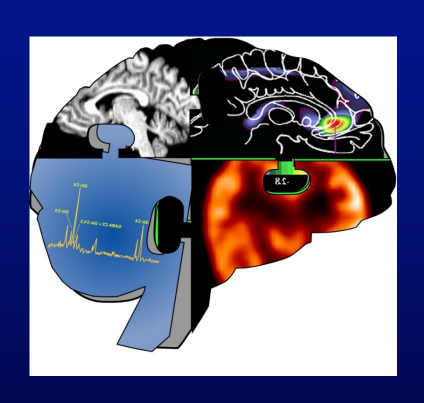
Positron Emission Tomography: Tool to Study Pharmacokinetics and to Facilitate Drug Development



Robert B. Innis, MD, PhD
Molecular Imaging Branch
National Institute Mental Health

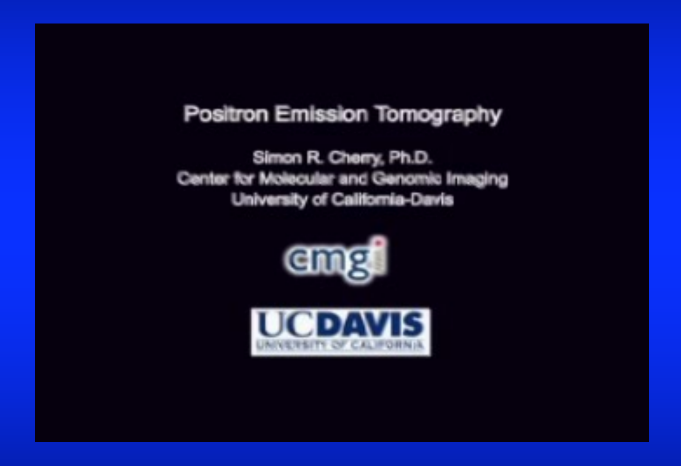
Outline of Talk

- 1. PET has high sensitivity and specificity
- 2. PET used in therapeutic drug development
- 3. Pharmacokinetic modeling of plasma concentration and tissue uptake can measure receptor density
- 4. Study drug distribution: block distribution to periphery and increase distribution to brain
- 5. Study drug metabolism: inhibit defluorination

Imaging Receptors with PET



Positron Emission Tomography



PET vs. MRI

	PET	MRI
Spatial Resolution	2 – 6 mm	<< 1 mm
Sensitivity	10 ⁻¹² M	10 ⁻⁴ M
Temporal Resolution	minutes	<1 sec

Radionuclide (11C): high sensitivity
Ligand (raclopride): high selectivity
Radioligand [11C]raclopride: high sensitivity
& selectivity

Radioligand = Drug + Radioactivity

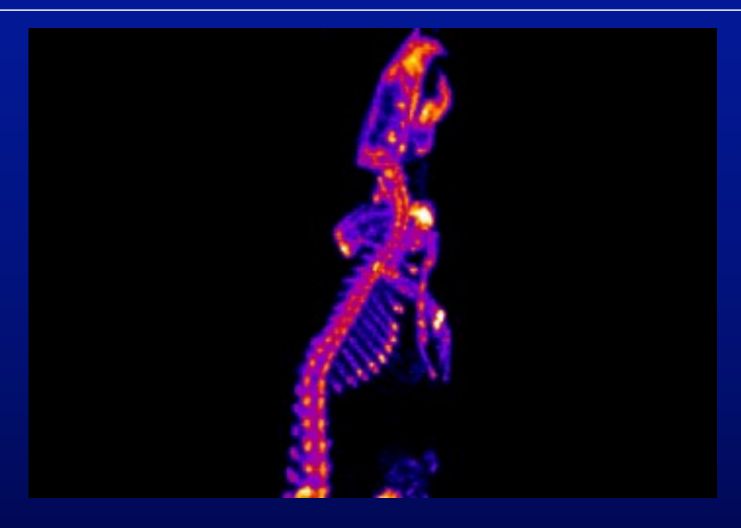
1. Drug administered at tracer doses

- a) No pharm effects
- b) Labels <1% receptors
- c) Labeled subset reflects entire population

2. Radioligand disposed like all drugs

- a) Metabolism & distribution
- 3. Radiation exposure

NIH Rodent PET Camera 18F bone uptake rat

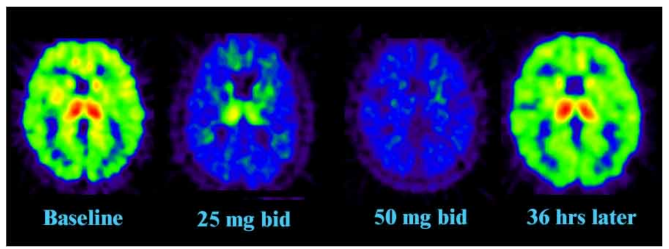


Developed By: Mike Green & Jurgen Seidel

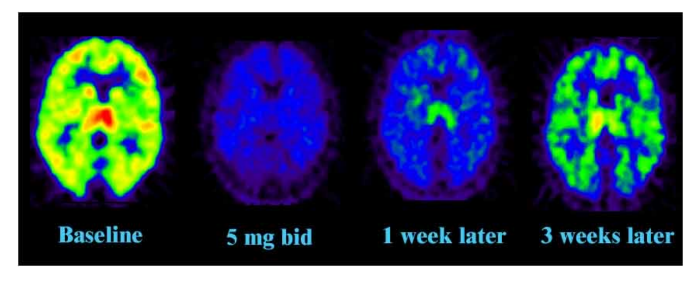
PET: Tool in Therapeutic Drug Development

- Determine dose and dosing interval
- Identify homogeneous group
- Biomarker for drug efficacy
- Monitor gene or stem cell therapy

Lazabemide blocks [11C]deprenyl binding to monoamine-oxidase-B (MAO-B)



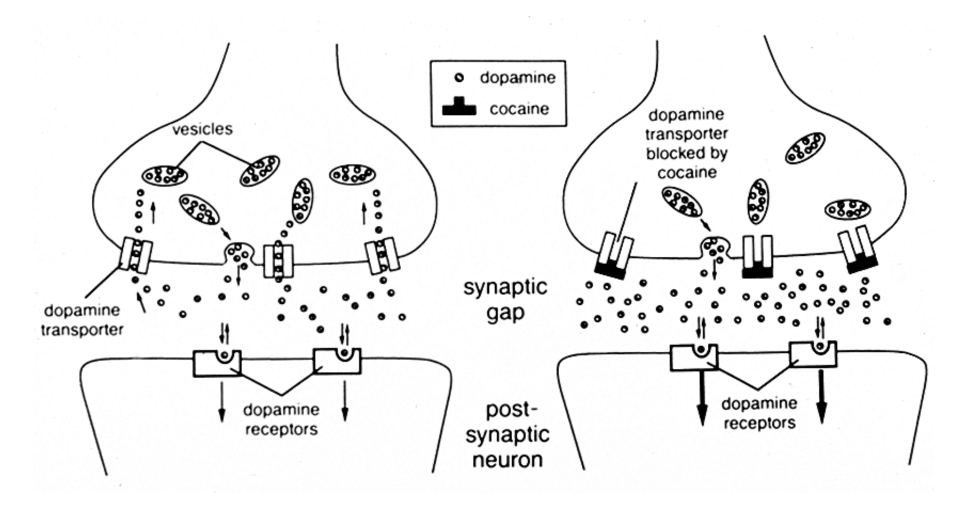
Selegilene is more potent and longer acting than lazabemide



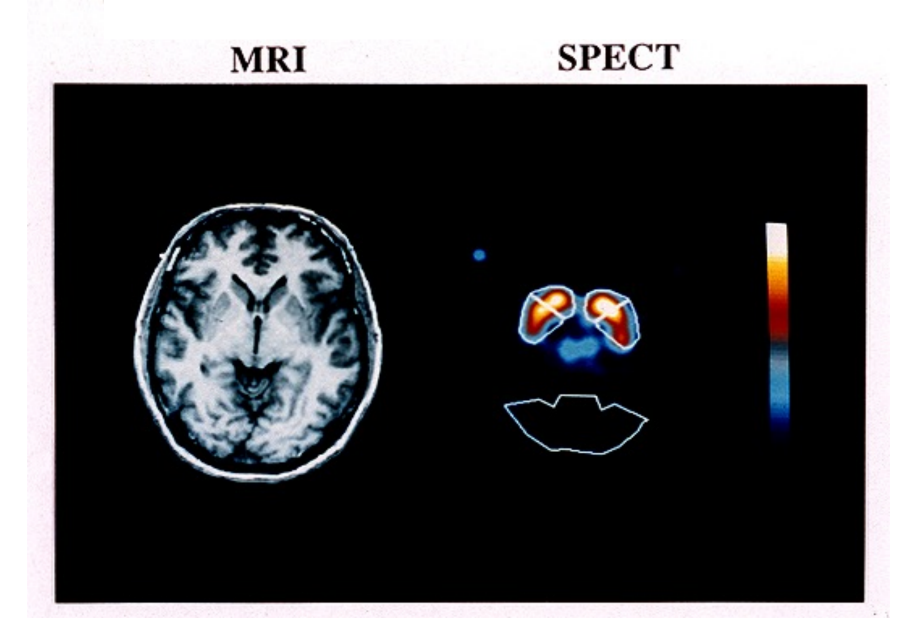
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Dopamine Transporter: Located on DA Terminals Removes DA from Synapse

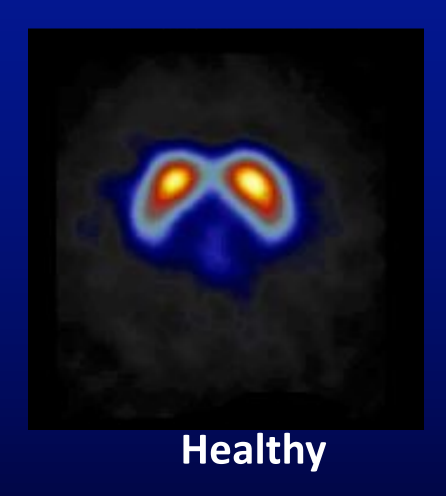


SPECT Imaging of Dopamine Transporter in Caudate and Putamen of Human Brain



Dopamine Transporter SPECT in Parkinson's Disease:

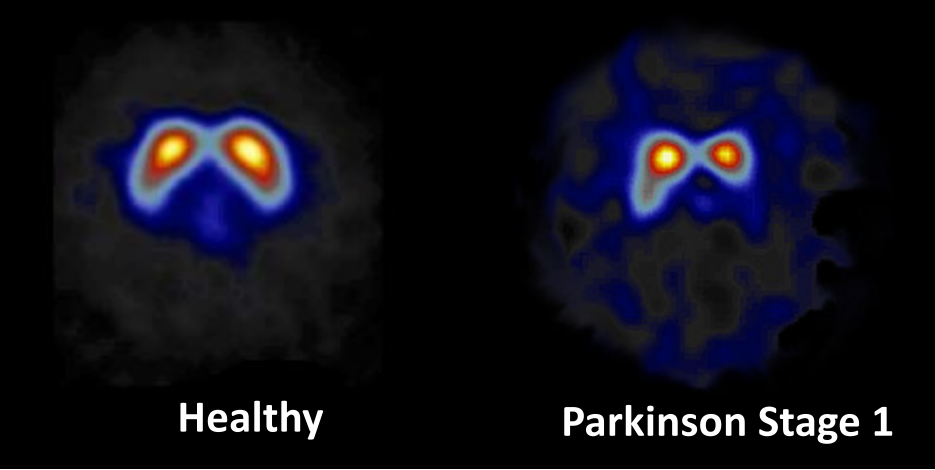
Decreased, asymmetrical, loss in putamen > caudate





Dopamine Transporter SPECT in Parkinson's Disease:

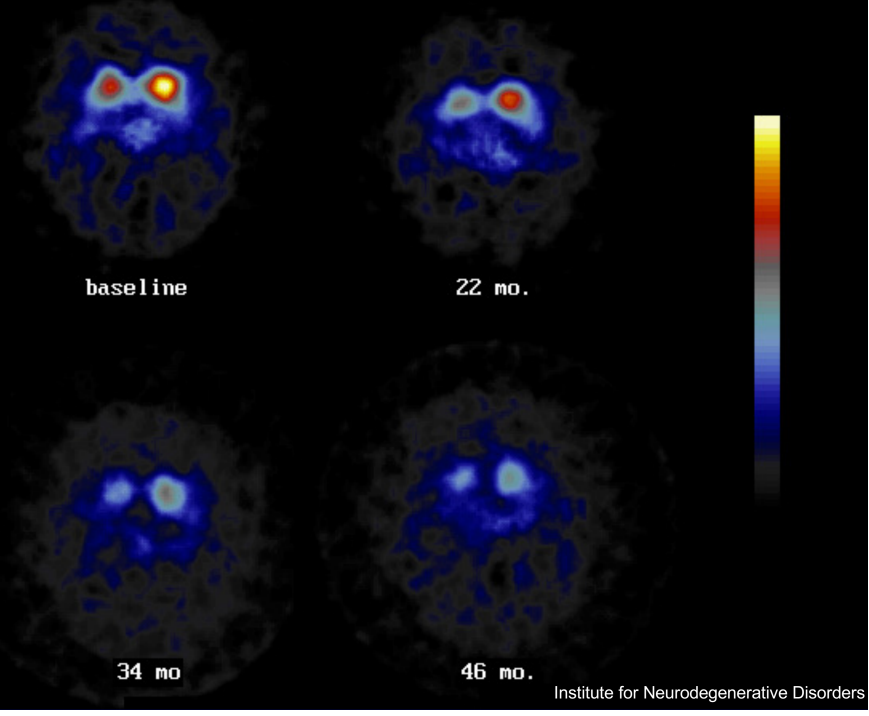
Decreased, asymmetrical, loss in putamen > caudate



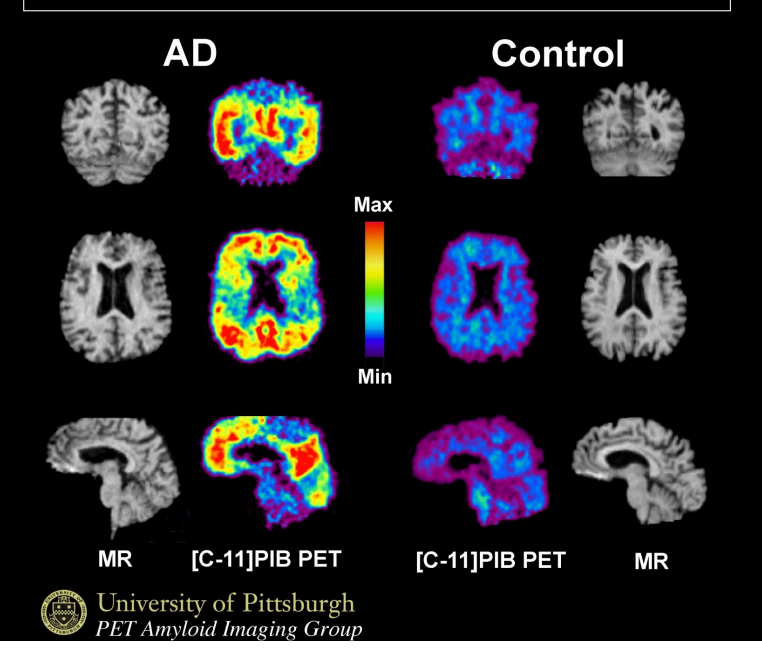
PET: Tool in Therapeutic Drug Development

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Serial Dopamine Transporter Imaging in a Parkinson Patient



PET Imaging of Amyloid: Biomarker for Alzheimer's Disease



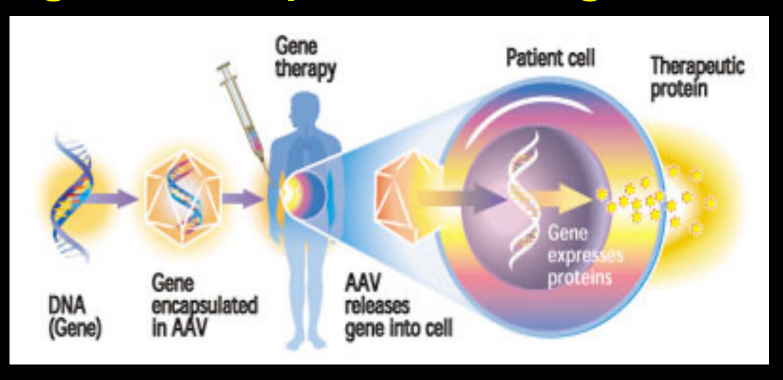
PET: Tool in Therapeutic Drug Development

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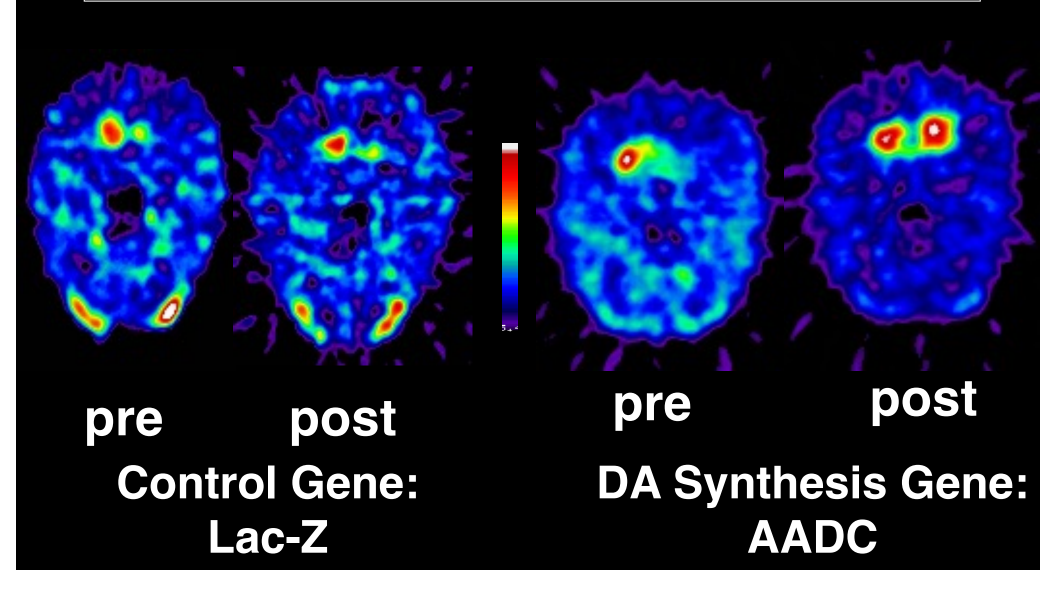
Gene Therapy Using Viral Vectors

Viral vectors deliver gene that synthesizes dopamine (DA) Infuse virus into striatum (target cells)

Target cells express the DA gene

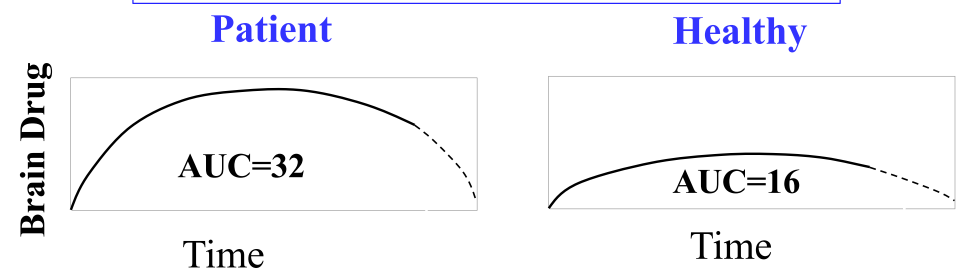


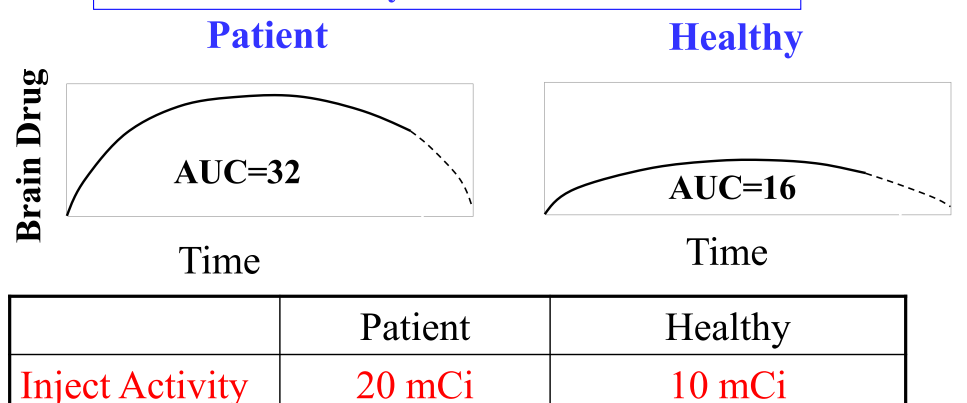
PET Dopamine Imaging in Hemi-Parkinson Monkey: Monitors gene for DA synthesis in right striatum

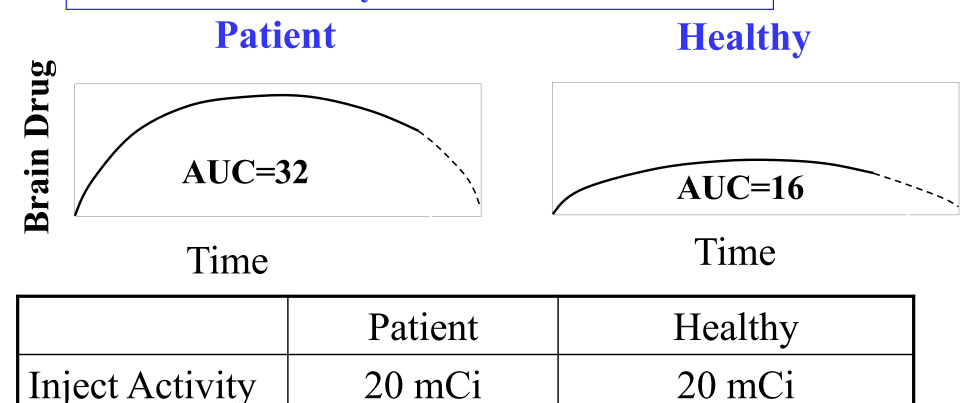


Outline of Talk

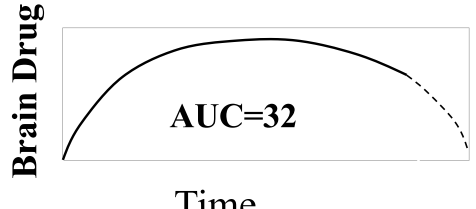
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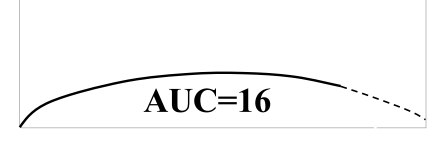






Patient Healthy



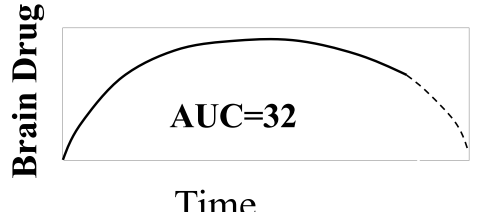


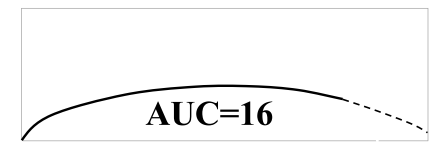
Time

Time

	Patient	Healthy
Inject Activity	20 mCi	20 mCi
Weight	50 kg	100 kg

Patient Healthy



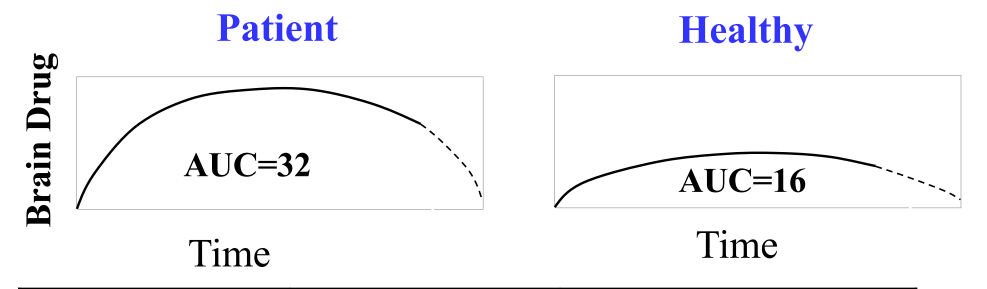


Time

Time

	Patient	Healthy
Inject Activity	20 mCi	20 mCi
Weight	100 kg	100 kg

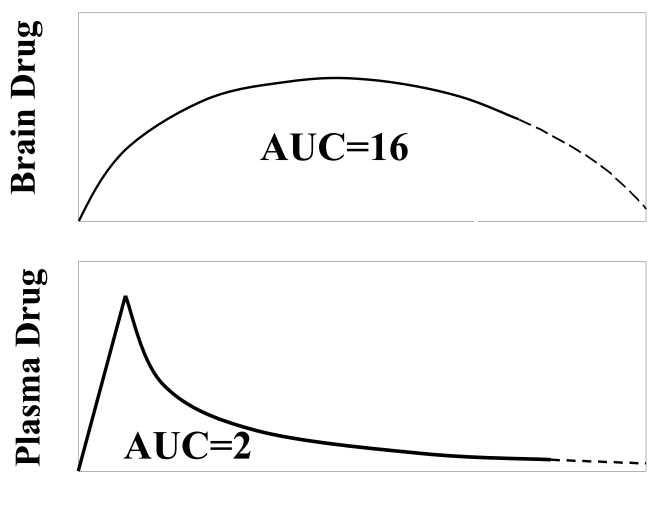
Brain Uptake of [18F]Fluoxetine: Measures Density of Serotonin Transporters



	Patient	Healthy
Inject Activity	20 mCi	20 mCi
Weight	100 kg	100 kg
Liver disease	Yes	No

Binding Potential (BP): Receptor Density * Affinity

BP equals uptake in brain relative to how much drug is delivered via arterial plasma.



Time

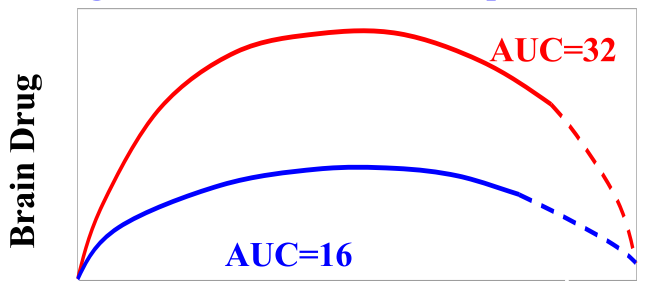
$$\mathbf{BP} = \frac{\mathbf{Area \ Brain \ Curve}}{\mathbf{Area \ Plasma \ Curve}}$$

$$BP = \frac{16}{2} = 8$$

Binding Potential: Independent of Injected Dose*

Double Plasma Input => Double Brain Response

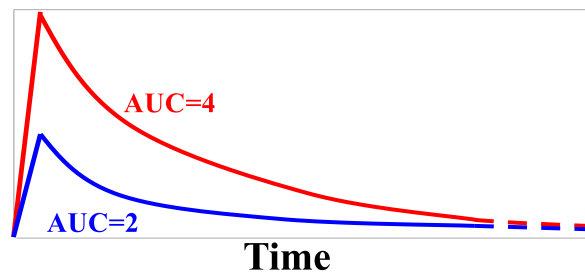
*If ligand does not saturate receptors - i.e., if tracer doses used



BP 1st Time =
$$\frac{16}{2}$$
 = 8

BP 2nd Time
$$=\frac{32}{4} = 8$$





BP can be calculated from the Area Under Curve (math integral) as well as rate constants (math differential).

From curves of plasma and brain radioactivity over time, estimate rate constants of entry and removal to/from tissue.

$$\begin{array}{c|c} K_1 \\ \hline \\ R_2 \end{array} \qquad \text{Brain}$$

$$BP = \frac{K_1}{k_2}$$

Tissue uptake is proportional to density of receptors and the affinity of the drug

Binding Potential

$$BP = \frac{B_{\text{max}}}{K_{\text{D}}} = B_{\text{max}} \times \frac{1}{K_{\text{D}}} = B_{\text{max}} \times \text{affinity}$$

 B_{max} = receptor density

 $K_{\rm D}$ = dissociation binding constant

$$\frac{1}{K_{\rm D}}$$
 = binding affinity drug

SUMMARY PET KINETICS

- Organ uptake is proportional to receptor density and affinity of drug
- Binding Potential (BP) = density X affinity
- "Drug Exposure" to tissue is AUC of: plasma concentration vs. time
- "Response" (uptake) of tissue is AUC of: tissue concentration vs. time

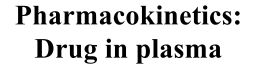
$$BP = \frac{\text{Response}}{\text{Exposure}} = \frac{AUC_{\text{tissue}}}{AUC_{\text{plasma}}}$$

 BP also equals ratio of rate constants of entry and removal to/from tissue

$$BP = \frac{K_1}{k_2}$$

Major Point of PET Pharmacokinetics (in words)

- Plasma pharmacokinetics provides a limited view of what's happening to drug in plasma.
- PET provides a limited view of what's happening to drug in tissue.
- Concurrent measurement of drug in plasma and of drug in tissue allows quantitation of the target of drug action
 - i.e., receptor.

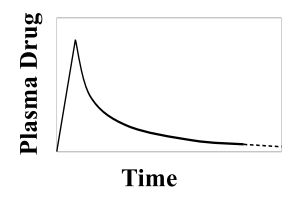


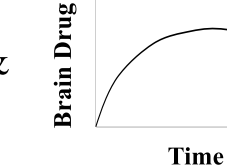
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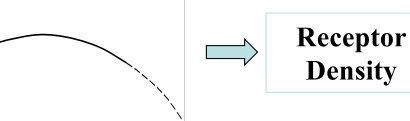
Pharmacodynamics: Drug acts at receptor



Receptor Density







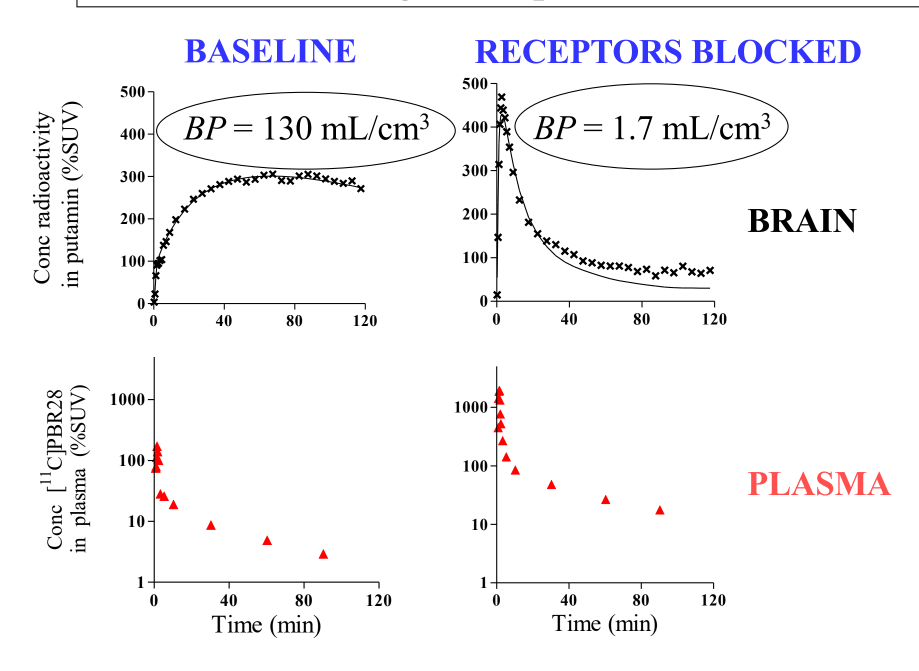
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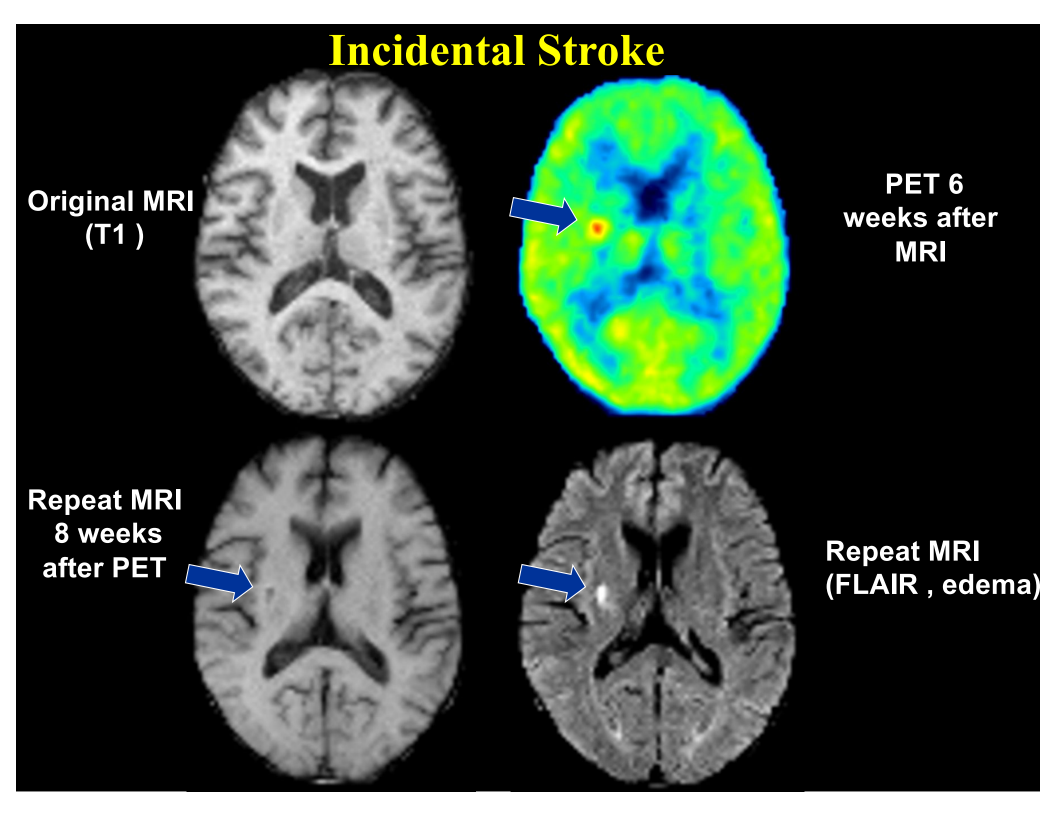
Translocator Protein (18 kDa) a.k.a. "peripheral benzodiazepine receptor"

- 1. Mitochondrial protein highly expressed in macrophages and activated microglia
- 2. Exists in periphery and brain
- 3. Multiple potential functions: steroid synthesis, nucleotide transport
- 4. Distinct from typical benzodiazepine GABA_A receptor in brain
- 5. Marker for cellular inflammation

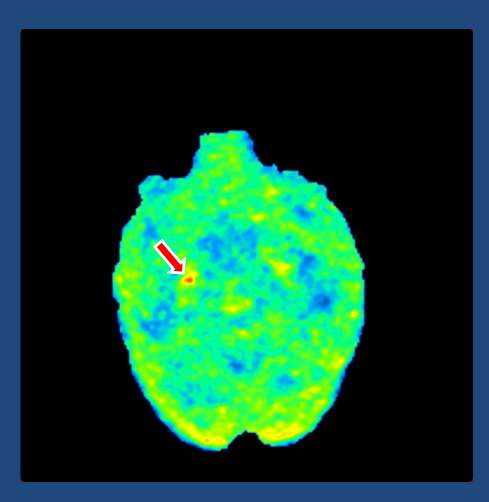
Receptor Blockade [11C]PBR28 in Monkey Brain: more radioligand in plasma and brain

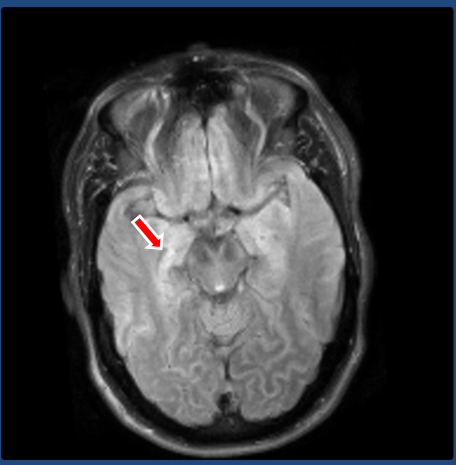


Receptor blockade displaces from lung & kidney. Drives more to brain but doesn't bind there.



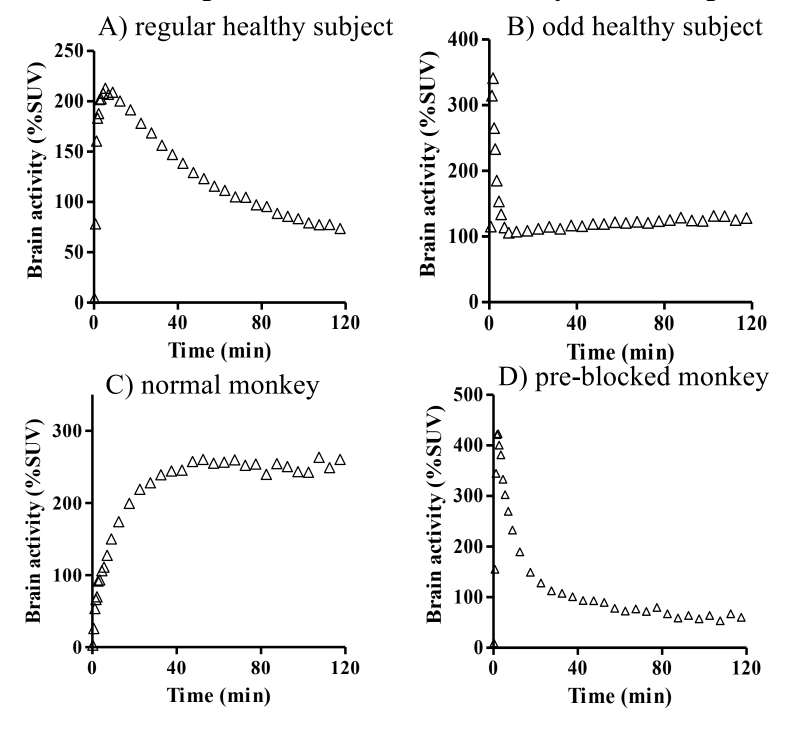
TSPO identifies epileptogenic focus in 15 of 16 patients.



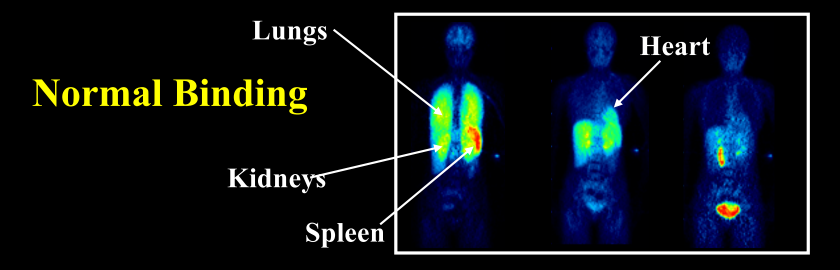


Hirvonen et al., JNM, 2012

Human with low uptake is similar to monkey with receptor blockade

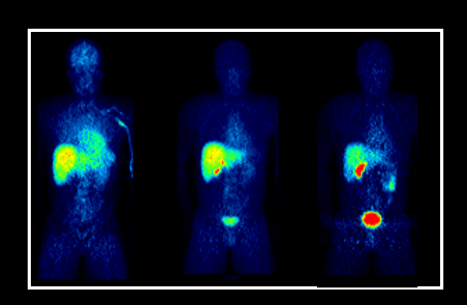


No Binding to [11C]PBR28 in Brain and Periphery



2 min 26 min 103 min

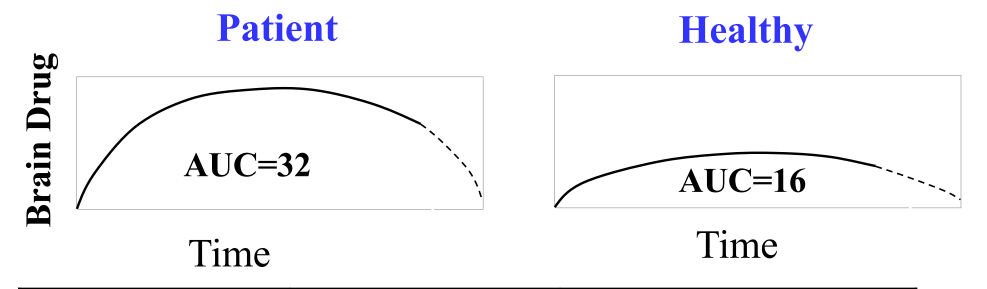
No Binding (~10% subjects)



TSPO rs6971 polymorphism causes differential affinity for PBR28

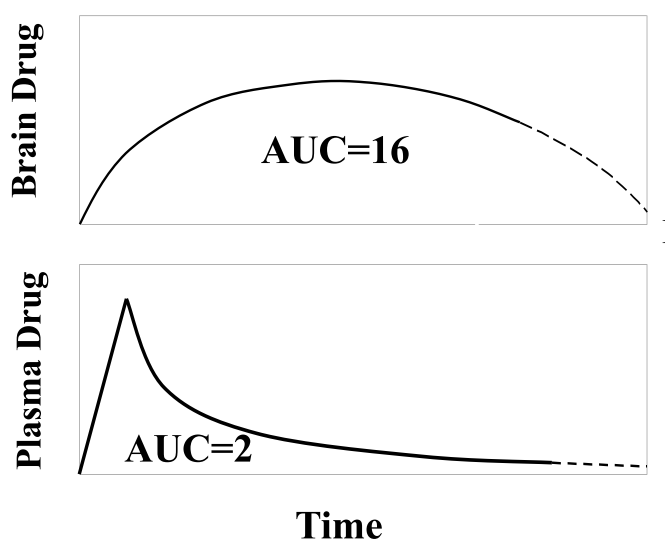
- Ala to Thr substitution
- Allelic frequency ~ 30%.
 - Prevalence of homozygotes ~ 9%
- Codominant expression
 - HAB high affinity binding
 - LAB low affinity binding
 - MAB reduced binding (mixed affinity states)

Brain Uptake of [18F]Fluoxetine: Measures Density of Serotonin Transporters



	Patient	Healthy
Inject Activity	20 mCi	20 mCi
Weight	100 kg	100 kg
Liver disease	Yes	No

Binding Potential (BP): Receptor Density * Affinity

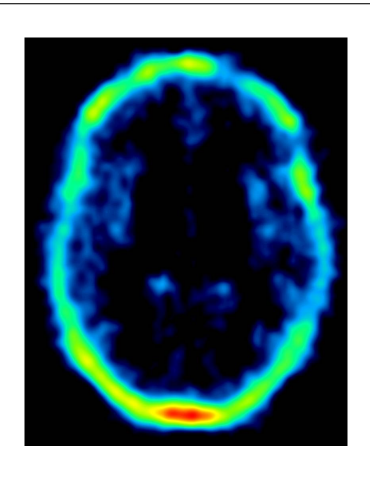


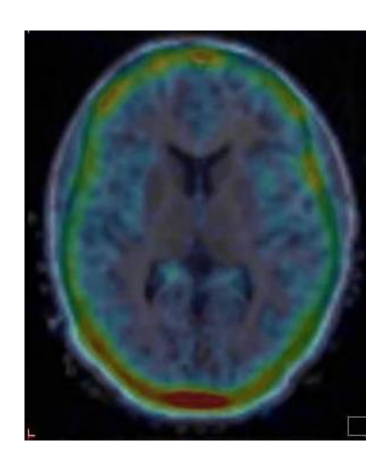
$$BP = \frac{16}{2} = 8$$

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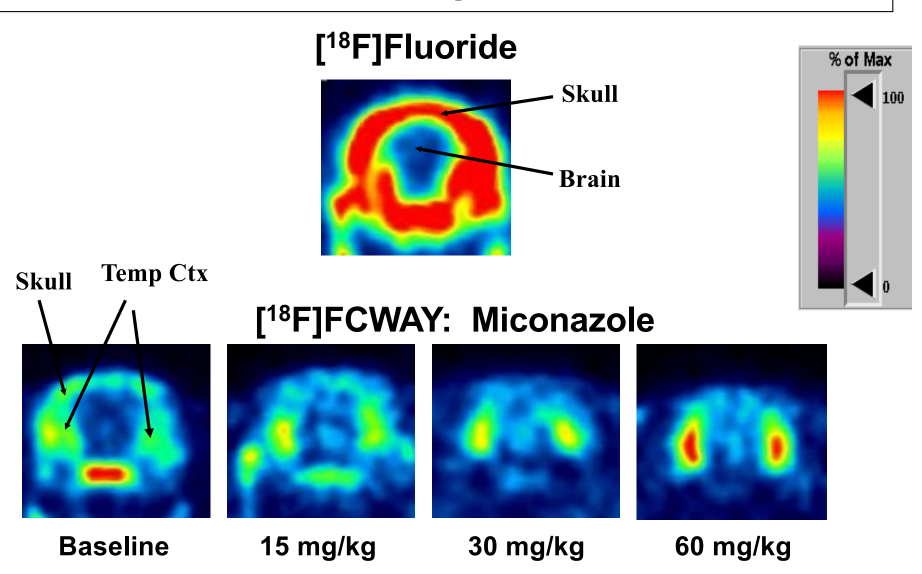
[18F]FCWAY: Defluorination Bone uptake: human skull at 2 h



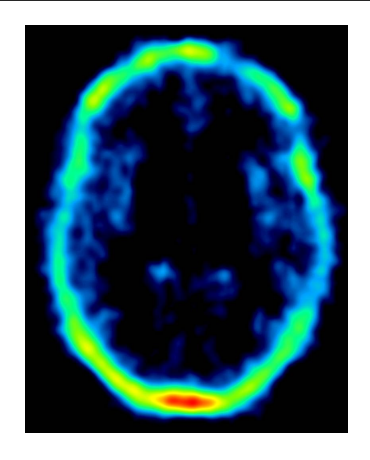


[¹⁸F]FCWAY: Defluorination ¹⁸F-fluoride ion accumulates in bone

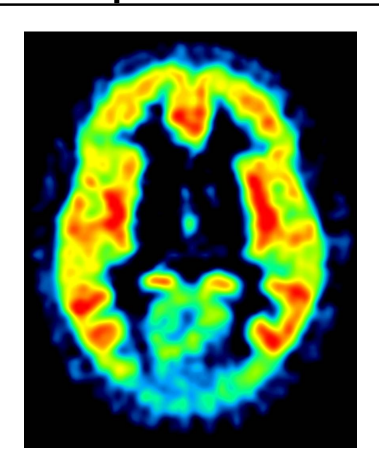
Miconazole Inhibits Defluorination & Bone Uptake



Disulfiram: Decreases Skull Activity & Increases Brain Uptake



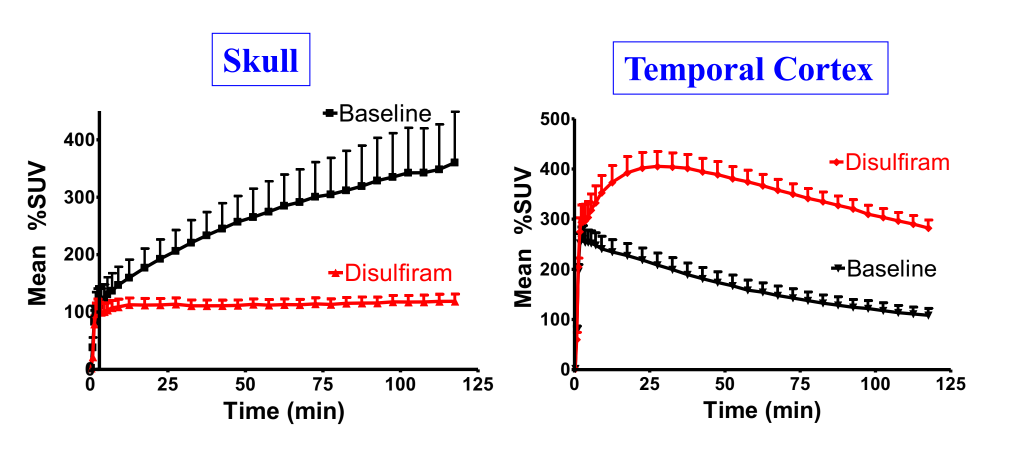
Baseline



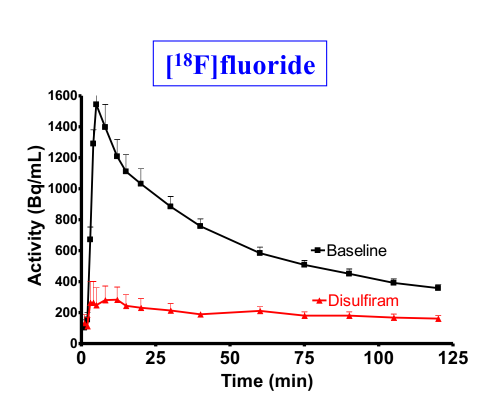
Disulfiram

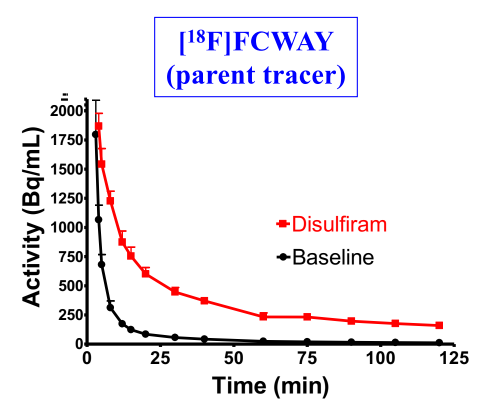
Images at 2 h in same subject. Disulfiram 500 mg PO prior night

Disulfiram: Decreases skull uptake of fluoride & Increases brain uptake of [18F]FCWAY



Disulfiram: Decreases plasma fluoride & Increases plasma radiotracer [18F]FCWAY





Summary

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- 4. Study drug distribution: block distribution to periphery and increase distribution to brain
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Self-Assessment Quiz: True or False?

- Imaging with positron emission tomography (PET) involves the injection of a radioactively labeled drug that emits a particle called a positron.
- PET shows the location of radioactivity in a cross section (or tomograph) of the body.
- PET can be used to quantify the density of specific proteins in the body.
- Compartmental modeling of PET data typically uses measurements over time of 1) PET images of the target tissue and 2) concentrations of unchanged parent radioligand in plasma.