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Fostrox – a smart chemotherapy with a unique, liver targeted & tumor selective treatment of HCC

Nucleotide prodrug, enabling oral administration & liver targeting >100-fold liver targeted exposure vs traditional chemotherapy¹

Promising signals of clinical benefit supports accelerated approval path

- First-in-class with OD designation in EU & US
- Fostrox + Lenvima provides additional clinical
 benefit to Lenvima alone
- Pivotal phase IIb with Accelerated Approval intent 2027/2028
- First-to-market opportunity in target population with annual market value of ~\$2.4bn in 2028*

Medivir – Oncology pipeline with in-house developed lead program in phase II & 3 out-licensed oncology programs

Fostrox



- Smart chemotherapy delivering cell-killing activity selectively to the liver tumor
- Orphan Drug Designation granted in USA and EU
- Promising clinical benefit with first-to market opportunity 2027

Partnering (programs

Out-licensed oncology programs with potential upside without further investment

- 3 out-licensed oncology programs; ongoing discussions for additional out-license
- Birinapant (IGM Biosciences) currently in phase 1 dose escalation with IGM-8444
- TNG348 (Tango Therapeutics) to enter phase 1 in H1 2024

Fostrox initial focus in 2L HCC where no treatments are approved and expected clinical benefit is low

Advanced stage HCC Treatment Algorithm

Slide 5

1L systemic therapy

Immunotherapy combination

- Only ~30% of patients respond to treatment¹
- Estimated time to progression ~6.5 months¹

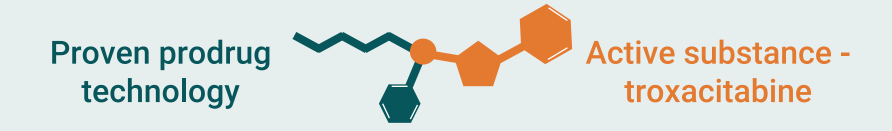
2L systemic therapy

No approved treatments – off-label Lenvima preferred

- Only ~5-10% of patients respond to treatment²
- Estimated time to progression ~3.5 months²
- Fostrox + Lenvima, the only novel combination in development



Fostrox – liver targeted, smart chemotherapy



- 1. Oral administration
- 2. Targeted (>100-fold) liver exposure vs IV chemotherapy¹
 - 3. Selective DNA damage in tumor vs normal liver tissue

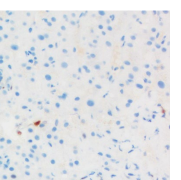


Fostrox – Patient biopsies confirming selective DNA damage & cell death in tumor cells while sparing normal liver tissue

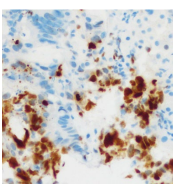
Tumor selective induction of DNA-damage¹

Cytotoxic in tumor tissue but not in normal liver tissue²

Normal liver tissue

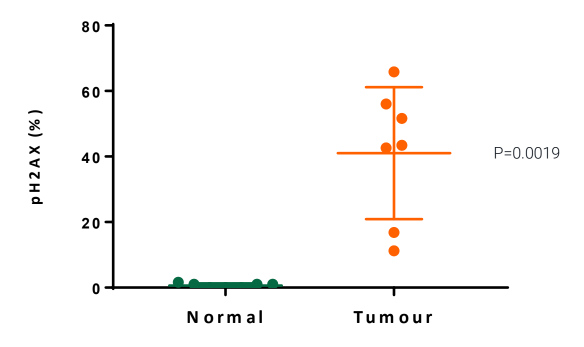


Tumor tissue



Fostrox-induced DNA-damage indicated by pH2AX immuno-histochemistry (IHC) staining of liver biopsy from phase 1b monotherapy

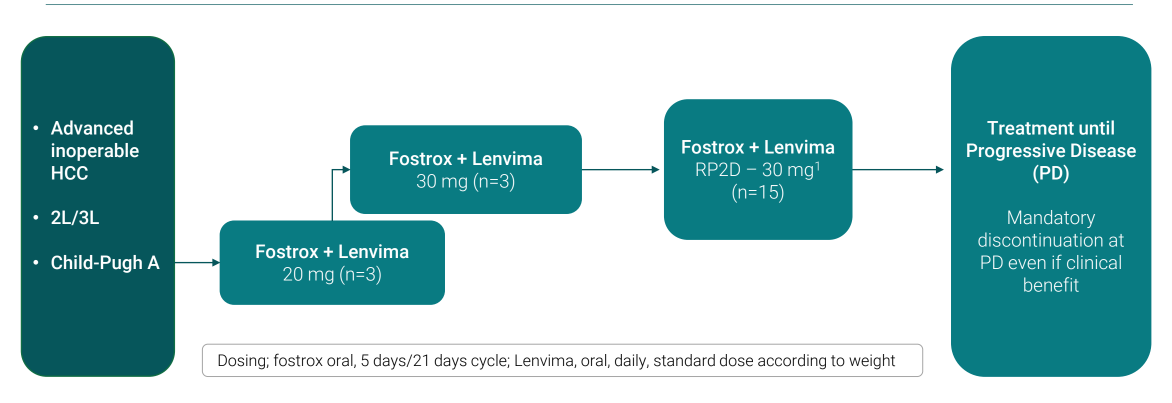
DNA-damage in normal liver vs tumour





Phase 1b/2a study fully recruited with >50% of patients still on treatment

Fostrox + Lenvima phase 1b/2a dose expansion study – 21 patients dosed



¹Maximal tolerated dose not reached with no DLTs reported. 30 mg selected with a focus on optimal dose ensuring balance between efficacy and tolerability

Previous studies in 2nd line HCC confirm difficult-to-treat population

RECIST 1.1	Efficacy Benchmarks – previous 2 nd line studies ¹
Overall response rate (ORR)	~10%
Clinical Benefit Rate (CBR/DCR)	~60%
Median Progression-free Survival/Time to Progression	~3.5 months

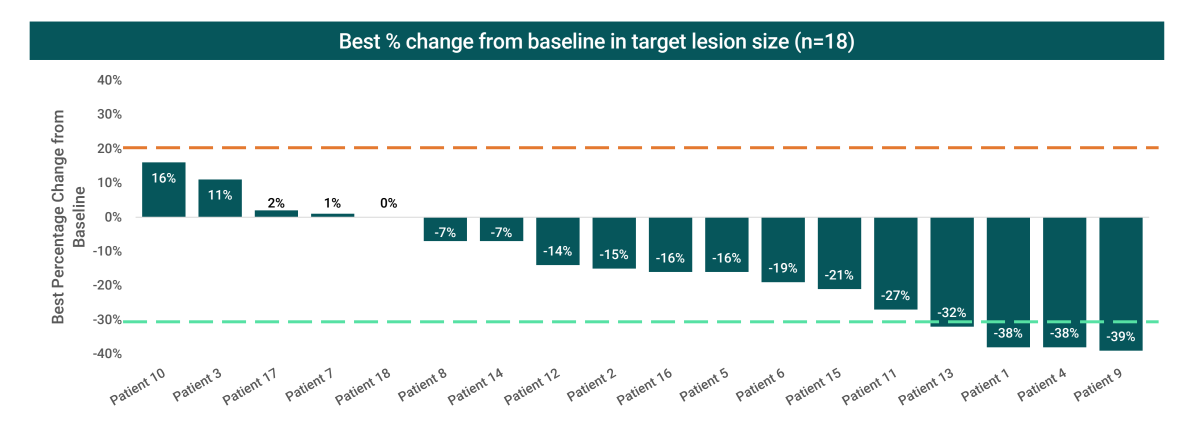
^{*}Preliminary results from Investigator review (18 patients with a minimum of 12 weeks follow-up)

Fostrox + Lenvima compares very favourably with benchmarks in 2nd line HCC

RECIST 1.1	Efficacy Benchmarks – previous 2 nd line studies ¹	Fostrox + Lenvima* (n=18)	
Overall response rate (ORR)	~10%	22%	
Clinical Benefit Rate (CBR/DCR)	~60%	78%	
Median Progression-free Survival/Time to Progression	~3.5 months	4.9 months	

^{*}Preliminary results from Investigator review (18 patients with a minimum of 12 weeks follow-up)

22% Overall Response Rate (ORR); more than two third of patients with tumor reduction* (Investigator review RECIST 1.1)



3 additional patients; all with ≥6 weeks follow-up & stable disease at 1st evaluation

Lenvima monotherapy data in 2nd line HCC confirms significant unmet medical need

RECIST 1.1	Lenvima ¹ (n=12) Independent & investigator review	Fostrox + Lenvima ² (n=18) Investigator review
ORR	8-17%	
Clinical Benefit Rate (at 12 weeks)	58%*	
Median Progression-free Survival/Time to Progression	2.8-4.1 months	
Median Treatment Duration	3.5 months	

^{*} Data only reported as mRECIST



Fostrox + Lenvima compares very favourably with benchmarks in 2nd line HCC

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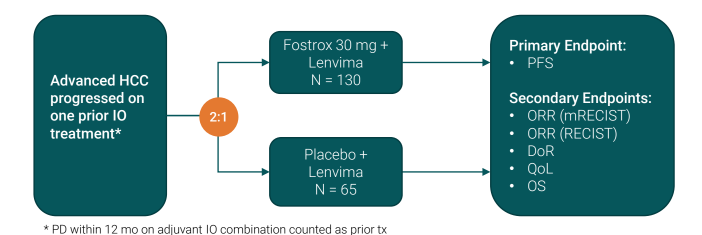
Fostrox + Lenvima study with comparable tolerability, no new safety events vs Lenvima study

	Lenvima ¹ (n=12)	Fostrox + Lenvima ² (n=18)
≥ Grade 3 AEs	67%	61%
Dose modifications Lenvima	92%	50%
Discontinuations due to AEs	25%	17%



Pivotal Phase IIb; randomized design with PFS as primary endpoint to enable accelerated approval 2027

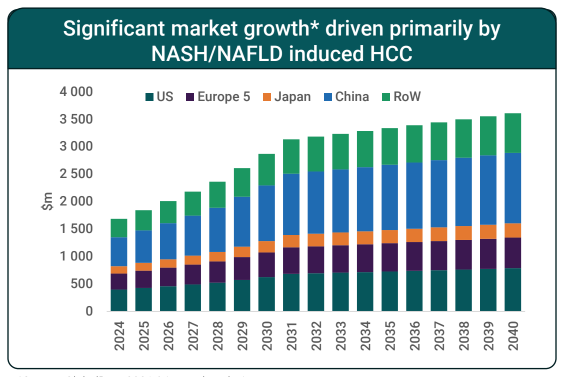
Phase IIb: randomized, double-blind study design

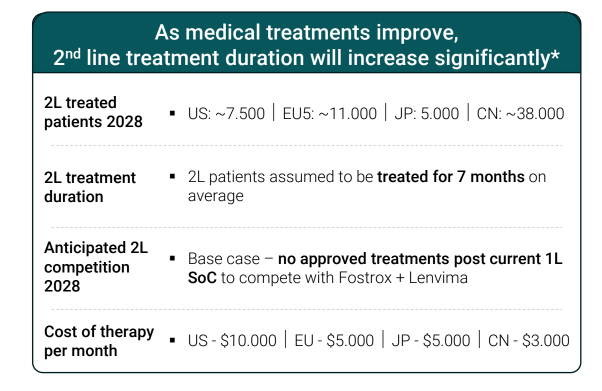


Key factors supporting accelerated approval process

- Serious, orphan disease with high unmet medical need
- ✓ Promising clinical benefit & safety profile
- Randomized study design with PFS as primary endpoint
- ✓ Appropriate patient safety database

First-to-market opportunity for fostrox in 2nd line HCC market worth \$2.4bn annually by 2028





Strategic evolution – Earlier treatment lines in HCC provides additional market opportunity of >\$5bn

^{*}Source: GlobalData 2021 & internal analysis

Fostrox – a smart chemotherapy with a unique, liver targeted & tumor selective treatment of HCC, potential for accelerated approval 27/28



Fostrox + Lenvima shows consistently improved efficacy compared with Lenvima alone



Continued development for fostrox + Lenvima in 2nd line HCC with Accelerated Approval intent 2027/2028



2nd line HCC post Tecentriq® + Avastin® lacks approved treatments & is a market valued at ~\$2.5bn annually

Key reasons underpinning Rights Issue



Keep maximum speed and momentum in development program for fostrox



Patients in ongoing fostrox + Lenvima study staying longer on treatment and data has continued to improve with increased maturity



Improved clinical benefit supports raised ambition & plan to enable accelerated approval as early as 2027, which will require accelerating critical activities with regards to regulatory interactions, clinical preparations and CMC

Thank You! **MEDIVIR** Slide 19

Fostrox + Lenvima combination uniquely targets key needs in 2nd line HCC

Ongoing studies in 2 nd line HCC post Tecentriq + Avastin			
	Fostrox + Lenvima	TKI monotherapy	IO combinations
Different mechanism of action than 1st line	✓	✓	
Combination treatment with potential for synergistic activity	✓		✓
Targeting tumor locally in the liver to minimize side effects	✓		