

Brainstem arousal systems adaptively shape large-scale cortical interactions for flexible decision-making

R.L. van den Brink^{1*}, K. Hagen¹, N. Wilming¹, P.R. Murphy^{1,2}, J. Calder-Travis¹, J. Finsterbusch³, C. Büchel³, & T.H. Donner^{1,4*}

¹Computational Cognitive Neuroscience Section, Department of Neurophysiology and Pathophysiology, University Medical Center Hamburg-Eppendorf, Hamburg, Germany.

²Trinity College Institute of Neuroscience and School of Psychology, Trinity College Dublin, Dublin, Ireland.

³Institute for Systems Neuroscience, University Medical Center Hamburg-Eppendorf, Hamburg, Germany

⁴Bernstein Center for Computational Neuroscience, Charité Universitätsmedizin, Berlin, Germany.

* Correspondence: r.van-den-brink@uke.de, t.donner@uke.de.

Abstract

Many cognitive tasks require a flexible mapping from specific features of sensory input to motor output. Such flexible input-output mapping is reflected in intrinsic correlated variability of activity within the cortical network that implements the decision process, and might rely on rapid plasticity mechanisms that are under neuromodulatory control. Here, we test for a role of neuromodulators in flexible decision-making by combining brainstem fMRI and pupillometry with time-resolved tracking of feature-specific intrinsic correlations within the human sensory-motor network. Human participants reported visual orientation judgments where the correct responses were contingent upon an active rule that could switch unpredictably. Rule switches evoked brainstem and pupil responses and changes in latent variables of behavior that were quantified with a computational model. Behavioral variables in turn were encoded in pupil dynamics. Brainstem activity and pupil dilation preceded fluctuations of stimulus-action coupling strength within the cortical network that implemented the decision. Brainstem arousal systems may thus instigate a context-dependent reorganization of selective cortical pathways for flexible decision-making.

Introduction

Decision-making involves the transformation of sensory signals into motor actions (Mante et al., 2013; Shadlen and Kiani, 2013). The transformation from sensation to action has been studied predominantly using tasks with static associations between stimulus and action (Bogacz et al., 2006; Gold and Shadlen, 2007; Donner et al., 2009; Hanks et al., 2015; Wilming et al., 2020; Murphy et al., 2021). However, the mapping between stimulus and action is often not fixed, but must be flexibly altered in accordance with environmental demands (Okazawa and Kiani, 2023). For example, in order to alternate between two stimulus-response (SR) mapping rules (Figure 1a), the brain must flexibly route stimulus information from visual cortex to the population of neurons in motor cortex that encode the correct action for a given stimulus and SR rule (Miller and Cohen, 2001) (Figure 1b). How the brain is able to flexibly remap SR-associations has remained a key question in decision neuroscience (Shadlen and Kiani, 2013).

We recently showed that arbitrary SR rules (Figure 1a) are instantiated in correlated variability of ongoing fluctuations of stimulus and action patterns amongst the regions that implement the decision process (van den Brink et al., 2022). That is, spontaneous fluctuations of orientation-selective fMRI signals in visual cortex were correlated with fluctuations in motor cortex that were selective to the appropriate action (in line with the schematic in Figure 1b). This held both in situations where the SR rule was explicitly instructed and where it switched in a hidden and unpredictable manner and needed to be inferred from noisy cues. In the latter case, the SR coupling patterns reflected the participants' internal belief about the active SR rule (gauged through a computational model). This observation supports the idea that task-specific SR pathways are constantly reconfigured when dictated by the environment, guided by higher-order inference processes. What mechanisms confer this flexible reconfiguration of SR pathways?

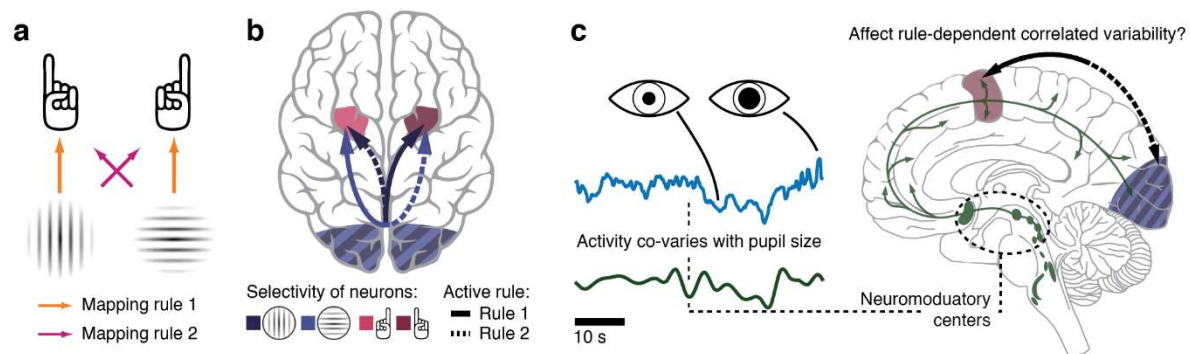


Figure 1. Rationale. **a.** Elementary perceptual choice task. Each visual grating is associated with a unique correct action, which depends on a currently active response rule. **b.** Schematic of rule-dependent correlated variability. Arrows indicate positive correlation. When rule 1 is active, the activity of the population of neurons in visual cortex that encodes a vertical grating correlates positively with those neurons that encode a left hand response. Similarly, horizontal-encoding neurons correlate with the right hand response. When rule 2 becomes active, this pattern flips sign. **c.** Experimental hypothesis. We expect signals from neuromodulatory brainstem nuclei (and their correlate in pupil responses) to (i) track changes in SR rule and (ii) modulate the strength of features-specific correlated variability in the sensory-motor network.

Theoretical work indicates that the brain can flexibly reconfigure SR pathways through short term synaptic plasticity mechanisms that are in turn governed by subcortical neuromodulatory inputs (Fusi et al., 2007). These nuclei project widely to the forebrain and are thus ideally situated to affect large-scale cortical activity dynamics (Berridge and Waterhouse, 2003; Aston-Jones and Cohen, 2005; van den Brink et al., 2019; Pfeffer et al., 2021; Podvalny et al., 2021;

Pfeffer et al., 2022), and control cortical state and arousal (Harris and Thiele, 2011; McGinley et al., 2015b). Moreover, neuromodulators are known to promote synaptic plasticity on multiple temporal scales (Bear and Singer, 1986; Rasmusson, 2000; Reynolds et al., 2001; Reynolds and Wickens, 2002; Berridge and Waterhouse, 2003; Vetencourt et al., 2008; Marzo et al., 2009; Nadim and Bucher, 2014). The activity of neurons that release neuromodulators is also sensitive to stimulus features that can signal the need to adjust behavior (Berridge and Waterhouse, 2003; Aston-Jones and Cohen, 2005; Bouret and Sara, 2005; Sarter et al., 2009), including specific computational variables entailed in complex inference strategies (Dayan and Yu, 2006; Nassar et al., 2012; Muller et al., 2019; Murphy et al., 2021). Combined, these characteristics indicate that neuromodulators may track hidden changes in SR rules and sculpt correlated variability of cortical signals that conditionally link stimulus with action (Figure 3c).

We tested these hypotheses by reanalyzing fMRI data from a previously published study (van den Brink et al., 2022). Here, we probe the relationship between the SR rule inference process, and brainstem fMRI signals, pupil diameter, and intrinsic correlations within the human sensory-motor network. Participants carried out an elementary perceptual choice task (Figure 1a) that was coupled to an inference problem: an active SR rule had to be inferred from ambiguous sensory evidence, and underwent hidden and unpredictable switches (van den Brink et al., 2022). The hidden rule switches evoked robust responses in several neuromodulatory nuclei as well as in the pupil (Figure 1c). These physiological markers of arousal in response to rule switches could be understood in terms of a concomitant increase in computational model variables, which signaled that a change in rule was likely to have occurred. These same markers of arousal also preceded an increase in SR coupling strength within the cortical network that implemented the decision. Our results suggest that brainstem arousal is driven by key computational variables for context inference, and helps shape the continuous and context-dependent reorganization of task-specific pathways within the cortex.

Results

Inferring volatile sensory-motor mapping rules under uncertainty

Participants (N=18) inferred an active sensory-motor mapping rule (Figure 1a) from a stream of ambiguous sensory evidence, which came in the form of rapidly presented dots (cues) along the horizontal meridian (Figure 2a). The rule itself could switch unpredictably with a low probability (1/70), and determined the correct response for elementary perceptual choices: orientation judgements of large and full-contrast visual gratings that were interspersed in the stream of dots at long and variable intervals. Despite ambiguity of the sensory evidence, participants were able to determine and apply the active rule well above chance level, and similarly across the two rules (Figure 2b,c; Table S1). Switches of the active rule resulted in a marked dip in performance, which recovered as sensory evidence in favor of the new rule accumulated (Figure 2d).

The normative (Bayesian) strategy for solving this rule inference task entails an adaptive, non-linear accumulation of sensory evidence in a way that balances build-up of stable belief states with sensitivity to hidden rule switches (Glaze et al., 2015; Piet et al., 2018; Murphy et al., 2021). We could model this behavior with a Bayesian model of the adaptive inference process (Glaze et al., 2015) fit to the participants' choice behavior. Adaptive sensitivity to rule switches was implemented in this model by a non-linear transformation of a prior before integrating it with newly arriving sensory evidence. The shape of this non-linear transformation depended on a

subjective estimate of the probability of rule switches. The fitted model quantified the ‘belief’ (L) of the participant in favor of one rule over the other. This belief has previously been shown to track patterns of correlated variability in the cortex (van den Brink et al., 2022), and here also systematically changed sign after rule switches (Figure 2e).

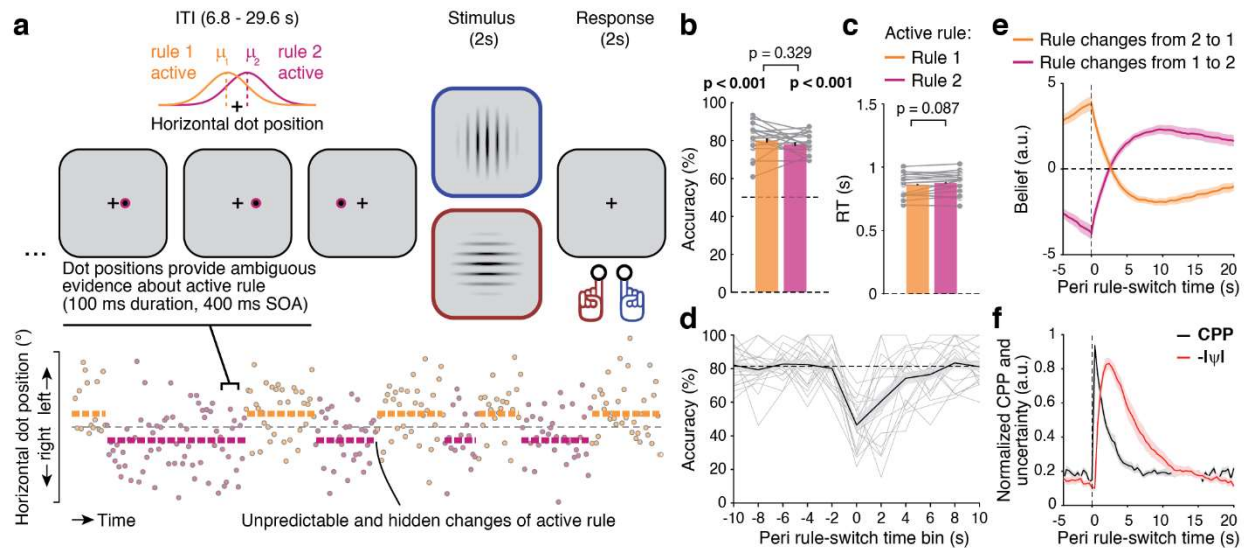


Figure 2. Task and behavior. **a.** Inferred rule task. Top: Example sequence of three evidence samples during the inter-trial-interval (ITI) preceding a task trial. Evidence samples were the horizontal positions of dots presented every 400 ms. Samples were drawn from one of two overlapping distributions, producing a noisy evidence stream. Bottom: example of noisy evidence stream. The generative distribution governed the active rule and could switch unpredictably between any two samples (probability: 1/70). **b.** Accuracy, and **c.** response time (RT) for each rule. The horizontal dotted line in **b.** shows chance level accuracy. Gray dots, individual participants. Bars, group average. Error bars, SEM. **d.** Accuracy for time bins centered on the onset of hidden rule changes. Accuracy drops substantially following a rule change and gradually recovers as participants accumulate evidence in favor of the rule change. Gray lines, individual participants. Black line, group average. Horizontal dotted line, average accuracy before the rule switch. Error bar, SEM. **e.** The belief-parameter from the model locked to rule switches. **f.** Model-derived behavioral variables locked to the onset of rule switches. Following a switch in rule, change point probability (CPP) and uncertainty ($-|L|$) increase. Error bar, SEM.

Here, we focus on two other latent variables of the model that have been shown to modulate behavioral evidence accumulation profiles and pupil-linked arousal responses during active inference (Murphy et al., 2021): change point probability (CPP), and uncertainty ($-|L|$). CPP indicated the probability that a switch of rule has just occurred, given the participants’ subjective estimate of the probability of rule change, the new sensory evidence, and current belief. CPP was high in cases where a new sample of evidence was inconsistent with the participants’ current belief – in particular, when both the previous belief and new (contradictory) evidence were strong. Uncertainty about the environmental state (i.e., active SR rule) before encountering a new sample of sensory evidence was tracked by $-|L|$. Switches of active rule were followed by a peak in both quantities, with a rapid rise in CPP, and a subsequent and more protracted rise in uncertainty (Figure 2f).

In sum, hidden rule switches elicited a sequence of latent processes that included the detection of sensory evidence that violated previously held beliefs (CPP), a rise in uncertainty about the active rule ($-|L|$), and a flip in the sign of the belief state L . Ultimately, these internal computational events led to an adjustment of behavior to the new rule and, consequently, a recovery of task accuracy.

Correlated variability of stimulus and action codes reflects active rule

In our prior work (van den Brink et al., 2022) we showed that correlated variability of population codes for stimulus and for action covaried, with a sign that depended on the model-derived belief of the participant. Here, we aimed to probe the mechanisms that underly the changes in correlated variability, elicited by the rule switch events that prompted changes of belief. We therefore examined if correlated variability aligned with the active rule directly.

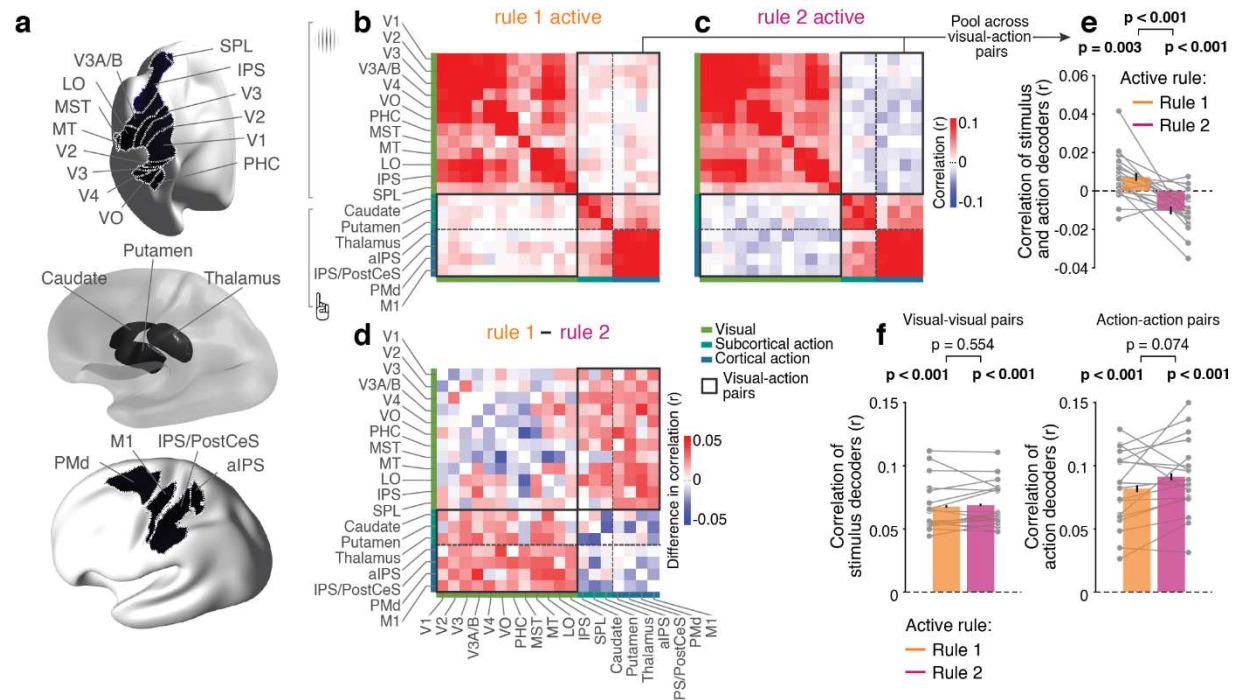


Figure 3. Correlated variability of population codes for stimulus and action. **a.** Regions of interest for the analysis of correlated variability. **b., c., d.** Correlation matrices for all stimulus and action decoders during rule 1 (b.), rule 2 (c.), and the difference between rules (d.). Gray rectangles, stimulus-action pairs. **e.** correlations from (a.) and (b.), collapsed across all pairs in the gray rectangles. **f.** Correlations from (a.) and (b.), collapsed across all visual-visual pairs and action-action pairs (excluding values on diagonals). Gray lines, individual participants; bars, group average; error bars, SEM.

We quantified ongoing fluctuations of population codes as the graded output of decoders that captured stimulus orientation (horizontal vs. vertical) or chosen action (left hand vs. right hand response) discriminant patterns. These decoders were applied to the patterns of spontaneous fMRI signal fluctuations within a set of cortical and sub-cortical regions (Figure 3a; Table 1) from which stimulus and action-evoked activity had been removed. The resulting time series indicated if the activity pattern at any given moment tended toward vertical or horizontal stimulus orientation (for visual cortical regions), or towards left hand or right hand button press (for sub-cortical and cortical motor-related regions). We then correlated the stimulus and action decoder outputs separately for intervals corresponding to the two rules (Figure 3b,c).

The sign of correlation of specifically the stimulus-action pairs was a product of the design of the decoders: positive for coupling consistent with rule 1, and negative for rule 2 (cf. van den Brink et al., 2022). The key prediction for rule-dependent correlated variability was that the sign of correlations of stimulus-action pairs should differ between rules. This is indeed what we observed (Figure 3d,e; Table S1). No such sign flip was observed for the correlations of

stimulus-stimulus or action-action region pairs (Figure 3f; Table S1). In sum, the active rule was instantiated in patterns of spontaneous correlations of stimulus and action codes across a task-relevant network of brain areas.

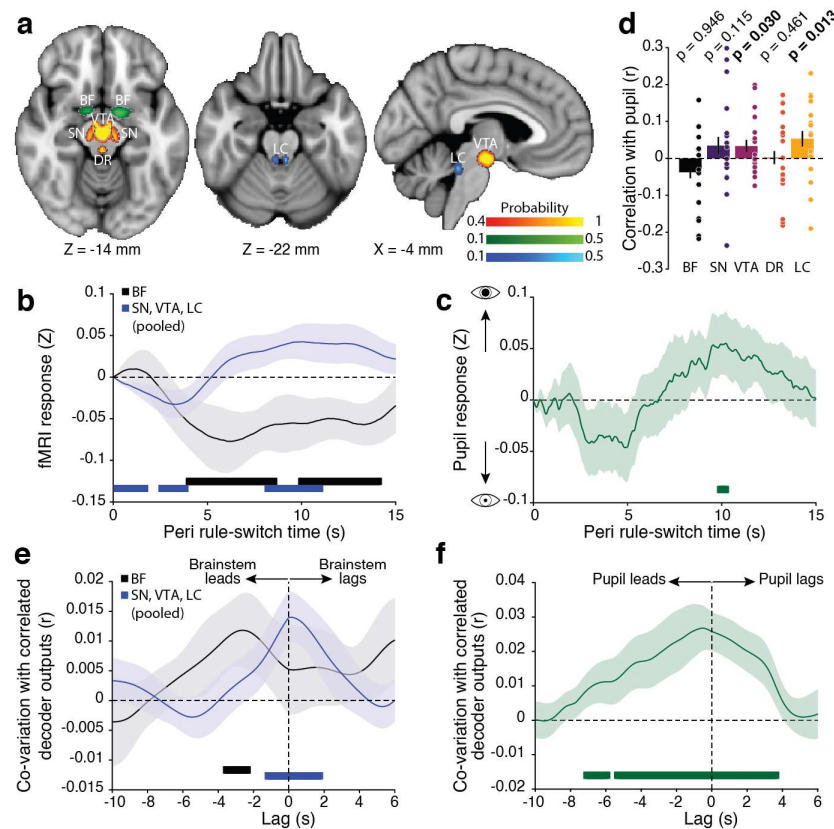


Figure 4. Relationship between brainstem neuromodulation, pupil dynamics, and correlated variability. a. Regions of interest within the brainstem, defined via neuromelanin sensitive scans of individual participants (LC), and anatomical atlases (other nuclei; Table 1). BF, basal forebrain. SN, substantia nigra. VTA, ventral tegmental area. DR, dorsal raphe. LC, locus coeruleus. **b.** Deconvolved response in the brainstem evoked by rule switches. **c.** Deconvolved pupil response evoked by rule switches. **d.** Covariation between the rule switch-evoked response magnitude between the brainstem and pupil. **e.** Cross-correlation between brainstem activity and stimulus-action decoder coupling within the cortex. Negative lags indicate activity in the brainstem preceding increases in decoder coupling. **f.** Cross-correlation between pupil diameter and stimulus-action decoder coupling within the cortex, corrected for hemodynamic delays. Negative lags indicate pupil dilation preceding increases in decoder coupling. In all panels, horizontal bars indicate $p < 0.05$, corrected for multiple comparisons with cluster-based permutation testing. Error bars, SEM.

Brainstem neuromodulatory centers and pupil-linked arousal are recruited by rule switches

Switches of the active rule prompted adjustments of behavior (Figure 2) as well as a sign flip of correlated variability among the regions that implemented the decision (Figure 3). We next examined if rule switches also evoked activity in brainstem centers that release neuromodulators throughout the brain.

We focused on five neuromodulatory centers (Figure 4a; Table 1): the cholinergic basal forebrain (BF), dopaminergic substantia nigra (SN) and ventral tegmental area (VTA), the serotonergic dorsal raphe (DR), and noradrenergic locus coeruleus (LC) (van den Brink et al., 2019). Imaging of the brainstem is difficult due to the size and location of the nuclei involved.

We thus applied physiological noise correction and controlled for additional noise contained in ventricular signals (Brooks et al., 2013; de Gee et al., 2017). For optimal precision in localization, we delineated the smallest nucleus, the LC, within each individual, based on neuromelanin sensitive anatomical scans (Keren et al., 2009; de Gee et al., 2017) that highlighted the LC as hyperintense spots (Figure S1). The location of the resulting masks corresponded well to the known location of the LC in the posterior part of the pons, at the base of the fourth ventricle (Figure S2a). For the other (larger) regions, we relied on publicly available atlases (Table 1).

Switches of the active rule caused prominent evoked activity in the catecholaminergic (i.e., dopaminergic and noradrenergic) nuclei (SN, VTA, and LC; Figure 4b; Table S1). These nuclei are strongly inter-connected (Sara, 2009) and have been implicated in orchestrating resets of cortical activity in response to contextual changes (Dayan & Yu, 2006; Fusi et al, 2007). We thus combined these three catecholaminergic brainstem nuclei in our analyses (Figure S2b shows individual nuclei). The DR did not show significant responses to rule switches (Figure S2b). Unexpectedly, the BF responded to rule switch events with a marked suppression of activity (Figure 4b; Table S1).

Non-luminance mediated fluctuations of pupil diameter are known to covary with activity in neuromodulatory centers of the brainstem, including the catecholaminergic nuclei (Murphy et al., 2014; McGinley et al., 2015a; Joshi et al., 2016; Reimer et al., 2016; de Gee et al., 2017; Larsen and Waters, 2018; Breton-Provencher and Sur, 2019). Indeed, we found pupil diameter to covary with the majority of nuclei as well (Figure S2c; Table S1). Critically, the pupil also dilated in response to the rule switches (Figure 4c; Table S1), and in a manner that was correlated with the rule-switch response of the VTA and LC (Figure 4d; Table S1).

In sum, hidden rule switches recruited catecholaminergic brainstem nuclei and pupil-linked arousal in a coordinated fashion. This suggests that brainstem arousal systems may orchestrate the coupling of population codes for stimulus and action within the cortex, which we examined next.

Brainstem and pupil fluctuations precede fluctuations of correlated variability

If brainstem arousal systems affect SR-coupling within the cortex, we expect to see increased activity in the brainstem prior to increases in cortical signatures of SR coupling strength. We thus cross-correlated the signals from all brainstem nuclei that responded to rule switches and covaried with the pupil (i.e. excluding the DR) and the pupil signal itself with the strength of stimulus-action decoder correlation. To this end, we evaluated the latter in a time-variant fashion. We used strength (i.e., absolute value) of the decoder correlations rather than the correlations per se, because we did not expect the brainstem signals to bias the decoder correlations in any particular direction (i.e., favor a specific SR rule), but rather to boost whichever coupling currently dominated (e.g., through plasticity mechanisms (Fusi et al., 2007)). We evaluated the correlations across a series of lags in order to chart temporal dependencies.

We found a pronounced peak at negative lags in the cross-correlation spectrum between BF and stimulus-action decoder coupling (Figure 4e; Table S1). This indicated that peaks of activity in the BF preceded peaks in stimulus-action decoder coupling. We also found significant correlations between activity of the pooled catecholaminergic nuclei and stimulus-action decoder coupling, including at negative lags, and peaking around zero lag (Figure 4e; Table S1). Finally, the pupil showed dilation before subsequent peaks in stimulus-action decoder

coupling strength (Figure 4f; Table S1). These findings, specifically the lags of the correlations, could not be explained by hemodynamic or pupillary delays because all delays were accounted for in the analysis. Moreover, because the pupil signal was sampled at a higher rate than the brainstem signal, it may be more sensitive to short (~ 1 s) lags between brainstem responses and changes in SR-coupling within the cortex.

Together, our results are consistent with the idea that activity in neuromodulatory nuclei (within the BF, and indexed by pupil diameter) tracked hidden rule switches, and modulated rule-specific coupling of stimulus and action codes amongst the network of brain regions that implemented the primary decision process. The brainstem receives input from brain areas that are involved in active inference, such as anterior cingulate (Aston-Jones and Cohen, 2005). We thus next asked if the arousal system encoded latent variables of behavior that are informative of rule switches.

Pupil fluctuations encode computational variables of behavior

Sensitivity of the arousal system to rule switches may be conferred by latent variables that track changes in environmental state: CPP and uncertainty ($-|\psi|$) (Nassar et al., 2012; Murphy et al., 2021). In our task, samples of evidence were presented without temporal jitter and at a rapid pace (Figure 2a), likely too quickly to relate brainstem fMRI responses to sample-wise estimates of CPP and $-|\psi|$. Nevertheless, the pupil was sampled at a faster rate than the fMRI signal. Given its close correspondence to brainstem activity, both expected from prior work (Murphy et al., 2014; McGinley et al., 2015a; Joshi et al., 2016; Reimer et al., 2016; de Gee et al., 2017; Larsen and Waters, 2018; Breton-Provencher and Sur, 2019), and observed here (Figure 4, Figure S2c), we used the pupil as a summary signal for probing relationships with computational variables of behavior.

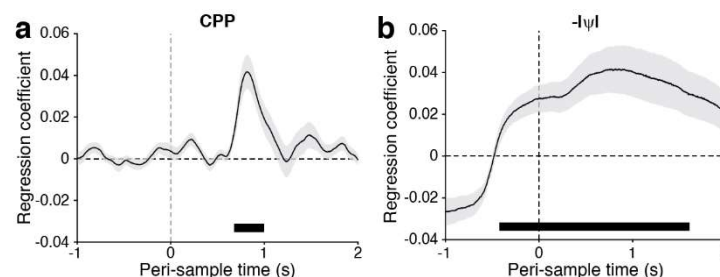


Figure 5. Relationship between pupil dynamics and behavior. **a.** Relationship between the derivative of pupil diameter and change point probability (CPP). **b.** Relationship between pupil diameter and uncertainty. In all panels, horizontal bars indicate $p < 0.05$, corrected for multiple comparisons with FDR correction. Error bars, SEM.

CPP, a latent variable in a model fit to participants' choices, increased rapidly following rule switches (Figure 2f). The temporal derivative of pupil diameter is maximally sensitive to such rapid changes (Murphy et al., 2021), and closely tracks activity in the noradrenergic LC (McGinley et al., 2015a; Reimer et al., 2016). We thus regressed CPP onto the derivative of diameter, while controlling for a range of other variables. Shortly after evidence sample onset, the derivative of diameter indeed encoded CPP (Figure 5a; Table S1). Uncertainty evolved more slowly than CPP (Figure 2f). We therefore expected its signature to become visible in diameter (rather than its derivative). This is indeed what we found (Figure 5b; Table S1). In conclusion, the pupil, a signal that closely matched the dynamics of brainstem neuromodulatory nuclei, tracked the two latent computational variables of the inference process that were sensitive to changes in environmental state (i.e., SR rule).

Discussion

It has long been established that neuromodulatory systems shape the global state and plasticity within the cortex. Much less is known about whether and how the interplay between the brainstem and cortex shapes the neural bases of specific cognitive behaviors – such as context-dependent sensory-motor decisions that we studied here. We found that unpredictable switches in the state of the environment (i.e., SR rule) triggered a boost of arousal: a response within the centers that release neuromodulators, which coincided with a dilation of the pupil. Pupil-linked arousal also encoded rapid fluctuations of key computational variables for detecting hidden rule switches. Finally, boosts in the activity of brainstem arousal systems were followed by stronger coupling of stimulus and action selective patterns within the cortical regions that implement decisions. Taken together, our results are consistent with the idea that brainstem arousal plays an important role in shaping large-scale cortical pathways for flexible cognitive behavior in uncertain environments.

A key aspect of adaptive behavior is that SR-associations must be able to be formed in an arbitrary manner. A number of previous studies on flexible decision-making have been done with non-arbitrary SR-associations (Heinzle et al., 2012; Sarafyazd and Jazayeri, 2019; Duan et al., 2021), where a stimulus is associated with a prepotent response that must be conditionally acted upon or suppressed, such as pro- and anti-saccade tasks (Munoz and Everling, 2004). By contrast, SR-associations in our task were inherently symmetrical, where neither stimulus had a prepotent response. Therefore, default anatomical pathways that establish SR-mapping within the cortex are unlikely to be in place before SR-associations are learned. These arbitrary SR-associations could come about through plasticity mechanisms that reshape existing SR-pathways or form new ones when no default pathways exist. This notion finds support in work in non-human primates and computational modeling, which has suggested that flexible SR-associations are the result of plasticity mechanisms operating on multiple time scales (Fusi et al., 2007).

Our current findings align well with this mechanistic interpretation. In this scenario, synaptic plasticity mechanisms reshape the pathways along which sensory information is directed to motor cortex (Fusi et al., 2007), prompted by neuromodulator release in the cortex in response to sensory evidence in favor of a new active rule. Neuromodulator release elicits plasticity of cortical (Bear and Singer, 1986; Rasmusson, 2000; Berridge and Waterhouse, 2003; Huang et al., 2004; Vetencourt et al., 2008; Marzo et al., 2009; Nadim and Bucher, 2014) and striatal connections involved in learned associations (Reynolds et al., 2001; Reynolds and Wickens, 2002). Moreover, neuromodulator release occurs in response to behaviorally relevant stimulus features (Berridge and Waterhouse, 2003; Aston-Jones and Cohen, 2005; Bouret and Sara, 2005; Sarter et al., 2009). Neuromodulators are thus well suited to restructure SR-association pathways when dictated by the state of the environment. Our findings that arousal responses followed in reaction to rule switches, but preceded the cortical signatures of remapped SR-associations, are consistent with this scenario. Our findings also align with theoretical proposals that link phasic arousal signals to cortical reorganization in response to changes of environmental state (Bouret and Sara, 2005; Dayan and Yu, 2006).

One prominent line of work focuses on the role of frontal and parietal associative regions in flexible SR mapping. Such regions have been found to encode rule information (Miller and Cohen, 2001; Woolgar et al., 2016; van den Brink et al., 2022), and they have been proposed to

flexibly route information from sensory to motor cortices (Miller and Cohen, 2001). These regions may do so by transiently activating in response to the conjunction of stimulus and rule information and acting as switches between the sensory and motor cortices (Cocuzza et al., 2020; Kikumoto and Mayr, 2020; Ito et al., 2022). Other findings indicate that prefrontal cortex encodes key computational variables for performance monitoring and tracking environmental state (Behrens et al., 2007; O'Reilly et al., 2012; McGuire et al., 2014). Frontal and parietal regions are known to innervate the brainstem arousal system (Schwarz and Luo, 2015; Schwarz et al., 2015; Breton-Provencher and Sur, 2019). Our present findings thus raise the possibility that prefrontal association cortex may control flexible SR information flow indirectly, through conveying computational variables such as change-point probability and uncertainty to the brainstem, which in turn sculpt the neural SR pathways accordingly.

The negative response to rule switches that we observed in the BF (Figure 4b) was unexpected, but interesting in light of recent findings. Positive fMRI transients across wide areas of the cortex, and especially in sensory cortices, have been shown to co-occur with drops of activity in the BF region (Liu et al., 2018). In addition, in non-human primates inactivation of the BF results in suppression of correlated fMRI signal fluctuations that are topographically aligned with its afferents (Turchi et al., 2018). Notably, although this region is the main source of cortical acetylcholine (Mesulam and van Hoesen, 1976; Mesulam et al., 1983; Mesulam and Changiz, 1988), it is diverse and also sends prominent long-range GABAergic projections to the cortex (Lin et al., 2015). Thus, it is possible that the here observed suppression of BF activity following rule switches represented a disinhibitory signal (Letzkus et al., 2015) that allowed correlated fluctuations in the cortex to emerge. This possibility is particularly appealing considering the proposed role of disinhibitory signals in flexible information routing within the cortex (Wang and Yang, 2018). Nevertheless, our findings also indicated that increases in BF activity preceded increases in SR-coupling (Figure 4e). It is thus also possible that a suppression of BF activity elicited by rule switches triggered a suppressive effect on SR-coupling within the cortex, and a new instantiation of SR-coupling was brought about by the subsequent activity in the catecholaminergic nuclei. Causal manipulations of specific neuron types within the BF, and concurrent electrophysiological recordings within the cortex, would be well suited to arbitrate between these alternatives.

Due to the nature of the fMRI signal, we cannot definitively distinguish between neuromodulatory signals and those signals originating from other sources. In addition, imaging of nuclei within the brainstem in general is difficult due to their size and proximity to noise sources (Brooks et al., 2013). We have used several methodological approaches to mitigate these issues: by optimizing slice alignment with respect to the brainstem and correcting for physiological noise. We also created masks for the LC on an individual participant basis for optimal localization (Eckert et al., 2010). Finally, we confirmed the expected covariation between signals extracted from individual nuclei and pupillary indices of arousal, for both ongoing signal fluctuations and the more specific rule-evoked responses (Figure S2c, Figure 4d). Together, these choices in experiment design, analysis, and findings, increased the likelihood that the signals we measured are truly related to neuromodulatory nuclei.

Pupil diameter covaried with uncertainty, which was significant before the onset of a new sample of evidence (Figure 5b). Although part of this pre-cue onset covariation can be accounted for by the fact that the prior is the result of sensory evidence that precedes the current sample, we cannot fully exclude that part of this effect is driven by correlation in the values of uncertainty at adjacent cues. In future studies this can be ruled out by using designs with larger lags and jitter between consecutive cues so that protracted pupil responses to

individual cues are more clearly separable. Nonetheless, the key finding that uncertainty covaries with pupil diameter holds regardless of the precise timing with respect to the evidence samples.

In conclusion, during flexible decision-making unpredictable switches in environmental state engage the arousal system. The arousal response in turn is followed by coordinated shifts of coupling between stimulus and action specific activity within the cortical regions that implemented the decision process. Thus, brainstem arousal may orchestrate the association of stimuli with their appropriate actions.

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Author contributions

RLvdB: Conceptualization, Methodology, Software, Formal analysis, Resources, Data Curation, Writing – Original Draft, Writing – Review & Editing, Visualization. **KH:** Methodology, Software, Formal analysis, Data Curation, Writing – Review & Editing. **NW:** Conceptualization, Methodology, Software, Formal analysis, Investigation, Data Curation, Writing – Review & Editing, Supervision. **PRM:** Methodology, Software, Formal analysis, Writing – Review & Editing. **JF:** Methodology, Writing – Review & Editing. **JC-T:** Methodology, Software, Formal analysis, Writing – Review & Editing. **CB:** Conceptualization, Writing – Review & Editing. **THD:** Conceptualization, Methodology, Resources, Writing – Review & Editing, Visualization, Supervision, Project administration, Funding acquisition.

Declaration of interests

The authors declare that no competing interests exist.

Materials and Methods

The current study involves the reanalysis of previously published data (although all presented findings are novel). For a full description of all task parameters and preprocessing, please see van den Brink et al. (2022).

Participants

A total of 22 healthy individuals with normal or corrected vision (median age 27, range 21 – 44, 8 male) took part in our experiment. All participants gave written informed consent and the study was approved by the ethics committee of the Hamburg Medical Association. Four participants were excluded from the current study: one for failure to complete all three sessions, another three for technical reasons (failure to record physiological signals or response data). The final N was thus 18.

Behavioral task

Participants performed two different versions of a flexible decision-making task, of which only one (the ‘inferred rule’ task) is analyzed in the current study. This task involved a basic visual orientation discrimination judgment (lower-order decision), combined with the selection of a volatile sensory-motor (SR) mapping rule (higher-order decision). The SR-mapping rule (Figure 1a) determined the correct action for a given orientation judgment (Figure 2a).

The active rule had to be continuously inferred from a sequence of noisy sensory evidence samples, which were small dots that appeared in rapid succession on the horizontal meridian (Figure 2a). The dots were drawn from one of two generative Gaussian distributions with equal SD and means that were equidistant from the central fixation point but on opposite sides. The generative distribution at any moment determined the active rule. Critically, the active rule could change unpredictably from one sample to the next, with a low probability (hazard rate) of 0.0143. Thus, in order to determine the active rule at a given moment, participants needed to continuously integrate the noisy rule evidence over time.

The appearance of the stimulus for the lower-order decision prompted participants to report their orientation judgement. ITIs for the lower-order decision were long and variable (uniform: 6.8 – 29.6 s). The accuracy of the action depended on both the selection of the correct rule, and on the correct orientation judgment.

MRI data collection

MRI scans were conducted using a Siemens PrismaFit 3T MRI scanner with a 64-channel head-neck coil. We collected 6 runs of the inference task, split over two sessions (T2*-weighted EPI data; Flip angle: 70°; TR: 1.9 s; TE: 28 ms; FOV: 224 x 224 mm², 62 slices (no gap) of 2.0 mm isotropic voxels; 328 volumes), and simultaneously recorded cardiac pulsation and breathing using a pulse oximeter and pneumatic belt. Slices were oriented perpendicular to the rostral-caudal axis of the brainstem as to maximize SNR in this region.

On both sessions, we collected B0 field homogeneity scans (Phase difference and magnitude image: flip angle: 40°; TR: 0.678 s; TE: 5.42/7.88 ms). At the end of the first MRI session, we collected a high resolution T1 neuromelanin sensitive scan, for the purpose of locating individual

participants' LC (T1-TSE; Flip angle: 120°; TR: 675 ms; TE: 12 ms; FOV: 175 x 224 mm², 14 slices (2.0mm, no gap); 0.70x0.70 mm² interpolated to 0.35x0.35 mm²). The slices were oriented perpendicular to the rostral-caudal axis of the brainstem in order to align with the longitudinal extent of the LC. At the end of the second MRI session, we collected a whole-brain T1 anatomical scan (MPRAGE; Flip angle: 9°; TR: 2.3 s; TE: 2.98 ms; TI 1.1 s; FOV: 192 x 256 mm², 240 slices; 1.0 x 1.0 x 1.0 mm).

Pupil recording and preprocessing

Pupil diameter and gaze position were recorded at 1000 Hz with an MRI-compatible EyeLink 1000 eye tracker and calibrated with a 9-point fixation routine. Blinks and missing data segments were linearly interpolated in the eye position data. Other artifacts were identified by the derivative of the pupil diameter exceeding a threshold of 25 pixels and interpolated across. This process was repeated iteratively to ensure all artifacts were identified and removed, in accordance with prior work (van den Brink et al., 2016). All sections containing artifacts were then similarly interpolated across in the gaze x and y position data.

Behavioral modeling

Normative model

The full details of the normative model and fit are provided in van den Brink et al. (2022). In brief, the rule selection (the higher-order decision) process was cast as a dynamic belief updating process (Glaze et al., 2015). In this model, a belief L for one possible state (rule 1) versus the alternative possible state (rule 2) was computed for each sample of evidence. This belief then underwent a non-linear transformation, dependent on a subjective hazard rate \hat{H} , in order to become a prior (ψ) on the presentation of the next sample, thus rendering the model adaptive to volatility of the rule.

Model-derived uncertainty and change-point probability

We used the model fits to produce two key computational parameters associated with each evidence sample: Uncertainty, and change-point probability (CPP). Uncertainty associated with the selected rule was computed as the negative of the magnitude of the prior ψ . CPP is a computational quantity that both the normative belief updating process and human participants are sensitive to in volatile decision-making contexts such as ours (Murphy et al., 2021). CPP was the posterior probability of a change having occurred, given \hat{H} , the previous posterior and the new evidence sample. CPP was computed as follows:

$$CPP = \frac{1}{1 + \Omega} \quad (1)$$

where,

$$\Omega = \frac{1 - \hat{H}}{\hat{H}} \frac{\cosh\left(\frac{1}{2} (LLR_n + L_{n-1})\right)}{\cosh\left(\frac{1}{2} (LLR_n - L_{n-1})\right)} \quad (2)$$

and where n was the current sample, LLR was the sensory evidence in favor of one rule over the other in the form of a log likelihood ratio. Eq. 1 and 2 represent an algebraic rearrangement of the equation for CPP derived by Murphy et al. (2021).

MRI data analysis

Pre-processing of fMRI data

The fMRI data were preprocessed identically to van den Brink et al. (2022). In brief, preprocessing included: motion correction, skull stripping, B0 unwarping, high-pass filtering at 100s, prewhitening, physiological noise correction with retrospective image correction (RETROICOR) (Glover et al., 2000), slice-time correction, and non-linear registration to MNI space.

No spatial smoothing was applied to the fMRI data for two reasons: i) To preserve spatial patterns that were the basis of multi-voxel pattern analysis (see below), and ii) to prevent signal mixing between the brainstem and closely adjacent 4th ventricle (Brooks et al., 2013).

Delineation of ROIs

Regions of interest (ROIs) were defined using a variety of sources (Table 1), including publicly available atlases as well as individually created masks based on neuromelanin sensitive scans. The set of ROIs that we used for all SR coupling-related analyses (see below) presented in this article were selected to span the sensory-motor pathway underlying the lower-order decision (Table 1; Figure 3a). Because the probabilistic masks of neuromodulatory brainstem centers (Zaborszky et al., 2008; Edlow et al., 2012; Murty et al., 2014) varied substantially in their spatial extent, we confined each of these masks to 18 peak probability voxels. Masks of comparable size for the LC were defined for individual participants based on neuromelanin sensitive TSE scans (see below).

Individual delineation of LC ROIs

Masks of the LC for individual participants were created using a semi-automated procedure. First, a mask was manually drawn on the approximate locations of bilateral LC in the T1-TSE scans (Sasaki et al., 2006; Keren et al., 2009), which coincided with the known anatomical location of the LC within the pons, along the floor of the fourth ventricle. An automated algorithm then identified the peak intensity voxel within the manually drawn mask, and an additional 14 contiguous voxels with the highest intensity in each hemisphere. Individual masks (Figure S1) were co-registered to 2 mm isotropic MNI space together with the high-resolution whole-brain T1 scans, max-unit normalized, and thresholded at values of 0.1 to reduce partial voluming effects.

A probabilistic map for the group was defined as the proportion of participants with non-zero values for each voxel in MNI space. This group mask was compared to a previously published mask of the LC (Keren et al., 2009) in order to verify the average location (Figure S2a).

Brainstem time-series extraction and post-processing

Imaging of the brainstem is difficult due to the size of the nuclei involved and their proximity to noise sources such as the 4th ventricle (Brooks et al., 2013; Turker et al., 2021). Physiological

noise in the functional data was mitigated using slice-specific RETROICOR (see: van den Brink et al. (2022)). Nevertheless, in order to maximize the anatomical specificity of the extracted signals, we weighted the time-series of each brainstem nucleus by the probability of voxels within the respective masks. Furthermore, we extracted a probability-weighted time series of voxels within a mask of the 4th ventricle. Variance associated with the ventricular signal was then removed from the signals of the individual brainstem neuromodulatory nuclei using linear regression.

Table 1. ROI definition.

| ROI labels | Functional areas | Source |
|-------------|--|--|
| V1 | Combination of dorsal/ventral parts of V1 | (Wang et al., 2015) |
| V2 | Combination of dorsal/ventral parts of V2 | |
| V3 | Combination of dorsal/ventral parts of V3 | |
| V3A/B | Combination of V3A and V3B | |
| V4 | Human analogue of V4 | |
| LO | Combination of LO1 and LO2 | |
| VO | Combination of VO1 and VO2 | |
| PHC | Combination of PHC1 and PHC2 | |
| MT | Human analogue of MT | |
| MST | Human analogue of MST | |
| IPS | Combination of IPS0-IPS5 | (Glasser et al., 2016) |
| SPL | SPL1 | |
| Caudate | Caudate nucleus | Harvard-Oxford structural Atlas (https://neurovault.org/collections/262/) |
| Putamen | Putamen | |
| Thalamus | Thalamus | |
| PMd | Areas 6a, and 6d | (de Gee et al., 2017) |
| M1 | Somatotopic hand-specific aspect subregion of M1 | (Zaborszky et al., 2008) |
| IPS/PostCeS | Junction of IPS/ and PostCeS | |
| aIPS | Anterior part of horizontal ramus of IPS | |
| LC | Locus coeruleus | This article (Edlow et al., 2012) |
| DR | Dorsal raphe | |
| VTA | Ventral tegmental area | (Murty et al., 2014) |
| SN | Substantia nigra | (Zaborszky et al., 2008) |
| BF | Ch4 region of the basal forebrain | |

Abbreviations: Vx, visual area X; LO, lateral occipital; VO, ventral occipital; PHC, parahippocampal; MST, middle superior temporal; MT, middle temporal; IPS, intraparietal sulcus; SPL, superior parietal lobule; PMd, dorsal premotor cortex; M1 primary motor cortex; PostCeS, postcentral sulcus.

Time-resolved correlation of stimulus and action population codes

We quantified the coupling between population codes for stimulus and action by means of time-resolved decoder output correlation (Anzellotti and Coutanche, 2018), following the methods described in van den Brink et al. (2022). Specifically, we trained sets of support vector machine (SVM) classifiers to decode stimuli (horizontal versus vertical orientation) and action (left versus right hand button press) based on evoked response patterns within cortical and subcortical regions (Figure 3a) that spanned the sensorimotor network.

We then removed feature-specific evoked responses that the decoders were trained on from the data using deconvolution, thus isolating ongoing fluctuations (i.e. activity pattern fluctuations that were not driven by choice grating onset or button presses). We next projected the decision functions (weight vectors) from the stimulus and action decoders onto these estimates of ongoing multi-voxel activity patterns. This produced vectors that compactly summarized ongoing pattern fluctuations for stimulus (S ; for visual cortical regions), or action (A ; for motor cortical and subcortical regions). For example, in V1, the time course S reflected the extent to which patterns of ongoing activity spontaneously tended towards vertical or horizontal orientations (Kenet et al., 2003).

The sign of S and A resulted from the specific decoder design: positive values indicated an ongoing activity pattern that resembled responses to vertical gratings (or left hand button presses) and negative values indicated an ongoing activity pattern that resembled responses to horizontal gratings (or right hand button presses). The schematic in Figure 1b in combination with this decoder design non-trivially predicted a positive correlation of decoder outputs for rule 1 and a negative correlation for rule 2, for the stimulus-action pairs specifically.

We Z-scored S and A across time and computed, for each time point t in the fMRI time series, the degree of covariation between the stimulus and action patterns, via:

$$IC_t = S_t A_t \quad (3)$$

Note that because S and A were Z-scored, IC , if averaged along the time dimension, was identical to the Pearson correlation coefficient of S and A . Therefore, IC represented the correlation of stimulus and action population codes unwrapped along the time dimension.

We tested if IC , averaged across all visual-motor ROI pair combinations, differed between sections of data that were split according to the active rule, using non-parametric permutation testing (10,000 iterations).

Rule switch-evoked brainstem and pupil responses

We used deconvolution to test for the sensitivity of the brainstem and pupil to switch events in the active rule, without making any assumptions about the shape of the hemodynamic or pupillary response (Dale, 1999). Specifically, the estimated response R (of shape $P \times 1$) was obtained via:

$$R = X^+ Y \quad (4)$$

where Y was an $N \times 1$ linearly detrended and z-scored time series (up-sampled fMRI data, or pupil diameter down-sampled to 50 Hz). X was an $N \times P$ design matrix, with value 1 in every element along those diagonals corresponding to the occurrence of switches in active rule, $^+$ denoted the pseudoinverse, and P was peri-stimulus time.

The deconvolved responses were compared to chance (zero) and corrected for multiple comparisons across time points using non-parametric cluster-based permutation testing (10,000 iterations, two-tailed) (Maris and Oostenveld, 2007).

Correlation between brainstem activity and pupil diameter

We assessed the correlation between brainstem and pupil activity in two ways: i) by correlating estimates of single-switch responses in the brainstem and in the pupil (Figure 4c), and ii) by correlation of the entire brainstem activity and pupil time series (Figure S2c).

Single-switch response amplitude scalars for individual brainstem nuclei were computed by regression of the individually estimated HRF (see section: *Rule switch-evoked brainstem and pupil responses*) onto the single rule switch locked data. Similarly, we computed single-switch responses in the pupil via regression, using the standard pupillary response function described by Hoeks and Levelt (1993), with a peak latency set for individual participants according to the deconvolved pupil response. The resulting vectors of response amplitude scalars were then correlated to estimate covariation in evoked responses in the brainstem and pupil.

Whole time-series correlation (Figure S2c) between the brainstem signals and pupil signal was done including a correction for HRF and pupillary response delays. That is, we estimated a peak HRF latency for individual participants from the stimulus-evoked response in V1 (van den Brink et al., 2022), and shifted the pupil signal forward by this latency. Because the pupil signal itself also lags neural activity, we subsequently shifted the pupil signal back by 1 s, following prior work (Yellin et al., 2015).

The significance of both single-switch response covariation as well as covariation of the full time series was assessed with non-parametric permutation testing (10,000 iterations). We tested one-tailed because we expected positive covariation of activity in the pupil and brainstem.

Cross-correlation between stimulus-action code covariation, brainstem activity, and pupil

We used cross-correlation to examine temporal relationships between stimulus-action code covariation (computed via Eq. 3) and two other signals: activity in the brainstem, and pupil diameter. We used cross-correlation because the temporal sequence of activity fluctuations may be indicative of directional relationships between these signals. That is, if the activity in the brainstem modulates coupling at the level of the cortex, then we expect the brainstem to become active prior to an increase in cortical coupling.

Stimulus-action code covariation was a signed signal, where the sign depended on the active task rule. We expected the brainstem to modulate the magnitude of this signal, regardless of sign, so in all related analyses we used the absolute of stimulus-action code covariation.

Because the pupil signal lags putative neural activity that underlies it, we corrected for this delay by shifting the pupil signal backward by the expected latency of 1s (Yellin et al., 2015), and

aligned it with the hemodynamic signal by shifting forward by the observed hemodynamic delay (identically to the procedure described in the section: *Correlation between brainstem activity and pupil diameter*). Stimulus-action code covariation and brainstem activity were both hemodynamic signals in origin, so no lag correction was required before cross-correlation.

Correlation coefficients of the cross-correlations were compared to chance (zero) and corrected for multiple comparisons across time points using non-parametric cluster-based permutation testing (10,000 iterations, two-tailed) (Maris and Oostenveld, 2007).

Relationship between pupil dynamics and model computational variables

We examined the relationship between pupil and behavior with linear regression of the variables resulting from the computational model fit to individual participants' behavior onto (the derivative of) pupil diameter. The regression had the following form:

$$\frac{dPupil(\tau)}{d\tau}_{t,c} = \beta_{0,t} + \sum_{l=-1}^0 (\beta_{1,l,t} \times CPP_{c+l} + \beta_{2,l,t} \times (-|\Psi_{c+l}|) + \beta_{3,l,t} \times |LLR_{c+l}|) + \beta_{4,t} \times |LLR_c - LLR_{c-1}| + \beta_{5,t} \times gaze_{x,t,c} + \beta_{6,t} \times gaze_{y,t,c} + \beta_{7,t} \times baseline_c \quad (5)$$

where t was time relative to cue (evidence sample) onset, c indexed cues, $\frac{dPupil(\tau)}{d\tau}_{t,c}$ gave the value of the pupil derivative time series at time t relative to the onset of cue c . CPP was change point probability, $-|\Psi|$ was uncertainty, LLR was the sensory evidence (i.e., the log-likelihood ratio), $gaze_x$ and $gaze_y$ were gaze positions on screen, and baseline was average pupil diameter in a 1 s window preceding cue onset. The terms relating to prior cues were included to account for auto-correlation in the pupil response and isolate responses to the current cue. The magnitude of the difference between the current and previous cue LLR was included to exclude that any relationship between pupil and CPP was driven by low-level visual differences between consecutive cues.

We also fit a variant of this regression model to baseline corrected pupil diameter (rather than the derivative). This regression model did not include baseline diameter, or terms for preceding cues:

$$Pupil_{t,c} = \beta_{0,t} + \beta_{1,t} \times CPP_c + \beta_{2,t} \times (-|\Psi_c|) + \beta_{3,t} \times |LLR_c| + \beta_{4,t} \times gaze_{x,t,c} + \beta_{5,t} \times gaze_{y,t,c} \quad (6)$$

Based on prior work (Murphy et al., 2021), we expected CPP following cue onset to covary positively with the derivative of pupil diameter. We also expected uncertainty to covary positively with pupil diameter.

Regression coefficients were compared to chance (zero) using non-parametric permutation testing (10,000 iterations, one-tailed) and corrected for multiple comparisons using the false discovery rate (FDR).

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Supplementary Figures and Tables

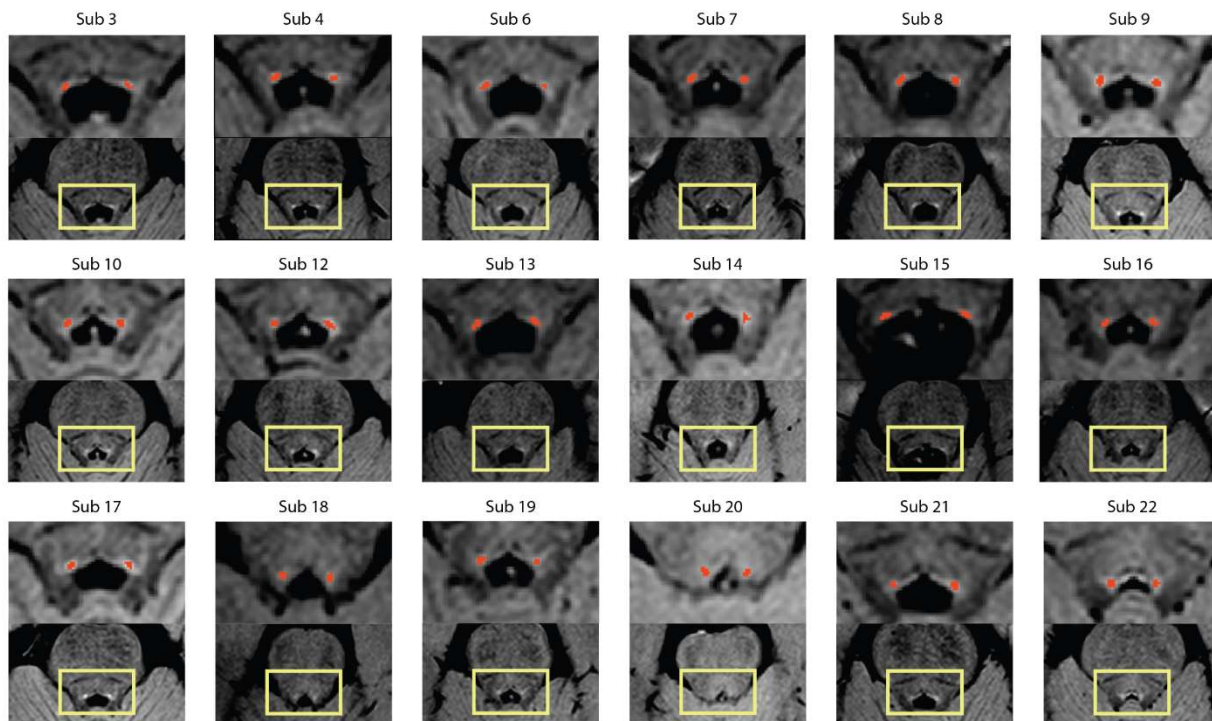


Figure S1. Neuromelanin sensitive TSE scans for all participants. The mask of the locus coeruleus (LC) is overlaid in red. The masks were created by manually selecting the maximal voxel within the LC region on each side, and subsequently automatically selecting 15 additional contiguous voxels with the highest intensity.

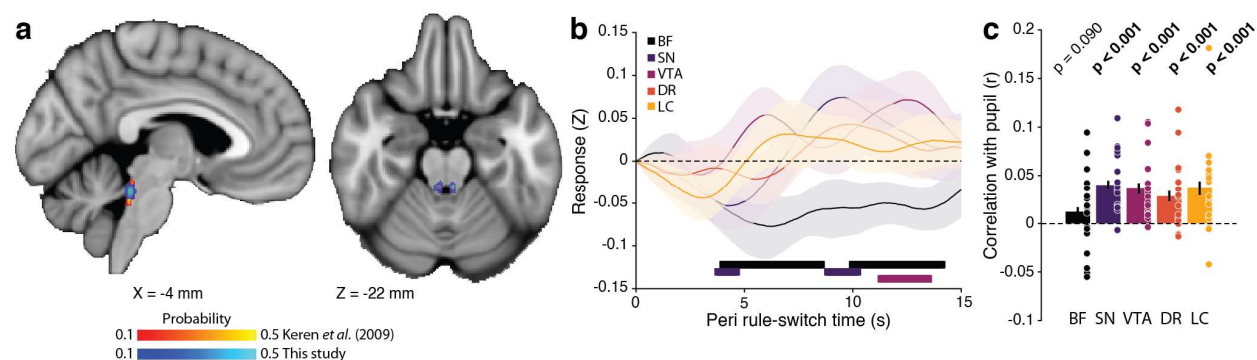


Figure S2. **a.** Overlap of LC mask of the current study with that of Keren *et al.*, (2009). **b.** Deconvolved response in the brainstem evoked by rule switches, shown for all individual nuclei. **c.** Correlation between brainstem signals and pupil diameter, across time. Covariation between the rule switch-evoked response magnitude between the brainstem and pupil. Dots, individual participants. Bars, group average. Error bars, SEM. Horizontal bars indicate $p < 0.05$, corrected for multiple comparisons with cluster-based permutation testing.

Table S1. Statistics summary.

| Figure | Label | Test | Tail | <i>p</i> -value | Cohen's <i>d</i> | 95% CI |
|------------|----------------------|-------------|------|-----------------|------------------|-----------------|
| 2b | Rule 1 | Permutation | Two | <0.001 | 3.410 | 18.798 - 40.995 |
| 2b | Rule 2 | Permutation | Two | <0.001 | 4.511 | 17.669 - 38.224 |
| 2b | Rule 1 vs rule 2 | Permutation | Two | 0.329 | 0.212 | -1.331 - 5.222 |
| 2c | Rule 1 vs rule 2 | Permutation | Two | 0.087 | -0.385 | -0.028 - 0 |
| 3e | Rule 1 | Permutation | One | 0.003 | 0.581 | 0.002 - 0.012 |
| 3e | Rule 2 | Permutation | One | <0.001 | -0.955 | -0.015 - -0.005 |
| 3e | Rule 1 vs rule 2 | Permutation | One | <0.001 | 1.005 | 0.008 - 0.027 |
| 3f (left) | Rule 1 | Permutation | Two | <0.001 | 3.408 | 0.042 - 0.093 |
| 3f (left) | Rule 2 | Permutation | Two | <0.001 | 3.800 | 0.043 - 0.094 |
| 3f (left) | Rule 1 vs rule 2 | Permutation | Two | 0.554 | -0.133 | -0.005 - 0.002 |
| 3f (right) | Rule 1 | Permutation | Two | <0.001 | 2.753 | 0.051 - 0.113 |
| 3f (right) | Rule 2 | Permutation | Two | <0.001 | 3.068 | 0.057 - 0.125 |
| 3f (right) | Rule 1 vs rule 2 | Permutation | Two | 0.074 | -0.395 | -0.019 - 0 |
| 4b | BF | Permutation | Two | 0.021 | 0.539 | 0.014 - 0.124 |
| | | | | 0.012 | 0.593 | 0.014 - 0.094 |
| 4b | SN, VTA, LC (pooled) | Permutation | Two | 0.018 | 0.523 | 0.007 - 0.072 |
| | | | | 0.029 | 0.474 | 0.001 - 0.02 |
| | | | | 0.037 | 0.449 | 0.002 - 0.06 |
| 4c | Pupil | Permutation | One | 0.044 | 0.429 | 0.002 - 0.106 |
| 4d | BF | Permutation | One | 0.946 | -0.354 | -0.076 - 0.001 |
| 4d | SN | Permutation | One | 0.115 | 0.260 | -0.012 - 0.081 |
| 4d | VTA | Permutation | One | 0.030 | 0.416 | 0.003 - 0.063 |
| 4d | DR | Permutation | One | 0.461 | 0.019 | -0.036 - 0.041 |
| 4d | LC | Permutation | One | 0.013 | 0.501 | 0.013 - 0.094 |
| 4e | BF | Permutation | One | 0.036 | 0.455 | 0.001 - 0.022 |
| 4e | SN, VTA, LC (pooled) | Permutation | One | 0.002 | 0.736 | 0.004 - 0.019 |
| 4f | Pupil | Permutation | One | <0.001 | 0.966 | 0.009 - 0.032 |
| | | | | 0.041 | 0.432 | 0 - 0.02 |
| 5a | CPP | Permutation | One | <0.001 | 4.911 | 0.03 - 0.037 |
| 5b | - η^2 | Permutation | One | <0.001 | 4.829 | 0.031 - 0.034 |

Cohen's *d*: Measure of effect size. $d > 0.2$: small effect size; $d > 0.5$: medium; $d > 0.8$: large; $d > 1.2$: very large; $d > 2.00$: huge. **CI:** confidence interval around mean (for comparisons to chance), or around difference of means (for comparisons between conditions). For figure panels that show multiple significant time points as bars, we report statistics for the data averaged across time, per significant cluster.

Abbreviations: BF, basal forebrain; SN, substantia nigra; VTA, Ventral tegmental area; DR, Dorsal raphe; LC, locus coeruleus; CPP, change point probability.