Collagen Injectables for Aesthetic and Regenerative Medicine Applications

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Soft tissues diseases significantly affect patients quality of life and usually require targeted, costly and sometimes constant interventions. With the average lifetime increase, a proportional increase of age-related soft tissues diseases has been witnessed. Due to this, the last two decades have seen a tremendous demand for minimally invasive one-step resolutive procedures. Intensive scientific and industrial research has led to the recognition of injectable formulations as a new advantageous approach in the management of complex diseases that are challenging to treat with conventional strategies. Among them, collagen-based products are revealed to be one of the most promising among bioactive biomaterials-based formulations. Collagen is the most abundant structural protein of vertebrate connective tissues and, because of its structural and non-structural role, is one of the most widely used multifunctional biomaterials in the health-related sectors, including medical care and cosmetics. Indeed, collagen-based formulations are historically considered as the "gold standard" and from 1981 have been paving the way for the development of a new generation of fillers. A huge number of collagen-based injectable products have been approved worldwide for clinical use and have routinely been introduced in many clinical settings for both aesthetic and regenerative surgery.

collagen injectable collagen medical devices

1. Introduction

Soft tissues loss could be due to iatrogenic, traumatic, pathological, or physiological reasons. Aside from significantly affecting patients' quality of life, their surgical management requires targeted, costly and sometimes constant interventions. With the average life increase, a proportional increase of age-related soft tissues diseases has been witnessed. Due to this, recent decades have seen a tremendous demand for soft tissue reconstruction strategies and one step resolutive procedures. Intense scientific and industrial research has been conducted to develop innovative approaches or optimize current solutions. Among them, in the last two decades injectable formulations have attracted even more interest for both aesthetic and regenerative surgery for their versatility and multifunctionality (**Figure 1**). Indeed, injectable scaffolds could be used in large and irregularly shaped lesions for a huge variety of damaged tissues, as well as providing temporary pain relief and functional improvement with a single treatment. Thus, injectable formulations could reduce the number of surgical procedures, costs, times and accelerate healing rate and quality.



Figure 1. The increasing research interest in injectable formulations and dermal filler. Articles indexed in Scopus (<u>www.scopus.com</u>) with the keywords 'injectable' and 'dermal filler' and published from 2000 to 2022 (last accessed on 27 May 2022).

The popularity of minimally invasive techniques increased rapidly for several reasons. A principal factor is the acceptance of soft tissue fillers among patients that are not ready for permanent treatments ^[1]. In the case of patients not wishing to undergo surgery, an easier procedure would generally be more accepted. Moreover, compared to undergoing more invasive surgery, fillers offer the patient less discomfort and a shorter recovery time, making them very practical in the resolution of minor-serious disease and allowing patients to return immediately to their daily routine ^{[1][2]}. Minimally invasive therapies would give a better quality of life also for that part of population that would otherwise not survive the trauma induced by conventional surgeries. Moreover, they could delay the execution of invasive surgical procedures for the implantation of permanent devices ^[3]. In the case of a staged surgical intervention, the use of injectable systems may avoid the need for multiple invasive operations, thus reducing the related morbidities and negative aesthetic effects associated with repeated procedures ^[4]. With regard to aesthetic treatments, minimally invasive therapies are preferred as they are less impacting and give a more natural look. Moreover, the lack of an external incision or an autologous tissue donor site is preferred because the absence of scarring is usually socially and psychologically more accepted.

From the surgeon's point of view, the advantages of minimally invasive procedures include principally the need for fewer resources (e.g., operating room, staff, equipment, and time). Being simpler, transcutaneous injections require less operating room staff and time. The pro-regenerative action of injectables would reduce operating room time also because they would be able to restore physiological conditions with a single injection. However, it should not be forgotten that simpler procedures are not less exhausting and do not require less experience. Like any surgical procedure, minimally invasive therapies require adequate knowledge in order to reach the best outcome and avoid unwanted adverse events.

Thus, not only clinicians' but especially patients' preference for fewer invasive and expensive procedures has undoubtedly promoted their use [4][5][6][7][8][9][10]. An injectable formulation for soft tissues reconstruction currently relies on two main approaches, involving autologous tissue displacement (e.g., lipofilling, platelet-rich plasma) or biomaterials-based filling ^[5]. Both approaches have some advantages and drawbacks. Autologous materials

provide the most physiological solution (no adverse events or immune reactions) but suffer from donor site morbidity, volume resorption rate variability, and double surgery requirements. Moreover, their harvesting is a time-consuming procedure that requires double intervention. Alternatively, biomaterials offer an off-the-shelf solution with immediate results and should be distinguished as non-resorbable and resorbable, depending on their half-life. Non-resorbable solutions (e.g., silicone, poly(methyl methacrylate), polyvinylpyrrolidone, polyacrylamide), are permanent (last more than 2 years) but usually suffer from mild-severe adverse reactions (i.e., granuloma, implant encapsulation, persistent pain or rejection) that limit patient satisfaction and could require implant removal surgery [6][7][8][9][10][11][12]. Contrarily, resorbable formulations are usually based on natural biomaterials (i.e., collagen, hyaluronic acid, calcium hydroxyl apatite) and last 6–18 months [13][14][15][16]. Their durability depends on many factors such as the raw material type, product cross-linking degree, lost tissue extension, disease site and etiology, and patient metabolism, age and co-morbidities. The most used resorbable dermal fillers are collagen or hyaluronic acid based.

Collagen is the most abundant structural protein of vertebrate connective tissues ^{[17][18][19][20][21][22][23][24][25]} and plays a crucial structural role for the maintenance of tissues' architecture, shape and mechanical properties ^[20]. Moreover, by mediating a fundamental inter- and intracellular signaling it dictates specialized regulatory functions, especially during development and repair processes ^{[26][27][28][29][30][31][32]}. Type I collagen is one of the most widely used biomaterials in the health-related sectors, including medical care and cosmetics ^{[17][18][19][20][21][22][23][24][25][33]}. Several collagen-based injectable products have been approved for clinical use and used in many clinical settings.

2. Collagen as Biomaterial

Collagen is the most abundant structural protein of vertebrate connective tissues, and accounts for about the 30% of the total body protein content [17][18][19][20][21][22][23][24][25]. The collagen family is a group of proteins that share a unique molecular fingerprint that is characterized by the presence of a right-handed triple-helical domain formed by three left-handed polyproline-II helices [26][34][35]. This superfamily accounts for 28 members, named from type I to XXVIII according to the discovery order [34][36]. Type I collagen was the first to be discovered and accounts for the 70% of the total collagen found in the human body ^[26]. This protein is a hetero trimer of about 400 kDa consisting of two identical $\alpha 1$ (≈ 139 kDa) chains and one $\alpha 2$ (≈ 129 kDa) chain of about 1000 amino acid residues $\frac{[20][37]}{3}$. Both chains are characterized by the repetition of the Glycine-X-Y triplet, where the X and Y positions are usually represented by proline and hydroxyproline, respectively [34][37]. Hydroxylation of proline residues is a typical modification of collagen and, because it accounts for about 11-14% of total residues, it is commonly used as a marker to detect and quantify collagen in tissues ^{[35][38]}. Another peculiarity of fibril-forming type I collagen molecules is their ability to spontaneously assemble to form fibrils in which molecules are quasi-hexagonally packed and super-twisted in a right-handed structure along the longitudinal axis of the fibril [39][40][41]. Thus, collagen molecules are aligned parallel to one another with a staggering of about 67 nm (D-banding) and can assemble into fibrils that can be greater than 500 µm in length and 500 nm in diameter ^{[25][34][42][43]}. Then, fibrils assemble in fibers whose 3D arrangement is tissue specific.

Type I collagen not only covers a crucial structural role in tissue architecture maintenance but is actively involved in several biological and pathological processes ^[44]. The involvement of collagen in numerous cellular processes prompted research towards the use of collagen as biomaterial for the development of simplified ECM-like structures ^{[20][35]}. To this, several companies isolate medical-grade type I collagen from several sources and manufacture collagen-based implantable devices that are currently used in many clinical settings. Besides its advantages in term of biocompatibility for its physiological structural and non-structural functions, the use of collagen as biomaterial offers several advantages including low immunogenicity, tunable properties, and biodegradability. The low evolutionary gap and the high conservation of type I collagen amino acid composition among vertebrates make that homology up to 95% ^{[19][45][46][47][48]}.

3. Collagen-Based Injectable Formulations

More than 60 kinds of collagen-based fillers are available on the market, according to the end-use and they have routinely been introduced in many clinical settings (Table 1). The most common collagen extraction sources for the manufacture of collagen based injectable formulations are bovine, swine, porcine, equine and human derived, whose advantages and disadvantages are described in depth elsewhere [19][20][25]. Bovine collagen is one of the most commonly used fillers for effectively reducing wrinkles and other facial imperfections. More famous branded bovine-based collagen fillers are Zyderm[®], Zyplast[®], Contigen[®] (Allergan Inc., Dublin, Ireland), Artefill[®] (Suneva Medical, San Diego, CA, USA), and Artecoll[®] (Canderm Pharma Inc., Saint-Laurent, QB, Canada). Others include CHondroGrid[®] (Bioteck Spa, Arcugnano, Italy), Integra Flowable Wound Matrix[®] (Integra LifeScience Corp., Princeton, NJ, USA), Resoplast[®] (Rofil Medical International, Breda, The Netherlands), Atelocell[®] (KOKEN Co., Ltd., Bunkyo-ku, Tokyo, Japan). However, bovine collagen is known to be exposed to zoonosis (e.g., the foot and mouth disease and the group of the bovine spongiform encephalopathies, among which the most dangerous for humans is the transmissible spongiform encephalopathy) and to trigger allergies (about 2–4% of population) [49][50] ^[51]. In addition to the strict regulation to which all implantable products are subjected, two consecutive negative patient skin tests at 6 and 2 weeks are required before use [51][52]. This sensitivity has been considered generally acceptable for implants for human use and actually bovine collagen is principally used for the treatment of the [6][53][54][55][56][57][58][59][60][61][62][63][64][65][66][67][68][69][70][71][72][73][74] integumental (NCT01060943) and musculoskeletal apparatus [75][76][77][78][79][80][81][82][83][84][85][86][87][88][89][90] and to a minor extent for the gastrointestinal [91][92][93][94][95][96][97][98], urinary [99][100][101][102][103][104] and cardiovascular [105][106][107] systems. Recently, bovine collagen in fibrillar form has been employed as an organ protection system during thermal ablation of hepatic malignancies [108].

Table 1. Summary of clinically available type I collagen-based injectable formulations.

Source	Manufacturer	Product	Additives	Applications	Ref.
Equine	Euroresearch S.r.l. (Milan, Italy)	Nithya	_	Integumental	[<u>109</u>]
	www.euroresearch.it,	Linerase	_	Integumental	[<u>110][111][112][113]</u> [<u>114]</u>

Source	Manufacturer	Product	Additives	Applications	Ref.
	accessed on 14 February Nearme 2028 aly S.r.I. (Como, Italy) <u>www.salvecoll.com</u> , accessed on 14 February 2023	Salvecoll-E	_	Integumental	[<u>115</u>]
		Biocollagen gel	Type III collagen, bone spongy powder	Musculoskeletal	_
		Biocollagen crunch	Type III collagen, bone powder, bone spongy chips	Musculoskeletal	_
		ActivaBone CLX gel	Bone powder, exur, Vitamin C	Musculoskeletal	_
	Bioteck Spa (Arcugnano, Italy) <u>www.bioteck.com,</u> accessed on 14 February	ActivaBone Injectable Paste	Demineralized bone matrix, bone powder, exur, Vitamin C	Musculoskeletal	_
	2023	ActivaBone modulable paste	Demineralized bone matrix, bone powder, bone cortical and spongy granules, exur, Vitamin C	Musculoskeletal	_
		ActivaBone Crunch	Demineralized bone matrix, bone powder, cortical and spongy chips, exur, Vitamin C	Musculoskeletal	_
Bovine	Bioteck Spa (Arcugnano, Italy) <u>www.bioteck.com,</u> accessed on 14 February 2023	CHondroGrid	-	Musculoskeletal	[<u>90]</u>
	Integra LifeScience Corp. (Princeton, NJ, USA) <u>www.integralife.com</u> ,	Integra Flowable Wound Matrix	Glycosaminoglycans	Integumental	[<u>66]</u>
	2023	Helitene	-	Soft tissues	[<u>108</u>]
	Rofil Medical International (Breda, The Netherlands)	Resoplast	Lidocaine hydrochloride	Integumental	_
	Suneva Medical (San Diego, CA, USA) <u>www.sunevamedical.com</u> ,	ArteFill	Polymethylmethacrylate, lidocaine	Integumental	[<u>53][55][56][57][58]</u> [<u>59][60][61][62][63]</u>

Source	Manufacturer	Product	Additives	Applications	Ref.
	accessed on 14 February 2023				
	Datascope Corp., (Montvale, NJ, USA)	VasoSeal	-	Cardiovascular	[<u>107</u>]
	BioMimetic Therapeutics, LLC (Franklin, TN, USA) www.biomimetics.com, accessed on 14 February 2023	Augment	β-tricalcium phosphate, recombinant human platelet-derived growth factor-BB	Musculoskeletal	[75][77][78][79][80] [81][82][83][84][85] [86][87][88][89]
	KOKEN Co., Ltd. (Bunkyo- ku, Tokyo, Japan) <u>www.kokenmpc.co.jp</u> , accessed on 14 February 2023	Atelocell	Type III collagen	Integumental, gastrointestinal	[<u>64][65][91][92]</u> , NCT01060943
	B. Braun (Crissier, Switzerland) <u>www.bbraun.com,</u> accessed on 14 February 2023	Gelofusine	_	Cardiovascular	[<u>105][106]</u>
	Allergan, Inc. (Dublin, Ireland) <u>www.abbvie.it</u> , accessed on 14 February 2023	Zyplast	Glutaraldehyde	Integumental	[6][54][61][67][68] [69][70][73][74][76] [94][95][97][116]
		Zyderm	-	Integumental	[<u>6][61][67][68][71]</u> [<u>72][96][98][116</u>]
		Contigen	glutaraldehyde	Gastrointestinal and genitourinary	[<u>93][100][101][102]</u> [<u>103][104]</u>
Swine	GUNA (Milan, Italy) <u>www.guna.com</u> , accessed on 14 February 2023	Dental Skin BioRegulation	Vitamin C, magnesium gluconate, pyridozine chlorhydrate, riboflavin, thiamine chlorhydrate	Skin	[<u>117</u>]
		Dental ATM BioRegulation	Hypericum	Musculoskeletal	[<u>118</u>]
		MD-HIP	Calcium phosphate	Musculoskeletal	[<u>119</u>]
		MD-ISCHIAL	Rhododendron	Musculoskeletal	[<u>120</u>]
		MD-KNEE	Arnica	Musculoskeletal	[121][122][123]
		MD-LUMBAR	Hamemelis	Musculoskeletal	[120][124][125]

Source	Manufacturer	Product	Additives	Applications	Ref.
		MD-NECK	Silicio	Musculoskeletal	_
		MD- SHOULDERS	Iris	Musculoskeletal	[<u>126][127]</u>
		MD-SMALL JOINTS	Viola	Musculoskeletal	-
		MD- THORACIC	Cimifuga	Musculoskeletal	_
		MD-MATRIX	Citric acid, nicotinamide	Soft tissues	[125][128][129]
		MD-MUSCLE	Hypericum	Musculoskeletal	[<u>118][120][121][124]</u> [<u>125][127][128][129]</u> [<u>130]</u>
		MD-POLY	Drosera	Musculoskeletal	-
		MD-NEURAL	Citrullus	Musculoskeletal	[120][124][129]
		MD-TISSUE	Ascorbic acid, magnesium gluconate, pyridoxine chlorhydrate, riboflavin, thiamine chlorhydrate	Soft tissues	_
	Joint Biomaterials S.r.l. (Mestre, Italy) <u>www.joint-biomateriali.it</u> , accessed on 14 February 2023	CartiRegen	Fibrin glue	Musculoskeletal	_
	Ubiosis (Gyeonggi-do, Republic of Korea)	COLTRIX CartiRegen	-	Musculoskeletal	-
	www.ubiosis.com, accessed on 14 February 2023	COLTRIX TendoRegen	-	Musculoskeletal	-
		CartiFill	Glucose, CaCl, amino acids, vitamin B, fibrin glue	Musculoskeletal	[<u>131][132]</u> , NCT02539030, NCT02519881
	Sewon Cenontech Co., Ltd. (Seoul, Republic of Korea) www.swcell.com, accessed	CartiZol	Glucose, CaCl, amino acids, vitamin B	Musculoskeletal	[<u>133]</u> , NCT02539095
	on 14 r cordary 2023	RegenSeal	-	Musculoskeletal	[<u>134]</u>
		TheraFill	_	Integumental	[<u>64][65]</u>

Source	Manufacturer	Product	Additives	Applications	Ref.
	Sunmax Biotechnology Co., Ltd. (Tainan, Taiwan)	Facial Gain	Lidocaine	Integumental	NCT03844529
	www.sunmaxbiotech.com, accessed on 14 February 2023	Collagen Implant I	-	Integumental	-
	Evolence (Skillman, NJ, USA)	Dermicol-P35	Ribose	Integumental	[<u>2][135][136][137</u>] _, NCT00929071, NCT00891774
	Mentor Corp. (Santa Barbara, CA, USA)	Fibrel	_	Integumental	[<u>138][139]</u>
	Tissue Science Labs. (Aldershot, UK)	Permacol	_	Gastrointestinal	[<u>140][141][142][143]</u> [<u>144][145][146]</u>
	EternoGen, LLC (Columbia, MO, USA)	RPC Pure Collagen	Ethylenediamine tetraacetic acid	Integumental	[<u>147]</u>
	Aspid S.A. de C.V. (Mexico City, Mexico) www.aspidpharma.com, accessed on 14 February 2023	Fibroquel	Polyvinylpyrrolidone	Musculoskeletal	[<u>148][149]</u> NCT04517162
	ColBar LifeScience Ltd. (Tel Aviv, Israel) <u>www.ortho-</u> <u>dermatologics.com</u> , accessed on 14 February 2023	Evolence	Ribose	Integumental	[<u>135][150]</u>
Human	Fascia Biosystem (Beverly Hills, CA, USA)	Fascian	Lidocain	Integumental	[<u>6][151][152]</u>
	Fibrocell Science (Exton, PA, USA) www.fibrocell.com, accessed on 14 February 2023	Isolagen therapy	_	Integumental	NCT00655356
	Inamed Corporation (Santa Barbara, CA, USA) <u>www.inamed-cro.com,</u>	Cosmoplast	Glutaraldehyde, lidocaine hydrochloride	Integumental	[<u>6][153]</u>
	accessed on 14 February 2023	Cosmoderm	lidocaine hydrochloride	Integumental	[6][153]
	Life Cell Corp. (Branchburg, NJ, USA)	Dermalogen	Type and VI collagen, elastin, fibronectin,	Integumental	[<u>154]</u>
		0140141 11004			<i>.</i>

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	Source	Manufacturer	Product	Additives	Applications	Ref.	ast.
				chondroitin sulfate, and other proteoglycans			
		ons to Injectable Soft T	Cymetra	Elascin, glycosaminoglycans, Lidocaine hydrochloride	Integumental	[6][96][155][156] [157][158]	verse
	1000	Collagenesis, Inc.,	Autologen	Elastin, fibronectin, glycosaminoglycans	Integumental	_	12, 11,
1	0 Chenr	(Beveny, MA, USA)	Dermologen	-	Integumental	[<u>156</u>]	kin
1	Plant	Vesco Pharmaceutical Co. Ltd. (Bangkok, Thailand) <u>www.vescopharma.com,</u> accessed on 14 February 2023	Collagen C 1000	Vitamin C	Integumental	_	
1	Silkworm	Monodermà (Milan, Italy) www.monoderma.com	Fillagen	Hyaluronic acid, carboxymethylcellulose	Integumental	[<u>159]</u>	f the
1	n. d.	Taumed (Rome, Italy) <u>www.taumed.it</u> , accessed on 14 February 2023	Karisma	Hyaluronic acid, carboxymethylcellulose	Integumental	_	t 1, 9,
1	n. d.	LABO International S.r.l. (Padova, Italy) <u>www.labosuisse.com,</u> accessed on 14 February 2023	Fillerina con 3D collagen	Hyaluronic acid	Integumental	_	st.
1	n. d.	Hebey Mepha Pharm Group Co., Ltd. (Shandong, Hebei, China) <u>www.mephacn.com</u> , accessed on 14 February 2023	Collagen Plus	_	Integumental	_	. Surg.
1	n. d.	Pierre Mulot Laboratories (Paris, France)	Neutroskin	Vitamin C	Integumental	_	27. Ources
1	n. d.	Elements Pharmaceuticals (Shijiazhuang, Hebei, China) <u>www.elementspharma.com,</u> accessed on 14 February 2023	Ele-collagen	Vitamin C, Vitamin B6	Integumental	_	en as

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2	Source	Manufacturer	Product	Additives	Applications	Ref.	เทด
2	n. d.	Globus Medical (Audubon, PA, USA) <u>www.globusmedical.com,</u> accessed on 14 February 2023	Kinex Bioactive gel	Bioglass, hyaluronic acid	Musculoskeletal	_	

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- 20136549045e0169661014tors 4400961ations 44009610119112111231133116011611, sprained knee pain ^[122], injured cartilage 2^[1311134] 2^[131134] 2^[1311134] 2^[1311135] 2
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2010, 11, 407-426.

To improve the appearance of aged skin many non-invasive (i.e., topical formulations, oral supplements), minimally 49. Lynn, A.K.; Yannas, I.V.; Bonfield, W. Antigenicity and Immunogenicity of Collagen. J. Biomed. invasive (i.e., dermal fillers) and surgical treatments (i.e., blepharoplasty) were developed. Although a multitude of Mater. Res. B Appl. Biomater. 2004, 71, 343–354. topical treatments are available for the improvement of aged skin appearance, these procedures appeared to have

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injectables became, more popular for their immediate effect. As previously noted, several biomaterials (i.e., 51. Charriere, G.; Bejot, M.; Schnitzler, L.; Ville, G.; Hartmann, D.J. Reactions to a Bovine Collagen collagen, hyaluronic acid, calcium hydroxyl apatite, carboxy methyl cellulose, poly (methyl methacrylate), poly(L-Implant: Clinical and Immunologic Study in 705 Patients. J. Am. Acad. Dermatol. 1989, 21, 1203– lactic acid) were employed for the development of skin filler, each of which has some advantages and drawbacks. 1208.

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skin aging processes. Wrinkles formation is caused by collagen density decrease due to its turn-over slowing-down 53, Lemperle, G.; Morhenn, V.; Charrier, U. Human Histology and Persistence of Various Injectable l lts decreased synthesis and replacement rate causes matrix loss and thus skin collapse and loss of elasticity, Filler Substances for Soft Tissue Augmentation. Aesthetic Plast. Surg. 2003, 27, 354–366.

which in turn leads to the appearance of wrinkles, folds, and facial contour changes, as masterfully described by

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Eight Years of Clinical Experience in 153 Patients. Can. J. Plast. Surg. 2012, 20, 28–32. **4.2. Musculoskeletal Apparatus**

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The loss of muscle or osteochondral mass with advancing age is the major public health problem for the elderly 57. Cohen, S.R.; Holmes, R.E. Artecoll: A Long-Lasting Injectable Wrinkle Filler Material: Report of a population. Thus, musculoskeletal apparatus-related medical treatments and costs increase with population age Controlled, Randomized, Multicenter Clinical Trial of 251 Subjects. Plast. Reconstr. Surg. 2004, (numbers over 50 years). Among invasive and non-invasive currently available treatments, collagen injections are 114, 964–976. revealed to be quite effective for the treatment of several musculoskeletal diseases such as hip [119] or knee 58st Hanakas E012211/2036/2031/40/16211 aspylated Niceospherespheres in replaganil Sential 1234 tan-if Maris Surgo 2001424 ank and and and thin about arthritis [81] or fusion [78][84][85][86][87], lumbar spinal fusion [77], myofascial pain syndrome [118][130] chronic pain [120], acute lumbar spine, pain [124] and in partial thickness rotator cuff tears [123][134][162] plantar 59. Kim, K.J., Lee, H.W., Lee, M.W., Choi, J.H., Moon, K.C., Koh, J.K. Artecoll Granuloma: A Rare fasciitis [163] and calcific supraspinatus tendinitis [126] and pain [118][120][124][130] Adverse Reaction Induced by Microimplant in the Treatment of Neck Wrinkles. Dermatol. Surg. 2004, 30, 545–547. Osteoarthritis is an inflammatory degenerative disease characterized by the progressive damage of articular 6 Gar Radia and Brosofti Tipsuga Augumonatation defining rate on 11: And erson abe xperience. Fracial Riastr Success fact2994N29 α_1 hard 146 seem to be the main proinflammatory cytokines involved in the pathophysiology of osteparthritis, even though others, including IL-15, IL-18, IL-21, leukemia inhibitory factor (LIF) and chemokines of the Anzona Experience and Lessons Learned. Dermator. Surg.

are implicated [161][197] The expression of these inflammation mediators in turn activates the cartilage-degrading 2005, 31, 1566–1576. enzymes, that are matrix metalloproteinases (MMPs) and A disintegrin metalloproteinase with thrombospondin 6/2013910/2020 ABAM TS; 10/2020 Linak progressive Rivel Rivel Recie Soft Tissing Augmentation with Bellatill A several studies were performed to prove the Fyperience in 212 Patients. Plastic Surgn 2021 agen 98-102 beneficial to

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Rullan, P.P.; Thaler, M.P.; Ubogy, Z.; et al. ArteFill: A Long-Lasting Injectable Wrinkle Filler Material **4.3. Urogenital System** —Summary of the U.S. Food and Drug Administration Trials and a Progress Report on 4- to 5-

Year Outcomes. Plast. Reconstr. Surg. 2006, 118, 64–76.

6401Mgen,irSectionLeenaVeJ.beeniereJeWedSuchbe.S.m@hmDhY.inLeenvel.engKinnievelfektive, SoMtionTuror Specific uro Common anatiene Bitued sest state Besteven neisen and Gataety cof 1991 201 201 201 201 201 201 201 201 201 incollaise labil 197,01ch @greetetiosus 111 aistri Surger Hacter Serigie 201, 57,0197,1147, 1652 prostatectomy incontinence [99][102][173][174][175][176], retrograde ejaculation [177] and ovarian function after premature ovarian failure [192] 65. Lee, J.H.; Choi, Y.S.; Kim, S.M.; Kim, Y.J.; Rhie, J.W.; Jun, Y.J. Efficacy and Safety of Porcine Collagen Filler for Nasolabial Fold Correction in Asians: A Prospective Multicenter, 12 Months Stress urmary incontinence affects 10–30% of women above 50 years of age 100. To solve this common issue, in Follow-up Study. J. Korean Med. Sci, 2014, 29, S217–S221. addition to surgical practices (i.e., retropubic bladder neck suspension or slings), biomaterials injections (i.e., teflon, 676t, Hinchee, Collageno) haase, Dee Fiselfermed, Hollerdaeekursu Tralksteringur, Tundkaesserul uha Noleak Use of gathem, collEdgewalDlenQgelregenneedeeosanasincenhaiteed WhetnixoentegraaTiMy. Flowaastley/WoundriMateix)a.Coenbeneedre or impwitherRencwaanechuevea.imetevea.ime protetand use Rectioning the skylention Fronto Front of the protect of the second state of the second stat continent after 2 months [166]. However, because of collagen absorption, stress urinary incontinence recurrence 67. Kligman, A.M.; Armstrong, R.C. Histologic Respose to Intradermal Zyderm and Zyplast occurred in 41% of patients who achieved continence after 7–8 months [166]. ((Glutaraldehyde Cross-Linked) Collagen in Humans. J. Dermatol. Surg. Collagen réportedly J. Oncol. 1986, 12, degraded completely within 10–19 weeks, although magnetic resonance imaging of the urethra showed the persistence of 357. the implant for as long as 22 months after injection $\frac{175}{175}$. Thus, repeated injections (2–5) may be necessary $\frac{166}{167}$ 62000 Stepheane Ridcii Ghuwse Bensched Kwichrmazign gallen Orleightnande Electric Migrescondie Evaluation 506 60%Zynder mo Envlagen and rzynła strunulanten in Adinen Human Eraciał Skibish, Rilet Studye Arced Permatola veals 2013 143 weber. it should be mentioned that elderly patients should be counseled that approximately 40% will

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Glottic dysfunctions due to glottic gap, atrophy, paresis, bowing, paralysis and scarring result in voice absence or
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phonation. Recently, to reach a better postoperative voice in the long term, biomaterials injection (i.e., autologous

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non20027jv28e4c85availability, and possibility to be performed under local anesthesia. Among them, collagen

jpjectable formulations proved to be effective for vocal fold management. Patients treated with 1–2 mL of selected 75. Nevins, M.; Giannobile, W.V.; McGuire, M.K.; Kao, R.T., Mellonig, J.T.; Hinrichs, J.E.; McAllister, collagen injectable formulations. (Koken[®], AlloDerm[®], Zyplast[®]), showed at least some improvement in vocal B.S.; Murphy, K.S.; McClain, P.K.; Nevins, M.L.; et al. Platelet-Derived Growth Factor Stimulates function after the treatment, according to the Grade, Roughness, Breathiness, Asthenia, Strain (GRBAS) scale, Bone Fill and Rate of Attachment Level Gain: Results of a Large Multicenter Randomized

Maximum phonation time. Mean flow rate. Relative glottal area. In particular, perceptual and objective voice quality Controlled Trial. J. Periodontol. 2005, 76, 2205–2215. 7ionphaveimentAllessreseak, Rnakticetationalveatsidegletsiregi, the Carlingerase of Zive last innervolood on the Tate at the effort aro End 8 dt 11 998 1241 56 10 at 61 3 reduction of the mean flow rate from 322-564 mL/s to 223-385 mL/s and of the

_glottal gap [91][92][179], for at least up to 2 years after operation [92]. Thus, from the moment in which the safety and 77. Gadomski, B.C.; Labus, K.M.; Puttlitz, C.M.; McGilvray, K.C.; Regan, D.P.; Nelson, B.; Seim, H.B.; efficacy of collagen injections for the treatment of the vocal cords was affirmed by Ford and Bless in 1993 [181], the Easley, J.T. Evaluation of Lumbar Spinal Fusion Utilizing Recombinant Human Platelet Derived injection of heterologous material started to be even more required, given the positive feedback and long-term Growth Factor-B Chain Homodimer (RhPDGF-BB) Combined with a Bovine Collagen/βresults ^[96]. Although collagen injections were quite effective, and serious adverse events were rare ^{[91][92][95][181]}, Tricalcium Phosphate (β-TCP) Matrix in an Ovine Model. JOR Spine 2021, 4, e1166. documented complications included local abscess, migration of the implant, hypersensitivity reactions, stiffening, 78 si SFAtte Raten Mrasistardules Brigo brincha Magaatu Bone pybeu je the Rieter State of States in Reviewed property inje Greewithe Footporcial Hindroatten de la de la file stande la de la

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Phosphate (RhPDGF-BB/β-TCP)-Collagen Matrix as an Alternative to Autograft. Foot Ankle Int.

All popes of dillersonal origination of the filler material, Regardless of the filler material,

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To the best of our knowledge, based on harvested and available data on adverse reactions registered after 32. Abidi, N.A.; Younger, A.; Digiovanni, C.W. Role of Platelet-Derived Growth Factor in Hindfoot collagen-based commercial product applications, severe adverse events accounted for 8.2% (211 cases on 2587 Fusion. Tech. Foot Ankle Surg. 2012, 11, 34–38. patients), while mild adverse events accounted for about 5.3% (137 cases on 2587 patients) of those receiving the

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With a focus on collagen extraction sources, it emerged that severe adverse events accounted for 12.1% (211

cases on 1742 patients) and mild events for 3.8% (67 on 1742 patients) when bovine collagen was used. In 84. DiGiovanni, C.W.; Petricek, J.M. The Evolution of RhPDGF-BB in Musculoskeletal Repair and Its particular, severe adverse events were addressed to the use of one collagen-based product that was Augment[®], an Role in Foot and Ankle Fusion Surgery. Foot Ankle Clin. 2010, 15, 621–640. injectable formulation composed of bovine collagen, β-tricalcium phosphate and recombinant human platelet-

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Atelocell, Zyderm, Zyplast, Contigen, Gelofusine, Flowable wound matrix and Helitene) were not associated with 86. DiGiovanni, C.W.; Baumhauer, J.; Lin, S.S.; Berberian, W.S.; Flemister, A.S.; Enna, M.J.; such issues [66][91][92][94][96][106][106][166][167][169][194][203] (NCT02808325, NCT04637308, NCT02715466, Evangelista, P.; Newman, J. Prospective, Randomized, Multi-Center Feasibility Trial of RhPDGF-NCT01515397, NCT02631356, NCT00868062). Since bovine collagen appeared to be safe, these events could be BB versus Autologous Bone Graft in a Foot and Ankle Fusion Model. Foot Ankle Int. 2011, 32, ascribable to other Augment components, without certainty. As regards mild adverse reactions, they were registered only when using Augment, Chondrogrid or Zyderm [79][84][96][160]

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