Marijuana has been used for centuries for medical reasons. In the early 20th century it was first linked to treatment for epilepsy. Over the last few decades researchers have been unravelling the truth behind the drug. **Prof. Giuseppe Di Giovanni** tells us more about using marijuana for medical research and his own research on this controversial drug.
When I was a high school student my dream was to become a university professor and carry out brain research. I achieved my dreams but I did not plan to end up as a marijuana researcher.

I only recently started studying the effects of chemicals found in Cannabis sativa, better known as marijuana. A few years ago, I was trying to establish a scientific collaboration with a friend and colleague of mine, Dr Roberto Di Maio (Neurology Department, Pittsburgh University), who is interested in epilepsy and marijuana. While researching the topic I found out that many epileptic patients have turned to marijuana to try to control their seizures. I had no idea of the extent of medical use of marijuana in the U.S.

Medical marijuana has many powerful uses. Apart from reducing epileptic seizures, medical marijuana can treat glaucoma, pain, and nausea in cancer and HIV patients, but it also protects the brain from the effects of trauma, eases the spasms of multiple sclerosis, slows the growth of tumours and reduces brain damage in Alzheimer’s disease. It appears that marijuana is effective for all sorts of disorders of the body, mind, and soul.

Considering the bounty of treatments linked to the drug, it is unsurprising that today 23 U.S. states allow the use of medical marijuana. Other states are expected to follow suit and other countries will probably also follow in America’s steps.

The situation is very different for the recreational use of marijuana. The drug has been decriminalised in many countries including Malta (up to 3.5 gr) but legalised in only a few parts of the world (four U.S. states, the Netherlands, and Bangladesh) and is ‘tolerated’ in many others. This is probably due to generalised governmental policy. Marijuana is still listed by the U.S. federal government as a Schedule 1 drug, alongside LSD and heroin. This class has no other drug that has already been accepted for medical use.

Marijuana is moving onto centre stage. Understanding the neurobiology and chemistry behind the increasingly claimed medical benefits of marijuana is vital for scientists and now governments worldwide. At present, the medical uses of marijuana as treatments are still controversial and anecdotal. All pharmaceutical drugs on the market need gold standard large double-blind controlled clinical studies that make sure a drug is safe, or at least safe enough. To date, no such clinical studies have been carried out to examine the beneficial effects of cannabis in different disorders.

Anecdotal evidence is not good enough for treatments. In my laboratory, I am imposing some scientific rigour on what has become a very big ad hoc experiment (Medical School, University of Malta). In Malta, the use of marijuana for research or medical reasons is allowed, but is strictly controlled by national legislation. To get some solid data, I focused part of my research on the effect of cannabinoids (after the plant’s formal name) on different types of epilepsy. I did not use the chemical found in marijuana called delta-9 tetrahydrocannabinol (Δ⁹-THC), which induces marijuana’s psychotropic effects, but a synthetic analogue named win 55,212-2, that is many times more powerful than Δ⁹-THC. To understand how this chemical can help epilepsy sufferers I will explain how the condition arises.

WHAT IT IS EPILEPSY?

Around 60 million people have epilepsy worldwide—in Malta, the number is 3,000. The common neurological condition is caused by recurring disruptions to the brain’s usual activity. The disruption is usually short-lived. The outward signs of epilepsy are known as seizures. Some go unnoticed while others lead to involuntary muscle spasms and loss of consciousness. The type of spasm depends on the part of the brain affected and how far the disruption has spread.

The brain is made up of billions of nerve cells that process information from our senses, thoughts, memories, emotions, and actions, and any (or all) of these activities can be affected by a seizure.
Most seizures are over within a few minutes or less, and the person recovers quickly. Epileptic syndromes are classified as either generalised seizures, which affect the entire brain, or partial seizures, which occur within specific brain regions. Conventional drugs are not fully effective in 30–40% of patients meaning that research is needed to find new treatments.

During my time in Palermo I started to study temporal lobe epilepsy (TLE), the most common form of epilepsy that results in partial seizure in humans. We showed the anti-epileptic role of nitric oxide—a gas produced naturally in our brain. When I moved to Cardiff, I joined Prof. Vincenzo Crunelli’s (University of Cardiff and now also University of Malta) group. We unexpectedly discovered that absence epilepsy is not largely a neuronal disease, but a defect in glial cells (the part of the brain that supports and modulates neurons)—work we published in *Nature Medicine*, one of the world’s most important journals. When I moved to Malta in 2009 I brought this research with me.

**CANNABIS TO TREAT EPILEPSY?**

Cannabis has been used to treat epilepsy for centuries (see table). Unfortunately, marijuana can have some serious cognitive side effects that have been the major obstacle in its medical use. Marijuana impairs perception, reaction time, and short-term memory. Marijuana can be addictive but only mildly so: 9% who try it will become addicted, according to a recent study. The same study listed alcohol at 15%, 17% for cocaine, 23% for heroin, and 32% for nicotine. So although marijuana may be addictive for some, 91% of those who try it do not get hooked.

For our study, we first focused on status epilepticus, a life-threatening condition in which one epileptic fit follows the other without the sufferer recovering consciousness. Dr Roberto Colangeli, a researcher in my group, (now at the University of Calgary, Canada), conducted many experiments that led to one important unexpected discovery. Synthetic cannabinoids only had a modest effect on the development of seizures. When another chemical called serotonin was activated, their effectiveness multiplied, stopping the fits.

Not all types of epilepsy are created equally. In temporal lobe epilepsy, synthetic cannabinoids were even more effective than the epileptic drug phenytoin. The only problem that we found with this treatment was that the dose of cannabinoid impaired the hippocampus, an important part of the brain. The process it blocks is needed for learning and memory. This side effect was very frustrating; our treatment was effective but could harm patients. Dr Stefania Butini (University of Siena, Italy) synthetised a new compound that could block the breakdown of the natural cannabinoids our brain normally makes. The new drug boosted the amount of our own marijuana in epileptics’ brains. The new compound was less effective in stopping epilepsy but was longer-lasting without major memory side effects. We are currently following this line of intervention, trying other drugs that increase the levels of our own cannabinoids when and where they are needed to avoid any possible
MEDICAL MARIJUANA FOR EPILEPSY: THE EVIDENCE

MEDIEVAL TIMES
Arab writer Ibn al-Badri Hashish reportedly uses marijuana to cure the sick son of the Caliphate Council’s chamberlain in Baghdad.

16TH CENTURY
German physician Leonhart Fuchs names the plant Cannabis sativa (A.D. 1542).

19TH CENTURY
British army surgeon William Brooke O’Shaughnessy introduces marijuana into medical practice as a treatment for pain, nausea and convulsions (A.D. 1842).

MODERN TIMES
• In a 1967 study by Martin Keeler and Clifford Reifler marijuana smoking was associated with an increase in seizure frequency.
• In the 1980s, Δ⁹-THC (active ingredient in marijuana) was characterised as an anticonvulsant in animal studies. Convulsions are a common symptom in epilepsy.
• A 1990 epidemiological study by Mervyn Susser and colleagues found that chronic marijuana use protects against seizures.
• According to a 2001 questionnaire conducted by Elisabeth Gordon and Orrin Devinsky and completed by 215 epileptic patients using marijuana regularly, 7.4% experienced a reduction, 2.3% an increase, and 90.2% no change in seizure frequency.
• Recent studies by Robert DeLorenzo’s group (Virginia Commonwealth University) show that endocannabinoids can block epilepsy (status epilepticus and temporal lobe epilepsy) in cell culture and animal models.
• In a 2014 survey, conducted by Orrin Devinsky on parents who had children with epilepsy and who were at the time using cannabis products, 16/19 respondents reported a reduction in seizure frequency while taking medical marijuana; others reported beneficial effects such as improved sleep and mood.
• In a 2015 retrospective survey, conducted by Craig Press and colleagues on paediatric patients with different types of epilepsy who were taking oral cannabis extracts, 57% of respondents reported improvement in seizure duration and frequency.

THE BRAIN’S OWN MARIJUANA
The active ingredient in marijuana comes from a plant, whose extracts would not normally have any affect on the human brain. However, this active ingredient is different. Why? The answer is simple; it happens that the active ingredient in cannabis is very similar to that which the human brain produces naturally. By taking the drug you elevate the chemical levels, switching on the brain to abnormal highs. Other drugs like LSD, opium, caffeine, and nicotine also have analogues that are produced naturally in our brains.

It seems incredible that everyone produces a form of these drugs in their own brain. As mentioned previously, the brain’s own marijuana compounds are called endogenous cannabinoids or endocannabinoids. In 1992 Raphael Mechoulam (Hebrew University, Israel) discovered the first endocannabinoid, naming it anandamide (n-arachidonoylethanolamine or AEA) after the Sanskrit word ‘ananda’ (bliss) and ‘mide’ (chemical). In 1997 Daniele Piomelli (University of California, USA)
discovered lipid 2-arachidonoyl glycerol (2-AG), the other major endocannabinoid in humans. The endocannabinoids (classified as neurotransmitters) pass on signals in our brains and bind to proteins called receptors found in neurons. These two cannabinoids bind to the CB1 and CB2 receptors that help to regulate appetite, mood, memory, and many other functions.

CB1 receptors, the more common type, are widely distributed in the brain, with high concentrations in regions of the brain linked to epilepsy, and are involved in pain perception and forming new memories (the cortex and hippocampus). There are low levels of CB1 in the brainstem, where cardiac and respiratory functions are regulated. This is one of the reasons why cannabis does not threaten your heart or breathing with no overdoses ever recorded from cannabis use.

CB2, the other main cannabinoid receptor, is found mostly in the immune system. The immune system triggers inflammation and studies show that marijuana can reduce inflammation. Its presence there interests scientists because the immune system triggers inflammation, and studies show marijuana can have an anti-inflammatory effect.

The active ingredient in marijuana, Δ9-THC, can bind to these CB1/ CB2 receptors to artificially trigger the same mechanisms in the brain. The cascade of events releases neurotransmitters (the molecules that brain cells use to communicate with one another). Suddenly, they make the world seem hilarious and normal foods taste delicious. Under the influence of marijuana, people generally feel happy, relaxed, and introspective, although paranoia and irritability are common, undesirable side effects. This is because the brain concentration of Δ9-THC reached – after smoking a joint or eating marijuana – is much higher than the normal level of endocannabinoids and greatly magnifies the effect. The result is the well-known marijuana high.

I find it difficult to answer the question: is marijuana good or bad for you? I do not think there will ever be a black or white answer, and it will always be somewhere in between. Although many scientists agree that marijuana is safe enough to temporarily alleviate the symptoms of certain medical conditions, both the short- and long-term use of the drug may harm the body and mind. Marijuana’s continued popularity among teenagers raises particular concern because the drug might hinder the ongoing maturation of the adolescent brain.

For medical use, trying to boost our own marijuana (endocannabinoids) levels will probably be safer. The best approach will probably be by blocking the enzymes that break it down. Research is needed to develop these new drugs to treat a host of different brain disorders. Several research groups in the world, including mine in Malta, are seeking answers to reinvent marijuana uses.

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Prof. Giuseppe Di Giovanni (giuseppe.digiovanni@um.edu.mt) is the Coordinator of the Malta Neuroscience Network, who organised the first Brain Awareness Week, from 14–20 March 2016, the global campaign engaging the public with brain research. Prof. Di Giovanni is also the Editor-in-Chief of Xjenza Online, the journal of the Malta Chamber of Scientists www.mcs.org.mt/index.php/xjenza

This is the first article in the Malta Neuroscience Network (MNN) series.
POTENTIAL APPLICATIONS OF MEDICAL MARIJUANA

EPILEPSY
Multiple animal studies have suggested that Δ⁹-THC may inhibit the brain processes thought to cause seizures. High-quality human studies are lacking and this leaves many questions unanswered.

CANCER
The active ingredient in marijuana, Δ⁹-THC, and its synthetic derivatives have been approved by the U.S. Food and Drug Administration to control nausea in cancer patients undergoing chemotherapy and to stimulate appetite in patients with AIDS. Yet it may not be as effective as other recently developed drugs, so marijuana is not considered as a first-line treatment for these symptoms. Numerous recent studies have suggested that cannabinoids might directly inhibit cancer growth.

MULTIPLE SCLEROSIS (MS)
A large trial published in 2012 found that a cannabis extract significantly decreased muscle stiffness and other symptoms associated with multiple sclerosis (MS). A smaller study found that smoking cannabis reduced spasticity and pain in participants resistant to other treatments. Given the few therapies available for MS, a 2011 review concluded that medical marijuana might help manage certain symptoms.

GLAUCOMA
Several studies have found that smoking marijuana lowers pressure inside the eye, relieving glaucoma-related discomfort for about three to four hours. Other pharmaceutical drugs have now been found to be a better treatment than medical marijuana.

PARKINSON’S DISEASE (PD)
Recent studies do not offer a complete understanding of the role of endocannabinoids and cannabinoid receptors in Parkinson’s disease (PD). Research supports the notion that the endocannabinoid system plays an important role in movement disorders like PD. The studies can lead to novel therapies.

STROKE
There is conflicting evidence on the usefulness of cannabinoids in the treatment of stroke—more research is needed.

PAIN AND INFLAMMATION
Marijuana is only slightly better than a placebo in reducing acute inflammation, and it may even increase the perception of pain in some patients. When taken in combination with other medications, various cannabis-derived drugs are moderately effective in reducing chronic pain.