



Clinical Registry Summary

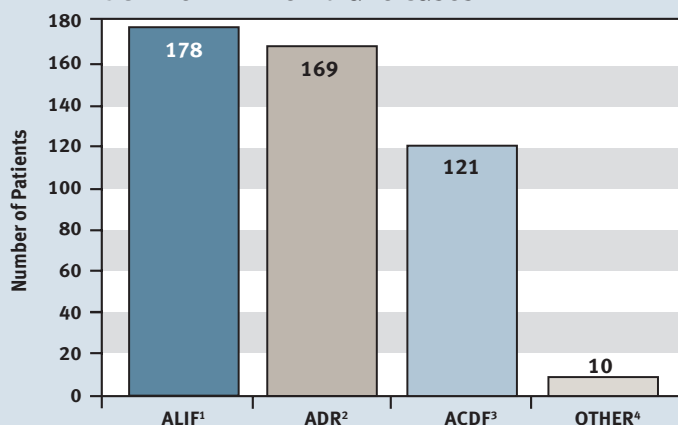
W. L. Gore & Associates conducted a Clinical Registry Program involving several centers in the US and one European center to support the safety of the GORE ePTFE membranes in anterior spinal surgery.

GORE ePTFE membranes were placed between vessels and various types of anterior spinal hardware to provide a permanent plane of dissection around vasculature to facilitate revisions.



GORE PRECLUDE®
Vessel Guard

GORE ePTFE Membrane Cases



- 478 documented implants
- ZERO reported device related complications

¹Anterior Lumbar Interbody Fusion
²Artificial Disc Replacement
³Anterior Cervical Discectomy and Fusion
⁴XLIF, Nucleus Replacement, Hybrid ADR/ALIF

Surgeon comments related to revisions reported at the time of data collection support the benefits of implanting GORE PRECLUDE® Vessel Guard during the index procedure.

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