**K041553: Arthrex Suture Graft Kit**

The purpose of this Memo is to document the addition of the SpeedGraft, suture construct, as a line extension to the Arthrex FiberWire family cleared via 510(k) in K041553 (Arthrex Suture Grafting Kit).

The SpeedGraft is comprised of donated human tendon(s) which are pre-sutured with FiberWire Suture.

The SpeedGraft is intended for use in soft tissue approximation or ligation.

The product labeling and intended use, and the technology, engineering and performance of the SpeedGraft are the same as the Arthrex Suture Grafting Kit. The mechanical testing data demonstrated that the tensile strength of the proposed devices meet or exceed the minimum acceptance criteria.

The proposed changes to Arthrex Suture Grafting Kit are considered minor and do not significantly affect the safety or efficacy of the device. Additionally, the proposed changes were not in response to adverse events in the field. Arthrex, Inc. has determined that the SpeedGraft does not require a 510(k) submission at this time.

Sincerely,

*Courtney Smith*, RAC
Manager, Regulatory Affairs