Genetic Signatures

Investor presentation

28 May 2020

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A global leader in the supply of molecular diagnostic solutions

A specialist molecular diagnostics company



Focused on becoming a global leader in the supply of molecular diagnostic solutions



Developing and commercialising its proprietary platform technology, *3base*™



Implementing its commercial strategy through teams in Australia, Europe and North America

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Scalable business which has achieved strong core revenue growth

Financial information

Enterprise value	A\$242.8m	
Debt (31-Mar-20)		
Cash (31-Mar-20)	A\$39.2m	
Market capitalisation	A\$279.0m	
Shares on issue	142.6m ¹	
Share price (27-May-20)	A\$1.96	



Top shareholders %

Asia Union (Chris Abbott private investment)	26.7%
Karst Peak (HK-based investment manager)	13.3%
Perennial Value Management	9.4%
Fidelity International	7.7%
Directors, management & advisors	3.6%

1: Excludes 3.28m unquoted options (various expiration dates and prices)





Infectious diseases continue to dominate global health threats



Competitive advantage underpinned by novel 3base[™] technology



Trusted and proven technology with third party validation



COVID-19 pandemic creating new opportunities for global expansion



Attractive and scalable revenue model



On track to achieve multiple commercial milestones in 2020



Significant global problem: infectious diseases continue to dominate the list of global health threats



Unprecedented impact of COVID-19 pandemic in 2020



Infectious diseases dominated WHO's List of 2019 Health Threats:

- Global Influenza pandemic
- Antimicrobial resistance
- Ebola and high-threat pathogens
- Dengue
- HIV
- Vaccine hesitancy

Infectious diseases continue to be one of the most serious threats to global health and the recent COVID-19 pandemic has highlighted the importance of diagnostic solutions

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Trusted and proven technology: *EasyScreen[™]* products built on **3base[™]** technology



Our Products

Transforming **molecular diagnostics** via streamlined sample processing methods linked to highly **multiplexed real-time PCR screening assays**.

Our automated **sample preparation** method is suitable for **bacterial**, **protozoan and viral** (DNA & RNA) targets.

The *EasyScreen*[™] Detection assays simultaneously detect a larger number of pathogen targets in a shorter time than conventional methods.









qPCR¹ detection methodology used, the gold standard for infectious disease diagnosis
 Rapid time to results, with results processed from 4 hours, for up to 188 specimens
 Screening for more targets per patient specimen increases accuracy of diagnosis
 Accelerates treatment path and reduces mortality and morbidity



Laboratories

Clear competitive advantage for target customer base of high throughput labs
 Reduces customer costs through accurate detection and minimising hands on time
 Reduced complexity in molecular testing



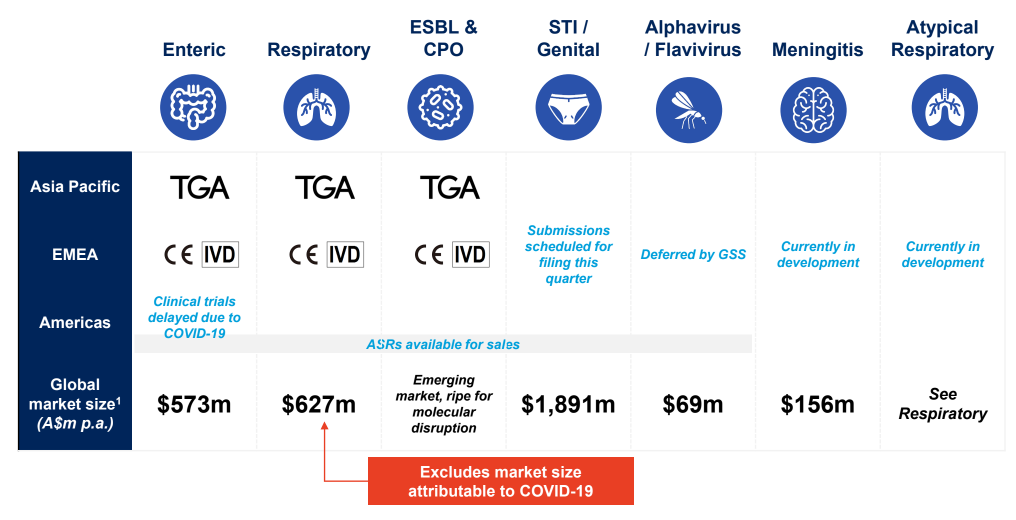
- Reduced hospital stays with broad and accurate detection of infectious disease
- Fast turnaround and accurate detection **reduces the spread of disease**
- Testing for more targets per specimen reduces repeat doctor visits
- Reduces overuse and misuse of antibiotics

1. Real-time polymerase chain reaction (real-time PCR), also known as quantitative polymerase chain reaction (qPCR)

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Large addressable market: new regulatory registrations and product launches imminent in APAC, EMEA and North America





1. World Market for Molecular Diagnostics, 5th. Edition (Infectious Disease, Oncology, Blood Screening, Pre-Natal and Other Areas) Kalorama Information, Published: 1/9/2013 & company estimates

COVID-19 global pandemic: as countries move towards reopening, the burden on testing will significantly increase



FMFA:

- In the UK, 40-50k cumulative deaths are likely to occur by August 2020 with over 100k tests required per day³
- Germany is forecast to see 10k deaths by August and is expecting to consume 184k tests each dav³
- 85% of testing in the UK is managed under NHS
- Germany has many mid-sized pathology labs spread geographically

APAC:

- In Australia, 7k cases have been confirmed, resulting in 102 deaths⁴
- 943k tests have been conducted to date (35% in NSW, 31% in Victoria)⁴
- Australian government has established a dedicated Medicare funded and bulk billed pathology test for COVID-19 worth \$170m⁵
- Market dominated by large private pathology groups and state government labs

The COVID-19 pandemic has created an opportunity for Genetic Signatures to accelerate international expansion and increase customer acquisitions

1.) The National Ensemble Forecast, Centers for Disease Control and Prevention (14 May, 2020) https://www.cdc.gov/coronavirus/2019-ncov/covid-data/forecasting-us.html; 2.) The Mechanics of the COVID-19 Testing Suppl Chain: Version 2.0, Harvard University https://ethics.harvard.edu/covid-supply-chain; 3.) Institution of Health Metrics and Evaluation, University of Washington 12 May, 2020) (http://www.healthdata.org/) 4.) Australian Government Department of Health (24 May, 2020); 5.) '\$2.4 billion plan to fight COVID-19', Prime Minister of Australia media release 11 March, 2020 (https://www.pm.gov.au/media/24-billion-health-plan-fightcovid-19)

North America:

- In the US, cumulative reported deaths are likely to exceed 100k¹ by June 1st and 147k³ by August
- More than 1m tests per week are currently being • delivered and this needs to be increased dramatically to reopen the economy²
- FDA EUA has been submitted for SARS-Cov-2 kit





Introducing the *EasyScreen[™]* SARS-CoV-2 Detection Kit

- In January 2020, GSS announced that the *EasyScreen[™]* Respiratory Pathogen targets included an assay for all known coronaviruses, including the new strain that originated from China (now known as SARS-CoV-2)
- The EasyScreenTM SARS-CoV-2 Detection Kit has been designed to provide rapid and accurate detection of SARS-CoV-2
- SARS-CoV-2 kit may be used alone or in conjunction with the current *EasyScreenTM* Respiratory Pathogen Detection Kit
- The kit is a PCR-based test, which detects the virus's genetic material this is increasingly being recognised as the 'gold standard' for COVID-19 testing¹

Vaccine development is likely to take >12 months creating challenges for labs around the world



Accurate and reliable diagnostic kits are vital for identifying and containing the spread of disease



Labs require tests from multiple suppliers to meet increasing demand and ensure they have access to a diversified supply



Labs are running thousands of tests per day - a high throughput solution is required with minimal hands-on time

1. The Royal College of Pathologists of Australasia (RCPA) supports the use of molecular tests or SARS-COV-2 and advises against the use of serological COVID-19 IgG/IgM rapid tests, such as a pin prick blood test, to detect early COVID disease.



Rapid response to COVID-19: Genetic Signatures has made rapid progress in bringing test kit to international market



Strongly positioned to test for SARS-CoV-2



Genetic Signatures' existing *EasyScreen™* Respiratory Pathogen Detection Kit **already included an assay for pan-coronaviruses**, **highlighting utility of 3base™**



3base[™] technology provides resistance to genetic drift or mutations of pathogens over time allowing preservation of clinical specificity



High throughput allows **testing** of up to 1,500 samples in a 24-hour period in batches of 94 to 188 samples per run

EasyScreen[™] SARS-CoV-2 Detection Kit update



CE-IVD and TGA received – kits can now be sold in Europe and Australia



FDA EUA¹ application - application submitted and clearance expected this quarter



Domestic testing underway – multiple customers are using the kits for routine testing



Site initiations underway for new customers in EMEA – in the final stages of validation



Driving global sales - international sales team and distributors in place in key regions



Expanded sales force - new appointments made to promote kits globally





Attractive revenue model

- High throughput with predictable orders
 - Target customers are **high throughput** pathology groups, hospitals or government run programs
 - Customers secure long-standing contracts with set prices and relatively predictable volumes
 - **Regular orders** (bi-monthly) with **fast payment terms** relatively low working capital needs

Sticky annuity revenue:

- "Printer & cartridge" model tests become embedded in workflow
- Customers may adopt new tests once workflow established

Attractive return on investment:

- Potential to fund new customer installations to **speed up customer** acquisition, particularly offshore
- Consumable revenue model customers pay per test

Contributing to attractive economics

100% customer retention since 2016

47% core revenue CAGR (FY15-19)

gross margins on 65% gross margins diagnostic kits

3-5

year contracts typically secured

Scalability supported by expanding pipeline of new customers / tenders

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Track record of success: year-on-year revenue growth with the potential to achieve record revenue in FY20



Key Highlights



Achieved **47%** core revenue CAGR from FY15-19

International sales

Focus on **international expansion** targeting new US and EU customers in 2H FY20

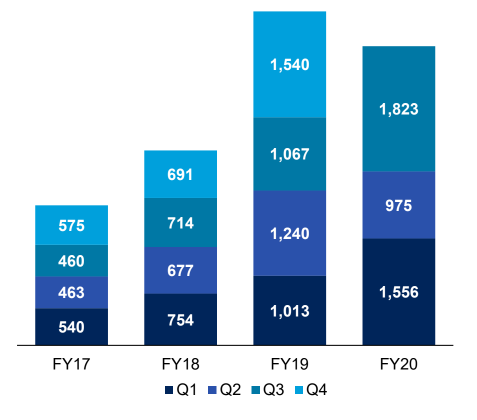
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Continued growth

Focused on growing core revenues where recent coronavirus outbreak has created both obstacles and new opportunities

Revenue (FY17 to Current)







Near term opportunities for SARS-CoV-2 commercialisation	 First US customer contract for SARS-CoV-2 product - targeting mid-2020
Launching <i>EasyScreen™</i> products in new markets	 TGA / CE-IVD submission for the STI / Genital kit scheduled to file this quarter Work set to recommence once COVID-19 restrictions are lifted: FDA submission for the Enteric Protozoan kit TGA / CE-IVD submissions for the Flavivirus / Alphavirus kit
Leverage growing international exposure to drive new contract wing	 Interest in the SARS-CoV-2 products likely to drive interest in Genetic Signatures' broader range of <i>EasyScreenTM</i> multiplex kits and facilitate <i>new contracts in US and Europe</i>

Active M&A space: a history of consolidation in the molecular diagnostics space with M&A activity expected to continue



Date	2020	2018	2018	2018	2017	2016
Target Company	QIAGEN (NYSE:QGEN)	STATdx* Private	(NASDAQ:CPHD)	Fast Track DIAGNOSTICS A Siemens Healthineers Company (Private)	EUROIMMUN a Perioditine company (Private)	(NYSE:DGX)
Acquired by	ThermoFisher SCIENTIFIC (NYSE:TMO)	QIAGEN (NYSE:QGEN)	(NYSE:DHR)	SIEMENS (ETR:SIE)	PerkinElmer For the Batter (NYSE: PKI)	DiaSorin (BIT:DIA)
Transaction	Takeover	Takeover	Takeover	Takeover	Takeover	Acquired molecular and immunoassay business
Size	US\$11.5bn	US\$147m upfront US\$44m milestone	US\$4bn	Not disclosed	US\$1.3bn	US\$300m

Strong strategic interest in multiplex panels such as 3base[™] technology





Infectious diseases continue to dominate global health threats: major contributor to global mortality and morbidity dominating WHO's¹ 2019 list of health threats and amplified by recent coronavirus outbreak



Competitive advantage underpinned by novel 3base™ technology: resistant to genetic mutations in the micro-organisms, and implemented in a simplified workflow processes, increasing throughput capacity, reducing time to results and creating cost saving benefits



Trusted and proven technology with third party validation: Genetic Signatures have achieved 100% customer retention since 2016 and the accuracy of the technology has been clinically validated



COVID-19 pandemic creating new opportunities for global expansion: increasing international recognition through the *EasyScreenTM* SARS-CoV-2 launch creates new avenues to expand customer base



Attractive and scalable revenue model: highly scalable business model with favourable unit economics expected to underpin growth through FY20 and beyond



On track to achieve multiple commercial milestones in 2020: international interest bolstered by the SARS-CoV-2 product release creates a platform for new contract wins for broader $EasyScreen^{TM}$ products

Genetic Signatures

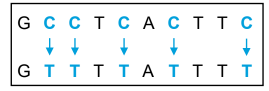
Transforming Molecular Diagnostics

Appendices

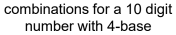
Novel proprietary technology: proprietary 3baseTM platform technology underpins the *EasyScreenTM* product range

Our proprietary 3base[™] solution...

- 3base[™] platform technology converts original 4-base microbial genome to 3-base
- 2 Conversion occurs during standard procedures with no additional steps for the technician
- 3 **3base™** MDx can identify a wider array of pathogens and provide greater testing accuracy by reducing complexity in a multiplex environment



1,048,576





combinations for a 10 digit number with 3-base

...improves workflow and increases throughput



EasyScreen[™] kits are compatible with most existing automated nucleic acid extraction and real-time PCR instruments and streamline the preparation process



High throughput labs can achieve **further workflow automation** with compatible hardware. Creates **workflow efficiencies and reduce costs**



Currently offering over 100 pathogen targets across enteric, respiratory, antimicrobial resistance, sexual health and tropical diseases

Genetic Signatures **Board of Directors:** proven track records of commercialisation success across key geographic regions



	Nick Samaras Non-Executive Chairman	 Significant experience in leading international sales expansions of biotech companies Former Managing Director of Applied Biosystems (acquired by ThermoFisher, US\$76.8bn market cap) Held senior roles with Perkin Elmer and AMRAD Corporation (now part of CSL)
	John Melki Managing Director & Chief Executive Office	 Led global commercialisation efforts of GSS since 2011 and the product development team since 2003 Successfully commercialised seven products globally Authored 20 peer-reviewed articles and listed as an inventor on eight patent applications
	Michael Aicher Executive Director	 Founder and former CEO of National Genetics Institute (subsidiary of LabCorp, US\$15.3bn market cap) Led Lab-Corp's Esoteric Business Units which generated over US\$1b revenue p.a. Former executive roles at Central Diagnostics Laboratory Recipient of Ernst & Young "Entrepreneur of the Year" award for emerging technologies
3	Tony Radford AO Non-Executive Director	 Former Co-Founder and CEO of Cellestis (ASX:CST, acquired by QIAGEN for c.A\$350m in 2011) Former member of CSIRO team that invented QuantiFERON Former Head of Development at AMRAD (later acquired by CSL)



Dr. Doug Millar Chief Scientific Officer	 One of the pioneers of the bisulphite genomic sequencing protocol with a PhD in Molecular Genetics Key inventor on over 30 patents or pending patent applications held by the company Authored 23 peer reviewed scientific papers and presented at 20+ international conferences
Peter Manley Chief Financial Officer & Company Secretary	 Led the recent Genetic Signatures capital raise, successfully securing \$37.5m Served as CFO and Company Secretary for AtCor Medical (now Cardiex) and Sirtex Medical Senior financial positions including 8 years with Dow Chemical and 4 years at Goodman Fielder
Jackson Jones Director of Global Sales & Marketing	 20+ years experience in clinical diagnostics, blood banking, and life sciences sector Joined Genetic Signatures in 2017 and brings significant commercial experience from working with several large US multinationals and roles across Australasia, Europe, and North America
Derek Joesting Director of Sales - North America	 20+ years of medical sales experience with broad sector experience Previously held leadership roles in molecular diagnostics and pathology sales in North America Holds a Bachelor of Science degree in Biology from Syracuse University
John Buckels Director of Sales & Support - Europe	 20+ years' experience in molecular biology and sales across the EMEA Former Senior Director and Head of Infectious Diseases sales at QIAGEN and 13 years experience in sales and marketing
Neralie Coulston Regulatory Affairs Manager	 Supported Genetic Signatures since 2002 and brings significant experiences in Quality System and Regulatory Affairs Former roles at the CSIRO and UNSW on both therapeutic development and research programs

Contact us

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