

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

October 13, 2023

Date of Report (Date of earliest event reported)

Aprea Therapeutics, Inc.

(Exact name of registrant as specified in its charter)

Delaware
(State or other jurisdiction
of incorporation)

001-39069
(Commission
File Number)

84-2246769
(IRS Employer
Identification No.)

3805 Old Easton Road
Doylestown, PA
(Address of principal executive offices)

18902
(Zip Code)

Registrant's telephone number, including area code: **(617) 463-9385**

(Former name or former address, if changed since last report): Not applicable

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- 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 registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common stock, par value \$0.001 per share	APRE	The Nasdaq Stock Market LLC

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

Emerging growth company

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.

Item 8.01 Other Events.

On October 13, 2023, Aprea Therapeutics, Inc. updated its corporate presentation slide deck. A copy of the corporate presentation slide deck is filed as Exhibit 99.1 hereto and incorporated herein by reference.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit Number	Description
99.1	Corporate Presentation (October 2023) .
104	Cover Page Interactive Data File (embedded within the inline XBRL document).

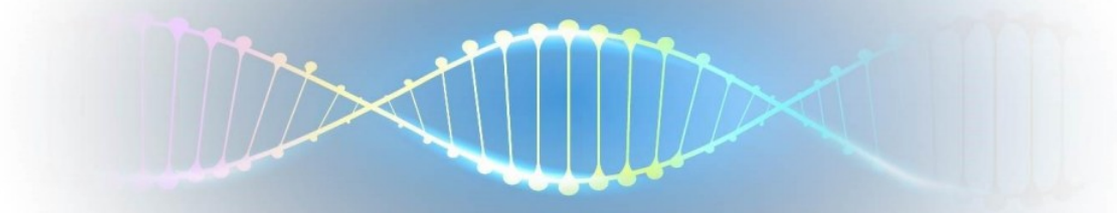
SIGNATURES

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.

Aprea Therapeutics, Inc.

Dated: October 13, 2023

By: /s/ Oren Gilad
Name: Oren Gilad, Ph.D.
Title: President and Chief Executive Officer



Precision Oncology Through Synthetic Lethality

October 2023

Certain information contained in this presentation includes “forward-looking statements”, within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended, related to our clinical trial regulatory submissions and strategic plans. We may, in some cases use terms such as “predicts,” “believes,” “potential,” “continue,” “anticipates,” “estimates,” “expects,” “plans,” “intends,” “may,” “could,” “might,” “likely,” “will,” “should” or other words that convey uncertainty of the future events or outcomes to identify these forward-looking statements. The forward-looking statements are based on current beliefs and expectations of our management team that involve risks, potential changes in circumstances, assumptions, uncertainties. Any or all of the forward-looking statements may turn out to be wrong or be affected by inaccurate assumptions our management team might make or by known or unknown risks and uncertainties. These forward-looking statements are subject to risks and uncertainties including, without limitation, risks related to the success and timing of our clinical trials or other studies and the other risks set forth in our filings with the U.S. Securities and Exchange Commission, including our Annual Reports on Form 10-K and Quarterly Reports on Form 10-Q. Forward-looking statements regarding our product candidates are also subject to additional risks and uncertainties, including without limitation, with respect to: our dependence on additional financing to fund our operations and complete the development and commercialization of our product candidates, and the risks that raising such additional capital may restrict our operations or require us to relinquish rights to our technologies or product candidates; our limited history and preclinical status of the assets we acquired from Atrin Pharmaceuticals Inc.; our business plan or the likelihood of the successful implementation of such business plan; the timing of initiation of planned clinical trials for our product candidates; the future success of such trials; the successful implementation of our research and development programs and collaborations and the interpretation of the results and findings of such programs and collaborations and whether such results are sufficient to support the future success of our product candidates; the success, timing and cost of our anticipated clinical trials for our current product candidates; the timing of initiation, utility analyses, data presentation, reporting and publication and receipt of interim results (including, without limitation, any preclinical results or data); any statements about our understanding of product candidates mechanisms of action and interpretation of preclinical and early clinical results from its clinical development programs and any collaboration studies; and other factors, including legislative, regulatory, political and economic developments not within our control. For all these reasons, actual results and developments could be materially different from those expressed in or implied by our forward-looking statements. You are cautioned not to place undue reliance on these forward-looking statements, which are made only as of the date of this presentation. We undertake no obligation to update such forward-looking statements to reflect subsequent events or circumstances, except to the extent required by law or regulation.

- Clinical stage precision medicine via novel synthetic lethality (SL) - based therapeutics
- All programs addressing significant unmet medical need

- ATR Inhibitor: ATRN-119
 - ◇ Clinical proof-of-concept
 - ◇ Phase 1/2a – Ongoing Dose Escalation
 - ◇ Patients 12 years of age or older with solid tumors harboring DDR mutation
 - ◇ Primary objective : Safety, MTD, RP2D and PK profile
 - ◇ Pre-clinical proof-of-principle
 - ◇ Demonstrated anti-tumor activity
 - ◇ Synergistic with anti-cancer therapies, including PARP inhibitors
 - ◇ Potential differentiation in safety and tolerability

- WEE1 Inhibitor: ATRN-1051
 - ◇ IND enabling studies
 - ◇ Anticipate submitting an IND by the end of 2023
 - ◇ Pre-clinical proof-of-principle
 - ◇ Demonstrated anti-tumor activity
 - ◇ Ovarian cancer with Cyclin E over expression
 - ◇ Synergistic with anti-cancer therapies, including ATR inhibitor
 - ◇ Potential differentiation in safety and tolerability

- DDR Inhibitor: Undisclosed
 - ◇ Lead optimization
 - ◇ Target identified from our RepliBiom discovery platform

Management

Oren Gilad, Ph.D. President and CEO	John Hamill CFO	Nadeem Mirza, M.D., MPH Senior Medical Advisor	Ze'ev Weiss, CPA, B.Sc. Chief Business Advisor	Mike Carleton, Ph.D. Translational Medicine Advisor

Board of Directors

Richard Peters, M.D., Ph.D. Chairman of the Board	Oren Gilad, Ph.D. President and CEO	Jean-Pierre Bizzari, M.D. Director
Marc Duey Director	Michael Grissinger Director	Gabriela Gruia, M.D. Director
John Henneman Lead Independent Director	Rifat Pamukcu, M.D. Director	Bernd R. Seizinger, M.D., Ph.D. Director



ATR Inhibitor : ATRN-119

ATR Inhibitor :
ATRN-119

Clinical
Proof-Of-Concept

AR-276-01: A PHASE 1/2a, OPEN-LABEL, SAFETY, PHARMACOKINETIC, AND PRELIMINARY EFFICACY STUDY OF ORAL ATRN-119 IN PATIENTS WITH ADVANCED SOLID TUMORS

Sites: 4 US sites for dose escalation

- ❖ University of Pennsylvania
- ❖ Mary Crowley Cancer Research
- ❖ University Hospitals Cleveland Medical Center
- ❖ Yale Cancer Center

Patient enrollment: 48 patients in total

- Escalation phase: up to 18 patients
- Expansion phase: up to 30 patients

IMP: ATRN-119 is an oral ATR kinase inhibitor given daily

Patient Population:

Male or female subjects 12 years of age or older with solid tumors harboring any DDR mutation per NGS

Part 1 (up to 18 patients)
Dose escalation (6 dose levels)
3+3 design



Part 2 (up to 30 patients)
Dose expansion, after MTD /
RP2D established

Primary objectives:

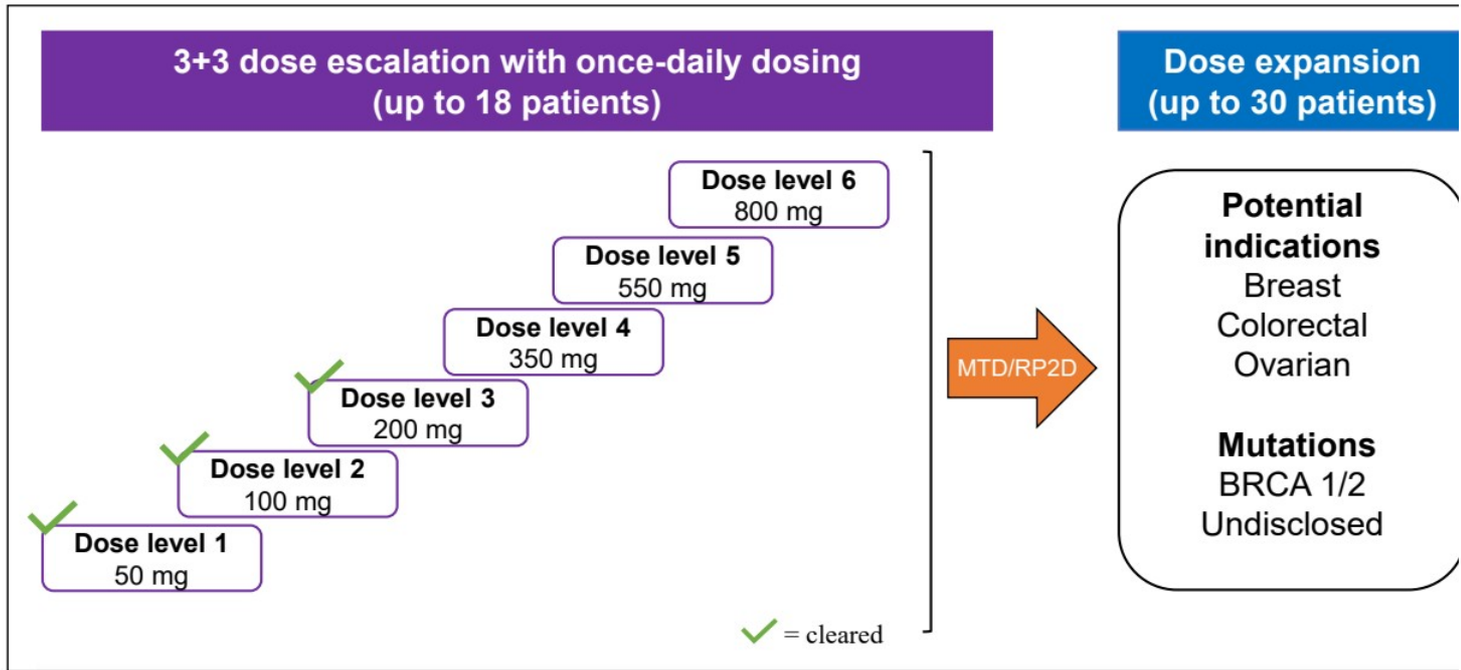
- Safety, MTD, RP2D
- Pharmacokinetics (PK profile of oral ATRN-119 and its active metabolite ATRN-157)

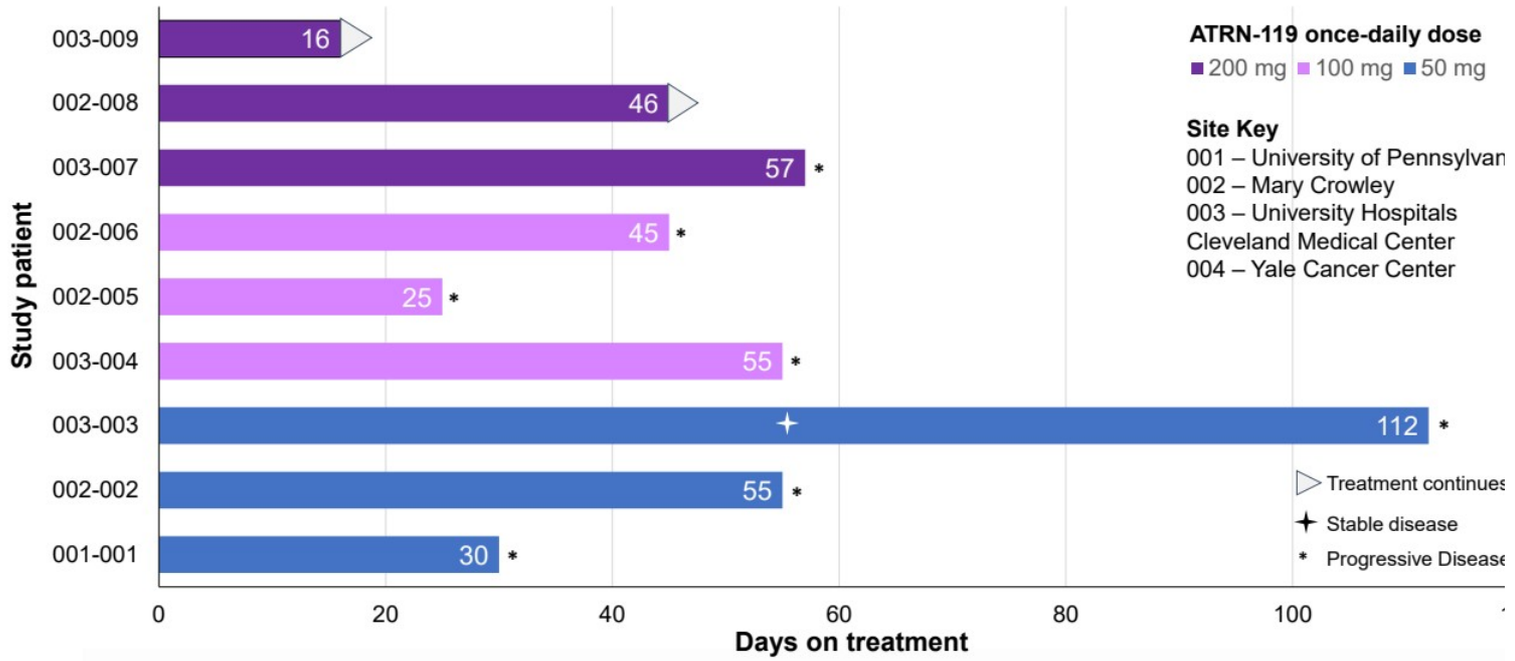
Secondary objectives:

- Antitumor activity (RECIST/PCWG3)

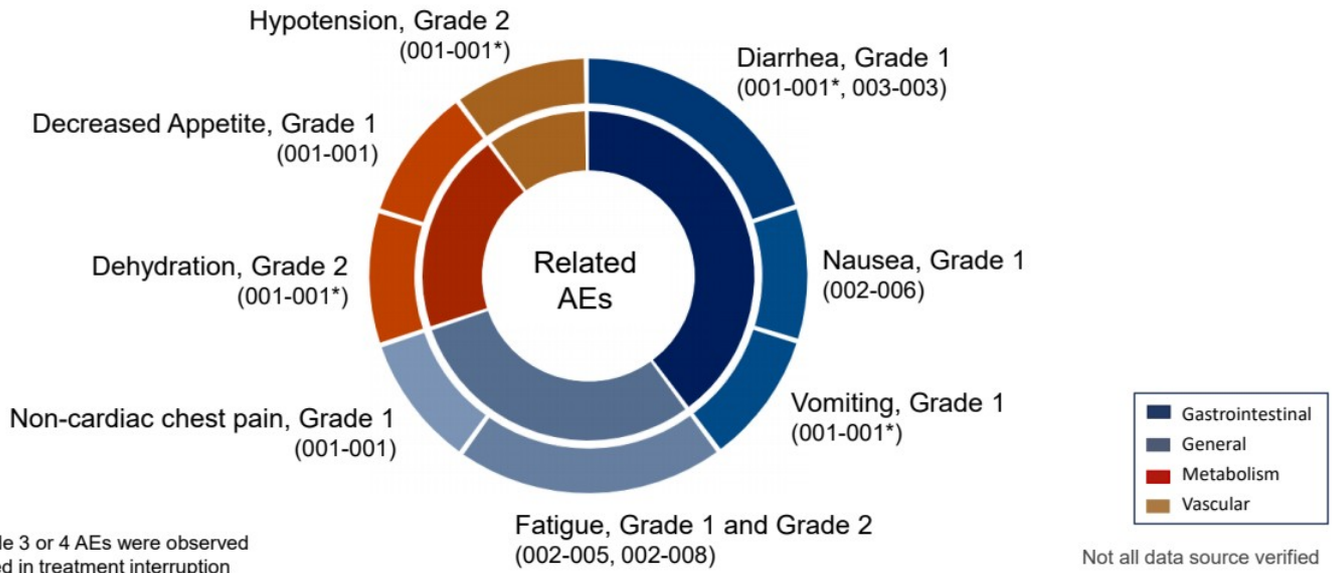
Exploratory objectives:

- Association between identified mutations and clinical outcomes





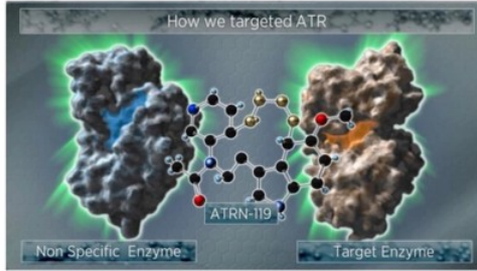
As of Sept 20, 2023: Five out of nine study patients have experienced AEs; assessed to be possibly/probably related to ATRN-119



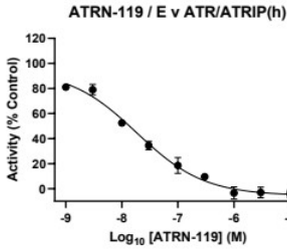
Milestone	Timeline
Phase 1/2a – Monotherapy Dose Escalation	
Preliminary clinical data	4Q 2023
Last Patient Enrolled	1Q 2024
Phase 1/2a – Monotherapy Dose Expansion	
First Patient Enrolled	2Q 2024
Last Patient Enrolled	2Q 2025
Phase 1/2a – Combination	
First Patient Enrolled	1H 2024
Last Patient Enrolled	2H 2025

ATR Inhibitor :
ATRN-119

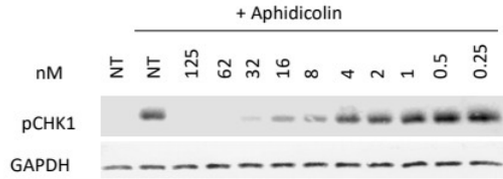
Preclinical
Proof-Of-Principal



Replication fork collapse
+ Double Strand Breaks

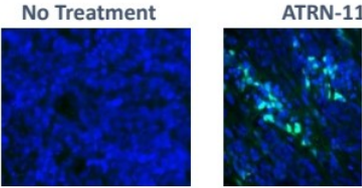


ATR-119 binds to ATR



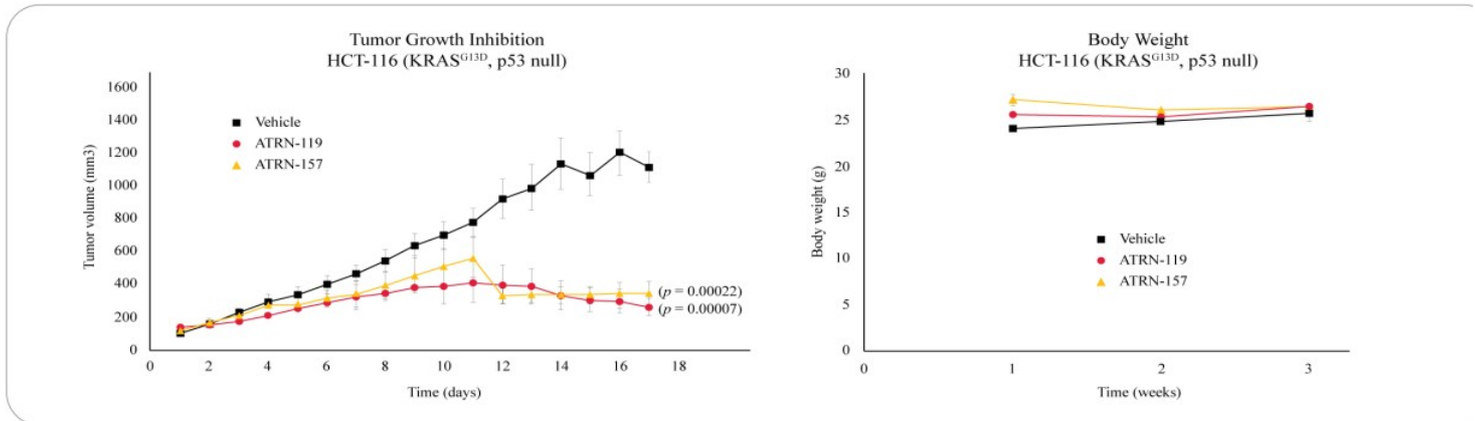
...inhibits its biological activity...

Tumor Samples Pharmacodynamics - γ H2A



...and triggers replication fork collapse and Double Strand Breaks

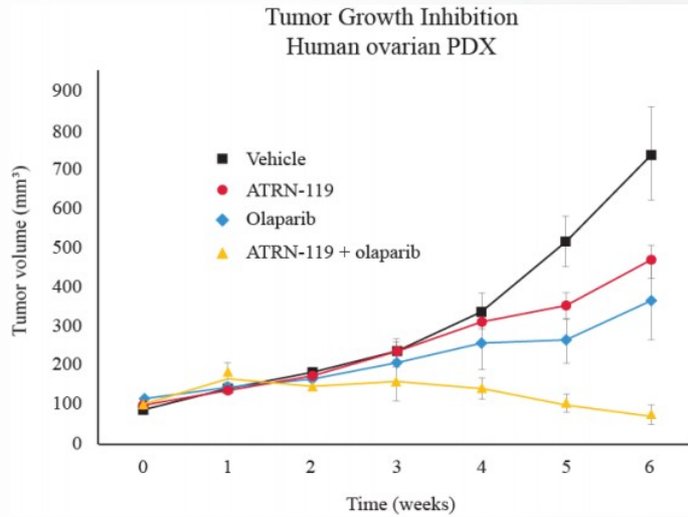
- Nanomolar potency in vitro across a broad spectrum of cancer cell lines
- Strong tumor control observed in vivo, including in challenging genetic backgrounds



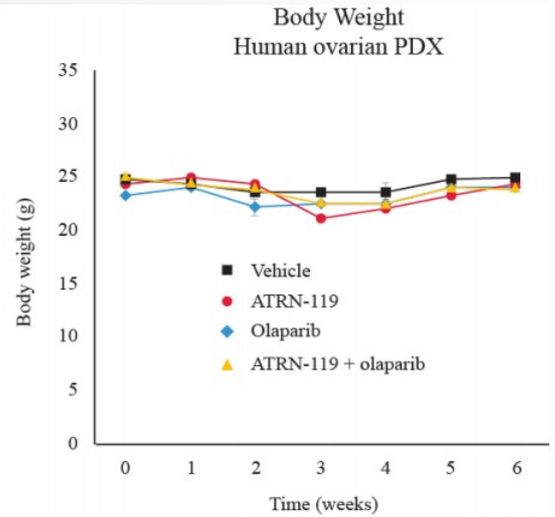
N=4 female mice per group, ATRN-119 - 100 mg/kg/day P.O, ATRN-157 - 20 mg/kg/day SQ.

ATRN-157 is an active metabolite identified in dogs receiving ATRN-119 P.O.. In vitro metabolism studies in dog and human hepatocytes and liver microsomes indicated formation of ATRN-157 in both species. Potency and selectivity of ATRN-157 was comparable to ATRN-119.

ATRN-119 + Olaparib Inhibits Ovarian Tumor Growth Over Time



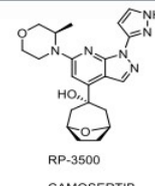
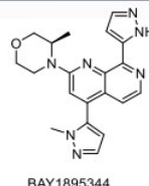
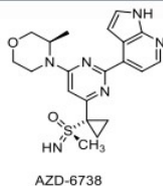
ATRN-119 + Olaparib Shows Negligible Weight Loss Over Time



N=6-8 mice per group, ATRN-119 - 90 mg/kg P.O BID, Olaparib - 50 mg/kg/day P.O, ATRN-119+Olaparib at the same doses and schedules

ATR Inhibitor :
ATRN-119

Potential Differentiation



Parameter	AstraZeneca AZD6738 ⁽¹⁾⁽²⁾	Bayer BAY1895344 ⁽³⁾	Repare / Roche ⁽⁴⁾ RP-3500 ⁽⁵⁾
Route Of Administration	Oral	Oral	Oral
Clinical Studies Chosen (MTD/RP2D), Dose Schedule	160mg BID, 2-weeks-on, 2-weeks-off, or: Continuous dosing ⁽¹⁾	40mg BID, 3-days-on/4-days-off	160mg QD, 3-days-on/4-days-off
Main Grade ≥3 Hematological toxicities reported at Chosen Dose Schedule (MTD/RP2D), in clinical studies	Patriot 1, Escalation Phase, 160mg, BID ⁽²⁾ : Anemia (1/6, 17%) Patriot 2, Expansion Phase ⁽¹⁾ : Fatigue, anemia, nausea & thrombocytopenia (not differentiated) ⁽¹⁾: (4/6, 67%) with continuous dosing (3/15, 20%) with 2-week-on, 2-week-off	Anemia (2/2, 100%) Neutropenia (1/2, 50%)	Anemia (23/95, 24%) Neutrophil count decreased (10/95, 11%) Platelet count decreased (5/95, 5%)

Note: Head-to-head studies with ATRN-119 have not been conducted

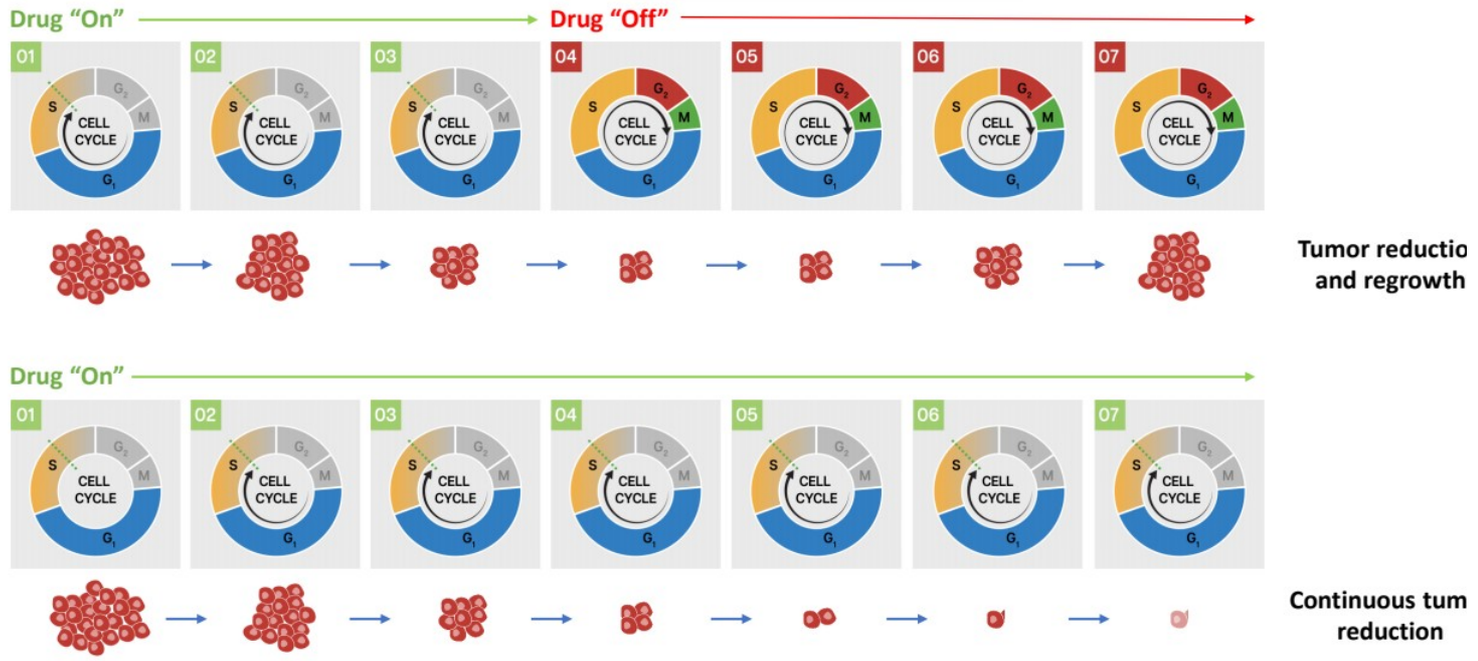
(1) Phase 1 study of ATR inhibitor, AZD6738, as monotherapy in advanced solid tumors (PATRIOT part A, B), Dillon et al, Volume 30, October 2019, Pages v165-v166

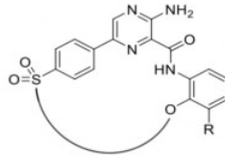
(2) Poster CT084: A Phase 1 dose-escalation study of ATR inhibitor monotherapy with AZD6738 in advanced solid tumors (PATRIOT Part A), AACR 2017

(3) First-in-Human Trial of the Oral Ataxia Telangiectasia and RAD3-Related (ATR) Inhibitor BAY 1895344 in Patients with Advanced Solid Tumors, Yap et al, Cancer Discov 2021;11:80-91 and 2019 ASCO Poster, De-Bono et al.

(4) Repare announced a worldwide license and collaboration agreement with Roche on June 1, 2022

(5) Preliminary Phase 1 Data From Ongoing First-in-Human Phase 1/2 TRESR Study of RP-3500, AACR 2022





Parameter	ATRN-119 ⁽¹⁾
Route Of Administration	Oral
Clinical Studies Chosen (MTD/RP2D), Dose Schedule	continuous once-daily dosing (dose TBD in Phase 1) ⁽¹⁾
Hematological toxicities in preclinical studies	<p>Pre-Clinical, Toxicology Studies:</p> <ul style="list-style-type: none"> In 28-day GLP tox study in dogs, hematological changes were of small magnitude with complete recovery In a head-to-head comparative tolerability study, ATRN-119 demonstrated significantly less toxicity than another oral ATRi that is currently in clinical development ⁽²⁾

ATRN-119 potential for reduced toxicity could make it a preferred ATR inhibitor as a single agent, as well as a candidate for combination with standard-of-care therapies.

Note: ATRN-119 has not yet been tested clinically

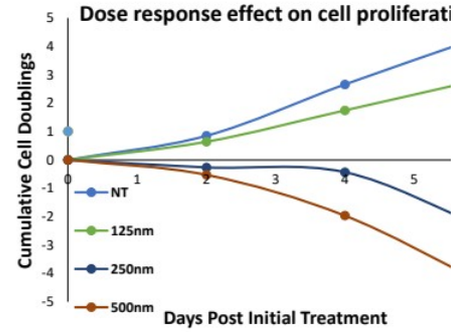
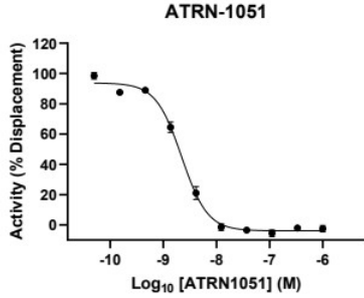
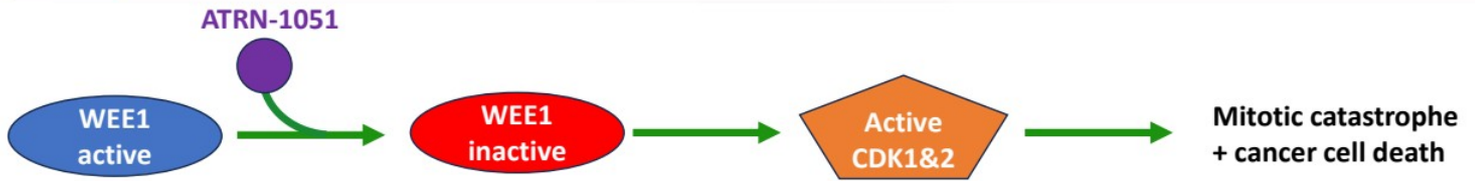
(1) ATRN-119, Phase 1/2a Clinical Study Protocol

(2) Internal pre-clinical head-to-head tolerability study in male beagle dogs. ATRN-119 - 20 mg/kg/day for 7 days, followed by 40 mg/kg/day for 7 days and finally 50 mg/kg/day for 7 days, all P.O. Competitor ATRi- administered at a clinically equivalent dose range during 21 days, P.O. By day 4 of dosing, the dog receiving the competitor ATRi exhibited severe reduction in multiple blood cell lineages including reticulocytes. Continued dosing of competitor ATRi for three weeks resulted in significant reduction of white blood cells, red blood cells and hemoglobin levels, and was accompanied by severe body weight loss (-15%). For the dog receiving ATRN-119, reduced levels of reticulocytes and neutrophils were noted with prolonged treatment but remained within normal ranges and body weight changes were negligible (-2% to +4%).



WEE1 Inhibitor : ATRN-1051

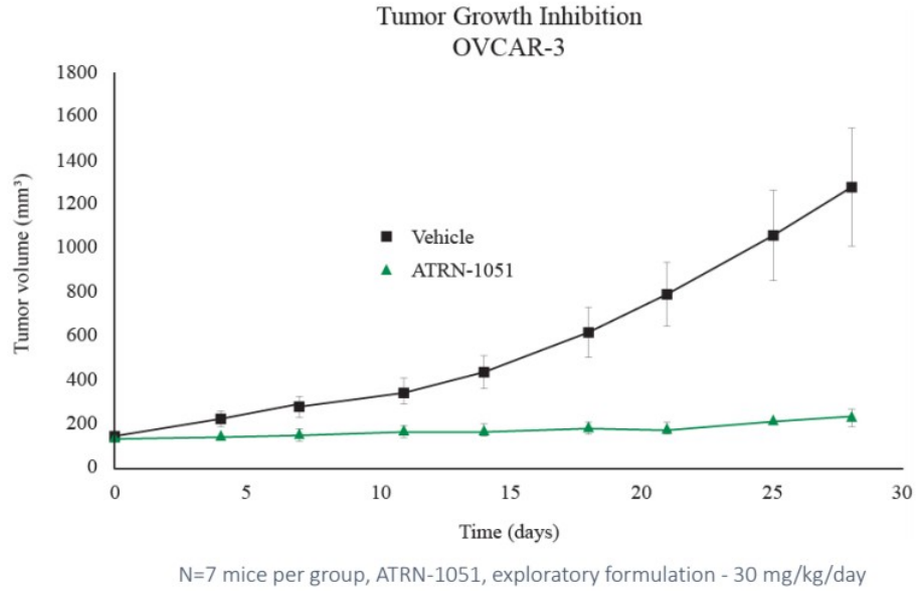
Preclinical proof-of-principle



ATRN-1051 binds to WEE1

...inhibits its biological activity...

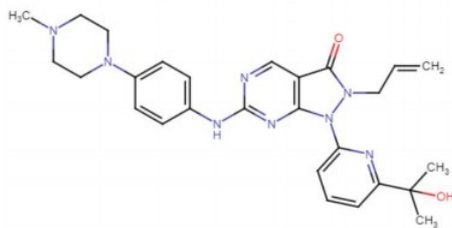
...and triggers mitotic catastrophe and cancer cell death



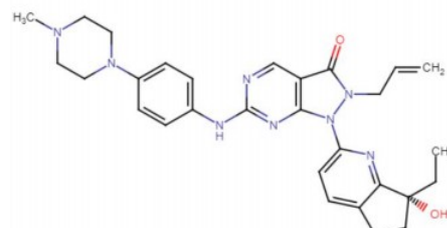
WEE1 Inhibitor : ATRN-1051

Potential Differentiation

ATRN-1051 shows potential to be potent and structurally differentiated, with high selectivity to limit off-target toxic



AZD-1775⁽¹⁾



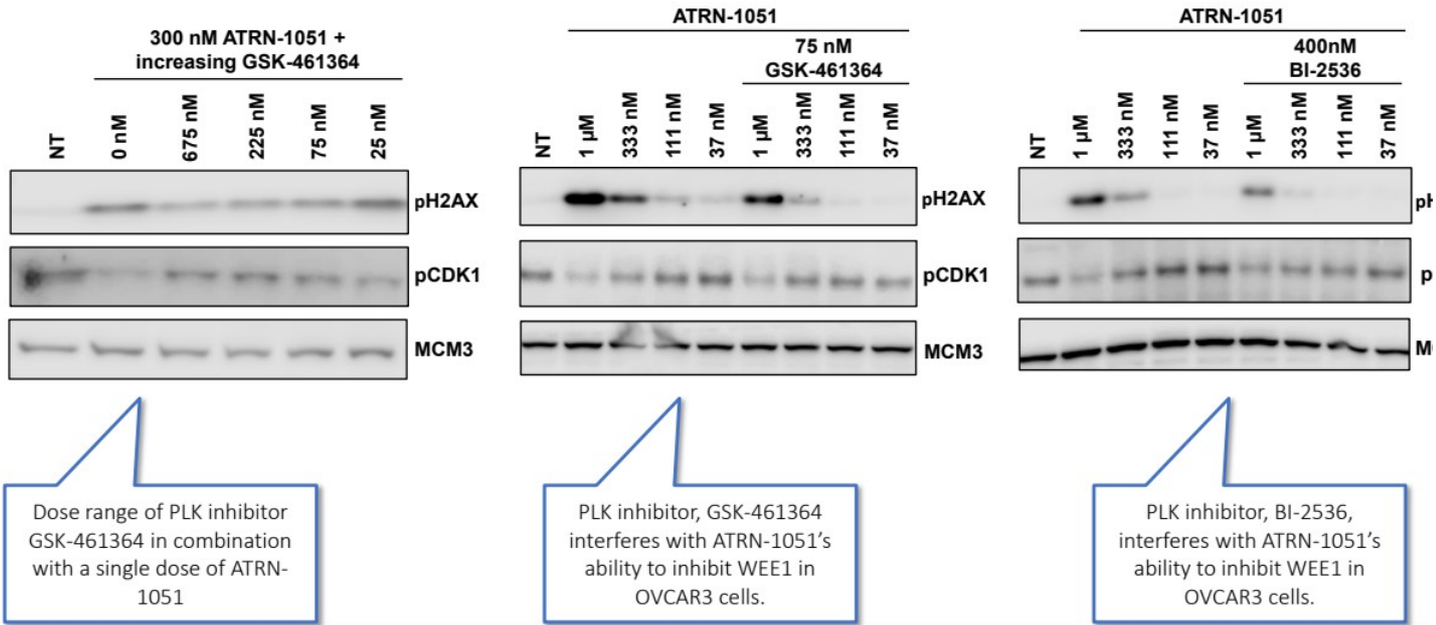
Azenosertib (ZN-c3)

	On-Target IC ₅₀ (nM)	Off-Target Inhibition at 1 μM (%)		
	WEE1	PLK1	PLK2	PLK3
Aprea: ATRN-1051	2.2	17	33	12
Zentalis: Azenosertib (ZN-c3) ⁽¹⁾	3.8	79	96	92
AstraZeneca: AZD-1775 ⁽¹⁾⁽²⁾	3.9	70	101	91

Note: Head-to-head studies have not been conducted

(1) Huang et al, (2021) J Med Chem

(2) AstraZeneca announced in July 2022 the discontinuation of development of AZD-1775



Based on pre-clinical studies, ATRN-1051 shows potentially favorable drug exposure:



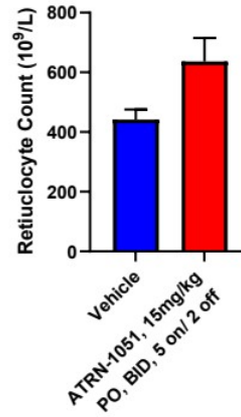
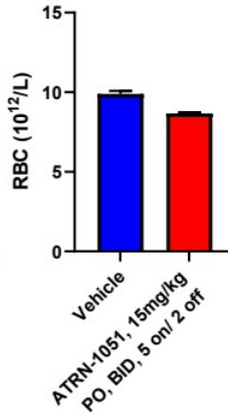
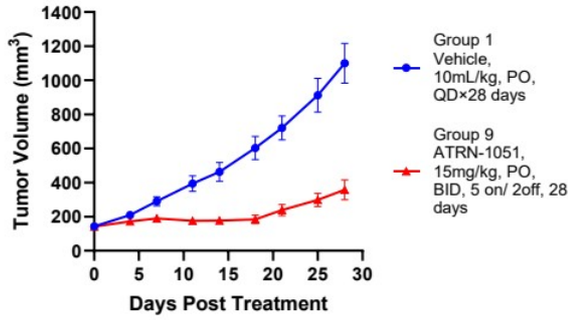
	ATRN 1051 ⁽¹⁾	Zentalis Azenosertib (ZN-c3) ⁽²⁾				AstraZeneca AZD-1775 ⁽²⁾		
Dose (mg/kg/d)	10	20	40	80	20	40	80	
C _{max} ng/ml	1,460	1,167	1,997	5,100	635	2,460	4,703	
T _{max} hr	3	1	1	1	1	1	1	
AUC _{0-24h} ng*hr/ml	16,739	4,863	17,088	39,722	1,494	6,313	13,408	

Note: Head-to-head studies have not been conducted

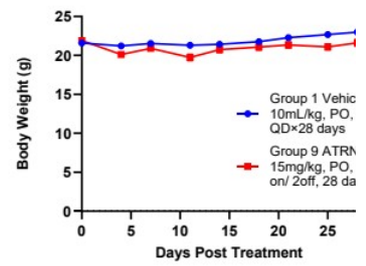
(1) Data from an exploratory formulation of ATRN-1051 administered to fasted Balb/c mice

(2) Data from study in A-427 NSCLC xenograft model as reported in Zentalis Corporate Overview, March 2022

OVCAR3 Xenograft Tumor Model in Female Nude Mice
Tumor Volume (mm³) (Mean±SEM)



OVCAR3 Xenograft Tumor Model in Female N
Body Weight (g) (Mean±SEM)



In vitro kinase assays		Average WEE1 kinase IC50	hERG inhibition		Average hERG IC50	Fold difference between kinase and hERG inhibition
LanthaScreen (Thermo)	Hotspot (Reaction Biology)	21.8 nM	<u>HEK293 cells (Medicilon)</u>	<u>CHO cells (WuXi)</u>	4,750 nM	<u>hERG inhibition over WEE1 kinase inhibition</u>
2.2 nM	41.4 nM		8,840 nM	660 nM		218-fold (range 16- to 3,946-fold)

Milestone	Timeline
Preclinical proof-of-principle	
Additional differentiation data	4Q 2023
IND	
Submission	4Q 2023
Clearance	1Q 2024
Phase 1/2a – Monotherapy Dose Escalation	
First Patient Enrolled	1H 2024
Last Patient Enrolled	2H 2025
Phase 1/2a – Combination	
First Patient Enrolled	2H 2024
Last Patient Enrolled	2H 2025



Summary

- Cash & Equivalents of \$27.7 million as of June 30, 2023
- Closed \$4.9M (net) public offering in February 2023
- Obtained \$2.0 million non-dilutive funding via research grant from National Cancer Institute (NCI)

Securities	Common Equivalents as of Aug. 10, 2023
Preferred Stock (as converted)	28,112
Common Stock	3,736,673
Options	558,141
Restricted Stock Units	20,870
Fully Diluted Equivalents	4,343,796

- Diversified portfolio with de-risked clinical and preclinical plans underway

- Opportunities in ovarian, CRC, prostate and breast cancers
 - ◇ Single agent and combination therapies

- Supportive follow-on strategy
 - ◇ IND submission by end of 2023
 - ◇ Undisclosed DDR asset

- Financed into Q4 2024
 - ◇ Reach short term inflection points and catalysts
 - ◇ Evaluate optimal strategic partnerships