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Isoflurane Impairs Low-Frequency Feedback but Leaves High-Frequency Feedforward Connectivity Intact in the Fly Brain

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Abstract

Hierarchically organized brains communicate through feedforward (FF) and feedback (FB) pathways. In mammals, FF and FB are mediated by higher and lower frequencies during wakefulness. FB is preferentially impaired by general anesthetics in multiple mammalian species. This suggests FB serves critical functions in waking brains. The brain of Drosophila melanogaster (fruit fly) is also hierarchically organized, but the presence of FB in these brains is not established. Here, we studied FB in the fly brain, by simultaneously recording local field potentials (LFPs) from low-order peripheral structures and higher-order central structures. We analyzed the data using Granger causality (GC), the first application of this analysis technique to recordings from the insect brain. Our analysis revealed that low frequencies (0.1–5 Hz) mediated FB from the center to the periphery, while higher frequencies (10–45 Hz) mediated FF in the opposite direction. Further, isoflurane anesthesia preferentially reduced FB. Our results imply that the spectral characteristics of FF and FB may be a signature of hierarchically organized brains that is conserved from insects to mammals. We speculate that general anesthetics may induce unresponsiveness across species by targeting the mechanisms that support FB.

Key words: directed connectivity; Drosophila; feedback; general anesthesia; Granger causality; local field potentials

Significance Statement

In mammalian brains, neurons communicate through fast (high-frequency) feedforward (FF) and slow (low-frequency) feedback (FB) pathways. Various theories of consciousness propose that FB is crucial for the maintenance of consciousness. Here we used Granger causality (GC) analysis and multi-electrode recordings to investigate FB in the fly brain. Surprisingly, we found that the temporal characteristics of FF and FB are conserved in flies. Further, we found that isoflurane anesthesia preferentially reduced FB, consistent with results from mammalian studies. Our results imply that the mediation of FF and FB by higher and lower frequencies is a signature of hierarchically organized brains and that FB is a conserved, system-level target for general anesthetics.

Introduction

Many complex networks including brains are hierarchically organized. In hierarchical systems information travels both from the bottom to the top of the hierarchy, known as feedforward (FF), and from the top to the bottom of the hierarchy, known as feedback (FB). It is only

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recently that the dynamic characteristics of neural FB and FF have been reported. For example, studies that investigated directed connectivity using Granger causality (GC) across the awake monkey visual hierarchy have shown that the alpha/beta and gamma bands preferentially mediate FB and FF, respectively (Bastos et al., 2014; van Kerkoerle et al., 2014). This pattern of low-frequency FB/high-frequency FF has also been reported across the visual (Michalareas et al., 2016) and auditory system (Fontolan et al., 2014) in awake humans. These studies imply that FB is mediated by lower frequencies than those that mediate FF.

The difference in frequency bands that mediate FB and FF is consistent with the functional explanation of these under the hierarchical predictive coding framework (Hohwy, 2013). According to predictive coding, compared to FF, FB necessarily operates at slower timescales because its main function is to provide prediction of incoming signals through the incorporation of memory and expectation. On the other hand, FF has to react to quickly changing sensory inputs, necessarily operating at faster timescales than FB. Note that these hypotheses may be true for all hierarchically organized biological brains, from insects to mammals, to the extent that these functional requirements are shared across species.

Anatomically, fly brains are hierarchically organized from peripheral sensory layers to central nuclei, which are responsible for more abstract computation (Paulk et al., 2013a; Shih et al., 2015; Haberkern and Jayaraman, 2016). Like mammals, wakeful insect brains also face functional challenges in adjusting fast sensory inputs based on prior experience on a slower time scale (Card and Dickinson, 2008; Combes, 2015; Kim et al., 2015; Mischiati et al., 2015). In addition, recent work suggests that the topological organization of the mammalian brain, which shares a number of similarities with the topological organization of the fly brain (Shih et al., 2015), influences the direction of information flow (Moon et al., 2015; Mejias et al., 2016; Moon et al., 2017). This evidence suggests that similar dynamic characteristics of FF and FB processing will be present in the insect brain.

A separate line of inquiry suggests that FB in particular is crucial for waking brain functions. A number of studies have demonstrated that general anesthetics preferentially reduce FB from frontal to posterior areas as measured using human scalp level electroencephalography (EEG; Lee et al., 2009; Ku et al., 2011; Boly et al., 2012; Jordan et al., 2013; Lee et al., 2013; Ranft et al., 2016). Although a subset of studies show differing results (Barrett et al., 2012; Nicolaou et al., 2012; Maksimow et al., 2014), the disrupted frontal-parietal connectivity has also been demonstrated using functional magnetic resonance imaging (fMRI; Hudetz and Mashour, 2016). In addition, general anesthetic suppression of FB has also been reported in rodents (Imas et al., 2006; Pal et al., 2016), ferrets (Wollstadt et al., 2017), and monkeys (Lamme et al., 1998; Papadopoulou et al., 2015). Note, however, that Wollstadt et al. (2017) show that FB (prefrontal cortex to V1) is reduced, but that this reduction may be a result of reduced “source information” within the prefrontal cortex. Also, Lamme et al. (1998) have shown that general anesthesia reduces contextual processing, which has been taken as an indirect support for the impairment of FB influences to V1.

Flies are also rendered unresponsive by general anesthetics and in similar concentrations to humans (van Swinderen and Kottler, 2014; Zulucki and van Swinderen, 2016). Indeed, the cellular and molecular machinery that makes up the nervous system is largely conserved across species (Littleton and Ganetzky, 2000), such that the cellular and molecular targets of general anesthetics are also likely to be conserved (Sonner, 2008; van Swinderen and Kottler, 2014; Zulucki and van Swinderen, 2016). Whether the conserved cellular and molecular targets of general anesthetics also translate to conserved system-level general anesthetic effects, such as reduced FB, is currently unknown.

Here, we investigated the dynamic characteristics of FB and FF in the small brains of fruit flies, *Drosophila melanogaster*. We analyzed directional connectivity in LFPs recorded from low-order peripheral structures and higher-order central structures. This showed that low frequencies (0.1–5 Hz) mediated FB from the center to the periphery, while higher frequencies (10–45 Hz) mediated FF in the opposite direction. Further, the general anesthetic isoflurane reduced FB while leaving FF intact. Our results imply that the preferential mediation of FB and FF by low and high frequencies, respectively, may be a signature of hierarchically organized brains and that general anesthetics induce unresponsiveness by targeting FB mechanisms.

**Materials and Methods**

**Overview**

We analyzed local field potentials (LFPs) data previously recorded from the brains of awake and anesthetized flies (Cohen et al., 2016). Whereas Cohen et al. (2016) focused on visually evoked responses, here we focus on the analysis of power, coherence, and GC of spontaneous LFPs, and how these are affected by isoflurane. In this section, we briefly recap the experimental setup. For the full experimental details, see Cohen et al. (2016).

Thirteen female laboratory-reared *D. melanogaster* (Can- ton S wild type) flies (3–7 d after eclosion) were collected...
under cold anesthesia, tethered and positioned on an air-supported Styrofoam ball. Linear silicon probes with 16 electrodes separated by 25 μm (Neuronexus 3mm-25-177) were inserted laterally to the eye of the fly until the most peripheral electrode site was just outside the eye. This probe covers approximately half of the fly brain and records neural activity from both peripheral and central brain structures (Fig. 1C). A fine tungsten wire was inserted in the thorax and used as a reference electrode.

Isoflurane was delivered through a rubber hose connected to an evaporator (Fig. 1A,B). The effects of isoflurane were tested at concentrations of 0% and 0.6% (for details, see Cohen et al., 2016). An olfactory stimulus controller was used to deliver six air puffs to gauge the behavioral responsiveness of the flies in each concentration of isoflurane. Fly movement activity in response to the air puff was recorded with a camera. We used the video data to confirm that 0.6% isoflurane abolished the behavioral responsiveness and that behavioral responsiveness returned after recovery (Cohen et al., 2016).

An experiment consisted of several blocks, each at a different concentration of isoflurane. Each block started with (1) a series of air puffs (light blue rectangles) followed by 18 s of rest; (2) presentation of visual stimuli (248 s); (3) 30 s of rest followed by a series of air puffs; and (4) isoflurane concentration change followed by 180 s of rest.
with the delivery of a series of air puffs, followed by −18 s of rest, which is the critical period of data for this paper. Visual stimuli were presented, lasting 248 s in total, followed by an additional 30 s of rest. A series of air puffs were delivered again and then the isoflurane concentration was changed. After 180 s of adjustment to the new isoflurane concentration the experimental block was repeated (Fig. 1D).

In Cohen et al. (2016), we focused on the analysis of the visually-evoked data, which corresponds to the 248 s period of visual stimuli presentation. Here we focused on an 18-s period before the presentation of visual stimuli (Fig. 1D). This no-visual stimulation period is better suited for coherence and GC analysis than the visual stimulation period because the stimulus-evoked responses can result in erroneous interpretation of coherence and GC (Truccolo et al., 2002; Wang et al., 2008). We focused on the period before rather than after visual stimuli presentation because we reasoned the latter period may also reflect residual visual activity.

The datasets analyzed in this study are available from the corresponding authors on request.

**LFP pre-processing**

LFPs were recorded at 25 kHz, downsampled to 1000 Hz, and the most peripheral electrode site was removed from the analysis. We rerereferenced the remaining 15 unipolar channels by subtracting adjacent channels (bipolar rerereferencing) to obtain a set of 14 differential signals which we refer to as “channels” throughout this paper (Bastos et al., 2014; Trongnetrpunya et al., 2016; Cohen and Tsuchiya, 2017).

We divided the 18-s period into eight consecutive epochs of 2.25 s. We removed line noise from each epoch and bipolar rerereferenced channel using three tapers, a window of 0.75 s and a step size of 0.375 s using the rmlinesmoving.m function from the Chronux toolbox (http://chronux.org/; Mitra and Bokil, 2007). We then linearly detrended and z-scored each channel and epoch by removing the mean and dividing by the standard deviation across temporal points within each epoch (Seth, 2010; Bastos et al., 2014). Figure 1C shows example of the resulting LFPs from one fly before (0% isoflurane) and during anesthesia (0.6% isoflurane).

**Analyzing power**

For power analysis we linearly detrended the preprocessed LFPs without the z-scoring operation, because z-scoring obscures overall shifts in power when comparing pre- and postanesthesia. We calculated power for each pre-processed channel (i = [1–14]), epoch e (1-8), k% isoflurane concentration (k = [0 (air), 0.6]) and frequency ω, $S_{kei}(ω)$ over the 2.25-s epochs. Using three tapers, the half bandwidth was 0.89 Hz (Mitra and Pesaran, 1999). We further averaged power across eight epochs to obtain one estimate of power for each channel and isoflurane concentration $S_{ke}(ω)$.

We analyzed the average power across channels within the center $S_{c}(ω)$ and periphery $S_{p}(ω)$ in units of 10log10μV², following the division of channels into peripheral channels=[1–6] and central channels=[9–14] as in Cohen et al. (2016)

$$S_{ci}^{c}(ω) = \frac{1}{6} \sum_{i=1}^{6} S_{kei}(ω)$$

$$S_{pi}^{c}(ω) = \frac{1}{14} \sum_{j=9}^{14} S_{kej}(ω)$$

In the fly brain, the locations of the peripheral channels correspond to the primary visual systems while those of the central channels to higher-order structures (Paulk et al., 2013b; Yap et al., 2017). Heartbeats are unlikely to affect the results presented here because we use bipolar rerereferencing which eliminates common input to neighboring channels, as may be expected from a muscle artifact (Cohen and Tsuchiya, 2017). To further rule out heartbeat as a potential confound we also analyzed GC (see Non-parametric GC analysis) in our data after excluding three flies in which we detected potential heartbeat in central channels by visual inspection of the power spectra. The results of this analysis closely matched those obtained with all thirteen flies, indicating the any heartbeat contamination is unlikely to affect our results.

**Analyzing coherence**

Coherence measures the strength of linear dependency between two signals in the frequency domain (Bendat and Piersol, 2000). It is defined as

$$C^c(ω) = \frac{\text{abs}(CS^{c}(ω))^2}{S(ω)S^c(ω)}$$

$CS^{c}(ω)$ is the cross-spectrum between channels i and j, and $S(ω)$ and $S^c(ω)$ are the power spectra of channels i and j, respectively. To estimate coherence for each bipolar rerereferenced channel pair i and j (i,j = [1–14]), epoch e (1-8) and k% isoflurane concentration (k = [0 (air), 0.6]) we used the multitaper method with nine tapers for each 2.25-s epoch, giving a half bandwidth of 2.22 Hz (Mitra and Pesaran, 1999). The auto and cross spectra for channels i and j at isoflurane concentration k% were averaged across the eight epochs to give one estimate of coherence at each isoflurane concentration.

We report center ($C^c$) and periphery ($C^p$) coherence as averaged across all non-neighboring center and periphery channel pairs.
We exclude neighboring channels pairs because these show artefactual high coherence due to the bipolar referencing (for details, see Shirhatti et al., 2016; Cohen and Tsuchiya, 2017).

We report center-periphery coherence (C_\text{CP}) as averaged across all center-periphery pairs

\[
C_{\text{CP}}(\omega) = \frac{1}{36} \sum_{i=1}^{9} \sum_{j=9}^{14} C_{ij}(\omega)
\]

**Estimating the coherence bias**

Because coherence is limited to the 0–1 range it has an inherent positive bias (Jarvis and Mitra, 2001). We used surrogate data statistics to estimate the empirical coherence bias in our data. The advantages of this method are (1) that it provides an empirical lower bound above which coherence may be considered significant and (2) that it accounts for potentially frequency-specific coherence biases sometimes observed for two time-series with similar power spectra (Faes et al., 2004).

For each of 13 flies we created surrogate data in which we randomized the phase of the signal while keeping the amplitude the same (Faes et al., 2004; Fasoula et al., 2013). We performed this 200 times for each channel (1–14), epoch (1–8) and isoflurane concentration ([0%, 0.6%]) resulting in surrogate data with the same power spectra as the original data but with near-zero cross spectra between channels. The surrogate data describes the null hypothesis that there is no coherence in the data.

To estimate the center, periphery and center-periphery coherence bias at k% isoflurane we calculated coherence as described above for each of the 200 surrogate data sets, and averaged the estimates across all 200 surrogates. We obtained the 95% confidence intervals by determining the 2.5 and 97.5 percentiles across the 200 surrogates for each fly.

When we compared the coherence bias in 0.6% and 0% isoflurane, we observed differences in some frequencies (0.1–5 Hz), but the magnitudes of the differences were very small (largest difference was 0.0038 at around 11 Hz for center coherence). Accordingly, our coherence results in 0.6% and 0% isoflurane with or without bias correction were nearly identical.

**Non-parametric GC analysis**

In simple terms, a signal X is said to Granger-cause a signal Y if past values of X improve predictions of future values of Y. This notion of causality is based on modeling X and Y as autoregressive processes. GC can be estimated in either the time or frequency domain. The latter provides spectral decomposition of GC influences. We hypothesized that examining GC influences at different frequencies is critical because recent work that studied mammalian brains showed that different directional interactions were mediated by different frequencies (Bastos et al., 2014; van Kerkoerle et al., 2014; Michalareas et al., 2016).

GC can be estimated parametrically by fitting an autoregressive model to the data (Geweke, 1982, 1984; Chen et al., 2006). The drawback of this approach is that it requires selection of the model order. Alternatively, GC can be non-parametrically estimated directly from the spectral density matrix (Dhamala et al., 2008; Wen et al., 2013). The spectral density matrix for channels i and j at isoflurane concentration k% Q_i(\omega) is obtained by setting the diagonal elements to the auto-spectra and the cross-diagonal elements to the cross-spectra (Wen et al., 2013)

\[
Q_{ij}(\omega) = \begin{pmatrix}
S_i(\omega) & C_{ij}(\omega) \\
C_{ji}(\omega) & S_j(\omega)
\end{pmatrix}
\]

We estimated the spectral density matrix using the same estimates of auto- and cross-spectra as those used for coherence analysis. To obtain Granger-causal influences from channel i to channel j at isoflurane concentration k% f_k^i(\omega) we first factorized the spectral density matrix using the `sfactorization_wilson.m` function and then estimated the Granger-causal influences using the `ft_connectivity_Granger.m` function. These functions are available from the FieldTrip toolbox (Oostenveld et al., 2011).

**Grouping FF and FB influences**

In the fly brain, visual information enters through the retina and is sequentially processed by the lamina, medulla and lobula, collectively referred to as the optic lobe (Sanes and Zipursky, 2010; Palk et al., 2013a). Visual projection neurons (VPNs) convey information from lower to higher-order visual processing centers, called optic glomeruli, which are located more centrally (Otsuna and Ito, 2006). Relatively little is known about computations in the optic glomeruli, but recent studies suggest a role in second-order motion processing (Zhang et al., 2013) and processing that mediates natural avoidance behaviors (Wu et al., 2016). Specific optic glomeruli, such as the ventrolateral protocereberum (Ito et al., 2014), send projections back to the lobula and medulla, as well as reciprocated projections to other central structures (Otsuna and Ito, 2006; Shih et al., 2015). Our simple summary of this connectome is that visual information enters in the periphery and sequentially conveyed through FF connections toward more central optic glomeruli, which convey FB back to the optic lobe.

We emphasize that our definition of FF and FB is based only on this anatomic organization. Because we did not manipulate bottom-up or top-down processing through stimulus characteristics or task demands, we do not claim that our FF and FB estimates directly reflect bottom-up sensory processing or top-down cognitive functions.

To estimate overall FF GC influences at k% isoflurane f_k^\text{FF}(\omega) we averaged across all GC influences f_k^i(\omega) from the periphery to the center.
\[ f_{ji}^k(\omega) = \frac{1}{36} \sum_{i=1}^{6} \sum_{j=9}^{14} f_{ki}^{-}(\omega) \]

where channel 1 is the most peripheral and channel 14 is the most central. Analogous derivations give FB GC influences

\[ f_{ij}^k(\omega) = \frac{1}{36} \sum_{i=1}^{6} \sum_{j=9}^{14} f_{kj}^{-}(\omega) \]

Estimating the Granger causal influences bias

GC influences are always positive and therefore have a positive bias. We checked whether the GC bias differed between FF and FB or across isoflurane concentrations. To estimate the GC bias, we used the same procedure for the estimation of the coherence bias through phase randomization. This procedure results in 200 estimates of FF and FB GC influences for each fly, at each isoflurane concentration \( k \). We obtained the final estimate of the FF and FB GC bias by averaging across all 200 surrogates.

The differences between FF and FB GC biases at each isoflurane concentration, as well as the differences in FF and FB GC biases between isoflurane concentrations were negligible. The maximum difference in group average bias was 0.0008 (difference between FF and FB at 0.6% isoflurane at around 5 Hz). The difference in GC bias averaged across 0.1–45 Hz were all negligible: 0.00006 (between FF and FB in 0% isoflurane), 0.0002 (between FF and FB in 0.6% isoflurane), 0.0002 (between 0% and 0.6% isoflurane for FF), and 0.0001 (between 0% and 0.6% isoflurane for FB).

Directed asymmetry index (DAI) analysis

Because the magnitude and variance of GC influences are dependent on the frequencies, some form of normalization is necessary for statistically comparing them. For this purpose, we adopt the DAI (Lee et al., 2013; Bastos et al., 2015; Michalareas et al., 2016), which is defined as

\[ DAI = \frac{FF - FB}{FF + FB} \]

The numerator of the DAI captures the predominant direction of the GC influence. The denominator provides normalization for the total GC influences at a given frequency. DAI allows comparisons across conditions and frequencies where there may be an overall net increase or decrease in GC influences.

Given our enumeration of channels (i.e., \( i > j \) indicates FF influences if \( i < j \), the DAI between channel \( i \) and \( j \) at isoflurane concentration \( k \% \) is given by

\[ DAI_{ij}^k(\omega) = \frac{f_{ki}^{-}(\omega) - f_{kj}^{-}(\omega)}{f_{ik}^{-}(\omega) + f_{ki}^{-}(\omega)}, i \in [1 - 6], j \in [9 - 14], \]

We report the DAI averaged over all center-periphery channel pairs

\[ DAI_k(\omega) = \frac{1}{36} \sum_{i=1}^{6} \sum_{j=9}^{14} DAI_{ij}^k(\omega) \]

Statistical analysis

To test for statistical differences between 0% and 0.6% isoflurane across frequencies (power, coherence, GC, and DAI spectrums) we used randomization-based, non-parametric statistical tests with cluster-based multiple comparison correction (Maris and Oostenveld, 2007). Each of the 13 flies provided one spectrum for each condition (0% and 0.6% isoflurane). We then performed paired t tests across 13 flies at each frequency to obtain t scores. If we found a cluster of continuous frequencies at \( p < 0.05 \) level, then we treated them as a first-level significant cluster. As a second-level statistic, we used the sum of the t scores across frequencies within each cluster. We then preformed 8191 randomizations corresponding to all possible randomizations of thirteen flies across the two conditions (\( 2^{13} - 1 = 8191 \)) and calculated the second-level statistic for each randomization. From each randomization, the largest second-level statistics was chosen and used for constructing a randomized distribution. We compared the observed second-level statistics against the randomized distribution to obtain p values.

Figures 2–4 show the results from the first-level (uncorrected) t tests as well as the second-level cluster-based statistics, which corrects for multiple comparisons across frequencies. When we compared DAI spectrums against zero (Fig. 4B,F,J), we randomized the FF and FB labels for each fly (Michalareas et al., 2016). For testing whether the DAIls averaged across low and high frequencies were different from zero (Fig. 4C,D,G,H) we randomized the FF and FB labels and used the group mean as the test metric. For testing the effect of isoflurane on the DAIls averaged across low and high frequencies (Fig. 4K,L), we randomized the isoflurane concentrations labels and used the group mean difference between the conditions as the test metric.

Results

Isoflurane reduces spontaneous power in the center and periphery

We investigated the dynamic characteristics of FB and FF and how these are affected by anesthesia by analyzing LFPs recorded from the center and periphery of awake and anesthetized fly brains (Fig. 1). Before analyzing FB and FF we first assessed whether neural activity is distinguishable across the center and periphery. To do this we analyzed the power of central and peripheral channels separately (Fig. 2A). We found that power in both the center and periphery was highest at low frequencies and decreased as frequency increased. In low frequencies, power in the center was significantly greater than power in the periphery, demonstrating that neural activity was indeed distinguishable across these two anatomic areas (Fig. 2A). At a concentration of 0.6% isoflurane, the flies were rendered unresponsive, as probed by air puffs (Cohen et al., 2016). We found that during anesthesia, power in both the center and periphery was reduced (0.6%...
isoflurane; Fig. 2B). For the center, power was significantly reduced across almost all frequencies tested. For the periphery the reduction was smaller and only significant in a subset of the frequencies tested (Fig. 2C). The reduction in power in the center was greater than the periphery for low frequencies. These observations are consistent with an overall suppression of neural activity but also suggest that this suppression is more consistent in the higher-order central structures of the fly brain.

Isoflurane reduces center, periphery, and center-periphery coherence

Next, we assessed the effects of isoflurane on local and global processing. To do this, we analyzed coherence, a spectral measure of linear dependency between signals (Bendat and Piersol, 2000). To assess the effects on local processing, we analyzed coherence within the center and periphery. We calculated center and periphery coherence using all non-neighboring center and periphery channel pairs (for details, see Materials and Methods).

Within-center coherence was highest for low frequencies and showed an additional peak around 10–20 Hz (Fig. 3A). Because coherence is confined to 0–1, it is inherently biased. Using surrogate data methods (see Materials and Methods), we found that the coherence was above the bias level for 0.1–45 Hz (Fig. 3A). We repeated the coherence analysis during anesthesia and found that the coherence remained above the bias for all frequencies (Fig. 3B). Isoflurane significantly reduced within-center coherence around 15 Hz (Fig. 3C).

Similarly, within-periphery coherence was highest for low frequencies, decreased as frequency increased and remained above the bias for 0.1–45 Hz (Fig. 3D). Unlike within-center coherence, however, within-periphery coherence had no prominent peak in the 10–20 Hz range (Fig. 3D). During anesthesia all frequencies remained above the bias (Fig. 3E). Isoflurane significantly reduced within-periphery coherence in the 15–25 Hz range (Fig. 3F).

Finally, we assessed the effects of anesthesia on more global processing by calculating center-periphery coherence, which we calculated using all center-periphery channel pairs (for details, see Materials and Methods). We found that similar to center and periphery coherence, center-periphery coherence was highest for low frequencies, decreased with increasing frequency and remained above the bias for 0.1–45 Hz (Fig. 3G). During anesthesia, low-frequency coherence remained above the bias, whereas high-frequency coherence was not distinguishable from the bias (Fig. 3H). Isoflurane significantly reduced center-periphery coherence around the 10–25 Hz range.

In sum, our coherence results indicate that isoflurane impaired localized processing within the center and periphery as well as global processing between the center and periphery.

Low- and high-frequencies preferentially mediate FB and FF, respectively

Finally, we analyzed directional connectivity. To dissect the directionality we used GC analysis (Dhamala et al.,

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**Figure 2.** Isoflurane reduces power in the center and periphery. A, Group average power in the center (orange) and periphery (green) in 0% isoflurane (before any anesthesia). A schematic representation of the fly head and inserted electrode showing the estimated positions of one central (orange) and one peripheral (green) channel is depicted above. Black dots and solid line mark uncorrected and corrected significant differences between center and periphery power at the $p < 0.05$ level, respectively (see Materials and Methods). B, Same as A in 0.6% isoflurane. C, The effect of isoflurane on power in the center and periphery, obtained by subtracting values in 0% isoflurane from values in 0.6% isoflurane. Negative values indicate that isoflurane reduced power for both the center and periphery. For central channels, orange thick lines above indicate significant reductions due to isoflurane (corrected, $p < 0.05$). For peripheral channels, green thick lines above indicate corresponding significant reductions (corrected, $p < 0.05$). Greater reductions in power in the center compared to the periphery are indicated by the black line (corrected, $p < 0.05$). Shaded area represents SEM across flies ($N = 13$) for all panels.
We analyzed Granger causal influences from the periphery to the center as FF GC influences, and influences from the center to the periphery as FB GC influences (Fig. 4). We found that both FF and FB GC were well above any bias level in the GC measure. FB GC appeared greater than FF in lower frequencies (Fig. 4A). However, as is apparent in Figure 4A, the variance of spectral GC influences also appears larger in lower frequencies. Thus, before comparing FF and FB GC we corrected for these statistical characteristics. To

Figure 3. The effect of isoflurane on center, periphery, and center-periphery coherence. A, Coherence between electrodes in the central brain in 0% isoflurane (black line). Gray shades represent SEM across flies (N = 13). Purple line and shades represents the bias level and its 95% confidence intervals. A schematic representation of the fly head and inserted electrode showing the estimated positions of two central channels is depicted above. B, Same as a in 0.6% isoflurane. C, The effect of isoflurane on center coherence, obtained by subtracting values in 0% isoflurane from values in 0.6% isoflurane. Dots and solid lines below the effect of isoflurane (C, F, I) indicate significant reductions (p < 0.05) uncorrected and corrected, respectively, for multiple comparisons. The same format is repeated in D–F for within-periphery coherence and G–I for center-periphery coherence.
Figure 4. Isoflurane reduces low-frequency (0.1–5 Hz) FB in the fly brain. A, Group level FF (blue) and FB (red) GC influences before any anesthesia (0% isoflurane). Black dots mark significant differences between FF and FB at the \( p < 0.05 \) level (uncorrected). The FF and FB GC bias levels (see Materials and Methods) are shown as dashed red and blue lines, which almost completely overlap near zero. B, Group level DAI before any anesthesia (0% isoflurane). Positive values indicate FF > FB and negative values indicate FF < FB. Black dots and solid line mark significant uncorrected and corrected differences from zero at the \( p < 0.05 \) level. Low (0.1–5 Hz) and high (10–45 Hz) frequencies are indicated by the pale pink and purple shadings. C, D, Group level DAI averaged for low (0.1–5 Hz; C) and high (10–45 Hz; D) frequencies before any anesthesia (0% isoflurane). Negative and positive DAI values indicate that FB > FF for low frequencies and that FB < FF for high frequencies. The fly head schemas above the graphs depict this as thicker FF (red, C) and FB (blue, D) arrows. E–H, Same as A–D for 0.6% isoflurane. I, The effect of isoflurane on FF (blue) and FB (red) GC, obtained by subtracting values in 0% isoflurane from 0.6% isoflurane. Blue (and red) dots and solid line mark significant uncorrected and
do this, we employed a normalized measure known as the DAI (Lee et al., 2013; Bastos et al., 2014; Michalareas et al., 2016). The DAI is defined as

$$DAI = \frac{FF - FB}{FF + FB},$$

which is positive when FF is stronger than FB and vice versa. For the DAI the variance was uniform across all frequencies (Fig. 4B).

During wakefulness (0% isoflurane), the DAI was negative for lower frequencies (~0.1–5 Hz), indicating that FB is stronger than FF. For higher frequencies (10–45 Hz) the DAI is predominantly positive, indicating that FF is stronger than FB. To summarize these results, we averaged the DAI for low (0.1–5 Hz; Fig. 4C) and high (10–45 Hz; Fig. 4D) frequencies separately. We found that in awake flies FF is stronger than FB for low frequencies (0.1–5 Hz; Fig. 4C, $p < 0.05$, group-level permutation tests, see Materials and Methods) and that FB is weaker than FF for high frequencies (10–45 Hz; Fig. 4D, $p < 0.01$). This global pattern of results, that is, the dominance of FB in lower frequencies and the dominance of FF in higher frequencies, is generally consistent with what has been reported in the awake mammalian brain (Bastos et al., 2014; Fontolan et al., 2014; van Kerkoerle et al., 2014; Michalareas et al., 2016).

### Isoflurane reduces FB

Next, we examined the effect of isoflurane on FF and FB GC. In humans, general anesthetics have been shown to preferentially reduce FB (Lee et al., 2009; Boly et al., 2012; Jordan et al., 2013; Lee et al., 2013; Ranft et al., 2016). In our data, we found that during anesthesia both FF and FB influences were reduced in magnitude (Fig. 4E). The DAI, which normalizes the magnitude and variance across frequencies, was no longer negative for low frequencies but remained positive for high frequencies (Fig. 4F). This meant that during anesthesia the dominance of FB over FF was lost in low frequencies (Fig. 4G), and that the dominance of FF over FB in high frequencies remained intact (Fig. 4H). Statistical testing confirmed a frequency- and direction-specific effects on GC (Fig. 4I), resulting in an increase in the DAI in low frequencies (Fig. 4JK; 0.1–5 Hz, $p < 0.05$). There was no change in the DAI in high frequencies (Fig. 4L; 10–45 Hz, $p = 0.8$).

### Discussion

In hierarchically organized brains, dedicated mechanisms may be necessary for routing information in an effective manner. In particular, the hallmark dynamics of hierarchical brains may be the modulation of lower-level, fast responses to the changing environment by higher-level, slower processing that incorporate prior information and expectation (Tani and Nolfi, 1999; Friston, 2010; Bastos et al., 2012; Hohwy, 2013). We found that, like human and monkey brains, FB and FF influences are mediated by lower and higher frequencies in fly brains. This result is surprising given the obvious differences in the number of neurons ($\sim 10^3$ neurons in flies (Chiang et al., 2011) and $10^{11}$ neurons in humans (Herculano-Houzel, 2009) and gross neuroanatomy between these brains. However, a recent study that analyzed the fly connectome has demonstrated a number of organizational similarities between the fly and mammalian brains, such as hierarchical structure and other graph theoretical characteristics (e.g., small-world and rich-club organization; Shih et al., 2015). These structural similarities may translate to similar dynamic characteristics of FB and FF, as recently demonstrated by analytical and computational studies (Moon et al., 2015, 2017; Mejias et al., 2016).

From a functional viewpoint, the dynamical characteristics of FF and FB may also serve similar roles across species, and possibly even hierarchically organized robots (Yamashita and Tani, 2008). Predictive coding suggests that “predictions” travel in the FB direction while “prediction errors” travel in the FF direction (Tani and Nolfi, 1999; Friston, 2010; Bastos et al., 2012; Hohwy, 2013). Because the updating of predictions depends on gathering prediction errors over time, predictions are inherently mediated through slower time scales (lower frequencies) than prediction errors (Bastos et al., 2012; Bastos et al., 2015; Michalareas et al., 2016). In insects, a number of recent studies have demonstrated predictive behaviors and begun to investigate their neural correlates (Card and Dickinson, 2008; Combes, 2015; Kim et al., 2015; Mischiati et al., 2015). Our finding that FB and FF influences are mediated by low and high frequencies in the fly brain is consistent with the idea that these animals are also engaging in a type of predictive coding. Predictive coding can be formulated under the more fundamental framework of free-energy minimization (Friston, 2010), which may apply to biological systems more broadly (Friston, 2012).

We found that isoflurane reduced both local coherence within the center and periphery as well as global coherence between the center and periphery. For directional connectivity, we found that both FF and FB were reduced, but that for low frequencies isoflurane preferentially reduced FB. A number of human studies have demonstrated that general anesthetics preferentially reduce frontal to posterior FB, as measured using EEG (Lee et al., 2009, 2013; Boly et al., 2012; Jordan et al., 2013; Ranft et al., 2016). Interestingly, a recent study using magneto-
encephalography (MEG) found that such anterior to posterior (FB) influences are mediated by lower frequencies than those that mediated posterior to anterior (FF) influences (Hillebrand et al., 2016). Our findings in flies parallel results previously reported for mammals. However, we acknowledge important limitations in regards to this comparison. In particular, we presented analyses of spontaneous, not sensory-evoked, LFPs. In the absence of a sensory signal to transmit, we do not know if the FF and FB we detected signal bottom-up and top-down processing. In this sense, cognitive processing or information transfer was not directly assessed. In Bastos et al. (2014), the presentation of a stimulus was necessary for clearly recruiting FF influences, which were interpreted as signaling bottom-up processing. While we did record data from periods of visual simulation (for details, see Cohen et al., 2016), these data were dominated by the stimulus frequency, violating the assumption of GC. It remains possible that in an experiment that uses sensory stimulation to engage bottom-up processes (yet left the data suitable for GC analysis), we would have detected more pronounced FF signaling and that this signaling would in fact be reduced by general anesthetics. We also emphasize that not all general anesthetic effects found in mammals translate to flies. Indeed, one of the clearest system-wide effects of general anesthetics (as well as sleep) is an increase in low-frequency (0.1–4 Hz) power (Steriade et al., 1993; Amzica and Steriade, 1998; Chauvette et al., 2011; Murphy et al., 2011; Lewis et al., 2012; Purdon et al., 2013, 2015). Such an increase has not been reported for insects (van Swinderen, 2006; Kirszenblat and van Swinderen, 2015), and we did not observe this here (the results did not change with longer time window (18 s) and/or using a unipolar reference scheme [data not shown]). Interestingly, a recent study reported that sleep induction in flies is accompanied by an increase in 7–10 Hz LFPs power (Yap et al., 2017). However, this was not observed for pharmacologically induced sleep, so any relevance to general anesthesia remains unknown. Further studies of population-level dynamics in flies will be required to fully understand the similarities between the way general anesthetics affect the fly and mammalian brain. Our study is a step in this direction, and suggests that low-frequency FB may be a system-wide target process for general anesthetics that is conserved across species.

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