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Why it hurts: with freedom comes the biological need for pain

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Abstract

We argue that pain is not needed to protect the body from damage unless the organism is able to make free choices in action selection. Then pain (including its affective and evaluative aspects) provides a necessary prioritising motivation to select actions expected to avoid it, whilst leaving the possibility of alternative actions to serve potentially higher priorities. Thus, on adaptive grounds, only organisms having free choice over action selection should experience pain. Free choice implies actions must be selected following appraisal of their effects, requiring a predictive model generating estimates of action outcomes. These features give organisms anticipatory behavioural autonomy (ABA), for which we propose a plausible system

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using an internal predictive model, integrated into a system able to produce the qualitative and affective aspects of pain. Our hypothesis can be tested using behavioural experiments designed to elicit trade-off responses to novel experiences for which algorithmic (automaton) responses might be inappropriate. We discuss the empirical evidence for our hypothesis among taxonomic groups, showing how testing for ABA guides thinking on which groups might experience pain. It is likely that all vertebrates do and plausible that some invertebrates do (decapods, cephalopods and at least some insects).

1 Introduction - the three faces of pain

It is still common for pain to be explained as a mechanism for protecting body parts from acute injury, even though it is accepted that reflex withdrawal is often sufficient for that purpose. Our question here is not about immediate responses to nociception; we seek a biological explanation for pain as defined by the International Association for the Study of Pain (IASP): “An unpleasant sensory and emotional experience associated with, or resembling that associated with, actual or potential tissue damage” – (Raja et al. 2020). For that we need to define several terms relating to emotional experience.

1.1 Working definitions

- *Sentience*: Crump et al. (2022) provide an excellent definition: “*Sentience is the capacity to feel. Understood broadly, sentience encompasses all felt experiences, including sensory experiences (e.g. visual, auditory, tactile, olfactory) as well as (for example) feelings of warmth, comfort, fatigue, hunger, thirst, boredom, excitement, distress, anxiety, pain, pleasure and joy. This capacity to feel should be distinguished from other, related capacities: a sentient being might not be able to reflect on its feelings or to understand others’ feelings*”. Sentience is one of several dimensions of consciousness, though often the words sentience and consciousness

are used interchangeably. Sentience is self-evidently needed for pain, but consciousness might not be: this is the crux of contention over whether animals of different kinds can feel pain, so we agree with Browning and Birch (2022), that a clear distinction is essential.

- *Feelings (qualia)*: are subjective (phenomenal) experiences, including pain. We cannot detect feelings by direct empirical study because they are definitively subjective and so bounded by the agent experiencing them: this fact has led to much philosophical debate over whether they even exist (Tye 2021). It is useful to consider them as emergent phenomena generated by (brain) information processing. Recently, Clark et al. (2019) made that idea concrete using the predictive processing theory of perception and consciousness, concluding that qualia are intermediate-level models generated by “Bayesian brains”. More generally, we take qualia to be mental constructs that can be functional and, crucially here, motivational (Hall 2008, Fulkerson 2021).
- *Consciousness*: includes several dimensions additional to sentience (Birch et al. 2020b), though sentience is one of its requirements (see Nani et al. 2021). Consciousness is notoriously difficult to identify and study scientifically, partly because it is subjective, but also because we still have no consensus on its definition (Michel 2020). Of greatest significance here are the dimensions of a) self-awareness, derived from a ‘meta-perception’ system that perceives the perception of internal and external stimuli and b) the integration of perception from internal and external stimuli, along with memory and any available outputs from internal generative models, to form a coherent whole ‘mental image’. Pain requires sentience because it is a feeling and it requires integration because it operates at the whole-organism level, but pain might not require the other dimensions of consciousness, though researchers differ over meta-perception: e.g. Key et al. (2021; 2022) consider it the

primary requirement for pain experience.

- *Emotion (affect)*: has been implicated in appraisal (Scherer et al. 2001), for action selection (Mendl and Paul 2020) and also direct motivation (Barlassina and Hayward 2019). Helm (2002) defined emotions as “not mere phenomenal states but evaluative responses to one’s situation”, though recognising that “emotions are feelings” as well. Affect is usually regarded as a top-level (system) phenomenon that sets the internal context for information processing and action selection: an internal psychological milieu (via neurohormones) modulating the parameters of judgement. To that extent emotions are evaluative in *function*. Confusion arises because we know from introspection that emotions have associated feelings (some say they *are* feelings): there is definitely something it is like to be joyful or disgusted, etc.. It is useful here to consider affect as a summarising self-appraisal of an organism’s situation as represented by an internal model, one that exists at the level of the integrated whole of the organism (including physiological responses and motor expressions) (Scherer 2022).

1.2 Approaches to pain

Following the pioneering model of Melzack and Casey (1968), pain is broadly recognised to have three dimensions: sensory-discriminative, affective-motivational and cognitive-evaluative (Corns 2014). Pains are feelings with perceptual specificity of location, intensity and quality that are generally noxious, draw attention and motivate those experiencing them to avoid them in future. A simple animal such as the protist *Stentor* can withdraw and guard itself following a noxious stimulus but we doubt it is capable of ‘feeling’ anything. Similarly, *Cnidarians* are usually assumed non-sentient, though capable of sensitisation (an escalating response to a stimulus (e.g. Cheng 2021)). By definition (Crump et al. 2022), feelings require sentience to create a phenomenal experience, so only sentient organisms can feel pain.

There are three broad approaches to establishing whether an organism can feel pain. Firstly behavioural responses, especially in experimental arrangements, can match our expectation for an organism feeling pain, but since pain is necessarily subjective, this can never provide a definitive answer. Most taking this approach carefully limit their interpretation as: observed behaviours are consistent with expectations for pain (Elwood 2019; 2021). Secondly, we may seek the neural circuitry thought to be necessary for pain (as in Key 2015, Key and Brown 2018, Key et al. 2021). However, we do not yet know what circuits are necessary and rely on either broad categories of processing, e.g. that there must be a subsystem to monitor and create awareness of the internal state of the perception system, or specific hypotheses about parts of the necessary circuits, e.g. that they must include feed-forward and comparator elements (Key et al. 2021). The problem with the former is that it can be too broad, leaving answers unclear. The problem with the latter is that any system proposed as necessary for generating the subjective feeling of pain remains an untested hypothesis until we know what is necessary. The third approach, which has received remarkably little attention, asks which evolved system (or behaviour) needs the subjective feeling of pain in order to work. If we can identify a system that requires pain for its functioning, together with the organisms that possess that system, then we might reasonably presume they will feel pain. This is the (philosophically functionalist) approach we adopt here.

The ‘imperativist’ account of pain (Hall 2008, Klein 2007, Martinez 2011; 2015) and the (related) realisation that pain could be interpreted as a part of a homeostatic regulation system for the body both provide valuable context. The imperativist account is that pain is not information about bodily damage or its potential, but rather is a command or motivation for taking action to protect the body from damage (actual or potential). This idea has been corroborated by animal studies showing lasting changes in motivation and behaviour following noxious experiences (Sneddon et al. 2014). At the heart of our present thesis is the realisation that this command may be functional only

for organisms that have freedom to choose among a range of options for action, that is, only if action-selection mechanisms are not pre-programmed (algorithmic), but rather are the result of the *evaluation* of possible action. Importantly, pain is not required if action selection is strictly reactive with no anticipation of possible futures — in such cases, a rigid relation between perception and action is always sufficient to appropriately respond to nociception.

For organisms able to anticipate future states, action selection is the result of an internally generated decision based on modelling and evaluating possible future states. The decision is based on the optimisation of some (hedonic) utility currency and is a free choice. We term this process *proactive autonomy* and organisms possessing the freedom it implies would benefit from a strong motivation to attend to injury when evaluating all the options. Conversely, an organism whose action selection is determined by a state-dependent information processing algorithm (if in state S do X), however complicated, gains no advantage from such motivation as pain could provide. The algorithm would instantiate the necessary and sufficient internal information for action to be taken, whether it is to protect against (further) injury, or to continue the current behaviour (e.g. fighting). The information constituting this algorithm could sufficiently be obtained by inheritance and may include sensor and activation threshold shifts in response to repetition of stimulus, enabling habituation or non-associative learning, without the need for modelling and evaluation. Such an algorithm would facilitate what we call *reactive autonomy* (Fig 1).

An organism that has an internal model predicting accessible future states could in principle initiate behaviours that do not depend on reacting to external stimuli. They could therefore be capable of action selection in which an appraisal of the desirability of future states enters the decision-making. We term this capability *anticipatory behavioural autonomy* (ABA). It is this capability that most readily justifies a cognitive-evaluative dimension to pain, since that dimension describes a comparison among the

expected outcomes of available actions. Pain’s role would be to motivate the organism to prioritise attention towards the source and cause of the pain (i.e. salience), but would leave open the possibility of attending to a more pressing matter, such as escape.

In this view, pain is part of the organism’s behaviour control system. In general, control is constraint (see Montévil and Mossio (2015)) and all constraint is the result of organising information (Bich et al. 2020, Farnsworth et al. 2013, Farnsworth 2022, Montévil and Mossio 2015, Mossio et al. 2016). This information is not merely the signal¹ of nociception, but crucially includes the causal structure of the cybernetic system responsible for the organism’s response. The operation of cybernetic systems that determine action selection is entirely one of information processing, i.e. computation, coupled to the physical world by actuators that physically perform the actions. Understanding this information basis for control is important in identifying the autonomy required for ABA .

2 Understanding systems that might use pain

It is widely thought that sentience requires an internal model of the self: “*subjective experience arises from [...] an integrated simulation of the state of the animal’s own mobile body within the environment*” – (Barron and Klein 2016). This internal model is an essential component of computer representations of animals in welfare research, conceived with widely differing perspectives (e.g. Budaev et al. 2020, Key et al. 2022). Within philosophy, such models are intrinsically implied by representational accounts of pain and are necessary for evaluative accounts beyond the strictly reactive (i.e. whenever options are to be evaluated for their future consequences). A self-model was conceived by Farnsworth (2017) as part of a mechanistic explanation for free choice in general

¹We use ‘signal’ in the standard engineering sense of variation indicating data concerning its source, rather than the special sense of an organism-generated sign conveying information, used in the study of animal communication.

systems, including organisms and AI systems and a conceptually similar system was proposed by Ridderinkhof (2017). The self-model forms part of an allostatic (predictive homeostatic) system that justifies and makes concrete the motivational aspect of felt experiences.

We propose that pain provides for evaluation of outcomes in anticipatory action selection via a common currency throughout the control system, one that can command salience and encode information in its qualitative character (as Cabanac (1992) describes in relation to pleasure). This strongly suggests a felt experience, implying sentience, but not necessarily the self-awareness, derived from a ‘meta-perception’ system (Cunningham 2001), as thought essential by Key and Brown (2018), and argued for by Brown et al. (2021) in response to Birch et al. (2020a). Higher-order-thought theories of consciousness imply that for awareness of pain there must be a subsystem (module) that ‘listens in’ to the universal signals and reports to a hypothetical executive centre, supposed to be the ‘theatre of consciousness’. This idea has been criticised (e.g. Dennett 1991) for falling into the ‘homunculus fallacy’ (Baltzer-Jaray 2018) and certainly strays from the principle of parsimony. We believe that the formation of an internal representation of the self can produce a phenomenal state with intrinsic evaluative character and that this is sufficient to explain the qualitative feeling of e.g. pain. What it is like to be in some degree of pain is the same as what it is like to have a particular self-model result. That is not a model output, since the result is a state of the internal model. In turn, the whole organism is in that state: a particular phenomenal state we term its Q-state. In this view, pain is a dispositional state of an organism having a predictive model of possible actions and using feelings as the arbiter of choice among them.

2.1 Autonomy and Action Selection

Autonomy is the property of a system undergoing state changes caused by internal events, so that it is at least partly controlled by internalised information rather than entirely

by external causes. Action selection is the resolution of conflicts between competing behavioural options. We define *proactive autonomy* as the ability of an agent to act in the physical world in a way that is determined by the free choice of the system. Since there is a choice, there must be at least two viable options and some sort of action-selection system that implements (on average) the expected fitness-enhancing decision, which in turn implies a system-level utility function to be maximised by the choice. Proactive autonomy implies proximate agent causation: the agent is the causal source of the action. Organisms possessing proactive autonomy display the ability to respond differently to the same external stimulus depending on their independent assessment, enabling appropriate responses to be made to novel circumstances and to take account of future possibilities such as deferred rewards. It is the freedom of choice, enabled by a-priori indeterminacy of outcome, that requires a normative (reward/punishment) evaluation of possible outcomes. The indeterminacy of outcome does not mean that it is random; rather, it is contingent upon some internal computation that is not preprogrammed. Proactive action selection solves an optimisation problem, for which it needs a common currency Y to represent the desirability of each competing behaviour. An arbitrary set of actions can be compared to find which maximises Y given the conditions. Y then acts as an objective function (in the optimality-theory sense), the maximisation of which will be the ‘goal’. The idea of common currency in this context was pioneered by McFarland et al. (1975), interpreted as biological fitness in the ecological context by McNamara and Houston (1986) and as pleasure by Cabanac (1992), who extended it to an explanation for emotion (Cabanac 2002).

In homeostasis (the most basic form of goal-dependent control shown in Fig. 1.a) the goal is embodied as a set-point. Different perception signals (S1 and S2) can be ‘hard wired’ to modulate one another to achieve a rudimentary form of action selection Fig. 1.b). If only one action is possible (e.g. in the escape reaction of *Paramecium* (Brette 2021)), then action-selection does not arise, but a homeostatic system comparing

multiple perceived signals with their corresponding goals may be used to switch the action **ON** or **OFF** using summation, or a winner takes all circuit (Tymoshchuk and Shatnyi 2015) (Fig. 1.c). Single-celled organisms are equipped with these sort of action-selection systems, e.g. for selecting between tumbling and swimming in the chemokinesis of *E. coli* (Berg 2004). When there are multiple perceptual signals and multiple possible actions, computation of the most appropriate response rapidly increases in complexity: a problem that could be alleviated using a global modulation signal that integrates the deviations on all the channels (Fig. 1.d). Organisms with small distributed neural networks may implement this sort of control architecture².

² *C. elegans* provides a clear example, where modulation and integration were found through molecular-level studies of individual neurons associated with specific behaviours such as chemokinesis, repulsion and aggregation. Cheung et al. (2005) showed the modulation of roaming behaviour by the aerokinetic (oxygen seeking) motive in *C. elegans*. A suite of similar cross-modulation systems and their integration was reviewed by Bargmann (2012), covering *C. elegans* and *Drosophila* neural circuits. In both cases, multiple behavioural motivation systems were found to be extensively cross-modulated by neurohormone control systems. At a higher level of behavioural integration, the ‘threat-reward’ decision system of *C. elegans* was found by Liu et al. (2020) to be cross-modulated by GABA secretion in reward biased motor neurons, with reception in cholinergic pre-motor neurons that control avoidance behaviour. The effect was that the D-AVA circuit integrates simultaneous attracting and repelling stimuli to produce an outcome that is “*dynamically regulated by the motor system*”. This finding corroborates the theory presented by Kaplan et al. (2018), in reviewing the evidence for inter-neuron integration and modulation of behaviour control (action selection) in *C. elegans*. Rather than segregated feed-forward sensory-to-motor control systems, they suggested that distributed integration of sensory and motor signals, in conjunction with neurohormones, performed computations to generate the observed behaviour (analogous to the computations of an artificial neural network). Further support for this comes from the entirely different approach of dynamic modelling of the complete neural network of *C. elegans* by Antonopoulos et al. (2016), where the information-theoretic measure ϕ , from Integrated Information Theory (Tononi 2008), revealed significant computation creating new information within the network.

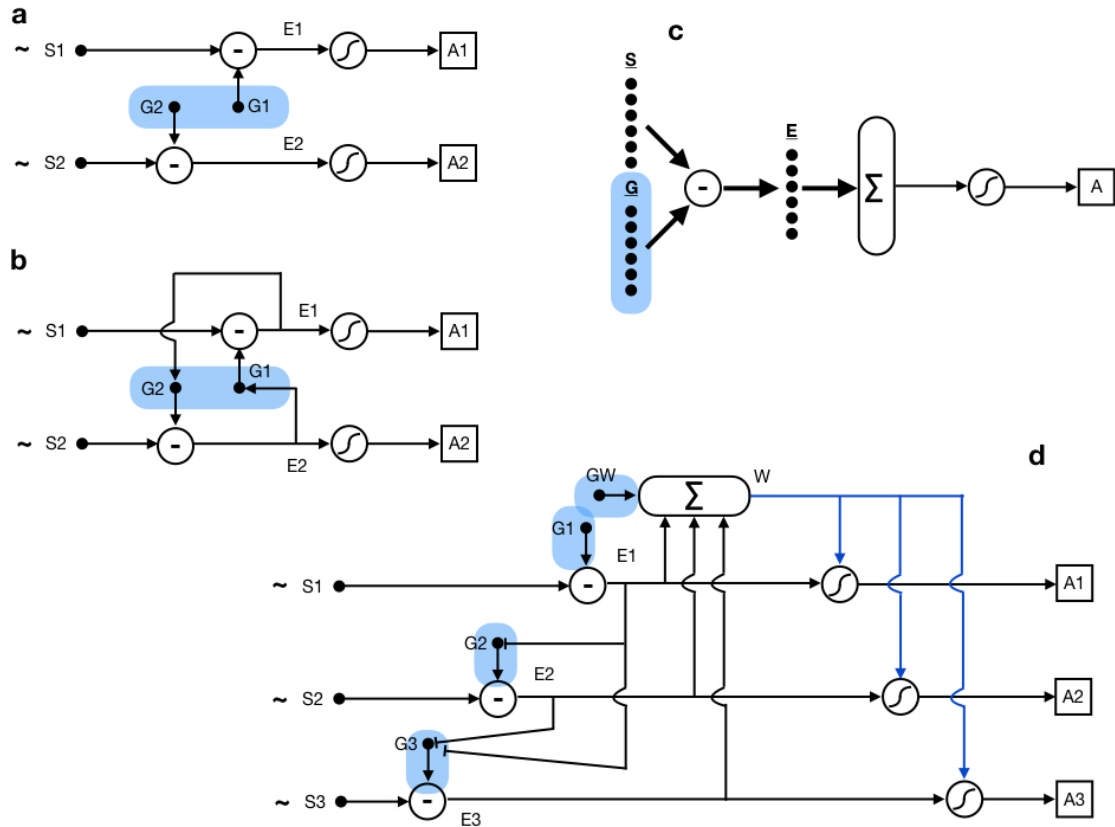


Figure 1: Reactive action-selection systems (autonomous control parts shaded). **a** is the simplest kind with two sensor – actuator channels acting independently. The actuators (A_1 and A_2) are triggered by exceeding a threshold in error signal (E_1 and E_2) which is the difference between the input signals (S_1 and S_2) and the set points (goal G_1 and G_2). **b** adds cross modulation for resolving conflict between A_1 and A_2 (e.g. E_1 could inhibit A_2 by increasing G_2). **c** shows multiple sensors and their associated set points combined by summation into a general action (arousal) signal for a single action A (e.g. escape); this could also be implemented through a ‘winner takes all’ algorithm instead of the summation. In **d**, three sensors add complication, especially in conflict resolution among actions (A_1 , A_2 and A_3). In principle, a complicated algorithm could embody a solution to all possible states for this system. In practice it is simpler to solve it by modulating the response thresholds of action signals with a general (arousal) signal W generated by summing the errors (E_1 , E_2 and E_3) along with an overall set point for arousal GW . This solution is generalisable to any number of sensors and actuators. In this example, E_1 has an inhibitory effect (reduces) G_2 and G_3 and E_2 reduces G_3 , but several other cross-modulations are possible here. (Blue shading for internally generated (free) signals).

Optimisation of a single (global) currency does not require a set-point since the maximum or minimum are self-evident extrema. Thus in principle, action selection does not need independently stored information (as the set point), but solving the optimisation problem simply by reaction (i.e. in the absence of anticipation as illustrated in Fig. 1.a-d)) would entail repeated cycling through the behavioural options to measure the realised value of the objective function in search of its optimum. That would likely be very inefficient and perhaps also risky. The alternative is to *anticipate* the objective function value for each candidate behaviour and select based on these predictions. This enables the action-selection system to be more general as it can solve the optimality problem in any conditions for which the outcomes of each behaviour can be estimated. There is now convincing evidence that this sort of anticipatory action selection is available to *Drosophila* flies (Barajas-Azpeleta et al. 2021, Cheriyaunkunel et al. 2021, Jiang and Pan 2022), as well as cephalopods (Ponte et al. 2022) and vertebrates such as corvids (Clayton et al. 2003).

2.2 Anticipatory action selection

Anticipatory action selection occurs when future states enter the decision-making. Since it uses unrealised future states in its determination, it necessarily implies proactive autonomy.

In allostatic systems (Sterling 2012), anticipation is built into the control system by an algorithm with pre-programmed information, for example the anticipation of diurnal temperature variation in physiological control (Pezzulo et al. 2022). The algorithm in these cases creates a link between stimulus and response that may be mediated by internal signal processing (e.g. by servomechanisms and internal oscillators (Cheng 2022)), but is causally necessary, i.e. part of a continuous uninterrupted chain. By contrast, an agent capable of *proactive autonomy* responds to a stimulus with an action *chosen* through evaluating the predicted outcome for each available option, using an internally

generated goal as a guide (Hoffmann 2003). This breaks the causal chain, introducing branching and optional causal paths (Ellis and Kopel 2019). The key difference between causally necessary linkage and proactive autonomy is captured by the idea that the former could be analysed using the engineering ‘black box’ approach to characterising systems by their input-output relations, while the latter produces outputs that cannot be understood from a knowledge of the inputs alone.

Predictions could, in principle, be provided by matching to memories of possible outcomes for every anticipated situation (a sort of database), but that would likely be cumbersome and inflexible. A strong competitive advantage can be gained from the ability to predict a possible future and select the action that maximises an objective function in novel circumstances, especially in an information-rich environment (Butz and Hoffmann 2002). The information system that fulfils the purpose, even for previously unanticipated circumstances, is a model of the self within the environment. It is a transformation (in the mathematical sense) between an input set of stimulus signals and an output that represents the desirability of an outcome (hedonic valuation). The transformation depends on both the action under evaluation and the state of the agent following the action, given the sensory inputs. Feed-forward models (systems that predict afferent signals, given the current efferent signals) are typically used to perform the transformation in anticipatory control systems (Fig. 2.A). Artificial neural networks are often used for the computation in engineering (e.g. Matsumoto and Tani 2020) and neural networks are known to implement it for the motor control of organisms (e.g. Jékely et al. 2021). This is extended to action selection by implementing a forward model for each potential action, predicting its outcome prior to realisation. Outcomes are generalised by a hedonic signal to be optimised for action selection. This signal may in practice be a neurohormone encoding valence information, which can then be used to select actions, e.g. by controlling the thresholds for actions to be realised Fig. 2.B). Since the information for appraisal results from training (by reinforcement learning) of

the forward model, it is internal and inherent to the control system (i.e. the organism) and to that extent free from exogenous control. It could function as a distress signal, but does not fulfil all the requirements for pain itself.

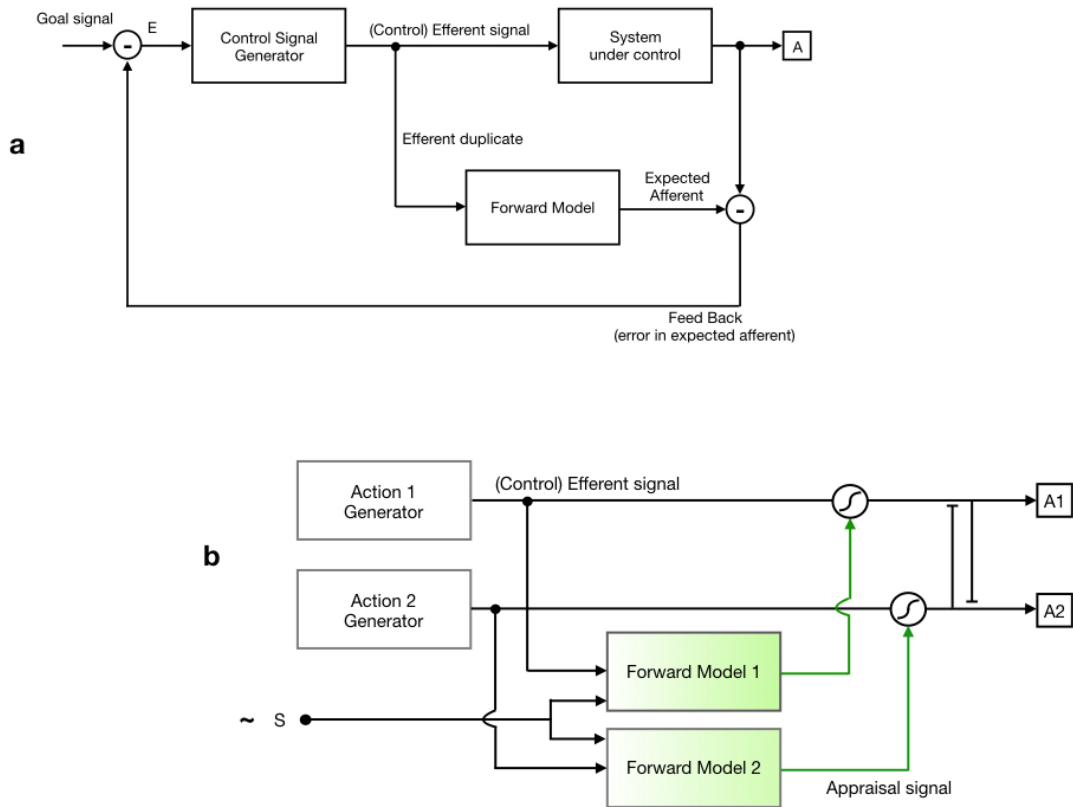


Figure 2: Control systems using forward models to predict the afferent result of actions. In **a**, the forward model continuously predicts the effect of the current control signals to enable feedback through which they are refined. This is a standard method for refining motor control. In **b**, this is adapted to predict a normative (hedonic) summary of the effect of each potential action (A1, A2) for use in action selection. Each forward model uses the corresponding efferent signal together with environmental perception, (S) for context, to generate an *appraisal signal* which may be implemented as a neuro-hormone level (indicated with green shading). This signal modulates the thresholds for enacting A1 and A2 (and the threshold gated action signals mutually inhibit to prevent indecision). The potential actions are realised as control signals generated by internal pre-programmed routines, but the forward models are trained by conditioning (reinforcement learning) to produce appropriate hedonic appraisal signals which therefore are internal (free) signals.

Forward models that can learn to generate a hedonic signal from potential actions, given a perceived context, can be implemented by recurrent neural networks with

hormone-secreting output neurons. In the active-inference approach, the idea of selecting an optimal action is replaced by finding optimal inferences (Bayesian beliefs) about likely future behaviours and their consequences (Friston et al. 2013). That uses an internal generative model, the states of which become probabilistic representations of external states (the physical world including the self). A hidden Markov model is a natural fit for it, readily implemented by a neural network. Active inference incorporates the goal as a minimisation of the divergence between the probability distribution of attainable states and states believed to confer high utility (Friston et al. 2013). It accounts for motivation (Clark 2020, Tate 2021), though not initiative (Klein 2018). *Initiative* is the ability to change or initiate a behaviour independent of external stimulus or ‘pre-programmed control’. It is the main emergent property of proactive autonomy having been derived from an internal model. Such proactive autonomy is the *anticipatory behavioural autonomy* (ABA) defined earlier. We use that term to emphasise the autonomy of decision-making and rational intention of the initiative — a point made by Hoffmann (2003), who termed the phenomenon “anticipatory behavioral control”. Proactive autonomy, based on evaluation of outcomes predicted by an internal model, enables *behavioural autonomy* (Schneider 2018). If the evaluation uses a common currency to represent the desirability of outcomes (e.g. a valence-informing hormone signal), then an arbitrary set of behaviours and outcomes can be compared in that common currency. Hormones can be accessible to the whole organism’s behavioural control system, with their concentration serving as a common currency. Then outcomes from different behaviours can be represented for evaluation in the common currency to find the solution to trade-offs such as between feeding and threat or reproductive opportunities. Action selection is then based on which prospective action produces the highest (or lowest) hormone level. Dopamine (generally involved in reward), serotonin (mediating anxiety) and cortisol or hyperglycemic hormone (arousal) are plausible candidates for this. ABA, then, is behavioural autonomy in which the choice is based on the organism’s prediction

of a global hedonic value under each of the available options. Because the decision is based on the anticipated value, rather than following prescribed rules (an algorithm), the organism's response is not entirely predictable from knowledge of the stimulus alone. Low predictability of behaviour, especially in novel circumstances, could therefore be an empirical indicator of ABA and by consequence, of the usefulness of pain.

2.3 A hypothetical model implementation

The “free-will machine” from Farnsworth (2017), taken as a hypothetical ABA generating system, can be implemented by a neural-hormonal control system that is consistent with the concept of pain. In Fig. 3, **S** represents perception inputs (signals from transducers, including nociceptors). They are compared to a model of expected inputs (**M**) by the comparator (-). This model is updated by e.g. Bayesian inference, and the modelled signals are compared to internally set goals (**G**) for the signals (desired or expected states). The difference between **M** and **G** on each channel (**E**) informs self-modelling about the current state. The result is the formation of a self-model that emerges in a particular Q-state. This Q-state may be interpreted as the informational embodiment of a quale. The model can generate as many different qualia as it has states: a number that increases rapidly with the number of neurons instantiating the model. Note the internal model does not have outputs *per se*, just its Q-states. The self-model has access to memories of Q-states, which it seeks to match. It is also connected with the rest of the body (soma), crucially including hormonal releasers and receptors. The integration of the self-model with the somatic system raises the Q-state to a state of affect: an emotional feeling (short term) or a mood (long term). This emotionally charged state of the combined model (neurons) and hormone system then modulates the drive to perform a finite set of actions (just two illustrated: A1,A2), each generated from a pre-programmed routine (R1, R2). The neurons that produce the routines are connected with the self-model such that the self-model modulates their thresholds for action. For

example, a particular Q-state may down-regulate the threshold for A1 and up-regulate the threshold for A2, with the result that A1 is performed. Note that attention (saliency) emerges from the somatic-self-model system as the hormones create the strength of the feeling of being in Q (that feeling being the quale). Thus, for example, if **S** is carrying substantial nociception, **E** will be large and the self-model will emerge in a pain Q-state, which will strongly stimulate hormones that thereby would be associated with pain and a state of suffering (emotional pain) will ensue, which will strongly down-regulate the thresholds for escape, guarding and other pain-related behaviours (turning them on), while simultaneously up-regulating the thresholds for all other behaviours, effectively stopping them.

This may seem complicated, but it is not unduly demanding of number and inter-connectivity of neurons. Greve et al. (2016) showed an artificial neural Turing machine could learn to solve a double T maze using just 70 nodes (artificial neurons), providing more than 10^{21} possible states. By contrast, drosophila has $\sim 100k$ neurons (Scheffer and Meinertzhagen 2019), each typically with ~ 100 synapses (estimate total of 2×10^7 synapses (Scheffer et al. 2020)); so even if only 1% of neurons implemented Q-states, there could be 10^{300} of them. The figures are beyond ‘astronomical’ for vertebrates.

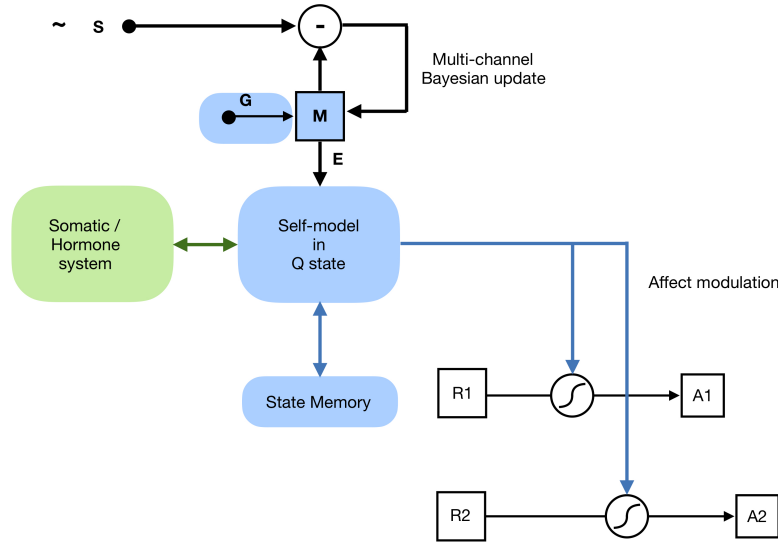


Figure 3: A hypothetical affect-driven action-selection system giving anticipatory behavioural autonomy. Bold symbols and lines represent vector (multi-channel) signals. **S** represents perception inputs (signals from transducers, including nociceptors). They are compared to a model of expected inputs (**M**) by the comparator (-). This model is updated by Bayesian inference, and the modelled signals are compared to internally set goals **G** for the signals (desired or expected states). The difference between **M** and **G** on each channel (**E**) informs self-modelling about the current state: the self-model emerges in a particular Q-state. It has access to memories (either experienced or pre-programmed) of Q-states, which it seeks to match. It is also multiply connected with somatic hormonal releasers and receptors. Integration of the self-model with the somatic system raises the Q-state to a state of affect resulting in an emotionally charged signal which modulates the drive to perform a finite set of actions (A1, A2), each generated from a pre-programmed routine (R1, R2). Modulation is achieved via action threshold modification. Further details in the text. (Blue shading for internally generated (free) signals on neurons, green for hormonal signals).

2.4 Anticipation, alone, does not require pain

Predictive processing is successful in explaining elementary cognition-response systems (e.g. Pezzulo et al. 2022). More generally, anticipatory action selection could, with relatively small systems, be implemented by a finite state automaton (FSA), leaving the organism absent of free choice and therefore not requiring pain. Examples of FSA-based anticipatory control systems typically depend on feed-forward models, as in Key et al. (2021), or internal models implementing active inference, as in Matsumoto and Tani (2020). Here we see the need for at least a memory, which may be elaborated into a model of the self, but having action selection still produced through the FSA architecture. Even if a global modulating signal (e.g. a neurochemical mediated state variable) were introduced to add nuance to the action selection, it could be implemented without recourse to agent freedom, still in principle leaving pain unnecessary.

This is roughly the conception of Key and Brown (2018), who developed a hierarchical predictive system in their search for the minimum system necessary to generate subjective experience. It consists of a nested pair of feed-forward predictive models, the inner model predicting the response to stimulus, the outer predicting the difference between this prediction and the realised response, given both signals together with ‘global input’ from other ‘brain’ areas. Key et al. (2021) argue that animals lacking a recognisably equivalent neural processing system would be incapable of the subjective experience that is pain. Since Key et al.’s (2021) two-level feed-forward model is only one of several plausible systems, that is a strong claim. One primary requirement, they claim, is that the higher-level prediction (or its error signal) is shared with the global system, for they say that the ‘3rd order awareness’ generated by their system is only sufficient for ‘pre-conscious’ awareness and it is the global availability of its output that produces conscious awareness, though they do not explain how or why. The whole system they propose remains reactive, since it does not incorporate any goal or desire and also has no action selection component (it was not intended for that purpose). We therefore need to add

goal-seeking to obtain a model of experience-driven autonomy. Key et al. (2021) distinguish their model from other predictive processing schemes, principally on the grounds that their predictive models are not “*embedded within* the internal sensory processing stream”, claiming that subjective experience cannot be supported without that separation of computational tasks. But computationally, it makes no difference whether the nested predictors are depicted as within the stimulus-reaction processing system or as a separate module sharing signals with it. What really makes the difference is the sharing of prediction signals with the global system. Though Key et al. (2021) recognise that necessity, they say nothing about what the global system has to do with these shared signals in order to generate subjective experience. Key et al. (2021) appear to imply that subjective experience is just ‘what it is like’ to have global availability of certain signals, or more generally what it is like to be in some particular states. We strongly agree to that: no mysterious, even metaphysical, experience-generating processor needs to be added to a brain in order to generate subjective experience; it is simply what it is like to be in a particular state (what we termed the Q-state). This is a philosophical position that avoids the homunculus fallacy.

In summary, subjective experience has an evaluative component — a normative character — arising from the difference between the current state and that sought by a goal-directed action selection system. It is therefore part of an anticipatory autonomy system. Pain feels bad because it is a state that is far from that desired and it motivates action in response. What it feels like to be in pain is the awareness of the gulf between a current state and the comfortable (homeostatic) state constantly sought. Thus, anticipation is necessary for pain, but only jointly with goals and a global-level evaluation.

2.5 Autonomous evaluation necessitates pain

The key difference between an automaton system and a free autonomous agent is that in the latter, actions are selected based on their evaluation in a common currency. It is the independent evaluation, a computational process isolated from the link between perception and response, that provides the freedom of free autonomy. Evaluation is subjective and context-dependent; it cannot be replaced with a FSA algorithm, not only because it entails an indeterminate number of states, but because it is necessarily a faculty of the whole organism³, which is the only organisational level to which we may accord the status of freedom. These choices are not determined by exogenous causes, nor by immutable internal causal structure (an inbuilt algorithm) in any component part of the organism; instead they are determined by the goal-seeking intention of the unified whole of the organism. This optimisation can be termed the 'will' of the organism only because the goal is instantiated at the highest level of causal organisation (Farnsworth 2018; 2017). The goal is the maximum of a global utility function which, by natural selection, should normally coincide with Darwinian fitness, but for the individual organism it may be represented by an effect-like signal on the pleasure/ pain axis as described by Hoffmann (2003) and (Schneider 2018) (noting this may be a simplification since pain and pleasure are thought to be separate systems (Pietri et al. 2013)).

Evaluation requires a universal currency to compare the value of each option regardless of its nature, similar to the economists' notion of 'utility', which enables comparison of cinema tickets with cheese. This universal currency needs the properties of valence (good/badness) and intensity (activation or arousal level). These are provided by the 'emotional space' defined by Russell (1978) and elaborated in Russell and Barrett (1999). This idea of evaluation on valence and arousal axes of a universal currency is compatible with the cognitive appraisal theories reviewed in Scherer et al. (2001). More recently, the

³ defined by closure to efficient causation, so not including symbiotic or parasitic organisms as in some interpretations of 'holobiont' organism.

evaluation component of emotion has gained broad acceptance within (human) emotion theory, brought together under an inclusive definition by Scherer (2022):

... “emotions 1) consist of an episodic process in response to a perceived event or situation of major significance, 2) which is characterised by recursive causal effects (forward and backwards) between several components that include the evaluation of the event in terms of its significance for the goals and values of the individual, 3) creating physiological reactions, motor expressions, and action tendencies and 4) that this process is partially accessible to consciousness, resulting in feelings that 5) can be categorised and subsequently labelled by the individual in terms of its subjective conceptual structure”.

The two-dimensional circumplex model of affect (Russell 1978, Posner et al. 2005) is the antithesis of the so-called ‘basic emotion’ model in which emotions are discrete separate sensations. The circumplex model has gained considerable empirical support and the idea that a wide range of emotions can be constructed from just two axes of latent variation (valence and arousal) is commonly invoked in human psychology, though contested (Ortony 2022). The practical (fitness) value of simple emotions has been shown using reinforcement learning in artificial intelligence systems, which can be enhanced by incorporating simulated emotions into action selection (Sequeira et al. 2015). This integrates current perception signals with memories and model expectations to produce an autonomous self-centred decision-making process. Affective signals are used as an overall hedonic objective function to be maximised, using both current and anticipated states for possible actions in the context of action selection. A typical arrangement involves a joy vs. distress axis, which may be further enhanced with a hope vs. fear axis, identified as the anticipation of joy or distress, respectively (Broekens et al. 2015).

At least for the present purpose, the single dimension of a valence is very suitable for the simplest motivational signal: the contrast between pleasure and pain, with their

associated general response of approach and withdrawal. In short, a single general signal of valence is enough to provide for the affect aspect of pain (and pleasure) and to function as a universal comparator of actions available for selection suggested by Hoffmann (2003). Further, we cannot ignore the obvious parallel between a universal signal of valence and the widely acting neurohormones, which are taken by many as an objective indicator of psychological stress or arousal, and in animal studies as a surrogate for pain: cortisol for vertebrates (Cerqueira et al. 2021, Stafford and Mellor 2005, Wagner 2010) and hyperglycaemic hormone for invertebrates, such as crustaceans (Chang 2005, Elwood and Adams 2015). Autonomous evaluation can be implemented using such hormones as a universal and integrating motivational quantity that is continuously variable and gives effect to the state of the internal model: in particular the feeling of pain. Thus, our key proposal is that pain is only adaptive for those animals able to make autonomous anticipatory decisions, i.e. animals that show ABA.

3 Empirical support for ABA implying the need for pain

The function of pain, distinct from nociception, is identified by Sneddon (2009) as enabling an organism to “quickly learn to avoid the noxious stimulus and demonstrate sustained changes in behaviour that have a protective function to reduce further injury and pain, prevent the injury from recurring, and promote healing and recovery”. In other words, pain should elicit persistent changes of behaviour through modulation of action selection. For example, we see conditioned place avoidance for areas associated with noxious stimuli in shore crabs (Magee and Elwood 2013) and octopuses (Crook 2021). Further, octopuses that could not avoid noxious stimuli preferred areas associated with a local anaesthetic. Other long-term changes in behaviour observed after noxious stimuli include alterations of shell preference in hermit crabs, which last at least 24 hours following electric shock (Appel and Elwood 2009, Elwood and Appel 2009) and the onset

of anxiety-like states in crayfish after shock (Fossat et al. 2014), which are also seen in fish (de Abreu et al. 2020) and amphibians (Brown et al. 2013). Anxiety-like states are usually associated with serotonin (Best et al. 2020, Curran and Chalasani 2012), but do not alone imply pain; e.g. a simple algorithmic mechanism for their manifestation, requiring only two neurons, has been found in *C. elegans* (Eliezer et al. 2019), but without evidence of evaluation. Anxiety-like states do demonstrate anticipation, and generalised modulation of action selection, but could, in principle, be generated by an automaton. So though the behavioural observations above are consistent with expectations of pain in a wide range of species (Sneddon et al. 2014), they do not conclusively support our hypothesis concerning autonomous and anticipatory behaviour. That hypothesis broadly suggests that pain is only useful, and hence likely to be present, in animals that can make a free choice between available responses in the presence of a noxious stimulus. To test this, we would need evidence of 1) mental models of the self and the environment to support anticipation; 2) flexibility in behavioural responses to stimuli (showing that options are available); 3) proactive choice and forward planning (actions based on anticipated consequences, rather than just the current state) and 4) free choice of response to noxious stimuli (not algorithmically pre-programmed) that is rational rather than random (shown by e.g. state-dependent trade-offs).

3.1 Models of self and the environment

Models of self presumably developed early in evolution, with examples emerging in a wide range of multicellular animals (Jékely et al. 2021). The most basic of these models involve reafference (von Holst and Mittelstaedt 1950), which is the term given to the ability of an animal to discriminate between sensory changes due to self-movement and those due to environment change (Jékely et al. 2021). This ability is important because the two are likely to have very different meanings for the animal. For example, an object taking up more space on the retina (or compound eye), i.e., a looming stimulus (Temizer

et al. 2015), could be due to the animal approaching the object or the object approaching the animal. In the first case there is little risk whereas in the latter case, looming might indicate danger to the animal. But reafference also applies to a wide range of stimuli, such as flow of water over the body surface due to own movement contrasted with that due to environmental flow, or to deformation of the body due to own movement or to some external force. That is, the animal has a model of self, and largely disregards inputs due to self-movement, whereas those due to external changes receive attention.

Animals also form models about the environment, demonstrated by the classic experiment in which chicks anticipated the timing of a light being switched on and off at regular intervals, showing startle responses when it turned off early or late (Broom 1968). Anticipatory modelling is of course central to predictive processing and active inference theories and the generation of associative learning. When animals learn about associations between two environmental changes, as in classical conditioning, or between an action and subsequent environmental event, as in instrumental conditioning, they form mental models that allow distinction between chance coincidence and true causal relations between neutral events and subsequent events of biological significance (Dickinson 1980), or at least allow for predictions Gallistel and Gibbon (2000). This process can be complex, even in invertebrates such as insects (reviewed by Perry and Barron 2013). Numerous groups of cells and circuits, and their interactions, have been identified as involved in learning about rewards and punishments. Further, the roles of octopamine and dopamine in reward and punishment learning have been discovered, as have more complex interplay between these two control circuits. Of particular interest, however, is the suggestion that with rewards there are subjective feelings of “liking” as separate from “wanting”, and again separate but interacting circuits seem to be involved (Perry and Barron 2013, Berridge and Robinson 2016). That is, the hedonic value of an unconditioned stimulus plays a role in the learning about a conditioned stimulus, such as an odour, and how it predicts the arrival of the unconditioned stimulus, such as sucrose or

a sweet substitute. Hedonic value may also play a part in avoidance of punishment, such as electric shock. For example, hermit crabs that receive a shock within their shell seem to value that shell less than do crabs that did not receive a shock, even though the shells remained the same (Appel and Elwood 2009, Elwood and Appel 2009). We conclude from these studies, and many others, that some animals are able to gather information about themselves and the environment to predict future events. This enables animals to better gain rewards and avoid risks. This integration of self and environment is key to the success of metazoans.

3.2 Flexibility of responses to stimuli

Identifying flexibility of response (following the classical definition of free will: “able to do otherwise”) depends on there being available response options and a demonstration of more than one response to equivalent stimuli from the same individual. The first criterion can be established within a species by observing inter-individual differences in response, because such differences would result from differences of internal state among individuals, i.e. a complex of genetic, developmental and accumulated experiences (Stamps 2016). Appel and Elwood (2009) demonstrated this with hermit crabs undergoing a standardised noxious stimulus (i.e. with minimum variation in magnitude and site of application). Crabs were induced to occupy empty gastropod shells wired to apply electric shocks to the abdomen of the crab within its shell. Of the 123 crabs that received a standardised shock treatment, 61 evacuated the shell and 29 of those groomed and tended to their abdomen at the site of the shock application. Four crabs attempted to climb the wall of the observation chamber and three engaged in shell-rapping, an activity normally seen in fights for ownership of shells. After evacuation, 57 crabs re-entered the shell, leaving four that stayed away from it. None of these activities were observed in unshocked controls. Evidently the observed behaviours show a variety of individual responses to the same noxious stimulus, demonstrating options for action

selection.

To identify flexibility within the individual (endogenous placticity), we must first distinguish between sources of variation in their response. We reject random processes as they negate autonomy. Developmental shifts (maturation and e.g. role differentiation in social insects) do not indicate coincident options for the animal (discussed by Jeanson 2019). Changes in response to a change of the environment might be generated by an automaton algorithm, so not free. For example, (Czaczkes et al. 2018) observed task switching between exploration and exploitation in forager ants in a T-maze with sucrose rewards at the end of each arm. Following a period of training in which ants learned to associate reward levels with various cues, ants were free to choose either arm over multiple trials. They showed little switching between arms, irrespective of their reward levels, as long as rewards remained constant. When reward levels, along with associated cues, in both arms were simultaneously increased, or decreased, then switching rate also increased between trials, showing a change from exploitation to exploration behaviour. If ants have an exploration algorithm, but otherwise default to exploitation, then a simple threshold switch, sensitive to reward change (Wilson 1976), would suffice to produce this apparent behavioural flexibility. Conversely, within-individual changes of behaviour, without a change in environmental stimulus, may result from learning, in which case internalised information gained by the organism, not inbuilt, is the source of change; hence free-choice flexibility is demonstrated. Jeanson (2019) discussed learning-dependent flexibility in social insects, e.g. that in ants able to perform multiple tasks, a successful foraging experience can increase the likelihood of repeated foraging (Ravary et al. 2007). Representing cephalopods, Chung et al. (2022) showed that cuttlefish changed their response to ambiguous prey choice following the experience of receiving an unexpected food reward. They interpreted this change as foraging strategy selection mediated by an internal state they identified as an emotion-like state. Magurran (1993) reviewed a substantial body of evidence of context-dependent behaviours within

teleosts, not least the ability of male guppies to choose between overt display for a mate, or “sneaky” mating tactics. Most supportive of ABA, Earley et al. (2013) found that mangrove killifish (*Kryptolebias marmoratus*) ‘perceive’ their own fighting ability (implying a self-model) and they “adjust contest strategy” when that perception is updated following wins or losses. The authors identified this behavioural flexibility with changes in three hormones, concluding it is “modulated by internal state”.

3.3 Proactive choice and forward planning

Examples of forward planning in action selection are available among invertebrates (Elwood 2022). One such study used terrestrial hermit crabs and allowed them to walk along a corridor with obstacles that partially blocked the passage of the crab’s shell (Sonoda et al. 2012). The obstacles varied in the degree to which they made passage difficult. However, crabs were proactive in this task, turning their shells to avoid the obstacle before encountering it. Further, they turned the shell to a greater degree if the space between the shell and obstacle was narrow. When crabs had plastic plates attached to the shells making the obstacle course more difficult, they turned the shell to a greater degree on their first attempt, enabling passage without the plastic extension colliding with obstacles. The behaviour (degree of turning) was selected prior to collision experience and in response to a novel situation, thus showing proactive selection of behaviour for optimal outcome. Other examples with hermit crabs showing proactive flexible responses are discussed in Elwood (2022). Ants following cues that predict a valuable reward show more pheromone marking of the trail than do those following cues to a weaker reward, which again suggests forward planning (Czaczkes et al. 2018). Spiders that live in a complex 3-dimensional environment can detect prey from a distance but reaching the prey might not be achieved in a straight line. Spiders have been seen to plan the route and on occasions might move further away from the prey in order to get to a branch that will then lead them closer (Tarsitano 2006). All together, action choice

with forward planning has been documented for arthropods. Among vertebrates, these faculties are well known, for example through reversal learning experiments, especially with birds (e.g. Bond et al. 2007).

3.4 Non-algorithmic, selection of response to a noxious stimulus

Probably the best evidence for proactive choice in response to a noxious stimulus comes from examples of trade-offs between avoidance of a noxious stimulus and any other goal (e.g. Balasko and Cabanac 1998). Evidence of this in fish (and cephalopod) species has been reviewed (Sneddon 2019). Goldfish (*Carassius auratus*), trained to feed in one region of an experimental aquarium, and subsequently subjected to electric shocks, would spend more time in this feeding/shock zone the more food-deprived they were. This trade-off shifted away from feeding attempts towards escape as the shock intensity was increased (Millsopp and Laming 2008). Fitness benefits were shown for squid (*Doryteuthis pealeii*) as they put extra effort into escape from predator cues when they were experimentally injured, leading to an almost doubling of survival rate compared to those that had been anaesthetised during the injury process (Crook et al. 2014). Hermit crabs evacuate their shell after an electric shock with a probability that depends on the quality of the shell (Elwood and Appel 2009) and also the presence of a predator odour Magee and Elwood (2016). Thus, these crabs displayed a flexible trade-off when responding to a noxious stimulus with respect to keeping a high-quality shell and avoidance of predation. Further, a recent study on bumblebees demonstrated a trade-off between avoiding a high temperature and obtaining a high-quality food source, with the bees using learned colour cues for their decisions, indicating both flexible responses and associative learning based on contextual information (Gibbons et al. 2022b). These demonstrations of trade-offs suggest proactive choice following noxious stimuli. It is doubtful if they could result from an inbuilt algorithm because of the complexity required and because they seem to occur in novel situations. Further, some authors have put considerable weight on trade-

offs as a key criterion of pain (Crump et al. 2022). Intriguingly, some crayfish subject to a heat stimulus from a soldering iron briefly touching the animal grabbed the shaft of the soldering iron in response, whilst others withdrew defensively (Puri and Faulkes 2015). This unexpected protective response replaced the reflex withdrawal, seen in some individuals, with a co-ordinated attack that seems to use freedom of action-selection to manifest.

3.5 Evolution of pain experience

So far, there is evidence for pain in three major phyla, the chordates (Sneddon et al. 2003), molluscs (Cooke 2021) and arthropods (Elwood 2019). These three phyla arose during the Cambrian explosion and the most recent common ancestor for these is likely to be a free-living worm-like organism from about 530–550 million years ago (Elwood 2011). The parsimonious explanation for the evolution of pain in the three phyla is that there was one evolutionary step that occurred in or before the most recent common ancestor. Against this, evidence for pain is restricted to specific groups, such as the cephalopods within molluscs, the decapod crustaceans and some insects and arachnids within the arthropods and the vertebrates within the chordates. Evidence for pain among many phyla remains weak or absent, but that might simply reflect lack of relevant studies. However, a patchy distribution of pain might occur if pain was lost in some lineages. For example, taxa that evolved from a free-living form to a sedentary lifestyle may have reduced their behavioural choices and, thus, there may be no need for free choice and pain. For example, bivalve molluscs, such as oysters and mussels, which are fixed to hard substrates and therefore limited in how they might respond to noxious stimuli, might not benefit from a pain system.

We might reasonably expect pain to be found in basal groups of these three phyla but within the arthropods, identifying basal groups with extant examples has proved difficult (Edgecombe and Legg 2014). Because pain has been suggested for the decapods, we

should examine basal crustaceans, for which ostracods or branchiopods represent extant early taxa, but we are not aware of any studies that might indicate sentience in these groups. There is also a paucity of relevant studies on primitive insects (Gibbons et al. 2022a). Basal molluscs, such as the worm-like aplousobranchs, provide no evidence for sentience because these are deep-sea burrowing animals and we are not aware of suitable studies on live specimens (Wanninger and Wollesen 2019). There is more information on early chordates, for example the protochordates, including *Amphioxus* (Lacalli 2022). There has been detailed comparison of the CNS of *Amphioxus* with those of vertebrates. This indicates that the brain of *Amphioxus* has some of the areas found in vertebrates, but *Amphioxus* lacks the major areas involved in the sensory experience of vertebrates. Lacalli (2022) concludes that sentience developed within the vertebrates rather than being a feature of the protochordates.

In general, the evidence points to the less parsimonious multiple origins of sentience and pain, as opposed to a common ancestor, with subsequent losses in some groups. One reason for this is suggested by Lacalli (2022), specifically for the chordates, but which might apply to the molluscs and arthropods. Early groups in these taxa lack well developed sensory systems. For example, light-sensitive cells may be found in early forms, but they likely only provide information about light levels. Whilst they might provide warning due to the shadow of a predator, they do not provide an image that came with the evolution of eyes. Eyes have evolved in some groups of molluscs, vertebrates and arthropods, and although these differ in composition, they are able to form images of distant objects and thus gather vast amounts of information (Godfrey-Smith 2020). If that is processed efficiently, it may be used to predict what will happen next. For example, improved sensory ability provides information about potential mates, potential competitors, potential predators and a myriad of other environmental changes that might impact fitness. This improved sensing is not restricted to vision but involves other modalities for which there has been marked development in the appropriate sense

organs. The integrated processing of this much larger amount of information has necessitated a parallel development of nervous systems. This was particularly likely in those animals that developed a highly mobile predatory lifestyle and the requirement for swift decision-making. Thus development of special senses leading to a substantial enlargement of information and potential action space may have stimulated the parallel development of ‘unlimited associative learning’, which was identified as the transition marker for the evolution of consciousness (Ginsburg and Jablonka 2019). and sentience. In engineering terms, the state-space of sensory information and potential actions completely outstripped the capabilities of automaton-based systems, necessitating autonomous affect-driven decision-making. We suggest that the resulting flexibility of behaviour and the vastly improved ability to predict has brought about the requirement for pain in the context of affect-driven decision-making. That is, pain may be a consequence of mobility and behavioural choice (anticipatory behavioural autonomy) that we see in fish (and other vertebrates), cephalopods and decapods and some insects and arachnids. Animals outside of these specific groups, but with similar sensory and behavioural properties, may be considered likely to also experience pain. One suggestion for this is the crustacean group of stomatopods, commonly called mantis shrimps, which so far appear to have been excluded from a consideration of pain-like states. Based on our arguments, we might also expect to find evidence for pain in other arthropods such as spiders, scorpions, millipedes and centipedes, and we encourage studies of such animals.

4 Conclusions

We have considered pain as an experiential phenomenon emergent from the neural processing of nociceptive signals in the context of a self-model which is integrated with a neurohormonal system that provides emotional valence. Pain causes suffering because

the pained state is remote from the goal (pain-free) state. In this view, the biological systems needed to cause suffering are no more than those needed to cause pain, so where pain is established, suffering is likely too. We consider the term psychological stress to be equivalent to suffering and note that psychological stress is the primary measure for animal welfare studies and normally quantified by surrogate stress-hormone assays.

Our proposition can be put rather simply: pain is adaptive only for organisms capable of anticipatory behavioural autonomy (ABA), which is the freedom to choose among available behaviours based on model-derived anticipation of the outcomes, so pain could reasonably be attributed to any organism capable of that. The hypothetical system we propose for achieving ABA is just one of presumably many biologically plausible systems, but the components and architecture of their assembly into a working action-selection system are all testable. It is consistent with previous models of anticipatory behavioural control (Hoffmann 2003), the imperativist account of pain (Martinez 2015), the ‘organisational approach’ (Mossio et al. 2009) explanation of autonomy (Bich and Damiano 2012, Froese et al. 2007, Farnsworth 2018), proposed hallmarks of consciousness (Ginsburg and Jablonka 2019) and empirical findings in neuroanatomy (Barajas-Azpeleta et al. 2021, Jiang and Pan 2022) and ideas about animal behaviour (Budaev et al. 2020, Clayton et al. 2003, Crump et al. 2022, Elwood 2019, Ponte et al. 2022, Sneddon et al. 2014). One advantage of the concept we propose is that it succeeds in explaining apparent free choice as well as the role of emotional pain (suffering) in the control system of organisms possessing it. Another important advantage is that it is in principle testable using animal behaviour experiments.

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