Naltrexone is an oral Opiate and Toll-Like Receptor Antagonist medication that modulates the immune system to improve health outcomes in many diseases: these include autoimmune & inflammatory illness, cancer, hormonal dysfunction & infertility, autism as well as chronic fatigue.

**The Naltrexone History**

**Opiates**
For six thousand years the opium poppy has been used for its sedative and pain relieving properties; botanically it was known as somniferum, the 'sleep-bringing poppy'. Opium is rich in alkaloids; as chemical research improved, its main component was extracted (in 1806) and named morphine after Morpheus the Greek God of dreams. Later diamorphine, also known as heroin, was synthesized. In 1939 pethidine was discovered and not many years later methadone formulated. They are all known as the opiates which relieve pain but in excess are harmful and extremely addictive, suppressing breathing and even cause death. Such adverse effects prompted researchers to find a molecule that could reverse the negative effects of the opiate drugs.

Naloxone was the first opiate antagonist to be discovered; when injected it immediately blocked the effects of morphine. In 1967 an oral medication, initially known as 'Endo1639A', was conceived; this had similar effects and later become known as naltrexone.

**Endorphins**

Opiate drugs mimic the action of the body’s 'natural neuropeptides' known as endorphins. The best known is beta-endorphin which has the greatest analgesic properties, influencing both the central nervous system (CNS, brain) and body's peripheral nervous system (PNS). However endorphines have other biological activities which are not as yet well understood. Under stress, the brain’s hormone control centre (hypothalamus and pituitary gland), releases a precursor protein POMC (known as proopiomelanocortin) which can be broken down in any part of the body to form endorphins.

Like electrical wiring, the peripheral nervous system (PNS) connects all parts of the body to the central nervous system (CNS). Where the nerve junctions or synapses meet, neurotransmitter chemicals are released, sending their information to specific receptor areas that continue the transmission of data. For example, when pain is experienced 'substance P' is released from the pre-synaptic terminal and travels across the nerve junction to the post-synaptic receptor to relay the pain-danger signal. In the PNS, opiate medications and endorphins bind to the pre-synaptic opiate receptors preventing substance-P being released and blocking the pain sensation. But nerve transmission is much more complex than this simple example. There are numerous other neurotransmitter substances that act like substance-P with a variety of opiate receptors (especially mu receptor) to modulate the pain signal.

In the brain-CNS, opiate receptors are everywhere but act differently, increasing the release of the happy chemical dopamine by inhibiting the neurotransmitter GABA (Gamma-Amino-Butyric Acid) which controls the release of dopamine. Increased dopamine has analgesic action and promotes the 'high' which opiate abusers desire.

Note: In 1979 proteins in wheat and milk were shown to have opiate-like effects, known as exorphins, which could be blocked by naloxone. This may explain the 'addictive' nature of these foods.

**Toll-Like-Receptors guardians of the immune system**

Back in 1985, researchers recognised that endorphines influence the immune system, and within a year Toll-Like-Receptors (TLR) were also discovered. TLR are found in white blood cells and the cells of various organs where they perform the first line of defense for the innate immune system against microbial invasion.

Once TLRs recognise a foreign microorganism, small inflammatory molecules, known as cytokines, are liberated to mobilise white cells that fight the invader. TLR is also able to release NF-kappa-B, one of the most potent signaling molecules involved in the response to severe stress, infection and inflammation, as well as autoimmune disease and cancer.

It is now recognised that opiate receptors are similar to TLR and both are inhibited by naltrexone.

Note: Vitamin D also modulates the Toll-like receptors of the innate immune system and can be used in conjunction with Naltrexone.

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**Current knowledge of Naltrexone**

Many diseases are an expression of a malfunctioning immune system.

The immune system is regulated by endorphins which have a primary action on opiate receptors.

The immune system identifies intruders with Toll-like receptors which release cytokines which initiate the white cell response.

A naltrexone capsule is a mixture of both LEFT (levo) and RIGHT (dextro) handed molecules - like our hands, mirror images of the each other.

Levo - Naltrexone action:
- Initially antagonizes the opiate/endorphins receptors
- Leading to upregulation of endorphin release and modify the immune system and
- Reduction in cell proliferation

Dextro ~ Naltrexone action:
- Antagonizes the Toll-like Receptors (TLR) suppressing cytokine response and modifying the immune system, and
- Antagonizes the TLR - 4 involved in release of NF kappa B reducing inflammation & potentially down-regulating oncogenes that initiate cancer growth.
**Naltrexone in clinical practice**

High dose naltrexone (HDN) was first used for opiate addiction, but proved ineffective due to troublesome side effects like anxiety, mood changes and agitation which led to poor patient compliance. However, the Finnish physician Dr Davis Sinclair had success using HDN in alcoholics wishing to stop drinking; this is commonly known as the Sinclair Method of treatment. The first clinician to demonstrate Low Dose Naltrexone (LDN) effectiveness was Dr Bernard Bihari in 1985. He began using LDN in immune-suppressed HIV patients and showed it could lessen susceptibility to infection, prevent the destruction of the immune system and lower death rate. At the same time Dr Ian Zagon starting decades of publishing - over 300 papers on LDN - unravelling the endorphin, opiate receptor and immune system connections.

**Endorphin Deficiency**

The low endorphin levels Dr Bihari found with his HIV patients may also be present in those who cry easily (e.g. watching a TV commercial), have low mood and depression, greater sensitivity of emotions and to pain, desire pleasurable rewards and crave chocolate, tobacco, sex, alcohol and drugs. Some have called this the endorphin deficiency syndrome.

As mentioned earlier naltrexone increases endorphin levels and may be a useful therapy, with lifestyle changes and medications, in conditions such as fibromyalgia, chronic fatigue, and depression and pain states.

Note: Aerobic exercise, massage, meditation, rhodila, chamomile & lavender, omega-3 oils and a healthy high protein diet are known to increase endorphins. The withdrawal of milk and gluten foods, that contain exorphins, might also be worth considering.

Hypothalamic-pituitary hormone disturbance and high cytokine levels from inflammation are known to disturb sleep, but both are normalized by naltrexone. A trial of LDN be useful for insomnia, thyroid disorders, premenstrual syndrome, polycystic ovarian disease, endometriosis, infertility and in obesity.

**Inflammatory and immune disorders**

It is mentioned above that naltrexone influences the Toll-like receptors that directly reduce cytokine-mediated inflammation and also inhibits NF-Kappa-B signaling which has been linked to the onset of infection, autoimmune & other inflammatory disease and the initiation of cancer. A number of LDN trials have shown positive response in the treatment of Crohn's disease, multiple sclerosis, thyroid disorders and cancer. See table below for diseases that may respond to LDN.

**Side effects**

The side effects of naltrexone therapy are minimal, the most common being sleep disturbance which usually settles when a lesser amount is taken.

**Dr Bernard Bihari’s Story**

In 1985 Dr Bihari was working in New York with HIV/AIDS patients with compromised immune systems. He had no available effective therapies to treat these people, many of whom were drug users. He was aware that endorphins, the natural morphine-like substance which are produced under pleasure and stress states, were involved in the regulation of the immune system.

Dr Bihari's brilliant insight was to initiate a twelve-week trial of LDN in 22 HIV patients with low endorphin levels, comparing them with a non-treated control group. During the trial no one in the LDN group developed opportunistic infections whereas 5 of the 16 controls became ill.

He then went on to actively use LDN in his patients and observed that when naltrexone was taken regularly, it appeared to prevent the destruction of the immune system and lower death rates.

When the new HIV anti-retroviral drugs became available, he noted that LDN acted synergistically to improve the clinical response.

In 2014 Bihari reported on 354 advanced cancer patients who had failed traditional therapy and then received low dose LDN with other possible complementary therapies. He found that 20% of patients had an objective response to LDN therapy and 25% stabilization of their cancer. He noted that most of those who died were terminally ill before they commenced LDN.

Many cancer types responded positively to LDN. An independent review confirmed that LDN alone did appear to invoke complete cancer remission in some patients. A recent study has also shown that LDN heightens the sensitivity of chemotherapy agents, increasing cancer cell death.

For more information watch the interview with Dr Bihari at [www.youtube.com/watch?v=x54Jccr8GT8](http://www.youtube.com/watch?v=x54Jccr8GT8).

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**Conditions that may respond to LDN Therapy**

**Inflammatory states**
- Autoimmune conditions - multiple sclerosis, lupus, rheumatoid arthritis etc
- GI inflammatory bowel disease - Crohn's disease, ulcerative colitis & IBS
- Lung - COPD, asthma
- Cancer

**Hormone dysfunctions**
- Thyroid - hyper- & hypo-thyroid states
- PMS, PCOD, infertility, obesity

**Brain connections**
- Insomnia, depression, ADHD, autism
- Parkinson’s, motor neuron disease, neuropathic pain, restless leg syndrome

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**General Reference:**

**Websites worth exploring:**
[www.lowdosenaltrexone.org](http://www.lowdosenaltrexone.org), [www.ldnresearchtrust.org](http://www.ldnresearchtrust.org)