Oxidative Stress and Response to Thymidylate Synthase-Targeted Antimetabolites

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Received April 23, 2015; accepted October 5, 2015

ABSTRACT

Thymidylate synthase (TYMS; EC 2.1.1.15) catalyzes the reductive methylation of 2'-deoxyuridine-5'-monophosphate (dUMP) by N5,N10-methylenetetrahydrofolate (CH2=CH2PteGlu), to form dTMP and dihydrofolate (Carreras and Santi, 1995; Berger and Berger, 2006). The enzyme is indispensable to DNA synthesis and is therefore critical for replication and repair of the cellular genome in growing cells. As a consequence, it has been an attractive target at which chemotherapeutic drugs are directed (Chu et al., 2003; Longley et al., 2003; Wilson et al., 2014). Inhibitors of TYMS, including both fluoropyrimidine and folate analogs, lead to inhibition of the enzyme, resulting in dTMP deficiency and cell death. In the current study, we have examined the role of oxidative stress in response to TYMS inhibitors. We observed that intracellular reactive oxygen species (ROS) concentrations are induced by these inhibitors and promote apoptosis. Activation of the enzyme NADPH oxidase (NOX), which catalyzes one-electron reduction of O2 to generate superoxide (O2·-), is a significant source of increased ROS levels in drug-treated cells. However, gene expression profiling revealed a number of other redox-related genes that may contribute to ROS generation. TYMS inhibitors also induce a protective response, including activation of the transcription factor nuclear factor E2-related factor 2 (NRF2), a critical mediator of defense against oxidative and electrophilic stress. Our results show that exposure to TYMS inhibitors induces oxidative stress that leads to cell death, while simultaneously generating a protective response that may underlie resistance against such death.

Introduction

Thymidylate synthase (TYMS, EC 2.1.1.15) catalyzes the reductive methylation of 2'-deoxyuridine-5'-monophosphate (dUMP) by N5,N10-methylenetetrahydrofolate (CH2=CH2PteGlu), to form dTMP and dihydrofolate (Carreras and Santi, 1995; Berger and Berger, 2006). The enzyme is indispensable to DNA synthesis and is therefore critical for replication and repair of the cellular genome in growing cells. As a consequence, it has been an attractive target at which chemotherapeutic drugs are directed (Chu et al., 2003; Longley et al., 2003; Wilson et al., 2014). Inhibitors of TYMS, including both fluoropyrimidine and folate analogs, lead to inhibition of the enzyme, resulting in dTMP deficiency and cell death. In the current study, we have examined the role of oxidative stress in response to TYMS inhibitors. We observed that intracellular reactive oxygen species (ROS) concentrations are induced by these inhibitors and promote apoptosis. Activation of the enzyme NADPH oxidase (NOX), which catalyzes one-electron reduction of O2 to generate superoxide (O2·-), is a significant source of increased ROS levels in drug-treated cells. However, gene expression profiling revealed a number of other redox-related genes that may contribute to ROS generation. TYMS inhibitors also induce a protective response, including activation of the transcription factor nuclear factor E2-related factor 2 (NRF2), a critical mediator of defense against oxidative and electrophilic stress. Our results show that exposure to TYMS inhibitors induces oxidative stress that leads to cell death, while simultaneously generating a protective response that may underlie resistance against such death.

This work was supported by the National Cancer Institute [Grant CA44013]; and the National Institutes of Health Centers of Biomedical Research Excellence program of the National Institute of General Medical Sciences [Grant GM103336]. The Microarray Core Facility receives partial support from South Carolina Idea Networks of Biomedical Research Excellence program of the National Institute of General Medical Sciences [Grant GM103336]. The Microarray Core Facility receives partial support from South Carolina Idea Networks of Biomedical Research Excellence [Grant P20GM103499]. dx.doi.org/10.1124/mol.115.099614.

This article has supplemental material available at molpharm.aspetjournals.org.

ABBREVIATIONS: APO, apocyanin; DCFDA, 2'ʹ,7'ʹ-dichlorodihydrofluorescein diacetate; DHE, dihydroethidium; DPI, diphenyleneiodonium; dThd, deoxythymidine; dUMP, 2'ʹ-deoxyuridine-5'ʹ-monophosphate; FeUrd, 5-fluoro-2'ʹ-deoxyuridine; FDXR, ferredoxin reductase; FUra, 5-fluorouracil; GW1843U89, (S)-2-(5-(((3-methyl-1-oxo-1,2-dihydrobenzo[f]quinazolin-9-yl)methyl)amino)-1-oxoisindolin-2-yl)pentanedioic acid; LY231514, (2'ʹR)-5-[(2-amino-4-oxo-1,7-dihydropyrrolo[2,3-d]pyrimidin-5-yl)ethyl]benzoyl]amino]pentanedioic acid; NAC, N-acetylcysteine; NOX, NADPH oxidase; NRF2, nuclear factor E2-related factor 2; O2·-, superoxide; qRT-PCR, quantitative reverse-transcription polymerase chain reaction; ROS, reactive oxygen species; RTX, raltitrexed; siRNAs, small interfering RNAs; TYMS, thymidylate synthase; VAS, VAS-2870, VAS-2870, 3-benzyl-7'-2-benzoxazolylthio-1,2,3-triazolo[4,5-d]pyrimidine; ZD9331, (2'S)-2-[(2'-7-dimethyl-4-oxo-1'H-quinazolin-6-yl)methyl-prop-2-ynylamino]l-2-fluorobenzoyle[amino]-4-(2'H-tetrazol-5-yl)butanoic acid.
The mechanism by which dTMP deficiencies lead to the death of cells exposed to TYMS inhibitors is of central interest. Over a decade ago, Hwang et al. (Hwang et al., 2001) demonstrated that induction of reactive oxygen species (ROS) in FUra-treated cells results from increased expression of the mitochondrial enzyme ferredoxin reductase (FDXR), and it is required for p53-dependent apoptotic response to drug. Studies by Liu and Chen (2002) similarly suggested that FDXR promotes ROS-mediated apoptosis in response to FUra in a p53-dependent fashion. Despite these interesting studies as well as others showing increased ROS levels after exposure to FUra (Akhdar et al., 2009; Matsunaga et al., 2010; Lamberti et al., 2012), we still lack a detailed mechanistic understanding of how exposure to TYMS inhibitors alters the intracellular redox environment, and how cells respond to such alterations. Currently, we have examined ROS-mediated oxidative stress and its function in apoptotic cell death induced by TYMS inhibitors. We have identified the superoxide-generating enzyme NADPH oxidase (NOX) as an important source of increased ROS levels during drug exposure. In addition, we have found that cells mount a protective response that includes activation of the transcription factor nuclear factor E2-related factor 2 (NRF2) and its downstream response that includes activation of the transcription factor nuclear factor E2-related factor 2 (NRF2) and its downstream target genes, which may diminish the cytotoxic impact of TYMS inhibitors.

Materials and Methods

Reagents. FUra, FdUrd, deoxythymidine (dTd), folinic acid, and N-acetylcysteine (NAC) were purchased from Sigma-Aldrich (St. Louis, MO). RTX was obtained from AstraZeneca (Macclesfield, Cheshire, UK). Diphenyleleniodonium (DPI) and NAPDH were purchased from Calbiochem/EMD Biosciences (San Diego, CA). Apocynin (APO) was obtained from Acros Organics (Geel, Belgium). Lucigenin and VAS-2870 [VAS (3-benzyl-7-(2-benzoxazolyl)thiobenzimidazolyl)] were purchased from Enzo Life Sciences (Plymouth Meeting, PA). Paraformaldehyde was from Alfa Aesar (Ward Hill, MA). Dihydroethidium (DHE) and 2′,7′-dichlorodihydrofluorescein diacetate (H2DCFDA) were purchased from Invitrogen/Molecular Probes (Eugene, OR). Oligonucleotides were purchased from Integrated DNA Technologies (Corvalle, IA).

Cell Culture. Human colon tumor cell line HCT116 (originally obtained from Dr. Michael G. Brattain), FdUrd-resistant HCT116/200 (Berger et al., 1988), HCT-15 (American Tissue Culture Collection, Manassas, VA), and SW480 (American Tissue Culture Collection, Manassas, VA) were maintained in RPMI-1640 medium (Mediatech, Manassas, VA) containing 4.5 g/l glucose and supplemented with 10% heat-inactivated fetal bovine serum (Atlanta Biologicals, Norcross, GA), 100 U/ml penicillin, 100 mg/ml streptomycin, and 250 ng/ml amphotericin B (Mediatech) at 37°C in a humidified 5% CO2 atmosphere. Cells were exposed to the indicated concentrations of TYMS inhibitors (FUra, FdUrd, RTX) based upon previous studies in our laboratory (Barbour and Berger, 2008); 10 μM folinic acid was included with FdUrd. Where indicated, NAC (10 mM), dTdh (10 μM), DPI (10 μM), APO (1 mM), or VAS (10 μM) were included in the culture medium; all were added along with TYMS inhibitors except for DPI, which was added 3 hours before addition of TYMS inhibitors.

Detection of ROS. Intracellular ROS production was measured by oxidation of cell permeable dyes 2′,7′-dichlorodihydrofluorescein diacetate (DCFDA) to detect H2O2 and DHE to detect superoxide (O2·−). Cells were grown to 60% confluence in chamber slides, fixed in 4% paraformaldehyde, and incubated at 37°C for 30 minutes in 5 μM DCFDA or DHE. After incubation, cells were washed twice with fluorescence-activated cell-sorting staining buffer (phosphate-buffered saline with 1% bovine serum albumin) at 37°C for 15 minutes in 50 nM DCFDA or DHE. After incubation, cells were washed twice with fluorescence-activated cell-sorting buffer and analyzed by flow cytometry (FC 500; Beckman Coulter, Fort Collins, CO). A shift to the right indicates increased ROS levels.

Measurement of Apoptotic Cell Death. Cells were treated with TYMS inhibitors and inhibitory agents at the indicated concentrations for 48 hours. Apoptotic indices were determined by TUNEL assays using the In Situ Cell Death Detection Kit, POD (Roche Applied Science, Indianapolis, IN). Cells were stained according to the manufacturer’s instructions, counterstained with hematoxylin, and viewed under a light microscope at 400× magnification. Apoptotic nuclei were counted manually, based on staining and morphology, and the apoptotic index was calculated as the ratio of apoptotic/total cells. At least 1000 cells from several microscopic fields were counted in each determination.

NADPH Oxidase Activity Assay. For each reaction, 105 cells were suspended in 500 μl reaction buffer (50 mM phosphate buffer (pH 7.0), 1 mM EGTA, 150 mM sucrose). NADPH oxidase activity was detected by lucigenin-derived chemiluminescence with 100 μM NADPH as substrate and 5 μM lucigenin. Cells were incubated at 37°C for 10 minutes. Chemiluminescence was measured using a luminometer (Promega, Madison WI) and expressed as arbitrary light units per 105 cells.

RNA Extraction and Quantitative Reverse-Transcription Polymerase Chain Reaction. Total RNA was isolated using RNase Mini Kit (Qiagen, Germantown, MD) with the addition of RNase Free DNase (Qiagen) to eliminate contaminating genomic DNA. The RNA concentration was determined by measuring the absorbance at 260 nm and 280 nm (Nanodrop ND-1000 Spectrophotometer, Thermo Fisher Scientific, Waltham, MA). For each reaction, 1 μg RNA was reverse transcribed using an iScript cDNA synthesis kit (Bio-Rad Laboratories, Hercules, CA) according to the manufacturer’s instructions. For quantitative reverse-transcription polymerase chain reaction (qRT-PCR), cDNA (1μl) prepared as described earlier was amplified using Power SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA) according to the manufacturer’s instructions. The PCR thermal profile was one cycle at 95°C for 10 minutes followed by 40 cycles of 95°C for 15 seconds, 50°C for 15 seconds, and 72°C for 40 seconds using an Applied Biosystems 7300 Real Time PCR System. Relative mRNA levels were normalized to GAPDH, and calculated with the ΔΔCT method. Relative changes in expression of each gene in response to TYMS inhibitors were expressed as fold-induction compared with the basal level of expression in nontreated cells. Gene-specific primer sets (Integrated DNA Technologies) used for qRT-PCR are listed in Table 1. Controls with no reverse transcriptase and no template RNA were used to monitor contamination.

Transfection of Small-Interfering RNAs. Small-interfering RNAs (siRNAs) were introduced into cells by transient transfection using the DharmaFECT 4 reagent (GE Dharmacon, Lafayette, CO) in accordance with the manufacturer’s instructions. HCT116 cells were transfected with ON-TARGETplus siRNA for NOX2 (Dharmafect J-011021-05-0005), NFR2 (Dharmafect J-003755-09-0005), or scrambled siRNA (Dharmafect D-001810-01-05) as a negative control. Twenty-four hours after transfection, the cells were treated with various drug combinations and harvested at appropriate times for assays.

Microarray Analysis. Whole-genome cDNA microarray analyses were performed using Agilent’s platform (Agilent, Santa Clara, CA). Total RNA samples isolated from HCT116 were treated with Pura for 24 hours, then amplified and labeled using Agilent’s Low Input Quick Amp Labeling Kit (cat. no. 5190-2206) according to the manufacturer’s recommendations. Briefly, mRNA contained in 200 ng of total RNA was converted into cDNA using a poly-dT primer that also contained the T7 RNA polymerase promoter sequence. Subsequently, T7 RNA for measurement of ROS by flow cytometry, cells were grown to 80% confluence in 60-mm plates and incubated in fluorescence-activated cell-sorting staining buffer (phosphate-buffered saline with 1% bovine serum albumin) at 37°C for 15 minutes in 50 nM DCFDA or DHE. After incubation, cells were washed twice with fluorescence-activated cell-sorting buffer and analyzed by flow cytometry (FC 500; Beckman Coulter, Fort Collins, CO). A shift to the right indicates increased ROS levels.
polymerase was added to cDNA samples to amplify original mRNA molecules and to simultaneously incorporate cytarine 3- or cytarine 5-labeled CTP (cRNA) into the amplification product. In addition, Agilent RNA spike-in controls (cat. no. 5188-5279) were added to samples before cDNA synthesis and were used as experimental quality control. In the next step, labeled RNA molecules were purified using Qiagen’s RNeasy Mini Kit (cat. no. 74104). After spectrophotometric assessment of dye incorporation and cRNA yield, samples were stored at −80°C until hybridization. Labeled cRNA samples were hybridized to Agilent Human GE 4x44K v2 Microarrays (cat. no. G4845A) at 65°C for 17 hours using Agilent’s Gene Expression Hybridization Kit (cat. no. 5188-5242) according to the manufacturer’s recommendations. Four (4) control sample replicates were hybridized against four (4) FUra-treated sample replicates in a dye swap design. After washes, arrays were scanned using an Agilent DNA Microarray Scanner System (cat. no. G2565CA).

Data were extracted from images with Feature Extractor Software version 10.7.3.1 (Agilent). In this process, background correction using additive and multiplicative detrending algorithms was performed. In addition, linear and LOWESS methods were used for dye normalization. Subsequently, data were uploaded into GeneSpring GX version 11.5.1 (Agilent) for analysis. In this process, data were log2 transformed, quantile normalized, and base line transformed using the median of all samples. Then, data were filtered by flags such that 100% of the samples in at least one of the two treatment groups have a detected flag.

Differentially expressed genes were determined by analysis of the data using an unpaired t test statistics that was corrected for multiple testing using the Benjamini-Hochberg algorithm. A cutoff of 1.5-fold change with a P value of 0.005 was used. To identify genes involved in redox metabolism, the GO (Gene Ontology) and KEGG (Kyoto Encyclopedia of Genes and Genomes) databases were queried with the terms “dehydro, dehydro, hydro, hydrog, oxid, oxidase, oxidation, redox, reduc, reductase”; identified genes were assessed for their presence in the microarray.

Preparation of Extracts and Western Immunoblotting. Cytoplasmic and nuclear protein extracts were prepared using NE-Per Nuclear and Cytoplasmic Extraction Reagents (Thermo Fisher Pierce, Rockford, IL). Cells were harvested by trypsinization, and separation of cytoplasmic and nuclear fractions was performed according to the manufacturer’s instructions, with the following modifications: 1X SDS loading buffer was added directly to the nuclear pellet at 125 μl x 107 cells, sonicated 3 times at 10-second pulses, and loaded on the gel. Immunoblotting was performed by standard techniques. After SDS/PAGE, the fractionated proteins were transferred to nitrocellulose membranes, probed with an antibody to NRP2 (Cell Signaling Technology, Beverly, MA) followed by anti-rabbit IgG-HRP secondary antibody (Bio-Rad Laboratories). The antigen–antibody complexes were visualized using an enhanced chemiluminescence kit (GE Healthcare Biosciences, Piscataway, NJ). To monitor the separation of cytoplasmic and nuclear fractions, and to control for equal loading, blots were reprobed with anti-lactate dehydrogenase (Rockland Immunoclonicals, Limerick, PA) followed by anti-goat IgG-HRP secondary antibody (Bio-Rad Laboratories), and anti-lamin (Santa Cruz Biotechnology, Dallas, TX) followed by anti-rabbit IgG-HRP secondary antibody.

Statistical Analysis. Results are reported as the mean ± S.E.M., based on at least two to five independent experiments. Significance was assessed by calculation of two-tailed P values by an unpaired Student’s t test.

Results

ROS Levels Are Induced during Exposure to TYMS-Directed Antimetabolites. Several studies have shown that intracellular ROS levels increase in response to FUra (Hwang et al., 2001; Liu and Chen, 2002; Akhdar et al., 2009; Matsunaga et al., 2010; Lamberti et al., 2012). However, it has not been determined whether such increases are due to inhibition of TYMS as opposed to other effects of the antime-

We measured the effects of TYMS inhibition on ROS levels in HCT116 cells treated with various concentrations of the TYMS-directed fluoropyrimidine FdUrd. Cells were analyzed by flow cytometry after the addition of the ROS-detecting fluorescent dye DCFDA to the media. As shown in Fig. 1A, a progressive increase in ROS levels over a drug concentration range of 1 nM to 10 μM was observed, with the maximum level being reached at ~1 μM. To verify and extend these initial findings, we examined the impact of three TYMS-directed antimetabolites, including two fluoropyrimidines (FUra, FdUrd) and a folate analog (RTX), on intracellular levels of ROS. The concentration of each was based upon earlier studies in our laboratory (Barbour and Berger, 2008). Using both fluorescence microscopy and flow cytometry, we observed marked increases in ROS concentrations in response to each inhibitor (Fig. 1B). Similar results were obtained with a different fluorescent probe, DHE (data not shown). Further analyses showed that addition of NAC to the culture media abrogated drug-induced fluorescence (Fig. 1C), indicating that the observed increases in fluorescence indeed reflect augmentation of ROS levels after drug exposure. Furthermore, supplementation with dThd caused a loss of drug-mediated ROS induction (Fig. 1D), demonstrating that dTMP deprivation due to TYMS inhibition is occurring. Finally, ROS levels were profoundly diminished in the TYMS-overproducing cell line HCT116/200 (Fig. 1E), which is resistant to fluoropyrimidines as a consequence of TYMS overproduction derived from amplification of the TYMS gene (Berger et al., 1988). In all, these observations show that ROS induction in response to TYMS-directed antimetabolites arises from inhibition of TYMS and reduction in dTMP pools.

FdUrd elicited increased ROS levels in other human colon tumor cell lines, such as HCT-15 and SW480 (Fig. 1F). These lines also responded to FUra and RTX (data not shown). Thus, the observed induction of ROS after exposure to TYMS inhibitors is not a specific property of HCT116 cells but rather is broadly observed among distinct colon tumor cell lines.
ROS Induction by TYMS Inhibitors Promotes Apoptosis.

The observed augmentation of ROS levels in cells treated with TYMS inhibitors may promote oxidative stress that contributes to apoptotic cell death. In previous studies, we used TUNEL assays to show that apoptosis is increased by 10- to 20-fold in cells exposed to TYMS inhibitors, and was
completely inhibited by the broad-spectrum caspase inhibitor Z-VAD-FMK \([N\text{-benzyloxycarbonyl-Val-Ala-Asp(O-Me) fluoro-}
\text{methyl ketone}]\) (Barbour and Berger, 2008). We assessed the role of ROS in apoptosis induced by TYMS inhibitors. Figure 2 shows that the apoptotic index rose ~11- to 17-fold in response to FUra \((P = 3.7 \times 10^{-5})\), FdUrd \((P = 2.5 \times 10^{-6})\), and RTX \((P = 1.6 \times 10^{-5})\). Furthermore, addition of NAC reduced the index by ~50% \((P = 0.017)\) for FUra, 4.0 \(\times 10^{-4}\) for FdUrd, and \(7.0 \times 10^{-4}\) for RTX), indicating that abrogation of ROS levels reduces apoptotic response to TYMS inhibitors. Thus, oxidative stress is important to the cytotoxicity of these inhibitors.

**NOX Contributes to ROS Induction after Exposure to TYMS Inhibitors.** Previous studies had indicated that the mitochondrial enzyme FDXR is an important source of expanded ROS pools after exposure to FUra (Hwang et al., 2001; Liu and Chen, 2002). Indeed, gene expression profiling conducted in our laboratory (described herein) identified FDXR mRNA as among the transcripts induced by FUra. However, given the abundance of genes that function in redox pathways, it is highly likely that there exist additional enzymes that generate ROS in response to TYMS inhibitors. We tested the role of one such enzyme, NOX, which catalyzes the formation of \(\text{O}_2^+\) via one-electron reduction of \(\text{O}_2\). The enzyme, which exists as a membrane-bound, multisubunit complex whose sole function appears to be generation of \(\text{O}_2^+\) (Sumimoto, 2008; Jiang et al., 2011; Brandes et al., 2014), is activated by a wide variety of chemical, physical, and inflammatory stress stimuli (Jiang et al., 2011). As such, it is of interest to test the contribution of NOX to expanded ROS pools in response to TYMS inhibitors.

We used a whole-cell chemiluminescence assay to measure total NOX enzyme activity and the effect of TYMS inhibitors. As shown in Fig. 3A, both FUra and FdUrd elicit a 5- to 7-fold activation of enzyme activity \((P = 2.7 \times 10^{-3})\) for FUra and \(1.6 \times 10^{-3}\) for FdUrd). To be assured that NOX is being measured in these assays, we added the NOX inhibitor DPI (Wind et al., 2010) to the culture medium and observed complete inhibition of the drug-induced activity \((P = 1.6 \times 10^{-3})\) for FUra and \(1.1 \times 10^{-3}\) for FdUrd) (Fig. 3A). Attenuation of the increase in enzyme activity was also observed in the presence of dThd \((P = 0.024)\) for FUra and 0.014 for FdUrd), and was significantly reduced in TYMS-overproducing HCT116/200 cells \((P = 0.021)\) for FUra and \(8.6 \times 10^{-3}\) for FdUrd) (Fig. 3A). Thus, NOX activation in response to TYMS inhibitors is primarily due to inhibition of TYMS.

We determined whether NOX functions in drug-mediated accumulation of ROS. DPI inhibited ROS induction in response to FUra, FdUrd, and RTX, as measured by fluorescence assays with DCFDA (Fig. 3B). Furthermore, another NOX inhibitor, APO, which has a mechanism of action distinct from DPI (Aldieri et al., 2008), also prevented ROS induction by the three TYMS-targeted drugs (Fig. 3B). Because both DPI and APO are nonspecific and exert NOX-independent effects (Aldieri et al., 2008), we tested the impact of VAS, a triazolo pyrimidine that exhibits a high degree of specificity for NOX (Wind et al., 2010). VAS, similar to DPI and APO, reduced drug-mediated ROS accumulation (Fig. 3B). Similar findings as these were made using DHE as the fluorescent probe for ROS detection (data not shown). These results implicate NOX activation as functioning in augmentation of ROS levels after exposure to TYMS inhibitors.

We measured the impact of drug-mediated NOX activation on apoptosis. As shown in Fig. 4, increases in the apoptotic index after exposure to FUra, FdUrd, or RTX were abolished by both DPI \((P = 9.7 \times 10^{-6})\) for FUra, \(1.0 \times 10^{-6}\) for FdUrd, and \(3.4 \times 10^{-6}\) for RTX) and APO \((P = 4.1 \times 10^{-5})\) for FUra, \(1.1 \times 10^{-5}\) for FdUrd, and \(1.6 \times 10^{-5}\) for RTX), indicating an important role for NOX in drug response. VAS also abrogated drug-induced increases in apoptosis, but only by ~50% \((P = 0.013)\) for FUra, 0.047 for FdUrd, and 0.063 for RTX) (Fig. 4). The effect of VAS was consistently less than that of either DPI or APO, which may reflect the fact that the latter two exhibit a broader range of specificity than the former (Aldieri et al., 2008; Wind et al., 2010). This implies that factors other than NOX may contribute to the apoptotic response to TYMS-targeted agents.

**Expression of NOX Regulatory Subunit p67phox Is Stimulated by TYMS Inhibitors.** The NOX family is organized into a number of distinct multisubunit complexes, each member of which is comprised of one of seven known core catalytic subunits along with several regulatory subunits (Sumimoto, 2008; Block and Gorin, 2012; Brandes et al., 2014). To gain insight into the mechanism by which NOX is activated in cells treated with TYMS-directed drugs, we examined expression of mRNAs encoding several NOX subunits, including catalytic (NOX1-4 and p22phox) as well as regulatory (p47phox, p67phox, p40phox, NOX01, NOXA1, and Rac1). For most mRNAs, there were no detectable changes in levels of mRNA expression after exposure to FdUrd, as measured by qRT-PCR (data not shown). The exception was the NOX2 regulatory subunit p67phox, which is required for maximal activity of the NOX2 isoform (Italiano et al., 2012). Expression of p67phox mRNA was up-regulated ~20- to 25-fold by FdUrd \((P = 5.1 \times 10^{-5})\) (Fig. 5A). This up-regulation was prevented by addition of dThd \((P = 5.0 \times 10^{-8})\), and was significantly attenuated in TYMS-overproducing HCT116/200 cells \((P = 2.0 \times 10^{-5})\) (Fig. 5A). Thus, TYMS inhibition and subsequent reduction of dTMP pool levels are crucial to the up-regulation of p67phox.
Interestingly, the antioxidant NAC as well as NOX inhibitors DPI and VAS reduced the drug-mediated increases in p67phox mRNA ($P = 1.3 \times 10^{-3}$ for NAC, $8.0 \times 10^{-5}$ for DPI, and $9.0 \times 10^{-4}$ for VAS) (Fig. 5B). This suggests a “feed-forward” mechanism whereby ROS generated by NOX activation leads to induction of p67phox and NOX activity.

The observation of induced p67phox expression in response to TYMS inhibitors, coupled with the fact that p67phox specifically regulates the NOX2 isoform (Sumimoto, 2008; Jiang et al., 2011; Block and Gorin, 2012), implies the latter as significant in activation of NOX activity and subsequent apoptosis accompanying drug exposure. Therefore, we assessed the impact of NOX2 down-regulation using siRNA transfection. Introduction of NOX2-specific siRNAs led to ∼50% reduction in NOX2 mRNA in both control and FdUrd-treated cells ($P = 1.5 \times 10^{-3}$ and $2 \times 10^{-2}$, respectively), while
a control nonspecific “scrambled” siRNA had no effect (Fig. 6A). Reduction of NOX2 expression had essentially no impact on basal NOX activity levels before drug exposure (Fig. 6B), indicating that the isoform contributes little to NOX activity in normal cells. In contrast, the NOX2-specific siRNA completely inhibited the FdUrd-mediated increase in enzyme activity ($P < 0.011$) (Fig. 6B), signifying that this increase is largely due to NOX2. A modest, yet significant, $\sim 30\%$ reduction in the apoptotic index was observed in FdUrd-treated cells transfected with the NOX2-specific siRNA ($P = 0.013$), while there was measurable effect of the siRNA in untreated cells (Fig. 6C). Thus, associated with down-regulation of NOX2 mRNA expression was attenuation of drug-mediated induction of both NOX activity and apoptotic cell death, implying that the NOX2 isoform plays a significant role in the apoptotic response to TYMS inhibitors.

**Alterations in the Redox Transcriptome by TYMS Inhibitors.** In light of the results reported in the previous sections, we were interested in determining the impact of TYMS-directed antimetabolites on the cellular transcriptome, particularly with regard to mRNAs involved in redox metabolism. We therefore performed a whole-genome cDNA microarray analysis of HCT116 cells both before and after 24 hours of exposure to 10 $\mu$M FUra. Data were filtered to include transcripts exhibiting at least a 1.5-fold change with $P < 0.005$. We found 1975 genes whose levels of expression were altered by FUra; among these, 1119 (56.6%) were induced in response to drug, while 856 (43.4%) were repressed (Supplemental Table 1). For the induced genes, 452 (41%) increased by 2-fold or more, 63 (5.6%) by 4-fold or more, and 16 (1.4%) by 6-fold or more; the largest induction was observed for CEACAM1, which increased by $\sim 14$-fold. Among the repressed genes, 243 (28%) decreased by 2-fold or more, 12 (1.4%) by 4-fold or more;
more, and 4 (0.5%) by 6-fold or more; the largest change was exhibited by HIST1H2BF, which decreased by ~8-fold.

To identify redox-related transcripts (the redox transcriptome), we queried both the GO (Gene Ontogeny) and KEGG (Kyoto Encyclopedia of Genes and Genomes) databases using the terms “dehydro, dehydro, hydro, hydro, oxid, oxidase, oxidation, redox, reduc, reductase.” Those present in the microarray were considered FUrA-regulated redox genes. This strategy led to identification of 54 genes, 27 of which were up-regulated after drug exposure (e.g., AKR1B10, ALDH3A1, FDXR, GGT1, GLRX2, SAT1, SOD2, HMOX1, LOXL4, CBR3), and 27 that were down-regulated (e.g., DHRS2, CYP5RL, HSD17B8, ACADSB, CRYL1, SORD) (Supplemental Table 2). The presence of FDXR among the up-regulated genes corroborates earlier findings by others (Hwang et al., 2001; Liu and Chen, 2002). We found that p67phox did not show up in the microarray, likely due to its low levels of expression. Our findings, overall, indicate that exposure to TYMS inhibitors modifies the redox transcriptome within the cell through altered expression of a variety of redox-related genes.

**NRF2-Regulated Genes Are among Those Induced by TYMS Inhibitors.** Notably, several of the redox-related transcripts whose expression is modulated by FUrA (e.g., AKR1B10, ALDH3A1, GGT1, HMOX1) are known to be regulated by transcription factor NRF2. NRF2 is a member of the cap’n collar family of transcription factors that controls expression of genes conferring protection against oxidative and electrophilic stress (Baird and Dinkova-Kostova, 2011; Jaramillo and Zhang, 2013). Table 2 lists 27 transcripts from the microarray whose expression levels have been identified in the literature as regulated by NRF2 (Agyeman et al., 2012; Suzuki et al., 2013; Choi et al., 2014; Hayes and Dinkova-Kostova, 2014). We tested 15 of these by qRT-PCR and confirmed all as inducible by FdUrd (Table 2). An additional six transcripts, representing known NRF2-regulated genes that did not appear in the microarray, were examined by qRT-PCR and shown to be induced by FdUrd (Table 3). In all, we identified 33 NRF2-modulated genes whose levels of expression change in response to TYMS inhibitors.

We determined the extent to which these findings apply to additional colon tumor cell lines. Figure 7, A–D, shows the results of qRT-PCR analysis of mRNA levels for four genes (AKR1B10, ALDH3A1, HSPB8, and SERPINE1) in cell lines SW480 and HCT-15, as compared with HCT116. Though considerable heterogeneity in expression of these genes was observed among the three cell lines, induction of expression by FdUrd was observed in all three, though to variable extents. It is thus apparent that increased expression of NRF2-regulated genes in the presence of TYMS inhibitors is not specific to cell line HCT116, but occurs among distinct colon tumor cell lines.

To verify that these genes are indeed NRF2-regulated, we examined their expression after siRNA-mediated down-regulation of the transcription factor in HCT116 cells. As shown in Fig. 7E, we observed a 40%–60% reduction in mRNA levels for AKR1B10 (P = 0.017), ALDH3A1 (P = 0.016), HSPB8 (P = 0.043), and SERPINE1 (P = 0.019) after introduction of an NRF2-specific siRNA.

**Nuclear Accumulation of NRF2 and Its Role in Apoptotic Response to TYMS Inhibitors.** Under normal circumstances, NRF2 is maintained at low levels within the cell, a status maintained by the transcription factor’s nuclear accumulation. For example, we showed that introduction of an NRF2-specific siRNA can induce the expression of the target genes. This finding is consistent with previous reports that NRF2 is a key regulator of oxidative stress responses and that its nuclear accumulation is a critical event in this process. Furthermore, we observed that induction of NRF2 expression is associated with increased levels of the target genes, suggesting a causal relationship between these events.

**Table 2: Identification of FUrA-inducible NRF2 target genes by gene expression profiling.**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Description</th>
<th>Fold Change</th>
<th>Corrected P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABCC2</td>
<td>ATP-binding cassette, subfamily C, member 2</td>
<td>1.5861826</td>
<td>0.000741887</td>
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<tr>
<td>AGPAT9</td>
<td>Acylglycerol-3-phosphate O-acyltransferase 9</td>
<td>1.6375676</td>
<td>0.000784155</td>
</tr>
<tr>
<td>AHR</td>
<td>Aryl hydrocarbon receptor</td>
<td>1.742765</td>
<td>0.001289602</td>
</tr>
<tr>
<td>AKR1B10</td>
<td>Aldo-keto reductase family 1, member B10</td>
<td>6.701502</td>
<td>2.20317E-05</td>
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<tr>
<td>AKR1B15</td>
<td>Aldo-keto reductase family 1, member B15</td>
<td>6.0408144</td>
<td>0.002556888</td>
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<tr>
<td>AKR1C1</td>
<td>Aldo-keto reductase family 1, member C1</td>
<td>2.3066108</td>
<td>0.001858692</td>
</tr>
<tr>
<td>AKR1C3</td>
<td>Aldo-keto reductase family 1, member C3</td>
<td>1.9071968</td>
<td>0.001267411</td>
</tr>
<tr>
<td>ALDH3A1</td>
<td>Aldehyde dehydrogenase family 3, member A1</td>
<td>2.0209148</td>
<td>0.00248864</td>
</tr>
<tr>
<td>ALDH4A1</td>
<td>Aldehyde dehydrogenase family 4, member A1</td>
<td>2.560456</td>
<td>0.000557924</td>
</tr>
<tr>
<td>ATF3</td>
<td>Activating transcription factor 3</td>
<td>3.492217</td>
<td>6.41362E-05</td>
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<tr>
<td>BLRVA</td>
<td>Biliverdin reductase B</td>
<td>2.0149345</td>
<td>0.001494992</td>
</tr>
<tr>
<td>GADD45A</td>
<td>Growth arrest and DNA-damage-inducible, alpha</td>
<td>2.0611608</td>
<td>0.000269189</td>
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<tr>
<td>GBE1</td>
<td>Glucan (1,4-alpha,)-branching enzyme 1</td>
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<td>6.9456E-05</td>
</tr>
<tr>
<td>GGT1</td>
<td>Gamma-glutamyltransferase 1</td>
<td>2.475309</td>
<td>0.000289259</td>
</tr>
<tr>
<td>GLRX2</td>
<td>Glutaredoxin 2</td>
<td>1.6844928</td>
<td>0.000622618</td>
</tr>
<tr>
<td>GSR2</td>
<td>Glutamine synthetase</td>
<td>2.4446866</td>
<td>0.000183384</td>
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<tr>
<td>HMOX1</td>
<td>Heme oxygenase 1</td>
<td>2.0041764</td>
<td>1.4295E-05</td>
</tr>
<tr>
<td>HSPB8</td>
<td>Heat shock 22-kDa protein 8</td>
<td>2.7588468</td>
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<td>KIFAP3</td>
<td>Kinesin-associated protein 3</td>
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<tr>
<td>MAFF</td>
<td>v-Mafavian murine sarcoma viral oncogene homolog F</td>
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<td>8.8772E-05</td>
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<td>MGST3</td>
<td>Microsomal glutathione S-transferase 3</td>
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<td>0.003199293</td>
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<tr>
<td>MTT1B</td>
<td>Metallothionein 1B</td>
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<td>0.000482686</td>
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<tr>
<td>MTT2A</td>
<td>Metallothionein 2A</td>
<td>1.7349892</td>
<td>0.00248864</td>
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<tr>
<td>SAT1</td>
<td>Spermidine/spermine N1-acetyltransferase 1</td>
<td>2.4873772</td>
<td>6.04632E-05</td>
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<tr>
<td>SERPINE1</td>
<td>Serpin peptidase inhibitor, clade E, member 1</td>
<td>5.988837</td>
<td>7.4388E-05</td>
</tr>
<tr>
<td>SPP1</td>
<td>Secreted phosphoprotein 1</td>
<td>3.0035641</td>
<td>5.6966E-05</td>
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</tbody>
</table>
intrinsic short half-life. NRF2 is degraded by ubiquitin-mediated proteasomal degradation facilitated by the adaptor molecule Kelch-like ECH-associated protein 1 (KEAP1) (Baird and Dinkova-Kostova, 2011; Ma and He, 2012; Jaramillo and Zhang, 2013; Kansanen et al., 2013; Geismann et al., 2014). Under conditions of oxidative or electrophilic stress, KEAP1 becomes disabled, allowing NRF2 to escape degradation and accumulate in the nucleus, where it binds antioxidant response elements within promoter regions and induces transcription (Baird and Dinkova-Kostova, 2011; Jaramillo and Zhang, 2013). To determine the concentration and intracellular locale of NRF2 both before and after exposure to TYMS inhibitors, we performed Western immunoblotting of cytoplasmic and nuclear fractions of HCT116 cells. Figure 8A shows that NRF2 was present in the nucleus, with little, if any, in the cytoplasm. Furthermore, its concentration in the nucleus increases after treatment with FdUrd.

The finding that induction of NRF2-regulated genes by TYMS inhibitors is associated with an increase in the nuclear NRF2 concentration prompted us to ask whether such induction impacts apoptosis in FdUrd-treated cells. Therefore, we analyzed apoptotic indices in cells in which NRF2 levels were down-regulated by siRNAs (see Fig. 7E, above). For this analysis, we reduced the FdUrd concentrations to 10 μM to maximize the chances of observing an increase in apoptosis after reduction of NRF2 concentrations; this resulted in slightly lower extents of apoptosis as compared with experiments in Figs. 2, 4, and 6. As shown in Fig. 8B, the apoptotic index increased by ~80% in FdUrd-treated HCT116 cells transfected with NRF2-specific siRNAs, as compared with cells with a control siRNA (P = 0.017).

We also examined TYMS-overproducing HCT116/200 cells, which are relatively resistant to TYMS inhibitors (see Fig. 1) (Berger et al., 1988). These cells, as expected, exhibited a reduced apoptotic index relative to parental HCT116 cells; similar to the parental cells, there was a 70% increase in apoptosis after exposure to FdUrd (P = 0.024). Thus, down-regulation of NRF2 levels results in sensitization to drug exposure.

Discussion

ROS are well-known mediators of both tumorigenesis as well as response to cytotoxic agents (Ozben, 2007; Trachootham et al., 2009; Gupta et al., 2012; Nogueira and Hay, 2013). They are generally maintained at higher concentrations in tumor cells relative to normal cells, and can function as protumorigenic signaling molecules that promote cell growth and proliferation. However, due to their highly reactive nature, ROS levels that are excessive can lead to intracellular molecular damage, which includes chemical modification of macromolecules and small metabolites, oxidative stress, and cell death. Thus, cancer cells must maintain levels of ROS that are high enough to sustain robust cell growth yet below the threshold that initiates intracellular oxidative damage. This requires an intricate balance between pro-oxidants and antioxidants. Augmented ROS levels in tumor cells puts them closer to this threshold, making them vulnerable to excessive oxidative stress. This creates an opportunity to augment the efficacy of therapeutic agents used in the control of cancer (Trachootham et al., 2009). Indeed, it is likely that modification of the intracellular redox environment in the direction of excess ROS will enhance the sensitivity of tumors to cytotoxic drugs used in cancer chemotherapy.

In the current study, we examined the role of oxidative stress in the cytotoxic effects of TYMS-targeted antimetabolites. We have shown that ROS levels are increased after exposure to TYMS inhibitors such as Fura, FdUrd, and RTX (Fig. 1). Such increases are attenuated by the antioxidant NAC, by dThd, and by overproduction of TYMS. Thus, it is TYMS inhibition, as opposed to some other drug effect, that underlies stimulation of ROS levels. Drug-mediated apoptotic cell death, like the ROS levels, is reduced by NAC, indicating that ROS induction is critical to promotion of apoptosis (Fig. 2). These observations, in total, strongly suggest that oxidative stress plays a central role in cell death mediated by TYMS-targeted agents. It is evident that TYMS inhibitors stimulate increases in ROS concentrations that exceed the threshold required for initiation of oxidative stress and eventual cell death.

We have identified NOX as a source of increased ROS levels in cells responding to TYMS inhibitors. NOX occurs as a family of at least seven multisubunit complexes that catalyze transfer of an electron from NADPH to O₂, generating O₂− (Sumimoto, 2008; Jiang et al., 2011; Block and Gorin, 2012; Brandes et al., 2014). Activities of NOX complexes respond to a wide range of hormonal, chemical, viral, and bacterial stimuli, and have been implicated in a variety of physiologic functions, including host defense, cell signaling, and the regulation of gene expression (Sumimoto, 2008; Jiang et al., 2011; Block and Gorin, 2012).

Several findings described in our report support the notion that NOX contributes to oxidative stress and apoptotic cell death brought on by exposure to TYMS inhibitors. First, total NOX enzyme activity was significantly increased in the presence of drug (Fig. 3A). Second, three NOX inhibitors (DPI, APO, and VAS) markedly attenuated increases in both ROS levels (Fig. 3B) and apoptotic indices (Fig. 4) in response to TYMS inhibitors. Third, expression of p67phox, a regulatory subunit for the NOX2 isoform, was up-regulated in cells exposed to TYMS inhibitors (Fig. 5). Finally, introduction of a NOX2-specific siRNA into cells completely abrogated increases in NOX activity and resulted in attenuation of apoptosis after drug treatment (Fig. 6). These findings, collectively, indicate that NOX, in particular the NOX2 isoform, is a source of increased ROS in response to TYMS-directed agents.

Interestingly, drug-induced NOX enzyme activity was completely inhibited by the NOX2-specific siRNA (Fig. 6B)
Fig. 7. NRF2-regulated genes are induced by TYMS inhibitors. HCT116, SW480, and HCT-15 cells were treated for 24 hours with 10 μM FdUrd, and expression of mRNAs corresponding to several NRF2 target genes were analyzed by qRT-PCR, including (A) AKR1B10, (B) ALDH3A1, (C) HSPB8, and (D) SERPINE1. For each mRNA, values were expressed relative to those in untreated HCT116 cells. (E) HCT116 cells were transfected for 24 hours with a scrambled, nonspecific control siRNA or the NRF2-specific siRNAs. Expression of the indicated mRNAs, as well as NRF2 (inset), was analyzed by qRT-PCR.
The high rate of proteasomal degradation, which is mediated by NRF2 (Zhang, 2013; Hayes and Dinkova-Kostova, 2014). NRF2 is involved in the detoxification processes (Baird and Dinkova-Kostova, 2011; Jaramillo and Zhang, 2013). Consistent with this model of NRF2 activation, we find that exposure to FdUrd leads to an accumulation of the transcription factor in the nucleus (Fig. 8), providing a mechanism by which TYMS-directed antimetabolites activate NRF2-regulated transcription. Furthermore, we found that siRNA-mediated down-regulation of NRF2 leads to increased apoptosis in FdUrd-treated cells, indicating that the transcription factor inhibits apoptotic response to TYMS inhibitors and generates a chemoresistant state.

Analysis of several colon tumor cell lines showed that while considerable heterogeneity in expression of the NRF2-regulated genes exists, induction of expression by FdUrd was observed in all three, though to variable extents (Fig. 7). This heterogeneity among cell lines is not unexpected. It should be recalled that the cDNA microarray experiment leading to the identification of these genes was performed in HCT116 cells; it is therefore feasible that an overlapping yet distinct set of drug-inducible genes exist in the other cell lines, and would have been recognized had the microarray analyses been performed in those lines.

In conclusion, our study shows that TYMS-targeted antimetabolites activate NRF2-regulated antioxidant response element-containing genes (Baird and Dinkova-Kostova, 2011; Jaramillo and Zhang, 2013). Consistent with this model of NRF2 activation, we find that exposure to FdUrd leads to an accumulation of the transcription factor in the nucleus (Fig. 8), providing a mechanism by which TYMS-directed antimetabolites activate NRF2-regulated transcription. Furthermore, we found that siRNA-mediated down-regulation of NRF2 leads to increased apoptosis in FdUrd-treated cells, indicating that the transcription factor inhibits apoptotic response to TYMS inhibitors and generates a chemoresistant state.

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In a number of circumstances, NRF2 has been shown to confer relative resistance to chemotherapeutic agents (Kensler et al., 2007; Wang et al., 2008; Ma and He, 2012). Such may be the case for TYMS inhibitors. Our observations invite speculation that expression of NRF2-regulated genes and their induction by TYMS inhibitors may constrain therapeutic response to this class of drugs. Further studies will be necessary to determine which of the many NRF2-modulated genes mediate reduced drug response. Furthermore, the cDNA microarray analysis presented in the current report makes it clear that many genes that are not regulated by NRF2 are induced by exposure to TYMS inhibitors and could be involved in drug response. Thus, it is likely that NRF2 is one of many molecular players that control cellular response to TYMS inhibitors.

In conclusion, our study shows that TYMS-targeted antimetabolites induce oxidative stress that promotes cell death; at the same time, these agents activate antioxidative mechanisms that inhibit or reverse such stress and constrain drug response. Thus, the relative balance between these two processes determines the cytotoxic impact of drug exposure. This has important implications for the clinical use of TYMS inhibitors in cancer treatment. By reducing NRF2 levels within the cell and preventing the nuclear accumulation of the transcription factor that occurs in the presence of TYMS inhibitors, it may be possible to increase the efficacy of these drugs, thereby enhancing their clinical impact. With this in mind, ongoing efforts are aimed at a deeper understanding of the role of NRF2 as a determinant of response to TYMS-directed antimetabolites.
Oxidative Stress and Thymidylate Synthase Inhibitors

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