A Perspective on Cocaine Induced Stroke - Its Mechanisms and Management

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Received date: September 20, 2019; Accepted date: October 15, 2019; Published date: October 16, 2019


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Abstract
Cocaine misuse has been in the news recently with the Department of Health in the UK, recognising the seriousness of its epidemic

Keywords: cerebrovascular disease; drugs; intoxication; ischaemia; haemorrhage; cocaine; management

Interaction
Cocaine misuse has been in the news recently with the Department of Health in the UK, recognising the seriousness of its epidemic. It is the most recurrent agent in drug-related strokes. [1,2] It has been linked to a poorer prognosis, with increased mortality and morbidity in young patients in comparison to other age-matched, drug-free stroke victims. [1, 3] The emergence of cocaine abuse as a major risk factor for stroke in young patients is linked to its consumption reaching worldwide epidemic proportions in the last two decades. In our own practice we have recognised it increasingly as a causative factor in young stroke patients. Its role in provoking stroke should be the subject of greater public awareness. Patients and health care professionals should be educated about the hazards and complications of cocaine use. This perspective contemplates the cerebrovascular effects of cocaine and their management.

Some facts and figures: The United States of America is reported to have have 1.9 million current cocaine users in 2008 with adults aged 18 to 25 being the most prominent age group [4]. Recent data from the United Kingdom reported that approximately 2.2% (or 0.7 million) aged between 16 to 59 abused cocaine in the year 2011 and 2012 rendering it the second most commonly used drug after cannabis [5]. In Australia the 20-29 year age group is the highest abuser of the drug [6]. There are a growing number of cocaine users in Asian countries such as India, Malaysia and Singapore; however detection is difficult as many abusers in these regions are from the higher socio-economic class [7].

Some historical facts: The natural alkaloid is extracted from the leaves of an Andean shrub, Erythroxylon coca or E. novgranatense. The leaves were chewed by indigenous populations of north-western South America to counter fatigue and hunger and also used to prepare a tea to treat mountain sickness. [8]

In Europe the first appearance of cocaine was in the form of extracts from the coca leaf. Vin Mariani, a coca wine was produced in France before 1870 and advocated for lassitude, melancholy and as a general tonic for the body and the mind. Another The coca drink, coca cola, contained 2.5 mg of cocaine per 100 ml in 1900, the last year of its presence in this product. Cocaine extracts were also available as pastilles, elixirs and chocolate bars prepared by pharmacists [9].

In 1880’s it was reported as a cure for alcoholism and morphine addiction. The general enthusiasm for cocaine and its actual euphoriant properties together with the absence of a legal frame regulating its marketing in the United States paved the way for the first cocaine epidemic in 1910. In the UK, cocaine fell under the Pharmacy Act of 1868, which limited its availability to pharmacists and physicians [9].

After an almost complete eradication of the cocaine epidemic by the 1930s, the drug started to reappear in the US in the 1960s and reached epidemic proportions in the 1980s with the appearance of cocaine free base or “crack”.

Pharmacokinetics: Cocaine is absorbable from all body mucous membranes. [10] It is metabolised into inactive water-soluble ecgonine methyl ester by liver esterases and serum pseudocholineseterase. Spontaneous non-enzymatic plasmatic hydrolysis leads to the formation of benzoylecgonine, then ecgonine. These hydrolytic processes account for 80-90% of urinary cocaine metabolites. Its metabolites remain detectable in blood or urine for 24 to 36 hours after ingestion.[12,13]

Cocaine and its metabolites can also be detected in human hair, using gas chromatography and mass spectrography. Hair analysis tests are described as highly sensitive and specific in detecting cocaine use even after disappearance of urinary metabolites. [14]

Mechanism of action: Cocaine has local anaesthetic properties and when given systemically, acts as a powerful sympathomimetic agent by blocking the presynaptic reuptake of norepinephrine (NE) and dopamine. [10,13] The rise in synaptic NE was traditionally held responsible for the sharp rise in blood pressure following cocaine use. [13] We now have a better understanding of this mechanism. The baroreceptor reflexes in healthy individuals indirectly regulate the degree of peripheral vasoconstriction by determining the level of central sympathetic outflow. Excessive adrenergic vasoconstriction is due to an inhibition of peripheral NE reuptake coupled with an unrestrained central sympathetic outflow due to absent altered baroreceptor reflexes. It has been speculated that the blood pressure-raising
The anti-muscarinic properties of cocaine on myocytes, explains the suppression of vagal reflexes that should accompany the sympathomimetic effect of cocaine. This also explains why the cocaine-related rise in blood pressure is accompanied by tachycardia instead of a compensatory bradycardia. [16]

Moreover, repeated episodes of vasoconstriction and subsequent ischaemia due to frequent cocaine snorting may cause nasal septal perforation. Our team has previously described several cases with septal perforation due to intra-nasal cocaine abuse [17]. Similarly, several cases of cocaine induced ischaemic myelopathy and spinal cord infarction have been reported. Cocaine as a risk factor for stroke: Drug abuse is one of the most common conditions predisposing to stroke in young adults; [2,18] The first report of a cerebral ischaemia following endocarditis or cocaine related stroke: [18,35]. A study utilised Magnetic Resonance Angiography (MRA) imaging to demonstrate that cocaine was actually responsible for the induction of cerebral vasoconstriction. This effect was dose-related. Repeated exposures result in a higher incidence of cerebral vasoconstriction [36]. Cerebral vasospasm has been suggested as an incriminating factor in cocaine-associated intra-cranial haemorrhage (ICH) with absent vascular pathology at autopsy [37]. Among designer drugs, cocaine is believed to be the one of the largest causes of strokes and it is difficult to separate its role in the event from that of cerebral vasospasm. The causes of cocaine associated vasculitis are not well understood but it has been associated with a progressive symptoms over weeks and an elevated Erythrocyte Sedimentation Rate (ESR) [31,33].

Pathophysiology of cocaine-related stroke: Cocaine has been linked to both ischaemic and haemorrhagic strokes. Alkaloidal cocaine has been associated to both types of stroke in equal proportions, whereas cocaine hydrochloride was associated with haemorrhagic stroke in approximately 80% of cases [18]. Overall cerebral haemorrhages are more frequent than cerebral infarctions independent of the type of cocaine or its route of administration [21]. Various mechanisms are outlined in Table 1.

<table>
<thead>
<tr>
<th>General</th>
<th>Long term</th>
<th>Specific</th>
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<tr>
<td><strong>History</strong></td>
<td><strong>Rehabilitation</strong></td>
<td>Diazepam, Hypothermia, Consideration of use of intravenous magnesium, corticosteroids,</td>
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<tr>
<td>(Patient, Medical records, collateral history)</td>
<td>(Physiotherapist, Speech &amp; language, Occupational therapist, Psychologist)</td>
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<td><strong>Imaging</strong></td>
<td><strong>Cardiovascular assessment</strong></td>
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<td>(CT/ MRI Brain)</td>
<td>ECG, Echocardiogram</td>
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<td>CT or MR angiogram, Carotid ultrasonography</td>
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<td><strong>Correct Hypertension</strong></td>
<td><strong>Neurological assessment with NIHSS (National Institute of Health Stroke Scale)</strong></td>
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<td>(IV Labetaolol, IV Glyeryl trinitrate)</td>
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<td><strong>Thrombolysis</strong></td>
<td><strong>Education / Counselling</strong></td>
<td>Therapy for drug abuse</td>
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<td>(if ischaemic and no contraindication; and within 4.5 hours of onset)</td>
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Table 1: Summary of management of cocaine related stroke

Thrombosis, whether linked to vasospasm, was proposed as a mechanism of cocaine related stroke [18,22]. Cocaine can increase platelet aggregation via an enhanced response to arachidonic acid [23]. Three cases of cocaine-related stroke associated with positive anti-cardiolipin antibodies have been described in two reports [24,25]. Embolism is mentioned as a possible mechanism [26], either following endocarditis2 or cocaine-induced cardiomyopathy [27]. Hypotension with cerebral ischaemia [28] following myocardial infarction [29] or ventricular fibrillation [30] caused by cocaine consumption has been incriminated. There are some reports of cerebral vasculitis associated with cocaine abuse [21,31,34]. Both angiography and brain biopsy were used to demonstrate vasculitis, although angiography does not seem to be very reliable [32]. Small vessels vasculitis has been described in very few cases of cocaine related strokes and it is difficult to separate its role in the event from that of cerebral vasospasm. The causes of cocaine associated vasculitis are not well understood but it has been associated with a progressive symptoms over weeks and an elevated Erythrocyte Sedimentation Rate (ESR) [31,33]. Cerebral vasospasm is thought one of the most likely mechanisms of cocaine-related ischaemic strokes [18,35]. A study utilised Magnetic Resonance Angiography (MRA) imaging to demonstrate that cocaine was actually responsible for the induction of cerebral vasoconstriction. This effect was dose-related. Repeated exposures result in a higher incidence of cerebral vasoconstriction [36]. Cerebral vasospasm has been suggested as an incriminating factor in cocaine-associated intra-cranial haemorrhage (ICH) with absent vascular pathology at autopsy [37]. Among designer drugs, cocaine is believed to be the one of the largest causes
of ICH in young adults [1]. Bleeds were thought to be a consequence of an outburst in blood pressure following the ingestion of the drugs. However cocaine-related ICH is more likely to occur in patients with underlying cerebral vascular abnormalities [1,22]. It is suggested that cocaine predisposes to aneurysmal rupture in smaller aneurysms and at a younger age [38]. In a small study of 13 patients with drug-related ICH, seven were found to have intracranial aneurysms and 3 had arteriovenous malformations. Cerebral angiography should be part of the evaluation of most young patients with ICH1. Moreover, cocaine induced ICH frequently results in intraventricular extensions [39]. Other contributing risk factors predisposing to stroke are platelet activation via CD40L (sCD40L), Neutrophil-Activating Peptide-2 (NAP-2) and 'regulated on activation, normal T cells expressed and secreted' (RANTES) resulting in higher intravascular thrombus formation and early onset atherosclerotic changes [40]. RANTES (CCL5 or Chemokine ligand 5) is a ligand expressed on many haematopoietic and non-haematopoietic cell types and plays an important part in homing and migration of memory T-cells. Vascular changes include abnormalities of the infolding of the tunica interna and disruption of the tunica externa in ischaemic strokes. Several colleagues have reported the presence of inflammatory mononuclear cells cuffing in cortical and meningeal venules with transmural lympho-monocytic infiltration in a 21-year old stroke patient with cocaine abuse: suggesting vessel inflammation secondary to ischaemic changes with no definitive evidence of vasculitis. Table 1:

Mechanisms of ischaemic stroke in Cocaine users:

Vasospasm – Disorder of Calcium-Magnesium homeostasis

Symptomimetic effect of cocaine
Platelet activation
Apoposis of cerebral vascular smooth muscle cells
Cardio-embolism (endocarditis, arrhythmia, cardio-myopathy)
Disorder of cerebral auto-regulation
PRES (Posterior reversible encephalopathy syndrome)
RCVS (Reversible cerebral vasoconstriction syndrome)

Conclusion
There is a great need to highlight cocaine as an incriminatory factor in the modern era in susceptible individuals. Cocaine-induced stroke is common in young people and increasingly recognised in middle-aged individuals as well. Although much is known about cocaine-induced stroke and its pathophysiology, there is still a great need for further research. There is a great need for specific pharmacological interventions at a molecular level to counter the cerebrovascular and other effects of cocaine and reduce the risks of strokes among cocaine abusers. Inevitably, healthcare physicians should encourage abstinence from cocaine abuse to prevent the widespread side-effects of this drug and recurrences of future intracranial events.

Disclosures
The authors have no conflict of interest(s) or financial interest(s) to declare.

Funding and Acknowledgement
The authors have not received any funding for the writing and publication of this manuscript. The authors would like to the Department of Stroke Medicine, Nottingham City Hospital for their experience in managing cocaine related stroke.

Author Contributions
HS Gendeh was involved in design, data analysis, drafting of article and critical revision of article. S Kamath was involved in data analysis, drafting of article and approval of article. SK Munshi and Abdul Rehman Chaudhary were involved in data analysis, drafting of article, critical revision and a
References:


