

Investor Presentation

October 2022

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Investment highlights

- **Proprietary 3base[®] technology platform** that provides a revolutionary approach for molecular diagnostics
- Dramatically simplifies multiple pathogen testing from a single sample (multiplexing)
 - More informative detect related pathogens/genes using fewer tests;
 - Simpler fewer reagents with better matched, ideal reaction conditions.
- Strong commercial adoption in Australian market expanding into European and US markets
 - 4 Diagnostic Test Kits cleared in one or more markets 5 new kits completing development;
 - Strong continued revenue growth FY22 revenue A\$35.4 million (+25% yoy), cash flow positive (\$6.7M) and profitable (\$3.1M).
 - Multiple drivers for growth funded from anticipated future cash flow and existing balance sheet
 - Commercial expansion into large international markets (Europe and US);
 - Product expansion multiple new products completing development or registration;
 - Instrument expansion embed 3base[®] technology in high-volume customers sites.



Recent Events

• US FDA clinical trial recruitment completed

- Targeting 4Q CY2022 application for clearance
- 3base[®] kit for antimicrobial resistance shows high detection rate
 - Independent study¹ showed excellent biological performance for the 5 most common carbapenemases
 - WHO has declared AMR as one of the top 10 global public health threats facing humanity > 5m deaths pa²

GSS commences commercial sales in Western Australia

 Two new sites trialling respiratory & gastrointestinal targets; first sales into WA

EasyScreen[™] Enteric Protozoan Detection Kit - application lodged with Health Canada

- Canadian market ~2.5% of world IVD market
- Will be 3rd *EasyScreen*[™] Detection Kit registered

¹ Gonzales, C et al, (2022), *Diagnostics* 2022, 12(9), 2223; <u>https://doi.org/10.3390/diagnostics12092223</u>
² Antimicrobial Resistance Collaborators (2022), The Lancet: https://doi.org/10.1016/ S0140-6736(21)02724-0



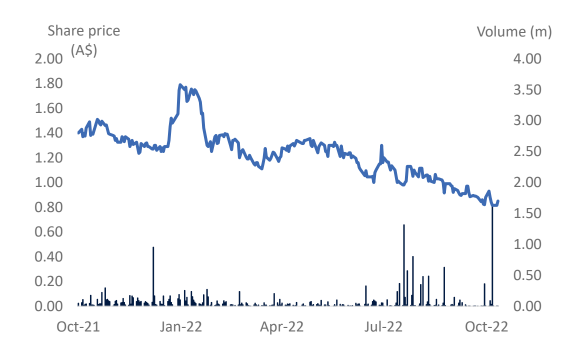


Financial information

Enterprise value	A\$85.0m
Debt (30-Jun-22)	Nil
Cash (30-Jun-22)	A\$36.9m
Market capitalisation	A\$121.9m
Shares on issue	143.4m
Share price (11-Oct-22)	A\$0.85

Top shareholders %

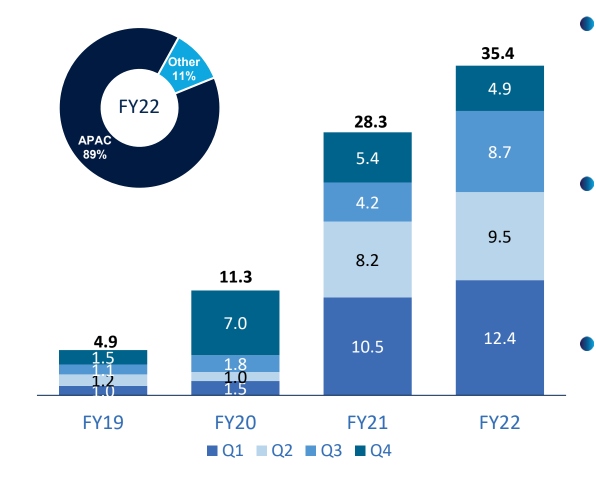
Asia Union (Chris Abbott private investment)	26.2%
Perennial Value Management	15.0%
Fidelity International	6.9%
Directors & management	3.0%



Track record of continued, strong revenue growth

Sales Revenue (A\$m)

FY22 sales revenue of \$35.4 million (+25% yoy, 89% 4yr CAGR)



Growing contribution from international sales

- Leveraging experience in Australian market;
- European orders for non-Covid Syndromic Kits;
- Significant US contributions to come once FDA clearance secured.

Strong demand for SARS-CoV-2 tests during FY21 & FY22

- Scale-back of molecular testing programs;
- Growing contribution from other *EasyScreen*[™] Kits;
- Shifting from COVID to Syndromic Respiratory.

Successful strategy of targeting high-volume customer groups

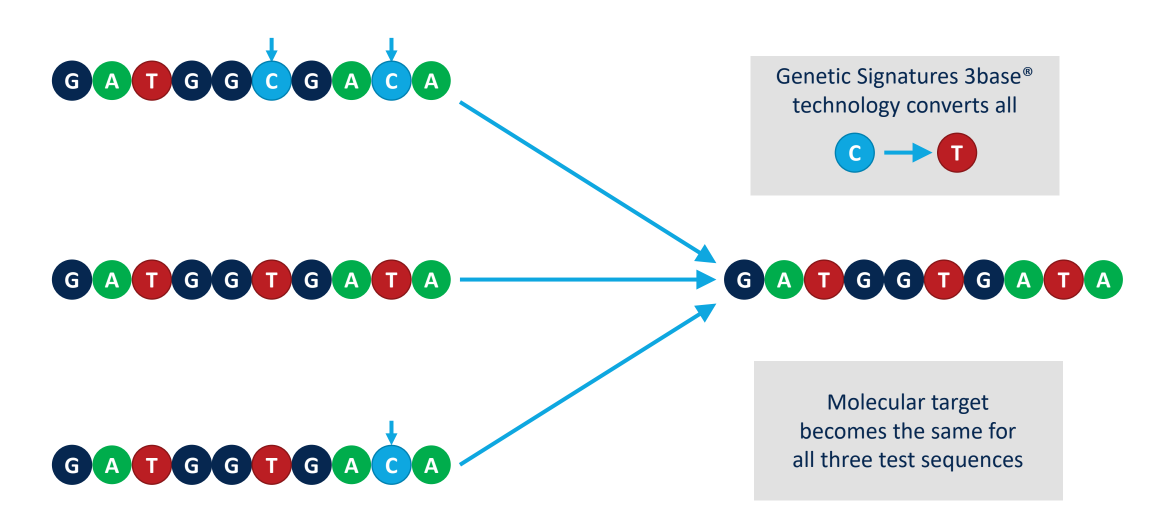
- High-throughput labs
- Multi-hospital groups
- Private pathology chains
- Government-led programs

3base[®] - a revolution in molecular diagnostics

- Molecular diagnostic tests are based on DNA/RNA sequences
 - DNA/RNA is unique to each organism.
- Molecular diagnostic tests are recognised as the 'gold standard'
 - Precisely targeted and highly specific PCR tests;
 - Can be less effective when:
 - Need to detect multiple pathogens or genes;
 - New strains or subtypes of pathogens emerge.
- Molecular diagnostic tests are often multiplexed
 - Multiplexing refers to conducting multiple tests simultaneously
- **Genetic Signatures 3base**[®] makes multiplexing easier:
 - More informative detect related pathogens/genes using fewer tests;
 - **Simpler** fewer reagents with better matched, reaction conditions.







* Human Papilloma virus sequences

How 3base[®] tests are performed

Proprietary method - patented until 2031+

1. Extraction and Conversion

• natural 4 bases to 3base®





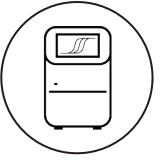
2. DNA Amplification (PCR)

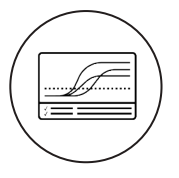
• uses 3base® DNA



3. Detection (primers & probes)

• uses 3base[®] DNA





Benefits

- 🔮 🛛 Rapid
- High throughput
- Informative
- Sensitive
- Specific
- Low manual involvement
- Reduced contamination risk

Equipment

- Run on standard equipment.
- Genetic Signatures' instruments further automate the process;
 - increase throughput
 - reduce labour.

3base[®] simplifies Syndromic Testing – *EasyScreen*[™] Kits



- Syndromic testing: simultaneously test for multiple pathogens that all can cause the same signs and symptoms
 - <u>Respiratory infections</u>: cough, runny nose, sore throat, headache, breathlessness;
 - <u>Gastrointestinal infections</u>: nausea, diarrhea, vomiting, abdominal cramps, fever.

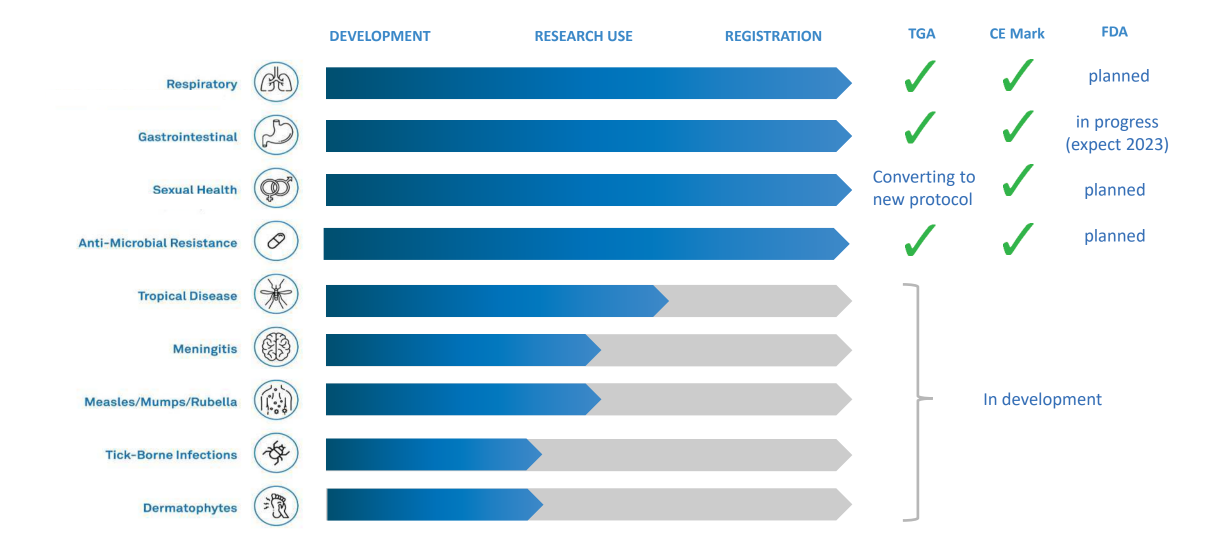
Syndromic testing

- allows single test to determine the potential cause of a disorder;
- avoids having to order separate tests for each possible pathogen.
- Genetic Signatures' *EasyScreen*™ is ideal for Syndromic Testing
 - Tests for over 100 different types of pathogens;
 - Able to detects variants (i.e. different strains or subtypes);
 - Combine tests to create *EasyScreen*[™] Syndromic Detection Test Kits;
 - Detect >20 different pathogens from a single sample.



Robust pipeline with multiple products cleared for sale





Growth initiatives

Leverage experience in Australian market to grow international sales

- Europe drive adoption of other 3base[®] products;
- US build 3base[®] franchise once Protozoan Detection Kit is cleared.
- Build and expand portfolio of commercially-available *EasyScreen*™ products
 - Expand menu of 3base[®] tests;
 - Develop new *EasyScreen*[™] Syndromic Test Kits;
 - Secure registration for new *EasyScreen*[™] products.

Embed 3base[®] technology in high-value customer's workflow

- Increase adoption of *EasyScreen*[™] kits for more applications;
- Broader range of commercial arrangements with customers.



Enteric Protozoan kit will provide entry to North America





North America accounts for 40% of the global molecular diagnostics market

Enteric Protozoan Screening Kit

- Completed recruitment for 1,500 subject clinical trial;
- Targeting 510(k) submission in Q4 CY2022;
- First *EasyScreen*[™] product for US

High need for Enteric Protozoan Kit

- 5.5 million tests conducted in the US pa;
- Primarily culture/microscopy: slow, labour intensive, unreliable;
- Detects leading protozoan infections.

US Market preparation activities underway

- KOL webinars;
- Sales & marketing presence in US;
- Warehousing facility in Los Angeles;
- Initial focus on 30 high-throughput, centralised labs;

First 3base[®] product for the US

• Regulatory dossier relevant for other *EasyScreen*[™] products.

New products and instruments





Expand available *EasyScreen*[™] Syndromic Kits

- 3 kits research use only (RUO) tropical diseases, MMR & meningitis;
- Other kits in development (tick-borne, skin infections, etc.);
- Advance additional 3 products through the FDA process

Improve and enhance 3base[®] technology platform

- Saliva-based protocol for SARS-CoV-2 cleared by TGA;
- Process improvements for amplification and time-to-result

Next-generation, "sample-to-result" instrument

- Highly automated, high-throughput;
- Ideally suited for high-volume commercial users;
- Embed use of 3base[®] with customers;
- Facilitates different commercial models;



Image is concept only

Upcoming milestones – 12 months

US Enteric Protozoan Kit

- File 510(k) application by end of CY2022;
- Launch product once clearance is granted.
- Increase sales and presence in UK and European markets
 - Contracts with new customers;
 - Direct sales force and distributor appointments.
- Initiation of US clinical trial for next *EasyScreen*[™] product

• R&D initiatives for new products

- New tests and *EasyScreen*[™] kits;
- Technology improvements;
- Development of Next Generation instrument prototype.
- Quarterly sales updates and progress reports



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