#### UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

#### FORM 8-K

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

Date of Report (Date of earliest event reported): June 8, 2022

#### PIERIS PHARMACEUTICALS, INC.

(Exact name of registrant as specified in its charter)

	Nevada (State or other jurisdiction of Incorporation)	001-37471 (Commission File Number	n	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, includ N/A					
		(Former name or former address, if	changed since last report.)				
Check th	e appropriate box below if the Form 8	s-K filing is intended to simultaneously satisfy t	he filing obligation of the re	gistrant under any of the following provisions:			
	Written communications pursuant to	Rule 425 under the Securities Act (17 CFR 23	0.425)				
	Soliciting material pursuant to Rule	14a-12 under the Exchange Act (17 CFR 240.1	4a-12)				
	Pre-commencement communication	s pursuant to Rule 14d-2(b) under the Exchange	e Act (17 CFR 240.14d-2(b))				
	Pre-commencement communication	s pursuant to Rule 13e-4(c) under the Exchange	Act (17 CFR 240.13e-4(c))				
Securities	s registered pursuant to Section 12(b) of t	he Act:					
	Title of each class	Trading Sym	bol(s)	Name of each exchange on which registered			
	Common Stock, \$0.001 par value per s	hare 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. $\Box$							

#### Item 7.01 Regulation FD Disclosure.

Furnished hereto as Exhibit 99.1 is the June 2022 Investor Presentation of Pieris Pharmaceuticals, Inc.

The information set forth under this "Item 7.01. Regulation FD Disclosure," including Exhibit 99.1 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.

# (d) Exhibits. 99.1 Investor Presentation, dated June 2022. 104 Cover Page Interactive Data File (embedded within the Inline XBRL document).

Item 9.01 Financial Statements and Exhibits.

#### 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: June 8, 2022

/s/ Tom Bures

Tom Bures

Chief Financial Officer

### PIERIS PHARMACEUTICALS



CORPORATE PRESENTATION
June 2022



#### **Forward-Looking Statements**

This presentation contains forward-looking statements as that term is defined in Section 27A of the Securities Act of 1933, 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 benefit in the treatment of IPF and PASC-related fibrosis; whether the combination of cinrebafusp alfa with other 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 seen in preclinical studies will be observed in clinical trials; the receipt of royalty payments 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 into and through the clinic and the expected timing for reporting data, making IND filings or achieving other milestones related to our programs, including PRS-060/AZD1402, cinrebafusp alfa, PRS-344/S095012, PRS-352/S095025, PRS-342/BOS-342, and PRS-400; our continued progress in the areas of co-stim bispecifics and inhaled therapeutics; the potential addressable market for our product candidates; 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 new products or technologies and operating as a development stage company; our ability to develop, complete clinical trials for, obtain approvals for and commercialize any of our product candidates, including our ability to recruit and enroll patients in our studies; our ability to address the requests of the U.S. Food and Drug Administration; competition in the industry in which we operate; delays or disruptions due to COVID-19 or geo-political issues, including the conflict in Ukraine; and market conditions. These forward-looking statements are made as of the date of this presentation, and we assume no obligation to update the forward-looking statements, or to update the reasons why actual results could differ from those projected in the forward-looking statements, except as required by law. Investors should consult all of the information set forth herein and should also refer to the risk factor disclosure set forth in the reports and other documents we file with the Securities and Exchange Commission available at www.sec.gov, including without limitation the Company's Annual Report on Form 10-K for the fiscal year ended December 31, 2021 and the Company's subsequent Quarterly Reports on Form 10-Q.



#### **Executive Summary**

### Proven Discovery Platform

- 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 immunooncology bispecifics

### Industry & Clinical Validation

- ~\$200M since 2017 in upfronts, milestones and equity investments
- Several co-developed and out-licensed programs
- Proven clinical activity for both focus areas

#### Value Proposition

- Four clinical-stage assets expected by year-end 2022
- Three are funded ~ ≥ 50% by partners or grant income
- · Retained US or WW rights for each program
- Five clinical readouts anticipated through 2023



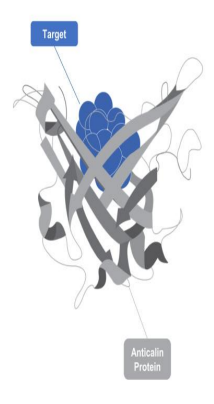
#### **Anticalin® Proteins as Therapeutic Modalities**

#### 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 Strong IP position on platform and derived products

#### Translational Science Expertise to Deploy Platform in Meaningful Way

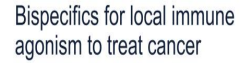
- Immunology expertise underpins IO and respiratory focus
- A leader in 4-1BB and costim biology
- Patient phenotyping efforts for improved stratification and novel intervention points in, e.g., asthma



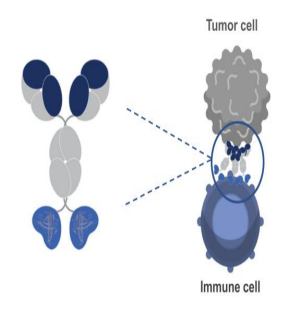


#### **Two-fold Focus of Anticalin Platform Deployment**

Inhalable formulations to treat respiratory diseases locally







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#### **Validating Partnerships & Non-Dilutive Capital**

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AstraZeneca	S





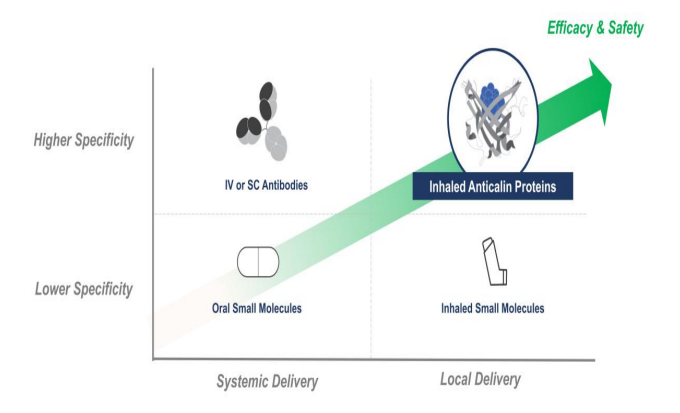




Number of Programs	Cash to Date	Cash Potential		
Four (three with co-dev options)	\$70.5M	>\$5B plus royalties		
Two	\$20M	>\$1.4B plus royalties		
Two (one co-dev program)	~\$41M	~\$230M plus royalties		
Three	\$35M	\$1.2B plus royalties		
One	\$10M	~\$353M		



#### **Combined Advantages of Higher Specificity with Local Delivery**



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#### **Respiratory Pipeline**

Program	Target	Indication	Discovery	Preclinical	Phase 1	Phase 2	Partner
PRS-060/AZD1402*	IL4Rα	Asthma	Phase 2a full	y sponsored by	AZ; co-dev option	on	AstraZeneca 🕏
PRS-220	CTGF	IPF, PF-ILD, PASC-PF#	>50% grant-fu	unded <sup>‡</sup>			
AstraZeneca Programs**	n.d.	n.d.					AstraZeneca 🕏
PRS-400	n.d.	n.d.					
Genentech (GENE1)	n.d.	n.d.					Genentech A Member of the Roche Group

<sup>#</sup>IPF - Idiopathic Pulmonary Fibrosis, PF-ILD - Progressive Fibrosing Interstitial Lung Diseases, PASC-PF - Post-Acute sequelae of SARS-CoV-2 infection (PASC) Pulmonary Fibrosis (PF)

<sup>\*\*</sup>Pieris has U.S. co-development options for two of three additional programs partnered with AstraZeneca



<sup>‡~\$17</sup> million grant from the Bavarian government to evaluate PRS-220 in PASC-PF covers more than half of early-stage and phase 1 development costs of PRS-220 Pieris has separate U.S. co-development and co-commercialization options on PRS-060/AZD1402

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





#### PRS-060/AZD1402 Phase 2a Study

Part 1 (Safety)

Part 1a: 1mg + 3 mg
Dose

☐ Part 1b: 10 mg Dose

Participant Population: Moderate asthmatics controlled on ICS/LABA

Primary Endpoint: Safety and tolerability compared to placebo from baseline until follow-up (approximately 56 days)

# of Participants: ~45 (randomized: 1:1:1 for part 1a; 2:1 for part 1b)

Part 2 (Efficacy)

□ Part 2a: 1mg + 3 mg Dose

☐ Part 2b: 10 mg Dose

Participant Population: Moderate uncontrolled asthmatics on ICS/LABA with blood EO count of ≥ 150 cells/µL and FeNO ≥ 25 ppb at screening\*

Primary Endpoint: Improvement of FEV1 at four weeks relative to placebo # of Participants: ~300 (randomized: 1:1:1 for part 2a; 2:1 for part 2b)

Parts 1b & 2a initiated 1Q 2022

Dry powder formulation, administered b.i.d. over four weeks on top of standard-of-care therapy (medium dose ICS with LABA)

Study is sponsored, conducted, and funded by AstraZeneca



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\*In addition to uncontrolled asthmatics with threshold EO count and FeNO profile, there are other enrollment criteria associated with part 2 not in part 1, including FEV1 range and a different ACQ score.

#### DPI Formulation of PRS-060/AZD1402 Passed Safety Review

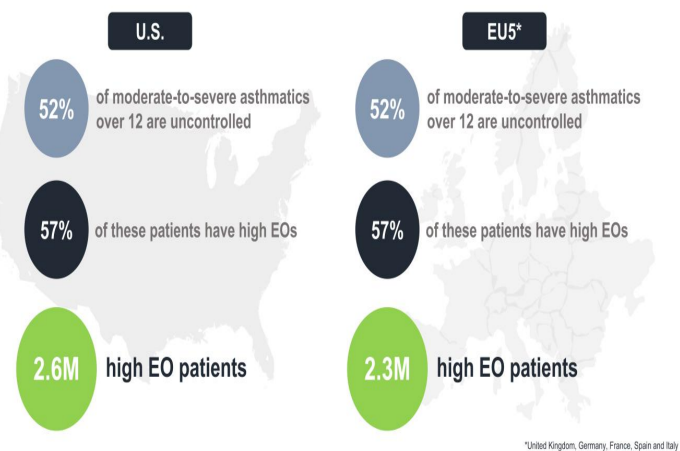
31 moderate asthmatics controlled on standard-of-care therapy (medium dose ICS with LABA) were dosed twice daily over four weeks randomized across two dose levels and placebo (1:1:1)

Safety review successfully completed for two dose levels (part 1a), triggering efficacy study (part 2a) 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



## Significant Market Opportunity in High EO Moderate-to-Severe Asthma





All numbers reflect 2022 estimates.

Sources: Artisan Healthcare Consulting analysis (2022), including the following: CDC, Journal of Asthma and Allergy,
The Journal of Allergy and Clinical Immunology, International Society of Pharmacoepidemiology

#### Co-Development Options for PRS-060/AZD1402

#### PIRS Opt-in Decision Point Three Possible Options No opt-in & no cost sharing \$1.9B in sales milestones **Phase 2a Primary Endpoint:** Improvement of FEV1 at 4 weeks relative to placebo 25% cost share with cost cap Single digit up to high-teen royalties for product lifetime Co-Dev Option Requirements: Development milestones approximating 50% of development costs 30-day opt-in period triggered upon both completion Potential \$3.5B+ in sales milestones of Phase 2a trial and notice by AZ (must include product development plan & budget) 50% cost share without cost cap Gross margin share percentage in mid-twenties for the product lifetime Development milestones approximating two-thirds of 25% option



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#### PRS-220: Inhaled CTGF Antagonist



<sup>\*</sup>IPF - Idiopathic Pulmonary Fibrosis

<sup>\*</sup>PASC-PF - Post-Acute sequelae of SARS-CoV-2 infection (PASC) Pulmonary Fibrosis (PF)

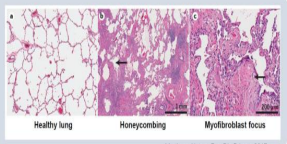


<sup>\*</sup>PF-ILD - Progressive Fibrosing Interstitial Lung Diseases

#### **IPF: High Unmet Medical Need and Significant Commercial Opportunity**

#### PF - a chronic lung disease:

ultimately fatal lung disease of unknown cause characterized by progressive scarring of the interstitial lung tissue



Martinez, Nature Rev Dis Primer, 2017

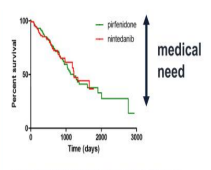
3 to 5 years

median survival from the time of diagnosis

Hopkins, European Respiratory Journal, 2016

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approved therapies nintedanib & pirfenidone providing modest benefit with significant side effects



Adapted from Cameli, Frontiers in Molecular Biosciences, 2020



Significant need for welltolerated and effective therapies



# Inhaled Delivery of PRS-220: A Novel Approach to Modulate CTGF Biology with Best-in-Class Potential

Key points of differentiation of inhaled PRS-220 compared to systemically delivered CTGF antagonists

# More Efficient Target Saturation

- Avoidance of systemic CTGF sink (in blood)
- Significantly higher affinity with superior binding profile

### Superior Lung Biodistribution

- Local delivery to the site of the disease in the lung via inhalation
- Increased concentration

# Increased Convenience

 Inhalation at home compared to regular visits to infusion centers for i.v. administrations



#### **Immuno-Oncology Pipeline**

Program	Target	Indication	Discovery	Preclinical	Phase 1	Phase 2	Partner
Cinrebafusp Alfa	a 4-1BB/HER2	HER2-High GC*			1000		
(PRS-343)		HER2-Low GC**			1000		
PRS-344/ S095012	4-1BB/PD-L1	n.d.	~50% co-de	ev cost share			* SERVIER
PRS-352/ S095025	OX40/PD-L1	n.d.					* SERVIER
Seagen Programs <sup>‡</sup>	Co-stim Agonist	n.d.					<b>O</b> Seagen
PRS-342/ BOS-342	4-1BB/GPC3	n.d.					BOSTON pharmaceuticals

<sup>‡3</sup> bispecific programs in collaboration with Seagen, with Pieris retaining a US co-promotion option for one of the three programs

<sup>\*\*</sup>Phase 2 study includes Cinrebafusp Alfa in combination with tucatinib (HER2-low arm)



<sup>\*</sup> Phase 2 study includes Cinrebafusp Alfa in combination with ramucirumab and paclitaxel (HER2-high arm)

#### 4-1BB & the Advantages of Anticalin-based Bispecifics

#### High-value target

- 4-1BB activation can drive massive proliferation and improved cytotoxic profile of tumor-specific T cells
- 4-1BB activation significantly increases mitochondrial load, improving metabolic fitness and overall survival of T cells

# Historical challenges of systemic mAbs

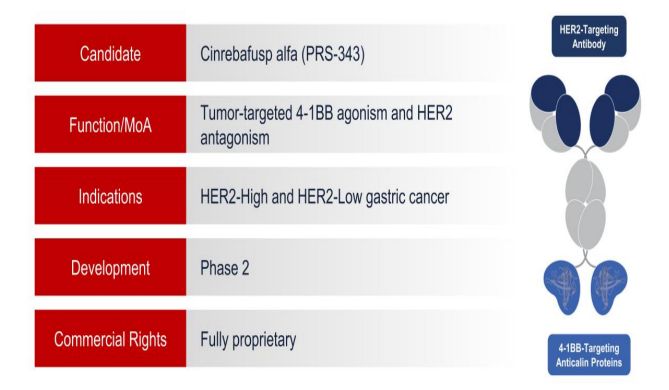
 Despite showing clinical activity, systemically active mAbs caused unmanageable hepatic toxicity and were discontinued

# Local activation solution

- Pieris' bispecifics are designed to efficiently activate 4-1BB on T cells and NK cells outside the liver, avoiding hepatic toxicity and driving improved therapeutic window
- Lead program validates this mode of action: well-tolerated and single-agent activity in heavily pre-treated patients



#### Cinrebafusp Alfa (PRS-343): Lead IO Asset





#### Cinrebafusp Alfa: 4-1BB/HER2 Bispecific

Cinrebafusp alfa drives 4-1BB agonism in the tumor microenvironment of HER2+ solid tumors

HER2-targeting moiety of the drug 4-1BB cross-linking ameliorates **BIOMARKERS** T-cell exhaustion and is critical localizes to the tumor microenvironment and facilitates 4-1BB cross-linking for T-cell expansion 4-1BB **Pathway** cinrebafusp alfa **Activation** HER2 Soluble 4-1BB targeting HER2 Antibody cinrebafusp alfa T-cell 4-1BB Co-Stimulation **Proliferation** targeting Anticalin® CD8+ and -1BB **Proteins** CD8+/Ki67+

**CLINICALLY-RELEVANT** 







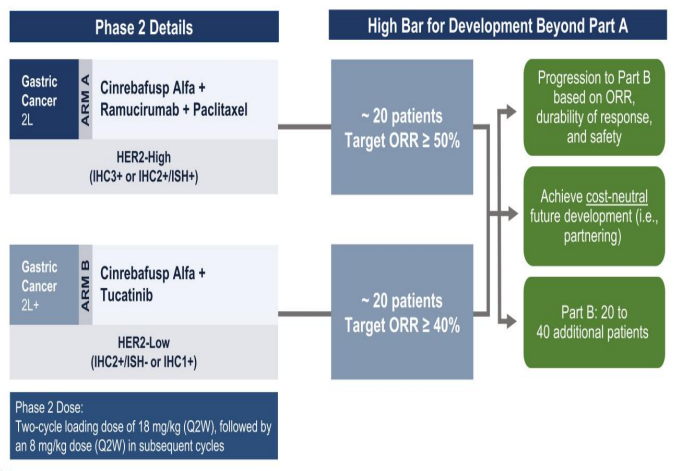
# Cinrebafusp Alfa Achieved Clinical POC in Phase 1 Monotherapy Study

- Acceptable safety profile observed at all doses tested with no doselimiting toxicities
- Clinical benefit at active dose levels (≥ 2.5 mg/kg), including confirmed complete response and several confirmed partial responses
- Dose-dependent immune activation and 4-1BB modulation in both HER2-high and HER2-low expressing patients
- Durable anti-tumor activity in heavily pre-treated patient population (5+ line on average), including "cold" tumors

As lead IO program, cinrebafusp alfa provides key validation of 4-1BB franchise, including PRS-344/S095012 and PRS-342/BOS-342



#### **Cinrebafusp Alfa Clinical Development Plan**



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

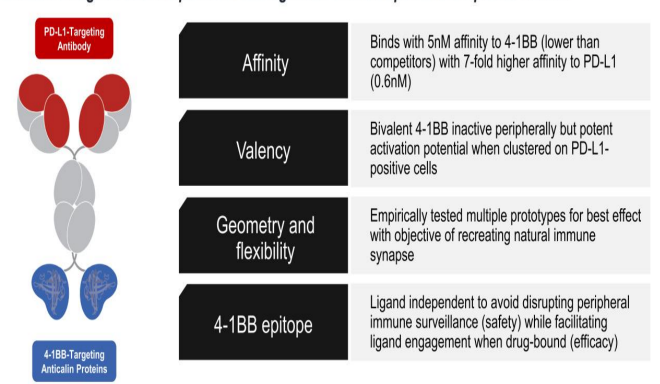




#### PRS-344/S095012: Why 4-1BB/PD-L1

PRS-344/S095012 is designed to activate 4-1BB on tumor-specific T cells when bridging to PD-L1-expressing tumors and dendritic cells

Molecule designed to drive potent 4-1BB agonism with an optimal therapeutic window





#### Financial Overview (as of 3/31/22)





255 State Street Boston, MA 02109 USA Zeppelinstraße 3 85399 Hallbergmoos Germany



Nasdaq: PIRS

