

ENDO ONE: Advanced Skin Rejuvenation for Aesthetic Clinics Through Italian Innovation and Clinical Insight



ENDO ONE

Why choose Endo One?

Endo-One is a state-of-the-art diode fiber system for providers seeking versatility, clinical excellence, and transformative patient outcomes. Featuring dual-wavelength diode fiber technology (980 nm & 1470 nm).

Endo One adapts to multiple clinical and aesthetic applications, delivering powerful results with maximum safety and precision. Two wavelengths, endless possibilities. Treat fat, skin laxity, vascular lesions, intimate wellness, and more with one compact, powerful device

FDA CLEARED

This FDA-cleared device integrates multiple treatment protocols, making it an essential tool for diverse clinical indications.

CORE TECHNOLOGY

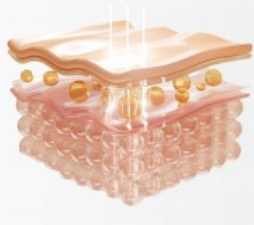
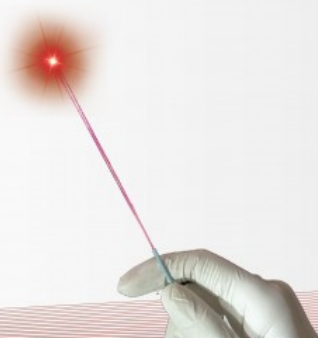
980 nm
Wavelength Superior fat emulsification
Coagulation of vessels
Ideal for lipolysis and contouring

1470 nm
Wavelength Optimal water absorption
Advanced skin tightening
Collagen remodeling with minimal thermal damage

Selectable Single or Combined Mode
Customizable energy delivery for patient-specific protocols and precise tissue targeting

Engineered by Aesthelab® Italian Excellence
Designed in Italy by Aesthelab®, a brand synonymous with reliability, style, and innovation, this laser combines ergonomic elegance with advanced functionality, ensuring both comfort and efficiency in daily use.

New concept of endolaser

Pozzuoli, Campania Sep 22, 2026 ([Issuewire.com](https://www.issuewire.com)) - Aesthetic medical practices operate in an increasingly sophisticated market where patients consistently demand noticeable structural lifting, skin renewal, and tissue tightening without the prolonged downtime, scarring, and risks associated with incisional surgery. [Advanced Skin Rejuvenation for Aesthetic Clinics](#) achieves optimal clinical efficacy when medical equipment design directly reflects procedural realities encountered by physicians during delicate subcutaneous interventions. Medical directors and aesthetic practitioners require treatment platforms that deliver predictable tissue retraction and dermal remodeling while maintaining seamless workflow efficiency throughout demanding clinical schedules.

Developed through a dedicated transatlantic partnership between frontline clinical practitioners and Italian medical device manufacturer ENDO ONE SRL, [ENDO ONE](#) provides modern aesthetic practices with an advanced diode endolaser platform. The system unites responsive optical energy control with ergonomic fiber handling to support high-precision subcutaneous rejuvenation across diverse patient demographics. By delivering targeted laser energy directly into the reticular dermis and subdermal tissue planes, the platform bridges the therapeutic gap between superficial non-invasive skin tightening and invasive plastic surgery.

Bridging Frontline Clinical Insight with Italian Engineering Excellence

Energy-based medical aesthetic devices achieve durable clinical adoption only when their engineering architecture solves genuine procedural limitations identified by working clinicians. Practitioners routinely encounter significant operational barriers when utilizing conventional transcutaneous energy systems, including variable dermal penetration depths, unpredictable bulk heating, superficial epidermal thermal

risks, and cumbersome handpieces that hinder precise anatomical vectoring. Superficial treatments often deliver modest, short-lived skin tightening because transcutaneous energy dissipates before reaching the deep fibroseptal networks responsible for structural tissue support.

The development of the platform originated from an extensive evaluation of clinical challenges experienced by aesthetic physicians, dermatologists, and plastic surgeons. Frontline treatment feedback gathered by EVO SKIN from active medical aesthetic practices highlighted the necessity of bypassing the epidermis to treat soft-tissue laxity at its anatomical source. Clinicians underscored the demand for an endolaser system that delivers reliable photothermal contraction directly within the reticular dermis and subcutaneous adipose planes while maintaining absolute safety and low patient recovery times.

To transform these practical clinical insights into an advanced medical device, the development team partnered with Aesthelab in Pozzuoli (Naples), Italy. Drawing upon a distinguished engineering heritage in medical aesthetic technologies, including the invention of Electroporation Needle-Free Mesotherapy, Italian designers engineered a balanced, high-performance GaAIAs semiconductor diode delivery system specifically configured for delicate subcutaneous tissue manipulation. This collaborative bridge between American clinical demand and Italian engineering heritage established a treatment platform designed by clinicians for clinicians.

The Evolution of ENDO ONE from Practitioner Demand to Clinical Platform

Translating frontline procedural requirements into durable, clinic-ready medical hardware demands rigorous technical discipline and repeated engineering iterations. Early development phases focused systematically on eliminating common operational frustrations through purposeful hardware and software integration. Medical practitioners frequently report that conventional laser systems suffer from optical power instability, excessive handpiece weight, and complex interface navigation that disrupt clinical treatment flow.

Engineering specialists addressed these limitations by refining every hardware element through iterative laboratory benchmarks and ergonomic assessments. The solid-state GaAIAs diode architecture was optimized to ensure stable laser output with minimal thermal drift during extended operational cycles, guaranteeing consistent photothermal dosing across every vector pass. The handpiece assembly and fiber coupling mechanisms were balanced to provide immediate tactile feedback, allowing the clinician to sense subtle variations in tissue density and resistance as the fiber navigates subcutaneous planes.

The resulting platform embodies a clear development philosophy centered on clinical relevance, operational reliability, and patient safety. Aesthetic medical practices benefit from a platform engineered to support demanding daily procedural schedules without performance degradation. By prioritizing mechanical durability and optical precision, the system establishes a dependable clinical foundation that enables aesthetic physicians to execute sophisticated rejuvenation protocols with consistent confidence.

Technical Precision Designed for Modern Aesthetic Clinic Workflows

Clinical efficiency requires treatment platforms that integrate effortlessly into existing treatment suites and outpatient surgical facilities. Bulky consoles, prolonged calibration sequences, and unintuitive user interfaces create procedural friction, increase staff preparation times, and reduce overall clinical productivity.

The system features an ergonomic form factor weighing eight and a half kilograms, housed within a compact chassis measuring 39 centimeters by 32 centimeters by 23 centimeters. This lightweight design allows clinical staff to transport the device smoothly between treatment rooms on a dedicated rolling cart. A high-resolution touchscreen interface streamlines parameter selection, enabling practitioners to configure patient-specific protocols, pulse durations, and energy levels within seconds.

Precision optical fibers deliver targeted photothermal energy through minimal puncture sites created with a fine-gauge needle, eliminating the need for surgical incisions, sutures, or visible scars. The system incorporates an adjustable 650 nanometer red aiming beam that transilluminates clearly through the skin, granting the operator continuous visual confirmation of the fiber tip position and subcutaneous depth throughout the procedure. Clinicians execute steady retrograde passes at roughly one centimeter per second. Continuous non-contact thermal monitoring confirms that epidermal temperatures stay safely below 40 to 42 degrees Celsius. Clinicians achieve controlled tissue tightening and collagen stimulation through direct subcutaneous vectoring, delivering immediate tissue retraction while stimulating sustained neocollagenesis across the treated vectors.

Expanding Practice Treatment Portfolios with Versatile Laser Capabilities

Practices seeking sustainable commercial expansion require versatile platforms capable of addressing diverse facial and body concerns without excessive equipment redundancy. Multi-application capability protects capital investment while empowering clinics to serve broader patient cohorts across both preventative and restorative aesthetic indications.

Clinical applications encompass comprehensive tissue tightening and soft-tissue remodeling across both facial and body contours. Practitioners routinely address skin laxity across the mid-face to elevate cheeks and refine malar contours, while simultaneously treating the mandibular border and submental tissues to sharpen the jawline. For delicate indications, ultra-fine vectoring enables periorbital fine line smoothing and subtle skin remodeling. Furthermore, high-power delivery supports substantial body firming across postpartum abdominal laxity, upper arm flaccidity, and inner thigh corridors.

By integrating multiple procedural pathways into a single console, practices can expand service menus and capture high-demand rejuvenation volume with predictable consumable costs. Delivering noticeable aesthetic improvements in a single outpatient session under local tumescent anesthesia enhances patient satisfaction, strengthens practice reputation, and supports long-term clinical retention.

International Quality Standards and Professional Collaboration

Global aesthetic practices demand verifiable manufacturing quality and dedicated technical support before introducing new energy-based platforms. International clinical adoption depends on establishing durable partnerships between equipment providers and clinical stakeholders.

Headquartered in Pozzuoli, Italy, the engineering and operational team at ENDO ONE SRL maintains direct technical communication with international distribution partners and clinical users, operating in alignment with ISO 13485 medical device quality management standards. Ongoing practitioner training programs and technical documentation support safe clinical onboarding and consistent procedural execution. Standardized clinical guidelines ensure that medical teams master treatment depths, fiber vectoring, and thermal safety parameters.

The brand continues to expand its clinical presence across key international markets, collaborating with aesthetic physicians to document clinical outcomes and refine procedural protocols. Practitioners gain

access to an established professional network dedicated to clinical excellence and procedural innovation. This continuous exchange of clinical data ensures that treatment recommendations evolve alongside real-world aesthetic advancements.

Connecting with ENDO ONE for Clinical Evaluation and Practice Integration

Selecting an energy-based rejuvenation platform requires careful analysis of clinical efficacy, equipment ergonomics, and post-purchase technical support. Prospective clinics can evaluate device specifications, procedural videos, and clinical documentation to determine operational fit.

Practices interested in integrating advanced skin rejuvenation for aesthetic clinics can connect directly with clinical advisors through the official digital consultation portal. Experienced specialists provide detailed platform overviews, consumable configurations, and commercial integration guidance tailored to professional aesthetic facilities. Prospective partners can visit the official website at <https://www.endo-one-prime.com/> to submit formal inquiries, evaluate system specifications, and access comprehensive procedural resources.

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