Sensory and Motor Systems

The Largest Response Component in the Motor Cortex Reflects Movement Timing but Not Movement Type

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Abstract

Neural activity in monkey motor cortex (M1) and dorsal premotor cortex (PMd) can reflect a chosen movement well before that movement begins. The pattern of neural activity then changes profoundly just before movement onset. We considered the prediction, derived from formal considerations, that the transition from preparation to movement might be accompanied by a large overall change in the neural state that reflects when movement is made rather than which movement is made. Specifically, we examined “components” of the population response: time-varying patterns of activity from which each neuron’s response is approximately composed. Amid the response complexity of individual M1 and PMd neurons, we identified robust response components that were “condition-invariant”: their magnitude and time course were nearly identical regardless of reach direction or path. These condition-invariant response components occupied dimensions orthogonal to those occupied by the “tuned” response components. The largest condition-invariant component was much larger than any of the tuned components; i.e., it explained more of the structure in individual-neuron responses. This condition-invariant response component underwent a rapid change before movement onset. The timing of that change predicted most of the trial-by-trial variance in reaction time. Thus, although individual M1 and PMd neurons essentially always reflected which movement was made, the largest component of the population response reflected movement timing rather than movement type.

Key words: condition-invariant signal; dPCA; movement initiation; movement triggering; reaction time; state space

Significance Statement

The activity of neurons often conveys information about externally observable variables, such as the location of a nearby object or the direction of a reach made to that object. Yet neural signals can also relate to “internal” factors: the thoughts and computations that link perception to action. We characterized a neural signal that occurs during the transition from preparing a reaching movement to actually reaching. This neural signal conveys remarkably accurate information about when the reach will occur, but carries essentially no information about what that reach will be. The identity of the reach itself is carried by other signals. Thus, the brain appears to employ distinct signals to convey what should be done and when it should be done.
Introduction

The responses of individual neurons are often characterized in terms of tuning: how the firing rate varies across different stimuli or behaviors (“conditions”). Additionally, neural responses may contain untuned features that are shared across many conditions, such as an abrupt rise in firing rate after the onset of any stimulus. These untuned response features may appear nonspecific, and thus of secondary interest. However, there is evidence that response features can be correlated across conditions, yet still carry computationally relevant information. Neural activity in the prefrontal cortex contains a large response component reflecting the passage of time (Machens et al., 2010), and time-varying signals have also been observed in the premotor cortex during anticipation of an informative cue (Confaïs et al., 2012). A related example is the time-encoding urgency signal observed during decision making, which is shared across neurons that encode different choices in both the oculomotor system (Churchland et al., 2008; Hanks et al., 2011) and the premotor cortex (Thura et al., 2012). Here we investigate another possible “untuned” signal in the motor/premotor cortex: one that arises after the desired target is known, at the time of the sudden transition from preparation to movement.

We were motivated by the observation that motor cortex neurons sometimes display broadly tuned movement-period responses (Fortier et al., 1993; Crammond and Kalaska, 2000)—e.g., a rise in rate for all directions—such that tuning models benefit from an omnidirectional term (Georgopoulos et al., 1986; Morani and Schwartz, 1999). More generally, many studies identify significant proportions of neurons with responses that are task-modulated yet not strongly selective for the parameter being examined (Evarts, 1968; Weinrich et al., 1984; Hocherman and Wise, 1991; Riehle et al., 1994; Messier and Kalaska, 2000). These findings argue that there must be some aspect of neural responses—i.e., some response “component”—that is at least moderately correlated across conditions. What are the temporal properties of such a signal and is its timing predictive of behavior? Does the signal make a small or large contribution to the overall population response? Is the signal merely correlated across conditions (“condition-correlated”)? Or might it be nearly identical across conditions (“condition-invariant”) and thus untuned in the traditional sense?

These questions derive both from a general desire to fully characterize the response during movement and from specific theoretical considerations. A condition-invariant signal (CIS) could, despite its seeming lack of specificity, be important to the overall computation performed by the population. Presumably there is a large change in computation just before movement onset, at the moment when the motor system transitions from preparing to move while holding a posture (Kurtzer et al., 2005) to generating the muscle activity that will drive the desired movement. Consistent with the idea of a change in computation, neural tuning changes suddenly and dramatically at a point ~150 ms before movement (Churchland et al., 2010) so that a neuron’s “preference” during movement can be quite unrelated to its preference during preparation (Wise et al., 1986; Crammond and Kalaska, 2000; Kaufman et al., 2010). A similar transition is observed at the population level: population dynamics are relatively stable and attractor-like during preparation but become strongly rotational just before movement onset (Churchland et al., 2012). This sudden change in network properties is presumably driven by an appropriately timed input (which could itself be the output of a computation that decides when to move; Romo and Schultz, 1987; Thaler et al., 1988; Schurger et al., 2012; Murakami and Mainen, 2015). One might initially expect a “triggering” input to be tuned (Johnson et al., 1999; Erhagen and Schönér, 2002). Yet theoretical considerations suggest that a simple, condition-invariant change in input is sufficient to trigger large changes in network dynamics and tuning (Hennequin et al., 2014). In particular, a recent neural network model of motor cortex (Sussillo et al., 2015) uses a condition-invariant input to trigger a change in dynamics that initiates movement. The model’s population-level responses resemble the empirical neural responses, and from inspection both clearly show at least some features that are invariant across conditions.

Critically, there are many ways in which activity patterns can be correlated across conditions. Only a minority of such possibilities involve a true CIS at the population level: that is, a signal that is nearly identical across conditions. Is a CIS present in motor cortex? On a trial-by-trial basis, does it exhibit timing locked to target onset, the go cue, or movement onset? Only the latter would be consistent with the role in movement triggering suggested by the model of Sussillo et al. (2015).

We found that a CIS was not only present but was the largest aspect of the motor cortex response—considerably larger than any of the condition-specific (tuned) re-
sponse components. The CIS resembled the previously reported omnidirectional or “speed-tuned” response component (Georgopoulos et al., 1986; Moran and Schwartz, 1999), but was essentially invariant with reach speed, distance, and curvature. In addition, the CIS underwent a large and sudden change ~150 ms before movement onset. The timing of this change was an excellent predictor of reaction time (RT) on a trial-by-trial basis. Finally, the dimensions in neural state space that were occupied by the CIS were almost perfectly orthogonal to the dimensions occupied by the condition-specific components. Overall, the profile, timing, and population-level manifestation of the CIS were remarkably similar to the structure naturally produced by the model of Sussillo et al. (2015). Our findings thus suggest a potential role for a large response component that initially appears nonspecific yet reflects movement timing very precisely.

Materials and Methods
The key features of the task and analyses are described in the Results. Below we detail all aspects of the apparatus, task, neural recordings, muscle recordings, data preprocessing, analyses, and controls.

Subjects and task
Animal protocols were approved by the Stanford University Institutional Animal Care and Use Committee. Experiments employed two adult male rhesus monkeys (Macaca mulatta), J and N, performing a delayed-reach task on a frontoparallel screen (Churchland et al., 2010, 2012; Kaufman et al., 2013). The monkey initially fixated a central spot with his eyes and touched it with a cursor. The cursor was projected slightly above the right fingertip, which was tracked optically. The task involved a large number of conditions—i.e., different target locations and reach paths—which was useful when attempting to identify response components that are invariant across conditions. On one-third of trials (“no-barrier” conditions) a lone target appeared within a frame around the workspace. On another one-third of trials (“maze” conditions) a target and ≤9 virtual barriers appeared. The remaining one-third of trials (“maze-with-distractor” conditions) were identical to the maze trials but included two distractor “targets” that were unreachable due to the barrier locations. The same set of target positions was used for the no-barrier, maze, and maze-with-distractor conditions. When barriers were present, the monkey had to perform a curved reach or the cursor would collide with and “stick” to the barrier. This paradigm evoked both a large response component that initially appears nonspecific yet reflects movement timing very precisely.

A randomized delay period separated target onset from a go cue. During the delay, targets jittered slightly (2–3 mm), indicating to the monkey that he could not yet reach or saccade. The go cue consisted of three simultaneous and salient cues: jitter ceasing, the targets changing from open to filled, and the central spot disappearing. Juice reward was delivered if the monkey swiftly reached to the target then held it for 450 ms (monkey J) or 700 ms (monkey N).

Delay-period statistics
The delay period lasted 0-1000 ms. Different datasets employed different delay-period statistics, depending on the analyses we wished to apply. Three datasets (JC, NAC, and NS) were collected with the primary goal of analyzing trials with longer delays. Longer delays enabled examination of the transition between a relatively stable plateau of preparatory activity and subsequent movement-related activity. To this end, delays of 450-1000 ms were approximately twice as probable as delays of 0-450 ms. Three further datasets (JAD1, JAD2, NAD) were recorded with the goal of characterizing the single-trial relationship between neural activity and response time (RT). For these datasets, delay durations of 0, 100, 200, and 500 ms were intentionally over-represented. These dataset names end with “D,” indicating that this set of discrete delays was over-represented. This allowed key analyses to be restricted to a set of trials with the same delay, removing the potential confound that RT can vary with delay. For these datasets most trials (78, 78, and 84% for datasets JAD1, JAD2, and NAD) used one of the discrete delays, with roughly equal probability. The remaining trials had random delays from 0 to 1000 ms as above. Because these datasets were each collected in a single day using implanted multielectrode arrays, the monkeys were not anticipating the over-represented delay durations.

Most analyses focused on the transition from movement preparation to movement and thus used only trials with >450 ms delays (datasets without discrete delays) or 500 ms delays (datasets with discrete delays). For analyses of the single-trial relationship with RT, we focused on datasets with discrete delay durations. For simplicity of presentation, for these analyses only trials with no delay (“zero delay”) or a 500 ms delay (“long delay”) are shown. All results were similar for delays of 100 or 200 ms.

Catch trials and trial counts
Several types of unanalyzed catch trials ensured the task was performed as desired. In particular, we presented novel mazes made by randomly removing barriers from a standard maze (10–15% of all trials), or randomly placing the target and two barriers (0–10% of all trials). These trials ensured that the monkey had to solve each trial independently, as similar-looking mazes could have different solutions.

Delay periods were randomly chosen on each trial. Conditions were organized in pseudorandom blocks. The array datasets had 3352, 2340, 2622, and 3590 successful trials (datasets JAD1, JAD2, NAD, and NAC) from a single session. For the “discrete delay” datasets (JAD1, JAD2, and NAD) there were ~250–500 usable trials for...
each of the four over-represented delays. Usable trials excluded catch trials, failed trials (e.g., if a barrier were struck), rare trials with an unusual velocity profile that did not allow a reliable RT measurement, and trials with a very short RT (in rare instances where the monkey “jumped the gun”) or an overly long RT (in rare instances where the monkey was presumably distracted). Datasets that included single-unit recordings (JC and NS) contained an average of 336 and 305 usable trials per unit.

Neural and muscle recordings
For both monkeys, we first performed single-electrode recordings (datasets JC and NS) using moveable tungsten microelectrodes (Frederick Haer) and a Plexon Multichannel Acquisition Processor (Plexon). These recordings included the caudal portion of the dorsal premotor cortex (PMd) and both surface and sulcal primary motor cortex (M1). All units recorded with single electrodes were well-isolated single neurons recorded from regions where microstimulation produced movement of the arm (typically the upper arm and/or shoulder). Each monkey was then implanted with two 96-electrode silicon arrays (Blackrock Microsystems), located in M1 and caudal PMd, as estimated from anatomical landmarks and previous mapping with microstimulation. Spikes were sorted off-line using custom software (MKsort, https://github.com/ripple-neuro/mksort). For array recordings, both single units and stable multiunit isolations (typically two neurons whose spikes could not be reliably separated) were analyzed. A strong CIS (see below) was present regardless of whether a dataset involved pure single-unit isolations or a mixture of single-unit and multiunit isolations. This is unsurprising: dimensionality reduction techniques, such as demixed principal component analysis (dPCA) or PCA, typically produce nearly identical results regardless of whether isolations involve one unit or a few units. These techniques are forgiving because the components needed to compose the responses of a single neuron are the same components needed to compose the summed response of >1 neuron. All neural recordings were from the left hemisphere. Array recordings produced datasets JAD1, JAD2, NAD, and NAC, and were included in dataset JC.

We analyzed all units where the firing rate range (over conditions and times) was greater than the maximal SEM (for all conditions and times). This signal-to-noise (SNR) criterion does not insist on any particular form of response or tuning—only that there be some response. For dataset JAD1, 116 of 123 units passed the SNR criterion; for dataset JAD2, 136 of 171 units passed; for dataset JC, 186 of 278 units passed; for dataset NAD, 172 of 188 units passed; for dataset NAC, 213 of 223 units passed; for dataset NS, 118 of 118 units passed. Of these, 67, 28, 108, 62, 58, and 118 were considered single units (datasets JAD1, JAD2, JC, NAD, NAC, and NS). For all analyses, results were similar when data from PMd and M1 were analyzed separately. These recordings were therefore pooled.

Data preprocessing involved three steps. First, spike trains were smoothed with a Gaussian (28 ms SD). Second, the firing rate was averaged across trials of the same type (excepting analyses of single trials; see below). We computed two averages: one with data aligned to target onset and one with data aligned to movement onset. Third, the firing rate of each neuron was normalized to prevent analyses from being dominated by a few high-rate neurons; this is especially important (Yu et al., 2009) when performing PCA-based analyses. To normalize without overamplifying the greater noise associated with low firing rates, we “soft normalized”: for each neuron we normalized the firing rate by its range (across all times and conditions) plus a constant, chosen to be 5 spikes/s. This choice follows our previous work, and was made before performing analyses. Results were extremely similar and sometimes stronger if we used a soft-normalization constant of zero.

Electromyographic (EMG) recordings used hook-wire electrodes (44 gauge with a 27 gauge cannula; Nicolet Biomedical), inserted percutaneously into the muscles of the right arm. Electrodes were inserted with the monkey awake and calm, with one recording per session. For monkey J, recordings were made sequentially from trapezius, latissimus dorsi, pectoralis, triceps brachii, medial and lateral aspects of the biceps, medial and anterior, medial, and posterior aspects of the deltoid. The recording from the triceps was excluded because it was not sufficiently modulated during the task. For monkey N, recordings were made from proximal, middle, and distal aspects of the trapezius, latissimus dorsi, pectoralis, triceps brachii, medial and lateral aspects of the biceps, and anterior, medial, and posterior aspects of the deltoid. Two recordings were made for each deltoid site. The recordings from the triceps and latissimus dorsi were excluded because they were not sufficiently modulated during the task. Raw EMG signals were band-pass filtered (150–500 Hz, four pole, 24 db/octave), differentiated, rectified, smoothed with a Gaussian (15 ms SD), and averaged across trials (Kaufman et al., 2013).

Projections of neural data
We identified response components by projecting the population response onto dimensions of interest. We began with a matrix, $R$, of trial-averaged neural responses (or EMG, for one analysis). Each of $n$ columns contained the normalized response of one neuron over time, with responses concatenated across conditions. To project the data onto a given dimension we computed $\hat{x} = R\hat{w}$, where $\hat{w}$ is a set of weights specifying the dimension. The projection $\hat{x}$ is therefore a weighted average of neurons’ firing rates. We refer to the projected activity pattern as a “component” of the population response, because the activity of any given neuron can be (approximately) composed of a weighted sum of multiple such components. This use of the term “component” follows the usage of Kobak et al. (2016) and others. Note that this use of “component” is not synonymous with “principal component,” which refers to a component of the neural covariance matrix and thus corresponds to a neural dimension.

Many studies have been concerned with how to best find projections given different goals and hypotheses. In
this study the most important projection method uses dPCA (Machens et al., 2010; Brendel et al., 2011) to find the dimensions \( \mathbf{w} \). This application of dPCA is detailed more thoroughly in the next section.

We also use a number of other projection methods, including standard PCA, and simply computing the mean across neurons (equivalent to setting all weights to 1/n). Two analyses use the pPCA method (Churchland et al., 2012), and in one case we used a classifier trained via a supervised algorithm. In every case it should be stressed that the projections shown (i.e., the response components) are simply linear weightings of the recorded neural responses. The use of multiple methods is desirable because no single method can capture all aspects of the response (e.g., the mean captures some aspects of the response and hides others).

All projection methods used here employ orthonormal dimensions. The orthogonality of these dimensions does not impose orthogonality on aspects of the neural response; it is simply a way of choosing a coordinate system. An orthonormal basis makes interpretation simpler: among other benefits, it allows each component to be independently quantified in terms of variance explained, making it harder to unintentionally interpret weak structure as meaningful. In all cases, when a percentage of variance is quoted, it is the fraction of the variance captured in the low-dimensional space (10–12 dimensions).

Identifying the CIS via dPCA

Many of our central analyses sought to determine whether there exist neural dimensions that segregate condition-specific (“tuned”) components from condition-invariant components of the population response. By “condition-specific” we mean that different conditions (reach directions, curvatures, etc.) evoke different responses when the population response is projected onto that dimension.

By “condition-invariant” we mean that the response varies with time but is similar across conditions when projected onto that dimension. To address this question, we applied dPCA (Machens et al., 2010; Brendel et al., 2011), a variant of PCA. dPCA leverages information normally discarded by PCA: each row of the data matrix \( \mathbf{R} \) is assigned labels. Here, those labels indicated the condition and time for which that set of firing rates was recorded. dPCA then finds a matrix \( \mathbf{W} \) that produces a projection \( \mathbf{X} = \mathbf{R} \mathbf{W} \). Each column of \( \mathbf{W} \) is a dimension and each column of \( \mathbf{X} \) is a component of the population response. Like PCA, dPCA attempts to find a projection that captures much of the variance in \( \mathbf{R} \), so that \( \mathbf{R} = \mathbf{XW} \). Unlike PCA, dPCA attempts to find \( \mathbf{W} \) such that the resulting columns of \( \mathbf{X} \) covary strongly with one label or the other. In the present case, dPCA attempts to find \( \mathbf{W} \) such that some columns of \( \mathbf{X} \) (some components) vary with time but not condition and other columns vary across conditions but not with time. As will be discussed below, such segregation is not necessarily possible: in general there will not exist \( \mathbf{X} \) with the desired properties. Indeed, in the present study, dPCA always found components that varied primarily with time (and not condition) but never found components that varied primarily with condition and not time. We therefore divided the components found by dPCA into two groups: condition-invariant (reflecting primarily time) and condition-specific (reflecting both condition and time). We refer to the group of condition-invariant components collectively as the CIS.

As a technical note, dPCA (unlike PCA) requires that the number of dimensions be specified in advance. Prior analyses indicate that 6–8 dimensions capture much of the condition-specific structure of the data (Churchland et al., 2010). We therefore wished that dPCA should capture a similar amount of condition-specific structure, in addition to any condition-invariant structure that might be present. We empirically picked the number of requested dimensions such that dPCA returned eight condition-specific dimensions (defined as containing <50% condition-invariant variance). In principle this might have necessitated requesting exactly eight dimensions (if all structure were tuned) or many more than eight (if little structure were tuned). In practice it was only necessary to request modestly more than eight total dimensions. For example, for dataset JAD1 we requested 10 total dimensions, which yielded two condition-invariant response components and eight condition-specific response components. The choice of eight condition-specific components is an arbitrary but reasonable cutoff. We always found a strong CIS regardless of the exact number of dimensions requested. dPCA identified dimensions \( \mathbf{W} \) based on the population response from \( -200 \) to +400 ms relative to target onset and \( -300 \) to +600 ms relative to movement onset. The data matrix being analyzed contained trial-averaged firing rates for long-delay trials (trials with delay periods >450 ms). For subsequent analyses of trial-to-trial variability in RT, we projected data from individual trials, including zero-delay trials, onto the same dimensions. The probabilistic-model version of dPCA was used (from the Python code available online associated with Brendel et al., 2011). We measured for each response component the marginal variances (Machens et al., 2010; Brendel et al., 2011), which indicate how much of a component’s variance was condition-specific (activity varying with condition or with both time and condition) versus condition-invariant (activity varying with time alone).

Because EMG responses were lower dimensional than neural responses, for the EMG datasets dPCA was performed at an overall dimensionality that returned three condition-specific dimensions. The resulting 4–5 dimensions (monkey J, N) accounted for 95–97% of the total variance in the EMG data. This reduced number of dimensions did not produce the differences between neural and muscle data: repeating the analysis on neural data using 4–5 dimensions yielded essentially identical results to those obtained with more dimensions.

Note regarding interpretation of the segregation produced via dPCA

Below we describe a key interpretational point regarding the dPCA method. The cost function optimized by dPCA attempts to find \( \mathbf{W} \) such that each column of \( \mathbf{X} \) (each response component) varies with exactly one of the pro-
vided labels (time and condition in this study) and not with the other(s). Yet as stated above, this segregation is not in general possible. In the present case, this has two implications. First, it is not guaranteed that dPCA will be able to find components that vary with condition but not with time; perhaps every component that strongly reflects condition also reflects time (this was indeed true of our data). Second, it is similarly not guaranteed that dPCA will be able to find components that vary with time but not condition; it may be that every component that strongly reflects time also reflects condition.

This last fact is worth stressing because many individual neurons exhibit what we refer to as “condition-correlated” structure: responses that are different across conditions, yet display an increase (or decrease) in firing rate that has a somewhat similar time course across conditions. Yet this structure at the single-neuron level is not sufficient, in and of itself, to indicate condition-invariant structure at the population level. Would dPCA, when applied to a population of such neurons, inevitably find condition-invariant components? In short, it would not. This can be demonstrated empirically (see Results) or formally via construction, as follows. Consider a simple case in which each neuron’s response \( r_n \) is a linear combination of two independent components \( x_1 \) (which will also be functions of condition \( c \) and time \( t \)) and \( x_2 \). Let both \( x_{1,c,t} \) and \( x_{2,c,t} \) be condition-specific, but suppose \( x_{1,c,t} \) contains an overall correlation between conditions. Due to the correlation of \( x_{1,c,t} \) across conditions, the responses \( r_n \) will also have shared response features across conditions. Nevertheless, it is not in general possible to find a linear combination of the \( x_{i,c,t} \)’s that is condition-invariant. A linear combination of the \( x_{i,c,t} \)’s is equivalent to a linear combination of \( x_{1,c,t} \) and \( x_{2,c,t} \). Since these components are independent, finding a condition-invariant linear combination is equivalent to solving the following system of \((C - 1)T\) equations, where \( C \) is the number of conditions (here, 2), and \( T \) is the number of time points:

\[
\sum_{i=1:2} p_i x_{i,c,t} = \sum_{i=1:2} p_i x_{i,c+1,t}
\]

for all times \( t \) and all pairs of conditions \( c \) and \( c + 1 \) (this is a sufficient constraint to ensure that all pairs of conditions are equal, since equality is transitive).

The number of free variables \( p_i \) is equal to the number of components \( D \), which in this example is 2. In general, then, this system is not solvable if \((C - 1)T > D\), which will be true for even modest numbers of times and conditions. The presence of correlated structure within \( x_{1,c,t} \) (and/or \( x_{2,c,t} \)) would not in general change this fact. In practice, then, it would be rare for condition-correlated responses to coincidentally produce a fully condition-invariant component. As one example, choose \( x_{1,c,t} = g_c \sin(t) \) and \( x_{2,c,t} = h_c \sin(3t) \), with \( g_c \) and \( h_c \) being positive scalars that vary with condition. Both \( x_{1,c,t} \) and \( x_{2,c,t} \) would be perfectly condition-correlated, yet no linear combination of \( x_{1,c,t} \) and \( x_{2,c,t} \) would be condition-invariant.

**Control: producing synthetic peristimulus time histograms with matched spectral content**

To illustrate empirically that condition-invariant components are not found in “generic” data, we generated synthetic peristimulus time histograms (PSTHs) with the same frequency content as the original neurons. Each unit was matched with a corresponding synthetic PSTH. The steps below were performed on the vector containing the trial-averaged firing rate over time for one condition. We first preprocessed each vector by smoothing lightly (10 ms SD Gaussian) to reduce the small discontinuity between target-aligned and movement-aligned data, then multiplying by a Hann window. The Fourier transform was performed, and the magnitude of the result was computed at each frequency (i.e., the square root of power spectral density). These curves were averaged over conditions to give the overall power-by-frequency curve for that unit. To construct a synthetic PSTH for each condition, we chose a random phase for each frequency component, then took the inverse Fourier transform.

**Control: removing the CIS from the neural responses**

To ask whether condition-invariant components (collectively the CIS) might result from the rectification of firing rates at zero, we removed the true condition-invariant components, rerectified firing rates, then applied dPCA. Specifically, we projected the population response onto the eight condition-specific dimensions identified by dPCA, then transformed the data back to the original \( n \)-dimensional space. This produced as many PSTHs as the original neurons. We rescaled and centered each “neuron’s” response to restore its original mean and range of firing rates. Finally, we set all negative firing rates to zero. This resulted in a population of surrogate neurons that are responsive and have positive firing rates, yet should have no “true” CIS. Thus, a strong CIS in this control population would indicate that rectified firing rates could create an artifactual CIS.

**Control: adding a condition-correlated component**

We constructed additional surrogate data that resembled the empirical data but lacked condition-invariant components. For each empirical condition-invariant component, we constructed a new component with the same time course, but with varying amplitude across conditions. That is, we created components that were condition-correlated but not condition-invariant. These components were recentered to have a zero mean during the baseline period (before target/maze onset), and then were added to the response of each neuron. Specifically, to each neuron’s response \( r_{n,c,t} \) we added \( w_{ni} k_c x_{i,t} \), where \( w_{ni} \) is the neuron’s original weight for the \( i^{th} \) condition-invariant component, \( x_{i,t} \) is the time course of the \( i^{th} \) new condition-correlated component, and the coefficients \( k_c \) were chosen randomly from a unit-variance Gaussian distribution. We rectified the resulting firing rates (setting all negative rates to zero). These operations largely preserved the time course of each neuron’s across-condition mean (because the \( k_c \)’s were zero-mean). Because the new components were condition-correlated, the responses of most neurons...
were strongly condition-correlated. Yet because the original condition-invariant components are now “contaminated” with condition-specific components of the same time course, the surrogate population should have no separable condition-invariant components.

Identifying a speed-predicting dimension
To identify a speed-predicting dimension, we began with the same neural data matrix $R$ used for PCA and dPCA. We then regressed the trial-averaged speed profile for each condition against $R$: $\hat{s} = R\hat{w} + b$, where $\hat{s}$ is the vector produced by taking the speed profile for each condition and concatenating the conditions, $b$ is the bias (a constant offset), and $\hat{w}$ specifies a set of weights. The speed profile was advanced by 150 ms before regression to accommodate known lags.

Trial-by-trial analysis
To assess how well projections onto different dimensions predict trial-by-trial movement onset we performed four steps: (1) we chose a potentially informative weighted sum of neurons (“dimension of interest”); (2) we binned and smoothed the spiking data on individual trials; (3) we projected the population neural response from each trial onto the dimension of interest; and (4) for each trial we found the time point at which that projection exceeded a criterion value. That time, relative to the go cue, was the predicted RT. These steps are explained in more detail below.

For the first step, we compared the performance of several different techniques for finding the dimension of interest. Three of these techniques were unsupervised: dimensions were identified based on the structure of the data without exploiting prior knowledge of the RT. These three methods—the CIS$_1$ method, the PC$_1$ method, and the mean-over-all-neurons method—used dPCA, PCA, and simple averaging, respectively. The CIS$_1$ dimension (producing the largest condition-invariant component) and the PC$_1$ dimension (the largest principal component) were found using the long-delay, trial-averaged data (as above). We also employed a linear decoder of reach speed (see above) and a supervised “classifier” method, described below.

For the second step, spikes were counted in 10 ms bins, from 60 ms before to 500 ms after the go cue. Each trial’s spike counts were convolved with a 30 ms Gaussian to produce a smooth spike rate. For the third step, we computed a weighted sum of the neurons’ spike rates. The weights depended on the dimension of interest, found during step one. We refer to the result of this third step as $z(t,r)$, the projection of the neural data as a function of time and trial.

For the final step, we wished to determine when $z(t,r)$ changed in advance of movement onset. To estimate that time, for each trial we asked when $z(t,r)$ first crossed a criterion value derived from the long-delay trials. To find that criterion value, we took the median of $z(t,r)$ across trials, producing $\tilde{Z}(t)$. We set the criterion value to be the midpoint of $\tilde{Z}(t): \frac{\max (\tilde{Z}(t)) + \min (\tilde{Z}(t))}{2}$. The midpoint is an arbitrary but reasonable choice to ensure robustness. For each trial, we found the time at which the criterion value was crossed. Trials that never exceeded the criterion value, or that exceeded it before the go cue, were discarded from the analysis. Such trials were uncommon, especially for the better prediction methods (0–9%, depending on dataset and method).

The three methods described above—the CIS$_1$ method, the PC$_1$ method, and the mean-over-all-neurons method—predict RT in an unsupervised manner. They were compared with a supervised method that was allowed to use knowledge of each trial’s RT. This “classifier” method was based on logistic regression. Single-trial data were first aligned to movement onset, then projected into the dPCA space (including both condition-specific and condition-invariant dimensions). Data were binned into a “premovement” time point (−360 to −150 ms relative to movement onset) and a “movement” time point (−150 to +60 ms relative to movement onset). The dividing point of 150 ms before movement onset was chosen to approximate the delay between when neural firing rates begin to change and when the hand begins to move. Logistic regression returned both a projection dimension and a criterion value that best discriminated between the pre movement and movement data.

As with the other projection methods, the classifier produces a projection vector $\hat{w}$ with as many coefficients as dimensions of the data (in this case, the number of components from dPCA). To characterize the classifier, we asked how much each dPCA component contributed to this projection. Specifically, we took the quantity $|\hat{w}_d|/\sqrt{\text{var}(RD_d)}$, where $|\hat{w}_d|$ is the absolute value of the $d^{th}$ element of $\hat{w}$, $D$ is the dPCA projection matrix (called $W$ in previous equations), $(RD)_d$ indicates the $d^{th}$ column of the matrix resulting from multiplying $RD$, and $\text{var}[]$ indicates taking the variance. This tells us how strongly each of the response components (returned by dPCA) contributed to the final classification.

Finally, we used a semisupervised method where RT was predicted as the time when the decoded reach speed crossed a 50% threshold. Importantly, for all the above methods, training employed only the long-delay data. Trial-by-trial prediction of RT for zero-delay data was entirely based on generalization. Analyses were based on 385/465 trials for dataset JAD1 (long-delay/zero-delay), 249/264 trials for dataset JAD2, 260/427 trials for dataset NAD, and 2982 long-delay trials for dataset NAC.

Finding a rotational plane
For some analyses, we wished to identify planes (two-dimensional projections of the population response) containing rotational structure. We performed dPCA and then applied jPCA (Churchland et al., 2012) to the condition-specific components, using an epoch when neural activity is changing rapidly (−200 to +150 ms relative to movement onset). As a technical detail, the PCA step and mean subtraction were disabled in the jPCA algorithm; dPCA served as a more principled way of focusing jPCA on the strongly condition-specific dimensions. Because both dPCA and jPCA produce linear projections, the final result is also a linear projection of the data.
Results

Behavior and neural recordings

Two monkeys (J and N) performed a variant of the standard delayed-reaching task: the maze task (Fig. 1A,B; Churchland et al., 2010, 2012; Kaufman et al., 2013). The monkey touched and fixated a central spot on a screen, then was presented with a target and, on most trials, a set of virtual barriers (magenta rectangles). After a randomized delay period, a go cue was presented, and the monkey was required to reach to the target, curving around barriers if present. We refer to each target/barrier configuration as a "condition." RTs were brisk: medians of 296 ms (monkey J) and 304 ms (monkey N).

We analyzed six datasets. Three datasets (JAD1, JAD2, and NAD) were collected specifically for this study. For these, recordings were from a single session, made via a pair of 96-electrode arrays, one in PMd and one in M1. To ensure robustness, we also reanalyzed three datasets that have been previously examined. One (NAC) was recorded using a pair of 96-electrode arrays, one (NS) was recorded over many days using single electrodes, and one (JC) combined 1 day of array recordings and many days of single-electrode recordings. These latter two datasets enabled us to analyze large populations that contained both surface PMd/M1 recordings and sulcal M1 recordings.

The firing rate versus time of a representative neuron is illustrated in Figure 1C (for ease of visualization, 4 of 27 conditions are shown). The neuron began responding ~50 ms after target onset, and achieved different firing rates, depending on which reach the monkey was preparing (Tanji and Evarts, 1976; Weinrich et al., 1984; Georgopoulos et al., 1986; Moran and Schwartz, 1999). More generally, the presence of such
neurons is consistent with many prior reports in which some reasonable percentage of neurons were modulated by the task yet not strongly selective for the parameter being tested: e.g., left versus right reaches (Weinrich et al., 1984), three curvatures (Hocherman and Wise, 1991), two or three distances (Riehle et al., 1994; Messier and Kalaska, 2000), or two loads (Evarts, 1968). The present results demonstrate that prior findings were not a trivial result of using few conditions. We employed 27 conditions (108 for dataset NAC) spanning different directions, distances, and reach curvatures, yet still found neurons whose responses were similar across all conditions. Nevertheless, we stress that while individual neurons often showed related structure across conditions—i.e., they were condition-correlated—they essentially never showed fully condition-invariant responses. For example, even the neuron in Figure 2C, which has unusually strong condition-correlated structure, displayed peak firing rates that differed between conditions by a factor of nearly two.

### Population-level structure

Given that single neurons can exhibit condition-correlated responses, some underlying population-level component must be correlated across conditions. To appreciate how this can happen, consider the standard model in which each neuron’s response is a weighted sum of population-level components. The response $r$ of neuron $n$ at time $t$ for condition $c$ is as follows (Eq. 1):

$$r_{n,c,t} = \sum_i w_{n,i} x_{i,c,t}$$

where $x_{i,c,t}$ is the $i^{th}$ response component (one element of the population state $x_c$) and $w_{n,i}$ determines the contribution of component $i$ to the response of neuron $n$. A component is “condition-correlated” if $\text{corr}(x_{i,c,\cdot}, x_{i,c',\cdot})$ is positive when averaged across all choices of conditions $c$ and $c'$.

The possible presence of a condition-correlated component has been considered in many contexts: e.g., decision variables are often modeled as reflecting evidence for a choice (which differs across conditions) plus a growing urgency to make some choice (which is shared across conditions; Churchland et al., 2008; Hanks et al., 2011; Thura et al., 2012; Thura and Cisek, 2014). In the case of reaching, many models include a nondirectional term reflecting hand speed (Georgopoulos et al., 1986; Moran and Schwartz, 1999). Since speed is always positive, and is by definition time-locked to movement onset, a component that reflects speed will be strongly condition-correlated.

In general a condition-correlated component can vary strongly across conditions; the temporal profile must be similar but the amplitude can vary. As a special case, though, such a component may be nearly identical for every condition and thus “condition-invariant.” That is, there might exist an $i^{th}$ component where $x_{i,c,t} = x_{i,c',t}$, for all choices of conditions $c$ and $c'$ and times $t$. This more constrained possibility is suggested by a recent model.

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**Figure 2.** Responses of four example neurons. Format is as in Figure 1C, but responses are shown for all 27 conditions. A, Unit with complex responses. This neuron showed both an overall increase in firing rate across conditions and a strong oscillatory component that was condition-specific (unit JAD1-98, same as in Fig. 1C). Scale bars same as Figure 1C. Inset in upper left shows reach trajectories, colored the same as their corresponding neural traces. B, Another unit with complex condition-specific responses, recorded from the other monkey (unit NAD-165). C, Unit with responses that were strongly condition-correlated (unit JAD1-70). D, Unit where the initial response was condition-correlated: a decline across all conditions. Later activity is more condition-specific (unit JAD1-114).
Sussillo et al., 2015) where the input that triggers movement generation produces population-level components that are close to condition-invariant.

The presence of a condition-invariant component versus a merely condition-correlated component can be determined only at the population level. To do so we applied dPCA (Machens et al., 2010; Brendel et al., 2011), a variant of PCA. Each component identified by dPCA is a pattern of responses across conditions and times (Eq. 1, $x_i ...$) from which the response of each neuron in the population is composed. dPCA exploits knowledge discarded by traditional PCA: the response of a neuron is not simply a vector of firing rates. Rather, each element of that vector is associated with a particular condition and time. dPCA attempts to find components that vary strongly with condition (but not time) or vary strongly with time (but not condition). In practice dPCA never found components of the first type; all components that varied with condition also varied with time. We term these components “condition-specific.” However, dPCA consistently found components that varied with time but not condition (i.e., that were condition-invariant).

Indeed, for every dataset the largest component found by dPCA was close to purely condition-invariant. Figure 3 quantifies the total variance captured by each component (length of each bar) and the proportion of that variance that was condition-invariant (red) versus condition-specific (blue). The largest component (top bar in each panel) exhibited 89–98% condition-invariant variance across datasets.

As a working definition, we term a component “condition-invariant” if $>50\%$ of the variance is condition-invariant. We term a component “condition-specific” if $<50\%$ of the variance is condition-invariant. Empirically...
components were either strongly condition-invariant (much greater than 50% condition-invariant variance) or strongly condition-specific (much less than 50% condition-invariant variance). Each bar plot in Figure 3 thus groups condition-invariant components at top and condition-specific components at bottom. All datasets contained multiple condition-invariant components: respectively two, three, four, four, and four for datasets JAD1, JAD2, JC, NAD, NAC, and NS. For a given dataset, we refer to the set of condition-invariant components as the CIS. We refer to the largest condition-invariant component as CIS1.

Time course of the largest condition-invariant component
CIS1, like all the components, is a linear combination of individual-neuron responses; it is a “protoneural” response strongly reflected in single-neuron PSTHs. The structure of CIS1 can thus be plotted using the format typically used for a single-neuron PSTH. Figure 3 does so for each dataset (colored traces below bar plots).

CIS1 displayed a large and rapid change before movement onset that was similar across conditions. This pattern was present for all datasets. The sudden change occurred ~150 ms before movement onset, corresponding to 50–100 ms before the first change in EMG activity (not shown). The condition-invariance of the signal can be visualized by noting that most individual traces (one per condition) overlap. In particular, during the moments before movement onset, CIS1 increases in a similar way and to a similar degree for every condition. Modest differences between conditions appeared primarily around the end of the movement and during the subsequent hold period (for reference, movement duration was on average 400 ms). Thus, while CIS1 was not identical across conditions, it was very close: on average 94% of its structure was dependent on time but not condition.

The CIS is large
For every dataset, CIS1 captured the most variance of any single component. That is, CIS1 was the component that made the largest contribution to the response structure of individual neurons. More generally, the set of condition-invariant components (CIS; Fig. 3, top grouping of bars within each panel) captured 49–77% of the total variance of datasets JAD1, JAD2, JC, NAD, NAC, and NS. Thus, not only is a CIS present, it typically comprises half or more of the data variance.

While each condition-specific component captured much less variance than CIS1, there were relatively more condition-specific components (Fig. 3, bottom groupings of bars) whose combined variance was 23–51% of the total variance captured by dPCA. These condition-specific components often contained preparatory activity followed by multiphasic responses during the movement. We return later to the structure captured by the condition-specific components.

We did not expect that such a large fraction of the structure in the data—half or more—would be condition-invariant. Most prior work (including our own) has concentrated on the tuned, condition-specific aspects of neural responses. This is reasonable: the presence of a large condition-invariant response component is not obvious at the single-neuron level. Essentially all neurons had contributions from condition-specific components and were therefore tuned for condition. Such tuning is the typical focus of analysis in most studies. Yet the fact that the CIS is so large argues that its properties should also be characterized.

While a few neurons (Fig. 2C,D) had an unusually large contribution from the condition-invariant components, we found no evidence for separate populations of condition-invariant and condition-specific neurons. Weights wni were continuously distributed, and could be positive (Fig. 2C) or negative (Fig. 2D). We also note that the average |wni| was similar for neurons recorded in PMd and M1, indicating that the CIS is of similar size in the two areas.

Assessing demixing
Importantly, dPCA cannot take condition-specific components and render them into condition-invariant components. This is true even if condition-specific components are strongly condition-correlated (see Materials and Methods for mathematical proof; empirical controls described below). Thus, the degree to which the population contains truly condition-invariant components can be assessed by the degree to which dPCA “demixes” responses; that is, the degree to which projecting onto orthogonal dimensions yields some response components that are close to purely condition-invariant. Demixing will be successful only if such condition-invariant structure is present in the data.

As noted above, demixing was successful for all datasets: most components were either strongly condition-invariant or strongly condition-specific. The condition-invariant components (Fig. 3, top grouping of bars in each panel) displayed 75–98% condition-invariant variance (mean, 88%). The condition-specific components (bottom grouping of bars) displayed 74–99% condition-specific variance (mean, 91%). As discussed above, the largest component—CIS1—was always very close to purely condition-invariant (mean, 94%). To put these findings in context, we analyze below a set of model and surrogate populations.

A CIS in a network model
In addition to the six physiological datasets, we analyzed two model populations. The models were recurrent neural networks trained (Sussillo and Abbott, 2009; Martens and Sutskever, 2011) to generate the empirical patterns of muscle activity for two monkeys (Sussillo et al., 2015). Model populations exhibited a CIS (Fig. 4) that closely resembled that of the neural populations. In particular, there was a sudden change in CIS1 shortly before movement onset that was almost purely condition-invariant, with a small amount of condition-specific structure appearing after that transition. Similar to the neural datasets, CIS1 was the largest component of the data and was overall very close (99 and 96%) to purely condition-invariant. As with the physiological data, demixing was successful: the model population response could be sep-
arated into components that were either nearly condition-invariant (Fig. 4, top grouping of bars in each panel) or strongly condition-specific (bottom grouping of bars). The model datasets exhibited, respectively, two and four condition-invariant components—similar to the range of 2–4 seen for the empirical datasets.

As will be shown below, a CIS is not a general feature of any large complex dataset. In the case of the model, the presence of a strong CIS is a consequence of the network inputs (which include a condition-invariant trigger signal) and of the “strategy,” found via optimization, by which the network solves the task. The network was designed such that condition-specific preparatory inputs produce network states (one per condition) appropriate to subsequent movement-period dynamics. Those movement-period dynamics are “turned on” by a strong triggering input that contains no condition-specific information. Because the network was optimized to achieve smooth dynamics, nonlinear interactions are modest, and the triggering input produces a signal that is nearly condition-invariant in the population response. Whether the neural data exhibit a CIS for similar reasons remains unknown, but the temporal structure of the CIS is remarkably similar for the model and for the data.

Controls: comparison of dPCA and PCA
One potential concern is that an algorithm such as dPCA might be able to “successfully” demix any high-dimensional data and find a condition-invariant component. As discussed above (and shown formally in the Materials and Methods), it is not in general mathematically possible to find a condition-invariant component if one is not truly present. Yet in practice, for a finite number of conditions, random smooth data will likely contain some (probably low-variance) signal that may be roughly condition-invariant. Is the empirical CIS larger than expected given this potential concern? Is the CIS found simply because dPCA attempts to find it?

One way to address this concern is to compare the performance of dPCA with that of PCA. PCA identifies dimensions that capture the most data variance possible. If dPCA achieved spurious demixing by finding components with the desired structure but little variance, then dPCA should capture much less variance than PCA. In fact, the dimensions found via dPCA captured almost as much variance as the dimensions found via PCA. Specifically, the set of dPCA dimensions captured 96–99% as much variance as the same number of PCA dimensions. Furthermore, the projections onto the first two PCA dimensions showed a structure that was naturally very close to condition-invariant. This was a simple consequence of the fact that the first few dimensions found by dPCA and PCA were very similar: the first dimension found via PCA formed an angle of only 5° on average with the first dimension found by dPCA. This was true for both the neural and model data. Thus, dPCA simply allows one to gain an ideal view of a condition-invariant structure that is naturally present in the data.

Controls: demixing of real and surrogate data
Despite the above control, one might remain concerned that perhaps any generic data will tend to contain a condition-invariant component that would become apparent when applying dPCA (or PCA). A related potential concern is that a CIS might be found simply because firing rates are constrained to be positive. We addressed these potential concerns by applying dPCA to various surrogate datasets.

First, for each empirical dataset, we replaced each neuron’s response with a random set of responses that was matched with that neuron for frequency content (see Materials and Methods). Across 1000 repetitions for each of the six datasets, dPCA never identified a component with >18% condition-invariant variance. In contrast, the original data contained components with up to 98% condition-invariant variance. This control thus demonstrates that “random” data is very unlikely to yield a strongly condition-invariant component, even when temporal smoothness is matched to that of the empirical data. However, although the randomized responses (not shown) are frequency-matched to the data, they do not form realistic-looking PSTHs because the phases have been randomized (they are essentially just filtered noise). The second and third controls below, in contrast, do result in surrogate responses that look realistic at the level of PSTHs.

For the next control, we produced surrogate datasets by removing the CIS from each real neuron’s response and then applying a firing-rate threshold at zero (see Materials and Methods). The goal was to determine whether it was possible to produce an artifactual CIS by constraining firing rates to be positive. None of these
surrogate populations exhibited a CIS. For example, for the original dataset JAD1, CIS1 contained >90% condition-invariant variance (Fig. 5A,D). The corresponding control dataset (Fig. 5B,E) had no CIS components; all components had <50% condition-invariant variance. For each of the six surrogate datasets, the first component found by dPCA had <21% condition-invariant structure (mean, 6%), in strong contrast to the data where the first component was always strongly condition-invariant. Thus, if a population response does not contain a CIS, a CIS is not created via the constraint that firing rates must be positive.

Finally, we wished to perform a control that could address both of the above concerns while preserving the surface-level features of the original data as closely as possible. To do so, we began with the original neural population (Fig. 5A) and added condition-correlated components (see Materials and Methods). These condition-correlated components had the same temporal profiles as the original condition-invariant components, but the response had a different magnitude for each condition. The surrogate population possessed single-neuron responses (Fig. 5C) that looked remarkably similar to the original responses, and exhibited changes in the average across-condition firing rate that were almost identical to the original responses. Yet the surrogate population lacked any CIS (Fig. 5F). There were no components with >50% condition-invariant variance for any of the surrogate populations, even though these are prominent in all the empirical datasets.

In summary, the presence of a CIS requires a very specific population-level structure and does not arise as a simple consequence of single-neuron response features. Of course, the presence of a CIS is fully consistent with prior work where fits to single-neuron firing rates (e.g., directional tuning curves) typically require a nondirectional component. However, a nondirectional component would also be required when fitting the surrogate responses in Figure 5B,C, which contain no CIS. Thus, the presence of a CIS is consistent with, but not implied by, prior results at the single-neuron level.

**Relationship of the CIS to reach speed and muscle activity**

For the model of Sussillo et al. (2015), the CIS plays an “internal role”: it reflects the arrival of a trigger signal that recruits strong dynamics. Might the CIS in the neural population play a similar internal role? Or might it be more readily explained in terms of external factors: for example, some aspect of kinematics or muscle activity that is invariant across conditions? In particular, tuning for reach speed has been a natural and reasonable way to model nondirectional aspects of single-neuron responses (Moran and Schwartz, 1999). However, for three reasons, the population-level CIS is unlikely to directly reflect reach speed. First, the CIS had a rather different profile from reach speed, which was more sharply phasic (lasting as little as ~200 ms, depending on the condition) and returned to zero as the movement ended (Fig. 6, red trace and blue trace have very different temporal profiles). Second, for the task used here, reach speed is not condition-invariant: it varies considerably (~2×) across the different distances and reach curvatures. Finally, even the small variations present in the CIS across conditions did not parallel variations in reach speed. For monkey J, peak speed and the peak magnitude of CIS1 were not significantly correlated (Fig. 6A,B; overall $r = 0.097$, $p = 0.63$ for JAD1; $r = −0.018$, $p = 0.93$ for JAD2). For monkey N, they were anticorrelated ($r = −0.502$, $p = 0.008$ for NAD, $r = −0.364$, $p < 0.001$ for NAC). Thus, the CIS and reach speed bore little consistent relation. As a side note, the dissimilarity between the CIS and hand speed does not imply that speed information could not be decoded. Using regression, we could identify a dimension that predicted speed fairly well (JAD1: $r = 0.663$; JAD2: $r = 0.743$; NAD: $r = 0.833$; NAC: $r = 0.720$), consistent with prior results that have found strong correlations between neural responses and reach speed (Moran and Schwartz, 1999).

The projection onto this dimension, however, captured much less variance (4–16% as much) than CIS1.

A related possibility is that the CIS might reflect nondirectional aspects of muscle activity. We performed dPCA on EMG recordings made from 9–11 key arm and shoulder muscles. The muscle populations did not exhibit a strong CIS. This can be seen by comparing the first component found via dPCA of the neural data (Fig. 7A,B) with the first component found via dPCA of the muscle data (Fig. 7C,D). The former is nearly condition-invariant while the latter is not. For each component found via dPCA, we measured the fraction of variance that was condition-invariant (the “purity” of condition-invariance) and the variance accounted for relative to the condition-specific components (the “strength” of that component). Unlike the neural populations (Fig. 7E, green) the muscle populations (purple) did not contain condition-invariant components that were both relatively pure and reasonably strong; there are no purple symbols in the upper right corner. Certainly the muscle population response contained some nondirectional aspects: there existed components in which there was an overall change that was mostly of the same sign across all conditions, resulting in a proportion of condition-invariant variance as high as 0.5–0.75 (purple symbols, left). This variance is not negligible, as evidenced by the fact that it could be further reduced via the control manipulations that were applied to the neural population in Figure 5 (muscle version not shown). However, the components in question captured relatively modest amounts of variance, and were not nearly as pure as the components found for the neural populations. Thus, the presence of condition-invariant structure in the neural population cannot be secondary to features of the muscle activity; only the neural population contained components that were both close to purely condition-invariant and captured a large percentage of the overall variance.

The muscle responses further underscore that the presence or absence of a CIS cannot be inferred from surface-level features. Individual muscle responses closely resembled neural responses in many ways, and often showed overall rises in activity across conditions.
Figure 5. Demixing performance for one empirical dataset and two surrogate control datasets. 

A, PSTHs of three example units from dataset JAD1. Scale bars: horizontal, 200 ms; vertical, 10 spikes/s. 

B, PSTHs for a surrogate dataset where we projected onto the condition-specific dimensions, then rectified so that all firing rates remained positive (see Materials and Methods). This surrogate dataset explores the possibility that a CIS might appear merely due to firing rates being constrained to be positive. The three PSTHs correspond to the same units shown in A, after modification. 

C, PSTHs for a surrogate dataset where we added condition-correlated components. The condition-correlated components had the same temporal profile as the projections onto the condition-invariant dimensions found by dPCA but had a different amplitude for each condition (see Materials and Methods). This surrogate dataset explores whether a condition-correlated structure at the single-neuron level is sufficient to yield condition-invariant components at the population level. The three PSTHs correspond to the same units shown in A, after modification. 

D–F. Quantification of the CIS as in Figure 3. Panels correspond to examples above. D is reproduced from Figure 3A for comparison.
Thus, fits to muscle activity would benefit from a nondirectional component just as do fits to neural activity. Yet as a population, the muscles showed only condition-correlated structure, and had little or no CIS.

**Trial-by-trial prediction of RT**

In all datasets, the sudden change in the CIS occurred ~150 ms after the go cue and ~150 ms before the onset of physical movement (50-100 ms before muscle activity began to change). The change in the CIS might thus be a visuospatial response locked to the go cue, consistent with the presence of other visuospatial signals in the premotor cortex (Crammond and Kalaska, 1994; Shen and Alexander, 1997). Alternatively, the change in the CIS could be locked to the transition from preparation to movement, consistent with the model of Sussillo et al. (2015). These two possibilities can be distinguished at the single-trial level. If the CIS reflects the visual go cue, it would have no ability to predict the subsequent variable RT between the go cue and movement onset. If the CIS reflects an internal transition related to movement onset, the CIS should be strongly predictive of RT.

We were able to address the trial-by-trial timing of the CIS in three datasets (JAD1, JAD2, and NAD) that were collected specifically for this purpose. These datasets involved simultaneous recordings (116–213 units) from two chronically implanted 96-electrode arrays, allowing single-trial estimates of the CIS. Critically, for these datasets we employed a task structure that allowed examination of trial-by-trial RT variability independent of delay-period duration. Over the course of training and most experiments, monkeys experienced a continuous distribution of delay-period durations from 0 to 1000 ms. It is well known that delay-period duration has an impact on RT (Rosenbaum, 1980; Riehle and Requin, 1989; Churchland et al., 2006a). To study RT variability independent of such effects, for these three datasets we interleaved additional trials with a discrete set of delay durations: 0, 100, 200, and 500 ms (see Materials and Methods). This allowed us to examine the relationship between neural and RT variability for sets of trials with a matched delay. Below we present data for trials with zero delay and trials with a “long” (500 ms) delay. Results were very similar when we analyzed the sets of trials with 100 and 200 ms delays. For comparison, we repeated these analyses of RT for dataset NAC (which did not contain discrete delays) using all trials with delays >150 ms. All results were very similar across all four datasets.

CIS$_1$ was readily resolved on individual trials (Fig. 8 shows data for JAD1 with analyses repeated in Fig. 9 for NAD). The neural weights defining CIS$_1$ were found using data from the long-delay trials. Example single-trial projections of the long-delay data are shown in Figures 8B and 9B. These same weights successfully generalized

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**Figure 6.** Comparison of the CIS and hand speed. Hand speed (blue) and the first component of the CIS (red) are shown for four reach conditions. For hand speed, light traces show all trials; heavy trace shows mean over trials. CIS$_1$ is the mean over trials. Insets show the maze for that condition and a prototypical reach path. A, A straight reach with a fast speed profile. Maze ID25. B, A straight reach with a slow speed profile. Maze ID7. C, A curved reach with a long speed profile. Maze IDS. D, A curved reach with an unusual speed profile. Maze ID14. The CIS was similar across all four examples, despite differences in the speed profile. Dataset NAD.
and revealed an essentially identical CIS1 for the zero-delay trials (Figs. 8A, 9A). The latency of the rise time of CIS1, relative to the go cue, varied from trial to trial. To estimate this latency, we measured when CIS1 crossed a criterion value following the go cue (Figs. 8A, B, 9A, B, gray line). We selected a 50% criterion that is simply a practical and robust criterion for estimating rise time (and should not be interpreted as suggesting a physiological threshold). The estimated rise time strongly predicted the subsequent RT on individual trials (Figs. 8C, 9C) for both long-delay (blue) and zero-delay (red) trials. This was true across all analyzed datasets: the average correlation was $r = 0.805$ for long-delay trials, and $r = 0.827$ for zero-delay trials.

The CIS strongly predicts RT on a single-trial basis, but does it do so more accurately than other reasonable...
methods? The projection of the data onto the first principal component of the data (PC1) predicted RT almost as well as did CIS1. This was especially true for monkey J (Fig. 8D) and somewhat less so for monkey N (Fig. 9D) due to a tendency for the projection onto PC1 to occasionally exceed the criterion early. Given the ability of CIS1 to predict RT, the similar success of the projection onto PC1 is unsurprising: as discussed above, the dimensions con-

**Figure 8.** Predicting RT using projections of the data for dataset JAD1. **A**, Each trace plots CIS1 over time on a single zero-delay trial. Fifty trials selected randomly at intervals throughout the day are shown. Black trace plots the median across all trials. **B**, Same as in **A** but for trials with a 500 ms delay period (long delay). The criterion value (gray line) was chosen using long-delay trials. The same value was used for zero-delay trials (**A**). **C–G**, Correlation of behavioral RT with the time when the neural criterion value was crossed. For each panel, data are shown for both long-delay trials (blue) and zero-delay trials (red). Lines show linear regressions; dashed lines show 95% confidence bounds of the fit. Each panel in **C–G** gives the correlation for a different linear projection of the population response: CIS1 (**C**), the projection onto the first PC (**D**), the mean over all neurons (**E**), the projection onto the dimension that best reconstructed speed according to a linear regression (**F**), and the projection onto the axis found by a logistic regression classifier (**G**). Trials where the neural data did not cross the criterion value were excluded. **H**, Coefficient influence for the classifier. Coefficients for condition-invariant dimensions shown in red; condition-specific dimensions shown in black.
taining PC₁ and CIS₁ were closely aligned. Nonetheless, CIS₁ always predicted RT at least slightly better than the projection onto PC₁, despite PC₁ capturing (by construction) slightly more variance. The average firing rate across all neurons (Figs. 8E, 9E) predicted RT less well than did CIS₁ or the projection onto PC₁. Finally, because RT was quantified based on measured hand speed, we considered the projection that best decoded hand speed (found via regression; see above). Decoded hand speed performed acceptably, but noticeably less well than CIS₁ (Figs. 8F, 9F; across all analyzed datasets, mean r = 0.666 for long delay, r = 0.674 for zero delay). Thus, CIS₁ predicted RT better than did other reasonable unsupervised and semisupervised methods.

Might there exist another signal in the data that could considerably outperform CIS₁? To address this, we trained a classifier based on logistic regression (see Materials and Methods) to distinguish neural data recorded before versus after the sudden transition in neural activity 150 ms before movement onset. The classifier—which

Figure 9. Predicting RT using projections of the data for dataset NAD. Same format as Figure 8. Regarding H, note that for this dataset there were four CIS components.
has the advantage of being optimized using knowledge of RT—predicted RT for zero-delay trials slightly better than CIS, for one dataset (Fig. 8G) and slightly worse for the other (Fig. 9G; note that when assessing generalization, a supervised method is not guaranteed to outperform an unsupervised method). We then asked which dimensions the classifier relied upon. The coefficients of the classifier (Figs. 8H, 9H) revealed that the condition-invariant dimensions (red) were used more strongly than the condition-specific dimensions (black); 74% of the classifier was based on the CIS (79% for dataset NAD). Thus, the CIS is a particularly good predictor of RT, and it is difficult to improve on the performance it provides. Results were similar for the other two datasets (for dataset JAD2: 69% of classifier based on CIS; dataset NAC: 82% of classifier based on CIS). Thus, the timing of the CIS reflects the pending onset of movement, rather than the arrival of a visual signal. Had the latter been true, the CIS would have had no ability to predict RT when data are time-locked to the go cue as they were here.

**Neural and model population trajectories**

We recently reported that the population response exhibits a strong ~2 Hz oscillatory component during movement, manifested as a rotation of the neural state (Churchland et al., 2012; Churchland and Cunningham, 2014). This oscillatory component is condition-specific: rotation amplitude and phase differ across reach directions, curvatures, speeds, and distances. As expected given these prior results, we found that the eight-dimensional condition-specific space identified via dPCA contained components with a strong rotational structure. This conveniently allows the population structure to be plotted as a neural trajectory in a state space, with one dimension capturing CIs, and two dimensions capturing the plane with the strongest rotations. The resulting three-dimensional projections captured 47 and 45% (for datasets JAD1 and NAD respectively) of the total variance captured by dPCA. The three-dimensional structure is best viewed in video format (Movies 1–4) but can also be appreciated via inspection of a set of two-dimensional projections (Fig. 10A,C).

Each trace in Figure 10 and Movies 1–4 plots the neural trajectory for one condition. Traces are colored gray during baseline, blue during the delay period, then shaded from red to green across conditions (to aid visualization) during a “perimovement period”: ~200 to +150 ms relative to movement onset. The overall structure carved out by the trajectories is roughly conical; neural activity is at the narrow end of the cone during the delay period, translates along the long axis of the cone just before movement onset, then exhibits rotations at the wide end of the cone during movement. Rotations begin with (or just at the end of) the translation and continue after the translation is over, tracing out a rough disk. The top row plots projections in which the cone is seen end-on. Middle rows plot projections in which the cone is seen from the side (the rotational disk being viewed from the edge) and the bottom row plots a projection that illustrates (as best as possible in two dimensions) the full three-dimensional structure.

Consistent with many reports demonstrating the existence of preparatory activity (Tanji and Evarts, 1976; Weinrich and Wise, 1982; Hocherman and Wise, 1991; Messier and Kalaska, 2000; Churchland et al., 2006b), condition specificity first develops during the delay period. For example, in the third row, blue traces spread out over a larger range of states than do gray traces. The subsequent rotations are also condition-specific. The CIS produces the long axis of the cone: a large translation of the neural state that is similar for every condition. This translation is almost perfectly orthogonal to the rotations. Such orthogonality is not a consequence of the analysis method: the axes are orthogonal by construction, but that in no way constrains the condition-invariant and condition-specific structure to be orthogonal. Indeed, demixing (Fig. 3) is successful precisely because the condition-invariant and condition-specific response structure is orthogonal, as revealed directly in Figure 10. The other four datasets showed the same structure.

A striking feature of the response structure is that condition-specific preparatory activity occurs in one region of state space, while condition-specific rotational structure during movement occurs in a different region of state space. Given the above results showing that the CIS predicts RT, a natural question is whether the transition from one region to another relates to the behavioral transition from preparing to move (while holding a steady posture) to actually moving. This is indeed how the network model of Sussillo et al. (2015) functions. Through optimization, that model adopted a strategy where an incoming “trigger signal” produced a large translation, bringing the population state near a fixed point where local dynamics were rotational and produced the multiphasic patterns of muscle activity. That study noted the general similarity between neural and model data, as revealed via canonical correlation analysis, and the presence of a change in the overall mean firing rate. That overall change is a natural product of the CIS, which as documented above is present in both neural (Fig. 3) and model (Fig. 4) populations.

To further compare, we projected the model population response (Fig. 10B,D) as we had the neural population response. Model and neural populations exhibited remarkably similar structure when viewed from all angles. Preparatory activity developed in one region of space, and the CIS then caused an overall translation to another region of space. The rotations of the neural state (at a little less than 2 Hz) began during that translation and continued to unfold after the translation was complete.

**Relative timing of the CIS and rotations**

The above results suggest that the CIS may relate to the transition from relatively stable preparatory dynamics to strongly rotational movement-period dynamics. This hypothesis makes a specific prediction: the CIS should begin to change with, or perhaps shortly before, the onset of rotational dynamics. The hypothesis would be falsified if the CIS began changing after rotations had already begun.
begun, or if the CIS began changing long before rotations began. To assess relative timing, we computed the “speed” of the neural trajectory: the rate of change of the neural state. This was done separately for the CIS dimensions and the two dimensions with the strongest rotations (Fig. 11). In all cases, for both the model and data, the peak speed in the CIS dimensions (red) slightly leads the peak speed in the rotational dimensions (blue). Thus, both the neural and model data showed the predicted effect.

Discussion
We found that the largest component of the population response in M1/PMd is consistently condition-invariant: it changes in an almost identical fashion regardless of reach direction, curvature, and distance. More generally, a small set of condition-invariant components (the CIS) contained half or more of the population-level variance. Thus, although essentially all individual motor cortex neurons are “tuned,” the population response is dominated by the CIS. This result could not be inferred from, but is consistent with, three prior findings. First, single neurons often exhibit an overall change in firing rate during movement (e.g., with most conditions showing an increase in rate, or most conditions showing a decrease in rate; Fortier et al., 1993; Crammond and Kalaska, 2000). Second, a strong nondirectional ensemble response is present in motor cortex (Moran and Schwartz, 1999; Churchland and Shenoy, 2007) such that fits are greatly aided by a nondirectional term (Georgopoulos et al., 1986; Moran and

Figure 10. Various projections that capture CIS1 and rotations of the neural state during movement. Data were first projected onto three dimensions: the dimension that yielded CIS1 and two condition-specific dimensions that captured strong rotational structure (see Materials and Methods). Each panel plots a different view of the data projected onto those three dimensions. A, Four different views of the three-dimensional projection for dataset JAD1. “Baseline” activity (before target onset) plotted in gray, preparatory activity plotted in blue, and activity after the go cue plotted in green and red (colors chosen arbitrarily for each condition). B, Same for the neural network model trained to produce EMG recorded from monkey J. C, Same for dataset NAD. D, Same for the neural network model trained to produce EMG recorded from monkey N.
Schwartz, 1999). Third, population summaries often show a rise in activity for both the “preferred” and “anti-preferred” direction around the time of the movement (Bastian et al., 2003). Yet importantly, the presence of the CIS could not be directly inferred from the above findings; they are all equally consistent with structure that is condition-correlated but far from condition-invariant. For example, the surrogate data in Figure 5 show all three of the above features yet lack any condition-invariant component. In summary, the current data and analyses reveal something that could not be inferred previously: the data contain condition-invariant components that constitute a very large percentage of the overall structure of the neural responses.

**Temporal properties of the CIS**

Although one might initially be tempted to view untuned response aspects as “nonspecific,” the CIS exhibits specific temporal structure. For all six neural datasets and both model datasets, there is a sudden change in the CIS ~150 ms before movement begins. The sudden change can be visualized on individual trials and is strongly predictive of trial-by-trial RT. This strong relationship reflects the fact that the CIS is tied to movement onset (rather than the appearance of the go cue) and is large enough to be readily measured on single trials. The CIS also has a specific population-level structure that was consistent across datasets: the CIS is manifested as a large translation of the neural state from one region of neural state space (occupied when the monkey is preparing the movement) to another region (occupied just before and during overt movement).

While neural responses are often interpreted in terms of their tuning for external factors, the CIS did not relate to any external factor we examined. The temporal profile of the CIS did not resemble that of hand speed, nor were
condition-to-condition variations in hand speed paralleled by the (very small) condition-to-condition variations in the CIS. This is consistent with the noisiness associated with decoding pure hand speed in neural prosthetics (Golub et al., 2014), and suggests that the CIS could be useful for applications seeking to decode a rest versus a move signal (Velliste et al., 2014). The CIS also did not relate to any measurable aspect of muscle activity. Although muscles often exhibited overall changes in activity that were correlated across conditions, the muscle population exhibited little to no CIS. This again underscores that condition-correlated structure typically does not imply a CIS. Finally, the CIS did not simply reflect the visual arrival of the go cue. As indicated by the ability to predict RT, the CIS was instead related to the time of movement onset. Furthermore, the sudden change in the CIS occurred well after (~150 ms) the visual go cue. This contrasts with the very rapid (~60 ms latency) response of neurons in M1 and PMd to the onset of the target (Ames et al., 2014; Fig. 2A,B,D). We also note that the visual go cue was far from condition-invariant: it involved salient changes in the appearance of the target(s), which had different visual locations across conditions. Thus, a natural interpretation is that the CIS relates to the go cue only indirectly, and reflects an internal transition from preparation to movement that follows the go cue with a long and variable latency. Still, we cannot rule out that the CIS is a long- and variable-latency visual response to the go cue, and that the reaction time inherits this variability. Addressing this possibility will require future experiments in which there is no sensory go cue.

Future experiments will also be required to address whether the timing of the CIS relates in any way to the last moment when movement can be suppressed. A recent hypothesis is that RTs are artificially long not because motor preparation is slow, but because “triggering” is conservative (Haith et al., 2016), leaving time for the movement to be altered or suppressed (Riehle et al., 2006; Scangos and Stuphorn, 2010; Mirabella et al., 2011). The relatively long ~150 ms time between the go cue and the sudden change in the CIS, relative to the ~60 ms latency of the first “preparatory” response, is consistent with this hypothesis.

An internal role for the CIS?

The properties of the CIS suggest that it likely relates not to a representation of external factors, but to some internal process—perhaps the transition from preparatory neural dynamics to movement-related neural dynamics. It is becoming increasingly appreciated that many motor cortex signals may not relate cleanly to external parameters, and are more naturally explained in terms of their internal roles in computation (Reimer and Hatsopoulos, 2009; Chase and Schwartz, 2011; Shenoy et al., 2013; Churchland and Cunningham, 2014). The hypothesis that the CIS might relate to the transition from preparation to
movement is further suggested by the finding that the network model of Sussillo et al. (2015) exhibits a very similar CIS—and similar overall population structure—to the neural data (Figs. 4, 10). In the case of the model, the CIS is a consequence of the externally delivered trigger signal, and is in turn the cause of the change in neural dynamics that generates movement. The original analyses in Sussillo et al. did not focus on or attempt to isolate a CIS. Yet a condition-invariant translation is clearly present in one key analysis (Sussillo et al., 2015, their Fig. 6) and can be seen to bring the set of network states close to a fixed point with rotational dynamics. Whether this interpretation is also correct for the data is of course still uncertain, but the population response structure is remarkably similar for the model and data. This interpretation is also supported by both the overall timing of the CIS (it occurs just as, or even slightly before, the onset of rotational dynamics; Fig. 11) and the remarkably strong correlation between the change in the CIS and the moment when movement begins (Figs. 8, 9).

Other, not necessarily exclusive explanations are also likely. For example, the CIS could activate, suppress, or alter how the local circuit processes feedback (Cluff et al., 2015). Similarly, the CIS could relate to an overall modulation of downstream reflexes or to a disengagement of postural control (Kurtzer et al., 2005; Cluff and Scott, 2016). After all, the initiation of activity that drives movement must presumably be accompanied by cessation of the activity that held the hand in place during the delay period. This is true even of the model of Sussillo et al., which is involved in a rudimentary form of postural control during the delay period: producing a constant pattern of muscle activity. For that model, the CIS produces the transition away from the stable dynamics that maintain constant outputs, and towards oscillatory dynamics that produce the movement-driving patterns of muscle activity.

What inputs might produce a CIS?
If motor cortex undergoes a large condition-invariant change prior to movement, what drives that change? What other area(s) might supply the relevant input? A number of candidate regions exist, including the basal ganglia (Romo et al., 1992; Hauber, 1998), superior colliculus (Werner, 1993; Philipp and Hoffmann, 2014), parietal cortex (Scherberger and Andersen, 2007; Pesaran et al., 2008), supplementary motor area (Orgogozo and Larsen, 1979; Eccles, 1982; Romero and Schultz, 1992), the dentate nucleus of the cerebellum (Meyer-Lohmann et al., 1977), and, in rodents, secondary motor cortex (Murakami et al., 2014). Moreover, the origin of the CIS may depend on the task: movements elicited by a strong sensory cue may be generated differently from self-initiated movements (Kurata and Wise, 1988) or movements that must be made very rapidly (Perfiliev et al., 2010). Along similar lines, RTs can be remarkably short when the go cue is provided by a mechanical perturbation of the limb (Evarts and Tanji, 1976; Pruszynski et al., 2008). These short RTs may be related to the finding that some neurons show a rapid perturbation-driven response that is invariant across perturbation directions (Herter et al., 2009, their Fig. 3C). It is thus vital that future studies address whether a similar CIS is present in motor cortex across the many possible sensory cues and internal events that can be responsible for causing movement initiation.

Summary
In summary, our results build upon the long-standing observation that responses are often correlated across conditions at the single-neuron level. Our results reveal that this general surface-level structure reflects a very particular kind of underlying structure: a large condition-invariant response component with timing closely tied to movement onset. This adds to a small but growing list of “untuned” response aspects that might initially appear incidental, but may in fact play important computational roles.

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