SHAPE Probing Reveals Human rRNAs Are Largely Unfolded in Solution

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ABSTRACT: Chemical probing experiments, coupled with empirically determined free energy change relationships, can enable accurate modeling of the secondary structures of diverse and complex RNAs. A current frontier lies in modeling large and structurally heterogeneous transcripts, including complex eukaryotic RNAs. To validate and improve on experimentally driven approaches for modeling large transcripts, we obtained high-quality SHAPE data for the protein-free human 18S and 28S ribosomal RNAs (rRNAs). To our surprise, SHAPE-directed structure models for the human rRNAs poorly matched accepted structures. Analysis of predicted rRNA structures based on low-SHAPE and low-entropy (lowSS) metrics revealed that, whereas ∼75% of Escherichia coli rRNA sequences form well-determined lowSS secondary structure, only ∼40% of the human rRNAs do. Critically, regions of the human rRNAs that specifically fold into well-determined lowSS structures were modeled to high accuracy using SHAPE data. This work reveals that eukaryotic rRNAs are more unfolded than are those of prokaryotic rRNAs and indeed are largely unfolded overall, likely reflecting increased protein dependence for eukaryotic ribosome structure. In addition, those regions and substructures that are well-determined can be identified de novo and successfully modeled by SHAPE-directed folding.

RNA is a central carrier of information in biological systems, and this information is encoded in both the primary sequence of the RNA and in higher-order structures that form when the RNA folds. Both highly stable, base-paired elements that populate single structures and unpaired or less stable structures play important biological roles, and the extent of structure can modulate RNA function by forming, making accessible, or sequestering interaction sites for proteins, other RNAs, or small-molecule ligands.1−4 Chemical probing technologies, especially the SHAPE (selective 2'-hydroxyl acylation analyzed by primer extension) strategy,5,6 have proven to be powerful tools for characterizing local nucleotide flexibility in an experimentally concise and accurate way. These nucleotide flexibility data can be parametrized7−10 and incorporated into a standard RNA folding algorithm. This melded SHAPE-directed approach, applied to many short and medium-sized RNAs and to long bacterial rRNAs, results in models that show good to outstanding accuracy when compared to accepted structures defined by comparative sequence analysis or high-resolution approaches.7−9 SHAPE-directed structure modeling can also accurately model the structures of individual functional elements in viral, bacterial, and human RNAs and has identified novel functional motifs in these long RNA.7,10−13

A current frontier in experimentally directed RNA secondary structure modeling lies in modeling complex RNAs that contain a mixture of regions with persistent stable structure intermixed with regions containing conformationally dynamic elements.14 Although it is clear that conformational dynamics are critical for the function of many RNAs, chemical probing-driven approaches have been largely validated by analysis of RNAs with very stable and well-defined structures.15,16 These test case RNAs are typically among the most highly structured RNAs known and include bacterial rRNAs and those RNAs that can be successfully crystallized.1,15−16 In fact, RNAs with such well-defined structures are highly unusual and likely represent outliers in the RNA world. Moreover, widely reported “whole transcriptome” structure probing experiments involving complex and time-intensive protocols often include minimal validation, especially as applied to large RNAs. Indeed, the RNAs or RNA motifs used to validate many “transcriptome-wide” probing experiments span ≤200 nucleotides, whereas the typical transcript in a eukaryotic cell exceeds 2000 nucleotides.

As part of an effort to validate and improve SHAPE data constraints and parameters used for analysis of long and complex RNAs and of transcriptome-wide experiments, we used the SHAPE-MaP (mutational profiling) chemical probing strategy to examine the structure of full-length human 18S and 28S rRNAs, which are 1869 and 5070 nucleotides in length, respectively. SHAPE-directed structure modeling of protein-free eukaryotic rRNAs in solution consistently showed poor overall accuracy (Figure 1A) compared to the accepted structures and this information is encoded in both the primary sequence of the RNA and in higher-order structures that form when the RNA folds. Both highly stable, base-paired elements that populate single structures and unpaired or less stable structures play important biological roles, and the extent of structure can modulate RNA function by forming, making accessible, or sequestering interaction sites for proteins, other RNAs, or small-molecule ligands.1−4 Chemical probing technologies, especially the SHAPE (selective 2'-hydroxyl acylation analyzed by primer extension) strategy,5,6 have proven to be powerful tools for characterizing local nucleotide flexibility in an experimentally concise and accurate way. These nucleotide flexibility data can be parametrized7−10 and incorporated into a standard RNA folding algorithm. This melded SHAPE-directed approach, applied to many short and medium-sized RNAs and to long bacterial rRNAs, results in models that show good to outstanding accuracy when compared to accepted structures defined by comparative sequence analysis or high-resolution approaches.7−9 SHAPE-directed structure modeling can also accurately model the structures of individual functional elements in viral, bacterial, and human RNAs and has identified novel functional motifs in these long RNA.7,10−13

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structures, taken to be base pairs visualized in crystallographic studies. This discrepancy prompted us to investigate whether the inability to model full-length human rRNAs reflects a limitation of SHAPE-directed structure modeling or represents a fundamental structural difference between deproteinized prokaryotic and eukaryotic rRNAs. Previous studies have shown that it is possible to identify well-structured regions in RNA genomes, in long noncoding RNAs, and in bacterial mRNAs using low SHAPE and low Shannon entropy metrics (lowSS regions). Using this metric, we discovered that, when examined in a protein-free solution, human rRNAs are structurally much less well-determined, and are much folded, than are bacterial rRNAs under comparable conditions. A subset of regions within the human rRNAs that passed lowSS filters likely do have stable persistent structure and could indeed be modeled accurately by SHAPE-directed folding.

## METHODS

**Reference Structures.** Structures were obtained from Ribovision curated models, revised from crystallographic and cryo-electron microscopy structures of complete ribosomes corresponding to Protein Data Bank entries 3R8S, 4GD1, 3J3A, 3J3B, 3J3D, and 3J3F, 17,19,20

**E. coli and Mammalian Cell Lysis and Protein Digestion.** A 25 ml aliquot of E. coli cells at an OD<sub>600</sub> of 0.5 was pelleted at 8000g and 4 °C for 10 min. Cells were lysed in 16.5 ml of E. coli lysis buffer [15 mM Tris (pH 8), 450 mM sucrose, 8 mM EDTA, and 0.4 mg/mL lysozyme] for 5 min at 23 °C and then for 10 min at 0 °C. Protoplasts were collected at 5000g and then resuspended in 2 ml of proteinase K buffer [50 mM HEPES (pH 8), 200 mM NaCl, 5 mM MgCl<sub>2</sub>, 1.5% sodium dodecyl sulfate (SDS), and 0.2 mg/mL proteinase K], vortexed for 10 s, and then incubated at 23 °C for 5 min and 0 °C for 10 min. For human RNA data, total RNA from HEK293 cells (80% confluency) was extracted under conditions designed to maintain the underlying RNA structure. Cells were lysed in cytoplasmic lysis buffer [40 mM Tris (pH 8.0), 40 mM NaCl, 6 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 250 mM sucrose, 0.5% Triton X-100, 250 units/mL RNase inhibitor, and 450 units/mL DNase I] for 5 min at 4 °C; nuclei were pelleted away, and SDS and proteinase K were added to final concentrations of 1.5% and 500 μg/mL, respectively. Cytoplasmic lysates were incubated at room temperature for 45 min. RNA from E. coli or human cell lysates was extracted twice with 1 volume of a phenol/chloroform/isoamyl alcohol mixture (25:24:1), pre-equilibrated in 1.1X RNA folding buffer [110 mM HEPES (pH 8.0), 110 mM NaCl, and 5.5 mM MgCl<sub>2</sub>], and twice with 1 volume of chloroform.

**SHAPE Probing of Cell-Extracted RNA.** E. coli and human RNA was exchanged into fresh 1.1X RNA folding buffer, incubated at 37 °C for 20 min, and split into two equal aliquots. One aliquot was treated with 1/9 volume of 5.2 mg/100 μL 5-nitrosourea anhydride, and the second was treated with the same volume of dimethyl sulfoxide (serving as an unmodified control). After incubation for 10 min at 37 °C, RNA was precipitated by adding 1/10 volume of 2 M ammonium acetate and 1 volume of isopropanol and incubating the mixture for 10 min at room temperature. Precipitated RNA was pelleted by centrifugation for 10 min at 12000g and 4 °C. The supernatant was removed, and the RNA pellet was washed with 1 volume of 75% ethanol and centrifuged at 7500g for 5 min at 4 °C. The samples were resuspended in water before being treated with DNase (TURBO DNase, Thermo Fisher) for 1 h at 37 °C and

**Reverse Transcription and Library Preparation of U1 RNA.** Reverse transcription was conducted on total human RNA from HEK293 cells using a gene specific primer for U1 RNA (5′-CAGGG GAAAG CGCGA A). The cDNA was purified, amplified and library preparation were performed using a two-step polymerase chain reaction (PCR), PCR products were purified, and libraries were sequenced (Illumina MiSeq instrument) as described previously.

**Structure Modeling.** Sequencing data were processed using ShapeMapper 2 software with a minimum required read depth of 1000. Superfold was used to model base pairs and pairing probabilities. RNA structures were modeled using constraints from experimental SHAPE reactivities, idealized data (1 for unpaired nucleotides and 0 for base-paired nucleotides based on the accepted structure), or no reactivity data (all reactivities set to −999). RNAs were modeled with maximum pairing distances of 600 and 800 nucleotides for the small and large subunits, respectively. Structures were scored by comparing predicted minimum free energy structures to accepted structures. Scoring allowed for a 1 bp offset. The sens value was calculated as the number of correctly predicted base pairs in the model divided by the number of canonical base pairs in the accepted structure, excluding pairs for which SHAPE data were not present at both positions. The ppv value was calculated as the number of correctly predicted base pairs divided by the number of predicted base pairs in the model, excluding no data pairs.

**Direct Reactivity Comparison.** For direct comparison of SHAPE reactivities across experiments performed under identical conditions, reactivity profiles were computed as ln(rateM/rateU), where rateM and rateU refer to MaP rates in SHAPE-modified and untreated samples, respectively. Positions with read depths below 1000 or untreated MaP rates above 0.05 were excluded.

**Identifying Low-SHAPE, Low-Entropy Regions.** Low-SHAPE, low-entropy regions were identified using a combination of SHAPE reactivity and Shannon entropy, using calculated base pairing probabilities. Median SHAPE reactivity and Shannon entropy were calculated over 51-nucleotide centered windows. Regions longer than 25 nucleotides with windowed SHAPE reactivities below 0.3 and windowed Shannon entropies below 0.08 were defined as lowSS regions. Regions were expanded to include nested base pairs that had base pairing probabilities of >90%. The selected SHAPE and entropy thresholds employed here balance region detection with model accuracy, identifying many well-structured regions in...
accurately modeled structures. Stricter (lower) thresholds increase structure modeling accuracy at the expense of detection rate, and less stringent (higher) thresholds recover more and longer lowSS regions but result in less accurate structures (Figure S1).

■ RESULTS

Structure Modeling of Full-Length rRNAs. As an initial point of comparison, the secondary structures and pairing probabilities of the *E. coli* 16S and 23S rRNAs were modeled using SuperFold with no SHAPE constraints or using experimental SHAPE data obtained for this study for RNAs gently extracted from bacterial cells (at 5 mM Mg2+) while the overall structure of the rRNA was maintained (Methods). In the absence of SHAPE constraints, the structures of the *E. coli* 16S and 23S rRNA are modeled with low accuracy in agreement with previous analyses of these and of other RNAs. The observed sensitivities (sens, percent accepted base pairs modeled correctly) and positive predictive values (ppv, percent modeled base pairs in the accepted structure) are in the range of 50–60% (Figure 1A, top; see entire rRNA columns). As expected, structure modeling accuracy notably increased with the use of SHAPE data as a pseudo-free energy change restraint to achieve a sensitivity of >90% for modeling of the 16S rRNA is obtained when the RNA is probed at 10 mM Mg2+ and regions not locally compatible with SHAPE data are omitted. For the sake of simplicity, this latter adjustment was not applied in this study, due to the complexity of defining omitted regions for the human rRNAs. True sens and ppv values are roughly 10% higher than the values reported here.

Next, we chemically probed human 18S and 28S rRNAs (the small and large subunit RNAs, respectively) in a similar cell-free state; the rRNAs were gently extracted from HEK293 cells. As expected, structure models were inaccurate without SHAPE constraints, with sens and ppv values in the range of 20–40%,...
compared to the accepted models (Figure 1A, bottom; entire rRNA columns). However, in contrast to the E. coli rRNA, the addition of SHAPE data restraints did not yield accurate secondary structure models for the full-length human rRNAs. The human 18S and 28S rRNAs were modeled with sensitivities of only 32% and 61%, respectively, meaning that the SHAPE-directed structures still deviated substantially from the accepted models (Figure 1A, bottom; Figure 1B).

Modeling rRNA Structure with Idealized Data. The SHAPE-directed pseudo-free energy change strategy was developed primarily using prokaryotic RNAs with compact structures, and one formal possibility was that the resulting parameters are not appropriate for modeling human rRNA. We therefore examined whether idealized restraints based on the accepted model of the human rRNAs would allow accurate folding. To create idealized data sets, we assigned a SHAPE reactivity of zero to nucleotides that are base-paired and a SHAPE reactivity of 1 to nucleotides that are single-stranded in the accepted secondary structures of the rRNAs. Models constrained by the idealized SHAPE data resulted in secondary structure models for the E. coli rRNAs with sens and ppv values in the range of 87−90% (Figure 1A, top). The idealized data thus yielded a significant increase in modeling accuracy.

The idealized restraints also notably improved modeling of the human rRNAs with sens values of 65% and 90% for the 18S and 28S rRNAs, respectively; some regions were modeled with very high accuracy (Figure 1B). These accuracies are not quite as high as those of E. coli rRNA models but are sufficiently high to strongly suggest that the observed disagreements between SHAPE-directed models and accepted structures are not due to a general limitation of SHAPE-constrained structure modeling but, instead, reflect fundamental biological structural differences between protein-free bacterial and human rRNAs. Indeed, there are many regions in the human rRNAs in which the observed experimental SHAPE data are simply incompatible with the accepted structure; these regions can be accurately modeled into the accepted structure when guided by idealized data, however (Figure 1B).

Direct Comparison of SHAPE Reactivities between Human and E. coli rRNA. As a second approach to establish whether the E. coli and human rRNAs show fundamentally different levels of folding, distinct from secondary structure modeling, we directly compared SHAPE reactivities between the two species. We examined SHAPE reactivity profiles over the full rRNA lengths and over selected regions in the small subunit with locally conserved sequence and structure, using raw
background-corrected SHAPE adduct-induced mutation rates from probing experiments performed under identical conditions. Global SHAPE reactivity profiles for human rRNA are substantially shifted to higher modification rates as compared to those of E. coli rRNA (Figure 2A). The human rRNAs are therefore overall intrinsically more reactive to SHAPE than are the E. coli rRNAs.

We also focused on reactivity profiles for four regions showing clear sequence and structure conservation between the human and E. coli small subunit rRNAs, defined as gap-free alignments over at least 15 nucleotides, with at least 50% sequence identity and one shared helix longer than 2 bp. In E. coli numbering, these regions were helix 8 (residues 144–178), helix 18 (500–545), helix 23 (673–717), and helix 24 (765–820), all of which have well-described roles in ribosome function. Helix 18 is a component of the mRNA entry site latch,24 and helices 8, 23, and 24 all form important bridging interactions with the large subunit.25

Helix 23 is fully or nearly fully folded in both E. coli and human rRNAs, as evidenced by shared low reactivities over base-paired positions and substantially similar reactivity profiles (Figure 2B). In contrast, the other three regions are folded in E. coli rRNA but strikingly unfolded in human rRNA, even though the expected secondary structures are similar (Figure 2C). Thus, isolated human rRNAs adopt structures notably different from those of their E. coli counterparts, even in regions with conserved sequences and functions.
Identification of Well-Structured (lowSS) RNA Regions. Prior work has emphasized that regions that are highly structured (as detected by a low local SHAPE reactivity) and have well-determined structures (supported by a low Shannon entropy) tend to be strongly correlated with function.\textsuperscript{10--12} These regions might comprise well-folded elements within the human rRNAs where structure can be more accurately predicted by SHAPE-directed structure modeling. Local regions of low SHAPE and low Shannon entropy (lowSS) within the \textit{E. coli} and human rRNAs were identified using a combination of SHAPE data and SHAPE-informed Shannon entropy, calculated over smoothed windows across the primary sequence.\textsuperscript{6,10,12} Although

Figure 4. Structural characterization of the human 28S rRNA. (A) Model of the full-length 28S rRNA with arcs representing correct, incorrect, and missing base pairs relative to the accepted structure (top). SHAPE reactivity (black) and entropy (brown) shown as medians over 51-nucleotide centered windows along the full-length 28S rRNA with the axes crossing at 0.3 and 0.08, respectively (middle). SHAPE-directed structure models for lowSS regions modeled as independent elements (bottom). (B) Nucleotide-level models of selected regions, highlighting plausible SHAPE-supported local refolding. Local structures that are clearly not compatible with the accepted structure are boxed (defined as nucleotides with high reactivity in base-paired regions or low reactivity in consecutive unpaired regions). Nucleotides are colored by SHAPE reactivity (see inset to Figure 1).
SHAPE reactivity influences the calculation of Shannon entropy, the SHAPE and entropy metrics provide orthogonal information, and the combination of the two terms identifies regions whose secondary structures are modeled with the highest accuracy (Figure S1). Reactivity and entropy thresholds were chosen to maximize the overall number of nucleotides modeled while maintaining high secondary structure modeling accuracy.

We identified well-structured regions in both E. coli and human rRNAs by the lowSS criteria and then modeled each using SHAPE restraints. For the E. coli rRNAs, focusing on the lowSS regions had a small, but positive, effect on the already high model accuracy (Figure 1A, top). LowSS regions comprise 81% and 69% of the E. coli 16S and 23S rRNAs, respectively, and these extensive regions of well-defined structure allow both RNAs to be modeled with high accuracy, even outside of the lowSS regions. In the human 18S rRNA, seven well-structured regions were identified, and these were modeled with a base pair sens of 76%, a dramatic improvement over the 32% sensitivity for the full-length rRNA (Figures 1A and 3). The human 28S rRNA model contained 14 lowSS regions, and these were modeled with 77% sens (Figures 1 and 4).

This analysis reveals that, although SHAPE data alone are ill-suited for recovering the accepted structures of full-length human rRNAs, the low-SHAPE, low-entropy strategy robustly identifies local well-defined secondary structures. Well-structured regions that pass the lowSS filters in human rRNA involve important functional elements such as the L1 stalk, the GTPase-associated center, the A site finger, and an element of the 18S rRNA that directly interacts with mRNA in the E site26−30 (Figure S2).

**Local rRNA Refolding.** The lowSS regions in the human rRNAs were generally modeled to good agreement with the accepted structures. However, SHAPE reactivities in some regions, including some lowSS regions, are clearly not consistent with the accepted structures in regions of both human 18S and 28S rRNAs (Figures 1B and 4B, boxed residues). We interpret these data as indicating that, for RNAs chemically probed in a cell-free (and protein-free) state, models for lowSS regions are better representations of these regions than are the accepted structures, likely because these regions locally refold when RNA is removed from the cellular (and native protein) environment.

**DISCUSSION**

This work supports three overarching conclusions. First, folding of the protein-free bacterial and human rRNAs is fundamentally different: in solution, the human rRNA structure is more
unfolded and less well-determined than that of the bacterial rRNAs (Figures 1 and 2). Second, algorithms that couple chemical probing data with empirical free energy change relationships cannot recover the accepted structure of all RNAs with high accuracy. This inability can be attributed to RNAs that rarely populate the accepted structure, as notably observed here for the human rRNAs, and to small errors in thermodynamic and chemical reactivity parameters that especially limit the accuracy of modeling long-range interactions.31,32 Third, the low-SHAPE reactivity, low-Shannon entropy (lowSS) metric is a powerful approach for characterizing the well-determinedness of folding for large RNAs and for addressing the challenge presented by conclusion 2. The lowSS metric identifies the subset of regions in long RNAs that can be modeled accurately by SHAPE-directed folding (Figures 1, 3, and 4).

SHAPE provides empirical information about local nucleotide flexibility that can be used to model the secondary structure of short RNAs, bacterial rRNAs, and numerous other RNAs— which have stable and well-defined structures—with high accuracy. However, SHAPE-directed folding alone can produce misleading or inaccurate models for long RNAs that contain poorly structured regions or regions capable of adopting multiple conformations. This challenge in de novo modeling of complex RNAs can be addressed, in part, by focusing on those regions within a large RNA that do fold to form a well-defined structure. The lowSS filter appears to be robust for both short and long RNAs and across RNAs with different levels of structure. For example, a majority of nucleotides in the highly structured human 18S rRNA (78%) meet the lowSS criterion (Figure SA). Both E. coli 16S and 23S rRNAs are well-structured with 81% and 69% of nucleotides, respectively, in lowSS regions (Figure SB). On the basis of these three RNAs, both short and long RNAs can have extensive lowSS regions. In contrast, the human 18S and 28S rRNAs, with only 34% and 43% of nucleotides in lowSS regions, respectively, have extensively poorly structured regions (Figure SC). Many regions of human rRNA are simply unfolded, despite sharing conserved sequence and structure with E. coli rRNA (Figure 2).

Our data emphasize that human rRNA is intrinsically less structured than E. coli rRNA under the conditions probed here. This study also suggests that the human rRNAs may not be well suited for use in validating modeling accuracies for transcriptome-wide studies. Eukaryotic rRNAs likely require more support from proteins for full structure formation than do bacterial rRNAs, consistent with models of ribosome evolution that show the accretion of a protein shell in eukaryotic lineages.33,34 Eukaryotic ribosome structure, assembly, and regulation are all more complex than those of prokaryotic ribosomes.35 Other aspects of the eukaryotic cellular environment, such as local ion concentrations or differences in macromolecular crowding, might play larger roles for eukaryotic ribosome assembly and folding than for assembly of the bacterial translation machinery. Unlike the relatively stable “RNA rocks” of prokaryotic ribosomes, only a small fraction of eukaryotic rRNA appears to be well folded in isolation.

Eukaryotic rRNAs, multikilobase mRNAs,12 viral RNAs,10,11 and noncoding RNAs1 appear to be less structured overall than well-understood short structured RNAs like bacterial riboswitches6,36 but do have regions of stable functional structures. In such cases, lowSS regions provide a starting point for locating well-folded and potentially functional structural elements. Indeed, lowSS regions identify the vast majority of well-characterized functional motifs in viral genomic RNAs,10,11,13 and the lowSS metric enabled de novo identification and validation of multiple novel regulatory elements in the E. coli transcriptome.12 Similarly, well-structured lowSS regions in the human rRNAs include important functional elements. The lowSS metric will likely prove to be a broadly useful tool for future modeling and functional studies of diverse classes of long RNA transcripts.

## ASSOCIATED CONTENT

† Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.9b00076.

Two figures and tables giving the full SHAPE data for the E. coli and human rRNAs (PDF)

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Notes

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