

Oxidation alters the architecture of the phenylalanyl-tRNA synthetase editing domain to confer hyperaccuracy

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ABSTRACT

High fidelity during protein synthesis is accomplished by aminoacyl-tRNA synthetases (aaRSs). These enzymes ligate an amino acid to a cognate tRNA and have proofreading and editing capabilities that ensure high fidelity. Phenylalanyl-tRNA synthetase (PheRS) preferentially ligates a phenylalanine to a tRNA^{Phe} over the chemically similar tyrosine, which differs from phenylalanine by a single hydroxyl group. In bacteria that undergo exposure to oxidative stress such as *Salmonella enterica* serovar Typhimurium, tyrosine isomer levels increase due to phenylalanine oxidation. Several residues are oxidized in PheRS and contribute to hyperactive editing, including against mischarged Tyr-tRNA^{Phe}, despite these oxidized residues not being directly implicated in PheRS activity. Here, we solve a 3.6 Å cryo-electron microscopy structure of oxidized *S. Typhimurium* PheRS. We find that oxidation results in widespread structural rearrangements in the β -subunit editing domain and enlargement of its editing domain. Oxidization also enlarges the phenylalanyl-adenylate binding pocket but to a lesser extent. Together, these changes likely explain why oxidation leads to hyperaccurate editing and decreased misincorporation of tyrosine. Taken together, these results help increase our understanding of the survival of *S. Typhimurium* during human infection.

INTRODUCTION

Accurate protein synthesis is essential for cellular survival and one important family of enzymes involved in this process are the aminoacyl-tRNA synthetases (aaRS). These enzymes add an amino acid to cognate tRNAs at the 3' terminal adenosine (A76) (1). The addition or charging of tRNAs is extremely accurate ($\sim 10^{-4}$) to ensure that correct amino acids are incorporated into proteins in a codon-specific manner (2). To enable this high fidelity, aaRSs discriminate between chemically-similar amino acids and have the ability to proofread and edit the incorrectly added amino acid (3). This proofreading capability allows for accurate translation to proceed.

aaRSs facilitate a two-step reaction to ligate an amino acid on a tRNA (4). First, the aaRS charges an amino acid with an adenylate group from ATP. Second, the amino-adenylate is ligated onto a tRNA. There are two structurally unrelated classes of aaRSs that differ in the location of the ligated aminoacyl group and in their editing functions (5,6). Class I synthetases aminoacylate the 2'-OH of the terminal A76 of a cognate tRNA (7,8), while Class II aaRSs generally aminoacylate tRNA at the 3'-OH of A76 (9). Class I aaRSs contain highly conserved editing domains and perform both pre-transfer editing, in which the misactivated amino acid-adenylate is hydrolyzed, and post-transfer editing, where the mischarged aminoacyl-tRNA is hydrolyzed (10–16). In contrast, class II aaRSs contain structurally diverse editing sites (17–19) and primarily rely on post-transfer editing. The diversity in class II aaRSs impart differences in their response to environmental stressors, such as increased thermal stability (20).

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Bacterial PheRS is a class II aaRS that functions as a heterotetramer, composed of two α subunits and two β subunits, ($\alpha\beta$)₂ (21). An amino acid, ATP, and tRNA bind in the aminoacylation domain of the α -subunits, and ligation occurs to the 2'-OH of tRNA^{Phe}, distinguishing PheRS from other class II aaRSs. The PheRS β -subunits contain the editing domain, where discrimination between Phe and Tyr occurs post-transfer via hydrolysis of misacylated tRNA^{Phe} to prevent misincorporation of Tyr at Phe codons in the growing peptide chain. Tyr misincorporation rates are high (21), but low error rates can be reconciled by post-transfer editing (8,9,22). One aspect of proofreading is the requirement for Tyr-tRNA^{Phe} to translocate ~ 30 Å to the editing domain (22,23). Conserved residue His265 interacts with the aminoacyl group (22), and discrimination between Phe and Tyr occurs by β Glu334 recognition of the Tyr hydroxyl (24). Conserved β Arg244 interacts with the tRNA^{Phe} backbone at C75, while two catalytic waters are adjacent to the aminoacyl group that assist with residues β Asn254 and β Thr354 to hydrolyze the ester bond between Tyr and the 2'-OH of tRNA^{Phe} A76 (24). Residues β Gly318 and β Ala356 are also implicated in proofreading activity, as mutations in these residues prevent Tyr-tRNA^{Phe} accommodation into the PheRS editing site (25).

PheRS proofreading activity is of particular importance when bacteria are subjected to oxidative stress (26,27). During oxidative stress, imbalances of reactive oxygen species damage various cellular components, including both proteins and nucleic acids (28–33). Damages include oxidation of free amino acids that alters endogenous amino acid pools whose limitation affects translational fidelity (29). Specifically, oxidative stress increases concentrations of *ortho*, *para*, and *meta*-Tyr isomers due to Phe oxidation, challenging PheRS proofreading by increasing the likelihood of tRNA^{Phe} misacylation, and subsequent mistranslation (29).

Salmonella enterica serovar Typhimurium is a pathogen that infects the gastrointestinal (GI) tracts of humans and survives exposure to oxidative stress both in the GI tract and in the lysosomes of macrophages (27,34). Previous studies show that oxidized PheRS in *Salmonella enterica* serovar Typhimurium is hyperaccurate during oxidative conditions, increasing translational fidelity (25). Upon hydrogen peroxide (H₂O₂) treatment, PheRS displays increased proofreading activity and lower misacylation rates without changes in canonical aminoacylation activity, protecting cells from mistranslation under adverse growth conditions. Several residues located throughout PheRS become oxidized resulting in changes to secondary structural features (25). Here, we solved a 3.6 Å cryogenic-EM (cryo-EM) structure of the oxidized *S. Typhimurium* editing-deficient PheRS β G318W to assess structural changes that may contribute to increased proofreading. We report that PheRS oxidation induces conformational rearrangements of both the α and β subunits providing a rationale for hyperaccuracy during oxidative stress.

MATERIALS AND METHODS

PheRS expression and purification

Salmonella enterica serovar Typhimurium PheS and PheT β G318W proteins, encoding the α - and β -subunit, respec-

tively, were purified as previously described (25). Briefly, PheRS was affinity purified (TALON), fractions containing protein were pooled, concentrated and dialyzed overnight in sample buffer (50 mM Tris pH 7.5, 100 mM KCl, 5 mM MgCl₂, 3 mM 2-mercaptoethanol, 5% glycerol). For storage at -80°C , PheRS was dialyzed 4 h in sample buffer containing 50% glycerol.

Cryo-EM sample preparation

Purified PheRS was dialyzed at 4°C for 4 h in a sample buffer containing 5 mM H₂O₂. Three μL of oxidized PheRS at 1.5 mg/ml was incubated for 10 s at 4°C on freshly glow-discharged grids (1.2/1.3 300 mesh Cu Quantifoil). Grids were blotted for 3 s at 100% humidity, plunged into liquid ethane with a Vitrobot Mark IV (FEI), and stored in liquid nitrogen.

Electron microscopy, image processing, and data analysis

A 1330 micrograph dataset was acquired using a Talos Arctica transmission electron microscope (ThermoFisher) operating at 200 keV with BioQuantum/K2 Summit direct electron detector (Gatan). Micrographs were collected at a pixel size of 1.04 Å/pixel at a defocus range of -0.5 to -3.5 μm . A total dose of 64.67 e/Å² per micrograph was fractioned into 48-frame movies with 12 s exposure time.

Image processing was conducted in Relion 3.0 (35). Motion correction and dose weighting was done using MotionCorr2 (36), and contrast transfer function parameters were estimated using Gctf (37). Laplacian of Gaussian algorithm was used to pick particles and generate templates for 2D-referenced based autopicking. Several rounds of 2D-reference based autopicking were conducted, with incorrectly selected particles discarded after reference-free 2D class averaging. An initial three-dimensional refinement with C2 symmetry was conducted using a previously published PheRS structure (PDB code 4P71 (38)) low-pass filtered to 45-Å in Chimera as a reference map (39) (Supplementary Figure S1). Three-dimensional refinement was followed by a 3D classification step, with poorly aligned particles discarded. A total of 204 902 particles were finally selected and subjected to particle polishing, 3D refinement, CTF refinement, and post-processing to yield a final reconstruction of 3.6 Å. Resolution estimations were calculated from Fourier shell correlations (FSC) at 0.143 between the two independently refined half-maps. Local resolution was calculated in Relion. The final map was refined and validated in PHENIX (40) and modeled with PDB code 3PCO (14). Coot (41) and UCSF ChimeraX (42) were used for model building.

Maps were analyzed using UCSF Chimera (39) and ChimeraX (42). Global structural alignments and RMSD calculations were conducted in ChimeraX, using the matchmaker tool which creates a global sequence alignment then fits aligned residue pairs (43). Distance calculations were all conducted in ChimeraX, from the C α carbon of a residue. Figures were created using UCSF Chimera and ChimeraX.

RESULTS

Global structural impact of oxidation on PheRS β G318W

Treatment of *S. Typhimurium* wild-type (WT) and editing-deficient PheRS β G318W with H_2O_2 results in different residues becoming oxidized (25). The PheRS β G318W variant has a 7.5-fold reduction in proofreading due to the prevention of Tyr-tRNA^{Phe} binding in the editing site, likely by steric hindrance (24). Both *S. Typhimurium* oxidized WT and PheRS β G318W show increased proofreading capacity, with PheRS β G318W displaying the most pronounced effect with a >3-fold increase in Tyr-tRNA^{Phe} proofreading (25). Several PheRS β G318W residues were oxidized across both the α - and β -subunits but outside of catalytic regions, and thus it is unclear how increased proofreading results. To define the structural changes that PheRS undergoes during oxidative stress, we solved a 3.6-Å resolution structure of oxidized PheRS β G318W by single particle cryo-EM (Figure 1A; Supplementary Table S1, Supplementary Figure S1). PheRS adopts a heterotetrametric conformation composed of two α -subunits and two β -subunits ($\alpha\beta$)₂ with the α -subunit containing the aminoacylation catalytic domain (residues 108–307) and the β -subunit containing the editing domain (residues 216–354) (14,44–47) (Figure 1C and D). Due to the resolution, it is not possible to resolve the oxidation of specific residues (Figure 1B), but mass spectrometry revealed extensive oxidation under treatment conditions (25).

PheRS undergoes a conformational change upon tRNA^{Phe} binding (44,45), including the ordering of the N-terminal coiled-coil domain of the α -subunit (residues 1–88) (Figure 1C) (48). This region is not resolved in our structure likely because this structure is in the absence of tRNA^{Phe} (Figure 1A). *S. Typhimurium* PheRS is similar to the previously published *Pseudomonas aeruginosa* apo-PheRS structure (apo-*Pa*PheRS) (PDB code 4P71; root mean square deviation (RMSD) of ~2.9 Å comparing the C α of 790 residues) (38). *S. Typhimurium* PheRS and *Pa* PheRS share high sequence identity between both subunits (51.5% for the β -subunits and 65.7% for the α -subunits) and adopt identical tertiary folds (Supplementary Figures S2A and S3). For these reasons, and because our structure is also apo, all future comparisons will be made to apo-*Pa*PheRS unless otherwise noted.

Oxidation induces conformational changes in the PheRS β G318W catalytic domains

Different residues are oxidized in wild-type versus the PheRS β G318W editing-deficient variant, likely due to differences in the overall architecture (25). Six residues become oxidized in PheRS β G318W, which are located across the enzyme domains, with five residues in the β -editing domain and one residue in the α -aminoacylation domain (25) (Figure 1B; Supplementary Table S2). Comparison of PheRS β G318W with apo-*Pa*PheRS reveals that oxidized β Met280 and β Lys282 undergo large changes in the position of their backbone (6.7 and 8.6 Å, respectively) while their side chain conformation is minimally impacted (Figure 2B and C). These residues are in the β -catalytic domain (Figure 2A) located adjacent to the editing site (Figure 1D). The en-

tire β -strand surrounding the editing domain moves in towards the tRNA binding site in the β -subunit, likely contributing to the shift and conformational changes to editing site residues. Other residues within the β -subunit become oxidized yet this has little to no impact on their conformations. For example, ox- β Lys184, ox- β Tyr60 and ox- α Asn222 all have limited movement (Supplementary Figure S4), so it is unlikely that these rotamer changes play a significant role in the hyperaccuracy phenotype. In contrast, ox- β Arg111 has a ~180° rotation compared to *Ec*PheRS complexed with Phe-AMP (Supplementary Figure S4F), and a 16.6 Å movement inwards towards the editing domain compared to apo-*Pa*PheRS (Supplementary Figure S4E). However, since β Arg111 is on the surface of the β -subunit, this difference may be a result from how apo-*Pa*PheRS packs in the crystal lattice (Supplementary Figure S5). Together, these data demonstrate that oxidation has a differential effect on local conformation of the PheRS residues, and oxidation of a given residue does not necessarily entail conformational change in that region.

PheRS oxidation causes structural rearrangements of its editing domain

After PheRS misacylates tRNA^{Phe} with a Tyr, Tyr-tRNA^{Phe} translocates to the editing domain in the β -subunit (Figure 3A) (22,24). In the editing domain, many residues surround Tyr to recognize and edit including β Arg244, β Asn254, β His265, β Glu334 and β Thr354 (Figure 3B). The tRNA is typically stabilized by interactions of β Arg244 with nucleotide C75, while β Asn254 and β Thr354 are in proximity to the aminoacyl ester bond and involved in substrate binding and hydrogen bonding with catalytic waters (Figure 3B and C) (24). These residues were not demonstrated to be oxidized yet they display altered conformations (Figure 3D). β Arg244, β Asn254 and β Thr354 all shift away from the tRNA binding location at distances between ~5.4 and 7.2 Å (Figure 3D; Supplementary Figure S6A). Notably, the β Arg244 flips ~180° towards the editing site (Figure 3D).

The changes in residues located in the PheRS editing domain alter how these residues interact with Tyr and the tRNA backbone of C75. β Glu334 plays an important role in tyrosine discrimination via the formation of hydrogen bonds with the hydroxyl group of Tyr as assessed by molecular docking studies (24). In our structure, β Glu334 shifts away from the editing site ~5.3 Å (Figure 3D). The β His265 stabilizes substrate binding through π -stacking interactions with the Tyr (Figure 3C) (23,24). Other well-conserved editing site residues form an antiparallel β -sheet that upon oxidation, appears to be disrupted (Figure 4A and B) (24). None of the editing site residues are oxidized but this global oxidation disrupts the hydrogen bonding network of the β -sheet, shifting the β -sheet away from the editing site by ~4.5 Å (Figure 4C; Supplementary Figure S6B) (38). This movement creates a larger binding surface (increases from ~204 Å² to ~295 Å²) (42) for the incorrect amino acid to be edited (Figure 4D and E). Despite the relatively moderate changes in the locations of the oxidized residues (Figure 2; Supplementary Figure S4), there is extensive rearrangement of β -

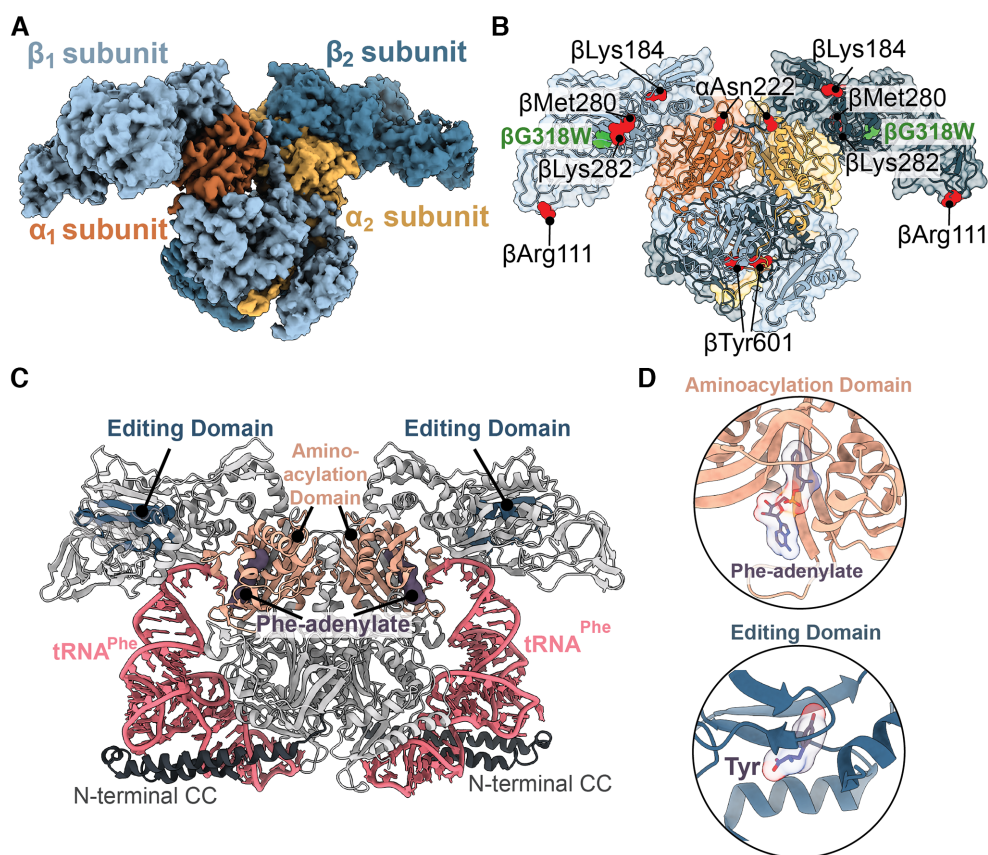


Figure 1. Overview of the PheRS β G318W structure. (A) The 3.6 Å map of the oxidized *S. Typhimurium* editing-deficient PheRS β G318W tetramer. PheRS is composed of two α -subunits (orange and yellow) and two β -subunits (light blue and dark blue). (B) Oxidized PheRS β G318W (green) with oxidized residues, as determined previously (25), shown as red spheres. (C) *T. thermophilus* PheRS (grey) containing Phe-adenylate (purple, surface) and tRNA^{Phe} (pink) (PDB code 2IY5). The N-terminal coiled-coil extensions (N-terminal CC (dark grey); residues 1–88) in the α -subunits are resolved when bound to tRNA. (D) Aminoacylation (peach) and editing domains (dark blue) are shown with respective ligands, Phe-adenylate and Tyr, bound (purple, PDB code 2AMC).

subunit residues involved with editing activity that result in a larger editing domain.

PheRS oxidation induces structural changes in the α -subunit active site

The catalytic domain of the PheRS α -subunits form interactions with Phe and AMP, allowing for the specific addition of a phenylalanyl-adenylate to tRNA^{Phe} (Figure 5A and B; Supplementary Figure S7A). The active site is composed of three regions: the motif 2 loop (residues 194–205), the second helical loop (h2, residues 145–158), and the phenylalanine-proline-phenylalanine (FPF) loop (residues 246–254) (Figure 5B–E; Supplementary Figure S7) (14,44,45). The only α -subunit residue that is oxidized is α Asn222, which is located outside the active site but its backbone conformation is not affected by oxidation (25) (Supplementary Figure S2B). However, global oxidation results in conformational differences within the phenylalanyl-adenylate binding pocket (Figure 5C–E).

In the motif 2 loop, α His202, which normally forms hydrogen bonds with AMP, moves ~ 2.1 Å away from the Phe-adenylate (Figure 5C; Supplementary Figure S7B). However, not all catalytic residues in the motif 2 loop are af-

fected. For example, α Phe206, which forms π -stacking interactions with the adenosine, adopts the same conformation as non-oxidized PheRS (Figure 5C). The h2 and FPF loop of the phenylalanyl-adenylate binding pocket also undergo conformational shifts (Figure 5). The second helical loop (h2), a segment that undergoes rearrangement upon Phe and AMP binding (14,45), undergoes conformational changes upon oxidation despite that none of the h2 residues are oxidized (Figure 5D; Supplementary Figure S7C). Both α His148 and α His149 shift away from the active site ~ 3.7 and ~ 3.8 Å, respectively (Figure 5D). Normally, tRNA^{Phe}, and specifically the last nucleotide A76, is recognized by contacts from the side chain of α Asp154 located within h2 (Figure 5D; Supplementary Figure S8). In oxidized PheRS β G318W, α Asp154 shifts ~ 4.5 Å away from the tRNA binding site. The FPF loop, which is conserved across all bacterial species (14), forms π -stacking interactions with Phe (Supplementary Figure S8). Within the FPF loop, α Phe248 shifts ~ 2.7 Å away from where Phe binds, which would prevent this interaction (Figure 5E; Supplementary Figure S7D). These movements in the motif 2, h2 and FPF loops enlarge the phenylalanyl-adenylate binding pocket from ~ 1560 to ~ 1980 Å² (Figure 5F and G).

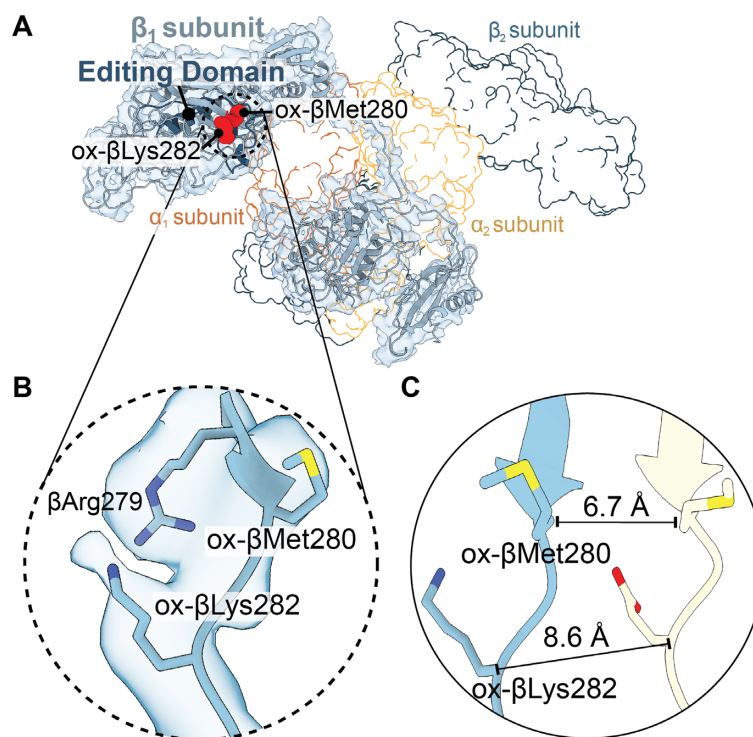


Figure 2. PheRS β Met280 and β Lys282 residues located outside the editing domain move substantially upon oxidation. (A) Overview of the β_1 subunit map and model (light blue) with the editing domain (dark blue) and oxidized residues (red spheres). The other PheRS subunits are shown as outlines. (B) Map and model of ox- β Met280 and ox- β Lys282 located in the editing domain. (C) Comparison of ox- β Met280 and ox- β Lys282 (corresponding residue in *P. aeruginosa* is Glu281) with *P. aeruginosa* apo-PheRS (yellow, PDB code 4P73) revealing significant movements of ~ 6.7 and ~ 8.6 Å, respectively.

Upon oxidation, there is a 30% decrease in misacylation of tRNA^{Phe} with Tyr (25). Discrimination between Phe and Tyr is facilitated by α Glu210 within h2 which hydrogen bonds to the bound amino acid to facilitate ligation (Figure 5B and D; Supplementary Figure S8). In our structure, the position of α Glu210 moves to a position that would directly overlap with the hydroxyl group of Tyr (14) (Supplementary Figure S9) as compared to a *Escherichia coli* PheRS structure containing Phe and AMP (PDB code 3PCO). This new α Glu210 position may explain why misacylation decreases upon PheRS oxidation.

Similarities between oxidized *S. Typhimurium* PheRS and the *E. coli* PheRS–Phe–AMP complex

Oxidation causes both larger editing and aminoacylation sites as compared to apo-*Pa*PheRS as previously described (an increase of 204–295 Å² (Figure 4D and E) and an increase of 1560–1900 Å² (Figure 5F and G), respectively). Although all comparisons to this point have been of oxidized apo-PheRS β G318W to apo-*Pa*PheRS, interestingly, specific regions of oxidized apo-PheRS are slightly more similar to the structure of *Ec*PheRS complexed with ligands Phe and AMP (PDB code 3PCO (14), RMSD of ~ 1.7 Å). While binding of Phe and AMP to *Ec*PheRS increases the editing (~ 272 Å²) and Phe-adenylate binding pocket (~ 1700 Å²), these sites are still smaller than when PheRS is oxidized (Supplementary Figures S10D and S11B).

One of the most striking changes between oxidized apo-PheRS and *Ec*PheRS bound to Phe and AMP is the place-

ment of β Met280 and β Lys282, residues that both become oxidized but are located outside the two catalytic sites (Figure 2). Normally the backbone of these residues dramatically changes upon binding of Phe and AMP (changes of 9.4 Å (β Met280) and 10.3 Å (β Lys282)). Oxidation of β Met280 and β Lys282 also influences their positions however, now these residues adopt similar positions as when Phe and AMP bind (Supplementary Figure S10A) (14). This movement places these residues closer towards the editing domains where tRNA binds despite not containing any ligands.

Comparison of Phe-AMP bound *Ec*PheRS with oxidized apo-PheRS also reveals changes of editing site residues that appear to enlarge the binding pocket and change how oxidized PheRS would engage either the amino acid or AMP (Supplementary Figure S10). First, the side chain of β His265 flips away from the amino acid binding site (Supplementary Figure S10B), likely influencing recognition of the incorrect Tyr. This flipping may occur as PheRS transitions to prepare for hydrolysis of incorrectly bound Tyr. The β Arg244 sidechain, which normally recognizes the tRNA backbone of nucleotide C75, is rotated $\sim 180^\circ$ away in the absence of tRNA (Figure 3C). Oxidation of PheRS causes β Arg244 to more resemble *Ec*PheRS-Phe-AMP despite lacking the tRNA (Supplementary Figure S10B). Lastly, oxidation causes the entire β -sheet comprising the editing domain to shift by ~ 2.7 – 3.7 Å and rotate $\sim 45^\circ$ thus placing it further away from the aminoacylated-tRNA binding site (Supplementary Figure S10C). Collectively, these changes create a larger editing site but with ac-

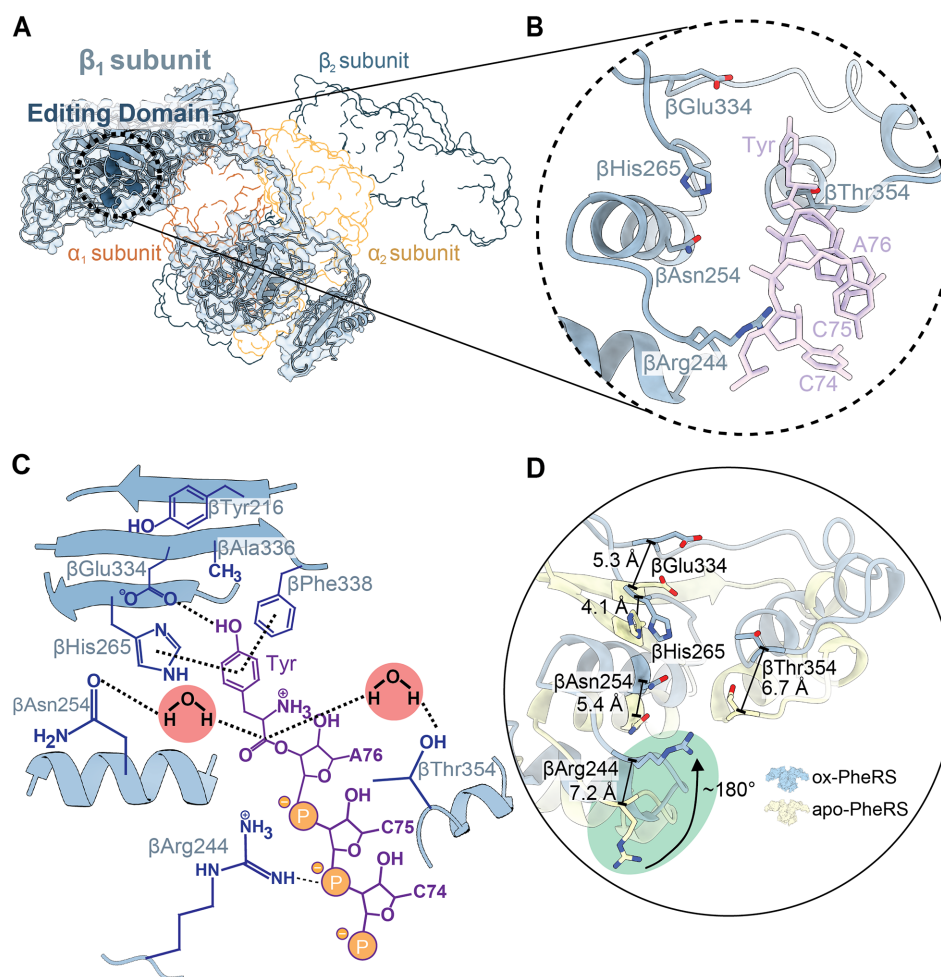


Figure 3. Localized active site conformational changes of the PheRS editing domain. (A) Oxidized *S. Typhimurium* editing-deficient PheRS β G318W β_1 -subunit with corresponding map (light blue) and active site residues in the editing domain shown as spheres (dark blue). The other PheRS subunits are shown in outline. (B) Closeup of editing site residues with Tyr-tRNA^{Phe} docked into the editing site (PDB code 3HFZ, purple). (C) Overview of the editing site residues (blue) and interactions with a Tyr-tRNA^{Phe} CCA end (purple), and the two catalytic waters that carry out the reaction. (D) Residues of the editing domain of the oxidized *S. Typhimurium* PheRS compared to the *P. aeruginosa* apo-PheRS (yellow, PDB code 4P73) undergo a significant shift, particularly β Arg244 which is shifted ~ 7.2 Å and flipped $\sim 180^\circ$ inwards towards the editing site (green circle).

tive site residues positioned in conformations to allow for the increase in editing activity of the oxidized PheRS.

DISCUSSION

To prevent protein mistranslation, aminoacyl-tRNA synthetases pair correct amino acids to their cognate tRNAs with high fidelity to ensure the correct addition of amino acids into proteins. Environmental stress, including oxidative stress, alters cellular amino acid pools and this change can affect the fidelity of gene expression. During oxidative stress, oxidation of Phe converts it to Tyr, increasing Tyr isomer levels and leads to an overall increase in tRNA^{Phe} misacylation (29). However, oxidation of PheRS compensates by increasing proofreading to reduce misacylated Tyr-tRNA^{Phe} from participating in protein synthesis. Several PheRS residues are oxidized and these modifications are found throughout PheRS and not localized solely at the active site (29). Since oxidation causes a hyperaccurate phenotype but there is no oxidation at the active sites, this sug-

gests that allosteric changes may contribute to PheRS acquiring hyperaccuracy consistent with previous results that demonstrated global changes (25). Based upon these results, we set out to determine the cryo-EM structure of oxidized PheRS. Overall, we find that oxidation results in structural changes mainly in the β -subunit editing domain and to a lesser extent, the α -subunit aminoacylation active site. The β -subunit editing domain is enlarged where mis-charged Tyr-tRNA^{Phe} would bind and its surface area increases by 50%. Oxidization also enlarges the phenylalanyl-adenylate binding pocket in the α -subunit by 30%. These enlargements are predicted to contribute to faster and efficient binding, while additional conformational changes of catalytic residues suggest an improvement in Tyr hydrolysis required to ligate the correct Phe. These results shed light on how *S. Typhimurium* is able to survive and grow in adverse environments.

Based on high structural homology between bacterial PheRS (49,50), it is likely that most bacterial PheRS enzymes have a similar hyperaccuracy phenotype under ox-

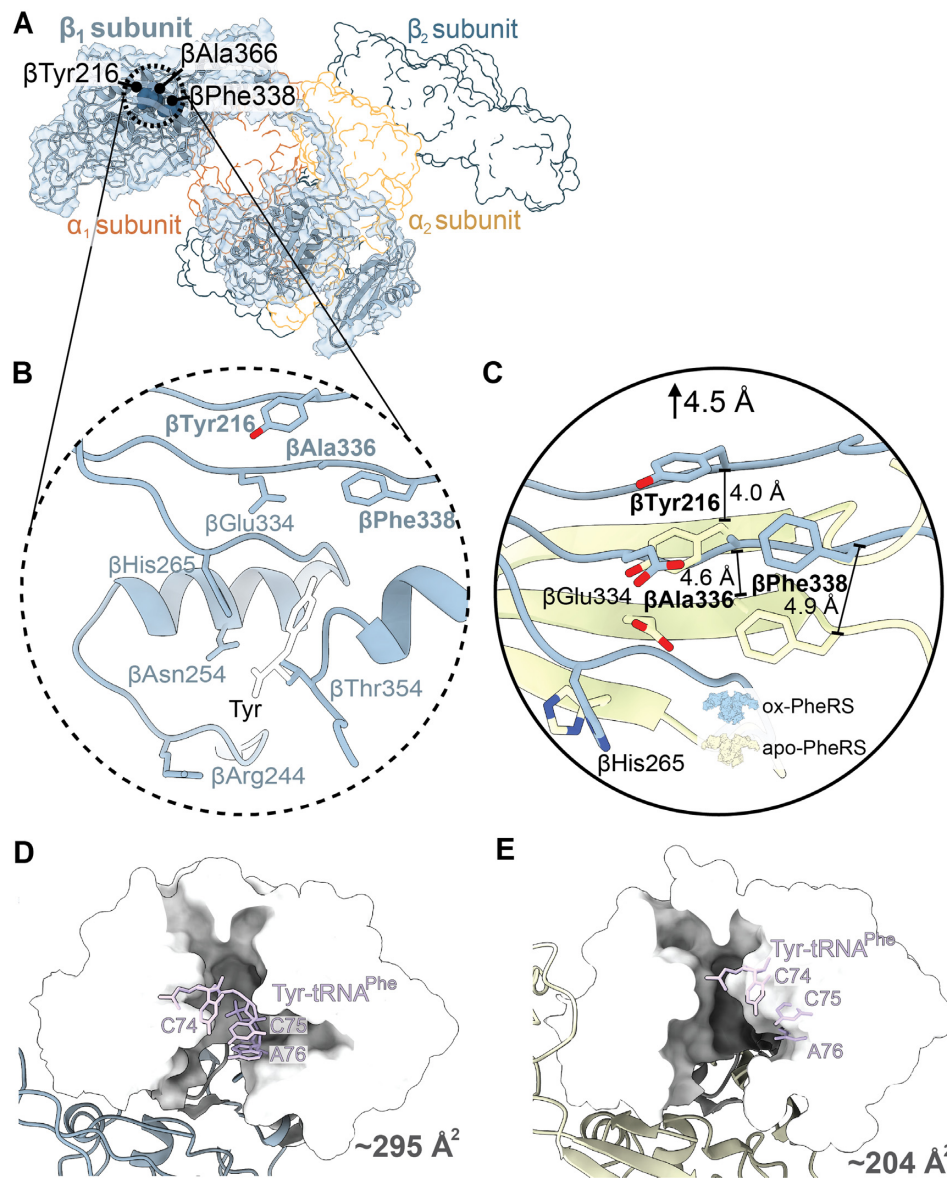


Figure 4. Oxidation of PheRS β G318W opens the editing domain leading to hyperaccuracy. (A) Structure of oxidized *S. Typhimurium* PheRS β G318W β_1 -subunit with corresponding map density (blue) and catalytic residues that are implicated in editing activity shown as dark blue spheres, with other subunits shown in outline. (B) Closeup of the catalytic region with Tyr overlaid from a *T. thermophilus* PheRS-Tyr structure (white, PDB code 3HFZ). (C) The catalytic residues β Tyr216, β Ala336, and β Phe388 of the oxidized *S. Typhimurium* PheRS undergo a shift averaging ~ 4.5 Å away from the editing domain compared to the *P. aeruginosa* apo-PheRS (PDB code 4P73, yellow). (D) Editing domain of oxidized PheRS β G318W shown as surface (white) with Tyr-tRNA^{Phe} docked in site (PDB code 3HFZ, purple). (E) Editing site residues of apo-PheRS shown as surface (white) with Tyr-tRNA^{Phe} docked into the editing site (PDB code 3HFZ, purple).

idative stress. Although there is typically high homology among mitochondrial and bacterial synthetases (51), PheRS is an exception as the mitochondrial and bacterial synthetase are divergent (52). Mitochondrial PheRS is a monomer and lacks an editing domain found in the tetrameric bacterial PheRS (52), and as a consequence, is unable to proofread misacylated tRNA^{Phe} (52). Although mitochondrial PheRS is oxidized during oxidative stress, this does not result in hyperaccuracy and oxidation can be reversed (25), restoring normal function.

While oxidation of other synthetases including ThrRS and AlaRS can occur, this does not result in hyperaccuracy, in contrast to PheRS. ThrRS is known to misacylate

tRNA^{Thr} with Ser and contains a catalytic cysteine in the editing domain (53). Under oxidative conditions, this reactive cysteine is oxidized and thus prevents ThrRS editing ability (54), the opposite phenotype to what is observed with PheRS (25,53). AlaRS also relies on a critical cysteine residue to carry out editing activity upon misacylation of a Ser- or Gly-tRNA^{Ala} (55), but changes in AlaRS editing activity upon oxidation have not yet been reported. Changes in AlaRS editing activity are significantly more determinantal for bacteria viability compared to ThrRS (56). From a fitness perspective, maintenance of AlaRS editing activity is more important for bacterial survival, suggesting possible divergence in editing response under oxidation as com-

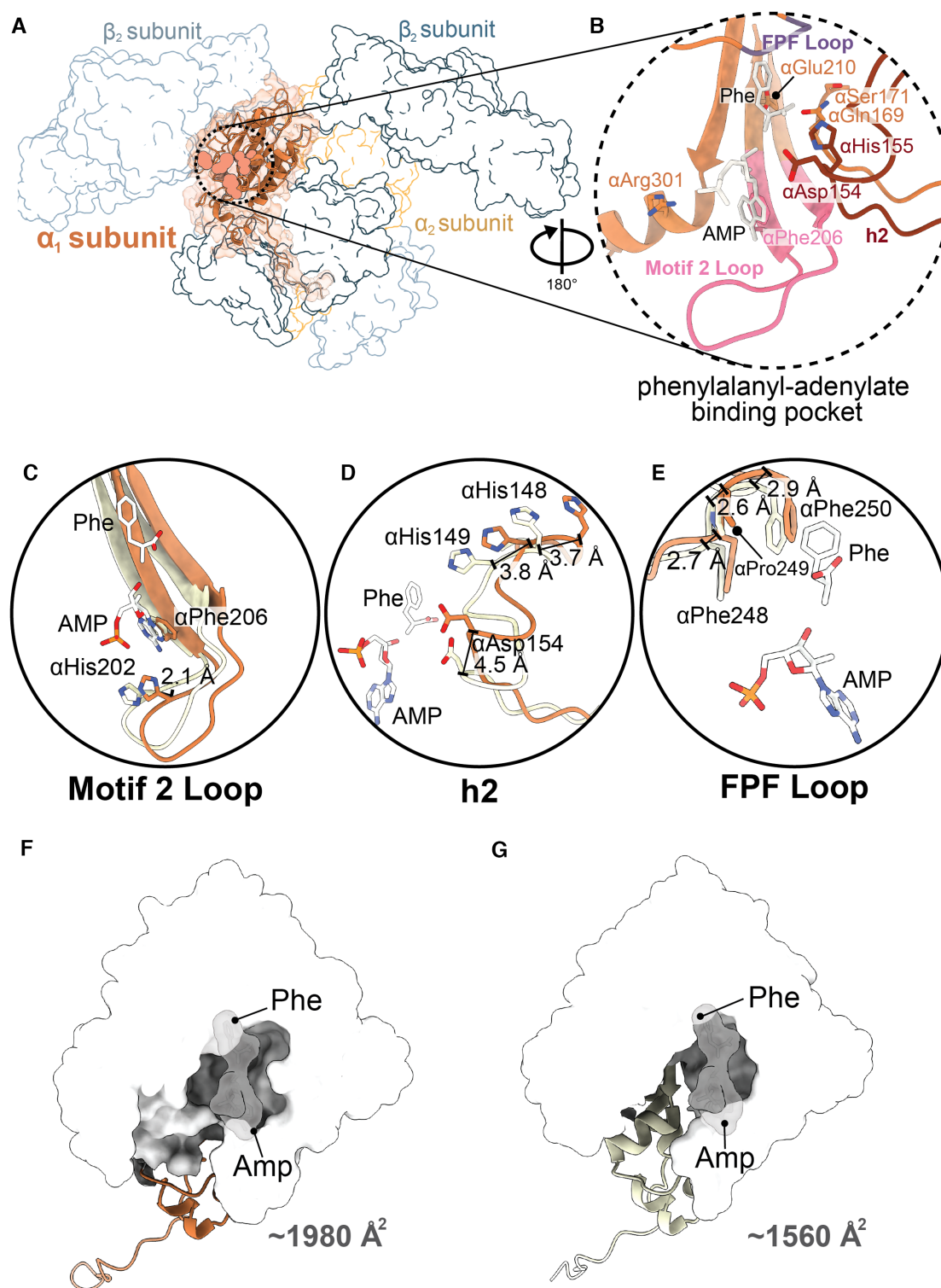


Figure 5. Conformational changes of the aminoacylation domain. (A) Structure of oxidized *S. Typhimurium* PheRS βG318W α₁-subunit and map (orange). Aminoacylation active site residues shown as spheres, with other subunits shown in outline. (B) Active site residues in the phenylalanyl-adenylate binding pocket (orange) with modeled AMP and Phe (PDB code 3PCO, white). The motif 2 loop (pink), helix 2 loop (h2, dark red), and FPF loop (purple) are shown. (C) The motif 2 loop shifts outwards by ~2.1 Å from *P. aeruginosa* apo-PheRS (PDB code 4P73, yellow). (D) The h2 of oxidized PheRS has conformational rearrangements, 3.7 Å and 3.8 Å at αHis148 and αHis149, respectively, and 4.5 Å at αAsp154, away from the phenylalanyl-adenylate binding pocket compared to apo-PheRS (PDB code 4P73, yellow). (E) The FPF loop contains minor conformational changes, with 2.7, 2.6 and 2.9 Å shifts of the αPhe248, αPro249 and αPhe250, respectively, away from the phenylalanyl-adenylate binding pocket compared to apo-PheRS (PDB code 4P73, yellow). (F) Enlargement of the aminoacylation domain shown with surface (white) of oxidized PheRS βG318W, with AMP and Phe groups docked in and shown with surfaces (PDB code 3PCO, grey). (G) Enlargement of the aminoacylation domain shown as surface (white) of the apo-PaPheRS (PDB code 4P73, yellow), with AMP and Phe groups docked in shown as surfaces (PDB code 3PCO, grey).

pared to ThrRS. Under normal growth conditions, PheRS editing activity is not vital for bacterial viability, but becomes required for growth under oxidative conditions (29). Translation accuracy under oxidative stress is vital for cellular survival, but among the class II aaRSs, only PheRS undergoes a gain of function change. The benefit of PheRS hyperaccuracy under conditions that directly increase the likelihood of mistranslation serves as a direct compensatory mechanism to decrease the likelihood of Tyr misincorporation; while ThrRS and possibly AlaRS are susceptible to oxidative stress, the Ser or Gly amino acid pools are not expected to change significantly, placing less selective pressure on these synthetases to compensate for misacylation under oxidative conditions.

Oxidation is a well characterized signal for proteins directly involved in the oxidative stress response including redox-operated switches that activate transcription of oxidative response elements (57). However, these mechanisms are indirect and contribute to a regulatory pathway that allows for increased cell survival under adverse conditions. PheRS represents one of the few examples of a functional protein that self-regulates as a feedforward mechanism to directly compensate against an adverse condition.

Responses to stress are typically analyzed at the level of gene expression (58,59). With oxidative stress, several genes including *oxyR*, *oxyS*, or *perR*, are directly regulated by oxidation of several cysteine residues allowing for transcription factor binding to promote the expression of proteins involved in metabolism or clearance of oxidative stress (60). The hyperaccuracy phenotype of PheRS under oxidative stress illustrates how important pathways in protein synthesis work to maintain proper gene expression. This ensures successful expression of oxidative response genes, which relies on accurate amino acid additions to growing nascent chain on the ribosome. Oxidized PheRS hyperaccuracy ensures that correct gene expression is not compromised by elevated and potentially toxic Tyr isomer misincorporation under oxidative stress. Hyperaccurate aminoacylation thus helps to increase survival, particularly for organisms regularly present in oxidative environments like *S. Typhimurium*, which can thrive in environments like a macrophage despite a host reactive oxygen species attack. Overall, the proper activation of stress responses, including increased proofreading activity of PheRS, increases *S. Typhimurium* survival and contributes to the overall fitness of the bacterium.

DATA AVAILABILITY

Atomic coordinates and maps for the reported cryo-EM structures have been deposited with the Protein Data bank under accession number 7N8Y and Electron Microscopy Data bank under accession number EMD-24249.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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