NOVELTIES IN DERMATOLOGY

Laser-Assisted Drug Delivery

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Abstract Absorption of topical products through the epidermis is limited by the skin’s barrier function. Numerous techniques and agents such as microneedling, dermabrasion, radiofrequency, and lasers have been used to increase penetration within an approach known as transdermal drug delivery. One of these techniques is laser-assisted drug delivery (LADD), which often uses ablative fractional lasers (CO2 or erbium:YAG lasers) because of their capacity to produce microscopic ablated channels. The parameters in LADD need to be adjusted to the patient, the skin condition and its location, and the drug. LADD has been used with various topical products, such as corticosteroids, photosensitizers, and immunotherapy agents (imiquimod or 5-fluorouracil) to treat numerous conditions, including scars, nonmelanoma skin cancer, and photodamage. LADD is a promising technique that enhances the absorption of topical molecules while adding the synergic effect of the laser.

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KEYWORDS Laser-assisted drug delivery; Ablative fractional laser; CO2 laser; Hypertrophic scars; Keloid scars

PALABRAS CLAVE Vehiculización de fármacos asistida por láser; Láser fraccional ablativo; Láser CO2; Cicatrices hipertróficas; Queloides

Resumen La absorción de productos tópicos a través de la epidermis está limitada por la función de barrera cutánea. Existen distintas técnicas, tales como el microneedling, la dermabrasión, la radiofrecuencia o los láseres, que se han empleado para aumentar la absorción de estas sustancias en una estrategia conocida como vehiculización transdérmica de fármacos. Entre estas técnicas destaca la vehiculización de fármacos asistida por láseres (VFAL), especialmente láseres fraccionales ablativos (CO2/Er:YAG), por su capacidad de generar canales de microablación. En la VFAL se deben ajustar los parámetros en función del tipo de paciente, la dermatosis, la localización y el fármaco empleado. Se ha estudiado la VFAL en el uso de...
Introduction

One of the most typical parts of the dermatologist’s work is that of prescribing topical treatments to induce a local effect against a skin disease. However, absorption of these agents is limited by the skin’s main function as a barrier to the external environment. Given its composition and location as the outermost layer, the stratum corneum is the key player in this defense function.1

It is estimated that around 1%-5% of topical preparations cross an intact epidermis.2 The epidermis can be crossed via 3 pathways: (1) the intercellular pathway, through which the most lipophilic substances are absorbed via the desmosomes in the spaces between cells; 2) the transcellular pathway, across the aqueous pores of the keratinocytes; and 3) the transappendageal pathway, which is mainly across hair follicles and, to a lesser extent, acrosyringia.3

We must also take account of the intrinsic properties of the drug that directly affect absorption. According to the Fick laws of diffusion, the flux of absorption is determined not only by differences in the concentration of drug absorbed between the external environment and the dermis, but also by the solubility and diffusion coefficient of the drug. The heaviest molecules, ie, those with the highest molecular weight, diffuse more slowly than smaller or neutral molecules. Lipophilic and hydrophobic substances make their way across the dermis more quickly.4

Topical drugs can be more easily absorbed by means of physical methods in what is known as transdermal delivery. One of the main methods involves lasers, which have led to exponential growth in the use of the technique known as laser-assisted drug delivery (LADD).1

Transdermal Drug Delivery

Transdermal drug delivery involves assisted transport of substances across the epidermis using various physical methods. This promising method of drug delivery to the body makes it possible to increase the absorption achieved with respect to topical delivery. In addition, oral delivery often entails problems of real bioavailability owing to the first-pass effect and may be limited by the greater risk of adverse effects. Studies on transdermal drug delivery have investigated scraping, dermabrasion, microneedling, pressure waves, vacuum effect, and lasers. These modalities have been used to deliver various types of drugs and substances, such as triamcinolone, methyl aminolevulinic acid (MAL), 5-fluorouracil, and methotrexate.5-11

Few studies have compared the efficacy of the various techniques available. One well-designed study compared the efficacy of topical MAL with that of photodynamic therapy (PDT) after application of MAL in addition to various physical methods, such as ablative fractional laser (AFL), nonablative fractional laser (NAFL), curettage, dermabrasion, and microneedling.12 The authors concluded that there were no differences in the formation of protoporphyrin IX after application of MAL alone or in combination with NAFL. However, the remaining techniques did prove superior with respect to formation of protoporphyrin IX. The most effective of all was AFL. These results have been confirmed, at least partially, by subsequent studies; therefore, AFLs are currently considered the most effective drug delivery technique.13,14

Laser-Assisted Drug Delivery: General Considerations

Lasers

Different types of laser have been used for LADD. Each type is more or less effective, although it is also important to know the individual safety profile. In practical terms, we can divide the lasers used in LADD into 4 groups: (1) Fully ablative lasers, such as the carbon dioxide (CO2) laser (10 600 nm) or the erbium-doped yttrium aluminum garnet (Er:YAG) laser (2940 nm), whose main chromophore is water and which lead to heating and total vaporization of the skin; (2) AFLs, which are the same as the previous group, although when used fractionally, they lead to columns of thermal injury, which are microscopic treatment zones (MTZs); (3) NAFL, which are similar to the previous group and include lasers such as the erbium fiber laser (1550 nm). These also produce MTZs, although in the columns, only heating of the skin is observed, with no ablation of the dermal-epidermal junction; and (4) Nonablative dermal remodeling lasers. This group includes all lasers with chromophores other than water but which have been used to induce greater drug absorption. They also include lasers for the treatment of vascular anomalies, such as the 585/595-nm pulsed dye laser or the neodymium-doped YAG laser (1064 nm).15 The characteristics of AFL have led it to be the most widely studied in LADD, followed by NAFL (Fig. 1).

While the risk of adverse effects such as erythema, vesiculation, crusting, or even scarring are lower with fractional lasers than with conventional ablative lasers, these effects may appear when high fluences and densities are applied.16-18 If AFLs are used for LADD, the settings must
be optimized to achieve the best results possible with the minimum risk.

The density of fractional lasers refers to the amount of tissue covered by the MTZs. The total density depends on both the number of channels and on the size of the laser beam. Initial studies with MAL, which were subsequently corroborated with other agents (diclofenac, tretinoin), showed that the concentrations of the product in skin increased with increases in density up to a maximum of 5%, at which point the concentrations stabilized. Thus, densities higher than 5% in treatment with AFL carry no clear benefits but do carry a greater risk of adverse events.

Another key parameter is the depth of the channel itself, which depends directly on laser fluence. As a general rule, we should apply higher fluences in order to reach the deeper layers of the dermis if we wish to treat conditions such as alopecia or scarring. The most superficial layers of the dermis would be appropriate for treatment of photodamage, melasma, or superficial scarring. In patients with melanoma, vitiligo, or superficial nonmelanoma skin cancer, ablation restricted to the epidermis could be sufficient.

In the case of AFLs, drug absorption is affected by the coagulation zone (CZ) surrounding the channels, as well as by total ablation depth. The area of the CZ ranges from around 50 mm to 150 mm, depending on fluence and, in particular, laser wavelength, and is greater for CO2 lasers (10 600 nm) than for Er:YAG lasers (2940 nm). The greater affinity of Er:YAG for water leads to “purer” ablation, with almost no heat given off around the MTZs, whereas in the case of CO2, a wider CZ is generated owing to the reduced affinity for water. The presence of CZs implies even greater absorption of the drug delivered across the channels, thus also explaining the superiority of AFL over modalities that do not involve the generation of heat around the channels, as is the case in microneedling. However, it is important to remember that the presence of CZs only reflects transmission of heat to the surrounding dermis, which may be associated with undesirable local reactions.

**Drugs and Application Techniques**

Given that lipophilic substances have a greater intrinsic ability to cross an intact epidermis, the more marked effect of LADD is observed with hydrophilic substances. Other relevant factors that affect absorption include the vehicle used, the type of preparation, and the presence of additives or excipients. Liquid and gel formulations cross the channels produced by AFL with greater affinity than more oily formulations in the form of creams or ointments. This observation should be taken into account owing to the greater efficacy and the greater risk of adverse effects when these formulations are used.

There is also some disparity between hydrophobic and hydrophilic substances and their association with penetration via the laser channels. In the case of hydrophilic substances, such as 5-fluorouracil or methotrexate, greater absorption has been detected in direct relation to AFL fluence and, therefore, the depth of the channels. However, this finding has not been confirmed in studies with hydrophobic substances such as imiquimod and lidocaine.

Various considerations must be taken into account with respect to the suitability of applying topical drugs directly to the dermis. First, we must ask whether the substances and excipients we intend to use are suitable for subcutaneous application. Local hypersensitivity has been reported with the application of vitamin C serum delivered via microneedling, which led to the formation of histologically
confirmed granulomas or even systemic hypersensitivity reactions. Under ideal conditions, we would use only highly sterile products, although not all topical products are available as such. AFL itself carries some risk of bacterial infection owing to direct exposure of the dermis, and this risk increases when we apply drugs such as 5-fluorouracil, corticosteroids, and MAL. It is important to insist on aseptic conditions for application of LADD procedures.

Another key factor in LADD is the interval between use of the laser and application of the product to the skin. It is important to remember that the microscopic channels close with time owing to debris such as fibrin, inflammatory mediators, and keratinocytes. One study showed that topical application 6 hours after use of the laser increases absorption, which is most successful during the first 30 minutes. However, no increase in absorption is observed when the product is applied after 24 hours. Figure 2 shows the appropriate steps for LADD and the considerations to be taken into account for each one.

Patients

As is the case with topical treatments, absorption of drugs using LADD depends on many characteristics that are intrinsic to the patient, the skin disease treated, and the area affected by the disease. Thus, absorption is more potent in cases of inflamed or eroded skin. Similarly, more hydrated skin exhibits greater affinity for the absorption of oily substances. Furthermore, the patient’s age is important, given that atrophy, erosion, and ulceration are more likely to affect older people, who are also at greater risk of infection and take longer to recover. Therefore, the ideal approach would be to make a correct evaluation of the patient’s skin type, since there may be some atrophy due to solar elastosis in patients with low skin types, even younger patients. In such cases, laser fluence should be reduced, and in CO₂ AFL devices that allow it, pulses with a lower heat deposit should be administered to prevent pathological scarring. As for location, the drug is absorbed mainly across the transappendageal pathway through the follicles in hairy areas; therefore, the parameters for using LADD should be applied with greater caution.

Laser-Delivered Drugs

For practical purposes, we focus on 2 of the main applications of LADD: delivery of corticosteroids for hypertrophic and keloid scars and laser-assisted PDT. In addition to these 2 indications, numerous drugs have been delivered in many other dermatologic indications. Data on these studies are shown in Table 1.

Aminolevulinic Acid and Methyl 5-Aminolevulinate: Laser-Assisted Photodynamic Therapy

Some studies report enhanced absorption of aminolevulinic acid and methyl 5-aminolevulinate assisted by various types of laser, and there are even studies that use devices such as pulsed dye lasers as light sources. In the present analysis, we are interested in those studies that use lasers to deliver photosensitizers to the dermis and thus ensure greater effectiveness with PDT. Thus, studies of the usefulness of LADD in the context of PDT—both for treatment of actinic keratosis and for superficial basal cell and squamous cell carcinoma—have reported greater rates of effectiveness, but also more adverse effects (eg, crusting, pruritus, pain, and erythema). Haedersdal et al demonstrated absorption rates for MAL that were 13.8 times greater with continuous-wave Er:YAG and up to 7.3 times greater with the fractional version. Similar results have been reported from animal models based on CO₂ lasers.
### Table 1  Medications Used in Laser-Assisted Drug Delivery.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Main Indications Assessed</th>
<th>Design of Main Studies</th>
<th>Outcome</th>
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</table>
| Aminolevulinic acid and 5-MAL: laser-assisted PDT<sup>48-57</sup> | 1. Actinic keratosis<sup>48</sup>  
2. Fine nodular basal cell carcinoma (< 2 mm)<sup>56</sup>  
3. Superficial squamous cell carcinoma<sup>57</sup> | 1. One session of fractional Er:YAG (550 μm; 22% density) + PDT with conventional 5-MAL vs 1 session of PDT with conventional 5-MAL  
2. One session of fractional Er:YAG (550 μm; 22% density) + PDT with 5-MAL vs 2 sessions with conventional PDT  
3. Two sessions of fractional Er:YAG (550 μm; 22% density) + PDT with 5-MAL vs 2 sessions of conventional PDT | 1. Response rate of 91% in laser-assisted PDT vs 65.6% in conventional PDT at 3 months  
2. Total response of 84% in laser-assisted PDT vs 50% in conventional PDT at 3 months  
3. Total response of 84.3% in laser-assisted PDT vs 52% in conventional PDT at 3 months |
<p>| TCA&lt;sup&gt;35,58,59&lt;/sup&gt; | Hypertrophic and keloid scars | Fractional Er:YAG (first pass 150-200 μm without coagulation; second pass 300-400 μm with coagulation; 22% density) + intralesional TCA 10% vs fractional Er:YAG + topical corticosteroids (desoximetasone 0.25%). Four sessions with intervals of 6 weeks&lt;sup&gt;35&lt;/sup&gt; | Improvement in both cases. No significant differences compared with postlaser intralesional or topical corticosteroids, although tolerance is much better with topical delivery and patient preference is greater |
| Methotrexate&lt;sup&gt;9&lt;/sup&gt; | Studies of absorption in an animal model (murine) | Topical application after continuous-wave Er:YAG 2940 nm vs electroporation | Better outcome for absorption with LADD (3-80 times greater than topical treatment alone depending on fluence) vs electroporation (2 times greater than topical treatment alone) |
| Imiquimod&lt;sup&gt;22&lt;/sup&gt; | Studies on absorption in an animal model (murine) with various types of laser | 0.4% aqueous formulation of imiquimod after fractional Er:YAG 2940 nm | Similar intradermal concentrations with the 0.4% aqueous formulation in LADD and with topical imiquimod 5% cream |
| Ingenol mebutate&lt;sup&gt;43&lt;/sup&gt; | Bowen disease | Ingenol mebutate 0.015% for 3 days on the face and 0.05% for 2 days on the rest of the body; preceded or not by fractional CO&lt;sub&gt;2&lt;/sub&gt; laser on the first day of treatment (100 mJ; 100 spot/cm&lt;sup&gt;2&lt;/sup&gt;) | Complete response in 88.9% of patients treated using LADD vs 12.5% with ingenol mebutate alone |</p>
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</tr>
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<tbody>
<tr>
<td>5-Fluorouracil(^{7,11})</td>
<td>1. Absorption studies in an animal model (murine) with various types of laser</td>
<td>1. Q-switched ruby laser (694 nm) vs continuous-wave Er:YAG (2940 nm) vs continuous-wave CO(_2) laser (0.12 mm spot; 10 mJ per pulse; density 5%) followed by 5-fluorouracil 5% under occlusion for 7 days</td>
<td>1. Better rates of absorption with continuous Er:YAG reaching (\times 156) 2. Histologically confirmed cure of 87% cases of Bowen disease and 71% of cases of basal cell carcinoma</td>
</tr>
<tr>
<td></td>
<td>2. Superficial basal cell carcinoma and Bowen disease on the trunk and limbs &lt; 2 cm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Timolol(^{41})</td>
<td>Deep infantile hemangioma</td>
<td>Timolol 0.5% in ophthalmic solution applied 4-5 times per day with weekly sessions of ablative fractional CO(_2) laser (0.12 mm spot; 25-30 mJ per pulse; density 5%)</td>
<td>Good/excellent response in more than 80% of patients with a mean of 14 weeks of treatment</td>
</tr>
<tr>
<td>Pimecrolimus, tretinoin, and bimatoprost(^{49})</td>
<td>Hypopigmented scars</td>
<td>Topical bimatoprost 0.03% + tretinoin 0.05%+ pimecrolimus 1% twice daily for 3 mo. Combined with nonablative fractional laser (1550 nm; 20-70 mJ; 20%-32% density), mean of 4.5 sessions at intervals of 4-8 wk</td>
<td>Partial repigmentation &gt; 50% in 86% of cases</td>
</tr>
<tr>
<td>Poly-L-lactic acid(^{48})</td>
<td>Atrophic scars</td>
<td>Sculptr (Valeant Aesthetics) reconstituted with saline solution, lidocaine, and epinephrine in combination with fractional CO(_2) laser (120 (\mu)m spot size, 375-500 (\mu)m depth, density 10%)</td>
<td>Improvement of more than 30% in most cases. Lower formation of delayed papules than with intralesional treatment</td>
</tr>
<tr>
<td>Finasteride(^{50})</td>
<td>Male pattern baldness</td>
<td>Combination of finasteride 0.05% and growth factors with nonablative fractional Er:Glass laser (1550 nm, 8 passes, 7 mJ, 3-9% density)</td>
<td>Improvement in follow-up photographs in 4 patients</td>
</tr>
<tr>
<td>Vitamins(^{46})</td>
<td>Recovery after rejuvenation treatment with ablative fractional laser</td>
<td>Formula with vitamin C, vitamin E, and ferulic acid applied with fractional CO(_2) laser daily during reepithelization. Randomized controlled study</td>
<td>Improved recovery with fewer adverse effects. In molecular terms, increase in expression of bFGF compared with placebo group</td>
</tr>
<tr>
<td>Anesthetics(^{28,42,43})</td>
<td>Studies on absorption of and preexposure to subcutaneous injections</td>
<td>Topical lidocaine 4% combined with ablative fractional Er:YAG laser</td>
<td>Increased serum levels of lidocaine and its metabolites in LADD vs topical administration. Lower pain scores in the group treated with lidocaine in LADD before subcutaneous injection</td>
</tr>
</tbody>
</table>

Abbreviations: bFGF, basic fibroblast growth factor; LADD, laser-assisted drug delivery; 5-MAL: methyl 5-aminolevulinate; PDT, photodynamic therapy; TCA, triamcinolone acetonide.
Hypertrophic Scars: Triamcinolone Acetonide

Triamcinolone acetonide (TCA) is one of the most widely used intralesional or subcutaneous corticosteroids in various skin diseases owing to its sustained release. The main adverse effects include atrophy, ecchymosis, telangiectasia, infection, and even ulceration at the injection site. Application of this agent using LADD has mainly been investigated in the treatment of pathologic scarring (hypertrophic and keloid scars) and in the sequelae of burns. While this drug has been investigated in several studies, that of Waibel et al. on hypertrophic scars causing functional restriction is noteworthy. The authors administered TCA using LADD (fractional CO\textsubscript{2} laser), with 3-5 treatments every 2-3 months. Improvement was recorded not only in texture and thickness, but also for dyschromia and scar functionality. Several studies have reported similar results. Of particular interest is a study comparing effectiveness and tolerability in the treatment of hypertrophic scars using fractional Er:YAG followed by intralesional corticosteroid injections (TCA 10%) or topical corticosteroids (desoximetasone 0.25% under occlusion), with 4 sessions every 6 weeks. The results show the topical corticosteroid to be slightly more effective than the intralesional drug, although the difference was not statistically significant. The authors also reported much better scores for pain and patient preference.

Our experience shows that the combination of CO\textsubscript{2} laser and TCA 10% applied with smooth massage during the 15 minutes following application of the laser yields very good results both for hypertrophic and keloid scars and for the sequelae of burns (Fig. 3). In addition, the simple application of topical anesthetic before laser treatment ensures better tolerance than combining laser treatment with subsequent TCA injections. In the case of very vascularized erythematous scars, this technique can be combined with a previous vascular approach based on techniques such as pulsed dye laser or intense pulsed light. For this purpose, we generally use pulsed dye laser (595 nm) with pulses of 0.5-6 ms and fluence of 5-7 J/cm\textsuperscript{2}, followed immediately by AFL.

Application of Laser-Assisted Drug Delivery in Fields Other Than Dermatology

LADD has been used to introduce drugs and substances into the body for many and various reasons, in addition to those we have seen in dermatology. Studies in animals and humans have shown the efficacy of fully ablative and fractional lasers for introducing small interfering RNA molecules (siRNA) and antibodies. siRNA can inhibit translation of various genes, and this has proven useful in many diseases. Although the only studies on LADD with siRNA to
date have been performed in mice, this pathway could be explored in the future in dermatological diseases using siRNA, as is the case with microRNA 21 in psoriasis, for example.

LADD has even proven effective for generating a favorable immune response with topical vaccines. Even more advanced are pilot studies with hematopoietic stem cells applied topically in combination with ablative laser in mice, thus paving the way for possible application of LADD as an alternative in hematopoietic progenitor transplant.

Conclusions

LADD can yield better results with topical treatments by facilitating absorption and achieving a synergic effect between medical treatment (drugs) and physical treatment (lasers). Safety and the most appropriate technique remain controversial, although there is no doubt that these issues will gradually be resolved in the coming years.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

References


