



Natrelle PERKS[™] is an offer exclusively for Alle Members who receive a breast augmentation with Natrelle® gel breast implants. Natrelle® breast augmentation patients are eligible to redeem a complimentary treatment with either JUVÉDERM® Ultra XC or BOTOX® Cosmetic (onabotulinumtoxinA). Joining Allē is easy. Plus, as an Allē Member, you will receive 500 points on your breast augmentation procedure with Natrelle® that can be redeemed for future Allergan Aesthetics treatments.

KASEY Actual Natrelle® patient. Individual results may vary.

Natrelle® Breast Implants IMPORTANT SAFETY INFORMATION AND APPROVED USES

Breast implants are not considered lifetime devices. The longer people have them, the greater the chances are that they will develop complications, some of which will require more surgery.

Breast implants have been associated with the development of a cancer of the immune system called breast implant-associated anaplastic large cell lymphoma (BIA-ALCL). This cancer occurs more commonly in patients with textured breast implants than smooth implants, although rates are not well defined. Some patients have died from BIA-ALCL.

Patients receiving breast implants have reported a variety of systemic symptoms, such as joint pain, muscle aches, confusion, chronic fatigue, autoimmune diseases, and others. Individual patient risk for developing these symptoms has not been well established. Some patients report complete resolution of symptoms when the implants are removed without replacement.

Who can get breast implants?

Natrelle® Breast Implants are approved for the following:

• Breast augmentation for women at least 22 years old for silicone-filled implants and for women at least 18 years old for saline-filled implants. Breast augmentation includes primary breast augmentation to increase the breast size and revision surgery to correct or improve the result of a primary breast augmentation

• Breast reconstruction. This includes primary breast reconstruction to replace breast tissue that has been removed due to cancer or trauma or that has failed to develop properly due to a severe breast abnormality. This also includes revision surgery to correct or improve the result of a primary breast reconstruction

Who should NOT get breast implants?

Breast implant surgery should NOT be performed in:

• Women with active infection anywhere in their body

- Women with existing cancer or precancer of their breast who have not received adequate treatment for those conditions
- Women who are currently pregnant or nursing

What should I tell my doctor?

Tell your doctor if you have any of the following conditions, as the risks of breast implant surgery may be higher:

- Autoimmune diseases (eg, lupus and scleroderma)
- A weakened immune system (eg, taking medications to decrease the body's immune response)
 Planned chemotherapy or radiation therapy following breast implant placement
 Conditions or medications that interfere with wound healing and blood clotting

- Reduced blood supply to breast tissue
- Clinical diagnosis of depression or other mental health disorders, including body dysmorphic disorder and eating disorders
- Those with a diagnosis of depression or other mental health disorders should wait for resolution or stabilization of these conditions prior to undergoing breast implantation surgery

Please see additional Important Safety Information throughout for Natrelle® and BOTOX® Cosmetic, including Boxed Warnings, and JUVÉDERM® Collection of Fillers.



implants removed and not replaced, you may experience dimpling, puckering, wrinkling, or other cosmetic changes, which may be permanent

• Breast implantation is likely not a one-time surgery. The longer implants are in place, the greater the potential risk for complications. You will likely need additional surgeries on your breasts due to complications or unacceptable cosmetic results. Thus, you should also consider the complication rates for later (revision) surgery since you may experience these risks in the future

• Cancer treatments and surgery will affect the outcome and timing of breast reconstruction

• Breast implants may affect your ability to breastfeed, either by reducing or eliminating milk production

• Rupture of a silicone-filled breast implant is most often silent. Even if you have no symptoms, you should have your first ultrasound or MRI at 5 to 6 years after your initial implant surgery and then every 2 to 3 years thereafter regardless of whether your implants are for augmentation or reconstruction. If you have symptoms of or uncertain ultrasound results for breast implant rupture, an MRI is recommended. Additional imaging may be required depending on your medical history and status. The health consequences of a ruptured silicone gel-filled breast implant have not been fully established

Here's How IT WORKS

- 1. Join Allē. Allē is the loyalty program by Allergan Aesthetics. Download the app to ioin. You must have the Alle app to claim your complimentary Natrelle Perks™ offer.
- 2. Checkout. Make sure your plastic surgeon has submitted your breast augmentation in Allē. You will earn 500 points to be used on a future Allergan Aesthetics treatment. You can check vour email for a treatment confirmation from Allē or go to "History" in your Allē Wallet to see if your treatment has been added.
- 3. Select. After you have joined Alle and your augmentation is submitted, text PERKS to 65190 to opt in for Alle text messages.* Two text messages will be delivered. Follow the second link to select your complimentary treatment. Before selecting, work with your provider on which complimentary treatment is right for you. Once you select your treatment, the offer will be deposited into your Wallet in 48 hours, and cannot be changed. Remember, you must have the Alle app to claim the offer.
- 4. Redeem. Once your offer has been deposited into your Alle Wallet, book your treatment with an Alle provider to redeem within 6 months of receiving.

*Standard text and data rates may apply. You can text "STOP" to opt out of Alle text at any time.

• Routine screening mammography for breast cancer will be more difficult, and implants may rupture during the procedure. Perform self-examination every month for cancer screening and ask your surgeon to help you distinguish the implant from your breast tissue. Lumps, persistent pain, swelling, hardening, or changes in implant shape should be reported to your surgeon and possibly evaluated

What are key complications with breast implants?

Key complications include reoperation, implant removal with or without replacement, implant rupture with silicone-filled implants, implant deflation with saline-filled implants, and capsular contracture (severe scar tissue around the implant). Other complications include breast pain, swelling, asymmetry, wrinkling/ rippling, implant malposition nipple complications, hypertrophic scarring, and implant palpability/visibility. Talk to your doctor about other complications.

For more information, see the patient brochures at www.allergan.com/products.

To report a problem with Natrelle® Breast Implants, please call Allergan® at 1-800-624-4261.

The sale and distribution of Natrelle® Breast Implants is restricted to licensed physicians who provide information to patients about the risks and benefits of breast implant surgery.

Please see additional Important Safety Information throughout for Natrelle® and BOTOX® Cosmetic, including Boxed Warnings, and JUVÉDERM® Collection of Fillers.



Enjoy the Full REWARDS OF ALLĒ

Allē is a premier consumer loyalty program from Allergan Aesthetics, where Members can earn and redeem points on Allergan Aesthetics brands, including BOTOX® Cosmetic (onabotulinumtoxinA), the JUVÉDERM® Collection of Fillers, SkinMedica®, and more. Members can also earn points on a variety of other brands and treatments. Visit Alle.com for more information.

Joining Alle is easy

To get started, visit Alle.com or download the app.

Once registered, you'll gain access to:

- Exclusive rewards
- Curated content
- Treatment information

Download the Alle app now!









JUVÉDERM® Injectable Gel Fillers Important Information APPROVED USES

JUVÉDERM® VOLUMA® XC injectable gel is for deep injection in the cheek area to correct age-related volume loss and for augmentation of the chin region to improve the chin profile in adults over 21.

JUVÉDERM® VOLLURE® XC, JUVÉDERM® Ultra Plus XC, and JUVÉDERM® Ultra XC injectable gels are for injection into the facial tissue for the correction of moderate to severe facial wrinkles and folds, such as nasolabial folds. JUVÉDERM® VOLLURE® XC injectable gel is for adults over 21.

JUVÉDERM® VOLBELLA® XC injectable gel is for injection into the lips for lip augmentation and for correction of perioral lines in adults over 21.

JUVÉDERM® Ultra XC injectable gel is for injection into the lips and perioral area for lip augmentation in adults over 21.

IMPORTANT SAFETY INFORMATION

Are there any reasons why I should not receive any JUVÉDERM® formulation?

Do not use these products if you have a history of multiple severe allergies or severe allergic reactions (anaphylaxis), or if you are allergic to lidocaine or the Gram-positive bacterial proteins used in these products.

What precautions should my doctor advise me about?

- Minimize strenuous exercise and exposure to extensive sun or heat within the first 24 hours following treatment. Exposure to any of these may cause temporary redness, swelling, and/or itching at the injection site
- Tell your doctor if you are pregnant or breastfeeding. The safety of these products for use during pregnancy or while breastfeeding has not been studied
- The safety of JUYÉDERM® VOLUMA® XC has not been studied in patients under 35 years or over 65 years for cheek augmentation, or under 22 years and over 80 years for chin augmentation. The safety of JUYÉDERM® VOLLURE® XC and JUVÉDERM® VOLBELLA® XC has not been studied in patients under 22 years, and the safety of JUYÉDERM® Ultra Plus XC and JUYÉDERM® Ultra XC has not been studied in patients under 18 years
- JUVÉDERM® VOLUMA® XC is intended for use in the chin and cheek areas. JUVÉDERM®
 VOLLURE® XC, JUVÉDERM® Ultra Plus XC, and JUVÉDERM® Ultra XC are intended for use in facial
 wrinkles and folds. JUVÉDERM® VOLBELLA® XC and JUVÉDERM® Ultra XC are intended for use in
 the lips and perioral area. The safety and effectiveness for treatment in other areas have not been
 established in clinical studies

Please see additional Important Safety Information throughout for Natrelle® and BOTOX® Cosmetic, including Boxed Warnings, and JUVÉDERM® Collection of Fillers.

Which Treatment IS RIGHT FOR YOU?

A consultation with your provider will determine which treatment is right for you!

JUVÉDERM® Ultra XC Before-and-After Photos (Kelly)









Actual patient. Results may vary.

Unretouched photos taken before treatment and 1 month after treatment with 1.2 ml of JUVÉDERM® Ultra XC in the smile lines and 1.7 ml of JUVÉDERM® Ultra XC in the lips.

JUVÉDERM® Injectable Gel Fillers IMPORTANT SAFETY INFORMATION (continued) What precautions should my doctor advise me about? (continued)

- Tell your doctor if you have a history of excessive scarring (thick, hard scars) or pigmentation disorders.
 The safety of JUVÉDERM® products has not been studied in these patients and may result in additional scars or changes in pigmentation
- Tell your doctor if you are on therapy used to decrease the body's immune response (immunosuppressive therapy). Use may result in an increased risk of infection
- Tell your doctor before treatment if you are using substances that can prolong bleeding, such as aspirin, ibuprofen, or other blood thinners. As with any injection, this may result in increased bruising or bleeding at the injection site
- Patients who experience skin injury near the site of injection may be at a higher risk for adverse events
- JUVÉDERM® VOLUMA® XC was not studied in patients with significant loose skin of the chin, neck, or jaw
- The effect of JUVÉDERM® VOLUMA® XC injection into the chin on facial hair growth has not been studied What are possible side effects?

The most commonly reported side effects with JUVÉDERM® injectable gels included redness, swelling, pain, tenderness, firmness, lumps/bumps, bruising, discoloration, and itching. For JUVÉDERM®

JUVÉDERM® is the #1 chosen dermal filler collection* that offers five different products to meet specific needs in different areas of the face. JUVÉDERM® Ultra XC is an injectable gel that temporarily adds more fullness in and plumps thin lips, and is also approved for smoothing and correcting moderate to severe facial wrinkles and folds, such as the smile lines around the nose and mouth.

*Based on provider survey data Sept 2021 (n=967)

JUVÉDERM® Ultra XC Before-and-After Photos (Clancy)









Actual patient. Results may vary.
Unretouched photos taken before treatment and 1 month after treatment with 1.2 ml of JUVÉDERM® Ultra XC in the lips.

VOLBELLA® XC, dryness was also reported. For JUVÉDERM® VOLUMA® XC, most side effects resolved within 2 to 4 weeks. For JUVÉDERM® VOLLURE® XC, JUVÉDERM® Ultra Plus XC, and JUVÉDERM® Ultra XC injectable gels, most resolved within 14 days or less. For JUVÉDERM® VOLBELLA® XC, most resolved within 30 days or less. These side effects are consistent with other facial injection procedures.

Most side effects will resolve with time. Your doctor may choose to treat side effects persisting over 30 days with antibiotics, steroids, or hyaluronidase (an enzyme that breaks down hyaluronic acid).

One of the risks with these products is unintentional injection into a blood vessel. The chances of this happening are very small, but if it does happen, the complications can be serious and may be permanent. These complications, which have been reported for facial injections, can include vision abnormalities, blindness, stroke, temporary scabs, or permanent scarring of the skin.

As with all skin injection procedures, there is a risk of infection.

Visit Juvederm.com or talk to your doctor for more information. To report a side effect with any JUVÉDERM® product, please call Allergan at 1-800-433-8871.

Products in the JUVÉDERM® Collection are available only by a licensed physician or properly licensed practitioner.

Please see additional Important Safety Information throughout for Natrelle® and BOTOX® Cosmetic, including Boxed Warnings, and JUYÉDERM® Collection of Fillers.



Only BOTOX® Cosmetic (onabotulinumtoxinA) is FDA-approved to temporarily make moderate to severe frown lines, crow's feet, and forehead lines look better in adults. BOTOX® Cosmetic delivers predictable results. It's the #1 selling product* of its kind in the US.

*Data collected through September 2021 (N = 970)

Treatment 1/Valerie Moderate to severe forehead lines





Real patient. Results may vary.

Photos taken at maximum eyebrow elevation before and 30 days after treatment with BOTOX* Cosmetic. In 2 clinical studies of healthy adults, 61% and 46% had a \geq 2-grade improvement in their forehead lines at day 30.

Treatment 2/Valerie Moderate to severe forehead lines





Real patient. Results may vary.

Photos taken at maximum eyebrow elevation before and 30 days after treatment with BOTOX* Cosmetic. In 2 clinical studies of healthy adults, 61% and 46% had a \geq 2-grade improvement in their forehead lines at day 30.

Side effects associated with the injection include localized pain, infection, inflammation, tenderness, swelling, redness, and/or bleeding/bruising.

BOTOX® Cosmetic (onabotulinumtoxinA) Important Information Approved Uses

BOTOX® Cosmetic is a prescription medicine that is injected into muscles and used to temporarily improve the look of moderate to severe forehead lines, crow's feet lines, and frown lines between the eyebrows in adults.

IMPORTANT SAFETY INFORMATION

BOTOX® Cosmetic may cause serious side effects that can be life threatening. Get medical help right away if you have any of these problems any time (hours to weeks) after injection of BOTOX® Cosmetic:

 Problems swallowing, speaking, or breathing, due to weakening of associated muscles, can be severe and result in loss of life. You are at the highest risk if these problems are pre-existing before injection. Swallowing problems may last for several months.

Treatment 1/Valerie Moderate to severe crow's feet lines





Real patient. Results may vary.

Photos taken at full smile before and 30 days after treatment with BOTOX* Cosmetic. In 2 clinical studies, 26.1% and 20.3% of adults had a \geq 2-grade improvement at day 30. In one of these studies, 67.9% had mild or no crow's feet lines at day 30 after treatment.

Treatment 2/Valerie Moderate to severe crow's feet lines





Real patient. Results may vary.

Photos taken at full smile before and 30 days after treatment with BOTOX* Cosmetic. In 2 clinical studies, 26.1% and 20.3% of adults had a \geq 2-grade improvement at day 30. In one of these studies, 67.9% had mild or no crow's feet lines at day 30 after treatment.

Spread of toxin effects. The effect of botulinum toxin may affect areas away from the injection
site and cause serious symptoms including: loss of strength and all-over muscle weakness, double
vision, blurred vision and drooping eyelids, hoarseness or change or loss of voice, trouble saying words
clearly, loss of bladder control, trouble breathing, and trouble swallowing.

${\tt BOTOX}^{\otimes}$ Cosmetic dosing units are not the same as, or comparable to, any other botulinum toxin product.

There has not been a confirmed serious case of spread of toxin effect when BOTOX® Cosmetic has been used at the recommended dose to treat frown lines, crow's feet lines, and/or forehead lines.

BOTOX® Cosmetic may cause loss of strength or general muscle weakness, vision problems, or dizziness within hours to weeks of taking BOTOX® Cosmetic. If this happens, do not drive a car, operate machinery, or do other dangerous activities.

Please see additional Important Safety Information throughout for *Natrelle*® and BOTOX® Cosmetic, including Boxed Warnings, and JUVÉDERM® Collection of Fillers.



FAQs

How do I redeem my Alle points?

Allē points can be used on Allergan Aesthetics products and treatments. Points are only redeemable at Alle provider offices. A consultation may be required to determine if a treatment is right for you. When you've decided on a treatment with your provider, ask to apply your available points at checkout for instant savings. Injection fees may apply.

How do I know how many Alle points I have?

You can check your points balance in your Alle Wallet online or through the Alle app.

How do I get my free JUVÉDERM® **Ultra XC or BOTOX® Cosmetic** (onabotulinumtoxinA) treatment?

Once you have had your breast augmentation procedure, your surgeon must submit your procedure in Allē. After it has been submitted, you must text "PERKS" to 65190 to claim your complimentary Natrelle Perks[™] treatment offer. Work with your provider to determine which complimentary treatment is right for you. Upon claiming, it will take 48 hours for your offer to be deposited into your Alle Wallet.

How long is my Natrelle PERKS™ complimentary offer valid for?

After claiming your offer, you have 6 months to redeem it in Allē.

What number do I call if I need help?

You can call the Alle support team at 1-888-912-1572, Monday - Friday, 8 AM - 6 PM CT.

BOTOX® Cosmetic (onabotulinumtoxinA) IMPORTANT SAFETY INFORMATION (continued)

Serious and/or immediate allergic reactions have been reported. They include: itching, rash, red itchy welts, wheezing, asthma symptoms, or dizziness or feeling faint. Get medical help right away if you are wheezing or have asthma symptoms, or if you become dizzy or faint.

Do not receive BOTOX® Cosmetic if you: are allergic to any of the ingredients in BOTOX® Cosmetic (see Medication Guide for ingredients); had an allergic reaction to any other botulinum toxin product such as Myobloc® (rimabotulinumtoxinB), Dysport® (abobotulinumtoxinA), or Xeomin® (incobotulinumtoxinA); have a skin infection at the planned injection site.

Tell your doctor about all your muscle or nerve conditions, such as ALS or Lou Gehrig's disease, myasthenia gravis, or Lambert-Eaton syndrome, as you may be at increased risk of serious side effects including difficulty swallowing and difficulty breathing from typical doses of BOTOX® Cosmetic.

Tell your doctor about all your medical conditions, including: plans to have surgery; had surgery on your face; have trouble raising your eyebrows; drooping eyelids; any other abnormal facial change; are pregnant or plan to become pregnant (it is not known if BOTOX® Cosmetic can harm your unborn baby); are breast-feeding or plan to (it is not known if BOTOX® Cosmetic passes into breast milk).



Natrelle PERKSSM Terms and Conditions

- Allē Members who undergo breast augmentation with Natrelle® gel implants ("Member") may qualify to receive either (a) one (1) complimentary treatment of BOTOX® Cosmetic (onabotulinumtoxinA) up to 50 units OR (b) one (1) complimentary treatment of JUVÉDERM® Ultra XC up to two .55 mL syringes at participating Allē provider offices only. Allergan Aesthetics is not responsible for any associated injection costs.
- Members enrolled in Medicare, Medicaid, or other federal or state healthcare programs are not eligible for this offer.
- Member must claim offer in the Allē app via text message link within 6 months of their Natrelle® breast augmentation. • Once claimed, offer must be redeemed within 6 months of the offer being deposited into Member's Allē Wallet. Offer
- expires 6 months after issue date into Allē Wallet.
- Limit 1 per Member.
- A healthcare provider will determine if Member is an appropriate candidate for a BOTOX® Cosmetic or JUVÉDERM®
- If Member is an appropriate candidate, offer can be redeemed at a participating provider's office.
- . Members will earn Allē points on all qualifying Earnings Eligible Product purchases, subject to applicable earnings caps.
- . The value of this offer cannot be redeemed or exchanged for cash.
- · Offer cannot be applied to past transactions.
- Offer cannot be combined with other Alle offers on BOTOX® Cosmetic or the JUVÉDERM® Collection of Fillers but can be combined with Allē and Allē brand-specific gift cards, Allē points offers, including Double Points offers, and other Allē brand-specific offers
- The complete value of this offer must be used in a single transaction.
- If you have questions, please contact Allē Customer Support at 1-888-912-1572 Monday Friday, 8 AM 6 PM CT.
- · Allergan Aesthetics, an AbbVie company, reserves the right to alter or cancel this offer at any time.



To learn more about Natrelle PERKS[™], ask your provider or visit Natrelle.com/rewards to get started.



To learn more about Allē, scan here.

BOTOX® Cosmetic (onabotulinumtoxinA) IMPORTANT SAFETY INFORMATION (continued)

Tell your doctor about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements, Using BOTOX® Cosmetic with certain other medicines may cause serious side effects. Do not start any new medicines until you have told your doctor that you have received BOTOX® Cosmetic in the past.

Tell your doctor if you have received any other botulinum toxin product in the last 4 months; have received injections of botulinum toxin such as $Myobloc^{\circ}$, $Dysport^{\circ}$, or $Xeomin^{\circ}$ in the past (tell your doctor exactly which product you received); have recently received an antibiotic by injection; take muscle relaxants; take an allergy or cold medicine; take a sleep medicine; take aspirin-like products or blood thinners.

Other side effects of BOTOX® Cosmetic include: dry mouth: discomfort or pain at the injection site: tiredness; headache; neck pain; and eve problems; double vision, blurred vision, decreased evesight. drooping eyelids and eyebrows, swelling of your eyelids and dry eyes.

For more information refer to the Medication Guide or talk with your doctor.

To report a side effect, please call Allergan at 1-800-678-1605.

Please see accompanying Summary of Information about BOTOX® Cosmetic.

Please see additional Important Safety Information throughout for Natrelle® and BOTOX® Cosmetic, including Boxed Warnings, and JUVÉDERM® Collection of Fillers. 15





To learn more about Natrelle PERKS[™], ask your provider or visit Natrelle.com/rewards to get started.









JUVÉDERM°



© 2022 AbbVie. NATRELLE® and its design are trademarks of Allergan, Inc., an AbbVie company. BOTOX® and its design are registered trademarks of Allergan, Inc., an AbbVie company. JUVÉDERM® and its design are trademarks of Allergan Holdings France SAS, an AbbVie company, or its affiliates. All rights reserved. Allē® and its design are service marks of Allergan, Inc., an AbbVie company. Natrelle.com PRT152761 04/22 21-84