Fluoroethoxy-Benzovesamicol, a PET Radiotracer for the Vesicular Acetylcholine Transporter and Cholinergic Synapses

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KEY WORDS autoradiography; cholinergic; dosimetry; imaging; PET; synapse; vesamicol

ABSTRACT Loss of cholinergic transmission in the cortex and hippocampus is a characteristic feature of Alzheimer’s disease, and visualization of functional cholinergic synapses in the brain with PET could be a useful method for studying this degenerative condition in living humans. We investigated [18F]fluoroethoxybenzovesamicol, (−)-[18F]FEOBV, (−)-(2R,3R)-trans-2-hydroxy-3-(4-phenylpiperidino)-5-(2-[18F]fluoroethoxy)-1,2,3,4-tetralin, a high affinity positron emitting ligand for the vesicular acetylcholine transporter, as a potential in vivo cholinergic synapse mapping agent. Rodent biodistribution, dosimetry, stereospecificity of biological effects, pharmacologic blocking studies, in vivo rodent brain autoradiography and metabolites were examined. (−)-[18F]FEOBV brain uptake following intravenous injection was robust, with 2.65% dose/brain in mice at 5 min, and the regional localization matched the known distributions of presynaptic cholinergic markers at later times. Both the cholinergic localization and curare-like effects of FEOBV were associated with the “(−)”-enantiomer exclusively. (−)-[18F]FEOBV regional brain distribution in rodents was changed little by pretreatment with haloperidol, (−)-3-PPP, or E-2020, indicating FEOBV, unlike other vesamicol analogs, did not interact in vivo with dopamine or acetylcholine receptor systems. Autoradiography of rat brain 3 h following i.v. injection of (−)-[18F]FEOBV showed high localization in brain areas rich in presynaptic cholinergic elements. Metabolic defluorination in rodents was modest, and analysis of brain tissue following tracer administration found FEOBV as the only extractable radioactive species. (−)-[18F]FEOBV dosimetry calculated from rat data estimate 10 mCi doses can be given to humans. These studies show FEOBV maps cholinergic areas with high specificity in vivo, and may provide a noninvasive means to safely and accurately gauge the functional integrity of cholinergic synapses in man using PET. Synapse 30:263–274, 1998. © 1998 Wiley-Liss, Inc.

INTRODUCTION

A dramatic loss of acetylcholine (ACh) and its metabolically related enzymes from the cortex and hippocampus is a characteristic feature of Alzheimer’s disease (AD) (Perry et al., 1994). In addition, loss of synaptic elements in the cortex and subcortical brain areas is correlated with the severity of cognitive decline in AD (Davies et al., 1987; DeKosky and Scheff, 1990; Masliah et al., 1992a,b; Samuel et al., 1994; Scheff et al., 1990; Terry et al., 1991). Some studies have found synaptic and neuronal loss to be even more strongly correlated with AD symptoms than the senile plaques and neurofibrillary tangles that are the diagnostic hallmarks of the disease (Gomez-Isla et al., 1997; Masliah et al., 1992b; Svennerholm and Gottfries, 1994; Terry et al., 1991). The synaptic pathology of AD involves neurons of multiple transmitter types, but loss of cholinergic presynaptic function in the neocortex appears to be a crucial component in the development of visuospatial deficits and impaired working memory that occurs with the disease (Perry et al., 1994). Several current pharma-

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cologic treatments for AD are aimed at enhancing cholinergic neurotransmission in the brain (Brennan, 1997). The root causes of neurodegeneration in AD are not understood (Izhaki, 1994), and a definitive diagnosis of disease can be made only after death. The ability to measure cholinergic synapses in the living human brain could be an important investigative tool for assessing neuronal damage, or for monitoring the efficacy of future neuron-sparing therapies for this devastating condition.

For these reasons, considerable efforts have been focused on developing radiotracer imaging techniques to visualize elements of presynaptic cholinergic function. One approach has employed radiolabeled ligands or substrates of acetylcholinesterase (AChE) as markers of cholinergic nerve terminals (Irie et al., 1994; Kilbourn et al., 1996; Pappata et al., 1996). Another method, and the basis for the present work, targets the vesicular ACh transporter (VAChT) in cholinergic terminals with radiolabeled forms of the drug vesamicol (Altar and Marien, 1988). VAChT is a glycoprotein located in cholinergic synaptic vesicle membranes which draws cytoplasmic ACh into the vesicles. Vesamicol binds on VAChT to a so-called vesamicol receptor site that is noncompetitive with ACh binding and, therefore, presumably not affected by changes in endogenous ACh levels. VAChT tissue distribution is highly correlated and uniquely distributed with other proteins involved in the presynaptic metabolism of ACh, including choline acetyltransferase (ChAT), high affinity choline uptake protein, and acetylcholinesterase. The close association of VAChT with other cholinergic markers was recognized originally from vesamicol binding studies (Altar and Marien, 1988), and has been confirmed recently using hybridization binding techniques with cloned VAChT gene cDNA probes (Erickson et al., 1994, 1996; Roghani et al., 1994), and VAChT immunocytochemistry (Gilmor et al., 1996). It has been discovered that the human VAChT gene lies nestled within the ChAT gene locus and these two genes are co-expressed in a tightly coupled fashion. A mounting body of evidence points to VAChT as being a very specific marker of cholinergic terminals. General recognition of this fact, and the discovery of many potent analogs based on the vesamicol template, has spurred the search for suitable radioligands for measuring VAChT in vivo (Efange et al., 1992; Parsons and Rogers, 1993; Efange et al., 1993, 1994, 1995a; Iung et al., 1996; Mach et al., 1997).

We studied a class of analogs called benzovesamicols, discovered by Rogers et al. (1989), and identified a number of novel radioiodinated benzovesamicol analogs which distribute in vivo in patterns that correlate closely with the densities of marker proteins of the cholinergic nerve terminal. Analogues with radiodine, carbon-11, or fluorine-18 labels at the 5 position of the 2-hydroxy-3-N-(4-phenyl)piperidino-tetralin framework, which we term 5-substituted-benzovesamicols (other investigators identify the same compounds as 4-benzovesamicols using a different nomenclature; see Parsons and Rogers, 1993) have shown particularly favorable imaging properties (Iung et al., 1990; Kilbourn et al., 1990; Mulholland et al., 1991a,b; Ingvar et al., 1993). Human SPECT (single photon emission computed tomography) imaging studies with the [123I]-labeled benzovesamicol analog 5-BVM show decreases in tracer binding in several brain regions of AD patients (Kuhl et al., 1994, 1996). These studies offer encouragement that appropriate benzovesamicol tracers can map cholinergic neurons and gauge the functional integrity of cholinergic synapses in man.

With the aim to extend human benzovesamicol imaging studies to a higher level of precision with positron emission tomography (PET), we developed a novel fluorine-18-labeled agent, 5-fluoroethoxy-benzovesamicol (FEOBV, Fig. 1). The synthesis of [18F]FEOBV (Mulholland et al., 1993) and an examination of its kinetics in the perfused rat heart (Degrado et al., 1994) were reported previously. The present communication examines distribution and pharmacologic properties of FEOBV in animals, compares them with those of related agents, and covers evaluation steps of FEOBV preliminary to its introduction as an investigational cholinergic synaptic tracer for human PET imaging studies.

MATERIALS AND METHODS

(--)-[18F]FEOBV was labeled in a “one-step” nucleophilic radiofluorination procedure from “no-carrier-added” [18F]fluoride ion and (--)-(2R,3R)-5-(2-tosyloxyethoxy)-benzovesamicol (Mulholland et al., 1993). The specific activity of the final tracer product at the end of synthesis exceeded 3 Ci/µmol (range 3–17 Ci/µmol, average 8 Ci/µmol) following HPLC purification, and the radiochemical purity was greater than 99%, as assayed by reverse phase HPLC. The inactive “(+)-” enantiomer of FEOBV was labeled by the same route to the same quality specifications as (--)FEOBV, starting from resolved (+)-5-(2-tosyloxyethoxy)-benzovesamicol.

Experiments with living animals were performed in accordance with the United States Department of Agriculture Animal Welfare Act.

Mouse biodistribution studies

Female CD-1 mice weighing 23–34 g were anesthetized lightly with diethyl ether and given tail vein injections of 7–600 µCi of (--)-[18F]FEOBV. The animals then resumed normal activity until the times of sacrifice, which ranged from 5 min to 15 h postinjection. Mice were killed under ether anesthesia by decapitation and tissues were quickly excised, weighed, and counted for radioactivity in an auto gamma counter. Samples from brains included striatum, whole cerebral cortex, hippocampus, hypothalamus, and cerebellum.
The results were corrected for radioactive decay and expressed in terms of percent injected dose per organ (%dose/organ), percent injected dose per gram of tissue (%dose/g), normalized to an animal weight of 25 g. Tissue concentration ratios were calculated from tissue pairs for each animal and then averaged together to yield tissue-to-cerebellum ratios. Each data point represented the mean ± standard deviation of values from at least three animals. Data were analyzed for significance using unpaired two-tailed t-tests.

**Pharmacologic pretreatment studies**

Female CD1 mice (n = 3 per data point) were given interperitoneal injections of either haloperidol (10 mg/kg), (+)-3-PPP (20 mg/kg), or E-2020 (donepezil) (10 mg/kg) 30 to 60 min prior to FEOBV radiotracer administration (30 µCi, ∼4 pmol FEOBV). Groups of mice were sacrificed at 5, 45, and 240 min posttracer and brain regions were dissected and counted for radioactivity.

**Rat biodistribution for dosimetry**

Male and female Sprague Dawley (SD) rats weighing 215–275 g were injected with 50–120 µCi of (−)-[18F]FEOBV via femoral vein while anesthetized with ether, and then allowed to recover and resume normal activity. Groups of animals were sacrificed by decapitation at 5, 15, 30, and 120 min following injection, and tissues and fluids were excised and counted for radioactivity in an auto-gamma counter. The radioactivity remaining in the carcass after removal of samples was measured in a dose calibrator. Values were corrected for decay and calculated as described above for mouse experiments. Each data point in the rat biodistribution data represents the mean ± standard deviation of at least four animal values, except for sex organs, for which n = 2–3. Dosimetry data were calculated using the rat biodistribution data, following the MIRD formalism (Loevinger and Berman, 1976). The percent administered dose per organ values were modified to reflect the different proportions of organ to total body mass in rat and man (Roedler, 1980). Residence times were obtained by integration under the organ time vs. activity curves, with the effective halflife of (−)-[18F]FEOBV assumed to be equal to the physical halflife of [18F] for times exceeding the last data point. Residence times were entered into the MIRDose2 program (Watson et al., 1988) for the generation of absorbed doses to selected target organs per unit of administered activity.

**Rodent brain and blood metabolite assays**

Groups of three female CD-1 mice (24–30 g), under ether anesthesia, were injected intravenously with ∼1 mCi of [18F]FEOBV. They were allowed to recover and resume normal activity until time of sacrifice at 30 and 90 min following tracer injection. The whole brains of the three animals were removed, pooled together, and homogenized in an equal volume of pH 7.6 TRIS/salt buffer (120 mM NaCl, 50 mM TRIS, 5 mM KCl, 2 mM CaCl2, 1 mM MgCl2). The supernatant extract was separated from sediment by low-speed centrifugation and collected. The sediment was then shaken with a 3 ml portion of ethanol, re-centrifuged, and the extract combined with the first. This step was repeated. The combined extracts, which contained more than 97% of the original brain radioactivity, were concentrated to near dryness under reduced pressure, then CH3CN (6 ml) was added and the resulting heterogeneous mixture was concentrated to a volume of 1 ml. The solution was decanted from the precipitated fatty solids, and the fats were rinsed once with 1 ml of fresh CH3CN. The combined final CH3CN solution (overall recovery 59% from brain) was assayed by thin layer chromatography (Analtech silica, 2:1:1:0.2 hexane: CH2Cl2: Et2O: N; Bioscan TLC plate reader; RfFEOBV = 0.75) and HPLC (column: 5 micron C-18 silica, mobile phase: 1.5 ml/min, 3:1:1 CH3CN: MeOH: 20 mM KH2PO4, pH 6.7, serial UV (220 nm) and γ radioactivity (CaF2) flow detectors; kFEOBV = 3.25).

The brain radioactivity from two female rats was examined separately in the same way 30 min following 2.3 and 4.3 mCi (femoral vein) injections of FEOBV. The overall recovery in these cases was 70% and 66%.

Two anesthetized female rats were injected in the tail vein with 6 and 8 mCi of (−)-[18F]FEOBV, respectively. Blood samples (0.1–0.3 mL) were withdrawn from a catheter in the vena cava at 5, 10, 30, and 60 min postinjection, diluted with 1 mL of isotonic saline, and centrifuged. In each sample the plasma was collected and 1 mL of 70% ethanol was added to the cells; the tube was vibrated briefly, centrifuged again, and the supernatant collected and combined with the plasma. The combined extract and residual solids from each blood sample were counted separately, and in cases where the extract recovery was less than 80%, the solids were subjected to a second extraction procedure in which they were washed twice with 1 mL portions of 10% trifluoroacetic acid in methanol. All extracts were concentrated, made basic with triethylamine, and analyzed by TLC.

**In vivo rat brain autoradiography**

Two restrained conscious female SD rats were injected intravenously with (−)[18F]FEOBV (4–9 mCi/rat), and then 5 min and 3 h later they were sacrificed by intravenous pentobarbital, followed by KCl injection. Brains were quickly excised, frozen in powdered CO2 ice, coated with frozen section imbedding medium (Lipshaw, Detroit, MI) and sectioned at –15°C into 20-µm thick slices. Brain slices were air-dried at 85°C on polylysine-treated slides and were exposed to LKB Betamax Ultralfilm for 12 h before development in Kodak D-19. Autoradiograms were analyzed by com-
puter assisted densitometry (MCID System, Imaging Research, St Catherines, ONT). Brain regions identified included areas with anticipated high and low levels of cholinergic markers on the basis of prior rodent and primate studies, and included the interpeduncular nucleus, basolateral amygdala, striatum, dentate gyrus, CA1 and CA3 fields of the hippocampus, frontal cortex, occipital cortex, superficial layers of the superior colliculus, thalamus, and cerebellar cortex.

RESULTS

Rodent brain uptake

(--)-[18F]FEOBV was rapidly taken up by the brain following intravenous injection. There was 2.68 ± 0.62% of the injected dose present in the whole brain of mice at 5 min, and nearly 1.5% remained in the brain at 6 h. In rats the peak brain content was 1.58 ± 0.20% of the administered dose at 15 min. Mouse brain concentrations of (--)-[18F]FEOBV radioactivity were examined at 5, 10, and 45 min, and 3, 4, 6, and 15 h following tracer administration. Time activity curves of [18F]FEOBV in selected brain regions are shown in Figure 2a. Tracer levels peaked within the first 5 min in all brain regions except striatum. Peak striatal accumulation in mice occurred between 45 and 180 min. Clearance was slowest from striatum, fastest from cerebellum, and intermediate from other brain areas. Figure 2b shows tissue [18F]FEOBV localization in mice expressed as a ratio relative to cerebellar concentrations. The regional brain localization ratio in mice at 15 h postinjection, the latest time point examined, was 41:10:12:1 striatum: cortex: hippocampus: cerebellum, respectively. (--)-[18F]FEOBV mouse brain uptake and localization was very similar to that of the closely related analogs [123I/125I]IBVM (Jung et al., 1990) and [11C]MABV (Kilbourn et al., 1990).

In vivo rat brain autoradiography

Rats sacrificed 5 min after tracer administration had fairly uniform (--)-[18F]FEOBV radioactivity levels throughout the brain, but (--)-[18F]FEOBV was distributed heterogenously in animals sacrificed at 3 h, with high localizations in certain tissue areas. Highest radioactivity densities appeared in interpeduncular nucleus, followed by basolateral amygdala, striatum, dentate gyrus, CA1 field of the hippocampus, CA3 field of the hippocampus, frontal cortex, occipital cortex, superficial layers of the superior colliculus, thalamus, and cerebellar cortex. This rank order is consistent with the levels of ChAT (Fuhrmann et al., 1985; Tucek, 1988) and high affinity choline transport-hemicholinium-3 binding (Bekenstein and Wooten, 1989; Happe and Murrin, 1993; Vickroy et al., 1984).
**In vivo cholinergic specificity and lack of interaction with dopamine neurons or sigma sites**

The tissue-to-cerebellum ratios of \((-\)\([18F]\)FEOBV in various brain regions of the mouse at 6 and 15 h postinjection showed close correspondence to the reported levels of ChAT and AChE and \([3H]\)Hemicholinium-3 binding in the same regions in rodents (Table I).

To investigate possible binding interactions of \((-\)\([18F]\)FEOBV with dopamine D2 or \(\sigma\) receptors, we examined the effects of large pretreatment doses of the potent D2 and \(\sigma\) ligand haloperidol, the \(\sigma\) ligand \((-\)\([3\)-PPP (3-(3-hydroxyphenyl)-N-propylpiperidine, Koe et al., 1989), or E-2020 (donepezil, Aricept); a \(\sigma\) ligand and AChE inhibitor, Brennan, 1997; Koe et al., 1991), on mouse brain distribution of the tracer. None of the pretreatments produced changes in the tissue/cerebellum ratios that were significantly different (\(P < 0.05\)) from 5, 45, or 240 min control values (Table II). Haloperidol pretreatment reduced the total \((-\)\([18F]\)FEOBV brain uptake by 20–30%, and blood activity by 25–50% at 5 min and 45 min following tracer administration. By 240 min, total brain uptake and blood activity in the haloperidol group was the same as for controls, and striatal uptake of \((-\)\([18F]\)FEOBV was elevated 36% relative to controls. No significant differences from control data were seen in the \((+)\)PPP and E2020 groups’ total brain uptake, regional brain uptake, or blood levels.

We examined the stereospecificity of \([18F]\)FEOBV tissue uptake by labeling the \((+)\)dextrorotatory enantiomer, \((+)\)\([18F]\)FEOBV and comparing its biodistribution in the mouse with \((-\)\([18F]\)FEOBV at 5, 45, and 180 min. Both isomers showed fairly homogeneous

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**Table I. Comparison of \([18F]\)FEOBV relative brain concentrations in vivo with cholinergic markers in rodents**

<table>
<thead>
<tr>
<th>Brain region</th>
<th>((-)([18F])FEOBV</th>
<th>((-)([18F])FEOBV</th>
<th>ChAT activity*</th>
<th>AChE activity*</th>
<th>([3H])Hemicholinium binding**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>((+))PPP</td>
<td>E2020</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Striatum</td>
<td>9.96 ± 4.21</td>
<td>9.61 ± 3.95</td>
<td>57.4 ± 5.26</td>
<td>23.2 ± 1.31</td>
<td>165 ± 9.3</td>
</tr>
<tr>
<td>Cortex</td>
<td>4.70 ± 1.82</td>
<td>10.3 ± 5.55</td>
<td>13.9 ± 1.50</td>
<td>4.12 ± 0.27</td>
<td>23.5 ± 3.8</td>
</tr>
<tr>
<td>Hippocampus</td>
<td>3.91 ± 1.51</td>
<td>12.2 ± 7.03</td>
<td>20.3 ± 0.88</td>
<td>5.52 ± 0.23</td>
<td>46.5 ± 4.3</td>
</tr>
<tr>
<td>Hypothalamus</td>
<td>2.16 ± 0.71</td>
<td>1.85 ± 0.64</td>
<td>9.14 ± 0.86</td>
<td>4.67 ± 0.86</td>
<td>n.a</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>1</td>
<td>1</td>
<td>1.09 ± 0.11</td>
<td>n.a</td>
<td>4.8 ± 0.8</td>
</tr>
</tbody>
</table>

*Data from Fuhrmann, 1985.
**Data from Vickroy, 1984.

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**Table II. Effect of pretreatments**

<table>
<thead>
<tr>
<th>%dose/brain</th>
<th>Control (n = 5–10/interval)</th>
<th>Haloperidol (n = 3/interval)</th>
<th>((+))PPP (n = 3/interval)</th>
<th>E2020 (n = 3/interval)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Striatum</td>
<td>2.11 ± 0.28</td>
<td>2.16 ± 0.14</td>
<td>2.19 ± 0.20</td>
<td>2.17 ± 0.20</td>
</tr>
<tr>
<td>Cortex</td>
<td>2.01 ± 0.22</td>
<td>2.05 ± 0.12</td>
<td>2.08 ± 0.19</td>
<td>2.06 ± 0.19</td>
</tr>
<tr>
<td>Hippocampus</td>
<td>2.00 ± 0.19</td>
<td>2.01 ± 0.18</td>
<td>2.03 ± 0.19</td>
<td>2.02 ± 0.19</td>
</tr>
<tr>
<td>Cerebellum</td>
<td>2.11 ± 0.28</td>
<td>2.16 ± 0.14</td>
<td>2.19 ± 0.20</td>
<td>2.17 ± 0.20</td>
</tr>
<tr>
<td>Blood</td>
<td>2.11 ± 0.28</td>
<td>2.16 ± 0.14</td>
<td>2.19 ± 0.20</td>
<td>2.17 ± 0.20</td>
</tr>
</tbody>
</table>

*\(P < 0.05\), **\(P < 0.005\), ***\(P < 0.0005\), different from controls by two-tailed t-test. Percent injected dose per gram of tissue, normalized to a 25 g mouse.
brain uptakes of similar magnitude at 5 min, but at later times the (+)-isomer brain distribution stayed uniform and comparable to the cerebellar level of (-)-[18F]FEOBV, whereas the “cholinergic” pattern of brain localization was specific to the (-)-isomer of [18F]FEOBV. The concentrations of (-)- and (+)-[18F]FEOBV enantiomers at 180 min in selected tissues of the mouse are shown in Figure 4.

**Rat biodistribution, dosimetry, and metabolism**

As a prerequisite to using [18F]FEOBV in human studies, rat whole body biodistribution studies (Table III) were carried out to evaluate tracer disposition and calculate expected radiation doses to man. [18F]FEOBV radioactivity was eliminated from the body mainly through gastrointestinal tract. Nearly 50% of the injected dose was found in the small bowel and contents at 120 min. The estimated human internal radiation dosimetry calculated from rat biodistribution using the MIRD formalism is shown in Table IV. The upper large intestine was estimated to receive the largest dose, 0.67 rad/mCi. This calculation allows up to a 10 mCi dose of [18F]FEOBV to be administered to humans.

Modest bone uptake of radioactivity was seen with [18F]FEOBV in rodents, which accumulated at a slow rate following injection. Bone uptake was most likely the result of metabolic defluorination of the tracer and deposition of inorganic [18F]fluoride ion. The inactive (+)- isomer of [18F]FEOBV showed a similar level of bone uptake as the active (-) form (Fig. 4).

Rodent brain tissue was analyzed for possible metabolites of [18F]FEOBV at two times following injection. The only radioactive species extractable from either mouse or rat brain 30 and 90 min after administration was unmetabolized FEOBV. The recovery of radioactivity from brain homogenates by the initial buffer/ethanol extraction process was 90–97%, but large amounts of lipid were present in the initial extract which interfered with thin layer chromatography (TLC). The fatty substances were precipitated by addition of CH3CN and the final extract was then analyzed by TLC without difficulty. This defatting step reduced total recovery to 60–70%, but no selective precipitation of FEOBV metabolites occurred with the fats. HPLC analysis of the fat precipitate resolubilized in CH2Cl2 found only authentic [18F]FEOBV present.

Radioactive metabolites of FEOBV appeared quickly in rodent blood, were highly polar in nature, and easily distinguishable from parent agent on TLC. They partitioned strongly with the blood cell solids and resisted extraction by the saline/ethanol combination, but the methanol/trifluoroacetic acid mixture solubilized them. When this extract was made basic with triethylamine or KOH and then analyzed by normal phase TLC, all radioactivity remained at the origin. Passage of the basified extract through a short alumina column and rinsing with methanol resulted in little trapping of radioactivity on the column. This suggested that the polar metabolite(s) was not inorganic [18F]fluoride ion. Based on analyses from two rats, the percentage of unmetabolized FEOBV in whole blood was ~80% at 5 min, 65–70% at 10 min, ~50% at 30 min, and less than 10% at 60 min. If the FEOBV blood metabolite profile in rats holds true for primates, it may be possible to do accurate blood activity metabolite corrections of the input function in FEOBV-PET studies without the need for chromatography, using a simple centrifugation of whole blood diluted with a saline alcohol extractant. In this case, the percentage of total blood activity in the decanted supernatant would be equal to the amount of unmetabolized FEOBV in the blood.

**Toxicity**

Unlabeled (-)-FEOBV was a strong neuromuscular blocking agent in rodents. Mice given parenteral doses quickly exhibited acute curare-like symptoms characteristic of vesamicol intoxication (Rogers et al., 1989). Intravenous doses as low as 30 µg/kg (-)-FEOBV were lethal within 30 min. By contrast, (+)-FEOBV did not kill mice at doses up to 4,000 µg/kg. Animals ventilated mechanically survived at least five times the mean lethal dose of (-)-FEOBV, suggesting that the primary lethal event was respiratory paralysis. Survivors recovered to apparent normalcy after 1–2 h, indicating (-)-FEOBV was eliminated or otherwise detoxified at a fairly rapid rate in rodents. No obvious behavioral or neurological signs of longer term neurotoxicity were seen in a small number of mice that were observed for 2 days following acute sublethal doses of (-)-FEOBV.

**DISCUSSION**

This work is a continuation of our earlier efforts to develop radiiodinated and carbon-11 labeled benzovesamicols [123I]IBVM and [11C]MABV as VACHT imaging agents for studying cholinergic function in Alzheimer’s disease (Jung et al., 1990; Kilbourn et al., 1990; Mulholland et al., 1991a; Mulholland and Jung, 1992). These agents have shown regional localization in vivo that closely matches the distributions of cholinergic markers such as ChAT, high affinity choline uptake, and acetylcholinesterase. The basic premise underlying the applicability of benzovesamicol radiotracers in studies of AD is that VACHT levels change in parallel fashion and magnitude with other cholinergic marker proteins, in particular ChAT, over the course of the disease. The validity of this assumption has been debated, based on results of some early studies showing mismatches between [3H]vesamicol binding and ChAT in postmortem AD brains (Kish et al., 1990; Ruberg et al., 1990), but more recent studies using higher affinity VACHT ligands (Efange et al., 1995b) or immunocytochemical probes (Gilmor et al., 1997) suggest VACHT and ChAT may in fact track closely together in AD and cholinergic...
Fig. 3. Autoradiographic distribution of \(^{18}\)FEOBV in rat brain at 5 min (A) and 180 min (B) following bolus i.v. tracer injection. Each time panel shows coronal sections at comparable rostrocaudal levels from the head of the caudate (upper left) to the cerebellum and brain stem (lower right). Tracer concentration is indicated by pseudocolor transformation, according to the scale at the lower right of each panel, with red and violet representing the highest and lowest levels, respectively. Note the laminar cortical and selective subcortical retention is apparent even at 5 min. At 180 min, strong retention is present in caudate (upper left section), basolateral amygdala (upper right), interpeduncular nucleus (lower left), and hypoglossal nucleus (lower right).
lesion models. It can be argued that vesamicol itself is not a particularly good ligand for VAChT, since it also binds other receptors (see below), and its affinity for VAChT, \( K_d = 11-36 \text{ nM}_{\text{rat brain}} \) (Marien et al., 1987); 1 nM\(_{\text{torpedo organ}}\) (Rogers et al., 1993a) is much weaker than analogs such as aminobenzovesamicol, ABV \( K_d = 6.5 \text{ pM} \) (Rogers et al., 1993a). IBVM, MABV, and FEOBV are very close analogs of ABV and have similar affinities as ABV for the VAChT, as indicated by the similar IC\(_{50}\) values for inhibition of \(^3\text{H}\)-MABV binding to rodent cortical membranes (Jung et al., 1996; Mulholland, unpublished).

Although it is very similar to \(^{123}\)I-BVM and \(^{11}\text{C}\)MABV in VAChT binding properties in vivo, \(^{18}\text{F}\) FEOBV may possess advantages over the other two agents as an imaging tracer for AD. When used with PET, \(^{18}\text{F}\) FEOBV has intrinsic quantification and resolution advantages over \(^{123}\)I-BVM/SPECT, which could result in improved sensitivity and precision in measuring cortical and hippocampal VAChT losses in

**Fig. 4.** Comparison of tissue uptakes of (−) and (+) isomers of \(^{18}\text{F}\) FEOBV in the mouse at 180 min. Values represent mean ± SD of at least three animals.

**TABLE III.** Whole body biodistribution of FEOBV in rats

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Brain</th>
<th>Eyeballs</th>
<th>Heart</th>
<th>Lung</th>
<th>Liver</th>
<th>Pancreas</th>
<th>Spleen</th>
<th>Adrenal</th>
<th>Kidney</th>
<th>Stomach</th>
<th>Stomach contents</th>
<th>Small intestine</th>
<th>Small intestine contents</th>
<th>Caecum</th>
<th>Caecum contents</th>
<th>Large intestine</th>
<th>Large intestine contents</th>
<th>Ovary</th>
<th>Muscle*</th>
<th>Bone*</th>
<th>Urine</th>
<th>Blood*</th>
<th>Testes</th>
<th>Rest of carcass</th>
<th>Total recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 min</td>
<td>1.47 ± 0.11</td>
<td>0.05 ± 0.01</td>
<td>0.77 ± 0.09</td>
<td>3.19 ± 0.45</td>
<td>11.58 ± 0.89</td>
<td>1.16 ± 0.23</td>
<td>0.60 ± 0.18</td>
<td>0.18 ± 0.06</td>
<td>3.39 ± 0.40</td>
<td>1.11 ± 0.13</td>
<td>0.28 ± 0.12</td>
<td>5.36 ± 1.53</td>
<td>1.80 ± 0.69</td>
<td>0.51 ± 0.10</td>
<td>0.54 ± 0.21</td>
<td>0.98 ± 0.26</td>
<td>0.28 ± 0.09</td>
<td>0.28 ± 0.01</td>
<td>0.32 ± 0.14</td>
<td>0.32 ± 0.01</td>
<td>0.07 ± 0.01</td>
<td>0.22 ± 0.03</td>
<td>0.71 ± 0.03</td>
<td>68.2 ± 2.2</td>
<td>101 ± 2.5</td>
</tr>
<tr>
<td>15 min</td>
<td>1.58 ± 0.20</td>
<td>0.05 ± 0.00</td>
<td>0.64 ± 0.07</td>
<td>2.12 ± 0.47</td>
<td>6.93 ± 0.64</td>
<td>2.01 ± 0.16</td>
<td>0.73 ± 0.11</td>
<td>0.06 ± 0.05</td>
<td>2.25 ± 0.04</td>
<td>1.34 ± 0.12</td>
<td>1.60 ± 1.00</td>
<td>6.50 ± 1.44</td>
<td>1.74 ± 0.61</td>
<td>0.45 ± 0.08</td>
<td>0.54 ± 0.07</td>
<td>0.89 ± 0.16</td>
<td>0.18 ± 0.10</td>
<td>0.14 ± 0.01</td>
<td>0.30 ± 0.04</td>
<td>0.42 ± 0.06</td>
<td>0.23 ± 0.05</td>
<td>0.22 ± 0.03</td>
<td>0.75 ± 0.02</td>
<td>67.7 ± 1.8</td>
<td>98 ± 1.9</td>
</tr>
<tr>
<td>30 min</td>
<td>1.15 ± 0.12</td>
<td>0.06 ± 0.00</td>
<td>0.46 ± 0.02</td>
<td>1.25 ± 0.10</td>
<td>6.38 ± 0.81</td>
<td>1.42 ± 0.40</td>
<td>0.53 ± 0.06</td>
<td>0.06 ± 0.02</td>
<td>1.53 ± 0.15</td>
<td>1.23 ± 0.26</td>
<td>3.43 ± 2.31</td>
<td>6.72 ± 2.62</td>
<td>2.26 ± 0.81</td>
<td>0.42 ± 0.10</td>
<td>1.21 ± 0.29</td>
<td>0.81 ± 0.11</td>
<td>0.65 ± 0.29</td>
<td>0.11 ± 0.01</td>
<td>0.27 ± 0.05</td>
<td>0.30 ± 0.01</td>
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<td>1.05 ± 0.36</td>
<td>0.026 ± 0.03</td>
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<td>68.9 ± 3.6</td>
</tr>
<tr>
<td>120 min</td>
<td>0.87 ± 0.13</td>
<td>0.09 ± 0.01</td>
<td>0.30 ± 0.03</td>
<td>0.53 ± 0.14</td>
<td>2.36 ± 0.32</td>
<td>0.40 ± 0.06</td>
<td>0.18 ± 0.04</td>
<td>0.03 ± 0.01</td>
<td>0.76 ± 0.08</td>
<td>0.54 ± 0.11</td>
<td>0.93 ± 0.01</td>
<td>5.81 ± 1.84</td>
<td>42.62 ± 3.37</td>
<td>0.48 ± 0.05</td>
<td>2.15 ± 1.15</td>
<td>0.70 ± 0.12</td>
<td>0.54 ± 0.24</td>
<td>0.05 ± 0.01</td>
<td>0.18 ± 0.02</td>
<td>0.33 ± 0.06</td>
<td>2.72 ± 0.12</td>
<td>0.28 ± 0.02</td>
<td>0.82 ± 0.12</td>
<td>37.67 ± 3.1</td>
<td>95 ± 4.0</td>
</tr>
</tbody>
</table>

Values are the mean percent injected dose per organ ± SD of at least four animals at each time point, except where noted otherwise. *Values expressed as percent injected dose per gram.
AD (Kuhl et al., 1996). While [12C]MABV is also a PET agent, [18F]FEOBV benefits from a longer halflife, which enables late imaging at a point where near steady-state binding levels are reached. Also, because of differences in labeling chemistries between 18F and 11C, [18F]FEOBV can be obtained routinely at much higher specific activity and lower dose masses than [12C]MABV, thus improving the safety factor of [18F]FEOBV in imaging studies.

A number of 18F-labeled benzovesamicols and vesamicols have been evaluated in the past as cholinergic neuron imaging agents (Fig. 5). Our earlier work with fluorobenzylamino-benzovesamicol (FBnABV) showed the compound to have favorable cholinergic regional brain distribution patterns in rodents and primates following i.v. injection, but it suffered from relatively low brain extraction, possibly due to the compound’s high lipophilicity (Mulholland et al., 1991a,b). Two [18F]fluoroacetyl derivatives of aminobenzovesamicol called NEFA and FAA also have been reported (Ingvar et al., 1993; Rogers et al., 1993b). FAA underwent rapid metabolic loss of the label, but the more stable analog NEFA showed a cholinergic pattern of regional brain localization, and its binding was stereoselective to the (+) isomer (Ingvar et al., 1993). In addition, two fluorinated vesamicols have been studied. [18F]Fluoromethylvesamicol had very high brain extraction but did not distribute in a pattern consistent with cholinergic innervation (Rogers et al., 1993b). Another agent called [18F]fluorobenzyltrozamicol (FBT) showed good brain uptake and moderate cholinergic localization, with a striatum/cerebellum ratio ~2 at 3 h in rodents as well as primates (Efange et al., 1994; Mach et al., 1997).

Several vesamicol tracers have been reported to interact with additional binding sites and neurotransmitter systems besides the VACHT. Vesamicol itself binds to σ receptors and a “vesamicol-binding-protein,” independent of the VACHT (Hicks et al., 1991). Possibly because of these secondary interactions, biodistribution studies using tritiated (+)-vesamicol showed poor in vivo localization in cholinergic brain areas (Jung et al., 1990). Some vesamicol analogs (Efange et al., 1995a) and (+)-benzovesamicol isomers (Jung et al., 1996) also show high affinity for σ receptors. Furthermore, dramatic increases in striatal tracer binding, up to 200% in some cases, have been reported with certain analogs following acute treatment with dopamine receptor antagonists (Ingvar et al., 1993; Efange et al., 1994, 1995a, 1996). These other interactions could complicate the interpretation of tracer data intended to assess cholinergic presynaptic function.

We were unable to find evidence of strong binding interactions in vivo of (−)-[18F]FEOBV with noncholinergic receptors. Pharmacological pretreatment of mice with large doses of the σ ligand (+)-3-PPP, or E-2020, a potent acetylcholinesterase inhibitor which also binds σ receptors, did not significantly alter the brain uptake or regional localization of (−)-[18F]FEOBV (Table II). Reductions in (−)-[18F]FEOBV total brain uptake and blood levels occurred following haloperidol, a potent antagonist of both D2 and σ receptors, but it is likely that blood flow changes caused by the near-stupefying haloperidol doses given were responsible for the alterations. In the 240-min haloperidol group, but not in the earlier treatment groups, we did observe a striatal %dose/g increase relative to controls that was significant, and a nonsignificant trend (P > 0.05) toward increased %dose/g levels for cortex, hippocampus and cerebellum. However, the relative brain (−)-[18F]FEOBV concentrations of the haloperidol groups, expressed as an average of tissue-to-cerebellum ratios from each animal, did not differ from controls (TableII). This suggests, further, that haloperidol pretreatment altered hemodynamic factors controlling tracer delivery to the brain, and did not modulate receptor availability at the cellular level.

Our results argue that FEOBV is not interacting competitively at sigma or D2-like receptor sites. In vitro, we found haloperidol to be a very weak inhibitor (IC50 > 1 μM) of benzovesamicol [3H]MABV binding to mouse striatal and cortical membranes (Mulholland, unpublished findings). Furthermore, the equivocal effects of haloperidol on striatal (−)-[18F]FEOBV binding we observed in mice indicate that acute D2 receptor antagonism did not modulate (increase) (−)-[18F]FEOBV binding to the VACHT through a mechanism involving blockade of normal dopamine inhibitory input onto cholinergic interneurons in the striatum. There is evidence for an inhibitory feedback circuit in the striatum and substantia nigra involving dopamine D2, acetylcholine, and GABA neurons (Lehmann and Langer, 1983), and several investigators have suggested this interneuronal mechanism to explain rapid, large increases they have observed in vivo striatal binding of certain vesamicol tracers following acute D2 antagonist pretreatments (Efange et al., 1994; Ingvar et al., 1993). We used haloperidol pretreat-
ment doses 10–30 times higher than those investigators employed, but saw little effect on ($^2$-$^{[18F]}$FEOBV striatal localization. It should be pointed out we used different species and intervals between haloperidol pretreatment and tracer injection in our study. Further investigation may be necessary to resolve whether our differing results are due to altered experimental conditions or basic differences in tracer properties between FEOBV and other vesamicol agents.

($^2$-$^{[18F]}$FEOBV was excreted primarily through the gastrointestinal tract. This excretion pattern results in the large bowel being the body tissue which receives the highest radioactive burden from intravenously administered ($^2$-$^{[18F]}$FEOBV, and limits the allowable dose to 10 mCi/year in adult humans, based on estimated dosimetry calculations from rat data (Table IV). Gastrointestinal excretion is also the major disposition pathway of [$^{123I}$]IBVM in rats and humans (Kuhl et al., 1994).

The absence of labeled brain metabolites of FEOBV in rodents is a desirable feature of this agent. Assuming this holds true in primates, it greatly simplifies the tasks of estimating tracer brain transport and VACHT binding site densities from PET studies with ($^2$-$^{[18F]}$FEOBV, since the total brain signal is contributed by a single radiochemical entity. Furthermore, the simple blood metabolism of ($^2$-$^{[18F]}$FEOBV makes it easy to do metabolite corrections of the plasma input function for quantitative imaging studies. The resistance of FEOBV to metabolic defluorination and bone deposition is also noteworthy. In general, [$^{18F}$]fluoroalkyl and [$^{18F}$]fluoroalkylamino groups tend to be more prone to loss of inorganic $^{18F}$– than are [$^{18F}$]fluoroaromatic groups. However FEOBV has a [$^{18F}$]fluoroethoxy group, and the presence of a vicinal oxygen atom in this group appears to stabilize the $^{18F}$-label against defluorination similar to the level of a [$^{18F}$]fluoroaromatic moiety. Relatively few [$^{18F}$]fluoroethyl ethers have been examined; however, in one case it was found this group conferred the greatest metabolic stability in a series of $^{18F}$-labeled estrogens (French et al., 1993).

In vivo blocking studies using unlabeled vesamicol analogs to prevent or displace ($^2$-$^{[18F]}$FEOBV uptake were not feasible because of their high toxicity. Previous in vivo ($^{123I}$]IBVM blocking attempts using vesamicol as the blocking treatment in rodents resulted in death at a vesamicol dose that blocked about 50% of control striatal IBVM uptake (Jung et al., 1990). However, in vitro ligand binding studies from this laboratory (Mulholland, unpublished findings) have found 5-substituted benzovesamicols to be very specific and saturable in their binding to striatal, cortical, or hippocampal membranes, and this binding was inhibited by nanomolar to subnanomolar concentrations of structurally related benzovesamicols. For example, ($^2$-$^{[18F]}$FEOBV inhibited specific binding of the probe ($^2$-$^5$-$^{[3H]}$methylaminobenzovesamicol ([3H]MABV, 80 Ci/mmol, 0.5 nM incubation concentration) to mouse cortical membranes with an apparent $IC_{50}$ of 0.4 nM.

Biological stereospecificity is characteristic of other benzovesamicols and vesamicol itself. Both the toxicity and the cholinergic specificity were attributable almost exclusively to the ($^2$-(2R,3R)-isomer of FEOBV. Mice survived comparatively large doses of the ($^1$)-isomer, 4,000 µg/kg i.v., that were 133 times greater than the mean lethal dose of the ($^2$)-isomer. The rodent biodistributions of the two enantiomers were very different. ($^1$-$^{[18F]}$FEOBV activity was uniform throughout the brain, and levels in all regions except cerebellum were lower than uptake of the ($^2$)-isomer. ($^2$-$^{[18F]}$FEOBV did accumulate in peripheral tissues rich in cholinergic terminals, such as atria, stomach, and pancreas, in which ($^2$-$^{[18F]}$FEOBV did localize (Fig. 4). Another indication of the biological stereospecificity of FEOBV was that the $IC_{50}$ of ($^1$)-FEOBV for inhibition of in vitro ($^2$-$^{[3H]}$MABV binding to cortical membranes was
nearly three orders of magnitude higher than that of the (−)-isomer. The clear relation between toxicity and cholinergic localization in FEOBV enantiomers suggests strongly that vesicular ACh uptake inhibition is the direct cause of toxicity seen with (−)-FEOBV.

Toxic levels of vesamicol analogs cause respiratory paralysis, probably through nondepolarizing blockade of peripheral cholinergic neurotransmission. Nearly all of the benzovesamicols we studied with high cholinergic localization, including IBVM, MABV, and other analogs (Mulholland et al., 1991b), share similar levels of toxicity. Since binding and inhibition of VACHT at the picomolar level (Rogers et al., 1993a) appears to be a common mechanism underlying both the high cholinergic localization and toxicity of these agents, it is unlikely these two properties can be separated in the benzovesamicol class of cholinergic tracers.

High toxicity does not necessarily preclude usefulness as an imaging agent. Other very toxic compounds, including thallium-201, carfentanil, (Dannals et al., 1985), IBVM and epibatidine analogs (Ding et al., 1996; Horti et al., 1997; Musachio et al., 1997; Villemagne et al., 1997), have been used safely and/or are being developed currently as human PET and SPECT imaging agents. The very high specific activities attainable routinely in fluorine-18-labeled agents, and the sensitive HPLC techniques used universally in tracer purification and analysis, make production of safe imaging doses of (−)-[^18F]FEOBV a reasonably straightforward process. For example, a 10 mCi dose of (−)-[^18F]FEOBV with specific activity of 5 Ci/µmol, an easily achievable value in our experience, contains only 2 nmol or 0.75 µg of (−)-FEOBV drug. This low mass quantity is less than 1/1,000 of the estimated human toxic dose based on mouse data, 1–2 mg (2.7–5.42 µmol, which incidentally is more than the total amount of labeling precursor we use per synthesis of (−)-[^18F]FEOBV), and provides a large margin of safety for human imaging studies. Nevertheless, it is important to recognize the potential risks of FEOBV and the vesamicol class in general, and practice care in the production and quality control standards for these tracers.

In conclusion, this work in rodents demonstrates (−)-[^18F]FEOBV to be a promising radiotracer for mapping cholinergic synapses in vivo. It has high brain extraction, favorable metabolism, a strong cholinergic pattern of localization which correlates highly with brain choline acetyltransferase distribution, and a lack of demonstrable extra-cholinergic binding interactions. Both the cholinergic tracer behavior and the pharmacological effects are stereospecific to the (−)-enantiomer.

Further work is needed to evaluate regional brain kinetics of (−)-[^18F]FEOBV in primates and develop imaging and analysis protocols to extract information from FEOBV-PET scans. It will then be possible to determine the utility of this new agent for measuring cholinergic synapse density and function in man using PET.