MicroRNA in Situ Hybridization in the Human Entorhinal and Transentorhinal Cortex

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MicroRNA in situ hybridization in the human entorhinal and transentorhinal cortex

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MicroRNAs (miRNAs) play key roles in gene expression regulation in both healthy and disease brains. To better understand those roles, it is necessary to characterize the miRNAs that are expressed in particular cell types under a range of conditions. In situ hybridization (ISH) can demonstrate cell- and lamina-specific patterns of miRNA expression that would be lost in tissue-level expression profiling. In the present study, ISH was performed with special focus on the human entorhinal cortex (EC) and transentorhinal cortex (TEC). The TEC is the area of the cerebral cortex that first develops neurofibrillary tangles in Alzheimer’s disease (AD). However, the reason for TEC’s special vulnerability to AD-type pathology is unknown. MiRNA ISH was performed on three human brains with well-characterized clinical and pathological parameters. Looked nucleic acid ISH probes were used referent to miR-107, miR-124, miR-125b, and miR-320. In order to correlate the ISH data with AD pathology, the ISH staining was compared with near-adjacent slides processed using Thioflavine stains. Not all neurons or cortical lamina stain with equal intensity for individual miRNAs. As with other areas of brain, the TEC and EC have characteristic miRNA expression patterns. MiRNA ISH is among the first methods to show special staining characteristics of cells and laminae of the human TEC.

Keywords: hippocampus, neurodegeneration, Alzheimer, brain, ncRNA, RNA, neuron, NFT

INTRODUCTION

Alzheimer’s disease (AD) is a prevalent neurodegenerative disease that culminates in severe deficits in cognition and autonomy. By definition, brains afflicted by AD contain two different neuropathological hallmarks — neurofibrillary tangles (NFTs) and neuritic amyloid plaques (NPs) The National Institute on Aging, and Reagan Institute Working Group on Diagnostic Criteria for the Neuropathological Assessment of Alzheimer’s Disease (1997). NFTs are ‘inclusion bodies’, composed of insoluble tau protein polymers that coalesce within neurons. NPs consists a roughly-spherical extracellular component that includes fibrillar polymers indistinguishable from those in NFTs.

Neuroanatomically, AD pathology manifests in a complex but well-characterized spatiotemporal sequence (Braak and Braak, 1991; Braak et al., 1993). Most clinico-pathological correlation studies indicate that cortical NFT density, assessed by Braak staging (Braak et al., 1993) or other means, is the parameter best correlated with the severity of AD cognitive impairment (Arriagada et al., 1992; Nelson et al., 2007b, 2008a,b; Sonnen et al., 2007). In the first stages of the disease, NFTs are observed in medial temporal lobe structures (Braak and Braak, 1991).

The specific cerebral cortical subfield with earliest NFT formation in AD is the transentorhinal cortex (TEC) (Braak and Braak, 1992). The TEC usually occupies the medial bank of the perirhinal collateral sulcus, comprising ~2–10 mm of the inexacty defined and phylogenetically variable Brodmann Area 35 (Schmidt et al., 1993; Taylor and Probst, 2008). As its name implies, the TEC constitutes a transitional zone between the entorhinal cortex (EC) and the more laterally situated six-layered cerebral isocortex (Braak and Braak, 1985). The TEC thus “can be considered a periallocortical field bordering a proisocortical field of the adjacent temporal cortex, that corresponds to the perirhinal region”, according to Heiko Braak and colleagues (Schmidt et al., 1993). NFTs are observed in the TEC long before clinical manifestations of AD: in a large autopsy series, scant numbers of NFTs were detected in the TEC of approximately one-fifth of persons dying in their thirties and over one-third of persons in their forties (Del Tredici and Braak, 2008).

During the progression toward end-stage AD, NFTs and/or cell death claim the large majority of neurons in some laminae of the TEC and nearby EC (Braak and Braak, 1985; Gomez-Isla et al., 1996; Garcia-Sierra et al., 2000; Hof et al., 2003). NFTs are also observed in the TEC and EC in neurodegenerative diseases other than AD: Parkinson’s disease, argyrophilic grain disease, and Huntington’s disease, for example (Braak and Braak, 1992; Ulrich et al., 1992; Braak et al., 2000; Nelson et al., 2009a). Unfortunately, the remarkable predisposition of TEC and EC neurons to develop pathological changes has not been adequately explained. Nor are there many reported markers providing clues about TEC neurochemistry.

In the present study, in situ hybridization (ISH) was performed in order to characterize the distribution of some neuronally-expressed microRNAs (miRNAs) in the human TEC and surrounding brain structures. MiRNAs are short (~22 nucleotide) RNA molecules that play fundamental roles in gene expression regulation in all known plants and animals. In particular, miRNAs are known
to serve key functions in neurodevelopment, synaptic plasticity, and neuroprotection (Kosik and Krichevsky, 2005; Cuellar et al., 2008; Smalheiser and Lugli, 2009). MiRNAs may have potenti-
ated mammalian brain evolution by amplifying the complexity of nervous system gene expression regulation (Nelson and Keller, 2007; Heimberg et al., 2008). On the other hand, miRNAs also contribute to human illnesses, particularly in the pathogenesis of human neurodegenerative disease (Nelson et al., 2008b; Hebert and De Strooper, 2009). ISH shows important cerebral cortical lamina-specific patterns of miRNA expression that would be lost on most tissue-level expression studies (Mellios et al., 2008; Nelson and Wilfred, 2009), and these lamina-specific miRNA expression patterns could be relevant to AD (Wang et al., 2008).

We chose to study four miRNAs that are expressed in human brain:

1. miR-107, which we have shown may be relevant to AD pathogenesis and traumatic brain injury, and which may be involved in metabolic regulation (Wilfred et al., 2007; Wang et al., 2008; Redell et al., 2009; Tang et al., 2009).

2. miR-124, which is highly enriched in neurons and plays many important roles in neuronal gene expression regulation (Smirnova et al., 2005; Krichevsky et al., 2006; Makeyev et al., 2007; Tang et al., 2007).

3. miR-125b, which is expressed in many different cell types including the mammalian brain and which has been proposed to play a number of complex nervous system roles (Smirnova et al., 2005; Lukiw and Pogue, 2007; Ferretti et al., 2008; Le et al., 2009a,b).

4. miR-320, which is highly expressed in neurons and glial cells and which is dysregulated in prion disease (Nelson et al., 2007a; Saba et al., 2008).

MATERIALS AND METHODS

Brain tissue was obtained from University of Kentucky ADC Brain bank using appropriate IRB protocols. Details of subject recruitment, autopsies, and other analyses using the University of Kentucky ADC autopsy series are described elsewhere (Nelson et al., 2007b, 2008a). Criteria for inclusion in this study included post-mortem intervals (PMIs) under 5 h. Brain sections from two individuals without antemortem cognitive decline were included (Cases 1 and 2), and a third person with early AD (Case 3). Demographic and pathological parameters of each of the three cases used for this study are shown in Table 1.

Table 1 | Information about cases used in the current study including demographic, cognitive testing, post-mortem interval, Consortium to Establish A Registry for Alzheimer’s Disease (CERAD) neuritic plaque, Braak neurofibrillary stages, and apolipoprotein E (ApoE) alleles.

| Case | Sex | Age at death | Final MMSE score | Months since final MMSE | Post-mortem interval (hrs) | CERAD score | Braak stage | ApoE |
|------|-----|-------------|------------------|------------------------|---------------------------|-------------|-------------|------|
| Case 1 | F   | 79          | 29               | 9                      | 1.75                      | No          | 0–1         | 3/3  |
| (Non-demented) |   |             |                  |                        |                           |             |             |      |
| Case 2 | F   | 94          | 27               | 5                      | 2.33                      | No          | II          | 3/3  |
| (Non-demented) |   |             |                  |                        |                           |             |             |      |
| Case 3 (AD) | M   | 76          | 27               | 32                     | 4.25                      | Definite    | VI          | 3/4  |

Human brain ISH methods have been published (Nelson et al., 2006; Wang et al., 2008; Nelson and Wilfred, 2009). Very briefly, post-mortem human brain sample was obtained via autopsy within 5 h of death. Tissue portions that included the EC were used. Brain tissues were fixed in 4% paraformaldehyde overnight at 4°C and then immersed in 20% sucrose (4°C) for an additional 24–48 h. Tissue was cut to 25 microns on a freezing microtome and mounted onto premarked Superfrost® Plus slides. Cut tissue sections were allowed to air-dry for 30 min. The slides were then transferred to a ~80°C freezer, until subsequent processing as described. Digoxigenin-labeled locked nucleic acid probes (Exiqon, Woburn MA) were used and their presence visualized via anti-digoxigenin immunohistochecmy using protocols described in detail previously (Wang et al., 2008; Nelson and Wilfred, 2009).

Histological stains were performed on near-serial sections that were fixed, cut, mounted, and frozen along with the sections used for ISH. Thioflavine S (Polysciences, Inc., Warrington, Pennsylvania) was used as a 1% aqueous solution followed by differentiation in two changes of 80% ethanol. Nissl staining was accomplished using 0.1% Cresyl violet solution that was filtered immediately before use. After staining for 5 min, sections were differentiated in 95% ethanol and cleared in xylenes.

RESULTS

Sections of human TEC and EC were evaluated using miRNA ISH and several histological stains. Photomicrographs from each of the three cases are presented in Figures 1–3. These show ISH results for miR-107, miR-124, miR-125b, and miR-320 in the TEC and nearby structures in correlation to AD pathology (Thioflavine S stained NFTs and NPs).

Since miR-320 and miR-124 showed distinctive cortical laminar staining in the TEC, a separate panel (Figure 4) shows the ISH results for near-serial section using miR-320 and miR-124 ISH and Thioflavine S from Case 3. Note that the Thioflavine S-stained NFT-bearing neurons are present in a band of cells that are relatively lacking in ISH stain for both miR-124 and miR-320. For both miR-124 and miR-320, there is an immediately more superficial band of cells that are labeled. As can be seen in Figures 1–3, the staining pattern in and near the TEC is relatively consistent with regard to this staining pattern of cells – the layer of miR-124 cells appears particularly characteristic. MiR-107 stains less strongly than the other probes. As expected, miR-125b, and to a lesser degree miR-107 and miR-320, appear to stain glial cells in addition to
FIGURE 1 | In situ hybridization, Nissl, and Thioflavine S stains for Case 1. (A) Nissl stain (B) miR-107 ISH (C) miR-125b ISH (D) miR-320 ISH (E) miR-124 ISH, and (F) Thioflavine S (amyloid plaques and neurofibrillary tangles are fluorescent.) In this case, there were no amyloid plaques and a few scattered neurofibrillary tangles, which are indicated with arrows in the transentorhinal cortex. Scale bar = 1000 microns. D = dorsal, M = medial, V = ventral, L = lateral.

FIGURE 2 | In situ hybridization, Nissl, and Thioflavine S stains for Case 2. (A) Nissl stain (B) miR-107 ISH (C) miR-125b ISH (D) miR-320 ISH (E) miR-124 ISH, and (F) Thioflavine S (amyloid plaques and neurofibrillary tangles are fluorescent.) TEC indicated with arrows. Scale bar = 1200 microns. D = dorsal, M = medial, V = ventral, L = lateral.
neurons. In some of the sections there was artifactual staining in the white matter that was trimmed out of the photomicrographs. To see what the staining in the white matter looks like, please see Supplementary Material.

Discussion

ISH using a set of probes against brain-enriched miRNAs was used to assess cellular miRNA expression in the human TEC and surrounding structures. These data have both technical and theoretical implications. From a technical standpoint, the present study further underscores the importance of ISH as a technique to complement tissue-level miRNA expression profiling. The pattern of miRNA expression also helps to refine the expectations for miRNA functions in the brain, in both normal and disease conditions. The pattern of ISH labeling in the human cerebral cortex affirms that defining individual miRNAs as “neuronal” or “non-neuronal” is overly simplistic, because different populations of neurons – even within a tiny cell layer – can have distinct miRNA expression profiles.

There are some limitations to the current study. ISH is a relatively low-throughput technique and thus we only were able to thoroughly evaluate the results using a handful of miRNA probes, and brain sections from only three individuals’ brains (all of these patients were elderly Caucasians). Although the results were consistent among the cases evaluated, it is possible that study of a larger population would result in greater variability or even completely disparate results. Also, it has been shown that post-mortem degradation of miRNAs in the human brain can be rapid and can affect different miRNAs at different rates (Sethi and Lukiw, 2009). In the current study we used brains that had relatively short PMIs (less than 5 h). However, using cases with even shorter PMIs may have revealed a different staining pattern. Finally, the technique that we used employs both ISH and...
immunohistochemistry (anti-digoxigenin antibody is used to detect the digoxigenin-labeled RNA probe). Like the great majority of histochemical staining methods, ISH is not a rigorously quantitative technique for evaluating gene expression. This is because of the many variables in tissue processing that are impossible to control between cases and even between different sections of the same case. As such it is more appropriate to evaluate critically the staining pattern, rather than the more detailed characteristics of staining intensity in any given section. We also are dissatisfied with employing near-serial sections to define correlative staining patterns. Hence we are investigating the use of double-label methods to enable, in the future, more definite determination of human brain miRNA co-expression patterns.

Despite the limitations inherent to this type of study, the ISH technique reveal consistent staining patterns in the human cerebral cortex. The TEC itself is a small (<1 cm) but highly intriguing cortical subfield, being most developed in the human relative even to other primate species (Braak et al., 2000; Taylor and Probst, 2008). Perhaps most interesting is the cell population that seems to “dive” from the superficial layer II neurons of the EC, through the entire depth of layer III in the TEC, to approximate layer V neurons in the “temporal proneocortex” (Braak and Braak, 1985, 1992). These are apparently the neurons most vulnerable to NFT development in the human cerebral cortex. These particular neurons show an apparent lack of high amounts of both miR-124 and of miR-320. The impact of these expression patterns are currently not understood. They may relate in some way with the remarkable dendritic plasticity shown by TEC and EC neurons (Arendt et al., 1998). However, neurons in the nearby pre-α (layer II) of the EC proper, which is also affected relatively early in AD, express high amounts of both miR-320 and miR-124. There are of course many other miRNAs with expression patterns overlaid on those of miR-107, miR-125b, miR-124, and miR-320. For all of these, the lamina-specific expression characteristics would not be possible to evaluate using most tissue-level expression profiling platforms.

In conclusion, we have demonstrated novel miRNA expression patterns in the human TEC and EC. Neurodegenerative diseases tend to affect these cell populations for reasons that currently are poorly understood. MiRNAs are important components of a neuron's gene expression repertoire. Each neuronal subpopulation may express a distinct fraction of the uniquely-human miRNome. Thus,
future directions for study of miRNAs should include more human brain ISH. With the help of these “descriptive” studies we can better understand the relevant mechanisms of miRNAs in health and in disease states.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at http://www.frontiersin.org/humanneuroscience/paper/10.3389/neo.009/0072010/

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