# Paltamira therapeutics

DELIVERING RNA - BEYOND THE LIVER

Investor Presentation Spring 2024

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# **Company Overview**



# Disruptive, Proprietary RNA Delivery Technology Platform



OligoPhore™ (siRNA) SemaPhore™ (mRNA)	<ul><li>Proprietary 21 amino acid peptide (nanoparticles)</li><li>Efficient delivery of RNA into extrahepatic target cells</li></ul>
RNA Market Taking Off	<ul><li>Rapidly growing number of RNA therapeutics</li><li>Active M&amp;A, licensing environment</li></ul>
'Picks and Shovels' Platform Strategy	<ul><li>Partner delivery platforms with pharma &amp; biotech</li><li>Initiated first collaborations</li></ul>
Two Flagship Programs for Demonstration	<ul> <li>KRAS-driven cancers (AM-401) - IND expected in 2025</li> <li>Rheumatoid arthritis (AM-411) - IND expected in 2025</li> </ul>
Divesting / Partnering Legacy Assets	<ul><li>Unlock intrinsic value of inner ear &amp; OTC assets</li><li>Extra, non-dilutive funding potential</li></ul>



OligoPhore/SemaPhore are nanoparticles comprising a proprietary peptide + RNA payload designed to enable safe and effective delivery by systemic administration.



Stability	RNA complexed in nanoparticle format and only released inside of cells after uptake
Extrahepatic delivery	Not sequestered in liver as is common with conventional RNA- based therapies; permeates inflamed pathological tissues (passive targeting)
Endosomal escape	Efficient release within target cell, about 10-fold increase over LNPs, the current industry standard
Selectivity	Acts on targets in diseased tissues only
Safety	No cellular or adaptive immune responsivity to nanoparticle components or RNA after multiple serial doses, and no organ toxicities in mice



Exemplary listing of companies active in RNA therapeutics and delivery (list not exhaustive)

Silend	ce gene expre	ession	Promo	te protein exp	pression	Deliver RN	A therapeutic	to target
<ul><li>Short interfe</li><li>Antisense c</li></ul>	ering RNA (siRNA oligonucleotides (	a) (ASOs)	• Messenger	r RNA (mRNA)		<ul> <li>Lipid nanopa</li> <li>Virus-based</li> <li>Ligand conju</li> <li>Peptide-base</li> </ul>	articles vectors ugates <b>sed nanoparticl</b>	es
Alnylam*		AstraZeneca	ARCTURUS	AstraZeneca	BIONTECH	Sirna mics Advancing RNAi Therapeutics	<b>Paltamira</b>	Arbutus
IONIS	Lilly	U NOVARTIS	CUREVAC the RNA people®	Lilly	S MERCK	\$70 million	\$3.3 million	\$501 million
novo nordisk	ProQR THERAPEUTICS		moderna	novo nordisk	<b>P</b> fizer			PepGen
SILENCE THERAPEUTICS	sylentis	STOKE THERAPEUTICS	sanofi	<b>iiii Translate</b> BIO	ultragenyx	\$411 million	\$3.3 billion*	\$372 million

\*Represents valuation of the company derived from 2021 acquisition Figures are sourced from Yahoo Finance as of April 17, 2024

# **Disruptive Technology Growth Opportunities**





# **Corporate RNA Strategy**



# Strong strategy based on external collaborations and in-house programs

Leverage versatility of technology

- Demonstrated to work in multiple disease areas (tested in 17 models....)
- Suitable for siRNA, mRNA, ASOs, circular RNA
- Particularly well-suited for

indications in oncology and inflammatory disorders

# Selecting two therapeutic

indications to showcase technology

- KRAS driven cancers AM-401
- Rheumatoid arthritis AM-411
- Partner upon IND or Phase 1

# OligoPhore has been tested in vivo...

- Pancreatic and colorectal cancer (KRAS)
- Ovarian cancer (TAM: AXL)
- Lung cancer (ETV-2)
- Metastatic melanoma (NF- $\kappa$ B)
- Adult T cell leukemia/lymphoma (NFκB)
- Sarcoma (MYCT-1)
- Sarcoma and breast cancer (MYCT-1)

- Necrotizing enterocolitis (NF-κB)
- Rheumatoid and osteoarthritis (NF- $\kappa$ B)
- Atherosclerosis (JNK2)
- Metabolic syndrome/Obesity (ASXL2)
- Aortic aneurysm (NF-κB)

# SemaPhore<sup>™</sup> has been tested *in vivo*...

- Osteoarthritis (WNT16)
- Atherosclerosis (p27<sup>Kip1</sup>)
- Aortic aneurysm (SOD2)

- Osteoarthritis (DNMT3B)
- Tumor microenvironment (ZBTB46)

# Leveraging the Platforms



# License technology to biotechs / pharmas for use with their own RNA molecules



# AM-401: Stop the "Beating Heart" of Tumors



# Knock down various KRAS mutations with *poly*KRAS<sup>mut</sup> OligoPhore nanoparticles

to inhibit cell proliferation in KRAS driven colorectal, pancreatic, or non-small cell lung cancer.

- Mutated KRAS may cause cancer to grow
- Found in 1/5 of all human cancers, particularly in:
  - Pancreatic cancer (85-90%)
  - Colorectal cancer (40%)
  - Non-small cell lung cancer (30-35%)
- 150,000 cases diagnosed in US p.a.
- ~1M deaths per year world-wide
- Considered "undruggable" for decades

#### Many mutations known, G12D, G12V, and G12C accounting for >50%



#### OligoPhore *poly*KRAS<sup>mut</sup> siRNA transfects tumor cells, not healthy or uninvolved cells





#### OligoPhore *poly*KRAS<sup>mut</sup> significantly reduces pancreatic tumor volume growth

KPC pancreatic tumor model in mice; Strand et al., 2019



# AM-401

# KRAS driven cancer IND targeted for 2025

- High unmet medical need most aggressive tumors
- Small molecule G12C inhibitors approved in NSCLC
  - Sotorasib (Lumakras, Amgen), Adagrasib (Krazati, Mirati)

Multiple other small molecule inhibitors under development (G12C, G12D...), but few competing RNA projects (G12D or KRAS modulators)

## AM-401 KEY DIFFERENTIATING FACTORS



*poly*KRAS<sup>mut</sup> allows to target different mutations and is thus **polyvalent** G12C, G12V, G12D, G12R, G12A, and A146T, covering 90.9% of KRAS mutations in pancreatic, 65,3% in colorectal, 80.0% in non-small cell lung cancer



Blocking production of KRAS by degrading mRNA to cause **less resistance** than inhibition of KRAS



Small molecule inhibitors have significant side effects, particularly when combined with other agents

OligoPhore targets specifically tumor cells

# AM-411: Block Inflammation in Rheumatoid Arthritis



#### Knock down NF-κB (p65), a key checkpoint in RA inflammation.

- Chronic autoimmune disease
- Causes joint swelling and pain
   Reduced QoL and productivity
- Affects 1 out of 28 women / 59 men
- No cure available, but various treatment options:
  - Disease-modifying anti-rheumatic drugs (DMARDs)
  - Non-steroidal anti-inflammatory drugs (NSAIDs)
  - Corticosteroids
- Major shortcomings of therapies:
  - Drug resistance (up to 50% of patients)
  - Systemic adverse reactions (e.g., rash, hair loss, altered liver function, low blood cell counts, nausea, weight loss, increased infections, and neuropathy)



# OligoPhore p65 stabilizes ankle swelling and reduces arthritis score



# OligoPhore p65 reduces inflammation and protects against bone erosion



Collagen-antibody induced arthritis model in mice, Zhou et al., 2014.



# AM-411

Rheumatoid arthritis IND targeted for 2025

#### High unmet medical need

- Global rheumatoid arthritis market = \$57.9 Billion in 2019 → \$62.9 Billion in 2027
  - Expiration of patents, biosimilars arriving
  - High hopes for novel Tx class of JAK inhibitors gave way to disappointment due to safety issues

# AM-411 KEY DIFFERENTIATING FACTORS



Mediators of inflammation play many physiological roles in healthy tissues – AM-411 targets only inflamed tissues Reduced systemic side effects



Blocking production of an NF- $\kappa$ B component by degrading mRNA to cause less resistance than inhibition of NF- $\kappa$ B Less likelihood of resistance

## **Intellectual Property**



(12) United States Patent Wickline et al.			(10) Patent (45) Date o	(10) Patent No.: US 9,987,371 B2 (45) Date of Patent: Jun. 5, 2018		
:4)	COMPOS POLYNU	ITIONS AND METHODS F CLEOTIDE TRANSFECTIO	DR 8,501,930 B2 N 8,617,516 B2 2005/0191746 A1	<ol> <li>8/2013 Rozema et al.</li> <li>12/2013 Wickline et al.</li> <li>9/2005 Van</li></ol>		
1)	Applicant:	Washington University, St. 1 (US)	ouis, MO 2007/0275923 A1 2011/0123438 A1	435/455 11/2007 Chen et al. 5/2011 Wickline et al.		
2)	Inventors:	Samuel A. Wiekline, St. Lou (US); Kirk Hou, St. Louis, M	s, MO FORE IO (US)	IGN PATENT DOCUMENTS		
3)	Assignce:	WASHINGTON UNIVERSI Louis, MO (US)	WO 2005/ TY, Saint WO 2007/ WO 2011/ WO 2011	085458 A2 9/2005 069090 A2 6/2007 020188 A1 2/2011 107596 A1 7/2014		
9	Notice:	Subject to any disclaimer, the patent is extended or adjuste U.S.C. 154(b) by 0 days. day	erm of this WO 2017 1 under 35	004512 A1 1/2017 DTHER PUBLICATIONS		
1)	Appl. No.:	14/790,408	Wu et al., 2012, R	ecent progress in copolymer-mediated siRNA		
2)	Filed:	Jul. 2, 2015	delivery, Journal of Noguchi et al., 200 Therapeutic Desene	Drug Targeting, 20(7): 551-560.* 6, Protein Transduction Technology: A Novel cline, 60(1): 1,11.*		
5)		Prior Publication Data	Examination Report	t for related CA application 2,896,834 dated		
	US 2015/0	314013 A1 Nov. 5, 2015	Aug. 23, 2016, 5 pr Partial Supplements	ages. 1ry European Search Report dated Aug. 9, 2016		
	Rel	ated U.S. Application Data	from related EP Ap International Search	plication No. 14735277.7, 10 pages. a Report and Written Opinion dated Oct. 4, 2016 Metert Application No. INTELESD16/04/0679-10.		
3)	Continuati PCT/US20	on-in-part of applicati 14/010212, filed on Jan. 3, 20	n No. pgs. 4. Salomone F. et al.,	"In Vitro Efficient Transfection by CM18-Tat11		
D)	Provisiona 3, 2013, pr on Aug. 61/873,18	application No. 61/748,615, f ovisional application No. 61/86 23, 2013, provisional appli , filed on Sep. 3, 2013.	Hybrid Peptide: A PLoS ONE, Jul. 29 A nove ation No. system for STAT3 si The FASEB Journa Units and Mithal	New Tool for Gene-Delivery Applications," 2013, pp. 1-11, vol. 8, No. 7, e70108. i melliti-devived psptide nanoparticle delivery iRNA mediated killing of B16 melanoma cells," 4, 2012, vol. 26, No. 1. Via Daviend Devider, for Nanoerlide, Based		
1)	Int. Cl. <i>C07K 19/</i>	Ø (2006.01)	Hou, et al., "Melli siRNA Transfection 34, No. 12.	tin Derived Peptides for Nanoparticle Based ," Biomaterials, Apr. 2013, pp. 3110-3119, vol.		
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8)	Field of C CPC C	lassification Search 07K 14/00; A61K 47/48315; A	61K 38/16 (74) Attorney, Ag	r — Amber D Steele gent, or Firm — Polsinelli PC		
	See applic	ation file for complete search l	530/320 istory.			
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#### WORLDWIDE EXCLUSIVE LICENSE FROM WASHINGTON UNIVERSITY Patent covering OligoPhore™ / SemaPhore™ platform



Compositions comprising a peptide-polynucleotide complex



Methods for delivering such nanoplexes



Coverage until 2034 (+ potential extension)



Generating further IP (filed e.g. *poly*KRAS<sup>mut</sup> – potential coverage until 2043)



Proprietary manufacturing process

# **Management Overview**







# **Bentrio® in Allergic Rhinitis**

#### Protection Against Airborne Particles

- Drug-free, preservative-free formulation, applied as nasal spray
- Four clinical trials demonstrating safety and efficacy in allergic rhinitis
  - Efficacy: close to medicated sprays
  - Tolerability: close to saline sprays
- Commercialized through distributors
- Significant growth expected
  - Launch in additional countries / regions
- Advanced discussions on North America, Europe and other key markets



#### First Step in Transition Process

- Sale of 51% of Altamira Medica AG
  - Cash consideration about \$2.3 million
  - Buyer is Swiss private equity investor
  - CYTO retaining 49% of capital
- CYTO also entitled to 25% of:
  - Future license income
  - Medica's value appreciation in case of a sale
- CYTO's overall share of upside: 62%
- Financial gain CHF 5.2 million
- Going forward: reduction in Bentrio related expenditures

# Legacy Programs: Inner Ear Assets to be Divested / Partnered





#### Become focused "Pure play" RNA delivery company

#### AM-125 in Acute Vestibular Syndrome

- Rx product, applied as nasal spray
- Reformulation of oral betahistine
  - Global market \$450M (ex US) standard of care for vertigo
  - Poor bioavailability
- Invested \$18 million to date
- Proof of concept in Phase 2, ready for Phase 3 trial
- No comparable product in US
- Structured partnering process initiated





# Monetize legacy assets through divestiture, out-licensing

#### Potential Other Indications

- Histamine plays important role in many behavioral and physiological functions:
  - Appetite, drinking, sleep, wakefulness, learning, attention and memory
- Clinical utility of betahistine shown, among others, in:
  - ADHD, cognitive function in dementia, memory loss, antipsychotic-induced weight gain
- Histamine as target, e.g.:
  - Narcolepsy, Tourette syndrome, Prader-Willi syndrome

# **Investor Summary**





## RNA technology coming of age

- Disruptive potential in human medicine
- Rapidly growing # of RNA therapeutics



#### Extensive proof of concept

- Successfully tested *in vivo* in 17 different disease models
- 30+ papers published



# Altamira has unique, versatile RNA delivery technology platform



- Patented, under license from Wash U
- Suitable for different types of RNA molecules
- OligoPhore<sup>™</sup> and SemaPhore<sup>™</sup>

# Flagship programs in oncology and rheumatoid arthritis

- First IND expected to be filed in 2024
- Technology platform out-licensing as business model



# Addressing major challenges in RNA delivery

- IV administration, reaching extrahepatic targets
- Strong endosomal release (10x compared to lipid nanoparticles)



### Divestiture/partnering of Legacy Assets

- Process started
- Unlock intrinsic value / non-dilutive funding

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