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Comparative Study of Purine and Pyrimidine Nucleoside Analogues Acting on the Thymidylate Kinases of Mycobacterium tuberculosis and of Humans


**Thymidine monophosphate kinase (TMPK) from Mycobacterium tuberculosis (TMPKmt) is an attractive target for the design of specific inhibitors. This fact is the result of its key role in the thymidine pathway and of unique structural features in the active site observed by X-ray crystallography, especially in comparison to its human counterpart (TMPKh). Different 5-modified thymidine derivatives, as well as purine and pyrimidine analogues or C-nucleosides were tested on TMPKmt and TMPKh, and the results were rationalized by docking studies. 5-Halogenated 2'-deoxyuridines are the best inhibitors of TMPKmt found and present the highest selectivity indexes in favor of TMPKmt.**

**KEYWORDS:** inhibitors • kinases • purine • pyrimidine • nucleosides

**Introduction**

Nucleoside analogues such as zidovudine and stavudine, or acyclovir and ganciclovir, have demonstrated strong antiviral properties and are successfully used in the treatment of HIV or herpes virus infections.[1] Administered as prodrugs, these compounds are phosphorylated by cellular or viral kinases, and the resulting nucleoside triphosphates have been shown to inhibit viral DNA synthesis by different mechanisms.[2] The enzymes involved in the first two steps of activation, thymidine kinase (TK) and thymidine monophosphate kinase (TMPK), are essential members of the thymine metabolic pathway. Similarly, deoxycytidine analogues,[3] such as zalcitabine and gemcitabine, are phosphorylated by deoxycytidine kinase and cytidine monophosphate kinase. Characterization of the enzymes involved in these phosphorylation steps is a prerequisite for the design of new specific prodrugs or inhibitors.

In the case of anti-herpes-virus agents, the most frequently used compounds are derivatives of guanine (acyclovir, ganciclovir) or 5-substituted pyrimidines (brivudin, idoxuridine, trifluridine).[1] To be active on the viral DNA polymerases, these compounds need to be phosphorylated to their triphosphate forms. The first phosphorylation step is achieved by a virus-encoded protein (HSV or VZV TK or CMV protein kinase) and is therefore restricted to infected cells. After conversion of these analogues into the corresponding monophosphates, they are further phosphorylated to nucleoside diphosphates, by TK in the case of the thymine derivatives or by cellular kinases in the other cases. The crystal structure of HSV-1 TK[4] reveals the presence of a fold common with the nucleoside monophosphate kinase (NMPK) family, made up of a five-stranded parallel β sheet and other additional structural elements. This fold, part of the protein core, contains the active site. Moreover, an insertion loop of HSV-1 TK is analogous to the LID domain of NMPKs, which covers the active site upon binding of adenosine triphosphate (ATP). HSV-1 TK and TMPKs exhibit an overall sequence identity below 16%. However, crystal structures of binary complexes of TK[5, 6] or TMPKs[7–10] revealed a similar orientation of the enzyme-bound substrate relative to the catalytic centre. The position of the substrate appeared shifted by about 2 Å after structure superimposition based on the strictly conserved residues.

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The recent characterization of TMPK from *Mycobacterium tuberculosis* (TMPKmt) in our laboratory revealed new structural and catalytic features, which make the enzyme a favored target for antituberculosis drugs.[11] Some thymidine-5′-O-monophosphate analogues were shown to be inhibitors of TMPKmt.[11–13] For therapeutic application, these analogues should be administered as nucleosides, which should be phosphorylated to their nucleotide monophosphate counterparts by an *M. tuberculosis* protein. However, there is no TK activity in this bacterium,[14, 15] which seemingly renders the use of TMPKmt as a target invalid. Our findings that 5-bromo-2′-deoxyuridine (5BrdU) and 3′-azido-3′-deoxythymidine (AZT) are inhibitors of TMPKmt as potent as their nonphosphorylated 5′-modified derivatives remove this barrier.[16] No metabolic processing would be necessary for these compounds to become pharmacologically active, which opens new avenues in the search for specific inhibitors of TMPKmt as antituberculosis drugs. Unlike in the treatment of anti-herpesvirus infections, these compounds would directly act on their target, TMPKmt.

In the study reported herein, the thymine moiety of thymidine (dT) was explored in particular. Different purine and pyrimidine nucleoside analogues, as well as C-nucleosides known to be more stable in vivo and nonheterocyclic nucleosides, were tested as inhibitors of purified TMPKmt. Some antiherpetic inhibitors were selected, as their anabolism and catabolism have been extensively studied. We also checked for specificity by carrying out parallel tests on human TMPK (TMPKh) and the experimental data were rationalized by docking studies.

**Results and Discussion**

The inhibitory potency of base-modified dT analogues was tested on recombinant TMPKh, purified as described in the Experimental Section, and on TMPKmt as described previously.[11] From the known crystal structure of TMPKmt complexed with dTMP (Protein Databank (pdb) accession number: 1G3U),[10] the potential effect of chemical substitutions was analyzed. A comparative analysis of the crystal structures of TMPKmt and TMPKh (pdb accession number: PDB1E2Q),[17] was performed, to assess the selectivity of the explored compounds on the two TMPKs.

The 5′-position of the pyrimidine ring

Substitution of the 5-methyl group with a halogen atom or an alkyl chain was explored first (Table 1). The 5-halogenated deoxyuridines exhibited affinities for TMPKmt similar to those of their corresponding nucleoside monophosphates, which are substrates of this enzyme.[11] However, these compounds are much less potent towards TMPKh in comparison to their nucleotide counterparts: the Michaelis constant $K_m$ for dTMP (5 µM) is 40 times lower than the inhibition constant $K_i$ for dT (180 µM). Despite significant structural similarities, TMPKmt and TMPKh show only 23% sequence identity. Analysis of the active site entrance in the vicinity of the 5′-position of the ribose moiety highlighted significant amino acid differences (Figure 1). We predicted that D163 in TMPKmt should be

### Table 1. Evaluation of the inhibitory potencies of 5-modified dU and AZdU analogues.[a]

<table>
<thead>
<tr>
<th>R₂ OH</th>
<th>Acronym</th>
<th>TMPKmt</th>
<th>Acronym</th>
<th>TMPKh</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₃</td>
<td>dT</td>
<td>27</td>
<td>AZT</td>
<td>28</td>
</tr>
<tr>
<td>H</td>
<td>dU</td>
<td>1020</td>
<td>AZdU</td>
<td>810</td>
</tr>
<tr>
<td>F</td>
<td>SFdU</td>
<td>212</td>
<td>N.I.[b]</td>
<td></td>
</tr>
<tr>
<td>Cl</td>
<td>SCH₂U</td>
<td>10</td>
<td>375</td>
<td>AZCldU</td>
</tr>
<tr>
<td>Br</td>
<td>SBrdU</td>
<td>5</td>
<td>214</td>
<td>AZBrdU</td>
</tr>
<tr>
<td>I</td>
<td>SIdU</td>
<td>33</td>
<td>350</td>
<td>N.I.[b]</td>
</tr>
<tr>
<td>CF₃</td>
<td>SCF₃dU</td>
<td>97</td>
<td>1020</td>
<td></td>
</tr>
<tr>
<td>OH</td>
<td>SHOdU</td>
<td>270</td>
<td>N.I.[b]</td>
<td></td>
</tr>
<tr>
<td>CH₂OH</td>
<td>SHOMedU</td>
<td>820</td>
<td>2700</td>
<td></td>
</tr>
<tr>
<td>CH₂=CH₂</td>
<td>BVdU</td>
<td>625</td>
<td>1100</td>
<td></td>
</tr>
<tr>
<td>CH₂CH₂</td>
<td>SEtU</td>
<td>1140</td>
<td>N.I.[b]</td>
<td></td>
</tr>
</tbody>
</table>

[a] Enzymatic assay conditions were as described in the Experimental Section and the figures given correspond to the $K_i$ values in µM. N.I., no inhibition detected at a final concentration of [b] 1 mM, [c] 3 mM, and [d] 4 mM.
involved in hydrogen bonding to the 5'-hydroxy group of nucleosides.\[16\] In TMPKh, this position is occupied by E149, which forms a salt bridge with R45. This latter residue is equivalent to Y39 in TMPKmt, which is connected to D163 through a bridging water molecule (W43). We propose that the 5'-hydroxy group moves to a position mimicking that of the bridging water molecule in the TMPKmt/dTMP crystal. In TMPKh, such a rearrangement would not be favorable because of the salt bridge (R45–E149), which leads to an affinity of a nucleoside for TMPKh lower than that of the corresponding nucleotide.

Consequently, the selectivity of nucleosides is in favor of TMPKmt, with selectivity indexes (SI = \(K_i \text{ TMPKmt}/K_i \text{ TMPKh}\)) between 2 (for 5-bromovinyl-2'-deoxyuridine (BVDU)) and 40 (for 5-chloro-2'-deoxyuridine (SCldU) and SBrdU).

The presence of a chlorine or a bromine atom at the 5-position of dU molecule increases its affinity, while that of an iodine atom (bigger than a methyl group) does not. The presence of a fluorine atom (smaller than a methyl group) results in a decreased affinity of the same order as produced by the hydroxy group. The most detrimental effect was observed on the substitution of the 5-methyl group with a hydrogen atom (dU).

The same tendency was observed in the 3'-azido series, when 5-ClAZdU and 5-BrAZdU were compared to AZdU and AZT. However, in contrast to TMPKmt, in which the 3'-azido group confers an affinity better than or similar to that of the 3'-hydroxyl counterpart, AZT exhibits a \(K_i\) value for TMPKh 2.5 times higher than that of dT. The introduction of a bulky alkyl group other than methyl at the 5-position (CF\(_3\), CH\(_2\)OH, CH=CHBr) also resulted in decreased affinity. Repulsion was expected from the substitutions of the methyl group by larger groups as a result of the numerous and close contacts observed in the dTMP/TMPKmt complex (Figure 2).

The improved affinity for TMPKmt observed with chloro and bromo derivatives is linked to the presence of the positively charged residue (R74) in the vicinity of the methyl binding pocket, otherwise made up only of hydrophobic side chains (F36, P37). While similar residues are observed at equivalent positions in TMPKh (R76, F42, and P43 respectively), some variability is observed in the second shell of residues surrounding the previously mentioned amino acids (e.g.: R45, N75, and D121 in TMPKmt instead of Y39, D73, and E124 in TMPKmt). These latter residues do not interact directly with the substrate but rather stabilize the side chains of amino acids directly involved in base recognition. The substitution of second shell residues might affect the flexibility of the substrate binding pocket and/or the environment characteristics (hydrophobicity or dielectric constant). These substitutions might explain the distinct ordering of the 5-substituted analogues according to their affinity for TMPKh compared to that for TMPKmt. Moreover, these results suggest only little rearrangement in the vicinity of the 5-methyl group upon binding to TMPKmt. Only isosteric and isoelectronic substituents (Cl or Br) with respect to the methyl group yielded inhibitors active below 10 \(\mu\)M. However a remarkable property of these inhibitors resides in their relative specificity, with the highest SI (around 40) in favor of TMPKmt.

Nature of the heterocyclic moiety

While the phosphate group was not essential for binding of dT analogues to TMPKmt, comparison with enterobacterial orthologues suggested that the aromaticity of the base moiety was important in the case of TMPKmt because of ring stacking with F70.\[16\] In TMPKh, the equivalent F72 is also believed to stabilize the planar and aromatic base moiety. As expected, a decrease in or the loss of the aromaticity of the base moiety results in lower affinity for TMPKmt: dhdT and dhdU are both less effective than dT (Table 2).

Purine and pyrimidine nucleosides were also tested on TMPKmt and TMPKh (Table 2): dC and 5MedC bind poorly to TMPKmt; the presence of a positively charged amino group at C4 instead of a carbonyl group appears to be detrimental. This effect is less important in the case of TMPKh, which has a better affinity for 5MedC (SI = 0.5), despite little structure change in the vicinity of the charged amino group (see above), and might be further enhanced in TMPKmt by the accompanying rearrangement of the neighboring N100, which is hydrogen-bonded to the hydrogen-bearing nitrogen N3 of dT. Again, the presence of the 5-methyl group seems to improve the inhibitory effect. Surprisingly the purine nucleosides dG and dA were equally effective inhibitors of TMPKmt, with \(K_i\) values 2.5–4.7 times lower than those of cytidine analogues (5MedC or dC, respectively). On the other hand, dI inhibits TMPKmt in the same concentration ranges as 5MedC. The inhibition of TMPKmt by dG was further characterized and shown to be noncompetitive with respect to ATP and competitive with respect to dTMP (data not shown), which indicates a specific interaction of dG with the dTMP binding site. As already reported for other TMPKs,\[16\] dGMP was not a substrate for TMPKmt but was a competitive inhibitor with a \(K_i\) value of 117 \(\mu\)M. Other guanosine analogues modified in

Figure 2. Orientation of dTMP in the active site pocket of TMPKmt. View from the 5-methyl group, showing the neighboring side chains of residues F36 and R74 and the main chain atom of P37. Van der Waals contacts involving the methyl group and hydrogen bonds with the nucleotide base atoms are represented as black lines, and the protein backbone as blue ribbon. Side chains of interacting residues are labeled, and the three water molecules involved in substrate binding are shown.
the sugar part of the molecule were tested on TMPKmt: acyclovir (ACV) and ganciclovir (GCV) are less potent than dG (by factors of 2 and 4, respectively). Positioning of the purine analogues in TMPKmt was based on conformations observed in their respective complexes with HSV-1 TK, with the ribose moiety as a main anchor. The impact of ribose moiety modification was analyzed independently by evaluation of the potential interaction changes. Little rearrangement was required for the binding to TMPKmt of acyclic polyols instead of ribose, while numerous interactions were lost, in agreement with the decreased affinity. In contrast, accommodation of the bulky purine base moieties implies significant rearrangements of the substrate-binding pocket, in particular reorientation of the N100 side chain. Favorable interactions might compensate for the cost of these conformational changes as well as discrimination between dI and dG (putative hydrogen bonding involving the amino group N4 and the hydroxy group of S103).

Three C-nucleosides (ψdU, ψdT, dmψdU) related to dT and dU were selected, as these compounds are more stable in vivo than their corresponding N-nucleotides. The carbon–carbon bond between the base and the sugar moiety has an increased stability over the glycosidic carbon–nitrogen bond. Consequently, C-nucleosides are not degraded by nucleosidases. Inhibitory data for these molecules are given in Table 2. dmψdU exhibits a $K_i$ value of over 5 mM, which might be explained by van der Waals clashes between one of the two methyl groups and the N100 amide group. ψdT and ψdU are known to be isosters of dU and dT, respectively,[19] with identical $pK_a$ values. Surprisingly, ψdT exhibits a poorer affinity for TMPKmt than ψdU. Previously, the presence of a methyl group or an isosteric and isoelectronic substituent at the S-position had yielded a better affinity, as in the case of dT versus dU or of dhdT versus dhdU. A conformational analysis in terms of sugar puckering (N/S population) and position of the base relative to the sugar (syn/anti orientation) was therefore conducted for ψdU and ψdT. The structural analysis, by high-resolution proton NMR spectroscopy, was based on evaluation of interproton distances and analysis of coupling constants. Interproton distances were evaluated by off-resonance ROESY experiments to avoid Hartmann–Hahn artefacts.[20] To evaluate the percentage of C2’ endo (S) and C3’ endo (N) conformers, a two-state analysis of sugar proton coupling constants, taking coupling constant sums into account, was conducted as described by Rinkel et al.[21] Both for ψdU and for ψdT, this leads to about 70–80% of C2’ endo (S sugar) conformers with a phase angle around 140° (anti conformers). Although the population of the S conformer is quite high, it is in the expected range for deprotonated C-nucleosides.[22] The similar conformations of ψdU and ψdT observed in solution did not shed more light on their respective inhibitory potencies. An induced-fit rearrangement of the more flexible C-nucleosides or of some protein side chains might therefore be involved. Indeed, ψdT is the only compound so far tested on TMPKmt that is capable of interacting suitably with R74 and N100 in both anti and syn conformations. In the syn conformation, the carbonyl oxygen atom O2 would point toward the 5’ hydroxy group of the nucleoside, at hydrogen bonding distance. In dC and dU, such a conformation would place a carbon atom in front of the N100 amide group.

**Table 2. Base moiety screening.**

<table>
<thead>
<tr>
<th>Compound</th>
<th>TMPKmt</th>
<th>TMPKh</th>
<th>Compound</th>
<th>TMPKmt</th>
<th>TMPKh</th>
<th>Compound</th>
<th>TMPKmt</th>
<th>TMPKh</th>
</tr>
</thead>
<tbody>
<tr>
<td>dU</td>
<td>1020</td>
<td>2550</td>
<td>dhdU</td>
<td>2850</td>
<td>N.D.</td>
<td>dA</td>
<td>280</td>
<td>N.I.[3]</td>
</tr>
<tr>
<td>dT</td>
<td>27</td>
<td>180</td>
<td>dhdT</td>
<td>475</td>
<td>N.I.[5]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>dC</td>
<td>1320</td>
<td>N.D.</td>
<td>ψdT</td>
<td>1320</td>
<td>N.I.[6]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5MedC</td>
<td>690</td>
<td>350</td>
<td>ψdU</td>
<td>94</td>
<td>N.L.[5]</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

[a] As in Table 1. N.D., not determined; N.I., no inhibition detected at a final concentration of [b] 2 mM, [c] 4 mM, and [d] 8 mM.
Conclusions

This study has established that, for different nucleosides with variations mainly on the base moiety, the best inhibitors of TMPKm are 5-halogenated thymidine analogues. Moreover, these compounds are highly specific for TMPKm in comparison with TPKh. Future investigations should aim to control the fate of these analogues in cells. 5-Halogenated 2'-deoxycytidine compounds are substrates for both TK1 and TK2.\(^{2,23}\) To avoid the phosphorylation of these compounds to their monophosphate derivatives by cellular TKs, other modifications on the sugar moiety would be necessary. One possibility would be modification at the 5'-position: in this case, the compounds would no longer be substrates for TKs, with the drawback that they might become inhibitors of TKs. Another possible way would be modification at the 2'-position. It is known that 2'-chloro-2'-deoxycytidine is not a substrate for TK1 and TK2, nor is it an inhibitor of these enzymes.\(^{2,24}\)

Experimental Section

Chemistry: \(^1\)H and \(^13\)C NMR spectra were recorded in DMSO-d\(_6\) on a Bruker 400 MHz instrument. Exact mass measurements were performed by J. Rozenski (Rega Institute for Medicinal Research, Leuven) on a quadrupole/orthogonal-acceleration time-of-flight tandem mass spectrometer equipped with a standard electro spray ionization (ESI) interface. Samples were injected in MeOH/H\(_2\)O (1:1). Thin layer chromatography (TLC) was run on Merck silica gel (F\(_254\)). Silica gel column chromatography was carried out on Merck 9385 silica gel. Unless otherwise stated, all chemicals were reagent grade, purchased from commercial sources and used without further purification.

2'-Deoxyadenosine (dA), 2'-deoxyctydine (dC), 2'-deoxyguanosine (dg), and thymidine (dT) were purchased from Chemimpex. 2'-Deoxycytosine (dCyd) was synthesized by treatment of cytosine with acetic anhydride, followed by bromination in acetic acid and subsequent chlorination with an electrophilic bromine reagent and sodium chloride. 2'-Deoxyguanosine (dG) was synthesized by treatment of guanine with acetic anhydride, followed by bromination in acetic acid and subsequent chlorination with an electrophilic halogen reagent and sodium chloride. 2'-Deoxycytosine (dCyd) was synthesized by treatment of cytosine with acetic anhydride, followed by bromination in acetic acid and subsequent chlorination with an electrophilic bromine reagent and sodium chloride. 2'-Deoxyguanosine (dG) was synthesized by treatment of guanine with acetic anhydride, followed by bromination in acetic acid and subsequent chlorination with an electrophilic halogen reagent and sodium chloride. 2'-Deoxythymidine (dThd) was synthesized by treatment of thymidine with acetic anhydride, followed by bromination in acetic acid and subsequent chlorination with an electrophilic halogen reagent and sodium chloride. 2'-Deoxyuridine (dU) was synthesized by treatment of uracil with acetic anhydride, followed by bromination in acetic acid and subsequent chlorination with an electrophilic halogen reagent and sodium chloride. 2'-Deoxyuridine is not a substrate for TK1 and TK2, nor is it an inhibitor of these enzymes.\(^{2,24}\)

3'-Azido-2'-3'-dideoxy-5-chlorouridine (SCIAzD) was synthesized by chemical modification of the 5-chlorouridine analogues previously described.\(^{25}\)

3'-Azido-2'-3'-dideoxy-5-bromouridine (SBBzAzD) was synthesized by treatment of azidolute with acetic anhydride, followed by bromination in acetic acid and subsequent removal of the acetyl group with methanolic ammonia.\(^{20}\) SBBzAzD was isolated by chromatography on a silica gel column, followed by HPLC on a C18 reversed-phase column (10–25% linear gradient of acetonitrile in 10 mM TEAA over 20 min) in 68% yield. \(R_t = 13.7\) min; \(^1\)H NMR (DMSO-d\(_6\)): \(\delta = 2.32\) (m, 1H; H2'), 2.45 (m, 1H; H2'), 3.60 (m, 1H; H5'), 3.70 (m, 1H; H5'), 3.85 (m, 1H; H4'), 3.48 (m, 1H; H3'), 5.30 (br s, 1H; 5'-OH); 6.03 (dd, \(J = 5.4, 1.0\) Hz, 1H; H1'); 8.30 (s, 1H; H6'); 11.8 (brs, 1H; NH) ppm; \(^{13}\)C NMR (DMSO-d\(_6\)): \(\delta = 37.79\) (C2), 61.03 (C5'), 96.57 (C5), 141.04 (C6), 150.56 (C2), 160.03 (C4) ppm. HRMS (ESI-MS): calc for C\(_9\)H\(_{10}\)BrN\(_5\)O\(_4\)Na\(^+\): 353.9814; found: 353.9815 (M+Na\(^+\)).

High-resolution NMR spectroscopy: High-resolution \(^1\)H NMR spectra for the conformational analysis of \(\psi\)-d\(_6\) and \(\psi\)-d\(_7\) were recorded on a Varian Inova 500 MHz spectrometer equipped with a triple resonance, pulsed field gradient probe with an actively shielded z gradient. The concentration of the samples was 15 m\(_\text{w}\) and the pH value (7.6) was in the same range as in the kinetics measurements. All spectra were recorded at 30 °C in D\(_2\)O. Chemical shifts are given with respect to sodium 2,2-dimethyl-2-silapentane-5-sulfonate (DSS) as internal reference. DQF-COSY, TOCSY, and off-resonance ROESY spectra were recorded with a spectral width of 4 kHz in both dimensions, a 90° pulse of 4.4 ms, 2 K (F1) × 2 K (F2) point data sets, zero-filled to 4K in the F1 dimension and by use of 16, 8, and 32 scans per increment, respectively. The off-resonance ROESY experiment was recorded with a 11.4 kHz effective spin lock field generated by a series of 30° pulses over 400 ms with a recycle delay of 5 s. To avoid Hartmann–Hahn artefacts,\(^{20}\) the offset of the spin lock carrier was shifted by approximately 8 kHz from the centre of the spectrum in order to create an angle of 54.7° between the effective spin lock axis and the static magnetic field. These \(^1\)H NMR experiments were processed by use of shifted sine-bell windows in both dimensions.

Cloning of the human tmk gene and purification of TMPK\(_h\) overexpressed in Escherichia coli: The 636-bp fragment corresponding to \(\text{tmk}\) gene coding for \(\text{TMPK}\) was amplified by PCR\(^{29}\) with cDNA from SK melanoma cells as the matrix. The two synthetic oligonucleotides used for amplification were 5'gatatgatagtcccggccgcccggg-3' and 5'ccgctgtcagtacctcatgagctccccccag-3'. During amplification, NdeI and Xhol restriction sites (in bold letters in the oligonucleotide sequences) were created at both ends of the amplified fragment. After digestion by NdeI and Xhol, the amplified gene was inserted into the pET28a plasmid (Novagen, Inc.) digested with the same enzymes. Two clones containing the \(\text{tmk}\) gene and overexpressing \(\text{TMPK}\) with a His-Tag at the N-terminal end were characterized. One of these clones was kept for further studies and the corresponding plasmid was named \(\text{pHLSO-7}\). The DNA insert was sequenced by the double-stranded dideoxynucleotide sequencing technique\(^{27}\) in order to verify the absence of any mutational events in the course of amplification. The BL21(DE3)/pDIA17 E. coli strain\(^{20}\) transformed with pHL50-7 plasmid, was grown in 2YT medium containing chloramphenicol (30 \(\mu\)g \(\text{mL}^{-1}\)) and kanamycine (70 \(\mu\)g \(\text{mL}^{-1}\)) until the absorbance at 600 nm, A\(_{600}\) reached 1.5. After induction with 1 \(\text{mM}\) isopropyl-\(-\beta\)-thiogalactopyranoside and growth for 3 h at 37 °C, cells were harvested, resuspended in buffer A (50 \(\text{mM}\) phosphate buffer, pH 7, containing 300 \(\text{mM}\) NaCl, 5 \(\text{mM}\) imidazole, and protease inhibitors (Complete ethylendiamine-tetraacetate-free, Roche)) and broken by sonication. After centrifugation at 14,000 rpm for 30 min at 4 °C, the supernatant was placed on a TALON column (Clontech Laboratories, Inc.) pre-equilibrated with 376-400 mM imidazole and washed with 600 mM imidazole. The protein fraction was eluted with 500 mM imidazole. The purity of the recombinant protein was determined by SDS-PAGE and determination of the absorbance at 260 nm. The protein was concentrated to 1 mg/ml using ultrafiltration (Amicon, Millipore Corp.).
with buffer A. The column was extensively washed with buffer A and the protein was eluted with phosphate buffer (50 mM, pH 7) containing NaCl (300 mM) and imidazole (150 mM). Fractions with enzymatic activity were pooled and dialysed against tris(hydroxymethyl)aminomethane (Tris)-HCl (50 mM, pH 7.4) and 20% glycerol and kept at −20 °C.

**Enzymatic assays**: Activity was determined as described previously\(^{13}\) by the coupled spectrophotometric assay at 334 nm in an Eppendorf ECOM 6122 photometer. The reaction medium (0.5 mL final volume) contained Tris-HCl (50 mM, pH 7.4), KCl (50 mM), MgCl\(_2\) (2 mM), NADH (0.2 mM), phosphoenol pyruvate (1 mM), and lactate dehydrogenase, pyruvate kinase, and nucleoside diphosphate kinase (2 units each). 1 unit activity corresponds to 1 mmole product formed in 1 min at 30 °C and pH 7.4. The concentrations of ATP and dTMP were kept constant at 0.5 mM and 0.05 mM, respectively, whereas the concentrations of analogues varied between 0.005 and 8 mM. Equation (1) was used to calculate the \(K_s\) values with the aid of Equations (2) and (3) (classical competitive inhibition model following the Lineweaver–Burk representation):

\[
K_s = \frac{K_m}{v_0 - v(1 + S/K_s)}
\]

\[
v = \frac{V_m[S]}{S + K_m}
\]

\[
v_i = \frac{V_m[S]}{S + K_m(1 + I/K_i)}
\]

where \(v\) and \(v_i\) are the reaction velocities in the absence and in the presence of the analogue at a concentration \([I]\), respectively, \(K_s\) is the \(K_s\) value for dTMP (4.5 μM for TMPKmt and 5 μM for TMPh), and \([S]\) is the concentration of dTMP (50 μM).

**Molecular modeling and docking studies**: Three-dimensional structure visualization and manual superimposition were performed with the XnMol software.\(^{32,32}\) Structure-driven sequence alignments and sequence-dependent structure superpositions were performed by use of in-house software.\(^{32,32}\) Simple substitutions were performed on the substrate molecule in the dTMP(TMPhKmt) complex.\(^{15}\) The more divergent analogues (compounds with a base or a sugar moiety different from thymine or ribose, respectively) which includes those crystallized in complexation with HSV-1 TK, were docked in the crystal structure of TMPhKmt by using the similarity with the original substrate. This allowed a visual inspection of the structures of the complexes and analysis of the ligand–protein contacts.

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