# Corticosteroid Tapering and Discontinuation in a Phase 2 Study of Rilonacept in Recurrent Pericarditis

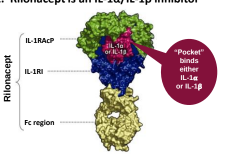
Allan Klein¹, S. Allen Luis², Martin M. LeWinter³, David Lin⁴, Paul Cremer¹, Saifullah Nasir⁵, Antonio Abbate⁵, Andrew Ertel७, Fang Fang®, Anna Beutler9, John F. Paolini® ¹Cleveland Clinic, Cleveland, Ohio, USA; ²Mayo Clinic, Rochester, Minnesota, USA; ⁵Stat! Cardiology, Chicago, Illinois, USA; ⁵Nirginia Commonwealth University, Richmond, Virginia, USA; ⁵Minneapolis, Minneapolis, Minneapolis, Minnesota, USA; ⁵Nirginia Commonwealth University, Richmond, Virginia, USA; ⁵Nirginia Commonwealth University, Richmond, VIII (VIII) (VIII

#### BACKGROUND

- Recurrent pericarditis (RP) is characterized by the recurrence of pericarditis signs and symptoms after a symptom-free period of ≥4 to 6 weeks<sup>1</sup> and affects 15% to 30% of patients after a first episode<sup>2,3</sup>
- Conventional treatment options include nonsteroidal antiinflammatory drugs (NSAIDs), colchicine, and corticosteroids (CS)<sup>4</sup>
- ESC guidelines<sup>1</sup> recommend NSAIDs and colchicine as first-line treatment, with CS added to NSAIDs and colchicine as escalation to triple therapy in case of no or incomplete response
- CS are often used long-term in patients (pts) with RP
- CS are associated with substantial comorbidities<sup>5</sup>
  - Cushingoid appearance, weight gain, increased risk for infection, skin fragility, corticosteroid induced diabetes, risk of adrenal insufficiency upon withdrawal, muscle weakness, elevated blood pressure, mood instability, and compression fractures due to osteoporosis<sup>7</sup>
  - Some CS-related morbidities may be irreversible or require a surgical intervention, as in the case of avascular necrosis or cataracts
- Treatment with CS with fast tapering may be associated with an increased risk of pericarditis recurrences8
- Interleukin-1 (IL-1) is a family of cytokines which mediate the pathophysiology of recurrent pericarditis (Figure 1)
- Tissue damage caused by inflammation as a result of IL-1 $\alpha$  and IL-1 $\beta$  in the pericardium stimulates additional IL-1 $\alpha$  and IL-1 $\beta$ , thereby creating a self-perpetuating cycle of pericardial inflammation
- Rilonacept inhibits IL-1 signaling by acting as a soluble decoy receptor that binds IL-1 $\alpha$  and IL-1 $\beta$ , thus preventing their interaction with IL-1 cell surface receptors (Figure 2)

#### Figure 1. Role of IL-1 $\alpha$ and IL-1 $\beta$ in the Autoinflammatory Cycle of Recurrent Pericarditis9, 10

Figure 2. Rilonacept is an IL- $1\alpha$ /IL- $1\beta$  inhibitor



#### **METHODS**

# Phase 2 Study

**Study Objective** 

- Evaluate the efficacy and safety of rilonacept in patients with RP,
- Improvement of pericarditis symptomatology with rilonacept Feasibility of weaning from CS while receiving rilonacept in patients
- with CS-dependent RF - Disease activity after CS taper in patients with CS-dependent RP
- Safety of rilonacept Study Design
- Open-label, single-active-arm, 5-part pilot study
- Eligible patients were adults (18 to 75 y) or children (≥6 to <18 y) with RP due to idiopathic or PPS etiology, presenting with at least a third pericarditis episode or with at least 3 prior episodes if not in an active episode but CS-dependent at the time of enrollment
- All patients at study entry were allowed concomitant NSAIDs and/or colchicine and/or CS (in any combination) as long as the dosages were stable for ≥7 days; CS-dependent patients must have been on CS at enrollment

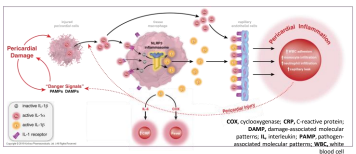
### Treatment and Procedures

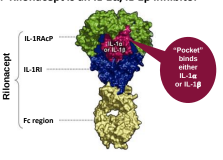
- In the 6-week open-label base treatment period (TP):
- Adults (≥18 y) received a loading dose of 320 mg (2 × 160 mg) rilonacept, administered via SC injection on day 0, followed by 160 m SC weekly for 5 additional doses
- No changes in concomitant medications were allowed during the TP unless medically indicated
- In the optional 18-week treatment extension period (EP), during which weekly rilonacept continued with the same schedule, investigators were given the option to wean patients from concomitant medications, including CS
- Patients who completed both the TP and the EP received rilonacent for 6 months
- Primary endpoints: Patients with active pericarditis: pain numeric rating scale (NRS) and C-reactive protein (CRP) levels at baseline and on treatment; patients with CS-dependent non-active pericarditis: disease activity during and after CS taper
- Secondary endpoints: Use of concomitant CS; changes in the use of other concomitant medications for pericarditis Safety Assessments
- Adverse events (AEs) were recorded with the level of severity and relationship to study drug based on investigators' judgement

# Patients on Corticosteroids or Colchicine at Baseline<sup>a</sup>

- In this retrospective analysis, patients were divided into groups based on their use of therapies at baseline
- CS-failure group: patients with active pericarditis despite ongoing treatment with CS at enrollment; prior history of at least 2 pericarditis
- CS-dependent group: patients with no active episode of pericarditis but defined by Investigator as dependent on CS to control their disease; history of at least 3 pericarditis episodes
- Colchicine-failure group: patients with active pericarditis despite ongoing treatment with colchicine enrolled in the study before escalating to CS therapy
- Analyses of CS tapering dose and duration excluded patients who did not enroll in the 18-week EP; other analyses (baseline demographics, efficacy, etc) included these patients

r continuous variables (e.g., change from baseline), summary statistics were calculated as mean and median; for categorical variables, frequency and percentage were calculated bPatients could be on any other standard of care medication in addition to CS to be included in this group, including Patients could be on any other standard of care medication, with the exception of CS, to be included in this grou





# **RESULTS**

**Baseline Patient Demographics and Characteristics** 

- Out of 25 patients enrolled in the study, 2 discontinued before entering the EP
  - 1 patient with active pericarditis discontinued during the TP (SAE of subcutaneous abscess; resolved with standard management)
  - 1 CS-dependent patient declined to enter the EP
- Out of 23 patients who completed the 24 weeks of study, 57% (n=13) of patients were receiving CS at baseline
  - CS-dependent: 35% (n=8); all except one patient were also receiving
  - CS-failures (active pericarditis): 22% (n=5); all patients were also receiving colchicine
- Out of 23 patients who completed the 24 weeks of study, 30% (n=7) were colchicine-failures (active pericarditis, no CS)

#### Table 1. Baseline Demographics

	General Characteristics	All Patients	CS- failure	dependent w/o active pericarditis	Colchicine -failure
	Unique patients, n	25	6	9	7
ng	Mean age (range), yrs	42.8 (26-62)	32.3 (26-42)	48.2 (36-62)	42.6 (26-58)
	Sex (male/female)	10/15	0/6	6/3	2/5
	Race (white/African American)	22/3	4/2	9/0	7/0
с	Mean # previous pericarditis recurrences <sup>a</sup> (range)	2.6 (1-8)	1.8 (1-3)	3.3 (2-5)	2.7 (1-8)
	Mean disease duration (range), yrs	2.2 (0.2-7.9)	2.9 (1.0-5.6)	1.4 (0.6-3.4)	2.9 (0.2-7.9)

### **Table 2. Clinical Characteristics**

	CS-failure CS-dependent w/o active n = 6 pericarditis n = 9		Colchicine- failure n = 7
Mean NRS <sup>a</sup> (SD)	5.7 (2.3)	1.4 (1.5)	4.3 (1.3)
Mean CRP (SD), mg/dL	4.6 (7.6)	0.2 (0.1)	4.2 (4.2)

<sup>a</sup>11-point numeric scale, ranging from zero (0, no pain) to ten (10, pain as bad as possible); CRP, C-reactive protein; NRS, numer

# **Corticosteroid Dose at Baseline in All Patients**

- In CS-failure patients with active pericarditis, mean baseline dose of prednisone was 20.6 mg per day
- In CS-dependent patients without an active pericarditis episode, mean baseline dose of prednisone was lower, at 8.5 mg per day
- The only CS used in the study for pericarditis treatment was

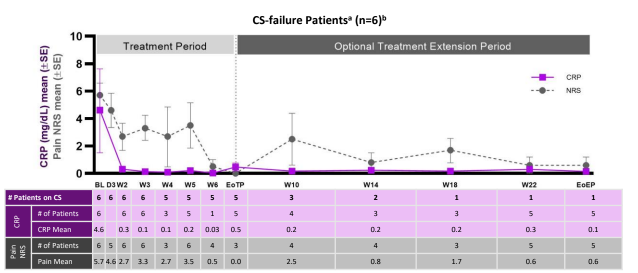
## Table 3. Baseline CS Dose in All Patients<sup>a</sup>

	C3-lallule	active pericarditis	
	n = 6	n = 9	
Prednisone dose (mg/day)			
Mean (SD)	20.6 (19.6)	8.5 (8.9)	
Median (range)	11.3 (1.0-50.0)	5.0 (2.5-30.0)	

#### RESULTS, continued

All CS-failure patients who completed the 24-week study experienced resolution of the acute pericarditis episode and were able to taper or discontinue CS without pericarditis recurrence while on treatment with rilonacept

Figure 3. All CS-failure patients experienced rapid, sustained, and clinically meaningful reductions in pericarditis pain and CRP with rilonacept



ents with active pericarditis episodes who were on CS at enrollment (safety population); results were similar in patients who failed CS and entered the EP (EP population; n=5) bOne CS-failure patient discontinued during the TP due to an SAE; EoTP, end of treatment period; EoEP, end of extension period

Table 4. CS dose and duration prior to enrollment in CS-failure patients who completed the EP (n=5)a

	Time on CS prior to enrollment (days)	CS dose at enrollment (mg/day)	CS dose at end of study (mg/day)
Patient 1	29	10	0
Patient 2	>320	1	0
Patient 3	19	12.5	0
Patient 4	9	30-40 <sup>b</sup>	0
Patient 5	13	50	30
30no CC failura natio	ant discontinued during the TD due t	o on CAE	

Figure 4. Median CS dose in CS-failure patients who completed the EP (n=5)a

Week of Stud

- Of 5 CS-failure patients who completed the 24 weeks of study
  - 4 patients discontinued CS during the EP

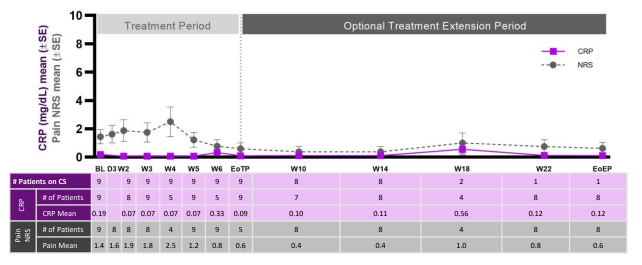
Patient was on 40 mg/day from day -9 to day -4, then on 30 mg/day until enrollmen

- 1 patient reduced CS dose from 50 mg/day at baseline to 30 mg/day at final visit There were no pericarditis recurrences during CS taper/discontinuation while on rilonacept treatment

All CS-dependent patients without active pericarditis who completed the study tapered and/or discontinued CS use with no pericarditis recurrence while on treatment with rilonacept

Figure 5. All CS-dependent patients without active pericarditis tapered and/or discontinued CS use with no pericarditis recurrence while on treatment with rilonacept

### CS-dependent Patients<sup>a</sup> (n=9)<sup>b</sup>

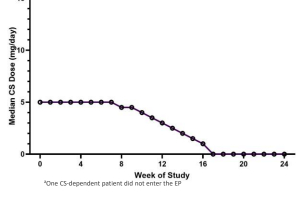


PALI CS-dependent patients without active pericarditis (safety population); results were similar in CS-dependent patients who entered the EP (EP population; n=8)
POne CS-dependent patient did not enter the EP; EoTP, end of treatment period; EoEP, end of extension period;

#### Table 5. CS dose and duration prior to enrollment in CS-dependent patients who completed the EP (n=8)<sup>a</sup> Time on CS prior CS dose at CS dose at end of

		to enrollment (days)	enrollment (mg/day)	study (mg/day)
	Patient 1	35	5	0
	Patient 2	17	2.5	0
	Patient 3	51	30	2.5
	Patient 4	33	8	0
	Patient 5	23	4	0
	Patient 6	63	3	0
	Patient 7	259	15	0
	Patient 8	142	5	0
	<sup>a</sup> One CS-dependent patient did not enter the EP			

Figure 6. Median CS dose during the study in CSdependent patients who completed the EP (n=8)<sup>a</sup>

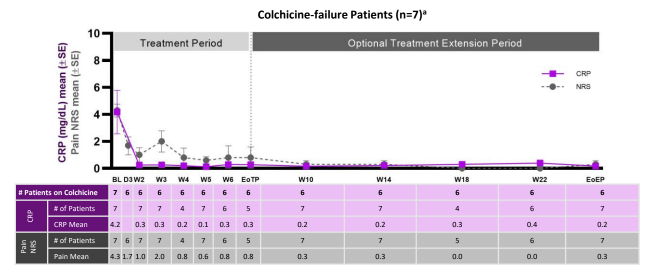


• 7 out of the 8 CS-dependent patients who completed the EP (87.5%) successfully stopped CS with no pericarditis recurrence while on

1 remaining patient successfully tapered from 30 mg/day at baseline to 2.5 mg/day by completion of the EP with no pericarditis recurrence while on treatment with rilonacept

#### All colchicine-failure patients had rapid, sustained, and clinically meaningful reductions in pericarditis pain and CRP with initiation of rilonacept

Figure 7. All colchicine-failure patients had rapid, sustained, and clinically meaningful reductions in pericarditis pain and CRP with initiation of rilonacept



EoTP, end of treatment period; EoEP, end of extension period

- Of the 7 colchicine-failure patients (active pericarditis episode despite colchicine and enrolled in lieu of CS initiation) who completed the EP:
- 6 out of 7 patients experienced successful treatment of the acute episode and no pericarditis recurrence while on treatment with rilonacept - 1 patient experienced a mild recurrence during the TP, 5 days duration, with NRS pain increase from 0 to 2 and CRP of 0.10 mg/dL, not requiring addition of
- new medication to treat pericarditis; patient stayed on rilonacept treatment until the end of the study with no other recurrence - 1 discontinued colchicine during the study, while the remaining 6 patients did not decrease colchicine dose. This investigator decision to not discontinue
- concomitant colchicine may be related to anticipation of a potential post-study recurrence in patients with long-standing RP once fixed-duration rilonacept treatment had been completed

#### Annualized incidence of pericarditis episodes decreased across all groups of patients during rilonacept treatment

Table 6. Annualized Incidence of Pericarditis Episodes Prior to and During the Study

	CS-failure	CS-dependent w/o active	Colchicine-failure
	n = 6	pericarditis n = 9	n = 7
Prior to the study <sup>a</sup>			
Pericarditis episodes per year, mean (SD)	1.9 (1.3)	4.2 (2.6)	5.7 (5.9)
During the study <sup>b</sup>			
Patients with pericarditis episodes, n	0	0	<b>1</b> <sup>c</sup>
Pericarditis episodes per year, mean (SD)	0	0	0.3 (0.8)

Episodes during the study include recurrences during TP and EP combined. Pericarditis recurrence during the study was based on Investigator's judgement;
Patient had a mild pericarditis recurrence in TP, 5 days duration, with NRS pain increase from 0 to 2, CRP 0.10 mg/dL, not requiring addition of new medication to treat pericarditis

# Rilonacept was generally well-tolerated: majority of AEs were mild

- There were 2 serious treatment-emergent AEs in patients presenting with an active pericarditis episode, both of which resolved
  - 1 serious adverse event (SAE, subcutaneous abscess) in week 5 of TP in a patient with a history of skin infections, still receiving concomitant prednisone at the dose of 10 mg/day. The abscess resolved with standard management; patient discontinued rilonacept.
- 1 patient with atypical chest pain (not related to study drug) continued rilonacept treatment AEs observed with rilonacept treatment are consistent with the known safety profile of rilonacept
- The most common AEs were injection site reactions (12 patients out of 25 [48%]), nasopharyngitis, arthralgia, and diarrhea

### **CONCLUSIONS**

- CS-failure patients with active pericarditis treated with rilonacept experienced rapid, sustained, and clinically meaningful reductions in pericarditis pain and CRP and tapered or discontinued CS use without pericarditis recurrences while on
- CS-dependent patients tapered or discontinued CS without pericarditis recurrences while on rilonacept treatment
- Colchicine-failure patients with active pericarditis treated with rilonacept experienced rapid, sustained, and clinically meaningful reductions in pericarditis pain and CRP
  - 6 out of 7 patients did not taper off colchicine. This investigator decision to not discontinue concomitant colchicine may be related to anticipation of a potential post-study recurrence in patients with long-standing RP once fixed-duration rilonacept treatment had been completed
- Safety data from this study are consistent with the known safety profile of rilonacept
- These data suggest a potential corticosteroid-sparing effect of rilonacept, i.e., supporting a reduction in corticosteroid dose or obviating the need for corticosteroid use while on treatment in the study. Novel therapies are needed which could eliminate or reduce the risk of significant corticosteroid-associated morbidity in recurrent pericarditis.

Results from this study support the design of RHAPSODY, a double-blind, placebo-controlled randomized withdrawal (RW) pivotal Phase 3 tudy with an open-label extension, designed to evaluate the efficacy and safety of rilonacept treatment in patients with RP

nterpretation of efficacy and safety outcomes is limited by small number of patients in each study part, open-label study design, and single-active-treatment arm design

# References

1. Adler Y, et al. Eur Heart J. 2015;36:2921-2964. 2. Cremer P, et al. JACC. 2016;68(21): 2311-2328. 3. Imazio M, et al. Lancet. 2014;383(9936):2232-2237. 4. Lilly LS. Circulation. 2013;127:1723-1726. 5. Chaudhry HS, et al. In: StatPearls. Treasure Island (FL): StatPearls Publishing, LLC; 2019. 6. Strehl C, et al. Ann Rheum Dis. 2016;75(6):952-957. 7. Alraies MC, et al. Am J Cardiol. 2015;115:542-547. 8. Imazio M, et al. Circulation. 2010;121:916-928. 9. Brucato A, et al. Int Emerg Med. 2018;13(6):839-844. 10. Dinarello CA,

Arcalyst® (rilonacept, Regeneron, Tarrytown, NY) is approved in the US for treatment of Cryopyrin-

Rilonacept is being investigated for the treatment of RP by Kiniksa Pharmaceuticals, Ltd.

#### **Disclosures and Acknowledgements** This study was sponsored by Kiniksa Pharmaceuticals, Ltd.

AK - research grant, scientific advisory board Kiniksa Pharmaceuticals, Ltd., advisory board Swedish Orphan Biovitrum AB, advisory board Pfizer, Inc.and royalties from Kluwers Lippincott and Elsevier; SAL – scientific advisory board Kiniksa Pharmaceuticals, Ltd., advisory board Swedish Orphan Biovitrum AB, consultant for Swedish Orphan Biovitrum AB; MML - one seminar for Kiniksa Pharmaceuticals, Ltd.; PC - advisory board Swedish Orphan Biovitrum AB, advisory board Kiniksa Pharmaceuticals, Ltd. ; SN - Advisory board member for Kiniksa Pharmaceuticals, Ltd.; consultant and advisory board member for Swedish Orphan Biovitrum AB; AA research grant and advisory board for Kiniksa; AB - Research grants from Kiniksa Pharmaceuticals, Ltd., Swedish Orphan Biovitrum AB, Olatec Therapeutics LLC, Serpin Pharma, LLC; consultant fees: Kiniksa Pharmaceuticals, Ltd., Olatec Therapeutics LLC, Serpin Pharma, LLC, Merck & Co., Inc.; FF and JFP – employees of Kiniksa Corp.; AB – employee of Kiniksa Ltd.; DL and AE – no disclo