1 Title:

2 Computational and neural mechanisms of affected beliefs

3 Authors:

- 4 Laura Müller-Pinzler¹, Nora Czekalla¹, Annalina V Mayer¹, Alexander Schröder¹, David S
- 5 Stolz¹, Frieder M Paulus¹, Sören Krach¹

6 Email:

- 7 Laura Müller-Pinzler* laura.muellerpinzler@uni-luebeck.de
- 8 Nora Czekalla <u>n.czekalla@uni-luebeck.de</u>
- 9 Annalina V Mayer <u>ann.mayer@uni-luebeck.de</u>
- 10 Alexander Schröder <u>a.schroeder@uni-luebeck.de</u>
- 11 David S Stolz <u>david.stolz@uni-luebeck.de</u>
- 12 Frieder M Paulus <u>frieder.paulus@uni-luebeck.de</u>
- 13 Sören Krach soeren.krach@uni-luebeck.de

14 Affiliations:

- 15 1: Department of Psychiatry and Psychotherapy, Social Neuroscience Lab, University of
- 16 Lübeck, Ratzeburger Allee 160, D-23538 Lübeck, Germany

17 Short Title:

18 *Corresponding Author:

- 19 Dr. Laura Müller-Pinzler
- 20 Department of Psychiatry and Psychotherapy, Social Neuroscience Lab

21 University of Lübeck, Ratzeburger Allee 160, D-23538 Lübeck, Germany

22 Phone: +49 45131017529

24

Abstract

The feedback people receive on their behavior shapes the process of belief formation 25 26 and self-efficacy in mastering a given task. The neural and computational mechanisms of how the subjective value of these beliefs and corresponding affect bias the learning process are yet 27 unclear. Here we investigate this question during learning of self-efficacy beliefs using fMRI, 28 29 pupillometry, computational modeling and individual differences in affective experience. 30 Biases in the formation of self-efficacy beliefs were associated with affect, pupil dilation and neural activity within the anterior insula, amygdala, VTA/SN, and mPFC. Specifically, neural 31 and pupil responses map the valence of the prediction errors in correspondence to the 32 experienced affect and learning bias people show during belief formation. Together with the 33 34 functional connectivity dynamics of the anterior insula within this network our results hint towards neural and computational mechanisms that integrate affect in the process of belief 35 formation. 36

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Introduction

40 Self-efficacy expectation is a person's subjective conviction that he or she can overcome challenging situations by own actions (Bandura, 1977). To successfully perform 41 goal directed actions, humans must learn from incoming information and thereby form beliefs 42 about the world and about themselves enmeshed in this world. According to economic theory, 43 learning should result in accurate beliefs representing an internal model of the world that is 44 suitable to inform decision making. Novel theoretical frameworks however emphasize that 45 besides its instrumentality (i.e. being accurate) the belief itself may carry intrinsic value 46 (Bromberg-Martin and Sharot, 2020) thereby shaping the learning process and how people 47 48 ultimately arrive at their beliefs (Sharot and Garrett, 2016). Here, affective states, e.g. 49 happiness about one's own good health prognosis, represent intrinsic values that individuals are inclined to optimize during belief formation (Bromberg-Martin and Sharot, 2020; Hughes 50 51 and Zaki, 2015). To prove this entanglement of affect and belief formation, we applied a learning task that induces affective reactions during the process of forming conceptually novel 52 beliefs about own abilities in mastering a task (Czekalla et al., 2021; Müller-Pinzler et al., 53 2019). We focused on the primary affective states elicited by self-related beliefs - the self-54 55 conscious emotions of embarrassment and pride – and their impact on the belief. By having 56 experimental control over failures and successes during the process of belief formation, we 57 were able to assess how experienced affect relates to computational mechanisms of belief formation and the underlying neural systems activity, explaining the shifts of preferences for 58 59 information of positive or negative valence during learning.

Affective states are considered to guide cognitive processing, representing embodied and experiential information about the positive or negative value of what people encounter (Frijda, 1987; Storbeck and Clore, 2008). These internal affective information are proposed to be integrated with external information shaping beliefs that, rather than being objective, are motivated and biased by subjective feelings towards the belief itself, which constitutes a

recursive influence of beliefs and affective states on each other (Bromberg-Martin and Sharot, 65 66 2020; Loewenstein, 2006). Previous studies support aspects of the Bromberg-Martin & Sharot framework by showing that internal beliefs and external feedback can elicit emotions like 67 happiness, pride, or embarrassment (Müller-Pinzler et al., 2015; Rutledge et al., 2014, 2016; 68 Stolz et al., 2020). Affective states also alter decision making (Charpentier et al., 2016a, 69 2016b; Stolz et al., 2020) and cognitive processes like situational judgments or learning styles 70 71 (Storbeck and Clore, 2008). Social anxiety, low self-esteem, or depression, which are likely associated with more negative affective reactions towards self-related beliefs, also bias belief 72 73 formation (Koban et al., 2017; Korn et al., 2014; Müller-Pinzler et al., 2019; Will et al., 2020) 74 supporting the overall rationale of the formation of "affected beliefs", that is, the notion that 75 beliefs are fundamentally shaped by affective experiences. The question however remains which neurophysiological mechanisms can explain how emotions brought up during learning 76 77 are associated with biases in belief formation.

Neuroscientific studies provided initial evidence that common brain areas map the 78 79 value of stimuli, actions, and their motivational relevance during social and non-social learning and decision making (Chib et al., 2009; Ruff and Fehr, 2014). Prediction errors, that is, the 80 81 mismatch of prior expectation and a situation's outcome, which are being minimized by 82 updating beliefs during learning, are generally processed in the dopaminergically innervated 83 ventral striatum, but also in the orbitofrontal cortex or the amygdala during learning (King-Casas et al., 2005; O'Doherty, 2004; Ruff and Fehr, 2014; Schultz et al., 1997). However, 84 85 more recent findings suggest that there are distinct and unique neural computations potentially reflecting the impact of motivational and emotional processes that are prominent during belief 86 87 formation. For example, studies could show that distinct value-related neural processes in subregions of the anterior cingulate cortex (ACC) are recruited depending on whether 88 89 information about oneself or another agent is processed (Lockwood and Wittmann, 2018; 90 Lockwood et al., 2016). In other studies, activity in the ventral striatum was modulated if the

91 social context changed from a private to a public situation, suggesting that the presence or 92 absence of others influenced sensitivity to the reward value of certain decisions (Izuma et al., 93 2010). Biases specific for self-related belief updating that are absent when people learn about 94 another person (Kuzmanovic et al., 2016), have been related to differences in tracking of 95 negative prediction errors (Sharot et al., 2011). Here, the ventromedial prefrontal cortex 96 (vmPFC) shows valence specific encoding of self-related information, which has been shown 97 to predict optimistic biases in updating (Kuzmanovic et al., 2016, 2018).

Affective states triggered after personal failures or successes are particularly important 98 when people acquire novel self-concepts (Hopkins et al., 2021) and develop an initial 99 100 understanding of themselves as being self-efficacious individuals in a novel task environment. 101 Central to the entanglement of affect and such self-efficacy beliefs is the assumption that people are highly motivated to perform well and maintain or even construct a positively shaped 102 103 self-image (Markus and Wurf, 1987; Sedikides and Gregg, 2008). Within this process, performance feedback elicits self-conscious emotions such as pride in case of success (Stolz et 104 al., 2020; Tangney et al., 2007; Williams and DeSteno, 2008), but also embarrassment if one 105 fails to achieve the expected outcome (Miller, 1996; Müller-Pinzler et al., 2015; Tangney et 106 107 al., 2007). It has already been shown that these emotions are not only a consequence of the 108 situation but also directly affect behavior. Pride experiences, associated with increased functional coupling between the ventral striatum and cortical midline structures (Stolz et al., 109 2020), thus functions as a motivator to continue one's effort (Williams and DeSteno, 2008). In 110 111 contrast, embarrassment experiences, as signified by increased functional connectivity between brain areas involved in "Theory of Mind" (Kanske et al., 2015) and arousal processing systems 112 within the ventral anterior insula (vAI) and amygdala (Müller-Pinzler et al., 2015), rather lead 113 individuals to stop the current behavior, withdraw, and appease others (Apsler, 1975; Feinberg 114 et al., 2012). For the process of belief formation, it has been argued that specifically the dorsal 115 116 mediofrontal cortex (dMFC), the ventral and dorsal anterior insula (vAI/ dAI), and amygdala,

brain areas involved during action monitoring as well as emotional processing, integrate affective states with outcome information (Koban and Pourtois, 2014). Therefore, among others, the AI has been regarded as an integrative hub for motivated cognition and emotional behavior (Koban and Pourtois, 2014; Wager and Feldman Barrett, 2017). Similarly, dopaminergic midbrain nuclei in the ventral tegmental area and substantia nigra (VTA/ SN) are associated with attention processes and at the same time events (i.e. reward cues) that are of motivational significance specifically during learning (Adcock et al., 2006; Schultz, 1998).

While current frameworks support the idea that intrinsic outcomes such as affective 124 125 states impact the process of belief formation (Bromberg-Martin and Sharot, 2020; Hughes and 126 Zaki, 2015), studies on this issue currently do not probe this framework as a whole. We aim 127 to bridge this gap by showing how emotional states relate to biases in self-related beliefs, that is, the formation of self-efficacy and how they shift preferences for information of positive or 128 129 negative valence. To do so, we tested the effects of individual differences in affective reactions to the task and learning behavior. Affective states were evoked during of learning self-efficacy 130 beliefs in a conceptually novel task environment. Using trial-by-trial updates of performance 131 expectations, we computed prediction error learning rates by fitting computational learning 132 models revealing valence specific learning biases. As predicted by current frameworks, 133 134 individual differences in the experience of the emotions embarrassment and pride were distinctively related to biases in learning of self-related beliefs. Biased belief updating and 135 affect were jointly related to neural processing of valence specific prediction errors in the AI, 136 137 amygdala, VTA/ SN and mPFC as well as pupillary reactivity in favor of information that is preferably used to update the belief. Increases in valence specific functional connectivity of the 138 dAI with the amygdala, VTA/ SN and mPFC support the idea of an integrative mechanism of 139 affective and attentional processes within the dAI. These findings provide insights into brain 140 networks involved in computational biases shaped by emotional experiences and coherently 141 142 support current theoretical frameworks integrating affective experiences in the process of belief

143 formation.

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Results

146 Experimental design

Of the participants in our overall sample more than half completed the task in the MRI 147 while additionally eve-tracking data were assessed during scanning, the other half completed 148 149 the task as a behavioral study (see methods section for details). In our self-efficacy learning experiment, the "learning of own performance" (LOOP) task (Müller-Pinzler et al., 2019), 150 participants were repeatedly confronted with manipulated feedback on their own and another 151 152 person's cognitive estimation ability. In different domains (e.g., estimating the weight of 153 animals) participants were led to form novel beliefs about their performance-related selfefficacy. We invited each participant together with a confederate to take part in the "cognitive 154 155 estimation" experiment in our neuroimaging lab. The participant performed the task in the MRI scanner, while the confederate (introduced as another participant) was located in an 156 adjacent room to simultaneously perform the task there. During the task, participants were 157 asked to estimate specific characteristics of different properties (e.g. heights of buildings or 158 weights of animals). After each trial, they received a manipulated performance feedback for 159 160 their last estimation (see **Figure 1a**). During the entire experiment participants took turns in performing the estimation task themselves ("Self" condition) or observing the other 161 participant performing ("Other" condition). At the beginning of each trial participants were 162 163 requested to rate either their own or the other person's expected performance for the upcoming trial, enabling us to examine the process of self- and other-related belief formation. The design 164 of the LOOP task provided a High Ability and a Low Ability condition which resulted in 165 overall four feedback conditions: Agent condition (Self vs Other) x Ability condition (High 166 Ability vs Low Ability; see Figure 1b and methods for a detailed description of the task). In 167 previous studies we showed that over time participants adapted their expected performance 168

169 ratings according to the feedback allowing for an assessment of valence specific self- and

170 other-related learning processes (Czekalla et al., 2021; Müller-Pinzler et al., 2019).

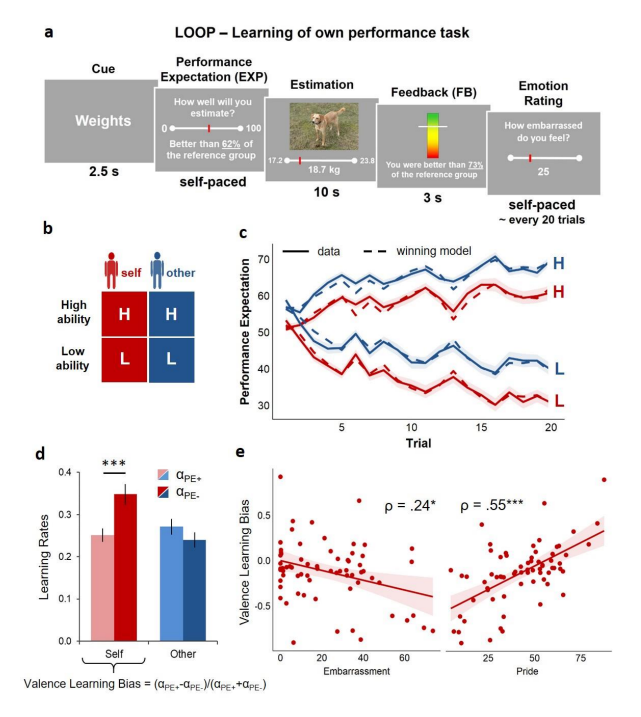




Figure 1. Trial sequence and timing, experimental conditions, modeling of learning behavior, 172 learning rates and their association with self-conscious emotions. a) A cue in the beginning of 173 each trial indicated the following estimation category as well as the agent who's turn it was. 174 After providing their performance expectation ratings (EXP) participants received an 175 estimation question, followed by the corresponding performance feedback. After 176 approximately every 20 trials participants were asked to rate their current emotional state 177 (pride, embarrassment, happiness, stress/ arousal). b) The LOOP task contained two 178 experimental factors, Ability level (High ability vs Low ability) and Agent (Self vs Other), 179 resulting in four feedback learning conditions that are distinguishable via different estimation 180

question types (e.g. estimation of weights of animals or heights of buildings). c) Predicted and 181 actual performance expectation ratings across time. The behavioral data indicate that 182 participants adapted their performance expectation ratings (solid lines) to the provided 183 feedback, thus learning about their allegedly distinct performance levels in the two ability 184 conditions for themselves and the other person. The winning valence specific learning model 185 captured the participants' behavior as indicated by a close match of actual performance 186 expectations with the predicted data (dashed lines). Shaded areas represent the standard errors 187 for the actual performance expectations for each trial. d) Learning rates derived from the 188 winning Valence Model indicate that there was a bias towards increased updating in response 189 to negative prediction errors (α_{PE-}) in contrast to positive prediction errors (α_{PE+}) for self-190 related learning. Bars represent mean learning rates, error bars depict +/- one standard error; 191 *** = p < .001, indicates a significant negativity bias during self-related learning. e) 192 Correlation plots and spearman correlations of self-related Valence Learning Bias and 193 embarrassment as well as pride experience during the experiment. * = p < .05, *** = p < .001. 194 195

196 Model free behavioral analyses reveal more negative self-evaluation

197 We first performed a model free analysis to capture the basic effects we observed in 198 our behavioral data. Analyses of behavioral data and learning rates are based on the combined fMRI (n=39) and behavioral sample (n=30; overall N=69). The Trial x Ability condition x 199 Agent condition x Group ANOVA revealed a significant main effect of Ability condition 200 $(F_{(1,67)}=175.51, p<.001)$ and interaction of Trial x Ability condition $(F_{(19,1273)}=108.87, p<.001)$ 201 indicating that participants adapted their performance expectation ratings over time according 202 to the feedback provided in each Ability condition (see Figure 1c). There was a significant 203 main effect of Agent condition ($F_{(1,67)}$ =44.70, p=.001) which indicated that participants 204 205 evaluated their own performance more negatively than the other's performance. There was no significant interaction of Agent condition x Ability condition ($F_{(1.67)}=0.67$, p=.415). The three-206 way interaction of Trial x Agent condition x Ability condition ($F_{(19,1273)}=1.60$, p=.047) showed 207 208 a significant effect hinting towards differential learning patterns between the Ability conditions for Self vs Other. There was a significant main effect of Group ($F_{(1,67)}$ =4.32, 209 p=.041) indicating slightly higher ratings in the fMRI sample, but there was no interaction of 210 Group with any of the effects reported above (p>.174). 211

212 Selection of computational models for self-related learning

In a next step, we modeled the participants' behavior by means of learning rates.

Therefore, we used the trial-specific expectation ratings for Self and Other to model prediction 214 215 error (PE) update learning and assess differences between updating behavior for information of positive vs negative valence. Our model space contained different models allowing us to 216 assess the importance of valence specific learning rates in contrast to unbiased learning 217 between conditions and agents (Figure S1). In line with our previous studies, an extended 218 version of the Valence Model, including separate learning rates for positive and negative PEs 219 220 for Self vs Other was the winning model (Model 8; for a more detailed description of this model and the whole model space see methods section). It received the highest sum PSIS-221 LOO score (approximate leave-one-out cross-validation (LOO) using Pareto-smoothed 222 223 importance sampling (PSIS)) (Vehtari et al., 2016) out of all models (for all PSIS-LOO scores 224 see **Supplementary Table S1**). In addition, Bayesian model selection (BMS) resulted in a protected exceedance probability of *pxp*>.999 for this model and a Bayesian Omnibus Risk 225 226 of BOR<.001. Thus, the extended Valence Model was selected for all further analyses of learning parameters allowing for a comparison of valence specific learning rates. The time 227 courses of performance expectation ratings predicted by our winning model successfully 228 captured trial-by-trial changes in the actual expectations due to PE updates within each of the 229 ability conditions at the individual subject level ($R^2=0.46\pm0.28$ [$M\pm SD$]) supporting the 230 231 validity of the model in describing the subjects' learning behavior. In addition to revealing PE valence specific learning, which could not directly be assessed via model free behavioral 232 analyses, posterior predictive checks also confirmed that the winning model captured the core 233 234 effects in our model free analysis (see Supplementary Results and Figure 1c).

235 Replication of the negativity bias for self-related learning

Participants showed higher learning rates when learning about themselves compared to learning about another person (main effect of Agent: $F_{(1,67)}$ =5.77, p=.019). There was also a main effect of PE valence [pos| neg] ($F_{(1,67)}$ =5.22, p=.025; indicating the categorical comparison of learning rates for positive vs negative PEs) and a significant interaction of Agent x PE valence [pos| neg] ($F_{(1,67)}$ =21.47, p<.001) which replicates earlier findings of a bias towards more negative updating when learning about one's own performance ($t_{(68)}$ =-4.85, p<.001, $M\alpha_{PE+Self}$ =0.25, SD=0.13; $M\alpha_{PE-Self}$ =0.35, SD=0.20) (Müller-Pinzler et al., 2019). Learning about the other person's performance did not show a significant bias towards more negative updating ($t_{(68)}$ =1.53, p=.128; $M\alpha_{PE+Other}$ =0.27, SD=0.16; $M\alpha_{PE-Other}$ =0.24, SD=0.15;

see **Figure 1d**). There was no significant main effect or interaction for Group (p>.097).

246 Associations of self-related learning with self-conscious emotions.

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Individual differences in the overall experience of embarrassment and pride during the 247 task were used as between-subject measures to quantify associations between learning 248 249 behavior and affect. Embarrassment and pride ratings were only weakly correlated ($\rho_{(68)}$ =-.10, p=.436), indicating that the experience of embarrassment and pride during the task represent 250 two rather independent affective components with respect to the self-related feedback. The 251 252 bias in self-related learning (Valence Learning Bias= $(\alpha_{Self/PE+} - \alpha_{Self/PE+})/(\alpha_{Self/PE+} + \alpha_{Self/PE-}))$ (Müller-Pinzler et al., 2019; Niv et al., 2012; Palminteri et al., 2017) was negatively linked to 253 the reported experience of embarrassment during the task ($\rho_{(68)}$ =-.24, p=.043; fMRI 254 subsample: $\rho_{(38)}$ =-.44, p=.005), that is, more negative updating behavior was associated with 255 increased embarrassment ratings. In contrast, the Valence Learning Bias was positively linked 256 257 to the emotion of pride ($\rho_{(68)}$ =.55, p<.001; fMRI subsample: $\rho_{(38)}$ =.47, p=.002). A regression predicting the Valence Learning Bias with both affect ratings simultaneously showed 258 independent effects of pride (β =0.56 $t_{(66)}$ =5.81, p<.001) and embarrassment (β =-0.22, $t_{(66)}$ =-259 2.30, p=.025; $R^2=.41$, $F_{(1,66)}=22.90$, p<.001). This indicates that the experience of self-260 conscious emotions during successful and unsuccessful performances are tied to the way 261 people updated their self-efficacy beliefs (see Figure 1e). Further, the way participants 262 processed the performance feedback in order to update their self-related ability beliefs was 263 associated with their self-esteem. People with higher self-esteem showed more positive 264 updating, $\rho_{(68)}$ =.33, p=.006 (fMRI subsample: $\rho_{(38)}$ =.35, p=.030), which strengthens the 265

assumption that prior beliefs about the self have a direct impact on how individuals learn novel

information about new abilities (Müller-Pinzler et al., 2019; Rouault et al., 2019).

268 Pupil dilation slopes are associated with surprise and valence of prediction errors in line

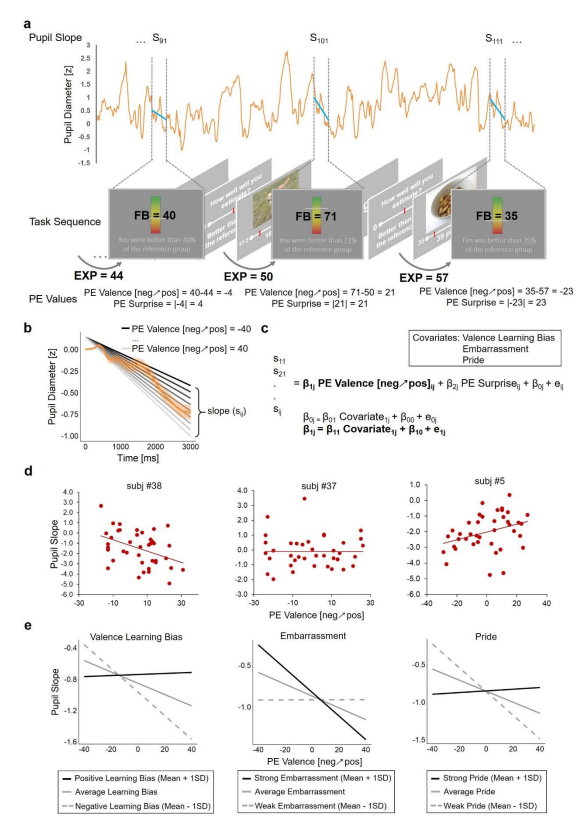
269 with a negative learning bias

Prior research has successfully linked changes in pupil diameter to surprise, PEs and 270 learning (Koenig et al., 2018; Preuschoff, 2011) as well as emotional experiences and arousal 271 (Bradley et al., 2008; Müller-Pinzler et al., 2015). To corroborate our assumption that changes 272 in pupil diameter, as indicated by the slope of the change in pupil size during self-related 273 feedback presentation, reflect increased arousal or attention associated with greater PEs we 274 275 regressed trial-by-trial variability in the pupil slope on PE surprise (continuous effect of 276 unsigned PEs) and PE valence $[neg \land pos]$ (continuous effect of signed PEs; see Figure 2a) (Rouhani and Niv, 2021). The linear mixed model showed a significant positive effect for PE 277 surprise $(t_{(1406)}=2.20, p=.028)$ and a negative effect for PE valence $[neg \nearrow pos]$ $(t_{(31.2)}=-2.50, p=.028)$ 278 p=.018; see **Figure 2b**). First, we observed an effect of PE surprise in the sense that the more 279 280 surprising the feedback was with respect to trial-by-trial prior expectations the more pupil dilation increased, in line with previous findings on pupil dilation in response to surprising 281 events (Preuschoff, 2011). Second, the results indicate that pupil dilation was greater with 282 decreasing PE values, linking negative PEs rather than positive PEs to greater dilation, 283 potentially indicating increased arousal and attention towards negative PEs in line with the 284 285 negativity bias we found in learning rates.

Pupil dilation response to prediction error valence is associated with affect and learning bias

It has been suggested that pupil dilation not only reflects differences between stimuli but similarly individual biases during decision making (see **Figure 2d** for examples of individual differences; de Gee, Knapen, & Donner, 2014). To corroborate our assumption that pupil slopes should reflect increased arousal or attention associated with PEs that are

preferably used for updating by each individual (i.e. individuals preferably learning from 292 negative vs positive PEs, experiencing more embarrassment/ less pride) we thus introduced 293 individual differences in learning and self-conscious emotions as between-subject covariates 294 into the linear mixed models assessing trial-by-trial pupil slopes (see Figure 2c). These 295 analyses demonstrated that individuals who experienced more embarrassment showed 296 297 stronger pupil dilations for negative compared to positive PEs while in individuals with lower 298 embarrassment pupil slopes did not differ between positive and negative PEs (significant interaction of embarrassment and PE valence [neg/pos]: $t_{(31,8)}$ =-2.57, p=.015; no main effect 299 for embarrassment: $t_{(34)}$ =-0.42, p=.680; see Figure 2e). Effects were reversed when pride 300 ratings were included in the model instead of embarrassment ratings (interaction pride and PE 301 valence $[neg pos] t_{(32.8)} = 3.14$, p = .004; main effect of pride: $t_{(34.1)} = 0.04$, p = .971). The Valence 302 Learning Bias modulated the relationship between PE valence [neg/pos] and pupil slopes in 303 the same way (interaction Valence Learning Bias and PE valence $[neg \nearrow pos] t_{(31,3)}=2.96$, 304 305 p=.006; main effect of Valence Learning Bias: $t_{(34,3)}=1.05$, p=.300) indicating that people with a more negative Valence Learning Bias had greater pupil dilation for negative PEs whereas 306 people with no or positive bias, showed less differentiation in the pupil dilation in response 307 to the valence of the PE. 308



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Figure 2. Association of pupil slopes with prediction error (PE) valence and individual pupil response differences explained by differences in Valence Learning Bias, embarrassment and pride experience. **a**) Exemplary pupil diameter trace over three trials for one subject (orange line) and trial specific fitted linear slopes (blue lines) for the feedback phase of each trial. PE values are calculated with the participant's current performance expectation (EXP) and the following feedback value (FB) and PE valence [neg \nearrow pos] represents the signed PE while PE

surprise represents the unsigned PE. b) More negative PEs are associated with greater pupil 317 slopes compared to positive PEs. The average pupil diameter trace during feedback is depicted 318 319 in orange, shaded area represents +/- one standard error. Pupil slopes for the different levels of PEs (from black = negative to grey = positive) were predicted by the multi-level model 320 containing PE valence $[neg \land pos]$ and PE surprise as predictors as described in the methods 321 section and depicted in c. c) Description of the multi-level model assessing the association of 322 PE valence $[neg \land pos]$ and PE surprise with within-subject trial-by-trial pupil slopes and the 323 impact of Valence Learning Bias, embarrassment and pride experience as between-subject 324 second level covariates explaining differences in the associations on the within-subject level 325 326 (cross-level-interaction; indicated in bold). d) Three exemplary scatter plots for three different subjects show the association of pupil slopes with PE valence $[neg \land pos]$ and illustrate the 327 variance between subjects. Subj#38 shows increased pupil slopes for negative PEs in line with 328 the group level effect. Subj#37 shows no association and subj#5 shows increased pupil slopes 329 for positive PEs. e) Illustration of the impact of the three between-subject covariates, Valence 330 Learning Bias (left), embarrassment (middle) and pride experience (right) explaining 331 differences in the associations of PE valence [neg/ pos] and pupil slope. The plots show the 332 333 data as predicted by the multi-level models for the mean covariate as well as the mean covariate +/-1 standard deviation (SD). The results show that, for example, individuals with 334 a more negative Valence Learning Bias (gray dashed line, left), increased experience of 335 336 embarrassment (black line, middle) and decreased experience of pride (gray dashed line, right) showed a stronger pupil response to negative vs positive PEs. 337 338

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9 Neural activations associated with feedback processing indicate a specific role of

340 feedback valence during self-related learning

In a next step, we examined the brain processes that underlie how people form self-341 and other-related ability beliefs. Therefore, we first compared neural activation during 342 feedback processing as measured with fMRI. We found that the bilateral insula, anterior 343 cingulate cortex, and thalamus (amongst others, see Figure 3a and Supplementary Table 344 345 S2) were activated significantly stronger if feedback was related to one's own performance as compared to the other person (i.e. Agent effect). This finding of heightened activity in brain 346 regions that have been linked to arousal but also self-agency potentially reflects a difference 347 in the subjective salience of self- vs other-related information (Craig, 2009; Späti et al., 2014; 348 Sperduti et al., 2011). Feedback for the Other as compared to the Self resulted in stronger 349 activation of the left and right middle temporal gyrus and precuneus/ middle cingulate gyrus 350 351 (Supplementary Table S2).

352

Second, we compared self-related positive vs negative feedback to examine how the

valence of information affected neural processing (categorial PE valence [posl neg] effect). 353 354 We found significantly stronger activations of the left and right nucleus accumbens/ ventral striatum (NAcc/VS), bilateral angular gyrus, medial prefrontal cortex (mPFC), and precuneus/ 355 posterior cingulate cortex (PCC) for positive vs. negative PE valence [pos| neg] (see 356 Supplementary Table S2). This valence effect was unique for processing of self-related 357 information and absent when feedback was related to the other person's performance (no 358 significant clusters for the PE valence [pos] neg] effect for Other; p < .001). The opposite 359 contrast, negative vs positive PE valence [pos| neg], yielded no significant activations, neither 360 for self-related nor for other-related information. When testing the interaction of Agent x PE 361 362 valence [pos| neg] we found increased activation for self-related positive vs negative feedback ([Self positive PE > Self negative PE] > [Other positive PE > Other negative PE]) in the 363 angular gyrus (see Supplementary Table S2), and on a more lenient threshold also the 364 bilateral NAcc/VS, the precuneus/ PCC, and precentral gyrus (cluster-wise FWE-corrected 365 with p < .05 at a cluster forming threshold of p < .001; see Figure 3a/ b and Supplementary 366 Table S3). 367

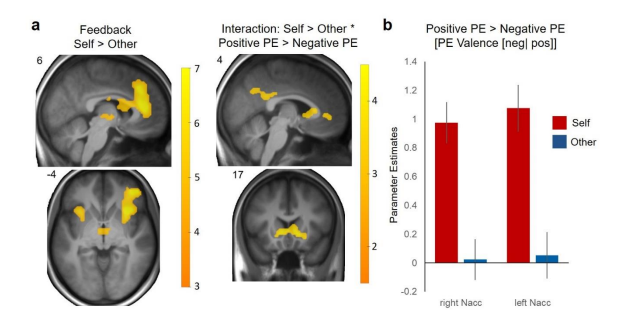




Figure 3. Neural activations associated with feedback processing. **a**) Self-related feedback vs other-related feedback (left) was associated with an increased activation of the mPFC/ ACC,

bilateral anterior insula and thalamus among other regions (p < .05, FWE-corrected). The 372 interaction of Agent and PE valence [pos| neg] (([Self positive PE > Self negative PE] > [Other 373 positive PE > Other negative PE]; right) resulted in activation of the angular gyrus, the 374 bilateral NAcc/VS, the precuneus/ posterior cingulate cortex, and precentral gyrus (cluster-375 wise FWE-corrected with p < .05 at a cluster forming threshold of p < .001). b) Parameter 376 377 estimates for the differences between feedback for positive vs negative PEs derived from the peak voxels of the interaction effect depict the interaction in the left NAcc/ VS [x, y, z: -9 20 378 -1] and right NAcc/ VS [x, y, z: 12 20 -1]. Self-related feedback resulted in a valence specific 379 activation while other-related feedback did not. 380

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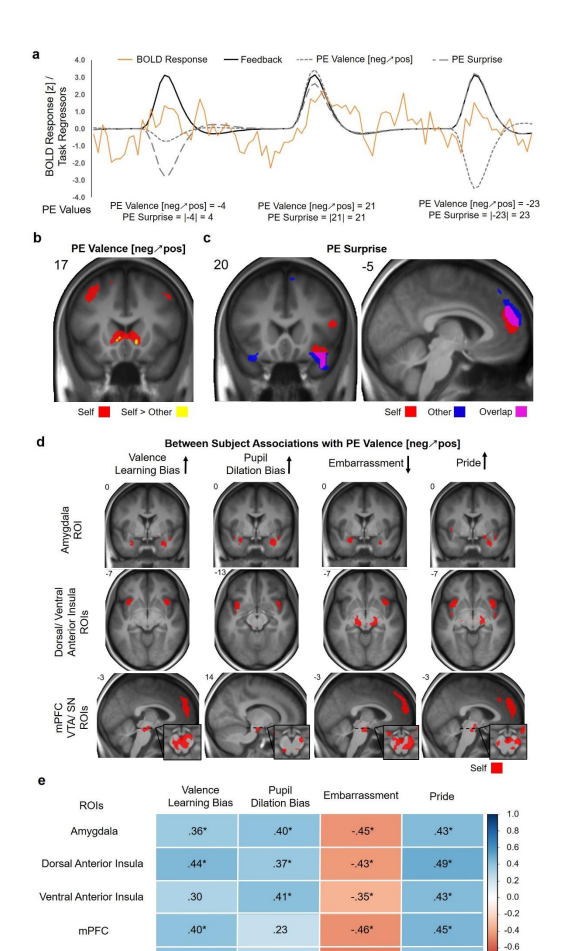
Common neural activations associated with prediction error surprise and distinct

383 activations for self-related prediction error valence

384 In a second step we assessed the effects of continuous trial-by-trial PE surprise and PE valence [neg/ pos] as parametric weights to assess neural aspects of learning more 385 specifically (see Figure 4a). Increased PE surprise was associated with greater activation of 386 the mPFC for Self and Other as well as clusters in the bilateral insula/ temporal pole/ frontal 387 orbital gyrus on a more lenient threshold (cluster-wise FWE-corrected with p < .05 at a cluster 388 389 forming threshold of p < .001; see Figure 4c and Supplementary Table S4). There was no significant difference between Self and Other (p < .001), potentially indicating that there is a 390 common process of error tracking mapped within the same brain regions independent of the 391 agent. 392

Assessing PE valence [neg/pos] revealed a distinct pattern for self- and other-related 393 394 learning. Self-related PE valence [neg / pos] was positively associated with increased activation of the NAcc/VS, mPFC, bilateral angular gyrus/ superior parietal lobule/ lateral 395 396 occipital gyrus and precentral gyrus, showing stronger activation for positive vs negative PEs (Figure 4b and Supplementary Table S5). There was no effect for other-related PE valence 397 $[neg \land pos]$ and a direct comparison of self vs other-related PE valence $[neg \land pos]$ effects 398 showed stronger associations in the NAcc/VS for Self (right: x, y, z: 12, 17, -4, $t_{(38)} = 5.23$; k 399 = 2; left: x, y, z: -9, 26, -1, $t_{(38)}$ = 5.77, k = 19; all coordinates in MNI space), supporting that 400 the valence of the feedback has a specific value when feedback refers to the self as compared 401

to another person. Although behavioral data and learning rates clearly stress the importance 402 403 of negative over positive PEs, there were no significant negative associations with PE valence [neg \nearrow pos] in the neural data (p < .001). To test if the activations associated with self-related 404 PE valence $[neg \nearrow pos]$ were actually related to PEs and not only to the feedback value alone 405 we ran an additional control analysis including parametric weights for trial-by-trial feedback 406 and performance expectation ratings instead of PE values (Zhang et al., 2020). This model 407 replicated the findings showing positive associations within the reported brain regions, 408 including the NAcc/ VS, with trial-by-trial self-related feedback values and negative 409 410 associations with prior expectations as it would be expected for PE-related effects (Supplementary Table S6). 411



412

VTA/ SN

.40*

-.51*

.27

-0.8

-1.0

.40*

Figure 4. Common neural activations associated with prediction error (PE) surprise, distinct 413 activations for self-related PE valence $[neg \land pos]$ and individual response differences to PE 414 valence [neg/ pos] explained by differences in Valence Learning Bias, embarrassment and 415 pride experience, and pupil dilation. a) Exemplary BOLD response over three trials for one 416 subject (orange line) and regressors for the feedback phase of each trial (back line; originally 417 separate regressors for self- and other-related feedback are here combined for displaying 418 419 purposes). PE valence [neg≁ pos] (small dashed) and PE surprise (large dashed) are added as parametric modulators in addition to the feedback regressors. PE values are calculated as 420 shown in **Figure 2**. b) PE valence $[neg \land pos]$ was associated with increased activation of the 421 422 NAcc/VS, mPFC, bilateral angular gyrus/ superior parietal lobule/ lateral occipital gyrus and precentral gyrus for Self. Activation of the NAcc/VS was stronger for Self vs Other. c) PE 423 surprise was associated with activation of the mPFC and the bilateral insula/ temporal pole/ 424 425 frontal orbital gyrus for Self and Other. d) Neural tracking of self-related PE valence $[neg \land$ pos] in the predefined regions of interest (ROIs: amygdala, ventral and dorsal anterior insula, 426 mPFC, VTA/SN) was modulated by between-subject variables. Specifically, individuals with 427 a more negative Valence Learning bias showed relatively stronger neural responses to 428 429 negative as compared to positive PEs, which was similar for individuals with stronger pupil dilation response to negative PEs, or increased ratings of embarrassment. In contrast, 430 individuals with higher pride ratings showed relatively stronger neural responses to positive 431 as compared to negative PEs. Black arrows indicate the direction the covariates are coded in 432 the analyses; embarrassment is coded negatively as high embarrassment was supposed to be 433 associated with increased activity for negative rather than positive PE valence [neg/ pos]. 434 Clusters refers to p < .005, uncorrected for displaying purposes; see **Supplementary Table** 435 436 S7 for FWE-corrected statistics. e) Pearson correlations for parameter estimates derived from the whole areas of our predefined ROIs with the Valence Learning Bias, Pupil Dilation Bias, 437 embarrassment and pride are color coded. * = p < .05, FDR corrected. 438 439

440 Neural activity in response to self-related prediction error valence is associated with

441 affect, learning bias, and pupil dilation

442 To assess how biases in learning as well as affective experience and pupil dilation were associated with valence specific PE processing on the single trial level, multiple general linear 443 444 models (GLMs) were implemented. These included the Valence Learning Bias, self-conscious emotions and a score representing a valence bias for pupil dilation responses towards positive 445 vs negative PEs (Pupil Dilation Bias = PupilSlope_{Self/PE+} - PupilSlope_{Self/PE+}) as between 446 447 subject covariates for PE valence [neg/pos] tracking. Analyses within our predefined regions of interest (ROIs) revealed that the more negative the Valence Learning Bias was, the more 448 449 neural activity increased in response to negative vs positive PEs in the bilateral dAI, vAI, amygdala, mPFC, and VTA/ SN (all results are p < .05 FWE-corrected within ROIs; see Figure 450 451 4d and Supplementary Table S7). Overall higher experience of embarrassment showed

similar associations with increased PE tracking for negative vs positive PEs in the right dAI, 452 453 bilateral amygdala, and VTA/ SN. Trendwise effects for embarrassment were found in the left 454 dAI, bilateral vAI, and mPFC. In line with this, lower experience of pride showed the same association in the dAI and vAI, amygdala, VTA/ SN and mPFC. Additional analyses revealed, 455 that effects for embarrassment and pride were mainly independent (see Supplementary 456 **Results**). Similarly, the more negative the Pupil Dilation Bias was, the stronger the increase 457 in activation of the dAI and vAI, amygdala and VTA/ SN towards negative vs positive PEs. 458 Thus, the greater the response of this neural system to PEs with negative valence the greater 459 was the preference for negative information during learning as well as the negativity of the 460 461 affective experience. This gained multi-modal support by similar associations of the Valence 462 Learning bias and affect with the pupil dilation response, which reflects the activity of this underlying neural system. In contrast, participants which showed a greater response of this 463 464 neural system towards positive PEs also had a preference for positive information during learning and reported more positive affect. 465

466 Functional connectivity of the dorsal anterior insula depends on prediction error valence

467 in line with the negativity bias

Functional connectivity dynamics of the left and right dAI were assessed as these were 468 469 activated during feedback processing for self- and other-related feedback, independent of Agent and PE valence. Here, we tested if the connectivity of the dAI differed depending on 470 the level of the valence of the PE, by using psychophysiological interaction (PPI) analyses. 471 472 We calculated the interaction of the continuous PE valence [neg 2 pos] and the timeseries extracted from the left and right dAI seed regions separately for Self and Other on the first 473 level and the two agents were contrasted against each other on the second level GLM as we 474 were specifically interested in connectivity dynamics that could reflect the differential 475 learning from negative PEs when processing self-relevant information. Contrasting the PPI 476 effects for PE valence [neg/pos] between the Self and Other demonstrated that during self-477

related learning, functional connectivity dynamics of the right dAI with the bilateral 478 479 amygdala, mPFC and VTA/ SN (p<.05, FWE-corrected within ROIs) more strongly aligned with the negativity of the PEs. The left dAI showed a weaker but similar spatial distribution 480 with significant differences between self- and other-related PE valence [neg/pos] for the left 481 amygdala and VTA/ SN (p <. 05, FWE corrected, see Figure 5a/b and Supplementary Table 482 S8). Thus, those brain regions that preferably tracked PEs of negative valence in individuals 483 with increased negative affect and learning biases also showed connectivity dynamics with 484 the dAI in a similar direction during self-related learning. Individuals who showed more 485 pronounced differences in functional connectivity, that is, stronger functional connectivity for 486 487 negative PEs during Self>Other, also showed a more negative Valence Learning Bias, 488 although this pattern was not fully consistent across all ROIs (see Figure 5c).

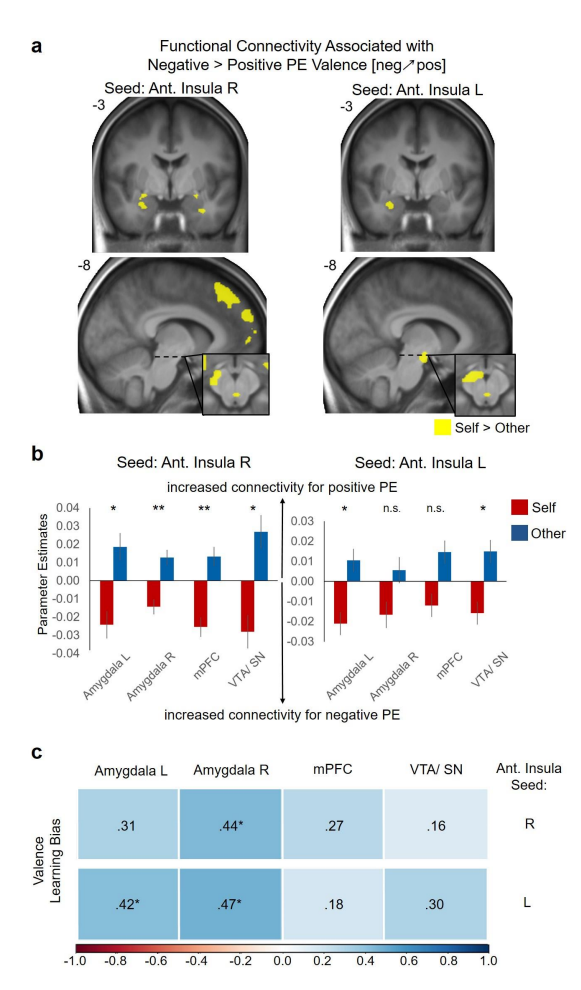


Figure 4. Differences in functional connectivity of the dorsal anterior insula during prediction 490 error (PE) valence $[neg \land pos]$ tracking for self- and other-related learning and associations 491 with the Valence Learning Bias. a) Increased functional connectivity of the dorsal anterior 492 insula for the negative effect of PE Valence [neg/pos] for self- vs other-related learning in 493 the predefined ROIs (amygdala, mPFC, VTA/ SN; p < .005 uncorrected for displaying 494 purposes). b) Functional connectivity dynamics of the dorsal anterior insula plotted separately 495 for self- and other-related learning. For displaying purposes parameter estimates are plotted 496 separately for Self and Other and refer to the peak voxels of the contrast Self vs Other that are 497 reported in Supplementary Table S8. * = p < .05, ** = p < .01. c) Spearman correlations of 498 the Valence Learning Bias with the functional connectivity dynamics between the dorsal 499 anterior insula (seed region reported on the right side) and the amygdala, mPFC and VTA/ SN 500 associated with PE valence [neg / pos] for self- vs other-related learning are color coded. * = 501 502 p < .05, FDR corrected.

503

504

Discussion

Belief formation is essentially biased and various studies have shown how motivations 505 shape belief formation (Elder et al., 2021; Sedikides and Gregg, 2008; Sedikides and Hepper, 506 2009; Sharot and Garrett, 2016). Our results demonstrate that the affect people experience 507 during learning is linked to the process of self-related belief formation on the level of neural 508 509 systems. Our computational modelling results imply that biases in the formation of selfefficacy beliefs in mastering a conceptually novel task are associated with the experience of 510 the self-conscious emotions of embarrassment and pride. Critically, on the level of neural 511 systems, the valence of PEs was associated with biases in self-related learning biases and 512 negativity of the affective experience. Individual differences in the response preference 513 towards negative PEs indicated by the pupil dilation response and activation of the AI, 514 amygdala, mPFC, and VTA/ SN were associated with a more negative learning bias and 515 negative affective experience, hinting towards a neurobiological system that integrates affect 516 517 during learning.

The novel framework on the "value of beliefs" proposed by Bromberg-Martin and Sharot (2020) nicely details how beliefs elicit emotions while at the same time these emotions shape how beliefs are updated in a reciprocal relation. Based on this framework and prior research on self-conscious emotions a negative belief about one's abilities should elicit

stronger embarrassment after failures and reduced pride after successes (Müller-Pinzler et al., 522 523 2015; Tangney et al., 2007). In the present data the association of the learning bias with the affective experience during learning supports this notion, with individuals who experience 524 more negative affect (embarrassment) and less positive affect (pride) when receiving self-525 efficacy feedback being also inclined to update their beliefs in a more negative way. Negative 526 emotions at the same time guide the information processing at various stages, including 527 528 perception, attention, and decision-making as discussed in the context of "motivated cognition" (Hughes and Zaki, 2015). This reciprocal relation finally results in biased belief 529 formation and beliefs that are both driver of affect as well as affected by emotional responses 530 531 to incoming information. Embarrassment in particular is one relevant example illustrating the 532 recursive relation between both, as the fear of failure as often discussed in the context of social anxiety (disorder) (Koban et al., 2017; Morrison and Heimberg, 2013; Müller-Pinzler et al., 533 534 2015, 2019), leads to shifts of expectations and attention (threat monitoring) towards negative information. At best this results in reparative behaviour and performance improvement (Darby 535 and Harris, 2010; Keltner and Potegal, 1997) and at worst results in a vicious cycle of fear and 536 pathologically increasing negative beliefs about the self (Heimberg et al., 2010) as it is 537 reflected in our results, when individuals who experience more intense embarrassment end up 538 539 with lower self-efficacy beliefs.

540 There are different ways on how emotions shape learning processes: First, emotions can imbue how information is processed in the brain by adaptively shifting attention towards 541 542 salient aspects of the situation (Christianson, 2014; Kaspar and König, 2012). Second, emotions entail arousal that intensifies internal rehearsal and evaluations leading to increased 543 learning (Christianson, 2014; Frijda, 1987; Storbeck and Clore, 2008), although these 544 processes often interact and are intricately related (Hughes and Zaki, 2015). The increased 545 pupil dilation in response to negative PEs in our study is in line with both increased salience 546 of and attentional shifts towards negative PEs (Koenig et al., 2018; Preuschoff, 2011) or 547

increased arousal elicited by negative PEs (Bradley et al., 2008; Müller-Pinzler et al., 2015).
Here, we think that the stronger impact of positive or negative information on pupil responses
and brain reactivity maps arousal and affect according to the valence of individual learning
biases and affective experiences.

Specifically the AI has been suggested to function as an integrative hub for motivated 552 cognition and emotional behavior (Koban and Pourtois, 2014; Wager and Feldman Barrett, 553 554 2017). While ventral aspects of the AI are associated with affective processing, emotions, and physiological arousal (Craig, 2003; Lindquist et al., 2012; Phan et al., 2002; Wager and 555 Feldman Barrett, 2017) dorsal aspects of the AI are strongly associated with the detection of 556 557 salient events, allocation of attention resources, executive working memory (Menon and 558 Uddin, 2010; Touroutoglou et al., 2012) and also (absolute) PEs and uncertainty during learning (Loued-Khenissi et al., 2020; Rutledge et al., 2010; Ullsperger et al., 2010). These 559 560 findings suggest that the functions of the AI provide a physiological basis for how emotions are translated into biased, motivated, or affected beliefs (Koban and Pourtois, 2014; Wager 561 and Feldman Barrett, 2017). A similar role as a link for the attention-emotion interaction has 562 also been suggested for the amygdala (Kaspar and König, 2012; Koban and Pourtois, 2014), 563 that shows similar responses in our task. The functional connectivity dynamics of the dAI, 564 565 matching the modelled learning rates with a stronger impact of self-related negative PEs, underline the insula's role as an integrative hub receiving and forwarding information that 566 affects information processing in other brain regions. 567

Tracking of PEs in the dopaminergically innervated VTA/ SN is influenced by motivational factors during learning (Adcock et al., 2006). The subjective value of self-related information significantly varies between subjects which is indicated by idiosyncratic response patterns of the VTA/ SN to gains or losses (Charpentier et al., 2018). In this line, we think that the present results reflect individual response tendencies at a very basic level of PE tracking. On higher layers of the computational hierarchy regions in the ACC and mPFC are also

associated with PE tracking and value representation (Hare et al., 2008; Lockwood and 574 575 Wittmann, 2018; Wallis and Kennerley, 2010) and have been previously associated with biases in belief updating (Korn et al., 2012; Kuzmanovic et al., 2016, 2018). Affect and arousal 576 could therefore bias learning on various stages of the computational hierarchy of PE 577 processing from more basic dopaminergic midbrain responses to more abstract value 578 representations in the neocortex (Diaconescu et al., 2017). While the directionality of the 579 580 effects remains to be determined the dynamics in the functional connectivity of the dAI suggest a modulatory role in this process. Here, information is forwarded to and/ or integrated 581 from VTA/ SN and mPFC, the same regions, whose response to the valence of PEs was also 582 583 modulated by differences in learning bias and affective experience. This strengthens the idea 584 that the AI plays a role in shifting responses to negative or positive information in other brain regions (e.g. by shifting attention and by affective tagging) or already receives stronger signals 585 586 in response to PEs of negative or positive valence from midbrain regions and mPFC.

The tracking of the absolute error, PE surprise (Rouhani and Niv, 2021), independent 587 of the agent, is in line with the "common currency" assumption (Izuma et al., 2008; Ruff and 588 Fehr, 2014) for the positive and negative value of one's own and others' performance 589 590 feedback. The common and valence independent coding of surprise in the insula and the 591 mPFC might therefore be sufficient to complete the learning task per se. Valence, however, matters when individuals learn about themselves as indicated by an additional shift in error 592 tracking in the same regions, AI and mPFC, which also track surprise in a valence independent 593 594 manner. As a consequence, across individuals we observe a robust effect for surprise, however, when people learn more negatively biased and experience more negative affect, this 595 596 signal is unbalanced and increases with more negative PEs. This pattern hints towards a neurocomputational mechanism of how affect shapes the formation of beliefs as proposed 597 earlier (Bromberg-Martin and Sharot, 2020). 598

599

In the current study, some of the key findings emerge at the level of individual

differences. We observed a wide inter-individual variance in the affective experience during 600 601 the task and in the Valence Learning Bias, that is, the kind of information participants preferably used to update self-related beliefs. While on average we find a negativity bias 602 during self-related belief formation, a little less than one third of the participants still shows a 603 positive learning bias, pointing out the importance of individual factors and meaningfulness 604 of variability. Studies do not only suggest that biases in belief formation differ between tasks 605 606 (Ertac, 2011; Müller-Pinzler et al., 2019; Sharot and Garrett, 2016) but also depend on situational factors like stress (Czekalla et al., 2021; Garrett et al., 2018). The individual's 607 ability to adjust the current information processing strategy to the context might be adaptive 608 609 (Bromberg-Martin and Sharot, 2020): for example, adaptation to an increased relevance of 610 negative or threat-related information during stress (Garrett et al., 2018) or coping with a negative self-concept following social stress by means of more self-beneficial belief updating 611 612 (Czekalla et al., 2021). It might also be adaptive for people who fear negative feedback to pay more attention to failure-related information in order to learn and circumvent potential future 613 failures (Sedikides and Hepper, 2009). However, it is not always straightforward to determine 614 under which conditions a strategy is adaptive or whether the affective experience can 615 ameliorate the individual's well-being. A maladaptive consequence of biased self-efficacy 616 617 beliefs becomes apparent in psychiatric disorders such as depression and social anxiety, in which amplified negative updating can lead to persistently distorted self-views and overly 618 negative beliefs about one's own capabilities in everyday life (Alden et al., 2008; Amir et al., 619 620 2012; Koban et al., 2017; Korn et al., 2014; Taylor and Brown, 1988).

621

Conclusions

Emotions experienced during learning affect computational mechanisms and manifest in distributed neural activity during belief formation. In particular, neural activity of the AI, amygdala, VTA/SN, and mPFC and pupil responses map the valence of PEs in correspondence to the experienced affect and learning bias people show during belief

formation. The more negative balancing in the functional connectivity dynamics of the dAI 626 627 during processing of self-related PEs within this network outline a scaffold for neural and computational mechanism integrating affect during belief formation. The results of the first 628 empirical spell-out of the "value of beliefs model" (Bromberg-Martin and Sharot, 2020) have 629 broader implications concerning any context which provides personal evaluations based on 630 behavioral performance. Here, the focus on the affective experience during learning provides 631 a deeper understanding on how feedback manifests in self-related beliefs which may then 632 significantly impact developmental processes and future behavior. 633

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- 635

Materials and Methods

636 **Participants**

The study was approved by the ethics committee of the University of Lübeck (AZ 18-637 066), has been conducted in compliance with the ethical guidelines of the American 638 Psychological Association (APA), and all subjects gave written informed consent. Participants 639 were recruited at the University Campus of Lübeck, were fluent in German, and had normal 640 or corrected-to-normal vision. Two independent samples were recruited, one for the fMRI 641 642 study and the other for a behavioral study that was added to increase the sample size for the 643 behavioral data. All participants received monetary compensation for their participation in the study. The final sample size for the fMRI sample was 39 participants (26 females, aged 18-28 644 years; M=22.3; SD=2.65). We initially recruited 48 participants and had to exclude six 645 646 participants because they did not believe the cover-story of the task and three participants because they did not attentively complete the task until the end (e.g. participants reported 647 having been too tired or the ratings indicated that they stopped responding to the estimation 648 task). The additional behavioral sample consisted of 30 participants (24 females, aged 18-32 649 years; M=23.3; SD=3.97). For more details on the sample characteristics see **Supplementary** 650 651 Table S9.

652 Learning of own performance task.

653 The Learning of own performance (LOOP) task enables participants to incrementally learn about their or another person's alleged ability in estimating properties. The LOOP task 654 has been previously introduced and validated in a set of behavioral studies (Müller-Pinzler et 655 al., 2019). For the LOOP task all participants were invited to take part in an experiment on 656 "cognitive estimation" together with a confederate who allegedly was another participant. In 657 658 contrast to the fMRI study, for the behavioral study two participants were invited and tested together instead of introducing a confederate. Participants were informed they would take 659 turns with the other participant/ confederate, either performing the task themselves (Self) or 660 661 observing the other person performing (Other). For the task participants were instructed to estimate different properties (e.g. the height of houses or the weight of animals). On a trial-662 by-trial basis participants received manipulated performance feedback in two distinct 663 664 estimation categories for their own estimation performance as well as for the other person's estimation performance. Unbeknownst to the participant, one of the two categories was 665 arbitrarily paired with rather positive feedback while the other was paired with rather negative 666 feedback (e.g. "height" of houses = High ability category and "weight" of animals = Low 667 ability category or vice versa; estimation categories were counterbalanced between Ability 668 669 conditions and Agent [Self vs Other] conditions). This resulted in four feedback conditions with 20 trials each (Agent condition [Self vs Other] x Ability condition [High Ability vs Low 670 Ability]). Trials of all conditions were intermixed in a fixed order with a maximum of two 671 672 consecutive trials of the same condition. Performance feedback was provided after every estimation trial, indicating the participant's own or the other person's current estimation 673 accuracy as percentiles compared to an alleged reference group of 350 university students 674 who, according to the cover-story, had been tested beforehand (e.g. "You are better than 94% 675 of the reference participants."; see Figure 1a). The feedback was defined by a sequence of 676 fixed PEs with respect to the participants' "current belief" about their abilities. The "current 677

belief" was calculated as the average of the last five performance expectation ratings per 678 679 category, which started at 50% before participants actually rated their performance expectation. This procedure led to varying feedback sequences between participants but kept 680 PEs mostly independent of the participants' performance expectations and insured a relatively 681 equal distribution of negative and positive PEs across conditions (Self: mean positive PE = 682 13.6, SD = 1.8 (average number = 20.3); mean negative PE = -12.6, SD = 1.4 (average number 683 = 19.7); Other: mean positive PE = 13.0, SD = 1.3 (average number = 19); mean negative PE 684 = -13.1, SD = 1.1 (average number = 21)). At the beginning of each trial a cue was presented 685 indicating the estimation category (e.g. "height") and participants were asked to state their 686 687 expected performance for this trial on the same percentile scale used for feedback. As part of 688 the cover story, participants were informed that accurate expected performance ratings would be rewarded with up to 6 cents per trial, that is, the better their expected performance rating 689 690 matched their actual feedback percentile the more money they would receive, to increase 691 motivation and encourage honest response behavior. Following each performance expectation rating, the estimation question was presented for 10 seconds. During the estimation period, 692 continuous response scales below the pictures determined a range of plausible answers for 693 each question. Participants indicated their responses by navigating a pointer on the response 694 695 scale with an MRI compatible computer mouse. Subsequently, feedback was presented for 3 696 seconds (see Figure 1a). Jittered inter-stimulus intervals were presented following the cue (2-6 * TR (0.992 secs), estimation (2.5 - 6.5 * TR) and feedback phase (4-8 * TR) for the fMRI 697 698 task. All stimuli were presented using MATLAB Release 2015b (The MathWorks, Inc.) and the Psychophysics Toolbox (Brainard, 1997). The fMRI task was completed in two separate 699 700 sessions of each 20 min with a short break in between.

Before starting the experiment all participants answered several questions about their
self-related beliefs and filled in a self-esteem personality questionnaire (SDQ-III; Marsh &
O'Neill, 1984). During the LOOP task participants were also asked to rate their current levels

of embarrassment, pride, happiness and stress/ arousal on a continuous scale ranging from not 704 705 at all (coded as 0) to very strong (coded as 100). Two emotion rating phases followed self-706 related feedback and two rating phases followed other-related feedback. The two emotion rating phases following self-related feedback were averaged to receive a rating for the 707 experience of self-conscious affect (embarrassment and pride) during self-related learning. 708 After the task participants completed an interview including ratings about self-related beliefs, 709 were debriefed about the cover-story, and reimbursed for their time before leaving. The whole 710 procedure took approximately 2 h. 711

712 Behavioral Data Analysis and Modeling

A model free analysis was performed on the participants' expected performance ratings for each trial to illustrate the basic effects we see in our behavioral data. A repeatedmeasures ANOVA was calculated with the factors Trial (20 Trials) x Ability condition (High ability vs Low ability) x Agent condition (Self vs Other) as well as Group as a between-subject factor to control for potential differences between the two samples. All statistical analyses on the behavioral data apart from the modeling procedure were performed using *jamovi* (Version 1.2.27, The jamovi project (2020). Retrieved from https://www.jamovi.org).

720 Dynamic changes in self-related efficacy beliefs, that is, performance expectation 721 ratings, were then modeled using PE delta-rule update equations (adapted Rescorla-Wagner model; Rescorla & Wagner, 1972). The model space contained three main models varying 722 with regards to their assumptions about biased updating behavior when learning about the self 723 724 (see Supplementary Figure S1). The simplest learning model used one single learning rate for all conditions for each participant, thus not assuming any learning biases (Unity Model). 725 726 The second model, the Valence Model, included separate learning rates for positive PEs (α_{PE+}) vs negative PEs (α_{PE}) across both ability conditions, thus suggesting that the valence (positive 727 vs negative) of the PE biases self-related learning. The third model, the Ability Model, 728 729 contained a separate learning rate for each of the ability conditions indicating context specific

learning. In addition, learning rates were either estimated separately for Self vs Other or across 730 731 Agent conditions. The Valence Model with separate learning rates for Self vs Other (Model 732 5), winning model in our previous studies (Czekalla et al., 2021; Müller-Pinzler et al., 2019), was further extended by adding a weighting factor reducing learning rates towards the ends 733 of the feedback scale (percentiles close to 0 % or 100 %), assuming that participants 734 experienced extreme feedback values as less likely than more average feedback (Kube et al., 735 736 2021). In the first model of these (Model 7) a linear decrease of the learning rates was assumed beginning at 50 % and ending at 0 % and 100 %. A weighting factor w was fitted for each 737 participant defining how strong the linear decrease was present for each individual. Since 738 739 many variables people encounter in every-day life (e.g. many test results) approximately follow a normal distribution with extreme values being less likely, for the second model of 740 this kind (Model 8) we assigned the relative probability density of the normal distribution to 741 742 each feedback percentile value. Again, a weighting factor w was fitted for each individual indicating how strongly the relative probability density reduced the learning rates for feedback 743 744 further away from the mean. In contrast to our previous studies implementing the LOOP task with fixed feedback sequences, here, feedback depended on the participants' current 745 746 expectations and thus differed between participants and conditions. Reduced learning rates 747 towards the ends of the feedback scale which could systematically confound learning rates between participants and conditions could thus be accounted for in Models 7 and 8. To test if 748 the participants' performance expectation ratings can be better explained in terms of PE 749 750 learning as compared to stable assumptions in each Ability condition, we included a simple Mean Model with a mean value for each task condition (Model 9; for more details see 751 Supplementary Methods). 752

753 Model Fitting.

For model fitting we used the RStan package (Stan Development Team, 2016. RStan:
the R interface to Stan. R package version 2.14.1.), which uses Markov chain Monte Carlo

(MCMC) sampling algorithms. All of the learning models in the model space were fitted for 756 757 each participant individually and posterior parameter distributions were sampled for each participant. A total of 2400 samples were drawn after 1000 burn-in samples (overall 3400 758 samples; thinned with a factor of 3) in three MCMC chains. We assessed if MCMC chains 759 converged to the target distributions by inspecting \hat{R} values for all model parameters (Gelman 760 and Rubin, 1992). Effective sample sizes (n_{eff}) of model parameters, which are estimates of 761 the effective number of independent draws from the posterior distribution, were typically 762 greater than 1500 (for most parameters and subjects). Posterior distributions for all parameters 763 764 for each of the participants were summarized by their mean as the central tendency resulting in a single parameter value per participant that we used in order to calculate group statistics. 765

766

Bayesian Model Selection and Family Inference.

For model selection we estimated pointwise out-of-sample prediction accuracy for all 767 fitted models separately for each participant by approximating leave-one-out cross-validation 768 (LOO; corresponding to leave-one-trail-out per subject; Acerbi et al., 2018; Vehtari et al., 769 770 2016). To do so, we applied Pareto-smoothed importance sampling (PSIS) using the log-771 likelihood calculated from the posterior simulations of the parameter values as implemented 772 by Vehtari et al. (2016). Sum PSIS-LOO scores for each model as well as information about \hat{k} values – the estimated shape parameters of the generalized Pareto distribution – indicating 773 774 the reliability of the PSIS-LOO estimate are depicted in Supplementary Table S1. As summarized in **Supplementary Table S1** very few trials resulted in insufficient parameter 775 values for \hat{k} and thus potentially unreliable PSIS-LOO scores (on average 1.1 trials per subject 776 with $\hat{k} > 0.7$ for the winning model; Vehtari et al., 2016). BMS on PSIS-LOO scores was 777 performed on the group level accounting for group heterogeneity in the model that best 778 describes learning behavior (Rigoux et al., 2014). This procedure provides the protected 779 780 exceedance probability for each model (*pxp*), indicating how likely a given model has a higher 781 probability explaining the data than all other models in the comparison set. The Bayesian omnibus risk (*BOR*) indicates the posterior probability that model frequencies for all models
are all equal to each other (Rigoux et al., 2014). We also provide difference scores of PSISLOO in contrast to the model that won the BMS that can be interpreted as a simple 'fixedeffect' model comparison (see Supplementary Table S1; Acerbi et al., 2018; Vehtari et al.,
2016). Model comparisons according to PSIS-LOO difference scores were qualitatively
comparable to the BMS analyses for our data.

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Posterior Predictive Checks and Statistical Analyses of Learning Parameters

First, posterior predictive checks were conducted by quantifying if the predicted data could capture the variance in performance expectation ratings for each subject within each of the experimental conditions using regression analyses. Additionally, we repeated the model free analysis we had conducted on the behavioral data with the data predicted by the winning model to assess if the winning model captured the core effects in the behavioral data (see **Supplementary Results**).

Model parameters, i.e. learning rates, of the winning models for all experiments were analyzed on the group level. A repeated-measures ANOVA was calculated on the learning rates with the factor Agent (Self [$\alpha_{Self/PE+}$, $\alpha_{Self/PE-}$] vs Other [$\alpha_{Other/PE+}$, $\alpha_{Other/PE-}$]) and factor PE valence [pos| neg] (PE+ [$\alpha_{Self/PE+}$, $\alpha_{Other/PE+}$] vs PE- [$\alpha_{Self/PE-}$, $\alpha_{Other/PE-}$]) as well as Group as a between-subject factor testing if learning about one's own performance was more valence specific as compared to learning about the other person's performance.

To associate learning biases with self-conscious affect, that is, embarrassment and pride, and self-esteem (SDQ-III subscale scores) we calculated a normalized learning rate valence bias score for self-related learning (Valence Learning Bias= $(\alpha_{PE+(S)} - \alpha_{PE-(S)})/(\alpha_{PE+(S)} + \alpha_{PE-(S)}))$ (Müller-Pinzler et al., 2019; Niv et al., 2012; Palminteri et al., 2017). Spearman correlations were calculated between Valence Learning Bias, affect ratings, and self-esteem scores.

807 **Pupil Data Analysis**

For the fMRI sample, eye-tracking data were assessed during scanning. Pupil diameter 808 809 and gaze behavior were recorded non-invasively in one eye at 500 Hz using an MRIcompatible Eyelink-1000 plus device (SR Research, Kanata, ON, Canada) with manufacturer-810 recommended settings for calibration and blink detection. Due to insufficient pupillometry 811 data quality three participants had to be excluded from analyses (final sample n=36). Pupil 812 data were preprocessed by cutting out periods of blinks and values in this gap were 813 interpolated by piecewise cubic interpolation. The pupil trace was subsequently z-normalized 814 over the whole session. To characterize the pupil dilation for each trial by a single value, we 815 calculated a linear slope for each feedback phase of three seconds. Pupil traces were only 816 817 analyzed for the Self condition as onsets during feedback strongly differed between Agent 818 conditions, which made a meaningful comparison between pupil slopes impossible. Pupil slopes during self-related feedback phases for each trial were then entered in linear mixed 819 models fitted by restricted maximum likelihood including PE valence [neg/pos] (continuous 820 signed PE values) and PE surprise (continuous unsigned/ absolute PE values) as fixed effects 821 822 and participant and PE valence [neg ≠ pos] as random effects. Additionally, separate linear mixed models including embarrassment ratings, pride ratings or the Valence Learning Bias as 823 well as their interaction with PE valence [neg/pos] were implemented to assess if variance 824 in individual pupil responses to positive and negative PEs (random PE valence [neg/pos] 825 slopes) was explained by with different emotional reactions and learning behavior (see Figure 826 **2c**). 827

828 fMRI Data

829 **fMRI Image Acquisition**

Participants were scanned using a 3T Siemens MAGENTOM Skyra scanner (Siemens,
München, Germany) at the Center of Brain, Behavior, and Metabolism (CBBM) at the
University of Lübeck, Germany with 60 near-axial slices. An echo planar imaging (EPI)
sequence was used for acquisition of on average 1520 functional volumes (min=1395, max=

1672) during each of the two sessions of the experiment, resulting in a total of on average 3040 functional volumes (TR=0.992s, TE=28ms, flip angle=60°, voxel size= $3 \times 3 \times 3$ mm³, simultaneous multi-slice factor 4). In addition, a high-resolution anatomical T1 image was acquired that was used for normalization (voxel size= $1 \times 1 \times 1$ mm³, $192 \times 320 \times 320$ mm³ field of view, TR= 2.300s, TE = 2.94ms, TI = 900ms; flip angle=9°; GRAPPA factor 2; acquisition time 6.55 min).

840 FMRI data analyses

FMRI data were analyzed using SPM12 (www.fil.ion.ucl.ac.uk/spm). Field maps were 841 reconstructed to obtain voxel displacement maps (VDMs). EPIs were corrected for timing 842 843 differences of the slice acquisition, motion-corrected and unwarped using the corresponding 844 VDM to correct for geometric distortions and normalized using the forward deformation fields as obtained from the unified segmentation of the anatomical T1 image. The normalized 845 846 volumes were resliced with a voxel size of $2 \times 2 \times 2$ mm and smoothed with an 8 mm fullwidth-at-half-maximum isotropic Gaussian kernel. To remove low-frequency drifts, 847 functional images were high-pass filtered at 1/384. 848

Statistical analyses were performed in a two-level, mixed-effects procedure. Three 849 main GLMs were implemented on the first level. The first fixed-effects GLM included four 850 851 epoch regressors modeling the hemodynamic responses to the different cue conditions (Ability: High vs Low \times Agent: Self vs Other), weighted with the performance expectation 852 ratings per trial as parametric modulator for each condition. Four regressors modeled the four 853 854 feedback conditions (PE valence [pos| neg]: Positive vs Negative × Agent: Self vs Other), each weighted with the PE value for each trial. The estimation periods for Self and Other as 855 two regressors, and emotion ratings phase and the instruction phase as separate regressors. To 856 account for noise due to head movement, six additional regressors modeling head movement 857 parameters were introduced and a constant term was included for each of the two sessions. 858 The second first-level GLM differed only with respect to the feedback regressors. Here, only 859

two regressors modeled feedback separately for Self and Other and two parametric modulators 860 were included per condition weighting feedback trials with PE valence [neg/pos] (continuous 861 effect of the signed PE values) and PE surprise (continuous effect of the unsigned PE values). 862 The third first-level GLM was set-up to show that activation found in response to PEs was 863 actually related to PEs and not only to feedback level alone. Therefore, the parametric weights 864 865 of the two feedback conditions in the second GLM were replaced by feedback level and performance expectation ratings, allowing us to assess if neural activity goes up with feedback 866 level and down with performance expectation ratings confirming a potential interpretation in 867 terms of PE tracking (Zhang et al., 2020). 868

On the second level for the first GLM model beta images for the four feedback 869 870 conditions were included in a flexible factorial design with two repeated-measurement factors (PE valence [pos| neg] and Agent). Beta images for the parametric weights of feedback were 871 extracted from the second and third first-level model for Self and Other. Separate repeated-872 measures ANOVAs and one sample t-tests (for baseline contrasts) were implemented for PE 873 874 valence $[neg \land pos]$ and PE surprise as well as feedback level and performance expectation level. Additional second level models for the PE valence [neg / pos] contrast included the 875 Valence Learning Bias, embarrassment and pride ratings as between subject covariates, 876 assessing differential tracking of PEs depending on biased learning and self-conscious affect. 877 A self-related Pupil Dilation Bias (average slope for positive PEs - average slope for negative 878 PEs; higher scores indicate stronger pupil dilation for positive PEs) was also included as 879 covariate in another second level model to assess if the neural response towards negative vs 880 881 positive PEs was associated with the pupil dilation response.

We additionally performed psychophysiological interaction (PPI) analyses on the first level, investigating whether functional connectivity of the dAI, that is commonly activated during feedback processing independent of agent and feedback valence (conjunction of baseline contrasts: feedback Self \land feedback Other) would differ depending on the PE valence

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886 $[neg \nearrow pos]$. PPI analyses were computed separately for Self and Other and the resulting contrast images for the PPI effects were aggregated on the second level using two-sample t-887 tests contrasting PPI effects for Self vs Other. For each participant, we defined 6-mm radius 888 889 spherical ROIs, centered at the nearest local maximum for the conjunction contrast feedback Self \land feedback Other and located within 10 mm from the group maximum within the dAI, 890 891 separately for the left dAI (x, y, z: -33 20 -4) and right dAI (x, y, z: 36 20 -7). By computing the first eigenvariate for all voxels within these ROIs that showed a positive effect for the 892 conjunction (p < .500), we extracted the time course of activations and constructed PPI terms 893 894 using the contrast for the parametric weights of PE valence [neg / pos] for Self or Other, respectively, resulting in four distinct PPI first level GLMs. One participant was excluded 895 from the PPI analyses for the right dAI, because no voxels survived the predefined threshold 896 897 for eigenvariate extraction. The PPI term, along with the activation time course from the (left 898 or right) dAI was included in a new GLM for each participant that also included all the regressors in the initial first level GLM (four regressors for the different cue conditions, each 899 weighted with the expected performance ratings; two feedback regressors for Self and Other 900 with each two parametric modulators for PE valence [neg 2 pos] and PE surprise; two 901 902 regressors for the estimation periods for self and other; one regressor for the emotion ratings phase; one regressor for the instruction phase; six regressors modeling head movement 903 parameters; a constant term for each session). On the second level we assessed if there was a 904 stronger functional coupling of the dAI with our predefined ROIs (Amygdala, mPFC, VTA/ 905 SN) for the Self in contrast to the Other when PE valence $[neg \neq pos]$ was more negative. 906 907 Functional connectivity dynamics were also associated with learning behavior by calculating Spearman correlations for the Valence Learning Bias and the parameter estimates for the PPI 908 909 effect of Self > Other derived from a sphere of 6mm around the peak voxels within our predefined ROIs. 910

911 Thresholding procedure and regions of interest

According to its suggested role as an integrative hub for motivated cognition and 912 913 emotional behavior the AI was defined as one of the regions of interest (ROIs) (Koban and Pourtois, 2014; Wager and Feldman Barrett, 2017). Due to their specific functional 914 associations, a bilateral ventral and a bilateral dorsal AI ROI was defined according the three 915 cluster solution by Kelly and colleagues (2012). The bilateral amygdala was defined as another 916 ROI and derived from the AAL atlas definition in the WFU PickAtlas (Tzourio-Mazover et 917 al., 2002) due to its similar role for the attention-emotion interaction (Kaspar and König, 2012; 918 Koban and Pourtois, 2014). The mPFC ROI was also derived from the AAL atlas in the WFU 919 PickAtlas (label: bilateral frontal superior medial) due to its specific role during social learning 920 921 and for biases in self-related belief updating in previous studies (Kuzmanovic et al., 2018; Sharot, 2011). Additionally, a VTA/ SN ROI, dopaminergic nuclei in the midbrain, was 922 included (Ballard et al., 2011; Murty et al., 2014) as dopamine signals motivationally important 923 924 events, e.g. during reward learning (Schultz, 1998), and has been associated with biases in memory towards events that are of motivational significance (Adcock et al., 2006). 925

FMRI results were family-wise-error (FWE) corrected on the whole brain level if not
mentioned otherwise and all coordinates are reported in MNI space. As our predefined ROIs
were chosen with respect to their involvement with the emotion-cognition link, we tested the
effects of our covariates on PE valence [neg *P* pos] tracking and PPI effects within the ROIs.
Anatomical labels of all resulting clusters were derived from the Automated Labeling Atlas
Version 3.0 (Eickhoff et al., 2005).

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