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Electrostatic Potential in the tRNA Binding Evolution of Dihydouridine Synthases

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ABSTRACT: Dihydouridine (D) is an abundant modified base of tRNA found in the majority of living organisms. This base is synthesized via an NADPH-dependent reduction of specific uridines by the dihydouridine synthases (Dus), a large family of flavoenzymes comprising eight subfamilies. Almost all of these enzymes function with only two conserved domains, an N-terminal catalytic domain (TBD) adopting a TIM barrel fold and a unique C-terminal helical domain (HD) devoted to tRNA recognition, except for the animal U20-specific Dus2 enzyme. Curiously, this enzyme is distinguished from paralogues and its fungi orthologues by the acquisition of an additional domain, a double stranded RNA binding domain (dsRBD), which serves as the main tRNA binding module. On the basis of a homology model of yeast Dus2 and the crystallographic structure of a human Dus2 variant (TBD + HD) lacking dsRBD, we herein show that the HD surface of the human enzyme is less electropositive than that of its yeast orthologue. This is partly due to two positively charged residues, K304 and K315, present in yeast and more broadly in fungi Dus2 that are replaced by E294 and Q305 in human and conserved among animals Dus2. By artificially reintroducing these positive charges in human Dus2 lacking dsRBD, we restored a functional tRNA binding in this enzyme variant. Altogether, these results suggest that the electrostatic potential changes of HD have likely played a key role in the emergence of a new tRNA binding mode among Dus2 enzymes.

Structural modularity has emerged as an essential adaptive feature as a new functional requirement in many enzymes.1−3 Typically, enzymatic modularity implies the existence of several domains, whose number increases according to the complexity of organisms.4 Although modularity can serve various purposes,3 it is often used by enzymes to separate the catalytic function from that of substrate recognition. Nucleic acid-targeting enzymes are known for carrying a particularly complex modularity, whose degree of complexity perfectly correlates with the biological complexity of organisms. Enzymes that mature and functionalize RNAs such as amino-acyl tRNA synthetases,5,6 RNases,7,8 or the large family of RNA editing and modification enzymes,9 to mention only a few of them, obey to this evolution’s principle.

Belonging to tRNA-modifying enzymes, dihydouridine synthases (Dus) catalyze the reduction of the C5–C6 double bond of uridines present at different sites on tRNAs to generate dihydrouridine (D) (Figure 1A,B).

Dihydouridine is important for stabilizing RNA functional fold,11,12 and it has been shown that tRNAs lacking this modified base in combination with others are rapidly degraded by cellular RNases.13 Dus form a broad family of modular flavoenzymes that use the flavin mononucleotide (FMN) as a prosthetic group and NADPH coenzyme as a source of hydride.14,15 There are eight Dus subfamilies distributed across the living world.15 Three are found in bacteria (DusA, B, and C), one in archaea, and four in eukaryotes (Dus1, 2, 3, and 4). So far, only a few of these enzymes have been studied, while most of them remain poorly characterized despite their significant biological relevance. Regarding the function, in vivo and in vitro studies of Escherichia coli16−18 Thermus thermophilus19,20 and Saccharomyces cerevisiae21−23 enzymes allowed the substrate specificity of Dus subfamilies to be established (Figure 1B). In addition, bioinformatics analysis15 and recent X-ray crystal structures10,18,19,24,25 established that the majority of Dus enzymes function only with two domains. These are, namely, an N-terminal domain serving as a catalytic module organized into a TIM-barrel wherein FMN lies in its center (TIM barrel domain = TBD) and a C-terminal domain formed by α helices and used as a tRNA binding module (helical domain = HD) (Figure 1C). Thus, TBD + HD constitutes the canonical Dus structure. Intriguingly, we have recently shown that human dihydouridine synthase 2 (hDus2)
and, based on sequence alignments, probably all animal Dus2 display a more complex modularity than its Dus paralogues or orthologues. In addition to the canonical Dus structure, we showed that this enzyme possesses a third module, the double stranded binding domain (dsRBD), appended at the C-terminus of HD, and an additional ∼50 amino acid peptide predicted as intrinsically disordered that ends the protein (Figure 1C,D). Moreover, this dsRBD seems to have evolved to function as the main tRNA binding domain of hDus2, suggesting that during evolution the tRNA binding function has been transferred, at least in part, from the TBD + HD part to the dsRBD. Why animal Dus2 specifically enjoys such a complex modularity while all other Dus and notably their closest relative fungi Dus2 function without dsRBD remains a mystery. It is more puzzling that this dsRBD does not seem to ensure a better thermodynamic stability of the tRNA/Dus2 complex since hDus2 and its yeast orthologue Dus2p (Dus2 from *Saccharomyces cerevisiae*) share a similar affinity for tRNA. However, one cannot exclude that this modularity could serve other cellular purposes, which remain to be demonstrated. Indeed, in addition to its tRNA modification function, human Dus2 has been shown to promote cell growth in some cancers mediated via its ability to interact with the amino-acyl tRNA synthetase complex EPRS or protein kinase R. However, its exact pro-oncotic mechanism is not yet elucidated.

Using a comparative approach combining bioinformatics, structural and functional analysis, we here identified that HD of bimodular Dus2, specific to fungi, presents a more electrostatic potential than that of its trimodular Dus2 orthologue, specific to animals. By reintroducing these lacking positive charges into the HD of a truncated hDus2 version (hDus2TBD+HD) in which both dsRBD and C-terminal region were removed, we managed to generate a new functional canonical Dus2 carrying a better affinity for tRNA than its parent counterpart. These results lead us to speculate that this important physicochemical electrostatic property has played a major role in the evolutionary process that led to the complex modularity of current animal Dus2.

### MATERIALS AND METHODS

**Protein Purification.** Wild-type human Dus2 (hDus2) and Δ(2–13)-hDus2<sup>TBD+HD</sup>, a construct lacking residue Ile2-Lys13 and Q341-W493, were cloned in pET11d vector between BamHI and NcoI while hDus2<sup>TBD+HD</sup>, a construct lacking Q341-W493, and Dus2p (Dus2 from *Saccharomyces cerevisiae*) were cloned in pET15b between BamHI and NcoI. All constructs were expressed in BL21(DE3)pLysS (Novagen) in LB medium. Induction with 0.5 mM isopropyl-1-thio-β-D-galactopyranoside (IPTG) was done at OD<sub>600</sub> ~ 0.6 and left overnight at 29 °C. Pelleted cells were suspended in buffer A (50 mM sodium phosphate pH 8, 2 M NaCl, 25 mM...
imidazole, and 10% glycerol) with the addition of phenylmethylene sulfonyl fluoride (PMSF) and lysed by sonication followed by centrifugation for 40 min at ~148 000g. The supernatant was loaded on a NiNTA column (GE healthcare) washed several times with buffer A and eluted with buffer A supplemented with 250 mM imidazole. All proteins were further purified by a size exclusion chromatography step on a Superdex HiLoad S200 16-600 unit (GE healthcare). Purity was assessed by SDS-PAGE. Point mutations were carried out with the Q5 site-directed mutagenesis kit (New England BioLabs) following the recommended procedure.

Preparation of Bulk tRNA. To extract bulk tRNA from yeast strain BY4741 Δdus2 derivative (MATa; his3Δ1; leu2Δ0; met15Δ0; ura3Δ0; YNR015w::kanMX4), cells were grown in 500 mL of YPD (peptone 2%; yeast extract 1% and glucose 2%) to an OD_{600} of 0.8. Pelleted cells were washed twice in 20 mL of TMN (10 mM Tris–HCl (pH 7.5), 10 mM MgCl2, 0.15 M NaCl). The final pellet was resuspended in 20 mL of TMN and 20 mL of acidic phenol (pH 4.5) and incubated for 20 min at room temperature on a rotating wheel. After centrifugation, the aqueous phase was recovered, supplemented with LiCl to a final concentration of 0.8 M, and incubated overnight at 4 °C to precipitate high molecular mass molecules. The precipitate was eliminated by centrifugation, and the supernatant was supplemented with two volumes of 100% ethanol and incubated at −20 °C for 2 h to precipitate tRNAs. After centrifugation, pellets of tRNAs were washed twice in 70% ethanol and resuspended in 1 mL of RNase-free water. tRNAs were desalted and concentrated four times to 50 μL in Centricon YM-3 devices (Millipore) using 100 mM ammonium acetate (pH 5.3) as a final buffer.

Crystallization, Data Collection, Structure Determination, and Refinement. Crystals of Δ(2–13)-hDus2TBD+HD E294K mutant were obtained by vapor diffusion by mixing 1 μL of freshly prepared protein in 50 mM Tris pH 8 and 150 mM NaCl with 1 μL of reservoir comprised of 30% PEG 2000 MME, 200 mM ammonium sulfate, and 50 mM sodium acetate pH 5.5. Crystallization of Δ(2–13)-hDus2TBD+HD Q305K mutant required 2.2 M ammonium sulfate, while Δ(2–13)-hDus2TBD+HD E294K/Q305K mutant crystallized in 30% PEG 2000 MME, 200 mM ammonium sulfate, and 50 mM sodium acetate pH 5. Crystals were obtained after 2 days and were cryoprotected using 15% glycerol before flash freezing in liquid nitrogen. The three structures were phased by molecular replacement using Phaser38 and as a starting model PDB entry 4XP7. All diffraction data were collected at 100 K on single crystals at the microfocused beamline PROXIMA-2 equipped with an Eiger X-27 M detector at SOLEIL synchrotron (Saint-Aubin, France). Data were indexed using XDS, scaled with STARANISO using the ap_scale module of autoPROC to correct the anisotropy of the data.29 The three structures were refined with autoBUSTER,30 and manual building into the density was performed in Coot.31 TLS were used with one TLS group per protein chain. STARANISO corrected intensities and structure factors as well as uncorrected intensities and structure factors without resolution cutoff have been deposited in the Protein Data Bank along with the coordinates and Fourier coefficients. (See Table S1 for the determination of best-fit ellipsoid to the anisotropic cutoff surface.)

Homology Modeling and Electrostatic Potential Analysis. Homology models of Dus2p were built by homology-modeling servers, I-TASSER,32 SWISS-MODEL,33 Phyre2,34 and ROBETTA,35 that all used PDB 4XP7 as a template. The resulting models were validated by the server ProQ3D.36 The calculation of protein electrostatic surfaces was performed by software APBS (v1.4).37 Before calculations, missing atoms from the side chains were modeled using the PDB2PQR software.38 Calculations were performed at 310 K with 150 mM NaCl with the same grid size (193, 193, 161) for all calculations. For all of the representations, we choose to show an electrostatic potential within ±3 kT e⁻¹.

tRNA Binding Assay. Electrophoretic mobility shift assays were carried out on a 6% native (19:1) PAGE at 4 °C with 100 V. Increased concentrations of proteins were added to a fixed concentration of bulk tRNA lacking D20 (1 μM) and incubated at room temperature for 20 min in 50 mM Tris pH 8, 10% glycerol, 5 mM DTT, and 150 mM ammonium acetate prior to migration. RNA was visualized by toluidine coloration and quantified using ImageJ (https://imagej.nih.gov/ij/index.html).

NADPH Oxidase Activity. Oxidase activities were assessed under steady state conditions with menadione as a final electron acceptor in 50 mM sodium phosphate pH 8, 150 mM NaCl, and 20% glycerol at 23 °C. Two μM of holoprotein was mixed with various concentrations of NADPH and 200 μM menadione. NADPH oxidation was followed by the decrease in absorption at 340 nm using a Cary-50 spectrophotometer. Data were analyzed using Michaelis–Menten formalism.

Dihydrouridine Activity Assay. In vitro activity was assayed for 30 min at 30 °C in 50 mM Tris pH 8, 150 mM ammonium acetate, 1 mM dithiothreitol, 2 mM MgCl2, 10% v/v glycerol. Bulk tRNAs (100 μM) were incubated with 5 μM of protein in a total volume of 50 μL, and the reaction was started with the addition of 2 mM NADPH. Quenching was performed by adding 50 μL of acidic phenol (Sigma-Alrich) followed by centrifugation at ~12 000g for 10 min, and tRNA in the aqueous phase was ethanol precipitated and further purified using a MicroSpin G-25 column (GE-healthcare). Dihydrouridine quantification was carried out by means of a colorimetric method as described previously.10 Briefly, samples were incubated at 40 °C for 30 min after the addition of 5 μL of 1 M KOH. The solutions were neutralized with 25 μL of 96% H2SO4 followed by 25 μL of a 3% solution of 2,3-butanediol monoxime (Sigma-Alrich) and 25 μL of a saturated solution of N-Phenyl-p-phenylenediamine (Sigma-Alrich). Samples were then heated at 95 °C for 10 min and cooled to 55 °C. Following the addition of 50 μL of 1 mM FeCl3, a violet-red coloration appeared allowing quantification via absorption at 550 nm. D content in tRNA was determined by using a standard curve obtained with variable amounts of dihydrouracil.

RESULTS

Human Dus2 Carries a More Electronegative HD than Its Yeast Orthologue. All nucleic acid enzymes acting on tRNA recognize their substrates primarily via positively charged residues such as arginine and lysine. This led us to speculate that the low affinity of hDus2TBD+HD for tRNA may result from the absence of one or more positively charged residue at the TBD or HD surface or both. One way to address this issue is to compare the electrostatic surfaces of both hDus2TBD+HD and Dus2p and identify regions that present different charge distributions. In the case of hDus2TBD+HD, we used the crystal structure of Δ(2–13)-hDus2TBD+HD as a
Table 1. NADPH Oxidase Activity of hDus2^{TBD+HD} and Its Mutants Using Menadione As a Final Electron Acceptor

<table>
<thead>
<tr>
<th>protein</th>
<th>$k_{cat}$ ($s^{-1}$)</th>
<th>$K_M$ (μM)</th>
<th>$k_{cat}/K_M$ (μM$^{-1}$ $s^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>hDus2^{TBD+HD}</td>
<td>0.8 ± 0.05</td>
<td>63 ± 5</td>
<td>0.013</td>
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<tr>
<td>E294K hDus2^{TBD+HD}</td>
<td>0.7 ± 0.08</td>
<td>70 ± 10</td>
<td>0.01</td>
</tr>
<tr>
<td>Q305K hDus2^{TBD+HD}</td>
<td>0.7 ± 0.04</td>
<td>68 ± 6</td>
<td>0.01</td>
</tr>
<tr>
<td>E294K/Q305KhDus2^{TBD+HD}</td>
<td>1 ± 0.06</td>
<td>55 ± 5</td>
<td>0.018</td>
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Table 2. Summary of Data Collection and Refinement Statistics

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<tr>
<th></th>
<th>HsDus2^{TBD+HD}</th>
<th>HsDus2^{TBD+HD}</th>
<th>HsDus2^{TBD+HD}</th>
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<tbody>
<tr>
<td></td>
<td>E294K mutant</td>
<td>Q305K mutant</td>
<td>E294K Q305K double mutant</td>
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<tr>
<td></td>
<td>PDB 6EZA</td>
<td>PDB 6EZA</td>
<td>PDB 6EZA</td>
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<tr>
<td>Data Collection</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>space group</td>
<td>P 2(1)</td>
<td>I 2 2 2</td>
<td>P 2(1)</td>
</tr>
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<td>cell dimensions $a$, $b$, $c$ (Å)</td>
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<td>72.08, 84.11, 144.35</td>
<td>45.99, 84.66, 100.39</td>
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<td>90, 90, 90</td>
<td>90, 90, 90</td>
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<td>43.61−2.25</td>
<td>43.18−2.00</td>
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<td>0.137 (1.609)</td>
<td>0.161 (2.521)</td>
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<tr>
<td>$R_{free}$</td>
<td>0.160 (1.344)</td>
<td>0.145 (1.714)</td>
<td>0.170 (2.665)</td>
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<tr>
<td>$I/σ(I)$</td>
<td>8.0 (1.3)</td>
<td>12.6 (1.4)</td>
<td>10.1 (1.1)</td>
</tr>
<tr>
<td>completeness (%)*</td>
<td>78.5 (29.3)</td>
<td>78.1 (18.6)</td>
<td>82.9 (27.0)</td>
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<tr>
<td>completeness (%)*</td>
<td>93.1 (93.2)</td>
<td>93.4 (60.8)</td>
<td>89.9 (43.5)</td>
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<td>9.2 (8.5)</td>
<td>9.3 (8.9)</td>
</tr>
<tr>
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<td>0.998 (0.385)</td>
<td>0.997 (0.413)</td>
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<td>42.33−2.00</td>
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<td>22 658 (454)</td>
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<tr>
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<td>0.182/0.223</td>
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<td>79</td>
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<td>no. of atoms, water</td>
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<td>230</td>
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<td>38.56</td>
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<td>48.02</td>
<td>47.67</td>
</tr>
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<td>rms deviation, bond lengths (Å)</td>
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<td>0.010</td>
<td>0.010</td>
</tr>
<tr>
<td>rms deviation, bond angles (deg)</td>
<td>1.06</td>
<td>1.13</td>
<td>1.03</td>
</tr>
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</table>

*Each data set was collected from a single crystal. Data statistics are calculated with STARANISO.
same place as those of Dus2p are also found in the HD of DusA but absent in both DusB and C (Figure S5). We propose that these two residues likely constitute a signature of the U20-specific Dus enzymes in both bacteria and fungi. Taken together, these results suggest that in the family of Dus2 enzymes specifically, the HD has evolved from an electro-positive domain in its closest relatives found in fungi to a more electronegative domain in animals, thereby leading to a domain displaying less tRNA binding capacity and loosening the overall binding of tRNA in hDus2TBD+HD.

Flavin Spectrum and NADPH-Oxidase Activity of E294K, Q305K, and E294K/Q305K hDus2TBD+HD Mutants.

To validate the evolutionary relevance and importance of these residues in the tRNA binding function and to obtain some clues on how the RNA binding function transfer from HD to dsRBD occurred, we proceeded by engineering three hDus2TBD+HD variants, namely, E294K, Q305K, and E294K/Q305K. Although residues 294 and 305 appear to be located far from the flavin binding site (Figure S5), we nonetheless ensured that their replacement did not affect the redox reactivity of FMN by first recording the UV–visible spectrum of each mutant and then assessing its NADPH oxidase activity under steady state conditions. As shown in Figure S6, wild-type and variant proteins display a similar flavin spectrum indicating that these mutations do not disturb the electronic environment of FMN. In addition, they catalyze menadione-dependent NADPH oxidation with apparent $k_{cat}$ and $K_M$ values at 23°C, comparable to those for the parent protein (Table 1).
Therefore, E294K, Q305K, and E294K/Q305K mutations do not disturb the FMN or NADPH binding properties of the proteins.

**X-ray Structure of E294K, Q305K, and E294K/Q305K hDus2^{TBD+HD} Mutants.** Next, we structurally characterized the mutants by solving their X-ray structures. Since hDus2^{TBD+HD} did not crystallize, we used Δ(2–13)-hDus2^{TBD+HD} to analyze the mutations. E294K, Q305K, and E294K/Q305K Δ(2–13)-hDus2^{TBD+HD} mutants crystallized in P 2₁(1), I 2 2 2, and P 2(1) 2(1) 2(1) space groups at 2, 2.25, and 2 Å resolution, respectively (Table 2).

The asymmetric unit of E294K protein is formed by two molecules, while Q305K and E294K/Q305K mutants contain only one molecule. Gel filtration analysis showed that like Δ(2–13)-hDus2^{TBD+HD}, all of the protein mutants are mainly monomers in solution (Figure S7), suggesting that the dimeric asymmetric unit of the E294K mutant is a result of crystal packing. All of the mutants display a structure identical to that of Δ(2–13)-hDus2^{TBD+HD}. Structural overlay shows no significant conformational changes induced by the mutations (Figure 2A).

Nonetheless, while the strictly conserved cys116 of the active site loop (from 115 to 129), which acts as an acid residue during dihydrouridine synthesis\(^{19,22}\) is too flexible to be observed in our Δ(2–13)-hDus2^{TBD+HD} crystal structure, its electronic density is clearly visible in the E294K mutant structure, allowing its manual building (Figure 2B). Lying in the TBD center, FMN is stabilized by several interactions that are all preserved in the mutants (Figure 2C). We then calculated the electrostatic potential of the surfaces of the four hDus2 protein variants. From the electrostatic surface representations, we can clearly observe that these mutations introduce positive charge(s) into HD (Figure 3), the effect being more prominent for the E294K/Q305K double mutant. Hence, mutations convert the HD into a more electropositive domain without altering the overall structure of hDus2^{TBD+HD}.

**E294K, Q305K, and E294K/Q305K hDus2^{TBD+HD} Mutants Increase the tRNA Binding Affinity and Dihydrouridine Activity.** To determine if such mutations enhance the tRNA binding function, we performed electrophoretic mobility shift assays on Dus2p from yeast, full-length hDus2, hDus2^{TBD+HD}, and its mutants (Figure S8). Consistent with our previous report, Dus2p and hDus2 seem to share a similar affinity for tRNA (Figures 4A and S9). The hDus2^{TBD+HD} has a ∼10-fold lower affinity than hDus2 and Dus2p (Figure 4A). Interestingly, introducing a positive charge at position 294 increases the tRNA affinity of hDus2^{TBD+HD} by ∼1.24-fold, while when introduced at position 305 we observed a ∼1.4-fold gain of tRNA binding capacity, which seems to be not a significant improvement. However, the remarkable effect is observed for the double mutant, which exhibits a ∼3.6-fold higher affinity than the parent protein. This result suggests that both charges introduced at positions 294 and 305 act synergistically in tRNA binding. As evidenced from the activity performed in vitro, the gain of the tRNA binding affinity translates into a better dihydrouridine synthase activity, which is comparable to that of full-length hDus2 (Figure 4B).

### DISCUSSION

We have here unambiguously established that the HD domain of hDus2 and more broadly of animals Dus2 are less electropositive than that of its yeast orthologue operating without dsRBD (Figure S1). This deficiency of positive charges explains in part why the canonical hDus2^{TBD+HD} structure is no longer able to ensure a stable enzyme/tRNA complex without dsRBD. By engineering an artificially electropositive HD via introduction of the missing positive charges at positions 294 and 305 conserved in fungi but absent in animals, in hDus2^{TBD+HD}, we were able to increase its affinity for tRNA. This was associated with an improvement of the dihydrouridine synthase activity, suggesting that these positive charges promote the formation of a catalytically competent enzyme/tRNA complex. Although the affinity was improved by ∼3.6-fold with the E294K/Q305K double mutant, we did not restore an affinity comparable to that observed with Dus2p (Figure 4A) or full-length hDus2.\(^{10}\) This suggests that other residues are certainly contributing to this difference, but in the absence of the Dus2p/tRNA complex structure, their identification remains challenging. Regarding hDus2, structural details on the full-length and tRNA-bound enzyme are also lacking. Nevertheless, based on a number of consistent evidence that includes (i) the structure of Δ(2–13)-hDus2^{TBD+HD} and hDus2^{dsRBD}, (ii) the general mode of action of dsRBD, which recognizes ∼12–16 bp dsRNA structure, and (iii) the structure of its bacterial orthologue DusA in complex with tRNA, a schematic model for the hDus2/tRNA complex
has been postulated. In the latter, the dsRBD together with the TBD and HD allow hDus2 to embrace almost the whole substrate. This new tRNA binding mode does not seem to present any obvious advantage in terms of stability for the complex since both Dus2p and hDus2 share a similar affinity for tRNA. Furthermore, we and other groups showed that DusA, Dus2p, and hDus2 are active only on mature tRNA bearing modifications and inactive on a tRNA transcript. This clearly suggests that the canonical Dus structure is endowed with a capacity to discriminate its substrate. Finally, having a dsRBD does not influence flavin reactivity and consequently the catalysis per se, comforting that its role is purely dedicated to tRNA binding. Thus, animal Dus2 represents a single class of modifying enzymes that has partially transferred its tRNA recognition function from a specialized domain, HD, to a rather common dsRBD. This domain already present in LUCA has emerged as a promiscuous RNA binding module for several enzyme family-catalyzing unrelated chemistries. Perhaps, beyond the RNA recognition, dsRBD acquisition may be needed for other unknown functions of Dus2, since in addition to being involved in dsRNA binding, dsRBBDs are known to serve as a protein/protein interaction module in RNA metabolism enzymes, but this remains to be proven.

In the absence of clear alternative functions, what appears unambiguous at this stage is the important role played by the dsRBD in the tRNA recognition function of hDus2. Although there is currently no phylogenetic study addressing such a specific question, we would still like to speculate on possible scenarios that could explain how a novel tRNA binding mode emerged in animal Dus2. Even though several possibilities could have happened, we propose two rather simplistic mechanisms that seem plausible (Figure 5).

In the first scenario, the ancestral animal Dus2 formed by TBD + HD, and closely related to its orthologous relatives in fungi, first acquires a dsRBD by gene fusion, a classical mechanism of protein modularity, leading to an intermediate protein 1 with a very high affinity for tRNA provided by the presence of two RNA binding domains. With this state being most likely not functional because of its excessive affinity that would affect the tRNA binding dynamic, intermediate 1 could have undergone mutations within the HD that decrease its electropositivity, generating animal Dus2. In the second scenario, HD first loses its electropositivity, weakening tRNA binding, and then functionality recovers after dsRBDs acquisition. In the latter case, the modularity could aim to rescue a compromised function. Although we have no evidence yet to support these hypotheses, we believe that ancestral enzyme reconstruction could clarify these fascinating issues and provide further insight into the evolutionary role of enzyme modularity.

**Figure 5. Speculative minimal evolutionary scenarios leading to the emergence of complex animal Dus2 modularity and a new tRNA binding mode among Dus enzymes. In this model, we propose that the common ancestor of both fungi and animal may have likely carried both the TBD and HD. The first pathway shows that the acquisition of a dsRBD occurred before the loss of positive charges in the HD, leading to a putative intermediate 1 carrying too much affinity for tRNA. Then, this intermediate evolved to a more functional state by converting HD into a less electropositive domain. Conversely, the second pathway proposes that the loss of positive charges occurred first and then acquisition of a dsRBD, which restored a better functionality.**

**ASSOCIATED CONTENT**

* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.8b00584.

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