



11 June 2026

ASX Announcement

Investor presentation

MELBOURNE Australia, 11 June 2026: Australian antiviral drug development company, Island Pharmaceuticals Ltd (**ASX: ILA; Island or the Company**) is pleased to provide the following presentation which will be used at the Gold Coast Investment Showcase from 11 June to 12 June 2026, and during the Company's investor webinar, scheduled for 11:00am AEST (9:00am AWST) on Friday, 12 June.

The presentation highlights the Company's growing momentum across regulatory, clinical and manufacturing initiatives, including progression under the US FDA Animal Rule pathway, execution of an expanded CRADA with USAMRIID, commencement of GMP manufacturing and the upcoming dose optimisation study.

Together, these initiatives are expected to further de-risk the Galidesivir program while supporting multiple potential value drivers, including a Tropical Disease Priority Review Voucher and future government procurement opportunities.

The presentation also discusses the increasing global focus on biodefence preparedness, the limitations of existing filovirus countermeasures and Galidesivir's potential role as a broad-spectrum antiviral capable of addressing multiple high-consequence viral threats.

Webinar details:

Island will host an investor webinar at 11:00am AEST (9:00am AWST) on Friday, 12 June 2026. During the webinar, CEO and Managing Director, Dr David Foster and Non-Executive Chairman, Mr Jason Carroll will provide a broader insight into the expanded CRADA with USAMRIID and provide an update on its broader biodefence engagement and other near-term opportunities associated with Ebola and Sudan virus.

Link: https://us02web.zoom.us/webinar/register/WN_rcig93IHSUCU2qyG507lww

Date and time: 11:00am AEST (9:00am AWST) on Friday, 12 June CY26

A recording of the webinar will be made available following the broadcast.

- Ends -

Approved for release to the ASX by:

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About Island Pharmaceuticals



Island (ASX: ILA) is focused on areas of unmet need for drugs that can address urgent viral diseases, public health or biosecurity threats. The Company is executing a dual development strategy for its assets, ISLA-101 and Galidesivir.

ISLA-101 has a well-established safety profile, being repurposed for the prevention and treatment of dengue fever and other mosquito (or vector) borne diseases. Galidesivir is a clinical-stage antiviral molecule with a broad spectrum of activity in over 20 RNA viruses, including high-priority threats such as Ebola, Marburg, MERS, Zika and Yellow fever – viruses with significant unmet medical needs and that may contribute to national security threats.

Island encourages all current investors to go paperless by registering their details with the Company's share registry, Automatic Registry Services, whose contact info is housed on the Shareholder Services page of the Company's website.

Visit www.islandpharmaceuticals.com for more on Island.

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ISLAND
PHARMACEUTICALS

COMBATTING URGENT VIRAL DISEASE THREATS

Advancing Galidesivir as a muflovirus countermeasure under the FDA
Animal Rule

Dr David Foster, CEO & Managing Director | Mr Jason Carroll, Non-Executive Chairman

June 2026

ASX: ILA

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ISLAND PHARMACEUTICALS (ASX: ILA)

TWO PROGRAMS TARGETING INFECTIOUS DISEASES



Two, well advanced clinical stage programs



Major market potential via both programs



Both assets have Priority Review Voucher potential



Phase 2a/b PROTECT clinical trial in dengue complete

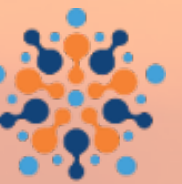


FDA Animal Rule pathway actively advancing through new USAMRIID CRADA



Multiple near term clinical trial, operational and regulatory catalysts

Company Overview



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Shares on issue ¹ :	295,916,682
Price per share ¹ :	\$0.365
Market capitalisation ¹ :	\$108.1m
Cash at bank (31 March 2026) ² :	\$14.2m
Potential additional capital from vested options where current share price exceeds exercise price:	~\$1.6m
Debt:	Nil

Substantial shareholders

Dr William James Garner ³	15.50%
Jason Alan Carroll ³	11.92%
MWP Partners Limited ⁴	8.25%
Dr Daniel Tillett ³	7.80%

Board of Directors

Jason Carroll, Non-Executive Chairman

Dr David Foster, CEO & Managing Director

Chris Ntoumenopoulos, Non-Executive Director

1. As at 10 June 2026
2. Does not take into consideration cash movement since reporting date
3. Per holding per Substantial interest notice lodged with ASX on 9 December 2025
4. Per holding per Substantial interest notice lodged with ASX on 3 June 2025

Price & volume (12 months)



Company Overview



- ✿ Two clinical stage assets Galidesivir and ISLA101 - both with Priority Review Voucher potential based on approval
- ✿ Galidesivir
 - Small molecule with broad antiviral activity against numerous high-priority threats
 - Robust development history with over US\$70m in funding to-date from US government
 - Opportunity to leverage FDA's Animal Rule to fast-track approval in Marburg
- ✿ ISLA101:
 - Pre-clinical work at Monash University highlighted antiviral promise
 - 40+ Phase I, II and III human trials in cancer and respiratory diseases, and deemed safe by regulators
 - Small molecule with activity against all 4 dengue serotypes and other mosquito borne viruses
 - Successfully completed Phase 2a/b clinical trial in dengue infected subjects
- ✿ Robust balance sheet allows for execution of program development

The Filovirus Threat Landscape



AN URGENT AND GROWING GLOBAL SECURITY CHALLENGE

Extremely high fatality rates: Marburg up to 88%, Ebola up to 90%, Sudan up to 47%, Bundibugyo ~~300%~~

Limited countermeasures
No approved therapeutics or vaccines for Marburg or Bundibugyo

BSL4 pathogens:
Require maximum containment; limited global capacity

Bioterror relevance:
Historical weaponisation research; persistent intelligence concern

Current situation:
Ongoing Bundibugyo outbreak with no available medical countermeasures

The US lacks a broad acting antiviral capable of addressing multiple filovirus threats

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The Preparedness Gap



THERE ARE NO FILOVIRUS THERAPEUTICS CAPABLE OF CROSS PROTECTION



Existing Ebola countermeasures are strain-specific (Zaire only)



No approved therapeutics for Marburg, Bundibugyo or Sudan virus



Outbreaks increasingly involve rare or divergent strains



Stockpile lacks a broad-acting antiviral with Animal Rule feasibility

GALIDESIVIR DIRECTLY ADDRESSES THIS GAP


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Why Galidesivir?



A BROADCASTING ANTIVIRAL WITH STRONG FILOVIRUS EFFICACY AND A BARRER PATHWAY

-  Demonstrated multi-filovirus activity (Marburg, Ebola, Sudan)
-  Strong in vivo efficacy with delayed dosing
-  FDA confirmed Animal Rule pathway is appropriate

-  IV and IM formulations with favorable safety profile
-  Manufacturing route improved with 2–3X yield increase
-  Developed with NIAID and BARDA support (>US\$70M historically)

GALIDESIVIR IS ONE OF THE FEW ANTIVIRAL CANDIDATES WITH CREDIBLE PROSPECTS

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Galidesivir Competitive Advantage



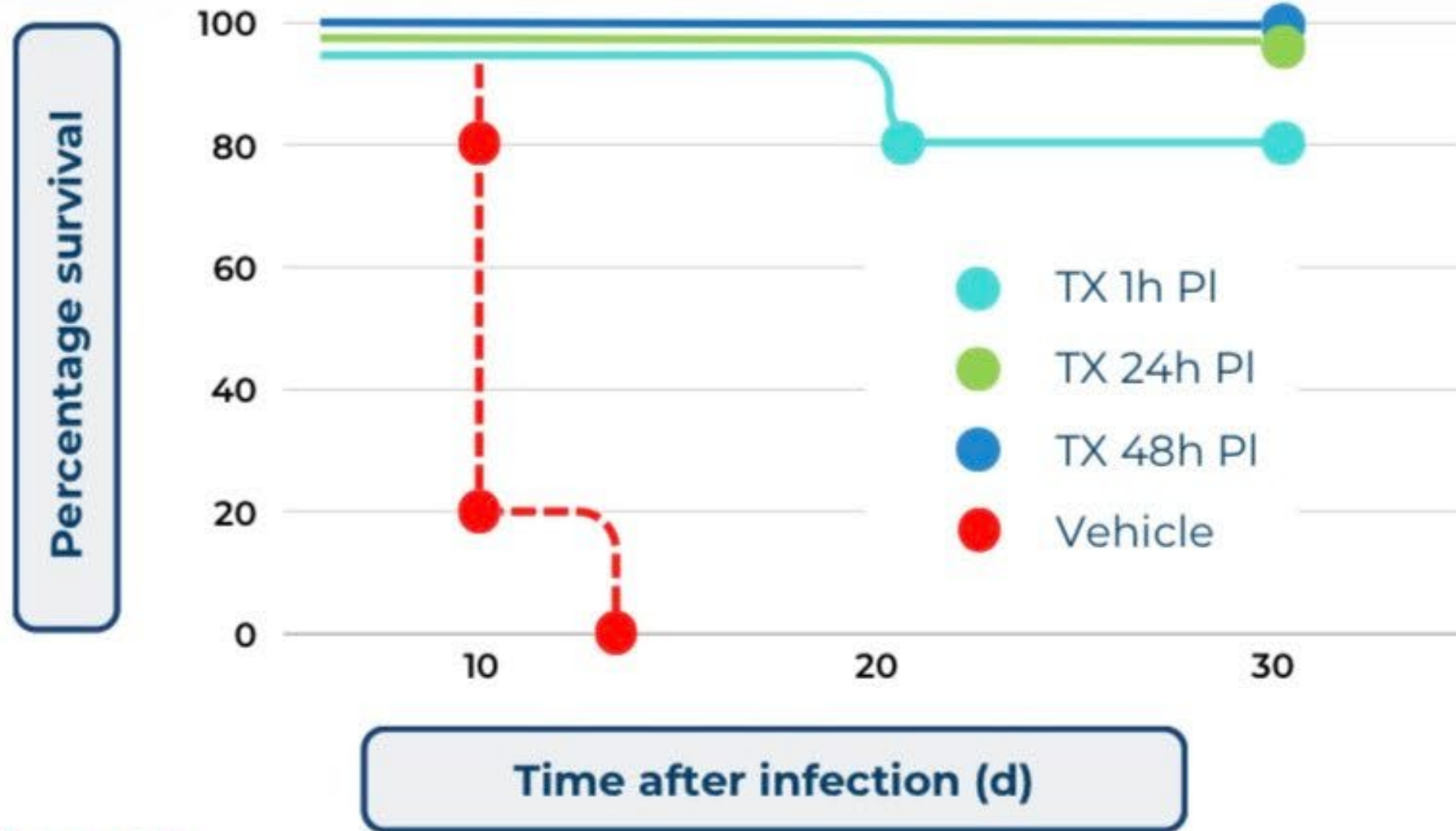
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FEATURE	GALIDESIVIR	OBELDESIVIR	REMDESIVIR	MAFTIVIMAB	MBP134	CHADOX-BDBV	RVS-BDBVGP	MRNALNP BDBV
BROADEST SPECTRUM ANTIVIRAL ACTIVITY (FILOVIRUSES AND OTHER FAMILIES)	✓	✗	✗	✗	✗	✗	✗	✗
PEP AND TX FLEXIBILITY (IM/IV)	✓	✗	✗	✗	✗	✗	✗	✗
ROOM TEMP STABLE	✓	✓	✗	✗	✗	✗	✗	✗
SMALL MOLECULE	✓	✓	✓	✗	✗	✗	✗	✗
PHASE 1 HUMAN SAFETY DATA	✓	✓	✓	✓	✓	✓	✓	✓

MARBURG NHP SURVIVAL



94% survival in Marburg NHP Model with treatment initiated up to 48 hours post infection



6/6 animals survived when dosed 48 hours post infection
6/6 animals survived when dosed 24 hours post infection
5/6 animals survived when dosed 1 hour post infection
0/6 untreated animals survived as part of the control group

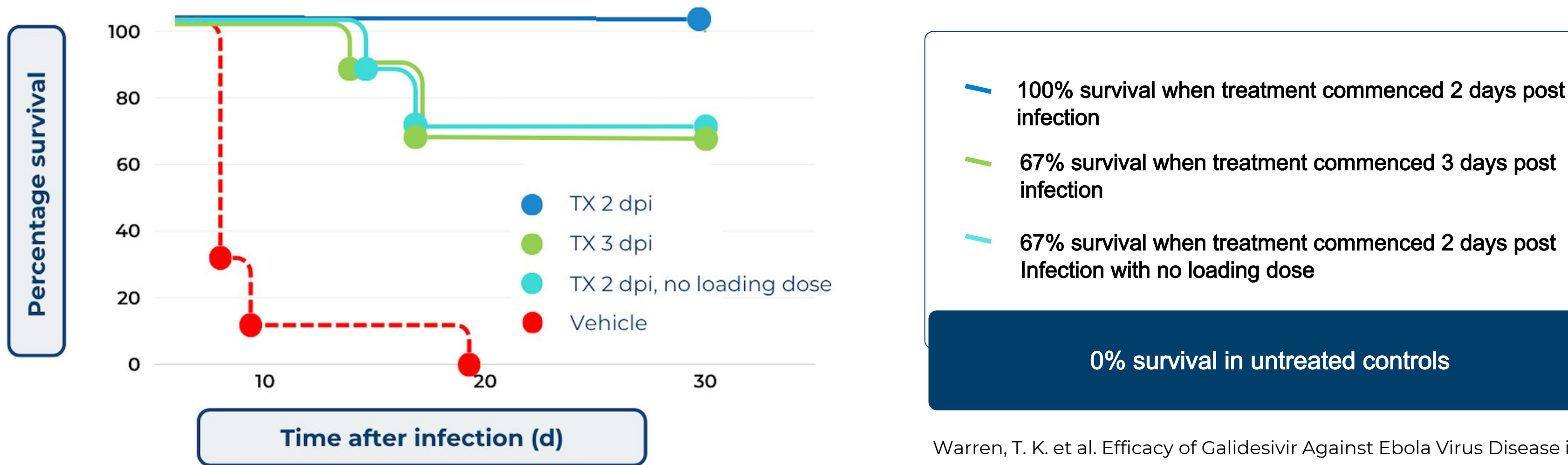
This level of efficacy, with a clinically meaningful therapeutic window, is rare in filovirus models and directly supports **IND Rule advancement**

EBOLA NHP SURVIVAL



Robust efficacy in Ebola NHP model with delayed dosing

Survival of rhesus non-human primates challenged with Ebola virus following intramuscular administration of 100 mg/kg BID loading dose followed by 25 mg/kg BID for 10 days.



Warren, T. K. et al. Efficacy of Galidesivir Against Ebola Virus Disease in Rhesus Monkeys. Poster Presentation ID Week 2017

Together with Marburg data, this positions Galidesivir as a ~~m~~filovirus antiviral candidate

FDA alignment significantly de-risks regulatory pathway



Island has the potential to become the first Australian company to gain drug approval via the FDA's Animal Rule



FDA confirmed the Animal Rule pathway is appropriate for developing countermeasures against Marburg virus



Clear guidance provided on clinical program design – enables Island to continue to engage with the FDA and finalise plans ahead of trial commencement



FDA advised that Galidesivir would qualify for a Tropical Disease Priority Review Voucher (PRV) on approval – Most recent PRV sold for US\$200m

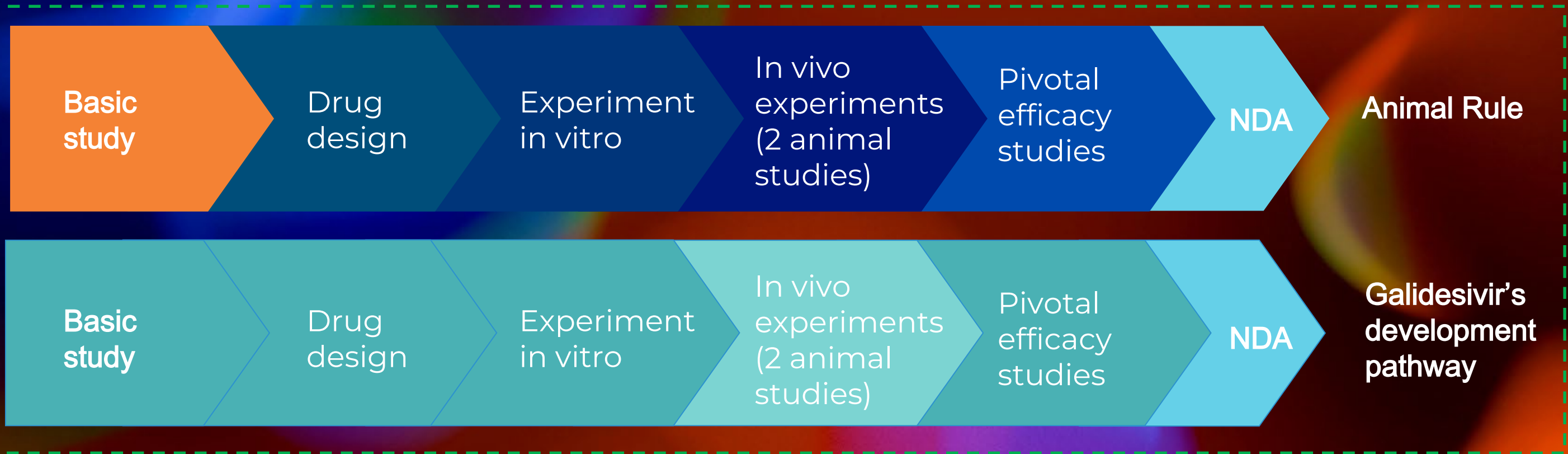
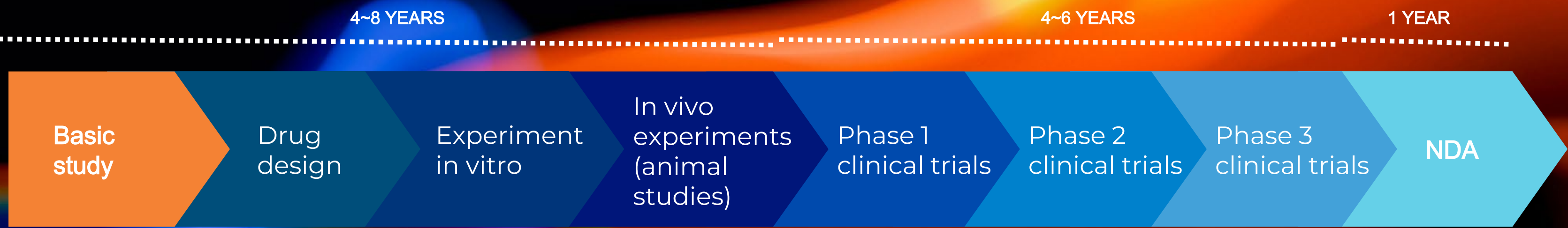


CRADA secured with USAMRIID to commence the next Galidesivir animal study in Marburg to advance clinical development pathway and potential approval

An Established Regulatory Pathway



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Confirmed PRV opportunity with a potential value of ~US\$200m

Island is now focused on advancing Galidesivir's clinical development pathway, which includes a dose optimisation study to commence shortly with USAMRIID

Expanded CRADA with USAMRIID



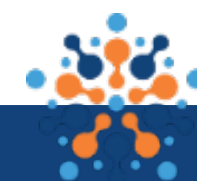
Agreement with US Army Medical Research Institute of Infectious Diseases and the Geneva Foundation to secured 23 non human primates and advance dose optimisation study



USAMRIID is the US Army's premier research institute and Biosafety Level 4 (BSL-4) facility for the study of infectious diseases and medical countermeasures against biowarfare threat agents



Amended CRADA confirms supply of 23 non-human primates and timeslot for dose optimisation study – Study to commence next quarter



Dose optimisation study designed to identify minimally effective dose required prior to pivotal confirmatory trial for potential FDA approval



Study to evaluate high loading dose on day one followed by various dose regimens and treatment initiations over five NHP cohorts

Expanded CRADA materially de-risks the Galidesivir program by securing non-human primate supply and working with one of the world's leading biodefence and high-containment infectious disease research organisations

NHP dose optimisation protocol



STUDY GROUP	# ANIMALS	GALDESIVIR DOSE	GALDESIVIR ADMINISTRATION (HR POST VIRUS EXPOSURE)	DOSING FREQUENCY	DISEASE BIOMARKER SPECIMEN COLLECTION	PK SPECIMEN COLLECTION	ENDPOINTS
1	4	0 MG/KG (VEHICLE CONTROL - 30MIN INFUSION)	24 HOURS	BID	DAY -14, 0 (PRE-CHALLENGE), 1, 2, 3, 4, 5, 7, 10, 14, 28	DAY -14, 0 (PRE-CHALLENGE); DAY OF FIRST TX (PRIOR TO TX; SIX INTERVALS POST TREATMENT TO MATCH PK CURVE); LAST DAY OF DOSING (PRIOR TO TX; SIX INTERVALS POST TREATMENT TO MATCH PK CURVE)	CLINICAL OBSERVATIONS, BODY WEIGHTS, FOOD CONSUMPTION, CLINICAL PATHOLOGY, VIREMIA (PCR AND PLAQUE ASSAY), BODY TEMPERATURE AND ACTIVITY, ANATOMIC PATHOLGY (GROSS NECROPSY AND HISTOPATHOLOGY)
2	4	15 MG/KG (30MIN INFUSION)	24 HOURS	BID			
3	4	10 MG/KG (30MIN INFUSION)	24 HOURS	BID			
4	4	3 MG/KG (30MIN INFUSION)	24 HOURS	BID			
5	4	15 MG/KG (30MIN INFUSION)	48 HOURS	BID			

NOTE: Each active treatment group will be administered two loading doses of 100mg/kg BID, 30min infusion, on treatment day 1. Following the BID loading doses, each cohort will be administered the defined maintenance doses listed in Table 1.

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Recent appointments strengthen clinical development



Combining deep Galidesivir expertise with extensive US biodefence and procurement experience to accelerate development, funding and commercialisation opportunities



Mark Herzog – Senior Global Health Security Advisor

- 25 years'+ experience in biodefence, biopharmaceuticals, government contracting and global health security initiatives
- Proven track record securing and managing US\$100m+ government contracts across biodefence and medical countermeasure programs
- Extensive experience engaging with key US agencies including the DoD, HHS, BARDA and allied procurement organisations
- Former US pharmaceutical executive supporting multiple US biodefence and medical countermeasure initiatives
- Executive Committee member of the Medical CBRN Defense Consortium (MCDC), a leading US biodefence organisation



Raymond Taylor – Senior Scientific Fellow

- Over 40 years' experience in drug development, antiviral therapeutics, biodefence and regulatory strategy
- Spent 19 years in senior leadership roles at BioCryst Pharmaceuticals, where Galidesivir was originally developed
- Direct historical experience with Galidesivir, including antiviral development, program execution and regulatory initiatives
- Secured and managed over US\$490m in US Government funding, including US\$125m+ in Strategic National Stockpile procurement contracts
- Extensive expertise working with BARDA, NIAID, CDC and other US Government agencies in countermeasure development

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Current Ebola outbreak highlights the opportunity



Emerging filovirus outbreaks continue to reinforce the importance of preparedness, rapid response capability and effective medical countermeasures

Active Ebola outbreak in East Africa:

- Budibugyo Ebola virus responsible for recent outbreak activity
- Highlights continued emergence of high-consequence threats
- Currently 101 dead and 550 confirmed cases

Governments continue to prioritise preparedness:

- Emerging outbreaks reinforce investment in biodefence
- There is an increasing focus on rapid-response countermeasures
- Island is actively engaging with industry participants to assist

Limited treatment options available:

- Existing countermeasure capabilities remain limited in Ebola
- Significant unmet need remains for broad-spectrum antivirals
- Global health agencies and governments are pursuing options

WHY GALIDESIVIR CAN BE THE DIFFERENCE:

- Designed to address high-consequence viral threats where treatment options remain limited
- Broad antiviral profile provides potential utility across multiple outbreak and biodefence scenarios
- Marburg development program provides the lead regulatory pathway, while creating optionality across other viral indications
- Increasing global focus on preparedness continues to support demand for effective medical countermeasures

Island is actively engaged in discussions with third parties to accelerate Ebola development and advancing engagement with key industry participants to expedite opportunities

Supply chain optimised to meet pending demand



Critical manufacturing and quality infrastructure being established to support FDA submission and future procurement opportunities

Manufacturing campaign commenced:

- Agreement with PI Health Sciences for a 5kg GMP manufacturing campaign
- Includes analytical method validation, reference standard preparation and stability studies
- GMP-grade Galidesivir to be delivered in the coming months
- Existing Galidesivir inventory available for planned dose optimisation studies

A transition to registration and SNS procurement:

- Establishing validated manufacturing, analytical and quality systems for late-stage development
- GMP product intended for planned pivotal initiatives under the FDA Animal Rule pathway
- Supports future regulatory submissions and government procurement initiatives
- Represents a key transition from development-stage asset to potential biodefence product

Supply for opportunities beyond Marburg:

- Maintains access to GMP-grade product for biodefence and outbreak response opportunities
- Supports engagement with government agencies and procurement stakeholders
- Broad-spectrum antiviral activity demonstrated across multiple high-consequence viral threats
- Enhances readiness to pursue opportunities with current and future outbreaks, including Ebola

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Multiple commercialisation opportunities



Galidesivir has the potential to fill multiple gaps in the US government's biodefence stockpile

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Virus	Cell culture data	Animal data	Non-human primate efficacy	PRV Eligible for first indication	Animal Rule Potential	Strategic National Stockpile Potential
Marburg	✓	✓	✓	✓	✓	✓
Ebola	✓	✓	✓	✓	✓	✓
Sudan	✓			✓	✓	✓
Zika	✓	✓	✓	✓		
Chikungunya	✓			✓		

Animal Rule is a proven path for SNS inclusion



Company	Product	Year Approved	Disease Treated	SNS Sales (AUD)	Under SNS Contract
Emergent BioSolutions	raxibacumab	2012	Inhalational Anthrax	~\$450M	Yes
Kaléo	AUVI-Q	2012	Anaphylaxis (emergency countermeasure)	~\$100M+	No (contract expired)
Emergent BioSolutions	BioThrax	2015	Anthrax (prophylactic vaccine)	~\$1.2B+ (multi-year)	Yes
Elusys Therapeutics	Anthim	2016	Inhalational Anthrax	~\$320M	Yes
SIGA Technologies	TPOXX	2018	Smallpox	~\$850M+ (ongoing)	Yes
Paratek Pharmaceuticals	Nuzyra	2018	Anthrax (post-exposure prophylaxis)	~\$120M (partial uptake)	Yes (limited scope)
varian Nordic	Jynneos	2019	Smallpox / Monkeypox	~\$300M+	Yes
imerix	Tembexa	2021	Smallpox	~\$400M	Yes

Since 2012, the FDA's Animal Rule approval has led to 8 bioterror countermeasures joining the US Strategic National Stockpile (SNS)

In 7 out of 8 cases, these medical countermeasures continue to remain under SNS contract and have generated 'lifetime sales' of between US\$100m - US\$1.2Bn at an average of US\$467m

~US\$600m has been provided through grants to develop a Marburg countermeasure with no tangible results

Marburg is the only Category A biothreat that has no treatment presently available in the Strategic National Stockpile

FDA approval of Galidesivir in Marburg provides a significant opportunity for a Priority Review Voucher as well as a multi-year SNS contract

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