

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): July 30, 2026

FULGENT GENETICS, INC.
(Exact Name of Registrant as Specified in Charter)

Delaware
(State or other jurisdiction of
incorporation)

001-37894
(Commission File Number)

81-2621304
(IRS Employer Identification No.)

4399 Santa Anita Avenue
El Monte, California
(Address of Principal Executive Offices)

91731
(Zip Code)

(626) 350-0537
(Registrant's telephone number, including area code)

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 \$0.0001 per share	FLGT	The Nasdaq Stock Market (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 July 30, 2026, Fulgent Genetics, Inc. (the “Company”) issued a press release announcing its financial results for the fiscal quarter ended June 30, 2026. A copy of the Company’s press release containing this information is being furnished as Exhibit 99.1 to this Current Report on Form 8-K.

Item 7.01 Regulation FD Disclosure.

From time to time, the Company presents and/or distributes slides and presentations to the investment community to provide updates and summaries of its business. On July 30, 2026, the Company updated its investor presentation, which is available on the Investor Relations section of the Company’s website at <http://ir.fulgentgenetics.com>. This presentation is also furnished as Exhibit 99.2 to this Current Report on Form 8-K.

The information in Items 2.02 and 7.01, including Exhibits 99.1 and 99.2, is being furnished and 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, nor shall it be deemed incorporated by reference into any registration statement or other filing under the Securities Act of 1933, as amended, or the Exchange Act, except as shall be expressly set forth by specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No.	Description
99.1	Press Release of Fulgent Genetics, Inc., dated July 30, 2026
99.2	Corporate Presentation of Fulgent Genetics, Inc.
104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

SIGNATURE

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.

Date: July 30, 2026

FULGENT GENETICS, INC.

By:	<u>/s/ Paul Kim</u>
Name:	Paul Kim
Title:	Chief Financial Officer

Fulgent Reports Second Quarter 2026 Financial Results

- Revenue of \$85.4 million, growing 20.1% sequentially
- GAAP gross profit of \$25.7 million, or GAAP gross margin of 30.1%; Non-GAAP gross profit of \$26.8 million, or Non-GAAP gross margin of 31.3%
- GAAP loss of \$29.5 million, or \$(1.05) per share; Non-GAAP loss of \$16.2 million, or \$(0.58) per share
- Executed on stock repurchase program; purchased approximately 1.5 million shares in the second quarter of 2026 totaling \$23.8 million in cash
- Ended the second quarter of 2026 with \$551.5 million of cash, cash equivalents, restricted cash, and investments in marketable securities, excluding an anticipated tax refund of approximately \$106.1 million
- Revised annual revenue guidance to be in the range of \$330.0-\$340.0 million

EL MONTE, CA, July 30, 2026 — Fulgent Genetics, Inc. (NASDAQ: FLGT) (“Fulgent,” or the “Company”), a technology-based company with a well-established laboratory services business and a therapeutic development business, today announced financial results for its second quarter ended June 30, 2026.

Second Quarter 2026 Results:

- Revenue of \$85.4 million
- GAAP loss of \$29.5 million, or \$(1.05) per share
- Non-GAAP loss of \$16.2 million, or \$(0.58) per share
- Adjusted EBITDA loss of \$17.1 million

Non-GAAP income (loss), non-GAAP income (loss) per share, adjusted EBITDA income (loss), non-GAAP gross profit and margin, and non-GAAP operating income (loss) and margin, are described below under “Note Regarding Non-GAAP Financial Measures” and are reconciled to the most directly comparable GAAP financial measure, GAAP income (loss), GAAP gross profit and margin, and GAAP operating income (loss) and margin, in the accompanying tables.

Ming Hsieh, Chairman of the Board of Directors and Chief Executive Officer, said, “In the second quarter, we had the honor of presenting the encouraging findings from Phase 2 data for FID-007 at the 2026 American Society of Clinical Oncology (ASCO) meeting, and we are encouraged by our clinical trial progress for both FID-007 and FID-022. We continue to execute on our strategic initiatives in our laboratory services business as we expand our AI portfolio and integrate digital pathology and are seeing a real impact from our AI tools with improved efficiency and quality.”

Paul Kim, Chief Financial Officer, said, “We are revising our revenue guidance for the year as a result of processing delays impacting our collection rate related to the final phase of the transition of our revenue cycle management and billing system. We are working to remedy the issue and remain confident in the fundamental strength of our business.”

Outlook:

For the full year 2026, Fulgent now expects:

- Revenue to be in the range of \$330.0-\$340.0 million
- Non-GAAP loss of approximately \$63.0-\$67.0 million
- Non-GAAP loss of approximately \$2.22-\$2.35 per share, which includes the impact of a reduction of 4.1 million shares used for the stock repurchase program, year to date as of June 30, 2026
- Cash, cash equivalents, restricted cash, and investments in marketable securities of approximately \$610.0 million, which includes \$63.9 million cash used for the stock repurchase program, year to date as of June 30, 2026*

*Cash expenditures may be higher or lower than currently estimated due to a variety of factors and circumstances, including as a result of the Company's ongoing stock repurchase program, or other expenditures outside the ordinary course of business, including M&A. This number further assumes receipt of approximately \$106.1 million in tax refunds prior to December 31, 2026, which have been delayed as a result of constrained resources at the IRS; and assumes capital purchases of \$12 million, no further repurchases under the stock repurchase program, and spend on the therapeutic development business of \$26 million. The timing and amount of any future repurchases pursuant to the stock repurchase program will depend on a variety of factors, including the market price of Fulgent's common stock, general market and economic conditions and other factors Fulgent's board of directors may deem relevant.

Conference Call Information

Fulgent will host a conference call for the investment community today at 4:30 PM ET (1:30 PM PT) to discuss its second quarter 2026 results. The call may be accessed through a live audio webcast in the Investor Relations section of the Company's website, <http://ir.fulgentgenetics.com>. An audio replay will be available at the same location.

Note Regarding Non-GAAP Financial Measures

Certain information set forth in this press release and/or to be discussed on the Company's earnings call, including non-GAAP income (loss), non-GAAP income (loss) per share, adjusted EBITDA income (loss), non-GAAP gross profit and margin, and non-GAAP operating income (loss) and margin, are non-GAAP financial measures. Fulgent believes this information is useful to investors because it provides a basis for measuring the performance of the Company's business, excluding certain income or expense items that management believes are not directly attributable to the Company's operating results. Fulgent defines non-GAAP income (loss) as net income (loss) calculated in accordance with accounting principles generally accepted in the United States of America, or GAAP, plus amortization of intangible assets, plus impairment loss, plus equity-based compensation expenses, plus acquisition-related costs, which include one-time banker fee, legal, valuation, due diligence, and closing costs, plus acquisition-related severance, plus or minus the non-GAAP tax effect, and plus or minus other charges or gains, as identified, that management believes are not representative of the Company's operations. The non-GAAP tax effect was calculated by excluding from the GAAP provision the impact of the amortization of intangible assets, impairment loss, equity-based compensation expenses, acquisition-related costs, and acquisition-related severance. Fulgent defines adjusted EBITDA income (loss) as GAAP income (loss) plus or minus interest (expense) income, plus or minus provisions (benefits) for income taxes, plus depreciation and amortization, plus equity-based compensation expenses, plus insurance expense related to transferable tax

credit, plus impairment loss, plus acquisition-related costs, plus acquisition-related severance, and plus or minus other charges or gains, as identified, that management believes are not representative of the Company’s operations. Fulgent defines non-GAAP gross profit as gross profit calculated in accordance with GAAP plus equity-based compensation included in cost of revenue as shown in the table below. Fulgent defines non-GAAP gross margin by taking non-GAAP gross profit and dividing it by GAAP revenue. Fulgent defines non-GAAP operating profit (loss) by taking GAAP operating profit (loss) and adding equity-based compensation expense, amortization of intangible assets, impairment of intangible assets, acquisition-related costs, and acquisition-related severance. Non-GAAP operating margin is calculated by taking non-GAAP operating profit (loss) and dividing it by GAAP revenue. Fulgent may continue to incur expenses similar to the items added to or subtracted from the GAAP financial measures, and, accordingly, the exclusion of these items in the presentation of these non-GAAP financial measures should not be construed as an implication that these items are unusual, infrequent or non-recurring. Management uses these non-GAAP financial measures along with the most directly comparable GAAP financial measure in evaluating the Company’s operating performance and for internal planning and budgeting. Non-GAAP financial measures should not be considered in isolation from, or as a substitute for, financial information presented in conformity with GAAP, and non-GAAP financial measures as reported by Fulgent may not be comparable to similarly titled metrics reported by other companies. The Company does not provide reconciliations of forward-looking non-GAAP measures to the most directly comparable GAAP measures because the information necessary to calculate such reconciliations is unavailable on a forward-looking basis without unreasonable effort. This is due to the inherent difficulty of forecasting the timing and amounts of items that would be included in the GAAP measures, including, but not limited to, equity-based compensation, tax effects, acquisition-related items, and potential impairments, any of which could be material. The Company is also unable to predict the probable significance of such items.

About Fulgent

Fulgent is a technology-based company with a well-established laboratory services business and a therapeutic development business. Fulgent’s laboratory services business includes technical laboratory and testing services and professional interpretation of laboratory results by licensed physicians. Fulgent’s therapeutic development business is focused on developing drug candidates for treating a broad range of cancers using a novel nanoencapsulation and targeted therapy platform designed to improve the therapeutic window and pharmacokinetic profile of new and existing cancer drugs. The Company aims to transform from a diagnostic business into a fully integrated precision medicine company.

Forward-Looking Statements

This press release contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Forward-looking statements are often identified by words such as “anticipate”, “believe,” “contemplate,” “continue,” “could,” “estimate,” “expect,” “guidance,” “intend,” “may,” “plan,” “project,” “should,” “target,” “will,” and similar expressions. Examples of forward-looking statements in this press release include statements about, among other things: future performance; guidance, including guidance regarding expected quarterly and annual financial results, revenue, GAAP loss, non-GAAP loss and related per share figures, and cash, cash equivalents, restricted cash, and investments in marketable securities; potential improvements in margins; evaluations and judgments regarding the strength of the Company’s business; the potential benefits of the continued integration of AI into the Company’s laboratory business and expansion of the Company’s AI portfolio; any references (express or implied) to the potential benefits of the StrataDx and Bako Diagnostics acquisitions, including

any potential or expected revenue and whether the revenue will offset loss of other revenues; the Company's ability to remediate operational issues relating to the transition of its billing and revenue cycle management systems; the Company's continued and future research and development efforts, including any implications that the results of earlier clinical trials will be representative or consistent with later clinical trials; and the Company's identification and evaluation of opportunities and its ability to capitalize on opportunities, capture market share, or expand its presence in certain markets; and the Company's ability to continue to grow its business.

Forward-looking statements are statements other than historical facts and relate to future events or circumstances or the Company's future performance, and they are based on management's current assumptions, expectations, and beliefs concerning future developments and their potential effect on the Company's business. These forward-looking statements are subject to a number of risks and uncertainties, which may cause the forward-looking events and circumstances described in this press release to not occur, and actual results to differ materially and adversely from those described in or implied by the forward-looking statements. These risks and uncertainties include, among others, risks regarding: the market potential for, and the rate and degree of market adoption of, the Company's tests; its ability to maintain turnaround times and otherwise keep pace with rapidly changing technology; the Company's ability to maintain the low internal costs of its business model; the Company's ability to maintain an acceptable margin; risks related to volatility in the Company's results, which can fluctuate significantly from period to period; risks associated with the composition of the Company's customer base, which can fluctuate from period to period and can be comprised of a small number of customers that account for a significant portion of the Company's revenue; dependence on a limited number of customers, including risks that any such customer may further reduce, delay, or internalize testing volumes; risks related to the Company's acquisitions, including Bako and StrataDx, such as integration challenges, costs, and the Company's ability to realize expected benefits on anticipated timelines; the Company's level of success in obtaining coverage and adequate reimbursement and collectability levels from third-party payors for its tests and testing services, including its ability to manage and to remediate billing software implementation issues and claims processing delays; the Company's level of success in establishing and obtaining the intended benefits from partnerships, strategic investments, joint ventures, acquisitions, or other relationships; the success of the Company's development efforts, including the Company's ability to progress its candidates through clinical trials on the timelines expected; the Company's compliance with the various evolving and complex laws and regulations applicable to its business and its industry; the Company's ability to obtain expected tax refunds on the timelines expected; and the Company's ability to protect its proprietary technology and intellectual property. As a result of these risks and uncertainties, forward-looking statements should not be relied on or viewed as predictions of future events.

The forward-looking statements made in this press release speak only as of the date of this press release, and the Company assumes no obligation to update publicly any such forward-looking statements to reflect actual results or to changes in expectations, except as otherwise required by law.

The Company's reports filed with the U.S. Securities and Exchange Commission, or the SEC, including its annual report on Form 10-K for the fiscal year ended December 31, 2025, filed with the SEC on February 27, 2026, its quarterly report on Form 10-Q for the fiscal year ended March 31, 2026, filed with the SEC on May 1, 2026, and the other reports it files from time to time, including subsequently filed annual, quarterly and current reports, are made available on the Company's website and on the SEC's website at www.sec.gov upon their filing with the SEC. These reports contain more information about the Company, its business and the risks affecting its business, as well as its results of operations for the periods covered by the financial results included in this press release.

Investor Relations Contact:
The Blueshirt Group
Lauren Sloane, lauren@blueshirtgroup.com

FULGENT GENETICS, INC.
Condensed Consolidated Balance Sheet Data
June 30, 2026, and December 31, 2025
(in thousands)

	June 30, 2026	December 31, 2025
ASSETS:		
Cash and cash equivalents	\$ 26,305	\$ 50,193
Investments in marketable securities	525,043	655,153
Accounts receivable, net	64,938	84,762
Property, plant, and equipment, net	113,068	112,549
Other assets	359,869	310,868
Total assets	\$ 1,089,223	\$ 1,213,525
LIABILITIES & EQUITY:		
Accounts payable, accrued liabilities and other liabilities	\$ 93,237	\$ 106,810
Total stockholders' equity	995,986	1,106,715
Total liabilities & equity	\$ 1,089,223	\$ 1,213,525

FULGENT GENETICS, INC.
Condensed Consolidated Statement of Operations Data
Three and Six Months Ended June 30, 2026, and 2025
(in thousands, except per share data)
(unaudited)

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Revenue	\$ 85,389	\$ 81,803	\$ 156,527	\$ 155,266
Cost of revenue (1)	59,710	47,368	109,358	92,485
Gross profit	25,679	34,435	47,169	62,781
Operating expenses				
Research and development (1)	14,554	13,480	28,730	25,875
Selling and marketing (1)	15,228	12,286	27,449	20,751
General and administrative (1)	27,195	26,392	54,879	51,683
Amortization of intangible assets	2,603	1,990	4,634	3,980
Impairment of intangible assets	2,172	—	2,172	—
Total operating expenses	61,752	54,148	117,864	102,289
Operating loss	(36,073)	(19,713)	(70,695)	(39,508)
Interest income	6,404	8,091	15,055	16,109
Interest expense	(59)	(17)	(76)	(31)
Impairment of equity securities	—	(9,926)	—	(9,926)
Other income, net	38	46	48	114
Total other income (expense), net	6,383	(1,806)	15,027	6,266
Loss before income taxes	(29,690)	(21,519)	(55,668)	(33,242)
Provision for (benefit from) income taxes	29	(2,263)	(675)	(2,087)
Net loss from consolidated operations	(29,719)	(19,256)	(54,993)	(31,155)
Net loss attributable to noncontrolling interests	189	299	637	668
Net loss attributable to Fulgent	\$ (29,530)	\$ (18,957)	\$ (54,356)	\$ (30,487)
Net loss per common share attributable to Fulgent:				
Basic	\$ (1.05)	\$ (0.62)	\$ (1.85)	\$ (0.99)
Diluted	\$ (1.05)	\$ (0.62)	\$ (1.85)	\$ (0.99)
Weighted-average common shares:				
Basic	28,005	30,544	29,435	30,687
Diluted	28,005	30,544	29,435	30,687
(1) Equity-based compensation expense was allocated as follows:				
Cost of revenue	\$ 1,085	\$ 1,737	\$ 2,546	\$ 3,517
Research and development	3,136	3,339	6,673	6,813
Selling and marketing	565	711	1,210	1,601
General and administrative	3,838	4,252	8,085	8,658
Total equity-based compensation expense	\$ 8,624	\$ 10,039	\$ 18,514	\$ 20,589

FULGENT GENETICS, INC.
Non-GAAP Income (Loss) Reconciliation
Three and Six Months Ended June 30, 2026, and 2025
(in thousands, except per share data)

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Net loss attributable to Fulgent	\$ (29,530)	\$ (18,957)	\$ (54,356)	\$ (30,487)
Amortization of intangible assets	2,603	1,990	4,634	3,980
Impairment loss (1)	2,172	9,926	2,172	9,926
Equity-based compensation expense	8,624	10,039	18,514	20,589
Acquisition-related costs (2)	17	—	2,649	—
Acquisition-related severance (3)	251	—	647	—
Non-GAAP tax effect	(303)	(919)	(1,403)	(763)
Non-GAAP (loss) income attributable to Fulgent	<u>\$ (16,166)</u>	<u>\$ 2,079</u>	<u>\$ (27,143)</u>	<u>\$ 3,245</u>
Net loss per common share attributable to Fulgent:				
Basic	\$ (1.05)	\$ (0.62)	\$ (1.85)	\$ (0.99)
Diluted	\$ (1.05)	\$ (0.62)	\$ (1.85)	\$ (0.99)
Non-GAAP (loss) income per common share attributable to Fulgent:				
Basic	\$ (0.58)	\$ 0.07	\$ (0.92)	\$ 0.11
Diluted	\$ (0.58)	\$ 0.07	\$ (0.92)	\$ 0.11
Weighted average common shares:				
Basic	28,005	30,544	29,435	30,687
Diluted	28,005	30,724	29,435	30,797

(1) Consists of non-cash charges related to (i) a 2026 impairment of an intangible asset previously acquired through a business combination and (ii) a 2025 impairment of a prior investment.

(2) Consists of acquisition-related costs related to the acquisition of Bako and StrataDx for the three and six months ended June 30, 2026.

(3) Consists of one-time severance payment to the former CEO & CFO of Bako and StrataDx.

FULGENT GENETICS, INC.
Non-GAAP Adjusted EBITDA Reconciliation
Three and Six Months Ended June 30, 2026, and 2025
(in thousands)

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Net loss attributable to Fulgent	\$ (29,530)	\$ (18,957)	\$ (54,356)	\$ (30,487)
Interest income, net	(6,345)	(8,074)	(14,979)	(16,078)
Provision for (benefit from) income taxes	29	(2,263)	(675)	(2,087)
Depreciation and amortization	7,676	6,054	13,676	11,973
Equity-based compensation expense	8,624	10,039	18,514	20,589
Insurance expense related to transferable tax credits	—	283	—	283
Impairment loss	2,172	9,926	2,172	9,926
Acquisition-related costs	17	—	2,649	—
Acquisition-related severance	251	—	647	—
Adjusted EBITDA	<u>\$ (17,106)</u>	<u>\$ (2,992)</u>	<u>\$ (32,352)</u>	<u>\$ (5,881)</u>

FULGENT GENETICS, INC.
Non-GAAP Operating Margin
Three and Six Months Ended June 30, 2026, and 2025
(in thousands, except percentages)

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Revenue	\$ 85,389	\$ 81,803	\$ 156,527	\$ 155,266
Cost of revenue	59,710	47,368	109,358	92,485
Gross profit	25,679	34,435	47,169	62,781
Gross margin	30.1%	42.1%	30.1%	40.4%
Equity-based compensation included in cost of revenue	1,085	1,737	2,546	3,517
Non-GAAP gross profit	26,764	36,172	49,715	66,298
Non-GAAP gross margin	31.3%	44.2%	31.8%	42.7%
Operating expenses	61,752	54,148	117,864	102,289
Equity-based compensation included in operating expenses	7,539	8,302	15,968	17,072
Amortization of intangible assets	2,603	1,990	4,634	3,980
Impairment of intangible assets	2,172	—	2,172	—
Acquisition-related costs	17	—	2,649	—
Acquisition-related severance	251	—	647	—
Non-GAAP operating expenses	49,170	43,856	91,794	81,237
Non-GAAP operating loss	\$ (22,406)	\$ (7,684)	\$ (42,079)	\$ (14,939)
Non-GAAP operating margin	-26.2%	-9.4%	-26.9%	-9.6%

Investor Presentation

July 30, 2026

Founded in 2011 | Located in El Monte, CA | NASDAQ:FLGT

Disclaimer



Forward-Looking Statements and Market Data

This presentation contains forward-looking statements, which are statements other than those of historical facts and which represent the estimates and expectations of Fulgent Genetics, Inc. (the "Company" or "Fulgent") about future events based on current views and assumptions. Examples of forward-looking statements made in this presentation include, among others, those related to long-term upside or value, management of risk, anticipated growth and positioning, addressable market estimates, the Company's mission, vision and strategies, the success of its business model and strategy, anticipated future revenue and guidance, evaluations and judgments regarding the Company's business, products, technologies, competitive landscape, scalability, plans regarding development and launch of potential future products, and any businesses the Company may seek to acquire or has acquired or has invested in or may seek to invest in, including statements regarding Fulgent Pharma Holdings, Inc. ("Fulgent Pharma"), Inform Diagnostics, CSI Laboratories, Bako and StrataDx acquisitions, and any potential synergies, or transformation of the Company's business, long-term visions and strategies, the clinical development of Fulgent Pharma's pipeline and related statements and assumptions regarding development timelines, any potentially accelerated pathway for regulatory approval, the potential safety and efficacy of the nanodrug delivery platform and any related therapeutic candidates, the potential market size for these candidates and platforms and the value of available data, including genomic data, the Company's research and development efforts, including any implications that the results of earlier clinical trials will be representative or consistent with later clinical trials, the expected timing or timing of enrollment for these clinical trials or that interim or preliminary data will be representative of the final data or results of these trials, and guidance regarding the Company's future performance and results of operations, including any cash or cash equivalent resource projections. The Company's views and assumptions on which these forward-looking statements are based may prove to be incorrect. As a result, matters discussed in any forward-looking statements are subject to risks, uncertainties and changes in circumstances that may cause actual results to differ materially from those discussed or implied by any forward-looking statements. Important factors that could cause actual results to differ materially from those implied by forward-looking statements are disclosed under "Risk Factors" and "Management's Discussion and Analysis of Financial Condition and Results of Operations" in the Company's reports filed with the Securities and Exchange Commission ("SEC"), including its annual report on Form 10-K filed on February 27, 2026, and other reports it files from time to time. Because of these factors, you should not rely upon forward-looking statements as predictions of future events. The forward-looking statements in this presentation are made only as of the date hereof, and, except as required by law, the Company assumes no obligation to update any forward-looking statements in the future. The Company's reports filed with the SEC, including its annual report on Form 10-K for the year ended December 31, 2025, filed with the SEC on February 27, 2026, and the other reports it files from time to time, including subsequently filed annual, quarterly and current reports, are made available on the Company's website upon their filing with the SEC. These reports contain more information about the Company, its business and the risks affecting its business, as well as its results of operations for the periods covered by the financial results included in this presentation.

This presentation also includes market data and forecasts with respect to the industry in which the Company operates. In some cases, the Company relies upon and refers to market data and certain industry forecasts that have been obtained from third-party surveys, market research, consultant surveys, publicly available information and industry publications that the Company believes to be reliable. These data and estimates involve a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates.

Non-GAAP Financial Measures

This presentation contains certain supplemental financial measures that are not calculated pursuant to U.S. generally accepted accounting principles ("GAAP"). These non-GAAP measures are in addition to, not a substitute for or superior to, measures of financial performance prepared in accordance with GAAP. A reconciliation of non-GAAP measures to GAAP measures is contained in this presentation. Fulgent believes this information is useful to investors because it provides a basis for measuring the performance of the Company's business, excluding certain income or expense items that management believes are not directly attributable to the Company's operating results. The Company does not provide reconciliations of forward-looking non-GAAP measures to GAAP measures, due to the inability to predict the amount and timing of impacts outside of the Company's control on certain items, particularly items related to equity-based compensation, tax effects and potential impairments, among other items, which could be material. Reconciling such items would require unreasonable efforts.

Leadership Team



Ming Hsieh
Chief Executive
Officer

Experienced operational leader, entrepreneur and philanthropist

Previously CEO, President, and Chairman of Cogent Systems, Inc.

Member of the National Academy of Engineering; Fellow of the National Academy of Inventors; Trustee of USC



Paul Kim
Chief Financial
Officer

Experienced financial leader and Certified Public Accountant

Previously CFO of Cogent Systems, Inc.; sold to 3M for \$943M in 2010

B.A. in Economics from University of California at Berkeley



Dr. Harry Gao
Lab Director and
Chief Scientific
Officer

Previously Lab Director at City of Hope

Clinical molecular genetics training fellowship and post-doctoral fellowship at Harvard Medical School

M.S. in Immunology, and M.D. and Ph.D. in Microbiology, Immunology, and Medical Genetics



James Xie
President and
Chief Operating
Officer

Responsible for managing all global operations, product vision and product engineering

Served as an SVP of Cogent Systems, Inc.

B.A. in Engineering, M.S. in Industrial Engineering and an M.S. in Computer Science



Brandon Perthuis
Chief Commercial
Officer

Extensive experience leading genetic testing commercialization programs since 2003

Previously VP of Sales and Marketing of the Medical Genetics Laboratory at Baylor College of Medicine

Prior to Baylor, held senior roles at PerkinElmer, Inc. and Spectral Genomics, Inc.

B.S. in Biomedical Science



Natalie Prescott
General Counsel &
Chief Privacy Officer

Seasoned legal and privacy professional with nearly two decades of legal experience

Privacy Law Specialist; Certified Information Privacy Manager; Certified Information Privacy Professional and an Advisory Board Member with the International Association of Privacy Professionals

J.D. from Duke University School of Law



Dr. Ray Yin
President, Pharma

Founder & CEO, ANP Technologies, Inc.

Former Team Leader of Nanobiotechnology for Chem/Bio Defense, U.S. Army Research Laboratory

Holder of 46 drug delivery/detection patents

Ph.D. in Chemistry, University of Southern California



About Fulgent

We are a premier global, technology-based genetic testing company focused on transforming patient care in oncology, infectious and rare diseases, and reproductive health.



Mission

Develop flexible and affordable diagnostics and therapeutics that improve the everyday lives of those around us.

Core Values

- Innovation
- Customer Service and Commitment
- Quality and Efficiency
- Our People

Strategy

- Leverage our proprietary technology platform for broad application
- Further clinical/regulatory program for Pharma
- Operational excellence
- Disciplined M&A

Laboratory Services



\$85M

Q2 Revenue

4%

Q2 Year-over-Year
Revenue Increase

18,400+ GENES | 900+ PANELS
CUSTOMIZABLE OFFERINGS

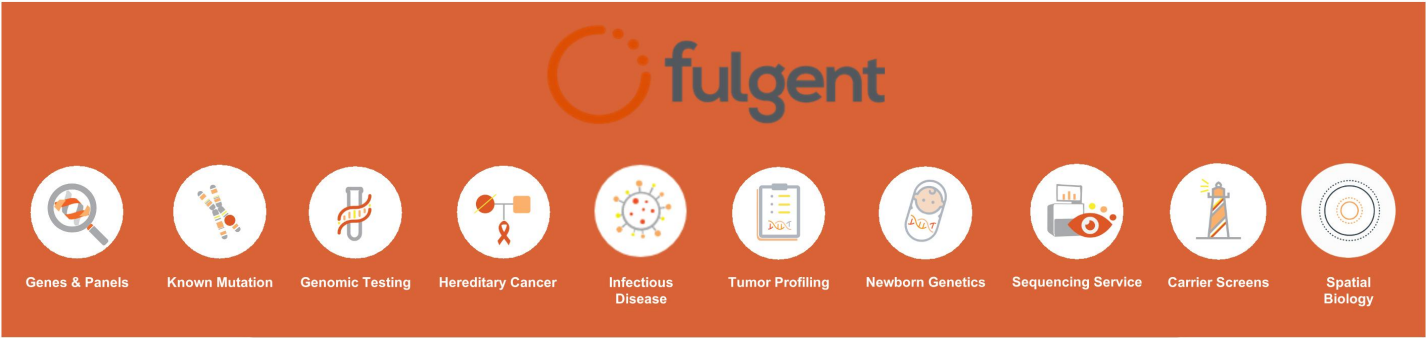
Positioned for Growth

- 1** Proprietary technology platform allows for rapid scaling of a **broad, flexible test menu**
- 2** **Next-generation sequencing (NGS)** platform complemented with growing portfolio of **emerging testing technologies** with a focus on oncology
- 3** Well-positioned to execute on a growth strategy that includes **organic and inorganic initiatives**, including:
 - Transformational acquisitions
 - Scaling partnerships
 - Acquisitions with a strategy of short- and long-term ROI, tangible synergies, and efficient capital deployment

Platform & Capabilities Across 3 Categories



Target Market Opportunity



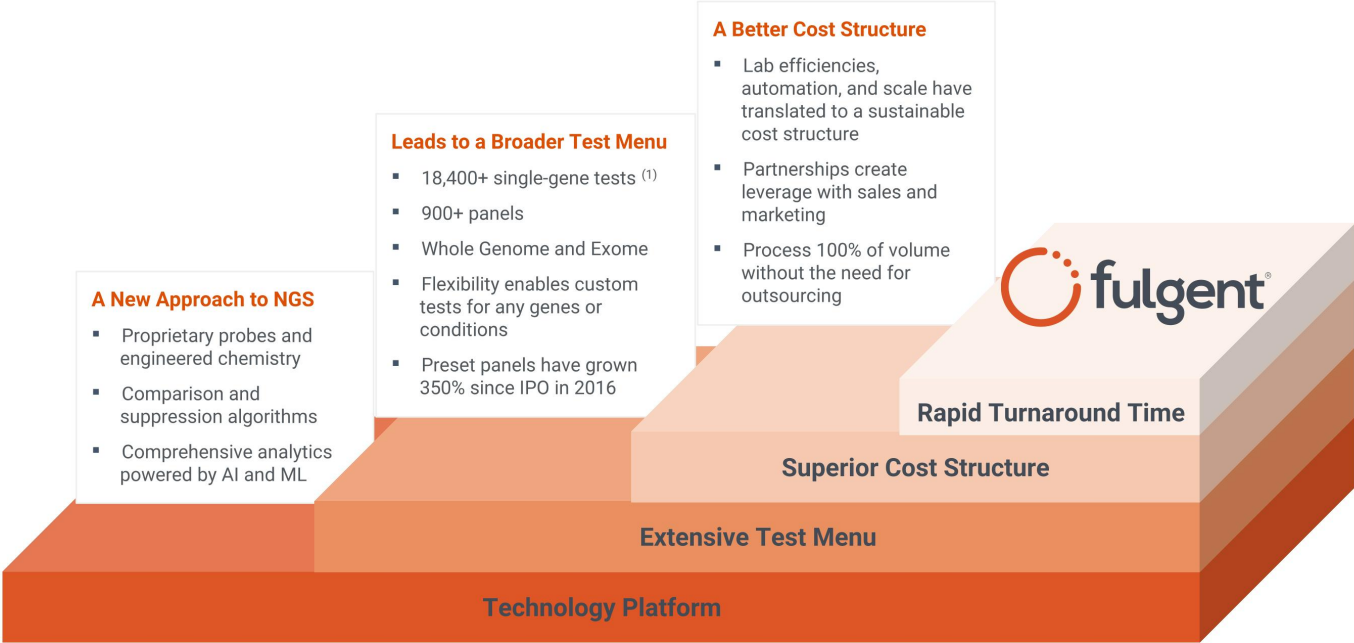
Cancer Diagnostics

**Early Detection/
Liquid Biopsy**

Reproductive Health

BioPharma Services

What Sets Fulgent Apart?



1) Represents genes covered by single-gene tests.

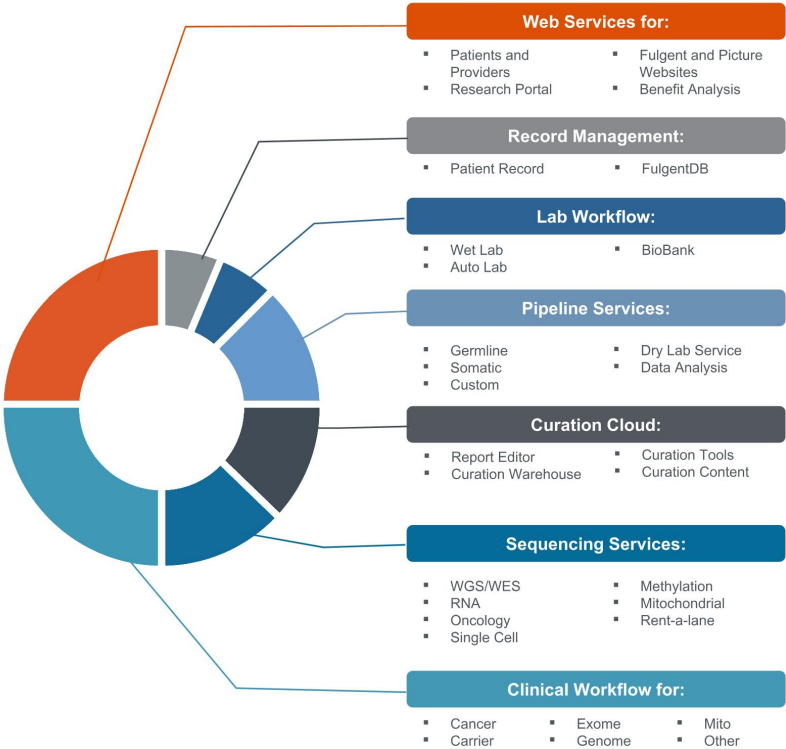
AI & Proprietary Technology Platform

Differentiated Technology...

- Engineered genetic biochemistry, including reagents and probes
- Data suppression and comparison algorithms
- Adaptive learning software
- Automated reporting

...Provides a Multitude of Advantages

- Broad test menu
- Ability to rapidly develop and launch new tests
- Customizable test offerings
- Lower costs per billable test
- High efficiency





Next Generation Sequencing Opportunities

Recent Traction with:

- Hereditary Cancer
- Cardiovascular Genetics
- Reproductive Health
- Neurodegenerative Genetics

Newly launched ultra rapid whole genome sequencing

Aggressively expanding sales and commercial organization



Specialized Oncology Testing

Wide Array of Technologies

Services Include:

- Flow cytometry
- Cytogenetic analysis
- Fluorescence in-situ hybridization (FISH)
- Immunohistochemistry
- Molecular genetics
- Consultations in hematopathology and surgical pathology
- NGS



Comprehensive Anatomic Pathology Services

Broad Capabilities

- Breast pathology
- Gastrointestinal pathology
- Dermatopathology
- Urologic pathology
- Neuropathology

Managed care contract network and **physician relationships** leveraged to provide diagnostic products and services **complementary to Fulgent's portfolio**

Expansive geographic presence with multiple **CLIA-licensed** laboratories across the United States

Expanding the Integrated Diagnostics Platform

Acquisition: Bako & StrataDx

Adjusted Purchase Price: \$56.4 Million

Closed: March 17, 2026

Test Menu Expansion: Adds premier national pathology assets, including PCR tests that optimize costs and offer rapid results.

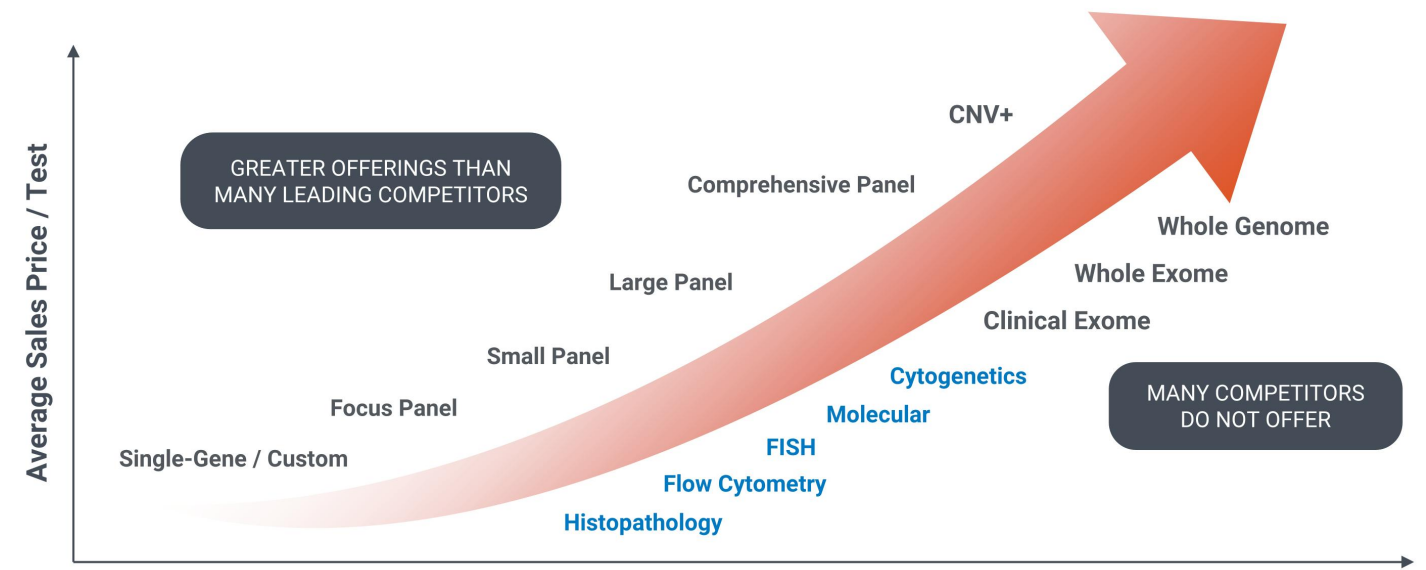
Drives commercial synergies: Nearly doubles pathology sales force headcount and introduces specialized dermatopathology expertise.

Expands national client and laboratory footprint: Addition of CLIA, CAP, and NY State certified laboratories in Georgia (Bako) and Massachusetts (StrataDx).

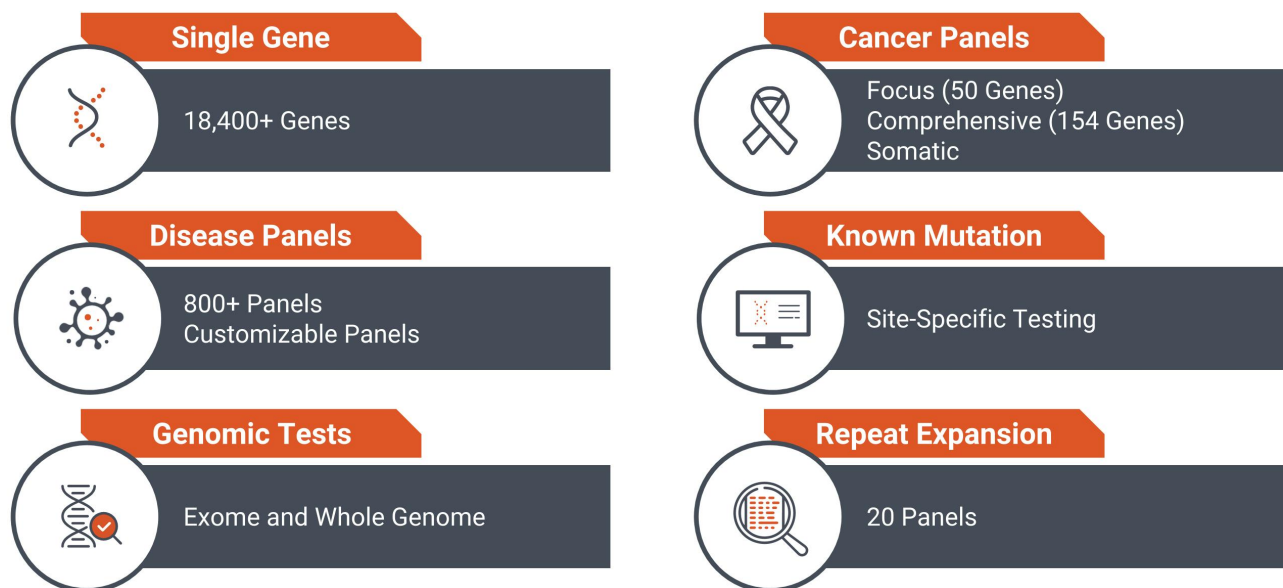
AI Integration: Leverages Eziopath, Fulgent's proprietary AI, to automate pathology workflows and improve diagnostic accuracy.



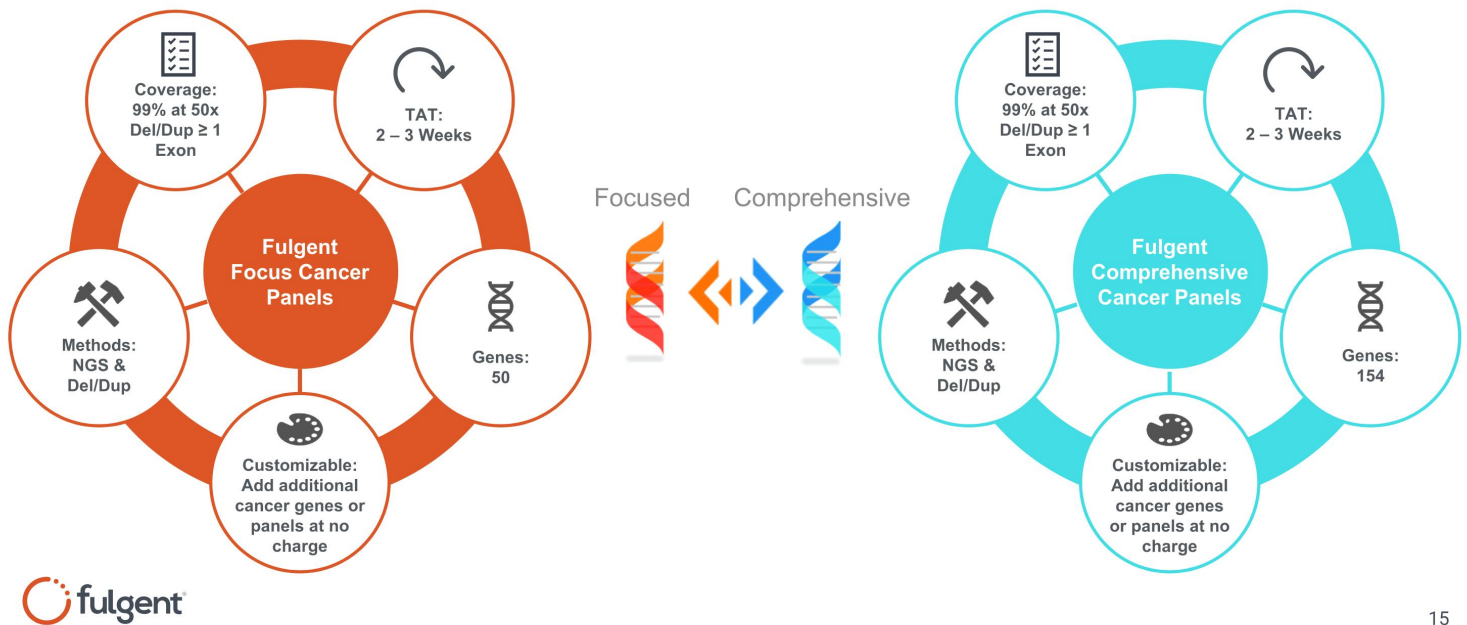
Scalable and Affordable Menu for Customers



NGS Testing – Offerings



NGS Testing – Germline Oncology Test Menu



Oncology Testing Platforms



FISH

- Expansive heme and solid tumor menu
- STAT testing available - PML/RARA <1 day turnaround time
- CD138 cell enrichment for PCM
- 3-5 day turnaround time



Histology

- 225+ stains
- Platform agnostic
 - Roche, Agilent, and Leica IHC
- Three levels of service – Tech, Global, Consultative
- PD-L1 - Various IVD platforms and indications
- <1-2 day turnaround time



Cytogenetics

- Oncology and constitutional
- >20% abnormality detection rate
- Mitogen stimulation/dual culture
- DSP30 (detection of B-cell disorders)
- Interleukin 4 for plasma cell myeloma
- Phytohemagglutinin and Interleukin 2 (detection of T-cell disorders)
- Children's Oncology Group approved
- 5-7 day turnaround time



Flow Cytometry

- 10-color platform
- Comprehensive panel design
- High-sensitivity for paroxysmal nocturnal hemoglobinuria
- Expert analysis and interpretation
- 12-24 hour turnaround time

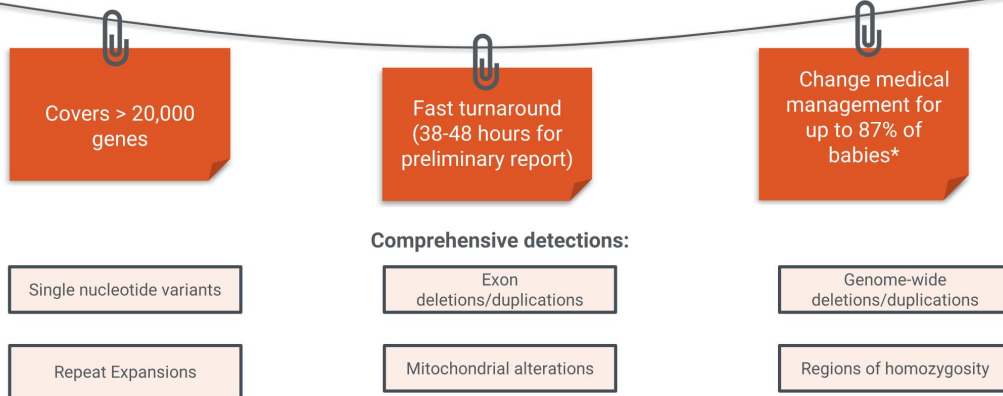


Molecular

- Hematology and solid tumor menu
- Extensive single gene menu
- NGS
- Solid tumor liquid biopsy NGS offering
- 5-7 day turnaround time

NGS Testing – Ultra Rapid Whole Genome

Designed for critically ill infants in the NICU/PICU to rapidly diagnose genetic disorders



Comprehensive detections:

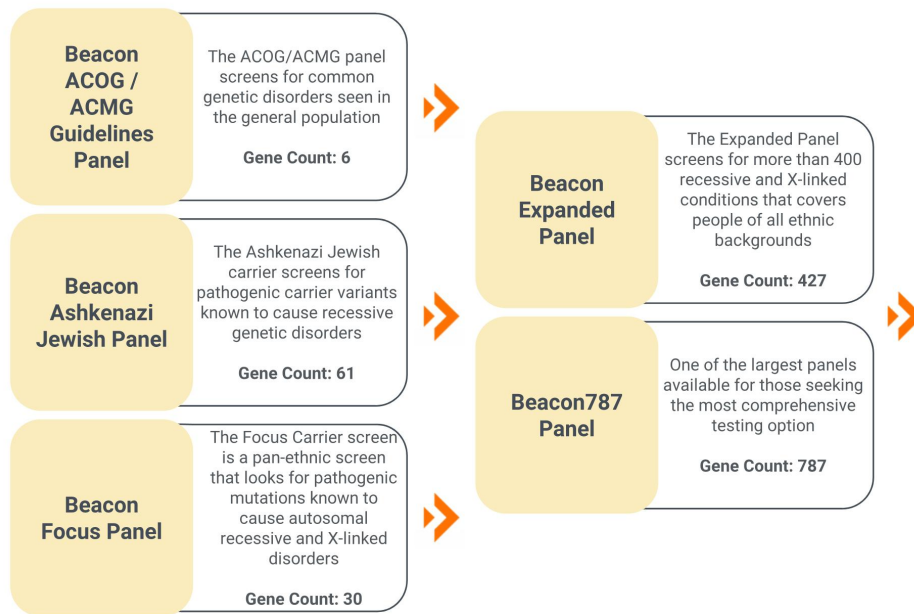
TRIO, DUO, or Proband +/- RNA-seq



* PMID 34089648 - Analysis limited to DNA analysis of SNVs and CNVs.

NGS Testing – Panel Deep Dive

Comprehensive Beacon Carrier Screening Tests



Beacon Carrier Screening

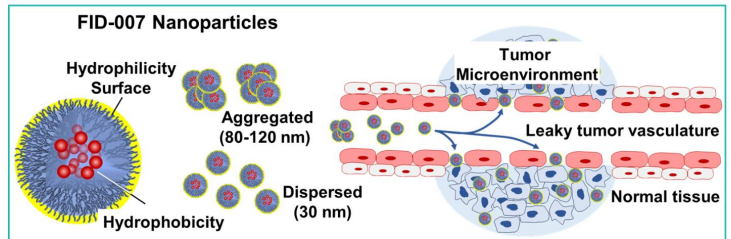
- NGS of entire genes, not just hotspots
- Deletion and duplication analysis
- Proprietary algorithms for pseudogenes
- TAT: 2 Weeks

Therapeutic Development



Background of FID-007

- Nanoencapsulated paclitaxel formulation to promote tumor penetration and retention
- PK showed a rapid and more extensive distribution to tissue/tumor with lower exposures of unbound paclitaxel when compared to solvent-based paclitaxel
- Enhanced chemotherapy agent as backbone for combination therapy



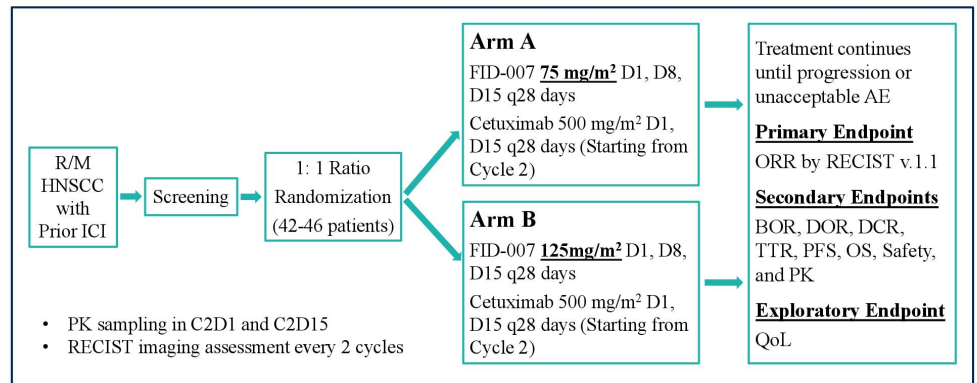
A first-in-human study of FID-007 monotherapy in advanced solid tumors (NCT03537690, N=50) demonstrated:

- a manageable safety profile without any grade ≥ 3 peripheral neuropathy
- an ORR of 45% in a subset of 11 heavily pre-treated patients with R/M HNSCC

FID-007: Study Design

Key Eligibility Criteria

- R/M HNSCC
- Disease progression after ≤ 1 prior line of systemic therapy in the R/M setting
- ECOG PS of 0 or 1
- Patients must have prior ICI treatment
- Patients with prior cetuximab or taxane treatment in the R/M setting is excluded.



Clinical Cut-Off Date: April 16, 2026

FID-007: Demographics and Baseline Disease Characteristics

		Arm A N=22 (%)	Arm B N=24 (%)
Age (years)	Median (min, max)	64.5 (49, 81)	66.0 (45, 79)
Gender	Male	17 (77.3)	20 (83.3)
	Female	5 (22.7)	4 (16.7)
Ethnicity	Hispanic or Latino	1 (4.5)	8 (33.3)
	Non-Hispanic or Latino	21 (95.5)	16 (66.7)
ECOG PS	0	4 (18.2)	10 (41.7)
	1	18 (81.8)	14 (58.3)
Primary Tumor Location	Oral cavity	12 (54.5)	13 (54.2)
	Oropharynx	8 (36.4)	8 (33.3)
	Larynx	2 (0.9)	2 (8.3)
	Paranasal sinuses	0	1 (4.2)
Prior ICI	Yes	22 (100)	24 (100)
Prior Platinum Tx	Yes	15 (68.2)	15 (62.5)
Prior Systemic Tx	Locally advanced (LA) only	8 (36.4)	7 (29.2)
	Recurrent / Metastatic (R/M) only	7 (31.8)	10 (41.7)
	Both LA and R/M	7 (31.8)	7 (29.2)
HPV (p16) Status	Positive, Oropharynx	7 (31.8)	6 (25.0)
	All other	15 (68.2)	18 (75.0)

Enrollment is completed as of Clinical Cut-Off Date April 16, 2026

- 46 patients were randomized and received ≥ 1 dose of FID-007 (ITT and safety populations)
- 42 patients were efficacy-evaluable

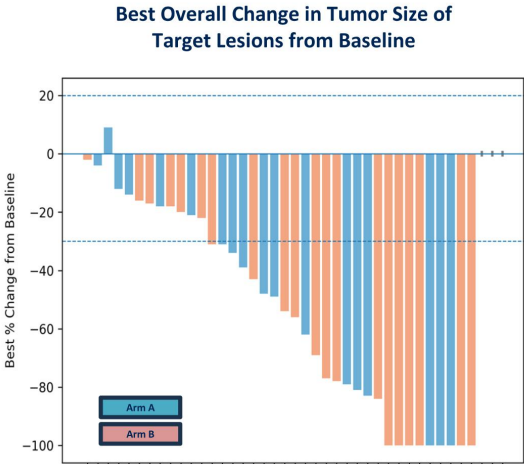
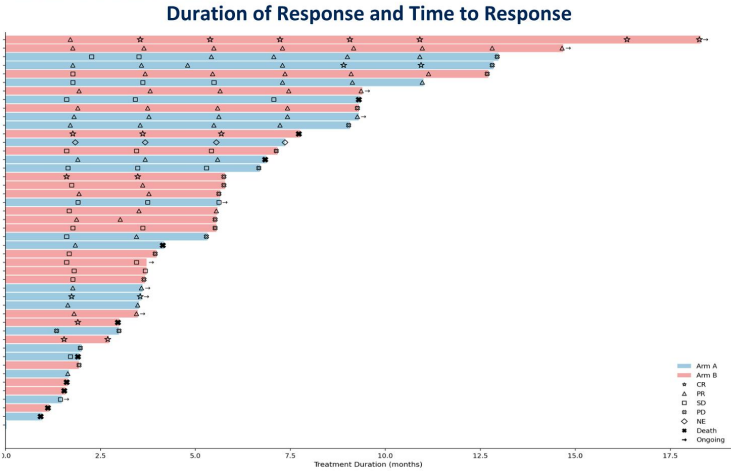
Platinum = carboplatin or cisplatin

FID-007: Best Overall Response

	Arm A (N=21)	Arm B (N=21)	Total (N=42)
BOR, n(%)			
CR	2 (9.5)	5 (23.8)	7 (16.7)
PR	10 (47.6)	9 (42.9)	19 (45.2)
SD	4 (19.0)	6 (28.6)	10 (23.8)
PD	3 (14.3)	1 (4.8)	4 (9.5)
NE	2 (9.5)	0	2 (4.8)
ORR, n (%)	12 (57.1)	14 (66.7)	26 (61.9)
DCR, n (%)	16 (76.2)	20 (95.2)	36 (85.7)

Subgroup	Category	ORR, n (%)
p16 status	p16 Positive OPC (N=11)	5 (45.5)
	Other HNSCC (N=31)	21 (67.7)
Prior Platinum Therapy	Yes (N=27)	17 (63.0)
	No (N=15)	9 (60.0)

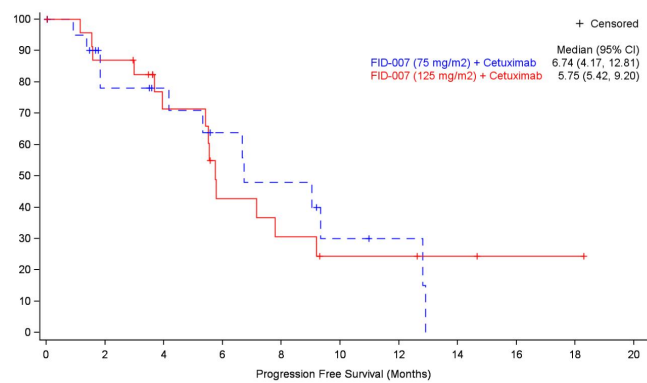
FID-007: Tumor Response and Treatment Duration



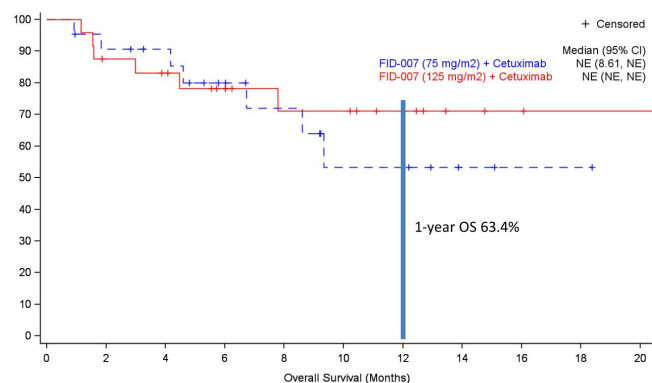
Responders	Arm A (N=12)	Arm B (N=14)
Median DOR (95% CI, mo)	7.36 (1.87, NE)	5.98 (2.17, NE)
Median TTR (Min, Max, mo)	1.77 (1.68, 7.29)	1.84 (1.64, 3.61)

FID-007: Tumor Response and Treatment Duration

Progression-Free Survival
(ITT Population)

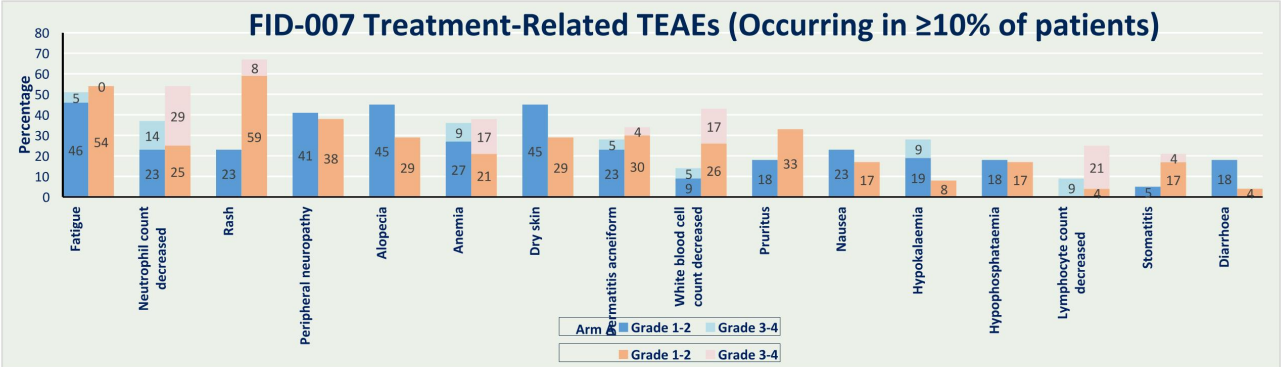


Overall Survival
(ITT Population)



ITT population	Arm A (N=22)	Arm B (N=24)	Total (N=46)
Median PFS (95% CI, mo)	6.74 (4.17, 12.81)	5.75 (5.42, 9.20)	6.67 (5.42, 9.20)

FID-007: Summary of Adverse Events



Adverse event category, n (%)	Arm A (N=22)	Arm B (N=24)	Total (N=46)
Any TEAEs	20 (90.9)	24 (100)	44 (95.7)
Any TEAEs of >= CTCAE Grade 3	15 (68.2)	22 (91.7)	37 (80.4)
Any TEAEs related to FID-007	20 (90.9)	24 (100)	44 (95.7)
Any TEAEs related to Cetuximab	18 (81.8)	21 (87.5)	39 (84.8)
Any serious TEAEs	10 (45.5)	13 (54.2)	23 (50.0)
Any TRAEs leading to dose modification of FID-007	11 (50.0)	19 (79.2)	30 (65.2)
Any TRAEs leading to withdrawal of FID-007	0	2 (8.3)	2 (4.3)
Any TRAEs leading to death*	0	1 (4.2)	1 (2.2)

* 1 Grade 5 TRAE in Arm B (pneumonia)

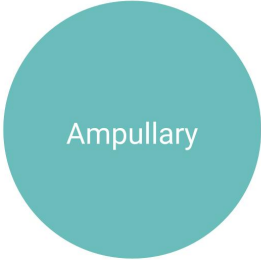
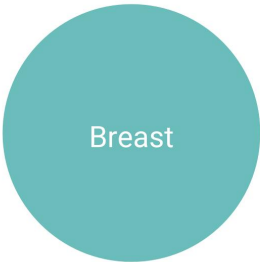
FID-007: Interim Clinical Data

- In an ongoing Phase 2, open-label trial and as of June 2026, FID-007 demonstrated encouraging clinical activity when combined with cetuximab in patients with R/M HNSCC
 - **Efficacy**
 - ORR: 61.9%
 - mDOR: 7.4 mo (95% CI: 3.7-11.1)
 - mPFS (ITT): 6.7 mo (95% CI: 5.4-9.2)
 - mOS: not mature; 1-Year OS: 63.4%
 - **Safety**
 - Manageable safety profile with no infusion-related reaction or grade ≥ 3 peripheral neuropathy
 - **Phase 3 study**
 - 125 mg/m² dose is proposed for Phase 3 study.
-

Potential Market Opportunity for FID-007



Initial Indication



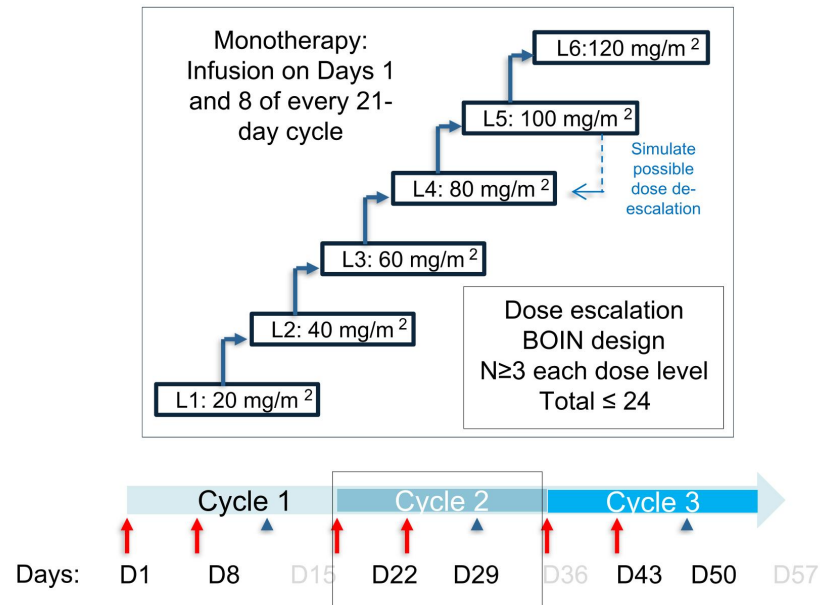
Subsequent Indications

Note: U.S. opportunity shown
Sources: Evaluate Pharma, Wall Street research, and management pricing expectations
1. Head & Neck, or H&N, market opportunity for both 2nd line and 3rd line therapy



FID-022: FIH Study Design (BOIN₁)

- FID-022-001 Clinical Update
- Dose level 1, 2 3 & 4 are completed



1. Bayesian Optimal Interval Design, a type of model-assisted dose finding design used to determine the maximum tolerated dose.

Pipeline

- FID-007: wholly-owned drug candidate initially focused on Head & Neck (H&N), Pancreatic/Ampullary cancers
 - Phase 2 trial ongoing for 2nd line treatment of H&N cancer
- FID-022 Phase 1 trial ongoing
- Potential FDA approval strategy uses 505(b)(2) studies, which may shorten clinical trial process and accelerate timeline to commercialization
- Developing a next generation antibody drug conjugate (ADC) technology platform that could potentially have better efficacy in various tumors having a broad range of target antigen expression levels when compared to current ADC benchmarks within the market

Drug Candidates	Target	Indication	Preclinical	Clinical P1	Clinical P2	Clinical P3	Milestones
FID-007	Cytotoxic	Head and Neck (H&N) (505(b)(2))					Interim Findings in June 2026
		Ampullary or ICI Resistant (505(b)(2))					Go/No-go Based on H&N Study
FID-022	Cytotoxic	Colon, Pancreatic, Ovarian, Bile Duct (505(b)(2))					Dosing completed for first 4 dose levels.
ADCs	Undisclosed	Solid Tumors					

Financials



Summary of Financial Performance

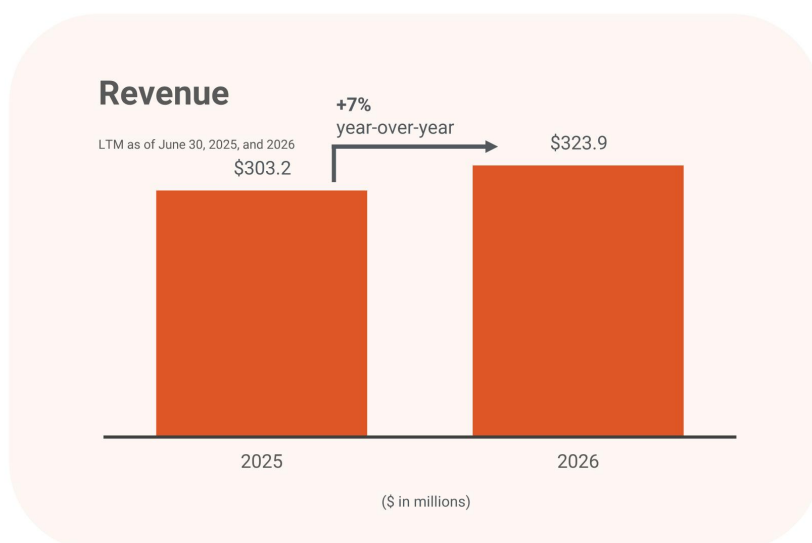
\$85M

Revenue in Q2 2026

4% increase year-over-year

\$(84)M *

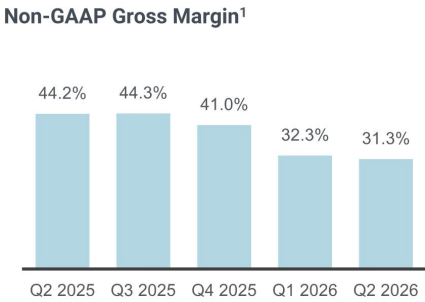
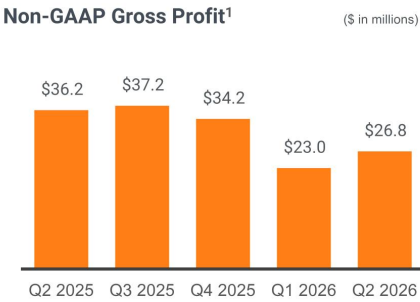
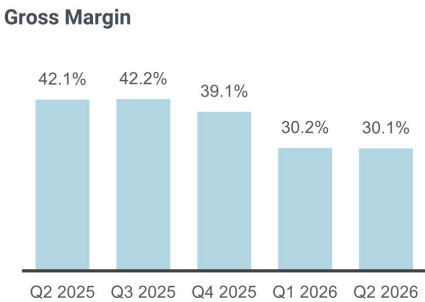
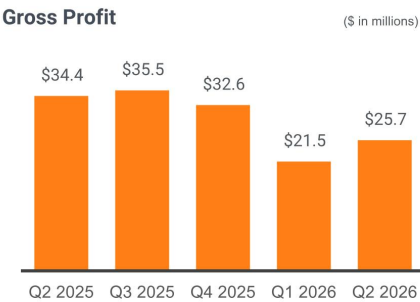
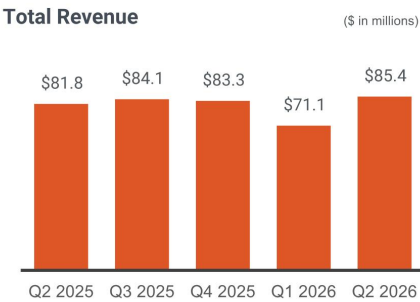
Last Twelve Months (LTM)
Operating Cash Flow as of Q2 2026



*As of quarter-end, we had not yet received the \$106M federal income tax refund which has been delayed due to constrained resources of the IRS.

Note: All figures are rounded.

Financial Performance: Revenue and Gross Margin



1) Fulgent defines non-GAAP gross profit as gross profit calculated in accordance with GAAP plus equity-based compensation included in cost of revenue, and Fulgent defines non-GAAP gross margin by taking non-GAAP gross profit and dividing it by GAAP revenue. See appendices for Non-GAAP reconciliations of these figures.

Note: All figures are rounded.

YTD 2026 Financial Performance Across Segments

GAAP (\$ in millions)	Laboratory Services		Therapeutic Development		Total	
	2026	YoY	2026	YoY	2026	YoY
Revenue	\$ 156.4	1%	\$ 0.1	*	\$ 156.5	1%
Operating expenses	\$ 101.8	14%	\$ 16.1	23%	\$ 117.9	15%
Operating loss	\$ (54.7)	-107%	\$ (16.0)	-22%	\$ (70.7)	-79%
Income (loss) before income tax	\$ (39.7)	-97%	\$ (16.0)	-22%	\$ (55.7)	-67%

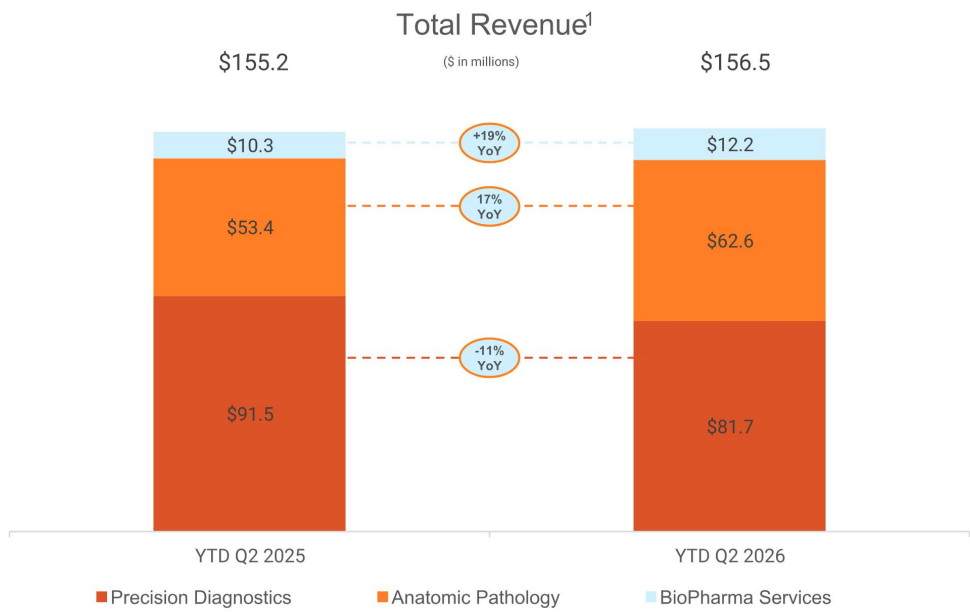
Non-GAAP ¹ (\$ in millions)	Laboratory Services		Therapeutic Development		Total	
	2026	YoY	2026	YoY	2026	YoY
Revenue	\$ 156.4	1%	\$ 0.1	*	\$ 156.5	1%
Operating expenses	\$ 82.3	14%	\$ 9.5	8%	\$ 91.8	13%
Operating loss	\$ (32.7)	-462%	\$ (9.4)	-7%	\$ (42.1)	-187%
Income (loss) before income tax	\$ (17.6)	-270%	\$ (9.4)	-7%	\$ (27.0)	-1845%

* Not meaningful

1) Non-GAAP metric excludes equity-based compensation, impairment loss, amortization of intangible assets, acquisition-related costs, and acquisition-related severance. See Appendices for Non-GAAP reconciliations of these figures.

Note: All figures are rounded.

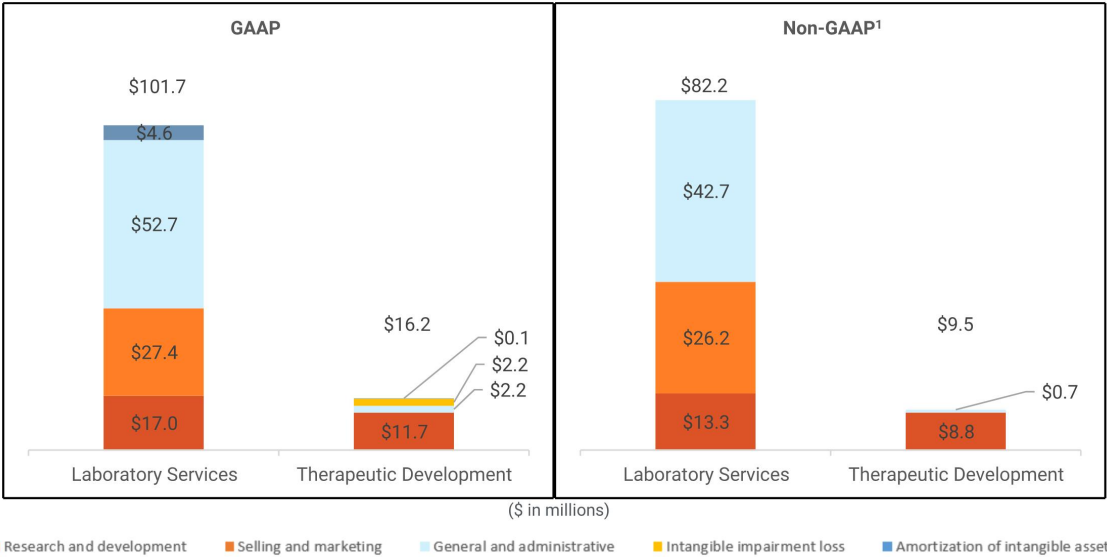
YTD Revenue by Category



1) Therapeutic development for drug candidates still in pre-commercial stage.

Note: All figures are rounded.

YTD 2026 Operating Expenses by Category



1) Non-GAAP metric excludes equity-based compensation, impairment loss, amortization of intangible assets, acquisition-related costs, and acquisition-related severance. See Appendices for Non-GAAP reconciliations of these figures.

Note: All figures are rounded.

Strategies for Success Across Segments



Laboratory Services

- ✓ Balanced growth strategy
- ✓ Enhancing operational efficiency
- ✓ Building integration readiness
- ✓ Investing in core capabilities
- ✓ Strengthening financial discipline

Target future Non-GAAP profitability



Therapeutic Development

- ✓ Completed enrollment for Phase 2 trial of FID-007 at YE 2025
- ❑ Meeting confirmed with the FDA for 2H 2026; Phase 3 protocol development is on-going
- ❑ Phase 1 clinical trial for FID-022 progressing through dose escalation

Target cash burn of ~\$26M

1) Non-GAAP profitability (income before income tax) excludes equity-based compensation, amortization of intangible assets, acquisition-related costs, any impairment loss, plus or minus other charges or gains, as identified, that management believes are not representative of the Company's operations.

2026 Financial Guidance

Metric	Full Year 2026		Expected Revenue Breakdown (\$M)		
	Low	High		Low	High
Total Revenue	\$330M	\$340M	Precision Diagnostics	\$ 161	\$ 166
% YoY Growth	+2.3%	+5.4%	Anatomic Pathology	146	150
Non-GAAP EPS ¹	\$ (2.22)	\$ (2.35)	BioPharma Services	23	24
			Total Revenue	\$ 330	\$ 340

Expected cash, cash equivalents, and investments in marketable securities of approximately \$610 million as of December 31, 2026²

1) This has accounted for the impact on weighted-average outstanding shares for shares repurchased up to date.

2) Cash expenditures may be higher or lower than currently estimated due to a variety of factors and circumstances, including as a result of the Company's ongoing stock repurchase program, or other expenditures outside the ordinary course of business, including M&A. This number further assumes receipt of approximately \$106.0 million in tax refunds prior to December 31, 2026, which have been delayed as a result of constrained resources at the IRS; and assumes capital purchases of \$12.0 million, no further repurchases under the stock repurchase program, and spend on the therapeutic development business of \$26.0 million. The timing and amount of any future repurchases pursuant to the stock repurchase program will depend on a variety of factors, including the market price of Fulgent's common stock, general market and economic conditions and other factors Fulgent's board of directors may deem relevant.

Note: All figures are rounded.

Balance Sheet

(\$ in millions)	Periods Ended	
	December 31, 2025	June 30, 2026
Assets		
Cash & cash equivalents	\$ 50.2	\$ 26.3 ⁽¹⁾
Marketable securities	285.9	230.7 ⁽¹⁾
Trade accounts receivable, net	84.8	64.9
Prepaid income taxes	107.1	110.6
Other current assets	22.5	22.2
Total current assets	550.5	454.7
Marketable securities, long-term	369.3	294.3 ⁽¹⁾
In-process research & development	68.5	68.5
Other intangible assets, net	64.8	80.8
Fixed assets, net	112.5	113.1
Goodwill	25.1	53.9
Other long-term assets	22.8	23.9 ⁽¹⁾
Total assets	\$ 1,213.5	\$ 1,089.2
Liabilities and Stockholders' Equity		
Accounts payable	\$ 18.7	\$ 15.6
Contract liabilities	3.6	3.4
Customer deposit	28.1	28.8
Other liabilities	56.4	45.4
Total liabilities	106.8	93.2
Stockholders' equity	574.5	524.1
Accumulated income	537.5	477.5
Total Fulgent stockholders' equity	1,112.0	1,001.6
Noncontrolling interest	(5.3)	(5.6)
Total stockholders' equity	1,106.7	996.0
Total liabilities and stockholders' equity	\$ 1,213.5	\$ 1,089.2

(1) \$551.5M in cash and investments including \$0.1M of restricted cash included in Other long-term assets.

Appendix





Prenatal Screening for Genetic Conditions

- NGS Comprehensive NIPS utilizing coordinative allele-aware target enrichment (COATE) suppresses allelic hybridization bias
- Dual end sequencing retains cfDNA fragmentation characteristics
- Multi-dimensional analyses for allelic ratios, read-depth, cfDNA fragmentation pattern

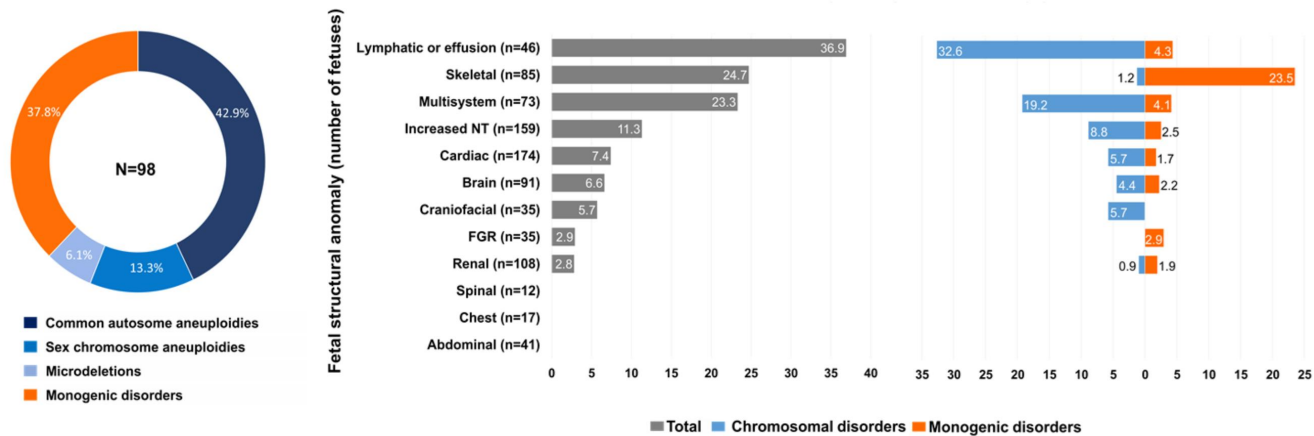


KNOVA technology is using features from both commonly used methods of NIPT (SNP-based and MPSS/counting methods). Additionally, we use proprietary technology that helps us better differentiate between maternal and fetal DNA. All of this increases the sensitivity and specificity of our test for both aneuploidies and monogenic conditions.

Fulgent NIPT/NIPS – Full Panel

Aneuploidies - 6	13, 15, 16, 18, 21, 22
Aneuploidies (sex chr)	Monosomy X (Turner), XXY (Klinefelter), XXX (Triple X), XYY (Jacob)
Microdeletions - 12	1p36; 2q33.1; 4p16; 5p15; 8q23; 9p; 11q23-25; 15q11.2-q13; 17p11.2; 18q; 18p; 22q11.2
Single genes - 56	ASXL1, BRAF, CBL, CD96, CDKL5, CHD7, COL10A1, COL11A1, COL1A1, COL1A2, COL2A1, EBP, EFNB1, ERF, FGFR1, FGFR2, FGFR3, FLNB, FREM1, GLI3, HDAC8, HNRNPK, HRAS, KAT6B, KMT2D, KRAS, LMNA, MAP2K1, MAP2K2, MECP2, NIPBL, NRAS, NSD1, NSDHL, PTPN11, RAD21, RAF1, RIT1, RUNX2, SHOC2, SKI, SLC25A24, SMC1A, SMC3, SNRPB, SOS1, SOS2, SOX9, SPECC1L, STAT3, TCF12, TRAF7, TSC1, TSC2, TWIST1, ZIC1

Detection Rates of KNOVA in High-Risk Pregnancies



The detection rate was increased by **60.7%** using KNOVA compared to standard NIPS in pregnancies with fetal anomalies.

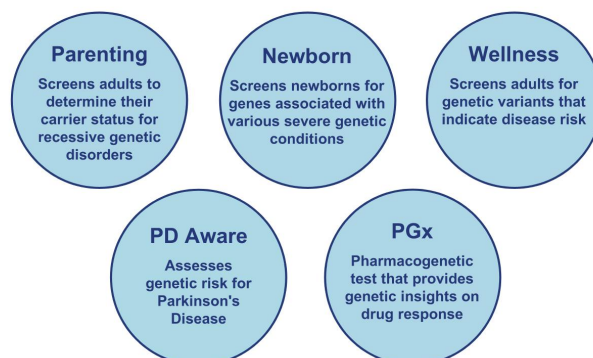
Consumer Initiated Tests – Picture Genetics

Targeting the Large Consumer Market with Picture Genetics

Launched in 2019 with significant growth amid COVID-19

- A consumer-focused offering that merges clinical utility with accuracy of an accredited lab
- Extends Fulgent's NGS capabilities to a broader market
- Validated by **successfully scaling to hundreds of thousands of tests** performed within months for COVID-19, after receiving an EUA
- Genetic tests utilizes complete sequencing (vs genotyping) by NGS analysis for better, more accurate results
- Patient-friendly with easy to use "order from home" model – no doctor office visits or insurance necessary, though many tests are eligible for reimbursement
- Select full service offering that includes analysis and genetic counseling support

Picture™
By Fulgent Genetics



Non-GAAP Financial Adjustments



(\$ in 000's)	2025				FY 2025	2026		YTD 2026
	Q1	Q2	Q3	Q4		Q1	Q2	
Revenue	\$73,463	\$81,803	\$84,069	\$ 83,336	\$322,671	\$ 71,138	\$ 85,389	\$156,527
Cost of revenue	45,117	47,368	48,557	50,754	191,796	49,648	59,710	109,358
Gross profit	28,346	34,435	35,512	32,582	130,875	21,490	25,679	47,169
Gross margin	38.6%	42.1%	42.2%	39.1%	40.6%	30.2%	30.1%	30.1%
Equity-based compensation included in cost of revenue	1,780	1,737	1,697	1,613	6,827	1,461	1,085	2,546
Non-GAAP gross profit (excluding equity-based compensation)	30,126	36,172	37,209	34,195	137,702	22,951	26,764	49,715
Non-GAAP gross margin	41.0%	44.2%	44.3%	41.0%	42.7%	32.3%	31.3%	31.8%
Operating expenses								
Research and development	12,395	13,480	13,860	14,170	53,905	14,176	14,554	28,730
Selling and marketing	8,465	12,286	11,642	10,978	43,371	12,221	15,228	27,449
General and administrative	25,291	26,392	23,335	41,646	116,664	27,684	27,195	54,879
Amortization of intangible assets	1,990	1,990	2,025	2,026	8,031	2,031	2,603	4,634
Impairment of intangible assets	—	—	—	—	—	—	2,172	2,172
Total operating expenses	48,141	54,148	50,862	68,820	221,971	56,112	61,752	117,864
Operating loss	(19,795)	(19,713)	(15,350)	(36,238)	(91,096)	(34,622)	(36,073)	(70,695)
Operating margin	-27.0%	-24.1%	-18.3%	-43.5%	-28.2%	-48.7%	-42.3%	-45.2%
Equity-based compensation included in operating expenses	8,770	8,302	8,020	7,663	32,755	8,429	7,539	15,968
Acquisition-related cost included in general and administrative ¹	—	297	90	1,537	1,924	2,632	17	2,649
Acquisition-related severance included in general and administrative	—	—	—	—	—	396	251	647
Professional liability expense included in general and administrative	—	—	—	14,500	14,500	—	—	—
Non-GAAP operating loss (excluding equity-based compensation, acquisition-related cost, acquisition-related severance, professional liability expenses, impairment loss and amortization)	\$ (7,255)	\$ (7,387)	\$ (3,518)	\$ (8,899)	\$ (27,059)	\$ (19,673)	\$ (22,406)	\$ (42,079)
Non-GAAP operating margin	-9.9%	-9.0%	-4.2%	-10.7%	-8.4%	-27.7%	-26.2%	-26.9%

1) Acquisition-related costs incurred in Q2 2025 were adjusted as the acquisition of ANP Technologies, Inc. was signed and closed during Q3 2025. Prior reporting included these costs in general and administrative operating expenses but did not remove them for Non-GAAP reporting.

Non-GAAP Financial Adjustments by Segment

(\$ in 000's)	2026		
	Laboratory Services	Therapeutic Development	Total
Revenue	\$ 156,410	\$ 117	\$ 156,527
Cost of revenue	109,343	15	109,358
Gross profit	47,067	102	47,169
Equity-based compensation included in cost of revenue	2,546	—	2,546
Non-GAAP cost of revenue (excluding equity-based compensation)	106,797	15	106,812
Non-GAAP gross profit (excluding equity-based compensation)	49,613	102	49,715
Operating expenses			
Research and development	17,037	11,693	28,730
Equity-based compensation included in research and development	3,753	2,920	6,673
Non-GAAP research and development (excluding equity-based compensation)	13,284	8,773	22,057
Selling and marketing	27,416	33	27,449
Equity-based compensation included in selling and marketing	1,177	33	1,210
Non-GAAP selling and marketing (excluding equity-based compensation)	26,239	—	26,239
General and administrative	52,711	2,168	54,879
Equity-based compensation included in general and administrative	6,666	1,419	8,085
Acquisition-related cost in general and administrative	2,649	—	2,649
Acquisition-related severance in general and administrative	647	—	647
Non-GAAP general and administrative (excluding equity-based compensation, acquisition-related cost, and acquisition-related severance)	42,749	749	43,498
Impairment of intangible assets	—	2,172	2,172
Amortization of intangible assets	4,570	64	4,634
Total operating expenses	101,734	16,130	117,864
Non-GAAP operating expenses (excluding equity-based compensation, acquisition-related cost, acquisition-related severance, impairment loss, and amortization)	82,272	9,522	91,794
Operating loss	(54,667)	(16,028)	(70,695)
Non-GAAP operating loss (excluding equity-based compensation, acquisition-related cost, acquisition-related severance, impairment loss, and amortization)	(32,659)	(9,420)	(42,079)
Other income (expenses)	15,027	—	15,027
Income (loss) before income tax	\$ (39,640)	\$ (16,028)	\$ (55,668)
Non-GAAP income (loss) before income tax (excluding equity-based compensation, acquisition-related cost, acquisition-related severance, impairment loss, and amortization)	\$ (17,632)	\$ (9,420)	\$ (27,052)

Thank You





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