Drugs Implicated In Seizures and Its Management

Pooja H Vaidya1* and Anup U Petare2
1Specialty Medical Officer, India
2Assistant Manager, Medical Services, India
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*Corresponding author: Pooja H Vaidya, Pharmacology, Specialty Medical Officer LITMMC & GH, Sion, Mumbai, Address: L-12/D-wing/504, Vighnaharta CHS, Pratiksha Nagar, Sion, Mumbai-22, India, Tel: 9664339836 Email: poojahvaidya@gmail.com

Abstract
Drug-induced seizures are a potentially serious adverse effect. This article provides a comprehensive review of drugs that are most commonly associated with seizures as an adverse drug reaction. The epidemiology, pathophysiology, various predisposing factors, and the drug classes implicated in seizures have been extensively reviewed to provide the physicians with treatment alternatives, especially in high-risk cases. Moreover, the review delineates the management of patients with drug-induced seizures briefly. We hope to assist the clinicians by revising data regarding drug class and individual medications associated with seizures and its management.

Keywords: Convulsion; Adverse drug reaction; Drug withdrawal; Generalized tonic-clonic seizures

Introduction
Seizures in a patient taking a medically prescribed drug are a serious adverse drug reaction [1,2]. Drug-induced seizures occur either due to exposure to or withdrawal from a medication, drug or a toxin [3]. The vital drug factors responsible for a seizure include class of drug, dose and route of administration. Whereas, the patient factors that predispose to drug-induced seizures include pre-existing neurological illness, concomitant medical illnesses, old age, liver failure, renal insufficiency, family history of epilepsy, non-compliance with anti-epileptic treatment, use of concomitant drugs, stress, sleep deprivation, and alcohol abuse [2]. These factors together lower the seizure threshold. Majority of these seizures are self-limited and do not cause permanent sequelae. Nonetheless, repeated or prolonged seizure activity can cause irreversible neurological injury and life-threatening complications including hypoxia, hypotension, pulmonary aspiration, hyperthermia, rhabdomyolysis and metabolic acidosis [3]. Data about drugs commonly responsible for drug-induced seizures is limited, particularly in children. Moreover, thorough acquaintance with the likely causative agents would be valuable to clinicians and could potentially guide the therapeutic approach.

Review
Epidemiology
According to the published literature, 6% of new-onset generalized tonic colonic seizures in individuals older than 16 years presenting to the emergency department at a single center over a five-year period were ascribed to drug exposures, excluding alcohol withdrawal which accounted for 17.6% cases [4]. In another study in an urban hospital, 9% of adults treated for status epilepticus had drug-induced seizures [5]. The drugs causing seizures do vary geographically. Two studies in the USA showed that antidepressants were the most common drug class implicated, with bupropion being the most commonly identified drug followed by anticholinergics [6,7]. In Switzerland, mefenamic acid and citalopram were the most commonly implicated drugs in seizures [8]. In Iran and Australia, tramadol overdose was the common cause of seizures and herbicides and insecticides were implicated in developing countries [3]. A study in the USA recognized that the drugs responsible for seizures showed rapid transition from cocaine, benzodiazepine withdrawal, and tricyclic antidepressants to atypical antidepressants [6].

Pathophysiology
Typically, seizure activity results when there is a sudden onset of disparity between the excitatory and the inhibitory forces in the cerebral cortex resulting in uncontrolled neuronal stimulation. The primary neurotransmitters involved are acetylcholine, gammaaminobutyric acid (GABA), and glutamate. Periodic oscillations of these neurotransmitters occur in the thalamocortical circuit and are regulated by serotonergic, noradrenergic and cholinergic brainstem pathways [9]. Drugs can decrease the inhibitory GABA neurotransmission and lead to overactivation resulting in seizures due to membrane depolarization.
[3]. For e.g. isoniazid or cephalosporin overdose causes seizures by decrease in GABAergic neurotransmission [10]. The release of excitatory neurotransmitter glutamate is mediated by three main receptors: NMDA, AMPA/kainite, and metabotropic. Chronic ethanol use leads to an increase in NMDA receptors. On sudden cessation of alcohol use, the increased neuroexcitatory tone is unmasked which may trigger convulsions [11].

However, no single mechanism can explain all cases of drug-induced seizures [3]. Certain drugs cause secondary seizures through indirect effects on brain perfusion, oxygenation or metabolic disturbances. Narcotics induce hypoxaemia and seizures by direct injury to lung parenchyma or by pulmonary aspiration of gastric contents [9]. Carbon monoxide and cyanide interfere with cellular oxygen utilization resulting in hypoxia and seizures. Even electrolyte disturbances such as hyponatraemia, hypomagnesaemia and hypoglycaemia can lead to seizures. For e.g. sulfonylureas induce seizures via hypoglycaemia, 3,4-Methylenedioxymethamphetamine (MDMA) via hyponatremia, salicylates via cerebral oedema [1,2,6].

Some toxins such as strychnine can induce spinal seizures characterized by involuntary muscle contraction, myoclonus, hyper reflexia and opisthotonus without loss of consciousness. Strychnine competively inhibits the action of glycine, a major inhibitory neurotransmitter in the spinal cord and brain stem and results in seizures. Similarly, tetanus toxin prevents the release of glycine from the pre-synaptic membrane and induces seizures [3].

### Drugs Frequently Implicated in Drug Induced Seizures

<table>
<thead>
<tr>
<th>Class of drugs</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-epileptics</td>
<td>Phenytoin, carbamazepine, lamotrigine, tiagabine, vigabatrin</td>
</tr>
<tr>
<td>Antidepressants and Antipsychotics</td>
<td>Tricyclic antidepressants, citalopram, escitalopram, bupropion, SRI, venlafaxine, lithium, chlorpromazine, phenothiazine, clozapine, olanzapine, quetiapine</td>
</tr>
<tr>
<td>Analgesics</td>
<td>Propoxyphene, tramadol, mefenamic acid, salicylates, meperidine, Phenylbutazone</td>
</tr>
<tr>
<td>Drugs of Abuse</td>
<td>Cocaine, amphetamines, MDMA, phenycyclidine, ketamine</td>
</tr>
<tr>
<td>Withdrawal</td>
<td>Ethanol, baclofen, sedatives – hypnotics</td>
</tr>
<tr>
<td>Miscellaneous agents</td>
<td>Methylxanthines, isoniazid, anticholinergics, organochlorine pesticides, organophosphate pesticides, camphor, lindane, nerve agents, carbamates, chlormequin, quinine, asphyxiants, iron</td>
</tr>
<tr>
<td>Natural Substances</td>
<td>Gyomitra esculenta (mushrooms), jimson weed (Datura stramonium), ephedra</td>
</tr>
</tbody>
</table>

The important drug related characteristics that contribute to the condition include intrinsic epileptogenicity of the substance, dose, route, and central nervous system (CNS) levels. Drugs with a high lipid solubility, low molecular weight, and low protein binding and weakly polar are more likely to enter the CNS [2,9]. Drugs implicated in instigating seizures are broadly classified as psychotropic and non-psychotropic agents. Seizures may also occur as an indirect effect of antiepileptic drugs, miscellaneous agents and drug-drug interactions. Psychotropic drugs include antidepressants, antipsychotics and anti-epileptics, whereas non-psychotropic agents include narcotics, methyldxantines, anticholinergics and several miscellaneous drugs as described herewith (Table 1) [1].

### Psychotropic agents inducing seizures

a) **Anti-epileptics**: Occurrence of seizures after anticonvulsant overdose is a rare finding and can occur after exposure to phenytoin, carbamazepine, vigabatrin, tiagabine, and lamotrigine [12]. The elderly are at a higher risk of confusion and medication misuse, thus increasing the likelihood of seizures. Moreover, exacerbation of pre-existing seizures may ensue because of acute or chronic toxicity, sudden withdrawal or by an indirect mechanism. For e.g. Carbamazepine can lead to seizures via inappropriate anti-diuretic hormone secretion and hyponatremia [12].

b) **Antidepressants and Antipsychotics**: Incidence of seizures at therapeutic doses of antidepressants and antipsychotics range from 0.11.5%. In overdose, the risk increases to 4 to 30%. Skowron et al. [13] has categorized the antidepressants in order of their probability to precipitate seizures (Table 2).

c) **Tricyclic antidepressants (TCA)**: Most of TCA induced seizures are associated with acute overdose with an incidence of 10% with drug overdose. The mortality rate from overdose remains significantly more up to 10% than other antidepressants. A study of TCA induced seizures has shown that seizures are generalized, brief, and occur within 1.5 hours of ingestion. Sustained seizures may occur in up to 17% cases of TCA overdose. Of all the TCA's, clomipramine has been reported to have a greater seizure risk at doses greater than 300 mg/day [14].

d) **Bupropion**: A monocyclic antidepressant was initially withdrawn from the U.S. market due to possibility of seizures, but was reintroduced later. Thundiyil et al [7] has demonstrated that bupropion is the most common cause of new onset seizures attributable to drug exposures. In case of bupropion overdose, the seizure incidence ranges

**Table 2: Seizures induced by Antidepressant.**

<table>
<thead>
<tr>
<th>Probability of inducing seizures</th>
<th>Antidepressants</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Clomipramine, amoxapine, maprotiline, bupropion</td>
</tr>
<tr>
<td>Intermediate</td>
<td>Amitryptiline, imipramine, desipramine, nortryptiline, protryptiline, doxepin</td>
</tr>
<tr>
<td>Low</td>
<td>Fluoxetine, sertraline, paroxetine, fluvoxamine, trazodone</td>
</tr>
<tr>
<td>Minimal</td>
<td>Tranykypromine, phenelzine</td>
</tr>
</tbody>
</table>

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from 11.15% and this rate was highest among those taking an extended release preparation. A significant number of patients experienced seizures greater than 8 hours post ingestion [15].

e) Citalopram and Escitalopram: Seizures occur in 515% of overdose cases. Citalopram induced QT prolongation; sedation, bradycardia, and hypotension indirectly cause convulsions [16].

f) Venlafaxine: Seizures are observed in 0.26% of patients at therapeutic doses, whereas in overdose, it has dose dependent proconvulsant effects with an incidence up to 14%. Doses in the range of 9001500 mg are associated with seizures.

g) Antipsychotics: First generation antipsychotics have a low incidence of seizures. A cohort study done retrospectively showed that second generation antipsychotics carry significant seizure risk. The seizure incidence appears to be highest with clozapine (2.8%) and olanzapine (2%). Clinical reports imply that haloperidol, molindone, pimozide, thioridazine, thiothixene and risperidone exhibit lowest seizurogenic effects. Factors implicated in the occurrence of seizures in patients receiving antipsychotics include elderly age group, high dose therapy and rapid titration. Moreover, risk of seizure is greater in epileptics as competitive enzyme inhibition by the anti-psychotics decrease anticonvulsant serum concentrations and can precipitate convulsions [1].

h) Specific stimulants: Cocaine, neither amphetamines, MDMA exhibit proconvulsant effect probably attributed to increase in nor epinephrine and serotonin levels. Stimulant induced seizures are associated with a higher mortality rate. A case series determined that 3 out of 7 deaths were attributed to stimulant abuse induced seizures [7]. Cocaine can trigger seizures in patients with epilepsy and alcoholic patients during the process of detoxification. Though less common, but a single dose of amphetamines or analogous substance (e.g., ephedra) can trigger seizures cluster. Amphetamines are the fifth leading cause of drug induced seizures due to their direct effects on serotonin and indirect effects of hypernatremia. Mortality due to MDMA is directly linked with number of seizures [17].

Non-Psychotropic Agents Inducing Seizures

i. Narcotics: An estimated incidence of seizures in opioid abusers in 12.5%. Seizures occur in as many as 20.2% of patients presenting with propoxyphene abuse. Of these seizures, 87% were generalized tonic-clonic and typically manifest in 2 hours. The metabolite of meperidine, normeperidine, is highly seizurogenic. There exists an increased risk with high doses or in patients with compromised kidney function [18]. In case of Tramadol, seizures occur not only in overdoses but even in therapeutic dose. The convulsions are not dose dependent. A study revealed 13.7% incidence of seizures in patients on tramadol and chronic use increased risk. Seizure activity also has been reported when reversing the effects of tramadol by using naloxone [19].

ii. dMethylxanthines: Seizures can occur even at therapeutic doses and are more likely to occur with serum levels of greater than 100 mg/dL in acute exposures and 60 mg/dL in chronic toxicity. Preclinical data suggest that blockade of the adenosine A1 receptor is the mechanism for theophylline induced seizures [20].

iii. Isoniazid: In case of overdoses, deficiency of pyridoxine which is required to convert glutamate to GABA is suggested as the causative factor for seizures. A retrospective review of 52 cases of INH overdoses reported that seizures were found in 100% cases. A natural substance commonly known as the false morel mushroom (Gyromitra esculenta) acts by similar mechanism. It is metabolized into monomethyl-hydrazine, which is structurally similar to isoniazid and results in functional depletion of pyridoxine and vitamin B6 [10].

iv. Anticholinergic Drugs: These are reported to account for up to 10% of drug implicated in seizures [7]. Diphenhydramine is the most commonly ingested anticholinergic agent and can cause seizures in overdose. Jimson weed (Datura stramonium) is a common weed consumed for its hallucinogenic and euphoric effects in the United States, and toxicity often results due to intentional ingestion by teenagers. It contain the belladonna alkaloids atropine, hyoscyamine, and L-scopolamine causing anticholinergic toxicity and seizures [7,9]. Household toxins like camphor and phenol can be ingested accidentally. Khine et al. [21] reported a cluster of seizures induced by camphor in children associated with illegally sold or imported camphor products.

v. Local Anaesthetics: Toxicity can be seen when excessive doses are administered eg. Lignocaine in doses exceeding 4.5 mg/kg. Higher dose symptoms include CNS excitation, seizures followed by respiratory depression, coma [22].

Drug withdrawal inducing seizures

The common drugs triggering seizures due to sudden withdrawal include mainly alcohol and sedative-hypnotics. Drug withdrawal results in decrease in GABA and subsequent loss of NMDA receptor inhibition. The result is increased glutamate stimulation, excitatory and a hyperadenergic state with potential for seizure activity. In ethanol withdrawal, seizures occur typically within 6-48 hours after cessation of drinking. The onset of symptoms after benzodiazepine withdrawal is not as predictable due to varying half-lives and pharmacokinetics of the various agents [17].

Clinical Presentation of Drug Induced Seizures

Differentiating drug and toxin induced seizures from other causes is a difficult task unless there is a history of overdose of any specific drug. In conditions wherein there is no history of...
epilepsy, patient is not hypoxic or hypoglycaemic, the physician should maintain a high index of suspicion to consider drugs or toxins as the aetiology. Conversely, in case of focal seizure, if there is no alteration in level of consciousness or a post ictal period, then the seizure is unlikely to be drug related. there is suspicion for drug induced seizures, the patient’s past and family history, current illness and certain clinical clues can prove useful. The patient’s access to medications should be assessed. Previous medical history of tuberculosis or epilepsy may suggest isoniazid or tiagabine induced seizures respectively. Prolongation of QRS on ECG points towards TCA, propoxyphene, venlafaxine, or diphenhydramine overdose. A sympathomimetic toxidrome prior to seizure activity suggest stimulant induced seizures or drug withdrawal. Serum levels of certain medications or toxins and sometimes drug screens may help clarify the aetiology
[2,9,23].

Management of Drug induced seizures

Majority of drug induced seizures clinically present as generalized tonic-clonic motor activity. It is often self-limited. However, prolongation of the convulsive muscle activity can lead to hypoxia, hypercarbia, pulmonary aspiration of gastric contents, lactic acidosis, hyperthermia and rhabdomyolysis. Initial treatment consists of airway management with adequate oxygenation and ventilation, stabilization of the blood pressure and heart rate and rapid testing of serum glucose concentration and benzodiazepines. Intravenous lorazepam is preferred initial benzodiazepine, although intravenous midazolam is also widely used. If intravenous access is not readily available, and then midazolam, being water soluble can be given intramuscularly [3]. Pyridoxine (vitamin B6) is an essential cofactor in GABA synthesis and effectively restores GABA synthesis. It is the drug of choice for seizures due to suspected isoniazid toxicity, and can also be used in poisoning by certain Gyromitra mushrooms [10].

If benzodiazepines are ineffective, barbiturates are recommended as the second line treatment. The drug of choice is phenobarbital. Studies have reported effectiveness of barbiturates in fluvoxamine-induced seizures resistant to benzodiazepines and phenytoin. Moreover, preclinical studies also suggest superiority of phenobarbital over phenytoin in prevention of theophylline-induced seizure and death [3,9,23]. Propofol is an alternative second line treatment as it may have an additive or synergistic effect when used with benzodiazepines or barbiturates. The efficacy to antagonize the NMDA receptor provides an advantage in case of seizures secondary to increased NMDA activity. However, relatively high cost, potential to cause hyperglycaemia, propofol infusion syndrome and neuroexcitatory events limit its use. Hence, it is usually reserved for patients with refractory status epilepticus [3]. The doses of the first and second line anticonvulsants for the treatment of drug induced seizures is as given in the (Table 3).

Table 3: Anticonvulsants for drug induced seizures.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Initial/Loading dose</th>
<th>Continuous infusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diazepam</td>
<td>5 – 10 mg IV (children: 0.2 to 0.5 mg/kg) over 2 to 5 min (max 10 mg/day); may repeat every 5 – 20 min</td>
<td>Note: contains propylene glycol</td>
</tr>
<tr>
<td>Lorazepam</td>
<td>2 - 4 mg IV (children: 0.05 to 0.1 mg/kg, max 4 mg/day); may repeat every 5 – 10 min (max rate: 2 mg/min)</td>
<td>Note: contains propylene glycol</td>
</tr>
<tr>
<td>Midazolam*</td>
<td>I.V.: 0.05 – 0.2 mg/kg (children: 0.1 – 0.3 mg/kg) over 20 – 30 sec (max 10 mg)</td>
<td>0.05 to 2 mg/kg/hr titrated to EEG</td>
</tr>
<tr>
<td>Pentobarbital</td>
<td>I.M.: 0.1 – 0.2 mg/kg (max 10 mg)</td>
<td>0.05 to 2 mg/kg/hr titrated to EEG</td>
</tr>
<tr>
<td>Phenobarbital</td>
<td>5 – 15 mg/kg I.V. (children: 3 - 15 mg/kg) no faster than 1 mg/kg/min</td>
<td>Note: contains propylene glycol</td>
</tr>
<tr>
<td>Propofol $</td>
<td>15 – 20 mg/kg I.V. no faster than 1 mg/kg/min. An additional 5 – 10 mg/kg dose may be given 10 min after initial dose</td>
<td>1.5 – 10 mg/kg titrated to EEG</td>
</tr>
<tr>
<td>Thiopental</td>
<td>1 -2 mg/kg I.V.</td>
<td>0.5 – 5 mg/kg/hr titrated to EEG</td>
</tr>
<tr>
<td>Thiopental</td>
<td>2 – 7 mg/kg I.V. no faster than 1 mg/kg/min</td>
<td>0.5 – 5 mg/kg/hr titrated to EEG</td>
</tr>
</tbody>
</table>

*Consider I.M. route when there is no I.V. access.

$ - Propofol is not recommended for infants and young children.

The role of phenytoin in drug induced seizures is questionable and not recommended. Pre-clinical studies and

Figure 1: Recommended treatment approach for drug induced seizures.
case reports have shown that phenytoin did not effectively terminate seizures produced by a variety of substances. Moreover, phenytoin may be harmful and exacerbate cardiac conduction abnormalities when used to treat seizures induced by lidocaine, theophylline, isoniazid, local anaesthetics and tricyclic antidepressants. Phenytoin was also ineffective in preventing recurrent alcohol withdrawal seizures in various studies [3]. Other anticonvulsant such as valproate is recommended for the prophylaxis of clozapine-induced seizures. However, it has been shown to increase the threshold for theophylline-induced seizures in a preclinical study. Ketamine was useful in tetramine poisoning in which seizures were refractory to benzodiazepines and thiopental. Levetiracetam has been reported effective in patients with nerve agent and pilocarpine neurotoxicity. Other potentially effective therapies still in development include adenosine analogues and cannabinoid receptor agonists [3,9].

Conclusion

Awareness about the potential of various drugs to cause seizures is important for all physicians, particularly those in emergency, neurology, or intensive care settings. Premarking studies, case reports of adverse outcomes, post-marketing surveillance and physician alert notices do educate physicians to be aware of the potential hazards of a drug. Moreover, in a therapeutic set-up, identification of patients at increased risk of seizures is an important and potentially preventative step. Those with history or presence of progressive neurological disease, extremes of age, renal impairment (where relevant to the drug pharmacokinetics e.g., antibiotics) and co-administration of other drugs with neurotoxic or epileptogenic potential have an increased risk of seizure precipitation. In an epileptic patient, it is prudent to optimise anticonvulsant drug therapy first. In some cases, anticipatory treatment may be possible. For example, using pyridoxine with isoniazid and avoiding theophylline or clozapine in patients with a history of epilepsy. It is easy to misattribute a changing mental status to other factors in acutely unwell, particularly older patients and in case of doubt, neurological clinical assessment and, if indicated, EEG should be considered. The patients should also be advised to inform their physicians of additions or changes in medications made either by other physicians or by patient using over the counter and/or complementary medicines. Additionally, periodic review of the necessity for, and the benefits and adverse effects of, the prescribed drugs should be made. Thus, rational prescribing and patient education are the best strategies to prevent drug-induced seizures [1, 2].

References
