Mapping social reward and punishment processing in the human brain: A voxel-based meta-analysis of neuroimaging findings using the social incentive delay task

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ABSTRACT
Social rewards or punishments motivate human learning and behaviour, and alterations in the brain circuits involved in the processing of these stimuli have been linked with several neuropsychiatric disorders. However, questions still remain about the exact neural substrates implicated in social reward and punishment processing. Here, we conducted four Anisotropic Effect Size Signed Differential Mapping voxel-based meta-analyses of fMRI studies investigating the neural correlates of the anticipation and receipt of social rewards and punishments using the Social Incentive Delay task. We found that the anticipation of both social rewards and social punishment avoidance recruits a wide network of areas including the basal ganglia, the midbrain, the dorsal anterior cingulate cortex, the supplementary motor area, the anterior insula, the occipital gyrus and other frontal, temporal, parietal and cerebellar regions not captured in previous coordinate-based meta-analysis. We identified decreases in the BOLD signal during the anticipation of both social reward and punishment avoidance in regions of the default-mode network that were missed in individual studies likely due to a lack of power. Receipt of social rewards engaged a robust network of brain regions including the ventromedial frontal and orbitofrontal cortices, the anterior cingulate cortex, the amygdala, the hippocampus, the occipital cortex and the brainstem, but not the basal ganglia. Receipt of social punishments increased the BOLD signal in the orbitofrontal cortex, superior and inferior frontal gyri, lateral occipital cortex and the insula. In contrast to the receipt of social rewards, we also observed a decrease in the BOLD signal in the basal ganglia in response to the receipt of social punishments. Our results provide a better understanding of the brain circuitry involved in the processing of social rewards and punishment. Furthermore, they can inform hypotheses regarding brain areas where disruption in activity may be associated with dysfunctional social incentive processing during disease

1. Introduction
The quest for social acceptance and belonging is considered a basic motive of humans. Positive social feedback, such as smiling faces, encouraging gestures and verbal praise are rewarding and can act as positive reinforcers, increasing the likelihood that the associated behaviour will be repeated in the future (Fehr and Camerer, 2007). In contrast, the prospect of socially undesirable outcomes, such as a face or a gesture of disapproval, may act as negative reinforcers, that is, increase the likelihood that the expected behaviour (in the case of the SID task, a

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fast reaction time to the target) will occur, so as to avoid them. Their occurrence may also act as punishments, e.g. reduce the likelihood that behaviours that led to them (e.g. lapses of attention during the task) occur. Social rewards or punishments are crucial for learning and adaptive behaviour (Fehr and Camerer, 2007). For instance, social actions (e.g. mutual face-to-face interactions) are often driven by the goal to increase one’s own social standing (Fehr and Camerer, 2007; Lieberman, 2007). Anticipation of social feedback in the form of praise (and reprimand) has been regarded as one of the most important educative modulator of social behaviour for centuries (Fehr and Camerer, 2007).

Disruption of the brain circuits processing social rewards (i.e. lack of motivation to invest in interactions with others (Fuldolf et al., 2018)) and punishments (i.e. hypersensitivity to social disapproval (Kumar et al., 2017a)) has been suggested to be at the heart of several neuropsychiatric disorders characterised by dysfunction in social interactions, such as autism spectrum disorder (Kohls et al., 2013), psychosis (Radua et al., 2015) or mood disorders (Naranjo et al., 2001). However, questions still remain about the exact neural circuitries underlying the processing of social rewards and punishments in the human brain.

The neural mechanisms prompting an organism to approach/avoid a potential reward/punishment have been extensively explored in several studies using monetary incentives (for an overview see (Dugre et al., 2018a; Oldham et al., 2018a; Wilson et al., 2018)). Among the various paradigms available, the monetary incentive delay (MID) task is among the most widely-used ones. Performance-dependent reward and punishment processing can be divided in two distinct temporal phases: an anticipation phase, where the prospect of a reward/loss is initially encountered, and an outcome phase (also called the receipt or consummatory phase), where the reward/loss is received or omitted (Oldham et al., 2018a). In the classical MID task, participants are asked to perform a simple motor reaction time task (i.e. press a button while a target is on the screen) in various incentive contexts defined by the presentation of cues indicating one of a range of positive or negative monetary rewards or punishments that may follow the response to the target. The cue is followed by a short delay period (the anticipation phase), before the target appears. Participants are instructed to respond to the target as quickly as possible, and the success rate is set to a fixed value (typically at 67 % of the trials) by adjusting the duration of the target presentation trial by trial and for each individual separately. Finally, in the outcome/feedback phase, the performance-dependent outcome is presented (attainment or loss of a potential monetary gain, or no monetary outcome in neutral trials).

The combination of the MID task design with functional magnetic resonance imaging (fMRI) offers the opportunity to assess the neural signatures specifically associated with the anticipation or receipt/avoidance of monetary wins/losses. Recent meta-analyses of fMRI studies using the MID task have identified several areas consistently involved in the anticipation and/or receipt of monetary rewards/punishments (Dugre et al., 2018a; Oldham et al., 2018a; Wilson et al., 2018). These meta-analyses have found considerable overlaps in the networks activated during monetary reward and loss anticipation, including the dorsal and ventral striatum, amygdala, insula and supplementary motor cortex. They have also identified neural networks engaged by monetary reward receipt — including the ventral striatum, the orbitofrontal cortex/ventromedial prefrontal cortex, amygdala, and the posterior cingulate.

Building on findings from the MID task, some studies have inspected if the same neural circuits that underpin the processing of monetary rewards and punishments also underpin the processing of their social counterparts. Studies have investigated the neural underpinnings of the anticipation and receipt of social rewards/punishments using a variant of the MID task known as the Social or Affective Incentive Delay task – henceforth referred to as SID task (for an overview see (Gu et al., 2019)). The SID task shares a similar structure as the MID task, but uses social stimuli that are inherently reinforcing or punishing (i.e. smiling or angry faces, positive/negative verbal messages, thumbs-up or down gestures) as stimuli. Fewer studies have used the SID task than those using the MID task though. Individually, studies using the SID task have started to delineate the brain regions involved in the processing of social rewards/punishments (Gu et al., 2019). On the one hand, there is evidence that the processing of social rewards frequently evokes the activity of generic brain regions involved in reward prediction and value encoding, such as the striatum and the orbitofrontal cortex (Barman et al., 2015b; Rademacher et al., 2014; Spreckelmeyer et al., 2009). On the other hand, some studies have also suggested that the processing of social rewards might involve additional brain regions thought to be critical for social-cognitive processes, such as the temporoparietal junction, dorsomedial prefrontal cortex, precuneus, and superior temporal gyrus (Barman et al., 2015b; Goerlich et al., 2017b; Spreckelmeyer et al., 2013b). At present, it is still under debate whether social rewards and punishment engage specific neural circuits related to their social nature or are rather encoded in an analogue manner to the neural encoding of other non-social rewards, such as money (common currency hypothesis) (Gu et al., 2019). In part, this debate stems from the lack of a clear picture of the brain circuits underpinning social reward and punishment processing in the human brain, which is still missing due to inconsistencies in the areas identified across individual studies. This lack of consistency further makes it difficult to inform targeted hypotheses about potentially disrupted areas in neuropsychiatric disorders, based on our understanding of the neural underpinnings of social reward/punishment processing in healthy individuals (Cremers et al., 2015; Delmonte et al., 2012; Kohls et al., 2018), limiting any research studies trying to understand how disruption of the brain processing of social rewards and punishments might contribute to social dysfunction during disease.

Meta-analyses can provide valuable help in our efforts to delineate a precise and fine-grained characterization of the neural substrates involved in both the anticipation and receipt phases of social incentive processing. They can help address one major limitation of individual studies, namely the lack of sufficient power due to small sample sizes. Indeed, lack of statistical power is a well-documented phenomenon in neuroimaging studies (Cremers et al., 2017; Poldrack et al., 2017) and comes at the cost of increased risks for both type I and type II errors, undermining replicability across independent studies (Button et al., 2013; Cremers et al., 2017; Kim, 2015). Therefore, meta-analyses of neuroimaging studies have become increasingly important (Muller et al., 2018).

To date, there has been only one attempt to quantitatively summarize neuroimaging findings from the SID task (Gu et al., 2019). This meta-analysis focused specifically on the anticipation of social rewards (it did not examine the receipt phase of social rewards, or the processing of social punishment at all). Given the role that differences in the processing of social punishments may play in a wide range of neuropsychiatric disorders (such as major depressive disorder (Kumar et al., 2017b) or antisocial personality disorder (Gong et al., 2019)), characterizing the neural underpinnings of social punishment processing in healthy samples is paramount to identify how disruption of these circuits might give rise to impairments in social cognition during pathology. Furthermore, this previous study used a method for the meta-analysis of neuroimaging data, Activation Likelihood Estimation (ALE) that, although popular, comes with some important limitations (Radua and Mataix-Cols, 2012). For instance, the ALE method is based on the reported peak coordinates in individual studies. However, reported peak coordinates may be biased due to the dependence on the statistical significance thresholds that were used, which may be arbitrary, and the power in individual studies, which tends to be less than the commonly accepted standard of at least 80 %, as highlighted above (Cremers et al., 2017).

Furthermore, an ALE meta-analysis does not produce a statistical measure of effect-size or its variance, or show the direction of the effect, thus making the interpretation of the biological significance of the results challenging. Instead, an ALE meta-analysis provides an informative summary of statistically significant fMRI results across a number of
studies in a field, based on the spatial convergence of neuroimaging findings across experiments (Radua and Mataix-Cols, 2012).

Here, we aimed to quantitatively synthesize the evidence from fMRI studies using the SID tasks to investigate social reward and punishment processing in humans. We decided to specifically focus on the SID task for three reasons. First, the design of this task allows us to disentangle neural correlates associated with the anticipation and receipt phases of social reward and punishment processing (Gu et al., 2019). Hence, by focusing on this specific task we will be able to provide detailed mapping of the brain regions engaged by each of these two phases of the task. Second, focusing on a homogenous well-established task design will allow us to avoid issues of heterogeneity, as would be the case if we pooled studies with large differences in task design and therefore in the underlying cognitive processes. Third, thanks to the similarity of the SID task with its monetary counterpart, the MID task, which has been widely used in many fMRI studies, we can perform straightforward comparisons between the neural substrates engaged in the anticipation and receipt of monetary and social outcomes without the additional confound of substantial differences in task design (Dugue et al., 2018a; Oldham et al., 2018a; Wilson et al., 2018).

As a result of our meta-analyses, we generate a fine-grained characterization of the neural circuitry implicated in the anticipation and receipt of both social rewards and punishments. To achieve this, we used Anisotropic Effect Size Signed Differential Mapping (AES-SDM) (Radua et al., 2012a, 2014), a meta-analytic method that can overcome some of the limitations of the previous meta-analysis on social reward anticipation using data from the SID task highlighted above (Gu et al., 2019). AES-SDM can combine reported peak information (coordinates and t-values) from some studies, with original statistical parametric maps (SPMs) from other studies, to produce an estimate of the magnitude and direction of the effect sizes of changes in the BOLD signal. The inclusion of unthresholded original statistical maps helps to address issues of low statistical power by preserving information from voxels that did not reach significance in the original studies because of power limitations. In addition, since AES-SDM is a method based on effect sizes, it preserves information on the sign of the effect (e.g. increases or decreases in the BOLD signal), therefore allowing a more straightforward and biologically plausible interpretation of the results (Radua and Mataix-Cols, 2012).

2. Materials and methods

2.1. Literature search

We conducted searches in PubMed, EMBASE and OVID using the terms (“social incentive delay” OR “affective incentive delay” OR “SID” OR “AID”) AND (“fMRI” OR “functional magnetic resonance imaging”) on 3rd May 2018. Our search query was adapted in accordance with the specification of each database. Our search strategy was tailored to include all human task-based fMRI studies conducted with healthy individuals, published up to this date (irrespective of publication date or age, sex and ethnicity of the subjects), and which were original reports on the neural substrates associated with social reward and punishment anticipation and receipt (as assessed by task-based fMRI using the social/affective incentive delay task).
these peaks, using an un-normalised Gaussian kernel, which is multiplied by the effect size of the peak. As part of an automatic step implemented by the AES-SDM software, all coordinates are first converted to a common space (Talairach) before effect sizes are calculated (Radua et al., 2012a, 2014). The use of effect sizes in the calculation has been shown to increase the accuracy of estimation of the true signal compared to alternative methods (Radua et al., 2012a). Additionally, the inclusion of statistical parametric maps has been shown to substantially increase the sensitivity of voxel-based meta-analyses. For example, in the initial validation study of the method, sensitivity increased from 55% to 73% and 87% with the inclusion of just one and two statistical maps, respectively (Radua et al., 2012a). The detailed AES-SDM approach has been described elsewhere (Radua et al., 2012a, 2014). The use of effect sizes in the calculation has been shown to increase the accuracy of estimation of the true signal compared to alternative methods (Radua et al., 2012a). Additionally, the inclusion of statistical parametric maps has been shown to substantially increase the sensitivity of voxel-based meta-analyses. For example, in the initial validation study of the method, sensitivity increased from 55% to 73% and 87% with the inclusion of just one and two statistical maps, respectively (Radua et al., 2012a). The detailed AES-SDM approach has been described elsewhere (Radua et al., 2012a, 2014).

We conducted four separate meta-analyses, one for each of our four contrasts of interest. For each of the meta-analyses, meta-analytic effect-sizes were voxel-wise divided by their standard error to obtain AES-SDM z-values. As AES-SDM z-values for each meta-analytic brain map may not follow a standard normal distribution, a null distribution was empirically estimated for each meta-analysis. Specifically, null distributions were obtained from 50 whole brain permutations (which, multiplied by the number of voxels, resulted in about four million values per null distribution); previous simulation work has found that permutation-derived AES-SDM thresholds are already stable with even 5 whole-brain permutations (Radua et al., 2012b). Voxels with AES-SDM z-values corresponding to p-values < 0.001 were considered significant, but voxels with AES-SDM z-values < 1, or in clusters with less than 10 voxels, were discarded in order to reduce the false positive rate. While these thresholds do not strictly apply family-wise correction for multiple comparisons, previous empirical validation work of this method has found these parameters to provide optimal sensitivity while maintaining the false positive rate below 5% (Radua et al., 2012a).

2.4.2. Sensitivity and sub-group analyses

We assessed the robustness of our findings by conducting Jackknife sensitivity analysis where we iteratively repeated the meta-analysis for each of the four contrasts leaving out one study at a time. In order to synthesize mean Jackknife maps for each contrast and meaningfully interpret them, we first thresholded each individual Jackknife map, for each contrast, using the significance thresholds outlined above, and then binarized them and combined their information into a single overlapping density map of significant voxel-wise data for each contrast. This allowed the visual identification of voxels in terms of density (the higher the density, the more the individual Jackknife maps that a given voxel reached significance), which provides an estimate of replicability across studies. Currently there is no standard definition of what a robust finding should be in AES-SDM leave-one-out meta-analyses. Hence, we decided to adopt an operational definition for our own appraisal of the data where we defined as robust every voxel that was present in more than 75% of the repetitions of the meta-analyses using the Jackknife leave-one-out procedure, showing that our main findings are not driven by the inclusion of specific studies.

The studies we considered used a variety of methodological approaches. Some studies used verbal feedback while other studies used emotional faces for feedback, some studies used static and others dynamic emotional faces, some studies examined the effects of a pharmacological intervention using a placebo-controlled design (we only considered data form the placebo condition). Therefore, we ran further sensitivity analyses where the number of included studies permitted it (that is, where >5 studies would be included in the respective sub-group meta-analysis). Specifically, for our anticipation and receipt of social reward meta-analyses, we ran meta-analyses on subgroups of studies: 1) including only studies using emotional faces as feedback; 2) including only studies using static emotional faces; 3) including only non-pharmacological studies. By default, we did not conduct any of these sub-group meta-analyses for the anticipation and receipt of social punishment conditions as the number of included studies would be less than 5. However, we did conduct one subgroup meta-analysis for the social punishment avoidance anticipation condition because one study used an implementation of the SID task that was conceptually different from the rest (Nawijn et al., 2017). Specifically, in Nawijn’s et al. implementation of the task, the success rate in the social punishment avoidance condition was set to approximately 34%, making social punishment avoidance the least frequently anticipated outcome (in all other studies, the success rate in the social punishment avoidance condition was set to approximately 66%). The subgroup analysis would allow us to test whether our results were unduly influenced by the inclusion of Nawijn’s et al. study.

2.4.3. Heterogeneity and publication bias

Heterogeneity is a problem that can arise when undertaking meta-analysis (Muller et al., 2018; Radua and Mataix-Cols, 2012). Ideally, a meta-analysis should combine results from studies using the same procedures and experimental protocols. Heterogeneity refers to the proportion of variability across studies that is due to methodological
variability relative to that from sampling error. Heterogeneity can be quantified and provide an index of whether the assumption that all studies use the same procedures and experimental protocols is met (Radua and Mataix-Cols, 2012). When considerable heterogeneity is found, it raises questions about how valid it is to combine all the data in one single meta-analysis. When possible, instances of substantial heterogeneity should be investigated further using subgroup and metaregression analysis (described below) to identify potential factors underlying this heterogeneity. We used the $I^2$ statistic maps provided by the software to assess heterogeneity. To identify areas of increased heterogeneity, we thresholded these maps for $I^2 > 40\%$ (since values of $I^2 < 40$ are typically assumed to not constitute important heterogeneity (Higgins et al., 2003)) and masked them to retain only voxels where we found significant increases/decreases in BOLD in the main meta-analyses. Voxels with $I^2$ between 40 and 60 indicate areas of moderate heterogeneity, between 60 and 90 substantial heterogeneity and higher than 90 strong heterogeneity (Higgins et al., 2003).

In turn, publication bias occurs when the outcome of an experiment or research study influences the decision to publish it (Jøsper et al., 2012). Increased publication bias means that a study is less likely to be published if the findings are null. We used funnel plots and the Egger test as implemented by the software to examine publication bias (Lin and Chu, 2018). Briefly, effect size estimates were extracted from the constructed effect size maps of each included study for the peak voxel of each of the clusters identified in each of the four meta-analyses we conducted. Using these, funnel plots were created and visually inspected. A funnel plot displays effect sizes (X-axis) against a measure of the study’s precision (i.e. sample size, Y-axis). In the absence of publication bias, studies with high precision will cluster around average effect sizes, while studies with low precision should be spread evenly on both sides of the average effect size, creating a roughly funnel-shaped distribution. Deviation from this symmetric shape can indicate publication bias (Lin and Chu, 2018) and should call for caution in interpreting such findings. We used the Egger regression test as a quantitative method of assessing asymmetry in the funnel plots. Potential publication bias is indicated if the intercept of the regression of effect size/standard error on 1/standard error significantly deviates from zero ($p < 0.05$) (Lin and Chu, 2018).

### 2.4.4. Meta-regressions

Previous studies have found that age (Rademacher et al., 2014) and gender (Greimel et al., 2018; Spreckelmeyer et al., 2009; Wang et al., 2017) differentially modulate the brain’s response during social and monetary incentive anticipation. For instance, one of our previous studies exploring the effects of age found that the nucleus accumbens response to cues of reward was higher for social when compared to monetary rewards in an older sample, but the opposite was true in a younger sample (Rademacher et al., 2014). Another study exploring the effects of gender showed that while in women the anticipation of monetary and social rewards engages the same brain areas, in men the anticipation of monetary rewards engaged a much wider network of brain areas in comparison to social rewards (Spreckelmeyer et al., 2009). Therefore, we considered that it was important to elucidate the impact of age and gender on the BOLD response to social reward and punishment anticipation or receipt. To achieve this, we conducted separate voxel-wise meta-regressions including mean age or the proportion of male participants across studies as moderator variables. Metaregression is a classical meta-analytic approach that allows the exploration of the impact of moderator variables on the effect sizes reported across studies using regression-based techniques (Lipsey, 2003). We took a conservative approach where we corrected the significance level for the number of meta-regressions conducted (2 metaregressions per meta-analysis), in order to contain the false positive rate. Hence, voxels with AES-SDM z-values corresponding to p-values < 0.0005 (0.001/2) were considered significant, but voxels with AES-SDM z-values < 1, or in clusters with less than 10 voxels, were discarded in order to reduce the false positive rate.

#### 2.4.5. Accounting for MRI signal dropout in brain areas afflicted by the susceptibility artefact

The ventromedial prefrontal cortex (vmPFC) is considered an important area for encoding social information (Hiser and Koenigs, 2018). This area is known to suffer from distortion and signal dropout during fMRI scanning due to its proximity to air and bone around the sinuses (Sutton et al., 2009). To examine whether our analyses may have missed effects due to partial coverage of this area in at least some of the studies, we examined the whole-brain coverage of the studies included in each of our main meta-analyses. This was implemented by binarising each of our effect size maps, after registration to a common template, based on whether there was signal in each voxel or not and summing these images to create coverage density maps. For those studies where we used peak coordinates, we assumed there was no signal drop out and used a whole-brain binarized template mask (note we have no way of confirming whether an area was covered, since in these cases the effect size may be null for those voxels just because they are far away from the peaks used to reconstitute the effect size maps). Since we found a decline in coverage around the anterior and inferior edge of the vmPFC in our data, we used the method developed by Cutler and Campbell-Meiklejohn to perform an adjusted analysishy modifying the calculations run by AES-SDM to include only studies with data present on a voxel by voxel basis (i.e. for each voxel, the adjusted meta-analysis only includes a study if it sampled that specific voxel; if no signal was present at that voxel, then this study would have not been included in the meta-analytic calculations for that specific voxel). This would allow us to check whether we might have missed significant effects in a given area simply because some of our statistical maps did not contain information in these voxels. For a detailed description of the method, please see (Cutler and Campbell-Meiklejohn, 2019).

#### 2.4.6. Conjunction analyses

We conducted two sets of separate conjunction analyses. First, in order to identify areas commonly recruited by i) the anticipation of social rewards and punishments, ii) the receipt of social rewards and punishment, iii) the anticipation and receipt of social rewards (social reward processing), and iv) the anticipation and receipt of social punishments (social punishment processing), we created overlap maps between the thresholded and binarized maps of each correspondent contrasts (the resulting map represents a conjunction of all the voxels present in both meta-analytic maps). However, it should be noted that the number of studies included in the meta-analysis for each contrast was variable, and particularly unbalanced across the contrasts focusing on social reward and social punishment. It is possible that a certain voxel reaches significance in a specific meta-analytic contrast that includes a higher number of studies (e.g. such as the social reward versus neutral feedback anticipation contrast), while the same voxel does not reach significance in a different contrast simply because the number of contributing studies is smaller. While overlapping voxels can be interpreted with confidence, these differences in the number of contributing studies across contrasts preclude further inferences on areas of non-overlap or subtraction analyses.

Second, we wanted to gain insight about whether the brain areas identified in our meta-analytic maps for the social incentive delay task broadly mapped onto the brain areas identified in a similar previous analysis for the monetary incentive task that was used the AES-SDM approach (Wilson et al., 2018). Since Wilson et al. only focused on the anticipation period of the monetary incentive delay task, we created overlap maps between the meta-analytic maps for the anticipation phase of social rewards/punishments that we obtained in our study, and the meta-analytic maps for the anticipation phase for monetary rewards/punishments obtained in the previous study (Wilson et al., 2018). Similar comparisons could not be conducted for the receipt phase because there is no meta-analysis using AES-SDM in the case of rewards,
and we could not retrieve the final meta-analytic maps from the authors in the case of punishment (Dugre et al., 2018b)).

2.4.7. Labelling and atlases

We labelled each significant cluster using the Harvard-Oxford atlas in FSL (FMRIB Software Library, www.fmrib.ox.ac.uk/fsl).

3. Results

3.1. Included studies

Our literature search process resulted in the identification of a total of 104 studies. We identified four additional references through screening of the reference lists of the included studies. After removing duplicates, 57 studies remained. Thirty-three studies were excluded during title, abstract and key-words screening, resulting in 24 full-texts to be assessed for eligibility. From these 24 full-texts, we end up excluding six studies. Reasons for exclusion are detailed in our PRISMA flow diagram (Fig. 1). Our final pool of studies included 18 references. We contacted the lead, senior and/or corresponding authors from these 18 studies to obtain the raw statistical maps, of whom 12 responded. We excluded two additional studies for which we could not retrieve from the paper or have provided by the authors statistical maps or peak coordinates for any of the reported contrasts. Therefore, the final pool of studies that was included in our meta-analyses consisted of 16 studies.

The combined sample size across all 16 included studies was 502 participants, mean age 25.95 years old (SD 5.95), mean percentage of males 52.90 % (SD 33.70), 62.5 % of the studies included only right-handed participants, while for 37.5 % of the studies, information on participants’ handedness was not available. Out of the 16 studies used MRI data acquired at 3 T scanners – the remainder two studies used data from 1.5 T scanners. For two studies, we included data acquired after the intranasal administration of a placebo in the context of pharmacological oxytocin studies. Nine studies used outcome feedback in the form of static emotional faces, three studies in the form of dynamic videos depicting positive/negative emotional reactions to performance (happy or angry face with thumbs up or down) and four studies used verbal feedback (either auditory or written). See Table 1 for an overview of the data included in each of our four meta-analysis and table S1 for details on sociodemographics, paradigm specification, MRI data acquisition and analysis.

4. Meta-analytical findings

4.1. Anticipation of social reward

Sixteen-studies reported results on the contrast of anticipation of social reward versus neutral feedback anticipation. We obtained statistical maps from nine of the 16 studies, and used peak coordinate statistics for the remaining seven studies. This meta-analysis included data from the full combined sample of 502 participants.

Table 1

| Contrast                               | N datasets | N statistical maps | Pooled sample size |
|----------------------------------------|------------|--------------------|--------------------|
| 1. Anticipation of social reward       | 16         | 9                  | 502                |
| 2. Receipt of social reward            | 9          | 6                  | 272                |
| 3. Anticipation of social punishment   | 5          | 3                  | 129                |
| 4. Receipt of social punishment        | 4          | 3                  | 107                |

4.1.1. Main meta-analytic findings

We present a detailed description of all the clusters and respective peaks corresponding to increases and decreases in the BOLD signal during the cued anticipation of social rewards versus neutral feedback in Table S2. Briefly, we found increases in the BOLD signal in regions spanning: i) subcortical regions, including the striatum, amygdala and insula bilaterally, the thalamus; ii) cortical regions, including the right olfactory cortex, the left inferior frontal gyrus, the precentral gyrus and the supplementary motor area bilaterally, the left middle occipital gyrus; iii) the brainstem; and iv) the cerebellum (vermian lobule). We found decreases in the BOLD signal in a network of cortical regions, including the superior frontal gyrus bilaterally, the left middle frontal gyrus, the superior frontal gyrus bilaterally, the right postcentral gyrus, the middle temporal gyrus bilaterally, the left fusiform and angular gyri, the right middle occipital gyrus, the left median/paracingulate gyri, in the right parahippocampal gyrus, and in the left cerebellum (Fig. 2).

4.1.2. Sensitivity analysis

Our sensitivity analysis showed that our main meta-analytic findings are overall robust (Fig. S1).

4.1.3. Heterogeneity and publication bias

We found some areas of heterogeneity in the basal ganglia, thalamus, right insula, superior temporal gyrus, midbrain and supplementary motor area (Fig. S2). We found some evidence for publication bias in five cluster peaks showing increases in the BOLD signal. These clusters included the right supplementary motor area, the left insula, the right precentral gyrus and the left middle occipital gyrus (Table S2).

4.1.4. Metaregressions

We did not find any voxels where changes in BOLD were moderated by the mean age or the percentage of men in the included studies.

4.1.5. Subgroup analyses

When we repeated our main meta-analysis including only studies using emotional faces as outcome (Fig. S3), or including only studies using static emotional faces (Fig. S4), or including only non-pharmacological studies (Fig. S5), we found no difference in the pattern of our results, but only observed a decrease in the size or extent of our significant clusters.

4.2. Receipt of social reward

Nine studies reported results on the contrast social reward feedback versus neutral feedback. We obtained statistical maps from 6 of these studies, and used peak coordinate statistics for the remaining three studies. This meta-analysis included data from 272 participants.

4.2.1. Main meta-analytic findings

We present a detailed description of all clusters and respective peaks corresponding to increases and decreases in the BOLD signal during receipt of social reward versus neutral feedback in Table S3. Briefly, we found increases in the BOLD signal in regions spanning: i) cortical regions, including the frontal medial cortex, the anterior cingulate and frontal orbital cortices bilaterally, the middle temporal gyrus bilaterally, the superior frontal gyrus bilaterally, the posterior cingulate/precuneus bilaterally, the lateral occipital cortex bilaterally and the right occipital pole; ii) subcortical regions, including the amygdala and hippocampus bilaterally and the right thalamus; and in iii) the brainstem. We found decreases in the BOLD signal in the superior frontal gyrus bilaterally, the right frontal pole, the left postcentral gyrus, the central operculum cortex/insula and the precuneus bilaterally (Fig. 3).

4.2.2. Sensitivity analysis

Our sensitivity analysis showed that our main meta-analytic findings were overall robust (Fig. S6).
4.2.3. Heterogeneity and publication bias

We found some areas of heterogeneity mostly in the frontal medial cortex, the anterior cingulate cortex, the amygdala/hippocampus, the frontal orbital cortex, the occipital cortex and the occipital pole, and the opercular cortex/insula (Fig. S7). We found evidence for publication bias in four cluster peaks showing decreases in the BOLD signal. These clusters included the right median cingulate/paracingulate gyri, the left insula and the right inferior parietal gyri (Table S3).

4.2.4. Metaregressions

We did not find any voxels where changes in the BOLD signal were moderated by the mean age or the percentage of men in the included studies.

4.2.5. Subgroup analyses

When we repeated our main meta-analysis including only studies using static emotional faces (Fig. S8), or excluding studies administering placebo (Fig. S9), we found no substantial difference in the pattern of our results, other than an overall decrease in the size/extent of our significant clusters. This decrease was more pronounced when we excluded the studies administering placebo, particularly with respect to decreases in the BOLD signal.
4.3. Anticipation of social punishment avoidance

Only six studies reported results on the contrasts anticipation of social punishment avoidance versus neutral. We obtained statistical maps from three of these studies, and used peak coordinate statistics from two studies. This meta-analysis included data from 129 participants.

4.3.1. Main meta-analytic findings

We present a detailed description of all clusters and respective peaks corresponding to increases and decreases in the BOLD signal during the anticipation of social punishment avoidance versus neutral feedback in Table S4. Briefly, we found increases in the BOLD signal in a set of regions spanning: i) cortical regions, including the frontal orbital cortex bilaterally, the superior and middle frontal gyri bilaterally, the right lateral and fusiform occipital cortices, and the occipital pole bilaterally; ii) subcortical regions, including the amygdala, thalamus, septal nuclei, striatum and pallidum bilaterally, the right insula; and iii) the brainstem. We found decreases in the BOLD signal mainly in the paracingulate gyrus, the frontal pole bilaterally, the left precentral gyrus, the left middle/inferior temporal gyri and the left temporal pole, the supramarginal gyri bilaterally, the left parietal operculum, and the right cerebellum (Crus I) (Fig. 4).

4.3.2. Sensitivity analysis

Our sensitivity analysis showed that our main meta-analytic findings were overall robust (Fig. S10).

4.3.3. Heterogeneity and publication bias

We found only a few areas of heterogeneity mostly in the occipital poles (bilaterally), the thalamus and the left striatum (Fig. S11). We did not find evidence for publication bias in any of the reported peaks (Table S4).

4.3.4. Metaregressions

We did not find any voxels where changes in the BOLD signal were moderated by the mean age of the participants included. We found positive associations between the percentage of men included in the studies and the BOLD signal change in the occipital poles bilaterally, the right thalamus, the striatum/pallidum bilaterally, and the right frontal orbital cortex, and negative associations with the BOLD signal change in the medial frontal cortex and the left middle/inferior temporal gyri (Fig. S12).

4.3.5. Sub-group analyses

When we repeated our main meta-analysis excluding the study from (Nawijn et al., 2017), we found no considerable difference in the pattern of our results, but only observed a decrease in the size or extent of our significant clusters in the basal ganglia.

4.4. Receipt of social punishment

Four studies reported results on the contrast social punishment feedback versus neutral feedback. We obtained statistical maps from three of these studies, and used peak coordinate statistics for one study. This meta-analysis included data from 107 participants.

4.4.1. Main meta-analytic findings

We present a detailed description of all clusters and respective peaks corresponding to increases and decreases in the BOLD signal during receipt of social punishment feedback versus receipt of neutral feedback in Table S5. Briefly, we found increases in the BOLD signal in a network of regions spanning the frontal orbital cortex bilaterally, the right superior/inferior frontal gyri, the right frontal pole, the left frontal operculum/insula and the right lateral occipital cortex. We found decreases in the BOLD signal in areas spanning the frontal pole and the superior/middle frontal gyri bilaterally, the precuneus, the angular gyrus and in the basal ganglia (the left caudate and right putamen) (Fig. 5).

4.4.2. Sensitivity analysis

Our sensitivity analysis showed that our main meta-analytic findings were overall robust for increases in the BOLD signal. However, it also showed that the robustness of the decreases in the BOLD signal was attenuated, with voxels reported in our main meta-analysis reaching significance in no more than 50 % of the Jackknife leave-one-out meta-analyses (Fig. S13).

4.4.3. Heterogeneity and publication bias

We found some minimal areas of heterogeneity mostly in the right Fig. 4. Anticipation of social punishment avoidance. Meta-analytic results for the social punishment anticipation contrast (5 studies, pooled sample size 129 participants). Colour bars represent Z-SDM scores. In red, we present increases and in the blue decreases in BOLD signal for the contrast anticipation of social punishment avoidance versus anticipation of neutral feedback. Results were considered significant for p < 0.001, Z-SDM > 1 and cluster extent > 10 voxels as per current standard recommendations for multiple comparisons control using this method.
Fig. 5. Receipt of social punishment. Meta-analytic results for the social punishment receipt contrast (4 studies, pooled sample size 107 participants). Colour bars represent Z-SDM scores. In red, we present increases and in the blue decreases in BOLD signal for the contrast receipt of social punishment versus receipt of neutral feedback. Results were considered significant for $p < 0.001$, Z-SDM $> 1$ and cluster extent $> 10$ voxels as per current standard recommendations for multiple comparisons control using this method.

Fig. 6. Conjunction analysis. In this figure, we present the results of a conjunction analysis examining the overlap between our four meta-analytic maps for the following pairs: 1. Anticipation of social reward AND anticipation of social punishment avoidance; 2. Receipt of social reward AND receipt of social punishment; 3. Anticipation of social reward AND receipt of social reward; 4. Anticipation of social punishment avoidance AND receipt of social punishment. Overlaps are depicted as binarized voxels coloured in red.
4.6. Conjunction analyses to identify areas of overlap in the processing of midbrain and the amygdala bilaterally (Fig. 6A).

4.6.1. Anticipation of social rewards and punishments

The contrasts for the anticipation of social rewards and punishment avoidance (versus the anticipation of neutral feedback) overlapped in a group of brain areas that included the caudate nucleus, putamen and pallidum (bilaterally), the thalamus (bilaterally), the frontal medial cortex (bilaterally), the right insular cortex, and to a small extent the midbrain and the amygdala bilaterally (Fig. 6A).

4.6.2. Receipt of social rewards and punishments

The contrasts for the receipt of social rewards and punishments (versus the receipt of neutral feedback) overlapped in the precuneus (bilaterally), the right lateral occipital cortex (right), and the frontal pole/orbitofrontal cortices (bilaterally) (Fig. 6B).

4.6.3. Anticipation and receipt of social rewards

The contrasts for the anticipation and receipt of social rewards (versus the anticipation and receipt of neutral feedback, respectively) overlapped in the frontal medial cortex (bilaterally), the anterior (left) and middle (right) cingulate cortex, the thalamus (bilaterally), the posterior cingulate (bilaterally) and to a small extent in the left amygdala, the lateral occipital cortex (bilaterally), the superior temporal gyrus (bilaterally), the left precentral/postcentral gyri, and the right superior frontal gyrus (Fig. 6C).

4.6.4. Anticipation and receipt of social punishments

We did not find any voxel where the contrasts for the anticipation and the receipt of social punishments (versus the anticipation and receipt of neutral feedback, respectively) overlapped (Fig. 6D).

4.7. Conjunction analyses to identify areas of overlap in the anticipation of social and monetary rewards

4.7.1. Anticipation of social and monetary rewards (versus neutral feedback)

Overall, we found little evidence that the anticipation of social punishment recruits modality-specific brain areas when qualitatively compared to the anticipation of monetary punishments (with the exception of the right cerebellar hemisphere – Crus I and the occipital poles). The brain areas we identified to be involved in the anticipation of social punishment avoidance broadly mapped onto the same anatomical regions identified in (Wilson et al., 2018) to be involved in the anticipation of monetary loss avoidance. It should be noted that in some cases the identified clusters were contiguous rather than overlapping exactly (Fig.S18).

5. Discussion

We present the results of the first comprehensive voxel-based meta-analysis using Anisotropic Effect Size Signed Differential Mapping (AES-SDM) to map the brain regions involved in the anticipation and receipt of performance dependent, probabilistic social rewards and punishments in the human brain using the MID task. All four meta-analytic maps can be freely downloaded from Neurovault (https://neurovault.org/collections/7793). We identify brain areas missed in individual studies due to a lack of power, including regions of the default-mode network that show decreases in the BOLD signal during the anticipation of both social rewards and punishment avoidance. Additionally, we characterise the effect size and direction of changes in the BOLD signal for each brain area. By uncovering brain regions missed in individual studies, our meta-analyses enhance our understanding of the neural circuits underlying social incentive processing in healthy humans. Furthermore, by providing a well-defined set of regions engaged by the anticipation and receipt of social rewards and punishments in healthy individuals, our results can inform targeted hypotheses in future studies investigating dysfunctional processing of social rewards and punishments during disease and its modulation by innovative potential treatments. We discuss each of our main findings below.

5.1. Brain regions involved in the anticipation of social rewards and punishment avoidance

Using a voxel-based meta-analytic method, we identified an extensive network of brain areas involved in the anticipation of social rewards. Our results consolidate findings from a previous coordinate-based meta-analysis (ALE) regarding the involvement of the basal ganglia, the midbrain, the dorsal anterior cingulate cortex, the supplementary motor area, the anterior insula and the occipital gyrus (Gu et al., 2019). However, thanks to the enhanced sensitivity of the AES-SDM meta-analytic method that we used, our results extend this network by including frontal, temporal, parietal and cerebellar regions that were not captured in the previous coordinate-based meta-analysis. Furthermore, for the first time, we identified the involvement of a robust network of brain regions typically regarded as part of the default-mode network, namely the posterior cingulate cortex, angular gyrus, inferior parietal lobe and medial prefrontal cortex (Alves et al., 2019). Specifically, we found that these brain regions showed a relative decrease in the BOLD signal during the anticipation of social reward, compared to neutral feedback. In a similar previous AES-SDM meta-analysis of fMRI data from the MID task, the authors also identified a network of brain regions (largely overlapping with the brain region we identified here) showing decreases in the BOLD signal during the anticipation of monetary rewards and losses (Wilson et al., 2018) that had not been identified in previous coordinate-based meta-analyses (Gu et al., 2019; Oldham et al., 2018b). Compared to coordinate-based approaches (i.e. ALE) which assess the convergence of foci reported from available experiments, AES-SDM is a mixed method which can combine reported peak information with original statistical parametric maps. The possibility of including original statistical maps increases sensitivity and
allows to address issues of low statistical power by preserving information from voxels that did not reach significance in the original studies because of power limitations (Radua and Mataix-Cols, 2012). This gain in sensitivity increases with the number of statistical parametric maps included (Radua et al., 2012a). In all of our meta-analyses, we gathered statistical parametric maps from at least 50% of the eligible studies. Hence, our results neatly illustrate some of the advantages of the AES-SDM meta-analytic approach.

Furthermore, our study provides the first meta-analysis of neuroimaging data focusing on the contrast comparing the anticipation of social punishment avoidance to the anticipation on neutral feedback. We identified a robust network of brain regions showing increases and decreases in the BOLD signal similar to that involved in the anticipation of social rewards. Our conjunction analyses suggested that the anticipation of both social rewards and punishment avoidance recruits a set of well described circuits typically known as the “reward system” (Arias-Carrion et al., 2010). This observation lends support to the idea that most these areas may be part of general motivational system linked to the anticipation of highly salient incentive stimuli independent of valence or incentive-type (i.e. social or monetary). Consistent with this idea, previous findings from an ALE meta-analysis also suggested that the same network of regions are involved in the anticipation of monetary gains and losses during the MID task (Oldham et al., 2018b). However, given that our social reward and punishment avoidance anticipation meta-analyses were highly unbalanced in terms of the number of included studies, which did not allow us to conduct subtraction analyses, we should not exclude that, at least, some of the areas we identified in each of these two meta-analyses might be specifically engaged by one process or the other.

Below we discuss some of the identified brain regions that are considered to play a key role in incentive processing. We identified increases in the BOLD signal during the anticipation of social rewards and punishment avoidance in the striatum, the anterior insula, the thalamus, the anterior cingulate, the supplementary motor cortex and the amygdala. During the anticipation phase of the SID task, participants are presented with a task-relevant cue that indicates one of a range of reward, punishment or neutral stimuli that may follow the response to the target depending on performance. This phase of the task entails a complex cognitive machinery which ultimately allow participants to: i) direct their attention to a salient task-relevant stimulus; ii) make predictions about the probabilities of one of the possible outcomes based on the anticipatory cue; iii) prepare an effective motor program that will allow them to respond to the target as quickly as possible, in order to obtain a reward or avoid a punishment. We argue the set of regions we found during the anticipation of both social rewards and punishment avoidance are broadly compatible with the cognitive processes engaged by this phase of the task. For instance, accumulating evidence associates the ventral striatum (and the mesolimbic dopamine pathway) to motivational processing independent of stimulus valence (Brooks and Berns, 2013; Lammel et al., 2014). Furthermore, many studies have suggested that the striatum plays a key role in computations that take place during social behaviour, including social reward prediction and learning - for an extensive review please see (Baez-Mendoza and Schultz, 2013). At the moment, it is still under debate whether the striatum solely responds to salience or might encode both salience and valence of a stimulus (Sarrira et al., 2013). In line with the first idea, we found increases in BOLD signal during anticipation of both social reward and punishment avoidance. However, it is also possible that it also exhibits a double pattern because during punishment blocks participants mostly anticipate punishment avoidance (which in most task setups is predefined to be the most frequent outcome) rather than punishment itself. While punishment has negative valence, punishment avoidance works as negative reinforcer which has positive valence. This hypothesis needs further exploration in future studies.

The anterior insula, which has dense interconnections with the striatum (Ghaziri et al., 2018), has been implicated in encoding outcome uncertainty (Gorka et al., 2016), which is a key process of the anticipation phase of the SID design (given that outcome receipt is probabilistic and dependent on performance). Furthermore, the insula is a key node of the salience network (Menon and Uddin, 2010) and processes ascending interoceptive and visceromotor signals (Ronchi et al., 2015). The insula is also regarded as a central area involved in emotional processing and social cognition (Couto et al., 2013), including the detection of salience of social stimuli or events (Chen et al., 2009; Feng et al., 2015; Luo et al., 2018). Therefore, the involvement of the insula during the anticipation of social rewards and punishment avoidance in the SID task is compatible with an overall role of the insula in the autonomic activation (arousal) associated with the processing of salient cues signalling the probabilistic receipt of one of three possible social outcomes, depending on performance (Schneider et al., 2018). The thalamus encodes an alerting signal to respond to salient stimuli (Wolff and Vann, 2019; Zhu et al., 2018) by conveying interoceptive information from the insula to the striatum, where an appropriate action response is then selected (Huang et al., 2018). Interestingly, social cognition impairments have been reported in stroke patients with unilateral thalamic lesions (Wilkos et al., 2015). Hence, together with the insula and the striatum, the thalamus might be part of the network of brain regions that work together to encode the salience of the task-relevant anticipatory cue participants are exposed to during this phase of the task.

Given that the SID is a motor response time task, the presence of the supplementary motor cortex, typically recruited during movement planning and control (Tanji, 1994), is also consistent with the characteristics of the task. Additionally, recent studies have shown that the motor and premotor cortices also encode reward signals related to both the anticipation and receipt of a reward (Ramakrishnan et al., 2017; Ramkumar et al., 2016). The dorsal cingulate is a region that subserves both cognition and motor control (Beckmann et al., 2009) and a plethora of studies have implicated this area in processes such as attention for action/target selection (Haken and Plott, 2010; Isomura et al., 2003), motivation (Monosov, 2017), motor response selection (Badgaiyan and Posner, 1998), performance monitoring (Gehring and Knight, 2000), novelty detection (Hayden et al., 2011) and social cognition (namely, in tracking others’ motivation) (Apps et al., 2016). Most of these processes are part of the cognitive processes engaged during the anticipation phase of the SID task. Lastly, the amygdala, which has been classically linked with negative emotional processing, has more recently been proposed to respond to stimulus salience/arousal rather than valence (Bonnet et al., 2015; Fastenrath et al., 2014; Zheng et al., 2017). In addition, the amygdala has also been proposed to be instrumental during the processing of emotional and socially relevant information, including the processing of emotions from faces which are often used as a feedback stimulus in the SID task (Todorov, 2012b).

For the first time, we also identified a group of brain regions showing decreases in the BOLD signal during the anticipation of both social rewards and punishment avoidance. These regions are part of what is commonly described as the default-mode network (Smith et al., 2009). There has been debate about the exact functions of the default-mode network (Crittenden et al., 2015; Raichle, 2015). Current views subdivide this network into a core subsystem, anchored in the posterior cingulate cortex and the anteromedial prefrontal cortex, which has been associated with self-referential processes; a medial temporal subsystem related to the processing of past and future autobiographical thoughts; and a ventral medial subsystem involved in the processing of reward to social cognition, story comprehension and semantic processing (Salomon et al., 2014). Importantly, the default mode network has been associated with decreases in the BOLD signal during external-oriented active tasks (Anticevic et al., 2012), thus leading some researchers to label the network as a task-negative network (this conceptualization of the default-mode network as a task-negative network has nevertheless been recently contested - see Spreng, 2012). The anticipation phase of the SID task arguably involves a range of cognitive processes, including
saliency processing, working memory (holding the cue-signalled state in mind), focused attention and motor preparation (participants are preparing to react as fast as possible when the target appears, as success is performance dependent) and predictions about potential outcomes. Therefore, the observed pattern of decreases in the BOLD signal is consistent with the idea that the brain should disengage from self-oriented processing to attend to and prepare to respond to relevant external stimuli (Scheibner et al., 2017). This idea is further supported by the activation of key nodes of the salience network such as the insula and the anterior cingulate cortex that we reported and discussed above. These nodes are typically assumed to participate in the switch between self- and external-oriented processing (Corbetta and Shulman, 2002).

One key question in the field regards the extent to which the processing of social rewards/punishments involves additional specific brain regions (Gu et al., 2019; Izuma et al., 2008) that are typically not engaged during the processing of other types of rewards/punishments (i.e. monetary). One prominent hypothesis states that the anticipation and receipt of social rewards/punishments engages both a generalist neural network that consists of brain regions involved in the processing of rewards/punishments irrespective of their type (such as the basal ganglia), and a specialist network of regions specifically involved in the processing of social information (such as the dopaminoceptive junction, the dorsomedial prefrontal cortex, the precuneus, and the superior temporal gyrus) (Barman et al., 2015a; Goerlich et al., 2017a; Spreckelmeyer et al., 2013a). This hypothesis has been challenged though by at least two recent meta-analyses, the results of which were consistent with the idea that a general-purpose brain system is involved in the anticipation of rewards/punishments irrespective of their type. Specifically, the first meta-analysis showed that the circuits engaged during the anticipation of monetary gains also encompass some of the brain areas often attributed to social information processing (Wilson et al., 2018). The second meta-analysis compared the anticipation of monetary and social rewards and failed to find any significant differences in the underpinning brain regions (Gu et al., 2019).

To help illustrate this question, we conducted an exploratory conjunction analysis between the meta-analytic effect size maps on the SID task from our study and meta-analytic effect size maps on the MID task that we obtained from another study that used the same AES/SDM meta-analytic method (Wilson et al., 2018). We only focused on the anticipation period as the previous study had not analysed data from the outcome phase of the MID task. Overall, we found little evidence of modality-specific processing, supporting the idea that, at least regarding the anticipation of rewards/punishments, a general-purpose anticipation system is recruited by both monetary and social rewards/punishments. However, in some cases we noted the recruitment of voxels across tasks that may have belonged to the same brain regions but were anatomically contiguous rather than overlapping. It is therefore tempting to speculate that rather than recruiting different neural systems, the anticipation of social and monetary rewards/punishments may be functionally encoded in the same regions but engage different neuronal ensembles. Indeed, something similar has been demonstrated for the brain encoding of physical and social rejection, which are characterized by differential multivariate voxel patterns despite common fMRI activity at the gross anatomical level (Woo et al., 2014). This is a hypothesis that should be addressed in future studies combining the application of both tasks in the same individuals with current state-of-art multivoxel pattern recognition techniques to ascertain whether modality can be decoded by a classifier trained on brain voxel-based responses (Corradi-Dell’Acqua et al., 2016; Mahmoudi et al., 2012).

5.2. Brain regions engaged during the receipt of social rewards and punishments

Our study provides the first meta-analysis of neuroimaging data focusing on the contrasts comparing the receipt of social rewards or punishments to neutral feedback. We identified a robust network of brain regions engaged during the receipt of social rewards, including the ventromedial frontal (vmPFC) and orbitofrontal cortices, the anterior cingulate cortex, the amygdala, the hippocampus, the occipital cortex and the brainstem. During the receipt phase of the SID task, participants are exposed to a positive, negative or neutral social stimuli. The frequency at which participants are exposed to each of these outcomes depends on their performance during a reaction time target selection motor task they performed immediately before. During this phase of the task a number of cognitive computations are thought to take place, such as: i) recognition of the sensorial features of the presented outcome; ii) informed by the previous, encoding of the subjective value of the outcome received; iii) computations of discrepancy between predictions about the outcome and the actual received outcome; iv) encoding of associative relationships between anticipatory cue, action and outcome. Therefore, we argue the brain regions we found for our receipt contrasts are broadly compatible with the implementation of these cognitive processes. Below, we will discuss in detail how each of the brain regions we found may map to each of these functions.

Increases in BOLD signal in the ventromedial prefrontal cortex and the orbitofrontal cortex have been consistently associated with the receipt of monetary outcomes (Oldham et al., 2018b). Previous work has suggested that the vmPFC and the orbitofrontal cortex encode subjective value (Piva et al., 2019) and the associative relationship between stimuli and outcome (de Wit et al., 2009). Furthermore, increasing evidence has implicated the vmPFC in multiple aspects of social cognition, such as facial emotion recognition, theory-of-mind ability, and the processing self-relevant information - for an extensive review and meta-analysis see (Hiser and Koenigs, 2018). Furthermore, we also observed increases in the BOLD signal in the amygdala, the brainstem, the anterior cingulate cortex, and the occipital cortex. The amygdala has been proposed to participate in the representation of the utility or affective value during monetary incentives receipt (Ernst et al., 2005; Hampton et al., 2007). According to this model, the utility encoded by the amygdala is thought to be then used to inform behavioural and physiological responses towards and away from positive and negative stimuli, respectively. This process has been suggested to include dispatches of the information encoding the affective value of stimuli from the amygdala to different brain systems, including the arousal circuits of the brainstem (Ernst et al., 2005; Hampton et al., 2007). Increased BOLD signal in the amygdala, the anterior cingulate cortex, and the occipital cortex is also a consistent finding of studies investigating brain responses to emotional faces (Fusar-Poli et al., 2009; Todorov, 2012a), which were the most frequently used outcome stimuli in the receipt phase of the SID tasks we included in our meta-analyses.

One notable absence was the lack of evidence for the involvement of the basal ganglia during the receipt of social rewards. In contrast, previous meta-analytic evidence has shown that the ventral striatum is engaged during the receipt of monetary rewards in the MID task (Oldham et al., 2018b) or in response to agreement with normative opinions during social conformity (Wu et al., 2016). Even though the MID and the SID tasks are not classical learning tasks, the ventral striatal response to uncertain monetary rewards has been commonly interpreted in the reinforcement learning literature as the neural correlate of the dopaminergic neurons coding discrepancy between expected and actual outcomes (prediction error) (Lockwood et al., 2016, 2020a; Lockwood and Klein-Flugge, 2020; Nasser et al., 2017; Wittmann et al., 2018). Similar prediction errors have been reported to occur in the striatum during social learning (Joiner et al., 2017). Therefore, we would have expected to have found an increase in the BOLD signal in the ventral striatum during the receipt of social rewards. Indeed, an increase in the BOLD signal in the ventral striatum was noted in some of the leave-one-out meta-analyses that were part of the sensitivity analysis; however, this increase was not consistent enough to be captured in the main meta-analysis. Ultimately, this could have resulted from the fact that most studies used high success rates (~66.6%). These high success rates might have biased the expectations of participants about the
outcomes towards high levels of certainty. According to the prediction error theory presented above, if participants were expecting to be rewarded with high levels of certainty and the outcome matched participants’ expectations about reward outcome most of the times, then one would expect the BOLD response in the basal ganglia to be reduced. While the same principle would apply for the MID task, we note that the number of studies included in previous MID meta-analyses was considerably higher than those of the studies using its SID counterpart we included here. Hence, it is possible we might have not had enough power to detect smaller changes in the BOLD signal in the basal ganglia during the receipt of social rewards in our meta-analysis.

Regarding the brain regions engaged during the receipt of social punishments, we identified a network of brain regions showing increases in the BOLD signal, including the orbitofrontal cortex, the superior and inferior frontal gyri, the lateral occipital cortex and the insula. Additionally, and in contrast to the receipt of social rewards, we also observed a decrease in the BOLD signal in the basal ganglia in response to the receipt of social punishments (compared to neutral feedback). This finding is in line with previous evidence that monetary loss, as compared to neutral feedback, results in a decrease in the BOLD signal in the striatum (in contrast to monetary gain, which results in an increase in the BOLD signal) (Delgado et al., 2000). Interestingly, one previous meta-analysis of the BOLD fMRI correlates of social conformity also found consistent decreases in the BOLD responses of the ventral striatum when people’s responses deviate from group opinions (Wu et al., 2016).

The integrated analysis of ours and these previous findings on social conformity suggests that BOLD decreases in the ventral striatum during disagreement with others might represent a punishment error signal to facilitate subsequent conforming behaviour. The sensitivity analyses indicated that we should consider the meta-analytic results from this contrast with caution, as they indicated lack of robustness (that is, the reported findings could be driven by the specific combination of studies). This is not unexpected given the small number of studies (N = 4) included in the meta-analysis for this contrast.

Our conjunction analysis identified clusters that were commonly engaged during the receipt of both social rewards and punishments. These clusters extended over the lateral occipital cortex, the precuneus, the cuneal cortex, the frontal pole and the orbitofrontal cortex. This observation is compatible with the hypothesis that certain regions may be engaged in shared sensory/cognitive processes irrespective of the valence of the feedback, and in encoding the subjective relevance of the feedback which does not necessarily depend on its valence (Oldham et al., 2018b).

5.3. Publication bias and heterogeneity

Very few neuroimaging meta-analyses consider the issues of publication bias and heterogeneity, which are commonly addressed in behavioural meta-analyses (Muller et al., 2018; Radua and Mataix-Cols, 2012). The consideration of publication bias in neuroimaging meta-analysis such as the ones we present here calls for reflection on one main issue. Voxels whose effect may have failed to survive multiple comparisons in individual studies would have been assigned an estimated effect size of zero when only peak coordinates are available, compared to approaches using whole brain statistical maps. With this in mind, we conducted and present the results of a publication bias analysis using both the inspection of funnel plots and the Egger’s test on the effects extracted for each peak of our four meta-analyses. We could only find some indication for publication bias for some of the peaks identified in the meta-analysis of the anticipation of social reward versus anticipation of neutral feedback contrast. For these peaks, the inspection of the respective funnel plots indicated the lack of smaller studies reporting small effect sizes for these peaks. Therefore, the findings regarding these peaks should be taken with caution. However, we should note that none of these peaks would have shown publication bias if we had applied statistical correction in our publication bias analyses for the number of peaks examined. Regarding the issue of methodological heterogeneity across the studies included in the reported meta-analyses, we found evidence suggesting the existence of considerable methodological heterogeneity in the meta-analytic maps for the anticipation of social reward or punishment avoidance versus the anticipation of neutral feedback contrast, and the receipt of social rewards versus neutral feedback contrasts. For the anticipation of social punishment avoidance versus the anticipation of neutral feedback contrast, this heterogeneity is likely to be explained by differences in the percentage of men included across studies, as suggested by our exploratory metaregression analyses using the percentage of men included in each study as predictor. This finding is largely in line with previous studies showing gender-related differences in the neurophysiological underpinnings of social incentive processing (Greimel et al., 2018; Spreeckmeyer et al., 2009; Wang et al., 2017). For instance, one study in adolescents showed that during the anticipation of potential social punishment, adolescent boys, compared with girls, exhibited a reduced stimulus-preceding negativity (Greimel et al., 2018). For the remaining contrasts, we found that neither differences in the mean age nor in the percentage of men could explain the observed heterogeneity. However, it should be noted that the inclusion of both statistical maps and coordinate-based data in AES-SDM may inflate heterogeneity estimates (Radua et al., 2012a).

5.4. Computational models in social incentive processing research: a way forward

The last decades of research in the field of the neuroscience of learning and decision making have witnessed an expansion of the number of studies combining computational modelling of behaviour with neuroimaging (Cohen et al., 2017; Corrado and Doya, 2007). Certain computations underlying our decisions, such as reward prediction errors or subjective values, are computed by our brains but often cannot be directly measured from participants’ responses to a task (Cohen et al., 2017; Corrado and Doya, 2007). Computational modelling offers an alternative to access these hidden quantities (Corrado and Doya, 2007). Computational models formalize cognitive processes in terms of precise mathematical algorithms that generate behaviour. Hence, fitting a computational model to participants’ responses in a task allows for a precise reconstruction of the trajectories of a number of neurocognitive processes underlying such responses (Cohen et al., 2017). These hidden variables or model parameters can in turn be regressed against a measure of brain activity during task performance, such as BOLD fMRI signal, giving insights into whether and where in the brain these computations are implemented (Cohen et al., 2017).

More recently, this type of computational or model-based neuroimaging experiments have also been conducted in the context of social neuroscience experiments to better understand the signals computed by the brain during social interactions - for in-depth reviews please see (Charpentier and O’Doherty, 2018; Dunne and O’Doherty, 2013; Suzuki and O’Doherty, 2020). Model-based fMRI studies on social feedback processing, such as social rewards and punishment, have proven that computational models of social reinforcement can be a powerful tool to dissect several aspects of social learning and decision making, for instance how we learn from social rewards and punishments (Jones et al., 2011; Will et al., 2017), how we learn to benefit (Lockwood et al., 2016) or avoid harm to others (Lockwood et al., 2020b), or even how others influence our own decisions (Zhang and Glaescher, 2020). While none of the SID studies we reviewed here employed computational modelling, one recent study employed a Rescorla–Wagner model of Reinforcement Learning to access trial-by-trial brain representations of subjective expected probabilities of gaining (anticipation phase) and prediction errors (outcome phase) during the MID task (Cao et al., 2019). This study showcases well how computational modelling can be applied in the context of an incentive processing task as simple as the MID or the SID tasks. As a way forward, future SID studies should consider including a similar model-based analytic approach in addition
to the conventional categorical analyses of anticipation and receipt.

5.5. Limitations

Our study has some limitations we should acknowledge. First, we note the relatively small number of studies investigating the processing of social punishment using the SID task. This aspect limits the power of our meta-analyses for the anticipation and receipt of social punishment contrasts. Nevertheless, we decided to proceed with the meta-analyses of these contrasts since we were able to retrieve statistical maps for more than half of the eligible studies. In fact, the pooled sample size of our social punishment meta-analyses exceeded 100 individuals, making it the largest dataset to date investigating this question. Given the role that differences in the processing of social punishments may play in a wide range of neuropsychiatric disorders (such as major depressive disorder (Kumar et al., 2017b) or antisocial personality disorder (Gong et al., 2019)), it seems imperative that future studies start to invest more in investigating the neural underpinnings of social punishment processing in both healthy and clinical samples. Second, our findings are limited by the reported sample demographics. Across all included studies, we were only able to systematically retrieve information on age and gender. While we did explore the impact of mean age and percentage of men using meta-regression, other potentially interesting variables such as substance use, education, intelligence quotient, socioeconomic status and ethnicity could not be consistently retrieved to allow for a thorough investigation of their impact. Third, our results may not generalize to the processes involved in the anticipation or receipt of social rewards or social punishments in contexts outside of the SID task, e.g. in neuroeconomic games involving the exchange of social rewards and punishments between individuals (such as the Prisoner’s dilemma (Sun et al., 2016)), indirect gains in social status or hierarchy (Gil et al., 2013; Munuera et al., 2018), or other tasks where feedback is not performance dependent (Hsu et al., 2018). Indeed, while designed to allow to disentangle brain processing associated with the phases of anticipation and receipt of social rewards and punishments in the lab, the SID task is a rather artificial setup that does not possess many of the characteristics social reward and punishment processing would entail in a naturalistic setting (i.e. interpersonal interaction, sensorial multimodality, direct impact of our own actions on others) (Reader and Holmes, 2016). This aspect limits the generalisability of the task itself and calls for further studies investigating what type of stimuli may better capture the social complexity of human interactions. Fourth, the cut-off date the studies that were included in our meta-analyses were identified was about 2 years ago, which raises the question of whether our meta-analyses might be missing a considerable number of fMRI studies using the SID task published in the past 2 years. To address this concern, we conducted a new literature search on 17/04/2020 which identified only a single study meeting our inclusion/exclusion criteria (He et al., 2019). We are thus confident that our meta-analyses still represent the vast majority of the published studies in this field.

Finally, we note that there was considerable heterogeneity across the various implementations of the SID task. For example, studies varied in terms of using static faces, dynamic faces or verbal feedback as the social stimulus, in terms of using scrambled faces, dysmorphic faces, or simple win/no-win symbols or landscape images in the neutral feedback condition. This heterogeneity can be both a blessing and a curse. This may result in a range of different neurocognitive processes experienced during anticipation and outcome presentation, including emotional responses (e.g. frustration), perception of self-efficacy, and even feelings of acquired learned helplessness driven by the enduring repeated experience of punishment that might be perceived to be beyond the participants’ own control (Wanke and Schwabe, 2019). We believe that future studies need to systematically investigate the impact of this aspect of heterogeneity at the neural level, as understanding the brain circuitry involved in the processing of negative reinforcers and punishments (as opposed to rewards) may be particularly useful for a range of neuropsychiatric disorders, such as depression.

6. Conclusion

This is the first voxel-based meta-analysis mapping the brain regions involved in the anticipation and receipt of social rewards and punishments in the human brain, as captured by the SID task. We identify brain areas missed in individual studies due to a lack of power, such as decreases in the BOLD signal in areas that are part of the default-mode network during the anticipation of both social rewards and punishment avoidance. We also characterise the effect size and direction of changes in the BOLD signal for each brain area. Qualitative anatomical comparisons showed little evidence supporting the involvement of domain-specific brain areas during the anticipation of social rewards/punishments, lending support to the hypothesis that a shared neural circuit underpins the anticipation of rewards/punishments irrespective of their type (i.e. social versus monetary). We noted the scarcity of studies focusing on the processing of social punishments despite the importance of this condition for several neuropsychiatric disorders. Our results provide a stereotaxic set of brain regions which could serve as regions-of-interest for future hypothesis-driven research seeking to investigate further how the human brain processes social rewards/punishments and how the disruption of these processes might contribute to the social dysfunction observed across many neuropsychiatric disorders. Ultimately, this knowledge may help us to identify potential target circuits that may be modulated by therapeutic interventions aiming to restore dysfunctional social incentive processing during disease.

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CRediT authorship contribution statement

D. Martins: Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Project administration, Writing - original draft, Writing - review & editing. L. Rademacher: Resources, Writing - review & editing. A.S. Gabay: Conceptualization, Methodology. R. Taylor: Investigation, Writing - review & editing. J.A. Richey: Resources, Writing - review & editing. D.V. Smith: Resources, Writing - review & editing. K.S. Goerlich: Resources, Writing - review & editing. L. Nawijn: Resources, Writing - review & editing. H.R. Cremer: Resources, Writing - review & editing. R. Wilson: Resources, Writing - review & editing. S. Bhattacharyya: Resources, Writing - review & editing. Y. Paloyelis: Conceptualization, Funding acquisition, Project administration, Supervision, Writing - original draft, Writing - review & editing.
Declaration of Competing Interest

The authors report no declarations of interest.

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Appendix A. Supplementary data

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