



Protein degraded.  
Disease targeted.  
Lives transformed.

43<sup>rd</sup> Annual J.P. Morgan Healthcare Conference  
January 2025



# Forward-looking Statements and Intellectual Property

## FORWARD-LOOKING STATEMENTS

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## Our Mission

To deliver on the promise of targeted protein degradation science to create a new generation of medicines that transform patients' lives.

Our portfolio of degrader medicines pursues targets that may benefit from a degrader approach:

### **Cemsidomide**

targeting IKZF1/3 for multiple myeloma and non-Hodgkin's lymphoma

### **CFT1946**

targeting BRAF V600 mutant for solid tumors including melanoma & colorectal cancer

### **CFT8919**

targeting EGFR L858R for non-small cell lung cancer

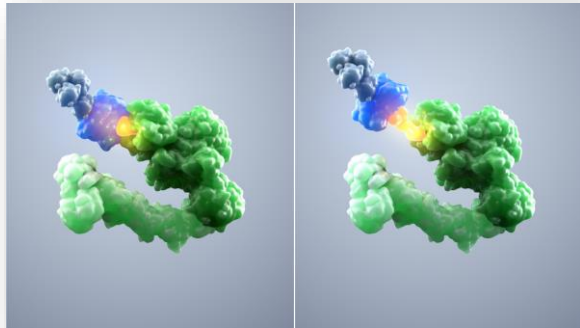
### **Internal Discovery Pipeline**

targets with unmet need and strong degrader rationale

# C4T Has Been at the Forefront of TPD Science and Is On the Path to Becoming a Fully Integrated Biotechnology Company

## Leading the Way in Designing Orally Bioavailable Degraders

2015 – 2020



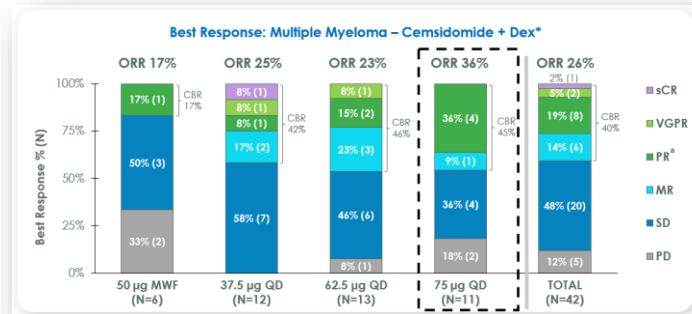
Built TORPEDO platform to design highly catalytic, orally bioavailable degraders

Established collaborations with leading global pharmaceutical companies, building expertise across a range of diseases and target classes

Assembled strong catalog of intellectual property

## Demonstrating Proof of Concept

2020 – 2025



Progressed four development candidates into clinical trials, including three trials underway today

Delivered two development candidates to a collaborator for non-oncology targets

Achieved blood-brain barrier penetration in several development candidates

## Delivering on the Promise of Targeted Protein Degradation

2025 and beyond



Advancing clinical programs to approval for patients with high unmet needs

Leveraging TORPEDO platform to develop a sustainable pipeline against targets of high unmet need with strong degrader rationale in large commercial markets

Expanding application of targeted protein degradation through high-value collaborations

# C4T Is Advancing a Portfolio of Rationally Designed Degraders Against Targets of High Unmet Need in Oncology

## Cemsidomide

Targeting IKZF1/3  
Transcription Factor

## CFT1946

Targeting BRAF V600X  
Scaffolding Kinase

## CFT8919

Targeting EGFR L858R  
Receptor Tyrosine Kinase

### Degrader Rationale

Approved IKZF1/3 degraders were not rationally designed and provide less than optimal potency and selectivity

Degradation has the potential to avoid resistance mechanisms and enable a longer duration of response

Degradation facilitates targeting an allosteric L858R-specific binding site distinct from current inhibitors leading to increased selectivity over wild-type, activity against inhibitor resistance mutations & an opportunity for use in conjunction with current SOC

### Clinical Progress

#### Data to date supports best-in-class profile:

- ✓ Differentiated safety profile
- ✓ Competitive ORR in combination with dex at 75 µg in MM
- ✓ Immune activity demonstrated as monotherapy
- ✓ Encouraging ORR and CMR rate in PTCL

#### Data to date demonstrates:

- ✓ Proof of mechanism established
- ✓ Early signs of anti-tumor activity in Phase 1 dose escalation

- ✓ Clinical trial initiated in Greater China<sup>1</sup>

### Potential Patient Population

#### Across U.S., EU4 and UK:

- MM: ~65,000<sup>2</sup>
- PTCL: ~16,000<sup>2</sup>

#### Across U.S., EU4 and UK:

- Melanoma: ~66,000<sup>3</sup>
- Colorectal cancer: ~33,000<sup>3</sup>

#### Across U.S., EU4, UK and China:

- EGFR L858R Mutated NSCLC: ~219,000<sup>4</sup>

### Commercial Rights



**TORPEDO platform has produced highly catalytic, brain penetrant, oral degraders across a broad range of target classes**

<sup>1</sup>License and collaboration agreement with Betta Pharmaceuticals for development and commercialization in Greater China

<sup>2</sup>NCI SEER, consulting engagements with Health Advances and Clearview.

<sup>3</sup>2024 Evaluate Ltd. US + EU4 + UK population

<sup>4</sup>EvaluatePharma, consulting engagements with Health Advances and Clearview. Complete metabolic response (CMR); dexamethasone (dex); multiple myeloma (MM); Overall response rate (ORR); peripheral T-cell lymphoma (PTCL); Germany, Italy, France, and Spain (EU4); Standard of care (SOC)

# Multiple 2025 Data Based Inflection Points to Unlock Value Across Entire Portfolio

## Cemsidomide

*IKZF1/3*

**2025:** Enable initiation of the next phase of clinical development for MM and PTCL. These new studies are expected to initiate in early 2026

**2H 2025:** Complete Phase 1 dose escalation trial in MM and NHL and present data

**2H 2025:** Open expansion cohort(s) in PTCL in the ongoing Phase 1/2 trial

## CFT1946

*BRAF V600 Mutant*

**1H 2025:** Complete monotherapy Phase 1 dose escalation trial in BRAF V600 mutant solid tumors

**2H 2025:** Generate data from Phase 1 cohorts evaluating CFT1946 as a monotherapy in melanoma, in combination with trametinib in melanoma, and in combination with cetuximab in CRC to define and enable next phase of development

**2H 2025:** Present Phase 1 data: monotherapy in BRAF V600 mutant solid tumors; monotherapy expansion cohorts in melanoma; combination with cetuximab in colorectal cancer

## CFT8919

*EGFR L858R*

**Year-end 2025:** Utilize data from Phase 1 dose escalation trial in Greater China to inform ex-China clinical development

## Discovery

**2025:** Present and publish preclinical work from internal pipeline and TORPEDO platform

**2025:** Advance internal and collaboration programs to key discovery milestones

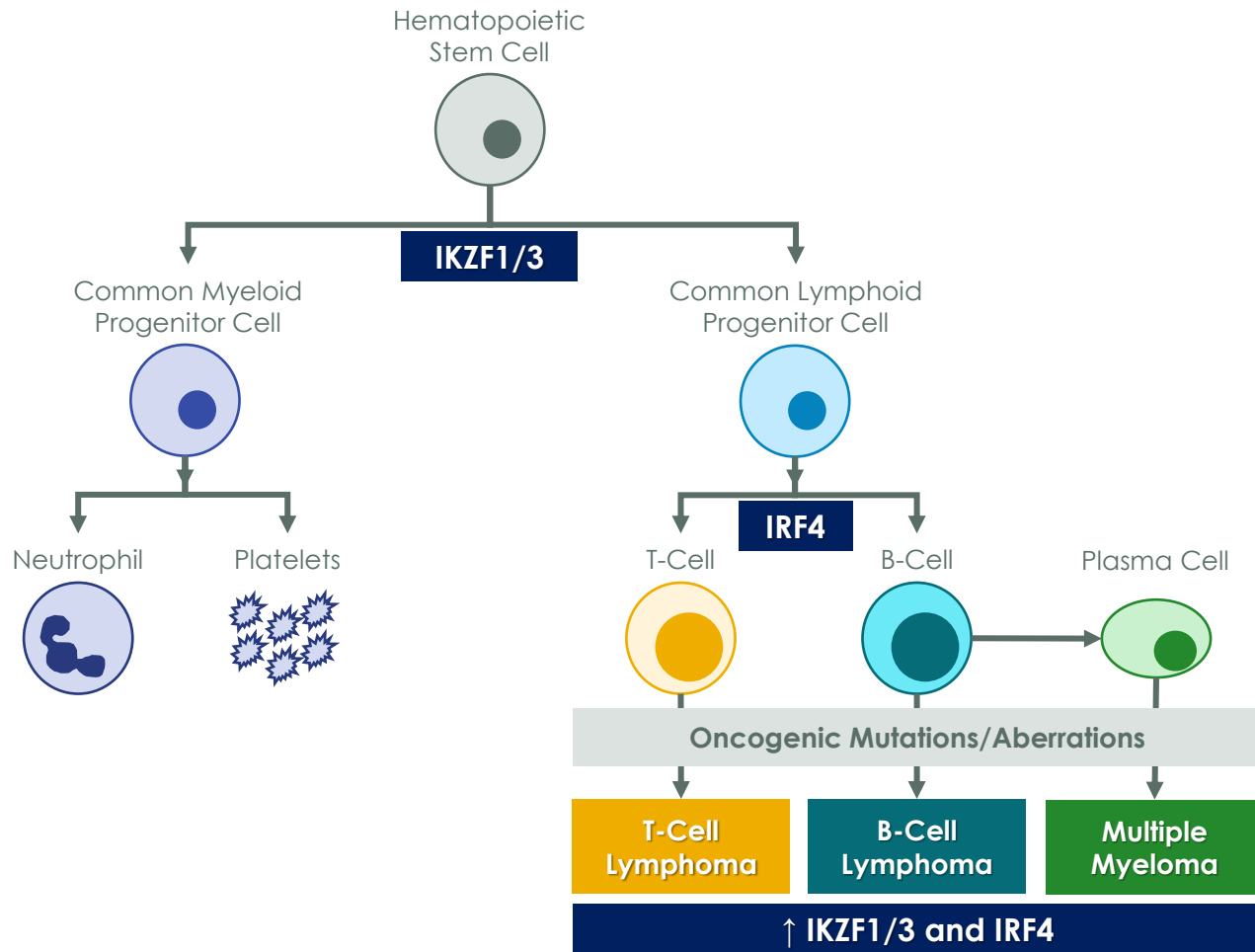
# Cemsidomide

IKZF1/3 Degradator

Multiple Myeloma & Non-Hodgkin's Lymphoma



# IKZF1/3 Are Key Promoters of Myeloma and Lymphoma Cell Survival and Will Remain Important Therapeutic Targets for These Indications



## Key Roles of IKZF1/3:

- Multiple myeloma and lymphoma cells rely on **IKZF1/3** and **IRF4** for survival
- Degrading **IKZF1/3** leads to down regulation of **IRF4**, promoting myeloma and lymphoma cell death and on-target neutropenia
- **IKZF1/3** degradation combined with MM immune-based regimens have potential to enhance activity through T-cell activation and cancer cell death by downregulation of **IRF4**

## Degrader Advantages of Cemsidomide:

- ✓ Cemsidomide is more potent than approved and development stage IKZF1/3 degraders
- ✓ Increased selectivity for IKZF1/3 resulting in reduced off-target toxicity

# Cemsidomide Phase 1 Dose Escalation Trial in MM and NHL Continues to Progress

## PHASE 1 DOSE ESCALATION TRIAL

**R/R MM**  
Monotherapy

**Dosing:** QD

21 days on/  
7 days off



**Status:** Complete

- Different dosing schedule required due to longer than anticipated half-life

**R/R MM**  
Monotherapy

**Dosing:** MWF & QD

14 days on/  
14 days off



**Status:** Complete

- Confirmed 14 days on/14 days off schedule
- Clinical evidence of anti-myeloma activity and immune T-cell activation

**R/R MM**  
Dex Combo

**Dosing:** MWF & QD

14 days on/  
14 days off

**Status:** Ongoing

- Currently enrolling at 100 µg QD
- Have not exceeded maximum tolerated dose

**R/R NHL**  
Monotherapy

**Dosing:** MWF & QD

14 days on/  
14 days off

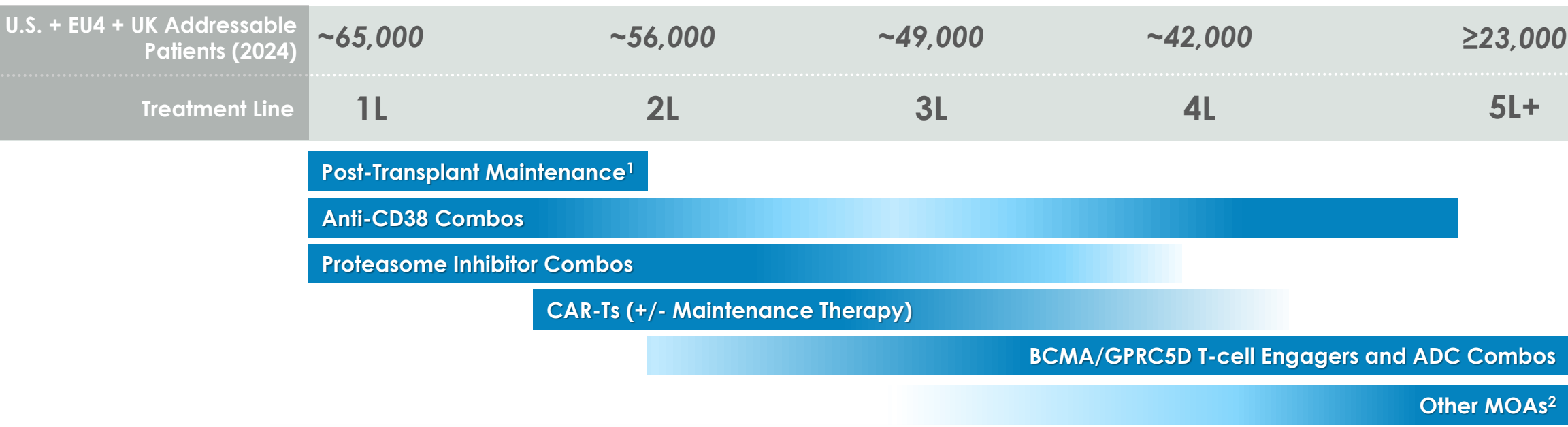
**Status:** Ongoing

- Currently enrolling at 75 µg QD
- Additional dose finding ongoing

Monday, Wednesday, Friday dosing (MWF); once daily (QD); relapsed refractory multiple myeloma (R/R MM); relapsed refractory non-Hodgkin's lymphoma (R/R NHL)

# With a Potentially Best-in-Class Profile, Cemsidomide Is Positioned to Be an IKZF1/3 Degradator of Choice Across Various Combinations

## EVOLVING MULTIPLE MYELOMA TREATMENT LANDSCAPE



## CEMSIDOMIDE OPPORTUNITY

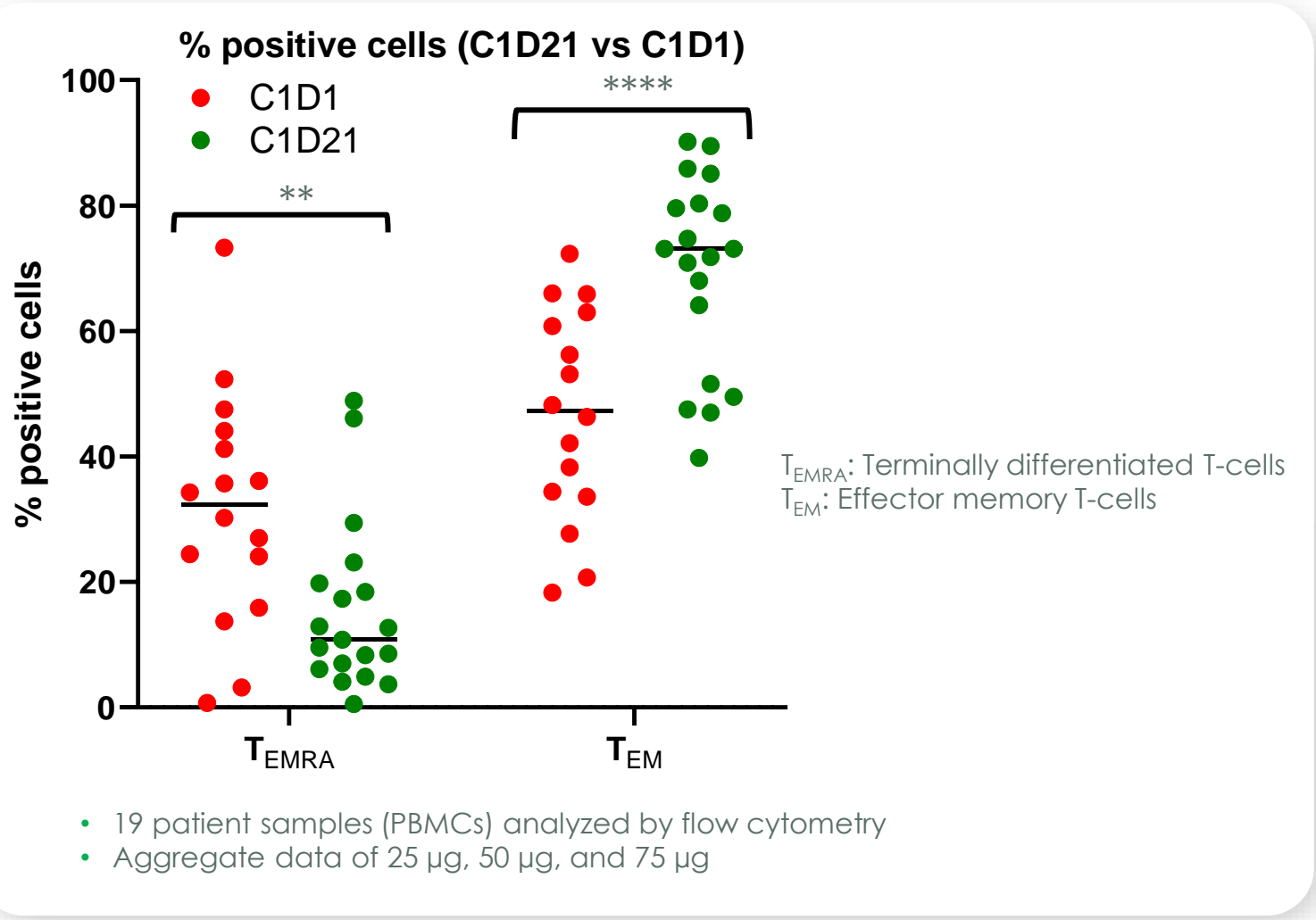
- IKZF1/3 degraders are currently used across lines of therapy, from post-transplant maintenance to 5L+ doublets, and will remain backbone therapies in various combination approaches
- **Cemsidomide has the potential to become the IKZF1/3 degrader of choice** in numerous regimens across lines of therapy given its potent anti-myeloma activity, differentiated safety profile, and immunomodulatory effects

<sup>1</sup> Approximately 30% of multiple myeloma patients undergo a hematopoietic stem cell transplant and receive post-transplant maintenance therapy.

<sup>2</sup> Other MOAs approved in MM include dexamethasone combos, anti-SLAMF7 mAbs and XPO1 inhibitors. Potential future treatment options include FcRH5 bispecific T-cell engagers, BCL-2 inhibitors, and others. Sources: EvaluatePharma (accessed 1/8/25), NCCN guidelines, consulting engagements with Health Advances and Clearview.

B-cell maturation antigen (BCMA); G protein-coupled receptor, class C, group 5, member D (GPRC5D); monoclonal antibodies (mAbs); mechanism of action (MOA); Germany, Italy, France, and Spain (EU4)

# Clinical Evidence of Immune T-cell Activation With Cemsidomide Monotherapy

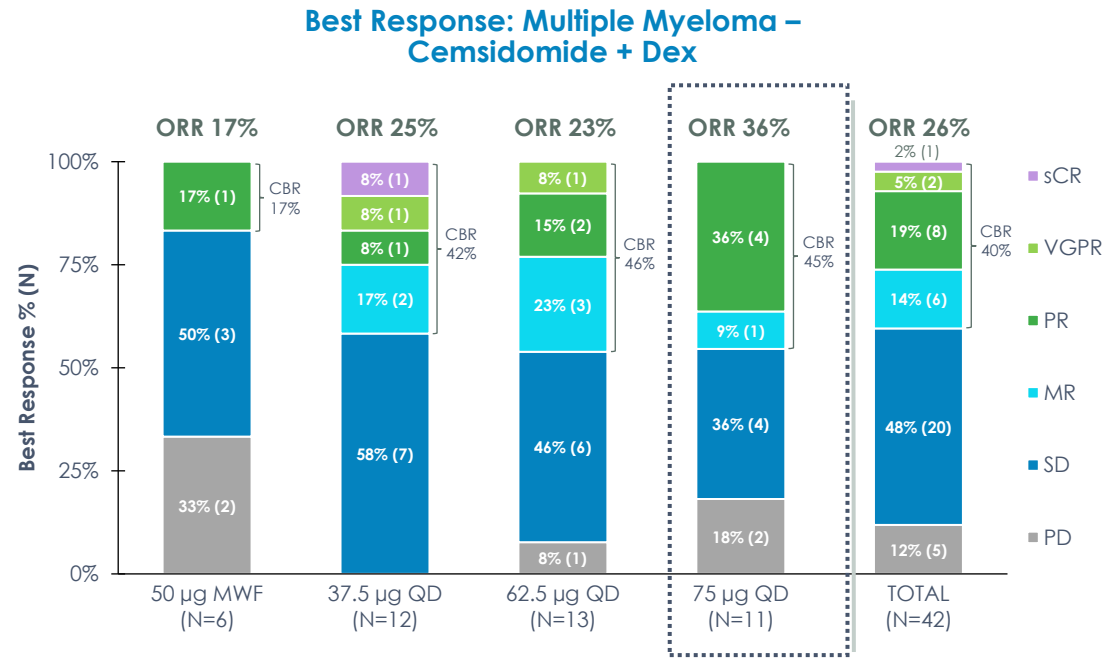


**Supports potential of cemsidomide as a maintenance therapy option and in combination with other MM agents to improve efficacy:**

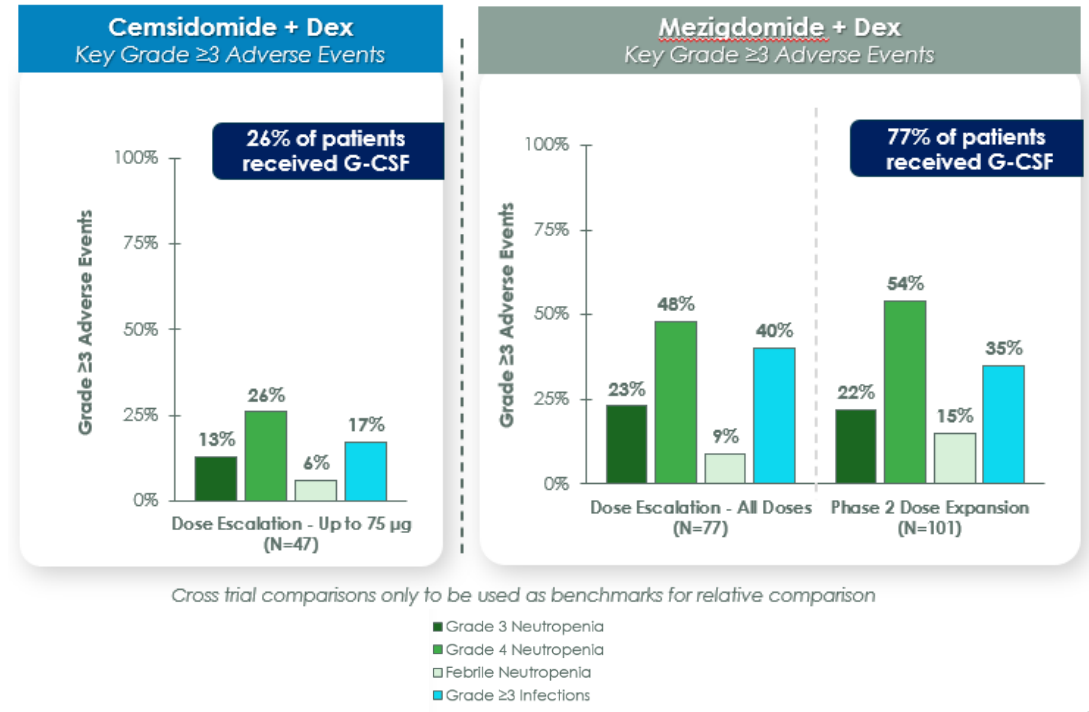
- ✓ Cemsidomide induces CD8+ T-cell activation by increasing effector memory T-cell subset
- ✓ T-cell activation is observed at well-tolerated monotherapy clinical doses
- ✓ Clinical data consistent with the preclinical *in vitro* data reported for cemsidomide

# Cemside + Dex Data Demonstrated a Potential Best-in-Class Profile

75 µg Cemside Dose Level Resulted in a 36% ORR and 45% CBR



Cemside's Differentiated Safety Profile Is Well Suited for Combination Studies, Potentially Resulting in Fewer Disruptive Adverse Events



**Profile supports cemside's development in combination with dexamethasone in the late-line setting and in combination with other MM agents in earlier lines of the treatment paradigm**

Source: ASH 2024; C4T data as of 10/11/2024 (<https://ir.c4therapeutics.com/static-files/32ae4fdb-d4d9-4a17-a77d-83289c66e91f>); see slide 20 of the ASH presentation for complete footnotes for the left panel; see slide 21 of the ASH presentation for complete footnotes for the right panel

# Cemsidomide Has the Potential to Become a Treatment Option Across Lines of Therapy and Address a Growing Patient Population

## INITIAL CEMSIDOMIDE DEVELOPMENT PATH IN MM

U.S. + EU4 + UK Addressable Patients (2024) <sup>1</sup>	~65,000	~56,000	~49,000	~42,000	≥23,000
Treatment Line	1L	2L	3L	4L	5L+

**Studies Currently Being Enabled**

**Phase 1/2 (Escalation/Expansion)**  
Cemsidomide/Dex + Anti-CD38 and Cemsidomide + BCMA Bispecific

**Phase 2 (Single Arm)**  
Cemsidomide + Dex (Post anti-BCMA)

*Potential Accelerated Approval*

**Studies for Registrational Intent**

**Randomized Phase 3**  
Cemsidomide + BCMA Bispecific (1-3 prior lines)

**Randomized Phase 3**  
Cemsidomide/Dex + Anti-CD38 (2-4 prior lines; post anti-BCMA)

**Development Rationale**

Potentially enhances response durability and treatment duration of BCMA bispecific by preventing T-cell exhaustion

Provides post anti-BCMA patients a potentially highly efficacious combo where there are limited proven options

Potential to provide highly refractory patients a treatment option that is tolerable and efficacious where there are limited options

<sup>1</sup> EvaluatePharma (accessed 1/8/25), consulting engagements with Health Advances and Clearview. EU4 = Germany, Italy, France, and Spain. B-cell maturation antigen (BCMA); dexamethasone (Dex); T-Cell engager (TCE); multiple myeloma (MM)

# Cemsidomide Has the Potential to be Developed Across NHL Subtypes and Lines of Treatment

	B-Cell Lymphomas				T-Cell Lymphomas
	DLBCL Diffuse Large B-Cell Lymphoma	FL Follicular Lymphoma	MZL Marginal Zone Lymphoma	MCL Mantle Cell Lymphoma	PTCL Subtypes Peripheral T-Cell Lymphoma
U.S. + EU4 + UK Annual Incidence (2024) <sup>1</sup>	~54,000	~33,000	~12,000	~8,000	~16,000
Lenalidomide FDA Approved <sup>2</sup>	✓	✓	✓	✓	
Lenalidomide in NCCN Guidelines	✓	✓	✓	✓	✓

## Cemsidomide Opportunity

- Lenalidomide is approved across NHL subtypes
- Cemsidomide has the potential to be developed as a **monotherapy in the R/R setting** and in **combination with front-line standard of care regimens**

<sup>1</sup> EvaluatePharma (accessed 1/8/25), American Cancer Society, Leukemia & Lymphoma Society. EU4 = Germany, Italy, France, and Spain.

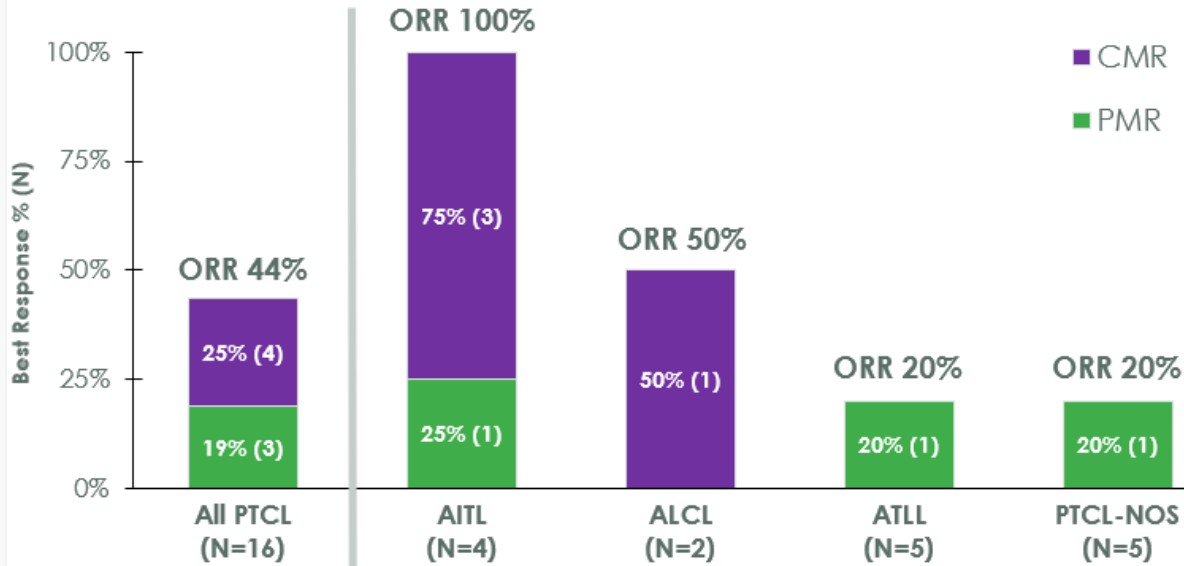
<sup>2</sup> FL, MZL, and MCL FDA approved in the Revlimid (lenalidomide) label and DLBCL approved in the Monjuvi (tafasitamab) label.

U.S. Food and Drug Administration (FDA); National Comprehensive Cancer Network (NCCN); non-Hodgkin's lymphoma (NHL); relapsed/refractory (R/R)

# Cemsidomide Data at ASH Demonstrated a Well-tolerated Profile, as Well as Compelling and Deep Responses in PTCL Patients

## Compelling & Deep Responses Achieved Across PTCL Subtypes

PET-CT-based Assessment of PMR or Better by PTCL Subtype<sup>1</sup> (N=16)



## Well-tolerated With Manageable Incidents of On-target Neutropenia

- 2 DLTs occurred at 100 µg QD (Grade 4 thrombocytopenia and Grade 3 febrile neutropenia)
- TEAEs leading to discontinuation: 9% (2/23)
- 39% (9/23) of patients received G-CSF
  - 3 of 9 patients received G-CSF in Cycle 1

Common (>20% All Grades) TEAEs and Events of Interest*, n (%)	All Grade (N=23)	Grade 3 (N=23)	Grade 4 (N=23)
<b>Infections</b>	15 (65)	4 (17)	2 (9)
Upper respiratory tract infection	4 (17)	0	0
Sepsis	1 (4)	0	1 (4)
Bacteremia	1 (4)	0	1 (4)
Pneumonia	2 (9)	2 (9)	0
<b>Neutropenia</b>	11 (48)	4 (17)	7 (30)
<b>Fatigue</b>	11 (48)	1 (4)	0
<b>Cough</b>	7 (30)	0	0
<b>Anemia</b>	6 (26)	4 (17)	0
<b>Peripheral edema</b>	5 (22)	0	0
<b>Febrile neutropenia*</b>	4 (17)	4 (17)	0
<b>Thrombocytopenia*</b>	4 (17)	1 (4)	2 (9)
<b>Maculopapular rash*</b>	3 (13)	2 (9)	0

One patient experienced a Grade 5 AE (hip fracture resulting in transfer to hospice)

**Profile supports cemsidomide’s development as a monotherapy in relapsed refractory settings and potentially as a combination in NHL subtypes across treatment lines**

<sup>1</sup> Investigator assessed response; 2 patients were evaluated based on CT scan and were PD but not evaluable based on PET-CT, both patients are included as PMD for PET-CT based assessment; 2 additional subjects that came off study prior to follow up scans were not considered efficacy evaluable. \*Events of Interest Angioimmunoblastic T-cell lymphoma (AITL); anaplastic large cell lymphoma (ALCL); adult T-cell lymphoma (ATLL); complete metabolic response (CMR); overall response rate (ORR); partial metabolic response (PMR); peripheral T-cell lymphoma (PTCL); peripheral T-cell lymphoma not-otherwise specified (PTCL-NOS); adverse event (AE); dose limiting toxicities (DLTs); granulocyte colony-stimulating factor (G-CSF); once daily (QD); treatment emergent adverse events (TEAEs)

Source: ASH 2024; C4T data as of 10/11/2024 (<https://ir.c4therapeutics.com/static-files/32ae4fdb-d4d9-4a17-a77d-83289c66e91f>)

# Cemsidomide NHL Data Supports Further Development in PTCL, Which Provides the Fastest Path to Market

## INITIAL CEMSIDOMIDE DEVELOPMENT PATH IN PTCL

U.S. + EU4 + UK Addressable Patients (2024) <sup>1</sup>	~16,000	≤12,000
Treatment Line	1L	2L+

Study Currently Being Enabled

**Phase 2 (Single Arm)**  
**Cemsidomide Monotherapy**  
 (2L+ R/R PTCL)

*Potential Accelerated Approval*

Study for Registrational Intent

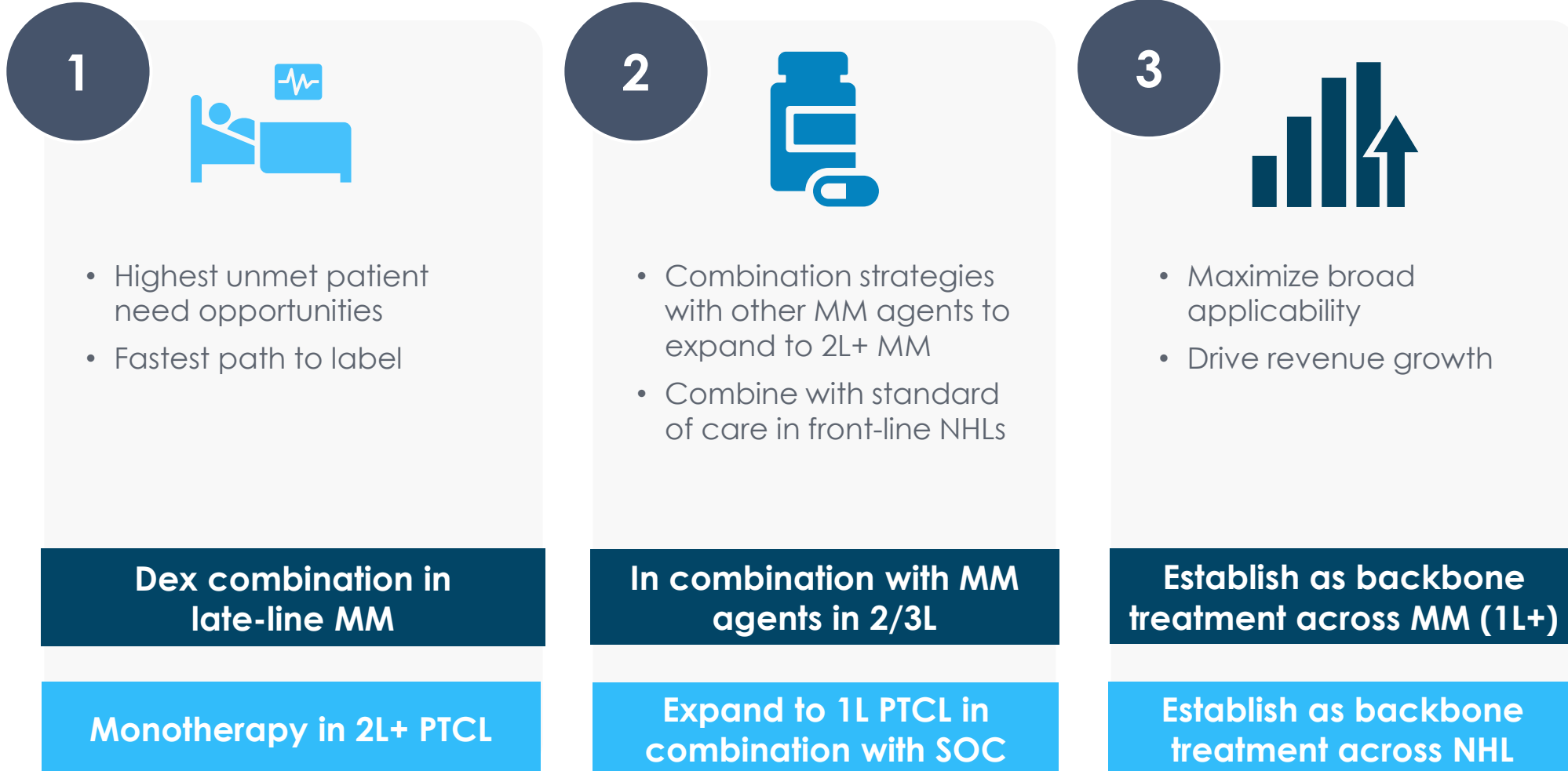
**Randomized Phase 3**  
**Cemsidomide + SOC<sup>2</sup>**  
 (treatment naïve)

<b>Development Rationale</b>	Potentially enhance response durability and decrease chemotherapy use, thus providing a more tolerable and durable option	Potentially provides R/R patients a treatment option that is tolerable and efficacious where there are limited options
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<sup>1</sup> EvaluatePharma, ACS, consulting engagements with Health Advances and Clearview.

<sup>2</sup> Standard of care (SOC) for 1L patients with CD30+ disease is brentuximab vedotin +/- chemotherapy and for CD30- patients it is the CHOP chemotherapy regimen (cyclophosphamide, doxorubicin, vincristine, prednisone) Germany, Italy, France, and Spain (EU4); peripheral t-cell lymphoma (PTCL); relapsed refractory (R/R); standard of care (SOC)

# Cemsidomide Has a Strategic Path to Become a Potential Backbone Therapy for MM and NHL Across Various Lines of Treatment



Multiple myeloma (MM); non-Hodgkin's lymphoma (NHL); peripheral T-cell lymphoma (PTCL); standard of care (SOC)

# CFT1946

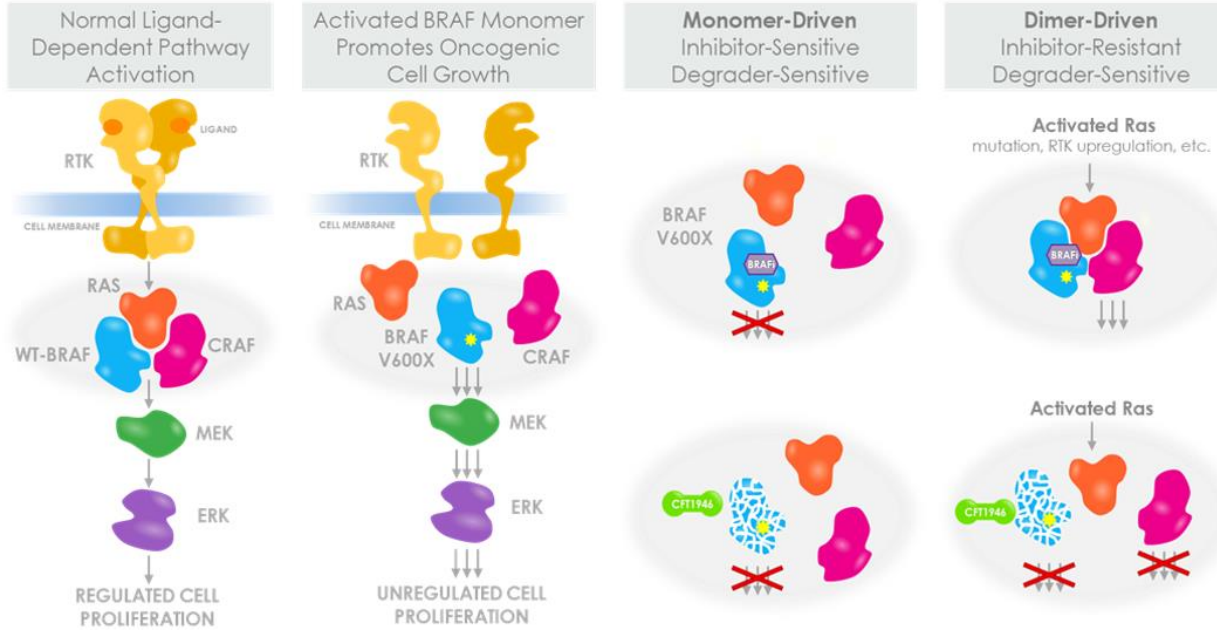
## BRAF Mutant V600 Degradator

Colorectal Cancer, Melanoma &  
Other BRAF Mutant Solid Tumors



# CFT1946 Is an Oral, Potent Degradator of BRAF V600 Mutants With Potential to Improve Outcomes for Patients

## Degrader Benefit in BRAF V600 Mutant Monomer and Dimer-dependent Diseases



### Causes of Dimer-Driven Resistance

**Melanoma:** Acquired NRAS mutation, BRAF V600 mutant amplification, BRAF V600 mutant splice variant

**CRC:** EGFR-mediated pathway reactivation

### Current Approved BRAF Inhibitors Have Limitations:

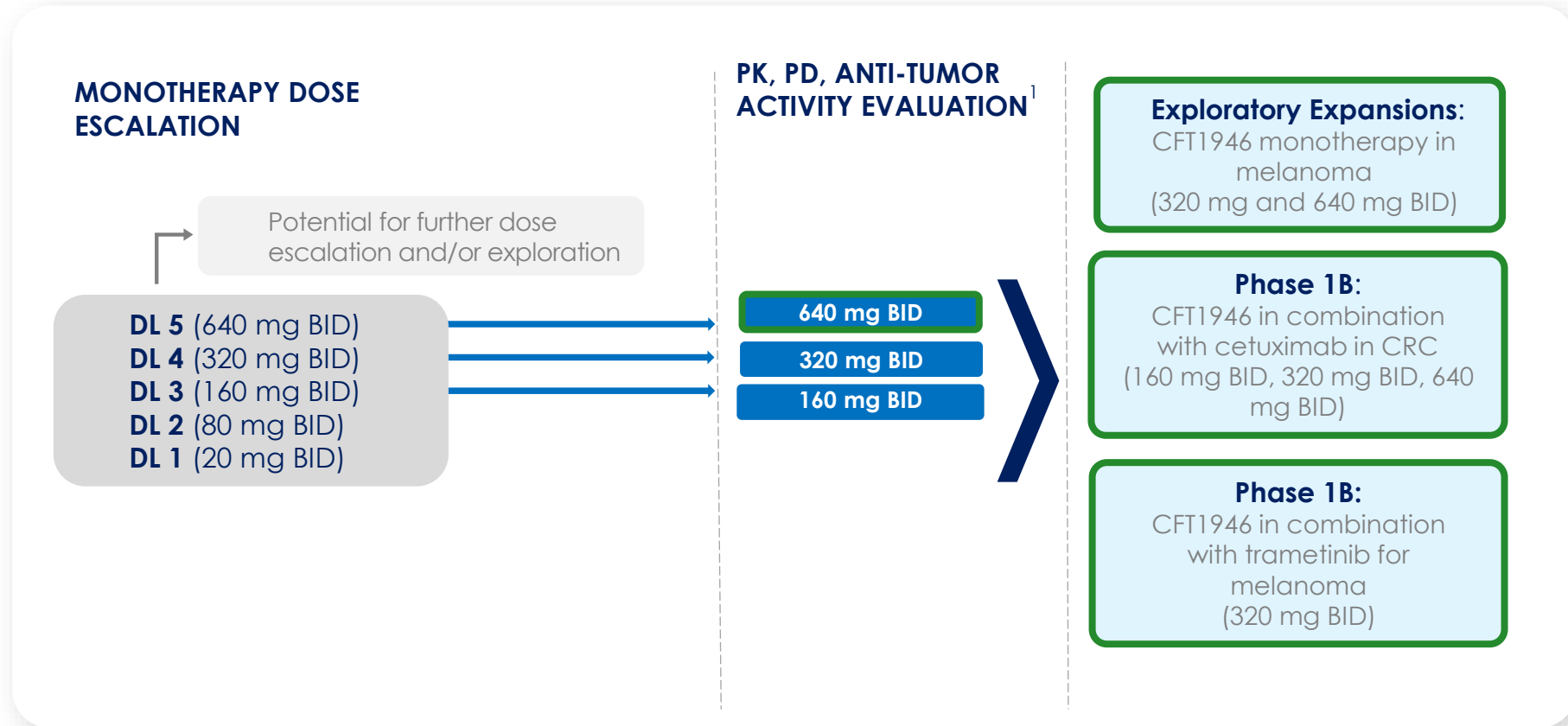
- Resistance mechanisms lead to **limited duration of response**
- Toxicities associated with inhibition of wild-type BRAF **limit tolerability**

### Potential Degradator Advantages of CFT1946:

- ✓ Enables deep elimination of mutant BRAF signaling to avoid resistance mechanisms and enable a longer duration of response
- ✓ Spares wild-type BRAF<sup>1</sup>, likely avoiding AEs associated with inhibition of wild-type BRAF
- ✓ Ability to cross the blood-brain barrier and highlight the potential for drug delivery to CNS tumors, with  $K_{p_{U,U}}$  values ranging from 0.34 to 0.88

<sup>1</sup>Kreger B et al. Abstract 1658, AACR 2024 Adverse event (AE); colorectal cancer (CRC); central nervous system (CNS); mitogen-activated protein kinase (MAPK); progression free survival (PFS); BRAF inhibitor (BRAFi).

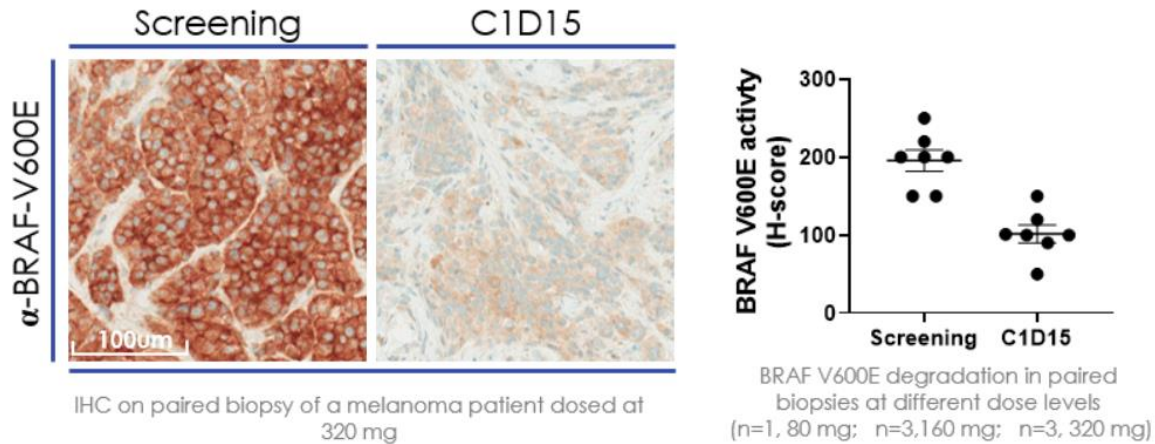
# CFT1946 Phase 1/2 Trial Continues to Progress Across BRAF V600 Mutant Driven Solid Tumors



<sup>1</sup>Evaluating additional patients for pharmacodynamic assessment via pre- and post-drug exposure biopsies  
Colorectal cancer (CRC); dose Level (DL); twice daily (BID); pharmacokinetic (PK); pharmacodynamic (PD)

# CFT1946 Monotherapy Phase 1 Data Presented at ESMO Demonstrate Proof of Mechanism and Provide Early Evidence of Proof of Degradation Concept

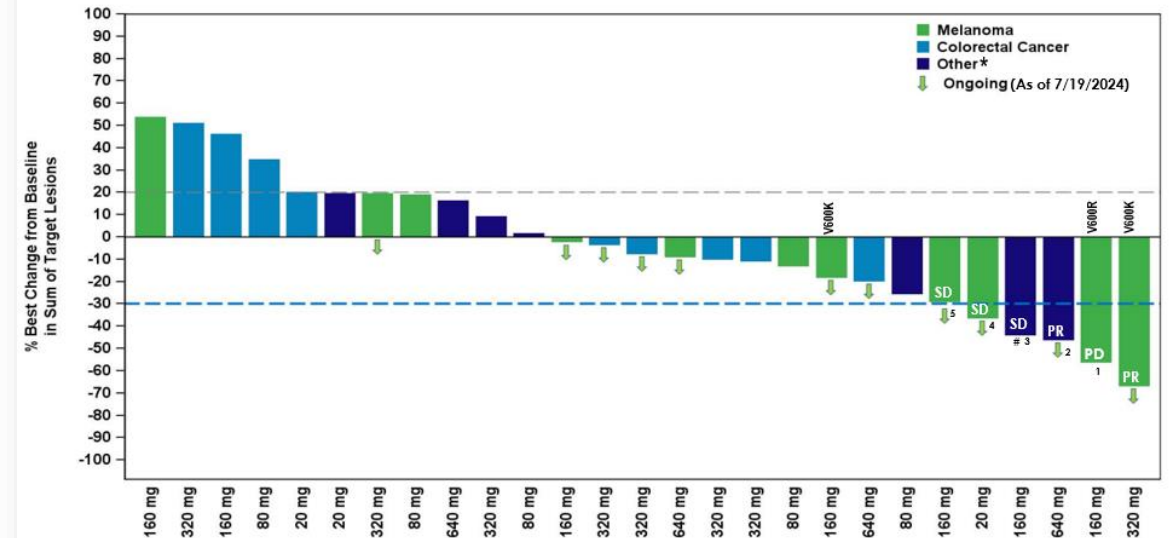
## BRAF V600E Degradation Determined by H-score of Paired Biopsies from Different Tumor Types



## Well-tolerated Monotherapy Safety Profile, Consistent with BRAF V600 Mutant Selectivity Design of CFT1946

- No DLTS
- Majority of TEAEs observed were mild to moderate
- No treatment-related SAEs

## Early Evidence of Monotherapy Anti-tumor Activity in Patients who Progressed After Treatment with BRAF inhibitors



\*Other tumor types include cholangiocarcinoma, non-small cell lung cancer, pancreatic carcinoma, and small intestine cancer; BRAF V600 mutation is V600E unless otherwise specified; #This patient did not receive prior BRAF inhibitor therapy, all other patients received prior BRAF inhibitor therapy. Dotted lines represent partial response (-30%, blue line) and progressive disease (20%, gray line) per RECIST v1.1.  
 1 Patient on 160 mg BID had 56.2% reduction on target lesion, progression on non-target lesion and a new lesion, hence assessed as PD for overall response; 2 Patient on 640 mg BID had PR confirmed after data cut off, and as of ESMO Congress (9/13/2024); 3 Patient on 160 mg BID had PD following first PR (-43.9%), hence assessed as SD for overall response; 4 Patient on 20 mg BID had unconfirmed PR, hence assessed as SD for overall response; 5 Patient on 160 mg BID had -29% reduction on target lesion, hence assessed as SD

**Multiple data readouts expected in 2H 2025 to define and enable next phase of development**

Serious adverse events (SAEs); dose limiting toxicities (DLTs); treatment-related adverse events (TRAEs); treatment-emergent adverse events (TEAEs)  
 Source: ESMO Congress 2024; C4T data as of 7/19/2024 (<https://ir.c4therapeutics.com/static-files/b648e3ae-c2db-4f5e-9d47-324b0c8a2b2b>)

# Potential for Multiple Paths to a Label for CFT1946 to Address High Unmet Needs in BRAF V600 Mutant Melanoma and CRC

## Data-Driven Decisions to Inform Next Steps



### Melanoma (BRAF V600 Mutation Rate: ~35%<sup>1</sup>)



*Monotherapy melanoma data will inform fastest path to label*

#### Development Rationale:

- Improvement on durability due to resistance mechanisms that emerge with BRAF inhibitors
- Large patient population (**~66,000 BRAF V600 mutant melanoma patients**)<sup>3</sup>



### Colorectal Cancer (BRAF V600 Mutation Rate: 5-10%<sup>2</sup>)

+ cetuximab CRC data will inform fastest path to label:



#### Development Rationale:

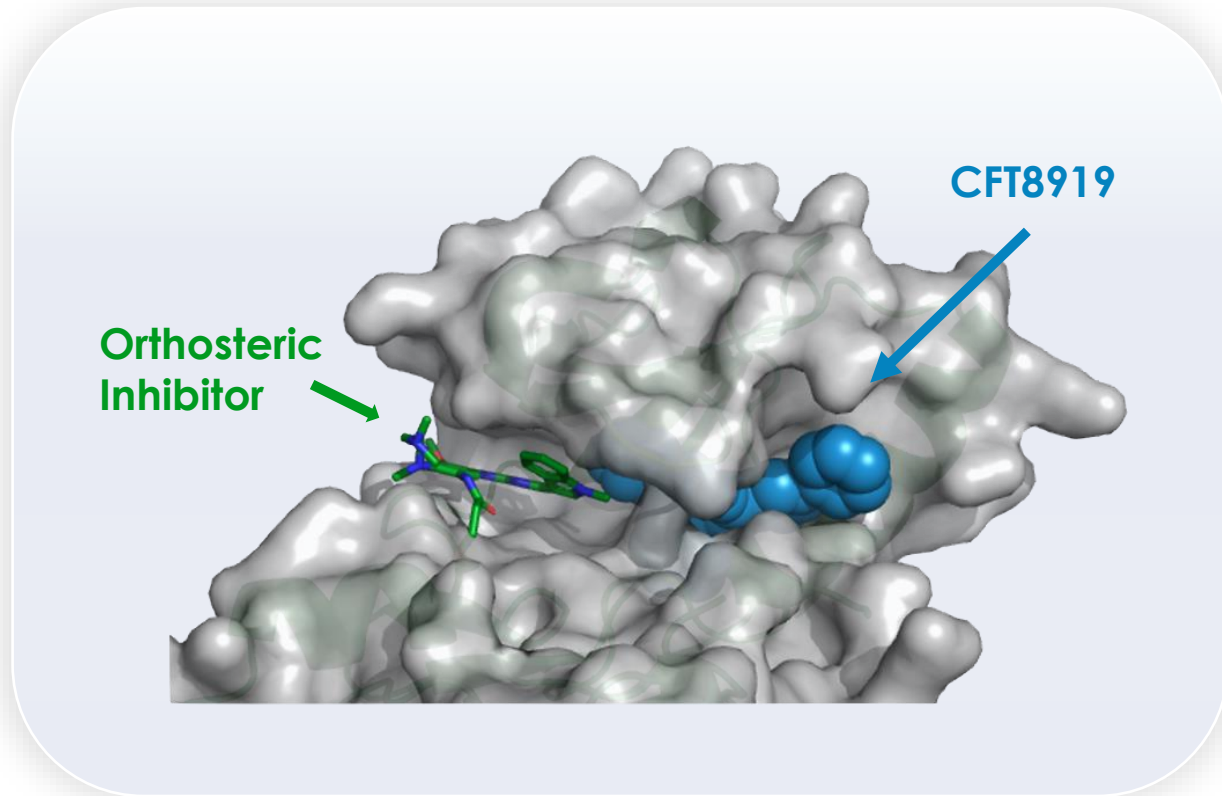
- Limited efficacious and tolerable treatment options for a large patient population (**~33,000 patients BRAF V600 mutant CRC patients**)<sup>3</sup>
- Patients typically treated with chemotherapy

<sup>1</sup> Owsley 2021 Exp Biol Med. <sup>2</sup> Paik 2011 J Clin Oncol. <sup>3</sup> 2024 EvaluatePharma (accessed 1/8/25); comprises the U.S. + EU4 + UK population. Germany, Italy, France, and Spain (EU4); Mek inhibitor (MEKi) BRAF inhibitor (BRAFi); colorectal cancer (CRC).

CFT8919  
EGFR L858R Degradator  
Non-Small Cell Lung Cancer



# CFT8919 Is a Potent, Oral, Allosteric, Mutant-selective Degradator of EGFR L858R With Potential to Improve Outcomes for NSCLC Patients



## Current Approved EGFR Inhibitors Have Limitations:

- Patients **become refractory due to secondary mutations**
- NSCLC patients with **L858R have inferior clinical outcomes**
- Toxicities associated with inhibition of wild-type EGFR **limit tolerability**

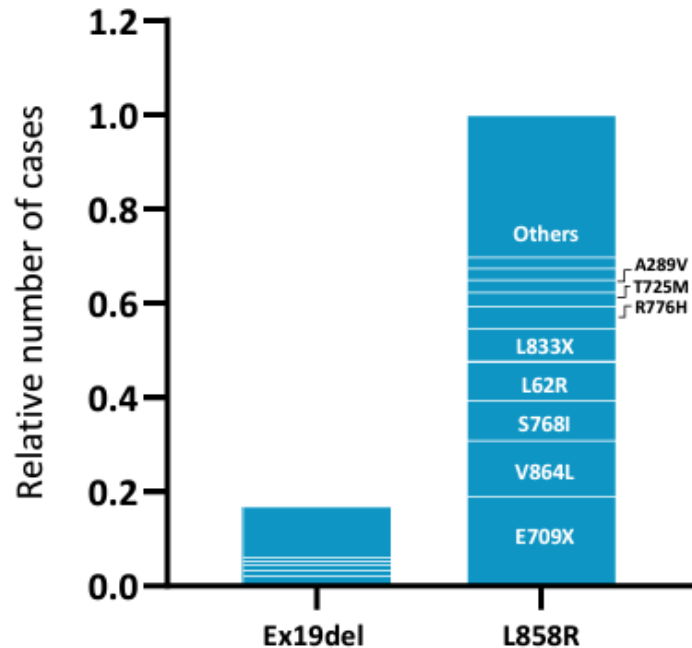
## Potential Degradator Advantages of CFT8919:

- ✓ CFT8919 exploits an allosteric binding site created by the L858R mutation, thereby avoiding resistance mutations to the orthosteric site
- ✓ Potent and selective against L858R regardless of secondary mutations with potential for more durable activity in this setting
- ✓ Does not hit wild-type, potentially resulting in better tolerability

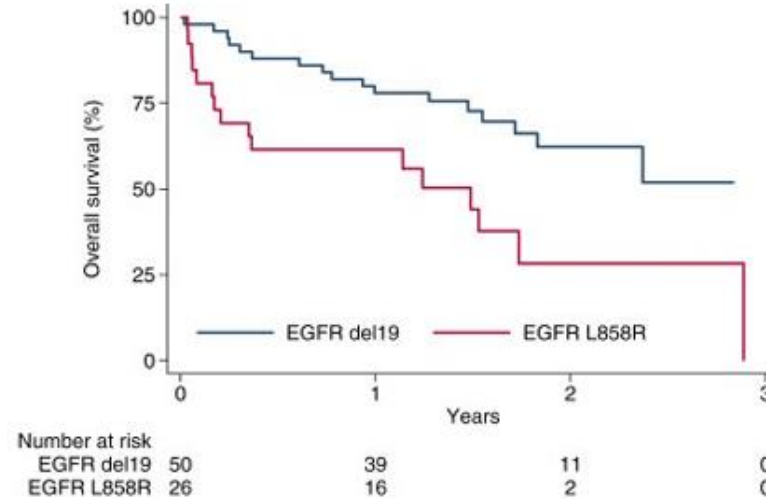


# CFT8919 Binds to Allosteric Site, Avoiding Impact of L858R Non-classical Co-mutations in the Orthosteric Binding Pocket

**EGFR-L858R Tumors More Frequently Co-express Non-classical EGFR Mutations Before Exposure to EGFR TKI<sup>1</sup>**



**Patients with L858R Do Less Well on Osimertinib Therapy vs Ex19del**



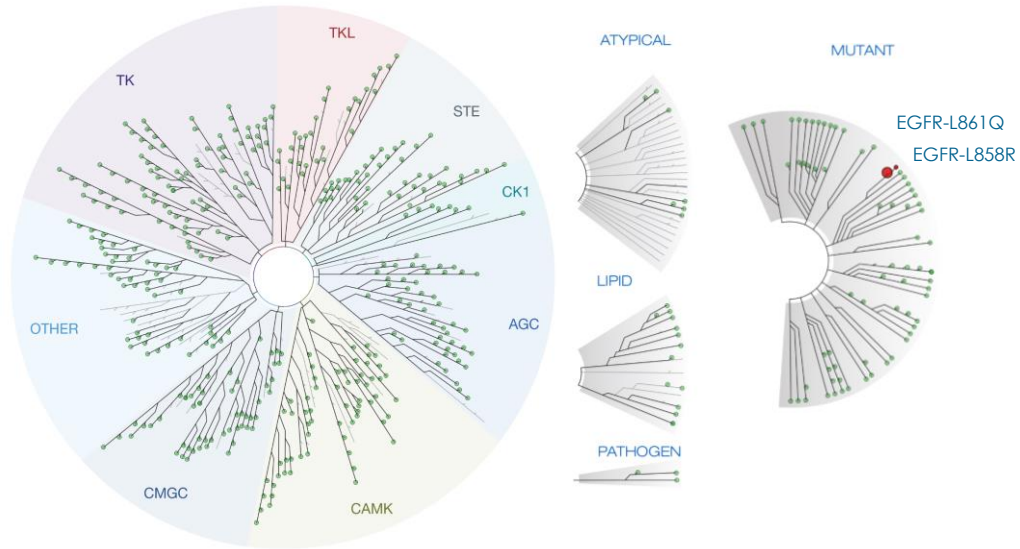
Overall survival by type of mutation in patients with Stage IV EGFR mutated NSCLC and brain metastasis who received first-line treatment with osimertinib

CFT8919 binds to the allosteric site, potentially avoiding the impact of non-classical co-mutations with L858R, where inhibitors demonstrate lower PFS in this patient population than those with EXON 19 deletion

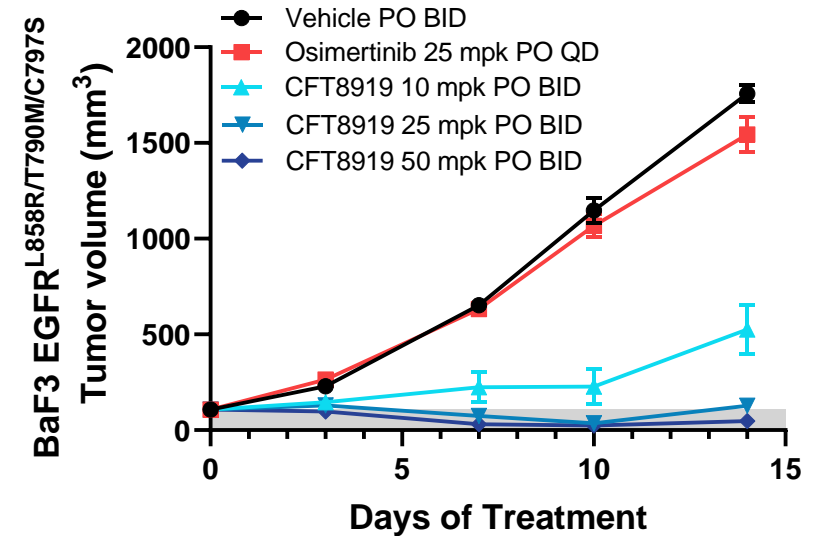
Sources: 1. From Black Diamond's analyses of 94,939 sequencing reports from treatment naive NSCLC (Guardant Health) presented at AACR 2024 ([https://blackdiamondtherapeutics.com/assets/files/AACR\\_2024\\_BDTX-1535\\_FINAL\\_Presentation\\_20240405.pdf](https://blackdiamondtherapeutics.com/assets/files/AACR_2024_BDTX-1535_FINAL_Presentation_20240405.pdf)) 2. Gitenbeek, et al. 2023 Progression free survival (PFS)

# CFT8919 Is Selective for EGFR L858R and Active in a Setting of Osimertinib Resistance in Preclinical Models

## Specific for EGFR Exon 21 Mutants



## Active in Setting of EGFR C797S

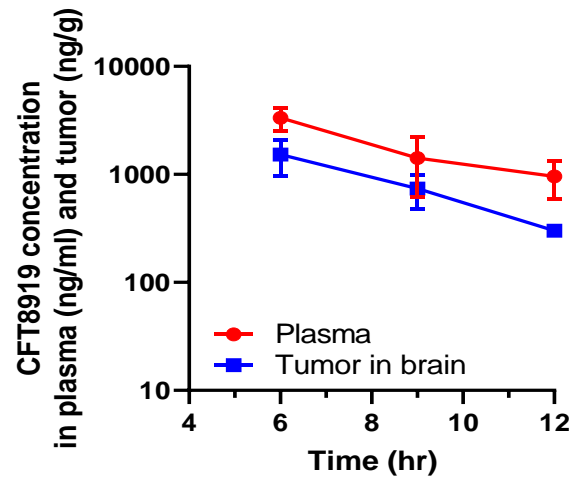


Source: C4T data on file; Presented at Keystone Symposium 2021 (<https://c4therapeutics.com/wp-content/uploads/Preclinical-Evaluation-of-CFT8919-as-a-Mutant-Selective-Degrader-of-EGFR-with-L858R-Activating-Mutations-for-the-Treatment-of-Non-Small-Cell-Lung-Cancer.pdf>)

Investigational new drug application (IND)

# CFT8919 Demonstrates Activity in Brain Metastasis Model

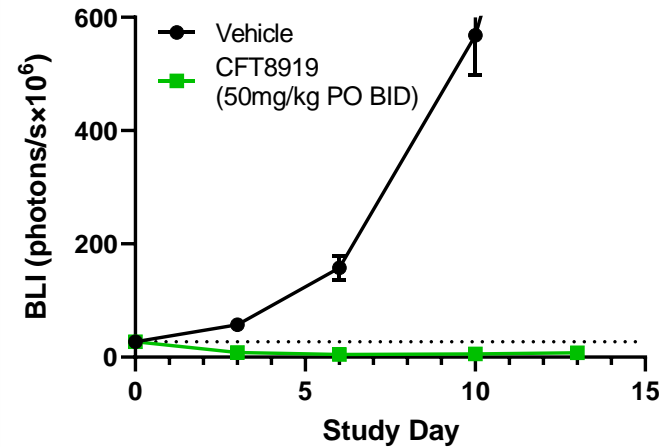
## Mean Plasma & Tumor Concentration



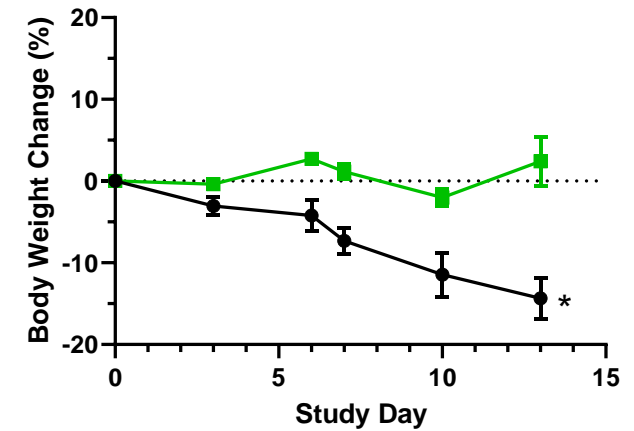
Plasma clearance  $t_{1/2} = 3.1$  hrs

50 mg/kg single dose PO

## In vivo Efficacy



## In vivo Body Weight Change



\*Body weight loss due to tumor burden

Source: C4T data on file; presented at TPD Summit 2021 ([https://c4therapeutics.com/wp-content/uploads/C4\\_CFT8919\\_TPD\\_Summit\\_Presentation.pdf](https://c4therapeutics.com/wp-content/uploads/C4_CFT8919_TPD_Summit_Presentation.pdf))  
By mouth (PO); twice daily (BID)

# CFT8919 Has the Potential to Address Multiple Opportunities with High Unmet Needs

## CFT8919's Fastest Path to Market Is in 2L+ With Potential to Expand Into Front-Line

### 2L+

#### Development Rationale:

- Fast path to market
- Lack of therapies after patients relapse with secondary mutation (i.e., C797S)

### Front-line

#### Development Rationale:

- Large patient opportunity
- Potential to increase responses and durability in L858R patients

**Dose escalation in Greater China is advancing; C4T to utilize data to inform ex-China clinical development**

### 2024 Annual Incidence of EGFR L858R Mutated NSCLC<sup>1</sup>:



- **U.S.:** ~17,000
- **China:** ~189,000
- **EU4 + UK:** ~13,000

<sup>1</sup> EvaluatePharma (accessed on 1/10/25), consulting engagements with Health Advances and Clearview. Germany, Italy, France, and Spain (EU4)

# Multiple 2025 Inflection Points to Unlock Value Across Entire Portfolio

## VALUE DRIVERS

## KEY CATALYSTS

**Cemsidomide**  
IKZF1/3

Further development in multiple myeloma and non-Hodgkin's lymphoma positions cemsidomide to potentially be best-in-class IKZF1/3 degrader

**CFT1946**  
BRAF V600 Mutant

Phase 1 data updates to further validate initial anti-tumor activity and safety profile in melanoma and colorectal cancer

**CFT8919**  
EGFR L858R

Phase 1 data from Greater China clinical trial to inform U.S. and rest-of-world development plans

**TORPEDO**  
Platform

Develop orally bioavailable degraders in oncology and non-oncology targets for internal research and collaborations

**C4T** is on a path to become a **fully integrated biotechnology** company focused on **orally bioavailable degraders**