

## Eukaryotic Transcriptional Regulation:

You are studying the regulation of a eukaryotic gene involved in memory called Absent Minded. Animals with mutations in this gene have severe short-term memory loss. You want to understand what controls the levels of Absent Minded gene expression to see if you can increase its expression in normal animals and therefore increase short-term memory. Using mutational mapping of the promoter, you identify two 10 bp regions of the promoter that when mutated strongly reduce expression of the Absent Minded promoter *in vivo*. You purify the proteins that bind these promoter elements and call the proteins ABR1 and ABR2.

To understand how these two proteins regulate Absent Minded gene expression, you do an *in vivo* experiment and an *in vitro* experiment

### *In vivo*

To be sure that these proteins are involved in activation of the Absent Minded promoter, you fuse the Absent Minded promoter (Upstream Activating Sequences and Promoter) to the LacZ gene and test the levels of expression in cells that have mutations that fail to produce the ABR1 and/or ABR2 proteins. You get the following results:

ABR1 Gene	ABR2 gene	Transcription Units
Wildtype	Wildtype	2000 Units
Mutant	Wildtype	50 Units
Wildtype	Mutant	200 Units
Mutant	Mutant	50 Units

- Based on these data, what can you conclude about the function of ABR1 and ABR2 at the Absent Minded promoter.

### *In vitro*

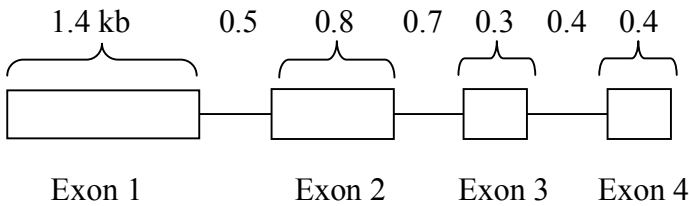
Your next step is to purify the ABR1 and ABR2 proteins. You test their ability to activate transcription *in vitro* by **first** adding the purified activators to a plasmid containing the Absent Minded Gene (Upstream Activating Sequences, Promoter, and Full Open Reading Frame). You then add purified RNA Polymerase II and auxiliary factors (TFIIA, TFIIB, TFIID, TFIIE, TFIIIF, and TFIIH). You get the following results:

ABR1 Protein	ABR2 Protein	Absent Minded Transcription (pmoles/minute)
-	-	500
+	-	500
-	+	3000
+	+	3000

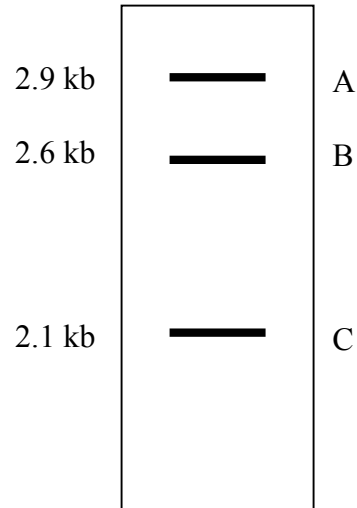
- Based on the differences between the reconstituted *in vitro* transcription experiments and the *in vivo* promoter fusion experiments, propose a function for the ABR1 regulator. **Hint:** Think about differences between prokaryotic and eukaryotic transcription and the kinds of functions eukaryotic transcriptional activators have. Explain your reasoning.

You are interested in studying the expression of the *mudblood* gene. The structure of the genomic coding region for the *mudblood* gene is shown below (Exons are boxes; Introns are lines). You isolate all the mRNAs from cells expressing this gene, run the mRNA on a gel, and then transfer the mRNAs to a membrane in order to do a northern blot. In order to study the mRNAs made by this gene, you make a radiolabeled DNA probe that corresponds to the DNA sequence found in exon 1. After hybridizing this probe to the membrane you made, you observe three different bands on the autoradiograph of your northern blot. Your results are shown on the right.

*Mudblood Genomic Region*



Northern Results with Exon 1 Probe:



- What process is occurring to produce this pattern?
- What can you conclude about the composition of bands A, B, and C that you observe on your northern blot?

We have seen in class that although RNA is single stranded, it has the ability to fold in to extremely complex structures. The following RNA sequence can fold up into a hairpin structure

5' CCACGUGACCUGCACGUCC 3'  
1 2 3

- Draw the structure of this hairpin.
- When the G residue marked 1 is mutated to a C, the RNA no longer folds into a stable hairpin. However, if a second mutation is introduced changing the C residue labeled 3 to a G, then a stable hairpin again forms. Briefly explain.
- If a deletion of the ACC sequence that begins at position 2 is made, then no folded structure is seen. Why do you think this occurs?