Restylane-L®
Patient brochure

Restylane-L®
Injectable Gel with 0.3% Lidocaine

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Q What is Restylane-L?
A Restylane-L® is a crystal clear injectable gel composed of hyaluronic acid, a natural substance that already exists in the body. Restylane-L is non-animal based and free from animal protein. Allergy pretesting is not necessary. Restylane-L contains 0.3% lidocaine. The lidocaine in Restylane-L has been added to reduce the discomfort associated with the treatment.

Q How does Restylane-L work?
A Restylane-L is injected into the skin with an ultrafine needle to plump the skin to smooth away wrinkles and folds such as the lines from your nose to the corners of your mouth (nasolabial folds) or into your lips for patients over the age of 21 for lip enhancement.

Q Why add Lidocaine to Restylane?
A Lidocaine was added to Restylane-L to reduce the pain and discomfort during and after injection. In a clinical study, 60 patients received Restylane® on one side of the face and Restylane-L® on the other side of the face. Restylane-L had an effect on reducing pain. At the time of injection, patients rated their pain about 45 on a scale of 0 to 100 for the side of the face treated with Restylane. In comparison, patients rated their pain about 15 on the same scale for the side of the face treated with Restylane-L. Patients reported less pain on the side of the face treated with Restylane-L up to 60 minutes after treatment.

Q How long does Restylane-L last?
A Restylane-L effects generally last about six months and gradually disappears from the body.

Q Has Restylane-L been studied?
A A clinical study was conducted with Restylane-L to evaluate the pain reducing effect up to 60 minutes after injection. This study enrolled 60 patients with moderate to severe nasolabial fold wrinkles. The study included 58 female patients and 2 male; 34 were White, 21 were Hispanic or Latino, 3 were African American, 1 was Asian, and 1 was “Other”. In this study 71.7% of patients experienced less pain after injection of Restylane-L than with Restylane alone. Please see the below table for additional information.

<table>
<thead>
<tr>
<th>Timepoint</th>
<th>Number of patients with pain reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No.</td>
</tr>
<tr>
<td>After Injection</td>
<td>43/60</td>
</tr>
<tr>
<td>15 Minutes</td>
<td>28/60</td>
</tr>
<tr>
<td>30 Minutes</td>
<td>17/60</td>
</tr>
<tr>
<td>45 Minutes</td>
<td>10/60</td>
</tr>
<tr>
<td>60 Minutes</td>
<td>4/60</td>
</tr>
</tbody>
</table>
In addition to evaluating the pain reducing effects, the study assessed patient satisfaction with Restylane-L treatment. All 60 subjects were asked to rate the level of improvement seen in their nasolabial folds after injection with Restylane-L. At day fourteen after injection 100% saw some improvement (Improved, Much Improved, and Very Much Improved). See below table for additional details.

<table>
<thead>
<tr>
<th>Category</th>
<th>Restylane-L</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Number of Patients</td>
</tr>
<tr>
<td>Very Much Improved (4)</td>
<td>17</td>
</tr>
<tr>
<td>Much Improved (3)</td>
<td>29</td>
</tr>
<tr>
<td>Improved (2)</td>
<td>14</td>
</tr>
<tr>
<td>No Change (1)</td>
<td>-</td>
</tr>
<tr>
<td>Worse (0)</td>
<td>-</td>
</tr>
</tbody>
</table>

**Safety**

**Q** **Who should not use Restylane-L (Contraindications)?**

**A** Safety has not been established and should not be used in people who are:
- Pregnant
- Breast feeding
- Trying to become pregnant
- Under the age of 22 for lip enhancement
- Under the age of 18 or over the age of 65
- Highly allergic (for example: gram positive bacteria)
- Prone to bleeding disorders

**Q** **What are some warnings to consider?**

**A** The use of Restylane-L at sites with skin sores, pimples, rashes, hives, cysts, or infections should be postponed until healing is complete. Use of Restylane-L in these instances could delay healing or make your skin problems worse.

You may experience skin discoloration (bruising), swelling, redness, tenderness, pain, itching, or small lumps in the area where you are injected. If any of these events occur, the majority usually last one to two weeks. If any symptom lasts longer than two weeks, call the doctor who administered the Restylane-L injection.

Red or swollen small bumps (inflammatory papules) may rarely occur. You may need antibiotics to treat them. In clinical studies swelling was higher in younger patients (28%) compared to older patients (18%) and bruising was higher in older patients (28%) compared to younger patients (14%). The majority of these events were mild.

If you are injected with Restylane-L into your lips, your physician should be able to feel the product when touching your lips.
Q  What are some potential risks you may encounter?
A  As with all procedures like this, the injection of Restylane-L carries a risk of infection and formation of scar tissue.

The safety and effectiveness of Restylane-L has not been established in pregnant or nursing mothers, and in patients under 18 or over 65 years of age. Restylane-L use while nursing could harm you or the nursing child. Restylane-L should not be used for lip enhancement in patients under the age of 22.

The use of Restylane-L in African-American patients can result in darkening of skin color (hyperpigmentation), which may take several weeks to correct.

If you have previously had facial cold sores, an injection can cause them to come back.

The safety of Restylane-L used with other skin therapies such as laser, mechanical or chemical peeling, and hair removal has not been established. The use of Restylane-L with these skin therapies may lead to other side effects such as inflammation.

You should avoid exposing the area(s) treated with Restylane-L to excessive sun or UV lamps, and extreme heat and cold until any redness or swelling has disappeared.

Clinical volunteers keeping diaries reported the following short-lived events: Restylane-L was evaluated in a clinical study of 60 patients. The below table shows what patients reported each day after injection of Restylane-L in the diary they kept. The most common events were: pain, swelling, redness, tenderness, bruising, itching and other. The reporting of these events decreased over time and by day 14 most events had resolved.

<table>
<thead>
<tr>
<th>Percentage of Patients Reporting Adverse Events After Treatment with Restylane-L</th>
<th>Number of days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (%)</td>
<td>1</td>
</tr>
<tr>
<td>Bruising</td>
<td>58.3%</td>
</tr>
<tr>
<td>Redness</td>
<td>50.0%</td>
</tr>
<tr>
<td>Swelling</td>
<td>66.7%</td>
</tr>
<tr>
<td>Pain</td>
<td>45.0%</td>
</tr>
<tr>
<td>Tenderness</td>
<td>68.3%</td>
</tr>
<tr>
<td>Itching</td>
<td>13.3%</td>
</tr>
<tr>
<td>Other</td>
<td>6.7%</td>
</tr>
</tbody>
</table>

Q  What are some benefits from clinical evaluation?
A  In one study in which 135 patients received Restylane injections in their lips, two weeks after the injection 96% of the patients said their lips were improved compared to before the injection. At least 74% of the patients still saw an improvement in their lips at 6 months after the injection.
Post-Marketing Surveillance:

**Q** Have there been adverse events reported through post-market surveillance?

**A** The adverse events received from post-marketing surveillance (voluntary reporting and published literature) for Restylane with and without lidocaine in the U.S. and other countries included: swelling and inflammatory reactions – immediate onset and onset up to several weeks after treatment, mass formation including lumps or bumps, induration (hardening), lack of effect, erythema (skin redness), hematoma (bruising), pain or tenderness, papules or nodules (small bumps), hyperpigmentation (skin discoloration), presumptive bacterial infections and abscess formation, inflammation, ischemia and necrosis (restricted blood flow leading to the death of skin) due to unintentional injection into a blood vessel, hypersensitivity, angioedema (swelling just below the skin), injection site reactions including burning sensation, warmth, irritation, extrusion of device, neurological symptoms including hypoesthesia (reduced sense of touch), paraesthesia (tingling sensation), facial nerve paralysis, pruritus (itching), atrophy/scarring, visual disturbance including blurred vision lasting for a short time, reduced vision, increased flow of tears, eyelid drooping, blindness, symptoms of reactivation of herpes infection, granuloma (mass formation), device dislocation, telangiectasia (spider veins/broken capillaries), rash, blisters/vesicles, acne, fistula, effusion/discharge, dermatitis, urticaria (hives), muscle twitching, dermatophytosis (fungal infection), encapsulation, fainting (vasovagal reactions) and other dermatological events including dry skin, skin exfoliation, skin wrinkling, alopecia (baldness), chapped lips, and non-dermatological events including headache, fever, dizziness, sinusitis, dyspnoea (shortness of breath) and anxiety.

Warning: One of the risks with using this product 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. If you have changes in your vision, signs of a stroke (including sudden difficulty speaking, numbness or weakness in your face, arms, or legs, difficulty walking, face drooping, severe headache, dizziness, or confusion), white appearance of the skin, or unusual pain during or shortly after treatment, you should notify your health care practitioner immediately.

About the Procedure

**Q** What are the serious side effects?

**A** Rarely, the doctor may accidentally inject the product into a blood vessel, which can cause injury to the blood supply and damage to the skin or lips.

Rarely, a few people have developed infections that must be treated with antibiotics or other treatment. Infection may be hard to treat, but will generally go away when the gel is absorbed.
Q What should patients do prior to treatment?
A *Restylane-L* requires no pretesting, but you should take a few precautions before being treated. Avoid using St. John’s Wort, high doses of Vitamin E supplements, aspirin, and other non-steroidal anti-inflammatory medications, such as ibuprofen prior to treatment, because these may increase bruising or bleeding at the injection site. Please speak to your doctor about when to stop these medications before your procedure. Also, if you have previously suffered from facial cold sores, discuss this with your physician. He or she may consider prescribing a medication to minimize recurrences.

Q What is the dose of *Restylane-L*?
A The amount used depends on your face and what you would like to have treated. The average patient who has all of the severe wrinkles around the mouth or lips corrected will use less than half a tablespoon.

Please speak to your doctor to determine the correct amount of product needed.

Q Do the injections hurt?
A *Restylane-L* is injected directly into the skin with an ultrafine needle. To help maximize your comfort, you should discuss the use of numbing medicines with your doctor before treatment.

Q How much does *Restylane-L* treatment cost?
A *Restylane-L* is a customized procedure based on your specific needs, so the cost will vary from patient to patient. In general, the cost of *Restylane-L* is similar to the cost of similar procedures. Please ask your doctor to give you an estimate of the cost.

Q Are there post-treatment instructions to follow after a *Restylane-L* treatment?
A Please observe the following after treatment with *Restylane-L*:

- A cloth dipped in cold water (cold compresses), wrung out, and applied to the injected area may be used immediately after treatment to reduce swelling.

- Avoid touching the treated area within six hours following treatment so you do not accidentally injure your skin while the area is numb. After that, the area can be gently washed with soap and water.

- Until there is no redness or swelling, avoid exposure of the treated area to intense heat such as sun lamps or sun bathing.

- If you have previously suffered from facial cold sores, there is a risk that the needle punctures could contribute to another occurrence. Speak to your physician about medicine to prevent this from happening again.

- Avoid taking aspirin, non-steroidal anti-inflammatory medications, St. John’s Wort, and high doses of Vitamin E supplements for one week after treatment. These agents may increase bruising and bleeding at the injection site.
Troubleshooting

Q  When should I call my doctor?
A  Most side effects like bruising, swelling, pain, tenderness, redness, and itching will usually go away within one to two weeks. Call your doctor if you have persistent problems beyond 14 days.

   Blisters or skin sores that recur may signal the presence of a herpes infection that must be treated.

   You can develop an infection that should be treated with antibiotics. If you experience any signs of infection such as fever, redness that spreads to surrounding areas, drainage, increasing tenderness, or increasing pain that does not go away you should call your doctor.

   Seek immediate medical attention if you develop symptoms such as unusual pain, vision changes, a white appearance of skin near the injection site, or any signs of a stroke (including sudden difficulty speaking, numbness or weakness in your face, arms, or legs, difficulty walking, face drooping, severe headache, dizziness, or confusion) during or shortly after the procedure (http://www.nlm.nih.gov/medlineplus/stroke.html).

Restylane-L®

User Assistance Information
Your questions about Restylane-L can be personally answered by contacting the Galderma Laboratories, L.P. toll-free call center between 8:00 a.m. and 5:00 p.m. Central Daylight Time, Monday through Friday.

1-855-425-8722

Company logo

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