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Crump, A, Bethell, EJ, Earley, R, Lee, VE, Mendl, M, Oldham, L, Turner, SP and Arnott, G (2020) Emotion in animal contests. Proceedings of the Royal Society B: Biological Sciences, 287 (1939). ISSN 0962-8452

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Emotion in animal contests

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Abstract: Emotions encompass cognitive and behavioural responses to reward and punishment. Using contests as a case-study, we propose that short-term emotions underpin animals' assessments, decision-making, and behaviour. Equating contest assessments to emotional "appraisals", we describe how contestants appraise more than resource value and outcome probability. These appraisals elicit the cognition, drive, and neurophysiology that governs aggressive behaviour. We discuss how recent contest outcomes induce longer-term moods, which impact subsequent contest behaviour. Finally, we distinguish between integral (objectively relevant) and incidental (objectively irrelevant) emotions and moods ("affective states"). Unlike existing ecological models, our approach predicts that incidental events influence contest dynamics, and that contests become incidental influences themselves, potentially causing maladaptive decision-making. As affective states cross contexts, a more holistic ethology (incorporating emotions and moods) would illuminate animal cognition and behaviour.

Keywords: Affective state, assessment, cognition, resource-holding potential, resource value, winner/loser effects.

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1. Introduction

Consider this: animal behaviour is underpinned by emotions and moods ("affective states"; see [1–11]). We define emotions as short-term states elicited by stimuli (or their predictors) that animals will work to acquire (rewards; e.g. prey) or avoid (punishments; e.g. predators [5,12,13]). Moods are longer-term states, which represent the cumulative average of emotions over time [14,15]. These functional definitions apply to any organism with a central nervous system [1]. Animal welfare scientists, neuroscientists, and psychopharmacologists now recognise that affective states play a key role in decision-making [6,7]. However, behavioural ecologists and fundamental ethologists have not yet embraced emotions and moods [16].

Two main dimensions characterise affective states: valence and arousal [6,17–19] (Figure 1). Valence, which ranges from positive to negative, encapsulates the fitness benefits and costs associated with a stimulus, either anticipated or actual [7]. Meanwhile, arousal (emotional intensity) indicates stimulus importance or urgency. High-arousal affective states divert attentional resources to the stimulus [20] and predispose vigorous action [21]. As well as emotions and moods, valence and arousal define sensations (e.g. pain) and interoception (which internal stimuli elicit; e.g. hunger [10]). Burgdorf and Panksepp [22] hypothesised that positive-valence, high-arousal states represent the activation of a reward acquisition system, whereas negative-valence, high-arousal states represent the activation of a punishment avoidance system. By conceptualising affective states in terms of reward and punishment, this dimensional approach captures their evolutionary function and avoids categorical labels that can lead to anthropomorphism (e.g. [8]).

Figure 1. Valence and arousal define affective states (grey box), which encompasses emotions and moods [6]. Moving from Q3-Q1 is increasingly appetitive; Q2-Q4 is increasingly aversive.

Anderson and Adolphs [1] identified two further characteristics of affective states. As well as having valence and arousal ("scalability"), emotions "generalise": various stimuli and situations can induce a particular affective state, and affective states can be associated with various behavioural responses. Affective states also "persist" after stimulus removal. Thus, emotions do not mediate fixed responses to specific stimuli, because fixed responses neither generalise nor persist. Examples of non-affective behaviours, therefore, include withdrawal reflexes (which are genetically encoded from birth) and sexual imprinting (which is learnt during development and subsequently invariant). Emotions, on the other hand, facilitate flexible behaviour in complex, variable environments [23].

We propose that animal contests are an example of affective behaviour. Contests are direct inter-individual interactions that determine access to resources, such as food, mates or territory (i.e. rewards [24]). Resource value (RV) is the fitness benefit of the resource [25]. Contest costs include energy and time expenditure, injury, and even death (i.e. punishments [26]). Greater potential benefits justify greater costs, so increasing RV increases investment [27,28]. However, contest costs and outcomes are not fixed. Resource-holding potential (RHP) is the ability to win contests, comprising traits like size, skill, and weaponry [29–31]. Animals with a higher RHP are better at winning, so they are more likely to gain resources. Contests involve acquiring resources and avoiding punishments (valence), vary in intensity and escalation (arousal), are elicited by diverse stimuli and exhibited in various ways (generalisation), and continue after the inciting event (persistence). These features imply an internal (i.e. affective) state mediating the link between reward, punishment, and contest behaviour.

Previous researchers have not comprehensively applied affective state theory to animal contests. However, conceptualised as responses to rewards, punishments, and their predictors, emotions cover contest information-gathering, decision-making, and behaviour. This novel approach extends and refines contest motivation models. For example, Elwood and Arnott [32] explained contest dynamics in terms of two dimensions: RV and costs. A contestant engages if RV exceeds costs and withdraws if costs exceed RV. Whereas RV usually remains stable, costs accumulate throughout the contest. If costs increase enough to exceed RV, a contestant's strategy switches

from engage to withdraw. This model approximates the valence dimension of affective states – RV representing positive valence and costs representing negative valence – except that valence is not specific to contests [6,7,14,15].

In this review, we use contests as a case-study for applying emotion theory to behavioural ecology and fundamental ethology. We argue that contestants evaluate contest benefits and costs, and that these "appraisals" elicit emotional episodes encompassing contest decisions and behaviour. We describe how the affective outcome of contests might produce experience effects: prior winners' tendency to initiate and win (and prior losers' tendency to avoid and lose) subsequent contests. Unlike traditional ecological models, our perspective predicts that affective states previously induced in other behavioural contexts will impact contest dynamics. These objectively irrelevant influences could mediate contest decisions and cause maladaptive behaviour.

2. Structure of Emotions

Emotions are elicited by appraisals: evaluations of stimuli, their context, and their personal significance [33]. Scherer [34] proposed that humans sequentially appraise stimulus novelty, intrinsic valence, congruence with personal goals, outcome probability, discrepancy from expectations, situation controllability, other individuals' responsibility, and whether potential responses are socially acceptable. Appraisal outcomes determine and differentiate emotions [35], with continuously-updated reappraisals regulating the response [36]. Other mammals, birds, and fish also appear to appraise stimuli [23]. In lambs (*Ovis aries*), for example, stimulus novelty, discrepancy from expectations, controllability, and social context impact physiology and behaviour [37]. These inferred appraisals elicit flexible emotional responses, which account for current conditions and personal circumstances, as well as intrinsic stimulus characteristics.

Emotions have multiple components that can be measured empirically [10,38] (Figure 2). These include changes in (1) cognition: information-gathering and processing; (2) drive: manifested as the work animals will invest to access reward or avoid punishment; and (3) neurophysiology: central and peripheral nervous system activity, and neuroendocrine function. Such changes facilitate the performance of (4)

behaviour, producing an organism-level response to reward and punishment [5,39,40]. Threatening stimuli, for instance, impact (1) cognition: increasing attention to the threat; (2) drive: maximising the work animals will invest in performing freeze, fight, or flight responses; and (3) neurophysiology: activating both the sympathetic nervous system and hypothalamic-pituitary-adrenal axis. These changes prepare the individual for (4) behaviour: avoiding, attacking or escaping the threat.

Figure 2. An emotional episode (white box). Appraisals of stimuli, their context, and their personal significance elicit the emotion (grey box), whose components include cognition, drive, and neurophysiology. These components govern the expression of behaviour. Conscious "feelings" (not shown) are another potential component, but not essential.

Conscious feelings, another potential emotion component, cannot be directly measured. Humans describe feelings through language, which animals cannot do. As a result, animal researchers usually study other emotion components and remain agnostic about feelings [4,9,10]. Indeed, many human psychologists recognise unconscious emotion, where the measurable components occur without corresponding feelings [41]. For example, Winkielman *et al.* [42] showed people positive or negative facial expressions. The images appeared too briefly for conscious awareness. When subsequently offered a novel drink, subjects shown the positive expression poured more, drank more, and paid more than subjects shown the negative expression. Self-reported affective states did not differ between treatments. In animals, the relationship between feeling and non-feeling emotion components is an important area for future research [43]. However, for present purposes, we view emotions as functional states that may or may not be accompanied by feelings.

Animal welfare scientists and psychopharmacologists investigate affective states through both experiment and observation. In emotion induction experiments, rewards (e.g. enrichment [44]) induce positive emotions, and punishments (e.g. social defeat [45]) induce negative emotions [13]. Pharmacological manipulations may also induce positive- and negative-valence states [46]. Observationally, the measurable components of an emotional episode can indicate valence [4]. This includes changes in (1) cognition: attention [47], judgement [48,49], and memory biases [50]; (2) drive: the work animals will invest to access reward or avoid punishment [51,52]; (3)

neurophysiology: brain and neuroendocrine circuits [5,8], and peripheral nervous system activity [2]; and (4) behaviour: approach, exploration, and play are often positively valenced, whereas avoidance and hiding are often negatively valenced [2]. Detailed discussion of empirical methods is beyond our scope, but we direct readers to previous reviews on measuring affective states in animals [2,4,6–8,10,11,47–49,51,52].

3. Initiating, Escalating, and Quitting Contests

Contest theorists emphasise two key assessments: animals assess RV (which determines fitness benefits and motivation) and RHP (which predicts fitness costs and outcome likelihood [25,29]). Contestants may assess only their own RHP (self-assessment [53,54]) or compare their RHP to their opponent's (mutual assessment [26,28,55]). In a meta-analysis of 36 species' assessment strategies, Pinto *et al.* [56] found that self-assessment is more common than mutual assessment.

Appraisal theory articulates and extends contest theory. The former predicts broader evaluations of the resource, opponent, and context, all related back to the individual's own goals. Under Scherer's [34] sequential theory, contestants would first appraise novelty. Familiar resources are valued above novel resources (e.g. residency effects [57,58]), whilst dominance hierarchies reduce aggression towards familiar rivals [59]. Second, contestants would appraise the resource's intrinsic valence (objective RV; e.g. the calories in food). Third, contestants would appraise whether the resource contributes to their goals (subjective RV; e.g. starving animals value food most [60]). Fourth, contestants would appraise outcome probability (which covers RHP assessments). Animals avoid or de-escalate contests they will probably lose [29]. Fifth, contestants would appraise discrepancy from expectations. Compared to unconditioned controls, animals trained that a stimulus signals reward become more aggressive when the stimulus is unrewarded [61–63]. Sixth, contestants would appraise their response's compatibility with social context. Observer presence can modify animals' behaviour (audience effects [64–66]) and watching contests can modify the observers' subsequent behaviour (bystander effects [64,67]). During ongoing contests, animals also reappraise assessments, adjusting their behaviour as

information and costs accumulate [31,55]. These appraisals have all been empirically documented, but several are not incorporated into current contest theory.

We further postulate that appraisals unify reward and punishment inputs into a decision-making common currency [68,69]. This facilitates cross-context comparisons between competing emotions, moods, sensations, and interoception. For instance, food-deprived goldfish (Carassius auratus) endure more electric shocks to feed than well-fed goldfish [60]. Following shocks, fewer hermit crabs (Pagurus bernhardus) evacuate preferred Littorina shells than non-preferred Gibbula shells [70]. We conceptualise valence as the common currency in these reward/punishment trade-offs. Contestants likewise weigh RV against potential contest costs and outcome likelihood [32]. In self-assessment, contestants' affective states integrate RV and own RHP information. Animals persist until they reach a negative-valence threshold: the maximum cost they will pay for the resource. This threshold may be energetic [71,72] or include injury costs as well [73]. In mutual assessment, affective states integrate RV, own RHP, and opponent RHP information. Animals withdraw when they establish that their opponent has a higher RHP [55], perhaps when they tip below neutral valence. Both self- and mutual assessment models require unidimensional (valence) comparisons of fitness-relevant information.

Affective states may also determine assessment strategy. Researchers traditionally viewed assessment strategies as fixed (e.g. [29,32,74]), but now recognise individual-and population-level variation [75–77]. For example, green anoles (*Anolis carolinensis*) [78], mangrove killifish (*Kryptolebias marmoratus*) [79], and fiddler crabs (*Uca mjoebergi*) [80] use mutual assessment when deciding whether to escalate a contest, and self-assessment during the fight. Humans in positive affective states rely on heuristics (i.e. rules of thumb) more than humans in negative affective states [81]. When assessing the strength of an argument, for instance, people experiencing positive emotions use the author's expertise, whereas people in neutral states judge the content (i.e. deeper processing [82,83]). In animal contests, positive valence may also promote less cognitively demanding assessment strategies, such as self-assessment or rules of thumb (e.g. "resident wins"; see [84]). Future research could manipulate affective states to test this. We hypothesise that prior reward will lead to self-assessment, whereas prior punishment will lead to mutual assessment.

Having defined emotions as functional responses to reward and punishment, we can say that contest assessments (i.e. appraisals) elicit emotions. We propose that positive emotions about potential contests indicate that fitness benefits outweigh perceived costs, activating a reward acquisition system [6,7]. This system covers (1) cognition: information gathering and decisions to enter and escalate contests; (2) drive: work invested to attack; (3) neurophysiology: dopamine and opioid activity; and (4) behaviour: threat displays and aggression. By contrast, negative emotions indicate that perceived contest costs outweigh fitness benefits, activating a punishment avoidance system. This system covers (1) cognition: information gathering and decisions to avoid and withdraw; (2) drive: work invested to escape; (3) neurophysiology: reduced serotonergic activity; and (4) behaviour: submission and retreat.

From a human perspective, linking positive valence and aggressive behaviour may seem counterintuitive. Anger, for instance, *feels* negative [85], but causes aggression [86,87]. However, this perspective is based on our conscious experience of emotion (i.e. the feeling component). The non-feeling components indicate that anger is a reward acquisition emotion (i.e. positive valence), not a punishment avoidance emotion (i.e. negative valence [88]). Anger drives approach towards the inducing stimulus, whereas negative-valence emotions drive withdrawal [88]. As a result, our functional definition of emotion – which does not require conscious feeling – categorises anger as positively valenced. Negative-valence emotions can lead to aggressive behaviour, but only when withdrawal is not an option (e.g. cornered animals lashing out). In the present manuscript, we only consider positive-valence aggression, where the aim is resource acquisition.

This review focuses on contest initiation, winning, and losing, but affective states might also govern behavioural transitions within contests, such as levels of display or escalated aggression (e.g. [78–80]). From an emotion standpoint, the transitions that bookend contests are more empirically tractable. Applying an emotional event precontest indicates how emotions influence initiation, for example, whereas applying an emotional event between contests indicates how emotions disrupt experience effects. Tracking emotions during contests is more challenging, as contests are ongoing emotional events. To resolve this issue, we propose startling contestants at set points during a contest [29,89]. Motivation theorists interpret faster contest resumption (i.e.

shorter startle latencies) as stronger motivation to fight [33]. However, affective state influences the startle reflex [47]. In humans [90], rhesus macaques (*Macaca mulatta*) [91], and rats (*Rattus norvegicus*) [90], negative-valence states increase startle duration and magnitude. Future researchers could use startle duration to understand how valence relates to within-contest behavioural transitions.

To summarise, emotion theory correctly predicts that contest assessments cover more than RV and RHP. Animals assess the resource, opponent, and context, in relation to individual circumstances. We hope researchers investigate whether additional human appraisals influence contest dynamics in other species. For example, perhaps agency appraisals (who was responsible? what did they intend?) influence contest decision-making. Under our definition of emotion, these appraisals elicit emotional responses that reflect personal circumstances and prevailing conditions. Conceptualising cognition, drive, and neurophysiology as a unified affective state underpinning behaviour explains existing results and generates new hypotheses.

4. Contest Outcome and Experience Effects

Contest outcomes indicate how an individual's RHP compares to the population's RHP [92,93]. Assuming self-assessment, wins signal relatively high personal RHP and losses signal relatively low personal RHP. Winners, therefore, initiate, escalate, and win more subsequent contests (winner effects), whereas losers avoid and lose more subsequent contests (loser effects [94–96]). We conceptualise contests as emotional events, so winning induces positive-valence emotions that increase aggressive behaviour and losing induces negative-valence emotions that reduce aggressive behaviour (even if actual RHP does not change). By reflecting cumulative emotional outcomes, winner and loser effects represent longer-term moods (Figure 3).

Figure 3. Cumulative emotional valence determines mood [11] (manifested in aggression). Considering only integral (objectively contest-relevant) influences, white dots are wins and black dots are losses. Considering both integral and incidental (objectively contest-irrelevant) influences, white dots are rewards and black dots are punishments.

Both emotions and moods cause cognitive changes, such as judgement and decision-making biases [97]. People in positive affective states interpret ambiguous stimuli

more optimistically than people in negative affective states [81], whereas pessimistic judgements characterise depression and anxiety [81,98–100]. Animals also exhibit judgement biases. Under ambiguity, mammals, birds, fish, and insects in positive affective states have higher expectations of reward and lower expectations of punishment than animals in negative affective states [7,46,48,49,101]. Assuming reward and punishment experience predicts likely outcomes in the present, moods indicate whether ambiguous stimuli signal positive or negative outcomes, leading to judgement biases [14,49]. We, therefore, suggest that mood-induced judgement bias underlies contest experience effects. Winners gain fitness-enhancing resources, so winning is positively valenced. Thus, previous winners should be relatively optimistic about unknown rewards (RV) and outcome likelihood (RHP), and correspondingly more aggressive. Losing, meanwhile, is negatively valenced, so losers should be more pessimistic and less aggressive. Indeed, perceived RHP, rather than actual RHP, influences winner and loser effects [94,95] (cf. [102]).

Empirical evidence suggests that contests induce judgement biases. Researchers have trained both dominant and subordinate animals to associate one stimulus with a highvalue reward (leading to shorter response latencies) and another stimulus with a lowvalue reward or no reward (leading to longer response latencies). When subsequently presented with untrained intermediate stimuli, dominant animals respond faster and more frequently than subordinates (rats: [103]; pigs, Sus scrofa: [104]; tufted capuchins, Sapajus apella: [105]). We interpret the dominant animals' higher reward anticipation as optimism, which may reflect wins inducing positive valence. In similar tasks, rats [45] and Murray cod (Maccullochella peelii) [106] that repeatedly lose contests exhibit lower reward anticipation towards ambiguous stimuli, which we interpret as pessimism. Equivalent opponent-directed behaviour – reduced likelihood of attacking an ambiguous rival – would constitute a loser effect. As judgement biases influence responses to ambiguity more than responses to predictable outcomes [7,49], we hypothesise that judgement biases impact behaviour in contests with unpredictable outcomes (where opponents have similar RHP) more than contests with predictable outcomes (where opponents' RHP differs markedly).

Experience effects also suggest that contests can be intrinsically rewarding [107]. In addition to yielding external reward, aggressive behaviour itself (and particularly

winning) seems to induce positive affective states, which may inform future decisions. For example, mice (*Mus musculus*) learn instrumental responses to access and attack submissive opponents [108]. Responses decline for non-submissive opponents, revealing that outcome matters. Moreover, winning induces conditioned place preference in mice [109], Syrian hamsters (*Mesocricetus auratus*) [110], and green anoles [111]. From an affective state perspective, positive emotions reward this conditioning. Affective reinforcement might also occur within a single contest. For instance, accurate strikes [30] or appropriate assessments [112] may be rewarding.

To recap, we suggest that moods, which reflect contest outcome experience, mediate expectations about unknown RV and future outcomes. Mood-induced judgement bias and affective reinforcement may underpin these experience effects. To investigate judgement bias, contest researchers could measure optimism pre- and post-contest (see [48,49]). We predict that wins induce optimism and losses induce pessimism, with state optimism producing winner effects and state pessimism producing loser effects. Exploring the role of neurotransmitters linked to reward, such as opioids, could reveal whether contests are intrinsically rewarding.

5. Crossing Behavioural Contexts

So far, we have considered adaptive affective states. There are clear fitness benefits to cumulative experience informing reliable assessments, but existing optimality models already predict these effects. How do emotions and moods advance our understanding?

Integral affective states are objectively relevant to a cognitive process. In humans, for example, sunshine (stimulus) induces positive valence (emotion) that causes a decision (cognition) to go outside (behaviour). Incidental affective states, on the other hand, influence objectively unrelated cognitive processes [38,81,113–115]. For example, people rate their overall life satisfaction higher on sunny days than rainy days [116]. Sunshine (stimulus) induces positive valence (emotion) that causes an objectively unrelated assessment (cognition) to be reported positively (behaviour). Incidental affective states, thus, distinguish optimal and affective decision-making. Optimality models only use integral information, whereas affective states incorporate incidental influences as well.

Although understudied in behavioural ecology, incidental affective states influence animal cognition and behaviour. Starlings (Sturnus vulgaris) with enriched housing judge unrelated temporal stimuli more optimistically [44], whilst honeybees (Apis *mellifera*) shaken aversively judge unrelated olfactory stimuli more pessimistically [117]. Moreover, isolating rats improves recall of unrelated light and sound stimuli [50]. It follows that incidental information may influence contest behaviour, and that rewards and punishments in general – not wins and losses specifically – induce "winner" and "loser" effects (Figure 3). For instance, positive-valence female interactions increase aggressive behaviour in male speckled wood butterflies (Pararge aegeria) [118] and wolf spiders (Venonia coruscans) [119], whereas negative-valence predator exposure decreases aggressive behaviour in daffodil cichlids (Neolamprologus pulcher) [120]. However, a note of caution: apparently incidental influences may be functionally integral. Presence of a potential mate, for example, increases contest benefits, and predation risk increases contest costs [118–120]. We must understand a species' ecology to determine whether cross-context variables are objectively relevant, and hence whether they are integral or incidental. We hope that new studies fill this knowledge-gap. Contest researchers could borrow affective state research methods from animal welfare science and psychopharmacology. Exposing fish to antidepressants and anxiolytics in wastewater has produced equivocal results: venlafaxine increases aggression [121], but fluoxetine reduces aggression [122]. To test whether incidental affective states influence contest behaviour, we need controlled interventions in more species.

Incidental affective states not only influence contests; contests might also induce incidental affective states and influence objectively unrelated cognitive processes (see [123]). For example, rats that repeatedly lose contests develop anhedonia: reduced reward sensitivity, expressed in non-contest situations and linked to depression in humans. Giving the rats unrelated but signalled food rewards reverses this effect [124]. Compared to tufted capuchins with subordinate bystanders, capuchins exposed to aggressive bystanders allocate more attention towards humans [125]. Dominant capuchins [105] and pigs [104] expect more positive outcomes from ambiguous spatial stimuli (i.e. optimism), whilst subordinate cod expect fewer positive outcomes from ambiguous spatial stimuli [106] (i.e. pessimism). Contest-induced incidental affective states may influence virtually any decision. Is brightly-coloured prey toxic

or a mimic? Are rustling leaves a predator or the wind? When moods bias decisions, the most encountered emotional stimuli with the longest duration and most polar valence might determine behaviour, regardless of objective relevance. It is possible that frequently winning contests, for example, may induce optimism that rare prey is edible, even if the prey is usually toxic. This example illustrates how decision-making using incidental information can negatively impact fitness. Incidental affective states may cause maladaptive behaviour [38].

Given their maladaptive potential, we suggest two reasons for incidental affective states. First, to be selected, cross-context affective states must increase fitness *on average* – not necessarily *every time*. Nettle and Bateson [14] noted that recent environment and physical condition persist across behavioural contexts. Lame animals, for instance, cannot fight, forage or flee from predators, so information from each of these contexts is integral to the others. Cross-context affective states will be selected if most are integral, even if some are incidental. In humans, various measures increase the likelihood that cross-context affective states only influence relevant cognition [114,115]. For example, people associate their affective states with concurrent cognitive processes [126]. Incidental emotional influences are also less common than incidental moods, because emotions usually have an obvious cause [114]. Animal research may reveal similar mechanisms to limit incidental affect.

The second possible explanation is that incidental affective states dominate when animals lack reliable information, or when acquisition and storage costs outweigh the benefits [59]. This is why humans evaluating ambiguous stimuli (e.g. brand names without product details) rely on incidental affective states [127]. In animal contests, a fight indicates rival RHP most accurately, but entails substantial investment and potential injury [64,67]. Assessments in other contexts carry their own cost/accuracy trade-offs. Bystander effects avoid fight costs and reflect individual RHP, but they require individual discrimination and recall [32]. Winner and loser effects are less cognitively demanding, but based on previous opponents' RHP. This measure will predict future opponents' RHP less accurately than individual assessments. We hypothesise that mood does not even distinguish between behavioural contexts, further reducing both cognitive requirements and accuracy. Incidental affective states may, therefore, influence decisions when contestants have less reliable information or

high information-gathering costs (e.g. intruders). From this perspective, incidental affective states are the "best of a bad job".

In summary, integral affective states are objectively relevant and adaptive, whereas incidental affective states are objectively irrelevant and potentially maladaptive. Incidental influences may nonetheless seep in when integral information is unavailable or costly. Despite preliminary evidence, we do not yet know the extent of incidental affective states in animal decision-making. We hope that future researchers test whether objectively unrelated stimuli impact contest dynamics. Without integral influences, we predict that generic rewards increase aggression and generic punishments decrease aggression.

6. Conclusion

An affective states approach generates novel predictions and opens new avenues for behavioural ecology (Table 1). Both emotions and contest behaviour rely on assessments of stimuli and their personal significance; both enlist cognition, drive, and neurophysiology; and both reflect reward and punishment experience. We equate contest assessments to emotional appraisals, which determine contest decisionmaking and behaviour. We explain experience effects as wins inducing positive moods and losses inducing negative moods. This hypothesis, and our conception of contests as emotional episodes, predicts that manipulating affective state will modify contest behaviour. As well as integral influences, incidental affective states may impact contests, and contest-induced affective states may impact objectively unrelated behaviours. We hypothesise that high-frequency, long-lasting, polar-valence events disproportionately influence animal decision-making and behaviour, even if incidental. Moreover, despite our focus on contests, emotion theory may underpin all non-reflexive behaviour – from signalling to mate choice to parental care. Behavioural ecologists and fundamental ethologists study these fields individually, but affective states transcend our boundaries. We need a more holistic ethology to understand affective cognition and behaviour.

Table 1. Major predictions and outstanding questions from applying emotion theory to animal contests

Funding: A.C. received a postgraduate studentship from Northern Ireland's Department for the Economy. This paper is part of a project that has received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme, Grant Number 851145.

Acknowledgements: We thank Robert Elwood, an anonymous reviewer, and Innes Cuthill for their useful comments and insights on this manuscript.

References

- 1. Anderson DJ, Adolphs R. 2014 A framework for studying emotions across species. *Cell* **157**, 187–200. (doi:10.1016/j.cell.2014.03.003)
- 2. Boissy A, Manteuffel G, Jensen MB, Moe RO, Spruijt B, Keeling LJ, Winckler C, Forkman B, Dimitrov I, Langbein J, Bakken M, Veissier I, Aubert A. 2007 Assessment of positive emotions in animals to improve their welfare. *Physiology & Behavior* **92**, 375–397. (doi:10.1016/j.physbeh.2007.02.003)
- 3. Gygax L. 2017 Wanting, liking and welfare: The role of affective states in proximate control of behaviour in vertebrates. *Ethology* **123**, 689–704. (doi:10.1111/eth.12655)
- 4. Kremer L, Klein Holkenborg SEJ, Reimert I, Bolhuis JE, Webb LE. 2020 The nuts and bolts of animal emotion. *Neuroscience and Biobehavioral Reviews* **113**, 273–286. (doi:10.1016/j.neubiorev.2020.01.028)
- 5. LeDoux J. 2012 Rethinking the Emotional Brain. *Neuron* **73**, 653–676. (doi:10.1016/j.neuron.2012.02.004)
- 6. Mendl M, Burman OHP, Paul ES. 2010 An integrative and functional framework for the study of animal emotion and mood. *Proceedings of the Royal Society B: Biological Sciences* **277**, 2895–2904. (doi:10.1098/rspb.2010.0303)
- 7. Mendl M, Paul ES. 2020 Animal affect and decision-making. *Neuroscience and Biobehavioral Reviews* **112**, 144–163. (doi:10.1016/j.neubiorev.2020.01.025)
- 8. Panksepp J. 2011 The basic emotional circuits of mammalian brains: Do animals have affective lives? *Neuroscience and Biobehavioral Reviews* **35**, 1791–1804. (doi:10.1016/j.neubiorev.2011.08.003)
- 9. Paul ES, Mendl MT. 2018 Animal emotion: Descriptive and prescriptive definitions and their implications for a comparative perspective. *Applied Animal Behaviour Science* **205**, 202–209. (doi:10.1016/j.applanim.2018.01.008)
- 10. Paul ES, Sher S, Tamietto M, Winkielman P, Mendl MT. 2020 Towards a comparative science of emotion: Affect and consciousness in humans and animals. *Neuroscience and Biobehavioral Reviews* **108**, 749–770. (doi:10.1016/j.neubiorev.2019.11.014)
- 11. Webb LE, Veenhoven R, Harfeld JL, Jensen MB. 2018 What is animal happiness?. *Annals of the New York Academy of Sciences* **1438**, 1–15. (doi:10.1111/nyas.13983)
- 12. Carver CS. 2001 Affect and the Functional Bases of Behavior: On the Dimensional Structure of Affective Experience. *Personality and Social Psychology Review* **5**, 345–356. (doi:10.1207/S15327957PSPR0504 4)
- 13. Rolls ET. 2005 Emotion Explained. Oxford: Oxford University Press.

- 14. Nettle D, Bateson M. 2012 The evolutionary origins of mood and its disorders. *Current Biology* **22**, 712–721. (doi:10.1016/j.cub.2012.06.020)
- 15. Trimmer PC, Paul ES, Mendl MT, McNamara JM, Houston AI. 2013 On the evolution and optimality of mood states. *Behavioral Sciences* **3**, 501–521. (doi:10.3390/bs3030501)
- 16. Fraser D. 2009 Animal behaviour, animal welfare and the scientific study of affect. *Applied Animal Behaviour Science* **118**, 108–117. (doi:10.1016/j.applanim.2009.02.020)
- 17. Posner J, Russell JA, Peterson BS. 2005 The circumplex model of affect: An integrative approach to affective neuroscience, cognitive development, and psychopathology. *Development and Psychopathology* **17**, 715–734. (doi:10.1017/S0954579405050340)
- 18. Russell JA. 1980 A circumplex model of affect. *Journal of Personality and Social Psychology* **39**, 1161–1178. (doi:10.1037/h0077714)
- 19. Russell JA. 2003 Core Affect and the Psychological Construction of Emotion. *Psychological Review* **110**, 145–172. (doi:10.1037/0033-295X.110.1.145)
- 20. Storbeck J, Clore GL. 2008 Affective arousal as information: How affective arousal influences judgments, learning, and memory. *Social and Personality Psychology Compass* **2**, 1824–1843. (doi:10.1111/j.1751-9004.2008.00138.x)
- 21. Bach DR, Dayan P. 2017 Algorithms for survival: A comparative perspective on emotions. *Nature Reviews Neuroscience* **18**, 311–319. (doi:10.1038/nrn.2017.35)
- 22. Burgdorf J, Panksepp J. 2006 The neurobiology of positive emotions. *Neuroscience and Biobehavioral Reviews* **30**, 173–187. (doi:10.1016/j.neubiorev.2005.06.001)
- 23. Faustino AI, Oliveira GA, Oliveira RF. 2015 Linking appraisal to behavioral flexibility in animals: implications for stress research. *Frontiers in Behavioral Neuroscience* **9**, 104. (doi:10.3389/fnbeh.2015.00104)
- 24. Briffa M, Hardy ICW. 2013 Introduction to Animal Contests. In *Animal Contests* (eds Hardy ICW, Briffa M.), pp. 1–4. Cambridge: Cambridge University Press.
- 25. Arnott G, Elwood RW. 2008 Information gathering and decision making about resource value in animal contests. *Animal Behaviour* **76**, 529–542. (doi:10.1016/j.anbehav.2008.04.019)
- 26. Enquist M, Leimar O. 1990 The evolution of fatal fighting. *Animal Behaviour* **39**, 1–9. (doi:10.1016/S0003-3472(05)80721-3)
- 27. Enquist M, Leimar O. 1987 Evolution of fighting behaviour: The effect of variation in resource value. *Journal of Theoretical Biology* **127**, 187–205. (doi:10.1016/S0022-5193(87)80130-3)
- 28. Hammerstein P, Parker GA. 1982 The asymmetric war of attrition. *Journal of Theoretical Biology* **96**, 647–682. (doi:10.1016/0022-5193(82)90235-1)
- 29. Arnott G, Elwood RW. 2009 Assessment of fighting ability in animal contests. *Animal Behaviour* **77**, 991–1004. (doi:10.1016/j.anbehav.2009.02.010)
- 30. Briffa M, Lane SM. 2017 The role of skill in animal contests: a neglected component of fighting ability. *Proceedings of the Royal Society B: Biological Sciences* **284**, 20171596. (doi:10.1098/rspb.2017.1596)
- 31. Parker GA. 1974 Assessment strategy and the evolution of fighting behaviour. *Journal of Theoretical Biology* **47**, 223–243. (doi:10.1016/0022-5193(74)90111-8)
- 32. Elwood RW, Arnott G. 2012 Understanding how animals fight with Lloyd Morgan's canon. *Animal Behaviour* **84**, 1095–1102. (doi:10.1016/j.anbehav.2012.08.035)

- 33. Moors A, Ellsworth PC, Scherer KR, Frijda NH. 2013 Appraisal theories of emotion: State of the art and future development. *Emotion Review* **5**, 119–124. (doi:10.1177/1754073912468165)
- 34. Scherer KR. 2001 Appraisal Considered as a Process of Multilevel Sequential Checking. In *Appraisal processes in emotion: Theory, methods, research* (eds Scherer, KR, Schorr A, Johnstone T), pp. 92–120. New York: Oxford University Press.
- 35. Moors A. 2013 On the Causal Role of Appraisal in Emotion. *Emotion Review* **5**, 132–140. (doi:10.1177/1754073912463601)
- 36. Uusberg A, Taxer JL, Yih J, Uusberg H, Gross JJ. 2019 Reappraising reappraisal. *Emotion Review* **11**, 267–282. (doi:10.1177/1754073919862617)
- 37. Veissier I, Boissy A, Désiré L, Greiveldinger L. 2009 Animals' emotions: studies in sheep using appraisal theories. *Animal Welfare* **18**, 347–354.
- 38. Lerner JS, Li Y, Valdesolo P, Kassam KS. 2015 Emotion and Decision Making. *Annual Review of Psychology* **66**, 799–823. (doi:10.1146/annurev-psych-010213-115043)
- 39. Damasio A, Carvalho GB. 2013 The nature of feelings: Evolutionary and neurobiological origins. *Nature Reviews Neuroscience* **14**, 143–152. (doi:10.1038/nrn3403)
- 40. Nesse RM, Ellsworth PC. 2009 Evolution, Emotions, and Emotional Disorders. *American Psychologist* **64**, 129–139. (doi:10.1037/a0013503)
- 41. Winkielman P, Berridge KC. 2004 Unconscious emotion. *Current Directions in Psychological Science* **13**, 120–123. (doi:10.1111/j.0963-7214.2004.00288.x)
- 42. Winkielman P, Berridge KC, Wilbarger JL. 2005 Unconscious affective reactions to masked happy versus angry faces influence consumption behavior and judgments of value. *Personality and Social Psychology Bulletin* **31**, 121–135. (doi:10.1177/0146167204271309)
- 43. Birch J, Schnell AK, Clayton NS 2020. Dimensions of animal consciousness. *Trends in Cognitive Sciences* **24**, 789–801. (doi:10.1016/j.tics.2020.07.007)
- 44. Matheson SM, Asher L, Bateson M. 2008. Larger, enriched cages are associated with 'optimistic' response biases in captive European starlings (Sturnus vulgaris). *Applied Animal Behaviour Science* **109**, 374–383. (doi:10.1016/j.applanim.2007.03.007)
- 45. Papciak J, Popik P, Fuchs E, Rygula R. 2013 Chronic psychosocial stress makes rats more "pessimistic" in the ambiguous-cue interpretation paradigm. *Behavioural Brain Research* **256**, 305–310. (doi:10.1016/j.bbr.2013.08.036)
- 46. Neville V, Nakagawa S, Zidar J, Paul ES, Lagisz M, Bateson M, Løvlie H, Mendl M. 2020 Pharmacological manipulations of judgement bias: A systematic review and meta-analysis. *Neuroscience and Biobehavioral Reviews* **108**, 269–286. (doi:10.1016/j.neubiorev.2019.11.008)
- 47. Crump A, Arnott G, Bethell EJ. 2018 Affect-driven attention biases as animal welfare indicators: Review and methods. *Animals* **8**, 136 (doi:10.3390/ani8080136)
- 48. Bethell EJ. 2015 A "How-To" Guide for Designing Judgment Bias Studies to Assess Captive Animal Welfare. *Journal of Applied Animal Welfare Science* **18**, S18–S42. (doi:10.1080/10888705.2015.1075833)
- 49. Mendl M, Burman OHP, Parker RMA, Paul ES. 2009 Cognitive bias as an indicator of animal emotion and welfare: Emerging evidence and underlying mechanisms. *Applied Animal Behaviour Science* **118**, 161–181. (doi:10.1016/j.applanim.2009.02.023)

- 50. Takatsu-Coleman AL, Patti CL, Zanin KA, Zager A, Carvalho RC, Borçoi AR, ... Frussa-Filho R. 2013 Short-term social isolation induces depressive-like behaviour and reinstates the retrieval of an aversive task: Mood-congruent memory in male mice?. *Journal of Psychiatry & Neuroscience* 38, 259. (doi:10.1503/jpn.120050)
- 51. Jensen MB, Pedersen LJ. 2008 Using motivation tests to assess ethological needs and preferences. *Applied Animal Behaviour Science* **113**, 340–356. (doi: 10.1016/j.applanim.2008.02.001)
- 52. Kirkden RD, Pajor EA. 2006 Using preference, motivation and aversion tests to ask scientific questions about animals' feelings. *Applied Animal Behaviour Science* **100**, 29–47. (doi:10.1016/j.applanim.2006.04.009)
- 53. Maynard Smith J. 1974 The theory of games and the evolution of animal conflicts. *Journal of Theoretical Biology* **47**, 209–221. (doi:10.1016/0022-5193(74)90110-6)
- 54. Mesterton-Gibbons M, Marden JH, Dugatkin LA. 1996 On wars of attrition without assessment. *Journal of Theoretical Biology* **181**, 65–83. (doi:10.1006/jtbi.1996.0115)
- 55. Enquist M, Leimar O. 1983 Evolution of Fighting Behaviour: Decision Rules and Assessment of Relative Strength. *Journal of Theoretical Biology* **102**, 387–410.
- 56. Pinto NS, Palaoro AV, Peixoto PEC. 2019 All by myself? Meta-analysis of animal contests shows stronger support for self than for mutual assessment models. *Biological Reviews* **94**, 12509. (doi:10.1111/brv.12509)
- 57. Fuxjager MJ, Mast G, Becker EA, Marler CA. 2009 The "home advantage" is necessary for a full winner effect and changes in post-encounter testosterone. *Hormones and Behavior* **56**, 214–219. (doi:10.1016/j.yhbeh.2009.04.009)
- 58. Kemp DJ, Wiklund C. 2004 Residency effects in animal contests. *Proceedings of the Royal Society of London. Series B: Biological Sciences* **271**, 1707–1711. (doi:10.1098/rspb.2004.2775)
- 59. Hobson EA. 2020 Differences in social information are critical to understanding aggressive behavior in animal dominance hierarchies. *Current Opinion in Psychology* **33**, 209–215. (doi:10.1016/j.copsyc.2019.09.010)
- 60. Millsopp S, Laming P. 2008 Trade-offs between feeding and shock avoidance in goldfish (Carassius auratus). *Applied Animal Behaviour Science* **113**, 247–254. (doi:10.1016/j.applanim.2007.11.004)
- 61. Duncan IJH, Wood-Gush DGM. 1971 Frustration and aggression in the domestic fowl. *Animal Behaviour* **19**, 500–504. (doi:10.1016/S0003-3472(71)80104-5)
- 62. Papini MR, Dudley RT. 1997 Consequences of surprising reward omissions. *Review of General Psychology* **1**, 175–197. (doi:10.1037/1089-2680.1.2.175)
- 63. Vindas MA, Folkedal O, Kristiansen TS, Stien LH, Braastad BO, Mayer I, Øverli Ø. 2012 Omission of expected reward agitates Atlantic salmon (Salmo salar). *Animal Cognition* **15**, 903–911. (doi:10.1007/s10071-012-0517-7)
- 64. Darden SK, May MK, Boyland NK, Dabelsteen T. 2019 Territorial defense in a network: audiences only matter to male fiddler crabs primed for confrontation. *Behavioral Ecology* **30**, 336–340. (doi:10.1093/beheco/ary169)
- 65. Miles MC, Fuxjager MJ. 2019 Social context modulates how the winner effect restructures territorial behaviour in free-living woodpeckers. *Animal Behaviour* **150**, 209–218. (doi:10.1016/j.anbehav.2019.02.011)
- 66. Montroy K, Loranger MJ, Bertram SM. 2016 Male crickets adjust their aggressive behavior when a female is present. *Behavioural Processes* **124**, 108–114. (doi:10.1016/j.beproc.2015.11.003)

- 67. Oliveira RF, Lopes M, Carneiro LA, Canário AVM. 2001 Watching fights raises fish hormone levels. *Nature* **409**, 475. (doi:10.1038/35054128)
- 68. Cabanac M. 1992 Pleasure: the common currency. *Journal of Theoretical Biology* **155**, 173–200. (doi:10.1016/S0022-5193(05)80594-6)
- 69. Levy DJ, Glimcher PW. 2012 The root of all value: A neural common currency for choice. *Current Opinion in Neurobiology* **22**, 1027–1038. (doi:10.1016/j.conb.2012.06.001)
- 70. Elwood RW, Appel M. 2009 Pain experience in hermit crabs? *Animal Behaviour* 77, 1243–1246. (doi:10.1016/j.anbehav.2009.01.028)
- 71. Payne RJH, Pagel M. 1996 Escalation and time costs in displays of endurance. *Journal of Theoretical Biology* **183**, 185–193. (doi:10.1006/jtbi.1996.0212)
- 72. Payne RJH, Pagel M. 1997 Why do animals repeat displays? *Animal Behaviour* **54**, 109–119. (doi:10.1006/anbe.1996.0391)
- 73. Payne RJH. 1998 Gradually escalating fights and displays: The cumulative assessment model. *Animal Behaviour* **56**, 651–662. (doi:10.1006/anbe.1998.0835)
- 74. Taylor PW, Elwood RW. 2003 The mismeasure of animal contests. *Animal Behaviour* **65**, 1195–1202. (doi:10.1006/anbe.2003.2169)
- 75. Camerlink I, Turner SP, Farish M, Arnott G. 2017 The influence of experience on contest assessment strategies. *Scientific Reports* **7**, 1–10. (doi:10.1038/s41598-017-15144-8)
- 76. Chapin KJ, Enrique P, Peixoto C, Briffa M. 2019 Further mismeasures of animal contests: a new framework for assessment strategies. *Behavioral Ecology* **30**, 1177–1185 (doi:10.1093/beheco/arz081)
- 77. Mesterton-Gibbons M, Heap SM. 2014 Variation between self- and mutual assessment in animal contests. *The American Naturalist* **183**, 199–213. (doi:10.1086/674443)
- 78. Garcia MJ, Paiva L, Lennox M, Sivaraman B, Wong SC, Earley RL. 2012 Assessment strategies and the effects of fighting experience on future contest performance in the green anole (Anolis carolinensis). *Ethology* **118**, 821–834. (doi:10.1111/j.1439-0310.2012.02072.x)
- 79. Hsu Y, Lee SP, Chen MH, Yang SY, Cheng KC. 2008 Switching assessment strategy during a contest: fighting in killifish Kryptolebias marmoratus. *Animal Behaviour* **75**, 1641–1649. (doi:10.1016/j.anbehav.2007.10.017)
- 80. Morrell LJ, Backwell PRY, Metcalfe NB. 2005 Fighting in fiddler crabs Uca mjoebergi: What determines duration? *Animal Behaviour* **70**, 653–662. (doi:10.1016/j.anbehav.2004.11.014)
- 81. Blanchette I, Richards A. 2010 The influence of affect on higher level cognition: A review of research on interpretation, judgement, decision making and reasoning. *Cognition & Emotion* **24**, 561–595. (doi:10.1080/02699930903132496)
- 82. Mackie DM, Worth LT. 1989 Processing Deficits and the Mediation of Positive Affect in Persuasion. *Journal of Personality and Social Psychology* **57**, 27–40. (doi:10.1037/0022-3514.57.1.27)
- 83. Worth LT, Mackie DM. 1987 Cognitive mediation of positive affect in persuasion. *Social Cognition* **5**, 76–94. (doi:10.1521/soco.1987.5.1.76)
- 84. Hutchinson JMC, Gigerenzer G. 2005 Simple heuristics and rules of thumb: Where psychologists and behavioural biologists might meet. *Behavioural Processes* **69**, 97–124. (doi:10.1016/j.beproc.2005.02.019)
- 85. Harmon-Jones E, Harmon-Jones C, Amodio DM, Gable PA. 2011 Attitudes toward emotions. *Journal of Personality and Social Psychology* **101**, 1332. (doi:10.1037/a0024951)

- 86. Cabral JCC, de Almeida RMM. 2019 Effects of anger on dominance-seeking and aggressive behaviors. *Evolution and Human Behavior* **40**, 23–33. (doi:10.1016/j.evolhumbehav.2018.07.006)
- 87. Veenstra L, Bushman BJ, Koole SL. 2018 The facts on the furious: a brief review of the psychology of trait anger. *Current Opinion in Psychology* **19**, 98–103. (doi:10.1016/j.copsyc.2017.03.014)
- 88. Carver CS, Harmon-Jones E. 2009 Anger is an approach-related affect: evidence and implications. *Psychological Bulletin* **135**, 183. (doi:10.1037/a0013965)
- 89. Elwood RW, Wood KE, Gallagher MB, Dick, JTA 1998. Probing motivational state during agonistic encounters in animals. *Nature* **393**(6680), 66–68. (doi:10.1038/29980)
- 90. Koch M 1999. The neurobiology of startle. *Progress in Neurobiology* **59**(2), 107–128. (doi:10.1016/S0301-0082(98)00098-7)
- 91. Winslow JT, Parr LA, Davis M 2002. Acoustic startle, prepulse inhibition, and fear-potentiated startle measured in rhesus monkeys. *Biological Psychiatry* **51**(11), 859–866. (doi:10.1016/S0006-3223(02)01345-8)
- 92. Fawcett TW, Johnstone RA. 2010 Learning your own strength: winner and loser effects should change with age and experience. Proce*edings of the Royal Society B: Biological Sciences* **277**, 1427–1434. (doi:10.1098/rspb.2009.2088)
- 93. Mesterton-Gibbons M. 1999 On the evolution of pure winner and loser effects: A game-theoretic model. *Bulletin of Mathematical Biology* **61**, 1151–1186. (doi:10.1006/bulm.1999.0137)
- 94. Hsu Y, Earley RL, Wolf LL. 2006 Modulation of aggressive behaviour by fighting experience: Mechanisms and contest outcomes. *Biological Reviews of the Cambridge Philosophical Society* **81**, 33–74. (doi:10.1017/S146479310500686X)
- 95. Hsu Y, Lee IH, Lu CK. 2009 Prior contest information: Mechanisms underlying winner and loser effects. *Behavioral Ecology and Sociobiology* **63**, 1247–1257. (doi:10.1007/s00265-009-0791-9)
- 96. Rutte C, Taborsky M, Brinkhof MWG. 2006 What sets the odds of winning and losing? *Trends in Ecology and Evolution* **21**, 16–21. (doi:10.1016/j.tree.2005.10.014)
- 97. Paul ES, Harding EJ, Mendl M. 2005 Measuring emotional processes in animals: The utility of a cognitive approach. *Neuroscience and Biobehavioral Reviews* **29**, 469–491. (doi:10.1016/j.neubiorev.2005.01.002)
- 98. Everaert J, Podina IR, Koster EHW. 2017 A comprehensive meta-analysis of interpretation biases in depression. *Clinical Psychology Review* **58**, 33–48. (doi:10.1016/j.cpr.2017.09.005)
- 99. Hirsch CR, Meeten F, Krahé C, Reeder C. 2016 Resolving Ambiguity in Emotional Disorders: The Nature and Role of Interpretation Biases. *Annual Review of Clinical Psychology* **12**, 281–305. (doi:10.1146/annurev-clinpsy-021815-093436)
- 100. Stuijfzand S, Creswell C, Field AP, Pearcey S, Dodd H. 2018 Research Review: Is anxiety associated with negative interpretations of ambiguity in children and adolescents? A systematic review and meta-analysis. *Journal of Child Psychology and Psychiatry* **59**, 1127–1142. (doi:10.1111/jcpp.12822)
- 101. Harding EJ, Paul ES, Mendl M. 2004 Cognitive bias and affective state. *Nature* **427**, 312. (doi:10.1038/427312a)
- 102. Kasumovic MM, Elias DO, Sivalinghem S, Mason AC, Andrade MCB. 2010 Examination of prior contest experience and the retention of winner and loser effects. *Behavioral Ecology* **21**, 404–409. (doi:10.1093/beheco/arp204)

- 103. Barker TH, George RP, Howarth GS, Whittaker AL. 2017 Assessment of housing density, space allocation and social hierarchy of laboratory rats on behavioural measures of welfare. *PLOS ONE* **12**. (doi:10.1371/journal.pone.0185135)
- 104. Horback KM, Parsons TD. 2019 Judgement bias testing in group-housed gestating sows. *Behavioural Processes* **159**, 86–92. (doi:10.1016/j.beproc.2018.12.021)
- 105. Schino G, Massimei R, Pinzaglia M, Addessi E. 2016 Grooming, social rank and 'optimism' in tufted capuchin monkeys: a study of judgement bias. *Animal Behaviour* **119**, 11–16. (doi:10.1016/j.anbehav.2016.06.017)
- Rogers L, Sales E, Shamsi S, Kopf RK, Freire R. 2020 Aggressive encounters lead to negative affective state in fish. *PLOS ONE* 15, e0231330. (doi:10.1371/journal.pone.0231330)
- 107. May ME. 2011 Aggression as positive reinforcement in people with intellectual disabilities. *Research in Developmental Disabilities* **32**, 2214–2224. (doi:10.1016/j.ridd.2011.05.029)
- 108. Falkner AL, Grosenick L, Davidson TJ, Deisseroth K, Lin D. 2016 Hypothalamic control of male aggression-seeking behavior. *Nature Neuroscience* **19**, 596–604. (doi:10.1038/nn.4264)
- 109. Martínez M, Guilli~n-Salazar F, Salvador A, Simon VM. 1995 Successful Intermale Aggression and Conditioned Place Preference in Mice. *Physiology & Behavior* **58**, 323–328 (doi:10.1016/0031-9384(95)00061-M)
- 110. Meisel RL, Joppa MA. 1994 Conditioned place preference in female hamsters following aggressive or sexual encounters. *Physiology & Behavior* **56**, 1115–1118. (doi:10.1016/0031-9384(94)90352-2)
- 111. Farrell WJ, Wilczynski W. 2006 Aggressive experience alters place preference in green anole lizards, Anolis carolinensis. *Animal Behaviour* **71**, 1155–1164. (doi:10.1016/j.anbehav.2005.10.006)
- 112. Reichert MS, Quinn JL. 2017 Cognition in Contests: Mechanisms, Ecology, and Evolution. *Trends in Ecology and Evolution* **32**, 773–785. (doi:10.1016/j.tree.2017.07.003)
- 113. George JM, Dane E. 2016 Affect, emotion, and decision making. *Organizational Behavior and Human Decision Processes* **136**, 47–55. (doi:10.1016/j.obhdp.2016.06.004)
- 114. Västfjäll D, Slovic P, Burns WJ, Erlandsson A, Koppel L, Asutay E, Tinghög G. 2016 The arithmetic of emotion: Integration of incidental and integral affect in judgments and decisions. *Frontiers in Psychology* **7**, 325. (doi:10.3389/fpsyg.2016.00325)
- 115. Wyer Jr RS, Dong P, Huang X, Huang Z, Wan LC. 2019 The Effect of Incidental Emotions on Judgments and Behavior in Unrelated Situations: A Review. *Journal of the Association for Consumer Research* **4**, 198–207. (doi:10.1086/701889)
- 116. Schwarz N, Clore GL. 1983 Mood, misattribution, and judgements of well-being informative and directive functions of affective states. *Journal of Personality and Social Psychology* **45**, 513–523. (doi:10.1037/0022-3514.45.3.513)
- 117. Bateson M, Desire S, Gartside SE, Wright GA. 2011 Agitated honeybees exhibit pessimistic cognitive biases. *Current Biology* **21**, 1070–1073. (doi:10.1016/j.cub.2011.05.017)

- 118. Bergman M, Olofsson M, Wiklund C. 2010 Contest outcome in a territorial butterfly: the role of motivation. *Proceedings of the Royal Society B: Biological Sciences* **277**, 3027–3033. (doi:10.1098/rspb.2010.0646)
- 119. Zhang S, Ai H, Li D. 2019 Mating changes a male contestant from a loser to a winner in male—male competition in a wolf spider. *Biological Journal of the Linnean Society* **128**, 83–92. (doi:10.1093/biolinnean/blz091)
- 120. Reddon AR, Dey CJ, Balshine S. 2019 Submissive behaviour is mediated by sex, social status, relative body size and shelter availability in a social fish. *Animal Behaviour* **155**, 131–139. (doi:10.1016/j.anbehav.2019.06.026)
- 121. Parrott JL, Metcalfe CD. 2018 Nest-defense behaviors in fathead minnows after lifecycle exposure to the antidepressant venlafaxine. *Environmental Pollution* **234**, 223–230. (doi:10.1016/j.envpol.2017.11.049)
- 122. Perreault HA, Semsar K, Godwin J. 2003 Fluoxetine treatment decreases territorial aggression in a coral reef fish. *Physiology & Behavior* **79**, 719–724. (doi:10.1016/S0031-9384(03)00211-7)
- 123. Niemelä PT, Santostefano F. 2015 Social carry-over effects on non-social behavioral variation: mechanisms and consequences. *Frontiers in Ecology and Evolution* **3**, 49. (doi:10.3389/fevo.2015.00049)
- 124. Van der Harst JE, Baars AM, Spruijt BM 2005. Announced rewards counteract the impairment of anticipatory behaviour in socially stressed rats. *Behavioural Brain Research* **161**, 183–189.
- 125. Boggiani L, Addessi E, Schino G. 2018 Receiving aggression triggers attention bias in tufted capuchin monkeys. *Animal Behaviour* **146**, 173–180. (doi:10.1016/j.anbehav.2018.10.021)
- 126. Clore, G. L., Wyer, R. S., Dienes, B. P. A., Gasper, K., Gohm, C., & Isbell, L. (2001). Affective feelings as feedback: Some cognitive consequences. In L. L. Martin & G. L. Clore (Eds.), Theories of Mood and Cognition: A User's Guidebook (pp. 27–62). Mahwah, NJ: Erlbaum.
- 127. Bakamitsos GA 2006. A cue alone or a probe to think? The dual role of affect in product evaluations. *Journal of Consumer Research* **33**, 403–412.

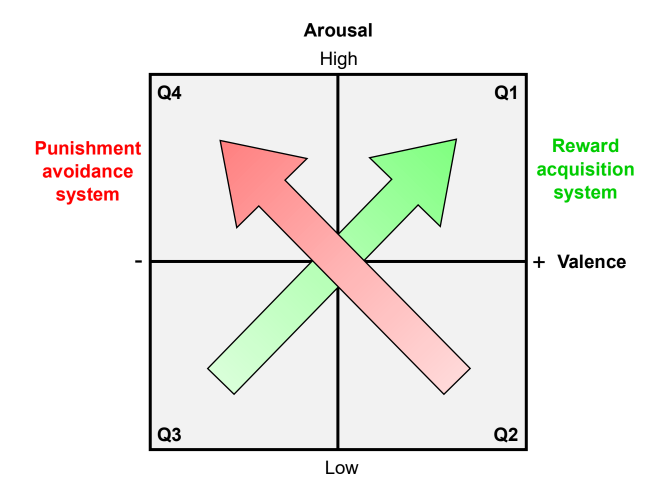


Figure 1. Valence and arousal define affective states (grey box), which encompasses emotions and moods [6]. Moving from Q3-Q1 is increasingly appetitive; Q2-Q4 is increasingly aversive.

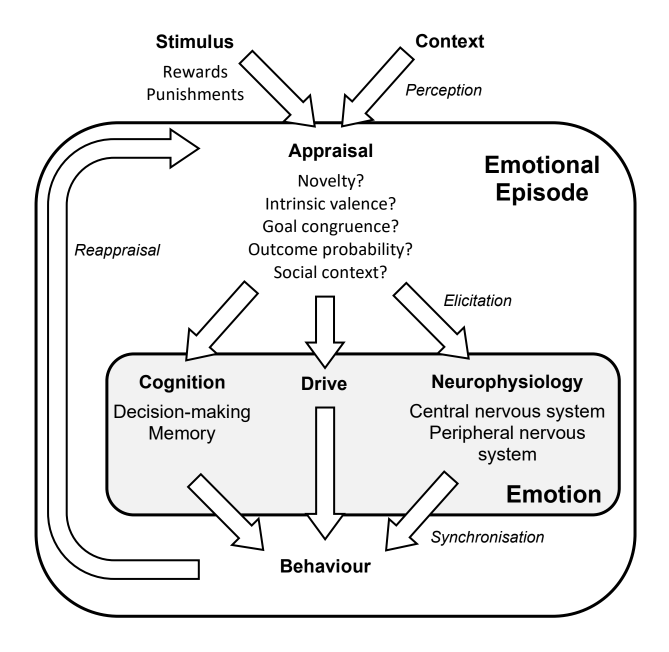


Figure 2. An emotional episode (white box). Appraisals of stimuli, their context, and their personal significance elicit the emotion (grey box), whose components include cognition, drive, and neurophysiology. These components govern the expression of behaviour. Conscious "feelings" are another potential component, but not essential.

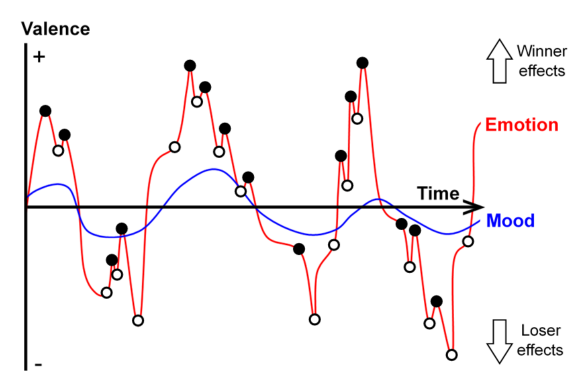


Figure 3. Cumulative emotional valence determines mood [11] (manifested in aggression). Considering only integral (objectively contest-relevant) influences, white dots are wins and black dots are losses. Considering both integral and incidental (objectively contest-irrelevant) influences, white dots are rewards and black dots are punishments.

Table 1. Major predictions and outstanding questions that arise from applying emotion theory to animal contests

Major Predictions

Contest appraisals cover more variables than traditionally recognised (i.e. RV and RHP)

Positive affective states induce selfassessment; negative states induce mutual assessment

Winner effects are associated with optimistic responses to judgement bias tasks; loser effects are associated with pessimistic responses

Incidental affective influences modify contest behaviour

Humans and animals share rules that increase the likelihood of incidental influences (e.g. concurrence, ambiguity, and link to moods)

The above predictions apply only to animals with a central nervous system

Outstanding Questions

Are contest appraisals sequential? Do untested human appraisals modify contest dynamics?

Do assessment strategies vary with affective state? How might this influence the outcome?

What neurocognitive mechanisms underpin judgement bias? Are they equivalent to the mechanisms underpinning winner/loser effects

Do incidental affective states commonly impact contests in nature? Why evolve a generalised (rather than domain-specific) affective system?

What mechanisms minimise incidental influences? How do these affect fitness?

Do all animals with a central nervous system have affective states? Are contest dynamics fundamentally different in organisms without a central nervous system?