Exosome component 1 cleaves single-stranded DNA and sensitizes kidney renal clear cell carcinoma cells to poly(ADP-ribose) polymerase inhibitor

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Abstract

Targeting DNA repair pathway offers an important therapeutic strategy for cancers. However, the failure of DNA repair inhibitors to markedly benefit patients necessitates the development of new strategies. Here, we show that exosome component 1 (EXOSC1) promotes DNA damages and sensitizes kidney renal clear cell carcinoma (KIRC) cells to DNA repair inhibitor. Considering that endogenous source of mutation (ESM) constantly assaul...
**Introduction**

DNA damages and subsequent mutations are central to development, progression, and treatment of nearly all cancers (Brown et al., 2017; Farmer et al., 2005; Jeggo et al., 2016; Pearl et al., 2015; Roos et al., 2016). Cancer cells frequently decreases DNA repair pathways and increases endogenous sources of mutation (ESM) to drive mutations (Brown et al., 2017; Farmer et al., 2005; Jeggo et al., 2016; Pearl et al., 2015; Roos et al., 2016). Hence, cancer cells are often more reliant on a subset of DNA repair pathway(s) to survive DNA damages. While, targeting critical DNA repair members, such as poly (ADP-ribose) polymerases (PARPs) (Brown et al., 2017; Tubbs and Nussenzweig, 2017), offers a therapeutic strategy for cancers (Tutt et al., 2010). Inhibition of PARPs by small-molecule compounds disrupts the ability of cancer cells to survive ongoing DNA damage and results in cell cycle arrest and/or cell death (Lord and Ashworth, 2012). However, failure of PARP inhibitors (PARPis) to markedly benefit patients suggests the necessity for developing new strategies. Due to the central role of ESM in ongoing DNA damages, there is a need for the identification and understanding of ESM.

ESM constantly assaults genomic DNA and almost inevitably leads to mutations (Figure 1A) (Jeggo et al., 2015; Roos et al., 2015). However, most of ESM studies were focused on the deamination. The significance of deamination as an ESM is supported mainly by two observations: (1) Transitions show higher frequency than expected by chance, although there are twice as many possible transversions. Nucleotide substitutions consist of two types: transition and transversion (Alexandrov et al., 2013a; Burgess, 2019; Petljak et al., 2019). Transition is a substitution in which one base is replaced by another of the same class (purine or pyrimidine), while transversion is a substitution in which a purine is replaced with a pyrimidine or vice versa (Figure 1B). (2) C>T transitions at methylated cytosine in CG base pairs display a higher frequency than expected (Alexandrov et al., 2013a; Alexandrov et al., 2013b; Burgess, 2019; Hutchinson, 2013; Petljak et al., 2019). Therefore, activation-induced cytidine deaminase (AID) (Barlow et al., 2013; Greaves, 2018; Petersen-Mahrt et al., 2002) and apolipoprotein B mRNA editing enzyme catalytic (APOBEC) family (Alexandrov et al., 2013b; Buisson et al., 2019; Hutchinson, 2013; McGranahan et al., 2017; Robertson et al., 2017) catalyzing the deamination of C were identified as ESMs. Unfortunately, some cancers such as kidney renal clear cell carcinoma (KIRC) show the low mutation proportion at CG base pairs and low APOBEC expression.
Due to advances in sequencing technology and the great efforts of The Cancer Genome Atlas (TCGA), it is now possible to explore the statistical relationship between mutations and the expression of individual genes in multiple cancer types (Weinstein et al., 2013). The majority of patients included in the TCGA database are accompanied by data regarding both mutations and genome-wide expression of individual genes (Weinstein et al., 2013). Because that DNA damages often comprise a major source of mutation, the relativity between DNA damage and mutation allows quantitative analyses of mutation to be taken as a proxy of DNA damage (Brown et al., 2017; Jeggo et al., 2016; Pearl et al., 2015). Furthermore, the gene-specific correlation between mRNA and protein levels allows quantitative analyses of individual gene expression as an indicator for the corresponding protein (Peng et al., 2015; Uhlen et al., 2017; Zhang et al., 2017). Hence, analyses of the cancer cohort may identify candidate ESMs (Tubbs and Nussenzweig, 2017).

The exosome is an evolutionarily conserved multiprotein complex formed by exosome components (EXOSCs) (Bousquet-Antonelli et al., 2000; Brown et al., 2000; Tomecki et al., 2010). In eukaryotes, the exosome complex has a “ring complex” (EXOSC4–EXOSC9) and a “cap” structure (EXOSC1–EXOSC3) (Figure 1C). The human exosome complex may also contain two additional subunits, EXOSC10 and EXOSC11 (Jonathan et al., 2006; Lorentzen et al., 2008; Quansheng et al., 2006; Wasmuth et al., 2014), which provide 3’ to 5’ exo- and/or endoribonuclease activities (Januszyk and Lima, 2014; Kilchert et al., 2016). The exosome is well known to degrade RNA (Januszyk and Lima, 2014; Kilchert et al., 2016; Ulmke et al., 2021). Hence, the exosome was reported to protect cells from genomic instability via degrading the DNA/RNA hybrids and restricting DNA strand mutational asymmetry (Lim et al., 2017; Pefanis and Basu, 2015; Pefanis et al., 2015). Interestingly, the cap unit EXOSC2 stably associated with the exosome complex, while EXOSC1 is not stably associated (Dai et al., 2018; Malet et al., 2010), suggesting that EXOSC1 might be involved in some functions independent of the complex.

In this study, we show that EXOSC1 acts as an ESM and sensitizes cancer cells to PARPi in KIRC. Due to the role of exosome in maintaining genomic stability, these results also indicate that a unit of multiprotein complex can play a role opposite to that of the complex.
Results

Identification of candidate ESMs in KIRC

Because that ESMs constantly assaults genomic DNA, we hypothesized that ESMs likely sensitized cancer cells to the inhibitors of DNA repair pathways. Considering that substitution is the most abundant mutation in all cancers, we initiated this study to identify candidate ESMs responsible for substitution mutations. To identify the candidate ESMs other than deamination, we focused on KIRC for three reasons: (1) KIRC shows the lowest proportion of mutations at CG in major cancer types (Burns et al., 2013b), suggesting that the deamination contributes less to the mutations in KIRC. (2) only low expressions of AID and APOBECs were detected in KIRC (Burns et al., 2013b). (3) The kidney potentially suffers less from exogenous source of mutations (EOSMs) (Loeb, 2011; Roberts and Gordenin, 2014). RNA-seq and exomic mutation data corresponding to 532 KIRC patients and 30,254 somatic substitution mutations in the TCGA were retrieved from The cBio Cancer Genomics Portal (http://cbioportal.org) (Figure 1D). Because that ESMs likely show similar impact on the template and code DNA strands, the 12 types of substitution were groups into 6 types of complementary substitution (c-substitution) to simplify the analyses (Figure 1D).

Spearman’s rank analysis was first performed to assess the correlation between each c-substitution type and the genome-wide expression of individual genes. Resultant p and r values were used for the further analyses. For example, GAPDH showed a $p = 0.0011$ and $r = 0.16$ correlation with C>A/G>T c-substitution, indicating that GAPDH expression displayed a positive correlation with C>A/G>T (Figure 1E). Similarly, CRB3 ($p = 0.0003$, $r = 0.17$) was positively correlated with A>T/T>A (Figure 1E). Although the $p$ values of multiple genes were lower than 0.05 (Figure 1F), only top ranked 200 genes (approximately 1% of the genome-wide genes) were taken as the candidates for each c-substitution type (Supplementary Table S1).

Student’s $t$-test analysis was then used to determine whether the expression difference of individual gene between the high and low c-substitution groups is significant. The expression of individual genes in each patient was normalized by a house keeping gene, TATA-binding protein (TBP) as previously described (Burns et al., 2013a; Burns et al., 2013b). According to each c-substitution, 532 KIRC patients were groups into 3 groups (high, medium and low). The difference of individual gene between high and low c-substitution mutation groups was then analyzed by student’s
\( t \)-test. Resultant \( p \) and fold of change (FC) values were used for the further analyses. For example, ASNS with \( p = 0.0005 \) and FC = 1.39, indicating that ASNS was increased in the high group (Figure 1G). Although the \( p \) values of multiple genes were lower than 0.05 (Figure 1H), only top 200 genes with high FC and \( p < 0.05 \) were taken as the candidates (Supplementary Table S2). Notably, none of APOBEC family members were identified as candidate by correlation or student’s \( t \)-test analyses, supporting that deamination contributes less to the mutations in KIRC.

Next, we performed meta-analyses to determine which of the 6 c-substitution types to be focused on. Mutation frequencies of c-substitution types were first analyzed in five major cancers: breast adenocarcinoma (BRCA), glioblastoma multiforme (GBM), bladder urothelial carcinoma (BLCA), acute myeloid leukemia (AML) and KIRC, which potentially suffer less from the EOSMs. As shown in Figure 1I, KIRC displayed higher frequencies of C>A/G>T, A>T/T>A, and A>C/T>G mutations. Using tumor protein p53 (TP53) as control, we then assessed the frequencies of c-substitution types in von Hippel-Lindau tumor suppressor (VHL), the most frequently mutated gene in KIRC. Consistent with previous studies (Burns et al., 2013a; Kandoth et al., 2013), the most frequent c-substitution type of TP53 mutations in BRCA was C>T/G>A (Figure 1J), while the most frequent type of VHL mutations in KIRC was C>A/G>T (\( p = 0.004 \), chi-squared test) (Figure 1K). Even after normalization according to the base frequency, this phenomenon was still observed (\( p = 0.021 \)) (Figure 1—figure supplement 1A and B). Further Kaplan–Meier (KM) analysis of overall survival (OS) indicated that patients with VHL C>A/G>T mutations showed poor OS (Figure 1L). These observations raised the significance of C>A/G>T c-substitutions in KIRC.

We then evaluated the expression difference of individual gene between the VHL C>A/G>T mutation positive and negative patients. Student’s \( t \)-tests analyses showed that 66 genes displayed \( p < 0.05 \) (Supplementary Table S3). Further overlap analyses demonstrated that cyclin B1 (CCNB1), exosome component 1 (EXOSC1), and RAB5 interacting factor (RAB5IF) were identified as candidate ESMs for C>A/G>T by all of the above analyses (Figure 1M).

**EXOSC1 promotes mutations in E. coli**

To evaluate the capability of the candidate gene to promote mutation, rifampicin-resistant assay in *E. coli* was performed as previously described.
Because that mutation of the rifampicin-targeted rpoB gene to rifampicin resistance (Rif\textsuperscript{R}) occurs at a low frequency, the capability of a gene to mutate rpoB to Rif\textsuperscript{R} can be evaluated by fluctuation analysis (Petersen-Mahrt et al., 2002). AID, an known ESM (Petersen-Mahrt et al., 2002), was used as a positive control. Four genes (CDK5, TARBP2, PSAT1 and NECAB3) were used as random controls (Figure 2—figure supplement 1A and Supplementary Table S4). These genes were expressed in \textit{E. coli} under the regulation of a trp/lac (tac) hybrid promoter, which could be activated by isopropyl β-D-1-thiogalactopyranoside (IPTG) (Figure 2A). Consistent with a previous study (Petersen-Mahrt et al., 2002), AID enhanced mutation in \textit{E. coli} (Figure 2B). Notably, EXOSC1 more significantly increased mutations than AID did ($p = 4.08 \times 10^{-5}$) (Figure 2B). We then evaluated the capabilities of EXOSC1 homologs (EXOSC2–EXOSC9) to promote mutations (Figure 2—figure supplement 1B). Among the members of exosome complex, EXOSC1 most notably enhanced mutation in \textit{E. coli} ($p = 3.5 \times 10^{-11}$) (Figure 2C). To determine whether the increase in mutation frequency stemmed from EXOSC1 protein itself, rifampicin-resistant assays were performed in the presence or absence of IPTG, the transcriptional inducer. As shown in Figure 2E–F, IPTG absence notably decreased the mutation frequency ($p = 1.29 \times 10^{-9}$), indicating that the protein of EXOSC1 promoted the mutation. Additionally, we evaluated the impact of EXOSC1 on the growth of \textit{E. coli}. As shown in Figure 2—figure supplement 1C, EXOSC1 expression only slightly decreased cell growth, which might be due to the increase in mutation burden (Schaaper and Dunn, 1987).

Next, the mutation spectra of Rif\textsuperscript{R} were analyzed by sequencing the rpoB gene PCR products from rifampicin-resistant clones. Sequencing of rpoB gene in 25 randomly selected rifampicin-resistant clones indicated that most of mutations in control clones were C>T/G>A transitions, while EXOSC1 frequently promoted C>A/G>T transversion mutations (Figure 2G–I). Moreover, most of C>A/G>T transversions in EXOSC1-transformed cells were clustered at C1576 (10/36 mutations) and C1699 (6/36 mutations), whereas C>T/G>A transitions in control cells showed a distinct distribution with major hot spots at C1565 (6/32 mutations) and C1721 (5/32 mutations) (Figure 2G). Hence, it was suggested that the mutations in control and EXOSC1-transformed cells were promoted by distinct mechanism. Interestingly, EXOSC1-transformed cells also showed a shift of C>A/G>T mutations from 6% (2/32 mutations) to 69% (25/36 mutations) ($p = 4.03 \times 10^{-7}$, chi-squared test) (Figure
2H–J). Even after normalization to the base frequency, this phenomenon was still significant ($p = 1.47 \times 10^{-6}$) (Figure 2—figure supplement 1D).

**EXOSC1 cleaves single-stranded DNA**

Considering that the exosome is well known to degrade RNA, we speculated that EXOSC1 might promote mutation through cleaving DNA. EXOSC1 was expressed and purified *in vitro* (Figure 3A). The resultant EXOSC1 protein was incubated with generic single-stranded DNA (ssDNA), double-stranded DNA (dsDNA) or the hybrid of DNA–RNA (Figure 3B). Polyacrylamide TBE-urea gel analyses of the resultant mixtures indicated that EXOSC1 notably cleaved ssDNA, while it displayed no detectable capability to cleave dsDNA or DNA-RNA hybrid (Figure 3C).

We then evaluated the capabilities of EXOSC1 homologs (EXOSC2–EXOSC9) to cleave ssDNA. The EXOSC protein was separately incubated with ssDNA. Gel analyses of the resultant mixtures indicated that, unlike EXOSC1, none of EXOSC2–EXOSC9 detectably cleaved ssDNA (Figure 3—figure supplement 1A). Considering that EXOSC1 is well known to form a complex with other exosome members, we also assessed the impact of the exosome members on the cleavage activity of EXOSC1. EXOSC1 was incubated with ssDNA in the presence of individual EXOSC1 homolog. Interestingly, EXOSC6, EXOSC7 and EXOSC8 decreased EXOSC1 cleavage activity (Figure 3D), while EXOSC2, EXOSC3, EXOSC4 and EXOSC9 showed no detectable impact (Figure 3—figure supplement 1B). We then evaluated the impact of the reaction components and pH on the cleavage activity of EXOSC1. It was found that K$^+$ and Mg$^{2+}$ enhanced the cleavage activity of EXOSC1 (Figure 3E), and EXOSC1 showed the highest cleavage activity at pH 7.0 (Figure 3F). Further analyses indicated that the cleavage rate of EXOSC1 was approximately $4 \times 10^{-4}$/min at 37 °C (Figure 3G and H).

**EXOSC1 prefers to cleave C sites in single-stranded DNA**

Considering that conserved exosome prefers to degrade the RNA with specific sequence (Cvetkovic et al., 2017), we determined whether EXOSC1 preferred to cleave some site(s) in ssDNA. EXOSC1 was incubated with DNAs containing unbiased sequence and distinct 3' end. Consistent with the result of the generic DNA, EXOSC1 cleaved unbiased ssDNA and displayed no detectable capability to cleave dsDNA or DNA-RNA hybrid (Figure 4A and Figure 4—figure supplement 1A). And...
only EXOSC1 displayed cleavage activity against unbiased ssDNA (Figure 4B).
Interestingly, the cleavage rate against unbiased ssDNA (approximately 1.2 X
10^3/min) was higher than that against generic ssDNA (Figure 4—figure supplement
1B), suggesting that the DNA sequence might show some impact on the activity of
EXOSC1. Notably, mass spectrometry (MS) analyses demonstrated that the resultant
mixtures contained more free C than the other three base types (Figure 4E),
suggesting that EXOSC1 preferred to cleave C sites in ssDNA. Since EXOSC1 was
correlated with the C>A/G>T c-substitution type, it was likely that EXOSC1 cleaved
C sites in ssDNA and subsequently resulted in C>A mutations through “A” rule DNA
repair.

To evaluate the above hypothesis, we then determined whether EXOSC1-promoted
mutations displayed strand asymmetries. Considering that EXOSC1 cleaved C sites in
ssDNA but not DNA-RNA hybrid, we speculated that “transcribed” template strands
likely bound by RNA were less cleaved by EXOSC1. As shown in Figure 4F, C>A
transversions in the “untranscribed” coding strand lead to C>A mutations in a gene,
while C>A transversions in the “transcribed” template strand result in G>T mutations.
Therefore, the capability of EXOSC1 to promote strand mutational asymmetry can be
evaluated by comparing C>A and G>T frequencies. We first analyzed the
distributions of C>A and G>T substitutions, instead of the distributions of C>A/G>T
c-substitution, in the EXOSC1-transformed E. coli cells described above. As shown in
Figure 4—figure supplement 1E and F, C>A substitution in rpoB gene were generated
at a much higher frequency than G>T. Compared with the control, EXOSC1 enhanced
C>A from 0% to 69% (p = 2.67 X 10^-7), while EXOSC1 only enhanced G>T from 6%
to 17% even without a significance (p = 0.27) (Figure 4—figure supplement 1E and
F). Next, we evaluated C>A strand asymmetry in KIRC using spearman’s rank and
student’s t-test analyses. Spearman’s rank analyses indicated that EXOSC1 showed
the highest correlation (r) with C>A, and the correlation between EXOSC1 and G>T
was even lower than that between EXOSC1 and C>A/G>T (Figure 4G and Figure
4—figure supplement 1G). To evaluate the impact of group number on the further
student’s t-test analyses, the KIRC patients were grouped into 2, 3, 4 or 5 groups
according to the mutation types C>A, G>T, C>A/G>T and total (12 substitution
types). As expected, the EXOSC1 differences between the low and high C>A/G>T
groups were more significant than those between the low and high groups of total (12
substitution types) mutations (Figure 4—figure supplement 1H and I). Importantly,
EXOSC1 showed more significant expression differences between the low and high C>A groups than those between the low and high G>T groups (Figure 4H and Figure 4—figure supplement 1J), suggesting that EXOSC1 prefers to cleave C sites in ssDNA in vivo.

EXOSC1 enhances DNA damage and mutations in KIRC cells

Considering that EXOSC1 cleaves DNA in vitro, we then evaluated the capability of EXOSC1 to promote intracellular DNA damage using γ-H2AX staining and neutral comet tail assays. The 769-P and TUHR14TKB KIRC cells were transfected with the plasmid encoding EXOSC1 using empty vector as a control (Figure 5—figure supplement 1A). γ-H2AX staining analyses of the resultant cells demonstrated that EXOSC1 increased the γ-H2AX foci in the cells (Figure 5A). The number of γ-H2AX-positive cells was increased approximately 7-fold by EXOSC1 (Figure 5B). While knockdown of EXOSC1 reduced the γ-H2AX foci (Figure 5C, and Figure 5—figure supplement 1B and C). Consistent with the results of γ-H2AX staining, comet tail analyses also indicated that EXOSC1 increased DNA damage (Figure 5D and E).

Due to the central role of DNA damage in mutations, we performed differential DNA denaturation PCR (3D-PCR) to determine whether EXOSC1 enhances mutation in KIRC cells. Because that DNA sequences with more A/T content can be amplified at lower denaturation temperatures than parental sequences, 3D-PCR enables qualitative estimates of genomic C/G>A/T mutations in a population of cells. As shown in Figure 5F, the enhanced expression of EXOSC1 (EXOSC1-OE) increased lower temperature amplicons (LTAs) of VHL, suggesting that EXOSC1 increased the mutations in VHL gene. Consistently, further sequencing analyses of the LTAs indicated that EXOSC1-OE cells showed more C>A mutations in VHL gene (Figure 5F).

Considering that the “A” rule DNA repair is dependent on X-ray repair cross-complementing 1 (XRCC1) (Sale et al., 2001), we knocked down XRCC1 to evaluate the role of XRCC1 in EXOSC1-promoted mutations (Figure 5—figure supplement 1D–F). 3D-PCR analyses of the resultant cells indicated that knockdown of XRCC1 impaired the capability of EXOSC1 to increase the LTAs (Figure 5G). Furthermore, both XRCC1 knockdown (XRCC1-KD) and EXOSC1 knockdown (EXOSC1-KD) decreased the LTAs (Figure 5H and I). Additionally, a subcutaneous
xenograft tumor model was used to determine whether EXOSC1 enhances DNA mutations in vivo (Figure 5J). Stable control (vector), EXOSC1-OE and EXOSC1-KD 769-P cells were subcutaneously implanted. After two weeks, 3D-PCR analyses of the resultant tumors indicated that EXOSC1 increased the LTAs of VHL, whereas knockdown of EXOSC1 reduced the LTAs (Figure 5K), suggesting that EXOSC1 enhanced mutations in KIRC.

EXOSC1 sensitizes KIRC cells to PARP inhibitor

Considering the central roles of mutation in the process of cancers, we evaluated the potential clinical significance of EXOSC1 in KIRC using KM analyses. KM analyses of disease-free survival (DFS) and OS were performed using the clinical data from 532 KIRC patients in TCGA. The fragments per kilo base per million mapped reads (FPKM) were used to evaluate the expression of EXOSC1 in KIRC (Figure 6—figure supplement 1A). The median-separation KM analyses indicated that high EXOSC1 group showed poor DFS and OS (Figure 6A and B). The median DFS in the low EXOSC1 group was 32.0-month longer than that in the high group (p = 9.78 X 10^-8, log-rank test) (Figure 6A). Consistently, the median OS in the low EXOSC1 group was 36.9-month longer than that in high group (p = 2.2 X 10^-8) (Figure 6B). As expected, the best-separation KM analysis also indicated that high EXOSC1 group significantly showed poor OS (p = 2.6 X 10^-12) (Figure 6—figure supplement 1B).

Due to the critical role of VHL mutation in KIRC, we then evaluated the potential clinical significance of EXOSC1 in the presence and absence of VHL mutation. KM analyses indicated that high EXOSC1 group showed poor OS in both presence (median OS (high vs low group) = 65.2 vs 98.5 months, p = 0.015) and absence (median OS (high vs low group) = 75.7 vs 104.5 months, p = 1.0 X 10^-5) of VHL mutation (Figure 6C and Figure 6—figure supplement 1C).

Considering that EXOSC1 increases DNA damage, we speculated that EXOSC1 potentially sensitizes KIRC cells to the inhibitor of poly(ADP-ribose) polymerase (PARP), which treats cancers via blocking DNA repair. As previously described (Li et al., 2020), colony formation assays were performed to evaluate the role of EXOSC1 in the response to a PARP inhibitor, niraparib. Stable control (vector) and enhanced EXOSC1 (EXOSC1-OE) KIRC cells were seeded and treated with serial dilutions of niraparib until colonies were notably formed. As shown in Fig 6D–F, niraparib more notably inhibited the 769-P and TUHR14TKB KIRC cells with enhanced EXOSC1,
suggesting that EXOSC1 sensitized the cells to the inhibitor. Consistent with the results in 769-P and TUHR14TKB cells, EXOSC1 also sensitized SNU-1272 and Caki-2 KIRC cells to the inhibitor (Figure 6—figure supplement 1D). Next, we determined whether EXOSC1 could sensitize KIRC cells to niraparib in xenograft mouse models. The control and EXOSC1-OE cells were subcutaneously injected. Resultant tumor-bearing mice were grouped and treated by vehicle or niraparib. Consistent with the ex vivo results, niraparib more notably inhibited the tumor with enhanced EXOSC1 (Figure 6G and I), indicating that EXOSC1 sensitized KIRC xenografts to the inhibitor. No significant weight loss was observed throughout the study, suggesting that the niraparib treatment was well tolerated (Figure 6H and J).

**Discussion**

The genomic integrity of human cells is constantly assaulted by ESMs. Although human cells possess multi DNA repair mechanisms to counteract these constant assaults, not all lesions are correctly repaired and almost inevitably result in mutations. The central roles of these acquired mutations in nearly all cancers (Jeggo et al., 2015; Roos et al., 2015) emphasize the identification and understanding of the ESMs. Here, we show that EXOSC1 cleaves ssDNA and acts as an ESM in KIRC. Consistent with the capability of EXOSC1 to promote DNA damage and mutations, KIRC patients with high EXOSC1 showed a poor prognosis, and EXOSC1 also sensitized cancer cells to the PARP inhibitor.

Our results show that a unit of multiprotein complex can play a role distinct from the function(s) of the complex. Biological processes frequently require fine control over the formation of a multiprotein complex in a particular region of the cell. The exosome complex is well known for its roles in RNA degradation (Januszyk and Lima, 2014; Kilchert et al., 2016). However, the role of exosome complex members other than RNA binding and degradation remains elusive. Interestingly, EXOSC1 can disassociate from the exosome complex (Dai et al., 2018; Malet et al., 2010), suggesting that EXOSC1 might be involved in some functions independent of the exosome. Our study demonstrated that EXOSC1 acts as an ESM to promote mutagenesis. Conversely, previous studies have described that the exosome, as a multiprotein complex, protects cells from genomic instability by preventing the formation of DNA/RNA hybrids and restricting DNA strand mutational asymmetry (Lim et al., 2017; Pefanis and Basu, 2015; Pefanis et al., 2015). This phenomenon can
at least partially be explained by the finding that some exosome members (EXOSC7 and EXOSC8) can block the activity of EXOSC1 to cleave DNA. Therefore, a single EXOSC1 protein can show different responses depending on the presence/absence of its interacting partners. Further studies are needed to better understand the roles of the individual exosome member.

The potential pathological significance of EXOSC1 is supported by its association with poor DFS and OS in KIRC. Due to the capability of EXOSC1 to cleave DNA and promote mutations, EXOSC1 might enhance mutations and consequently provide genetic fuel for cancer development, metastasis, and even therapy resistance. Therefore, EXOSC1 might represent not only a KIRC marker but also a target to decrease the rate of KIRC evolution and stabilize the targets of existing therapeutics. Furthermore, targeting DNA repair in cancers by inhibiting PARPs offers an important therapeutic strategy (Cleary et al., 2020). Unfortunately, the failure of PARP inhibitors to markedly benefit patients enforces the necessity for developing new strategies to improve their efficacy (Cleary et al., 2020; Dizon, 2017; Lord and Ashworth, 2017). Our study demonstrated that EXOSC1 sensitized KIRC cells to PARP inhibitor, suggesting inhibition of PARPs might be a penitential strategy to treat KIRC patients with high EXOSC1. We also noticed that KIRC patients with high EXOSC1 and VHL mutations showed the poorest OS. Considering the DNA cleavage activity of EXOSC1 and the role of VHL in stabilizing the genome (Thoma et al., 2009; Zhang et al., 2018), we speculate that patients with the VHL mutation and high EXOSC1 might show higher possibility to benefit from PARP inhibitor(s).

However, several limitations of this study should be noted. First, we observed a notable variation in terms of the correlation with a different c-substitution type for a given gene, implying the need for further studies. Second, although we showed that EXOSC1 could cleave ssDNA and act as an ESM, we did not directly identify the mechanism responsible for turning the DNA cleavages into mutations. The role of XRCC1 in EXOSC1-promoted mutations was only briefly evaluated. Hence, we cannot exclude the possibility that other proteins might contribute to this process. Third, because that DNA cleavage by EXOSC1 should be independent of the cancer type, we propose that EXOSC1 likely contributes to the mutations in the cancers other than KIRC. And more work is still needed. Despite these limitations, our results still indicate that EXOSC1 acts as an ESM in KIRC.
Materials and methods

Sample preparation

Samples of 532 KIRC patients from TCGA used for expression and mutation analyses were collected through The cBio Cancer Genomics Portal (http://cbioportal.org) as described in our previous studies (Li et al., 2020; Zhou et al., 2019).

Cell culture

All cell lines were obtained from the American Type Tissue Collection. The 769-P, SNU-1272 and Caki-2 cells were maintained in RPMI 1640 medium containing 10% heat-inactivated FBS, 100 U/ml penicillin and streptomycin at 37°C under a humidified atmosphere of 5% CO₂. TUHR14TKB cells were maintained in DMEM containing 10% heat-inactivated FBS, 100 U/ml penicillin and streptomycin at 37°C under a humidified atmosphere of 5% CO₂. Transfections were performed using lipofectamine 2000 (Thermo Fisher Scientific Inc., Waltham, MA, USA).

Antibodies, reagents and plasmids

Anti-Flag (Cat. number: F7425) and anti-His (Cat. number: SAB1306085) antibodies were from Sigma-Aldrich (St. Louis, MS, USA). Anti-phospho-γ-H2AX (Ser139) (Cat. number: 05-636) and anti-XRCC1 (Cat. number: SAB1306085) were from Millipore (Billerica, MA, USA). Anti-EXOSC1 (Cat. number: EPR13526) was from Abcam (Cambridge, MA, USA). Niraparib (Cat. number: HY-10619) and rifampicin (Cat. number: R3501) were from MedChem Express and Sigma-Aldrich, respectively.

Full-length EXOSC1 was cloned into the Xba I and Nhe I sites of the lentivirus vector pCDH-CMV-MCS-EF1-Puro (System Biosciences, Mountain View, CA, USA) to construct pCDH-Flag EXOSC1. CCGAGTTTCTACAGACCTAAG and CGAGAAGCTATCCCGAAAGAA sequences were cloned into pLKO.1 to construct the pLKO shEXOSC1-1 and pLKO shEXOSC1-2 plasmids, respectively. Similarly, CCAGGCTCCAGGAAGATATA and CGATACGTCACAGCCTTCAAT sequences were cloned into pMKO.1 to construct the pMKO shXRCC1-1 and pMKO shXRCC1-2 plasmids, respectively. According to the knockdown efficiency (Figure 5—figure supplement 1), pLKO shEXOSC1-1 and pMKO.1 shXRCC1-1 with higher knockdown efficiencies were used to generate EXOSC1-KD and XRCC1-KD cells.

AID (NM_020661.4), CDK5 (NM_004935.4), TARBP2 (NM_134323.1), EXOSC1 (NM_016046.5), RAB5IF (NM_018840.5), CCNB1 (NM_031966.4), PSAT1
(NM_058179.4), NECAB3 (NM_031232.3), EXOSC2 (NM_014285.7), EXOSC3 (NM_016042.4), EXOSC4 (NM_019037.3), EXOSC5 (NM_020158.4), EXOSC6 (NM_058219.3), EXOSC7 (NM_015004.4), EXOSC8 (NM_181503.3) and EXOSC9 (NM_005033) were amplified and cloned into the pET-28a(+) vector (Novogen Limited, Hornsby Westfield, NSW, Australia) to construct the pET-28a-Gene-6XHis E. coli expression plasmids. PCR primers for the amplification of above genes are described in Supplementary Table S5.

**Immunoblotting and immunofluorescence**

Immunoblotting and immunofluorescence were carried out as described in our previous study (Song et al., 2018; Wang et al., 2020).

**Rifampicin-resistant assay in E. coli**

Rifampicin-resistant assays were carried out as described previously (Petersen-Mahrt et al., 2002). Briefly, rifampicin-resistant assays for each gene were performed using 30 independent cultures grown overnight to saturation in rich medium supplemented with 50 mg/L kanamycin and 1 mM IPTG. RifR mutants were selected on medium containing 50 mg/L rifampicin. Mutation frequencies were assessed by determining the median number of rifampicin-resistant clones per $10^9$ viable plated cells. The mutation spectra of RifR were analyzed by sequencing the amplified rpoB 627-bp PCR products using 5'-TTGGCGAAATGGCGGAAAACC-3' and 5'-CACCGACGGATACCACCTGCTG-3' primers.

**Expression and purification of EXOSC proteins**

Expression and purification of EXOSC proteins were carried out as described in our previous study (Wang et al., 2020). pET-28a-EXOSCs-6XHises encoding His-tagged EXOSCs were introduced into BL21 (DE3)-pLysS, which were grown in nutrient-rich medium with 32Y (containing 3.2% (w/v) yeast extract, 0.8% (w/v) peptone and 0.58% (w/v) NaCl) in 10 mM Tris–HCl at 30°C and pH 7.4. Protein expression was induced with 0.4 mM IPTG at 20°C for 20 h after the cells reached an OD$_{600}$ of 0.4–0.5. Induced BL21 (DE3)-pLysS host cells without any plasmid were used as a negative control. The resultant cells were harvested by centrifugation at 5000 g for 10 min and washed twice with ice-cold PBS. The collected cells were resuspended in
PBS (1 g of wet weight cells per 10 mL of PBS) containing 1 mM MgCl₂, 20 mM imidazole, 1 tablet/50 mL protease inhibitor cocktail, and 100 U/mL DNase. Resuspended cells were broken by an ultrasonic wave. Cell lysates were centrifuged at 20,000 g at 4°C for 30 min to remove unbroken cells and debris.

After pre-equilibration with 10 column volumes (CV) of binding buffer (PBS containing 10% (v/v) glycerol and 20 mM imidazole, pH 7.6), Ni sepharose 6 Fast flow (GE Healthcare, New York, NY, USA) was applied for the purification of EXOSCs. The resins were washed 5 times and eluted using elution buffer (binding buffer containing 300 mM imidazole). EXOSCs were concentrated using an Amicon Ultrafree centrifugal filter (Millipore Corporation, Billerica, MA, USA) and pre-equilibrated with 10 mM HEPES buffer (pH 7.4) containing 150 mM NaCl and 10% glycerol. Size-exclusion chromatography (SEC) with a Superdex-200 HiLoad 10/600 column (GE Healthcare, New York, NY, USA) was used to further purify EXOSCs. The purity of the fractions was analyzed by coomassie blue staining. The protein concentration was determined using a BCA assay according to the manufacturer’s instructions (Pierce, Rockland, IL, USA).

**Cleavage activity assay in vitro**

The cleavage assays of EXOSCs were carried out in reaction buffer modified from a previous study (Quansheng et al., 2006). Briefly, 50 µL of reaction mixture containing 1 µM oligonucleotides, 1 µM EXOSC protein, 70 mM KCl, 700 µM MgCl₂, 1 mM DTT, and 20 mM Tris-HCl pH 7.0 was incubated at 37°C for 4 h. The reaction was stopped by addition of 10 µM proteinase K at 58°C for 10 min and heating at 90°C for 30 s. The resultant samples were then analyzed using 15% polyacrylamide TBE-urea gels.

**LC-MS/MS analysis**

LC-MS/MS analyses of deoxyadenine (A), deoxythymidine (T), deoxyguanine (G) and deoxycytocine (C) were carried out as described in our previous studies (Song et al., 2018; Wang et al., 2020).

**Generation of stable cell lines**

Stable cell lines were generated as described in our previous study (Song et al., 2018;
Briefly, the TUHR14TKB, SNU-1272, 769-P and Caki-2 cells were infected with pCDH-CMV-MCS-EF1-Puro (empty vector used as control), pCDH-Flag EXOSC1 (EXOSC1-OE), pLKO.1-scramble shRNA (empty vector used as control), pLKO shEXOSC1-1 (EXOSC1-KD) or pLKO shEXOSC1-2 lentiviral particles, which were generated following the manufacturer’s protocol (System Biosciences, Mountain View, CA, USA). The resultant cells were selected with puromycin for 2 weeks. These stable cells were then infected with virus encoding pMKO.1, pMKO.1 shXRCC1-1 (XRCC1-KD) or pMKO.1 shXRCC1-2. The resultant cells were selected with hygromycin B for 2 weeks to generate stable XRCC1 knockdown cells. According to the knockdown efficiency (Figure 5—figure supplement 1), shEXOSC1-1 and shXRCC1-1 with higher knockdown efficiencies were used as EXOSC1-KD and XRCC1-KD in this study.

γ-H2AX staining and neutral comet tail assays

γ-H2AX staining and neutral comet tail assays were performed as described previously (Li et al., 2020).

3D-PCR and sequencing

3D-PCRs of VHL mutations were carried out as described previously (Burns et al., 2013a) using first (5’-GAGTACGGCCCTGAAGAAGA-3’ and 5’-TCAATCTCCCATCCGTTGAT-3’) and nested (5’-TGGCCTAGGTAACCTGC-3’ and 5’-GCGGCAGCGTTGGTAGG-3’) PCR primers. PCR products were then analyzed by gel electrophoresis, cloned into pMD20-T vector, and sequenced.

Colony-forming assay

The colony-forming assays were performed as described in our previous study (Li et al., 2020).

Subcutaneous xenograft tumor growth in vivo

The following animal-handling procedures were approved by the Animal Care and Use Committee of Dalian Medical University. Xenograft models were carried out as described in our previous studies (Li et al., 2020; Song et al., 2018; Yang et al., 2010). Briefly, 2 × 10^6 stable control/EXOSC1-OE 769-P and Caki-2 cells were suspended
and injected subcutaneously into the flank of 6-week-old nude mice. After 7 days, these tumor-bearing mice were randomized into 4 groups (6 mice per group) and treated by oral gavage twice a day with vehicle or niraparib (4 mg/kg). The mice were observed daily and weighed once per week. Tumor size was measured using a caliper, and the tumor volume was calculated using the following formula: \(0.52 \times L \times W^2\), where \(L\) is the longest diameter and \(W\) is the shortest diameter. Mice were euthanized when the tumors reached 1500 mm\(^3\) or showed necrosis.

**Statistical analyses**

\(P\) values were calculated by the two-tailed student’s \(t\)-test, log-rank test, fisher’s exact-test, chi-squared test, or spearman correlation analyses as noted. \(P\) values < 0.05 were considered statistically significant.

**Availability**

All data associated with this study are available in the main text or the supplementary materials.

**Experimental replicates and reproducibility**

All data presented in this paper are representative of 2–4 independent experiments with comparable results.

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**Authors’ contributions**

Q.Y. and C.S. designed the experiments and wrote the paper. Q.X., Q.L. and L.W. performed most experiments. Q.X. and Q.L. carried out EXOSC protein extraction. C.S. and W.L. performed bioinformatics analyses. N.W. and K.W. maintained the cells and performed some
of the western blot analyses. Q.L. and L.W. carried out the γ-H2AX staining and comet assays.

Conflict of interest

The authors declare no competing interests.

References

Alexandrov, L.B., Nik-Zainal, S., Wedge, D.C., Aparicio, S.A., Behjati, S., Biankin, A.V., Bignell, G.R., Bolli, N., Borg, A., Borresen-Dale, A.L., et al. (2013a). Signatures of mutational processes in human cancer. Nature 500, 415-421.

Alexandrov, L.B., Nik-Zainal, S., Wedge, D.C., Aparicio, S.A., Behjati, S., Biankin, A.V., Bignell, G.R., Bolli, N., Borg, A., Borresen-Dale, A.L., et al. (2013b). Signatures of mutational processes in human cancer. Nature 500, 415-421.

Barlow, J.H., Faryabi, R.B., Callen, E., Wong, N., Malhowski, A., Chen, H.T., Gutierrez-Cruz, G., Sun, H.W., McKinnon, P., Wright, G., et al. (2013). Identification of early replicating fragile sites that contribute to genome instability. Cell 152, 620-632.

Bousquet-Antonelli, C., Presutti, C., and Tollervey, D. (2000). Identification of a regulated pathway for nuclear pre-mRNA turnover. Cell 102, 765-775.

Brown, J.S., O’Carrigan, B., Jackson, S.P., and Yap, T.A. (2017). Targeting DNA Repair in Cancer: Beyond PARP Inhibitors. Cancer Discov 7, 20-37.

Brown, J.T., Bai, X., and Johnson, A.W. (2000). The yeast antiviral proteins Ski2p, Ski3p, and Ski8p exist as a complex in vivo. Rna-a Publication of the Rna Society 6, 449.

Buisson, R., Langenbucher, A., Bowen, D., Kwan, E.E., Benes, C.H., Zou, L., and Lawrence, M.S. (2019). Passenger hotspot mutations in cancer driven by APOBEC3A and mesoscale genomic features. Science 364.

Burgess, D.J. (2019). Switching APOBEC mutation signatures. Nat Rev Genet 20, 253.

Burns, M.B., Lackey, L., Carpenter, M.A., Rathore, A., Land, A.M., Leonard, B., Refsland, E.W., Kotandeniya, D., Tretyakova, N., Nikas, J.B., et al. (2013a). APOBEC3B is an enzymatic source of mutation in breast cancer. Nature 494, 366-370.
Burns, M.B., Temiz, N.A., and Harris, R.S. (2013b). Evidence for APOBEC3B mutagenesis in multiple human cancers. Nat Genet 45, 977-983.

Cleary, J.M., Aguirre, A.J., Shapiro, G.I., and D'Andrea, A.D. (2020). Biomarker-Guided Development of DNA Repair Inhibitors. Mol Cell 78, 1070-1085.

Cvetkovic, M.A., Wurm, J.P., Audin, M.J., Schutz, S., and Sprangers, R. (2017). The Rrp4-exosome complex recruits and channels substrate RNA by a unique mechanism. Nat Chem Biol 13, 522-528.

Dai, L., Zhao, T., Bisteau, X., Sun, W., Prabhu, N., Lim, Y.T., Sobota, R.M., Kaldis, P., and Nordlund, P. (2018). Modulation of Protein-Interaction States through the Cell Cycle. Cell 173, 1481-1494 e1413.

Dizon, D.S. (2017). PARP inhibitors for targeted treatment in ovarian cancer. Lancet 390, 1929-1930.

Farmer, H., McCabe, N., Lord, C.J., Tutt, A.N., Johnson, D.A., Richardson, T.B., Santarosa, M., Dillon, K.J., Hickson, I., Knights, C., et al. (2005). Targeting the DNA repair defect in BRCA mutant cells as a therapeutic strategy. Nature 434, 917-921.

Greaves, M. (2018). A causal mechanism for childhood acute lymphoblastic leukaemia. Nat Rev Cancer 18, 471-484.

Hutchinson, L. (2013). Genetics: Signatures of mutational processes in cancer—a big step closer. Nat Rev Clin Oncol 10, 545.

Januszyk, K., and Lima, C.D. (2014). The eukaryotic RNA exosome. Curr Opin Struct Biol 24, 132-140.

Jeggo, P.A., Pearl, L.H., and Carr, A.M. (2015). DNA repair, genome stability and cancer: a historical perspective. Nature Reviews Cancer 16, 35.

Jeggo, P.A., Pearl, L.H., and Carr, A.M. (2016). DNA repair, genome stability and cancer: a historical perspective. Nat Rev Cancer 16, 35-42.

Jonathan, H., John, L.C., and David, T. (2006). RNA-quality control by the exosome. Nat Rev Mol Cell Biol 7, 529-539.

Kandoth, C., McLellan, M.D., Vandin, F., Ye, K., Niu, B., Lu, C., Xie, M., Zhang, Q., McMichael, J.F., Wyczalkowski, M.A., et al. (2013). Mutational landscape and significance across 12 major cancer types. Nature 502, 333-339.

Kilchert, C., Wittmann, S., and Vasiljeva, L. (2016). The regulation and functions of the nuclear RNA exosome complex. Nat Rev Mol Cell Biol 17, 227-239.

Li, S., Zhang, Y., Wang, N., Guo, R., Liu, Q., Lv, C., Wang, J., Wang, L., and Yang,
Q.K. (2020). Pan-cancer analysis reveals synergistic effects of CDK4/6i and PARPi combination treatment in RB-proficient and RB-deficient breast cancer cells. Cell Death Dis 11, 219.

Lim, J., Giri, P.K., Kazadi, D., Laffleur, B., Zhang, W., Grinstein, V., Pefanis, E., Brown, L.M., Ladewig, E., Martin, O., et al. (2017). Nuclear Proximity of Mtr4 to RNA Exosome Restricts DNA Mutational Asymmetry. Cell 169, 523-537 e515.

Loeb, L.A. (2011). Human cancers express mutator phenotypes: origin, consequences and targeting. Nat Rev Cancer 11, 450-457.

Lord, C.J., and Ashworth, A. (2012). The DNA damage response and cancer therapy. Nature 481, 287-294.

Lord, C.J., and Ashworth, A. (2017). PARP inhibitors: Synthetic lethality in the clinic. Science 355, 1152-1158.

Lorentzen, E., Basquin, J., Tomecki, R., Dziembowski, A., and Conti, E. (2008). Structure of the active subunit of the yeast exosome core, Rrp44: diverse modes of substrate recruitment in the RNase II nuclease family. Molecular Cell 29, 717-728.

Malet, H., Topf, M., Clare, D.K., Ebert, J., Bonneau, F., Basquin, J., Drazkowska, K., Tomecki, R., Dziembowski, A., Conti, E., et al. (2010). RNA channelling by the eukaryotic exosome. EMBO Rep 11, 936-942.

McGranahan, N., Rosenthal, R., Hiley, C.T., Rowan, A.J., Watkins, T.B.K., Wilson, G.A., Birkbak, N.J., Veeriah, S., Van Loo, P., Herrero, J., et al. (2017). Allele-Specific HLA Loss and Immune Escape in Lung Cancer Evolution. Cell 171, 1259-1271 e1211.

Pearl, L.H., Schierz, A.C., Ward, S.E., Al-Lazikani, B., and Pearl, F.M. (2015). Therapeutic opportunities within the DNA damage response. Nat Rev Cancer 15, 166-180.

Pefanis, E., and Basu, U. (2015). RNA Exosome Regulates AID DNA Mutator Activity in the B Cell Genome. Adv Immunol 127, 257-308.

Pefanis, E., Wang, J., Rothschild, G., Lim, J., Kazadi, D., Sun, J., Federation, A., Chao, J., Elliott, O., Liu, Z.P., et al. (2015). RNA exosome-regulated long non-coding RNA transcription controls super-enhancer activity. Cell 161, 774-789.

Peng, L., Bian, X.W., Li, D.K., Xu, C., Wang, G.M., Xia, Q.Y., and Xiong, Q. (2015). Large-scale RNA-Seq Transcriptome Analysis of 4043 Cancers and 548 Normal Tissue Controls across 12 TCGA Cancer Types. 5, 13413.
Petersen-Mahrt, S.K., Harris, R.S., and Neuberger, M.S. (2002). AID mutates E. coli suggesting a DNA deamination mechanism for antibody diversification. Nature 418, 99.

Petljak, M., Alexandrov, L.B., Brammeld, J.S., Price, S., Wedge, D.C., Grossmann, S., Dawson, K.J., Ju, Y.S., Iorio, F., Tubio, J.M.C., et al. (2019). Characterizing Mutational Signatures in Human Cancer Cell Lines Reveals Episodic APOBEC Mutagenesis. Cell 176, 1282-1294 e1220.

Quanshe Ng, L., Greimann, J.C., and Lima, C.D. (2006). Reconstitution, activities, and structure of the eukaryotic RNA exosome. Cell 127, 1223-1237.

Roberts, S.A., and Gordenin, D.A. (2014). Hypermutation in human cancer genomes: footprints and mechanisms. Nat Rev Cancer 14, 786-800.

Robertson, A.G., Kim, J., Al-Ahmadie, H., Bellmunt, J., Guo, G., Cherniack, A.D., Hinoue, T., Laird, P.W., Hoadley, K.A., Akbani, R., et al. (2017). Comprehensive Molecular Characterization of Muscle-Invasive Bladder Cancer. Cell 171, 540-556 e525.

Roos, W.P., Thomas, A.D., and Kaina, B. (2015). DNA damage and the balance between survival and death in cancer biology. Nature Reviews Cancer 16, 20.

Roos, W.P., Thomas, A.D., and Kaina, B. (2016). DNA damage and the balance between survival and death in cancer biology. Nat Rev Cancer 16, 20-33.

Sale, J.E., Calandrini, D.M., Takata, M., Takeda, S., and Neuberger, M.S. (2001). Ablation of XRCC2/3 transforms immunoglobulin V gene conversion into somatic hypermutation. Nature 412, 921-926.

Schaaper, R.M., and Dunn, R.L. (1987). Spectra of spontaneous mutations in Escherichia coli strains defective in mismatch correction: the nature of in vivo DNA replication errors. Proceedings of the National Academy of Sciences of the United States of America 84, 6220-6224.

Song, C., Wang, L., Wu, X., Wang, K., Xie, D., Xiao, Q., Li, S., Jiang, K., Liao, L., Yates, J.R., 3rd, et al. (2018). PML Recruits TET2 to Regulate DNA Modification and Cell Proliferation in Response to Chemotherapeutic Agent. Cancer Res 78, 2475-2489.

Thoma, C.R., Toso, A., Gutbrodt, K.L., Reggi, S.P., Frew, I.J., Schraml, P., Hergovich, A., Moch, H., Meraldi, P., and Krek, W. (2009). VHL loss causes spindle misorientation and chromosome instability. Nat Cell Biol 11, 994-1001.
Tomecki, R., Drazkowska, K., and Dziembowski, A. (2010). Mechanisms of RNA degradation by the eukaryotic exosome. Chembiochem 11, 938-945.

Tubbs, A., and Nussenzweig, A. (2017). Endogenous DNA Damage as a Source of Genomic Instability in Cancer. Cell 168, 644-656.

Tutt, A., Robson, M., Garber, J.E., Domchek, S.M., Audeh, M.W., Weitzel, J.N., Friedlander, M., Arun, B., Loman, N., Schmutzler, R.K., et al. (2010). Oral poly(ADP-ribose) polymerase inhibitor olaparib in patients with BRCA1 or BRCA2 mutations and advanced breast cancer: a proof-of-concept trial. Lancet 376, 235-244.

Uhlen, M., Zhang, C., Lee, S., Sjostedt, E., Fagerberg, L., Bidkhori, G., Benfeitas, R., Arif, M., Liu, Z., Edfors, F., et al. (2017). A pathology atlas of the human cancer transcriptome. Science 357.

Ulmke, P.A., Xie, Y., Sokpor, G., Pham, L., Shomroni, O., Berulava, T., Rosenbusch, J., Basu, U., Fischer, A., Nguyen, H.P., et al. (2021). Post-transcriptional regulation by the exosome complex is required for cell survival and forebrain development via repression of P53 signaling. Development 148.

Wang, L., Song, C., Wang, N., Li, S., Liu, Q., Sun, Z., Wang, K., Yu, S.C., and Yang, Q. (2020). NADP modulates RNA m(6)A methylation and adipogenesis via enhancing FTO activity. Nat Chem Biol 16, 1394-1402.

Wasmuth, E.V., Kurt, J., and Lima, C.D. (2014). Structure of an Rrp6-RNA exosome complex bound to poly(A) RNA. Nature 511, 435-439.

Weinstein, J.N., Collisson, E.A., Mills, G.B., Shaw, K.R., Ozenberger, B.A., Ellrott, K., Shmulevich, I., Sander, C., and Stuart, J.M. (2013). The Cancer Genome Atlas Pan-Cancer analysis project. Nature genetics 45, 1113-1120.

Yang, Q., Deng, X., Lu, B., Cameron, M., Fears, C., Patricelli, M.P., Yates, J.R., 3rd, Gray, N.S., and Lee, J.D. (2010). Pharmacological inhibition of BMK1 suppresses tumor growth through promyelocytic leukemia protein. Cancer Cell 18, 258-267.

Zhang, J., Wu, T., Simon, J., Takada, M., Saito, R., Fan, C., Liu, X.D., Jonasch, E., Xie, L., Chen, X., et al. (2018). VHL substrate transcription factor ZHX2 as an oncogenic driver in clear cell renal cell carcinoma. Science 361, 290-295.

Zhang, Y., Ng, K.S., Kucherlapati, M., Chen, F., Liu, Y., Tsang, Y.H., Velasco, G.D., Kang, J.J., Akbani, R., and Hadjipanayis, A. (2017). A Pan-Cancer Proteogenomic Atlas of PI3K/AKT/mTOR Pathway Alterations. Cancer Cell 31, 820.
X.R., Shi, Y.H., Wang, Z., et al. (2019). Genomic sequencing identifies WNK2 as a driver in hepatocellular carcinoma and a risk factor for early recurrence. J Hepatol 71, 1152-1163.
Figure legends

**Figure 1.** Identification of candidate ESMs in KIRC by statistical analyses. (A) Schematic of this study. (B) Illustration of base substitutions. (C) Schematic showing the archaeal and eukaryotic exosome complexes viewed from the top. (D) Summary statistics for the 6 types of c-substitutions in KIRC. (E) Scatter plots showing the correlation between the rank of mutation and gene expression. Each plot represents one KIRC sample. The orange dashed line shows the best fit for visualization. P values were calculated by spearman’s rank correlation. (F) Volcano plots of p and r values calculated by spearman correlation analyses. Each plot represents one gene. The top 1% of genes were taken as candidates and marked in red. (G) Box plots showing ASNS expression in the high and low C>A/G>T mutation groups. The expression was normalized to TBP. (H) Volcano plots showing the p and fold change (FC) values calculated by the two-tailed student’s t-test. Each plot represents one gene. FC was calculated by the formula: FC = the mean gene expression in the high group/that in the low group. The top 1% genes were taken as candidates and marked in red. (I) C-substitution mutation frequencies in 5 types of major cancers. (J, K) Mutation spectra of the TP53 gene in BRCA (J) and VHL gene in KIRC (L). (L) Kaplan-Meier (KM) analyses of OS between VHL C>A/G>T and C>T/G>A mutation groups. The median OSs in the C>A/G>T and C>T/G>A groups were 72.95 and 108.91 months, respectively. The p value was obtained from the log-rank test. (M) Venn diagram showing the overlap of the candidate genes identified by three types of statistical analyses as noted.
Figure 1

A

- Genomic mutation analysis
- VHL mutation analysis
- Overlap
- Mutator screening in E. coli
- Nuclease activity determination
- Mutator screening in vivo
- Clinical relevance of EXOSC1 in KIRC

B

- Adenine
- Cytosine
- Guanine
- Thymine
- Transitions vs. Transversions
- Purines vs. Pyrimidines

C

Archaeal
Eukaryotic

D

|                  | Total | Missense | Sense |
|------------------|-------|----------|-------|
| Number of patients| 532   | 532      | 532   |
| C>T/G>A (mean)   | 19194 (19.2) | 6462 (12.0) | 3792 (7.1) |
| C>A/G>T (mean)   | 5461 (10.3)  | 4272 (8.0)  | 1189 (2.2)   |
| A>G/T/C (mean)   | 4096 (0.4)   | 3219 (6.1)  | 1777 (3.3)   |
| A>T/T>A (mean)   | 4260 (6.6)   | 3571 (8.7)  | 689 (1.3)    |
| C>G/G>C (mean)   | 3121 (5.9)   | 2542 (4.8)  | 570 (1.1)    |
| A>C/T>G (mean)   | 2222 (4.2)   | 1805 (3.4)  | 417 (0.8)    |
| Mutation (mean)  | 42429 (79.8) | 33986 (63.9) | 6443 (15.9) |

E

- GAPDH
- CRB3

F

- Range of Expression vs. Rank of C>A/G>T mutation

G

- ASNS

H

- Normalized Expression

I

- TTV frequency (%)

J

- TP53 in BRCA

K

- VHL in KIRC

L

- VHL

M

- C>A/G>T
- C>T/G>A
- A>G/T<C
- A>T/T>A
- C>G/G>C
- A>C/T>G

CCNB1
EXOSC1
RAB51F
Figure 2. EXOSC1 promotes mutations in *E. coli*. (A) Western blot showing His-tagged protein levels in *E. coli*. (B, C) The frequencies of the Rif<sup>R</sup> mutation in the *E. coli* cells expressing candidate ESMs (B) or exosome family members (C). The vector and AID were used as negative and positive controls, respectively. Each plot represents the mutational frequency of an independent overnight culture (n=30). Median mutational frequency of the gene is noted. (D) Frequencies of the Rif<sup>R</sup> mutation in the *E. coli* cells treated with and without IPTG (n=30). (E, F) Representative images of *E. coli* cells treated with (E) and without (F) IPTG. (G) The mutational distribution in 25 independent Rif<sup>R</sup> colonies transformed by vector or EXOSC1. (H, I) Summary of the c-substitutions in Rif<sup>R</sup> colonies transformed by vector (H) and EXOSC1 (I). (J) The mutational frequencies of each c-substitution in Rif<sup>R</sup> colonies. The *p* value was calculated by fisher's exact-test.
Figure 2

A

![Image of gel electrophoresis with bands labeled AID, CDK5, TARBP2, EXOSC1, RAB5IF, CCNB1, PSAT1, NCEAB3, 100KD, 55KD, 35KD, 25KD]

B

![Graph showing mutants/10^9 cells for different conditions: Vector, AID, CDK5, TARBP2, EXOSC1, RAB5IF, CCNB1, PSAT1, NCEAB3, IPTG+]

C

![Graph showing mutants/10^9 viable cells for different conditions: Vector, AID, CDK5, TARBP2, EXOSC1, RAB5IF, CCNB1, PSAT1, NCEAB3, IPTG+]

D

![Graph showing mutants/10^9 cells for different conditions: Vector, AID, CDK5, TARBP2, EXOSC1, RAB5IF, CCNB1, PSAT1, NCEAB3, IPTG-]

E

![Images of plates with different conditions: IPTG+, RIF+, Vector, AID, EXOSC1, EXOSC3]

F

![Images of plates with different conditions: IPTG-, RIF-, Vector, AID, EXOSC1, EXOSC3]

G

![Sequence alignment for Vector and EXOSC1]

H

| From | A  | G  | C  | T  |
|------|----|----|----|----|
| A    | 4  | 2  | 2  | 2  |
| G    | 1  |    |    | 2  |
| C    | 1  | 12 |    |    |
| T    | 2  | 2  | 4  |    |

I

| To   | A  | G  | C  | T  |
|------|----|----|----|----|
| EXOSC1 | 1 | 2 | 2 | 2 |

J

![Bar graph showing mutation frequency for different conditions: A-G/C-T, A-T/C-G, A-G/T-C, A-T/G-C, C-G/T-A, C-T/G-A]
Figure 3. EXOSC1 cleaves ssDNA. (A) Coomassie blue staining of in vitro purified EXOSC1 protein. (B) Schematic of synthetic DNA substrates. (C) In vitro cleavage assays of EXOSC1 using generic ssDNA, dsDNA and DNA-RNA hybrid as substrates. The assays were performed in the presence of 1 μM oligonucleotides, 1 μM EXOSC1, 70 mM KCl, 700 μM MgCl₂, 1 mM DTT, and 20 mM Tris-HCl pH 7.0. After incubation at 37°C for 4 h, resultant samples were analyzed by 15% polyacrylamide TBE-urea gels. (D) Cleavage assays of EXOSC1 in the presence or absence of EXOSC3, EXOSC6, EXOSC7 and EXOSC8 using generic ssDNA as substrates. (E) Cleavage assays in the presence of the components as noted. (F) Cleavage assays of EXOSC1 at the pH as noted. (G) Time course cleavage assays of EXOSC1 using generic ssDNA as substrates. (H) Rate curve of EXOSC1 cleavage at 37°C and pH 7.0.
**Figure 4.** EXOSC1 prefers to cleave C sites in ssDNA. (A, B) Cleavage assay of EXOSC1 using unbiased DNA, dsDNA and DNA-RNA hybrid as substrates. (C, D) Cleavage assays of EXOSC1, EXOSC3, EXOSC7 and EXOSC8 using unbiased ssDNA as substrates. (E) Mass spectrometry (MS) analyses of the resultant mixtures described in (B). C, G, A and T were detected by MS using nucleoside to base ion mass transitions of 228.1 to 112.2 (C), 268.1 to 152.1 (G), 252.2 to 136.1 (A), and 243.1 to 127.2 (T) (Supplementary Figure 3E). Standard curves were generated by a serial dilution of C, G, A and T (Supplementary Figure 3F). Free C, G, A and T contained in the reaction mixtures were quantified by standard curves. (F) Schematic showing that the consequence of C>A mutation in the coding strand is distinct from that in the template strand. (G) Correlation between EXOSC1 expression and the c-substitution mutation as noted in KIRC. Each line represents one best fit for visualization. P values were from spearman’s rank correlation. P and r values of C>A mutations were 0.0001 and 0.19, respectively; C>A/G>T: p = 0.0006, r = 0.17; G>T: p = 0.0271, r = 0.10; total mutations: p = 0.0594, r = 0.09; C/G>G/C: p = 0.5730, r = 0.03. (H) Student’s t-tests analyses of the expression difference of EXOSC1 between the high and low mutation groups. FC = the mean gene expression in the high group/that in the low group.
Figure 5. EXOSC1 enhances DNA damages and mutations in KIRC Cells. (A) Representative fluorescent images of γ-H2AX foci in 769-P cells transfected with control (pCDH, empty vector) or pCDH-Flag EXOSC1 plasmids for 2 days. Scale bar = 10 μm. (B) Percentage of cells with more than 20 γ-H2AX foci in the KIRC cells transfected with control or pCDH-Flag EXOSC1 plasmids for 2 days. (C) Percentage of cells with more than 20 γ-H2AX foci in the KIRC cells infected with lentivirus encoding shRNAi control (pLKO scramble) or pLKO sh-EXOSC1 for 2 days. (D) Representative images of DNA comets in 769-P cells transfected with control or pCDH-Flag EXOSC1 plasmids for 2 days. (E) Comet tail moment of the 769-P and TUHR14TKB cells transfected with control or pCDH-Flag EXOSC1 plasmids for 2 days. (F) 3D-PCR and subsequent sequencing analyses of the VHL mutations in the TUHR14TKB cells stably expressing control (vector) or EXOSC1 (pCDH-Flag EXOSC1, EXOSC1-OE). (G) 3D-PCR analyses of VHL in TUHR14TKB cells stably expressing control (pLKO.1 vector) or shRNA against EXOSC1 (pLKO shEXOSC1-1, EXOSC1-KD). (H) 3D-PCR analyses of VHL in TUHR14TKB cells stably expressing control, EXOSC1-OE and/or shRNA against XRCC1 (pMKO.1 shXRCC1-1, XRCC1-KD). (I) 3D-PCR analyses of VHL in TUHR14TKB cells stably expressing control (pLKO.1 vector), shRNA against EXOSC1 (pLKO shEXOSC1-1, EXOSC1-KD) and/or shRNA against XRCC1 (pMKO.1 shXRCC1-1, XRCC1-KD). (J) Schematic showing the subcutaneous xenograft tumor models. The 5 x 10⁶ control, EXOSC1-OE and EXOSC1-KD 769-P cells were implanted subcutaneously. After 2 weeks, the resultant tumors were analyzed by 3D-PCR. (K) 3D-PCR analyses of VHL in the tumors described in (J).
Figure 5

A

Flag  H2AX  DAPI  Merged

Control

EXOSC1-OE

B

769-P

TUHR14TKB

Y-H2AX* cells (%)

Vector  EXOSC1-OE  Vector  EXOSC1-OE

C

769-P

TUHR14TKB

Y-H2AX* cells (%)

Vector  EXOSC1-KD  Vector  EXOSC1-KD

D

EXOSC1-OE

E

769-P

TUHR14TKB

Comet tail moment

Vector  EXOSC1-OE  Vector  EXOSC1-OE

F

VHL

90°C  87°C

Vector  EXOSC1-OE

C>A/G>T per sequence

G

VHL

90°C  87°C

Vector  EXOSC1-OE  EXOSC1-OE (XRCC1-KD)

H

VHL

90°C  87°C

Vector  EXOSC1-KD

I

VHL

90°C  87°C

Vector  EXOSC1-KD  (XRCC1-KD)

EXOSC1-OE

J

EXOSC1-OE cells

Tumor

Vector cells

EXOSC1-KD cells

Tumor

3D-PCR

K

VHL

90°C  87°C

Vector  EXOSC1-OE

EXOSC1-KD

EXOSC1-OE

EXOSC1-KD  XRCC1-KD
**Figure 6.** EXOSC1 sensitizes KIRC cells to PARP inhibitor. (A, B) KM analyses of DFS (A) and OS (B) in KIRC patients with different EXOSC1 levels. *P* values were obtained from the log-rank test. (C) KM analysis of OS in KIRC patients with different EXOSC1 expression levels and VHL mutations. (D) Colony formation of control and EXOSC1-OE TUHR14TKB cells treated with 600 nM niraparib. (E, F) Clonogenic survival of control and EXOSC1-OE TUHR14TKB (E) and 769-P (F) cells treated with niraparib. (G, I) Tumor volumes of 769-P (G) and Caki-2 (I) xenografts treated with or without niraparib. *n* = 4 groups; *n* = 6 mice/group; ± SEM. (H, J) Body weight change percentage of 769-P (H) and Caki-2(J) xenografts treated as described in (G).
Figure 6

A

B

C

D

E

F

G

H

I

J
Supplementary Figure Legends

Figure 1—figure supplement 1. Identification of candidate ESMs in KIRC by statistical analyses. (A and B) Mutation spectra of TP53 (A) and VHL (B) normalized by the nucleotide abundance.
Figure 2—figure supplement 1. EXOSC1 promotes mutations in E. coli. (A) Setup plot showing the overlap numbers of the candidate genes of the 6 types of c-substitutions. More details can be found in Supplementary Table S4. (B) Western blot showing the His-tagged protein levels of the exosome complex members in the E. coli cells. (C) Growth rates of control and EXOSC1-transformed cells in the presence of IPTG. n = 5. (D) Frequencies of c-substitutions in Rif\(^{R}\) colonies were normalized to the nucleotide abundances. The \(p\) value was calculated by fisher's exact test.
Figure 3—figure supplement 1. EXOSC1 cleaves ssDNA. (A) In vitro cleavage assays of EXOSC1 homologs (EXOSC2–EXOSC9) using generic ssDNA, dsDNA and DNA-RNA hybrid as substrates. (B) Cleavage assays of EXOSC1 in the presence/absence of EXOSC2, EXOSC4, EXOSC5 or EXOSC9.
**Figure 4—figure supplement 1.** EXOSC1 prefers to cleave C sites in ssDNA. (A) Cleavage assays of EXOSC2-EXOSC9 using unbiased ssDNA, dsDNA and DNA-RNA hybrid as substrates. (B) Time course cleavage assays of EXOSC1 using unbiased ssDNA as substrates. (C) Base ion mass transitions for LC-MS/MS analyses of C, G, A and T. Free nucleosides were detected using nucleoside to base ion mass transitions of 228.1 to 112.2 (C), 268.1 to 152.1 (G), 252.2 to 136.1 (A), and 243.1 to 127.2 (T). (D) LC-MS/MS standard curves of nucleoside as noted. (E and F) Frequencies (E) and normalized frequencies (F) of 12 types of substitutions in Rif\(^R\) colonies. \(P\) values were calculated using fisher's exact test. (G) Correlation between EXOSC1 expression and the mutation as noted in KIRC. Each plot represents one KIRC sample. (H) Student’s \(t\)-test analyses of EXOSC1 expression difference between the high and low mutation groups. The 532 KIRC patients were grouped into 2, 3, 4 or 5 groups to evaluate the impact of group number. (I and J) Summary statistics for the mutation as noted in KIRC. \(P\) values were calculated by the two-tailed student’s \(t\)-test.
EXOSC1 enhances DNA damages and mutations in KIRC cells. (A) Western blot analyses of EXOSC1 in stable control, enhanced EXOSC1, and EXOSC1 knockdown cells. (B) Quantitative PCR analyses of EXOSC1 mRNA in the 769-P cells stably expressing shRNA (pLKO EXOSC1-1 or pLKO EXOSC1-2) against EXOSC1. (C) Representative fluorescence images of γ-H2AX foci in the 769-P cells stably expressing shRNA against control (empty vector) or EXOSC1 (pLKO EXOSC1-1, EXOSC1-KD). Scale bar = 10 μm. (D and F) Western blot analyses of EXOSC1 and XRCC1 protein levels in the stable cells as noted. (E) Quantitative PCR analyses of the mRNA in the cells stably expressing shRNA against EXOSC1 and/or XRCC1.
Figure 6—figure supplement 1. EXOSC1 sensitizes KIRC cells to PARP inhibitor. (A) The fragments per kilo base per million mapped reads (FPKM) of the noted gene in 532 KIRC. Each dot represents one KIRC patient. (B) Best separation KM analyses of OS in KIRC patients with different EXOSC1 levels. The median OS (the low vs high EXOSC1 group) = 106.40 vs 71.91 months. (C) Summary statistics for Figure 6C. (D) Clonogenic survival of control/EXOSC1-OE SNU-1272 and Caki-2 cells treated with niraparib.
Figure 4

A

5'-TTAGGCTTAAGGCCTAAAGCC-3'

ssDNA dsDNA DNA+RNA hybrid

EXOSC1 - + - + - +

50 nt

25 nt

5 nt

B

5'-TTAGGCTTAAGGCCTAAAGCC-3'

No prot EXOSC1 EXOSC3 EXOSC7 EXOSC8

hr 0 4 4 4 4 0 4 4

50 nt

25 nt

5 nt

C

5'-TTAGGCTTAAGGCCTAAAGCC-3'

ssDNA dsDNA DNA+RNA hybrid

EXOSC1 - + - + - +

50 nt

25 nt

5 nt

D

5'-TTAGGCTTAAGGCCTAAAGCC-3'

No prot EXOSC1 EXOSC3 EXOSC7 EXOSC8

hr 0 4 4 4 4 0 4 4

50 nt

25 nt

5 nt

E

Unbiased sequence

dC
dG
dA
dT

Standard ssDNA + EXOSC1 ssDNA + EXOSC1 ssDNA + EXOSC1 ssDNA + EXOSC1

Control Control Control Control

RT

RT

F

Unbiased sequence

dC
dG
dA
dT

Standard ssDNA + EXOSC1 ssDNA + EXOSC1 ssDNA + EXOSC1 ssDNA + EXOSC1

Control Control Control Control

RT

RT

G

5 Groups

C>A

G>T

C>A/G>T

C>G/G>C

2 Groups

3 Groups

4 Groups

5 Groups

Rank of Expression

Rank of Mutation

G>T

C>A

Rank of Expression

Rank of Mutation

G>T

C>A

-1.0 1.0 2.0 3.0

-log_{10}(p-value)

1.10 1.25 1.40

FC
Figure 1—figure supplement 1

A

TP53 in BRCA

B

VHL in KIRC

| Mutation          | BRCA Mutations per Patient | KIRC Mutations per Patient |
|-------------------|---------------------------|---------------------------|
| C>A/G>T           |                           |                           |
| C>T/G>A           |                           |                           |
| A>G/T>C           |                           |                           |
| A>T/T>A           |                           |                           |
| C>G/G>C           |                           |                           |
| A>C/T>G           |                           |                           |

Mutations per patient
Figure 2—figure supplement 1

A

Number of gene

45 40 35 30 25 20 15 10 5 0

A>C/T>G
A>G/T>C
A>T/T>A
C>A/G>T
C>G/G>C
C>T/G>A

B

IPTG
63 KD → 48 KD → 35 KD → 25 KD →

EXOSC1
EXOSC2
EXOSC3
EXOSC4
EXOSC5

EXOSC6
EXOSC7
EXOSC8
EXOSC9

C

Viable cells per mL

Vector
Exosc1

Time (hr)

0 5 10 15 20 25

6×10^9
4×10^9
2×10^9
0

D

Normalized mutation frequency (%)

Vector
Exosc1

A>G/T>C
A>C/T>G
A>T/T>A
C>A/G>T
C>G/G>C
C>T/G>A

p 1.47e-06

****
Figure 3—figure supplement 1

A

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC2 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC3 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC4 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC5 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC6 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC7 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC8 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

| ssDNA | dsDNA | DNA+RNA hybrid |
|-------|-------|----------------|
|      |      |                |
| EXOSC9 |      |                |
| 50 nt |      |                |
| 25 nt |      |                |
| 5 nt  |      |                |

B

| No prot | EXOSC1 | EXOSC2 | EXOSC1+EXOSC2 |
|---------|--------|--------|---------------|
| 50 nt   |        |        |               |
| 25 nt   |        |        |               |
| 5 nt    |        |        |               |

| No prot | EXOSC3 | EXOSC4 | EXOSC1+EXOSC4 |
|---------|--------|--------|---------------|
| 50 nt   |        |        |               |
| 25 nt   |        |        |               |
| 5 nt    |        |        |               |

| No prot | EXOSC5 | EXOSC6 | EXOSC1+EXOSC6 |
|---------|--------|--------|---------------|
| 50 nt   |        |        |               |
| 25 nt   |        |        |               |
| 5 nt    |        |        |               |

| No prot | EXOSC7 | EXOSC8 | EXOSC1+EXOSC8 |
|---------|--------|--------|---------------|
| 50 nt   |        |        |               |
| 25 nt   |        |        |               |
| 5 nt    |        |        |               |

| No prot | EXOSC9 | EXOSC10 | EXOSC1+EXOSC9 |
|---------|--------|---------|---------------|
| 50 nt   |        |         |               |
| 25 nt   |        |         |               |
| 5 nt    |        |         |               |
