

Genome-wide measurement of RNA secondary structure in yeast

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The structures of RNA molecules are often important for their function and regulation^{1–6}, yet there are no experimental techniques for genome-scale measurement of RNA structure. Here we describe a novel strategy termed parallel analysis of RNA structure (PARS), which is based on deep sequencing fragments of RNAs that were treated with structure-specific enzymes, thus providing simultaneous *in vitro* profiling of the secondary structure of thousands of RNA species at single nucleotide resolution. We apply PARS to profile the secondary structure of the messenger RNAs (mRNAs) of the budding yeast *Saccharomyces cerevisiae* and obtain structural profiles for over 3,000 distinct transcripts. Analysis of these profiles reveals several RNA structural properties of yeast transcripts, including the existence of more secondary structure over coding regions compared with untranslated regions, a three-nucleotide periodicity of secondary structure across coding regions and an anti-correlation between the efficiency with which an mRNA is translated and the structure over its translation start site. PARS is readily applicable to other organisms and to profiling RNA structure in diverse conditions, thus enabling studies of the dynamics of secondary structure at a genomic scale.

Existing experimental methods for measuring RNA structure can only probe a single RNA structure per experiment and are typically limited in the length of the probed RNA (Supplementary Note 1). To measure structural properties of many different RNAs simultaneously, we extracted polyadenylated transcripts from yeast growing in the log phase, renatured the transcripts *in vitro* and treated the resulting pool with RNase V1 and, separately, with S1 nuclease. RNase V1 preferentially cleaves phosphodiester bonds 3' of double-stranded RNA, whereas S1 nuclease preferentially cleaves 3' of single-stranded RNA⁷. Thus data from these two complementary enzymes should allow us to measure the degree to which each nucleotide was in a single- or double-stranded conformation (Fig. 1). We chose renaturation and enzymatic cleavage conditions under which the cleavage reactions occur with single-hit kinetics (Supplementary Fig. 1a, b) and where intramolecular, but not intermolecular, RNA–RNA interactions are dominant (Supplementary Fig. 1c, d). As a control, we also added two short RNA domains from HOTAIR, a human non-coding RNA⁸, and from the structurally known *Tetrahymena* group I intron ribozyme⁹.

We devised a ligation method specifically to ligate V1- and S1-cleaved RNA to adaptors, and converted them into complementary DNA (cDNA) libraries suitable for deep sequencing (Supplementary Fig. 2). As both enzymes leave a 5' phosphate at the cleavage point and because only 5' phosphoryl-terminated RNAs are capable of ligating to our adaptors, we enrich for V1- and S1-cleaved fragments and select against random fragmentation and degradation products

that typically have 5' hydroxyl (Supplementary Fig. 3). Thus each observed cleavage site provides evidence that the cut nucleotide was in a double-stranded (for V1-treated samples) or single-stranded (for S1-treated samples) conformation. As a quantitative measure at nucleotide resolution representing the degree to which a nucleotide was in a double- or single-stranded conformation, we took the log ratio between the number of sequence reads obtained for each nucleotide in the V1 and S1 experiments. A higher (lower) log ratio, or PARS score, thus denotes a higher (lower) probability for a nucleotide to be in a double-stranded conformation.

We performed four independent V1 experiments and three independent S1 experiments, which were highly reproducible across replicates (correlation = 0.60–0.93, Supplementary Table 1), resulting in over 85 million sequence reads that map to the yeast genome, of which approximately 97% mapped to annotated transcripts (Supplementary Table 2). At an average nucleotide coverage above 1.0, we obtained structural information for over 3,000 yeast transcripts (Supplementary Table 3 and Supplementary Fig. 4a), covering in total over 4.2 million transcribed bases, which is approximately 100-fold more than all published RNA footprints to date.

We used several tests to check for biases in our method. We found that RNase cleavage, adaptor ligation and cDNA conversion do not introduce significant sequence biases (Supplementary Fig. 5), that our protocol has a very small bias towards particular regions along the transcript (Supplementary Fig. 6) and that we capture RNA fragments in proportion to their abundance in the initial pool (Supplementary Fig. 4b, c). We also confirmed that signals generated by RNase V1 are highly distinct from those generated by S1 nuclease. Global inspection across all transcripts revealed that approximately 7% of the V1 and S1 peaks are shared (Methods, Supplementary Table 4 and Supplementary Fig. 7). These joint peaks could be the result of experimental noise introduced by non-specific enzymatic activity, but could also correspond to dynamic RNA regions or transcripts that fold into more than one stable conformation.

To test whether PARS accurately measures RNA structures, we first confirmed that its signals are similar to those obtained with traditional footprinting. To this end, we performed ten separate footprinting experiments with either RNase V1 or S1 nuclease, on two domains from the *Tetrahymena* ribozyme, two domains from the human HOTAIR non-coding RNA, which we doped into our samples, and two domains of endogenous yeast mRNAs. In all cases, we found high agreement between our PARS signals and footprinting (correlations = 0.40–0.97; Fig. 2 and Supplementary Figs 8–10). Notably, owing to length limitations of footprinting, we had to select short domains from each of the above transcripts, transcribe them *in vitro* and then apply footprinting. Thus footprinting may be inaccurate,

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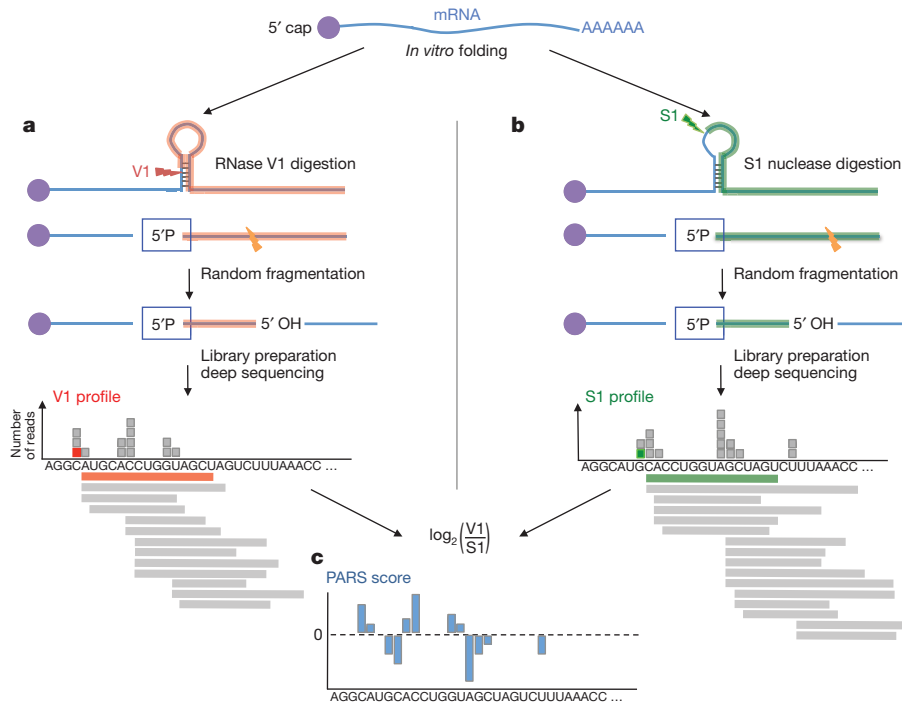


Figure 1 | Measuring structural properties of RNA by deep sequencing. **a**, RNA molecules are cleaved by RNase V1, which cuts 3' of double-stranded RNA, leaving a 5' phosphate (5'P). One such cut is illustrated by a red arrow. After random fragmentation, V1-generated fragments are specifically captured and subjected to deep sequencing. Each aligned sequence provides structural evidence about a single base. The marked red square illustrates the evidence obtained from one mapped sequence (red). Additional evidence (grey boxes) is collected by mapping more sequences (grey horizontal bars).

because lacking long-range interactions, the excised fragment could fold differently when taken out of context. In contrast, PARS can probe RNAs in their full-length context.

Next, we compared PARS with reported structures of yeast coding and non-coding RNAs, and found that it correctly reproduces the known secondary structure of three structured RNA domains of ASH1 (ref. 10), of a structural element in URE2 mRNA¹¹ and of the glutamate transfer RNA (Fig. 2e, f and Supplementary Figs 11 and 12). This suggests that PARS can provide structural information of transcripts in their full-length context and endogenous abundance from within a complex RNA pool. Taken together, our analyses demonstrate that PARS recapitulates results obtained by low-throughput methods with high accuracy, and has advantages over existing methods, stemming from its ability to probe structures of long RNAs.

As another independent validation of PARS, we compared it with computational predictions of RNA structure, by applying the Vienna package¹² to the 3,000 transcripts that we analysed. We found a significant correspondence between these predictions and our PARS scores, whereby nucleotides with high (low) double-stranded PARS score had a significantly higher (lower) average predicted pairing probability ($P < 10^{-200}$; Fig. 3a and Supplementary Fig. 13). Despite this significant global correspondence, there are large differences between PARS and predictions, in part owing to noise in our approach but also because of known inaccuracies of folding algorithms. We thus suggest that genome-wide PARS data can be used to constrain folding algorithms and improve their accuracy, as previously shown for specific RNAs^{13,14} (Supplementary Fig. 15).

We used the obtained structural profiles to investigate five global properties of yeast transcripts. First, examining the average PARS score across the coding regions and untranslated regions (UTRs), we found that coding regions exhibit significantly more pairing than 5' and 3' UTRs ($P < 10^{-30}$ and $P < 10^{-50}$, respectively; Fig. 3c).

A large number of reads aligned to the same base indicates that the base is cleaved many times by RNase V1 and is thus more likely to be in double-stranded conformation. **b**, Same as **a**, but the RNA sample is treated with S1 nuclease, which cuts 3' of single-stranded RNA. Collected reads in this case suggest that the base was unpaired in the original RNA structure. **c**, By combining the data extracted from the two complementary experiments **a** and **b**, we obtain a nucleotide-resolution score representing the likelihood that the inspected base was in a double- or single-stranded conformation.

Notably, the start and stop codons each exhibit local minima of PARS scores, indicating reduced tendency for double-stranded conformation and increased accessibility. These findings agree with previous computational predictions for mouse and human genes¹⁵. The evolutionary conservation of this global organization of mRNA secondary structure suggests that it may have functional importance. An overall unstructured background in UTRs may allow functional elements to stand out and, conversely, highly paired domains along coding regions may protect against ectopic translation initiation, or regulate ribosome translocation and protein folding, as recently postulated¹³.

Second, aligning our measured transcripts about their start codon and applying a discrete Fourier transform analysis to the average PARS signal, we detected a periodic structure signal across coding regions with a cycle of three nucleotides, such that, on average, the first nucleotide of each codon is least structured and the second nucleotide is most structured. Notably, this periodic signal is only found in coding regions and not in UTRs (Fig. 3b), and the degree of three-nucleotide periodicity in transcripts is significantly associated with ribosome density *in vivo*¹⁶ (Supplementary Fig. 14), suggesting that this periodicity may directly or indirectly facilitate translation.

Third, we tested whether there is a correlation between mRNA structure around the translation start site and translation efficiency. Such a relation has long been hypothesized¹⁷ and recently shown for one reporter protein in *E. coli*¹⁸. We found a small but significant anti-correlation between PARS scores at the region located approximately 10 base pairs (bp) upstream of the translation start site and ribosome density throughout the transcript¹⁶, a proxy for translational efficiency (correlation = -0.1 , $P < 10^{-4}$; Fig. 4a). Intriguingly, the -10 -bp region corresponds to the 5' position of the first ribosome on yeast mRNAs¹⁶. To examine this relation in more detail, we applied *k*-means clustering ($k = 4$) to the PARS structural profile of the ± 40 bp surrounding the translation start site. Notably, genes from clusters 3 and

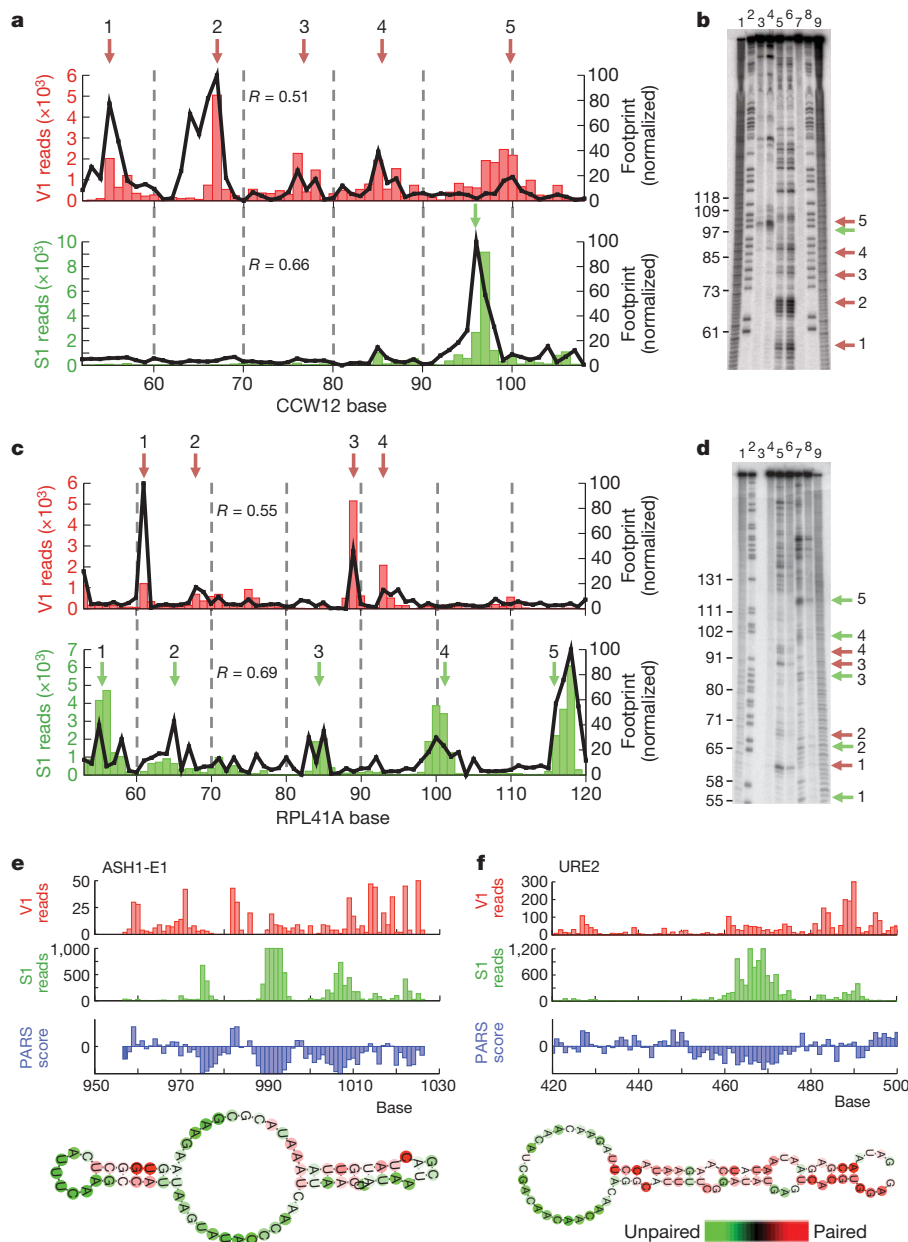


Figure 2 | PARS correctly recapitulates results of RNA footprinting and known structures.

a, The PARS signal obtained for bases 50–110 of the yeast gene *CCW12* using the double-stranded cutter RNase V1 (red bars) or the single-stranded cutter S1 nuclease (green bars) accurately matches the signals obtained by traditional footprinting of that same transcript domain (black lines). The PARS signal is shown as the number of sequence reads that mapped to each nucleotide; footprinting results are obtained by semi-automated quantification of the RNase lanes shown in **b**. The red arrows indicate RNase V1 cleavages, the green arrows indicate S1 nuclease cleavages as shown in the gel (**b**). **b**, Gel analysis of RNase V1 (lanes 5, 6) and S1 nuclease (lanes 3, 4) probing of *CCW12*. Additionally, RNase T1 ladder

(lanes 2, 8), alkaline hydrolysis (lanes 1, 9) and no RNase treatment (lane 7) are shown. **c**, The PARS signal obtained for bases 50–120 of the yeast gene *RPL41A* matches the signals obtained by traditional footprinting. **d**, RNase V1 (lanes 5, 6) and S1 nuclease (lanes 7, 8) probing of *RPL41A*, RNase T1 ladder (lane 2), alkaline hydrolysis (lanes 1, 9) and no RNase treatment (lane 4). **e**, **f**, Raw number of reads obtained using RNase V1 (red bars) or S1 nuclease (green bars) and the resulting PARS score (blue bars) along one inspected domain of *ASH1* (**e**) and *URE2* (**f**). Also shown are the known structures of the inspected domains with nucleotides colour-coded according to their computed PARS score. The Pearson correlations (R) between PARS and traditional footprinting are indicated.

4 exhibit significantly less structure in their 5' UTR than in the beginning of their coding region, as well as a higher ribosome density (Fig. 4b). Overall, these results provide the first genome-wide experimental validation for the suggestion that mRNA secondary structure around the start codon may reduce translational efficiency¹⁷, although the low correlation we found implies that, *in vivo*, translational efficiency is determined by additional factors.

Fourth, we asked whether genes with shared biological functions or cytoplasmic localizations¹⁹ tend to have similar PARS scores, indicative of similar degrees of secondary structures. We found a rich picture of biological coordination (Supplementary Fig. 16 and Supplementary

Table 5), including increased RNA structure, especially in coding regions, in transcripts whose encoded proteins localize to distinct cellular domains or participate in distinct metabolic pathways. We also found that mRNAs with the least secondary structure in their 5' UTR and coding sequences encode subunits of the ribosome.

Finally, we examined the PARS score of transcripts predicted to encode a signal peptide, because a recent study showed that RNA sequences encoding the signal sequence (termed the signal sequence coding region (SSCR)) of secretory proteins function as RNA elements that promote RNA nuclear export²⁰. We found that the 5' UTR region and approximately first 30 coding nucleotides of signal

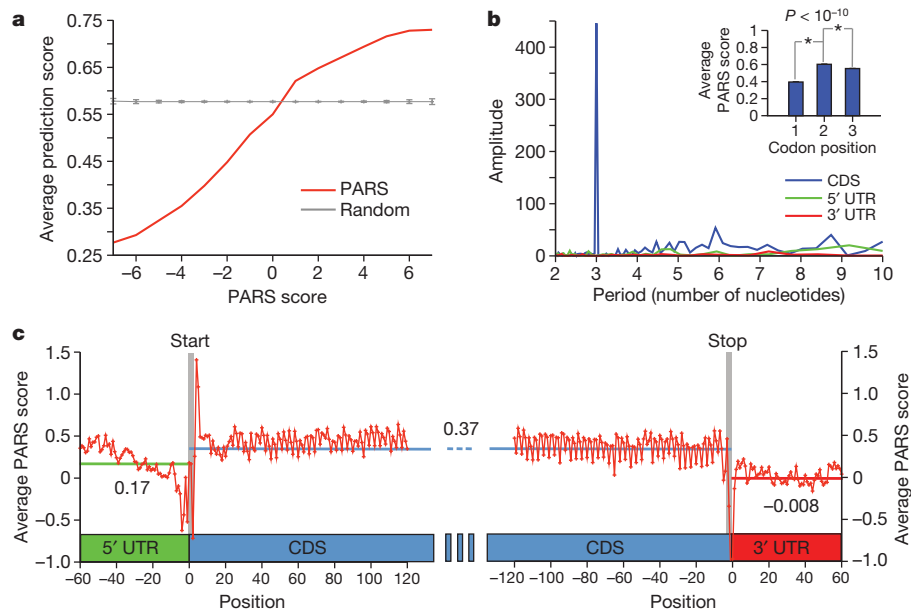


Figure 3 | Functional units of the transcript are demarcated by distinct properties of RNA structure. **a**, Significant correspondence between PARS and computational predictions of RNA structure. We used the Vienna package¹² to fold the 3,000 yeast mRNAs used in our analysis, and extracted the predicted double-stranded probability of each nucleotide. The average predicted double-stranded probability of each nucleotide (y axis) is shown, where nucleotides were sorted by their PARS score (x axis). Average and standard deviation from 1,000 shuffle experiments in which a random prediction score was assigned to

each probed base are shown in grey. **b**, Discrete Fourier transform of average PARS score across the coding region, 3' UTR and 5' UTR. Inset shows PARS score obtained for each of the three positions of every codon, averaged across all codons. **c**, PARS score across the 5' UTR, the coding region and the 3' UTR, averaged across all transcripts used in our analysis. Transcripts were aligned by their translational start and stop sites for the left and right panel, respectively; start and stop codons are indicated by grey bars; horizontal bars denote the average PARS score per region.

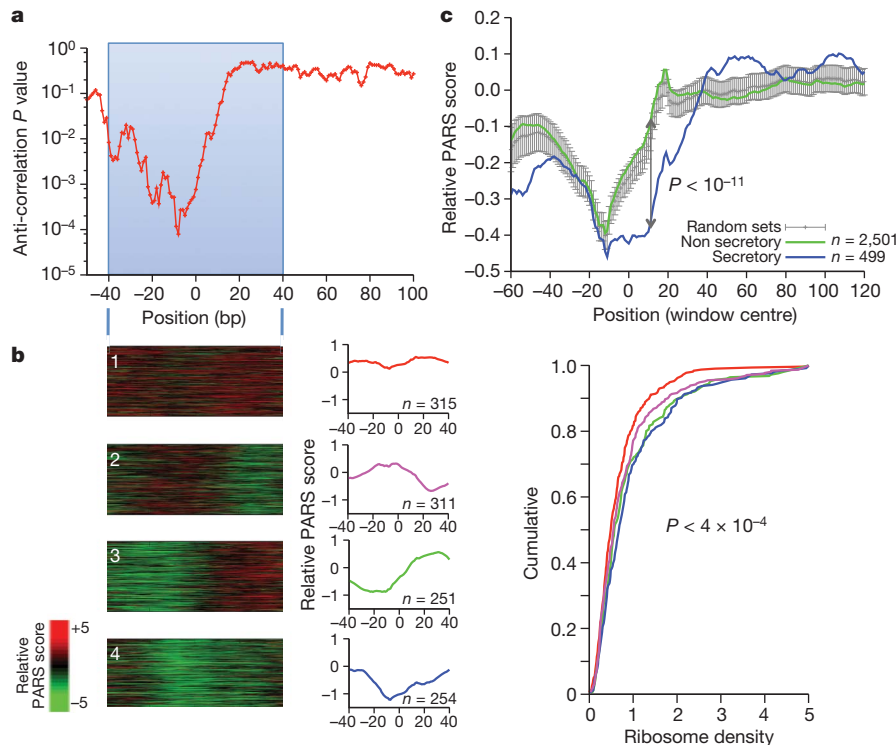


Figure 4 | Structure around start codons correlates with low translational efficiency. **a**, Sliding window analysis of local PARS score and ribosome density¹⁶. The significance (P value) of the anti-correlation between average PARS score along a 40-bp-wide window and the reported ribosome density is shown. **b**, Left, k-means clustering of PARS scores across the ± 40 -bp window surrounding the translation start site of all transcripts for which enough coverage was obtained. The average structural profile and number of member genes is shown to the right of each cluster. Right, Cumulative distribution plot of ribosome occupancy for each cluster and the associated Kolmogorov–Smirnov test P value between the distribution of clusters 1 and

4. **c**, Tendency for less RNA structure in the first 30 bases of open reading frames encoding predicted secretory proteins. Although structure typically builds up immediately upon entry to the coding sequence, genes predicted to code for secretory proteins retain low structure in approximately the first 30 bases of the coding sequence, consistent with a dual function of SSCRs in both protein coding and targeting of the mRNA²⁰. The figure shows average relative PARS scores (Methods) across a 30-bp sliding window for the 499 genes coding for secretory proteins (blue), the remaining 2,501 genes (green) and the mean and standard deviation obtained from 1,000 shuffle experiments in which sets of 499 genes were randomly selected (grey).

peptide transcripts have a lower PARS signal, indicating increased single-stranded propensity compared with other transcripts ($P < 10^{-11}$; Fig. 4c). Because RNA sequences encoding the signal sequence typically reside in the beginning of the coding region, these results suggest that specific secondary RNA structure around gene starts may assist in the cytotopic localization of mRNAs and their resulting proteins. More generally, we suggest that PARS can be used both to generate and test hypotheses of signals of secondary structure that may characterize and have functional importance for classes of mRNAs.

In summary, we introduced PARS, the first high-throughput approach for genome-wide experimental measurement of RNA structural properties, and showed that it recovers structural profiles with high accuracy and at nucleotide resolution. Like most existing methods, one limitation of PARS is that it maps RNA structures *in vitro*, and its reported structures may thus differ significantly from the *in vivo* conformations. This may be addressed in the future by using reagents that can probe RNA structure in living cells⁷, but it will require new methods to adapt to deep sequencing. Overall, PARS transforms the field of RNA structure probing into the realm of high-throughput, genome-wide analysis and should prove useful both in determining the structure of entire transcriptomes of other organisms and in systematically measuring the effects of diverse conditions on RNA structure. Probing RNA structure in the presence of different ligands, proteins or in different physical or chemical conditions may provide further insights into how RNA structures control gene activity.

METHODS SUMMARY

Sample preparation. Total RNA was extracted from yeast grown at 30 °C to exponential phase in yeast peptone dextrose (YPD) medium by using hot acid phenol. Poly(A)⁺ RNA was obtained by purifying it twice using the Poly(A) Purist Kit. Supplementary Fig. 2 shows the PARS protocol.

Sequencing library construction. RNA was folded and probed for structure using 0.01 U RNase V1 (Ambion), or 1,000 U of S1 nuclease (Fermentas), in a 100- μ l reaction volume. A modified version (see Supplementary Methods) of the SOLiD Small RNA Expression Kit was used to convert fragments into a sequencing library.

SOLiD sequencing and mapping. cDNA libraries were amplified onto beads and subjected to emulsion PCR, according to the standard protocol described in the SOLiD Library Preparation Guide. Obtained sequences were truncated to 35 bp, and required to map uniquely to either the yeast genome or transcriptome, allowing up to one mismatch and no insertions or deletions.

Computing the PARS score. The PARS score is defined as the \log_2 of the ratio between the number of times the nucleotide immediately downstream of the inspected nucleotide was observed as the first base when treated with RNase V1 and the number of times it was observed in the S1 nuclease treated sample. To account for differences in overall sequencing depth between the V1- and S1-treated samples, the number of reads for each nucleotide is normalized before the computation of the ratio.

Periodicity. Periodicity was analysed by applying a discrete Fourier transform to the average PARS score collected from the following genomic features: the last 100 bases of the 5' UTR, the first 200 bases of the coding sequence and the 100 first bases of the 3' UTR.

Online resources. Nucleotide-resolution raw reads and PARS scores for the 3,000 genes included in our analysis can be visualized and downloaded at <http://genie.weizmann.ac.il/pubs/PARS10> or using the PARS iPhone App.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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