Computational dissection of human episodic memory reveals mental process-specific genetic profiles

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Episodic memory performance is the result of distinct mental processes, such as learning, memory maintenance, and emotional modulation of memory strength. Such processes can be effectively dissociated using computational models. Here we performed gene set enrichment analyses of model parameters estimated from the episodic memory performance of 1,765 healthy young adults. We report robust and replicated associations of the amine compound SLC (solute-carrier) transporters gene set with the learning rate, of the collagen formation and transmembrane receptor protein tyrosine kinase activity gene sets with the modulation of memory strength by negative emotional arousal, and of the L1 cell adhesion molecule (L1CAM) interactions gene set with the repetition-based memory improvement. Furthermore, in a large functional MRI sample of 795 subjects we found that the association between L1CAM interactions and memory maintenance revealed large clusters of differences in brain activity in frontal cortical areas. Our findings provide converging evidence that distinct genetic profiles underlie specific mental processes of human episodic memory. They also provide empirical support to previous theoretical and neurobiological studies linking specific neuromodulators to the learning rate and linking neural cell adhesion molecules to memory maintenance. Furthermore, our study suggests additional memory-related genetic pathways, which may contribute to a better understanding of the neurobiology of human memory.

learning and memory | computational model | gene set enrichment analysis | functional neuroimaging | frontal lobe

uman memory is a highly heritable and complex trait that is the outcome of numerous neurocognitive processes (1, 2). Although many genetic variations associated with human memory performance have been identified (3-9), most were based on general memory scores, such as the number of recalled items. Such scores may serve as good correlates of overall memory ability, but they offer little insight into the genetic underpinnings of specific memory-related mental processes, such as encoding, memory maintenance, modulation by emotional arousal, or guessing strategies. Importantly, neurobiological and molecular profiles of such processes are known to be partly distinct (10-12). Because some of the mental processes involved in memory are not readily amenable to direct observation, computational modeling can be used to make inferences about them (13, 14). Modelbased analyses provided insights into neurocomputational mechanisms of reward-based learning and decision making (15–17), related model parameters such as the learning rate to genetic polymorphisms (18, 19), and provided a computational explanation for the inverted-U-shaped relation between stress intensity and behavioral performance (20). As a relatively recent development,

the model-based analysis approach has largely been missing from studies of human episodic memory and genome-wide association studies (GWAS).

GWAS already had identified a number of associations between single genetic markers and cognitive traits, such as episodic memory (7–9) and modulation of memory strength by negative emotional arousal (21). However, because many such traits are complex and multigenic, it is unlikely that a single genetic variant will explain a sufficient amount of behavioral variability. As a statistically more powerful and biologically meaningful alternative, gene set-based methods of linking sets of biologically related genes with traits of interest have been developed (22, 23). They test whether a group of genes (gene set) is more enriched in associations with a studied phenotype than would be expected by chance. Such studies identified gene sets related to neuropsychiatric phenotypes (24, 25) and to cognition in healthy individuals, including working memory (26) and emotionally aversive memory (27).

Significance

Human memory is a highly heritable and complex trait that is the outcome of several neurobiologically distinct mental processes, such as learning, memory maintenance, and emotional modulation of memory strength. Such processes, which are not always amenable to direct observation, can be dissociated using computational models. Here we provide converging evidence linking computational model parameters and related behavioral and neuroimaging phenotypes to sets of biologically related genes in several populations of healthy young and elderly adults. Our findings suggest that distinct genetic profiles underlie specific mental processes of human episodic memory. Understanding the genetic and neural bases of such processes is essential for the development of novel therapeutic approaches that could lead to a better treatment of neuropsychiatric disorders.

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Table 1. Samples and core phenotypes used

Sample name	Phenotypes of	
(location)	interest	Sample size
Discovery (Zurich and Basel)	Computational model parameters	1,239
Replication (Basel)	Computational model parameters	526
Words/pictures* (Basel)	Free recall of pictures	493
Pictures/fMRI (Basel)	Free recall of pictures	835
	Picture recognition	822
	Working memory	825
	fMRI data on picture recognition	795
	fMRI data on working memory	797
AgeCoDe elderly (Bonn)	Free recall of words	743

*This is a subsample of the Replication sample. All other samples are independent.

In the present study, we performed a gene set enrichment analyses (GSEA) of specific parameters of episodic memory performance estimated from a computational model of verbal memory in young adults (21). We found four distinct gene sets associated with different computational parameters in independent samples, three of which also were linked to related measures of picture memory in young adults and verbal memory in elderly individuals. The L1 cell adhesion molecule (L1CAM) interactions gene set also was associated with differences in picture recognition-related brain activity in a number of frontal cortical areas.

Results

The verbal memory task contained neutral, positive, and negative words, which the participants had to recall freely first after their presentation and later after an ~5 min delay. Participants in the discovery sample (n = 1,239; Table 1) and the replication sample (n = 526) performed the identical verbal memory task. Eight performance measures (PM₁₋₈; Table 2) indicated the number of correctly recalled words in each valence category (i.e., neutral, positive, and negative) and confabulative errors (words not on the learning list) in the immediate and the delayed recall. Because performance measures were partly correlated, the effective dimensionality of data was smaller: principal component analysis (PCA) revealed five major components (Fig. S1).

Computational Modeling. To study specific cognitive processes relevant for memory performance, we used a simple computational model of learning in the verbal task (21). Our model had the following parameters, which we aimed to infer from the behavioral data: the learning rate α and the Gaussian noise σ , the repetition-based memory improvement *c* (related to memory maintenance) and the forgetting rate γ , the modulation of

memory strength by positive or negative emotional arousal ε_{pos} and ε_{neg} , the decision threshold β , and the sigmoidal steepness s (for a detailed description see Materials and Methods). Because the effective dimensionality of the behavioral data was lower than the number of model parameters, we empirically selected five of these parameters (α , β , c, ε_{pos} , ε_{neg}) to be estimated individually, keeping the three remaining ones fixed. After the estimation (Fig. S2), more than 99% of the resulting individual parameter sets passed the χ^2 test of goodness-of-fit [mean χ^2 (discovery) = 1.5057 and χ^2 (replication) = 1.4955], indicating that the model was sufficiently flexible. High correlations [mean $\rho_{\text{Spearman}}(\text{discovery}) = 0.981 \text{ and } \rho_{\text{Spearman}}(\text{replication}) = 0.986]$ among the 10 best parameter sets (hill climbing end points in stage 2 of the estimation) suggested high estimation reliability, and the presence of 99.8% of individual parameter sets within the middle 90% of the value ranges (except the bound of c = 1, because repetition should not weaken memories) indicated that parameter estimation bounds did not constrain the results.

GSEA. We first performed a GWAS with the estimated model parameters (α , β , c, ε_{pos} , ε_{neg}) with 1,239 individuals in the discovery sample (for quality control and statistics, see Materials and Methods). Based on the resulting P values of association with these parameters, we performed a GSEA using MAGENTA (23). Of the 1,411 gene sets used for input, we found false-discovery rate (FDR)-corrected enrichment for seven gene sets (Table 3). A second GSEA was performed for these gene sets in an independent sample of 526 individuals (Table 1), again using GWAS-derived P values of association with the estimated model parameters. Of the seven gene sets significantly enriched in the discovery sample, four largely nonoverlapping gene sets were also enriched in the replication sample (Table 3): amine compound solute-carrier (SLC) transporters associated with the learning rate α , L1CAM interactions associated with the repetition-based memory improvement c, and collagen formation and transmembrane receptor protein tyrosine kinase activity associated with ε_{neg} . Although behavioral measures of verbal memory were correlated with model parameters (Table S1), none of the gene sets reported above would have been detected if these behavioral measures had served as starting points for GSEA (using the same multiple testing correction threshold as for the model parameters); however, five of the seven gene sets identified in the discovery sample were nominally associated with the relevant behavioral measures.

We next examined if the four replicated gene sets showed enrichment in associations with related, albeit less specific, behavioral measures in different tasks and in additional populations (Table 1). Computational modeling was not practical for these tasks because of the low dimensionality of the behavioral data. In the sample of nondemented elderly individuals from the German Study on Aging, Cognition and Dementia in Primary Care Patients (AgeCoDe) (n = 743), participants performed a word-list verbal recall task from the Consortium to Establish a Registry for Alzheimer's Disease (CERAD) neuropsychological test battery (28). In a picture task, performed by two populations of healthy young adults, participants were presented neutral, positive, and negative pictures, which they had to recall freely 10 min after presentation. After a longer delay, participants of the

Table 2. Description of performance measures (PM₁₋₈) in the verbal memory task and their population statistics

Performance measure	Discovery sample, mean \pm SEM	Replication sample, mean \pm SEM
PM ₁ : Positive words correctly recalled immediately	7.666 ± 0.042	7.645 ± 0.058
PM ₂ : Negative words correctly recalled immediately	8.318 ± 0.036	8.364 ± 0.050
PM ₃ : Neutral words correctly recalled immediately	7.392 ± 0.047	7.345 ± 0.065
PM ₄ : Mistakes made immediately after encoding	1.302 ± 0.038	1.164 ± 0.050
PM ₅ : Positive words correctly recalled after 5 min	2.948 ± 0.043	3.055 ± 0.061
PM ₆ : Negative words correctly recalled after 5 min	2.645 ± 0.041	2.882 ± 0.059
PM ₇ : Neutral words correctly recalled after 5 min	2.448 ± 0.041	2.618 ± 0.056
PM ₈ : Mistakes made 5 min after encoding	1.681 ± 0.048	1.727 ± 0.071

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Table 3. Associations between computational model parameters for the verbal task and different gene sets

		Gene set	Discovery $(n = 1,$	Replication sample $(n = 526)$	
Model parameter	Database	(number of genes in gene set)	P _{nominal}	P _{FDR}	$P_{\rm nominal}$
Learning rate α	Reactome	Amine compound SLC transporters (27)	$2.0 imes 10^{-4}$	0.0405	0.0031
Decision threshold β	KEGG	Axon guidance (129)	$4.4 imes 10^{-5}$	0.0067	0.405
Repetition-based	Reactome	L1CAM interactions (86)	$3.0 imes 10^{-6}$	0.0011	0.0231
memory improvement c		Interaction between L1 and ankyrins (23)	1.0×10^{-3}	0.0365	0.176
Modulation by negative	Reactome	Collagen formation (58)	$1.0 imes 10^{-4}$	0.0129	0.0025
emotional arousal $\epsilon_{\rm neg}$	Gene Ontology	Transmembrane receptor protein tyrosine kinase activity (43)	4.1×10^{-5}	0.0124	0.0212
		Transmembrane receptor protein kinase activity (51)	$1.0 imes 10^{-4}$	0.0398	0.059

Only the gene sets surviving MAGENTA FDR correction in the discovery sample are shown. The replicated gene sets are shown in bold. Interaction between L1 and ankyrins is a subset of L1CAM interactions, and transmembrane receptor protein tyrosine kinase activity is a subset of transmembrane receptor protein kinase activity. L1CAM interactions contains four genes [EGFR, FGFR1, and neuropilin 1 and 2 (NRP1, and NRP2)] shared with transmembrane receptor protein tyrosine kinase activity.

Basel pictures/functional MRI (fMRI) sample also were asked to perform picture recognition: provided with both previously shown and new pictures, they had to answer whether they had seen each picture. For all of these tasks, we selected behavioral measures most closely related to each computational parameter. For the learning rate α , immediate recall is more appropriate than delayed recall, because the latter also accounts for forgetting. However, because immediate recall was not available in the picture task, we used free delayed recall (see also Table S1). For the repetition-based memory improvement c, only delayed recall was appropriate, because immediate recall did not account for memory maintenance. In the picture task both free recall and recognition measured memory over delays; however, because we also aimed to study the neural correlates of memory maintenance using functional neuroimaging, which were available only for picture recognition data, we chose the latter phenotype. For negative emotional modulation factor ε_{neg} , only picture data were available; therefore we chose the difference between freely recalled negative and neutral pictures as a relevant phenotype. Based on the GWAS results of these phenotypes (population statistics are shown in Table 4), we used MAGENTA for testing the enrichment of the four gene sets associated with different computational parameters of the (original) verbal task. We found that amine compound SLC transporters were associated with immediate recall of words in the elderly sample and with free recall of pictures in both young samples; L1CAM interactions were associated with delayed recall of words in the elderly sample and with picture recognition in the pictures/fMRI sample; collagen formation was associated with free recall of negative minus neutral pictures in the words/pictures sample; and there were no significant associations with transmembrane receptor protein tyrosine kinase activity (Table 5). Because the arousal of negative pictures [2.36 (mean) \pm 0.06 (SEM)] was higher than that of positive pictures (1.94 \pm 0.04; Student *t* test; P = 0.000002), the association of collagen formation with picture recall may be mediated by arousal rather than by negative valence. We also checked if sex was correlated with any of the significant SNPs (one per gene) in any of the significantly enriched gene sets (Tables 3 and 5). No significant correlations, corrected for the number of SNPs per gene set, were found (*Ps* > 0.05).

Neural Correlates of the Discovered Gene Sets. Because the amine compound SLC transporters and L1CAM interactions gene sets were associated with free recall and recognition of pictures, respectively, in the pictures/fMRI sample, we examined neural correlates of these associations using functional neuroimaging. Participants in this sample performed the encoding and recognition parts of the picture task in the fMRI scanner; free recall was performed outside the scanner. We studied association with amine compound SLC transporters (related to the learning rate α) by using two encoding contrasts: all pictures vs. scrambled controls and the difference due to memory (Dm, i.e., subsequently remembered vs. not remembered pictures; 29). Because neural correlates of memory maintenance may not be apparent during encoding, we examined the association with L1CAM

Table 4. Population statistics of relevant phenotypes for the picture task and for the word task used in the German elderly sample

Data sample	Mean \pm SEM
Words/pictures ($n = 493$)	39.58 ± 0.49
Words/pictures ($n = 493$)	17.17 ± 0.59
Pictures/fMRI ($n = 835$)	42.57 ± 0.39
Pictures/fMRI ($n = 835$)	17.67 ± 0.45
Pictures/fMRI ($n = 822$)	
	80.74 ± 0.55
	15.68 ± 0.49
	3.57 ± 0.15
	1.02 ± 0.07
	9.41 ± 0.28
	89.57 ± 0.31
AgeCoDe elderly ($n = 743$)	19.55 ± 0.13
AgeCoDe elderly ($n = 743$)	5.88 ± 0.07
	Data sample Words/pictures (n = 493) Words/pictures (n = 493) Pictures/fMRI (n = 835) Pictures/fMRI (n = 835) Pictures/fMRI (n = 822) AgeCoDe elderly (n = 743) AgeCoDe elderly (n = 743)

*Previously seen pictures rated as known.

Table 5.	Association between	the four replicate	ed gene sets and	other related	phenotypes
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Database	gene set, associated model parameter)	Relevant phenotype	Data sample	Association P value
Reactome	Amine compound SLC	Percentage of correctly	Words/pictures	0.0302
	transporters (27, α)	remembered pictures in the free recall	Pictures/fMRI	0.0001
		Number of correctly remembered words in the immediate recall	AgeCoDe elderly	0.032
	L1CAM interactions (86, c)	Percentage of correctly recollected pictures in picture recognition	Pictures/fMRI	0.0419
		Number of correctly remembered words in the delayed recall	AgeCoDe elderly	0.0068
	Collagen formation (58, ε_{neg})	Percentage of correctly	Words/pictures	0.0206
		remembered negative pictures minus neutral pictures in the free recall	Pictures/fMRI	0.145
Gene Ontology	Transmembrane receptor	Percentage of correctly	Words/pictures	0.147
	protein tyrosine kinase activity (43, ϵ_{neg})	remembered negative pictures minus neutral pictures in the free recall	Pictures/fMRI	0.153

Gene set (number of genes in

interactions (related to the repetition-based memory improvement c) using the recognition contrast of previously seen vs. new pictures, a task that engages the brain areas involved in memory maintenance and retrieval, such as prefrontal cortex and medial temporal lobe (30, 31). To capture multiallelic effects of each gene set on the differences in brain activity, we generated individual multilocus genetic scores (*Materials and Methods*). For L1CAM interactions, the score was built using 28 significant SNPs from the same number of genes (one most significant SNP per gene; Table S2); the genetic score for amine compound SLC transporters comprised 14 SNPs.

Genetic score-dependent analysis revealed no whole-brain familywise error (FWE)-corrected correlations with differences in brain activity for amine compound SLC transporters. However, the L1CAM interactions genetic score was negatively correlated at the whole-brain FWE-corrected level with differences in the activity of numerous frontal regions: large clusters in the left superior frontal cortex and the left inferior frontal gyrus (Fig. 1, A and B and Table 6) and small clusters in the bilateral orbitofrontal cortices and insulae (Table 6). The results were very similar if the two most significant SNPs per gene were used to build a genetic score (Table S3) or if only correctly rated pictures were included in the contrast (recollected previously seen pictures vs. new pictures rated as new). We also performed a genetic score-independent analysis of brain activity, which revealed substantial differences in activity at all cluster peaks dependent on the L1CAM interactions genetic score (Fig. 1, C and D), suggesting that higher values of the genetic score reflect smaller increases in frontal activity during the recognition of previously seen pictures as compared with the viewing of new pictures. To ensure that the observed effect specifically reflected differences in the episodic memory of previously seen pictures, we also used the contrast between recollected previously seen pictures vs. previously seen pictures rated as familiar or new (n =786). Of the 238 voxels significant in the contrast between previously seen vs. new pictures, the great majority also passed small-volume correction (SVC) for the latter contrast [162 voxels in superior frontal cortex, peak coordinates (0, 17, 48), T = -4.93; four voxels in left lateral orbitofrontal cortex, peak coordinates (-30, 22, -8), T = -5.07], with the peak voxels also surviving whole-brain correction.

Because the differences in brain activity at all cluster peaks dependent on the genetic score of L1CAM interactions also were negatively associated with recognition performance (Table S4, n = 795), we performed a mediator analysis to exclude the possibility that the association between the genetic score and the differences in brain activity could be explained fully by the correlated behavioral phenotype. We examined whether the differences in activity in these brain regions were mediating the correlation

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between the genetic score and picture recognition performance. We found that all four clusters acted as significant mediators of the genetic score–behavior relationship (for more details about mediation, see *Materials and Methods* and Fig. S3).

As two additional controls, we first extracted average beta values for the two largest clusters, whose activity was associated with the picture recognition phenotype at the whole-brain FWE-corrected level [cluster 1: peak coordinates (0, 28, 40), T = -12.36, number of voxels = 2,917; cluster 2: peak coordinates (-47, 41, 16), T = -10.57, number of voxels = 1,541] and subjected them to GSEA using MAGENTA. The L1CAM interactions gene set was significantly enriched in both cases, with



Fig. 1. Functional neuroimaging of picture recognition (the contrast between previously seen vs. new pictures). (*A* and *B*) Differences in brain activity dependent on the L1CAM interactions genetic score. (*C* and *D*) Genotype-independent differences in brain activity. Color-coded t values are shown. The maps are centered at (-3, 28, 40) in the left superior frontal cortex (*A* and *C*) and at (-47, 44, -8) in the left inferior frontal gyrus (*B* and *D*). Only voxels surviving whole-brain FWE correction are shown (*P* wholebrain FWE-corrected < 0.05; |T| > 4.77; n = 795). Activations are overlaid on sections of the study-specific group template.

Table 6. Association between the L1CAM interactions genetic score and brain activity differences during picture recognition

Regional correspondence of		No. of voxels	Pe coc	Peak MNI coordinates		Peak statistics		
the maximum	Left/right		х	Y	Z	Т	P _{nominal}	Empirical P value
Superior frontal cortex (79%), caudal anterior cingulate cortex (2%)	Left	187	-3	28	40	-6.19	9.4 × 10 ⁻¹⁰	0.0001
Pars orbitalis cortex (34%), pars triangularis cortex (28%), rostral middle frontal cortex (16%)	Left	39	-47	44	-8	-5.70	1.6 × 10 ⁻⁸	0.0021
Lateral orbitofrontal cortex (20%), insula (23%)	Right	6	30	22	0	-5.16	3.2×10^{-7}	0.018
Lateral orbitofrontal cortex (73%), insula (8%)	Left	4	-28	22	-4	-4.87	$1.4 imes 10^{-6}$	0.053

Previously seen vs. new pictures fMRI contrast was used. Only clusters with at least three voxels surviving whole-brain FWE correction are shown (P whole-brain FWE-corrected < 0.05; |T| > 4.77; n = 795). Region names are in accordance with the FreeSurfer nomenclature; probabilities are in accordance with the in-house atlas. For each cluster we also provide empirical P values of the peak voxel based on 10,000 randomly generated genetic scores (null distribution).

nominal P = 0.013 for each cluster. Second, we generated 10,000 genetic scores based on randomly selected SNPs (28 SNPs per genetic score, as for L1CAM interactions) and the picture recognition phenotype, using the same procedure as for the real genetic scores. None of the random scores showed statistically significant, whole-brain-corrected correlation with the neuro-imaging phenotype (the contrast between previously seen vs. new pictures) after correction for the number of genetic scores (threshold $P_{\text{nominal}} = 2.2 \times 10^{-10}$). Based on the null distribution computed with the 10,000 random scores, we also provide empirical P values for the four clusters, which describe the probabilities of their peak voxel significance being reached for the most significant voxel across the whole brain using the random scores (Table 6).

To evaluate the specificity of our imaging genetics finding, we examined if the correlation between the L1CAM interactions genetic score and differences in frontal activity also could be observed during encoding (the subsequent memory contrast, Dm). No such correlation was found at either the whole-brain FWE-corrected level or when applying SVC for 238 voxels significant in the recognition contrast, suggesting either that the relation between L1CAM interactions and differences in frontal activity was not evident during encoding and developed later or that it was specific to the cognitive demands of the recognition task. Because the participants in the pictures/fMRI sample also performed a working memory task (N-back), we examined if the reported association with L1CAM interactions was related to working memory. The MAGENTA analysis performed using Pvalues of GWAS of N-back performance revealed significant enrichment for L1CAM interactions (P = 0.023, n = 825). The multilocus genetic score based on N-back performance was negatively correlated with the one based on picture recognition performance ($\rho_{\text{Spearman}} = -0.140$, $P = 5.9 \times 10^{-5}$, n = 815), suggesting that, with regard to this gene set, better working memory was related to worse picture recognition. Because common frontal mechanisms may mediate both effects, we examined the neural correlates of working memory using fMRI (2-back vs. 0-back contrast, n = 797) and found no correlation between differences in brain activity and the working memory-based L1CAM interactions genetic score at either the whole-brain or SVC levels of significance (number of voxels for the latter, 238). We also examined if the multilocus genetic score based on N-back performance was related to differences in brain activity in the recognition of previously seen vs. new pictures. Again no significant voxels were found at either whole-brain FWE-corrected or SVC thresholds, suggesting that there was no neural evidence implicating L1CAM interactions in working memory or common mechanisms with picture recognition.

Discussion

In this study we show robust associations between multiple gene sets and different computational parameters of verbal memory as well as other relevant neurobehavioral traits in four samples of young and elderly individuals (Table 1). Specifically, amine compound SLC transporters were associated with the learning rate α in two samples of young individuals performing the verbal task, with free recall of pictures in two samples of young individuals, and with immediate recall of words in an elderly sample. L1CAM interactions were linked to the repetition-based memory improvement c (reflecting memory maintenance) in two young samples, to picture recognition and large clusters of differences in recognition-related frontal activity in a third young sample, and to delayed recall of words in the elderly sample. Finally, collagen formation and transmembrane receptor protein tyrosine kinase activity were associated with modulation of verbal memory strength by negative emotional arousal (ε_{neg}) in two young samples, with the former gene set also linked to free recall of negative minus neutral pictures in young individuals.

At this point, it is important to specify the term replication in the context of gene set-based association studies. In contrast to SNP-based studies, in which the object of replication is the significant SNP (or variants in high linkage disequilibrium), the object of replication in gene set-based studies is the significant gene set. Across different samples, replicated gene sets typically are composed of different tagging SNPs (32, 33), a phenomenon related to the allelic and locus heterogeneity of complex traits (22, 34). In our study we did not observe substantial tagging SNP overlap between replicated gene sets across samples (Tables S5 and S6).

The correspondence of significant genes between different samples and phenotypes shows that for amine compound SLC transporters genes encoding GABA, norepinephrine, choline, and vesicular amine transporters are significant in multiple samples, including the discovery sample (Table 7 and Table S5). These genes have been implicated in epilepsy, substance abuse, attention deficit hyperactivity disorder, schizophrenia, and neurodegenerative disorders (35-37). On a neurocomputational level, acetylcholine plays an important role in the encoding of new memories, may act as a switch between encoding and retrieval (38), and has been suggested as a neural correlate of the learning rate (39). Variations in several dopaminergic genes have been linked to the learning rate in the reinforcement learning framework (18, 19) and in economic games (40). Therefore, our results support the notion that the relations between certain neuromodulators and the learning rate generalize for various types of learning models.

For L1CAM interactions, genes encoding fibroblast growth factor receptor (FGFR), ankyrin at node of Ranvier (ANK3), contactins, integrins, and important neural cell adhesion molecules (NRCAM and NCAM1) were significant in multiple samples including the discovery sample (Table 8 and Table S6). These molecules have been implicated in neuronal development and signaling and are essential for synaptic plasticity and its stability in many different brain regions, spatial learning, and memory consolidation (41–44), supporting our discovered association with

Table 7. Correspondence of significant amine compound SLC transporters genes among samples

Gene name	Chromosome	Description	Discovery sample	Replication sample	Pictures/fMRI sample	Words/pictures sample	AgeCoDe sample
SLC6A11	3	GABA transporter 3	0.0184	0.0167	0.0084	0.0004	0.0707
SLC14A2	18	Urea transporter 2	0.0179	0.0159	0.0141	0.0049	0.0619
SLC6A20	3	X transporter protein 3	0.0059	0.1808	0.0092	0.2848	0.0024
SLC6A2	16	Norepinephrine transporter	0.0491	0.1976	0.0141	0.0097	0.0162
SLC44A3	1	Choline transporter-like protein 3	0.0184	0.0529	0.0235	0.0047	0.2009
SLC6A6	3	Taurine transporter	0.0287	0.0371	0.0437	0.0932	0.0235
SLC6A7	5	Brain-specific L-proline transporter	0.0184	0.0275	0.0295	0.1263	0.1680
SLC6A18	5	Sodium channel-like protein	0.0377	0.0482	0.0417	0.0234	0.1872
SLC18A2	10	Vesicular amine transporter 2	0.0491	0.1651	0.0388	0.0205	0.1940

Only genes with significant SNPs in the discovery sample and at least two other samples are shown (the SNP rsIDs and linkage disequilibrium between them are provided in Table 55). The P values of association with relevant phenotypes (Table 5) are provided. Genes are sorted by the product of these values.

a memory maintenance-related computational parameter. Some of them, e.g., *ANK3* and *NCAM1*, also have been related to neurodevelopmental and psychiatric disorders (45) and to chronic stress-related cognitive disturbances (41).

Among collagen formation genes significant in both discovery and replication samples, *TLL2* was linked to mouse avoidance behavior and human bipolar disorder (46), and *ADAMTS14* was linked to multiple sclerosis (47). Collagens are essential for connective tissue and mobility, but they also have been implicated in anxiety- and fear-related behaviors in humans and mice (48–50). Interestingly, we found that the collagen formation gene set was significantly enriched (P = 0.005) in the AgeCoDe elderly sample with Geriatric Depression Scale score (51) as phenotype. Finally, for the transmembrane receptor protein tyrosine kinase activity genes significant in both discovery and replication samples, we did not find references implicating them in emotional memory, anxiety, or affective disorders.

Despite the convergence of our results, particularly for amine compound SLC transporters and L1CAM interactions, the exact processes and gene systems influencing the learning rate and memory maintenance cannot be inferred directly, because GSEA are fundamentally based on gene sets defined in different databases [Reactome, Kyoto Encyclopedia of Genes and Genomes (KEGG), Gene Ontology, BioCarta] that are in the process of being refined and extended. Our use of related behavioral phenotypes rather than computational model parameters in the picture task and the AgeCoDe elderly sample also is a notable limitation. Because a variety of relevant behavioral measures are necessary for fitting nontrivial computational models, it is essential for modelers and experimentalists to collaborate during the process of task design. Furthermore, we note that none of our previously reported GWAS SNPs associated with memory performance, such as those of *KIBRA* (7) and *CTNNBL1* (9), was significantly associated with the model parameters. Memory recall performance and model parameters are not identical phenotypes; thus SNPs significantly associated with a model parameter can be missed when a behavioral measure serves as a starting phenotype, and vice versa.

Functional neuroimaging results revealed a robust negative correlation between the L1CAM interactions genetic score and the differences in activity in left superior and inferior frontal gyri, which have been implicated in the recognition of words and pictures and in working memory in healthy individuals (31, 52–54) and Alzheimer's patients (55, 56). The superior frontal gyrus has been implicated in cognitive control (57), and the inferior frontal gyrus has been implicated in retrieval attempt and effort (58, 59) and depth of memory (60). Reverse inference analysis using Neurosynth (a database of fMRI coordinates and associated

Table 8.	Correspondence of	f significant L1CAM	interactions genes	between samples

Gene name	Chromosome	Description	Discovery sample	Replication sample	Pictures/fMRI sample	AgeCoDe sample
RPS6KA2	6	Ribosomal S6 kinase 3	0.0005	0.0177	0.0011	0.0020
KCNQ3	8	Potassium channel subunit Kv7.3	0.0017	0.0044	0.0020	0.0048
DPYSL2	8	Collapsin response mediator protein 2	0.0059	0.0062	0.0247	0.0009
ITGA9	3	Integrin alpha-9	0.0027	0.0008	0.0049	0.0858
ANK3	10	Ankyrin 3, node of Ranvier	0.0023	0.0197	0.0235	0.0013
CNTN6	3	Contactin 6	0.0009	0.0131	0.0110	0.0235
SH3GL2	9	Endophilin A1 BAR domain	0.0032	0.0031	0.0130	0.0578
SCN2A	2	Sodium channel, type II, α SUBUNIT	0.0024	0.0072	0.0676	0.0087
FGFR1	8	Fibroblast growth factor receptor 1	0.0125	0.0018	0.0185	0.0260
NRCAM	7	Neuronal cell adhesion molecule	0.0004	0.0409	0.0149	0.0578
ITGA1	5	Integrin, alpha 1	0.0079	0.0345	0.0267	0.0024
LAMA1	18	Laminin, alpha 1	0.0310	0.0879	0.0318	0.0003
EGFR	7	Epidermal growth factor receptor	0.0011	0.0328	0.0281	0.0459
NCAM1	11	Neural cell adhesion molecule 1	0.0144	0.0037	0.0352	0.0840
SPTBN1	2	Beta-II spectrin	0.0273	0.0045	0.1314	0.0162
CSNK2A2	16	Casein kinase II subunit alpha	0.0059	0.0807	0.0318	0.0190
ANK1	8	Ankyrin 1, erythrocytic	0.0152	0.0114	0.0254	0.0788
ITGA2	5	Integrin, alpha 2	0.0161	0.0312	0.0352	0.0247
CNTN1	12	Contactin 1/F3	0.0140	0.0290	0.0632	0.0235
NRP2	2	Neuropilin 2	0.0266	0.0181	0.0335	0.0504
NRP1	10	Neuropilin 1	0.0240	0.0275	0.0407	0.0823

Only genes with significant SNPs in the discovery sample and at least two other samples are shown (the SNP rsIDs and linkage disequilibrium between them are provided in Table S6). P values of association with relevant phenotypes (Table 5) are provided. Genes are sorted by the product of these values.

keywords automatically extracted from the literature; 61) showed that peak coordinates of the largest cluster dependent on the L1CAM interactions genetic score (-3, 28, 40) were related mostly to the term "response conflict" (Z-score = 5.03, posterior probability = 0.82). Thus, one explanation for the negative correlation between the differences in the activity in these regions and the genetic score is that these frontal activity patterns reflect increased effort or conflict between different options in the case of poorer recognition performance and less effort/conflict in the case of good memory. The negative correlation between the L1CAM interactions genetic score and the differences in activity in the left superior and inferior frontal gyri also may indicate that the reported regions code specifically for familiarity rather than recollection of pictures, as suggested by dual-process models (62, 63). However, because of the ceiling effect in our picture recognition data (with less than 4% of previously seen pictures rated as new and more than 80% as recollected), the percentages of correctly recollected pictures were strongly inversely correlated with percentages of previously seen pictures rated as familiar $(\rho_{\text{Spearman}} = -0.96)$. Therefore, based on our data, it was impossible to dissociate recollection and familiarity effectively at either the behavioral or neural level (where contrasts based on familiar pictures were significantly underpowered), which remains an important question for future studies.

In conclusion, by computationally dissecting episodic memory performance into specific components, we were able to identify distinct genetic profiles that underlie specific mental processes of human episodic memory.

Materials and Methods

Participants, Data Preprocessing, and GWAS Quality Control. We had three samples of healthy, young Swiss subjects: the discovery sample (used for associations with model parameters in the verbal task) consisted of the Zurich subsample (192 males, 514 females, age mean \pm SD = 21.92 \pm 2.95 y), and the Basel subsample (261 males, 504 females, age 22.47 \pm 3.62 y). The Basel words/pictures sample (200 males, 410 females, age 22.48 \pm 3.51 y) was used for the direct replication of associations with model parameters in the verbal task and for testing associations with performance in the picture task; the Basel pictures/fMRI sample (404 males, 588 females, age 22.47 \pm 3.35 y) was used for testing associations with performance in the picture task and studying their neural correlates with neuroimaging. Participants' physical and mental health was assessed by standard questionnaires and telephone interviews. Indications for exclusion were the recent use of medications influencing the central nervous system, pregnancy, visual system disorders, hormonal irregularities, a body mass index outside the 18.0-35.0 range (20.0–35.0 for men), recent or regular use of cannabis, excessive use of alcohol, use of other psychoactive drugs, history of mental illness, cancer, brain surgery, and for the fMRI sample, the presence of metal implants and claustrophobia. Written informed consent was obtained from the subjects after a complete description of the study. The ethics committees of the Cantons of Zurich and Basel, Switzerland approved the study protocols.

A total of 930,856 SNPs were genotyped. For association testing, markers with call rate less than 0.95, with minor allele frequency less than 0.05, and with Hardy-Weinberg equilibrium P < 0.05 were excluded. Population stratification was assessed by analyzing all genomewide, array-based autosomal SNPs passing the quality-control (QC) criteria with EIGENSTRAT (64). PCA was applied to each population (Zurich words subsample, Basel words subsample, Basel words/pictures sample, and Basel pictures/fMRI sample) to reduce genetic variation to a few dimensions. For PCA, default parameters were used (i.e., definition of 10 principal components in five iterations; the outlier criterion was 6 SDs). EIGENSTRAT identified a number of individuals deviating from a large, genetically homogenous population cluster: 112 in the Zurich words subsample, 68 in the Basel words subsample, 55 in the Basel words/pictures sample, and 112 in the Basel pictures/fMRI sample. These outliers were removed from further analyses. Because learning and memory abilities deteriorate with age (65), we also excluded the outliers (individuals with age more than 2 SDs from the mean in each population) from further analyses: 18 in the Zurich words subsample, 44 in the Basel words subsample, 29 in the Basel words/pictures sample, and 47 in the Basel pictures/fMRI sample. After outliers based on population stratification and age were excluded, the following numbers of participants remained for the final analysis: 1,239 in the discovery sample (582 in the Zurich words subsample, 657 in the Basel words subsample), 526 in the Basel words/pictures sample (picture

data available for 493 subjects), and 835 in the Basel pictures/fMRI sample [picture recognition data available for 822 subjects and fMRI data for 795 subjects (for the remaining subjects the fMRI data were corrupted); working memory data were available for 825 subjects, and fMRI data were available for 797 subjects; picture recognition and working memory data were available for 815 subjects].

GWAS Statistics and GSEA. All GWAS were run under the assumption of an additive model and using the Spearman rank correlation test. Sex chromosomes were excluded from the analysis. The resulting P values of autosomal SNPs in the GWAS (one list per sample and phenotype) served as input for the GSEA, which was performed using MAGENTA (23). Briefly, the method first maps SNPs to genes (each SNP can be counted toward only one gene) and then assigns each gene an SNP association score (i.e., the maximum SNP P value within \pm 0 kb of the annotated gene; 26). By applying a stepwise multiple linear regression, the analysis is corrected for the following confounders: gene size, number of SNPs, number of independent SNPs, number of recombination hotspots, linkage disequilibrium, and genetic distance (measured in centimorgans). Last, a gene set enrichment-like statistical test is applied to determine whether a gene set is enriched for highly ranked P values compared with a gene set of identical size randomly drawn from the genome. Based on recommendations in the literature (23, 26), we used a 75th percentile cutoff to determine the significance of a gene, because that cutoff has the optimal power for weak genetic effects that are expected for complex, polygenic traits. The gene sets used were extracted and curated from the MSigDB v3.1 database (www.broadinstitute.org/gsea/msigdb), including gene sets from different online databases (KEGG, Gene Ontology, BioCarta, and Reactome). We used a gene set size of 20 to 200 genes to avoid both overly narrow and overly broad functional gene set categories (26), resulting in 1,411 gene sets to be analyzed. Furthermore, to reduce bias, genes mapping in the high linkage disequilibrium (LD) extended major histocompatibility complex were excluded from the respective gene sets (66). FDR multiple testing correction was used to correct for the number of aene sets.

Memory Testing-the Verbal Task. Subjects viewed six series of five semantically unrelated nouns presented at a rate of one word/s with the instruction to learn the words for immediate free recall after each series. The words were taken from the collections of Hager and Hasselhorn (67) and consisted of 10 neutral words such as "angle," 10 positive words such as "happiness," and 10 negative words such as "poverty." The order of words was pseudorandom, with each group of five words containing no more than three words per valence category. In addition, subjects underwent an unexpected delayed free-recall test of the learned words after ~5 min. The free recall of a word was considered successful only if it was spelled correctly or a with single-letter typo that did not make it become a different valid word. The relevant performance measures (PMs) are described in Table 2. In the Basel words/pictures sample, the picture task was performed more than 3 h after the end of the verbal task. Word arousal ratings were available in the Zurich sample (on a scale of 1-5), and the mean arousal ratings did not differ significantly between positive and negative words: arousal(positive) = 3.41 (mean) \pm 0.44 (SEM); arousal(negative) = 4.10 \pm 0.16; Student t test; P = 0.16.

Computational Model for the Verbal Task. To dissociate specific mental processes involved in learning and memory, we used a computational model to describe individual performance in the verbal memory task (21). Depending on how well individuals remember a word, they may or may not try to write it down in the free recall, and if they try, their recall may or may not be correct. The probability that the attempted recall is correct depends on memory strength m of each word as follows: $p_{correct}(m, s) =$ $1/(1 + \exp(-s(m - m_{50\%})))$, with sigmoidal steepness s and center of the sigmoid $m_{50\%}$ = 1. The decision to attempt the recall of weak memories depends on the participant's willingness to risk making errors. We modeled this decision-making aspect using the decision threshold $\boldsymbol{\beta},$ where recall was attempted for words with memory strength $m > \beta$ but not for those with $m < \beta$. During the encoding, initial memory strength for each word was assigned as $m = \alpha \times \varepsilon + N(0, \sigma)$, with α being the learning rate, ε the emotional modulation of memory strength (ϵ = ϵ_{neg} for negative words, $\varepsilon = \varepsilon_{pos}$ for positive words, $\varepsilon = 1$ for neutral words), and N(0, σ) the Gaussian noise with mean 0 and SD σ , reflecting randomness in learning different words. Because the memory strength of words that have been recalled correctly in the immediate recall is likely to increase through repetition (i.e., the participants writing them down), we multiplied the memory strength m of immediately recalled words by the repetition-based memory improvement c (c \geq 1). To model forgetting during the 5-min delay, we multiplied all memory strengths by the forgetting rate γ (γ <1). It is important to note that, because no external measure could distinguish the repetition-based memory improvement from the forgetting rate in our experiments, these two parameters were closely related and together reflected memory maintenance.

The described model had eight parameters: α , β , γ , ε_{pos} , ε_{neg} , s, c, and σ . Because our behavioral phenotype consisted of only eight partly correlated measures per individual, it was impossible to estimate all eight parameters reliably. Because the PCA indicated five substantial components (Fig. S1), we chose to keep five of the eight parameters in the model flexible (different between individuals) and the three remaining ones fixed (same for all individuals). The selection of the most appropriate model (i.e., which parameters would be flexible and which fixed) was performed empirically, based on the corresponding mean goodness-of-fit values. Such selection ensured that closely related parameters, such as c and γ , would not both be flexible, whereas relatively independent parameters, which can account for more variance, would.

Estimation and Evaluation of Best-Fitting Model Parameters. To estimate the best-fitting model parameters, we computed the expected values of all performance measures (PM₁₋₈; Table 2) as a function of eight model parameters (α , β , γ , ϵ_{posr} , ϵ_{negr} , s, c, σ). We numerically computed integrals over probability distributions of memory strength *m*, because this method was more efficient and robust than simulating the model with random numbers and computing averages over multiple simulation runs. As a control, we also simulated the model stochastically: The averages of PMs over 100,000 simulations were almost exactly the same as those using the expected value based method.

To evaluate how well the model with a particular set of parameters fits individual behavioral performance, we used the following goodness-of-fit function (20, 21, 68):

$$\chi^{2} = \sum_{i=1}^{8} \frac{\left(PM_{i}^{exp} - PM_{i}^{mod}[parameters] \right)^{2}}{\left(\sigma_{i}^{exp}\right)^{2}}$$

where PM_i^{exp} and PM_i^{mod} are experimental and modeled performance measures of that individual, respectively, and $(\sigma_i^{exp})^2$ is the variance in the experimental data of PM_i .

With χ^2 as the objective function to minimize, we performed the estimation of best-fitting parameters in several stages (21):

- i) Model selection: To determine which five parameters should be flexible, we evaluated all 56 possible five-of-eight combinations. Because of the high computational cost of running 56 full estimation procedures, at this stage we performed only a moderately accurate estimation of the three fixed parameters.
- *ii*) Using the two best models, we performed a more refined estimation of fixed parameters, thereby improving the χ^2 values. We note that although the improvements in χ^2 values were substantial, they were small compared with the differences between the initial χ^2 values of the two best models and those of the other, worse models; therefore, it is very unlikely that any of those other models would become comparatively better with refinement.
- iii) For the final refinement, we evaluated the averages of all $2^{10} 1 = 1,023$ combinations of the 10 best parameter sets for each model, thereby further improving the χ^2 values. Finally, parameter sets from the model with the best goodness-of-fit were used for GWAS.

In all parameter estimation steps the search was performed in the following ranges: $(\alpha, \beta, \varepsilon_{\text{pesr}}, \varepsilon_{\text{negr}}, \sigma) \in (0.3, 3.5), c \in (1, 4.2), \gamma \in (0, 0.8), and s \in (0, 16)$. In choosing the ranges we balanced two partially opposing aims: to keep them as similar as possible to avoid bias to the estimation results and to keep them as close as possible to the likely distribution of each parameter to maximize estimation accuracy. The default range of 0.3–3.5 ensured that fewer than 1% of estimated parameter values were near the boundaries. Other ranges were used either because of fundamental constraints ($c \ge 1$ and $\gamma < 1$) or because the likely spread of parameter values was very different from the default. To evaluate how well the model fits individual data, we used the χ^2 test with $\nu = 8 - 5 =$ three degrees of freedom (five flexible parameters and eight PMs). Values of P > 0.05 indicate no statistical difference between modeled and observed PMs, meaning that the model fits the data well.

The Picture Task and Controls. After receiving general information about the study and giving their informed consent, participants were instructed and

trained on the picture task as well as the N-back working memory task. They performed the encoding of pictures for 20 min and then performed the N-back task for 10 min.

In the picture task, stimuli consisted of 72 pictures that were selected from the International Affective Picture System (IAPS; 69) and from in-house standardized picture sets that allowed us to equate the pictures for visual complexity and content (e.g., human presence). Participants were instructed to view the pictures passively and subsequently rate them according to emotional valence/arousal. Pictures received from IAPS were classified according to the IAPS valence rating; the remaining eight neutral pictures were rated based on an in-house valence rating. Positive stimuli were selected initially to match the arousal ratings of negative stimuli based on data from a pilot study in 20 subjects. Examples of pictures are as follows: erotica, sports, and appealing animals for the positive valence; bodily injury, snake. attack scenes for the negative valence; and neutral faces, household objects, and buildings for the neutral condition. On the basis of normative valence scores (from 1 to 9), pictures were assigned to the emotionally negative (2.3 \pm 0.6), emotionally neutral (5.0 \pm 0.3), and emotionally positive (7.6 \pm 0.4) conditions, resulting in 24 pictures for each emotional valence. Four additional pictures showing neutral objects were used to control for primacy and recency effects in memory. Two of these pictures were presented in the beginning and two at the end of the picture task. They were not included in the analysis. In addition, 24 scrambled pictures were used. The background of the scrambled pictures contained the color information of all pictures used in the experiment (except primacy and recency pictures), overlaid with a crystal and distortion filter (Adobe Photoshop CS3). In the foreground, a mostly transparent geometrical object (rectangle or ellipse of different sizes and orientations) was shown. The pictures were presented for 2.5 s in a quasi-randomized order so that no more than four pictures in the same category occurred consecutively. A fixation-cross appeared on the screen for 500 ms before each picture presentation. The stimulus onset time was jittered within 3 s [1 repetition time (TR)] per valence category with regard to the scan onset. After a picture was presented, participants subjectively rated the picture according to valence (negative, neutral, positive) and arousal (large, medium, small) on a three-point scale (self assessment manikin) by pressing a button with a finger of their dominant hand. For scrambled pictures, participants rated the form (vertical, symmetric or horizontal) and size (large, medium, small) of the geometrical object in the foreground. A maximum 6 s was allowed for rating each picture. Participants were not told that they should remember the pictures for later recall.

The participants also performed the 0- and 2-back versions of the N-back task (70). The task consisted of 12 blocks (six 0-back, six 2-back) in which 14 test stimuli (letters) were presented. The 0-back condition required participants to respond to the occurrence of the letter "x" in a sequence of letters (e.g., N-I-X-g ...), and the 2-back condition required subjects to compare the currently presented letter with the penultimate letter to decide whether they are identical or not (e.g., S-f-s-g ...) Performance was recorded as the number of correct responses, and the difference between the 2-back and the 0-back conditions served as a control phenotype of working memory.

After finishing the tasks, participants performed free recall of the pictures, which required them to write down a short description (a few words) of the previously seen pictures. Remembered primacy and recency pictures and training pictures were excluded from the analysis. No time limit was set for this task. Two trained investigators independently rated the descriptions for recall success (interrater reliability >99%). Depending on the computational model parameter with which each gene set was associated, different behavioral phenotypes were used. For the learning rate α , the relevant phenotype was the percentage of pictures remembered in the free recall (Table S1). For the negative emotional modulation parameter ε_{neg} , the relevant phenotype was the percentage of negative pictures minus the percentage of neutral pictures remembered in the free recall.

Multilocus Genetic Scores. To capture the multiallelic effect of gene sets associated with different model parameters, we generated individual multilocus genetic scores for each of these gene sets. The score comprised SNPs from an equal number of genes (i.e., one most significant SNP per gene) that were associated with the relevant phenotype of the picture task (or the control N-back task) in the pictures/fMRI sample. The score was computed by summing up the individual number of reference alleles over all of these SNPs, weighted by the direction of effect on the phenotype) or "-1" (the reference allele positively correlated with the phenotype). Each missing SNP was assigned the average value of that SNP over all participants of the sample for whom it was not missing. There were 0.41% of SNPs missing for L1CAM interactions, and 0.69% were missing for amine compound SLC transporters. To test the

Construction of a Population-Average Anatomical Probabilistic Atlas. Automatic segmentation of 1,000 subjects' T1-weighted images, including the participants from the current study, was used to build a population-average probabilistic anatomical atlas. More precisely, each participant's T1-weighted image first was automatically segmented into cortical and subcortical structures using FreeSurfer (version 4.5, surfer.nmr.mgh.harvard.edu; 71). Labeling of the cortical gyri was based on the Desikan–Killiany Atlas (72), yielding 35 regions per hemisphere.

The segmented T1 image then was normalized to the study-specific anatomical template space using the subject's previously computed warp field and was affine-registered to the MNI space. Nearest-neighbor interpolation was applied to preserve labeling of the different structures. The normalized segmentations finally were averaged across subjects to create a populationaverage probabilistic atlas. Each voxel of the template consequently could be assigned a probability of belonging to a given anatomical structure, based on the individual information from 1,000 subjects.

Functional Neuroimaging and Picture Recognition. Participants of the Basel pictures/fMRI sample performed the picture and N-back tasks in the scanner. They were positioned in the scanner after training. The participants received earplugs and headphones to reduce scanner noise. Their heads were fixated in the coil using small cushions, and they were told not to move their heads. Functional MRIs were acquired during the performance of the picture task for ~20 min, followed by the 10-min N-back task. After that task, participants were taken from the scanner to a different room for free recall of the presented pictures. Pictures were presented in the scanner using MR-compatible LCD goggles (Visuastim XGA; Resonance Technology). Eye correction was used when necessary.

Approximately 80 min after the presentation of the last picture during encoding, participants were repositioned in the scanner and performed a recognition task for 20 min. The recognition task consisted of two sets of stimuli that were either new (i.e., not presented before) or old (i.e., presented during the picture encoding task). Each of the two sets contained 72 pictures (24 pictures of each valence). During recognition, the pictures were presented for 1 s in a quasi-randomized order so that no more than four pictures of the same category appeared consecutively. A fixation cross appeared on the screen for 500 ms before each picture presentation. The stimulus onset time was jittered within 3 s (1 TR) per valence and previously seen/new category with regard to the scan onset. After picture presentation, participants subjectively rated the picture as recollected, familiar, or new by pressing a button with a finger of their dominant hand. Picture rating was possible in a time window of maximum 3 s.

Depending on the computational model parameter with which each gene set was found to be associated, different fMRI contrasts were used. For the learning rate α , we used two encoding contrasts: pictures of all valence categories vs. scrambled controls and subsequently remembered pictures (of all valence categories) vs. subsequently not-remembered pictures; the latter contrast is known as difference due to memory (Dm; 29). For the repetitionbased memory improvement c, which reflects memory maintenance, the relevant behavioral phenotype was the percentage of recollected pictures that had been seen previously, and the appropriate fMRI contrast was previously seen pictures vs. new pictures during recognition. Recognition data were used for this parameter because free recall was not performed in the scanner, and memory maintenance processes may not be reflected during the encoding. To ensure that the effects observed using contrast between the previously seen vs. new pictures reflected the episodic memory component of previously seen, not new, pictures, we also used the contrast of previously seen pictures that were recollected vs. those rated as familiar or new. To investigate neural correlates of working memory, the 2-back vs. 0-back contrast was used.

The contrasts described above were calculated individually using a fixedeffects model (first-level analysis). Because we were using such contrasts, we controlled for all factors that were constant between the two conditions, whether of neuronal or nonneuronal origin. The resulting contrast parameters then were used for genotype-dependent analyses in a random effects model (second-level analysis). Specifically, we used a regression model to analyze gene-dose-dependent differences in brain activity, with the multilocus genetic score, based on the related behavioral phenotype, as covariate. We controlled for the effect of sex by including it as a covariate. The significance threshold for score-dependent analysis was set at P < 0.05, FWE-corrected for multiple comparisons in the whole brain. We also performed second-level analysis with the related behavioral phenotype itself as a covariate.

Mediation Analysis. To test whether the observed effect of the L1CAM interactions gene set on behavioral performance was mediated by the activation of specific brain regions, we ran a whole-brain mediation analysis using the Multilevel Mediation and Moderation toolbox (wagerlab.colorado.edu/ wiki/doku.php/help/mediation/m3_mediation_fmri_toolbox; 73). We looked for brain regions whose activation during recognition (the contrast between previously seen vs. new pictures) mediated the association between the multilocus genetic score (called the "predictor") and recognition performance. To be considered as mediator, a voxel had to satisfy the following criteria: the predictor variable must be related to the mediator (path a); the mediator must be directly related to behavior, controlling for the predictor (path b); the mediation effect must be significant (effect a*b), meaning that the predictor-behavior relationship is reduced significantly by including the mediator in the model. Bootstrapping was used to obtain corresponding P values, and an FDR threshold of q < 0.05 across all effects of interest (a, b, and a*b coefficients) was applied to correct for multiple comparisons. Furthermore, sex was included as covariate.

Because causality between correlated brain activity and behavior cannot be inferred, we also tested using the same procedure to determine whether the association between the multilocus genetic score and differences in brain activity during recognition (the contrast between previously seen vs. new pictures) was mediated by picture recognition.

Sample from the German AgeCoDe Study. The sample consisted of elderly participants in the German AgeCoDe study, an ongoing primary care-based prospective longitudinal study on the early detection of mild cognitive impairment and dementia established by the German Competence Network Dementia. The sampling frame and sample selection process of the AgeCoDe study have been described in detail previously (74). After the application of the selection criteria (SI Materials and Methods), a total of 1,244 subjects remained in the sample. Sufficient DNA samples for genomewide genotyping were available for 782 subjects. Additionally, 39 subjects were excluded because of sex-check inconsistencies, low call rate, or extreme values of heterozygosity rate, which may indicate a genotyping bias. The final sample comprised 743 subjects (241 males and 502 females; mean age: 79.53 \pm 3.21 y). As part of the structured clinical interviews, subjects were presented a list of 10 words from the German version of the CERAD Word List Learning Task (28) three times (presentation per word: 2 s); each trial was presented in a different order. Participants were asked to read each word aloud as it was presented. After each trial, subjects were asked to recall freely as many words as possible. The number of correctly remembered items over the three trials (immediate recall) served as a relevant phenotype for the gene set associated with the learning rate α , whereas the number of correctly recalled items after a 10-min delay (filled with other questions of the structured interview) was relevant for the gene set associated with the repetitionbased memory improvement c.

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