Collective decision-making by rational agents with differing preferences

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Collective decisions can emerge from individual-level interactions between members of a group. These interactions are often seen as social feedback rules, whereby individuals copy the decisions they observe others making, creating a coherent group decision. The benefit of these behavioral rules to the individual agent can be understood as a transfer of information, whereby a focal individual learns about the world by gaining access to the information possessed by others. Previous studies have analyzed this exchange of information by assuming that all agents share common goals. While differences in information and differences in preferences have often been conflated, little is known about how differences between agents’ underlying preferences affect the use and efficacy of social information. In this paper, I develop a model of social information use by rational agents with differing preferences, and demonstrate that the resulting collective behavior is strongly dependent on the structure of preference sharing within the group, as well as the quality of information in the environment. In particular, I show that strong social responses are expected by individuals that are habituated to noisy, uncertain environments where private information about the world is relatively weak. Furthermore, by investigating heterogeneous group structures, I demonstrate a potential influence of cryptic minority subgroups that may illuminate the empirical link between personality and leadership.

The choices made by others serve as a conduit for information about the world that other individuals possess, and therefore enable a focal individual to make decisions with greater expected utility or fitness than they could do alone. This transmission of information, frequently labeled as “social information,” has been posited as an important factor driving collective cohesion in animal and human groups (1–3), alongside other benefits of aggregation such as dilution of predation risk (4).

The tendency of individuals to follow the decisions of others has been comprehensively demonstrated across many taxa (e.g., insects (5), fish (6, 7), birds (8, 9), and mammals (10–12), including humans (13, 14)). Many studies have posited simple social feedback rules as models for collective decision-making, demonstrating how individual behavioral heuristics could generate cohesive collective decisions (15–17). Other studies have sought to reveal the form of these social interaction rules via a data-driven approach (5, 18). However, neither of these methods identify the source of the behavioral rules in terms of the direct benefit to the individual, an important evolutionary principle in unrelated groups (19). More recent work has investigated how social interaction rules could be justified from the rational self-interest of the decision-making agents, either through an evolutionary analysis (20) or by direct calculation (1–3).

Identifying the individual fitness or utility motivations of social interactions is important for understanding how behavior will vary with context, and thus how to extrapolate from empirical observations to make predictions about social behavior in other environments (3).

Studies that have sought to directly evaluate rational rules of social interaction (1–3) have addressed the fundamental question of social information: What can one individual learn by observing the choices made by others? The general approach to this question has centered on two conceptual points: 1) The choices of other agents are of interest because those agents have information that the focal individual may lack; and 2) the focal agent can only infer what that information might be by considering why the other agent made the choice it did. In considering why another agent may have made the choice they did, the focal agent must have a model for how that agent responds to different information. A simple model is to assume that the other agent is identical to oneself, and therefore responds identically to any set of information (3). However, groups are not, in general, composed of identical individuals, and agents may have different preferences based on their particular needs or tastes. This may be the case within same-species groups, as individuals can differ in, for example, age, genetic profile (21), or current nutritional needs. Even more clearly, groups may be composed of multiple species (8, 22), with potentially very different needs and goals between individuals from different species.

How then might agents make use of social information when their preferences differ? It is important here to distinguish between differences in preference and differences in information. Information and preference differences are frequently conflated in models inspired by physical systems, where both are typically expressed in terms of attractive forces or potentials (e.g., refs. 23–26), but, in reality, these are different sources of possible conflict, with quite different effects upon a rational agent. Consider two individuals, one of which is trained to...

Significance

For animals choosing where to forage or sleep, or for humans choosing a restaurant, a hotel, or a career, the choices made by others can provide useful information. If a restaurant is consistently full of customers, we might assume it must be serving good-quality food. However, the choices that others make are only useful to us if they are made by agents with similar preferences; a vegan can learn very little from the popularity of a steak house. In this paper, I show how differences in preferences change how agents should respond to each other to maximize their personal fitness or utility, and thus how population structures might determine the interaction rules that evolve in social groups.

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associate food with red markings, the other with blue markings. This may result in a difference in revealed color preference between them when both markings are displayed. However, this color preference is, from a rational perspective, a cipher for both individuals’ common preference for food. It is therefore better understood as a difference of information: The two individuals may differ in where they believe food to be located, but both still share a desire to find the food, and should therefore be interested in what the other knows. Conversely, two individuals with exactly the same information may make different choices if they have different goals, and be entirely uninterested in the choices each other makes—for example, a vegetarian and a committed carnivore choosing where to eat based on the same set of restaurant reviews. I have previously considered differences in information when agents have identical preferences (3), where agents can view each other as more or less direct proxies in terms of response to information. Differences in preferences matter because they change what one individual can learn from observing another, as the actions of those other agents may be different than what the focal agent would have done with the same information.

Here I develop a model for collective decision-making by rational agents with differing preferences, that seek to maximize differing utility functions (27). This is based on a model of utility structured as an array of environmental factors characterizing different choices, which can encode differing degrees of correlation between individual preferences. Using this framework of utility functions, I then derive the rational decision-making rules for individuals to follow, assuming that individuals have common acquired knowledge about the expected range of environmental characteristics and the typical degree to which their preferences align. Finally, I derive the expected behavior of the individuals and the group when faced with a binary decision between two alternatives, A and B, which can represent different foraging patches, different movement directions, or any other mutually exclusive activities. Following ref. 3, I evaluate the expected behavior from the perspective of an external observer (as opposed to the agents themselves), and thus make predictions about the likely characteristics of collective decision-making seen in empirical studies and in real human and animal groups.

Results

In this section, I explore the decision-making of rational agents under a variety of different environmental and social contexts, by varying several key parameters of the model described in Materials and Methods: 1) the habitual environmental noise level (\(\epsilon\)). This is the noise to signal ratio in the environment to which the agent is habituated, and thus represents the agent’s prior belief regarding the reliability of environmental information—note that all agents are assumed to share the same belief about this environmental noise level; 2) the experimental noise level, \(\eta\). This is the noise to signal ratio under which the agents are being observed, and will typically be lower than \(\epsilon\) in laboratory experiments; and 3) the magnitude and structure of the correlation between agent preferences \(\rho\). Agents with similar preferences have high values of \(\rho\), while strongly dissimilar preferences imply very low or even negative values of \(\rho\). All of these parameters are discussed in more detail in Materials and Methods.

**Differing Preferences Induce Environmental Dependence in Decision-Making.** I evaluated the probability that a focal individual will choose option A, under natural conditions where the experimental noise level (\(\eta\)) matches the habitual environmental noise level (\(\epsilon\)), conditioned on different sequences of previous decisions, and within a group where individual preferences have a characteristic correlation \(\rho\). I further calculated how these probabilities varied with the habitual environmental noise level \(\epsilon\). The results of these calculations are shown in Fig. 1, with each line corresponding to a different sequence of up to three previous decisions. Each line is labeled by the sequence of previous decisions by up to three other individuals, across a range of environmental noise levels, with (A) weakly aligned preferences (\(\rho = 0.5\)) or (B) strongly aligned preferences (\(\rho = 0.9\)).

![Fig. 1. The probability for a focal agent to choose option A, conditioned on possible sequences of previously observed decisions by up to three other individuals, across a range of environmental noise levels, with (A) weakly aligned preferences (\(\rho = 0.5\)) or (B) strongly aligned preferences (\(\rho = 0.9\)).](image-url)
in addition to this general trend: As the noise level $\epsilon$ is increased, there is a transition from weakly following the majority opinion to strongly following the most recent decisions in cases where the most recent decision conflicts with the majority. This can be seen clearly in the crossing of the lines representing the decision sequences AAB and BBA in Fig. 1B. A complementary effect can be observed for the sequences AAB, ABA and BAA: Here the sequence of previous decisions makes a substantial difference to the focal agent’s decision when the environmental noise level is high, but, when the environmental noise is low, all three cases converge to the same decision probability. These results should be compared with those of ref. 3, which correspond to the special case of $\rho = 1$. In that case, there is no dependence on environmental noise levels, and the most recent decision is always favored.

**Relative Social Weighting.** The theory described in Materials and Methods results in a complex, recursive decision-making rule. However, a simpler perspective on the importance of the model parameters can be obtained if one focuses only on the decision faced by the second individual, who must balance the social information provided by the first decision against their own private information. How are these sources of information weighted? One can see from Eq. 12 that the private information of agent 2 enters in the second term, weighted by a factor of $1/\epsilon$, while the social information is contained in the third term and weighted by $\rho/\sqrt{1 + \epsilon^2 - \rho^2}$. From the ratio of these two weightings, we can define the “relative social weighting” (RSW),

$$\text{RSW} = \rho \epsilon/\sqrt{1 + \epsilon^2 - \rho^2}. \quad [1]$$

This relationship contains several important special cases. When individuals share identical preferences ($\rho = 1$), the choices made by others are a direct proxy for the decision the focal agent would have made themselves, and thus the RSW is one. When preferences are not identical ($\rho < 1$), private information dominates in cases where that information is reliable ($\epsilon << 1$), and thus the RSW is close to zero. However, when private information is not reliable ($\epsilon >> 1$), the choices of others are weighted in proportion to how strongly preferences are aligned (RSW $\sim \rho$).

If there is no correlation between two individuals’ utility functions ($\rho = 0$), then the choices made by one convey no social information to the other, and the RSW is zero. If $\rho$ is negative, then agents have opposed preferences, and the resulting RSW is negative, implying that one agent will actively disfavor the option chosen by the other. Note that a homogeneous value of $\rho$ below zero is not possible in groups of more than two agents.

**Sequence Ordering in Previous Decisions.** I investigated further how preference alignment and environmental noise determine how a focal individual uses social information, focusing on the possible tension between a majority of previous decisions and the most recent of those decisions. To explore this tension, I consider a simple conflict: a sequence of previous decisions of the form BBA. That is, the focal individual is presented with two weightings, we can define the “relative social weighting” (RSW),

$$\text{RSW} = \rho \epsilon/\sqrt{1 + \epsilon^2 - \rho^2}. \quad [1]$$

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**RSW Drives Consensus.** Having determined that preference alignment and environmental noise have a strong effect on how a focal individual makes use of social information, we need to understand the consequences of this for collective behavior. Of particular interest is the degree to which groups can come to consensus decisions and remain cohesive.

Assuming a group of eight individuals making sequential choices according to the rational decision-making rule, I evaluated the probability for each possible collective outcome in the binary decision-making scenario, in terms of the eventual number of individuals selecting option $A$, $n_A$, and those choosing option $B$, $n_B$. This calculation considered the probability of every sequence of decisions that could give rise to a given collective

![Fig. 2. Resolving conflicts between majority decisions and recent decisions.](image)

(A) The probability that a focal individual will select option $A$, conditioned on observing three previous decisions of the form BBA, as a function of the utility function correlation and the environmental noise level, tested under natural conditions ($\eta/\epsilon = 1$). (B) The equivalent probabilities, assuming the focal individual is tested under low-noise laboratory conditions ($\eta/\epsilon = 1/2$). In both cases, the focal agent favors recent information when utility alignment is strong and environmental noise is high. (C) The results for the natural setting (black points) and the laboratory setting (red points) plotted against the RSW.
outcome and summed over these to give the final probabilities. Each of these collective outcomes is given a “consensus score,” defined as

$$\text{consensus} = \frac{|n_A - n_B|}{n_A + n_B}$$

such that a consensus score of one indicates that all individuals chose the same option, and a consensus score of zero indicates an equal split between A and B.

I performed this calculation for the same range of environmental noise levels, preference alignments, and experimental conditions as in the analysis of sequence ordering above. The results, shown in Fig. 3, demonstrate that consensus depends strongly on these factors. When individuals are habituated to a noisy environment or have strongly aligned preferences, then the group tends to come to a consensus decision, with a high consensus score under these conditions and in the naturalistic experimental environment ($\eta/\epsilon = 1$). Conversely, consensus scores are low when the habitual environment has little noise, or when preferences have very low alignment. Low-noise experimental conditions ($\eta/\epsilon = 1/2$) create greater consensus for any given values of $\rho$ and $\epsilon$ (Fig. 3B); this effect of experimental noise is in line with those explored in ref. 3. In Fig. 3C, the expected consensus score for each experimental condition is plotted as a function of the RSW (black points for the $\eta/\epsilon = 1$, red points for $\eta/\epsilon = 1/2$). Here one can see that RSW explains almost all of the variation in consensus as a function of $\rho$ and $\epsilon$.

In this experiment, agents made decisions once, in an ordered sequence. If agents are allowed to revise their choices after seeing the choices made by all others, this will, in general, lead to greater consensus values. For agents with identical preferences ($\rho = 1$), this iterative process must eventually lead to perfect consensus by virtue of Aumann’s agreement theorem (28). For value of $\rho$ below one, persistent disagreement is possible, but allowing agents to revise their choices nonetheless typically leads to greater consensus. Therefore, this should not be seen as a precise prediction of the quantitative degree of consensus in groups, but rather a demonstration of how consensus levels vary with environmental and experimental noise and preference alignment.

The Effect of Cryptic Group Substructures. The analysis, so far, has looked at the behavior of groups from a population that is undifferentiated, having a fixed utility correlation between any pair of individuals. Now I consider a group drawn from a population with further substructure: the existence of two distinct types of individual, $\alpha$ and $\beta$. I assume that individuals of the same type share a high correlation between their utility functions, whereas individuals of differing types have a low correlation. Furthermore, I assume that these individual types are cryptic, that is, they are not directly observable by other individuals, except through decision-making behavior. To illustrate the relevance of such a structure, consider a population with two distinct genotypes, with the result that individuals with each genotype require different relative amounts of carbohydrate and protein, but where the genotype has no other, directly perceivable phenotypic effect. In such a population, we would expect the interactions between individuals to reflect the reality that some conspecifics more closely share their nutritional preferences than others.

First, I consider a population of two equally numerous types, with those of a given type having identical preferences ($\rho = 1$), and individuals of differing types having no correlation in their utility function ($\rho = 0$). I calculated the probability that a focal decision maker of type $\alpha$ will choose option A conditional on a variety of possible sequences of previous decisions, and for a range of possible environmental noise levels (by symmetry, the results are the same for a focal individual of either type). These probabilities are shown in Fig. 4A. These results show a strong dependence on the environment. This pattern is close to that seen for agents with identical preferences (see ref. 3 and Fig. 1). However, in cases where previous decisions have not been unanimous, there is a strong dependence on the environmental noise, resembling that seen in Fig. 1A. The intuition behind this superposition is that, when a focal agent observes all previous decision makers in agreement, they conclude that it is likely they are all of the same type, and thus share identical preferences. Conversely, observing disagreement implies a high probability that the previous decision makers were of differing types. The consequences of this are particularly striking for the case of low environmental noise: In such environments, social influence is relatively low, except in the case of unanimity. That is, the difference between a unanimous set of previous decisions compared to a majority is qualitative rather than merely quantitative.

In the example above, the two distinct types were equally prevalent in the population as a whole. In general, this will not be the case: Frequency-dependent evolution can select for substantially unequal proportions of differing types of individual (e.g., ref. 20). What is the effect of such an imbalance on social behavior? I considered a population composed of 90% type $\alpha$ individuals and 10% type $\beta$, where $\rho_{\alpha,\alpha} = \beta_{\beta,\beta} = 0.9$ and $\rho_{\alpha,\beta} = 0.25$. I calculated the probability for a focal agent of either type to choose option A, based on the same set of putative previous decision sequences as in the previous example. The results for an individual of the majority type $\alpha$ are shown in Fig. 4B, and those for an individual of the minority type $\beta$ are shown in Fig. 4C. From these plots, one can see a substantial difference in the social behavior of the two types of individual: Type $\alpha$ individuals display strong social responses close to those of individuals in an undifferentiated and strongly aligned population (Fig. 1B),
while type $\beta$ individuals have a weak social response similar to those from an undifferentiated and weakly aligned population (Fig. 1A). This difference arises from the different utility correlation between individuals of each type and the population as a whole: Those in the majority can assume that most other individuals have closely aligned preferences and therefore are worth following, while those in the minority understand that most other individuals have differing preferences and therefore convey little useful social information.

**Discussion**

In this paper, I have described a model of collective decision-making by rational individuals with differing preferences, utility functions, or fitness outcomes when faced with decisions that depend on many factors. Developing this model has focused attention on precisely how the choices of one individual convey information about the likely utility of possible options to another individual, and when that information is likely to be reliable. This model goes beyond previous efforts to understand the foundations of social information by removing any assumption that individuals are identical, while distinguishing between differences in knowledge and differences of preference.

The calculations performed by rational agents in this model are complex and require making inferences about the inferences made by all other observed agents, in a known sequence of actions. Even in relatively small groups, this would be extremely cognitively demanding for any organism. Furthermore, the restriction to a single well-ordered sequence of actions is unrealistic for many real-world collective decisions, where agents may choose simultaneously and/or repeatedly. The intention of this model is not to derive the precise rules that real agents will apply, but to reveal the qualitative features of rational decision-making that any evolved, heuristic decision-making rule can be expected to emulate. I expect that cognitively sophisticated organisms will display behaviors that correspond to more of the fine details of the rational choice theory (such as sequence-specific behavior), while cognitively simpler organisms may use heuristics that mimic only the broader patterns seen in these results (such as the overall degree of sociality).

I have shown that the rational use of social information depends strongly upon the degree to which the focal decision maker believes that others share its preferences, with stronger social interactions between similar agents. This intuitive result provides an informational basis for several observed tendencies in animal groups, for example, for the observed stronger social response to conspecifics compared to heterospecifics in mixed-species bird groups (8, 22), for the tendency of baboons to follow movements initiated by close social affiliates (29), and for the preference of true conspecifics over a robotic imitation in zebrafish (30). It is also likely to contribute to homophily in human societies, for example, in housing choice (31), where the presence of many households with similar characteristics to oneself in an area provides useful information that your own needs and preferences can be met locally. In this way, collective patterns such as neighborhood segregation may be driven by the different information values of other individuals as well as other explanations such as the dislike of being in a local minority (32).

The use of social information also depends strongly on the quality of information from the environment. This, in turn, has important consequences for the collective behavior of social groups in different contexts. Different species of animal, for example, are habituated to widely differing levels of environmental noise. The same outcomes may have quite different fitness consequences for different individuals within an animal group. In human society, too, one finds contexts of varying uncertainty and agreement over preferences. The model predicts that these

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**Fig. 4.** Scenarios with cryptic group substructures. (A) The probability for a focal individual to choose option A as in Fig. 1, when individuals are drawn from a population of two equally prevalent subtypes with identical within-type preferences and zero between-type correlation. (B and C) The equivalent probabilities for individuals of the (B) major type and (C) minority type in the case where 90% of individuals are of the major type, with high within-type utility alignment ($\nu = 0.9$) and low between-type correlation ($\nu = 0.25$).
differing environmental and social contexts will lead to dramatically different social behavior. Previous work has shown that rational individuals with identical preferences should exhibit similar social responsiveness across a wide range of environmental conditions (3); in a noisy environment, private and social information both become less reliable to the same degree, such that the balance between the two remains constant. Here I have shown that, when preferences are not identical, this symmetry is broken. In a low-noise environment, private information is more trustworthy than social information, since it is not corrupted by differences in utility function. In a noisy environment, the uncertainty an individual has about others’ preferences is relatively low compared to the general uncertainty of all information, and thus the relative strength of social information increases. On the group level, this should lead to greater consensus in collective decision-making when individual preferences are strongly aligned and when uncertainty is high. These results imply that different species of animal (or the same species habituated to different environments) will display different social behavior in the wild in a predictable fashion. Groups composed of individuals habituated to noisier environments should be more cohesive and responsive to the actions of others. In addition to a stronger overall social response, I also found that high-noise environments and strong commonality in preferences led to a strong order-dependent use of social information, wherein a focal decision maker was more likely to follow a minority of recent decisions rather than a majority of preceding ones. This suggests that recent information will be dominant in cases where uncertainty about the world is generally high, and where a focal decision maker can assume that others want the same things as it does.

As noted in ref. 3, there is comparative ecological evidence to support the greater social responsiveness of agents habituated to noisier environments. However, this is complicated by the effect of context dependency where animals have been tested under conditions that differ from their habitual environment, and the effect is likely to be amplified by studies in the laboratory under common conditions. In common with earlier predictions by ref. 3, here I found that reducing the experimental noise level relative to the habitual environmental level strengthened social response, increased consensus, and made recent social information more important. Thus, individuals transposed from a noisy environment to the laboratory are subject to the increased sociability imposed by adaptation to a noisy environment and also the heightened social responsiveness from a reduced experimental noise level. There is also likely to be a strong selection bias in the experimental literature within the field of collective behavior toward instances where individuals display strong social interactions. Examples of trivially nonsocial behavior in low-noise environments, such as selecting the phone number for a known contact in a telephone directory, are therefore unlikely to be considered for study. These likely issues with existing experimental and observational data mean that further comparative studies are needed to assess how far the predictions in this paper are reflected in real populations.

Where agents are able to adapt their behavior to a variety of different contexts, one can expect to see individuals change their social responsiveness as the situation demands. A consequence of this would likely be to induce strong social feedback and a high salience for the most recent social information when environmental uncertainty is high and individuals transparently share the same objectives. This is a possible contributing factor in several examples of destructive herding behavior: Stock markets consisting of agents with strongly aligned goals (investment returns) tend to eliminate the value of private information for most participants (33), leading to high uncertainty for the typical investor, conducive to bubbles and crashes; crowd disasters are often associated with situations where individual utilities are strongly aligned (to escape), as private information becomes less reliable [e.g., through the presence of smoke (34), or through alcohol consumption (35)]. While, in these instances, such social feedback is ultimately maladaptive, such behavior may be the result of a social heuristic with a rational basis: When private information is scarce, follow others, especially those similar to yourself.

The results also demonstrate a potential source of leadership in heterogeneous groups. Where a group is composed of cryptic subtypes with differing preferences, we should expect those in the minority group to attend less to social information, and those in the majority to be more strongly social. On the individual level, this may contribute to the consistent differences in social response observed between individuals in some animal groups (e.g., refs. 36 and 37). At the group level, it would create an emergent leadership role for those in the minority, who would attend relatively more strongly to environmental cues, while those in the majority act preferentially as followers and thus ensure group cohesion. Since cohesive groups can be led by a relatively small number of relatively less social individuals (20, 23, 25, 38), these minority groups could have a disproportionately large effect on the collective decision-making. However, rather than the group being led by a subset of “informed individuals,” in this model, leadership would be conferred on those with unusual preferences rather than greater information. Heterogeneity of preferences could result from genetic or ontogenetic causes, and one may speculate on whether these may be connected with “personality” features such as boldness that have been linked to leadership (9, 39, 40). Conflicting preferences could also result from transient circumstances such as different hunger levels, creating “leadership by need” (25). Phenotypic and behavioral heterogeneity plays an important role in how social groups function (41), and the potential role of minority-preference groups as leaders adds another mechanism whereby individual-level variation can be translated into group-level behavioral differences.

I also found that heterogeneous groups created a special role for unanimity in decision-making, especially in relatively low-noise environments. When private information is of high quality (low noise), individuals would ordinarily tend to follow this information rather than attend to the choices of others, unless preference alignment is extremely high. However, in groups where some individuals have near-identical preferences, the existence of a consensus among previous decision makers can convince the focal individual to follow this consensus quite strongly (thus also reinforcing the consensus). Not only does this provide a mechanism for generating surprising degrees of consensus in groups that may have a low degree of average preference alignment, it also makes any deviation from that consensus especially powerful. Such a deviation from previous unanimity “breaks the spell” and drives further decision makers back to trusting their own private information.

It is important to reiterate that the model developed in this paper considers only the information value of other agents’ decisions. Aggregation may also carry intrinsic benefits (42), such as predation dilution (4), mate availability, and temperature regulation (43). Conversely, competition between agents for localized resources may discourage aggregation (44, 45). These additional factors complicate the identification of social information use in real animals, especially in the wild (46). Nonetheless, the results here suggest concrete predictions regarding collective behavior: 1) stronger social interactions between individuals with similar preferences, 2) greater aggregation in noisy environments, 3) stronger salience of recent social information in the presence of uncertainty, and 4) a correlation between leadership and minority types in heterogeneous groups. These predictions are potentially testable both through comparative analysis in wild
populations and through intervention studies in the laboratory, especially using artificial conspecifics (47–49) [although bearing in mind the prediction that social responses may be systematically stronger under laboratory conditions (3)]. The effect of genuinely differing preferences (as opposed to differing information) on collective decision-making has received relatively little experimental attention. The results in this paper show how future work may test whether any such effects are based on rational individual decision-making principles.

Materials and Methods

Utility Functions. I assume that any two possible choices can be distinguished by a set of values, \( x_1, x_2, \ldots, x_n \) characterizing the difference between the options across a set of independent factors. I further assume that the utility difference, \( U_k \), between the two options for any individual \( k \) is a weighted sum over these values,

\[
U_k = \sum_{i=1}^{n} \alpha_{i,k} x_i, \tag{3}
\]

where the \( \alpha_{i,k} \) are the weighting coefficients for individual \( k \) and thus specify their utility function. I assume that individuals are familiar with making decisions within a habitual environment and thus with the general properties of how the factors \( x_1, x_2, \ldots, x_n \) vary between decisions. I further assume, without loss of generality, that each factor \( x_i \) is measured on a standard scale, with mean zero and unit variance. Therefore, if the number of factors is large, or if the factors are themselves normally distributed, the prior distribution over the utility difference for individual \( k \) follows a normal distribution,

\[
p(\hat{U}_k) \propto \phi \left( \frac{U_k}{\sqrt{\sum_{i=1}^{n} \alpha_{i,k}^2}} \right), \tag{4}
\]

where \( \phi(\cdot) \) is the standard normal probability density function. Although individuals may have differing utility functions, I assume that these operate on the same scale, which I set, without loss of generality, to one. This is equivalent to specifying that \( \sum_{i=1}^{n} \alpha_{i,k}^2 = 1 \) for all \( k \) and therefore I retrieve the same prior distribution over utility differences as specified in ref. 3.

\[
p(U_1, U_2, \ldots, U_n) = \mathcal{N}(0, 0, \Sigma), \tag{5}
\]

where \( U = [U_1, U_2, \ldots, U_n]^\top \), and

\[
\Sigma = \begin{bmatrix}
1 & \rho_{1,2} & \ldots & \rho_{1,N} \\
\rho_{2,1} & 1 & \ldots & \rho_{2,N} \\
\vdots & \vdots & \ddots & \vdots \\
\rho_{N,1} & \rho_{N,2} & \ldots & 1
\end{bmatrix},
\]

with \( \rho_{ij} \) being the correlation between any two utility functions: \( \rho_{ij} = \sum_{k=1}^{n} \alpha_{i,k} \alpha_{j,k} \). At this point, I introduce the concept of an undifferentiated population. In such a population, I assume that all pairs of agents are equally aligned in their preferences, averaged over possible character characteristics \( x \), which corresponds to setting \( \rho_{ij} = \rho \) for all pairs \( i, j \). This corresponds to a social environment in which agents know that others have somewhat differing preferences, and know the general degree of overlap between those preferences and their own, but do not keep track of which individuals may be more or less closely aligned to themselves.

Private Information. Each agent receives private information about the value of \( x \) from the environment, for example, through physical sensory mechanisms. This information is assumed to be imperfect, and corrupted by noise residuals \( \epsilon_{ijk} \) which are independent between individuals and between factors, such that the measured value of \( x \), denoted as \( \hat{x}_i \), takes the form

\[
\hat{x}_k = x_k + \epsilon_{ijk}. \tag{7}
\]

For convenience, I define a "privately estimated utility" for agent \( k \), \( \hat{U}_k \),

\[
\hat{U}_k = \sum_{i=1}^{n} \alpha_{i,k} \hat{x}_i = U_k + \sum_{i=1}^{n} \alpha_{i,k} \epsilon_{ijk}. \tag{8}
\]

Defining \( \epsilon \) as the environmental noise level relative to the scale of utilities, with \( \text{cov}(\epsilon_{ij}) = 0 \) and \( \text{var}(\epsilon_{ij}) = \epsilon^2 \), one can therefore specify the conditional probability distribution of the privately estimated utility for agent \( k \),

\[
p(\hat{U}_k | U_k) = \phi((\hat{U}_k - U_k)/\epsilon), \tag{9}
\]

where I assume that \( \epsilon \) is the same for all agents (i.e., agents are homogeneous in their sensory sensitivity). Taken alongside Eq. 6, this implies that the true and privately estimated utilities for all agents are jointly normally distributed, with

\[
\begin{align*}
\text{cov}(U_1, \hat{U}_1) & = \text{cov}(U_2, \hat{U}_2) = \rho + \delta_{1,2}(1 - \rho), \\
\text{cov}(U_1, \hat{U}_2) & = \rho + \delta_{1,2}(1 + \epsilon^2 - \rho),
\end{align*}
\]

where \( \delta_{ij} \) is the Kronecker delta function.

In the case of the first individual to make a decision, their belief about the relative utilities of the two options is entirely determined by the combination of their prior expectations and their private information,

\[
p(U_1 | \hat{U}_1) \propto \phi(U_1) \phi(|\hat{U}_1 - U_1|/\epsilon). \tag{10}
\]

This implies that the first individual will choose option A if and only if \( \hat{U}_1 > 0 \).

Social Information. Having observed the decision made by the first individual, \( C_1 \), the second decision maker must combine this social information with their private information \( \hat{U}_2 \) to update their belief about the relative utilities based on their own preferences, \( U_2 \). Recalling that the true and privately estimated utilities of all agents are jointly normally distributed, the second individual’s beliefs are updated as

\[
p(U_1 | \hat{U}_1, C_1) \propto p(U_1 | \hat{U}_1) p(C_1 | U_2) \propto \phi(U_1 - U_2)/\epsilon \phi(U_2 | C_1 \rho/\sqrt{1 + \epsilon^2 - \rho^2}) \tag{12}
\]

This belief over \( U_1 \) further implies a critical value of the privately estimated utility, \( U_2^* \) such that

\[
\mathbb{E}(U_2 | U_2^*, C_1) = 0. \tag{13}
\]

If \( U_2 > U_2^* \), then the second individual will choose option A; otherwise, they will choose option B.

Further Decisions. The third decision maker can observe the social information from the first decision available to the second agent. Since I assume that agents are undifferentiated and share a common pairwise preference alignment \( \rho \), the third decision maker can also determine the critical value \( U_2^* \) that is calculated by the second individual as above. Having observed the second decision, they are therefore able to determine whether the privately estimated utility of individual 2 is greater or less than this critical value. The third individual should then combine their own private information, summarized by their privately estimated utility \( \hat{U}_3 \), with the knowledge they have of the bounds on what individuals 1 and 2 have observed. For example, if the third individual observes the first decision in favor of option A and the second observes in favor of option B, they should update their belief as follows:

\[
\begin{align*}
p(U_3 | \hat{U}_3, C_1 = 1, C_2 = -1) & \propto p(U_3 | \hat{U}_3) p(C_1 = 1, C_2 = -1 | U_3) \\
& \propto \phi(U_3) \phi(|U_3 - \hat{U}_2|)/\epsilon \times \\
& \int_{-\infty}^{U_2^*} \int_{0}^{\infty} \mathcal{N}' \left( \hat{U}_2, \rho U_3 \right) \mathcal{N}' \left( \hat{U}_2, \rho U_3 \right) dU_1 dU_2 \tag{14}
\end{align*}
\]

This updated belief structure further specifies a critical value \( U_3^* \) such that \( \mathbb{E}(U_3 | U_3^* \geq C_1, C_2 = -1) = 0 \). Subsequent agents can thus determine a bound on \( U_3 \) from the choice of individual 3, and this process can be followed recursively in turn for each further decision maker, with the
limits of integration chosen based on whether each individual chose option A or B.

Observation. In common with the model described in ref. 3, I consider the perspective of an external observer recording the decisions individuals make. This observer may record decisions in a natural environment, where the noise level is the same as the habitual environmental noise, \( \epsilon \). However, they may also record decisions under laboratory conditions that differ from those in the wild. Therefore, I define an experimental noise level, \( \gamma \). Low values of \( \gamma \) indicate that the laboratory is much less noisy than the natural environment, which is likely a common feature of behavioral experiments where the many environmental stimuli are removed or simplified so as to focus on the social response. An agent that continues to apply a decision-making rule based on the habitual noise level will calculate a critical value of the privately estimated utility as above. However, the probability that the actual privately estimated utility that the agent receives will exceed this critical value (which ultimately determines behavior) actually depends on the experimental noise level,

\[
P(U_i > U_j | U_i, \gamma) = \phi \left( \frac{U_i - U_j}{\gamma} \right).
\]

[15]

Populations with Cryptic Subtypes. So far, I have considered groups of undifferentiated individuals, making the approximation that every pair of individuals has the same degree of preference alignment. Now I consider groups drawn from a population in which there are two cryptic subtypes. Individuals have a strong preference alignment with those of the same subtype but a weak preference alignment with those of the other type. That these subtypes are cryptic implies that the focal decision maker is not aware of the specific type identities of previous decision makers, only the overall prevalence of the two types in the population.

Consider a population composed of two types, \( \alpha \) and \( \beta \), and let \( \gamma \) be the proportion of the population that is of type \( \alpha \). I assume that individuals of the same type have a preference alignment of \( \rho_{\alpha, \alpha} = \rho_{\alpha, \beta} = \rho_{\beta, \beta} \) while those of different types have \( \rho_{\alpha, \beta} = \rho_{\alpha, \beta} \text{ with } \rho_{\alpha, \beta} < \rho_{\text{high}} \). If the focal decision maker can determine the types of the previous decision makers, then it can apply the model previously described, but with an adapted covariance between different individuals,

\[
\text{cov}(U_i, U_j) = \begin{cases} 
\rho_{\alpha, \alpha} & \text{if } k, l \text{ same type} \\
\rho_{\alpha, \beta} & \text{if } k, l \text{ different types.}
\end{cases}
\]

[16]

To determine its own utility belief function, a focal decision maker must evaluate the probability that the observed decisions were made, based on each possible sequence, \( s \), of types among previous decision makers, and weight these by their relative probabilities,

\[
p(U_i | U_1, C_1, \ldots, C_{k-1}) \propto p(U_i | U_j) \prod_{k \in S} p(C_k, \ldots, C_{k-1} | U_k, s)p(s),
\]

[17]

where \( S \) represents all possible sequences of individual types, and \( p(s) = \gamma^n(1 - \gamma)^{n-k} \), with \( n_\alpha \) and \( n_\beta \) being the number of individuals of each type in the sequence \( s \).

This process can be carried out recursively as for the undifferentiated population, with the focal agent calculating a different critical value of \( U_j^* \) for a previous decision maker \( j \) depending on whether that agent was type \( \alpha \) or \( \beta \) in a given putative sequence. Since the types of each individual are cryptic, the focal agent knows that other individuals are also ignorant of the types of the earlier decision makers.

Code Availability. Code to reproduce the results of this paper in R is included as Dataset S1.

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