



QUIDEL®

INVESTOR PRESENTATION

Q1 2020



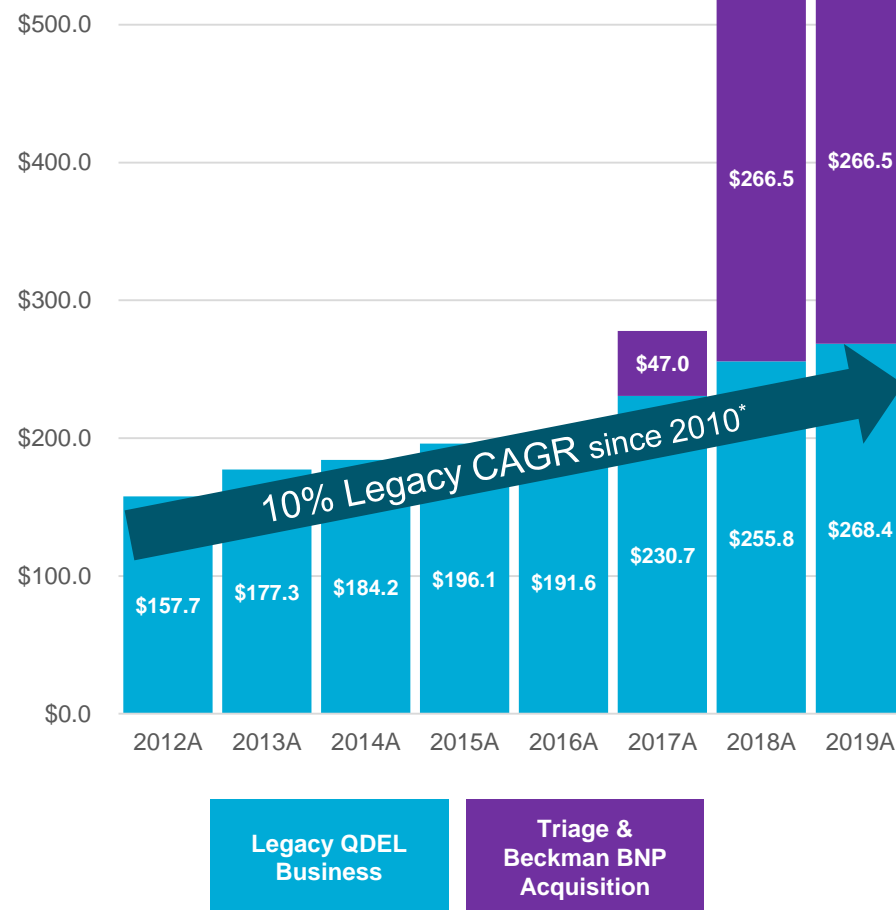
Forward-Looking Statements

This presentation contains forward-looking statements within the meaning of the federal securities laws that involve material risks, assumptions and uncertainties. Many possible events or factors could affect our future financial results and performance, such that our actual results and performance may differ materially from those that may be described or implied in the forward-looking statements. As such, no forward-looking statement can be guaranteed. Differences in actual results and performance may arise as a result of a number of factors including, without limitation, our reliance on sales of our influenza diagnostic tests; fluctuations in our operating results resulting from the timing of the onset, length and severity of cold and flu seasons, seasonality, government and media attention focused on influenza and the related potential impact on humans from novel influenza viruses, adverse changes in competitive conditions in domestic and international markets, the reimbursement system currently in place and future changes to that system, changes in economic conditions in our domestic and international markets, lower than anticipated market penetration of our products, the quantity of our product in our distributors' inventory or distribution channels, changes in the buying patterns of our distributors, and changes in the healthcare market and consolidation of our customer base; our exposure to claims and litigation, including the ongoing litigation between the Company and Beckman Coulter, Inc.; our development and protection of proprietary technology rights; our development of new technologies, products and markets; our reliance on a limited number of key distributors; intellectual property risks, including but not limited to, infringement litigation; our need for additional funds to finance our capital or operating needs; the financial soundness of our customers and suppliers; acceptance of our products among physicians and other healthcare providers; competition with other providers of diagnostic products; adverse actions or delays in new product reviews or related to currently-marketed products by the U.S. Food and Drug Administration (the "FDA") or other regulatory authorities or loss of any previously received regulatory approvals or clearances; changes in government policies; costs of or our failure to comply with government regulations in addition to FDA regulations; compliance with government regulations relating to the handling, storage and disposal of hazardous substances; third-party reimbursement policies; our failure to comply with laws and regulations relating to billing and payment for healthcare services; our ability to meet demand for our products; interruptions in our supply of raw materials; product defects; business risks not covered by insurance; our exposure to cyber-based attacks and security breaches; competition for and loss of management and key personnel; international risks, including but not limited to, compliance with product registration requirements, exposure to currency exchange fluctuations and foreign currency exchange risk sharing arrangements, longer payment cycles, lower selling prices and greater difficulty in collecting accounts receivable, reduced protection of intellectual property rights, political and economic instability, taxes, and diversion of lower priced international products into U.S. markets; changes in tax rates and exposure to additional tax liabilities or assessments; risks relating to the acquisition and integration of the Triage and BNP Businesses; Alere's failure to perform under various transition agreements relating to our acquisition of the Triage and BNP Businesses; that we may incur substantial costs to build our information technology infrastructure to transition the Triage and BNP Businesses; that we may have to write off goodwill relating to our acquisition of the Triage and BNP Businesses; our ability to manage our growth strategy; the level of our indebtedness; the amount of, and our ability to repay, renew or extend, our outstanding debt and its impact on our operations and our ability to obtain financing; that the Senior Credit Facility is secured by substantially all of our assets; our prepayment requirements under the Senior Credit Facility; the agreements for our indebtedness place operating and financial restrictions on the Company; that an event of default could trigger acceleration of our outstanding indebtedness; that we may incur additional indebtedness; increases in interest rate relating to our variable rate debt; dilution resulting from future sales of our equity; volatility in our stock price; provisions in our charter documents, Delaware law and the indenture governing our Convertible Senior Notes that might delay or impede stockholder actions with respect to business combinations or similar transactions; and our intention of not paying dividends. Forward-looking statements typically are identified by the use of terms such as "may," "will," "should," "might," "expect," "anticipate," "estimate," "plan," "intend," "goal," "project," "strategy," "future," and similar words, although some forward-looking statements are expressed differently. The risks described in reports and registration statements that we file with the Securities and Exchange Commission (the "SEC") from time to time, should be carefully considered. You are cautioned not to place undue reliance on these forward-looking statements, which reflect management's analysis only as of the date of this presentation. Except as required by law, we undertake no obligation to publicly release the results of any revision or update of these forward-looking statements, whether as a result of new information, future events or otherwise.

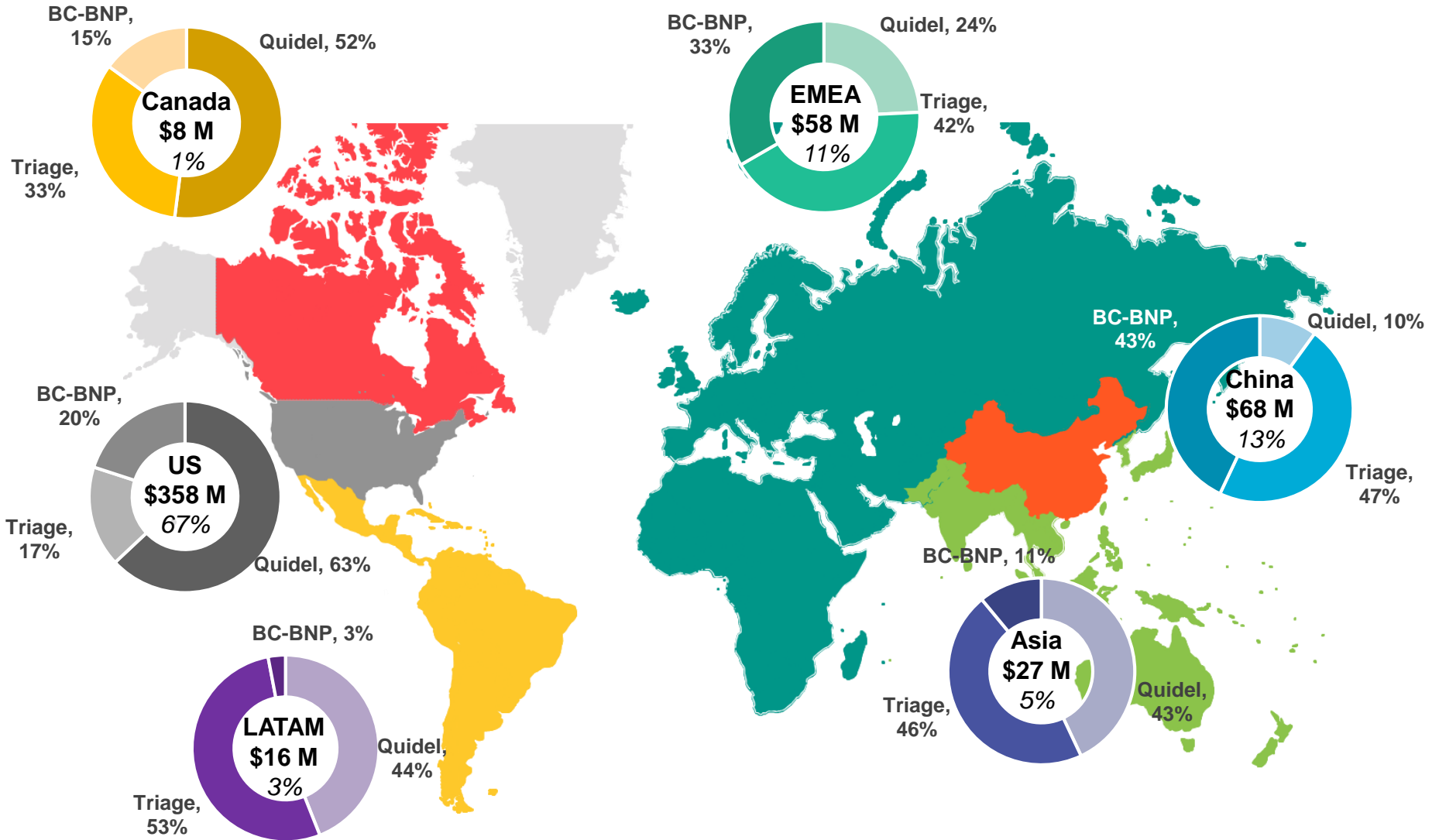
Quidel is committed to delivering strong performance by investing in growth and innovation.

2009 - Present			
Acquired Triage and BNP Businesses from Alere	Submitted over 30 510(k) packages to the FDA	Over 25 FDA clearances and expanded product claims, 4 CLIA waiver designations	
Launched Sofia 2 platform, 3 FDA-cleared assays	Launched Solana instrumented MDx platform	Launched Sofia platform and 4 FDA-cleared assays	
Launched 2 first generation molecular systems (AmpliVue & Lyra)	Launched Virena wireless system	Acquired DHI (\$45 million annual direct business)	
Acquired BioHelix (proprietary HDA technology)	Acquired Immutopics direct bone health business	Acquired RPS POC direct Eye Care businesses	Acquired ViroMed, CellPro direct Virology businesses
1989 - 2009			
Launched QuickVue Influenza A/B rapid diagnostic assay		First company to receive CLIA waiver for Flu, Strep and H. pylori tests	
1979 - 1989			
Company founded		Primarily a provider of Strep A and Pregnancy rapid diagnostics	

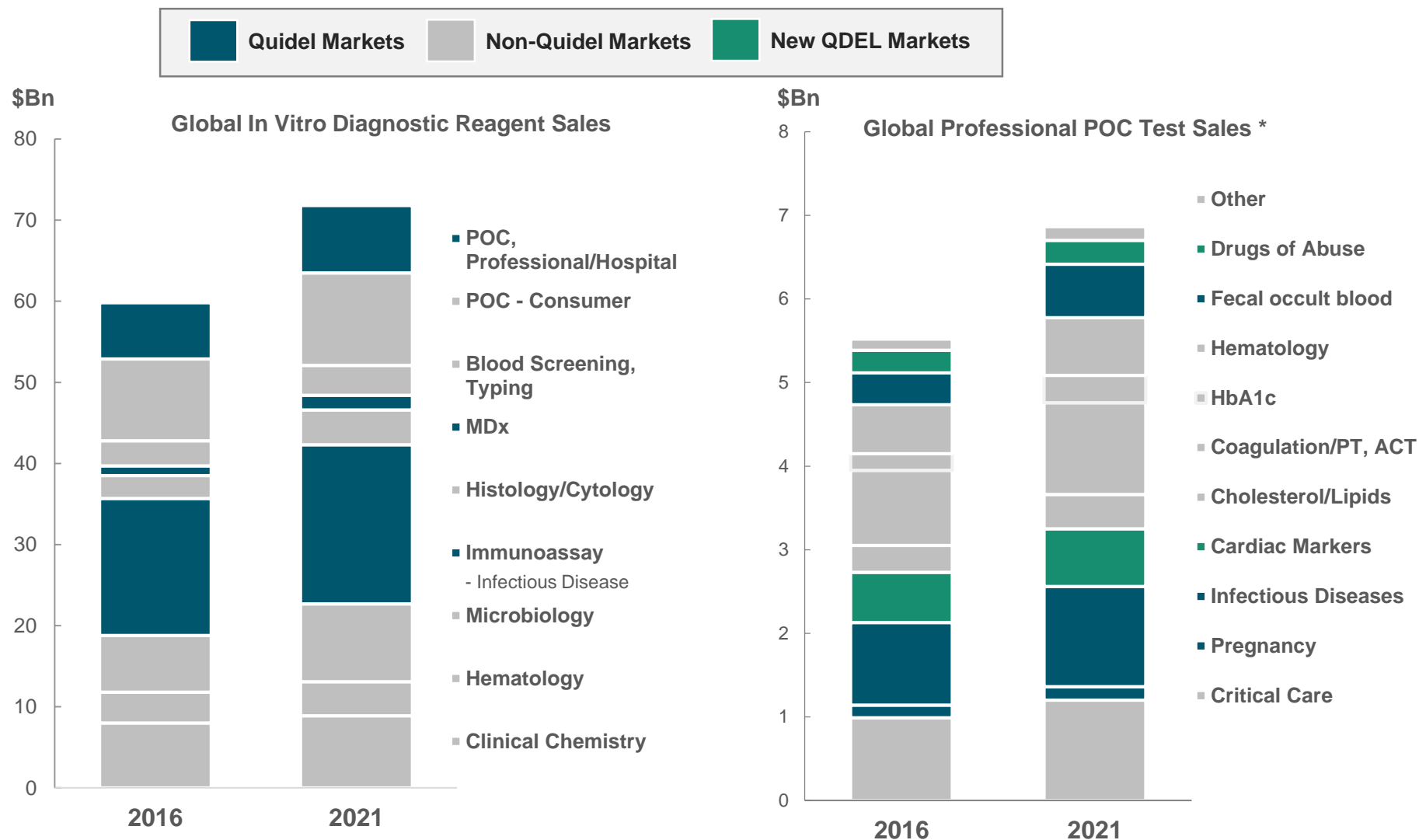
Revenues



Quidel sells products across the globe through the Quidel Network - a broad channel mix composed of direct account managers and distribution partners.

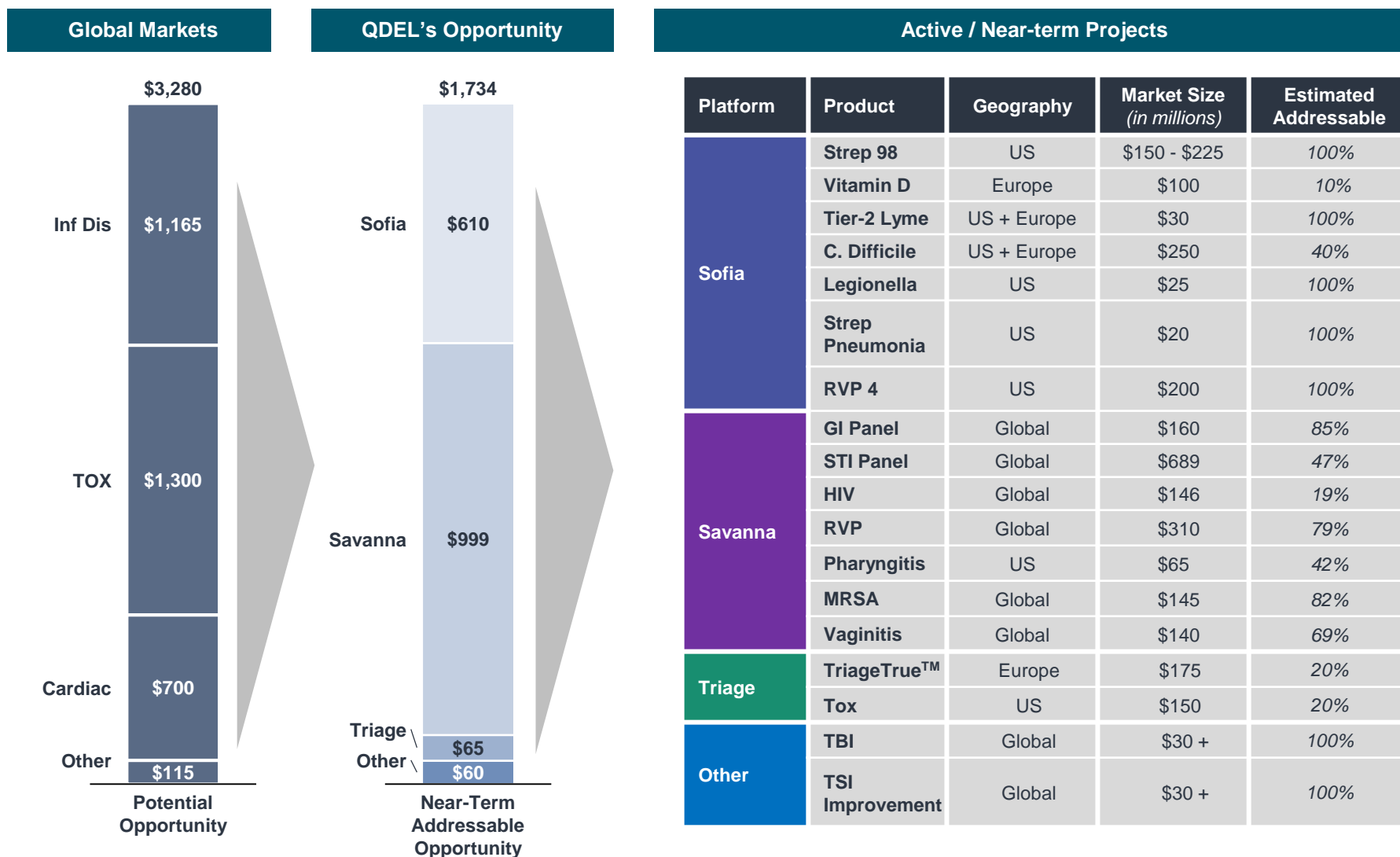


Our Cardiac acquisition has unlocked new markets.



* Does not include "New Market" category as defined by Quidel on subsequent slides

Active projects represent \$1.7B in addressable market opportunity.



Quidel's 3 Main Objectives over 3-5 years



Leverage Assets Fully:

- Sofia
- Global Infrastructure
- R&D Expertise



Play a meaningful role in the MDx space:

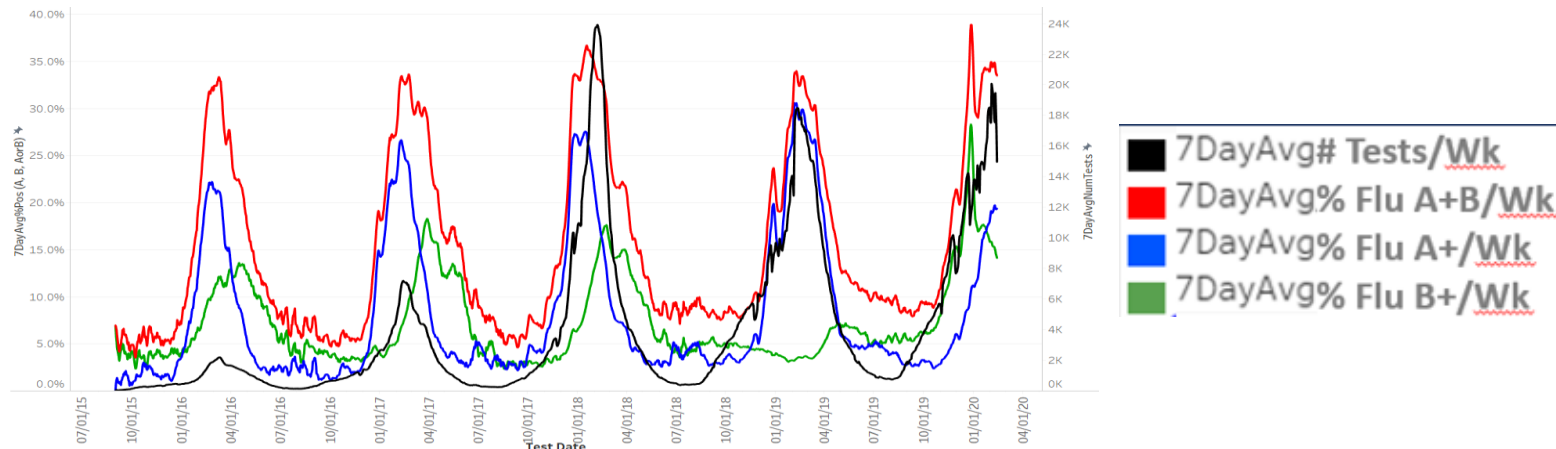
- Solana
- Savanna



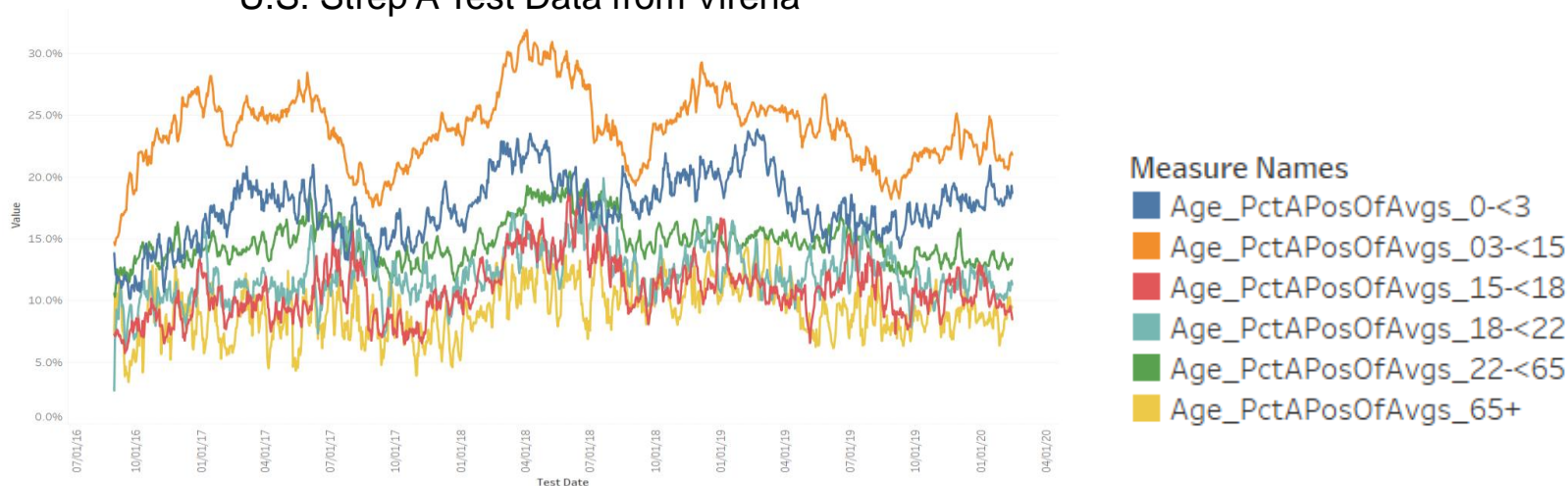
Deploy capital wisely

Influenza and Strep A

U.S. Influenza Test Data from Virena



U.S. Strep A Test Data from Virena



Our Instrumented Immunoassay systems have become flagship products, combining quality, high sensitivity, low cost and fast diagnosis.

Rapid Immunoassay



FDA-Cleared, CLIA-Waived

Sofia® Immunoassay platform is the next generation, objectively-read system in support of our legacy lateral flow business, designed to deliver more value and a higher **gross margin**.

Sofia captures market share with the first wave of assays (for Respiratory Disease) in physician offices, hospitals and alternate sites.



FDA-Cleared, CLIA-Waived

Sofia® 2, at a fraction of the cost of the original Sofia, can further penetrate the lower-volume and higher-volume segments of the POC testing market.

Sofia 2's integrated wireless options will expand connectivity with the potential to increase overall diagnostic testing.

Cardiac Immunoassay



FDA-Cleared

Triage® MeterPro® is our cost effective, easy to use instrument for cardiovascular and toxicology diagnostic assays. Multiple immunoassays can run on the same platform, with rapid results in about 15-20 minutes

The **MeterPro** instrument can run multiple sample types, such as whole blood, plasma, or urine, with process controls built into the meter, software and test device

Our innovative molecular diagnostics products are designed to meet the various needs of the customer in any POC setting.

Molecular



FDA-Cleared, Mod-Complex

AmpliVue®, a non-instrumented, hand-held disposable molecular device requires no thermocycler or upfront costs. Moderately complex claim provides an easy-to-use entry point into molecular testing for smaller hospitals and lower-volume users wishing to convert from legacy testing methods.



FDA-Cleared, Mod-Complex

Solana®, our first instrumented molecular system, extends AmpliVue's proprietary HDA technology to offer a low-cost, medium volume molecular solution that can multiplex, running up to 12 samples at a time in approx. 30 minutes.

Solana is designed for samples that do not require extraction or quantitation in the mod-complex setting.



FDA-Cleared, Highly-Complex

Lyra®, our Real-time molecular PCR assays, are designed to run on a hospital lab's existing thermocycler, and provides a real-time testing solution to higher throughput labs using a hospital's established systems – with no upfront costs.

Lyra molecular assays can be paired with other **Lyra** assays to create customized multiplex Respiratory Disease panels.



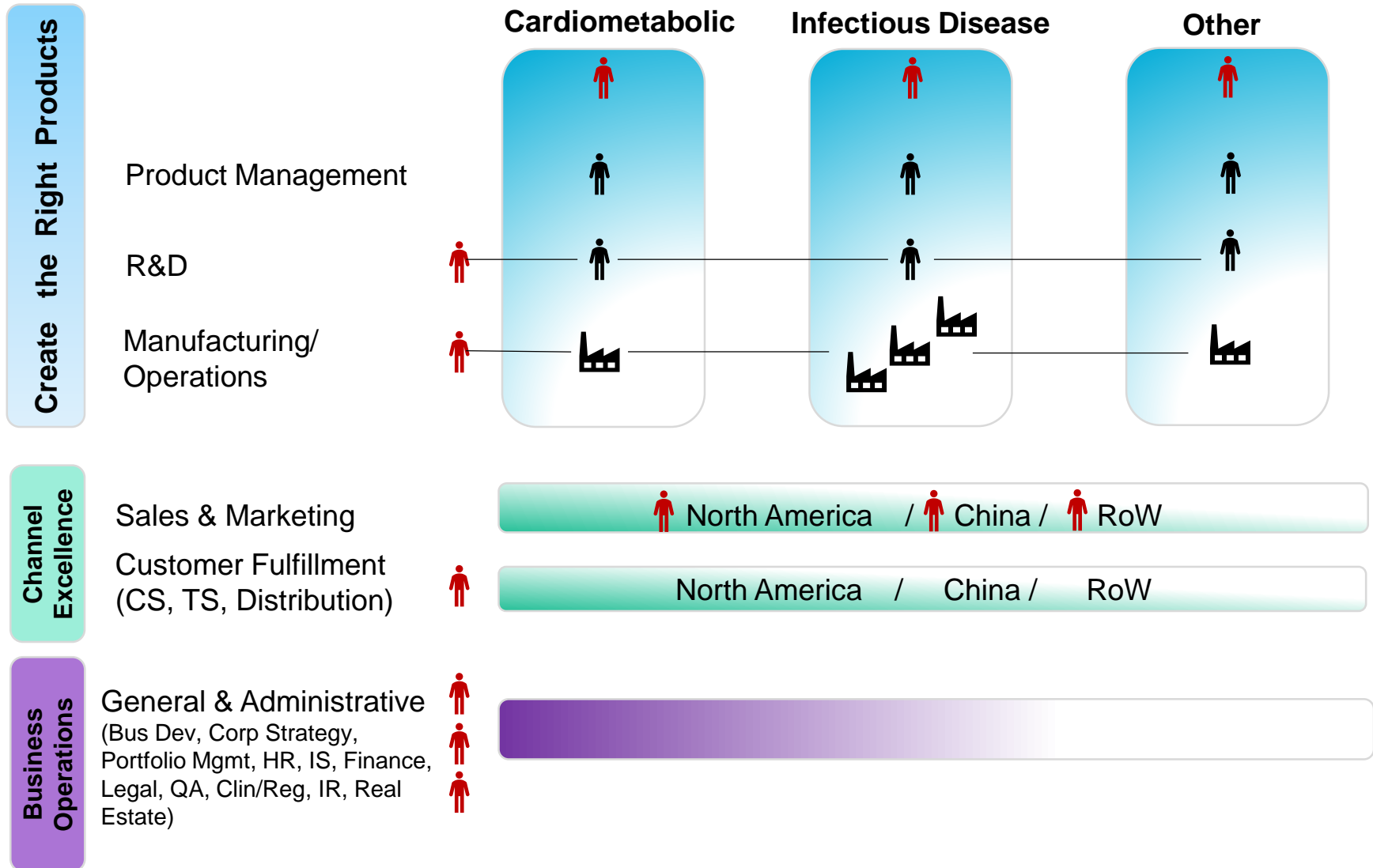
In Development

Savanna® is our low-cost, fully integrated “sample-to-answer” molecular diagnostic system.

Target cartridge cost is less than \$5 and target instrument cost is \$10,000, designed for low-volume and potentially CLIA-waived settings.

Savanna can run either traditional PCR or HDA assays for samples that do not require extraction.

Our company structure will evolve to better support growth in our key businesses



Quidel has aggressively reduced its leverage, which better positions the company for additional M&A opportunities.

Debt Type	Transaction Date October 6, 2017	Current	Actions
Convertible Bond Debt	\$167 million	\$13 million	Re-paid \$154 million through opportunistic convertible bond exchange transactions.
Credit Facility	\$255 million	\$0	Paid in full within 2 years of transaction close.
Total Debt	\$422 million	\$13 million	
Leverage Ratio	4.0X	< 1.0X	
Balance on Deferred and Contingent Consideration	\$280 million \$8 million per year x 5 years \$40 million per year x 6 years	\$184 million \$8 million per year x 3 years \$40 million per year x 4 years	Made second \$48 million payment to Abbott in April 2019.

Maximizing Shareholder Value:



Achieve 7% or higher in annual revenue growth through new product introductions



Invest in the development and commercialization of new products driving gross margins to 65%



Leverage operational efficiency driving EBITDA margins to 35%



Maximize cash flow from the core business



Pay down Abbott obligation; deploy capital wisely.



Our M&A Strategy:

Use cash on hand and access to capital to acquire assets that fit

What we're looking for...



Targets range from tuck-ins up to equal size



New technology that accelerates the advancement of in-house MDx abilities



Products that leverage our assets



Companies that can leverage our global infrastructure



Accretive within 12-24 months

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