Alzheimer’s disease (AD) and age-related macular degeneration (AMD) are complex and progressive inflammatory degenerations of the human neocortex and retina. Recent molecular, genetic and epigenetic evidence indicate that at least 4 micro RNAs (miRNAs) - including the NF-κB-regulated miRNA-9, miRNA-125b, miRNA-146a and miRNA-155 - are progressively up-regulated in both AD and AMD. This quartet of up-regulated miRNAs in turn down-regulate a small brain- and retinal-cell-relevant family of target mRNAs, including that encoding complement factor H (CFH), a major negative regulator of the innate immune and inflammatory response. Together miRNA-146a and miRNA-155 recognize an overlapping miRNA regulatory control (MiRC) region in the CFH 3'-untranslated region (3'-UTR; 5'-TTTAGTATTAA-3') to which either of these miRNAs may interact. Progressive, pathogenic increases in specific miRNA binding to the entire 232 nucleotide CFH 3'-UTR appears to be a major regulator of CFH expression down-regulation, and the inflammatory pathology that characterizes both AMD and AD. The data presented in this report provides evidence that up-regulation of brain- and retinal- abundant miRNAs, including miRNA-9, miRNA-125b, miRNA-146a and miRNA-155, are common to the pathogenetic mechanism of CFH deficiency that drives inflammatory neurodegeneration, and for the first time indicates multiple, independent miRNA-mediated regulation of the CFH mRNA 3'-UTR.

Keywords: Brain, complement factor H, evolution, miRNA regulatory control (MiRC) region, micro RNA, miRNA-9, miRNA-125b, miRNA-146a, miRNA-155, retina, small non-coding RNA

Complement factor H (CFH) in AD and AMD

Alzheimer’s disease (AD) and age-related macular degeneration (AMD) are multifactorial, late onset, progressive neurodegenerative diseases of the human brain and retina, and represent the leading cause of cognitive and visual impairment in aging populations of the industrialized world [1-6]. Despite considerable research efforts, the causes of AD and AMD are not well understood, and currently there are no effective treatments for these progressive degenerations of human cognition and vision. There are many shared pathological characteristics of AD and AMD, including (a) the appearance of dense, insoluble, polymorphic amyloid beta (Aβ) peptide-, oxidized lipid- and complement factor-enriched pathological lesions known as senile plaques (SP) in AD and drusen in AMD, (b) elevated cholesterol, 24S-hydroxycholesterol and other oxidized lipid metabolites, (c) amyloidogenesis, (d) heightened inflammatory and pro-apoptotic signaling, and (e) decreases and/or dysfunction in complement factor H (CFH), a major regulator of the innate immune and inflammatory response [8-30]. For instance, it has recently been shown that in AD, amyloidogenic, pro-inflammatory and pro-apoptotic gene expression spreads from the limbic system into the primary visual sensory cortex (Brodmann area A17) as AD advances, rostrally through the thalamus/superior colliculus to the retina, along the entire anteroposterior axis of the retinal-primary visual cortex pathway. This spreading correlates with progressive visual disturbances reported in late-stage AD.
miRNAs of the aging human brain and retina

patients [7-9]. Further, transgenic models of AD, such as the 5xFAD model (that contains 5 familial amyloid transgenes regulated by neural-specific elements of the mouse Thy1 promoter to drive human amyloid over-expression) show age-related deposition of amyloid peptides, innate immune and inflammatory defects and CFH-associated pathology in both the brain and retina [9-13]. As discussed more fully below, at the molecular-genetic level pathogenic up-regulation of miRNA-9, miRNA-125b, miRNA-146a and miRNA-155 appear to be involved in a comparable progressive, CFH-mediated inflammatory degeneration that is characteristic of both AD and AMD.

Consistent observations concerning changes in CFH signaling have been described in both AD and AMD [1-12, 16-25]. CFH (also known as AC3b1RNA, adrenomedullin binding protein-1, AMBP-1, AM binding protein-1factor H, β1H globulin, C3b inactivator accelerator, H factor, HF, H factor-1, HF1) is an important member of the regulator of complement activation (RCA) group of proteins encoded within the RCA gene locus spanning chromosome 1q21-1q32. In the human brain and retina CFH is transcribed from a single copy gene that generates one major ~4100 nucleotide mRNA transcript [23, 25; ~95% of the CFH mRNA signal in a single band on Northern gels; Genbank BC142699; unpublished observations]. A major amount of CFH is also normally generated in, and secreted by, the liver, and CFH reaches blood plasma concentrations of 500-800 ug/ml in the systemic circulation [3, 15, 25]. Hence CFH is the second most abundant serum plasma protein, after human serum albumen, and normally performs a systemic sentinel function against unscheduled or spontaneous activation of the innate immune system [18-20]. Systemic CFH deficits are conducive to excessive and pathogenic complement pathway activation associated with increased complement activity on otherwise healthy host cells, autoimmunity, host tissue damage and a sustained and often chronic inflammatory response [7, 9, 10, 18-20]. At the current time it is not clear if CFH, a soluble, hydrophilic 155 kDa glycoprotein, is permeable to the blood-brain barrier; such large glycosylated serum proteins are usually prevented from access to CNS compartments, and the brain and retina may have an independent CFH supply secreted by neurons, astroglia, microglial and/or endothelial cells [2, 12, 19]. Otherwise, CFH may contribute to SP and drusen formation via a diseased or leaky neocortical or retinal vasculature [47-50]. CFH expression is significantly down-regulated in the degenerating brain and retina, suggesting that insufficient quantities of this RCA regulator lead to excessive activation of the innate immune response and pro-inflammatory signaling [5-8, 18, 20-25, 32, 79]. In both of these tissues CFH functions as a co-factor in the inactivation of its C3b ligand by factor 1 in the alternative complement pathway [2-5]. As a molecular constituent of the SP and drusen that accumulate in AD and in AMD, CFH co-localizes with its ligand C3b in sub-structural spherules within drusen that also contain Aβ42 peptide and additional related toxic βAPP-derived fragments [14-17, 31, 32]. Extensive inverse abundance interrelationships between high Aβ42 levels and low CFH levels have been described in degenerative disorders of retinal and brain cells [9, 16]. Moreover, the significant statistical relationship between the CFH Y402H “risk” variant and AMD further suggests that either a defective ‘mutant’ CFH, or low CFH abundance may have analogous pro-inflammatory effects in activation of the innate immune response [2-5, 20, 33].

Pathogenic, pro-inflammatory micro RNAs (miRNAs) in the brain and retina

Micro RNAs (miRNAs) constitute a family of ~22 nucleotide (nt), non-coding, single-stranded labile RNAs [14, 21-25, 34, 42, 55, 60, 63, 76-79] The major mode of biological action of miRNA is to bind to complementary RNA sequences in the 3′ un-translated region (3′-UTR) of messenger RNA (mRNA), and thereby act as a repressor of that mRNA’s expression. It is now widely accepted that up-regulated miRNAs predominantly act to decrease their target mRNA levels, and hence down-regulate genetic information encoded by that target mRNA [14, 21-25, 34]. Of the approximately 1898 human miRNAs currently identified, less than 10^2 miRNAs are abundantly expressed in brain and retina [40-42; unpublished]. Interestingly, while up-regulated miRNAs have the potential to reduce the transcript levels of their target miRNAs on a genome-wide scale, in disease processes only certain miRNAs are preferentially affected. Figure 1A and 1B shows expression of a family of pathogenic miRNAs significantly up-regulated in aging human neocortex and in AMD retina. As discussed more fully below, these results were
miRNAs of the aging human brain and retina

Independent confirmation using RT-PCR and/or Northern dot-blot techniques (Figure 1C) [5-8, 24, 25]. Common to aged degenerating brain and retina are up-regulated miRNA-9, miRNA-125b, miRNA-146a and miRNA-155, and up-regulation these 4 miRNAs has been shown to be involved with an up-regulation of inflammatory signaling in large part due to their specific targeting of the CFH mRNA 3'UTR and down-regulation of CFH expression [7, 9, 16, 21-25, 31, 42, 45, 68, 76]. Interestingly, certain of these up-regulated miRNAs also target other pathology-relevant miRNAs including tetraspanin 12 (TSPAN12; also known as NET-2 or trans-membrane 4 superfamily member 12), a membrane spanning protein that associates with ADAM10, a membrane secretaise responsible for the cleavage of beta amyloid precursor protein (βAPP) into neurotrophic sAPPox [25, 30, 31, 37, 38]. Hence miRNA-125b-, miRNA-146a- and miRNA-155-mediated down-regulation of TSPAN12 impairs ADAM10 activity, shunting βAPP processing into amyloidogenic (Aβ42 generating) pathways [9, 10, 25, 31, 37, 38]. As further discussed below, several recently published reports indicate a significant up-regulation of miRNA-146a and CFH and TSPAN12 down-regulation in human primary cells and tissues [16, 25, 31].

Mathematical considerations of brain and retinal miRNA complexity

As previously mentioned, degenerating human neocortical and retinal cells display an intriguing overlap in the speciation and complexity of a small family of specific miRNA’s expressed within those tissues (Figure 1). Using the most advanced MRA-1001 miRNA microfluidic chip analytical platform currently available (LC Sciences, Houston TX) to date only 1898 miRNAs have been detected in the cells of all human tissues. However, depending on cell type, nervous tissues only contain a small fraction of all known miRNAs and some are far more abundant than others [39, 40-42]. Bioinformatics and sequence analysis further indicates that a 22 nucleotide single stranded RNA composed of
4 different ribonucleotides can have over $10^{13}$ possible sequence combinations, so the fact that there typically only about $2 \times 10^9$ abundant miRNAs in all types of human cells suggests a very high developmental and evolutionary selection pressure to utilize only specific miRNA oligonucleotide sequences that will yield biologically useful miRNA-mRNA interactions. Put another way, miRNA-mRNA pairings have been strongly selected over evolution to maximize their efficiency in regulating brain and retinal gene expression as well as expression in other tissues. Further, miRNAs are highly developmental stage-, tissue- and cell-specific, even in adjacent cell types, and in human brain neocortical and retinal cells high abundance miRNAs number probably less than $10^2$ individual species [31, 39-42, 45; unpublished observations]. The fact that out of the top 12 changed miRNAs in degenerating neocortex and retina that 5 are common to each tissue was the impetus for these studies, which collectively suggest that certain of these miRNAs might be significant in genetic regulatory mechanisms common to inflammatory neurodegenerative mechanisms characteristic of both AD and AMD.

**AD and AMD; pathological similarities - senile plaques (SP) and drusen**

AD and AMD are characterized pathologically by the progressive appearance within the brain parenchyma and retinal macular region of pro-inflammatory, highly insoluble, amyloid beta (Aβ) peptide enriched pathological lesions known, respectively, as neocortical senile plaques (SP) and retinal drusen. Although there are several other disease-related components and lesions associated with AD and AMD, SP and drusen represent major pathological hallmarks of these age-related degenerative disorders of the neocortex and retina [44-50, 75]. SP are roughly spherical 20-30 μm in diameter, non-crystalline, argyrophilic globular masses composed of a diffuse-to-dense core of various amyloid peptides and other oxidized protein and lipid components; drusen are approximately 5-15 μm diameter, circular, yellow-white lesions located throughout the fundus of the retina but more so in the macular region, around the optic disc or the periphery [46-50]. The biological environment in which these lesions initiate and grow may be their only major difference in establishing their final size and ultimate morphology. In AD SP usually form exterior to the neuron in an ‘unrestricted’ brain parenchyma free of connective tissue, allowing these lesions to form into globular aggregations considerably larger than drusen [47]. In AMD drusen form in a far more restricted and delineated region usually between the basement membrane of the retinal epithelial (RPE) cell layer and Bruch’s membrane [47-50]. In the study of neuropathology and pathobiology, strongly similar constituents of both SP and drusen are apparent; SP and drusen each contain upwards of about 100 different end-stage protein, lipid and other constituents including toxic trace metals [32, 46-51]. Major common components of SP and drusen include beta-amyloid precursor protein (βAPP), the α-, β- and γ-secretases which cleave βAPP to generate a wide range of βAPP catabolites including self-aggregating Aβ peptides 37-42 amino acids in length (Aβ37-Aβ42) [17, 32]. Other important, common constituents of SP and drusen include α- and β-crystallin, serum related molecules such as amyloid P, clusterin, collagen, complement proteins such as C3, C5, complement factor H (CFH) and the C3b and C5b-9 complex, apolipoproteins B and E, esterified and non-esterified cholesterol, oxidized lipids and phospholipids, 24S-hydroxycholesterol, neutral vitronectin, neprilysin, vasogenic factors such as VEGF, vimentin, and the secretase-accessory proteins which further process βAPP into multiple amyloid peptides [47, 48]. Interestingly, the major neurotrophic βAPP metabolite sAPPα attains some of its highest concentrations in the retina, the RPE and particularly in the vitreous humor located just adjacent to the retina [9, 30; unpublished observations]. SP and drusen also contain trace metals such as aluminum and zinc which are thought to assist in the aggregation of the constituents contained within these insoluble lesions [46-48; unpublished observations]. In both cases, waste products from neurons, astroglia, photoreceptors, RPE and retinal ganglion or blood plasma components via the highly vascularized neocortex and retinal choriocapillaris provide a steady source of extracellular serum-derived material for SP and drusen formation that over time support their enlargement. Indeed oxidative and immune-mediated modifications appear to further support their progressive aggregation, growth and expansion [46-50]. Interestingly, both highly insoluble pathological lesions are progressively deposited as a function of age, and many of their components represent ‘end-stage’ or metabolic waste.
products highly resistant to degradation by the endoplasmic-reticulum-associated protein degradation (ERAD) and ubiquitin-proteasome systems [46-52].

AD and AMD; similarities in specifically up-regulated miRNAs

Tissue- and cell-specific families and sub-families of miRNAs are emerging as extremely important post-transcriptional regulators of messenger RNA (mRNA) complexity and in shaping the transcriptome of a cell in development, health, aging and disease [16, 23-26, 34-36]. The human brain and retina rely on a specific subset of these miRNAs to shape their gene expression patterns and it is remarkable that the total number of abundant miRNAs they utilize to do this are probably less than 10^2 [53, 54; unpublished observations]. Recently, using miRNA arrays, DNA arrays, Northern micro-dot blots, RT-PCR, Western analysis, ELISA-based assays, bioinformatics and miRNA-mRNA pairing algorithms it has been shown that diseased brain and retina share a number of abundant pathology-relevant miRNAs, and related families of expressed pro-inflammatory and amyloidogenic genes are similarly targeted (Figure 1) [16, 23, 38-42]. As previously mentioned these include at least four inducible miRNAs known to be up-regulated by NF-κB that include miRNA-9, miRNA-125b, miRNA-146a and miRNA-155 (Figure 1) [22-25, 31]. These up-regulated miRNAs in turn appear to down-regulate multiple target mRNAs and thus regulate a family of potentially pathogenic genes including 15-lipoxygenase (15-LOX), synapsin-2 (SYN-2), tetraspanin-12 (TSPAN12) and complement factor H (CFH) [7, 9, 12, 14, 16, 21-25]. These data further indicate that as few as four abundant NF-κB-mediated miRNAs have tremendous potential to contribute to many interrelated aspects of brain and retinal cell function and dysfunction, including non-homeostatic neurotrophic support (15-LOX), synaptogenesis (SYN-2), amyloidogenesis (TSPAN12) and innate immune signaling and inflammation (CFH) [21-25, 28, 30, 31].

miRNA control of complement factor H (CFH) expression

Importantly, up-regulated miRNA-125b, miRNA-146a, miRNA-155 and perhaps miRNA-9, and subsequent down-regulation of CFH gene expression may in part explain the association of CFH deficiencies with innate immune system dysfunction and inflammatory degeneration in both AD and AMD (Figure 2) [2-5, 7-12, 14-16, 42, 45]. As previously mentioned, a mutant, defective or ‘loss of function’ CFH, as generated by the Y402H polymorphism, may have analogous pathophysiological effects as insufficient CFH [2-5, 33, 41, 45; unpublished observations]. Targeting of the CFH mRNA 3’-UTR by miRNA-125b, miRNA-146a and miRNA-155, either separately or as a group within the MiRC region, has recently been demonstrated; a MiRC region is a miRNA regulatory control region found within a 3’-UTR to which multiple miRNAs may bind (Figure 2) [14, 16, 22-25, 45]. Note that miRNA-9 also has a binding site in the CFH mRNA 3’-UTR however its elevation in AD and AMD was much more variable than miRNA-125b, miRNA-146a and miRNA-155. An overlap of the miRNA-155 and miRNA-146a binding sites in the CFH mRNA 3’-UTR suggests evolutionary relationships or different, or overlapping, modes of CFH expression control in the brain and retina (Figure 2) [14, 16, 23, 73, 74]. Additional experiments using a 3’UTR-luciferase expression assay that utilizes a luciferase gene fused to the 232 nt human CFH 3’UTR (pLightSwitch-3’UTR Luciferase assay system; Cat # S801178; Switchgear Genomics, Menlo Park CA) shows that indeed the CFH mRNA 3’-UTR is highly sensitive to miRNA-125b, miRNA-146a and miRNA-155 inhibitory actions and that these may be restored using anti-miRNA strategies using nanomolar cocktails of anti-miRNAs (antagomirs; Figure 3) [23, 25, 45]. Lastly, the small size of miRNAs and the recent identification of miRNA-protective protein and miRNA-containing vesicles suggests that miRNAs may be a novel means for paracrine and related forms of inter-cellular and inter-tissue communication which may in part explain the propagation of the AD and AMD process over time [31, 55, 56].

Up-regulated miRNAs, their mRNA targets and effects on gene expression in the brain and retina

miRNA-9

Although miRNA-9 was easily detected in both the neocortex and retina it was found to be one of the most variably abundant miRNA in these tissues, perhaps indicative of its highly inducible nature [59, 60, unpublished observations]. With
miRNAs of the aging human brain and retina

Figure 2. Overlapping miRNA-155 and miRNA-146a high affinity binding sites in the CFH mRNA 3’-UTR (energy of association of less than -22 kcal/mol) defines an exceptionally stable miRNA-mRNA interaction and a potential CFH mRNA 3’-UTR miRNA regulatory control (MiRC) region 5’-TTTAGTATTAA-3’ (see text) [23, 31, 25, 45]; (A) entire nucleotide sequence of miRNA-155 and miRNA-146a interaction is highlighted in yellow; overlapping recognition for both miRNA-155 and miRNA-146a is highlighted in red; vertical black arrows indicate boundaries of miRNA-146a-miRNA-155-CFH mRNA 3’-UTR potential region of interaction; (B) shows entire 232 nucleotide sequence of the human CFH mRNA 3’-UTR; as in (A) entire region of miRNA-155 and miRNA-146a interaction is highlighted in yellow; overlapping MiRC region is highlighted in red; black vertical arrows indicate boundaries of miRNA-mRNA potential region of interaction; the single miRNA-9 site is upstream to this site; the miRNA-125b recognition region is distal to this site; white arrow = polyadenylation site; a 28 nucleotide polyadenylation feature is shown.

miRNA-125b

One of the most human brain and retinal abundant, if not the most abundant CNS miRNA and intensively studied miRNA is the inducible miRNA-125b, first shown to be up-regulated in differentiating mouse and human neurons, and since implicated in mammalian neuronal develop-
miRNAs of the aging human brain and retina

Development and function in the brain [63]. Both miRNA-125b and miR-9 play central roles in neuronal differentiation during retinal development [60]. The abundance of miRNA-125b has been shown to be significantly induced by neurotoxic metal sulfates that generate robust levels of oxidative stress, and miRNA-125b is also up-regulated in brain cancers where it apparently targets and down-regulates CDKN2A, a negative regulator of cell growth [41, 42, 64, 65]. Up-regulated miRNA-125b further associates with glial cell proliferation and astrogliosis in inflammatory neurodegenerative conditions such as AD and Down’s syndrome, as well as in glioma and glioblastoma multiforme [64]. Interestingly up-regulation of miRNA-125b is associated with down-regulation of both the 15-lipoxygenase (15-LOX) and the synaptic vesicle-associated phosphoprotein synapsin-2 (SYN-2) [21-25]. The 15-LOX enzyme is essential in the conversion of the essential omega-3 fatty acid docosahexaenoic acid (DHA) into the potent neuroprotectin D1 (NPD1), and deficits in 15-LOX correlate with NPD1 deficits in AD brain and other human tissues [46, 66]. The neuronal -enriched phosphoprotein SYN-2 that associates with the cytoplasmic surface of synaptic vesicles is also a miRNA-125b target [67]. MiRNA-125b up-regulation is further associated with SYN-2 down-regulation in inflammatory neurodegeneration and CFH down-regulation in human primary astroglial cells [16, 22, 64, 67].

miRNA-146a

miRNA-146a was first described as an NF-κB-regulated pro-inflammatory miRNA that was found to target signaling proteins of innate immune responses, and more specifically the 3' UTR of complement factor H (CFH) in human monocytes [68]. Elevated miRNA-146a in AD brain was subsequently shown to also target CFH and the interleukin-1 associated kinase 1 (IRAK-1) miRNAs, and is believed to contribute to altered innate immune responses and neuroinflammation in degenerating human brain cells and tissues in inflammatory neurodegenerative diseases including AD, AMD, prion disease, in experimental and human temporal lobe epilepsy and in experimental diabetes in retinal microvessel endothelial cells [14, 21, 23, 25, 41, 45, 69-71]. Although CFH was classically regarded as highly abundant human serum protein of hepatic origin, abundant CFH presence in brain and retinal tissues suggests CFH involvement in the innate immune response and inflammatory regulation within the ‘privileged immunology’ of these tissues [2-5, 14, 23]. While miRNA-146a is the least basally abundant miRNA when compared to miRNA-9, miRNA-
125b and miRNA-155, it is the most inducible and up-regulated miRNA in human neuronal and astroglial cells compared to all other NF-κB-regulated miRNA species so far indentified (Figure 1) [21-23, 45]. Interestingly, miRNA-146a may be the most induced of these up-regulated miRNAs due to the presence of 3 tandem, canonical NF-κB binding sites in the human pre-miRNA-146a promoter [23, 45, 68].

miRNA-155

MiRNA-155 is a cytokine, NF-κB and cell cycle-regulated miRNA also abundant in the human neocortex and retina, and has approximately 45% sequence homology to miRNA-146a (Figure 2) [25, 31, 72]. Interestingly, miRNA-146a and miRNA-155 have partially overlapping binding (recognition) sites in the CFH mRNA 3’-UTR, and miRNA-146a and miRNA-155 together define an overlapping miRNA regulatory control (MiRC) region in the CFH 3’-untranslated region (3’-UTR; 5’-TTTAGTATTA-3’) wherein either of these miRNAs may interact (Figure 2). As is true for miRNA-146a, inflammatory cytokines increase miRNA-155 expression in human retinal pigment epithelial cells by activation of the JAK/STAT signaling pathway [72]. Several additional studies indicate that miRNA-146a and miRNA-155 together play a key role in regulating several critical pathways that orchestrate innate immune responses and chronic inflammatory processes that are conserved across many different human tissue systems [77, 81].

Summary and conclusions

The five main conclusions of the data presented in this report are: (a) that normally aging brain neocortex and retinal tissues exhibit overlap in the speciation of their resident miRNAs; (b) that several brain and retinal miRNAs are similarly up-regulated during age-related inflammatory degeneration, and these include miRNA-9 and more significantly miRNA-125b, miRNA-146a and miRNA-155; (c) that miRNA-125b, miRNA-146a and miRNA-155 all have high affinity binding sites in the CFH mRNA 3’-UTR, supportive of their roles in regulation of CFH and the immune response; (d) that miRNA-146a and miRNA-155 both recognize an overlapping MiRC region in the CFH mRNA 3’-UTR that is conducive to down-regulation in the expression of CFH in both the brain and retina; and (e) that multiple miRNAs are capable of targeting and down-regulating an important repressor of the innate immune and inflammatory response. It is tempting to speculate that a subfamily of human miRNAs that include miRNA-125b, miRNA-146a, miRNA-155, and perhaps miRNA-9 and others, may help coordinate innate immune and inflammatory signals across the entire anteroposterior axis of the retinal-primary visual cortex pathway, and perhaps also in other immune-responsive cells and tissues. Other brain and retinal mRNA 3’-UTR targets, and ensuing down-regulated gene expression are probably affected by this miRNA-mRNA control system. This work in progress presents several testable hypotheses. For example it will extremely interesting to expand our understanding of the role of NF-κB with miRNAs and with other transcription factors, chromatin-remodeling mechanisms, and other epigenetic influences on specific miRNA-mRNA activation pathways to further understand their surprisingly dynamic interactive roles, and their coordinate contribution to the neurogenetic control of brain cell aging in health, aging and degenerative disease. It will be further interesting to see what individual brain and retinal cell types might be contributing to these multiple miRNA and CFH aberrations in AD and AMD [16, 22, 31, 45], and how this knowledge can be profitably manipulated using anti-miRNA and related pharmacological approaches that have not yet been considered in the effective management of these diseases.

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References

[22] Pogue AI, Percy ME, Cui JG, Li YY, Bhattacharjee S, Hill JM, Kruck TPA, Zhao Y, Lukiw WJ. Up-regulation of NF-kB-sensitive miRNA-125b and miRNA-146a in metal sulfate-stressed human
miRNAs of the aging human brain and retina


[31] Lukiw WJ. NF-kB-regulated micro RNAs (miRNAs) in primary human brain and retinal cells. Exp Neurol 2011; [Epub ahead of print].


miRNAs of the aging human brain and retina


[72] Kutty RK, Nagineni CN, Samuel W, Vijayasara-thy C, Hooks JJ, Redmond TM. Inflammatory cytokines regulate microRNA-155 expression in human retinal pigment epithelial cells by acti-
miRNAs of the aging human brain and retina


