Default and control network connectivity dynamics track the stream of affect at multiple timescales

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Abstract

In everyday life, the stream of affect results from the interaction between past experiences, expectations and the unfolding of events. How the brain represents the relationship between time and affect has been hardly explored, as it requires modeling the complexity of everyday life in the laboratory setting. Movies condense into hours a multitude of emotional responses, synchronized across subjects and characterized by temporal dynamics alike real-world experiences. Here, we use time-varying intersubject brain synchronization and real-time behavioral reports to test whether connectivity dynamics track changes in affect during movie watching. The results show that polarity and intensity of experiences relate to the connectivity of the default mode and control networks and converge in the right temporoparietal cortex. We validate these results in two experiments including four independent samples, two movies and alternative analysis workflows. Finally, we reveal chronotropic connectivity maps within the temporoparietal and prefrontal cortex, where adjacent areas preferentially encode affect at specific timescales.

Key words: default mode network; control network; naturalistic stimulation; fMRI; affect

Introduction

Emotions are intense and immediate reactions of the body and the brain to an external or internal event occurring in the present, happened in the past or may occur in the future. In daily life, the temporal trajectory of emotions is characterized by regularities in features, such as the time to rise, the duration or the probability of resurgence (Kuppens and Verduyn, 2017), which inform mental models of emotion transitions and support affective forecasting (Thornton and Tamir, 2017). Whence, it appears that the study of diachronicity is crucial for building a more comprehensive understanding of emotion processing and conceptualization in humans. Indeed, over the last years, increasing attention has been dedicated to the study of the temporal characteristics of affect using behavioral and experience sampling methods (Kuppens et al., 2010; Trampe et al., 2015; Verduyn et al., 2015), allowing researchers to track emotions over days, weeks or months. In sharp contrast, studies on the brain correlates of affective dynamics are still limited (Waugh and Schirillo, 2012; Costa et al., 2014; Verduyn et al., 2015; Résibois et al., 2017, 2018) and have employed static or relatively brief stimuli (Posner et al., 2009; Baucom et al., 2012; Kim et al., 2017) that may not be adequate to account for the complex temporal trajectory of lifelike experiences (Waugh and Schirillo, 2012).

A possible strategy to overcome this limitation is measuring brain activity elicited by movies using functional magnetic resonance imaging (fMRI). Indeed, movies are ecological and dynamic stimuli that synchronize brain response across individuals (Hasson et al., 2004) and elicit a wide variety of emotional states in a few hours (Schaefer et al., 2010). In addition, movies represent an effective tool for tracking time-varying brain connectivity (Nastase et al., 2019) and revealing the interplay among large-scale networks. Promising evidence in this regard comes from a work that, using relatively brief movie excerpts (∼8 min), demonstrated the association between the connectivity of salience and amygdala-based networks and the perceived intensity of sadness, fear and anger (Raz et al., 2016).

Also, brain connectivity dynamics well exemplify the tight relationship between emotion and cognition, as the same brain region can interact with distinct functional networks over time, and the same network can support multiple mental processes (Pessoa, 2017, 2018). In line with this, studies combining naturalistic stimulation with brain connectivity measurements show
that the default mode network (DMN) contributes to information encoding (Simony et al., 2016), memory representations (Chen et al., 2017) and prediction of future events (Antony et al., 2021). Interestingly, it has been argued that the same network is also crucial for constructing emotional experiences (Satpute and Lindquist, 2019) and integrating the self with the social world (Yeshurun et al., 2021).

Here, we test the hypothesis that changes in DMN connectivity would reflect the temporal properties of emotion dynamics experienced by individuals. In a first experiment, we explore whether, during the viewing of a 2h movie (Forrest Gump; Hanke et al., 2016), subjective reports of the affective experience explain brain connectivity dynamics in independent participants watching the same film. To rule out the possibility that results depend on the selected stimulus, we validate our findings in a second experiment by acquiring emotional reports and brain data in other individuals presented with a different live-action movie (i.e. 101 Dalmatians). Also, given the impact of analytical choices on brain imaging results (Botvinik-Nezer et al., 2020), we test the generalizability of findings by pre-processing fMRI data using two alternative workflows. Finally, in light of topographies revealed in high-order associative areas (Baldassano et al., 2017), we test the existence of a chronotopic organization (Protopapa et al., 2019) in regions encoding the stream of affect.

Materials and methods
Experiment 1: behavioral study

Behavioral data have been previously reported (Lettieri et al., 2019) and made freely available at https://github.com/psychoinformatics-de/studyforrest-data-perceivedemotions. In brief, 12 Italian native speakers (5F; mean age 26.6 years, range 24–34) provided ratings of their moment-by-moment emotional experience while watching the Forrest Gump movie (Zemeckis, Paramount Pictures, 1994; for details see Supplementary Data).

They were presented with an Italian dubbed and edited version of Forrest Gump, split into eight movie segments ranging from 11 to 18 min. Subjects were asked to report (i.e. 10 Hz sampling rate) the type and intensity of their inner emotional state using six emotion categories (happiness, surprise, fear, sadness, anger and disgust, Ekman, 1992). As the complexity of the experience elicited by Forrest Gump cannot be merely reduced to six emotions, we allowed participants to report combinations of categories throughout the movie simultaneously. Our approach revealed that behavioral ratings captured 15 different affective states (see Lettieri et al., 2019). Starting from categorical descriptions, we then obtained the underlying affective dimensions—i.e. polarity, complexity and intensity—using principal component analysis (see Supplementary Data).

For each subject, the behavioral data acquisition’s overall duration was 120 min (72 000 timepoints). Stimulus presentation and recording of the responses were implemented in Matlab (R2016b; MathWorks Inc., Natick, MA, USA) and Psychtoolbox v3.0.14 (Kleiner et al., 2007).

Experiment 1: fMRI study

Brain activity elicited by the watching of Forrest Gump was obtained from the studyforrest project—phase II (n = 14, German native speakers; Hanke et al., 2016). Details on brain data acquisition and pre-processing can be found in Supplementary Data.

To track time-varying brain connectivity, we opted for the intersubject functional correlation (ISFC) approach, which computes the correlation between the signal change of a specific brain region in one subject with the activity of all other regions in other subjects (Simony et al., 2016). Compared to other functional connectivity measures, ISFC demonstrates a higher signal-to-noise ratio as it inherently filters out activity not related to stimulus processing and artifacts not correlated across brains. In addition, ISFC can be used to study the dynamic reconfigurations of brain networks using a moving-window approach. We estimated the static and time-varying ISFC (tISFC) associated with the watching of Forrest Gump using custom MATLAB scripts, freely available at https://osf.io/v8r9w.

As a first step, we selected the atlas provided by Schaefer et al. (2018) and parcellated each subject’s brain into 1000 cortical regions of interest (ROIs). We opted for this atlas as, compared to other parcellations (e.g. Shen et al., 2013), it provides a finer-grained segmentation of the cerebral cortex that, at the same time, reflects the seven resting state networks (Yeo et al., 2011): visual, somatomotor, ventral attention, dorsal attention, limbic, control and DMN. In addition, as Schaefer parcellation does not include subcortical regions, we used Harvard-Oxford subcortical atlas (Makris et al., 2006) to define 14 additional ROIs (7 per hemisphere): nucleus accumbens, amygdala, caudate, hippocampus, pallidum, putamen and thalamus. Thus, in total, the parcellation we adopted comprises 1014 areas.

To compute static ISFC, the time course of brain activity was extracted from each ROI of one subject and correlated using Pearson’s coefficient with the remaining 1013 regions of all other subjects. By repeating this procedure for each of the 14 participants, we generated 91 (i.e. number of all possible combinations of subjects) correlation matrices. We then applied Fisher z-transformation and averaged correlation values across subjects to produce a group-level ISFC map.

To estimate tISFC, one has first to determine the width of the moving-window and the degree of overlap between adjacent windows. Both parameters are often chosen arbitrarily, even though they may significantly affect the results. To limit arbitrariness, we estimated the optimal window width as a function of the intersubject correlation in affective dimensions obtained from the behavioral experiment. Specifically, we computed the average intersubject correlation of affective dimension timeseries for all the possible window widths, ranging from (1) 20 to 250, (2) 20 to 500, (3) 20 to 750 and (4) 20 to 1000 timepoints (i.e. from 40 s to 33 m) and having 33% of overlap (Figure 1A and B). Of note, although the upper bound of the explored intervals (i.e. 250, 500, 750, 1000) is arbitrarily selected, the optimal window size is determined by a data-driven approach. Specifically, the optimal timescale was defined for each range as the point at which the intersubject correlation curve starts to flatten (i.e. knee point), namely the point representing a reasonable trade-off between the agreement across participants and the window width (Figure 1B). Depending on the dynamics of emotional ratings, this procedure provides up to four optimal window widths (i.e. one for each range), and it was repeated for each affective dimension separately. The reason for exploring four timescales is that distinct brain regions and process narratives at multiple temporal resolutions (Baldassano et al., 2017), and a similar mechanism could be hypothesized for the processing of affective information.

Once optimal window widths were established, we computed the tISFC as the Pearson’s correlation between each ROI of one subject and all the other ROIs of the other subjects in each window.
The primary aim of our study was to relate changes in tISFC to variations in the perceived affective state during movie watching. Therefore, for each ROI, we computed connectivity strength as the sum of the z-transformed correlation values at each timepoint (Figure 1D). We thus obtained 1014 timeseries expressing the relationship of each ROI with the rest of the brain throughout the movie. These timeseries were then correlated using Spearman’s \( \rho \) coefficient with the group-level timeseries of affective dimensions, after those were downsampled to the optimal window widths using a moving-average procedure. To assess the significance of the association (Figure 1E), we used a non-parametric permutation test based on timepoint shuffling of affective dimensions (200,000 iterations; minimum two-tailed \( P \)-value: 1.0e–5). At each iteration, randomized affective dimensions were correlated with ROIs connectivity strength, providing null distributions of Spearman’s correlation values. The position of the actual association in the null distribution determined the two-tailed level of significance. To correct for multiple comparisons, we pooled the obtained timeseries with the timecourse of group-level affective dimensions (panel D). The assessment of significance was performed using a non-parametric permutation test, and false discovery rate correction (Benjamini and Yekutieli, 2001) for multiple comparisons was then applied (panel E).

Finally, to test the existence of a chronotopic organization of affect associated with connectivity dynamics, we exploited the multiple timescales derived from the optimal window estimate. In brief, we first estimated the significance of the associations between connectivity and affective dimensions at all explored timescales, as described above. A logical or across timescales was then computed for each affective dimension and a winner-takes-it-all criterion on the log-transformed \( P \)-value maps established the preferred timescale of each brain region (Figure 4).

### Experiment 2: behavioral study

Twenty-one Italian native speakers (13F; mean age 29.4 years, range 23–44) provided ratings of their emotional experience while watching, for the first time, an edited version of the live-action movie 101 Dalmatians (Herek, Walt Disney Pictures, 1996). All participants were clinically healthy and had no history of any neurological or psychiatric condition. Each subject was informed about the nature of the research and gave written informed consent to participate in the study. The local Ethical Review Board approved the experimental protocol and procedures (CEAVNO: Comitato Etico Area Vasta Nord Ovest; protocol no. 1485/2017), and the research was conducted in accordance with the Declaration of Helsinki.

Differently from experiment 1, in the validation dataset, participants were asked to report moment-by-moment (10Hz sampling rate) the pleasantness or unpleasantness and intensity of the experience on a continuous scale ranging from −100 (extremely negative) to +100 (extremely positive). For each subject, raw ratings were interpreted as a measure of polarity, whereas the absolute values were considered a measure of intensity. For stimulus preparation and task implementation, please see Supplementary Data.

### Experiment 2: fMRI study

Brain activity elicited by the watching of 101 Dalmatians was obtained from 10 healthy Italian subjects (8F; 35 ± 13 years), instructed to simply inhibit any movement and enjoy the movie. Individuals participating in the MRI study did not take part in the
Fig. 2. (Panel A) Results of the connectivity strength of brain regions associated with changes in the polarity of the affective experience at all explored timescales. Grading of the color red indicates the number of window widths in which the connectivity strength of a region was associated with polarity. In the MDS plot, located in the lower part of the figure, each brain area (i.e. dot) is color-coded depending on network membership, whereas its size and transparency are scaled according to the $-\log(P$-value) of the relationship with polarity. In panel B, the connectivity strength of brain regions associated with changes in the intensity of the affective experience at all explored timescales is reported. Grading of the color blue indicates the number of window widths in which the connectivity strength of a region was associated with intensity. In the MDS plot, each brain area (i.e. dot) is color-coded depending on network membership, whereas its size and transparency are scaled according to the $-\log(P$-value) of the relationship with intensity. LH = left hemisphere; RH = right hemisphere; CS = central sulcus; arCING = ascending ramus of the right cingulate sulcus; Amyg = amygdala; Hipp = hippocampus; precUN = precuneus; TPJ = temporoparietal junction; INS = insula; dPFC = dorsal prefrontal cortex; vPFC = ventral prefrontal cortex; mCING = mid-cingulate cortex; FusG = fusiform gyrus; MFG = middle frontal gyrus; dmPFC = dorsomedial prefrontal cortex; MOG = middle occipital gyrus; PFC = prefrontal cortex; preCG = precentral gyrus; postCG = postcentral gyrus; postCS = postcentral sulcus; dpreCS = dorsal precentral sulcus; vpreCS = ventral precentral sulcus; SPL = superior parietal lobule; SMG = supramarginal gyrus; OFC = orbitofrontal cortex; DorsAttn = dorsal attention network; VentAttn = ventral attention network; SomMot = somatomotor network; SubCort = subcortical network.

Fig. 3. Results of the validation dataset. Panel A depicts the set of regions whose connectivity is associated with either polarity or intensity at all explored timescales in the exploratory dataset (i.e. conjunction analysis, see Formula A). The Manhattan plot in panels B and C summarizes the significance of the association between tISFC and affective dimensions in the validation dataset (i.e. $-\log(P$)). Connectivity of the R-TPJ, a relevant node of the DMN, is associated with changes in polarity and intensity, irrespectively of the fMRI pre-processing pipeline ($P_{FDR} < 0.05$ for workflows A and B). Also, the connectivity of the right dorsal prefrontal cortex (control network) is associated with changes in polarity ($P_{FDR} < 0.05$ for workflows A and B). Panels D and E show whole-brain results. Green outlines delineate significant regions in the exploratory dataset; red (polarity; panel D) and blue fill (intensity; panel E) highlight regions associated with affect in the validation dataset after applying whole-brain FDR correction. Findings obtained from the whole-brain approach confirm ROI-based ones. LH = left hemisphere; RH = right hemisphere; Amyg = amygdala; precUN = precuneus; TPJ = temporoparietal junction; INS = insula; dPFC = dorsal prefrontal cortex; vPFC = ventral prefrontal cortex; SPL = superior parietal lobule; SMG = supramarginal gyrus; OFC = orbitofrontal cortex; DorsAttn = dorsal attention network; VentAttn = ventral attention network; SomMot = somatomotor network; SubCort = subcortical network.
behavioral experiment. Also, they watched the movie 101 Dalmatians for the first time inside the scanner. The fMRI acquisition lasted 1 h and was split into six runs (3T Philips Ingenia scanner; gradient recall echo echo-planar imaging (GRE-EPI): repetition time (TR) = 2000 ms, echo time (TE) = 30 ms, flip angle (FA) = 75°, 3 mm isotropic voxel). The Ethical Committee of the University of Turin (protocol no. 195874/2019) approved the fMRI study.

Considering the high variability in results related to analytical choices (Botvinik-Nezer et al., 2020), we used two alternative pre-processing pipelines to analyze the validation dataset (i.e. workflows A and B), which are detailed in Supplementary Data.

To evaluate the reliability and generalizability of the association between tISFC and the timescourse of affective dimensions reported in experiment 1, we tested whether brain connectivity dynamics explains changes in polarity and intensity in 101 Dalmatians. To this aim, we followed the procedure detailed in the ‘Experiment 1—fMRI study’ section (see also Supplementary Data) and used a conjunction analysis of results obtained from experiment 1 to restrict the search for significance. Specifically, we selected brain areas whose connectivity significantly tracked changes of one or more affective dimensions at all explored timescales. This conjunction analysis can be summarized as follows:

Formula A.

\[(P_{w_1} \cap P_{w_2} \cap P_{w_3} \cap P_{w_4}) \cup (C_{w_1} \cap C_{w_2} \cap C_{w_3} \cap C_{w_4}) \cup (I_{w_1} \cap I_{w_2} \cap I_{w_3} \cap I_{w_4})\]

where \(P_{w_i}\), \(C_{w_i}\) and \(I_{w_i}\), represent regions significantly encoding polarity, complexity and intensity at the \(i\)-th timescale.

This entire procedure was repeated for workflows A and B. To control for false positives, we pooled together \(P\)-values obtained from both workflows and all affective dimensions and applied the false discovery rate algorithm under dependency assumptions (Benjamini and Yekutieli, 2001; \(P\)-value < 0.05).

Results

Experiment 1: single-subject affective dimensions obtained from behavioral ratings

Affective states elicited by watching Forrest Gump are described by three orthogonal affective dimensions (Lettieri et al., 2019): (i) polarity defines whether the current experience is perceived as pleasant (negative scores) or unpleasant (positive scores); (ii) complexity represents cognitively mediated states (positive scores) and instinctual responses (negative scores); and (iii) intensity denotes the emotional impact of the experience (monotonic increase in positive scores; for further details, see Supplementary Data).

The results of the estimate of the optimal window width show that a trade-off between the across-participant agreement and window size is reached at 273 (9 m:6 s), 239 (7 m:58 s), 143 (4 m:46 s) and 89 timepoints (2 m:58 s) for polarity; also, the optimal window size is 205 (6 m:50 s), 143 (4 m:46 s), 109 (3 m:38 s) and 88 timepoints (2 m:56 s) for complexity and 331 (11 m:2 s), 247 (8 m:14 s), 163 (5 m:26 s) and 100 timepoints (3 m:20 s) for intensity.
**Experiment 1: association between tISFC and polarity of the emotional experience**

The connectivity strength of three regions is significantly associated with the changes in the polarity of the affective experience at all explored timescales \( P_{FDR} < 0.05 \); **Figure 2A; Supplementary Table S1**). These three regions are the right dorsal prefrontal cortex (R-dPFC) pertaining to the control network, the right insula (R-INS), a node of the ventral attention network and the left precuneus (L-preCUN), a node of the DMN. Additionally, the connectivity of three other DMN regions, the right temporoparietal junction (R-TPJ), the right precuneus (R-preCUN) and the right dorsomedial prefrontal cortex (R-dmPFC), encodes polarity at three out of four timescales. Other brain areas, such as the right middle frontal gyrus (R-MFG), the right orbitofrontal cortex (R-OFC), the right postcentral gyrus (R-postCG), the left mid-cingulate cortex (L-mCING), the left middle occipital gyrus, the left insula and the left central sulcus, are significantly associated with changes in polarity at two or only one time window. Except for L-MOG, all other significant regions are negatively associated with polarity scores, meaning that the more unpleasant the experience is (i.e. positive scores), the weaker the connectivity of these nodes with the rest of the brain (Supplementary Figure S1A).

Overall, the connectivity strength of DMN and control network nodes plays a major role in representing the pleasantness of the emotional experience during naturalistic stimulation (**Figure 2A, Supplementary Table S1**).

**Experiment 1: association between tISFC and intensity of the emotional experience**

Findings demonstrate that connectivity strength of four regions is associated with the perceived intensity of the emotional experience \( P_{FDR} < 0.05 \); **Figure 2B; Supplementary Table S2**). These four parcels are the right ventral prefrontal cortex (R-vPFC), a node of the limbic network, the R-TPJ—pertaining to the DMN, the right superior frontal gyrus (R-SFG), belonging to the control network, and the left amygdala (L-Amyg). Intensity is also related to the connectivity of the right precentral gyrus, of the posterior portion of the right supramarginal gyrus (R-SMG) and of the right superior parietal lobule (R-SPL), at three out of four time windows. These brain regions pertain to the control, dorsal attention and somatomotor networks. Other areas, such as the R-OFC, the right dorsal and ventral precentral sulcus (R-dpreCS and R-vpreCS, respectively), the R-INS, the right medial superior frontal gyrus, the ascending ramus of the right cingulate sulcus (R-arCING), the L-preCG, the left postcentral sulcus (L-postCS), the left fusiform gyrus (L-FusG) and the left hippocampus (L-Hipp) encode intensity at two or only one timescale. Except for the bilateral precentral gyrus, the L-FusG, the L-Amyg and L-Hipp, all other significant regions are positively associated with intensity values, meaning that the more intense the experience, the stronger the connectivity of these nodes with the rest of the brain (Supplementary Figure S2A). Altogether, the connectivity strength of control and dorsal attention networks plays a major role in tracking the intensity of the emotional experience (**Figure 2B, Supplementary Table S2**).

**Experiment 1: chronotopic maps of affective dimensions**

Here, we show the preferred timescale at which the connectivity of significant regions relates to changes in polarity, complexity, and intensity. This analysis reveals the existence of chronotopic maps (**Protopapa et al., 2019**), where adjacent cortical regions to complexity at three out of four timescales. Two parcels within the right inferior temporal gyrus (R-ITG), pertaining to the control and the dorsal attention networks, the right intraparietal sulcus (R-IPS), also included in the control network, and the left dorsal prefrontal cortex (L-dPFC), a node of the DMN, track changes in complexity at one timescale (Supplementary Figure S4). Concerning the interpretation of these relationships, R-ITG and R-IPS increase the strength of their connectivity with the rest of the brain during cognitively mediated states (e.g. mixed emotions; Lettieri et al., 2019), whereas R-mCING and L-dPFC are more strongly connected during movie scenes eliciting more instinctual responses (e.g. fear).

**Experiment 2: validation of the relationship between tISFC and affective dimensions**

To test the reliability and generalizability of results obtained from **Forrest Gump** (i.e. exploratory dataset), we collected and analyzed emotion dimension ratings of the live-action movie **101 Dalmatians** (i.e. validation dataset, for further details, please see Supplementary Data). We measured the association between connectivity strength and affective dimensions after restricting the search for significance to regions encoding at least one dimension in the exploratory dataset at all explored timescales. This set of brain areas includes the R-TPJ, the R-SFG, the R-INS, the R-dPFC, the R-vPFC, the L-preCUN and the L-Amyg (**Figure 3A**). As no brain region encodes complexity at all explored timescales, none of the selected regions is obtained from the analysis of this dimension. Therefore, validation analyses are limited to polarity and intensity. Moreover, the selected areas pertain to the default, the control, the limbic, the ventral attention, and the subcortical networks.

The results show that, in the validation dataset, both polarity and intensity ratings are associated with the connectivity strength of R-TPJ, a DMN node \( P_{FDR} < 0.05 \); **Figure 3B–E**. Importantly, this finding does not depend on the fMRI data pre-processing pipeline, and the relationship between affect and brain connectivity is interpreted as in the exploratory dataset: the more pleasant and intense the experience, the stronger the connectivity between this DMN area and the rest of the brain (Supplementary Figures S6 and S7).

In addition, the connectivity strength of R-dPFC, a control network region, is significantly associated with changes in polarity in the validation dataset \( P_{FDR} < 0.05 \) for both workflows A and B; **Figure 3B and D**. Similarly to R-TPJ, the interpretation of the relationship between R-dPFC connectivity and polarity is identical across the two experiments: the more pleasant the experience is, the more R-dPFC is connected to the rest of the brain (Supplementary Figures S6 and S7).

In summary, the results of the validation dataset demonstrate that two main findings of the exploratory dataset generalize to another movie, sample, different fMRI pre-processing workflows and setup for the collection of behavioral data. Specifically, in both datasets we found a significant relationship between the polarity and connectivity of R-dPFC (control network), as well as an association between R-TPJ (DMN) and both polarity and intensity.
preferentially map the stream of affect at different time intervals (Figure 4).

For the polarity dimension, chronotopic organizations are revealed in the right temporoparietal and the right prefrontal cortex (Figure 4A). Within TPJ, the connectivity of dorsocaudal territories is associated with polarity scores at shorter timescales (~3 min), while ventrorostral portions preferentially track polarity in ~8 min segments. In the right prefrontal cortex, connectivity is mainly associated with the scores of the polarity component at shorter timescales (~3 min) in its central region, and at intermediate intervals (~5 min) in its ventral and dorsal parts. In addition, the connectivity of the R-dmPFC preferentially represents polarity in 5 min segments, whereas L-MOG shows a marked preference for longer timescales (from ~8 to ~9 min). The connectivity of all other brain areas is maximally associated with this affective dimension at shorter intervals (~3 min).

For the intensity component, we show timescale topographies within the right parietal lobe, the right frontopolar cortex, the right cingulate region and the R-SFG (Figure 4B). In the right parietal lobe, the territories of the R-SPL preferentially map the intensity of the emotional experience in longer segments (from ~8 to ~11 min), whereas regions pertaining to the inferior parietal lobule in shorter time windows (from ~3 to ~5 min). The chronotopic arrangement in the right frontopolar region indicates that the connectivity of its lateral portion preferentially encodes intensity in ~5 min segments, whereas its medial part in ~3 min intervals. Also, within the right cingulate region, we observe an abrupt change in timescale preference, with connectivity of the inferior portion of the R-arcCING preferentially mapping emotional intensity at the longest timescale (~11 min) and its superior region at the shortest one (~3 min). For the R-SFG, the medial part optimally represents intensity in ~3 min segments and the lateral portion in ~8 min ones. Interestingly, the connectivity of L-Hipp and L-Amyg shows opposite behaviors: the hippocampus encodes emotional intensity at the longest timescale (~11 min), whereas the amygdala at the shortest one (~3 min). All other brain regions, besides the R-dprC5S, the L-postC5S and the L-FusC5, are tuned to shorter time windows.

Finally, no region significantly associated with the complexity dimension shows a chronotopic organization (Supplementary Figure S5).

**Discussion**

In the current study, we combined the real-time reports of the affective experience with recordings of fMRI activity during naturalistic stimulation to reveal how brain connectivity encodes the stream of affect. We collected approximately 45 h of behavioral reports and 38 h of brain imaging data over two experiments. In experiment 1, ratings of emotion categories associated with the watching of Forrest Gump were used to obtain time series of affective dimensions and to explain brain connectivity dynamics in independent individuals. In experiment 2, the 101 Dalmatians movie was used to test the reliability and generalizability of results obtained from the exploratory dataset: moment-by-moment scores of the polarity and intensity of the emotional experience were correlated with changes in the connectivity of brain regions identified in experiment 1. Also, fMRI data were pre-processed according to two alternative workflows. Therefore, our study protocol ensures that findings cannot be merely ascribed to (i) the selection of a specific movie, (ii) participants’ characteristics, (iii) the method used to collect behavioral reports and (iv) the pipeline adopted to pre-process fMRI data.

The results show that the connectivity strength of the R-TPJ and of the R-dPFC, respectively, DMN and control network nodes, is positively associated with the polarity of the affective experience. Moreover, affective dimensions converge in the right temporoparietal cortex, as this region’s connectivity tracks changes in emotional intensity as well. Together with the right prefrontal cortex, the R-TPJ represents the dynamics of affect in a chronotopic manner, meaning that adjacent portions of these areas preferentially map the stream of affect at different time intervals, ranging from few to several minutes.

In everyday life, affect varies as a function of time, and this complex relationship is hardly recreated in the laboratory setting. To elicit a variety of affective states, track their evolution in time and maintain a relatively high ecological validity, an optimal solution may be represented by movie watching. Movies produce strong affective responses over a few hours and mimic everyday life situations with an alternation of sudden and predictable events (Schaefe et al., 2010). Also, changes in affect induced by movies are based on rich contextual information and are related to the viewer’s expectations. Building upon this, we recorded brain activity and affective ratings during the watching of two movies and demonstrated that changes in the subjective emotional experience explain brain connectivity dynamics in independent participants.

Over the last decade, researchers have emphasized the importance of moving from a region- to a network-based approach when studying the neural correlates of emotion (Lindquist and Barrett, 2012; Raz et al., 2016; Kragel et al., 2016; Pessoa, 2017). Changes in affect are associated with modifications in bodily responses (e.g. autonomic and musculoskeletal) and mental processes (e.g. memory and decision making), which are likely reflected in a distributed functional reorganization of brain modules (Pessoa, 2017). In this regard, the DMN is thought to play a crucial role in interoception (Kleckner et al., 2017) and in constructing and representing emotions (Satpute and Lindquist, 2019). The connectivity of this network is also altered in affective disorders, such as anxiety (Liao et al., 2010) or depression (Whitfield-Gabrieli and Ford, 2012). Our findings support the central role of this network in the processing of affect, as we demonstrate that its connectivity tracks changes in the subjective judgment of emotional polarity and intensity in a naturalistic environment.

Furthermore, we show across two experiments that the connectivity of a prefrontal control network node is also related to the polarity of affect. The control network is involved in empathy (Christov-Moore et al., 2020) and seems crucial in selecting sensations for conscious awareness during the construction of a mental state (Lindquist and Barrett, 2012). In our study, participants of the behavioral experiment were asked to constantly report their conscious emotional experience, indeed, a task that involves the processing and selection of sensations. It is interesting to note that, although fMRI participants were not explicitly asked to focus on their emotions, watching an evocative movie modulates the connectivity of the control network, and such modulation is associated with conscious reports of affect.

When we look closely at nodes encoding affective dimensions, we notice that connectivity between the R-TPJ and the rest of the brain represents both polarity and intensity. This cortical area is one of the core regions of the social brain (Saxe and Kanwisher, 2019).
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Supplementary data

Supplementary data are available at SCAN online.

Conflict of interest

The authors declare no conflict of interest.

Author contributions

G.L., G.H., P.P. and L.C., conceived the study. G.L., G.H. and L.C. designed the behavioral experiment, collected behavioral data, developed the code, performed behavioral and fMRI data analysis, interpreted the obtained results and drafted the manuscript. E.M.C. acquired behavioral data of the validation dataset. F.S., V.B., M.D. and A.L. acquired fMRI data of the validation dataset. G.L., G.H., L.C., F.S. and A.L. analyzed fMRI data of the validation dataset. G.L., G.H., E.M.C., F.S., V.B., M.D., A.L., E.R., P.P. and L.C. critically revised the manuscript. All the authors approved the final version of the manuscript.

References

Antony, J.W., Hartshorne, T.H., Pomeroy, K., et al. (2021). Behavioral, physiological, and neural signatures of surprise during naturalistic sports viewing. Neuron, 109(2), 377–90.

Baldassano, C., Chen, J., Zadbood, A., Pillow, J.W., Hasson, U., Norman, K.A. (2017). Discovering event structure in continuous narrative perception and memory. Neuron, 95(3), 709–21.

Baucom, L.B., Edell, D.H., Wang, J., Blitzer, D.N., Shinkareva, S.V. (2012). Decoding the neural representation of affective states. NeuroImage, 59(1), 718–27.

Benjamini, Y., Yekutieli, D. (2001). The control of the false discovery rate in multiple testing under dependency. Annals of Statistics, 29(4), 1165–88.

Botvinik-Nezer, R., Holzmeister, F., Camerer, C.F., et al. (2020). Variability in the analysis of a single neuroimaging dataset by many teams. Nature, 582(7810), 84–8.

Chen, J., Leong, Y.C., Honey, C.J., Yong, C.H., Norman, K.A., Hasson, U. (2017). Shared memories reveal shared structure in neural activity across individuals. Nature Neuroscience, 20(1), 115–25.

Christov-Moore, L., Beggente, N., Douglas, P.K., Feusner, J.D., Iacoboni, M. (2020). Predicting empathy from resting state brain connectivity: a multivariate approach. Frontiers in Integrative Neuroscience, 14, 3.

Costa, T., Cauda, F., Crini, M., et al. (2014). Temporal and spatial neural dynamics in the perception of basic emotions from complex scenes. Social Cognitive and Affective Neuroscience, 9(11), 1690–703.

Ekman, P. (1992). An argument for basic emotions. Cognition and Emotion, 6(3–4), 169–200.

Etkin, A., Büchel, C., Gross, J.J. (2015). The neural bases of emotion regulation. Nature Reviews Neuroscience, 16(11), 693–700.

Hanke, M., Adelhöfer, N., Kottke, D., et al. (2016). A studyforest extension, simultaneous fMRI and eye gaze recordings during prolonged natural stimulation. Scientific Data, 3, 160092.
Hasson, U., Chen, J., Honey, C.J. (2015). Hierarchical process memory: memory as an integral component of information processing. Trends in Cognitive Sciences, 19(6), 304–13.

Hasson, U., Nir, Y., Levy, I., Fuhrmann, G., Malach, R. (2004). Inter-subject synchronization of cortical activity during natural vision. Science, 303(5664), 1634–40.

Kim, J., Shinkareva, S.V., Wedell, D.H. (2017). Representations of modality-general valence for videos and music derived from fMRI data. NeuroImage, 148, 42–54.

Kleckner, I.R., Zhang, J., Touroutoglou, A., et al. (2017). Evidence for a large-scale brain system supporting allostatics and interoception in humans. Nature Human Behaviour, 1(5), 1–14.

Kleiner, M., et al. (2011). What’s new in psychtoolbox-3. Perception, 36(1).

Kohn, N., Eickhoff, S.B., Scheller, M., Laird, A.R., Fox, P.T., Habel, M., et al. (2016). Spatially localized connectivity of resting state networks in social anxiety disorder. NeuroImage, 148, 345–55.

Kragel, P.A., Knodt, A.R., Hariri, A.R., LaBar, K.S. (2016). Decoding spontaneous emotional states in the human brain. PLoS Biology, 14(9), e2000106.

Kuppens, P., Oravecz, Z., Tuerlinckx, F. (2010). Feelings change: accounting for individual differences in the temporal dynamics of affect. Journal of Personality and Social Psychology, 99(6), 1042.

Kuppens, P., Verdun, P. (2017). Emotion dynamics. Current Opinion in Psychology, 17, 22–6.

Lamm, C., Nusbaum, H.C., Meltzoff, A.N., Decety, J. (2007). What are you feeling? Using functional magnetic resonance imaging to assess the modulation of sensory and affective responses during empathy for pain. PloS One, 2(12), e1292.

Lettieri, G., Handjarias, G., Ricciardi, E., et al. (2019). Emotionotopy in the human right temporo-parietal cortex. Nature Communications, 10(1), 1–13.

Liao, W., Chen, H., Feng, Y., et al. (2010). Selective aberrant functional connectivity of resting state networks in social anxiety disorder. NeuroImage, 52(4), 1549–58.

Lindquist, K.A., Barrett, L.F. (2012). A functional architecture of the human brain: emerging insights from the science of emotion. Trends in Cognitive Sciences, 16(11), 533–40.

Makris, N., Goldstein, J.M., Kennedy, D., et al. (2006). Decreased volume of left and total anterior insular lobe in schizophrenia. Schizophrenia Research, 83(2–3), 155–71.

Nastase, S.A., Gazzola, V., Hasson, U., Keysers, C. (2019). Measuring shared responses across subjects using intersubject correlation. Social Cognitive and Affective Neuroscience, 14(6), 667–85.

Pessoa, L. (2017). A network model of the emotional brain. Trends in Cognitive Sciences, 21(5), 357–71.

Pessoa, L. (2018). Understanding emotion with brain networks. Current Opinion in Behavioral Sciences, 19, 19–25.

Posner, J., Russell, J.A., Gerber, A., et al. (2009). The neurophysiological bases of emotion: an fMRI study of the affective circuitry using emotion-denoting words. Human Brain Mapping, 30(3), 883–95.

Protopapa, F., Hayashi, M.J., Kulasekhar, S., et al. (2019). Chronotopic maps in human supplementary motor area. PLoS Biology, 17, 3.

Raz, G., Touroutoglou, A., Wilson-Mendenhall, C., et al. (2016). Functional connectivity dynamics during film viewing reveal common networks for different emotional experiences. Cognitive, Affective and Behavioral Neuroscience, 16(4), 709–23.

Réspois, M., Verdyn, P., Delaveau, P., et al. (2017). The neural basis of emotions varies over time: different regions go with onset- and offset-bound processes underlying emotion intensity. Social Cognitive and Affective Neuroscience, 12(8), 1261–71.

Réspois, M., Rotgé, J.Y., Delaveau, P., et al. (2018). The impact of self-distancing on emotion explosiveness and accumulation: an fMRI study. PLoS One, 13(11), e0206889.

Satpute, A.B., Lindquist, K.A. (2019). The default mode network’s role in discrete emotion. Trends in Cognitive Sciences, 23(10), 851–64.

Saxe, R., Kanwisher, N. (2003). People thinking about thinking people: the role of the temporo-parietal junction in ‘theory of mind’. NeuroImage, 19(4), 1835–42.

Schaefer, A., Nils, F., Sanchez, X., Philippot, P. (2010). Assessing the effectiveness of a large database of emotion-eliciting films: a new tool for emotion researchers. Cognition and Emotion, 24(7), 1153–72.

Schaefer, A., Kong, R., Gordon, E.M., et al. (2018). Local-global parcellation of the human cerebral cortex from intrinsic functional connectivity MRI. Cerebral Cortex, 28(9), 3095–114.

Shen, X., Tokoglu, F., Papademetris, X., Constable, R.T. (2013). Group-wise whole-brain parcellation from resting-state fMRI data for network node identification. NeuroImage, 82, 403–15.

Simony, E., Honey, C.J., Chen, J., et al. (2016). Dynamic reconfiguration of the default mode network during narrative comprehension. Nature Communications, 7, 12141.

Thornton, M.A., Tamir, D.I. (2017). Mental models accurately predict emotion transitions. PNAS, 114(23), 5982–7.

Torgerson, W.S. (1952). Multidimensional scaling: I. Theory and method. Psychometrika, 17(4), 401–19.

Trampe, D., Quoidbach, J., Taquet, M. (2015). Emotions in everyday life. PLoS One, 10(12), e0145450.

Verdun, P., Delaveau, P., Rotgé, J.Y., Fossati, P., Van Mechelen, I. (2015). Determinants of emotion duration and underlying psychological and neural mechanisms. Emotion Review: Journal of the International Society for Research on Emotion, 7(4), 330–5.

Waugh, C.E., Schirillo, J.A. (2012). Timing: a missing key ingredient in typical fMRI studies of emotion. Behavioral and Brain Sciences, 35(3), 170–1.

Whitfield-Gabrieli, S., Ford, J.M. (2012). Default mode network activity and connectivity in psychopathology. Annual Review of Clinical Psychology, 8, 49–76.

Yeo, B.T., Krienen, F.M., Sepulcre, J., et al. (2011). The organization of the human cerebral cortex estimated by intrinsic functional connectivity. Journal of Neurophysiology, 106(3), 1125–65.

Yeshurun, Y., Nguyen, M., Hasson, U. (2021). The default mode network: where the idiosyncratic self meets the shared social world. Nature Reviews Neuroscience, 22(3), 181–92.