



Fitness Effects of Single Amino Acid Insertions and Deletions in TEM-1 β -Lactamase

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Abstract

Short insertions and deletions (InDels) are a common type of mutation found in nature and a useful source of variation in protein engineering. InDel events have important consequences in protein evolution, often opening new pathways for adaptation. However, much less is known about the effects of InDels compared to point mutations and amino acid substitutions. In particular, deep mutagenesis studies on the distribution of fitness effects of mutations have focused almost exclusively on amino acid substitutions. Here, we present a near-comprehensive analysis of the fitness effects of single amino acid InDels in TEM-1 β -lactamase. While we found InDels to be largely deleterious, partially overlapping deletion-tolerant and insertion-tolerant regions were observed throughout the protein, especially in unstructured regions and at the end of helices. The signal sequence of TEM-1 tolerated InDels more than the mature protein. Most regions of the protein tolerated insertions more than deletions, but a few regions tolerated deletions more than insertions. We examined the relationship between InDel tolerance and a variety of measures to help understand its origin. These measures included evolutionary variation in β -lactamases, secondary structure identity, tolerance to amino acid substitutions, solvent accessibility, and side-chain weighted contact number. We found secondary structure, weighted contact number, and evolutionary variation in class A beta-lactamases to be the somewhat predictive of InDel fitness effects.

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Introduction

Insertions and deletions (InDels) are an important source of genetic variation in nature. They occur nearly as frequently as point mutations in some genomes [1,2]. For example, 15%–21% of polymorphisms can be attributed to short InDels in the human genome [3]. Indels can result in dramatic effects on the properties of a protein and how it evolves [2,4–6] and are known to be the cause of diseases such as cystic fibrosis and numerous types of cancer [7,8].

The metaphor of an adaptive walk across a fitness landscape works well for substitution mutations [9]. Distance can be easily measured as the number of single substitutions necessary to convert one sequence to another. However, distance is not so cleanly defined for InDels because InDels add or subtract dimensions to a protein's accessible sequence space. One way to conceptualize InDels is

that they represent a “leap” across sequence space rather than a step [10]. As such, InDels have the potential to open up new pathways for adaptation. For example, amino acid substitutions appear to be enriched around the site of InDel events in evolving proteins, either because InDel events make accumulation of substitution mutations near such sites more likely [10] or because substitutions make InDel accumulation in their vicinity more likely via “neutral roaming” [2]. This enrichment of substitutions near InDels suggests that how the surrounding protein region changes during selection may be substantially impacted by InDels.

InDels also represent a potentially underutilized source of variation in protein engineering. Although routine engineering of backbone modifications has been challenging, InDels have long been recognized as important tools for altering protein structure and properties [2,11]. Because InDels add or remove atoms from the polypeptide backbone, they can

cause major structural modifications not available through substitutions alone. They may be particularly important when seeking to dramatically change active-site structure, as they have been found to propagate long-range effects on catalytic activity [11]. However, despite their importance in nature and the laboratory, InDels remain understudied compared to substitutions.

The fitness effects of point mutations and substitutions have been extensively studied in recent years [12,13]. Previously, we comprehensively characterized the fitness effects of single amino acid substitutions in TEM-1 β -lactamase [14]. Other large-scale mutagenesis studies have been reported for over 14 proteins, characterizing the effects of single amino acid substitutions on function or fitness [13]. Such studies have advanced our understanding of the genetic code, protein structure, epistasis, and predictive models. However, we lack a similar systematic, large-scale analysis on the fitness effects of InDels.

Multiple studies have offered insight into the effects of deletions on a smaller scale. For example, a 2007 study of TEM-1 β -lactamase assayed 53 single amino acid deletions occurring throughout the protein and found that 14 (26.4%) of the variants had a minimum inhibitory concentration (MIC) that was <1% of that conferred by TEM-1, while the remaining variants varied in activity, including four that retained wild-type levels, as measured by a MIC assay [15]. The majority of debilitating deletions occurred in secondary structure elements and buried/core residues. Similarly, a 2014 study on enhanced green fluorescent protein (EGFP) characterized the tolerance to 87 random single amino acid deletions throughout the protein [16]. They found that the majority of tolerated deletions occurred in loops, while the rest were found equally distributed in helices and β -strands, with the termini of β -strands being more tolerant than the middle. Computational analysis of the EGFP found that structural properties such as relative solvent accessibility and weighted contact number (WCN) can be used to predict tolerance to deletions to some extent [17].

Insertion studies are even more limited, generally examining only a few rationally chosen insertion sites in a protein. For example, 2006 study in TEM-1 assessed the impact of random peptide insertion into three loops and found that tolerance depended largely on the insertion site [15]. Based on their findings, they also suggested that tolerance to insertions was not well correlated to tolerance to substitutions in the same region.

While these studies provide important insights into the effects of InDels, they are limited by their scale. Here, we present a near-comprehensive analysis of the fitness effects of single amino acid InDels in TEM-1 β -lactamase, a widely studied antibiotic resistance protein. We find that while InDels are

largely deleterious compared to substitutions, partially overlapping regions of tolerance to InDels exist throughout the protein.

Results and Discussion

Fitness effects of single amino acid InDels

TEM-1 β -lactamase is a commonly studied protein and convenient model for protein evolution experiments. It confers high resistance to penicillin antibiotics, such as ampicillin, which can be used as a proxy for protein fitness [14,18,19]. We use our previously described band-pass, MIC-like approach for measuring antibiotic resistance in a high-throughput, high-resolution manner [14,20,21]. This method uses a synthetic biology approach to quantify Amp resistance as a proxy for fitness. Unlike growth competition experiments and standard MIC assays, the fitness measures are ampicillin concentration independent and low fitness values are as precisely measured as high fitness values.

We focused on in-frame insertions or deletions of three nucleotides. We did not study insertions or deletions that are one or two nucleotides in length, as such mutations are frame-shifting mutations with drastic changes to protein sequence and nearly always inactivate proteins, although such InDels can be bypassed by transcriptional or translational slippage to give full-length functional proteins [22]. We did not study three-nucleotide insertions or deletions that are out of frame, as they cause substitutions in the amino acid sequence in addition to the amino acid insertion or deletion. We wanted to be able to isolate the effect of the single amino acid insertion or deletion away from any substitution effects.

We used inverse-PCR to create a plasmid library designed to code for every possible single amino acid insertion (5720 variants) and every possible single amino acid deletion (286 variants) in TEM-1. For insertions, we used degenerate primers in which the 5'-end of the forward primer had an additional (NNN) sequence. For deletions, we used primers in which the 5'-end of the forward primer had a 3-base-pair deletion. We transformed SNO301 *Escherichia coli* cells with each library of InDel alleles and plated on LB plates supplemented with tetracycline and 13 different Amp concentrations ranging from 0.25 to 1024 μ g/ml. Amp prevents growth if the Amp concentration is too high relative to the amount of Amp resistance conferred. Tetracycline prevents growth if the concentration of Amp is too low relative to the amount of Amp resistance conferred. This behavior is the key feature of the band-pass gene circuit in SNO301 cells. As a result, a particular allele will confer growth only in a narrow range of Amp

concentrations (see Firnberg *et al.* [14] for a detailed explanation), and the higher the midpoint of that range, the fitter the allele. After incubation at 37 °C overnight, we recovered the 13 sublibraries and performed deep-sequencing to determine how often each allele appeared on each plate. Sequencing reads of alleles containing synonymous codon insertions were grouped together, with the exception of the stop codons. The amber (UAG) stop codon exhibits nonsense suppression in SNO301 *E. coli* via the *supE44* tRNA allele, which results in glutamine incorporation at UAG codons with variable efficiency depending on the nucleotides immediately flanking UAG [14]. To avoid convolution, we included only non-amber stop codons in our analysis. The reported fitness values are calculated as the Amp concentration at which the mutant allele appeared most frequently relative to the wild-type allele (see [Materials and Methods](#) for a more detailed description).

We obtained fitness values for 77.9% (4457/5720) of possible amino acid insertions and 97.9% (280/286) of possible amino acid deletions in TEM-1 (Fig. 1). As expected, we find that InDels are largely deleterious. Over half of insertions (51%) and deletions (59%) resulted in at least a 100-fold decrease in fitness relative to TEM-1. In contrast, only 9.8% of insertions and 11% of deletions retained 50% of wild-type fitness, although close to half (40.9%) of these were in the signal sequence, which is cleaved and not part of the mature protein. Although we measured 74 InDels alleles with fitness values nominally greater than 1, only 27 were statistically different than 1. However, we suspect that many of these are not actually beneficial insertions. For example, the highest “significant” beneficial insertions cluster in the region 257–272. The insertions W and K at 257, I and R at 261, T at 267, A at 271, and T and D at 272 all have a peculiar distribution of sequencing counts that is defined by an abnormally high number of sequencing count on the plate with 512 $\mu\text{g/ml}$ Amp with comparatively very few counts on 256 $\mu\text{g/ml}$ and no counts on 1024 $\mu\text{g/ml}$. The abnormally high counts on just one plate suggest an artifact. For this reason, we focus our attention on broad patterns in our fitness effects rather than the fitness values associated with particular InDels.

Relationship between InDel Fitness Effects and TEM-1 Sequence/Structure

Visual examination of the heatmap depicting the fitness effects (Fig. 1) suggests a higher tolerance to InDels outside of secondary structures. It also appears that the fitness effect of an insertion depends more on the site of the insertion than on the amino acid identity. To examine this quantitatively, we looked at the distribution of mean fitness

values per position and compared it to the distribution of mean fitness values grouped by amino acid (Supplementary Fig. 1). We found that the mean fitness effects per position have a wider distribution of values than the mean fitness effects grouped by the amino acid inserted ($P = 0.009$, Brown–Forsythe test).

Examining the median fitness of alleles containing insertions and the fitness of alleles containing deletions across TEM-1, we observed “hot spots” of tolerance for InDels in TEM-1 (Figs. 2 and 3). The pattern suggests some correlation between where InDels are tolerated and indicates higher tolerance in the signal sequence and in unstructured regions of the protein. Higher tolerance to InDels in loops compared to helices and strands is widely observed across many families of proteins [23]. Our results also agree with previous observations in TEM-1 in particular. For example, visual examination of Figs. 1 and 2 suggests a notable tolerance to insertions in the loop connecting the final β -strand to the C-terminal helix, which is a location previously found to be broadly tolerant to random sequences of insertions [15].

We also examined the relationship between evolutionary variations in class A β -lactamases and InDel fitness effect in TEM-1. We aligned a published set of 157 class A β -lactamase sequences from different bacterial species (including TEM-1) [24] by progressive multiple alignment using a Gonnet scoring matrix in MATLAB. We chose this data set because it consists of close homologs of TEM-1 where there were enough examples of InDels, but the size of the InDels was general small (our analysis found 77% of the InDels were single amino acids and 94% were not more than 2 amino acids). We wanted to compare to homologs with small InDels, because our fitness measurements were of single amino acid InDels. We identified the positions at which these 157 sequences [24] contained an insertion or deletion relative to TEM-1. We find that these positions generally overlap insertion-tolerant regions in TEM-1, but several regions in TEM-1 that tolerate insertions and especially deletions are not observed in natural class A β -lactamases, at least in this data set (Fig. 2).

We find that the 23-amino-acid signal sequence is the most InDel-tolerant region in TEM-1 (Fig. 4). This sequence directs TEM-1's export to the periplasm via the Sec export pathway. The signal peptide is removed upon export to the periplasm and is not part of the mature protein. Presumably, mutations in the signal sequence affect fitness through changes of TEM-1's export efficiency to the periplasm. The signal sequence of TEM-1 is also the most tolerant region to missense mutation [15]. This tolerance is consistent with the loose sequence constraints for Sec-dependent signal sequences and its lack of secondary structure elements [25].

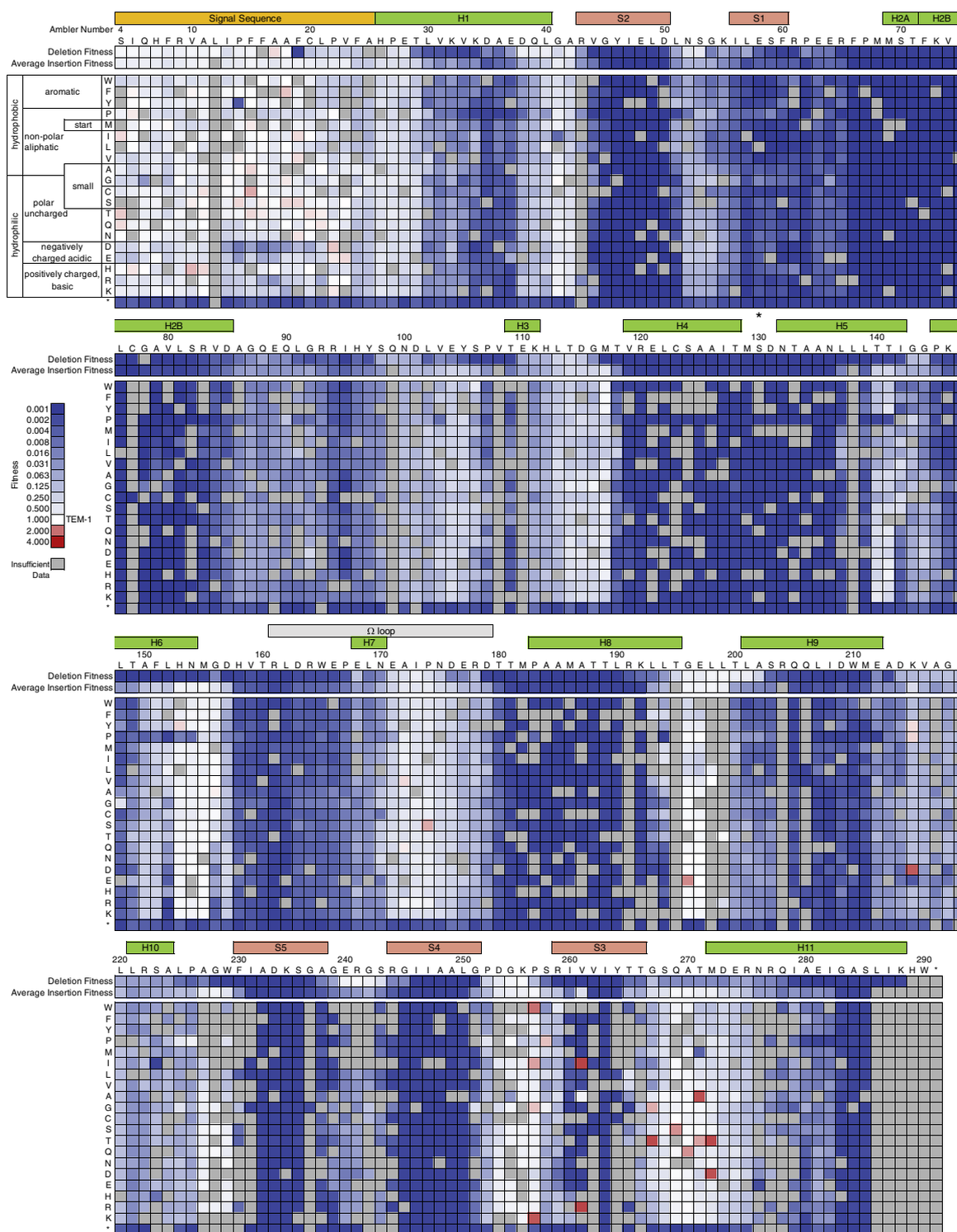


Fig. 1. The fitness effects of amino acid InDels in TEM-1. The heatmap indicates relative fitness values as calculated based on ampicillin resistance. Insertion position is defined by the new position of the inserted amino acid (e.g., an insertion denoted at position 50 was inserted between residues 49 and 50 in TEM-1). Ambler consensus numbering for beta-lactamases is used. The signal sequence (yellow), α helices (green), β strands (orange), Ω loop (gray), and active sites (*) are indicated. Tabulated sequencing counts and fitness data are provided as Supplementary Data S1 (insertions) and S2 (deletions).

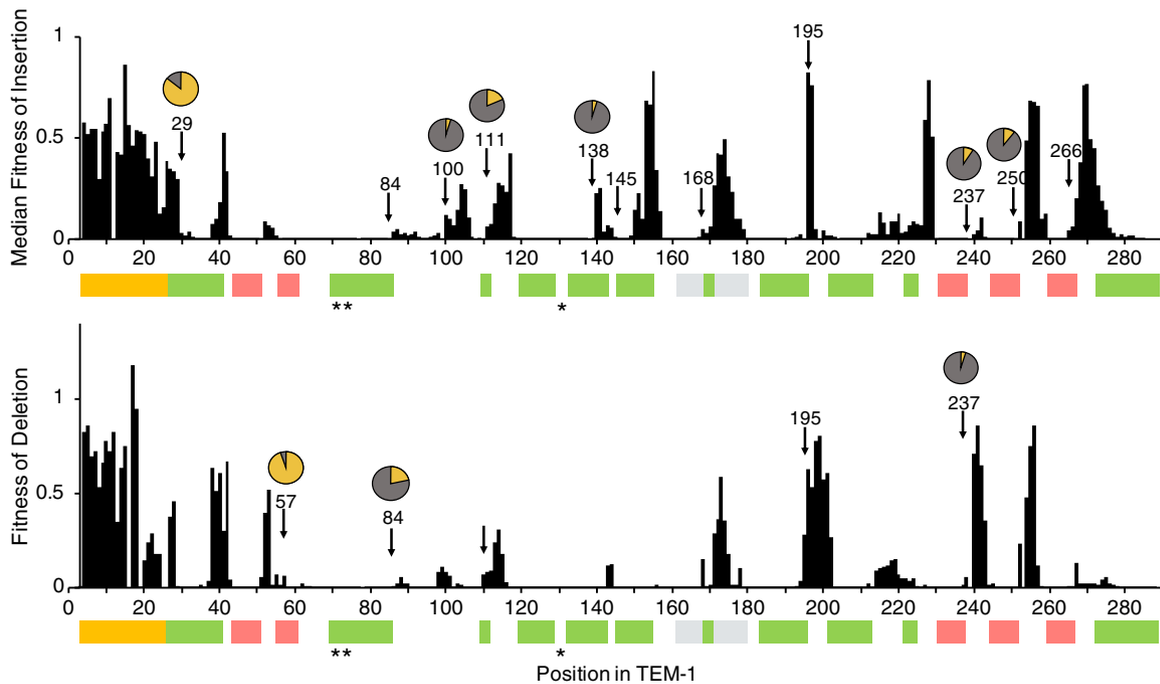


Fig. 2. Fitness of TEM-1 containing InDels as a function of primary sequence. Median fitness values are presented for insertions. Arrows indicate positions at which other class A β -lactamases contain an insertion or deletion (based on a multiple sequence alignment of 156 class A β -lactamase and TEM-1). Pie charts indicate in yellow the fraction of sequences out of 156 that contain an insertion (top chart) or deletion (bottom chart) at that position. Pie charts are omitted for fractions less than 3%. The colored bars indicate the signal sequence (yellow), α helices (green), β strands (pink), Ω loop (gray), and active sites (*).

In the mature protein, helices and strands are the least tolerant to InDels. For both InDels, the mean fitness of mutant alleles in loop regions is higher than in secondary structure elements ($P < 0.0001$ for insertions, $P < 0.001$ for deletions, Student's *t*-test). This is not surprising given that backbone modifications can cause structured regions to fold incorrectly and have dramatic effects on the protein [26]. However, we found some exceptions to this overall pattern. For example, the loop region between β -strand S1 and α -helix H2A shows no tolerance for insertions or deletions. We also found that 2.9% (51/1765) of insertions in α -helices, often at the ends of the structure element, resulted in less than a 50% decrease in fitness.

Tolerance to insertions was greatest at the ends of both α -helices and β -strands and decreased as one moved into the structural element (Fig. 5). The median fitness for an insertion immediately before or after an α -helix was 4.3-fold to 28-fold higher than the median values for insertions two to eight residues from the end ($P < 2 \times 10^{-5}$ for all pairwise comparisons, Mann-Whitney). When the inserted residue was one residue away from the end of the α -helix, the median fitness value was 2.6- to 17-fold higher than that for insertions located three to eight residues from the end ($P < 0.0006$ for all pairwise comparisons, Mann-Whitney). A notable exception to this trend was the 17-residue C-terminal α -helix H11, the longest α -helix

in the protein, which showed significant tolerance for the first 11 positions before abruptly dropping to complete intolerance. The median fitness values for insertions immediately before or after a β -strand were 8- to 16-fold higher than the median values for insertions within a β -strand ($P < 2 \times 10^{-14}$ for all pairwise comparisons, Mann-Whitney). When the inserted residue was one residue away from the end of the β -strand, the median fitness value was 1.8-fold higher than those for insertions at other locations in the β -strand ($P < 2 \times 10^{-9}$ for all pairwise comparisons, Mann-Whitney).

To more specifically examine the difference between tolerance to insertions and deletions, we calculated the ratio of the mean fitness of alleles with insertions to the fitness of an allele with a deletion at each position across TEM-1 (Fig. 6). Overall, we find more regions where insertions are preferred over deletions, but a few regions where deletions are preferentially tolerated. For example, the C-terminal α -helix is dominated by preference to insertions, while the N-terminal α -helix contains several positions where deletions are relatively preferred. The extended loop between β -strands S5 and S4 (G238–S243) was the region that most preferred deletions over insertions (Fig. 6). Deletions relative to TEM-1 in this region are observed evolutionarily, although they are uncommon (Fig. 2).

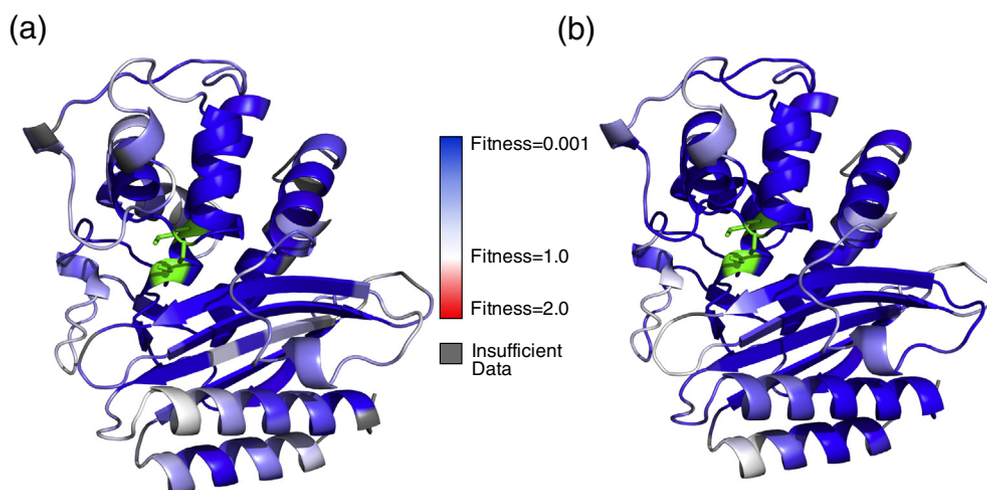


Fig. 3. InDel fitness mapped onto TEM-1 structure. (a) TEM-1 secondary structure colored by median fitness of insertions. Positions for which we obtained fewer than 4 fitness values are indicated in gray. (b) TEM-1 secondary structure colored by fitness of deletions. In both figures, the active site residues are colored in green. No mean fitness values > 1 are observed.

We also examined the distribution of fitness effects of InDels compared to substitutions (measured in our previous study [14]). Unsurprisingly, we found that InDels were more often deleterious than substitutions (Supplementary Fig. S2). The distributions of InDel fitness values were similar, and the mean fitness of alleles containing an insertion (0.14 ± 0.26) was not significantly different than the mean fitness of alleles containing a deletion (0.11 ± 0.23) (Supplementary Fig. S2).

To explore the comparison between InDels further, we examined the correlation between the mean fitness of alleles with an insertion at a given position and the fitness of an allele with a deletion at the corresponding position (Supplementary Fig. S3a) and found a weak correlation ($R^2 = 0.32$). We also compared the mean fitness change of an insertion of a given amino acid against the mean fitness change of a deletion of the same amino acid and found almost no correlation ($R^2 = 0.07$) (Supplementary Fig. S3c). This further

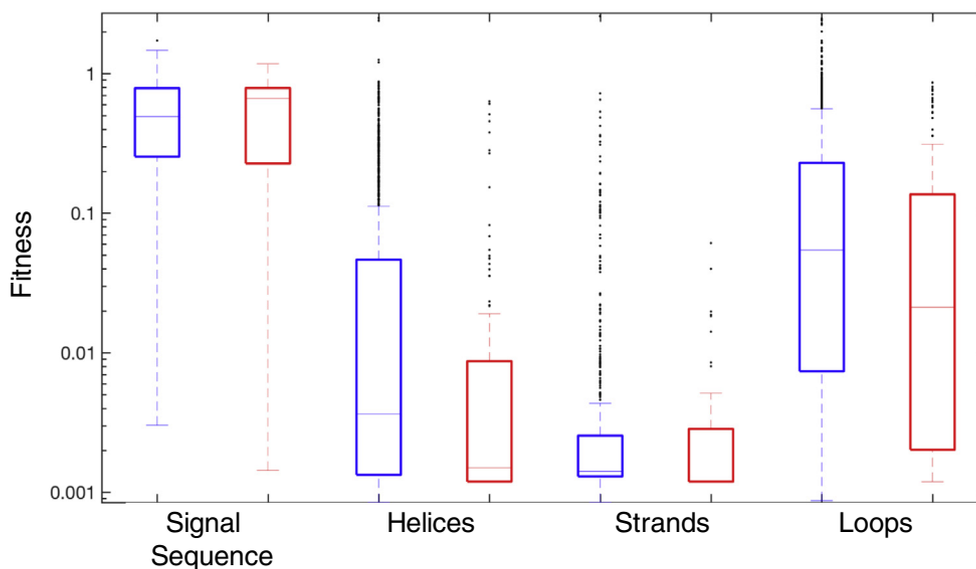


Fig. 4. Relationship between InDel fitness and secondary structure. Box plots of fitness values for insertions (blue) and deletions (red) are shown for the signal sequence and secondary structure elements. The central line indicates the median, and the bottom and top edges of the box indicate the 25th and 75th percentiles. The median fitness value for deletions in strands is at the 25th percentile and therefore not visible on the plot. The whiskers extend to the most extreme data points not considered outliers, which are represented by circles. Outliers are defined as values more than 1.5 times the interquartile range away from the top or bottom of the box.

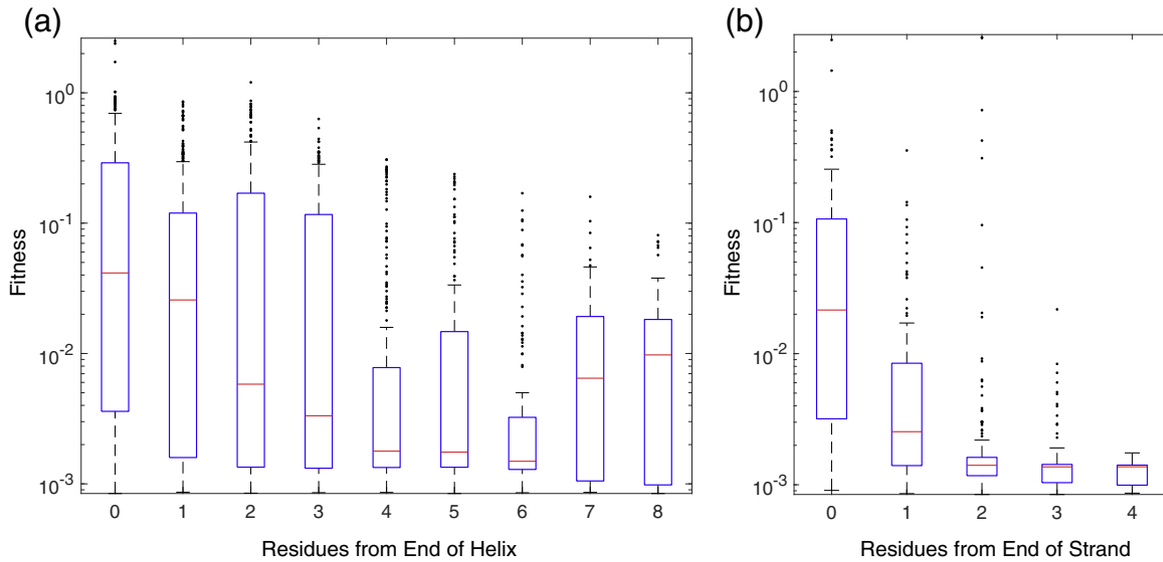


Fig. 5. Relationship between insertion fitness and distance from the closest end of secondary structure element. (a) Box plots of fitness values for insertions within helices. (b) Box plots of fitness values for insertions within strands. The number 0 corresponds to insertions immediately before or immediately after the structure element, the number 1 refers to insertions after the first or before the last residue in the structure element, and so on.

indicates that the location of the InDel is more predictive than the identity of the amino acid inserted or deleted.

Next, we examined the correlation between fitness values when comparing insertions and substitutions. Specifically, we wondered if the fitness effect of an amino acid inserted before position N would correlate to the fitness effect of having position N mutated to the same amino acid. In this comparison, we included only fitness values of insertions at positions with a mean fitness of ≥ 0.1 . We do this to account for the predominance of insertions that result in complete loss of function. By excluding those positions, we instead ask the question: where insertions are tolerated to some degree, what is the correlation between the effects of insertions and substitutions? We find no significant correlation when we compare insertions to substitutions at the corresponding position ($R^2 = 0.07$) (Supplementary Fig. S3b); however, the mean fitness change of an amino acid substitution is somewhat predictive of the mean fitness effect of the same amino acid insertion ($R^2 = 0.39$) (Supplementary Fig. S3d). For example, the two least tolerated amino acid insertions (Pro and Trp) are also the least tolerated substitutions, and the two most tolerated insertions (Ser and Thr) are among the most tolerated substitutions (Supplementary Fig. S3d).

We further explored TEM-1's tolerance to insertions by determining the effective number of amino acid insertions at each position (k_{INS}^*). This measure is analogous to a measure of substitution tolerance (k^*) that derives from information-theoretical entropy, which was originally proposed to quantify the variability at a given position in a set of aligned sequences [27]. As we showed previously, k^* can be

adapted to quantify the tolerance of substitutions based on measured fitness values [14]. For substitutions, a k^* value of 1 indicates a position at which all missense mutations result in complete inactivation of the protein, and a k^* value of 20 indicates that all amino acid substitutions result in the same fitness as wild type. Here, we define a similar measure for insertions (k_{INS}^*), which includes the possibility of no insertion (i.e., wild type) in the distribution of protein fitness values at each position (Eqs. (1)–(4)).

$$k_{\text{INS}}^* = \frac{21k_{0,\text{INS}}^*}{n} \quad (1)$$

$$k_{0,\text{INS}}^* = 2^S \quad (2)$$

$$S = -\sum_{i=1}^k p_i \log_2 p_i \quad (3)$$

$$p_i = \frac{w_i}{\sum_{j=1}^n w_j} \quad (4)$$

A k_{INS}^* value of 1 indicates a position at which no amino acid insertion is tolerated (i.e., the fitness values of all amino acid insertions are zero) and a k_{INS}^* value of 21 indicates a position at which all insertions retain wild-type fitness values.

Over 30% of positions do not tolerate a single amino acid insertion of any kind ($k_{\text{INS}}^* < 2.0$) (Fig. 7a). The peak in the distribution of k_{INS}^* values between 17 and 20 indicates that there is a fraction of positions (19.3%) for which most insertions are

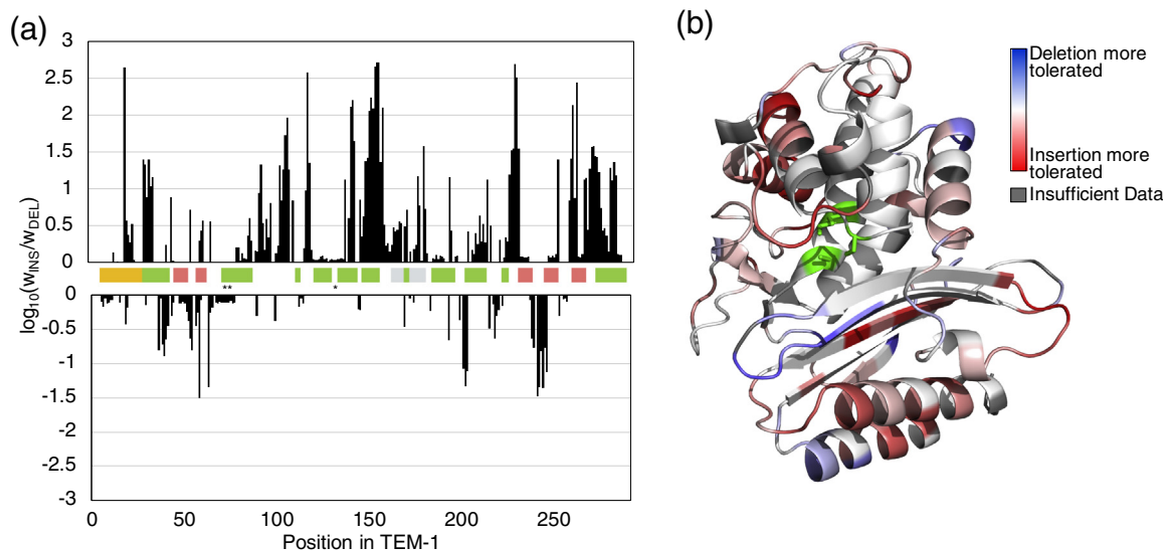


Fig. 6. Differences in tolerance to InDels across TEM-1. (a) The \log_{10} of the ratio between mean fitness of insertions and the fitness of a deletion at each position across TEM-1. The colored bars indicate the signal sequence (yellow), α helices (green), β strands (pink), Ω loop (gray), and active sites (*). (b) TEM-1 structure colored by the same ratio values. Blue indicates positions with higher tolerance to deletions, white indicates the same tolerance to both InDels, and red indicated higher tolerance to insertions.

well tolerated. However, there are no positions for which every inserted amino acid retains wild-type fitness ($k^*_{INS} = 21$).

All 23 positions in the signal sequence had k^*_{INS} values above 13, but 5 positions had a k^* for substitutions less than 13 (Fig. 7a). In the entire protein, a position's tolerance for insertion, as measured by k^*_{INS} , weakly correlated its tolerance for substitutions (Fig. 7b). We found that tolerance to insertions correlates weakly with distance from the active site (Fig. 7c). Positions less than 10 Å away from the active site are almost completely unaccepting of insertions. We observed a slightly stronger correlation between k^*_{INS} and percent solvent accessible surface area, with buried residues being less amenable to insertions (Fig. 7d). We found that side-chain WCN, a measure of how densely packed a residue is [28], best predicts how well an insertion is tolerated (Fig. 7e), although the R^2 is only 0.27. WCN was also the single best predictor of whether a deletion is tolerated in eGFP [17]. The lack of simple correlations between properties and insertion fitness were consistent with our expectation that the fitness effects of insertions have complex origins. Analogous correlations between these properties and substitution mutations were notably stronger, with ~2-fold higher R^2 values [14].

Conclusions

Our analysis of InDels in TEM-1 provides the first systematic and near-comprehensive study of their fitness effect on a single protein and insight into a common yet understudied source of genetic

variation. We found InDels to be largely deleterious, although regions of tolerance were observed, particularly in unstructured regions of the protein and at the ends of helices and strands. While regions of tolerance to InDels partially overlapped, we found that most regions of the protein tolerated insertions more than deletions. Of the measures we examined, we found secondary structure, WCN, and evolutionary variation in class A beta-lactamases to be somewhat predictive of InDel fitness effects. A broader understanding the fitness effects of InDels and how they relate to structural properties should allow for more informed protein engineering strategies, more robust computational prediction of protein structure, and a deeper understanding of the role that different types of mutations play in protein evolution.

Materials and Methods

Insertion library creation

The TEM-1 gene was expressed on pSkunk2, a 4.36-kb plasmid containing spectinomycin resistance and the *p15* origin of replication, under the IPTG-inducible *tac* promoter in *E. coli*. We used inverse PCR with oligo primers (IDT) designed to create every possible single amino acid insertion in TEM-1, using primers with a degenerate nucleotide (NNN) sequence on the 5' end of the forward primer and a compatible reverse primer designed for each position. PCR products were visualized using gel electrophoresis, to confirm the creation of a

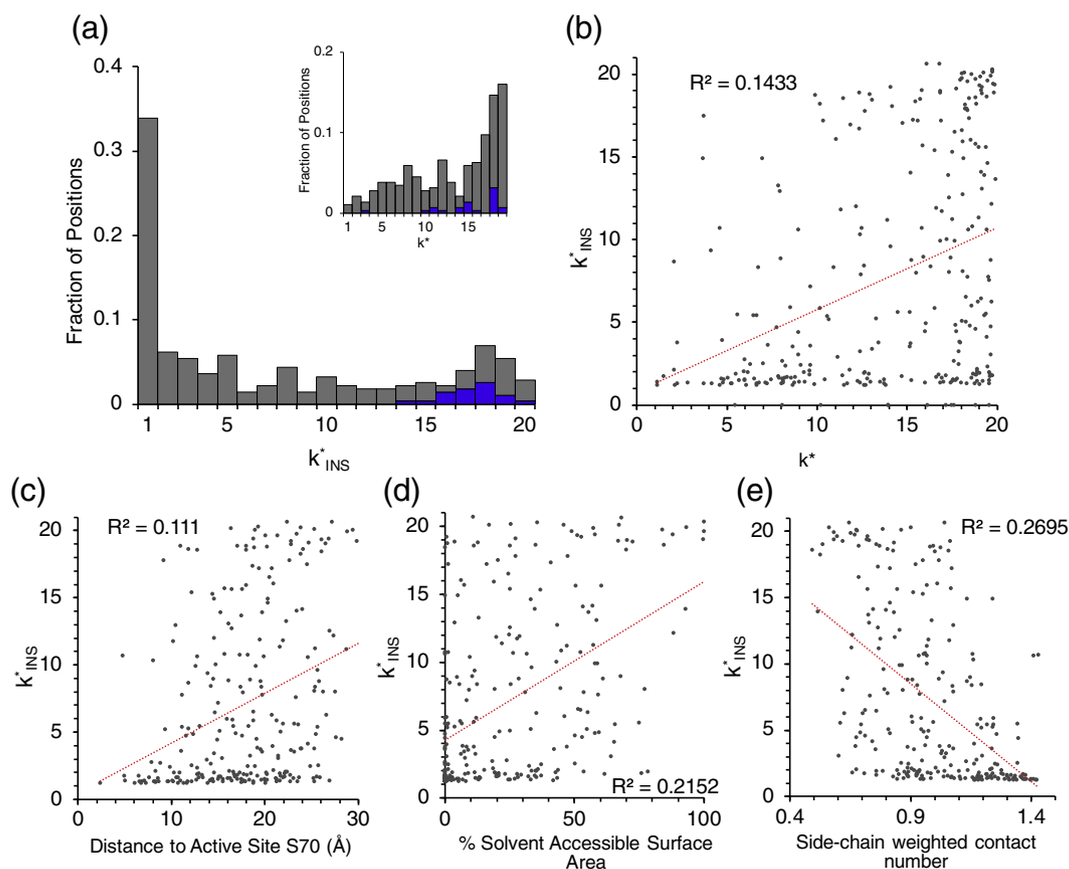


Fig. 7. Determinants of tolerance of TEM-1 to amino acid insertions. (a) The distribution of k_{INS}^* values in TEM-1. k_{INS}^* values for the mature protein are colored in gray, and k_{INS}^* values for the signal sequence are colored in blue. The inset shows the corresponding distribution of k^* values for substitutions [14]. (b) Correlation of k_{INS}^* with k^* of substitutions [14]. (c) Correlation of k_{INS}^* with distance from the active site. (d) Correlation of k_{INS}^* with percent solvent accessibility surface area. (e) Correlation of k_{INS}^* with side-chain WCN.

linearized plasmid product at each of the 286 positions. We were unable to create a product for a small number of positions, despite troubleshooting efforts. We pooled the PCR products, creating a library for each third of the gene, to be compatible with Illumina MiSeq 2×300 bp sequencing. We isolated the ~ 4 -kb band from an agarose electrophoresis gel for each third, phosphorylated the DNA at 37 °C (NEB T4 PNK), and ligated it overnight at 16 °C. NEB 5- α F' lacIq *E. coli* were transformed with the ligation product and plated on LB-agar plates containing 50 μ g/ml spectinomycin and 2% glucose. At least 500,000 transformants were obtained for each library (i.e., each third of the gene).

We recovered each library from the plate in LB media and isolated the plasmid library. We transformed electrocompetent SNO301 *E. coli* cells with each library and plated on LB-agar plates containing 50 μ g/ml spectinomycin, 50 μ g/ml chloramphenicol, and 2% glucose. At least 100,000 transformants were obtained from each third. We recovered each library from the plate in LB media and made glycerol stocks.

Deletion library creation

The deletion library was made in the same way as the insertion library with a few exceptions. The forward primer for inverse-PCR contained a 3-bp deletion on the 5' end, to create a deletion at every position in TEM-1. The same reverse primers were used. The deletion library was not created in thirds, as it was subsequently sequenced using PacBio, which can accommodate longer reads.

Selection and sequencing

High-throughput selection for resistance to ampicillin (Amp) was performed using a band-pass genetic circuit, described in previous work [14]. Briefly, *E. coli* SNO301 cells containing each library were plated on LB-agar plates containing 20 μ g/ml tetracycline and 13 different Amp concentrations, ranging from 0.25 to 1024 μ g/ml, in 2-fold increments. Plates were incubated for 21 h at 37 °C. Each library was plated in triplicate on each Amp concentration and

the CFUs from each plate were counted to determine the frequency of colonies appearing on each plate. Based on these counts, a proportional amount of DNA from each plate was deep sequenced. For the insertion library, barcoded amplicons were prepared by recovering the cells from each selection plate, isolating the plasmid DNA, and performing PCR with appropriate primers as described previously [14,20]. Barcodes to identify each plate and adapters compatible with Illumina MiSeq platform were added in this PCR step. Amplicons were pooled and sequenced using Illumina MiSeq with 300-base-pair, paired-end reads. For the deletion library, we recovered cells from each selection plate, isolated the plasmid DNA, linearized it with the *SphI* restriction enzyme, and separately sequenced each of the 13 linearized plasmid libraries using PacBio.

Data analysis

The de-multiplexed MiSeq reads and the PacBio reads were analyzed using custom MATLAB scripts. For MiSeq reads, paired-end reads were trimmed and concatenated to yield full-length reads. Each read was then aligned to *TEM-1* using a Smith–Waterman algorithm with the lowest possible gap opening penalty of 1 and a gap extending penalty of 0.1. Reads with an alignment score lower than 100 were filtered out and only reads containing a single amino acid insertion (or deletion) were used for analysis. Fitness was calculated for each unique InDel mutant based on the counts from each plate (Amp concentration). For insertions, synonymous codons were grouped together and total counts were used to calculate the single amino acid fitness. Amber codons (UAG) were excluded from the stop codon analysis.

For each allele, counts were first adjusted based on the number of sequencing reads obtained from each plate relative to the CFUs observed on that plate, as described previously [20]. Detailed description of the fitness calculation can be found in our previous studies [14,20], which we followed with just a few differences. For the insertion library, we excluded alleles with fewer than 20 counts and alleles with a maximum single plate count less than one third the total count. For the deletion library, we excluded alleles with fewer than 5 counts. We used a lower cutoff for the deletion library because we used PacBio sequencing, which sequenced the entire *TEM-1* gene. Thus, the sequencing data for deletions with low counts could not have artifactual data arising from *TEM-1* base-substitution mutations arising spontaneously or during library creation, unlike the analogous data for the insertion library sequenced by MiSeq. Inspection of the sequencing counts per plate for the 17 deletions with between 5 and 20 counts supported the use of this lower threshold. Supplemental Data S1 and S2 contains sequencing counts and fitness values.

For each allele (i), the plate with the highest adjusted counts and the four plates on either side (i.e., two plates with higher Amp and two plates with lower Amp) were used to calculate an unnormalized fitness value, representing the midpoint resistance to Amp:

$$f_i = \frac{\sum_{p=1}^{13} c_{i,p} \log_2(a_p)}{\sum_{p=1}^{13} c_{i,p}} \quad (5)$$

where $c_{i,p}$ is the adjusted count of allele i on plate p , and a_p is the Amp concentration on plate p (in $\mu\text{g/ml}$). The reported fitness values are normalized to wild-type *TEM-1*:

$$w_i = \frac{2^{f_i}}{2^{f_{\text{TEM-1}}}} \quad (6)$$

Wild-type fitness was calculated in the same way (i.e., using adjusted sequencing counts) and verified separately by plating wild type in triplicate during the bandpass selection step. Both colony counts and sequencing counts revealed a midpoint Amp resistance of $\sim 215 \mu\text{g/ml}$.

Error in fitness (σ_{w_i}) was estimated via Eqs. (7) and (8), using our previously determined correlation between sequencing counts (n_i) and the standard deviation of the difference in fitness between synonymous alleles [14].

$$\sigma_{w_i} = w_i \times e_i \quad (7)$$

where e_i , the upper-level estimate of the fraction error in fitness, is given by:

$$e_i = 0.667 n_i^{-0.387} \quad (8)$$

Fitness values were determined to be significantly different than 1 if they were greater or less than 1 by twice the error estimate.

CRedit authorship contribution statement

Courtney E. Gonzalez: Formal analysis, Investigation, Writing - original draft, Writing - review & editing. **Paul Roberts:** Investigation. **Marc Ostermeier:** Conceptualization, Formal analysis, Supervision, Writing - review & editing.

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Author Contributions: C.E.G. performed all experiments except some initial test libraries, which were constructed by P.R. C.E.G. and M.O. conceived and designed the experiments, analyzed the data, and wrote the paper.

Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jmb.2019.04.030>.

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InDels, insertions and deletions; MIC, minimum inhibitory concentration; EGFP, enhanced green fluorescent protein; WCN, weighted contact number.

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