

## Video Article

# Brain Imaging Investigation of the Neural Correlates of Emotional Autobiographical Recollection

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## Abstract

Recollection of emotional autobiographical memories (AMs) is important to healthy cognitive and affective functioning<sup>1</sup> - remembering positive AMs is associated with increased personal well-being and self-esteem<sup>2</sup>, whereas remembering and ruminating on negative AMs may lead to affective disorders<sup>3</sup>. Although significant progress has been made in understanding the brain mechanisms underlying AM retrieval in general (reviewed in<sup>4,5</sup>), less is known about the effect of emotion on the subjective re-experience of AMs and the associated neural correlates. This is in part due to the fact that, unlike the investigations of the emotion effect on memory for laboratory-based *microevents* (reviewed in<sup>6,7-9</sup>), often times AM studies do not have a clear focus on the emotional aspects of remembering *personal* events (but see<sup>10</sup>). Here, we present a protocol that allows investigation of the neural correlates of recollecting emotional AMs using functional magnetic resonance imaging (fMRI). Cues for these memories are collected prior to scanning by means of an autobiographical memory questionnaire (AMQ), therefore allowing for proper selection of emotional AMs based on their phenomenological properties (i.e., intensity, vividness, personal significance). This protocol can be used in healthy and clinical populations alike.

## Video Link

The video component of this article can be found at <http://www.jove.com/details.php?id=2396>

## Protocol

### 1. Collection and Selection of Memories, Imaged Task, and Experimental Protocol

#### Collection of the Emotional AMs

1. Personal memories are elicited from each participant during an interview performed prior to the fMRI session, similar to the procedure employed for non-emotional AMs (e.g.,<sup>11,12</sup>). Unlike most previous techniques, our AMQ is specifically constructed to target the assessment of emotional personal episodes and their recollective properties. A commonly used method in behavioural research and adapted for use in neuroimaging studies involves the Crovitz and Schiffman technique<sup>13-15</sup>, where subjects remember specific, personally experienced events in response to cue words. A photo-paradigm has been also used to investigate AMs in more controlled settings in fMRI studies<sup>16,17</sup>. However, these techniques do not typically target examination of the emotional component of AMs, that are matched in terms of recollective qualities.
2. Our AMQ comprises a list of 115 verbal cues for distinct life events (e.g., the birth of a family member, the death of a relative), which is a combination and extension of lists used by others<sup>18,19</sup>. For each cue, participants are asked to remember a unique episode from their life, that occurred in a specific place and time (e.g., one instance when s/he played in a specific basketball game), rather than remembering general or repeated events (e.g., playing high school basketball). Importantly, the memories must be accompanied by the recollection of being personally involved, rather than hearing about them from others.
3. Upon recollection, participants are asked to provide a brief description of the memory (see Figure 1), which will then be used as a personalized memory cue during fMRI scanning (participants are naïve to the specific purpose of the pre-scanning interview).
4. Each memory is also dated and rated on six Likert scales<sup>20,21</sup>, to assess its phenomenological properties (see Figure 1). Our scales included *emotional valence* (using a 7-point scale: -3 = very negative, 0 = neutral, +3 = very positive), *emotional intensity*, *personal significance*, the amount of *contextual details*, the amount of *visuo-perceptual details* (i.e., vividness), and the *frequency of retrieval* (all of the latter using a 7-point scale: 1 = not at all, 7 = extremely).

**Figure 1. Illustration of AMQ Administration.** For each cue, participants remember and briefly describe a specific event, and then date and rate it on 6 scales.

**Selection of Highly Emotional AMs**

5. Next, the 40 most emotional memories (20 positive and 20 negative) are selected for each participant, based on the ratings provided on the AMQ (i.e., rated 2 or 3 and -2 or -3, respectively). The positive and negative AMs are then matched in terms of age and phenomenological properties, to ensure that any differences in brain activity during recollection are not confounded by differences in these basic properties.
6. If necessary, the descriptions of the selected memory cues are slightly adapted to be matched as closely as possible for length and grammatical complexity; it is also recommended to select a few additional memories for practice purposes.

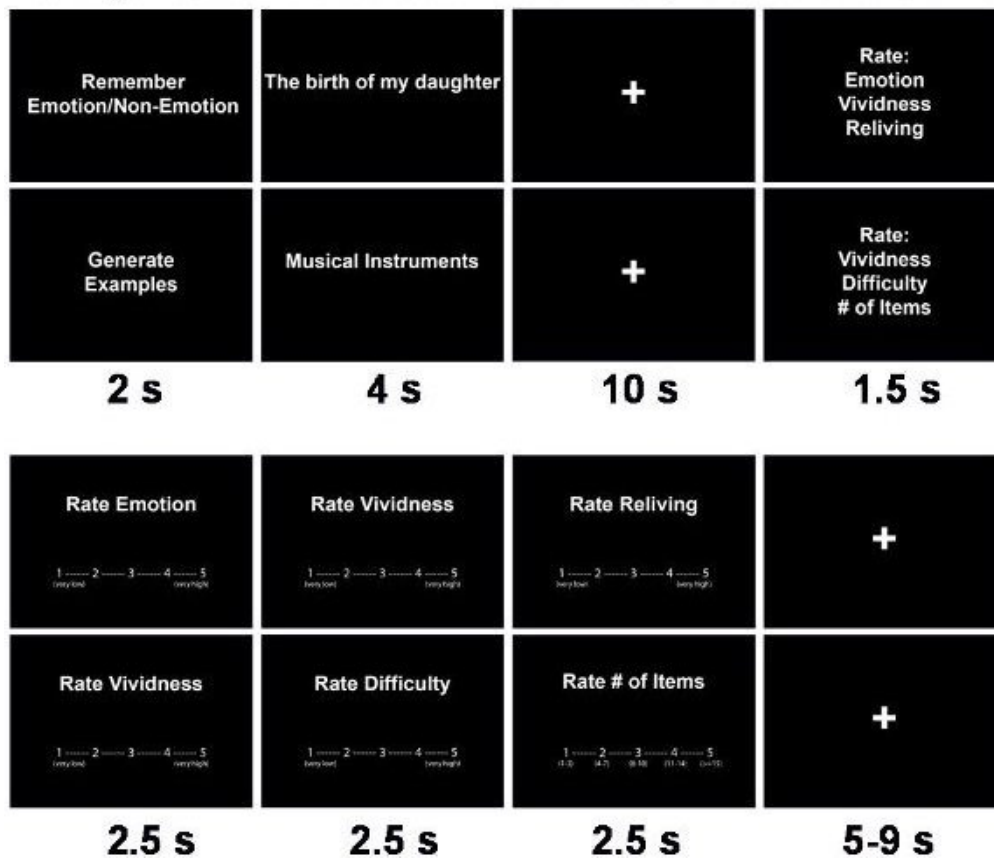
**The fMRI Task**

7. The fMRI task is designed to allow comparison of the AM task with a semantic memory (SM) control task; the AM and SM retrieval trials have a similar general structure (see Figure 2A). We used CIGAL (<http://www.nitrc.org/projects/cigal/>) for stimulus presentation in the MR scanner, but other stimulus presentation software may also be used.
8. The AM task is based on the personalised cues collected prior to scanning. Each trial begins with a cue that triggers AM recollection, which is indicated by the participant with a button press. Then, participants continue remembering details of the event until cued again to rate the recollected memory (Figure 2B).
9. The SM task involves generation of exemplars from different semantic categories (e.g., musical instruments, sports), which like AM retrieval involves search in memory and extended retrieval time<sup>22</sup> (Figure 2B). After a semantic category is cued, participants press a button as soon as they start recalling exemplars from that category and then continue recalling until cued again for memory ratings.

**A. General Structure of the Trials**

Task Instruction AM/SM	Memory Cue	Recollection	Ratings Instruction
Rating Cue 1	Rating Cue 2	Rating Cue 3	Inter-Trial Interval

## B. Specific Structure of the AM and SM Trials



**Figure 2. Structure of the fMRI Trials.** A. General Structure of the trials. B. Specific structure of the AM and SM trials.

10. In addition to the basic comparison of AM and SM trials, other manipulations can also be involved. For instance, the focus of the AM retrieval can be manipulated by directing the participants to pay attention to emotional or non-emotional aspects of the remembered events (Figure 2B). This manipulation allows investigation of how retrieval focus may influence the experiencing of AMs and any associated changes in the associated neural correlates.
11. Each AM or SM trial is followed by rating screens presented in a counterbalanced order, using 5-point Likert scales (see Figure 2B).
12. The experiment is split into runs/blocks of trials both to allow participants time to rest and to avoid data loss in case of equipment malfunction. Run order is counterbalanced between participants. Each run begins with six seconds of fixation, to allow stabilization of the MR signal. The AM and SM conditions are presented in a random order separated by an inter-trial interval of variable duration (5-9 sec, average = 7sec.).

## 2. Preparing the Subject for the Scan

All participants provide written informed consent prior to running the experimental protocol, which is approved by an Ethics Board. Typically, to avoid confounds in the lateralization of brain activations, scanned participants are right-handed.

### Prior to Entering the Scanning Room

1. On the day of scanning, participants' current affective state is assessed<sup>23</sup>, to control for the effect of mood on recollection of emotional AMs. In conjunction with post-scanning assessments, these initial evaluations can also be used to screen for changes in mood as a result of the study participation, and as covariates in the fMRI analyses to investigate brain activations influenced by current states. Similarly, assessments of personality traits can also be made (e.g., neuroticism), to investigate their possible biases on the AM task<sup>24</sup> and the associated neural correlates.
2. Prior to the scan, participants are informed in detail of the scan procedures, and are given specific instructions for the behavioral task. Participants also complete a short practice session, to familiarize themselves with the task.

### Entering the Scanning Room

3. Participants are instructed to lie supine on the scanning bed, and are provided with additional head cushioning, to ensure comfort during the scan and to minimize movement. To further minimize head movement, the non-adhesive side of a length of tape may be wrapped lightly around the subjects' forehead. Subjects are given ear protection as well as isolation headphones to communicate with the experimenter during the MRI scan.
4. Subject's right hand is positioned comfortably on the response box, thus allowing the left hand to be used for support or for other measurements (e.g., skin conductance responses). An emergency stop button is placed nearby so that the subject may indicate any urgent need to stop the scanner.
5. Before starting data collection, it is critical to make sure that subjects can see the screen projection clearly for stimulus presentation and that the response buttons work properly.

### 3. Data Recording and Processing

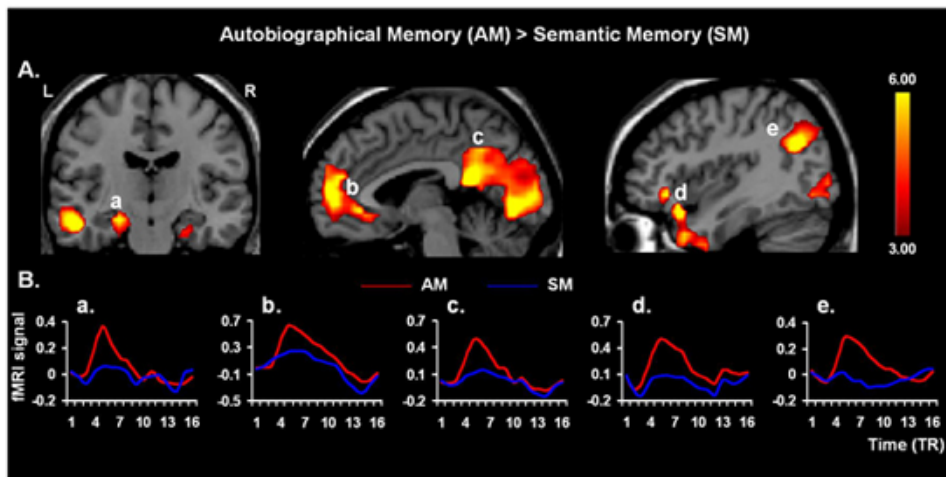
#### Scanning Parameters

We collected MRI data using a 1.5 Tesla Siemens Sonata scanner for MRI recordings. Anatomical images were 3D MPRAGE anatomical series (repetition time (TR) = 1600 ms, echo time (TE) = 3.82 ms; number of slices = 112; voxel size = 1x1x1 mm), and functional images were series of 28 functional slices, acquired axially using an echoplanar sequence (TR = 2000 ms; TE = 40 ms; field of view FOV = 256x256 mm; voxel size = 4x4x4 mm), thus allowing for full-brain coverage.

#### Data Analysis

We used Statistical Parametric Mapping (SPM: <http://www.fil.ion.ucl.ac.uk/spm>) in combination with in-house Matlab-based tools. Pre-processing involved typical steps: quality assurance, TR alignment, motion correction, co-registration, normalization, and smoothing (8 mm<sup>3</sup> Kernel). Individual and group-level statistical analyses may involve comparisons of brain activity according to memory type (AM vs. SM), emotional valence (positive vs. negative), and retrieval focus (emotional vs. non-emotional content).

### 4. Representative Results



**Figure 3. Neural Correlates of AM Retrieval.** Validating the present protocol, retrieval of AMs yielded increased activity in the AM retrieval network<sup>4, 25</sup>, including hippocampal areas (a), involved in general memory retrieval, the medial prefrontal cortex (b), associated with personal engagement, the cuneus/precuneus regions and parieto-occipital junction (c, e, respectively), associated with processing visuo-spatial representation, and the frontal-temporal junction (d), involved in affective AM retrieval; similar effects to the latter were also found in the amygdala (not shown). The "activation maps" are superimposed on high resolution brain images displayed in coronal (left panel) and sagittal (middle and right panels) views; the color bars indicate the gradient of *t* values of the activation maps ( $p < 0.005$ , 10 contiguous voxels<sup>26</sup>), reflecting brain activity time-locked to the onsets of the memory cues. The line graphs illustrate the time courses of the fMRI signal (% signal change), for each trial type and TR (1 TR = 2 seconds). L = Left; R = Right.

#### Discussion

The experimental design introduced here allows investigation of the neural correlates of remembering emotional autobiographical memories. This design has the potential to advance our knowledge of how the brain generates affective biases (positive or negative) in remembering personal memories, and how those biases may be modulated by the retrieval focus (on emotional or non-emotional aspects). This protocol has additional benefits in that it can also be used with clinical populations (e.g., in patients with depression and post-traumatic stress disorder), which allows investigation of alterations associated with negative affective biases in AM retrieval (e.g., rumination on negative experiences and uncontrollable recollection of traumatic events, respectively). Overall, the success of this design depends on careful AM collection and selection and proper experimental manipulations.

#### Disclosures

No conflicts of interest declared.

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