Interaction of anthracyclines with iron responsive element mRNAs

Joshua C. Canzoneri and Adegboyega K. Oyelere*

School of Chemistry and Biochemistry, Parker H. Petit Institute for Bioengineering and Biosciences, Georgia Institute of Technology, Atlanta, GA 30332-0400, USA

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ABSTRACT

Double-stranded sections of mRNA are often inviting sites of interaction for a wide variety of proteins and small molecules. Interactions at these sites can serve to regulate, or disrupt, the homeostasis of the encoded protein products. Such ligand target sites exist as hairpin–loop structures in the mRNAs of several of the proteins involved in iron homeostasis, including ferritin heavy and light chains, and are known as iron responsive elements (IREs). These IREs serve as the main control mechanism for iron metabolism in the cell via their interaction with the iron regulatory proteins (IRPs). Disruption of the IRE/IRP interaction could greatly affect iron metabolism. Here, we report that anthracyclines, a class of clinically useful chemotherapeutic drugs that includes doxorubicin and daunorubicin, specifically interact with the IREs of ferritin heavy and light chains. We characterized this interaction through UV melting, fluorescence quenching and drug–RNA footprinting. Results from footprinting experiments with wild-type and mutant IREs indicate that anthracyclines preferentially bind within the UG wobble pairs flanking an asymmetrically bulged C-residue, a conserved base that is essential for IRE–IRP interaction. Additionally, drug–RNA affinities (apparent $K_d$s) in the high nanomolar range were calculated from fluorescence quenching experiments, while UV melting studies revealed shifts in melting temperature ($\Delta T_m$) as large as 10 °C. This anthracycline–IRE interaction may contribute to the aberration of intracellular iron homeostasis that results from anthracycline exposure.

The unique tertiary structures attainable by RNA, and the burgeoning knowledge of the ligand specificity of those tertiary structures, have revealed RNA to be an inviting target for a variety of proteins and small molecules (1–7). The interaction of these ligands with mRNA can alter the translation of the encoded protein. The iron responsive elements (IREs) of proteins involved in iron metabolism serve as the premier regulatory site of translation due to their unique tertiary structure. IREs are ~30-nt long hairpin–loop structures in the untranslated regions (UTRs) of several iron metabolism proteins including both ferritin subunits (Figure 1), ferroportin and the erythroid isofrom of aminolevulinate synthase, eALAS (8,9). The structure and sequence of IREs is highly conserved throughout all of the animal kingdom including early metazoan (10). The homeostasis of IRE containing proteins is regulated by the iron-dependent availability of the IRP proteins (8). Ligand interaction in the UTRs of mRNA can serve to stabilize or destabilize the mRNA, and thus increase or decrease translation. In general, binding of a protein or small molecule in the 5′-UTR inhibits translation by deterring the association of translation factors and preventing the assembly of the ribosome. Conversely, the binding of a ligand in the 3′-UTR upregulates translation by protecting against nuclease degradation (8). In the case of IRE–IRP interaction, at low intracellular iron concentrations, IRP1 is converted into its active binding state through the dissociation of its 4Fe-4S cluster. This active state then binds to the 5′ IRE of ferritin heavy and light chain mRNAs, serving to downregulate translation and reduce the iron-storage capacity of ferritin (8). Disruption of this IRE/IRP interaction could greatly affect intracellular iron handling.

Anthracyclines are powerful antineoplastic agents that are widely used for the treatment of leukemia, lymphoma, solid tumors and Hodgkin’s disease (11). However, the clinical use of anthracyclines is limited by the occurrence of both chronic and acute cardiotoxicity that can often lead to cardiomyopathy, myocardial infarction and congestive heart failure (11–13). The mitigation of these often fatal side effects is of high interest and many attempts have been made to characterize the mechanism of anthracycline cardiotoxicity. However, there is much disagreement on this point in the literature. Theories include the suppression of transcription factors, production of reactive...
oxygen species, free radical-induced oxidative stress, decreased levels of intercardiac protective proteins and impaired intracellular iron handling and homeostasis (14–22). Elegant attempts have been made to tie all of the popular pathways listed above into a single complex model for anthracycline cardiotoxicity (23). However, the exact method by which anthracyclines lead to cardiomyopathy must still be elucidated in order to rationally design new therapeutic options for the successful treatment of cancer with anthracyclines.

The antitumor effects of anthracyclines are attributed to the ability of the molecules to inhibit topoisomerase by intercalating DNA, and the specifics of this interaction are very well-characterized (24–32). However, the ability of these drugs to intercalate double-stranded RNA has largely been neglected in the literature. It has previously been observed that anthracyclines cause an imbalance in the regulation of IRP and its binding to IREs, but it has yet to be investigated whether this perturbation is a direct result of anthracyclines interacting with the IREs (17,18). Toward unraveling the mechanism of anthracycline-induced disruption of IRE binding to IRP we investigated the possibility of a direct interaction between anthracyclines and IREs harbored at the 5'-UTR of ferritin heavy and light chain mRNAs. A direct interaction of a small molecule, such as an anthracycline, with the IRE could change the tertiary structure of the RNA drastically and alter the effectiveness of the IRE/IRP interaction. In this study, we characterized the isolated IREs of ferritin heavy and light subunit mRNAs in the absence and presence of doxorubicin (DOX) and daunorubicin (DAU), two widely used anthracyclines, via biophysical techniques including UV melting, fluorescence quenching and drug–RNA footprinting. Results from these experiments suggest that anthracyclines do indeed interact with the IRE hairpin–loop structures in a strong and specific manner. This interaction may contribute to the intracellular aberration of iron regulatory protein homeostasis upon anthracycline exposure.

**EXPERIMENTAL PROCEDURES**

**Drugs and reagents**

Doxorubicin hydrochloride and daunorubicin hydrochloride were procured from Oakwood Products Incorporated. All reagents were of molecular biology grade (Sigma, St. Louis, MO). Tubes and tips were sterile and autoclaved prior to use. Water was ultra pure, autoclaved for 1 h, and filtered through a 0.2μm filter prior to use. All buffers used were also filtered prior to use.

**mRNA sequence selection and preparation**

The mRNA sequences of the human proteins of interest were retrieved from the Entrez Nucleotide Database (www.pubmed.org). The UTRs of these mRNA sequences were identified, and the folding of these sequences was modeled in the RNAstructure Version 4.3 program (rna.urmc.rochester.edu/eastructure.html). The resulting secondary structures were surveyed to identify hairpin loop structures. The isolated sequence of the loop and stem region of these structures, on average 30nt in length, was then folded via the RNAstructure program once again to ensure that the secondary structure was maintained in the absence of the rest of the sequence (Figure 1). The mRNA sequences of these structures were purchased from Dharmacon Research Inc., Lafayette, CO, deprotected according to the manufacturer’s protocol, and their purity was verified via denaturing urea polyacrylamide gel electrophoresis. RNA concentrations were determined by UV-vis spectroscopy in 1 cm path length quartz cuvettes using the molar extinction coefficients provided by Dharmacon.

**UV-vis melting and absorption experiments**

UV melting experiments were performed in the absence of drug and in the presence of DOX or DAU at final drug concentrations of 5 μM and 20 μM. RNA and drug complexes were formed by the addition of an aliquot of stock drug solution in DMSO to a volume of RNA in BPE buffer (6 mM Na2HPO4, 2 mM Na2H2PO4, 1 mM EDTA, pH 7.1). The mixture was allowed to equilibrate for at least 20 min at 4°C to ensure complete association. The drug stock was prepared so that upon reaching the desired concentration of drug, the amount of DMSO in the sample was no >1% of the total volume. For absorption studies, a constant concentration of drug (7μM) was titrated with increasing amounts of RNA (0.5–20 μM) and the absorbance was measured over the range of 400–600 nm on a Cary 50 Bio UV-vis spectrophotometer. Melting studies were performed over a temperature range of 25–90°C.
using a melting rate of 1°C/min and data were collected every half degree. In order to favor hairpin formation and avoid self-association, constructs were heated at 99°C for 1 min, then slowly cooled (1°C/min) to 10°C prior to being melted from 25°C to 90°C (33). For melting experiments, absorbance was measured at 260 nm, the temperature was controlled with a Quantum Northwest TC 125 temperature control unit, drug concentration varied between 5 and 20 μM, and the concentration of RNA was held constant at 2.5 μM. The \( T_m \) was taken as the midpoint of the melting transition as determined by the maxima of first derivative plots. An example of the first derivative plots for ferritin H-chain IRE RNA in the absence and presence of both DOX and DAU can be found in the Supplementary Material (Supplementary Figure S1). All \( T_m \)'s were reproducible to within ± 1°C. Repeated recording of heating and cooling profiles indicates that no hysteresis occurs in the absence or presence of ligand.

**Fluorescence quenching experiments**

Quenching of the drug's natural fluorescence upon binding to RNA was monitored via titrations of a constant concentration of drug (7 μM) with RNA of increasing concentration (0.1–20 μM). Quenching experiments were performed in a 96-well black microplate following the protocol described by Hergenrother et al. (2). Fluorescence emission at 556 nm was recorded on a Molecular Devices SpectraMAX Gemini microplate reader after excitation at 480 nm and 485 nm for DOX and DAU respectively. All experiments were repeated at least four times. The data collected in the fluorescence quenching titrations was fit to a second-order decay curve utilizing TableCurve 2D v5.01, equation 8107 (2):

\[
y = \frac{a \times b}{b + x}
\]

Where \( b \) is the apparent dissociation constant, \( x \) is the RNA concentration and \( a \) is the asymptotic limit.

Fluorescence quenching plots and curve fits are available in the Supplementary Material.

A stock solution of 4 M NaCl in BPE buffer was diluted with BPE to achieve the various salt concentrations listed in the paper (34).

**Rnase T1 footprinting**

The protocol for drug–RNA footprinting was adapted from previously reported methods (35). RNA constructs were 5'-end-labeled with \( \gamma^{32} \)P ATP (PerkinElmer, Waltham, MA) utilizing T4 polynucleotide kinase (Invitrogen, Carlsbad, CA) according to the manufacturer’s protocol. Labeled RNA (1.25 μM), unlabeled RNA (11.5 μM) and drug (1–20 μM) were combined in a final volume of 9 μl of footprinting buffer (11 mM Tris-HCl, pH 7.0) and were allowed to equilibrate for 30 min at 25°C. The equivalent of 1 U of RNase T1 enzyme (USB) was added and allowed to proceed for 10 min at 25°C. The reaction was quenched by the addition of 5 μl of urea loading buffer (8 M urea, 20 mM EDTA and 2 mM Tris, pH 7.5). Samples were then heated at 95°C for 5 min, and 11 μl of each sample was run on a 12% denaturing acrylamide gel for 90 min at 290 V. The gel was then dried, exposed to an IP plate (Fuji Film) and was visualized on a Fuji Film FLA-3000 IP plate scanner. Quantification of band intensity was performed with the Fuji Film program Multi Gauge version 3.0 utilizing profile quantification in the direction of migration and a polygonal baseline for background compensation.

**RESULTS**

**UV melting and absorbance experiments**

The magnitude of ligand-induced change in nucleic acid \( T_m \) \( \Delta T_m = T_m(RNA + ligand) - T_m(RNA) \) is an indication of the affinity of the ligand for the nucleic acid target. Hence, we performed UV melting experiments to obtain the first indication of a direct association between the drugs and the IRE RNAs.

The UV melting curves of both ferritin H-chain and L-chain IREs (Figure 2) display bi-phasic melting profiles in the absence and presence of drug. The higher temperature transition \( T_m = 61^\circ \)C is a minor transition and was found to remain constant under all conditions. The lower temperature transition revealed a concentration-dependent increase in \( T_m \) in the presence of DOX and DAU. The observed \( \Delta T_m \) for all construct and drug combinations are presented in Table 1. It was found that the addition of drug to the IRE RNAs caused an increase in \( T_m \) not only in a concentration-dependent manner but also in

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Overlay of the melting profiles of (a) ferritin H-chain mRNA and (b) ferritin L-chain mRNA in the absence (solid) and presence (dashed) of 20 μM DOX.
Table 1. The average $\Delta T_m$s observed for all drug and IRE combinations

| Ligand | H-chain | L-chain | G23I-IRE | U10C-IRE | U6C-IRE | C8A-IRE | U6C-U10C-IRE |
|--------|---------|---------|----------|-----------|---------|---------|---------------|
| 5$\mu$M DOX | 2.5     | 2.7     | 4.7      | 3.7       | 2.6     | 3.9     | 1.0           |
| 20$\mu$M DOX | 9.9     | 10.1    | 10.3     | 6.8       | 6.5     | 7.9     | 5.5           |
| 5$\mu$M DAU | 0.5     | 3.5     | 3.8      | 1.7       | 1.5     | 2.9     | 0.5           |
| 20$\mu$M DAU | 5.7     | 6.5     | 7.3      | 4.7       | 4.0     | 5.9     | 1.5           |

Average $T_m$ values can be found in the Supplementary Material (Table S1).

a drug-dependent manner, with DOX consistently causing a more substantial shift than DAU (Table 1, Supplementary Figure S1). It was also observed that the slight changes in sequence and structure of the constructs led to a graded degree of $T_m$ shift in response to the presence of drug. In the presence of 20$\mu$M DOX, both the ferritin heavy and light IREs produced a $\Delta T_m$ of around 10°C (Table 1). Comparatively, the HIV-1 TAR RNA construct (6,36), which is also a hairpin-loop structure that has been shown to interact with small molecules, produced a maximum shift of 5°C under the same conditions (data not shown).

Guided by the results from the RNase footprinting experiments on wild-type IREs (see details below), mutations were made to the native H-chain IRE that altered or removed certain structural features associated with drug binding and recognition (37,38). We constructed four point mutants designated U6C-IRE, C8A-IRE, U10C-IRE and G23I-IRE (I = inosine) with mutations at nucleotide positions 6, 8, 10 and 23, respectively; and one double mutant designated U6C-U10C-IRE, bearing double mutations that replaced the two UG pairs with CG pairs. These mutations were predicted by RNAstructure to preserve the IRE secondary structure. The difference in $\Delta T_m$ between these mutants and the wild-type indicated the importance of both UG wobble base pairs found in the ferritin IRE. The observed $\Delta T_m$ for the double mutant, U6C-U10C-IRE, which replaces both UG wobble base pairs, was 5.5°C and 1.5°C in the presence of 20$\mu$M DOX and DAU, respectively. For the G23I-IRE mutant, the magnitudes of the observed $\Delta T_m$ are slightly higher than that of the wild-type for both DOX and DAU at all concentration studied (see Supplementary Material for melting curves). Interestingly, U6C-IRE and U10C-IRE, mutants in which one of the UG pairs is replaced with a CG, have identical ligand-dependent $\Delta T_m$. However, their observed $\Delta T_m$ is about 3°C less than that observed with the wild-type RNA in the presence of 20$\mu$M DOX.

Figure 3a shows the changes that occur in the absorption spectrum of DOX in the visible region during the titration of a fixed concentration of DOX with increasing concentrations of the wild-type H-chain IRE. The resulting absorbance profile of DOX yielded two distinct absorbance maxima at ~472 nm and 499 nm and a marked bathochromic shift was observed for both peaks resulting in a $\Delta \lambda$ of 5 nm and 10 nm, respectively. Additionally, a hypochromism of 33.7% for the first maxima and 24.4% for the second was observed.

Fluorescence quenching

The natural fluorescence of DOX and DAU upon excitation was monitored as they were titrated with increasing amounts of RNA. The fluorescence spectra show a quenching of DOX fluorescence signal upon titration with IRE RNA (Figure 3b). The decay curves of the fluorescent signal were fitted, and apparent $K_q$s were calculated. It is important to note that this apparent $K_q$ is a total dissociation constant, and does not differentiate between the contribution of direct drug binding (e.g. intercalation) and the contribution of electrostatic interaction due to the charged nature of the anthracycline molecule. The apparent $K_q$s and associated standard errors of RNA constructs are presented in Table 2.

Fluorescence quenching data at 25, 28, 30 and 35°C was used to perform a Stern–Volmer analysis (34,39,40) of ferritin H-chain IRE RNA quenching DOX according to the equation:

$$I_0/I = 1 + K_{SV} [Q]$$

Where $I_0$ and $I$ are the fluorescence intensities in the absence and presence of RNA, respectively, $[Q]$ is the concentration of quencher (RNA) and $K_{SV}$ is the dynamic quenching constant related to the bimolecular collision process. For the quenching experiment at 25°C, this analysis revealed a linear relationship between $I_0/I$ and $[Q]$ (Figure 3c). The linearity of the Stern–Volmer plot indicates that the mode of quenching is either collisional or static. Collisional and static quenching can be distinguished by their different dependencies on temperature. An increase in temperature leads to an increase in the diffusion constant of the quencher and will result in an increase in collisional quenching. In contrast, an increase in temperature will lead to a decrease in the binding constant of quencher for fluorophore and will result in a decrease in quenching for a static quencher (40). To identify the quenching mechanism, we repeated the quenching experiments at 25, 28, 30 and 35°C. Stern–Volmer analysis of the resulting data revealed linear relationships between $I_0/I$ and $[Q]$ and also showed that the $K_{SV}$ is inversely correlated with temperature (Figure 3c). Based on this result, it could be concluded that the probable quenching mechanism of the DOX-H-chain IRE–RNA complex is initiated by static quenching (39,40).

To determine the drug/RNA stoichiometry, we use the fluorescence quenching data to construct Job plots (Figure 4) (41,42). The x-axis of the Job plot, labeled as the mole fraction, is the ratio of the concentration of RNA to the total concentration of RNA and drug, or: Mole
Fraction = [RNA]/([RNA] + [drug]). In this case, the sum of [RNA] and [drug] was held constant at 7 mM. The intersection of the two linear portions of the Job plot yield the mole ratio corresponding to the approximate binding stoichiometry between drug and RNA. This analysis revealed a mole fraction corresponding to 3.0 molecules of drug associating with each molecule of the H-chain IRE RNA.

To further confirm the participation of the UG pairs in anthracycline binding, we also constructed Job plots using the fluorescence quenching data of the mutants IREs. We obtained a binding stoichiometry of 2.0 molecules of drug associating with each molecule of either U10C or the double mutant U6C, U10C (See Supplementary Figure S3).

Binding energy

In an effort to partition the binding free energies of anthracyclines with the IRE RNAs into their nonelectrostatic and polyelectrolyte contributions, fluorescence
quenching was monitored over a range of Na⁺ concentrations, and a Stern–Volmer plot for each was plotted (Figure 3d) (25). The quenching constant ($K_{sv}$) was determined for each curve and was then used to calculate the standard Gibbs free energy ($\Delta G^\circ$) using the equation:

$$\Delta G^\circ = -RT \ln K_{sv}$$

Where R is the gas constant (1.987 cal/K/mol) and T is the temperature in Kelvin. The quenching constants estimated at 25, 28, 30 and 35 °C are $6.2 \times 10^5$ M$^{-1}$, $5.0 \times 10^5$ M$^{-1}$, $4.6 \times 10^5$ M$^{-1}$ and $3.8 \times 10^5$ M$^{-1}$, respectively. A subsequent plot of log $K_{sv}$ versus log [Na⁺] yielded the salt dependence of the binding constant defined by the slope (SK) of such plots. The SK value (−0.83 for DOX and ferritin H-chain IRE) was used in the following equation to determine the polyelectrolyte contribution ($\Delta G_{pe}$) at a given salt concentration (25,43):

$$\Delta G_{pe} = (SK) RT \ln [Na^+]$$

To determine the nonelectrostatic contribution to binding energy, the thermodynamic free energy change ($\Delta G_t$) was calculated according to the equation (25):

$$\Delta G_t = \Delta G^\circ - \Delta G_{pe}$$

The values obtained for the binding of DOX to H-chain IRE RNA (BPE buffer, 16 mM Na⁺, 25°C) are as follows: $\Delta G^\circ = -7.7$ kcal mol$^{-1}$, $\Delta G_{pe} = -2.0$ kcal mol$^{-1}$ and $\Delta G_t = -5.7$ kcal mol$^{-1}$. The error associated with these values is estimated to be ± 0.1 kcal mol$^{-1}$ (25). A comparable value of $\Delta G_{pe}$ has been reported for the binding of DOX to DNA under similar conditions (25).

**Drug–RNA footprinting**

RNase footprinting was performed in order to probe the binding site(s) of DOX and DAU on the RNA constructs. Of the several RNases investigated, RNase T1 was found to provide the cleanest and most useful cleavage pattern. RNase T1 cleaves single-stranded RNA at the 3'-end of guanylic residues and, as expected, it caused strong cleavage of the RNA at G7, G16 and G18, all residues predicted by RNAstructure to be within the single-stranded region of the IRE secondary structure (Figure 5a). The most striking observation from this autoradiograph is that T1 cleavage at G22/G23 and G26/27, the nucleotide positions corresponding to the two UG sites, displayed a drug concentration-dependent increase in band intensity (Figure 5a). While G16 also showed some variation in band intensity, comparison of the quantitated G16 and G23 band intensities, after the footprinting of the H-chain IRE with DOX, revealed that G23 displayed a consistent and marked concentration-dependent increase in cleavage intensity compared to G16 (Figure 5b).
We then used RNase T1 footprinting to follow the effects of the aforementioned mutations on cleavage at G23, the guanylic residue of one of the UG pairs (U10G23) that showed T1 cleavage enhancement in the presence of DOX. We chose to monitor G23 because it is the better resolved of the two G residues on the autoradiograph (Figure 5a). In a manner similar to the wild-type H-chain IRE, the footprinting of the G23I mutant with DOX displayed a drug concentration dependent increase in band intensity at G23 (data not shown). Conversely, the footprint of the U10C-IRE, the mutant which replaced the top UG pair, lacks the band corresponding to G23 when compared to that of the wild-type H-chain IRE (Figure 5a). Validation of the cleavage band designations can be found in the supplementary material (Supplementary Figure S4).

**DISCUSSION**

Anthracycline–DNA interactions are among the best characterized in the literature (24–32). Despite their chemical similarity to DNA and direct role in gene expression, little is known about the extent to which RNA interactions with anthracyclines contribute to anthracyclines’ biological activities. In this article, we demonstrate that DOX and DAU directly interact with the IREs hairpins (Figure 1a and b) located in the 5′-UTR of ferritin H- and L-chains mRNAs. DOX and DAU caused a concentration-dependent shift in the melting profiles of the H- and L-chains IREs (Figure 2 and Supplementary Figure S1). In most cases, we observed higher \( \Delta T_m \)s for DOX compared to DAU for all IRE RNAs studied under the same conditions (Table 1). Since the magnitude of \( \Delta T_m \) is an indication of the affinity of a probe molecule for a nucleic acid target (44), the noted differences in the values of \( \Delta T_m \)s suggest that DOX binds tighter to these RNAs compared to DAU. The lower \( \Delta T_m \)s observed for DAU (Table 1, bottom half) is consistent with the literature report indicating that DAU binds to DNA between 1.5- to 4-fold less strongly than DOX (31). Furthermore, the only structural difference between the two drugs is the presence of a primary OH-group at C14 of DOX (Figure 1c); it is therefore possible that the OH-group at C14 is playing a role in the binding of anthracyclines to the RNAs, perhaps in a similar manner to its involvement in DNA binding and recognition (45).

The drug induced changes in the melting profiles of the mutant IREs gave further evidence for the base sequence preference of the interactions between the anthracyclines and the IRE RNAs. While the asymmetric bulge in the H-chain IRE is somewhat tolerant of a base change, the replacement of either or both of the UG wobble pairs flanking the asymmetric bulge with a CG significantly impaired DOX and DAU binding (Table 1). These results point to the requirement of the UG base pairs in facilitating the binding of DOX and DAU to the IRE RNAs. However, the UG base pair recognition by anthracyclines may be RNA sequence dependent. Preliminary evidence to that effect was seen in the melting profile of a C22U mutant TAR RNA. Relative to the wild-type TAR RNA, we observed almost no change in \( \Delta T_m \) of the C22U mutant in the presence of 20 \( \mu \)M DOX. This result may suggest that the microenvironment of the UG pairs within the RNA sequence may play important role in anthracycline recognition (see Supplementary Figure S2, for the melting profiles of wild-type and C22U mutant TAR RNA).

The ability of IRE RNAs to alter the spectral characteristics of anthracyclines gives further indication of an interaction of drug and RNA (Figure 3a and b). We observed a quenching of the natural fluorescence of DOX and DAU with increasing RNA concentration (Figure 3b). The quenching of DOX fluorescence by DNA has been ascribed to its intercalation into DNA (27,46). Thus, we attribute the extensive fluorescence quenching (>90% at a 2:1 ratio of DOX to H-chain IRE RNA) to the intercalation of one or more anthracycline molecules per RNA molecule (Figure 3b). A similar interpretation has recently been proposed to explain DOX fluorescence quenching by a SELEX generated DOX RNA aptamer (5). Additionally, the titration of a fixed concentration of DOX with increasing concentrations of the wild-type H-chain IRE results in a hypochromic effect accompanied by a shift of the absorption maximum (\( \lambda_{\text{max}} \)) to a longer wavelength (bathochromic shift). The observed maximal absorbance decrease is \( \sim 33.7\% \) (Figure 3a). These results are indicative of changes in the drug’s environment and a perturbation of the DOX chromophore system upon binding to the IRE RNA, consistent with intercalative binding of DOX within the IRE RNA. Furthermore, the apparent \( K_d \) (Tables 2) extracted from the fluorescence quenching data give an indirect clue about the affinity of DOX for the IREs. Although we did not have sufficient information to differentiate between the contribution of direct drug binding and that of electrostatic interaction for all IRE constructs, the magnitudes of the apparent \( K_d \)’s suggest strong association of the drug with the IREs.

We performed RNase footprinting experiments to gain more insight into the binding site(s) of DOX and DAU on the ferritin H-chain IRE and a series of mutant RNAs (47,48). Results from the drug–RNA footprinting experiments show that the T1 cleavage at two sites, G22/G23 and G26/27, is enhanced upon addition of DOX (Figure 3a). Chaires et al. (48) have attributed an analogous enhancement in nuclease cleavage of DNA, in the presence of DAU, to the perturbation of DNA structure in the immediate vicinity of a DAU intercalation site. If this explanation holds for RNA, our results suggest the existence of at least two main drug binding sites at or near the two UG wobble base pairs of the ferritin H-chain IRE (Figure 5). Interaction of anthracycline molecules at both of these proposed binding sites would not violate the nearest neighbor exclusion rule when applied to anthracycline–nucleic acid binding (27).

Wobble pairs, especially GU or UG, are ubiquitous building blocks of higher order RNA structures. Compared to regular Watson–Crick base pairs, wobble pairs differ in both the array of functional groups displayed in the helical grooves and the orientation of the bases with respect to the phosphodiester backbone.
The consequence of this is that a wobble pair causes weakening of the helix and local distortions which significantly increase the electronegative environment at or near the mismatch pair (49,50). The two UG pairs in the ferritin H-chain IRE could each provide an electronegative environment for nestling the electropositive DOX. In fact, the wobble pair has been shown to be a commonly conserved ligand binding site (49). To further probe this observation, we extensively studied the effects of mutation of the two UG pairs and bulge C on the binding affinity of DOX for the IRE RNA. Results from these experiments show that all mutations that preserved the U10-G23 wobble displayed essentially the same cleavage pattern as in the wild-type RNA. This suggests that these mutants are able to induce the structural changes necessary for DOX binding within the vicinity of U10-G23. More importantly, however, the G23I mutant sheds light on the specifics of the UG pair recognition by DOX. The G23I mutation results in the deletion of the guanosine N2 exocyclic amine, a very distinct functional group displayed in the minor groove of a wobble that often serves as a specific receptor recognition signal. The fact that the G23I-IRE mutant also displayed increase in the magnitudes of the observed ΔTm for the G23I-IRE mutant are slightly higher than that of the wild-type in the presence of either DOX or DAU at all concentration studied (Table 1). These data suggest that the substitution of G23 for I may have resulted in slightly increased RNA ligand-binding affinity. Previous literature observation indicates that an IU base pair is slightly less stable than a GU pair (50). The subtle duplex instability as a result of the G23I mutation may enhance the structural changes essential for ligand binding, thereby resulting in increased interaction of ligand with the RNA. Second, the G23 band in the U10C, a mutant IRE in which the G23-U10 is substituted by a GC base pair, is protected from T1 cleavage at G23 (Figure 5). The protection of G23 from T1 digestion may be due to diminished or lack of DOX binding, thereby preventing the necessary conformational changes that expose G23 to T1. Alternatively, it may also be that DOX binding persists but is not accompanied by a conformational change that is sensitive to T1 cleavage.

The Stern–Volmer analysis of the drug fluorescence quenching data supports a static mechanism for the quenching of the DOX fluorescence by the H-chain IRE RNA (Figure 3c and d). The static quenching could occur through the formation of a complex, stabilized mainly by hydrophobic interactions, between DOX (fluorophore) and RNA (quencher) (39,40). This complex could result from the intercalation of DOX into the IRE bases as suggested by the aforementioned evidence. In addition, the Stern–Volmer analysis allows an estimation of the equilibrium constant for the formation of the DOX-H-chain IRE RNA complex. Applying the polyelectrolyte theory (25,43), these equilibrium constants enable the dissection of the RNA binding free energies of anthracyclines into their nonelectrostatic and polyelectrolyte contributions. The calculated value of ΔGpe is very similar to that reported for the binding of DOX to DNA under similar conditions (25). This result suggests a similar polyelectrolyte contribution to the overall ΔG of binding of DOX to DNA and RNA. A Job plot of the fluorescence quenching of DOX with wildtype IRE RNA gives a drug to RNA binding ratio of ~3.0:1 (Figure 4). A similar plot of the U10C IRE, a mutant in which one of the proposed binding sites has been replaced with a GC pair, resulted in an drug to RNA stoichiometry of 2.0:1 (Supplementary Figure S3). This result is consistent with our original inference on the participation of U10-G23 wobble pair in DOX binding to the IRE. However, the drug to RNA stoichiometry remained 2.0:1 when both UG pairs are replaced (Supplementary Figure S3). The lack of further change in the drug to RNA ratio upon the replacement of the proposed second DOX binding site may be due to the fact that the second mutation did not completely abolish DOX binding. Although footprinting only identified the two UG sites as possible binding sites, the nonelectrostatic binding energy, ΔGpe, was found to account for only 74% of the total observed change in ΔG upon DOX binding to IRE RNA. Taken together, these data suggest that in addition to the intercalation of two anthracycline molecules at the UG sites, there is at least one additional site of interaction, most likely electrostatic in nature, on the IRE RNA.

The crystal structure of IRP1 bound to frog ferritin IRE has recently been solved (51). The structure reveals almost twice the number of specific interactions found in most protein–RNA associations. Despite the level of interaction, single mutations to just a few of the residues involved have been shown to greatly diminish the IRE/IRP interaction (37,52,53). This high degree of structural and sequence selectivity reveals the seemingly robust interaction to be in fact very sensitive to minute disturbances. The binding of anthracycline molecules to the IRE structure could cause changes to the tertiary structure of the IRE RNA that could diminish the affinity of IRP for the IRE. This may cause an aberration in the intracellular regulation of the translation of mRNAs harboring these IREs. Supporting this hypothesis is a recent observation from the Thorp group on up-regulation of the cellular expression of ferritin upon exposure to yohimbine, a small molecule that binds to the ferritin IRE (3).

The biochemical evidence presented in this study suggests that anthracyclines form strong and specific interactions with IRE RNAs. Furthermore, it sheds light on some of the structural requirements for this interaction. Although anthracyclines such as DOX and DAU have been reported to show a requirement for GC sequences in DNA binding (48,54), we have presented evidence that...
the binding of anthracyclines to ferritin IRE RNA demonstrates a strong preference for UG pairs. The prevalence of wobble base pairs in higher order RNA structures (49) suggests that this non-canonical pair, when structurally available, could serve as a ubiquitous recognition site for the binding of anthracyclines to other RNAs as well. Further studies are necessary to better understand the possible biological significance of the anthracycline-IRE interaction. Part of our ongoing effort is to probe the effects of DOX and DAU on IRE-IRP complex formation both in vitro and in vivo.

SUPPLEMENTARY DATA
Supplementary Data are available at the NAR Online.

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