Bridging neuronal correlations and dimensionality reduction

Graphical abstract

Highlights

- Pairwise correlation and dimensionality reduction characterize how neurons covary
- Pairwise and population metrics are closely related mathematically and empirically
- Decrease in V4 pairwise correlation corresponds to multiple population-level changes
- Multiple activity statistics should be used when describing population covariability

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In brief

Pairwise correlations and dimensionality reduction are widely used approaches for measuring how neurons covary. Umakantha, Morina, Cowley, et al. establish concrete mathematical relationships between the two approaches and empirically investigate these relationships for visual cortical neurons. The findings provide a cautionary tale for summarizing population-wide covariability using any single activity statistic.

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Article

Bridging neuronal correlations and dimensionality reduction

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SUMMARY

Two commonly used approaches to study interactions among neurons are spike count correlation, which describes pairs of neurons, and dimensionality reduction, applied to a population of neurons. Although both approaches have been used to study trial-to-trial neuronal variability correlated among neurons, they are often used in isolation and have not been directly related. We first established concrete mathematical and empirical relationships between pairwise correlation and metrics of population-wide covariability based on dimensionality reduction. Applying these insights to macaque V4 population recordings, we found that the previously reported decrease in mean pairwise correlation associated with attention stemmed from three distinct changes in population-wide covariability. Overall, our work builds the intuition and formalism to bridge between pairwise correlation and population-wide covariability and presents a cautionary tale about the inferences one can make about population activity by using a single statistic, whether it be mean pairwise correlation or dimensionality.

INTRODUCTION

A neuron can respond differently to repeated presentations of the same stimulus. These variable responses are often correlated across pairs of neurons from trial to trial, measured using spike count correlations ($r_{sc}$, also referred to as noise correlation; Cohen and Kohn, 2011). Studies have reported changes in spike count correlation across various experimental manipulations and cognitive phenomena, including attention (Cohen and Maunsell, 2009; Mitchell et al., 2009; Herrero et al., 2013; Gregoriou et al., 2014; Ruff and Cohen, 2014a; Snyder et al., 2018), learning (Gu et al., 2011; Jeanne et al., 2013; Ni et al., 2018), task difficulty (Ruff and Cohen, 2014b), locomotion (Erisken et al., 2014), stimulus drive (Maynard et al., 1999; Kohn and Smith, 2005; Smith and Kohn, 2008; Miura et al., 2012; Ponce-Alvarez et al., 2013; Ruff and Cohen, 2016b), decision making (Nienborg et al., 2012), task context (Bondy et al., 2018), anesthesia (Ecker et al., 2014), adaptation (Adibi et al., 2013), and more (Figure 1A). Spike count correlation also depends on timescales of activity (Sair et al., 2001; Kohn and Smith, 2005; Smith and Kohn, 2008; Mitchell et al., 2009; Runyan et al., 2017), neuromodulation (Herrero et al., 2013; Minces et al., 2017), and properties of the neurons themselves, including their physical distance from one another (Lee et al., 1998; Smith and Kohn, 2008; Smith and Sommer, 2013; Ecker et al., 2014; Solomon et al., 2015; Rosenbaum et al., 2017), tuning preferences (Lee et al., 1998; Romo et al., 2003; Kohn and Smith, 2005; Huang and Lisberger, 2009), and neuron type (Qi and Constantinidis, 2012; Snyder et al., 2016). Theoretical work has posited that changes in correlations affect neuronal computations and sensory information coding (Zohary et al., 1994; Shadlen and Newsome, 1998; Abbott and Dayan, 1999; Averbeck et al., 2006; Moreno-Bote et al., 2014; Sharp and Berkes, 2019; Rumyantsev et al., 2020; Bartolo et al., 2020). Given such widespread empirical observations and theoretical insight, spike count correlation has been and remains instrumental in our current understanding of how neurons interact.

Most studies compute the average spike count correlation over pairs of recorded neurons for different experimental
conditions, periods of time, neuron types, etc. A decrease in this mean correlation is commonly attributed to a reduction in the size (or gain) of shared co-fluctuations (Shadlen and Newsome, 1998; Rabinowitz et al., 2015; Lin et al., 2015; Eckert et al., 2016; Huang et al., 2019; Ruff et al., 2020), e.g., a decrease in the strength of “common shared input” that drives each neuron in the population. However, other distinct changes at the level of the entire neuronal population can manifest as the same decrease in mean pairwise correlation (Figure 1B). For example, a common input that drives the activity of all neurons up and down together could be altered to drive some neurons up and other neurons down. Alternatively, that first common input signal might remain the same, but a second input signal could be introduced that drives some neurons up and others down. It is difficult to differentiate these distinct possibilities using a single summary statistic, such as mean spike count correlation.

Distinguishing among these changes to the population-wide covariability might be possible by considering additional statistics that measure how the entire population of neurons co-fluctuates together. In particular, one may use dimensionality reduction to compute statistics that characterize multiple distinct features of population-wide covariability (Cunningham and Yu, 2014). Dimensionality reduction has been used to investigate decision making (Harvey et al., 2012; Mante et al., 2013; Kiani et al., 2014; Kaufman et al., 2015), motor control (Churchland et al., 2012; Gallego et al., 2017), learning (Sadler et al., 2014; Ni et al., 2018; Vyas et al., 2018), sensory coding (Mazor and Laurent, 2005; Pang et al., 2016), spatial attention (Cohen and Maunsell, 2010; Rabinowitz et al., 2015; Snyder et al., 2018; Huang et al., 2019), interactions between brain areas (Perich et al., 2018; Ruff and Cohen, 2019a; Ames and Churchland, 2019; Semedo et al., 2019; Veuthey et al., 2020), and network models (Williamson et al., 2016; Mazzucato et al., 2016; Recanatesi et al., 2019), among others. As with mean spike count correlation, the statistics computed from dimensionality reduction can also change with attention (Rabinowitz et al., 2015; Huang et al., 2019), stimulus drive (Churchland et al., 2010; Cowley et al., 2016; Snyder et al., 2018), motor output (Gallego et al., 2018), learning (Athayre et al., 2017), and anesthesia (Ecker et al., 2014). However, unlike mean spike count correlation (henceforth referred to as a “pairwise metric”), which averages across pairs of neurons, the statistics computed from dimensionality reduction (henceforth referred to as “population metrics”) consider the structure of population-wide covariability (Figure 1C). Although dimensionality reduction is often applied to trial-averaged activity (removing trial-to-trial variability), here, we focus on using dimensionality reduction to study trial-to-trial variability (around the trial-averaged mean). An example of a commonly reported population metric is dimensionality (Yu et al., 2009; Rabinowitz et al., 2015; Cowley et al., 2016; Williamson et al., 2016; Mazzucato et al., 2016; Gao and Ganguli, 2015; Gallego et al., 2017; Stringer et al., 2019a; Recanatesi et al., 2019). Dimensionality is used to assess whether the number of population co-fluctuation patterns (possibly reflecting the number of common inputs) changes across experimental conditions (Figure 1B, condition 1 versus condition 2, right panel). Thus, population metrics could help to distinguish among the distinct ways in which population-wide covariability can change, especially those that lead to the same change in mean spike count correlation (Figure 1B).

Both pairwise and population metrics aim to characterize how neurons covary, and both can be computed from the same spike count covariance matrix (Figure 1C). Still, studies rarely report both, and the relationship between the two is not known. In this study, we establish the relationship between pairwise metrics and population metrics both analytically and empirically using simulations. We find that changes in mean spike count correlation could correspond to several distinct changes in population metrics, including (1) the strength of shared variability (e.g., the strength of a common input), (2) whether neurons co-fluctuate together or in opposition (e.g., how similarly a common input drives each neuron in the population), or (3) the dimensionality (e.g., the number of common inputs). Furthermore, we show that a rarely reported statistic—the standard deviation of spike count correlation—provides complementary information to the mean spike count correlation about how a population of neurons co-fluctuates. Applying this understanding to recordings in area V4 of macaque visual cortex, we found that the previously reported decrease in mean spike count correlation with attention stems from multiple distinct changes in population-wide covariability. Overall, our results demonstrate that common ground exists between the literatures of spike count correlation and dimensionality reduction and provides a cautionary tale for attempting to draw conclusions about how a population of neurons covaries using one, or a small number of, statistics. Our framework builds the intuition and formalism to navigate between the two approaches, allowing for a more interpretable and richer description of the interactions among neurons.

RESULTS

Defining pairwise and population metrics

We first define the metrics that we will use to summarize (1) the distribution of spike count correlations (i.e., pairwise metrics) and (2) dimensionality reduction of a population covariance matrix (i.e., population metrics). For pairwise metrics, we consider the mean and standard deviation (SD) of $r_{sc}$ across all pairs of neurons, which summarize the $r_{sc}$ distribution (Figure 1C, bottom left panel). For population metrics, which are derived from factor analysis (FA), we consider loading similarity, percent shared variance (abbreviated to %sv), and dimensionality (described below and in more detail in STAR Methods). These metrics each describe some aspect of population-wide covariability and thus represent natural, multivariate extensions of $r_{sc}$.

To illustrate these three population metrics, consider the activity of a population of neurons over time (Figure 2A, spike rasters). If the activity of all neurons goes up and down together, we would find the pairwise spike count correlations between all pairs of neurons to be positive. A more succinct way to characterize this population activity is to identify a single time-varying latent co-fluctuation that is shared by all neurons (Figure 2A, blue line). The way in which neurons are coupled to this latent co-fluctuation is indicated by a loading for each neuron. In this example, because the latent co-fluctuation describes each neuron’s activity going up and down together, the loadings have the same sign (Figure 2A, green rectangles). We refer to the latent co-fluctuation’s corresponding
set of loadings as a co-fluctuation pattern. A co-fluctuation pattern can be represented as a direction in the population activity space, where each coordinate axis corresponds to the activity of one neuron (Figure 2A, right panel).

The first population metric is loading similarity, a value between 0 and 1 that describes to what extent the loadings differ across neurons within a co-fluctuation pattern. A loading similarity close to 1 indicates that the loadings...
have the same sign and are of similar magnitude (Figure 2A, green rectangles). A loading similarity close to 0 indicates that many of the loadings differ, either in magnitude, sign, or both (Figure 2B, green and pink squares). In this case, some neurons may have positive loadings and co-fluctuate in the same direction as the latent co-fluctuation (Figure 2B, top rows of neurons show high firing rates when blue line is high and low firing rates when blue line is low), whereas other neurons may have negative loadings and co-fluctuate in the opposite direction as the latent co-fluctuation (Figure 2B, bottom rows of neurons show low firing rates when blue line is high and high firing rates when blue line is low). One can view changing the loading similarity as rotating the direction of a co-fluctuation pattern in population activity space (Figure 2B, bottom plot).

The second population metric is percent shared variance or %sv, which measures the percentage of spike count variance explained by the latent co-fluctuation. This percentage is computed per neuron and then averaged across all neurons in the population (Williamson et al., 2016). A %sv close to 100% indicates that the activity of each neuron is tightly coupled to the latent co-fluctuation, with a small portion of variance that is independent to each neuron (Figure 2A). A %sv close to 0% indicates that neurons fluctuate almost independently of each other and their activity weakly adheres to the time course of the latent co-fluctuation (Figure 2C). By changing %sv, one does not change the co-fluctuation pattern in population activity space (Figure 2, blue lines are the same in panels A and C) but rather the strength of the latent co-fluctuation (Figure 2C, blue line has smaller amplitude than in panel A).

Figure 2. Intuition about population metrics: loading similarity, percent shared variance (%sv), and dimensionality

(A) Population activity (where each row is the spike train for one neuron over time; simulated data) is characterized by a latent co-fluctuation (blue) and a co-fluctuation pattern made up of loadings (green rectangles). Each neuron’s time-varying firing rate is a product of the latent co-fluctuation and that neuron’s loading (which may either be positive or negative). One may also view population activity through the lens of the population activity space (right plot), where each axis represents the activity of one neuron (n1, n2, n3 represent neuron 1, neuron 2, and neuron 3). In this space, a co-fluctuation pattern corresponds to an axis whose orientation depends on the pattern’s loadings (right plot, blue line).

(B) Population activity with a lower loading similarity than in (A). The loadings have both positive and negative values (i.e., dissimilar loadings), leading to neurons that are anti-correlated (compare top rows with bottom rows of population activity). Changing the loading similarity will rotate a pattern’s axis in the population activity space (bottom plot, “rotate axis”).

(C) Population activity with a lower %sv than in (A). The latent co-fluctuation shows smaller amplitude changes over time than in (A), which leads to a lower %sv. Changing %sv leads to no changes of the co-fluctuation pattern (bottom plot, axis is same as that in A).

(D) Population activity with a dimensionality of 2, compared to a dimensionality of 1 in (A). Adding a new dimension leads to a new latent co-fluctuation (orange line) and a new co-fluctuation pattern (“added new pattern”). Each neuron’s time-varying firing rate is expressed as a weighted combination of the latent co-fluctuations, where the weights correspond to the neuron’s loadings in each co-fluctuation pattern. Here, each dimension corresponds to a distinct subset of neurons (top rows versus bottom rows); in general, this need not be the case, as each neuron typically has non-zero weights for both dimensions. In the population activity space (bottom plot), the activity varies along the two axes (i.e., a 2D plane) defined by the two co-fluctuation patterns. See also Figure S5. The spike trains shown in this figure were created for the sole purpose of illustrating the population metrics in this figure and were not used in subsequent analyses. The spike trains were generated by first creating latent co-fluctuations using Gaussian processes. These latent co-fluctuations were then linearly combined using loading weights (drawn from a standard normal distribution), yielding a time-varying firing rate for each neuron. Spike trains were generated according to an inhomogeneous Bernoulli process based on the time-varying firing rates. The intended duration of each spike train plotted is around 10 s.
Figure 3. Relationship between population metrics and pairwise metrics

(A–D) The simulation procedure to assess how systematic changes in population metrics lead to changes in pairwise metrics. 

(A) We first systematically varied one of the population metrics while keeping the others fixed. For example, we can increase the loading similarity from a low value (left, blue) to a high value (right, green), while keeping %sv and dimensionality fixed.

(B) Then, we constructed covariance matrices corresponding to each value of the population metric in (A) (see STAR Methods), without generating synthetic data. 

(C) For each covariance matrix from (B), we directly computed the correlations (i.e., the $r_{sc}$ distributions).

(D) We computed $r_{sc}$ mean and $r_{sc}$ SD from the $r_{sc}$ distributions in (C) and then assessed how the change in a given population metric from (A) changed pairwise metrics. In this case, the increase in loading similarity increased $r_{sc}$ mean and decreased $r_{sc}$ SD (blue dot to green dot).

(E) Varying loading similarity with a fixed %sv of 50% and dimensionality of 1. Each dot corresponds to the $r_{sc}$ mean and $r_{sc}$ SD of one simulated covariance matrix with specified population metrics (dots are close together and appear to form a continuum). The color of each dot corresponds to the loading similarity (see STAR Methods), where a value of 1 indicates that all loading weights have the same value.

(F) Varying %sv. The same setting as in (E), except we consider two different values of percent shared variance (50% and 30%). 

(G) Varying dimensionality (i.e., number of co-fluctuation patterns) while sweeping loading similarity between 0 and 1 and keeping %sv fixed at 50%. In this simulation, the relative strengths of each dimension uniform across dimensions (i.e., flat eigenspectra; see STAR Methods).

The third population metric is dimensionality. We define dimensionality as the number of co-fluctuation patterns (or dimensions) needed to explain the shared variability among neurons (see STAR Methods). The variable activity of neurons may depend on multiple common inputs, e.g., top-down signals like attention and arousal (Rabinowitz et al., 2015; Cowley et al., 2020) or spontaneous and uninstructed behaviors (Stringer et al., 2019b; Musall et al., 2019). Furthermore, these common inputs may differ in how they modulate neurons. This may result in two or more dimensions of the population activity (Figure 2D, blue and orange latent co-fluctuations). For illustrative purposes, each dimension might correspond to a single group of tightly coupled neurons (Figure 2D, neurons in top rows have non-zero loadings for pattern 1, whereas neurons in bottom rows have non-zero loadings for pattern 2). However, in general, each neuron can have non-zero loadings for multiple patterns. In population activity space, adding a new dimension adds a new axis along which neurons covary (Figure 2D, orange line). We use the term “dimension” to refer either to a latent co-fluctuation or its corresponding co-fluctuation pattern, depending on context.

Varying population metrics to assess changes in pairwise metrics

Given that both pairwise and population metrics are computed from the same spike count covariance matrix (Figure 1C), a connection should exist between the two. We establish this connection by deriving mathematical relationships and carrying out simulations. In simulations, we assessed how systematically changing one of the population metrics (e.g., increasing loading similarity; Figure 3A) changes the spike count covariance matrix (Figure 3B) and the corresponding $r_{sc}$ distribution (Figure 3C), which we summarized using its mean and standard deviation (Figure 3D). The covariance matrix was parameterized in a way that allowed us to create covariance matrices with specified population metrics (see STAR Methods). Thus, our simulation procedure does not simulate neuronal activity but rather creates covariance matrices that are consistent with the specified population metrics. Loading similarity has opposing effects on $r_{sc}$ mean and SD

We first asked how the loading similarity of a single co-fluctuation pattern (i.e., one dimension) affected $r_{sc}$ mean and SD. Intuitively, a high loading similarity indicates that the activity of all neurons...
increases and decreases together (Figure 2A), resulting in values of $r_{sc}$ that are all positive and similar in value. Indeed, in simulations, we found that high loading similarity corresponded to large $r_{sc}$ mean and $r_{sc}$ SD close to 0 (Figure 3E, green dots near horizontal axis). On the other hand, a low loading similarity indicates that, when some neurons increase their activity, others decrease their activity (Figure 2B), resulting in some positive $r_{sc}$ values (for pairs that change their activity in the same direction) and some negative $r_{sc}$ values (for pairs that change their activity in opposition). In simulations, a low loading similarity indeed corresponded to an $r_{sc}$ mean close to 0 and a large $r_{sc}$ SD (Figure 3E, blue dots near vertical axis). By varying the loading similarity, we surprisingly observed an arc-like trend in the $r_{sc}$ mean versus $r_{sc}$ SD plot (Figure 3E). In Math Note A, we derive the analytical relationship between loading similarity and $r_{sc}$. In Math Note B, we show mathematically why the $r_{sc}$ mean versus $r_{sc}$ SD relationship follows a circular arc.

**Decreasing %sv reduces $r_{sc}$ mean and SD**

We next asked how %sv, which measures the percentage of each neuron’s variance that is shared with other neurons in the population, is related to $r_{sc}$ mean and SD. Intuitively, one might expect %sv and $r_{sc}$ mean to be closely related because $r_{sc}$ measures the degree to which the activity of two neurons is shared (Cohen and Kohn, 2011). We investigated this in simulations and found that how closely %sv and $r_{sc}$ mean were related depended on the loading similarity. When loading similarity was high (Figure 3F, green dots), there was a direct relationship between %sv and $r_{sc}$ mean (specifically, %sv equals $r_{sc}$ mean). However, when loading similarity was low (Figure 3F, blue dots), the relationship between %sv and $r_{sc}$ mean was less direct. Namely, $r_{sc}$ mean remained close to zero, regardless of %sv. This illustrates that $r_{sc}$ mean and %sv are not the same. It is possible for a population of neurons with high %sv (e.g., Figure 3F, blue dots in outer arc) to have smaller $r_{sc}$ mean than a population with lower %sv (e.g., Figure 3F, green dots in inner arc).

These relationships that we have shown through simulation can be captured mathematically. First, if we have knowledge of the loading weights in the co-fluctuation pattern, the $r_{sc}$ between a pair of neurons can be expressed in terms of the %sv and loading values of the two neurons (Math Note A),

$$r_{ij} = \sqrt{\phi_i \phi_j \text{sign}(w_i w_j)},$$

(Equation 1)

where $r_{ij}$ is the $r_{sc}$ between neurons $i$ and $j$, $\phi_i$ and $\phi_j$ are the %sv of each neuron (expressed as a proportion per neuron, in contrast to %sv in Figure 3F, which shows the average %sv across all neurons), and $w_i$ and $w_j$ are the loading weights of the neurons in the co-fluctuation pattern. The $r_{sc}$ mean is the average of $r_{ij}$ values across all neuron pairs. From Equation 1, we observe that, when loading similarity is high (i.e., most loading weights have the same sign), %sv and $r_{sc}$ mean are directly related (i.e., $r_{ij} = \sqrt{\phi_i \phi_j}$). However, when loading similarity is low (i.e., some loading weights are positive and others are negative), $r_{sc}$ mean is small, regardless of %sv, because some pairs have $\text{sign}(w_i w_j) = +1$ and others have $\text{sign}(w_i w_j) = -1$.

Second, if we have information about the $r_{sc}$ SD (instead of loading weights), we can establish the following relationship between %sv, $r_{sc}$ mean, and $r_{sc}$ SD (Math Note B):

$$%sv = \sqrt{(r_{sc} \text{ mean})^2 + (r_{sc} \text{ SD})^2}.$$
Figure 4. Relative strengths of dimensions affect r_sc distributions

With dimensionality of 2, we systematically varied the relative strengths of the two dimensions with a fixed total %sv of 50%. We considered two scenarios: (1) one dimension has high loading similarity and the other has low loading similarity (A) and (2) both dimensions have low loading similarity (B). Each dot represents one simulated covariance matrix and r_sc distribution. The colors of the dots indicate different relative strengths between the two dimensions, and numbers next to each cloud of dots indicate the ratio between the relative strength associated with each dimension. For example, in (A), red dots correspond to the high loading similarity dimension being 19 times stronger (95:5) than the low loading similarity dimension. Black dots correspond to the low loading similarity dimension being 19 times stronger (95:5) than the high loading similarity dimension. In (B), because both patterns have low loading similarity, clouds for 80:20 and 95:5 are very similar to clouds for 20:80 and 5:95, respectively, and are thus omitted for clarity. See also Figure S1.

The relative strength of each dimension impacts pairwise metrics

In the previous simulation (Figure 3G), we assumed that each dimension explained an equal proportion of the overall shared variance (e.g., for two dimensions, each dimension explained half of the shared variance; see STAR Methods). However, it is typically the case for recorded neuronal activity that some dimensions explain more shared variance than others; in other words, neuronal activity co-fluctuates more strongly along some patterns than others (Sadler et al., 2014; Williamson et al., 2016; Mazzucato et al., 2016; Gallego et al., 2018; Huang et al., 2019; Stringer et al., 2019a; Ruff et al., 2020). We sought to assess the influence of the relative strength of each dimension on pairwise metrics.

We reasoned that stronger dimensions would play a larger role than weaker dimensions in determining the r_sc distribution and pairwise metrics. Extending Equation 1 to multiple dimensions, we show that the r_sc between a pair of neurons can be expressed as the sum of a contribution from each constituent dimension (Math Note C). The stronger a dimension, the larger the magnitude of its contribution to r_sc and thus the larger its impact on r_sc mean and SD.

To test this empirically, we performed a simulation with two dimensions while systematically varying the relative strength of each dimension. We considered two scenarios: (1) one dimension has a pattern with high loading similarity and one dimension has a pattern with low loading similarity (Figure 4A) and (2) both dimensions have patterns with low loading similarity (Figure 4B). Note that both dimensions cannot have patterns with high loading similarity because they would not be orthogonal (Math Note E).

In scenario (1), where one dimension’s pattern has high loading similarity and the other has low loading similarity, r_sc mean and r_sc SD reflect the loading similarity of the dominant dimension (Figure 4A). When the dimension with a high loading similarity pattern dominated, r_sc mean was large and r_sc SD was small (Figure 4A, red dots are close to horizontal axis). When the dimension with a low loading similarity pattern dominated, r_sc mean was small and r_sc SD was large (Figure 4A, black dots are close to vertical axis). When the two dimensions were of equal strength (i.e., neither dimension dominated), r_sc mean and r_sc SD were both intermediate values (Figure 4A, light gray dots are between red and black dots). Thus, the dimensions along which neuronal activity co-fluctuates more strongly have a greater influence on pairwise metrics (Figure S1).

In scenario (2), where both dimensions have patterns of low loading similarity, r_sc mean was low and r_sc SD was high (Figure 4B), similar to when there is one dimension with low loading similarity (Figure 3E, blue dots). When we made one dimension stronger than the other, r_sc mean remained low and r_sc SD remained high (Figure 4B, light gray dots and black dots are both close to vertical axis) because both patterns had low loading similarity. However, the radius of the arc increased (Figure 4B, black dots farther from the origin than light gray dots) and was close to the arc that would have been produced with a single dimension (Figure 3G, 1 dim.). Thus, whereas changing the number of dimensions causes discrete jumps in the arc radius (Figure 3G), changing the relative strength of each dimension allows for r_sc mean and r_sc SD to vary continuously between the arcs for different dimensionalities. Put another way, changing the relative strength of each dimension varies the “effective dimensionality” of population activity in a continuous manner. Neuronal activity for which one dimension dominates another (Figure 4B, black dots) has a lower effective dimensionality than when both dimensions have equal strength (Figure 4B, light gray dots).
Figure 5. Summary of relationship between pairwise and population metrics
A change in $r_{sc}$ mean and $r_{sc}$ SD may correspond to changes in loading similarity, %sv, dimensionality, or a combination of the three. Shaded regions indicate the possible $r_{sc}$ mean and $r_{sc}$ SD values for different dimensionalities; increasing dimensionality tends to decrease $r_{sc}$ mean and $r_{sc}$ SD (shaded regions for larger dimensionalities become smaller). Within each shaded region, decreasing %sv decreases both $r_{sc}$ mean and SD radially toward the origin. Finally, rotating co-fluctuation patterns such that the loadings are more similar (going from low to high loading similarity) results in moving clockwise along an arc such that $r_{sc}$ mean increases and $r_{sc}$ SD decreases. We also note two subtle trends. First, there are more possibilities for loading similarity to be low than high (Math Note E), suggesting that $r_{sc}$ SD will generally tend to be larger than $r_{sc}$ mean if neuronal activity varied along a randomly chosen co-fluctuation pattern (shading within each region is darker near the vertical axis than the horizontal axis). Second, this effect becomes exaggerated for higher dimensional neuronal activity, as many dimensions can have low loading similarity but only one dimension can have high loading similarity (Math Note E). Thus, it becomes progressively unlikely for $r_{sc}$ SD to be 0 as dimensionality increases (shaded regions for larger dimensionalities lifted off the horizontal axis).

Reporting only a single statistic provides an incomplete description of population covariability
Figure 5 summarizes the relationships that we have established between pairwise metrics and population metrics. Rotating a co-fluctuation pattern from a low loading similarity to a high loading similarity increases $r_{sc}$ mean and decreases $r_{sc}$ SD along an arc (Figure 5, arrow outside pink arc). Decreasing %sv decreases both $r_{sc}$ mean and SD (Figure 5, arrow pointing toward origin), and increasing dimensionality also tends to decrease $r_{sc}$ mean and SD (Figure 5, pink to yellow shaded regions).

These results provide a cautionary tale that using a single statistic on its own provides an opaque description of population-wide covariability. For example, a change in $r_{sc}$ mean could correspond to changes in loading similarity, %sv, dimensionality, or a combination of the three. Likewise, reporting dimensionality on its own would be incomplete because the role of a dimension in explaining population-wide covariability depends on how much shared variance it explains and the loading similarity of its co-fluctuation pattern. For example, consider a decrease in dimensionality by 1. This would have little impact on population-wide covariability if the removed dimension explains only a small amount of shared variance, whereas it could have a large impact if the removed dimension explains a large amount of shared variance.

Considering multiple statistics together provides a richer description of population-wide covariability. For example, in the case where population activity co-fluctuates along a single dimension, $r_{sc}$ mean and $r_{sc}$ SD can be used together to approximate %sv (using distance from the origin) and deduce whether loading similarity is low ($r_{sc}$ SD > $r_{sc}$ mean) or high ($r_{sc}$ mean > $r_{sc}$ SD), whereas $r_{sc}$ mean alone would not provide much information about %sv or loading similarity (cf. Figure 5). In the next section, we further demonstrate using neuronal recordings how relating pairwise and population metrics using the framework we have developed (Figure 5) provides a richer description of how neurons covary than using a single statistic (e.g., $r_{sc}$ mean) alone.

Case study: V4 neuronal recordings during spatial attention
When spatial attention is directed to the receptive fields of neurons in area V4 of macaque visual cortex, $r_{sc}$ mean among those neurons decreases (Cohen and Maunsell, 2009; Mitchell et al., 2009; Gregoriou et al., 2014; Snyder et al., 2016, 2018). This decrease has often been attributed to a reduction in shared modulations among the neurons. However, we have shown both mathematically and in simulations that several distinct changes in population metrics (e.g., decrease in loading similarity, decrease in %sv, or an increase in dimensionality) could underlie this decrease in $r_{sc}$ mean (Figure 5). Here, we sought to assess which aspects of population-wide covariability underlie, and how each of them contribute to, the overall decrease in $r_{sc}$ mean.

We analyzed activity recorded simultaneously from tens of neurons in macaque V4 while the animal performed an orientation-change detection task (Figure 6A; previously reported in Snyder et al., 2018). To probe spatial attention, we cued the animal to the location of the stimulus that was more likely to change in orientation. As expected, perceptual sensitivity increased for orientation changes in the cued stimulus location (Figure 6A, inset, red dot above black dot). “Attend-in” trials were those in which the cued stimulus location was inside the aggregate receptive fields (RFs) of the recorded V4 neurons, whereas “attend-out” trials were those in which the cued stimulus location was in the opposite visual hemifield.

For pairwise metrics, $r_{sc}$ mean decreased when attention was directed into the RFs of the V4 neurons (Figure 6B, left panel), consistent with previous studies (Cohen and Maunsell, 2009; Mitchell et al., 2009; Gregoriou et al., 2014; Snyder et al., 2016, 2018). We further found that $r_{sc}$ SD was lower for attend-in trials than for attend-out trials, an effect not reported previously (Figure 6B, right panel).

The decrease in both $r_{sc}$ mean and $r_{sc}$ SD could arise from several different types of distinct changes in population-wide covariability (Figure 5). To compute the population metrics, we applied FA separately to attend-out and attend-in trials (see STAR Methods). FA is the most basic dimensionality reduction method that characterizes shared variance among neurons.
and is consistent with how we created covariance matrices in Figures 3 and 4. We found three distinct changes in population metrics. First, neuronal activity during attend-in trials had lower %sv than during attend-out trials (Figure 6C, left), consistent with previous interpretations that attention reduces the strength of shared modulations (Rabinowitz et al., 2015; Ecker et al., 2016; Huang et al., 2019; Ruff et al., 2020). Second, we also found lower loading similarity for attend-in trials than attend-out trials for the dominant dimension (i.e., the dimension that explains the largest proportion of the variance). In (A)–(C), dots indicate means and error bars indicate 1 SEM, both computed across recording sessions. See also Figure S2.

Figure 6. An observed decrease in $r_{sc}$ mean of macaque V4 neurons during a spatial attention task corresponds to changes in multiple population metrics

(A) Experimental task design. On each trial, monkeys maintained fixation while Gabor stimuli were presented for $400 \text{ ms}$ (with $300–500 \text{ ms}$ in between presentations). When one of the stimuli changed orientation, animals were required to saccade to the changed stimulus to obtain a reward. At the beginning of a block of trials, we performed an attentional manipulation by cuing animals to the location of the stimulus that was more likely to change for that block (dashed circle denotes the cued stimulus and was not presented on the screen). The cued location alternated between blocks. Animals were more likely to detect a change in stimulus at cued rather than uncued locations (inset in bottom right, $p < 0.002$ for both animals; data for monkey 1 are shown). During this task, we recorded activity from V4 neurons whose receptive fields (RFs) overlapped with one of the stimulus locations.

(B) $r_{sc}$ mean (left panel) and $r_{sc}$ SD (right panel) across recording sessions for two animals. Black denotes “attend-out” trials (i.e., the cued location was outside the recorded V4 neurons’ RFs), and red denotes “attend-in” trials (i.e., the cued location was inside the RFs). Data were pooled across both animals to compute $p$ values reported in titles for comparison of attend-out (black) and attend-in (red). For individual animals, $r_{sc}$ mean was lower for attend-in than attend-out ($p < 0.001$ for each animal), and $r_{sc}$ SD was also lower for attend-in than attend-out ($p < 0.05$ for monkey 1 and $p = 0.148$ for monkey 2).

(C) Population metrics identified across recording sessions for two animals (same data as in B). Black denotes attend-in trials; red denotes attend-out trials. Data were again pooled across animals to compute $p$ values reported in titles for comparing attend-out and attend-in. %sv was lower for attend-in than attend-out ($p < 0.001$ for monkey 1 and $p < 0.02$ for monkey 2). Loading similarity was lower for attend-in than attend-out ($p < 0.001$ for monkey 1 and $p = 0.162$ for monkey 2). Dimensionality was lower for attend-in than attend-out ($p = 0.113$ for monkey 1 and $p = 0.174$ for monkey 2). In (A)–(C), dots indicate means and error bars indicate 1 SEM, both computed across recording sessions. See also Figure S2.

(D) Summary of the real data results. Attention decreases both $r_{sc}$ mean and $r_{sc}$ SD (black dot to red dot). These decreases in pairwise metrics correspond to a combination of decreases in %sv, loading similarity, and dimensionality (dashed arrows).

See also Figures S3, S4, and S6.
shared variance; Figure 6C, middle; see also Figure S2B). This implies that, with attention, neurons in the population co-fluctuate in a more heterogeneous manner (i.e., more pairs of neurons co-fluctuate in opposition and fewer pairs co-fluctuate together). Third, we found that dimensionality was slightly lower for attend-in than attend-out trials (Figure 6C, right). Thus, on average, a smaller number of distinct shared signals were present when attention was directed into the neurons’ RFs. The small change in dimensionality is consistent with the relative strength of each dimension (i.e., eigenspectrum shape) being similar for attend-in and attend-out (Figure S2A). Taken together, this collection of observations of both pairwise and population metrics leads to a more refined view of how attention affects population-wide covariability.

The pairwise (Figure 6B) and population (Figure 6C) metrics are computed based on the same recorded activity, and each represents a different view of population activity. The central contribution of our work is to provide a framework by which to understand these two perspectives and five different metrics in a coherent manner. Using the relationships between pairwise and population metrics we have established in the $r_{sc}$ mean versus $r_{sc}$ SD space (Figure 5), we can decompose the decrease in $r_{sc}$ mean and SD into (1) a small decrease in dimensionality (Figure 6D, small dashed arrow), (2) a decrease in loading similarity (Figure 6D, medium dashed arrow), and (3) a substantial decrease in %sv (Figure 6D, large dashed arrow). We quantify these contributions in Figure S3. The $r_{sc}$ mean and SD decreased despite the decrease in dimensionality (which alone would have tended to increase $r_{sc}$ mean and SD) because of the larger contributions of loading similarity and %sv to pairwise metrics in these V4 recordings. We have also applied the same analysis to population recordings in visual area V1 (Zandvakili and Kohn, 2015; available on http://crcns.org) and found that, although $r_{sc}$ mean and SD both decreased (like in the V4 recordings), the population metrics changed in a different way compared to the V4 recordings (Figure S4). Together, these analyses demonstrate the need for considering both pairwise and population metrics together when studying correlated variability, with a bridge that allows one to navigate between the two.

**DISCUSSION**

Coordinated variability in the brain has long been linked to the neural computations underlying a diverse range of functions, including sensory encoding, decision making, attention, learning, and more. In this study, we sought to relate two major bodies of work investigating the coordinated activity among neurons: studies that measure spike count correlation between pairs of neurons ($r_{sc}$) and studies that use dimensionality reduction to measure population-wide covariability. We considered three population metrics and established analytically and empirically that (1) increasing loading similarity corresponds to increasing $r_{sc}$ mean and decreasing $r_{sc}$ SD, (2) decreasing %sv corresponds to decreasing both $r_{sc}$ mean and SD, and (3) increasing dimensionality tends to decrease $r_{sc}$ mean and SD. Applying this understanding to recordings in macaque V4, we found that the previously reported decrease in mean spike count correlation associated with attention stemmed from a decrease in %sv, a decrease in loading similarity, and decrease in dimensionality. This analysis revealed that attention involves multiple changes
in how neurons interact that are not well captured by a single statistic alone. Overall, our work demonstrates that common ground exists between the literatures of spike count correlation and dimensionality reduction approaches and builds the intuition and formalism to navigate between them.

Our work also provides a cautionary tale for attempting to summarize population-wide covariability using one, or a small number of, statistics. For example, reporting only $r_{sc}$ mean is incomplete because several distinct changes in population-wide covariability can correspond to the same change in $r_{sc}$ mean. In a similar vein, reporting only dimensionality is incomplete because it does not indicate how strongly the neurons covary or their co-fluctuation patterns. For this reason, we recommend reporting several different pairwise and population metrics (e.g., the five used in this study along with the eigenspectrum of the shared covariance matrix), as long as they can be reliably measured from the data available. This not only allows for a deeper and more complete understanding of how neurons covary, but also it allows one to make tighter connections to previous literature that uses the same metrics. Future work may seek to revisit previous results of correlated neuronal variability that are based on a single statistic (e.g., $r_{sc}$ mean) and reinterpret them within a framework that considers multiple perspectives and statistics of population-wide covariability, such as that presented here.

There are some situations where it is not feasible to reliably measure population statistics, such as recording from a small number of neurons in deep brain structures (Nevet et al., 2007; Liu et al., 2013) or when the number of trials is small relative to the number of neurons recorded (Wainwright, 2019). In such situations, the $r_{sc}$ can be measured between pairs of neurons recorded in each session and then averaged across sessions to obtain the $r_{sc}$ mean. Based on our findings, we recommend that studies that report $r_{sc}$ mean also report $r_{sc}$ SD because the latter provides additional information about population-wide covariability. For example, in the special case of one latent dimension (typically not known in advance for real data), measuring $r_{sc}$ mean and $r_{sc}$ SD allows one to estimate the loading similarity and %sv (cf. Figures 3E and 3F). In general, even when there is more than one latent dimension in the population, $r_{sc}$ SD provides value in situating the data in the $r_{sc}$ mean versus $r_{sc}$ SD plot (Figure 5). Changes in $r_{sc}$ mean and SD can then inform changes in population metrics based on the relationships established in this work (cf. Figure 6D).

The reason that our work, and many previous studies, have focused on trial-to-trial variability is that it has important implications for information coding. Early work on information-limiting correlations typically focused on $r_{sc}$ mean (e.g., Zohary et al., 1994; Shadlen and Newsome, 1998; Cohen and Maunsell, 2009; Cohen and Kohn, 2011), which reflects the strength of shared variability among neurons. Recent theoretical work (Averbeck et al., 2006; Moreno-Bote et al., 2014; Kohn et al., 2016) and experimental evidence (Ni et al., 2018; Ruff and Cohen, 2019a; Cowley et al., 2020; Rumyantsev et al., 2020; Bartolo et al., 2020) have shown that it is not only the strength of shared trial-to-trial variability but also the directions of shared variability relative to stimulus tuning (Figure 7A) that need to be considered for information coding. These properties of shared trial-to-trial variability are precisely what are measured by the population metrics used here. In particular, the %sv measures how strongly trial-to-trial variability is shared among neurons (Figure 7B), loading similarity measures the direction(s) of variability (Figure 7C), and dimensionality measures how many different directions of variability exist in the data (Figure 7D). By considering these three population metrics together, along with the way in which mean population responses vary across conditions (i.e., the stimulus-encoding directions), we can more incisively characterize how trial-to-trial variability impacts information coding than by using $r_{sc}$ mean alone. Understanding how patterns of shared variability are related to (e.g., align with or are orthogonal to) patterns of stimulus encoding and downstream readouts will be likely critical for understanding information coding in the brain.

We considered three population metrics—dimensionality, %sv, and loading similarity—that summarize the structure of population-wide covariability and are rooted in well-established concepts in existing literature. First, dimensionality has been used to describe how neurons covary across conditions (i.e., an analysis of trial-averaged firing rates; Churchland et al., 2012; Rigotti et al., 2013; Mante et al., 2013; Cowley et al., 2016; Kobak et al., 2016; Sohn et al., 2019), as well as how neurons covary from trial to trial (Yu et al., 2009; Santhanam et al., 2009; Sadler et al., 2014; Rabinowitz et al., 2015; Mazzucato et al., 2016; Williamson et al., 2016; Bittner et al., 2017; Athalye et al., 2017; Williams et al., 2018; Stringer et al., 2019a; Recanatesi et al., 2019). We focused on the latter in our study to connect with the $r_{sc}$ literature, which also seeks to understand the shared trial-to-trial variability between neurons. To focus on the shared variability among neurons, we used FA to measure dimensionality. Another commonly used dimensionality reduction method, principal-component analysis (PCA), although appropriate for studying trial-averaged activity, does not distinguish between variability that is shared among neurons and variability that is independent to each neuron. Second, investigating the loading similarity has provided insight about whether shared variability among neurons arises from a shared global factor that drives neurons to increase and decrease their activity together (Ecker et al., 2014; Okun et al., 2015; Lin et al., 2015; Rabinowitz et al., 2015; Williamson et al., 2016; Huang et al., 2019) or whether the co-fluctuations involve a more intricate pattern across the neuronal population (Snyder et al., 2018; Insanally et al., 2019; Cowley et al., 2020). Third, we have previously reported %sv for area V1 (Williamson et al., 2016), area M1 (Henning et al., 2018), and network models (Williamson et al., 2016; Bittner et al., 2017). Conceptually, %sv and $r_{sc}$ mean are both designed to capture the strength of shared variability in a population of neurons. Thus, we might initially think that there should be a one-to-one correspondence between the two quantities. Indeed, if the population activity is described by one co-fluctuation pattern with a high loading similarity, there is a direct relationship between %sv and $r_{sc}$ mean (Figure 3F). However, in general, %sv and $r_{sc}$ mean do not have a one-to-one correspondence between them (Figure 3F, moderate or low loading similarity).

We focus here on studying trial-to-trial activity fluctuations that are shared between neurons. Many studies have considered the...
source of these shared fluctuations in the context of pairwise correlations (Cohen and Kohn, 2011). Most commonly, pairwise correlations have been suggested to originate through common input (Zohary et al., 1994; Shadlen and Newsome, 1998). However, there are, in fact, numerous mechanisms that can shape the trial-by-trial shared variability of neuronal populations, including neuromodulation (Harris and Theile, 2011; Herrero et al., 2013; Minces et al., 2017), coupled inhibition (Haider et al., 2006), or distinct patterns of neuronal connectivity (Mazzucato et al., 2016; Williamson et al., 2016; Huang et al., 2019; Recanatesi et al., 2019). These mechanisms likely produce distinct signatures in population metrics, such as %s, loading similarity, and dimensionality. The framework that we have developed here can be applied to spiking network models with different underlying mechanisms of shared cortical variability to identify signatures in population metrics (Mazzucato et al., 2016; Williamson et al., 2016; Huang et al., 2019; Recanatesi et al., 2019). We can then assess whether any of those signatures are present in neuronal recordings to gain insight into the underlying mechanisms of shared variability in the brain.

Although pairwise correlation and dimensionality reduction have most commonly been computed based on spike counts, several studies have also computed these metrics on neuronal activity recorded using other modalities, such as calcium imaging (Harvey et al., 2012; Ahrens et al., 2012; Dechery and Mageean, 2018; Stringer et al., 2019a; Rumyantsev et al., 2020). The relationships that we established here between pairwise and population metrics are properties of covariance matrices in general and do not rely on or assume recordings of neuronal spikes. Thus, the intuition built here can be applied to other recording modalities.

Our work here focused on studying interactions within a single population of neurons. Technological advances are enabling recordings from multiple distinct populations simultaneously, including neurons in different brain areas, neurons in different cortical layers, or different neuron types (e.g., Ahrens et al., 2013; Jiang et al., 2015; Jun et al., 2017). Studies are dissecting the interactions between these distinct populations using pairwise correlation (Smith et al., 2013; Pooresmaeili et al., 2014; Oemisch et al., 2015; Zandvakili and Kohn, 2015; Ruff and Cohen, 2016a; Snyder et al., 2016) and dimensionality reduction (Semedo et al., 2014; Buesing et al., 2014; Bittner et al., 2017; Perich et al., 2018; Semedo et al., 2019; Ames and Churchland, 2019; Ruff and Cohen, 2019a; Veuthey et al., 2020; Cowley et al., 2020). As we have shown here for a single population of neurons, considering a range of metrics from both the pairwise correlation and dimensionality reduction perspectives and understanding how they relate to one another will provide rich descriptions of how different neuronal populations interact.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

- **KEY RESOURCES TABLE**
- **RESOURCE AVAILABILITY**

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.neuron.2021.06.028.

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**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

The authors declare no competing interests.
REFERENCES


Neuron


STAR METHODS

KEY RESOURCES TABLE

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RESOURCE AVAILABILITY

**Lead contact**
Further information and requests for resources should be directed to and will be fulfilled by the Lead Contact, Byron M. Yu (byronyu@cmu.edu).

**Materials availability**
This study did not generate new unique reagents.

**Data and code availability**
Original code has been deposited at Zenodo and is publicly available as of the date of publication. DOIs are listed in the key resources table. Additional information or data are available upon request from the lead contact (byronyu@cmu.edu).

METHOD DETAILS

**Spike count covariance matrix**
Both pairwise metrics and population metrics are computed directly from the spike count covariance matrix Σ of size $n \times n$ for a population of $n$ neurons. Each entry in Σ is the covariance between the activity of neuron $i$ and neuron $j$:

$$\Sigma_{ij} = \text{cov}(x_i, x_j) = \mathbb{E}[(x_i - \mu_i)(x_j - \mu_j)]$$

(Equation 2)

where $x_i$ and $x_j$ represent the activity of neurons $i$ and $j$, respectively, and $\mu_i$ and $\mu_j$ represent the mean activity of neurons $i$ and $j$, respectively. The variance of the $i$th neuron is equal to $\Sigma_{ii}$.

**Pairwise metrics**
We computed the spike count correlation ($r_{sc}$) between neurons $i$ and $j$ directly from the spike count covariance matrix:

$$\rho_{ij} = \frac{\Sigma_{ij}}{\sqrt{\Sigma_{ii}\Sigma_{jj}}}$$

(Equation 3)

We then summarized the distribution of $r_{sc}$ values across all pairs of neurons in the population with two pairwise metrics: the $r_{sc}$ mean and $r_{sc}$ standard deviation (SD).
Population metrics
The metrics we use for characterizing population-wide covariability are based on factor analysis (FA; Santhanam et al., 2009; Yu et al., 2009; Churchland et al., 2010; Harvey et al., 2012; Williamson et al., 2016; Bittner et al., 2017; Athalye et al., 2017; Huang et al., 2019), a dimensionality reduction method. We chose FA because it is the most basic dimensionality reduction method that explicitly separates variance that is shared among neurons from variance that is independent to each neuron. This allows us to relate the population metrics provided by FA to spike count correlation, which is designed to measure shared variability between pairs of neurons. One might consider using principal component analysis (PCA), but it does not distinguish shared variance from independent variance. Thus, FA is more appropriate than PCA for studying the shared variability among a population of neurons.

Decomposing the spike count covariance matrix
FA decomposes the spike count covariance matrix $\Sigma$ into a low-rank shared covariance matrix, which captures the variability shared among neurons in the population, and an independent variance matrix, which captures the portion of variance of each neuron unexplained by the other neurons (Figure S5A):

$$\Sigma = \Sigma_{\text{shared}} + \Psi$$  
(Equation 4)

where $\Sigma_{\text{shared}} \in \mathbb{R}^{n \times n}$ is the shared covariance matrix for $n$ neurons, and $\Psi \in \mathbb{R}^{n \times n}$ is a diagonal matrix containing the independent variance of each neuron. The low-rank shared covariance matrix can be expressed using the eigendecomposition as (Figure S5A):

$$\Sigma_{\text{shared}} = U \Lambda U^T$$  
(Equation 5)

where $U \in \mathbb{R}^{n \times d}$ and $\Lambda \in \mathbb{R}^{d \times d}$, with $d < n$. The rank (i.e., dimensionality) of the shared covariance matrix, $d$, indicates the number of latent variables. Each column of $U$ is an eigenvector and represents a co-fluctuation pattern containing the loading weights of each neuron (i.e., how much each neuron contributes to that dimension). The matrix $\Lambda$ is a diagonal matrix where each diagonal element is an eigenvalue and represents the amount of variance along the corresponding co-fluctuation pattern (e.g., in Figure 2A has larger eigenvalue than 2C).

Based on this matrix decomposition, we defined the three metrics that describe the population-wide covariability:

- **Loading similarity**: the similarity of loading weights across neurons for a given co-fluctuation pattern. Scalar value between 0 (the weights are maximally dissimilar, defined precisely below) and 1 (all weights are the same).

- **Percent shared variance (%sv)**: the percentage of each neuron’s variance that is explained by other neurons in the population. Percentage between 0% and 100%.

- **Dimensionality**: the number of dimensions (i.e., co-fluctuation patterns). Integer value.

We give the precise definitions of these population metrics below and in Figure S5B.

Loading similarity
We sought to define loading similarity such that, for a given co-fluctuation pattern, if the weights for all neurons are the same, we would measure a loading similarity of 1. When the weights are as different as possible, we would measure a loading similarity of 0. We define the loading similarity based on the variance across the $n$ weights (for $n$ neurons) in a co-fluctuation pattern $\mathbf{u}_k$. The smallest possible variance is 0; the largest possible variance, for a unit vector, is $1/n$ (Math Note F). Thus, we define loading similarity for a co-fluctuation pattern $\mathbf{u}_k \in \mathbb{R}^n$ as:

$$\text{loading similarity}(\mathbf{u}_k) = 1 - \frac{\text{var}(\mathbf{u}_k)}{\max_{\mathbf{v}_k} \text{var}(\mathbf{v}_k)} = 1 - \frac{\text{var}(\mathbf{u}_k)}{1/n}$$  
(Equation 6)

where the loading similarity is computed on unit vectors (i.e., $\mathbf{u}_k$ has a norm of 1). The notation $\text{var}(\mathbf{u}_k)$ denotes that the variance is being taken across the $n$ elements of the vector $\mathbf{u}_k$. The denominator of Equation 6 acts as a normalizing factor, bounding the loading similarity value between 0 and 1.

The loading similarity distinguishes between a co-fluctuation pattern along which all neurons in the population have the same weight in which case they change their activity up and down together (Figure 2A: loading similarity of 1), from one in which weights are different and some neurons increase their activity when others decrease their activity (Figure 2B: loading similarity of 0). The loading weights we use here are closely related to ‘population coupling’ (Okun et al., 2015) and ‘modulator weights’ (Rabinowitz et al., 2015). For some types of shared fluctuations, these weights are similar across neurons in a population (i.e., high loading similarity; Okun et al., 2015; Rabinowitz et al., 2015; Huang et al., 2019). For other types of shared fluctuations, the weights vary substantially across neurons in the population (i.e., low loading similarity; Snyder et al., 2018; Cowley et al., 2020).

We show in Math Note E why, if one dimension has high loading similarity, the other dimensions must have low loading similarity. The reason is that co-fluctuation patterns are defined to be mutually orthogonal. If one co-fluctuation pattern has all weights close to the same value (i.e., high loading similarity), then all other co-fluctuation patterns must have substantial diversity in their weights (i.e., low loading similarity) to satisfy orthogonality.
Percent shared variance
The percent shared variance (%sv) measures the percentage of each neuron’s spike count variance that is explained by other neurons in the population (Williamson et al., 2016; Bittner et al., 2017; Hennig et al., 2018). Equivalently, we can think of %sv in terms of latent co-fluctuations. Because latent co-fluctuations capture the shared variability among neurons, the %sv measures how much of each neuron’s variance is explained by the latent co-fluctuations. The activity of neurons may be tightly linked to the latent co-fluctuation (e.g., Figure 2A), in which case a large percentage of each neuron’s variance is shared with other neurons, or may only be loosely linked to the latent co-fluctuation (e.g., Figure 2C), in which case a small percentage of each neuron’s variance is shared with other neurons. Mathematically, we define the %sv for a neuron $i$ according to

$$%sv \text{ for neuron } i = \frac{\sum_{j=1}^{n} \frac{S_{ij}}{S_j}}{\sum_{j=1}^{n} 100\%} = \frac{S_i}{S_i + \psi_i} \times 100\%$$

(Equation 7)

where $s_i$ is the $i$th entry along the diagonal of the shared covariance matrix ($\Sigma_{shared}$, $\Sigma_{shared}$), and $\psi_i$ is the $i$th entry along the diagonal of the independent covariance matrix ($\Sigma_{independent}$, $\Psi$). A %sv of 0% indicates that the neuron does not covary with (i.e., is independent of) other neurons in the population, whereas a %sv of 100% indicates that the neuron’s activity can be entirely accounted for by the activity of other neurons in the population. To compute %sv for an entire population of neurons, we averaged the %sv of the individual neurons. All %sv values reported in this study are the %sv for the neuronal population.

Dimensionality
Dimensionality refers to the number of latent co-fluctuations needed to describe population-wide covariability. For example, the population-wide covariability can be described by one latent co-fluctuation (Figure 2A) or by several latent co-fluctuations (Figure 2D). In the population activity space, dimensionality corresponds to the number of axes along which the population activity varies (see Figure 2D, bottom inset). Mathematically, the dimensionality is the rank of the shared covariance matrix (i.e., the number of columns in $U$, Figure S5A).

Creating the spike count covariance matrices with specified population metrics
To relate pairwise and population metrics, we created spike count covariance matrices of the form in Equation 4 with specified population metrics. Importantly, we did not simulate spike counts, nor fit a factor analysis model to simulated data. Rather, we created covariance matrices using (4) and computed pairwise correlations directly from the entries of the covariance matrix, as shown in (3). Across simulations (Figures 3 and 4), we simulated with $n = 30$ neurons and set independent variances (i.e., diagonal elements of $\Psi$ in Equation 4) to 1.

Specifying co-fluctuation patterns to obtain different loading similarities
Each co-fluctuation pattern $u_k$ is a vector with $n = 30$ entries (one entry per neuron). We generated a single co-fluctuation pattern by randomly drawing 30 independent samples from a Gaussian distribution with a mean of 2.5. We choose a nonzero mean so that we could obtain co-fluctuation patterns with loading similarities close to 1 when drawing from the Gaussian distribution (i.e., a mean of 0 would have resulted in almost all co-fluctuation patterns having a loading similarity close to 0). To get a range of loading similarities between 0 and 1, we used different standard deviations for the Gaussian. For a small standard deviation value, all entries in the co-fluctuation pattern are close to 2.5, resulting in a high loading similarity. For larger standard deviations, some loading weights are positive and some negative, with large variability in their values, resulting in co-fluctuation patterns with low loading similarity. We increased the Gaussian standard deviation from 0.1 to 5.5 with increments of size 0.1. For each increment, we generated 50 patterns and normalized them to have unit norm. In total, we created a set of 2,750 random patterns.

The following procedure describes the construction of shared covariance matrices with one co-fluctuation pattern. We chose a single pattern $u_k \in \mathbb{R}^{30 \times 1}$ (i.e., $U$ has only 1 column) from the set of 2,750. We constructed the shared covariance matrix by computing $U \Lambda U^T$, where $\Lambda$ was chosen to achieve a desired percent shared variance (see below). The covariance matrix was then computed according to Equation 4. We created a covariance matrix, yielding a spread of loading similarities between 0 and 1 (Figures 3E and 3F). In the next section, we describe the procedure for creating a covariance matrix with more dimensions.

Specifying the percent shared variance
To achieve a given %sv, either the independent variance or the amount of shared variability (i.e., the eigenvalues) of each dimension can be adjusted. In the main text, we set the independent variance of each neuron to $\Psi_i = 1$, and changed the total amount of shared variability by multiplying each eigenvalue (each diagonal element in $\Lambda$ from Equation 5) by the same constant value, $a$. To obtain a specified %sv, we identified $a$ by searching through a large set of possible values (from $10^{-4}$ to $10^3$ with step size $10^{-3}$). We allowed for a tolerance of $e = 10^{-3}$ between the desired %sv and the %sv that was achieved after scaling the eigenvalues by $a$. In other analyses (not shown), we allowed the independent variances to be different across neurons (e.g., drawn from an exponential distribution), and the relationships between pairwise and population metrics were qualitatively similar to those in the main text.
Increasing dimensionality

To assess how changing dimensionality affects pairwise metrics, we created covariance matrices whose shared covariance matrix comprised more than one dimension. To create a shared covariance matrix with \( d \) dimensions, we randomly chose \( d \) patterns from the set of 2,750 we had generated above (see ‘Specifying co-fluctuation patterns to obtain different loading similarities’). We then orthonormalized the chosen patterns using the Gram-Schmidt process to obtain \( d \) orthonormal (i.e., orthogonal and unit length) co-fluctuation patterns \( U \in \mathbb{R}^{m \times d} \). We formed the shared covariance matrix using \( U \Lambda U^T \), where \( \Lambda \in \mathbb{R}^{d \times d} \) is a diagonal matrix containing the eigenvalues (i.e., the strength of each dimension; see ‘Specifying the relative strengths of each dimension’ below). We repeated this procedure to produce 3,000 sets of \( d \) orthonormal patterns (i.e., 3,000 different \( U \) matrices), each of which was used to create a shared covariance matrix. The spike count covariance was computed according to Equation 4.

Specifying the relative strengths of each dimension

In simulating shared covariance matrices with more than one dimension, we chose the relative strength of each dimension by specifying the eigenspectrum (diagonal elements of \( \Lambda \) in Equation 5). We worked with three sets of eigenspectra. First, a flat eigenspectrum had eigenvalues that were all equal (Figure 3G). Second, for two dimensions, we varied the ratio of the two eigenvalues between 95:5, 80:20, 50:50, 20:80, and 5:95 (Figure 4). Third, we considered an eigenspectrum in which each subsequent eigenvalue falls off according to an exponential function (Figure S1). Only the relative (and not the absolute) eigenvalues (i.e., the shape of the eigenspectrum) affect the results, because the eigenspectrum was subsequently scaled to achieve a desired \( \%sv \) (see ‘Specifying the values of percent shared variance’).

Analysis of V4 neuronal recordings from a spatial attention task

Electrophysiological recordings

We analyzed data from a visual spatial attention task reported in a previous study (Snyder et al., 2018). Briefly, we implanted a 96-electrode “Utah” array (Blackrock Microsystems; Salt Lake City, UT) into visual cortical area V4 of an adult male rhesus macaque monkey (data from two monkeys were analyzed; in our study, monkey 1 corresponds to “monkey P” and monkey 2 corresponds to “monkey W” from Snyder et al., 2018). After recording electrode voltages (Ripple Neuro.; Salt Lake City, UT), we used custom software to perform offline spike sorting (Kelly et al., 2007, freely available at https://github.com/smithlabvision/spikesort). This yielded 93.2 ± 8.9 and 61.9 ± 27.4 candidate units per session for monkey 1 and 2, respectively. Experiments were approved by the Institutional Animal Care and Use Committee of the University of Pittsburgh and were performed in accordance with the United States National Research Council’s Guide for the Care and Use of Laboratory Animals.

To further ensure the isolation quality of recorded units, we removed units from our analyses according to the following criteria. First, we removed units with a signal-to-noise ratio of the spike waveform less than 2.0 (Kelly et al., 2007). Second, we removed units with overall mean firing rates less than 1 Hz, as estimates of \( \tau_{rc} \) for these units tends to be poor (Cohen and Kohn, 2011). Third, we removed units that had large and sudden changes in activity due to unstable recording conditions. For this criterion, we divided the recording session into ten equally-sized blocks and for each unit computed the difference in average firing rate between adjacent blocks. We excluded units with a change in average firing rate greater than 60% of the maximum firing rate (where the maximum is taken across the ten equally-sized blocks). Fourth, we removed an electrode from each pair of electrodes that were likely electrically-coupled. We identified the coupled electrodes by computing the fraction of threshold crossings that occurred within 100 µs of each other for each pair of electrodes. We then removed the fewest number of electrodes to ensure this fraction was less than 0.2 (i.e., pairs with an unusually high number of coincident spikes) for all pairs of electrodes. Fifth, we removed units that did not sufficiently respond to the visual stimuli used in the experiment. Evoked spike counts (i.e., a neuron’s response after stimulus presentation) were taken between 50 ms to 250 ms after stimulus onset, and spontaneous spike counts (i.e., a neuron’s response during a blank screen) were taken in a 200 ms window that ended 50 ms before stimulus onset. For each unit, we computed a sensitivity measure \( d’ \) between evoked and spontaneous activity:

\[
d’ = \frac{\mu_{\text{evoked}} - \mu_{\text{spontaneous}}}{\sqrt{\frac{\sigma^2_{\text{evoked}} + \sigma^2_{\text{spontaneous}}}{2}}}
\]

for mean spike counts \( \mu_{\text{evoked}} \) and \( \mu_{\text{spontaneous}} \) and spike count variances \( \sigma^2_{\text{evoked}} \) and \( \sigma^2_{\text{spontaneous}} \). We removed units with \( d’ < 0.5 \) from analyses, as these units had spontaneous and evoked responses that were difficult to distinguish.

After applying these five criteria, 44.5 ± 11.3 and 18.8 ± 6.7 units per session (mean ± s.d. over sessions) remained for monkeys 1 and 2, respectively. Although these remaining units likely contained both single-unit and multi-unit activity, we refer to each unit as a neuron for simplicity. In this study, we restricted analyses to sessions with at least 10 neurons remaining after applying the above criterion (23 sessions for monkey 1, and 14 sessions for monkey 2).

Visual stimulus change-detection task

Animals were trained to perform a change-detection task with a spatial attention cue to the location of the visual stimulus that was more likely to change (Snyder et al., 2018). In the visual change-detection task (Figure 6A), animals fixated a central dot while Gabor stimuli were presented in two locations on a computer screen. One location was chosen to be within the aggregate receptive fields
(RFs) of the recorded V4 neurons (mapped prior to running the experiment), and the other location was placed at the mirror symmetric location in the opposite hemifield. Animals maintained fixation while a sequence of Gabor stimuli were presented. Each drifting Gabor stimulus (oriented at either 45° or 135°) was presented for 400 ms, followed by a blank screen presented for a random interval (between 300 and 500 ms). The sequence continued, with a fixed probability for each presentation, until one of the two stimuli changed orientation when presented (i.e., the ‘target’). Upon target presentation, animals were required to make a saccade to the target to earn a juice reward. We manipulated spatial attention in the experiment by cueing the more probable target location in blocks. At the beginning of each block, the cue was denoted by presenting only one Gabor stimulus at the more probable target location (90% likely), and requiring animals to detect orientation changes at this location for 5 trials. Consistent with the results of previous studies, we found that animals had greater perceptual sensitivity for orientation changes at the cued (i.e., attended) location than the uncued location (Figure 6A, inset in the bottom right) and shorter reaction times (Snyder et al., 2018).

Data processing and computing spike counts
We first separated the trials into two groups: (1) “attend in” trials, for which the cued stimulus was inside the recorded neurons’ RFs and (2) “attend out” trials, for which the cued stimulus was outside the RFs. Since the initial orientation of the stimulus at the cued location could be one of two values (i.e., 45° or 135°), we further divided trials, resulting in a total of 4 groups of trials per session (attend in & 45°, attend out & 45°, attend in & 135°, attend out & 135°). Each combination of cued location and stimulus orientation was treated as an independent sample. The same neurons were used for each of the 4 groups within each session, ensuring a fair comparison between the attend-in and attend-out conditions.

We analyzed all stimulus presentations for which the target stimulus did not change. For each stimulus presentation, we took spike counts in a 200 ms window starting 150 ms after stimulus onset. For each of the 4 groups, we formed a spike count matrix $X \in \mathbb{R}^{n \times t}$, containing the spike counts of the $n$ recorded neurons for the $t$ trials belonging to that group. These spike count matrices were then used to compute both the pairwise and population metrics (described below). For all analyses (Figure 6), we excluded recording sessions with fewer than 10 neurons. Additionally, because population metrics depend on the number of trials (Williamson et al., 2016), for each session we equalized the number of trials across the 4 groups by randomly subsampling from groups with larger numbers of trials.

Computing pairwise metrics for V4 spike counts
We computed pairwise metrics on each combination of attention state (‘attend in’ and ‘attend out’) and stimulus orientation. We computed the correlation as described above in ‘Pairwise metrics’ and then computed $r_{sc}$ mean and $r_{sc}$ SD. For each attention state, we averaged the $r_{sc}$ mean and $r_{sc}$ SD over sessions and different stimulus orientations.

Computing population metrics for V4 spike counts
We fit the parameters of a factor analysis model (see Figure S5A) to each spike count matrix $X$ (as described above) using the expectation-maximization (EM) algorithm (Dempster et al., 1977). For each session, this was performed separately for each attention state and stimulus orientation. Using the FA parameters, we then computed the three population metrics (Figure S5B). For dimensionality, we first found the number of dimensions $d$ that maximized the cross-validated data likelihood. We fit an FA model with $d$ dimensions, and then found the number of dimensions required to explain 95% of the shared variance, termed $d_{shared}$ (Williamson et al., 2016). We report $d_{shared}$ because it tends to be a more reliable estimate of dimensionality than the number of dimensions that maximizes the cross-validated data likelihood. We computed %SV as described by Equation 7. We report the loading similarity as defined in Equation 6 for the co-fluctuation pattern that explained the most shared variability (i.e., the eigenvector with the largest eigenvalue; see Figure S1 for why the loading similarity of this dimension is most informative), since it contributes most to describing the population-wide covariability. For ‘attend in’ and ‘attend out’ conditions, we averaged the population metrics across sessions and stimulus orientations.

Much of our work focuses on systematically changing a single population metric and assessing changes in pairwise metrics (Figures 3A–3D). When analyzing neuronal recordings, one needs to fit factor analysis to the recordings in order to estimate the population metrics. When estimating the population metrics together, it could be the case that changes in one population metric impacts or biases the estimation of another population metric. We characterized these estimation errors in Figure S6. Moreover, in Figure S7, we show that our main findings (Figure 5) are the same when estimating population metrics from Poisson simulated data, which resembled realistic neuronal activity.

Statistics
We employed paired permutations tests for all statistical comparisons of pairwise metrics and population metrics between ‘attend-in’ and ‘attend-out’ conditions (Figures 6B and 6C). First, for a given metric, we computed its value separately for each stimulus type (i.e., 45° or 135°), condition (i.e., attend-in or attend-out), and session. We then averaged the difference between attend-in and attend-out across stimulus types and sessions. To compute a null distribution, we randomly permuted the pair of attend-in and attend-out labels for each stimulus type and condition combination and recomputed the average difference. We ran 10,000 permutations to obtain a
null distribution of 10,000 samples. We computed p-values as the proportion of samples in the null distribution that were more extreme than the average difference in the data, corresponding to p < 0.0001 as the highest attainable level of significance in our statistical analyses.

Math Notes

A) Relationship between correlation, loading similarity, and %sv (one latent dimension)

We establish here the mathematical relationship between $r_{sc}$, loading similarity, and %sv. This will provide the formalism for understanding why decreasing %sv decreases both $r_{sc}$ mean and SD (Figure 3F), that a high loading similarity corresponds to large $r_{sc}$ mean and low $r_{sc}$ SD (Figure 3E), and that a low loading similarity corresponds to small $r_{sc}$ mean and large $r_{sc}$ SD (Figure 3E).

Let $n$ be the number of neurons, and let $w$ be the co-fluctuation pattern (i.e., loading vector $[w_1, w_2, \ldots, w_n]^T \in \mathbb{R}^{n \times 1}$), $\lambda \in \mathbb{R}$, be the strength of the co-fluctuation pattern (i.e., eigenvalue of the shared covariance matrix), and $\Psi \in \mathbb{R}^{n \times n}$ be a diagonal matrix specifying the independent variance of each neuron $(\psi_1, \psi_2, \ldots, \psi_n)$. Then the covariance matrix of the population activity is (see STAR Methods and Figure S5):

$$
\Sigma = \Sigma_{shared} + \Psi = w \lambda w^T + \Psi
$$

From this, we observe that $\Sigma_{ij} = \Sigma_{shared, ij} = \lambda w_i w_j$ for the off-diagonal entries (i.e., if $i \neq j$). Along the diagonal, $\Sigma_{shared, ii} = \lambda w_i^2$ and $\Sigma_{ii} = \lambda w_i^2 + \psi_i$. The correlation (i.e., $r_{sc}$ if $\Sigma$ is a spike count covariance matrix) between neurons $i$ and $j$ can be written as:

$$
\rho_{ij} = \frac{\Sigma_{ij}}{\sqrt{\Sigma_{ii} \Sigma_{jj}}} = \frac{\lambda w_i w_j}{\sqrt{\left(\lambda w_i^2 + \psi_i\right) \left(\lambda w_j^2 + \psi_j\right)}}
$$

$$
= \sqrt{\frac{\varphi_i \varphi_j \text{sign}(w_i w_j)}{\lambda w_i^2 + \psi_i}}
$$

(Equation 8)

where $\varphi_i$ and $\varphi_j$ represent the %sv (as proportions) for neurons $i$ and $j$, respectively, and $\text{sign}(w_i w_j) = +1$ if $w_i w_j > 0$ or $-1$ if $w_i w_j < 0$. The last line follows from the fact that %sv for neuron $i$ is defined in Equation 7 as:

$$
\varphi_i = \frac{\Sigma_{shared, ii}}{\Sigma_{ii}} = \frac{\lambda w_i^2}{\lambda w_i^2 + \psi_i}
$$

(Equation 9)

Equations 8 and 9 provide a basis for understanding the relationships between $r_{sc}$, %sv, and loading similarity. The $r_{sc}$ mean and SD are computed across all pairs of neurons $\rho_{ij}$, for $i < j$.

For establishing a relationship between pairwise metrics and %sv, consider decreasing the overall %sv of the population while keeping the loadings $w_i$ fixed. This corresponds to decreasing $\lambda$ in Equation 9, which implies $\varphi_j$ for each neuron decreases, and thus the product $\sqrt{\varphi_i \varphi_j}$ decreases for all pairs. The magnitude of each $\rho_{ij}$ decreases (i.e., each $\rho_{ij}$ moves closer to 0). As such, decreasing %sv of the population decreases the distance of a point from the origin in the $r_{sc}$ mean versus $r_{sc}$ SD plot, all else being equal (Figure 3F).

For establishing a relationship between pairwise metrics and loading similarity, consider two extreme cases: 1) when loading similarity is 1 (as high as possible) when 0 is 0 (as low as possible). We first assume that each neuron has the same independent variance $\psi_i$, for simplicity, as we did in Figure 3. A loading similarity of 1 corresponds to each $w_i = 1$ or each $w_i = -1$. In either case, $\text{sign}(w_i w_j)$ is always +1. Furthermore, $\psi_i$ is the same for every neuron and $\sqrt{\varphi_i \varphi_j} = %sv$ (i.e., the %sv of the population, expressed as a proportion) for every pair of neurons. Thus, all $\rho_{ij} = %sv$ for all pairs of neurons $i$ and $j$. In this case, $r_{sc}$ mean = %sv and $r_{sc}$ SD = 0. If the independent variances $\psi_i$ are different across neurons, we can still get each $\text{sign}(w_i w_j) = +1$ and each $\varphi_i$ to be the same by setting each $w_i = +\sqrt{\varphi_i}$ or each $w_i = -\sqrt{\varphi_i}$. This would also result in $\rho_{ij} = %sv$ for all pairs of neurons $i$ and $j$, and thus $r_{sc}$ mean = %sv and $r_{sc}$ SD = 0. In this case, the loading similarity is still high (all $w_i$ are the same sign; we can show that load. sim. > 0.5), but not equal to 1.

Now, consider a scenario in which half the loadings are $+\frac{1}{\sqrt{n}}$ and the other half are $-\frac{1}{\sqrt{n}}$ (and assume again that $\psi_i$ are the same for every neuron). This is one way to obtain a loading similarity of 0. In this case, $\varphi_i$ are still the same for every neuron, so $\sqrt{\varphi_i \varphi_j} = %sv$ for all pairs. However, $\text{sign}(w_i w_j) = -1$ for $\frac{n}{2}$ pairs, and $\text{sign}(w_i w_j) = +1$ for $\frac{n}{2} - \frac{1}{4}$ pairs. We can show that $r_{sc}$ mean = $-\frac{\%sv}{\sqrt{n-1}}$ and, by using Equation 10 from Math Note B below, $r_{sc}$ SD = %sv $\sqrt{\frac{1}{n-1}}$. Thus, for a large number of neurons $n$, this case (where loading similarity = 0) corresponds to small negative $r_{sc}$ mean (close to 0), and large $r_{sc}$ SD (close to the %sv). As an example, for 30 neurons and %sv = 50%, this corresponds to $r_{sc}$ mean = -0.0172 and $r_{sc}$ SD = 0.4997.

With this analysis, we have established that for one latent dimension:
Decreasing %sv decreases the magnitudes of correlations (i.e., each $\rho_{ij}$ closer to 0), $r_{sc}$ mean and SD both decrease (as seen empirically in Figure 3F).

Starting from a loading similarity near 1, a decrease in loading similarity involves flips in the signs of some correlations (i.e., some $\rho_{ij}$ become $-\rho_{ij}$). $r_{sc}$ mean decreases but $r_{sc}$ SD increases (as seen empirically in Figure 3F).

Both $r_{sc}$ mean and %sv measure shared variance among neurons, but they are not always equal. Equation 8 shows that the two quantities are equal if all sign($w_i, w_j$) are the same (i.e., when loading similarity is high). However, in general $r_{sc}$ mean and shared variance (%sv) are not the same—e.g., when loading similarity is low, or when there are multiple dimensions (Math Note C).

In this section, we consider the extremes of loading similarity. In the next section, we analyze how gradual changes in loading similarity affect $r_{sc}$ mean and SD for a fixed %sv.

B) Circular arc in $r_{sc}$ mean versus $r_{sc}$ SD plot for one latent dimension and fixed %sv

We establish here mathematically that gradually varying the loading similarity for one latent dimension and fixed %sv results in an arc-circular arc. We can write variance matrices with one latent dimension and fixed %sv. This would correspond to a point being equidistant from the origin, and thus a circular arc. Let $c$ be the distance of a point (corresponding to one instance of the population activity covariance matrix) from the origin in the $r_{sc}$ mean versus $r_{sc}$ SD plot (i.e., $c = \sqrt{(r_{sc} \text{ mean})^2 + (r_{sc} \text{ s.d.})^2}$). We want to know whether $c$ is the same for all population activity covariance matrices with one latent dimension and fixed %sv. This would correspond to a point being equidistant from the origin, and thus a circular arc. We can write $c$ as:

$$c^2 = (r_{sc} \text{ mean})^2 + (r_{sc} \text{ s.d.})^2$$

$$= E[\rho]^2 + \text{Var}(\rho)$$

$$= E[\rho]^2 + E[\rho^2] - E[\rho]^2$$

$$= E[\rho^2]$$

Thus, the squared distance (i.e., squared radius) is equal to $E[\rho^2]$, the mean of $\rho_{ij}^2$ across all pairs in the population. Let $m$ be the number of pairs (i.e., $m = \binom{n}{2}$). Now, using Equations 8 and 9 derived in Math Note A:

$$E[\rho^2] = \frac{1}{m} \sum_{i \neq j=1}^{n} \rho_{ij}^2$$

$$= \frac{1}{m} \sum_{i \neq j=1}^{n} \frac{(\lambda w_i^2)(\lambda w_j^2)}{(\lambda w_i^2 + \psi_i)(\lambda w_j^2 + \psi_j)}$$

$$= \frac{1}{m} \sum_{i \neq j=1}^{n} \frac{\phi_i \phi_j}{\psi_i + \psi_j}$$

where $\phi_i$ and $\phi_j$ are the %sv of neurons $i$ and $j$ (expressed as proportions), as defined in Math Note A. We can show that $2\sum_{i=1}^{n} \sum_{j=1}^{n} \phi_i \phi_j = \sum_{i=1}^{n} \sum_{j=1}^{n} \phi_i \phi_j - \sum_{i=1}^{n} \phi_i^2$. Intuitively, if we have a symmetric matrix $\Phi$ with entries $\Phi(i, j) = \phi_i \phi_j$, and we want to find the sum of the off-diagonal elements ($2\sum_{i=1}^{n} \sum_{j=1}^{n} \phi_i \phi_j$), then we can take the sum of all elements and subtract the diagonal elements ($\sum_{i=1}^{n} \sum_{j=1}^{n} \phi_i \phi_j - \sum_{i=1}^{n} \phi_i^2$). Using this equivalence, it follows:

$$E[\rho^2] = \frac{1}{m} \sum_{i \neq j=1}^{n} \phi_i \phi_j$$

$$= \frac{1}{2m} \left( \sum_{i=1}^{n} \sum_{j=1}^{n} \phi_i \phi_j - \sum_{i=1}^{n} \phi_i^2 \right)$$

$$= \frac{1}{2m} \left( \sum_{i=1}^{n} \phi_i \sum_{j=1}^{n} \phi_j - \sum_{i=1}^{n} \phi_i^2 \right)$$

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This provides an equation for the squared radius (i.e., squared distance from the origin) of a point in the loading vector values. In the above derivation, $E[\phi]$ and $\text{Var}(\phi)$ are taken across the percent shared variance of each neuron in the population $\phi_i$. Thus, $E[\phi]$ is equal to our population metric $\%sv$. Now, we will bound $\text{Var}(\phi)$, which by definition is greater than or equal to 0. Since $0 \leq \phi_i \leq 1$, one instance where the maximum variance occurs is when there are an equal number of $\phi_i = 0$ and $\phi_i = 1$ (and $E[\phi] = 0.5$). Then,

$$\text{Var}(\phi) = \frac{1}{n} \sum_{i=1}^{n} (\phi_i - 0.5)^2 = \frac{1}{n} \left( \frac{n}{2} \right) (1 - 0.5)^2 + \frac{n}{2} (0 - 0.5)^2 = \frac{1}{n} (0.25n) = 0.25$$

So $0 \leq \text{Var}(\phi) \leq 0.25$. For a small number of neurons $n$, the second term in Equation 10 is non-negligible. For example, for a model with 6 neurons and $\%sv = 50\%$, the radius of the data points may vary between 0.4472 and 0.5. As the number of neurons increases, the second terms becomes negligible, and data points lie approximately along an arc with radius equal to $\%sv$. For example, for 30 neurons as in our simulations and a $\%sv$ of 50%, the radius of the data points may vary between 0.4913 and 0.5. As the number of neurons increases, the radius becomes asymptotically closer to $\%sv$.

To summarize, Equation 10 computes the distance from the origin of a point for a given population of neurons. For a fixed $\%sv$, $\text{Var}(\phi)$ can be the same or differ across many simulation runs. If $\text{Var}(\phi) = 0$ or is the same across runs, then the points will lie perfectly along an arc, with radius specified by Equation 10. However, if $\text{Var}(\phi)$ is different across runs, the distances of each point from the origin will differ slightly, so they will lie close to, but not exactly along, an arc.

With this analysis, we have shown that in the case of one latent dimensions:

- A point (i.e., corresponding to a given population of neurons, simulated or real) on the $r_{sc}$ mean versus $r_{sc}$ SD plot has distance from the origin (i.e., radius) less than or equal to $\%sv$.
- If the $\%sv$ for individual neurons (\(\phi_i\)) are all the same (see Math Note A), then the radius equals $\%sv$.
- As the number of neurons increases, the radius becomes asymptotically closer to $\%sv$.

### C) Relationship between correlation, loading similarity, and $\%sv$ (multiple latent dimensions)

In Math Note A, we established a mathematical relationship between $r_{sc}$, loading similarity, and $\%sv$ in the case of one latent dimension. Here, we generalize Equation 8 to include multiple dimensions in order to better understand the relationship between $r_{sc}$ and dimensionality. We demonstrate here that the general relationships between $r_{sc}$, $\%sv$, and loading similarity for one latent dimension also hold true for multiple latent dimensions. For multiple latent dimensions, the relative strengths of each dimension is an important consideration—a stronger dimension plays a bigger role in determining the $r_{sc}$ distribution. Finally, we consider the relationship between dimensionality itself and $r_{sc}$. We will discover below that increasing dimensionality tends to decrease the magnitude of $r_{sc}$ values.

First, consider the case of two latent dimensions. Again, let $n$ be the number of neurons, let $w$ be the co-fluctuation pattern (i.e., loading vector $[v_1, v_2, ..., v_n]^T \in \mathbb{R}^{n \times 1}$) with eigenvalue $\lambda_{sv}$, let $v$ be another pattern orthogonal to $w$ ($[v_1, v_2, ..., v_n]^T \in \mathbb{R}^{n \times 1}$; $v \perp w$), with
eigenvalue $\lambda_v$, and let $\Psi \in \mathbb{R}^{n \times n}$ be a diagonal matrix specifying the independent variance of each neuron ($\psi_1, \psi_2, \ldots, \psi_d$). Then the covariance is $\Sigma = \Sigma_{\text{shared}} + \Psi = \Sigma_w + \Sigma_v + \Psi = \mathbf{w}\mathbf{w}^T + \mathbf{v}\mathbf{v}^T + \Psi$. On the off-diagonals entries (i.e., if $i \neq j$), $\Sigma_{ij} = \lambda_w w_i w_j + \lambda_v v_i v_j$. Along the diagonal, $\Sigma_{ii} = \lambda_w w_i^2 + \lambda_v v_i^2 + \psi_i$.

Because the shared covariance matrix $\Sigma_{\text{shared}}$ can be expressed as a sum of two component matrices $\Sigma_w + \Sigma_v$, we can express the %sv of neuron $i$ ($\phi_i$) as

$$\phi_i = \frac{\Sigma_{\text{shared} ii}}{\Sigma_{ii}} = \frac{\Sigma_w ii + \Sigma_v ii}{\Sigma_{ii}} = \frac{\lambda_w w_i^2 + \lambda_v v_i^2}{\lambda_w w_i^2 + \lambda_v v_i^2 + \psi_i} = \phi_i^{(w)} + \phi_i^{(v)}$$

where $\phi_i^{(w)}$ is the %sv variance of neuron $i$ explained by dimension $w$ and $\phi_i^{(v)}$ is the %sv variance of neuron $i$ explained by dimension $v$.

With this decomposition of $\psi_i$, and following similar steps as in Equation 8:

$$\rho_i = \sqrt{\phi_i^{(w)} \phi_i^{(w)} \text{sign}(w_i w_j) + \phi_i^{(v)} \phi_i^{(v)} \text{sign}(v_i v_j)}$$

(Equation 11)

where %sv values ($\phi$) are represented as proportions. Equation 11 relates $r_{sc}$, %sv, and loading similarity for the case of two latent dimensions. Next, we compare these relationships for one versus two latent dimensions.

We will show that, for two latent dimensions, the relative strength of each dimension (i.e., the ratio $\lambda_w : \lambda_v$) is an important consideration. For two latent dimensions, decreasing the overall %sv by decreasing both $\phi^{(w)}$ and $\phi^{(v)}$ equally (e.g., $\lambda_w = \lambda_v$ and both decrease equally) pushes each $\rho_{ij}$ closer to 0; $r_{sc}$ mean and SD will decrease. This is similar to what happens for one latent dimension when %sv is decreased. On the other hand, even if the overall %sv is held constant, but $\phi^{(w)}$ increases relative to $\phi^{(v)}$ (i.e., increase the strength of $\mathbf{w}$ relative to $\mathbf{v}$), pairwise correlations could change. Each $\rho_{ij}$ will largely be determined by $\phi^{(w)}$ and $\mathbf{w}$; $r_{sc}$ mean and SD will be more similar to what they would be if only $\mathbf{w}$ existed (Figure 4A). In other words, each $\rho_{ij}$ for two latent dimensions is the sum of the $\rho_{ij}$ that would have been produced by each of the two constituent dimensions on their own. The dimension with larger relative strength $\lambda$ will have larger $\phi$; the stronger dimension will play a larger role in determining each value of $\rho_{ij}$ and thus the resulting $r_{sc}$ distribution.

Using this logic, we can deduce that increasing the loading similarity of one of the dimensions would increase $r_{sc}$ mean and decrease $r_{sc}$ SD for the same reasons as for one latent dimension (Math Note A). Doing so for a relatively stronger dimension would result in larger changes in $r_{sc}$ than doing so for a relatively weaker dimension.

We have shown how having multiple latent dimensions can affect the relationship between $r_{sc}$, %sv, and loading similarity. Now, we show that dimensionality itself and $r_{sc}$ are related—a larger dimensionality tends to decrease $r_{sc}$ mean and SD. To see this, we can generalize Equation 11 for $d < n$ orthogonal latent dimensions $\mathbf{u}_1, \ldots, \mathbf{u}_d \in \mathbb{R}^n$.

$$\rho_i = \sqrt{\sum_{k=1}^d \phi_i^{(u_k)} \phi_i^{(u_k)} \text{sign}(u_{ki} u_{kj})}$$

Considering the sign of one term, $\rho_i$ could have the same sign for $\text{sign}(u_{ki} u_{kj})$ across all dimensions $\mathbf{u}_1, \ldots, \mathbf{u}_d$; in this case, a larger dimensionality acts to increase the correlation between neurons $i$ and $j$ ($\rho_{ij}$) above the level corresponding to a single dimension. However, because the loading vectors $\mathbf{u}_1, \ldots, \mathbf{u}_d$ are orthogonal, a pair of neurons $i$ and $j$ is likely to have many sign($u_{ki} u_{kj}$) of opposite sign across dimensions; in this case, a larger dimensionality pushes the correlation between neurons $i$ and $j$ ($\rho_{ij}$) closer to 0. Thus, we would expect the magnitude of correlations to decrease as more dimensions are added (i.e., a tendency for $r_{sc}$ mean and SD to decrease; Figure 3G). In the next section, we show this relationship mathematically.

**D) Increasing dimensionality decreases arc radius**

We establish here that increasing dimensionality results in a decrease in the radius of the arc in the $r_{sc}$ mean versus $r_{sc}$ SD plot (Figure 3G). We extend the math for an arc for one latent dimension (Math Note B) to multiple latent dimensions. We will refer to the one latent dimension as the ‘1-d case’ and multiple ($k$) latent dimensions as the ‘$k$-d case’.

We use the same notation as in Math Note C. Consider the distance $c$ of a point (corresponding to one instance of the population activity covariance matrix) from the origin in the $r_{sc}$ mean versus $r_{sc}$ SD plot. From Math Note B, $c^2 = E[l^2]$. For this 2-d case, the correlation between neurons $i$ and $j$ is $\rho_{ij} = \frac{\Sigma_{ij}}{\sqrt{\Sigma_{ii} \Sigma_{jj}}} = \frac{\lambda_w w_i w_j + \lambda_v v_i v_j}{\sqrt{(\lambda_w w_i^2 + \lambda_v v_i^2 + \psi_i)(\lambda_w w_j^2 + \lambda_v v_j^2 + \psi_j)}}$. Thus we can write $\rho_{ij}^2$ as:

$$\rho_{ij}^2 = \frac{(\lambda_w w_i w_j + \lambda_v v_i v_j)^2}{(\lambda_w w_i^2 + \lambda_v v_i^2 + \psi_i)(\lambda_w w_j^2 + \lambda_v v_j^2 + \psi_j)}$$
where the % shared variance of neuron \( i \) in this 2-d case is

\[
\rho_i^2 = \frac{(\lambda_w w_i^j + \lambda_v v_i^j)^2}{(\lambda_w w_i^j + \lambda_v v_i^j)^2} = \frac{\lambda_w^2 w_i^j + 2 \lambda_w \lambda_v w_i^j v_i^j + \lambda_v^2 v_i^j}{(\lambda_w^2 w_i^j + \lambda_v^2 v_i^j + \psi_i^j)}
\]

Then letting \( m \) be the number of pairs in the population, and following similar steps to Equation 10 in Math Note B, we arrive at:

\[
E[\rho^2] = \frac{1}{m} \sum_{i,j=1}^{n} \sum_{i,j=1}^{n} \rho_i^2 = \frac{1}{n-1} \text{Var}(\phi) - \frac{1}{m} \sum_{i,j=1}^{n} \sum_{i,j=1}^{n} \lambda_w \lambda_v (w_i^j - v_i^j)^2 (\lambda_w^2 w_i^j + \lambda_v^2 v_i^j + \psi_i^j)
\]

(Equation 12)

Not including the negative sign in front, note that this final term is non-negative (given that \( \lambda_w \) and \( \lambda_v \) are non-negative, as for any covariance matrix). Thus, comparing the final line in Equation 12 to the final line from Equation 10, we observe that the distance of the point for the 2-d case in the \( r_{sc} \) mean versus \( r_{sc} \) SD plot is necessarily smaller than or equal to the distance for the corresponding 1-d case.

More generally, for a \( k \)-dimensional case we can show that:

\[
E[\rho^2] = (\%sv)^2 - \frac{1}{n-1} \text{Var}(\phi) - \frac{1}{m} \sum_{w,v} \sum_{i,j=1}^{n} \sum_{i,j=1}^{n} \lambda_w \lambda_v (w_i^j - v_i^j)^2 (\lambda_w^2 w_i^j + \lambda_v^2 v_i^j + \psi_i^j)
\]

(Equation 13)

where the sum \( \sum \) is taken over all unique pairs of loading vectors \( (w, v) \). Indeed, as more latent dimensions are subsequently added, the radius of the \( r_{sc} \) mean versus \( r_{sc} \) SD plot decreases (Figure 3G). Intuitively, this final term accounts for how population activity covaries along many different dimensions in the high-d firing rate space. As more orthogonal dimensions are added, population activity is further pulled in different directions in the high-d space, more interaction terms come into play, and the magnitude of correlations is further decreased. This tends to decrease both \( r_{sc} \) mean and \( r_{sc} \) SD, explaining why the radius of the arc in the \( r_{sc} \) mean versus \( r_{sc} \) SD plot tends to decrease as dimensionality increases.

We note that \( r_{sc} \) mean and \( r_{sc} \) SD do not necessarily both need to decrease. For example, consider a pattern with a loading similarity of 1; loading weights for all neurons would have the same value, \( r_{sc} \) across all pairs would be the same value, and thus \( r_{sc} \) SD would be 0 (see Math Note A). When a second pattern of necessarily low loading similarity (see Math Note E) is added, \( r_{sc} \) values across pairs of neurons would differ, and \( r_{sc} \) S.D. would be larger than 0. Therefore, \( r_{sc} \) SD can increase when going from the 1-d case to the 2-d case. However, the corresponding decrease in \( r_{sc} \) mean would be larger in magnitude than the increase in \( r_{sc} \) SD, resulting in an overall decrease in arc radius (Figure 3G, 1 to 2 dimensions, data points closest to the horizontal axis).

The third term in Equation 13 can also help explain variability of the radius \( (E[\rho^2]) \) across different random instantiations with the same population metrics (Figures 3G and 4). Consider a fixed \( \%sv \). For the 1-d case, the radius is determined by the first two terms of the above equation, and any variability in radius will be caused by different values of \( \text{Var}(\phi) \) across different instantiations. For the 2-d case, the third term also plays a factor in determining the radius, and this term varies across different random instantiations, typically to a larger degree than the second term for large numbers of neurons \( n \) (see Math Note B). Thus, the 2-d and \( k \)-d cases have greater variability in \( E[\rho^2] \) than 1-d cases (Figures 3G and 4). Other subtle factors can affect the variability of \( E[\rho^2] \). For example, variability in \( E[\rho^2] \) can increase or decrease depending on the relative strengths of each dimension and their corresponding loading similarities (Figures 4 and S1). This can be explained by the third component of Equation 13, in particular by the terms involving \( \lambda_w \) and \( \lambda_v \).

**E) Properties of loading similarities across different co-fluctuation patterns**

We asked whether there was a relationship between the loading similarities of different co-fluctuation patterns in the same model. In our simulations and V4 data analysis, we ensured that we obtain unique co-fluctuation patterns by constraining dimensions to be orthogonal. Thus, we might conjecture that if one pattern has high loading similarity (e.g., \( [1, \ldots, 1] \)), then another pattern in the same model necessarily has low loading similarity (e.g., \( [1, -1, 1, \ldots, -1, 1] \)). Indeed, this is true because the sum across the loading similarities of each pattern in a model is at most 1. We show this property of loading similarity here.

Let \( w \) and \( v \) be vectors representing two co-fluctuation patterns in the same model. We use the notation \( w \cdot v \) to refer to the element-wise product between \( w \) and \( v \), resulting in a vector that is the same size as \( w \) and \( v \). Furthermore, we use \( E[w_j], \text{Var}(w_j), \text{Cov}(w) \)
as shorthand to refer to computations across the elements of a vector (and not as operations on a random variable): e.g., \(E[w] = \frac{1}{n} \sum_{i=1}^{n} w_i\), and \(\text{Cov}(w, v) = E[wv] - E[w]E[v] = \frac{1}{n} \sum_{i=1}^{n} wi - \left(\frac{1}{n} \sum_{i=1}^{n} wi\right) \left(\frac{1}{n} \sum_{i=1}^{n} vi\right)\). Also, in this section we refer to the loading similarity of vector \(w\) as \(ls(w)\) for shorthand.

We first show a constraint on loading similarities for a model with two co-fluctuation patterns (i.e., loading vectors for each dimension). Let \(n\) be the number of neurons and let \(w, v \in \mathbb{R}^d\) be two loading vectors. As in our simulations and data analysis (see Methods), \(w\) and \(v\) are orthogonal unit vectors: \(\sum_{i=1}^{d} w_i^2 = 1\), \(\sum_{i=1}^{d} v_i^2 = 1\), and \(\sum_{i=1}^{d} wi v_i = 0\). Then, using these constraints,

\[
\text{Cov}(w, v) = E[wv] - E[w]E[v] \\
= \frac{1}{n} \sum_{i=1}^{n} wi - \frac{1}{n} \sum_{i=1}^{n} wi \frac{1}{n} \sum_{i=1}^{n} vi \\
= -E[w]E[v] \\
\text{Var}(w) = E[w^2] - E[w]^2 \\
= \frac{1}{n} \sum_{i=1}^{n} w_i^2 - \frac{1}{n} \left(\frac{1}{n} \sum_{i=1}^{n} w_i\right)^2 \\
= \frac{1}{n} \sum_{i=1}^{n} w_i^2 - \frac{1}{n} \left(\frac{1}{n} \sum_{i=1}^{n} w_i\right)^2
\]

(Equation 14)

Because correlation is bounded between \(-1\) and \(1\), we know that \(|\text{Cov}(w, v)| \leq \sqrt{\text{Var}(w) \text{Var}(v)}\). It follows that:

\[
\text{Cov}^2(w, v) \leq \text{Var}(w) \text{Var}(v) \\
E[w^2]E[v^2] \leq \left(\frac{1}{n} \sum_{i=1}^{n} w_i^2\right) \left(\frac{1}{n} \sum_{i=1}^{n} v_i^2\right) \\
0 \leq \frac{1}{n^2} \sum_{i=1}^{n} wi v_i \leq \frac{1}{n} \sum_{i=1}^{n} w_i^2 + \frac{1}{n} \sum_{i=1}^{n} v_i^2 \\
\text{Var}(w) \leq \frac{nE[w^2]}{n} + \frac{nE[v^2]}{n} \\
ls(w) + ls(v) \leq 1
\]

(Equation 15)

The last step follows from the definition of loading similarity:

\[
ls(w) \equiv 1 - \frac{\text{Var}(w)}{\frac{1}{n} \sum_{i=1}^{n} wi} = 1 - \frac{\frac{1}{n} \sum_{i=1}^{n} wi^2}{\frac{1}{n} \sum_{i=1}^{n} wi} = \frac{nE[w^2]}{n}
\]

The final inequality in Equation 15 proves the intuition provided at the beginning of this section— if \(ls(w)\) is large, then \(ls(v)\) must be small (at most \(1 - ls(w)\)). More strongly, if \(ls(w) = 1\), then \(ls(v) = 0\).

Generally, for a model with \(d\) dimensions and patterns \(u_1, \ldots, u_d \in \mathbb{R}^d\), we can show that \(\sum_{i=1}^{d} ls(u_i) \leq 1\). To see this, we can construct a matrix \(C\) with entries \(c_{ij} = \text{Cov}(u_i, u_j) = -E[u_iu_j] \text{ for } i \neq j\), and \(c_{ii} = \text{Var}(u_i) = \frac{1}{n} - E[u_i]^2\). Note that \(C \in \mathbb{R}^{d \times d}\), with variances on the diagonal and covariances off-diagonals, is a covariance matrix, which implies \(\text{det}(C) \geq 0\). For a 3-d model,

\[
\text{det}(C) = \frac{1}{n^3} \left(1 - nE[u_1]^2 - nE[u_2]^2 - nE[u_3]^2\right) \geq 0
\]

which implies \(ls(u_1) + ls(u_2) + ls(u_3) \leq 1\). In general, for a \(d\)-dimensional model (with \(d \leq n\)):

\[
\text{det}(C) = \frac{1}{n^d} \left(1 - \sum_{i=1}^{d} nE[u_i]^2\right) \geq 0
\]

(Equation 16)

Equation 16 has several implications:

- If one knows the loading similarities of all dimensions \(u_1, \ldots, u_d\) in a model, then the maximum possible loading similarity of any new dimension is \(1 - \sum_{i=1}^{d} ls(u_i)\). It follows that two dimensions with high loading similarity cannot co-exist in the same model.
- If one dimension has \(ls = 1\), then all other dimensions in the model (or that would be added to the model) necessarily have \(ls = 0\). Note that there is only one possibility for a pattern to have \(ls = 1\) (i.e., \(u = \left[\frac{1}{\sqrt{n}}, \ldots, \frac{1}{\sqrt{n}}\right]^T\), such that \(\text{Var}(u) = 0\)). This implies that there are many possibilities for a pattern to have \(ls(u) = 0\). More loosely, there are relatively few ways for a pattern to have high loading similarity, but many more ways for a pattern to have low loading similarity.
**F) Maximum variance of a unit vector**

We defined loading similarity for a co-fluctuation pattern \( u \) (normalized to have norm 1) of \( n \) neurons to be \( 1 - \frac{\text{Var}(u)}{n^2} \), where the variance is computed along the elements of \( u \). This value lies between 0 and 1 because the maximum variance across the elements of \( u \) is \( 1/n \). We now show this mathematically.

Let \( u \in \mathbb{R}^n \) be a unit vector. Because \( u \) is a unit vector, \( \sum_{i=1}^{n} u_i^2 = 1 \). Using these facts:

\[
\text{Var}(u) = E[u^2] - E[u]^2
= \frac{1}{n} \sum_{i=1}^{n} u_i^2 - E[u]^2
= \frac{1}{n} - E[u]^2
\leq \frac{1}{n}
\]

This holds with equality when \( E[u] = 0 \) (i.e., when the mean across the elements in a co-fluctuation pattern is 0). This implies that the smallest loading similarity is 0 (when \( \text{Var}(u) = 1/n \)), and the largest loading similarity is 1 (when \( \text{Var}(u) = 0 \)).
Supplemental information

Bridging neuronal correlations
and dimensionality reduction

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Supplementary Information

Supplementary Figure 1: Relationship between pairwise metrics, loading similarity of each latent dimension, and the relative strengths of each dimension. Related to Figure 4.

In Fig. 3e and Math Note A, we considered the relationship between loading similarity and pairwise metrics when population activity was one dimensional. Here, we asked about the informativeness of loading similarity when population activity varies along multiple dimensions, and the impact of the relative strengths of each dimension (i.e., the shape of the eigenspectrum of $\Sigma_{shared}$, which specifies the amount of shared variance explained by each dimension).

We considered two cases. First, we considered an eigenspectrum that decays quickly, as has been widely reported in population recordings (Sadtdler et al., 2014; Williamson et al., 2016; Mazzucato et al., 2016; Gallego et al., 2018; Huang et al., 2019; Stringer et al., 2019a; Ruff et al., 2020). In this case, we found that the loading similarity of the strongest dimension (i.e., dimension with largest eigenvalue) was most informative about pairwise metrics, while the loading similarities of the other dimensions were less informative. Second, we considered a flat eigenspectrum. In this case, the loading similarities of each dimension were equally informative.

a. Loading similarity for a decaying eigenspectrum of the shared covariance matrix ($\Sigma_{shared}$ in Supplementary Fig. 5a). We reproduced the simulation in Fig. 3 for a latent dimensionality of 3 and $\%sv=50\%$. For each 3-d model, we evaluated the $r_{sc}$ mean and s.d., and then plotted the same point in 3 separate panels colored by loading similarity of each of the 3 different dimensions. The loading similarity of strongest dimension (‘Dim 1’) is very informative—high loading similarity implies high $r_{sc}$ mean and low $r_{sc}$ s.d. (green dots), whereas low loading similarity implies low $r_{sc}$ mean and high $r_{sc}$ s.d. (blue dots). This is the same relationship as shown in Fig. 3e for the case of one dimension. The loading similarities of ‘Dim 2’ and ‘Dim 3’ are less informative—in both cases, low loading similarity points (blue dots) are scattered throughout the arc. The only case when the loading similarity of ‘Dim 2’ or ‘Dim 3’ is informative is when either of them have a high loading similarity (green dots). This is informative because it implies that ‘Dim 1’ must have low loading similarity (‘Dim 1’ is blue for dots where ‘Dim 3’ is green; see Math Note E), implying low $r_{sc}$ mean and high $r_{sc}$ s.d. (continued on next page...)
b. Same as panel a but for flat eigenspectrum across the three dimensions. In this case, $r_{sc}$ mean will tend to be small and $r_{sc}$ s.d. will tend to be large because: 1) all three dimensions contribute equally, and 2) it is not possible for all three dimensions to have high loading similarity, while multiple dimensions can have low loading similarity (Math Note E). However, knowing whether any of the three dimensions have high loading similarity can provide more specific information about $r_{sc}$ mean and s.d. within this limited range (green dots tend to have high $r_{sc}$ mean and lower $r_{sc}$ s.d. in each panel).

Because most studies of population neuronal recordings have shown quickly decaying eigenspectra as in panel a (Sadatler et al., 2014; Williamson et al., 2016; Mazzucato et al., 2016; Gallego et al., 2018; Huang et al., 2019; Stringer et al., 2019; Ruff et al., 2020), we recommend considering the loading similarity of the strongest dimension for concision and simplicity (as we do in Fig. 6c; and see eigenspectra in Supplementary Fig. 2). However, if it happens that the data have an eigenspectrum that decays slowly or has multiple dimensions that are very strong, then one may benefit by considering the loading similarities of additional dimensions as well.

This analysis also highlights how the shape of the eigenspectrum influences pairwise metrics. First, an exponentially-decaying eigenspectrum tended to have a higher $r_{sc}$ mean and s.d. compared to its corresponding flat eigenspectrum (dots in panel a are farther from origin here than in panel a). This occurs because, for an exponentially-decaying eigenspectrum, an added dimension explains relatively little shared variance. Thus, the added dimension tends to result in only a small decrease in $r_{sc}$ mean and s.d. On the other hand, adding a dimension to the flat eigenspectrum affects $r_{sc}$ mean and s.d. as much as any other dimension, leading to larger changes (i.e., decreases) in $r_{sc}$ mean and s.d. than in the case of an exponentially-decaying eigenspectrum.

Second, we observed a greater radial and angular spread for exponentially-decaying eigenspectra (panel a) compared to flat eigenspectra (panel b). This occurs because, when the eigenspectra are not flat, there is greater diversity in how the co-fluctuation patterns of different dimensions can contribute to $r_{sc}$. In other words, permuting the eigenvectors of three dimensions with equal eigenvalues (i.e., both dimensions explain the same amount of shared variance) results in the same model and same covariance matrix—yielding the same values for $r_{sc}$ mean and s.d. However, permuting the eigenvectors of three dimensions with different eigenvalues will likely result in a different covariance matrix and different values of $r_{sc}$ mean and s.d. Thus, for non-flat eigenspectra, the greater diversity by which co-fluctuation patterns can contribute to the shared covariance matrix leads to greater spread in the $r_{sc}$ mean vs s.d. plots. The mathematical details regarding this observation are provided in Math Note D.

An implication of this analysis is that it is important to report the eigenspectrum shape whenever one reports dimensionality. Thus, considering both dimensionality and the eigenspectrum curve, instead of dimensionality alone, will lead to a more complete picture of the structure of population activity. Inspecting the eigenspectrum will also help determine whether assessing loading similarity in the strongest dimension is sufficient (panel a), or whether one needs to consider the loading similarities of other dimensions as well (panel b).
Supplementary Figure 2: Eigenvalues and loading similarity by dimension for V4 population activity. Related to Figure 6.

Although we observed only a modest change in dimensionality with attention (Fig. 6c), our simulations showed that the relative strength of each dimension (i.e., shape of the shared eigenspectrum) could alter the “effective dimensionality” of population activity and have large effects on pairwise metrics (Fig. 4a). Here, we asked whether the relative strengths of each dimension changed with attention. We also considered the loading similarities across different dimensions.

a. We found that the shape of the eigenspectra was qualitatively similar for ‘attend in’ and ‘attend out’ conditions (red and black curves have similar shape). In both conditions, the eigenvalues of the shared covariance matrix decayed (dot for each subsequent dimension was below dot for the previous dimension), indicating that a small number of dimensions were needed to explain the population-wide covariability.

When comparing eigenspectra (i.e., the amount of shared variance explained by each dimension), one also needs to consider the firing rates under each condition. Mean firing rates tend to be higher for attend-in than attend-out trials. Higher firing rates typically correspond to higher spike count variance due to the Poisson-like firing of neurons. All else being equal, the higher mean firing rates imply higher levels of both shared variance and independent variance (Churchland et al., 2010). Thus, a direct comparison of the eigenspectra should be done with caution. Nonetheless, we plotted attend-in and attend-out together to relate our results to previous reports (Huang et al., 2019; Ruff et al., 2020). Consistent with these studies, we found that attention decreased the strength of the strongest dimension (red below black dot for dimension index 1), though the magnitude of the decrease we observed was more consistent with Ruff et al. (2020) than Huang et al. (2019). Had we been able to equalize the mean firing rate across the two conditions, we likely would have observed an even greater difference between attend-in and attend-out. We note that the caveat described here for comparison of eigenspectra (i.e., the amount of shared variance) does not apply to comparisons of %sv (Fig. 6c) because %sv is normalized by the overall spike count variance.

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Supplementary Figure 2: (...continued from previous page)

The eigenspectra were computed in the following way. We decomposed the V4 spike count covariance matrix into shared and independent components using factor analysis (see Methods). We then computed the eigendecomposition of the shared covariance matrix (Supplementary Fig. 5, $\Sigma_{\text{shared}} = U\Lambda U$). We found that eigenvalues (diagonal of $\Lambda$) tended to increase linearly with the number of neurons recorded; therefore, in order to combine across sessions, we normalized the eigenvalues by dividing by the number of neurons recorded in each session. After normalizing, we computed the eigenspectrum averaged across sessions and stimulus orientations. Because the dimensionality identified by cross-validation differed across sessions, there were a different number of sessions that contributed to each average. We did not plot mean eigenvalues when there were fewer than 5 sessions to average (i.e., dimensions $\geq 6$ for monkey 1; dimensions $\geq 4$ for monkey 2). Error bars indicate standard error. Data points have been jittered horizontally for visual clarity.

b. Loading similarity for ‘attend-in’ (red) and ‘attend-out’ (black) by dimension. Pooled across monkeys, the loading similarity for the first (i.e., strongest) dimension was larger for ‘attend-out’ than ‘attend-in’ (same result as Fig. 6c). We also observed some differences in loading similarity across the other dimensions in both monkeys. These differences could be important for specific scientific questions (see Fig. 7, for example). However, as we show in Fig. 4, Supplementary Fig. 1, and Math Note C, the first dimension plays the largest role in determining the $r_{sc}$ distribution because it explains the greatest amount of shared variance.
Supplementary Figure 3: Quantifying the extent to which each population metric contributes to changes in pairwise metrics. Related to Figure 6.

In Fig. 6c, we observed changes in several population metrics with attention in V4 population responses. However, it was unclear to what degree the change we observed in each population metric contributed to the overall changes in pairwise metrics (Fig. 6b). In order to quantify this, here we used a population metric matching procedure to assess how much each individual change in a population metric contributed to the changes in a pairwise metric. We found that for these V4 data, %sv contributes the most, followed by loading similarity, and finally dimensionality. We illustrate these results in Fig. 6d (also reproduced here as panel a for convenience).

a. Reproduction of Fig. 6d to aid the interpretation of panels b and c here. For pairwise metrics, we observed decreases in both rs_c mean and s.d. with attention. For population metrics, we observed decreases in %sv, loading similarity, and dimensionality with attention.

b. Contribution of population metrics to changes in rs_c mean. For each recording session, we assessed how allowing all population metrics to vary (“all”) or only a single population metric to vary between “attend-out” (unatt.) and “attend-in” (att.) influenced rs_c mean. The procedure for assessing this is detailed at the end of the caption (“Details of population metric matching procedure”). When only %sv or only loading similarity were allowed to vary, rs_c mean decreased with attention; when only dimensionality was allowed to vary, rs_c mean increased. When all population metrics were allowed to vary, rs_c mean decreased, consistent computations directly from data (Fig. 6b). Results for both monkeys were consistent; means and standard errors across sessions are shown.

(continued on next page...)
Supplementary Figure 3:

c. Contribution of population metrics to changes in $r_{sc}$ s.d. Same format as b. When only $\%sv$ was allowed to vary, $r_{sc}$ s.d. decreased with attention. When only loading similarity was allowed to vary, $r_{sc}$ s.d. slightly increased (not significant). Also, when only dimensionality could vary, $r_{sc}$ s.d. increased. When all population metrics were allowed to vary, we found that $r_{sc}$ s.d. decreased with attention, consistent with our computations from data (Fig. 6b).

These results provide a systematic quantification of the illustration that relates pairwise and population metrics in V4 (panel a). Based on direction and magnitude of contributions, we conclude that for overall changes in pairwise metrics in these data: 1) $\%sv$ is most important, 2) followed by loading similarity, 3) followed by dimensionality. More generally, the population metric matching procedure (described below) provides a framework for assessing how changes in population metrics contribute to changes in pairwise metrics in recorded neuronal population activity.

Details of population metric matching procedure. Given two factor analysis (FA) models (e.g., fitted to two different experimental conditions), we first assess the overall change in pairwise metrics by computing $r_{sc}$ mean and s.d. directly from the two fitted models (see Methods). In this case, all three population metrics are allowed to change between the two conditions, and contribute to the overall observed change in pairwise metrics (labeled “all” in the plots above).

Next, we use population metric matching to assess the contribution of each individual population metric change to the overall change in pairwise metrics. To do so, we choose one of the two fitted FA models (e.g., the model fitted to “attend-out”) and systematically change the model such that one of its population metrics matches that of the other FA model (e.g., “attend-in”), while the other two population metrics remain the same. We then assessed the change in pairwise metrics between the base FA model (i.e., “attend-out”) and the “matched” FA model (i.e., modified “attend-out” model). This allowed us to assess the change in pairwise metrics that would have resulted from a change in a single population metric.

For systematically modifying $\%sv$, we scaled the eigenspectrum (see Methods) of the base FA model in order to match the $\%sv$ of the other FA model. For systematically modifying loading similarity, we replaced the co-fluctuation patterns ($U$ in Supplementary Fig. 5a) in the base FA model (e.g., “attend-out”) with the co-fluctuation patterns from the other FA model (e.g., “attend-in”). In cases where the dimensionality of the two models was different, we swapped the top $k$ co-fluctuation patterns, where $k$ is equal to the smaller dimensionality in the two models. For systematically modifying dimensionality, we removed dimensions from the base FA model if it had higher dimensionality than the other FA model, or added dimensions (after orthogonalization) from the other model to the base FA model if it had lower dimensionality. Because adding or removing dimensions changes the $\%sv$, we then scaled the eigenspectrum to match the original $\%sv$ of the base FA model. These procedures allowed us, using two FA models fit to real data, to systematically vary one of the population metrics while keeping the other two the same and assess the contribution to a change in pairwise metrics.
Supplementary Figure 4: Relationship between pairwise and population metrics in V1 population responses. Related to Figure 6.

In Fig. 6, we assessed the relationships between pairwise and population metrics in V4 population recordings where a decrease in $r_{sc}$ mean with spatial attention had been widely reported (Cohen & Maunsell, 2009; Mitchell et al., 2009; Gregoriou et al., 2014; Luo & Maunsell, 2015; Snyder et al., 2018). To demonstrate the applicability of the identified relationships to other brain areas, we applied the same analysis to population recordings in primary visual cortex (V1). Previous studies have shown that the $r_{sc}$ mean is lower after stimulus onset (i.e., evoked activity) than before stimulus onset (i.e., spontaneous activity) in V1 (Smith & Kohn, 2008; Churchland et al., 2010). Here, we analyzed population activity recorded using Utah arrays in V1 (88 to 159 units per session, 112.2 on average) in three macaque monkeys (previously reported in Zandvakili & Kohn, 2015; Semedo et al., 2019, http://dx.doi.org/10.6080/K0B27SHN). Two monkeys had two recording sessions each, while the third monkey had a single recording session, for a total of 5 recording sessions. Animals were presented with 1.28s of oriented gratings (1 of 8 possible orientations) interleaved with 1.5s of a blank screen.

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In V1, evoked activity had smaller $r_{sc}$ mean and s.d. than spontaneous activity. We also found that evoked activity had smaller $%sv$ and loading similarity, but larger dimensionality than spontaneous activity. Most changes in pairwise and population metrics between V1 “spontaneous” and “evoked” activity matched the direction of the changes in V4 “attend-out” and “attend-in”, except for the change in dimensionality (cf. panel c and Fig. 6d). Taken together, our analyses of V1 and V4 population activity demonstrate that similar changes in pairwise metrics need not correspond to precisely the same changes in population metrics. In this case, measuring population metrics provided insight about the dimensionality of the population-wide variability that would not have been gleaned from changes in pairwise metrics alone.

a. The $r_{sc}$ mean was smaller in evoked activity than in spontaneous activity (left panel; $p < 0.0001$) (Smith & Kohn, 2008; Churchland et al., 2010). We also found that $r_{sc}$ s.d. was smaller in evoked activity than in spontaneous activity (right panel; $p < 0.0001$), which has not been previously reported.

b. Next, we assessed how population metrics changed between evoked and spontaneous V1 activity. Consistent with (Churchland et al., 2010), we found that $%sv$ was smaller for evoked activity than spontaneous activity (left panel; $p < 0.0001$). We also found that loading similarity for the dominant dimension was smaller for evoked activity than spontaneous activity (middle panel; $p < 0.0001$). Finally, we found that dimensionality was higher for evoked activity than spontaneous activity (right panel; $p < 0.01$). This result differed from a previous study in which dimensionality was lower for evoked activity than spontaneous in neural recordings from rat gustatory cortex and in a clustered network model (Mazzucato et al., 2016). This could be explained by a difference in sensory modality or the way in which dimensionality was measured.

c. Using the framework we developed to understand the relationships between pairwise and population metrics (Fig. 5), the decrease in both $r_{sc}$ mean and s.d. with evoked V1 activity corresponds to: 1) a decrease in $%sv$, 2) a decrease in loading similarity, and 3) an increase in dimensionality. The direction of the changes in pairwise metrics between spontaneous and evoked activity are the same as those we observed between “attend-out” and “attend-in” in V4 (Fig. 6b), as are the changes in $%sv$ and loading similarity population metrics (Fig. 6c). However, the increase in dimensionality from V1 spontaneous to evoked is in the opposite of what we observed from “attend-out” to “attend-in” in V4 (Fig. 6c, right panel).

Methods. For evoked activity, we computed spike counts for each trial in the time period 160-260 ms after stimulus onset. For spontaneous activity, we computed spike counts during the blank screen in the 100 ms immediately prior to stimulus onset. We chose to use 100 ms bin sizes to match those used in Semedo et al. (2019). We define spike counts during these two time periods during a trial as a “spont-evoked pair”. Each recording session consisted of 400 repeats of a spont-evoked pair for each of the 8 oriented stimuli. For each session, we assessed changes in metrics for each orientation and computed the mean and standard error of the metric across the 8 orientations (transparent colored data points connected by lines). We also plot the average across the 5 sessions (thick black line). To compare metrics for spontaneous and evoked activity, we computed p-values across all 40 datasets (5 sessions, 8 orientations per session) using a paired t-test.
Supplementary Figure 5: Decomposition of the spike count covariance matrix and defining population metrics. Related to Figures 1 and 2, and STAR Methods. a. We use factor analysis to decompose the spike count covariance matrix $\Sigma$ into the sum of a low-rank shared covariance matrix $\Sigma_{\text{shared}}$ and a diagonal independent variance matrix $\Psi$. The $i$th diagonal entry of $\Sigma_{\text{shared}} (s_i)$ corresponds to the spike count variance that neuron $i$ shares with other neurons in the population (i.e., shared variance), while the $i$th diagonal entry of $\Psi_i$ corresponds to spike count variance of neuron $i$ that cannot be explained by the other neurons (i.e., independent to neuron $i$). We can further decompose $\Sigma_{\text{shared}}$ via an eigendecomposition to extract the co-fluctuation patterns (i.e., the eigenvectors) and the strength of each latent co-fluctuation (i.e., the eigenvalues). b. The population metrics used in this study are loading similarity, percent shared variance ($\%sv$), and dimensionality.

<table>
<thead>
<tr>
<th>Population Metric</th>
<th>Mathematical Definition</th>
<th>Intuition</th>
</tr>
</thead>
<tbody>
<tr>
<td>loading similarity</td>
<td>$\text{loading sim.} = 1 - \frac{\text{var}(u_k)}{1/n}$</td>
<td>similarity in weights in co-fluctuation pattern $u_k$</td>
</tr>
<tr>
<td>percent shared variance</td>
<td>$%sv = \frac{1}{n} \sum_{i=1}^{n} \frac{s_i}{s_i + \psi_i} \cdot 100%$</td>
<td>% of neuron's variability explained by other neurons in the population</td>
</tr>
<tr>
<td>dimensionality</td>
<td>$\text{dim.} = \text{rank}(U)$</td>
<td>number of co-fluctuation patterns</td>
</tr>
</tbody>
</table>
Supplementary Figure 6: Characterizing how changes in one population metric can impact the estimates of another population metric. Related to Figure 6.

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In the main text, we related population metrics to pairwise metrics by systematically changing a single population metric and measuring the resulting changes in $r_{sc, \text{mean}}$ and $r_{sc, \text{s.d.}}$ (Figs. 3 and 5). However, in real neuronal data, multiple population metrics could change together between experimental conditions (e.g., see Fig. 6 and Supplementary Fig. 4). When we measure that multiple population metrics changed, it could be the case that a change in one population metric impacted the estimates of the other population metrics (e.g., we could have measured a change in multiple population metrics when only one metric truly changed). This can affect the precision by which we can distinguish population metric changes in real neuronal data.

Here, we assessed this by systematically changing one population metric while keeping the other two population metrics constant. We then simulated data and fit factor analysis (FA) to the data to obtain the population metrics. We examined in turn each of the three population metrics under conditions when they did not actually change (but one of the other metrics did). If there were no dependencies between estimates of population metrics, then all the vertical values in panels a-c would be 0. We found that this was the case for estimates of $\%sv$ were under conditions in which the true loading similarity changed (a). However, estimates of loading similarity and dimensionality were affected by true changes in $\%sv$. Increasing the number of simulated trials reduced the estimation error caused by true changes in $\%sv$ (b, c). These findings allow us to better interpret changes in population metrics estimated from neuronal activity.

a. Estimation error in $\%sv$ due to changes in loading similarity. “Model 1” and “model 2” had the same dimensionality (1) and $\%sv$ (20%). The only difference between the two models was their loading similarity. We varied how different the loading similarity was between the two models (horizontal axis), while assessing how different was the estimated $\%sv$ across the two models (vertical axis). We found that $\%sv$ estimates remained unaffected in the presence of true changes in loading similarity (all changes in $\%sv$ are near 0). This was true for both low loading similarities where “model 1” had loading similarity of 0.05 (blue) and high loading similarities where “model 1” had loading similarity of 0.55 (green). As we simulated more trials, estimates of $\%sv$ became more precise (error bars decrease in size going from left to right panels). Error bars show means and standard deviations across simulations.

b. Estimation error in loading similarity due to changes in $\%sv$. “Model 1” and “model 2” had the same dimensionality (1) and loading similarity. The only difference between the two models was their $\%sv$. We varied how different the $\%sv$ was between the two models (horizontal axis), while assessing how different was the estimated loading similarity between the two models (vertical axis). We found little changes in estimates of loading similarity when the true loading similarity was low (blue points). However, we found larger changes in estimates of loading similarity when the true loading similarity was high (green points). The the size of the change increased with larger true changes in $\%sv$. To understand this, recall that there are relatively few ways to have high loading similarity (e.g., all loadings must be the same to have loading similarity of 1), while there are many ways to have low loading similarity (Math Note E). Thus, high loading similarities are more likely to be underestimated than low loading similarities. This underestimate tends to be larger when $\%sv$ is low than when $\%sv$ is high. However, increasing the trial counts reduced the estimation error of loading similarity (vertical values closer to 0 going from left to right panels).

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c. Estimation error in dimensionality due to changes in $\%sv$. “Model 1” and “model 2” had the same dimensionality (5; with eigenspectrum defined as $\lambda_k = e^{-0.75k}$) and loading similarity. The only difference between the two models was their $\%sv$. We varied how different the $\%sv$ was between the two models (horizontal axis), while assessing how different was the estimated dimensionality ($d_{shared}$; see Methods) between the two models (vertical axis). We found changes in the estimates of dimensionality between “model 1” and “model 2”, and the size of the change increased with larger true changes in $\%sv$. This can be understood by the fact that dimensions with small eigenvalues can be difficult to recover when fitting FA to data, particularly when $\%sv$ is low. However, increasing the trial counts reduced the estimation error of dimensionality (vertical values closer to 0 going from left to right panels).

These results have important implications for interpreting estimated changes in population metrics in real neuronal data. First, because estimation error depends on trial count, one should equalize the number of trials across conditions in order to make fair comparisons across conditions using population metrics. Second, when changes in $\%sv$ are large and trial counts are small, one may need to be careful in interpreting estimated changes in loading similarity and dimensionality. For trial count, the key quantity to consider is the ratio of observed trials to the number of recorded neurons (Wainwright, 2019). In the simulations above, we used 30 neurons—the left column represented a ratio of 5x trials to neurons, the middle column represented 10x, and the right column represented 20x.

In our V4 data (Fig. 6), most sessions had 10 times (or more) the number of trials as the number of neurons (ratio of trials to neurons: 9.90 ± 0.66 for monkey 1, 27.60 ± 2.68 for monkey 2). We observed a difference in $\%sv$ of ≈3% between “attend-out” and “attend-in”. Based on the results in panels a and b, the differences we measured in $\%sv$ and loading similarity in our V4 data are unlikely to be due to estimation error. Based on panel c, the small difference we measured in dimensionality in our V4 data could potentially be explained by a change in $\%sv$, if the only true change between conditions was in $\%sv$ (and not loading similarity or any other aspect of the population activity).
Supplementary Figure 7: Relationships between pairwise and population metrics hold for metrics estimated from Poisson simulated data. Related to Figure 3.

In our simulations and analytical derivations, we created covariance matrices with specified population metrics from which we computed pairwise metrics (Fig. 3). However, when assessing population metrics in neuronal recordings, one needs to fit a factor analysis (FA) model to data. Here, we simulated Poisson data to assess whether the relationships between pairwise and population metrics were impacted by: 1) needing to estimate metrics from data, and 2) the mismatch between the linear-Gaussian assumption of FA and the Poisson-like statistics of neuronal activity. We found that the relationships between pairwise and population metrics were very similar to those shown in Figs. 3 and 5.

a. Estimating loading similarity and %sv. We simulated data from a model with a single co-fluctuation pattern and Poisson observations (see details at the end of the caption). In the ground truth models, we varied loading similarity smoothly between 0 and 1 and chose %sv equal to 30% or 50%. We estimated $r_{sc}$ mean and s.d. from the simulated data. To estimate population metrics, we fit the FA parameters to the same simulated data. We then plotted estimates of pairwise metrics and colored or labeled points according to the estimated population metrics (as opposed to the ground truth population metrics used to generate the data). We found that as estimated loading similarity increased, $r_{sc}$ mean increased and $r_{sc}$ s.d. decreased (blue to green). We also found that as estimated %sv increased, $r_{sc}$ mean and s.d. both increased (inner arc with %sv=32.6±1.0% to outer arc with %sv=55.1±3.2%). These results are consistent with Fig. 3e-f.

b. Same as a, but fixing %sv=50% and varying the dimensionality of the ground truth model, with a flat eigenspectrum (corresponding to Fig. 3g). We colored points according to estimated dimensionality as opposed to the ground truth dimensionality. We found that as the estimated dimensionality increased, $r_{sc}$ mean and s.d. both tended to decrease (purple outer arc with dim=1 to salmon inner arc with dim=5), consistent with Fig. 3g.

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Supplementary Figure 7: (...continued from previous page)

The results here, based on estimating factor analysis parameters from Poisson simulated data, are qualitatively the same as those in the main text (Fig. 3e-g) and analytical derivations (Appendices). This indicates that the relationships between pairwise and population metrics are robust to: 1) having to estimate these metrics from data and 2) the Poisson-like variability of neuronal activity.

Simulating from a Poisson observation model. According to FA, the observations $x$ (i.e., spike counts) have a linear-Gaussian relationship with latent variables $z$ (which represent shared activity among neurons; see Fig. 2): $z \sim N(0, I)$ and $x|z \sim N(Lz + \mu, \Psi)$. We fit the FA parameters to data simulated from a Poisson observation model. We generated Poisson spike counts for 30 neurons as follows. For neuron $i$, we sample from $x_i|z \sim \text{Poisson}(\text{ReLU}(L_i::z + \mu_i))$, where $\text{ReLU}$ indicates a rectified linear unit, and $L \in R^{30 \times d}$ is the loading matrix with $L_{i,:}$ as the $i^{th}$ row. We set $\mu_i = 10$ for each neuron, a typical average firing rate (10 Hz) for neurons across many areas of macaque cortex (assuming a 1 second time bin). We consider the asymptotic case by simulating many trials for each model (corresponding to a single dot in panels $a$ and $b$; see Methods for how model parameters are randomly chosen). We consider estimation from limited data in Supplementary Fig. 6. We drew 6000 samples (i.e., 6000 trials) of $x$ from the Poisson observation model. Thus, this procedure generated a data matrix $X \in R^{30 \times 6000}$ of simulated spike counts, which we then used to estimate pairwise and population metrics.