

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

August 12, 2026

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: **(215) 948-4119**

(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 2.02 Results of Operations and Financial Condition.

On August 12, 2026, Aprea Therapeutics, Inc. (the “Company”) issued a press release announcing its financial results for the three and six months ended June 30, 2026, and provided an update on the Company’s operations for the same period. The Company is furnishing a copy of the press release, which is attached hereto as Exhibit 99.1.

In accordance with General Instruction B.2 of Form 8-K, the information included in this Item 2.02, including Exhibit 99.1 hereto, shall not be deemed “filed” for the purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the “Exchange Act”), or otherwise subject to the liabilities of that section, nor shall it be deemed incorporated by reference into any filing made by the Company under the Exchange Act or Securities Act of 1933, as amended, except as shall be expressly set forth by specific reference in such a filing.

Item 8.01 Other Events.

On August 12, 2026, the Company updated its corporate presentation slide deck. A copy of the corporate presentation slide deck is filed as Exhibit 99.2 hereto and incorporated herein by reference.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit Number	Description
99.1	Press release issued by Aprea Therapeutics, Inc. dated August 12, 2026.
99.2	Corporate Presentation (August 2026).
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: August 12, 2026

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

Aprea Therapeutics Announces Second Quarter 2026 Financial Results

Enrollment in the ongoing Phase 1 dose escalation ACESOT-1051 trial of oral WEE1 inhibitor APR-1051 is accelerating as dose escalation advances

Aprea plans to advance APR-1051 in uterine serous carcinoma (USC), and Cyclin E-overexpressing platinum-resistant ovarian cancer (PROC) while also evaluating combination regimens with standard of care in HPV-positive head and neck cancer and colorectal cancer

Updated Phase 1 data presented at ASCO 2026 showed early single-agent activity for APR-1051, including partial responses and disease stabilization, with a manageable tolerability profile

\$41.2 million in cash and cash equivalents as of June 30, 2026

DOYLESTOWN, PA, August 12, 2026 (GLOBE NEWSWIRE) — Aprea Therapeutics, Inc. (Nasdaq: APRE) (“Aprea”, or the “Company”), a clinical-stage precision medicine oncology company focused on the discovery and development of targeted therapies for patients with biomarker-defined cancers,

today reported financial results for the second quarter ended June 30, 2026, and provided a business update.

"Having observed early signs of clinical activity for APR-1051, we are now expanding the ongoing ACESOT-1051 trial," said Oren Gilad, Ph.D., President and Chief Executive Officer of Aprea. "We are accelerating enrollment, advancing dose escalation to inform dose selection, and preparing to evaluate APR-1051 both as a single agent in uterine serous carcinoma (USC) and Cyclin E-overexpressing platinum-resistant ovarian cancer (PROC) and also in combination with standard of care in HPV-positive head and neck cancer and colorectal cancer. We look forward to sharing our next clinical data update at a medical meeting in the fourth quarter of 2026. This ongoing progress reflects our commitment to developing targeted cancer therapies that have the potential to improve outcomes and quality of life for patients, while also creating value for our shareholders."

Key Business Updates and Upcoming Key Milestones

ACESOT-1051: A Biomarker-Focused, Phase 1 Trial of Oral WEE1 Inhibitor, APR-1051

- APR-1051 is a potent and selective, oral small molecule WEE1 inhibitor designed to potentially address therapeutic window limitations observed with earlier WEE1 programs. It is currently being evaluated in ACESOT-1051, a multi-center, open-label Phase 1 study. The primary objectives of this study are safety, dose-limiting toxicity, maximum tolerated or maximum administered dose, and
-

RP2D. Secondary objectives include pharmacokinetics and antitumor activity assessed by RECIST/PCWG3.

- **Aprea expects to report the next clinical data update from ACESOT-1051 at a medical meeting in the fourth quarter of 2026.** Enrollment is accelerating ahead of this anticipated clinical catalyst, with the number of active clinical sites expanding from three to ten. The Company expects enrollment to reach 6 to 10 patients per month by Q4 of 2026, potentially increasing the pace of clinical data generation.
- Supported by the \$30 million private placement that closed in the first quarter of 2026, the Company is expanding enrollment in ACESOT to include at least 50 patients with uterine serous carcinoma or cyclin E-overexpressing, platinum-resistant ovarian cancer. Completion of dose escalation and backfill expansion is anticipated in the second quarter of 2027. This expansion is intended to further characterize the clinical activity of APR-1051 in biomarker-defined tumor populations with a mechanistic rationale for WEE1 inhibition.
- Aprea also plans to evaluate APR-1051 in combination settings, supported by preclinical synergy observed in relevant disease models. In HPV-positive head and neck squamous cell carcinoma, APR-1051 will be combined with immune checkpoint therapy. In colorectal cancer, APR-1051 will be combined with a standard-of-care chemotherapy. Advancing clinically evaluated, active doses are intended to support the tolerability and dosing of APR-1051 when added to standard-of-care treatment.
- For more information on ACESOT-1051, refer to ClinicalTrials.gov NCT06260514.

APR-1051 Presentation at ASCO 2026

- On May 30, 2026, Aprea presented updated data from ACESOT-1051 in a poster titled “Early results from the first-in-human phase 1 study of WEE1 inhibitor APR-1051 in patients with advanced solid tumors (ACESOT-1051)” (Abstract #3107) at the American Society of Clinical Oncology (ASCO) 2026 Annual Meeting in Chicago, Illinois. The presentation summarized data as of a May 6, 2026 cutoff. A copy of the poster can be found on the Aprea corporate website [here](#).

ATR inhibitor, ATRN-119

- ATRN-119 is a potent and highly selective potentially first-in-class macrocyclic ATR inhibitor designed for patients with tumors harboring mutations in DDR-related genes. Cancers with mutations in DDR-related genes represent a high unmet medical need, and these patients often have a poor prognosis and currently lack effective therapeutic options.
 - During 2025, Aprea determined the recommended Phase 2 monotherapy dose (RP2D) of 1,100 mg once daily for ATRN-119 in the ABOYA-119 Phase 1/2a dose-escalation study and subsequently closed this study to focus resources on the clinical development of APR-1051. Building on the completion of dose escalation, the Company is considering further ATRN-119 development in combination approaches that could expand its therapeutic potential. Aprea believes ATRN-119's mechanism of action, potentially favorable safety profile, and pharmacologic characteristics could make it an ideal candidate for combination with other anti-cancer therapies, including radiation therapy, chemotherapy, antibody-drug conjugates (ADCs) and immune checkpoint inhibitors.
 - Aprea is currently in discussions with leading academic centers to explore various combinations for ATRN-119. These include investigator-initiated studies evaluating ATRN-119 in combination with I/O
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agents, chemotherapy, ADCs and/or radiation. Potential indications for these combinations include advanced solid tumors (e.g. HPV+ head and neck cancers, sarcomas, ovarian, colorectal, lung) and hematologic malignancies (e.g. Acute Myeloid Leukemia, Myelodysplastic syndromes).

Pipeline

- Aprea also has an early-stage program, a macrocyclic DYRK1A/B inhibitor, that potentially could enter IND enabling studies in the fourth quarter of 2026, subject to available resources.

Financial Results for the Second Quarter Ended June 30, 2026

- As of June 30, 2026, the Company reported cash and cash equivalents of \$41.2 million, compared to \$14.6 million as of December 31, 2025. The Company believes that its cash and cash equivalents as of June 30, 2026 will be sufficient to meet its currently projected operating expenses and capital expenditure requirements into the first quarter of 2028.
- For the second quarter ended June 30, 2026, the Company reported an operating loss of \$4.0 million, compared to an operating loss of \$3.4 million in the second quarter of 2025.
- Research and Development (R&D) expenses were \$2.5 million for the quarter ended June 30, 2026, compared to \$1.9 million for the second quarter of 2025. The increase in R&D expense was primarily related to higher expenses in ACESOT-1051, our Phase 1 dose-escalation study for APR-1051, partially offset by lower expense in the ABOYA-119 clinical trial to evaluate ATRN-119, which has been closed.
- General and Administrative (G&A) expenses were \$1.6 million for each of the quarters ended June 30, 2026 and 2025.
- The Company reported a net loss of \$3.6 million, or \$(0.07) per basic share, on approximately 53.5 million weighted-average common shares outstanding for the quarter ended June 30, 2026, compared to a net loss of \$3.2 million, or \$(0.53) per basic share, on approximately 6.1 million weighted-average common shares outstanding for the comparable period in 2025.

About Aprea

Aprea is a clinical-stage precision medicine oncology company focused on the discovery and development of targeted therapies for patients with biomarker-defined cancers. The Company is pioneering a new approach to treat cancer by exploiting vulnerabilities associated with cancer cell mutations. This approach was developed to kill tumors while minimizing the effect on normal, healthy cells. Aprea's technology has potential applications across multiple cancer types, enabling it to target a range of tumors, including ovarian, endometrial, colorectal and head and neck squamous cell carcinoma. The company's lead programs are APR-1051, an oral, small-molecule inhibitor of WEE1 kinase, and ATRN-119, a small molecule ATR inhibitor, both in clinical development for solid tumor indications. For more information, please visit the company website at www.aprea.com.

The Company may use, and intends to use, its investor relations website at <https://ir.aprea.com/> as a means of disclosing material nonpublic information and for complying with its disclosure obligations under Regulation FD.

Forward-Looking Statement

Certain information contained in this press release 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 study analyses, clinical trials, regulatory submissions, development plans and projected cash position. We may, in some cases, use terms such as “future,” “predicts,” “believes,” “potential,” “continue,” “anticipates,” “estimates,” “expects,” “plans,” “intends,” “targeting,” “confidence,” “may,” “could,” “might,” “likely,” “will,” “should” or other words that convey uncertainty of future events or outcomes to identify these forward-looking statements. Our forward-looking statements are based on current beliefs and expectations of our management team and on information currently available to management, and involve risks, potential changes in circumstances, assumptions and uncertainties. All statements contained in this press release other than statements of historical fact are forward-looking statements, including statements regarding our ability to develop, commercialize and achieve market acceptance of our current and planned product candidates, our research and development efforts, including timing considerations and other matters regarding our business strategies, use of capital, results of operations and financial position, projected cash runway, plans and objectives for future operations, the timing, progress, enrollment and completion of our ongoing and planned clinical trials, the timing of expected clinical data updates and medical meeting presentations, the potential expansion of ACESOT-1051, the potential development of APR-1051 in single-agent and combination settings, the potential further development of ATRN-119, and the potential advancement of our early-stage programs. Any or all of the forward-looking statements may turn out to be wrong or be affected by inaccurate assumptions we 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, timing and cost of our ongoing and anticipated clinical trials for our current product candidates, including the timing of initiation, pace of enrollment and completion of trials, our ability to fully fund our disclosed clinical trials, which assumes no material changes to our currently projected expenses, the receipt and timing of interim or preliminary results, which are not necessarily indicative of final results, our understanding of product candidate mechanisms of action and interpretation of preclinical and early clinical results from our clinical development programs, our ability to predict clinical outcomes based on such preclinical and early clinical results, the timing and outcome of regulatory interactions, and the other risks, uncertainties and factors described under “Risk Factors,” “Management’s Discussion and Analysis of Financial Condition and Results of Operations” and elsewhere in the documents we file with the U.S. Securities and Exchange Commission. 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 press release. We undertake no obligation to update such forward-looking statements for any reason, except as required by law.

Investor Contact:

Mike Moyer
LifeSci Advisors
mmoyer@lifesciadvisors.com

**Aprea Therapeutics, Inc.
Consolidated Balance Sheets**

	June 30, 2026	December 31, 2025
	(unaudited)	
Assets		
Current assets:		
Cash and cash equivalents	\$ 41,233,428	\$ 14,599,347
Prepaid expenses and other current assets	679,686	961,899
Total current assets	41,913,114	15,561,246
Property and equipment, net	64,033	59,807
Restricted cash	41,615	41,186
Other noncurrent assets	271,162	271,162
Total assets	\$ 42,289,924	\$ 15,933,401
Liabilities and Stockholders' Equity		
Current liabilities:		
Accounts payable	\$ 935,139	\$ 713,668
Accrued expenses	2,157,163	2,050,690
Total current liabilities	3,092,302	2,764,358
Commitments and contingencies		
Series A convertible preferred stock, \$0.001 par value, 40,000,000 shares authorized; 31,194 shares issued and outstanding at June 30, 2026 and December 31, 2025	727,361	727,361
Stockholders' equity:		
Common stock, \$0.001 par value, 400,000,000 shares authorized, 12,391,136 and 8,192,538 shares issued and outstanding at June 30, 2026 and December 31, 2025, respectively	12,391	8,192
Additional paid-in capital	389,596,149	356,709,645
Accumulated other comprehensive loss	(10,617,617)	(10,634,714)
Accumulated deficit	(340,520,662)	(333,641,441)
Total stockholders' equity	38,470,261	12,441,682
Total liabilities and stockholders' equity	\$ 42,289,924	\$ 15,933,401

Aprea Therapeutics, Inc.
Consolidated Statements of Operations and Comprehensive Loss
(Unaudited)

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Grant revenue	\$ —	\$ 118,111	\$ —	\$ 280,574
Operating expenses:				
Research and development	2,462,042	1,912,213	4,073,209	4,395,279
General and administrative	1,570,138	1,593,671	3,389,383	3,358,650
Total operating expenses	4,032,180	3,505,884	7,462,592	7,753,929
Loss from operations	(4,032,180)	(3,387,773)	(7,462,592)	(7,473,355)
Other income (expense):				
Interest income, net	358,923	178,027	493,707	382,753
Other income	79,000	—	79,000	—
Foreign currency gain (loss)	2,953	(29,124)	10,664	(80,927)
Total other income	440,876	148,903	583,371	301,826
Net loss	<u>\$ (3,591,304)</u>	<u>\$ (3,238,870)</u>	<u>\$ (6,879,221)</u>	<u>\$ (7,171,529)</u>
Other comprehensive loss:				
Foreign currency translation	8,083	(1,681)	17,097	(1,038)
Total comprehensive loss	<u>\$ (3,583,221)</u>	<u>\$ (3,240,551)</u>	<u>\$ (6,862,124)</u>	<u>\$ (7,172,567)</u>
Net loss per share attributable to common stockholders, basic and diluted	<u>\$ (0.07)</u>	<u>\$ (0.53)</u>	<u>\$ (0.20)</u>	<u>\$ (1.19)</u>
Weighted-average common shares outstanding, basic and diluted	<u>53,483,504</u>	<u>6,083,329</u>	<u>34,191,653</u>	<u>6,038,845</u>



APREA
THERAPEUTICS
WE CAN'T WAIT TO CURE CANCER

A clinical-stage precision
medicine oncology company
focused on the discovery and
development of targeted
therapies for patients with
biomarker-defined cancers

August 2026



Forward-Looking Statements

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 study analyses, clinical trials, regulatory submissions, and projected cash position. We may, in some cases use terms such as “future,” “predicts,” “believes,” “potential,” “continue,” “anticipates,” “estimates,” “expects,” “plans,” “intends,” “targeting,” “confidence,” “may,” “could,” “might,” “likely,” “will,” “should” or other words that convey uncertainty of the future events or outcomes to identify these forward-looking statements. Our forward-looking statements are based on current beliefs and expectations of our management team and on information currently available to management that involve risks, potential changes in circumstances, assumptions, and uncertainties. All statements contained in this presentation other than statements of historical fact are forward-looking statements, including statements regarding our ability to develop, commercialize, and achieve market acceptance of our current and planned product candidates, our research and development efforts, including timing considerations and other matters regarding our business strategies, use of capital, results of operations and financial position, project cash runway, plans and objectives for future operations, the timing, progress, enrollment and completion of our ongoing and planned clinical trials, the timing of expected clinical data updates and medical meeting presentations, the potential expansion of ACESOT-1051, the potential development of APR-1051 in single-agent and combination settings, the potential further development of ATRN-119, and the potential advancement of our early-stage programs. Any or all of the forward-looking statements may turn out to be wrong or be affected by inaccurate assumptions we 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, timing and cost of our ongoing and anticipated clinical trials for our current product candidates, including the timing of initiation, pace of enrollment and completion of trials, our ability to fully fund our disclosed clinical trials, which assumes no material changes to our currently projected expenses, the receipt and timing of interim or preliminary results, which are not necessarily indicative of final results, our understanding of product candidate mechanisms of action and interpretation of preclinical and early clinical results from our clinical development programs, our ability to predict clinical outcomes based on such preclinical and early clinical results, the timing and outcome of regulatory interactions, and the other risks, uncertainties, and factors described under “Risk Factors,” “Management’s Discussion and Analysis of Financial Condition and Results of Operations” and elsewhere in the documents we file with the U.S. Securities and Exchange Commission. 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 for any reason, except as required by law. This presentation shall not constitute an offer to sell or the solicitation of an offer to buy any securities, nor shall there be any sale of any securities in any state or jurisdiction in which such offer, solicitation or sale would be unlawful prior to registration or qualification under the securities laws of any such state or jurisdiction. This presentation may not be reproduced, forwarded to any person or published, in whole or in part.

APR-1051 and ATRN-119 are investigational drugs and have not yet been approved by the U.S. Food and Drug Administration or any other regulatory authority.



Aprea Therapeutics (NASDAQ: APRE) One Critical Pathway - Multiple Targets



Positioned at the forefront of synthetic lethality and precision medicine

Targeted Oncology

Transition from broad, toxic chemotherapy to potentially safer, precision-guided targeted therapies

Precision-Driven Development

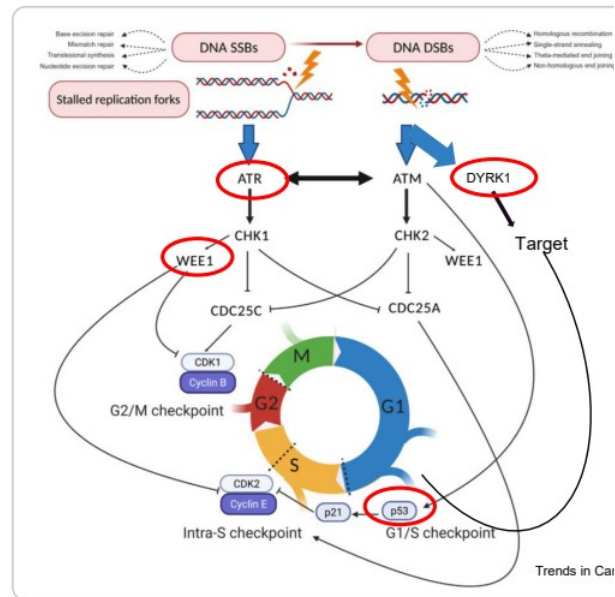
Develop highly selective cancer therapies that exploit tumor-specific mutations to maximize cancer cell killing while sparing healthy tissue

Pipeline with Clinical Momentum

All programs are designed to address significant unmet medical needs across genetically defined cancer populations

Early Clinical Proof-of-Concept

Preliminary signals of monotherapy activity observed in ongoing Phase 1 trial

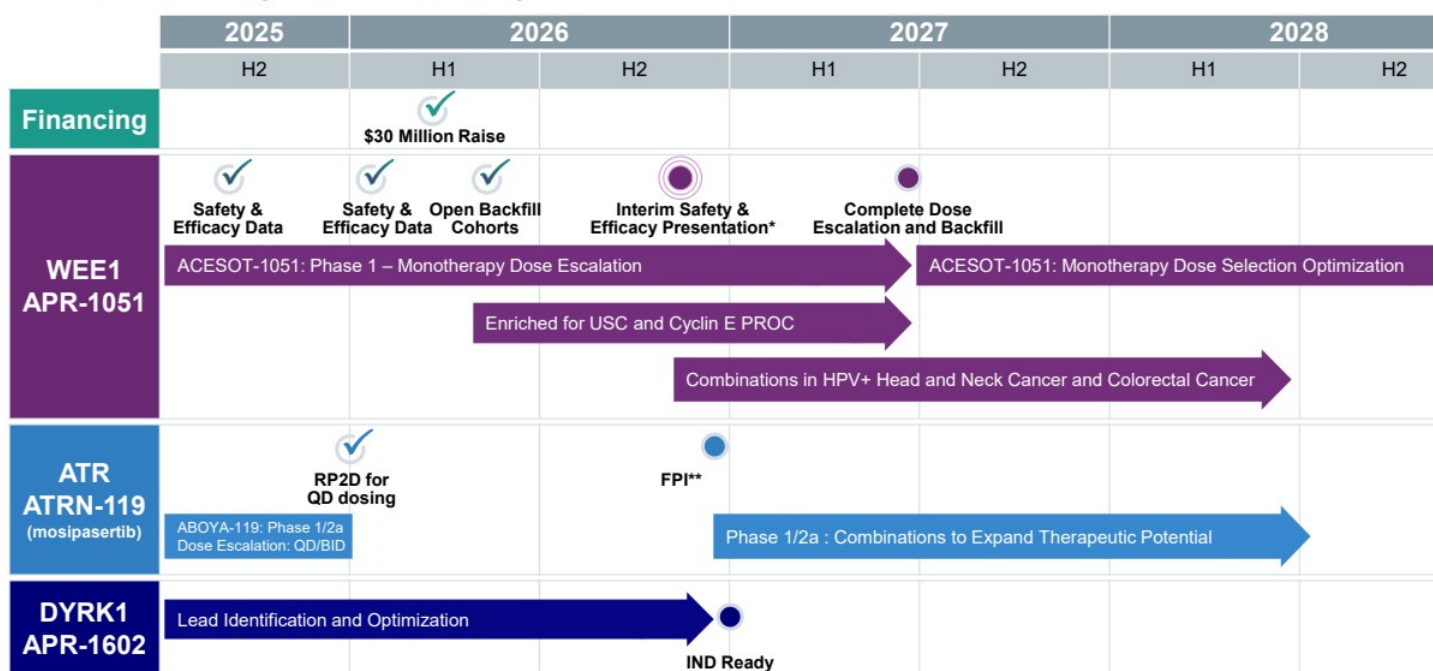


1. Ngoi N, *et al.* Targeting the replication stress response through synthetic lethal strategies in cancer medicine. *Trends in Cancer*. (2021); 7(10):930-957

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Robust DDR Development Pipeline and Corporate Milestones

2025-2028 Accomplished and anticipated milestones



DNA Damage Response (DDR)

*Planned major medical conference presentation

**FPI contingent upon execution of a research collaboration or partnering agreement; study initiation is expected to be externally supported without use of current company resources.

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Strong Drug Development and Commercial Expertise

Experienced team in synthetic lethality and targeted therapy

Management

Oren Gilad, PhD President and CEO	John P. Hamill SVP and CFO	Eugene Kennedy, MD Chief Medical Advisor	Ze'ev Weiss, CPA, BSc Chief Business Advisor	Mike Carleton, PhD Translational Medicine Advisor	Brian Wiley SVP, Corporate Strategy

Board of Directors

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



**APREA
THERAPEUTICS**
WE CAN'T WAIT TO CURE CANCER

WEE1 Inhibitor: APR-1051

ACESOT-1051:

Clinical Proof-Of-Concept

WEE1 Has Emerged as a Therapeutic Target of Significant Industry Interest

Clinically validated, prior WEE1 inhibitors have been challenged by narrow therapeutic windows
Aprea is applying key insights to advance APR-1051 as a potentially best-in-class WEE1 inhibitor

Program	Clinical Limitations	Strategic Outcome	What It Signals
Adavosertib (AstraZeneca)	Hematologic & GI toxicity limited dose intensity	Terminated further clinical development Returned by AstraZeneca to Merck & Co.	Biology works, narrow therapeutic window
Azenosertib (Zentalis)	Continuous dosing not tolerated ¹	TRAEs leading to dose reductions, interruptions and discontinuation	Biology works, therapeutic window still being defined
Debio 0123 (Debiopharm)	QT prolongation liability at high doses ²	Limited single-agent activity – no responses up to MTD ³ ; activity seen in combination with PKMYT1 inhibitor ⁴	Potential future as combination agent
Program	Engineered Profile	Strategic Outcome	What It Signals
APR-1051 (Aprea Therapeutics)	Structurally differentiated, highly potent, limited off-target inhibition	Preliminary signals of monotherapy activity without class-limiting tox to date	Potential to expand therapeutic window

No head-to-head studies have been conducted. Trial information is based on publicly available data and should be interpreted cautiously



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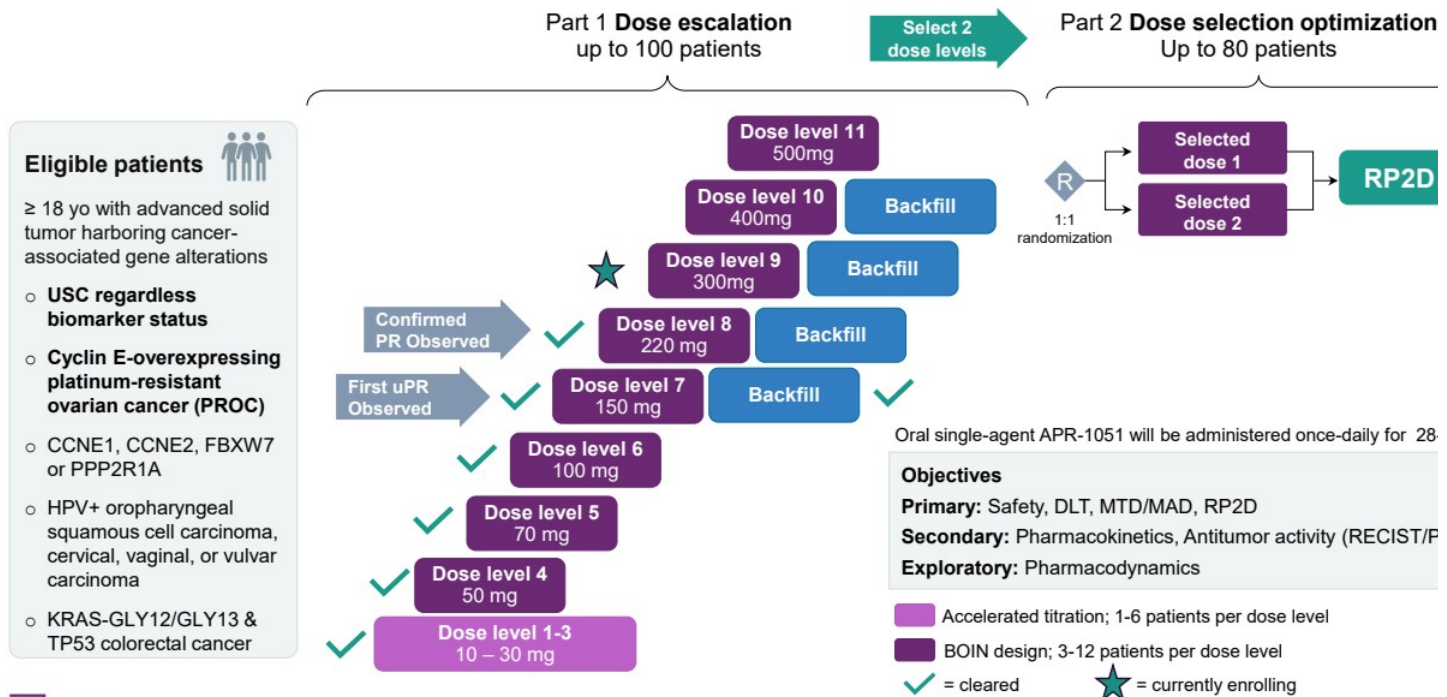
TRAEs – Treatment related adverse events

1. Zentalis Corporate Presentation, April 2026
2. Debio 0123-101, A Phase 1 Trial of Debio 0123 In Combination With Carboplatin In Advanced Solid Tumors: Safety, Pharmacokinetic, And Preliminary Antitumor Activity Data, Poster ASCO 2023
3. Abstract 3120, ASCO 2024
4. Poster CT022, AACR 2026

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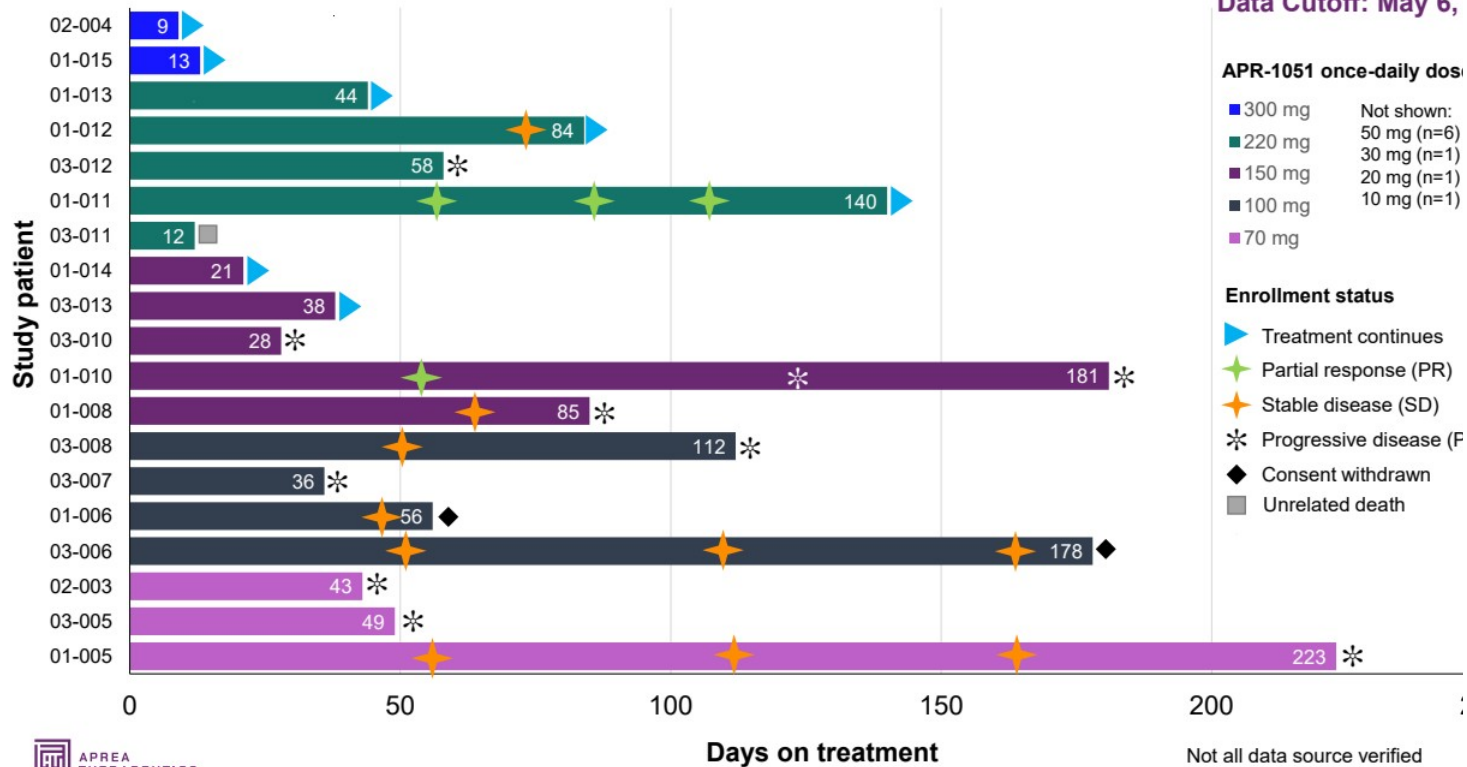
ACESOT-1051: Phase 1 Study Design

Multi-center, open-label Phase 1 single-agent dose escalation and dose selection optimization



APR-1051 Summary of Duration of Treatment (n=28 for all dose levels)

Data Cutoff: May 6,



Treatment-related AEs in Patients Treated with APR-1051 (N=28)

Data Cutoff: May 6, 2026

MedDRA Preferred Term	APR-1051 All dose levels (N=28)	
Treatment-related AEs, n (%) ^a	All Grades	Grade ≥ 3 ^b
Nausea	10 (35.7)	0 (0)
Fatigue	4 (14.3)	0 (0)
Vomiting	3 (10.7)	0 (0)
Alanine aminotransferase increased	1 (3.6)	1 (3.6) ^c
Anemia	1 (3.6)	0 (0)
Aspartate aminotransferase increase	1 (3.6)	1 (3.6) ^c
Blood bilirubin increased	1 (3.6)	0 (0)
Constipation	1 (3.6)	0 (0)
Dehydration	1 (3.6)	0 (0)
Dysgeusia	1 (3.6)	0 (0)
Dyspepsia	1 (3.6)	0 (0)
Eczema	1 (3.6)	0 (0)
Gastroesophageal reflux disease	1 (3.6)	0 (0)
Hypokalemia	1 (3.6)	0 (0)
Lymphocyte count decreased	1 (3.6)	1 (3.6)
Platelet count decreased	1 (3.6)	0 (0)

Confirmed Partial Response in Patient 01-011

220mg QD – Currently on treatment

Demographics: 63-year-old Black Female

Site: MD Anderson Cancer Center

Diagnosis: Uterine carcinosarcoma

Key Mutations: PPP2R1A

Treatment History (4 prior lines)

- Line 1: Carboplatin + Paclitaxel → 126 days, PD
- Line 2: Doxorubicin → 56 days, PD
- Line 3: Topotecan → 70 days, PD
- Line 4: Pembrolizumab + Lenvatinib → 5months, PD

APR-1051 – Treatment Outcome

- C1D1: Dec 18, 2025
- Current Status: **On treatment 140 days (C6D1) – May 6, 2026**
- Best Response: **Confirmed Partial Response (PR)**
 - PR (-50%) at first assessment Feb 10, 2026
 - Confirmed PR with additional -9.5% reduction from C3D1 Mar 10, 2026; scan April 8, 2026 showed 0% change
- Tumor marker: CA-125 reduction from BL 362.4 U/mL to C3D1 46.8 U/mL (87% decrease); 40.2 U/mL Mar 11, 2026; 53 U/ml April 08
- Adverse Events: C1D22 Gr 1 rash. C1D15 Gr1 thrombocytopenia , possibly related to IP; intermittent nausea Gr1 probably related; increase Gr1 unlikely related. No DLT

Disease Control Observed in Early Patient Outcomes

APR-1051 shows preliminary evidence of disease control in rectal cancer with mutated FBXW7

100 mg Cohort Case Report

Stable disease maintained for 178 days in patient with FBXW7 mutation (100 mg QD) (elected to stop study participation)

Patient: 86-year-old Asian Female

Diagnosis: Rectal Cancer

Key Mutations: FBXW7 (Drives Cyclin E accumulation and overexpression)

Treatment History: 5 prior lines - heavily pretreated

- **Line 1:** Capecitabine/oxaliplatin → 191 days, PD
- **Line 2:** Capecitabine/oxaliplatin/bevacizumab → 45 days, PD
- **Line 3:** FOLFIRI + bevacizumab → 43 days, PD
- **Line 4:** Local XRT (lung mets) → 12 days, not evaluable
- **Line 5:** Tretinoin/bevacizumab/Tecentriq (ATRTR trial) → 50 days, PD

APR-1051 Activity:

- **Current Status:** Consent withdrawn after 178 days
- **Best Response:** SD at third scan (-15% tumor response)

Notes: Durable SD maintained 181 days in a heavily pretreated 86-year-old patient; well tolerated with minimal toxicity. FBXW7 mutation may be relevant to response

Disease Control Observed in Early Patient Outcomes

APR-1051 shows preliminary evidence of disease control in HPV+ head and neck cancer

70 mg Cohort Case Report

Stable disease maintained for 223 days in patient 01-005 HPV+ head and neck cancer (70 mg QD) (PD)

Patient: 62-year-old White Male

Diagnosis: HPV+ Oropharyngeal Squamous Cell Carcinoma (base of tongue)

Key Mutations: P16+

Treatment History: 3 prior lines

- **Line 1:** Concomitant cisplatin/XRT → 49 days, PD
- **Line 2:** Pembrolizumab → 84 days, PD
- **Line 3:** Paclitaxel/carboplatin → 184 days, PD

APR-1051 Activity

- **Current Status:** PD after 223 days of SD treatment
- **Best Response:** SD at first scan (-5% tumor response)

Notes: Stable disease maintained for 223 days.

Biomarker Defined Registration Path

Preliminary signals of monotherapy activity across cohorts in genomically defined tumors including uterine cancer, endometrial cancer, CRC and HNSCC

Clinical Activity by Indication and Biomarker

Indication	Biomarker / Altered Genes	Clinical Activity in ACESOT-1051*
Uterine/ Endometrial Cancer	PPP2R1A	2 PR
	CCNE1 overexpressed TP53	2 SD
CRC	KRAS & TP53	3 SD
	FBXW7	
HNSCC	HPV+	1 SD

Path to Registrational Cohort

- 1 Expand USC and CycE-PROC
- 2 Add additional cohorts
 - CRC FBXW7-mutated
 - HPV+ cancers
 - PPP2R1A mutated
- 3 Confirm durability, consistency of response and safety

- Responses across dose levels support a biomarker enriched expansion path
- Activity beyond endometrial cancer supports additional biomarker-defined cohorts



*Data as of May 6, 2026 and are preliminary from an ongoing dose-escalation study. Responses and stable disease require confirmation in additional patients and may change as follow-up matures. Safety and efficacy outcomes may vary by dose, schedule, and patient characteristics.

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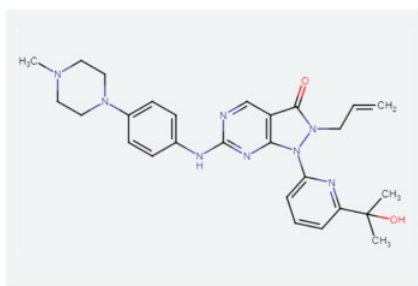
APR-1051:

Potentially Differentiated WEE1 Inhibitor
Pre-Clinical

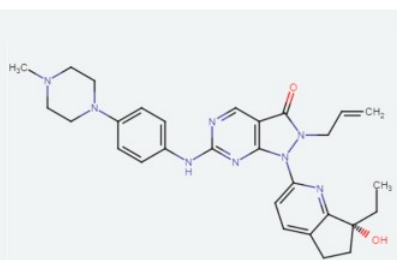
APR-1051: Potentially Best-in-Class WEE1 Inhibitor

Structurally differentiated: high potency, limited off-target inhibition design compared to other molecules

APR-1051 is based on a different molecular structure than AZD-1775 and ZN-c3 (not an analogue)



AstraZeneca
Adavosertib (AZD-1775)



Zentalis
Azenosertib (ZN-c3)



Apra
APR-1051



No head-to-head clinical studies have been conducted.

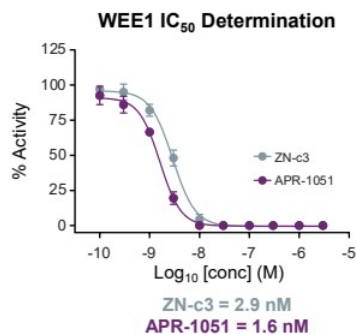
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APR-1051: Potentially Best-in-Class WEE1 Inhibitor

Potent inhibitor of WEE1

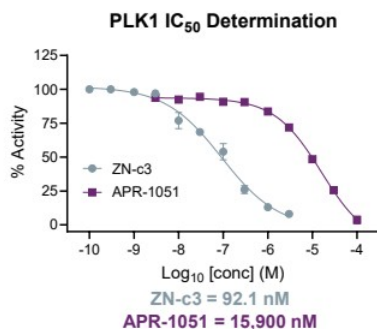
Does not substantially inhibit structurally and functionally related PLK1, PLK2 or PLK3

On-target WEE1 potency¹

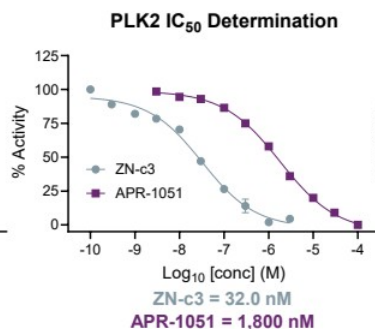


WEE1 Inhibition
IC₅₀ similar to ZN-c3

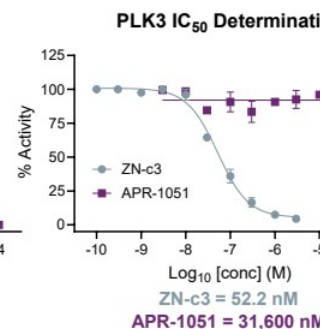
Important difference in off-target inhibition between APR-1051 and ZN-c3



PLK1 Inhibition
IC₅₀ >150-fold difference



PLK2 Inhibition
IC₅₀ >50-fold difference



PLK3 Inhibition
IC₅₀ >600-fold difference

APR-1051 specificity for WEE1 opens potential for greater therapeutic window

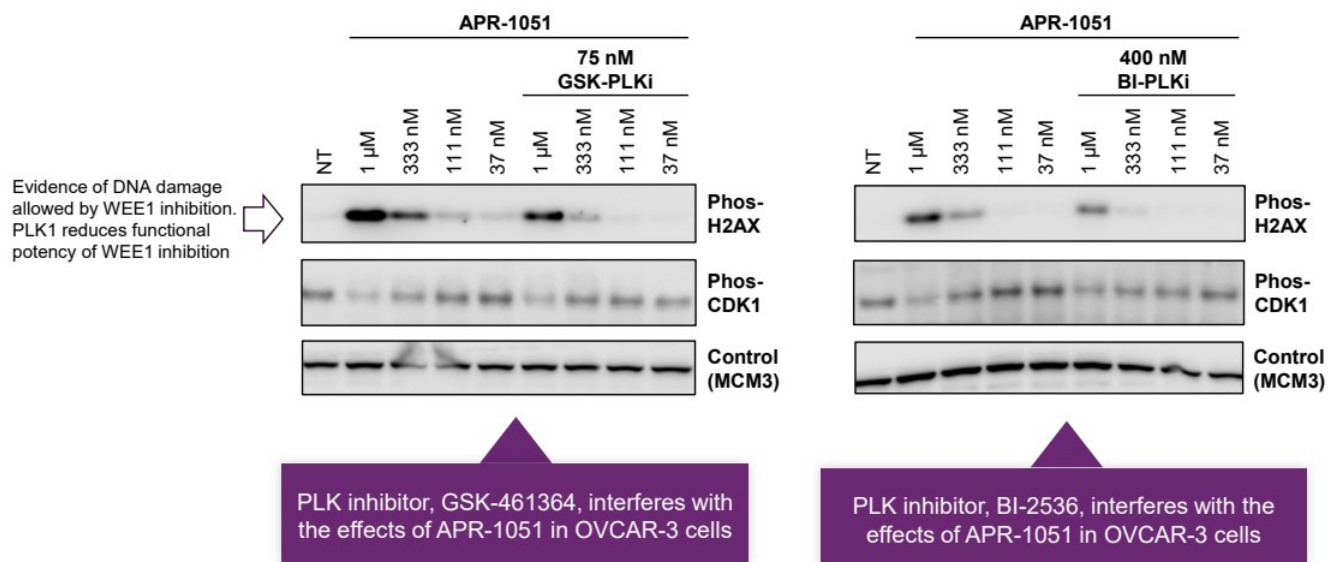


¹Data from exploratory in-vitro studies. No head-to-head clinical studies have been conducted.

AACR-NCI-EORTC Meeting, Poster B323, :
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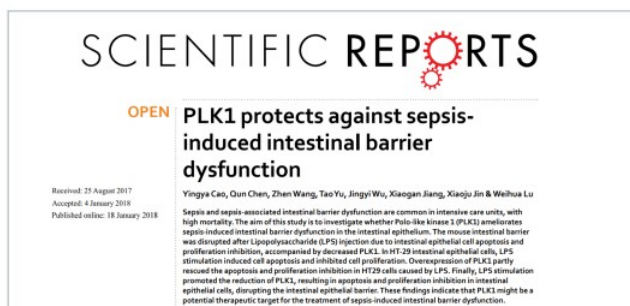
PLK1 Inhibition Counteracts Effect of WEE1 Inhibitors¹

Minimal PLK1 co-inhibition enhances therapeutic window for APR-1051



Inhibition of PLK1 reduces efficacy of WEE1 inhibition. Results in requiring higher doses of WEE1 inhibitors and introduces PLK1 related toxicity

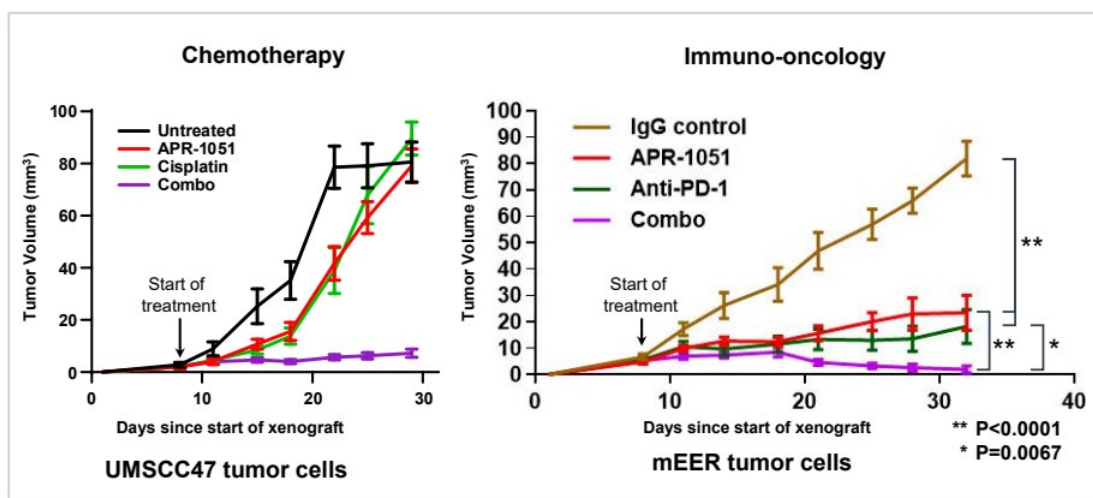
Studies Show PLK1 Suppression is Associated with Sepsis-Induced Loss of Intestinal Barrier Function



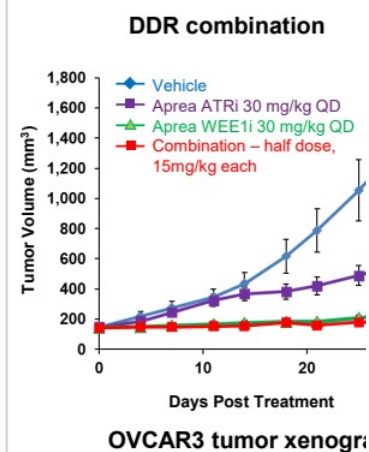
- 1 PLK1 protects against sepsis-induced intestinal barrier dysfunction, Cao *et al*, *Scientific Reports* (2018).
- 2 PLK1 protects intestinal barrier function in sepsis: A translational research, Cao *et al*, *Cytokine* (2023).
- 3 PLK1 protects intestinal barrier function in sepsis: A translational research, Cao *et al*, *Molecular Medicine* (2022).
- 4 LncRNA DANCER improves the dysfunction of the intestinal barrier and alleviates epithelial injury by targeting the miR-1306-5p/PLK1 axis in sepsis, Wang *et al*, *Cell Biology International* (2021).

APR-1051 Showed Preclinical Activity in Combination with Chemo, IO and ATRi Across Multiple Cancer Models

APR-1051 suggests synergistic potential preclinically with standard oncology agents



HPV+ Cancer – Collaboration with MD Anderson



Ovarian Cancer*



* Data on file.

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APR-1051 WEE1 Summary

A next-generation WEE1i with the potential to deliver meaningful clinical benefit and durable value

1

Clinically validated target

- WEE1 inhibitors have shown promising activity in genomically defined tumors
- Competitor programs constrained by low therapeutic window

2

APR-1051 clinical highlights

- Preliminary clinical proof-of-concept at 150 mg and 220 mg dose levels with two partial responses and six patients with stable disease presented at ASCO 2026
- Potentially favorable safety profile at active dose levels
- Enrollment accelerating with expansion from three to ten active clinical sites
- **Interim clinical data to be presented Q4 2026**

3

APR-1051 preclinical differentiation and potentially best-in-class opportunity

- Novel structure with high potency, limited off-target inhibition design
- Minimal PLK1 inhibition enhances therapeutic window
- Potential for synergy demonstrated in combination with standard oncology agents



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Intellectual Property Portfolio
Financial Summary & Capitalization
Investment Highlights



Robust Global Intellectual Property Protection

Family 1	Ataxia Telangiectasia and Rad3-Related (ATR) Protein Kinase Inhibitors	• Macrocytic inhibitors of ATR & methods of using them to treat various cancers, filed on Oct. 13th, 2015 • Patents granted in AU, BR, CA, CN, EP, IL, IN, JP, KR, MX, HK. • 1.1: Issued on May 30, 2017 as U.S. Patent 9,663,535; 1.2: Issued on May 29, 2018 as U.S. Patent 9,981,989; 1.3: Issued on Feb. 5, 2019 as U.S. Patent 10,196,405
Family 2	ATR Inhibitor and Methods of Use	• Carboxylic acid-containing macrocyclic ATR inhibitors, and prodrugs; methods of using these inhibitors to treat various cancers; filed on Apr. 12, 2017 • Issued on May 28th, 2019 as U.S. Patent 10,301,324
Family 3	ATR Inhibitor Pharmaceutical Composition and Methods	• International application filed on Apr. 14, 2023 • Pharmaceutical formulation and composition of our lead ATR inhibitor in the clinic • Patent granted in JP and EA; Applications pending US, AU, BR, CA, CN, EP, HK, IL, IN, KR, MX, NZ, PH, SG, ZA
Family 4	WEE1 Inhibitor Pharmaceutical Compositions and Methods	• International Application filed on Jun. 3, 2022 • Composition of our lead WEE1 inhibitor compounds • Patent granted in AU; Applications pending in US, AU, BR, CA, CN, EP, HK, IL, IN, JP, KR, MX, ZA
Family 5	Methods of Treating Cancer	• Clinical methods of treating advanced solid cancer tumors using lead ATR inhibitor • International application filed on Sept. 19, 2025
Family 6	Macrocytic DYRK1A-B Inhibitors and Methods of their Preparation and Use	• International application filed on Jan. 22, 2026 • Macrocytic inhibitors of DYRK1A-B and methods of using them to treat disease
Family 7	Macrocytic DDR Target Inhibitors and Methods of their Preparation and Use	• International application filed on Jan. 25, 2026
Family 8	Methods of Treating Cancer	• Provisional application filed on Jan. 9, 2026 • Clinical methods of treating cancer using lead Wee1 inhibitor

Aprea Therapeutics (NASDAQ: APRE)

Financial Summary and Capitalization

Cash and Equivalents of ~\$41.2M as of June 30, 2026

\$30.0M in gross proceeds raised in private placement that closed on March 31, 2026

Securities	Common Equivalents as of August 12, 2026
Preferred Stock (as converted)	15,596
Common Stock ⁽¹⁾	12,591,136
Warrants ⁽²⁾	89,438,517
Options	1,137,333
Restricted Stock Units	65,438
Fully Diluted Equivalents	103,248,020



1. 400,000,000 common shares authorized
2. Total warrants include pre-funded, Tranche A, Tranche B and Purchase

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Investment Highlights



Technology developed by pioneers in synthetic lethality

- Management with strong drug development and commercial expertise
- Focused on addressing unmet needs for patients with biomarker defined cancers



Highly potent and selective design, potential best in class inhibitors

- Diversified portfolio including WEE1 (APR-1051) and ATR (ATRN-119) inhibitors
- Preliminary evidence of clinical activity including PRs with APR-1051
- Single agent and combination potential therapies



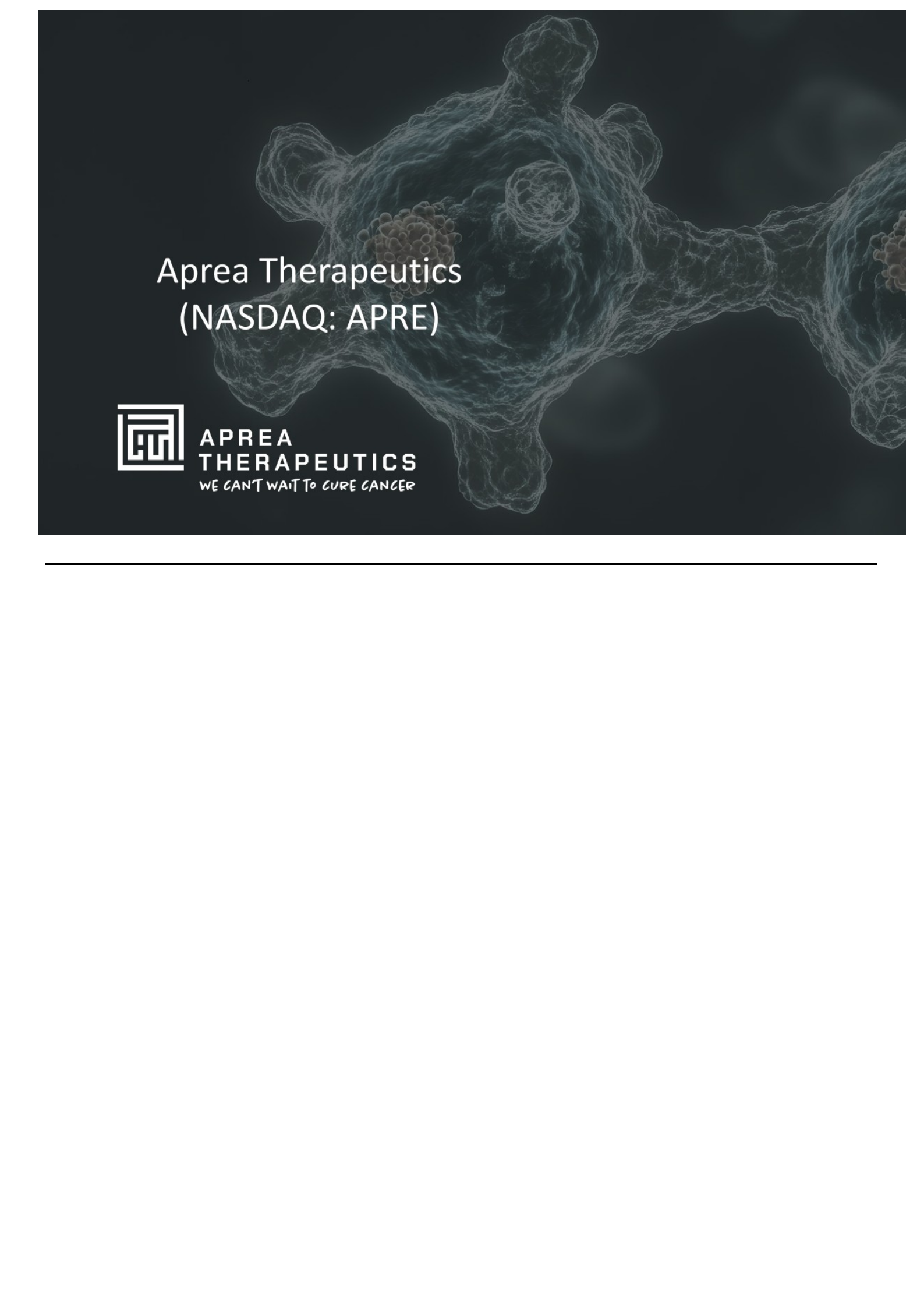
Near term catalysts

- APR-1051: Q4 2026 Interim clinical data; Q2 2027 Complete dose escalation
- ATRN-119: October 2025 RP2D ✓ H2 2026 Potential collaborations on combinations



Expected cash runway into Q1 2028

- Achieve near term inflection points and catalysts
- Evaluate optimal strategic partnerships

A detailed, high-magnification microscopic image of a cell, likely a cancer cell, showing a complex, irregular shape with a prominent, glowing nucleus. The cell is rendered in a semi-transparent, wireframe-like style, revealing internal structures. The background is dark, making the cell stand out.

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