A general theoretical framework to design base editors with reduced bystander effects

Qian Wang, Jie Yang, Zhicheng Zhong, Jeffrey A. Vanegas, Xue Gao & Anatoly B. Kolomeisky

Base editors (BEs) hold great potential for medical applications of gene therapy. However, high precision base editing requires BEs that can discriminate between the target base and multiple bystander bases within a narrow active window (4 – 10 nucleotides). Here, to assist in the design of these optimized editors, we propose a discrete-state stochastic approach to build an analytical model that explicitly evaluates the probabilities of editing the target base and bystanders. Combined with all-atom molecular dynamic simulations, our model reproduces the experimental data of A3A-BE3 and its variants for targeting the "TC" motif and bystander editing. Analyzing this approach, we propose several general principles that can guide the design of BEs with a reduced bystander effect. These principles are then applied to design a series of point mutations at T218 position of A3G-BEs to further reduce its bystander editing. We verify experimentally that the new mutations provide different levels of stringency on reducing the bystander editing at different genomic loci, which is consistent with our theoretical model. Thus, our study provides a computational-aided platform to assist in the scientifically-based design of BEs with reduced bystander effects.
The development of genome editing tools associated with the clustered regularly interspaced short palindromic repeat (CRISPR) systems has revolutionized biomedical studies. Holding great potential for the treatment of genetic diseases, diverse precise genome editing tools based on CRISPR-Cas9 have been developed, such as homology-directed repair (HDR) based systems, as well as cytosine and adenine base editors (BE)\textsuperscript{1–3}. While the HDR method requires double-stranded DNA breaks (DSBs) and causes unpredictable editing outcomes, BEs use nickase Cas9 (nCas9), enabling more precise modifications without generating DSBs\textsuperscript{4–8}. For example, cytosine BEs (CBEs) are constructed by fusion of a cytidine deaminase domain with nCas9. This fusion protein forms a complex with the guide RNA and performs site-specific deamination to convert cytosine (C) to uracil (U) in the deaminase activity window. The base U is subsequently replaced with thymine (T) by the endogenous cellular repairing machinery, resulting in an overall C-to-T substitution at the defined genomic site. Since point mutations are responsible for more than half of human disease-associated genetic variants\textsuperscript{9}, BEs are superior than HDR based systems due to their higher editing efficiency in the correction of pathogenic single nucleotide polymorphisms\textsuperscript{9}, avoidance of unwanted DSBs, and prevention of insertions and deletions\textsuperscript{4,5}.

While engineering of several BE variants has improved product purity and overall editing efficiency\textsuperscript{8,9}, one of the major challenges in base editing is the discrimination of multiple identical bases located within the deaminase activity window of 4–10 nucleotides. As a result, the target base and other bystander bases will all be modified, negatively impacting the precision of genome editing outcomes. To address this issue, the introduction of beneficial mutations to deaminase has further advanced BEs\textsuperscript{10–12}. For example, compared to the wild-type APOBEC3A (A3A)-BE3, an engineered A3A CBE with the mutation N57G maintained high editing activity at the target C in the TCR motif with greatly reduced activity against bystanders\textsuperscript{11}. Also, followed by several rounds of screening and validation of rational mutagenesis, we previously engineered an APOBEC3G (A3G)-CBE that preferentially edits the second C in the “C C” motif with 6000-fold improvement in perfectly modified alleles compared to the original BE4max\textsuperscript{12}. Despite these successes, a general theoretical framework to guide the design of mutations that can lead to high editing activity at the target base and low activity at bystanders (defined as BE high editing selectivity) is still missing. Mutation selections in these previous studies were mostly suggested by structural considerations: starting from the identification of key residues in the deaminase near the DNA binding motif and then mutating those residues to form a candidate library for experimental validation. The design process could be greatly accelerated with a comprehensive theoretical model that could quantitatively explain and predict the effect of specific mutations on editing activity at the target base and bystanders. In addition, such a theoretical model would also improve our fundamental understanding of the biochemical and biophysical processes that take place during base editing.

Molecular dynamic (MD) simulations have been used to study the activity of BE complexes and the role of beneficial mutations to enhance overall editing activity (both at target and bystanders)\textsuperscript{13,14}. Herein we present a comprehensive multi-scale theoretical approach to describe the molecular processes taking place during BE editing, explaining at the microscopic level the role of beneficial mutations in discriminating the target base over bystanders. To fulfill this goal, we built a general theoretical framework combining a discrete-state stochastic (chemical-kinetic) model and MD simulations, explicitly calculating the base editing probability at both the target base and bystanders. In our model, we include an important parameter, $\Delta E_m$, the binding affinity between deaminases and ssDNA. This parameter was modulated by introducing various mutations into BE and its values were measured through MD simulations. This framework helps establish a relationship between mutations and BE editing selectivity. We then proposed a theoretical principle arguing that the BE selectivity is non-monotonically dependent on $\Delta E_m$. It is argued that the highest BE selectivity can be obtained by varying the binding affinity. In addition, other relevant kinetic parameters are included in the model, such as the binding rate between Cas9 and ssDNA and the deamination rate of BE, allowing us to discuss how $\Delta E_m$ correlates with those parameters to affect BEs editing selectivity. Our model successfully explains how the mutations influence the editing selectivity of A3A-BE3. Finally, we designed mutations to further improve the selectivity of the A3G-BE system and we verified the improved editing selectivity experimentally. Thus, the framework we propose opens multiple opportunities for future efficient engineering of BE using theory-driven methods.

**Results**

**Kinetic model of base editing.** We developed a discrete-state stochastic model to describe the dynamics of target and bystander editing. This is a minimal chemical-kinetic approach that considers the most relevant chemical states and features of base editing. For convenience, unless noted otherwise, we will use A3A-BE3 to edit the EGFP site 1 as an example.

In this theoretical model (Fig. 1), it is assumed that the Cas9 domain of CBE can bind to ssDNA with a rate $w_0$, initiating the base editing (transition from state 0 to state 2). Alternatively, the protein complex can go to an unproductive state where editing cannot take place, with a rate of $u_2$ (transition from state 0 to state 1). Next, either the Cas9 domain dissociates from DNA with a rate $w_0$ (Transition from state 2 to state 0), or the target cytidine binds to the deaminase catalytic site with a rate $u_1$ (transition from state 2 to state 3). Then the cytidine can either dissociate from the site with a rate $w_1$ without being edited (backward step from state 3 to state 2), or it can be chemically transformed to uridine with a rate $u_3$ (transition from state 3 to state 5). Similarly, the bystander cytidine may bind to the deaminase with a rate $u_2$ (transition from state 2 to state 4), and subsequently, it can either unbind with a rate $w_2$ without being edited (transition from state 4 back to state 2), or it can be chemically transformed with a rate $u_4$ (transition from state 4 to state 6). After that, while Cas9 is still bound to DNA (being in the state 5 or 6), the deaminase can continue editing other cytidines in this region with the same sequence of events (transition to the states 9–12). Alternatively, if Cas9 dissociates from DNA, uridine will be transformed to thymidine through DNA repair (transitions from state 5 to states 7 and 13, or transition from state 6 to states 8 and 14). This U-to-T editing decreases the rebinding rate of Cas9 to ssDNA (transition $7 \rightarrow 5$) if the endogenous DNA repair and replication machinery has changed the DNA sequence from G:C pair to A:T pair. In this case, the new DNA sequence does not perfectly match the spacer sequence of sgRNA. Because the repairing rate is unknown, the rebinding rate is assumed to be $m t_{d0}$ with $0 \leq m \leq 5$. The parameter $m$ reflects how much the rebinding ability of the BE complex is lowered in comparison with the original substrate. If the DNA repairing rate is slow then $m$ tends to be closer to 1; otherwise, $m$ tends to be closer to 0. Note that the kinetic network in Fig. 1 is a minimal description of complex chemical processes that take place during base editing.

To evaluate the dynamics of base editing, we explored the first-passage probabilities method successfully used in various problems in chemistry, physics, and biology\textsuperscript{15–18}. In the case of EGFP site 1 editing by A3A-BE3 there are four possible outcomes as shown in Fig. 1: CTC (state 1, failed editing), CTT (state 13,
only the target base is edited), TTC (state 14, only the bystander is edited) and TTT (state 12, both the target base and the bystander are edited). The explicit solution for the probability for the system to end up in one of these products is given below (see derivations in the Supplementary Information Appendix):

Similarly, the overall probability of editing the bystander cytidine, \( P_b \), was calculated as:

\[
P_b = P_{TTC} + P_{TTT}
\]  

Our goal is to parameterize the model by reproducing experimentally measured probabilities, \( P_t \) and \( P_b \). Here, we assume that the binding between the cytidine (both target and bystander) and the deaminase is mainly a diffusion-controlled process. Therefore, considering that target and bystander cytidine are chemically identical and very close spatially, we added an additional approximation:

\[
u_2 = u_1
\]

\[
w_2 = w_1 e^{\Delta \Delta E_b/kT} = w_1 e^{\left[\Delta E_b(\text{bystander}) - \Delta E_b(\text{target})\right]/kT}
\]
The physical meaning of these expressions is the following: the binding rate to the target or the bystander are the same, but the unbinding is governed by the strength of the interactions between the DNA substrate and the protein complex. In Eq. (9), the term $\Delta E_o$ represents the binding free energy between the ssDNA binding motif and the deaminase. $\Delta E_b$ represents the difference in $\Delta E_o$ between the dissociation from the target base and the dissociation from the bystander base. This difference arises from the sequence shift in the binding interface. An example is shown in Fig. 1, where the sequence of ssDNA binding motif changes from "T1C0" in the case of target editing, to "G1C0" in the case of bystander editing. This change can be formalized by a mutation from thymine to guanine at position -1, which perturbs the binding free energy and further influences the unbinding rate $w$ of the cytidine from the catalytic site. Note that this approximation can also be explained using thermodynamic arguments, since the ratio between rates of binding and unbinding is related to the free energy difference between two states: the state where the protein-RNA complex is bound to the DNA chain and the state where both DNA and protein complex are free, $u_{\text{w}} = e^{-\Delta E_0/\kappa B T}$, $u_{\text{b}} = e^{-\Delta E_b/\kappa B T}$. Using Eq. (8) one can derive the result in Eq. (9).

Similarly, any deaminase mutation can be represented as a perturbation in binding free energy relative to the wild type, $w_{1, \text{mutation}} = w_{1, \text{WT}} e^{\Delta E_1/\kappa B T} = w_{1, \text{WT}} e^{(\Delta E_1(\text{mutation}) - \Delta E_1(\text{WT}))/\kappa B T}$

$\Delta E_m$ represents the difference in free energy due to mutations. Substituting Eqs. (8–10) into Eqs. (6–7), we obtain:

$$P_t = \frac{\left( y_1 + m + y_1 y_2 + y_1 y_3 e^{\Delta E_1/\kappa B T}\right) \left( 1 + y_1 e^{\Delta E_0/\kappa B T} \right) \left( 1 + y_2 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right)}{\left( y_1 + m + y_1 y_3 + y_1 y_3 e^{\Delta E_1/\kappa B T}\right) \left( 1 + y_1 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_1 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right)}$$

$$P_b = \frac{\left( y_2 + m + y_2 y_1 + y_2 y_3 e^{\Delta E_1/\kappa B T}\right) \left( 1 + y_2 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_2 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right)}{\left( y_2 + m + y_2 y_3 + y_2 y_3 e^{\Delta E_1/\kappa B T}\right) \left( 1 + y_2 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_2 e^{\Delta E_1/\kappa B T} \right) \left( 1 + y_3 e^{\Delta E_1/\kappa B T} \right)}$$

Equations (11–15) give the full analytical expressions in terms of kinetic rates and binding affinities that can be used to calculate the editing probability. There are six free parameters to describe the base editing process ($y_1$, $y_2$, $y_3$, $m$, $\Delta E_0$, $\Delta E_m$) but this number can be reduced using additional information. For example, previous binding experiments have indicated that A3A binds to ssDNA with $K_d = 57\mu M$, $K_M = 62\mu M$ and $k_{cat} = 1.1/s$. From these values, one can infer that $w_{1, \text{WT}} = 12.54/s$ and $u_{\text{b}} = 1.1/s$. Therefore, after the cytidine binds to the catalytic site, the relative probability between unbinding and the chemical transformation step, $y_3$, is 11.4. Next, if the changed ssDNA sequence no longer perfectly matches the sgRNA sequence, we assume that successful editing prevents rebinding of Cas9 to ssDNA, therefore $m=0$. Nevertheless, we show below that this assumption only has a minor effect on the final results. Lastly, we performed all-atom computational simulations to estimate $\Delta E_0$ and $\Delta E_m$, as shown in the next section. As a result, only two free parameters remain in the model, $y_1$ and $y_3$ (Eqs. 13 and 15), both of which are parameterized by reproducing experimental values of $P_t$ and $P_b$.

Computational estimates of binding free energy changes. We chose four CBEs developed by the Joung group: A3A(S99A), A3A(Y130F), A3A(N57Q), and A3A(N57A) to calculate the binding free energy changes between ssDNA and A3A. These CBE variants reduce the bystander effect to different extents while maintaining a high probability of on-target editing. The binding interface in the wild type A3A-ssDNA binding complex is shown in the crystal structure (PDB ID: 5KEG) (Fig. 2a). The carbonyl oxygen of Ser99 forms a hydrogen bond with the N4 atom of the cytidine in the catalytic site (dC0). The hydroxyl group of Tyr130 forms a hydrogen bond with the S'-phosphate of dC0. Lastly, the nitrogen atom in the sidechain of Asp57 forms a hydrogen bond with the O3 atom of dC0. Therefore, all four CBE variants appear to destabilize the binding interface. To quantitatively calculate $\Delta E_0$ and $\Delta E_m$, we utilized the so-called "alchemical free-energy calculations" based on MD simulations, ($\Delta G_2$). A thermodynamic cycle was constructed to convert $\Delta E_0$ and $\Delta E_m$ (Fig. 2b, $\Delta G_2 - \Delta G_1$) to the difference between two slow chemical transitions (Fig. 2b, $\Delta G_2 - \Delta G_4$). One transition is the free energy change for the A3A-ssDNA complex due to mutations (Fig. 2b, $\Delta G_4$) whereas the other is the free energy change for A3A alone due to mutations (Fig. 2b, $\Delta G_2$). Calculated values indeed show that mutations cause an apparent increase in the deaminase-ssDNA binding free energy (Fig. 2c), consistent with predictions based on the structural data.

The rationale for A3A mutants that reduce the bystander effect. To check whether our model can reproduce the experimentally measured on-target and bystander editing probability, we substituted $\Delta E_0$ and $\Delta E_m$ calculated above into Eqs. (11–15) and adjusted $y_1$ and $y_3$. The resulting theoretical prediction is in very good agreement with the experimental measurements (Fig. 3a), with values $y_1 = \frac{\Delta G_1}{\kappa B} = 2.1$ and $y_3 = \frac{\Delta G_4}{\kappa B} = 2.9 \times 10^{-5}$. The value of $y_1$ indicates that there is a significant fraction of BEs failing to initiate editing, whereas the value of $y_3$ suggests that the residence time of Cas9 on ssDNA is sufficient for the deaminase to function. We note here that the choice of $m$, which quantifies the effect of sgRNA mismatch on the rebinding rate of Cas9 and ssDNA, does not significantly affect the result (Fig. S1). The model also well produced the editing patterns at multiple genomic loci (Fig. S3), demonstrating the generality of this model.
Weakening the binding interface up to 4 \( k_{\mathrm{BT}} \) and that of the reversed events (Fig. 1, state 12). The outcome is largely in favor of the bystander until the target cytidine is edited (Fig. 1, state 5), the system has the choice of getting released with the product CTT (Fig. 1, state 13) or to continue editing the bystander, leading to the product TTT (Fig. 1, state 13). The outcome is largely influenced by the ratio between \( \gamma_2 = \frac{u_2}{u_3} \) and \( \gamma_1 = \frac{u_1}{u_3} \). If \( \gamma_2 \) is significantly larger than \( \gamma_1 \), bystander editing will be blocked because the residence time for the bystander cytidine in the catalytic site is too short to complete the transition to thymidine. Analytically, after the target cytidine gets edited, the probability ratio between editing the target and bystander cytidine, respectively.

\[
R_2 = \frac{P(\text{state} 5 \rightarrow 13)}{P(\text{state} 5 \rightarrow 12)} = \frac{w_0}{u_2} \left( 1 + \frac{w_3}{u_3} \right) = \gamma_3 \left( 1 + \gamma_2 e^{\frac{-6 k_{\mathrm{BT}} \Delta \Delta E}{k_{\mathrm{BT}}}} \right)
\]

For A3A(WT), \( R_2 \) is 0.17, meaning that the dominant product is TTT (Fig. 3b, purple square at \( \Delta \Delta E_m = 0 \)). This explains that for the wild-type A3A the editing efficiency of the target cytidine is similar to that of the bystander. In sharp comparison, as \( \Delta \Delta E_m \) increases by 4.5 \( k_{\mathrm{BT}} T \) for A3A(N57G), \( R_2 \) is 14.8. Now the dominant edited product changes to CTT (Fig. 3b, green line at \( \Delta \Delta E_m = 4.5 k_{\mathrm{BT}} T \)). In this case, A3A(N57G) minimizes the bystander effect while maintaining a high probability of editing the target base.

Similar arguments can be presented for the other three A3A mutants (S99A, Y130F, and N57Q). Calculations (Table S1) show that \( R_2 \) gradually increases from 0.42 for S99A to 4.91 for N57Q mutants, indicating that the bystander effect gradually decreases. This is consistent with experimental findings (Fig. 3). However, it is critical to note that to gain high editing selectivity, mutated residues have a non-monotonic effect in the deaminase-ssDNA interface. Here, selectivity is defined as the difference in probabilities between editing the target and editing the bystander. Weakening the binding interface up to 4–6 \( k_{\mathrm{BT}} T \) (depending on...
the system) greatly improves selectivity (Fig. 3a), but it drops when \( \Delta \Delta E_m \) continues further increasing. This result can be explained using the following physical considerations. Increasing \( \Delta \Delta E_m \) leads to faster-unbinding rates between cytidine and deaminase. At moderate values of \( \Delta \Delta E_m \), target editing is less affected (Fig. 1, state 2→5) but bystander editing is blocked (Fig. 1, state 5→12). However, for very large values of \( \Delta \Delta E_m \), both editing pathways are essentially blocked and the system prefers to go into the inactive state (Fig. 1, state 1). Therefore, proper modulation of the binding interface is the key to optimize base editing selectivity. We further prove this point in the next section.

The computational model helps design new A3G-BEs with improved editing selectivity. In this section, we employ our theoretical model to optimize the editing selectivity of the base editor A3G3.1 (Fig. 4a). First, the editing profiles at both target and bystander bases were calculated by using Eqs. (11–15). Our calculations show that improving the editing selectivity of A3G3.1 requires mutations that increase \( \Delta \Delta E_m \) by 2–3 kT (shaded area in Fig. 4a). Second, specific mutations were designed and \( \Delta \Delta E_m \) was calculated for each mutation by alchemical free-energy calculations as described above. Four mutations (T218S, T218N, T218I and T218G) fell into these selection criteria. A failure example is T218W, which loses the editing activity at the target base owing to overly increased \( \Delta \Delta E_m \) (Fig. 4a). We then experimentally verified these four mutations at three genomic loci containing the “TCC” motif, including EMX1 #a3, PPP1R12C #a1, and ATM #1. We chose these target sites with the “TCC” motif, which are generally more challenging over “ACC” or “GCC” for selectively editing the second C since “T” and “C” are structurally more similar. In the “TCC” case, the deaminase tends to treat “T” as a “C” and preferentially edits the bystander first “C” as well. In our tests, A3G3.14 (A3G3.1 with T218S) and A3G3.15 (A3G3.1 with T218N) generally show much improved editing selectivity (Fig. 4b), with marginally or modestly decreased editing efficiency. Therefore, A3G3.14 and A3G3.15 were further tested at other five genomic loci, including MMS22L #1, FANC E #1, MRPL44 #1, FANCF #c1, and MRPL40 #1 (Fig. 4c). Compared to the original A3G3.1, the target-to-bystander editing ratio increases from average 2.9 to 8.6-fold with mutations.

Our results indicate that mutagenesis stringency and genomic sites are tightly coupled in determining the target-to-bystander editing ratio. Mutagenesis stringency influences the overall editing patterns while specific genomic sites dictate the mutation with the best performance. The basic rule is that relatively large mutagenesis stringency (i.e., high \( \Delta \Delta E_m \)) is needed for genomic sites with low editing selectivity, and vice versa. We tested eight genomic loci can be divided into two types in regards to the A3G3.1 editing selectivity. The first group, including EMX1 #a3, FANCE #1, and MRPL40 #1 sites, showed low selectivity, as expressed by the target-to-bystander editing ratio around 1.07–1.23; whereas the second group, including PPP1R12C #a1, ATM #1, MMS22L #1, MRPL44 #1 and FANCF #c1 sites, showed selectivity to some extent, with the target-to-bystander editing ratio ranging from 2.87 to 5.97. This natural site-dependent variation in selectivity arises from multiple sequence factors such as DNA accessibility. Therefore, the first group needs mutations with higher \( \Delta \Delta E_m \) than the second group. The theoretical model predicts that \( \Delta \Delta E_m \) follows an order of S < N (Fig. 4a). As a result, for the first group, A3G3.15 (T218N) generally performs better than A3G3.14 (T218S). In contrast, for the second group, A3G3.14 (T218S) performs better. These results indicate that mutagenesis stringency and genomic sites should be considered simultaneously during the designing process.

Currently, one difficulty in designing BE is that there are few methods to predict the editing pattern for a novel mutation before experimental validation. In addition, the same mutation can function differently at different genomic loci. Using our model, the editing patterns of those two mutations on A3G3.1 are computationally predictable, and well-validated by experiments (Fig. 4b and 4c). This result demonstrates the power of combining theoretical and experimental approaches. EMX1 #a3 site was also tested in three cell lines, K562, Jurkat, and HeLa (Supplementary Fig. 4). Although these cell lines generally have low transfection efficiency, we still observed an increase of the target-to-bystander editing ratio in A3G3.15 treated cells, compared to those treated by A3G3.1 (Supplementary Fig. 4 and Fig. 4b), for two cell lines, K562 (two-tailed \( p = 0.0005 \) with unpaired t-test) and Jurkat (\( p = 0.0001 \)). The improvement for Hela cells is less significant and needs further optimization in the future.

Discussion

In this work, we developed a theoretical framework to understand the molecular mechanisms of base editing. Our approach suggests several general rules to design BEs with improved editing selectivity. Our goal is fulfilled by modulating the binding affinity between deaminase and ssDNA using mutagenesis (\( \Delta \Delta E_m \)). The principle is to guarantee that the residence time of deaminase on ssDNA is sufficiently long to complete the editing of the first target site, while being too short for editing the second (bystander) site. Our theoretical method predicts optimal values for \( \Delta \Delta E_m \). Away from these optimal values, selectivity decreases. Therefore, instead of testing experimentally a set of candidate BE mutants, one can instead set up a computational pre-screening process by estimating the \( \Delta \Delta E_m \) of those variants, and only candidates near the optimal value can then be tested experimentally. Herein, we used alchemical free-energy calculations to estimate \( \Delta \Delta E_m \). The accuracy of this method has been validated in the A3A and A3G system (Figs. 3 and 4). Future work will help to develop carefully parameterized scoring functions for ssDNA-protein interactions or combine machine learning techniques so that the prediction of \( \Delta \Delta E_m \) can be accelerated. In addition, when estimating \( \Delta \Delta E_0 \) and \( \Delta \Delta E_m \), our model only considers the local sequence near the target base and neglects the long-range contributions from other bases in the same editing window. Because the local sequence context is a major chemical factor in determining the relative outcomes of bystander edits vs target site edits, with such approximations our model can still explain the existing experimental data (Fig. 3) and guide the design of new mutations (Fig. 4). However, the long-range contributions might serve as additional regulators and requires more detailed investigations in the future.

Equations (11–15) indicate that for a given system the editing probability is regulated by two other parameters, \( \gamma_1 \) and \( \gamma_2 \), in addition to \( \Delta \Delta E_m \). Therefore, we plotted the editing probability for different values of \( \gamma_1 \) (Fig. 5a) and \( \gamma_2 \) (Fig. 5b). We first reduced the parameter \( \gamma_1 \) (Fig. 5a) which can be achieved by increasing the on-rate of Cas9 to ssDNA. It turns out that the editing selectivity for the WT system is not affected by \( \gamma_1 \) (Fig. 5a, solid blue line vs dashed blue line at \( \Delta \Delta E_m = 0 \)) as the efficiencies of both target and bystander editing increase synchronously. However, the selectivity greatly improves when \( \Delta \Delta E_m \) is 4–6 kT, meaning that \( \gamma_1 \) amplifies the deaminase mutation regulation effect. This suggests an effective combination strategy in the design of highly selective BE: optimization of \( \Delta \Delta E_m \) first, then reducing \( \gamma_1 \) to amplify this effect. We then reduced the parameter \( \gamma_2 \) (Fig. 5b). Our calculation indicates that reducing \( \gamma_2 \) does not change the maximum editing selectivity but induces a...
Fig. 4 Engineering of A3G-BEs. a Theoretical calculation. C6 represents the target base. C5 represents the bystander. The shaded area represents the region with improved editing selectivity. 

b Experimental measurements at three genomic loci for four mutations picked by theoretical model; A3G3.1 is the full-length APOBEC3G deaminase with a set of mutations which increase the catalytic efficiency. A3G3.8, 3.9, 3.14, and 3.15 are A3G3.1 with T218G, T218I, T218S, and T218N, respectively. Bar plots represent the mean ± s.d. of three independent biological replicates.

c Experimental measurements at other five genomic loci for A3G3.14 (T218N) and A3G3.15 (T218S). Bar plots represent the mean ± s.d. of three independent biological replicates, except for the bar representing the editing efficiency of A3G3.14 at FANCF #c1 site, which shown the mean ± s.d. from four biological replicates. Source data are provided as a Source Data file.
The variants are represented by dashed lines (case 2: γ3, since γ4 comes, since 44). For example, A3A mutations N57G and Y315 greatly reduce the observed editing selectivities. In fact, even for the same BE, editing differences may vary. Therefore, each BE may require a unique optimization scheme because at different loci the neighboring bases of the bystander motif changes from equivalent mutations. In summary, a general design strategy would be (a) employing the chemical-kinetic model (Eqs. (10,13-15)) to determine the binding free energy changes required to achieve the maximum editing selectivity, ΔE peak; (b) designing mutations in the deaminase, estimating ΔΔE_m and selecting the ones near ΔE peak; (c) keeping mutations picked in the previous step and designing extra mutations that increase the binding rate of Cas9 to DNA substrate; (d) experimental validation of these changes.

Methods

Free energy calculations by molecular dynamic simulations. We utilized MDs based on chemical free-energy calculations20,21 (Fig. 2) to estimate the binding free energy changes under various conditions41. All simulations were carried out using the Gromacs package23, Amber99sb-ILDN force field with bsc0 correction for nucleic acids was used24-27. The integration time step was set to 2 fs. The initial states of the A3A-ssDNA30 and the A3G-ssDNA96 binding complex were taken from their crystal structures (PDB ID: 5ekg and 6bux, respectively). Then we used the pmx webserver30,31 to generate hybrid structures and find the most energetically favorable mutations. Each system was solvated in a cubic box with TIP3P water molecules. The ions concentration was set to 0.1 M. The dimension of the box is 9 nm. The temperature was maintained at 300 K by the Berendsen thermostat32 while the pressure was maintained at 1 atm by using the Berendsen barostat33. Electrostatic interactions were calculated by the Particle Mesh Ewald method44. The soft-core function was used for the nonbonded interactions during the allochemical transitions35. For each system, energy minimization was first performed, followed by 1 ns NVT and 1 ns NPT equilibration with the protein configuration restrained. Then the system was further equilibrated for 5 ns without any restrain. The last snapshot of the trajectory served as the starting configuration for the following allochemical transitions.

The allochemical transition (λ = 0 → λ = 1) was divided into 21 consecutive windows with the bin size of 0.05. For each window i, λ was first increased from 0 to λi (λi = 0.05, 0.1, 0.15, ... ) with a slow rate 10-8/step, then was fixed to λi for 40 ns production run. dH/dλ values were recorded every 100 steps. The free energy and error bar were estimated by Bennett’s acceptance ratio method46.

Experiment

Mammalian cell culture. HEK293T cells (American Type Culture Collection, CRL-3216) were cultured in GlutaMAX™ high-glucose Dulbecco’s modified Eagle’s medium (DMEM, Thermo, cat. 10569044). Hela (ATCC, CCL-2), K562 (ATCC, CRL-3343), and Jurkat (ATCC, TIB-152) lines were maintained in GlutaMAX™ RPMI 1640 medium (HEPES buffered, Thermo, cat. 72400146). The culture media were all supplemented with 10% fetal bovine serum (FBS, Thermo, cat. 10437028) and 100 U/mL penicillin-streptomycin (Thermo, cat. 15140122). Cells were grown in a humid atmosphere at 37 °C with 5% CO2. HEK293T and Hela cells were passaged at a ratio of 1:4 when reaching 90% confluence using TrypLE Express (Thermo, cat. 12605028). Jurkat and K562 cells were subcultured and added with fresh medium every 2 or 3 days to keep the density below 106 cells/mL. Mycoplasma testing was performed monthly using a mycoplasma PCR detection kit (abm, cat. G238).

Plasmid construction. The full-length human codon-optimized wild-type A3G with a set of mutations (P200A + N236A + P247K + Q318K + Q323K) was synthesized as gBlock and inserted into the BEmaxm construct (Addgene #112093) to replace the rAPOBEC1 region, resulting in A3G3.1. To do so, both insertion and vectors were amplified using primers with overhangs containing Esp3I recognition sites, which would generate compatible sticky ends after cutting. Then the one-pot Golden Gate assembly was employed to cut and ligate two ampli

Cell transfection, genomic DNA extraction, amplicon sequencing, and analysis. Cell transfection was performed as previously described with slight modifications15. Briefly, HEK293T or HeLa cells were seeded into a poly-D-lysine-coated 48-well plate (Corning, cat. 354090) at a density of 4.5 x 10^4 cells per well in 250 μL antibiotic-free culture medium supplemented with 10% FBS. In about 12–16 h, upon reaching 70% confluence, cells of each well were transfected. K562 and Jurkat cells were reverse-transfected at a density of 2 x 10^4 cells per well. Four cell types were all transfected with 750 ng BE plasmids and 250 ng sgRNA plasmids using 1.5 μL Lipofectamine 2000 (Thermo, cat. 11668019) dispersed in 25 μL Opti-MEM (Thermo, cat. 31985062) according to the manufacturer’s instructions. Three days later, the genomic DNAs were collected by removing the medium by aspiration or centrifugation, washing the cells gently with PBS (Thermo, cat. 10018049), and lysing the cells at 37 °C for 1–2 h with 100 μL per well of lysis buffer containing 10 mM Tris-HCl (pH 7.5), 500 mM NaCl, 0.05% SDS (Thermo, cat. 71725), and 25 μg/mL protease K (Fisher BioReagents, cat. BP1700). The cell lysates containing genomic DNA were then subjected to heat

Fig. 5 Theoretical calculations on regulating the base editing pattern of A3A-BE3. Base editing pattern of A3A-BE3 regulated by (a) γ1 and (b) γ3. The definition of ΔΔE_m, γ1 and γ3 can be found in Eqs. (10,13-15). P_a and P_b are the overall probabilities of editing the target and bystander cytidine, respectively. The difference between P_a and P_b is shown in blue. The setting with original parameters is represented by solid lines (case 1) whereas variants are represented by dashed lines (case 2: γ1 divided by five; case 3: γ3 divided by five). Source data are provided as a Source Data file.

|  |  |
|---|---|
| Ph, case 1 |  |
| Pb, case 1 |  |
| Pt, case 1 |  |
| Pb, case 2 |  |
| Pt, case 2 |  |
| Pb, case 3 |  |
| Pt, case 3 |  |
| Pt - Pb, case 1, 2 |  |
| Pt - Pb, case 2, 3 |  |

ΔEm (kJ/mol)
inactivation of the proteasine K at 80 °C for 0.5–1 h. For Sanger sequencing, the genomic DNA amplification primers and the Sanger sequencing primers are listed in Supplementary Table 2. The 20 μl PCR reactions containing 0.4 U QS High-Fidelity DNA polymerase (New England Biolabs, cat. M0491L), 0.5 μM of forward and reverse primers, and 100 ng of genomic DNA, were performed using a 35-cycle PCR program. The Sanger sequencing results were analyzed using EditR online software (https://www.baseeditr.com/).

A total of 100 ng genomic DNA was amplified at the EAMX target site by using primers attached with the partial illumina adapters and 8 bp compatible and nucleotide-balanced indices on both 5’ and 3’ end. The forward and reverse primers are as follows: for 5’-CACCTTCTTCTACGAGCTTCCTCATCCTCTGGCTATCTNNNNNNNTGTTGGTCCAGAACCAGGAG-3’; rev: 5’-GACTGGAGGTCAAGGCTGTGCTCTCCAGCTNNNNNNNCTGCTCCTGGTGGTTTT-3’.

The protospacer sequence is 5’-GAGTCGCGACGAGAAGAAGA-3’. The amplicon sequence is 5’-TGTGTTTCCAGACGCGGAGGACAACTGACAACTCAACCGACGGACGGAGGAAAGTACAAA

GCGAGAAGCTGGAGGAGGAAGGGCCTGAGTCCGAGCAGAAGAAGAAGGGCCTTGAGCTCCATCACCAGGAGGACTGGAGGTCAAGGCTGTGCTCTCCAGCTNNNNNNNCTGCTCCTGGTGGTTTT-3’.

Amplicons were pooled, column purified (Qiagen), recovered in nuclease-free water (Thermo, cat. 1077012), and quantified by the Qubit dsDNA HS assay (Thermo, cat. Q2831). A volume of 25 μl sample with the final concentration adjusted to 20 ng/μl was submitted for Ampliton-EX sequencing (GeneWiz). Fastq files were then downloaded from Genewiz ftp server and analyzed by using CRISPResso2 (https://github.com/pinelolab/CRISPResso2) to align reads and quantify the base editing efficiency and frequency.

All gblocks and primers were synthesized by Integrated DNA technologies.

Statistical analysis. All experiments were performed with 2–4 independent biological replicates. Bar plots in Fig. 4, and Supplementary Fig. 4 represent means ± standard derivation (s.d.). Bar plot in Supplementary Fig. 2a represents means ± standard error of the mean (s.e.m.).

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. All data generated or analyzed during this study are included in this published article (and its supplementary information files). Plasmids encoding A3G CBVs used in this study will be made available upon reasonable request to the corresponding authors. Targeted ampiclon sequencing data have been deposited at the Sequence Read Archive (SRA): https://www.ncbi.nlm.nih.gov/bioproject(PRJNA700693). Source data are provided with this paper.

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**Author contributions**
Q.W., X.G., and A.B.K. designed the research; Q.W. and Z.C.Z. performed the theoretical study; J.Y. and J.A.V. performed the experimental study; Q.W., J.Y., X.G., and A.B.K. analyzed the data; Q.W., J.Y., X.G., and A.B.K. wrote the paper.

**Competing interests**
The authors declare no competing interests.

**Additional information**

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**Correspondence**
and requests for materials should be addressed to Qian Wang, Xue Gao or Anatoly B. Kolomeisky.