

IJPBS |Volume 5| Issue 2|APR-JUN|2015|134-144

R<u>eview Article</u> P<u>harmaceutical Sciences</u>

IN VIVO IMAGING TECHNIQUES FOR DETERMINATION OF THE FATE OF DRUG DELIVERY SYSTEMS

Abhishek Bhattacharjee*

Department of Pharmaceutical Sciences, Assam University Silchar, Pin- 788011, India. *Corresponding Author Email: <u>abhishek18jan@yahoo.co.in</u>

ABSTRACT

The various bio imaging techniques provides a non invasive method to monitor the in vivo fate of the different pharmaceutical formulations post administration into the body. The different imaging techniques including gamma scintigraphy, near infra red fluorescent dyes helps to predict the pharmacokinetic behaviour of dosage form in subjects under investigation. The transit of dosage form through its intended site of delivery can be non-invasively imaged in vivo via the judicious introduction of an appropriate short lived gamma emitting radioisotope and non isotopic florescent dyes. Near-infrared (NIR) fluorescent probes offer advantages of high photon penetration, reduced light scattering and minimal autofluorescence from living tissues, rendering them valuable for noninvasive mapping of molecular events, assessment of therapeutic efficacy, and monitoring of disease progression in animal models. These studies provide an insight into the outcome of delivery systems, its integrity as well as enable the relationship between in vivo performance and resultant pharmacokinetic parameters. This review provides an overview of the recent development of the design and optical property of the different classes of NIR fluorescent nanoprobes as well gamma scintigraphic technique associated with in vivo imaging applications.

KEY WORDS

Bioimaging, Gamma Scintigraphy, Radiolablling, Fluroscent probes, NIR.

INTRODUCTION

The first medical imaging was realized in the late 1895 by Wilhelm Roentgen shortly after he discovered X-ray and applied to capturing the images of the bones of a hand on film. From the first anatomical charts to the most recent development of Magnetic Resonance Imaging (MRI) or Positron Emission Tomography (PET), setting into images what is normally unseen within living systems has been a major issue for the understanding of biological processes [1–3]. These imaging technologies differ predominantly in the following aspects: resolution, penetration depth, temporal resolution and energy expended for generation of the image. With the exception of MRI, which relies on magnetic properties of atom nuclei with half-integer spin, most of the imaging modalities rely on photons with wavelengths ranging from gamma emission to infrared. Any photon has specific interactions with tissues or bones that will define its uses and drawbacks. On one hand, high-energy photons such as gamma emission or X-rays can go through the tissues with low interactions, making them suitable for deep tissue or whole body imaging, while being potentially harmful because of the reactions they can trigger within cells. Their use is thus strictly restricted to dedicated areas and under the supervision of trained personnel. On the other hand, photons in the visible and

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*

Int J Pharm Bio Sci

 $_{\rm Page}134$



infrared wavelengths do not penetrate deeply within organic tissues, but they cause no or little damage to cells. Moreover, light sources and imaging apparatus are widespread, relatively cheap, their handling requires moderate levels of training and protection, and does not generate radioactive wastes.

Contrary to tissue auto-fluorescence recording, near-infrared/visible fluorescence is an optical imaging modality that relies on the injection of an exogenous probe that will emit light when excited with suitable wavelength. This method can be used from the macroscopic to the microscopic range, goes as further as sub-cellular resolution, and is highly sensitive. Targeted fluorescent probes can be designed to specifically mark and visualize different biological targets, which can be imaged with high contrast using the appropriate set of optical filters to minimize autofluorescence contribution of the surrounding tissues. Fluorescence is particularly suitable for in vitro and superficial in vivo imaging applications. However, fluorescence imaging using the near infrared range is also now routinely used for whole body imaging of small animals in preclinical studies [4-7], and the first human clinical trials have been performed in 2008/2009 for sentinel lymph node mapping in oncology by different groups [8-12]. The next challenge to address is to pursue the development of fluorescence imaging techniques in medical applications. Since the instrumentation tools now exist [13-18], the main blocking point to overcome for the expansion of clinical trials remains the availability of performing fluorescent tracers [19,20]. Nanotechnologies could facilitate their design by allowing modular and flexible constructions. This review will focus on the *in vivo* imaging and the coming challenges for their clinical translation.

Among the different imaging modalities, optical imaging, which owes its origin to single-cell *in vitro* studies, is attractive for small animal

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

imaging because of its lower cost, portability, and potentially high spatial resolution with customizable fine-tuning. Despite being one of the most attractive techniques to provide noninvasive and nonionizing in vivo visualization [21], optical imaging is impeded by the tendency of living biological tissues to absorb and scatter photons and generate strong autofluorescence, which interferes with signal collection and processing[22]. In addition, living tissues also contain other major NIR absorbers, such as water, lipids oxyhemoglobin and deoxyhemoglobin [23] that prove challenging for optical imaging. To overcome these barriers, intense research has focused on developing highly sensitive and efficient fluorescent probes that function in the biologically transparent window of the first and second NIR region (NIR I, 650-950 nm, and NIR II, 1000-1350 nm)[24].

Information about the in vivo behaviour of dosage forms when obtained using radionuclide tagged with dosage forms is known as gamma scintigraphy. This radiolabelled method is a well established technique in medical practice for the past several decades. Moreover, instead of relying on pharmacokinetic data alone, it is better to combine it with the technique of gammascintigraphy or pharmacoscintigraphy to assess the performance of dosage form in man. The coupling of gamma scintigraphy with usual pharmacokinetic methods makes it possible to correlate the information obtained on the distribution of medicament with complete absorption profile. Radionuclide tagged with dosage forms can provide vital information regarding the extent, rate, site and mode of drug release and morphology of the drug delivery system during release in human under ethical norms. Conclusions can also be drawn about the performance of the formulation, including: the ability of a formulation to target a specific location; the rate of erosion in comparison with in

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



vitro dissolution data; the impact of absorption window on bioavailability.

Today, these modern imaging technologies coupled with newly developed imaging probes are widely used in monitoring disease progression, interrogating cellular and molecular events, evaluating safety and toxicology in drug discovery and development, and assessing therapeutic efficacy *in vivo* [25-33]

Gamma Scintigraphy Technique

Nuclear medicine images are primarily functional in nature, providing substantial information on physiology and more limited information on anatomy. The technique commonly termed as scintigraphy involves imaging the bio distribution of a radiopharmaceutical. Gamma camera images of the in vivo distribution of pharmaceutical formulation radiolabelled with a suitable gamma emitting radionucleotide, may be used to quantify the biodistribution of the formulation. No other technique can locate so precisely the site of disintegration of the formulation in the gastro intestinal tract. Scintigraphic technique also allows correlation between observed pharmacological effects and the precise site of delivery and thus facilitating drug targeting studies.

Radiolabelling by Gamma-Emitting Isotopes

Prior to imaging by this technique, the dosage form should be radiolabelled. Drug/formulations can be radiolabelled either by gamma emitting isotopes or by neutron activation technique. Emitted radiations are further captured by external detectors such as gamma cameras.

Radiolabelling of formulation involves utilization of a short-lived radioisotope that can spontaneously emit gamma radiation. The isotope is incorporated in the formulation as a salt (e.g., sodium pertechnetate) in normal saline.

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

In this type of labelling, a reducing agent is used to reduce technetium, Tc (VII), into a lower valence state. The final solution is maintained in a buffered system, which is followed by incubation to enable labelling [34]. The most widely used reducing condition is the acidic stannous chloride or other stannous salts [35]. Various approaches for radiolabelling can be classified into whole dose radiolabelling, point radiolabelling and surrogate marker technique. Commonly used radionuclides are shown in table 1 along with their half-life [36].

•	
Radionucleotides	Approximate Half Life
81mKr (Krypton)	13 seconds
⁹⁹ mTc (Technetium)	6.02 hours
¹¹¹ In (Indium)	2.8 days
¹²³ I (Iodine)	13 hours
¹³¹ l (lodine)	8.05 days

Labelling by Neutron Activation Technology

This approach involves the incorporation of a stable isotope into the dosage form prior to its manufacture, followed by neutron irradiation of the intact dosage form (table 2). Neutron flux exposure is conducted for a very short time, generally 5-30 seconds. This exposure time has been shown to maintain the characteristics and integrity of the dosage form. Short period of time also prevents the degradation of drug under conditions of bombardment. Longer exposures may result in cross-linking of the polymers used in the dosage form. During this technique, thermal neutron irradiation converts the carefully selected stable isotopes (¹⁵²Sm or ¹⁷⁰Er) into radioactive gamma emitting isotopes (¹⁵³Sm or ¹⁷¹Er) that can be detected by external imaging devices [37-40].

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



Table 2: Commonly used radionucleotides used in neutron activated scintigraphy

Radionucleotide	Natural abundance (%)	Approximate Half Life	Stable Nuclide
¹³⁹ Ba (Barrium)	71.7	83 minutes	138Ba (Barrium)
¹⁷¹ Er (Erbium)	14.9	7.5 hours	170Er (Erbium)
¹⁵³ Sm (Samarium)	26.7	47 hours	151Sm (Samarium)

Gamma Camera

Gamma scintigraphy relies on the detection of radiations emitted from a radionuclide. In this technique, gamma camera is used to image gamma radiation emitted from radioisotopes Nuclear imaging is predominantly carried out with planar or SPECT (single photon emission computed tomography) cameras and by using radionuclides that emit gamma radiation with energies between 100 and 250 KeV. The single photon emitting radioisotopes such as ⁹⁹mTc and ¹¹¹In are widely used with these instruments. Gamma camera is composed of an array of photomultiplier tubes coupled to a sodium iodide crystal. The interaction of a gamma photon from the source with the crystal leads to the production of a flash and it is detected by photomultiplier. To ensure that radiation from the source is detected in straight line, a lead collimator is placed between the subject and the crystal. The camera provides two dimensional or planar images of the distribution of radioactivity in the subject. The planar image provides a good depiction of the position of the radiotracer. For this reason, radiolabelled drug delivery systems are best studied with planar camera [41]. If planar imaging cannot provide the required deposition details, then SPECT should be considered. SPECT is a technique for producing crosssectional images of radionuclide distribution in the body. This is achieved by imaging the organ at different angles (e.g. 64 or 128 images/ 180⁰ or 360°) using a rotating gamma-camera. The acquired raw data are then processed by highspeed computers [42].

Applications of Gamma Scintigraphy

The use of imaging techniques particularly gamma scintigraphy to follow the behavior of drug formulations has revolutionized our knowledge of absorption and distribution in drug delivery. Following are the few applications of Gamma Scintigraphy technique for determination of the fate of the dosage forms under study-

- 1. Pulmonary drug delivery
- 2. Oral drug delivery
- a. Gastroretentive
- b. Colon targeting
- 3. Ocular drug delivery
- 4. Rectal drug delivery
- 5. Site specific targeting
- a. Brain targeting
- b. Tumor targeting
- c. Liver & spleen targeting

Further, formulations involving colloidal particles, liposomes, micro-emulsions, and dendrimers can also be effectively be evaluated. The major advantage of gamma scintigraphy is that it is free of animal scarification and can therefore be especially useful in exploring sources of intersubject variation. This method can be applied to both experimental animals and humans alike. Hence, it is a versatile technique for the investigators who can use this method to generate real time pharmacokinetic and pharmacodynamic data during various stages of drug development.

Fluorescence imaging technique using NIR probes

Molecules that absorb in the near-infrared (NIR) region, 700–1000 nm, can be efficiently used to visualize and investigate *in vivo* molecular targets

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



because most tissues generate little NIR fluorescence [43, 44]. The most common organic NIR fluorophores are polymethines. Among them, pentamethine and heptamethine cyanines comprising benzoxazole, benzothaizole, indolyl, 2-quinoline or 4-quinoline have been found to be the most useful [45].

An emerging new class of probes for in vivo fluorescence imaging is semiconductor nanocrystals or quantum dots. Quantum dots (QDs) typically have a core/shell structure of 2–8 nm in diameter with size-dependent fluorescence emission. The unique optical properties of QDs for in vivo optical imaging include high absorbency, high quantum yield, narrow emission bands, large Stokes shifts, and high resistance to photobleaching. QDs that emit at several different wavelengths can be excited with a single wavelength, and thus are suitable for multiplex detection of multiple targets in a single experiment. Several recent reviews summarize the synthesis, bioconjugation chemistry, optical features and applications of

QDs for in vivo imaging [46, 47].

Labeling Mechanism of Fluorescent Probes

Optical imaging has increasingly been used for dynamic noninvasive imaging of biological events in mouse models. Due to the lack of NIR fluorescence contrast generated by most tissues in the biological transparency window, exogenous fluorescent probes have to be administered for *in vivo* studies in order to visualize living tissues in its native physiological state. Ideally, the fluorescent probes to be administered should be biologically stable in the *in vivo* environment, and accumulate and produce imaging contrast at the target site. Fluorescent probes are classified according to their mechanism of contrast generation, collectively as non-specific, targeting and activatable.

Non-specific probes simply have differential distribution and are used to assess physiological

processes such as changes in blood volume, permeability and perfusion in angiogenesis. Typically, they achieve only low target-tobackground signals due to the non-binding circulating probes producing significant background fluorescence within the compartment.

Tissue- or cell-specific contrast is created by coupling a targeting moiety to a fluorescent robe that binds specifically to a receptor thus generating the readout signal. These probes can report more detailed information about the biological events than non-specific probes. Targeting probes can achieve high target-tobackground signals provided the targeting moiety has a high affinity for the receptor and any unbound probes are thoroughly removed from the system, thereby reducing the background fluorescence.

Activatable probes comprise of donor-acceptor fluorophores that are coupled to each other in close proximity to maintain a quenched state. The fluorescence emission is activated by enzymemediated cleavage that releases the fluorescent probes. Activatable probes can attain high targetto-background signals as these probes in their native injected state are relatively undetectable.

Non-Specific Organic-Dye Probes

The choice of a suitable dye for *in vivo* imaging depends on many factors. The most important consideration would be the molar extinction coefficient and quantum yield of the dye in the NIR region [48, 49]. Of all the dyes, cyanine makes up the majority of commercial fluorescent probes for *in vivo* applications. Cyanine is a synthetic dye family of the polymethine group with a conjugated chain of odd number of carbon atoms linked between two nitrogen centers [50]. Among this class of compounds, carbocyanine dyes with indolic groups are readily available commercially and have also been synthesized in a large variety of analogues. The structure of carbocyanine is

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



formed by reaction of two indolic isomers (either identical or different) linked on each end of a C1, C3 or C5 methane. The absorption and fluorescence emission wavelength is determined both by the chain length of the methane and the side chains attached to the indolic groups. Carbocyanine dyes emits in the visible to NIR range. These dyes generally exhibit high molar extinction coefficients but only have moderate fluorescence quantum yields up to 30%. A prominent representative of the NIR cyanine dye is Indocyanine green (ICG). ICG was first synthesized in the fifties [51] and is the only clinically approved dye available commercially. It has been used in optical imaging of changes in blood-brain barrier permeability after thrombus formation in a mouse model of cerebral venous thrombosis [52] and liver perfusion in mouse [53]. Clinically, ICG has been used in clinical retinal angiography [54], hepatic function testing [55] and imaging of human brain [56, 57]. Since ICG binds tightly to plasma proteins and becomes confined to the vascular system, they are widely used for intraoperative assessment of vascular flow in cardiovascular surgery [58-60]. Because of its amphophilicity and a shortage of functional groups available for conjugation, ICG could only function as a non-specific probe for in vivo imaging.

To improve the pharmacokinetics of in vivo contrast agents, a possible approach is to reduce the conjugate molecular size while still preserving the targeting affinity of the labeled dyes. Neri et al. demonstrated this idea by conjugating antibody single chain fragments selected from phage display libraries against an angiogenesisassociated oncofetal fibronectin isoform to cyanine dyes and applied to imaging of angiogenesis in a variety of animal models [61, 62]. Alternative strategy is also shown in the work by Achilefu et al. and Licha et al. highlighting the successful application of receptor-specific

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

peptide-dye conjugates for fluorescent imaging of tumors [63–65]. As a whole, fluorescent dyes can be readily functionalized to achieve targeting.

Targeting Organic-Dye Probes

Covalent attachment of targeting moieties such as antibodies, antibody fragments, proteins and peptides to fluorescent dyes can significantly improve the target-to-background signal. The strategy of conjugating antibodies to cyanine dyes were first demonstrated by Folli et al. and Ballou et al. [66, 67] and applied to fluorescence imaging of tumor in mice. Antibodies, however, have found limited utility due to their unfavorable pharmacokinetics. The typical circulating half-life of antibodies is much shorter than the time required to access a small tumor with reasonable accumulation, thus rendering effective targeting rare using antibodies [68].

Activatable Organic-Dye Probes

The concept of activatable NIR probes was introduced by Weissleder et al. for in vivo imaging of enzyme activity [69]. Enzyme-activatable probes contain either two identical or different fluorophores linked in close proximity to each other by a specific peptide linker. The fluorescence of the probes is essentially undetectable in the guenched state. After enzymatic cleavage, the fluorophores are separated to restore the fluorescence emission. A large number of activatable probes have been documented, among them the enzyme activation of NIR targeting dyes is mediated mainly by tumor associated proteases, such as cathepsins, caspases and matrix metalloproteinases [70-75]. The ability of activatable fluorescent probes to detect gene expression is particularly useful as a means to diagnose malignant molecular process in the early disease state [76].

Nanomaterial-Based Fluorescent Probes

Despite the long history of *in vivo* optical imaging, organic dyes have suffered from small Stokes shifts, poor photostability, high plasma protein

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



binding rate, aggregation and background fluorescence in aqueous medium [77]. Attention has been increasingly channeled toward nanotechnology to search for new class of more effective probes. Many nanomaterials are already widely used as catalysts, and energy storage and electronic devices [78-82]. Their application to the biomedical arena has attracted a similarly enthusiastic following. Interest in the nanomaterials arises from the fact that at nanoscale the properties of materials can be very different from their bulk counterparts. Firstly, nanomaterials when compared to the same mass of material existing in a larger form have a much higher surface area. As such, nanomaterials are more chemically reactive because of the high surface energy. In addition, at nanoscale range, quantum effects begin to dominate the behavior of matter, affecting the optical, electrical and magnetic behavior of materials [83, 84].

Lately, the integration of nanotechnology with medicine has found many novel applications of nanomaterials. Nanoparticles can be engineered to overcome biological barriers for effective and targeted delivery of drugs, genes, and contrast agents [85]. Active moieties such as proteins, peptides and nucleic acids can easily be conjugated to the surface of the nanoparticles for use as non-specific, targeting or activatable nanoprobes. The following sections are devoted to the recent development of some particulate NIR fluorescent nanoprobes for potential *in vivo* imaging.

CONCLUSION

From the foregoing discussion it is evident that the bioimaging of Pharmaceutical dosage form for determination of the *in vivo* efficacy past administration in the body can be done effectively by using the above mentioned techniques. Pharmacoscintigraphic technique is an elegant approach to gain insight of the actual

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

in vivo distribution pattern of dosage form along with the assessment of pharmacokinetic information. Conventional methods for evaluating these formulations in vivo are laborious and time-consuming. The data obtained from scintigrams, although qualitative, are quite reliable in predicting the extent to which a formulation has been targeted to its destined site. As a tool to examine drug delivery to the lung and to the eye, scintigraphy is the method of choice and needs to be exploited to the maximum for its potentials in the evaluation of molecular entities, drug delivery new systems/formulations and in therapeutic drug monitoring. The potential of NIR fluorescent nanoprobes for in vivo application is unlimited. In particular the versatile, nanomaterials-based NIR fluorescent probes offer ample opportunity to optimize their optical and targeting properties. nanomaterial-based fluorescent Among all probes, Si QDs is the only system that can be directly loaded with drugs and biodegraded into silicic acid that can be excreted efficiently through renal clearance. Like other nanomaterialbased fluorescent probes, improvement of their quantum efficiency, however, will still require significant effort. Robust and environmentally friendly synthetic routes that can produce these nanomaterials-based NIR nanoprobes with uniform size and colloidal stability represent other significant challenges. The widespread application of NIR fluorescent nanoprobes awaits demonstration of their long term biocompatibility, in vivo targeting efficacy as well as parallel development of advanced optical instrumentation for deep-tissue imaging capability. Nevertheless, their role in advancing the field of nanomedicine will undoubtedly be essential.

REFERENCES

1. Massoud, T.F.; Gambhir, S.S. Molecular imaging in living subjects: Seeing fundamental biological

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



Available Online through

www.ijpbs.com (or) www.ijpbsonline.com

processes in a new light. Genes Develop. 2003, 17, 545–580.

- Massoud, T.F.; Gambhir, S.S. integrating noninvasive molecular imaging into molecular medicine: An evolving paradigm. Trends Mol. Med. 2007, 13, 183– 191.
- Frangioni, J.V. New technologies for human cancer imaging. J. Clin. Oncol. 2008, 26, 4012–4021.
- Leblond, F.; Davis, S.C.; Valdès, P.A.; Pogue, B.W. Preclinical whole-body fluorescence imaging: Review of instruments, methods and applications. J. Photochem. Photobiol. B: Biol. 2010, 98, 77–94.
- Hassan, M.; Klaunberg, B.A. Biomedical applications of fluorescence imaging *in vivo*. Comp. Med. 2004, 54, 635–644.
- Koo, V.; Hamilton, P.W.; Williamson, K. Non invasive in vivo imaging in small animal research. Cellular Oncol. 2006, 28, 127–139.
- Licha, K.; Olbrich, C. Optical imaging in drug discovery and diagnostic applications. Adv. Drug Deliv. Rev. 2005, 57, 1087–1108.
- Miyashiro, I.; Miyoshi, N.; Hiratsuka, M.; Kishi, K.; Yamada, T.; Ohue, M.; Ohigashi, H.; Yano, M.; Ishikawa, O.; Imaoka, S. Detection of sentinel node in gastric cancer surgery by indocyanine green fluorescence imaging: Comparison with infrared imaging. Ann. Surg. Oncol. 2008, 15, 1640–1643.
- Ogasawara, Y.; Ikeda, H.; Takahashi, M.; Kawasaki, K.; Doihara, H. Evaluation of breast lymphatic pathways with indocyanine green fluorescence imaging in patients with breast cancers. Word J. Surg. 2008, 32, 1924–1929.
- Sevick-Muraca, E.M.; Sharma, R.; Rasmussen, J.C.; Marshall, M.V.; Wendt, J.A.; Pham, H.Q.; Bonefas, E.; Houston, J.P.; Sampath, L.; Adams, K.E.; Blanchard, D.K.; Fischer, R.E.; Chiang, S.B.; Elledge, R.; Mawad, M.E. Imaging of lymph flow in breast cancer patients after microdose administration of a near-infrared fluorophore. Radiology 2008, 246, 734–741.
- Tagaya, N.; Yamazaki, R.; Nakagawa, A.; Abe, A.; Hamada, K.; Kubota, K.; Oyama, T. Intraoperative identification of sentinel lymph nodes by near-infrared fluorescence imaging in patients with breast cancer. Am. J. Surg. 2008, 195, 850–853.
- Troyan, S.L.; Kianzad, V.; Gibbs-Strauss, S.L.; Gioux, S.; Matsui, A.; Oketokoun, R.; Ngo, L.; Khamene, A.; Azar, F.; Frangioni, J.V. The FLARE intraoperative nearinfrared fluorescence imaging system: A first-in-human clinical trial in breast cancer sentinel lymph node mapping. Ann. Surg. Oncol. 2009, 16, 2943–2952.
- 13. Ntziachristos, V.; Ripoll, J.; Wang, L.V.; Weissleder, R. Looking and listening to light: The evolution of whole-

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

body photonic imaging. Nat. Biotechnol. 2005, 23, 313–320.

- Laidevant, A.; Hervé, L.; Debourdeau, M.; Boutet, J.; Grenier, N.; Dinten, J.-M. Fluorescence time-resolved imaging system embedded in an ultrasound prostate probe. Biomed. Opt. Express 2011, 2, 194–206.
- Liu, Y.; Bauer, A.Q.; Akers, W.J.; Sudlow, G.; Liang, K.; Shen, D.; Berezin, M.Y.; Culver, J.P.; Achilefu, S. Handsfree, wireless goggles for near-infrared fluorescence and real-time image-guided surgery. Surgery 2011, 149, 689–698.
- Qin, C.; Zhu, S.; Tian, J. New optical molecular imaging systems. Curr. Pharm. Biotechnol. 2010, 11, 620–627.
- Pierce, M.C.; Javier, D.J.; Richards-Kortum, R. Optical contrast agents and imaging systems for detection and diagnosis of cancer. Int. J. Cancer 2008, 123, 1979– 1990.
- Gioux, S.; Choi, H.S.; Frangioni, J.V. Image-guided surgery using invisible near-infrared light: Fundamentals of clinical translation. Mol. Imaging 2010, 9, 237–255.
- Velde, E.A.; Veerman, T.; Subramaniam, V.; Ruers, T. The use of fluorescent dyes and probes in surgical oncology. Eur. J. Surg. Oncol. 2010, 36, 6–15.
- Kobayashi, H.; Ogawa, M.; Alford, R.; Choyke, P.L.; Urano, Y. New strategies for fluorescent probe design in medical diagnostic imaging. Chem. Rev. 2010, 110, 2620–2640.
- Hickson, J. In vivo optical imaging: Preclinical applications and considerations. Urol. Oncol. Semin.Orig. Investi. 2009, 27, 295–297.
- 22. Frangioni, J.V. In vivo near-infrared fluorescence imaging. Curr. Opin. Chem. Biol. 2003, 7,626–634.
- 23. Weissleder, R. A clearer vision for in vivo imaging. Nat. Biotechnol. 2001, 19, 316–317.
- 15. Smith, A.M.; Mancini, M.C.; Nie, S.M. BIOIMAGING: Second window for in vivo imaging. Nat. Nanotechnol. 2009, 4, 710–711.
- Rudin, M.; Weissleder, R. Molecular imaging in drug discovery and development. Nat. Rev. DrugDiscov. 2003, 2, 123–131.
- Hargreaves, R.J. The role of molecular imaging in drug discovery and development. Clin.Pharmacol. Ther. 2008, 83, 349–353.
- Weissleder, R. Scaling down imaging: Molecular mapping of cancer in mice. Nat. Rev. Cancer 2002, 2, 11–18.
- Contag, P.R. Whole-animal cellular and molecular imaging to accelerate drug development. Drug Discov. Today 2002, 7, 555–562.

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*

 $_{\rm Page}14$



- Gross, S.; Piwnica-Worms, D. Molecular imaging strategies for drug discovery and development.Curr. Opin. Chem. Biol. 2006, 10, 334–342.
- Rudin, M. Noninvasive structural, functional, and molecular imaging in drug development. Curr. Opin. Chem. Biol. 2009, 13, 360–371.
- Dufort, S.; Sancey, L.; Wenk, C.; Josserand, V.; Coll, J.L. Optical small animal imaging in the drug discovery process. BBA. Biomembranes 2010, 1798, 2266–2273.
- Sivaraman, D.; Biswas, P.; Cella, L.N.; Yates, M.V.; Chen, W. Detecting RNA viruses in living mammalian cells by fluorescence microscopy. Trends Biotech. 2011, 29, 307–313.
- Koba, W.; Kim, K.; Lipton, M.L.; Jelicks, L.; Das, B.; Herbst, L.; Fine, E. Imaging devices for use in small animals. Semin. Nucl. Med. 2011, 41, 151–165.
- DE Reichert; JS Lewis; CJ Anderson. Coordination Chemistry Reviews. 1999,184, 1, 3-66.
- 35. A Maitra; T Banerjee; A Singh; R Sharma. International Journal of Pharmaceutics, 2005, 289, 1-2, 189-195.
- 36. G Meseguer; R Gurny; P Buri. Journal of Drug Targeting, 1994, 2, 4, 269-288.
- IR Wilding; AJ Coupe; SS Davis. Advanced Drug Delivery Reviews, 2001, 46, 1, 103-124.
- A Parr; M Jay. Pharmaceutical Research, 1987, 4, 6, 524-526.
- GA Digenis; EP Sandefer; RC Page; WJ Doll. Pharmaceutical Science and Technology Today, 1998, 1, 3, 100-107.
- 40. GA Digenis; AF Parr; M Jay. In: Drug delivery to the gastrointestinal Tract, 1st ed., Ellis Horwood, Chichester (UK), 1989; 111-120.
- NK Jain. In: Advances in controlled and novel drug delivery, 1st ed., CBS Publishers & Distributors, New Delhi, India, 2006; 104-109.
- HK Chan. In: Encyclopedia of pharmaceutical technology, Marcel Dekker, New York, 2002; 2365-2371.
- 43. Frangioni JV: In vivo near-infrared fluorescence imaging. Curr Opin Chem Biol 2003, 7: 626-634.
- 44. Ballou B, Ernst LA, Waggoner AS: Fluorescence imaging of tumors in vivo. Curr Med Chem 2005, 12: 795-805.
- 45. Licha K: Contrast agents for optical imaging. Top Curr Chem 2002, 222:1-29.
- Michalet X, Pinaud FF, Bentolila LA, Tsay JM, Doose S, Li JJ, Sundaresan G, Wu AM, Gambhir SS, Weiss S: Quantum dots for live cells, in vivo imaging, and diagnostics. Science 2005, 307:538-544.
- 47. Medintz IL, Uyeda HT, Goldman ER, Mattoussi H: Quantum dot bioconjugates for imaging, labeling and sensing.Nat Mater 2005, 4:435-446.

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

- Gioux, S.; Choi, H.S.; Frangioni, J.V. Image-guided surgery using invisible near-infrared light: Fundamentals of clinical translation. Mol. Imaging 2010, 9, 237–255.
- Schaafsma, B.E.; Mieog, J.S.D.; Hutteman, M.; van der Vorst, J.R.; Kuppen, P.J.K.; Lowik, C.; Frangioni, J.V.; van de Velde, C.J.H.; Vahrmeijer, A.L. The clinical use of indocyanine green as a near-infrared fluorescent contrast agent for image-guided oncologic surgery. J. Surg. Oncol. 2011, 104, 323–332.
- Mishra, A.; Behera, R.K.; Behera, P.K.; Mishra, B.K.; Behera, G.B. Cyanines during the 1990s: A review. Chem.Rev. 2000, 100, 1973–2011.
- Licha, K. Contrast agents for optical imaging. In Contrast Agents Ii; Krause, W., Ed.; Springer-Verlag: Berlin, Germany, 2002; 1–29.
- Kim, D.E.; Jaffer, F.A.; Weissleder, R.; Tung, C.H.; Schellingerhout, D. Near-infrared fluorescent imaging of cerebral thrombi and blood-brain barrier disruption in a mouse model of cerebral venous sinus thrombosis. J. Cerebr. Blood Flow Metabol. 2005, 25, 226–233.
- Liu, X.; Guo, X.L.; Liu, F.; Zhang, Y.; Zhang, H.; Hu, G.S.; Bai, J. Imaging of indocyanine green perfusion in mouse liver with fluorescence diffuse optical tomography. IEEE Trans. Biomed. Eng. 2011, 58, 2139– 2143.
- Herbort, C.P.; LeHoang, P.; Guex-Crosier, Y. Schematic interpretation of indocyanine green angiography in posterior uveitis using a standard angiographic protocol. Ophthalmology 1998, 105, 432–440.
- Achilefu, S.; Dorshow, R.B. Dynamic and continuous monitoring of renal and hepatic functions with exogenous markers. Contrast Agent. II 2002, 222, 31– 72.
- Liebert, A.; Wabnitz, H.; Obrig, H.; Erdmann, R.; Moller, M.; Macdonald, R.; Rinneberg, H.; Villringer, A.; Steinbrink, J. Non-invasive detection of fluorescence from exogenous chromophores in the adult human brain. Neuroimage 2006, 31, 600–608.
- Liebert, A.; Sawosz, P.; Milej, D.; Kacprzak, M.; Weigl, W.; Botwicz, M.; Maczewska, J.; Fronczewska, K.; Mayzner-Zawadzka, E.; Krolicki, L.; Maniewski, R. Assessment of inflow and washout of indocyanine green in the adult human brain by monitoring of diffuse reflectance at large source-detector separation. J. Biomed. Opt. 2011, 16, 046011.
- Yamamoto, M.; Sasaguri, S.; Sato, T. Assessing intraoperative blood flow in cardiovascular surgery. Surg. Today 2011, 41, 1467–1474.
- Raabe, A.; Beck, J.; Gerlach, R.; Zimmermann, M.; Seifert, V. Near-infrared indocyanine green video angiography: A new method for intraoperative

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*

 $_{\rm Page}142$



Available Online through

www.ijpbs.com (or) www.ijpbsonline.com

assessment of vascular flow. Neurosurgery 2003, 52, 132–139.

- Taggart, D.P.; Choudhary, B.; Anastasiadis, K.; Abu-Omar, Y.; Balacumaraswami, L.; Pigott, D.W. Preliminary experience with a novel intraoperative fluorescence imaging technique to evaluate the patency of bypass grafts in total arterial revascularization. Ann. Thorac. Surg. 2003, 75, 870– 873.
- Neri, D.; Carnemolla, B.; Nissim, A.; Leprini, A.; Querze, G.; Balza, E.; Pini, A.; Tarli, L.; Halin, C.; Neri, P.; Zardi, L.; Winter, G. Targeting by affinity-matured recombinant antibody fragments of an angiogenesis associated fibronectin isoform. Nat. Biotechnol. 1997, 15, 1271–1275.
- Birchler, M.; Neri, G.; Tarli, L.; Halin, C.; Viti, F.; Neri, D. Infrared photodetection for the in vivo localisation of phage-derived antibodies directed against angiogenic markers. J. Immunol.Method. 1999, 231, 239–248.
- Bugaj, J.E.; Achilefu, S.; Dorshow, R.B.; Rajagopalan, R. Novel fluorescent contrast agents for optical imaging of in vivo tumors based on a receptor-targeted dyepeptide conjugate platform. J. Biomed.Opt. 2001, 6, 122–133.
- Achilefu, S.; Jimenez, H.N.; Dorshow, R.B.; Bugaj, J.E.; Webb, E.G.; Wilhelm, R.R.; Rajagopalan, R.; Johler, J.; Erion, J.L. Synthesis, in vitro receptor binding, and in vivo evaluation of fluorescein and carbocyanine peptide-based optical contrast agents. J. Med. Chem. 2002, 45, 2003–2015.
- Becker, A.; Hessenius, C.; Licha, K.; Ebert, B.; Sukowski, U.; Semmler, W.; Wiedenmann, B.; Grotzinger, C. Receptor-targeted optical imaging of tumors with near-infrared fluorescent ligands. Nat. Biotechnol. 2001, 19, 327–331.
- Folli, S.; Westermann, P.; Braichotte, D.; Pelegrin, A.; Wagnieres, G.; Vandenbergh, H.; Mach, J.P. Antibodyindocyanin conjugates for immunophotodetection of human squamous-cell carcinoma in nude-mice. Cancer Res. 1994, 54, 2643–2649.
- Ballou, B.; Fisher, G.W.; Waggoner, A.S.; Farkas, D.L.; Reiland, J.M.; Jaffe, R.; Mujumdar, R.B.; Mujumdar, S.R.; Hakala, T.R. Tumor labeling in-vivo using cyanineconjugated monoclonalantibodies. Cancer Immunol. Immunother. 1995, 41, 257–263.
- Ballou, B.; Fisher, G.W.; Hakala, T.R.; Farkas, D.L. Tumor detection and visualization using cyanine fluorochrome-labeled antibodies. Biotechnol. Progr. 1997, 13, 649–658.
- 69. Weissleder, R.; Tung, C.H.; Mahmood, U.; Bogdanov, A. In vivo imaging of tumors with protease-activated

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

near-infrared fluorescent probes. Nat. Biotechnol. 1999, 17, 375–378.

- Bullok, K.E.; Maxwell, D.; Kesarwala, A.H.; Gammon, S.; Prior, J.L. Snow, M.; Stanley, S.; Piwnica-Worms, D. Biochemical and in vivo characterization of a small, membrane-permeant, caspase-activatable far-red fluorescent peptide for imaging apoptosis. Biochemistry 2007, 46, 4055–4065.
- Edgington, L.E.; Berger, A.B.; Blum, G.; Albrow, V.E.; Paulick, M.G.; Lineberry, N.; Bogyo, M. Noninvasive optical imaging of apoptosis by caspase-targeted activity-based probes. Nature Med. 2009, 15, 967–973.
- Bremer, C.; Ntziachristos, V.; Weissleder, R. Opticalbased molecular imaging: Contrast agents and potential medical applications. European Radiol. 2003, 13, 231–243.
- Figueiredo, J.L.; Alencar, H.; Weissleder, R.; Mahmood, U. Near infrared thoracoscopy of tumoral protease activity for improved detection of peripheral lung cancer. Int. J. Cancer 2006, 118, 2672–2677.
- Blum, G.; von Degenfeld, G.; Merchant, M.J.; Blau, H.M.; Bogyo, M. Noninvasive optical imaging of cysteine protease activity using fluorescently quenched activity-based probes. Nat. Chem. Biol. 2007, 3, 668– 677.
- 75. Weissleder, R.; Ntziachristos, V. Shedding light onto live molecular targets. Nature Med. 2003, 9, 123–128.
- Marten, K.; Bremer, C.; Khazaie, K.; Sameni, M.; Sloane, B.; Tung, C.H.; Weissleder, R. Detection of dysplastic intestinal adenomas using enzyme-sensing molecular beacons in mice. Gastroenterology 2002, 122, 406– 414.
- Escobedo, J.O.; Rusin, O.; Lim, S.; Strongin, R.M. NIR dyes for bioimaging applications. Curr. Opin. Chem. Biol. 2010, 14, 64–70.
- Lordi, V.; Yao, N.; Wei, J. Method for supporting platinum on single-walled carbon nanotubes for a selective hydrogenation catalyst. Chem. Mater. 2001, 13, 733–737.
- 79. Huynh, W.U.; Dittmer, J.J.; Alivisatos, A.P. Hybrid nanorod-polymer solar cells. Science 2002, 295, 2425– 2427.
- Baughman, R.H.; Zakhidov, A.A.; de Heer, W.A. Carbon nanotubes—The route toward applications. Science 2002, 297, 787–792.
- Fischer, J.E.; Dai, H.; Thess, A.; Lee, R.; Hanjani, N.M.; Dehaas, D.L.; Smalley, R.E. Metallic resistivity in crystalline ropes of single-wall carbon nanotubes. Phys. Rev. B 1997, 55, R4921–R4924.
- 82. Porti, M.; Blasco, X.; Nafria, M.; Aymerich, X. Electrical characterization and fabrication of SiO2 based metal-oxide-semiconductor nanoelectronic devices with

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*



Available Online through

www.ijpbs.com (or) www.ijpbsonline.com

atomic force microscopy. Nanotechnology 2003, 14, 584–587.

- 83. Roduner, E. Size matters: Why nanomaterials are different. Chem. Soc. Rev. 2006, 35, 583–592.
- 84. Buhro, W.E.; Colvin, V.L. Semiconductor nanocrystals— Shape matters. Nat. Mater. 2003, 2, 138–139.

IJPBS |Volume 5| Issue 2 |APR-JUN|2015|134-144

 Bruchez, M.; Moronne, M.; Gin, P.; Weiss, S.; Alivisatos, A.P. Semiconductor nanocrystals as fluorescent biological labels. Science 1998, 281, 2013– 2016.



 $_{Page}144$

International Journal of Pharmacy and Biological Sciences (e-ISSN: 2230-7605)

Abhishek Bhattacharjee*

www.ijpbs.com or www.ijpbsonline.com