Spontaneous neural encoding of social network position

Carolyn Parkinson1*, Adam M. Kleinbaum2 and Thalia Wheatley3

Unlike many species that enact social behaviour in loose aggregations (such as swarms or herds), humans form groups comprising many long-term, intense, non-reproductive bonds with non-kin1. The cognitive demands of navigating such groups are thought to have significantly influenced human brain evolution2. Yet little is known about how and to what extent the human brain encodes the structure of the social networks in which it is embedded. We characterized the social network of an academic cohort (N=275); a subset (N=21) completed a functional magnetic resonance imaging (fMRI) study involving viewing individuals who varied in terms of 'degrees of separation' from themselves (social distance), the extent to which they were well-connected to well-connected others (eigenvector centrality) and the extent to which they connected otherwise unconnected individuals (brokerage). Understanding these characteristics of social network position requires tracking direct relationships, bonds between third parties and the broader network topology. Pairing network data with multi-voxel pattern analysis, we show that information about social network position is accurately perceived and spontaneously activated when encountering familiar individuals. These findings elucidate how the human brain encodes the structure of its social world and underscore the importance of integrating an understanding of social networks into the study of social perception.

Relationships are intrinsic to human behaviour. Everyday interactions are shaped not only by our own relationships, but also by knowledge of bonds between third parties and the broader social networks in which we are embedded. Well-connected individuals can effectively threaten or bolster one's reputation3, those who bridge otherwise disparate groups can efficiently seek and spread information4, and knowledge of mutual ties influences information-sharing and trust5. Human social intelligence rests, in part, on a calculus that inheres in an understanding of social network structure.

Is knowledge about others’ social network positions activated only when explicit goals require it, or spontaneously, whenever we encounter familiar individuals? It may be efficient to process such information only when our goals require it (for example when determining how to obtain information, or forecasting the repercussions of a social misstep). Alternatively, it may be beneficial to activate such knowledge spontaneously when encountering others, given the importance of social network position to many aspects of behaviour and to impressions of status and competence6,7. Humans spontaneously register a great deal of information when perceiving other people (such as intentions, traits and emotions)8,9, presumably to aid appropriate, beneficial social interactions. Thus, the brain may run several social ‘daemons’ — efficient, background processes that spontaneously register information useful for predicting the social repercussions of potential actions, and, more broadly, for informing cognition and behaviour.

To test whether the brain spontaneously encodes the social network positions of familiar others, we scanned (using fMRI) members of a real-world social network (see Fig. 1; Methods) as they viewed brief videos of 12 classmates (Fig. 2). The only task was to indicate when the same video was presented twice in a row (see Methods), to ensure attention without any instructions to retrieve social relationship knowledge or person knowledge more generally. Therefore, we consider any information about social network position encoded while participants performed this task to be retrieved spontaneously (that is, without instruction).

Each classmate in each participant’s stimulus set was characterized according to three metrics derived from the social network data: geodesic social distance from the participant; eigenvector centrality; and constraint, an inverse measure of brokerage. Geodesic social distance refers to the minimum number of intermediary social ties required to connect two individuals. Eigenvector centrality is a prestige-based centrality metric that considers not only how many connections a given individual has, but also the centralities characterizing each contact8. High eigenvector centrality (high EC) implies that an individual is well-connected to well-connected others; low EC implies that an individual has few friends and that these friends tend to be unpopular. Prestige-based centrality metrics are particularly useful for characterizing social status, given that being named as a friend by a popular individual should increase one’s sociometric status (that is, the extent to which someone is liked by others) more than being named by someone less popular9. Individuals who connect others who would not otherwise be connected occupy network positions low in constraint, and have the capacity to serve as ‘brokers’ of resources (for example, information) in the network. Because of the structure of their local social ties, brokers can coordinate behaviour and translate information across structural holes in networks4.

To probe for the spontaneous encoding of social network position information, we used representational similarity analysis (RSA), which distills fMRI response patterns into representational dissimilarity matrices (RDMs) that indicate the degree to which particular brain regions distinguish between sets of stimuli or mental states10. Because RDMs are abstracted away from the spatial layout of neuroimaging data (they are indexed by experimental condition; Fig. 3), RSA affords the evaluation of the degree to which similarity structures characterizing the representational content of particular brain regions reflect those characterizing data acquired using other modalities of measurement or computational models11 (here, the social network data). Specifically, in the current study, we used a

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As social distance between people increases, their relevance to each other given the potential reputation costs of bad behaviour. Similarly, sharing mutual friends may enhance trust, with such individuals could damage relationships with one’s direct be relatively important to identify and monitor: negative interactions usu‐ ally most self-relevant. Given the importance of reputation man‐ 
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reliably across participants (about specific characteristics of social network position was carried
of the degree to which the representational content of local neural
positions of the individuals in that participant’s stimulus set (Fig. 3).
Neural RDMs were iteratively extracted within 9-mm-radius spheres centred at each point in each participant’s brain. Within
each participant, each local neural RDM was modelled as a weighted combination of RDMs based on properties of the social network positions of the individuals in that participant’s stimulus set (Fig. 3).
Using this technique, participants’ brains were mapped in terms of the degree to which the representational content of local neural responses to familiar others could be explained by those individuals’ positions in their social network, and in terms of where information about specific characteristics of social network position was carried reliably across participants (Fig. 4; Supplementary Fig. 2).
We hypothesized that geodesic social distance would be spontaneously encoded, given the importance of this information for determining self-relevance. One’s immediate social ties are obviously most self-relevant. Given the importance of reputation man‐ 
agement for human behaviour, individuals two ‘degrees away’ may be relatively important to identify and monitor: negative interactions with such individuals could damage relationships with one’s direct connections. Similarly, sharing mutual friends may enhance trust, given the potential reputation costs of bad behaviour. As social distance between people increases, their relevance to each other decreases. We predicted that social-distance-related information would be carried in the lateral superior temporal cortex (STC) and inferior parietal lobule (IPL), as well as the medial prefrontal cortex (MPFC), given past research implicating these regions in encoding social distance and self-relevance more generally.

Social distance was reliably signalled in a large cluster centred in the lateral posterior STC and extending inferiorly throughout the posterior lateral temporal cortex (LTC), and superiorly to the anterior aspect of the IPL (see Fig. 4; Supplementary Table 1). Past research demonstrated that multi-voxel response patterns in this region encode egocentric spatial and abstract (for example, social) distances when explicitly judging or mentally navigating such distances; the current findings suggest that this region also encodes egocentric distances spontaneously (that is, in the absence of any explicit distance task). Thus, when encountering a familiar individual, knowledge of agent-to-agent relationships seems to be spontaneously retrieved, such that representations of other people in this region are organized in terms of whether someone is a friend, a friend-of-a-friend, or farther removed from oneself in social ties. It has been suggested that some regions within the posterior parietal cortex, such as the anterior IPL, which have well-established roles in representing and navigating physical space, analogously represent more abstract relationships (such as social ties between agents). The current results indicate that when encountering familiar individuals, humans may spontaneously retrieve knowledge of where they are located, relative to oneself, in a mental map of ‘social space’.

Although the LTC and IPL regions that carried information about social distance here have previously been implicated in encoding social distance, some regions previously implicated in signalling social distance were not implicated in the current study. For instance, previous research has implicated MPFC in distinguishing friends from strangers, and a recent study implicated the hippocampus and posterior cingulate cortex (PCC) in tracking social distances between participants and characters in an interactive game. Differences between the current results and those observed in previous investigations
Eigenvector centrality and constraint would be encoded in brain regions encoding individual identity when imagining others’ actions (for example, the temporal pole and fusiform gyrus; see Fig. 4).

Whereas social distance is inherently relative to the perceiver, other aspects of familiar others’ social network positions, such as the degree to which each person ‘bridges’ different areas of the network and the number of friends someone has, are increasingly thought to be largely stable, possibly heritable, dispositional tendencies that shape social behaviour. Therefore, we hypothesized that eigenvector centrality and constraint would be encoded in brain regions involved in encoding others’ traits and behavioural tendencies more generally, such as the MPFC, which is widely implicated in inferring and encoding person knowledge and in integrating knowledge of personality traits in order to signal individual identity.

Information about eigenvector centrality was reliably carried in brain regions that encode individual identity when imagining others’ actions (for example, MPFC) and viewing faces (for example, the temporal pole and fusiform gyrus; see Fig. 4 and Supplementary Table 2), suggesting that sociometric status may constitute a dimension of meaning for organizing mental representations of others. Eigenvector centrality was also encoded in medial parietal cortex (precuneus, PCC), a region previously shown to encode extraversion, which is modestly correlated with eigenvector centrality, suggesting that this region may encode dispositional tendencies common to both extraversion and eigenvector centrality.

In addition, recent work has shown that the medial parietal cortex, as well as other regions involved in inferring others’ mental states, intentions and traits (for example, MPFC; temporoparietal junction), spontaneously responds to well-liked individuals in a real-world social network, which is thought to reflect that perceivers are preferentially motivated to understand the internal states of popular others. The current findings are consistent with the notion that brain regions that represent others’ internal states and behavioural tendencies (such as PCC and MPFC) track sociometric status, and suggest that like other facets of social status (for instance, dominance or prestige), eigenvector centrality may modulate attention to the internal states of others. Future behavioural studies should directly test the impact of eigenvector centrality on social attention.

Information about eigenvector centrality was also reliably carried in unexpected regions, such as extrastriate visual cortex (EVC). This result is unlikely to be due to low-level visual characteristics of stimuli, as each participant had a unique stimulus set, and because videos corresponding to each individual in each stimulus set were horizontally mirrored on half of the trials (see Methods). This finding...
may nonetheless reflect the effects of social status in terms of social ties on visual attention. People tend to orient preferentially toward high-status individuals and to the loci of their attention, presumably to obtain behaviourally relevant information about our surroundings\(^{29-31}\). Given that eigenvector centrality is reliably signalled in EVC response patterns, future research should test whether visual attention is also preferentially allocated to central actors in one’s social network.

Eigenvector-centrality–based RDMs were also significantly related to neural RDMs in brain areas previously implicated in evaluating social status in terms of dominance, prestige and morality, such as the ventral MPFC and ventrolateral prefrontal cortex (VLPFC)\(^{32-34}\). The involvement of the ventral MPFC in social status encoding has been suggested to reflect a more general role in assessing the value of stimuli\(^{35}\), whereas the VLPFC has been suggested to encode social status in order to appropriately modulate behavioural responding, which is thought to be a primary function of status cues\(^{3}\). We suggest that these regions probably encode eigenvector centrality for similar reasons, as high-EC individuals have high behavioural relevance and value as social partners. For example, individuals connected to well-connected others may be protected from mistreatment because they are more likely to be defended by others, who themselves are more likely to be defended. Less risk is associated with wronging a low-EC individual, given that low-EC individuals have little influence on the spread of information and other resources\(^{3}\).

The current results suggest that when encountering a familiar individual, the degree to which that individual is well-connected to well-connected others shapes processes related to valuation, behavioural modulation, attention, and encoding others’ internal states, dispositional characteristics and identities. Many of these findings echo the known effects of other dimensions of social status (for example status conferred by dominance). Although a great deal of past psychological and neuroimaging research on social status has focused on physical dominance, we note that overt physical violence is relatively rare in contemporary human groups\(^{36}\) and that social support and reputation management are central to everyday human life\(^{37,38}\). Social power in such groups may be relatively less contingent on individual strength and physical aggression, and more dependent on group dynamics and affiliative relationship maintenance. Thus, sociometric status is likely to be especially relevant to modern humans, and merits further attention in social perception and neuroscience research.

In addition to social distance and eigenvector centrality, diverse aspects of social cognition and behaviour (for example, deciding how to effectively seek or spread information; trust decisions) would benefit from encoding network constraint. Low-constraint individuals can broker the flow of information between groups, and thus exert a disproportionate influence on the flow of ideas and resources\(^{4}\). Additionally, individuals in relatively closed local networks, characterized by high constraint, suffer greater reputation costs for bad behaviour; correspondingly, constraint can foster trust and cooperation\(^{5}\). Given the dearth of previous research investigating the perception of constraint, we made no specific predictions about which brain regions would be involved in encoding this facet of social network position.

Large clusters spanning both right and left lateral STC carried information about constraint (Supplementary Table 3), as did a smaller cluster in the supplementary motor area. Although the lateral STC and supplementary motor area are implicated in biological motion processing\(^{39}\) and action understanding\(^{40}\), respectively, this finding was not attributable to the amount of movement in videos (see Supplementary Information). A perceiver’s knowledge of the network constraint of an individual, or of associated dispositions, may affect how that perceiver attends to that individual’s movements. Because brokers may be perceived as exceptionally charismatic or interesting (for example, because they often serve as sources of new information or opportunities\(^{5}\)), they may command differential amounts of attention to their expressions and gestures. Brokers may also differ in the amount of social meaning carried in their facial and bodily movements (for example, using movement to express oneself coherently versus fidgeting aimlessly). The latter explanation would be consistent with evidence that the STC responds to the social meaning, rather than amount, of movement in dynamic displays\(^{41}\). Future studies could arbitrate between these hypotheses by testing whether strangers are able to differentiate between individuals high and low in constraint based on their observed movements. If so, this would suggest that network constraint is encoded in lateral STC because this aspect of social network position is apparent in how individuals carry themselves. If not, this would be consistent with the interpretation that perceivers’ knowledge of an individual’s network constraint, or of qualities related to this aspect of social network position, influences how perceivers attend to that individual’s expressions, gestures and bodily movements.

After scanning, participants were asked about their perceptions of each social network analysis–derived metric of interest for each individual in their stimulus set (see Supplementary Information). This allowed us to test the accuracy of participants’ perceptions of others’ social network positions, and to evaluate how well participants’ perceptions matched the data used to construct their stimulus sets. Post-scan ratings indicated that participants’ explicit perceptions of the social network positions of the individuals in
their stimulus sets closely matched reality. Veridical constraint had a significant effect on perceived constraint ($β = 19.44, \text{standard error} (SE) = 2.01, p < 0.0001$), and veridical eigenvector centrality had a significant effect on perceived eigenvector centrality ($β = 14.95, \text{SE} = 0.95, p < 0.0001$). Further, subjective ratings of social closeness ($β = -31.00, \text{SE} = 1.62, p < 0.0001$), proportion of social time spent together ($β = -22.74, \text{SE} = 1.84, p < 0.0001$), and frequency of discussions ($β = -33.77, \text{SE} = 1.89, p < 0.0001$) varied as a function of geodesic network distance (see Methods and Supplementary Fig. 1).

Although participants had consciously accessible knowledge of the characteristics of social network position studied here (Supplementary Fig. 1), the task used in the fMRI study (a one-back memory task) did not require participants to retrieve that knowledge. Nevertheless, up to 40% of the variance in similarity structures of local IMRI responses to personally familiar others could be explained merely by characteristics of those individuals’ positions in the perceivers’ social network (Supplementary Fig. 2). These findings are consistent with behavioural evidence that humans spontaneously activate knowledge about other people upon encountering them in order to inform cognition and behaviour13, and suggest that humans spontaneously activate complex knowledge about other people’s positions in their social networks when viewing them. The findings are also consistent with psychologists’ mounting appreciation of the importance of both direct and indirect relationship knowledge to everyday cognition and behaviour. Everyday interactions are influenced not only by information that would be available to any observer, but also by patterns of personal and third-party relationships. By adopting an interdisciplinary approach combining theory and methods from neuroscience, psychology and social network analysis, we can begin to uncover a deeper understanding of how the human brain negotiates the intricacies of everyday social life.

Methods

Part 1: Social network characterization. Participants. Participants in Part 1 of the study were 275 first-year Masters of Business Administration (MBA) students at a private university in the United States who participated as part of their coursework on leadership (91 females; 184 males). The total class size was 277 students; two students failed to complete the questionnaire (response rate 99.3%). All procedures were completed in accordance with the standards of the Dartmouth Committee for the Protection of Human Subjects.

Social network characterization. To characterize the social network of all first-year students, an online social network survey was administered. Participants followed an e-mailed link to the study website where they responded to a survey designed to assess their position in the social network of first-year students in their academic programme. The survey question was adapted from Burt14 and has been previously used in the modified form used here10,11. It read, “Consider the people with whom you like to spend your free time. Since you arrived at [institution name], who are the classmates you have been with most often for informal social activities, such as going out to lunch, dinner, drinks, films, visiting one another’s homes, and so on?”

A roster-based name generator was used to avoid inadequate or biased recall. Classmates’ names were listed in four columns, with one column corresponding to each section of students in the MBA programme. Names were listed alphabetically within section. Participants indicated the presence of a social tie with an individual by placing a checkmark next to his or her name. Participants could indicate any number of social ties and had no time limit for responding. The result is a friendship network based on voluntary social interactions.

Social network analysis was performed using the R package igraph14,15. Three social-network-derived metrics were extracted for each node: constraint, eigenvector centrality and geodesic distance from each classmate, as described in greater detail below.

Constraint. The constraint of actor i is given by the following equation, where $P_i$ corresponds to the proportion of i’s direct social ties accounted for by his/her tie to actor j. The inner summation approximates the indirect constraint imposed by i on j by other actors, q, who are socially connected to both i and j (mutual friends of i and j):

$$\text{Constraint}_i = \sum_{j=1}^{n} P_{ij} + \sum_{q=1}^{n} P_{iq} P_{pq}$$

Eigenvector centrality. A graph consisting of nodes connected by edges can be characterized by an adjacency matrix $A$, populated by elements such that $a_{ij} = 1$ if nodes i and j are directly connected, and $a_{ij} = 0$ if these nodes are not connected. The eigenvector centrality of each node is given by the eigenvector of $A$ in which all elements are positive. The requirement that all elements of the eigenvector must be positive yields a unique eigenvector solution (that is, that corresponding to the greatest eigenvalue). Here, when computing eigenvector centrality, the directionality of the graph was preserved; in the event of asymmetric relationships, only incoming, rather than outgoing, ties were used to compute eigenvector centrality.

Social distance. Geodesic social distance refers to the smallest number of intermediary social ties required to connect two individuals in a network. Individuals whom a participant named as friends have a distance of one from him/her. Individuals whom a participant’s friends named as friends (but who each named as friends by the participant) have a distance of two from the participant. Individuals who were named as friends by classmates at a distance of two from the participant (but not by the participant or his/her friends) have a distance of three, and so on.

Part 2: Neuroimaging study. Participants. A subset of individuals who had completed Part 1 participated in a subsequent neuroimaging experiment. Participants were informed during class about the opportunity to participate in an fMRI study that was ostensibly unrelated to the online questionnaire in Part 1, and that they would receive $20 per hour as compensation and images of their brains. All participants were right-handed, fluent in English, and had normal or corrected-to-normal vision. Participants provided informed consent in accordance with the policies of the Dartmouth College Committee for the Protection of Human Subjects. Twenty-four participants (12 females) completed the fMRI study. The sample size was chosen based on previous fMRI studies using similar paradigms and RSA methods16,17. One participant was excluded owing to image artifact, and two were excluded because they scored less than 65% correct on the one-back memory task used in the scanner (this threshold was based on what has been used previously in similar studies7). Consequently, we analysed data from 21 participants (10 females, aged 25–33, mean 27.9 years, standard deviation 2.16 years). As a within-subjects design involving no group allocation was used, blinding investigators to between-subjects conditions and random assignment of participants to conditions was not applicable.

Image acquisition. Participants were scanned at the Dartmouth Brain Imaging Center using a 3T Philips Achieva Intera scanner with a 32-channel head coil. An echo-planar sequence (35 ms TE; 2000 ms TR; resolution 3.0 mm × 3.0 mm × 3.0 mm; matrix size 80 × 80; field of view 240 × 240 mm; 35 interleaved transverse slices with no gap; slice thickness 3.0 mm) was used to acquire functional images. Functional runs consisted of 180 dynamic scans, for a total acquisition time of 360 s per run. A high-resolution T1-weighted anatomical scan was acquired for each participant (8.2 s TR; 3.7 ms TE; field of view 240 × 187; resolution 0.938 mm × 0.938 mm × 1.0 mm) at the end of the scanning session. Foam padding was placed around subjects’ heads to minimize motion.

Stimuli. Each participant’s customized stimulus set consisted of short videos of four individuals at each of three geodesic distances (one, two and three) from the participant in the social network of first-year MBA students. The two highest- and lowest-EC individuals at each social distance were included in the stimulus set (Fig. 2).

The videos used as stimuli consisted of individuals introducing themselves (for example, “Hi, my name is [first name], and you can call me [first name]/[nickname]”). A video of this kind was made involving the participant at the beginning of the academic year as a resource for other students and faculty. Videos were truncated to 2 s, beginning when the subject began to say the word “Hi”, and were presented without sound. Prior to entering the fMRI scanner, participants were shown each video with sound to familiarize themselves with the stimuli.

fMRI paradigm. The fMRI study consisted of ten runs and followed a rapid event-related design with an inter-trial interval consisting of 4 s of fixation (Fig. 2c). Four null events, each consisting of an additional 2 s of fixation, were randomly inserted into each run. In each run, four replications of 14 event categories (12 identity trials; 1 null event; 1 catch trial) were pseudo-randomized such that there were no consecutive repeats of the same category. Horizontal mirroring was randomly applied to half the presentations of each stimulus within each run to reduce similarities within identities due to local low-level visual features. Catch trials involved seeing the same stimulus at the same mirroring level as the immediately previous stimulus (or two trials back if a catch trial followed a null event). Participants were instructed to press a button when an identical video was presented twice in a row (that is, for catch trials).
Post-scan questionnaire. After scanning, participants were asked about their subjective perceptions of each social network metric of interest for each individual in their stimulus set, as well as questions assessing tie strength (see Supplementary Information). Because the constraint question asked about brokerage (that is, which individuals were low in network constraint), responses to this item were multiplied by −1. To alleviate skew in the network data, eigenvector centralities and network constraint values were log-transformed prior to analysis.

The corresponding parameters used in fMRI post-scan ratings and the social network position characteristics of the individuals in their stimulus sets was assessed using linear mixed models using the R package lme4\(^4\). For each of the five questions (see Supplementary Information), a model was constructed with participants’ ratings as the dependent measure and the relevant social network position characteristics as a fixed effect, as well as random intercepts and slopes for each participant. To test the significance of the relationship between participants’ ratings and social network data, p-values were computed using Satterthwaite’s approximation for degrees of freedom\(^\alpha\) as implemented in lmerTest\(^4\).

fMRI data preprocessing. For fMRI data analysis, data were preprocessed and average voxel-wise haemodynamic responses to each identity were estimated using the AFNI 3dDeconvolve\(^1\). Pre-processing steps included applying AFNI’s 3dDespik function to remove transient, extreme values in the signal not attributable to biological phenomena, slice-time correction to correct for interleaved slice acquisition order, alignment of the last volume of the final run to the high-resolution anatomical scan, registration of all functional volumes to the anatomically aligned functional volume using a six-parameter algorithm for 3D motion correction, spatial smoothing using a 4-mm full-width at half-maximum Gaussian kernel, and scaling each voxel time series to have a mean amplitude of 100. Prior to regression, consecutive volumes for which the Euclidean norm of the derivatives of the motion parameters exceeded 0.3 mm were excluded from further analysis, as were volumes in which more than 10% of brain voxels were identified as outliers by the AFNI programme 3dIdentify\(^\alpha\).

Parameter estimates were extracted for each voxel using a GLM that consisted of gamma-variate convolved regressors for each of 13 predictors (one for each of the 12 identities in the participant’s stimulus set; one for each trial), as well as 12 regressors for each of the six de-meaned motion parameters extracted during volume registration and their derivatives, and three regressors for linear, quadratic and cubic signal drifts within each run. This procedure removed variance caused by regressors of no interest and resulted in an estimate of the response of each voxel to each trial type.

GLM decomposition searchlight. Using PyMVPA\(^19\) and SciPy\(^17\), a GLM decomposition searchlight\(^1\) was performed within each participant’s data. A radius (radius \(r\)) was defined, such that, throughout each participant’s brain, at each point in the brain, the local distributed patterns of neural responses to each person in the stimulus set were extracted within a sphere centred on that point, and the pairwise correlation distances between them were calculated to construct a local neural RDM (Fig. 3a–c), which was decomposed into a weighted combination of predictor RDMs using ordinary least squares regression. Thus, the resultant predictor RDMs were used orthogonal to one another prior to performing the GLM decomposition searchlight.

Because RDMs are symmetric about a diagonal of zeros, all RDMs were flattened to form vectors of their above-diagonal elements prior to performing GLM decomposition searchlight. At each searchlight centre (each voxel), the GLM decomposition searchlight procedure yielded a \(\beta\) value corresponding to how much the variance in the neural activity of interest was explained by the corresponding predictor RDM (equation 1). This procedure removed variance caused by the local distributed patterns of neural responses to each person in the stimulus set.

Group analysis. Each subject’s maps of regression coefficients and \(R^2\) values were transferred to standard (Talairach\(^1\)) space using AFNI anatomical scans were linearly aligned to the subject’s respective template using the 3dAFNE template using the 3dTemplate algorithm in AFNI, and the same transform was used to align each participant’s searchlight results to standard space prior to group analysis. To identify areas that reliably contained information about each specific aspect of social network position across participants, the regression coefficients for each RDM derived from social network position were tested against 0 across participants within a diluted grey matter mask using one-tailed (that is, whether a regression coefficient was greater than 0), one-sample \(t\)-tests. More specifically, FSL’s ‘randomise’\(^52\) programme was used to perform permutation tests and to generate a null distribution of cluster masses for multiple comparisons correction (cluster-forming threshold: \(p < .01\), two-tailed; 5,000 permutations; 10-mm variance smoothing). All reported results have been thresholded at a family-wise error rate of 5%.

Data availability. The data that support the findings of this study are available from the corresponding author upon reasonable request. Interactive 3D visualizations of the data depicted in Fig. 4 are available online at http://neurovault.org/collections/1BPYPUU/.

Code availability. The code used for the analyses is available from the corresponding author upon request.

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Author contributions
C.P., A.M.K. and T.W. conceived and designed the study. C.P. and A.M.K. collected the data. C.P. analysed the data. C.P., A.M.K. and T.W. wrote the paper.

Additional information
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Competing interests
The authors declare no competing interests.
Spontaneous neural encoding of social network position

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Supplementary Figures

Supplementary Figure 1. Associations between perceived and actual social network characteristics. Black dashed lines depict the relationships between perceived and actual social network characteristics across all participants (fit using an ordinary least squares linear model). Solid purple, orange and green lines depict these relationships for each subject for social distance, eigenvector centrality, and constraint, respectively. (A) Neuroimaging study participants’ subjective ratings of social closeness, proportion of social time spent together, and frequency of discussions with the individuals in their stimulus sets varied according to geodesic network distance from them in the network (all \( p’s < .0001; \) see main text). (B) Participants’ estimates of the eigenvector centrality of the individuals in their stimulus sets were closely related to those individuals’ actual eigenvector centralities \( (p < .0001; \) see main text). (C) Participants’ estimates of the network constraint of individuals in their stimulus sets were also associated with the actual constraint of those individuals’ positions in the social network \( (p < .0001; \) see main text). As described in the main text, self-report data was obtained after scanning; network constraint and eigenvector centrality were log-transformed prior to plotting and analysis to alleviate skew. Perceived network constraint ratings were multiplied by -1 prior to plotting because the relevant question...
asked participants to rate perceived brokerage (which is inversely related to network constraint). Analyses of behavioral ratings were conducted using linear mixed models that included by-subject random slopes and intercepts.
Supplementary Figure 2. Voxel-wise $R^2$ values, averaged across subjects, are depicted within clusters that reliably signaled one or more of the tested aspects of social network position. The $R^2$ value corresponding to the GLM decomposition performed at each searchlight center indicates the extent to which the information contained in local multi-voxel response patterns can be explained by the social network positions of the classmates being viewed. Results are projected onto a cortical surface model of the Talairach\textsuperscript{1} N27 brain using PySurfer (https://github.com/nipy/PySurfer). Brain regions the reliably signaled one or more of the tested aspects of social network position are shown in Fig. 4 of the main text.
Supplementary Figure 3. Relationship between network constraint and movement during videos. The amount of movement of the 88 individuals whose videos were used as stimuli was not significantly related to the constraint characterizing those individuals’ positions in the social network of first-year MBA students, $r = -0.12$, $p = 0.28$. Solid line indicates an ordinary least squares fit to the data; shaded region indicates 95% bootstrapped confidence interval of the slope of the regression line.
Supplementary Figure 4. Post-scan questionnaire. Following scanning, participants responded to questions about their subjective perception of each aspect of social network position of interest for each individual in their stimulus set. A screenshot of the question corresponding to network constraint (reverse-scored) is shown.
### Supplementary Tables

**Supplementary Table 1**: Brain regions where local neural information content is associated with the social distance of the individuals being viewed.

<table>
<thead>
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<th>Size (mm$^3$)</th>
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<th>COG y</th>
<th>COG z</th>
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</tbody>
</table>

Hemi = hemisphere; COG = center of gravity; L = left; R = right; IPL = inferior parietal lobule; SMG = supramarginal gyrus; STG = superior temporal gyrus; STS = superior temporal sulcus; MTG = middle temporal gyrus. All reported results are significant at a statistical threshold of $p < .05$, FWE-corrected. All coordinates are in Talairach space.

**Supplementary Table 2**: Brain regions where local neural information content is associated with the eigenvector centrality of the individuals being viewed.

<table>
<thead>
<tr>
<th>Hemi</th>
<th>Size (mm$^3$)</th>
<th>COG x</th>
<th>COG y</th>
<th>COG z</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>L</td>
<td>24,483</td>
<td>-42.4</td>
<td>-18.5</td>
<td>33.6</td>
<td>IPL, IFG, Ins., pre-central gyrus</td>
</tr>
<tr>
<td>R</td>
<td>8,768</td>
<td>21.7</td>
<td>26.8</td>
<td>-4.9</td>
<td>MPFC, IFG, aIns., ant. PHG, TP</td>
</tr>
<tr>
<td>L</td>
<td>7,716</td>
<td>-32.7</td>
<td>12.9</td>
<td>-6.1</td>
<td>aIns., IFG</td>
</tr>
<tr>
<td>L, R</td>
<td>7,552</td>
<td>-9.4</td>
<td>-41.8</td>
<td>40.4</td>
<td>PCC, precuneus</td>
</tr>
<tr>
<td>R</td>
<td>6,802</td>
<td>20.4</td>
<td>-48.0</td>
<td>-2.8</td>
<td>PHG, LG, FG</td>
</tr>
<tr>
<td>L, R</td>
<td>6,056</td>
<td>48.0</td>
<td>-29.4</td>
<td>38.4</td>
<td>IPL, precuneus, post-central gyrus</td>
</tr>
<tr>
<td>L, R</td>
<td>5,233</td>
<td>-0.8</td>
<td>-44.4</td>
<td>65.4</td>
<td>Precuneus, post-central gyrus</td>
</tr>
<tr>
<td>R</td>
<td>4,961</td>
<td>-0.6</td>
<td>-83.3</td>
<td>28.5</td>
<td>EVC</td>
</tr>
</tbody>
</table>

Hemi = hemisphere; COG = center of gravity; L = left; R = right; a = anterior; IPL = inferior parietal lobule; IFG = inferior frontal gyrus; Ins. = insula; MPFC = medial prefrontal cortex; PHG = parahippocampal gyrus; TP = temporal pole; PCC = posterior cingulate cortex; LG = lingual gyrus; FG = fusiform gyrus; EVC = extrastriate visual cortex. All reported results are significant at a statistical threshold of $p < .05$, FWE-corrected. All coordinates are in Talairach space.

**Supplementary Table 3**: Brain regions where local neural information content is associated with the constraint of the individuals being viewed.

<table>
<thead>
<tr>
<th>Hemi</th>
<th>Size (mm$^3$)</th>
<th>COG x</th>
<th>COG y</th>
<th>COG z</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>R</td>
<td>11,872</td>
<td>51.5</td>
<td>-16.0</td>
<td>-3.0</td>
<td>STS, STG, MTG, ITS, pIns.</td>
</tr>
<tr>
<td>L</td>
<td>7,739</td>
<td>-51.6</td>
<td>-38.5</td>
<td>7.1</td>
<td>STS, STG, MTG, pIns.</td>
</tr>
<tr>
<td>R</td>
<td>4,363</td>
<td>11.5</td>
<td>-5.7</td>
<td>58.4</td>
<td>SMA, dorsal premotor cortex</td>
</tr>
</tbody>
</table>

Hemi = hemisphere; COG = center of gravity; L = left; R = right; STS = superior temporal sulcus; STG = superior temporal gyrus; MTG = middle temporal gyrus; ITS = inferior temporal sulcus; pIns. = posterior insula; SMA = supplementary motor area. All reported results are significant at a statistical threshold of $p < .05$, FWE-corrected. All coordinates are in Talairach space.
Supplementary Methods

Optical flow analysis. To quantify the amount of movement within each video clip used as a stimulus in the neuroimaging experiment, the average optical flow (i.e., the pattern of apparent motion between consecutive video frames) was computed for each video that was shown in the fMRI study. Given that the videos used as stimuli were recorded by a stable camera against a plain, static background, optical flow estimates for these videos capture of the amount that each individual moved his or her facial features and head in the video clip. Farneback’s algorithm for motion estimation\(^2\) as implemented in OpenCV\(^3\) was used to estimate the average magnitude of optical flow in each video. This method extracts a pixel-wise motion vector for each pair of sequential frames in which each pixel is characterized by a magnitude and a direction. To estimate the magnitude of motion within each frame pair, the magnitude values (without respect to direction) were summed across pixels. To compute the mean magnitude of optical flow for a given video, the motion magnitude estimates were averaged across frames within that video.

In order to test whether or not individual differences in network constraint are related to movement in the videos used as stimuli, the correlation between network constraint and average motion magnitude was assessed among the 88 individuals whose videos were used as stimuli in the fMRI study. Given that distributions of both variables were highly skewed, data were log-transformed prior to analysis. The results of this procedure suggest that in the stimuli used in the current study, network constraint
and amount of movement were not significantly correlated, $r = -0.12, p = 0.28$ (see Supplementary Fig. 3).

**Post-scan questionnaire.** Participants performed the post-scan questionnaire on a 13" MacBook laptop using Psychopy\(^4\). Participants first viewed an instruction screen that read, “Now you will see the same people who you saw in the scanner. You will be asked questions about each person. These questions relate only to this person’s interactions within the [institution name] MBA cohort. We understand that people have many social circles that they participate in (perhaps including family, friends outside of [the institution], other contacts, etc.). For these questions, please just consider interactions within the MBA cohort. You will be presented with a continuous rating scale for each question. You can choose any point along the continuum to respond. Press any key to continue.” During the survey, videos of the 12 individuals from the participant’s stimulus set were presented in a random order. Participants responded to all questions about a given individual sequentially, and the same video that had played in the scanner repeated on a loop (without sound) above the question text and response scale (see Supplementary Fig. 4).

Participants were presented with questions concerning lay definitions of eigenvector centrality (“In social network analysis, scientists assess a construct that measures how many friends a person has, and how many friends a person’s friends have. How would you rate this person on this construct?” Responses ranged from “Low (few friends who have few friends)” to “High (many friends who have many friends)”) and constraint (“Social network analysts also assess a construct called ‘brokerage’ that
measures how much a person connects groups of people who wouldn’t otherwise be connected. Using this definition, how high is this individual in ‘brokerage’?” Responses ranged from “Low (this person never connects distinct groups of people)” to “High (This person often connects distinct groups of people)”). Responses to the item assessing brokerage were reverse scored in order to estimate perceived network constraint.

Participants were also presented with the name generator that had originally been used to construct the network (“Consider the people with whom you like to spend your free time. During the last month, is this one of the classmates who you have been with most often for informal friendship activities, such as going out to lunch, dinner, drinks, films, visiting one another’s homes, and so on?” Responses ranged on a continuum from “None of my social activities in the past month have included this person” to “All of my social activities in the past month have included this person”), as well as questions designed to assess tie strength (“How close are you with this person?” Responses ranged from “Distant” to “Less than close” to “Close” to “Especially Close”) and frequency of interactions (“On average, how often do you talk to this person (any social or business discussion)?” Responses ranged from “Less often” to “Monthly” to “Weekly” to “Daily”).
Supplementary References