The introduction of brain imaging technology revolutionized the study of psychiatric illnesses and continues to advance our understanding of the biological abnormalities that underpin psychiatric disorders. Aside from ruling out 'organic' brain pathology, neuroimaging modalities do not yet have a major role in the diagnosis of psychiatric disorders; however, neuroimaging has begun to aid in the identification of disease specific intermediate phenotypes (or endophenotypes) which may be useful in aiding diagnosis and in the prediction of treatment response in psychiatric illnesses. The greatest potential of neuroimaging as an approach lies in its combined use with carefully defined psychiatric phenotypes, genotyping, neuropsychopharmacology and post-mortem studies. In this way, neuroimaging can provide a means by which preclinical data can be translated into clinically useful information for application in psychiatry. This editorial is intended to provide a brief overview of the range of imaging modalities and methods currently being applied in psychiatry and includes examples of resulting data.

Thirty years ago the advent of x-ray computed tomography (CT) transformed medical diagnostics. In psychiatry, these first CT studies assessed cerebral and ventricular volume and sulcal size. The most consistent findings from these studies were that patients with schizophrenia and mood disorders demonstrated ventricular enlargement and sulcal widening, albeit to differing degrees. In addition, early MRI studies identified increasing white matter hyperintensities (WMH; bright foci observed on T2 MRI scans thought to represent injury to white matter tracts), with age and their association with cerebrovascular disease, bipolar disorder and late-onset depression. Unlike these early imaging studies, more recent studies have begun to identify disease specific pathophysiological abnormalities.

Structural MRI studies have more commonly examined the differences in grey matter volume in psychiatric populations. Areas of the brain strongly implicated in emotion regulation and cognitive processing have been reported to have reduced grey matter volume during depression including for example, the subgenual anterior cingulate cortex and hippocampus. After grey matter volume has also been identified in schizophrenia in prefrontal and temporal cortical areas and in bipolar disorder in several areas implicated in mood regulation. Structural MR images can also be used to assess cortical thickness, curvature, surface area and white matter volume. First episode schizophrenia has recently been associated with reduced thickness of the paralimbic area of the anterior cingulate cortex concomitant with an increase in the surface area. These newer methods of examining brain structure add significantly to the pool of knowledge that can be gained from structural MR imaging studies in psychiatry and are likely to be more widely used in upcoming studies.

In addition to structural MR-based studies, the MRI scanner can also be used to perform spectroscopy, and diffusion tensor imaging (DTI). The most commonly used form of MR spectroscopy is proton spectroscopy (1H-MRS). 1H-MRS can be used to assess levels of a number of molecules and neurotransmitters in the brain including glutamate, GABA, and markers of neuronal and glial integrity. Reductions in GABA levels in occipital and frontal cortices have been reported during depression, however, it is less clear whether this deficit exists following remission. In addition, the majority of MRS studies during depression have reported increased membrane turnover without neurodegeneration and in contrast to bipolar disorder (BD), a hypoglutamatergic state in major depressive disorder (MDD). DTI is a relatively new and rapidly developing diffusion imaging method which exploits the tendency of water molecules in the axonal fibres of the myelinated or white matter tracts to diffuse preferentially in one direction, a property known as anisotropy. Changes in white matter tract integrity can thereby be detected and disruptions to the tracts are thought to represent altered connectivity between regions. However the microstructural abnormalities underlying diffusion differences are unknown. The majority of DTI studies in schizophrenic samples have detected relatively widespread alteration in brain white matter integrity. The most rapidly developing aspect of DTI currently is the computational modeling of the diffusion signal which may have contributed to inconsistencies between studies observed to date.

The combined discoveries and efforts of chemists, physicists and medical investigators have lead to the development of single photon emission computed tomography (SPECT), and positron emission tomography (PET). These revolutionary new technologies have permitted the direct visualisation of tumours, neuronal degeneration, and localisation of origin points of seizures in epileptic subjects. The first functional neuroimaging studies exploited the ability of PET with radio-labelled water (15O-H2O), and deoxyglucose (18F-FDG) to measure the rate of cerebral blood flow (CBF) and the rate of glucose metabolism, respectively. These represented the first imaging methods of assessing the neurobiological correlates of human behaviour. These functional studies have given us a myriad data, for example implicating reduced blood flow and metabolism during clinical depression in the subgenual...
anterior cingulate cortex. The most common functional imaging method applied presently, however, is functional MRI (fMRI), which affords greater temporal resolution of cerebral activity than PET, for example it can take an entire brain measurement at the time a volunteer is experiencing anticipation of reward and a second entire brain measurement moments later during the subjective experience of reward itself. One of the most novel discoveries made recently using fMRI was of awareness in a vegetative state. A young female subject in a vegetative state as a result of a car accident was asked to imagine playing tennis or moving about her home: the cortical areas activated in response to this instruction were indistinguishable from those activated by healthy volunteers.

Functional imaging studies assessing cerebral blood flow, glucose metabolism and the fMRI signal are global measures representing synaptic activity but in a non-specific manner. A vital step to capitalize on the evidence derived from these methods is to examine what may underpin altered neuronal activity at an anatomical and/or molecular level. The widening array of targets one can assess using molecular PET and SPECT make this possible. Molecular PET / SPECT radioligands have binding characteristics that enable estimation of an outcome parameter that is proportional to the density of the target, such as a receptor or a transporter. This molecular imaging has lead to the discoveries of altered levels of several proteins during depression including the serotonin transporter, the serotonin-1A receptor either reduced dopamine-2 receptor levels or increased dopamine release and reduced cholinergic muscarinic-2 receptors in several brain areas involved in emotion regulation in each case. The meaningfulness of these data are heavily dependent on certain vital sample characteristics including homogeneity of mood state, medication-free status, diagnostic subtype and the inclusion/exclusion of comorbid states. Controlling for these factors can lead to the detection of molecular differences between phenotypically similar disorders. The level of the serotonin transporter in the brainstem of depressed unmedicated individuals with MDD differs from that in depressed unmedicated individuals with BD. This could conceivably underlie the observed difference in antidepressant efficacy of serotonin reuptake inhibitors in MDD, which is approximately 75%, relative to that in BD, about 35%. Evidence supporting this has recently emerged. A better response to the serotonin reuptake inhibitor, citalopram in the STAR-D study was associated with a polymorphism in the gene that codes for the 5-HT2A receptor and this same polymorphism was associated with having greater levels of the serotonin transporter binding in the above mentioned depressed sample (pending publication). The extension of these types of preliminary data to larger samples may help to identify endophenotypes or a set of characteristics that indicate the likelihood that a subgroup of individuals possessing such characteristics will respond preferentially to a given treatment.

In molecular PET and SPECT imaging, certain radioligands are sensitive to displacement by the endogenous neurotransmitter. This permits the measurement of neurotransmitter release, such as in the case of C-11 labelled raclopride which can be used to assess dopamine release. Martin-Soelch et al (2006) have recently shown that a natural reward task involving gambling with no pharmacological intervention, results in adequate dopamine release to measurably displace raclopride binding. These data demonstrate that naturalistic paradigms may be combined with imaging to further extend our understanding of the biochemical alterations associated with addiction or anhedonia. (These data will be presented at the ‘Neuroscience Ireland’ meeting in August 2008 in NUI Galway.) Unfortunately there is no molecular radioligand sensitive to displacement by serotonin currently available for use in humans. One can additionally measure target occupancy using PET which has proven useful in determining the level of serotonin transporter occupancy necessary to achieve clinical efficacy. These data are subsequently helpful in the early elimination of agents which have inappropriate levels of side effects in animals when administered at the level of occupancy necessary to achieve an antidepressant response.

To date, the discovery of therapeutic agents in psychiatry has been somewhat serendipitous such as the mood stabilizing effects of lithium and the anticonvulsants in the treatment of bipolar disorder. It is critical that novel therapeutic targets are identified and that further investigations into the prediction of treatment response be carried out. Many individuals have an unsatisfactory response to existing treatment options and/or become disheartened with multiple pharmacological trials. The majority of imaging studies aimed at identifying predictors of treatment response to date have focused on MDD and obsessive compulsive disorder (OCD). These studies have most commonly compared glucose metabolism assessed using FDG-PET in response to task performance before and after treatment response (for example see ref 27). Although these data are of limited clinical utility currently, they represent some of the first steps toward the development of neuroimaging as a clinical tool in psychiatry. The potential for clinical application of PET is illustrated further by a recently developed PET radioligand, Pittsburgh Compound-B (PiB). This radioligand binds amyloid and therefore serves as a means to monitor progression of Alzheimer’s disease, and consequently treatment performance in retarding disease progression.

It is unlikely that preclinical studies alone, such as animal models and probing second messenger systems, will lead to a comprehensive understanding of the pathophysiology of psychiatric illnesses. However, these studies critically provide the evidence necessary to design translational studies. Neuroimaging provides a means by which such preclinical data can be translated into clinically applicable information. The major advantages of the neuroimaging approach are that it can be carried out in a relatively non-invasive manner, can obtain data throughout the entire brain, and that these data can be obtained during subjective experiences in human volunteers. The resolution of PET and SPECT and the number of molecular targets that can be assessed, while not ideal has improved greatly, but these do not match that of a post-mortem approach. On the other hand, the time consuming post-mortem approach can select only a small area of the brain to assess a given molecular target. Thus it appears optimal to combine the use of neuroimaging tools and post-mortem approaches to identify molecular abnormalities associated with the disease state. In this way, the molecular pathology underlying abnormalities identified using molucu-
lar neuroimaging can be examined further. For instance, functional imaging studies have strongly implicated the amygdala in anxiety and depression and subsequent post-mortem histological studies identified reductions in glial cells with no change in neuronal cell numbers in MDD. Similarly, reduced grey matter volume of the hippocampus during depression detected using MRI, has been informed by post-mortem studies revealing changes in the mRNA levels of factors important to neuronal cell growth and survival. In combination with genetic information and functional correlates these combined approaches represent a powerful way to biologically characterise the complex array of contributors to a disease state in psychiatry.

There is currently a burgeoning degree of neuroimaging research in Ireland. Trinity College Institute of Neuroscience (TCIN, www.tcd.ie/Neuroscience/) operates a research dedicated MRI scanner for a range of functional MRI studies. In Galway, our Clinical Neuroimaging Lab at NUIG (www.nuigalway.ie/psychiatry/research/neuroimaging_lab/) is involved in structural MRI studies, MR spectroscopy, and is developing DTI based research in psychiatric populations. A research dedicated MRI scanner is planned at St. James’ Hospital. Researchers in Beaumont Hospital, St. Vincent’s University Hospital and the AMNCH in Tallaght are also engaged in neuroimaging studies.

The neuroimaging community in Ireland is expanding expertise in a growing number of imaging modalities. In addition to adult psychiatry these are currently being applied to disciplines including cognitive psychology, neurology, pediatric, and old-age psychiatry through a number of productive national and international collaborations. Structural MR studies of individuals experiencing their first episode of psychosis or with euthymic bipolar disorder are underway at NUI Galway. These studies additionally aim to examine the genetic contribution to morphometric abnormalities in collaboration with Prof. Michael Gill in Trinity College Dublin. Molecular PET imaging collaborations with the Drs. William Theodore and Wayne Drevets at NIH in the US, involve assessing serotonin transporter levels in subjects with temporal lobe epilepsy. Kinetic modeling of the PET data to produce images of serotonin transporter binding is performed by the Clinical Neuroimaging Lab at NUI Galway.

The data described in this editorial are intended to convey that neuroimaging can provide a critical methodological bridge between preclinical and clinical investigations, and to identify biologic targets for the quantitative assessment of treatment response and of variables influencing illness progression. The Irish neuroimaging community are well positioned presently, to advance the understanding of the biological basis of clinical disorders including psychiatric illnesses through initiating and funding multidisciplinary studies in translational medicine.
In obsessive-compulsive disorder the dose is 20–240 mg daily in divided doses. The maximum daily dose is 80 mg. Deliberately reduced or alternate-day dosing is recommended for patients who may be able to reduce the dose. Do not use with MAOIs: fluoxetine may cause serious reactions when used in combination with monoamine oxidase inhibitors (MAOIs). The use of irreversible MAOIs is recommended for patients who have started therapy before starting fluoxetine. If fluoxetine is taken with MAOIs, it should be discontinued and the MAOIs should be discontinued at least 5 weeks before fluoxetine is started. In overdose, symptomatic treatment is recommended.

Undesirable effects: Most common effects are nausea, diarrhoea, constipation, and changes of taste. Headache, dizziness, impotence, reduced libido, chest pain, and hot flushes may also occur. Uncommon effects include the development of hives or angioedema, fever, or unexplained abdominal pain. Rare effects include fever, convulsions, and changes of taste. Headache, dizziness, impotence, reduced libido, chest pain, and hot flushes may also occur. Uncommon effects include the development of hives or angioedema, fever, or unexplained abdominal pain.

PROZAMEL is the Clonmel Brand of Fluoxetine.