Acetylation promotes TyrRS nuclear translocation to prevent oxidative damage

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Tyrosyl-tRNA synthetase (TyrRS) is well known for its essential aminoacylation function in protein synthesis. Recently, TyrRS has been shown to translocate to the nucleus and protect against DNA damage due to oxidative stress. However, the mechanism of TyrRS nuclear localization has not yet been determined. Herein, we report that TyrRS becomes highly acetylated in response to oxidative stress, which promotes nuclear translocation. Moreover, p300/CREB-associated factor (PCAF), an acetyltransferase, and sirtuin 1 (SIRT1), a NAD+-dependent deacetylase, regulate the nuclear localization of TyrRS in an acetylation-dependent manner. Oxidative stress increases the level of PCAF and decreases the level of SIRT1 and deacetylase activity, all of which promote the nuclear translocation of hyperacetylated TyrRS. Furthermore, TyrRS is primarily acetylated on the K244 residue near the nuclear localization signal (NLS), and acetylation inhibits the aminoacylation activity of TyrRS. Molecular dynamics simulations have shown that the silico acetylation of K244 induces conformational changes in TyrRS near the NLS, which may promote the nuclear translocation of acetylated TyrRS. Herein, we show that the acetylated K244 residue of TyrRS protects against DNA damage in mammalian cells and zebrafish by activating DNA repair genes downstream of transcription factor E2F1. Our study reveals a previously unknown mechanism by which acetylation regulates an aminoacyl-tRNA synthetase, thus affecting the repair pathways for damaged DNA.

\textbf{Results}

Nicotinamide and Oxidative Stress Enhance the Acetylation Level of TyrRS. Several recent proteomic studies have shown that lysine residues on TyrRS are often acetylated (10, 12). To confirm that TyrRS was acetylated, human full-length TyrRS was overexpressed in HEK293T cells, which were subsequently treated with nicotinamide (NAM), a common deacetylase inhibitor that is effective in HEK293T cells, which were subsequently treated with nicotinamide (NAM), a common deacetylase inhibitor that is effective. TyrRS was indeed acetylated and that the acetylation level of TyrRS was approximately fourfold higher after treatment with 5 mM NAM for 4–6 h (Fig. L2), indicating that an NAD+-dependent deacetylase of the sirtuin family, most likely cytosolic sirtuin 1 (SIRT1) or SIRT2 (21), was the deacetylase of TyrRS.

The acetylation of TyrRS in the nucleus is a posttranslational modification that occurs in response to oxidative stress. TyrRS, a member of the family of aminoacyl-tRNA synthetases (AARSs), comprises a family of 20 enzymes that catalyze the first step of protein synthesis by activating specific amino acids to the 3′ ends of their cognate tRNAs (1). Recently, AARSs have been implicated in specific physiological responses, such as apoptosis (2), cellular growth (3), and angiogenesis (4). Tyrosyl-tRNA synthetase (TyrRS or YARS) is conserved in eukaryotes ranging from insects to humans, and multiple mutations have been identified in its catalytic domain. These mutations are associated with dominant intermediate Charcot-Marie-Tooth (CMT) disease, a common peripheral nervous system disorder caused by axonal degeneration and demyelination (5).

In response to oxidative damage or serum starvation stress, TyrRS translocates to the nucleus, where it protects against DNA damage by activating the transcription factor E2F1 and subsequent downstream DNA repair genes (6, 7). Notably, resveratrol directly binds to the active site of TyrRS, thereby directing TyrRS to the nucleus, and stimulates the activation of NAD+-dependent auto-poly-ADP ribosylation of poly(ADP ribose) polymerase 1 (PARP1) (6, 7). Interestingly, a hexapeptide motif in the anticondon recognition domain of TyrRS has been identified as a nuclear localization signal (NLS) (8). However, the mechanism by which TyrRS is redistributed between the cytoplasmic and nuclear compartments during stress and the effect of TyrRS localization on its aminoacylation and activity toward DNA damage protection remain unknown.

Protein acetylation, a posttranslational modification process that has been conserved through evolution, has been extensively investigated in recent years (9–12). Acetylation regulates diverse cellular processes, including gene silencing (13), oxidative stress (13, 14), DNA repair (15), cell survival and migration (16, 17), and metabolism (9, 18, 19). Most acetylated proteins act as transcription factors in the nucleus and as metabolic enzymes outside the nucleus (9). Strikingly, the acetylation of multiple aminoacyl-tRNA synthetases, including tyrosyl-tRNA synthetase, has been reported in a number of proteomic studies (10, 12). However, the link between acetylation and AARSs remains to be established.

Herein, we propose a model in which acetylation of the NLS in TyrRS enhances its nuclear transport under oxidative stress.

\textbf{Author contributions:} W.Y. conceived the study; W.Y. designed the experiments; X.C., C.L. and W.Y. analyzed data; and X.C., C.L. and W.Y. wrote the paper.

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Because TyrRS has been reported to be involved in the oxidative stress response (7), its acetylation level was assessed after inducing oxidative stress. Strikingly, upon H₂O₂ treatment, the acetylation level of TyrRS significantly increased in a concentration-dependent manner (Fig. 1B). To further validate this result, cells overexpressing TyrRS were treated with the oxidative stressor menadione, a polycyclic aromatic ketone that generates intracellular reactive oxygen species (ROS) at multiple cellular sites via futile redox cycling (14). Menadione treatment also significantly increased the acetylation level of TyrRS (Fig. 1C), indicating that TyrRS becomes highly acetylated in response to oxidative stress.

**NAM and Oxidative Stress Promote TyrRS Nuclear Localization.** Despite its central role in the translational machinery of the cytoplasm, TyrRS rapidly translocates to the nucleus in response to oxidative stress (7). Having shown that the acetylation levels of TyrRS significantly increased in response to stress (H₂O₂ and NAM), we determined whether acetylation induced nuclear translocation, by treating HEK293T, HeLa, and human osteosarcoma U2OS cells with H₂O₂ and NAM, and assessed the distribution of endogenous TyrRS after treatment by cell fractionation and immunofluorescence microscopy. In all three different types of cells, NAM and H₂O₂ dramatically enhanced the nuclear import of TyrRS (Fig. 1D and E).

![Fig. 1. NAM and oxidative stress promote TyrRS nuclear translocation through acetylation.](image)

**Oxidative Stress-Mediated SIRT1 and PCAF Regulate the Nuclear Localization of TyrRS in an Acetylation-Dependent Manner.** Lysine acetylation is a reversible process catalyzed by acetyltransferases and deacetylases (9). To identify the acetyltransferase responsible for TyrRS acetylation, we overexpressed p300/CBP-associated factor (PCAF), Tip60, and GCN5 into HEK293T cells, because PCAF, Tip60, and GCN5 have been shown to respond to oxidative stress (22–25), and found that the acetylation of TyrRS was elevated only after the ectopic expression of PCAF, not the other acetyltransferases (Fig. 2A). We also detected a specific interaction between TyrRS and PCAF (Fig. 2B). Furthermore, PCAF knockdown using siRNA significantly reduced the acetylation of TyrRS (Fig. 2C). To determine whether PCAF influenced the nuclear localization of TyrRS, the effect of ectopic PCAF expression on TyrRS localization was analyzed. As expected, ectopic PCAF expression promoted the nuclear localization of TyrRS (Fig. 2D). Together, these results demonstrated that PCAF acetylates TyrRS, thus facilitating nuclear translocation.

Next, the deacetylase responsible for TyrRS regulation was investigated. We confirmed that NAM (a sirtuin inhibitor) and oxidative stress treatment significantly increased the acetylation level of TyrRS (Fig. 1A). Previous studies have shown that oxidative stress reduces the protein expression of SIRT1 (26, 27). To determine whether SIRT1 interacted with TyrRS as its deacetylase, we coexpressed TyrRS with either SIRT1 or SIRT2 in HEK293T cells and then measured the acetylation level of TyrRS. We found that the ectopic expression of SIRT1, but not SIRT2, decreased TyrRS acetylation (Fig. 2E). We further verified the SIRT1–TyrRS interaction in HEK293T cells by coimmunoprecipitation. However, SIRT2, a sirtuin deacetylase that is abundant in the cytoplasm, did not interact with TyrRS (Fig. 2F and Fig. S24). Moreover, SIRT1 knockdown via transfection of two siRNAs significantly increased the acetylation level of TyrRS (Fig. 2G). To further confirm that SIRT1 regulated the acetylation and subsequent subcellular relocation of TyrRS, we coexpressed SIRT1 with TyrRS in HEK293T cells and found that ectopic SIRT1 inhibited the nuclear translocation of TyrRS in response to H₂O₂ treatment (Fig. 2H). We also constructed a SIRT1 knockout (KO) HeLa cell line using the iCRISPR method (28) to further verify the regulation of TyrRS by SIRT1 (Fig. S28). Through cell fractionation and immunofluorescence staining experiments, we discovered that the loss of SIRT1 promoted the nuclear translocation of TyrRS (Fig. S2 and C). The knockdown of SIRT1 in HEK293T with two siRNAs further confirmed this result (Fig. S2D). Collectively, these results demonstrated that SIRT1 prevented TyrRS localization to the nucleus via deacetylation.

To understand whether oxidative stress regulated SIRT1 and PCAF to affect TyrRS acetylation levels, we investigated the SIRT1 expression levels and its deacetylation activity using an assay under oxidative stress. Consistent with previous reports, H₂O₂ and menadione treatment decreased SIRT1 protein levels (Fig. 2F and Fig. S2E) and reduced its deacetylation activity (Fig. 2F), resulting in significantly increased TyrRS acetylation (Fig. 2A). Moreover, we found that PCAF levels were elevated under H₂O₂ treatment, enhancing the acetylation of TyrRS. Taken together, these results suggest that oxidative stress-mediated PCAF and SIRT1 regulate TyrRS acetylation and its subsequent translocation to the nucleus.
considered the role of K244, a lysine residue in the NLS motif that is the most conserved lysine residue across different TyrRS proteins, as a putative acetylation site (Fig. 3A). To identify the primary acetylation site of TyrRS, we replaced several lysine residues with glutamine (Q), an acetyl-mimetic amino acid (18), and transfected 293T cells with individual mutants. We found that substitution of the K244 residue significantly decreased the overall acetylation of TyrRS compared with that of the WT protein, suggesting that K244, but not K197 or K206, was the primary acetylation site (Fig. 3B). The acetylation of TyrRS was not completely abolished in the K244 mutant, indicating that other sites may be acetylated in TyrRS.

To test the hypothesis that K244 acetylation promoted TyrRS nuclear translocation in response to oxidative stress, we transfected HEK293T cells with either WT TyrRS or mutant versions (K244Q, acetylated mimic, and K244R, deacetylated mimic) of TyrRS and detected their subcellular localization. The acetylation mimic mutant K244Q showed greater nuclear accumulation than both the WT and K244R mutant, demonstrating the importance of acetylation for this change in localization (Fig. 3C). We also found that the phenotype of the K244Q mutant was identical to that of the H2O2-treated WT, showing that substantial TyrRS accumulation occurred in the nucleus. Moreover, the phenotype of the K244R mutant was identical to that of the untreated WT (Fig. 3C). To further investigate how the acetylation of K244 modulates the nuclear import of TyrRS, we performed molecular dynamics simulations on the deacetylation mimetic mutation K244R and the acetylation mimetic mutation K244Q in human TyrRS (PDB ID code 4BQT). First, acetylation of K244 affected the conformation of TyrRS around the NLS, indicating a positive charge on the lysine side chain. Moreover, the acetylation-induced changes on K244 affected the electrostatic surface potential of the NLS in human TyrRS (Fig. 3E). Taken together, these simulations suggested that acetylation of K244 induced changes in TyrRS that affected its conformation and the electrostatic field around the NLS, which may have promoted TyrRS nuclear translocation. Therefore, we decided to generate an antibody specific to acetylated K244 and verify its specificity to further investigate the function of K244 acetylation. The proposed antibody readily detected K244-acetylated TyrRS (Fig. 3F). Furthermore, Western blotting analysis of a whole-cell extract from HEK293T cells with the anti-acetyl-K244 antibody detected a band with a substantial increase in intensity after treatment with either NAM or H2O2, thereby demonstrating endogenous acetylation of TyrRS at K244 in cultured cells (Fig. 3G).

Given that SIRT1 regulates TyrRS acetylation, we measured the endogenous TyrRS K244 acetylation level in SIRT1-KO iCRISPR HeLa cells and found that K244 acetylation was markedly enhanced (Fig. 3H), indicating that K244 was the major site of TyrRS acetylation and that SIRT1 was responsible for the deacetylation of TyrRS. Moreover, we confirmed that the ectopic expression of PCAF increased K244 acetylation and that the knockdown of PCAF using siRNA significantly reduced K244 acetylation (Fig. S3A and B). These results demonstrated that PCAF was the major acetyltransferase for TyrRS at lysine 244 and that TyrRS K244 was the major acetylation site regulated by SIRT1 and PCAF.

**Acetylation at K244 Promotes TyrRS Nuclear Translocation but Inhibits Enzyme Activity.** To further show that the nuclear translocation of TyrRS was directly regulated by K244 acetylation, we conducted cell fractionation and immunofluorescence experiments and discovered that K244-acetylated TyrRS proteins were highly enriched in the nucleus after NAM and H2O2 treatment (Fig. 3 I and J). In addition to promoting the nuclear translocation of TyrRS (6), resveratrol also enhanced the protein level of PCAF and the acetylation level of K244, thus facilitating the

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**Fig. 2.** Oxidative stress-mediated PCAF and SIRT1 regulate the nuclear localization of TyrRS. (A) PCAF acetylates TyrRS. TyrRS-Flag was coexpressed with different acetyltransferases and purified by Flag beads. **P < 0.01.** (B) Coimmunoprecipitation assay detecting TyrRS-PCAF binding in HEK293 cells transiently transfected with Flag-tagged PCAF or TyrRS. (C) PCAF knockdown decreases TyrRS acetylation. Acetylation levels of TyrRS-Flag expressed and purified from 293T cells with or without PCAF knocked down by siRNA were detected by Western blotting. (D) PCAF overexpression promotes TyrRS translocation. The subcellular localization of TyrRS was examined in human HEK293T cells expressing the indicated plasmids or subjected to the indicated treatments. (E) SIRT1 overexpression decreases TyrRS acetylation. The acetylation levels of TyrRS in HEK293T cells expressing the indicated plasmids and then treated with H2O2 were analyzed by Western blotting. (F) Coimmunoprecipitation assay detecting TyrRS-SIRT1 binding in HEK293 cells transiently transfected with Flag-tagged SIRT1 or TyrRS. (G) SIRT1 knockdown promotes TyrRS acetylation. Acetylation levels of TyrRS-Flag expressed and purified from 293T cells with or without SIRT1 knocked down by siRNA were detected by Western blotting. (H) SIRT1 overexpression prevents oxidative stress-mediated TyrRS translocation. The subcellular localization of TyrRS was examined in human HEK293T cells expressing the indicated plasmids or subjected to the indicated treatments. (I) H2O2 treatments decrease SIRT1 level and increase both PCAF level and TyrRS acetylation. TyrRS-Flag was transfected into HEK293T cells followed by indicated treatments. Average quantified relative protein abundance from all three repeats is shown with SD. **P < 0.05; **P < 0.01; ***P < 0.001. (J) SIRT1 deacetylase activity is significantly decreased under oxidative stress. HEK293T cells transfected with HA-tagged SIRT1 were performed with indicated treatments. Then HA-tagged SIRT1 was immunoprecipitated from cells and used in the in vitro deacetylase assay. The mean value of triplicate experiments and SD are presented. **P < 0.05; **P < 0.01.
K244 acetylation affects TyrRS nuclear translocation under the regulation of PCAF and SIRT1. (A) Alignment of the TyrRS sequence surrounding K244 from various species, including human (Homo sapiens, National Center for Biotechnology Information (NCBI)) reference number NP_001006314.1), zebrafish (Danio rerio, NP_958473.1), chicken (Gallus gallus, NP_001006314.1), and fruit fly (Drosophila melanogaster, NP_648895.1), budding yeast (Saccharomyces cerevisiae, NP_011701.3), and fission yeast (Schizosaccharomyces pombe, NP_587876.1). Bold blue text indicates lysine 244. (B) K244 is the primary acetylation site of TyrRS. The indicated plasmids were transfected into HEK293T cells, and the proteins were immunoprecipitated before being subjected to Western blotting for acetylation analysis. ***p < 0.001. (C) Exogenous K244Q mutant locates in nucleus. Subcellular localization of exogenous TyrRS-Flag and K244 mutations was examined in HEK293T cells with indicated treatments or expression plasmids. (D and E) Molecular modeling of acetylation of K244 in TyrRS. TyrRS K244R caused a hydrogen-bond disruption between K244 and D236 (TyrRS). TyrRS K244R caused a hydrogen-bond disruption between K244 and D236 (TyrRS). The indicated plasmids were transfected into HEK293T cells, and the proteins were immunoprecipitated before being subjected to Western blotting for acetylation analysis. ***p < 0.001. (F) Characterization of acetyl-TyrRS (K244) antibody. The acetylation level of HEK293T cells was detected by Western blotting after treatment with resveratrol at the doses shown. Shown are representative Western blot results of three replicates with SD. **p < 0.01; ***p < 0.001. **p < 0.01; ***p < 0.001.

The Acetylation-Mimic K244Q Mutant TyrRS Protects Against DNA Damage Under Oxidative Stress. Nucleus-localized TyrRS has previously been shown to protect cells from DNA damage under certain conditions (7). We hypothesized that the acetylation mimic K244Q mutant TyrRS would be able to perform the same function by accumulating in the nucleus at the cost of decreasing its enzymatic activity. To test this hypothesis, we transfected HEK293T cells and U2OS cells with WT, K244Q, and K244R TyrRS and detected γ-H2A.X levels by both Western blotting and immunofluorescence after treatment with 200 μM H2O2 for 1 h. The K244Q mutant, but not the K244R mutant, showed significantly less DNA damage under oxidative stress conditions (Fig. 4A and Fig. S4A). To further test these results, we transfected HEK293T cells with WT, K244Q, and K244R TyrRS and used quantitative real-time PCR (qRT-PCR) to measure the expression level of the DNA damage-response gene cluster, which is activated by the transcription factor E2F1. In support of our hypothesis, we found that the K244Q-mutant TyrRS significantly increased the DNA damage repair and sensing functions of BRCA1 and RAD51L1, respectively (30, 31), in the absence of stress (Fig. 4B). In contrast, the phenotypes of the K244R mutant and vector-transfected cells were identical. Specifically, the transcription of DNA repair genes was not up-regulated (Fig. 4C). We next investigated whether mutants of TyrRS would affect homologous recombination (HR) during the DNA damage repair process. Using a reporter assay for HR (32), we found that reconstituting cells with shRNA-resistant K244Q and K244R TyrRS mutants resulted in compromised HR in TyrRS-depleted HEK293T cells compared with that in cells reconstituted with WT or K244Q TyrRS mutants (Fig. 4D and Fig. S4B and C). These results support the idea that K244Q TyrRS has a stronger effect on the promotion of DNA damage repair.

TyrRS is an essential component of the translation machinery (1). However, severe knockdown of TyrRS expression in human cells does not result in clear cytotoxicity (7). To eliminate the possibility that the overexpression of exogenous TyrRS did not mimic authentic intracellular conditions, we stably transfected U2OS cells with TyrRS shRNA (Fig. S4B) and reconstituted the cells with shRNA-resistant K244Q or K244R mutants using a lentivirus (Fig. S4C). We detected the subcellular localization of TyrRS in these cell lines and found that the K244Q-mutant TyrRS accumulated in the nucleus, even in the absence of stress (Fig. 4E). In contrast, the K244Q mutant accumulated in the cytoplasm and did not translocate to the nucleus in response to H2O2 treatment (Fig. 4E), which is consistent with the results shown in Fig. 3. We also found that the K244Q mutant promoted translocation and protected against DNA damage after H2O2 treatment, as detected by γ-H2A.X staining (Fig. 4E). Specifically, γ-H2A.X foci in positive nuclei were counted and quantified, and the counts were normalized to the total number of analyzed nuclei (Fig. 4F). These results further indicated that TyrRS K244 acetylation protects cells against DNA damage due to oxidative stress through TyrRS nuclear translocation.
Acetylation-mimic K244Q mutant TyrRS protects against DNA damage under oxidative stress. (A) K244Q TyrRS protects cells from DNA damage caused by oxidative stress. HEK293T cells were transfected with the indicated plasmids and then subjected to the indicated treatments. Results were detected by Western blotting. (B) Overexpression of K244Q TyrRS in HEK293T cells up-regulates DNA damage response genes relative to the WT, as detected by qRT-PCR. Error bars represent the error for triplicate experiments. *(P < 0.05); **(P < 0.01); ***P < 0.001. (C) Over-expressing K244R TyrRS in HEK293T cells did not activate DNA damage response gene clusters. *P < 0.05. (D) K244R TyrRS mutant resulted in compromised HR. The HR assay is well-described in SI Materials and Methods. Data are presented as mean ± SD of three independent experiments. ###P < 0.001. (E) K244 acetylation promotes TyrRS nuclear translocation, thereby protecting against DNA damage. The subcellular localization of TyrRS was examined in shTyrRS cells and in K244Q and K244R cell lines with indicated treatments. The γ-H2A.X level was also measured. (F) Cells positive for γ-H2A.X foci were defined as cells with one nucleus containing at least 10 foci. γ-H2A.X focus-positive cells were randomly counted from N total cells subjected to the indicated treatments (n = 50 to 100). Error bars represent the error for triplicate experiments. ###P < 0.001; ***P < 0.001. (G) and (H) K244Q protects zebrafish embryos from oxidative stress-mediated DNA damage. The embryos of zebrafish were injected with the indicated mRNA and morpholinos. The γ-H2A.X levels of 28-h postfertilization fish were examined by immunofluorescence microscopy. The images (G) and fluorescence intensities (H) were analyzed and quantified with ImageJ. Error bars represent the error for N independent fish embryos (n = 3 to 6). **P < 0.01. (I) Schematic illustration of how acetylation regulates TyrRS relocalization in response to oxidative stress.

Discussion

TyrRS has been recently shown to translocate to the nucleus, exhibiting protective effects against DNA damage due to oxidative stress (7). A potential mechanism of the aforementioned protective effects was analyzed in the present study. Specifically, we identified factors that promote the nuclear translocation of acetylated TyrRS and established a physiological role for acetylation in the prevention of DNA damage (Fig. 4F). Although acetylation plays an important role in regulating the function of several metabolic enzymes (14, 18, 19), acetylation has been shown to modulate several proteins, such as PKM2 and FOXO1, enhancing their translocation into the nucleus (18, 33). Our study provided further insights into the mechanism of acetylation-regulated TyrRS nuclear localization. TyrRS is a member of the aminoacyl-tRNA synthetase family, whose members are among the most abundant proteins in cells and are known for their essential aminoacylation roles in protein synthesis. In the present study, we discovered that oxidative stress markedly increased the acetylation level of TyrRS, which promoted the nuclear translocation of TyrRS. Moreover, acetylation at K244 in TyrRS regulated both its cytoplasmic aminoacylation activity and nuclear translocation, thereby increasing protection against DNA damage. Indeed, K244 acetylation strongly protected cell cultures and zebrafish against DNA damage by activating DNA repair genes located downstream of E2F1. More importantly, we demonstrated that PCAF and SIRT1 were the acetylation enzymes that acted on TyrRS and regulated its nuclear translocation in response to oxidative stress. Along with previous findings, our data were indicative of a unique mechanism in which acetylation regulates TyrRS, thus protecting cells from DNA damage (Fig. 4F). These findings suggest that targeting the K244 residue of TyrRS may be a therapeutic strategy for physiological diseases characterized by DNA damage.

SIRT1, an NAD+-dependent deacetylase, is the best-studied member of the sirtuin family and has been implicated in signaling pathways in various diseases, including cancer (34), cardiovascular diseases (35), diabetes (36), and neurodegeneration (37). DNA damage due to oxidative stress is an important hallmark of many diseases, and SIRT1 plays several critical roles in the DNA damage response, as previously discussed (15, 38). Notably, PARP1, another protein that responds to DNA damage, is part of an intricate network that includes SIRT1 (16, 38). Specifically, SIRT1 and PARP1 share a common cofactor, NAD+, as well as several common substrates (39). More strikingly, Sajish and Schimmel recently found that serum starvation or resveratrol addition promotes the nuclear translocation of TyrRS, which enhances the interaction of TyrRS with PARP1 and increases the PARP1 activity (6). However, serum starvation increases SIRT1 protein expression, thereby inhibiting the function of PARP1 (16, 40). Thus, the function of this regulatory network remains controversial. Because resveratrol has been shown to promote the localization of TyrRS in the nucleus, we verified herein that resveratrol increased the amount of acetylated K244 of TyrRS in a dose-dependent manner by up-regulating the protein expression.

Finally, we explored the protection conferred by the acetylation-mimic K244Q TyrRS against DNA damage in vivo using the zebrafish as a model organism and the well-established method described by Wei et al. (7). Briefly, 0 to 1 h postfertilization, we injected zebrafish embryos with two TyrRS-targeting antisense morpholinos to suppress the endogenous fish TyrRS and reconstructed the embryos with WT, K244Q, and K244R TyrRS mRNAs to induce the expression of exogenous human TyrRS (Fig. S4E). We then examined the number of γ-H2A.X foci by immunofluorescence after treatment with 20 mM H₂O₂ for 2 h. H₂O₂ treatment significantly increased DNA damage in non-injected, WT-injected, and K244R mutant-injected fish (Fig. 4G and Fig. S4F). In comparison, DNA damage was much less extensive in K244Q mutant-injected fish after H₂O₂ treatment (Fig. 4G and Fig. S4F), suggesting that K244Q TyrRS was strongly protective against DNA damage in vivo. These results suggested that TyrRS-mediated protection was strongly correlated with subcellular localization, which is regulated by acetylation at K244 (Fig. 4F).
levels of PCAF (Fig. 3K). We also found that SIRT1 deacetylated TyrRS and prevented nuclear localization. Interestingly, the acetylation level of TyrRS significantly decreased after 2 h of serum starvation (Fig. S2F), similar to the effects of PARP1 activation (6). Moreover, the mRNA level of SIRT1 increased when the K244Q mutant of TyrRS was overexpressed in cells (Fig. S4D). Collectively, these results suggest that regulation occurred due to negative feedback from TyrRS, PARP1, and SIRT1 under oxidative stress.

In addition to their aminoaacylation functions in protein synthesis, many aminoaacyl-tRNA synthetases, including TyrRS, have been shown to take on multiple roles (41–43). Specifically, TyrRS was found to act as a sensor for oxidative stress after translocating to the nucleus, where it activates DNA damage repair genes that are downstream of E2F1 (7). Our study suggests that acetylation of K244 in the NLS of TyrRS results in the relocalization of this protein to the nucleus and prevents DNA damage by significantly activating DNA repair genes located downstream of E2F1, as observed in our in vivo models. Therefore, the present study not only reveals a regulatory mechanism for the tRNA synthetase TyrRS due to posttranslational modification but also enriches our knowledge of the regulatory pathway between SIRT1 and the repair response to damaged DNA. These findings provide a basis for the development of drugs that target the K244 residue of TyrRS to treat diseases characterized by DNA damage.

Materials and Methods

All experiments were also approved by the Animal Care Committee at the Fudan University, China. HeLa cells were cultured in DMEM and transfectected with a total of 9 μg of pAAV1-TALEN-L, pAAV1-TALEN-R, Puro-Cas9, and Neo-MzRTTA donor (1:1:8:8). After transfection, cells were treated with G418 and puromycin. After antibiotic selection, 12 to 24 colonies were randomly picked based on HeLa morphology, mechanically disaggregated, and replated into individual wells of 96-well plates. CRISPR HeLa cells were treated with doxycycline before and during transfection. The DNA cassette targets SIRT1 were used to generate guide RNA (gRNA). After the gRNA transfection, CRISPR HeLa cells were dissociated into single cells and replated at 2,000 cells per 10-cm dish. Cells were allowed to grow until colonies from single cells became visible (∼10 d). Then single colonies were randomly picked mechanically to amplify in a 24-well dish. Detailed descriptions of all materials and methods are provided in SI Materials and Methods.

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