Adopted by the OBCE at their March 21, 2013 meeting

**BAX 3000 and Similar Devices**

The BAX 3000 is marketed as device to diagnose and treat allergies and food sensitivities. The device was reviewed by the OBCE’s ETSDP Committee and on January 17, 2013, this policy was adopted by the OBCE in accordance with the ETSDP rule.

The BAX 3000 and similar devices are disapproved (outside the scope) as a diagnostic procedure.

As a treatment modality, the BAX 3000 and similar devices are considered Investigational with Moderate Risk for use with chiropractic patients. This rating requires a written Informed Consent statement signed by the patient. This rating also recommends the chiropractic physician participate in or conduct a formal investigation of the procedure.

The written informed consent must at a minimum address or include:
- The risks of ingesting food or substances which may provoke an anaphylaxis reaction.
- A statement that the use of this treatment could cause an exacerbation.
- An acknowledgement that there is currently a lack of peer reviewed evidence and other evidence such as case studies.
- If the patient is to be part of a research or case study, consents to that participation.
- An understanding that this treatment is considered ‘Investigational with Moderate Risk” by the Oregon Board of Chiropractic Examiners.
- This device/procedure is not used to diagnose allergies or other conditions and that other procedures are used for that purpose.

Chiropractic physicians using the BAX 3000 or similar devices must adhere to the OBCE’s advertising rules and policies. They must refrain from making advertising claims which cannot be supported. (3/21/13)