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# Preclinical Characterization of ATYR1923 (iMod.Fc), an Immune-Modulatory Therapeutic With Potentially Broad Application in Interstitial Lung Diseases

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INTRODUCTION: During the evolution of complex organisms, aminoacyl-tRNA synthetase genes evolved to incorporate new sequences and generate multiple splice variants, which lose their tRNA synthetase activity and take on novel functions (Lo et al. Science. 2014;345(6194):328-32). Histidyl-tRNA synthetase (HARS) and its splice variants are secreted and exhibit extracellular activity, which we have termed the Resokine pathway. Based on the overexpression in the lung of a splice variant encoding the N-terminal domain of Resokine, we hypothesized that it modulates the activity of immune cells in interstitial lung diseases (ILDs) and consequently ameliorates disease

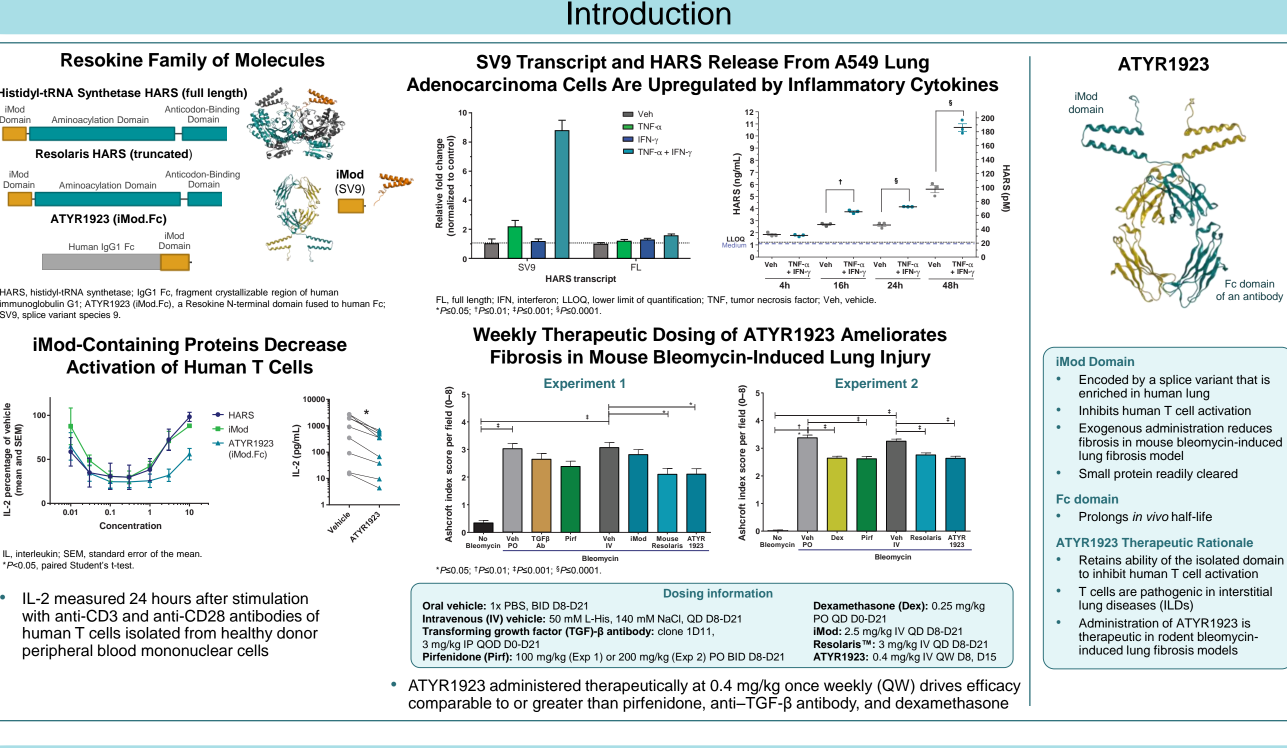
RATIONALE: In previous work, we showed that administration of Resokine proteins containing the N-terminal immunomodulatory (iMod) domain reduced bleomycin-induced lung fibrosis in mice, demonstrating the functional significance of the Resokine pathway in the lung. Based on these observations, we sought to engineer and characterize a clinical candidate with appropriate pharmaceutical properties for clinical study in ILD. Specifically, we sought to extend the duration of action of the iMod by fusion to the fragment crystallizable region (Fc) of human immunoglobulin G1 (IgG1 Fc).

METHODS: ATYR1923 (iMod.Fc), a Resokine N-terminal domain fused to human Fc, was expressed in Escherichia coli and purified to homogeneity, confirming low endotoxin (limulus amebocyte lysate [LAL] assay)

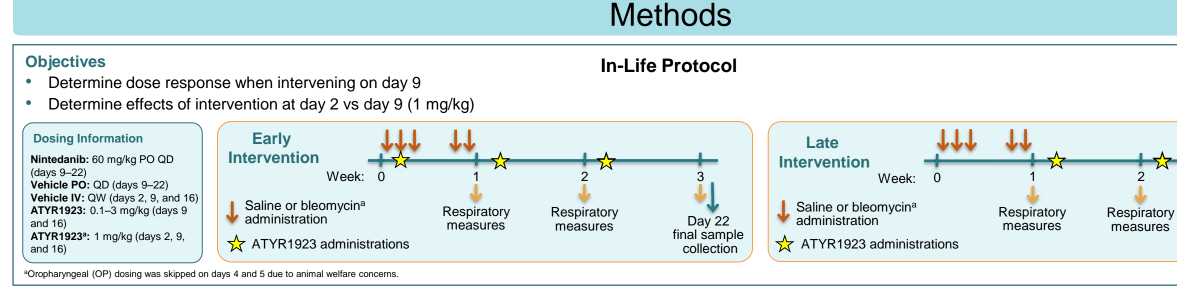
and pathogen-associated molecular pattern signals by a novel cell-based method. A rat model of bleomycin-induced lung fibrosis was employed to explore the effects ATYR1923 in vivo, including whole body plethysmography and histological disease scoring on day 22. Pharmacokinetic studies and Good Laboratory Practice (GLP)-compliant 1- and 3-month toxicology studies were conducted in rats and nonhuman primates (NHPs)

RESULTS: ATYR1923 exhibited the therapeutic potential of the iMod domain while having a long in vivo half-life ATYR1923 had a terminal half-life of ~3 days in rats and ~4.5 days in NHPs, in contrast to the isolated iMod domain that had a terminal half-life of ~20 minutes in rats. In rat bleomycin-induced lung fibrosis, ATYR1923 at 0.1-3 mg/kg weekly beginning on day 9 exerted therapeutic activity as revealed by reversal of bleomycin-induced changes in respiratory parameters and decreased histological fibrosis (Ashcroft score) and immune infiltration. One- and 3-month GLP-compliant studies found no adverse test article-related findings. The no-observed-adverse-effect level was 60 mg/kg in both species.

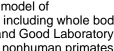
CONCLUSIONS: ATYR1923 has been engineered to have a long duration of action and is efficacious in bleomycin-induced lung fibrosis preclinical models when administered weekly. Based on the preclinical data, clinical testing is planned.

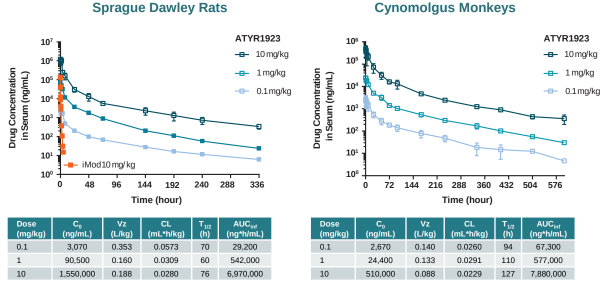


Abstract



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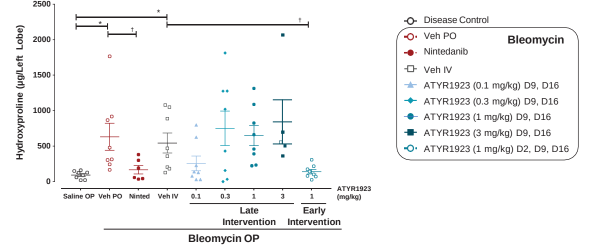




Fusion of iMod to Human Fc Sustains Exposure

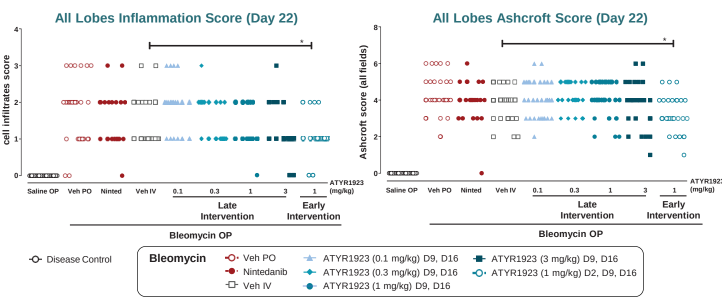
AUC<sub>10</sub>, area under the curve extrapolated to infinity: C<sub>0</sub>, initial plasma drug concentration: CL, total clearance: T<sub>10</sub>, half-life Vz. apparent volume of distribution during terminal phase.

# Collagen Content Decreased by ATYR1923 and Nintedanib



\*P<0.05: Kruskal-Wallis analysis of variance (ANOVA) followed by Dunn's multiple comparisons test of intended comparisons <sup>†</sup>P<0.05: Mann-Whitney U test

# **ATYR1923 Reduces Histological Inflammation and Fibrosis**



BLM +

Late

ATYR1923

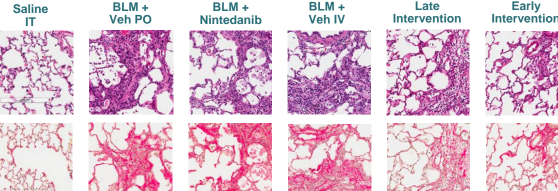
BLM +

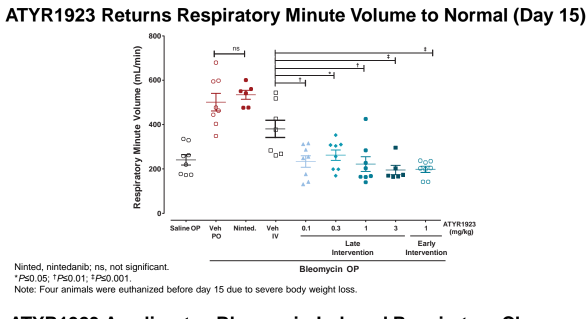
**ATYR1923** 

\*P<0.05, 2-way repeated measures ANOVA followed by Dunnett's multiple comparisons test

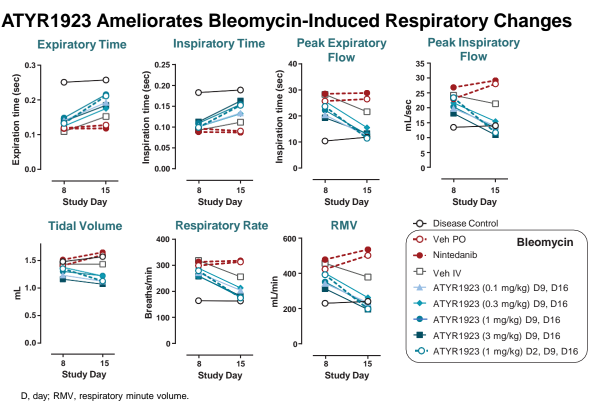
# Day 22 final sample collection

## Representative Images BLM + BLM + BLM + Veh PO Nintedanib Veh IV





# \**P*≤0.05; †*P*≤0.01; ‡*P*≤0.001



# Day 8 Respiratory Measures

				•					
		Bleomycin OP							
	Disease				ATYR1923 (mg/kg)/Treatment Days				
	Control (Saline OP)	Vehicle PO	Nintedanib	Vehicle IV	0.1 Days 9, 16	0.3 Days 9, 16	1 Days 9, 16	3 Days 9, 16	1 Days 2, 9, 16
Tidal volume	1.5	1.4	1.5	1.4	1.2	1.4	1.3	1.2	1.3
(mL)	(0.06)	(0.05)	(0.05)	(0.05)	(0.05)	(0.09)	(0.11)	(0.07)	(0.11)
Respiratory	164.4 <sup>‡</sup>	298.9	313.0	318.2	270.8	288.6	259.5	257.9	279.3
rate (bpm)	(20.2)	(15.9)	(17.8)	(16.6)	(17.4)	(18.2)	(21.0)	(18.9)	(13.5)
Respiratory minute volume (mL/min)	230.5 <sup>‡</sup> (28.0)	423.0 (33.4)	477.5 (36.3)	456.3 (28.7)	342.4 (33.5)	400 (39.3)	351.2 (55.5)	312.6* (45.9)	391.9 (44.3)
Expiratory time (s)	0.25 <sup>‡</sup>	0.12	0.12	0.11	0.14	0.12	0.15	0.13	0.13
	(0.017)	(0.008)	(0.008)	(0.008)	(0.012)	(0.011)	(0.014)	(0.014)	(0.007)
Inspiratory	0.18 <sup>‡</sup>	0.10	0.09	0.09	0.10	0.10	0.11	0.11	0.10
time (s)	(0.016)	(0.004)	(0.005)	(0.005)	(0.005)	(0.007)	(0.008)	(0.006)	(0.005)
Peak expiratory	10.3 <sup>‡</sup>	25.6	28.4	28.2	20.3	25.6	22.2	20.5	23.6
flow (mL/s)	(1.5)	(2.5)	(1.9)	(1.8)	(2.1)	(3.5)	(4.0)	(2.7)	(3.1)
Peak inspiratory flow (mL/s)	13.4‡	22.9	26.7	24.1	19.6	21.6	203	18.1	23.3
	(1.5)	(1.1)	(2.1)	(1.4)	(1.8)	(1.8)	(3.2)	(2.2)	(2.7)

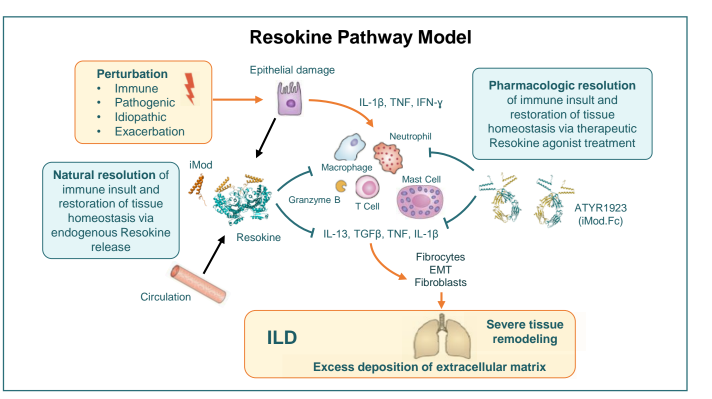
# **Day 15 Respiratory Measures**

			-						
		Bleomycin OP							
	Disease				ATYR1923 (mg/kg)/Treatment Days				
	Control (Saline OP)	Vehicle PO	Nintedanib	Vehicle IV	0.1 Days 9, 16	0.3 Days 9, 16	1 Days 9, 16	3 Days 9, 16	1 Days 2, 9, 16
Tidal volume	1.6	1.6	1.6	1.4	1.1*	1.2	1.2	1.1 <sup>†</sup>	1.1*
(mL)	(0.06)	(0.04)	(0.04)	(0.05)	(0.06)	(0.05)	(0.08)	(0.08)	(0.04)
Respiratory	163.1 <sup>‡</sup>	312.5	317.3	255.5	204.2	213.5	178 <sup>†</sup>	183.1*	175.1 <sup>†</sup>
rate (bpm)	(17.1)	(20.0)	(10.5)	(20.0)	(14.8)	(15.5)	(15.1)	(14.5)	(8.5)
Respiratory minute volume (mL/min)	241.1* (23.6)	501.2 (39.6)	534.6* (20.1)	378.4 (39.0)	233.5* (26.2)	262.2 (23.5)	222.1 <sup>†</sup> (33.3)	195 <sup>†</sup> (20.9)	198 <sup>†</sup> (13.3)
Expiratory time (s)	0.26 <sup>‡</sup>	0.13	0.12	0.16	0.19	0.18	0.22 <sup>†</sup>	0.18	0.21 <sup>†</sup>
	(0.016)	(0.013)	(0.006)	(0.012)	(0.015)	(0.010)	(0.021)	(0.013)	(0.010)
Inspiratory	0.19 <sup>‡</sup>	0.09	0.09	0.11	0.13	0.13	0.15 <sup>†</sup>	0.16 <sup>†</sup>	0.15*
time (s)	(0.016)	(0.004)	(0.002)	(0.010)	(0.010)	(0.009)	(0.011)	(0.014)	(0.008)
Peak expiratory	11.8*	26.4	28.8	21.6	12.3*	15.5	12.8*	13.3	11.4*
flow (mL/s)	(1.1)	(2.8)	(1.2)	(2.1)	(1.1)	(1.5)	(2.0)	(2.3)	(0.5)
Peak inspiratory flow (mL/s)	14.0*	27.9	29.0*	21.3	13.8*	15.4	13.1*	10.8 <sup>†</sup>	11.7 <sup>†</sup>
	(1.5)	(1.8)	(1.0)	(2.1)	(1.4)	(1.4)	(2.0)	(1.6)	(0.7)

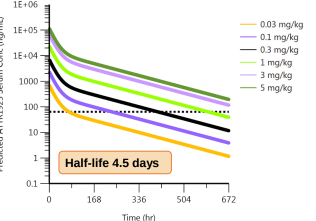
\*P≤0.05; †P≤0.01; ‡P≤0.001 Each respiratory endpoint was subject to 2-way repeated measures ANOVA followed by Dunnett's multiple comparisons test vs vehicle IV

# Exhibit 99.1

# Results



Pharmacokinetic predictions potentially enable once-monthly dosing in patients



# Single-Dose Simulations: 0.3–3 mg/kg

ATYR1923 Single Ascending IV Infusion Dose Level (mg/kg)	Duration of Time Plasma Conc is >1 nM (64 ng/mL) <sup>a</sup> [Half-life 4.5 days]	Duration of Time Plasma Conc is >1 nM (64 ng/mL) <sup>a</sup> [Half-life 7 days]
0.03	3.1 days	4.8 days
0.1	9.8 days	15.1 days
0.3	16.9 days	26.2 days
1.0	24.8 days	38.3 days
3.0	>29.2 days	49.4 days
10.0	>33.3 days	>58 days

<sup>a</sup>1 nM is assumed to be the concentration above which therapeutic benefit is expected

# Favorable Good Laboratory Practice (GLP) Safety Profile

## Nonhuman Primate

- 2 weekly IV doses of 3 mg/kg
- No increase in ~30 serum immune markers
- 1- and 3-month weekly IV dose at 0, 10, 30, and 60 mg/kg
- No adverse test article-related findings
- Systemic exposure increased with increasing dose and did
- not appear to change with repeated dosing
- Anti-drug antibodies (ADA) did not appear to have an impact on systemic exposure
- No-observed-adverse-effect level (NOAEL) = 60 mg/kg  $(C_{trough} = 228 \text{ nM})$

- 1- and 3-month weekly IV dose
- at 0, 10, 30, and 60 mg/kg No adverse test article-related finding
- Systemic exposure increased with
- increasing dose and did not appear to change with repeated dosing
- ADA did not appear to have an impact on systemic exposure
- NOAEL = 60 mg/kg ( $C_{trough}$  = 75 nM)

# Summary

- ATYR1923 had a terminal half-life of ~3 days in rats and ~4.5 days in NHPs
- ATYR1923 at 0.1–3 mg/kg weekly beginning on day 9
- Reversed bleomycin-induced changes in respiratory parameters
- ATYR1923 at 1 mg/kg weekly beginning on day 2
- Decreased histological fibrosis (Ashcroft score) Decreased histological inflammation
- Reduced lung collagen content
- 1- and 3-month GLP-compliant studies found no adverse test article-related findings. The NOAEL was 60 mg/kg in both species.

# Conclusions

 ATYR1923 has been engineered to have a long duration of action and is efficacious in bleomycin-induced lung fibrosis preclinical models when administered weekly. Based on the preclinical data, a phase 1 clinical study is under way.

## Acknowledgments

The authors thank the scientists at Charles River Laboratories (bleomycin-induced lung fibrosis experiment), Edinburgh, UK; Shin Nippon Biomedical Laboratories (cynomolgus monkey PK study), Seattle, WA, USA; MPI Research (toxicology), Mattawan, MI, USA; and SD Scientific, Inc. (PK predictions), San Diego, CA, USA, for the data that they contributed