Accepted Manuscript

Functional brain networks for learning predictive statistics

Joseph Giorgio, Vasilis M. Karlaftis, Rui Wang, Yuan Shen, Peter Tino, Andrew Welchman, Zoe Kourtzi

PII: S0010-9452(17)30271-X

DOI: 10.1016/j.cortex.2017.08.014

Reference: CORTEX 2097

To appear in: *Cortex*

Received Date: 17 May 2017

Revised Date: 1 August 2017

Accepted Date: 3 August 2017

Please cite this article as: Giorgio J, Karlaftis VM, Wang R, Shen Y, Tino P, Welchman A, Kourtzi Z, Functional brain networks for learning predictive statistics, *CORTEX* (2017), doi: 10.1016/j.cortex.2017.08.014.

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.



Functional brain networks for learning predictive statistics

Joseph Giorgio^a, Vasilis M Karlaftis^a, Rui Wang^b, Yuan Shen^c, Peter Tino^d, Andrew Welchman^a, Zoe Kourtzi^a

^a Department of Psychology, University of Cambridge, Cambridge, United Kingdom

^b Key Laboratory of Mental Health, Institute of Psychology, Chinese Academy of Sciences,

Beijing, China

^c Department of Mathematical Sciences, Xi'an Jiaotong-Liverpool University, Suzhou, China

^d School of Computer Science, University of Birmingham, Birmingham, United Kingdom

Correspondence: Zoe Kourtzi Department of Psychology University of Cambridge Cambridge, UK Email: <u>zk240</u>@cam.ac.uk

Abstract

Making predictions about future events relies on interpreting streams of information that may initially appear incomprehensible. This skill relies on extracting regular patterns in space and time by mere exposure to the environment (i.e. without explicit feedback). Yet, we know little about the functional brain networks that mediate this type of statistical learning. Here, we test whether changes in the processing and connectivity of functional brain networks due to training relate to our ability to learn temporal regularities. By combining behavioral training and functional brain connectivity analysis, we demonstrate that individuals adapt to the environment's statistics as they change over time from simple repetition to probabilistic combinations. Further, we show that individual learning of temporal structures relates to response strategy. Our fMRI results demonstrate that learning-dependent changes in fMRI activation within and functional connectivity between brain networks relate to individual variability in strategy. In particular, extracting the exact sequence statistics (i.e. matching) relates to changes in brain networks known to be involved in memory and stimulus-response associations, while selecting the most probable outcomes in a given context (i.e. maximizing) relates to changes in frontal and striatal networks. Thus, our findings provide evidence that dissociable brain networks mediate individual ability in learning behaviorally-relevant statistics.

Keywords: brain plasticity, fMRI, functional network connectivity, individual differences, statistical learning.

1. Introduction

Successful interactions in a new environment entail interpreting initially incomprehensible streams of information and making predictions about upcoming events. The brain is thought to succeed in this challenge by finding regular patterns and meaningful structures that help us to predict and prepare for future actions. This skill is thought to rely on our ability to extract spatial and temporal regularities, often with minimal explicit feedback (Perruchet and Pacton, 2006; Aslin and Newport, 2012). For example, previous behavioral studies have shown that structured patterns become familiar after simple exposure to items (shapes, tones or syllables) that co-occur spatially or follow in a temporal sequence (Saffran et al., 1996; Saffran et al., 1999; Chun, 2000; Fiser and Aslin, 2002; Turk-Browne et al., 2005).

Functional imaging studies have identified key brain regions involved in the learning of statistical regularities. In particular, striatal and hippocampal regions have been implicated in the learning of temporal sequences (Rauch et al., 1997; Schendan et al., 2003; Aizenstein et al., 2004; Gheysen et al., 2011; Rose et al., 2011; Hsieh et al., 2014). Further, the medial temporal cortex has been implicated in learning of probabilistic associations (Turk-Browne et al., 2010; Schapiro et al., 2012). However, we know little about the functional brain networks and their interactions that mediate statistical learning.

Recent functional connectivity studies provide accumulating evidence for learningdependent changes in human brain networks due to training in a range of tasks including visual perceptual learning (Lewis et al., 2009; Baldassarre et al., 2012), motor learning (Sun et al., 2007; Bassett et al., 2011; Ma et al., 2011), auditory learning (Ventura-Campos et al., 2013) and language learning (Veroude et al., 2010). These studies typically involve prolonged training with feedback. Here we ask whether mere exposure to streams of information (i.e. without trial-by-trial feedback) changes processing in functional brain networks that mediate our ability to extract statistical regularities.

We combine behavioral measurements and multi-session fMRI (before and after training) to investigate processing in functional brain networks that mediate statistical learning of temporal structures. Event structures in the natural environment typically contain regularities at different scales from simple repetition to probabilistic combinations. To investigate the brain networks involved in extracting such structures unencumbered by past experience, we generated temporal sequences based on Markov models of different orders (i.e. context lengths of 0 or 1 previous item) (Figure 1). We exposed participants to sequences of unfamiliar symbols and varied the sequence structure unbeknownst to the participants by increasing the context length. To facilitate learning, sequences were first determined by frequency statistics (i.e. occurrence probability per symbol), and then by context-based statistics (i.e. the probability of a given symbol appearing depends on the preceding symbol). Participants performed a prediction task, indicating which symbol they expected to appear next in the sequence. Following previous statistical learning paradigms, participants were exposed to the sequences without trial-by-trial feedback. We tested for improvement in the prediction task and fMRI activation changes in functional networks due to training (i.e. before vs. after training on frequency and context-based statistics).

Further, we asked whether learning-dependent changes in functional brain networks relate to the participants' ability to learn temporal structures. Previous work (Shanks et al., 2002; Erev and Barron, 2005; Lagnado et al., 2006; Wozny et al., 2010; Eckstein et al., 2013; Acerbi et al., 2014; Murray et al., 2015) has highlighted the role of strategies in probabilistic learning and decision making and suggests that previous experience shapes the selection of response strategies (Rieskamp and Otto, 2006; Fulvio et al., 2014). That is, observers are shown to match their choices stochastically according to the underlying input statistics or maximize their reward by selecting the most probable positively rewarded outcomes. Here,

we tested whether learning-dependent changes in functional brain networks relate to the participants' response strategy when learning frequency and context-based statistics.

Our behavioral results show that individuals adapt to the environment's statistics; that is they are able to extract predictive structures that change over time. Further, we show that individual learning of structures relates to response strategy; that is individuals differed in their response strategies, favoring probability maximization (i.e. extracting the most probable outcome in a given context) or matching the exact sequence statistics. We used this variability in response strategy to interrogate fMRI activity in functional brain networks. Our results demonstrate that distinct brain networks mediate these two strategies. In particular, learning-dependent fMRI changes in functional brain networks relate to individual variability in response strategy: matching relates to fMRI activation changes in brain networks involved in memory and stimulus-response associations (including precuneus, sensorimotor, middle temporal and the right central executive), while maximizing relates to activation changes in frontal and striatal brain networks (including basal ganglia and the left central executive). Further, increased functional connectivity due to training between networks involved in memory and stimulus-response associations relates to matching, while between frontal and striatal networks relates to maximization. Thus, our findings provide evidence for distinct functional brain networks that mediate individual ability to extract behaviorally-relevant statistics in variable environments.

Figure 1

2. Material and Methods

2.1 Observers

Twenty-three participants (mean age = 21.8 years) were tested in multiple scanning and behavioral training sessions. The data from four participants were excluded from further

imaging analysis due to excessive head movement or signal variance. A single run from six of the remaining nineteen participants was also removed due to excessive head movement or signal variance. All participants were naive to the aim of the study, had normal or correctedto-normal vision and signed an informed consent. This study was conducted in the School of Psychology, Birmingham and was approved by the University of Birmingham Ethics Committee.

2.2 Stimuli

Stimuli comprised four symbols chosen from Ndjuká syllabary (**Figure 1a**). These symbols were highly discriminable from each other and were unfamiliar to the participants. Each symbol subtended 8.5° of visual angle and was presented in black on a mid-grey background. Experiments were controlled using Matlab and the Psychophysics toolbox 3 (Brainard, 1997; Pelli, 1997). For the behavioral training sessions, stimuli were presented on a 21-inch CRT monitor (ViewSonic P225f 1280 x 1024 pixel, 85 Hz frame rate) at a distance of 45 cm. For the pre and post-training fMRI scans, stimuli were presented using a projector and a mirror set-up (1280 x 1024 pixel, 60 Hz frame rate) at viewing distance of 67.5 cm. The physical size of the stimuli was adjusted so that angular size was constant during behavioral and scanning sessions.

2.3 Sequence design

To generate probabilistic sequences that differed in their structure, we used a temporal Markov model and manipulated the memory order of the sequence, which we refer to as the context length. The Markov model consists of a series of symbols, where the symbol at time *i* is determined probabilistically by the previous '*k*' symbols. We refer to the symbol presented at time *i*, s(i), as the *target* and to the preceding *k*-tuple of symbols (s(i-1), s(i-2), ..., s(i-k)) as the *context*. The value of '*k*' is the order or level of the sequence:

$$P(s(i) | s(i-1), s(i-2), ..., s(1)) = P(s(i) | s(i-1), s(i-2), ..., s(i-k)), k < i$$

In our study, we used two levels of memory length; for k=0,1. The simplest $k=0^{th}$ order model is a memory-less source. This generates, at each time point *i*, a symbol according to symbol probability P(s), without taking account of the previously generated symbols. The order k=1 Markov model generates symbol s(i) at each time *i* conditional on the previously generated symbol s(i-1). This introduces a memory in the sequence; that is, the probability of a particular symbol at time *i* strongly depends on the preceding symbol s(i-1). Unconditional symbol probabilities P(s(i)) for the case k=0 are replaced with conditional ones, P(s(i)|s(i-1)).

At each time point, the symbol that follows a given context is determined probabilistically, making the Markov sequences stochastic. The underlying Markov model can be represented through the associated context-conditional target probabilities. We used 4 symbols that we refer to as stimuli A, B, C and D. The correspondence between stimuli and symbols was counterbalanced across participants.

For level-0, the Markov model was based on the probability of symbol occurrence: one symbol had a high probability of occurrence, one low probability, while the remaining two symbols appeared rarely (**Figure 1b**). For example, the probabilities of occurrence for the four symbols A, B, C, and D were 0.18, 0.72, 0.05 and 0.05, respectively. Presentation of a given symbol was independent of the items that preceded it. For level-1, the target depended on the immediately preceding stimulus (**Figure 1b**). Given a context (the last seen symbol), only one of two targets could follow; one had a high probability of being presented and the other a low probability (e.g., 80% vs. 20%). For example, when Symbol A was presented, only symbols B or C were allowed to follow, and B had a higher probability of occurrence than C.

To test whether participants adapt to changes in the temporal structure, we ensured that the sequences across levels were matched for properties other than context-length. That is, sequences across levels were matched for the number of symbols presented (i.e. all four

symbols were presented for both level-0 and level-1 sequences). To ensure that for level-1 participants learned context-target contingences rather than individual symbols, all symbols in level-1 were presented with equal frequency (i.e. marginal probability of each symbol was 0.25). These constraints resulted in differences in the probability distributions between level-0 and level-1. However, we designed the stochastic sources from which the sequences were generated so that the context-conditional uncertainty remained highly similar across levels. In particular, for the zero-order source, only two symbols were likely to occur most of the time; the remaining two symbols had very low probability (0.05); this was introduced to ensure that there was no difference in the number of symbols presented across levels. Of the two dominant symbols, one was more probable (probability 0.72) than the other (probability 0.18). This structure is preserved in Markov chain of order 1, where conditional on the previous symbol, only two symbols were allowed to follow, one with higher probability (0.80) than the other (0.20). This ensures that the structure of the generated sequences across levels differed predominantly in memory order (i.e. context length) rather than context-conditional probability.

2.4 Procedure

Participants were initially familiarized with the task through a brief practice session (8 minutes) with random sequences (i.e. all four symbols were presented with equal probability 25% in a random order). Following this, participants took part in multiple behavioral training and fMRI scanning sessions that were conducted on different days. Participants were trained with structured sequences and tested with both structured and random sequences to ensure that training was specific to the trained sequences.

In the first scanning session, participants were presented with zero- and first-order sequences and random sequences. Participants were then trained with zero-order sequences, and subsequently with first-order sequences. For each level, participants completed a

minimum of 3 and a maximum of 5 training sessions (840-1400 trials). Training at each level ended when participants reached plateau performance (i.e. performance did not change significantly for two sessions). A post-training scanning session followed training per level (i.e. on the following day after completion of training) during which participants were presented with structured sequences determined by the statistics of the trained level and random sequences (90 trials each). The mean time interval (\pm standard deviation) between the pre-training and the post-training test sessions was 21.6 (\pm 3.3) days.

2.5 Psychophysical training

Each training session comprised five blocks of structured sequences (56 trials per block) and lasted one hour. To ensure that sequences in each block were representative of the Markov model order per level, we generated 10,000 Markov sequences per level comprising 672 stimuli per sequence. We then estimated the Kullback-Leibler divergence (KL divergence) between each example sequence and the generating source. In particular, for level-0 sequences this was defined as:

$$KL = \sum_{target} Q(target) \log(\frac{Q(target)}{P(target)}),$$

and for level-1 sequences this was defined as:

 $KL = \sum_{context} Q(context) \sum_{target} Q(taget|context) \log(\frac{Q(target|context)}{P(target|context)}),$

where P() refers to probabilities or conditional probabilities derived from the presented sequences, and Q() refers to those specified by the source. We selected fifty sequences with the lowest KL divergence (i.e. these sequences matched closely the Markov model per level). The sequences presented to the participants during the experiments were selected randomly from this sequence set.

For each trial, a sequence of 8-14 stimuli appeared in the center of the screen, one at a time in a continuous stream, each for 300ms followed by a central white fixation dot (ISI) for 500ms (**Figure 1a**). This variable trial length ensured that participants maintained attention

during the whole trial. Each block comprised equal number of trials with the same number of stimuli. The end of each trial was indicated by a red dot cue that was presented for 500ms. Following this, all four symbols were shown in a 2x2 grid. The positions of test stimuli were randomized from trial to trial. Participants were asked to indicate which symbol they expected to appear following the preceding sequence by pressing a key corresponding to the location of the predicted symbol. Participants learned a stimulus-key mapping during the familiarization phase: key '8', '9', '5' and '6' in the number pad corresponded to the four positions of the test stimuli - upper left, upper right, lower left and lower right, respectively.

After the participant's response, a white circle appeared on the selected item for 300ms to indicate the participant's choice, followed by a fixation dot for 150ms (ITI) before the start of the next trial. If no response was made within 2s, a null response was recorded and the next trial started. Participants were given feedback (i.e. score in the form of Performance Index, see Behavioral analysis) at the end of each block –rather than per-trial error feedback– that motivated them to continue with training.

2.6 Scanning sessions

The pre-training scanning session (Pre) included six runs (i.e. three runs per level) the order of which was randomized across participants. Scanning sessions after training per level (denoted as Post-0, Post-1) included nine runs of structured sequences determined by the same statistics as the corresponding trained level and random sequences. Each run comprised five blocks of structured and five blocks of random sequences presented in a random counterbalanced order (2 trials per blocks; a total of 10 structured and 10 random trials per run), with an additional two 16s fixation blocks, one at the beginning and one at the end of each run. The trial design was adjusted to afford modeling of fMRI signals within the scanning timing constraints. In particular, each trial comprised a sequence of 10 stimuli that were presented for 250ms each, separated by a blank interval during which a white fixation

dot was presented for 250ms. Following the sequence, a response cue (central red dot) appeared on the screen for 4s before the test display (comprising four test stimuli) appeared for 1.5s. Participants were asked to indicate which symbol they expected to appear following the preceding sequence by pressing a key corresponding to the location of the predicted symbol. A white fixation was then presented for 5.5s before the start of the next trial. In contrast to the training sessions, no feedback was given during scanning.

2.7 fMRI data acquisition

The experiments were conducted at the Birmingham University Imaging Centre using a 3T Philips Achieva MRI scanner. T2*-weighted functional and T1-weighted anatomical (175 slices; $1 \times 1 \times 1$ mm³ resolution) data were collected with a 32-channel head coil. Echo planar imaging (EPI) data (gradient echo-pulse sequences) were acquired from 32 slices (whole brain coverage; duration=6min; TR=2s; TE=35 ms; $2.5 \times 2.5 \times 4$ mm³ resolution; SENSE).

2.8 Behavioral analysis

2.8.1 Performance index

We assessed participant responses in a probabilistic manner. We computed a performance index per context that quantifies the minimum overlap (min: minimum) between the distribution of participant responses and the distribution of presented targets estimated across 56 trials per block by:

 $PI(context) = \sum_{target} min (P_{resp}(target|context), P_{pres}(target|context))$

where the sum is over targets from the symbol set A, B, C and D.

The overall performance index is then computed as the average of the performance indices across contexts, PI(context), weighted by the corresponding stationary context probabilities:

$$PI = \sum_{context} PI(context) \cdot P(context)$$

To compare across different levels, we defined a normalized PI measure that quantifies participant performance relative to random guessing. We computed a random guess baseline; i.e. performance index PI_{rand} that reflects participant responses to targets with a) equal probability of 25% for each target per trial for level-0 ($PI_{rand} = 0.53$); b) equal probability for each target for a given context for level-1 ($PI_{rand} = 0.45$). To correct for differences in random-guess baselines across levels, we subtracted the random guess baseline from the performance index ($PI_{normalized} = PI - PI_{rand}$).

2.8.2 Strategy choice and strategy index

To quantify each participant's strategy, we compared individual participant response distributions (response-based model) to two baseline models: (i) probability matching, where probabilistic distributions are derived from the Markov models that generated the presented sequences (Model-matching) and (ii) a probability maximization model, where only the single most likely outcome is allowed for each context (Model-maximization). We used Kullback-Leiber (KL) divergence to compare the response distribution to each of these two models. KL is defined as follows:

$$KL = \sum_{target} M(target) \log(\frac{M(target)}{R(target)})$$

for level-0 model, and

$$KL = \sum_{context} M(context) \sum_{target} M(target|context) \log(\frac{M(target|context)}{R(target|context)})$$

for level-1 model, where R() and M() denote the probability distribution or conditional probability distribution derived from the human responses and the models (i.e. probability matching or maximization) respectively, across all the conditions.

We quantified the difference between the KL divergence from the response-based model to Model-matching and the KL divergence from the response-based model to Model-maximization. We refer to this quantity as strategy choice indicated by Δ KL(Model-

maximization, Model-matching). We computed strategy choice per training block, resulting in a strategy curve across training for each individual participant. We then derived an individual strategy index by calculating the integral of each participant's strategy curve and subtracting it from the integral of the exact matching curve, as defined by Model-matching across training. We defined the integral curve difference (ICD) between individual strategy and exact matching as the individual strategy index. Negative strategy index indicates a strategy closer to matching, while positive index indicate a strategy closer to maximization.

2.9 fMRI data analysis

2.9.1 Data pre-processing

We pre-processed the fMRI data in Matlab R2013a and SPM12 software package (http://www.fil.ion.ucl.ac.uk/spm/software/spm12/) following the pipeline described in (Taylor et al., 2015). We first processed the T1 weighted anatomical images by applying brain extraction and segmentation. From the segmented T1 we created a white matter (WM) mask and a cerebrospinal fluid (CSF) mask. For each fMRI run, we corrected the EPI data for slice scan timing (i.e. to remove time shifts in slice acquisition) and motion (least squares correction). We co-registered each run to the T1 image and calculated the mean CSF and WM signal per volume. We then aligned the T1 image to MNI space (affine transformation) and applied the same transformation to the EPI data. Finally, we resliced the aligned EPI data to 3 x 3 x 4 mm³ resolution and applied spatial smoothing with a 5mm isotropic FWHM Gaussian kernel.

2.9.2 Independent Component Analysis (ICA)

We used group spatial ICA (GICA) (McKeown et al., 1998; Haberecht et al., 2001; Calhoun et al., 2009; Calhoun and Adali, 2012) to extract participant- and session-specific hemodynamic source locations using the Group ICA fMRI Toolbox (GIFT) (<u>http://mialab.mrn.org/software/gift/</u>). Pre-training sessions comprised of 3 runs, whereas

post-training sessions comprised of 9 runs. To account for the difference in number of runs between sessions, we matched the post-training session runs to the pre-training session in their acquisition order and therefore, we included the matched 3 runs for subsequent analyses. Following pre-training processing of each run, we performed intensity normalization and dimensionality reduction. We used the Minimum Description Length criteria (MDL) (Rissanen, 1978) to estimate the dimensionality and determine the number of independent components. We used a two-level dimensionality reduction procedure using PCA; first at the participant level and then at the group level. The ICA estimation was run 20 times and the component stability was estimated using ICASSO (Himberg et al., 2004).

This procedure resulted in 28 independent components. We generated participant- and session-specific spatial maps and timecourses for each component using GICA3 back reconstruction. Participant spatial maps were not scaled and, as intensity normalization was performed prior to ICA, they represent percent signal change. For further analysis, we extracted the timecourse per participant per component and regressed out the six motion parameters (translation and rotation) as well as the mean WM and CSF signal. We then removed slow drifts by applying linear detrending on the regressed timecourse (Van Dijk et al., 2010).

2.9.3 Component selection

We used a quantitative method, as described in (Stevens et al., 2007), to remove components of non-neuronal origin. We first converted each component's spatial map to a z-map and thresholded it at z=1.96 to calculate its spatial correlation with grey matter (GM) and CSF probabilistic maps (as extracted from the MNI template). We rejected any component with a spatial correlation of $R^2 > 0.025$ with CSF or WM and of $R^2 < 0.025$ with GM. To supplement this method, we visually inspected all rejected components to verify that they were not of neuronal origin. This method resulted in 13 rejected components: 7 components had a high

spatial correlation with CSF, 1 component had a high correlation with WM and 5 components had a low correlation with GM.

We labeled the selected components based on spatial correlation with known restingstate networks, as the brain's functional architecture at rest has been shown to relate to taskbased networks (Fox and Raichle, 2007; Smith et al., 2009). We correlated the thresholded spatial maps with network templates (Allen et al., 2011) and labeled each component based on its highest correlation value to the network templates. In further analysis, we used only the selected components. To further denote the areas included in each selected component, we created participant-specific maps per component by averaging the maps across runs and sessions per participant. We then generated a group map based on one sample t-test on the participant average map (FWER corrected at p<0.005). We visualized the significant clusters in xjView toolbox (<u>http://www.alivelearn.net/xjview</u>) and labeled them based on their peak voxels (**Table 1**).

2.9.4 GLM-based analysis

We generated a GLM event-related (epoch) design and ran a multiple regression analysis on each component's timecourse (treated for nuisance variables: 6-motion, CSF and WM) per participant per run. The GLM design was composed of random and structured trial blocks convolved with the hemodynamic response function. The output of the regression is a set of β weights (i.e. parameter estimates) for the task conditions (random, structured sequences); where the β weights represent the degree to which the component timecourse is modulated by each task condition. We then averaged the β weights of each task condition across runs resulting in a single value for each condition per participant per component per session.

To test whether component activation changes in relation to individual behavior (i.e. strategy), we correlated strategy for frequency (level-0) and context-based (level-1) statistics with change of β weight (i.e. post- minus pre-training) per component, separately for each

task condition (random, structured). We used the Robust Correlation Toolbox (Pernet et al., 2013) and Pearson's skipped correlation to account for potential outliers. We accepted as significant the correlations where the bootstrapped 95% confidence interval (CI) after 1000 permutations doesn't cross the zero origin.

2.9.5 Functional Network Connectivity (FNC)

To investigate the functional interaction of the networks identified in the analysis of temporal dynamics, we calculated the between network connectivity of these components (Jafri et al., 2008). We defined as FNC the correlation of each component's timecourse (after nuisance regression and detrending) with every other component's timecourse, per participant. We converted the correlation coefficients to z-scores (Fisher z-transform) and averaged the values across runs for each pair of components; deriving one connectivity value per participant per session. We then correlated the change in average z-score (post- minus pre-training) with strategy for frequency (level-0) and context-based (level-1) statistics using the Robust Correlation method.

3. Results

3.1 Behavioral performance

Previous studies have compared learning of different spatiotemporal contingencies in separate experiments across different participant groups (Fiser and Aslin, 2002; Fiser and Aslin, 2005). Here, to investigate whether individuals extract changes in structure, we presented the same participants with sequences that changed in structure unbeknownst to them (**Figure 1a**). We parameterized sequence structure based on the memory-order of the Markov models used to generate the sequences (see 2.3); that is, the degree to which the presentation of a symbol depended on the history of previously presented symbols (**Figure 1b**). We first presented participants with simple zero-order sequences (level-0) followed by

more complex first-order sequences (level-1), as previous work has shown that temporal dependencies are more difficult to learn as their length increases (van den Bos and Poletiek, 2008) and training with simple dependencies may facilitate learning of more complex contingencies (Antoniou et al., 2016). Zero-order sequences (level-0) were context-less; that is, the presentation of each symbol depended only on the probability of occurrence of each symbol. For first-order sequences (level-1), the presentation of a particular symbol was conditionally dependent on the previously presented symbol (i.e. context length of one).

As the sequences we employed were probabilistic, we developed a probabilistic measure to assess participants' performance in the prediction task. Specifically, we computed a performance index (PI) that indicates how closely the probability distribution of the participant responses matched the probability distribution of the presented symbols. This is preferable to a simple measure of accuracy because the probabilistic nature of the sequences means that the 'correct' upcoming symbol is not uniquely specified; thus, designating a particular choice as correct or incorrect is often arbitrary.

Comparing normalized performance (i.e. after subtracting performance based on random guessing) before and after training per level (**Figure 2a**) showed that participants improved substantially in learning probabilistic structures (i.e. mean improvement higher than 20% for all levels). A two-way repeated measures ANOVA with Session (Pre, Post) and Level (level-0, level-1) showed a significant main effect of Session (F(1,18)=58.7, p<0.001), but no significant effect of Level (F(1,18)=0.6, p=0.459) nor a significant interaction (F(1,18)=0.6, p=0.459), indicating that participants improved similarly at both levels through training. Further, we asked whether these learning effects were specific to the trained structured sequences. We contrasted performance on structured vs. random sequences before and after training sessions. A repeated-measures ANOVA showed a significant interaction of Session (Pre, Post) and Sequence type (structured, random) for level-0 (F(1,18)=20.5,

p<0.001) and level-1 (F(1,18)=58.6, p<0.001), suggesting that learning improvement was specific to the structured sequences.

3.2 Response strategies: matching vs. maximization

Previous work (Shanks et al., 2002; Lagnado et al., 2006; Rieskamp and Otto, 2006; Eckstein et al., 2013; Acerbi et al., 2014; Fulvio et al., 2014; Murray et al., 2015) on probabilistic learning and decision making has proposed that individuals use two possible response strategies when making a choice: matching vs. maximization. Observers have been shown to either match their choices stochastically according to the underlying input statistics or to maximize their reward by selecting the most probable positively rewarded outcomes. In the context of our task, as the Markov models that generated stimulus sequences were stochastic, participants needed to learn the probabilities of different outcomes to succeed in the prediction task. It is possible that participants used probability maximization whereby they always select the most probable outcome in a particular context. Alternatively, participants might learn the relative probabilities of each symbol (e.g. p(A)=0.18; p(B)=0.72, p(C)=0.05; p(D)=0.05) and respond so as to reproduce this distribution, a strategy referred to as probability matching.

To quantify participants' strategies across training, we computed a strategy index that indicates each participant's preference (on a continuous scale) for responding using probability matching vs. maximization (**Figure 2b**). **Figures 2b**, **2c** indicate variability in strategy index across participants. Comparing individual strategy across levels showed significantly higher values for level-1 compared to level-0 (t(18)=2.2, p=0.04), suggesting that participants adopted a strategy closer to maximization when learning context-based rather than frequency statistics (**Figure 2c**). Note, that this relationship was not confounded by differences in performance, as there were no significant correlations (level-0: r=0.34, p=0.19; level-1: r=0.04, p=0.88) between performance after training and strategy index.

Further, we conducted two additional analyses to control for the possibility that the differences we observed in strategy index between levels may be confounded by differences in the probability distributions between levels (i.e. 72% vs. 80% probability for the most frequent target for level-0 vs. level-1) and PI. First, we observed significantly higher strategy index for level-1 compared to level-0 (t(18)=2.19, p=0.042) after scaling the strategy index in level-0 by 0.8/0.72 (i.e. the ratio of maximum PI for exact maximization for level-1 vs. level-0). Second, strategy index remained higher for level-1 than level-0 (t(18)=2.36, p=0.030) after regressing out the post-training PI from strategy index per level. Thus, our result showing higher strategy index for level-1 than level-0 is unlikely to be confounded by differences in PI or the probability distributions between levels.

Finally, participants were exposed to the sequences without trial-by-trial feedback, but were given block feedback about their performance that motivated them to continue with training. A control experiment during which the participants were not given any feedback showed similar results to our main experiment; that is, higher strategy index for level-1 than level-0, suggesting that differences in the strategy between levels could not be simply attributed to feedback. Taken together, these results suggest that participants adopt a strategy closer to maximization for learning higher–order sequences (i.e. context-based statistics) than simple frequency statistics. This is consistent with previous studies showing that participants adopt a strategy closer to matching when learning a simple probabilistic task in the absence of trial-by-trial feedback (Shanks et al., 2002). However, for more complex probabilistic tasks, participants weight their responses towards the most likely outcome (i.e. adopt a strategy closer to maximization) after training (Lagnado et al., 2006).

Figure 2

3.3 fMRI analysis: functional brain networks

To identify functional brain networks that mediate our ability to adapt to changes in temporal statistics, we performed fMRI on participants before and after training on each level with structured and random sequences. First, we decomposed the fMRI timecourse into functionally connected components (i.e. components comprising voxel clusters with correlated fMRI time course) using ICA and selected components of neuronal origin using a spatial correlation method with known brain networks (Allen et al., 2011) (**Figure 3, Table 1**). We then tested whether learning-dependent changes in fMRI activation in these brain networks relate to individual strategy when learning frequency and context-based statistics. For each component we extracted a β weight across voxels for structured and random sequences per session (pre-, post-training). We correlated learning-dependent changes in fMRI signal (post-pre β weight) for structured sequences with individual strategy. Positive correlations indicate increased activation after training that relates to maximization, while negative correlations indicate increased activation that relates to matching, as negative strategy index indicates strategy towards matching.

Figure 3, Table 1

First, we observed significant negative correlations between learning-dependent fMRI changes and strategy index in functional brain networks known to be involved in memory processes and stimulus-response associations. In particular, for learning frequency statistics (**Figure 4a**), we found significant negative correlations of fMRI activation change in the Precuneus (CP_20, peak activations in bilateral precuneus and cingulate; r=-0.70, CI=[-0.88, -0.48]), the Sensorimotor (CP_6, peak activations in bilateral precentral and postcentral gyri; r=-0.70, CI=[-0.90, -0.42]) and the Right Central Executive (CP_17, peak activations in right inferior parietal and right inferior frontal gyrus; r=-0.42, CI=[-0.73, -0.07]) networks with strategy. For learning context-based statistics (**Figure 4b**), we found significant negative correlations of fMRI activation change in the Precuneus (CP_20; r=-0.37, CI=[-0.68, -0.03])

and the Middle Temporal (CP_26, peak activations in bilateral precuneus and middle temporal gyrus extending medially into parahippocampal cortex; r=-0.44, CI=[-0.74, -0.01]) networks with strategy. These results suggest that increased functional activation in these brain networks after training relates to matching the exact sequence statistics. This is consistent with the role of precuneus and cingulate in memory retrieval (Wagner et al., 2005; Cabeza et al., 2008; St. Jacques et al., 2011) in the context of episodic and working memory tasks (Nyberg et al., 2002). Further, sensorimotor areas have been implicated in the consolidation of stimulus-response associations, mainly at early stages of motor consolidation (Muellbacher et al., 2002). Similarly, the Right Central Executive Network has been implicated in the initial stages of learning (Seger et al., 2000). Thus, these networks contribute at the initial raining on frequency statistics, as this brain network has been implicated in episodic memory and mnemonic tasks involving longer memory length (Nyberg et al., 2002; Vincent et al., 2006; Cabeza et al., 2008).

Figure 4

In contrast, we observed significant positive correlations between learning-dependent fMRI changes and strategy in basal ganglia and the Left Central Executive Network. In particular, for learning frequency statistics, we found a significant positive correlation of fMRI activation change in the Basal Ganglia Network (CP_13, peak activation in bilateral caudate) with strategy (r=0.43, CI=[0.04, 0.72]) (**Figure 5a**), suggesting involvement of basal ganglia in learning by maximizing. This is consistent with previous work suggesting that basal ganglia is involved in the consolidation of the stimulus-response mapping (Albouy et al., 2008; Shohamy et al., 2008) and category learning (Ashby and Maddox, 2005; Seger and Cincotta, 2005). In particular, previous work on humans and animals emphasizes the role of the caudate in switching between strategies (Monchi et al., 2001; Cools et al., 2004; Seger

and Cincotta, 2005), and learning after a rule reversal (Cools et al., 2002; Pasupathy and Miller, 2005). For learning context-based statistics, we found a significant positive correlation of fMRI activation change in the Left Central Executive Network (CP_21, peak activations in left inferior parietal and left middle frontal gyrus) with strategy (r=0.63, CI=[0.29, 0.84]) (**Figure 5b**), suggesting that higher activation after training in this region relates to maximization. Executive networks have been implicated in holding and updating task rules (Ridderinkhof et al., 2004; Vincent et al., 2008; D'Ardenne et al., 2012). In particular, increased activation in the Left Central Executive Network has been shown after training in the context of category learning (Seger et al., 2000). This is consistent with our behavioral results showing that participants adopt a stronger maximization strategy during later training on context-based statistics.

Figure 5

Finally, we tested whether our results were specific to the learned structured sequences. We computed fMRI activation for random sequences in brain networks that showed significant correlations with strategy for structured sequences. For frequency statistics, fMRI activation change in the Precuneus Network showed a significant negative correlation with strategy (r=-0.53, CI=[-0.81, -0.11]). For context-based statistics: a) activation change in the Middle Temporal Network (r=-0.59, CI=[-0.87, -0.14]) and the Precuneus Network (r=-0.57, CI=[-0.81, -0.25]) showed a significant negative correlation with strategy b) activation change in the Left Central Executive Network showed a significant positive correlation with strategy (r=0.61, CI=[0.25, 0.79]). To compare correlations for structured vs. random sequences, we used a Steiger z-score comparison (Lee and Preacher, 2013), for comparing of dependent correlations with a shared variable (i.e. strategy index). We found significantly higher negative correlations for structured vs. random trials in: a) Precuneus (z=-2.19, p=0.029), b) Right Central Executive (z=-2.43, p=0.015) and

c) Sensorimotor (z=-2.92, p=0.004) Networks. These results suggest differences in the processing of structured vs. random sequences primarily when participants learn by matching, as this strategy requires learning the exact sequence statistics that differ between these two sequence types.

3.4 Functional Network Connectivity (FNC)

Our analyses so far identified brain networks that show learning-dependent changes in functional processing that relate to individual strategy for learning temporal structure. Next, we asked whether learning-dependent changes in the connectivity between these networks relate to individual strategy when learning frequency and context-based statistics. We calculated pairwise correlations between the six brain networks (Precuneus, Sensorimotor, Right Central Executive, Middle Temporal, Basal Ganglia, Left Central Executive) that showed significant correlations with strategy (see 3.3). We calculated these correlations for each session (Pre, Post-0, Post-1) and converted them to z-scores (Fisher z). We then correlated change (post- minus pre-training z-score) in Functional Network Connectivity (FNC) with strategy index to assess the relationship of strategy with changes in between network connectivity (**Figure 6**).

For frequency statistics, we found that a) connectivity change between Left Central Executive and Middle Temporal network correlated negatively with strategy (r=-0.62, CI=[-0.86, -0.18]), and b) connectivity change between Precuneus and Sensorimotor networks correlated negatively with strategy (r=-0.62, CI=[-0.88, -0.15]). These results suggest that increased connectivity between these networks with training relates to learning by matching the exact sequence statistics. For context-based statistics, we found that connectivity change between Right Central Executive and Basal Ganglia networks correlated positively with strategy (r=0.55, CI=[0.01, 0.85]), suggesting that increased connectivity between these

networks with training relates to maximization. These results are consistent with previous work highlighting the role of Central Executive networks in controlling learning of contextual and stimulus-response associations (Ridderinkhof et al., 2004; D'Ardenne et al., 2012). Further, recent neurophysiology findings (Antzoulatos and Miller, 2014) show enhanced connectivity between prefrontal cortex and basal ganglia in the context of category learning, suggesting that fast learning in the basal ganglia may train slower learning in the frontal cortex that may facilitate the generalization and abstraction of learned associations.

This functional connectivity analysis is consistent with our previous analyses showing fronto-striatal networks involved in maximization, the strategy for which participants showed stronger preference when learning context-based statistics (Figure 2b, 2c). Our results provide complementary evidence that learning-dependent changes in the connectivity of brain networks known to be involved in memory and stimulus-response associations mediate learning by matching the exact sequence statistics, while connectivity changes in frontal and striatal networks mediate learning by maximizing (i.e. extracting the most probable outcome in a given context).

Figure 6

4. Discussion

Here, we investigate the functional brain networks that mediate our ability to adapt to changes in the environment's statistics and make predictions. Our behavioral results demonstrate that individuals adapt to changes in temporal structure and extract the relevant frequency or context-based statistics for making predictions of upcoming events. Our fMRI results provide evidence for dissociated functional brain networks that mediate our ability to extract behaviorally-relevant statistics.

Our modeling approach allows us to track participants' predictions and their strategies during training. We demonstrate that learning predictive structures relates to individual

variability in response strategies: that is individuals favored either probability maximization (i.e. extracting the most probable outcome in a given context) or matching the exact sequence statistics. Previous behavioral studies have reported individual variability in response strategy in the context of probabilistic learning tasks and suggested that strategies change during the course of training with feedback (Gluck et al., 2002; Shanks et al., 2002; Lagnado et al., 2006). Here we show that response strategy relates to sequence structure; that is, learning context-based statistics relates to stronger maximization than learning simple frequency statistics. Further, we provide evidence that these response strategies engage distinct functional brain networks: matching relates to changes in fMRI activation within and functional connectivity between brain networks involved in memory and stimulus-response associations, while maximizing relates to changes in frontal and striatal brain networks.

Previous studies have implicated these brain networks in reinforcement learning (e.g. for reviews (Robbins, 2007; Balleine and O'Doherty, 2010)). Previous brain imaging and neurophysiology studies have demonstrated learning-dependent changes in functional brain connectivity in a range of tasks: visual perceptual learning (Lewis et al., 2009; Baldassarre et al., 2012), category learning (Antzoulatos and Miller, 2014), motor learning (Sun et al., 2007; Bassett et al., 2011; Ma et al., 2011), auditory learning (Ventura-Campos et al., 2013) and language learning (Veroude et al., 2010). However, most of this work has focused on reward-based learning that involves training with trial-by-trial feedback. Here, we show that learning temporal statistics may proceed without explicit trial-by-trial feedback and involve interactions between brain networks similar to those known to support reward-based learning (Alexander et al., 1986; Lawrence et al., 1998).

Finally, we considered whether the learning we observed occurred in an incidental manner or involved explicit knowledge of the underlying sequence structure. Previous studies have suggested that learning of regularities may occur implicitly in a range of tasks:

visuomotor sequence learning (Nissen and Bullemer, 1987; Seger, 1994; Schwarb and Schumacher, 2012), artificial grammar learning (Reber, 1967), probabilistic category learning (Knowlton et al., 1994) and contextual cue learning (Chun and Jiang, 1998). This work has focused on implicit measures of sequence learning, such as familiarity judgments or reaction times. In contrast, our paradigm allows us to directly test whether exposure to temporal sequences facilitates the observers' ability to explicitly predict the identity of the next stimulus in a sequence. Although, our experimental design makes it unlikely that the participants memorized specific stimulus positions or the full sequences, debriefing the participants suggests that most extracted some high probability symbols or context-target combinations. Thus, it is possible that prolonged exposure to probabilistic structures (i.e. multiple sessions in contrast to single exposure sessions typically used in statistical learning studies) in combination with prediction judgments (Dale et al., 2012) may evoke some explicit knowledge of temporal structures, in contrast to implicit measures of anticipation typically used in statistical learning studies.

5. Conclusions

Our findings provide evidence that functional brain connectivity changes with learning in dissociable networks to support our ability to extract behaviorally-relevant statistics. This network connectivity relates to individual response strategies when learning temporal structures. Our paradigm tested learning of structures that increased in context-length over time; thus, it does not allow us to dissociate learning time course from changes in sequence structure over time. In future work, it would be interesting to investigate the time course of learning temporal statistics using dynamic connectivity analysis that allows us to track changes in brain connectivity over time.

Acknowledgements: We would like to thank Caroline di Bernardi Luft for help with data collection; Matthew Dexter for help with software development; David Samu, Loraine Tyler, Deniz Vatansever and Emmanuel Stamatakis for helpful discussions and feedback.

Funding: This work was supported by grants to ZK from the Biotechnology and Biological Sciences Research Council [H012508], the Leverhulme Trust [RF-2011-378] and the [European Community's] Seventh Framework Programme [FP7/2007-2013] under agreement PITN-GA-2011-290011, AEW from the Wellcome Trust (095183/Z/10/Z) and the [European Community's] Seventh Framework Programme [FP7/2007-2013] under agreement PITN-GA-2012-316746, PT from Engineering and Physical Sciences Research Council [EP/L000296/1].

CEP CEP

References

- Acerbi L, Vijayakumar S, Wolpert DM (2014) On the Origins of Suboptimality in Human Probabilistic Inference. PLoS Computational Biology 10:e1003661.
- Aizenstein HJ, Stenger AV, Cochran J, Clark K, Johnson M, Nebes RD, Carter CS (2004) Regional Brain Activation during Concurrent Implicit and Explicit Sequence Learning. Cerebral Cortex 14:199-208.
- Albouy G, Sterpenich V, Balteau E, Vandewalle G, Desseilles M, Dang-Vu T, Darsaud A, Ruby P, Luppi PH, Degueldre C, Peigneux P, Luxen A, Maquet P (2008) Both the hippocampus and striatum are involved in consolidation of motor sequence memory. Neuron 58:261-272.
- Alexander GE, DeLong MR, Strick PL (1986) Parallel Organization of Functionally Segregated Circuits Linking Basal Ganglia and Cortex. Annual Review of Neuroscience 9:357-381.
- Allen EA et al. (2011) A baseline for the multivariate comparison of resting-state networks. Frontiers in systems neuroscience 5:2.
- Antoniou M, Ettlinger M, Wong PC (2016) Complexity, Training Paradigm Design, and the Contribution of Memory Subsystems to Grammar Learning. PloS one 11:e0158812.
- Antzoulatos EG, Miller EK (2014) Increases in functional connectivity between prefrontal cortex and striatum during category learning. Neuron 83:216-225.
- Ashby FG, Maddox WT (2005) Human Category Learning. Annual Review of Psychology 56:149-178.
- Aslin RN, Newport EL (2012) Statistical learning: From acquiring specific items to forming general rules. Current directions in psychological science 21:170-176.
- Baldassarre A, Lewis CM, Committeri G, Snyder AZ, Romani GL, Corbetta M (2012) Individual variability in functional connectivity predicts performance of a perceptual task. Proceedings of the National Academy of Sciences 109:3516-3521.
- Balleine BW, O'Doherty JP (2010) Human and Rodent Homologies in Action Control: Corticostriatal Determinants of Goal-Directed and Habitual Action. Neuropsychopharmacology 35:48-69.
- Bassett DS, Wymbs NF, Porter MA, Mucha PJ, Carlson JM, Grafton ST (2011) Dynamic reconfiguration of human brain networks during learning. Proceedings of the National Academy of Sciences 108:7641-7646.
- Brainard DH (1997) The psychophysics toolbox. Spatial vision 10:433-436.
- Cabeza R, Ciaramelli E, Olson IR, Moscovitch M (2008) The parietal cortex and episodic memory: an attentional account. Nature reviews Neuroscience 9:613-625.
- Calhoun VD, Adali T (2012) Multisubject independent component analysis of fMRI: A decade of intrinsic networks, default mode, and neurodiagnostic discovery. IEEE Reviews in Biomedical Engineering 5:60-73.
- Calhoun VD, Liu J, Adali T (2009) A review of group ICA for fMRI data and ICA for joint inference of imaging, genetic, and ERP data. NeuroImage 45:163-172.
- Chun M, Jiang Y (1998) Contextual cueing: implicit learning and memory of visual context guides spatial attention. Cognitive psychology 36:28-71.
- Chun MM (2000) Contextual cueing of visual attention. Trends in cognitive sciences 4:170-178.
- Cools R, Clark L, Robbins TW (2004) Differential Responses in Human Striatum and Prefrontal Cortex to Changes in Object and Rule Relevance. Journal of Neuroscience 24:1129-1135.
- Cools R, Clark L, Owen AM, Robbins TW (2002) Defining the neural mechanisms of probabilistic reversal learning using event-related functional magnetic resonance imaging. The Journal of neuroscience : the official journal of the Society for Neuroscience 22:4563-4567.
- D'Ardenne K, Eshel N, Luka J, Lenartowicz A, Nystrom LE, Cohen JD (2012) Role of prefrontal cortex and the midbrain dopamine system in working memory updating. Proceedings of the National Academy of Sciences 109:19900-19909.
- Dale R, Duran ND, Morehead JR (2012) Prediction during statistical learning, and implications for the implicit/explicit divide. Advances in Cognitive Psychology 8:196-209.

- Eckstein MP, Mack SC, Liston DB, Bogush L, Menzel R, Krauzlis RJ (2013) Rethinking human visual attention: Spatial cueing effects and optimality of decisions by honeybees, monkeys and humans. Vision Research 85:5-9.
- Erev I, Barron G (2005) On Adaptation, Maximization, and Reinforcement Learning Among Cognitive Strategies. Psychological review 112:912-931.
- Fiser J, Aslin RN (2002) Statistical learning of higher-order temporal structure from visual shape sequences. Journal of Experimental Psychology: Learning, Memory, and Cognition 28:458-467.
- Fiser J, Aslin RN (2005) Encoding multielement scenes: statistical learning of visual feature hierarchies. Journal of Experimental Psychology: General 134:521.
- Fox MD, Raichle ME (2007) Spontaneous fluctuations in brain activity observed with functional magnetic resonance imaging. Nature reviews Neuroscience 8:700-711.
- Fulvio JM, Green CS, Schrater PR (2014) Task-Specific Response Strategy Selection on the Basis of Recent Training Experience. PLoS Computational Biology 10:e1003425.
- Gheysen F, Van Opstal F, Roggeman C, Van Waelvelde H, Fias W (2011) The Neural Basis of Implicit Perceptual Sequence Learning. Frontiers in Human Neuroscience 5.
- Gluck MA, Shohamy D, Myers C (2002) How do people solve the "weather prediction" task?: individual variability in strategies for probabilistic category learning. Learning & memory 9:408-418.
- Haberecht MF, Menon V, Warsofsky IS, White CD, Dyer-Friedman J, Glover GH, Neely EK, Reiss AL (2001) Functional neuroanatomy of visuo-spatial working memory in turner syndrome. Human Brain Mapping 14:96-107.
- Himberg J, Hyvärinen A, Esposito F (2004) Validating the independent components of neuroimaging time series via clustering and visualization. NeuroImage 22:1214-1222.
- Hsieh LT, Gruber MJ, Jenkins LJ, Ranganath C (2014) Hippocampal Activity Patterns Carry Information about Objects in Temporal Context. Neuron 81:1165-1178.
- Jafri MJ, Pearlson GD, Stevens M, Calhoun VD (2008) A method for functional network connectivity among spatially independent resting-state components in schizophrenia. NeuroImage 39:1666-1681.
- Knowlton BJ, Squire LR, Gluck MA (1994) Probabilistic classification learning in amnesia. Learning & memory 1:106-120.
- Lagnado DA, Newell BR, Kahan S, Shanks DR (2006) Insight and strategy in multiple-cue learning. Journal of Experimental Psychology: General 135:162.
- Lawrence AD, Sahakian BJ, Robbins TW (1998) Cognitive functions and corticostriatal circuits: insights from Huntington's disease. Trends in Cognitive Sciences 2:379-388.
- Lee IA, Preacher KJ (2013) Calculation for the test of the difference between two dependent correlations with one variable in common.
- Lewis CM, Baldassarre A, Committeri G, Romani GL, Corbetta M (2009) Learning sculpts the spontaneous activity of the resting human brain. Proceedings of the National Academy of Sciences 106:17558-17563.
- Ma L, Narayana S, Robin DA, Fox PT, Xiong J (2011) Changes occur in resting state network of motor system during 4weeks of motor skill learning. NeuroImage 58:226-233.
- McKeown MJ, Makeig S, Brown GG, Jung TP, Kindermann SS, Bell AJ, Sejnowski TJ (1998) Analysis of fMRI data by blind separation into independent spatial components. Human Brain Mapping 6:160-188.
- Monchi O, Petrides M, Petre V, Worsley K, Dagher A (2001) Wisconsin Card Sorting revisited: distinct neural circuits participating in different stages of the task identified by event-related functional magnetic resonance imaging. The Journal of neuroscience : the official journal of the Society for Neuroscience 21:7733-7741.
- Muellbacher W, Ziemann U, Wissel J, Dang N, Kofler M, Facchini S, Boroojerdi B, Poewe W, Hallett M (2002) Early consolidation in human primary motor cortex. Nature 415:640-644.

- Murray RF, Patel K, Yee A (2015) Posterior Probability Matching and Human Perceptual Decision Making. PLOS Computational Biology 11:e1004342.
- Nissen MJ, Bullemer P (1987) Attentional requirements of learning: Evidence from performance measures. Cognitive Psychology 19:1-32.
- Nyberg L, Forkstam C, Petersson KM, Cabeza R, Ingvar M (2002) Brain imaging of human memory systems: between-systems similarities and within-system differences. Cognitive Brain Research 13:281-292.
- Pasupathy A, Miller EK (2005) Different time courses of learning-related activity in the prefrontal cortex and striatum. Nature 433:873-876.
- Pelli DG (1997) The VideoToolbox software for visual psychophysics: transforming numbers into movies. Spatial Vision 10:437-442.
- Pernet CR, Wilcox R, Rousselet GA (2013) Robust Correlation Analyses: False Positive and Power Validation Using a New Open Source Matlab Toolbox. Frontiers in psychology 3.
- Perruchet P, Pacton S (2006) Implicit learning and statistical learning: one phenomenon, two approaches. Trends in cognitive sciences 10:233-238.
- Rauch SL, Whalen PJ, Savage CR, Curran T, Kendrick A, Brown HD, Bush G, Breiter HC, Rosen BR (1997) Striatal recruitment during an implicit sequence learning task as measured by functional magnetic resonance imaging. Human Brain Mapping 5:124-132.
- Reber AS (1967) Implicit learning of artificial grammars. Journal of Verbal Learning and Verbal Behavior 6:855-863.
- Ridderinkhof KR, van den Wildenberg WP, Segalowitz SJ, Carter CS (2004) Neurocognitive mechanisms of cognitive control: the role of prefrontal cortex in action selection, response inhibition, performance monitoring, and reward-based learning. Brain and cognition 56:129-140.
- Rieskamp J, Otto PE (2006) SSL: A Theory of How People Learn to Select Strategies. Journal of Experimental Psychology: General 135:207-236.
- Rissanen J (1978) Modeling by shortest data description. Automatica 14:465-471.
- Robbins TW (2007) Shifting and stopping: fronto-striatal substrates, neurochemical modulation and clinical implications. Philosophical Transactions of the Royal Society B: Biological Sciences 362:917-932.
- Rose M, Haider H, Salari N, Buchel C (2011) Functional Dissociation of Hippocampal Mechanism during Implicit Learning Based on the Domain of Associations. Journal of Neuroscience 31:13739-13745.
- Saffran JR, Aslin RN, Newport EL (1996) Statistical Learning by 8-Month-Old Infants. Science 274:1926-1928.
- Saffran JR, Johnson EK, Aslin RN, Newport EL (1999) Statistical learning of tone sequences by human infants and adults. Cognition 70:27-52.
- Schapiro AC, Kustner LV, Turk-Browne NB (2012) Shaping of object representations in the human medial temporal lobe based on temporal regularities. Current Biology 22:1622-1627.
- Schendan HE, Searl MM, Melrose RJ, Stern CE (2003) An fMRI Study of the Role of the Medial Temporal Lobe in Implicit and Explicit Sequence Learning. Neuron 37:1013-1025.
- Schwarb H, Schumacher E (2012) Generalized lessons about sequence learning from the study of the serial reaction time task. Advances in Cognitive Psychology 8:165-178.
- Seger CA (1994) Implicit learning. Psychological Bulletin 115:163-196.
- Seger CA, Cincotta CM (2005) Dynamics of Frontal, Striatal, and Hippocampal Systems during Rule Learning. Cerebral Cortex 16:1546-1555.
- Seger CA, Poldrack RA, Prabhakaran V, Zhao M, Glover GH, Gabrieli JD (2000) Hemispheric asymmetries and individual differences in visual concept learning as measured by functional MRI. Neuropsychologia 38:1316-1324.
- Shanks DR, Tunney RJ, McCarthy JD (2002) A re-examination of probability matching and rational choice. Journal of Behavioral Decision Making 15:233-250.

- Shohamy D, Myers CE, Kalanithi J, Gluck MA (2008) Basal ganglia and dopamine contributions to probabilistic category learning. Neuroscience and biobehavioral reviews 32:219-236.
- Smith SM, Fox PT, Miller KL, Glahn DC, Fox PM, Mackay CE, Filippini N, Watkins KE, Toro R, Laird AR, Beckmann CF (2009) Correspondence of the brain's functional architecture during activation and rest. Proceedings of the National Academy of Sciences of the United States of America 106:13040-13045.
- St. Jacques PL, Kragel PA, Rubin DC (2011) Dynamic neural networks supporting memory retrieval. NeuroImage 57:608-616.
- Stevens MC, Kiehl KA, Pearlson G, Calhoun VD (2007) Functional neural circuits for mental timekeeping. Human Brain Mapping 28:394-408.
- Sun FT, Miller LM, Rao AA, D'esposito M (2007) Functional connectivity of cortical networks involved in bimanual motor sequence learning. Cerebral cortex 17:1227-1234.
- Taylor JR, Williams N, Cusack R, Auer T, Shafto MA, Dixon M, Tyler LK, Cam-CAN X, Henson RN (2015) The Cambridge Centre for Ageing and Neuroscience (Cam-CAN) data repository: Structural and functional MRI, MEG, and cognitive data from a cross-sectional adult lifespan sample. NeuroImage 144:262-269.
- Turk-Browne NB, Junge J, Scholl BJ (2005) The automaticity of visual statistical learning. Journal of experimental psychology General 134:552-564.
- Turk-Browne NB, Scholl BJ, Johnson MK, Chun MM (2010) Implicit perceptual anticipation triggered by statistical learning. The Journal of neuroscience : the official journal of the Society for Neuroscience 30:11177-11187.
- van den Bos E, Poletiek FH (2008) Effects of grammar complexity on artificial grammar learning. Memory & Cognition 36:1122-1131.
- Van Dijk KRa, Hedden T, Venkataraman A, Evans KC, Lazar SW, Buckner RL (2010) Intrinsic Functional Connectivity As a Tool For Human Connectomics: Theory, Properties, and Optimization. Journal of neurophysiology 103:297-321.
- Ventura-Campos N, Sanjuan A, Gonzalez J, Palomar-Garcia M-A, Rodriguez-Pujadas A, Sebastian-Galles N, Deco G, Avila C (2013) Spontaneous Brain Activity Predicts Learning Ability of Foreign Sounds. Journal of Neuroscience 33:9295-9305.
- Veroude K, Norris DG, Shumskaya E, Gullberg M, Indefrey P (2010) Functional connectivity between brain regions involved in learning words of a new language. Brain and language 113:21-27.
- Vincent JL, Kahn I, Snyder AZ, Raichle ME, Buckner RL (2008) Evidence for a frontoparietal control system revealed by intrinsic functional connectivity. Journal of neurophysiology 100:3328-3342.
- Vincent JL, Snyder AZ, Fox MD, Shannon BJ, Andrews JR, Raichle ME, Buckner RL (2006) Coherent spontaneous activity identifies a hippocampal-parietal memory network. Journal of neurophysiology 96:3517-3531.
- Wagner AD, Shannon BJ, Kahn I, Buckner RL (2005) Parietal lobe contributions to episodic memory retrieval. Trends in Cognitive Sciences 9:445-453.
- Wozny DR, Beierholm UR, Shams L (2010) Probability Matching as a Computational Strategy Used in Perception. PLoS Computational Biology 6:e1000871.

Figure 1: Trial and sequence design. (a) The trial design: 8-14 symbols were presented sequentially followed by a cue and the test display. (b) Sequence design: Markov models of the two context-length levels. For the zero-order model (level-0): different states (A, B, C, D) are assigned to four symbols with different probabilities. For the first-order model (level-1), diagrams indicate states (circles) and conditional probabilities (solid arrow: high probability; dashed arrow: low probability). Transitional probabilities are shown in a four-by-four (level-1) conditional probability matrix, where rows indicate temporal context and columns the corresponding target.

Figure 2: Behavioral performance. (a) Mean normalized performance index (PI) across participants per level during pre-training (grey bars) and post-training (black bars) test sessions. Error bars indicate standard error of the mean. (b) Strategy index boxplots for level-0 and level-1 indicate individual variability. The upper and lower error bars display the minimum and maximum data values and the central boxes represent the interquartile range (25th to 75th percentiles). The thick line in the central boxes represents the median. (c) Scatterplot of strategy index for level-0 against strategy index for level-1.

Figure 3: Spatial maps of ICA task-related components. 15 task-related components are shown organized into known functional groups (Allen et al., 2011). Spatial maps are thresholded at p<0.005 (FWER corrected) and displayed in neurological convention (left is left) on the MNI template. The x,y,z coordinates per component denote the location of the sagittal, coronal and axial slices, respectively.

Figure 4: ICA components related to matching strategy. Average spatial maps showing significant negative correlation of BOLD change (post- minus pre-training) with strategy

index for (a) learning frequency statistics: Precuneus, Sensorimotor and Right Central Executive. (b) learning context-based statistics: Precuneus and Middle Temporal. Spatial maps are averaged across sessions, thresholded at p<0.005 (FWER corrected) and displayed in neurological convention (left is left) on the MNI template. Open circles in the correlation plots denote outliers.

Figure 5: ICA components related to maximization strategy. Average spatial maps showing significant positive correlation of BOLD change (post- minus pre-training) with strategy index for: (a) learning frequency statistics: Basal Ganglia. (b) learning context-based statistics: Left Central Executive. Spatial maps are averaged across sessions, thresholded at p<0.005 (FWER corrected) and displayed in neurological convention (left is left) on the MNI template.

Figure 6: Functional Network Connectivity (FNC) change related to strategy. Average correlation matrix of FNC change (post- minus pre-training) with strategy index for: (a) frequency statistics and (b) context-based statistics. Black dots indicate significant positive, while black diamonds significant negative correlations (at 95% bootstrapped confidence intervals) of FNC change with strategy index. ICA components included in this analysis are: left central executive network (ICEN), right central executive network (rCEN), middle temporal (MT), precuneus (PRCUN), basal ganglia (BG) and sensorimotor (SM).

Tables

Table 1: ICA components. Clusters within the 15 task-related components are extracted from the group maps and are organized into known functional groups (Allen et al., 2011). The table shows the number of voxels within each cluster, the x,y,z coordinates of the peak voxel in MNI space and the t-statistic of the peak voxel.

Network	Component	Areas	voxels	x,y,z (mm)	t-value
Attentional	CP 17	Inferior Parietal R	387	48,-61,42	23.91
		Cerebellum Posterior L	151	-12,-73,-34	18.51
		Inferior Frontal Gyrus R	817	45,41,14	16.35
		Thalamus R	67	9,-22,6	15.95
		Putamen R	155	30,14,-6	14.96
		Inferior Parietal L	57	-48,-46,50	11.75
	CP 21	Inferior Parietal L	414	-36,-58,42	16.32
		Cerebellum Posterior R	147	21,-67,-34	15.29
		Middle Frontal Gyrus	658	-45,23,30	15.04
		Putamen L	46	-33,-16,-6	14.08
		Insula L	25	-27,17,-2	11.76
	CP 24	Cingulate Gyrus BL	3742	6,20,38	28.71
		Cerebellum Posterior R	23	36,-67,-26	10.32
	CP 13	Caudate BL	1548	18,17,2	28.46
	CP 27	Putamen BL	1321	-24,2,-6	25.81
Dasal Galiglia		Cingulate Gyrus	86	6,-1,46	12.07
		Cerebellum Anterior	20	-3,-58,-38	11.17
Default Mode	CP 20	Precuneus BL	819	12,-67,30	21.61
		Cingulate BL	251	12,32,18	15.51
		Superior Frontal Gyrus L	26	-24,41,22	11.86
		Inferior Parietal R	20	48,-55,42	10.72
	CP 23	Anterior Cingulate BL	1834	-6,44,10	24.39
		Posterior Cingulate BL	121	-3,-46,30	18.83
		Cingulate Gyrus	32	-3,-16,38	11.45
		Superior Temporal Gyrus L	68	-48,-58,26	12.18
		Cerebellum Posterior R	52	27,-79,-30	11.02
		Superior Temporal Gyrus R	26	54,-61,38	10.61
		Putamen R	24	30,8,2	10.48
	CP 26	Precuneus BL	1011	6,-61,18	21.88
		Middle Temporal Gyrus R	237	45,-64,22	21.75
		Middle Temporal Gyrus L	233	-48,-67,14	13.61
		Postcentral Gyrus R	93	48,8,26	14.71
Sensorimotor	CP 5	Superior Temporal Gyrus L	516	-45,-19,6	19.38
		Superior Temporal Gyrus R	654	48,-10,6	17.27
		Medial Frontal Gyrus	24	-3,-1,62	12.41

			1 1		
		Postcentral Gyrus I	448	-42 -28 54	25.15
		Precentral Gyrus R	91	57 -13 34	14 48
	CP 6	Cerebellum Anterior R	72	215226	12.48
		Postcentral Gyrus R	30	36,-7,62	10.96
		Parietal Superior L	20	-21,-61,58	10.03
	CP 10	Paracentral BL	1653	21,-31,66	24.7
		Cerebellum Anterior	125	-6,-46,-18	12.75
		Insula L	139	-39,-13,-2	17.42
		Supramarginal R	167	57,-28,26	17.17
		Insula R	177	42,-10,-6	16.19
	CD 10	Supramarginal L	114	-63,-31,22	15.48
	CP 19	Cingulate Gyrus BL	87	12,-34,38	13.3
		Precuneus BL	33	-6,-49,58	13.21
		Postcentral R	20	21,-46,66	11.2
		Middle Temporal Gyrus L	22	-54,-61,6	9.83
Visual	CP 7	Lingual Gyrus BL	1197	5,-63,2	31.77
		Cerebellum Declive	47	-3,-73,-26	15.43
	CP 12	Middle Occipital Gyrus BL	1730	30,-85,18	22.85
		Posterior Cingulate BL	107	1,-31,26	15.94
Cerebellum	CP 16	Cerebellum Anterior Lobe	3013	30,-58,-34	36.86
		Precuneus BL	30	3,-58,38	10.51









a. Frequency statistics









Highlights

- Learning temporal structure without feedback alters functional brain networks
- Functional brain connectivity relates to individual strategies for learning
- Dissociable functional brain networks mediate learning of predictive statistics