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**Neurochemistry of response inhibition and interference in gambling disorder: A preliminary study of  $\gamma$ -aminobutyric acid (GABA+) and glutamate-glutamine (Glx)**

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### Abstract

**Objective.** Neurobehavioural research on the role of impulsivity in gambling disorder (GD) has produced heterogeneous findings. Impulsivity is multifaceted, with different experimental tasks measuring different sub-processes, like response inhibition and distractor interference. Little is known about the neurochemistry of inhibition and interference in GD.

**Methods.** We investigated inhibition with the Stop Signal Task (SST) and interference with the Eriksen Flanker Task, and related performance to metabolite levels in individuals with and without GD. We employed magnetic resonance spectroscopy (MRS) to record glutamate-glutamine (Glx/Cr) and inhibitory,  $\gamma$ -aminobutyric acid (GABA+/Cr) levels in the dorsal ACC (dACC), right dorsolateral prefrontal cortex (dlPFC), and an occipital control voxel.

**Results.** We found slower processing of complex stimuli in the Flanker task in GD ( $p < .001$ ,  $\eta^2_p = .78$ ), and no group differences in SST performance. Levels of dACC Glx/Cr and frequency of incongruent errors were correlated positively in GD only ( $r = .92$ ,  $p = .001$ ). Larger positive correlations were found for those with GD between dACC GABA+/Cr and SST Go error response times ( $z = 2.83$ ,  $p = .004$ ) as well as between dACC Glx/Cr and frequency of Go errors ( $z = 2.23$ ,  $p = .03$ ), indicating general Glx-related error processing deficits. Both groups expressed equivalent positive correlations between post-error slowing and Glx/Cr in the right dlPFC (GD:  $r = .74$ ,  $p = .02$ ; non-GD:  $r = .71$ ,  $p = .01$ ).

**Conclusion.** Inhibition and interference impairments are reflected in dACC baseline metabolite levels and error processing deficits in GD.

45

46 **Introduction**

47 Gambling disorder (GD) is a psychiatric condition characterised by irritability and  
48 failing to stop gambling, recurrent thoughts about gambling and gambling as a coping  
49 mechanism, loss chasing, and hiding gambling behaviours from others or exploiting others  
50 for gambling money<sup>1, 2</sup>. The health-harming behaviours indicative of GD are now widely  
51 recognised as a public health issue<sup>3, 4</sup>.

52 Gambling disorder has long been associated with deficits in self-reported  
53 impulsivity<sup>5, 6</sup> and impaired task performance on behavioural indices of impulsive  
54 behaviour<sup>7, 8</sup>. A range of cognitive-behavioural domains have assessed the broad construct  
55 of impulsivity such as attentional inhibition, motor inhibition, discounting, decision-making,  
56 and reflection impulsivity<sup>9</sup>. As a result, observed deficits are heterogeneous across studies  
57 and individuals and warrant further investigation into the contribution of different  
58 impulsivity-related subprocesses in GD<sup>5, 10</sup>. This may include, for example, inhibitory control  
59 understood in terms of prepotent response inhibition and resistance to interference from  
60 distractors<sup>11, 12</sup>. Disentangling the separate and/or combined influence of specific  
61 impulsivity-related processes in GD might aid understanding of the various trajectories that  
62 lead to excessive gambling behaviour and enable future treatment development.

63 One subprocess, response inhibition or the ability to inhibit prepotent responses, is  
64 often assessed using the Stop Signal Task (SST)<sup>13</sup>. In the SST, a manual button press is  
65 required on most trials upon visual presentation of an arrow. The minority of arrow  
66 presentations are followed by an auditory stop signal, indicating the requirement to  
67 withhold the prepotent button press. Importantly, the time at which the auditory stop signal

68 is delayed in respect to the visual arrow, the stop signal delay (SSD), is adjusted in a  
69 stepwise manner, which computes the stop signal response time (SSRT).

70 A second subprocess, response interference or resistance to interference from  
71 distractors, is commonly assessed using the Eriksen Flanker task<sup>14</sup>. Like the SST, a central  
72 arrow is presented and the direction it faces determines the required button press. In the  
73 Flanker task, the central task-relevant stimulus is either flanked by congruent (C) or  
74 incongruent (IC) arrows and presentation of IC flankers induces response competition which  
75 increases response times and error rates<sup>14</sup>.

76 While response interference using the Flanker task has to our knowledge not yet  
77 been studied in populations with GD, previous research on problem gambling and response  
78 inhibition using the SST has produced mixed results. Inhibition-related variables, such as the  
79 SSD and SSRT, often do not dissociate between gambling and non-gambling participants<sup>15-20</sup>.  
80 Similarly, response time on Go trials fails to distinguish between those with and without  
81 GD<sup>21, 22</sup>. However, both prolonged response time and SSRTs are seen in participants with  
82 high gambling severity, whereas at-risk gamblers do not differ in their SST performance  
83 compared to non-gambling participants<sup>17, 23</sup>. Studies of response inhibition in GD show  
84 increased SSRTs with moderate to large effect sizes and increased Go response times during  
85 Go/No-go tasks with small to moderate effect sizes<sup>24</sup>. In addition to inhibition-related SST  
86 measures, only two SST studies, to date, have investigated error frequency and post-error  
87 slowing (PES) in gamblers. Lorains et al.<sup>18</sup> found enhanced error responses on Go trials in a  
88 sample of treatment-seeking gamblers, while Lawrence, et al.<sup>21</sup> found no differences to  
89 controls when investigating a moderate to severe disordered gambling sample. However, it  
90 is noteworthy that both studies also investigated the effects of previous trial types (correct

91 go, correct stop, failed stop) on current, within-session Go responses. Usually, behavioural  
92 responses that are preceded by an error are slower than behavioural responses that are  
93 preceded by correct trials, which might reflect an adaptive mechanism to reduce future  
94 errors or increased salience of errors<sup>25-27</sup>; however, both studies found no differences  
95 between gamblers and non-gamblers on PES.

96         Despite measuring purportedly different aspects of impulsivity, the neural networks  
97 recruited during the Flanker Task and SST overlap. Indeed, a recent activation likelihood  
98 estimation (ALE) meta-analysis compared the neural networks involved in cognitive  
99 inhibition, composed of Stroop and Flanker task data, to those involved in response  
100 inhibition, consisting of SST and Go/No-go tasks, and found overlap among task-based  
101 functional magnetic resonance imaging (fMRI) activity in dorsal anterior cingulate cortex  
102 (dACC), right-, but not left-hemispheric, dorsolateral prefrontal cortex (dlPFC) and the left  
103 anterior insula<sup>28</sup>. In a Go/No-go version of the Flanker task, interference-related dACC  
104 activation correlated positively with response times and error rates during the IC condition,  
105 while the number of inhibition errors correlated negatively with response inhibition-related  
106 activity in the right, but not left-hemispheric, dlPFC activity<sup>29</sup>.

107         Functional abnormalities of the overlapping brain areas supporting response  
108 inhibition and response distractor interference have been reported in disordered gambling.  
109 For example, a recent SST fMRI study assessed high-frequency poker players and revealed  
110 increased dACC activity during successful response inhibition compared to non-gambling  
111 controls in the absence of SSRT differences<sup>15</sup>. However, it is likely that gambling-related  
112 abnormalities in the dACC extend beyond neural activation. In previous work conducted  
113 with the present sample of males with GD, we showed that baseline glutamate-glutamine

114 (Glx) levels in the dACC negatively correlate with gambling severity<sup>30</sup>. This supports related  
115 findings showing that medication acting on glutaminergic transmission reduces gambling  
116 severity<sup>31, 32</sup>. It is noteworthy that optimal response inhibition and interference task  
117 performance is assumed to depend on optimally balancing excitatory and inhibitory  
118 neurometabolites, such as glutamate and  $\gamma$ -aminobutyric acid (GABA)<sup>33</sup>. Consistent with  
119 this, correlations between levels of these metabolites and behavioural performance have  
120 been reported previously. For instance, percentage of inhibition errors correlate negatively  
121 with GABA levels in the dACC<sup>34</sup>, as does self-reported impulsivity, which additionally  
122 correlates negatively with GABA+ (+ indicates contributions from unsuppressed  
123 macromolecules) levels in the right dlPFC<sup>35</sup>. Similarly, Chowdhury, et al.<sup>20</sup> reported a positive  
124 correlation between GABAergic transmission in the motor cortex and SSRTs. Interestingly,  
125 despite the absence of group differences in SST performance, Chowdhury, et al.<sup>20</sup> also found  
126 evidence for reduced GABA<sub>A</sub> receptor activity and increased glutamate receptor activity in a  
127 GD sample compared to non-gamblers and at-risk gamblers, respectively. Additionally,  
128 exogenous dopamine administration reduced prefrontal GABA<sub>A</sub> receptor availability less in  
129 treatment-seeking problem gamblers than in healthy volunteers<sup>36</sup>.

130 In terms of distractor interference, one previous investigation into the relationship  
131 between response time differences between IC and C Flanker trials and metabolite levels in  
132 the medial/dorsal ACC found no correlation with Glx, while GABA was unassessed<sup>37</sup>. Little is  
133 known, therefore, about the role of GABA in response interference, in GD compared or non-  
134 GD populations. It is possible, however, that GABAergic processes are involved during  
135 Flanker task interference control: Faßbender et al.<sup>38</sup> investigated the effects of Lorazepam, a  
136 benzodiazepine binding to the GABA<sub>A</sub> receptor and thereby enhancing GABA release, on

137 Flanker performance and reported increased error rates as well as response times when  
138 dosage was increased. On the other hand, performance on the related interference Stroop  
139 Task where the distracting stimulus dimension is dominant<sup>12</sup>, did not significantly correlate  
140 with glutamate or GABA in the dACC or parieto-occipital cortex<sup>34</sup>.

141 In sum, the existing evidence reveals conflicting findings on the range and type of  
142 impulsive deficits in GD, while little is known about the underlying neurochemistry of  
143 impaired response inhibition and interference. The present preliminary investigation  
144 therefore sought to undertake a combined behavioural and MRS study utilizing GD and non-  
145 GD samples. Baseline GABA+/Cr, and Glx/Cr were assessed in the dACC, right dlPFC (given  
146 its' role in Flanker as well as SST tasks<sup>28, 29</sup> and self-reported impulsivity<sup>35</sup>) and an occipital  
147 control voxel and then related to performance indices of response inhibition, using the SST,  
148 and distractor interference, using the Flanker task.

## 149 Methods

### 150 Participants

151 Twenty-six right-handed male participants were allocated into GD and non-GD  
152 groups based on their past year gambling severity scores on the *Problem Gambling Severity*  
153 *Index* (PGSI)<sup>39</sup>. This resulted in  $n = 12$  in the GD group (i.e., PGSI score  $> 8$ ;  $M = 15.2$ ,  $SD =$   
154  $5.1$ ;  $M_{\text{age}} = 36.3$ ,  $SD = 9.5$ ) and 14 age-matched, non-GD participants (i.e., PGSI score  $< 1$ ;  $M$   
155  $= .071$ ,  $SD = .027$ ;  $M_{\text{age}} = 35.7$ ,  $SD = 8.7$ ). The study was approved by the Department of  
156 Psychology Ethics Committee, Swansea University and all participants provided signed,  
157 informed consent. All methods were carried out in accordance with relevant guidelines and  
158 regulations (Declaration of Helsinki). Further data corresponding to the demographics and  
159 MRS measures of this sample are reported in Weidacker, et al.<sup>30</sup>. In brief, we previously

160 reported significant negative correlations for the GD sample between Glx/Cr in two  
161 locations, the dACC as well as occipital voxel, and gambling severity in terms of the PGSI<sup>39</sup> as  
162 well as the DSM-5<sup>2</sup> scores for problematic gambling behaviour. Further, no significant  
163 between-group differences were found regarding MRS measures, but the GD group scored  
164 significantly lower on full scale intelligence (FSIQ; assessed with the Wechsler Abbreviated  
165 Scale of Intelligence<sup>40</sup> subtests for Matrix Reasoning and Vocabulary) and higher on  
166 Attention deficit hyperactivity disorder (ADHD) symptoms (assessed with the World Health  
167 Organization Adult ADHD Self Report Scale version 1.1; ASRS<sup>41</sup>) compared to the non-GD  
168 group. Assessed with the Alcohol, Smoking, and Substance Involvement Screening Tests  
169 version 3<sup>42</sup>, GD participants scored also significantly higher on alcohol usage. Importantly,  
170 age, other substance use such as for tobacco, cannabis, cocaine and amphetamine as well as  
171 the presence of Axis 1 disorders (assessed with the MINI International Neuropsychiatric  
172 Interview version 5.0.0<sup>43</sup>) were not statistically different between the groups<sup>30</sup>. Due to the  
173 significant between-group differences relating to ASRS and FSIQ scores, Pearson  
174 correlations were performed to assess the necessity of including them as covariates; none  
175 of the correlations with Flanker and SST variables reached significance ( $|r| < .35$ ,  $ps > .08$ ),  
176 revealing no indication for inclusion.

## 177 **Assessments**

178 **Gambling severity.** The PGSI<sup>39</sup> assesses the severity of gambling problems via nine  
179 items, on a Likert scale from *never* (= 0; 92.9% of the non-GD group), *sometimes* (= 1; 7.1%  
180 of the non-GD group [1 participant scored 1]), *most of the time* (= 2) to *almost always* (= 3).  
181 All GD participants were categorized as *problem gamblers* (> 8 on the PGSI). The PGSI has



182 high internal consistency (Cronbach's  $\alpha = .90$ ) and adequate validity for both GD and non-GD  
183 groups<sup>44, 45</sup>.

184 The *Diagnostic and Statistical Manual of Mental Disorders 5 (DSM-5)*<sup>2</sup> states nine  
185 criteria for problematic gambling behaviour leading to significant past year distress  
186 categorized as *mild* (4-5 criteria apply; 33.3% of the gamblers), *moderate* (6-7; 25%) or  
187 *severe* gambling problems (8-9, 41.7%).

188 The *South Oaks Gambling Screen (SOGS)*<sup>46</sup> assesses gambling risk via 20 items.  
189 Participants were characterised as *no problems* (= 0; 92.9% of the non-GD group), *some*  
190 *problems* (1-4; 7.1% of the non-GD group [1 participant scored 1]) or *probable pathological*  
191 *gambling* (> 5; 100% of the GD group).

## 192 Procedure

193 Pre-screening for eligibility utilized the PGSI, SOGS, and DSM-5 as well as magnetic  
194 resonance exclusion criteria and participants were invited to the Imaging Centre at Swansea  
195 University upon meeting the inclusion criteria (i.e., PGSI score  $\leq 1$  or  $> 8$ , right handedness,  
196 and safety criteria for scanning). The behavioural and MRS assessments took place on  
197 separate days (mean number of days between testing sessions = 15.7). Before MRS testing,  
198 participants' blood alcohol levels were measured with single use breathalysers (none of the  
199 participants had consumed alcohol before testing). Behavioural tasks were administered in a  
200 counterbalanced order across participants.

201 **Flanker Task.** The flanker task was presented using Psychtoolbox<sup>47</sup> in combination  
202 with MATLAB R2010b (Mathworks Inc., Massachusetts, USA). In the 200 stimuli Flanker task,  
203 either congruent (C; 70%) arrows (e.g., > > > > >) or incongruent (IC; 30%) arrows (e.g., > >  
204 > < > >) were presented. Participants were instructed to press as fast and accurately as

205 possible in the direction where the middle arrow pointed to (button Z on the keyboard for  
206 middle arrows pointing to the left; button M for middle arrows pointing to the right) while  
207 ignoring all arrows on the sides. Within each stimulus type (C and IC), arrows pointing to the  
208 right and left were presented in equal proportions. The presentation of stimuli was pseudo-  
209 randomized with the restrictions to not have an IC trial presented at the first trial, exclude  
210 the possibility of two IC trials in a row, and to have between two and five C trials in between  
211 IC trials. In the intertrial interval, a centred fixation cross was presented, with randomized  
212 durations between 900 ms and 1200 ms, in steps of 50 ms. Before the start of the  
213 experimental task, 30 practice stimuli were shown to make participants familiar with the  
214 arrow design and task requirements. During this practice part, feedback was presented  
215 when response times exceeded 750 ms (“Please try to press faster”), upon wrong button  
216 presses (“Wrong direction”) and following correct button presses (“Well done!”).

217 **Stop Signal Task.** The Stop Signal task (SST) was presented using Psychtoolbox<sup>47</sup> in  
218 combination with MATLAB R2010b (Mathworks Inc., Massachusetts, USA). In the SST, one  
219 arrow is presented centrally per trial and the participants are asked to press the button  
220 corresponding to the direction the arrow is pointing to (right pointing arrows required the  
221 button M, left pointing arrows the button Z on the keyboard) as fast and accurately as  
222 possible. On the minority of trials, an auditory stop signal is presented following the visual  
223 arrow, in these trials the participants are asked to inhibit their already initiated motor  
224 response as quickly as possible. The experiment was programmed in three experimental  
225 blocks with a self-paced break in between blocks, per block 100 stimuli were presented  
226 (30% of stop trials). Stimulus presentation was pseudo-randomized with the only restriction  
227 to prevent two consecutive stop trials. Within each stimulus type (stop, go), left and right

228 arrows were presented equally often. In between trials a centred fixation cross was  
229 presented, with randomized durations between 900 ms and 1200 ms, in steps of 50 ms. The  
230 stop signal delay (SSD), the delay between the visual presentation of the arrow and the  
231 auditory stop signal, was initially set to 250 ms at task begin. Thereafter, each correct  
232 withholding of button presses in response to stop trials decreased the SSD by 50 ms  
233 (minimum was set to 50 ms), incorrectly pressing a button at stop trials increased the SSD  
234 by 50 ms. Before the start of the experimental task, 40 practice stimuli (12 stop trials) were  
235 presented. One participant was excluded from the non-GD group due to recording issues.

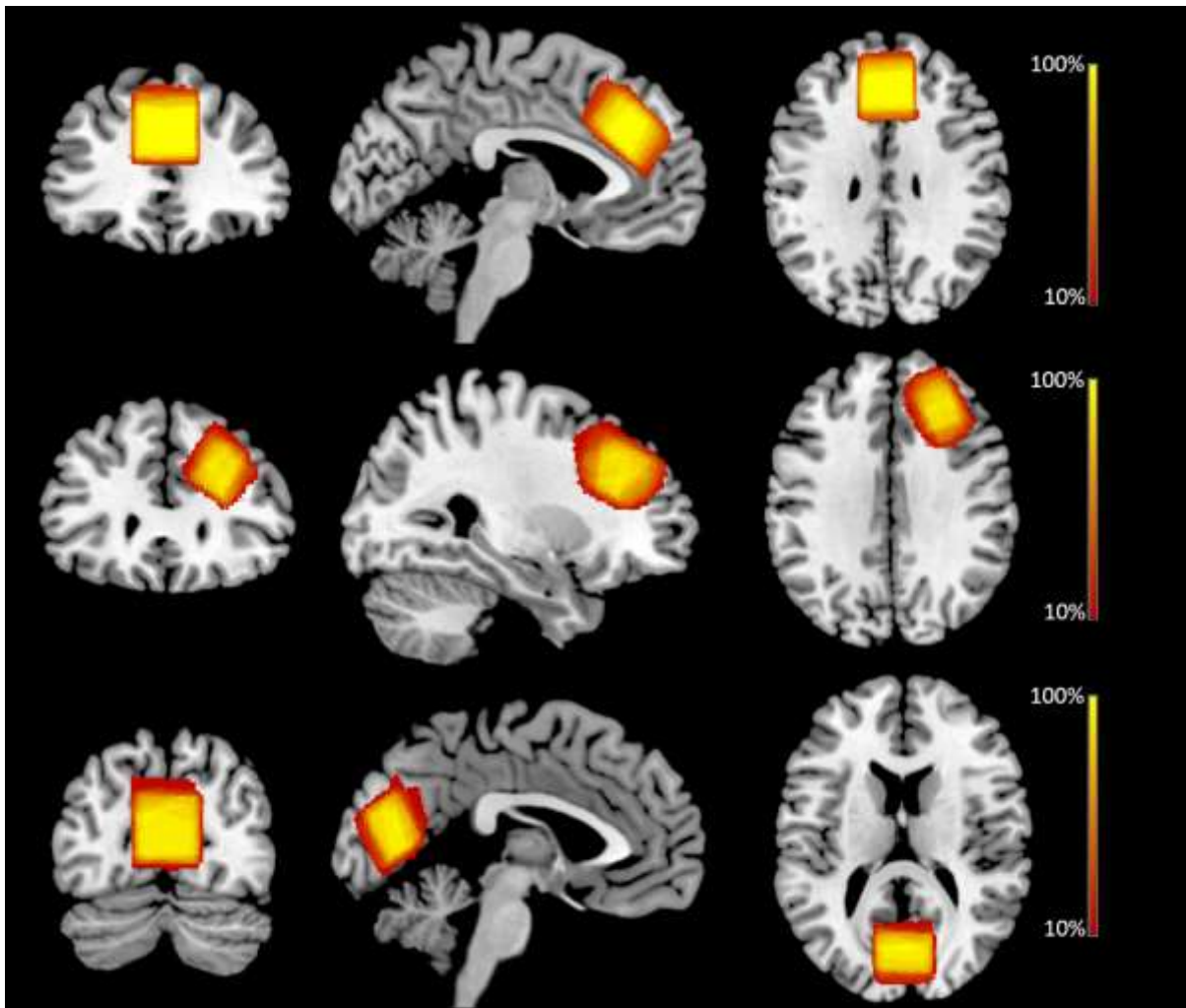
### 236 **MR acquisition**

237 MR was acquired using a 3-T Siemens Magnetom Skyra scanner (Siemens Medical  
238 Solutions, Erlangen, Germany; software version VD13) in combination with a 32-channel  
239 head coil. The MPRage sequence was used to obtain a T1-weighted image with the following  
240 parameters: repetition time (TR = 2200 ms), echo time (TE = 2.45 ms), inversion time (TI =  
241 900ms), flip angle (8 deg), 192 slices, 1 mm slices.

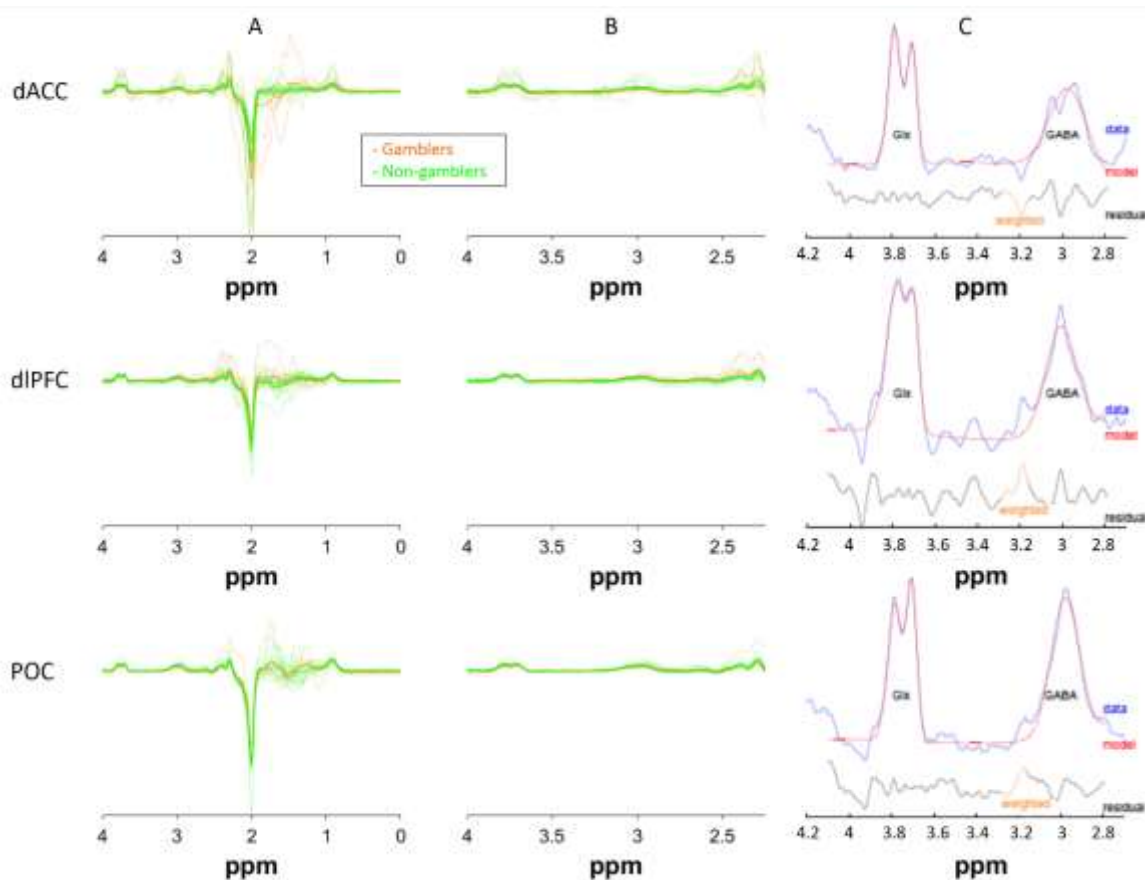
242 Single voxel MRS was based on the MEGA-PRESS MRS package<sup>48</sup> (provided by the  
243 University of Minnesota under a C2P agreement). The following VOIs were acquired in  
244 sequence: the dorsal ACC (30x30x20 mm), the right dlPFC (30x20x20 mm) and occipital,  
245 between the calcarine fissure and the parieto-occipital sulcus (20x30x25 mm). GABA+ was  
246 utilized as an edited estimate of gabaergic concentration (i.e., concentration/level of GABA)  
247 in the absence of macromolecule suppression and acquired with the following parameters:  
248 TR = 1800 ms, TE = 68 ms, 200 averages (per ON and OFF spectra), 1024 complex data  
249 points, editing pulse frequency = 1.90 ppm (4.70 ppm center frequency), editing pulse  
250 bandwidth = 52 Hz, offset frequency set to 3.00 ppm (reflecting the offset, relative to water,

251 of the carrier frequency of the slice-selective pulses). Higher-order shimming was performed  
252 manually to reduce local field inhomogeneities in each voxel of interest (VOI) and VAPOR  
253 was used for water suppression. No outer voxel suppression was applied. See Figure 1 for  
254 voxel locations, overlap and Figure 2 for corresponding mean and individual spectra per  
255 group. Recommended minimum reporting details for the MRS details are also included in  
256 appendix 1 as set out in the Minimum Reporting Standards for In Vivo Magnetic Resonance  
257 Spectroscopy (MRSinMRS): Experts' Consensus Recommendations<sup>49</sup>. Five participants  
258 produced no adequate MRS data for any of the three MRS voxels during acquisition (e.g.  
259 hardware failure, excessive motion, inadequate shimming) and were therefore excluded  
260 prior to this report.

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262  
263 *Figure 1.* Voxel locations for the dACC, dlPFC, and occipital voxels. Shown is the percentage  
264 overlap across all participants (from 10 to 100%) per location. Each participant's voxel  
265 location was transformed into MNI space before calculating the percentages. dACC = dorsal  
266 anterior cingulate, dlPFC = right dorsolateral prefrontal cortex, POC = posterior occipital  
267 cortex.  
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269  
 270 *Figure 2.* Spectra and example model fit for the dACC, dlPFC, and occipital MRS voxels. The  
 271 first column (A) shows the individual MRS spectra (from 0 to 4 ppm), the second column (B)  
 272 shows only the critical signal region (from 2.25 to 4 ppm). Both (A) and (B) are colour coded  
 273 with orange representing participants with and green representing participants without  
 274 gambling disorder. The respective group average MRS plots are added as a thicker line  
 275 following the same colour coding. The third column (C) shows an example GannetFit output  
 276 per MRS voxel. dACC = dorsal anterior cingulate, dlPFC = right dorsolateral prefrontal cortex,  
 277 POC = posterior occipital cortex.

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## 280 Spectral Quantification

281 MRS quantification was conducted via GANNET 3.0<sup>50</sup> (Baltimore, MD, USA) in  
 282 MATLAB on Siemens .rda files (averaged spectra) using the standard processing steps,  
 283 inbuilt models and assumptions for this software (details at <http://www.gabamrs.com>). The  
 284 edited spectrum was based on the subtraction of the 'ON' and 'OFF' spectra following  
 285 alignment of sub-spectra based on the spectral registration algorithm<sup>51</sup>. The GANNET  
 286 pipeline models GABA+, Glx and the creatine (Cr) reference as a single-Gaussian, doublet,

287 and singlet, respectively. Data is reported as a raw ratio of area under the fitted curve  
288 referenced to Cr (aligned with our previous report on this sample), for each metabolite, and  
289 does not account for differential proton densities, metabolite-specific relaxation properties,  
290 or tissue make up. In addition to GABA+/Cr and Glx/Cr, we utilized the GABA+/Glx ratio for  
291 MRS-related analyses in line with our previous report on this sample. While cerebrospinal  
292 fluid (CSF) correction is not necessary when using Cr as reference, grey matter (GM)  
293 contribution may however be of influence and is therefore controlled for in all MRS-related  
294 analysis via partial Pearson correlations using GM fraction of the respective MRS voxel, GM  
295 / (GM + CSF + white matter), as covariate. Per voxel, GM tissue fractions were obtained using  
296 unified segmentation<sup>52</sup> of the T1-weighted image in SPM12  
297 (<https://www.fil.ion.ucl.ac.uk/spm/>). Within each voxel (e.g. dACC), task performance  
298 indices (e.g. SST Go error RTs) were correlated with MRS measures (e.g. dACC GABA+/Cr)  
299 and corrected for GM contribution (e.g. GM in dACC MRS voxel) using partial Pearson  
300 correlations. However, producing visual representations (scatterplots) of partial Pearson  
301 correlations included a few more steps. First, both variables in a correlation pair were  
302 corrected for GM contribution by performing linear regressions (e.g. linear regression 1:  
303 predicting dACC GABA+/Cr from dACC GM; linear regression 2: predicting SST Go error RTs  
304 from dACC GM) and saving the corresponding residuals. These residuals are fully corrected  
305 for GM contribution and were used to create the corresponding scatterplot per significant  
306 correlation, and are as such simply a visual representation of a partial Pearson correlation,  
307 correcting for GM contribution.

308 From the 26 participants included in this report, individual MRS voxels were  
309 discarded due to inadequate MRS voxel acquisition during scanning, e.g. due to excessive

310 motion, inadequate shimming, hardware/recording issues (applies to 2 dACC, 1 dIPFC, and 4  
311 occipital voxels), bad model fit (applicable to 1 dACC voxel), presence of subtraction artifact  
312 (1 dACC, 1 occipital voxel), phase issues (1 dIPFC, 2 occipital voxels), and presence of  
313 truncation artifact (1 occipital voxel). GANNET Model fit was assessed based on visual  
314 inspection and FWHM (Full-width at half-maximum) within 3 *SDs* from the group mean per  
315 metabolite (e.g. within the dACC: GABA+, Glx). Due to excessive FWHM, one additional  
316 occipital voxel was excluded from analyses concerning GABA+. Data was included from 9  
317 participants in GD and 13 in non-GD for the dACC voxel, leading to mean (and *SDs*) of the  
318 signal-to-noise ratios (SNR) for Glx of 21.77 (14.99) and 29.10 (11.96) and for GABA+ of  
319 15.54 (10.69) and 18.65 (8.30), respectively for GD and non-GD. For the right dIPFC voxel, 11  
320 GD and 13 non-GD participants were included, with SNRs for Glx equalling 18.61 (8.47) and  
321 22.13 (7.27) and for GABA+ of 12.77 (5.09) and 16.35 (5.24). For the occipital voxel, 8 GD  
322 and 10 (Glx) or 9 (GABA+) non-GD participants were included, leading to SNRs for Glx of  
323 18.14 (6.23) and 19.12 (5.45) and SNRs for GABA+ of 19.44 (6.76) and 19.38 (5.90),  
324 respectively. The mean FWHM (and corresponding *SDs*) of included dACC MRS data  
325 equalled 14.42 (2.35) and 16.32 (3.15) for Glx, and 16.50 (6.16) and 20.43 (3.18) for GABA+,  
326 respectively for GD and non-GD. FWHMs for included dIPFC MRS data equalled 14.51 (1.92)  
327 and 15.11 (3.21) for Glx, and 16.55 (4.15) and 19.18 (4.25) for GABA+, respectively for GD  
328 and non-GD. FWHMs for included occipital MRS data equalled 13.99 (1.21) and 14.95 (1.12)  
329 for Glx, and 20.55 (4.01) and 21.76 (3.71) for GABA+, respectively for GD and non-GD. The  
330 mean Gannet Fit Error (*SD*) for included dACC MRS data equalled 8.29 (9.66) and 5.36 (1.71)  
331 for Glx/Cr, and 9.42 (5.71) and 8.40 (3.75) for GABA+/Cr, respectively for GD and non-GD.  
332 The mean Gannet Fit Error (*SD*) for included dIPFC MRS data equalled 6.60 (3.09) and 5.96



333 (2.09) for Glx/Cr, and 8.93 (3.44) and 8.16 (3.69) for GABA+/Cr, respectively for GD and non-  
334 GD. The mean Gannet Fit Error (*SD*) for included occipital MRS data equalled 6.58 (2.99) and  
335 6.02 (1.42) for Glx/Cr, and 6.47 (3.75) and 5.63 (1.60) for GABA+/Cr, respectively for GD and  
336 non-GD.

### 337 **Statistical Analysis**

338 **Flanker Task.** First an rmANOVA was conducted on correct response times with  
339 group as between-subject factor and trial type (C vs IC) as within-subjects factor. For error  
340 processing, two separate one-way ANOVAs were conducted with group as between-subject  
341 factor, the first on IC error response times, the second on error percentages. We confined  
342 the error analyses to IC trials (%  $M = 9.04$ ,  $SD = 8.92$ , range = 0 to 40), since few participants  
343 made errors in C trials (%  $M = .77$ ,  $SD = .88$ , range = 0 to 3.57). One participant per group  
344 made no IC errors and both were therefore excluded from the error response time analyses.

345 Post-error slowing (PES) analyses were based on correct trials preceding and  
346 following IC errors as suggested for calculating robust PES<sup>53, 54</sup>. The functional role of PES,  
347 the observation that trials following an error produce longer response times than trials  
348 being preceded by a correct trial<sup>25</sup> is under debate with arguments for reducing future error  
349 responses or being a result of the increased salience of errors among others<sup>26, 27</sup>. Earlier  
350 investigations into PES and gambling behaviour were based on only post-error response  
351 times, termed the traditional method to calculate PES<sup>54</sup>. Comparing approaches to PES  
352 calculation however, showed that the traditional method is affected by global changes in  
353 attention and motivation, therefore underestimates PES, and is outperformed by the robust  
354 method which compares post-error responses to pre-error responses<sup>53</sup>. These trial types

355 were subjected to an rmANOVA as within-subject factors, adding group as between-subject  
356 factor, and the resultant sample size was 11 for GD and 13 for the non-GD group.

357 The Flanker variables (response time differences between IC and C trials, percentage  
358 and response times of IC error trials, and PES) were correlated to dACC, dlPFC and occipital  
359 MRS variables (Glx/Cr, GABA+/Cr, GABA+/Glx ratio) using partial Pearson correlation  
360 coefficients, correcting for grey matter (GM) content within each voxel, , first using all  
361 participants and thereafter separately per GD and non-GD. The derived significant partial  
362 correlation coefficients were statistically compared following Fisher's  $r$  to  $z$  transformation.  
363 As this was an exploratory study, data are reported using exact  $p$ -values without correction.

364 **Stop Signal Task.** The mean SSD was calculated as the average of SSDs stemming  
365 from successful stop trials and trials with premature responses (button presses that  
366 occurred before the stop signal). The probability of responses occurring when a stop signal  
367 was presented was calculated as inverse of % correct stop trials. To estimate SSRT (stop  
368 signal response time), the go response time that matches this probability within the  
369 distribution of response times to go trial (including wrong trials and imputing the response  
370 times of missed go trials with the maximum of that distribution) was selected, and mean  
371 SSD was subtracted from it.

372 The SST exclusion criteria were as follows: SSRTs indicating waiting for the stop signal  
373 (e.g., negative SSRT, applicable to one GD participant), probabilities outside the range of 24.4  
374 to 75% (not met by participants in this study), response time higher at unsuccessful stop  
375 trials than the mean of the go trial distribution (not applicable to these participants),  
376 recording issues (applicable to one non-GD participant). Applying these criteria resulted in a  
377 behavioural sample comprised of 11 GD and 13 non-GD participants. Calculations of mean

378 SSD, probability and SSRT are in line with recent suggestions on the use of the integration  
379 method for SST studies<sup>55</sup>.

380 Group-differences were assessed using separate one-way ANOVAs with group as  
381 between-subjects factor on response times to correct Go trials, % errors to Go trials (%  $M =$   
382 2.82,  $SD = 3.71$ , range = 0 to 17.14), SSDs and SSRTs. Error response times for choice errors  
383 (Go: pressing the wrong direction) and inhibition failures (Stop: pressing during stop trials)  
384 were analysed using an rmANOVA with trial type (Go vs. Stop) as within- and group as  
385 between-subjects factor. Post-error slowing was analysed as described above for the  
386 Flanker task, making use of trials preceding and following failed inhibition errors, in a  
387 rmANOVA with group as between-subject factor. Two GD participants did not make errors  
388 to Go targets and were therefore not incorporated in the associated response time  
389 analyses. The MRS variables were correlated to the SST variables (response times to correct  
390 go, wrong go and wrong stop trials, percentages go errors, PES, SSD and SSRT) in the same  
391 manner as outlined for the Flanker task. Given the additional outlier criteria applied to the  
392 SST, the correlations between SST variables and dACC MRS measures were based on 8 GD (7  
393 for Go Error response times) and 12 non-GD participants, while correlations between dlPFC  
394 MRS measures and SST task-data were based on 10 GD and 12 non-GD participants, and  
395 correlations between POC MRS measures and SST performance were based on 8 GD and 9  
396 (Glx) or 8 (GABA+) non-GD participants.

397 Across tasks, significant rmANOVA results are accompanied by  $\eta^2_p$  as effect size,  
398 while Cohen's  $d$  is used for independent-sample  $t$ -tests. G\*Power 3.1.9.2<sup>56</sup> was used for  
399 sensitivity analysis and the smallest detectable effect size  $d$  for between-group effects  
400 equalled 1.20, given our SST sample sizes, a two-sided  $\alpha$  of .05 and 80% power. Regarding  $t$ -

401 tests, Levene's Test for Equality of Variances was performed and corrected statistics are  
402 reported when applicable. Multivariate normality (of all three variables within a partial  
403 Pearson correlation) was ascertained using Chi-square generalized distance plots obtained  
404 via the software Statgraphics (Version 18, Statistical Graphics Corporation, Rockville, USA)  
405 and all variables included in significant correlations fell within the 95% confidence interval,  
406 consistent with the hypothesis of an underlying multivariate normal distribution.

## 407 **Results**

### 408 **Demographics**

409 As expected, the GD and non-GD groups differed significantly on PGSI scores,  
410  $t(11.05) = 10.19, p < .001, d = 4.01$ , SOGS scores,  $t(11.12) = 11.68, p < .001, d = 4.59$ , and  
411 number of endorsed DSM-5 criteria,  $t(11) = 12.45, p < .001, d = 4.90$  (see Weidacker et al.<sup>30</sup>  
412 for further details).

### 413 **Response Interference (Flanker task)**

414 A rmANOVA on response times across trial types revealed a main effect of trial-type  
415 ( $F(1,24) = 87.03, p < .001, \eta^2_p = .78$ ), group ( $F(1,24) = 6.09, p = .02, \eta^2_p = .20$ ) and a non-  
416 significant interaction between group and trial type ( $F(1,24) = 3.92, p = .06$ ). Response times  
417 to IC trials were significantly slower ( $M = 535.27, SD = 108.30$ ) than to C ( $M = 424.80, SD =$   
418  $73.70$ ) trials, regardless of gambling status. Gamblers had significantly longer response times  
419 ( $M = 521.40, SD = 105.52$ ) than the non-GD group ( $M = 444.58, SD = 46.22$ ) when averaged  
420 across trial type.

421 No significant effects of group were found when analysing the IC error response  
422 times ( $F(1,23) = 2.32, p = .14$ ) and the percentage of IC errors ( $F(1,25) = 1.82, p = .19$ ). The  
423 rmANOVA on trials preceding and following IC errors revealed significant post-error slowing

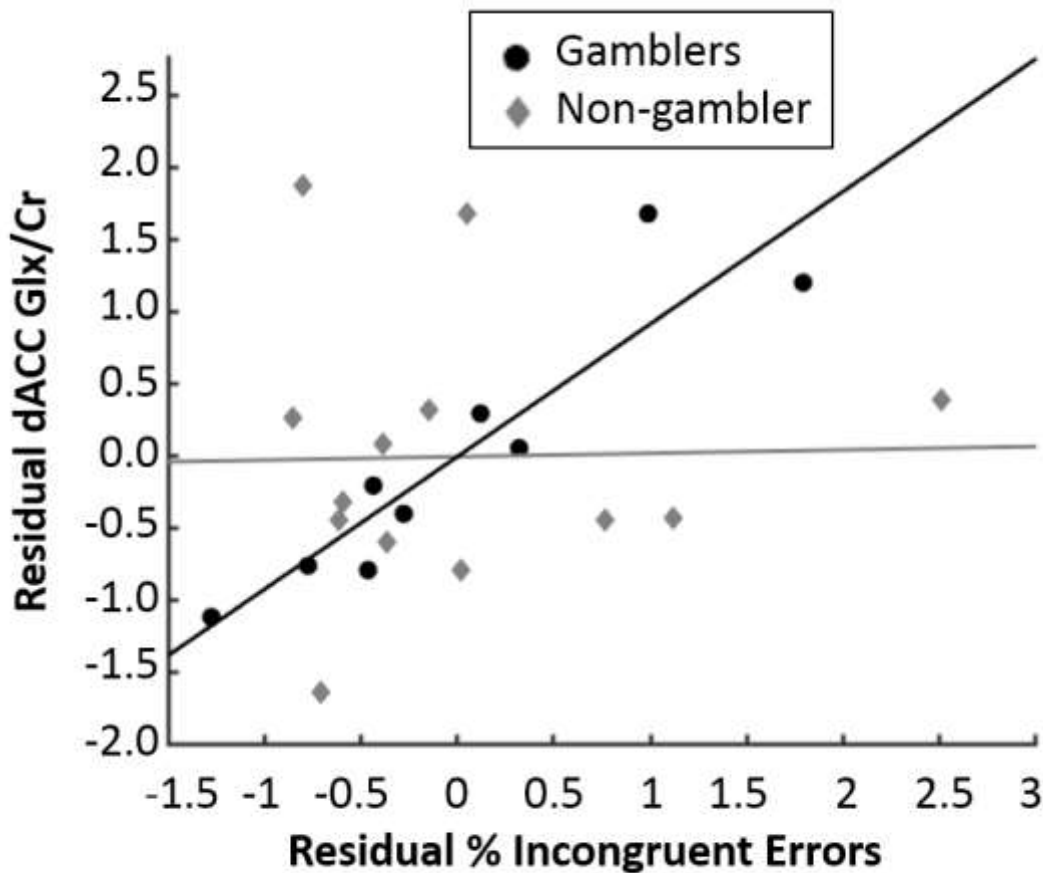
424 ( $F(1,22) = 11.88, p = .002, \eta^2_p = .35$ ), but no significant main effect of group ( $F(1,22) = 11.11, p$   
425  $= .30$ ) or interaction between group and trial type ( $F(1,22) = .01, p = .93$ ).

426 **Correlations between response interference and MRS measures.** Analysing  
427 associations between dACC MRS variables and Flanker variables did not reveal significant  
428 correlations in the whole sample ( $|r_s| < .38, p_s > .12$ ). When assessing the correlations  
429 within the GD group, dACC Glx/Cr was significantly positively correlated with the proportion  
430 of IC Errors ( $r = .92, p = .001$ ) the remaining correlations were not statistically significant  
431 ( $|r_s| < .69, p_s > .05$ ). Within the non-GD group, this correlation, between dACC Glx/Cr and  
432 proportion of IC Errors, was not significant ( $r = .02, p = .94$ ), as were the remaining  
433 correlations ( $|r_s| < .27, p_s > .40$ ). Using Fisher's  $r$  to  $z$  transform, the difference between the  
434 correlation coefficient obtained for the relationship between dACC Glx/Cr and proportion of  
435 IC Errors was significantly larger in GD than the non-GD group ( $z = 3.03, p = .002$ ), see Figure  
436 3.

437

438

\*\*\*FIGURE 3\*\*\*



439

440 *Figure 3.* Scatterplot of the significant correlation (adjusted for grey matter content)  
 441 between Glx in the dACC and percentage errors to incongruent trials in the Flanker task.  
 442 This relationship is shown in black for gambling ( $r = .92, p = .001$ ) and in grey for non-  
 443 gambling participants ( $r = .02, p = .94$ ). dACC = anterior cingulate cortex. Lines represent the  
 444 least squares fit to the data.

445

446 In the dIPFC voxel, no correlations between MRS and Flanker variables were  
 447 significant in the whole sample ( $|r_s| < .25, p_s > .26$ ), the GD group ( $|r_s| < .52, p_s > .12$ ) and  
 448 the non-GD group ( $|r_s| < .59, p_s > .05$ ). In the occipital voxel, no correlations were significant  
 449 for the whole sample ( $|r_s| < .35, p_s > .21$ ), the GD group ( $|r_s| < .36, p_s > .48$ ) and the non-GD  
 450 group ( $|r_s| < .67, p_s > .05$ ).

451

452 **Response Inhibition (Stop Signal Task)**

453           Analysing the effect of gambling status on correct response times to Go trials in the  
454 SST revealed no significant effect ( $F(1,23) = 3.18, p = .09$ ). The rmANOVA on response times  
455 for choice errors for Go and inhibition errors on Stop trials produced a significant main  
456 effect of trial type ( $F(1,20) = 34.25, p < .001, \eta^2_p = .63$ ), due to longer response times when  
457 performing errors of inhibition ( $M = 379.09, SD = 58.50$ ) than errors of choice ( $M = 195.57,$   
458  $SD = 47.26$ ). The interaction between group and trial type ( $F(1,20) = .63, p = .44$ ) as well as  
459 the main effect of group were not significant ( $F(1,20) = 1.5, p = .23$ ). Further, the percentage  
460 of choice errors on Go trials did not differ significantly between groups ( $F(1,23) = 1.80, p =$   
461  $.19$ ).

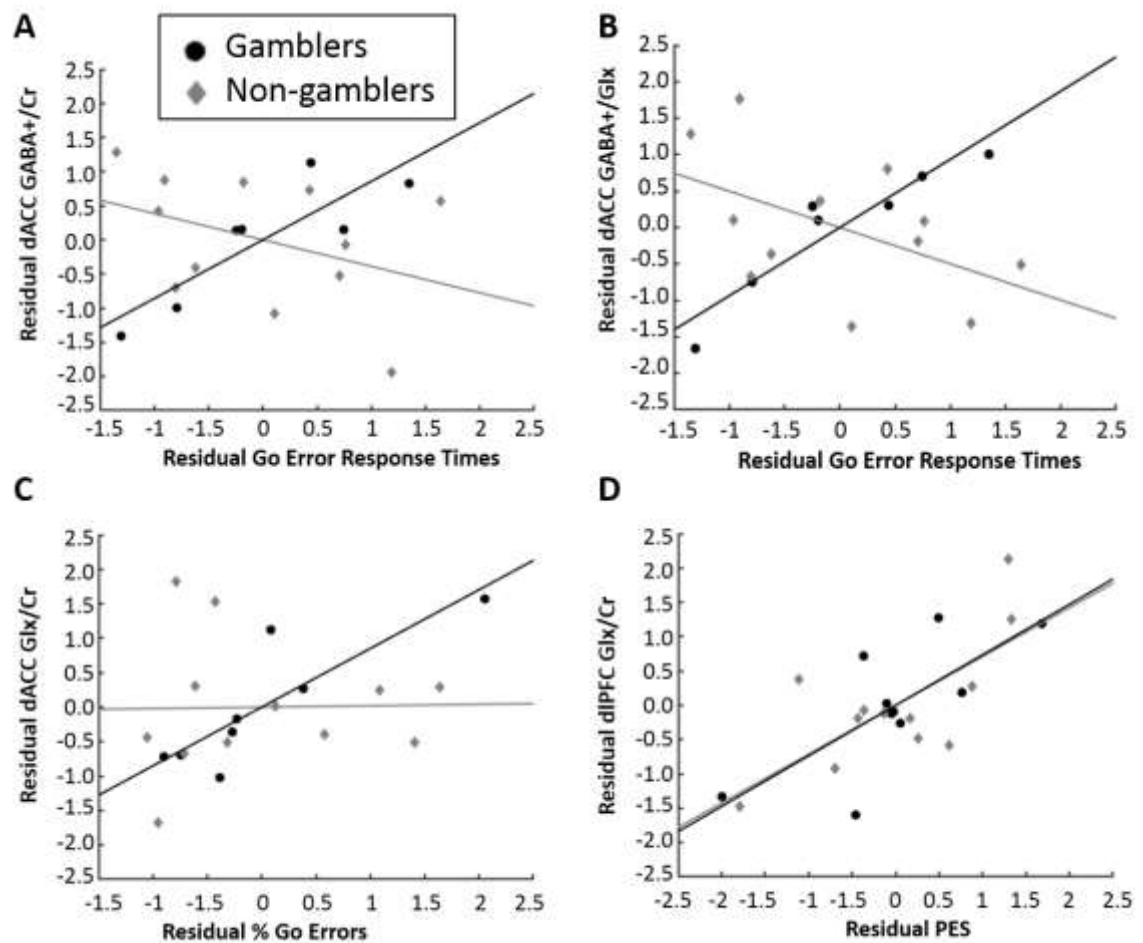
462           Analysing the SST inhibition-related variables, mean SSDs ( $F(1,23) = 2.71, p = .11$ ) and  
463 SSRTs ( $F(1,23) = 1.42, p = .25$ ) revealed no significant group differences between GD and  
464 non-GD groups. The rmANOVA on post-error slowing revealed no significant main effect of  
465 trial type ( $F(1,22) = 3.94, p = .06$ ), group ( $F(1,22) = 1.65, p = .21$ ) or interaction between  
466 them ( $F(1,22) < .01, p = .99$ )

467           **Correlations between response inhibition and MRS measures.** Analysing  
468 associations between dACC MRS variables and SST variables did not reveal any significant  
469 correlations in the whole sample ( $|r_s| < .39, p_s > .09$ ). When assessing the correlations  
470 within the GD group, the response times on Choice (Go) errors correlated positively with  
471 GABA+/Cr ( $r = .86, p = .03$ ) and the GABA+/Glx ratio ( $r = .936, p = .006$ ). Both correlations  
472 were not significant in non-GD, with  $r = -.39, p = .24$  for the correlation between Go error  
473 response times and GABA+/Cr, and  $r = -.50, p = .12$  for the correlation with GABA+/Glx.  
474 Comparing the obtained correlation coefficients for the relationship between GABA+/Cr and

475 Go Error response times across groups revealed a significantly stronger correlation in GD  
476 than non-GD ( $z = 2.83, p = .004$ ), see Figure 4A. Similarly, the correlation between Go Error  
477 response times and the GABA+/Glx ratio was significantly larger in GD than non-GD ( $z =$   
478  $3.75, p < .001$ ), see Figure 4B. Further, the % Go Errors correlated positively with dACC  
479 Glx/Cr ( $r = .85, p = .015$ ) in GD, while this correlation was not significant in non-GD ( $r = .02, p$   
480  $= .95$ ). The remaining correlations were not significant ( $|r_s| < .58, p_s > .17$ ) in GD and none of  
481 the correlations was significant within the non-GD sample ( $|r_s| < .50, p_s > .11$ ). Comparing  
482 the groups on their correlation coefficients obtained for the association between % Go  
483 Errors and Glx/Cr revealed a significantly stronger correlation in GD than non-GD ( $z = 2.23, p$   
484  $= .03$ ), see Figure 4C.

485





486

487 *Figure 4.* Scatterplot of the significant correlations (adjusted for grey matter content)  
 488 obtained for the Stop Signal Task. Data from gamblers are shown in black and data from  
 489 non-gamblers are depicted in grey. Lines represent the least squares fit to the data. A)  
 490 Positive, significant, correlation between Go Error response times and dACC GABA+/Cr in  
 491 gamblers ( $r = .86, p = .03$ ); this correlation was not significant in non-gamblers ( $r = -.39, p =$   
 492  $.24$ ). B) Positive, significant, correlation between Go Error response times and ACC  
 493 GABA+/Glx ratio in gamblers ( $r = .936, p = .006$ ); this correlation was not significant in  
 494 nongamblers ( $r = -.50, p = .12$ ). C) Positive, significant, correlation between % Go Error  
 495 responses and dACC Glx/Cr in gamblers ( $r = .85, p = .015$ ); this correlation was not significant  
 496 in non-gamblers ( $r = .02, p = .95$ ). D) Positive, significant, correlations between post-error  
 497 slowing (PES) and dlPFC Glx/Cr in gamblers ( $r = .74, p = .02$ ) and non-gamblers ( $r = .71, p =$   
 498  $.01$ ). dACC = dorsal anterior cingulate cortex, dlPFC = dorsolateral prefrontal cortex.  
 499

500 Analysing the partial correlations between dlPFC MRS variables and SST variables in  
 501 the whole sample, revealed a significant correlation between Glx/Cr and PES ( $r = .69, p <$   
 502  $.001$ ), all remaining correlations were not significant ( $|r_s| < .37, p_s > .13$ ). The significant

503 correlation between Glx/Cr and PES was confirmed in both, the GD ( $r = .74, p = .02$ ) and  
504 non-GD group ( $r = .71, p = .01$ ), see Figure 4D. Within the GD group, dIPFC Glx/Cr also  
505 correlated negatively with the percentage Go/choice errors ( $r = -.68, p = .04$ ) while this  
506 correlation was not significant in the non-GD group ( $r = .14, p = .68$ ). When comparing  
507 correlation coefficients for the association between dIPFC Glx/Cr and the percentage  
508 Go/choice errors across groups, no significant difference was obtained ( $z = 1.94, p = .05$ ).  
509 The remaining correlations were not significant within the GD ( $|r_s| < .67, p_s > .10$ ) and non-  
510 GD groups ( $|r_s| < .45, p_s > .18$ ).

511         Assessing the significance of the partial correlations between occipital MRS and SST  
512 variables revealed no significant correlations in the whole sample ( $|r_s| < .51, p_s > .06$ ), GD  
513 ( $|r_s| < .68, p_s > .22$ ), and non-GD ( $|r_s| < .75, p_s > .05$ ).

514  
515

### Discussion

516         The present study is the first investigation of distractor interference and response  
517 inhibition performance in GD, with *in vivo* GABA+/Cr and Glx/Cr metabolic measurements  
518 obtained from three brain areas (dACC, right dIPFC, and an occipital control voxel).  
519 Gambling disorder individuals' behavioural performance evidenced prolonged response  
520 times in the Flanker Task, regardless of stimulus congruency. On the other hand, SST  
521 performance did not suggest prolonged response times or inhibition deficits in those with  
522 GD. It is possible therefore that the complex stimuli used in the Flanker task might at least  
523 partially explain the reduced processing speed in GD that we observed. Despite error  
524 responses in the Flanker Task not differentiating between groups, GD participants expressed  
525 a positive correlation between dACC Glx/Cr and the number of errors in response to IC

526 targets, which was significantly larger than the correlation coefficient found within non-GD  
527 participants.

528         This is the first report of a positive correlation between dACC Glx/Cr and error rates  
529 on the Flanker task; the only previous related investigation focussed on response times and  
530 Glx and found no significant association<sup>37</sup>. These different results suggest that dACC Glx/Cr  
531 may play a more prominent role in terms of error rates<sup>57-59</sup> than response times, in line with  
532 previous reports on increased glutamate-glutamine ratio levels in the dACC being associated  
533 with increased self-reported impulsivity as well as increased error rates on a Go/No-go  
534 task<sup>57</sup>. Similarly, decreased ACC Glutamate/Cr was previously associated with increases in  
535 cognitive control-related striatal activation when contrasting Stroop IC to C trials, and this  
536 activation in turn was correlated positively with error rates<sup>59</sup>. In sum, despite few  
537 behavioural differences between GD and non-GD on distractor interference measures, the  
538 metabolic differences suggest potentially abnormal dACC function related to error  
539 processing. This warrants further investigation of any associated striatal abnormalities in GD  
540 during interference-related errors.

541         Response inhibition, in terms of SST performance, has been subject to several  
542 investigations in GD, with individual studies finding heterogeneous results and meta-  
543 analyses indicating either no or moderate to large effects on SST response inhibition indices,  
544 respectively<sup>16, 24</sup>. Previous research also supports the idea that response inhibition deficits in  
545 GD might emerge at higher gambling severity levels<sup>17, 23</sup>, but this hypothesis was not  
546 supported by the present investigation which focussed solely on participants with high  
547 gambling severity. However, due to the restrictive inclusion criteria, the current study

548 suffers from a relatively small sample size and might therefore not be perfectly suited to  
549 identify smaller effects and should be a starting point for larger scale research.

550 Like previous investigations on SST-type tasks, PES was unaffected by the presence  
551 of GD in both the SST and Flanker tasks, despite both tasks producing significant PES.  
552 However, SST and Flanker PES seem to involve different neural aspects, only SST PES  
553 correlated positively with Glx/Cr levels in the right dIPFC, and no dissociation in the strength  
554 of correlation as a function of gambling addiction status was observed. Previous research on  
555 neural involvement during PES found a positive correlation between PES and left anterior  
556 midcingulate white matter, a region which supports connectivity to frontopolar and  
557 dorsolateral frontal brain regions<sup>60</sup>. However, dIPFC involvement in post-error slowing  
558 shows task-dependent variations<sup>60</sup> and might represent a subprocess of PES<sup>61</sup>. PES in terms  
559 of the Flanker task was found to be unaffected by Lorazepam and gamma-hydroxybutyrate,  
560 two GABA agonists working on different receptor types<sup>62, 63</sup>, but PES was less pronounced in  
561 Flanker when compared to Stroop and Go/No-go tasks<sup>64</sup>. Within the Stroop task, Moeller, et  
562 al.<sup>65</sup> investigated the effect of methylphenidate on PES and reported enhanced PES  
563 following administration of the drug, which is thought to excite GABAergic interneurons as  
564 well as increase glutamate uptake<sup>66, 67</sup>. As such, the finding that the neurochemical  
565 involvement in PES differs between Flanker and SST tasks might be due to task design and  
566 associated differences in pronunciation of PES. In the SST, it was notable that PES was  
567 positively associated with dIPFC Glx/Cr, regardless of gambling status.

568 While our small samples of GD and non-GD showed consistent correlations between  
569 PES and dIPFC Glx/Cr levels, analysis of GD participants revealed additional associations  
570 between MRS measures and SST error processing indices that differed in directionality and

571 significance to non-GD participants. Previous research on GD and error processing in the SST  
572 is limited, with Lorains, et al.<sup>18</sup> revealing enhanced Go error frequency in treatment-seeking  
573 problem gamblers, while Lawrence, et al.<sup>21</sup> found no between-group differences. The  
574 current investigation did not reveal behavioural differences in SST error processing, but did  
575 suggest between-group correlation differences between SST error processing and MRS  
576 measures. In the dACC, GABA+/Cr as well as the GABA+/Glx ratio correlated positively and  
577 significantly with Go error response times in GD, whereas both correlations were negative  
578 and did not reach significance in non-GD. Similarly, in GD, baseline dACC Glx/Cr correlated  
579 positively and significantly with the frequency of Go errors, a correlation which was also not  
580 significant in non-GD. This positive association between dACC Glx/Cr and SST Go errors in  
581 GD resembles that found between dACC Glx/Cr levels and error rates for the Flanker task,  
582 perhaps indicating a general influence of baseline Glx on error processing deficits in GD.

583         In contrast to the Flanker task, response times in the SST did not depend on  
584 gambling status, but the positive correlations between Go error response times and dACC  
585 GABA+/Cr and the GABA+/Glx ratio indicated GD-specific abnormalities. Previous research  
586 with non-gambling populations showed that enhancing GABA levels via agonists, such as  
587 Diazepam or Lorazepam, prolongs response times across tasks<sup>38, 68</sup>, like the positive  
588 association between baseline GABA+/Cr and SST response times found in the current  
589 investigation. While GABA agonists induce widespread increases in cortical GABA, our  
590 investigation found the relationship between GABA+/Cr and response times significant  
591 within the dACC voxel. Previous neuroimaging research suggests hypo- or hyper-activation  
592 in the dACC during SST in GD and frequent poker players, respectively<sup>15, 19</sup>. Together, this  
593 suggests that in GD dACC function may be affected and accompanied by neurochemical



617 Additionally, response inhibition did not differ statistically between GD and non-gamblers.  
618 Neurochemically, GD expressed enhanced correlations between baseline dACC GABA+/Cr  
619 and Go error response times as well as between dACC Glx/Cr and frequency of Go errors in  
620 the SST and the frequency of IC errors in the Flanker task. Further, GD and non-GD  
621 participants expressed equivalently efficient PES in both response inhibition and distractor  
622 interference tasks, while neural involvement of baseline dlPFC Glx/Cr levels in the SST-based  
623 PES did not vary depending on gambling status.

624

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626

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633

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634

635

## Disclosure information

636

Kathrin Weidacker, Stephen J. Johnston, Paul G. Mullins, Frederic Boy, and Simon Dymond

637

have nothing to disclose.

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