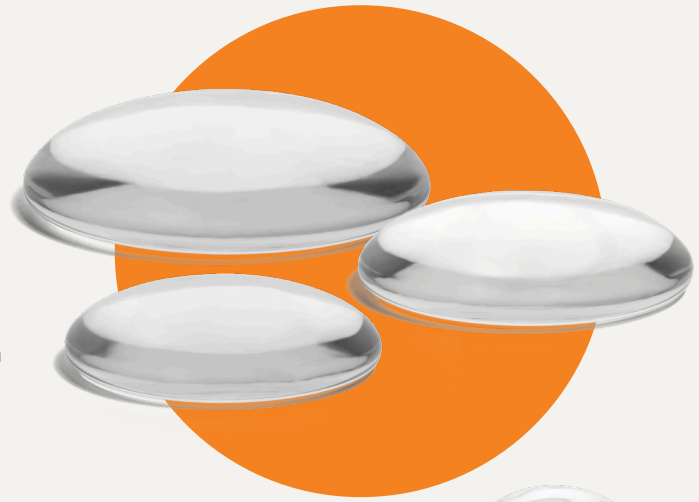


introducing THE LOW PLUS PROFILE PROJECTION

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- 36 new round, smooth implant options (80cc – 440cc)
- Available in both HSC and HSC+ gel for optimal shape retention and fracture resistance while maintaining a natural feel
- Exclusively available to board-certified plastic surgeons



WTH (cm)	PROJ (cm)	VOL (cc)	HSC Catalog #	HSC+ Catalog #
8	2.1	80	10610-080LPP	10710-080LPP
9	2.4	110	10610-110LPP	10710-110LPP
9.75	2.6	140	10610-140LPP	10710-140LPP
10	2.7	155	10610-155LPP	10710-155LPP
10.5	2.8	175	10610-175LPP	10710-175LPP
10.75	2.9	185	10610-185LPP	10710-185LPP
11	3.0	195	10610-195LPP	10710-195LPP
11.25	3.0	210	10610-210LPP	10710-210LPP
11.5	3.1	225	10610-225LPP	10710-225LPP
11.75	3.2	235	10610-235LPP	10710-235LPP
12	3.3	255	10610-255LPP	10710-255LPP
12.5	3.4	280	10610-280LPP	10710-280LPP
12.75	3.5	300	10610-300LPP	10710-300LPP
13	3.6	320	10610-320LPP	10710-320LPP
13.25	3.6	335	10610-335LPP	10710-335LPP
13.5	3.7	355	10610-355LPP	10710-355LPP
14	3.8	390	10610-390LPP	10710-390LPP
14.5	3.9	440	10610-440LPP	10710-440LPP

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BREAST IMPLANTS

Sientra breast implants are indicated for breast augmentation in women at least 22 years old and for breast reconstruction. Breast implant surgery is contraindicated in women with active infection anywhere in their body, with existing cancer or precancerous conditions who have not received adequate treatment for those conditions, and who are currently pregnant or nursing. Prior to use, plastic surgeons should review all risk information, which is found in the Directions for Use. Key complications associated with the use of silicone gel breast implants include capsular contracture, implant removal, rupture, and reoperation. The Directions for Use and detailed information regarding the risks and benefits of Sientra breast implants can be found at sientra.com. The sale and distribution of this device is restricted to users and/or user facilities that provide information to patients about the risks and benefits of this device in the form and manner specified in the approved labeling provided by Sientra, Inc.