

Non-homologous Recombination of Deoxyribonucleoside Kinases from Human and *Drosophila melanogaster* Yields Human-like Enzymes with Novel Activities

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In antiviral and cancer therapy, deoxyribonucleoside kinases (dNKs) are often the rate-limiting step in activating nucleoside analog (NA) prodrugs into their cytotoxic, phosphorylated forms. We have constructed libraries of hybrid enzymes by non-homologous recombination of the pyrimidine-specific human thymidine kinase 2 and the broad-specificity dNK from *Drosophila melanogaster*; their low sequence identity has precluded engineering by conventional, homology-dependent shuffling techniques. From these libraries, we identified chimeras that phosphorylate nucleoside analogs with higher activity than either parental enzyme, and that possess new activity towards the anti-HIV prodrug 2',3'-didehydro-3'-deoxythymidine (d4T). These results demonstrate the potential of non-homologous recombination within the dNK family for creating enzymes with new and improved activities towards nucleoside analogs. In addition, our results exposed a previously unknown role for the C-terminal regions of these dNKs in determining substrate selectivity.

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Introduction

Here we describe the use of directed evolution to reprogram the catalytic activity and selectivity of human thymidine kinase 2 (hTK2), through non-

homologous recombination with the deoxyribonucleoside kinase from *Drosophila melanogaster* (*DmdNK*). Deoxyribonucleoside kinases (dNKs) phosphorylate deoxyribonucleosides to their monophosphorylated forms, and are often the rate-limiting enzymes in the activation of anticancer and antiviral nucleoside analog (NA) prodrugs. In tumor cell lines, the administration of exogenous dNKs *via* gene therapy has been shown to potentiate the biological activity of several NA prodrugs.^{1–4} However, the kinetic properties of dNKs constitute a limiting factor in the efficacy of these therapies, and there is a need to identify enzymes with higher efficiency and improved specificity towards NAs for further development of this strategy.

Based on sequence similarity, hTK2 and *DmdNK* belong to a large subfamily of dNKs (type-1) that includes other human enzymes (deoxycytidine kinase, hdCK; and deoxyguanosine kinase, hdGK) and dNKs from insects.⁵ Type-1 dNKs share the same fold, but vary widely in their substrate specificities and overall catalytic efficiencies. *DmdNK* possesses the highest catalytic rates of any type-1 dNK, and is a multisubstrate enzyme that

Abbreviations used: dNK, deoxyribonucleoside kinase; NA, nucleoside analog; d4T, 2',3'-didehydro-3'-deoxythymidine; hTK2, human thymidine kinase 2; *DmdNK*, *Drosophila melanogaster* deoxyribonucleoside kinase; hdCK, human deoxycytidine kinase; hdGK, human deoxyguanosine kinase; TK, thymidine kinase; HSV TK, herpesvirus thymidine kinase; dT, thymidine; dC, deoxycytidine; dA, deoxyadenosine; dG, deoxyguanosine; FARA, 2-fluoro-9 β -D-arabino-furanosyladenine; dFdC, 2',2'-difluorodeoxycytidine; 3TC, 2'-deoxy-3'-thiacytidine; ddT, 2',3'-dideoxythymidine; ddC, 2',3'-dideoxycytidine; GCV, ganciclovir; ACV, acyclovir; ddI, 2',3'-dideoxyinosine; aa, amino acid residue.

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phosphorylates pyrimidine and purine deoxyribonucleosides, as well as several NAs. *DmdNK* has been the subject of both rational and random mutagenesis approaches to probe its substrate specificity.^{6,7} In contrast with *DmdNK*, the less studied hTK2 is ~30 to 80-fold less efficient at phosphorylating pyrimidine nucleosides, and has no activity towards purines. Despite their functional differences, the active site residues of these enzymes are highly conserved. However, our previous research has shown that the narrow specificity of hTK2 is not a result of the primary shell differences with *DmdNK*.⁸ Furthermore, other studies suggest that non-active site features such as flexible loops⁹ and C-terminal extensions¹⁰ have subtle effects on the specificity of dNKs.

Given the conservation of active site residues, the lack of structural information for hTK2, and the limitations of rational design in investigating non-local interactions, we have utilized a directed evolution approach in this study. Recombination can be an effective tool for sampling sequence space and generating enzymes with new catalytic activities, and recent studies have also emphasized the utility of recombination techniques in identifying regions of functional importance in protein frameworks.¹¹ One of the most successful examples of engineering dNKs utilized DNA family shuffling to recombine the closely related herpes virus thymidine kinases (HSV TK-1 and HSV TK-2, 78% DNA identity).¹² In theory this approach is a good strategy for other suicide gene/prodrug combinations; however, the sequence similarity between type-1 dNKs is below the threshold at which DNA shuffling techniques are effective. Homologous recombination-based methods can only shuffle sequences of >70% identity,¹³ and preferentially generate crossovers in high-homology regions. Because *DmdNK* and hTK2 are only 48% identical at the DNA level, we have employed the non-homologous recombination techniques of ITCHY¹⁴⁻¹⁶ and enhanced crossover SCRATCHY^{17,18} as a route to their combinatorial diversification. From single and triple-crossover libraries, we have isolated hybrids with novel activities, including the introduction of activity towards the anti-HIV prodrug d4T (2',3'-didehydro-3'-deoxythymidine), and we have imparted purine activity on a predominantly hTK2 scaffold. These results provide insight into the structure/function relationships of hTK2 and *DmdNK*, and demonstrate the utility of non-homologous recombination within the dNK family for creating enzymes with new and improved activities towards nucleoside analogs.

Results

Library construction and analysis

Two single-crossover ITCHY libraries of hTK2 and *DmdNK* were generated to cover both possible orientations, consisting of 4×10^6 transformants (*DmdNK*-hTK2; DH) and 5×10^6 transformants

(hTK2-*DmdNK*; HD). The theoretical diversity is 4.9×10^5 (648 bp \times 750 bp); therefore, these libraries provide essentially full coverage of available sequence space. The distribution of crossover locations was checked by sequencing randomly selected clones from each library; a broad distribution of crossover locations was observed (Supplementary Data Figure S1).

Because crossovers can occur at any base, two-thirds of all ITCHY library members will contain frameshifts, which result in premature termination or non-functional, frame-shifted progeny. Before recombining the ITCHY libraries to generate multiple-crossover chimeras, the libraries were subcloned into pInSAlect¹⁹ and plated on carbenicillin for selection of clones in the correct reading frame. As expected, approximately one-third of each library (DH, 33%; HD, 29%) survived this selection. Post-selection sequence analysis of 24 clones confirmed that all hybrid genes possessed in-frame crossovers.

For the creation of multiple-crossover chimeras, we divided the in-frame ITCHY libraries into three approximately equal sections (~250 bp) with short overlaps between each section, then amplified the sections by PCR using skewed primers. The resulting fragments, each containing a single crossover, were assembled by overlap extension to generate two complementary triple-crossover libraries, each with $>10^6$ transformants: hTK2-*DmdNK*-hTK2-*DmdNK* (HDHD) and *DmdNK*-hTK2-*DmdNK*-hTK2 (DHDH). This strategy is a variation on the enhanced SCRATCHY protocol¹⁸; however, we assembled the multiple-crossover chimeras by overlap extension PCR, instead of by DNA shuffling, in order to control the number of crossovers, and to minimize the frequency of single-crossover variants and/or revertant wild-type sequences. With the previous protocol, 58% of clones had ≤ 1 crossover.¹⁸ Similar levels of background would have overwhelmed the genetic selection used herein, making it difficult to isolate functional, multi-crossover enzymes. We chose to design the libraries as triple-crossovers to increase the sequence diversity while limiting the structural disruption that >3 crossovers may introduce. Sequencing of 30 clones (14 from DHDH; 16 from HDHD) showed that each contained three crossovers, implying that any wild-type or single-crossover background was present at <3%. Each analyzed clone represented a unique sequence, and a broad diversity of crossover locations was observed.

Selection for thymidine kinase activity

Chimeras that retained thymidine kinase activity were selected from the HD, DH, HDHD and DHDH libraries by functional complementation of the *tdk*⁻ *Escherichia coli* strain KY895.²⁰ Less than 0.1% of each ITCHY library survived this functional selection. DNA sequencing of these chimeras yielded a majority of clones that were *DmdNK*-like, though a small number of clones were found that more closely resembled hTK2. All functionally selected single-

crossover clones examined by DNA sequencing showed a similar pattern, with the recombination site near the N or C terminus.

The two triple-crossover libraries were also subjected to functional selection in *E. coli* KY895. Eighteen HDHD clones survived the selection (<0.01%). DNA sequencing revealed that six of these clones contained three crossovers; however, the other 12 possessed only a single crossover. In the complementary DHDH library, only one of 24 chimeras was identified with three crossovers; the remaining 23 were single-crossover hybrids. These single-crossover clones were similar to the functionally selected ITCHY clones, with all crossovers occurring near the termini. No wild-type parental sequences were observed. From these results, it appears that despite the library design, single-crossover products were present at very low levels in each library. Furthermore, these single-crossover clones were more likely to retain function and survive the functional selection than the multiple crossover clones.

Modeling of selected chimeras

Homology models of selected chimeras were constructed, based on the crystal structures of *DmdNK*^{21,22} (Figure 1), to elucidate the effects of crossovers on key structural features. Full sequence details for each clone are available in Supplementary Data 1. Four representative clones from the single-crossover libraries were chosen for further study (Figure 2). DH-03 and HD-15 represent the predominantly *DmdNK* chimeras, with crossovers occurring near the C terminus and N terminus, respectively. DH-03 contains a crossover in a non-homologous region of $\alpha 8$, and based on alignment of the parental sequences, possesses a net insertion of 10 aa. HD-15 has only the first 11 aa from hTK2, followed by a

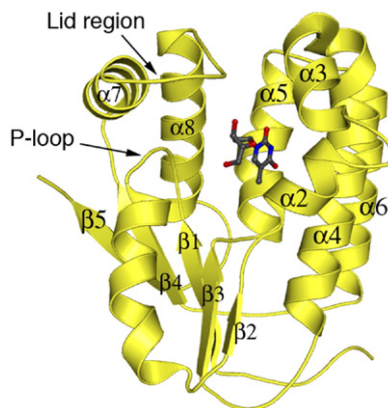


Figure 1. Conserved features of type-1 dNKs. The monomeric subunit of *DmdNK*²² is used as an archetypal example. The overall structure is a three-layer $\alpha\beta\alpha$ Rossmann fold. Two conserved features are present in all type 1-dNKs: the phosphate binding P-loop (GX₄GKS/T) located after strand $\beta 1$; and the Lid region, a connecting loop between $\alpha 7$ and $\alpha 8$, which contains conserved arginine residues at the phosphate donor site.

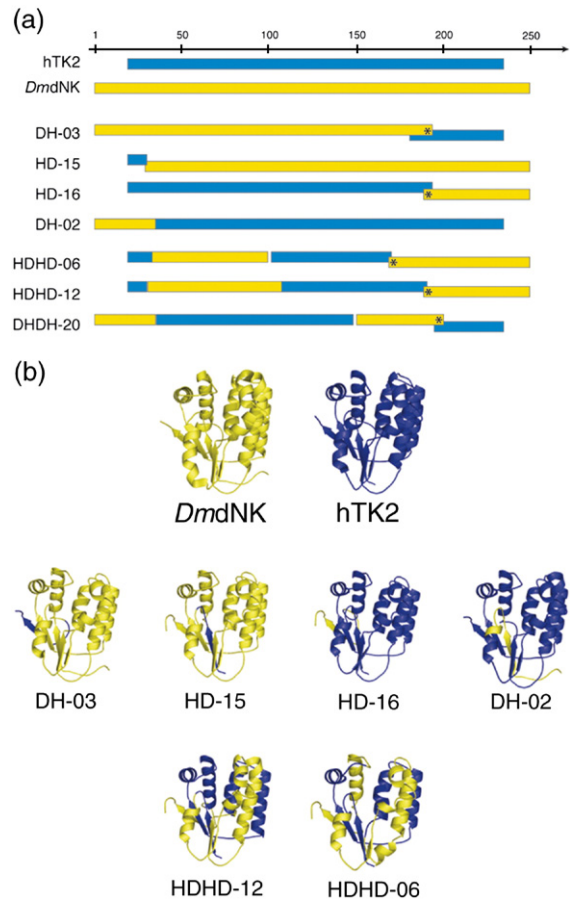


Figure 2. Comparison of parental enzymes and selected chimeras. (a) Schematic representation of selected chimeric protein sequences based on a CLUSTALW sequence alignment of the parental enzymes. Segments from hTK2 and *DmdNK* are represented as blue and yellow bars, respectively. Amino acid substitutions (point mutations) are marked by asterisks. (b) Structural models of the parental enzymes and selected chimeras. The structure of *DmdNK* (PDB entry 1OT3²²) is shown in yellow. All homology models were generated using SWISS-MODEL.³⁴ The homology model of hTK2 is colored blue. The homology models of the single and multiple-crossover chimeras (DH-02, DH-03, HD-15, HD-16, HDHD-06 and HDHD-12) are colored as the parental enzymes, with segments from hTK2 and *DmdNK* shown in blue and yellow, respectively.

crossover leading to a 1 aa insertion in the section of the P-loop (SGKTT) that is identical in the two parents. Clones HD-16 and DH-02 represent the chimeras with predominantly hTK2 sequence. HD-16 consists of the first 193 residues from hTK2, and residues 190–250 from *DmdNK*. A mid-codon crossover introduces a mutation (W190R, *DmdNK* numbering) at the beginning of the *DmdNK* segment. This crossover also yields a 6 aa insertion. Like HD-15, DH-02 has a crossover located in the P-loop.

Two of the six functionally selected HDHD triple-crossover chimeras were analyzed further: HDHD-06 and HDHD-12 (Figure 2). In each case, the first crossover is located in the P-loop. In HDHD-06, the second crossover creates a 2 aa deletion, which

in the homology model appears to disrupt $\alpha 4$ and $\beta 3$. The third crossover introduces a point mutation (S170A, *DmdNK* numbering) and extends the Lid by insertion of 2 aa between $\alpha 7$ and $\alpha 8$. In HDHD-12 there is no insertion or deletion at the second crossover. The third crossover occurs after the Lid region, at the end of $\alpha 8$, and results in a 2 aa insertion as well as the same point mutation (W190R) observed in HD-16.

Only one triple-crossover chimera from the DHDH library, DHDH-20, survived functional selection (Figure 2(a)). As observed in the other libraries, the first crossover was in the P-loop. The second crossover occurs after $\alpha 6$ and results in a single-residue deletion, while the third occurs at the end of $\alpha 8$ and introduces both a Phe to Ile mutation, and also a 5 aa insertion.

Enzyme purification and activity with natural deoxyribonucleosides

The seven selected chimeras were subcloned into pET14b to incorporate an N-terminal (His)₆-tag for purification *via* metal affinity chromatography. Yields varied from 0.1–4 mg protein per liter of culture, with DH clones having lower yields than HD clones. DHDH-20 proved to be insoluble, and therefore could not be characterized further.

The catalytic activities of the selected hybrids and parental enzymes towards the four natural deoxyribonucleosides were measured spectrophotometrically (Table 1). Of these substrates, the pyrimidines are substrates for both parental enzymes, while the purines are substrates for *DmdNK* only. These differences allow the chimeras to be classified as *DmdNK*-like or hTK2-like, based on their substrate specificities.

Three of the four single-crossover chimeras have substrate-activity profiles that mirror the predominant parental enzyme (Table 1). The two chimeras with the majority of their sequence derived from *DmdNK* (DH-03 and HD-15) exhibited *DmdNK*-like substrate profiles, demonstrating activity towards pyrimidines and purines. For HD-15, the introduction of 11 hTK2 amino acid residues at the N terminus had little effect on purine activity, but reduced the k_{cat}/K_M for dT and dC ~16-fold. The other chimera, DH-02, had 77% of its amino acid residues derived from hTK2 and exhibited hTK2-like, pyrimidine-specific activity. All three chimeras had reduced catalytic activity towards pyrimidine substrates, compared with *DmdNK*. DH-02 and DH-03 had k_{cat}/K_M values at least two orders of magnitude worse than *DmdNK* for every substrate.

In contrast, HD-16 shares 78% sequence identity with hTK2 and yet has a *DmdNK*-like substrate profile (Table 1). Surprisingly, this chimera gained the ability to phosphorylate the purines dA and dG, an activity not detectable in hTK2. Furthermore, HD-16 has two to fivefold higher catalytic efficiency than hTK2 for pyrimidines.

Two triple-crossover hybrids, HDHD-06 and HDHD-12, were also profiled. Both have approximately equal contributions from each parent (HDHD-06, 44% hTK2; HDHD-12, 40% hTK2), but differ in their substrate specificities (Table 1). HDHD-06 is a pyrimidine-specific, hTK2-like enzyme. It possesses catalytic efficiencies for dT and dC that are threefold lower than hTK2, and >100-fold lower than *DmdNK*, due to decreases in k_{cat} for both substrates. The *DmdNK*-like hybrid, HDHD-12, has improved k_{cat}/K_M values relative to hTK2 for all four substrates. The Michaelis constants with pyrimidine substrates are comparable to *DmdNK*, though the

Table 1. Kinetic parameters of wild-type and chimeric enzymes

	dT			dC		
	k_{cat} (s ⁻¹)	K_M (μM)	k_{cat}/K_M (M ⁻¹ s ⁻¹)	k_{cat} (s ⁻¹)	K_M (μM)	k_{cat}/K_M (M ⁻¹ s ⁻¹)
A. Kinetic parameters of wild-type and chimeric enzymes with pyrimidine nucleosides						
hTK2 ^a	1.8±0.2	6.7±1.1	2.7×10 ⁵	2.0±0.2	16.2±1.1	1.2×10 ⁵
<i>DmdNK</i> ^a	17.5±0.6	1.9±0.3	9.1×10 ⁶	19.3±0.5	2.2±0.3	9.0×10 ⁶
DH-03	0.54±0.01	10.4±1.3	5.2×10 ⁴	0.48±0.01	23.5±1.7	2.0×10 ⁴
HD-15	3.6±0.1	6.2±0.9	5.9×10 ⁵	4.3±0.1	8.2±0.6	5.2×10 ⁵
HD-16	2.5±0.1	1.9±0.3	1.3×10 ⁶	3.2±0.1	14.1±2.2	2.3×10 ⁵
DH-02	0.69±0.02	9.5±1.6	7.3×10 ⁴	0.98±0.01	74.0±3.1	1.3×10 ⁴
HDHD-06	0.43±0.01	5.4±0.4	8.0×10 ⁴	0.72±0.02	19.5±1.8	3.7×10 ⁴
HDHD-12	2.6±0.1	1.8±0.3	1.4×10 ⁶	3.5±0.1	3.8±0.3	9.1×10 ⁵
	dA			dG		
	k_{cat} (s ⁻¹)	K_M (μM)	k_{cat}/K_M (M ⁻¹ s ⁻¹)	k_{cat} (s ⁻¹)	K_M (μM)	k_{cat}/K_M (M ⁻¹ s ⁻¹)
B. Kinetic parameters of wild-type and chimeric enzymes with purine nucleosides						
hTK2 ^a	nd	nd	nd	nd	nd	nd
<i>DmdNK</i> ^a	26.2±0.5	315±23	8.3×10 ⁴	12.5±0.4	710±70	1.8×10 ⁴
DH-03	0.45±0.01	1300±190	1.7×10 ²	0.23±0.01	610±70	7.4×10 ²
HD-15	4.6±0.1	200±14	2.2×10 ⁴	5.6±0.1	310±20	1.8×10 ⁴
HD-16	1.2±0.1	1240±170	9.3×10 ²	0.23±0.1	1530±110	1.5×10 ²
DH-02	nd	nd	nd	nd	nd	nd
HDHD-06	nd	nd	nd	nd	nd	nd
HDHD-12	2.5±0.1	1100±60	2.2×10 ³	1.1±0.1	2340±200	4.5×10 ²

^a Data from Gerth *et al.*⁸

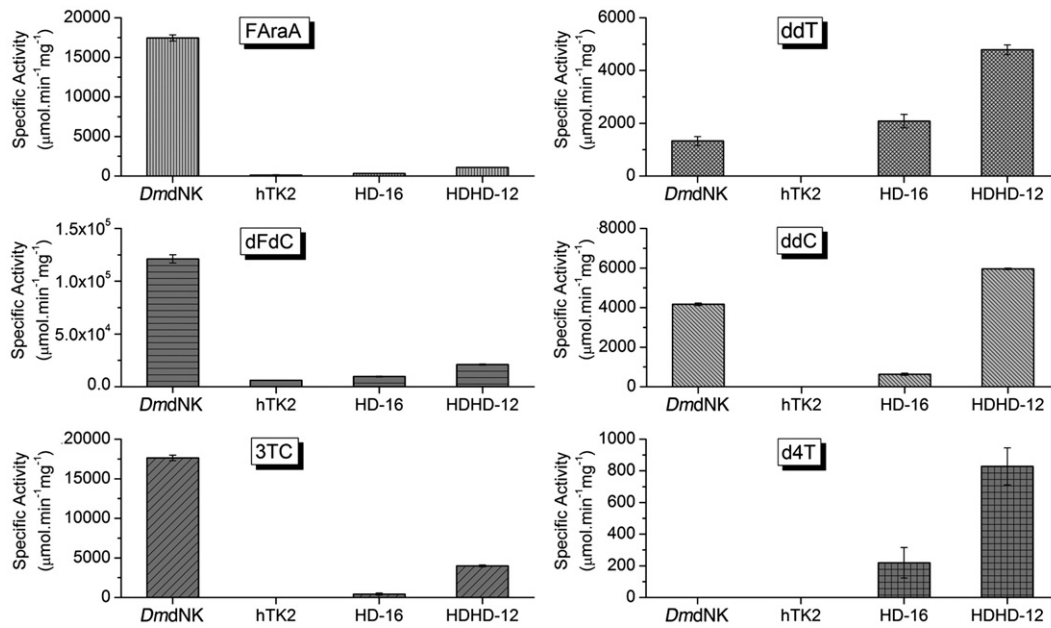


Figure 3. Specific activities of *DmdNK*, hTK2, HD-16 and HDHD-12 for phosphorylation of the following nucleoside analogs: FArA (2-fluoro-9 β -D-arabinofuranosyladenine), dFdC (2',2'-difluorodeoxycytidine), 3TC (2'-deoxy-3'-thiacytidine), ddT (2',3'-dideoxythymidine), ddC (2',3'-dideoxycytidine) and d4T (2',3'-didehydro-3'-deoxythymidine). No catalytic activity towards the NAs GCV (ganciclovir), ACV (acyclovir) and ddI (2',3'-dideoxyinosine) was detected.

k_{cat} values mirror the lower values of hTK2. Relative to *DmdNK*, the catalytic efficiency of HDHD-12 is reduced by 38-fold and 40-fold towards dA and dG, respectively.

Chimeras HD-16 and HDHD-12 had *DmdNK*-like substrate profiles, despite having large segments from hTK2 (74% and 40% of total sequence, respectively). Interestingly, both enzymes possessed the W190R mutation (*DmdNK* numbering), introduced at a crossover. To control for the effect of this mutation on substrate specificity, we constructed the revertant tryptophan variants of each clone. These mutants were purified and characterized with all four natural deoxyribonucleosides. The catalytic activities of the revertant enzymes were the same as the W190R-containing chimeras (data not shown).

Specific activities with nucleoside analogs

Based on their broadened specificities for the natural nucleosides, the specific activities of HD-16 and HDHD-12 were determined with purine and pyrimidine NAs (Figure 3). As with the natural nucleosides,

the parental enzymes have distinct NA substrate profiles. Of the nine analogs tested, *DmdNK* showed activity towards FArA, dFdC, 3TC, ddC and ddT. hTK2 was able to phosphorylate dFdC and FArA, albeit poorly. Neither *DmdNK* nor hTK2 showed activity against d4T, and no phosphorylation of GCV, ACV or ddI was detected for any parental or chimeric enzyme.

Both chimeras could phosphorylate FArA, dFdC and 3TC, though in each case the specific activity was reduced compared to *DmdNK* (Figure 3). Such reductions in activity, relative to the most active parent, appear common for enzymes derived from homology-independent recombination.²³ HD-16 retained 2–8% of *DmdNK*'s activity for these three substrates; for HDHD-12, the residual activities were 6–23%. The activity of each chimera was substantially higher than hTK2 (Figure 3).

The chimeras displayed their highest specific activities with the dideoxynucleosides ddT and ddC (Figure 3). Both HDHD-12 and HD-16 showed greater activity towards ddT (by 3.6-fold and 1.6-fold, respectively) than *DmdNK*. HDHD-12 also

Table 2. Kinetic parameters of wild-type enzymes and selected chimeras with nucleoside analogs

	ddC			d4T		
	k_{cat} (s ⁻¹)	K_{M} (μM)	$k_{\text{cat}}/K_{\text{M}}$ (M ⁻¹ s ⁻¹)	k_{cat} (s ⁻¹)	K_{M} (μM)	$k_{\text{cat}}/K_{\text{M}}$ (M ⁻¹ s ⁻¹)
hTK2	nd	nd	nd	nd	nd	nd
<i>DmdNK</i>	1.5±0.1	380±20	3.9×10 ³	nd	nd	nd
HD-16	0.42±0.01	1240±110	3.4×10 ²	0.07±0.01	66.3±10.3	1.1×10 ³
HDHD-12	1.5±0.1	135±11	1.1×10 ⁴	0.29±0.01	148±17	1.9×10 ³

nd: not detectable.

possessed 1.5-fold greater activity towards ddC than *DmdNK*, although HD-16 only retained 15% specific activity for this substrate. Most strikingly, both hybrid enzymes were able to phosphorylate d4T, an activity that neither *DmdNK* nor hTK2 possesses.²⁴

ddC and d4T are both important components of the therapy regimens used for the treatment of AIDS. Therefore, the kinetic parameters of HD-16 and HDHD-12 with these NAs were analyzed in detail (Table 2). The catalytic efficiency of HD-16 for ddC phosphorylation was 11-fold lower than *DmdNK*, due to k_{cat} and K_M both worsening. For HDHD-12, a reduction in K_M led to a 2.8-fold improvement in k_{cat}/K_M for ddC. Furthermore, the catalytic efficiency of HDHD-12 for ddC phosphorylation is 28-fold greater than that of hdCK,²⁵ the enzyme responsible for phosphorylating ddC *in vivo*.²⁶ The new d4T activity is interesting because d4T is a poor substrate for dNKs; the catalytic efficiencies of our evolved enzymes are two orders of magnitude higher than those previously reported for natural or engineered dNKs.^{27,28} The activities of HD-16 and HDHD-12 with d4T were similar ($k_{\text{cat}}/K_M \approx 10^3 \text{ M}^{-1}\text{s}^{-1}$). Substrate turnover was slow ($k_{\text{cat}} < 1 \text{ s}^{-1}$), but the hybrids had surprisingly low K_M values of 66.3 and 148 μM , respectively. Remarkably, these values indicate that the active sites of the chimeras are more optimized for d4T binding than *DmdNK* is for purines ($K_M(\text{dA}) = 315 \mu\text{M}$; $K_M(\text{dG}) = 710 \mu\text{M}$; Table 1B).

Discussion

Human dNKs are important targets for engineering, due to their role in activating NA prodrugs and also their potential utility for suicide gene therapy. We have used non-homologous recombination to identify regions accounting for functional differences between hTK2 and *DmdNK*, and to improve the activity of hTK2 towards clinically relevant NAs.

Selection for thymidine kinase activity from the DH and HD libraries yielded a majority of clones that were predominantly comprised of *DmdNK* sequence. This appears to reflect the impaired ability of wild-type hTK2 to fold in *E. coli*.¹⁹ Every functional single-crossover clone had its crossover situated near the N or C terminus. Single crossovers in the interior of these dNKs are clearly too disruptive for maintaining function, a bias previously observed for other enzymes.¹³ Crossovers near the C terminus occurred in or beyond $\alpha 8$, and typically fell into non-homologous regions. In contrast, N-terminal crossovers always occurred in the conserved P-loop. This loop is located between $\beta 1$ and $\alpha 1$ (Figure 1) and is critical for binding and positioning the phosphoryl donor, therefore the introduction of crossovers into this key structural and catalytic feature was unexpected. The crossover in the P-loop may be more tolerated than in the neighboring α helix or β sheet, due to the greater flexibility of loop secondary structure elements. Furthermore, crossovers in the P-loop region may also be favorable, as many of the residues in the adjacent $\beta 1$ and $\alpha 1$ are conserved between

the parental enzymes, thus minimizing functional disruption.

Most of the single-crossover hybrids characterized kinetically had specificity profiles that mirrored the predominant parent; the exception was HD-16, which has an hTK2-like sequence but displayed a *DmdNK*-like substrate/activity profile. The crossover in HD-16 is located in the loop region connecting $\alpha 8$ and $\beta 5$ near the C terminus of the enzyme. Beyond $\beta 5$, this portion of the protein sequence shows low overall sequence homology among type-1 dNKs and none of the critical active site residues are located in this region. Further complicating the evaluation of its functional relevance, no structures of *DmdNK* are available that include this region; the structures have been determined using a recombinant, C-terminal truncation mutant lacking the last 20 residues and in these structures, the last 14–16 residues have no ordered electron density.^{21,22} Crystal structures of full-length sequences from other type-1 dNKs,^{21,25} on the other hand, show that this portion of the protein folds into a helical segment ($\alpha 9$) but otherwise has no apparent critical contacts with the rest of the protein. In a separate study, it has been shown that C-terminal truncations of *DmdNK* affect its substrate specificity; the C-terminal 20 amino acid residues were not required for phosphorylation of deoxyribonucleosides but were necessary for full activity with dideoxy- and purine-ribonucleosides.¹⁰ The deletion of ten additional residues reduced the activity to 1% that of wild-type. A similar C-terminal truncation of hdGK also resulted in a virtually inactive enzyme.²⁹ Enzyme inactivation upon deletion of the C terminus could be the result of misfolding or a lack of necessary conformational constraints. Based on our substitution experiments, we suggest that subtle structural changes in protein topology might be responsible for the observed changes in catalytic activity and broadened substrate specificity. Moreover, comparison of DH-03 and HD-16 (Figure 3, Table 1) suggests that the C-terminal subdomain of *DmdNK* activates hTK2, but not that the C-terminal region of hTK2 is incompatible with purine activity, *a priori*. Further biochemical and biophysical studies will be required to clarify the role of this subdomain.

HDHD-12 was the most active chimera isolated from the triple-crossover libraries. In addition to the first crossover in the N-terminal P-loop region, HDHD-12, like HD-16, possesses the 52 aa C-terminal subdomain derived from *DmdNK* and a *DmdNK*-like substrate profile. The second crossover occurs at the beginning of $\alpha 5$ that forms part of the deoxyribonucleoside-binding pocket. As the region shows high amino acid sequence conservation between the two parental enzymes, we predict only small local-structure perturbations resulting from the crossover. On the other hand, the substitution with the 78 amino acid residue segment of *DmdNK* (Ser31–I107) divides the kinase scaffold, including the active site binding pocket, into two equal segments. Given the extensive interface between these segments and the low overall sequence

homology between *DmdNK* and hTK2, protein crystallography might be one way for obtaining more detailed structural information and for identifying potential conformational changes that translate into altered enzyme function.

Upon testing for NA activity, HDHD-12 and HD-16 both displayed improved activity (relative to hTK2) for six of the nine compounds screened. Quite unexpectedly, the specific activities of HDHD-12 towards ddC and ddT, as well as of HD-16 towards ddC, were actually greater than either parental enzyme. Even more remarkably, the two chimeras are able to phosphorylate d4T, an activity that both parents lack. Moreover, this activity is greater than that exhibited by any known natural or engineered dNK. While it is not surprising that the chimeras have activity towards ddC and ddT, given the known role of the *DmdNK* C terminus in dideoxyribonucleoside phosphorylation, it is interesting that the catalytic efficiency of hybrid enzymes surpasses that of *DmdNK*. The reason for the improved activity for dideoxyribonucleosides, as well as the new activity towards d4T, is not clear from available structural information. In the case of ddC, the increase in the k_{cat}/K_M for HDHD-12 results from improvement in the apparent binding constant, possibly caused by small conformational changes as a result of chimeragenesis. A similar analysis for d4T is not possible as the parental enzymes have no detectable activity for the NA.

Though it is an oft-repeated rule in directed evolution that "you get what you select for", our results suggest that even when a direct selection for novel or improved activity is not available, enzymes with these properties can be found with minimal screening. Furthermore, the chimeras obtained in this study contain successful combinations that would have been impossible to predict from the available structural and phylogenetic information. Future developments, such as the implementation of direct screens or selections for NA phosphorylation and recombination with other type-1 dNKs, should enable further improvement of the chimeras described. Moreover, as the percentage of active chimeras isolated was very low, creation of alternative crossover libraries is also a potential subject of future work. In a general sense, alternate crossovers could sample a greater portion of the sequence-function diversity. More specifically, it would be interesting to explore crossover combinations that result in N and C termini from the same parent. It has been observed in recombination experiments of other enzyme families, that functional chimeras tend to have the N and C termini from the same parent.³⁰ The ultimate goal of our experiments is to identify humanized enzymes with higher activities towards NAs than towards natural nucleosides, making them even more promising candidates for suicide gene therapy. The results described here constitute a major step towards this goal.

Materials and Methods

Materials

Pfu Turbo DNA polymerase from Stratagene (La Jolla, CA), was used for all cloning, unless otherwise indicated. The α -phosphorothioate dNTP mix and Exonuclease III were purchased from Promega (Madison, WI). Pyruvate kinase and lactate dehydrogenase were from Roche Biochemicals (Indianapolis, IN). Plasmid DNA was isolated using the Qiaprep Spin Miniprep kit and PCR products were purified with the Qiaquick PCR Purification kit (Qiagen, Valencia, CA). Enzymes were purchased from New England Biolabs (Beverly, MA), and all other reagents were from Sigma-Aldrich (St. Louis, MO), unless noted. Constructs were confirmed by DNA sequencing.

ITCHY library construction

The single-crossover DH and HD libraries were generated using full-length *DmdNK* (DNA encoding amino acid residues 1–250) and a truncated variant of hTK2 (DNA encoding amino acid residues 20–234).⁸ The libraries were prepared using the thio-ITCHY protocol described,¹⁴ though instead of using *DmdNK* and hTK2 cloned in tandem, each gene resided on separate pDIM plasmids, which were linearized, mixed and assembled *via* PCR as described below. *Taq* DNA polymerase (Promega) was used for the PCR amplification. The template and primer combinations used in constructing the DH ITCHY library were: pDIM-hTK2 linearized with *Nde*I; pDIM-*DmdNK* linearized with *Spe*I; forward primer 5'-CGC CAT ATG AGT GGG AAG ACG ACA TGC-3'; and reverse primer 5'-GCG ACT AGT TCA CTT GCA CGA CTG CGG TCG TCT CTG-3'. For the HD library the templates and primers used were: pDIM-*DmdNK* linearized with *Nde*I; pDIM-hTK2 linearized with *Spe*I; forward primer 5'-CGC CAT ATG AGC GGG AAG ACC ACG TAT TTG AAC-3'; and reverse primer 5'-CGC ACT AGT TCA GGC TGC CAT GGG GAA AAG G-3'. Thermocycling conditions were: 95 °C for 3 min; two cycles of 94 °C for 10 s and 68 °C for 60 s; add primers and α -dNTPs; 30 cycles of 94 °C for 10 s, 55 °C (DH) or 58 °C (HD) for 60 s, 68 °C for 210 s; one cycle 68 °C for 4.5 min. The PCR products were used to generate the DH and HD ITCHY libraries as described.¹⁶ The resulting ligation mixture was used to transform *E. coli* KY895. Libraries were plated on LB-agar containing carbenicillin (Carb; 100 μ g/ml). After overnight growth, the colonies were recovered by scraping the plates with 2YT medium, supplemented with glucose (2% w/v) and glycerol (15% v/v). The recovered libraries were divided into aliquots; one portion was saved at -80 °C as a frozen stock, and another portion was isolated in plasmid form.

Reading frame selection

The single-crossover libraries were amplified by PCR to replace the stop codon from pDIM-DH and pDIM-HD with a codon for glycine (shown underlined), using the forward primer 5'-ATT AAC CCT CAC TAA AGG GA-3' (T3), and the reverse primers 5'-AGC ACT AGT TCC TGG GCA ATG-3' (DH library) and 5'-GGC ACT AGT TCC TCT GCG ACC CTC TG-3' (HD library). The PCR products were purified, digested with *Nde*I and *Spe*I, and ligated into pInSAlect¹⁹ that had been digested with the same enzymes. The ligation products were used to

transform *E. coli* DH5 α -E by electroporation. Cells were plated on LB-agar containing chloramphenicol (Cm; 50 μ g/ml) and incubated at 37 °C. Colonies were recovered in 2YT medium supplemented with glucose (2%) and glycerol (15%). Reading frame selection was carried out by replating the libraries on LB-agar containing carbenicillin (100 μ g/ml) and incubating at 21 °C. Libraries were recovered as described above, and the plasmid pools (pInSALect-DH and pInSALect-HD) were isolated.

SCRATCHY library construction

Each ITCHY library was divided into three approximately equal and overlapping sections. The sections were amplified by PCR using skewed primers (i.e. a *DmdNK*-specific forward primer and an hTK2-specific reverse primer, or *vice versa*), to generate products that each contained one crossover. The pInSALect-DH library plasmids were used to generate the three DH segments, with the following primers: *DH-frag1*, DF1 (5'-AGT TGT CAG AGG TCA TAT GGC GGA GGC AGC-3') and HR1 (5'-GCG TAA GAC CCC AGC GAG AGG C-3'); *DH-frag2*, DF2 (5'-CAC ACC GCC CCA ACC AAC AAG AAG-3') and HF2 (5'-CTG GTA ACA AGT CTC AGG ATT G-3'); *DH-frag3*, DF3 (5'-CCG GCA GCG GGC TCG TTC TG-3') and HR3 (5'-TGC GAA TAC CAC TAG TTC CTG GGC AAT G-3'). Using pInSALect-HD as the template, the following HD fragments were generated: *HD-frag1*, HF1 (5'-AGT TGT CAG AGG TCA TAT GTC AGT G-3') and DR1 (5'-CTT CTT GTT GGT TGG GGC GGT GTG-3'); *HD-frag2*, HF2 (5'-GCC TCT CGC TGG GGT CTT ACG C-3') and DR2 (5'-CAG AA CGA GCC CGC TGC CGG-3'); *HD-frag3*, HF3 (5'-CAA TCC TGA GAC TTG TTA CCA G-3') and DR3 (5'-GAATAC CAC TAG TTC CTC TGG CGA C-3'). The amplifications were carried out using the following cycling conditions: 95 °C for 2 min; 30 cycles of 95 °C for 30 s, 57 °C for 30 s, 72 °C for 45 s; one cycle of 72 °C for 10 min.

Overlap assembly was used to generate the full-length, triple-crossover products. For each library, three gel-purified DNA segments from the previous step were mixed in equal amounts to a final concentration of 15 ng/ μ l. Fragments *DH-frag1*, *HD-frag2* and *DH-frag3* were mixed for assembly of the DHDH library. The HDHD library was assembled using fragments *HD-frag1*, *DH-frag2* and *HD-frag3*. The primerless assembly reaction used the following conditions: 95 °C for 2 min; 35 cycles of 95 °C for 30 s, 52 °C for 30 s, 72 °C for 1 min; one cycle of 72 °C for 10 min.

Next, the reassembled full-length templates were purified from the PCR mix and used as templates for PCR amplification using gene-specific, flanking primers, which incorporate NdeI and SpeI restriction sites to facilitate cloning. For amplification of the DHDH library the following primers were used: a *DmdNK*-specific forward primer, 5'-CTA TCA TAT GGC GGA GGC AGC ATC CTG TGC-3'; and a hTK2-specific reverse primer, 5'-GCG ACT AGT TCA TGG GCA ATG CTT-3'. Similarly, amplification of the HDHD library was performed with an hTK2-specific forward primer, 5'-CAG CCA TAT GTC AGT GAT CTG TGT CG-3', and a *DmdNK*-specific reverse primer, 5'-CTC ACT AGT TCA TCT GGC GAC CCT CTG-3'. The cycling conditions were: 95 °C for 2 min; 15 cycles of 95 °C for 30 s, 54 °C for 30 s, 72 °C for 1 min; one cycle of 72 °C for 10 min. The resulting products were purified, digested with NdeI and SpeI, and ligated with the pDIM vector. The resulting plasmids were used to transform *E. coli* KY895.

Selection for thymidine kinase activity

E. coli strain KY895²⁰ was transformed by plasmids and plated on LB-agar containing Carb (100 μ g/ml). For library selections, the cells were recovered and replated on thymidine kinase selection plates³¹ containing 2% Casamino acids, 1.5% (w/v) noble agar (both Difco, BD Biosciences, Sparks MD), 0.5% (w/v) NaCl, 0.2% glucose, 50 μ g/ml carbenicillin, 10 μ g/ml 5'-fluorodeoxyuridine (Acros Organics, Morris Plains, NJ), 12.5 μ g/ml uridine and 2 μ g/ml thymidine.

Site-directed mutagenesis

The R190W mutations were introduced by overlap extension PCR. For mutagenesis of HD-16, two overlapping PCR products were formed with the primer pairs (mutation sites are underlined): forward primer 5'-TAA TAC GAC TCA CTA TAG GG-3' (T7F) and reverse primer 5'-GTG TAT CAG CCA GCT GCC TTT GAT GAG C-3'; forward primer 5'-CAT CAA AGG CAG CTG GCT GAT ACA CC-3' and reverse primer 5'-TAT GCT AGT TAT TGC TCA G-3' (T7R). The purified products were assembled in a PCR reaction with the primers T7F and T7R. For mutagenesis of HDHD-12, the primer pairs were: T7F with reverse primer 5'-GGT GTA TCA GCC AGA GCC ACT CCT CAT GG-3'; and forward primer 5'-CAT GAG GAG TGG CTC TGG CTG ATA CAC C-3' with T7R. The products were also assembled using T7F and T7R.

Enzyme purification

Genes of interest were subcloned into the NdeI and SpeI sites of pET14b (vector modified to include SpeI site). The resulting plasmids were used to transform *E. coli* BL21 (DE3) pLysS (Novagen). LB cultures (2 l) were used for IPTG-induced protein expression, as described.⁸ Enzymes were purified by batch/gravity flow column purification using Talon resin (Clontech, Mountain View, CA), also as described previously.⁸ The pH of each lysis and storage buffer was adjusted such that it was at least one unit away from the pI (estimated using the ExPASy ProtParam tool³²) of the (His)₆-tagged protein being purified; this was done to maximize solubility. The buffers used were: pH 7.0 for HD-15, HD-16 and HDHD-06; pH 7.5 for *DmdNK* and HDHD-12; and pH 8.0 for hTK2, DH-02 and DH-03. Amicon Ultra centrifugal filter units (Amicon Bioseparations, Billerica, MA) were used to exchange the purified proteins into storage buffer (50 mM Tris-HCl, 150 mM NaCl, 1 mM β -mercaptoethanol, 10% glycerol). Yields were 0.1–4 mg per liter culture, depending on the enzyme, and preparations were typically >90% pure as analyzed by SDS-PAGE. Protein concentrations were quantified by measuring $A_{280\text{nm}}$ using molar extinction coefficients calculated as described by Pace *et al.*³³ Aliquots of purified proteins were frozen in liquid nitrogen and stored at -80 °C until use.

Steady-state kinetic assays

Spectrophotometric assays to measure nucleoside and NA phosphorylation were performed as described.⁸ All experiments were performed in triplicate. Data were fit to the Michaelis-Menton equation using Origin (OriginLab, Northampton, MA). Specific activities towards nucleoside analogs were measured with 1 mM each substrate, and 0.4 μ g of each enzyme.

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.jmb.2007.05.021](https://doi.org/10.1016/j.jmb.2007.05.021)

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