UNITED STATES SECURITIES AND EXCHANGE COMMISSION

WASHINGTON, D.C. 20549

FORM 8-K

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

Date of Report (Date of earliest event reported): June 12, 2024

DAY ONE BIOPHARMACEUTICALS, INC. (Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation)	001-40431 (Commission File Number)	83-2415215 (IRS Employer Identification No.)
2000 Sierra Point Parkway, Suite 501 Brisbane, California (Address of principal executive offices)		94005 (Zip Code)
Registrant's tele	phone number, including area code: (650)	484-0899
(Former	N/A name or former address, if changed since last report)	
eck the appropriate box below if the Form 8-K filing is owing provisions:	intended to simultaneously satisfy the filing	obligation of the registrant under any of the
Written communications pursuant to Rule 425 unde	r the Securities Act (17 CFR 230.425)	
Soliciting material pursuant to Rule 14a-12 under th	ne Exchange Act (17 CFR 240.14a-12)	
Pre-commencement communications pursuant to Ru	ule 14d-2(b) under the Exchange Act (17 CFF	R 240.14d-2(b))
Pre-commencement communications pursuant to Ru	ale 13e-4(c) under the Exchange Act (17 CFR	240.13e-4(c))
Securities r	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 \$0.0001 per share	DAWN	Nasdaq Global Select Market
icate by check mark whether the registrant is an emerg pter) or Rule 12b-2 of the Securities Exchange Act of		of the Securities Act of 1933 (§ 230.405 of this
		Emerging growth company \square

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any

new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. \Box

On June 12, 2024, Day One Biopharmaceuticals, Inc. (the "Company") updated its corporate presentation in connection with recent updates to the Company's pivotal Phase 3 trial ("FIREFLY-2") evaluating tovorafenib as a front-line therapy in patients aged 6 months to 25 years with pediatric low-grade glioma ("pLGG").

A copy of the updated presentation is attached as Exhibit 99.1 to this Current Report on Form 8-K.

The information furnished in this Item 7.01, including Exhibit 99.1 to this Current Report on Form 8-K, shall not be deemed to be "filed" for 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 or Sections 11 and 12(a)(2) of the Securities Act of 1933, as amended (the "Securities Act"). The information contained in this Current Report on Form 8-K and in the accompanying Exhibit 99.1 shall not be incorporated by reference into any other filing under the Exchange Act or under the Securities Act, except as shall be expressly set forth by specific reference in such filing.

Item 8.01. Other Events.

On June 12, 2024, the Company announced the following changes with respect to its FIREFLY-2 trial:

- The primary endpoint of objective response rate will be assessed according to the Response Assessment in Pediatric Neuro-Oncology Low-Grade Glioma ("RAPNO-LGG") criteria
- Key secondary endpoints of progression free survival and duration of response will be assessed according to RAPNO-LGG criteria
- New patients will be initiated on a starting dose of $380 \text{ mg/m}^2/\text{dose}$ once weekly
- Addition of a once-monthly carboplatin regime as a fourth standard of care option for arm 2

 $Those \ updates \ to \ the tovorafenib \ program \ were \ made \ based \ on \ feedback \ from \ the \ U.S. \ Food \ and \ Drug \ Administration \ during \ review \ of \ the \ New \ Drug$ Application forming the basis of the approval of OJEMDATM (tovorafenib) for the treatment of pediatric patients 6 months of age and older with relapsed or refractory pLGG harboring a BRAF fusion or rearrangement, or BRAF V600 mutation.

Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No.

Description

99.1 Corporate Presentation.

104 Cover Page Interactive Data File (embedded within the Inline XBRL document) SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, as amended, the registrant has duly caused this report to be signed on its behalf by the undersigned thereunto duly authorized.

DAY ONE BIOPHARMACEUTICALS, INC.

Date: June 12, 2024

By: /s/ Charles N. York II, M.B.A.
Charles N. York II, M.B.A.
Chief Operating Officer and Chief Financial Officer



Disclaimer

This presentation and the accompanying oral commentary contain forward-looking statements that are based on our management's beliefs and assumptions and on information currently available to our management. Forward-looking statements are inherently subject to risks and uncertainties, some of which cannot be predicted or quantified. In some cases, you can identify forward-looking statements by terminology such as "may," "will," "should," "could," "expect," "plan," anticipate," "believe," "estimate," "predict," "intend," "potential," "would," "continue," "ongoing" or the negative of these terms or other comparable terminology. Forward-looking statements include all statements of the nan statements of historical fact contained in this presentation, including information concerning our future financial performance, including the sufficiency of our cash, cash equivalents and short-term investments to fund our operations, business plans and objectives, timing and success of our planned nonclinical and clinical development activities, the results of any of our strategic collaborations, including the potential achievement of milestones and provision of royalty payments thereunder, timing and results of nonclinical studies and clinical trials, efficacy and safety profiles of our products and product candidates, the ability of tovorafenib to treat pediatric low-grade glioma (pLGG) or related indications, the potential therapeutic benefits and economic value of our products and product candidates, potential growth opportunities, competitive position, industry environment and potential market opportunities, our ability to protect intellectual property and the impact of global business or macroeconomic conditions, including as a result of inflation, changing interest rates, cybersecurity incidents, potential instability in the global banking system, uncertainty with respect to the federal debt ceiling and budget and potential government shutdowns related thereto and global regional conflicts, on our business and operations.

Forward-looking statements are subject to known and unknown risks, uncertainties, assumptions and other factors. It is not possible for our management to predict all risks, nor can we assess the impact of all factors on our business or the extent to which any factor, or combination of factors, may cause actual results to differ materially from those contained in any forward-looking statements we may make. These factors, together with those that are described under the heading "Risk Factors" contained in our most recent Quarterly Report on Form 10-Q filed with the Securities and Exchange Commission ("SEC") and other documents we file from time to time with the SEC, may cause our actual results, performance or achievements to differ materially and adversely from those anticipated or implied by our forward-looking statements.

In addition, statements that "we believe" and similar statements reflect our beliefs and opinions on the relevant subject. These statements are based upon information available to us as of the date of this presentation, and although we believe such information forms a reasonable basis for such statements, such information may be limited or incomplete, and our statements should not be read to indicate that we have conducted a thorough inquiry into, or review of, all potentially available relevant information. These statements are inherently uncertain and investors are cautioned not to unduly rely upon these statements. Furthermore, if our forward-looking statements prove to be inaccurate, the inaccuracy may be material. In light of the significant uncertainties in these forward-looking statements, you should not regard these statements as a representation or warranty by us or any other person that we wild achieve our objectives and plans in any specified time frame, or at all. We undertake no obligation to publicly update any forward-looking statements, whether as a result of new information, future events or otherwise, except as required by law.

This presentation also contains estimates and other statistical data made by independent parties and by us relating to market size and growth and other data about our industry. This data involves a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates. In addition, projections, assumptions and estimates of our future performance and the future performance of the markets in which we operate are necessarily subject to a high degree of uncertainty and risk.





Cancer Therapies for People of All Ages



Our Approach

- Develop medicines for genomically-defined cancers
- Establish first-in-class position through rapid registration pathways
- Expand to adolescent and adult populations in parallel and pursue those opportunities with the same commitment we do for children





Nasdaq: DAWN IPO: 2021 Founded: 2018 Financial Position: Runway into 2026



Our Pipeline

Product Candidate	Therapeutic Area	Preclinical	Phase 1	Phase 2	Phase 3/ Registrational	Approved	Recent & Anticipated Milestones
Tovorafenib Type II RAF Inhibitor	BRAF-altered Relapsed pLGG	FIREFLY-1 (pivo	tal Phase 2)			(*) ojemda* (tovorafenio)	FDA approval: April 2024
OJEMDA brand name in U.S. ¹	Frontline RAF- altered pLGG	FIREFLY-2 (pivo	tal Phase 3)				First patient dosed: March 2023
Pimasertib MEK 1/2 Inhibitor	MAPK-altered solid tumors [†] (Combo w/ tovorafenib)	FIREFLIGHT-1†	t				Recommended Phase 2 dose & schedule expected: 2H 2024
VRK1 Program VRK1 Inhibitor	Pediatric and adult cancers						In-licensed [§] : August 2023



¹OJEMDA has received accelerated approval by the U.S. Food and Drug Administration. † Pimasertib Phase 1 dose escalation and expansion trial previously completed. † † Includes patients ≥12 years of age. ⁵ Research collaboration and license agreement with Sprint Bioscience AB for exclusive worldwide rights to a research-stage program targeting VRK1. pLGG, pediatric low-grade glioma. The safety and efficacy of investigational agents and/or investigational uses of approved products have not been established.





OJEMDA Now Approved In The U.S.



OJEMDA is the **first and only FDA Approved therapy** for the treatment of pediatric patients 6 months of age and older with relapsed or refractory pediatric low-grade glioma harboring a BRAF fusion or rearrangement, or BRAF V600 mutation



Day One

pLGG Impact On Patients' Lives

Lily was diagnosed with an operable brain tumor at 5 months of age













Pediatric Low-Grade Glioma: The Most Common Type Of Brain Tumor In Children

pLGGs are chronic and relentless, with patients suffering profound tumor and treatment-associated morbidity that can impact their life trajectory over the long term¹

A Serious and Life-Threatening Disease

- For the majority of pLGG patients in the relapsed setting, there is no standard of care and no approved therapies
- Up to 75% of pLGGs have a BRAF alteration*, of those ~80% are BRAF fusions and ~20% are BRAF V600 mutations²⁻⁶
- Despite surgery playing a significant role in treatment, the vast majority of patients still require systemic therapy^{7,8}
- Due to high rate of disease recurrence, most patients will undergo multiple lines of systemic therapy over the course of their disease





*Incidence of BRAF atterations varies across pLGG subtypes. *Sievert AI, Fisher MJ. Pediatric low-grade gliomas. J Child Neurol. 2009;24(11):1397-1408. doi:10.1177/0883073809342005.*Penman CL et al. Front Onc 2015;554.*Cohen AR, N Engl J Med. 2020;368(20):1922-1931. *Lassaletta A, et al. J Clin Oncol. 2017;35(25):2934-2941.*Faulkner C, et al. J Neuropathol Exp Neurol. 2015;74(9):867-872. *Packer RJ, et al. Neuro Oncol. 2017;19(6):750-761.*Cost pm Cf et al. Neur

Conventional Treatments Can Be Disruptive To Childhood And Can Have Significant Long-Term Consequences

Surgery

- Significant recovery times
- Risks of complications
- Resection may be limited by location of tumor
- Potential for functional deficits based on location of tumor and extent of resection

Chemotherapy

- Requirement for indwelling catheter and weekly infusions
- Risk of neutropenia, hypersensitivity reactions, nausea and vomiting and peripheral neuropathy

Radiation

- Risk of secondary malignancy
- Risk of malignant transformation
- Risk of vascular proliferation and stroke
- Neurocognitive impact, depending on location of tumor and radiation field

Goal of therapy is to control the tumor, minimize the burden of surgery, chemotherapy, and radiation, and reduce the risk of life-long treatment and disease-related effects



Fourter 1. Hettzer AM, Raghtubar K, Ris MD, et al. Neuropsychological functioning following surgery for podiatric low-grade gillomar, a prospective longificational study. J Neuropsy Podiatri. 2019;1-9. doi:10.3717/2019.9.PCD519397. 2. Sypart R. Hanaging side effects of childhood cancer concurrence, treatment and risks of second primary malignancies. Cancerage (Basel). 2021;113;207. 6. doi:10.3717/2019.9. doi:10.3717/2019.9. doi:10.3717/2019.0. A lattice of the concurrence (Basel). 2021;113;207. 6. doi:10.3717/2019.9. doi



Overview U.S. Prescribing Information For OJEMDA™ (tovorafenib)

Available in tablet formulation and pediatric-friendly powder for oral suspension

INDICATION

OJEMDA is indicated for the treatment of pediatric patients 6 months of age and older with relapsed or refractory pediatric low-grade glioma harboring a BRAF fusion or rearrangement, or BRAF V600 mutation

RECOMMENDED DOSE

380 mg/m² administered orally once weekly (not to exceed a dose of 600mg once weekly); OJEMDA can be taken with or without food



For full prescribing information, visit dayonebio.com



'This indication is approved under accelerated approval based on response rate and duration of response. Continued approval for this indication may be contingent upon verification of clinical benefit in a confirmatory trial.

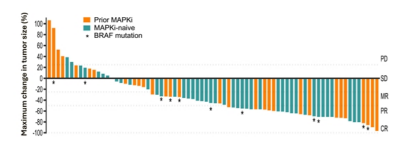


Efficacy Summary From OJEMDA™ (tovorafenib) Prescribing Information



51%

Overall response rate (RAPNO-LGG) in 76 evaluable patients



	RAPNO-LGG				
Response (IRC)		n (%)	95% CI		
ORR, n (%)	76	39 (51)	40-63		
BRAF fusion or rearrangement	64	33 (52)	39-64		
BRAF V600 mutation	12	6 (50)	21-79		
Prior MAPKi use	45	22 (49)	31-64		
MAPKi-naïve	31	17 (55)	36-73		
Median DOR, months	39	13.8	11.3-NR [†]		
Median TTR, months	39	5.3			
Range		1.6-11.2			



June 5, 2023 data cutoff. CI, confidence interval; DOR, duration of response; IRC, independent radiology review committee; LGG, tow-grade glioma; NR, not reached; ORR, overall response rate; RAPNO, Response Assessment in Pediatric Neuro-Oncology; TTR, time to response; CR, complete response; PR, partial response; MR, minor response; SD, stable disease; PD, progressive disease. 1 As of the data cutoff, 68% remain on tovorafenib.



Safety Summary From OJEMDA™ (tovorafenib) Prescribing Information



Warnings and Precautions

- Hemorrhage
- · Skin toxicity, including photosensitivity
- Hepatotoxicity
- · Effect on growth
- · Embryo-fetal toxicity
- Use in NF1- associated tumors

No boxed warnings or contraindications

	TEAEs (≥ 30% of patients [n=137])				
Preferred Term, n (%)	Any Grade	Grade ≥3			
Any AE	137 (100)	86 (63)			
Hair color changes	104 (76)	0			
Anemia	81 (59)	15 (11)			
Elevated CPK	80 (58)	16 (12)			
Fatigue	76 (55)	6 (4)			
Vomiting	68 (50)	6 (4)			
Hypophosphatemia	64 (47)	0			
Headache	61 (45)	2 (1)			
Maculo-papular rash	60 (44)	11 (8)			
Pyrexia	53 (39)	5 (4)			
Dry skin	49 (36)	0			
Elevated LDH	48 (35)	0			
Increased AST	47 (34)	4 (3)			
Constipation	45 (33)	0			
Nausea	45 (33)	0			
Upper RTI	43 (31)	2 (1)			
Dermatitis acneiform	42 (31)	1 (1)			
Epistaxis	42 (31)	1 (1)			







ett F, van Tilburg CM, Bison B, et al. Response to trametinib treatment in progressive pediatric low-grade glioma patients. I Neurooncol. 2020;149(3):499-510. doi:10.1007/s11060-020-03640-3. ² Ryall S, Tabori U, Hawkins C. Pediatric low-grade more and the earl of molecular diagnostics. Acta Neuropathol Commun. 2020;81(1):30. doi:10.1186/sa04078-020-00902-2. ² SEER US complete prevalence counts of patients aged under 25 with lemin and Other Nervous Systems tumors as of January 1, 10-68TRUS, Qaddoumi et al (2005, Schreck et al 2019, ClearView Analysis. ² US Gensus. Estimated annual incidence, estimated prevalence, and estimated treverent/progressive total addressable patient population are Boy Onc esclusions base publicly available data. ² Source: Internal market research conducted by EpidSrategies, A Division of ToxStrategies, Inc. on behalf of Day Onc. ² Ryall S, et al. Acta Neuropathol Commun. 2020;8(1):30. ² Behalfer, F, et al. Cancers (Basel). 1911(6):748. ² Pornama CL, et al. Front Oncol. 2015;55:54. ³ Packer RJ, et al. Neuropathol Co. 2017;19(6):75-75. ⁵ Choen Art, et al. N. Frag I Med. 2023;238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-238(2):1922-23



What Physicians & Caregivers Are Looking For In A Therapy

What HCP's are Seeking

Effective in stopping or shrinking tumors
Manageable safety profile
Minimal disruption to child's life



"The goal is not interfering with the child's life." – Ped Onc, Chicago Ad Board

What Caregivers are Seeking

Live as normal of a childhood as possible Minimal impact from the disease Minimal disruption to child's life



"Our time with our kids is precious and not guaranteed, so the less time with meds and doctors the better."

– Caregiver for a child under 5 yrs





Product Profile Aligns With What Physicians Are Looking For In A Therapy



Efficacy	Meaningful tumor stabilization or shrinkage may be possible with OJEMDA. In the clinical trial: 51% of children experienced tumor shrinkage by at least 25% 82% of children saw their tumors shrink or remain stable
Safety	Generally well-tolerated therapy, with 9 out of 10 patients staying on treatment in the clinical trial Most common grade 3 / 4 adverse events include: anemia, elevated CPK, maculo-papular rash, fatigue & vomiting
Dosing	Once-weekly, taken with or without food conveniently from home can mean fewer daily interruptions

OJEMDA is indicated for the treatment of patients 6 months of age and older with relapsed or refractory pediatric low-grade glioma (LGG) harboring a BRAF fusion, rearrangement, or BRAF V600 mutation.



Comprehensive Approach For A Successful Launch



Physicians

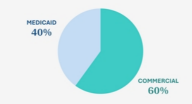
Objective: Establish OJEMDA[™] as 1st choice in relapsed / refractory *BRAF*-altered pLGG patients



 Dedicated & experienced sales team to engage HCPs

Payers

Objective: Rapidly establish coverage



- Pre-launch engagement to establish Day
 One & provide background information
- Plans in place for rapid engagement postapproval

Patients & Families

Objective: Provide a positive & supportive experience when initiating therapy





- SP distribution enables consistent patient experience
- Comprehensive patient support programs address patient needs and accelerates access to drug





Targeted Launch With Highly Experienced Field Team



Targeting ~200 centers where 90% of pLGG patients receive treatment

Deep oncology experience with relationships at top-tier accounts



18 Account Managers fully-dedicated to OJEMDA

Average experience:

- 13 years of oncology
- 4 years of rare disease
- **2** years of pediatric oncology clinical experience

Institutional experience and existing relationships with key accounts





Patient Support Program Supporting Access

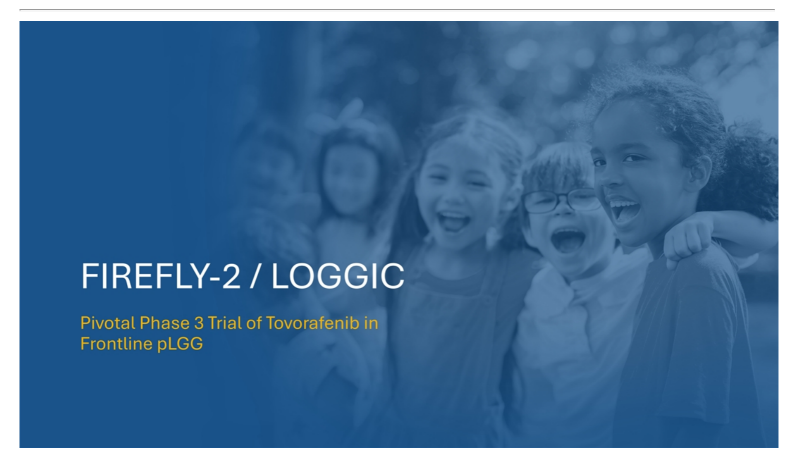












FIREFLY-2/LOGGIC Pivotal Phase 3 Trial Of Tovorafenib In Frontline pLGG



Trial Design

- Randomized, global, registrational Phase 3 trial of monotherapy tovorafenib vs SoC chemotherapy
- Eligibility: Patients aged 6 months to <25 years with LGG harboring a RAF alteration and requiring first-line systemic therapy
- Tovorafenib available as tablets and pediatric-friendly liquid suspension
- Patients who progress after stopping tovorafenib may be re-challenged
- Patients who progress in the SoC arm during or post-treatment may crossover to receive tovorafenib

Endpoints

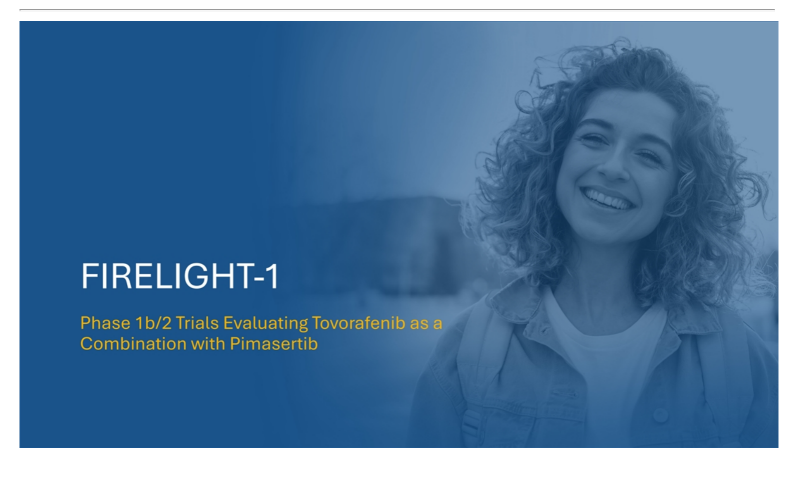
- Primary endpoint: ORR based on RAPNO-LGG criteria, assessed by blinded independent central review
 - The ORR primary analysis is expected to occur ~12 months after the last patient randomized
- Key secondary endpoints: PFS and DoR by RAPNO-LGG criteria
- Other secondary endpoints: changes in neurological and visual function, safety, and tolerability
- Key exploratory objectives: QoL and health utilization measures





* COG or SIOPe-LGG regimen. Abbreviations: CHG, chiasmatic, hypothalamic glioma; DoR, duration of response; LGG, tow-grade glioma; ORR, objective response rate; QoL, quality of life; QW, once weekly; Society of the control of the c





Pimasertib: Investigational Allosteric MEK1/2 Inhibitor With Demonstrated Activity In MAPK-Driven Solid Tumors

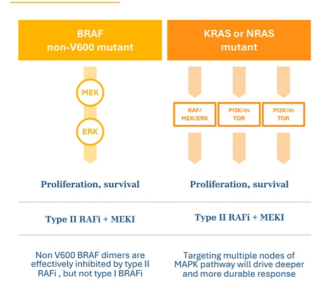
- Pimasertib is an investigational orally-bioavailable, selective, noncompetitive MEK1/2 inhibitor in-licensed from Merck KGaA in February 2021
- Extensive non-clinical and clinical development work through Phase 2, including a solid tumor trial in Japan and combinations with other MOAs
- Main AEs typical for all in-class allosteric MEK inhibitors (GI, CPK elevation, skin rash, visual disturbances)
- Nearly three-fold higher CNS penetration than other MEKi inhibitors (trametinib or selumetinib)
- Pimasertib showed monotherapy clinical activity, including an improvement in median PFS versus dacarbazine in NRAS mutant melanoma
- Combination with tovorafenib and other targeted therapies may unlock the full value of pimasertib in advanced solid tumors

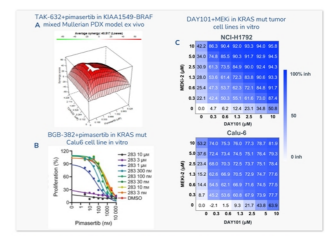






Vertical MAPK Pathway Inhibition With Tovorafenib And Pimasertib May Unlock Potential Synergy For Adult Solid Tumors





- Type II RAFi + MEKi is synergistic in BRAF fusion melanoma PDX model ex vivo (internal data)
- Sensitivity of KRAS Q61 mutant cells to pimasertib is enhanced when cells are treated with the type II BRAF inhibitor BGB-283 (Yuan et al., Mol Onc 2020)
- Tovorafenib + MEK inhibitor is synergistic in KRAS G12C and Q61 mutant tumor cells (Venetsanakos et al., 2021 AACR poster presentation)





Tovorafenib / Pimasertib Combination In Solid Tumors (FIRELIGHT-1)

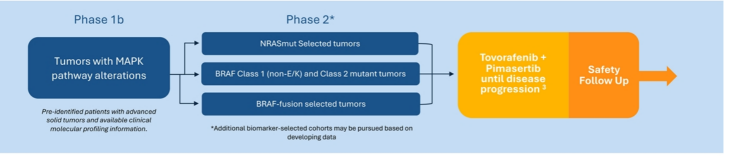


Trial Design¹

- Combination dose escalation, global phase 1b/2 trial²
- Phase 1b, BOIN (adaptive), n = 10/cohort (approximately)
- Phase 2, Simon 2-stage, n = 25/cohort (approximately)
- Eligibility: Patients aged 12 years and older, dose escalation will be performed in advanced solid tumor patients with any MAPK alteration. Expansion cohorts will focus on indications with a potential path to accelerated approval

Endpoints

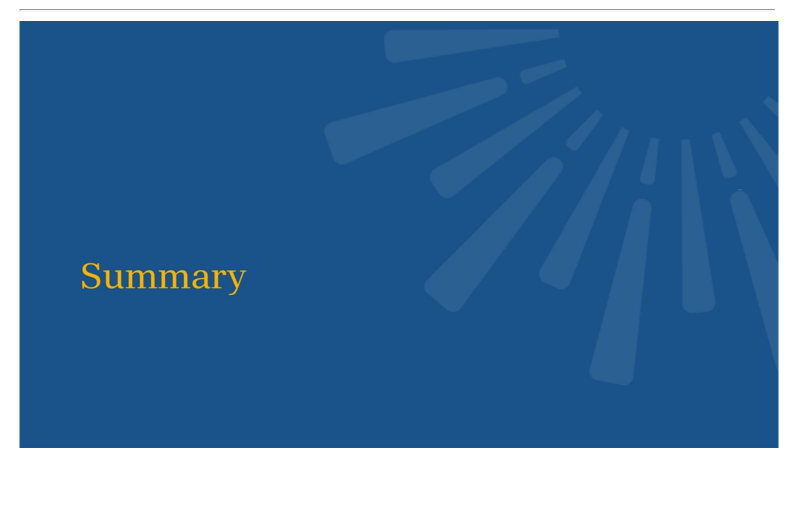
- Phase 1b: PK, PD and Safety, MTD/RP2D
- Phase 2: Efficacy (ORR, DOR)





Abbreviations: BOIN, Bayesian Optimal Interval Design; BRAF, B-Raf proto-oncogene, serine/threonine kinase; MAPK, mitogen-activated protein kinase; NRAS, neuroblastoma rat sarcoma viral oncogene. "Umbrella master study – DAY101-102 (main protocol) DAY101 and MAPK pathway aberration, Sub-study 1 monotherapy (DAY101-102a), Sub-study 2 MEK combo (DAY101-102b). "Intend to open U.S. and ex-U.S. clinical sties." 30/4/101 + Pimasertiu buttil disease progression, intolerable toxicity, withdrawal of consent, or death





Financial Summary: DAWN

Cash, cash equivalents and short-term investments as of March 31, 2024: \$317.9 million (no debt)

PRV sale in May 2024: \$108.0 million in gross proceeds

~87.4 million shares of common stock outstanding as of May 1, 2024

\$ Millions	Three Months Ended 3/31/24	Three Months Ended 3/31/23
R&D Expense	\$40.2	\$27.8
G&A Expense	\$26.6	\$18.0
Net Loss	\$62.4	\$42.4

Projected Cash Runway into 2026

FIREFLY-1: Pivotal Phase 2 clinical trial of tovorafenib

- Data published in Nature Medicine and oral presentations at SNO in November 2023
- OJEMDATM (tovorafenib) approved in the U.S. in April 2024
- Sale of PRV for \$108 million in gross proceeds in May 2024

FIREFLY-2/LOGGIC: Pivotal Phase 3 clinical trial of tovorafenib in newly diagnosed pLGG

First patient dosed in March 2023



All financial and share information is unaudited. PRV, Priority Review Voucher. As part of the PRV transaction, \$8.1 million of the total consideration received from the sale of the PRV pursuant to the PRV Transfer Agreement will be paid to Viracta Therapeutics, Inc. pursuant to the Company's License Agreement with Viracta, dated December 16, 2019, as amended.



Priorities as a Commercial-Stage Company

Launch OJEMDA[™] (tovorafenib)

- Expand awareness amongst physicians and establish broad coverage to enable patient access
- Establish OJEMDA as the standard of care for relapsed or refractory pLGG harboring a BRAF alteration
- Provide a positive and supportive experience when initiating OJEMDA therapy for patients and families

Advance Portfolio

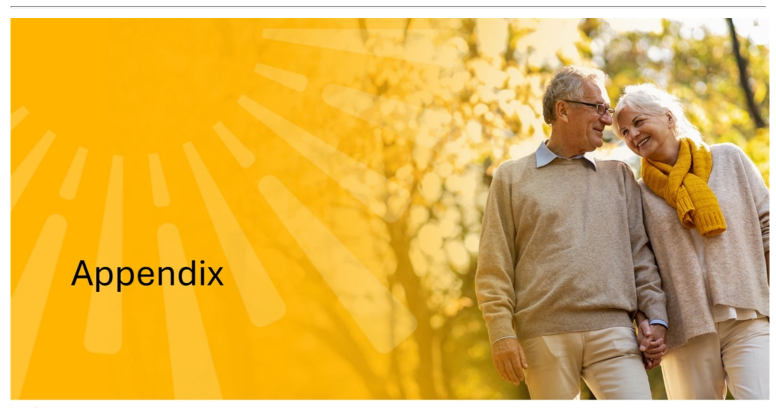
- FIREFLY-2: Study tovorafenib as a frontline therapy for treatmentnaive patients with pLGG
- FIRELIGHT-1: Evaluate tovorafenib in combination with pimasertib in adolescent and adult populations
- Advance early stage VRK1 program to clinical development

Expand Pipeline

- Grow Day One into a leading, biopharmaceutical company that is the partner of choice for oncology drug development
- Explore selective partnerships as a source of capital and risk sharing
- Further invest in business
 development activities to expand our
 multiple asset portfolio for both
 children and adults

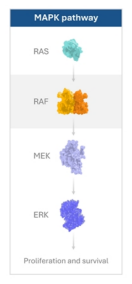


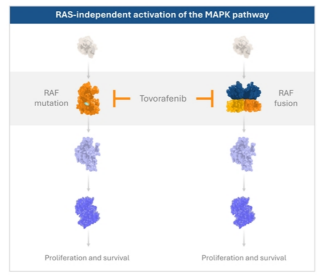






Tovorafenib Inhibits Both BRAF Fusions And BRAF V600 Mutations





Tovorafenib is an investigational, oral, selective, CNS-penetrant, type II RAF inhibitor that was designed to inhibit both monomeric and dimeric RAF kinase

- Activity in tumors driven by both RAF fusions and BRAF V600E mutations
- Tablet and pediatric-friendly liquid suspension
- Once weekly dosing

Currently approved type I BRAF inhibitors are indicated for use in patients with tumors bearing BRAF V600 mutations

 Type I BRAF inhibitors cause paradoxical MAPK activation in the setting of wild-type RAF, increasing the risk of tumor growth in BRAF fusion-driven



Source: 1. Sun Y et al., Neuro Oncol. 2017; 19: 774-85; 2. Sievart AJ et al., PNAS. 2013; 110:5957-62; 3. Karajannis MA et al., Neuro Oncol. 2014;16(10):1408-16.



Pivotal Phase 2 Trial Of Monotherapy Tovorafenib In Relapsed Or Progressive pLGG (FIREFLY-1)

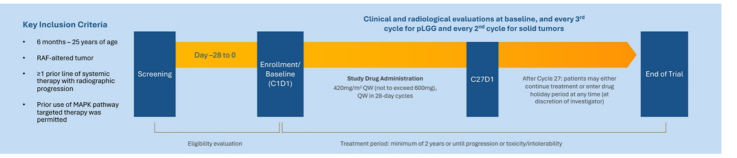


Trial Design

- Three arm, open-label, global registrational phase 2 trial
 - Pivotal Arm 1 (recurrent/progressive pLGG, n=77): harboring a KIAA1549-BRAF fusion or BRAF V600E mutation
 - Arm 2 (expanded access recurrent/progressive LGG, n=60): harboring an activating RAF alteration
- Arm 3 (extracranial solid tumors): harboring an activating RAF fusion

Endpoints (Pivotal Arm 1)

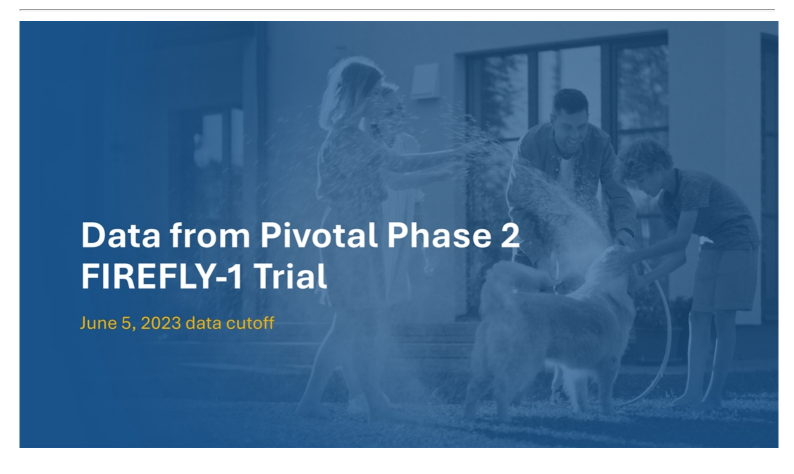
- Primary endpoint: ORR based on RANO-HGG¹, assessed by blinded independent central review
- Secondary endpoints: ORR by RAPNO-LGG² assessed by blinded independent central review; PFS, DoR; TTR, CBR; safety
- Exploratory analyses: ORR and CBR by RANO-LGG³ assessed by blinded independent central review





June 5, 2023 data cutoff. 1 Wen PY, et al. J Clin Oncol. 2010;28(11):1963-1972. Fangusaro J, et al. Lancet Oncol. 2020;21(6):e305-316. 3 van den Bent MJ, et al. Lancet Oncol. 2011;12(6):583-593. Abbreviations: CBR, clinical benefit rate; IRC, independent review committee; C, cycle; D, day; LGG, low-grade glioma; ORR, objective response rate; PFS, progression-free survival; DoR, duration of response; QW, once weekly; TIR, time to response; PANNO, Response Assessment in Neuron-Oncology; MAPNO, Response Assessment in Neuron-Oncology; MAPNO

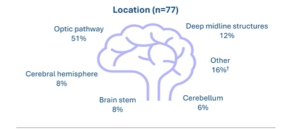


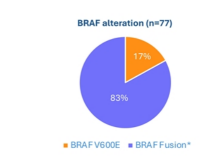


FIREFLY-1 Baseline Patient Characteristics



Characteristic	Arm 1 (n=77)
Median age, years (range)	8 (2-21)
Sex, n (%) Male Female	40 (52) 37 (48)
Race, n (%) White Asian Black Multiple Other Not specified	41 (53) 5 (6) 2 (3) 3 (4) 6 (8) 20 (26)
Number of lines of prior systemic therapy Median (range) 1, n (%) 2, n (%) ≥3, n (%)	3 (1-9) 17 (22) 21 (27) 39 (51)
Prior MAPK pathway targeted therapy, n (%) Prior MEK inhibitor Prior BRAF inhibitor Prior BRAF and MEK inhibitors [‡] Any MAPK inhibitor	43 (56) 8* (10) 5 (7) 46 (60)





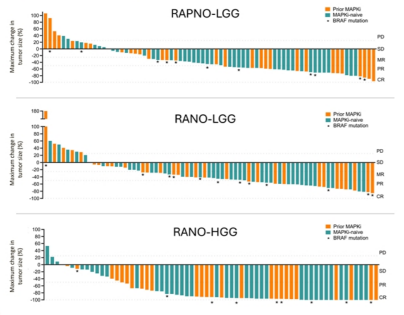


June 5, 2023 data cutoff. 'Includes 6 patients with BRAF duplication and 2 with BRAF rearrangement per fluorescence in situ hybridization or in situ hybridization. 'Includes tumors that were extending into multiple regions of the brain, leptomeningeal disease, and/or spinal disease. 'The 5 patients that had previously received both a MEK inhibitor and also a BRAF inhibitor are recorded in both the "Prior MEK inhibitor" and "Prior BRAF inhibitor" groups. MAPK, mitogen-activated protein kinase.



Tumor Response To Tovorafenib Using RAPNO-LGG, RANO-LGG and RANO-HGG





Response (IRC)	RAPNO-LGG n=76	RANO-LGG N=76	RANO-HGG N=69
ORR,* n (%)	39 (51)	40 (53)	46 (67)
95% CI	40-63	41-64	54-78
CBR,* n (%)			
SD of any length of time	62 (82)	63 (83)	64 (93)
SD ≥12 months	43 (57)	46 (61)	54 (78)
BOR,* n (%)			
CR	0	0	12 (17)
PR	28 (37)	20 (26)	34 (49)
MR	11 (14)	20 (26)	n/a
SD	23 (30)	23 (30)	18 (26)
SD <12 months	19 (25)	17 (22)	10 (14)
SD≥12 months	4 (5)	6 (8)	8 (12)
PD	13 (17)	11 (14)	4 (6)
NE	1 (1)	2 (3)	1 (1)
Median DOR, months	13.8	14.4	16.6
95% CI	11.3-NR	11.0-NR	11.6-NR
Median TTR, months	5.3	5.5	3.0
Range	1.6-11.2	1.6-11.3	2.6-16.6

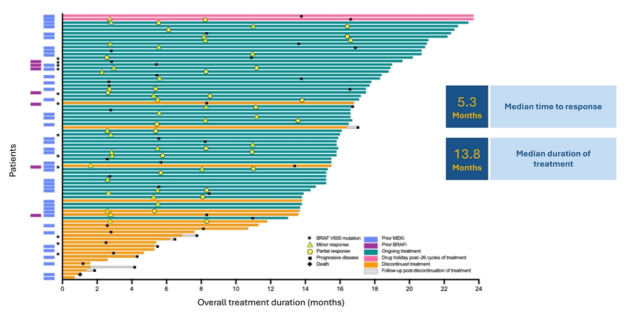


ne 5, 2023 data cutoff. BOR, best overalt response; CBR, clinical benefit rate; Cl. confidence interval; CR, complete response; DOR, duration of response; HGG, high-grade glioma; IRC, independent radiology view committee; LGG, low-grade glioma; MR, minor response; n/a, not applicable; NE, not evaluable; NR, not eached; ORR, overall response rate; PD, progressive disease; PR, partial response; RANO, Responsessment in Neuro-Oncology; RAPNO, Response Assessment in Pediatric Neuro-Oncology; SD, stable disease; TR, time to response. * ORR, CBR and BOR for RAPNO-LGG and RANO-LGG included MRs.



Duration Of Tovorafenib Therapy For All Patients With RAPNO-LGG Evaluable Lesions



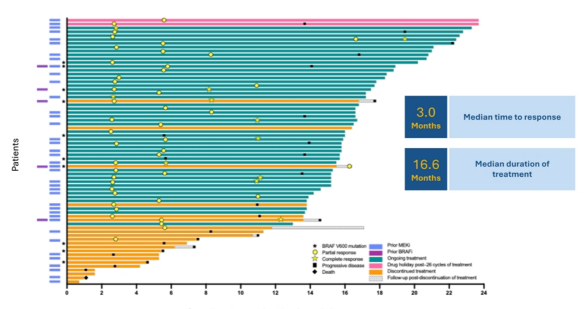






Duration Of Tovorafenib Therapy For All Patients With RANO-HGG Evaluable Lesions





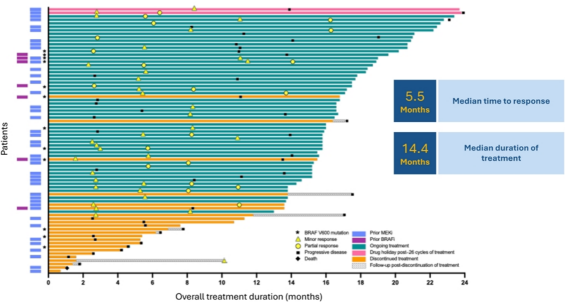
Overall treatment duration (months)





Duration Of Tovorafenib Therapy For All Patients With RANO-LGG Evaluable Lesions









Tumor Response To Tovorafenib Across Three Assessment Criteria Were Consistent Across BRAF Fusion And Mutation Patients, and Patients With Prior MAPK Treatment



	RAPNO-LGG ²			RANO-LGG ^{3,4}		RANO-HGG ¹	
Response (IRC)	n		n		n		
ORR,* n (%)	76	39 (51)	76	40 (53)	69	46 (67)	
BRAF fusion	64	33 (52)	64	33 (52)	59	41 (69)	
BRAF mutation	12	6 (50)	12	7 (58)	10	5 (50)	
Prior MAPKi	45	22 (49)	45	23 (51)	41	29 (71)	
MAPKi-naive	31	17 (55)	31	17 (55)	28	17 (61)	
CBR,* n (%) (SD of any length of time)	76	62 (82)	76	63 (83)	69	64 (93)	
BRAF fusion	64	53 (83)	64	53 (83)	59	55 (93)	
BRAF mutation	12	9 (75)	12	10 (83)	10	9 (90)	
Prior MAPKi	45	38 (84)	45	38 (84)	41	37 (90)	
MAPKi-naive	31	24 (77)	31	25 (81)	28	27 (96)	
CBR,* n (%) (SD ≥12 months)	76	43 (57)	76	46 (61)	69	54 (78)	
BRAF fusion	64	37 (58)	64	39 (61)	59	49 (83)	
BRAF mutation	12	6 (50)	12	7 (58)	10	5 (50)	
Prior MAPKi	45	25 (56)	45	26 (58)	41	33 (80)	
MAPKi-naive	31	18 (58)	31	20 (65)	28	21 (75)	
Median DOR, months (95% CI)**	39	13.8 (11.3-NR)	40	14.4 (11.0-NR)	46	16.6 (11.6-NR)	
BRAF fusion	33	13.8 (11.3-NR)	33	16.3 (11.0-NR)	41	16.8 (11.6-NR)	
BRAF mutation	6	NR (8.4-NR)	7	12.0 (8.4-NR)	5	15.1 (8.3-NR)	
Prior MAPKi	22	13.8 (11.3-NR)	23	12.0 (8.5-NR)	29	15.1 (9.0-16.8)	
MAPKi-naive	17	NR (8.4-NR)	17	16.3 (8.4-NR)	17	NR (11.6-NR)	



June 5, 2023 data cutoff. ¹ Fangusaro J, et al. Lancet Oncol. 2020;21(6):e305–316. ² Fangusaro J, et al. Lancet Oncol. 2020;21(6):e305–316. ³ van den Bent MJ, et al. Lancet Oncol. 2011;12(6):583–593. 4. Wen PY, et al. J. Clin Oncol. 2017;35(21),2439-2449. * ORR, CBR for RAPNO-LGG and RANO-LGG included MRs. ** the 95% CI were calculated using Kaplan-Meier method.

Tovorafenib Safety Data (n=137)



	TEAEs		TRAEs		
Preferred Term, n (%)	Any Grade	Grade ≥3	Any Grade	Grade ≥3	
Any AE	137 (100)	86 (63)	134 (98)	58 (42)	
Hair color changes	104 (76)	0	104 (76)	0	
Anemia	81 (59)	15 (11)	67 (49)	14 (10)	
Elevated CPK	80 (58)	16 (12)	77 (56)	16 (12)	
Fatigue	76 (55)	6 (4)	60 (44)	6 (4)	
Vomiting	68 (50)	6 (4)	28 (20)	3 (2)	
Hypophosphatemia	64 (47)	0	48 (35)	0	
Headache	61 (45)	2 (1)	29 (21)	0	
Maculo-papular rash	60 (44)	11 (8)	56 (41)	11 (8)	
Pyrexia	53 (39)	5 (4)	17 (12)	1 (1)	
Dry skin	49 (36)	0	45 (33)	0	
Elevated LDH	48 (35)	0	42 (31)	0	
Increased AST	47 (34)	4 (3)	41 (30)	4 (3)	
Constipation	45 (33)	0	31 (23)	0	
Nausea	45 (33)	0	25 (18)	0	
Upper RTI	43 (31)	2 (1)	2 (1)	0	
Dermatitis acneiform	42 (31)	1 (1)	41 (30)	1 (1)	
Epistaxis	42 (31)	1 (1)	27 (20)	0	
Decreased appetite	39 (28)	5 (4)	28 (20)	4 (3)	
Paronychia	36 (26)	2 (1)	32 (23)	2 (1)	
Pruritus	35 (26)	1 (1)	32 (23)	1 (1)	
COVID-19	34 (25)	0	0	0	

- The most common reasons for discontinuation were tumor hemorrhage (3 patients) and decrease in growth velocity (2 patients)
- 33 patients (24%) had TRAEs leading to dose reduction; 50 patients (37%) had TRAEs leading to dose interruption
- Median duration of dose interruption was 2 weeks
- 9 patients (7%) had TRAEs leading to discontinuation



June 5, 2023 data cutoff. Treatment-emergent AEs ≥25% any grade in arms 1 & 2. AE, adverse event; ALT, Alanine transaminase; AST, aspartate aminotransferase; COVID-19, Coronavirus disease 2019; CPK, creatine phosphokinase; LDH, lactate dehydrogenase; RTI, respiratory tract infection; TEAEs, treatment-emergent adverse events; TRAEs, treatment-related adverse events

