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Ribonucleotide Reductases: Structure, Chemistry, and Metabolism Suggest New Therapeutic Targets

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Abstract
Ribonucleotide reductases (RNRs) catalyze the de novo conversion of nucleotides to deoxynucleotides in all organisms, controlling their relative ratios and abundance. In doing so, they play an important role in fidelity of DNA replication and repair. RNRs’ central role in nucleic acid metabolism has resulted in five therapeutics that inhibit human RNRs. In this review, we discuss the structural, dynamic, and mechanistic aspects of RNR activity and regulation, primarily for the human and *Escherichia coli* class Ia enzymes. The unusual radical-based organic chemistry of nucleotide reduction, the inorganic chemistry of the essential metallo-cofactor biosynthesis/maintenance, the transport of a radical over a long distance, and the dynamics of subunit
interactions all present distinct entry points toward RNR inhibition that are relevant for drug discovery. We describe the current mechanistic understanding of small molecules that target different elements of RNR function, including downstream pathways that lead to cell cytotoxicity. We conclude by summarizing novel and emergent RNR targeting motifs for cancer and antibiotic therapeutics.

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INTRODUCTION
The availability of adequate and balanced deoxynucleotide pools is essential for accurate DNA replication and repair and, consequently, for genome stability. Deoxynucleotides are supplied universally in all organisms by a de novo pathway catalyzed by ribonucleotide reductases (RNRs) that convert RNA building blocks to DNA building blocks (1–3). Deoxynucleotides can also be generated in an organism-, environment-, and disease-specific fashion by nucleoside (or nucleotide) salvage pathways (4). Our current understanding of the unique organic (5) and
inorganic chemistry (6) of RNRs has been revealed, in part, by our understanding of clinically used therapeutics that target the universal radical-mediated nucleotide reduction mechanism and of the specific metallo-cofactor biosynthetic and repair pathways. An ensemble of studies led to the first structures of class I RNRs at low resolution (7–10) and, more recently, to high-resolution structures in trapped active and inhibited states (8, 11, 12). These recent studies suggest, in combination with inhibitors of specific signaling pathways downstream of RNR, that the time is right to revisit RNRs as a target for antibacterial, antiviral, as well as anticancer agents.

All RNRs catalyze the conversion of nucleoside 5′-diphosphates (NDPs) or triphosphates (NTPs) to deoxynucleotides (dNDP or dNTP) (Figure 1a). The RNRs share a common active-site architecture located in subunit α that houses three essential cysteines (Figure 1b) (13). Two cysteines (on the top face of the RNR) provide the reducing equivalents to make dNDPs, and the third cysteine (on the top face of the RNR) is transiently oxidized to a thyl radical (–S•) that initiates NDP reduction (14). Distinct metallo-cofactors catalyze this oxidation (Figure 1c), and they are the primary basis for RNR classification (Ia–e, II, III), although a recently discovered non-metallo-cofactor, 2,3-dihydroxy-phenylalanine radical (DOPA•), breaks this paradigm (15–17). This review focuses on the class I RNRs, which share a distinct mechanism by which a transient –S• is generated and whose formation requires a second subunit, β, that houses the cofactor oxidant (Figure 1c) (18, 19).

Docking Model and Radical Transfer Pathway

In 1969, Reichard and colleagues discovered the class Ia Escherichia coli RNR and proposed that the active enzyme is an αβ2 complex (20, 21). However, not until 1994 was the X-ray structure of α2 reported, by Uhlin & Eklund (13) (Figure 2b); this structure, together with their earlier structure of β2 (Figure 2a) (19), led to a symmetrical docking model based on subunit shape complementarity (Figure 2c) (13). This model guided experimentation until recently. A fascinating feature of the docking model is that the differic-tyrosyl radical cofactor (Fe3+·-Y122•) (Figure 2c) in β is ~35 Å away from C439 (in E. coli numbering), which is oxidized in the α subunit. The turnover frequency for dNDP production (2–10 s⁻¹), together with the long distance between Y122• and C439, engenders a radical transfer (RT) pathway (7, 18): Y122•[β] = [W48[β]] = Y136[β] to Y131[α] = Y159[α] = C439[α] (Figure 3) (note that [W48] involvement has not yet been demonstrated).

One or more rate-limiting physical steps mask both NDP reduction and RT chemistry. These processes are conformationally gated by proper substrate and effector binding to α2 and its association with β2 (22). The stable Y122• in β2 is transiently reduced and reoxidized on each turnover, and RT through the pathway involves distinct proton-coupled electron transfer (PCET) steps at each pathway residue (Figure 3). The first step in RT is proposed to occur at the metal cofactor in β2, triggered by substrate and effector binding in α2 more than 35–40 Å away. Studies using site-specifically incorporated tyrosine analogs with altered reduction potentials; high-field, multifrequency electron paramagnetic resonance (EPR) methods; structural analysis; and RT photoinitiation in photosensitized RNRs (photo-RNRs) have provided insight into each of the proposed steps (7). Our current understanding of this pathway suggests that the thermodynamic landscape of the RT process (Y122• to C439) is uphill by greater than 200 mV and that the NDP reduction reaction, which also involves an uphill 3′-H atom abstraction, is driven to the right by rapid and irreversible loss of water during NDP reduction (Figure 1a). This pathway design avoids buildup of highly reactive protein radical intermediates such as tyrosyl radical (Y•), which has a reduction potential of 0.96 V versus the normal H electrode. Reduction of any of the Y• intermediates in the pathway (Figure 3) would inactivate the RNR, leading to catastrophic consequences for the organism (23, 24). Accordingly, the RT pathway provides a target of opportunity for future drug design.

α: the large subunit of RNR (80 kDa) that in active class Ia RNR is likely α2

–S•: thyl radical

β: the small subunit of class Ia RNR (47 kDa) that is always β2

Radical transfer (RT): the reversible 35 Å pathway between Y122• in β and C439 to be oxidized in α that involves multiple amino acid radical intermediates

Proton-coupled electron transfer (PCET): a central mechanism involved in oxidation/reduction chemistry in biology

Photosensitized RNR (photo-RNR): an αβ2 complex in which β is modified with a photooxidant that generates, with light, the Y136•-β in the RT pathway; this method allows the uncoupling of conformational gating and measurement of the chemistry within α

Y•: tyrosyl radical, a one-electron-oxidized tyrosine with proton loss
Figure 1
(a) Ribonucleotide reductases (RNRs) catalyze the conversion of nucleoside 5′-diphosphates (NDPs) or triphosphates (NTPs) to deoxynucleotides (dNDP or dNTP). (b) The reduction occurs in the active site in subunit α, composed of a 10-stranded β-barrel with three cysteines and conserved placement of the oxidant (gray circle) involved in thyl radical formation (–S•; top face in panel a) that initiates NDP reduction. The bottom-face thiols in panel a deliver the reducing equivalents and themselves become oxidized. (c) The oxidants are distinct among the RNR classes (I, II, and III), represented here by a gray circle juxtaposed with the –S• loop. The substrate and four essential residues, including the three essential cysteines and E441, are shown as sticks. The class Ia RNRs use a diferric-tyrosyl radical (Y•) cofactor (M1, M2 = Fe³⁺) that is located in subunit β (left, bottom) to generate a radical species in the active site in subunit α. The oxidation occurs over a distance of ~35 Å by long-range radical transfer (RT) to first generate a Y• in subunit α (under the gray circle in panel b) and then generate –S• on an adjacent cysteine (top face in panel a). In other class I RNRs (Ib–Ie), oxidation also occurs by long-range RT across α and β but involves distinct metallo-oxidants (X, M1, M2). In class II and III RNRs, the 5′-deoxyadenosyl radical generated from adenosylcobalamin (class II), and the glycyld radical (class III) generated from S-adenosylmethionine and an FeS cluster are located adjacent to the cysteine to be oxidized (gray circle in panel b). Abbreviations: A, adenine base; TRR, thioredoxin reductase.

Evidence for the docking model (Figure 2c) has been provided by trapping pathway radicals using tyrosine analogs with perturbed reduction potentials (Figures 3 and 4a) and active-site radicals in α2 generated using mechanism-based inhibitors (MBIs) (Figure 4b) (25). Pulsed electron–electron double-resonance (PELDOR) spectroscopy and negative-stain electron
Figure 2
Structural models of the class Ia ribonucleotide reductase (RNR) from *Escherichia coli*. (a) X-ray structure of β2 (19), a homodimer (red/orange) with disordered C-terminal tail residues (341–375; dashed lines). (b) X-ray structure of α2 (13), a homodimer (light/dark blue) with disordered C-terminal tail residues (737–761; dashed lines) that houses the two cysteines (red balls) that rereduce the active-site disulfide formed on NDP reduction (see Figure 1a). α2 also houses the A site (activity site or cone domain) that binds ATP (which activates the RNR) or dATP (which inactivates the RNR) (green); the C site (catalytic site) that binds cytosine, uridine, guanosine, and adenosine 5′-diphosphates (CDP, UDP, GDP, and ADP; collectively, NDPs) (magenta); and the S site (specificity site) that binds the effectors deoxyadenosine, adenosine, thymidine, and deoxyguanosine 5′-triphosphates (dATP, ATP, TTP, and dGTP) (yellow). (c) The Eklund docking model of α2β2 (13) with the long-range radical transfer pathway (left) (18). Also shown is a peptide (residues 360–375 of β2; gray) that is proposed to represent the tail of β2 responsible for α2 binding. (d) A cryo–electron microscopy (cryo-EM) structure of an active α2β2 complex with two mutations in β2: F3Y122 and E52Q (11). The asymmetric complex forms when F3Y122/E52Q-β2 interacts with α2, GDP, and TTP. The 3.6-Å-resolution cryo-EM density structure is shown in transparent gray. This structure of the active α2β2 can be compared with the symmetric docking model in panel c.

Microscopy (EM) (Figure 2c) have enabled spectroscopic and structural analysis, respectively, of these trapped α2β2 complexes (see the next section).

3-Aminotyrosine–RNR and Pulsed Electron–Electron Double-Resonance Analysis: Low-Resolution Evidence for the Docking Model
3-Aminotyrosine (NH₂Y) is easier to oxidize than Y by 590 mV (Figure 4a). When NH₂Y replaces a pathway Y in α or β and is incubated with the second subunit, substrate, and specificity effector,
Mechanism-based inhibitor (MBI): nucleotide that binds to the active site of an RNR, leading to its \( \beta \)-C-H bond cleavage by hydrogen atom abstraction and subsequently to distinct radical chemistry and enzyme inactivation

Pulsed electron-electron double resonance (PELDOR) spectroscopy: a paramagnetic resonance method that allows measurement of distances (15 to 100 Å) between two paramagnetic species that experience weak dipolar interactions
Structures of unnatural amino acids and nucleotide analogs used to study class Ia ribonucleotide reductases (RNRs). (a) Unnatural amino acids that have been site-specifically incorporated in place of the tyrosines (Ys) or cysteine within the radical transfer pathway (see Figure 3) as well as their reduction potentials versus Y for NO\textsubscript{2}Y, 2,3-dihydroxy-phenylalanine (DOPA), 3-aminotyrosine (NH\textsubscript{2}Y), and fluoride-substituted tyrosines (F\textsubscript{n}Ys) (24) and versus cysteine for seleocysteine (Sec) at pH 7. NH\textsubscript{2}Y is 590 mV easier to oxidize than Y. Fluorinated Ys (F\textsubscript{n}Ys, where \( n = 2 \) or 3) enable tuning of the reduction potential over 170 mV depending on the number of Fs and their substitution pattern. (b) Nucleoside 5'-diphosphates can be irreversible and reversible inhibitors of RNR. The irreversible inhibitors are mechanism-based, as the 3' C-H bond (red) of the inhibitor must be cleaved, as with the normal substrate (see Figure 1a), before distinct radical chemistry in each case occurs, causing enzyme inactivation. The nucleosides (FC, N\textsubscript{2}C, F\textsubscript{2}C, VFC, CIF, CIA, FIU) are utilized therapeutically and are metabolized to the diphosphates (PPO). The inhibitors of RNR are shown.
**Figure 5**
Support for the Eklund docking model (see Figure 2c). (a) Pulsed electron–electron double-resonance spectroscopy used to measure distances between Y_{122}• in the unreacted α/β pair (right) and the trapped radicals (NH$_2$Y• or N•) in the reactive α/β pair where Y_{122} is reduced (YOH) (left). (b) Representative negative-stain electron microscopy 2D class averages of the structures of the NH$_2$Y$_{730}$• trapped in an α$_2$β$_2$ complex (25). The view with the yellow star resembles the Eklund docking model shown in Figure 2c.

**HIGHER-RESOLUTION STRUCTURES OF RNRs**

**Inhibited Structures In Vitro**

dATP is a universal inhibitor of all class Iα RNRs. It binds to the N-terminal domain of α (Figure 2b). Two independent studies by the Dealwis and Walz groups (8) and the Drennan and Asturias groups (27) in 2011 and 2018, respectively, revealed structures of eukaryotic dATP-inhibited states. In a study of *Saccharomyces cerevisiae* RNR, Fairman et al. (8) observed an α$_6$ hexameric ring structure crystallographically (6.6 Å). In a study of human α with CDP, dATP, and a small amount of ATP, Brignole et al. (27) observed by cryo–electron microscopy (cryo-EM) a similar but higher-resolution hexameric ring structure (3.3 Å) (Figure 6b). Both groups also reported negative-stain EM studies (28 and 30 Å) of a dATP-inhibited state in the presence of both α and β (1:1). Despite the stoichiometry of the subunits, in both cases less than one β$_2$/α$_6$ was observed. Both eukaryotic dATP-inhibited states are a trimer of dimers, with the cone domains responsible for the dimer interfaces. In addition, in human RNRs, small-angle X-ray scattering data on this state suggested that β$_2$ could not enter the hole in the α$_6$ ring structure, implying that an active α$_2$β$_2$ is not accessible (28).

The structure of the dATP-inhibited *E. coli* Iα RNR generated from α:β (1:1) in the presence of dATP is distinct from its eukaryotic counterparts. Using a variety of biophysical methods, Drennan and colleagues (29, 30) reported an α$_4$β$_4$ ring complex with alternating α$_2$s and β$_2$s and a hole in the center (Figure 6a). In this structure, in contrast with the eukaryotic inhibited state, the cone domain interacts with β$_2$. The most intriguing result is that the distance between Y$_{122}$•-β$_2$ and C$_{439}$ in α$_2$ (Figure 2c) has increased from 35 to 60 Å, shutting down RT and, consequently,

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**Abbreviations**

- EM: electron microscopy
- NH$_3$Y: 3-aminotyrosine
- NH$_2$Y•: one-electron-oxidized 3-aminotyrosine
Table 1  Pathway radicals trapped in the *Escherichia coli* class Ia ribonucleotide reductase by site-specifically incorporating unnatural amino acids or reaction with N₃UDP; pulsed electron–electron double-resonance distances (±1 Å) are given in the last column. The red dot corresponds to forward radical transfer (RT) and the blue dot to reverse RT, as indicated by the direction of the arrows on the pathway shown below the table.

<table>
<thead>
<tr>
<th>α</th>
<th>β</th>
<th>Y₁₂₂</th>
<th>Y₃₅₆</th>
<th>Y₇₃₁</th>
<th>Y₇₃₀</th>
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Abbreviations: NDP, nucleoside diphosphate; WT, wild-type.

nucleotide reduction. Thus, despite the distinct quaternary structures of the dATP-inhibited states, a common mechanism of inhibition emerges that involves inability of β₂ to form an active α₂β₂ state.

**Inhibited Structures In Vivo**

The presence of these inhibited states in cells (Figure 6) is important to establish. Drennan and colleagues (31) used their structural insight from the *E. coli* α₄β₄ complex and site-directed mutagenesis to disrupt the α₂/cone domain/β₂ interface. Activity assays and negative-stain EM analysis of several mutants showed that dATP no longer inhibited RNR, and no α₄β₄ was detected. This study, in concert with genetic experiments on *E. coli* using a random mutagenesis protocol, a screen
Figure 6
Structures of dATP-inhibited states of class Ia ribonucleotide reductases (RNRs). (a) The X-ray structure of a dATP-inhibited *Escherichia coli* class Ia RNR (29) is an α4β4 ring structure with a hole in the middle, composed of alternating α2 (light/dark blue, with the cone domains in green) and β2 (orange/red) subunits. Note the importance of the cone domain in the α/β interaction. (b) The cryo-electron microscopy (cryo-EM) structure of a dATP-inhibited human class Ia RNR (27) is a hexameric α6 ring with a hole in the middle. The α subunits are in light and dark blue, the cone domain is in light and dark green, and a three-helix insertion is in purple (residues 638–681). Note the importance of the cone domain in the α/α interactions. (c) Cryo-EM structure of an α/β (1:1) clofarabine triphosphate (CIFTP)-inhibited human class Ia RNR (27). (Top) Representative cryo-EM 2D class-average images generated from α, β, and CIFTP that show β (arrow) interacting with α. (Middle, bottom) Two views of the 3D reconstruction of the same data set. The bottom image is rotated 90° from the middle image. Only a fraction of the α6 rings in these images have a single and variably positioned β.

for altered dNTP pools, and genome sequencing, identified RNR with mutations at the same interface (32). Together, these biochemical and genetic studies suggest that the dATP-inhibited state of *E. coli* α4β4 occurs in vivo. Studies with clofarabine (CIF) (33) and other nucleoside therapeutic inhibitors of RNR [cladribine (CIA) and fludarabine (FlU)] (34) (Figure 4b) have demonstrated α6 formation in several human cell lines treated with sublethal doses of the nucleosides (35). The distinct inhibitory structures of the Ia RNRs (Figure 6) are discussed further in the section titled Use of Mechanism-Based Inhibitors and Reversible Inhibitors to Understand the Mechanism and Design of New Therapeutics, below.

**Toward Active α2β2 Structures**

Our studies using fluorinated (F) tyrosine analogs (Figure 4a), combined with bioinformatics and the docking model of α2β2 (Figure 2c) to identify residues within the α/β subunit interface including E52 in β, led us to investigate the *E. coli* double mutant of β2 (E52Q and F1Y122*) (11). The F1Y122 substitution allowed trapping of the Y356* on the RT pathway (Figure 3) that resulted in a tighter subunit affinity, that is, the *Kd* < 0.4 nM as compared with 0.2 μM for the control with E52Q-β2 (36). Although incubation of E52Q-β2 with any substrate and effector resulted in a completely inactive RNR, incubation of the double mutant of β2 (E52Q and F1Y122*) with α2 (or His6-α2), GDP, and TTP unexpectedly produced 0.5 equivalents of Y356* and 0.5 equivalents of dGDP, consistent with half-site reactivity (36). The resulting α2 (E52Q and F1Y122*β2) complex gave rise to a near-atomic-resolution (3.6-Å) cryo-EM structure (Figure 2d) that is asymmetric, consistent with half-site reactivity. In line with the biochemistry, the structure of α shown in the left side of Figure 2d has generated a disulfide in the active site that we presume gave rise to dGDP. With the α/β pair shown on the right side of Figure 2d, residues 341–375 in β have been visualized for the first time (Figure 2a). In addition, GDP and TTP are apparent, and the location of the
Resetting ribonucleotide reductases for single and multiple turnovers for 2'-deoxynucleoside diphosphate (dNDP) formation. The model assumes an α:β ratio of 1:1, that Y_{356} is distributed equally between each β of β2, and that the wild-type α2β2 complex is asymmetric. In the absence of external reductants, two dNDPs generated at a rate of 2 s⁻¹ (substrate only) to 5–10 s⁻¹ (substrate and effector) arise from chemistry at each α of α2 (a). Steps b and c are rate limiting and conformationally gated. In an assay in the absence of an external reductant, two additional dNDPs are formed at 0.1 s⁻¹ (g). In the presence of an external reductant such as thioredoxin (TR) and thioredoxin reductase (TRR), under steady-state conditions, step b becomes rate limiting. Other steps include (a) binding of substrate- and effector-bound α2 with β2; (c) rapid radical transfer to generate the C_{419}S•⁻; (d) active site chemistry, dNDP formation, and reverse radical transfer; and (f) reversible reduction of the active site disulfide by the cysteines in the C-terminal tail of α and binding of two additional dNDPs.

Pathway residue Y_{356} is revealed for the first time as part of the entire RT pathway (11) (Figures 2c and 3). Our ability to trap radicals at different residues within the pathway summarized in Table 1 and the increased subunit affinity observed under these conditions suggest that this approach may lead to additional cryo-EM structures that will provide insight into the dynamics of this amazing machine and the switching mechanism between the two α/β pairs.

NEW MECHANISTIC INSIGHT INTO THE CHEMISTRY OF NDP REDUCTION

Model for Disulfide Rereduction and Conformational Gating

In the *E. coli* RNR, the rate-limiting step (or steps) for dNDP formation (Figure 7b,e) is physical, involving conformational changes that mask the chemistry of long-range reversible RT and dNDP.
issues affecting screens to discover RNR inhibitors

The model for RNR-mediated NDP reduction shown in Figure 7 encompasses weak and dynamic subunit interactions that change with subunit and dNTP concentrations. These changes alter RNR’s quaternary structure(s) and activity. These issues are essential to understand to successfully develop E. coli RNR assays in vitro and in vivo for high-throughput screens for RNR inhibitors. The same issues are likely to be encountered with other class Ia and Ib RNRs.

NDP Reduction Mechanism

Figure 8 depicts our current proposed mechanism for nucleotide reduction (5, 43). Steps designated a–e describe the main chemical transformation, whereas the numbers show the proposed chemistry that occurs to the NDP and the active site protein residues, 1–6. The important features are that an –S• (C439, E. coli) initiates reduction of NDP by removal of its 3′-H to form a 3′-nucleotideradical (2) (Figure 8a) (44). E441 facilitates this step by functioning as a general base catalyst for 3′-OH deprotonation (45). This reaction is driven to the right by the rapid, irreversible loss of water (2 to 3) catalyzed by C225 (Figure 8b). The proposal for the reductive half-reaction (Figure 8c,d) is that the 3′-keto-2′-radical generated subsequent to water loss is reduced by PCET to generate the 3′-ketodeoxynucleotide and the three-electron, disulfide radical anion (Figure 8c, conversion of 3 to 4). The disulfide radical anion then reduces 3′-ketone by another PCET step (Figure 8d), in which the proton is supplied by the protonated E441. Finally, the H atom abstracted from the 3′ position of NDP in the first step is returned to the same position (step e) to form dNDP and the C439• that reoxidizes Y122 in β2 on each turnover.

Role of multiple thiyl radicals. The role of the –S• initiator (C439) (Figure 8) was previously established on the basis of studies of the class II adenosylcobalamin-dependent ribonucleotide triphosphatase reductase. An exchange-coupled –S•-cob(II)alamin species, detected by EPR and UV-visible absorption stopped-flow spectroscopies, was shown to be chemically and kinetically competent in deoxynucleotide formation (46, 47). The structural homology and conserved residues in the active site of all RNRs (Figure 1b) have thus been used to infer the universal involvement of –S• in initiating 3′-H atom abstraction (14). Support for the involvement of –S• in the reductive half-reaction (Figure 8c,d) comes from studies with E441Q-α/β/CDP/TTP in formation (22, 37). The observed rate constants (kcat) for dNDP formation in the absence of an external reductant (Figure 7a–e) range from 2 to 5–10 s⁻¹ with substrate and substrate/effector, respectively. In the presence of the physiological reductants (in cells or in steady-state assays), additional conformational changes become rate limiting (1–2 s⁻¹; Figure 7h) and likely involve α/β subunit dissociation, conformational changes associated with the reoxidation of the active-site disulfide by the C-terminal tail of α (Figure 7f–h), or both, and are protein concentration dependent. Different αs have distinct cysteine configurations within their C-terminal tails (Figure 2b) and require organism-specific reductants [e.g., thioredoxin (TR), NrdH, glutaredoxin, thioredoxin reductase (TRR), and glutaredoxin reductase] (38–41).
Mechanism of \(-S^*\) mediated NDP reduction by most ribonucleotide reductases (5). a–f indicate the proposed steps in the reaction, and numbers 1–6 indicate the active-site participants in each step. (a) An \(-S^*\) (C439, Escherichia coli) initiates reduction by removal of the NDP 3′-H. (1, 2) E441 facilitates this step by functioning as a base catalyst for 3′-OH deprotonation. (b) This reaction is driven to the right by the rapid, irreversible loss of water catalyzed by C225. (c, d) The 3′-keto-2′-radical (3) is reduced by proton-coupled electron transfer (PCET) to generate the 3′-ketodeoxynucleotide and the three-electron, disulfide radical anion (4). (d) This species then reduces 3′-ketone by another PCET step. (e) The H atom abstracted from the 3′ position of NDP is returned to the same position to form dNDP and the C439r that reoxidizes Y122 in β2 on each turnover. (f) The rereduction of the active site disulfide is proposed to occur, as described in Figure 7. Note that steps a, c, d, and e involve either H atom transfer (HAT) or PCET processes. Other abbreviation: TR, thioredoxin.

which a disulfide radical anion was spectroscopically identified due to the absence of a required proton from E441 for the PCET (Figure 8d) (48). Although \(-S^*\) chemistry has been proposed for many enzymatic reactions, RNR is the only enzymatic system in which this intermediate has been detected (46, 47).

**Photo-RNRs unmask rate constants for NDP reduction chemistry.** The development of methods to uncouple conformational gating (49, 50) and unmask chemistry (23, 26) has allowed unprecedented insight into active-site chemistry, including \(-S^*\) mediated H atom abstraction (Figure 8a) and the subsequent rate-limiting 3′-ketodeoxynucleotide reduction (Figure 8d, 4 to 5). In the former case, we designed a method for photosensitization of RNRs (Figure 9a) in which a photooxidant, bromomethylpyridyl rhenium(I) tricarbonyl phenanthroline ([Re]), is covalently attached to a single surface-exposed cysteine in the S355C-β2 mutant; the Y122• in β2 is reduced and Y356 in β2 is replaced with a fluorinated tyrosine (F,Y356) (Figure 4a). This photo-β2, in
complex with α2, substrate, and effector, can be rapidly (within nanoseconds) oxidized to an F335Yβ−β2 state upon illumination (Figure 9b). The photochemically generated radical rapidly equilibrates with the RT pathway in α2, ultimately oxidizing C339 and initiating cleavage of the 3′ C–H bond of NDP. Comparison of the F335Yβ− decay, observed by transient absorption spectroscopy in the presence of 3′-[3H]-CDP or independently in the presence of 3′-[3H]-CDP, established a lower limit for the −S• mediated H atom abstraction (Figure 8a) of 1.3 × 10^3 s⁻¹ and an isotope effect of ≥7 (51). Note that the kcat for RNR is 2–10 s⁻¹. The RT chemistry is thus very fast and was revealed for the first time using this method.

The subsequent rate-limiting 3′-ketodeoxynucleotide reduction (Figure 8d) has been examined by incorporating tyrosine analogs with altered reduction potentials in place of Y122 in β. Use of these “hotter” oxidants drives RT and also uncouples conformational gating. Specifically, F1Y122• and NO3Y122• (Figure 4a) have higher reduction potentials than the native Y122• (β2) by 80 and >200 mV, respectively (23, 26), as determined from independent measurements of formal reduction potentials in a small three-helix-bundle protein (24). These β2 mutants have been studied in an effort to observe the slow step(s) within the proposed chemistry, specifically PCET reduction of the 3′-ketone by the disulfide radical anion (Figure 8d). When NO3Y122•−β2 (or F1Y122•−β2) is mixed with α2/CDP/ATP, dCDP formation occurred at ~150 s⁻¹ (or 30 s⁻¹). This rate constant is similar to the 50 s⁻¹ measured for dCTP formation; the latter is catalyzed by class II RNRs in incubated with the E. coli RNR resulted in time-dependent release of Cl⁻ and cytosine and that the

**USE OF MECHANISM-BASED INHIBITORS AND REVERSIBLE INHIBITORS TO UNDERSTAND THE MECHANISM AND DESIGN OF NEW THERAPEUTICS**

The MBIs 2′-halo-(X)-2′-deoxyNDPs (XNDP, where X = Cl or F) (Figures 4b and 10) have played a pivotal role in our current understanding of the mechanism of nucleotide reduction (53). In 1976, Thelander et al. (54) reported that 2′-chloro-2′-deoxycytidine diphosphate (ClCDP) incubated with the E. coli RNR resulted in time-dependent release of Cl⁻ and cytosine and that the
Figure 10

Generic mechanism for 2'-X-dNDP (where X = F, Cl, N₃, or F₂) (see Figure 4) mechanism-based inhibition of ribonucleotide reductases with loss of X⁻ in step b (43, 53). 3 is formed with the bottom-face protonation states of the thiols unknown. The intermediate 2'-nucleotide radical can be reduced from the top face by SH of residue 439 to produce 7 (right) or the bottom face to produce 7 (left) through intermediates 4 and 4'; both of these dissociate from the active site and decompose to generate the products (PPᵢ, base, and 8). 8 can alkylate the α subunit. Alternatively, if in conversion of 2 to 3 (step b) XH is eliminated, then the same product produced by the reduction of NDP, that is, deoxynucleotide (dNDP), is formed. a–c describe the steps proposed for the conversion of the inhibitor to intermediate 7. The numbers 1–4 and 4' represent the proposed structures of the nucleotide intermediates and the protein environment. Other abbreviation: XH, protonated form of the leaving group.

α subunit was inactivated. These observations provided the impetus for studies using isotopically labeled nucleotide analogs, which led to the general model for inhibition shown in Figure 10. As with the NDP substrate, the –S• abstracts the 3'-H (Figure 10a) to generate 2. The outcome of the reaction depends on whether and how the loss of X at 2'-C is catalyzed by the enzyme. From 3, the 2'-delocalized radical can be reduced from the top face (4') by H atom transfer mediated by C₄₃₉ or the bottom face (4) facilitated by C₂₂₅. With Cl(F)NDP, a 3'-ketodeoxynucleotide is generated (7) that dissociates from the active site (when X is not protonated). Intermediate 7 can decompose on a minute timescale to a nucleic acid base, pyrophosphate (PPᵢ), and a furanone (8) that nonspecifically alkylates the α subunit. If the reduction of the nucleotide intermediate in 3 occurs from the top face by C₄₃₉ (4'), then reverse RT can effectively regenerate the Y₁₂₂• in β2. However, if reduction of this same intermediate occurs from the bottom face (4), Y₁₂₂ remains
reduced and β2 is inactivated. Thus, α and/or β can be inactivated via distinct mechanisms. With both XNDPs, if X is protonated (Figure 10b), then dNDP is formed. The details of RNR inactivation in vitro and in vivo depend on the identities of the substrate and effector, leaving group (X), and reductant. In all cases, α inactivation requires Yr reduction. These inhibitors inactivate class I, II, and III RNRs by a common mechanism, suggesting similar active sites (Figure 1b). The involvement of the α C-terminal cysteines in enzyme inhibition (Figure 2b) is not well understood, as their covalent linkage to 8 is reversible, precluding isolation and characterization of alkylated α.

In contrast to XNDPs (X = Cl or F), a number of MBIs (X = N3, F2, or VF) (55–57) (Figure 4b) share similar chemistry in steps a and b (Figure 10), but then undergo distinct chemistry controlled by X and the residues and their protonation states in the active-site cavity. Unraveling the mechanism by which N3NDP inactivates all RNRs has defined the strategy to study the mechanism of action of the clinically used nucleoside therapeutics gemcitabine (F2C) and ClF. F2CDP is an irreversible inhibitor (58), and ClFDP and ClFTP are reversible, noncovalent inhibitors of RNRs (59–61) (Figure 4b).

2′-Azido-2′-Deoxynucleotide

N3NDPs (N = C, U, or A) are MBIs first reported by Thelander et al. (54). Extensive studies with N3UDP (Figure 11a, b) revealed that its incubation with α2β2 resulted in rapid loss of ∼90% RNR activity concomitant with the loss of only 0.5 equivalents of Y122•. The total Y• loss was biphasic; the fast phase was accompanied by formation of N2 gas and a nucleotide-based N•, derived from the N1 moiety of N3UDP. The N• was structurally characterized using isotopically labeled N3UDPs and EPR methods. The nucleotide N• species then slowly decomposes to form a nucleoside base (blue N in Figure 11a), PPi, and 8 (62, 63). The α/β subunits then dissociate, and subsequent to α2β2 complex reformation, additional Y• is lost and more N• is formed. The recent examples of half-site reactivity (Table 1) and RNR asymmetry (Figure 2d) suggest an explanation for the observation that 1 equivalent of inhibitor per α2β2 results in >90% loss of enzyme activity (53).

In vitro, N3CDP inhibits β2 by reduction of the essential Y122•, whereas in vivo, the nucleoside analog 2′-azido-2′-deoxycytidine (N3C) is not cytotoxic. In cells, N3C is not readily

Figure 11

2′-Azido-2′-deoxynucleoside diphosphate (N3NDP) (Figure 4b) is a potent inhibitor of all class Ia ribonucleotide reductases (RNRs) (62, 63). Studies of this inhibitor provided a glimpse of unprecedented chemistry associated with reactive radical species in an active-site cavity and the challenges associated with radical structure elucidation. These studies also provided early evidence for half-site reactivity. The products formed during the inactivation include: N2, NH3, PPi, and the furanone 8, all initiated by 3′–C–H bond cleavage, resulting in 3H2O. (b) The kinetics of loss of RNR activity and tyrosyl radical during the inactivation are biphasic.
phosphorylated to N3CMP by deoxycytidine kinase, demonstrating the importance of the specificity of this and other kinases in generating nucleotide (di- and triphosphate) therapeutics.

**Gemcitabine and Clofarabine: Clinically Used Nucleoside Therapeutics That Inhibit Human RNRs**

The nucleosides F2C and ClF are used clinically as cancer therapeutics (Figure 4b). F2C targets a broad spectrum of solid tumors (pancreatic, metastatic breast, lung) and hematological cancers. In the clinic, this compound is used in combination with DNA damaging agents such as cisplatin or small-molecule inhibitors of signaling pathways that affect the cellular response to DNA replication stress (3, 64–66). ClF is limited to hematological cancers (acute myelocytic leukemia, acute lymphocytic leukemia). Both agents inhibit DNA synthesis. RNR is the upstream target of the diphosphate forms of these compounds (F2CDP, ClFDP), whose inhibition alters dNTP pools. Additionally, the triphosphate forms of these compounds (F2CTP, ClFTP) inhibit DNA polymerases by incorporation into DNA. The mechanisms by which these compounds inhibit RNR and DNA synthesis, however, are distinct.

**F2CDP**

F2C was synthesized independently by two research groups (58, 67). Studies by Plunkett and colleagues (67–69) demonstrated that F2C inhibited growth of a variety of tumor cell lines and that cytotoxicity resulted from inhibition of multiple targets, including DNA polymerases and RNR. Biochemical studies on *E. coli* and human RNR established that F2CDP is a time-dependent irreversible inhibitor and that inactivation occurs with 0.5 equivalent per α subunit. Studies using isotopically labeled F2CDPs established that the products of the inactivation were distinct depending on whether the inhibition studies were carried out in the presence or in the absence of reductant (70–72).

In the presence of reductant (TR/TRR or DTT) subsequent to cleavage of the 3′ C–H bond of the inhibitor, 2F–, cytosine, PPi, and one alkylated α-cysteine (C225) per α2 were identified, and no Y122β in β2 was lost (Figure 12a). Under these conditions, while only 0.5 equivalents of α are inactivated and β remains active, all enzymatic activity is lost. Analysis of the inhibited reaction mixture by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) with no heating of samples revealed that α migrated as a 60:40 ratio of 80 kDa (endogenous α molecular weight) to 110 kDa (modified α) (74). In cancer cell lines incubated with F2C, SDS-PAGE analysis of cell lysate revealed that the α subunit also migrated in a 60:40 ratio (Figure 12b), similar to what was found in the in vitro studies (73). Similar experiments in the absence of reductant resulted in 50% loss of the β2-Y122α and in the formation of an equivalent amount of a new, nucleotide-based radical (Figure 12c). This radical slowly breaks down to cytosine and PPi. Inhibition was accompanied by loss of 2F–, but the α subunit was not covalently labeled. Thus, in both the presence and absence of reductant, 1F2CDP/α2 is sufficient for inhibition, although the underlying mechanisms of inactivation are distinct.

To account for the complete inactivation of RNR with only 0.5 equivalents F2CDP/α, we proposed that the α/β subunit affinity increased and switching to the second α/β pair for additional chemistry is prevented. To test this possibility, inactivated *E. coli* and human RNR were subjected to size-exclusion chromatography (SEC) analysis. The former showed a species consistent with an apparent molecular weight for α2β2 and the latter with α6β6 (74). In the control, in the absence of F2CDP, the subunits separate with β eluting as a dimer and α as a mixture of monomers and dimers consistent with weak subunit interactions. On the basis of our recent EM analyses of a mixture of α and β (1:1) with ClFTP (Figure 6; see also the next section) (27), some α6β2 and no
Gemcitabine diphosphate (F₂CDP) (Figure 4b) inhibits human ribonucleotide reductase by distinct mechanisms in (a,b) the presence and (c) the absence of reductant. (a) Shown are the products of the inhibition carried out in vitro in the presence of reductant (protein, thioredoxin, or small molecule, dithiothreitol) (70–72). (b) SDS-PAGE of RNRs from studies in vitro (a) and cell lysate without boiling after incubation with F₂CDP (in vitro) or gemcitabine (F₂C) (in cells; graphic representation of actual data). In vitro and in cell lines (73), α migrates as a 60:40 mixture of an 80-kDa and a 110-kDa α. (c) Shown are the products of F₂CDP inhibition carried out in the absence of reductant. Abbreviations: RNR, ribonucleotide reductase; SDS-PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis.

α6β6 was observed. Altered molecular weights using SEC analysis can be attributed to unusual, nonglobular shapes (α6β2) or altered quaternary structure(s). For example, fibril structures have been reported with human α and ATP (27) as well as with Bacillus subtilis class Ib (α/β) RNRs (75).

**CIFDP and CIFTP (Reversible)**

Early studies by Plunkett and colleagues established the toxicity of CIF towards many cell lines (CEM, K562, Hep2). In cell-free systems, CIFTP inhibits RNR and DNA polymerases α and ε (59, 60, 66, 76). The observation that the CIFTP:CIFDP ratio in some cells was 7:1 led to the proposal that CIFTP was a reversible inhibitor of ATP binding to the A site of α (Figure 2b). To better understand how RNR is targeted, kinetic and biochemical studies were undertaken with both CIFDP and CIFTP (61). CIFDP was shown to be a reversible, time-dependent, slow-binding inhibitor of the C site. The kinetic analysis revealed a two-step binding mechanism with a $K^*_1$ of 17 nM. CIFTP exhibits reversible, time-independent A-site binding. With CIFTP in five-fold excess relative to RNR under physiological conditions, RNR activity was rapidly and completely lost with a $K_i$ of 40 nM. With sample dilution and follow-up assays, enzyme activity was recovered over 30 min, but only to 50% of the initial value. The half-life ($t_{1/2}$) of human Y• in β2 is 30 min at 37°C (61), and the α subunit is prone to oxidation, making the kinetic measurements challenging; further studies are required. To determine whether the observed inhibition was associated with changes in the RNR’s quaternary structure, CIFTP (CIFDP) with α and with and without an
allosteric effector (dGTP) were each examined by SEC. In the absence of the corresponding nucleotide in the elution buffer, α migrated as α6 in the presence of either CIF(T)P or CIFD(P). This result is distinct from dATP-α6, which reverts to the monomer when dATP is not present in the elution buffer during SEC. Thus, the presence of CIFD(P) or CIF(T)P alters α’s quaternary structure such that, even subsequent to CIFD(T)P dissociation, α6 remains trapped in the inactive state. SEC analysis further showed that β2 had no effect on inhibition or migration. Structures of CIF(T)P mixed with human α/β (1:1) were examined by cryo-EM and solved to 30-Å resolution (Figure 6c). Fewer than one β2 per three α2 was observed, and it appeared randomly positioned outside or on top of a hexameric ring structure (Figure 6c) (27). In support of this model, experiments with D57N-α in which the mutation in the cone domain prevents hexamerization of α revealed that neither CIFD(T)P-treated mutant RNR nor CIF-treated cells with mutant RNR were inhibited. Finally, E. coli RNR, which does not form α6 structures, is not inhibited by CIFD(P) (61).

The dynamics of quaternary structure interconversions offer an opportunity to inhibit RNRS through unconventional mechanisms. The flexible cone domains (Figures 2b and 6a,b) (27, 30) play critical but distinct roles in these states. Strengthening or weakening the interactions responsible for these quaternary structures with small molecules could alter RNR activity. To assess the importance of the hexameric state of human RNR, investigators studied His6-α expressed at 3- or 30-fold (3× or 30×) endogenous levels in COS cells that were then treated with noncytotoxic levels of CIF for 3 h. Analysis of the 30× material purified by Ni-affinity chromatography revealed that the α6 state was present; with 3× endogenous levels, cross-linking was required to detect α6. The α6 state from these and other studies is likely the inhibited state inside the cell in the presence of CIF(T)P and dATP (33).

An extension of this strategy to other adenosine analog therapeutics, CIA and FIU (Figure 4b), has recently been reported (34). In vitro studies of the di- and triphosphates of cladribine (CIADP and CIA(T)P) interactions with human α revealed α6 formation. Further assessment of the hexameric structures and their relationship to cell cytotoxicity is an ongoing challenge. Collectively, results obtained in cells and in vitro with these adenosine inhibitors suggest a potential new way to target RNRS: trapping α in an inhibited state with a small molecule.

**Pleiotropic Modes of Cytotoxicity of Gemcitabine and Clofarabine**

With both F2C and CIF, the mechanisms of cytotoxicity require nucleoside uptake and metabolism (64, 65, 77). As noted above, the diphosphates and triphosphates of F2C and CIF inhibit RNR and DNA polymerases, respectively (60, 78), the latter by chain termination. The consequences of DNA inhibition involving both targets are DNA replication stress that manifests as stalled or collapsed DNA replication forks and DNA single- or double-strand breaks, which can lead to cell cycle arrest, DNA repair, or programmed cell death (64) (Figure 13).

F2CDP, a potent MBI of RNRS, results in lower dNDP and, consequently, dNTP pools. Reduced dCTP, a feedback inhibitor of deoxycytidine kinase (3, Figure 13), enhances production of F2CCTP, leading to elevated levels of F2CCTP. As a result, F2CCTP can more effectively compete with lowered dNTP pools to inhibit DNA synthesis. F2C’s broad spectrum of solid tumor inhibition, distinct from that of other nucleoside therapeutics such as araC, may be associated with pleiotropic metabolic effects (Figure 13), resulting in its self-potentiation (69).

CIF is also phosphorylated by deoxycytidine kinase (Figure 13, 3) and subsequently by distinct kinases to afford CIFD(P) and CIF(T)P. Its stability (due to F/Cl substitution) is increased relative to that of other adenosine analogs (CIA, FIU) (Figure 4b) by its resistance to metabolism by purine nucleoside phosphorylase and adenosine deaminase. Downstream consequences of DNA synthesis inhibition by F2C and CIF are actively being studied. F2C is being investigated in
Cellular responses

**Mechanism of action**

- **F₂U**
- **F₂C**
- **dCTP**
- **F₂CTP**
- **F₂CMP**
- **F₂CDP**
- **F₂CTP**
- **F₂CMP**
- **dCTP**
- **dNDP**
- **dNTP**

**Stabilized forks**
- **Cell cycle arrest**

**Strand breaks**
- **DNA repair**

**Collapsed forks**
- **Cell death**

**Figure 13**

A general scheme for metabolism of nucleosides using gemcitabine (F₂C) as an example (64). F₂C and clofarabine (ClF) therapeutics require cellular uptake and phosphorylation to the appropriate state recognized by target enzymes. The former is mediated by nucleoside transporters ENT1, ENT2, and CNT (1). Once inside the cell, both F₂C and ClF are phosphorylated to the monophosphate by deoxycytidine kinase (3) and subsequently to the di- and triphosphates by cellular kinases (4, 6). Deoxycytidine kinase has unusual specificity in that it phosphorylates both pyrimidines and purines. The concentrations of the monophosphates are in general greater than the triphosphates and much greater than the diphosphates, are cell type distinct, and influence therapeutic outcomes. (2) Cytidine deaminase. (5) Ribonucleotide reductase. (7) DNA polymerase. Inhibition of DNA synthesis (purple box) results in a variety of cellular responses (orange and green boxes).

In combination with DNA damage response inhibitors of checkpoint kinase 1 (Chk1) (64, 68, 79, 80); with inhibitors of ATR, a nuclear kinase that controls S phase progression in response to DNA damage and replication fork stalling, in the same pathway (81); and with DNA repair enzyme inhibitors (65, 82). In addition, F₂C is often used in combination with cisplatin, which enhances DNA damage and alters the downstream consequences. The ability to monitor the consequences of treatment with combinations of therapeutics using genomics, phosphotranscriptomics, and metabolomics has aided and will continue to aid in the development of new approaches (65, 83). For a recent report of alternative functions of h-RNR α and potential effects in therapeutic design, see the sidebar titled A Moonlighting Function of α.

**Reversible C-Site Binders Lacking Phosphoryl Groups**

Two compounds, I and II (Figure 14), have recently been reported to inhibit human RNR by binding reversibly to the C site of α. In contrast to CI-FDP and CIADP (Figure 4b), these small molecules lack the diphosphate moiety thought to be essential for substrate recognition. A 5′-substituted amine of F₂C (I), for example, is reported to inhibit RNR in vitro and in vivo (85). The unusual diphosphate binding site for NDP in α (no lysine, arginine, or Mg²⁺) suggests that amine substitution might avoid issues associated with cellular uptake and phosphorylation (Figure 13).

**A MOONLIGHTING FUNCTION OF α**

A recent report (35) and review (84) provide support for a moonlighting function of α, independent of its ability with β to make dNDP in the cytosol of the cell. Through the use of a C→S mutant that inactivates formation of −Sα in α (Figure 1a), a small amount of α was detected in the nucleus of the cell in an α6 state. Yeast two-hybrid experiments with cDNA from HeLa cells revealed that α interacts with ZRANB3, a protein that forms a complex with PCNA, the sliding clamp that, together with DNA polymerase, promotes DNA synthesis in nonstressed cells. Nucleus-localized α inhibits the interaction of ZRANB3 with PCNA, resulting in inhibition of DNA synthesis. This study (35) may explain the tumor suppressor activity reported for α (3).
Dealwis and colleagues (86, 87) reported a molecule with a naphthyl salicylic acyl hydrazone scaffold that also targets the C site. Medicinal chemistry approaches to reduce hydrolyzability of the acyl hydrazone and increase interaction with the phosphate binding region led to compound II (Figure 14). Decorating the scaffold with an appropriately placed electrophilic moiety such as ClCH₂CO-(R') could result in alkylation of one of the C-site cysteines, analogous to the mechanism of covalent protein kinase inhibitors (88).

**Reversible Inhibitors That Disrupt α/β Subunit Affinity**

The C-terminal tails (30 to 35 amino acids) of all β₂ subunits are disordered (Figure 2a) (18, 89–93), distinct, and predominantly responsible for subunit affinity (Figure 15). Early studies of herpes simplex viruses (HSV-1, HSV-2) (94, 95), which encode for their own RNRs, provide an example of how researchers have successfully developed peptidomimetics of their C-terminal tail that disrupted the α/β subunit interface in vitro and in a murine ocular model of HSV-1-induced keratitis (96). The new structure of *E. coli* α₂β₂ (Figure 2d), which reveals, for the first time, the interaction between the tail (residues 341 to 375) and the α subunit (11), may suggest new ways to disrupt this interface.

A second example of subunit disruption was reported by the Yen group (82, 97), who used the structure of human p53β₂ and computer modeling to identify a pocket in each β subunit close to the C-terminal tail but removed from the buried Fe³⁺-Y* cluster (Figure 1c) essential for β₂ stability. Virtual screening and additional experiments led to the identification of COH29 (Figure 15), which exhibited cytotoxicity to many of the NIH 60 cancer cell lines and caused S-phase cell cycle arrest. COH29 enhanced cytotoxicity of BRAC1-deficient HCC1937 cells. This peptide inhibition of α/β interactions

<table>
<thead>
<tr>
<th>Organism</th>
<th>C-terminal sequence β</th>
<th>Peptide inhibition of α/β interactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herpes simplex virus</td>
<td>YAGAVVNDL</td>
<td>HN2, HO, OOH, 3-AP</td>
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<tr>
<td>Mammals</td>
<td>NSFTLDADF</td>
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<tr>
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</tr>
<tr>
<td><em>S. cerevisiae</em></td>
<td>AGAFTFNEDF</td>
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</table>

**Figure 15**

Targeting the α/β interface of active ribonucleotide reductase to prevent active complex formation with peptidomimetics and COH29. Targeting formation and repair of the M₂⁺⁻Y* (M = Fe; see Figure 1) cofactor of β₂ with hydroxyurea (HU) and triapine (3-AP).
observation provides an example of DNA repair inhibition (98), in this case genetically, that potentiates the effects of the RNR inhibitor COH29.

**BIOSYNTHESIS AND REPAIR OF THE ESSENTIAL DIFERRIC-Y•
COFACTOR OF CLASS Ia RNRs: TARGETING THE β2 COFACTOR**

Whereas ClF and F₂C target α and the α/β subunits of RNRs, respectively, hydroxyurea (HU) and triapine (3-AP; a thiosemicarbazone) (Figure 15) target reduction of the essential Y• cofactor (83, 99, 100) in β2 and/or interfere with cofactor assembly and/or its repair if the essential Y• gets reduced (Figure 1c). HU is used clinically, predominantly in combination with other therapeutics (65), although recent studies have suggested that the RNR is not a key target of its cytotoxicity (99, 101, 102). 3-AP continues to be examined in clinical trials but has not yet been approved for clinical use (83, 100). Although the upstream target of both compounds is the RNR, the downstream pathways that lead to cytotoxicity are pleiotropic and distinct in different organisms. In this section, we focus on HU and 3-AP inhibition of RNRs in vitro and in the early stage of cell culture, when cell viability remains high. Even under these conditions, the detailed mechanism(s) of RNR inhibition requires further exploration.

**Background for Metallo-Cluster Metabolism**

The class Ia RNRs require a Fe³⁺-Y• cofactor in β2 to initiate NDP reduction in α₂, with activity being directly proportional to the concentration of Y• (Figures 1c and 3). The t½ of the Y• in the cluster of different class Ia β2s is variable, ranging from 4 days in *E. coli* at 4°C to 30 min in humans at 37°C. In addition, recombinant expression of β from different organisms results in variable amounts of active cofactor (0 to 1 Fe³⁺-Y•/β2) (6, 103). In general, therefore, the β2 cofactor must be loaded by self-assembly by use of Fe²⁺ and O₂, with variable outcomes (6, 104).

In the past 2 decades, the importance of biosynthetic pathways has been established for FeS cluster cofactor assembly that, in turn, has been linked to formation of mono- and dinuclear nonheme iron cofactors, including the RNR cofactor (105). Although much remains to be learned, genetic studies in *E. coli* and *S. cerevisiae*, as well as in vitro biochemical studies on these class Ia β2s, suggest that there are pathways not only for cofactor biosynthesis but also for its maintenance and activity regulation (Figure 16). Our general model for Fe³⁺-Y• cofactor biosynthesis indicates a requirement for one or more chaperone proteins (106) to alter the apo-β2 conformation for optimized Fe²⁺ loading, an Fe²⁺ carrier protein or small molecule that delivers Fe²⁺ to apo-β2, and a reducing equivalent delivery mechanism required for cluster assembly with O₂ as the oxidant (107). Studies in vivo in *E. coli* (108) and *S. cerevisiae* (109) reveal that cluster assembly can yield β2 in which each β subunit has two Fe³⁺ and one Y•, that is, quantitative loading. In vitro, however, *E. coli* β2 loading gives rise to ~66% active cofactor and 34% inactive diferric clusters with no Y•. In both in vivo and in vitro loading, the activity of the RNR per Y• is the same, suggesting identical cofactor structures.

**Hydroxyurea**

HU (Figure 15) has been studied since the 1960s. On the basis of EPR analyses of prokaryotic and eukaryotic cells and of purified β2 with a self-assembled Fe³⁺-Y•, HU treatment reduces Y• to YOH. In vitro, the iron cluster of human β2 is also reduced (Fe³⁺-YOH), whereas in *E. coli* it remains in the Fe³⁺ state (Fe³⁺-YOH). HU reduction of β2 alone is slow (0.45 M⁻¹ s⁻¹), and there is no evidence that it binds to either *E. coli* or human β2 (107, 109, 110). The chemical mechanism of Y• reduction and the structure of the resulting cluster remain unknown (112).
Figure 16
Model for Me$_2^{3+}$-$\mathbf{Y}$ (Me = Fe or Mn) cofactor biosynthesis (black), maintenance (green), and regulation (red). Factors identified from *Escherichia coli* are in orange, and those from *Saccharomyces cerevisiae* are in blue (6). *S. cerevisiae* counterparts are found in humans. Regulation can occur by endogenous reductants or by therapeutics such as hydroxyurea (HU) and triapine (3-AP) (see Figure 15).

In vitro study by the Sjöberg laboratory (113) has shown that HU-mediated loss of RNR activity is potentiated 10-fold by complexation of β2 with α2, substrate, and effector. This result led these authors to suggest that HU reduction of Y$_{122}$$^•$ is not direct but rather might involve trapping of a transient pathway radical at the α/β subunit interface (Figure 3). Studies of the reduction by HU of the Mn$^{4+}$Fe$^{3+}$-$\beta$2 cofactor in the *Chlamydia trachomatis* class Ic RNR (Figure 1c), an Fe$^{3+}$-$\mathbf{Y}$ surrogate, were also interpreted to suggest that HU intercepts a pathway radical at the α/β subunit interface (112). Furthermore, in the presence of α2, CDP, and ATP, Mn$^{4+}$Fe$^{3+}$-$\beta$2 is reduced by HU to an Mn$^{3+}$–Fe$^{3+}$ cluster with half-site reactivity involving a fast phase and a slow phase, with apparent saturation by HU for the fast phase. These studies support HU binding and targeting of the RT pathway (112, 114). The consequences of the HU-reduced cofactor state in *E. coli* and mammalian cells are still unclear; however, since the proteins identified in *S. cerevisiae* for β2 cofactor biosynthesis and maintenance are also found in mammalian cells (107), Y$^•$ regeneration is a possible fate (Figure 16) and requires further investigation.

RNR inhibition by HU blocks DNA replication. Two papers have suggested that cytotoxicity from extended HU exposure of *E. coli* (101) or *S. cerevisiae* (102) cells is linked to reactive oxygen species (such as HO$^•$)-mediated damage. Vernis and colleagues (102) have shown that HU resistance in *S. cerevisiae* leads to enhanced production of the cytosolic FeS cluster biosynthetic machinery, including Dre2/Tah18. We have demonstrated the importance of these two proteins in the assembly of the β2 cofactor in *S. cerevisiae* (107).

**Triapine**

3-AP (Figure 15) has been extensively investigated since its introduction in the 1990s, and its cytotoxic effects have inspired the synthesis of many additional thiosemicarbazones. However, studies of these analogs reveal that the mechanism of cytotoxicity changes with structure. The complexity arises from their distinct abilities to bind Fe$^{2+}$ and Fe$^{3+}$ (as well as Cu$^{2+}$ and Zn$^{2+}$) and the resultant ligand field–imposed iron redox chemistry (100). Results reported by different
groups (115, 116) in different mammalian cell lines, primarily at late stages of 3-AP treatment, have thus made it challenging to compare and evaluate the outcomes of the different studies.

Our recent studies in cell culture in the early stages of 3-AP treatment provide a framework for thinking about the issues and evaluating its potential as a therapeutic that targets β2 (115). Three mammalian cell lines (K565, COS-1, and HU-resistant TA3) treated with 3-AP and analyzed by whole-cell EPR revealed loss of the RNR Y•, and assays of the corresponding cell lysates revealed loss of RNR activity. Immunoprecipitation of β2 from 55Fe-treated and nontreated cells revealed similar iron content. These and additional studies suggest that Y• loss is the major mode of RNR inhibition, with iron loading remaining unchanged. Although the oxidation state of the bound iron is unknown, we know from in vitro studies that Fe^{2+}-loaded β2 can assemble rapidly into the native Fe^{3+}-Y• cofactor, consistent with a maintenance pathway (Figure 16). In our model, Fe^{2+}-(3-AP) is the active species involved in β2 inhibition, and in the continued presence of Fe^{2+}/Fe^{3+}, the RNR is susceptible to Fe^{2+}-(3-AP) inhibition by direct Y• reduction. In a recent study by Gräslund and colleagues (116), the use of [3H]-(3-AP) and a docking model of 3-AP to mouse β2 resulted in the proposal of a specific 3-AP binding site. However, neither 3-AP nor Fe^{2+}-(3-AP) binding to β2 has been observed. In our opinion, the mechanism of action of these compounds requires further study. Finally, our studies at early stages subsequent to 3-AP treatment, in contrast to other researchers’ later-stage studies, indicate that reactive oxygen species are not responsible for loss of RNR activity.

How 3-AP and HU inhibit RNR and the nature of the relationship between their RNR inhibition and cell cytotoxicity remain a mystery. Although interference with cluster assembly and maintenance might yield effective therapeutics, a better understanding of the biology of Fe^{3+}-Y• pathways is required. However, the recent discovery of Mn^{3+}-Y• cofactors in β2 of class Ib RNRs (Figure 1c) and the identification of a NrdI-β2 interaction essential for both oxidant delivery (O2•−) and active cofactor formation (Figure 16) (6, 117) suggest that disruption of this protein/protein interface could provide proof of principle for targeting of cofactor pathways in pathogenic bacteria. The link among the class Ia Fe^{3+}-Y• pathway, iron homeostasis, and oxidative stress will make selective targeting difficult. However, for pathogenic organisms with Mn and or Mn/Fe clusters (Figure 1c), interference with cluster assembly may well provide a new therapeutic target.

### SUMMARY POINTS

1. The quaternary structures of the class Ia RNR α subunit are nucleotide dependent and distinct. The α structures detected to date are α, α2, noncanonical α2 (118), α4, α6, and fibrils (27, 75).
2. dATP-inhibited RNR structures include α6 (human), α4β4 (E. coli), α4 (Pseudomonas aeruginosa) (119), and a double-helical fibril of canonical and noncanonical α2s (class Ib; B. subtilis) (75).
3. dATP-inhibited states appear to interfere with the RT pathway and –S• formation by preventing β2 from forming a productive α2β2 complex.
4. CIFDP (CIFTP) binds to human RNR and forms conformationally stable α6 state(s), even after dissociation.
5. N1NDP and F2CDP are mechanism-based inhibitors of class Ia RNRs with one inhibitor/α2 in the α2β2 complex, half-site reactivity.
6. Incorporation of unnatural amino acids (F\textsubscript{Y}122\textsuperscript{r} or NO\textsubscript{Y}122\textsuperscript{r} to replace Y\textsubscript{122} in \( \beta \) or NH\textsubscript{Y} to replace three residues: Y\textsubscript{346} in \( \beta \), Y\textsubscript{711} in \( \alpha \), or Y\textsubscript{730} in \( \alpha \)) and incubation with the second subunit, substrate, and effector trap radicals within the pathway and increases \( \alpha/\beta \) subunit affinity.

7. The reaction of F\textsubscript{Y}122\textsuperscript{r}/E52Q-\beta\text{2} with \( \alpha \text{2} \), substrate, and effector results in an asymmetric, active, and kinetically trapped \( \alpha\beta\text{2} \) complex, whose structure has been determined by cryo-EM (11). The RT pathway is revealed for the first time as is the C-terminal tail of one \( \beta \) 2 bound to \( \alpha \) 2.

**FUTURE ISSUES**

1. Trapping of additional \( \alpha\beta\text{2} \) complexes of RNRs using mechanism-based inhibitors and unnatural amino acid mutants to trap radicals in a forward or reverse radical transfer pathway may provide distinct and higher-resolution structures.

2. Cryo-EM analyses have shown that the active form of a mutant \( \text{E. coli} \) class Ia RNR is an asymmetric and dynamic \( \alpha\beta\text{2} \). The relationship of this structure to the wild-type enzyme and the structure of the human active complex remain to be established.

3. Identification of small molecules that can trap human and bacterial RNRs in distinct inhibited quaternary structures represents a new way to target RNRs.

4. The discovery of biosynthetic pathways for dimetallo-Y\textsuperscript{r} cluster assembly in class Ia and Ib RNRs suggests that targeting the metal center formation, such as disruption of NrdI/NrdF interaction in the assembly of the class Ib Mn\textsuperscript{3+}\textsubscript{2}-Y\textsuperscript{r} cofactor, might be possible.

5. The omics revolution (proteomics, phosphomics, transcriptomics) and a refined understanding of nucleotide metabolism are providing new insight into RNR regulation. This knowledge will lead to combination chemotherapies using RNR inhibitors in conjunction with inhibitors of downstream signaling pathways.

**DISCLOSURE STATEMENT**

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

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Errata

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