

MINIREVIEW

SnoPatrol: how many snoRNA genes are there?

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Abstract

Small nucleolar RNAs (snoRNAs) are among the most evolutionarily ancient classes of small RNA. Two experimental screens published in *BMC Genomics* expand the eukaryotic snoRNA catalog, but many more snoRNAs remain to be found.

What are snoRNAs?

The biosynthesis of eukaryote ribosomes is complex, involving numerous processing events to generate mature ribosomal RNAs (rRNAs) and the subsequent assembly of processed rRNAs with dozens of ribosomal proteins. Small nucleolar RNAs (snoRNAs) are central to ribosome maturation, being required in key cleavage steps to generate individual rRNAs, and in their capacity as guides for site-specific modification of rRNA. In the rRNA of the budding yeast *Saccharomyces cerevisiae*, on the order of 100 snoRNA-guided modifications are made during the biosynthesis of a single ribosome; this number is approximately double in humans. Around half of these modifications are methylations of the 2' position on ribose, and are carried out by C/D-box small nucleolar ribonucleoproteins (snoRNPs), which consist of a guide snoRNA acting in concert with several proteins, including Nop1p, the RNA methylase component of the snoRNP. The remaining modifications produce pseudouridine, an isomer of uridine, and are guided by H/ACA-box snoRNPs, with the Cbf5p subunit performing the pseudouridylation reaction [1]. Figure 1 illustrates the interaction between the two types of snoRNA and their respective RNA targets.

Over the past decade, the snoRNA universe has expanded rapidly. H/ACA- and C/D-family RNAs have been discovered in Archaea (where they are dubbed sRNAs, as Archaea lack nucleoli), and likewise modify

rRNA, and in the Cajal body of the eukaryote cell (small Cajal body scaRNPs), where they modify small nuclear RNAs (snRNAs), the RNA constituents of the spliceosome [2]. Recently, HBII-52, a human C/D snoRNA, has been shown to regulate splicing of serotonin receptor 2C mRNA, indicating a wider role in gene regulation [3], and another C/D snoRNA has been shown to be expressed from the Epstein-Barr virus genome [4]. As our knowledge of snoRNAs expands beyond RNA modification and hints at wider regulatory roles, there is a need to identify the full repertoire of snoRNAs in a genome and establish when and on what RNAs they act. Against this backdrop, experimental screens that trawl organism-by-organism for snoRNAs are vital, as bioinformatic screens have so far failed to provide a robust computational alternative to labour-intensive experimental methods of RNA identification. Two recent papers in *BMC Genomics* by Zhang *et al.* [5] and Liu *et al.* [6] report the identification of novel snoRNAs from the rhesus monkey *Macaca mulatta* and the filamentous fungus *Neurospora crassa*, respectively. Both sets of authors experimentally investigated snoRNA pools by sequencing cDNAs derived from RNA extracted from their species of interest. Subsequent bioinformatics analysis was used by each group to classify sequences as either of the two snoRNA classes or otherwise. These approaches netted 48 H/ACA and 32 C/D box snoRNAs in the monkey and 20 H/ACA and 45 C/D box snoRNAs in the fungus. Studies like these are vital to the extension of our knowledge of how complements of snoRNAs vary through evolution. Given the intense effort required for such analyses, it is worth taking stock and asking, where are the current gaps in our knowledge of snoRNAs?

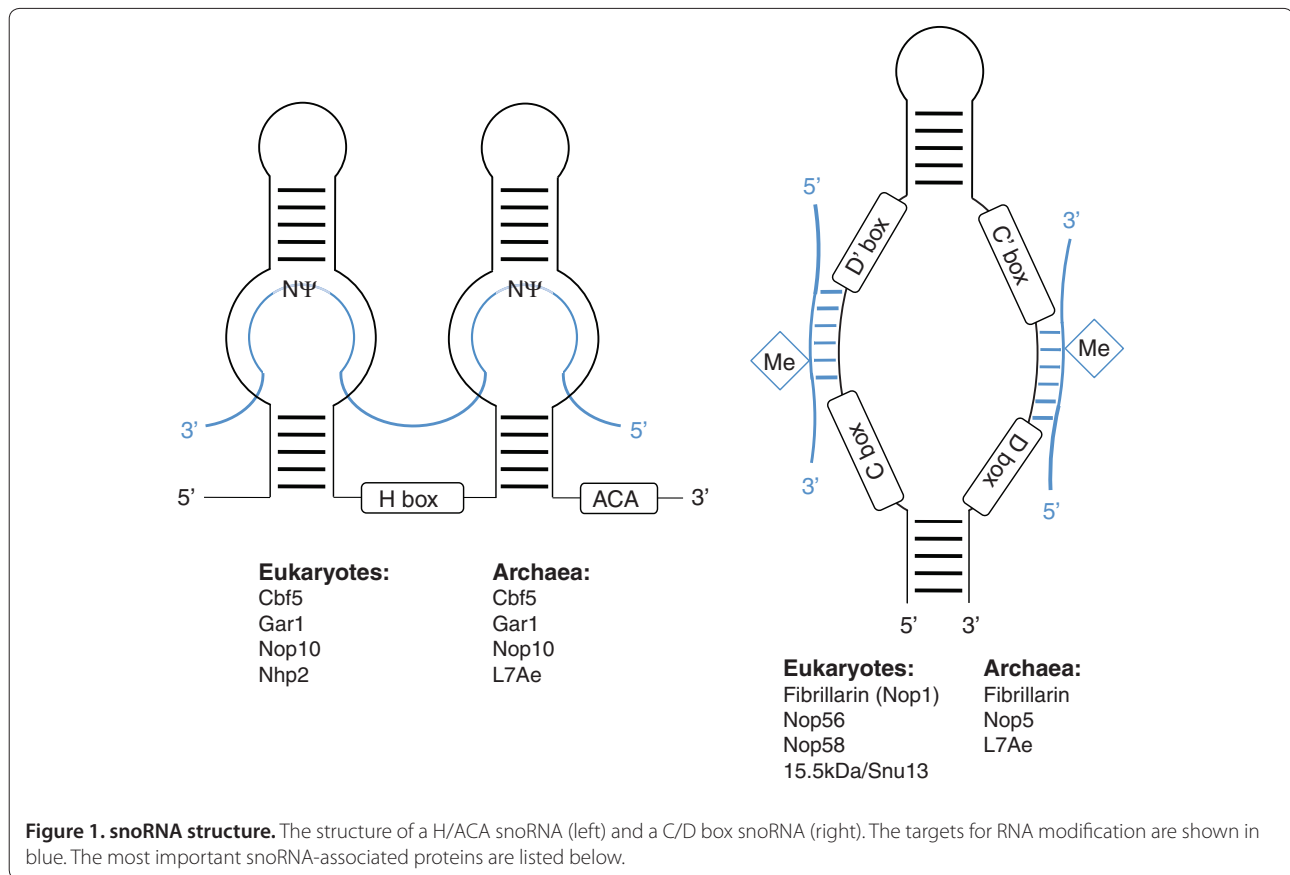
The taxonomic distribution of known snoRNAs

To investigate the taxonomic distribution of the known snoRNAs and highlight where potential new discoveries can be made, we have gathered data from the Pfam (protein families), Rfam (RNA families), Genomes Online (GOLD) and EMBL databases (Figure 2). The Rfam database uses experimentally validated ncRNA sequences that have been deposited in EMBL to search for homologous sequences across all nucleotide sequences (see the red and pink bars in Figure 2). The results show

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that for many major taxonomic clades there are few or no known snoRNAs annotated.

In the Archaea, annotated snoRNAs are notably absent from the taxon *Halobacterium*, for which a genome sequence has been available for nearly 10 years and which has been proposed to contain snoRNAs on the basis of the presence of the snoRNP-associated proteins fibrillarin and Nop56/58 [7]. In fact, only 33% of the crenarchaeal and 60% of the euryarchaeal groups carry known or predicted snoRNAs, and numbers of snoRNAs are very low in the Euryarchaeota. Still within the Archaea, snoRNAs have been annotated in some methanococcal genomes, predicted on the basis of homology to experimentally validated snoRNAs from members of the Thermoprotei [8].

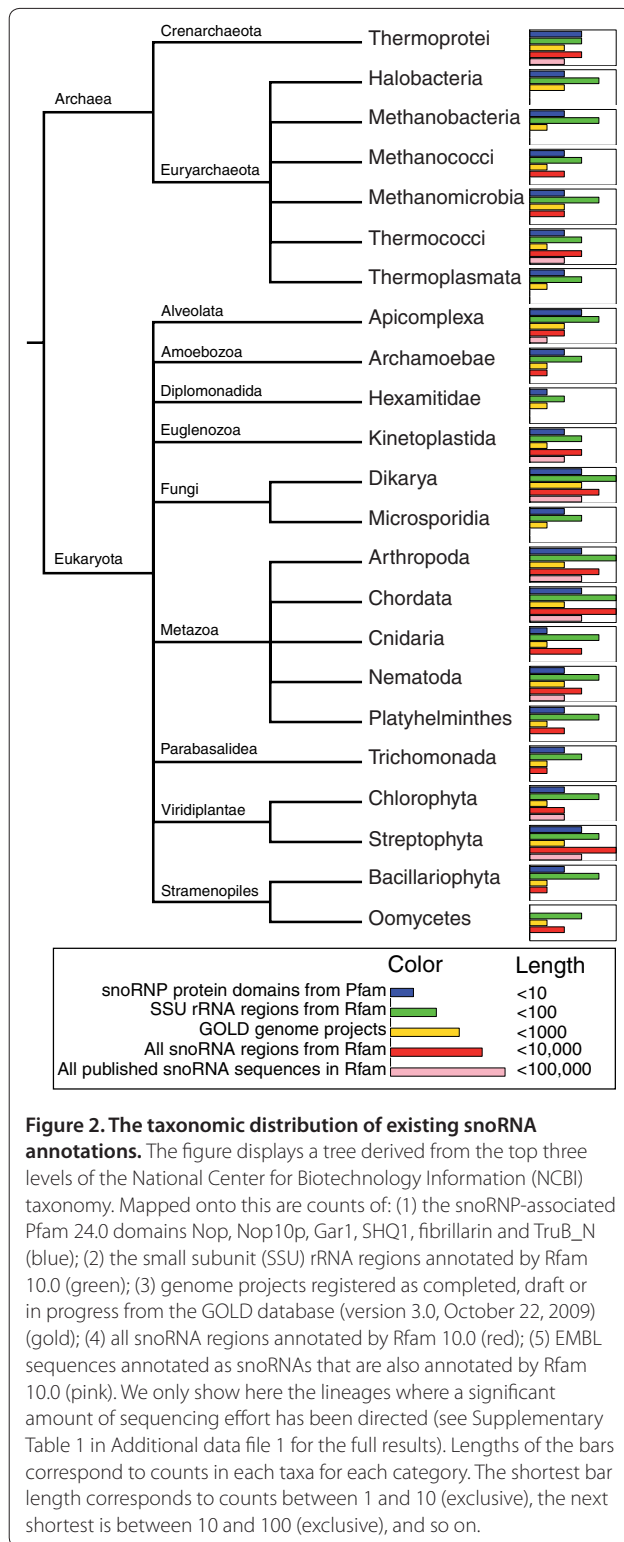
Some eukaryotic taxa fare little better. For example, in the unicellular diplomonads (Diplomonadida; Figure 2), such as *Giardia lamblia*, there are no snoRNA families listed in Rfam, although putative snoRNA-like RNAs have been reported from *G. lamblia* [9,10]. Databases such as Rfam inevitably lag behind the current literature; we expect that these missing snoRNAs will be included in future releases.

The case of the microsporidia (unicellular organisms allied to the fungi) is interesting in that one genome

sequence was published nearly a decade ago and eight further projects are in progress, yet despite this apparent wealth of information no snoRNAs have been identified. But like diplomonads, microsporidia clearly have components of the snoRNA machinery and almost certainly utilize snoRNAs. The absence, therefore, is due to the fact that snoRNAs have not been experimentally determined, and current bioinformatics methods are not sensitive enough to reliably identify snoRNAs in these taxa from sequence analyses alone, so none have been inferred by homology.

As expected, the Metazoa are comparatively well studied; there is a host of supporting experimental and bioinformatics evidence for snoRNAs across the metazoa, with the exception of the Cnidaria and the Platyhelminthes, which currently only have bioinformatically predicted snoRNAs based upon sequence similarity to other metazoan snoRNAs.

The genome sequence for the parasitic protozoan *Trichomonas vaginalis* (a parabasalid; Figure 2) bears one lonely C/D-box snoRNA annotation for a homolog of the fungal snoRNA snR52/Z13. Furthermore, this is a rather low-scoring hit (26.12 bits, E-value = 1.04e+02) to an otherwise exclusively fungal family and the *Trichomonas* sequence has some differences from the canonical C- and



D-box motifs, suggesting that the prediction may be spurious (Additional file 1). In contrast, the two main groups of green plants (Viridiplantae), the Streptophyta (multicellular green plants and some green algae) and

Chlorophyta (green algae) (Figure 2), both have good snoRNA coverage, which is based on both bioinformatics and intensive experimental study of green plant snoRNAs.

Finally, the Stramenopiles (Figure 2) have five completed and one draft genome project according to the GOLD database. Both the two main lineages of stramenopiles, Bacillariophyta and Oomycetes, have reasonable numbers of predicted snoRNAs based on homology to other lineages (9 and 75, respectively), though none has been experimentally validated. Whereas counts of Pfam domains and rRNAs indicate that the snoRNP machinery is present in all known taxa of Archaea and Eukaryota, surprisingly it seems to be absent from Oomycetes. However, this lack is likely to be due to the protein sequences not yet being included in the public sequence databases rather than *bona fide* loss of the snoRNP machinery.

Future directions for snoRNA research

Up to now, bioinformatics approaches for *de novo* prediction of snoRNAs have not been a great success. As shown by Figure 2, a homology search using experimentally verified snoRNAs, as performed by the Rfam database, has some success in identifying snoRNAs in taxonomic lineages where no experiments have yet been performed. But many of these predictions need further validation before they can be entirely trusted. Using additional information such as genomic context and target information could prove quite useful in this regard [11,12]. The growing host of orphan snoRNAs - that is, snoRNAs lacking a target-modification site - are especially interesting in that several lines of evidence hint at a possible regulatory role, as with human HBII-52 [3]. The snoRNA universe is thus likely to expand in function, phylogenetic diversity, and through the discovery of new snoRNAs. Fortunately, discovery has never been easier, thanks to the growing power of new sequencing technologies.

Additional file 1: Supplementary methods and results. It contains details of how the data for Figure 2 were collected, the full dataset summarized in Figure 2 in a tabular format, and an alignment of a *T. vaginalis* candidate snoRNA and the fungal homologs.

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