Multi-level gene expression profiles affected by thymidylate synthase and 5-fluorouracil in colon cancer

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Abstract

Background: Thymidylate synthase (TS) is a critical target for cancer chemotherapy and is one of the most extensively studied biomarkers for fluoropyrimidine-based chemotherapy. In addition to its critical role in enzyme catalysis, TS functions as an RNA binding protein to regulate the expression of its own mRNA translation and other cellular mRNAs, such as p53, at the translational level. In this study, a comprehensive gene expression analysis at the levels of both transcriptional and post-transcriptional regulation was conducted to identify response markers using human genome array with TS-depleted human colon cancer HCT-C18 (TS-) cells and HCT-C18 (TS+) cells stably transfected with the human TS cDNA expression plasmid.

Results: A total of 38 genes were found to be significantly affected by TS based on the expression profiles of steady state mRNA transcripts. However, based on the expression profiles of polysome associated mRNA transcripts, over 149 genes were affected by TS overexpression. This indicates that additional post-transcriptionally controlled genes can be captured with profiling polysome associated mRNA population. This unique approach provides a comprehensive overview of genes affected by TS. Additional novel post-transcriptionally regulated genes affected by 5-fluorouracil (5-FU) treatment were also discovered via similar approach.

Conclusion: To our knowledge, this is the first time that a comprehensive gene expression profile regulated by TS and 5-FU was analyzed at the multiple steps of gene regulation. This study will provide candidate markers that can be potentially used for predicting therapeutic outcomes for fluoropyrimidine-based cancer chemotherapy.

Background

Thymidylate synthase (TS) is a folate-dependent enzyme that catalyzes the reductive methylation of dUMP by 5,10-methylenetetrahydrofolate to form dTMP and dihydrofolate [1,2]. Because the TS-catalyzed enzymatic reaction provides the sole intracellular de novo source of thymidylate, an essential precursor for DNA biosynthesis, this enzyme has been an important target for cancer chemotherapy for the past 50 years [3-5]. TS is also one of the most extensively investigated biomarkers in recent years...
were loaded and resolved with 10% from both control and 5-FU treated HCT-C18 (TS+) cells.

In this regard, it is critical to develop more comprehensive molecular expression profiles to provide candidate genes that can potentially be used for predicting clinical outcomes for colorectal cancer.

The goal of this study is to provide global comprehensive gene lists that were deregulated by over-expression of TS. In this study, a systems biology approach was used to investigate genes altered by the overexpression of TS at both transcriptional and post-transcriptional levels using human genome expression array in TS-depleted human colon cancer HCT-C18(TS-) cells and HCT-C18 (TS+) cells stably transfected with the human TS cDNA expression plasmid. In addition, both cell lines were treated with 5-FU for different time periods in an attempt to identify both acute response genes as well as delayed response genes affected by the 5-FU exposure. A number of genes were identified by comparing the expression profiles obtained from both steady state total mRNAs and polysome associated mRNA transcripts isolated from 5-FU treated HCT-C18 (TS-) and HCT-C18 (TS+) cells. Comprehensive gene lists were generated that were relevant for both TS dependent and independent cytotoxicity by 5-FU exposure. We also discovered genes and pathways that may be involved for the oncogenic function of TS.

In this study, a systems biology approach was used to investigate genes altered by the overexpression of TS at both transcriptional and post-transcriptional levels using human genome expression array in TS-depleted human colon cancer HCT-C18(TS-) cells and HCT-C18 (TS+) cells stably transfected with the human TS cDNA expression plasmid. In addition, both cell lines were treated with 5-FU for different time periods in an attempt to identify both acute response genes as well as delayed response genes affected by the 5-FU exposure. A number of genes were identified by comparing the expression profiles obtained from both steady state total mRNAs and polysome associated mRNA transcripts isolated from 5-FU treated HCT-C18 (TS-) and HCT-C18 (TS+) cells. Comprehensive gene lists were generated that were relevant for both TS dependent and independent cytotoxicity by 5-FU exposure. We also discovered genes and pathways that may be involved for the oncogenic function of TS.

The rational for investigating genes affected by TS and 5-FU at both transcriptional and post-transcriptional levels is that TS, in addition to its critical enzymatic function, may be involved for the oncogenic function of TS.

Previous studies have demonstrated that the mechanism for the decreased p53 expression occurred at the translational level [16]. We use p53 as a positive control gene for monitoring our high throughput expression profiling approach. Our microarray gene expression analysis indicates that there is no difference in the expression level of p53 based on steady state total mRNA (Additional file 1). However, the expression level of actively translated p53

be particularly important to systematically investigate other potential post-transcriptional regulated genes via TS protein. This may be especially vital for the discovery of additional chemotherapeutic response related markers that otherwise would be missed by simply profiling steady state total mRNAs [18]. A recent report suggested that TS may also function as an oncogene to transform NIH3T3 cells [19]. A comprehensive gene expression profiling analysis may also gain new insights into signaling pathways that were deregulated by over-expression of TS. In this regard, it is critical to develop more comprehensive molecular expression profiles to provide candidate genes that can potentially be used for predicting clinical outcomes for colorectal cancer.

The rational for investigating genes affected by TS and 5-FU at both transcriptional and post-transcriptional levels is that TS, in addition to its critical enzymatic function, functions as a RNA binding protein [11]. The translation of human TS mRNA is regulated by its own protein product via a negative autoregulatory mechanism whereby the binding of TS protein to at least two distinct sequences on its own TS mRNA results in translational repression [12,13]. TS is also capable of interacting with several other cellular mRNAs such as p53 mRNA and c-Myc mRNA [14,15]. Previous studies demonstrated that TS protein regulates p53 gene expression at least in part, at the translational level [16]. In this case, TS may be involved in coordinating the regulation of expression and/or function of cellular growth and proliferation and it is conceivable that TS may play an essential role as a regulator of cell cycle related events. More importantly, this study will have direct clinical relevance in that the mechanism of acute and long term 5-FU related drug resistance is distinct. The acute induction of TS expression after 5-FU treatment was regulated at the translational level and long term resistance for 5-FU is related to transcriptional activation and gene amplification of TS [17]. Thus, it would
Figure 2
Clustering analysis of genes expression affected by TS overexpression with HCT-C18 (TS+) (Lane 1) vs. HCT-C18 (TS-) (Lane 2) based on the profiles of steady state total mRNAs. One-way ANOVA analysis was used for the gene clustering with $p < 0.05$. 
Clustering analysis of genes expression affected by TS overexpression with HCT-C18 (TS+) (Lane 1) vs. HCT-C18 (TS-) (Lane 2) based on the profiles of polysome associated mRNAs. One-way ANOVA analysis was used for the gene clustering with p < 0.05.

Figure 3
Clustering analysis of genes expression affected by TS overexpression with HCT-C18 (TS+) (Lane 1) vs. HCT-C18 (TS-) (Lane 2) based on the profiles of polysome associated mRNAs. One-way ANOVA analysis was used for the gene clustering with p < 0.05.
mRNA in HCT-C18 (TS+) cells was found to decrease by 4.5-fold compared to the p53 mRNA level in HCT-C18 (TS-) cells (Additional file 2). This is consistent with our previous results that p53 was regulated at the translational level via TS [16].

**Effect of 5-FU on the expression of TS and p53**

The effect of 5-FU on the expression of both TS and p53 was investigated using Western immunoblot analysis. HCT-C18 (TS+) cells were treated with 10 µM 5-FU for 4 hrs and 24 hrs. TS protein expression was increased by nearly 4-fold (lane 2) at 4 hrs and 10-fold at 24 hrs (lane 3) (Figure 1). The TS-FdUMP-tetrahydrofolate covalent ternary complex was clearly visible after 5-FU treatment. The expression of wild type p53 was also increased in response to 5-FU treatment (Figure 1, lane 2–3).

**Effect of TS on steady state total mRNAs expression**

The global gene expression profile of HCT-C18 (TS-) and HCT-C18 (TS+) cells based on steady state total mRNA expression was analyzed using human high density CodeLink oligo array (20 K). Although the growth characteristics of the two cell lines are similar in terms of doubling time, many genes are already affected by overexpressing TS protein. Our analysis revealed that over 38 genes were changed in their expression in response to TS expression (n = 3, p < 0.05 and 4-fold cut-off) (Figure 2). The partial gene list and the clustering analysis are shown in Figure 2 (For complete gene list see Additional file 2). Genes such as CENTB1 and MT3, which are involved in cell proliferation and signaling, were up-regulated by TS. CASP1, a gene involved in regulation of I-kB/NF-κB cascade, apoptosis and signal transduction, was decreased 22-fold by TS overexpression.

**Effect of TS on polysome associated mRNAs**

To discover novel genes post-transcriptionally regulated by TS protein, we isolated the polysome associated mRNAs from both HCT-C18 (TS-) and HCT-C18 (TS+) cells and performed expression analysis using human high density CodeLink oligo array (20 K). Over 149 genes were affected in TS overexpressing HCT-C18 (TS+) cells (n = 3, p < 0.05 with 4-fold cut-off) (Figure 3; Additional file 2). Genes involved in protein biosynthesis (FLJ10989, PEX1, FLJ20450, PSMC6, MRPL3) and cell cycle control (MAD2L1, INHBA, APPL, D123) were up-regulated at the post-transcriptional level. The overlapping genes (such as INHBA, and CASP1) between steady state total mRNA and polysome associated mRNA profiles are listed in Table 1.

**Steady State mRNAs affected by 5-FU treatment in HCT-C18 (TS+) Cells**

To identify acute and delayed response genes following 5-FU treatment in HCT-C18 (TS+) cells, gene expression profiling analysis was performed on steady state total mRNAs isolated from control and 5-FU treated samples at 4 hrs and 24 hrs time points. The expression analysis reveals that over 46 genes were affected by 5-FU treatment by One-way ANOVA analysis (n = 3, p < 0.05 with 4-fold cut-off). The clustering analysis is shown in Figure 4. (For complete gene list see Additional file 3). The expression analysis clearly showed a different gene expression profiles between 4 hrs and 24 hrs 5-FU exposure. Based on the comparison of the expression profiles among the HCT-C18 (TS+) control and 5-FU treated samples, a number of genes were acutely increased in response to 5-FU. Genes (shown in orange color in Figure 4, lane 2) such as ORCL6 (DNA replication), PRPS1 (nucleotide

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**Table 1: Overlapping genes between steady state and polysome associated mRNAs affected by TS overexpression**

| Genebank accession | Gene ID | Fold change | Biological function |
|--------------------|---------|-------------|---------------------|
| **Increased genes** |         |             |                     |
| NM_002192          | INHBA   | +21.81      | +33.02              |
| NM_000930          | PLAT    | +11.93      | +6.73               |
| ABO32261           | SCD     | +5.48       | +6.62               |
| **Decreased genes**|         |             |                     |
| NM_001442          | FABP4   | -100.00     | -100.00             |
| NM_002153          | HSD17B2 | -54.95      | -100.00             |
| NM_033292          | CASP1:COPI | -22.22   | -11.68              |
| NM_003641          | IFITM1  | -18.08      | -23.58              |
| NM_002274          | KRT13   | -7.81       | -11.90              |
| NM_002638          | PI3     | -5.75       | -10.41              |

- **Increased genes**: INHBA +21.81 +33.02; PLAT +11.93 +6.73; SCD +5.48 +6.62.
- **Decreased genes**: FABP4 -100.00 -100.00; HSD17B2 -54.95 -100.00; CASP1:COPI -22.22 -11.68; IFITM1 -18.08 -23.58; KRT13 -7.81 -11.90; PI3 -5.75 -10.41.

- **Biological function**: Cell cycle arrest; cell differentiation; signal transduction; cell-cell signaling; Blood coagulation; protein modification; proteolysis and peptidolysis; Fatty acid biosynthesis; Transport; Estrogen biosynthesis; metabolism; Regulation of I-kB kinase/NF-κB cascade; apoptosis; signal transduction; Signal transduction; immune response; cell proliferation and cycle; Epidermis development; Copulation.
Clustering analysis of TS dependent cytotoxicity in HCT-C18 (TS+) cells (Lane 1) treated with 10 µM 5-FU for 4 hrs (Lane 2) and 24 hrs (Lane 3) based on the expression profiles generated from steady state total mRNAs via One-way ANOVA analysis with $p < 0.05$.

**Figure 4**
Clustering analysis of TS dependent cytotoxicity in HCT-C18 (TS+) cells (Lane 1) treated with 10 µM 5-FU for 4 hrs (Lane 2) and 24 hrs (Lane 3) based on the expression profiles generated from steady state total mRNAs via One-way ANOVA analysis with $p < 0.05$. 

SPRR1A
SCNN1B
DDB2
SPRR3
KRT17
S100A2
AKAP12
DKFZp762F0713
CDKN1A
LOC57822
KRT6A
PLAU
SPRR1B
IL8
FLJ23311
C4BPB
HCC4
Dkk4
ORC6L
SEI1
CXCR4
HSD17B2
FAM20C
CDIPT
PRPS1
CHC1L
GP1BB
RRM2
SPRR
SPINK4
STARD7
MFN2
MCM10
RPS26
CD38
SPN
CST8
LOC51659
SAMHD1
H4F2
H2AFX
EIF4E
DDX15

5-FU Treatment
Control 4 hours 24 hours
Clustering analysis of TS dependent cytotoxicity in HCT-C18 (TS+) cells (Lane 1) treated with 10 µM 5-FU for 4 hrs (Lane 2) and 24 hrs (Lane 3) based on the expression profiles generated from polysome associated mRNAs via One-way ANOVA analysis with p < 0.05.

**Figure 5**
Clustering analysis of TS dependent cytotoxicity in HCT-C18 (TS+) cells (Lane 1) treated with 10 µM 5-FU for 4 hrs (Lane 2) and 24 hrs (Lane 3) based on the expression profiles generated from polysome associated mRNAs via One-way ANOVA analysis with p < 0.05.
To further define the genes that were associated with TS independent toxicity, we compared the expression clusters (Figures 4 and 6) from both HCT-C18 (TS-) and HCT-C18 (TS+) cells treated with 5-FU using Venn diagram analysis. The overlapping gene list is shown in Table 2. Genes such as RRM2 (DNA replication), CDKN1A (G1 arrest), SEI1 (cell proliferation), DDB2 (nucleotide-excision repair), SFN (cell cycle regulation) were associated with TS independent cytotoxicity to 5-FU treatment. Gene expression analysis from Figure 4 was used for the construction of the cell cycle control pathway after 5-FU treatment illustrated in Figure 7.

Discussion
In this study, a systems biology approach was used to analyze global gene expression and its networks affected by TS and 5-FU at multiple levels of control. To the best of our knowledge, this is the first report to address genes that are affected at both transcriptional and post-transcriptional levels affected by elevated TS protein expression and 5-FU treatment. We believe that this will provide a better understanding of complex network related to the multi-functionalities of TS protein as an RNA binding protein or as a potential oncogene. It also provides a technology platform to systematically discover potential post-transcriptionally controlled genes by TS and 5-FU treatment. As our comparative model, we used a pair of human colon cancer cell lines: the mutant HCT-C18(TS-) subline in which the TS protein had been rendered marginally active and the TS over-expressing subline, HCT-C18(TS+), created by stable expression of a human TS cDNA plasmid. Both cell lines express wild-type p53 protein [16]. p53 is one of the known genes affected by the over-expression of TS at the translational level due to its RNA binding function [16]. Using our expression profiling approach, the levels of p53 mRNA are not significantly changed based on steady state total mRNA profile. However, the polysome associated p53 mRNA was decreased by 4.5-fold in HCT-C18 (TS+) cells compared with HCT-C18 (TS-) cells (listed in Additional file 2). These results, taken together, confirmed that our comprehensive gene profiling analysis approach is capable of discovering new post-transcriptionally regulated genes.

The gene profiling analysis based on steady state total mRNA transcripts revealed over 38 genes that were deregulated by TS over-expression (Figure 2 and Additional file 1). Although this study does not provide any direct evidence for TS as a cell cycle regulator, a number of cell cycle control and apoptotic control genes were perturbed by TS over-expression. One of the genes is inhibin/activin A (INHBA) that was up-regulated by nearly 22-fold in HCT-C18 (TS+) cells. INHBA is a growth factor that is involved in cell proliferation [21]. INHBA does this through type I and type II receptor serine kinases [21]. This may be a potential pathway by which TS is able to influence the cellular transformation process [21-23]. In fact, it has been...
Clustering analysis of predictive marker genes for TS independent cytotoxicity in HCT-C18 (TS-) cells (Lane 1) treated with 10 µM 5-FU for 4 hrs (Lane 2) and 24 hrs (Lane 3) via One-way ANOVA analysis with p < 0.05.

**Figure 6**
Clustering analysis of predictive marker genes for TS independent cytotoxicity in HCT-C18 (TS-) cells (Lane 1) treated with 10 µM 5-FU for 4 hrs (Lane 2) and 24 hrs (Lane 3) via One-way ANOVA analysis with p < 0.05.
reported recently that INHBA is over-expressed in stage IV colorectal cancer [24]. Additionally, single stranded RNA interacting protein (RBMS3), a gene responsible for mRNA splicing, was also up-regulated by 28-fold. The gene is a member of a small family of proteins which bind single stranded DNA/RNA. These proteins are characterized by the presence of two sets of ribonucleoprotein consensus sequence (RNP-CS) that contain conserved motifs, RNP1 and RNP2, originally described in RNA binding proteins, and required for DNA binding. These proteins have been implicated in such diverse functions as DNA replication, gene transcription, cell cycle progression and apoptosis [25]. The encoded protein was isolated by virtue of its binding to an upstream element of the alpha2 (I) collagen promoter. The observation that this protein localizes mostly in the cytoplasm suggests that it may be involved in a cytoplasmic function such as controlling RNA metabolism, rather than transcription. Thus, RBMS3 may play an important role as part of the post-transcriptional control mediated by TS. Conversely, genes such as caspase 1 (CASP1) that control cell cycle arrest and apoptosis, were down-regulated some 22-fold. Hence, future studies will focus on determining whether TS protein regulates the expression of these genes directly by binding to the mRNAs or indirectly through downstream mechanisms.

Using our systems biology approach of polysome associated mRNA comparisons, we have identified a list of genes that are over- or under-expressed due to TS protein overexpression (Figure 3 and gene list in Additional file 2). Many of them play key roles in cell-cell signalling, protein biosynthesis, RNA processing, DNA repair, cell cycle control and translational regulation. Some of them are previously known translational controlled genes such as EGFR [26], one of the key targets for the latest targeted anticancer drug development for colon cancer treatment [27,28]. In this study, the expression of EGFR was found to
increase by nearly 8-fold by TS over-expression. Many other genes on this list are novel potential post-transcriptionally regulated genes. Among the down regulated genes, cyclin dependent kinase inhibitor p21 (CDKN1A) was decreased by over 5-fold. Apoptosis inhibitor BAK1 was also decreased by 6-fold in TS over-expressing cells. It seems that over expressing TS has set the stage for cells to have many growth advantages using multiple mechanisms.

To identify genes regulated at both transcriptional and post-transcriptional level, we performed an overlapping analysis by comparing the gene lists generated from steady state total mRNAs and polysome associated mRNAs using Venn diagram. There are 15 overlapping genes between the two lists (Table 1). D123 (Gdc123), up-regulated by 5-fold in HCT-C18 (TS+) cells, has been shown to be regulated at the translational level via elf2γ [29]. We have identified an additional 136 genes based on polysome associated mRNAs which could be easily missed if only steady state mRNA profiling is performed. This approach provides a technical platform to systematically analyzing gene expression at post-transcriptional level. It also reveals the importance of analyze gene expression at multiple levels.

Based on the clinical observation and detailed molecular investigations [11,17], TS can be acutely induced with a short time 5-FU exposure and the mechanism of TS induction was regulated, at least in part, at the translational level [11]. With this notion, we designed a strategy that allows us to capture both acute and delayed response genes in response to 5-FU exposure. To achieve this, both HCT-C18 (TS+) and HCT-C18 (TS-) cells were treated with 10 μM 5-FU for 4 hrs and 24 hrs. Figures 4 and 6 listed genes with known functions based on One-way ANOVA clustering analysis from steady state total mRNA profiling from each cell line, respectively. A clear and unique expression pattern revealed a dynamic response to 5-FU treatment (Figure 4). There are 46 genes that were proved to be significant factors for acute 5-FU treatment (Genes listed in Additional file 3). They include CXCR4, FLJ23311, RSP26, AKAP12, ORC6L, PRPS1, DDX15, DKK4, and EIF4E. DKK4 functions as an agonist in the Wnt signaling pathway to regulate cell proliferation during brain development [30]. However, little is known about the function of DKK4 in cancer. This is the first report to show that DKK4 may play an important role in colorectal cancer possibly via the Wnt signaling pathway [31]. elf4E, one of the key translational initiation factors, has been shown to play key roles in cell cycle control and is an important marker for determining chemosensitivity [32,33]. We speculate that the acute induction of elf4E expression by 5-FU may be contributing to the resistance mechanism. Following 24 hr 5-FU exposure, nearly all of the acutely up-regulated genes had returned to baseline levels. However, additional genes such as SPRR1A, DDB2, and CDKN1A were only up-regulated after 24 hrs. These differential gene expression profiles suggest that single time point microarray data from clinical samples should be viewed with caution and not over-interpreted.

Interestingly, we discovered an additional 67 genes associated with for 5-FU treatment based on profiling of polysome associated mRNA transcripts (Figure 5). The induction of p53 will contribute a significant part for cell cycle arrest and apoptosis based on previous reports [34,35]. In addition, we also found genes such as, G-protein coupled receptor 132 (G2A), colon cancer antigen 33 (SDCCAG33), TSC2, and p34 (SEI1), were perturbed with 5-FU exposure. SEI1 is a CKD4-binding protein to regulate CKD4 activity [36] and was up-regulated following 5-FU exposure. G2A and SDCCAG33 were acutely up-regulated by 5-FU treatment whereas TSC2 was a delayed response gene affected by 5-FU. It has been recently shown that TSC2 is regulated at the translational level by hypoxia [37]. TSC2 is also involved in mTOR signaling pathway to regulate cell growth, proliferation, and cell death [38]. mTOR is one of the key pathways for translational regulation via elf4E [39]. These gene expression profiles may be helpful in elucidating additional resistance mechanisms related to 5-FU treatment.

We also determined the marker genes that were associated with TS independent cytotoxicity using HCT-C18 (TS-) cells treated with 5-FU in the presence of 10 μM thymidine. The main cytotoxicity caused by 5-FU was the inhibition of TS to block the sole de novo synthesis of thymidylate, an essential precursor for DNA biosynthesis. However, in the presence of 10 μM thymidine, the cytotoxicity caused by 5-FU in HCT-C18 (TS-) cells was mainly by the direct incorporation of 5-FU metabolite into DNA and RNA. The genes involved in TS independent cellular toxicity have not been systematically investigated in the past. In this study, we discovered a number of genes that were associated with TS independent cytotoxicity of 5-FU. Genes involved in DNA replication (NFI, CDC45L, TREX1, UBE1), DNA repair (TREX1, BTG2, XPC), DNA metabolism (TK1), RNA catabolism (ELAVL1), RNA processing (U3-55K), cell signaling (DDK2, CAPR1, TFAP2C), and apoptosis (NFkB1A, TRAF4) are on this list (Figure 6 and Additional file 5). It’s not surprising to see these genes on the list because direct incorporation of 5-FU metabolite to RNA and DNA is the major part of cytotoxicity via TS independent mechanism [40,41]. It has been shown that incorporation of 5-FU into RNA affects pre-mRNA splicing process [42]. This study provides the molecular targets and regulatory network that are potentially responsible for future enhancement of such cellular toxicity to tumors.
Genes associated with TS independent cytotoxicity caused by 5-FU were listed in Table 1. This list includes genes such as RRM2, SEI1, GPIBB, DDB2, and CDKN1A. A recent report showed that RRM2 (ribonucleotide reductase subunit M) was regulated at the translational level by the upstream AUGs [43]. This finding provides further validation to our approach as a systematic discovery platform for novel translational regulated genes. It has been shown recently that decreasing RRM2 level in HCT-116 (p53-/-) cells sensitize cells to DNA damaging agents and ribonucleotide reductase inhibitors [44]. It is clear that multiple genes will be responsible for determining the sensitivity in response to 5-FU other than TS itself [45]. These marker genes are important for us to understand the complex network regulated by TS and 5-FU. To go a step further, we also attempted to analyze them in the context of the regulatory network. As an example, we try to put these genes into a biological context via analyzing the cell cycle related genes affected by TS over-expression using pathway analysis, and the results are shown in Figure 7. It’s quite clear that TS over-expression decreases p53 level, thereby affecting its downstream genes such as p21 and the rest of the cell cycle related genes. This approach also helps to visualize the inter-relationship between TS and cell cycle related genes. We believe this is a starting point to show the power of systems biology approach for dissecting the mechanism and network of gene regulation.

Table 2: 2

| Genebank accession | Gene ID | P value | Biological function |
|--------------------|---------|---------|---------------------|
| S73288             | SPRR1A  | 0.0073  | 0.0078              |
| N001034            | RRM2    | 0.0143  | 0.0133              |
| N005554            | KRT6A, B, C | 0.0168 | 0.0030             |
| N003125            | SPRR1B  | 0.0211  | 0.0014              |
| N000407            | GPIBB   | 0.0229  | 0.0094              |
| N000389            | CDKN1A  | 0.0265  | 0.0137              |
| N013376            | SEI1    | 0.0274  | 0.0477              |
| N000422            | KRT17   | 0.0278  | 0.0044              |
| N002658            | PLAU    | 0.0362  | 0.0019              |
| N000107            | DDB2    | 0.0441  | 0.0049              |
| N006142            | SFN     | 0.0482  | 0.0115              |

Overlapping genes between HCT-C18 (TS+) and HCT-C18 (TS-) cells affected by 5-FU treatment.

Conclusion

In conclusion, the studies presented here provide a comprehensive expression profile of transcriptional and post-transcriptional levels affected by over-expression of TS protein and 5-FU treatment. This work expands our current understanding of the complex networks regulated by TS protein. In addition, our study also discovered candidate genes associated with TS independent cellular cytotoxicity. In particular, the newly discovered post-transcriptionally regulated genes by TS and 5-FU will be good candidates for further investigating the molecular and cellular mechanism of such regulation. This study further demonstrates the importance of understanding translational control in a global context in response to genotoxic stresses such as exposure to chemotherapeutic agents.

Methods

Cell culture

The human colon cancer cell lines, HCT-C18 (TS-) and HCT-C18 (TS+), have been previously described [16,20]. HCT-C18 (TS-) cells were maintained in 75-cm² plastic tissue culture flasks with growth medium consisting of RPMI 1640 medium containing 10% fetal bovine serum and supplemented with 10 µM thymidine (Sigma, MO) at 37°C and 5% CO₂. The HCT-C18 (TS+) cells were growing in the same medium without thymidine.

Western immunoblot analysis

Equal amounts of protein (25 µg) from each cell line were resolved by SDS/PAGE on 10% gels by the method of Laemmli [46]. Proteins were probed with mouse anti-TS monoclonal antibody (1:5,000 dilution) (Zymed Laboratories, CA), or anti-p53 mouse monoclonal antibody
(1:1,000 dilution) (Santa Cruz, CA), anti-α-tubulin mouse monoclonal antibody (1:1,000 dilution) (Santa Cruz, CA) followed by secondary antibody (Bio-Rad, CA). Proteins were visualized with a chemiluminescence detection system using the Super Signal substrate (Pierce, IL).

**Isolation of polysome associated mRNA transcripts**

For preparation of cytoplasmic extracts, cells from three 175 cm² tissue culture plates (60% confluency) were treated with 100 µg/ml cycloheximide (Sigma, MO) for 5 min at 37°C, washed with ice-cold phosphate-buffered saline containing cycloheximide (100 µg/ml) and harvested by scraping [47]. The fresh cells were pelleted by centrifugation, swollen for 2 min in 375 µl of low salt buffer (LSB) (20 mM Tris pH 7.5, 10 mM NaCl and 3 mM MgCl₂) containing 1 mM dithiothreitol and 50 U recombinant RNAsin (Promega, WI) and lysed by addition of 125 µl of lysis buffer (1× LSB containing 0.2 M sucrose and 1.2% Triton X-100) (Sigma) followed by vortexing. The nuclei were pelleted by centrifugation in a microcentrifuge at 13,000 rpm for 2 min. The supernatant (cytoplasmic extract) was transferred to a new 1.5 ml tube on ice. Cytoplasmic extracts were carefully layered over 0.5–1.5 M linear sucrose gradients (in LSB) and centrifuged at 45,000 rpm in a Beckman SW40 rotor for 90 min at 4°C. Gradients were fractionated using a pipette and the absorbance at 260 nm was measured for each fraction by UV spectrometry. Fractions 6 to 13 containing polysomes based on the positive absorbance profiles at 260 nm were pooled for RNA isolation.

**Sample preparation, array hybridization and gene expression analysis**

CodeLink UniSet Human 20 K Oligo Bioarray (Amersham Biosciences, NJ), containing approximately 20,289 genes probes, was used to generate gene expression profiles of both steady state total mRNAs and actively translated mRNAs isolated from control HCT-C18 (TS-) and HCT-C18 (TS+) cells and cells treated with 10 µM 5-FU (n = 3). Steady state total mRNAs from untreated HCT-C18 TS (-) and HCT-C18 TS (+) cells or after treatment with 10 µM 5FU for 4 and 24 hrs were isolated using Trizol Reagent (n = 3) (Invitrogen, CA). The corresponding polysomal fractions from each sample were pooled together and actively translated mRNAs were isolated using Trizol LS Reagent (Invitrogen, CA). All reagents were provided in the CodeLink expression assay kit (Amersham Biosciences, NJ), except where noted. cRNA synthesis was performed as per manufacturer’s instructions. Using 2 µg of total RNA, first-strand cDNA was generated by reverse transcriptase and a T7 primer. Subsequently, second-strand cDNA was produced using DNA polymerase I and RNase H. The resulting double-stranded cDNA was purified on a QIAquick column (Qiagen, CA) and cRNA was generated via an in vitro transcription reaction using T7 RNA polymerase and biotin-11-UTP (Perkin-Elmer, MA) at 37°C for 14 hrs. cRNA was purified on an RNeasy column (Qiagen, CA), quantified by UV spectrophotometry, and 10 µg biotin-labelled cRNA was then fragmented by heating at 94°C for 20 min in the presence of magnesium buffer. The fragmented cRNA was hybridized overnight at 37°C in hybridization buffer to a UniSet Human 20 K Bioarray in a shaking incubator at 300 rpm.

After hybridization, the arrays were washed in 0.75× TNT buffer [1× TNT: 0.1 M Tris-HCl (pH 7.6), 0.15 M NaCl, and 0.05% Tween 20] at 46°C for 1 hr followed by incubation with Cy5-streptavidin (Amersham Biosciences, NJ) at room temperature for 30 minutes in the dark. Arrays were then washed in 1× TNT four times for 5 min each. The slides were then dried by centrifugation and kept in the dark until scanning. Images were captured on an Axon GenePix 4200 A scanner. The resulting image was quantified and the intensity of each spot divided by the median spot intensity to provide a scaled and comparable number across multiple arrays. Bacterial spots provide both positive and negative controls. After dot grid and QC, CodeLink software generates export files for analysis by GeneSpring software 7.2 (Aglient, CA).

Gene expression analysis was carried out on GeneSpring software version 7.2, which allows multifilter comparisons using data from different experiments to perform the normalization, generation of restriction lists and the functional classification of the differentially expressed genes. Under Cross-Gene Error Model, normalization was applied in two steps: (a) "per chip normalization" in which each measurement was divided by the 50th percentile of all measurements in its array; and (b) "per gene normalization" in which all the samples were normalized against the specific samples (controls). Then data were filtered by flags and 4-fold change. The expression profiles of the different groups were compared using One-way ANOVA with cut-off p < 0.05. Comparisons of gene lists across different groups were performed using Venn Diagrams and clustering was performed with the Condition Tree algorithm. In addition, the Gene Ontology groupings and Gen Maps 2.0 program were used in conjunction with GeneSpring to identify pathways and functional groups of genes.

**Authors’ contributions**

YX performed the tissue culture and microarray gene expression profiling, data analysis and confirmation. GN performed the Western immunoblot analysis and assistance to the project. JS provided the critical review and scientific input of the manuscript. EC provided the cell lines and the critical scientific design and review of the project. JJ conceived, designed and led the project. All authors read and approved the manuscript.
Additional material

Additional File 1
Gene expression affected by TS over-expression based on profiling steady state mRNAs in HCT-C18 (TS-) and HCT-C18 (TS+) cells. This file contains the global gene expression profile of HCT-C18 (TS-) and HCT-C18 (TS+) cells based on steady state total mRNA expression using human high density CodeLink oligo array (20 K). Over 149 genes were changed in their expression in response to TS expression (n = 3, p < 0.05 and 4-fold cut-off).

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Additional File 2
Gene expression affected by TS over-expression based on profiling polypeptide associated mRNAs in HCT-C18 (TS-) and HCT-C18 (TS+) cells. This file contains the gene list of potential novel genes controlled at the post-transcriptionally by TS protein. Polypeptide associated mRNAs from both HCT-C18 (TS-) and HCT-C18 (TS+) cells were isolated and expression analysis was performed using human high density CodeLink oligo array (20 K). Over 149 genes were affected in TS overexpressing HCT-C18 (TS+) cells (n = 3, p < 0.05 with 4-fold cut-off).

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Additional File 3
Effect of 5-FU treatment on steady state mRNAs expression in HCT-C18 (TS+) cells. This file contains a gene list associated with both acute and delayed response genes following 5-FU treatment in HCT-C18 (TS+) cells by gene expression profiling analysis using steady state total mRNAs isolated from control and 5-FU treated samples at 4 hrs and 24 hrs time points. The expression analysis reveals that over 46 genes were affected by 5-FU treatment by One-way ANOVA analysis (n = 3, p < 0.05 with 4-fold cut-off). The clustering analysis is shown in Figure 4.

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Additional File 4
Effect of 5-FU treatment on polypeptide associated mRNAs expression in HCT-C18 (TS+) cells. This file contains both acute and delayed response genes that were affected at post-transcriptional level by 5-FU treatment. Polypeptide associated mRNAs were isolated from control, and 5-FU treated samples at 4 hrs and 24 hrs time points. Gene expression analysis via microarray revealed that over 67 genes were affected in response to 5-FU (n = 3, p < 0.05 with 4-fold cut off). The clustering analyses are shown in Figure 5.

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Additional File 5
Effect of 5-FU treatment on steady state mRNAs expression in HCT-C18 (TS-) cells. This file contains gene list based on gene expression profiles generated from these samples. These genes are mainly associated with TS independent toxicity in response to 5-FU by direct incorporation to RNA and DNA. The clustering analysis is shown in Figure 6. This file contains a gene list of 185 genes (n = 3, p < 0.05 with 4-fold cut-off) of 5-FU exposure in HCT-C18 (TS-) cells in the presence of thymidine.

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