

Collagen Hydrolysates, from Beauty Supplements to Supporting Wound Healing: How Solid is the Rationale? An Overview and Insights for Regenerative Medicine Practitioners

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Abstract

Introduction and purpose: Aquatic organisms, including invertebrates such as sponges, mollusks, and jellyfish, are sources of environmentally friendly marine collagen and low-molecular-weight bioactive oligopeptides, which are purified using advanced technologies. Since the early 2010's, numerous high-quality experimental and human studies have explored the properties of hydrolyzed marine collagen fragments as systemic functional ingredients in regenerative medicine. This review discusses these properties and the rationale behind them, with a focus on wound healing. Mechanisms underlying chronic wound healing may provide a strong foundation for the still-missing, high-level clinical studies, particularly in patients with diabetic and pressure ulcers.

Methods: This review examines only academically significant studies published in PubMed-indexed journals with significant impact factors, supplemented by a few contributions from Google Scholar for methodologically sound *in vitro* and animal studies.

Results: Activation of skin fibroblasts and other mesenchymal cells underpins the systemic regenerative properties of highly purified hydrolyzed marine collagens. For example, 50 µg/mL of hydrolyzed marine collagen peptides nearly replicated *in vitro* the accelerated cell migration induced by 10 µg/mL of recombinant human epidermal growth factor. Faster wound healing, associated with increased collagen neosynthesis, is

accompanied by increased immunohistochemical expression of platelet-endothelial cell adhesion molecule-1, basic fibroblast growth factor, and transforming growth factor β -1. The potential supportive role of collagen hydrolysates in managing insulin resistance could benefit the treatment of chronic diabetic and pressure ulcers.

Conclusion: An increasing number of preclinical and human studies highlight the systemic regenerative properties of hydrolyzed marine collagens and their excellent safety profile. The evidence for their regenerative properties in aesthetic skin rejuvenation appears solid. Pre-clinical evidence supporting wound healing is also growing. Unfortunately, sound clinical studies confirming the experimental evidence in everyday wound care practice are still lacking, and long-term safety remains a primary concern.

Keywords: Collagen hydrolysates; Diabetic foot care; Fibroblast activation; Hydrolyzed marine collagen; Regenerative medicine; Wound care

Introduction

Marine collagen hydrolysates and their growing role as beauty supplements

Collagen derivatives from cowhide and other land mammals, along with more recent marine collagen supplements, have gained prominence in the global beauty supplement industry, despite the limited number of well-designed studies. The decline in native collagen with age after early adulthood, which contributes to visible signs of skin aging, such as wrinkles and sagging, especially among women after menopause, has fueled the recent widespread acclaim for collagen derivative supplements [1].

Seawater and freshwater organisms including invertebrates like sponges, mollusks, echinoderms, and jellyfish are sources of marine collagen [2-4]. Unlike bovine, porcine, or human recombinant Type 1 collagens, all marine collagens are naturally less cross-linked, making them biophysically unique. They exhibit lower tensile strength, which is less critical in the low-gravity aquatic environment, but show greater flexibility [4]. Collagen from aquatic sources addresses religious concerns and avoids the risks of prion disease transmitted by land-mammal collagens; additionally, it avoids the risks of contamination with heavy metals and marine pollutants [2,5]. Highly purified marine collagens are also environmentally friendly because they originate from the estimated 20 million tons of waste organic residues produced annually by the global industrial fishing industry [2,5]. Moreover, published studies and a recent review of their regenerative potential indicate that marine collagen hydrolysates have safety profiles similar to placebo, with only occasional mild nausea and negligible risks of Type 1 immediate hypersensitivity [2].

The high bioavailability of hydrolyzed marine collagens after oral intake explains their widespread distribution throughout the body [2]. Several studies discuss their skin-rejuvenating effects and potential to reactivate functions in joints and other organs. However, these studies are outside the scope of this review, which focuses on the potential benefits of marine collagen fragments for wound healing, especially in slow-healing wounds.

Because they were introduced only recently, the number of published papers on hydrolyzed marine collagens indexed in PubMed is limited. This is especially true in high-impact, peer-reviewed journals. A PubMed search conducted in early January 2026, using the search string “marine collagen peptides” and limited to English-language articles, returned only 30 citations, including a review in aesthetic regenerative medicine by two authors of this manuscript [2]. Many of these papers mainly focus on purification procedures and other technical, non-medical issues.

Wound care dressings often use hydrolyzed collagen-based matrices, and a substantial body of literature supports their use. Still, as of early January 2026, no paper in PubMed-indexed journals appeared to have reviewed the effectiveness of orally administered marine collagen hydrolysates for wound care in human patients.

A role for marine collagen oligopeptides in chronic wound healing is soundly based

Despite the lack of clinical studies on marine collagen bioactive fragments in wound care, especially for chronic, difficult-to-heal wounds such as diabetic and pressure ulcers, the rationale for their widespread use appears solid. Fibroblast dysfunction and abnormally high levels of Matrix Metalloproteinases (MMPs) disrupt extracellular matrix remodeling by reducing collagen content [6-8]. Histochemical studies in experimental diabetic models confirm this disruption, showing delayed appearance of sparse, poorly stained precollagen (reticulin) fibers and weak staining for acid and neutral mucopolysaccharides [6]. Histologically, scar formation is delayed, with abnormal persistence of early vascularization and cellular activity. In fibroblasts, wound-healing mediators such as *COL16A1* are poorly expressed. Likewise, several intracellular signaling pathways, including the transcription factors *PLAGL1*, *RUNX2*, and *ZKSCAN7*, which appear downregulated in poorly healing lesions, respond favorably to regenerative treatments such as platelet-rich plasma [9,10].

The purpose of this review is to discuss the properties of bioactive hydrolyzed marine collagen oligopeptides, with a particular focus on wound healing, and to present the rationale for their potential effectiveness in this indication. The mechanisms underlying chronic wound healing, as demonstrated preclinically and discussed herein, may provide a strong foundation for the high-level clinical studies that remain warranted.

The rationale appears especially well documented for diabetic ulcers, but may extend to other hard-to-heal chronic wounds, as suggested, in patients with chronic tissue ischemia and pressure ulcers, usually over bony prominences, through the oral administration of bioactive oligopeptides derived from gelatin or heat-denatured collagen from land mammals [11].

Materials and Methods

The authors adopted selective criteria for their review of published papers on hydrolyzed marine collagens in wound management:

- Inclusion only of well-described, methodologically sound, and reproducible studies, published in PubMed-indexed journals with notable impact factors.
- Exclusion of all studies of questionable quality published in non-indexed, non-peer-reviewed journals.

These selection criteria aimed to minimize the risk of promotional bias. The early 2010's mark the period when the first indexed, academically credible papers on hydrolyzed marine collagens appeared in the literature; this period was set as the search cutoff for the consulted databases. The authors made the final decision about which studies to include in their review during an online consensus meeting held in mid-November 2025. They adhered as closely as possible to the PRISMA guidelines for reviews while maintaining a general narrative style to make their manuscript easier to read.

Results and Discussion

Even if still limited, evidence from the literature on marine collagen-derived oligopeptides indicates a notable oral bioavailability and systemic distribution (Figure 1) [12]. After oral intake and systemic distribution, the low-molecular-weight ^{14}C -labelled fragments persist in target tissues for more than two weeks—skin, muscles, and mesenchymal tissues involved in slow-healing wounds [12,13].

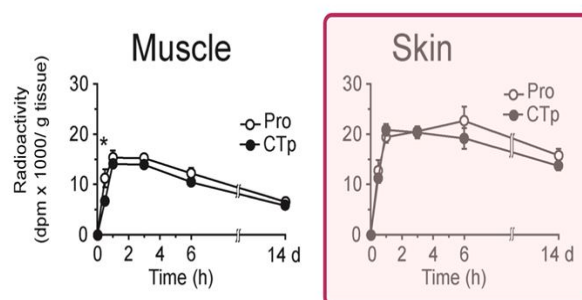


Figure 1: Distribution of radioactivity in the skin and muscle, normalized to 1 g of wet tissue, in laboratory Wistar rats during the first six hours and after 14 days following oral administration of ^{14}C -labelled hydrolyzed collagen fragments (CTp) and proline (Pro); mean $\pm$ standard error. Reproduced and adapted from Sibilla S, et al. under a Creative Commons CC BY-NC 4.0 License [12].

Efficient absorption, mediated by transporters in the intestinal brush border membrane, such as the proton-coupled oligopeptide transporter 1 (PepT1), prevents digestion of marine collagen fragments into individual amino acids [2]. The high systemic bioavailability of marine collagen oligopeptides, which is indistinguishable from that of the amino acid proline, sets hydrolyzed marine collagens apart from collagen fragments derived from land mammals. Absorption is up to 50% more efficient compared to mammalian collagens, likely owing to the higher content of glycine, serine, and threonine and the lower levels of proline and hydroxyproline [14,15].

Safety and oral-route bioavailability of hydrolyzed marine collagens

In preclinical investigations, marine collagen oligopeptide hydrolysates showed negligible impact on pro-inflammatory mediators in animal studies [2]. The low histidine content, a histamine precursor, indicates the low antigenic potential of marine fragments and reduced risk of type-1 immediate hypersensitivity [16]. These hydrolysates also act as potent antioxidants, with free radical scavenging activity comparable to E321, a widely used food preservative (Figure 2) [5].

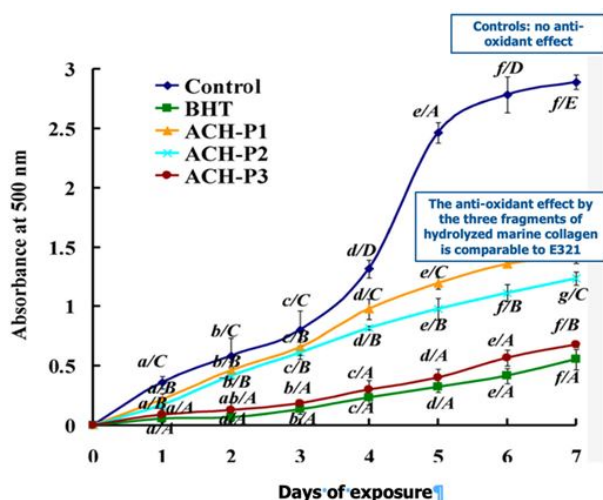


Figure 2: The decrease in spectrophotometric absorbance by cell membranes over seven days directly correlates with the oxidative damage caused by free radicals. The antioxidant effectiveness of the positive control E321 and three peptides from the same batch of marine collagen is comparable. Control: inert negative control. Reproduced and adapted from Geahchan S, et al. under a Creative Commons CC BY-NC 4.0 License [5].

The biological properties of hydrolyzed marine collagens

Activation of skin fibroblasts underpins the regenerative properties of marine collagen hydrolysates after oral intake, both in the skin and systemically. These properties are unique to marine breakdown fragments, compared with dermally infiltrated intact collagen [2,16-19]. Fibroblast activation requires a polypeptide structure, as exposure to proline and hydroxyproline, the amino acids typical of all collagens, fails to exhibit the same biological behavior (Figure 3) [12].

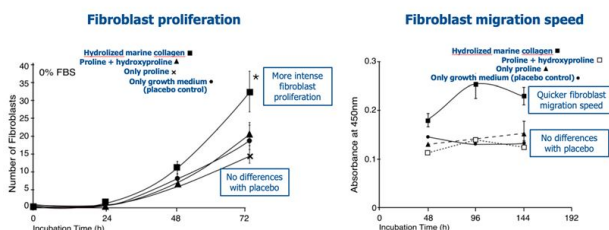


Figure 3: Effects on *in vitro* proliferation and migration speed of human fibroblasts after exposure to marine collagen fragments for 72 hours, compared with the growth medium (inert placebo), the amino acid proline, and a proline-hydroxyproline mixture; * $p < 0.05$ versus placebo. Reproduced from Sibilla S, et al. under a Creative Commons CC BY-NC 4.0 License [12].

The effect on fibroblasts appears direct. For example, the hydrophobic peptide F5 from Mozambique tilapia (*Oreochromis mossambicus*) fish scales not only scavenges reactive oxygen species and prevents oxidative stress-induced damage to fibroblasts, but also docks to the active sites of matrix metalloproteinases MMP-1, MMP-3, and MMP-12, which break down collagen and elastin [3]. Furthermore, in laboratory animals supplemented with oral marine collagen hydrolysate for 24 months, there is an increase in the expression of Type I and III collagen compared to

aged untreated controls. This increase in collagen deposition occurs alongside a decrease in Metalloproteinase-1 (MMP-1) expression and higher tissue levels of the MMP-1 inhibitor, as revealed by immunohistochemical and Western blot analysis [20].

These changes are linked to the activation of the SMAD signaling pathway mediated by the SMAD group of related proteins acting as the primary signal transducers for receptors of the Transforming Growth Factor beta (TGF- β) superfamily and the up-regulated expression of the SMAD activator TGF- β RII (T β RII), supporting the direct effect on fibroblasts [21]. The activation of the Hyaluronan Synthase 2 (HAS2) transcription pathway stimulates the synthesis of high-molecular-weight hyaluronan [20].

The rationale behind these properties is sound and unsurprising, aligning with the innovative concept of transitioning from inflammatory repair to non-inflammatory tissue regeneration (Anti-Inflammatory Regenerative Medicine, AIMED) [22]. The small peptides, known as matrikines, released during the breakdown of extracellular matrix collagen, are highly homologous to marine collagen fragments and act as good models of the biological effects for marine collagen hydrolysates (Figure 4). Molecular docking studies demonstrate the high affinity of several matrikines for receptors for chemokines, cytokines, ion channels, and growth factors [2,23]. Examples include the collagen matrikine α 1 C-1158/59, which promotes wound healing and cell migration, and the glycine- and proline-containing matrikine PGP, which binds at the chemokine receptors C-X-C motif chemokine receptors 1 and 2 (*CXCR1* and *CXCR2*), thereby regulating both the inflammatory and epithelialization phases of wound healing [23,24].

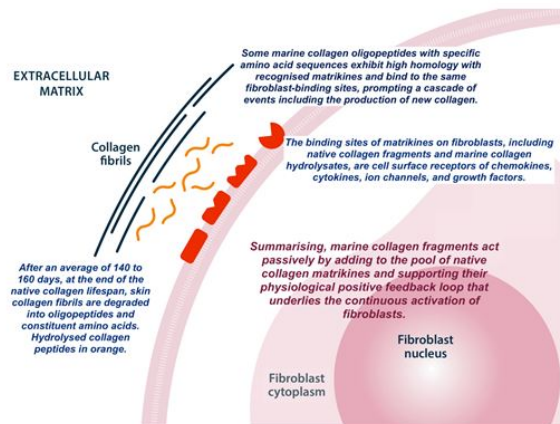


Figure 4: The effects of hydrolyzed marine collagens on fibroblasts resemble and complement those of physiological matrikines as native collagen reaches the end of its lifespan in the dermis and subdermal tissues. Reproduced and adapted from Bartoletti E, et al. under a Creative Commons CC BY-NC 4.0 License [2].

The promising rationale for hydrolyzed marine collagens in wound care

A recent review examined the emerging benefits of orally administered marine collagen hydrolysates in aesthetic medicine and plastic surgery [2]. Building on those findings, the authors present some ideas about the rationale and expected advantages of marine collagen hydrolysates in wound care.

Experimental *in vitro* exposure to marine collagen peptides promotes wound healing in a dose-dependent manner, as demonstrated in several early preclinical studies from the mid-2010's. In a 2015 study based on the *in vitro* scratch closure assay, with marine collagen proteolysate concentrations ranging from 6.25 to 50 $\mu\text{g/mL}$, there was no difference in the rate of cell migration and repair ability of the injured cell monolayer between fibroblasts exposed to 50 $\mu\text{g/mL}$ of marine collagen fragments and those exposed to 10 $\mu\text{g/mL}$ of recombinant human Epidermal Growth Factor (rhEGF), used as positive controls [25]. The same year, a study in young male Balb/c mice injected in the tails (test solutions: 80 μL) with Acid-Soluble Collagen (ASC) from haddock (*Melanogrammus aeglefinus*) skin and four ASC hydrolysates with different average molecular weights H1 — (53.8 kDa), H2 (23 kDa), H3 (14.4 kDa), and H4 (6.5 kDa) — found that ASC and the hydrolysates with higher molecular weights (H1 and H2) shortened clotting and bleeding times compared to controls (saline) [26]. Moreover, during experimental burn wound healing, skin cellularity and maturity increased in both the epidermal and dermal layers compared to the control group. After a week, blood capillaries appeared in scalded skin treated with hydrolysates but not in controls (Figure 5) [26].

In another similar study conducted two years later, chum salmon skin collagen fragments accelerated wound healing after cesarean section. This was associated with increased immunohistochemical expression of platelet-endothelial cell adhesion molecule-1, basic fibroblast growth factor, and transforming growth factor β -1 [27]. The dose-dependent increase in fibroblast proliferation following exposure to marine collagen peptides is also chronologically connected to the dose-dependent activation of the fibroblast intracellular NF- κ B p65 signaling pathway, followed by enhanced signaling along other secondary intracellular pathways [28].

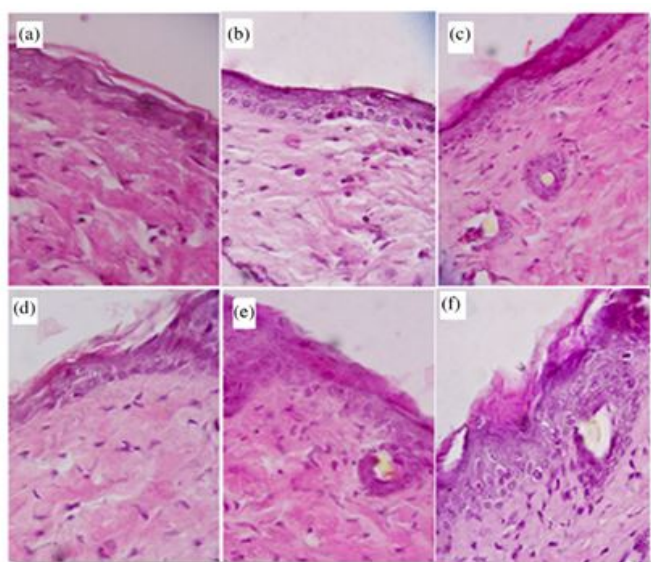


Figure 5: Increased cellularity and maturation of cells in the epidermis and dermis of scalded mouse skin ($\times 400$) after treatment with (a) saline (control group); (b) haddock skin ASC; (c) H1; (d) H2; (e) H3; (f) H4. Reproduced from Dang QF, et al. with permission from the publisher [26].

Monolayer scratch wounds close quickly. In a previously discussed wound-healing scratch assay, the repair rate of the scarified monolayer of Human Umbilical Vein Endothelial Cells (HUVEC) after 48 hours

was about 74-75% following exposure to 6.25 $\mu\text{g/mL}$ of two slightly different formulations of marine collagen peptides from the jellyfish *Rhopilema esculentum*. The repair rate of vehicle-treated, scarified control HUVEC cells was less than 50%. Starting at 18 hours, the difference in wound closure was significant compared to controls (Figures 6 and 7) [19].

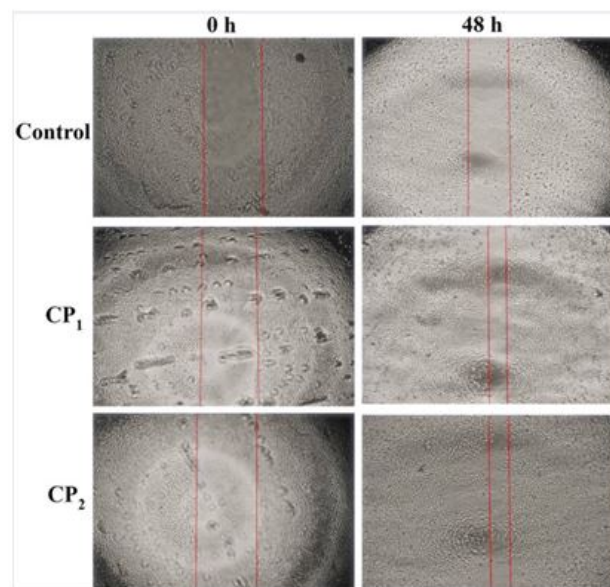


Figure 6: Migration of HUVEC cells in a wound-healing scratch assay after exposure to two slightly different formulations of jellyfish collagen peptides (6.25 $\mu\text{g/mL}$) compared with migration after exposure to vehicle (control) at baseline and after 48 hours. Red lines mark the edges of the wound gap. Reproduced from Hu Z, et al. under a Creative Commons CC BY-NC 4.0 License [19].

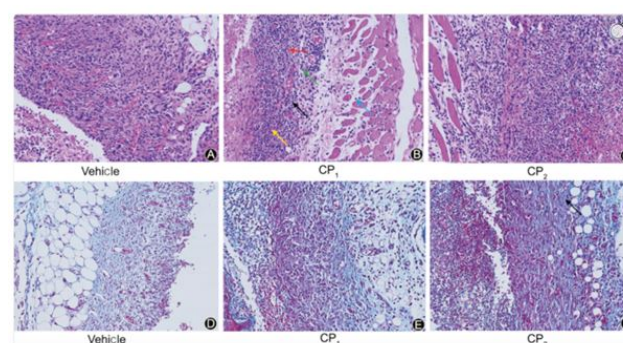


Figure 7: Skin sections from mice treated with 0.9 g/kg body weight collagen peptides or vehicle on day 6; magnification 200X. A to C micrographs: Skin sections showing fibroblasts (black arrows), fiber cells (red arrow), new blood vessels (yellow arrow), inflammatory cells (green arrow), and interstitial connective tissue around muscle cells (blue arrow). Hematoxylin and eosin stain. D to F micrographs: Blue-stained collagen (black arrow). Masson trichrome stain. Reproduced from Hu Z, et al. under a Creative Commons CC BY-NC 4.0 License [19].

Most of the limited number of published papers on hydrolyzed marine collagens indexed in PubMed are experimental, conducted *in vitro* and in laboratory models, including studies on wound healing.

Surprisingly, despite the sound rationale for marine collagen hydrolysates, there are no studies specifically focused on pressure injuries in laboratory models, nor have human studies examined the potential of hydrolyzed marine collagen in wound management. The results of the only (as far as the authors know) double-blind placebo-controlled study investigating the effectiveness of non-marine collagen hydrolysates could approximate the expected positive outcomes of the still-absent human studies [11]. Patients with pressure ulcers receiving standard care were randomized to twice-daily, high-dose oral intake of low-molecular-weight hydrolysates from heat-denatured pork skin gelatin containing either low or high levels of the prolylhydroxyproline (Pro-Hyp) and hydroxyprolylglycine (Hyp-Gly) dipeptides or placebo (maltodextrin TK-16). The primary efficacy parameters, the two scoring scales Pressure Ulcer Scale for Healing (PUSH) and Pressure Sore Status Tool (PSST), as well as the wound area, were significantly lower at the end of 16 weeks compared to placebo, with only minor differences between the two formulations. The PUSH scores for the slightly more effective formulation were 6.46 ± 0.98 and 9.26 ± 2.09 for the placebo ($p < 0.01$), while the PSST scores were 19.71 ± 3.08 and 23.38 ± 3.85 ($p = 0.01$), and the wound areas were 3.19 ± 2.88 and 5.00 ± 3.88 ($p = 0.027$). Over 71% of individuals showed improvements in their PUSH and PSST scores. According to the authors [11], the significant improvements in PUSH and PSST scores and wound area could be attributed to the physiological activity of the systemically absorbed oligopeptides.

Conversely, evidence is accumulating on the potential role of collagen hydrolysates in modulating insulin resistance, an interesting finding given the rising prevalence of diabetic ulcers [29-33]. Helping control insulin resistance and regulate glycemic metabolism could be beneficial, as the prevalence of diabetic foot, a serious complication of diabetes, is estimated at 4-10%. It primarily affects older patients, with a lifetime risk of about 15% for diabetic individuals to develop this complication [34].

Although not strictly derived from an aquatic organism, exposure to the skin collagen polypeptide GPAGAP from *Andrias davidianus*, a giant, critically endangered amphibious salamander from the Yangtze River basin in central China, improved insulin resistance in the HepG2 cell line through multiple-target, multi-pathway mechanisms. Molecular docking studies revealed interference with several intracellular signaling pathways involved in type 2 diabetes mellitus and its complications, including the PI3K-Akt and AGE-RAGE pathways, as well as α -glucosidase inhibition [29]. Marine collagen polypeptides from the bones of *Thunnus albacares*, a yellowfin tuna from tropical and subtropical oceans, also demonstrated competitive inhibition of α -glucosidase [30]. Even more convincing is the evidence that marine collagen hydrolysates inhibit Dipeptidyl Peptidase IV (DPP-IV), which degrades the intestinal incretin hormones GLP-1 (Glucagon-Like Peptide-1) and GIP (Glucose-Dependent Insulinotropic polypeptide), thereby promoting insulin secretion, inhibiting glucagon release, and delaying gastric emptying [30]. Already in 2014, the tripeptide GPA from fish scale collagen proved to be a competitive inhibitor ($K_i = 4.5$ mM) of DPP-IV; in contrast, collagen hydrolysates from pig and cattle skin and chicken feet showed no inhibitory activity against DPP-IV [31]. Collagen polypeptides from the Far East fish *Selar crumenophthalmus* also increased GLP-1 expression [32]. Molecular docking analysis identified nine collagen oligopeptides from bighead carp skin with potential DPP-IV inhibitory activity, with the fragment PPGF exhibiting the highest potency ($IC_{50} = 4.63$ nM) [33]. Emerging clinical

evidence from well-designed, double-blind, actively controlled trials indicates that patients with type 2 diabetes treated with fish collagen fragments experience significantly better short- and medium-term control, including reductions in fasting blood glucose and glycosylated Hemoglobin (HbA1c) over a three-month period [34].

Building on this preclinical evidence, one of the authors, M.R., is currently coordinating a pilot study sponsored by his company that uses a patented hydrolyzed marine collagen supplement (NUXIDIO, Asteria Pharma Srl, Milan, Italy) to explore the potential of oral marine collagen hydrolysates in the early stages of diabetic foot ulcers as an adjunct to standard diabetic ulcer treatment in the early stages of diabetic foot ulcers [35-40].

Conclusion

The rationale for the value of collagen oligopeptides, particularly their potential to support wound healing *via* oral intake, appears trustworthy. This conclusion is primarily based on methodologically sound, reproducible preclinical studies. However, there are no clinical studies specifically addressing wound care. This lack of human studies hinders a balanced discussion of the advantages and disadvantages of hydrolyzed marine collagens in wound management. It also prevents drawing definitive conclusions beyond a cautious acknowledgment of the consistent preclinical rationale and the potential for clinical benefits in support of standard conservative wound management, such as surgical debridement, correction of peripheral arterial insufficiency, offloading the wound, daily saline or similar dressings to maintain a moist wound environment, and infection control.

Conversely, clinical investigations for the indication of skin rejuvenation for aesthetic purposes are available and discussed in systematic reviews and meta-analyses. However, even in that indication, acceptance of generally favorable outcomes is not universal, as suggested by subgroup meta-analyses by funding source corporate-dependent or corporate-independent.

Although persuasive, experimental data on wound management from *in vitro* and animal models cannot escape debate and doubts about their predictive value for humans. These considerations led the Corresponding Author to oversee an academic clinical research program on early diabetic foot care with medium-term follow-up. Based on the described preclinical evidence, short-term benefits are likely; however, only long-term studies will confirm their sustainability and prevent overestimating benefits. Verifying the long-term safety of marine collagen fragments is another critical issue. Chronic wound care is a prolonged medical and surgical endeavor, and currently, there is limited information on long-term safety, including for the anti-aging and skin rejuvenation indications.

In conclusion, although it may still be too early to provide definitive recommendations to wound care specialists about adjunctive biologically active marine collagen hydrolyzed oligopeptides, pending confirmation from the well-designed clinical trials currently underway, the substantial experimental evidence to date suggests potential benefits in clinical settings. Clinicians may consider the existing evidence when making decisions on a case-by-case basis.

Ethical Statement

The submitted review covers previously published information, thereby waiving any formal preliminary requirement for Institutional Review Board approval.

Data Sharing Statement

The minutes of board discussions are archived and can be provided by the Corresponding Author upon reasonable request.

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Authors' Contribution

All authors made significant contributions to the manuscript, whether in conception, study design, execution, data acquisition, analysis, and interpretation, or across all these areas; participated in drafting, revising, or critically reviewing the article; gave final approval of the manuscript for submission; and agreed to be accountable for all aspects of the work. Dr. Mauro Raichi was the primary medical writer.

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