Dr. Robert Innis Interview

Claudia Wassman: This is Claudia Wassman and today’s date is Monday, March 3, 2005. I’m conducting an interview with Dr. Robert Innis.

[break in audio]

CW: You became chief a new laboratory at NIMH with an emphasis on brain imaging using PET in 2001. That is correct. So maybe you could begin with telling me what brought you here.

Robert Innis: The -- I do brain imaging particularly with PET, Positron Emission Tomography. The things that attracted me here were largely the resources that were available to do this state-of-the-art brain imaging because it’s a very expensive methodology requiring extensive infrastructure like cyclotrons and cameras and a multi-disciplinary research team.

CW: So maybe you can tell a little bit more about the state of the art facilities that you have available here. Be more--

RI: At the NIH, four PET, Positron Emission Tomography, you must first be able to make a radioactive atom or a radionuclide and that’s done in a cyclotron. And at NIH, in the Clinical Center they have three cyclotrons which is three times as many as most places have. Then, after the cyclotron makes this radioactive atom, it’s a Carbon-11 CO2, carbon 11 labeled radioactive carbon -- Carbon Dioxide, it has to be quickly, that radioactive carbon has to be quickly incorporated synthetically with chemistry into some radio labeled probe that will go to the brain and label some specific protein in the brain or someplace in the body as a whole. In addition to having this cyclotron there has to be an extensive radiochemistry facilities in order to make the radio label probe that is subsequently injected into an animal or human subject. NIMH in particular was very interested -- the National Institute of Mental Health was very interested in this methodology because it allows a way to look at proteins that may be involved in normal brain function as well as how, what goes wrong in the brains of people with neuro-psychiatric disorders. So if you want to try to measure these proteins, this is one of the major ways, if not the only way in order to be able to do that in a living human brain.
Because NIMH was interested they were willing to make a large investment in a new radiochemistry laboratory here. Then after you make it -- after you make it! After you make the radio labeled probe, the radio tracer, the radioligand which binds to a specific protein in the brain, then you have to inject it into an animal or human and do imaging. The imaging facilities here for PET are really extensive. They currently they have two cameras just for imaging in rats, they have 1-2-3-4, four to five cameras for imaging in monkeys and humans. So it’s really an extensive facilities and also the experience -- personnel who are multi-disciplinary and skilled in many different areas.

CW: Your focus, one of the things you mentioned, is proteins. Is this the most promising approach to brain imaging?

RI: For measuring specific proteins in the living human brain it may be the only one right now really, for the mast majority. And it’s an issue of sensitivity. Many of the proteins that we’re interested in, that we think may be involved in psychiatric illnesses or neurological illnesses, many of those proteins are present in very low concentration and in order to be able to measure them we need a very highly sensitive method to be able to do so. PET is currently unique in the imaging methodologies for doing that. If you want some specific numbers, I would say that if you compare MRI methods, magnetic resonance imaging as a whole, their sensitivity, they can measure lets say the targets that are maybe present at 10^-4 molar. But PET can probably go down 10^-12 molar. So it’s just many, many orders of magnitude higher sensitivity, and many of the proteins that we’re interested in measuring are present at 10^-9, 10^-10 molar. So it is out of bounds for being possible to measure, at least currently, with MRI techniques and we need some other higher sensitivity method. The downside is that you have to use radioactivity, but the advantage is you have much greater sensitivity.

CW: Which tracers do you focus on?

RI: The purpose this lab is to develop new tracers, so because of the unique sensitivity of the PET, and also I might say because of the specificity -- you can make drugs that will bind to just one specific protein. So because you can measure specific proteins at low concentrations in vivo, this seems like a very valuable technique that could be. So NIMH put in a lot of money in order to make this radiochemistry lab and the purpose of this lab is to develop many new ligands because it’s -- it has great potential but there has been relatively little payoff so far. As we develop the new ligands, the idea is to make the knowledge about them available to other PET centers throughout the United States. We are currently working on perhaps twenty different targets at various stages of work, in terms of cold chemistry to radio labeling to imagining in animals going on to humans. And to some patient disorders. It’s a broad number, we’re working on about twenty different targets right now -- target proteins.

CW: So you say it has a great potential but little payoff so far. What makes it so difficult?
RI:  It’s -- this is developing a radio labeled drug so it’s called a radio pharmaceutical. Everyone knows, most people, many people know that pharmaceutical development is very risky. Less than one out of a hundred compounds turn out to work. Well it’s a similar success rate, maybe a little, it’s a better success rate than that, but it is a low success rate because you have to make a radio pharmaceutical that is going to work and the characteristics of that are different from a therapeutic agent. Therapeutic agents, many of those don’t work out. So it’s an inherently risky business and difficult in order to make it.

The other is, I mean there has been some successes -- I would say the difficulties are the high failure rate in developing pharmaceuticals, including radio pharmaceuticals, and also the technically complex nature of it. Few places have cyclotrons available and radio chemistry facilities, but that is expanding. I think one of the reasons why this is a good time to get into it is that PET is now beginning to expand in a manner that is very similar to MRI in, let’s say, the ‘80s. In the past few years insurance agents, including the government’s Medicare, has started to reimburse for selected PET scans, and these in particular are PET scans to measure metabolism with FTG, a glucose analog. And many tumors and their metastases will have increased glucose uptake so that you can localize the tumor and its metastases with this PET tracer. So it’s been re-approved, it’s been approved by Medicare for reimbursement and money drives the capitalist medical system in the United States. Not only that, but the charge for the FTG scan is pretty high, so that you can make a profit, so that cash starved hospitals say, “Profit? Well, we’ll set up a PET too.” Companies where they set up a PET camera, this two million dollar camera, in a truck and bring it around from hospital to hospital like a mammogram for people to have their scans.

As medial centers get PET cameras and get set up to do it, then they have a significant component of the infrastructure there in order to be able to do the research studies. And the same time happened with MRI where it was limited utility, and when there was reimbursement for it, every hospital gets an MRI. While the MRI is there you have the infrastructure and you can work on research, and functional MRI is sort of invented, and then the infrastructure was there and you could do it. So it seems like it could expand now.

CW:  Yeah. So you say have about twenty different tracers that you are working on. Is there one that you think is most promising or maybe most specific?

RI:  Well they’re all specific. These are twenty different targets, and for any target we might have several ligands. There are many targets that may have potential clinical utility. I can give you an example of one that I’ve worked on in the past. When I was previously at Yale University, I helped to make a probe for a particular target that’s a marker in Parkinson’s Disease. That is now, is used in the European Union to aid in the diagnosis of Parkinson’s Disease. Specifically we radio labeled a chemical analog of cocaine, and it
binds to the cocaine receptor and it just so happens that patients with Parkinson’s Disease lack the cocaine receptor. With this radio labeled probe, or with the absence of that, you can aid in the diagnosis of Parkinson’s disease. There has been a lot of work now done on, in the United States, on a probe for amyloid, a protein which builds up in Alzheimer’s disease. Two other research groups have developed probes that could measure this protein and may be able to diagnosis Alzheimer’s disease before the onset of symptoms.

CW: When you chose these tracers, is your research driven by a specific disease? You mentioned Parkinson’s, you mentioned Alzheimer’s, or what is it that makes you go into a specific direction?

RI: If we’re fortunate enough to know a very clear connection to a disease then that can be one of the reasons for going into it. So in the case of Alzheimer’s disease, we really do need to know the dopamine neurons that are gone. So we are trying to make a new and improved Amyloid probe and we have other animal and human studies looking at Parkinson’s disease. But unfortunately in the majority of cases we don’t have a clear connection like that. Because it isn’t like, let’s say for schizophrenia, that we have one target that we know that if we could measure that, would be sure to get valuable information for the diagnosis or treatment of schizophrenia. That’s not known. So for the vast majority of the targets these are proteins for which animal or limited human studies may suggest, may be involved in the passive physiology. So the vast majority wouldn’t be “diagnostic agents”, like in Parkinson’s or Alzheimer’s, but really research tools to be able to explore pathophysiology. So these are more like tools to be used to understand what may be going wrong, to look at the time course, to look at the effect of treatment, etcetera. But it would be judged, we only work on a tracer if we think it would extend to humans, and we would only do it if there is some reasonable evidence that that target could be involved in an important way in physiology of pathophysiology.

CW: One receptor you focus on is dopamine T2, and one other is nicotine, that’s a 2-coline receptor and specific areas in the brain that you target in the brain are the thalamus and the temporal cortex. But for the moment you say it’s too early to tell which one would work out and which one wouldn’t. Is that right?

RI: Well maybe we could take an example of the nicotinic receptor. This is the receptor in the brain that nicotine binds to, it’s thought to be the receptor that mediates the addictive proprieties of smoking. There could be reasons to look at that in terms of addiction, let’s say quite clearly with regard to nicotine addiction, but there could be other utility too in looking at cognitive function and agents which enhance nicotine receptor function may enhance cognition and that could be helpful in schizophrenia or Alzheimer’s disease, and in Parkinson’s disease. There are some drugs that act at this nicotine receptor now which enhance cognition.
We worked to develop a probe of the nicotine receptor, and then we’ve looked at it recently in Parkinson’s disease and there is a substantial loss in Parkinson’s disease. We might want to follow that up now to look at Parkinson’s disease with or without dementia. So it would be sort of studying, it’s a research tool to study pathophysiology. We have a good probe for that receptor; now there’d be questions on its use as a clinical research tool. For that we always collaborate with a clinical group that’s a specialist in Parkinson’s disease, or cognitive function in Parkinson’s disease.

CW: So maybe you can -- since much of your work involves animal studies, maybe you can say a little bit about what you’re doing at the NIH right now [inaudible]?

RI: Okay, I’ll give you an example. We’re collaborating with Ron McKay, in the Neurology Institute, who studies embryonic stem cell transplantation. Stem cells might be a useful treatment, they say, of Parkinson’s, Alzheimer’s, diabetes. One of the issues that will come up is that if some stem cells are developed for the treatment, we’ll have to have some way of monitoring whether the stem cells have survived and whether they’re growing and doing what they’re supposed to do.

So, in a project in collaboration with Ron McKay, we looked at -- we did the imaging; Ron McKay and his group did all the molecular biology and interesting stuff. We’re sort of the toolmaker -- we’re the maker of the biomarker. But he has shown that you can poison a rat and give it Parkinson’s disease in one half of its brain so it is hemi-Parkinsonian and it will have abnormalities. Then he treats the animals with embryonic stem cells which can be genetically modified so a lot of them turn into the dopamine neurons, the neurons that are destroyed in Parkinson’s disease, and when he does this he -- the animals get better. So this is a great animal model of Parkinson’s disease and it could be a good one, if it were ever extended to human subjects, to try to develop a marker to see if you can measure the survival and growth of that transplant and we’ve done that. So we can do head imaging of a marker on the dopamine neuron, and we can monitor the survival and growth, and that’s correlated with behavioral improvement in those animals. So that would be an example of an animal study in which we’re demonstrating the feasibility of this as a biomarker.

CW: What other examples are there?

RI: We’re doing a lot of different studies. Here’s -- we’ve done a study in collaboration with the Alcohol Institute to look at monkeys that have had maternal deprivation. The monkeys that have maternal deprivation -- they’re raised by peers, they’re not raised by their mother -- when they grow up have abnormalities; they don’t socialize, they’re very shy, they don’t explore, when exposed to alcohol they will have bouts of alcoholism. So, the analogy of all this to the human situation is very clear and it could well be that that maternal
deprivation is doing something to the brain which then continues into adulthood to lead to these behaviors. That could be responsible for alcoholism, let’s say shyness and withdrawal, and there was evidence that the investigators, Drs. Higley and Sumi [spelled phonetically] had, that there’s a decreased amount of one particular transmitter in the brain, serotonin -- measured if they measured in cerebral spinal fluid, the fluid around the brain.

So we did a study in these animals and looked at regularly reared animals and maternally deprived, and sure enough we had a marker for the serotonin neurons; it was actually the serotonin transporter. It’s the target site where Prozac works, so it’s the “Prozac Receptor” and it was markedly depleted in many areas of the brain. So one could think about some interventions which are being thought about now; maybe they could be treated to enhance that and that would reverse those problems. Maybe that could be a model for disorders in humans; it’s also thought to be a potential model in alcoholism, so we are now doing a study of the same Prozac Receptor in alcoholic subjects. So there’s a direct connection to look and see if we see similar things in the human disorder of alcoholism.

By the way, our results are coming back exactly the opposite. It’s really quite strange. The alcoholics are markedly increased in their level of the serotonin transporters whereas these maternally deprived monkeys are in the other direction, so [laughs] we are left with more questions than answers.

CW: So you were really surprised by your results!

RI: Yeah, yeah.

CW: That’s fascinating. So do you have an idea why that would be?

RI: No. It’s going to take some more work, but what’s nice is that you could think about additional studies that could be done in both animals and humans to try to explore that. For example, in those maternally deprived monkeys they’re also very aggressive. They’re shy but aggressive. So, we’re going to look now at alcoholics at those alcoholics with aggression -- some have markedly increased aggression compared to others, and we’ll look at some other parameters. That may be one aspect of it that we would have to look at, and that could be refined or done in both humans and animals. So that’s another great, great advantage about the work here. Beyond any technical facility resources, the capability of doing bedside -- bench-to-bedside work with very, very outstanding researchers in those areas to examine these questions in a broad way; it’s a big positive at NIH.

CW: So if you say just the two years [?] -- the monkeys are shy or aggressive, so are there specific regions of the brain that you are also alerted to that are active?

RI: So one would think that, yes, but the changes that I mentioned were actually broad over all regions of the brain. There are other
things besides the serotonin transporter that are involved in aggression.

CW: So I think now you have mentioned -- you have mentioned Parkinson’s, Alzheimer’s, schizophrenia, addiction. Is there another field, another study, that would be worth mentioning?

RI: Anxiety disorders and depression are certainly active areas of research here. And again, some of those cases -- in the case of anxiety disorders and depression we do not know like a single protein that we definitely want to measure in order to know that we’re going to have diagnostic information. It isn’t like measuring amyloid in Alzheimer’s disease, or measuring dopamine neurons in Parkinson’s disease. So it is developing tools that we think could be useful in understanding the pathophysiology or the treatment of these disorders, and we have several markers in development or early studies on that.

I guess one that I might mention here is there’s strong evidence in the case of depression that the mechanism action of many antidepressants from drugs to electroconvulsive therapy, that they all work by a common mechanism increasing a second messenger system in the brain -- increasing the cyclic AMP messenger system in the brain. So we have implemented a probe here that will measure cyclic AMP phosphodiesterase which will measure a particular aspect of this second messenger system. So in addition to measuring receptors that are located on the membranes we are trying to move beyond that to second messenger systems. We’ve implemented that tracer here, we’ve developed methods to appropriately quantitate it in rats, and we hope this year to start studies in patients with depression. But again, it would be -- just because we think that it would be involved in either the pathophysiology or the treatment of depression.

CW: Then I would like to know why is it so difficult to measure dopamine transmission in the living human brain.

RI: Well, there are different aspects of it that are -- what you want to be able to do to measure dopamine transmission -- dopamine is a chemical neurotransmitter in the brain -- is that of the various aspects of how that system is mediated, dopamine is a chemical transmitter, it’s released from one neuron, diffuses, travels across a gap, and then interacts with a post-synaptic receptor, like a D2 receptor, and then after that there might be a second messenger system on cyclic AMP dependent systems. So you’d want to measure the synthesis of dopamine, you’d want to measure the release of dopamine, you’re going to measure its interaction with the receptor, and you’d want to measure what effects binding to that receptor has. So you have to develop in vivo, just like has been developed in vitro, probes all along the way in order to be able to do that.

There are probes already developed to measure synthesis, and there are probes available to measure some of the receptors like the D1 and the D2 receptor. There are some novel ways of trying to measure the amount of dopamine released, which I might talk about more -- you might be
interested -- and then the dopamine signal is terminated, is ended, by
the dopamine being taken back up into the first neuron, and that’s done
via the dopamine transporter which is also the cocaine receptor. So
you have markers on the first neuron pre-synaptically and on the seoncd
neuron post-synaptically. And all of these aspects could be of
interest or be abnormal in one area or another of the brain.

An example of the dopamine released can be valuable here. When
amphetamine, or speed, the way speed works is to release dopamine. It
causes dopamine to be released into the synapse and that release of
dopamine causes the euphoria and the high and probably is related to
the addictive properties of amphetamine and some other stimulants like
that. So what we’ve done is to -- and others -- is to label the D2
receptor that is post-synaptic that is on the second neuron -- when
dopamine comes out it binds to that -- we put in a radio tracer that
also binds there so we can measure this D2 receptor before and after we
give amphetamine. We’ve done a series of studies in monkeys and in
humans and in patients. And when the amphetamine causes the dopamine
to be released it displaces the radio-ligand, it knocks it off the
receptor, so therefore the binding goes down, and the amount that the
binding goes is proportional to how much dopamine is released.

Well there’s one of the leading hypotheses of schizophrenia is that the
psychotic symptoms -- hallucinations, delusions -- are caused by
increased dopamine release in the brain. So in order to test this we
did this D2 receptor imaging before and after giving amphetamine and
we’re able to show that patients with schizophrenia had a much greater
release of amphetamine than healthy subjects. In addition, in healthy
subjects the amount of dopamine released was related to the high. They
got higher if they had more dopamine -- even though they had the same
dose of amphetamine they had different amounts released and different
effect -- the higher they got by certain measures or activation, the
more dopamine that was released. In the schizophrenic patients there
can be a transient -- short lasting, one hour, two hour -- increase in
psychotic systems, paranoid symptoms typically, and the transient
increase in psychotic symptoms was related to the amount of dopamine
released. So that was really some of the first evidence clearly
linking dopamine release in the brain to psychotic symptoms in
patients. So that was able to help explore the pathophysiology.

Now, in addition, I mean probably some of the greatest disability that
patients with schizophrenia have is not just the psychotic symptoms but
other symptoms called negative symptoms of withdrawal -- social apathy,
lack of initiative, lack of emotional responsiveness, and also
cognitive impairment, sometimes full-blown dementia -- and that these
cognitive deficits may impair, overall, life even greater than the
psychotic symptoms. So there’s a lot of interest in looking at why is
there this cognitive decline in patients with schizophrenia, and one of
the leading causes is having to do with dopamine transmission in the
cortex. So we and others are trying to develop probes to measure this
in the cortex, not just in the inner portion of the brain called the
stratum, and we have some studies to look at that now, but it’s
difficult to do in general because there’s much less dopamine
transmission in the cortex quantitatively, let’s say, than in the strata, so you have very low density sites that you’re measure and it becomes important to develop probes to be able to measure dopamine transmission in the cortex as well as the sub cortical areas, and that’s a need for a new tool so I’m a toolmaker!

[laughter]
And toolmakers have their roles.

CW: Yeah. [laughs] That’s great. Okay. So these technologies that are available to look at the brain, so far have they already to get a different understanding of emotion and cognition or is this still something that’s in the future?

RI: I think vis-à-vis -- I think there’s much more in the future than there’s been in the past, at least I hope so. It’s just the tip of the iceberg. The -- and there are other imaging methodologies that may be better for some of the questions that you have with regard to cognition and emotion like functional Magnetic Resonance Imaging, fMRI, which has much -- you can do things more quickly and you have no radiation exposure and you have higher resolution, and you can do these studies repeatedly so that fMRI has been far more valuable in, so far, really, than PET, in studying emotion and cognition. But in terms of trying to look at a specific protein I think there you’re at with PET, and insofar as the drugs that are likely to be developed for neuropsychiatric disorders will be targeting specific proteins, there can be extra utility for this sort of measure for the therapeutic development.

The drug companies recognize that and most of the big drug companies have either set up PET within their own facilities or are collaborating with other sites. We have several collaborations ourselves with big pharma because they want to develop the radio labeled compounds that will help them -- when they make the therapeutic agent it helps them in the therapeutic drug development, and we work with them to make the radio labeled probe because it can be a useful research tool to study pathophysiology. So that so-called public/private partnership is strongly encouraged and has been very beneficial to us.

CW: Okay. Well, thank you very much.

RI: You’re welcome!

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