UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 8-K/A

CURRENT REPORT Pursuant to Section 13 OR 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): May 11, 2022

PIERIS PHARMACEUTICALS, INC.

(Exact name of registrant as specified in its charter)

	Nevada (State or other jurisdiction of Incorporation)	001-374 (Commis File Numl	ion	30-0784346 (IRS Employer Identification No.)				
		255 State Street, 9th Floor	02109					
		Boston, MA						
		(Address of principal executive offices)	(Zip Code)					
		Registrant's telephone number, incl	uding area code: 857-246-8998					
		N/A						
		(Former name or former address,	if changed since last report.)					
Check the	appropriate box below if the Form 8-K filing is intend	ed to simultaneously satisfy the filing obligation of the	registrant under any of the following provision	ons:				
	□ Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)							
	□ Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)							
	Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))							
	Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))							
Securities re	egistered pursuant to Section 12(b) of the Act:							
	Title of each class	Trading	Symbol(s)	Name of each exchange on which registered				
	Common Stock, \$0.001 par value per share		PIRS	The Nasdaq Capital Market				

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (17 CFR §230.405) or Rule 12b-2 of the Securities Exchange Act of 1934 (17 CFR §240.12b-2).

Emerging Growth Company \square

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

EXPLANATORY NOTE								
The sole purpose of this Form 8-K/A is to amend the Form 8-K, filed by Pieris Pharmaceuticals, Inc. with the Securities and Exchange Commission on May 11, 2022, to include an updated Ex. 99.2 - Investor Presentation,								
dated May 2022. No other changes have been made.								

Item 7.01 Regulation FD Disclosure.

Furnished hereto as Exhibit 99.2 is the May 2022 Investor Presentation of the Company.

The information set forth under this "Item 7.01. Regulation FD Disclosure," including Exhibit 99.2 furnished hereto, shall not be deemed "filed" for any purpose, and shall not be deemed incorporated by reference into any filing under the Securities Act of 1933, as amended, or the Securities Exchange Act of 1934, as amended, regardless of any general incorporation language in any such filing except as shall be expressly set forth by specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

99.2 <u>Investor Presentation, dated May 2022</u>.

104 Cover Page Interactive Data File (embedded within the Inline XBRL document).

SIGNATURE

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

PIERIS PHARMACEUTICALS, INC.

Dated: May 11, 2022

/s/ Tom Bures
Tom Bures
Chief Financial Officer

PIERIS PHARMACEUTICALS



CORPORATE PRESENTATION
May 2022



Forward-Looking Statements

This presentation contains forward-looking statements as that term is defined in Section 27A of the Securities Act of 1934, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. Statements in this presentation that are not purely historical are forward-looking statements. Such forward-looking statements include, among other things, the timing for initiation of clinical trials of PRS-220, whether PRS-220 will provide a clinical need in HER2 gastric cancer patients who do not respond to traditional HER2-targeted therapies; whether therapies could address a high medical need in HER2 gastric cancer patients who do not respond to traditional HER2-targeted therapies; whether the effects of the combination of cinrebafusp alfa with other therapies some in precinical studies will be observed in clinical trials; the receipt for royal yaments provided for in our collaboration agreements; the expected timing and potential outcomes of the reporting by the Company of key clinical data from its programs, references to novel technologies and methods and our business and product development plans, including the Company's cash resources, the advancement of our proprietary and co-development programs including PRS-660/AZD1402, cinrebafusp affa, PRS-344/S09512, PRS-352/S095025 and PRS-342/BOS-342; our continued progress in the areas of co-stim bispecifics and inhaled therapeutics; and the advancement of our developmental programs generally. Actual results could differ from those projected in any forward-looking statements due to numerous factors. Such factors include, among others, our ability to raise the additional funding we will need to continue to pursue our business and product development plans; the inherent uncertainties associated with developing mey products or technologies and operating as a development stage company; our ability to evelope, complete clinical trials for, obtain approvals for and commercialize any of our product candidates, including our ability to recruit and enroll patients in



Our Formula for Success

We combine leading protein engineering capabilities and deep insights into molecular drivers of disease to develop medicines that drive local biology to produce superior clinical outcomes for patients.





Executive Summary

Superior Medicines via Efficient Biology

- Protein therapeutics that exploit biology validated by mAbs yet are engineered for focused activity at disease locus
 Improved activity, reduced side effects, increased convenience

Two Focus Areas

- Oral inhaled antagonists for respiratory disease
 Locally activated immuno-oncology bispecifics
 Multiple near-term catalysts

Supportive Partnerships

- ~\$200M since 2017 in upfronts, milestones and equity investments
- Several co-developed and out-licensed programs
 Clinical supply for combination studies and development expertise



Anticalin® Proteins as Therapeutic Modalities

A Novel Therapeutic Class with Favorable Drug-Like Properties

- Human Derived from lipocalins (human extracellular binding proteins)
- Small Monomeric, monovalent, small size (~18 kDa vs. ~150kDa mAbs)
- Stable Inhalable delivery
- Simple Bi/multispecific constructs
- Proprietary Broad IP position on platform and derived products

Translational Science Expertise to Deploy Platform in Meaningful Way

- Immunology expertise underpins IO and respiratory focus
- A leader in 4-1BB and costim biology
- Patient phenotyping efforts for improved stratification and novel intervention points in, e.g., asthma



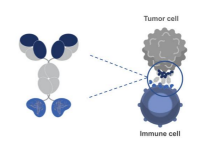


Two-fold Focus of Anticalin Platform Deployment

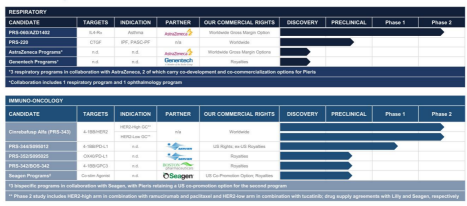
Inhalable formulations to treat respiratory diseases locally



Bispecifics for local immune agonism to treat cancer



Our Pipeline





Validating Partnerships with Leading Companies



- PRS-060/AZD1402 + 3 additional programs
 Upfront & milestones to date: \$70.5M
 \$10M equity investment from AstraZeneca
 Eligible to receive over \$5.0B in potential
 milestone payments plus royalties
 Retained co-development and cocommercialization (US) options on PRS-060
 and up to 2 additional programs

BOSTON

- Boston Pharmaceuticals holds exclusive license for PRS-342/BOS-342
 Upfront & milestones to date: \$10M
 Eligible to receive up to approximately \$353M in potential milestone payments
 Entitled to tiered royalties

Genentech

- 1 respiratory program + 1 ophthalmology program
 Upfront & milestones to date: \$20M Eligible to receive over \$1.4B million in potential milestone payments
 Entitled to liered royalities
 Genentech has option to select additional targets in return for an option exercise fee

Seagen

- 3-program IO bispecific partnership
 Upfront & milestones to date: \$35M
 Eligible to receive up to approximately
 \$1.28 in potential milestone payments plus royalties
 \$13M equity investment from Seagen
 Tucatinib drug supply for phase 2 combination trial of cinrebafusp alfa in HER2-low gastric cancer



- PRS-344/S095012: PD-L1/4-18B antibody-Anticalin bispecific, for which Pieris holds full U.S. rights
 Upfront & milestones to date: ~\$41M
 Eligible to receive up to approximately \$230M in potential milestone payments
 Entitled to tiered royalties



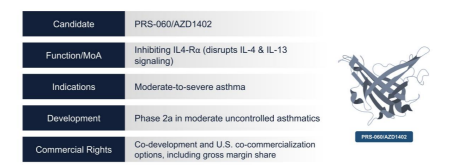


- Lipocalin templates deployed by Pieris in respiratory programs are abundant in the human lung and can permeate lung epithelium
- Stable, monovalent molecules with high melting temperatures and insensitivity to mechanical stress
- Inhalation pharmacokinetics suitable for once or twice daily administration and compatible with flexible treatment regimens
- Control of particle size distribution in critical size range in both "wet" and "dry" formulations to enable tailored delivery to discrete lung regions



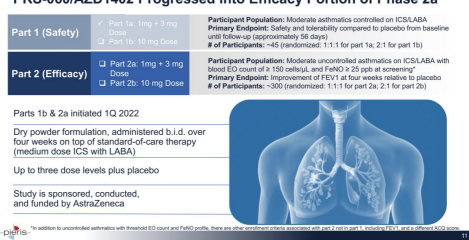


PRS-060/AZD1402: Inhaled IL-4Rα Antagonist





PRS-060/AZD1402 Progressed into Efficacy Portion of Phase 2a



DPI Formulation of PRS-060/AZD1402 Passed Safety Review

31 moderate asthmatics controlled on standard-of-care (medium dose ICS with LABA) asthma therapy were dosed twice daily over four weeks randomized across two dose levels and placebo arm (1:1:1)

Safety review successfully completed for two different dose levels that will now be explored for efficacy in participants with asthma uncontrolled on medium dose ICS-LABA Safety review performed of the following (compared to placebo):

Incidence of adverse events

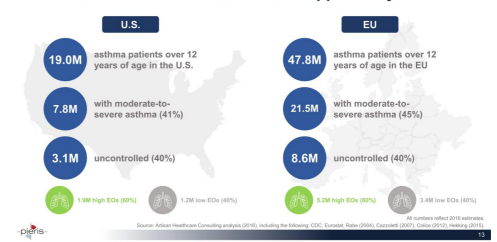
Changes in laboratory markers (immune biomarkers, clinical chemistry, and hematology)

Forced expiratory volume in 1 second (FEV1)

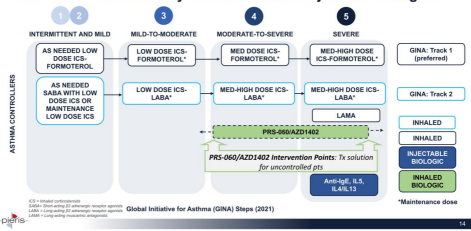
Pharmacokinetics



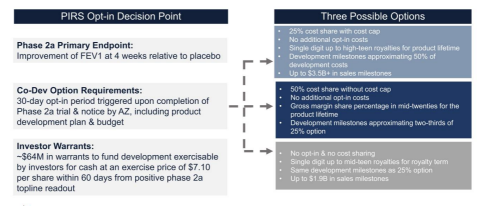
Moderate-to-Severe Asthma Market Opportunity



Potential Large Market Opportunity in Moderate-to-Severe Asthma not Addressed by ICS/LABA before Injectable Biologics



Co-Development Options for PRS-060/AZD1402





PRS-220: Inhaled CTGF Antagonist



*Idiopathic pulmonary fibrosis and post-acute sequelae of SARS-CoV-2 infection (PASC) pulmonary fibrosis



IPF: High Unmet Medical Need and Significant Commercial Opportunity

IPF is a chronic, progressive, and ultimately fatal lung disease of unknown cause characterized by chronic lung inflammation and progressive scarring (fibrosis) of the tissues between the alveoli of the lung



3 to 5 million

people affected worldwide with increasing global incidence, with ~130K affected in the US each year^{1,2}

2 to 5 mean survival from the time of diagnosis²



current market in sales

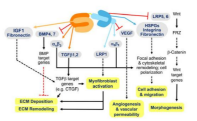
Currently approved treatments provide modest benefit, in addition to having side effects that require management



1 - Glassberg, AJMC 2015

CTGF: Clinically Validated Intervention for IPF

- Connective tissue growth factor (CTGF), or CCN2, a protein localized in the extracellular matrix, is a driver of fibrotic remodeling as consequence of an aberrant wound healing process
- Over-expression of the protein in lung tissue is observed in patients suffering from IPF
- Competitor clinical data indicate inhibition of CTGF reduces the decline in lung function among patients with IPF
- Competitor compound requires highdose infusions to effectively target lungresident CTGF



CTGF affects multiple signaling pathways and processes important in pathophysiology. CTGF interacts with a variety of molecules, including cytokines and growth factors, receptors and matrix proteins. These interactions alter signal transduction pathways, either positively or negatively, which results in changes in collutar reargners.

(Lipson, Fibrogenesis & Tissue Repair, 2012)



PRS-220: Inhaled Solution

The only CTGF inhibitor in clinical trials for IPF is a monoclonal antibody administered by IV infusion, 30 mg/kg every three weeks

The objective of PRS-220 is to more efficiently engage a clinically validated target via oral inhalation directly to the lung epithelium and interstitium

Benefits of inhaled administration:

- Inhaled administration eliminates the need for additional clinic visits required for systemic drug administration
- Direct administration into the lungs may result in more efficient CTGF inhibition in the site of the disease
- Patients with IPF frequently take inhaled medications and thus no additional training required
- This approach supports add-on to SOC, whereas patients on SOC are excluded from current studies of reference mAh



Grant from Bavarian Government to Support Program Acceleration and Evaluation of Efficacy in PASC-PF



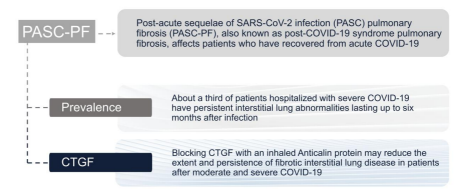
approximately 14 million euro grant from the Bavarian government for the research and development of PRS-220 for post-acute sequelae of SARS-CoV-2 infection (PASC) pulmonary fibrosis (PASC-PF)

Grant will:

- Allow Pieris to accelerate development of the program IND planned 2022
- Support clinical-readiness activities and initial clinical development for the program, including GLP tox studies, GMP manufacturing, and phase 1 clinical development
- Broaden scope of the program beyond the original IPF indication by including the evaluation of PRS-220 for the treatment of post-COVID-19-related pulmonary fibrosis

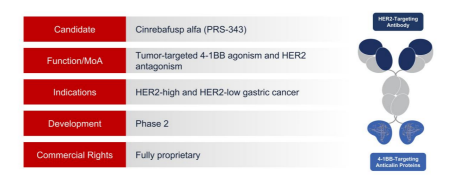


PRS-220 for PASC-PF





Cinrebafusp Alfa (PRS-343): Lead IO Asset

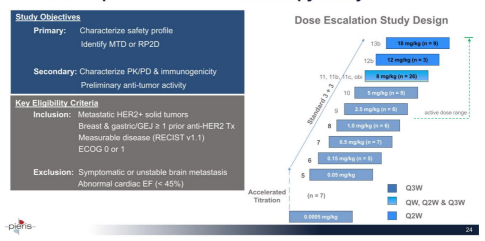








Cinrebafusp Alfa Phase 1 Monotherapy Study



Phase 1 Monotherapy Treatment-related Adverse Events at Active Doses (≥ 2.5 mg/kg)

reatment-related Adverse Events (TRAEs occurring in > 1 patient; n = 53)	All Grades n (%)	Grade 1-2 n (%)	Grade 3-4 n (%)
Infusion-related reaction	13 (25%)	9 (17%)	4 (8%)
Nausea	7 (13%)	7 (13%)	1 (070)
Chills	6 (11%)	6 (11%)	
Vomiting	6 (11%)	6 (11%)	
Dyspnea	4 (8%)	4 (8%)	
Fatigue	4 (8%)	4 (8%)	
Arthralgia	3 (6%)	2 (4%)	1 (2%)
Decreased appetite	3 (6%)	3 (6%)	
Non-cardiac chest pain	3 (6%)	3 (6%)	
Asthenia	2 (4%)	2 (4%)	
Diarrhea	2 (4%)	2 (4%)	
Dizziness	2 (4%)	2 (4%)	
Headache	2 (4%)	2 (4%)	
Paresthesia	2 (4%)	1 (2%)	1 (2%)
Pruritus	2 (4%)	2 (4%)	
Pyrexia	2 (4%)	2 (4%)	
Rash	2 (4%)	2 (4%)	

1 Gr 3 Ejection Fraction dec and 1 Gr 3 Heart Failure; both events occurred in one patient and resolved w/o sequelae.

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Data cut-off: 25-Feb-21

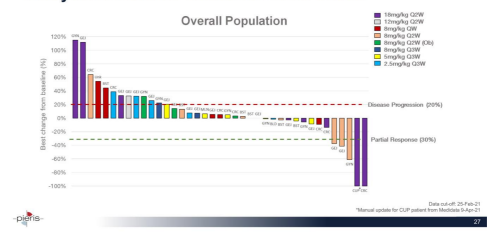
Summary of Responses in Phase 1 Monotherapy Study

Cohort	13b 18 mg/kg, Q2W	12b 12 mg/kg, Q2W	11c 8 mg/kg, QW	Obi 8 mg/kg, Q2W	11b 8 mg/kg, Q2W	11 8 mg/kg, Q3W	10 5 mg/kg, Q3W	9 2.5 mg/kg, Q3W	Total
Best Response									
Evaluable Patients	8	2	5	4	7	4	7	5	42
CR	1	÷	-	-	-	-	-	-	1
PR	1	-	-	-	3	-	-	-	4
SD	3	-	1	2	3	3	3	2	17
ORR	25%	0%	0%	0%	43%	0%	0%	0%	12%
DCR	63%	0%	20%	50%	86%	75%	43%	40%	52%

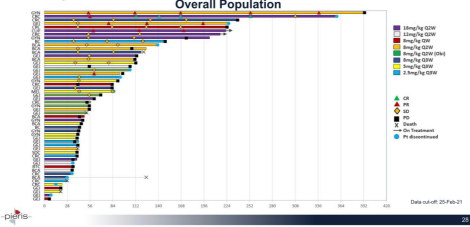


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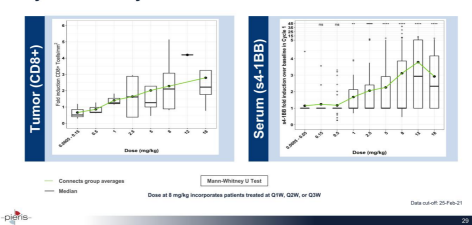
Cinrebafusp Alfa Phase 1 Monotherapy Efficacy Data: Analysis of Patients Treated at Active Doses



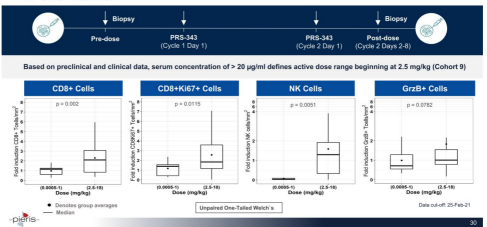
Durable Responses with Cinrebafusp Alfa Among Heavily Pre-treated Population Overall Population



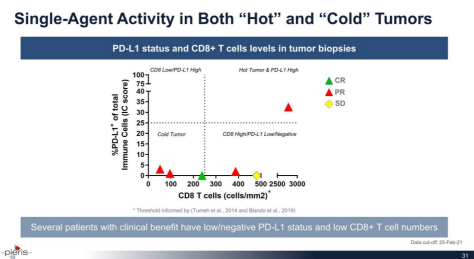
Cinrebafusp Alfa Shows Dose-dependent Activity Across Key Pharmacodynamic Parameters



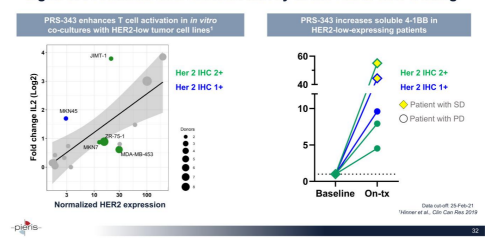
Cinrebafusp Alfa Activates Adaptive and Innate Immunity in the Tumor



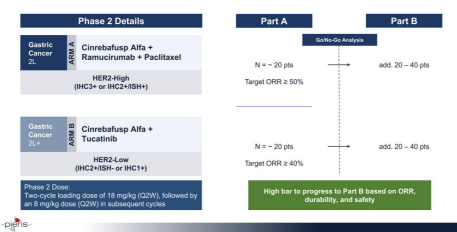
Single-Agent Activity in Both "Hot" and "Cold" Tumors



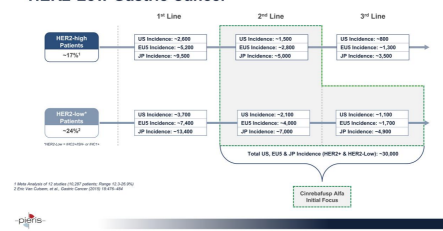
Signs of Preclinical and Clinical Activity in the HER2-Low Setting



Cinrebafusp Alfa Clinical Development Plan



Cinrebafusp Alfa Opportunity in HER2-High & HER2-Low Gastric Cancer

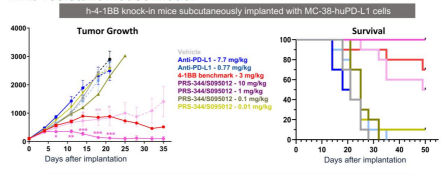


PRS-344/S095012: Meaningfully Building on Localized MoA of Cinrebafusp Alfa





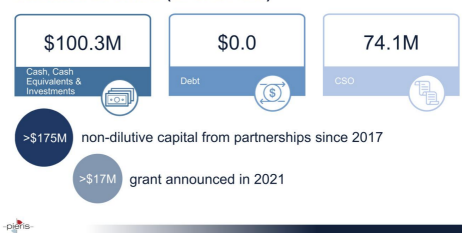
PRS-344/S095012 Drives Strong Anti-tumor Activity in Anti-PD-L1 mAb-resistant Mouse Model



Dose-dependent anti-tumor response that leads to significant extension of survival
 Superior to equimolar doses of anti-PD-L1 mAb treatment alone



Financial Overview (as of 3/31/22)







PRS-060 Phase 1 Multiple Ascending Dose Trial

Strategic Objectives Ascertain PK/PD with a reliable biomarker to confirm local target engagement and inform Phase 2 dosage regimen

Trial Design Highlights

Dosing patients with mild asthma with elevated FeNO levels (≥35 ppb), to receive inhaled PRS-060 or pbo b.i.d.* over a 10-day period

*q.d. on Day 10

Initiated in July 2018

Evaluating safety, tolerability, PK, and PD and will also evaluate FeNO reduction vs. placebo

Measuring safety, tolerability and FeNO changes days 1-10, 17, and

Pieris is sponsoring the trial; AstraZeneca is reimbursing Pieris for all associated costs





Phase 1b Interim Results: Favorable Safety Profile

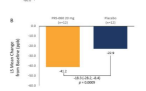
- All doses of AZD1402/PRS-060 tested in the study were well tolerated
- No treatment-related serious AEs were observed

System organ class AE Preferred Terms ^b	Placebo (N = 12) n (%) m	AZD1402/PRS-060° (N = 30) n (%) m	Overall (N = 42) n (%) m
Gastrointestinal disorders	4 (33.3) 4	13 (43.4) 14	17 (40.5) 18
Dry mouth	1 (8.3) 1	2 (6.7) 2	3 (7.1) 3
Nausea	1 (8.3) 1	3 (10.0) 3	4 (9.5) 4
Infections and infestations Upper respiratory tract infection	1 (8.3) 1	7 (23.3) 8	8 (19.0) 9
	1 (8.3) 1	3 (10.0) 4	4 (9.5) 5
Nervous system disorders Headache Presyncope	5 (41.7) 9 3 (25.0) 6 0	13 (43.4) 18 5 (16.7) 7 4 (13.3) 6	18 (42.9) 27 8 (19.0) 13 4 (9.5) 6
Respiratory, thoracic and mediastinal disorders	6 (50.0) 6	14 (46.7) 15	20 (47.6) 21
Cough	1 (8.3) 1	4 (13.3) 4	5 (11.9) 5
Rhinorrhoea	2 (16.7) 2	1 (3.3) 1	3 (7.1) 3
Wheezing	2 (16.7) 2	4 (13.3) 5	6 (14.3) 7



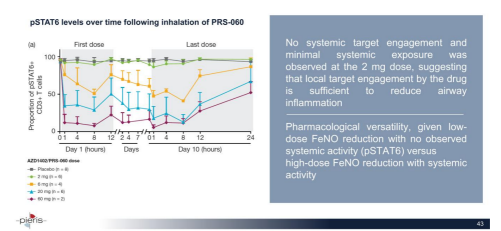
Phase 1b Interim Results: Robust FeNO Reduction

PRS-060 Relative FeNO Reduction (ANCOVA Analysis)





Phase 1b Interim Results: Pharmacological Versatility



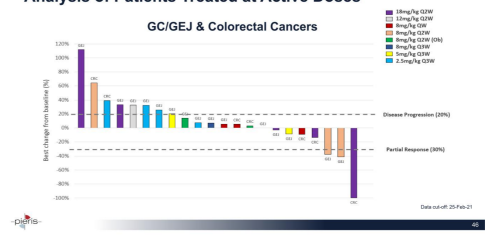


Phase 1 Monotherapy Baseline Characteristics (N = 78)

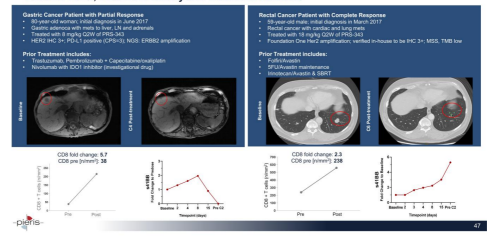
Characteristic	n (%)	Primary Cancer Type	n (%)	
Age, Median (range)	63 (24–92)	Gastroesophageal	34 (44%)	
Gender		Gastioesopriagear	34 (44%)	
F	46 (59%)	Breast	16 (21%)	
M	32 (41%)	D. Oddi	10 (2170)	
ECOG PS		Colorectal	12 (15%)	
0	19 (24%)			
1	59 (76%)	Gynecological	9 (12%)	
Prior Therapy Lines		Province:		
1	11 (14%)	Bladder	2 (3%)	
2	10 (13%)	Pancreatic	4 (49/)	
3	16 (21%)	Pancieatic	1 (1%)	
4	12 (15%)	Other - Cancer	2 (3%)	
5+	29 (37%)	of Unknown Origin		
Median # of anti-HER2 Tx		Other - Salivary Duct	1 (1%)	
Breast	6			
Gastric	2	Melanoma	1 (1%)	
			Data cut-off: 25-Feb	

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Cinrebafusp Alfa Phase 1 Monotherapy Efficacy Data: Analysis of Patients Treated at Active Doses



Case Studies: PR in Gastric Cancer and CR in Rectal Cancer Patient Profile, Treatment History and Treatment Outcome



Case Study: PR in Cancer of Unknown Primary Patient Profile, Treatment History and Treatment Outcome



Lesions	Lesion Site	Lesion Size (mm)			
		Pre-treatment	Post-treatment		
		Pre-treatment	Cycle 2	Cycle 4	Cycle 6
Target 1	Lung, right lower lobe mass	25	13	0	0
	Total	25	13	0	0
	% Change from Baseline		-48%	-100%	-100%
Non-target 1	Lung, bilateral pulmonary masses	Present	Not assessed	Present	Present
Non-target 2	Lymph nodes, mediastinal and hilar	Present	Not assessed	Present	Present
Overall Response			PR	PR	PR

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Data cut-off: 25-Feb-21

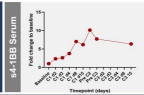
Case Study: SD in Colorectal Cancer Patient Profile, Treatment History and Treatment Outcome

Patient Profile

56-year-old female Initial diagnosis Jan 2009 Stage 4 Colorectal Adenocarcinoma Cancer Archival HER2 3+ MSI stable; KRAS, NRAS, BRAF wt

Treatment History

9 prior lines of therapy, including: Folfiri Folfox + Avastin 5-FU + bevacizumab trastuzumab/pertuzumab Investigational agent (immune stimulator antibody conjugate (ISAC) with antibody similar to trastuzumab



	Lesion Site	Lesion Size (mm)			
Lesions		Pre-treatment	Post-treatment		
		Pre-treatment	Cycle 2	Cycle 4	Cycle 6*
Target 1	Lung, right upper lobe pulmonary nodule	10	8	8	
Target 2	Lung, right lower lobe pulmonary nodule	12	11	11	
	Total	22	19	19	-
	% Change from Baseline		-14%	-14%	
Non-target 1	Lung, multiple pulmonary nodules	Present	Present	Present	
CEA		<1.9	1.1	1.3	

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Soluble 4-1BB (s4-1BB): Blood-based Biomarker of Cinrebafusp Alfa Engagement

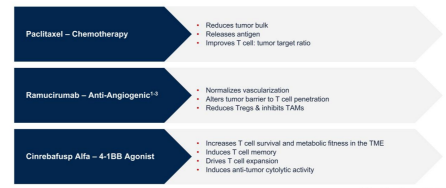
- s4-1BB is an alternatively spliced form of 4-1BB receptor lacking the transmembrane encoding exon (Setareh et al., 1995; Shao et al., 2008)
- s4-1BB is **released by leukocytes in an activation-dependent manner** (Michel et al., 2000; Salih et al., 2001; Schwarz et al., 1996)
- s4-1BB is **produced with a slightly delayed kinetic to pathway activation**. Hypothesized role as a negative regulator, keeping 4-1BB-mediated co-stimulation in check

s4-1BB utility as a pathway specific biomarker provides ability to track cinrebafusp target engagement and activity using serum samples





Scientific Rationale for Combining Cinrebafusp Alfa & SoC



Allen et al., Science Translational Medicine 201
 Juang et al., Front Immunology 201
 Tada et al., Journal for Immunotherapy of Cancer 201



Scientific Rationale for Combining Cinrebafusp Alfa & Tucatinib

Tyrosine kinase inhibitors
(tucatinib)

- Upregulates or stabilizes tumor cell surface HER2
expression^{2.3.4}
- Increases clustering potential of cinrebafusp alfa on tumor
cells to drive enhanced 4-1BB cross-linking

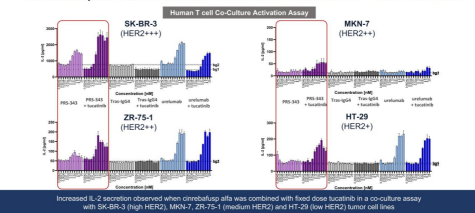
- Inhibits HER2 signaling AND activates tumor-specific T cells
in tumor microenvironment

- Enhances inhibition of HER2 signaling by concurrent binding
to HER2 on the tumor cell surface and TKI inhibition of the
internal kinase signaling domain¹
- In vitro, combination leads to significantly increased T cell
activation in the presence of HER2-Low cell lines

Baselga J., Lancet, 201
 Haruyama T., et al, Anticancer Res., 201
 S- Scalirtii M., et al, Oncogene, 200
 Hartmans, et al. Oncotarget, 201

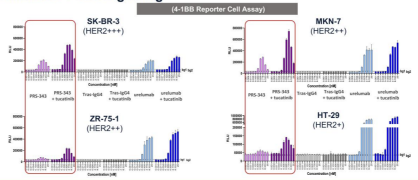


Cinrebafusp Alfa and Tucatinib Combination Enhances T cell Activation



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Cinrebafusp Alfa and Tucatinib Combination Leads to Enhanced 4-1BB Signaling



Increased 4-1BB signaling observed when cinrebafusp alfa was combined with fixed dose tucatinib in a reporter assay with SK-BR-3 (high HER2), MKN-7, ZR-75-1 (medium HER2) and HT-29 (low HER2) tumor cell lines



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