Metal and ligand binding to the HIV-RNase H active site are remotely monitored by Ile556

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ABSTRACT

HIV-1 reverse transcriptase (RT) contains a C-terminal ribonuclease H (RH) domain on its p66 subunit that can be expressed as a stable, although inactive protein. Recent studies of several RH enzymes demonstrate that substrate binding plays a major role in the creation of the active site. In the absence of substrate, the C-terminal helix E of the RT RNase H domain is dynamic, characterized by severe exchange broadening of its backbone amide resonances, so that the solution characterization of this region of the protein has been limited. Nuclear magnetic resonance studies of 13C-labeled RH as a function of experimental conditions reveal that the δ1 methyl resonance of Ile556, located in a short, random coil segment following helix E, experiences a large 13C shift corresponding to a conformational change of Ile556 that results from packing of helix E against the central β-sheet. This shift provides a useful basis for monitoring the effects of various ligands on active site formation. Additionally, we report that the RNase H complexes formed with one or both divalent ions can be individually observed and characterized using diamagnetic Zn2+ as a substitute for Mg2+. Ordering of helix E results specifically from the interaction with the lower affinity binding to the A divalent ion site.

INTRODUCTION

HIV-1 reverse transcriptase (RT) converts single-stranded viral RNA into double-stranded DNA for subsequent incorporation into the host cell genome. The RT p66 subunit contains a C-terminal ribonuclease H (RH) domain that is responsible for degrading the RNA template of the hybrid RNA•DNA form of the viral genome (1). Concepts of the catalytic mechanism and active site structure of RNase H enzymes have evolved significantly in recent years as a result of the availability of structural data for RNase H complexes with the RNA•DNA substrate (2–4). These results, combined with theoretical analyses (5,6), provide compelling support for a catalytic model involving two divalent cations that function to activate the nucleophile and to stabilize the transition state (4). As a result of the emergence of drug-resistant mutated forms of RT, the targeting of additional sites such as the RNase H domain of RT has attracted considerable attention (7–10).

Kinetic, mutational and structural studies have established the functional importance of residues located in the C-terminus of the RNase H domain (2,11–13). Residue Asp549 on C-terminal helix E (residues 543–554) is one of the four acidic residues of the DEDD motif that bind the two catalytically important divalent metal ions (2). Deletion of the 16 C-terminal residues (545–560) has been reported to eliminate RH activity (11). There is also evidence for a functional role of the C-terminal residues that follow helix E; deletion of the eight C-terminal residues on the p66 subunit of RT (553–560) does not eliminate RH activity, but leads to a loss of strand transfer activity (11). Mutations of Lys558 are statistically associated with a number of thymidine analog resistance mutations (TAMs), suggesting that these mutations reinforce nucleoside drug resistance (13). In addition to its importance as a drug target, the RNase H domain of RT has been studied as a model for protein folding (14,15).

Despite the functional importance of divalent ions and the C-terminal residues of the RNase H domain, nuclear magnetic resonance (NMR) studies indicate that, even in the presence of physiological levels of Mg2+ ions, these residues are dynamic and not well ordered (15–18). For the isolated RH domain the backbone amide resonances of the C-terminal helix E exhibit significant exchange broadening so that the region between residues 545 and 554 is difficult to observe (15,16,18). In contrast, amide resonances from 555 to 560 are fairly intense and their shifts are indicative of a random coil structure. The exchange broadening of the helix E resonances suggests that there is some pre-assembly of the active site helix...
since, unlike the resonances of the six C-terminal residues, they do not exhibit random coil characteristics. Similarly, crystallographic studies of apo RT structures sometimes do not reveal helix E and rarely reveal any of the post-helical residues, also consistent with a dynamic structure. Thus, the structure of the RNase H active site in the presence of its substrate differs substantially from that of the apo enzyme, which is incompletely formed and considerably more dynamic. This behavior contrasts with many enzymes that utilize a well-formed active site to achieve high substrate selectivity and catalytic efficiency. A complete understanding of the catalytic behavior and, from a practical standpoint, the development of RNase H inhibitors requires an understanding of the transformation of the RNase H active site from the dynamic, partially disordered, incompletely coordinated state of the apo enzyme to the ordered state characterizing the substrate complex. In view of the limited insight available from analysis of the backbone resonances, we have examined the behavior of resonances of the amino acid sidechains. There has been substantial recent progress in utilizing sidechain resonance shifts for conformational analysis (19–22). Remarkably, we demonstrate that the δ1 methyl resonance arising from Ile556, located in the post-helical random coil segment of the sequence, provides significant insights into the behavior of helix E and the assembly of the active site. Furthermore, this resonance can serve as a useful indicator for investigating the dependence of active site formation on divalent metal ions and other experimental conditions, and for the identification of other ligands that influence RNase H active site stability.

MATERIALS AND METHODS

The constructs used were identical to those used in previous studies and the labeled RNase H domain was produced following a protocol modified from that described previously (18,23). Briefly, the collected cell pellets were suspended in 50 mM Tris–HCl, 5% Glycerol pH 8.0 and lysed by sonication. The lysate was centrifuged at 30 000g for 30 min. The lysate supernatant was loaded onto a HiTrap SP HP column in tandem with a HiTrap QP HP column, and washed with 50 mM Tris–HCl pH 8.0 until the OD280 value of flow through approached zero. The QP HP column was gradient eluted with 1 M NaCl in 50 mM Tris–HCl, 5% Glycerol pH 8.0. The RNase H-containing fractions were pooled based on SDS–PAGE results. The pooled RNase H was concentrated to a small volume and loaded on the pre-equilibrated HiLoad 26/60 Superdex-200 column with 50 mM Tris–HCl, 200 mM NaCl 1 mM EDTA pH 8.0 buffer. All of the above procedures were carried out at 4°C. The RNase H domain mutants, H539S and R557S, were produced using the Quickchange mutagenesis kit (Stratagene, Inc., La Jolla, CA, USA). Expression and purification of the mutants was identical to procedures used for the wild-type enzyme. Subunit-specific labeling of HIV-1 RT was implemented as described previously (23) except that in the present case, we utilized a [4,13C,3,3,-2H2]-2-oxobutyrate labeling precursor in a deuterated growth medium (24).

The RNase H sequence used in this and our previous studies (18,25) corresponds to the HXB2 sequence contained in the HIV sequence compendium (26) with an additional N-terminal MNEL leader sequence that precedes Tyr427. In the present study, we have followed the standard numbering of the RNase H domain in the p66 subunit, while in our earlier BMRB entries, the residues were numbered from the N-terminus of the isolated construct. The residue identifications in the BMRB entries can be converted into the standard numbering by adding 422. The exact sequence is included as Supplementary Figure S1. Assignments of the isoleucine δ-13CH3 resonances were based on comparison of the 1H–13C HMQC spectrum of [6-13CH2-Ile]RNase H obtained in 80 mM Mg2+ with the previous entries determined under similar conditions (BMRB 5347), subject to two changes: first, the specific isoleucine Cδ-labeled sample indicates that the previous assignments of the Ile521 Cδ and Cγ2 resonances need to be reversed. Second, the unusual shift and exchange broadening of Ile556 Cδ resonance apparently resulted in an incorrect assignment of this resonance in BMRB 5347. The correct assignment can be made by exclusion, since each of the other nine Ile Cδ resonances is assigned using the above procedure.

The Ile434 and Ile556 δ–methyl resonances in the p66 subunit of HIV-1 RT were assigned based on their positions in the isolated RNase H domain and their relative isolation, which greatly limits the possibility of misassignment. Analogous domain-based assignment strategies have been used in similar studies of very large proteins (27,28). A more extensive study of isoleucine-labeled RT is currently in preparation.

Synthesis of the RNase H inhibitor 2-hydroxyisoquinoline-1,3(2H,4H)-dione is described by Billamboz et al. (29), and was obtained as a custom synthesis product from MRIGlobal (Kansas City, MO). 1H–13C HMQC spectra were obtained at 25°C using Varian’s gChmqc sequence on a UNITY INOVA 500 MHz spectrometer with 128 × 1024 complex points and acquisition time of 128 ms in both t1 and t2. Sixteen scans were acquired per increment, with a 1.0 s delay between scans. The 1H–15N HSQC spectra of the arginine NHε region were acquired at 25°C using the Varian gNHsqc sequence. The 15N offset was set at 90 ppm. All NMR data were processed by NMRpipe (30) and analyzed with NMRViewJ program (31). Titration studies were performed on 15 μM RNase H domain in 20 mM Tris–d11. Structures were analyzed and visualized with PyMol (www.pymol.org).

The pH and Mg-induced Ile556 titration data was fit using a single KD according to the relations:

\[
\delta^{13}C = c_1 + (c_2 - c_1) \frac{10^{pK-pH}}{1+10^{pK-pH}} (1)
\]

\[
\delta^{13}C = c_1 + (c_2 - c_1) \times \frac{([Mg]+[RH]+K_D) - \sqrt{([Mg]+[RH]+K_D)^2 - 4[Mg][RH]}}{2[RH]} (2)
\]
where \( c_1 \) and \( c_2 \) are the limiting Ile shifts for the apo enzyme under high or low pH conditions, or low and high Mg\(^{2+}\), respectively, and the p\( K \) and \( K_D \) values are also allowed to vary. For the Mg\(^{2+}\) titration, the data were insufficient to allow fitting to a more complex binding model.

The \( P_{\text{HelixE}} \) values calculated from the Ile556 C\( \delta \) methyl shift using the relation in the text, were subsequently fitted to Equation (1); however in this case, the \( c_1 \) and \( c_2 \) parameters correspond to limiting helix probabilities. Similarly, the \( P_{\text{HelixE}} \) values determined as a function of Mg\(^{2+}\) from the measured Ile556 C\( \delta \) methyl shift values and the relation in the text were subsequently fit to the relation:

\[
P_{\text{HelixE}} = c_1 + (c_2 - c_1) \frac{[\text{Mg}^{2+}]}{[\text{Mg}^{2+}] + K_D} \tag{3}
\]

in order to evaluate the limiting behavior of \( P_{\text{HelixE}} \) as a function of Mg\(^{2+}\).

RESULTS

The limited information available from analysis of the backbone resonances of the RT RNase H domain, particularly in the region of the C-terminus, led us to investigate the behavior of the sidechain resonances. Surprisingly, the spectrum of the [\(^{13}\)C\( \delta \)-Ile] RNase H domain revealed a significant sensitivity of the Ile556 methyl resonance to pH (Figure 1 and Supplementary Figure S2). A fit of the titration data for Ile556 yields total titration shifts of \( \Delta^{13}\text{C} = -2.34 \) ppm; \( \Delta^1\text{H} = -0.146 \) ppm, both corresponding to a p\( K \) = 5.77, with the resonances moving upfield as the pH is reduced. There are no titratable residues in the sequence surrounding Ile556: LVSAGI556RKVL560 consistent with the observed p\( K \) value, and the influence of residues more distant in sequence but in spatial propinquity is inconsistent with the disordered conformation of this segment.

There is increasing evidence that the \(^{13}\)C shift values for many amino acid sidechain resonances appear to be dominated by conformationally-dependent ‘\( \gamma \)-effects’ first characterized in studies of aliphatic hydrocarbons by Grant and coworkers (32,33). According to this analysis, the upfield shift observed for Ile556 \( \delta_1 \) methyl is typical of Ile residues with \( \chi_2 \) in the g- conformation (19). Using the \( \chi_2 \) distribution changes from \( \sim 45\% \) g- conformer at neutral pH, to \( \sim 84\% \) g- conformer at low pH, so that its pH dependence may be qualitatively described by the relation:

\[
\text{Ile556}_{\chi_2 \sim g} \xrightleftharpoons{\text{pH}^\uparrow}{\text{pH}^\downarrow} \text{Ile556}_{\chi_2 \sim g}
\]

where \( rc \) represents a random coil mixture. Since the g- conformation represents a less energetically favorable rotamer than the predominant \( t \) conformation, the high g- probability that characterizes the low pH behavior of Ile556 is inconsistent with a random coil state. Thus, this behavior indicates that at low pH, Ile556 experiences significant inter-residue stabilizing interactions that compensate for the higher intrinsic energy of the g- rotamer.

Crystallographic studies of RT provide limited information on the behavior of the RNase H C-terminus; however, recent studies of RNase H–inhibitor complexes show the positions of both bound Mg\(^{2+}\) ions as well as most of the C-terminal residues (34–37). Interestingly, although none of the structures of apo RT show an Ile556 residue with a \( \chi_2 = g- \) conformation, all of the RNase H–inhibitor complexes show this conformation to be present (Figure 2). In each of these structures, the sidechain of Ile556 packs against the central \( \beta \)-sheet and continues the amphiphilic motif of Val548 and Val552. These structures reveal the role of the bound Mg\(^{2+}\) ions, and particularly Mg\(_A\), in stabilizing the orientation of helix E by bridging Asp443, located on the central \( \beta \)-sheet, and Asp549, located on helix E (Figure 2a). It is reasonable to hypothesize that the observed pH dependence of the Ile556 shift results from formation of a hydrogen bonding interaction between the same two aspartyl residues, such that the repulsive interaction at higher pH is replaced by a positive interaction as the hydrogen bond forms at lower pH. The p\( K \) value of 5.77 obtained from the Ile556 shifts is well above that expected for an isolated aspartyl sidechain (38), but is consistent with a shared hydrogen bond.

We also considered the alternative hypothesis that the pH-dependent shifts of Ile556 and Ile542 might result from a conformational transition induced by titration of His539. This hypothesis is based on the proximity of His539 to Glu546 and Asp549 in some of the structures of RNaseH–inhibitor complexes, as well as the similarity of the p\( K \) values obtained above to typical histidine p\( K \) values.

In order to evaluate the role of such interactions, we analyzed the pH behavior of the isoleucine residues in an RNase H (H539S) mutant. The titration behavior of Ile556 in the mutant is essentially unaffected by this mutation, while there is a somewhat larger perturbation of the Ile542 titration, consistent with its proximity to the mutated residue (Supplementary Figure S3). Thus, titration of His539 has no effect on the behavior of Ile556, consistent with the above interpretation that the pH-dependent effect on Helix E results from an effect on the electrostatic repulsion of the active site aspartyl residues.

It is likely that both the integrity of the helix, as well as its position is also destabilized by loss of the interactions with the \( \beta \)-sheet. However, the exchange broadening of the amide resonances as well as the Ile556 \(^{13}\)C\( \delta \) shift behavior discussed above indicate that the structure does not unwind and produce a random coil, but equilibrates between the packed conformation observed in the inhibitor complex and a partially disordered conformation. In summary, comparison of the titration data with the structures of RNase H–inhibitor complexes strongly suggests that lowering the sample pH is exerting a conformational effect that is in some respects analogous to the conformation observed for the Mg\(^{2+}\)–inhibitor complexes.
Effects of divalent metal ions

Based on the above analysis, we also titrated the RNase H sample with MgCl2. The binding of Mg2+ influences the Ile556 Cδ shift similarly to a pH reduction (Figure 1b). Interestingly, the Ile556 and (smaller) Ile542 shift responses to both H+ and Mg2+ are qualitatively similar, while the responses of the remaining methyl groups are very different (Figure 1a and b). This result is consistent with the conclusion that Ile542 and Ile556 are sensitive to the changing position/stability of helix E, while for the remaining isoleucine resonances, H+ and Mg2+ exert different conformational effects. Analysis of the Mg2+ titration study yielded an apparent $K_{\text{Mg}}$ of 6.3 mM (Supplementary Figure S4), falling between the values of 3.2 and $\sim$35 mM previously obtained for Mg2+ binding to the apo RNase H domain (18). This intermediate apparent $K_{\text{Mg}}$ value probably results from partial occupancy of both the A and B divalent ion sites, since A-site binding would be expected to have a much more significant stabilizing effect on helix E.

Crystal structures of the isolated HIV-1 RT RNase H domain (39) and the Escherichia coli RNase HI (40) in the presence of Mn2+ indicate that both divalent ion positions are occupied. In order to achieve an analogous occupation of both divalent ion sites without the associated paramagnetic effects on resonance linewidth, we utilized another transition metal ion, Zn2+. The ionic radii of Zn2+ and Mg2+ are nearly identical, and it has been reported that optimal RT RNase H activity is achieved at $\sim$25 μM Zn2+, much lower than the concentration of Mg2+ required for optimal activity (although the maximum rate obtained with Zn2+ is substantially below that determined in the presence of Mg2+) (41). Addition of Zn2+ at a 1:1 molar ratio with RNase H results in small shift changes for several isoleucine methyl resonances, particularly Ile482 and Ile505 but produces a minimal perturbation of the Ile556 resonance (Figure 3). This result is consistent with the conclusion that the initially bound Zn2+ ion exhibits sufficient differential affinity for the A and B sites such that only the higher affinity B site is significantly occupied, and selective occupation of site B by a divalent ion is insufficient to significantly influence the stability of helix E (Figure 2a). Further addition of Zn2+ to produce a 2:1 complex results in a large, upfield shift of Ile556, with $\delta^{13}\text{C} = 9.8$ ppm (Figure 3b), lower than the values observed at low pH or at 64 mM Mg2+ ($\sim$10.3 ppm). These results clearly indicate that at a 2:1 ratio, both divalent ion sites are largely occupied, with the occupation of site A resulting in significant helix E stabilization. The more readily resolved Ile505 resonance also shows clearly the effects of the two separate Zn2+ complexes that are formed. Formation of the 1:1 complex produces a shift to a slightly lower $\delta^{13}\text{C} \sim 11.9$ ppm value. Subsequent formation of the 2:1 complex results in a distinct resonance with a larger $\delta^{13}\text{C} \sim 12.4$ ppm (Figure 3b). Further increases in the Zn2+ concentration resulted in only very small additional shifts of the Ile556 resonance, although the resonance becomes more intense, consistent with a reduction in the exchange broadening (Supplementary Figure S5).

The $^1H$–$^{13}C$ HMQC spectrum of the $^{[13}\text{C}^\text{d}_3\text{H}_3$-Ile]RNase H–Mn2+ complex was also obtained; however, the paramagnetic broadening produced by the Mn2+ ion prevents...
observation of the isoleucine methyl resonances arising from the closer Ile482, Ile542 and Ile556 residues (Supplementary Figure S6).

**Effect of a Mg\(^{2+}\)-inhibitor complex and evaluation of a helix probability factor**

As is apparent from Figure 1b, even at 64 mM Mg\(^{2+}\), the Ile556 resonance exhibits substantial exchange broadening, indicating that even very high Mg\(^{2+}\) concentrations are insufficient to strongly stabilize the completely folded state of the RNase H domain. Similarly, the backbone amide resonances of helix E remain difficult to observe even in the presence of high Mg\(^{2+}\) (18). In order to evaluate the behavior of a more completely stabilized ternary complex, we obtained a representative dual Mg\(^{2+}\)-binding ligand, 2-hydroxyisoquinoline-1,3(2H,4H)-dione, the synthesis of which has been described by Billamboz et al. (29).

The 2-hydroxyisoquinolone and structurally related ligands interact with both Mg\(_A\) and Mg\(_B\), forming a stable RNaseH\(\bullet\)Mg\(^{2+}\)isoquinolone complex. Addition of 4 mM Mg\(^{2+}\)+0.5 mM isoquinolone ligand resulted in further spectral changes, such that the Ile556 \(^{13}\)C\(_\beta\) is shifted to a more extreme position (\(\delta^{13}\)C = 8.6 ppm), and the exchange broadening is dramatically reduced (Figure 4a). Thus, we assign this Ile556 shift to Ile556 locked in the \(\chi_2 = \gamma\) conformation characteristic of the fully folded state (21).

Since Ile556 is not located directly in the active site, the shift behavior of the Ile556 C\(_{\delta}\) resonance can be used to provide an estimate for the formation of helix E as a function of different experimental conditions that is largely independent of the details of active site structure and ligands. Defining the probability of a well-formed helix to be 1 in the presence of both Mg\(^{2+}\) and the isoquinolone inhibitor, and estimating that in the absence of a helix, the shift parameters for this Ile556 should be similar to those of a random coil (42), we obtain:

\[
p_{\text{HelixE}} = \frac{\delta_{\text{obs}}^{13} \text{C}_{\delta} - \delta_{\text{Mg}^{2+}-\text{Inh}}^{13} \text{C}_{\delta}}{\delta_{\text{obs}}^{13} \text{C}_{\delta} - \delta_{\text{Mg}^{2+}-\text{Inh}}^{13} \text{C}_{\delta}} = \frac{12.9 - 8.6}{4.3} = 1.29
\]

Using this expression, the helix E orientational probability varies from \(\sim 14\%\) at pH 7.1 up to \(\sim 63\%\) at pH 4.5 (Figure 5). These values are generally consistent with the observed exchange broadening that makes the backbone amide resonances difficult to observe. Even a pH of 4.5 or a Mg\(^{2+}\) concentration of 64 mM is insufficient to stabilize the helix by \(\sim 65\%\). We note that the calculation of Equation (4) provides an estimate of the helix E orientational probability in RNase H based on the behavior of the Ile556 resonance, and is distinct from the \(\gamma\) estimate that can be made using the relation given by Hansen et al. (21), which is based on an analysis of the relationship between \(\delta(C^\delta)\) and \(^{3}J(C^\delta,C^\beta)\) values for the residues in a set of six proteins.

**Behavior of the RNase H domain in full RT**

In order to evaluate whether a similar transition can be observed for the full RT molecule, we prepared \([^{13}\text{C}_0\text{H}_3\text{-Ile}]_{66}\text{RT}\), containing isoleucine C\(_\delta\) methyl labels in the p66 subunit. In the spectrum of RT obtained in the absence of Mg\(^{2+}\), the resonance for Ile556 was not readily apparent, possibly due to overlap with other resonances or to more severe exchange broadening. Surprisingly, addition of 4 mM Mg\(^{2+}\) was found to produce a much more extreme shift for Ile556 in RT than in the isolated RNase H domain (9.7 versus 11.4 ppm) (Figure 4b). In the full RT molecule, the N-terminus of helix E including Ile542

![Figure 2. Crystal structure of the active site of the HIV RT RNase H domain. (a) Isolated RT RNase H domain–inhibitor complex [pdb code: 3K2P (34)]. The inhibitor, which binds to both Mg ions, is not shown. The four acidic residues that complex the two Mg ions are shown, along with residues Val548, Val552 and Ile556 that mediate the interaction of helix E with the central \(\beta\)-sheet. Residue Ile556, indicated in orange, adopts a \(\gamma\) conformation for \(\chi_2\). (b) Active site region of RT RNase H domain with bound RNA\(\bullet\)DNA hybrid modeled into the active site. The model is based on an overlay with the structure of the human RNase HI RNAs\(\bullet\)DNA complex [pdb code 2QKK(3)].](image-url)
interacts with the p51 subunit. The interface includes a hydrophobic-binding pocket on p51 that interacts with Ile542 and, in some structures, additional hydrogen bonding interactions between Arg284_{p51} and residues Gln547 and/or Glu546 on the RNase H domain (Figure 6). We thus conclude that the interaction of the RNase H domain on the p66 subunit with the thumb subdomain on the p51 subunit, helps to orient helix E,

![Figure 3](image1.png)

**Figure 3.** Effect of Zn^{2+} on RT RNase H Ile methyl resonances. (a) Overlay of the {^{1}H}-{^{13}C} HMQC spectra of the 15\textmu M [^{13}C\text{H}_{3}-\text{Ile}]RNase H domain (black), 20\textmu M RNase H plus 20\textmu M Zn^{2+} (red). (b) Overlayed spectra of 20\textmu M [^{13}C\text{H}_{3}-\text{Ile}]RNase H with 20\textmu M Zn^{2+} (red) and with 40\textmu M Zn^{2+} (blue). c) Overlayed spectra of 15\textmu M [^{13}C\text{H}_{3}-\text{Ile}]RNase H (black), 20\textmu M [^{13}C\text{H}_{3}-\text{Ile}]RNase H with 40\textmu M Zn^{2+} (blue) and 20\textmu M [^{13}C\text{H}_{3}-\text{Ile}]RNase H with 16\textmu M Zn^{2+}-ATP (magenta).

![Figure 4](image2.png)

**Figure 4.** Effect of Mg^{2+} and 2-Hydroxyisoquinoline-1,3(2H,4H)-dione ligands on RNase H Ile resonances. (a) Overlay of the {^{1}H}-{^{13}C} HMQC spectra of the isolated [^{13}C\text{H}_{3}-\text{Ile}]RNase H domain (black), plus 4 mM Mg^{2+} (red), after subsequent addition of 0.5 mM isoquinolone (blue). (b) Overlayed {^{1}H}-{^{13}C} HMQC spectra of 20\textmu M [^{13}C\text{H}_{3}-\text{Ile}]RNase H with 20\textmu M Zn^{2+} (red) and with 40\textmu M 2-Hydroxyisoquinoline-1,3(2H,4H)-dione (blue). The inset in Figure 3b shows the Ile556 resonances plotted with a lower threshold. In addition to the dramatic shift of Ile556, a few other isoleucine resonances experience small shift perturbations.
leading to a more completely preformed MgA-binding site with higher Mg2+ affinity, so that addition of a fixed concentration of Mg2+ leads to a greater shift of 9.2 ppm for Ile556. Based on the observed Ile556 13C shift of 9.2 ppm, we obtain $P_{\text{HelixE}} = 86\%$ for the RH domain of RT, compared with 35% for the isolated RNase H domain, both evaluated in the presence of 4 mM MgCl2. Addition of the isoquinolone inhibitor leads to a more extreme shift of the Ile556 methyl resonance and eliminates the exchange broadening, consistent with a well-defined helix orientation. In contrast with the result obtained in the presence of Mg2+, the Ile556 resonance shifts observed for the ternary complexes of both RT and the isolated RNase H domain with Mg2+-isoquinolone are similar. In general, the relative rarity of the $\chi^2 = g-$ shift conformation makes it easy to identify the corresponding resonances even in enzymes such as RT that contain many isoleucine residues.

An analogous difference of pH sensitivity was also observed between the isolated RH domain and intact RT. At pH 6.1, Ile556 $\delta^{13}C = 11.7$ ppm for the isolated RH domain, and 9.6 ppm in RT (data not shown). It thus appears likely that the additional structural features present in RT that help to preform the MgA-binding site also lead to elevation of the pK value that characterizes the Ile556 shift. This result is consistent with the presence of a bridging interaction in which Asp443 and Asp549 share a proton.

**Structural and catalytic role of Arg557**

The data presented above support the conclusion that the conformations of the C-terminal residues observed in the recently determined inhibitor-complex structures, which are not apparent in structures of the apo enzyme, can be significantly populated at low pH or in the presence of divalent ions. In addition to the role of Ile556 in stabilizing the orientation of helix E, this structural analysis also supports a functional role for Arg557, in orienting the active site Asp549 residue, stabilizing the
helix E structure, and presumably interacting with the RNA strand of the substrate.

In order to better characterize the behavior of Arg557, we obtained $^1$H-$^{15}$N HSQC spectra of the region containing the arginine NH$_e$ resonances at pH 5.5, at which the slower exchange of H$_e$ facilitates the observations (Figure 7). The well-resolved resonance is assigned to Arg463 on the basis of structures showing a side-bonded guanidino group of this residue with the carbonylate of Glu438 [e.g. pdb 3K2P, (34)]. This salt bridge plays an important role in limiting the accessibility of the Phe440–Tyr441 protease cleavage site in the RNase H domain (43). Two other Arg NH$_e$ resonances are observed, with an intensity ratio of 2:1. The Arg557 resonance was assigned based on the spectrum of an RNase H (R557S) mutant as indicated in Figure 7, and the remaining resonance, with approximately double the intensity of each resolved peak, corresponds to Arg448 and Arg461. The resolution of the Arg557 NH$_e$ resonance, despite the fact that H$_e$ is not involved in a hydrogen bond, is consistent with the positional constraint of the Arg557 guanidino group. In contrast, the NH$_e$ resonances for Arg448 and Arg461 are degenerate. A small upfield shift for the H$_e$ resonance of Arg463 is also observed for Arg463 on the basis of structures showing a side-bonded guanidino group of this residue with the carbonylate of Glu438 [e.g. pdb 3K2P, (34)]. This salt bridge plays an important role in limiting the accessibility of the Phe440–Tyr441 protease cleavage site in the RNase H domain (43). Two other Arg NH$_e$ resonances are observed, with an intensity ratio of 2:1. The Arg557 resonance was assigned based on the spectrum of an RNase H (R557S) mutant as indicated in Figure 7, and the remaining resonance, with approximately double the intensity of each resolved peak, corresponds to Arg448 and Arg461. The resolution of the Arg557 NH$_e$ resonance, despite the fact that H$_e$ is not involved in a hydrogen bond, is consistent with the positional constraint of the Arg557 guanidino group. In contrast, the NH$_e$ resonances for Arg448 and Arg461 are degenerate. A small upfield shift for the H$_e$ resonance of Arg463 is also observed for the R557S mutant. Presumably, this results from a longer range influence of the helix E structure on the adjacent $\beta$-sheet, which includes Arg463.

The limiting shift of Ile556 observed in the presence of both Mg$^{2+}$ and the isoquinolone inhibitor is also somewhat smaller for the R557S mutant ($\delta^{13}$C = 9.8 ppm; Supplementary Figures S7). We interpret this difference as a consequence of the backbone restraint imposed by positioning of the Arg557 sidechain. In the absence of this conformational restraint, the backbone of the adjacent Ile556 is also less constrained and this will also influence the sidechain conformation as it packs against the $\beta$-sheet and the Lys454 sidechain.

**Identification of a new RNase H ternary complex**

Coupling of the $^{13}$C shift of Ile556 C$\beta$ with the position of helix E and assembly of the active site makes this resonance well suited as a sensitive probe for the discovery of new RNase H active site complexes. The location of the probe $>$10 Å from the active site itself suggests that the shift should be sensitive to how well positioned the helix is, without being strongly dependent on the detailed nature of the ligand complex. After surveying a series of divalent cations and available ligands, we found that Zn$^{2+}$-ATP was also capable of producing a shift of the Ile556 C$\beta$ resonance that was similar to that produced by the Mg$^{2+}$-isoquinolone inhibitor complex (Figures 3c and 4b).

We also investigated the effect of this ternary complex on the behavior of the $^1$H-$^{15}$N HSQC spectrum containing the arginine NH$_e$ resonances. Addition of 200 µM Zn$^{2+}$ to the sample at pH 5.5 resulted in small decreases in intensity of several resonances. In contrast, addition of 200 µM Zn$^{2+}$ + 100 µM ATP decreased the intensity of the Arg557 resonance by ~50% (Figure 7c). The most probable interpretation of this result is that the bound Zn-ATP helps to restrain the position of the Arg557 sidechain, resulting in a decreased $T_2$ value, reduced polarization transfer and hence reduced resonance intensity, although several alternative interpretations are also possible. Regardless of the basis for the intensity perturbation, this observation is consistent with the suggestion above that the Arg557 guanidinium sidechain helps to stabilize the interaction of the negatively charged Asp549 sidechain and a negatively charged ligand—either the substrate or in this case, ATP.

**DISCUSSION**

Concepts regarding the structure and catalytic mechanism of RNase H enzymes have undergone a considerable transformation in recent years as a result of the availability of crystal structure data for several RNase H–RNA•DNA complexes (2,3). These structures demonstrate the critical contribution of the substrate to the divalent ion-binding sites. In contrast with these results, the structure of an RNase H complex with dsDNA lacks both of the bound divalent ions (44). In combination with the characterization of the dynamic state of the apo RT RNase H domain, these structures demonstrate the major role that the hybrid RNA•DNA substrate plays not just for divalent ion binding, but for the creation of the active...
site. Although the active sites of most enzymes are highly pre-organized to make optimal and highly discriminating contacts with the substrate, the results presented above for the RT RNase H domain demonstrate that the active site is quite dynamic and lacks significant structural integrity in the absence of the substrate. Even Mg\(^{2+}\) concentrations well in excess of normal physiological levels of \(~1\text{mM}\) (45,46) only are able to partially overcome the lack of pre-organization, while the addition of a specifically designed inhibitor results in a more significant stabilization of the active site structure. Under physiologic conditions, this low Mg\(^{2+}\) affinity is alternatively overcome as a result of synergistic binding of Mg\(^{2+}\) with the hybrid RNA\(\bullet\)DNA substrate. These studies thus support the conclusion that substrate-induced active site stabilization plays a major role in the determination of enzymatic substrate specificity. Recognition and activation are thus not achieved by a preformed active site, but by requiring the substrate nucleotide to complete the formation of the active site of the enzyme.

Although there is little data on the solution behavior of the *Bacillus halodurans* RNase H, the more significant lack of stabilizing elements in the structure suggests that it too relies on substrate binding for formation of the active site complex. Nowotny and Yang (4) propose that product release is likely to require dissociation of Mg\(^{2+}\), so that the poor active site stability may contribute to product release. Since the Tyr441 sidechain interacts directly with Gly544 in helix E, partial disorder of helix E in the absence of ligands may also facilitate the unfolding of one RNase H domain, which is required to allow HIV protease access to the F440–Y441 cleavage site (39).

These studies also suggest a functional role for Arg557 in the catalytic activity of RT RNase H. A structure of the complex of RT RNase H domain with an RNA\(\bullet\)DNA hybrid (Figure 2b) was generated by superposition with the structure of the human RNase HI in complex with an RNA\(\bullet\)DNA (pdb code: 2QKK) (3). The sidechain of Lys454 interrupts the progress of helix E, forcing the backbone to loop around so that Arg557 is positioned to form a salt bridge with Asp549. This leads to a positional similarity of the guanidine sidechain of Arg278 in the human enzyme with Arg557 in RT RNase H, suggesting a functional equivalence that is not apparent from the sequence alignment (3). It is likely that this residue facilitates substrate binding by reducing the electrostatic repulsion between Asp549 and the negatively charged phosphate groups on the substrate. The effect of the Zn\(^{2+}\)-ATP ligand on the Arg557 NH\(\epsilon_1\) resonance intensity (Figure 7c) supports the significance of an analogous interaction between Arg557 guanidinium group and the negatively charged substrate.

As demonstrated here, Mg\(^{2+}\) possesses a limited ability to stabilize the active site; its limiting effect on helix stability is in fact similar to the effect of low pH; both Mg\(^{2+}\) and H\(^+\) produce asymptotic limits of the helix E formation probability of \(\sim2/3\). Based on the concentration dependence of helix E stabilization, at least some of the effect of the Mg\(^{2+}\) appears to result from partial occupancy of the two divalent ion-binding sites. Mn\(^{2+}\) has been widely used as a Mg\(^{2+}\) substitute in X-ray crystallographic studies due to its chemical similarity and its anomalous scattering at the copper K\(\alpha\) wavelength. The tighter binding of Mn\(^{2+}\) to both the *E. coli* and RT RNase H enzymes is reflected in the ability to form two-ion complexes, while no complexes containing both catalytic Mg\(^{2+}\) ions have been observed in the absence of substrate or inhibitor. As shown in this study, Zn\(^{2+}\) provides an attractive diamagnetic alternative to Mn\(^{2+}\) for some NMR studies, with the tighter binding leading to more complete helix stabilization and allowing a clean discrimination of the two divalent ion-binding sites. Typical Mg\(^{2+}\) dissociation constants often exceed the conveniently accessible protein concentrations used for NMR studies, making it difficult to identify specific binding interactions. Alternatively, the Zn\(^{2+}\) dissociation constants will often be lower than the enzyme concentrations used for NMR studies, allowing better identification of specific binding interactions. Consistent with these observations, Zn\(^{2+}\) is able to support RT RNase H activity at \(\sim25\mu\text{M}\), significantly below the millimolar concentrations of Mg\(^{2+}\) that are typically required (41).

The differential effects of the metal ions on helix E stability provide additional insight into one of the more puzzling observations related to RNase H domain activity. As summarized above, the isolated RT RNase H domain has negligible activity (47–49). However, modifications that enhance substrate-binding affinity such as the addition of His-tags or the inclusion of substrate-binding residues derived from the *E. coli* enzyme are able to restore Mn\(^{2+}\) activity, but not Mg\(^{2+}\)-dependent activity. The significantly greater abilities of Zn\(^{2+}\) and Mn\(^{2+}\) compared with Mg\(^{2+}\) to bind to both divalent ion sites in the absence of substrate suggests a weaker requirement for substrate-induced positioning of the ions. Thus, consistent with the substrate-dependent specificity behavior outlined above, correctly positioned Mn\(^{2+}\) ions may be able to compensate for poorer substrate positioning, while the proper orientation of the Mg\(^{2+}\) ions is fully dependent on correct substrate positioning. The inability of bound dsDNA to support divalent metal ion binding to the *B. halodurans* RNase H (44), is also consistent with this general view that correct substrate positioning is critical to the correct binding of Mg\(^{2+}\).

**Ile556 shift as a basis for ligand screening**

Another significant implication of the present study is the utility of conformationally-induced shifts such as those observed for Ile556, as a basis for ligand screening. Isotopically labeled ILV methyl groups provide a useful basis for identifying ligand-binding sites by chemical shift mapping (50). Interestingly, the total \(^{13}\text{C}\) shift of \(\sim3.6\text{ppm}\) observed for the Ile556 methyl resonance is larger than any of the reported direct methyl shift contributions observed in a study designed to map ligand-binding sites (50). In general, it is probably straightforward to distinguish these effects since, as in the present case, the conformationally mediated \(g/t\) transitions occur for residues that are remote from the binding site and typically buried in the protein interior, while surface residues are more likely to adopt the lower energy \(\text{trans}\) conformation and to experience direct ligand-induced
shifts. For the RNase H domain, either isolated or as part of RT, the Ile 556 resonance provides a direct readout of the stabilization of helix E that is expected to be essentially independent of the details of ligand binding. The feasibility of using the Ile556 shift in this way is illustrated by our discovery of an RNase H $\text{HeZn}^{2+}$-ATP complex, the details of which are currently under investigation.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online: Supplementary Figures 1–7.

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