Human spatial memory implicitly prioritizes high-calorie foods

Rachelle de Vries1,2,4*, Paulina Morquecho-Campos1,4, Emely de Vet2, Marielle de Rijk1, Elbrich Postma3, Kees de Graaf2, Bas Engel3 & Sanne Boesveldt1

All species face the important adaptive problem of efficiently locating high-quality nutritional resources. We explored whether human spatial cognition is enhanced for high-calorie foods, in a large multisensory experiment that covertly tested the location memory of people who navigated a maze-like food setting. We found that individuals incidentally learned and more accurately recalled locations of high-calorie foods – regardless of explicit hedonic valuations or personal familiarity with foods. In addition, the high-calorie bias in human spatial memory already became evident within a limited sensory environment, where solely odor information was available. These results suggest that human minds continue to house a cognitive system optimized for energy-efficient foraging within erratic food habitats of the past, and highlight the often underestimated capabilities of the human olfactory sense.

A recurring fitness-relevant task faced by all species is the efficient pursuit of nutritional resources1. A central theorem of optimal foraging theory is that an individual's fitness is a direct function of the efficiency with which one acquires energy, and natural selection pressures favour foraging traits that maximize the net rate of energy gain1,2. Although this theory has been extensively referenced in relation to the foraging strategies of other animals2, the question of whether humans also inherently carry adaptations geared toward energy-efficient foraging has not been thoroughly assessed to date.

For about 99 percent of human evolution, our ancestors were hunter-gatherers inhabiting a highly complex and variable physical food environment, where food sources varied on both spatial and temporal availabilities3,4. A cognitive adaptation that could have evolved to optimize foraging efforts within such erratic food habitats of the past is a high-calorie bias in spatial memory5,6. Such an inbuilt spatial bias entails the automatic registration and prioritization in memory of high-calorie food locations. This would have enabled foragers to efficiently navigate toward valuable calorie-dense resources – without competing for limited attentional capacities required in other important activities such as avoiding predation4,6. Indeed, a similar mechanism has been observed in other animal species7–9. Using an innovative and ecologically valid experimental set-up that covertly tested the food location memory of more than 500 individuals, we provide first-hand evidence that human spatial processing is implicitly biased toward high-calorie foods.

To mirror real-world navigation within a heterogeneous food environment as closely as possible, we created a maze-like setting where participants followed a specific route within a room to sample an assortment of (sweet and savory) high- and low-calorie food stimuli at dispersed pillar locations (Fig. 1). We emulated two sensory environments in separate rooms, each of which engaged sensory modalities fundamental to the processes of spatial navigation and eating behavior10–12: In the multisensory environment (i.e. vision + taste + olfaction), stimuli consisted of actual food products that individuals had to eat, whereas individuals were instructed to only smell food odors in the olfactory environment. Importantly, participants were not informed that their (spatial) memory would be tested afterwards, to ensure that the encoding of food locations would be purely incidental. We then compared performance, expressed as the proportion of correct food-to-pillar relocations in a surprise spatial memory task, for high-calorie versus low-calorie food stimuli in both sensory environments.

1Division of Human Nutrition and Health, Wageningen University and Research, P.O. Box 17, 6700 AA Wageningen, The Netherlands. 2Consumption and Healthy Lifestyles, Wageningen University and Research, Wageningen, The Netherlands. 3Mathematical and Statistical Methods (Biometris), Wageningen University and Research, Wageningen, The Netherlands. 4These authors contributed equally: Rachelle de Vries and Paulina Morquecho-Campos. *email: rachelle.devries@wur.nl
Results

Human spatial memory automatically prioritizes high-calorie food. In the multisensory environment, individuals relocated high-calorie foods to correct pillar locations significantly more frequently than low-calorie alternatives (High-calorie: $M = 0.63$, 95% CI = [0.58,0.67]; Low-calorie: $M = 0.57$, 95% CI = [0.52,0.62]), $\chi^2 (1) = 9.35$, $p = 0.002$, OR = 1.27, 95% CI = [1.09, 1.48] (Fig. 2). This effect occurred regardless of demographics, relevant state characteristics (e.g. hunger and alertness), hedonic evaluations of foods (i.e. liking and desirability ratings; Fig. 3), and familiarity with foods. Similarly, individuals in the olfactory environment more frequently relocated odors signaling high-calorie foods to correct pillar locations relative to low-calorie odor counterparts (High-calorie: $M = 0.36$, 95% CI = [0.33,0.39]; Low-calorie: $M = 0.30$, 95% CI = [0.27,0.34]), $\chi^2 (1) = 6.88$, $p = 0.009$, OR = 1.28, 95% CI = [1.06, 1.54] (Fig. 2), while controlling for the same set of potential confounders – although the likelihood of a correct relocation increased with a greater familiarity with an odor stimulus, $\chi^2$. 

Figure 1. Heterogeneous food environment. Example of the spatial distribution of food stimuli and navigation route within the maze-like experimental setting.
Figure 2. Food spatial memory accuracy. Human spatial memory for high-calorie and low-calorie food stimuli in two sensory environments, expressed as the proportion of correct food-to-pillar relocations. Error bars represent 95% confidence intervals.

Figure 3. Food ratings across sensory environments. Liking (a), Desirability (b), and Familiarity (c) ratings (on a 100 mm Visual Analogue Scale) for all food stimuli in the multisensory and olfactory environment. Error bars represent 95% confidence intervals.
The high-calorie bias in human spatial memory manifests with limited sensory information. In a combined analysis of both sensory conditions, a better overall food relocation performance was observed in the multisensory compared to the olfactory environment (Multisensory: $M = 0.58, 95\% CI = [0.54,0.61]$; Olfactory: $M = 0.36, 95\% CI = [0.33,0.39]$). The chi-squared test ($\chi^2 (1) = 62.95, p < 0.001, OR = 2.43, 95\% CI = [1.95,3.03]$) indicated that the high-calorie spatial memory bias was equally expressed in both sensory environments – even where solely odor information was available.

Discussion
In a naturalistic multisensory experiment, individuals incidentally learned and more accurately recalled locations of high-calorie food stimuli. These results are compatible with the notion of "adaptive memory", which contends that memory systems – much like other biological systems – were shaped by the forces of natural selection and should therefore show sensitivity to fitness-relevant content\(^{13,14}\). Indeed, alternative interpretations of our findings that are grounded in more traditional memory frameworks, which champion the primacy of content-insensitive general learning mechanisms, can be ruled out by our data\(^{19}\). The possibility that the high-calorie spatial memory bias resulted from a greater "depth" of processing or motivational salience of high-calorie stimuli is minimal, given that we controlled for an individual's personal familiarity with a food, as well as their explicit liking and desire to consume an item\(^{25}\). In addition, high- and low-calorie food products were equivalent in their composition of important macronutrients (i.e. protein to carbohydrate and fat ratios), rendering it unlikely that differences in nutritional balance – rather than caloric content – is what drove the mnemonic advantage in the high-calorie condition\(^{16}\). However, the observation that (odor) familiarity predicted a higher frequency of overeating\(^{16}\) highlights the importance of considering both content-sensitive and content-insensitive learning processes for human spatial cognition\(^{2}\).

Remarkably, the expression of the high-calorie bias in human spatial memory required only a limited presence of sensory information – granted that available sensory cues (such as odors) can communicate the relative value (e.g. caloric content) of potential foods – which further speaks to the processing efficiency of the mechanism\(^{1,17}\). We speculate that this could be due to an overlap in underlying (hippocampal) neural coding processes, despite variations in the (dominant) sensory modality used to explore the external world and significant objects contained within them\(^{19}\). For instance, it is feasible that hippocampal place cells show enhanced activity during recognition of objects (or cues) that flag a high-priority resource, independently of the type of sensory input received\(^{18}\). However, a sizeable difference in overall spatial memory performance was evident between sensory conditions, which may have resulted from a greater variety of sensory information present in the multisensory environment. Individuals in the multisensory environment had a wider availability of sensory modalities (e.g. visual information) to utilize as spatial cues during encoding, which could have yielded a richer construction of mental spatial representations\(^{9,20}\). Going forward, research efforts would benefit from additionally documenting or matching participant samples on individual abilities to mentally represent and flexibly manipulate spatial information (i.e. between the viewer-centered perspective during navigation and the aerial map perspective during spatial recall)\(^{21}\), for a more refined comparison of (food) location memory between sensory conditions.

In turn, differences in the expression of the high-calorie spatial memory bias may offer a novel explanation for why some individuals are less successful in maintaining a healthy energy balance within the modern food landscape\(^{22}\). An enhanced memory for high-calorie food locations could make high-calorie options relatively easier to obtain within a diverse food environment, especially for those with a greater expression of the bias\(^{22}\). In this manner, the cognitive bias may facilitate high-calorie food choice, by capitalizing on the tendency of individuals to prefer convenient easily-accessible items when making food decisions\(^{25}\). Similarly, it could stimulate individuals to visit calorie-laden food locations (e.g. fast food outlets) on a wider scale of space. Given the paucity of literature on the high-calorie spatial memory bias and its potential behavioral effects, further investigation is merited on what other cognitive processes are associated with the bias, and how it may influence the manner in which people navigate contemporary food replete settings.

Finally, our findings add to a growing literature that highlight the relevance of olfaction for eating behavior in humans, which is known to be the case across other species\(^{1,12}\). The human sense of smell is often depicted to be inferior to those of other mammals, such as dogs or rodents\(^{24}\). However, our observations showcase the intact ability of individuals to distinguish different odor types, deduce caloric properties of signaled foods from odor cues, and localize odor objects in space\(^{1,17,25}\). Indeed, a well-developed olfactory sense is thought to have conferred a survival advantage to (ancestral) hunter-gatherers\(^{26,27}\).

Taken together, we find that human minds may continue to house an implicit cognitive system optimized for energy-efficient foraging within the fluctuating ancestral food environments in which memory evolved.

Materials and methods
Participants. This experiment was part of the three-day Lowlands Science 2018 festival program (the Netherlands). A total of 512 attendees were analyzed: 258 participants (47% female; $M_{\text{age}} = 28.2$ years, $SD = 9.1$; $M_{\text{BMI}} = 24.0$ kg/m$^2$, $SD = 3.6$) in the multisensory environment and 254 participants (50% female; $M_{\text{age}} = 28.5$ years, $SD = 9.0$, $M_{\text{BMI}} = 23.8$ kg/m$^2$, $SD = 3.4$) in the olfactory environment. Data from 539 individuals were initially collected, but 21 files contained missing values and 6 files originated from individuals who participated in both sensory conditions which was an exclusion criterion. All participants (and/or their legal guardians) provided
written informed consent prior to testing. This study was approved by the Social Sciences Ethics Committee of Wageningen University and was performed in accordance with relevant ethical guidelines and regulations. The hypothesis, full research protocol and analysis plan were preregistered, and can be accessed alongside reported data at https://osf.io/2rwmt/.

Spatial memory task. Participants were brought to a starting point within a room (area of 12 m²). They navigated between eight pillars at a fixed pre-determined order that was indicated by arrow signs on the floor. Although navigation schemes remained constant, the assignment of food stimuli to pillar locations (i.e. encoding order of caloric density—taste conditions) was randomized every hour and pillar frequencies did not differ between conditions. Participants tasted (or smelled) and provided ratings (i.e. liking, desire to eat, familiarity; Fig. 3) on a food stimulus at all pillars. Participants then completed a surprise spatial memory task in a separate area. During recall, participants were randomly presented with a sequence of previous food stimuli and had to indicate the pillar location of each item on a (two-dimensional) digital map of the relevant room. The total number of possible pillar locations (N = 8) was displayed anew each recall round, and a pillar location could be selected more than once.

Food stimuli. Four high-calorie (M = 498.5 kcal/100 g, SD = 35.8) and low-calorie (M = 34.3 kcal/100 g, SD = 18.9) food products and odor equivalents were used, with an equal number of sweet (e.g. High-calorie: chocolate brownie; Low-calorie: apple) and savory (e.g. High-calorie: potato chip; Low-calorie: cherry tomato) options for each. Food odors were matched on perceived intensity (i.e. 55–75 mm on a 100 mm Visual Analogue Scale) between caloric density—taste conditions and validated in previous research. Food products were placed in bowls and refilled at regular time intervals to maintain a consistent presentation volume. Food odors were presented in (screw-capped) brown bottles (50 ml) containing scented cotton pads, which participants had placed in bowls and refilled at regular time intervals to maintain a consistent presentation volume. Food odors were also replaced regularly to uphold the desired odor intensity.

All food stimuli were placed atop pillars and covered by identical cloths that participants had to open during navigation.

Statistical analysis. For data from each sensory environment, a generalized linear mixed model (GLMM) with a random slope was formulated. A GLMM was chosen to flexibly model for correlated errors in the (non-normal) binary outcome variable, and linearity of covariates (on the logit scale) was shown to sufficiently capture their effects. The GLMM comprised fixed main and interaction effects for experimental factors Caloric Density and Taste, and random effects for the factor Participant. All effects were introduced on the logit scale. Additionally, in the fixed part of the model and also on the logit scale, Gender, Age (in tertiles), Subjective SES, Food Allergies, Hunger ratings, hours of Sleep, Alertness, Alcohol consumption, Drug use, Smoking, Liking, Desirability, and Familiarity were entered as covariates. Binary observations, conditional upon the random effects for participants, were assumed to follow a Bernoulli distribution. To test whether the type of sensory environment (i.e. multisensory versus olfactory) moderates food spatial memory accuracy and expression of the high-calorie bias, observations from both sensory rooms were combined into a single analysis, adding fixed main and interaction effects (e.g. with Caloric Density of Sensory Environment) to the GLMM. Ordinary likelihood ratio tests (using the -2LL test statistic) were used for testing, with p values derived from an approximation with the chi-square distribution. Inference was based on Laplacian integration employing the lme4 package from R. Detailed information on the measurement of covariates and the model selection process can be found at https://osf.io/2rwmt/.

Data availability
The data that support the findings of this study are available on the Open Science Framework repository with the identifier https://doi.org/10.17605/OSF.IO/2RWMT. Received: 17 April 2020; Accepted: 2 September 2020
Published online: 08 October 2020

References
1. Schoener, T. W. Theory of feeding strategies. Annu. Rev. Ecol. Evol. Syst. 2, 369–404 (1971).
2. Pyke, G. H., Pulliam, H. R. & Charnov, E. L. Optimal foraging: a selective review of theory and tests. Q. Rev. Biol. 52, 137–154 (1977).
3. Winterhalder, B. Optimal foraging strategies and hunter-gatherer research in anthropology: theory and models. In Hunter-Gatherer Foraging Strategies: Ethnographic and Archaeological Analyses (ed. Smith, E. A.) 13–35 (University of Chicago Press, Chicago, 1981).
4. New, J., Krasnow, M. M., Truxaw, D. & Gaulin, S. J. Spatial adaptations for plant foraging: women excel and calories count. Proc. R. Soc. B 274, 2679–2684 (2007).
5. de Vries, R., de Vet, E., de Graaf, K. & Boesveldt, S. Foraging minds in modern environments: high-calorie and savory-taste biases in human food spatial memory. Appetite 152, 104718 (2020).
6. Krasnow, M. M. et al. Cognitive adaptations for gathering-related navigation in humans. Evol. Hum. Behav. 32, 1–12 (2011).
7. Cunningham, E. & Janson, C. Integrating information about location and value of resources by white-faced saki monkeys (Pithecia pithecia). Anim. Cogn. 10, 283–304 (2007).
8. Janson, C. H. Experimental evidence for spatial memory in foraging wild capuchin monkeys, Cebus apella. Anim. Behav. 55, 1229–1243 (1998).
9. Janmaat, K. R., Polansky, L., Ban, S. D. & Boesch, C. Wild chimpanzees plan their breakfast time, type, and location. Proc. Natl. Acad. Sci. USA 111, 16343–16348 (2014).
10. Yarmolinsky, D. A., Zaker, C. S. & Ryba, N. J. Common sense about taste: from mammals to insects. Cell 139, 234–244 (2009).
11. Jacobs, L. F. From chemotaxis to the cognitive map: the function of olfaction. Proc. Natl. Acad. Sci. USA 109(Supplement 1), 10693–10700 (2012).
12. Boesveldt, S. & de Graaf, K. The differential role of smell and taste for eating behavior. Perception 46, 307–319 (2017).
13. Nairne, J. S. Adaptive memory: evolutionary constraints on remembering. In The Psychology of Learning and Motivation: Vol. 53 (ed. Ross, B. H.) 1–32 (Elsevier Academic Press, Amsterdam, 2010).
14. Nairne, J. S. & Pandeirada, J. N. Adaptive memory: remembering with a stone-age brain. JVI/V 11, 671–684 (1972).
15. Simpson, S. J. & Raubenheimer, D. Obesity: the protein leverage hypothesis. Obes. Rev. 6, 133–142 (2005).
16. Zoon, H. F., De Graaf, C. & Boesveldt, S. Food odours direct specific appetite. Foods 5, 12 (2016).
17. Schiller, D. et al. Memory and space: towards an understanding of the cognitive map. J. Neurosci. 35, 13904–13911 (2015).
18. Schifferstein, H. N., Smeets, M. A. & Postma, A. Comparing location memory for 4 sensory modalities. Chem. Senses. 35, 135–145 (2009).
19. Downs, R. M. & Stea, D. Cognitive maps and spatial behavior: process and products. In The Map Reader: Theories of Mapping Practice and Cartographic Representation (eds Dodge, M. et al.) 312–317 (Wiley, New York, 2011).
20. Wolbers, T. & Hegarty, M. What determines our navigational abilities? Trends. Cogn. Sci. 14, 138–146 (2010).
21. Allan, K. & Allan, J. L. An obesogenic bias in women's spatial memory for high calorie snack food. Appetite 67, 99–104 (2013).
22. Furst, T., Connors, M., Bisogni, C. A., Sobal, J. & Falk, L. W. Food choice: a conceptual model of the process. Appetite 26, 247–266 (1996).
23. McGann, J. P. Poor human olfaction is a 19th-century myth. Science 356, eaam7263 (2017).
24. Wu, Y., Chen, K., Ye, Y., Zhang, T. & Zhou, W. Humans navigate with stereo olfaction. Proc. Natl. Acad. Sci. USA 117, 16065–16071 (2020).
25. Bastir, M. et al. Evolution of the base of the brain in highly encephalized human species. Nat. Commun. 2, 588–595 (2011).
26. Majid, A. & Kruspe, N. Hunter-gatherer olfaction is special. Curr. Biol. 28, 409–413 (2018).
27. Bates, D. et al. Package ‘lme4’. Available at https://github.com/lme4/lme4/ (2019).
28. McGann, J. P. Poor human olfaction is a 19th-century myth. Science 356, eaam7263 (2017).
29. Wu, Y., Chen, K., Ye, Y., Zhang, T. & Zhou, W. Humans navigate with stereo olfaction. Proc. Natl. Acad. Sci. USA 117, 16065–16071 (2020).
30. Bastir, M. et al. Evolution of the base of the brain in highly encephalized human species. Nat. Commun. 2, 588–595 (2011).
31. Majid, A. & Kruspe, N. Hunter-gatherer olfaction is special. Curr. Biol. 28, 409–413 (2018).
32. Bates, D. et al. Package ‘lme4’. Available at https://github.com/lme4/lme4/ (2019).
33. McGann, J. P. Poor human olfaction is a 19th-century myth. Science 356, eaam7263 (2017).
34. Wu, Y., Chen, K., Ye, Y., Zhang, T. & Zhou, W. Humans navigate with stereo olfaction. Proc. Natl. Acad. Sci. USA 117, 16065–16071 (2020).
35. Bastir, M. et al. Evolution of the base of the brain in highly encephalized human species. Nat. Commun. 2, 588–595 (2011).
36. Majid, A. & Kruspe, N. Hunter-gatherer olfaction is special. Curr. Biol. 28, 409–413 (2018).
37. Bates, D. et al. Package ‘lme4’. Available at https://github.com/lme4/lme4/ (2019).
38. McGann, J. P. Poor human olfaction is a 19th-century myth. Science 356, eaam7263 (2017).
39. Wu, Y., Chen, K., Ye, Y., Zhang, T. & Zhou, W. Humans navigate with stereo olfaction. Proc. Natl. Acad. Sci. USA 117, 16065–16071 (2020).
40. Bastir, M. et al. Evolution of the base of the brain in highly encephalized human species. Nat. Commun. 2, 588–595 (2011).
41. Majid, A. & Kruspe, N. Hunter-gatherer olfaction is special. Curr. Biol. 28, 409–413 (2018).
42. Bates, D. et al. Package ‘lme4’. Available at https://github.com/lme4/lme4/ (2019).
43. McGann, J. P. Poor human olfaction is a 19th-century myth. Science 356, eaam7263 (2017).