Here’s the skinny on compounded “bioidentical” hormone therapy—popular among women but absolutely data-free.

Bio

PLANT- DERIVED BUT CHEMICALLY ALTERED
All bioidentical hormones—both individually compounded formulations and pharmaceutical products—come from the same soy or wild yam precursors before they are chemically converted to the different hormones.
identical” hormones

What you (and your patient) need to know

The Women’s Health Initiative (WHI) caused a sea change in women’s attitudes toward menopausal hormone therapy and aroused many fears—not always rational—that remain almost palpable today. One study of the aftermath of the WHI found that 70% of women who were taking hormone therapy discontinued it, and 26% of women lost confidence in medical recommendations in general.

Into the chaos stepped Suzanne Somers, Michael Platt, and other celebrities, touting the benefits of a new kind of hormone: bioidentical. You don’t have to read Somers’ bestseller, The Sexy Years, to encounter the claims it makes on behalf of bioidenticals; the cover itself makes them clear:

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OBG MANAGEMENT Senior Editor Janelle Yates contributed to this article.
10 erroneous beliefs patients have about compounded hormones

- “They’re identical to the hormones in my body”
- “They occur naturally”
- “They are safer and more effective than conventional hormone therapy”
- “They’re risk-free”
- “They are monitored by the FDA”
- “They are the fountain of youth”
- “They prevent breast cancer”
- “Celebrities know more about them than physicians and menopause and hormone experts do”
- “Doctors oppose bioidentical hormone therapy because they are in the pocket of Big Pharma”
- “Bioidentical hormones are not a huge money-making enterprise”

It’s a false claim that all bio identical hormones are bioengineered to contain the same chemical structure as natural female sex hormones.

What is “bioidentical”?

**OBG MANAGEMENT:** Let’s start with the basics. What does the word “bioidentical” mean? Is it a legitimate medical term?

**DR. PINKERTON:** Bioidentical hormones are exogenous hormones that are biochemically similar to those produced endogenously by the body or ovaries. These include estrone, estradiol, estriol, progesterone, testosterone, dehydroepiandrosterone (DHEA), and cortisol. The FDA has approved many prescription products that contain bioidentical hormones. However, the term “bioidentical” is often used to refer to custom-compounded hormones. The major difference between the FDA-approved prescription bioidentical hormone products and custom-compounded products is that the former are regulated by the FDA and tested for purity, potency, efficacy, and safety.

Bioidenticals are not “natural” hormones, although many consumers think they are. In reality, compounded bioidentical hormones and FDA-approved bioidentical hormones all come from the same precursors. They begin as soy products or wild yam and then get converted to the different hormones in a laboratory in Germany before finding their way to the various world markets.

The claim that all bioidentical hormones are bioengineereed to contain the same chemical structure as natural female sex hormones is false. As one expert noted, “the term ‘bioidentical’ has become inappropriately synonymous with ‘natural’ or ‘not synthetic’ and should be redefined to correct patient misconceptions.”

Common misconceptions

**OBG MANAGEMENT:** What are some of the other false impressions you encounter among patients who ask for bioidenticals?

**DR. PINKERTON:** That the hormones are safer or more effective than hormone therapy, that they carry no risks, and that they are as well-monitored as FDA-approved products, to name a few. (For more, see “10 erroneous beliefs patients have about compounded hormones”).

**OBG MANAGEMENT:** Where do these ideas originate?

**DR. PINKERTON:** They are propagated by self-proclaimed experts and celebrities or by laypersons and physicians who devote the bulk of their time to promoting these hormones, usually at considerable cost to the patient.

**OBG MANAGEMENT:** What are the risks of compounded bioidentical hormones?

**DR. PINKERTON:** According to FDA guidance for industry, in the absence of data about these hormones, the risks and benefits should be assumed to be identical to those of FDA-approved hormone therapies, with the caveat that we don’t know from batch to batch what a woman is receiving. However, they are not regulated or monitored by the FDA and tested for purity, potency, efficacy, and safety.
The problems with compounded hormones

What’s in a name?
“Bioidentical” isn’t a bona fide term. There is no definition of it in any medical dictionary; it’s just a name the industry cooked up, a catchy one at that. And when bioidenticals are advertised and promoted, the term “natural” is usually in close proximity. Most patients equate the word natural with plant-derived substances that have not been chemically altered. The fact is, many compounded prescription drugs are derived from plants—but they are also chemically altered.

Some applications are legitimate
A number of women use compounded medications because they make it possible to obtain hormone combinations that are not readily available in cream form. For example, if a patient wants testosterone as part of a cream of estrogen and progesterone, a compounded product is the only option.

Show me the data
No studies have compared compounded drugs with commercial drugs—and such studies are exceedingly unlikely. Compounding pharmacies have no incentive to conduct or participate in such studies. The pharmaceutical compounding industry is a multibillion-dollar enterprise in this country, and compounded prescription drugs for menopausal conditions are probably the biggest product outside of the oncology arena. Proponents of compounded hormones have a captive audience, so to speak, made up of women who don’t like commercial drug manufacturers or who prefer products that appear to be natural, or both.

The problem is that these women receive no package insert or prescription drug label with their hormones. Warning labels are not required because compounded drugs are not regulated by the FDA. Consumers are basically at the mercy of whatever claims they read on the Internet or in the lay literature, which tends to be written by people who have a financial interest in affiliating with the compounding industry. It’s a very frustrating situation for a lot of people.

Unintended consequences of the WHI
The Women’s Health Initiative (WHI) stirred demand for bioidentical hormones by casting the safety claims for some commercial hormone therapy products in a less than favorable light. That wasn’t the investigators’ intent, of course, and some of the findings of the WHI have since been questioned.

The goal of the WHI was to critically evaluate some of the touted health benefits of commercial hormone therapy prescription drugs, but, by questioning some of these claims, it inadvertently pushed a significant percentage of patients toward compounded prescription drugs—and we have no safety data on them.

No one knows exactly how many women were swayed, but the consensus is that they were, and no one’s been happy about that.

The FDA has nothing against compounding pharmacies per se. Individualized preparation of a customized medication for a patient, based on a valid prescription, is an essential part of the practice of pharmacy. However, some actors in the pharmacy compounding business have taken the practice to a different level, not just in terms of the volume of business they do, but in the way compounded hormones are advertised and promoted. The courts aren’t necessarily interested in intervening in cases involving high volume alone. And when it comes to unsubstantiated claims of benefit, the FDA has found it difficult to assert jurisdiction over pharmacy compounding in general, making it hard to assert control over the advertising claims these pharmacies make on behalf of compounded drugs.

The result? The FDA has been unable to rein in claims that compounded prescription drugs are safer or better than commercially prepared medications. These drugs are probably as safe and effective as their manufactured counterparts, but there are no data to confirm this assumption.

FDA, so we are lacking testing for purity, potency, efficacy, and safety. When the FDA did analyze compounded bioidentical hormones, a significant percentage (34%) failed one or more standard quality tests. In comparison, FDA-approved drugs fail analytical testing at a rate of less than 2%.

The main problem with the compounded hormones, as I see it, is that women who use them do not receive any written information from the compounding pharmacist about risks and benefits. Nor do they receive...
Women who use compounded hormones do not receive the black box warnings that appear on FDA-approved estrogen therapy.

ACOG, NAMS, and The Endocrine Society agree: Compounded hormones are not safer

The American College of Obstetricians and Gynecologists (ACOG), North American Menopause Society (NAMS), and The Endocrine Society have all issued statements noting the lack of safety data on compounded bioidentical hormones. Here’s what they say:

ACOG
“Most compounded products have not undergone rigorous clinical testing for safety or efficacy, and issues regarding purity, potency and quality are a concern. Compounded hormone products have the same safety issues as those associated with hormone therapy agents that are approved by the US Food and Drug Administration and may have additional risks intrinsic to compounding. There is no scientific evidence to support claims of increased efficacy or safety for individualized estrogen or progesterone regimens.”

NAMS
“NAMS does not recommend custom-compounded products over well-tested, government-approved products for the majority of women—and does not recommend saliva testing to determine hormone levels” (www.menopause.org/bioidentical_NAMS.aspx).

The Endocrine Society
“‘Bioidentical hormones,’ particularly estrogen and progesterone, have been promoted as safer and more effective alternatives to more traditional hormone therapies, often by people outside of the medical community. In fact, little or no scientific and medical evidence exists to support such claims about ‘bioidentical hormones.’ Additionally, many ‘bioidentical hormone’ formulations are not subject to FDA oversight and can be inconsistent in dose and purity....”

The black box warnings on FDA-approved estrogen therapy. I believe women need to be adequately educated about the potential risks and benefits, as well as the lack of efficacy data and quality control, if compounded products are requested. That means it’s up to the prescriber to educate the patient about the potential risks and benefits.

Rosenthal states that symptomatic menopausal women or those who fear breast cancer or heart disease can be considered a vulnerable population: "Patients do not have the background to decipher credible sources from noncredible sources.” False claims present convincing arguments for laypersons. A woman may be vulnerable to unsubstantiated claims by virtue of her symptoms and the anxiety and even depression that they can produce. Without comprehensive education, these women cannot be assumed to be adequately informed.

Let me put it in perspective. If a patient with a history of breast cancer complains about severe vaginal dryness that interferes with her sex life, I might decide to give her the smallest amount of topical estrogen that I can—for example, a dime-sized amount of estrogen to apply to her vulvar area twice a week. This amount of estrogen can’t be detected in her system with current assays. I know that some of it will be systemically absorbed, but it cannot be detected. When the patient buys that commercially prepared cream from the pharmacist, she will receive the same black box warning that comes with all systemic hormones since the WHI. However, if she goes to a compounding pharmacist with a prescription for bioidentical hormone therapy, she will not get the warning, regardless of the ingredients or dosage.

Are compounded bioidenticals ever justified?

OBG MANAGEMENT: According to the FDA, compounding of drug products is justified...
Bioidentical hormones

only when a practitioner finds that an FDA-approved drug does not meet the patient’s needs. Do you think this is ever really the case, given the availability of FDA-approved bioidentical hormone preparations?

**DR. PINKERTON:** In rare cases, compounding of bioidentical hormones is justified, such as when a patient cannot tolerate an FDA-approved product. The problem is that women have been especially concerned about the safety of hormone therapy since the WHI, and bioidentical hormones have been promoted as being safer than FDA-regulated preparations, despite the lack of evidence of their safety or efficacy in peer-reviewed literature. So many women request them.

In a recent commentary, Boothby and Doering call bioidentical hormone therapy “a panacea that lacks supportive evidence.” They say, “It’s our belief that pharmacists are compounding these with the best intentions, but they are ill informed regarding the lack of scientific underpinning associated with efficacy and safety.”

**OBG MANAGEMENT:** Do you ever prescribe bioidentical hormones?

**DR. PINKERTON:** Yes, but rarely, and primarily for women who can’t tolerate FDA-approved hormones or who, after adequate information and education, refuse FDA-approved hormone therapy.

Is salivary hormone testing informative?

**OBG MANAGEMENT:** Many clinicians who prescribe bioidentical hormones base the dosage on salivary hormone testing. They claim that this allows them to offer individualized formulations. Is this a reliable claim?

**DR. PINKERTON:** No, it isn’t. Although compounded bioidentical hormone therapy is often prescribed on the basis of salivary hormone testing, there is no scientific evidence

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that a correlation exists between a patient’s symptoms and salivary hormones, or that salivary hormone testing reflects what is happening at the tissue level. As Fugh-Berman and Bythrow have observed, this type of testing is often used to convince asymptomatic consumers to use hormones—or symptomatic women to take higher dosages. That practice is likely to lead to adverse events. The practice also directly contradicts evidence-based guidelines, which recommend that hormone therapy be individualized on the basis of symptoms, not hormone levels.

There are no published studies in the peer-reviewed literature that show that salivary testing is a reliable measure on which to safely and effectively base dosing decisions. Indeed, The Endocrine Society issued a position statement that notes, among other issues, that salivary hormone tests are “inaccurate and should not be considered reliable measures of hormones in the body.” The American College of Obstetricians and Gynecologists also advises against salivary testing, observing that:

1) there is no biologically meaningful relationship between salivary sex steroidal hormone concentrations and free serum hormone concentrations
2) there is large within-patient variability in salivary hormone concentrations. Salivary hormone levels vary depending on diet, time of day of testing, the specific hormone being tested, and other variables.

Do bioidenticals protect against cancer?

OBG MANAGEMENT: Some reports mention the fact that many women believe that bioidentical hormones—specifically, estriol—can reduce their risk of breast and endometrial cancer. Is there any truth behind this belief?

DR. PINKERTON: Estriol is a weak estrogen. There is no evidence that, if it is given at a dosage high enough to relieve symptoms, it is any safer than estradiol.

In regard to endometrial cancer, if the exogenous estrogen—bioidentical or otherwise—is unopposed or inadequately opposed, the risk of endometrial cancer is elevated. The problem is that it is hard to determine whether estrogen is being adequately opposed, particularly when transdermal compounded progesterone is given, because the progesterone molecule is too large to be well-absorbed systemically.

In regard to breast cancer, estriol is a less potent estrogen than estradiol, but it is believed to carry the same risks if it is dosed at effective levels. There is nothing about estriol per se in the peer-reviewed literature that shows that it protects against breast cancer.

The data on risk of breast cancer with estrogen therapy is confusing, with potentially higher risks if estrogen is combined with progestogen. Most of the data we have on estradiol come from animals, but a study from 1980 in humans showed that, when older women with breast cancer were treated with estradiol, 25% had increased growth of metastases.

How do you monitor use of bioidentical hormones?

OBG MANAGEMENT: When you do prescribe a compounded bioidentical hormone, how do you monitor the patient?

DR. PINKERTON: First, I want to reiterate that I prescribe these hormones after considerable patient education about FDA-approved options and their potential risks. Second, when a patient needs or requests hormone therapy, I recommend conventional therapy. Only when she cannot tolerate or refuses FDA-approved drugs do I consider prescribing compounded bioidentical hormones—which, as I said earlier, are assumed to carry risks identical to those of FDA-approved hormones.

In some cases, I provide gynecologic care for patients who obtain compounded bioidentical hormones from other sources. What I will sometimes do, just to give myself some idea of how much estrogen they are getting, is to measure the peak and trough estradiol and estrone levels. That is, I measure the hormone level within 4 hours of the patient taking the drug to see how high it goes, and again about 12 hours later to see how low it
goes. I measure both because estradiol may be peripherally converted to estrone.

Regrettably, we don’t know what to do about the various hormone levels. It isn’t like treating thyroid disorders; we normally dose estrogen therapy based on symptoms.

Who pays?

**OBG MANAGEMENT:** Who pays for salivary testing and compounded bioidentical hormones? Does health insurance cover them?

**DR. PINKERTON:** Like other “natural” products, compounded bioidenticals may cost more than their commercially prepared counterparts and are not covered by insurance. In addition, prescribers may charge more for a “consultation” than do practitioners who accept insurance; they also may recommend salivary testing, which is expensive. Patients can end up paying large sums out of pocket.

As Rosenthal noted, many women do not appear to be concerned about the added costs. That may be because compounded bioidentical hormone therapy is usually offered to economically advantaged patients.

Ethical considerations

**OBG MANAGEMENT:** That raises an important question: What ethical considerations are inherent in the prescribing of compounded bioidenticals?

References


**Instant Poll Results**

What did your colleagues say about compounded bioidentical hormones? See Instant Poll Results on page 15.