RNA Recognition by the DNA End-Binding Ku Heterodimer

Andrew B. Dalby  
*University of Colorado Boulder*

Karen J. Goodrich  
*University of Colorado Boulder*

Jennifer S. Pfingsten  
*University of Colorado Boulder*

Thomas R. Cech  
*University of Colorado Boulder, thomas.cech@colorado.edu*

Follow this and additional works at: [https://scholar.colorado.edu/chem_facpapers](https://scholar.colorado.edu/chem_facpapers)  
Part of the [Biochemistry, Biophysics, and Structural Biology Commons](https://scholar.colorado.edu/chem_facpapers)

**Recommended Citation**  
[https://scholar.colorado.edu/chem_facpapers/28](https://scholar.colorado.edu/chem_facpapers/28)

This Article is brought to you for free and open access by Chemistry & Biochemistry at CU Scholar. It has been accepted for inclusion in Chemistry & Biochemistry Faculty Contributions by an authorized administrator of CU Scholar. For more information, please contact cuscholaradmin@colorado.edu.
RNA recognition by the DNA end-binding Ku heterodimer

Andrew B. Dalby, Karen J. Goodrich, Jennifer S. Pfingsten, et al.

RNA 2013 19: 841-851 originally published online April 22, 2013
Access the most recent version at doi:10.1261/rna.038703.113
RNA recognition by the DNA end-binding Ku heterodimer

ANDREW B. DALBY, KAREN J. GOODRICH, JENNIFER S. PFINGSTEN,† and THOMAS R. CECH‡
Howard Hughes Medical Institute, Department of Chemistry and Biochemistry, University of Colorado BioFrontiers Institute, Boulder, Colorado 80309-0596, USA

ABSTRACT
Most nucleic acid-binding proteins selectively bind either DNA or RNA, but not both nucleic acids. The Saccharomyces cerevisiae Ku heterodimer is unusual in that it has two very different biologically relevant binding modes: (1) Ku is a sequence-nonspecific double-stranded DNA end-binding protein with prominent roles in nonhomologous end-joining and telomeric capping, and (2) Ku associates with a specific stem–loop of TLC1, the RNA subunit of budding yeast telomerase, and is necessary for proper nuclear localization of this ribonucleoprotein enzyme. TLC1 RNA-binding and dsDNA-binding are mutually exclusive, so they may be mediated by the same site on Ku. Although dsDNA binding by Ku is well studied, much less is known about what features of an RNA hairpin enable specific recognition by Ku. To address this question, we localized the Ku-binding site of the TLC1 hairpin with single-nucleotide resolution using phosphorothioate footprinting, used chemical modification to identify an unpredicted motif within the hairpin secondary structure, and carried out mutagenesis of the stem–loop to ascertain the critical elements within the RNA that permit Ku binding. Finally, we provide evidence that the Ku-binding site is present in additional budding yeast telomerase RNAs and discuss the possibility that RNA binding is a conserved function of the Ku heterodimer.

Keywords: Ku heterodimer; RNA binding; telomerase; footprinting; chemical modification

INTRODUCTION
Telomerase is the enzyme that synthesizes telomeric DNA repeats to cap chromosome ends in eukaryotic cells. Telomerase is a ribonucleoprotein complex, and the conserved core enzyme is comprised of a reverse transcriptase (TERT) and an RNA component (TER) (Greider and Blackburn 1987; Lingner et al. 1997). TER contains an internal template, which is used by TERT to catalyze the addition of telomeric DNA repeats to chromosome termini during the late stages of semiconservative DNA replication. The core enzyme is sufficient for catalysis in vitro. However, additional protein cofactors are necessary in vivo to mediate essential functions such as nuclear localization, recruitment of telomerase to the telomere, and stimulation of telomerase repeat addition processivity (Egan and Collins 2012; Nandakumar and Cech 2013).

The telomerase core in the budding yeast Saccharomyces cerevisiae consists of the large 1.1-kb TLC1 RNA and Est2 (Ever shorter telomeres 2), the yeast TERT (Singer and Gottschling 1994; Lingner et al. 1997). TLC1 is present at ~30 copies per haploid cell, which is less than the number of telomeric ends (Mozdy and Cech 2006). Consistent with limiting telomerase, only a subset of telomeres are extended by a few repeats each cell cycle, with preferential extension at telomeres with shorter tracts (Teixeira et al. 2004). Structurally, TLC1 contains three long helical arms that emanate from a conserved pseudoknot (Dandjinou et al. 2004; Lin et al. 2004; Zappulla and Cech 2004). Est2 binds to the central pseudoknot and template region, and each of the arms serves as a scaffold for one of the protein cofactors that comprise the holoenzyme (Zappulla and Cech 2004). The terminal arm contains the binding site for the Sm7 complex, another arm contains the Est1-binding site, and the arm extending from the template boundary helix contains the Ku-binding site (KBS) (Seto et al. 1999, 2002; Peterson et al. 2001).

The Ku heterodimer is comprised of two subunits that share the same overall topology, namely, an N-terminal α/β domain followed by a β-barrel domain and a variable C-terminal arm (Walker et al. 2001). The structural basis for the recognition of duplex DNA by Ku is illustrated in the cocrystal structure of human Ku bound to DNA (Walker et al. 2001). The two β-barrel domains of each subunit associate at a large dimerization interface. A preformed ring structure is formed by two loops protruding from each subunit to bridge the interface. DNA binds in the central cavity of the ring and makes sequence-independent electrostatic contacts with the surfaces of both subunits adjacent to the dimerization interface. In yeast, both Ku subunits are ~70 kDa and show sequence similarity with the subunits of the human Ku heterodimer (Feldmann et al. 1996). The Ku70 subunit, which faces toward the DNA terminus, is crucial for NHEJ function, and the Ku80 subunit, which faces away from the terminus, maintains telomeric silencing functions (Bertuch and Lundblad 2003;...
In yeast, knockout of the Ku heterodimer contributes to telomere shortening and elongated 3’ overhangs as a result of increased Exo1 access to the C-strand (Boulton and Jackson 1999; Gravel et al. 1998; Bertuch and Lundblad 2004; Vodenicharov et al. 2010). Ku’s well-established role as a multifaceted DNA end-binding protein raises questions about how and why it binds to the TLC1 RNA.

The association of Ku with TLC1 was originally identified in an overexpression screen for disruption of telomeric silencing, and deletion analysis localized the KBS to a 48-nt stem-loop at the distal end of the template boundary arm (Peterson et al. 2001). The terminal KBS (nt 288–312) is predicted to form a hairpin with a two-base bulge at the center. The fold of the terminal KBS is retained in subsequent structural modeling of TLC1, but the conformation of the helix adjacent to the KBS varies among the proposed structures (Dandjinou et al. 2004; Zappulla and Cech 2004). Disruption of the KBS structure perturbs Ku binding both in vivo and in vitro, resulting in a telomere shortening phenotype (Peterson et al. 2001; Stellwagen et al. 2003).

Ku has been proposed to serve as a bridging factor to recruit telomerase to the telomere during G1 of the cell cycle, and thereby help to facilitate telomerase action during late S-phase (Peterson et al. 2001; Stellwagen et al. 2003; Fisher et al. 2004). Ku-dependent recruitment is thought to act in concert with the established Est1-Cdc13 recruitment mechanism (Evans and Lundblad 1999; Fisher et al. 2004). Loss of Ku or TLC1 results in a substantial decrease in de novo telomere addition upon DNA damage, adding support to the Ku-dependent telomerase recruitment hypothesis (Myung et al. 2001; Stellwagen et al. 2003). Notably, deletion of the KBS within TLC1 does not affect telomere capping or telomeric silencing (Peterson et al. 2001; Vodenicharov et al. 2010).

More recent experiments indicate that the Ku–TLC1 interaction may play a more complex role in telomerase recruitment than originally thought. Fluorescent in situ hybridization experiments demonstrate that the deletion of the KBS or either of the Ku subunits results in decreased TLC1 nuclear localization, explaining in part the telomere shortening phenotype (Gallardo et al. 2008; Pfingsten et al. 2012). Live cell imaging experiments indicate that telomerase only transiently associates with telomeres during G1 (Gallardo et al. 2011). Additionally, Ku binds the KBS RNA and duplex DNA in a mutually exclusive and competitive manner in vitro (Pfingsten et al. 2012). These experiments call into question the hypothesis that Ku serves as a simple bridging factor to recruit telomerase. Nonetheless, the insertion of an additional KBS into the TLC1 RNA results in a telomere hyperelongation phenotype, indicating that the Ku–TLC1 interaction may contribute to telomere homeostasis in capacities beyond nuclear localization (Zappulla et al. 2011). For example, the Ku–TLC1 interaction is also thought to contribute to anchoring of the telomerase to the nuclear envelope (Schober et al. 2009).

In spite of the clear Ku–TLC1 association in S. cerevisiae, it is unclear whether Ku’s role as a protein cofactor for telomerase is conserved among other eukaryotes. Deletion of YKU70 in a S. paradoxus strain with long telomeres results in a telomere shortening phenotype, providing indirect evidence for a TLC1–Ku interaction (Liti et al. 2009b). KBSs are proposed to exist in both sensu stricto and sensu lato Saccharomyces, and a putative KBS that varies in both hairpin size and sequence appears to be present in the Candida glabrata TER (Chappell and Lundblad 2004; Kachouri-Lafond et al. 2009). Knockout of YKUBO in Kluyveromyces lactis does not result in a telomere-shortening phenotype, and structural modeling suggests that K. lactis TER does not contain a KBS (Brown et al. 2007; Carter et al. 2007; Kabaha et al. 2008). Outside of the Saccharomyces clade, KBSs have not been detected in the TERs of the fission yeast Schizosaccharomyces pombe or the filamentous ascomycete Neurospora crassa (Webb and Zakian 2008; Qi et al. 2012). Although the precise Ku-binding sites have not been identified, Ku is reported to bind to both human TER and specific TER isoforms in Arabidopsis thaliana (Ting et al. 2005, 2009; Cifuentes-Rojas et al. 2012) (see also Chai et al. 2002).

Ku also appears to bind to a host of other cellular RNAs. Prior to DNA damage, human Ku resides in the nucleolus and associates with a host of RNA-binding proteins in an RNA-dependent manner; following DNA damage, Ku disperses throughout the nucleus and incorporates into DNA damage complexes (Adelmant et al. 2012). Immunoprecipitation experiments demonstrate that Ku interacts with a pool of RNAs in HeLa nuclear extracts (Zhang et al. 2004). Ku is reported to bind the HIV TAR element stem–loop with high affinity in vitro (Kaczmarski and Khan 1993). Thus, mounting evidence suggests that RNA binding may be a conserved function of the Ku heterodimer.

In order to better understand the RNA-binding function of Ku, we set out to characterize the interaction between Ku and the TLC1 KBS. We first identify the site of Ku binding on the TLC1 KBS with nucleotide resolution. We next provide evidence that the KBS secondary structure folds into an unusual bulged hairpin. Through mutagenesis of the KBS, we identify the critical determinants for Ku binding and define a minimal KBS motif. Finally, we provide evidence that KBSs are likely retained in several Saccharomyces TERs.

RESULTS

Ku interacts with the terminal stem–loop of the template boundary arm

To identify the site of Ku binding on the TLC1 RNA with nucleotide resolution, we carried out phosphorothioate footprinting experiments. Protein contacts on an RNA molecule can be identified by protection of phosphorothioate linkages from chemical cleavage by iodine (Schatz et al. 1991; Rudinger et al. 1992). In vitro-transcribed RNAs, with
randomly incorporated phosphorothioate nucleotide analogs, were subjected to iodine-mediated cleavage in the presence or absence of the purified Ku heterodimer (Fig. 1A). The cleavage of a number of phosphorothioate linkages was reduced in the presence of Ku (Fig. 1B). The most stable protections were clustered in the terminal loop of the KBS RNA at nt C300, A302, A303, and U306. Protections in the terminal loop had roughly twofold reductions in cleavage throughout the duration of the time course (Fig. 1C). In contrast, less stable protections extended from the loop into the adjacent stems surrounding the AU bulge. For example, nt U290, A294, G295, and G311 all had greater than twofold reductions in cleavage in a number of phosphorothioate linkages was reduced in the presence of Ku (Fig. 1B). The most stable protections were clustered in the terminal loop of the KBS RNA at nt C300, A302, A303, and U306. Protections in the terminal loop had roughly twofold reductions in cleavage throughout the duration of the time course (Fig. 1C). In contrast, less stable protections extended from the loop into the adjacent stems surrounding the AU bulge. For example, nt U290, A294, G295, and G311 all had greater than twofold reductions in cleavage in a number of phosphorothioate linkages was reduced in the presence of Ku (Fig. 1B). The most stable protections were clustered in the terminal loop of the KBS RNA at nt C300, A302, A303, and U306. Protections in the terminal loop had roughly twofold reductions in cleavage throughout the duration of the time course (Fig. 1C). In contrast, less stable protections extended from the loop into the adjacent stems surrounding the AU bulge. For example, nt U290, A294, G295, and G311 all had greater than twofold reductions in cleavage in a number of phosphorothioate linkages was reduced in the presence of Ku (Fig. 1B). The most stable protections were clustered in the terminal loop of the KBS RNA at nt C300, A302, A303, and U306. Protections in the terminal loop had roughly twofold reductions in cleavage throughout the duration of the time course (Fig. 1C). In contrast, less stable protections extended from the loop into the adjacent stems surrounding the AU bulge. For example, nt U290, A294, G295, and G311 all had greater than twofold reductions in cleavage at 10 sec, but not 10 min (Fig. 1C).

To further test the interaction of Ku with the terminal stem-loop of the KBS, nuclease V1 footprinting was carried out on the Ku–KBS complex (Supplemental Fig. 1A). Nuclease V1 cleaves duplexed or structured RNAs in a sequence-nonspecific manner to release products with a 3′ hydroxyl and a 5′ phosphate (Ehresmann et al. 1987). The KBS was most strongly protected by Ku from V1 cleavage from nt 284–310 (Supplemental Fig. 1B). We note that V1 nuclease gives a bigger and less-resolved footprint than iodine, as expected, because an enzyme is large and much more subject to steric hindrance than a small-molecule probe. Given the overlap between the iodine and V1 nuclease footprinting, it is likely that the protections observed here are due to direct protein contacts. Nevertheless, it must be kept in mind that changes in iodine reactivity may be due to dynamics in the RNA–protein contacts or protein-dependent changes in the structure of the RNA (Rose and Weeks 2001; Webb et al. 2001). Our results agree with previous in vivo mutagenesis studies that identified the AU bulge and surrounding base pairs as important for the Ku–TLC1 interaction (Peterson et al. 2001) and show that the interaction extends into the terminal loop.

The TLC1 KBS forms a hairpin with a bulge motif

In order to experimentally test the previously predicted terminal KBS secondary structure (Peterson et al. 2001; Dandjinou et al. 2004; Zappulla and Cech 2004), chemical modification experiments were carried out to identify unpaired nucleotides. TLC1 KBS RNA was treated with either dimethyl sulfate (DMS) or 1-cyclohexyl-3-(2-morpholinoethyl)carbodiimide metho-p-toluenesulfonate (CMCT). Accessible imino groups at position 1 on adenosine and position 3 on cytosine can be methylated by DMS, and accessible imino groups at position 3 of uracil and position 1 of guanosine can be modified by CMCT (Ehresmann et al. 1987). Modifications can be detected as stops in a reverse-transcription reaction 1 nt before the site of modification (Inoue and Cech 1985).

Treatment of the KBS RNA with DMS resulted in a number of strong stops in the reverse-transcription reaction. Modification occurred at adenosines 302–304 in the terminal loop of the KBS RNA (Fig. 2A,C). Likewise, treatment of the KBS with CMCT modified U301 and U293 (Fig. 2B,D). Nucleotides in the putative stem immediately adjacent to the terminal loop were not heavily modified upon treatment with CMCT or DMS, given that the band intensities at these positions were within one standard deviation of the mean band intensity. The modification pattern of bases in the penultimate stem (nt 288–291 and 309–312) showed more variability than the terminal stem, suggesting a more complex motif. Additionally, stops corresponding to weak modification (less than one standard deviation above the mean band intensity) of U301 and U287 were observed in the DMS data. DMS is not known to react with uracil, and U287 is not modified by CMCT, so the meaning of these data is unknown. Collectively, the strong modification of the terminal

FIGURE 1. Phosphorothioate footprinting of the Ku–KBS RNA interaction. (A) Coomassie-stained SDS-PAGE analysis of the Ku heterodimer purified from S. cerevisiae. (B) Ku-dependent protection from iodine-mediated cleavage of a TLC1 KBS RNA construct transcribed in the presence of A/U or C/G phosphorothioate NTPs. Cleavage times are denoted above the gel and lasted for 10 sec, 34 sec, and 10 min. Seq indicates the sequencing lanes, and −I2 is a no-iodine control. Protected nucleotides are denoted by arrows. (C) Heatmaps of the fold-change in band intensity on a log2 scale are plotted on the secondary structure of the TLC1 KBS. KBS numbering corresponds to the numbering for full-length TLC1 RNA.
loop and low reactivity of the majority of nucleotides in the putative stems are consistent with the hypothesis that nt 288–312 form a stem-loop.

As mentioned above, the modification pattern for nt 291–293 did not appear to be consistent with a simple two-base bulge structure. U293 showed very high reactivity toward CMCT (more than three standard deviations above mean band intensity), while A292 showed virtually no reactivity toward DMS (below the mean band intensity). U291 had a moderate band intensity, greater than one standard deviation above the mean, while U290 and U308 were weakly reactive with band intensities that were above average (Fig. 2C,D). The modification data suggest that this portion of the KBS may fold into a specific structural motif. Varying chemical reactivity has been observed in the 2-nt bulge of domain V in the group II intron (Costa et al. 1998), which can likely be explained by the formation of a base-triple including one of the nucleotides in the bulge (Keating et al. 2010). Factors such as ion-binding sites or water-mediated hydrogen bonds may contribute to the protection of single-stranded nucleotides as well (Leontis and Westhof 1998). These modification data are consistent with previous secondary structure predictions in which the TLC1 KBS forms a hairpin; additionally, our data suggest that the hairpin contains a noncanonical bulge motif.

Ku from *S. cerevisiae* binds to the KBSs of noncognate TERs

An alignment of TLC1 sequences from 34 *S. cerevisiae* strains and 25 *S. paradoxus* strains, in addition to single TLC1 sequences from *S. kudriavzevi*, *S. cariocanus*, *S. bayanus*, *S. pastorianus*, *S. mikatae*, *S. arboricola*, and *S. castellii* demonstrated that the KBS is highly conserved from nt 289–311 (*S. cerevisiae* numbering, Supplemental Fig. 2A,B). Variation between the *Saccharomyces* spp. occurred in the predicted terminal loop (e.g., larger loops for *S. bayanus* and *S. castellii*). The length of the stem preceding the AU bulge varies between 3 and 4 bp (e.g., *S. castellii* compared with *S. cerevisiae*), and one A-to-G transition can be observed in the presumptive AU bulge (cf. *S. cerevisiae* and *S. arboricola*). Overall, the high degree of sequence conservation in the KBS relative to the weak conservation in other portions of this arm of TLC1 (Zappulla and Cech 2004) suggests that the RNA sequence is critical for structure and for recognition by Ku.

To verify the importance of conserved sequence in the KBS, affinity measurements were made using Ku from *S. cerevisiae* and the putative noncognate KBSs from selected sensu stricto strains, a sensu lato strain *S. castellii*, and the ascomycete *K. lactis* (Fig. 3A,B). Phylogenomic analyses indicate that *S. paradoxus* and *S. cerevisiae* are very closely related organisms (Fitzpatrick et al. 2006); correspondingly, the KBSs from these species show 100% identity from nt 288–312. Consistent with the importance of the conservation of the terminal KBS, Ku from *S. cerevisiae* bound the noncognate *S. paradoxus* KBS with an apparent *K*ₐ similar to that of the cognate interaction (Fig. 3B). Conversely, *K. lactis* is an outgroup to both the sensu stricto and sensu lato yeasts and is not predicted to have a KBS (Fitzpatrick et al. 2006; Brown et al. 2007). As expected, Ku from *S. cerevisiae* had a severely diminished affinity for the distal stem–loop of the template boundary arm from *K. lactis*.

Noncognate KBSs with intermediate sequence conservation revealed additional determinants of Ku recognition. Apparent *K*ₐ’s from *S. cerevisiae*, *S. bayanus*, and *S. castellii* KBSs revealed that the size of the terminal loop, but not the sequence, is important for recognition. In *S. castellii*, the terminal loop of the KBS increased from 7 to 11 nt and changed in sequence, but had only a slight decrease in affinity for the *S. cerevisiae* Ku (Fig. 3B). In contrast, the *S. bayanus* KBS...
Terminal stem length and bulge motif are critical for Ku recognition

Binding assays with mutant *S. cerevisiae* KBS constructs were carried out to further elucidate the elements critical for Ku binding (Fig. 4A; Table 1). The complete set of mutant KBS constructs is shown in Supplemental Figure 4. Due to the high number of uridines in the conserved region of the KBS, we reasoned that mutants would have a high propensity to improperly fold and thereby artificially increase the $K_d$. To try to identify potentially misfolded mutants, and hence exclude them from our study, RNA secondary structures were analyzed by mFold. Then to experimentally confirm the mFold predictions and verify that sequence changes did not affect overall structure, the mFold predictions and verify that sequence changes did
not cause RNAs to adopt unanticipated conformations, we carried out V1 nuclease digests of selected mutants (Supplemental Fig. 3A). We found that when the lowest free energy structure for a mutant was misfolded (i.e., did not maintain the mFold-predicted 4 bp stems on either side of the bulge), the mutant RNA typically misfolded as judged by V1 nuclease digests (Supplemental Fig. 3B,C). Although our chemical modification data suggest that the native KBS may contain a more complex bulge motif than is represented by the predicted mFold secondary structure, we found mFold a useful tool for eliminating mutants prone to misfolding.

Previous models of the *S. cerevisiae* KBS secondary structure differ slightly in the conformation of the helices formed by nt 276–287 and 313–323 (Peterson et al. 2001; Dandjinou et al. 2004; Zappulla and Cech 2004). We tested two KBS constructs predicted to fold into these secondary structures (Supplemental Fig. 4; cf. Zappulla: TLC1-96 and Dandjinou: TLC1-56). We found that Ku bound both constructs with the same affinity (Fig. 3; Table 1). The construct with the predicted Zappulla fold (TLC1-96) bound Ku with a $K_d$ of 1.3 ± 0.5 nM, and the construct with the predicted Dandjinou (TLC1-56) fold bound with a $K_d$ of 1.13 ± 0.02 nM. In addition, both the sequence and predicted folds for the noncognate *S. paradoxus* and *S. castellii* KBSs differed from that of *S. cerevisiae* KBS, yet both noncognate KBSs bound *S. cerevisiae* Ku with only slightly diminished affinities. In sum, these data

### TABLE 1. Binding data for mutant KBS RNAs

<table>
<thead>
<tr>
<th>Mutant number</th>
<th>Region of hairpin mutated</th>
<th>Original base pair or sequence</th>
<th>Mutated base pair or sequence</th>
<th>Nucleotides changed</th>
<th>$K_d$ nM</th>
<th>$\Delta \Delta G^\circ$ kcal/mol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>1.3 ± 0.5</td>
<td>–</td>
</tr>
<tr>
<td>1</td>
<td>Reverse complement terminal loop</td>
<td>GGCUAA</td>
<td>CCGAUUU</td>
<td>298–304</td>
<td>3.0 ± 0.9</td>
<td>0.48</td>
</tr>
<tr>
<td>2</td>
<td>Reverse complement portion of terminal loop</td>
<td>CUAA</td>
<td>GAUUU</td>
<td>300–304</td>
<td>2.4 ± 0.5</td>
<td>0.35</td>
</tr>
<tr>
<td>3</td>
<td>Compensatory double mutant</td>
<td>A/U, U/A</td>
<td>C/G, C/G</td>
<td>296–297, 305–306</td>
<td>3.9 ± 0.5</td>
<td>0.62</td>
</tr>
<tr>
<td>4</td>
<td>Combined changes from mutants 2 and 3</td>
<td>A/U, U/A, CUAA</td>
<td>C/G, C/G, GAUUU</td>
<td>296–306</td>
<td>5 ± 1</td>
<td>0.76</td>
</tr>
<tr>
<td>5</td>
<td>AU bulge deletion</td>
<td>AU</td>
<td>No sequence change</td>
<td>292–293</td>
<td>&gt;270</td>
<td>&gt;2.91</td>
</tr>
<tr>
<td>6</td>
<td>Internal loop deletion</td>
<td>CU, UUA</td>
<td>No sequence change</td>
<td>286–287, 312–314</td>
<td>2.5 ± 0.4</td>
<td>0.36</td>
</tr>
<tr>
<td>7</td>
<td>GUG bulge deletion</td>
<td>GUG</td>
<td>No sequence change</td>
<td>280–282</td>
<td>1.5 ± 0.4</td>
<td>0.08</td>
</tr>
<tr>
<td>8</td>
<td>AU bulge sequence inversion</td>
<td>AU</td>
<td>UA</td>
<td>292–293</td>
<td>7 ± 2</td>
<td>0.96</td>
</tr>
<tr>
<td>12</td>
<td>Compensatory mutant</td>
<td>U/A</td>
<td>G/C</td>
<td>290, 310</td>
<td>29 ± 4</td>
<td>1.70</td>
</tr>
<tr>
<td>16</td>
<td>Terminal loop to GNRA tetraloop</td>
<td>GCUAA</td>
<td>GAAA</td>
<td>299–301</td>
<td>8 ± 1</td>
<td>1.05</td>
</tr>
<tr>
<td>17</td>
<td>Wobble pair to Watson-Crick pair</td>
<td>U/G</td>
<td>U/A</td>
<td>309</td>
<td>17 ± 4</td>
<td>1.40</td>
</tr>
<tr>
<td>18</td>
<td>Compensatory mutant</td>
<td>U/G</td>
<td>G/U</td>
<td>291, 309</td>
<td>35 ± 8</td>
<td>1.80</td>
</tr>
<tr>
<td>19</td>
<td>Watson-Crick pair to wobble pair</td>
<td>A/U</td>
<td>G/U</td>
<td>294</td>
<td>61 ± 12</td>
<td>2.11</td>
</tr>
<tr>
<td>20</td>
<td>Compensatory mutant</td>
<td>A/U</td>
<td>G/U</td>
<td>294, 308</td>
<td>6 ± 2</td>
<td>0.87</td>
</tr>
<tr>
<td>21</td>
<td>Compensatory mutant</td>
<td>A/U</td>
<td>G/C</td>
<td>295, 307</td>
<td>1.5 ± 0.4</td>
<td>0.09</td>
</tr>
<tr>
<td>22</td>
<td>Compensatory mutant</td>
<td>A/U</td>
<td>G/C</td>
<td>288, 312</td>
<td>1.6 ± 0.3</td>
<td>0.13</td>
</tr>
<tr>
<td>23</td>
<td>Lengthening of terminal stem</td>
<td>AGAU:AUCU</td>
<td>AGAGAU: AUCUCU</td>
<td>Ins 293–294, 308–309</td>
<td>164 ± 17</td>
<td>2.64</td>
</tr>
<tr>
<td>24</td>
<td>Lengthening of terminal stem</td>
<td>AGAU:AUCU</td>
<td>AGAGAU: AUCUCU</td>
<td>Ins 293–294, 308–309</td>
<td>&gt;270</td>
<td>&gt;2.91</td>
</tr>
<tr>
<td>25</td>
<td>Compensatory mutant</td>
<td>U/A</td>
<td>C/G</td>
<td>290, 310</td>
<td>17.1 ± 0.6</td>
<td>1.41</td>
</tr>
<tr>
<td>26</td>
<td>Compensatory mutant</td>
<td>G/C</td>
<td>C/G</td>
<td>295, 307</td>
<td>1.6 ± 0.4</td>
<td>0.12</td>
</tr>
<tr>
<td>27</td>
<td>Compensatory mutant</td>
<td>A/U</td>
<td>U/A</td>
<td>288, 312</td>
<td>1.5 ± 0.5</td>
<td>0.13</td>
</tr>
<tr>
<td>28</td>
<td>AU bulge sequence change</td>
<td>AU</td>
<td>G</td>
<td>292–293</td>
<td>&gt;81</td>
<td>&gt;2.26</td>
</tr>
<tr>
<td>29</td>
<td>AU bulge sequence change</td>
<td>AU</td>
<td>AA</td>
<td>293</td>
<td>1.2 ± 0.3</td>
<td>−0.04</td>
</tr>
<tr>
<td>30</td>
<td>AU bulge sequence change and insertion</td>
<td>AU</td>
<td>AAA</td>
<td>Ins 292–293, 293</td>
<td>17 ± 2</td>
<td>1.40</td>
</tr>
<tr>
<td>31</td>
<td>AU bulge sequence change and insertion</td>
<td>AU</td>
<td>AAA</td>
<td>Ins 292–293, 293</td>
<td>17 ± 6</td>
<td>1.43</td>
</tr>
</tbody>
</table>

*Compensatory mutations correspond to changes between Watson–Crick base pairs in the paired regions of the stem.*

*For mutants 5–7, nucleotides were deleted; however, the remaining sequence was not changed.*

*Ins designates an insertion between the bases listed.*

*$K_d$ values reported represent an average of three determinations normalized to the wild-type $K_d$ from that set of experiments.

*$\Delta \Delta G^\circ = \Delta G^\circ(\text{mutant}) - \Delta G^\circ(\text{wild-type}) = RT \ln(\text{mutant} / \text{wild-type} K_d)$, calculated from the mean $K_d$ values, measurements made under the conditions described in Materials and Methods. Mutants 9–11 and 13–15 were suspected of misfolding, and therefore not included in the analysis for this study.
support the conclusion that the specific sequence or structure of the template boundary arm preceding the terminal hairpin is not a factor in Ku recognition, but that the terminal conserved region of the KBS (nt 289–311) is the critical Ku recognition element.

Considering the terminal hairpin loop, KBSs with sequence changes in the terminal loop bound Ku with near wild-type affinity (mutants 1 and 2) (Fig. 4A; Table 1). In contrast, mutant 16, which shortened the terminal loop to four bases, had a moderate fivefold reduction in affinity. These data were consistent with the findings of the noncognate binding studies, which indicated that the size of the terminal loop was important for high-affinity interactions with Ku. Deletions of internal loops and bulges outside of the conserved region of the KBS (nt 289–311) had minor effects on Ku recognition (mutants 6 and 7).

Our structure-probing data suggested that the conserved portion of the KBS formed a hairpin with a noncanonical bulge motif. In order to test this prediction, we made a series of mutants to produce compensatory base-pair substitutions within the helix. We reasoned that compensatory mutations in the helix should have little or no impact, provided that both the KBS structure and potential KBS–Ku contacts were preserved. Base-pair substitutions at three of the four base pairs in the terminal helix and at base-pair A288:312U in the penultimate helix had very little impact on KBS affinity for Ku (mutants 3, 21, 22, and 26) (Fig. 4A; Table 1; Supplemental Fig. 4). In contrast, mutations of base pairs surrounding the bulge at positions U290:310A, U291:309G, and A294:308U had moderate to severe reductions in affinity (∼5–60-fold, see mutants 12, 17–20, 25). To test whether the length of the helix between the AU bulge and terminal loop is important, the stem was lengthened by 2 or 4 bp while preserving the native KBS sequence immediately adjacent to the bulge. Lengthening of the stem resulted in increasingly drastic reductions in Ku affinity (mutants 23 and 24).

To determine the importance of the bulge motif for Ku association, a series of mutants was made to probe the size and sequence requirements within the bulge. Deletion of both bulge nucleotides resulted in a severe reduction in affinity (>230-fold, see mutant 5), indicating that this bulge is essential for proper Ku binding (Fig. 4A; Table 1; Supplemental Fig. 3A). Other mutations within the bulge had variable effects. For example, reduction of the AU bulge to 1 nt also resulted in a severe binding defect (mutant 28). On the other hand, mutations that altered the sequence at positions 292 or 293 resulted in much less severe binding defects. Inversion of nt 292 and 293 resulted in a sixfold reduction in affinity, while the transversion U293A had no impact on KBS–Ku binding (mutants 8 and 29, respectively). Finally, increasing the size of the bulge by the addition of adenosines resulted in moderate reductions in affinity (mutants 30 and 31), indicating that increases in bulge size were tolerated more than decreases in bulge size. We conclude that the terminal helix length and the nucleotides in and surrounding the noncanonical bulge motif (nt 290–294, 308–310) are crucial in mediating the KBS–Ku interaction.

Based on the results of the mutagenesis studies, noncognate binding assays, and footprinting experiments, we propose a minimum consensus for the KBS (Fig. 4B). Mutations resulting in greater than fivefold reductions in affinity are identified as positions in the stem–loop that are critical for Ku binding. The sequence of the two base pairs preceding the bulge, and the first base pair following the bulge, is important for recognition by Ku or the overall fold of the hairpin. The ability to substitute but not delete bulged nucleotides argues for a specific fold being important for Ku binding, but sequence-dependent contacts within the bulge cannot be entirely dismissed. The nucleotide identity at position 292 appears to be marginally important, while position 293 can be mutated without significantly impacting affinity (mutant 29). Based strictly on our mutagenesis, an adenosine appears to be important for Ku recognition; however, previous mutagenesis studies indicate that a guanosine is tolerated much better than a cytosine at this position in vivo (Peterson et al. 2001). With the exception of the compensatory mutations listed above, mutations that preserved helical structure, changed loop sequence, or disrupted internal loops had little effect on affinity and were deemed not to be critical determinants of Ku binding. In summary, we can now more precisely define the KBS for yeast Ku as a hairpin containing a noncanonical bulge motif that is capped by a 4-bp stem and terminal loop with 7–11 nt.

**DISCUSSION**

Previous studies identified a 48-nt hairpin as important for the TLC1–Ku interaction in vivo in *S. cerevisiae* (Peterson et al. 2001; Stellwagen et al. 2003). Here we extend these studies with footprinting, which indicates where Ku physically interacts with the KBS RNA, and with mutational analysis, which quantifies the contributions of individual RNA structural features to the binding energy. We identify a noncanonical bulge motif within the highly conserved portion of the KBS, and show that spacing between the bulge and terminal loop is critical for Ku recognition. Additionally, we provide direct evidence that *S. cerevisiae* Ku is able to recognize several noncognate KBSs. We suggest that the KBS is a conserved element within both sensu stricto and sensu lato *Saccharomyces* species and discuss the possibility that Ku recognizes additional cellular RNAs.

Phosphorothioate footprinting shows that the Ku interaction is restricted to 25 nt in the KBS and identifies putative contacts at nucleotide resolution. The contacts are in the terminal loop and stems on either side of the AU bulge. The footprint encompasses the portion of the KBS that is highly conserved between putative noncognate KBSs, and the noncanonical bulge motif that is particularly susceptible to mutation. The time-dependent contacts in the stem may be the result of transient KBS–protein interactions or Ku-dependent
changes in the KBS structure (Rose and Weeks 2001; Webb et al. 2001). Although we cannot definitively distinguish between these possibilities, the complex was equilibrated prior to footprinting and any Ku-induced changes in local KBS structure to promote binding likely occurred prior to iodine-mediated cleavage. We favor a model in which Ku directly interacts with the terminal loop as well as the stems on either side of the bulge, as supported by the V1 nuclease protection data (Supplemental Fig. 1).

Our footprinting data show that Ku interacts with the highly conserved portion of the KBS (nt 289–311). Covariations to support the RNA secondary structure model are not present among currently identified yeast KBSs. Therefore, we utilized chemical probing experiments, which provided support for the proposed bulged hairpin structure. In addition, the modification pattern suggested that the KBS adopts a non-Watson–Crick helical fold from positions 290 to 293. The structure of this bulge motif cannot be predicted solely from the modification pattern. Some possibilities for the protection of A292, but not U293, include the formation of a base-triple including A292, intercalation of bases into the helical stack, ion-coordination, or water-mediated hydrogen bonding (Costa et al. 1998; Leontis and Westhof 1998). Further investigation will be necessary to distinguish between these possibilities. The importance of this bulge is underscored by the detrimental effect that mutations have in this region of the RNA, as discussed below.

Our direct binding data are consistent with previous in vivo studies, which show that disruption of the proposed KBS hairpin secondary structure perturbs the Ku–TLC1 genetic interaction (Peterson et al. 2001). Mutations in and surrounding the bulge region have the greatest impact on KBS function in vivo (Peterson et al. 2001). Because mutations in the terminal loop have a small impact on KBS function (Peterson et al. 2001) and binding (mutants 1, 2, and 4, this study), contacts between Ku and the terminal loop of the KBS are likely to be sequence nonspecific and may involve the RNA backbone. Mutations in and directly adjacent to the bulge have a strong impact both in vivo (Peterson et al. 2001) and in vitro (positions 291 and 292, this study). Our data suggest that mutations to the two terminal base pairs adjacent to the loop do not significantly affect the ability of the KBS to bind Ku (mutant 3); in agreement, mutations at these positions do not seem to perturb the KBS–Ku interaction in vivo. In contrast, our data show that binding is not compromised in mutant 21 (G295A;C307U); however, there appears to be a minor impact on KBS function for this mutant in vivo (Peterson et al. 2001). Finally, we show a moderate binding defect when making compensatory mutations at the U290:310A base pair. In vivo, mutations at this position do not have a detrimental impact. The differences in the impact of mutations to the KBS may be a reflection of the different assays used to monitor the Ku–KBS interaction. The in vivo analysis measures the ability of the mutant KBS to disrupt telomeric silencing in the context of overexpression, while we measure the impact on Ku–KBS binding equilibria in vitro.

The identification of KBSs competent for specific association with Ku in TERs of several Saccharomyces species adds to the growing evidence that RNA binding is a conserved function of the Ku heterodimer. Ku associates with cellular RNAs in human cells (Zhang et al. 2004; Ting et al. 2009; Adelant et al. 2012). Ku interacts with noncoding RNAs such as TERRA and Y-RNAs (Zhang et al. 2011; Pfeiffer and Lingner 2012). RNA aptamers have been selected to bind human Ku with high affinity (Yoo and Dynan 1998). The association of human Ku with HIV-1 TAR RNA provides a specific example of another hairpin that is competent for binding Ku (Kaczmarski and Khan 1993). The TAR hairpin shows little or no sequence similarity with the TLC1 KBS; however, they share similar secondary structures and both are able to compete with double-stranded DNA for Ku binding (Kaczmarski and Khan 1993; Pfingsten et al. 2012). The TAR hairpin has a 6-nt terminal loop that is separated from a 3-nt bulge by an intervening 4-bp helix. The lack of sequence similarity between these RNAs may reflect differences between the human and S. cerevisiae Ku heterodimers; however, the conserved bulge hairpin with specific spacing requirements suggests that Ku recognizes RNAs with a conserved fold. We therefore favor a model in which Ku recognizes the hairpin structure rather than making sequence-specific contacts. One possibility is that the KBS has the ability to mimic a B-form DNA helix and thereby associate with Ku’s preformed ring; B-DNA mimicry by RNAs to associate with proteins has been previously observed (Reiter et al. 2008; Bullock et al. 2010). The identification of additional Ku-binding RNAs and an increased understanding of how Ku recognizes these hairpins will certainly shed light on how this nucleic acid-binding protein mediates a myriad of distinct cellular functions.

MATERIALS AND METHODS

KBS constructs and in vitro transcription of RNAs

TER sequences and alignments for representative Saccharomyces strains were obtained from the Telomerase Database (Podlevsky et al. 2008); TER sequences from additional Saccharomyces strains and species were manually added to the alignment using the program BioEdit V7.1.3.0. S. cerevisiae and S. paradoxus sequences were obtained by BLAST from the Saccharomyces genome resequencing project (Liti et al. 2009a). The S. arboricola putative TER sequence was recovered by BLAST from the whole-genome assembly (GCA_000292725.1) using the S. mikatae TER as a query sequence.

For a complete list of RNA constructs used see Supplemental Figure 4. RNAs were designed and checked for misfolding using mfold version 2.3; RNAs were folded at 30°C (Zuker 2003). Mutant RNA constructs were prepared by quick-change mutagenesis (Agilent) of the previously reported TLC1-96 KBS construct (Pfingsten et al. 2012). The TLC1 KBS constructs used for phosphorothioate footprinting and mutagenesis experiments were
prepared by T7 in vitro transcription from PCR products as previously described (Pfingsten et al. 2012). The TLC1 KBS construct used for the chemical modification experiments, as well as putative Ku arms from other *Saccharomyces* TERS, were made by annealing and cloning complementary oligos containing the putative Ku arm sequence, a T7 promoter, and FokI site. The RNAs were prepared by T7 polymerase run-off transcription from the FokI linearized plasmid. All in vitro-transcribed RNAs were purified on a denaturing 10% polyacrylamide gel. Prior to 32P-labeling RNAs, calf intestinal alkaline phosphatase (Roche) was used to remove the 5′ phosphate. RNAs were then 5′ labeled with [γ-32P]ATP (Perkin Elmer) and T4 polynucleotide kinase (NEB) according to the manufacturer’s protocol. 5′-labeled RNAs were then repurified on a denaturing 10% polyacrylamide gel.

**Purification of the Ku heterodimer**

Protein constructs and purification were previously described (Pfingsten et al. 2012) or had the following modifications. Cells were grown to a final density of 1–4 × 10^7 cells/mL. Lysate was passed over a 5-mL HisTrap HP column (GE Healthcare). Bound protein was eluted first in shallow gradient ranging from 20 to 80 mM imidazole over 10 column volumes (CV) and second in a steep gradient from 80 to 500 mM imidazole over five CV. The elution buffer also contained 50 mM Tris (pH 8.0) and 500 mM NaCl. Fractions containing Ku were dialyzed overnight into a buffer containing 50 mM Tris (pH 8.0), 50 mM NaCl, and 2 mM DTT. Ku was then exchanged over a HiTrap Q HP column (GE Healthcare) under the same buffer conditions used for dialysis. Bound protein was eluted from the Q column in a shallow gradient ranging from 50 mM to 1 M NaCl over 10 CV and then in a steep gradient ranging from 1 to 2 M NaCl over five CV. Fractions containing Ku were then recaptured on a 1-mL HisTrap HP column. The nickel column was then directly connected to a Superdex 200 (GE Healthcare), and protein was injected from the nickel column onto the sizing column and complexed at room temperature for 20 min. Footprinting reactions were initiated by the addition of I2, and reactions were subsequently quenched at various time points in an equal volume of 100 mM β-mercaptoethanol. The no-I2 controls were treated identically, but the RNA was removed prior to I2 addition. Sequencing reactions were prepared by carrying out I2-mediated cleavage of the substituted RNAs at 65°C for 2 min. Reactions were precipitated by the addition of 2 vol of RNase inactivation/precipitation buffer (Life Technologies), precipitated overnight, and washed once with 70% ethanol. RNAs were then suspended in equal volumes of H2O and 2X formamide loading dye (0.5X TBE, 93.5% formamide, 30 mM EDTA, 0.5% xylene cyanol and bromophenol blue). Samples were heated to 85°C for 1 min, cooled on ice, and equal numbers of scintillation counts for each sample were loaded onto a 10% polyacrylamide sequencing gel. Gels were dried and visualized by Phosphorimager (GE Healthcare). The computer program SAFA was used to align the gel and calculate band intensities in order to determine the level of protection (Das et al. 2005). SAFA-calculated band intensities were normalized to the total number of counts per lane. The fold-change in band intensity upon addition of Ku was calculated with the normalized values. Intensities were plotted onto the secondary structure on a log2 scale.

**Nuclease V1 footprinting assays**

RNAs for footprinting reactions were prepared by in vitro transcription as described above. Final footprinting reactions contained 5′-32P-labeled TLC1 KBS RNA, V1 footprinting buffer (15 mM Tris at pH 8.0, 12% glycerol, 130 mM NaCl, 5 mM MgCl2, 1.6 mM DTT), 0.05 μg of yeast tRNA (Sigma), 1.08 μM Ku, and 0.005 or 0.0013 units of RNase V1 (Life Technologies) diluted in Ku storage buffer (25 mM Tris at pH 8.0, 20% glycerol, 200 mM NaCl, 0.5 mM EDTA, and 2 mM DTT) prior to initiation of the reaction. RNAs were annealed as described for the footprinting reactions, and Ku-KBS complexes were reconstituted for 30 min at room temperature prior to the addition of nuclease V1. Digestions were carried out at room temperature for 10 min and then quenched in RNase inactivation/precipitation buffer (Life Technologies). RNAs were precipitated and processed as described above for the phosphorothioate footprinting reactions. The alkaline hydrolysis ladder was prepared by incubating 5′-32P-labeled TLC1 KBS RNA and 2 μg of yeast tRNA (Sigma) in 50 mM sodium bicarbonate (pH 9.2) at 85°C for 4 min. The RNase T1 ladder reactions contained 5′-32P-labeled TLC1 KBS RNA, 2 μg of yeast tRNA (Sigma), 1X RNA sequencing buffer (Life Technologies), and 0.1 unit of RNase T1 (Life Technologies). RNAs were heated to 55°C for 1 min prior to the addition of RNase T1, and then incubated at 55°C for 3 min. RNase T1 and hydrolysis ladders were quenched in 1X formamide loading dye as described above. Samples were prepared, run on sequencing gels, and quantitated as described for phosphorothioate footprinting.

**Chemical modification assays**

Chemical modification and subsequent reverse transcription experiments were carried out as previously described (Zaug and Cech 1995; Brunel and Romby 2000), with minor modifications. Final reactions contained ~1 μg of TLC1-38 KBS primer extension RNA (2.7 μM), 150 mM NaCl, 25 mM HEPES (pH 8.0), and 5 mM MgCl2, and either 90 mM DMS (Sigma, diluted in 100% ethanol) or 18 mg/mL CMCT (Sigma, dissolved in H2O). RNAs were annealed as described for the footprinting experiments in the absence of DMS or CMCT.
Modification reactions were initiated by the addition of either DMS or CMCT, and reactions were quenched at various time points. For the premix negative control reactions, annealed RNA was added to the appropriate quench solution prior to adding DMS or CMCT. Denature controls consisted of modifying the RNA with the appropriate chemical reagent for 30 sec at 85°C. DMS modification was quenched by addition of 9 vol of DMS quench buffer consisting of 90 mM β-Mercaptoethanol, 333 mM NaOAC (pH 5.2), and 20 μg of glycerogen (Roche). The CMCT quench buffer was identical to the DMS quench buffer except that it lacked β-mercaptoethanol. Quenched reactions were then precipitated by the addition of 2.5 vol of ice-cold ethanol. RNAs were washed once with 70% ethanol and then resuspended in H2O.

Final reverse transcription reactions contained ~0.2 μg of modified RNA, trace amounts of 32P-labeled primer, 4 mM dNTPs, AMV reverse transcriptase (NEB), and 1X AMV buffer (NEB). Primers were annealed to RNA in 1X AMV buffer by heating to 85°C for 2 min, followed by snap cooling on ice. Reverse-transcription reactions were initiated by the addition of an equal volume of mix containing the dNTPs, 1X AMV buffer, and the reverse transcriptase. Primer extension reactions were allowed to proceed at 37°C for 40 min. Sequencing reactions were carried out with the same reverse-transcription conditions described above and the addition of a single ddNTP to a final concentration of 0.5 mM. Reverse-transcription reactions were quenched by the addition of 2X formamide loading dye and heating to 95°C for 5 min. Reactions were loaded onto a 10% sequencing gel and the gel was dried and band intensities were calculated as described for the phosphorothioate footprinting experiments. Modification levels were calculated by normalizing the SAFA calculated band intensities to the total counts per lane. The mean and standard deviation of the band intensities in a lane were used to calculate a z-score for each band in the lane.

**Determination of dissociation constants**

Dissociation constants for the mutant and additional putative yeast KBS RNAs were measured as previously described (Pfingsten et al. 2012). Briefly, trace amounts of KBS RNA were incubated with Ku at room temperature for 1 h in a buffer containing 21 mM HEPES (pH 7.5), 6 mM Tris (pH 8), 150 mM NaCl, 16% glycerol, 5 mM MgCl2, 1 mM EDTA, 25 μg/ml trNA, 0.1 mg/ml BSA, and 1 mM DTT. Ku-bound KBS RNAs were separated from free KBS RNA on a native 4%–20% acrylamide TBE gel (Life Technologies); gel boxes were placed in ice for the electrophoretic run. All measurements represent the average of three independent replicates. Mutant Kd values were normalized to the wild-type Kd for a given set of experimental replicates by multiplying the measured mutant Kd by the ratio of the measured wild-type Kd over the average wild-type Kd for all experiments. All measured wild-type KBS Kd values were within 2.5-fold of the average wild-type Kd reported in Table 1.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

**ACKNOWLEDGMENTS**

We are grateful to the members of the Cech laboratory for helpful discussions, to Jayakrishnan Mandakumar for the critical reading of this manuscript, and to Art Zaug for technical advice. T.R.C. is an investigator of the HHMI.

**Received February 6, 2013; accepted March 18, 2013.**

**REFERENCES**


Galard F, Olivier C, Dandjinou AT, Wellinger RJ, Chartrand P. 2008. *TLC1* RNA nucleo-cytoplasmic trafficking links telo-


