

**UNITED STATES
SECURITIES AND EXCHANGE COMMISSION**
Washington, D.C. 20549

FORM 8-K

CURRENT REPORT
Pursuant to Section 13 or 15(d)
of The Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): August 9, 2023

TELA Bio, Inc.

(Exact name of registrant as specified in its charter)

Delaware
(State or other jurisdiction of incorporation)

001-39130
(Commission File Number)

45-5320061
(I.R.S. Employer Identification No.)

1 Great Valley Parkway, Suite 24
Malvern, Pennsylvania
(Address of principal executive offices)

19355
(Zip Code)

Registrant's telephone number, including area code: (484) 320-2930

Not Applicable
(Former name or former address, if changed since last report.)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
- Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
- Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
- Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock, par value \$0.001 per share	TEL A	Nasdaq Global Market

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (\$230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (\$240.12b-2 of this chapter).

Emerging growth company

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 2.02 Results of Operations and Financial Condition.

On August 9, 2023, TELA Bio, Inc. (the “**Company**”) issued a press release announcing its financial results for the second quarter ended June 30, 2023. A copy of this press release is furnished as Exhibit 99.1 hereto.

The information furnished pursuant to this Item 2.02, including Exhibit 99.1, shall not be deemed “filed” for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the “**Exchange Act**”), or otherwise subject to the liabilities of that section, and shall not be deemed to be incorporated by reference in any filing under the Securities Act of 1933, as amended (the “**Securities Act**”), or the Exchange Act, except as expressly set forth by specific reference in such filing.

Item 7.01 Regulation FD Disclosure.

On August 9, 2023, the Company updated information reflected in a corporate slide deck, which representatives of the Company will use in various meetings with investors from time to time. A copy of the presentation is attached hereto as Exhibit 99.2, and incorporated herein by reference.

The information furnished pursuant to Item 7.01, including Exhibit 99.2, shall not be deemed “filed” for purposes of Section 18 of the Exchange Act or otherwise subject to the liabilities of that section, and shall not be deemed to be incorporated by reference in any filing under the Securities Act or the Exchange Act, except as expressly set forth by specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits

The following exhibits are being furnished herewith:

Exhibit No.	Document
99.1	Press Release of TELA Bio, Inc., dated August 9, 2023.
99.2	Corporate Slide Deck, dated August 9, 2023.
104	Cover Page Interactive Data File (embedded within the Inline XBRL document).

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

TEL A BIO, INC.

By: /s/ Antony Koblisch

Name: Antony Koblisch

Title: President, Chief Executive Officer and Director

Date: August 9, 2023



TELABIO Reports Second Quarter 2023 Financial Results

MALVERN, PA, August 9, 2023 -- TELABIO, Inc. ("TELABIO"), a commercial-stage medical technology company focused on providing innovative soft-tissue reconstruction solutions that optimize clinical outcomes by prioritizing the preservation and restoration of the patient's own anatomy, today reported financial results for the second quarter ended June 30, 2023.

Recent Highlights

- Revenue of \$14.5 million in the second quarter, representing growth of 39% over the second quarter of 2022;
- Revenue of \$26.4 million for the first half of 2023, representing growth of 42% over the same period of 2022;
- Demand for OviTex® and OviTex PRS Reinforced Tissue Matrix increased in the second quarter of 2023, resulting in year-over-year revenue growth of approximately 43% and 31%, respectively;
- Results from a consumer survey highlighted strong consumer desire for shared-decision making and surgeon expertise in innovative, more natural hernia repair options;
- Underwritten public offering yielded net proceeds of approximately \$46.4 million; and
- Full year 2023 revenue guidance reaffirmed, with a range of \$60.0 million to \$65.0 million.

"We are pleased to report another quarter of sequential growth for TELABIO, driven by increased momentum in the sales organization and improving macroeconomic conditions for elective procedure volumes," said Antony Kobilish, co-founder, President and Chief Executive Officer of TELABIO. "In the first half of 2023, we grew our top line, increased market share, and launched innovative products and product enhancements to address the varying needs of surgeons and patients in an expanding soft tissue market. We look forward to leveraging our strong first-half performance into the third and fourth quarters and beyond."

Second Quarter 2023 Financial Results

Revenue was \$14.5 million in the second quarter of 2023, an increase of 39% compared to the same period in 2022. The increase was due to the expansion of our commercial organization, the addition of new customers, increased penetration within existing customer accounts, and growing international sales.

Gross profit was \$10.2 million in the second quarter of 2023, or 70% of revenue, compared to \$6.6 million, or 63% of revenue, in the same period in 2022. The increase in gross margin was primarily due to better inventory management practices resulting in a decrease in the reserve for excess and obsolete inventory as a percentage of revenue as compared to the prior year period.

Operating expenses were \$20.6 million in the second quarter of 2023, compared to \$16.8 million in the same period in 2022. The increase was due to higher compensation and employee-related expenses from additional headcount as we continue to expand our organization, along with increased travel expenses, increased consulting fees and higher study costs.

Loss from operations was \$10.4 million in the second quarter of 2023, compared to a loss from operations of \$10.2 million in the same period in 2022.

Net loss was \$10.8 million in the second quarter of 2023, compared to a net loss of \$12.7 million in the same period in 2022.

Cash and cash equivalents on June 30, 2023 totaled \$65.3 million following a public offering which raised approximately \$46.4 million dollars after commissions and expenses.

2023 Financial Guidance

We continue to expect full year 2023 revenue to range from \$60.0 million to \$65.0 million, reflecting growth of 45% to 57% over full year 2022.

Conference Call

TEL A Bio will host a conference call at 4:30 p.m. Eastern Time on Wednesday, August 9, 2023 to discuss its second quarter 2023 financial results. Investors interested in listening to the conference call should [register online](#). Participants are required to register a day in advance or at minimum 15 minutes before the start of the call. A replay of the webcast can be accessed via the [Events & Presentations](#) page of the investor section of TEL A Bio's website.

About TEL A Bio, Inc.

TEL A Bio, Inc. (NASDAQ: TELA) is a commercial-stage medical technology company focused on providing innovative technologies that optimize clinical outcomes by prioritizing the preservation and restoration of the patient's own anatomy. The Company is committed to providing surgeons with advanced, economically effective soft-tissue reconstruction solutions that leverage the patient's natural healing response while minimizing long-term exposure to permanent synthetic materials. For more information, visit www.telabio.com.

Caution Regarding Forward-Looking Statements

This press release contains forward-looking statements within the meaning of The Private Securities Litigation Reform Act of 1995. Words such as "may," "might," "will," "should," "believe," "expect," "anticipate," "estimate," "continue," "predict," "forecast," "project," "plan," "intend" or similar expressions, or statements regarding intent, belief, or current expectations are forward-looking statements and reflect the current beliefs of TELA Bio's management. Such forward-looking statements include statements relating to our expected revenue and revenue growth for the full year 2023. These statements are not guarantees of future performance and are subject to certain risks, uncertainties and other factors that could cause actual results and events to differ materially and adversely from those indicated by such forward-looking statements including, among others: the impact to our business from macroeconomic conditions, including the COVID-19 pandemic and other public health crises, recessionary concerns, banking instability, increasing market interest rates, and inflationary pressures, potentially impacting our ability to market our products, demand for our products due to the deferral of elective procedures, the labor and staffing environment in the healthcare industry, disruption in our supply chain, or pricing pressures concerning our products or the procedures using our products; our ability to achieve or sustain profitability; our ability to gain market acceptance for our products and to accurately forecast and meet customer demand; our ability to compete successfully; that data from earlier studies related to our products and interim data from ongoing studies may not be replicated in later studies or indicative of future data; that data obtained from clinical studies using our product may not be indicative of outcomes in other surgical settings; our ability to enhance our product offerings; development and manufacturing problems; capacity constraints or delays in production of our products; maintenance of coverage and adequate reimbursement for procedures using our products; product defects or failures. These risks and uncertainties are described more fully in the "Risk Factors" section and elsewhere in our filings with the Securities and Exchange Commission and available at www.sec.gov, including in our Annual Report on Form 10-K and Quarterly Reports on Form 10-Q. Any forward-looking statements that we make in this announcement speak only as of the date of this press release, and TELA Bio assumes no obligation to update forward-looking statements whether as a result of new information, future events or otherwise after the date of this press release, except as required under applicable law.

Investor Contact

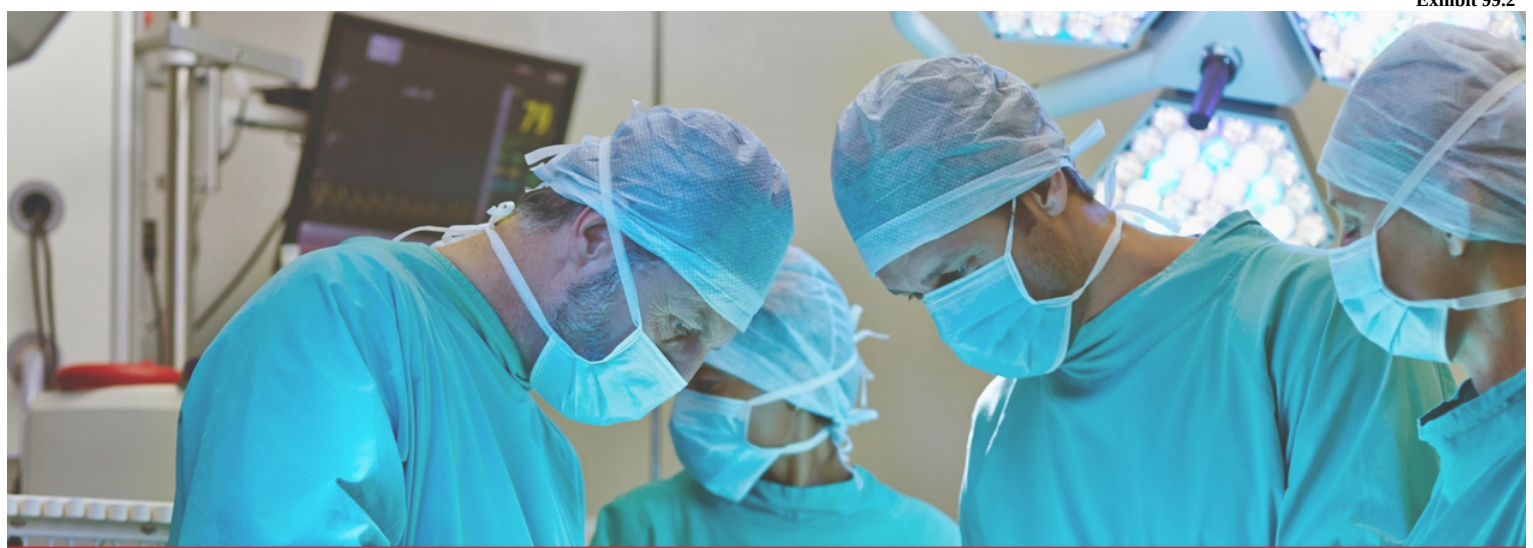
Greg Chodaczek
332-895-3230
ir@telabio.com

TEL A Bio, Inc.
Consolidated Balance Sheets
(In thousands, except share and per share amounts)
(Unaudited)

	June 30, 2023	December 31, 2022
Assets		
Current assets:		
Cash and cash equivalents	\$ 65,266	\$ 42,019
Accounts receivable, net	7,894	6,621
Inventory	14,098	11,792
Prepaid expenses and other assets	1,910	2,015
Total current assets	<u>89,168</u>	<u>62,447</u>
Property and equipment, net	1,764	1,682
Intangible assets, net	2,309	2,499
Right-of-use assets	1,145	1,227
Total assets	<u>\$ 94,386</u>	<u>\$ 67,855</u>
Liabilities and stockholders' equity		
Current liabilities:		
Accounts payable	\$ 2,250	\$ 1,534
Accrued expenses and other current liabilities	10,795	10,869
Total current liabilities	<u>13,045</u>	<u>12,403</u>
Long-term debt	40,212	39,916
Other long-term liabilities	1,124	1,231
Total liabilities	<u>54,381</u>	<u>53,550</u>
Stockholders' equity:		
Preferred stock; \$0.001 par value; 10,000,000 shares authorized; no shares issued and outstanding	—	—
Common stock; \$0.001 par value; 200,000,000 shares authorized; 24,475,504 and 19,165,027 shares issued and outstanding at June 30, 2023 and December 31, 2022, respectively	24	19
Additional paid-in capital	336,939	288,361
Accumulated other comprehensive income	84	150
Accumulated deficit	(297,042)	(274,225)
Total stockholders' equity	<u>40,005</u>	<u>14,305</u>
Total liabilities and stockholders' equity	<u>\$ 94,386</u>	<u>\$ 67,855</u>

TEL A Bio, Inc.
Consolidated Statements of Operations and Comprehensive Loss
(In thousands, except share and per share amounts)
(Unaudited)

	Three months ended June 30,		Six months ended June 30,	
	2023	2022	2023	2022
Revenue	\$ 14,494	\$ 10,406	\$ 26,403	\$ 18,637
Cost of revenue (excluding amortization of intangible assets)	4,198	3,318	8,114	6,474
Amortization of intangible assets	95	538	190	614
Gross profit	<u>10,201</u>	<u>6,550</u>	<u>18,099</u>	<u>11,549</u>
Operating expenses:				
Sales and marketing	14,577	11,055	28,043	20,433
General and administrative	3,472	3,630	7,106	7,088
Research and development	2,514	2,102	4,566	4,109
Total operating expenses	<u>20,563</u>	<u>16,787</u>	<u>39,715</u>	<u>31,630</u>
Loss from operations	(10,362)	(10,237)	(21,616)	(20,081)
Other expense:				
Interest expense	(1,298)	(934)	(2,544)	(1,845)
Loss on extinguishment of debt	—	(1,228)	—	(1,228)
Other income (expense)	870	(342)	1,343	(449)
Total other expense	<u>(428)</u>	<u>(2,504)</u>	<u>(1,201)</u>	<u>(3,522)</u>
Net loss	\$ (10,790)	\$ (12,741)	\$ (22,817)	\$ (23,603)
Net loss per common share, basic and diluted	\$ (0.46)	\$ (0.88)	\$ (1.08)	\$ (1.62)
Weighted average common shares outstanding, basic and diluted	<u>23,239,262</u>	<u>14,557,453</u>	<u>21,223,639</u>	<u>14,548,210</u>
Comprehensive loss:				
Net loss	\$ (10,790)	\$ (12,741)	\$ (22,817)	\$ (23,603)
Foreign currency translation adjustment	(36)	134	(66)	181
Comprehensive loss	<u>\$ (10,826)</u>	<u>\$ (12,607)</u>	<u>\$ (22,883)</u>	<u>\$ (23,422)</u>



A Soft-Tissue Preservation and Restoration Company

INVESTOR PRESEN

Forward Looking Statements

This presentation contains forward-looking statements within the meaning of The Private Securities Litigation Reform Act of 1995. All statements other than historical facts contained in this document, including but not limited to statements regarding possible or assumed future results of operations, business development plans, regulatory activities, market opportunity competitive position, potential growth opportunities, and the effects of competition, are forward-looking statements. These statements involve known and unknown risks, uncertainties and other important factors that may cause TELA Bio, Inc.'s (the "Company") results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. In some cases, you can identify forward-looking statements by terms such as "may," "will," "should," "expect," "plan," "aim," "anticipate," "could," "target," "project," "contemplate," "believe," "estimate," "predict," "potential" or "continue" or the negative of these terms or other similar expressions. The statements in this presentation are only predictions. The Company has based these forward-looking statements largely on its current expectations and projections about future events and financial trends that it believes may affect the Company's business, financial condition and results of operations. These forward-looking statements speak only as of the date of this presentation and are subject to a number of risks, uncertainties and assumptions, some of which cannot be quantified and some of which are beyond the Company's control, including, among others: the impact to the Company's business from macroeconomic conditions including the COVID-19 pandemic or other public health crises, recessionary concerns, banking instability, increasing market interest rates, and inflation potentially impacting our ability to market our products, demand for our products due to the deferral of elective procedures, the labor and staffing environment in the healthcare industry, disruption in our supply chain, or pricing pressures concerning our products or the procedures using our products; our ability to achieve profitability; the Company's ability to gain market acceptance for the Company's products and to accurately forecast and meet customer demand; the Company's ability to compete successfully; that data from earlier studies related to the Company's products and interim data from ongoing studies may not be replicated in future studies and may not be indicative of future data, that data obtained from clinical studies utilizing the Company's products may not be indicative of outcomes in other surgical settings; the Company's ability to enhance the Company's product offerings; development and manufacturing problems; capacity constraints or delays in production of the Company's products; maintenance of coverage and adequate reimbursement for procedures using the Company's products; product defects or failures. Other risks and uncertainties are described more fully in the "Risk Factors" section and elsewhere in the Company's filings with the Securities and Exchange Commission and available at www.sec.gov. You should not rely on these forward-looking statements as predictions of future events. The events and circumstances reflected in the Company's forward-looking statements may not be achieved or occur, and actual results could differ materially from those projected in the forward-looking statements. Moreover, the Company operates in a dynamic industry and economy. New risk factors and uncertainties may emerge from time to time that are not possible for management to predict all risk factors and uncertainties that the Company may face. Except as required by applicable law, we do not plan to update or revise any forward-looking statements contained herein, whether as a result of any new information, future events, changed circumstances or otherwise.

TEL A Bio, Inc.

- Advanced reinforced tissue matrix portfolio supported by compelling clinical evidence
- \$2.2B US market opportunity¹ – still in early stages of growth
- Driving commercial adoption with targeted direct-sales approach
- Recent product launches in growing markets: robotic hernia surgery, plastic and reconstructive surgery
- Broad intellectual property portfolio
- Established DRG-based reimbursement pathway for hernia repair and robust GPO access
- Highly accomplished executive team with proven track record

*Redefining soft tissue pre-
restoration with a differentiat-
tissue reinforcement m-
and supportive prod-*

OVITE
REINFORCED TISSUE M

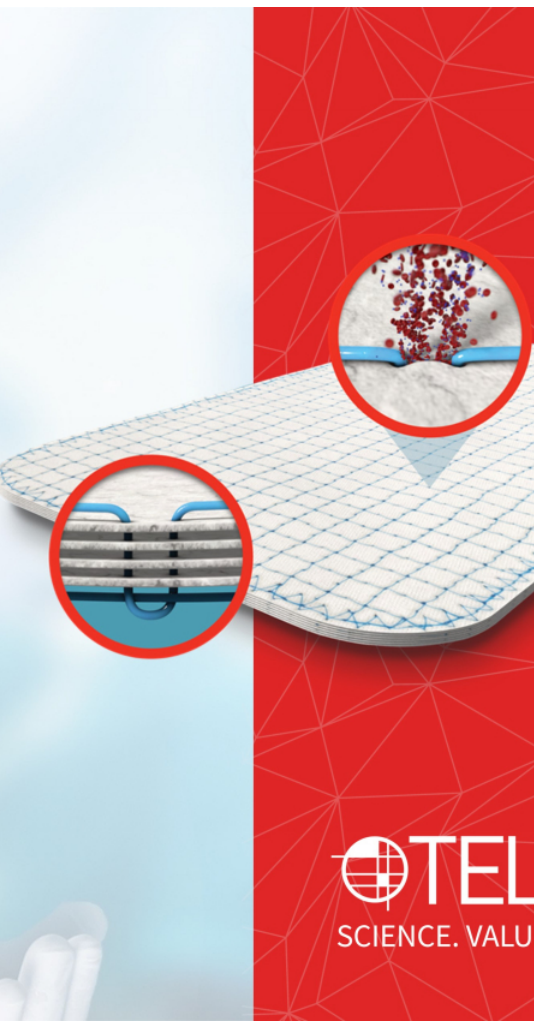
OVITEX®
REINFORCED TISSUE M

¹ Management estimate. \$2.2B total includes \$1.5B hernia & abdom-
plastic reconstructive surgery.



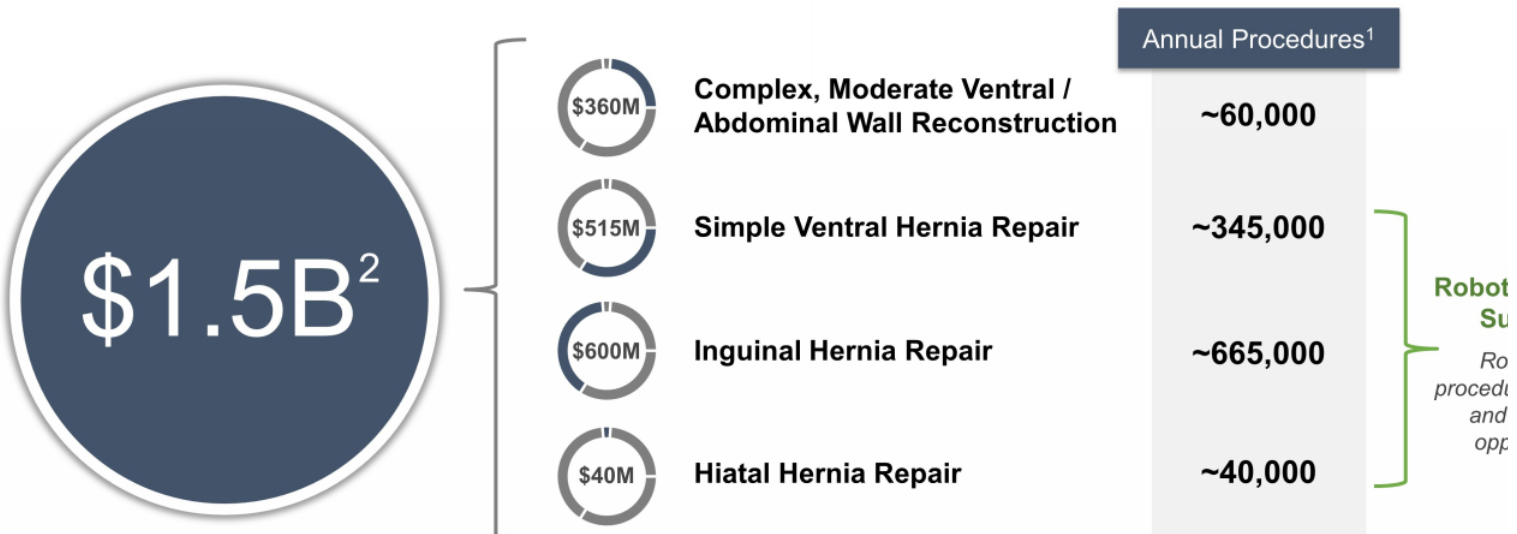
OVITEX®

REINFORCED TISSUE MATRIX



OTEL
SCIENCE. VALUE.

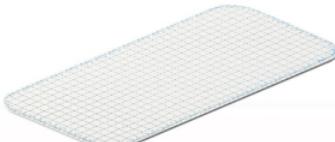
U.S. Hernia Surgery Market: ~\$1.5 Billion Annual Opportunity



¹Sources: Millennium Research Group Reports, IMS Health Data; iData Research MedSKU

²Management estimate. Market size based on volume and weighted average selling price for OviTex

OviTex Portfolio: Designed for a Range of Hernia Patients and Surgical Techniques



CONFIGURATION

OviTex LPR

4-layer device, with "smooth side" suitable for intraperitoneal placement

Robot Compatible¹: Yes

Strength²: +

Viscera Contact³: Yes, smooth side

OviTex

4-layer device, not intended for intraperitoneal placement

Robot Compatible¹: Yes

Strength²: +

Viscera Contact³: Not recommended

OviTex 1S

6-layer device, with "smooth side" suitable for intraperitoneal placement

Robot Compatible¹: Yes

Strength²: ++

Viscera Contact³: Yes, smooth side

OviTex 2S

8-layer device, not intended for intraperitoneal placement

Robot Compatible¹: Yes

Strength²: +++

Viscera Contact³: Not recommended

COMPETITIVE SET

- Coated resorbable synthetic meshes



- Resorbable synthetic meshes



- Biologic meshes



- Biologic meshes



- Coated resorbable synthetic meshes



- Biologic meshes



- Biologic meshes



Images represent permanent polymer OviTex products. Resorbable polymer products have clear polymer. All trademarks and registered marks are property of their respective owners.

1. Robot compatibility based on use of 10mm trocar. Robot compatibility of LPR and OviTex include sizes 400 cm² or less. Robot compatibility of OviTex 1S includes sizes 200 cm² or less

2. Biomechanical data on file.

3. Devices with a smooth side were shown to not adhere in an animal model. Rabbit data on file. Correlation to results in humans has not been established. Animal test results may not necessarily be indicative of human clinical performance.

Need for Alternative to Permanent Synthetic Mesh

59%

of surgeons agree that use of permanent synthetic mesh puts patients at long-term risk of complications¹

3 of 4

Hernia patients want proactive control in their care²

~24K

Lawsuits against permanent synthetic meshes estimated to be assembled across the U.S.³

1. Hernia and Abdominal Surgeries Survey (Oct 2020). A group of 71 surgeons were surveyed regarding use of mesh in various hernia repair surgeries.

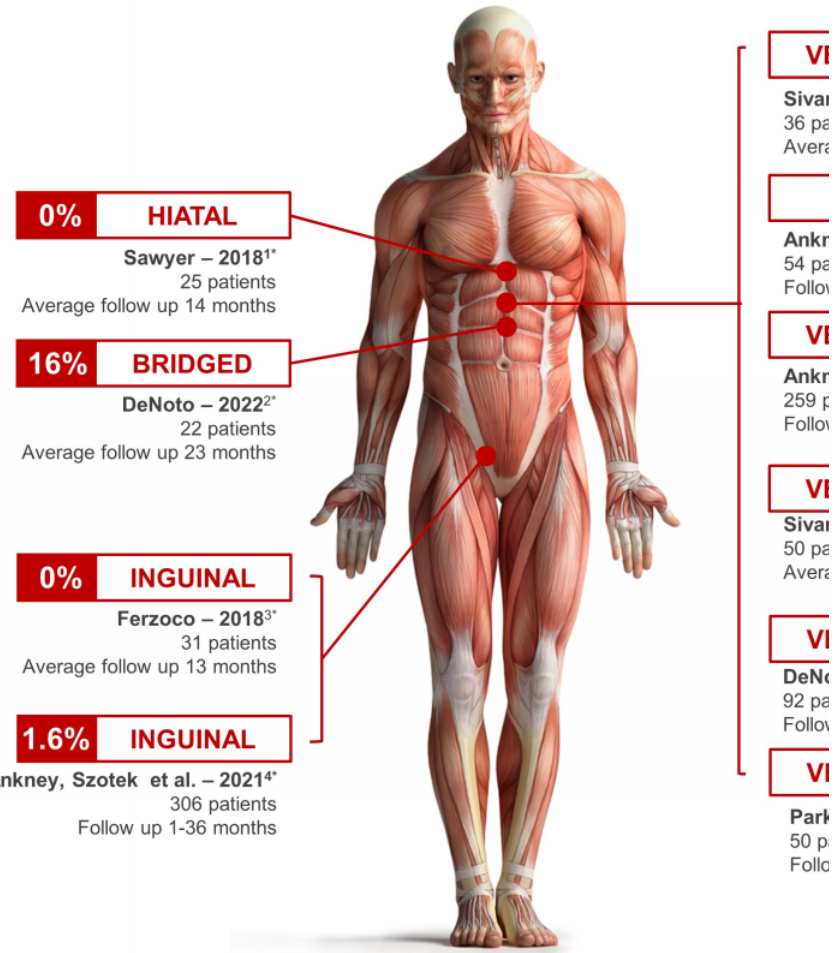
2. Figures derived from Company-sponsored poll of approximately 1,100 potential patients for hernia procedures.

3. www.drugwatch.com (September 2022)

LOW RECURRENCE ALL APPLICATIONS WITH OVITEX



Source: Refer to "Ovitex Clinical References" in this presentation.
 * Indicates one or more surgeons are paid consultants of Tela Bio, Inc.



Favorable Results of OviTex in Ventral Hernia Repair Comparisons to synthetic mesh and leading generation one biologics

	Parker et al. ⁸		Sivaraj et al. ⁵		
Total enrolled patients	50 OviTex	50 Polypropylene	36 OviTex	51 Strattec	17 Permacol
Length of follow-up	12 months	12 months	28.6 months (median)	34.6 months (median)	58.4 months (median)
mVHWG	32% grade 2 68% grade 3 ^a	94% grade 2 6% grade 3	33% grade 1 58% grade 2 8% grade 3	17% grade 1 79% grade 2 4% grade 3	18% grade 1 71% grade 2 12% grade 3
CDC wound class	70% CDC class II+ ^a	94% CDC class I	89% class I-II	86% class I-II	94% class I-II
Incidence of SSO	36%*	22%*	16.7%*	47.1%*	52.9%*
Incidence of SSI	-	-	2.8% ^b	12.5%	11.8%
Recurrence rate	6%	12%	2.8% ^c	13.7% ^c	29.4%

*Overall complications including SSI

a – OviTex patients were more complicated with a significantly higher mVHWG distribution and CDC wound classification compared to polypropylene patients

b – OviTex patients experienced significantly less complications than patients receiving the other three biologics

c - OviTex and Strattec patients had a statistically lower recurrence rate than patients receiving the other two biologics

Source: Refer to "Clinical References" in this presentation.

Positive 24-month BRAVO results in ventral hernia OviTex performance contextualized alongside recent publication leading competitive products

	DeNoto et al. (BRAVO) ⁷	Harris et al. (PRICE) ¹⁰		Roth et al. ¹¹
Total enrolled patients	92 OviTex	82 Stratite	83 Ventralight ST or Bard Soft Mesh	121 Phaxis
Length of follow-up	24 months	26 months		36 months
mVHWG	78% grade 2-3	-	-	-
CDC wound class	95% class I-II	90% class I-II	93% class I-II	100% class I
Surgical technique	Open (65%) Laparoscopic (13%) Robotic (22%)	Open	Open	Open
Incidence of SSO	38% (includes SSI)	21% (excludes SSI)	22% (excludes SSI)	-
Incidence of SSI	20.7%	39%	34%	9%*
Recurrence rate	2.6%*	40% (overall) 34% (class I wounds)	22% (overall) 28% (class I wounds)	17.9%*

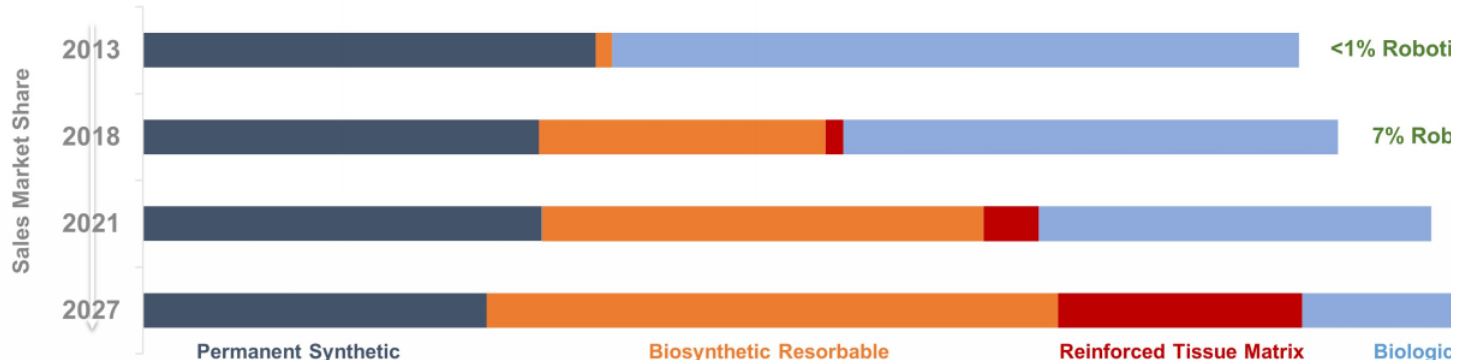
* Kaplan-Meier survival estimate

**No head-to-head clinical studies have been conducted. Due to differences in patient population, surgeons, surgical technique, and other variables, no direct comparisons of results can be made. For a comparative discussion of these studies, please see G. DeNoto, E.P. Ceppa, S.J. Pacella, M. Sawyer, G. Slayden, M. Takata, G. Tuma, J. Yunis, 24-Month results of the BRAVO study: A prospective, multi-center study evaluating the clinical outcomes of a ventral hernia cohort treated with OviTex® 1S permanent reinforced tissue matrix, Ann Medicine Surg 2022, 83, 104745.

Source: Refer to "Clinical References" in this presentation.

Hernia Market Evolution

TELA Bio is gaining from a market shift by providing our reinforced “natural repair” solutions as an alternative to traditional Permanent Synthetics or Biologics



***Biosynthetic Resorbable* and *Reinforced Tissue Matrix* strengths:**



Clinical Evidence



Robot Compatibility



Cost-effectiveness



Patient Choice & Shared Decision-making

Sources for Sales Market Share (%): 2009 - 2013 = IMS Hospital Supply Index; 2018 - 2021 = iData Research MedSKU; 2027 = Management Estimate

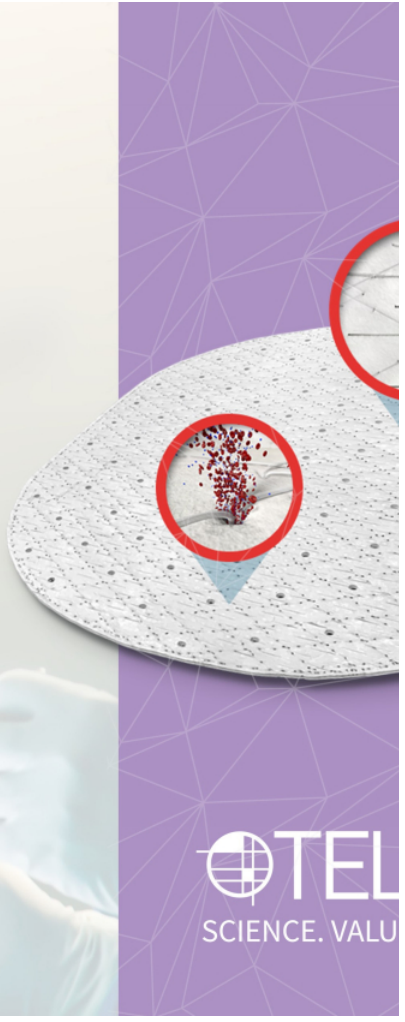
Sources for Total US Market Size: 2021 - 2027 = DRG Hernia Repair Devices Report – 2021; 2013 - 2018 = Management Estimate

Sources for % Robotic Procedures (Px): 2018 - 2027 = DRG Hernia Repair Devices Report – 2021; 2013 = Management Estimate



OVITEX® PRS

REINFORCED TISSUE MATRIX



OATEL
SCIENCE. VALUE.

U.S. Plastic and Reconstructive Surgery Market: ~\$700 Million Annual Opportunity

\$600M²

Surgeons use products to reinforce soft tissue during various reconstructive surgeries¹, including:

- Head and neck surgery
- Chest wall reconstruction
- Pelvic reconstruction
- Extremities reconstruction
- Breast reconstruction

Market dominated by human acellular dermal matrices (HADMs):

- Prone to high degree of stretch
- Expensive, putting pressure on hospital systems
- Can experience supply shortages, particularly when large pieces of material are required

¹OviTex PRS is for implantation to reinforce soft tissue where weakness exists in patients requiring soft tissue repair or reinforcement in plastic and reconstructive surgery. The device is supplied sterile and is intended for one-time use. OviTex PRS has not been tested in breast surgical procedures.

²Management estimate. Source: iData Research MedSKU, Q3 2021. Market size based on sales of current biologics

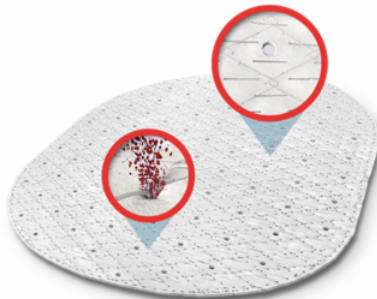
Cosme
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\$1

OviTex PRS: Specifically Designed for Plastic and Reconstructive Surgery

Available in both **2-layer resorbable (polyglycolic acid) polymer, 3-layer permanent (polypropylene) polymer, or 3-layer resorbable (polylactic-co-glycolic acid) polymer** reinforcing the same biologic material



An innovative reinforced tissue matrix designed to improve outcomes by facilitating fluid management and controlling degree and direction of soft tissue expansion.

Product Features:

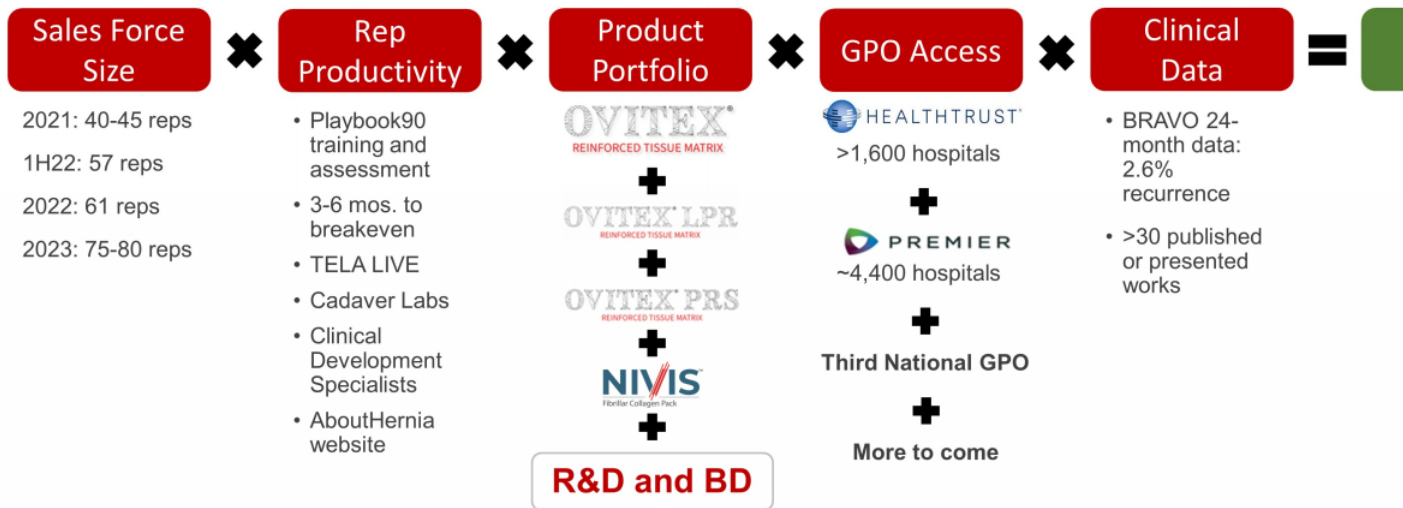
- Layers composed of biologic building block retain significant macromolecules for tissue regeneration
- Diamond embroidery pattern and stents allow for flexibility or sawtooth embroidery pattern to accommodate directional stretch while providing stretch resistance
- Distinct permeability elements in various configurations e.g., micropores, macropores, and stents – designed to facilitate fluid management

OviTex PRS compared to market leading products

- Exhibited earlier host cell proliferation, collagen deposition, and neovascularization
- Demonstrated tissue remodeling into mature, functional organized collagen

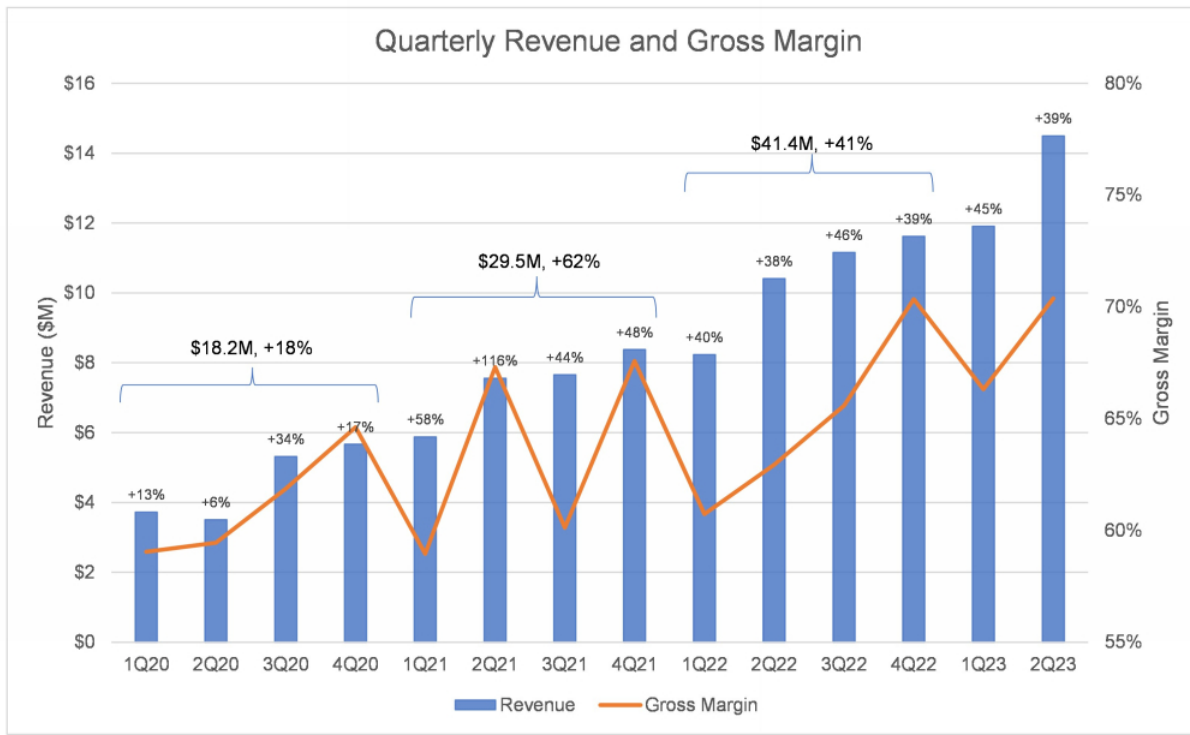
1. Certain configurations available in two or three layers, see product catalog for more information. 2. Lun S, Irvine S.M., Johnson K.D., Fisher N.J., Floden E. W., Negron L., Dempsey S.G., McLaughlin R.J., Vasudevanamurthy M., Ward B.R., May B.C., A functional extracellular matrix biomaterial derived from ovine forestomach, *Biomaterials* 31(16) (2010) 4517-29.
3. ADM: Acellular Dermal Matrix. Overbeck N, Beierschmitt A, May BC, Qi S, Koch J. In-Vivo Evaluation of a Reinforced Ovine Biologic for Plastic and Reconstructive Procedures in a Non-human Primate Model of Soft Tissue Repair. *Elasty*. 2022 Sep 14;22:e43. PMID: 36160663; PMCID: PMC9490877. Animal testing results may not be indicative of clinical performance.

Driving Revenue Growth



TELA Bio is growing each factor that contributes to sales, providing multi-year, long-term growth

Delivering Revenue Growth and Margin Improvement



Q2 2023 Performance

- Revenue of \$14.5M, +39% over corresponding period in 2022
- 70% Gross Margin
- Cash and Cash Equivalents as of June 30, 2023: \$1.2B

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