How to Have Dual Lives: Proteins that Bind DNA and RNA

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Editorial

Gene expression comprises transcription, splicing, mRNA export, mRNA stability and translation, which come together to control the abundance of any protein. These steps are not independent but are linked by proteins that act on multiple levels of gene regulation by interacting with the nucleic acids.

Traditionally, Nucleic Acid Binding Proteins (NABPs) were classified in several types according to the nature of the nucleic acid they bound, as well as to their sequence specificity. Thereby, it is frequent to find NABPs classified as DNA-binding proteins, DNA-binding proteins that specifically bind single-stranded DNA, RNA-binding proteins that bind to single- or double-stranded RNA, as well as proteins that bind specific or non-specifically to nucleic acids. On the other hand, proteins binding to DNA or RNA have been often considered and studied as separate functional classes of NABPs since Transcription Factors (TF) bind to genomic promoters and control target gene expression by activating or repressing RNA polymerase activity, and RNA binding proteins modulate protein expression by regulating the degradation, stability, and translation of target mRNAs. However, the consideration of DNA- and RNA-binding functions within proteins as discrete units has been rapidly changing in recent times. Several articles have demonstrated the dual DNA- and RNA-binding capability and cytoplasm-nucleus location of a growing group of NABPs. This duality creates an important additional level of gene regulation that serves to connect transcriptional and post-transcriptional control, and suggests a key role of these proteins in modulating gene expression, survival, and cellular homeostasis.

Dual NABPs may participate in gene expression by acting in different ways. TFs with dual binding abilities can establish an autoregulatory feedback loop in gene expression. TF with similar affinity for DNA and RNA may be kidnapped by nascent RNA molecules, thus reducing promoter occupancy and transcription of target genes. In addition to binding to the promoter of target genes, several NABPs may participate in several steps of the RNA metabolism (miRNA processing, mRNA stability, degradation, and translation). Further, some dual NABPs can bind and modulate secondary structures adopted by DNA and RNA during transcription and/or RNA processing, thus promoting or impeding transcription/translation processes.

The typical TF responsible for autoregulatory feedback loop in gene expression regulation is TFIIIA. TFIIIA is perhaps the first protein identified having dual nucleic acid binding capability. It was initially isolated from a 7S RNP complex as bound to 5S rRNA in Xenopus laevis oocyte cell extracts [1]. Later, it was also found as a transcription factor capable of activating 5S rRNA gene expression [2,3]. TFIIIA escorts 5S rRNA during nuclear export as 7S RNP complex and remains in the cytoplasm until the 5S rRNA is released for ribosome assembly. Release of the 5S rRNA allows TFIIIA to return to the nucleus, where it can activate the expression of additional 5S rRNA. As 5S rRNA transcripts are produced, TFIIIA progressively becomes sequestered in the cytoplasm thus leading to reduced transcription of the 5S rRNA gene.

TF also regulate genes post-transcriptionally through the regulation of miRNA biogenesis. Several TF have been shown to regulate primary miRNA processing by Drosha, a key step in the biogenesis of functional miRNAs. Often, identical sequences in DNA and RNA recruit NABP for transcriptional and posttranscriptional gene regulation.

During transcription, double-stranded DNA is open exposing single-stranded regions that can adopt secondary structures that may block or favor transcription initiation. In the past few decades a large number of scientific papers, both computational and experimental in vitro and in vivo; have demonstrated that G-rich DNA sequences can fold as intra-strand quadruple helix structures called G-quadruplexes or G4-DNA. G4-structures can be also formed in RNA molecules, usually in the 5'UTR. Several NABPs have been identified binding to G4-DNA and G4-RNA of the same gene and modulating their foldings. The presence of G4 might serve as a mark for recruiting specific TF or other RNA-binding proteins, thus promoting or avoiding the progress of transcription or translation as well.

Elucidating the mechanisms of action of these multifunctional proteins is difficult and could explain the slow progress in the field to date. The identification of the pool of dual NABPs in eukaryotic cells is not easy. For human cells, the QuickGO gene ontology database (European Bioinformatics Institute) yields low amount of proteins having experimental evidences for both RNA binding and DNA binding. Despite the PROTEOME database (BioBase) returns a higher amount of these proteins, direct experimental evidences regarding this dual activity is lacking for many of these entries. Recently, it was reported the ‘mRNA interactome’ of HeLa cells consisting of 860 mRNA-binding proteins [4]. Among them, approximately one half had been previously characterized as double-stranded DNA binding proteins [5], thus suggesting a relatively high representation in the human proteome. The number of dual NABPs would likely increase when including proteins present in other cell types. Certainly, the advent of new technologies ensures that more dual-NABPs and their mediated functions will be discovered in the coming years. This could be a focus of future research to enable a better understanding of how these highly abundant proteins fit into the broader theme of gene regulation.
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