Recruitment of the default mode network during a demanding act of executive control

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Declaration of Conflicting Interests
We declare no conflicting interests
Abstract

In the human brain, a default mode or task-negative network shows reduced activity during many cognitive tasks, and is often associated with internally-directed processes such as mind wandering and thoughts about the self. In contrast to this task-negative pattern, we show increased activity during a large and demanding switch in task set. Furthermore, we employ multi-voxel pattern analysis and find that regions of interest within default mode network are encoding task-relevant information during task performance. Activity in this network may be driven by major revisions of cognitive context, whether internally or externally focused.
Introduction

Functional magnetic resonance imaging (fMRI) has repeatedly demonstrated that cognitive tasks of many kinds decrease activity in a large-scale cortical network, variously termed the task-negative or default mode network (DMN) (Raichle et al. 2001; Shulman et al. 1997; Andrews-Hanna et al. 2010). The DMN consistently shows reduced activity during task performance compared to rest (Raichle and Snyder 2007) and often reduced activity for harder compared to easier task versions (Gilbert et al. 2012). In contrast to this general pattern, increased activity has been reported in a cluster of mental states involving thinking about the self, one’s own perspective, or the perspective of others (Buckner and Carroll 2007). Examples include recollecting previous experiences (Vincent and Snyder 2006) or imagining future ones (Addis, Wong, and Schacter 2007), mind-wandering (Mason et al. 2007) and theory of mind tasks (Young, Dodell-Feder, and Saxe 2010). The DMN has thus been linked to a number of high-level cognitive processes, such as self-referential processing (Gusnard et al. 2001) and imaginary scene construction (Hassabis and Maguire 2007).

Recently, Andrews-Hanna et al. (2010) have argued that the DMN separates into three sub-networks. Using graph theoretical analytic approaches to resting-state fMRI data, Andrews-Hanna et al. identified a core sub-network comprising bilateral posterior cingulate cortex (PCC) and anterior medial prefrontal cortex (AMPFC) (Figure 2b, yellow ROIs), a medial temporal lobe (MTL) sub-network made up of ventromedial prefrontal cortex (VMPFC), bilateral hippocampal formation (HF), parahippocampus (PHC), retrosplenial cortex (Rsp), and posterior inferior parietal lobule (pIPL) (green ROIs), and a dorsomedial prefrontal cortex (DMPFC) sub-network which includes the DMPFC, bilateral temporal parietal junction (TPJ), lateral temporal cortex (LTC) and the
temporal Pole (TempP) (blue ROIs). Andrews-Hanna et al. argue for a degree of
functional segregation between these sub-networks, with the MTL sub-network
especially linked to construction of mental scenes based on memory, and the DMPFC
network more involved in mentalising. These segregations, however, are likely to be
relative, with many experiments, for example, linking all three DMN sub-networks to
conscious recollection (Vilberg and Rugg 2012).

Here we consider a simple conceptualisation of DMN function and apply it within the
Andrews-Hanna framework. Arguably, imagination, mind-wandering, and taking
another’s perspective have in common a substantial change from the current cognitive
context. Similarly, conscious recollection is typically conceived as reactivation of a
previously-experienced episode, with components linked into a complex surrounding,
context. Substantial shifts of context may be common in everyday activity, e.g. a shift
from cooking dinner to giving directions to guests over the phone, but less common in
the constrained setting of typical laboratory tasks. For example, in a recent review of
neuroimaging studies of task switching (Kim et al. 2011), tasks that they argue required
a contextual switch involved either a change in simple attended features or binary
categorization rules using a fixed, small set of possible stimuli. Irrespective of specific
high-level processes involved, we reasoned that the DMN may be involved in any large
switch of cognitive context – an operation presumably calling for relaxation of many
aspects of a current attentional focus, with concomitant activation of representations
and processes relevant to the new context.

To test this hypothesis, we used a novel experimental paradigm that required
participants to switch between similar and dissimilar tasks, within a relatively large set
of 6 tasks (Fig. 1). The 6 tasks were each associated with a different rule, as determined
by the colour border surrounding the task stimulus. The tasks were split into three groups defined by stimulus category, with 2 possible tasks per stimulus type. A no-switch trial occurred when participants had to apply the same rule that was applied on the previous trial. A similar-task-switch trial – resembling switches in typical neuroimaging studies - occurred when participants had to apply the other rule from the same category as the previous trial. A dissimilar-task-switch occurred when participants had to apply a rule from a different category compared to the previous trial.

Contrary to the common concept of a task-negative system, we predicted increased DMN recruitment for the most difficult condition of switching between dissimilar tasks. We found this to be the case and furthermore, that the activity increase was selectively found in the Core and MTL sub-networks. In addition, we provide evidence that all sub-networks of the DMN represented task-related information during task performance.

**Results**

**Behavioral Results**

Accuracy on all tasks was high (median accuracy for all tasks > 95%, inter quartile range < 6%). As predicted, response time (RT) was significantly longer when switching between dissimilar tasks (2043 ms) compared both to trials when no task switch occurred (1670 ms; $t_{17}=8.6$, $p<0.001$), and to trials with switches between similar tasks (1746ms; $t_{17}=8.1$, $p<0.001$). Switches between similar tasks also produced significantly longer RTs compared to no-switch trials ($t_{17}=2.8$, $p=0.006$).

**Task-switch related activity in the DMN**

Preprocessing steps for fMRI data included realignment of the raw echo-planar images (EPI), slice-time correction, coregistration of the EPI images with the structural image,
normalisation to the Montreal Neurological Institute (MNI) template brain, smoothing
with an 8mm full-width at half-maximum Gaussian kernel and filtering with a high-pass
filter (see Materials and Methods). Univariate analysis of fMRI data was used to
compare dissimilar-task-switch with no-switch trials through the standard general
linear model (GLM) approach. A regressor was constructed for each switch type with
events modelled from stimulus onset until response and convolved with the
haemodynamic response function. The resulting beta values for each switch type were
compared and thresholded at $p<0.05$, correcting for the false discovery rate. We
identified widespread activation predominantly in regions of in the DMN (Fig. 2a), with
peaks found in bilateral HF, PHC, Rsp, PCC, AMPFC and left pIPL (Table 1). It is worth
noting that all of these regions fall within either the Core or MTL sub-networks. In
comparison, no regions from the DMPFC sub-network showed significant activation at
the whole-brain level. A contrast of similar-task-switch against no-switch trials revealed
no significant activation across the whole brain.

To examine changes in activation from the perspective of the DMN sub-networks, we
used individual DMN regions of interest (ROIs) previously defined (Buckner et al. 2009;
Andrews-Hanna et al. 2010). The mean beta value was extracted from each ROI
following each switch type. Planned, paired two-tailed t-tests revealed significant
increase in activity during dissimilar-task-switch compared to no-switch in core
(bilateral AMPFC, PCC) and MTL (Rsp, PHC) sub-networks, with a tendency to de-
activation in the DMPFC sub-network (significant in right TPJ) (Fig. 2c). Again, no ROIs
revealed a significant difference between the similar-task-switch trials and no-switch
trials. Two-way repeated measures ANOVAs were performed separately for each sub-
network, with factors of ROI (Core: 4, MTL: 9, DMPFC: 7) and switch type (no-switch,
dissimilar-task-switch). Main effects of task-switching were found for the Core 
sub-networks, showing increased 
activity for dissimilar switches. In contrast, the DMPFC sub-network showed a 
marginally significant de-activation \((F_{(1,17)}=4.1, p=0.06)\). Corresponding ANOVAs were 
performed to test for the difference between similar-task-switch and no-switch, but 
these revealed no main effect in any sub-network. To investigate differences at the sub-
network level, beta values were averaged across the ROIs within each sub-network each 
of the three trial types and a two-way repeated measures ANOVA (factors of sub-
network and switch type) was performed on the mean beta values. This analysis 
revealed a main effect of sub-network \((F_{(2,34)}=18.9 p<0.001)\) and an interaction of 
switch type and sub-network \((F_{(4,68)}=17.8 p<0.001)\). These data therefore show a 
dissociation within the DMN: While the DMPFC sub-network displayed the 
characteristic pattern of reduced activity during executive control, switching between 
dissimilar tasks showed an opposite pattern of increased activity in Core and MTL sub-
networks.

In an exploratory analysis we looked at the univariate activation associated with 
dissimilar switches between specific categories compared to no-switch trials, in the 
three DMN sub-networks, e.g. from a semantic task to a lexical task, compared with 
repetition of the lexical task. This was performed for all 6 between category switch 
types. Figure 3 shows that all types of switch showed a relative increase in Core sub-
network activation and decrease in DMFPC sub-network activation. The MTL sub-
network shows increases for 4 of the 6 switch types and a marginal decrease when 
switching from the perceptual task. Especially for the Core and DMPFC sub-networks, 
these data suggest little variation in the pattern of results across different task types.
Multivariate decoding demonstrates task representation across the DMN

For the multivariate analysis the same preprocessing pipeline was followed, with the omission of the smoothing step. We reasoned that if the DMN was involved in switching between tasks, then the differences between tasks might be represented within the network. To test this hypothesis we performed a multivariate pattern analysis on the same ROIs. For each ROI, classifiers were trained to discriminate between the voxel-wise pattern of activation for each task pair (6 tasks, therefore 15 task pairs) and these classifiers were subsequently tested on independent data using leave-one-run-out cross-validation (see Materials and Methods). The matrices in Fig. 4a show the classification accuracy (CA) for each task pair in each ROI, averaged across participants. The strongest decoding of task was found in bilateral HF, pIPL and the PCC, while bilateral TPJ, Rsp, AMPFC and DMPFC on the midline showed weaker but still significant decoding.

Decoding of task between dissimilar task pairs is likely driven by many differences in task features. In contrast, differences between similar tasks will be predominantly driven by the internal representation of the specific decision rule. To quantify the extent to which rule and other features were driving the CA scores, CAs for ROIs within the Core, MTL and DMPFC sub-networks were averaged separately for similar task pairs and dissimilar task pairs. This analysis (Fig. 4b) revealed significant decoding of dissimilar task pairs in all DMN sub-networks, and weaker but significant decoding of similar task pairs in the DMPFC sub-network.
Recently concern has arisen that differences in RT may be driving differences in CA (Todd, Nystrom, and Cohen 2013; for contrary arguments see Woolgar, Golland, and Bode 2014). We performed a regression analysis of CA against absolute difference in RT in each of the three sub-networks separately. First, we extracted the CA associated with each task pair in each ROI in each subject. We then calculated the mean CA across the component ROIs of the Core, MTL and DMPFC sub-networks in each individual, producing a 3-dimensional matrix of CA values for 3 sub-networks x 18 subjects x 15 task pairs. A similar matrix was produced for absolute RT differences. We segregated the dissimilar-task pairs and similar-task pairs and conducted a Spearman's correlational analysis of RT against CA for each task-pair type within each sub-network. The results showed strong discrimination of dissimilar task pairs, and weak discrimination of similar task pairs, irrespective of RT difference (Fig. 5a).

A second analysis considered data from each subject separately, with a separate regression analysis for similar and dissimilar task pairs in each ROI. A general linear model was constructed with a regressor for the absolute RT difference for either the similar-task-pairs or dissimilar-task-pairs, which was fit to the corresponding CA data. This produced a beta estimate for similar and dissimilar-task-pairs in each ROI, in each subject. Beta estimates were subsequently averaged across the component ROIs of each sub-network. Fig. 5b shows the mean beta estimate for the similar and dissimilar task pairs in each subject for the Core, MTL and DMPFC sub-networks. Within each graph, data from the 18 subjects are sorted in ascending order. Overall, the graphs suggest that response time does not systematically predict CA across participants, especially for the dissimilar task pairs (bottom row) for which CA was highest.
Together these analyses demonstrate that simple response time differences were not strongly driving the classification accuracies.

**Discussion**

In previous work, DMN activity has been associated with a variety of complex, often self-referential mental processes, such as retrieving past events from one’s life, imagining possible future events, or considering the beliefs of oneself and others (Buckner and Carroll 2007). Here we argue that a simple variable may relate these complex processes – the degree of change in cognitive context. To address this hypothesis we modified a typical task switching study to include both small shifts – similar to those of many previous studies – and much larger shifts. In contrast to the common idea of the DMN as a task-negative system, the activity of which progressively decreases with increasing task difficulty, our results show the opposite for a large change of cognitive context, in particular for Core and MTL sub-networks.

As cognitive context is changed, elements of the old context must be suppressed and elements of the new context must be retrieved or activated, producing a reconfiguration appropriate to the new circumstances. Our data suggest DMN activation only when the change is sufficiently large, perhaps analogous to many of the shifts taking place in everyday cognition. It is uncertain whether large and small shifts differ qualitatively or only quantitatively. In our task, for example, a shift between rules within the same category may have been executed without reference to the link between frame color and categorization rule; to perform such a switch, it was necessary only to see that frame color had changed and to retrieve the other rule relevant to the current set of
stimuli. In contrast, a shift between categories likely required reference back to a broader set of task rules, including the remembered list of color-rule combinations. More broadly, one possibility is that the DMN is involved in relaxing a current attentional focus, allowing new cognitive contents to arise. When cognitive operations are largely similar across successive trials, the DMN may be suppressed, but as increasingly more of the current focus must be dissolved, suppression may shift to activation.

Thus it is the magnitude of switch involved that distinguishes this study from much of the previous work in the literature. The point is illustrated by a detailed consideration of the task-switching review by Kim et al. (2011), as described in the Introduction. Kim et al. (2011) identified three major types of task-switch found in the literature: perceptual switch, response switch and contextual switch, the latter of which is the most comparable to the current work. Of the 20 tasks that they identified as involving a contextual switch, 17 used fixed small stimulus sets, with the switches (like our within-category switches) concerning just the relevant stimulus feature such as colour or shape. Two studies used a task that required participants to make one of two possible binary decisions on serially presented letter strings. The remaining task was essentially our semantic task, using text instead of images. Unlike these studies, we propose that it is our use of much more substantial switches, requiring larger revisions of cognitive context and operations, that leads to the novel finding of DMN activation.

Much previous work links activity in the DMN to conscious recollection, commonly defined by the ability to link a remembered item to the surrounding context of a specific event. A role in memory for individual events fits well with the proposal that the DMN binds together the components of a complex cognitive context. Beyond traditional
studies of episodic and autobiographical memory, our results show the importance of
DMN context processing in cognitive control. In contrast to the common finding of
deactivation linked to executive control, our findings show increased activity when the
change of context is sufficiently large. Establishing a new, complex context, we suggest,
may be common to recollection of specific previous events and to major revisions of
complex task rules.

An intriguing aspect of the present results is the difference between the regions that
show switch-related activity and those that encoded task-related information. The
regions that demonstrate switch-related activity show relatively strong correspondence
to the pattern of DMN fractionation presented by Andrews-Hanna et al. In contrast, the
DMN regions that show strong encoding extend across all three sub-networks,
suggesting only partial functional segregation. Encoding analysis is likely more sensitive
than univariate activity analysis, reflecting both overall activity within a broad region,
and the exact pattern of activity within that region (Davis et al. 2014). Though DMPFC
showed no univariate signal linked to switching task category, our results suggest some
involvement in task representation.

Our results show very different levels of CA for discrimination of similar vs. dissimilar
tasks. For the comparison of similar tasks, other than the different colour of border
surrounding the task item, the only difference is the internal representation of task rule.
In contrast, comparing dissimilar tasks involves a myriad of differences such as visual
properties of the stimuli, cognitive domain (semantic knowledge, lexicon, visual
discrimination) as well as task rule. Correspondingly, much stronger decoding is seen
across the DMN between dissimilar tasks. This is unsurprising for a network of regions
hypothesised to link components of a broad cognitive context.
Many studies suggest sustained DMN activity in rest compared to active task performance. It is unclear how such sustained activity relates to the switch-related activity we have shown here. On the one hand, it is plausible that, when a participant lies in a scanner at rest, there are periodic large shifts in the content of cognition, and in part, “sustained” DMN activity may reflect averaging across shifts occurring at variable, unknown times. Indeed, traditional studies of resting state functional connectivity depend on temporal variation in network activity, as expected for a signal in part linked to transient cognitive events. That said, if a core aspect of DMN function is relaxing an attentional focus, sustained enhancement is plausible during a period of relatively unfocused cognitive activity.

An unexpected aspect of our results is the lack of switch-associated activity when changing between similar tasks, which does show a robust behavioral cost compared to no switch trials. In previous studies, apparently similar cases of task switching have been associated with widespread recruitment of a fronto-parietal, executive control network (Braver, Reynolds, and Donaldson 2003; Kim et al. 2011; Yeung et al. 2006; Sohn et al. 2000; Monsell 2003), and it is unclear why no similar activity was seen in our data. One contributing factor may be our explicit modelling of RT differences between conditions, convolving the canonical haemodynamic response with the duration of each trial from stimulus presentation until response. As this procedure is designed to correct for activation differences due simply to longer RT on switch trials, it may reduce or remove differences seen in studies that do not adopt such a correction. Our results also raise the possibility, however, that traditional task-switching results may not generalise to the more complex conditions of our experiment. Future work will be needed to resolve this discrepancy.
Conclusion

In conclusion, we propose that the DMN may be recruited whenever large changes of cognitive context are required. This may apply in complex cases of self-referential processing, mind wandering etc., but also in relatively simple acts of cognitive or executive control. The DMN, widely seen as a ‘task-negative’ network, may respond positively to any task which demands a switch from one broad context to another.

Materials and methods

Participants

18 right-handed participants (10 females) aged between 18 and 40 were recruited from the Medical Research Council Cognition and Brain Sciences Unit subject panel. Of 21 original subjects scanned, 3 had to be removed for excessive head-movements (over 10 mm translation and/or 6° rotation). No participant had a history of neurological or psychiatric illness. Participants were reimbursed for their time. Ethics approval was given by the Cambridge Psychology Research Ethics Committee.

Task Description

All three tasks were created using the Psychophysics Toolbox for MATLAB (Brainard 1997). Within the scanner, the stimulus display was projected onto a mirror mounted to a 32-channel head-coil.

Participants were required to learn 6 different tasks (Fig. 1a). Each task was associated with a different rule, with the appropriate rule determined by the colour border in which the task stimulus appeared. The 6 tasks rules are shown in Fig. 1a. Prior to
scanning, participants practised the task until they had completed at least 20 trials with
an accuracy exceeding 80%. Importantly, the 6 tasks were grouped into three categories
of 2 tasks each, where the stimuli within a category could be relevant to either rule
within that category, but not to rules of other categories. Furthermore, all categories
included trials which required a positive answer for both rules, for one rule but not the
other, or for neither rule; therefore subjects needed to remember and apply the correct
rule on all trials. All questions were framed in a true/false format, so that arbitrary
response mappings for each rule did not have to be learned in addition to the rules
themselves.

Each trial began with the simultaneous appearance of the colour border (visual angle =
7.9°) and the task stimulus (Fig. 1b). Participants were requested to respond as quickly
as possible with a true or false answer (right thumb button press = true, left thumb
button press = false). The border and stimulus remained on screen until the subject had
made their response. A low tone was played to participants if they made an incorrect
response. There was a jittered interval between the response to one trial and the onset
of the next. Interval jittering followed an exponential distribution between 1s and 11s,
with a mean of 4.1s.

Participants learned the tasks prior to scanning. An event-related design was adopted,
with 73 trials per run. Each run had at least 12 trials of each of the 6 task types. Task
switch type was also balanced within a run: 24 no-switch trials, 24 similar-task-
switches and 24 dissimilar-task-switches. Post scanning, when questioned, no
participants reported having any sense of what task to expect on a given trial.

fMRI Acquisition
Scans were acquired with a 3T Siemens Trim Trio scanner. 32 3-mm slices (0.75mm interslice gap) in axial orientation gave an in-plane resolution of 3x3mm and were acquired using a TR of 2s. T2*-weighted echo-planar imaging (EPI) capturing blood oxygen level dependent (BOLD) contrast was employed with a flip angle of 78°. For both experiments the first 8 images were discarded to avoid T1 equilibration effects.

**Univariate Analysis**

For the univariate analysis, images were preprocessed and analysed with SPM5 (Wellcome Department of Cognitive Neurology). In the first preprocessing step, data were checked for obvious artefacts, and all images were realigned to the first image. Next we performed slice time correction and coregistration of the structural with the functional EPI images. Finally, data were normalized to the standard Montreal Neurological Institute (MNI) template, smoothed with an 8mm full-width at half-maximum Gaussian kernel and subjected to a high-pass filter with cut-off at 128s.

Fixed-effects analyses were performed on each individual's data using a general linear model. In the first univariate analysis investigating switching related activity 3 regressor functions were used (no-switch trials, similar-task-switch trials and dissimilar-task-switch trials). Each regressor was modelled as a rectangular function from the onset of each stimulus to the moment of response and convolved with the canonical hemodynamic response function. Beta weight images were contrasted for the conditions dissimilar-task-switch > no-switch and similar-task-switch > no-switch. Contrasts were further examined by random-effects analysis. Activation maps (threshold 0.05, FDR-corrected) were visualised using MRIcroGL (Rorden, Karnath, and Bonilha 2007).
For ROI analysis, mean contrast values within each ROI were extracted for each subject, using the MarsBaR SPM toolbox (Brett et al. 2002). ROIs were spherical, with 8mm radius, based around peak coordinates (Fig. 2b) taken from Andrews-Hanna et al. (2010).

For the exploratory analysis into the activation associated with switches between specific categories a separate GLM was constructed. 36 regressors were used (one regressor for each possible switch type) and modelled as before. The resulting beta estimates were then processed using the same ROI analysis method as before with the same ROIs.

**Multivoxel pattern analysis**

Multivoxel pattern analysis (MVPA) was performed using the Decoding Toolbox (Christophel, Hebart, and Haynes 2012; Görgen, Hebart, and Haynes 2012). Preprocessing of the data was the same as for the univariate whole-brain analysis, except for the omission of the smoothing step. Again, a fixed effects analysis was performed on each participant’s data using a general linear model. For this GLM, each task was modelled as a separate regressor, constructed as a rectangular function from the onset of each stimulus to the moment of response and convolved with the canonical hemodynamic response function. The same ROIs as previously (Fig. 2b) were used.

Prior to pattern analysis within each ROI, beta values were Z-scored across all voxels within the ROI, separately for each task. This step was intended to reduce any impact of task differences in overall ROI activity. Pattern discrimination between tasks was then estimated using pairwise classification, i.e. only 1 of the 15 possible task pairs was
decoded at a time. A support vector machine (LIBSVM) (Fan, Chen, and Lin 2005) was used to train and classify data from three of the four runs, with the remaining run used to test the classifier. Test and training runs were always kept separate and each run was used to test the classifier once, i.e. fourfold cross-validation. The classification accuracy (CA) for a given ROI was averaged across test-train splits, yielding a single CA for each ROI, in each individual, for each task pair.
Acknowledgements:

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Table 1. Peak coordinates of DMN regions that showed significantly greater activation for dissimilar-task-switch over no-switch. Coordinates are in MNI space.

HF = hippocampal formation, PHC = parahippocampus, Rsp = retrosplenial cortex, PCC = posterior cingulate cortex, AMPFC = anterior medial prefrontal cortex, pIPL = posterior inferior parietal lobe.

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Figure 1. Task Description. The experiment required participants to learn 6 task tasks prior to scanning. (a) The 6 tasks were each associated with a different rule, as determined by the colour border. The tasks were split into three groups defined by stimulus category, with 2 possible tasks per stimulus type. (b) Experimental design. Within each run, trials using the 6 tasks occurred in random order. A no-switch trial occurred when participants had to apply the same task that was applied on the previous trial. A similar-task-switch trial occurred when participants had to apply the other task from the same category as the previous trial. A dissimilar-task-switch occurred when participants had to apply a task from a different category compared to the previous trial.

Figure 2. Activation of the DMN for dissimilar task switches. Region labels and ROIs are color-coded according to the sub-network to which they belong: yellow for Core, green for MTL, blue for DMPFC. (a) Whole brain rendering in axial slices: the numbers above each slice indicate z-coordinate of that slice. The contrast of dissimilar-task-switch > no-switch ($T=3.23$, $p<0.05$, FDR corrected) shows activations in regions previously identified as the DMN. (b) Locations of DMN ROIs distinguished by Andrews-Hanna et al. (c) Change in activation of similar-task-switch (darker colours) and dissimilar-task-switch (lighter colours) relative to no-switch trials in the DMN ROIs. APMFC: anterior medial prefrontal cortex, PCC: posterior cingulate cortex, pIPL: posterior inferior parietal lobe, Rsp: retrosplenial cortex, PHC: parahippocampal cortex, HF: hippocampal formation, VMPFC: ventromedial prefrontal cortex, TPJ:
temporoparietal junction, LTC: lateral temporal cortex, TempP: temporal pole, DMPFC: dorsomedial prefrontal cortex. * indicates p<0.05.

**Figure 3. Activation associated with each between category switch.** An exploratory analysis looking at the activation/deactivation associated with switching between each of the three task categories. Core and MTL sub-networks predominantly show increased activation following a dissimilar task switch across switch types, whereas DMPFC shows a relative decrease in activation. Abbreviations: sem = semantic category, per = perceptual, lex = lexical. * denotes p<0.05 from a paired, two-tailed t-test.

**Figure 4. Classification accuracy within the DMN sub-network ROIs.** (a) Classification accuracies between different task pairs in all DMN ROIs. Large correlation matrix used as example is the same as left HF. The colour of the circle in the key matches with the colour borders used to indicate each task in Fig. 1a: Red for 'bigger than a shoebox?', blue for 'living?', brown for 'same shape?', pink for 'same height?', green for 'does adding A make a word?', purple for 'does adding I make a word'? Matrices show the classification accuracy of decoding each task pair; values below the diagonal show classification accuracy for all task pairs, while non-grey values above the diagonal show only decoding that survived the threshold for statistical significance. The colour borders indicate the sub-network that the ROIs belong to: core (yellow), MTL (green) and DMPFC (blue). ROIs on the left side of each box are from the left hemisphere, those on the right are from the right hemisphere. (b) All three sub-networks demonstrated above-chance classification accuracy when decoding dissimilar tasks, while only the DMPFC sub-network demonstrated significant decoding of similar task pairs. Error bars indicate standard error. * indicates p<0.05.
Figure 5. The influence of response time on classification accuracy. (a) Correlation between classification accuracy and RT difference in the three DMN sub-networks. Each point represents data for a single task pair in a single subject, with mean CA across ROIs of the named sub-network plotted against absolute RT difference. The darker shades in each graph are taken from the similar task pairs, while the lighter shades are taken from the dissimilar task pairs. (b) Beta estimates for the association of CA and RT in each subject for similar and dissimilar task pairs of the three DMN sub-networks. In each graph subjects’ beta estimates are sorted in ascending order. Top row, a-c, displays beta estimates for similar task pairs in the Core, MTL and DMPFC respectively. Bottom row, d-f, shows beta estimates for dissimilar task pairs in the Core, MTL and DMPFC respectively. The p-value from a 2-tailed, 1-way t-test of each graph’s beta values is shown.
(a) Is it larger than a shoebox? Does 'A' fit to make a word? Are they the same height?

(b) similar rule switch

no switch
dissimilar rule switch

B_TH