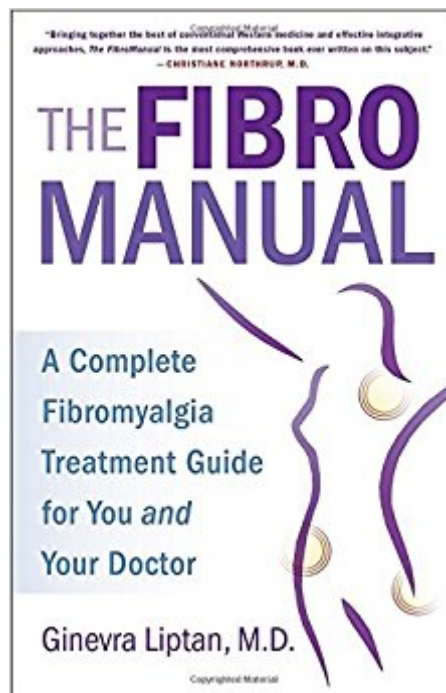




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The FibroManual: A Complete Fibromyalgia Treatment Guide For You And Your Doctor



Synopsis

The most up-to-date, comprehensive treatment guide to fibromyalgia, by a renowned physician who herself has the condition. If you suffer from fibromyalgia and are struggling to get help from your doctor, you're far from alone. Ten million Americans experience the widespread muscle pain, profound fatigue, and fuzzy brain (a fibrofog) that have long frustrated both patients and doctors. In this unique resource, Ginevra Liptan, M.D., shares a cutting-edge new approach that goes far beyond mainstream medical knowledge to produce dramatic symptom improvement. Dr. Liptan's program incorporates clinically proven therapies from both alternative and conventional medicine, along with the latest research on experimental options like medical marijuana. Since many health care providers have limited fibromyalgia expertise, The FibroManual includes a thoroughly sourced health care provider guide that enables readers to help their doctors help them. Alleviate fibromyalgia symptoms in four simple steps (Rest, Repair, Rebalance, and Reduce) and you will restore deep, restful sleep achieve long-lasting pain relief optimize hormone and energy balance reduce fatigue. This accessible and empowering resource provides essential information about understanding and treating fibromyalgia from a physician who, as both patient and provider, understands the illness from the inside.

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Customer Reviews

Ginevra Liptan, M.D., is a graduate of Tufts University School of Medicine, board certified in internal medicine and trained in functional medicine, a holistic approach that blends both Western and

alternative medicine. After developing fibromyalgia as a medical student, Dr. Liptan spent many years using herself as a guinea pig to find successful treatments, and she has fine-tuned her approach by treating thousands of patients. She is one of the few clinical specialists in the world to focus solely on fibromyalgia. She directs the Frida Center for Fibromyalgia and serves as medical advisor to the Fibromyalgia Information Foundation.

9781101967201|excerptLiptan / THE FIBROMANUALChapter 1Figuring Out Fibromyalgia for MyselfAt the end of my first year of medical school, I was doing crunches at the gym when I felt a muscle in the front of my neck rip, causing intense pain. I had injured muscles before, and I figured it would heal quickly. But as days turned to weeks, it didn't. My neck burned constantly and felt like it could no longer hold up the weight of my head. The only relief I got was when I wrapped a heating pad around it. A chiropractor found mild abnormalities in my cervical spine and made some adjustments. I started getting regular chiropractic treatments, because they were all I could think to do, and they helped a little. That summer, I was achy and tired all the time. I took a part-time babysitting job, and working for just a few hours left me exhausted for days. I woke up in the morning with a sore neck and back, which hurt all day. I frequently felt stiff, weak, and lightheaded. It was as if all the energy of my body and mind had been sucked out. I went to see my primary care doctor, who prescribed a muscle relaxant, which did not help. The pain progressed to my upper back - an ache between my shoulder blades that would not go away. My spine hurt, my skin hurt, everything hurt. I was sleeping poorly, tossing and turning, and I woke every morning feeling more tired than before I went to bed. I felt too weak to even lift my arms to wash my hair in the shower. One day my hips started aching so much that I couldn't do anything but lie in bed and cry. I was sure there was something really wrong with me, so I went again to my doctor. She drew blood and sent me to a rheumatologist. The rheumatologist ordered X-rays of my neck and hips and assured me they were completely normal. "So why do I hurt all the time?" I asked. "I don't know, but you don't have arthritis," he responded. I returned to my doctor to find that my labs were all normal. My doctors had no answers for me and nothing to offer. A DiagnosisI started my second year of medical school as a total wreck. And then a bit of grace fell my way. My chiropractor sold her practice. I tearfully told my entire story to the chiropractor who took it over, and he suggested that I might have fibromyalgia - the first time anyone had even mentioned it to me. He checked my tender points, and registered exquisite tenderness in at least twelve of them, confirming the diagnosis. He recommended that I pick up what he called the bible of fibromyalgia, Fibromyalgia and Chronic Myofascial Pain Syndrome: A Survival Manual, by Devin

Starlanyl and Mary Ellen Copeland. I spent weeks reading and rereading it, trying to convince myself that I did not have the condition, but unable to avoid the fact that it described me perfectly. "Chronic" was such an awful word. And fibromyalgia had such a stigma among doctors - even among my fellow medical students - that I couldn't quite believe I had it. I began doing research in my medical textbooks, on the Internet, and in bookstores. I quickly became discouraged. Western medicine had little to offer beyond antidepressants and exercise. I was already on antidepressants, and exercise made me feel worse. So I ventured into the confusing world of alternative medicine, perusing countless theories and treatments online, many of which conflicted directly with what I was learning in school. I was overwhelmed by the different ideas I encountered about the causes of fibromyalgia and how to treat it. Was it yeast overgrowth? Low thyroid hormone production? Did I need to cleanse my body of toxins? Do a raw-juice fast? Use guaifenesin to reduce calcium phosphate deposits in the muscles? Was I deficient in some vitamin that I needed in megadoses? Did I need a macrobiotic diet? Raw foods? Alkaline foods? Compounding my frustration was the fact that I was so fatigued I didn't have the energy to contemplate major lifestyle or dietary changes. Acupuncture didn't help, and massage made me feel more achy and tired. The latter half of my second year of medical school was basically a very expensive correspondence course. I rarely felt well enough to make it to class, and survived by getting lecture notes from friends, studying at home, and showing up only for the occasional required seminar and exams. I finally accepted that I had fibromyalgia. I felt hopeless and helpless. I told myself that it wasn't fatal - it wasn't cancer - but it still felt like a death sentence. It became clear that there was no way I could make it through the rigors of the third year of medical school, with its eighty-hour workweeks and high levels of physical and emotional stress. I took a leave of absence. During my yearlong leave I read every book I could get my hands on about fibromyalgia and tried every treatment I could. I knew exercise was supposed to help, so I kept trying new regimens, only to stop after injuring myself. After diligently following the guaifenesin protocol described in *What Your Doctor May Not Tell You About Fibromyalgia: The Revolutionary Treatment That Can Reverse the Disease*, by R. Paul St. Amand and Claudi Craig Marek, I was devastated when I got no benefit from it. I took thousands of dollars' worth of useless supplements. I tried IV therapies with a naturopath, and saw a top holistic MD in Boston. I did colonic hydrotherapy and fasted and did detoxifying diets. I tried every type of massage I could find, but each session seemed to make my pain worse. I journaled and saw a therapist and tried to meditate. None of it helped, and midway through the year I was feeling hopeless and preparing to drop out of school completely. Then I read Claire Musickant's *Fibromyalgia: My Journey to*

Wellness. Musickant describes her dramatic reduction in symptoms after testing for food sensitivities and eliminating the offenders. I knew it was something I needed to do; it felt right. I had a rare rush of hope and excitement as I called the lab to find a practitioner in my area, a naturopath. She told me the test was expensive. I hesitated and asked her if she had found that it helped people with fibromyalgia. "Oh, yes," she said, "and it helped me. I had fibromyalgia and am now ninety percent better." She had fibromyalgia "À- past tense. These were possibly the most beautiful words I had ever heard. She was the first person I had talked to who had even suggested that fibromyalgia was something one could recover from. I did the blood test and started to avoid the recommended foods and chemicals. After about two weeks, I realized that the all-À- over body ache I had grown so accustomed to was gone. Now it only hurt when someone actually pressed on my muscles. It didn't feel as if I constantly had the flu. I had the energy to grocery shop, to cook, to exercise. I was thrilled. But my sleep was still light and restless and I woke up exhausted every morning. My neck and arm pain flared up easily with exertion or repeated motion. A massage therapist recommended I try a manual therapy technique, the John F. Barnes's Myofascial Release Approach, which involves slow, prolonged stretching that releases restrictions in the fascia, the connective tissue around the muscle (see chapter 12 for further details). After several sessions I experienced dramatic pain reduction in my arms and neck. With ongoing myofascial release therapy, I felt well enough to return to medical school, but I remained fatigued all the time. Things really fell apart during a month of night shifts in the hospital. I couldn't sleep during the day and developed severe insomnia. I dragged myself once again to my primary doctor, who prescribed a sleep medication. Suddenly everything shifted. I was getting deeper sleep and feeling more rested when I woke up. The combination of improving sleep quality with medication, avoiding inflammation-À- producing foods, and myofascial release treatments got me through the grueling remainder of medical school and residency training. And it forms the core of my treatment approach to this day. Over the years I fine-À- tuned my technique, learning other ways to improve deep sleep, treat the fascia, and warm up so that exercise helps and doesn't hurt. As I continued my medical training, I was able to better assess the thousands of studies and articles written about fibromyalgia. I began to put together why certain treatments had helped me so much, while others did nothing at all. Based on my own relief with myofascial release I was convinced that the fascia was the source of fibromyalgia pain, and my research has focused on it ever since. At Oregon Health and Science University I conducted a study that found myofascial release therapy was more helpful than standard massage for fibromyalgia symptoms (Liptan 2013). I've also published articles on exercise and self-À- management strategies (Jones 2012; Liptan 2010; Jones 2009). In 2011 I

founded the first private practice in the United States dedicated exclusively to fibromyalgia, the Frida Center for Fibromyalgia, in Lake Oswego, Oregon. For years after my diagnosis I felt bitter toward all the doctors who had not been able to help me. It was so frustrating that again and again I had to find my own way, wasting money on ineffective treatments because I had no guidance. My resentment faded, though, as I continued my medical training and practice and realized that physicians in this country are working within an untenable system. Our current medical framework is just not set up well to deal with chronic and complex illnesses like fibromyalgia. And while most doctors do really try to help their fibromyalgia patients, they simply lack the expertise or tools to do it. I told no one about my diagnosis, even my closest friends. I only revealed it during my final presentation to colleagues and teachers on the last day of my residency. I felt as if I was *œcoming out* • and revealing a whole secret life. I wanted to show my fellow doctors it is a real disease that can happen to anyone. Since then, I have focused a large part of my career on educating other health care professionals about this invisible illness. I am done feeling ashamed. My hypervigilant nervous system keeps me attuned to subtle changes in other people's emotional states and in my environment. It has helped me to become a better doctor. And my intimate experiences with pain and suffering have made me a better human being.

I was hesitant about this book, as I already have tons of info about FM floating around my head (so I didn't feel I needed an introduction to the topic). But not only is the Fibro Manual comprehensive and current, it provides a very trustworthy roadmap for all aspects of management (pain, sleep, exercise, energy, doctors, etc). I realized while reading it that my approach has been pretty scattershot for the last several years -- and that I was often slowing down my own progress because I didn't realize ways in which my experimentation was undermining my overall efforts. Liptan does a wonderful job of explaining how everything connects (as best we now know), so that we can effectively prioritize and organize our management strategies. Her section on sleep aids and supplements (with advice on how to combine them safely and effectively) is alone worth the price of the book. I've been following the manual pretty religiously for a couple of months now. Of course I'm not "cured," but I feel the most stable I've felt in years, not desperately trying new things every day. Highly recommended.

This is one of the best books available on treating fibromyalgia. A must read for every fibromyalgia patient and treatment provider. This book lives by my desk because I look things up in it constantly. It's required reading for my Fibromyalgia Coach training class. There are a few things in particular I

love about this book: First, Dr. Liptan includes all of the little things that have made the biggest difference in my fibro healing. These are all things that took me years to discover on my own -- and you'll have them all right here! Second, there is a section at the end of each chapter that outlines the things you can do for yourself, as well as the things your doctor needs to help you with. This makes it a very effective book for you to use on your own. Third, there is a bullet point guide at the back of the book specifically for your doctor. It includes the research and studies that support her treatment recommendations. This means that our busy doctors can read the "cliff notes" version to know how to treat you. Many people have asked me how this differs from her first book, *Figuring Out Fibromyalgia*. I'm also often asked if they should buy *The FibroManual* if they've already read the first one. If you're wondering the same thing, here's your answer: *Figuring Out Fibromyalgia* is limited to what we knew about fibromyalgia at the time it was published (January 2011). After six years, we know a lot more. Studies have been done and drugs approved (or not approved!) that have changed our treatment strategies. While *Figuring Out Fibromyalgia* includes more of Dr. Liptan's personal fibromyalgia journey, *The FibroManual* was written with your doctor in mind. As I mentioned above, Dr. Liptan includes bullet points at the end of every chapter and a "cliff notes" section in the back for your doctor with all the research links and information he or she needs to treat you effectively. I highly recommend getting a copy of *The FibroManual*. It truly is a complete fibromyalgia treatment guide that is written for both you and your doctor.

I bought Dr. Ginevra Liptan's new book after reading her first book and benefitting greatly from the information it contained. Her new book is accessible, well-organized, concise, and really covers fibromyalgia treatment, using the overview of Rest, Repair, Rebalance, and Reduce, and breaking it down to the fine details that really make a difference. I am a nurse case manager and have been working in the healthcare field for many years, so I can really appreciate the comments she makes about the healthcare system and the need to be your own best advocate. Her advice regarding how to help your health care providers by organizing and carrying your own records and labs, and providing symptom tracker logs/data summaries among other ideas, is priceless. Many healthcare providers are not very knowledgeable about working with fibromyalgia patients and tend to rely on the very few FDA approved medications as treatment. They do so without an understanding of the importance of effectively addressing sleep and other issues in fibro in a step-wise fashion. The references contained in the book, and the abridged Healthcare Provider Guide to Fibromyalgia Management in the Appendix, provide an evidence basis for the proposed treatments, which will be much appreciated by providers seeking a grounding in evidence-based

practice. The book also contains practical references for patients in terms of their own education and self-care that are very useful, as well as resources. Any fibromyalgia patient working with a willing provider should be able to make substantial progress with symptoms using the information in this book. Dr. Liptan has fibromyalgia herself, so her suggestions and information come from years of heartfelt search for solutions. An absolutely essential guide to fibromyalgia for patients and providers that goes straight to practical treatment strategies. The book is also a useful adjunct for any patient or provider working with chronic pain issues of any sort.

This book has been a huge help to me as I travel the road to find balance with fibromyalgia. My Rheumatologist recommended this book, and I checked it out from the library initially. I found that I had so many bookmarks and notes inserted, because of the large amounts of helpful information, that I just needed to purchase my own copy! Simple to understand, but yet based on Scientific evidence, this book is a great resource.

Very well organized and contains a lot of helpful information. Everyone has a different experience with this disease, but there is good info here.

I felt like I finally met someone who truly understood the frustration of daily life with fibro and could give me advice based on experience. I like that she balanced Western medical knowledge with alternative practices and admits there is no one right way to deal with this for everyone. This made me really want to listen to what she had to say. Highly recommend!

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