<u>COVER SHEET</u>

Instructions for completing Allergan's Device Tracking Form for NATRELLE® Silicone and Saline Breast Implants

IMPORTANT: Read and remove these instructions prior to completing the Device Tracking Form. For Device Tracking purposes the physician/health care facility MUST notify Allergan upon implantation, explantation or when a NATRELLE® Silicone or Saline Breast Implant is discarded or destroyed. Failure to comply could result in violation of Federal law.

Healthcare facility, please complete the following sections of the form. Device tracking information can also be provided electronically via the **AbbVie Device Management Portal**, the National Breast Implant Registry, or the Aesthetic One app. **To use the AbbVie Device Management Portal**, **go to devicemanagement.abbvie.com to register and start data entry.** For the National Breast Implant Registry, go to https://www.surgery.org/downloads/microsite/aestheticone/index.php to register and start data entry.

I. Complete upon Implantation

a. Device and Surgery Information

For implantation surgery, affix the breast implant label attached to the inner product box labeling to page 1 of the forms and place the device tracking label to page 2 of the forms, L for the left breast implant and R for the right breast implant. If labels are not available, please record the catalog number (REF) and serial number (SN) in the space provided for each page of the form.

- b. Implanting/Explanting Physician Information
- c. Attending /Following Physician Information
- d. Patient Information

II. Complete only for new devices opened and discarded/destroyed

Complete this section if *NATRELLE*® Breast Implant(s) were opened and not used for any reason during surgery and discarded or destroyed. If not applicable, mark N/A. If applicable, provide serial number(s), date of occurrence and reason/comments in the space provided. Allergan requires the device to be returned if defective, otherwise if available, you may choose to return it to: 1800 Waters Ridge Drive Suit 100, Lewisville, TX 75057, Attn: Allergan Device Analysis Laboratory with contact information, serial number(s) and reason for return.

III. Complete ONLY if NATRELLE® Breast Implant(s) were removed

After completing the Device Tracking Form and ensuring that the serial number information is on each form, remove page 1 and send by mail or provide electronically through the AbbVie Device Management Portal, NBIR, or Aesthetic One app (detailed above). Provide page 2 to the patient, for completion of enrollment in the Allergan Device Tracking program.

Upon receipt of the first page of the form by Allergan, patient specific information is entered in the Device Tracking database. Patients who do NOT wish to participate in the Device Tracking Program or choose NOT to release their information to any third parties unless legally required, check the appropriate box and unless legally required to retain their personal information, Allergan will remove their personal information from the database upon receipt of their form.

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SHEET 1

Allergan _{th} He science of rejuvenation th					Silicone Vie Device	and Salir Tracking 1 N.	ne Breas Waukegan	TRACKIN st Implant Rd. Bldg. J23 nicago, IL 600 1.800.972.93
I. Complete Upon Implant								
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REF 10-270	ILFE	R)			(Left) S	SN		
3N 20-1 3070	ALLERG	AN			☐ Rec	onstruction	☐ Aug	mentation
	ALLERG	AN			☐ Rev	ision		
Affix RIGHT breast implan	t label here. If lab	el is n	ot avail	able, recor	d REF and	d Serial Nu	mber belo	w.
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II. Complete Only For New De	vices Onened	and	Disca		vallable		1 N/Δ	
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III. Complete ONLY If NATREL	LLE® Breast Im	plan	ts Wei	re Remov	ed		N/A	
•	Explanted	Devi	ce Inf	ormation				
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Left) REF #	Unk	nown	(Right)	REF #				☐ Unknow
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f no, please provide the cause:								
Original implant date: mm/dd			_					
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