

Running Head: Value or Worth? (Un-)Certainty, Sign- or
Goal-tracking, Appropriations in dyn4-TAM-Space

For Value or for Worth? Part 2: A Neuroeconomical Thought-Action-Mood-Space Modulated by (Un-)Certainty as Sign- or Goal-tracking

Gottfried Reinhold Sebastian Treviranus

BipoSuisse, Independent Research, Vereinsweg 11, CH-3012 Berne, Switzerland

*Corresponding Author: bipoSuisse@bluewin.ch

Copyright©2017 by authors, all rights reserved. Authors agree that this article remains permanently open access under the terms of the Creative Commons Attribution License 4.0 International License

Abstract Epistemology requires trans-disciplinary logics for convergence. Here a logico-geometrically expanded cyclical version of the classical French temperamental and anxio-affective thought-action-mood-model, “*dyn4-TAM-cube*”, harboring Appropriation Waves (AWs), encounters an *effort*-related node of present neuro-economical debates: the cyclical relation between “value” and “worth”. Accordingly, as a fundamental of the brain, this essay’s second part continues to explore the alternation between symbolic frontal 4-dimensional (Halford) processing (“*4D- Thought*”), and high-dimensional parietal (Rizzolatti) intel- lectual intuition (“*5D+-Action*”), as balanced according to Richard Sorrentino’s prime motivator trait (Un-)Certainty Orientation (“*UO-versus-CO*”) interacting with “Mood”. The two mentally processed transitions between these low- and high-dimensional domains, “*4D-T~*” and “*5D+-A~*”, are complexity-reductive Perception $P\{A \rightarrow T\}$ and -expan- sionist Intention ($I\{T \rightarrow A\}$), from which two kinds of lear- ning feed into “*4D-T~*” for decision. Specifically in early AW the not just predictive, but *incentive* among Pavlovian cues putatively *promises discounts in effort* which foster intentions for *worth-appropriative moves*: $I\{T \rightarrow A\}$. As “*UO-versus-CO*” to date seems homologous to the more sign- or more goal-tracking dimension (“*ST-versus-GT*”) in rats. Since ST is controlled by the thalamic paraventricular nuclei and GT e.g. by the hippocampus, putative intruding *mast cells* might cause the depressive reversals in orientation, establishing mast cells as bio-economical agents.

Keywords (Un-)Certainty Orientation, Sign-tracking, Goal-tracking, Computational Psychiatry, Relational Complexity, Neuro-economics, Prediction Error, Cortico-subcortico-thalamo-cortical Circuits, Mast Cells, Mixed Bipolar Disorder

1. Introduction

The second part of this essay [2] proposing a more integrated and developed classical framework related to temperamental and anxio-affective “appropriations” in recurrent sequential waves, and attracted by permuted states, “*dyn4-TAM-cube*”, (see abbreviation code¹), provides an encounter with currently rapidly unfolding convergent neuro-economics.

2. Interfacing 4DT~ and 5+DA~: “Value” and “Worth”

The *opposite* cognitive attitudes in facing unexplained complexity, which manifest as opposite human performances [1], in the *dyn4-TAM*-model [2] occur at the *interfaces* between low- and high-dimensional processing. The present proposal states, that *in analogy* to *uncertainty-oriented* humans (UOs), *sign-tracker* rats (STs), after engaging in the “perception of value” $P\{A \rightarrow T\}$, perform vigorously in exploring *anticipated potential discounts in effort*, and secondary advantages in exchange of *values*, and *goal-tracker*-rats (GTs) instead in the “intuitive intention of *worth-appropriative moves*” $I\{T \rightarrow A\}$. Where the former *sanguinics* excel in *reducing* complexity, and thereby hopefully *effort*, into more sophisticated models of appropriation, the latter *choleric*s excel in keeping or *expanding* complexity. In order to substantiate this claim this convergent review aggregates more detailed structural analogies as a first step to hypothesis-testing.

Such temperamental variety, as other intra-group diversities [3], may favor fitness of groups and societies’ welfare [4] and are at the core of conflict and cooperation.

Such value/worth-*dualism* also complies with the old stance, that the market *value* “exchanged (...) is the quantity

of labor (...) commonly taken in producing them.” [5]. Labor in the individual appears as “effort” - as reflected in e.g. tonic extracellular striatal dopamine [6]. Thus abstract value represents production costs, which ultimately consist of e.g. human effort.

This duality appears as biologically hard-wired. The phylogenetically newer anterior lateral OFC processes such abstract (monetary) “value”-rewards [7,8], which overcome subjective effort as incentive appetite probably signaling the opportunity of unusual bargain, while appropriated “worth” results in anticipatable, utility- reflecting affective “hot-spots” [9]. Hereby even the temperamental shapes of “utility” curves seem to be accounted for, yet “value” and “worth” are commonly considered as interchangeable [10].

3. Dual vs. Mixed Motivational Updates

Reward prediction error (RPE) signals, which update about the reliability (“precision”) of cues (for opportunities), were first discovered in midbrain dopamine (DA) neurons [8, 11-13], and then in additional sites including - for punishment - the habenulae [14], striatum – also in humans [15].

The scope-specific single-step updating via eligibility-targeted plasticity [11] accordingly is not a backward modulation, albeit still connected to specific needs. In fact pairing a CS(1) first with US(a) and then with US(b) is equally efficient for appropriate action, but blocks learning from a new CS(2) about US(a)¹. Rather than supporting an additional Konorskian general activation [16], such a block, deviating to another US(b), would allow to stick to an established joint cue about less effort in appropriating both.

A recent simulation [17] of sign-tracking (ST) underscores an antegrade capacity of the cue to reawaken a reward pertinent to a need, which then engages in a second, interrelating process. The niche-dependent relative efficacy of goal- or sign-tracking is not considered and these “rewards” are added and not multiplied.

Only the reminder that appropriations are motivated by two processes [18], which in dyn4 occur at “Intention” and “Perception” interfacing between 4DT~ and 5+DA~ (Figure 1), which therefore ought not to be collapsed, seems to clarify several neuro-economical issues.

4. Disincentivizing at the Interfaces

Besides cognitive efforts, and many other intricacies of motivation [19], motor efforts impact future “rational” best-reward-for-least-effort-choices and subjective evaluations (like regret) through a same network. This correlates with reward and inversely with effort and involves SMA, itself corresponding to avoidance, and dorsal ACC’s caudal portion for calculating motor costs. Not decision for action, but seeking of reward is activated by the vmPFCⁱ [20].

As dissociated appear the shape and site of the action-devigoring impacts a) of expenditure of effort, which is lesionable at the ACC (in the cortico-subcortico-thalamo-cortical circuit, CSTC, for 5+DA) or by unilateral sites of mPFC in rats [21], and b) of delayed reward, which is lesionable at OFC (in the CSTC of Mood) [22]. While ACC and the anterior insula (aINS) perceive, $P\{A \rightarrow T\}$, efforts [23] from 5+DA, delays might allow worries about extra efforts (4DT), to dis-incentivize, less $I\{T \rightarrow A\}$, at the ventral striatum and the vmPFC. At least for STs effort-related fatigue usefully prevents further losses by stopping exploration of tempting cues [24].

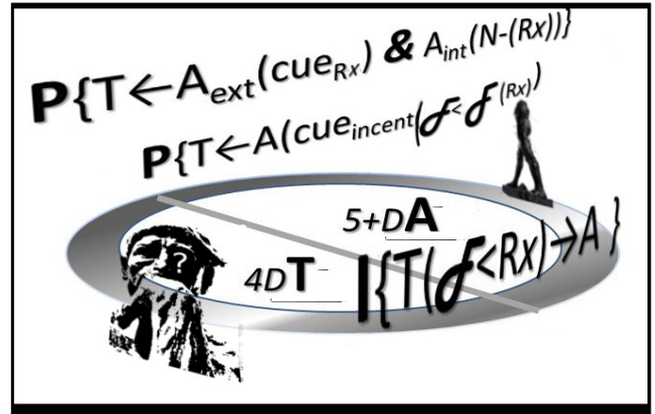


Figure 1. The cycle alternating 4D-Thought and 5+D-Action, as illustrated through the “Thinker” and “Marcher” by Auguste Rodin, hides Appropriate Waves (AWs). At its first transition of Perception of external and internal feedbacks from effected change, some signals become contingent reward-x-specific cues, $A(\text{cue}(R_x))$, inspiring the early AW. Some of these become incentive cues of a different kind, which hint to a learned discount in the expected appropriate effort (F). Intention urges for action when the expected cost, felt as effort (F), is expected to be more than matched by the perceived reduction of Need by reward R_x as $5+D-A_{int}(N-R_x)$.

Less effort as a negative reinforcement might provide a further “facilitation of a specific form of neural computation that results in conditioned approach behavior” also by mesolimbic dopamine, which here would not act as an anticipator of reward [25].

The principle of least effort also guides many decisions when building Intentions ($I(T \rightarrow A)$) to act. As a loss-minimizing strategy – under interacting conditions of exhaustion and scarcity [26] – it is computed at the frontal pole as effort under risk [27]. In dyn4 cues instill “Hope” to overcome the effort (transition “e” in part 1, Figure 3). Maximal own expendable specific need-proportionate effort (“Effme”) therefore remains a crucial ceiling parameter, and the inverse of “value to me”.

While decisions localize to the OFC, foragers rapidly adapt their means or goals to changing surrounds through such a function of possible maximal effort and attainment through least effort, i.e. of “resources and opportunities.” Herein the dorsal ACC [28] and the posterior cingulate cortex may at times signal the “pressure to pursue unlikely choices” – preferred by uncertainty-orientated STs [29].

5. Prediction of less Effort or of Reward

The ubiquitous “reward prediction error” (RPE) in most current accounts (but not in all experiments) could probably be smoothly replaced by a RPE *normalized* into $RPE(-\Delta\%Effme)^i$ through multiplication with a ratio expressing the reward-specific *maximal expendable effort*, integrating past specific expenditures and regrets, and also the present “energetic state” [26], divided by the *presently anticipated*, at times negative, *discount*, as indicated by incentive cues.

The perception of the *cue* is temporally *closer to the evaluation of anticipated effort*, and therefore more easily associated with it, than the outcome in terms of *utility* („worth”), apparent only after consumption. This *worth* divided by *recent effort* answers to the question “Was it *worth the effort?*” - risking *regret* [31] and updates the net RPE [32].

5.1. Common and Individual Economic Comparison

The need for comparison of rewards, often *without* a “common quality”, calls for a ranking on a “common scale of values”, on which the encoding of the RPE as “subjective value” is proposed as a, questionably, “ideal way” of steering economic decisions [31]. This “value” in *reality* though, as we can see, is a variable *composite* of *worth* (utility) and, at times incentive, *value* (effort), which is hidden behind of- ten dichotomic decisions. The variation of rewards in only one attribute is said *not* to allow the “isolation” of subjective preference - as far as sooner, more certain, and more should be preferred as better [31]. But beyond this we want what we like at *discounted* objective and subjective *effort*, thus the RPE changes to the residual $RPE(-\Delta\%Effme)$.

The closing e.g. consummatory end of the **AW** perceives the feedback $P\{A \rightarrow T\}$ from $5+DA$ -led actions through evoked external and internal reactions, and activates *instrumental* learning. The update of $RPE(-\Delta\%Effme)$ in $4DT$ ~ instead learns from spared, or increased, momentarily anticipated *effort*, not from outcome [8,11].

Accordingly ventral striatal neurons signal “reward” or rather $RPE(-\Delta\%Effme)$ before the rat’s decision, the OFC only afterwards.

In market *economics* goal-specific objective effort, being the essential aversive variable related to appropriation, can be called “*value*”, and sellers initially cash in on *unaccustomed* economies in effort since hereby they can buy “any” good representing *more effort* (value) with less effort, which on average holds true after transformation into subjective residual value represented by $RPE(-\Delta\%Effme)$. In *dyn4* this momentarily advantageous (positive) reduction of *value* figures as perception of a resource-saving cumulative *1-D-index* of net appropriative experience relevant to the early **AW**-module **Worry-T~/a~/m~**. The dynamics of actions and the individual utility of things and services (“worth”) instead attain to $(5+D-)$ Action.

That such acquired evaluative neo-Pavlovian “reflexes” obey to a *ceiling of effort* may find support in the resurrection

of memory and learning in the study of primary sensory cortices [31]. It also seems even compatible with the various *trait-like marginal utility curves* [31] generated from “rational choice” via the unlikely “definition of value” as that from which “one can’t get enough of” [11] in the pure monetary form. Nor anything formally scrunches thereafter when “utility is taken to be correlative to (...) want” - or by *collapse* with “worth” - what “a person is willing to pay” for satisfaction [33], in some other currency. Finally, since **UOs**, and presumably **STs**, are *attracted by the unknowns of computable risk* calling for modeled discounts in effort, they can be expected to enjoy the pure unsaturable monetary shape, of which imponderable worths only cause impure inflections.

5.2. The Divisionary Focus of Habits

This *dualism* of processing styles, here reflected through “**UC-CO**-orientation”, is primarily *non-habitual* and thus different from the two “systems”: either “quick, intuitive, and effortless” or innovative “slow, analytical, and deliberate” – thereby “overcoming intuition.” [34]. In *dyn4* these map to, at first *non-habitual, intuitive* ways of $5+DA$ and to more conscious symbolic processing in $4DT$, often struggling with what it perceives ($A \rightarrow T$) of the complexity of $5+D$ -Action and reaction.

Such *model-based learning (MBL)* besides the testing of hypotheses by relational thinking nevertheless also involves more Gestaltian [35] *4D-pattern* searches, than even intractably costly searches through trees [36].

5.3. When Scare Prefers Grasp over Model

Only the dorsal ACC and the aINS [37] related to $4DT$ ~ and **M~**, not the parietal cortex for $5+DA$, harbor the highly intuitive rapid Von Economo-spindle cells [38], which might have a role in the reductionist *grasping* “Thought-about-Action” $I\{T \rightarrow A\}$ that provides *newly* adaptive fast-and-frugal heuristics. Such prove successful under both habitual and erratic circumstances [39] and are triggered by often just one cue providing “model-free” learning (**MFL**). *Unexplained* adverse complexity via input from locus coeruleus can block experienced model-based functioning in the ACC in favor of such *erratic* behavior, whereby, at times “naturally” simple-minded **CO**-actors [1] become as unpredictable as their surrounds.

5.4. When Modeling Becomes Useful

Yet when routines stop working, **UOs** are needed for their renewal. Then even the *liking* system adapts: once cake is lacking you’ll like bread! (Un-)Certainty-orientation theory [1] shows, that the related affective valences (**M~**) cannot be taken for granted, but reflects a stable *cognitive trait*.

The Uncertainty- oriented (**UOs**) are enabled to maintain the effort required for the conceptual reduction of the complexity of reality by their higher threshold for *cogniti-*

ve-effort-related dysphoria, and by their, at times even *high* intellectual pleasure, Flow [41] denoting the process from Bliss to Interest in *dyn4*.

The Certainty-oriented (COs), being unblest by opportunity-optimizing modeling tasks, instead use rapid reductive “chunking” into essentially *simple commands*, to govern, intrinsically complex (!), habits.

While *lack of mental flexibility* can lead to apparent CO, this usually comes with some loss of intelligence, which is *not* a feature of COs [1], but rather due to disturbances at the medio-dorsal thalamus [42]. The focus on such cognitive efforts presently rejuvenates psychiatry [43, 44].

5.5. Collapsing Duality by Arithmetical Lumping

One account of the ST-problem [36] avoids this double character, or “disunity” [45] of “value” or “worth”, by proposing a *halfway mixed* dual system, whereby the STs are judged by the standards of the cage to be underperforming. MFL through sequential “hot-or-cold”-attempts, here represents “*both* habits and incentive salience” of Pavlovian “reflexes”, while tree-comprehensive permutational, alias model-based, 4D-T-learning (MBL) supposedly relates “to goal-directed valuation, be it instrumentally or in Pavlovian settings”. This commixture, ensuing from a crude reduction of intentional planning, cannot represent habits as entrenchment of instrumental learning nor decide the issue.

5.6. Mood Interacts with Attitude towards the Not-yet-explained

When it comes to positive, or negative, *affect*, what is interesting therefore is, that in the COs the valence of M~ decreases with increasing 4DT~, whereas the UOs enjoy intellectual adventure proportionally [1]. Actually M~ *interacts* with (Un)-Certainty-orientation: while UOs or COs are defined under the premises of a longing for “maximal achievement”, under the clinically depressive premises of a *minimization of further losses*, the two extreme types, at a certain point of prudence, switch into their mirror cognitive style [46, p.6]. Rats who have lost goals [47], or which know them for longer [48], switch from GT to ST, and certain “animals shift their preference from stable to variable food sources under (...) increased physical effort or falling energetic(s).” [26]. Furthermore both orientations *gain with high spirits*, which steepen their opposed regressions.

Yet the *yet-to-be-explaineds* in foraging become dramatic only on arid, not on lush meadows, so at the end of despair COs, after *transient* UOs-ness, would become COs again, sticking to live-saving solutions. Yet goal-tracking rats (GTs) with empty pots would start searching farther away than desperate STs.

Teasing through contextual unknowns increases ST further [50-52], maybe because then cues signal lesser efforts unreliably [53]. Tracking-attitude may also depend on *status*, which is partly inborn: Dominant members often eat first and submissive members are more successful if they use

innovative hypotheses, i.e. if they look more out for cues, than for the food proper, since in the first case they may eat unnoticed by the dominants, while in the second case, they may end up only knowing where the food is others eat.

5.7. Hiding Circularity

The *circular* account is obviously lacking in the recent Bayesian “Active Inference” model, which *seemingly solved circular explanations* of “reward”, whereas this circularity just mirrors the essence of AWs informed by attributes of homeostasis or growth and as such *should not* be solved. The model in fact makes *unpredictability, precision and salience collapse* into “Active Inference” or what midbrain DA supposedly codes for, conveying how (active) “perception minimizes exteroceptive prediction errors and action minimizes proprioceptive prediction errors.” [54]. The model in fact reflects the again truly marvelous discovery, that certain dopamine neurons under conditioned stimuli proportionally code for *unpredictability* of reward [55], but *unpredictability* unduly replaced reward and salience, whereas reality is more complex [13].

At least in the cortex response variability furthermore seems not be a solid foundation, since any stimulus causes its decline [56]. Albeit *unpredictability, precision and salience* determine the *value* in finance industry, they do not account for all facts in the life of rats.

6. Mast Cells, Histamine, and Thalamus

Mast cells (MCs) enter the brain during development, and these cells are replenished [57] or augmented by additional MCs which rapidly immigrate upon signals, which reflect social events, germs, drugs or physical changes [58] or, why not, sexual rubbing. Cerebral MCs are usually found to be scarce, yet most densely present at the thalamus, the habenulae, the olfactory bulb, and within the meninges. Via the braking habenulo-mesencephalic loops [59] MCs guard the blood-brain barrier [60,61] or trans-granulate into neurons [62], and thus probably modulate incentives.

The *perivascular access* to the parenchyma of these sites is wide open in the subcortical [63], but obliterated in the cortical locations. The induction of MC degranulation in the *thalamus* of rats caused excitation (70% in females, 11% in males), or inhibition (7% in females, 33% in males) of thalamic neurons [64]. Positive affect accompanies the behavioral invigoration triggered by MCs under several social circumstances, e.g. during courtship [65] in the medial habenulae. Female rats after cohabitation increase thalamic MCs within the medial geniculate and four other thalamic nuclei [66], whereas in mice not thalamic, but meningeal degranulation of MCs correlates with wakefulness and stimulatory tone in the CNS [67].

6.1. Mast Cells and Thalamo-frontal Driving Feedbacks

Dominant *thalamo-frontal* influences have been recognized in several domains [68-70]. These occur within the

largely *segregated*, and thus dimensionally orthogonal, CSTCs [71], wherein cortical inputs to the basal ganglia are conveyed back to the cortex via the thalamus. Several of the many MC mediators [72] and effects could plausibly modulate the thalamus. While within the CSTCs the striatum is driven through glutamate by plentiful excitatory input from the PFC, and by thalamo-striatal connections, MCs intriguingly only potentiate excitotoxicity via histamine (HA) [73], but are not glutamate-releasers, while requiring it for degranulation [74].

Nevertheless HA, commonly of MC origin, selectively *potentiates* N-methyl-d-aspartate receptors (NMDARs¹) allosterically on a magnesium-sensitive NR2B¹-site [75] also involved in hallucinogenicity [76]. Such an increase of *glutamatergic* activity could plausibly impact on the functioning of CSTCs. While brain HA was normal in mice deficient in MCs, HA in rats was shown to stem from MCs up to 90% in the thalamus and to half in the brain [77]. The latter findings presumably also reflect a more activated state of MCs, wherein large amounts are secreted. Rat cerebral MCs were nearly all thalamic and specifically found in three areas and in the *sign-tracking-related* paraventricular nucleus of thalamus (PVT) [78].

6.2. Mast Cells as Agents with Destination and Destiny

It is tempting to investigate, with cohorts of MCs, after peripheral priming of *destination and destiny*, would often migrate to the brain, where they would lastingly influence subcortical and cortical modules. In the striatum they interact with *perivascular* nerves, which are the fastest *first responders for midbrain DA* [79], or with cells of the neuro-vascular unit. Thereby physiological inflammatory processes could be pushed beyond temperament to “affective temperaments” [80], anxio-phobo-affective diagnoses or to soft, yet often deteriorating, bipolar mixed states [81].

Specifically the depressive switch into opposite tracking or (Un-)Certainty [1,46] mode could be modulated by MCs primed to cause “depression” (instead of hypomanic lesional “vigor”) when reaching the PVT, via the thalamo-perforate and thalamo-geniculate arteries.

The PVT in any case achieves its importance for *cue-reward pairing* through its broad subcortical and prelimbic cortical afferents and glutamergic efferents [82].

7. Scopes and Outlooks

The achievement of *logically expressed* convergent high-quality research is a daunting task, especially if practical *clinical utility* remains a goal. Huge global projects like the Research Domain Criteria (RDoC) initiative are under way [83], which astonishingly removed movement from the classical triad [84]. Furthermore especially the understanding of intrinsically *dimensional* topics, e.g. of CSTCs providing *dimensional data*, seems relevantly hampered by the habitual avoidance of *direct* acquisitions of

“unplugged” i. e. *intact* patterns, conservable e.g. through the Configural Frequency Analysis of Gustav Lienert [85, 86]. This intriguing neglect could be related to UC-orientation [87].

This essay, which takes advantage of the author’s tiny context, strives to contribute a sketch of an anxio-affective framework for such convergence to epistemology and clinical *talk* alike, *dyn4* being also progressively expanded to dyadic or family system relations. The essay focuses on an *Aristotelian geometric classically triadic dyn4-T-A-M*-model newly interpreted as contrast between private low-, and public high-*dimensional processing*. The comprehensive **AW**, as inscribed in the cube and attracted by its dichotomic permutations, apparently has not been proposed as the basic sequence of behavior before.

7.1. Triadic Models in Clinical Psychology

An entwined “adolescent” *triadic* model [88,89], which centered on maturing balances of self-control [90], had instead cut across the T-A-M-dimensions in a not comprehensively orthogonal way. It generated a three composite factor balance between a) a mainly *subcortical cognitively* (actually movingly) *impulsive non-delaying approach driven by reward*, stemming even from “risk taking” (Joy-t~A~M~), b) a *prefrontal cognitively reflected overall control* (T~), and, beyond “dual systems”, c) an *amygdaloid emotionally deranged avoidance* (a~, m~). All the same e.g. it doesn’t accommodate *amygdalar salience* or *striatal expectancy* [91] or the *affective temperaments* [92] well - nor the “inextricable” “interactive dexterity” emerging from studies, not allowed to be constrained into *mechanistic orthogonality* [93], albeit maybe just this would support the sought independence from valence, besides providing systematic complex predictions amenable to non-tautological falsification. Classically triadic instead is the influential *associative, sensori-motor, and limbic* tripartite division [59], albeit some sensations are low-dimensional.

7.2. Are Neuro-economics “Dual or Not”?

Intervening also into the present (*neuro*-)economical debate this essay calls for “circular” experiments avoiding undue “collapse” between low-dimensional “value”/effort and high-dimensional “worth”/utility. This is now strongly supported by the first localization of the positive human *interactive* “value-to-utility transformation” to the inversely activated dorsal anterior *mid-cingulate cortex* (dmPFC). The connectivity of this is positive with the probably uphill *inferior frontal gyrus* [94], related e.g. to *perceptive* “confidence” (certainty) [95], and negative with the probably downhill Nac providing *intentional* subjective “valuation”. Within the OFC instead “value” seems stored behind, and “worth” in front [96]. Others shed doubt on the necessity of emotions as mediators of mesolimbic dopaminergic effects e.g. on feeding, but explore a more abstract “facilitation of a specific form of neural computation” [97], maybe an

expansion of complexity.

Impressive reviews of *transfer* [98,99] concede that despite guaranteed rewards, cues still work, whatever the “worth” [100], while others focus on “efforts” [101]. Importantly the “dual” arguments also apply when “worth” equals *inviolacity* [102].

By conceiving the *incentivity* of cues to be due to opportune *effort*-reducing and thus facilitating means signaled by the cue, their three [103] attributes – *attractivity* (for agent’s attention or approach), *instigation* of effort to reach them, *energizing* of appropriation of reward – can be accounted for.

An *orthogonal* conceptualization of the segregated albeit cortico-cortically linked CSTCs and their “intentional” function is maintained in some frameworks, at least for *A*~ and *M*~ [104], yet in a *collapsed* way avoiding *T*~. Also the “orthogonalization” between motricity and reward again avoids the duality of *4D-T*~ and *5+D-A*~ and just deals with *5+D-A*~ and its first derivative: in fact more experimental distinction of *MBL* and *MFL* is being asked for [105].

7.3. Habits Are Not Always Rapid or the Primary Issue

Within the *dyn4*-framework, “models” are created in *4D-T*~ with *cognitive effort*, but *incentivated* by the perspective of a variously [106, 107] tempting discount in *antici-pated* appropriative *5+D-A*~ related mainly *motor effort*, as signaled by cues functioning in such a model.

Motor-related *habits* [108], albeit dopaminergically [109] crystallized, are *still complex* programs reformatted away from globus pallidus [110] through various processes, e.g. TGFβ- [111] or NMDAR- [112] activity on striatal DA-neurons. The lack of the latter *glutamatergic* input slows down learning, social contacts and forced swimming, but *not effortful* performance [113].

Some neuro-economically engaged clinicians say themselves not yet content [114] with their differently dual complexity *MBL/MBF*-approach contrasting “more complex”, “goal-directed” *MBL*-based behavior with alternative habitual *MFL*-based decision making [115]. They e.g. showed that rises in ventral striatal (VS) DA correlate with *MBL*-related “signatures” in dlPFC and inversely with *MFL*-related encoding in VS [115] - both being conceived as *4D-T*~ in *dyn4*. Their alternative habitual *MBL*-processing instead as such in *dyn4* would remain “complex” as related to *5+D-A*~ Action, despite requiring less *4D-T*~ related conscious steering or being shielded from outcome-perceiving feedback. This delayed habitual reformatting by repetition may actually *detract* from the problems collapsed neureconomics encounter in mental care.

The sufficient checking of a purportedly rapid less effortful intuitive system-1 (*5+DA*) by a more reflective system-2 in otherwise biased decisions, has been strongly complemented by a core role of even *less* intuitive numerical abilities (*4DT*) [116].

7.4. Neurobehavioral Complexity Changes and *dyn4*

Following *dyn4* it would have to be explored wither the CSTCs involving the ACC assigned to *5+D-A* really show a higher e.g. fractal dimensionality than the one involving the dlPFC mapped to *4D-T*. In fact the dorsal ACC itself already produces *neuro-economical* [117] *reductive* models of conflicting past and present experience [118] ready to feed decisions to be taken in dlPFC, linking contexts with *appropriative*, and therefore lastly *motor strategies* by producing a rich “task space” [119].

Since *appropriation* is the organizing principle in *dyn4 motricity* (*5+DA*) is in command of secondary parietal or primary motor areas. Similarly the CSTC involving OFC / vmPFC [120], and not the “limbic systems”, represents *Mood*, as they master economical emotions [121] and integrate emotionally valenced “worth” to command appropriations [122].

The CSTCs themselves being feed-forward structures show an about 500-fold quantitative neuronal reduction in “complexity” between striatum and the pre-thalamic inhibiting output components. Of these the substantia nigra (SNr) e.g. may “gain control” over cortical feedback when sparing explorative efforts [123], braking “complicated” *5+DA*. Rodent-primate homologies of CSTCs are many and also related to psychiatric models [124].

The present *dyn4*-account also implies that the *alternating coordination* between *4D-T*- and *5+D-A*-processing is an enlightening prerequisite, beyond basic divergence and funneling, for any functional brain activity. This occurs within *4D-T* and is often dealt with as top-down *attention*. Recent theoretical shifts towards considering the dorsal attention network (DAN) within the *fronto-parietal cortices* as a *common* substrate of “internal attention” sustaining as variegate functions as working memory, episodic retrieval of *percepts*, and *intentionally complex* mental imagery [125] supports this crudely mechanistic prediction on a high level of sophistication.

7.5. Effortful Controls of Thought, Action, Mood

We tend to fuse the concepts of *subjective effort* - the emotion of cost - as the felt *passive* brake on expenditure of resources, and again the *overcoming active* effort throughout the initiation and maintenance of effort-full processes. The steering of motor-costs is primordial, while the pleasure-systems and their hot-spots in evolution are small and marginal [126]. In humans though the costs of emotions, like the one from the urge of want, and their cognitive costs can become predominant. Thus, besides some focus on duration [127, 128], mainly the cost of *suppressing* emotions is monitored [129]. Fortunately *affect dynamics* are taking momentum also in the case of a likely alias of increased *emotional effort* suspected to be a pre-depressive signature of decoupling from usual functional connectivity: rigid *emotional inertia* [130].

These modules related to effort-ful appropriation have just been assembled in a formidable review [131]: The *ventral striatum* (VS) including the nucleus accumbens (NAc) activity likely is due to a momentaneous internal Perception of the opportune *simple* ratio of utility to effort, which continues when utility reaches a ceiling, helped by the midbrain. The VS *invigorates* appropriative action and the *momentaneous changes in dopamine* correlate with the *willingness* to work, which correlates with *incentive* cueing, even in absence of reward! Demanded *high-effort* choices, which need permission by the ACC, instead activates the *amygdala*, which seems to aid in encoding of *relevance* of the former inner and external Perceptions. It gives rise to *urges*, also in associative learning, whereby DA is released in the NAc, and the ACC is instructed to allow for high-effort expenditures. The amygdala overall acts as a conservative or investing *expense controller* in front of the *temptations* provided by the VS.

The *dorsal striatum* plans, decides, and automatizes *motor* behavior often into *habits*, while it also monitors internal *metabolic* and even external *nutritional* resources. As a result it encodes specific energetic prerequisites for appropriation. Mice without DA instead die from aphagy, while hedonics and spatial learning of food remained intact. In *dyn4* this preparation of Action corresponds to *Intention*. Within the dopaminergic midbrain the VTA and SNr interact with the striatum and thereby seem to provide the expected average *opportunity* on appropriation with the specific effort. Yet the amphetamine-sensitive emotional *drive*, as computed from the latter costs and delays combined with „subjective value“ (*worth*) and the variable confidence in consequential *Intentions*, is provided by the vmPFC. The ultimate *decision* is taken around the intraparietal sulcus.

In *dyn4* this corresponds to Interest or Worry leading via Application or Remediation to Pursuit [2, Table 1].

The supplementary motor areas (SMAs) monitoring *muscle* contraction interestingly feeds into subjective *effort*, while through preparedness it may invigorate response or maybe inversely spare effort.

7. 6. Momentaneous Fluctuations Could Confirm dyn4

This same review [131] then attracts attention to the meaningful information hidden in the momentaneous *fluctuations* in cognitive and physical *effort*. Albeit the *speed-accuracy trade-off* is pervasive, it becomes hidden in the context of higher rewards, by which *both* increase. In psychiatry instead the new validated concentration deficit disorder (CDD), former „sluggish cognitive tempo“, which has replaced most of ADHD-inattentive type, is *not* an executive disorder, but strangely reminiscent of a *coupled inertia* of $T\sim$, $A\sim$, and $M\sim$, related to depression and refraining from higher intensities in all three dimensions [132].

In *dyn4* accurate *distances* are a result of $5+DA\sim$, while $4DT\sim$ is related to appropriations in a vage *future* and their *speed*. Reward-induced *invigoration* along the Appropria-

tion-axis „from Need to Pursuit“ in fact causes a symmetric intensification in $T\sim$, $A\sim$ and $M\sim$. Through this analogy we start to consider coupled, usually skewed, *simple harmonic oscillators* (SHO as a *mass-on-a-spring* with $(-k/m)\cdot x = d^2x/dt^2$ obeyed by $\sin(x)$ or $\cos(x)$) as a biaxial [133], not mono-axial [134] model of „mood swings“. Hereby the *above momentaneous fluctuations* of $T\sim$, $A\sim$, and $M\sim$ are modeled, which putatively correspond to the three „affective“ CSTCs. The CSTCs by virtue of their direct and (negative) indirect paths, in fact could be approximated as SHOs.

Silvain Tomkins modeling of emotions as *analogue amplifiers of intensity and its first derivative* over time with as prime role for muscle sensibility [135] here appears as very much to the point and compatible with *dyn4*.

7.7. Biopersonology and dyn4-TAM

While *dyn4-TAM* can probably be best mapped to a bio-amine-centered model of personality e. g. by Richard Depue [136], rapidly-acting ketamine-related or cholinergic antidepressants have deviated attention from these systems (see 7.5.).

The specifically cholinergic molecular loss of function in STs [137] points strongly to the fact that *cholinergic* systems support *anti-distractive* cognitive control, whilst also allowing for *attentive shifts* with reorientation to cues and cue-re- sponsive action [113], like approaching the goal!

Present psychological research on humans applies the concept of „ST-to-GT“ [107] and could use cross-validating tests for „UC-to-CO“ [1,46], while studying resistance to *temptation* or *effort* [106] would also test the here exposed hypothesis of homology.

7.8. Mast Cells at the Reins of Appropriation?

As to the own hypothesis, that the anatomical convergence of the three “affective” CSTCs at the *thalamus* might provide access especially for *mast cells* [138] intruding along the posterior arteries to modulate subcortical logistics, some few observations concur. Since STs are high in ventral HC myo-inositol, and hereby dopaminergically incentivize Nac in Pavlovian approach [139], putative roles both of hippocampal mast cells [140] and of lithium [141], inhibiting IMPase [142], emerge, which hint to how the convergent framework *dyn4* could operate in affective disorders. In fact cues become *less* incentive under ketamine [143], the miracle antidepressant pro-drug which acts by upregulating AMPA-receptors [144], which happens to incite [145] or to calm mast cells [146], but does not affect midbrain DA [147].

The latest review on STs [148] points also to the *lateral habenulae* (LHb) [149-151], and thereby, see below, also to *mast cells* (MCs), as a part of the food-cue-induced “motive circuit”, and its rapid adaptations. Within a larger network [152] they help in attributing *salience* [153] to the point, that

the LHb *drives* the VTA and SNr during RPE [154]. The LHb specifically act as indirect [155,156] strong *inverse* modulators [153] of the DA of the midbrain's SEEKING system [157], and the playfulness of STs is supported by the centrality of LHb for *social play* [158, 157]. Conversely LHb and the medial habenula (MHb) are sensitive in the non-depressed to present [159] or future *punishment* [160] up to *learned helplessness* [161, 162], produce *vegetative costs* of emotions [163], and *shrink* [164, 165] especially in bipolar depression. Drugs inhibiting LHb reverse resistant depression [166].

As to MCs [58] they rapidly intrude after psychosocial events as acute activators e.g. into the LHb after repeated defeats [167], and with parenthood [168] into the reinforcing MHb of which silencing is aversive [169]. Mastocytosis finally is depressiogenic [170].

8. Conclusions

An essay "takes things from many sides without comprehending it fully." [171], and as a vivid genre of troubled times it takes high risks to fall victim to its own boldness by loosening cognitive control. Furthermore Karl Jaspers justly warned: "Theorizing has an atmosphere of its own." [172]. Progress in fact mostly, but not entirely, occurs through painstaking continuity of endeavors critical in seeking better lives, and the Ann Kelley's saga, certainly testifies to this. In the commemorative volume to her and also by her lab, which added successes with the STs and GTs rats, John D. Salamone contributed insights into the central role of Nac in bringing about *effort*-related choices [173]. This would also explain the common failures of RPE to behave in schizophrenia as computational psychiatrist, which build around it, were hoping for [176], and in this their "orthogonal" tautological Bayesian relations, which are akin to any reciprocal falsification couple between theory and hypothesis, will not be of any comfort. Till date highly erudite accounts on "cues" [177] can still make it without "effort", but the two lines of enquiry will not continue on parallel tracks with little convergence for long.

The conceptual skipping of Thought and Intention in the Perception-Action models, even in their most erudite form [178] remains puzzling (to me), while the intricacies of *relational* cognitive processing explore the limits of complexity of Thought [179] - till now without a factoring-in of the "UC-to-CO"-algorithm. The Intention to think, act, and feel is certainly enriched by their "economical" braking by *effort*, and the second response component in midbrain DA-neurons, which codes reward value as a "numeric, quantitative utility prediction error", [180] could be a predictor of *opportunity* of less effort, and not of utility (worth), since it starts early enough to prevent "confusion with unrewarded stimuli and objects."

Especially in the NAc DA participates in *effort*-based choices among often many opportunities in the surrounds of

the niche. Variability in active effort has been referred mostly to fluctuations in subjective anticipated effort [131]. Research on *emotional* effort is centered on the control or *suppression* of emotions, and maybe today research on the neuroeconomy of *confidence* [181] is most on track in this area, since confidence in a cheap simple heuristic model comes at a rarely relevant [39] price of error. Also *dyn4* knows the processes of **D**oubt or **C**onfidence [2: Table 1] dealing with *opposite* interactions between modeling-in-4DT and **M**ood. The proposal that the incentivivity of such cues are about *specific hope*, that the expected required effort will be discounted, and that **H**ope in *dyn4* is the passage from **N**eed-tam to **I**nterest-TaM in *dyn4*, is compatible with positive psychotherapy using *incentive hope*.

Survival depends on movement, movement on motivation, and motivation on cost-benefit analyses of active effort expressed in passively anticipated effort? Optogenetics on *freely moving* rodents will soon tell us more about this [181]. The SMA's feeding of muscle *contraction* into subjective effort [129] reminds us of the *cutaneo-muscularly felt* emotion theory of Silvain S. Tomkins [135] or recalls oro-facial mimics of „li(c)king“ [182] or the *clenching of teeth* to increase active effort, but it can't sustain the claim, that the felt quality of the „SEEKING“ system model would allow to collapse the *duality* of learning in approach [157].

A recent assembly of research on *circuits of positive emotions* [184], to which this paper originally was submitted as an elaboration of a poster, confirmed, that the important, because extremely basic hypothesis of a reinforcement or reversal of the *prime motivator* (Un-)Certainty-orientation of Richard M. Sorrentino [1,46] by valenced emotion, which reveals a constituting evolutionary link between cognition and emotion, and the mast cells are just only starting to attract noticeable interest in the mainstream of neurosciences. Therefore pioneers in ST-research [126] and MC-research related to the brain [58], Rae Silver, now leader in circadian rhythms, not fully by chance are immediate neighbors in a monograph on motivation [185], and the links likely also run via "clocked" [186] and "clocking" [187] MCs in brain [58] and other tissue [188] in relation to bio-economic metabolism [189] and its central "subjective" variable *effort* [190].

Acknowledgements

This work became possible through the legacy from my sister Fiorenza Hohl.

Conflicting Financial Interests

None.

REFERENCES

- [1] R. M. Sorrentino. Uncertainty orientation: A theory where the exception forms the rule. *Revista de Motivación y Emoción* 1,

- 1 – 11, 2012 Online available from http://reme.uji.es/reme/1-sorrentino_pp_1-11.pdf
- [2] G. R. S. Treviranus. For value or for worth? Part 1: Speaking syllogistic attractors of appropriations in a thought-action-mood-space. *Universal Journal of Psychology*, 5(2), 42-49, 2017.
- [3] Trubenová, R. Hager. Social selection and indirect genetic effects in structured populations. *Evolutionary Biology* 41(1), 123-133, 2014. doi:10.1007/s11692-013-9252-5
- [4] Plato. *The Republic*. Transl. B. Jowett. J. Manis (Ed.). The Electronic Classics Series Hazleton, PA: PSU-Hazleton. Online available from <http://worldlibrary.org>
- [5] Anonymous. Some thoughts on the interest of money in general, and particularly in the publick funds,.. London, 1739 or 1740. Cited in: K. Marx, *Capital*, vol. one. Chp.1. Sect.4. The fetishism of commodities and the secret thereof. Annot. 9. [Transl. Ben Fowkes 1990. London: Penguin Books.], 1867.
- [6] S. B. Ostlund, K. M. Wassum, N. P. Murphy, B. W. Balleine, N. T. Maidment. Extracellular dopamine levels in striatal subregions track shifts in motivation and response cost during instrumental conditioning. *Journal of Neuroscience* 31(1), 200-207, 2011. doi: 10.1523/JNEUROSCI.4759-10.2011
- [7] Y. Li, G. Vanni-Mercier, J. Isnard, F. Mauguière, J. C. Dreher. The neural dynamics of reward value and risk coding in the human orbitofrontal cortex. *Brain* 139(Pt 4), 1295-1309, 2016. doi:10.1093/brain/awv409
- [8] W. Schultz, P. Dayan, P. R. Montague. A neural substrate of prediction and reward. *Science* 275(5306), 1593-1599, 1997. doi:10.1126/science.275.5306.1593
- [9] D. C. Castro, K. C. Berridge. Advances in the neurobiological bases for food 'liking' versus 'wanting'. *Physiology, Behavior* 136(0), 22-30, 2014. doi:10.1016/j.physbeh.2014.05.022/
- [10] J. P. O'Doherty. The problem with value. *Neuroscience, Biobehavioral Reviews* 43(1), 259-268, 2014. doi:10.1016/j.neubiorev.2014.03.027/
- [11] W. Schultz. Neuronal reward and decision signals: from theories to data. *Physiological Reviews* 95(3), 853-951, 2015. doi:10.1152/physrev.00023.2014
- [12] H. M. Bayer, P. W. Glimcher. Midbrain dopamine neurons encode a quantitative reward prediction error signal. *Neuron* 47(1), 129-141, 2005. doi:10.1016/j.neuron.2005.05.020
- [13] C. D. Fiorillo, M. R. Song, S. R. Yun. Multiphasic temporal dynamics in responses of midbrain dopamine neurons to appetitive and aversive stimuli. *Journal of Neuroscience* 33(11), 4710-4725, 2013. doi:10.1523/JNEUROSCI.3883-12.2013
- [14] E. S. Bromberg-Martin, O. Hikosaka. Lateral habenula neurons signal errors in the prediction of reward information. *Nature Neuroscience* 14(9), 1209-16, 2011. Erratum (print only) in: *Nature Neuroscience* 14(12), 1617. doi:10.1038/nn.2902/
- [15] H. W. Chase, P. Kumar, S. B. Eickhoff, A.Y. Dombrovski. Reinforcement learning models and their neural correlates: an activation likelihood estimation meta-analysis. *Cognitive, Affective, & Behavioral Neuroscience* 15, 435-459, 2015. doi:10.3758/s13415-015-0338-7
- [16] L. H. Corbit, B. W. Balleine. *Learning and Motivational Processes Contributing to Pavlovian-Instrumental Transfer and Their Neural Bases: Dopamine and Beyond*. In: E. H. Simpson, P. D. Balsam (eds.). *Behavioral Neuroscience of Motivation*. Current Topics in Behavioral Neurosciences. Vol. 27. (pp 259-289). Springer Int., Cham ZG, 2016. doi:10.1007/7854_2015_388
- [17] S. Kaveri, H. Nakahara. Dual reward prediction components yield Pavlovian sign- and goal-tracking. *PLoS One* 9(10), e108142, 2014. doi:10.1371/journal.pone.0108142
- [18] R. A. Rescorla, R. L. Solomon. Two-process learning theory: relationships between Pavlovian conditioning and instrumental training. *Psychological Review* 74,151-183, 1967.
- [19] M. R. Bailey, E. H. Simpson, P. D. Balsam. Neural substrates underlying effort, time, and risk-based decision making in motivated behavior. *Neurobiology of Learning and Memory*. 133, 233-256, 2016. doi:10.1016/j.nlm.2016.07.015
- [20] M. C. Klein-Flügge, S. W. Kennerley, K. Friston, S. Bestmann. Neural signatures of value comparison in human cingulate cortex during decisions requiring an effort-reward trade-off. *Journal of Neuroscience* 36(39), 10002-10015, 2016. doi:10.1523/JNEUROSCI.0292-16.2016
- [21] P. L. Croxson, M. E. Walton, M. F. Rushworth, D. M. Bannerman. Unilateral medial frontal cortex lesions cause a cognitive decision-making deficit in rats. *European Journal of Neuroscience* 40, 3757-3765, 2014. doi:10.1111/ejn.12751
- [22] M. C. Klein-Flügge, S. W. Kennerley, A. C. Saraiva, W. D. Penny, S. Bestmann. Behavioral modeling of human choices reveals dissociable effects of physical effort and temporal delay on reward devaluation. *PLoS Computational Biology* 11(3), e1004116, 2015. doi:10.1371/journal.pcbi.1004116
- [23] C. Prévost, M. Pessiglione, E. Météreau, M. L., Dreher, J. C. Cléry-Melin. Separate valuation subsystems for delay and effort decision costs. *Journal of Neuroscience* 30(42), 14080-14090, 2010. doi:10.1523/JNEUROSCI.2752-10.2010
- [24] R. Kurzban, A. Duckworth, J. W. Kable, J. Myers. An opportunity cost model of subjective effort and task performance. *Behavioral and Brain Sciences* 36(6), 661-679, 2013. doi:10.1017/S0140525X12003196
- [25] S. M. Nicola. Reassessing wanting and liking in the study of mesolimbic influence on food intake. *American Journal Physiology Regulatory, Integrative, and Comparative Physiology*. 311, 5, R811-R840, 2016. doi:10.1152/ajpregu.00234.2016
- [26] M. A. Miller, A. Thomé, S. L. Cowen. Intersection of effort and risk: ethological and neurobiological perspectives. *Frontiers in Neuroscience* 7, 208, 2013. doi:10.3389/fnins.2013.00208
- [27] C. J. Burke, C. Brünger, T. Kahnt, S. Q. Park, P. N. Tobler. Neural integration of risk and effort costs by the frontal pole: only upon request. *Journal of Neuroscience* 33(4), 1706-1712, 2013. doi:10.1523/JNEUROSCI.3662-12.2013/
- [28] B. Y. Hayden, J. M. Pearson, M. L. Platt. Neuronal basis of sequential foraging decisions in a patchy environment. *Nature Neuroscience* 14(7), 933-939, 2011. doi:10.1038/nn.2856
- [29] N. Kolling, M. Wittmann, M. F. Rushworth. Multiple neural mechanisms of decision making and their competition under

- changing risk pressure. *Neuron* 81(5), 1190-1202, 2014. <https://doi.org/10.1016/j.neuron.2014.01.033>.
- [30] C. Giorgetta, A. Grecucci, N. Bonini, G. Coricelli, G. Demarchi, C. Braun, A. G. Sanfey. Waves of regret: a meg study of emotion and decision-making. *Neuropsychologia* 51(1), 38-51, 2013. doi:10.1016/j.neuropsychologia.2012.10.015
- [31] W. R. Stauffer, A. Lak, W. Schultz. Dopamine reward prediction error responses reflect marginal utility. *Current Biology* 24(21), 2491-2500, 2014. doi:10.1016/j.cub.2014.08.064
- [32] B. W. Balleine. *Sensation, Incentive Learning, and the Motivational Control of Goal-Directed Action*. In: J. A. Gottfried (Ed.). *Neurobiology of Sensation and Reward*. Boca Raton, FL: CRC Press, 2011.
- [33] A. Marshall. *Principles of Economics*. An introductory volume. 8th ed. London, UK: Macmillan, 1920. (p.78).
- [34] A. L. Alter, D. M. Oppenheimer, N. Epley, R. N. Eyre. Overcoming intuition: metacognitive difficulty activates analytic reasoning. *Journal of Experimental Psychology: General* 136(4), 569-576, 2007. doi:10.1037/0096-3445.136.4.569
- [35] P. R. Montoro, D. Luna, J. J. Ortells. Subliminal Gestalt grouping: evidence of perceptual grouping by proximity and similarity in absence of conscious perception. *Consciousness and Cognition* 25(1), 1-8, 2014. doi:10.1016/j.concog.2014.01.004
- [36] Q. J. Huys, P. N. Tobler, G. Hasler, S. B. Flagel. The role of learning-related dopamine signals in addiction vulnerability. *Progress in Brain Research* 211, 31-77, 2014. doi:10.1016/B978-0-444-63425-2.00003-9
- [37] L. Q. Uddin, J. Kinnison, L. Pessoa, M. L. Anderson. Beyond the tripartite cognition-emotion-interoception model of the human insular cortex. *Journal of Cognitive Neuroscience* 26(1), 16-27, 2014. doi:10.1162/jocn_a_00462
- [38] H. C. Evrard, T. Forro, N. K. Logothetis. Von Economo neurons in the anterior insula of the macaque monkey. *Neuron* 74(3), 482-489, 2012. doi:10.1016/j.neuron.2012.03.003
- [39] G. Gigerenzer, H. Brighton. Homo heuristicus: Why biased minds make better inferences. *Topics in Cognitive Science* 1(1), 107-143, 2009. doi:10.1111/j.1756-8765.2008.01006.x
- [40] D. G. Tervo, M. Proskurin, M. Manakov, M. Kabra, A. Vollmer, K. Branson, A. Y. Karpova. Behavioral variability through stochastic choice and its gating by anterior cingulate cortex. *Cell* 159(1), 21-32, 2014. doi:10.1016/j.cell.2014.08.037.
- [41] J. Nakamura, M. Csikszentmihalyi, M. *Flow theory and research*. In C. R. Snyder and S. J. Lopez, (Eds.), *Handbook of Positive Psychology*. (pp. 195–206). Oxford: Oxford University Press, 2009.
- [42] S. Parnaudeau, K. Taylor, S. S. Bolkan, R. D. Ward, P. D. Balsam, C. Kellendonk. Mediodorsal thalamus hypofunction impairs flexible goal-directed behavior. *Biological Psychiatry* 77(5), 445-453, 2014. doi:10.1016/j.biopsych.2014.03.020
- [43] A. Culbreth, A. Westbrook, D. Barch. Negative symptoms are associated with an increased subjective cost of cognitive effort. *Journal of Abnormal Psychology* 125(4), 528-536, 2016. doi:10.1037/abn0000153
- [44] W. P. Horan, L. F. Reddy, D. M. Barch, R. W. Buchanan, E. Dunayevich, J. M. Gold, (...) M. F. Green. Effort-based decision-making paradigms for clinical trials in schizophrenia: part 2 - External validity and correlates. *Schizophrenia Bulletin* 41(5), 1055–1065, 2015. doi:10.1093/schbul/sbv090
- [45] S. B. Ostlund, B. W. Balleine. The disunity of Pavlovian and instrumental values. *Behavioral and Brain Sciences* 31(4), 456-457, 2008. doi:10.1017/S0140525X0800492
- [46] R. M. Sorrentino, C. J. R. Roney. *The uncertain mind: Individual differences in facing the unknown*. Philadelphia: Psychology Press., 2000.
- [47] M. J. Robinson, P. Anselme, A. M. Fischer, K. C. Berridge. Initial uncertainty in Pavlovian reward prediction persistently elevates incentive salience and extends sign-tracking to normally unattractive cues. *Behavioural Brain Research* 266(0), 119-130, 2014. doi:10.1016/j.bbr.2014.03.004
- [48] C. S. Srey, J. M. Maddux, N. Chaudhri. The attribution of incentive salience to Pavlovian alcohol cues: a shift from goal-tracking to sign-tracking. *Frontiers in Behavioral Neurosciences* 9, 54, 2015. doi:10.3389/fnbeh.2015.00054
- [49] D. J. S. Costa, R. A. Boakes. Context blocking in rat autoshaping: Sign-tracking versus goal-tracking. *Learning and Memory* 40(2), 178-185, 2009. <http://dx.doi.org/10.1016/j.lmot.2008.11.001>
- [50] P. Anselme, M. J. Robinson, K. C. Berridge. Reward uncertainty enhances incentive salience attribution as sign-tracking. *Behavioural Brain Research* 238, 53-61, 2013. doi:10.1016/j.bbr.2012.10.006
- [51] M. J. F. Robinson, P. Anselme, K. Suchomel, K. C. Berridge. Amphetamine-induced sensitization and reward uncertainty similarly enhance incentive salience for conditioned cues. *Behavioral Neuroscience* 129(4), 502–511, 2015. doi:10.1037/bne0000064
- [52] P. Anselme. Does reward unpredictability reflect risk? *Behavioural Brain Research* 280, 119-127, 2015. doi:10.1016/j.bbr.2014.12.003/
- [53] A. Funamizu, M. Ito, K. Doya, R. Kanzaki, H. Takahashi. Condition interference in rats performing a choice task with switched variable- and fixed-reward conditions. *Frontiers in Neuroscience*. 9, 27, 2015. doi:10.3389/fnins.2015.00027
- [54] K. J. Friston, T. Shiner, T. FitzGerald, J. M. Galea, R. Adams, H. Brown, R. J. Dolan, R. Moran, K. E. Stephan, S. Bestmann. Dopamine, affordance, and active inference. *PLoS Computational Biology* 8, e1002327, 2012. doi:10.1371/journal.pcbi.100232
- [55] C. D. Fiorillo, J. K. Kim, S. Z. Hong. The meaning of spikes from the neuron's point of view: predictive homeostasis generates the appearance of randomness. *Frontiers in Computational Neuroscience* 8, 49, 2014. doi:10.3389/fncom.2014.00049
- [56] M. M. Churchland, B. M. Yu, J. P. Cunningham, L. P. Sugrue, M. R. Cohen. Stimulus onset quenches neural variability: a widespread cortical phenomenon. *Nature Neuroscience* 13(3), 369-378, 2010. doi:10.1038/nn.2501
- [57] K. M. Nautiyal, C. Liu, X. Dong, R. Silver. Blood-borne donor mast cell precursors migrate to mast cell-rich brain regions in

- the adult mouse. *Journal of Neuroimmunology* 240-241, 142-146, 2011. doi:10.1016/j.jneuroim.2011.09.003
- [58] R. Silver, J. P. Curley. Mast cells on the mind: new insights and opportunities. *Trends in Neuroscience* 36(9), 513-521, 2013. doi:10.1016/j.tins.2013.06.001
- [59] S. Ikemoto, C. Yang, A. Tan. Basal ganglia circuit loops, dopamine and motivation: A review and enquiry. *Behavioural Brain Research* 290, 17-31, 2015. doi:10.1016/j.bbr.2015.04.018
- [60] P. Esposito, N. Chandler, K. Kandere, S. Basu, S. Jacobson, R. Connolly, D. Tutor, T. C. Theoharides. Corticotropin-releasing hormone and brain mast cells regulate blood-brain-barrier permeability induced by acute stress. *Journal of Pharmacology and Experimental Therapeutics* 303(3), 1061-1066, 2002. doi:10.1124/jpet.102.038497
- [61] D. Ribatti. The crucial role of mast cells in blood-brain barrier alterations. *Experimental Cell Research* 338(1), 119-125, 2015.
- [62] M. Wilhelm, R. Silver, A. J. Silverman. Central nervous system neurons acquire mast cell products via transgranulation. *European Journal of Neuroscience* 22(9), 2238-2248. doi: 10.1111/j.1460-9568.2005.04429.x
- [63] Jones, E. G. On the mode of entry of blood vessels into the cerebral cortex. *Journal of Anatomy* 106(Pt 3), 507-520, 1970.
- [64] P. Kovács, I. Hernádi, M. Wilhelm. Mast cells modulate maintained neuronal activity in the thalamus in vivo. *Journal of Neuroimmunology* 171(1-2), 1-7, 2006. doi:10.1016/j.jneuroim.2005.07.026
- [65] X. Zhuang, A. J. Silverman, R. Silver. Reproductive behavior, endocrine state, and the distribution of GnRH-like immunoreactive MCs in dove brain. *Hormones and Behavior* 27(3), 283-295, 1993. doi:10.1111/j.1365-2826.1992.tb00160.x
- [66] L. Asarian, E. Yousefzadeh, A. J. Silverman, R. Silver. Stimuli from conspecifics influence brain mast cell population in male rats. *Hormones and Behavior* 42(1), 1-12, 2002. doi:10.1006/hbeh.2002.1799
- [67] A. A. Larson, M. J. Thomas, A. McElhose, K. J. Kovács. Spontaneous locomotor activity correlates with the degranulation of mast cells in the meninges rather than in the thalamus: disruptive effect of cocaine. *Brain Research* 1395(0), 30-37, 2011. doi:10.1016/j.brainres.2011.04.033
- [68] J. de Bourbon-Teles, P. Bentley, S. Koshino, K. Shah, A. Dutta, P. Malhotra, T. Egner, M. Husain, D. Soto. Thalamic control of human attention driven by memory and learning. *Current Biology* 24(9), 993-999, 2014. doi:10.1016/j.cub.2014.03.024
- [69] B. B. Theyel, D. A. Llano, S. M. Sherman. The cortico-thalamocortical circuit drives higher-order cortex in the mouse. *Nature Neuroscience* 13(1), 84-88, 2010. doi:10.1038/nn.2449
- [70] A. Nakhnikian, G. V. Rebec, L. M. Grasse, L. L. Dwiell, M. Shimono, J. M. Beggs. Behavior modulates effective connectivity between cortex and striatum. *PLoS One* 9(3), e89443, 2014. doi:10.1371/journal.pone.0089443
- [71] M. S. Mega, J. L. Cummings. Frontal-subcortical circuits and neuropsychiatric disorders. *Journal of Neuropsychiatry and Clinical Neurosciences* 6(4), 358-70, 1994. doi:10.1176/jnp.6.4.358
- [72] T. C. Moon, A. D. Befus, M. Kulka. Mast cell mediators: their differential release and the secretory pathways involved. *Frontiers in Immunology* 5, 569, 2014. doi:10.3389/fimmu.2014.00569
- [73] S. D. Skaper, L. Facci, W. J. Kee, P. J. Strijbos. Potentiation by histamine of synaptically mediated excitotoxicity in cultured hippocampal neurones: a possible role for mast cells. *Journal of Neurochemistry* 76(1), 47-55, 2001. doi:10.1046/j.1471-4159.2001.00008.x
- [74] E. K. Hamasato, A. P. Ligeiro de Oliveira, A. Lino-dos-Santos-Franco, A. Ribeiro, V. Ferraz de Paula, J. P. Peron, A. S. Damazo, W. Tavares-de-Lima, J. Palermo-Neto. Effects of MK-801 and amphetamine treatments on allergic lung inflammatory response in mice. *International Immunopharmacology* 16(4), 436-443, 2013. doi:10.1016/j.intimp.2013.04.019
- [75] A. Burbán, R. Faucard, V. Armand, C. Bayard, V. Vorobjev, J. M. Arrang. Histamine potentiates N-methyl-D-aspartate receptors by interacting with an allosteric site distinct from the polyamine binding site. *Journal of Pharmacology and Experimental Therapeutics* 332, 912-921, 2010. doi:10.1124/jpet.109.158543
- [76] E. K. Lambe, G. K. Aghajanian. Hallucinogen-induced UP states in the brain slice of rat prefrontal cortex: role of glutamate spillover and NR2B-NMDA receptors. *Neuropsychopharmacology* 31(8), 1682-1689, 2006.
- [77] R. C. Goldschmidt, L. B. Hough, S. D. Glick. Rat brain mast cells: contribution to brain histamine levels. *Journal of Neurochemistry* 44(6), 1943-1947, 1985. doi:10.1111/j.1471-4159.1985.tb07191.x
- [78] R. C. Goldschmidt, L. B. Hough, S. D. Glick, J. Padawer. Mast cells in rat thalamus: nuclear localization, sex difference and left-right asymmetry. *Brain Research* 323(2), 209-217, 1984. doi:10.1016/0006-8993(84)90291-9
- [79] D. Afonso-Oramas, I. Cruz-Muros, J. Castro-Hernández, J. Salas-Hernández, P. Barroso-Chinea, S. García-Hernández, J. L. Lanciego, T. González-Hernández. Striatal vessels receive phosphorylated tyrosine hydroxylase-rich innervation from midbrain dopaminergic neurons. *Frontiers in Neuroanatomy* 8, 84, 2014. doi:10.3389/fnana.2014.00084
- [80] J. M. Himmelhoch. The sources of characterological presentations of mixed bipolar states. *Clinical Neuropharmacology* 15 (Pt. A) 630A-631A, 1992. Available online from <http://journals.lww.com/clinicalneuropharm/Citation/1992/01001>
- [81] G. S. Malhi, L. Lampe, C. M. Coulston, M. Tanius, D. M. Bargh, G. Curran, S. Kuiper, H. Morgan, K. Fritz. Mixed state discrimination: a DSM problem that won't go away? *Journal of Affective Disorders* 158, 8-10, 2014. doi:10.1016/j.jad.2014.01.008
- [82] J. L. Haight, S. B. Flagel. A potential role for the paraventricular nucleus of the thalamus in mediating individual variation in Pavlovian conditioned responses. *Frontiers in Behavioral Neuroscience* 8, 79, 2014. doi:10.3389/fnbeh.2014.00079
- [83] L. M. Williams, A. N. Goldstein-Piekarski, N. Chowdhry, K. A. Grisanzio, N. A. Haug, Z. Samara, A. Etkin, R. O'Hara, A. F. Schatzberg, T. Suppes, J. Yesavage. Developing a clinical translational neuroscience taxonomy for anxiety and mood

- disorder: protocol for the baseline-follow up Research domain criteria Anxiety and Depression ("RAD") project. *BMC Psychiatry* 16, 68, 2016. doi:10.1186/s12888-016-0771-3
- [84] J. A. Bernard, V. A. Mittal. Updating the research domain criteria: the utility of a motor dimension. *Psychological Medicine* 45(13), 2685-2659, 2015. doi:10.1017/S0033291715000872
- [85] A. von Eye. *Configural Frequency Analysis - Methods, Models, and Applications*. Mahwah, NJ: Lawrence Erlbaum, 2002.
- [86] G. A. Bogat, A. Von Eye, L. R. Bergman. *Person-oriented approaches*. 797- 845. In: D. Cichetti (Ed.). D. J. Cohen. *Developmental Psychopathology, Theory and Method*. 2nd Edition. ISBN: 978-0-471-23736-5. 1104 pages.
- [87] G. R. S. Treviranus. Not grasping patterns as you should? Person-centered Configural Frequency Analysis, its bio-psychological avoidance & the (high) costs of this. SSPE biannual meeting 2016. Poster session. University of Zurich. 24.06.2016. Online available from <https://www.researchgate.net/publication/304580615>
- [88] M. Ernst, D. S. Pine, M. Hardin. Triadic model of the neurobiology of motivated behavior in adolescence. *Psychological Medicine*, 36(3), 299–312, 2006. doi:10.1017/S0033291705005891
- [89] M. Ernst. The triadic model perspective for the study of adolescent motivated behavior. *Brain and Cognition* 89(0), 104-111, 2014. doi:10.1016/j.bandc.2014.01.006
- [90] A. S. Heller, A. O. Cohen, M. F. Dreyfuss, B. J. Casey. Changes in cortico-subcortical and subcortico-subcortical connectivity impact cognitive control to emotional cues across development. *Social Cognitive and Affective Neuroscience* 11(12), 1910-1918, 2016. doi:10.1093/scan/nsw097
- [91] L. H. Somerville, B. G. van den Bulk, A. C. Skwara. Response to: "the triadic model perspective for the study of adolescent motivated behavior". *Brain and Cognition* 89, 112-113, 2014. doi:10.1016/j.bandc.2014.01.003
- [92] X. Gonda, G. H. Vázquez. Theoretical and clinical overview of affective temperaments in mood disorders. *Psicodebate*, Vol. 14, N° 2, 2014. doi:10.1010.18682/pd.v14i2
- [93] B. J. Casey, A. Galván, L. H. Somerville. Beyond simple models of adolescence to an integrated circuit-based account: A commentary. *Developmental Cognitive Neuroscience* 17, 128-30, 2016. doi:10.1016/j.dcn.2015.12.00
- [94] Y. A. Kurnianingsih, O. A. Mullette-Gillman. Neural mechanisms of the transformation from objective value to subjective utility: converting from count to worth. *Frontiers in Neuroscience* 10, 507, 2016. doi:10.3389/fnins.2016.00507
- [95] M. T. Sherman, A. K. Seth, R. Kanai. Predictions shape confidence in right inferior frontal gyrus. *Journal of Neuroscience* 36(40),10323-10336, 2016. doi:10.1523/JNEUROSCI.1092-16.2016
- [96] Y. Li, G. Sescousse, C. Amiez, J. C. Dreher. Local morphology predicts functional organization of experienced value signals in the human orbitofrontal cortex. *Journal of Neuroscience* 35(4), 1648-58, 301, 2015. doi:10.1523/JNEUROSCI.3058-14.2015
- [97] S. M. Nicola. Reassessing wanting and liking in the study of mesolimbic influence on food intake. *American Journal Physiology Regulatory, Integrative, and Comparative Physiology*. 311, 5, R811-R840, 2016. <https://doi.org/10.1152/ajpregu.00234.2016>
- [98] L. H. Corbit, B. W. Balleine. *Learning and Motivational Processes Contributing to Pavlovian-Instrumental Transfer and Their Neural Bases: Dopamine and Beyond*. In: E. H. Simpson, P. D. Balsam (Eds.). *Behavioral Neuroscience of Motivation. Current Topics in Behavioral Neurosciences*. Vol. 27. (pp 259-289). Springer Int., Cham ZG, 2016. doi:10.1007/7854_2015_388
- [99] E. Cartoni, B. Balleine, G. Baldassarre. Appetitive Pavlovian-instrumental transfer: a review. *Neuroscience & Biobehavioral Reviews* 71, 829-848, 2016. doi:10.1016/j.neubiorev.2016.09.020
- [100] R. Lehner, J. H. Balsters, A. Herger, T. A. Hare, N. Wenderoth. Monetary, food, and social rewards induce similar Pavlovian-to-instrumental transfer effects. *Frontiers in Behavioral Neuroscience*. 10, 247, 2017. doi:10.3389/fnbeh.2016.0024
- [101] B. Studer, S. Knecht (Eds.). *Motivation: Theory, Neurobiology and Applications*. Progress in Brain Research 229, 2016. 1st Edition. Elsevier: London UK. 486 p. doi:10.1016/bs.pbr.2016.06.003
- [102] K. Lloyd, P. Dayan. Safety out of control: dopamine and defence. *Behavioral and Brain Functions* 12(1), 15, 2016. doi:10.1186/s12993-016-0099-7
- [103] T. E. Robinson, L. M. Yager, E. S. Cogan, B. T. Saunders. On the motivational properties of reward cues: Individual differences. *Neuropharmacology* 76(pt B), 450-459, 2014. doi:10.1016/j.neuropharm.2013.05.040
- [104] Y. Saga, Y. Hirata, D. Takahara, K. Inoue, S. Miyachi, A. Nambu, J. Tanji, M. Takada, E. Hoshi. Origins of multi-synaptic projections from the basal ganglia to rostrocaudally distinct sectors of the dorsal premotor area in macaques. *European Journal of Neuroscience* 33(2), 285-297, 2011. doi:10.1111/j.1460-9568.2010.07492.x
- [105] M. Guitart-Masip, E. Duzel, R. Dolan, P. Dayan. Action versus valence in decision making. *Trends in Cognitive Sciences* 18(4),194-202, 2014. doi:10.1016/j.tics.2014.01.00
- [106] B. T. Saunders, T. E. Robinson. Individual variation in resisting temptation: implications for addiction. *Neuroscience & Biobehavioral Reviews* 37(9 Pt A), 1955-1975, 2013. doi:10.1016/j.neubiorev.2013.02.008
- [107] S. Garofalo, G. di Pellegrino. Individual differences in the influence of task-irrelevant Pavlovian cues on human behavior. *Frontiers in Behavioral Neuroscience* 9, 163, 2015 doi:10.3389/fnbeh.2015.00163
- [108] K. S. Smith, A. M. Graybiel. Habit formation. *Dialogues in Clinical Neuroscience* 18(1), 33-43, 2016.
- [109] M. Aggarwal, J. R. Wickens. A role for phasic dopamine neuron firing in habit learning. *Neuron* 72(6), 892-894, 2011. doi:10.1016/j.neuron.2011.12.006
- [110] E. Schechtman, M. I. Noblejas, A. D. Mizrahi, O. Dauber, H. Bergman. Pallidal spiking activity reflects learning dynamics and predicts performance. *Proceedings of the National Academy of Sciences of the U. S. A.* 113(41), E6281-E6289, 2016. doi:10.1073/pnas.1612392113

- [111] L. P. Wang, F. Li, D. Wang, K. Xie, D. Wang, X. Shen, J. Z. Tsien. NMDA receptors in dopaminergic neurons are crucial for habit learning. *Neuron* 72(6), 1055-1066, 2011. doi:10.1016/j.neuron.2011.10.019
- [112] S. X. Luo, L. Timbang, J. I. Kim, Y. Shang, K. Sandoval, A. A. Tang, J. L. Whistler, J. B. Ding, E. J. Huang. TGF- β signaling in dopaminergic neurons regulates dendritic growth, excitatory-inhibitory synaptic balance, and reversal learning. *Cell Reports* 17(12), 3233-3245, 2016. doi:10.1016/j.celrep.2016.11.068
- [113] K. Jastrzębska, M. Walczak, P. E. Cieślak, Ł. Szumiec, M. Turbasa, D. Engblom, T. Błasiak, J. R. Parkitna. Loss of NMDA receptors in dopamine neurons leads to the development of affective disorder-like symptoms in mice. *Science Reports* 6, 37171, 2016. doi:10.1038/srep37171
- [114] L. Deserno, F. Schlagenhauf, A. Heinz. Striatal dopamine, reward, and decision making in schizophrenia. *Dialogues in Clinical Neuroscience* 18(1), 77-89, 2016.
- [115] L. Deserno, Q. J. Huys, R. Boehme, R. Buchert, H. J. Heinze, A. A. Grace, R. J. Dolan, A. Heinz, F. Schlagenhauf. Ventral striatal dopamine reflects behavioral and neural signatures of model-based control during sequential decision making. *Proceedings of the National Academy of Sciences of the U. S. A.* 112(5), 1595-1600, 2015. doi:10.1073/pnas.1417219112
- [116] A. Sinayev, E. Peters. Cognitive reflection vs. calculation in decision making. *Frontiers in Psychology*, 6, 532, 2015. doi:10.3389/fpsyg.2015.00532
- [117] A. Shenhav, M. M. Botvinick, J. D. Cohen. The expected value of control: an integrative theory of anterior cingulate cortex function. *Neuron* 79(2), 217-240, 2013. doi:10.1016/j.neuron.2013.07.007.
- [118] J. Scholl, N. Kolling, N. Nelissen, C. J. Stagg, C. J. Harmer, M. F. Rushworth. Excitation and inhibition in anterior cingulate predict use of past experiences. *Elife* 6, pii: e20365, 2017. doi:10.7554/eLife.20365
- [119] S. R. Heilbronner, B. Y. Hayden. Dorsal anterior cingulate cortex: A bottom-up view. *Annual Review of Neuroscience* 39, 149-170, 2016. doi:10.1146/annurev-neuro-070815-013952
- [120] M. Roy, D. Shohamy, T. D. Wager. Ventromedial prefrontal-subcortical systems and the generation of affective meaning. *Trends in Cognitive Sciences* 16(3), 147-156, 2012. doi:10.1016/j.tics.2012.01.005
- [121] S. M. Levens, J. T. Larsen, J. Bruss, D. Tranel, A. Bechara, B. A. Mellers. What might have been? The role of the ventromedial prefrontal cortex and lateral orbitofrontal cortex in counterfactual emotions and choice. *Neuropsychologia* 54, 77-86, 2014. doi:10.1016/j.neuropsychologia.2013.10.026
- [122] N. Cooper, D. S. Bassett, E. B. Falk. Coherent activity between brain regions that code for value is linked to the malleability of human behavior. *Scientific Reports* 7, 43250, 2017. doi:10.1038/srep43250
- [123] J. Brown, K. A. Martin, J. Dudman. Behavioral evidence for feedback gain control by the inhibitory microcircuit of the substantia nigra. *bioRxiv* preprint server. Accessible at: <http://biorxiv.org/content/early/2016/11/28/090209>. doi:10.1101/090209
- [124] S. R. Heilbronner, J. Rodriguez-Romaguera, G. J. Quirk, H. J. Groenewegen, S. N. Haber. Circuit-based corticostriatal homologies between rat and primate. *Biological Psychiatry* 80(7), 509-521, 2016. doi:10.1016/j.biopsych.2016.05.012
- [125] H. C. Lückmann, H. I. Jacobs, A. T. Sack. The cross-functional role of frontoparietal regions in cognition: internal attention as the overarching mechanism. *Progress in Neurobiology* 116, 66-86, 2014. doi:10.1016/j.pneurobio.2014.02.002
- [126] M.J.F. Robinson, A.M. Fischer, A. Ahuja, E.N. Lesser and H. Maniates. *Roles of "Wanting" and "Liking" in Motivating Behavior: Gambling, Food, and Drug Addictions*. In: E. H. Simpson, P. D. Balsam (Eds.). *The Behavioral Neuroscience of Motivation: An Overview of Concepts, Measures, and Translational Applications*. *Current Topics in Behavioral Neuroscience*. 2016;27: 105-136. doi:10.1007/7854_2015_402
- [127] K. Brans, P. Verduyn. Intensity and duration of negative emotions: comparing the role of appraisals and regulation strategies. *PLoS ONE* 9, e92410, 2014. doi:10.1371/journal.pone.0092410
- [128] S. M. Ng, W. K. Hou. S. M. Ng, W. K. Hou. Contentment duration mediates the associations between anxious attachment style and psychological distress. *Frontiers in Psychology*, 8, 258, 2017. <http://doi.org/10.3389/fpsyg.2017.00258>
- [129] D. Cutuli. Cognitive reappraisal and expressive suppression strategies role in the emotion regulation: an overview on their modulatory effects and neural correlates. *Frontiers in Systems Neuroscience* 8, 175, 2014. doi:10.3389/fnsys.2014.00175
- [130] P. Koval, S. Sütterlin, P. Kuppens. Emotional inertia is associated with lower well-being when controlling for differences in emotional context. *Frontiers in Psychology* 6, 1997, 2016. doi:10.3389/fpsyg.2015.01997
- [131] N. B. Kroemer, C. Burrasch, L. Hellrung. To work or not to work: Neural representation of cost and benefit of instrumental action. In: B. Studer and S. Knecht (Eds.). *Motivation — Theory, Neurobiology and Applications*. *Progress in Brain Research* 229, 125-157, 2016. doi:10.1016/bs.pbr.2016.06.009
- [132] S. P. Becker, D. R. Leopold, G. L. Burns, M. A. Jarrett, J. M. Langberg, S. A. Marshall, K. McBurnett, D. A. Waschbusch, E. G. Willcutt. The internal, external, and diagnostic validity of Sluggish Cognitive Tempo: A meta-analysis and critical review. *Journal of the American Academy of Child and Adolescent Psychiatry* 55(3), 163-78, 2016. doi:10.1016/j.jaac.2015.12.006
- [133] G. R. S. Treviranus. A synopsis of cyclical biaxial (affective) models: from Kraepelin to Askland's genetics and evolutionary neuroanatomy. Poster. 11th International Review Bipolar Disorders, 2010, Rome. *International Clinical Psychopharmacology* 26, e48, 2011. doi:10.1097/01.yic.0000405714.06608.40 Online available at <https://prezi.com/kosqybvz8aqx/>
- [134] E. Koutsoukos, E. Angelopoulos. Mood regulation in bipolar disorders viewed through the pendulum dynamics concept. *International Journal of Bipolar Disorder* 2(1), 9, 2014. doi:10.1186/s40345-014-0009-6

- [135] S. S. Tomkins, R. McCarter. What and where are the primary affects? Some evidence for a theory. *Perceptual Motor Skills* 18(S1), 119-158, 1964. In: E. V. Demos (Ed.) *Exploring affect. The selected writings of Silvan S. Tomkins*. Cambridge/Paris: Cambridge UP/Paris: Maison de l'Homme. 1995.
- [136] M. Lenzenweger, R. Depue. *A Neurobehavioral Model of Personality Disorders*. In D. Cicchetti (Ed.). *Developmental Psychopathology: Risk, Disorder, and Adaptation*. NY: Wiley-Interscience. 2015.
- [137] G. Paolone, C. C. Angelakos, P. J. Meyer, T. E. Robinson, M. Sarter. Cholinergic control over attention in rats prone to attribute incentive salience to reward cues. *Journal of Neuroscience*. 33(19), 8321-8335, 2013. doi:10.1523/JNEUROSCI.0709-13.2013
- [138] M. Krystel-Whittemore, K. N. Dileepan, J. G. Wood. Mast cell: A multi-functional master cell. *Frontiers in Immunology* 6, 620, 2015. doi:10.3389/fimmu.2015.00620
- [139] C. J. Fitzpatrick, S. A. Perrine, F. Ghodoussi, M. P. Galloway, J. D. Morrow. Sign-trackers have elevated myo-inositol in the nucleus accumbens and ventral hippocampus following Pavlovian conditioned approach. *Journal of Neurochemistry* 2016 Jan 4. [Epub ahead of print] doi:10.1111/jnc.13524.
- [140] S. Hendrix, K. Warnke, F. Siebenhaar, E. M. Peters, R. Nitsch, M. Maurer. The majority of brain mast cells in B10.PL mice is present in the hippocampal formation. *Neuroscience Letters* 392(3), 174-177, 2006. doi:10.1016/j.neulet.2005.09.029
- [141] B. P. Forester, C. T. Finn, Y. A. Berlow, M. Wardrop, P. F. Renshaw, C. M. Moore. Brain lithium, N-acetylaspartate and myo-inositol levels in older adults with bipolar disorder treated with lithium: a lithium-7 and proton magnetic resonance spectroscopy study. *Bipolar Disorders* 1(6), 691 – 700, 2008. doi:10.1111/j.1399-5618.2008.00627.x
- [142] J. W. Chang, H. Choi, S. L. Cotman, Y. K. Jung. Lithium rescues the impaired autophagy process in CbCln3-(Δ ex7/8/ Δ ex7/8) cerebellar cells and reduces neuronal vulnerability to cell death via IMPase inhibition. *Journal of Neurochemistry* 116(4), 659-668, 2011. doi:10.1111/j.1471-4159.2010.07158.x
- [143] C. J. Fitzpatrick, J. D. Morrow. Subanesthetic ketamine decreases the incentive-motivational value of reward-related cues. *Journal of Psychopharmacology* 31(1), 67-74, 2017. doi:10.1177/0269881116667709
- [144] W. Zhou, N. Wang, C. Yang, X. M. Li, Z. Q. Zhou, J. J. Yang. Ketamine-induced antidepressant effects are associated with AMPA receptors-mediated upregulation of mTOR and BDNF in rat hippocampus and prefrontal cortex. *European Psychiatry* 29(7), 419-423, 2014. doi:10.1016/j.eurpsy.2013.10.005
- [145] C. L. Lee, Y. H. Jiang, H. C. Kuo. Increased apoptosis and suburothelial inflammation in patients with ketamine-related cystitis: a comparison with non-ulcerative interstitial cystitis and controls. *BJU International* 112(8), 1156-1162, 2013. https://doi.org/10.1111/bju.12256
- [146] T. Fujimoto, T. Nishiyama, K. Hanaoka. Inhibitory effects of intravenous anesthetics on mast cell function. *Anesthesia & Analgesia* 101(4), 1054-1059, 2005. doi:10.1213/01.ane.0000166955.97368.80
- [147] A. Can, P. Zanos, R. Moaddel, H. J. Kang, K. S. Dossou, I. W. Wainer, J. F. Cheer, D. O. Frost, X. P. Huang, T. D. Gould. Effects of ketamine and ketamine metabolites on evoked striatal dopamine release, dopamine receptors, and monoamine transporters. *The Journal of Pharmacology and Experimental Therapeutics* 359(1), 159–170, 2016. http://doi.org/10.1124/jpet.116.235838
- [148] S. Fligel, T. E. Robinson. Neurobiological basis of individual variation in stimulus-reward learning. *Current Opinion in Behavioral Sciences* 13, 178-185, 2017. doi:10.1016/j.cobeha.2016.12.004
- [149] O. Hikosaka. The habenula: from stress evasion to value-based decision-making. *Nature Reviews in Neuroscience*, 11(7), 503–513, 2010. doi:10.1038/nrn2866
- [150] V. M. Namboodiri, J. Rodriguez-Romaguera, G. D. Stuber. The habenula. *Current Biology* 26(19), R873-R877, 2016. doi:10.1016/j.cub.2016.08.051
- [151] S. Torrisi, C. L. Nord, N. L. Balderston, J. P. Roiser, C. Grillon, M. Ernst. Resting state connectivity of the human habenula at ultra-high field. *Neuroimage* 147, 872-879, 2017. doi:10.1016/j.neuroimage.2016.10.034
- [152] S. K. Peters, K. Dunlop, J. Downar. Cortico-striatal-thalamic loop circuits of the salience network: A central pathway in psychiatric disease and treatment. *Frontiers in Systems Neuroscience*, 10, 104, 2016. doi:10.3389/fnsys.2016.00104
- [153] C. L. Danna, P. D. Shepard, G. I. Elmer. The habenula governs the attribution of incentive salience to reward predictive cues. *Frontiers in Human Neuroscience* 7, 781, 2013. doi:10.3389/fnhum.2013.00781
- [154] J. S. Ide, C. S. Li. Error-related functional connectivity of the habenula in humans. *Frontiers in Human Neuroscience* 5, 25, 2011. doi:10.3389/fnhum.2011.00025
- [155] O. Hikosaka, S. R. Sesack, L. Lecourtier, P. D. Shepard. Habenula: crossroad between the basal ganglia and the limbic system. *Journal of Neuroscience* 28(46), 11825-11829, 2008. doi:10.1523/JNEUROSCI.3463-08.2008
- [156] J. J. Balcita-Pedicino, N. Omelchenko, R. Bell, S. R. Sesack. The inhibitory influence of the lateral habenula on midbrain dopamine cells: ultrastructural evidence for indirect mediation via the rostromedial mesopontine tegmental nucleus. *Journal Comparative Neurology*. 519(6), 1143-1164, 2011. doi:10.1002/cne.22561
- [157] J. S. Wright, J. Panksepp. An evolutionary framework to understand foraging, wanting, and desire: The neuropsychology of the SEEKING system. *Neuropsychanalysis* 14(1), 5-75, 2012.
- [158] L. W. van Kerkhof, R. Damsteegt, V. Trezza, P. Voorn, L. J. Vanderschuren. Functional integrity of the habenula is necessary for social play behaviour in rats. *European Journal of Neuroscience* 38(10), 3465-3475, 2013. doi:10.1111/ejn.12353
- [159] R. P. Lawson, B. Seymour, E. Loh, A. Lutti, R. J. Dolan, P. Dayan, N. Weiskopf, J. P. Roiser. The habenula encodes negative motivational value associated with primary punishment in humans. *Proceedings of the National Academy of Science of the U.S.A.* 111(32), 11858-11863, 2014. doi:10.1073/pnas.1323586111

- [160] D. J. Furman, I. H. Gotlib. Habenula responses to potential and actual loss in major depression: preliminary evidence for lateralized dysfunction. *Social and Cognitive Affective Neuroscience* 11(5), 843-851, 2016. doi:10.1093/scan/nsw019
- [161] M. M. Mirrione, D. Schulz, K. A. Lapidus, S. Zhang, W. Goodman, F. A. Henn. Increased metabolic activity in the septum and habenula during stress is linked to subsequent expression of learned helplessness behavior. *Frontiers in Human Neuroscience* 8, 29, 2014. doi:10.3389/fnhum.2014.00029
- [162] K. Svenningsen, M. T. Venø, K. Henningsen, A. S. Mallien, L. Jensen, T. Christensen, J. Kjems, B. Vollmayr, O. Wiborg. MicroRNA profiling in the medial and lateral habenula of rats exposed to the learned helplessness paradigm: Candidate biomarkers for susceptibility and resilience to inescapable shock. *PLoS One*. 11(8), e0160318, 2016. doi:10.1371/journal.pone.0160318
- [163] Y. Ootsuka, M. Mohammed. Activation of the habenula complex evokes autonomic physiological responses similar to those associated with emotional stress. *Physiological Reports* 3(2), pii: e12297, 2015. doi:10.14814/phy2.12297
- [164] F. M. Schmidt, S. Schindler, M. Adamidis, M. Strauß, A. Tränkner, R. Trampel, M. Walter, U. Hegerl, R. Turner, S. Geyer, P. Schönknecht. Habenula volume increases with disease severity in unmedicated major depressive disorder as revealed by 7T MRI. *European Archives of Psychiatry and Clinical Neurosciences* 267(2), 107-115, 2017. doi:10.1007/s00406-016-0675-8
- [165] R. P. Lawson, C. L. Nord, B. Seymour, D. L. Thomas, P. Dayan, S. Pilling, J. P. Roiser. Disrupted habenula function in major depression. *Molecular Psychiatry* 22(2), 202-208, 2017. doi:10.1038/mp.2016.81
- [166] C. Winter, B. Vollmayr, A. Djodari-Irani, J. Klein, A. Sartorius. Pharmacological inhibition of the lateral habenula improves depressive-like behavior in an animal model of treatment resistant depression. *Behavioural Brain Research* 216(1), 463-465, 2011. doi:10.1016/j.bbr.2010.07.034
- [167] F. Cirulli, L. Pistillo, L. de Acetis, E. Alleva, L. Aloe. Increased number of mast cells in the central nervous system of adult male mice following chronic subordination stress. *Brain, Behavior, and Immunity* 12(2), 123-133, 1998. doi:10.1006/brbi.1998.0505
- [168] M. Wilhelm, B. King, A. J. Silverman, R. Silver. Gonadal steroids regulate the number and activational state of mast cells in the medial habenula. *Endocrinology* 141(3), 1178-1186, 2000. doi: 10.1210/endo.141.3.7352
- [169] Y. W. Hsu, S. D. Wang, S. Wang, G. Morton, H. A. Zariwala, H. O. de la Iglesia, E. E. Turner. Role of the dorsal medial habenula in the regulation of voluntary activity, motor function, hedonic state, and primary reinforcement. *Journal of Neuroscience* 34(34), 11366-11384, 2014. doi:10.1523/JNEUROSCI.1861-14.2014
- [170] D. S. Moura, S. Sultan, S. Georjin-Lavialle, N. Pillet, F. Montestruc, P. Gineste, S. Barete, G. Damaj, A. Moussy, O. Lortholary, O. Hermine. Depression in patients with mastocytosis: prevalence, features and effects of masitinib therapy. *PLoS One* 6(10), e26375, 2011. doi:10.1371/journal.pone.0026375
- [171] R. Musil. The man without Qualities. The 1st Vol. of 3: A Sort of Introduction - The Like of It Now Happens (I). New York, N.Y. Perigee Books, 1980. *Chapter 62*.
- [172] K. Jaspers. *General Psychopathology. Volume II. Transl. Hoening J, Hamilton MW. Baltimore: Johns Hopkins Univ Pr, 1997, p. 547*
- [173] E. J. Nunes, P. A. Randall, S. Podurgiel, M. Correa, J. D. Salamone. Nucleus accumbens neurotransmission and effort-related choice behavior in food motivation: effects of drugs acting on dopamine, adenosine, and muscarinic acetylcholine receptors. *Neuroscience and Biobehavioral Reviews* 37(9 Pt A), 2015-2025, 2013. doi:10.1016/j.neubiorev.2013.04.002
- [174] J. D. Salamone, M. Correa, E. J. Nunes, P. A. Randall, M. Pardo. The behavioral pharmacology of effort-related choice behavior: Dopamine, adenosine and beyond. *Journal of the Experimental Analysis of Behavior* 97(1), 125-146, 2012. doi:10.1901/jeab.2012.97-125
- [175] J. D. Salamone, S. E. Yohn, L. López-Cruz, N. San Miguel, M. Correa. Activational and effort-related aspects of motivation: neural mechanisms and implications for psychopathology. *Brain* 139(Pt5), 1325-1347, 2016. doi:10.1093/brain/aww050
- [176] M. P. Paulus, Q. J. Huys, T. V. Maia. Computational Psychiatry: From mechanistic insights to the development of new treatments. *Biological Psychiatry: Cognitive Neuroscience and Neuroimaging* 1(5), 382 - 385, 2016. doi:10.1016/j.bpsc.2016.08.001
- [177] E. Cartoni, B. Balleine, G. Baldassarre. Pavlovian-instrumental transfer: A review. *Neuroscience & Biobehavioral Reviews* 71, 829-848, 2016. doi:10.1016/j.neubiorev.2016.09.020
- [178] K. R. Ridderinkhof. Neurocognitive mechanisms of perception-action coordination: a review and theoretical integration. *Neuroscience & Biobehavioral Reviews* 46 Pt 1, 3-29, 2014. doi:10.1016/j.neubiorev.2014.05.008
- [179] N. K. Hansell, G. S. Halford, G. Andrews, D. H. Shum, S. E. Harris, G. Davies, S. Franic, (...), M. J. Wright. Genetic basis of a cognitive complexity metric. *PLoS One* 10(4), e0123886, 2015. doi:10.1371/journal.pone.0123886
- [180] W. R. Stauffer, A. Lak, S. Kobayashi, W. Schultz. Components and characteristics of the dopamine reward utility signal. *Journal of Comparative Neurology* 524(8), 1699-1711. doi:10.1002/cne.23880
- [181] F. Zhang, H. C. Tsai, R. D. Airan, G. D. Stuber, A. R. Adamantidis, L. de Lecea, A. Bonci, K. Deisseroth. Optogenetics in Freely Moving Mammals: Dopamine and Reward. *Cold Spring Harbor Protocols* 2015(8), 715-724, 2015. doi:10.1101/pdb.top086330
- [182] A. Kepecs, B. D. Mensh. Emotor control: computations underlying bodily resource allocation, emotions, and confidence. *Dialogues in Clinical Neuroscience*, 17(4), 391-401, 2015
- [183] Y. Li, C. Lindemann, M. J. Goddard, B. I. Hyland. Complex multiplexing of reward-cue- and licking-movement-related activity in single midline thalamus neurons. *Journal of Neuroscience* 36(12), 3567-3578, 2016. doi:10.1523/JNEUROSCI.1107-15.2016
- [184] C. Calanchini, C. Bassetti, M. R. Celio. Special edition on positive emotions. *Journal of Comparative Neurology* 524(8),

1529-1531, 2016. doi:10.1002/cne.23974 [Papers from the Congress “Brain Circuits of Positive Emotions”, 19-23 October 2014, Ascona, Switzerland, 19-23 October 2014. Accessible at: <http://www.unifr.ch/anatomy/emotions/>.

- [185] M. C. Antle, R. Silver. *Circadian Insights into Motivated Behavior*. In: E. H. Simpson, P. D. Balsam (Eds.). *The Behavioral Neuroscience of Motivation: An Overview of Concepts, Measures, and Translational Applications*. Current Topics in Behavioral Neuroscience 27, 105-136, 2016. doi:10.1007/7854_2015_384
- [186] T. Kawauchi, K. Ishimaru, Y. Nakamura, N. Nakano, M. Hara, H. Ogawa, K. Okumura, S. Shibata, A. Nakao. Clock-dependent temporal regulation of IL-33/ST2-mediated mast cell response. *Allergology International pii: S1323-8930(17)30008-4*, 2017. doi:10.1016/j.alit.2017.02.004
- [187] Y. Nakamura, K. Ishimaru, S. Shibata, A. Nakao. Regulation

of plasma histamine levels by the mast cell clock and its modulation by stress. *Science Reports* 39934, 2017. doi:10.1038/srep39934

- [188] M. Spencer, L. Yang, A. Adu, B. S. Finlin, B. Zhu, L. R. Shipp, N. Rasouli, C. A. Peterson, P. A. Kern. Pioglitazone treatment reduces adipose tissue inflammation through reduction of mast cell and macrophage number and by improving vascularity. *PLoS One* 9(7), e102190, 2014. doi:10.1371/journal.pone.0102190
- [189] G. Treviranus. Mast cell bio-economies: of barriers, routes, neuronal fates (2017), *Frontiers in Molecular Neuroscience*, submitted.
- [190] J. D. Salamone, S. E. Yohn, L. López-Cruz, N. San Miguel, M. Correa. Activational and effort-related aspects of motivation: neural mechanisms and implications for psychopathology. *Brain* 139(Pt5), 1325-1347, 2016. doi:10.1093/brain/aww050

i Abbreviation codes:

1. **Neuroanatomy, Neurochemistry:** ACC Anterior cingulate cortex; aINS Anterior insula; **CSTC:** Cortico-striato-thalamo-cortical circuits; Nac Nucleus accumbens; OFC Orbitofrontal cortex; vmPFC ventromedial Pre-Frontal Cortex. **PVT** paraventricular nucleus of thalamus. MC Mast cell; HA Histamine; NMDAR *N*-methyl-D-aspartate receptor for glutamate with e.g. NR2B-subunits; SNr substantia nigra (basal ganglia).
2. **Psychology / Ethology:** **UOs** Uncertainty-Oriented Individuals; **COs** Certainty-Oriented Individuals (Richard M. Sorrentino); **UO-versus-CO** Uncertainty versus Certainty “orientation”; **GTs** goal-trackers; **GT** goal-tracking; **STs** sign-trackers; **ST** sign-tracking. **CS** conditioned stimulus; **US** unconditioned stimulus. **MBL** Model-based learning; **MFL** Model-free learning.
3. **Neuro-economics:** **Effme** maximal expendable effort; **RPE** Reward Prediction Error; **PE (Effme)** RPE normalized to the present discount of previous maximum expendable effort.
4. **dyn4-TAM modeling.** *dyn4*TAM modified classic mixed bipolar disorder model; **T~**, *4DT* symbolic 4-dimensional cognitive processing; *4D*-Thought symbolic 4-dimensional cognitive processing; **A~**, *5+DA* intuitive 5- or higher-dimensional cognitive processing; *5+D*-Action intuitive 5- or higher-dimensional cognitive processing; **Mood** dichotomic negative or positive valence; **M~** dichotomic negative or positive valence; **T/t**, **A/a**, **M/m** dichotomic realizations of T, A, and M in triples. **P{A→T}** Perception, i.e. transitions from *5+D*-Action to *4D*-Thought; **I{T→A}** Intention, i.e. transitions from *4D*-Thought to *5+D*-Action; **AW**, **AWs** Appropriation Wave, Appropriation Waves; **SHO** Simple Harmonic Oscillator; **Worry-T~a~m~** Worry with much Thought, low Action and Mood.