

Advances in Imaging: Brain Tumors to Alzheimer's Disease



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Editor Note:

Professor Black and colleagues have been working to improve the quality and sensitivity of imaging in the early detection of conditions from brain tumors to Alzheimer's disease to enhance treatment protocols and patient management. Professor Black et al introduced nanoparticles to improve MRI imaging. These nanoparticles consist of poly (b-L- malic acid (PMLA)) conjugates with monoclonal antibodies ((mAbs)) and Gd-DOTA. These are known as MRI nano-imaging agents (NIA). Most importantly, they can penetrate the endothelial blood-brain barrier (BBB) to reach brain tumors (primary or metastasis). This is effective in cases of brain tumors or breast cancer or other cancers such as lung cancer and gastric cancer having HER2 and/or EGFR positive crossing BBB. By the covalent conjugation of MR contrast (NIA), the MRI virtual biopsy can differentiate brain tumors from infections or other brain pathological conditions. The brain's intrinsic natural fluorescence such as NADH, FAD, lipopigments and porphyrin in the brain tissue can be identified by using time resolved fluorescence spectroscopy (TRFS) which is operated through the use of ultra-short laser. TRFS produces various color bands to differentiate the tumor from normal brain tissue in real time and registers the data on a 3D map. This is significant, as this will provide a greatly improved assessment methodology of tissue type. Consequently, this will potentially result in shorter operation times as well as more satisfactory tumor removal. In the detection of Alzheimer disease, amyloid plaque is deposited in retina tissue (including the RGC, RNFL and inner plexiform layer) which can produce a fluorescence effect by using curcumin as a contrast. This is then shown by human retina amyloid imaging device. Immunotherapies with glatiramer acetate (GA) have been shown to reduce amyloid deposits in brain and retinal AB deposits in mice. The study of advanced imaging technology and techniques including NIA, TRFS and the detection of amyloid plaque in Alzheimer disease are very important approaches to create a new era for diagnostic and therapeutic management of brain tumors and other cancers (HER2 and/or EGFR positive). This pioneering work by Professor Black, and colleagues, gives rise to a new hope for cancer patients for targeted therapy and for immunotherapies in Alzheimer's disease.

Keywords: nanoconjugate, brain metastasis, MRI imaging, nano-medicine, blood-brain barrier, tumor treatment, tumor targeting, glioma, time-resolved fluorescence spectroscopy, in-vivo diagnosis, neurodegenerative disease, retina, amyloid plaques, senile plaques, eye, curcumin, optical imaging, prognosis, immunotherapy, vaccination

Technical and biological advances are improving early detection, classification, surgical, and medical management of a wide range of human disorders. In this publication, we will present advances using nano-molecular imaging to provide an imaging MRI molecular classification of brain tumors and time-resolved fluorescence spectroscopy to improve differentiation of normal from tumor infiltrated tissue during surgery in real-time. In this same communication, we will also present results demonstrating early detection of one of the hallmarks of Alzheimer's disease, amyloid plaques, using a non-invasive retinal scanning technique.

1. Brain tumors-Nano imaging

Brain tumors include primary and secondary deposits, and the primary deposit is relatively rare. The majority of brain metastasis (BM) are lung and breast which accounts for around 20-30% of cancer deaths annually.¹⁻⁶ 25-40% of BM originate from breast and lung cancer respectively.⁷⁻⁸ Non-small cell lung cancer shows the highest incidence followed by small cell carcinoma.⁸ In the case of breast cancer with overexpressing epidermal growth factor receptor-2 (HER2/neu) and triple-negative breast cancers (TNBC) but over expressing epidermal growth factor receptor-1 (EGFR) developing BM is a poor survival outcome.⁹⁻¹² The incidence of BM is increased even when the primary site is progressing well with treatment.^{13,14}

Management of BM is complex because contrast enhancing lesions in the brain may represent several things, from a primary tumor metastasis, to an infection in an immunocompromised patient after chemotherapy, from posttreatment radiation necrosis to a mimic (or actual) tumor recurrence. The therapeutic monoclonal antibodies (mAbs) e.g. trastuzumab (to HER2) for breast, ovarian and gastric cancer, cetuximab (to EGFR) for lung and breast cancer and rituximab (to CD20) for lymphoma are effective only for primary tumor treatment. However, they cannot penetrate the endothelial blood-brain barrier (BBB)¹⁵ to reach brain tumors, and this is one of the causes of treatment failure.^{13,16} These mAbs can, however, be used for drug delivery to the brain when they join with nano vehicles capable of efficiently crossing the BBB.¹⁷⁻²¹ Nano

vehicles are made up of MultiHance, a gadolinium (Gd)-based MRI contrast agent which is routinely used and is highly effective in detecting brain tumors. We have engineered a second, a new class of MR nanoimaging agents (NIA) based on poly (beta-L-malic acid)(PMLA) as a platform with a lack of toxicity and immunogenicity.²² It can demonstrate high versatility for on-demand covalent conjugation of proteins, chemotherapeutic agents and imaging agents.^{17,23,24} We engineered a NIA with various mAbs and conjugated these with Gd-DOTA (Figure 1) that can penetrate through the BBB. The composition of the nanoconjugate and hydrodynamic diameter/zeta potential is summarized in Table 1. When compared, MultiHance, Targeted NIA cetuximab and non-targets NIA trastuzumab in case of EGFR+++ the findings reveal a higher contrast enhancement on Gd-DOTA/acetuximab (Alexa-680) and less enhancement on Gd-DOTA/trastuzumab (Alexa-680), with the highest contrast in the tumor at 60 minutes (Figure 2 and Figure 3).

In HER2 positive cases, with targeted NIA with trastuzumab, the findings reveal increased contrast in the tumor within 50 minutes and this slightly declines at 180 min (Figure 4). This diagram shows that the nanoconjugates efficiently cross the BBB and there is high probability of a mechanism of nanodrug action on the tumor cell (Figure 5). Figure 6 demonstrates that free transtuzumab is inefficient for the treatment of metastatic brain tumors due to its inability to pass through the BBB, but when transtuzumab conjugates with PMLA, it can penetrate BBB and reach the tumor

Table 1: Summary of nanoconjugates: their abbreviations and physicochemical characterization.

Imaging and treatment agents	Hydrodynamic diameter (nm) ^a	Zeta potential (mV) ^b
poly(β-L-malic acid) (PMLA)	6.6 (±0.1)	-22.9 (±1.7)
P/MEA(4%)/Gd-DOTA(10-12%) ^c (Preconjugate-1)	8.8 (±0.7)	-16.8 (±1.8)
P/cetuximab(0.12%)/MsTfR-mAb(0.12%)/Alexa-680(1%)/Gd-DOTA(10-12%) (P/Gd-DOTA/cetuximab/MsTfR-mAb/Alexa-680)	16.0 (±1.5)	-8.7 (±1.2)
P/trastuzumab(0.12%)/MsTfR-mAb(0.12%)/Alexa-680 (1%)/Gd-DOTA(10-12%) (P/Gd-DOTA/trastuzumab/MsTfR-mAb/Alexa-680)	16.3 (±1.6)	-6.9 (±1.1)
P/mPEG(5%)/LOEt(40%)/MEA(4-6%) (Preconjugate-2)	8.1 (±0.5)	-8.45 (±1.2)
P/mPEG(5%)/LOEt(40%)/HuTfR-mAb(0.2%) MsTfRmAb(0.2%)/EGFR-AON(2-4%) (P/Hu/MsTfR-mAb/EGFR-AON)	15.7 (±2.0)	-2.61 (±0.8)
P/mPEG(5%)/LOEt(40%)/trastuzumab(0.2%)/MsTfR (0.2%)/HER2 AON(3-4%) (P/trastuzumab/MsTfR-mAb/HER2-AON)	15.7 (±2.0)	-4.09 (±0.8)

^aHydrodynamic diameter by number distribution at 25°C measured in PBS at a concentration of 2 mg/ml, calculated from DLS data by Malvern Zetasizer software (Malvern Instruments, Malvern, UK), which assumes spherical shapes of particles.

^bzeta potential at 25°C in aqueous solution of 10 mM NaCl at 150 mV.

^ccomposition of nanoconjugates; percentage refers to total number (100%) of pendant carboxyl groups in unsubstituted PMLA.

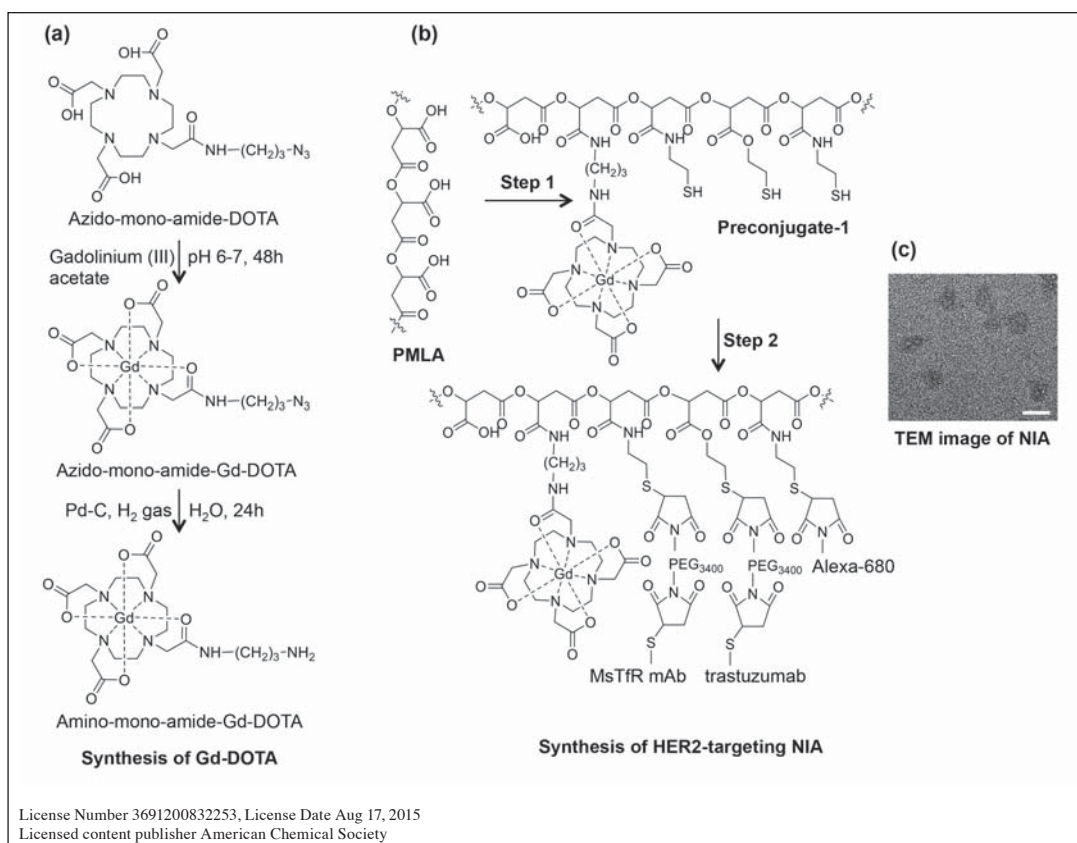


Figure 1

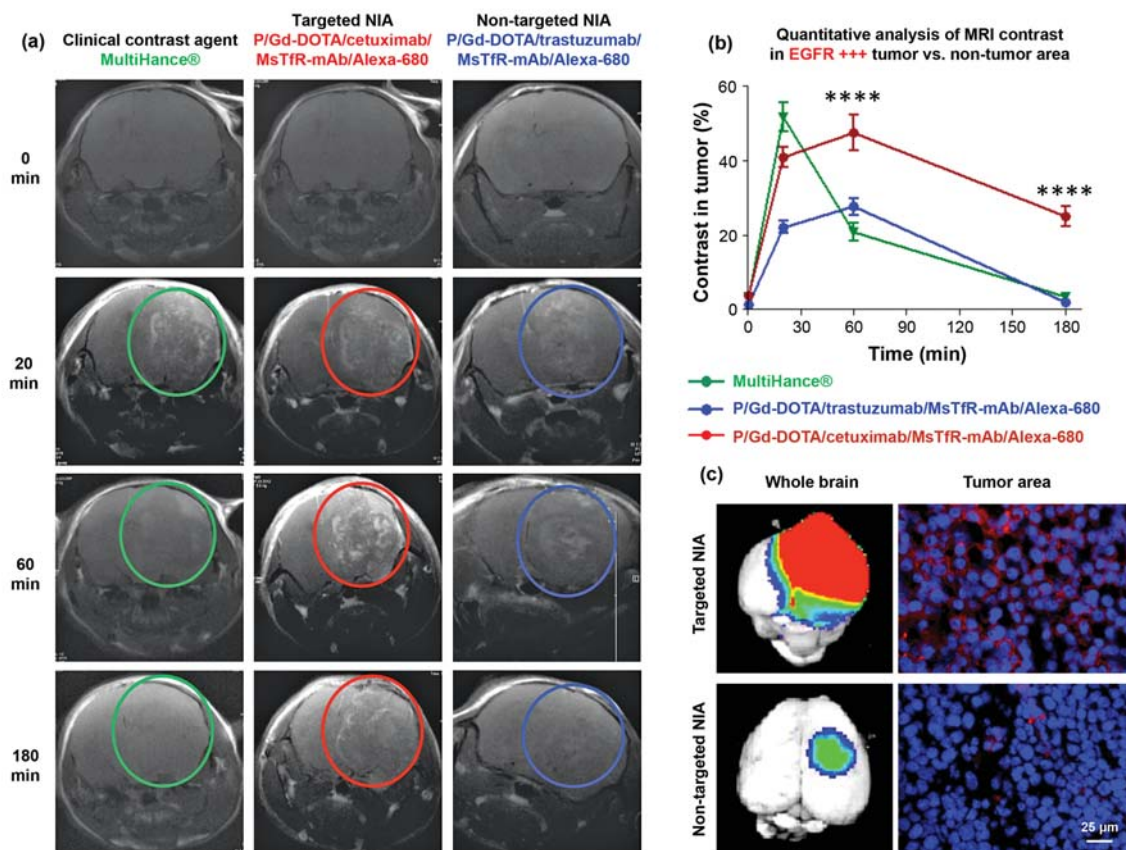


Figure 2

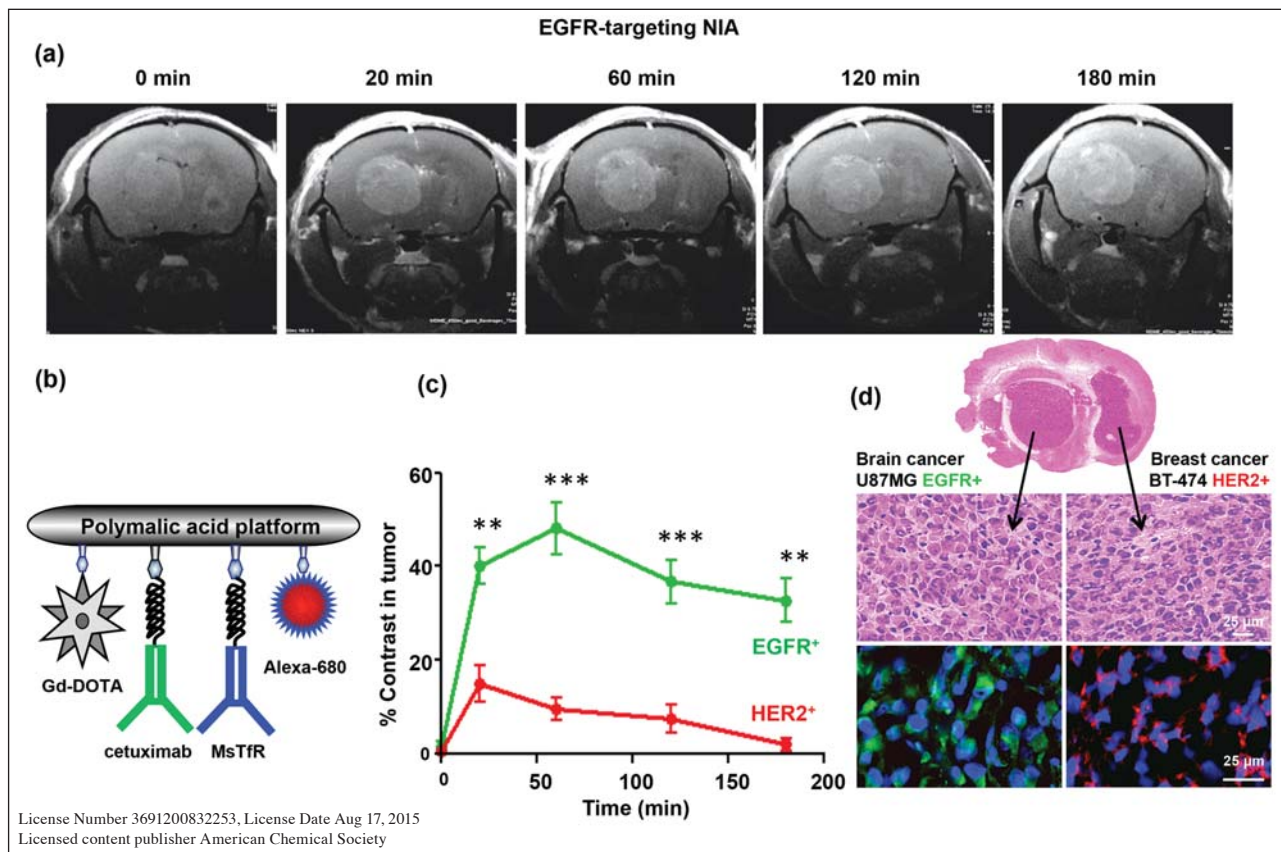


Figure 3

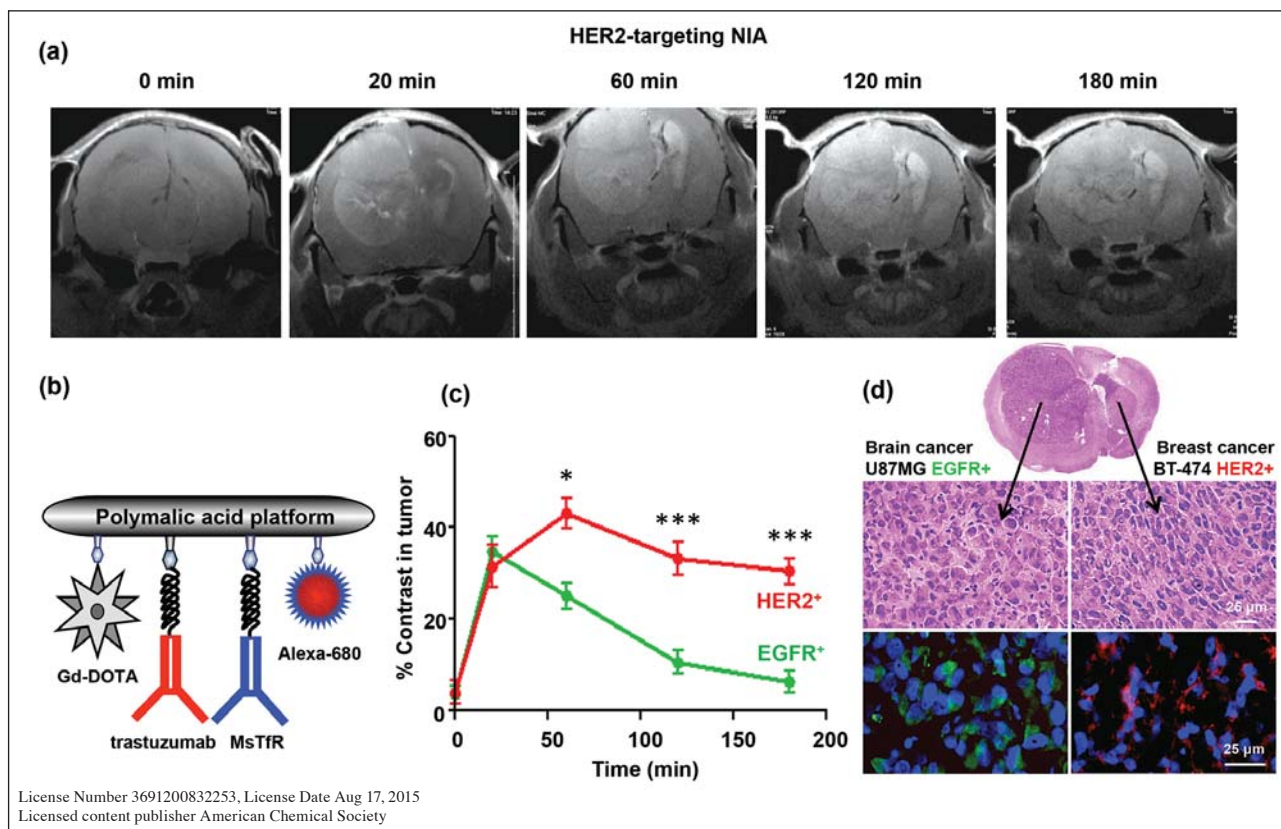


Figure 4

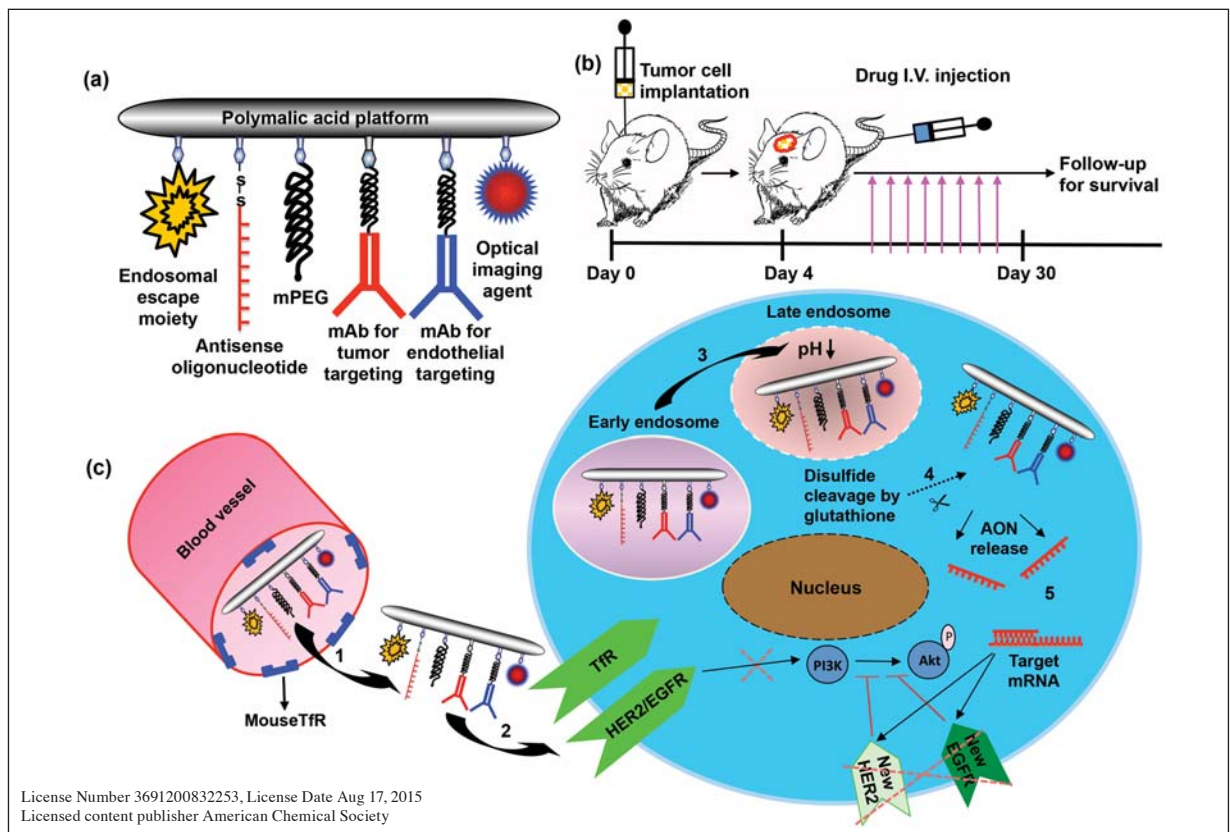


Figure 5

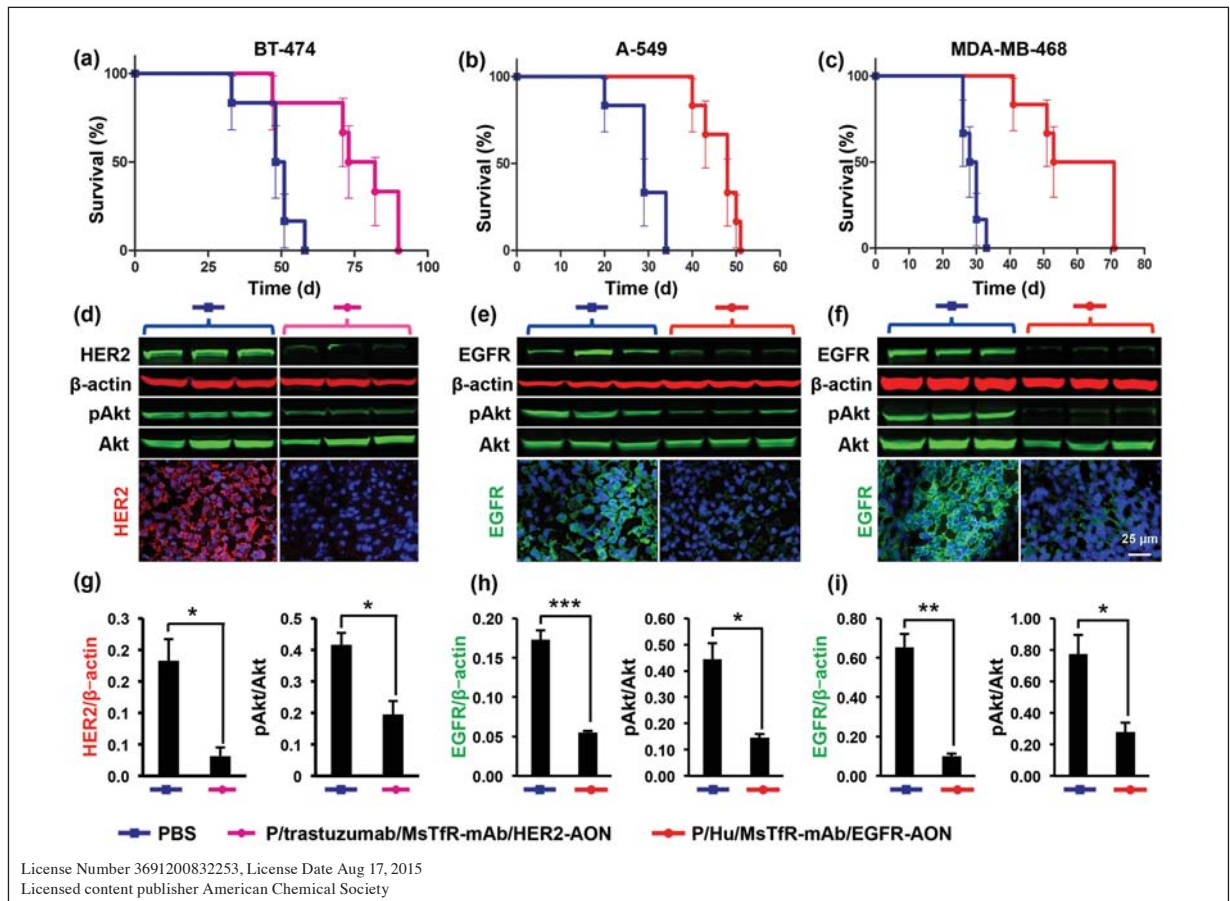


Figure 6

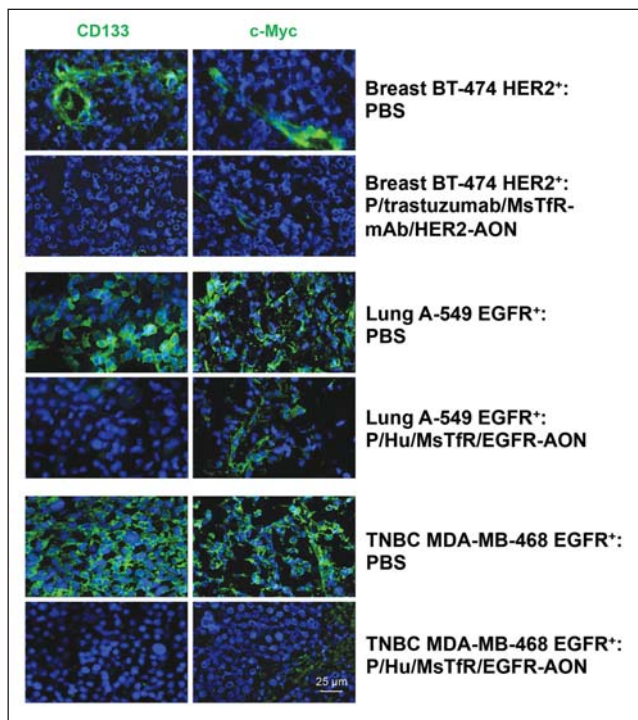


Figure 7

cells acting as a specific targeting antibody of HER2 and as an inhibitor to the activation of existing HER2 receptors.²⁴⁻²⁸ The survival of the nanodrug treated in mice increased. Figure 7 demonstrates that the nanodrug can inhibit receptors in both wild and mutated EGFR.

2. Time resolved fluorescence spectroscopy (TRFS)

In its natural state, the brain tissue has natural fluorescence properties.³⁵⁻³⁷ The main fluorophores are NADH, FAD, lipopigments and porphyrins^{38,39} (Figure 8). In general, tumors have lower fluorescence emissions when compared to normal tissue at the excitation light of 355nm wavelength and with a ratio of NADH. FAD as a signature for brain tumor differentiation.⁴⁰

We introduced an instrument, the time resolved fluorescence spectroscopy. This operates where the tissue is excited using an ultrashort laser and the corresponding fluorescence intensity is captured, based on the fluorescence spectrum and the decay characteristics of various color bands from TRFS. The differentiation of the tumor tissue from normal brain tissue is possible in real time as

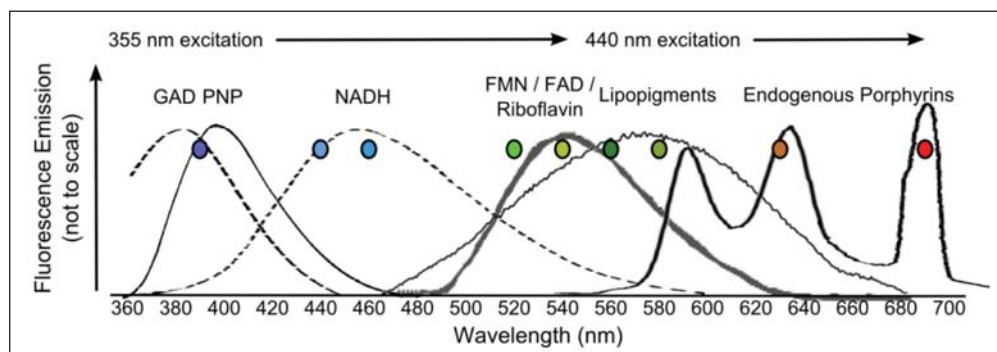


Figure 8: Natural fluorophores of biological tissue.

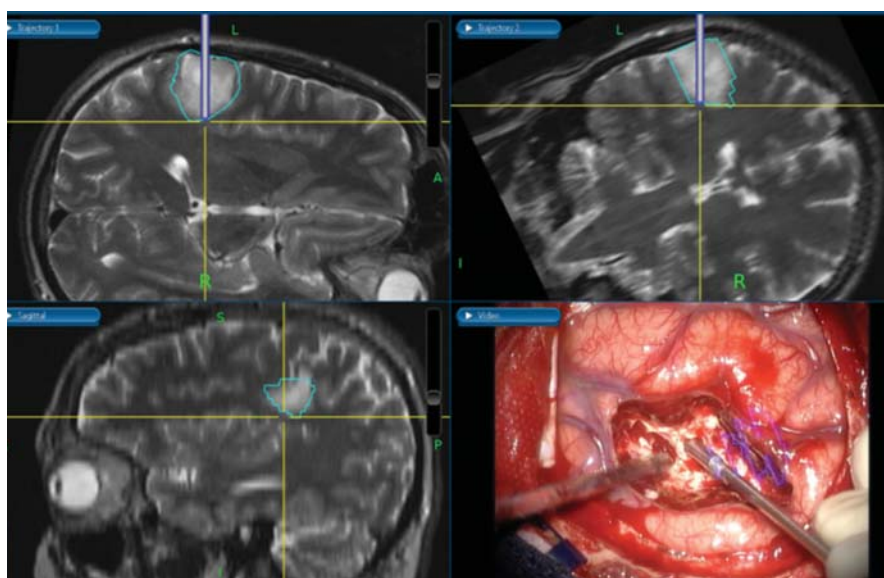


Figure 9: TRFS measurement registered by neuro-navigation.

TRFS is a single point detection tool. It works by incorporating a neuro-navigation system and registering 3D space. Neurosurgeons can continue carving the tumor boundary during an operation (Figure 9), and this information will allow the surgeon to identify areas the suction cannula is carving in, whether normal brain tissue or tumor.

This results in a significant impact on the assessment of tissue type and leads to improvements such as reducing operating time and waiting time for the frozen section. The TRFS system uses a 355nm laser pulse (400 ps,

5uJ/pulse) to excite the tissue fluorescence (Figure 10). We experimented with 9 patients of glioma. During the operation, the TRFS fiber optic probe was positioned above the areas of interest, glioma, normal cortex (NC), and normal white matter (WM). A total number of 35 points included 10 points of NC, 12 points of normal WM and 13 points for GBM. The linear discriminant analysis (LDA) classification has been implemented and integrated in a three-dimensional scatterogram. See Figure 11, which is identical to the areas of NC, WM and GBM by a 3D scatterogram of all 9 patients.

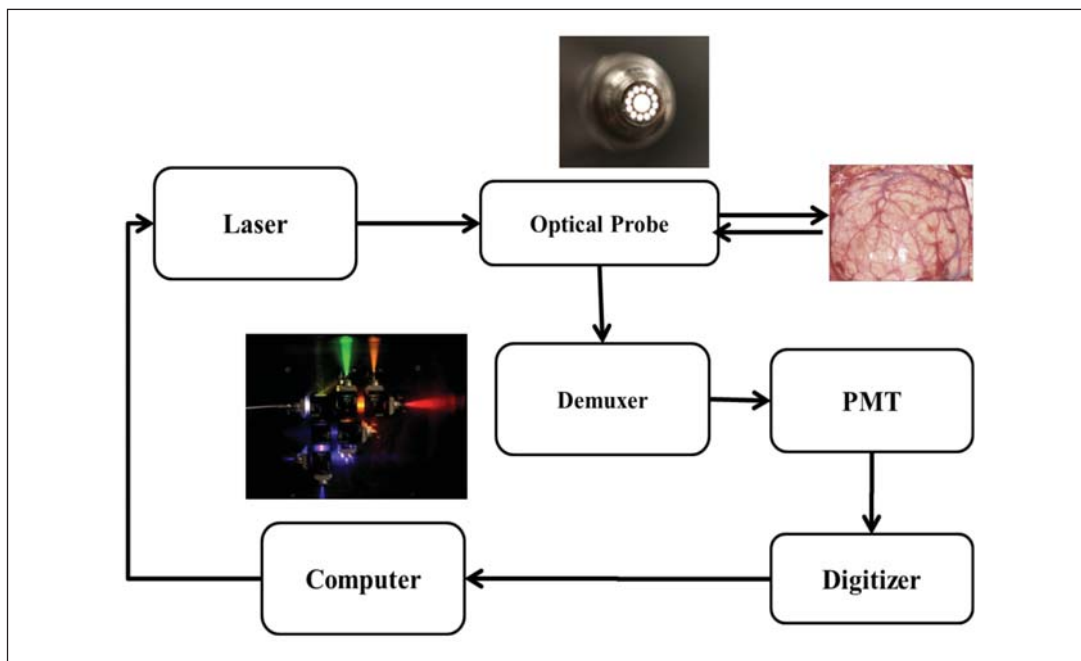


Figure 10: TRFS system diagram.

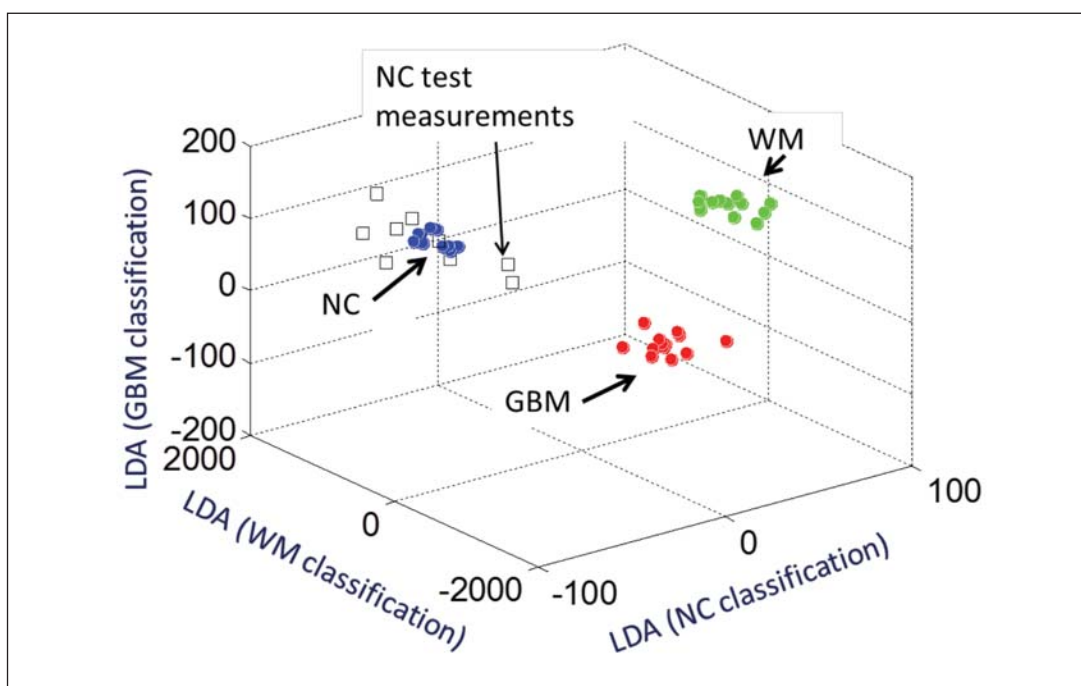


Figure 11: TRFS Classification output for differentiating, Normal Cortex, White Matter, and GBM tissue types.

3. Retinal Imaging for Alzheimer's disease

Alzheimer's disease (AD) is the leading cause of dementia in the United States, and the fifth leading cause of death among the elderly,⁶³ claiming more than 5 million Americans.⁶³ It is estimated that the number of Americans affected by AD could triple by 2050.⁶³ The cost of medical care for AD in the US is more than \$200 billion a year currently and this would increase to up to \$1.1 trillion in 2050.⁶³ It is estimated that 44 million people are suffering from Alzheimer's disease or other dementias globally⁴¹ and this number could increase to 76 million in 2030 and 135 million in 2050.^{42,43}

Early diagnosis of AD could potentially result in early treatment or timely prevention of disease progression.

Today, a confirmation of definitive AD is only possible with an autopsy of brain tissue, with an examination for the presence of two pathological hallmark signs of AD: amyloid (senile) plaques predominantly composed of proteinaceous A β and neurofibrillary tangles (NFT) formed by an aggregation of the hyperphosphorylated microtubule-associated protein tau.^{44,45} Cognitive tests such as the Mini Mental State Examination (MMSE) can determine level of impairment, but not the type of disease. Clinical, genetic, physiologic, and biochemical evidences suggest that the primary pathological event leading to AD is an age-dependent accumulation of toxic alloforms of A β in the brain.^{46,47} This elevation in A β levels results in a self-association, formation of neurotoxic A β assemblies, and deposition of fibrillary A β in amyloid plaques. Elevated cerebral levels of A β seem to be a very early and specific event in AD that likely occurs 10-20 years prior to symptom manifestation.⁴⁷⁻⁵⁰ Although methods are currently being developed to detect amyloid-related changes directly within the skull-shielded brain (i.e. PET, MRI), there are many difficulties and limitations inherent in these techniques such as low specificity, poor spatial resolution and high costs.⁵¹⁻⁵⁸ Presence of A β plaques in the retina of people with AD, possibly at pre-symptomatic stages, and the ability to detect them with high detail and accuracy offers a new improved opportunity for the early detection of AD. The retina is a part of the central nervous system, originating as an outgrowth of the developing brain and shares many similarities with the brain. The retina is easily accessible for direct and noninvasive imaging with high spatial resolution and sensitivity.⁵⁸

In retinal imaging for AD patients, our research group at Cedars-Sinai, also founding members of Neurovision Imaging, made a groundbreaking discovery-the first identification of A β plaque pathology in retinas of AD patients: with definite, probable and possible diagnosis, comparable with that observed in their brains.⁶⁰ Our studies with AD transgenic mice have further shown that A β plaques may appear in the retina before they appear in

the brain. Our group has developed an approach to detect those plaques in live rodent models with unprecedented high detail.^{59,60} Retinal plaques in mice were mainly detected in the nerve fiber layer (NFL) throughout the outer plexiform layer, and their prevalence correlated with disease progression, retinal ganglion cell (RGC) degeneration, microglial activation, and functional impairment.⁶¹ Therefore it is essential to develop a method to detect A β plaques in live patients by a noninvasive and direct retinal imaging with high resolution and less expense compared to brain imaging techniques, which will enable a detailed evaluation of AD-specific pathology at an early stage.

The retina may offer a better target to detect AD pathology since existing brain-imaging technologies are radioactive, invasive and cannot provide sufficient detail about early changes in amyloid build up. A direct retinal-imaging test may offer details related to early AD pathology in real-time, including smaller-size plaque accumulation, individual plaque dynamic (appearance and clearance), plaque distribution and geographical location near retinal structures (i.e. blood vessels) that neither brain imaging nor the highly invasive CSF A β levels currently provide. In collaboration with Carol A. Miller, MD (Pathology and Neurology, USC), we continue to collect rare retinal and brain samples isolated from AD patients at different stages of the disease. This is a unique collection of well-characterized AD patients that have both pre-mortem cognitive/behavioral and postmortem complete brain pathological assessments.

Results and Discussion

Our Cedars-Sinai research group is a pioneer in identifying the feature A β plaque pathology in postmortem retinas of AD patients (Figure 12), and in early-stage cases (Figure 13), comparable with plaque load observed in their brains. Retinal A β plaques in AD patients were found to be located mostly within the inner layers. Retinal plaques location and morphology were qualitatively documented and compared with those found in the brain. In AD patients, retinal A β plaques were mostly found abluminal and near blood vessels, and had typical morphology of presumably earlier-stage; condensed hard-core(s) containing aggregated A β and lack of radiating fibril arms, less frequently seen in the brains. These findings suggest that A β plaques may appear sooner in the retina than in the brain. A quantitative analysis (Figure 14) of curcumin-labeled retinal A β plaques in AD-Tg mice indicated a significant decrease in their number and size following immunotherapy and to the same extent as their brain counterparts.⁵⁹⁻⁶⁰ These findings provided further evidence for a correlation between retinal and brain plaques. Moreover, curcumin was found to cross the blood-retinal and brain barriers and label A β with high affinity and specificity in vivo in mouse model retinas and brains. More importantly, in early stage of the disease, A β plaques appeared in the

retina before their appearance in the brain (Figure 15). A novel noninvasive retinal imaging approach developed by our group, enabled the detection of A β plaques with high sensitivity, specificity, and repeatability in live AD transgenic (AD-Tg) mouse models, after intravenous (iv.) injection of curcumin or when given orally (Figure 16). A follow up study from our group demonstrated the

feasibility to noninvasively monitor individual A β -plaque appearance and clearance in the retina of transgenic AD models in response to an immunotherapy (Figure 17). Phase I and II clinical trials using curcumin showed low toxicity profiles in humans even at high doses (8-12 gr/day), and when given over extended periods of time.⁶²

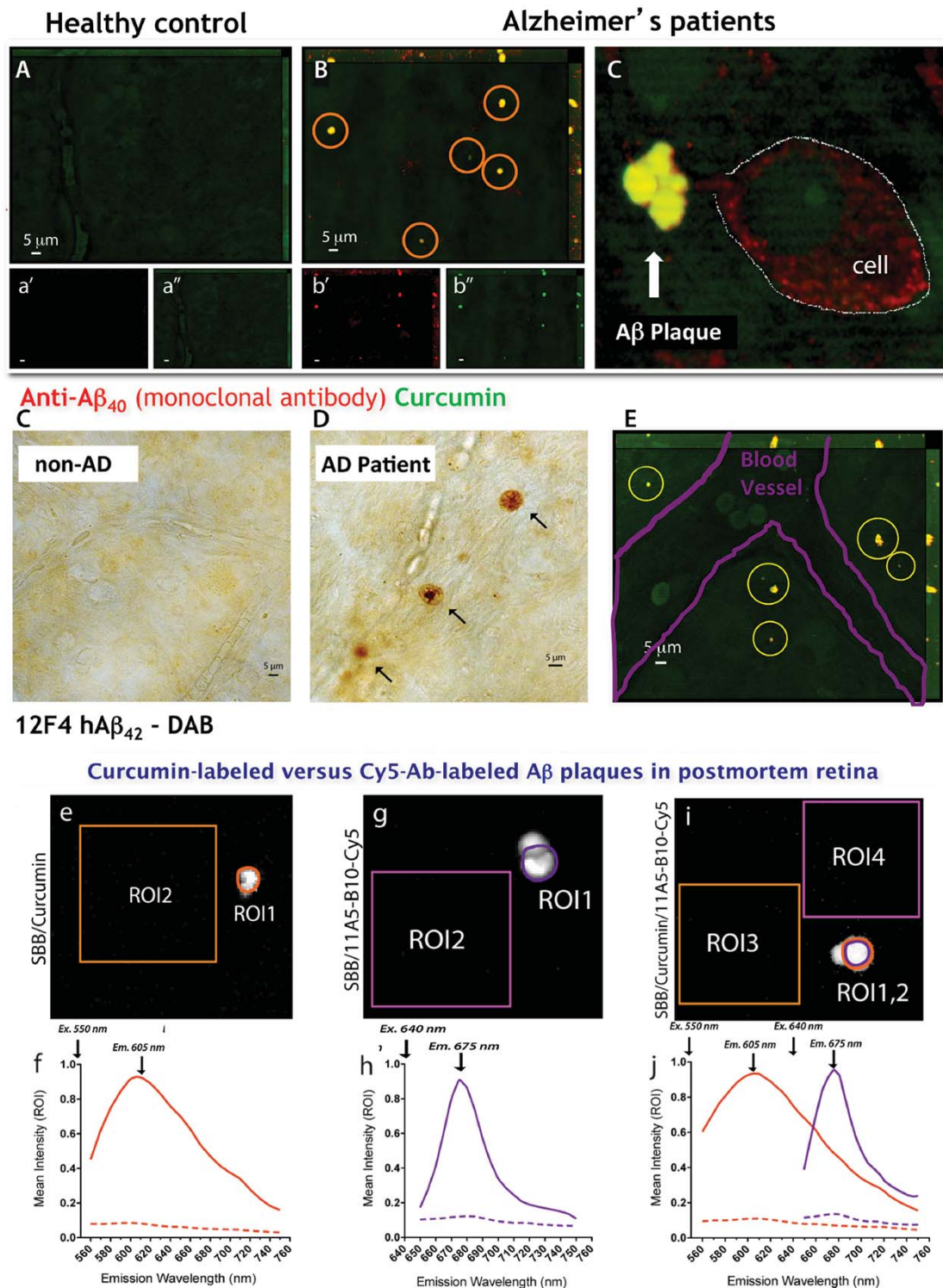
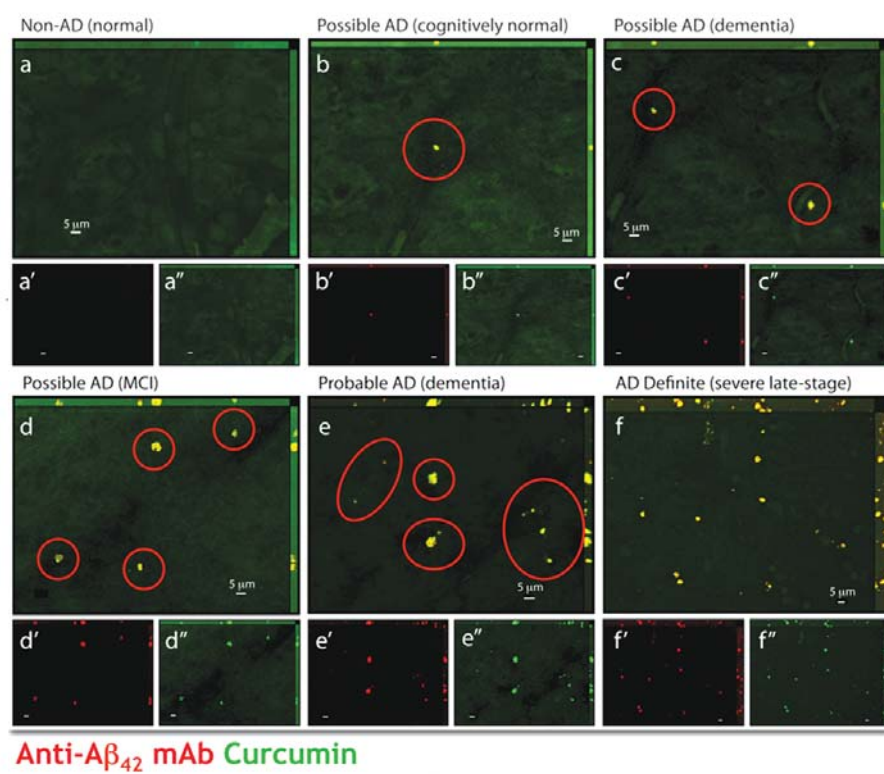
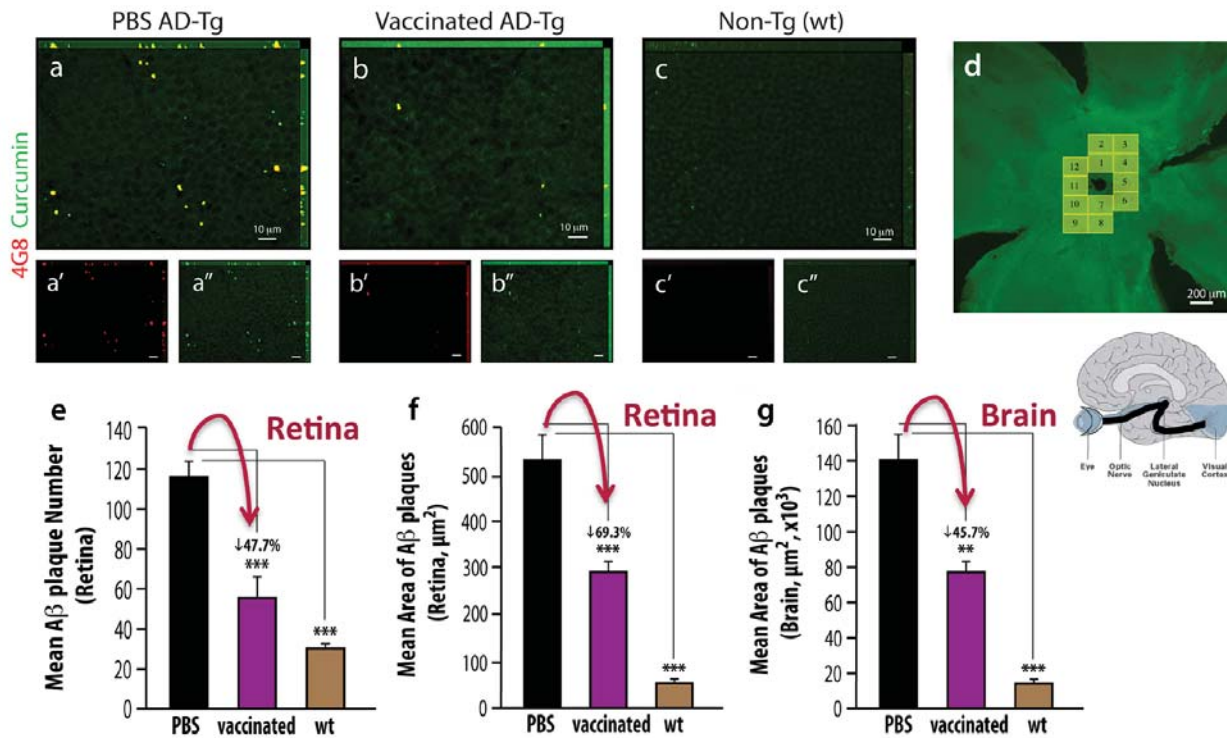


Figure 12: Koronyo-Hamaoui et al., Neurolmage 2011.



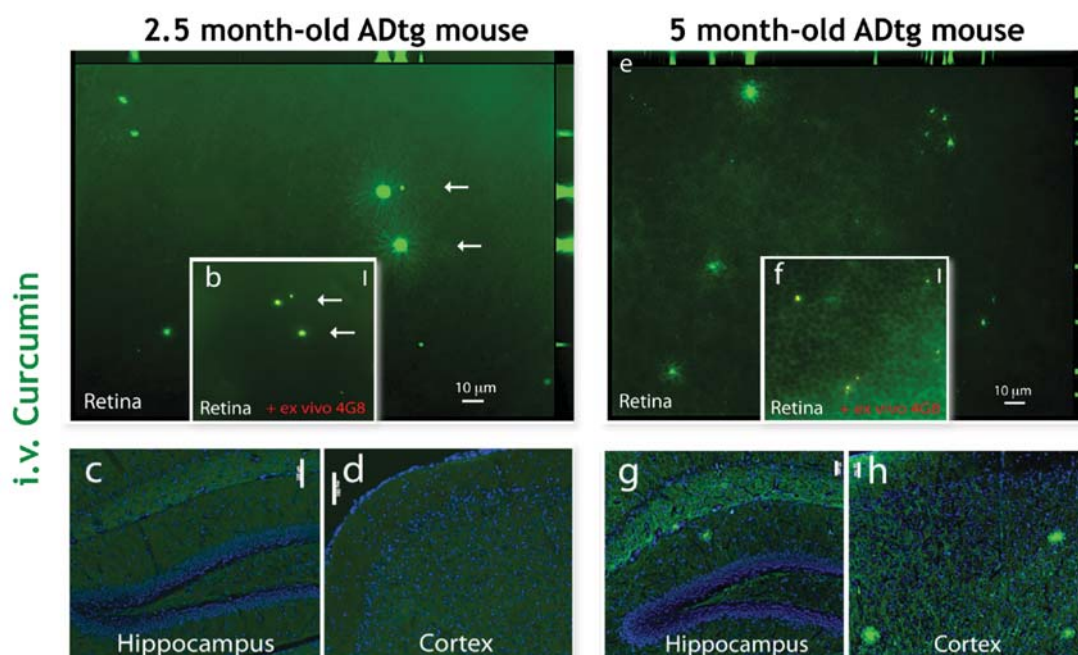
Koronyo-Hamaoui et al., Neurolmage 2011

Figure 13: Hallmark plaques detected early in the retina of AD patients.



Koronyo-Hamaoui et al., Neurolmage 2011

Figure 14: Quantitative Assay: amyloid-beta plaque reduction in the retina to the same extent as in the brain in response to immunomodulation therapy.



Koronyo-Hamaoui et al., NeuroImage 2011

Figure 15: Accumulation of amyloid-beta deposits in the retina with disease progression in ADtg mice; Earlier detection of plaques in the retina than in the brain.

Conclusion

We demonstrate several examples of imaging technologies which aim to improve detection and classification, as well as surgical and medical management of brain tumors and Alzheimer's disease.

1. Brain tumors-Nano imaging. A significant advantage of our nanosystem is that it is possible to use not only for imaging but also for the delivery of specific mRNA suppressors (AONs) or other drugs with simple structural manipulations of nanoplateforms for efficient tumor treatment.^{31,32} This versatility was illustrated here by successfully treating BMs from breast and lung tumors using different tumor-targeting mAbs (e.g., trastuzumab for HER2+ tumors and anti-human TfR for EGFR+ tumors (Figure 6). For BM treatment, Gd-DOTA in NIAs was just substituted by HER2- or EGFR-specific AONs to block the synthesis of these tumor markers along with endosomal escape units for cytoplasmic delivery and PEG for protection. These AONs were delivered to the tumor cell cytoplasm while sparing healthy brain tissue (Figure 18). They efficiently blocked the synthesis of new HER2 or EGFR, and suppressed the activation of a downstream PI3K/Akt signaling pathway. This pro-survival pathway may contribute to apoptosis prevention, regulation of cell cycle progression, and enhanced tumor cell survival.^{29,30,33-34}

2. Time resolved fluorescence spectroscopy (TRFS). TRFS has the potential to be an extremely helpful surgical tool that will allow surgeons to more precisely identify tumor margins for a wide variety of tumors, thus, improving surgical outcomes.

3. Retinal Imaging for Alzheimer's disease. The retina is part of the central nervous system. Both the optic nerve and the retina have been shown to succumb to neurodegeneration in AD. A β accumulation in the retina has been well documented in AD mouse models.^{60,64-66} We have successfully documented A β -deposits in the retina of patients with AD and MCI and APP/PS1 Tg mouse models NDD76. The original discovery of amyloid-beta plaques in the retina of AD patients at early stages⁶⁰ was followed by the demonstration of increased Abeta1-42 peptides in AD patients' retina⁶⁷ and further detection of plaque-like structures in the postmortem retina of an AD patient.⁶⁸

The retina is easily accessible for direct and non-invasive imaging with high spatial resolution and sensitivity. Our group was the first to report that pathology in definite, probable, and possible AD patient retinas correlate with brain amyloidosis.⁶⁰ We also reported that A β deposits appear very early in the retina and along the blood vessels of AD transgenic mice-seemingly even before plaques appeared in the brain.⁶⁰ Several groups have shown therapeutic amyloid changes in the retinas of AD and glaucoma rodent models.^{60,69,70}

Since the retina can contain autofluorescence spots, the use of amyloid binding contrast such as curcumin is critical to reliable identification of amyloid deposits in the retinal tissue. The extensive studies performed by the Koronyo-Hamaoui team to identify and image A β plaque thru the retina provided the grounds to develop the human retinal amyloid imaging device.

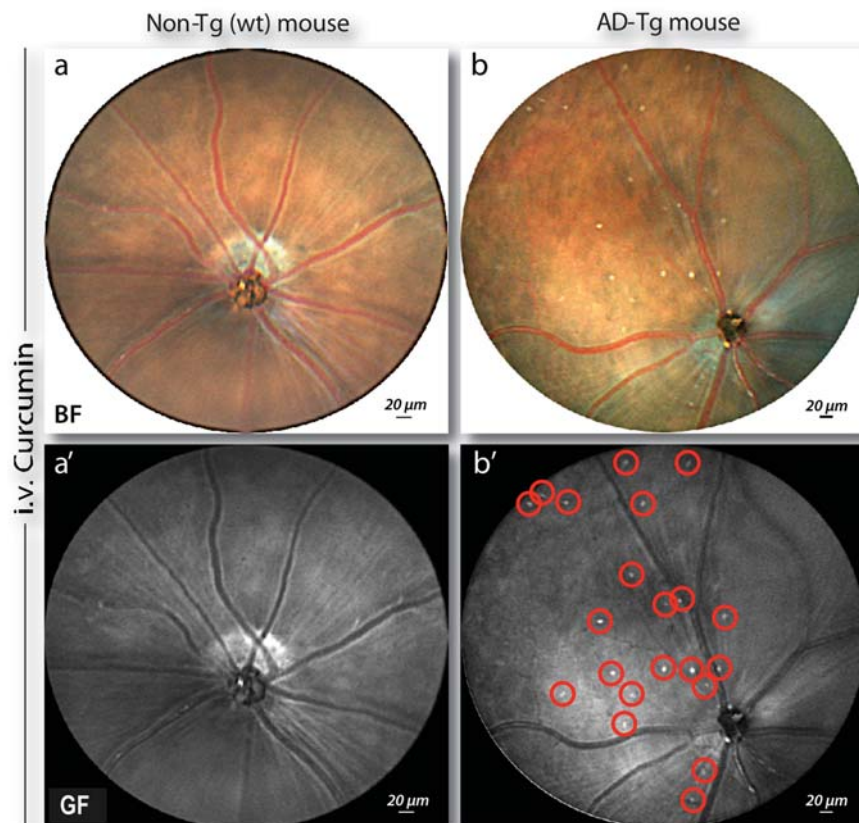
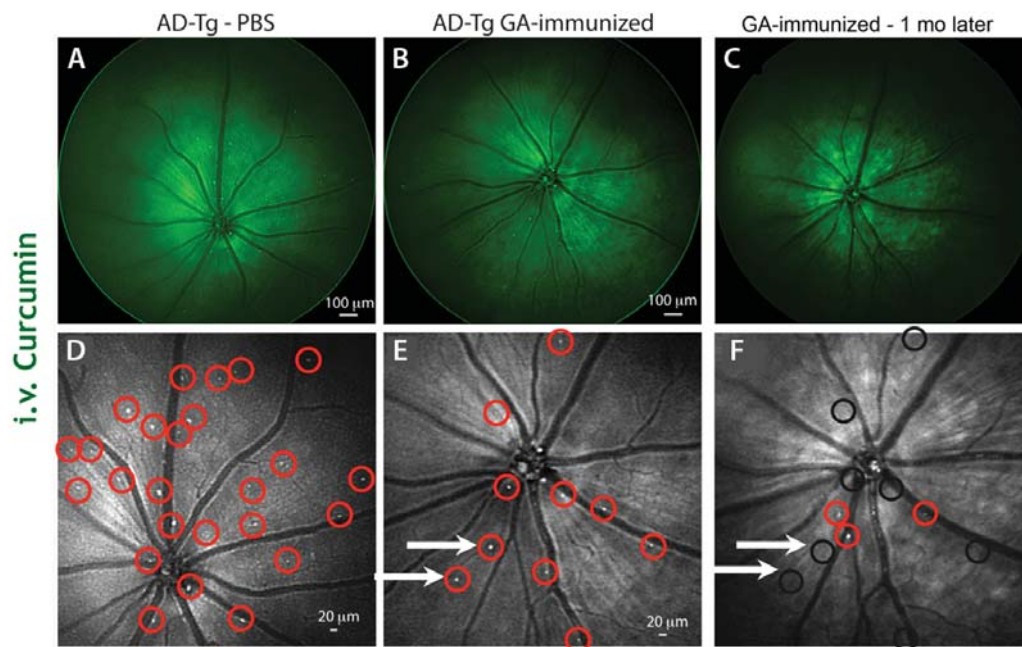


Figure 16



Longitudinal study of curcumin-labeled A β plaques in the mouse retina following glatiramer acetate (GA)-immunotherapy. Intravenous (i.v.) injection of curcumin to ADtg mouse allows visualizing *in vivo* retinal A β plaques. Images A – C are in green fluorescence (GF) channels, D – F are the same images magnified and in gray scale. White spots are A β plaques. Image D shows AD-Tg mouse treated with PBS where A β plaques are indicated by red circles. After a month of GA treatment a reduction in A β plaques is observed (Image E). After another month of GA, we monitored dynamic changes in distinct retinal A β plaques (Image F). Noticeable decrease in A β plaques as indicated by black circles and as marked by white arrows and appearance of new A β plaques as indicated by red circles.

Koronyo et al., *Neurodegenerative Dis.* 2012

Figure 17

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