



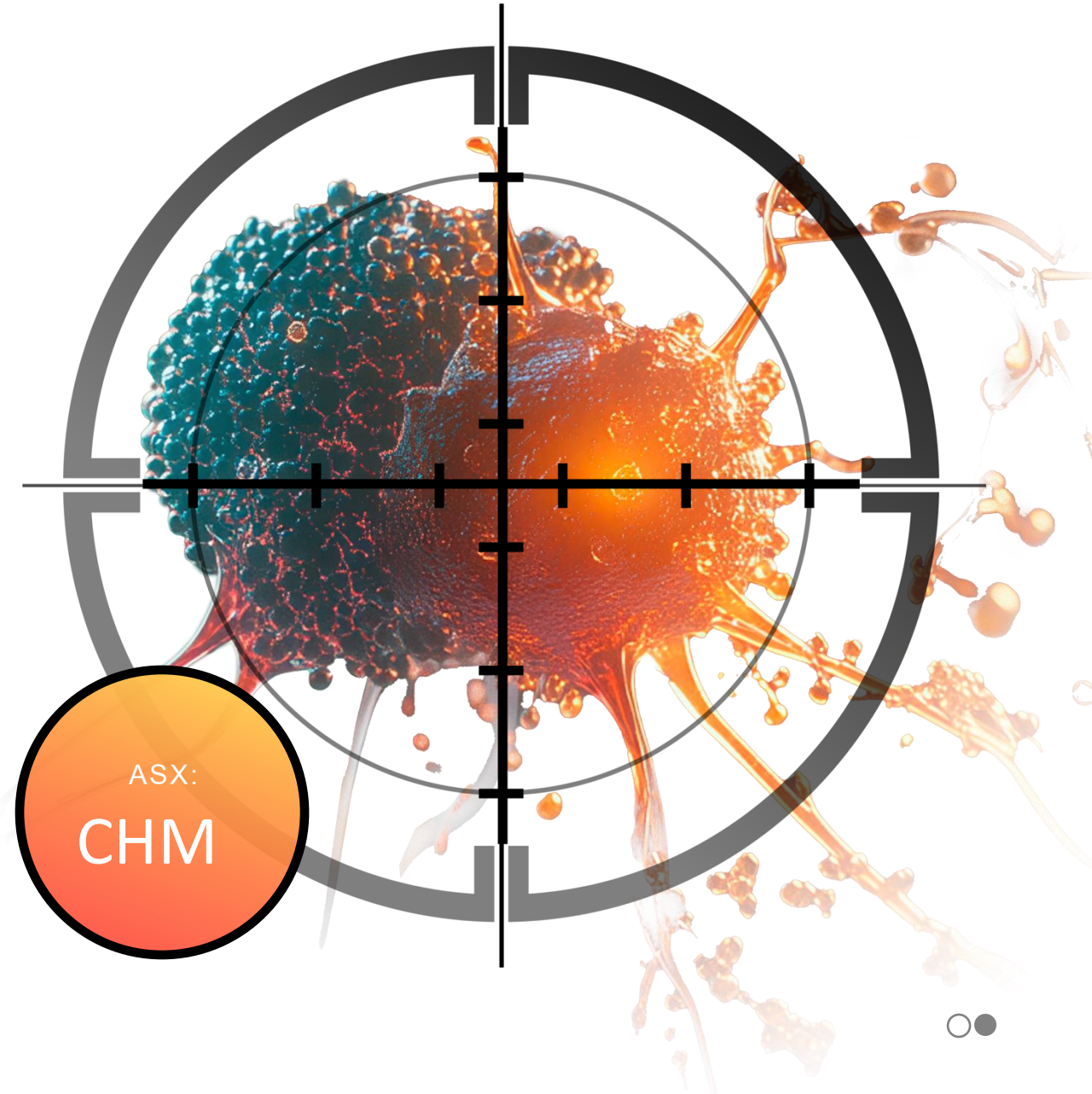
CHIMERIC THERAPEUTICS

20
25

CHIMERIC

PIONEERS IN CELL THERAPY

AGM 2025





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CHM: BROAD PORTFOLIO

3 Novel cell therapies; 4 Clinical Trials





CHM CDH17 PROGRESS AS AT NOV 24TH, 2025

FIVE ONGOING STABLE DISEASE

Dose Level	Status	Notes
1 50m	Completed	4 pts treated 1 pt remains on study 12months ongoing Stable Disease
2 150m	Recruitment Ongoing	5 patients treated 4 pts achieved ongoing Stable Disease 1 pt awaiting results
3 450m	Recruitment underway	1 pt awaiting dosing assignment
Manufacturing	FDA GMP compliant	10 out of 10 successful runs

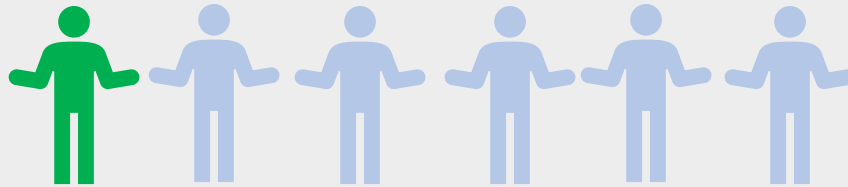
FDA FASTRACK GRANTED



CHM CORE-NK FRONTLINE PHASE 1 STUDY RESULTS

Four Completes Responses, 1 Partial Response

Dose escalation
completed



Relapse Refractory Patients
1 Complete Response

First line
Dosing
Underway



FrontLine Patients with no prior treatment
3 Complete Responses
1 Partial Response

Complete Response means that all visible or measurable signs of cancer have disappeared after treatment

Partial Response means the cancer has reduced by at least 50% but not entirely disappeared in response to treatment



SUMMARY

CHM CDH17

CHM 2101

9 patients treated
10/10 manufacturing runs
Tumour shrinkage observed
Manageable safety profile
No off-target effects

10

CHM CORE-NK

'OFF THE SHELF' NK

Investigator Sponsored Trial at MD ANDERSON
Dose Escalation completed
First Line Dosing Underway – 7 dosed up to 20 patients
No safety issues to date



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PIONEERS IN CELL THERAPY

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ASX:
CHM

