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Asymmetric mutations in the tetrameric R67 dihydrofolate reductase reveal high tolerance to active-site substitutions

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Abstract: Type II R67 dihydrofolate reductase (DHFR) is a bacterial plasmid-encoded enzyme that is intrinsically resistant to the widely-administered antibiotic trimethoprim. R67 DHFR is genetically and structurally unrelated to *E. coli* chromosomal DHFR and has an unusual architecture, in that four identical protomers form a single symmetrical active site tunnel that allows only one substrate binding/catalytic event at any given time. As a result, substitution of an active-site residue has as many as four distinct consequences on catalysis, constituting an atypical model of enzyme evolution. Although we previously demonstrated that no single residue of the native active site is indispensable for function, library selection here revealed a strong bias toward maintenance of two native protomers per mutated tetramer. A variety of such “half-native” tetramers were shown to procure native-like catalytic activity, with similar K_M values but k_{cat} values 5- to 33-fold lower, illustrating a high tolerance for active-site substitutions. The selected variants showed a reduced thermal stability ($T_m \sim 12^\circ\text{C}$ lower), which appears to result from looser association of the protomers, but generally showed a marked increase in resilience to heat denaturation, recovering activity to a significantly greater extent than the variant with no active-site substitutions. Our results suggest that the presence of two native protomers in the R67 DHFR tetramer is sufficient to provide native-like catalytic rate and thus ensure cellular proliferation.

Keywords: trimethoprim resistance; combinatorial mutations; thermostability; active site engineering; homotetramer

Abbreviations: CD, circular dichroism; DDM, R67 DHFR mutant library, doubly mutated; DHF, dihydrofolate; DHFR, dihydrofolate reductase; DNM, R67 DHFR dimer, non-mutated; DSM, R67 DHFR mutant library, singly mutated; THF, tetrahydrofolate; TMP, trimethoprim.

Additional Supporting Information may be found in the online version of this article.

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Introduction

Dihydrofolate reductases (DHFRs) play a critical role in proliferation of all living cells. They catalyze the reduction of dihydrofolate (DHF) to tetrahydrofolate (THF), using NADPH as the hydride-donating cofactor. Among its metabolic roles, THF is required for the biosynthesis of purines and thymidylate. As a result, inhibition of DHFR leads to cell death, making this enzyme a main target of antibiotic and anticancer drugs over the past six decades.

Trimethoprim (TMP) is a low-cost and effective bacteriostatic antibiotic that is used worldwide in the prophylaxis and treatment of bacterial infections, both

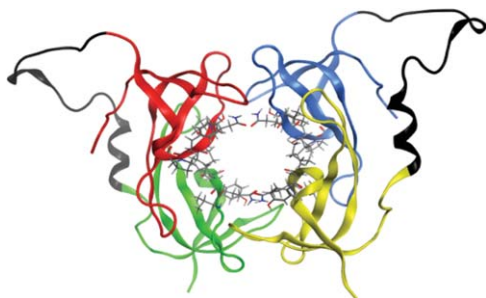


Figure 1. R67 DHFR: The four protomers (A–D) are illustrated in red, green, blue and yellow, respectively. The active site residues 66–69 are shown in sticks representation. The linker introduced for the dimer construct and the *N*-terminal tail of R67 connecting protomers A,B and C,D are shown in black. The linker conformation was predicted using the loop-modelling tool of the MOE software.⁵

in humans and livestock.¹ It is an effective competitive inhibitor of the chromosomally-encoded DHFR in many bacteria and parasites; for example, it inhibits *E. coli* DHFR with $K_i = 0.4$ nM.² Its widespread use has resulted in the development of microbial resistance to TMP, in the form of highly transmissible plasmid-encoded DHFRs. TMP-resistant DHFRs (type I–XVII) are classified into two gene families: *dfrA* and *dfrB*.^{3,4} The *dfrA* family DHFRs are close genetic variants of the monomeric chromosomally-encoded DHFRs that are 160–190 amino acids long. In contrast, *dfrB* genes encode shorter sequences, on the order of 78 amino acids, that homotetramerize to form the active DHFR species. They are genetically and structurally unrelated to chromosomal DHFRs. Type II R67 DHFR belongs to this second class. Its single, symmetrical active-site pore characterized by 222 symmetry lies at the heart of a channel formed by the assembly of four protomers that are largely constituted of a SH3-like β -sandwich fold (Fig. 1). Short loops and both termini extend out of each β -sandwich, forming the inter-protomer interfaces.^{6–8} As R67 DHFR is unrelated to its chromosomal counterparts, it has a different catalytic mechanism^{4,9–13} as well as a different response to antifolate inhibitors.⁴ It has been qualified as being “primitive,” in that its catalytic strategy lacks in binding specificity, suggesting that it may not yet be evolutionarily optimized.¹⁴ Importantly, as a result of being unrelated to *dfrA* family DHFRs, it confers resistance to high concentrations of TMP ($K_i = 0.15$ mM).¹⁵

Here, we investigate the tolerance to active-site modification of this compact reductase. One β -strand per protomer (V66, Q67, I68, and Y69) constitutes the major portion of the active site channel (Fig. 1). The DHF substrate and NADPH cofactor have been proposed to slip into the channel from opposite ends to juxtapose their reactive atoms at the center of the channel, each forming contacts mainly with one protomer, though additional stabilizing contacts are contributed by the three remaining protomers.^{16,17}

For this reason, residues involved in ligand binding can have up to four different consequences within a single binding/catalytic event.⁹ This makes the active site poorly tolerant to point mutations because of a cumulative effect of each fourfold repeated residue. Despite this observation, we have previously demonstrated that the homotetrameric active site can be heavily modified, including up to four simultaneous mutations per protomer (16 in the tetrameric active site).¹³ In that selection experiment, the vast majority of modified homotetrameric active sites (99.8% of the variants tested) did not give rise to native-like function, yet this and other studies showed that the simultaneous variation of more than one residue in R67 DHFR can provide compensatory effects, thus allowing multiple mutations to complement each other.^{13,18–20} This study also revealed that no single residue of the native active site is indispensable for function.

Because our previous active-site mutation efforts using the monomeric species resulted in a low yield of functional variants, here we investigate the tolerance of the active site to more conservative modifications. To reduce the complicating effect of the active-site symmetry, R67 DHFR has previously been genetically linked to provide a covalent dimeric or tetrameric species, using the unstructured *N*- and *C*-termini to provide an inter-domain linker (Fig. 1).^{19,21} This breaks the active-site symmetry by introduction of substitutions on only one (for the covalent tetramer) or two (for the covalent dimer) of the four protomers.^{18,19,22} In this work, we have modified the active site in the context of the dimeric species so that any substitution is present twice rather than four times. The *N*-terminal protomer carried multiple active-site substitutions and the *C*-terminal protomer was either native or carried an independent set of active-site mutations. Screening the library for the native catalytic activity allowed us to further evaluate the tolerance of this antibiotic-resistance enzyme to active-site variation, providing further insight into the potential for this antibiotic-resistance enzyme to adapt its function in response to selective pressure.

Results

Selection of TMP-resistant dimeric, doubly-mutated (DDM) R67 DHFR variants yields singly mutated dimers

To introduce asymmetric mutations in the R67 DHFR active site, a library of R67 DHFR protomers mutated at positions 66–69 was linked to a second, similarly mutated library of R67 DHFR protomers to form a library of covalent dimers, doubly mutated (DDM). The DDM dimer library encoded mutated protomer 1 with an *N*-terminal 6-histidine tag for protein purification and a *C*-terminal Glu-Leu linker

(encoded by an inserted, unique *SacI* restriction site) connecting to the *N*-terminus of mutated protomer 2. Residues 66–69 of both protomers were combinatorially mutated according to a previously described strategy for partial randomization.¹³ Briefly, most mutations were designed to participate in reactivity according to the following mechanisms: (1) The DHF substrate must be protonated at N5 to promote hydride transfer at C6. Contrary to chromosomally-encoded DHFRs, the active site channel of R67 DHFR does not contain an evident proton donor. We thus replaced the central Gln67 by Asp or Glu. (2) The dielectric environment in the *E. coli* chromosomal DHFR promotes hydride transfer by favoring substrate tautomerization.²³ We thus altered the dielectric environment in the active site of R67 DHFR by encoding Val66Asp, Gln67Lys, Ile68Lys, Ile68Arg and Tyr69Phe. (3) The active-site channel is relatively wide; we included bulkier residues in an attempt to increase the proximity between the substrate and cofactor. Because a single primer set was designed to capture all desired features, some additional mutations were also encoded, adding additional sequence diversity.

The library was restricted to eight amino acids encoded at position 66, four at position 67, twelve (and the stop codon) at position 68, and four at position 69 (Supporting Information Table SI). This encoded a maximum of 2.4×10^6 unique, combinatorially mutated dimer variants. The native sequence is not encoded, because no Gln is allowed at codon 67. A silent mutation at codon 74 [CTT (Leu) to TTG (Leu)] was also encoded to detect any potential contamination by native sequence. To confirm that the library construction was successful, 25 clones were picked for DNA sequencing from the obtained library of 10^6 unselected clones. Slight biases above the expected 1/8 ratio were observed for both protomers at position 66 for Asn and in position 67 a slight bias below the expected 1/4 ratio was observed for Glu. Similarly, position 68 was biased slightly above or below the expected 1/12 ratio for Ser, Asn and Trp; no Leu or Trp was observed among those 25 unselected clones and for protomer 2 no Met or stop codon was observed (Supporting Information Table SII). At position 69 a ratio slightly below the expected 1/4 was observed for Phe. No contamination with the native sequence was observed. In two instances, the *C*-terminal protomer included unforeseen residues (Supporting Information Table SII: I67 and S69 in DDM clone 14NS; K66 in DDM clone 19NS; compare to amino acids encoded, Supporting Information Table SI), which may result from primer impurity or errors introduced by the polymerase during PCR. Overall, library quality was deemed to be good.

The 10^6 DDM clones of R67 DHFR were selected for TMP resistance by plating equal aliquots on non-

selective and on selective minimal medium plates containing TMP. DNA sequencing of 20 TMP-resistant clones revealed a disparity of the DNA sequence compared to unselected constructs: 25% of the selected clones had only one mutated protomer (Supporting Information Table SIII). The Leu74 silent mutation was present in both protomers, demonstrating that this was not the result of contamination with native monomer or with DSM library members. Observation of a non-mutated protomer was unexpected, because the native Gln67 was not encoded in the library; its presence may result from contamination of the mutagenic primers with undesired nucleotides at a level that is too low to be observed in the unselected library, or may result from bacterial mutational activity as a response to strong selective pressure. We note that a single base change can suffice to alter the encoded RAK, to the native CAG.

To evaluate the genetic stability of the selected clones, we re-cloned the 20 individual dimeric inserts into fresh vector and transformed anew. Following two more selection rounds, the majority of the selected variants (14/20) held one protomer with native sequence at residues 66–69 and one mutated protomer; again, the Leu74 silent mutation was present in both protomers (Supporting Information Table SIII). Thus, clones with two mutated protomers after the first round of selection acquired a non-mutated protomer in subsequent selection rounds. We note that, in all cases but one, it is the *C*-terminal protomer that reverted to native sequence. Previous studies of a heavily mutated, covalent tetrameric species of R67 DHFR, incorporating up to 55 mutations, showed long-lasting genetic stability even after 7 days of subculturing under similar conditions.¹⁸ In that case, all mutations were positioned in surface or disordered regions of the construct rather than within the central active-site channel. Here, mutagenesis within the active site appears to have triggered molecular mechanisms to “correct” the sequence into a native form, preserving the bacteria from loss of activity of R67 DHFR under selective conditions.

Notwithstanding the mode of its appearance during selection, this indicates that the presence of one protomer with native sequence at residues 66–69 provided a significant survival advantage. This is in good agreement with our previous study of mutated monomers, where only three variants with native-like activity were identified among to 1,536 possible variants screened.¹³ The selection of the DDM library was repeated in different *E. coli* strains under similar conditions. Results consistently showed the tendency to enrich clones with one native protomer (data not shown).

Among the non-native protomer sequences that were observed following the rounds of selection, an

Table I. Deduced Amino Acid Sequence of the Active Site Residues 66–69 of the Mutated N-terminal Protomer Within the Dimeric DSM Library Following Selection for TMP Resistance

Selected clone	Residues 66–69	Selected clone	Residues 66–69
1S	NKSH	17S	TEFL
2S	TENH	18S	SDCF
3S	TENH	19S	GNNL
4S	ANTF	20S	DDNH
5S	TEFL	21S	GNNF
6S	NDTF	22S	SEYF
7S	TNTY	23S	IETY
8S	SKKY	24S	TEFL
9S	INSF	25S	AEFL
10S	DETL	26S	DDNY
11S	NNRH	27S	HEYF
12S	TEFY	28S	NKNY
13S	IKYH	29S	VDNH
14S	DESH	30S	NKNY
15S	NDTY	31S	DDNY
16S	SDWL	32S	INNf

important bias was observed at positions 66 and 67, which held the non-native Asn66/Lys67 pair in 9/20 variants, one occurrence of which was in a dimer where both protomers remained mutated. We note that the DDM sequences that held no native protomer following three rounds of selection had generally varied throughout the selection rounds.

Selection of TMP-resistant singly mutated dimers (DSM) R67 DHFR variants

On the basis of that preliminary observation, we sought to determine the breadth of sequence diversity that can be functionally accommodated in dimeric variants forming a tetramer with two native and two mutated protomers. To this effect, a library of mutated protomers was genetically linked to a native protomer to form a covalent dimer, singly mutated (DSM) library. As above, residues 66–69 of the N-terminal protomer were combinatorially mutated. To confirm that the library construction was successful, 32 clones were picked for DNA sequencing from the total library of 1.7×10^4 clones obtained. No deviations from the encoded codons were detected and no contamination with the native sequence was observed (Supporting Information Table SIV).

Some biases above the expected 1/8 ratio were observed at position 66 for Ile (9/32), Asn (7/32), and Asp (6/32), and no Ala was observed. Similarly, position 68 was slightly biased above the expected 1/12 ratio for Thr (5/32) and the stop codon (4/32); no Phe or Arg were observed among those 32 unselected clones, though they were identified following selection. Position 67 and 69 showed no strong bias towards any residue. Overall, library quality was deemed to be good.

The DSM dimer library was then selected for TMP resistance as described above. A 79% selection rate (number of selected clones/number of unselected) was obtained. DNA sequencing of 32 TMP-resistant clones confirmed that no native sequence had arisen in the mutated N-terminal protomer (Table I). Furthermore, the native C-terminal protomer showed no unexpected mutations. We note that the observed 79% selection rate of the DSM library does not necessarily indicate that nearly four out of five of the possible sequence variations in the N-terminal protomer give rise to an active enzyme species. Bacteria may be multiply transformed, thus harbour more than one variant at a time, which increases the apparent selection rate. Nonetheless, this selection rate is much $>0.2\%$ observed upon selection of the same mutated monomeric species.¹³ As all transformations and selections were undertaken similarly, this indicates that there is a greater fraction of functional variants when the mutated protomer is combined with a native one. This is further supported by the fact that only three variants among that mutated monomeric library had provided TMP resistance upon selection in *E. coli*,¹³ while a broad variety of variants were functionally selected here. Among the 32 clones sequenced, 27 unique sequences were identified in the N-terminal protomer (Table I). This represents roughly 2% of all possible combinations. The distribution of codons/amino acids observed for each mutated residue (66–69) was compared for unselected (Supporting Information Table SIV) and selected clones (Table I). For the selected clones, a slight (twofold) enrichment of Thr was observed at position 66 and a similar enrichment of Glu was observed at position 67. Only three occurrences of the non-native Asn66/Lys67 pair, which was observed in nearly 50% of the selected DDM variants, were recorded. Interestingly, a new pair emerged at the same position in this selection: the combination of Thr66/Glu67 was observed in nearly 20% of selected DSM variants (Table I). At position 68, a ninefold enrichment of Asn was observed as well as a sixfold enrichment of Phe. The distribution of codons for position 69 was similar for unselected and selected clones. The only unexpected mutation was His66 in mutant 27S, which was not designed into the mutagenic primers. We note that neither the exact sequences of the three functional monomeric variants (SKIH, INRY and GELH¹³), nor any closely related homolog, were observed among the 32 selected dimers. For those variants, all four protomers were identical such that the mutated sequences needed to be fourfold tolerated in the context of function. Here, the sequence at residues 66–69 for two of the protomers may be complementary, to the native residues on the two other protomers. Thus a greater sequence variation can be expected and, indeed, was observed.

Table II. Kinetic Parameters for the Non-Mutated DNM Dimer and Selected Mutated DSM Dimer Variants

Residues 66–69 in <i>N</i> -terminal protomer	k_{cat} (s^{-1})	$K_{\text{M}}^{\text{DHF}}$ (μM)	$K_{\text{M}}^{\text{NADPH}}$ (μM)	$k_{\text{cat}}/K_{\text{M}}^{\text{NADPH}}$ ($\text{s}^{-1} \mu\text{M}^{-1}$)	$k_{\text{cat}}/K_{\text{M}}^{\text{DHF}}$ ($\text{s}^{-1} \mu\text{M}^{-1}$)
DNM: VQIY (native)	0.28 ± 0.03	13 ± 4	13 ± 5	0.02	0.02
DSM: TENH (3S)	0.18 ± 0.01	28 ± 4	19 ± 4	0.01	0.007
ANTF (4S)	0.28 ± 0.02	51 ± 8	44 ± 6	0.006	0.005
TEFL (5S)	0.03 ± 0.01	35 ± 23	ND	—	0.001
INSF (9S)	0.06 ± 0.01	16 ± 4	>200	< 0.0003	0.004
TEFY (12S)	0.12 ± 0.01	18 ± 5	84 ± 20	0.001	0.007
IKYH (13S)	0.050 ± 0.004	15 ± 4	13 ± 2	0.004	0.003
AEFL (25S)	ND	ND	ND	—	—
HEYF (27S)	0.10 ± 0.01	8.5 ± 3.0	23 ± 7	0.005	0.012
NKNY (30S)	0.11 ± 0.01	13 ± 4	16 ± 5	0.007	0.009

The selected mutated dimeric DSM variants 4S, 9S, and 12S were expressed at levels similar to the non-mutated dimer (DNM), whereas variants 3S, 13S, and 30S were expressed about 50% less; variants 5S, 25S, and 27S expressed only in the range of 25% of the DNM level. The molecular weight of DNM and a subset of four selected DSM variants (3S, 5S, 12S, and 13S) were verified by mass spectrometry (Supporting Information Table SV), confirming that the dimeric species were expressed with no unexpected modifications.

Kinetic characterization of the selected DSM

The kinetic parameters of nine selected DSM variants were determined and compared to the native-like DNM (Table II). The k_{cat} obtained for DNM was similar to that previously reported for a similar dimer construct (0.28 s^{-1} vs. 1.2 s^{-1}).²¹ The K_{M} values obtained for DHF and NADPH (12.9 and 13.3 μM , respectively) were also similar to values reported for the similar dimer construct (6.3 and 2.7 μM , respectively).²¹ The differences may be attributed to experimental conditions (here, measured at ambient temperature in absence of β -mercaptoethanol rather than at 30°C with β -mercaptoethanol) and the different inter-dimer linker. For the selected DSM variants, the k_{cat} values were similar or 2- to 28-fold lower than for the DNM dimer, except for variant 25S where no activity was

detected; this variant was not further analyzed. In contrast, the K_{M} values for DHF and NADPH were generally similar to those of the DNM dimer, except for $K_{\text{M}}^{\text{NADPH}}$ of variants 5S and 9S, which were significantly increased.

Functional DSM variants assemble predominantly as “dimers of dimers”

For all subsequent experiments we focused on the selected DSM variants 3S, 9S, 12S, 13S, and 30S, based on their diversity of sequence and kinetic activity. Size exclusion chromatography was carried out to determine the oligomeric state of the native-like DNM dimer as well as the *N*-terminally mutated DSM variants. For DNM, a single species was observed at $\sim 36 \text{ kDa}$, confirming the formation of a dimer of the dimeric species, equivalent to the native tetrameric state of R67 DHFR (Table III). Similar results were observed for the variants 9S, 12S, and 13S, where the “dimer of dimers” species was predominant [Fig. 2(b)].

For variants 3S and 30S, the main peak on the chromatogram is consistent with the formation of a “dimer of dimers” yet a shoulder observed at $\sim 53 \text{ kDa}$ is consistent with the formation of a “trimer of dimers.” The formation of “trimers of dimers” opens new possibilities for assembly, which are illustrated in Figure 2. This model is imperfect because the “pending” monomers may not be adequately stabilized; we currently have no direct insight into the mode of assembly. We note that higher-order assembly of R67 DHFR multimers has previously been suggested in relation to the intermolecular association of a covalent tetrameric construct.¹⁸

The potential for the dimers to assemble in various configurations to form a functional active site may underlie the high selection rate observed. In a trimeric arrangement, the active site may be constituted of three non-mutated and one mutated protomer, reducing the deleterious effect of some mutations [Fig. 2(c)]. The presence of a non-polar residue at position 66 may favor the formation of a dimer–dimer assembly (9S and 13S). A polar residue at that position may increase the prevalence of the

Table III. Native Molecular Weight Determined by Size Exclusion Chromatography for the Non-Mutated DNM Dimer and Selected Mutated DSM Dimer Variants

Residues 66–69 in <i>N</i> -terminal protomer	Main peak (kDa)	Second peak (kDa)	Peak ratio (% main/second)
DNM: VQIY (native)	34.0	N/O ^a	100
DSM: TENH (3S)	33.4	52.9	61.5
INSF (9S)	33.3	45.5	92.6
TEFY (12S)	31.1	45.8	87.5
IKYH (13S)	33.3	N/O ^a	100
NKNY (30S)	35.3	52.9	74.1

^a N/O: none observed.

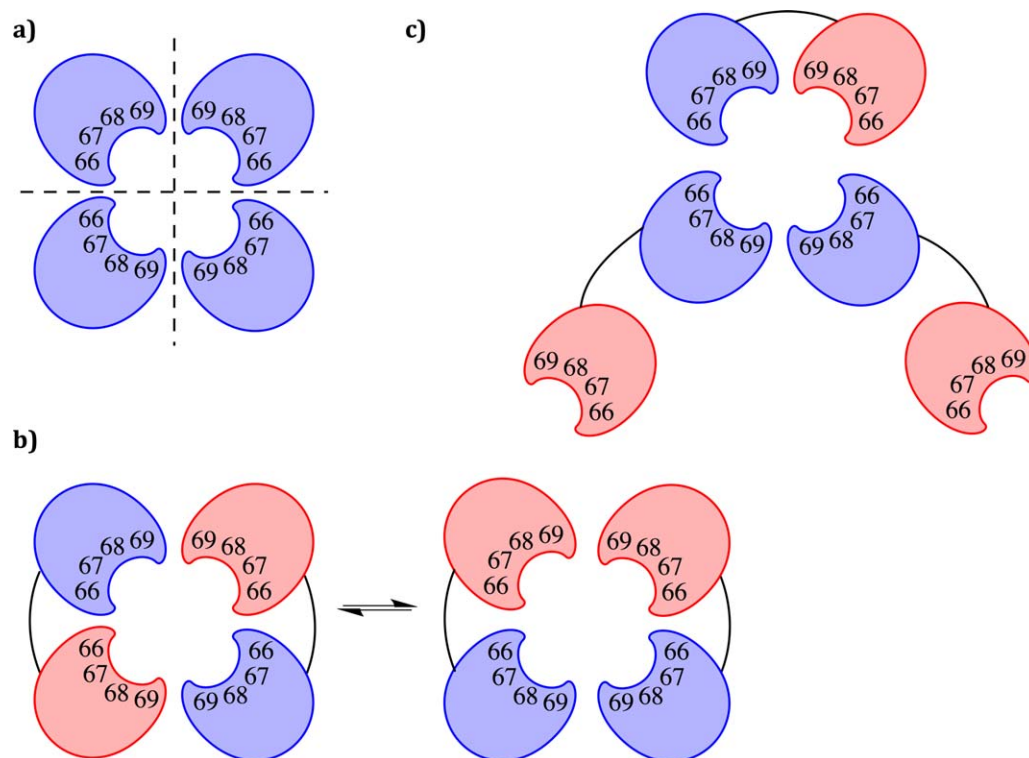


Figure 2. Schematic representation of protomer assembly for R67 DHFR variants. (a) Assembly of the native, monomeric R67 DHFR into a tetramer with a central active-site pore. (b) Assembly of a dimer of dimers, where the red protomers are mutated. (c) Putative assembly of a trimer of dimers to yield an active-site where only one protomer is mutated.

trimeric arrangement up to 37.5% in the most distant variant from the WT sequence, 3S.

Secondary structure is native-like

Circular dichroism (CD) allowed an assessment of the extent of secondary structure formation in functional dimers. The CD spectrum of the native DNM dimer is consistent with that reported for a previously constructed dimer, and is also very similar to that of the native homotetrameric R67 DHFR [see Zhuang *et al.*, Fig. 4(B)].²¹ The CD scans for DSM variants 3S, 9S, 12S and 30S all yield a CD spectrum that is similar to the non-mutated DNM, and are characteristic of high β -sheet content [Fig. 3(a)]. This is consistent with the native structure of R67 DHFR and indicates that the overall secondary structure was not significantly disrupted by the introduction of the mutations. Only slight differences in steepness were observed among variants.

The thermal stability of the dimeric variants was evaluated with CD between 25 and 95°C, to observe any perturbation of the secondary structure [Fig. 3(b)]. Surprisingly, monitoring the secondary structure showed no clear temperature denaturation for any R67 DHFR variant. Only variants 12S and 30S showed a small increase in ellipticity in the range of 90 to 95°C. This result was in apparent contradiction with the previously published T_m value for native R67 DHFR and a dimer construc-

tion,²¹ determined using differential scanning calorimetry (DSC). DSC reveals not only changes to secondary structure, but also changes to higher order structure. Therefore, we speculated that the previously reported T_m may have reported on changes to the tertiary and quaternary structure, while the secondary structure elements appear to be thermally stable up to at least 90°C.

Protomer assembly into the active structure is weaker in the selected DSM variants

To investigate this possibility we performed a thermal scanning fluorimetry shift assay. We used SYPRO orange as the hydrophobic fluorophore to bind hydrophobic patches of the protein as they become increasingly solvent-exposed upon thermal denaturation.²⁴ Similarly to DSC, this approach can reveal changes in secondary, tertiary, and quaternary structure, as observed by an increase in fluorescence. The resulting melting curve for the native DNM dimer yielded a T_m of 64.5°C, consistent with the value of 61.3°C reported by DSC for a similar native dimer construct [Fig. 3(c,d)].²¹ A difference of 1–3°C in T_m between the two methods is consistent with previously published data using other proteins.^{25,26} These results indicate that the secondary structure elements of the native DNM dimer—which is likely to be stabilized as a β -barrel spanning both protomers 6—remain essentially native-like up to

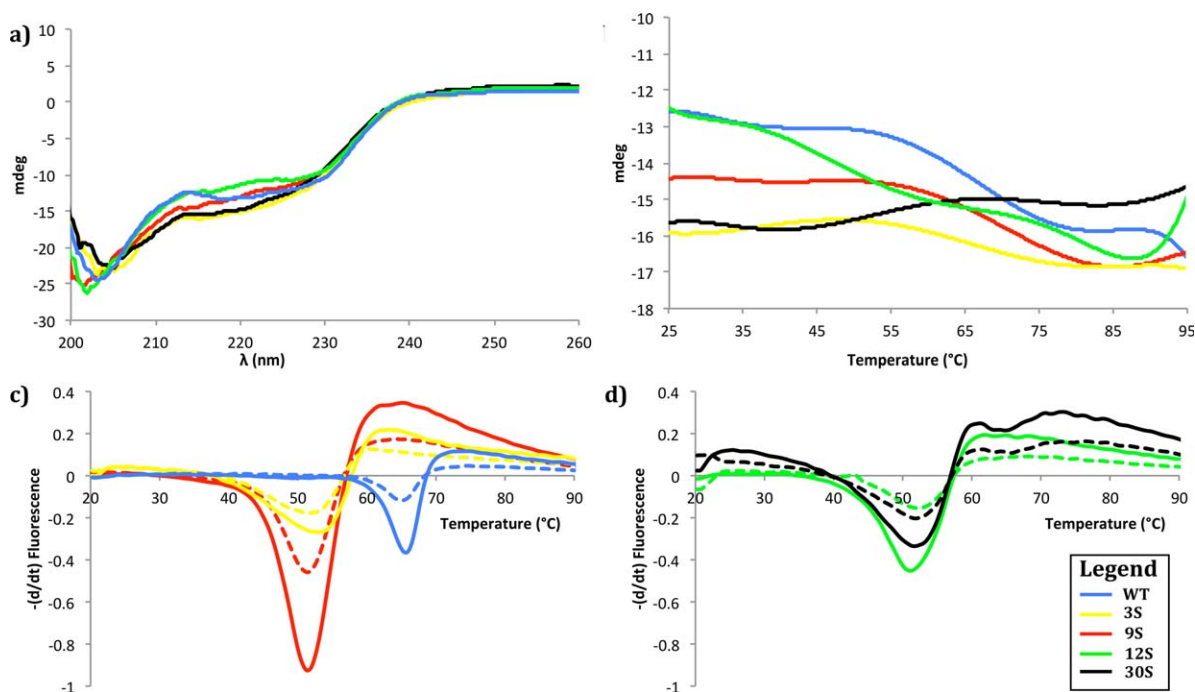


Figure 3. Thermal denaturation of R67 DHFR dimer mutants monitored by CD spectrometry and thermal scanning fluorimetry. (a) Far-UV CD spectra at 25°C of WT, 3S, 9S, 12S, and 30S. (b) Thermal denaturation measured at 215 nm at a rate of 25°C h⁻¹ for WT, 3S, 9S, 12S, and 30S. (c,d) First derivative analysis of representative thermal melting curves observed by fluorescence of SYPRO Orange. Melting curves are shown for a ratio of 3.33 x SYPRO to 8 μ M (full line) or 4 μ M (dashed line) protein, for (c) WT, 3S, 9S and (d) 12S and 30S.

$\geq 90^\circ\text{C}$, while the quaternary subunit assembly (association of two covalent dimers) breaks down above 60°C. It remains to be investigated whether the high thermostability at the level of secondary structure is an intrinsic property of the native R67 DHFR protomers, or whether it has been increased by linkage of the protomers into a covalent dimer.

The melting curves for the four DSM variants yielded T_m values $\sim 10^\circ\text{C}$ lower than for the DNM dimer, indicative of a perturbation to the quaternary subunit assembly (Table IV). Residues 66–69 are contained in the inner-most β -strand of the active site and do not participate directly in the interface of protein–protein interaction between the two dimers. Nonetheless, their modification, even in only two of the four protomers of the assembled R67 DHFR variants, appears to destabilize the tetrameric core assembly. Interestingly, the T_m value does not differ significantly among the DSM variants, whereas the introduced mutations differ greatly in physical and electrostatic properties.

To further investigate the high thermal stability of the secondary structure elements, we analyzed the stability of the protomers with respect to controlled proteolysis. Digestion with α -chymotrypsin over 24 h revealed a similar digestion pattern for the native DNM dimer and all DSM variants; variant 3S is shown as a representative example (Fig. 4). We also verified that no significant difference in digestion pattern was observed during the first 2 h

(data not shown). α -Chymotrypsin can theoretically recognize up to 22 cleavage sites in the R67 DHFR dimers studied. In all cases, the predominant proteolysis bands observed upon analysis by SDS PAGE following 24 h digestion were equivalent to the molecular weight of a single protomer (monomer) and a slightly shortened dimer, though additional bands indicate further preferred proteolysis sites. Cleavage is most likely to occur in the unstructured *N*- and *C*-terminal regions and in the inter-protomer linker as a result of accessibility. We note that the band of highest molecular weight (see “Dimer” arrow, Fig. 4) was digested significantly more rapidly, and to completion, in the DNM sample than for the variants examined, where a significant undigested band remained even after 24 h digestion.

For all variants, the core of the R67 DHFR protomer remained intact and inaccessible to α -chymotrypsin even following 24 h digestion. This indicates that the reduction in T_m by $\sim 10^\circ\text{C}$ for the selected variants did not destabilize packing to the extent where the core becomes accessible for proteolysis. To verify this, MS/MS analysis was performed on the main band of the DSM 3S dimer remaining following overnight proteolysis, corresponding to a single protomer. Both the non-mutated and the mutated protomers were observed by MS in that sample. Because this method is not quantitative, it was not possible to determine the ratio of non-mutated:mutated protomer resulting from digestion.

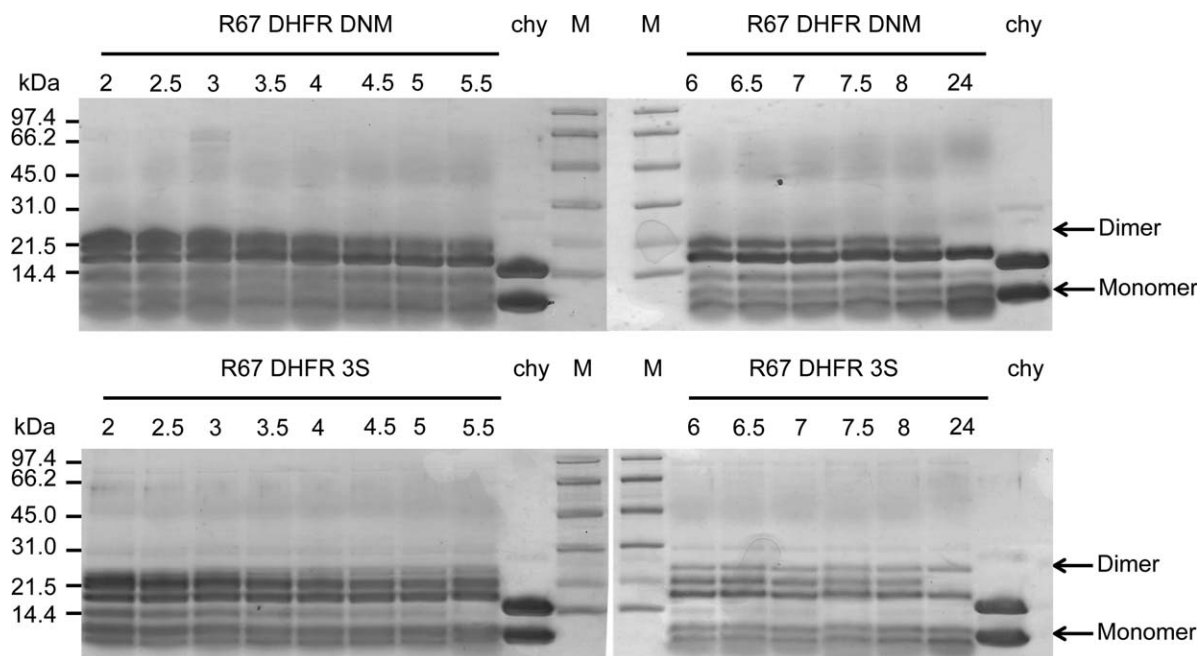


Figure 4. Limited chymotryptic digestion of R67 DHFR DNM (upper panel) and DSM variant 3S (lower panel), observed by tricine SDS PAGE. R67 DHFR DNM or DSM 3S were digested with chymotrypsin (1:100 chymotrypsin : R67 DHFR) for 2–24 h at 4°C, as indicated above each lane. Chy: chymotrypsin reference; M: molecular weight markers. Arrows on the right indicate the bands corresponding to the monomeric and dimeric species.

Nonetheless, this result indicates that the tight fold of each protomer is conserved upon introduction of mutations at the active site, consistent with the maintenance of secondary structure observed by CD upon thermal denaturation.

Recovery of activity following thermal denaturation is enhanced in the selected DSM variants

We further investigated the high thermal stability of the secondary structure elements and the multimeric assembly by verifying the activity of the variants following incubation at 95°C, under conditions mimicking the CD thermal denaturation experiment. After incubating the samples at 95°C for 90 minutes, slow cooling for 45 min allowed partial recovery of the activity (Table V, cf., conditions A and B). All DSM variants showed a high recovery ($\geq 45\%$) of the initial activity, where the variants 3S and 9S showed the highest recovery ($>70\%$). Interestingly, the recovery of DNM was significantly

lower ($<25\%$), consistent with a previous report of 15% activity recovered following heating to 75°C for a similar dimeric construct.²¹ Shorter incubation at 95°C (conditions C and D) was generally less disruptive to activity. Thus the selected DSM variants have acquired an increased resilience to heat denaturation, despite having reduced T_m values relative to the non-mutated DNM.

Discussion

Overall, we have found that the active, tetrameric form of R67 DHFR is very tolerant to active site variation, particularly in a context where two among the four protomers are native. In most multimeric enzymes, this would not be surprising, since a single protomer can often independently provide catalytic function. This is not the case in R67 DHFR, where the relatively small active site tunnel formed at the center of the tetramer can accommodate a single pair of ligands (DHF substrate + NADPH cofactor) at a time; all four protomers contribute simultaneously to ligand binding.

Through the process of selection, we have obtained DSM variants of R67 DHFR that exhibit a lower T_m . This illustrates that assembly of the dimers into an active structure (generally consisting of 4 protomers, though 6 protomers were also observed according to size-exclusion chromatography) was not as thermally stable in the selected DSM variants as in the non-mutated DNM dimer. Nonetheless, those selected DSM variants showed enhanced recovery of activity following thermal

Table IV. T_m Values Obtained by Thermal Scanning Fluorimetry for the Non-Mutated DNM Dimer and Selected Mutated DSM Dimer Variants

Residues 66–69 in <i>N</i> -terminal protomer	T_m (°C)
DNM: VQIY (native)	64.5
DSM: TENH (3S)	53.0
INSF (9S)	51.7
TEFY (12S)	51.8
NKNY (30S)	52.8

Table V. Recovery of Enzymatic Activity Following High-temperature Incubation

Condition	DNM (%)	3S (%)	9S (%)	12S (%)	30S (%)
A: 95°C, 90 min; RT, 1 min	11	18	18	26	17
B: 95°C, 90 min; RT, 45 min	23	72	72	46	47
C: 95°C, 4 min; RT, 1 min	78	95	90	50	11
D: 95°C, 4 min; RT, 45 min	71	>99	90	53	70

Samples were incubated at 95°C then at room temperature for the periods indicated, then activity was measured. Activity is expressed as percent of activity for that variant prior to incubation.

denaturation, suggesting an increased rate of dimer assembly into the active enzyme form. We note that the selection strategy (growth on agar plates at 37°C) was not designed to elicit this trait. We hypothesize that, as 37°C is largely inferior to the T_m of the selected variants (near 52°C), reduced thermal stability was not a selective disadvantage, while the improved dimer assembly into the active tetrameric form appears to have conferred a selective advantage.

Our observation of some selected DDM sequences that contain no native active-site sequence supports our earlier report of active-site modification in R67 DHFR, and confirms that no specific residue in the active-site tunnel formed by residues 66–69 is indispensable for catalytic activity.¹³ Importantly, we observed no partial enrichment toward native residues with retention of accompanying mutations: the residue 66–69 stretch either became entirely native, or else remained highly variable with only slight sequence biases. There may be a synergy between residues 66–69 that provides one robust solution for catalysis and structural integrity. We do not know if there is a functional or structural significance to the predominance of C-terminal reversion to the native sequence during the selection process; this may reflect different constraints on the two protomers as a result of their non-native fusion and/or the presence of the N-terminal 6-His tag.

The non-native Asn66/Lys67 residue pair was frequently observed in replacing the native Val66/Gln67 residue pair when the starting point was the doubly mutated dimer library DDM (9/20), but was less frequent when the starting point was the DSM singly mutated dimer library (3/32). In the DSM library the Thr66/Glu67 residue pair was predominant (6/32). This suggests that the Asn66/Lys67 or the Thr66/Glu67 substitution pair confers a selective advantage under the more stringent conditions of selecting for activity in a fully mutated active-site environment (DDM library) relative to the active sites that are part native, part mutated (DSM library). It is tempting to speculate on functional relevance of those pairwise substitutions. However, uncertainty with respect to the “dimer–dimer”

arrangement (do the likewise mutated monomers face each other in the tetramer or not?)¹⁹ and hence the resulting environment of substrate/co-factor binding opens a number of possible interaction profiles. Further investigation of this question would require the structural resolution of a ligand-bound dimer complex; earlier attempts to resolve the structure of a covalently linked R67 DHFR dimer in our laboratory have not been successful.⁸

The observation of high sequence diversity at the active site of one protomer in the presence of the second protomer with native sequence suggests that the enzymatic activity of R67 DHFR can be readily supported by an active site that is 50% native. This is supported by the observation of native-like enzyme kinetics for a number of DSM variants. Assembly of the four protomers to form a central active-site channel allows DHF and NADPH to slip in from opposite mouths of the tunnel¹⁷; their reactive atoms are juxtaposed near the center of symmetry of the tetramer. Catalysis is proposed to result from proper positioning of the reactive atoms to allow hydride transfer to occur on the pre-protonated DHF molecule. Again, no residue of the enzyme is indispensable for catalysis. Our results suggest that the native 66–69 residues of two protomers are sufficient to provide a native-like catalytic rate enhancement in R67 DHFR.

We also observed an additional structural dimension with the formation of “trimer–dimer.” This evidence for a functional assembly other than the “dimer–dimer” assembly increases the complexity of the system and is likely to reduce the impact of the introduced mutations. Along with the observed reversion of one protomer to the native sequence in the DDM dimers, assembly into trimers can provide an additional route for rapid recovery of trimethoprim resistance in the bacterial cell during selection.

In conclusion, we have found that R67 DHFR can explore a large active-site sequence space and various quaternary structures while maintaining native-like activity. It thus appears that this enzyme, postulated to have recently acquired DHFR activity as a response to selective pressure, is not in a highly specialized, evolutionary dead-end; instead, it appears to have a great potential for continued functional evolution. This finding is highly interesting because newly emerged and still evolving proteins might behave similarly to ancient proteins, and may thus serve as models with which to conduct studies on ancestral proteins.

Our results have direct implications on the threat that R67 DHFR holds with respect to antibiotic resistance. Already natively resistant to trimethoprim as a result of its fold being unrelated to that of DfrA DHFRs, R67 DHFR may continue to evolve even as we direct inhibitors to it,²⁷ and will need to be closely monitored.

Materials and Methods

Materials

All reagents were of the highest available purity. Restriction enzymes and DNA-modifying enzymes were from New England Biolabs and MBI Fermentas. Synthetic oligonucleotides were obtained from Integrated DNA Technologies (Coralville, USA). The infrared dye-labeled sequencing primers were from Li-Cor Biotechnology Division. The Sequenase 2.0 DNA sequencing kit was purchased from GE Healthcare. All aqueous solutions were prepared using water purified with a Millipore BioCell system. The 7,8-dihydrofolate (DHF) was prepared as described previously.²⁸

Bacterial strains and plasmids

E. coli SS320 was used to verify library quality by DNA sequencing. *E. coli* BL21 containing plasmid pRep4 (Qiagen, Mississauga) was used for expression. Chloramphenicol (Chl) was used at 10 $\mu\text{g mL}^{-1}$ and kanamycin (Kan) at 50 $\mu\text{g mL}^{-1}$ for maintenance of plasmids derived from pQE32Chl¹³ and for pRep4, respectively. Ampicillin (Amp) was used at 100 $\mu\text{g mL}^{-1}$ for maintenance of plasmids derived from pQE32 (Qiagen). Isopropyl β -D-1-thiogalactopyranoside (IPTG) was used at 1 mM for expression and TMP was used at 70 $\mu\text{g mL}^{-1}$ during selection.

Construction of the “dimer, doubly mutated” (DDM) and “dimer, singly mutated” (DSM) libraries

Starting from the construction described in Ref. 13, a covalent R67 DHFR dimer mutated at residues 66–69 was created by inserting a second mutated R67 gene between *SacI* and *HindIII* restriction sites on pQE32Chl (library DM, where DM stands for “dimer, mutated”). A covalent dimer containing two native (WT) R67 DHFR genes was also created by ligating the *BanII*-*XbaI* digested products of the native construct described in Ref. 13 and a construct containing the native R67 DHFR gene in pQE32 (Amp resistant) between *SacI* and *HindIII* sites, yielding construct DNM (where DNM stands for “dimer, non-mutated”). Library DM and construct DNM were digested with *SacI* and *XbaI*. This generated two fragments of 2500 and 1500 bp for each construct. The 2500 bp fragment of library DM (encoding the N-terminal mutated protomer) and the 1500 bp fragment from the DNM or the DM construct (encoding the C-terminal non-mutated or mutated protomer, respectively) were extracted using a QiaexII extraction kit (Qiagen). The fragments were ligated yielding the dimer libraries, where protomer 1 is mutated at residues 66–69 and protomer 2 is either non-mutated (“dimer, singly mutated,” DSM) or mutated (“dimer, doubly mutated,” DDM). Apart from the mutations, this dimer construct differs slightly from a previously reported

dimer, where protomer 1 was immediately connected to protomer 2²¹; the Glu-Leu linker was adopted here because it introduces a *SacI* restriction site for recombination of protomers into dimers.

The libraries were transformed by electroporation of 0.4–0.5 μg ligated material, into *E. coli* SS320. The bacteria were plated under non-selective conditions, on 1.2% agar Terrific Broth (TB) containing Chl. Over 2.4×10^6 colonies were obtained for the DDM library and 1.7×10^4 colonies for the DSM library. DNA sequencing of individual clones served to evaluate library quality prior to selection. Sequencing was performed using the Sequenase 2.0 DNA sequencing kit (GE Healthcare) with a Li-Cor IR2 automated system, using dye-labeled primers. Sequences were analyzed using the AlignIR V2.0 software (Li-Cor).

Library selection

The DDM and DSM dimer libraries in *E. coli* SS320 were harvested, propagated in Terrific Broth (TB), and the DNA retransformed into *E. coli* BL21/pRep4 for selection. Equal aliquots of transformation mixture were plated on TB plates (+ Chl and Kan) and on M9c minimal medium plates (+ Chl and Kan) containing IPTG and TMP. Colonies were grown at 37°C for 18 h. The plasmid inserts from selected clones were sequenced. Those clones were retransformed into *E. coli* BL21/pRep4 and subjected to two further rounds of selection and sequencing to confirm that activity was not due to cellular acquisition of TMP resistance or unforeseen changes in their DNA sequences.

Small-scale protein overexpression and purification

An overnight culture (0.2 mL) was used to inoculate 20 mL TB (+ Kan, and Chl or Amp for DNM). Cultures were propagated at 37°C with shaking until OD₆₀₀ reached 0.7. Protein expression was induced for 3 h following the addition of IPTG (1 mM final concentration). The cells were harvested by centrifugation (30 min, 3000g, 4°C). The cell pellet was resuspended in 4 mL of lysis buffer (100 mM potassium phosphate, pH adjusted to 8.0) and was disrupted by sonication on ice for 3 \times 30 s with 1 min intervals. Following centrifugation at 3000g for 30 min at 4°C, 1 mL of Ni-NTA agarose resin was added to the supernatant. The mixture was gently shaken for 30 min at 4°C and transferred into a gravitation column. The resin was washed twice with 5 mL lysis buffer, and twice with the same containing 10 and 20 mM imidazole, respectively. The enzyme was eluted with elution buffer 100 mM potassium phosphate buffer, pH 8 containing 250 mM imidazole.

Large-scale protein over-expression and purification

Induction of protein expression was undertaken as above, in 500 mL of TB. The cell pellet was

resuspended in 100 mM Tris-Cl buffer pH 8, and lysed by one passage through a cell disrupter (Constant Systems) adjusted to 27 kpsi. Following centrifugation (30 min, 3800g, 4°C), the supernatant was filtered over a 0.2- μ m filter. Purification was performed following a two-step purification protocol on an Äkta FPLC (GE Healthcare, Piscataway, NJ) at 4°C. The supernatant was applied to a 5 mL His-trap column pre-equilibrated with 100 mM Tris-Cl, pH 8. The protein was eluted with a gradient of imidazole (0–300 mM). Fractions were analyzed for DHFR activity¹³ and by tricine SDS-PAGE.²⁹ Those containing the most target protein were pooled and concentrated using an Ultra Amicon Ultracel 3.5K (Millipore). The concentrate was applied to a 1.5 \times 28 cm² Superose 12 column equilibrated in 100 mM TrisCl, pH 8 and eluted in the same buffer. The fractions were collected and analyzed for DHFR activity, by tricine SDS-PAGE and using the Bio-Rad protein assay.

Mass spectral characterization of R67 DHFR variants

Mass spectral analysis of R67 variants was achieved by HPLC-MS. Separations were performed on an 1100 LC system coupled to an ESI-TOF mass spectrometer (Agilent Technologies, Santa Clara, CA). The chromatographic column was a Poroshell 300SB-C8, 75 \times 2.1 mm², 5 μ particle size (Agilent Technologies) operated at 0.4 mL min⁻¹. The eluents used were 0.1% formic acid in water and 0.1% formic acid in acetonitrile. A linear acetonitrile gradient from 20 to 95% was used for elution. The mass spectrometer was operated in positive electrospray mode and mass spectra were acquired from m/z 110 to 3200. Deconvolution on the multiply charged species mass spectrum was performed using BioConfirm, an integrated program from MassHunter Qualitative Analysis.

Determination of multimerization state

Size exclusion chromatography was carried out at 4°C using a 1.5 \times 28 cm² Superose 12 column equilibrated in 100 mM Tris-Cl buffer pH 8. Ribonuclease A (13.7 kDa \pm 15%, 10 mg mL⁻¹), chymotrypsinogen A (25.0 kDa \pm 25%; 3 mg mL⁻¹), ovalbumin (43.0 kDa \pm 15%, 7 mg mL⁻¹) and albumin (67.0 kDa \pm 10%; 7 mg mL⁻¹) were used as standards to determine the oligomeric state of the R67 DHFR variants. The flow rate was 1.5 mg mL⁻¹. Standard curves were produced by plotting the logarithm of the molecular mass of standards versus K_{av} . $K_{av} = (V_E - V_v)/(V_B - V_v)$ where V_E is the elution volume, V_v is the void volume and V_B is the bed volume of the column matrix.

Kinetic characterization

Spectrophotometric enzyme activity assays were carried using a Varian Cary 100 Bio spectrophotometer.

The assay buffer was MTA (50 mM MES, 100 mM Tris-Cl, 50 mM acetic acid), pH 7. Assays were performed in the presence of 50 nM TMP to inhibit any residual *E. coli* chromosomal DHFR. The extinction coefficients used were 28,400 M⁻¹ cm⁻¹ at 282 nm for DHF, 6230 M⁻¹ cm⁻¹ at 340 nm for NADPH and 12,300 M⁻¹ cm⁻¹ at 340 nm to determine product formation.¹³ To determine K_M^{DHF} , DHF was held at 100 μ M while NADPH was varied from 0 to 100 μ M. Similarly, NADPH was fixed at 100 μ M to determine K_M^{NADPH} while DHF was varied from 0 to 100 μ M. The data were fitted to the Michaelis–Menten equation using non-linear regression analysis with the GraphPad software. Protein concentration was determined with the BioRad assay.

Circular dichroism

Circular dichroism (CD) spectra were recorded on a Jasco J-815 spectropolarimeter with a Peltier element for temperature control. Protein solutions were prepared at a final concentration of 0.25 μ g μ L⁻¹ in 50 mM potassium phosphate, pH 7. The path length of the quartz cell was 0.2 cm. Spectra were acquired at 20°C over the range 190–280 nm at a scan rate of 20 nm min⁻¹. Three scans were averaged. Buffer background recorded under the same conditions was subtracted. A single protein preparation was tested for all proteins except for DNM, the non-mutated R67 DHFR dimer (two preparations). Thermal denaturation experiments were performed by monitoring changes in ellipticity at 215 nm, at a scan rate of 25°C h⁻¹. The Savitzky–Golay method was used for curve smoothing.³⁰ The T_m values were determined using the derivatives function of the Jasco Spectra Manager software using the Savitzky–Golay algorithm for determination of first order derivative, where T_m is given as the maximum of this derivative.

Thermal scanning fluorimetry shift assays

T_m values were also determined by thermally induced incorporation of SYPRO Orange into the unfolding protein, with analysis using a Q-PCR thermal cycler as previously described.²⁴ Briefly, each combination of 10 \times , 5 \times , and 3.33 \times SYPRO Orange solution (Invitrogen) with 4 μ M or 8 μ M test protein was probed in a 96-well LightCycler plate (Sarstedt). SYPRO Orange and the protein were diluted with 50 mM potassium phosphate, pH 7, to a final volume of 20 μ L per well. Controls contained SYPRO Orange in buffer. The plates were sealed using Optically Clear Sealing Tape (Sarstedt) and heated from 20 to 95°C in a LightCycler 480 apparatus (Roche) with a ramp speed of 0.04°C s⁻¹ and 10 acquisitions/°C. Fluorescence was monitored with a CCD camera, using λ_{exc} = 483 nm and λ_{em} = 568 nm and a 1 s exposure time. Any curve showing a maximum

fluorescence plateau during denaturation was excluded from the T_m calculation.

For the T_m calculations, both temperature and fluorescence data were smoothed.³⁰ The first derivatives dFluo or dTemp were calculated using the cubic spline interpolation. The preliminary maximum was determined to obtain the half-values to the left and right of it. The linear fit for the curve outside the half-values was calculated, followed by the calculation of the average deviation from the fit. If the maximum was below the detection limit (fit value + 3 × deviation), the T_m determination was considered uncertain. The quadratic fit around the maximum was then calculated as follows to obtain T_m . The first derivative of the quadratic fit function (y-value) was set to 0 and the x-axis value (temperature) was resolved. Then, the average deviation of the curve points around the maximum from the quadratic fit was calculated. If the relative deviation was >5%, the T_m values were rejected if the corresponding maximum was below the detection limit. However, T_m values with a maximum above the detection but a relative deviation >5% were defined as uncertain.

Thermal denaturation and refolding assay

To simulate the thermal denaturation conditions of the CD experiment, the same temperature ramp was programmed into a BioRad PTC-200 PCR cycler. The total volume in each PCR tube was 70 μ L with a protein concentration at 1 μ g μ L⁻¹ in 50 mM potassium phosphate, pH 7. The activity of the R67 DHFR dimer variants was measured as described above for the different conditions. Condition A: Ramp to 95°C, hold for 90 min, keep at RT for 1 min and determine activity. B: Ramp to 95°C, hold for 90 min, keep at RT for 45 min and determine activity. C: Ramp to 95°C, hold for 4 min, keep at RT for 1 min and determine activity. D: Ramp to 95°C, hold for 4 min, keep at RT for 45 min and determine activity. Control: immediately determine activity. Specific activity was determined as described above.

Proteolysis by chymotrypsin

The packing of the mutated protomers was assessed by digesting with α -chymotrypsin (MP Biomedicals). Each R67 DHFR species was purified and quantitated as described above. The protein concentration was 3 μ g μ L⁻¹ and chymotrypsin was added in a mass ratio of 1%. Incubation was at 4°C and samples were taken at various time points up to 24 h. The progress of the proteolysis was analyzed by tricine SDS-PAGE.

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