Diffusion Tensor Imaging of the Corpus Callosum in Addiction

Danilo Arnone²,³  Mohammed T. Abou-Saleh²  Thomas R. Barrick¹

¹Division of Mental Health and ²Centre for Clinical Neuroscience, Division of Cardiac and Vascular Sciences, St. George’s University of London, London, and ³Department of Psychiatry, University of Oxford, Warneford Hospital, Oxford, UK

Introduction

Interest in the corpus callosum (CC) in addictions began in 1985 with post-mortem neuropathological studies of patients with alcohol dependence (AD) consistently showing volumetric reduction in the white matter [1]. Most MRI studies have also confirmed white matter volume reduction in patients with AD although restricted to gross estimations. However, less is known about the mechanisms of alcohol-related volume loss and its functional significance [2, 3], nor are the mechanisms underlying volume restoration observed in alcohol abstinence [2, 4] clear. There is also controversy over the effect of gender and advancing age on white matter pathology in alcoholism [5, 6]. Neuropathology is even more uncertain in other types of substance misuse [7]. Voxel-based morphometry studies support the hypothesis that MDMA (Ecstasy) users show reduction in cortical grey matter concentration in multiple brain regions including neocortex, brainstem, cerebellum, and anterior cingulate gyrus [8], which may translate into white matter abnormalities characterised by ‘pruning’ of serotoninergic neurons [9]. In cocaine users volumetric changes have been demonstrated [10] and white matter deficiencies could derive from ‘perfusion defects’ attributable to long-lasting alter-
ations in cerebral blood flow, enduring alterations in cerebral glucose metabolism persisting after cocaine discontinuation or neurotoxicity to brain dopamine neurons [11]. In chronic cannabis users, although there are signs of mild cognitive impairment, there is little evidence that such impairments are accompanied by drug-induced neuropathology [12]. Diffusion tensor imaging (DTI) is a novel, non-invasive magnetic resonance technique capable of providing a more sensitive detection of white matter pathology. DTI quantifies in vivo the directionality and coherence of white matter fibre tracts [13] by detecting the freedom with which water molecules move within a tissue type and the amount and predominant intravoxel orientation of their diffusion [14]. Mean diffusivity (MD), a measure computed from DTI, quantifies the magnitude of (isotropic) water diffusivity in each image voxel and increases when boundaries to water diffusion are reduced. In particular, increased MD has been reported in lesions attributable to different neuropsychiatric conditions which may reflect oedema, demyelination, and axonal loss [15]. Fractional anisotropy (FA) is a further measure computed from DTI and quantifies how restricted (anisotropic) water diffusion is in each image voxel. Specifically, restrictions to water diffusion are caused by tissue microstructure, with a reduction in FA indicating a decline in tissue structural integrity. Finally, as DTI also provides a direction of predominant intravoxel orientation that corresponds to gross tissue orientation, the intravoxel coherence (IC) may be computed. This measure determines the coherence of local intravoxel diffusion orientations such that if local orientations are similar the IC is greater, with a reduced IC possibly providing evidence of disruption in tract organisation.

In summary, increased MD, reduced FA and reduced IC provide evidence of structural damage and may reflect a disruption in the organization of tracts [16]. These complementary measures are considered to be sensitive indices of axonal integrity [17]. For these reasons, a review of published data on CC volume in addictions was carried out. The CC was chosen because of its functional significance [18] and it is a large enough structure to allow reliable identification with region-of-interest methods. The aims of this review were (1) to establish whether DTI studies support current knowledge about volumetric changes in CC in alcoholism, (2) to clarify the role of ethanol-induced damage with reference to mechanisms of action, age, gender differences, lifetime alcohol consumption, time since last use, and (3) to determine the capability of DTI in detecting abnormalities in CC in other substance users.

Methods

A comprehensive search from a range of electronic databases, including BNI, CancerLit, Cochrane Library, EMBASE, Medline, Psychinfo, and PubMed was conducted for the period from the introduction of DTI to July 2006. Key words used to identify the studies were: diffusion tensor imaging, magnetic resonance imaging, DTI, RMI, alcoholism, marijuana, cannabis, cocaine, Ecstasy, MDMA, methamphetamine, and substance misuse. The search was also complemented by a manual search of bibliographic cross-referencing. Researchers who had expressed an interest in the subject were contacted for any non-published information. Papers were included if they presented original data and addressed the question, ‘use of DTI in substance misuse’. Studies were screened for diagnosis according to ICD-10 or DSM-IV, inclusion/exclusion criteria, demographic variables (e.g., age, gender), clinical settings, length of exposure to substance of abuse, length of abstinence prior to scanning, presence of physical illness and co-morbid psychiatric illness. Studies with results that did not reach statistical significance or reported results for areas other than the CC were excluded.

Results

Eight published and one unpublished reports met criteria for inclusion (table 1), five on alcoholism [19–23], one on cocaine use [24] and three on cannabis [25–27]. One study by Lim et al. [28] was excluded because it did not include the CC. The studies were all case-control comparisons. Meta-analytic evaluation was not possible because group differences were presented in a form not amenable to effect size calculation. Details of the reports are given below.

Alcohol Dependence

All the studies [19–23] had a similar design: patients recruited were chronic users with a large consumption of ethanol assessed with semi-structured alcohol consumption assessment interviews and were abstinent prior to scanning; caseness was identified with structured clinical interviewer tools and diagnosis of AD was given according to standard diagnostic criteria; major mental and physical illnesses were excluded; confounding variables were minimised by assessing pre-morbid IQ, analysing gender-related data separately, excluding other substance misuse, and matching for age. Only one study matched for handedness [22], and another for body mass index [23]. Pfefferbaum et al. [19] recruited 15 alcoholic men and 19 controls. CC volume showed lower FA in the alcoholic group in the genu (p < 0.05) with a significant correlation with time from last drink (p < 0.001) but not age (p > 0.05). IC failed to yield significant group or interac-
Diffusion Tensor Imaging of the Corpus Callosum in Addiction

Table 1. Studies of corpus callosum in addiction

<table>
<thead>
<tr>
<th>Study name</th>
<th>Cases mean consumption/ duration of use</th>
<th>abstinence days</th>
<th>n (m/f)</th>
<th>mean age years</th>
<th>Controls mean consumption</th>
<th>n (m/f)</th>
<th>mean age years</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pfefferbaum et al. [19]</td>
<td>1,484.6 kg</td>
<td>322</td>
<td>15/0</td>
<td>55.4</td>
<td>65.9 kg</td>
<td>19/0</td>
<td>54.4</td>
<td>tFA in G (p &lt; 0.05)</td>
</tr>
<tr>
<td>Pfefferbaum and Sullivan [20]</td>
<td>m: 301 kg</td>
<td>322</td>
<td>15/12</td>
<td>m: 55.4</td>
<td>m: 65.9 kg</td>
<td>31/18</td>
<td>44.7</td>
<td>f: tFA in G (p &lt; 0.05)</td>
</tr>
<tr>
<td></td>
<td>f: 1,484.6 kg</td>
<td>677</td>
<td>f: 45.8</td>
<td>m: 13.3 kg</td>
<td>f: 43.3</td>
<td></td>
<td></td>
<td>f: tIC in G (p ≤ 0.02)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>f: tIC in G (p ≤ 0.02) and S (p &lt; 0.05)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>f: tIC in S (p = 0.0001)</td>
</tr>
<tr>
<td>Pfefferbaum and Sullivan [21]</td>
<td>1,484.6 kg</td>
<td>322</td>
<td>15/0</td>
<td>55.4</td>
<td>65.9 kg</td>
<td>19/0</td>
<td>54.4</td>
<td>tMD in CC (p = 0.0698) tFA in CC (p &lt; 0.05)</td>
</tr>
<tr>
<td>Schulte et al. [22]</td>
<td>511.3 kg</td>
<td>NS</td>
<td>7/4</td>
<td>42.6</td>
<td>48.5 kg</td>
<td>8/5</td>
<td>51.6</td>
<td>tFA in CC (p &lt; 0.03) tMD in CC (p &lt; 0.005)</td>
</tr>
<tr>
<td>Pfefferbaum et al. [23]</td>
<td>m: 1,032.8 kg</td>
<td>NS</td>
<td>40/17</td>
<td>m: 52.8</td>
<td>m: 66.6 kg</td>
<td>32/42</td>
<td>m: 52.2</td>
<td>tFA in CC (p &lt; 0.001) (men &gt; women), tMD in CC (p &lt; 0.0008)</td>
</tr>
<tr>
<td></td>
<td>f: 411.5 kg</td>
<td></td>
<td>f: 50.4</td>
<td>f: 32.2 kg</td>
<td>f: 54.4</td>
<td></td>
<td></td>
<td>f: tFA in CC (p &lt; 0.007)</td>
</tr>
<tr>
<td>Moeller et al. [24]</td>
<td>9.7 years</td>
<td>NS</td>
<td>14/4</td>
<td>33.1</td>
<td>nil</td>
<td>14/4</td>
<td>26.2</td>
<td>tFA in CC</td>
</tr>
<tr>
<td>Gruber and Yurgelun-Todd [25]</td>
<td>age of onset 14.1 years, ≥ 4,000 times</td>
<td>currently using</td>
<td>8/1</td>
<td>26.8</td>
<td>nil</td>
<td>8/1</td>
<td>26.2</td>
<td>tFA in CC</td>
</tr>
<tr>
<td>DeLisi et al. [26]</td>
<td>not current use, ≥1 year of use when aged ≥ 18</td>
<td>NS</td>
<td>9/1</td>
<td>21.1</td>
<td>nil</td>
<td>9/1</td>
<td>23.0</td>
<td>f: tFA in CC</td>
</tr>
<tr>
<td>Arnone et al. [27]</td>
<td>8.6 years duration, mean first use aged 15.3 years</td>
<td>currently using (daily)</td>
<td>11</td>
<td>25.0</td>
<td>nil</td>
<td>11</td>
<td>23.0</td>
<td>f: tFA in CC</td>
</tr>
</tbody>
</table>

G = Genu; S = splenium; NS = not specified.

Abnormal alcohol intake correlated positively with MD and negatively with FA in the total CC (p = 0.005 and 0.01, respectively). Pfefferbaum et al. [23] studied 57 alcoholics (40 men and 17 women) and 74 controls (32 men and 42 women). Both alcoholic men and women had FA deficits in the all CC (p = 0.0001) with lower FA in alcoholic men compared to women. MD was higher in both alcoholic groups in the all CC (p < 0.0008). Correlational analyses revealed that in the alcoholic groups, the smaller the CC, the lower the FA and the higher the MD. The genu was the CC area more affected (p < 0.009). Advancing age predicted smaller callosal area, lower FA, higher MD in the CC of the all alcoholic group (p = 0.018 and 0.0003, respectively) and alcoholic men (p < 0.026 and 0.0007, respectively), but not in alcoholic women (p > 0.05). The genu and splenium areas were generally more affected. In the men, greater lifetime alcohol consumption correlated with smaller splenium area (p = 0.02).
Cocaine Dependence
Moeller et al. [24] studied 18 cocaine-dependent subjects and 18 healthy controls (14 men and 4 women in each group). All subjects were screened for medical disorders, psychiatric illness and severity of drug use using structured interviews. Results of the DTI showed significantly reduced FA in the genu (p = 0.007) and rostral body (p = 0.004) of the anterior CC in cases. The initially significant negative correlation between FA in the genu and years of self-reported cocaine use (p = 0.03) and years of alcohol use (p = 0.007) lost its significance when controlled for age (p > 0.05). Controlling for use of amphetamines did not affect results (p > 0.05).

Cannabis Use
Gruber and Yurgelun-Todd [25] compared 9 cannabis users (8 male and 1 female) with 9 healthy matched controls. Psychopathology was excluded by using standardised criteria. Medical conditions and current use of other substances than marijuana were excluded. Users were selected on the basis of lifetime consumption of cannabis (≥4,000 times) and positive urine test to cannabinoids. The study aimed to elicit possible abnormalities detectable with functional MRI in executive functions with the use of the Stroop test. Also DTI analysis was performed as an adjunct. DTI results showed no difference in FA in the CC between cases and controls but a trend towards an increased trace in marijuana smokers (p = 0.09). DeLisi et al. [26] studied 10 subjects (9 female and 1 male) who used cannabis during their adolescence (≤18 years) for ≥1 years between 2-7 times a week (≥21 times/year) and 10 healthy controls matched for sex and age and social class. All 10 cases were non-users at the time of scanning. Three cases also used other illegal drugs in the past and were frequent alcohol consumers. All subjects were selected according to diagnostic interviews and mental illness was excluded. Results of the DTI did not show significantly reduced FA in the CC in adolescent cannabis users but FA was increased in the left anterior cingulate, right medial frontal gyrus, left precentral gyrus, right inferior parietal, right cingulated gyrus and left superior frontal gyrus (p < 0.01, cluster size > 200 mm³).

The authors concluded that findings did not support pathological changes in the white matter of individuals who used marijuana at least moderately during adolescence. Arnone et al. [27] studied 11 male, right-handed, heavy cannabis users with a ‘whole brain’ approach. Subjects were well matched for gender, age, handedness, and other substances, including alcohol, were controlled for. Strict exclusion criteria included physical and mental illness. To measure whole brain white matter structural integrity, a histogram analysis was used. In marijuana users, median FA was decreased in comparison with controls (p = 0.03). Preliminary findings using region-of-interest analysis in the whole CC did not find statistical difference between cases and controls (p > 0.05). These preliminary results do not support the possibility of white matter damage in the CC of heavy cannabis users but further analysis is currently underway.

Discussion
CC in Chronic Alcoholism
Neuropathology. DTI results across the studies confirmed white matter pathology in the CC [19–23]. Consistent findings were a reduction in FA and IC in the CC and more significantly in the genu and the splenium (genu > splenium). Studies which also considered MD reported increased values in alcoholics with a negative correlation between MD and FA particularly in the genu [21–23]. The genu and the splenium connect left and right frontal sites and parietal and occipital sites, respectively [29]. These observations are in agreement with white matter shrinkage reported in post-mortem [30] and MRI [31] studies. It is therefore possible that DTI changes in genu and splenium result from compromised primary directionality of fibres in these areas which can be caused by oedema, demyelination, and axonal loss [16]. Animal models of stroke postulated that a strong negative correlation between FA and MD would support extracellular white matter damage; conversely a less strong correlation would be in keeping with intracellular damage [31]. In this context, white matter disruption in the genu of the CC would be commonly a consequence of extracellular pathology, e.g. oedema, whereas the splenium could be damaged by intracellular cytoskeletal damage. The latter would be in agreement with down-regulation of genes responsible for cytoskeletal proteins as observed in animals [32].

Gender Differences. Women alcoholics showed decreased FA [20, 23], IC [20] and a higher MD [23] in the CC compared to controls particularly in the genu [20]. Compared to men, women alcoholics showed no involvement of the splenium of the CC [20, 23]. These findings suggest that the degree of white matter abnormality in the CC may be present in wider regions of the CC and that in women alcoholics it is likely to be less pronounced compared to men and possibly more difficult to detect with conventional MRI techniques. Sex differences in axonal

Arnone/Abou-Saleh/Barrick
density have been described in some areas of the CC [33]. The genu in women may be more prone to focal damage and therefore detectable with DTI. Differences could also be explained by a number of other variables not controlled for, e.g. different magnitude of lifetime alcohol consumption in women, younger age, and possible differences in the metabolism of alcohol.

**Lifetime Alcohol Consumption.** Lifetime alcohol consumption generally correlated positively with MD, and negatively with FA and C [20, 22, 23].

**Time to Last Drink.** One study reported a significant positive correlation between genu FA and time to last drink [19]. This finding underlies a possible acute toxic effect of alcohol which may be detected with DTI rather than chronic changes. Another important effect which could account for this result is the documented increase in brain volume associated with abstinence capable of affecting MD and FA [34].

**Effect of Age on Alcoholism.** DTI showed that in the genu and splenium of the CC of older alcoholics had greater shrinkage for their age than did younger alcoholics [21–23]. As reported elsewhere [35], age is an independent variable correlated with a natural decline in FA and enhanced MD in normal subjects in the CC equally in men and women. This was observed in controls [e.g. ref. 22]. In alcoholic patients, age was positively correlated with alcoholism, suggesting that the combined effect of increasing age and chronic alcoholism can potentiate the effect of either alone, but also suggesting a greater effect of alcohol consumption over age in alcoholics on the disruption of brain white matter.

**CC in Cocaine Users**

Moeller et al. [24] found significantly lower FA in the genu and rostral body of the CC compared to controls. The initial negative correlation between FA in the genu and years of self-reported cocaine use/years of alcohol use lost its significance when age was controlled for. Although the authors controlled for the effect of amphetamine use, use of other substances was not controlled for and may have potentially affected the validity of their findings. Also, other variables, e.g. handedness and gender, were not considered in the analysis. With all these limitations, this study suggests reduced integrity of anterior CC white matter, supporting frontal cortical impairment in cocaine-dependent subjects.

**CC in Cannabis Users**

Gruber and Yurgelun-Todd [25] published the first study using DTI in cannabis users which showed no differences between cases and controls in FA in the CC. There was a trend towards increased FA in the CC suggestive of the effect of possible confounders. Alternatively, since the study aimed to show a possible derailment in executive functions in cannabis users with fMRI, it might have suffered the limitations of a non-ideal design or underpower. DeLisi et al. [26] did not find significant differences in FA in the CC of subjects who used marijuana during their adolescent years. Some users also consumed significant quantities of alcohol and other substances potentially able to affect the overall results. Subjects were only mild to moderate users and it was not possible to control for physical and psychiatric co-morbidity. It is arguable that based on the time lapsed from the last use to DTI scanning, at least some of the neurotoxic effect of cannabis might have recovered. A cluster size of >200 mm³ might have significantly introduced sufficient background noise to justify the detection of an increased FA in some brain regions. Arnone et al. [27] did not find decreased FA in the CC despite encouraging evidence using a whole brain histogram approach. If confirmed with further analysis, this finding is suggestive of the fact that the magnitude of white matter damage in the CC of heavy cannabis users could be small if present at all. This study not only included subjects who were current users but also targeted individuals who began to use marijuana during their adolescence, the most sensitive neurodevelopmental phase to neuropsychiatric changes.

**Implications for Further Studies**

DTI shows good potential for improved detection of white matter pathology in vivo which antecedes gross morphological changes identifiable with conventional MRI techniques. However, it is conceivable that these studies did not have enough power to detect consistent results as shown by the modest number of subjects included in these reports (table 1). In chronic alcoholism the majority of the studies were published by the same group (table 1). The main shortcoming of this observation is the overall decreased generalisability of the results. Most of the studies attempted matching at least for age when IQ, gender, education, polysubstance use, and handedness are all capable of affecting FA parameters [36] and were not always controlled for. Patient samples might have been different with respect to the severity and duration of addiction, variation in length of abstinence. There may also be some differences in DTI methodology and only very few studies used ‘optimised’ gradient schemes [e.g. ref. 23]. All the studies adopted a region-of-interest approach. Although this is an acceptable ap-
proach, the demarcation of the regions selected is subjective; this should be guided by unambiguous criteria and with demonstrated interrater/intrarater reliability. The risk of a systematic placement bias was minimised by choosing a very identifiable region like the CC. Effect size is difficult to assess because DTI is a novel technique and still under experimentation. Future research could employ larger, homogeneous patient groups selected according to clearly defined diagnostic criteria, whose substance use could be quantified with validated scales. The time from last use of the substance to time of scanning may potentially be crucial in differentiating acute from longer-lasting changes and should be controlled for. A whole brain approach may prevent occurrence of systematic bias but would probably require a larger sample size. To further clarify the mechanisms of white matter damage, its possible reversibility, and to exclude/confirm illness progression, DTI could be employed in longitudinal studies to show modification of neuropathological findings during different stages of dependence (current use/abstinence/relapse). This would also allow the study of the effectiveness of treatment in ameliorating white matter integrity like, for instance, adequate nutrition, vitamin supplementation, and pharmacological treatment.

**Conclusion**

In patients with AD, DTI showed white matter abnormalities in the CC consistent with compromised primary directionality of fibres. A regional pattern was reported with genu and the splenium particularly affected. Women appeared less vulnerable to white matter disruption. The effect of alcohol was amplified by increasing age and longer lifetime consumption. Time to last drink and length of abstinence prior to scanning demonstrated potentials for affecting results. In cocaine dependence, results showed reduced FA in the genu and rostral body of the CC which lost significance when age was controlled for. There is not sufficient evidence to support white matter damage in the CC of marijuana users.

**References**

Diffusion Tensor Imaging of the Corpus Callosum in Addiction


