



August 2, 2024

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#### **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.

# Leadership Team



Mina 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 postdoctoral 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.



Dr. Lawrence Weiss Chief Medical Officer

Esteemed background in

Most recently Chief Medical

Laboratory, Inc.; prior senior

Officer at NeoGenomics

role at Clarient Inc.

Chairman Emeritus of

Pathology at City of Hope

National Medical Center

molecular science and

pathology



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



















(NEO







# 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

### Strategic Vision – A One-Stop Solution for Cancer Care



To build a vertically integrated solution to combat cancer early detection | clinical diagnostics | post treatment monitoring | drug discovery and cancer treatment







#### **FULGENT PHARMA**

Leading Genetic Testing Company Offering Tech-Enabled Diagnostic Solutions

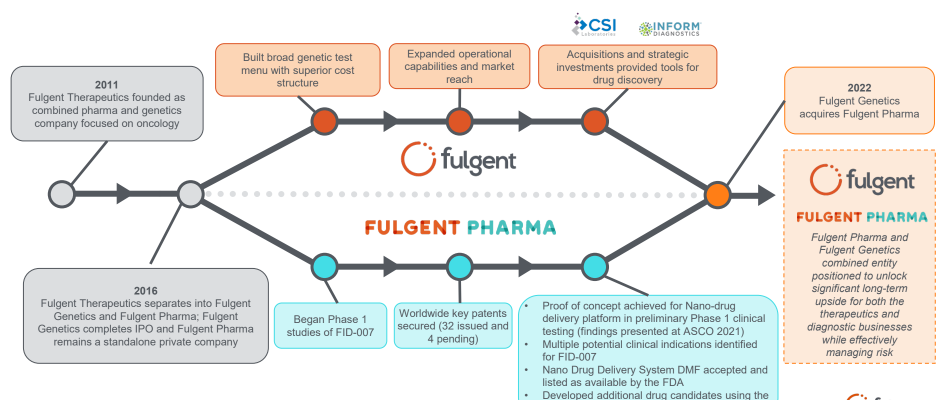
Nano-Drug **Delivery Platform**  Exciting Cancer Therapeutic Opportunity Realizing Precision Medicine Potential

#### Therapeutic and Diagnostic Entity Providing Comprehensive Solutions Across the Cancer Care Continuum

- Vertically integrated "one-stop" solution across the healthcare chain following the CSI, Inform Diagnostics, and Pharma acquisitions
- Proprietary nano drug delivery technology platform serves as an underpinning technology between diagnostic and pharm to help create a more sustainable and profitable business model in precision medicine for years to come
- Addition of a talented scientific team creates a strong synergy and competitive advantage that may be leveraged across the combined business
- Potential near-term opportunity includes shortened 505(b)(2) drug development and commercialization timelines and potential long-term opportunity leverages large data insights and novel analytical tools from diagnostics business to enable additional precision medicine pipeline through organic or partnered development strategies
- Commitment to continue growing diagnostic and therapeutic opportunities through organic investments and M&A
- Seasoned management team along with strong cash position allow Fulgent to enter therapeutic opportunities while managing risk



# History of Fulgent



same drug delivery platform

# Long-Term Vision: Fulgent Continuum of Care

Diagnosis

Therapy

Database + Drug Discovery + Patient Care



. .

- Transforms from a service company to a combined diagnostic and therapeutic company focused on precision medicine
- Large oncology market opportunity
- Careful pipeline management will responsibly engage therapeutic opportunities while managing potential risks
- Valuable, diverse assets and future sustainable revenue and margins



#### **FULGENT PHARMA**

- Well-funded by diagnostics to advance H&N therapy trials and commercialization process
- Genomics data accelerates pipeline therapeutics development, e.g. spatial biology for tumor microenvironment profiling
- Long-term opportunity to leverage data insights from diagnostics to enable precision medicine through proprietary or partnered development strategies
- Manufacturing capability to aid drug development

# LABORATORY SERVICES



+5%

**Q2** Year-over-Year Core Revenue Increase

18,400+ GENES | 900+ PANELS CUSTOMIZABLE OFFERINGS

### Positioned for Growth

- Proprietary technology platform allows for rapid scaling of a **broad**, **flexible test menu**
- Next-generation sequencing (NGS) platform complemented with growing portfolio of emerging testing technologies with a focus on oncology
- Well-positioned to execute on a growth strategy that includes organic and inorganic initiatives, including:
  - Transformational acquisition of Inform Diagnostics
  - Ramping of CSI Labs
  - Scaling partnerships
  - Potential future acquisitions with a strategy of short- and longterm ROI, tangible synergies, and efficient capital deployment



# Platform and Capabilities Across 3 Divisions



**Laboratory Services** 

### **Precision Diagnostics**

- **Reproductive Health**
- **Oncology / Liquid Biopsy**
- Rare Disease
- **Neurogenetics**



### **Anatomic Pathology**

- **Dermatopathology**
- GI
- GU
- **GSP**



### **BioPharma Services**

- **Spatial Phenotyping**
- **Exome/Genome sequencing**
- **RNA** sequencing
- Single Cell sequencing

# Target Market Opportunity



**Cancer Diagnostics** \$80B market<sup>1</sup>

**Early Detection / Liquid Biopsy** 

\$18B market

**Reproductive Health** \$8B market<sup>2</sup>

**BioPharma Services** \$50B market<sup>3</sup>

Market sizes sourced from Wall Street equity research

Market size sourced from Frost & Sullivan, October 2022 Market size sourced from Research and Markets, April 2022

# What Sets Fulgent Diagnostics Apart?

### Leads to a Broader Test Menu

- 18,400+ single-gene tests (1)
- 900+ panels
- Whole Genome and Exome
- Flexibility enables custom tests for any genes or conditions
- Preset panels have grown 350% since IPO in 2016

#### **And a Better Cost Structure**

- Lab efficiencies, automation and scale have translated to a sustainable cost structure
- Partnerships create leverage with sales and marketing
- Process 100% of volume without the need for outsourcing



#### Comparison and suppression algorithms

A New Approach to NGS

 Proprietary probes and engineered chemistry

 Comprehensive analytics powered by AI and ML

**Superior Cost Structure** 

**Extensive Test Menu** 

**Technology Platform** 



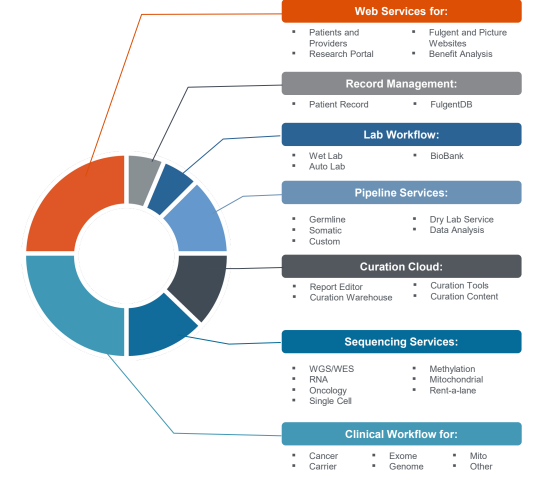
### Proprietary Technology Platform

### **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





# **Broad Capabilities**



Next Generation Sequencing Opportunities

#### **Recent Traction with:**

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

Newly launched pharmacogenetic test

**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
- thology Urologic pathology
- Gastrointestinal pathology
- Neuropathology
- Dermatopathology

Hematopathology

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

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



# Technology Platform Case Study: COVID-19

Fulgent deployed its technology platform to rapidly respond to the COVID-19 Pandemic, scaling operations to provide tests with reliable results and rapid turnaround time



### Next Generation Sequencing for COVID-19

- Research driven platform worked with local and federal government on genomic studies
- CDC contract awarded to Fulgent, worth up to \$47M to study SARS-CoV-2 using Fulgent's NGS platform
- Capacity of 10,000 NGS tests per day
- Used to identify new strains and mutations



### Commercialized COVID-19 Testing Primarily RT-PCR Based Testing

#### **Contracts with:**

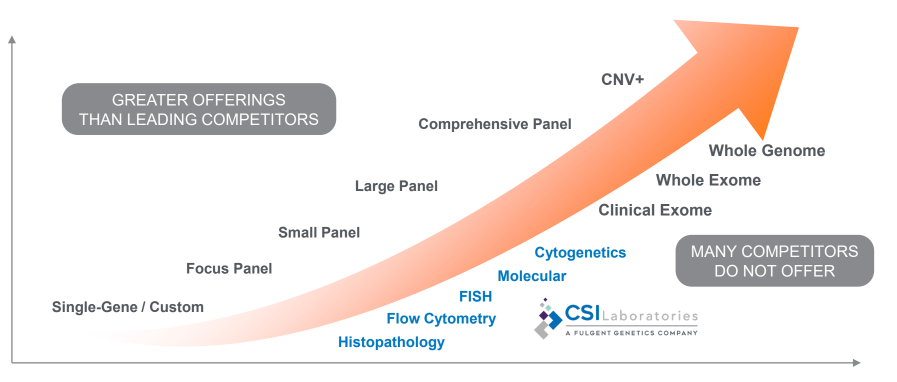
- School systems
- Nursing homes
- Athletic organizations
- Specialty health clinics
- Travel organizations
- Government agencies

#### Offered through:

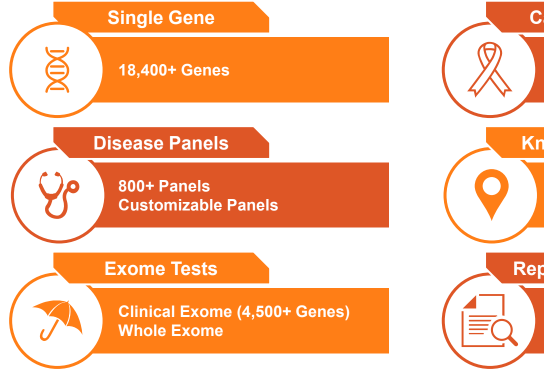
- Drive-through sites
- Picture at-home kits
- Managed on-site programs

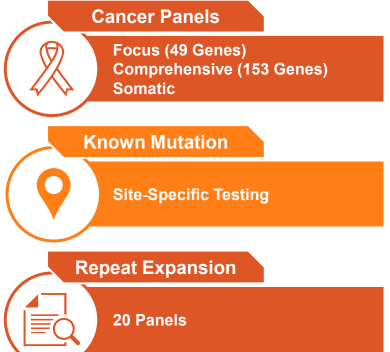
Result: Over 19M COVID-19 tests delivered between 2020-2023, generating >\$1.7B in revenue for Fulgent

### Scalable and Affordable Menu for Customers

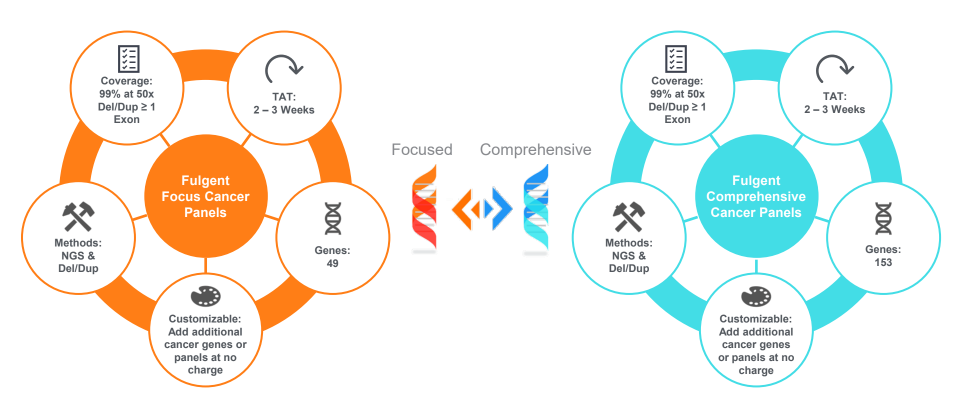


# NGS Testing – Offerings





# NGS Testing – Germline Oncology Test Menu



# Oncology Testing Platforms



### **FISH**

- Expansive heme and solid tumor menu
- STAT testing available
   PML/RARA <1 day</li>
   TAT
- 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 Tcell 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 8-10 days]



# NGS Testing – Panel Deep Dive

### Comprehensive Beacon Carrier Screening Tests

Beacon ACOG / ACMG Guidelines Panel

The ACOG/ACMG panel screens for common genetic disorders seen in the general population

Gene Count: 6

Beacon Ashkenazi Jewish Panel The Ashkenazi Jewish carrier screens for pathogenic carrier variants known to cause recessive genetic disorders

Gene Count: 61

Beacon Focus Panel

The Focus Carrier screen is a pan-ethnic screen that looks for pathogenic mutations known to cause autosomal recessive and X-linked disorders

Gene Count: 30



Beacon Expanded Panel The Expanded Panel screens for more than 400 recessive and X-linked conditions that covers people of all ethnic backgrounds

Gene Count: 427



Beacon787 Panel One of the largest panels available for those seeking the most comprehensive testing option

Gene Count: 787



### **Beacon Carrier Screening**

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



# NGS Testing – Reproductive Services: PGT-A

### PGT-A Can Expand a Patient's Prospects of a Successful Pregnancy



Embryos with a normal number of chromosomes have a better chance of resulting in successful pregnancy

**Pregnancy** 

### Reduced Risk of Miscarriage

Embryos with an abnormal number of chromosomes (aneuploid) typically do not result in successful pregnancy or may result in birth defects





More Confidence in Transferring a Single Embryo

Avoid health risks associated with twin or triplet pregnancies that can occur from multiple embryo implantation



Reduce the amount of time to pregnancy and the costs of additional IVF cycles



Preimplantation Genetic Testing for Aneuploidy (PGT-A) can identify potentially abnormal embryos for transfer in IVF, thereby expanding a patient's prospects of a successful pregnancy

Who is PGT-A testing for?

Women 35+

Those who have experienced miscarriages

Those who want to reduce the likelihood of having multiples

Couples experiencing male factor infertility

Those who have experienced IVF failure



# NGS Testing – Rapid Whole Genome

Designed for critically ill infants in the NICU/PICU to rapidly diagnose genetic disorders Covers >4.000 Focused reporting single gene Fast turn around of diagnostic findings only (7-10 days) Ideal for Infants Experiencing: Multiple congenital Inborn errors of metabolism Immunodeficiency Respiratory distress Epilepsy anomalies In a Retrospective Analysis of Diagnostic and Clinical Finding with 35 Acutely III Infants (2015): 13 out of the 20 diagnosed infants (65%) had clinical 20 out of the 35 infants (57%) received a diagnosis usefulness for treatment TAT of 7-10 Days

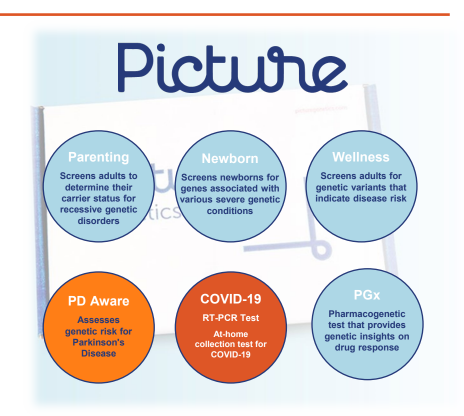
### Consumer Initiated Tests – Picture Genetics

# Targeting the Large Consumer Market with Picture Genetics

Launched in 2019 with significant growth amid COVID-19

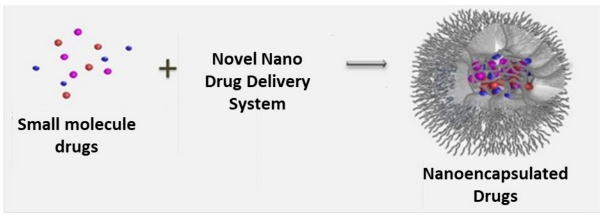
- A consumer-focused offering that merges clinical utility with accuracy of an accredited lab
- Extends Fulgent's NGS capabilities to a broader market
- Validated by successfully scaling to hundreds of thousands of tests performed within months for COVID-19, after receiving an EUA
- Genetic tests utilizes 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



# THERAPEUTIC DEVELOPMENT

# Nano-Drug Delivery Platform Overview



### **Platform Advantage:**

Soluble in both water and various organic solvents and capable of hot melt mixing with APIs

- Many drug candidates in the industry failed during preclinical and clinical development and testing due to poor water solubility
- Nanoencapsulation produces amorphous drug candidates with improved solubility and potentially enhanced absorption, drug PK profiles, safety and efficacy
- · Broadly applicable to both IV and oral drug delivery formulations
- Potentially shortened development timeline
- Plug and play drug delivery platform provides multiple shots on goal
- Simple and low-cost production process



# FID-007 Program Overview

### FID-007 Phase 1/1b First in Human Clinical Trial – Preliminary Findings (n=46 patients)

- Dose levels up to 160 mg/m²/week with manageable safety profile
  - RP2D at 125 mg/m<sup>2</sup>/week
- There is preliminary evidence of anti-tumor activity in 46 heavily pre-treated patients across different tumor types (ORR = 17%)
- No high-grade neuropathy often seen in other taxanes
- Updated clinical data presented at ASCO 2024

### FID-007 Phase 1/1b Preliminary Highlights (as of 6/2/24):

#### **H&N Cancer**

 45% ORR and 72% DCR were observed in 11 heavily treated HNSCC patients. Among them, 3 out of the 5 patients who achieved a PR had received prior taxane.

### FID-007 Plus Cetuximab Phase 2 Update (as of 7/30/24):

#### **H&N Cancer**

Multiple clinical sites activated (USC, Moffitt, etc.) with 3
 Patients enrolled and dosed

# Abstract # 6042: Efficacy from the phase 1 study of FID-007, a novel nanoparticle paclitaxel formulation, in patients with head and neck squamous cell carcinoma



Lydia Chow<sup>1</sup>, Robert Hsu<sup>1</sup>, Jorge Nieva<sup>1</sup>, Denice Tsao-Wei<sup>1</sup>, Ming Hsieh<sup>2</sup>, Ray Yin<sup>2</sup>, Anthony El-Khoueiry<sup>1</sup>, Jacob Thomas<sup>1</sup>.

<sup>1</sup>University of Southern California, Norris Comprehensive Cancer Center; <sup>2</sup>Fulgent Pharma. Contact: Jacob Thomas@med.usc.edu



### FID-007 Clinical Data Presented at ASCO 2024

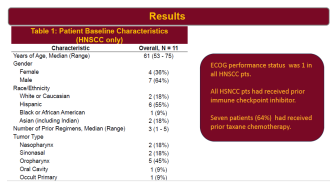


Table 2: Treatment-related select AE categories (>= 10%) (All patients)						
Toxicity	Number Of Patients With Maximum Grade Toxicity Experienced (N=46)					
	Grade 1 or 2	Grade 3	Grade 4			
Alopecia	24 (52%)	0	0			
Pruritus	20 (43%)	0	0			
Rash maculo-papular	17 (37%)	16 (35%)	0			
Fatigue	17 (37%)	0	0			
Nausea	13 (28%)	0	0			
White blood cell decreased	12 (26%)	6 (13%)	3 (7%)			
Anorexia	12 (26%)	1 (2%)	0			
Neutrophil count decreased	10 (22%)	3 (7%)	6 (13%)			
Dry skin	10 (22%)	1 (2%)	0			
Dysgeusia	10 (22%)	0	0			
Anemia	9 (20%)	8 (17%)	0			
Peripheral sensory neuropathy	9 (20%)	0	0			
Palmar-plantar erythrodysesthesia syndrome	9 (20%)	0	0			
Constipation	6 (13%)	0	0			
Vomiting	6 (13%)	0	0			
Diarrhea	6 (13%)	0	0			

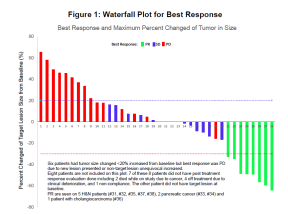
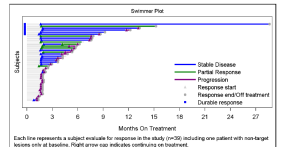


Figure 2: Swimmer Plot for Responses over Time



A durable responder is a patient whose response>6 months.

Table 3: Tumor Responses and Outcomes						
Characteristic	Overall, N = 46	HNSCC, N = 11				
Total Courses Completed, Median (Range)	2 (1 - 30)	5 (2-16)				
Best Response*						
PR .	8 (17%)	5 (45%)				
SD	16 (35%)	3 (27%)				
PD	21 (46%)a	3 (27%)				
Inevaluable	1 (2%)	0 (0%)				
Duration of Follow-up (Months), Median (Range)	12.1 (1.1, 45.9)	4.0 (1.0-15.0)				

- a PD includes 4 patients who had clinical deteriorations prior to RECIST evaluation.
- One patient with inevaluable response; off-treatment due to non-compliance. No response evaluation was performed.

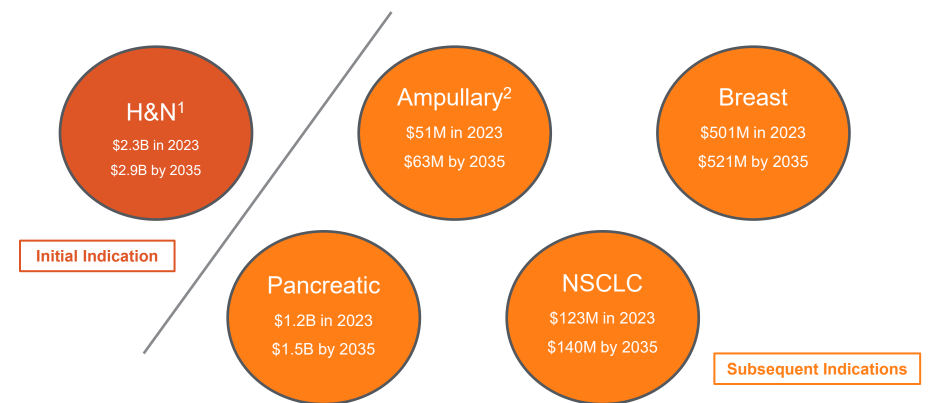


#### **Conclusions**

- FID demonstrates preliminary evidence of anti-tumor activity in heavily pretreated HNSCC pts across different primary tumor sites, with an ORR 45%.
- 3 out of the 5 patients who achieved a PR had received prior taxane.
- There has been no grade 3 or higher peripheral neuropathy.
- Phase 2 study of FID combination with cetuximab in pts with HNSCC has begun enrollment.



# Potential Market Opportunity for FID-007



Note: U.S. opportunity shown

Sources: Evaluate Pharma, Wall Street research, and management pricing expectations

H&N market opportunity for both 2<sup>nd</sup> line and 3<sup>rd</sup> line therapy
 Ampullary market opportunity for 2<sup>nd</sup> line therapy

# Pipeline Progress

- FID-007: wholly-owned drug candidate initially focused on Head & Neck (H&N), Pancreatic/Ampullary cancers
  - Seeking initial therapeutic indication for 2<sup>nd</sup> line treatment of H&N cancer
  - Potential FDA approval strategy uses 505(b)(2) studies, which may shorten clinical trial process and accelerate timeline to commercialization
- FID-022 moving toward IND
- Developing a next generation antibody drug conjugate (ADC) technology platform that could potentially provide even broader killings towards heterogeneous cancer cells than those ADCs with the bystander killing effect

Drug Candidates	Target	Indication	Pre-Clinical	Clinical P1	Clinical P2	Clinical P3	Milestones
FID-007	Cytotoxic	Head and Neck (H&N) (505(b)(2))					Began P2 Enrollment in 2Q24
Cytotoxic		Ampullary or ICI Resistant (505(b)(2))					Go/No-go Based on HN Study
FID-022	Cytotoxic	Colon (505(b)(2))					IND Filing by YE24
ADCs	Undisclosed	Solid Tumors					

# FINANCIALS

# Summary Financial Performance

\$70M Core Revenue<sup>1</sup> in Q2'24 5% growth year-over-year

\$37M Last Twelve Months (LTM) Operating Cash Flow as of Q2'24





### Financial Performance: Revenue Profile



### 2024 Financial Guidance

Metric	Full Year 2024	Expected Revenue Breakdown			
Core Revenue	\$280M	Precision Diagnostics	\$173M		
Core Revenue	+7% y/y <sup>1</sup>	Anatomic Pathology	\$96M		
GAAP EPS	(\$1.95) <sup>2</sup>	BioPharma Services	\$11M		
Non-GAAP EPS	on-GAAP EPS (\$0.30) <sup>2</sup>		\$280M		

Expected Cash, cash equivalents, and investments in marketable securities of approximately \$800 million as of December 31, 2024<sup>3</sup>

Cash expenditures may be higher or lower than currently estimated due to a variety of facts and circumstances, including as a result of the Company's ongoing stock repurchase program or other expenditures outside of ordinary course.



Core Revenue excludes NGS COVID-19 test revenue for more accurate year over year comparison purposes.

Improvements from prior guidance of (\$2.25) and (\$1.05), respectively

# Balance Sheet

(in 000's)	Periods Ended				
	Dece	mber 31, 2023	June 30, 2024		
Assets					
Cash & cash equivalents	\$	97,473	\$	65,111 <sup>(1</sup>	
Marketable securities		326,681		246,595 <sup>(1</sup>	
Trade accounts receivable, net		51,132		56,573	
Other current assets		32,559		30,825	
Total current assets		507,845		399,104	
Marketable securities, long-term		423,571		526,163 <sup>(1</sup>	
Intangible assets, net		143,053		138,973	
Fixed assets, net		83,464		93,368	
Goodwill, net		22,055		22,055	
Redeemable preferred stock investment		20,438		20,438	
Other long-term assets		34,902		32,138	
Total assets	\$	1,235,328	\$	1,232,239	
Liabilities and Stockholders' Equity					
Accounts payable	\$	15,360	\$	19,873	
Contract liabilities		2,874		2,744	
Customer deposit		22,700		26,297	
Other liabilities		61,108		54,477	
Total liabilities		102,042		103,391	
Stockholders' equity		501,721		522,423	
Accumulated income		634,380		610,126	
Total Fulgent stockholders' equity		1,136,101		1,132,549	
Noncontrolling interest		(2,815)		(3,701)	
Total stockholders' equity		1,133,286		1,128,848	
Total liabilities and stockholders' equity	\$	1,235,328	\$	1,232,239	
(1) \$838M in cash and investments.					

# Non-GAAP Financial Adjustments

(in 000's)	2023			FY 2024			
	Q1	Q2	Q3	Q4	2023	Q1	Q2
Revenue	\$66,168	\$67,853	\$84,687	\$70,505	\$289,213	\$64,485	\$71,028
Cost of revenue	47,357	47,281	44,843	45,276	184,757	42,381	44,537
Gross profit	\$18,811	\$20,572	\$39,844	\$25,229	\$104,456	\$22,104	\$26,491
Gross margin	28.4%	30.3%	47.0%	35.8%	36.1%	34.3%	37.3%
Equity-based compensation included in cost of revenue	2,394	2,359	2,621	2,375	9,749	2,009	1,999
Non-GAAP gross profit (excluding equity-based compensation)	\$21,205	\$22,931	\$42,465	\$27,604	\$114,205	\$24,113	\$28,490
Non-GAAP gross margin	32.0%	33.8%	50.1%	39.2%	39.5%	37.4%	40.1%
Operating expenses							
Research and development	\$9,782	\$9,692	\$10,014	\$11,952	\$41,440	\$11,434	\$13,486
Selling and marketing	10,083	10,723	10,161	10,500	41,467	8,989	8,595
General and administrative	21,802	17,993	17,498	31,706	88,999	21,489	21,326
Amortization of intangible assets	1,968	1,962	1,957	1,958	7,845	1,990	1,990
Goodwill impairment loss	_	_	_	120,234	120,234	_	_
Total operating expenses	43,635	40,370	39,630	176,350	299,985	43,902	45,397
Operating profit (loss)	(\$24,824)	(\$19,798)	\$214	(\$151,121)	(\$195,529)	(\$21,798)	(\$18,906)
Operating margin	-37.5%	-29.2%	0.3%	-214.3%	-67.6%	-33.8%	-26.6%
Equity-based compensation included in operating expenses	7,871	7,964	8,281	9,057	33,173	9,509	9,636
Non-GAAP operating profit (loss) (excluding equity-based compensation,							
amortization and goodwill impairment)	(\$12,591)	(\$7,513)	\$13,073	(17,497)	(\$24,528)	(\$8,290)	(\$5,281)
Non-GAAP operating margin	-19.0%	-11.1%	15.4%	-24.8%	-8.5%	-12.9%	-7.4%

# THANK YOU

