the dATP off-path pre-insertion structure in Fig. 2B lower right), the template TN is even pushed ‘backward’ (to an intermediate configuration between the post- and pre-translocation state), while the Y639 side ring stacks with the upstream DNA template nucleotide and forms a ‘fake’ base pairing with dATP [30,64,36]. That is to say, by competing with the template TN in association with the incoming NTP, Y639 plays a critical role in deterring the non-cognate NTP at pre-insertion. Interestingly, comparing the rATP pre-insertion structure with that of dATP (off-path), one can see that a ‘bridge’ water molecule forms hydrogen bonds with the –OH group of Y639 side chain and that on the ribose of rATP, respectively; in contrast, no such a water bridge can be found in the dATP case due to lack of the ribose –OH for dATP [30] (see Fig. 2B lower panel).

Indeed, the non-cognate nucleotide pre-insertion complexes modeled above are denoted as the ‘off-path’ pre-insertion structures [64], which are at disadvantage for further nucleotide insertion but are easy for the non-cognate dissociation. In the off-path pre-insertion structure, Y639 directly associates with the non-cognate nucleotide to prevent it from closely interacting with the template TN. Energetically, it has been calculated from the MD simulations that the non-cognate nucleotide (rGTP e.g.) dissociation free energy almost vanishes at the off-path pre-insertion site, while the dissociation free energy for the cognate nucleotide from the pre-insertion site appears to be ~ 4 kBT [65]. Upon such a kind of cognate ‘trapping’, and non-cognate off-path Y639 ‘grabbing’ but energetically ‘non-trapping’ at pre-insertion, ~98% of the non-cognate nucleotide species can be already screened and removed from accessing further to the active site.

To compare, we also modeled the so-called on-path pre-insertion complexes for the non-cognate nucleotide species, by starting from the equilibrated pre-insertion complex of the cognate nucleotide, and then algebraically converting the cognate nucleotide into the non-cognate species (from rATP to rGTP and dATP e.g.) [64,36]. Accordingly, the modeled on-path pre-insertion structures appear similar to the cognate pre-insertion structure: In particular, the non-cognate nucleotide has its base closely associated with that of the template TN, even though in the absence of the WC base pairing (see Fig. 2B for rGTP on-path or Fig. 2D bottom config 1 for dATP on-path). From the alchemical free energy calculations, we got to see that the non-cognate rGTP on-path pre-insertion structure was energetically less stable (~3 kBT) than that of the cognate rATP [64], while the non-cognate dATP on-path pre-insertion structure was slightly more stable than that of the rATP structure (~1 kBT) [36]. Further calculations reveal that the rGTP on-path pre-insertion is indeed highly non-accessible due to a very high rGTP association barrier (with template TN a dTTP), while the dATP on-path pre-insertion does accommodate for a fairly easy dATP association, and allow for further nucleotide insertion and selection following the on-path (see Fig. 2D bottom).

5. Selective Ratcheting Proceeds Through Slow Nucleotide Insertion With Substantial Selection

The nucleotide insertion accompanied by a conformational transition from a ‘semi-open’ to the closed form has been suggested to be a slow step during the NAC in T7 RNAP from previous biochemical studies [22]. It is highly likely that the slow insertion step is employed by the enzyme to substantially scrutinize against non-cognate nucleotide species to achieve sufficiently high transcription fidelity [35]. In viral T7 RNAP elongation, an elongation error rate reaches to ~10^{-4} to ~10^{-6} for the base mismatch incorporation (e.g. rGTP replacing rATP) [66,67], or ~10^{-2} for the deoxyribonucleotide mis-incorporation (e.g. dATP replacing rATP) [20]. Note that T7 RNAP has not been detected with proofreading or editing activities yet, hence, its fidelity control may fully rely on the nucleotide selectivity, from pre-insertion to insertion, and likely also during catalytic reaction.

To probe the insertion structural dynamics and energetics, we conducted systematic umbrella-sampling simulations in aggregation to microseconds for several nucleotides, along a highly collective coordinate for each nucleotide insertion [36]. The collective coordinate is defined according to the rmsd changes of a highly relevant set of atoms that are essentially involved from the nucleotide pre-insertion to insertion, which include five helices from the fingers subdomain, the insertion nucleotide, and the template DNA nucleotide. The free energy profiles along the i rmsd (in reference with the pre-insertion and insertion structures), or say, the potential of mean forces (PMFs) of the cognate rATP and non-cognate rGTP off-insertion are shown representatively (see Fig. 2C). In the cognate rATP insertion, an activation barrier ~3 kBT occurs at the intermediate conformation (config 3 in Fig. 2D top), where the O-helix opens up to ~25° (~15° in config 1) to resist the nucleotide insertion. In the end, the insertion state is reached as ~3 kBT more stabilized than the pre-insertion state, while the O-helix closes to ~5° on average. It is worth pointing out that the thumb subdomain also rotates back (non-published results), along with the closing motions of the fingers subdomain, from the pre-insertion to the insertion state. The on-path insertion free energy profile of the non-cognate rGTP appears quite similar to that of the cognate rATP, although in reality, only a prohibitively low population (~1%) of non-cognate rGTP can be loaded at on-path pre-insertion for the further on-path insertion [36].

In comparison, the off-path insertion PMF of rGTP incurs a quite high barrier (~7 kBT; see Fig. 2C right), as the O-helix opens up ~30° in the intermediate state (config 3 in Fig. 2D middle). In addition, the template DNA nucleotide TN deviates its base from the incoming nucleotide all the way from the beginning (config 1–4), until finally the insertion state (config 5) is reached. Thus, the large deviation of the template base from the insertion nucleotide together with the O-helix full opening contributes to the significant high insertion barrier, which is ~4 kBT higher than that of the cognate rATP and serves strongly for the mismatch insertion inhibition. The non-cognate rGTP insertion state is also ~3 kBT more stabilized than the pre-insertion off-path, and ~3 kBT less stabilized than the cognate rATP insertion state. The O-helix closes ~5° into the rGTP insertion structure [36].

Upon the pre-insertion rejection and further insertion inhibition, the elongation error rate (i.e., for rGTP replacing rATP) would be reduced to ~10^{-3}, according to our calculations via a chemical master equation (CME) approach [35,68]. Hence, to reduce the error rate further down to ~10^{-4}, selection into the catalytic stage seems to be required, for which we estimate that a selection energy ~7 kBT is necessary. That means during the catalytic reaction, the non-cognate rGTP may face with an activation barrier ~7 kBT higher than that of the cognate rATP. Hence, even though our classic MD simulation approach cannot compute the catalytic energetics, we could still predict quantitatively the catalytic selection energetics, combining MD simulation results with experimentally measured error rate information [36].

On the other hand, the energetics for the insertion of deoxy-ribonucleotide dATP appears quite differently from the base mismatch rGTP. Firstly, we could infer that the pre-insertion on-path for dATP is quite accessible as the off-path pre-insertion configuration. Then the calculated PMF of the on-path dATP shows a significant high barrier (~6 kBT or 3 kBT higher than that of rATP; see Fig. 2D bottom), which also corresponds to the O-helix opening above 30° at the transition intermediate (config 3). Interestingly, we could find that two magnesium ions switch positioning of leave and stay during the dATP on-path insertion, but not in other cases. According to a two magnesium catalysis mechanism suggested for the polymerase action [69,70], MgA remains close to the 3′-end of RNA before and after the catalysis, while MgB comes with the insertion NTP and leaves with the product PPI release. However, varying for several types of simulation force field settings, we could always find that MgA and MgB switch their positions during the on-path dATP insertion (Fig. 2D bottom), likely due to the lack of a negative charge on the dATP ribose, so that MgA soon drifts toward the
beta-phosphate to ‘collide’ with MgB, which binds there early during the insertion [36]. Accordingly, the high energetics involved in the dATP on-path insertion appears to be related to the two-magnesium ion switching, which may serve for the selectivity against the deoxy-ribonucleotide incorporation in T7 RNAP.

The off-path insertion of dATP, on the other hand, does not incur a very high barrier (~1 k_BT higher than the cognate rATP); along this path, however, the O-helix cannot close below ~10° to reach a proper insertion state, until it possibly relaxes or merges with the on-path near the final insertion stage. Anyhow, due to the prominent selection coordinated by Y639 at the dATP pre-insertion off-path, together with the significant inhibition on-path, the error rate (i.e. dATP replacing rATP) can be achieved at 10⁻² as experimentally detected [20,71], according to our CME calculation. Hence, no further selection seems to be needed at the catalytic stage. That is to say, the non-cognate dATP, once being properly inserted, can be catalytically added to the growing end of the RNA as easy as the cognate rATP. Interestingly, previous experimental studies had found that by simply mutating Tyr to Phe at residue 639, the mutant Y639F RNAP can allow the dNTP incorporation as efficient as rNTP [20,71]. Our simulation study accordingly suggested the underlying mechanism by attributing the differentiation between dNTP and rNTP at pre-insertion to the −OH hydroxyl group of Tyr639 (see Fig. 2B bottom for the water bridge), which is lack of for Phe [30,64,36]. Early mutant studies indicated additional amino acids such as H784 in regulation of the nucleotide selectivity [20,72]. We also noticed that H784 could not approach sufficiently close to an incoming dNTP as to an rNTP into the insertion state in the MD simulations. How exactly H784 regulates the selectivity deserve further investigations by possibly studying the double mutant Y639F/H784A and/or including additional modifications to the non-cognate species.

6. Conclusion

Combining atomistic MD simulations with further quantitative analyses, we surveyed a complete NAC during T7 RNAP elongation and dissected detailed structural dynamics mechanisms of the Brownian movements of RNAP along DNA and the selective ratcheting processes for fidelity control. The Brownian-alike motions are thermally activated, with quasi-equal free energetics pre- and post-translocation at the equilibrium condition. The biased ratcheting then requires the system to work at non-equilibrium, i.e., driven by chemical free energy from the polymerization reaction. The template-bound nucleotide selectivity is accordingly achieved at the non-equilibrium steady state during the RNAP elongation, which leads to substantially improved fidelity in comparison with the equilibrium condition or with the slow polymerization in the absence of the enzyme. The nucleotide selection is actually conducted stepwise from pre-insertion to insertion, and to catalytic reaction, prior to potential proofreading and further editing. For single-subunit polymersomes similar to T7 RNAP, both the pre-insertion rejection and insertion inhibition of the non-cognate nucleotides play significant roles in the fidelity control, which are coordinated by the fingers subdomain opening and closing motions together with key residue interactions. Accordingly, only for those nucleotides that pass through the above selection checkpoints from initial association to final incorporation, successful ratcheting steps of the RNAP can be achieved. In summary, the nature’s design of this small but highly capable viral transcription machine appears to be: The mechanical movements of RNAP along DNA mean to be thermally activated without much bias; the RNAP ratcheting forward along DNA is energetically supported by chemical reaction of polymerization, which is tightly coupled with genetic information interrogation for transcription fidelity control, and subjects to evolution pressure of biochemical synthesis. Furthermore, it is important to be aware that the transcription activation and ratcheting energetics involved in each NAC are closely tied to sequence stability variations upon displacing the transcription bubble along DNA [73,74]. Notably, genome-wide transcription dynamics and accuracy measurements could reveal sequence motifs for RNAP pausing, backtracking, or editing [75–77]. Comparative studies between non-backtracking and backtracking RNAP species, for example, may possibly contribute to elucidate consensus or enzyme-dependent motifs and mechanisms. As a minimum transcription machine, T7 RNAP had been laboratory evolved, engineered or redesigned for various purposes [78–80]. The redesigning of T7 RNAP variants with potential functions such as backtracking thus brings another trial of the synthetic approaches to the living systems.

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