

Expression of PER, CRY, and TIM genes for the pathological features of colorectal cancer patients

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Department of General Surgery, Department of Gastroenterology, The Second Affiliated Hospital of Anhui Medical University, Hefei, People's Republic of China **Abstract:** As typical clock gene machinery, period (*PER1*, A Q, and PERcryptochrome (CRY1 and CRY2), and timeless (TIM), could control protectation, co amage, key functions, such as recognition and repair of DN vsfunct RC). I could result in tumorigenesis of colorectal cancer his study, the expression levels of M in the PER1, PER2, and PER3, as well as CRY1 X2, and mor tissue and apparently nined and col a quantitative real-time polyhealthy mucosa from CRC patients were are merase chain reaction. Compared with the hear v mucosa from CRC patients, expression levels of PER1, PER2, PER3, and CRY2 in their tumo. ssue are much lower, while TIM level was Agnificant difference is the CRYI expression level. High levels of TIM mRNA were much prelent in the tund mucosa with proximal lymph nodes. CRC patients with lower expression of PE and PER3 in the tumor tissue showed significantly poorer survival rates. The abnormal expression vels of R and TIM genes in CRC tissue could be related to the genesis proce tumor, innuencing host-tumor interactions.

Keywords: colore all car cs., ser chronotherapy, period genes, cryptochrome genes, timeless general

trocking

The ofe-dependent pattern of variation exists in most biological processes and function of living organisms. The circadian rhythm is defined as the rhythmic patterns of oscillation with a period of ~24 hours. The circadian timing system, containing contral and peripheral oscillators, is responsible for the generation of the rhythmic variation. The central pacemaker and master oscillator, located at the suprachiasmatic nuclei of the brain, are entrained to the environmental light—dark cycle via photon, which is conveyed by the retinohypothalamic tract and synchronizes slave oscillators in peripheral tissues. 3,4

Up to now, several biological clock genes have been confirmed, such as ARNTL1, clock (circadian locomotor output cycles kaput), period 1, 2 and 3 (*PER1*–3), cryptochrome (*CRY1* and *CRY2*), timeless (*TIM*), timeless-interacting protein (TIPIN), and CSNK1E (encoding casein kinase 1-epsilon, CK1ε). The molecular components, which generate circadian rhythms of the circadian/biological clock, constitute a unique collaboration mechanism of genes and proteins via transcriptional, translational, and posttranslational procedures.

PER1–3, *CRY1* and *CRY2* can be transcriptionally activated by the CLOCK and ARNTL1, which are the basic helix–loop–helix/Period, Aryl-hydrocarbon-receptor, Single minded (PAS) transcription factors, heterodimerizing and binding to the elements of E-box enhancer in the gene promoters. In contrast, *CRY* and *PER* proteins form the repression complex, translocating back into the nucleus and interacting with



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CLOCK and ARNTL1, resulting in the loss of their activity.^{5,6} As a core circadian clock gene in *Drosophila melanogaster*, *TIM* is also retained in mammals. However, the influence of mammalian circadian system on clock function is not yet clear. By interacting with the components of the DNA replication system, *TIM* could regulate DNA replication processes, which are essential for ataxia telangiectasia and Rad3-related-checkpoint kinase 1 (ATR-Chk1)-mediated as well as ataxia telangiectasia mutated—checkpoint kinase 2 (ATMChk2)-mediated signaling and S-phase arrest.⁷ As more and more incidence of various cancers has been reported by epidemiological studies, the temporal organization variation in body, defined as the circadian disruption, is nowadays considered to be an important risk factor for cancer.^{8,9}

Approximately 5%-15% genome-wide mRNA expressions, including key cell cycle regulators, tumor suppressors genes, and oncogenes, are circadian rhythmic, which are driven by biological clock genes, whose expression regulates the timing of DNA repair, apoptosis, and cell proliferation. The cycle progression of the cells is tightly regulated by the circadian system, including the control of cell proliferation and apoptosis, as well as clocking the transcription and posttranslational modification for key proteins. The deregulated cell proliferation may occur in case of the circadian clo disruption, implicating in many types of cancers. 10,11 Over expression of PER1 in cancer cell lines of huma the reduction of colony formation and clonog and altered expression of transcriptional targeties MYC and p21. For the PER2, a mutation as bee √estigated al polyp in to accelerate the generation of in mice, and an increase in hypernasia an neoplasia under γ-radiation has been shown PER2-null m. 12,13 Based on nt mous models of breast cancer, the analysis of two diffe PER3 was putatively side d to be a tumor suppressor patien with estracen receptor-positive gene. Moreover d especially for those tumors treat with moxin without chaother breast cancer recurrence has been ated to the deletion and/or reduction in the observed to be gene.14 expression of PE.

In humans, deregalation or polymorphism of the *PER*, *CRY*, and *TIM* genes is associated with a number of neoplastic and hemolymphoprolipherative diseases. ^{15,16} Recently, there is a great deal of interest in the alteration of clock gene and clock-controlled gene expression in colorectal cancer (CRC) patients. ^{17,18} CRC accounts for \sim 10% among all kinds of cancers, becoming the third most common cancer and the fourth most common for death all over the world. ^{19,20} In this study, the expression levels of *PER* and *CRY* genes in human CRC tissues and matched normal mucosa were evaluated to explore

the relationship between their expressions in tissue with cancer and the clinical and pathological features of CRC patients.

Patients and methods

Patients

Primary tumor samples and matched normal tissues were obtained from 19 recently diagnosed CRC patients (13 men and six women) in our hospital who had had curative surgery. Clinical data, tumor characteristics, location, and staging of these patients are listed in Table 1. All tissue specimens

Table I Clinical and pathological for tires of color tal cancer patients (N=19)

patients (IN=19)			
Features of patients	n	%	Mfemale
Age, mean ± SD (years)	68.9±8.8		1.1±8.7/72.8±8.6
Sex			•
Male (M)/female (F			13/6
Tumor location			
Cecum	2	10.5	0/2
Ascending colon			0/0
Proy ar uravisverse		10.5	1/1
Digal transverse	2	10.5	1/1
Decending colon	1	5.3	1/0
Signid colon	11	57.9	10/1
Rectu sigmoid ion	1	5.3	0/1
listologic type			
sin-producing	15	78.9	11/4
adenocarcinoma			
Mucin-producing	4	21.1	2/2
adenocarcinoma			
rading			
GI-G2	17	89.5	13/4
G3-G4	2	10.5	0/2
Depth of tumor invasion	-	. 0.0	V/ -
TI-T2	3	15.8	2/1
T3–T4	16	84.2	11/5
Lymph node metastasis	10	01.2	11/3
N0	12	63.2	8/4
NI-N2	7	36.8	5/2
Metastasis	•	50.0	3/2
No	17	89.5	12/5
Yes	2	10.5	1/1
AJCC/TNM stage	2	10.5	1/1
	3	15.8	2/1
	9	47.4	6/3
 	5	26.3	4/1
IV	2	10.5	1/1
Vascular invasion	2	10.5	1/1
No No	12	63.2	6/6
Yes	7	36.8	7/0
	,	36.6	770
MSI status	,	217	2/2
Missing	6	31.6	3/3
MSI-H	3	15.8	1/2
MSI-L	4	21.1	2/2
MSS	6	31.6	3/3

Abbreviations: AJCC, American Joint Committee on Cancer; MSI, microsatellite instability; MSI-L, low microsatellite instability; MSI-H, high microsatellite instability; MSS, microsatellite stable; SD, standard deviation; TNM, tumor, node, metastasis.

were collected between 9 am and 5 pm in a same day (9 am to 11 am five samples, 11 am to 1 pm four samples, 1 pm to 3 pm six samples, and 3 pm to 5 pm four samples) and immediately put into liquid nitrogen and stored at -80°C. This study was approved by the Ethics Committee in the Second Affiliated Hospital of Anhui Medical University. All patients offered their informed written consent.

Extraction of RNA from fresh frozen tissue and synthesis of the first-strand cDNA

The total RNA from ~150 mg to 200 mg fresh frozen tissues was isolated by phenol extraction. The amount of extracted RNA was weighted by Nano Drop Spectrophotometer, while Agilent 2100 Bioanalyzer was applied for monitoring RNA integrity after subsequent digestion by DNaseI. Then, 1.0 μ g of total RNA was reverse transcribed by using the High-Capacity cDNA Archive Kit.

Quantitative real-time reverse transcriptasepolymerase chain reaction assay

The differential expressions of the genes in tumor tissue matched to normal mucosa of CRC samples were assessed by quantitative real-time reverse transcriptase-poly chain reaction (qRT-PCR) assay. Human Quan Primers Assay was applied for the determination of Ph PER2, PER3, CRY1, CRY2, and TIM. Al ART-CRs w a 25 µL final volume and three replices were derformed on a QuantiFast SYBR Green PC kit an in an ABI PRISM 7700 Sequence Detect System up r conditions at 50°C for 2 minutes and C to 10 minutes as well as 40 cycles at 95°C for 1 seconds an 60°C for 1 minute. The threshold cycle (2,) values were accorded. Expression levels of the target ones we normalized by glutaraldehyde-3-phosphate dehydro ase how keeping control gene,21 and the rative mount of RNA in each target gene company de de lutaral de la vide nated by average $2^{-\Delta\Delta Ct}$ method.²² The results nase was are shown in Table 2.

Microsatellite instability

The Bethesda panel of microsatellite (BAT25, BAT26, D5S346, D17S250, and D2S123) was applied for the microsatellite instability (MSI) analysis, which was evaluated by means of multiplex PCR and polyacrylamide gel electrophoresis analysis. Then, the tumors were accordingly classified into the following categories: microsatellite stable, low microsatellite instability (MSI-L), and high microsatellite instability (MSI-H).²³

Statistical analysis

The SPSS Statistical Package was used for the statistical analyses. Comparing with gene expression levels of the normal mucosa, those from the adjacent CRC tissues were calculated by the formula $2^{-\Delta\Delta Ct}$, with the reported values of median, 25th percentile (Q1), and 75th percentile (Q3). The normal distribution of the continuous variables was verified by the Shapiro-Wilk test and the one-sample Kolmogorov-Smirnov test. By addressing with a nonparametric Wilcoxon signed-rank test, the statistical significance of the up- or downregulation for non-normal $2^{-\Delta\Delta Ct}$ transformation was assessed the differences among the groups for normally distribute variables were compared by Student's t-test or translysis variant, while those for non-normally di ariable ere compared by the Mann-Whitney nk am test or the Kruskal–Wallis rank sum ter The cornection beween mRNA expression levels and and patho cal features was evaluated by Spearman's test, and the survival rates were calculated by the Meier metho via censored data analysis. A P-value 0.05 was considered as statistically significant.

sults and discussion

The expression levels of the core clock genes (*PER1*, *PER2*, *CRY1*, *CRY2*, and *TIM*) from 19 paired normal and colon cancer tissues were evaluated, in order to identify those differentially expressed in primary CRC. Figure 1 presents the relative expression levels of clock genes in CRC tissue samples according to the data in Table 2. Comparing with normal tissue after normalizing to 1, the following five genes in the tumor samples were down-regulated: *PER1* (median =0.42, Q1–Q3=0.24–0.85, *P*=0.002), *PER2* (median =0.51, Q1–Q3=0.34–0.88, *P*=0.011), *PER3* (median =0.36, Q1–Q3=0.15–0.58, *P*=0.003), and *CRY2* (median =0.55, Q1–Q3=0.27–0.90, *P*=0.012). The *CRY1* gene did not show different expression (median =0.93, Q1–Q3=0.65–1.48, *P*=0.600). It is obviously shown in Figure 1 that *TIM* was upregulated (median =1.22, Q1–Q3=0.92–1.63, *P*=0.044).

The association between gene expression levels (PER, CRY, and TIM) and clinical and pathological features, such as patient age and sex, tumor location and stage, and MSI status, is listed in Table 3. A significant association was observed for the CRYI and TIM genes. In particular, lower expression level of CRYI in the tumor mucosa was found in the 68–75-year-old subjects (P=0.026) and female patients (P=0.005), whereas higher expression level of CRYI in the tumor mucosa was found in cancers located in the distal colorectal segments (P=0.015), which was confirmed by Spearman's correlation (r=0.521, P=0.02, slope =0.519) (Figure 2).

Table 2 Relative and normalized expression levels of clock genes in colorectal cancer and mRNAs were expressed as mean $2^{-\Delta\Delta Ct}$ and SD

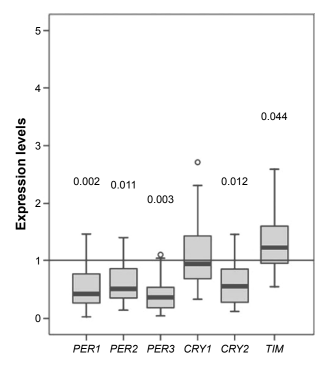
	PERI		PER2		PER3		
	Mean 2 ^{-ΔΔCt}	SD	Mean 2 ^{-∆∆Ct}	SD	Mean 2 ^{-ΔΔCt}	SD	
Sample PS I	0.148593313	0.003329629	0.140636383	0.001378578	0.041369743	0.005547747	
Sample PS2	0.686256067	0.059030876	0.837356315	0.08115278	0.44794733	0.084363612	
Sample PS3	0.293233832	0.021471752	0.360234712	0.024152627	0.579212347	0.054530476	
Sample PS4	0.517660447	0.046787661	0.483264041	0.01729745	0.440295024	0.014052806	
Sample PS5	0.615409643	0.030150945	0.562359068	0.021690964	0.49138551	0.066455953	
Sample PS6	0.405616491	0.020982013	0.189677498	0.018320315	0.15215594	0.006157781	
Sample PS7	0.062033048	0.006675992	0.243125077	0.01070676	0.06347467	0.008683778	
Sample PS8	0.421673014	0.045380419	0.337789071	0.014045216	0.36097168	0.058131258	
Sample PS9	0.676397751	0.031561495	0.50870249	0.034245053	0.077369856	0.006263042	
Sample PS10	0.238319619	0.018670048	0.936377508	0.047961853	0.249940	0.041435853	
Sample PS11	0.846401991	0.205716657	0.292537606	0.014757011	0.931/ 1729	378102	
Sample PS12	0.047178752	0.003438641	0.453037804	0.021973839	0.23-, 5017	01149226	
Sample PS13	0.026752629	0.003197484	0.664332366	0.010683014	.05447- 7	08898889	
Sample PS14	0.300053079	0.021065332	0.448304223	0.021745458	0.21043458	0.017448073	
Sample PS15	0.910213643	0.231713871	0.829084909	0.178008842	0.25 88173	0.022470912	
Sample PS16	0.288833852	0.049316316	0.876888235	0.024310168	41743796	0.033459043	
Sample PS17	1.333893837	0.057954476	1.411384224	0.07352 /5	2.625848	0.18764957	
Sample PS18	1.471514504	0.074230307	3.407718958	0.102 2464	340 891	0.128939674	
Sample PS19	1.370382169	0.236537711	1.370840431	0 46873 8	1.0 499345	0.071021113	
	CRYI		CRY2		TIM		
	Mean 2 ^{-ΔΔCt}	SD	Mean 2 ^{-∆∆Ct}	JD	Mean 2 ^{-ΔΔCt}	SD	
Sample PS I	0.649289748	0.059934764	0.117900145	0.00664726	0.545659652	0.075202149	
Sample PS2	0.715427505	0.034935461	0.898393284	0.1022	0.606442156	0.085315813	
Sample PS3	0.389462167	0.051389512	0.441259597	0.0578	0.922434233	0.064052995	
Sample PS4	0.857378971	0.029410076	0714777702	0.027 38623	0.582847524	0.062303115	
Sample PS5	0.934737143	0.281852036	0. 850	0.037949085	0.866423092	0.04148866	
Sample PS6	0.328891017	0.038016805	0.2 24570	0.010453803	0.97474993	0.014107983	
Sample PS7	0.5833602	0.03740551	0.210 17 24	0.011059941	2.590559252	0.02738315	
Sample PS8	0.813085982	0.0615856	0.2330 503	0.032872414	1.572210757	0.028702739	
Sample PS9	0.555555039	0.03857 _37	0.28047 47	0.040498653	1.219881211	0.04902701	
Sample PS10	0.722475552	0.12230 53	19345	0.063735334	1.585920607	0.056514161	
Sample PS10 Sample PS11			0.268831945	0.063735334 0.020482531	1.585920607 1.267166976		
•	0.722475552	0.12230 53				0.035374096	
Sample PS11	0.722475552 2.307305573	0.12236 53 0 .1873396	0.268831945	0.020482531	1.267166976	0.035374096	
Sample PS11 Sample PS12	0.722475552 2.307305573 2.263095958	0.12230 53 01.873390 169538304	0.268831945 0.769572585	0.02048253 I 0.053583003	1.267166976 2.32439189	0.035374096 0.274518286	
Sample PS11 Sample PS12 Sample PS13	0.722475552 2.307305573 2.263095958 1.900991943	0.12236 53 0.4873396 169538304 6.1072431	0.268831945 0.769572585 0.551525369	0.020482531 0.053583003 0.034847113	1.267166976 2.32439189 1.235656923	0.035374096 0.274518286 0.013061341 0.029969761	
Sample PS11 Sample PS12 Sample PS13 Sample PS14	0.722475552 2.307305573 2.263095958 1.900991943 1.109515431	0.12236 53 0.1873398 1169538304 0.1072431 0.021 33253	0.268831945 0.769572585 0.551525369 0.434306719	0.020482531 0.053583003 0.034847113 0.020559688	1.267166976 2.32439189 1.235656923 1.032233245	0.035374096 0.274518286 0.013061341 0.029969761 0.085031924	
Sample PS11 Sample PS12 Sample PS13 Sample PS14 Sample PS15	0.722475552 2.307305573 2.263095958 1.900991943 1.109515431	0.12236 53 9 1873396 1169538304 0 1072431 0.02 33253 0.07970 44	0.268831945 0.769572585 0.551525369 0.434306719 0.800736942	0.020482531 0.053583003 0.034847113 0.020559688 0.072909877	1.267166976 2.32439189 1.235656923 1.032233245 1.632193291	0.035374096 0.274518286 0.013061341	
Sample PS11 Sample PS12 Sample PS13 Sample PS14 Sample PS15 Sample PS16	0.722475552 2.307305573 2.263095958 1.900991943 1.109515431 1.4791219 0.2888 852	0.12236.53 0.1873396 169538304 0.1072431 0.024.33253 0.07976.44 0.049316316	0.268831945 0.769572585 0.551525369 0.434306719 0.800736942 0.876888235	0.020482531 0.053583003 0.034847113 0.020559688 0.072909877 0.024310168	1.267166976 2.32439189 1.235656923 1.032233245 1.632193291 0.441743796	0.035374096 0.274518286 0.013061341 0.029969761 0.085031924 0.033459043	

M, timeles PER, period, , cryptochrome; SD, standard deviation. **Abbreviations**

A significant assocition could be observed between high *TIM* mRNA expression level in tumor mucosa and stages III-IV (P=0.005), involving lymph nodes (P=0.005), in particular, of proximal lymph nodes (P=0.013), and DNA mismatch repair proficiency and MSI (P=0.015) (Table 3).

CRC patients with lower expression of *PER1* (*P*=0.010) and PER3 (P=0.010) in the tumor tissue showed significantly poorer survival rates, according to the Kaplan-Meier method for the analysis of censored data. No statistically significant decrease in survival could be evidenced in patients with low expression level of PER2 (P=0.143), CRY1 (P=0.143), CRY2 (P=0.236), and TIM (P=0.491) in the tumor tissue (Figure 3).

The core clock genes drive and activate downstream clock-controlled genes and the control of tissue/organ function.^{24,25} In addition, altered expression levels of PER and CRY genes have been evidenced in intestinal biopsies of diseased intestinal segments of patients affected by ulcerative colitis and Crohn's disease, and inflammation is considered to favor the development of neoplastic disease. Considering the reported involvement of the circadian clock in several cancers, qRT-PCR was applied to examine the expressions



 $\textbf{Figure I} \ \, \textbf{Expression of core clock genes in colorectal cancer tissue.}$

Notes: Box and whisker plots of the expression of the core clock genes in CRC tissue as analyzed by qRT-PCR and compared with matched normal tissue, with the expression in GAPDH used as calibrator. For each gene, a box plot shows the median, quartiles, minimum, and maximum values, as well as outliers. Boxes represent the interquartile range (IQR) and the horizontal bar the median relative expression. Expression values that do not fall within 1.5× IQR are outlied the indicated by circles where appropriate.

Abbreviations: CRC, colorectal cancer; qRT-PCR, quantitative real-time transcriptase-polymerase chain reaction; GAPDH, glutaraldehyde-3-phos ard dehydrogenase; TIM, timeless; PER, period; CRY, cryptochron

of PER1, PER2, PER3, CRY1, CI I in CRC and 2, and matched normal colorectal s. The exp of some clock genes in tuntor tiss was found to be significantly decreased, and the case of \$\infty R1, PER2, PER3, and CRY2, whereas Lat of TM (P=0.044) was higher, and poorer survival ra. sociated with lower expression nd Pb. in the tracor tissue in a statistically levels of PEP significa way.

Otheresults from agreement with previous reports describing terations in the expression of clock genes in CRC. In particular, the decreased expression of PER genes seems to be most relevant for the process of oncogenesis and tumor progression. PER1 and PER2 are related to the pathways for ATM-Chk1/Chk2 DNA damage response, β -catenin modulate, and proliferation of colon and noncolon cancer cells. In contrast, clock function may be altered by intestinal tumorigenesis, leading to increase in β -catenin-destabilizing PER2 protein. PER1 could directly interact with ATM in vitro as a response to radiation. In mice, PER2 mutation would result in change in the temporal gene expression for regulation of the cell cycle and tumor suppression

(C-MYC, CYCLIN A, CYCLIN D1, GADD45A, and MDM-2), DNA-damage response deregulation, accelerating intestinal polyp formation in APC^{Min/+} mice, and increase in neoplastic growth. ^{26,27} We found no statistically significant difference in the *CRYI* expression level between CRC and matched normal tissue. CK1ε phosphorylates *PER* and *CRY* proteins and β-catenin, thereby facilitating their ubiquitination and proteasomal degradation. In vitro and in vivo studies strongly confirm that CKIε is a key factor in the early stages of tumorigenesis, predisposing to colon cancer. As tumor cells have more dependence on the factor of CK1ε than normal ones, a specific kickse inhibite to CK1ε could induce tumor cell-selective cythoxicity. ^{28,29}

A correlation between Now explain of ER1 gene and liver metastasis and ne relationship een high expression of *PER2* gane ificantly better outcomes were revious dy. 30 Leaur study, CRY1 mRNA reported in a C were significantly related expressio le s in tissues the lowest levels detected in the age to patient age (w ex (with the lowest levels detected in emale patients), and cancer location (with the highest levels ors located in the distal colon). etected in tu

The close gene machinery, which controls the system functions of hepatic, intestinal, renal detoxification, and a biotic detoxification, exhibits circadian variation of activity, determining time-dependent toxicity of xenobiotics and drugs. Time-of-day-dependent variation in drug toxicity, metabolism, and effectiveness provides the basis for the specific 24-hour period timing of drug administration during the chronomodulated cancer chemotherapy of advanced-stage CRC.

For example, diurnally active patients deliver oxaliplatin in the afternoon and fluorouracil and leucovorin at night.³¹ For the sex dependency, female CRC patients have been reported to have shorter survival and greater toxicities, according to European Organization for Research and Treatment of Cancer (EORTC) Chronotherapy Group trial.³² The male and female mammals (including humans) have different xenobiotic detoxification and metabolic pathways. The change of sexual dimorphism in hepatic drug metabolism could be observed in double mutant CRY1^{-/-} CRY2^{-/-} male mice. Once the CRY genes are inactivated, the male and female mice have similar expression levels of sex-specific liver products, such as several cytochrome P450 enzymes.33 By transferring this evidence to humans, it could be suggested that the decreased expression level of the CRY1 gene in female CRC patients might lead to the different median survival and the increase in the toxicity after the administration of chronic-modulated chemotherapy.

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Table 3 Relationship between expression levels of CRY1 and TIM genes and clinicopathological features of CRC patients

Category	n	CRYI				TIM			
		Median	QI	Q3	P-value	Median	QI	Q3	P-value
Age (years)									
<60	4	1.05	0.75	1.28	0.026	0.99	0.76	1.12	0.336
60–67	6	1.87	1.40	2.31		1.70	0.61	2.32	
68–75	3	0.56	0.39	0.72		1.57	1.12	1.92	
>75	6	0.83	0.58	1.11		1.13	0.92	1.59	
Sex									
Male	13	1.32	0.86	1.90	0.005	1.24	0.97	1.63	0.430
Female	6	0.60	0.39	0.72		1.07	0.87	1.57	
Tumor location									
Proximal colon	2	0.64	0.56	0.72	0.02	1.52	1.12	92	0.711
Transverse colon	4	0.62	0.49	0.68		1.25	د ۲	l.c	
Distal colon	13	1.32	0.93	1.90		1.22	1.97	1.59	
Grading						_			
G1/G2	17	1.02	0.72	1.48	0.201	1.17	0.91	1.59	0.273
G3/G4	2	0.47	0.39	0.56		1,9	1.57	V	
AJCC/TNM stage									
1	3	0.72	0.33	1.11	0.560	l.	0.92	1.59	0.028
II	9	0.93	18.0	1.32		0.97	0.6	1.12	
III	5	1.40	0.58	2.26		1.92	ر 33	2.32	
IV	2	1.15	0.39	1.90		1.40	1.22	1.57	
Histologic type									
Nonmucin-producing adenocarcinoma	15	1.11	0.72	1.48	317		0.97	1.63	0.841
Mucin-producing adenocarcinoma	4	0.68	0.52	I	•	1.25	0.73	2.08	
Lymph node positive									
No positivity	12	0.90	0.72	3	0.083	0.99	0.74	1.18	0.013
Proximal lymph node	5	1.40	0.58	2.2		1.92	1.63	2.32	
Distal lymph node	2	1.15		1.90		1.40	1.22	1.57	
Lymph node involvement									
N0	12	0.90	0.	.26	0.672	0.99	0.74	1.18	0.005
NI-N2	7	1.40	0.5	2.26	•	1.63	1.27	2.32	
Vascular invasion									
No	12	J.87	0.62	1.26	0.150	1.07	0.92	1.77	0.799
Yes	7	1.90		2.31		1.24	0.90	1.57	
MSI status									
Missing	6								
MSI-H		0.56	0.39	1.90	0.735	1.57	1.22	1.92	0.015
MSI-L	4	0.96	0.71	0.91		1.61	1.43	1.88	
MSS	6	0.90	0.72	1.24		1.02	0.97	1.12	

Note: P-value: Spearman tests.

Abbreviations: TIM, timeless (RC, color al cancer; AJCC, American Joint Committee on Cancer; MSI, microsatellite instability; MSI-L, low microsatellite instability; MSI-H, high microsatellite instability instability; MSI-H, high microsatellite instability instability; MSI-L, low microsatellite instability; MSI-L, low microsatell

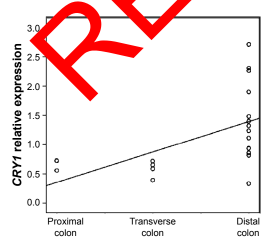


Figure 2 The correlation between *CRY1* expression levels in tumor tissue and cancer location, evaluated by Spearman's test (*r*=0.521, *P*=0.02, slope =0.519).

Comparing with the normal mucosa, the *TIM* mRNA levels observed in the CRC tissue were significantly related to American Joint Committee on Cancer (AJCC)/TNM stage (highest levels in TNM stages III–IV), lymph node involvement (highest levels in the case of positive lymph nodes, especially proximal lymph node involvement), and MSI (highest levels in MSI-H and MSI-L). Most of the cytotoxic anticancer drugs would damage DNA and activate DNA checkpoints for approving the attempted DNA repair, which is important to the survival of the cells. However, the cytotoxicity of anticancer drugs may be reduced. For ATM-dependent Chk2-mediated signaling of doxorubicin-induced DNA double-strand breaks, *TIM* plays an essential role. Moreover, the arresting of

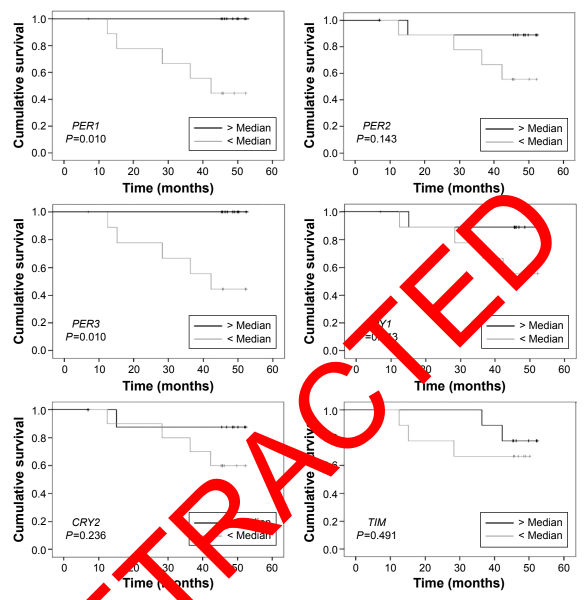


Figure 3 Cumulative survival patients according to the pression levels of PER1, PER2, PER3, CRY1, CRY2, and TIM in tumor tissue of CRC patients.

Notes: Patients with lower pression of PER1 and PER3 showed significantly poorer survival rates (P=0.010 PER1 and P=0.010 PER3). No statistically significant difference was found in cumulative solvial rate of CRC patients according to the tumor tissue mRNA levels of PER2 (P=0.143), CRY1 (P=0.143), CRY2 (P=0.236), and TIM (P=0.491). P-value: log-rank test by Kapin Market method.

Abbreviations: neless; C, colore cancer; PER, period; CRY, cryptochrome.

doxor icin-in (al. G(2)/M cell cycle would be significantly attended by downregulation of *TIM* siRNA, sensitizing cance cells to doxorubicin-induced cytotoxicity. Hence, the variation in drug sensitivity could be predicted by *TIM* mutation in human cancers. In order to enhance the cytotoxic effectiveness of chemotherapeutic drugs for activating DNA response pathways in cancer cells, the *TIM* inhibition becomes a potential novel target for anticancer drugs.^{34,35} In our study, *TIM* expression was also significantly associated with MSI. Approximately 15% of CRCs are diagnosed by defects in mismatch repair system of DNA, leading to MSI and generating many substitution, insertion, deletion, and mutations. As mainly targeting to

the microsatellite sequences, these mutations could lead to reading frame variation, further resulting in truncation or alterations in protein. The mRNA expressions with such frameshift mutations could be decreased with the presence of premature stop codons, leading to some mutant mRNA degradation via the nonsense-mediated decay pathway.^{7,36} As better outcome with irinotecan-containing regimens was shown in MSI-H tumors than with 5-fluorouracil-containing treatments, the MSI status of CRC patients could affect the response to adjuvant chemotherapy.³⁷ In MSI-H and MSI-L patients, the increase in the *TIM* expressions may be associated with the tumorigenesis process in CRC and reduction in the response to adjuvant chemotherapy.

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Conclusion

In conclusion, there are differences in the expression levels of PER, CRY, and TIM genes in CRC tissues compared with matched normal ones, and the altered expression might influence the process of carcinogenesis and various aspects of host-tumor characteristics and interactions.

Disclosure

The authors report no conflicts of interest in this work.

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