The aim of this study was to assess human striatal dopamine receptor 2 (D2) and cortical 5-hydroxytryptamine receptor 2A (5-HT2A) occupancy of SB-773812 to demonstrate brain penetration and binding to the target receptors and assess the pharmacokinetics-receptor occupancy relationship over time to aid dose selection and dosage regimen, in preparation for the phase II trials. Methods: D2 and 5-HT2A occupancy were measured over time (both at the time of maximum [Tmax; 6 ± 2 h] and at the time of minimum [Trough; 24 ± 4 h] plasma concentration after dosing) by means of 123I-iodobenzamide and 123I-4-amino-N-[1-[3-(4-fluorophenoxy)propyl]-4-methyl-4-piperidinyl]5-iodo-2-methoxybenzamide (123I-R91150) SPECT in 3 studies. Study A consisted of SB-773812 single doses in healthy volunteers—D2 occupancy measured at 48 (n = 9) and 56 mg (n = 9) and 5-HT2A occupancy at 56 mg (n = 9); study B consisted of D2 and 5-HT2A occupancy measured in 12 stabilized schizophrenia patients on stable doses (16–18 d of 56 mg/d) after washout of previous medication; and study C included D2 occupancy measured in a double-blind study of patients with acutely exacerbated schizophrenia (n = 10) on stable doses (18–21 d) of SB-773812 (100 mg/d; n = 7) or risperidone (6 mg/d; n = 3).

Results: Study A showed less than 30% D2 occupancy at Tmax, maintained at Trough, 5-HT2A occupancy was 74%–97% and also maintained over time. Study B revealed that 8 of the 12 schizophrenia patients showed more than 40% D2 occupancy. 5-HT2A occupancy ranged from 91% to 100%. In study C, SB-773812–induced D2 occupancy was 60.3% ± 13.3% at Tmax and 55.1% ± 4.9% at Trough. The pharmacokinetics-receptor occupancy relationship was assessed in each study and strengthened, combining all data to yield a concentration associated with 50% occupancy (EC50) of 92.7 ± 13.5 ng/mL for D2 and 2.11 ± 0.50 ng/mL for 5-HT2A. Conclusion: In all subjects, SB-773812 showed penetration into the brain, reaching its target receptors. In patients with schizophrenia, D2 occupancy levels induced by a single dose were maintained over time, indicating that once-daily dosing regimens are appropriate. Pharmacokinetics-receptor occupancy analysis provided guidance for the selection of a clinically effective dose, supporting progression in phase II.

Key Words: antipsychotic; schizophrenia; single photon emission computed tomography; receptor occupancy; SB-773812

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The antipsychotics currently available for the treatment of schizophrenia predominantly are efficacious in ameliorating positive (psychotic) symptoms but less effective in treating negative symptoms and cognition. Additionally, these treatments may cause side effects (e.g., neurologic and metabolic), which often lead to noncompliance within the schizophrenia population (1). Thus, there is a high unmet need in this population, especially for treatments that may have increased efficacy against negative symptoms and cognition while reducing the side effect burden. SB-773812 has been specifically designed to target antagonism at those receptors believed to be associated with antipsychotic efficacy but to eliminate affinity at receptors suggested to be linked to the side effects of current antipsychotics. SB-773812 is a moderate-affinity antagonist at dopamine receptor 2 (D2) (–log expression of Ki, the inhibitor constant [pKi], 7.4) and a high-affinity antagonist at the dopamine receptor 3 (D3) (pKi, 8.5) and at the serotonin 5-hydroxytryptamine receptors 2A (5-HT2A) (pKi, 9.0), 2C (pKi, 8.1), and 6 (pKi, 8.1). It has no affinity for histamine receptor 1, muscarinic receptors 1–4, dopamine receptor 1, adrenergic receptor 1B, or adrenergic receptors 1–3. This unique receptor interaction profile is predicted to deliver efficacy against positive symptoms, with the potential for efficacy against the cognitive and negative or mood symptoms of the disease while minimizing side effects associated with other antipsychotics, including sedation, weight gain, cognitive impairment, dysphoria, and extrapyramidal symp-

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**REFERENCES:**


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toms. SB-773812 is characterized by a linear pharmacokinetics profile, oral clearance of approximately one third of the liver blood flow, and a large apparent volume of distribution conductive to an average terminal half-life of approximately 55 h. As such, steady state is generally achieved within 2 wk of dosing, and the steady-state pharmacokinetics profile is characterized by limited peak-to-trough fluctuations. The CYP450 isoform 3A4 appears to be a major metabolic pathway for SB-773812 (GlaxoSmithKline, unpublished data, April 2009).

Neurotransmission imaging techniques, such as PET and SPECT, play a significant role in the development process of drugs (2)—particularly relevant for drugs acting at the central nervous system, for which animal models are limited and translation to humans is extremely complicated. There are examples in the literature on how these technologies can provide evidence of drug–brain penetration and of drug action at the target and can contribute to dose selection based on the relationship between drug plasma concentrations (pharmacokinetics) and receptor occupancy in the brain (3). However, only few of these works have used SPECT. Moreover, published evidence of the contribution of these technologies to the whole development process of a particular single drug is still rare.

Independent of the imaging modality, it is important that the PET or SPECT ligand to be used in drug development trials is well characterized in humans to allow accurate interpretation of the data. SPECT ligands for 2 of the main SB-773812 targets, D2 ([123I-iodobenzamide] and 5-HT2A ([123I-iodobenzamide and 123I-4-amino-N-[1-[3-(4-fluoro-phenoxy)propyl]-4-methyl-4-piperidinyl]5-iodo-2-methoxy-benzamide [123I-R91150]), were available at the time clinical development of this drug was being planned, and several benchmark studies were performed before SB-773812 was selected as a candidate for first-time-in-humans administration. First, [123I-R91150 was fully characterized in healthy volunteers (4) and shown to be a selective ligand for 5-HT2A measurements, demonstrating a dose-dependent displacement after ketanserin challenge (5). Second, the test–retest variability of a single bolus injection of [123I-iodobenzamide was reported for the first time (6). Finally, the D2 occupancy induced by marketed atypical antipsychotics was measured over time in stabilized patients to estimate the expected levels of D2 occupancy to achieve efficacy, using exactly the same methodology to be applied in the subsequent planned SPECT studies with SB-773812 (7).

The present work aimed to contribute to the development of the antipsychotic SB-773812 by investigating human brain penetration and confirming the target of action in early development as well as pharmacokinetics–receptor occupancy relationships in both healthy volunteers and patients with schizophrenia by means of [123I-iodobenzamide and [123I-R91150 SPECT.

MATERIALS AND METHODS

Three studies were performed on a group of healthy volunteers and a group of patients with schizophrenia to measure SB-773812–induced D2 occupancy, 5-HT2A occupancy, and their relationships with plasma concentration (Table 1). First, SB-773812–induced D2 and 5-HT2A occupancy was measured over time in healthy volunteers at the 2 highest doses (48 and 56 mg), previously shown to be safe and well tolerated in the first-time-in-humans single-dose escalation study (study A). Second, D2 and 5-HT2A occupancy was measured at a single time point at stable SB-773812 plasma levels after repeated administration of the 56-mg dose in patients with stable schizophrenia (study B). Finally, the D2 occupancy at stable SB-773812 plasma levels after repeated administration of 100 mg/d was measured over time in patients with acute schizophrenia (study C). All the studies were approved by the local Ethics Committees and the Spanish Ministry of Health. All subjects provided written informed consent before being included in the study.

Study Subjects

All subjects had normal laboratory evaluations, including hematology; clinical chemistry; urinalysis; serology for HIV, hepatitis B, and hepatitis C; 12-lead electrocardiography; and 24-h Holter monitoring. All subjects had negative pre-study alcohol breath and urine drug screening for amphetamines, barbiturates, cocaine, opiates, cannabinoids, and benzodiazepines.

Twenty-seven healthy male volunteers (mean age ± SD, 26 ± 5 y), with no clinically significant abnormality identified on the medical and laboratory evaluations as specified, were included in the study. From the total of 26 patients with schizophrenia (24 men), 14 (mean age ± SD, 33 ± 7 y) were diagnosed with chronic schizophrenia by DSM-IV criteria (8) and were included in study B. The remaining 12 patients (mean age ± SD, 36 ± 8 y) were diagnosed with schizophrenia or schizophrainiform disorder by DSM-IV criteria, showing moderate disease exacerbation with a positive and negative syndrome scale (PANSS) score of more than 70 and a score on the PANSS-positive scale of 25 or more, and were included in study C (Table 1).

Study Designs

Radiation exposure of the study subjects was kept below 10 mSv per the guidelines of the International Commission on Radiological Protection (ICRP) (category IIb in ICRP publication 62 (9)). For this reason, study designs were adapted so that a maximum of 2 SPECT scans were obtained for each subject, at least 48 h apart. A summary of the study designs is provided in Table 1.

Study A. The whole sample of healthy volunteers (n = 27) was divided into 3 cohorts of 9 subjects each. Single SB-773812 doses were administered: 48 mg to subjects in cohort 1 and 56 mg to subjects in cohorts 2 and 3. To allow receptor occupancy estimates, each subject underwent 2 scans, one at baseline and another after drug administration. For each cohort, 5 subjects were scanned at 6 ± 2 h after drug administration (time of maximal plasma concentration, or Tmax), and 4 subjects were scanned at 24 ± 4 h after drug administration (time of minimal plasma concentration, or Trough). In cohorts 1 and 2, D2 occupancy was measured, and in cohort 3, 5-HT2A occupancy was measured. The selection of the time frame for scanning the subjects around Tmax was based on the pharmacokinetic characteristics of SB-773812 in previous studies on healthy volunteers and patients with schizophrenia (GlaxoSmithKline, unpublished data, April 2009), showing that SB-773812 peak plasma concentration appears to be independent of the dose or duration of dosing and is typically observed at 4–8 h after dosing. The ±2-h window allowed for some flexibility in the scanning execution and was deemed appropriate given the range of expected Tmax for SB-773812.
HT2A occupancy assessments were then obtained for each subject at T max and T trough.

After scans were completed, patients underwent a 1-wk washout and restabilization hospitalized period and a follow-up visit. 6 months after the last SB-773812 dose.

In Study B, patients were included in the study and a washout of previous antipsychotic medication, they were hospitalized and received 56 mg of SB-773812 daily for 16–18 d to achieve steady-state plasma levels. SPECT scans for D2 and 5-HT2A occupancy assessments were then obtained for each subject at a single time point, 12 h after administration of the last dose. After scans were completed, patients underwent a 1-wk washout and restabilization hospitalized period and a follow-up visit at T max.

In Study C, patients with acute schizophrenia were hospitalized and randomized to receive either SB-773812 (100 mg/d; n = 9) or risperidone (6 mg/d; n = 3) until steady-state plasma concentration were achieved and SPECT scans acquired (18–21 d). The risperidone arm was included only to keep the investigators masked. A 1-wk washout and restabilization hospitalized period followed, and at the end of the last SB-773812 dose, D2 occupancy was calculated for each subject at T max and T trough.

### Plasma Sampling

SB-773812 plasma concentrations were measured in all studies at different time points. On the day of the SPECT scan, additional samples were taken immediately before and after the SPECT scan. In study C, risperidone and its metabolite 9-hydroxyrisperidone plasma concentrations were also measured at the same time points as those for SB-773812. Plasma samples for SB-773812 were analyzed by Worldwide Bioanalysis, using validated analytic methods based on protein precipitation with acetonitrile, followed by high-performance liquid chromatography mass spectrometry or mass spectrometry analysis. Drug analysis for the determination of risperidone or 9-OH-risperidone was performed using a validated assay methodology (liquid chromatography mass spectrometry or mass spectrometry) by York Bioanalytical Solutions, under the guidance of Worldwide Bioanalysis, DMPK, GlaxoSmithKline.

### SPECT Methodology

123I-iodobenzamide and 123I-R91150 SPECT scans were acquired as described elsewhere (4,10). A 3-head camera (Prism 3000S; Philips) fitted with ultra-high-resolution fanbeam collimators was used. Subject preparation included administration of potassium perchlorate (8 mg/kg) up to 20 min before ligand administration to minimize radiation exposure to the thyroid gland. Four multimodality SPECT/MRI markers (MM3003; IZI Medical Products Corp.), each filled with approximately 0.074 MBq of123I, were firmly stuck to the subject’s head on the mastoids and at the corners of the eyes and remained in the same position until both SPECT and MRI acquisitions were finalized. For D2 measurements, 60-min scan acquisitions started at 120 min after intravenous injection of123I-iodobenzamide (153.8 ± 19.2 MBq). For 5-HT2A measurements, 60-min scan acquisitions started at 180 min after intravenous injection of123I-R91150 (146.6 ± 18.6 MBq). Ligand injections were followed by a 20-mL saline serum flushing. In all the SPECT explorations, the following acquisition parameters were used: step-and-shoot mode, 360° circular orbit, 120 steps, 3°/step, and a 128 × 128 matrix.

All subjects underwent a T1-weighted 3-dimensional MRI scan on the same day as the SPECT scan using a superconductive 1.9-T system (Prestige 2T; GE Healthcare) equipped with a head coil. An axial 3-dimensional spoiled gradient-echo slab was positioned until both SPECT and MRI acquisitions were finalized.}

### Summary of Human SPECT Studies Performed to Contribute to SB-773812 Development

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Study A, healthy volunteers (n = 27; all men; 26 ± 5 y)</th>
<th>Study B, patients with chronic schizophrenia (n = 14; 10 men; 33 ± 7 y)</th>
<th>Study C, patients with exacerbated schizophrenia (n = 12; all men; 36 ± 8 y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SB-773812 dose</td>
<td>48 mg (single)</td>
<td>56 mg/d, 16–18 d (steady state)</td>
<td>100 mg/d, 18–21 d (n = 9)</td>
</tr>
<tr>
<td>Risperidone dose</td>
<td>D2</td>
<td>D2 + 5-HT2A</td>
<td>Postdose Cmax, postdose Ctrough</td>
</tr>
<tr>
<td>Target (receptor)</td>
<td>Baseline postdose</td>
<td>5-HT2A</td>
<td>Postdose Cmax, postdose Ctrough</td>
</tr>
<tr>
<td>Scans (2/subject, 48 h apart)</td>
<td>Baseline postdose</td>
<td>5-HT2A</td>
<td>Postdose Cmax, postdose Ctrough</td>
</tr>
<tr>
<td>Scan time after dose</td>
<td>T max (6 ± 2 h) (n = 5/cohort), T trough (24 ± 4 h) (n = 4/cohort)</td>
<td>12 ± 4 h</td>
<td>T max (6 ± 2 h) (n = 10), T trough (24 ± 4 h) (n = 8)</td>
</tr>
<tr>
<td>% receptor occupancy formula (×100)</td>
<td>(1 – [SURpostdose/SURbaseline])</td>
<td>(1 – [SURpostdose/SURtmax])</td>
<td>(1 – [SURpostdose/SURtmax])</td>
</tr>
<tr>
<td>Plasma concentration over time (pharmacokinetics)</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

HV = healthy volunteers.
The Chang algorithm, with a manually fitted elliptic attenuation map on each slice ($\mu = 0.1 \text{ cm}^{-1}$), was used for attenuation correction. SPECT and MRI scans were registered using fiducial markers with software implemented in-house. In short, the corresponding external markers were manually identified on the MRI and SPECT scans. Then, rigid-body transformation (3 translations and 3 rotations) was estimated automatically by minimizing the sum of the squared distances between the corresponding marker positions ($\mathbf{I}$).

Regions of interest (ROIs), including the cerebral regions with specific and nonspecific uptake, were manually drawn on each subject’s MR image and translated (copied) to the corresponding SPECT slices after coregistration. For $^{123}$I-iodobenzamide, ROIs were drawn on the whole striatum (S) and the occipital cortex (O). For $^{123}$I-R91150, ROIs were drawn on cortical regions (Ctx) and the cerebellum (C) (Fig. 1).

The specific uptake ratio (SUR) was calculated as the ratio of specific-to-nondisplaceable activity according the following equations: for $^{123}$I-iodobenzamide, $\text{SUR} = (S - O)/O$; for $^{123}$I-R91150, $\text{SUR} = (\text{Ctx} - C)/C$. The proportion of receptors occupied by the drug (RO) was calculated using the following equation:

$$\% \text{RO} = (1 - [\text{SUR}_{\text{postdose}}/\text{SUR}_{\text{baseline}}]) \times 100,$$

where $\text{SUR}_{\text{postdose}}$ is the SUR calculated after dose administration and $\text{SUR}_{\text{baseline}}$ is the SUR obtained from a sample of healthy volunteers without dose administration.

For both studies B and C, baseline values from the healthy volunteers in study A were used for receptor occupancy estimates. In all studies, plasma samples were drawn to allow a full pharmacokinetic profile.

**Pharmacokinetics–Receptor Occupancy Relationships**

The relationship between SB-773812 plasma concentration and D2 or 5-HT$_{2A}$ occupancy was assessed using a population pharmacokinetic pharmacodynamic approach as implemented in NONMEM (version VI; Globomax). The data were analyzed with the classic hyperbolic equation (derived from the law of mass action):

$$\text{RO} = \frac{100 \times C_p}{C_p + \text{EC}_{50}}.$$

Where $C_p$ is plasma concentration, EC$_{50}$ is the concentration associated with 50% occupancy (i.e., drug potency), and the maximal effect ($E_{\text{max}}$) was constrained to the theoretic value of 100% across all studies, given that smaller $E_{\text{max}}$ values were tested without improvements in model performance. Nonlinear mixed-effect modeling was used to gather an estimate of intersubject variability in drug potency and statistically test a potential time or study effect on model parameters and was deemed as the natural approach for analyzing and interpreting the sparse observations (1–3 scans per subject) obtained in the SPECT experiments.

**RESULTS**

**Study A**

Individual D$_2$ and 5-HT$_{2A}$ occupancy data over SB-773812 plasma concentrations from study A are shown in Figure 2, and Figure 3 shows representative baseline and postdrug SPECT images. After single 48- and 56-mg SB-773812 doses, healthy volunteers showed low D$_2$ occupancy ($<30\%$) both at $T_{\text{max}}$ and $T_{\text{trough}}$ (Table 2), whereas a high 5-HT$_{2A}$ occupancy was measured at $T_{\text{max}}$ ranging from 74.4% to 97.1% across the regions analyzed. This high receptor occupancy was maintained at $T_{\text{trough}}$, ranging from 72.1% to 97.8% across regions.

**Study B**

Only 12 of the 14 patients completed all the scans. Mean D$_2$ occupancy in the 12 patients with chronic schizophrenia at stable plasma levels of SB-773812 (56 mg) measured

![FIGURE 1. Selected MRI and coregistered SPECT slices at basal ganglia level (top and middle rows) and at cerebellum level (bottom row), showing representative ROIs drawn on basal ganglia and occipital cortex for D$_2$ quantification (top row) and on cortical regions and cerebellum for 5-HT$_{2A}$ quantification (middle and bottom rows). IBZM = iodobenzamide.](image1.png)

![FIGURE 2. Individual values of D$_2$ and 5-HT$_{2A}$ occupancy from study A. 5-HT$_{2A}$ occupancy values are mean of all cortical regions.](image2.png)
12 h after last dose administration was 42.7% ± 25.2% (range, 6%–81%) (Fig. 3). Eight patients showed more than 40% D₂ occupancy. Only 10 of the 12 patients undergoing the first SPECT session with ¹²³I-iodobenzamide underwent the second SPECT session with ¹²³I-R91150. Mean values of 5-HT₂A occupancy ranged from 91.4% to 100% across the cortical regions analyzed. D₂ occupancy and 5-HT₂A occupancy data from studies A and B were fitted to the corresponding Eₘₐₓ models as shown in Figures 4 and 5.

**Study C**

Only 10 of the 12 subjects had SPECT scans completed, 7 on SB-773812 (n = 5 with Tₘₐₓ and Tₜₐₕₜₜ scans completed), and 3 on risperidone (Table 1). D₂ occupancy and plasma concentration results at Tₘₐₓ and Tₜₐₕₜₜ from patients on repeated doses of SB-773812 (100 mg/d) or risperidone (6 mg/d) are presented in Table 2 and Figures 5 and 6. Data from the patients in this study were added to the previous D₂ occupancy data from studies A and B to fit the Eₘₐₓ model with all the data together. At high plasma concentration, D₂ occupancy values were lower for SB-773812 (range, 43%–83%) than for risperidone (range, 81%–88%). Furthermore, SB-773812 showed more stability over time (SB-773812 range, 33%–75%, vs. risperidone range, 60%–67%, at Tₜₐₕₜₜ) (Fig. 6).

**Pharmacokinetics–Receptor Occupancy Relationship**

Although each individual study was analyzed separately, in all cases SB-773812 plasma concentrations and receptor occupancy were fitted using the classic hyperbolic equation (i.e., an Eₘₐₓ model), and the estimated drug potency was generally consistent across the studies (Table 3; Figs. 4 and 5). R² of model fitting was 0.7 for D₂ occupancy and 0.4 for 5-HT₂A occupancy data. When all data were pooled and analyzed together, EC₅₀ calculations were precise (SEE of estimated value, 15% [D₂ occupancy] and 24% [5-HT₂A occupancy]) but associated with a substantial predicted between-subject variability of around 70% (Table 3). The nonlinear mixed-effect approach was used to test for the potential effect of subject type on D₂ drug potency and on scan time or day of dosing. None of these possible covariates resulted in a significant improvement in model fitting (P > 0.58 in all cases). Similarly, estimating Eₘₐₓ rather than fixing it to 100% did not improve significantly the D₂ fitting (P = 0.18) because the Eₘₐₓ estimate was fairly close to 100% (93% ± 14%).

**TABLE 2**

<table>
<thead>
<tr>
<th>Study</th>
<th>Drug</th>
<th>Drug dose</th>
<th>Plasma concentration during scan~Tₘₐₓ (ng/mL)</th>
<th>% D₂ occupancy at Tₘₐₓ (6 ± 2 h)</th>
<th>Plasma concentration during scan~Tₜₐₕₜₜ (ng/mL)</th>
<th>% D₂ occupancy at Tₜₐₕₜₜ (24 ± 4 h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>SB-773812</td>
<td>48/56 mg</td>
<td>21.0 ± 6.7 (n = 10)</td>
<td>16.4 ± 8.4 (n = 10)</td>
<td>17.8 ± 5.3 (n = 8)</td>
<td>21.3 ± 7.8 (n = 4)</td>
</tr>
<tr>
<td>B</td>
<td>SB-773812</td>
<td>56 mg/d</td>
<td>89.0 ± 27.1 (n = 12)</td>
<td>42.7 ± 25.2 (n = 12)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>C</td>
<td>SB-773812</td>
<td>100 mg/d</td>
<td>125.1 ± 23.7 (n = 7)</td>
<td>60.3 ± 13.3 (n = 7)</td>
<td>98.8 ± 34.5 (n = 5)</td>
<td>55.1 ± 14.9 (n = 5)</td>
</tr>
<tr>
<td></td>
<td>Risperidone</td>
<td>6 mg/d</td>
<td>55.5 ± 31.9 (n = 3)</td>
<td>83.6 ± 3.4 (n = 3)</td>
<td>26.1 ± 20.0 (n = 3)</td>
<td>64.4 ± 3.6 (n = 3)</td>
</tr>
</tbody>
</table>

Risperidone plasma concentration reported as active moiety (i.e., risperidone + 9-OH-risperidone).

NA = not available.
DISCUSSION

This paper shows how $^{123}$I-iodobenzamide and $^{123}$I-R91150 SPECT was used to contribute to the development of the antipsychotic SB-773812 from early phase I single-dose trials in healthy volunteers to phase II repeated-dose trials in patients with schizophrenia. To the best of our knowledge, this is the first time that SPECT has been used as a decision-making tool throughout the drug development process from first time in humans to proof of concept.

Despite the complexity and high cost of the PET technology, compared with SPECT, most neuroimaging studies for drug development have used PET because of its better imaging resolution, more accurate quantification, and lower variability. It is also more feasible to develop PET ligands than SPECT ligands for new target molecules. However, in those cases in which a SPECT ligand for a target is available, SPECT is a valuable alternative that provides the same information at a lower cost ($^{11}$). Nevertheless, data on antipsychotic pharmacokinetics–receptor occupancy relationships are far more limited with SPECT than with PET. Sigmoidal $E_{\text{max}}$ models using plasma concentration data are usually available in PET studies ($^{12}$–$^{17}$), whereas the most usual case in the SPECT literature is to use the antipsychotic dose as a variable for the study of pharmacokinetics–receptor occupancy relationships ($^{18}$–$^{23}$). To the best of our knowledge, only 2 SPECT studies have used plasma concentrations for pharmacokinetics–D$_2$ occupancy relationship profiles ($^{7}$,$^{18}$). Similarly, the influence of the time of scanning after last antipsychotic dose intake on D$_2$ occupancy variability has been reported using PET ($^{24}$–$^{26}$), but there is only 1 SPECT study reporting the importance of antipsychotic-induced D$_2$ occupancy assessments over time to fully characterize the pharmacokinetics–receptor occupancy profiles ($^{7}$). A carefully designed prospective within-subject comparison of the D$_2$ occupancy values measured with $^{11}$C-raclopride PET and $^{123}$I-iodobenzamide SPECT was recently reported ($^{10}$,$^{28}$). Although D$_2$ occupancy measurements with both techniques were well correlated, SPECT measurements were consistently 9%–14% lower than PET measurements. These data offer further evidence that both techniques provide the same information, but to compare D$_2$ occupancy measurements, this bias should be considered.

D$_2$ and 5-HT$_{2A}$ occupancy results after single doses of SB-773812 in healthy volunteers (study A) provided evidence of the compound brain penetrability in humans and confirmed its action at the 2 receptor targets. Moreover, a high 5-HT$_{2A}$ occupancy (>70%) was measured at a low exposure associated with a single 56-mg dose, supporting the in vitro profile of SB-773812 high affinity for this receptor. This high 5-HT$_{2A}$ occupancy was measured at both $T_{\text{max}}$ and $T_{\text{trough}}$, independently of the fact that lower
plasma concentrations were measured at $T_{\text{rough}}$, thus suggesting that a 56-mg dose was nearly saturating 5-HT$_{2A}$. Similar levels of D$_2$ occupancy were found after single doses of both 48 and 56 mg, which were below 30% in all cases. Taken altogether, these data supported a go decision at this early stage of SB-773812 development, and the preliminary estimate of drug potency for D$_2$ and 5-HT$_{2A}$ was used to design subsequent clinical exploration (including studies B and C).

A wide range of both D$_2$ occupancy and D$_2$ plasma concentrations was found in patients with schizophrenia from studies B and C (repeated administration of SB-773812, 56 and 100 mg/d, respectively), but most showed a D$_2$ occupancy between 40% and 80%. A wide range of both D$_2$ occupancy and plasma concentrations was also found in our benchmark study in stabilized patients with schizophrenia on marketed antipsychotics (risperidone, olanzapine, clozapine, and quetiapine) (7). That study also showed that at therapeutic doses of those antipsychotics, clinical response may be maintained, with SPECT-measured D$_2$ occupancy values below 65% D$_2$ occupancy (7), in agreement with data from Frankle et al. (29), showing D$_2$ occupancy levels of 55% ± 11% induced by olanzapine (10 mg/d) and of 69% ± 8% induced by risperidone (6 mg/d), measured by $^{123}$I-iodobenzamide SPECT. Evidence of 40%–80% D$_2$ occupancy in schizophrenic patients from studies B and C gave confidence about the likelihood of proving efficacy in later development phases at both 56- and 100-mg/d doses, thus minimizing risk of dose selection for phases II and III.

The estimated EC$_{50}$ for 5-HT$_{2A}$ occupancy (2.11 ± 0.50 ng/mL) was about 45-fold lower than that for D$_2$ occupancy (mean, 92.7 ± 13.5 ng/mL), confirming in vivo the much higher affinity of SB-773812 for 5-HT$_{2A}$ blockade than for D$_2$ blockade (as per in vitro design, GlaxoSmithKline, unpublished data, April 2009). Therefore, at expected therapeutic SB-773812 doses (i.e., doses delivering at least 50% D$_2$ occupancy) 5-HT$_{2A}$ was predicted to be fully blocked. Nonlinear mixed-effect models were used as the natural approach for analyzing the sparse SPECT receptor occupancy measurements (collected at different scan times in different populations and with different dosing regimens), allowing the assessment of between-subject variability in drug potency, which was found to be moderate to high for both targets (Table 3). Potential covariates were tested to explain variability, but none yielded a significant improvement in model fitting. These covariates included subject type (healthy volunteer, acute or chronic patient), chronic versus acute experiment, and timing of SPECT scan. Timing supported the fact that SB-773812 showed stable plasma concentrations and D$_2$ occupancy over time.

The high variability found for D$_2$ occupancy values is a limitation of this study. A higher variability in D$_2$ occupancy estimates has been reported using SPECT than PET (10). Moreover, the use of a healthy volunteer group for receptor occupancy estimations in patients with schizophrenia may have contributed to this variability, given the reported differences in D$_2$ density between the 2 groups (30). These differences have not been found in 5-HT$_{2A}$ availability (31), supporting the lower variability found in 5-HT$_{2A}$ occupancy estimations. Age-related differences between the healthy volunteer and patient groups might also account for the high D$_2$ occupancy variability. An approximately 5% decrease per decade in D$_2$ R availability above the age of 30 y has been reported (32). In our study, the mean age of the healthy volunteer group was lower than that of the patient groups (Table 1). However, the latter were still on their third decade, so the effect of age in the D$_2$ occupancy estimations was predicted to be minimal.

A risperidone arm was included in this study to keep the investigators masked, and D$_2$ occupancy comparisons between SB-773812 and risperidone were not planned. Subjects administered risperidone were scanned at the same time frames as subjects administered SB-773812 to keep the study masked, although risperidone $T_{\text{max}}$ has been reported to be around 1.5 h (7,33). Therefore, the D$_2$ occupancy values at 6 ± 2 h are representative of high plasma levels for both drugs, but risperidone peak D$_2$ occupancy values would probably be higher than those observed in our study at approximately 6 h after dosing, as previously reported (7). Nevertheless, the D$_2$ occupancy profiles from 6 to 24 h after dosing showed that SB-773812–induced occupancy was more stable with time. This stability is likely driven by the SB-773812 stable plasma profile, which, coupled with its moderate D$_2$ affinity, makes excessively high D$_2$ blockade in patients an unlikely achieve-

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Study A</th>
<th>Study B</th>
<th>Study C</th>
<th>Pooled</th>
</tr>
</thead>
<tbody>
<tr>
<td>EC$_{50}$ for D$_2$ (ng/mL)</td>
<td>90.7 ± 63.8</td>
<td>98.3 ± 15.6</td>
<td>81.2 ± 8.9</td>
<td>92.7 ± 13.5</td>
</tr>
<tr>
<td>Between-subject variability in D$<em>2$ EC$</em>{50}$</td>
<td>NA</td>
<td>82%</td>
<td>45%</td>
<td>73%</td>
</tr>
<tr>
<td>EC$<em>{50}$ for 5-HT$</em>{2A}$ (ng/mL)</td>
<td>2.3 ± 0.4</td>
<td>1.6 ± 0.8</td>
<td>NA</td>
<td>2.1 ± 0.5</td>
</tr>
<tr>
<td>Between-subject variability in 5-HT$<em>{2A}$ EC$</em>{50}$</td>
<td>56%</td>
<td>147%</td>
<td>NA</td>
<td>71%</td>
</tr>
</tbody>
</table>

NA = not available.
CONCLUSION

SPECT is a valuable tool for drug development when the appropriate radioligands are available and well characterized. The single-dose study in healthy volunteers demonstrated SB-773812 brain penetration and binding to target receptors in humans, supporting a go decision at early stages of development and providing preliminary estimates of target plasma concentrations and doses for further clinical development. Subsequent studies in patients with schizophrenia at repeated doses confirmed a high 5-HT$_{2A}$ occupancy, with moderate D$_2$ occupancy maintained over time, supporting a once-a-day dose regimen. Finally, the pharmacokinetics–receptor occupancy predictions were in reasonable agreement across studies and provided guidance for the selection of therapeutic doses to be tested in the clinical setting.

APPENDIX

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REFERENCES


Contribution of SPECT Measurements of D_2 and 5-HT_2A Occupancy to the Clinical Development of the Antipsychotic SB-773812

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