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Arterial spin labeling versus BOLD in pharmacological fMRI

A carefully controlled study allowed us to compare the sensitivity of ASL (arterial spin labeling) and BOLD (blood oxygen level dependent) fMRI for detecting the effects of the adenosine A2a antagonist tozadenant in Parkinson disease. Only ASL detected the direct effect of tozadenant. BOLD was more sensitive to a cognitive task, which (unlike most drugs) allows on-off comparisons over short periods of time. Neither ASL nor BOLD could detect a cognitive-pharmacological interaction. These results are consistent with the known relative advantages of each fMRI method, and suggest that for drug development, directly imaging pharmacodynamic effects with ASL may have advantages over cognitive-pharmacological interaction BOLD, which has hitherto been the more common approach to pharmacological fMRI.
Arterial spin labeling versus BOLD in pharmacological fMRI

Running Title: ASL vs BOLD in pharmacological fMRI

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Abstract

A carefully controlled study allowed us to compare the sensitivity of ASL (arterial spin labeling) and BOLD (blood oxygen level dependent) fMRI for detecting the effects of the adenosine A2a antagonist tozadenant in Parkinson disease. Only ASL detected the direct effect of tozadenant. BOLD was more sensitive to a cognitive task, which (unlike most drugs) allows on-off comparisons over short periods of time. Neither ASL nor BOLD could detect a cognitive-pharmacological interaction. These results are consistent with the known relative advantages of each fMRI method, and suggest that for drug development, directly imaging pharmacodynamic effects with ASL may have advantages over cognitive-pharmacological interaction BOLD, which has hitherto been the more common approach to pharmacological fMRI.

Introduction

Pharmacological magnetic resonance imaging (phMRI) uses fMRI to determine drug-induced changes in brain activity and has multiple applications for pharmaceutical development and efficacy testing. Before the development of functional MRI (fMRI), pharmacological brain imaging most often directly compared brain activity on drug to brain activity off drug (Herscovitch, 2001; McCulloch, 1982). Generally, phMRI studies have avoided this direct approach. Some used drugs with rapid onset and rapid decay of action, and correlated brain BOLD (blood oxygen level dependent) signal with noticeable transient physiological effects, e.g. repeated ratings of cocaine-induced “high” (Breiter et al., 1997). Other phMRI studies used drugs with rapid uptake and rapid elimination, with sequential measurements of plasma concentration, to detect brain changes with the expected pharmacokinetics (Bloom et al., 1999). Drug effects on functional connectivity have also been examined (Schwarz et al., 2007). The most common phMRI approach examines the interactive effects of a drug on the...
BOLD signal changes induced by a cognitive or sensory stimulus (Cole et al., 2012; Moeller et al.; Wise et al.). All of these study designs were motivated in part by limitations of BOLD fMRI, whose signal is nonquantitative and fluctuates artifactually over space and time (Iannetti et al., 2007).

By contrast, ASL (arterial spin labeling) is an fMRI method that produces a temporally stable signal. Additionally, ASL images reflect regional cerebral blood flow (rCBF) and thus allow relatively straightforward physiological interpretation. These advantages have led some recent drug discovery phMRI studies to use ASL (Wang et al., 2011; Zelaya et al., 2014 [in press]).

These considerations, and our experience with pharmacological PET (positron emission tomography) blood flow imaging (Black et al., 1997; Black et al., 2000; Black et al., 2005; Black et al., 2002; Hershey et al., 2003; Hershey et al., 2000; Hershey et al., 1998), led us to choose a pure pharmacological challenge approach with perfusion fMRI for a pharmacological challenge MRI study in Parkinson disease (Black et al., 2010b). However, we designed the study so that we would also have data from the more prevalent BOLD drug-task interaction design. The resulting data set allows a fair comparison of these two methods, i.e. subjects provided imaging data for both methods during the same imaging sessions, with similar drug concentrations, the same task, and similar total MRI acquisition times. Here we report the results of that comparison.

Materials & Methods

Study participants

Fourteen nondemented, nondepressed, ambulatory adults age 40–75 with idiopathic Parkinson disease, treated with a stable dose of levodopa but no dopamine agonists, participated in the clinical trial (registered at http://clinicaltrials.gov with identifier NCT00605553). Detailed inclusion and exclusion criteria were reported previously (Black et al., 2010a). The study was approved by the Washington University Human Research Protection Office (IRB) approval # 08-0059, and all subjects provided
written documentation of informed consent prior to participation.

**Study protocol**

In this single-subject crossover study, subjects were randomly assigned to one of two treatment groups: those assigned to group 1 took 60 mg of the adenosine A2a antagonist tozadenant (SYN115) twice daily for one week, waited for a one week washout period and then took a matching placebo twice daily for one week; those assigned to group 2 participated in the reverse order. The original report included additional subjects allocated to 20mg vs placebo, but for this report we focus only on the 60mg arms.

Subjects and investigators were blind to the group assignments. Neuroimaging was performed on the last day of each treatment week. On the morning of the scan day, they did not take their usual antiparkinsonian medications, but did take the last dose of tozadenant or placebo at approximately 6:00 AM. The timing of the fMRI assessments was planned to approximately bracket the time to maximal plasma concentration of tozadenant after chronic dosing. Subjects took 200 mg of carbidopa on arrival to the imaging center and then underwent two sets of MRI assessments, once before and once during an infusion of levodopa (LD). The study design was optimized for tozadenant rather than levodopa, and the dose of levodopa was relatively low by design, so analyses examining the effect of levodopa were secondary (see Supplementary Materials).

**Subject behavior**

Each scanning session included two perfusion MRI (ASL) runs while the subject performed the 2-back memory task, two control ASL runs while the subject fixated on a crosshair, and two block-design BOLD runs, each with 3 fixation blocks bracketing 3 task blocks. In each session, scans were obtained in the following order: fixation ASL, 2-back ASL, 2 BOLD runs, fixation ASL, and 2-back ASL. Thus in each session the ASL scans bracketed the BOLD runs. One subject was excluded from all analyses presented here because his 2-back task performance was less than 80% accurate.
Tozadenant had no statistically significant effect on 2-back performance (Campbell et al., 2010).

**MR image acquisition**

Both BOLD and ASL MRI data were acquired on the Siemens 3T Tim Trio with matrix head coil. BOLD-sensitive echo-planar images (EPI) were obtained with flip angle $90^\circ$, echo time (TE) 27 ms, repetition time (TR) 2000 ms, 36 planes with interleaved slice acquisition, field of view $256 \times 256$ mm, and voxel size $(4.0 \text{mm})^3$. Over a period of 4.33 min for each run, 130 volumes (frames) were acquired; the first 4 frames were discarded to ensure steady-state magnetization.

ASL images were acquired with the commercial Siemens pASL sequence (Wang et al., 2003b). Fifteen echo-planar readout slices with center-to-center slice distance 7.5 mm were acquired in the AC-PC plane with $64 \times 64 \times 3.4375 \text{mm}^3$ voxels in each plane, TR 2600 ms, TE 13.0 msec, and flip angle $90^\circ$. An $M_0$ image was followed by 31 tag-control pairs for a total acquisition time for each ASL run of 2.73 min.

Brain structure was assessed from sagittal MP-RAGE acquisitions with voxel size $(1.0 \text{mm})^3$, TR = 2400 msec, TE = 3.08 msec, TI = 1000 msec, flip angle = $8^\circ$. The structural images for each subject were inspected visually, images of lower quality were rejected, and the remaining 1-4 MP-RAGE images for each subject were mutually registered and averaged using a validated method (Black et al.).

**Image preprocessing**

BOLD images from each subject were preprocessed to reduce artifacts, including correction for intensity differences due to interleaved acquisition, interpolation for slice time correction, correction for head movement, and alignment to atlas space (Hershey et al., 2004). Image intensity was adjusted on a frame-by-frame basis so that each frame had a whole-brain modal value of 1000 (Ojemann et al., 1997). Frames were smoothed using a
6mm (FWHM) Gaussian filter and resampled to (3mm)$^3$ cubic voxels. To minimize motion-related artifact, frames were removed if framewise displacement exceeded 0.9mm (Siegel et al., 2014).

The 63 frames of the ASL run were smoothed using a 5.7mm (FWHM) Gaussian filter (resolution chosen to best match the final smoothing estimated from the BOLD images) and rigidly aligned using a validated method (Black et al., 2001a). Cerebral blood flow (CBF) was computed in each voxel for each tag-control EPI pair as described (Wang et al., 2003b). The aligned EPI images were also summed to facilitate later alignment steps, and the summed, aligned EPI images from each run were mutually aligned within each subject and summed across runs. The resulting summed EPI images from each subject were affine registered to a target image in Talairach and Tournoux space made using validated methods from these subjects’ structural MR images (Hershey et al., 2004). The products of the registration matrix from this step and the matrices from the within-run mutual registration step were used to resample the 31 tag–control pair CBF images from each run into atlas space images with (3mm)$^3$ cubic voxels in a single resampling step. To minimize motion-related artifact we removed tag–control pairs if framewise displacement in either EPI image exceeded 0.9mm (Siegel et al., 2014). One subject’s data was excluded from further analysis because over half of his frame pairs were removed due to head motion. The CBF images in atlas space from the remaining pairs were averaged to create one atlas-registered CBF image for each ASL run. Each CBF image was corrected to an idealized modal global (whole-brain) CBF of 50 mL/hg/min (Stewart et al., 2014).

**Statistical analysis**

**Analysis strategy**

The analyses were designed so that each ASL–BOLD comparison included the same scan sessions from the same group of subjects, and as nearly as possible the same image smoothness. Furthermore, the images used to compare the modalities were $t$ images from the same sample, and hence...
were commensurate. Statistical images were created for each imaging modality to examine the 2-back task effect, the interaction of the 2-back task with tozadenant, and a direct comparison of tozadenant versus placebo.

Statistical images

To identify regions of activation and deactivation, we used a mixed-effects approach with partitioned variance (Penny et al., 2007). First, for each study subject, we used a voxelwise general linear model (GLM) that included main effects of task (2-back vs. fixation), levodopa (during vs. before infusion) and drug (tozadenant vs. placebo). For each effect analyzed (drug, 2-back task, infusion and their interactions), SPM12b software (www.fil.ion.ucl.ac.uk/spm/) generated a contrast image for each subject from ASL data, and fIDL (http://www.nil.wustl.edu/~fidl/) did the same for BOLD images (also correcting for linear drift within each run). Note for each subject, every contrast image for ASL data was derived from the same set of scans, and similarly for the BOLD data. These single-subject contrast images were used as input to second-level SPM analyses based on a voxelwise general linear model with a covariate for subject age and a factor for sex. One-tailed one-sample t tests at each voxel tested whether the single-subject contrast images at that voxel were significantly less than or greater than zero, across subjects. After thresholding at the t value corresponding to uncorrected p=.001, multiple comparisons correction was performed with the cluster false discovery rate set at p=.05. Approximate anatomical locations of peaks in the statistical images were provided by the Talairach Daemon client (www.talairach.org) (Lancaster et al., 1997; Lancaster et al., 2000).

Results

Cross-modality image comparison

The final resolution of the 3×3×3mm ASL and BOLD images was similar (Table 1). Total acquisition time was about 25% longer for ASL than BOLD, but acquisition time for the data actually submitted to statistical analysis
was much more similar (Table 1), largely because each head movement lost 5.2 sec of data in the ASL data versus 2.0 sec in the BOLD data.

**Task activation**

The working memory task serves as a positive control, and significant regional activations were identified. The analysis using the ASL data identified one significant activation cluster (22 voxels = 0.6 ml, corrected p = 0.030, peak t = 5.88 at -32, -3, 57, left middle frontal gyrus, Brodmann area [BA] 6). The analysis using the BOLD data identified 12 significant clusters; the largest cluster also included -32, -3, 57 (515 voxels = 13.9 ml, corrected p < .001, peak t = 12.29 at -40, 3, 33 (left precentral gyrus, BA6) (see Suppl. Table 1). There were no significant deactivations in the ASL data, while the analysis using the BOLD data identified 11 significant deactivation clusters (the largest had volume 2142 voxels = 57.8 ml, corrected p < .001, peak t = 12.70 at -4, -54, 12, left posterior cingulate, BA29) (see Suppl. Table 2).

**Drug effect**

The task-drug interaction (tozadenant × 2-back) showed no significant results for ASL or BOLD. However, the same ASL data revealed significant rCBF decreases on tozadenant in the thalamus bilaterally (Table 2, Suppl. Figure 1). There were no significant clusters of increased rCBF. As expected, the same contrast with the BOLD data found no significant clusters of activation or deactivation. Table 3 summarizes all these contrasts.

**Discussion**

Cognitive-pharmacological interaction is a common phMRI approach. However, in this study neither ASL nor BOLD analyses detected significant clusters for the interaction of tozadenant with 2-back task activation, whereas directly comparing rCBF on versus off drug using ASL did reveal significant differences. The drug-induced rCBF decreases detected by ASL are in the thalamus, consistent with animal studies suggesting that
Adenosine A2a receptor antagonists inhibit neuronal activity in the indirect pathway, including in pallidal afferents to thalamus (Black et al., 2010b).

Positive controls built into the experiment confirm that the absence of significant drug effects in the BOLD analysis cannot be comfortably attributed to inadequate image quality or limited data: these same scans were quite adequate to detect significant cognitive (2-back task) effects in a pattern consistent with previous functional imaging studies on working memory (Barch et al., 2012; Bledowski et al., 2010). BOLD is generally more sensitive than ASL for comparisons like this one that can be made over very brief time intervals (a minute or so) (Wang et al., 2003a). However, noise in BOLD data worsens as the time between activation and control acquisitions increases (Aguirre et al., 2002; Ollinger et al., 2001), and this temporal instability likely explains why the BOLD data could not detect direct drug effects between sessions. By contrast, the temporal stability of ASL may suit it better to measure the effects of medications, which after all often have been optimized to require only a few doses a day, and hence have slow onset and wearing off of action (Aguirre et al., 2002; Wang et al., 2011; Zelaya et al., 2014 [in press]).

Comparing scans from different sequences was feasible here because both BOLD and ASL data were acquired during the same scan sessions in the same subjects, and because the images submitted to statistical analysis were of similar spatial smoothness. Also, in each scan session, half of the ASL scans came before and half after the two BOLD runs, so that any slowly evolving effects of practice, fatigue or drug should be similar on average for the two modalities. Limitations of this study include the imperfect matching between ASL and BOLD of total acquisition time and original voxel size. The different original voxel size is in part a technical limitation because ASL is best suited to acquiring read-out planes in inferior-to-superior order, whereas BOLD can be acquired with even and odd read-out planes interleaved.
Decreased thalamic rCBF with tozadenant was also the most significant result of the previously published analysis of ASL data from this study (Black et al., 2010b), but the present analysis detected fewer significant voxels. This is probably because in order to match the BOLD data, the present analysis excluded half the ASL data (acquired during additional behavior states for which were no comparable BOLD data) and smoothed the data less than in the published analysis. We now also excluded subjects with excessive movement or poor 2-back task performance, censored frames for head motion, and improved the correction for global CBF.

One additional advantage of this study comes from the following consideration. A drug that produces symptomatic effects, for instance a feeling of calm, may cause secondary effects on neuronal activity via the effect on emotional state in addition to any direct neuronal effects (including the neuronal effects that themselves produce the sense of calm). The same reasoning applies to any placebo effect that may be heightened if the subject notices any drug effect. In this study, most subjects were unable to distinguish whether they were taking active drug or placebo, allowing more straightforward interpretation of the drug’s effects on neuronal activity.

Conclusions

In summary, these data offer direct, head-to-head evidence that phMRI using ASL and pure pharmacologic activation may be more sensitive than task-interaction BOLD phMRI.

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### Table 1: Comparison of BOLD and ASL images

|                        | BOLD      | ASL       |
|------------------------|-----------|-----------|
| Total acquisition time per scanning session | 8.7 min   | 10.9 min  |
| Acquisition time per session, limited to frames retained after motion censoring (mean ± SD) | 8.5 ± 0.1 min | 9.2 ± 1.1 min |
| FWHM (x × y × z) *    | 10.1 × 10.5 × 9.0 mm | 9.4 × 10.5 × 11 mm |

* Average of the FWHM estimates across SPM analyses.

### Table 2: Significant clusters of decreased rCBF on tozadenant

| Significant clusters | cluster volume, voxels (cm³) | 25 (0.68) | p (FDR) | .004 | peak t | 5.67 | atlas location | 8, -15, 9 | anatomical location of peak t | Right medial dorsal nucleus of thalamus |
|----------------------|-------------------------------|-----------|---------|------|-------|-----|----------------|-----------|---------------------------------|--------------------------------------|
|                      | cluster volume, voxels (cm³) | 10 (0.27) | p (FDR) | .049 | peak t | 5.17 | atlas location | -8, -21, 9 | anatomical location of peak t | Left medial dorsal nucleus of thalamus |

* Table includes all clusters with FDR-corrected p<.05.
Table 3: Summary of activation clusters for all contrasts

| Task Contrast                  | Number of Significant Clusters |
|-------------------------------|--------------------------------|
|                               | ASL | BOLD |
| 2-back activation             | 1   | 12   |
| 2-back deactivation           | 0   | 11   |
| Tozadenant × 2-back activation| 0   | 0    |
| Tozadenant × 2-back deactivation| 0  | 0    |
| Tozadenant activation         | 0   | 0    |
| Tozadenant deactivation       | 2   | 0    |
Supplementary Material

Supplementary Table 1: Significant activations during 2-back task (BOLD)

| #  | cluster volume, voxels | cluster volume, cm³ | p (FDR) | peak t value | anatomical location * |
|----|------------------------|----------------------|---------|--------------|-----------------------|
| 1  | 515                    | 13.9                 | <.00 1  | 12.29       | left precentral gyrus (BA 6) |
| 2  | 471                    | 12.7                 | <.00 1  | 9.80        | right superior frontal gyrus (BA 6) |
| 3  | 327                    | 8.8                  | <.00 1  | 10.75       | right inferior temporal gyrus (BA20) |
| 4  | 224                    | 6.0                  | <.00 1  | 9.40        | left posterior lobe |
| 5  | 223                    | 6.0                  | <.00 1  | 8.70        | right middle frontal gyrus (BA9) |
| 6  | 166                    | 4.5                  | <.00 1  | 7.53        | left caudate |
| 7  | 163                    | 4.4                  | <.00 1  | 6.38        | right postcentral gyrus (BA2) |
| 8  | 142                    | 3.8                  | <.00 1  | 13.42       | right insula (BA 13) |
| 9  | 127                    | 3.4                  | <.00 1  | 12.94       | left claustrum |
| 10 | 108                    | 2.9                  | <.00 1  | 8.41        | left cerebellum |
| 11 | 47                     | 1.3                  | <.00 1  | 7.69        | left superior parietal lobule (BA7) |
| 12 | 22                     | 0.6                  | .016    | 6.30        | left superior frontal gyrus (BA10) |
Supplementary Table 2: Significant deactivations during 2-back task (BOLD)

| #  | cluster volume, voxels | cluster volume, cm³ | p (FDR) | peak t | atlas location of peak t value | anatomical location * |
|----|------------------------|---------------------|---------|--------|--------------------------------|----------------------|
| 1  | 2142                   | 57.8                | <.001   | 12.7   | 4 -54 12                       | right posterior cingulate (BA29) |
| 2  | 507                    | 13.7                | <.001   | 8.03   | 4 12 0                         | right caudate         |
| 3  | 360                    | 9.7                 | <.001   | 7.76   | -38 -18 21                     | left insula (BA13)    |
| 4  | 132                    | 3.6                 | <.001   | 8.78   | -44 -75 30                     | left angular gyrus (BA39) |
| 5  | 104                    | 2.8                 | <.001   | 6.72   | 52 -75 21                      | right middle temporal gyrus (BA19) |
| 6  | 65                     | 1.8                 | <.001   | 6.81   | -56 0 -15                      | left middle temporal gyrus (BA21) |
| 7  | 59                     | 1.6                 | <.001   | 7.57   | 26 6 -21                       | right uncus (BA28)    |
| 8  | 46                     | 1.2                 | .001    | 9.74   | 10 -51 -42                    | right cerebellar tonsil |
| 9  | 42                     | 1.1                 | .001    | 6.50   | 32 -72 -33                    | right pyramis         |
| 10 | 40                     | 1.1                 | .001    | 6.68   | -34 -18 0                      | left lentiform nucleus |
| 1  | 29                     | 0.8                 | .006    | 7.18   | 14 39 54                       | right superior frontal gyrus (BA8) |

* BA, Brodmann area
**Supplementary Figure 1:** Coronal, axial and sagittal sections showing the significant CBF decreases on tozadenant 60mg twice daily. Colored voxels indicate p<.001 uncorrected; the corrected p value is .004 for the cluster in right thalamus and .049 for the left (see also Table 2).

**Supplementary Material (continued)**

**Materials & Methods (secondary levodopa analyses)**

The data come from the same scans as reported in the main body of the paper. The study design was optimized for tozadenant rather than levodopa (LD), and the LD dose was relatively low, so analyses examining the effect of levodopa were secondary.

The approach was identical to that reported for the task and tozadenant analyses in the main body of the paper. To investigate the effects of LD we created statistical images of the LD effect (comparing scans acquired during the LD infusion to scans prior to infusion), of the interaction of the 2-back task with LD, and of the 3-way interaction of the 2-back task, LD and tozadenant.

**Results (secondary LD analyses)**

There were no significant clusters for the pure LD effect, the task-LD interaction, or the 3-way interaction in either the ASL or the BOLD images.
