# In Vivo Imaging of Neuromodulation Using Positron Emission Tomography: Optimal Ligand Characteristics and Task Length for Detection of Activation

Evan D. Morris, Ronald E. Fisher, Nathaniel M. Alpert, Scott L. Rauch, and Alan J. Fischman

Radiology (E.D.M., R.E.F., N.M.A., A.J.F.) and Psychiatry (S.L.R.) Departments, Massachusetts General Hospital and Harvard Medical School, Boston, Massachusetts

Abstract: Considerable evidence suggests that cognitive state affects local levels of neurotransmitter in the brain. We introduce a compartment model of neuroreceptor ligand kinetics to describe the effect of change in cognitive state on positron emission tomography (PET) signal dynamics. The model is used to establish optimal experimental conditions, timing of activation, and ligand characteristics, for detecting cognitive activation. The model, which follows free and bound endogenous neurotransmitter, describes the PET curve predicted for a single injection of radioligand in the presence or absence of activation. Activation was conceptualized as the performance of a task that raises the level of neurotransmitter that competes for receptor sites with the radioligand. Simulating the dopamine system, for example, required making assumptions regarding the kinetic rate constants for binding/dissociation of endogenous dopamine to/from the receptor and dopamine concentrations in the synapse. Simulations suggest that activation of dopamine should be detectable with PET and the D2 receptor ligand [11C]raclopride, although this ligand might not be optimal. Aspects of experimental design can be modified to optimize the likelihood of detecting neurotransmitter changes. The ideal radioligand for these studies should bind irreversibly to its receptor. Furthermore, the task should commence at injection time and last for at least 7 minutes. Optimal task duration depends on the dynamics of free radioligand in the tissue and can be determined via model simulations for any well-characterized receptor ligand. Flow effects were shown to be distinguishable from those of neurotransmitter activation. General principles regarding desirable ligand characteristics and activation timing held for both the D2 receptor and the dopamine transporter site. 0 1995 Wiley-Liss, Inc.

Key words: endogenous, dopamine, receptor, activation, mathematical model, optimization

#### INTRODUCTION

The basic idea of this work is that it may be possible to have subjects perform a cognitive, sensory, or motor task that would increase the level of a neurotransmitter in the synaptic cleft, which would, in turn, chang the kinetics of a corresponding receptor ligand. I assess the likelihood of detecting the activation of particular neurotransmitter system with positron emi sion tomography (PET), we performed simulations a compartmental model of receptor binding that we modified to account for the competition betwee endogenous transmitter and radioactive ligand for the same class of (receptor) binding sites.

The evolution of receptor-binding models in PE began with the Mintun formulation, which propose

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Address reprint requests to Dr. Alan Fischman, Division of Nuclear Medicine, MZ-TIL-201, Massachusetts General Hospital, Fruit Street, Boston, MA 02114.

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compartments for plasma, free, and bound species while assuming that nonspecific binding was instantaneous [Mintun et al., 1984]. The standard model was subsequently modified [Bahn et al., 1989; Huang et al., 1989] by the introduction of a bound-to-specific activity ratio as a means of accounting for the effect of unlabeled ligand as a competitor for radioligand binding sites. Delforge et al. [1990] proposed parallel models for following both the "hot" and "cold" exogenous ligands separately (i.e., with separate compartments). We chose to begin with the formulation of Delforge et al. [1990] because it is the most comprehensive approach [Morris et al., submitted]. In essence, what needed to be added to the standard model were the equations that describe the effect of yet another competitor-the endogenous neurotransmitter-for the binding sites of interest. Logan et al. [1991] have attempted a similar analysis using a different formulation than the one proposed herein, adding compartments to the conventional model and specifying a loss rate for endogenous material from the free compart-

We present a series of simulations of a deterministic compartmental system subjected to transient step changes in endogenous competitor concentration to model the effect of cognitive activation. Neurochemical transmission involves intermittent synaptic release of transmitter on the millisecond and micron scale. Before we could predict the effect of synaptic activation on a PET signal, these microphenomena had to be related to macroscopic ones on the time and spatial scales of phenomena that are measured with PET. That process of translating published data from in vivo and in vitro results into receptor modeling assumptions is discussed in detail in the accompanying paper [Fisher et al., this issue]. The simulations presented herein were based on data for existing ligands (raclopride, CFT) (WIN 35, 428) and extended to hypothetical ligands whose kinetic properties were varied. In the course of our theoretical investigations, we discovered that certain characteristics of timing and kinetics could amplify the predicted effect of activation, leading to greater detectability with PET. We attempt to formalize the optimization of future PET studies geared to identifying task-driven activation of specific neurochemical systems using the dopamine system as an example.

#### **METHODS**

#### Theory

To the Delforge model (Equations 1-6) we added a differential equation describing the receptor binding

(and dissociation) of free endogenous neurotransmitter. The free endogenous chemical concentration presented to the receptor (i.e., in the synaptic cleft) was modeled simply as a step function; step up for activation of the neurotransmitter and step down for return of the endogenous level to the resting state. As suggested by Logan et al. [1991], the actual fluctuation of endogenous neurotransmitter in the synaptic cleft is far more complicated than a step function and certainly varies on the millisecond time scale; however, the low temporal resolution of PET (seconds to minutes) requires that such changes be detected as shifts in average level. Logan et al. [1991] made model modifications similar to ours to study the effect of transient changes in endogenous chemical on the measured density of dopamine (D2) receptors. Their concern was that unintended transients in endogenous dopamine concentration might alter the estimated value of available D2 receptors, B' max, as measured with [18F]N-methylspiroperidol ([18F]NMS). Consequently, they modeled the possible rapid change in dopamine concentration in the free space via an additional differential equation with a first order term. Since we are concerned with intentional and sustained alterations in endogenous transmitter level via a motor or cognitive task, a step function was chosen to describe these alterations.

The model used for simulations is described schematically in Figure 1. Both labeled ("hot") and unlabeled ("cold") exogenous ligand are introduced into the system via the plasma. Together they enter and exit the brain according to the transport parameters,  $K_1$  and  $k_2$ . These are the only model parameters that are influenced by blood flow. Once in the brain the ligand experiences three possible fates: diffusion back to the plasma compartment, binding nonspecifically as indicated by the NS and NSc compartments, or binding at the saturable receptor site B or Bc. Note that the superscript c means cold and superscript en means endogenous. The dotted box drawn around the binding compartments indicates that the cold and hot ligand compete for a limited number of common sites, B + B<sup>c</sup>. Of course, only the hot ligand is detected by PET, as indicated by the solid box labeled "PET pixel." The inclusion of the endogenous chemical in the model is diagrammed by the two compartments below the dotted line. For the purposes of our model, the endogenous chemical is introduced into the system at the free compartment, Fen, whose value is specified at any time (by equation 8). The endogenous chemical can bind to the receptor sites, Ben. As indicated by its inclusion in the dotted box, the pool of possible binding sites for endogenous material are the same pool as those for exogenous ligand, [B<sub>max</sub> – B + B<sup>et</sup> + B<sup>ett</sup>]. Mathematically, this means that the differential equations describing accumulation of the exogenous species (labeled and unlabeled) are coupled to the equations for the endogenous species via the three binding compartment variables (see equations 1, 2, 4, 5, and 7). It is important to note that the total number of receptor sites B<sub>max</sub> specified by this enhanced PET model is not the same as the receptor density term measured in studies that assume the endogenous transmitter to be in steady state. That term, which refers only to the number of sites not bound on average to endogenous material, is appropriately designated B'<sub>max</sub> to distinguish it from the theoretical total number of receptor sites.

# Model equations

Compartments of labeled ligand:

$$\frac{dF}{dt} = K_1 C_p(t) - k_2 F - k_{on} F [B_{max} - B - B^c - B^{en}]$$

$$+ k_{off}B - k_5F + k_6NS - \lambda F$$
 (1)

$$\frac{dB}{dt} = k_{on}F[B_{max} - B - B^{c} - B^{en}] - k_{off}B - \lambda B \quad (2)$$

$$\frac{dNS}{dt} = k_5F - k_6NS - \lambda NS. \quad (3)$$

Compartments of unlabeled ligand:

$$\frac{dF^{c}}{dt} = K_{1}C_{p}^{c}(t) - k_{2}F^{c} - k_{on}F^{c}[B_{max} - B - B^{c} - B^{on}]$$

$$+ k_{off}B^c - k_5F^c + k_6NS^c$$
 (4)

$$\frac{dB^{c}}{dt} = k_{on}F^{c}[B_{max} - B - B^{c} - B^{en}] - k_{off}B^{c} \qquad (5)$$

$$\frac{dNS^c}{dt} = k_5 F^c - k_6 NS^c. \quad (6)$$

Compartments of endogenous transmitter:

$$\frac{dB^{en}}{dt} = k_{on}^{en} F^{en} [B_{max} - B - B^{c} - B^{en}] - k_{od}^{en} B^{en} \qquad (7)$$

$$F^{en} = F^{en}(0) + A \cdot U(t - t_{up}) - A \cdot U(t - t_{down}).$$
 (8)

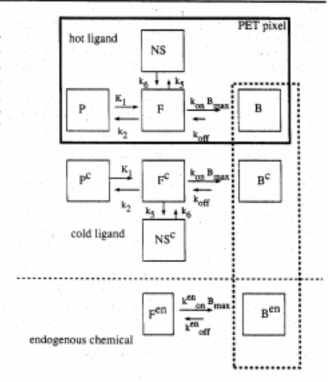


Figure I.

Schematic corresponding to model described in Equations 1-8. The diagram is composed of three sections corresponding to the three chemical species of interest: radiolabeled exogenous ligand (hot), unlabeled exogenous ligand (cold) and endogenous transmitter. Each section contains boxes (i.e., compartments) representing the possible states of the particular species. Hot ligand can exist in the plasma state (P), free state (F), receptor-bound state (B), or nonspecifically bound state (NS). Cold ligand can exist in the plasma state (Pc), free state (Fc), receptor-bound state (Bc), or nonspecifically bound state (NSc). Endogenous transmitter is modeled as existing either in the free state (Fer) or the bound state (Ber). The heavy dashed box around the three bound compartments indicates that all three chemical species are in competition for the same binding sites, which are limited in number. Appropriate first order rate constants (labeling each arrow) govern the transition of ligand from one state to another. Free ligand is transformed to bound ligand via its bimolecular association with available receptor sites, hence the shorthand label keeBress for the corresponding arrow. The receptor modeling approach of Delforge et al. [1990], diagrammed above the dashed line, has been appended by the inclusion of compartments for endogenous transmitter, below the line.

Rate constants governing transfer of ligand between compartments  $(k_2, k_{off}, k_5, k_6)$  are expressed in  $(min^{-1})$ ,  $K_1$  is measured in  $(ml/min)(g^{-1})$ , and  $k_{on}$  is in  $(pmol/ml)^{-1}(min^{-1})$ ;  $\lambda$   $(min^{-1})$  is the radioactive decay constant.  $B_{max}$  is measured in concentration units (pmol/ml) as are all the state variables in the above equations  $(F, F^c, F^{en}, B, B^c, B^{en}, NS, NS^c)$ .  $F^{en}(0)$  is the

basal level of endogenous transmitter in the free compartment.  $U(t-t_{up})$  and  $U(t-t_{down})$  are the unit step functions at times  $t_{up}$  and  $t_{down}$ , respectively. The magnitude of the step change is A (pmol/ml). Cp and  $Cp^c$  are the metabolite-corrected plasma concentrations for the labeled and unlabeled ligand, respectively. For a single injection, the plasma functions are related by the specific activity of the injected material.

# Protocol for simulation experiments

All of the computer simulations presented below follow a uniform protocol. Each simulation is based on a single injection of high specific activity radiolabeled receptor ligand. The specific activity and activity of the injectate are known at the time of injection. The dynamic PET activity is predicted for 60 minutes following injection in 1 minute time intervals. The metabolite-corrected plasma activity during the duration of the scan is assumed to be known. The activation task, which is modeled as having an immediate effect on the free endogenous neurotransmitter level, starts at the time of injection unless otherwise specified.

#### Parameter values used for simulations

For detecting activation at the post-synaptic D2 receptor, the model time-activity curves were generated using parameter values reported by Farde et al. [1989] for [11C]raclopride binding in the putamen of normal subjects. These values are based on a threecompartment model that ignores nonspecific binding. Thus, in our simulations of raclopride and other D2 receptor ligands, the nonspecific binding rate constants,  $k_5$  and  $k_6$ , were set to 0. The other raclopride parameter values are given in the last footnote of Table I; the specific activity of the injection was set at 900 mCi/µmol. Simulations of studies to detect activation at the dopamine transporter site were generated using parameters determined for [11C]CFT in monkey studies in our lab (unpublished data). The parameter values are given in the last footnote of Table II; the specific activity of the injection was set at 1,000 mCi/μmol.

As we discussed at length elsewhere [Fisher et al., 1995], we had to estimate a steady state concentration of endogenous dopamine and its K<sub>D</sub> from the literature [see Table I in Fisher et al., 1995, for a summary of sources]. When modeling radioligand behavior at the D2 receptor site, the free compartment in our model corresponds to the micro-environment surrounding the post-synaptic receptor. Our interpretation of published in vivo studies is that the average concentration

of dopamine in that region is in the 100 nM range at rest and 200 nM during activation [Fisher et al., 1995]. By varying the value of the dopamine concentration, we checked the sensitivity of our results to this interpretation. The KD for endogenous neurotransmitter is taken to be 100 nM [Gingrich and Caron, 1993]. Because we assume that binding and release of ligand at the D2 site must be fast, we chose ken and ken values of 0.25 ml/(pmol·min) and 25.0 min-1, respectively. When modeling binding at the dopamine transporter site, which is likely to be both intra- and extra-synaptic [Iversen and Kelly, 1975; Cerruti et al., 1991], the free endogenous concentration is estimated to be 30 nM at rest and 90 nM during activation [Fisher et al., 1995]. For transporter site simulations we chose values of 0.75 ml/(pmol·min) and 25.0 min-1, respectively, for ken and ken so that action of the endogenous transmitter would still be fast relative to the exogenous ligand and the transporter operates near its KD when the organism is at rest ( $K_D = k_{off}^{en}/k_{on}^{en} = 33.3 \text{ nM}$ ).

#### Numerical, statistical methods

The model depicted in equations 1-8 was solved numerically using an ordinary differential equation solver employing the Adams-Gear method as provided in the IMSL/IDL programming language (IMSL Inc., Sugar Land, TX). The plasma input function, Cp(t), used for the simulations was an analytical bi-exponential decay function based on prior measurements. The chi-square statistic  $(\chi^2)$ , was used to rank activated PET curves by how different they were from their corresponding resting PET curve. The parameter was calculated as the weighted sum of the squares of the distances between the activated (a) and control (c) curves:  $\chi^2 = \sum [(a - c)_i/\sigma_i]^2$  [Press et al., 1992]. The variance of each simulated data point, σ<sub>i</sub>, was modeled as  $\sigma_i = f \cdot c_i / \Delta t_i$ , where f is a proportionality factor that was kept constant for a set of simulations to be compared with each other, c is the value of the control curve at the ith time point, and  $\Delta t_i$  is the length of the PET scan for the ith time point.

# Ligand nomenclature

Each simulated dynamic PET activity curve reported below requires a set of kinetic parameters for a given ligand. Some of the ligands discussed, such as [11C]raclopride, or [11C]CFT, are currently in use and therefore the necessary parameter estimates are known. In order to establish optimal kinetic characteristics, we also present model-based predictions for hypothetical ligands that do not currently exist. Thus,

TABLE I. Optimization of ligand characteristics for D2 receptor site

Simulation description	k <sub>on</sub> (raclo.)	k <sub>off</sub> (raclo.)	K <sub>D</sub> (raclo.)	B <sub>max</sub> a	% Occupation at rest <sup>b</sup>	[Resting endogenous dopamine] (nM)	[Activated endogenous dopamine] (nM)	Time of activation (min)	x <sup>2</sup>
A. Raclopride			7.7						
<ol> <li>Raclopride with activation</li> </ol>	0.01	0.099	9.9	56	50	100	200	0.7	
B. Vary ligand binding	0.01	0.055	7.7	30	50	100	200	0-7	5,249
1. 2× larger k <sub>an</sub>	0.02	0.099	4.9	56	50	100	200	0-7	6,136
2. 4× larger kon	0.04	0.099	2.5	56	50	100	200	0-7	4,623
3. High kon	0.1	0.099	0.99	. 56	50	100	200	0-7	1,579
4. Low kon	0.001	0.099	99.0	56	50	100	200	0-7	432
<ol> <li>High k<sub>off</sub></li> </ol>	0.01	0.99	99.0	56	50	100	200	0-7	212
6. Low koff	0.01	0.0099	0.99	56	50	100	200	0-7	15,470
<ol><li>Irreversible</li></ol>	0.01	0.0	0.0	56	50	100	200	0-7	18,210
C. Keep K <sub>D</sub> constant									20,220
<ol> <li>High k<sub>on</sub>, k<sub>off</sub></li> </ol>	0.1	0.99	9.9	56	50	100	200	0-7	2,122
2. Low kon, koff	0.001	0.0099	9.9	56	50	100	200	0-7	3,315
D. Effect of endog- enous [dopamine] 1. 20% of chosen									
[dopamine] 2. 40% of chosen	0.01	0.099	9.9	32.67	~17	20	40	0-7	1,591
[dopamine] 3. 5-fold higher	0.01	0.099	9.9	39.2	~ 29	40	80	0-7	2,151
[dopamine] 4. 10-fold higher	0.01	0.099	9.9	160	~83	500	1000	0-7	14,238
[dopamine]	0.01	0.001	9.9	295	~91	1000	2000	0-7	15,163

As noted in text, B<sub>max</sub> when modeling the effect of endogenous neurotransmitter is a superset of the available number of receptors, B<sub>max</sub> measured in conventional receptor imaging with raciopride where the endogenous chemical level is ignored.

the reader is advised that a "lower kon version of raclopride" (i.e., a ligand whose kinetic parameters are the same as the published raclopride numbers except that the association rate constant is one-tenth the published value) does not exist but is a theoretical construct for the purposes of experimental design. The particular variations in kinetic characteristics implied by "high kon raclopride" or "irreversible CFT" are catalogued in Tables I and II. As a point of interest, we note that CFT is one of many fluorophenyltropane analogs of cocaine referred to as WIN 35,XXX that have been evaluated previously in vitro [Madras et al., 1990] for their specificity to the dopamine transporter site. It is possible that one or more of these compounds may resemble the hypothetical ligands that we evaluate below.

#### RESULTS

# Model simulation for binding of ["C]raclopride at the D2 receptor with activation of endogenous dopamine

Our primary interest was in determining whether activation of a neurotransmitter system could be detected with PET and a currently available radioligand. Figure 2 shows model predictions related to a dynamic PET study with [11C]raclopride that includes activation of the dopamine system for the first 7 minutes of the study (via a continuous cognitive or movement task). The predictions in Figure 2 reflect our assumptions, discussed above, that the average endogenous dopamine concentration presented to

<sup>&</sup>lt;sup>b</sup> The percentage occupation of the receptor sites is determined by knowing K<sub>D</sub><sup>so</sup> for dopamine, which has been set at 100 nM, and the resting endogenous dopamine level, which is given in column 7. Our B<sub>max</sub>, which is not the value measured by Farde et al. [1989], is set by augmenting B'<sub>max</sub> of footnote c, so the percentage of B<sub>max</sub> that is unoccupied equals the previously determined parameter B'<sub>max</sub>.

 $<sup>^</sup>c$  [ $^{11}$ C]raclopride kinetic parameters taken from subject C of Farde et al. [1989]:  $K_1 = 0.199 \, \text{min}^{-1}$ ,  $k_2 = 0.36 \, \text{min}^{-1}$ ,  $k_{an} = 0.01 \, \text{ml/(pmol · min)}$ ,  $k_{off} = 0.099 \, \text{min}^{-1}$ ,  $B'_{max} = 28 \, \text{pmol/ml}$ . The endogenous transmitter parameters were set at  $k_{cn}^{an} = 0.25 \, \text{ml/(pmol · min)}$  and  $k_{cff}^{an} = 25.0 \, \text{min}^{-1}$ .

TABLE II. Optimization of ligand characteristics for dopamine transporter site

Simulation description	k <sub>on</sub> (CFT)	k <sub>off</sub> (CFT)	Ko	B <sub>max</sub> s−c	% Occupation at rest	[Resting endogenous dopamine] (nM)	[Activated endogenous dopamine] (nM)	Time of activation (min)	x <sup>2</sup>
A. CFT <sup>d</sup>									
<ol> <li>CFT with activation</li> </ol>	0.00153	0.062	40.5	1,358	~50	30	90	0-7	1,310
B. Vary ligand binding							P100 1 2 2		
1. High kon	0.0153	0.062	4.05	1,358	~50	30	90	0-7	46
2. Low ken	0.000153	0.062	405	1,358	~50	30	90	0-7	763
3. High k <sub>off</sub>	0.00153	0.62	405	1,358	~50	30	90	0-7	. 400
4. Low kor	0.00153	0.0062	4.05	1,358	~ 50	30	90	0-7	1,592
5. Irreversible	0.00153	0.0	0.0	1,358	~ 50	30	90	0-7	5,433

As noted in text, B<sub>max</sub> when modeling the effect of endogenous neurotransmitter is a superset of the number of receptors, B'<sub>max</sub>, measured with conventional receptor imaging.

the post-synaptic receptors is 100 nM at rest and can be doubled by the increased frequency of firing of dopaminergic neurons. The dynamic concentration curve for each model compartment (p, plasma; f, free; b, bound) multiplied by its respective volume fraction is shown in Figure 2a. The total curve (t), which is proportional to the predicted PET activity curve, is the sum of these curves. This figure is corrected for radioactive decay. The raclopride parameters for this simulation were taken from Farde et al. [1989; see footnote to Table I] who were able to model PET activity without a nonspecific binding compartment, hence the absence of a nonspecific curve in Figure 2b. It is important to note that the concentration of labeled ligand in the free compartment (f) rises sharply after injection and peaks by 5 minutes. This amount of exogenous radioligand in the free compartment is the sole pool of radioactive signal that can respond to the abrupt change in concentration of endogenous competitor that we have hypothesized.

All the rate constants determined by Farde et al. [1989] should apply to the model presented above with the exception of the receptor density. As mentioned previously, the term B<sub>max</sub> in our model represents the total possible binding sites for any of three species, labeled and unlabeled raclopride as well as endogenous dopamine in the synapse. Thus, the

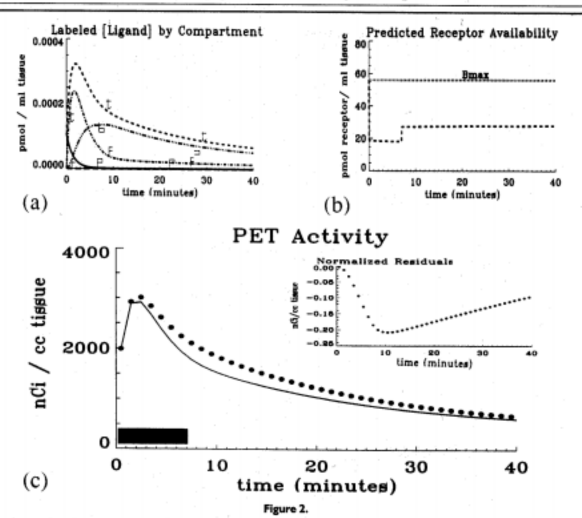
model must be constrained so that the difference between Bmsx and the number of receptors occupied by endogenous dopamine at steady state equals the B'mex value determined by other groups. This constraint on B<sub>max</sub> is displayed graphically in Figure 2b. The total number of receptor sites that potentially can be bound by any of the three species is designated by the constant curve at 56 pmol/ml. The rise in number of available receptors (dashed curve) reflects the abrupt drop in endogenous dopamine that we expect with termination of the cognitive activation task at 7 minutes. Activation tasks of other durations are analyzed in a section below. The model indicates that once the activation has ceased (i.e., subject is in the resting state), the number of available receptors is 28 pmol/ml [Farde et al., 1989]. The fraction of occupied receptor sites at rest (50%) is consistent with our assumption that the endogenous ligand is operating at its KD (average post-synaptic dopamine concentration = 100  $nM = K_D$ ).

The effect of activation on the dynamic PET curve is illustrated in Figure 2c; the period of activation is indicated by the horizontal bar on the time-axis. The solid curve is the activity corresponding to the total concentration curve in Figure 2b. The dotted curve is the control curve, that is, the PET signal predicted by either the conventional receptor model (Equations

The percentage occupation of the receptor sites is determined from K<sub>0</sub><sup>m</sup> for dopamine transporter, which has been estimated at 30 nM, and the resting endogenous dopamine level at the (extra-synaptic) site, given in column 7. B<sub>max</sub> is set by augmenting B'<sub>max</sub> of footnote d, so the percentage of B<sub>max</sub> that is unoccupied equals the previously determined parameter B'<sub>max</sub>.

c B<sub>max</sub> value of 1,358 pmol/ml may seem quite high but it should be noted that the corresponding B'<sub>max</sub> value of 679 that we measured in beboons via PET is not too far from the in vitro estimate of ~350 pmol/g published by Madras et al. [1989].

<sup>&</sup>lt;sup>d</sup> [<sup>11</sup>C]CFT kinetic parameters taken from model fits to preliminary monkey data obtained in our lab:  $K_1 = 0.3 \text{ min}^{-1}$ ,  $k_2 = 0.118 \text{ min}^{-1}$ ,  $k_{en} = 0.00153 \text{ ml/(pmol·min)}$ ,  $k_{eff} = 0.062 \text{ min}^{-1}$ ,  $k_{en} = 679 \text{ pmol/ml}$ ,  $k_5 = 0.00217$ ,  $k_6 = 0.000175$ . The endogenous transmitter parameters were set at  $k_{en}^{en} = 0.75 \text{ ml/(pmol·min)}$  and  $k_{eff}^{en} = 25.0 \text{ min}^{-1}$ .

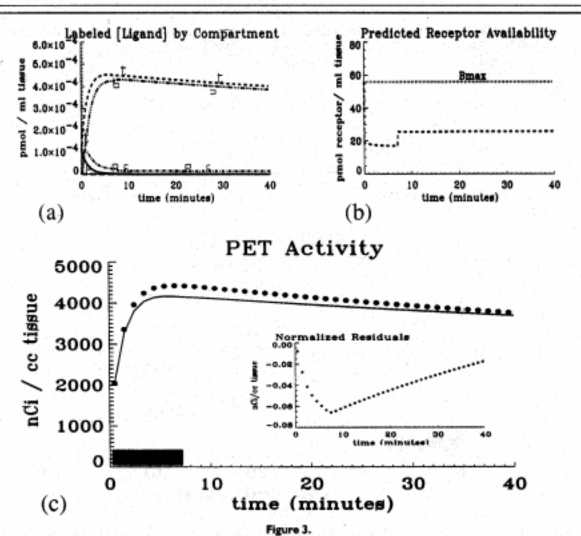


(a) Simulation of total concentration (pmol/ml) of labeled ligand, [¹¹C]raclopride, in brain tissue (t), which is the sum of the ligand in each individual compartment (p, plasma; f, free; b, bound), during the first 40 minutes of a cognitive activation study. (b) Availability of dopamine (D2) receptor sites (pmol/ml) remaining during activation of endogenous dopamine. After initial drop in available sites due to 7 minutes of activation, receptor availability returns to 50% of total receptors. Horizontal line at B<sub>max</sub> represents the theoretical maximum number of available receptor sites, which is specified.

here as 56 pmol/ml. (c) Predicted PET activity curves (nCI/cc) for study (solid curve), which include cognitive activation for first 7 minutes and for resting study with no activation (filled circles). Activation refers to increased endogenous dopamine level from 100 to 200 nM and is indicated by solid bar on x-axis. Inset: Normalized residuals ([activated-rest]/activated) corresponding to the curves in c. All figures except for the normalized residual plot are corrected for radioactive decay.

1-6) with the Farde parameters (i.e.,  $B'_{max} = 28 \text{ pmol/ml}$ ) or our augmented model (Equations 1-8) where the free endogenous concentration is fixed at  $F^{en}(0) = 100 \text{ nM}$  (and  $B_{max} = 56 \text{ pmol/ml}$ ). The normalized residuals for Figure 2 (inset), show that, in relative terms, the difference between the two PET curves is maximal at 21% of the activated curve at 10 minutes. The residual plot (which consists of all negative values) also indicates that the PET signal for the

activated case is always lower than the corresponding resting PET curve. The residual plot is not corrected for decay. The difference between these curves can be quantified by the  $\chi^2$  parameter, given for this simulation in section A of Table I. The control curve and the activated curve are nearly coincident by 40 minutes (i.e., 33 minutes after the end of activation), which suggests that any scanning after 40 minutes may be superfluous.

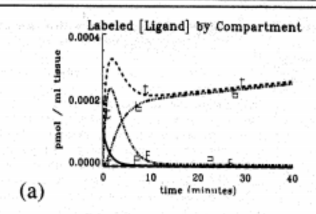


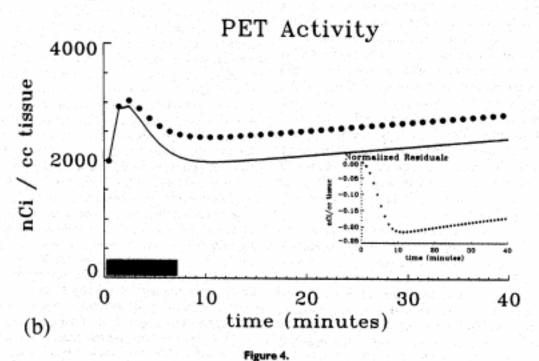
(a) Simulation of total concentration (pmol/ml) of high k<sub>ce</sub> version of labeled ligand, [<sup>11</sup>C]raclopride, in brain tissue (t), which is the sum of the ligand in each individual compartment (p, plasma; f, free; b, bound), during the first 40 minutes of a cognitive activation study. (b) Availability of dopamine (D2) receptor sites (pmol/ml) remaining during activation of the endogenous dopamine. After initial drop in available sites due to 7 minutes of activation, receptor availability returns to 50% of total receptors as in Figure 2. Horizontal line at B<sub>max</sub> represents the theoretical maximum number of available.

receptor sites, which is specified here as 56 pmol/ml. (c) Predicted PET activity curves (nCi/cc) for study (solid curve), which include cognitive activation for first 7 minutes and for resting study with no activation (filled circles). Activation refers to increased endogenous dopamine level from 100 to 200 nM and is indicated by solid bar on x-axis. Inset: Normalized residuals ([activated-rest]/activated) corresponding to the curves in c. All figures except for the normalized residual plot are corrected for radioactive decay.

# Optimal characteristics for raclopride-like ligand

Further simulations were performed with the augmented receptor model to investigate additional ways of optimizing the difference between control and activated curves and thus to maximize the chance of detecting neurotransmitter activation at the postsynaptic D2 receptor. These simulations are summarized in sections B and C of Table I. These two sections of Table I catalogue the  $\chi^2$  value for simulations where the  $k_{on}$  and/or  $k_{off}$  of a theoretical raclopride-like ligand have been changed. Counter to our intuition, the most important characteristic of an optimal ligand for these receptor activation studies appears to be its irreversibility, that is, a small dissociation rate constant,  $k_{off}$ . Comparing the entries in section B of Table I with section A shows that  $\chi^2$  is no better (i.e., equal or smaller) than native raclopride for any modulation of





(a) Simulation of total concentration (pmol/ml) of irreversible (k<sub>eff</sub> = 0) version of labeled ligand, [11C]raclopride, in brain tissue (t), which is the sum of the ligand in each individual compartment (p. plasma; f, free; b, bound), during the first 40 minutes of a cognitive activation study. Availability of dopamine (D2) receptor sites (pmol/ml) remaining during activation of the endogenous dopamine (not shown) is comparable to the plots in Figures 2b and

3b. (b ) Predicted PET activity curves (nCl/cc) for study (solid

curve), which include cognitive activation for the first 7 minutes and for resting study with no activation (filled circles). Activation refers to increased endogenous dopamine level from 100 to 200 nM and is indicated by solid bar on x-axis. Inset: Normalized residuals ([activated-rest]/activated) corresponding to the curves in b. All figures except for the normalized residual plot are corrected for radioactive decay.

 $k_{on}$  or  $k_{off}$  except for a lower  $k_{off}$  version of raclopride. The ideal case of a ligand that is identical to raclopride except for complete irreversibility ( $k_{off} = 0$ ) maximizes the difference between control and activated curves,  $\chi^2 = 18,210$ . It is interesting to note that the  $K_D$  of the ligand for the "High  $k_{on}$ " and "Low  $k_{off}$ " cases are the same but the latter yields a much larger  $\chi^2$ . In other

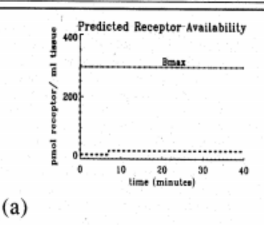
words, K<sub>D</sub> appears not to be the critical ligand characteristic for predicting the separation of the PET curves. Section C of Table I confirms that K<sub>D</sub> of the radioligand is not the determinant of potential success of the proposed experiments. The two simulations summarized here were run with the K<sub>D</sub> of native raclopride but the individual rate constants were either increased or decreased simultaneously. Neither change gave  $\chi^2$  values as large as the original raclopride simulation.

Figure 3 is analogous to Figure 2 for a ligand identical to [11C]raclopride in all respects except that its kon is 10-fold higher (see entry B-3 in Table I). The resting and activated levels of dopamine are identical to those for Figure 2. Figure 3a shows the predicted radioligand concentrations by compartment. As would be expected with a very fast binding ligand, the total signal is primarily made up of bound ligand. The reserve of free radioligand in the crucial first 10 minutes of the study is much less than in Figure 2b. The appearance of the PET curves in Figure 3c confirms the  $\chi^2$  entry in Table I, namely, the activated (solid curve) and control curve (dotted) are less different than are the curves for native raclopride in Figure 2c. The plot of receptor availability (Fig. 3b) for this theoretical ligand is nearly identical to that of native raclopride because the rate constants for endogenous dopamine have not been changed and the majority of occupied receptor sites in a high specific activity injection study are occupied by endogenous transmitter. As with the native raclopride simulation, the residual plot (inset) is entirely negative and peaks early although it reaches a maximum value of only 6.5% of the activated PET signal.

The results of simulating the comparable activation study with an irreversible form of raclopride are presented in Figure 4. The endogenous input function is the same 7 minute step function as in the previous two studies. The compartment concentrations (Fig. 4a) show that the free compartment is completely and irrevocably depleted by 15 minutes, after which the slowly rising total tissue concentration is comprised entirely of bound ligand. The figure is corrected for radioactive decay. The receptor availability plot (not shown) is also similar to previous ones (refer to Fig. 2b). The normalized residuals plot is all negative; it peaks at approximately 10 minutes (inset) and is similar to the native raclopride simulation. However, unlike Figure 2, the normalized residuals for the irreversible ligand never get smaller than 15% of the activated signal within the first 60 minutes of scanning. This phenomenon is borne out by the PET curves in Figure 4b, which show a greater separation between the activated and control curves than in the previous simulations. Entry B-7 in Table I confirms that the sum of squared distances between activated and control curves,  $\chi^2$ , is maximal for this theoretical ligand.

# Sensitivity of results to estimate of endogenous dopamine levels

As discussed elsewhere [Fisher et al., 1995], there is some question as to the appropriate average intrasynaptic dopamine levels for the rest and activated states to input into the model. Examination of published results leads us to estimates of 100 nM at rest and 200 nM during activation. This estimate is in accordance with the intuitively attractive presumption that the receptor operates in an environment whose average endogenous ligand concentration is at or near the Kon and thus equally sensitive to circumstances that cause the level of endogenous transmitter to drop or rise. To ascertain the sensitivity of our results to the particular endogenous concentration levels chosen we ran additional simulations at alternative levels. Section D of Table I indicates that based on calculation of the χ<sup>2</sup> statistic, resting average concentrations of 500 or 1,000 nM dopamine in the vicinity of the post-synaptic receptor would enhance the effect of activation compared with the assumed level of 100 nM. On the other hand, if the average dopamine concentration were only 20 or 40 nM but the Kp were to remain unchanged, the detection of activation with raclopride would be more difficult ( $\chi^2 = 1,591$  or 2,151, respectively). If the endogenous dopamine is set at 40 nM the  $\chi^2$  value can still be increased by more than three times by using an irreversible version of raclopride (data not shown). Because the Kon of dopamine was held constant at 100 nM throughout all the simulations summarized in Table I, stipulating higher levels of dopamine dictates that a higher percentage of receptors be occupied at all times. Note that the modified levels of receptor occupancy are reflected in column 6 of section D of Table I. Figure 5 displays the simulation corresponding to entry 4 in section D of Table I. The endogenous input to the system is a step function that steps up to 2,000 nM for the 7 minutes of activation and then returns to a basal level of 1,000 nM of endogenous chemical after the activation is completed. The compartmental curves (not shown) look very similar to the native raclopride simulations. Nevertheless, there are slight differences from the curves in Figure 2. The receptor availability plot in Figure 5a shows graphically the effect of the endogenous concentration level being 10 times the equilibrium dissociation constant of dopamine, Ken. According to the definition of equilibrium dissociation constant, approximately 91% of all receptors must be occupied at rest. Recall that Farde et al. [1989] have already determined with raclopride that the available



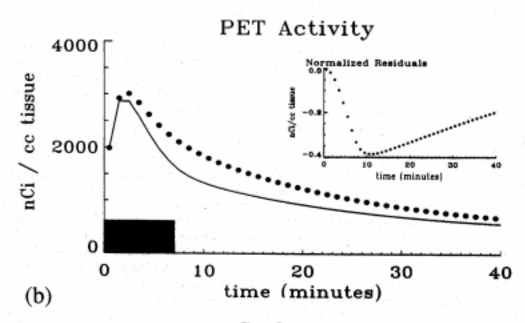


Figure 5.

Simulation of labeled ligand, [11C] raclopride, in brain tissue during the first 40 minutes of a cognitive activation study assuming higher levels of resting and activated dopamine than in previous figures. Concentration curves by compartment (not shown) are reminiscent of Figure 2a. (a) Availability of dopamine (D2) receptor sites (pmol/mf) remaining during activation of endogenous dopamine. After initial drop in available sites due to 7 minutes of activation, receptor availability returns to approximately 10% of total receptors (i.e., simulation assumes 90% occupation of receptors at rest). Horizontal line at B<sub>max</sub> represents the theoretical maximum num-

ber of available receptor sites, which is specified here as 295 pmol/ml. (b) Predicted PET activity curves (nCi/cc) for study (solid curve), which include cognitive activation for the first 7 minutes and for resting study with no activation (filled circles). Activation refers to increased endogenous dopamine level from 1,000 to 2,000 nM and is indicated by double thickness solid bar on x-axis. Inset: Normalized residuals ([activated-rest]/activated) corresponding to the curves in b. All figures except for the normalized residual plot are corrected for radioactive decay.

number of D2 receptors at steady state (B'max) is 28 nM. To satisfy both of these criteria, the total concentration of receptors (Bmax) for this simulation had to be specified as 295 nM. Although the compartmental concentrations (not shown) are similar to the values in Figure 2, there is slightly less receptor binding of the

labeled ligand in this case—presumably due to increased competition with more endogenous dopamine. Less labeled ligand in the bound compartment means more in the free compartment, particularly during the critical activation period. Like Figure 2, the PET curves for control and activated in Figure 5b converge by 40 minutes; however, the differences between them are greater during those 40 minutes than for the simulations with 100 nM endogenous dopamine. The shape of the residuals plot (inset) is similar to the plot of raclopride residuals but the scale in Figure 5 is larger.

#### Flow effects versus neurotransmitter activation

In practice, it is very likely that the tasks that will be employed to raise levels of neuromodulatory chemical will also cause an increase in local blood flow to the brain region of interest. Of course, this is the basis for conventional activation studies with PET. Thus, it is important to be able to demonstrate that we will be able to distinguish flow effects from neurotransmitter activation. As we discuss elsewhere [Fisher et al., 1995], we expect that increased local flow will lead to increased delivery of injected ligand and therefore a larger PET signal over time compared with the resting case. Such a change would be opposite in direction to the effect of competition between endogenous transmitter and exogenous radioligand. Figure 6 confirms our intuition for an [11C]raclopride study. We compared the theoretical case of increased blood flow during the "activation" period of the study but no neurotransmitter activation during that time to the resting case. Figure 6a shows that the receptor availability during the study is a constant 28 pmol/ml (an initial dip in the curve is the normal effect on receptor sites of a high specific activity injection). Increased blood flow transiently was simulated as a threefold increase in the blood flow parameters, K1 and k2, from 0 to 7 minutes. These conditions yield a PET curve (Fig. 6b) that is consistently higher and sharper than the control (filled circles). Significantly, the plot of the residuals (inset) in Figure 6 is quite different in shape and in sign (i.e., + or -) from plots for all the simulations we have examined (Figs. 2-5) that include neurotransmitter activation. Rather than peaking at or near the time at which the activation task is discontinued, the residuals in this simulation are maximal at time 0 and are fairly constant after 10 minutes.

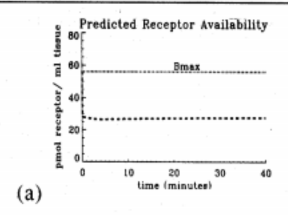
# Model simulation for uptake of [11C]CFT at the transporter site with activation of endogenous dopamine

We also investigated whether the activation of the dopamine system might be detectable at the site of dopamine reuptake by its transporter molecule on the presynaptic terminal. An activation study with the

transporter ligand [11C]CFT and a 15 minute activation period is simulated in Figure 7. Kinetics of the transporter ligand were satisfactorily modeled with the same receptor model used for raclopride (Fig. 1), specifically including some nonspecific binding and some reversible specific binding. The activation was modeled, conservatively, as inducing a step in endogenous dopamine concentration from 30 to 90 nM in the neighborhood of the dopamine transporter site. These levels of dopamine are lower than in other simulations because the transporter site may be extrasynaptic. Although the total curve in Figure 7a is primarily made up of bound radioligand, there is still a noticeable pool of free radioligand that is affected by the activation. Note the drop in free concentration and concomitant rise in binding when activation is turned off at 15 minutes. As in previous simulations, the free endogenous concentration at rest is set approximately equal to the equilibrium constant for endogenous dopamine, Kon. Thus, slightly less than half of the receptors are occupied after activation is turned off (refer to section A, Table II). Figure 7b indicates that Bms has been set at 1,358 nM. The predicted PET curves are broader than those predicted for raclopride and the difference between the control and activated [11C]CFT curves is less impressive for comparable activation times, hence our decision to consider longer activation times for CFT. Nevertheless, it is important to observe that the normalized residuals plot (inset) displays the same characteristic (negative) peak when the activation is terminated as do the simulations for raclopride and related ligands. The residuals for the 15 minute activation paradigm peak at approximately 11 percent of the activated curve.

# Optimal characteristics for CFT-like ligand

Section B of Table II contains  $\chi^2$  results for simulations with theoretical ligands that derive from the native [\$^{11}\$C]CFT. The activation length in this series of tests was held at 7 minutes, beginning at time 0. The only change to the CFT binding parameters that increases the value of the  $\chi^2$  measure is significant reduction in the dissociation rate constant,  $k_{\text{off}}$ . When  $k_{\text{off}}$  is reduced by at least an order of magnitude the resulting simulations yield roughly 20% improvement in  $\chi^2$  value compared with the native CFT simulation (listed in section A, Table II). The optimal quality of the irreversible ligand is reminiscent of our findings with raclopride.



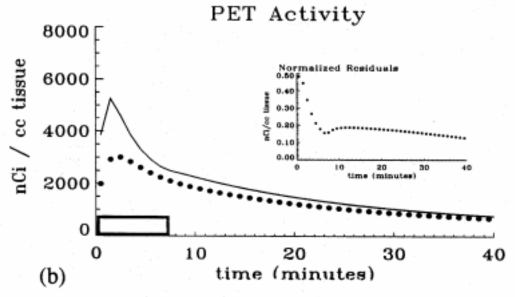


Figure 6.

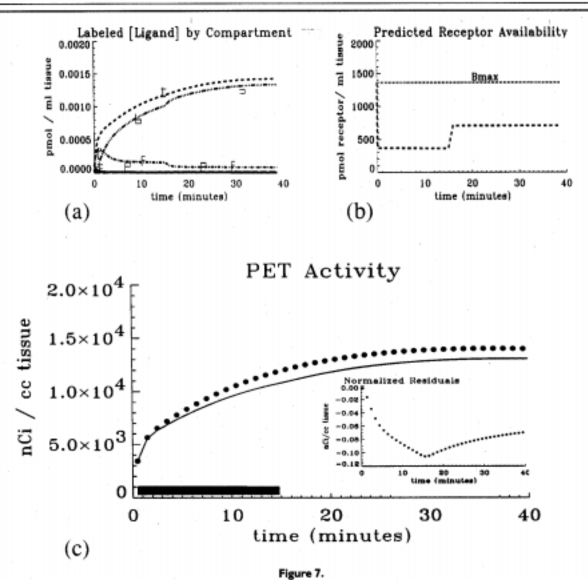
Simulation of labeled ligand, [11C]raclopride, in brain tissue during the first 40 minutes of an "activation" study that assumes a threefold increase in plasma-tissue transport parameters, K<sub>1</sub> and k<sub>3</sub>, for the period 0–7 minutes, but no increase in levels of activated dopamine relative to rest. (a) Plot of availability of receptor sites. Near constancy of the receptor availability at 50% of B<sub>max</sub> confirms that there is no modulation of endogenous transmitter in this

simulation. (b) Dynamic PET activity curve for resting state (filler circles) is plotted on same axis as PET activity for increased blood flow but no change in endogenous dopamine during 0–7 minute activation period (solid curve). Inset: Normalized residuals ([activated-rest]/activated) corresponding to the PET curves. All figure except for the normalized residual plot are corrected for radioactive decay.

# Optimization of activation paradigm

Having found what appears to be a crucial characteristic of a good receptor/activation ligand, namely, irreversibility, we sought to examine the role of the start and stop times for the activation task in order to enhance detectability. Figure 8a displays the dependence of detectability of activation (via the  $\chi^2$  statistic) on length of activation for raclopride-derived ligands.

Specifically, we plotted this relationship for the native compound as well as the "low  $k_{cd}$ " and irreversible versions of raclopride. As one might expect, in all cases the  $\chi^2$  parameter grows as the length of activation grows. There appears to be a definite phenomenon of diminishing returns in  $\chi^2$  for the more irreversible ligands but less so for raclopride itself. Each of the activation tasks represented in Figure 8 was specified to begin at the time of injection. At first, it may seem



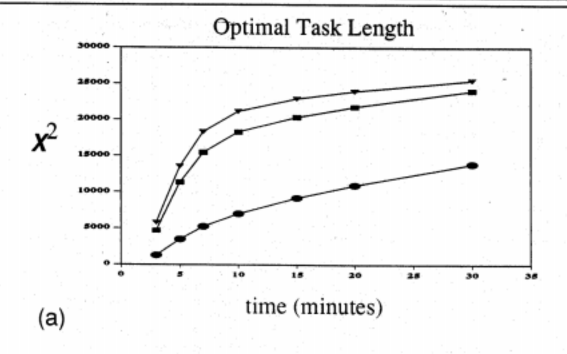
Simulation of first 40 minutes of cognitive activation and resting studies with the labeled ligand, ["C]CFT, in brain tissue. (a) Simulation of total concentration (pmol/ml) of ligand in brain tissue (t) in response to 15 minute activation. Total curve is the sum of the ligand in each individual compartment (p, plasma; f, free; b, bound, n, nonspecific). (b) Receptor availability rises to roughly half the B<sub>max</sub> value once activation is discontinued. (c) Predicted PET activity curves (nCi/cc) for study (solid curve), which include

cognitive activation for first 15 minutes and for resting study with no activation (filled circles). Activation refers to increase in extrasynaptic endogenous dopamine level from 30 to 90 nM and is 
indicated by half-thickness solid bar on x-axis. Inset: Normalized 
residuals ([activated-rest]/activated) corresponding to the PET 
curves. Normalized residuals are largest at about 15 minutes, 
coinciding with the end of activation. All figures except for the 
normalized residual plot are corrected for radioactive decay.

simple to design a cognitive or motor task that could be sustained for longer than 15 minutes. However, if one considers the possibility of training effects coming into play, it might not be reasonable to model increased neurotransmitter levels as a step function in the case of lengthy tasks. For these reasons, our standard simulation protocol for comparing ligand characteristics and task delay times incorporated a 7

minute task. Still, it is clear from Figure 8a (for raclopride) that regardless of task length, irreversibility is a highly desirable characteristic.

Figure 8b displays the comparable plots of the  $\chi^2$ statistic versus activation length for CFT-derived ligands. Once again, increased task length correlates with larger separation between rest and activation. However, for native CFT this relationship is linear;



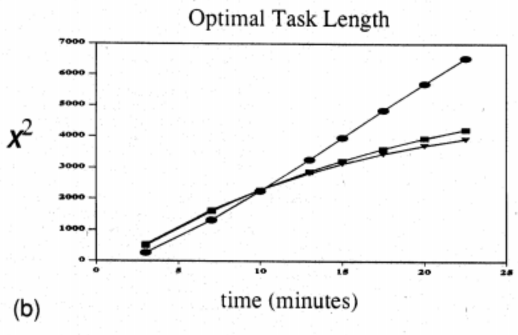


Figure 8

(a) Plot of  $\chi^2$  value (measuring difference between activated and control curves) for increasing lengths of activation. All activations begin at time 0. Individual curves correspond to the ligand [ $^{11}$ C]raclopride and its theoretical derivatives—native raclopride ( $\bullet$ ), low  $k_{cd}$  ( $\blacksquare$ ), and irreversible ( $\blacktriangledown$ ). (b) Plot of  $\chi^2$  values for

increasing lengths of activation, each beginning at time 0. Curves correspond to the ligand [ $^{11}$ C]CFT and its theoretical derivatives—native CFT ( $\bullet$ ), low  $k_{eff}$  ( $\blacksquare$ ), and irreversible ( $\blacktriangledown$ ). Note: the scales in a and b are arbitrary because of how the variance is calculated and thus the two plots cannot be compared directly.

there is no apparent point of diminishing returns. The more irreversible forms of the transporter ligand are more similar to the plots in Figure 8a in that they begin to saturate (albeit slowly) at long activation times. Although the scales are somewhat arbitrary for  $\chi^2$ (based on our model of the error in the PET signal) and the two graphs for the respective families of ligands cannot be compared directly, there is some suggestion that CFT would be a preferable ligand for use with long activation paradigms because the separation of the PET curves will continue to grow even at long times.

For a given task length (7 minutes) we also examined the effect on  $\chi^2$  of delaying the beginning of the task for some time after the injection of radioligand and start of scanning. In contrast to the effect of activation length, the effect of task delay is essentially the same for raclopride and CFT. Figure 9a (raclopridederived ligands) and Figure 9b (CFT-derived ligands) both indicate that delaying the activation task by as little as 1, 2, or 3 minutes from the moment of the injection can be expected to reduce the difference between activation and control curves. Longer delays lead to progressively smaller  $\chi^2$  values. Finally, it must be emphasized that delaying onset of the task by just 10 minutes was enough to obliterate completely the effect of neurotransmitter activation for any of the ligands examined.

#### DISCUSSION

# How is what we are proposing different from other studies?

There has been some recent attention in the field of receptor-ligand imaging with PET directed toward detecting or at least accounting for changes in neurotransmitter levels and their effect on conventional receptor imaging assays. Much of this work was initially motivated by the desire to resolve the controversy over D2 receptor measurements with raclopride and spiperone in schizophrenics [Wong et al., 1986; Farde et al., 1987; Andreasen et al., 1987]. In contrast to the current study, Logan et al. [1991] viewed variable levels of dopamine as a potential source of error in attempts to estimate D2 receptor density. Nevertheless, through extensive simulations they contributed greatly to our understanding of the interaction between exogenous and endogenous receptor ligands. In that study, a model similar to the one presented here was used to assess the impact of either different constant levels or transient changes in endogenous

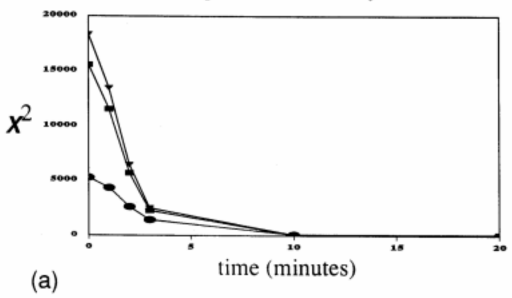
dopamine on [18F]NMS measures of receptor availabil ity and density. The authors concluded that the halftime for a radioligand in the tissue (along with its equilibrium constant) is an important factor in determining its sensitivity to changes in endogenous neuro transmitter levels. In turn, they claimed that halftime is a function of the tissue to plasma rate constant, K and thus cited it as an important parameter in deter mining the sensitivity of the ligand to changes ir endogenous neurotransmitter levels. In general, we agree that the issue of competition between an endog enous chemical and a short-lived exogenous radioli gand is a kinetic one. However, because this is a kinetic rather than an equilibrium phenomenon, we believe our results show that the equilibrium constant Kr., has little or no effect on our ability to detec competition while the rate of dissociation, koff, it highly correlated with the degree of separation be tween the activated and control curves. This issue it explained more fully below.

The researchers from Brookhaven National Lab followed their simulation studies with a series o studies focusing on inducing changes in endogenou neurotransmitter levels pharmacologically [Dewey e al., 1991, 1992, 1993]. Researchers from Hammersmitl Hospital have observed neuromodulation indirectly by imaging blood flow changes with PET during cognitive tasks before and after pharmacologic inter vention [Friston et al., 1992]. Studies have been re ported that tried to look at either the effects of activating the dopamine system via SPECT imaging o [123I]IBZM [Childress et al., 1993] or the release of endogenous opioids with PET [Duncan et al., 1992] The former study proposed using a reversible liganand commencing activation 1 hour post injection and the latter study proposed activating 30-40 minute post injection of [11C]diprenorphine. Our simulation results suggest that these experimental designs ma have been suboptimal for detecting activation. W hope to be able to model, detect, and quantify th average change in endogenous transmitter levels b PET to investigate the connection between specifi brain functions and activity of neuromodulatory neu rons. In order to predict whether such an assay woulbe possible, and even optimal, we turned to mode simulation studies.

# Can we detect activation of a particular neurotransmitter system with PET?

Our simulations have not accounted for noise that i certain to be included in the actual control an

# Optimal Task Delay



# Optimal Task Delay

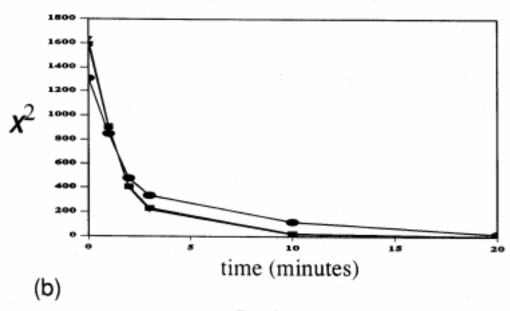


Figure 9.

(a) Plot of  $\chi^2$  value (measuring difference between activated and control curves) for increasing lengths of delay between time of injection and time of activation. All activations are 7 minutes long. Individual curves correspond to the ligand [ $^{11}$ C]raclopride and its theoretical derivatives—native raclopride ( $\bullet$ ), low  $k_{of}$  ( $\blacksquare$ ), and

irreversible ( $\nabla$ ). (b) Plot of  $\chi^2$  value for increasing delay time activation length is 7 minutes. Curves correspond to the ligani [ $^{11}$ C]CFT and its theoretical derivatives—native CFT ( $\Phi$ ), low  $k_e$  ( $\blacksquare$ ), and irreversible ( $\nabla$ ). Note: the scales in a and b are arbitrar; and cannot be compared directly.

activated curves. Nevertheless, based on the difference in simulated curves presented above we believe there is reason to be optimistic that endogenous neurotransmitter activation could be detectable. Our PET camera, (PC4096, Scanditronix) has a typical signal to noise ratio of 10 at the peak concentration (10% error) whereas the differences conservatively predicted herein between activation and control using native raclopride were on the order of 20%. Although the results of simulations with the D2 receptor ligand (raclopride) predicted more dramatic effects of activation than did our simulations of the dopamine transporter ligand, there seems to be sufficient change in the predicted PET curves for [11C]CFT to warrant pilot studies with either ligand given an appropriate task. It is also apparent that detectability can be enhanced by properly timing the activation task in accordance with the dynamics of the radioligand in the free space and by choosing a ligand whose binding is nearly irreversible.

# Why is an irreversible ligand desirable?

Ideally, the labeled injectate in a receptor activation study should create a large pool of free label during the activation period while the competition for binding sites between exogenous and endogenous ligand is occurring. This transient competition with endogenous chemical should produce a PET curve that is lower than the corresponding control PET curve during the activation interval. If at the moment the activation were terminated, the remaining free pool of hot ligand could be eliminated and the bound ligand prevented from dissociating from the receptor, then the separation of the two curves would be maintained, notwithstanding the ongoing decay of signal from the tissue. This is the scenario for maximal detectability of neurotransmitter activation and it corresponds closely to the case of the irreversible version of raclopride  $(k_{off} = 0)$  we have simulated. Figure 10 displays the compartmental curves for native raclopride, a high kon version, and an irreversible version of the ligand, respectively. As demonstrated in Figure 10, during the first 7 minutes of the activation study the free pool of labeled ligand (cross-hatched area under the free curve) is large for the native and irreversible raclopride cases (Fig. 10a,c) and inadequately small for the case of a high kon ligand (Fig. 10b). Once the activation is ended at 7 minutes, normal levels of neurotransmitter prevail and binding of hot ligand can resume unfettered. However, because the dissociation of li-

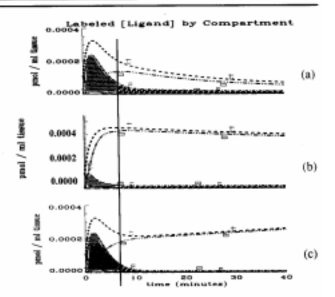


Figure 10.

Comparison of labeled ligand concentrations by compartment (t, total; p. plasma; b. bound; f. free) for the three ligands examined in Figures 2a, 3a, and 4a. (Native raclopride, a high kan version of raclopride, and an irreversible version of raclopride are depicted in Fig. 8a, b, and c, respectively). The vertical line through all three plots indicates the termination of cognitive activation, which raises endogenous dopamine concentration from time 0 to 7 minutes of the study. The area under the free ligand concentration curve is filled in to indicate the size of the pool of ligand over the course of the study that is available for binding. To the left of the vertical line (i.e., during activation) the area under the free curve corresponds to the size of the free pool of labeled ligand that is in competition with activated levels of endogenous dopamine for binding to the D2 receptors. To the right of the vertical line is the remaining pool of labeled ligand that remains either to bind to receptors or to leave the system via the plasma after the activation of endogenous dopamine has been terminated. (a) The total curve is comprised primarily of free ligand during activation whereas after activation the free ligand is a smaller but significant part of the total tissue concentration. (b) The free ligand makes up a small fraction of the total during and after activation. (c) The free ligand is a large part of the total during activation, similar to a. After activation, though, almost no ligand exists in the free state, thus ensuring that the total curve remains at a nearly constant level, which was determined by the competition between free labeled ligand and activated endogenous transmitter during the first 7 minutes of the study. All figures are corrected for radioactive decay.

gand is disallowed in the irreversible case, the pool of free ligand is more quickly depleted (diagonally hatched area under curves), the difference between the activated and control PET curves is preserved and thus more easily detectable than in the native raclopride case.

# How to chose the right ligand and the right timing of activation

As we have demonstrated, the likelihood of success of an activation study seems to correlate very closely with the area under the free compartment curve during the time of activation. On the other hand, one may be able to prolong the persistence of a significant free pool via extended activation using a ligand (such as [11C]CFT), with a slower k2 value and a much larger binding capacity (Bmax = 1,358 pmol/ml) than raclopride (Fig. 7). The fact that these results are not intuitive is an argument in favor of the type of simulation study presented here. In all cases, longer activations lead to greater detectability but an experimenter must factor in the ability of a subject to sustain a task, which we have not considered in our simulations. Furthermore, it is important to continue characterizing new receptor ligands (and re-evaluating old ligands) via compartmental modeling and parameter estimation, since our simulations would be meaningless without reliable estimates of receptor model parameters. Regardless of the ligand, though, the finding that the activation task must begin at the same time as the injection and not be delayed seems universal.

#### Spiperone versus Raclopride

The finding that irreversibility of the ligand is optimal for detection of neurotransmitter levels suggests that N-methyl-spiperone, which is often modeled as irreversible, may be preferable to raclopride for studies of dopamine activation. Unfortunately, spiperone has been shown to bind to the serotonin 5HT site and therefore would be of dubious value in trying to establish the relative dopaminergic versus serotonergic qualities of a particular cognitive task.

#### How to account for flow effects of activation?

We have demonstrated (in Fig. 6) that the flow effects typically associated with cognitive activation run counter to the expected effect on the PET curve from increased competition with endogenous chemical. Thus, it is unlikely that increased flow of blood to a region of the brain could be confused with increased concentration of a neurotransmitter due to activation. Nevertheless, this issue brings out an important consideration for the design of experiments. A possible receptor activation study might involve dynamic scanning after injection of an irreversible D2 receptor ligand and a motor task that could be expected to

activate the basal ganglia unilaterally. In this way, the ipsilateral nonactivated brain region could be used to generate the control curve in the same study. However, in such a case the control region would have to be modeled with resting values for transport parameters, K1 and k2, while the activated, contralateral region would be correctly described by parameters adjusted for increased regional cerebral blood flow (rCBF). Thus, it might be necessary to perform additional PET studies with a blood flow tracer (e.g., H<sub>2</sub>O) under the same activation conditions in order to gauge the increase in rCBF between the two sides during activation. If a lateralized task were not possible, a stepwise procedure for identifying increases (or decreases) in rCBF during activation via H25O imaging followed by model-based simulation of control curves might be necessary. In outline form, the general procedure for any task might be as follows: 1) perform resting and activation PET receptor studies that yield "activation" and "control" curves; 2) using a model—sans activation—fit a curve to the control data yielding model parameters for no rCBF change and no synaptic activation; 3) adjust the blood flow parameters (K1 and k2) from step 2 in accord with the flow changes detected via H25O PET; 4) simulate the theoretical control curve corresponding to increased (or decreased) CBF and no change in synaptic activity; and 5) compare the curve in step 4 with the activation curve from step 1. If need be, multiple (1 minute) rCBF measurements could be made in order to construct a piecewise continuous profile of the blood flow throughout a 7 minute activation.

The second important point to be taken from the examination of blood flow effects on the simulations of receptor activation is the appearance of the normalized residual plot. The residual plot for the simulation of increased K1 and k2 during the first 7 minutes of the study (Fig. 6) is dramatically different from the residual plots for simulations containing neurotransmitter activation. In the residual plot of the same figure (inset), the residuals at all times are positive, reflecting the increased PET activity characteristic of increased flow, not increased competition with endogenous ligand. The characteristic (negative) peak in the residuals that corresponds closely to the time of task termination is absent. When we followed the scheme outlined above for comparing an activated PET curve with its proper control, the resulting residual plot displayed the characteristic minimum at or near the end of the activation period (data not shown).

# Use of residual plot as confirmation of augmented model

In all activation cases the normalized residuals consisted of a unimodal curve that was necessarily 0 at time 0, a minimum at approximately the time of cessation of activation, and asymptotically approached 0 at long time. We suggest that the appearance of such a peak in experimental data might be considered a confirmation of the simple model proposed in this paper to describe the contribution of change in endogenous neurotransmitter levels to the conventional receptor-binding model. Furthermore, we acknowledge that the reliance on  $\chi^2$  values alone as a means of detecting differences in curves due to activation is limited and ignores temporal information in the data. We therefore propose a two-part process for identifying neurotransmitter activation: sufficiently high  $\chi^2$ values and the identification of the single peak in the normalized residuals at or near the end of activation.

For very high kon simulations we note that the (minimum) peak in the residuals coincided exactly with the end of activation whereas in the case of slower  $k_{on}$  values, the peak was slightly delayed. This behavior agrees with our intuition that the kon value for the hot ligand determines how fast the system can respond to the abrupt drop in endogenous neurotransmitter level. Neglecting the effect of noise for the moment, it is exciting to speculate that for a wellcharacterized ligand whose kom is known one might use the residual plot to identify the end of activation of one particular neuromodulatory system among many systems that are activated by a complex cognitive task. Alternatively, in the case of a kon that is not. well known, but a task that is well characterized, the time of the residuals peak in relation to the known end of activation time might be used to infer the kon value.

#### B'\_v versus B\_v

Inherent in the standard receptor model is the assumption that endogenous neurotransmitter levels are in steady state and therefore can be ignored when modeling the availability of binding sites. As such, the number of available receptor sites prior to the injection of radioligand, B'max is taken to be a constant property of the system. That is, B'max is a parameter of the standard receptor model to be determined by fitting model curves to experimental data. The augmented receptor model, which includes the effect of nonsteady levels of endogenous neurotransmitter, must consider all possible binding sites for ligand and

endogenous competitor in the tissue. Thus, the mode for analyzing receptor activation studies (Equation: 1–8) has a parameter B<sub>max</sub>. A possible advantage of the studies proposed herein is that they may lead to ar estimate of the total number of receptors, Bmas, which is not available from analysis of a conventional recep tor imaging study with PET. Similarly, the mode presented introduces two other parameters not found in the standard receptor model. Fen(0) and A are respectively, the average basal level of neurotransmit ter in the free space and the change in level during activation. In theory at least, it should be possible to estimate these parameters from the type of recepto: activation study we have proposed, provided that enough of the remaining model parameters are sufficiently well known or could be constrained based or prior knowledge.

#### CONCLUSIONS

To answer the question of whether or not PET could be used to detect changes in neurotransmitter levels we constructed a model of receptor binding in the presence of endogenous neurotransmitter. Using pub lished rate constants, we were able to simulate and compare predicted PET curves assuming a high spe cific activity injection of [11C]raclopride into subject either at rest or in a state of dopamine activation. We found that the PET curves for activated dopaming levels were predicted to be different from control Additionally, these differences, and hence detectabil ity, could be maximized through the choice of ligans characteristics and the proper activation task. We now believe that PET studies designed to detect cognitive or motor activation of a specific neurotransmitte system are worth considering. Such experiments could certainly be optimized by timing activation tasks to coincide with the presence of a large pool of radioli gand in the free compartment (always starting at time 0) and through the use of irreversibly binding receptor ligands.

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