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): January 13, 2020

EyePoint Pharmaceuticals, Inc.

(Exact name of registrant as specified in its charter)

Delaware (State or Other Jurisdiction of Incorporation) 000-51122 (Commission File Number) 26-2774444 (IRS Employer Identification No.)

480 Pleasant Street

Watertown, MA 02472 (Address of Principal Executive Offices, and Zip Code)

(617) 926-5000 Registrant's Telephone Number, Including Area Code

(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 communication 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)

D Pre-commencement communication pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

Pre-commencement communication 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:

	Trading	Name of each exchange
Title of each class	Symbol(s)	on which registered
Common Stock, par value \$0.001	EYPT	The Nasdaq Stock Market LLC

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 \Box

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 8.01. Other Events.

On January 13, 2020, EyePoint Pharmaceuticals, Inc. posted an updated corporate presentation on its website at www.eyepointpharma.com. A copy of the presentation is filed herewith as Exhibit 99.1 and is incorporated by reference herein.

Item 9.01. Financial Statements and Exhibits.

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.

EYEPOINT PHARMACEUTICALS, INC.

Date: January 13, 2020

/s/ Nancy Lurker Nancy Lurker President and Chief Executive Officer



INVESTOR PRESENTATION

JANUARY 2020

NASDAQ: EYPT



FORWARD LOOKING



SAFE HARBOR STATEMENTS UNDER THE PRIVATE SECURITIES LITIGATION ACT OF 1995: Various statements made in this release are forward-looking, and are inherently subject to risks, uncertainties and potentially inaccurate assumptions. This presentation is intended for communication for investors only. Nothing in this presentation should be construed as promoting the use of YUTIQ®, DEXYCU® or other product candidates. All statements that address activities, events or developments that we intend, expect, plan or believe may occur in the future, including but not limited to statements about our expected financial results for the fourth quarter and full fiscal year ended December 31, 2019 and longer-term financial and business goals, are forward-looking statements. Our preliminary fourth quarter and full year 2019 revenue results are preliminary and subject to adjustment in the ongoing review procedures by our independent registered public accounting firm. In addition, any financial projections and other estimates contained herein are forward-looking statements with respect to the anticipated performance of the Company. Such financial projections and estimates are as to future events and are not to be viewed as facts, and reflect various assumptions of management of the Company and are subject to significant business, financial, economic, operating, competitive and other risks and uncertainties and contingencies (many of which are difficult to predict and beyond the control of the Company) that could cause actual results to differ materially from the statements included herein. Some of the factors that could cause actual results to differ materially from the anticipated results or other expectations expressed, anticipated or implied in our forward-looking statements are risks and uncertainties inherent in our business including, without limitation: our ability to achieve profitable operations and access to needed capital; fluctuations in our operating results; our ability to successfully produce sufficient commercial quantities of YUTIO and DEXYCU and to successfully commercialize YUTIQ and DEXYCU in the U.S.; our ability to sustain and enhance an effective commercial infrastructure and enter into and and better of the use of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affecting the posterior segment of the every potential off-label sales of ILUVIEN for non-infectious uveits affe consequences of fluocinolone acetonide side effects for YUTIQ; consequences of dexamethasone side effects for DEXYCU; successful commercialization of, and receipt of revenues from, ILUVIEN for diabetic macular edema, or DME; Alimera's ability to obtain additional marketing approvals and the effect of pricing and reimbursement decisions on sales of ILUVIEN for DME; Alimera's ability to commercialize ILUVIEN for non-infectious uveits affecting the posterior segment of the eye in the territories in which Alimera is licensed to do so; declines in Retisert royalties; our ability to market and sell products; the success of current and future license agreements; termination or breach of current license agreements; our dependence on contract research organizations, contract sales organizations, vendors and investigators; effects of competition and other developments affecting sales of products; market acceptance of products; effects of guidelines, recommendations and studies; protection of intellectual property and avoiding intellectual property infringement; retention of key personnel; product liability; industry consolidation; compliance with environmental laws; manufacturing risks; risks and costs of international business operations; volatility of stock price; possible dilution; absence of dividends; and other factors described in our filings with the Securities and Exchange Commission. We cannot guarantee that the results and other expectations expressed, anticipated or implied in any forward-looking statement will be realized. A variety of factors, including these risks, could cause our actual results and other expectations to differ materially from the anticipated results or other expectations expressed, anticipated or implied in our forward-looking statements. Should known or unknown risks materialize, or should underlying assumptions prove inaccurate, actual results could differ materially from past results and those anticipated, estimated or projected in the forward-looking statements. You should bear this in mind as you consider any forward-looking statements. Our forward-looking statements speak only as of the dates on which they are made. We do not undertake any obligation to publicly update or revise our forward-looking statements even if experience or future changes makes it clear that any projected results expressed or implied in such statements will not be realized.

AN EMERGING LEADER IN OPHTHALMOLOGY





Chronic non-infectious uveitis affecting the posterior segment of the eye

 Transitioned into commercial growth company with the launches of YUTIQ® and DEXYCU® in Q1 2019

 Pipeline focused on innovative therapies improving standard of care in areas of unmet need, including wet AMD

 Focused on organic and inorganic product growth leveraging proprietary technologies and select product acquisition to build top-line revenue and expand pipeline

 Strategic focus to become go-to partner for commercialization in ophthalmology

Veteran executive team with deep experience in commercial product launches and clinical development

PRODUCT PIPELINE



Program	Preclin.	Phase 1	Phase 2	Phase 3	Commercial	Rights
DEXYCU [®] post-operative inflammation	-			Launo	hed/J-Code In-Place	ww
YUTIQ [®] three-year treatment for chronic non-infectious uveitis affecting the posterior segment				Laund	hed/J-Code In-Place	U.S. ⁽¹⁾
YUTIQ° 50 (sNDA) 6-month treatment for chronic non-infectious uveitis affecting the posterior segment	~45 patie	nt 6-month study pl	anned			ww
EYP-1901 - TKI Durasert [™] 6-month wet AMD treatment	IND enabling studies					ww

PROPRIETARY OCULAR DELIVERY TECHNOLOGIES



Durasert[™] and Verisome[™]: Proven, FDA Approved Technologies

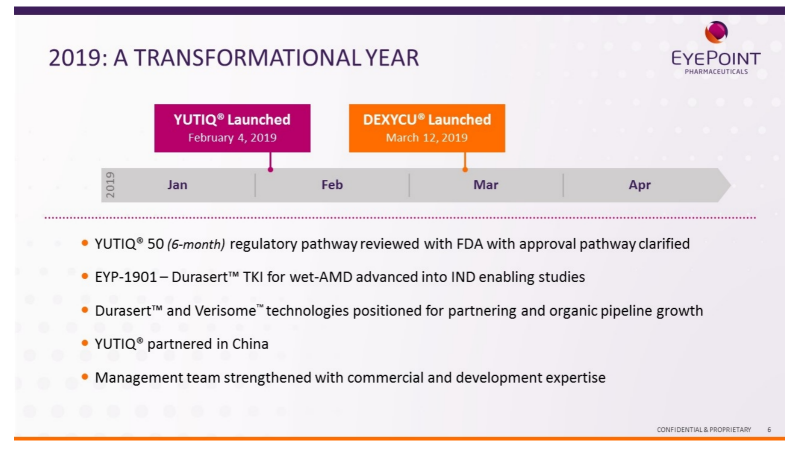
• RETISERT ® (2005, B&L) - Uveitis

• VITRASERT® (1996, B&L) -CMV retinitis

Durasert™	Verisome™
Ocular insert for long-term delivery of small molecules	Microsphere suspension short-term delivery of small and potentially large molecules
Approved products:	Approved products:
 YUTIQ[®] (2018, EyePoint) 	 DEXYCU[®] (2019, EyePoint)
 ILUVIEN[®] (2012, Alimera) - DME 	

CONFIDENTIAL & PROPRIETARY

5



TWO COMMERCIAL PRODUCTS LAUNCHED IN 2019

-





Postoperative inflammation following ocular surgery

- 33 dedicated KAMs targeting high-volume ambulatory surgical centers (ASCs)
- Permanent and specific J-Code with minimal reimbursement issues



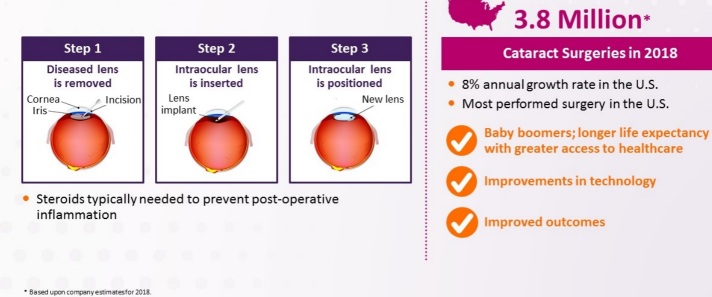
Chronic non-infectious uveitis affecting the posterior segment of the eye

- 12 dedicated KAMs targeting uveitis specialists – increasing to 18 in 2020
- Permanent and specific J-Code effective as of October 1, 2019

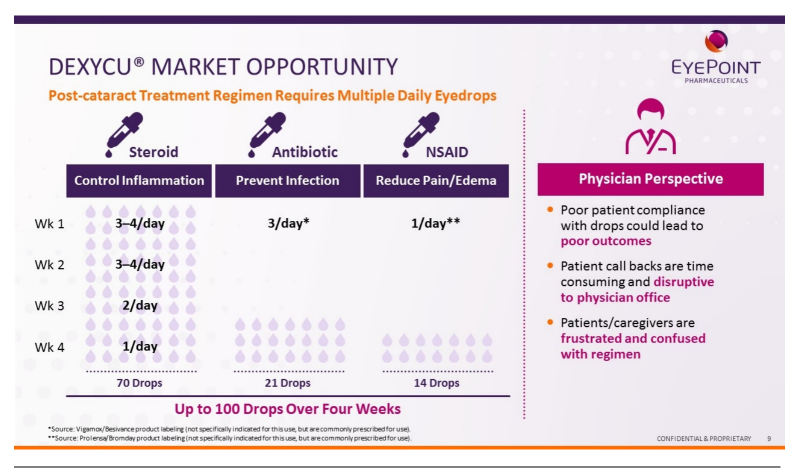
DEXYCU[®] - Significant Market Opportunity ~\$2B



U.S. Cataract Surgery Very Large and Growing



Source: imaged from the American Optometric Association.

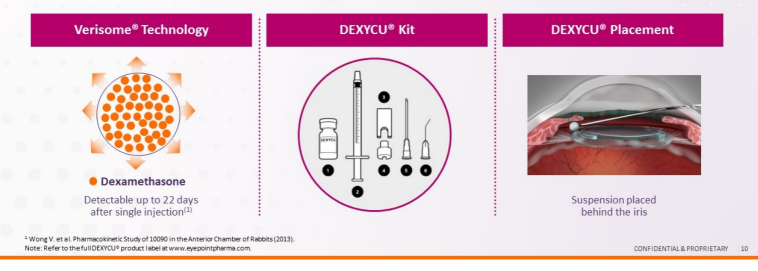


DEXYCU®



First and Only FDA-approved Single-dose, Sustained-release, Intracameral Steroid for the Treatment of Postoperative Inflammation Following Ocular Surgery

- Single dose (5µL) administered in the posterior chamber (behind the iris) at the end of surgery
- Encapsulated in bioerodible Verisome® technology for extended release of dexamethasone



DEXYCU DEMONSTRATION

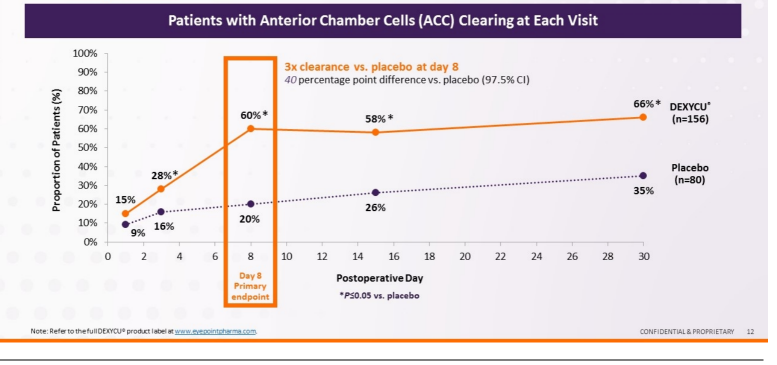


Please see video at company website: https://eyepointpharma.com/case-study-series/

DEXYCU[®]



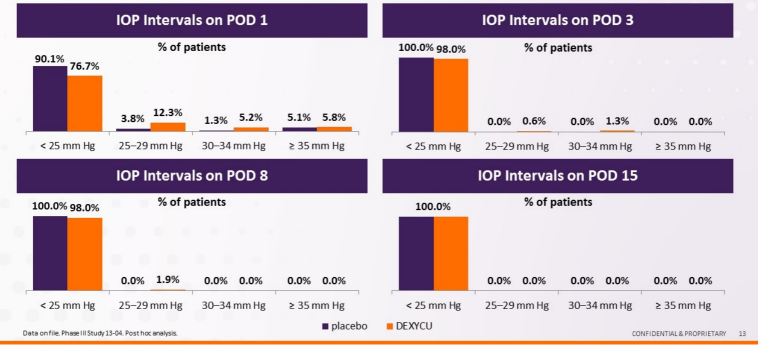
Statistically Significant Inflammation Reduction



DEXYCU[®]

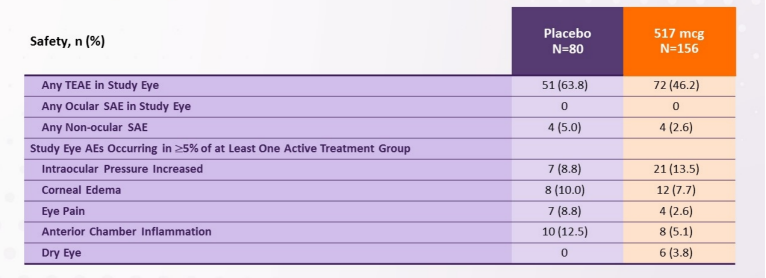
EYEPOINT PHARMACEUTICALS

IOP Elevation Versus Placebo Not Clinically Significant



DEXYCU®

Phase 3 Study - Safety Results

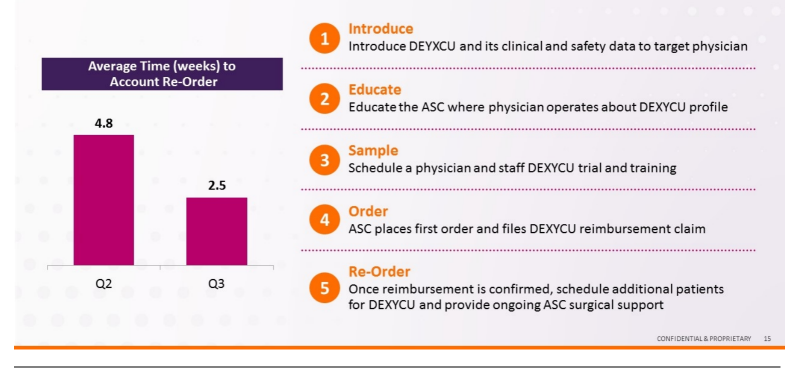


Note: Refer to the fullDEXYCU® product label at <u>www.eyepointpharma.com</u>.



DEXYCU® SALES PROCESS





DEXYCU[®] LAUNCH PROGRESS UPDATE

4Q19 Compared to 3Q19





September 2019 Represented the Highest Volume Month for Repeat Orders to Date

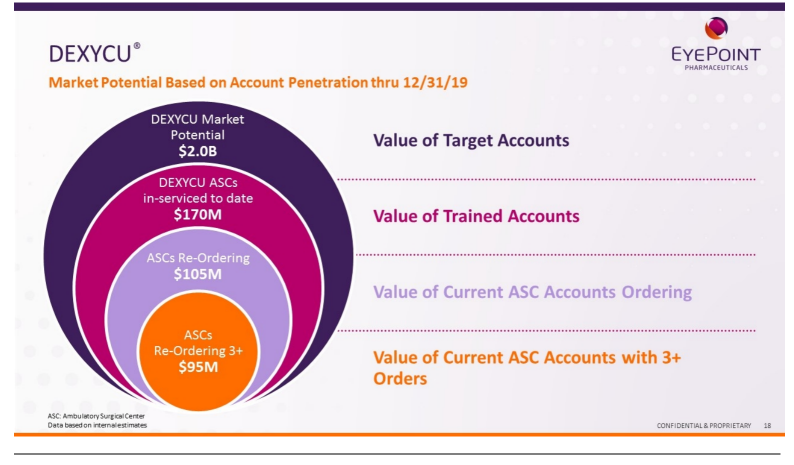
ASC: Ambulatory Surgical Center

DEXYCU® CUMULATIVE ORDERS SINCE LAUNCH



Month Over Month Growth Accelerating

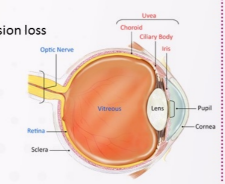




YUTIQ[®] Market



- Uveitis *is*:
 - Inflammation of the Uveal tract (iris, ciliary body, choroid), or adjacent structures (lens, retina, vitreous, optic nerve),
 - Acute or Chronic,
 - A precursor to severe vision loss or blindness
 - Often lifelong



🖤 ~55К–120К

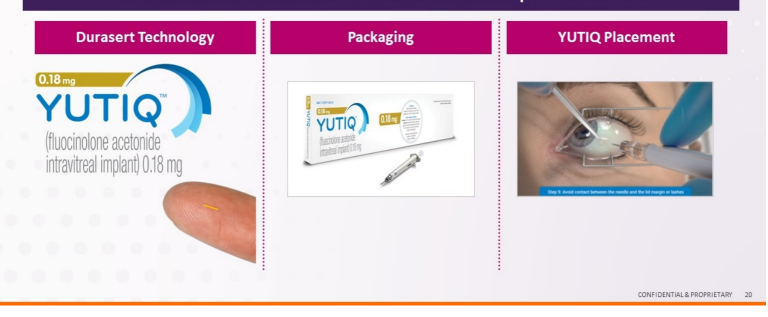
Patients in the U.S. with Chronic Non-infectious Posterior Segment Uveitis

- ~30,000 new cases of blindness per year in the U.S.
- 3rd leading cause of blindness in the U.S.

YUTIQ®



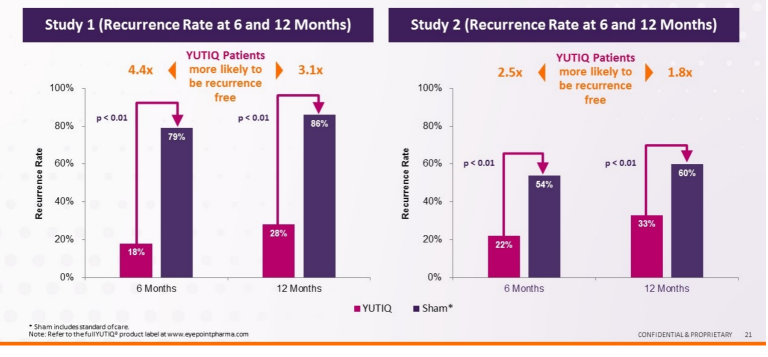
YUTIQ[®] Is Designed to Deliver a Sustained Release of Fluocinolone for Patients with Chronic Noninfectious Posterior Uveitis for Up To 36 Months





YUTIQ®

Recurrence Rate at Six and Twelve Months vs Sham

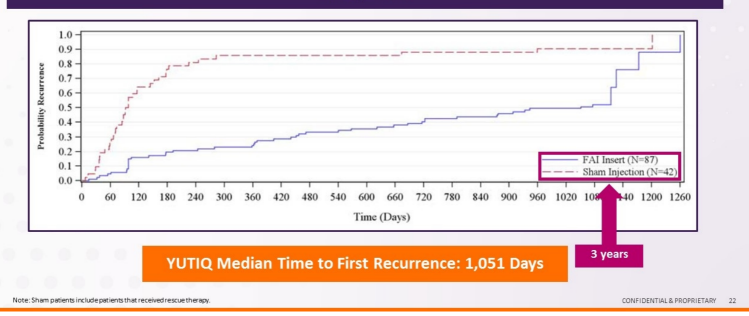


YUTIQ®



Single Insert Reduced Probability of Uveitis Recurrence Through 36 Months

ITT Population



YUTIQ® Safety – Select Adverse Events



Safety, n (%)	YUTIQ [®] n=226	Placebo n=94
Visual Acuity Reduced	33 (15%)	11 (12%)
Macular Edema ¹	25 (11)	33 (35)
Uveitis	22 (10)	33 (35)
Conjunctival Hemorrhage	17 (8)	5 (5)
Eye Pain ²	17 (8)	12 (13)
Hypotony of the Eye ³	16 (7)	1 (1)
Anterior Chamber Inflamm.	12 (5)	6 (6)
Dry Eye	10 (4)	3 (3)

Includes macular edema and cystoid macular edema
 Includes eye pain and procedural pain
 Includes hypotony, intraocular pressure decreased and procedural hypotension Note: Refer to the fullYUTIQ[®] product label at www.eyepointpharma.com

YUTIQ[®] LAUNCH PROGRESS UPDATE

Q319 compared to Q219

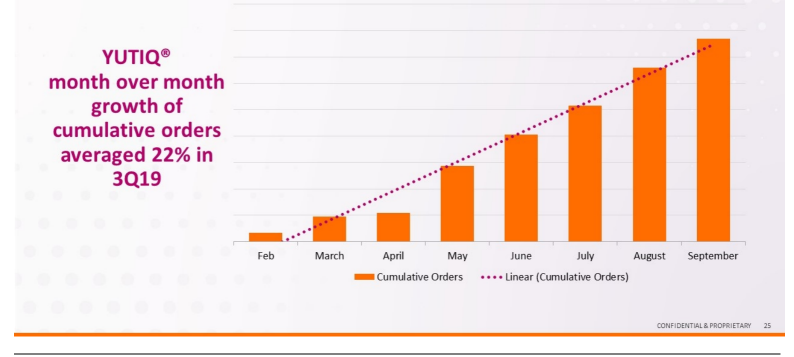




Continued Strong Reception of the YUTIQ® Product Profile from Uveitis Specialists

YUTIQ® CUMULATIVE ORDERS SINCE LAUNCH





KEY ACCESS AGREEMENTS TO EXPAND PRODUCT REACH







- DEXYCU[®] and YUTIQ[®] added to the Federal Supply Schedule
- Access to U.S. veterans and other federal agencies
- Nine Million VA beneficiaries added



- Three-year agreement for DEXYCU[®]
- Vizient's network includes more than 50% of the nation's acute care providers, including 95% of the nation's academic medical centers, and more than 20% of ambulatory care providers

One of Largest Integrated Delivery Systems in the U.S.

- DEXYCU[®] available to its 8.5 million patients
- 2 year contract includes California, Washington, Georgia, Colorado and Mid-Atlantic states

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