



PRECISE-HA Filler PATIENT PAGE

IMPLANT CARD

IMPLANT CARD: WHAT IS IT FOR?

Your healthcare professional has given you your personalized implant card because you have been treated with a **PRECISE-HA dermal filler**.










You must keep it and it is recommended to scan it to ensure preservation of information.

Your implant card is both-sided. The front side is intended to be completed with your personal information while the back side includes information about your device and its traceability.

IMPLANT CARD: CONTENT

IMPLANT CARD FRONT SIDE:

Make sure your healthcare professional has filled in the following fields (1), (2), (3) on the front side:

SKINCEUTICALS		
 (1) Your name	 EN - Person identification FR - Identification de la personne	 EN - Health care center or doctor FR - Centre de santé ou médecin
 (2) Date of injection		
 (3) The name and full address of the healthcare professional where the injection took place	 EN - Date of implantation FR - Date d'implantation	 EN - Subject information website FR - Site web d'informations à destination du sujet
 https://www.symatase.com/patient/precisehafillers/ (4)		
 SYMATESE SAS - ZI les Troques - 69630 Chaponost - France (5)		

(4) Link to the website where all necessary information on your device is available











(5) Name and postal address of the manufacturer of your device



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PRECISE-HA DERMAL FILLERS

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IMPLANT CARD BACK SIDE:

<p> EN - Manufacturer FR - Fabricant</p> <p> EN - Device name FR - Nom du dispositif</p>	<p> EN - Unique device identifier FR - Identifiant unique des dispositifs</p> <p> EN - Batch code FR - Code du lot</p> <p><small>ICS00A</small></p>	<p>(1) EN - Dermal filler - Non medical purpose FR - Produit de comblement dermique - Usage non médical</p> <p>(2)  PRECISE-HA Filler SCULPT</p> <p>(3)  401008</p> <p>(4)  (01) 03760172160489</p> <p>(5)  S2232230223</p> <p>(6)  </p> <p><small>441100</small></p>
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- (1) Device type
- (2) Device name
- (3) Catalogue number
- (4) Unique Device Identifier – Device identifier
- (5) Batch code
- (6) Unique Device Identifier



IMPORTANT INFORMATION CONCERNING YOUR PRECISE-HA Filler DEVICES

PRECISE-HA fillers are intended for non-medical use only.

PRECISE-HA fillers devices are intended to modify the skin anatomy and facial appearance.

PRECISE-HA Filler EYE is indicated to correct infraorbital hollows. PRECISE-HA Filler EYE is injected into dermis to hypodermis and supraperiosteal.
PRECISE-HA Filler SOFTEN is indicated to correct nasolabial folds and perioral lines. PRECISE-HA Filler SOFTEN is injected into dermis to hypodermis.
PRECISE-HA Filler LIPS is indicated to correct the volume and the shape of the lips. PRECISE-HA Filler LIPS is injected into dermis, hypodermis and labial mucosa.
PRECISE-HA Filler SMOOTH is indicated to correct nasolabial folds. PRECISE-HA Filler SMOOTH is injected into dermis to hypodermis.
PRECISE-HA Filler SCULPT is indicated to restore cheek volume loss. PRECISE-HA Filler SCULPT is injected into hypodermis and supraperiosteal.

PRECISE-HA fillers shall only be administered by appropriately trained healthcare professionals who are qualified or accredited in accordance with national law in injection techniques for the modification of facial anatomy.

PRECISE-HA fillers are used in adult subjects excluding pregnant and breastfeeding women.

PRECISE-HA fillers are not to be used in persons who are less than 18 years old.

due to limited clinical data, caution should be exercised when envisaging to use PRECISE-HA Filler EYE on: - Male - Population aged from 18 to 29 years old and older than 69.
Due to limited clinical data, caution should be exercised when envisaging to used PRECISE-HA Filler SOFTEN on: - Male - Population aged from 18 to 34 years old and older than 75.
Due to limited clinical data, caution should be exercised when envisaging to used PRECISE-HA Filler LIPS on: - Male - Population aged from 18 to 34 years old and older than 75.
due to limited clinical data, caution should be exercised when envisaging to use PRECISE-HA Filler SMOOTH on: - Male - Population aged from 18 to 37 years old and older than 70.



due to limited clinical data, caution should be exercised when envisaging to use PRECISE-HA Filler SCULPT on:

- Male
- Population aged from 18 to 35 years old and older than 65.

EXPECTED PERFORMANCE AND LIFETIME

PRECISE-HA fillers act by adding volume to the tissue, thereby restoring the skin contours and the volume and shape of the face to the desired level. The volume and lifting capacity originate from the ability of the crosslinked hyaluronic acid gel to hold its shape in the tissue, thus maintaining over time the volume and projection obtained at injection. The duration of effect depends on the area treated and the depth of injection, and may vary from one subject to another.

PRECISE-HA Filler EYE	<p>On average, with 0,5 mL injected per infraorbital hollow in one session, between 68.4 and 71.9% of subjects are noticeably improved at 9 months (clinical study results obtained with the Global Aesthetic Improvement Scale).</p> <p>The expected lifetime for PRECISE-HA Filler EYE is 12 months.</p>
PRECISE-HA Filler SOFTEN	<p>On average, with 1,2 mL injected into the nasolabial folds and 0.5 ml injected into lip fine lines, in one session, 93% of subjects are noticeably improved at 9 months (clinical study results obtained with the Global Aesthetic Improvement Scale).</p> <p>Note: During the clinical study, the treatment of perioral fine lines with PRECISE-HA Filler SOFTEN was performed in conjunction with the treatment of the lips. The proximity of these two areas may potentially have had a slight influence on the doses used and the performance results obtained. It is recommended to take into account the proximity of neighbouring treated areas to avoid overcorrections.</p> <p>The expected lifetime for PRECISE-HA Filler SOFTEN is 12 months.</p>
PRECISE-HA Filler LIPS	<p>On average, with 1 mL injected into the lips in one session, 93% of subjects are noticeably improved at 9 months (clinical study results obtained with the Global Aesthetic Improvement Scale).</p> <p>The expected lifetime for PRECISE-HA Filler LIPS is 12 months.</p>



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PRECISE-HA Filler SMOOTH	On average, with 0.9 mL injected per nasolabial fold in one session, 82% of subjects are noticeably improved at 9 months (clinical study results obtained with the Global Aesthetic Improvement Scale). The expected lifetime for PRECISE-HA Filler SMOOTH is 9 months.
PRECISE-HA Filler SCULPT	On average, with 1,1 mL injected per cheek in one session, between 58.2 and 60% of subjects are noticeably improved at 9 months (clinical study results obtained with the Global Aesthetic Improvement Scale). The expected lifetime for PRECISE-HA Filler SCULPT is 15 months.

PRECISE-HA Filler devices contain local anaesthetic lidocaine in order to reduce pain during and post-injection procedure.

CONTRAINDICATIONS

PRECISE-HA Filler devices are contraindicated in:

- Minors.
- Subjects with a known allergy to hyaluronic acid, lidocaine or amide local anesthetics.
- Subjects with porphyria.
- Subjects with an autoimmune disorder, or using an immunosuppressant medication.
- Pregnant or breastfeeding women.
- Subjects with inflammation, infection or cutaneous disorders at the treatment site or nearby.
- Subjects with bleeding disorders or in subjects receiving thrombolytic or anticoagulant treatment.
- Areas other than those recommended.

RESIDUAL RISKS AND POTENTIAL UNDESIRABLE SIDE-EFFECTS

According to SYMATESE policy the residual risks have been identified, reduced as far as possible. The main method of mitigation is the design control of the device, validation of usability of the device to the foreseeable use errors, manufacturing controls and then the information provided to the user (labelling, instruction for use,...). These residual risks are all considered as acceptable under normal conditions of use, which is consistent with a high level of protection for the safety and health of persons. This is in accordance with the state of science and medical knowledge.

The adverse events related to PRECISE-HA Filler devices are minor, reversible and similar to other hyaluronic filler available on the market.



The most common side-effects are injection site reactions (swelling, tenderness/pain, firmness, bruising, lumps/bumps, redness, itching, discoloration). Most of these reactions disappear spontaneously after a few hours or days without any treatment. When these symptoms persist, they can be treated with analgesics, topical steroids or non-steroidal anti-inflammatory drugs (NSAIDs). Lumps and bumps can also be treated with hyaluronidase. In case of doubt or complication, patient should contact the healthcare professional to make a monitoring. Appropriate treatment is at the discretion of the healthcare professional and is in accordance with the laws of the territories.

The following adverse events have been reported with regard to hyaluronic acid fillers: abnormal hepatic function, acne, bleeding, bruising /hematoma /ecchymosis /contusion, capillary disorders /telangiectasia, discoloration, discomfort or pain /tenderness, dryness /xerosis, erosion /exfoliation, erythema /redness, eyes disorders /vision disorder /blindness, facial asymmetry, granuloma, headache, hyperbilirubinaemia, induration /firmness, infection /folliculitis, inflammation, ischemia /necrosis, mass /lumps /bumps /nodules /irregularities, migration of the device, muscle tightness /twitching, needle track marks, nerve injury, numbness /paraesthesia /anaesthesia /hypoesthesia, pruritus /itching, rosacea, scarring /scar hyperpigmentation and hypertrophy, sensation of foreign body /burning sensation /hyperaesthesia, seroma, severe allergic reaction /hypersensitivity /angioedema /rash, superficial wound, swelling /oedema /lymphadenopathy, syncope /dizziness, Tyndall effect, vascular damage /vascular compromise, vesicles, warmth.

Vascular compromise may occur due to an inadvertent intravascular injection or as a result of vascular compression associated with implantation of any injectable product. This may manifest as ischemia or necrosis at the implant site or in the area supplied by the blood vessels affected, or very rarely as ischemic events in other organs due to embolization. Following facial esthetic treatments, isolated rare cases have been reported regarding ischemic events affecting the eyes (leading to loss of vision) and the brain (resulting in cerebral infarction). After injection closed to the nose, ischemia or necrosis may occur, especially in subjects who have previously undergone rhinoplasty.

Symptoms of inflammation at the implant site commencing either shortly after injection or after a delay of up to several weeks have been reported on hyaluronic acid fillers. In case of unexplained inflammatory reactions, infections should be considered and treated if necessary, since inadequately treated infections may progress into complications.

For subjects who have experienced clinically significant reactions, any decision to repeat treatment must take into consideration the cause and severity of previous reactions.

Rare cases of granuloma have been reported in the literature and may occur after a few months or even several years.



Adverse Events related to Lidocaine: Overdosage toxic reactions / Drug interactions (antiarrhythmics, amide type anaesthetics), Hypersensitivity / Allergy, Porphyria.

Precautions related to Lidocaine: Systemic or local adverse events may occur for patients with epilepsy, impaired cardiac conduction, severely impaired hepatic function or severe renal dysfunction.

DECLARATION

The sterile gel of PRECISE-HA Filler devices does not contain CMR (carcinogenic, mutagenic or toxic for reproduction) and/or endocrine-disrupting substances.

WARNING AND PRECAUTIONS

No device other than hyaluronic acid must be implanted in the area to be treated.

If laser treatment, chemical peeling or any other procedure based on active dermal response is considered after treatment with PRECISE-HA Filler devices, there is a theoretical risk of eliciting an inflammatory reaction at the implant site. This also applies if PRECISE-HA Filler devices are administered before the skin has healed completely after such a procedure.

In the event of local inflammation, you must avoid extreme temperature conditions (intense cold, sauna, etc.) until any signs have disappeared.

Considering the composition of PRECISE-HA Filler devices, there is no warning, precaution and/or measure to be taken as regards the exposure to reasonably foreseeable external influences or environmental conditions, such as magnetic fields, external electrical and electromagnetic effects, electrostatic discharge, radiation associated with diagnostic or therapeutic procedures

QUALITATIVE AND QUANTITATIVE INFORMATION ON THE MATERIAL USED FOR THE MANUFACTURING OF PRECISE-HA Filler DEVICES

Here are the details on the composition of your PRECISE-HA Filler devices.

PRECISE-HA Filler devices contain a sterile gel in an injection device. The injectable gel is based on hyaluronic acid, crosslinked with 1,4-butanediol diglycidyl ether (BDDE), containing viscous non-crosslinked hyaluronic acid and lidocaine, packed in a 1 mL plastic syringe.

The components of **PRECISE-HA Fillers** are:



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	Reference	PRECISE-HA Filler EYE	PRECISE-HA Filler SOFTEN	PRECISE-HA Filler LIPS	PRECISE-HA Filler SMOOTH	PRECISE-HA Filler SCULPT
Main ingredients	Total Sodium Hyaluronate	15 mg/g	20 mg/g	20 mg/g	22 mg/g	23 mg/g
	Lidocaine hydrochloride	3 mg/g	3 mg/g	3 mg/g	3 mg/g	3 mg/g
Other ingredients	Sodium chloride NaCl	9 mg/g	9 mg/g	9 mg/g	9 mg/g	9 mg/g
	Sodium dihydrogenophosphate dihydrate NaH ₂ PO ₄ · 2H ₂ O	0.02 mg/g	0.02 mg/g	0.02 mg/g	0.02 mg/g	0.02 mg/g
	Disodium phosphate dihydrate Na ₂ HPO ₄ · 2H ₂ O	0.22 mg/g	0.22 mg/g	0.22 mg/g	0.22 mg/g	0.22 mg/g
	Water for Injection (WFI)	QSP 1g	QSP 1g	QSP 1g	QSP 1g	QSP 1g
Volume per syringe		1 mL	1 mL	1 mL	1 mL	1 mL

MAXIMUM DOSAGE

PRECISE-HA Filler EYE	<p>Based on the clinical data, per session, it is recommended to not inject more than 1.0mL of PRECISE-HA Filler EYE per side.</p> <p>As PRECISE-HA Filler EYE gel contains lidocaine, the practitioner must consider the total dose of lidocaine and other anesthetics of the amide type administered if the subject is using a device or medicinal treatment that delivers lidocaine, as the systemic toxic effects can be additive. High doses of lidocaine (more than 200 mg/day) can cause acute toxic reactions manifesting as symptoms affecting the central nervous system and cardiac conduction.</p>
PRECISE-HA Filler SOFTEN	<p>Based on the clinical data, per session, it is recommended to not inject more than 1.2 mL of PRECISE-HA Filler SOFTEN per nasolabial fold and to not inject more than 1.4 mL of PRECISE-HA Filler SOFTEN for perioral lines during one session</p>



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	<p>As PRECISE-HA Filler SOFTEN contains lidocaine, the practitioner must consider the total dose of lidocaine and other anesthetics of the amide type administered if the subject is using a device or medicinal treatment that delivers lidocaine, as the systemic toxic effects can be additive. High doses of lidocaine (more than 200 mg/day) can cause acute toxic reactions manifesting as symptoms affecting the central nervous system and cardiac conduction.</p>
PRECISE-HA Filler LIPS	<p>Based on the clinical data, per session, it is recommended to not inject more than 1.0 mL of PRECISE-HA Filler LIPS into the upper lip and to not inject more than 1.3 mL of PRECISE-HA Filler LIPS into the lower lip.</p> <p>As PRECISE-HA Filler LIPS gel contains lidocaine, the practitioner must consider the total dose of lidocaine and other anesthetics of the amide type administered if the subject is using a device or medicinal treatment that delivers lidocaine, as the systemic toxic effects can be additive. High doses of lidocaine (more than 200 mg/day) can cause acute toxic reactions manifesting as symptoms affecting the central nervous system and cardiac conduction.</p>
PRECISE-HA Filler SMOOTH	<p>Based on the clinical data, per session, it is recommended to not inject more than 2.0 mL of PRECISE-HA Filler SMOOTH per nasolabial fold.</p> <p>As PRECISE-HA Filler SMOOTH gel contains lidocaine, the practitioner must consider the total dose of lidocaine and other anesthetics of the amide type administered if the subject is using a device or medicinal treatment that delivers lidocaine, as the systemic toxic effects can be additive. High doses of lidocaine (more than 200 mg/day) can cause acute toxic reactions manifesting as symptoms affecting the central nervous system and cardiac conduction.</p>
PRECISE-HA Filler SCULPT	<p>Based on the clinical data, per session, it is recommended to not inject more than 3.0 mL of PRECISE-HA Filler SCULPT per side.</p> <p>As PRECISE-HA Filler SCULPT gel contains lidocaine, the practitioner must consider the total dose of lidocaine and other anesthetics of the amide type administered if</p>



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the subject is using a device or medicinal treatment that delivers lidocaine, as the systemic toxic effects can be additive. High doses of lidocaine (more than 200 mg/day) can cause acute toxic reactions manifesting as symptoms affecting the central nervous system and cardiac conduction.

- ✓ **Information about medicinal substance in the device:** PRECISE-HA Filler devices contain lidocaine hydrochloride (3 mg/mL). It is added in order to reduce pain during injection and improve comfort during and after the injection procedure.
- ✓ The sterile gel of PRECISE-HA Filler devices **does not contain CMR (carcinogenic, mutagenic or toxic for reproduction), endocrine-disrupting substances, phthalate in the device.**
- ✓ PRECISE-HA Filler devices **do not contain and are not manufactured with tissue of neither animal origin nor any animal tissue derivative.**
- ✓ PRECISE-HA Filler devices **do not include any derivatives of tissues or cells of human origin which are non-viable or are rendered non-viable.**

INFORMATION ON WHEN AND HOW TO REPORT UNDESIRABLE SIDE-EFFECTS

Please be aware that any serious incident that might have been caused by PRECISE-HA fillers should be reported directly or through the distributor to the manufacturer or the Competent Health Authority of the country where the event occurred.

INFORMATION ON WHEN TO CONTACT HEALTHCARE PROFESSIONAL

Contact your healthcare professional if you believe that you are experiencing side-effects related to the device or if you are concerned about risks.

LINKED DOCUMENTATION

SSCP- Section B (Patient)

SSCP PRECISE-HA Filler EYE	FASY MDR 24-006
SSCP PRECISE-HA Filler SOFTEN	FASY MDR 24-003
SSCP PRECISE-HA Filler LIPS	FASY MDR 24-004
SSCP PRECISE-HA Filler SMOOTH	FASY MDR 24-005
SSCP PRECISE-HA Filler SCULPT	FASY MDR 24-007