Phylogenetic and Structural Analysis of BC200 and Hominoidea Homologs

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I. INTRODUCTION

A. What is BC200?

Alzheimer's disease (AD) is a neurodegenerative disease which greatly affects the lives of those who are diagnosed and those who care for the diagnosed. In the United States, 6.2 million patients are estimated to be living with AD and it is the sixth leading cause of death for American adults [1]. In Canada, over 747,000 patients are living with AD, or another form of dementia [2]. Symptoms of AD range from memory loss and poor judgement in mild cases to the inability to communicate and seizures in severe cases [3].

AD involves multiple cell types and signaling pathways [4], as such, the collective knowledge of AD is spread across many different domains. This spread of knowledge means that fully understanding AD in humans is difficult, let alone understanding the disease in other species of the hominoidea superfamily. Finch and Austad argue that with our current understanding of AD, it is not possible to determine if AD is uniquely human [5]. As such, this paper explores one facet of AD pathogenesis, the long non-coding RNA (lncRNA) BC200. BC200 has been implicated in AD as it upregulates the expression of b-site APP-cleaving enzyme1 (BACE1) [4], [6]. The upregulation of BACE1 in turn leads to higher levels of beta-amyloid (A β) in the brain, thus, disrupting cell function [4], [6].

In hopes of better understanding how AD may affect other species in the hominoidea superfamily, specifically, the role that hominoidea homologs of BC200 play in AD, we answer the following research questions:

- RQ1. What does the phylogenetic tree of BC200 look like?
- RQ2. What structural differences exist between BC200 and its four most closely related hominoidea homologs?

The following paper is structured as follows: section II gives background into BC200 and the role it plays in AD, section III lays out the methods used for selecting BC200 homologs and performing both the phylogenetic structural analysis, section IV presents the results of our analysis, section V discusses the relevancy of the results, and section VI concludes the paper.

Brain Cytoplasmic 200 lncRNA RNA (BC200) is a 200 nucleotide long RNA transcript which is found mostly in the brain [7]. As a non-coding RNA, BC200 is not translated into protein but can be used as a potential therapeutic target and biomarker due to its regulatory role in biological processes involved in disease development [4], [8]. This lncRNA has recently been studied extensively because of its role in regulating translation and inhibiting its initiation, as well as its impacts in the pathogenesis of Alzheimer's disease and cancer [4], [7]. These [Derek: these being BC200?] non-coding RNAs are involved in translation control, thus, they impact the synthesis of dendritic proteins which facilitates long-term plastic changes at the synapse [8].

II. BACKGROUND

B. The Relation Between AD and BC200

Alzheimer's disease (AD) is a neurodegenerative disease resulting from synaptic plasticity failure in neurons [8]. It is a complex disease, meaning that it involves multiple cell types and signaling pathways [4].

AD is thought to occur due to the accumulation of two proteins in the brain. One of them is beta-amyloid $(A\beta)$ which accumulates in neurons, forms plaques, and disrupts cell functions. The other one is hyper-phosphorylated tau protein which in abnormal levels can form neurofibrillary tangles in neurons and block synaptic transmissions [4].

 $A\beta$, a cleavage product of the amyloid precursor protein (APP), is generated by b-site APP-cleaving enzyme1 (BACE1) and γ -secretase complex, and it strongly influences the pathogenesis of AD. Inhibition of BACE1 activity and the subsequent reduction in $A\beta$ levels may cure or prevent AD [4], [6].

BC200 facilitates AD pathogenesis by up-regulating $A\beta$ production through the modulation of BACE1 expression. The inhibition of BC200 significantly suppresses BACE1 expression, increases cell viability and reduces cell apoptosis in an AD model, and these effects are reversed by BC200 over-expression [4], [6].

Many researches have demonstrated the important role of BC200 in AD. El Mus et al. [8] show that there are steady decline in BC200 level from age 49 to 86, but, in AD brain its level was substantially higher. They also observe that BC200 expression is increased in brain areas that are involved in AD and it is parallel with severity of disease. Huanyen Li et al. [6] establish an AD cell model overexpressing A β 1-42 to observe the effects of BC200 on the cell viability and apoptosis and to investigate the associated underlying mechanisms. They observe that BC200 and BACE1 were increased upon treatment with A β 1-42, and inhibition of BC200 rescued this A β 1-42-mediated dysfunction, as indicated by the interaction of BC200 directly targeting BACE1. Moreover, inhibition of BC200 increased AD cell growth and reduced cells apoptosis. They demonstrate that BC200 is a potent positive regulator of BACE1 in AD cells and in conclusion, lncRNA BC200 facilitates AD pathogenesis by up-regulating $A\beta$ through BACE1.

III. MATERIALS AND METHODS

A. Selection of lncRNAs

The lncRNA BC200 was selected for phylogenetic analysis due to the role it plays in AD as discussed in section II. The homologs of BC200 were selected as a result of an NCBI Blast [9]. Specifically, Megablast [10] with default parameters was used as it is able to compare closely related sequences [11]. From the Blast results, sequences which are known BC200 homologs as indicated by the inclusion of BC200 in their name were chosen. Table I outlines the sequence name including the organism and the accession number of each of the chosen sequences.

B. Phylogenetic Analysis

TODO

C. Structural Analysis

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IV. RESULTS

A. Phylogenetic Tree of BC200

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B. Structural difference between BC200 and its homologs

Brain cytoplasmic 200 long non-coding RNA (or BC200 lncRNA) is a 200 nucleotide RNA transcript that is found predominantly in the brain. It's primary function is regulating translation by inhibiting its initiation. It's role in AD is not fully understood, but research shows that lncRNA BC200 facilitates AD pathogenesis by upregulating AB through BACE1. Pathologically, AD is characterized by an imbalance

in the production and clearance of amyloid-beta in the brain leading to plaque formation [6]. The BC200 structure consist of three main parts: A-rich domain, Alu domain and unique domain [12].

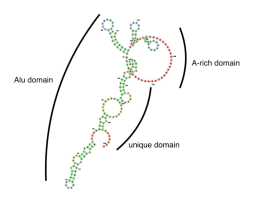


Fig. 1: BC200 RNA Secondary Structure

As we can see from the RNA secondary structures, all of the homologs have a nearly simialr A-rich domain, but on the other hand the the Alu domain is different between the homologs and BC200. The Alu domain of the mammalian signal recognition particle (SRP) comprises the heterodimer of proteins SRP9 and SRP14 bound to the 5 and 3 terminal sequences of SRP RNA [13]. So, their function in brain should be very different from one another, even though there may be high similarity between their sequences. Figures 2, 3, and 4 show the RNA secondary structure of the three most closely related BC200 homologs from great apes (hominidae). Additionally, we have chosen to depict the RNA secondary structure of the Hylobates lar BC200 homolog in Figure 5. While Hylobates lar is not a great ape, it is still part of the hominoidea superfamily, and Phylogenetic analysis revealed that it was closely related to BC200.

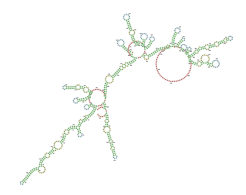


Fig. 2: BC200 RNA Secondary Structure in Gorilla

TABLE I: Sequence information for BC200 and hominoidea homologs

Sequence Name	Accession Number
Homo sapiens brain cytoplasmic RNA 1 (BCYRN1), long non-coding RNA	NR_001568.1
Pongo pygmaeus BC200 alpha scRNA gene, complete sequence	AF067778.1
Pan paniscus BC200 alpha scRNA gene, complete sequence	AF067778.1
Gorilla gorilla BC200 alpha scRNA gene, complete sequence	AF067779.1
Macaca mulatta BC200 alpha scRNA gene, complete sequence	AF067784.1
Hylobates lar BC200 alpha scRNA gene, complete sequence	AF067781.1
Papio hamadryas BC200 alpha scRNA gene, complete sequence	AF067782.1

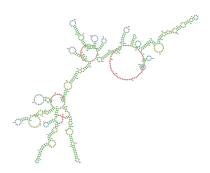


Fig. 3: BC200 RNA Secondary Structure in Pan

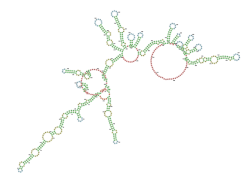


Fig. 4: BC200 RNA Secondary Structure in Pango

C. Comparing BC200 sequence with its homologs

One of the best ways for finding the conserved portions of a gene during evolution, is sequence alignment. Here, we aligned BC200 and the three most related homologs with each other to find the conserved parts of the sequence.

Here are the alignment results:

As it is obvious from the results, most parts of the sequence are conserved. Which shows that there should be a high similarity between BC200 RNA in human body and in other species. So, the following result does not support our hypothesis, and they suggest that Alzheimers can be shared between human and great apes.



Fig. 5: BC200 RNA Secondary Structure in Hylobates lar

		00002.12:47335315-47335514 Homo sapiens chromosome 2, Primary Assembly Query ID: lcl Query_43861 Length: 200	
>AF067778.1 Fan paniscus BC200 alpha scRNA gene, complete sequence Sequence ID: Query 43864 Length: 713 Range 1: 431 to 635			
Score:340 bits(184), Expect:le-97, Identities:199/205(97%), Gaps:5/205(2%), Strand: Plus/Plus			
Query 60	1	$\tt GGCCGGGCGCGGTGGCTCACGCCTGTAATCCCAGCTCTCAGGGAGGCTAAGAGGCGGGAG$	
60			
Sbjct 490	431	GGCCGGGCGCGGTGGCTCACGCCTGTAATCCCAGCTCTCAGGGAGGCTAAGAGGCGGGAG	
Query	61	GATAGCTTGAGCCCAGGAGTTCGAGACCTGCCTGGGCAATATAGCGAGACCCCGTTCTCC	
120			
Sbjct 550	491	GATAGCTTGAGCCCAGGAGTTCGAGACCTGCCTGGGCAATATAGCGAGACCCCGTTCTCC	
Query 175	121	AGaaaaaggaaaaaaaacaaaagacaaaaaaaaTAAGCGTAACTTCCCTC	
	551	AGAAAAA GGAAAAAAAAAAAAAAAAAAAAAAAAAAAA	
Sbjct 610	551	AUGARARAN UGARARARARARARARARARARARARARARARARARARAR	
Query	176	AAAGCAACAAccccccccCTTT 200	
Sbjct	611	AAAGCAACAACCCCCCCCCCTTT 635	

Fig. 6: BC200 Sequence Alignment with BC200 in pan

V. DISCUSSION

TODO

VI. CONCLUSION

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Fig. 7: BC200 Sequence Alignment with BC200 in Pango

Fig. 8: BC200 Sequence Alignment with BC200 in Gorilla

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