Black Diamond Therapeutics, Inc.

Developing MasterKey Therapies to Defeat Cancer Resistance

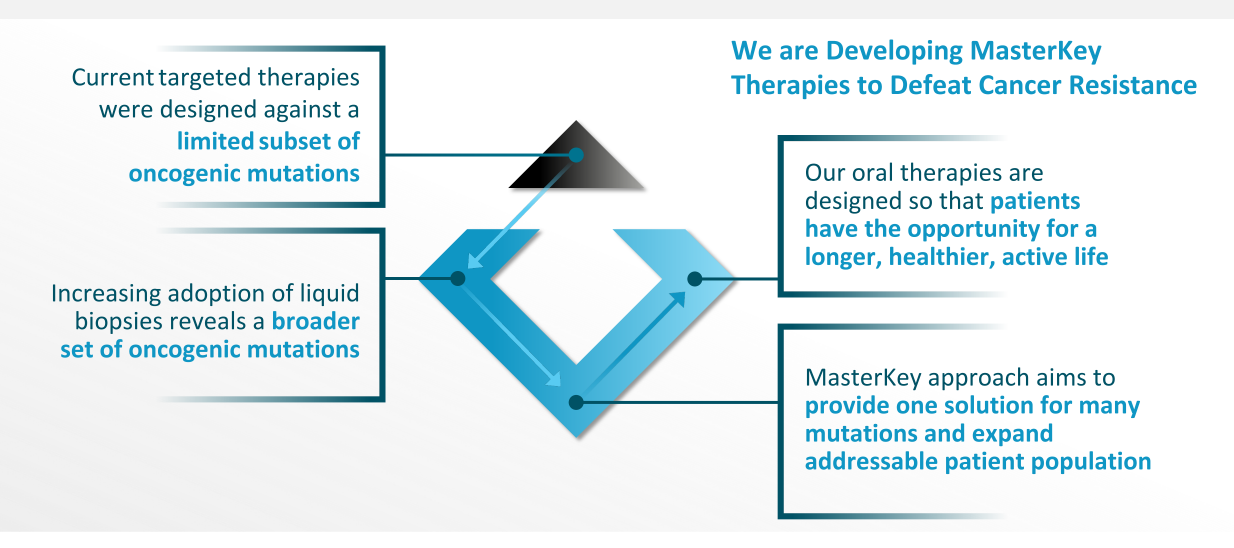


Forward-Looking Statements

Statements contained in this presentation regarding matters that are not historical facts are "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995. Because such statements are subject to risks and uncertainties, actual results may differ materially from those expressed or implied by such forward-looking statements. Such statements include, but are not limited to, statements regarding: the continued development and advancement of BDTX-1535 and BDTX-4933, including the ongoing clinical trials and the timing of clinical updates for BDTX-1535 in patients with NSCLC and in patients with recurrent GBM, and for Phase 1 clinical trial results for BDTX-4933, the potential of BDTX-1535 to benefit patients with NSCLC in earlier lines of therapy, potential future development plans for BDTX-1535 in NSCLC and GBM, including in first-line settings, and the Company's expected cash runway. Any forward-looking statements in this statement are based on management's current expectations of future events and are subject to a number of risks and uncertainties that could cause actual results to differ materially and adversely from those set forth in or implied by such forward-looking statements. Risks that contribute to the uncertain nature of the forward-looking statements include those risks and uncertainties set forth in its Annual Report on Form 10-K for the year ended December 31, 2023, filed with the United States Securities and Exchange Commission and in its subsequent filings filed with the United States Securities and Exchange Commission. All forward-looking statements contained in this presentation speak only as of the date on which they were made. The Company undertakes no obligation to update such statements to reflect events that occur or circumstances that exist after the date on which they were made.



Cancer is a Complex and Ever-Evolving Disease





Black Diamond Therapeutics At-a-Glance



Clinical-stage company
advancing MasterKey
therapies designed to
expand the addressable
patient population



Experienced team with deep understanding of cancer biology and oncology drug development



Pipeline of oral, brain penetrant drug candidates selectively targeting families of oncogenic mutations



Lead asset BDTX-1535 shows durable clinical responses in NSCLC, with additional opportunity in GBM



Multiple clinical catalysts across the pipeline in 2024

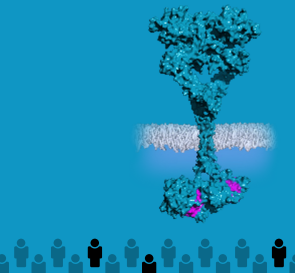


Strong balance sheet, with runway into Q2 2025; ended Q4 2023 with \$131.4M in cash



MasterKey: One Solution for Many Mutations

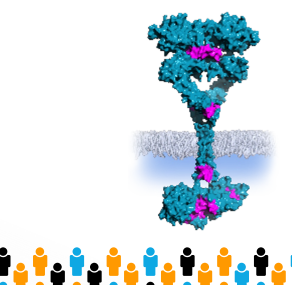
Traditional Approach: Targeting single mutations in individual tumor types



Limited addressable patient population

Black Diamond Approach:

Targeting families of oncogenic mutations



Expanded addressable patient population

Potent against broad mutation families (including drug resistance mutations)

Brain-penetrant to treat CNS disease

Selective targeting to deliver well-tolerated therapies

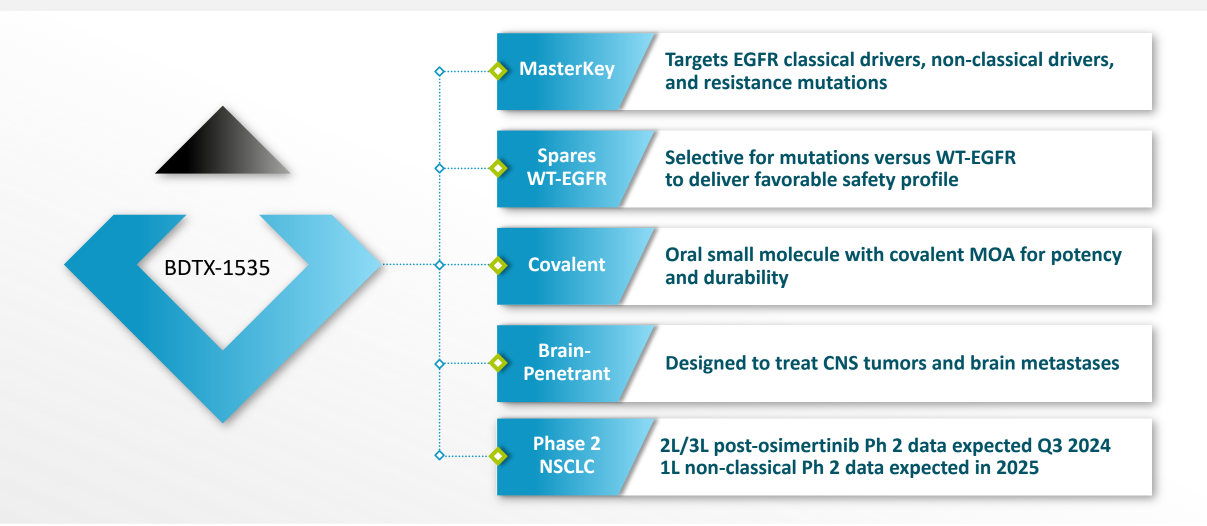


Advancing Wholly Owned Pipeline Across Multiple Oncology Indications

Target	Drug Candidate	Indication	Pre-clinical	Phase 1	Phase 2	Phase 3
EGFR	BDTX-1535	NSCLC	Phase 2 data in 2L/3L Phase 2 data in 1L pat	patients expected Q3 202 ients expected 2025	4	
		GBM	Phase 1 and "window data expected Q2 202			
RAF	BDTX-4933	KRAS mutant NSCLC	Phase 1 enrolling data expected Q4 2024			
		RAF/RAS mutant solid tumors		4		
FGFR2/3	BDTX-4876	Achondroplasia or solid tumors	Partnering candidate			
Undisclosed	Undisclosed	Multiple Solid tumors	Partnering candidate			



BDTX-1535: Most Advanced 4th Gen EGFR Inhibitor with Potential Best-In-Class Profile

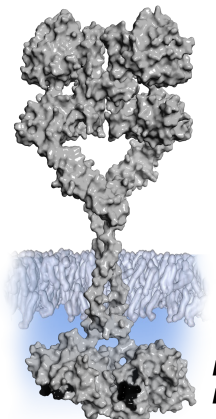




Real-World Data Describe a Broad EGFR Mutational Landscape in NSCLC & Reveal New Opportunities for EGFR Targeting

2004

Two classical EGFR oncogenic mutations first described¹



L858R & Ex19del (classical)

Today

Real-World Data reveals a broad landscape of classical & non-classical EGFR oncogenic mutations in NSCLC



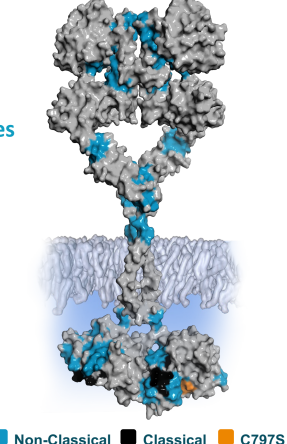


Black Diamond examined 235,761 sequenced NSCLC cases



- >100 EGFR mutations (classical & non-classical*)
- Distinct co-mutational profiles for L858R vs Ex19del
- Resistance mutations evolve with TKI therapies

*non-classical EGFR mutations also referred to as: atypical, second-site, uncommon, intrinsic, or PACC mutations

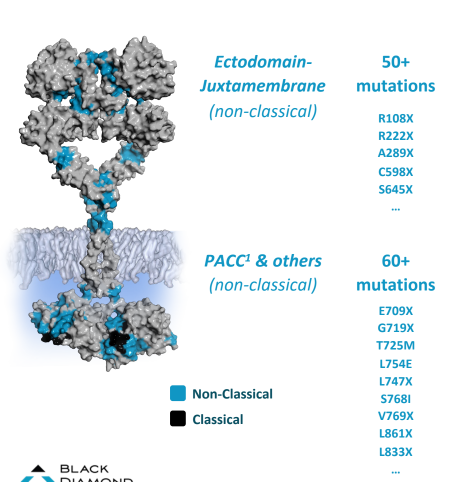




1. Paez et a. Science 2004 BDTX AACR 2024 oral presentation

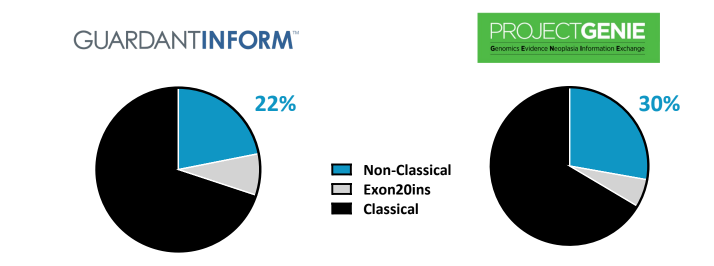
20-30% of Newly Diagnosed EGFRm NSCLC Patients Carry Non-Classical Mutations; Are Not Adequately Addressed by Current Therapies¹

Classical and non-classical driver mutations are distributed across EGFR structure



HERAPEUTICS

22-30% of newly diagnosed EGFRm NSCLC express non-classical mutations



Black Diamond Therapeutics analyses of 94,939 sequencing reports from <u>treatment naïve NSCLC</u>

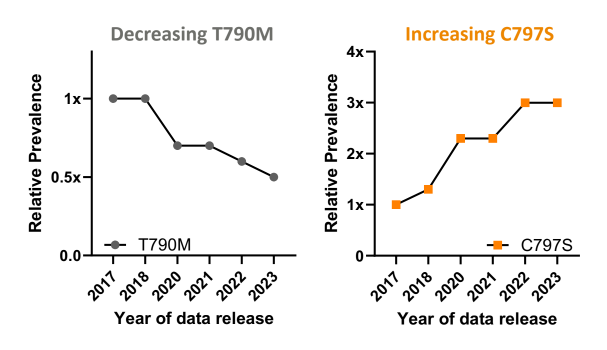
Adapted from Du et al. 2023. Analysis of the AACR GENIE database of 22,050 cases of NSCLC

Current therapies do not adequately address non-classical EGFR mutations¹

1. Borgeaud M. JTO 2024 BDTX AACR 2024 oral presentation

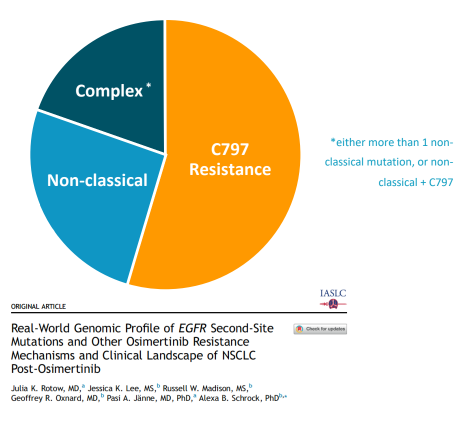
C797S and Non-Classical Mutations are Major Mechanisms of Acquired On-Target EGFR Resistance

On-target EGFR resistance mechanisms have shifted since approval of 3G TKI osimertinib in 1L setting



Black Diamond Therapeutics analyses of Foundation Medicine's FoundationInsights™ platform

C797S & non-classical mutations are major mechanisms of on-target EGFR resistance in patients post-osimertinib¹



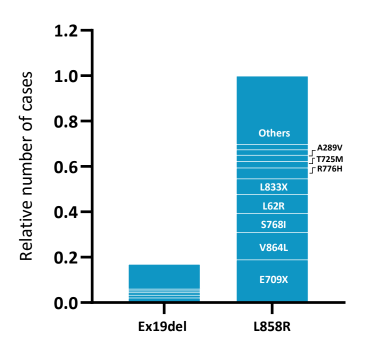
1. Adapted from Rotow, JK., et al., Journal of Thoracic Oncology, 2023. (non-classicals represented as L792, G796, G724, L718).



BDTX AACR 2024 oral presentation

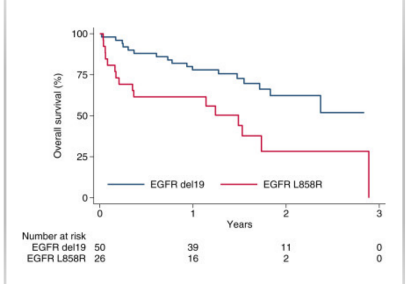
BDTX-1535 Potently Targets Non-Classical EGFR Mutations Commonly Co-Expressed with L858R Which are Insensitive to Osimertinib

EGFR-L858R tumors more frequently coexpress non-classical EGFR mutations before exposure to EGFR TKI



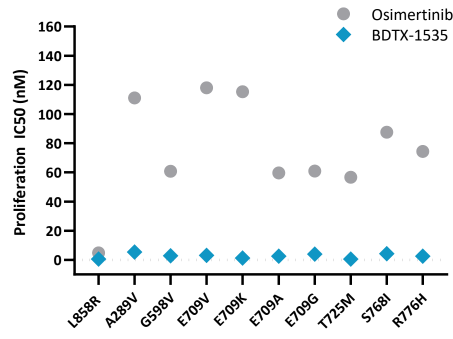
Black Diamond Therapeutics analyses of 94,939 sequencing reports from treatment naïve NSCLC (Guardant Health)

Patients with L858R do less well on osimertinib therapy vs Ex19del



Gijtenbeek et al 2023. Overall survival by type of mutation in patients with stage IV EGFR mutated NSCLC and brain metastasis who received first-line treatment with osimertinib

Non-classical mutations co-expressed with L858R are insensitive to osimertinib but sensitive to BDTX-1535



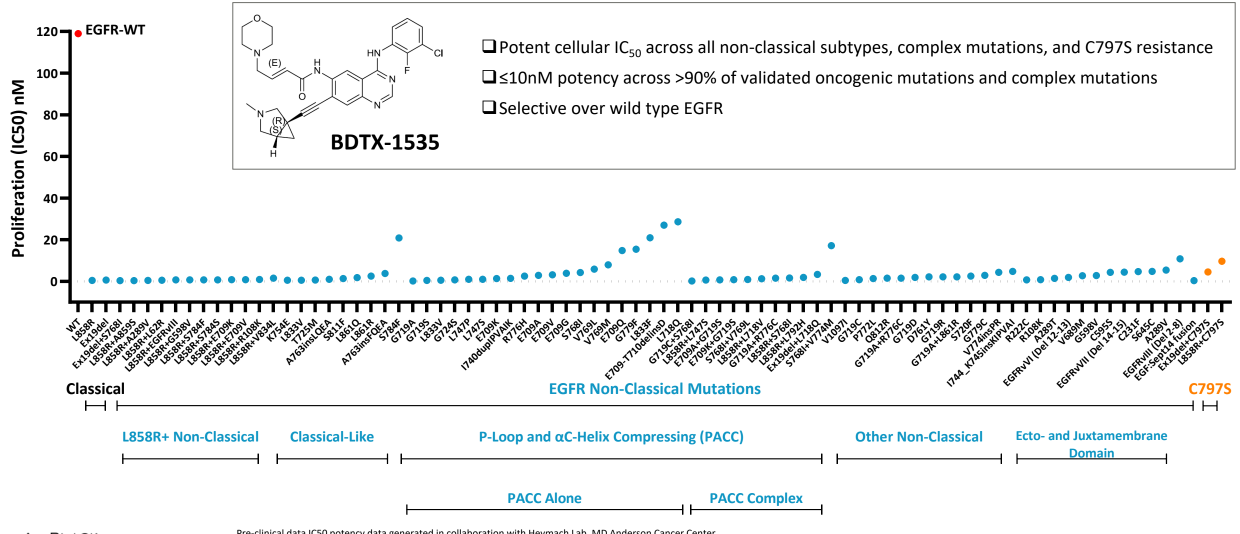
EGFR mutation frequently associated with L858R

Pre-clinical data; EGFR mutations are engineered in Ba/F3 cells



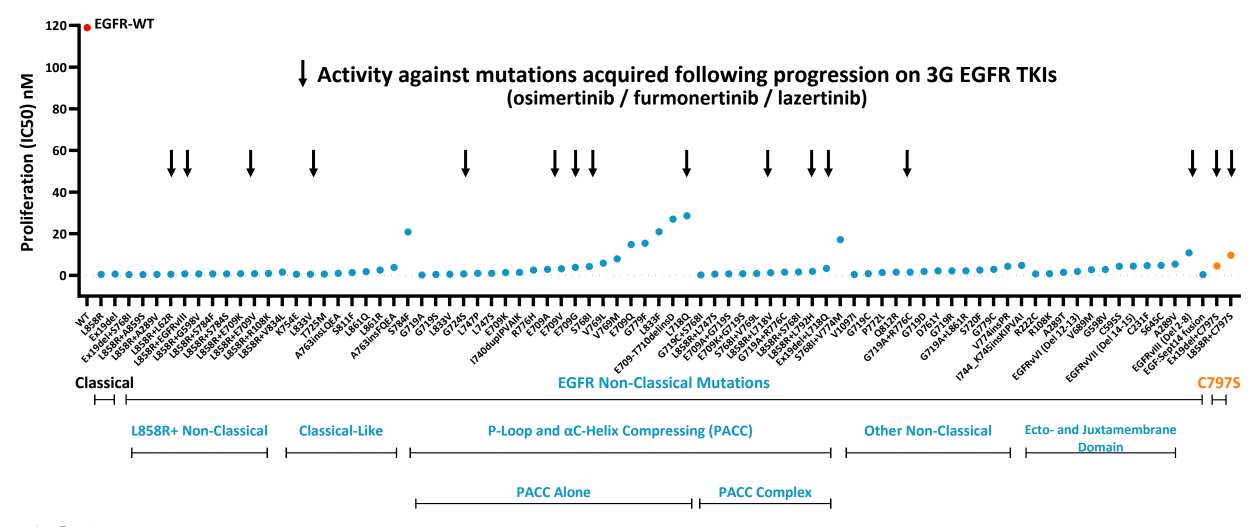
BDTX AACR 2024 oral presentation

BDTX-1535: A MasterKey EGFR Inhibitor Achieving Potent Inhibition of *EGFR Classical* (Ex19del and L858R) & *Non-Classical* Mutations while Sparing EGFR-WT





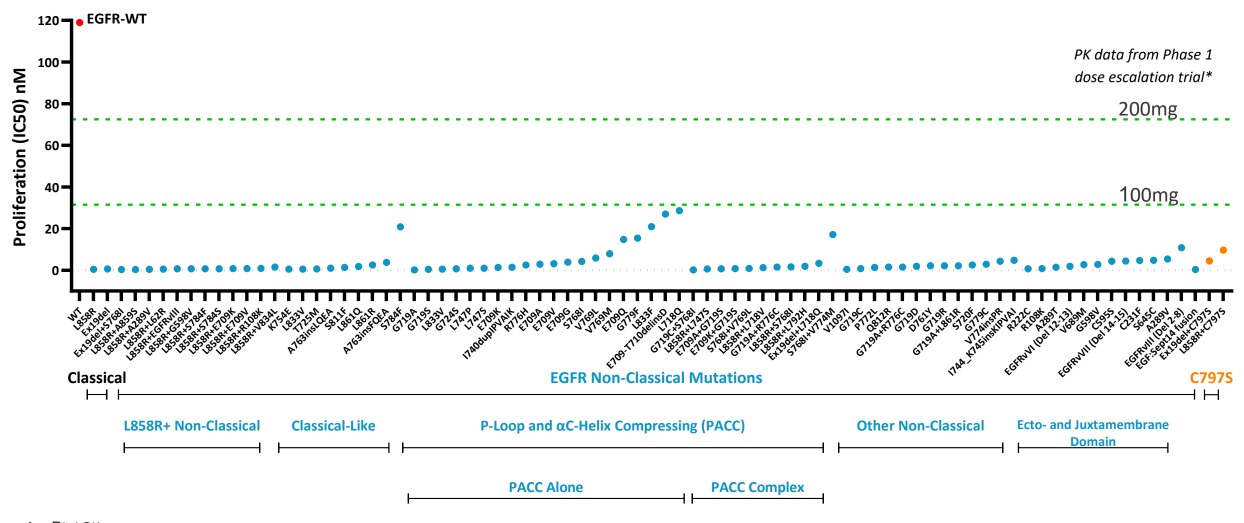
BDTX-1535: A MasterKey EGFR Inhibitor Achieving Potent Inhibition of *EGFR Classical* (Ex19del and L858R) & *Non-Classical* Mutations while Sparing EGFR-WT



BLACK
DIAMOND
THERAPEUTICS

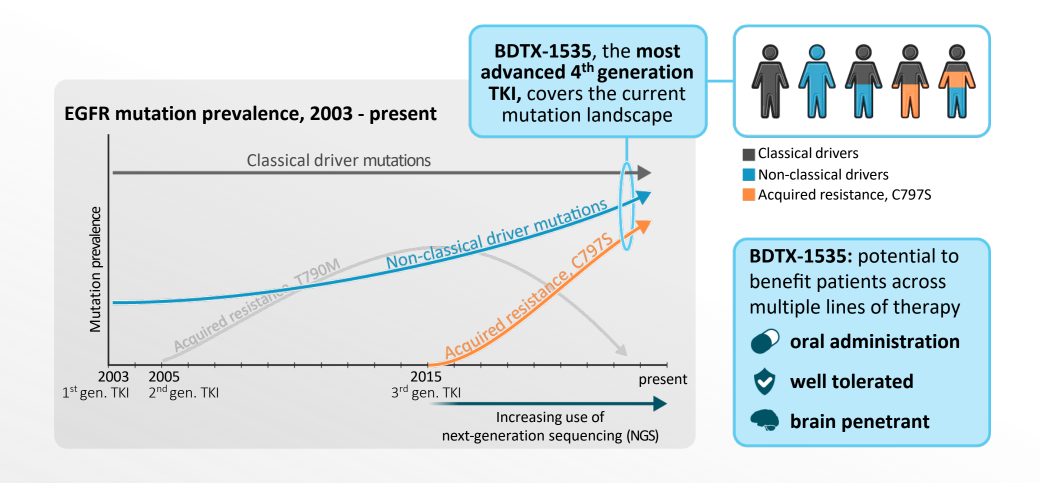
BDTX AACR 2024 oral presentation

BDTX-1535 Achieves Coverage of the EGFR Mutation Spectrum at the Well-Tolerated Oral Doses of 100-200mg QD



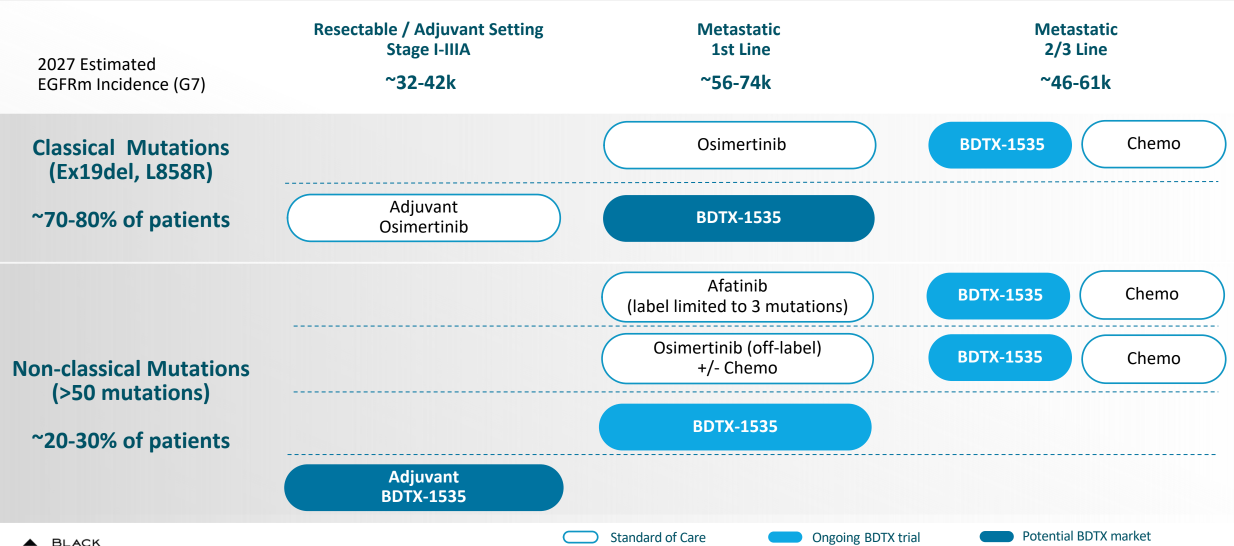


BDTX-1535 Addresses the Most Clinically Relevant EGFR Mutations in NSCLC: Classical / Non-Classical Drivers and C797S Resistance





BDTX-1535 Potential Position: A Well-Tolerated Oral Therapy In Early-Line EGFRm NSCLC





BDTX-1535 Phase 1 Dose Escalation: Summary

Mutation Matched Phase 1 Study Inclusion Criteria

Red	current NSCLC Coh	Recurrent GBM Cohort		
EGFR mutations at the time of progression: - Non-classical driver, OR - Acquired resistance C797S	Progression after EGFR TKI	Exclusion of EGFR T790M, Ex20ins, KRAS mutations, cMET amplification	EGFR alterations at resection/diagnosis	Wild-type isocitrate dehydrogenase (IDH)

Dose Escalation Completed: 15 mg QD to 400 mg QD

- Primary objective:
 PK and safety
- Secondary objective: Anti-tumor activity



- Target coverage and clinical activity at ≥ 100 mg, MTD at 300 mg
- Phase 2 in 2L/3L NSCLC enrolling at 100 mg QD and 200 mg QD



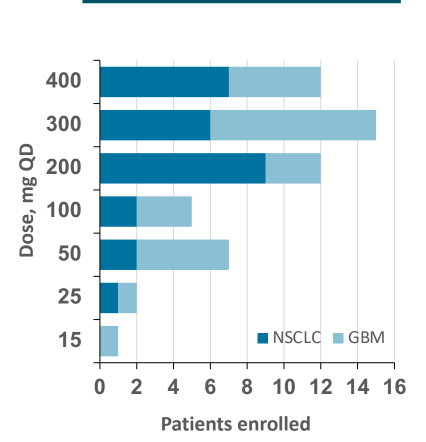
NSCLC Key Data Takeaways

- Once-daily dosing delivers sufficient exposure to inhibit EGFR mutations
- Manageable EGFR TKI safety profile at 200 mg (similar to osimertinib)
- Radiographic responses and durable anti-tumor activity across multiple mutation families
- of mutant alleles, which is predictive of clinical benefit¹
- Phase 2 data expected Q3 2024



BDTX-1535-101 Phase 1 Dose Escalation Patient Characteristics

Patient Enrollment

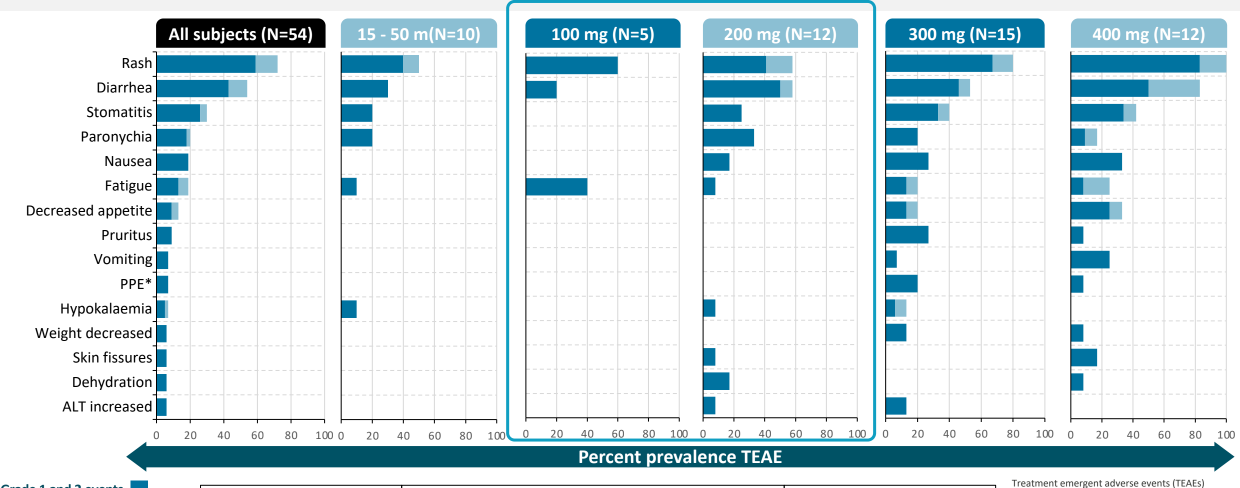


NSCLC			
Patient Characteristics	All Treated		
Age, mean (range)	64 (46, 81)		
Female	18 (67%)		
ECOG PS			
0	7 (26%)		
1	20 (74%)		
Prior lines of therapies			
median (min, max)	2 (1, 9)		
Prior anti-cancer agents			
EGFR TKI	27 (100%)		
Chemo	19 (70%)		
Anti-angiogenic or CPIs	11 (41%)		
HER3-ADC	2 (7%)		
Prior EGFR TKIs			
Osimertinib	23 (85%)		
1st line treatment	17 (74%)		
Erlotinib or gefitinib	9 (33%)		
Afatinib	3 (11%)		
Dacomitinib	1 (4%)		
BLU-701	1 (4%)		

GBM^1				
	All Treated			
Age, mean (range)	58.7 (41, 85)			
Female	10 (37%)			
Karnofsky PS				
90	4 (15%)			
80	12 (44%)			
70	5 (19%)			
60	3 (11%)			
Prior lines of therapies				
median (min, max)	2 (1, 4)			
Prior anti-cancer agents				
TMZ	26 (96%)			
Anti-angiogenic or CPIs	11 (41%)			
Chemo	7 (26%)			



Dose Escalation Treatment Emergent Adverse Events Related to BDTX-1535: Well-Tolerated Profile



Grade 1 and 2 events
Grade 3 events

No Grade 4 AEs were reported

- No dose limiting toxicity (DLTs) were observed at ≤200 mg
- Maximum tolerated dose is 300mg QD

Dose reductions: 1 (8%) pt at 200 mg QD; 5 (33%) pts at 300 mg QD

occurring in ≥6% patients;

All patients in 300 mg cohort received rash prophylaxis

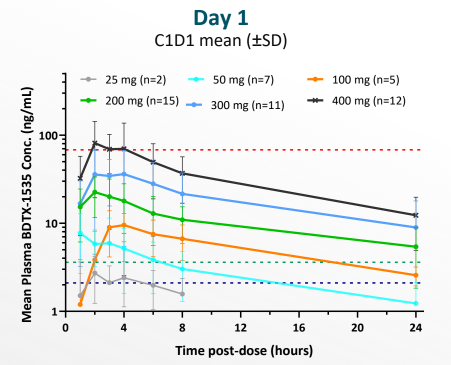
Rash group terms: rash, rash maculo-papular, dermatitis acneiform

*PPE = Palmar-plantar erythrodysaesthesia syndrome

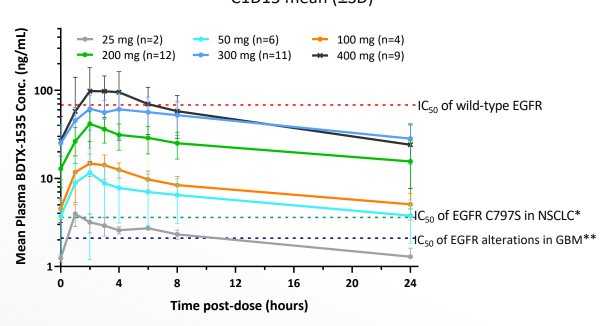


BDTX-1535 Linear Plasma PK Profile Supports Daily Dosing To Achieve 24-Hour Target Coverage

Mean plasma concentration-time profile of BDTX-1535







- Target blockade based on preclinical IC50 was achieved at BDTX-1535 ≥ 100 mg QD
- Exposure was dose proportional with half-life ~15 hours to support daily dosing
- Clinical anti-tumor activity observed at ≥ 100 mg QD

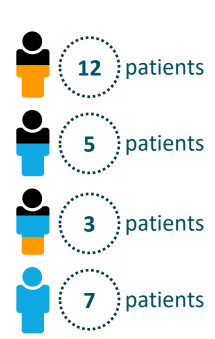


^{*}IC₅₀ of EGFR C797S is average of IC₅₀ of Exon19del/C797S and L858R/C797S mutations tested in Ba/F3 cells

NSCLC Dose Escalation Patients Reflect Real-World EGFR Mutation Landscape Post Osimertinib

20 of 27 patients with 2 or more mutations

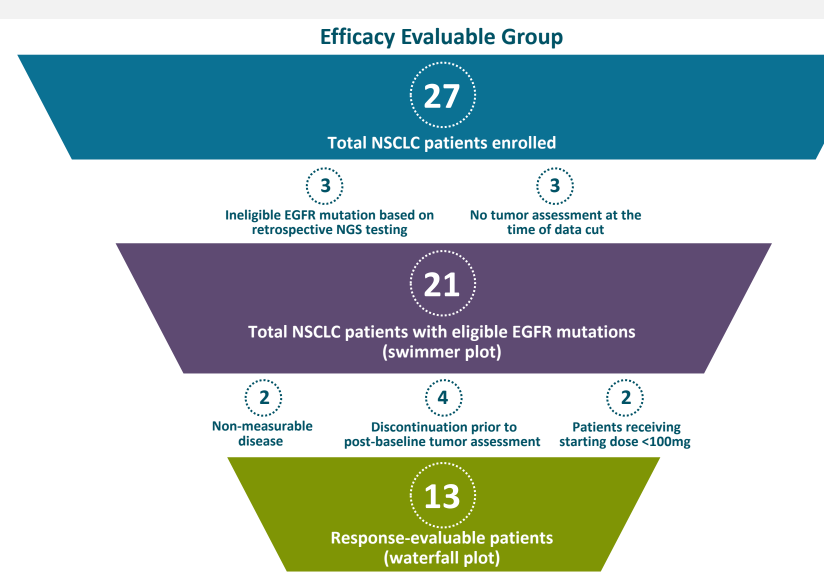




Classical driver mutations	Non-classical driver mutations	Acquired resistance mutation	
Exon 19del L858R	E709A/V L718Q G724S L833V G719A L861Q L747P S768I T751K K754E L747_E759del E746_T751delinsA L747_T751delInsP	C797S	

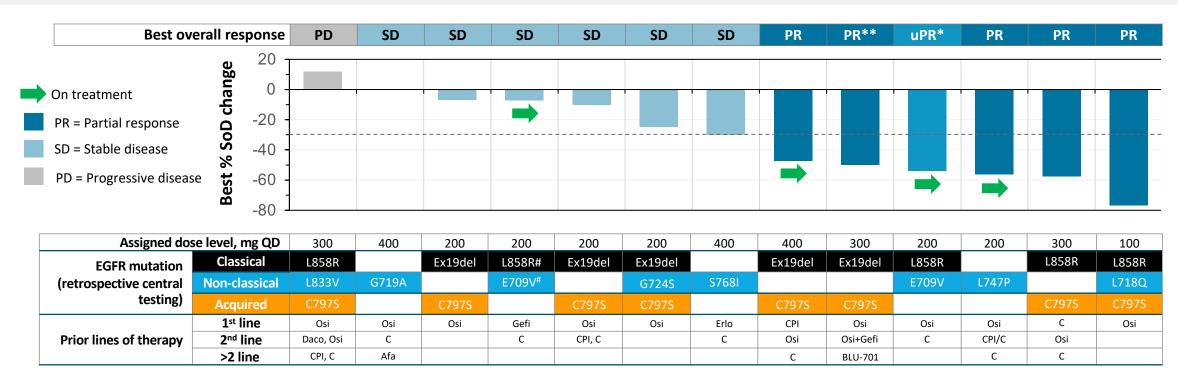


BDTX-1535 Phase 1 Dose Escalation: 13 Response-Evaluable NSCLC Patients





Radiographical Responses in Efficacy-Evaluable NSCLC Patients Across All Relevant Mutations



Osi = Osimertinib; Afa = Afatinib; Gefi = Gefitinib; Daco = Dacomitinib; Erlo = Erlotinib; CPI = Checkpoint inhibitor, C = Chemotherapy; # - mutations were absent on confirmatory test; * uPR=unconfirmed partial response-patient had a PR on a post-baseline scan, but a radiologist was unable to confirm a response on a subsequent scan; this patient remains on study treatment without evidence of PD. **%SoD was updated to -50% from prior data release 24July2023 BDTX-1535-101 clinical data extract

Data from Poster at EORTC/AACR/NCI International Conference on Molecular Targets and Cancer Therapeutics October 2023

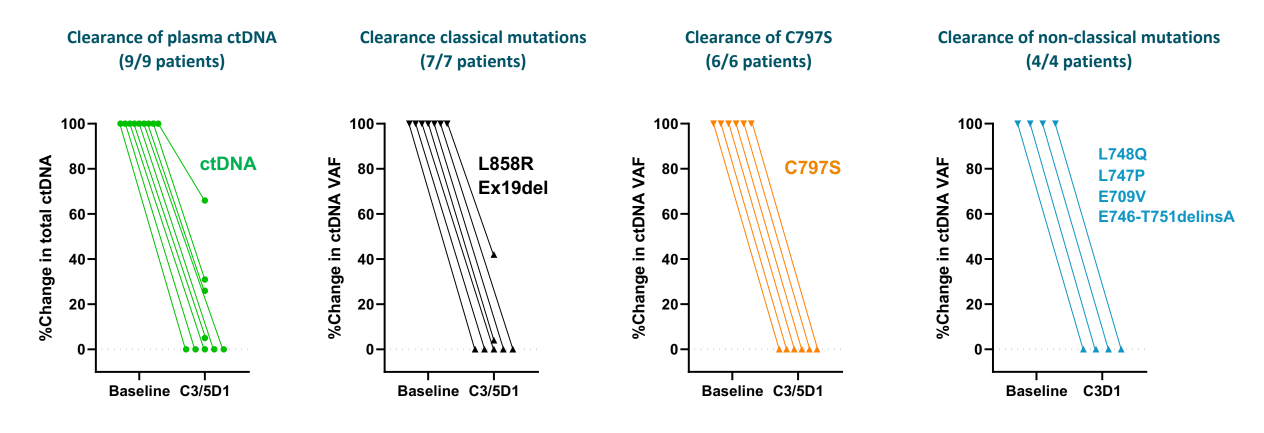
Efficacy-Evaluable Patients 5 cPR, 1 uPR of 13 by RECIST



Post-Osimertinib Patients
5 cPR, 1 uPR of 11 by RECIST



BDTX-1535 Drives Clearance of Mutant EGFR VAF and ctDNA in Phase 1 Trial

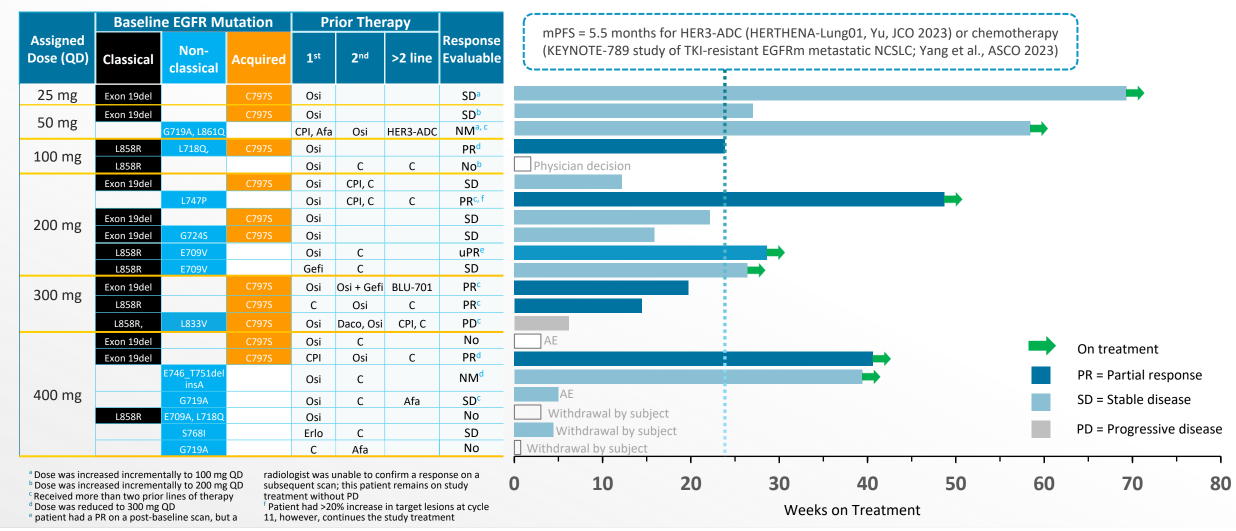




Clearance of plasma ctDNA as well as clearance of EGFR Classical, Non-Classical, and C797S observed with BDTX-1535



BDTX-1535: Emerging Evidence of Durable Tumor Response in NSCLC

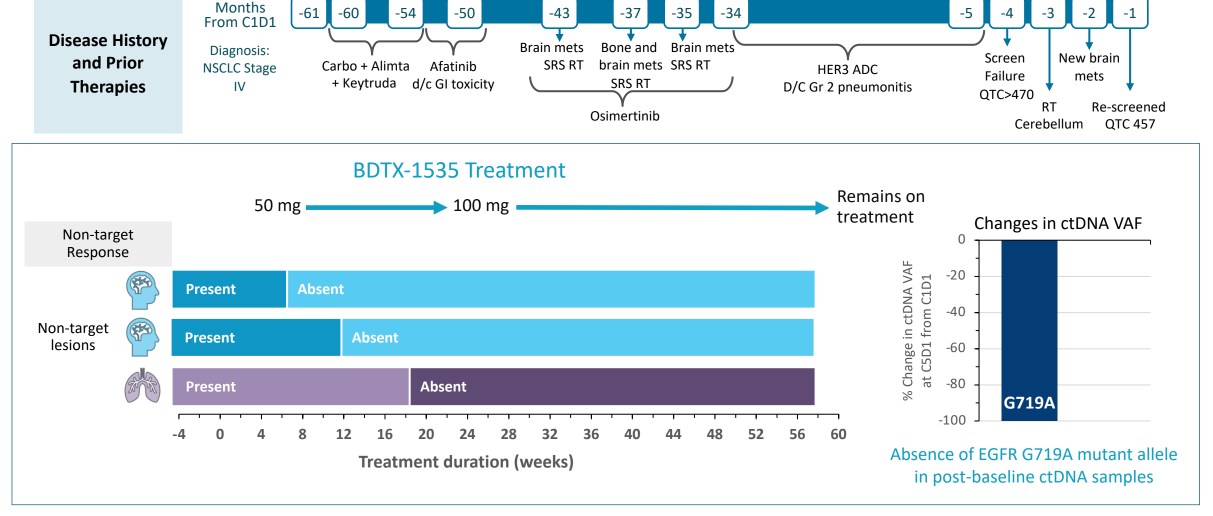




Clinical Benefit in a Non-Response Evaluable Patient With CNS Disease: Remains on Therapy for > 1 year

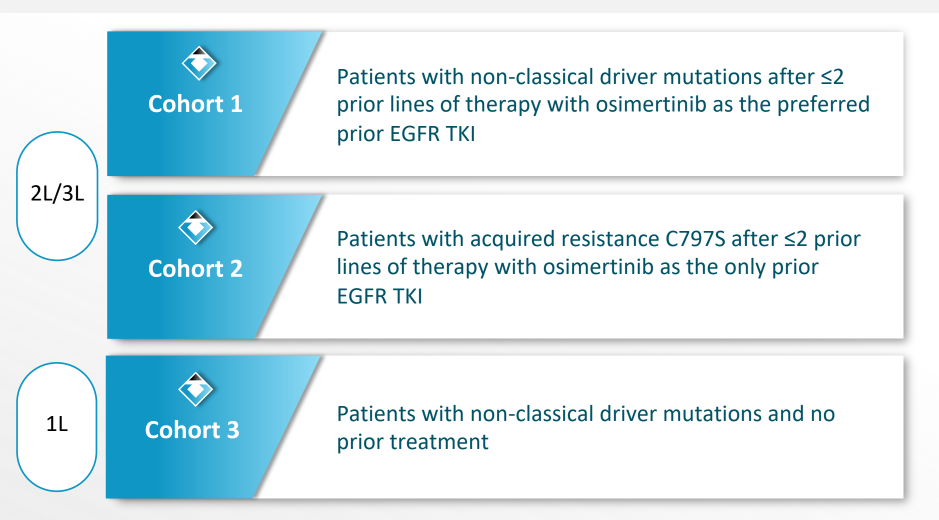
Non-Classical Driver G719A

Non-Classical Driver L861Q¹





BDTX-1535: Phase 2 Trial Enrolling in 1L and 2L/3L NSCLC

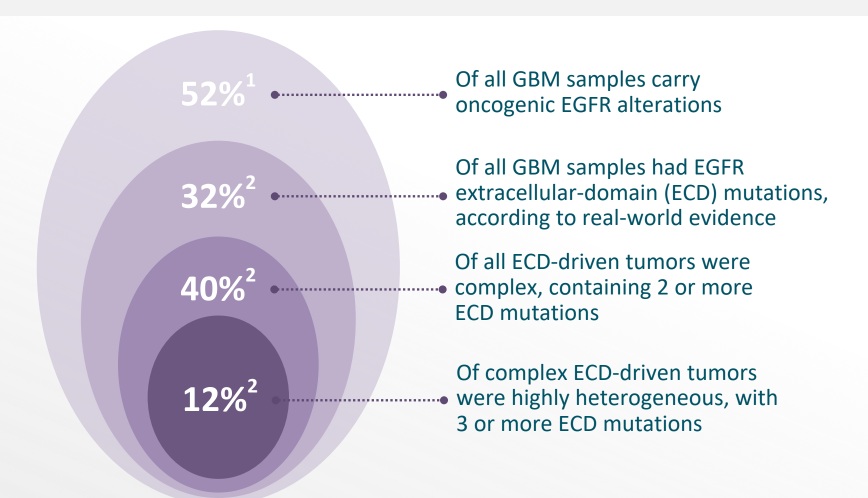


- Enrolling up to 40 patients each cohort
- 2L/3L patients enrolling at 100 mg QD and 200 mg QD
- ORR primary endpoint
- Phase 2 data in 2L/3L patients expected Q3 2024
- FDA Fast Track Designation granted for 2L EGFRm/C797S
- End of Phase 1 FDA feedback received, 1L cohort initiated





Treatment of EGFR-Driven GBM Requires Inhibition of Complex EGFR Mutations: Potent Preclinical Inhibition by BDTX-1535

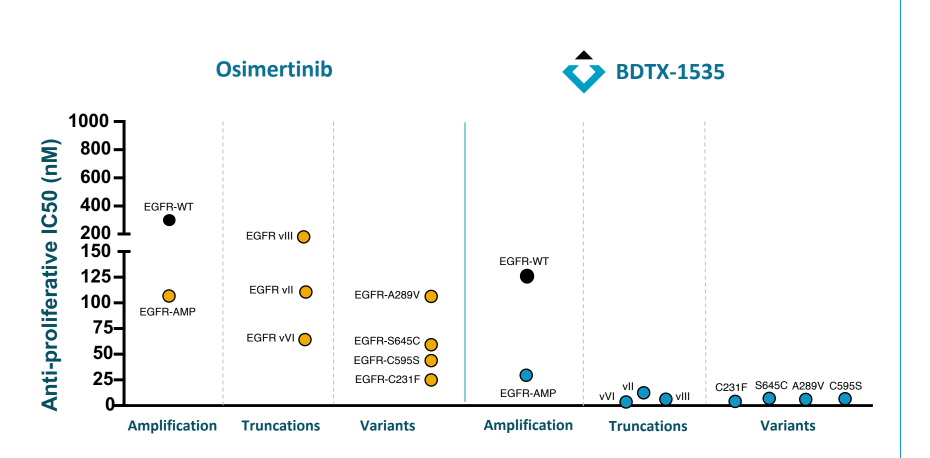


GBM patients in the US are diagnosed each year with EGFR mutations that have been shown in preclinical studies to be inhibited by

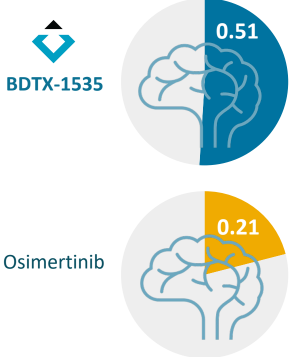
BDTX-1535



BDTX-1535 Demonstrates Potent Preclinical Inhibition of Oncogenic GBM EGFR Alterations vs. Osimertinib and Superior Brain Exposure



BDTX-1535 Exhibits Superior Brain Exposure Kpuu (rat)





BDTX-1535: Potential to Overcome Limitations of Prior Attempts to Drug EGFR in GBM

Lessons From Past Failures



Heterogenic expression of EGFR oncogenic alterations within tumors



Potent MasterKey inhibition of co-occurring EGFR alterations and amplification

Paradoxical activation of EGFR GBM oncogenes induced by reversible inhibitors



Covalent MOA and no paradoxical activation

Poor tolerability driven by on target WT-EGFR activity



Spares WT-EGFR in normal cells while retaining potent activity against EGFR alterations

Low brain exposure due to a lack of CNS penetrance



Designed to be brain-penetrant to treat CNS tumors

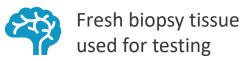


BDTX-1535 Opportunity in Newly Diagnosed EGFRm GBM Patients

EGFR Driver Status Often Evolves During Treatment Newly Diagnosed Temozolomide + Radiation Recurrence Tumor evolves with Fresh biopsy not Fresh biopsy time and treatment¹⁻⁵ available for majority of recurrent patients^{6,7} EGFR alteration status Treatment not may change in up to matched to **EGFR** status characterized 40% of cases¹⁻³ mutational profile BDTX-1535 BDTX-1535 Opportunity Ph1 Complete

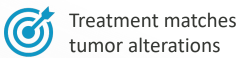
GBM Treatment Paradigm

Opportunity for BDTX-1535 in Newly Diagnosed Patients





Up-to-date test results guide treatment





Promising Clinical Activity in Heavily Pre-treated GBM Patients From Dose Escalation Portion of Phase 1 Study



- Recurrent, heavily pre-treated (2L/3L+)
- Historical PFS ~ 2-4 months
- EGFR status *not* confirmed at dosing

- ✓ Well tolerated, with favorable plasma PK
- ✓ Of 22 efficacy evaluable heavily pre-treated GBM patients
 - 3 patients on therapy longer than 10 months
 - 1 remains on therapy at 15 months (100mg QD), had progressed on TMZ after 3 months
 - 1 patient on therapy longer than 6 months
 - 5 patients on therapy longer than 4 months
- ✓ Of 19 patients with measurable disease assessable by RANO criteria
 - 1 confirmed partial response (200 mg QD), on therapy > 4 months
 - 8 patients with stable disease



BDTX-1535 in GBM: Next Steps and 2024 Milestones



Window of Opportunity (WOO) Study Overview

Sponsored by Ivy Brain Tumor Center

- Patients receive 5 days of dosing with BDTX-1535 as monotherapy
- Surgical resection following day 5 of dosing
- Tumor PK and EGFRm status evaluated
- If PK threshold met, patient can remain on therapy

Anticipated Upcoming Milestones

Phase 1 full data set at medical meeting Q2 2024

"Window of Opportunity" trial currently enrolling, data expected at a medical meeting Q2 2024

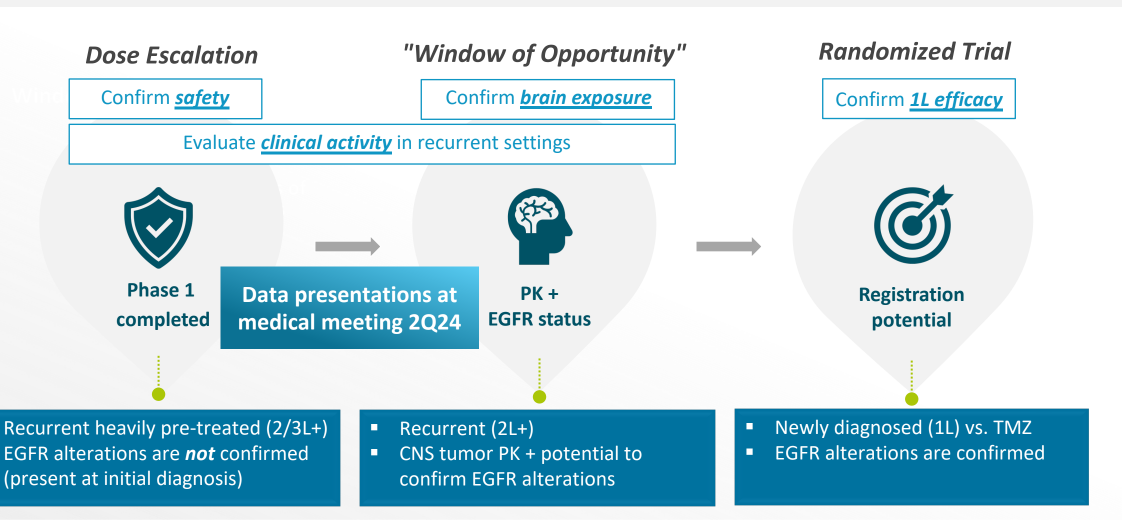


If go forward criteria met:

Opportunity to benefit 1L patients with confirmed EGFR status at diagnosis



BDTX-1535 GBM Development Path Designed for Sequential De-Risking





BDTX-1535 Summary: NSCLC and GBM Opportunities

Leading Position



First and potential best-in-class
4th gen EGFR TKI

Robust clinical POC in a heavily pre-treated Phase 1 NSCLC population

- 5 cPR + 1 uPR out of 13 efficacy-evaluable patients
- Durable responses and clinical evidence of CNS anti-tumor activity
- Well tolerated with manageable (similar to osimertinib), on-target EGFR TKI AEs
- Phase 2 enrolling, 2L/3L data expected in Q3 2024

Compelling Asset



Differentiated profile

Clear differentiation against standard of care and emerging treatment options

- WT-EGFR sparing with favorable clinical tolerability vs chemo/ADC-based combos
- Brain-penetrant to address CNS disease
- Highly potent against all major, clinically relevant EGFR mutations
- Once daily oral administration

Large Markets



Robust near-term commercial opportunity in 1L and 2L NSCLC

Emerging potential in GBM

Real-world evidence¹⁻³ demonstrates a growing commercial opportunity in NSCLC

- 1L: Non-classical driver mutations
- 1L: Post-osimertinib adjuvant therapy
- 2L: Post osimertinib C797S, classical/non-classical drivers and complex mutations

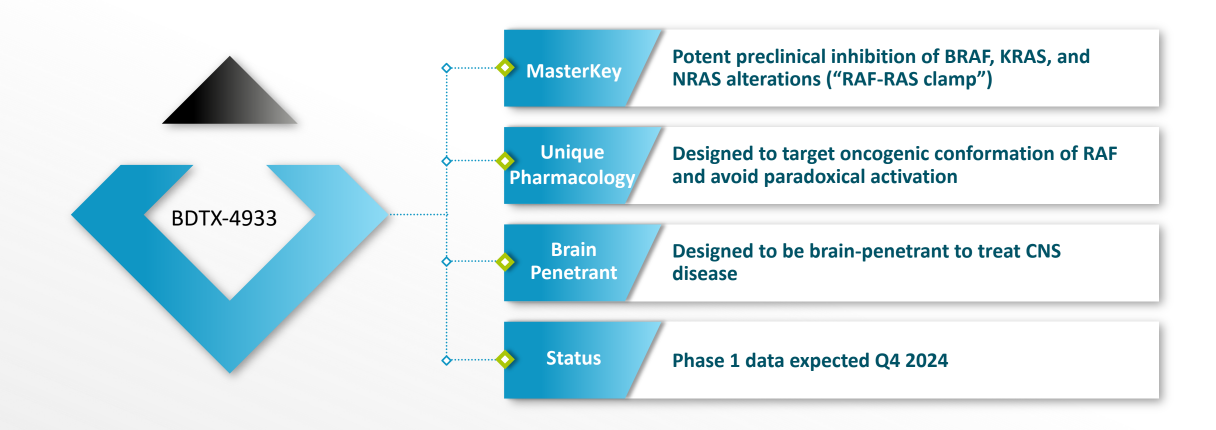
Potential in 1L GBM with EGFR alterations (>50% of all newly diagnosed GBM)







BDTX-4933: Oral, Brain-Penetrant, RAF MasterKey Inhibitor



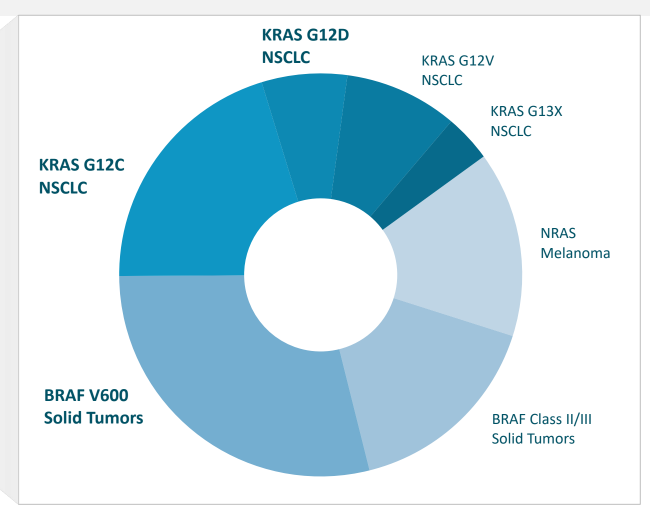


MAPK Pathway Mutations Affecting KRAS/NRAS/BRAF are Among the Most Common Oncogenic Mutations in Cancer

Addressable US / EU5 / JP Patient Population

~318,660

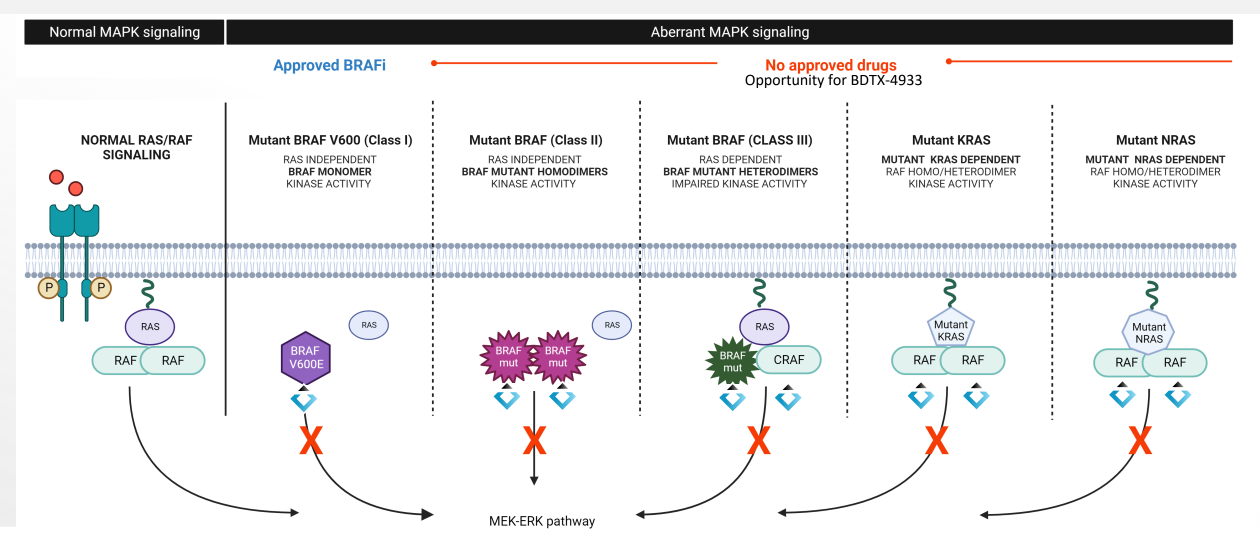
~112,00 with CNS metastasis



Source: EvaluatePharma, TCGA, GENIE-11



BRAF Alterations Drive Oncogenesis Through Hyperactivation of the RAS-MAP Kinase Pathway: Multiple Opportunities for BDTX-4933





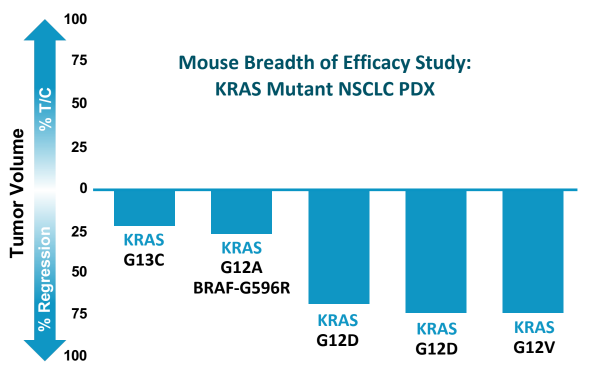
1. Zhang, W., Cell Res. (2002); 2. Yuan, J., J Hematol Oncol 13, 113 (2020); 3. Yao Z, Cancer Cell (2015); 4. Karoulia Z, Cancer Cell (2016); 5. J. Wang, Pharmacol. Res. 129, 414–423 (2018); 6. H. Ellens, Drug Metab. Dispos. 45, 646–656 (2017); 7. R. K. Mittapalli, J. Pharmacol. Exp. Ther. 344, 655–664 (2013); 9. Belum VR, Ann Oncol. (2015); 10. Su F., N Engl J Med. (2012); 11. Hatzivassiliou G, Nature. (2010); 12. Poulikakos Pl., Nature, (2010)

BDTX-4933 Demonstrates Potent Preclinical Inhibition of a Spectrum of BRAF/RAS and KRAS Mutations in Cell Lines and PDX Models

Potential Best-in-Class Potency Compared to Other RAF Inhibitors

Potent and selective inhibition of proliferation across tumor cell lines with MAPK pathway mutations

		Cell Proliferation IC50				
Mutation		BDTX-4933	Naporafenib	Belvarafenib	Exarafenib	Encorafenib
BRAF Class I	V600E					
	BRAF fusion					
DDAE Glove II	BRAF fusion					
BRAF Class II & non-V600	L597V					
	L245F					
	BRAF indel					
NRAS	NRAS Q61K					
MAS	NRAS Q61L				Not available	
NRAS BRAF	WT					Paradoxical Activation

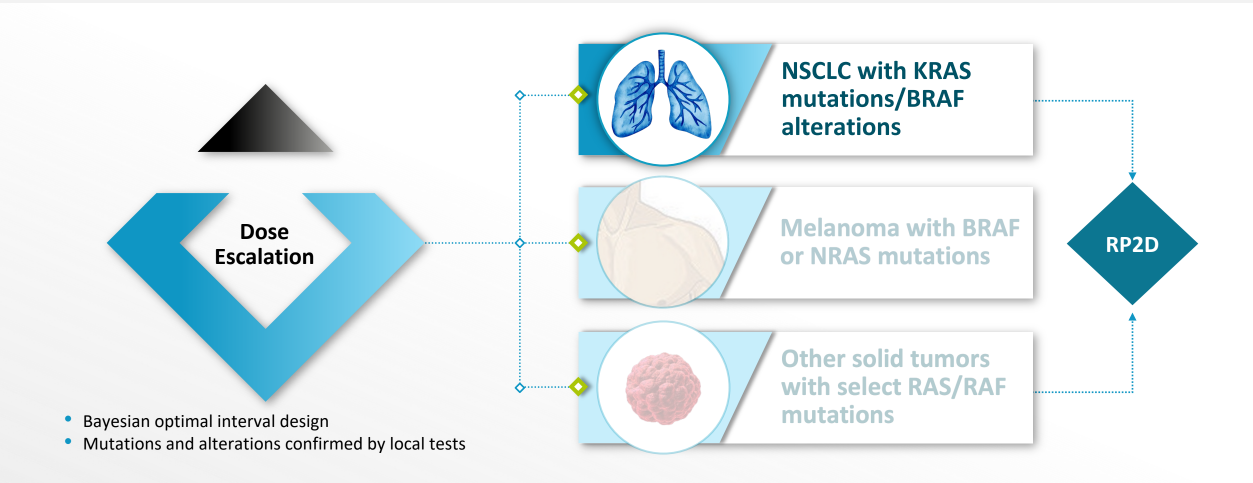


No significant body weight loss observed BDTX-4933: 10 mg/kg QD x 28 or 5 mg/kg BID x 56





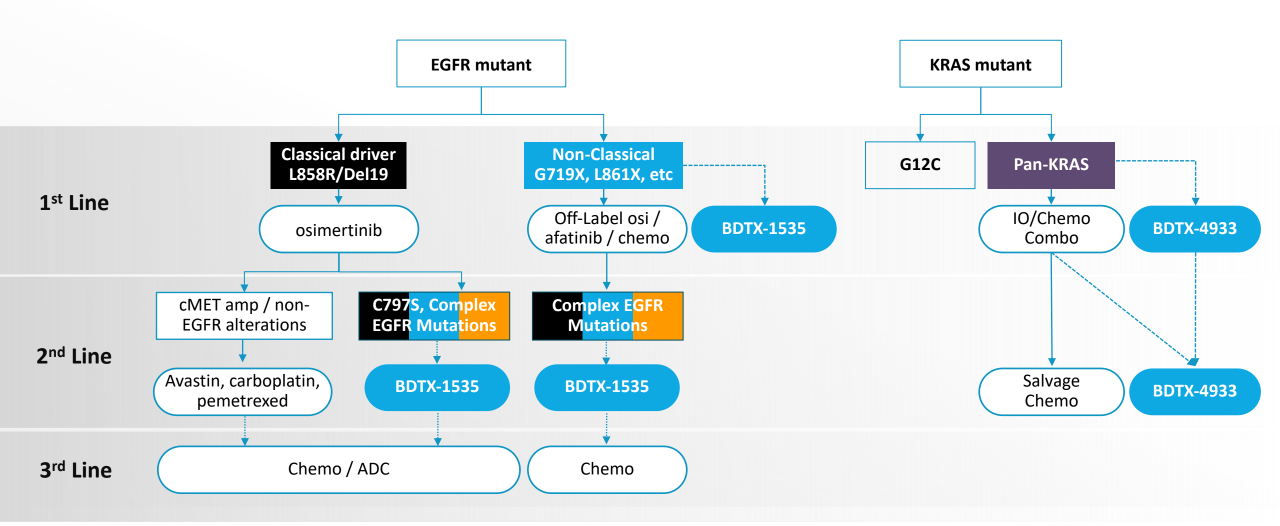
BDTX-4933: Focused, Biomarker-Driven Phase 1 Trial Initiated Data Anticipated in Q4 2024





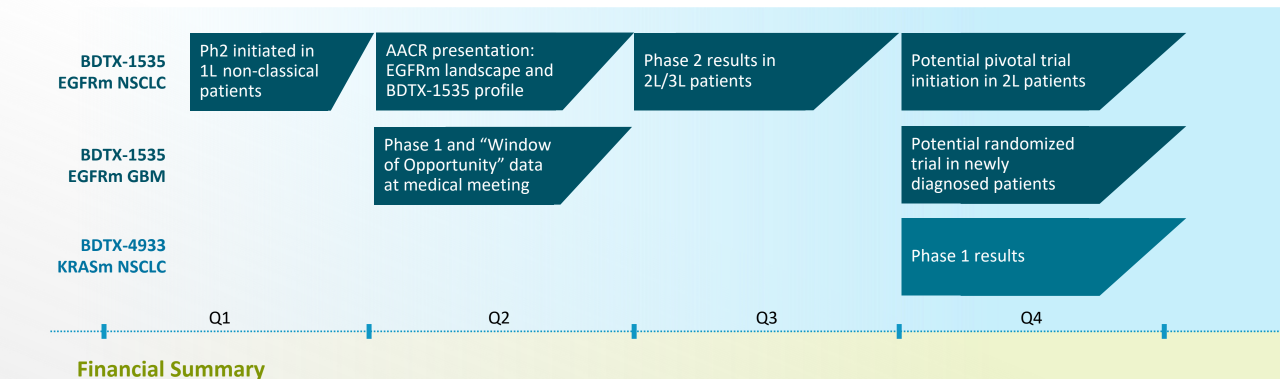
BDTX-1535 and BDTX-4933: Potential for NSCLC Franchise







Anticipated 2024 Key Milestones





\$131.4m

as of December 31, 2023

Cash runway into Q2 2025

