Inducible nitric oxide synthase (NOS2) produces high local concentrations of nitric oxide (NO), and its expression is associated with inflammation, cellular stress signals, and cellular transformation. Additionally, NOS2 expression results in aggressive cancer cell phenotypes and is correlated with poor outcomes in patients with breast cancer. DNA hypomethylation, especially of noncoding repeat elements, is an early event in carcinogenesis and is a common feature of cancer cells. In addition to altered gene expression, DNA hypomethylation results in genomic instability via retrotransposon activation. Here, we show that NOS2 expression and associated NO signaling results in substantial DNA hypomethylation in human cell lines by inducing the degradation of DNA (cytosine-5)–methyltransferase 1 (DNMT1) protein. Similarly, NOS2 expression levels were correlated with decreased DNA methylation in human breast tumors. NOS2 expression and NO signaling also resulted in long interspersed noncoding element 1 (LINE-1) retrotransposon hypomethylation, expression, and DNA damage. DNMT1 degradation was mediated by an NO/p38-MAPK/lysine acetyltransferase 5–dependent mechanism. Furthermore, we show that this mechanism is required for NO-mediated epithelial transformation. Therefore, we conclude that NOS2 and NO signaling results in DNA damage and malignant cellular transformation via an epigenetic mechanism.

Nitric oxide (NO) is an endogenously produced signaling molecule with a broad array of biological functions, ranging from neurotransmission to viral immune response (1–3). Inducible nitric oxide synthase (NOS2) is an inflammatory enzyme that produces high NO levels and initiates S-nitrosothiol (SNO) signaling, a term that we use herein to refer to signaling induced by NO-dependent cellular thiol oxidation (4, 5). SNO signaling results in altered protein function and signaling pathways (6–8), in addition to protein disulfide bond formation (9). NOS2 expression activates multiple oncogenic signaling pathways via SNO signaling (10, 11) and modulates the tumor immune response (12). Furthermore, NOS2 expression is associated with poor outcomes in patients with cancer and aggressive cancer cell phenotypes (13–18), while NOS2 inhibition reduces tumor burden and metastasis (19, 20).

DNA hypomethylation is frequently altered in human cancer cells compared with normal cells, as cancer cell genomes are predominantly hypomethylated with hypermethylated promoter regions of tumor suppressor genes (21–23). In addition to altered gene expression, global DNA hypomethylation results in increased DNA mutation rates (24), endogenous retrotransposon and retrovirus expression, genomic and microsatellite instability, and insertional mutagenesis (25–27). The most abundant retrotransposon in the human genome is the long interspersed noncoding element 1 (LINE-1), which is normally suppressed by promoter methylation but commonly is expressed in many human tumors. (26, 28). LINE-1 expression results in multiple genomic alterations, including indel mutations, transcriptional interference, and DNA strand breaks (29). Notably, both genome-wide hypomethylation and LINE-1 expression are observed in precancerous lesions, implicating DNA demethylation as an early event in tumorigenesis (30–32). Furthermore, the extent of DNA hypomethylation and LINE-1 expression also correlates with tumor stage (33, 34). Therefore, DNA hypomethylation and LINE-1 expression appear to be involved in both tumor initiation and progression.

DNA (cytosine-5)–methyltransferase 1 (DNMT1) is the critical epigenetic writer responsible for maintaining DNA methylation during replication (35), as well as de novo methyltransferase activity at retrotransposons (36). Two molecular processes achieve DNA demethylation: TET dioxygenase-mediated “active” and replication-dependent “passive” demethylation (37). While active demethylation is, in part, regulated by TET dioxygenase expression and cofactor availability, passive demethylation occurs during DNA replication in the absence of DNMT1 (37). DNMT1 protein

Significance

Nitric oxide is a multifaceted signaling molecule that affects multiple pathways and cellular systems. Here, we report that inducible nitric oxide synthase expression, which is strongly correlated with inflammation and poor outcomes in patients with cancer, substantially alters DNA methylation to regulate cellular plasticity. Our data connect inflammation mechanistically with stress signals, DNA demethylation, and genotoxic retrotransposon expression. Passive DNA demethylation occurs during conditions of reduced DNA (cytosine-5)–methyltransferase 1 (DNMT1) activity; however, the cellular pathways that control passive demethylation are not clear. Our results show that sustained cellular stress signals result in DNMT1 protein loss and DNA hypomethylation, similar to DNMT1 inhibition by 5-azacytidine. This implies that chronic inflammation drives cellular transformation via DNA hypomethylation and retrotransposon activation.
stability is governed by multiple posttranslational modifications, such as acetylation, methylation, and phosphorylation (38, 39). Reduced DNMT1 expression results in global hypomethylation, loss of genomic imprinting (40), and spontaneous tumors in mice (41). Furthermore, DNMT1 mutations that result in the deletion of the catalytic domain can be found in human colorectal cancers, implicating loss of DNMT1 activity during tumorigenesis (42). Therefore, decreased DNMT1 expression and activity are associated with tumorigenesis.

As tumor DNA hypomethylation and NOS2 expression are both associated with aggressive cancer phenotypes and poor survival rates among patients, we investigated the role of NOS2 on DNMT1 activity, expression and DNA methylation. Here, we show that NOS2 expression and SNO signaling caused the proteasomal degradation of DNMT1 and global DNA hypomethylation. Mechanistic studies revealed that NO-mediated DNMT1 degradation and epithelial cell transformation was p38-MAPK dependent. Consistent with DNA hypomethylation, we also show that NOS2 expression resulted in LINE-1 expression and DNA damage. These data indicate that NOS2 activity, via SNO signaling, dramatically alters cellular DNA methylation profiles to promote genomic instability and cellular transformation.

Results

NOS2 and SNO Signaling Induces Passive DNA Demethylation.

Cancer cell NOS2 expression is correlated with a basal-like breast cancer gene signature in estrogen receptor–negative (ER−) breast tumors (14). However, a majority of genes up-regulated in NOS2high compared with NOS2low ER− breast tumors have a repressive CpG island located at the transcriptional start site (SI Appendix, Table S1), which led us to hypothesize that NOS2 activity may alter DNA methylation levels. DNMT1 catalysis requires a cysteine at the active site and is, therefore, potentially inhibited by SNO signaling (5). To determine if methyltransferase activity is inhibited by SNO bond formation, recombinant human DNMT1 protein was incubated with the rapid-releasing, NO-donor 1,1-diethyl-2-hydroxy-2-nitroso-hydrazine sodium (DEANO; half-life [t1/2] = 2 min). NO significantly reduced rDNMT1 activity and resulted in DNMT1 S-nitrosation (SI Appendix, Fig. S1A). DEANO-treated isolated nuclei also exhibited reduced (cytosine-5)-methyltransferase activity (SI Appendix, Fig. S1B). However, in a whole-cell context using the slow-releasing NO-donor 2,2-(hydroxynitroso-hydrazono)bis-ethanimine (DETANO; t1/2 = 22 h), a seminal and intriguing observation in multiple human cell lines was made, namely that physiologically relevant concentrations of NO led to a significant loss of DNMT1 protein (Fig. 1A and SI Appendix, Fig. S1 C–F). DETANO reduced DNMT1 levels similar to the clinically used DNMT1 inhibitor, 5-azacytidine, which also results in DNMT1 protein loss (43). NO-mediated loss of DNMT1 has profound implications for cell lineage survival and identity, as we describe later, and occurs without significantly reducing cell proliferation or viability (SI Appendix, Fig. S2A). To examine the temporal regulation of DNMT1 by NO, cells were exposed to DETANO for the indicated times and DNMT1 protein was measured by immunoblotting. DETANO exposure decreased DNMT1 expression between 6 and 12 h and remained low during NO exposure (Fig. 1B). DNMT3A protein levels were not affected by NO signaling (SI Appendix, Fig. S2B). Similar to DNMT1 protein levels, total DNA methylation levels measured by liquid chromatography–mass spectrometry (Fig. 1C) and Illumina BeadChip arrays (Fig. 1D and E and SI Appendix, Fig. S2C and Dataset S1) were significantly reduced in NO-treated cells compared with control cells and were statistically similar to results after 5-azacytidine treatment. The decrease of DNMT1 expression followed an NO threshold–concentration effect, similar to the previously described SNO-mediated oncogenic pathway activation (10), NO-mediated loss of DNMT1 protein was not sensitive to soluble guanylyl cyclase inhibition (SI Appendix, Fig. S2D), and a stable cGMP analog did not significantly alter DNMT1 protein levels (SI Appendix, Fig. S2E). These data indicate that canonical NO/cGMP signaling does not contribute to reduced DNMT1 protein expression and suggest that NO mediates DNA demethylation by SNO signaling.

In addition to passive demethylation, TET iron dioxygenases catalyze the oxidation of 5mC during active DNA demethylation (37). However, NO treatment decreased TET expression, activity, and 5hmC content compared with vehicle-treated cells (SI Appendix, Fig. S2F–I). As NO is a pleiotropic signaling molecule that affects multiple pathways (5), it is conceivable that NO results in decreased DNA 5mC by an alternative mechanism that is independent of DNMT1 protein status and TET activity. However, the effect of NO-mediated DNMT1 protein degradation and DNA demethylation was rescued by DNMT1 overexpression (SI Appendix, Fig. S2J), indicating that SNO signaling caused the shift in global 5mC content by decreasing DNMT1 protein levels and inducing passive demethylation.

Human NOS2 transgene expression in NOS2-null breast cancer cell lines (MDA-MB-231 and MCF7) or a noncancerous cell line (HEK293) resulted in decreased DNMT1 protein levels (Fig. 2A and SI Appendix, Fig. S2K), which was partially reversed by NO inhibition (Fig. 2B), indicating that NOS2 activity and subsequent SNO signaling decrease DNMT1 protein expression. NO2 expression also resulted in a significant decrease in 5mC content, which was attenuated by NO2 inhibition (Fig. 2C). Additionally, IFN-γ treatment resulted in NOS2 expression and significant DNMT1 downregulation, which was partially reversed by the NOS2-selective inhibitor, 1400W (Fig. 2D). Similarly, high NO2 expression in human breast tumors is associated with decreased global DNA methylation levels compared with tumors with low NOS2 expression (Fig. 2E and SI Appendix, Table S2). Therefore, NOS2 expression and activity are associated with decreased DNA methylation.

SNO Signaling Induces DNMT1 Degradation.

To account for the reduced DNMT1 protein expression in response to SNO signaling, DNMT1 transcript expression was measured in MDA-MB-231 and MCF7 cell lines exposed to vehicle or 500 μM DETANO. In both cell lines, SNO signaling resulted in increased DNMT1 mRNA levels, indicating that SNO signaling does not acutely regulate DNMT1 protein at the transcriptional level (SI Appendix, Fig. S3A). However, DNMT1 protein levels did not change in response to SNO signaling in cells treated with the proteasome inhibitor MG-132 (SI Appendix, Fig. S3B), indicating that SNO signaling induces the degradation of DNMT1. To determine if SNO signaling resulted in DNMT1 ubiquitination (Ub), HEK293 cells coexpressing Myc-DNMT1 and HA-ubiquitin were treated with DETANO or vehicle. SNO signaling resulted in increased DNMT1-Ub compared with control cells (SI Appendix, Fig. S3C), indicating that SNO signaling reduces DNMT1 protein expression by inducing its proteasomal degradation.

p38-MAPK Activity Is Required for SNO-Mediated DNMT1 Degradation.

SNO signaling activates multiple intracellular pathways such as Src, Akt, and β-catenin (5, 10). Gene set enrichment
analysis of the NOS2-associated signature from patients with ER− breast cancer (14) revealed an enrichment of p38-MAPK–regulated transcription factors present in the NOS2 up-regulated genes using the TRANSFAC database (SI Appendix, Fig. S4 A and Table S3), and similar results were obtained from oPOSSUM transcription factor binding-site cluster analysis (SI Appendix, Fig. S4 B and C and Table S4) (44, 45). These data indicate that p38-MAPK activity is a dominant driver of NOS2-associated gene expression in human ER− breast tumors; therefore, the role of p38 in regulating DNMT1 protein levels was examined. NOS2 transgene expression and DETANO, at concentrations that reduce DNMT1 protein levels, resulted in significant p38 phosphorylation (Fig. 3 A and B and SI Appendix, Fig. S4 D and E). SNO signaling did not reduce DNMT1 protein levels in either CRISPR-mutated, kinase-inactive p38 (D168A) cells (Fig. 3 C), p38-inhibited or p38 kinase-dead mutant-expressing cells (SI Appendix, Fig. S4 F and G). These data strongly indicate that p38 activity is critical for SNO-signaling-induced DNMT1 degradation.

The requirement of p38-MAPK activity for SNO-mediated DNMT1 loss prompted us to examine if activation of this kinase is sufficient for DNMT1 loss and passive DNA demethylation. Direct p38 activation by expression of constitutively active MKK3b (Ser218Glu/Thr222Glu) double mutant (Glu) resulted in a plasmid concentration–dependent decrease of DNMT1 protein expression (Fig. 3 D and SI Appendix, Fig. S5A). MKK3b (Glu) expression failed to reduce DNMT1 levels in HEK293 cells coexpressing a kinase-dead p38 mutant (SI Appendix, Fig. S5B). Similar to SNO signaling, MKK3b (Glu) expression resulted in increased DNMT1 mRNA expression, DNMT1–Ub, and loss of DNMT1 protein that was blocked by proteasome inhibition (SI Appendix, Fig. S5 C–E). MKK3b (Glu) expression in HEK293 and MCF7 cells resulted in decreased DNA 5mC content, compared with control cells (Fig. 3 E). DNA demethylation by MKK3b (Glu) was blocked in HEK293 cells expressing a kinase-dead p38 mutant, indicating that p38 activity is required for both DNMT1 degradation.
and DNA demethylation (SI Appendix, Fig. S5F). MKK3b (Glu) expression resulted in replication-dependent passive DNA demethylation, as the DNA polymerase inhibitor aphidicolin blocked the ability of MKK3b (Glu) expression to reduce DNA 5mC content (SI Appendix, Fig. S4G). MKK3b (Glu) expression did not alter TET activity or 5hmC levels (SI Appendix, Fig. S5H and I), and TET2 protein expression was not significantly altered (SI Appendix, Fig. S5J), demonstrating that p38 signaling does not induce active DNA demethylation.

Transient p38 activation achieved by CD3/CD28 activation of Jurkat cells or EGF stimulation of breast cancer cells did not alter DNMT1 protein levels (SI Appendix, Fig. S5K and L). Thus, we conclude that prolonged p38 activation is sufficient to induce passive DNA demethylation.

**p38 Signaling Induces Passive DNA Demethylation Via KAT5 Activation.** DNMT1 protein stability is partially controlled by lysine acetylation (38) and p38 activates lysine acetyltransferase 5 (KAT5) to drive cellular senescence and p53-PUMA signaling (46, 47). To determine if KAT5 mediates DNMT1 degradation, cells were transiently transfected with either wild-type (WT) KAT5 (isoform 3) or a (T191A) mutant that cannot be phospho-activated by p38. Coexpression of MKK3b (Glu) with KAT5-WT in HEK293 cells resulted in decreased DNMT1 protein levels; however, MKK3b (Glu) expression did not alter DNMT1 protein levels in cells coexpressing mutant KAT5 (T191A) (SI Appendix, Fig. S6A). MKK3b (Glu) caused an increase in Myc-DNMT1 lysine acetylation in KAT5-WT–expressing cells, but not in KAT5 (T191A)–expressing cells (SI Appendix, Fig. S6B). Furthermore, MKK3b (Glu) expression significantly increased KAT5 activity toward rhDNMT1 in an in vitro acetyltransferase assay but did not increase KAT5 (T191A) activity (SI Appendix, Fig. S6C). Additionally, the KAT5-specific inhibitor TH1834 (48) blocked DNMT1 degradation by SNO signaling or MKK3b (Glu) expression (SI Appendix, Fig. S6D and E). These data indicate that p38-mediated KAT5 phosphorylation and activation are required for DNMT1 acetylation and subsequent degradation.

**SNO/p38 Signaling Induces LINE-1 Retrotransposon Expression and Double-Strand Breaks.** A consequence of global DNA demethylation is the activation of LINE-1 retrotransposons, which are normally suppressed by promoter methylation (29). LINE-1 repression is alleviated during global DNA demethylation...
events, and LINE-1 expression is commonly observed in human cancer cells (49). SNO signaling induced LINE-1 promoter demethylation similar to 5-azacytidine (SI Appendix, Fig. S7A). Consistent with L1 promoter demethylation, DETANO treatment resulted in LINE-1 Orf2p mRNA expression (Fig. 4A) and SNO signaling caused a concentration threshold effect on LINE-1 Orf2p expression, similar to DNMT1 degradation (Fig. 4B and SI Appendix, Fig. S7B). Additionally, NOS2 expression in HEK293 cells resulted in LINE-1 Orf2p expression, compared with mock transfected cells, and similar to cells expressing full-length L1.3 element (Fig. 4C). These data indicate that NOS2-mediated SNO signaling induces LINE-1–promoter demethylation and expression.

L1 expression results in DNA double-strand breaks (DSBs) (50); therefore, the effect of SNO signaling on DSBs was examined by γ-H2AX expression. Similar to DNMT1 degradation and L1 Orf2p expression, SNO-signaling levels of DETANO resulted in significantly increased γ-H2AX expression (Fig. 4D and SI Appendix, Fig. S7C) and punctate nuclear staining (Fig. 4E). NOS2 expression resulted in significant γ-H2AX expression (Fig. 4F and G) and was similar to cells expressing full-length L1.3 element (SI Appendix, Fig. S7D). Furthermore, DNA damage was strongly associated with proximal NOS2-GFP expression. Immunofluorescent imaging shows NOS2 and γ-H2AX expression had a positive Pearson correlation coefficient \( r = 0.874; P < 0.0001 \), suggesting that NOS2 expression is causally linked to γ-H2AX expression (Fig. 4G and SI Appendix, Fig. S7E).

To determine if p38 signaling is required for SNO-mediated DNA damage, CRISPR-mutated kinase-dead p38 (D168A)–expressing MCF10A cells were treated with DETANO or vehicle and compared with parental MCF10A breast epithelial cells. SNO signaling failed to induce γ-H2AX expression in p38 (D168A) mutant cells, in contrast to significant γ-H2AX expression in parental cells (Fig. 5A).

DNMT1 overexpression blocked SNO-mediated demethylation. To examine if DNMT1 overexpression rescues NO-induced DNA damage, Myc-tagged DNMT1-overexpressing cells were treated with DETANO and compared with mock transfected cells. While DETANO induced γ-H2AX expression in control cells, compared with vehicle-treated control cells, SNO signaling did not alter γ-H2AX expression in DNMT1-overexpressing cells (Fig. 5B), indicating SNO signaling induced DNA damage via DNMT1 protein degradation. Additionally, SNO-induced γ-H2AX expression was significantly attenuated in transiently LINE-1–silenced cells (Fig. 5C), indicating that SNO signaling and reactive nitrogen species induce DNA cleavage by an epigenetic-retrotransposition mechanism.

**Chronic SNO Signaling Transforms Breast Epithelial Cells Via a p38/DNMT1 Mechanism.** To determine if SNO signaling induced cellular transformation via DNA demethylation, a normal, human breast epithelial cell line (MCF10A) was chronically treated with DETANO or vehicle. Control cells retained a cuboidal, epithelial structure, whereas DETANO-treated cells exhibited an altered cellular morphology similar to a mesenchymal phenotype (Fig. 6A). However, in MCF10A p38 inactive-kinase cells, long-term exposure of DETANO resulted in cellular toxicity, while vehicle-treated mutant cells retained a cuboidal epithelial...
morphology. The toxicity observed in DETANO-treated p38-inactive cells is similar to the induction of apoptosis by hydrogen peroxide in p38-null cells (51).

After chronic exposure to DETANO, cellular proliferation was measured in real time via electrical impedance, which showed that DETANO-treated cells initially showed reduced proliferation compared with control cells (Fig. 6B). However, after approximately 60 h of recovery, DETANO-pretreated cells showed rapid proliferation compared with control cells, whereas inactive p38-mutated cells exposed to DETANO showed minimal growth over 4 d in culture. DNMT1 expression remained low after NO stress and returned to normal levels 3 d after removing DETANO from the culture medium, and this correlated with increased growth rates at 72 and 96 h after NO treatment (Fig. 6C). In parallel experiments, glucose and lactate levels were measured after 96 h by 1H-NMR (52). DETANO-treated cells showed increased glucose consumption and elevated lactate production compared with control cells (Fig. 6D and SI Appendix, Fig. S7). Long-term DETANO- and vehicle-treated cells were cultured for 72 h before assessing cellular migration.

Fig. 4. NOS2 and SNO signaling induce LINE-1 expression and DNA damage. (A) Relative LINE-1 Orf2p mRNA in DETANO-treated cells compared with vehicle controls. Data shown are mean ratios normalized to HPRT (±SEM; n = 6). Significance was calculated by two-tailed, unpaired t test with Welch’s correction. (B) Immunoblot of LINE-1 Orf2p and GAPDH expression in MDA-MB-231 cells cultured with DETANO for 48 h. (C) Immunoblot of LINE-1 Orf2p and GAPDH expression in HEK293 cells transiently transfected with either full-length human L1.3 element or NOS2-GFP expression plasmids and compared with mock transfected cells. (D) Immunoblot of γ-H2AX expression in DETANO-treated MDA-MB-231 cells. Graph represents mean densitometric analyses of γ-H2AX normalized to an α/β-tubulin from DETANO-treated MDA-MB-231 and MCF10A cell lines (±SEM; n = 3–5). Significance from vehicle controls was calculated by one-way ANOVA with Dunnett’s multiple comparisons test. (E) Immunofluorescent images of γ-H2AX expression in DETANO or vehicle for 48 h. γ-H2AX was visualized with Alexa488-conjugated secondary, and cells were counterstained with DAPI. (F) Immunoblot of γ-H2AX expression in mock or NOS-GFP–transfected HEK293 or MDA-MB-231 cells. Graph represents γ-H2AX densitometric analysis as fold change (fc) of control (±SEM; n = 6). Significance was calculated by two-tailed, unpaired t tests. (G) Pearson correlation analysis of proximal NOS2 and γ-H2AX expression in NOS2-GFP–transfected HEK293 cells. Regions of interest (ROI) from immunofluorescent images (SI Appendix, Fig. S7E) were analyzed for green and red channel intensities, normalized to mock-transfected ROI. Graph shows ROI signal intensities and linear correlation with calculated Pearson coefficient and two-tailed P value inset.
Consistent with a mesenchymal phenotype and cellular transformation, WT cells exposed to long-term SNO signaling exhibited increased chemotactic migration compared with vehicle-treated controls. In contrast, CRISPR-mutated inactive p38 kinase–expressing breast epithelial cells exposed to long-term SNO signaling showed decreased migration compared with vehicle-treated p38 inactive cells (Fig. 6E). To measure cellular transformation, vehicle- or DETANO-pretreated MCF10A cells were grown in a soft agar matrix. Consistent with altered metabolism and growth kinetics, DETANO pretreatment resulted in increased colony formation compared with vehicle-treated cells, whereas SNO signaling decreased the colony-forming ability of p38 inactive cells (Fig. 6F). NO resulted in the hypomethylation of cancer stem cell markers (e.g., Notch, ALDH1) and progrowth genes (e.g., ERK1) in a breast cancer cell line (Dataset S1). Thus, the expression of breast cancer stem cell markers was examined in NO-transformed breast epithelial cells. NO-transformed breast epithelial cells had significantly higher expression of breast cancer stem cell markers (CD44, Notch1, and ALDH1A3) and ERK1, a finding that correlates with increased growth rates and metabolic activity (Fig. 6G). Therefore, these results indicate that sustained SNO signaling resulted in cellular transformation via a p38-dependent, epigenetic mechanism.

Discussion

NO has numerous cellular signaling effects, some of which are due to protein SNO bond formation. Here, we tested the hypothesis that SNO signaling would inhibit DNMT1 activity to reduce DNA methylation levels. While S-nitrosation of the active-site cysteine residue resulted in inhibited DNMT1 activity, the cellular response of coordinated DNMT1 degradation was unexpected. Our results are consistent with recent reports that cellular oxidants inhibit S-adenosylmethionine formation to limit DNA methylation (53). Therefore, SNO signaling limits DNMT1 cofactor availability while simultaneously...
degrading and inhibiting this enzyme required for copying the methylome to the nascent DNA strand during replication.

NO is a potent driver of gene expression and can alter the transcriptome by modulating histone methylation (54). Here, we extend the known epigenetic functions of NO to include the regulation of DNA methylation. However, the effect of NO on DNA methylation has a detrimental outcome, as LINE-1 demethylation and expression induced by NO resulted in DNA...
resulted in DNMT1 degradation (Fig. 3). Required as direct p38 activation via mutant MKK3b also
more, we conclude that DNMT1-SNO formation is not
the cellular reducing capacity of cytosolic glutathione. Further-
ly, NO forms potent oxidants capable of DNA damage and
mutagenesis (61, 62). However, NO-derived oxidants, such as
NO2, are rapidly quenched by abundant cytosolic thiols such as glutathione. Furthermore, S-nitrosation of cellular thiols is
generally resolved to disulfides (9). Therefore, it is not clear
how NO-derived oxidants can penetrate the reductive capacity
of the cytosol to oxidize DNA. The p38-dependent DNMT1-
degradation mechanism is an elegant way to chemically bypass
the cellular reducing capacity of cytosolic glutathione. Further-
more, we conclude that DNMT1-SNO formation is not
required as direct p38 activation via mutant MKK3b also
resulted in DNMT1 degradation (Fig. 3 D and E); therefore, the
loss of DNMT1 is an indirect effect caused by NO2 and
SNO signaling. In the mechanism proposed here, the nitrosa-
tive signal is sensed by p38 and/or upstream activator(s) and is
transduced via a series of PTMs (i.e., KAT5 phosphorylation
and DNMT1 acetylation, Ub) to degrade DNMT1 protein,
allowing for passive DNA demethylation and the expression of
LINE-1 retrotransposons that causes genomic alterations and
DNA DSBs (Fig. 7). In addition to SNO signaling, reactive
oxygen species, nickel, and IL-6 also result in LINE-1 expres-
sion (26, 63). Therefore, oxidants such as NO may drive DNA
damage and genomic instability via an epigenetic and molecular
mechanism instead of direct chemical oxidation of DNA bases.

Our data indicate that NO2 expression and associated
SNO signaling promote aggressive tumor phenotypes by driv-
ing DNA hypomethylation and genomic instability via LINE-1
retrotransposon expression. This is consistent with the link
between tumors with high NO2 expression and epigenomic
instability in breast cancer with higher tumor grades and poor
outcomes in patients (14, 64). While NO2 inhibition has
therapeutic efficacy in breast cancer models (19), our mecha-
nism points to multiple targets for cancer chemoprevention or
intervention, as p38, KAT5, and protein acetylation can be
pharmacologically manipulated. Studies will be required to elu-
cidate if these targets limit tumorigenesis.

The loss of DNMT1 protein expression and passive DNA
demethylation caused by SNO signaling was dependent on
p38-MAPK activity. In addition to controlling cell cycle and
differentiation, p38 has both tumor suppressor and oncogene
functions. Interestingly, cellular differentiation and tumorigene-
sis both require DNA demethylation events, and the duration
of p38 activation appears to modulate its demethylation function,
as transient p38 activation did not alter DNMT1 levels. This is
consistent with differential signaling effects depending on whether
cells encounter chronic stress or acute mitogenic signals (65), sug-
uggesting that p38 signaling can be distinguished by source and
duration of stimulation. Given that cellular stress and chronic
p38 signaling is induced by carcinogens (66), it is plausible that
p38-mediated DNA demethylation may be a common event in
tumorigenesis, as we have linked the chronic cellular stress
response to epigenetic changes associated with tumor initiation.

A potential limitation of this study is the physiological relevance
of NO levels from NO2 transgene expression or a NO donor
compound. While both scenarios resulted in p38-MAPK activa-
tion and DNMT1 protein degradation, it is difficult to compare
the nitrosative environments of each scenario. NO2 activity
produces high, local, intracellular NO concentrations, while
DETANO releases NO into the bulk media, where it autoxidizes
to form nitrosating species. However, IFN-γ–induced NO2
expression also resulted in decreased DNMT1 protein expres-
sion, indicating that this effect is physiologically relevant. This is further
supported by DNA methylation levels being significantly decreased in breast tumors with high NO2 expression compared
with those with low NO2 expression. While the use of NO2 or
DETANO may have disadvantages, the results demonstrate that
DNMT1 is a target of nitrosative signaling in human cells.

To conclude, our data reveal an epigenetic function for NO2 and
cellular nitrosative signaling. NO2 activity and SNO signal-
ing induced passive DNA demethylation by chronically activating

Fig. 7. NOS2 promotes passive DNA demethylation and LINE-1 expression. (Left) Under normal physiological conditions, DNMT1 methylates the nascent
DNA strand during replication and LINE-1 retrotransposons are silenced by promoter CpG methylation. (Right) NOS2 expression results in sustained p38-
MAPK activation, which leads to DNMT1 protein degradation. Replication in the absence of DNMT1 activity results in passive DNA demethylation, LINE-1
expression, and genomic instability.
a p38/KAT5-signaling pathway to degrade DNMT1 protein. Consistent with genome-wide DNA hypomethylation, NOS2 and SNO signaling also resulted in LINE-1 retrotransposon expression. We conclude that NOS2 promotes cellular transformation and an aggressive cancer cell phenotype by inducing genomic hypomethylation, retrotransposition, and instability.

Materials and Methods

Cell Lines and Cell Culture. MDA-MB-231, MCF7, Jurkat E6.1 (European Collection of Authenticated Cell Cultures), HEK293, HL-60, and RAMOS (ATCC) were cultured in RPMI 1640 supplemented with 10% fetal bovine serum (FBS), penicillin (50 units/mL), and streptomycin (50 μg/mL) in a humidified incubator at 37 °C with 5% CO2. MCF10A breast epithelial cell line (ATCC) was cultured in mammary epithelial cell growth medium (MEGM) with MEGM SingleQuot Supple-
ments (hydrocortisone, rEGF, insulin, and bovine pituitary extract) as instructed (Lonza) and 100 ng/mL cholera toxin (Sigma). Cell lines were pas-
saged by 0.05% trypsin and 0.5 mM EDTA (Gibco) and quenched with serum-containing medium. MCF10A cells were trypsinized and quenched with Defined Trypsin Inhibitor (Thermo Fisher Scientific). Cell viability was measured by trypsin blue exclusion assay. Briefly, 10 μL of cell suspension was mixed with 10 μL of 4% trypan blue solution and total and trypan-stained cells were quantified by Countess II Automated Cell Counter (Thermo Fisher Scientific).

Chemicals and Reagents. DEANO and DETANO were obtained from Sigma and stock solutions were prepared in 10 mM NaOH. NO-donor purity was con-
firmed by measuring ultraviolet absorbance, ensuring a single symmetrical peak at 250 nm without a shoulder or nitrite peak at approximately 225 nm. Concentration was calculated using Beer’s law using 25 mM absorbance values (ε250 = 8000 M-1 cm-1). DETANO treatment of cells was done in RPMI + 10% FBS. We prepared stock solutions of 1400W (Sigma), 1H-[1,2,4]oxadiazolo[4,3-a]quinazine sodium (DEANO) or vehicle for 30 min at 37 °C in a 100-
μg of nuclear pro-
cell suspension was mixed with 10 μL of 4% trypan blue solution and total and trypan-stained cells were quantified by Countess II Automated Cell Counter (Thermo Fisher Scientific).

Plasmids and Transfections. The following constructs were obtained from Addgene: pCDNA3/Myc-DNMT1 (plasmid 36939, from A. Riggio), pRKS-HA-Ubiquitin-WT (plasmid 17608, from T. Dawson), pCDNA Flag MKK3b(Glu) (plasmid S0449, from R. Davis), and pCDNA Flag HA (plasmid 10792, from W. Selltes). We obtained pCDNA4-CGP-GFP-NOS2 (NOS2-GFP) from OriGene (RG211819), and pCDNA3.1 HA-KAT5 (isoform 3) and pCDNA3.1-HA-KAT5 (isoform 3, T191A) were synthesized by GenScript. Cells (0.25 x 106 in 3 mL of medium) were seeded into 6-well plates and cultured overnight. Plasmids were diluted in Opti-MEM medium (125 μL/well) and supplemented with P3000 (5 μL/well) and added to Lipofectamine 3000 (5 μL/well) diluted in Opti-MEM medium (125 μL/well). DNA complexes were incubated for 15 min at room temperature and added dropwise to cells. Mock-transfected cells were treated with Lipofectamine and P3000 diluted in Opti-MEM as described. Cells were then incubated for the times indicated before analysis.

DNMT Activity and S-Nitrosation. Full-length human recombinant DNMT1 (Reaction Biology Corp.) was desalted using a Zeba spin micro column and spayed to 1.5 mM in 20 mM Tris HCl at pH 7.5, aliquoted into microcentrifuge tubes, and incubated with 1,1-diethyl-2-hydroxy-2-nitroso-hydra-
zine sodium (DEANO) or vehicle for 30 min at 37 °C in a 100 μL reaction. DNMT1 activity was then measured using EpiQuik DNMT1 Activity/Inhibitor Screening Assay Core Kit (EpiGentek) as instructed using a final DNMT1 concent-
ration of 100 nM. Formation of SNO bond was determined by a biotin-switch method, as previously described (9). HL-60 nuclear fractions were prepared using Subcellular Protein Fractionation Kit for Cultured Cells (Thermo) as instructed, and protein concentrations were measured using bicinchoninic acid (BCA) assay. Nuclear fractions were then incubated with DEANO for 30 min at 37 °C, and total nuclear DNMT activity was measured using 1 μg of nuclear protein, as described.

Immunoblotting. Cells were lysed with 2x RIPA buffer containing complete protease inhibitor mixture and phosphatase inhibitor mixture 2 (Sigma). Lysates were denatured with 4x SDS sample buffer supplemented with 5% (vol/vol) 2-mercaptoethanol and incubated at 85 °C for 5 min. Lysates were then sepa-
rated on 4 to 20% or 4 to 15% sodium dodecyl sulfate polyacrylamide gel electrophoresis gels and transferred onto polyvinylidene fluoride or polyvinylidene difluoride membranes using the Bio-Rad Trans-Blot Turbo system. Membranes were blocked with 2% bovine serum albumin (BSA) in phosphate-buffered saline (PBS) containing 0.1% Tween-20 (PBS-tween) and incubated overnight at 4 °C with primary antibodies in 2% BSA in PBS-tween. Primary antibodies and dilu-
tions used are listed in SI Appendix, Table S5. After washing, membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibodies in 2% BSA PBS-tween for 1 h at room temperature. After washing with PBS-
tween, membranes were exposed to Pierce ECL reagent and imaged with Amersham Hyperfilm ECL or iBright FL1500. Immunoblots were quantified with Gel-Pro Analyzer v3.1 software.

Flow Cytometry Analysis. Cells (1.0 x 106 in 15 mL of medium) were seeded into 75 flasks and cultured overnight before vehicle, DETANO (500 μM), or 5-azacytidine (1 μM) was added and cultured for 48 h. Cells were then trypsi-
nized and quenched in defined trypsin inhibitor and washed in PBS. Cells were then fixed in 4% formaldehyde for 15 min at room temperature and washed in PBS. Cells were permeabilized with 90% cold methanol for 30 min and washed in PBS, followed by resuspension in PBS + 0.5% BSA. IgG-PE or anti-DNMT1-PE (3 μg/mL) was added to cells and incubated for 1 h in the dark at room temperature. Cells were then washed in PBS + 0.5% BSA before analysis on a BD FACSCan II analyzer. Data were then further analyzed using FlowJo software.

qPCR Analysis. RNA was isolated from cells using RNeasy Mini Kit (Qiagen) and quantified on a P200 Pipetman spectrophotometer. Complementary DNA (cDNA) was prepared using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) as instructed. qPCR assays for DNMT1 and H3F3B utilized the Universal ProbeLibrary System (Roche), and primers were designed using the Universal ProbeLibrary (UPL) Assay Design Center. Human LINE-1 was mea-
sured using iTaq Universal SYBR Green Supermix (Bio-Rad). UPL probes and pri-
ers used are listed in SI Appendix, Table S6. qPCR reactions were carried out using 5.1 ng of cDNA, 10 μM left and right primers, 10 μM UPL probe, and TaqMan Universal Master Mix II, with Uracl-N-glycosylase in a 20 μL reaction vol-
ume. qPCR reactions were performed on an Applied Biosystems StepOnePlus Real-Time PCR system. Relative expression normalized to HPRT was calculated using the comparative cycle threshold method.

DNA Methylation. DNA was isolated from cells using DNeasy Blood & Tissue Kit (Qiagen) and 1 μg of DNA was digested with DNA Degrades Plus (Zymo Research) for 4 h at 37 °C in a 25 μL reaction. After digestion, 0.1% formic acid was added to each sample and stored at −20 °C until liquid chromatography–tandem mass spectrometry analysis. Genomic SmCm, 5Hmc2, and total cytosine content was mea-
sured on an Agilent 1100 LC system interfaced with a Waters Quattro LC triple quadrapole mass spectrometer as previously described (67). Genomic methylation was measured with MethylhationEPIC Beadchip, using the manufacturer’s standard protocol (Illumina, Inc.). Genomic DNA was extracted from cells using DNeasy Blood & Tissue Kit and bisulfite converted with EZ-96 DNA Methylation Kit (Zymo Research Corp.). The converted DNA was processed by the Biomedical Research Centre based at Guy’s and St Thomas’ National Health Service Foundation Trust and King’s Col-
lege London. Genomic methylation data were analyzed using Partek Genomics Suite and only differentially methylated positions with a Δβ = ±0.2 relative to control samples were included for some analyses, as indicated in figure legends.

Human Tumor NOS2 and DNA Methylation Analysis. DNA methylation data (Illumina 450K BeadChip arrays) were obtained from a publicly available data set (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE37754) and corresponding NOS2 immunohistochemistry and patient data were generously provided by Dr. Stefan Ambs (National Cancer Institute, NIH, Bethesda, MD) (68). Mean M-values for each genome were plotted according to NOS2 expres-
sion level. Corresponding patient characteristics are shown in SI Appendix, Table S3.
TET Activity Assay. HEK293 cells (0.25 × 10^6 well in 3 mL of RPMI 1640 + 10% FBS) were seeded into 6-well plates and incubated overnight. Cells were then transiently transfected with FLAG-MKK3b (Glu) expression plasmid or FLAG-HA control plasmid, as described previously. Nuclear fractions were isolated 24 h after transfection and protein concentrations were determined by BCA assay. TET activity was measured using the Epigenase 5mC Hydroxylase TET Activity/Inhibition Assay Kit as instructed and normalized to the protein assessed.

KAT5 Activity Assay. HEK293 cells were transiently transfected with HA-KAT5 (WT) or HA-KAT5 (T191A) and FLAG-MKK3b (Glu), and cells were lysed after 24 h with RIPA buffer containing protease and phosphatase inhibitors. Lysates were then centrifuged to remove debris and supernatants were incubated overnight with anti-HA-tag-conjugated magnetic beads (Cell Signaling Technology, 11846) at 4 °C. Beads were then isolated in a magnetic field and washed three times with acetyltransferase activity buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 100 μM dithiothreitol). Recombinant human DNM1T (10 ng/reaction) and acetyl-CoA (50 μM; Sigma) were added to immuno-captured KAT5 and incubated at 37 °C for 30 min. Sample buffer (4X) was added to terminate reactions, and samples were analyzed by immunoblotting as described.

Immunofluorescence and Brightfield Microscopy. γ-H2AX expression was visualized by immunofluorescent imaging. Cells were grown on sterile glass coverslips and treated with DETANO or vehicle or transiently transfected with NOS2-GFP plasmid for 48 h. Cells were then incubated overnight at 4 °C, and permeabilized with PBS containing 0.3% Triton X-100 and blocked with 1% BSA in PBS containing 0.3% Triton X-100 for 1 h at room temperature. Coverslips were then incubated overnight at 4 °C with anti-γ-H2AX (Cell Signaling, #9718; 1:500 dilution) in blocking buffer. Coverslips were then washed with PBS containing 0.3% Triton-X100 and incubated with Alexa488- or Alexa594-conjugated secondary antibody, and developed with a colorimetric substrate. Coverslips were then incubated with 5mC antibody, washed, incubated with an HRP-conjugated secondary antibody, and bound to a streptavidin-coated, 96-well plate. Immobilized LINE-1 promoters were then centrifuged to remove debris and supernatants were incubated overnight with anti-RNA pol III promoter-conjugated magnetic beads (Cell Signaling Technology) and bound to a streptavidin-coated, 96-well plate. Immobilized LINE-1 promoters were then solubilized, and cells stained with a fluorescent dye and quantified.

Statistical Analysis. All statistical analyses were calculated using Prism GraphPad 9 unless otherwise stated. Comparisons between treatment groups and control were determined by one-way ANOVA with Dunnett’s post hoc test. Comparisons between two conditions were determined by two-way unpaired t test, while comparisons between two conditions and two variables were determined by two-way ANOVA with Sidak’s post hoc test. The effect of tumor NOS2 levels on DNA methylation was analyzed by two-tailed Welch’s t test (low NOS2 levels, n = 10; high NOS2 levels, n = 43). Hierarchical clustering was analyzed in RStudio running the pvclust package, an implementation of multiscale bootstrap resampling to calculate an approximately unbiased P value. Pearson correlation between NOS2-GFP and H2AX intensity was calculated with a two-tailed t value. Statistical significance was defined as P < 0.05.

Data Availability. All study data are included in the article and/or supporting information.

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Briefly, cells were seeded into 24-well plates and incubated for indicated times. Conditioned medium was passed through 0.22-μm filters and stored at –80°C until NMR analysis, and cells were assayed for protein content using the BCA assay with a BSA standard curve. Proton peaks corresponding to glucose and lactate in conditioned medium were measured and normalized to protein content of the well. Control (0 h) conditions were aliquots of fresh medium placed into 24-well plates and assayed as described.
