Site specific target binding controls RNA cleavage efficiency by the Kaposi’s sarcoma-associated herpesvirus endonuclease SOX

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ABSTRACT

A number of viruses remodel the cellular gene expression landscape by globally accelerating messenger RNA (mRNA) degradation. Unlike the mammalian basal mRNA decay enzymes, which largely target mRNA from the 5′ and 3′ end, viruses instead use endonucleases that cleave their targets internally. This is hypothesized to more rapidly inactivate mRNA while maintaining selective power, potentially though the use of a targeting motif(s). Yet, how mRNA endonuclease specificity is achieved in mammalian cells remains largely unresolved. Here, we reveal key features underlying the biochemical mechanism of target recognition and cleavage by the SOX endonuclease encoded by Kaposi’s sarcoma-associated herpesvirus (KSHV). Using purified KSHV SOX protein, we reconstituted the cleavage reaction in vitro and reveal that SOX displays robust, sequence-specific RNA binding to residues proximal to the cleavage site, which must be presented in a particular structural context. The strength of SOX binding dictates cleavage efficiency, providing an explanation for the breadth of mRNA susceptibility observed in cells. Importantly, we establish that cleavage site specificity does not require additional cellular cofactors, as had been previously proposed. Thus, viral endonucleases may use a combination of RNA sequence and structure to capture a broad set of mRNA targets while still preserving selectivity.

INTRODUCTION

Viral infection dramatically reshapes the gene expression landscape of the host cell. By changing overall messenger RNA (mRNA) abundance or translation, viruses can redirect host machinery towards viral gene expression while simultaneously dampening immune stimulatory signals (1–3). Suppression of host gene expression, termed host shut-off, can occur via a variety of mechanisms, but one common strategy is to accelerate degradation of mRNA (1–3). This occurs during infection with DNA viruses such as alphaherpesviruses, gammaherpesviruses, and vaccinia virus, as well as with RNA viruses such as influenza A virus and SARS and MERS coronaviruses (1,4,5). In the majority of these cases, a viral factor promotes endonucleolytic cleavage of target mRNAs. This strategy bypasses the normally rate limiting steps of deadenylation and decapping to effect rapid mRNA degradation by host exonucleases (1).

Virally encoded host shut-off endonucleases are usually specific for mRNA, yet broad-acting in that they target the majority of the mRNA population. This is exemplified by herpesviral nucleases, including the SOX endonuclease encoded by Kaposi’s sarcoma-associated herpesvirus (KSHV), an oncogenic human gammaherpesvirus that causes Kaposi’s sarcoma and B cell lymphoproliferative diseases (6,7). KSHV SOX is a member of the PD-(D/E)\(_{\text{\kappa}}\)K type II restriction endonuclease superfamily that possesses mechanistically distinct DNase and RNase activities (8–10). The RNase activity of the gammaherpesvirus SOX protein has been shown to play key roles in various aspects of the viral lifecycle, including immune evasion, cell type specific replication, and controlling the gene expression landscape of infected cells (11–14). However, the mechanism by which SOX targets mRNAs remains largely unknown.

Sequencing data indicate that within the mRNA pool there appears to be a range of SOX targeting efficiencies; some transcripts are efficiently cleaved in cells, while others are partially or fully refractory to cleavage (15–19). Additionally, SOX has been shown to cut within specific locations of mRNAs in cells, further emphasizing that there must be transcript features that confer selectivity (16,20). Indeed, a transcriptome-wide cleavage analysis indicated that SOX targeting is directed by a relatively degenerate motif, often containing an unpaired polyadenosine stretch.

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shortly upstream of the cleavage site, which is located in a loop structure (20). Cleavage within an unpaired loop was confirmed in a recent crystal structure of SOX with RNA, although additional contacts that could confer sequence specificity were not observed (21).

Thus, a major outstanding question is how RNA sequence and/or structure contribute to SOX target recognition. In this context, it is unclear how sequence features surrounding the RNA cleavage site might impact SOX targeting, for example by changing its affinity for a given RNA or the efficiency with which cleavage occurs. To address these questions, we sought to reconstitute the SOX cleavage reaction in vitro using purified components. Using an RNA substrate that is efficiently cleaved by SOX in cells, we revealed that specific RNA sequences within and outside of the cleavage site significantly contribute to SOX binding efficiency and target processing. In particular, we found that the polyadenosine stretch adjacent to the cleavage site is critical for SOX binding, and we experimentally verified the importance of an open loop structure surrounding the cleavage site. Finally, we demonstrated that this in vitro system faithfully recapitulates the initial endonucleolytic cleavage event that is an essential component of mRNA target specificity in vivo. Collectively, our data reveal that specific sequence features potently impact SOX binding, and thus provide key insight into the breadth of SOX targeting efficiency observed across the transcriptome. More broadly, this information provides a framework for better understanding the target specificity of endonucleases, which play central roles in mammalian quality control processes and viral infection outcomes.

MATERIALS AND METHODS

Recombinant protein expression and purification

KSHV SOX was codon optimized for Sf9 expression and synthesized from GENEWIZ. SOX was then subcloned using restriction sites BamHI and SalI (New England BioLabs) into pFastBac HTD. This vector was modified to carry a GST affinity tag and PreScission protease cut site as described (22). All SOX mutants were generated using single primer site-directed mutagenesis (23). Sequences were validated using standard pGEX forward and reverse primers. Generation of viral bacmids and transfections were validated using standard pGEX forward and reverse priming, and resuspended in RNase-free ddH2O.

Ribonuclease assays

Ribonuclease assays

$K_{\text{obs}}$ and Hill coefficients of SOX were determined from the cleavage kinetics of $[^{32}\text{P}]$-labeled RNA substrates as previously described (23). Briefly, 1 μl (≤1 pM) of $[^{32}\text{P}]$-labeled RNA was added to 9 μl of premixture containing 20 mM HEPEs pH 7.1, 70 mM NaCl, 2 mM MgCl2, 1 mM TRCEP, 1% glycerol, and increasing concentrations of purified SOX. Reactions were performed at room temperature under single turnover conditions, and quenched at the indicated time intervals with 8 μl stop solution (10 M urea, 0.1% SDS, 0.1 mM EDTA, 0.05% xylene cyanol, 0.05% bromophenol blue). Samples were resolved by 15% urea–PAGE, imaged using a Typhoon variable mode imager (GE Healthcare), and quantified using ImageQuant and GelQuant software packages (Molecular Devices). The data were plotted and fit to exponential curves using Prism 7 software package (GraphPad) to determine observed rate constants.

A FRET probe with excitation at 646 nM and emission at 660 nM (LIMDI 54 Flo) was purchased from Dharmacon (Supplementary Table S1). The RNA FRET probe was
added at a final concentration of 100 nM to 9 μl of pre-mixture containing 20 mM HEPES pH 7.1, 70 mM NaCl, 2 mM MgCl₂, 1 mM TCEP, 1% glycerol with 2 μM of SOX (23). Terminator 5’ exonuclease (Lucigen) was added to reactions using a 1:1000 dilution of the enzyme. Reactions were quenched at indicated time intervals with equal volumes of stop solution containing 95% formamide and 10 mM EDTA, then resolved using urea-PAGE and visualized using a Typhoon variable mode imager (GE Healthcare). The data were plotted using Prism 7 software package (GraphPad). All experiments were repeated >3 times and mean values were computed.

For assays designed to detect endonucleolytic cleavage intermediates, 1 μl of labeled RNA substrate was combined with 9 μl of reaction solution (20 mM HEPES pH 7.1, 70 mM NaCl, 0.200 mM CaCl₂, 0.700 mM MgCl₂, 1% glycerol, 0.5 mM TCEP) in the presence or absence of 2 μM SOX for 10 min at room temperature. RNA was then ethanol precipitated, resuspended in 95% formamide solution containing 10 mM EDTA, and resolved on a 12% urea-PAGE analytical grade sequencing gel together with a ss-RNA Decade ladder (Ambion Life Technologies) for 1.5 h at 22 W before imaging as described above.

In-line probing

The sequence surrounding the cut site in LIMD1 was inserted into a pBSSK (−) backbone using the BamHI and XhoI restriction sites. Mutations were introduced by the Quickchange site directed mutagenesis protocol (Agilent). The 100 nt sequence surrounding the GFP cut site was inserted using the BamHI and XhoI restriction site.

In-line probing was performed as previously described (24). Briefly, pBSSK(−) plasmids containing the indicated sequences (see Supplementary Table S2) were linearized by digestion with XhoI and ScaI for GFP or BlpI and SacI (NEB) for LIMD1, gel purified, phenol/chloroform extracted, and ethanol precipitated. The fragments were then used as templates for in vitro transcription with the HiScript T7 High Yield RNA synthesis Kit (NEB) and afterwards subjected to Turbo DNase (Ambion Life Technologies) treatment. RNA was resolved by 8% Urea PAGE, and full length transcripts were excised from the SYBR Gold stained gel (Thermo Fisher Scientific), eluted overnight in G50 buffer (20 mM Tris HCl pH 7.5, 300 mM NaOAc, 2 mM EDTA, 0.2% SDS), phenol/chloroform extracted, and ethanol precipitated. The RNA (~40 pmol) was dephosphorylated using shrimp alkaline phosphatase (rSAP, NEB), labeled with 1 μl [γ-32P]-ATP (150 mCi/ml) using USB Optokinase (Affymetrix), then gel purified as described above and dissolved in 20 μl of nuclease free water. For the in-line probing reaction, 1 μl RNA (≥20 000 cpm) was incubated in 2× reaction buffer (100 mM Tris-HCl pH 8.3, 40 mM MgCl₂, 200 mM KCl) at room temperature for 24 or 48 h. The reaction was quenched with 2× loading buffer (10 M urea, 1.5 mM EDTA pH 8.0). To generate ladders, 1 μl of the purified RNA was separately subjected to hydrolysis using the Next Magnesium RNA Fragmentation module (−OH) or RNase T1 digestion (T1) (NEB). Reactions were resolved by 8% urea–PAGE, exposed on a phosphorimager screen, and scanned using the Storm 820 imaging system (GE Healthcare). Deduced RNA structures were drawn using the RNA secondary structure visualization tool forna (Vienna RNA Web Services).

Electrophoretic mobility shift assays (EMSA)

RNA probes used in EMSA experiments were radiolabeled using the protocol described for ribonuclease activity assays. Reactions were incubated at RT for 30 min in buffer containing 20 mM HEPES pH 8.0, 30 mM KCl, 5 mM CaCl₂, 0.01% Tween-20, 0.5 TCEP, 0.2 mg/ml BSA (Sigma-Aldrich), 40 μg/ml of yeast tRNA (Ambion Thermo Fisher), and the indicated amount of purified SOX protein. Calcium Chloride was used in these binding assays to prevent substrate processing and stabilize RNA–protein interactions. Reactions volumes were kept at 10 μl and stopped with 3 μl 7× EMSA loading dye (70 mM HEPES pH 8.0, 420 mM KCl, 35% glycerol). Reactions were resolved by 8% native PAGE, and gels were imaged on a Typhoon multivariable imager (GE Healthcare) and quantified using GelQuant software package (Molecular Dynamics).

SOX RNA footprinting assay

LIMD1-54 RNA was 5′ end labeled with γ-[32P]-ATP-6000 Ci/mmol 150mCi/ml (PerkinElmer) using T4 PNK (New England BioLabs). RNA was then gel purified as stated previously. EMSA gel shifts were first used to determine optimal binding conditions (>90% binding, homogeneous complexes of RNA-protein). Binding buffer contained 0.01% Tween 20 (Sigma-Aldrich), 5 mM CaCl₂, 5 mM KCl, 50 mM NaCl, 0.5 mM TCEP, 20 mM HEPES pH 8.0, 0.04 mg/ml yeast tRNA (Ambion), 0.2 mg/ml nuclease free Bovine Serum Albumin (BSA) (Ambion). A dilution series of SOX (8–0.5 μM) was incubated with 1 μl of radiolabeled LIMD1-54 in the presence of 0.1 unit of RNase T1 (Epistem Illumina). Reactions were incubated at RT for a total of 10 min before being ethanol precipitated. RNA pellets were then resuspended in 5 μl of 95% formamide solution containing 10 mM EDTA and boiled for 5 min. Samples were then loaded on a 10% analytical grade urea–PAGE gel and run at 22 W for 1.5 h. Gels were imaged and analyzed as stated above. In order to produce an RNase T1 ladder, 1 μl of LIMD1-54 was incubated with 0.1 units of RNase T1. Reactions were incubated at RT for 5 min before being quenched and prepared as stated previously. The LIMD1-54 hydrolysis ladder was generated as stated in the in-line probing methods.

Bio-layer interferometry (BLI) real-time binding kinetics

RNA probes (3′ end labeled with biotin) were synthesized from Dharmacon (GE healthcare) and HPLC and PAGE purified (See Supplementary Table S1). The Octet RED96e Bio-Layer Interferometry instrument and Streptavidin (SA) Biosensors were available from ForteBio (Menlo Park, CA, USA). All steps were performed in reaction buffer similar to EMSA binding conditions. Biosensors were incubated with 200 nM of the biotinylated RNA substrate for 150 sec and free RNA was washed away in EMSA buffer.
containing no RNA. SOX protein was incubated with the 
RNA conjugated biosensors for 200–400 s in order to reach 
saturation. Indicated protein concentrations for each bio 
sensor are located on corresponding binding curves. Com-
plexes were dissociated for minimum of 15 min. Response 
curves for each biosensor were normalized against biosen-
sors conjugated to RNA in the absence of SOX (buffer only 
control). Normalized response curves were processed using 
Octet Software version 7 by fitting the group of selected bio-
sensors to a nonlinear regression model (25). Dissociation 
constants ($K_d$) were determined from $k_{on}$ and $k_{dis}$ values de-
duced from the fitted curves. A complete table of all values 
is provided in Supplementary Table S2.

RESULTS

KSHV SOX cleaves RNA substrates endonucleolytically as 
a monomer

In cells, the mRNA fragments resulting from the primary 
SOX endonucleolytic cleavage are predominantly cleared by 
the host 5′-3′ exonuclease XRN1, while in vitro, RNA frag-
ments are rapidly degraded by 5′-3′ exonuclease activity 
intrinsic to purified SOX (9). Thus, it has been challenging to 
analyze the initial endonucleolytic cleavage event that is 
an essential component of mRNA target specificity in vivo. 
Here, we sought to develop a biochemical system to address 
these questions.

Our prior analysis of SOX targets in cells identified the human 
*LIMD1* mRNA, which codes for a protein essential for 
P body formation and integrity, as being highly suscep-
tible to cleavage by SOX (20). The minimum sequence re-
quired to directly cut the putative cleavage site in *LIMD1* 
in cells was mapped to a 54-nucleotide segment (*LIMD1*- 
54), and we therefore chose this as our model substrate to 
study SOX targeting in vitro (20). We first expressed and pu-
rified KSHV SOX to greater than 95% purity from Sf9 in-
sect cells (Supplementary Figure S1A). Using the *LIMD1*-54 
substrate, we plotted the observed rate constant ($k_{obs}$) 
as a function of SOX concentration, yielding a Hill coeffi-
cient of $n = 1.1$ (Figure 1A). Thus, in agreement with previ-
ous observations (9,10), SOX appears to function predom-
nantly as a monomer. Under conditions of half maximal 
activity (2 μM; Figure 1A), SOX displayed a strong prefer-
ce for the ‘hard’ divalent metal Mg$^{2+}$ and a weaker prefer-
ce for the ‘softer’ and larger metals Mn$^{2+}$, Co$^{2+}$ and Zn$^{2+}$ 
(Figure 1B). This is again consistent with other character-
ized members of the P/DDexK family of enzymes (9,26). No-
tably, SOX activity in the presence of Mg$^{2+}$ was inhibited in 
a dose-dependent manner upon competitive addition of 
Ca$^{2+}$ (Figure 1C and Supplementary Figure S1D). This is 
likely the result of increased coordination partners engaged 
by Ca$^{2+}$, which decreases the ability of catalytic residues to 
promote proper base hydrolysis (27–29). Finally, increasing 
the NaCl concentration above 100 mM led to substantially 
decreased SOX activity (Figure 1D), in accordance with the 
observation that high salt concentrations frequently inhibit 
nuclease activity by disrupting protein-protein or protein-
substrate interactions (29). Given that recombinant SOX 
displays robust 5′-3′ exonuclease activity (9,10), we sought 
to confirm that *LIMD1*-54 was subject to endonucleolytic 
SOX cleavage, as this is the predominant event that directs 
mRNA turnover in SOX expressing cells (3,16). Both the 5′ 
and 3′ ends *LIMD1*-54 were blocked by capping the 5′ 
end with a Cy5 fluorophore and the 3′ end with an Iowa 
Black quencher (*LIMD1*-54 Flo). We confirmed this RNA 
was resistant to degradation by the 5′-phosphate dependent 
exonuclease terminator (Figure 1E, lane 3). However, in the 
presence of SOX, a cleavage product was observed that cor-
related with an endonucleolytic cut (Figure 1E, lane 2).

To confirm this processing event was not a result of 
contamination, we purified a SOX mutant containing mu-
tations within two key residues of the SOX active site 
(D221N/E244Q). Incubation of this mutant with *LIMD1*- 
54 over the course of 1.5 h yielded no RNA cleavage (Sup-
plementary Figure S1E). Thus, recombinant SOX appears 
to target *LIMD1*-54 for endonucleolytic cleavage in vitro, as 
has been observed for this substrate in cells.

KSHV SOX shows RNA substrate selectivity in vitro

To analyze RNA substrate selectivity using our in vitro as-
day, we first compared SOX degradation of *LIMD1*-54 to a 
51-nucleotide sequence of the mRNA encoding GFP (*GP-
F'). We have previously shown that GFP mRNA is cleaved 
by SOX in cells, and that GFP-51 is the minimal sequence 
required to elicit cleavage (15,16). The cleavage sites for 
*LIMD1*-54 and GFP-51 are predicted to occur in an open 
loop region (Figure 2A, red arrow). Upon direct compar-
ison of these two RNAs, we observed a ∼6-fold increase in 
the catalytic efficiency of SOX for the *LIMD1*-54 sub-
strate compared to GFP-51 (Figure 2B). This difference was 
not exclusively due to the fact that the GFP substrate was 
slightly shorter than *LIMD1*-54, as SOX also displayed a 5-
fold reduction of catalytic efficiency on a longer, 100 nt GFP 
substitute (GFP-100; Figure 2B). Electrophoretic mobility 
shift assays (EMSA) further revealed a 10-fold increase in 
SOX binding to *LIMD1*-54 compared to GFP-51 (Figure 2C). 
Given that both substrates contain the requisite un-
paired bulge at the predicted cleavage site (see Figure 2A 
and Supplementary Figure S2), these observations suggest 
that additional sequence or structural features impact SOX 
targeting efficiency on individual RNAs.

Two SOX point mutants, P176S and F179A, located in 
an unstructured region of the protein that bridges domains 
I and II have been shown to be selectively required for its 
endonucleolytic processing of RNA substrates (Supplemen-
tary Figure S3A and S3B) (8,21). Structural data indicate 
that residue F179 forms a stacking interaction with an ade-
nine base in the RNA, likely stabilizing the protein-RNA in-
teraction, while P176 is hypothesized to contribute to struc-
tural rearrangements required for F179 engagement (21). 
We purified both mutants to evaluate their relative RNA 
processing and RNA binding activity against the optimal 
*LIMD1*-54 substrate. Both mutants displayed purity and 
elution profiles similar to wild type (WT) SOX (see Supple-
mentary Figure S1A–C). However, the catalytic efficiency 
of each mutant was >10-fold less than WT SOX (Figure 2D). 
Furthermore, RNA binding was severely perturbed; the 
binding kinetics of WT SOX for *LIMD1*-54 are in the 
single digit nanomolar range ($K_d = 7$ nM), while P176S and 
F179A display >2 log defects ($K_d = 702$ nM and 831 nM, 
respectively) (Figure 2E and Supplementary Figure S4A–
Figure 1. Kinetic characterization of recombinant SOX. (A) The observed rate constant ($k_{\text{obs}}$) was plotted as a function of SOX concentration using the 5′-32P-labeled LIMD1-54 RNA substrate, showing a hill coefficient ($n$) of 1.11. (B) The catalytic efficiency of SOX (2 µM) in the presence of MgCl$_2$, MnCl$_2$, CoCl$_2$, and ZnCl$_2$ was plotted as a function of cofactor concentration. (C) The impact of adding increasing concentrations of CaCl$_2$ on SOX-induced degradation of a 5′-32P-labeled LIMD1-54 RNA probe. Reactions were carried out in the presence of 0.7 mM MgCl$_2$. (D) SOX catalytic efficiency was determined under increasing concentrations of NaCl. (E) The 5′ and 3′ ends of LIMD-54 were blocked with Cy5 and Iowa Black, respectively, to prevent exonucleolytic degradation. Reactions were incubated for 30 min in the presence of SOX or, as a control, the Terminator 5′ exonuclease (EXO). Input refers to RNA in reaction buffer without enzyme.

C). Thus, the large defect in RNA binding likely explains the decreased efficiency of RNA processing. Notably, while there was a dramatic decrease in the relative affinities of the two mutants for LIMD1-54, there was not a complete loss of binding or RNA processing. This could be a result of secondary nonspecific interactions and/or nonspecific exonucleolytic degradation by SOX from the 5′ monophosphorylated end of the probe.

Secondary structure determination of the LIMD1-54 substrate

In silico RNA folding predictions of SOX targeting motifs, coupled with RNA mutagenesis experiments, have indicated that an RNA stem loop structure is an important determinant in SOX targeting both in vitro and in vivo (20,21). Given the importance of this predicted motif, and in partic-
C

0 = 2 µM

B

E0 = 2 µM

D

A

G

G

C

U

G

G

A

U

G

A

G

C

U

G

G

A

C

G

C

U

G

C

G

U

C

G

U

A

3'

5'

Predicted RNA fold

LIMD1 54

Predicted RNA fold

GFP 51

Figure 2. Substrate specificity, salt concentration and the bridge motif play important roles in SOX activity. (A) The predicted folding of the LIMD1-54 and GFP-51 RNAs was determined using mFold (35). Red arrows mark the predicted SOX cleavage site. (B) Catalytic efficiencies were determined for SOX (2 µM) in the presence of GFP-51, GFP-100 or LIMD1-54 substrates. Reactions were performed in triplicate. (C) Binding curves of SOX with GFP-51 and LIMD1-54 RNA. Percent binding of substrates was determined by EMSA, whereupon curves were fit to a single binding model from three independent measurements. (D) Catalytic efficiency of WT SOX or the host shutoff mutants P176S and F179A was determined at a constant enzyme concentration (2 µM) using a 5′ 32P-labeled LIMD1-54 RNA probe. Experiments were performed in triplicate. (E) EMSAs were used to determine percent binding of 5′ 32P-labeled LIMD1-54 RNA probe to WT SOX, P176S, and F179A. Curves were fit to a single binding model from three independent measurements.

ular the proposed requirement for unpaired sequence at the cut site, we sought to experimentally determine the structure of LIMD1-54 using chemical based in-line probing (Figure 3A). This showed that the LIMD1-54 structure contains a largely base paired stem region, followed by a loop at positions 15–27 that encompasses the predicted SOX cleavage site between nt 26 and 27, and a short hairpin structure at positions 29–40 (Figure 3B). Notably, some differences exist between the predicted and observed structures of LIMD1-54, including a larger loop region and the subsequent short stem-loop (compare Figure 3B to Figure 2A). However, in both cases the predicted cleavage site of SOX resides in a loop region.
SOX binds to a region encompassing the unpaired stretch of adenosine repeats

Recently, a high-resolution crystal structure was solved of SOX bound to a 31nt fragment of the KSHV premicroRNA K12-2 (K2-31). In this structure, the only observed contacts between SOX and K2-31 occurred between the four active site residues of SOX (Y373, R248, C247, F179) and the UGAAG motif surrounding the cleavage site of the RNA (21). It was therefore hypothesized that no other residues beyond this unpaired UGAAG motif were involved in transcript recognition (21). However, the binding affinity we observed for LIMD1-54 was 200-fold stronger than what was previously reported for K2-31 (21), suggesting that a more extended interaction surface might distinguish optimal from sub-optimal RNA substrates. We therefore used RNA footprinting to map the SOX binding sites on LIMD1-54. Indeed, SOX protected a region of LIMD1-54 that included the three adenosine stretch (positions 20–24) from RNase T1 digestion in a dose dependent manner (Figure 4). Notably, this mapped binding region is the same region predicted from in vivo PARE-seq data to be important for SOX targeting, although the reason for its importance remained unknown (20). We also observed a modest protection of base 27 (G) located directly adjacent to the predicted cleavage site of SOX, which represents the region detected in the crystal structure of K2-31 bound to SOX. Collectively, these findings suggest that while SOX may interact with residues directly adjacent to the cut site, a more extensive interaction interface exists for its preferred in vivo targets.

Base pairs surrounding the SOX binding and cleavage sites contribute to efficient substrate degradation

To explore the importance of the residues involved in SOX binding and cleavage, we engineered 3 mutants of the LIMD1-54 substrate (Figure 5A). First, we preserved the loop structure but replaced the three adenosines bound by SOX (residues 39–43) with guanosines (LIMD1-54 3xA-G). Second, we largely abolished the loop structure by providing complementary base pairing (LIMD1-54 Zipper). Third, we mutated the residue located at the predicted SOX cut site that was also protected in the footprinting assay (LIMD1-54 A-G). This mutant has been previously identified to block SOX cleavage in vivo (20). The predicted structures of the LIMD1-54 3xA-G and LIMD1-54 zipper mutants were verified by in-line probing (Figure 5). Supplemen-
Figure 4. SOX binds to a stretch of adenosines upstream of the cleavage site. (A) An RNA footprinting assay was carried out by incubating 5′-32P-labeled LIMD1-54 with RNase T1 in the presence (lanes 3–7) or absence (lane 2) of a dilution series of SOX (0.5–8 µM). Hydrolysis (–OH, lane 1) and RNase T1 (T1, lane 8) ladders of the RNA were also generated in order to map the location of protected sites. Lines on the right denote protected base pairs. (B) Diagram of LIMD1-54 indicating sites protected from RNase T1 cleavage by SOX. The upstream SOX binding site is colored orange while the protected residues surrounding the cut site are shown in red.

We next quantitatively measured the catalytic efficiency of SOX towards each of the above RNA substrates. Despite SOX having WT binding affinity for the predicted cleavage site mutant LIMD1-54 A-G, there was a 7-fold defect in binding site, we also engineered an additional zipper mutant (LIMD1-54 zipper 2) that did not disrupt the polyadenosine sequence. In agreement with the loop structure playing a critical role in target recognition, this LIMD1-54 zipper 2 mutant also displayed a substantial defect in binding ($K_d = 2.09$ µM; Supplementary Figure S7A, B, Supplementary Table S2). To rule out the possibility that the effect on binding affinity to the LIMD1-54 zipper mutant was a result of altered residues within the binding site, we also engineered an additional zipper mutant (LIMD1-54 zipper 2) that did not disrupt the polyadenosine sequence. In agreement with the loop structure playing a critical role in target recognition, this LIMD1-54 zipper 2 mutant also displayed a substantial defect in binding ($K_d = 2.09$ µM; Supplementary Figure S7A, B, Supplementary Table S2). Notably, the affinity of SOX for K2-31 was within the range of the LIMD1-54 structural mutants ($K_d = 1.08$ µM), suggesting that despite having an UGAAG motif upstream of a predicted bulge, this is unlikely to be a SOX target (Figure 5B, Supplementary Figure S5E and Supplementary Table S2).

We next quantitatively measured the catalytic efficiency of SOX towards each of the above RNA substrates. Despite SOX having WT binding affinity for the predicted cleavage site mutant LIMD1-54 A-G, there was a 7-fold defect in
Figure 5. Mutations that disrupt LIMD1-54 structure reduce SOX binding affinity. (A) Diagram showing the location of mutations made within LIMD1-54, with the WT binding site and cut site residues labeled in orange and red, respectively. Binding site mutants and cut site mutants are labeled in green. (B) The binding affinity of SOX for each of the LIMD1-54 mutants was tested in parallel using Bio-Layer Interferometry (BLI). (C) SOX catalytic efficiency was tested at a constant concentration (2 µM) for all of the LIMD1-54 mutant substrates. All assays were performed in triplicate.

its ability to degrade this mutant (Figure 5C). Even more marked defects in SOX catalytic efficiency were observed for the binding site mutant LIMD1-54 3XA-G, the loop mutants LIMD1-54 zipper and LIMD1-54 zipper 2, and the pre-miRNA K2-31 (Figure 5C and Supplementary Figure S7). Collectively, these data indicate that efficient RNA cleavage requires both an appropriate SOX binding site and a suitable cut site.

Site-specific endonucleolytic cleavage of target RNA occurs in vitro

In cells, SOX cleaves its mRNA substrates site-specifically. Mutagenesis of residues in mapped cleavage sites generally abolishes SOX cleavage at that location (20). To determine if our in vitro assay faithfully recapitulated the site specificity of SOX endonucleolytic targeting observed in cells, we established reaction conditions that enabled trapping of
the early cleavage events. By combining Ca\textsuperscript{2+} and Mg\textsuperscript{2+} in our reaction buffer, we were able to sufficiently slow SOX processing to visualize cleavage products derived from 5'-32P labeled substrates. Indeed, we observed a predominant 27 nt band, which is the size of the product released upon \textit{LIMD1}-54 cleavage at the predicted cut site (Figure 6A, lane 3). Additional bands also appeared, likely representing subsequent processing events. Importantly, when we incubate SOX with the cut site mutant \textit{LIMD1}-54 A-G, there is a complete loss of this 27 nt product, as well as the additionally processed intermediates (Figure 6A, lane 4). Production of these cleavage intermediates required SOX, as no decay was observed in the RNA-only controls (Figure 6A, lanes 1–2).

Finally, we sought to verify that the predominant 27 nt cleavage product we observed was a result of an endonucleolytic cleavage and not 5' end processing. To this end, we generated a \textit{LIMD1}-54 substrate containing a 3'-32P pCp label and a free 5' OH to block 5' end processing. Again, in the presence of SOX, WT \textit{LIMD1}-54 but not the A-G mutant produced a cleavage product whose size corresponded to cleavage at the predicted site (Figure 6B). Taken together, these data confirm that our \textit{in vitro} assay faithfully recapitulates SOX cleavage site specificity on a true substrate.

**DISCUSSION**

Endonuclease-directed mRNA degradation plays key roles in the lifecycle of gammaherpesviruses, yet the fundamental principles governing target specificity by SOX and other viral endonucleases are not well understood. Here, through the development of the first biochemical system to faithfully recapitulate the internal cleavage specificity observed for SOX in cells, we revealed how both RNA sequence and structure contribute to targeting. These findings resolve a central feature of the current model of SOX activity (Figure 7). Previous observations established that sequences flanking the cut site were required to direct cleavage by SOX (16,20). However, it was unresolved whether they played a strictly structural role in presenting an exposed loop for cleavage, served as a platform for SOX binding, or created a binding site for one or more cellular factors that then indirectly recruited SOX to its targets. Through a combination of mutational analyses, RNA structure probing, and RNA footprinting assays, we showed that efficient SOX targeting requires both an exposed loop structure and upstream sequences that serve as a SOX binding platform. This combination of sequence and structural features within the targeting motif helps explain why some mRNAs are efficiently cleaved by SOX, whereas others are weaker substrates.

A key open question related to SOX function is how it can target the majority of mRNAs in cells, yet with significant site specificity. Our observations suggest that there must be specific mRNA features that influence targeting. Indeed, PARE-seq analyses of cleavage intermediates in SOX expressing cells revealed that cleavage sites were associated with a degenerate sequence motif (20). Sequences proximal to the cleavage site were predicted to be un-base paired and frequently contained a polyadenosine stretch followed by a purine (20). The requirement for these sequence features for SOX targeting was validated for the \textit{LIMD1} transcript in cells (20). Because \textit{LIMD1} has been established as a particularly robust SOX target in cells (20), we reasoned that it must contain features optimal for SOX processing and therefore would be an ideal substrate to dissect biochemically why these features are important. Indeed, SOX binding to \textit{LIMD1}-54 was 10-fold better than to the commonly used reporter substrate GFP, and ~100-fold better than to the \textit{K2-31} pre-miRNA, which has not been demonstrated to be processed by SOX in cells. Importantly, these binding differences correlated with the efficiency of SOX cleavage in \textit{in vitro}, arguing that the ability to bind the targeting motif is a key step in target recognition. Through RNA footprinting assays, we were able to show that SOX binds to a bulge structure proximal to the cleavage site containing the polyadenosine stretch previously predicted to be important for mRNA cleavage by SOX in cells (20). Mutating either just the bulge structure (\textit{LIMD1}-54 zipper 2) or maintaining the bulge but mutating the polyadenosine stretch (\textit{LIMD1}-54 3xA-G) resulted in a ~100-fold reduction in binding affinity, correlating with a dramatic decrease in cleavage efficiency. Collectively, these data demonstrate that variability in the efficiency of SOX targeting observed in cells is likely due to differences in RNA sequences that mediate SOX binding.

A recent crystal structure of SOX bound to the \textit{K2-31} pre-miRNA captured the importance of the exposed loop region for SOX cleavage (21). However, the structure did not reveal additional interactions between SOX and the RNA beyond the three residues surrounding the cut site. Our data suggest that this is likely because the \textit{K2-31} RNA lacks the additional residues necessary for SOX binding site found in both \textit{LIMD1} and \textit{GFP}. While the \textit{K2-31} RNA does contain adenosines upstream of the cleavage site, structural predictions indicate these residues are within a stem region (21), rather than in an exposed loop as is the case for \textit{LIMD1} and \textit{GFP}. Together, these observations indicate that while upstream adenosines are important for binding, they must be present in an unpaired state to promote SOX binding.

It is notable that prior studies reported much weaker interactions between SOX and RNA (K\textsubscript{d} = 75 \textmu M) compared to its DNA substrates (K\textsubscript{d} = 1 \textmu M) (9,10,21). However, in these cases binding assays were conducted with scrambled RNA sequences. We found that SOX binding affinities to RNA substrates vary over several orders of magnitude, in a manner that correlates with cleavage efficiency. Interestingly, the crystal structure of SOX bound to DNA showed more dynamic interactions along the length of the protein (~480 \AA interaction surface), when compared to the \textit{K2-31} RNA bound structure (~240 \AA interaction surface). It is therefore possible that more interaction along the length of SOX protein might occur with optimal substrates such as \textit{LIMD1} that are more tightly bound.

The fact that purified SOX endonucleolytically cleaved \textit{LIMD1}-54 at the precise site observed in SOX-expressing cells demonstrates that cleavage site selection on an mRNA is not mediated by a cellular cofactor. Instead, targeting at particular RNA motifs is strongly influenced by the strength of SOX binding. Our observation that the P176S and F179A SOX mutants display significant RNA binding defects indicates that their failure to cleave mRNAs in cells is due to an inability to efficiently bind the targeting motif.
Figure 6. SOX endonucleolytically cleaves LIMD1-54 in a site-specific manner. (A) 5′ 32P-labeled LIMD1-54 and LIMD1 A-G were incubated for 10 min in 0.1 mM CaCl2 and 0.7 mM MgCl2 in the presence or absence of 2 µM SOX. The 5′ cleavage product with a size corresponding to cleavage at the mapped cut site (27 nt) is shown. (B) Cleavage assay was performed as in (A), except using 3′ 32P pCp-labeled LIMD1-54. The 3′ cleavage product with a size corresponding to cleavage at the mapped cut site (28 nt) is shown.

Figure 7. Model of mRNA targeting by SOX. SOX is able to distinguish mRNA from other types of RNA in cells by an as yet unknown mechanism. Subsequently, it endonucleolytically cleaves its targets at specific sites, whereupon the fragments are degraded by host exonucleases such as XRN1 and DIS3L2. Here, we revealed that in addition to the requirement for an unpaired loop at the cleavage site, additional upstream RNA sequences increase the affinity of SOX for individual targets, thereby controlling cleavage efficiency.

The mechanism by which SOX initially distinguishes RNA polymerase II transcribed mRNAs from other types of RNA in cells remains an important open question, as this feature of SOX selectivity is not preserved in vitro. We hypothesize that cellular co-factors, perhaps though interactions with SOX, enable this distinction. More broadly, endonucleases are instrumental in RNA processing and degradation. Nuclease processing defects lead to several human pathologies ranging from cancer to neurodegeneration (30–34), and our study provides a framework for better understanding the mechanistic features governing endonuclease targeting.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.
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