



H.C WAINWRIGHT BIOCONNECT CONFERENCE

JANUARY 10-13, 2022

MEDIVIR

Today's presenter

Interim CEO,
Chief Financial Officer



Magnus Christensen

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Table of content

1. Executive summary
2. MIV-818
3. Clinical portfolio and partnerships

Medivir and recent events

Clinical portfolio

- Lead asset - MIV-818 – a prodrug that selectively targets cancer in the liver, currently in clinical phase 1/2a development
- Three clinical stage assets; one fully financed by partner and two open for partnering/outlicensing

Recent events

- Supporting clinical data from the MIV-818 phase 1b monotherapy presented at ESMO
- Jens Lindberg appointed new CEO of Medivir
- Birinapant clinical study initiated by IGM Biosciences – milestone MUSD 1.5
- First patient dosed in the MIV-818 combination study

Founded: 1988
Listed: Nasdaq OMX
Location: Stockholm
Cash position: SEK 226M ¹⁾
Market Cap: SEK 624M ²⁾
FTE: 9

1) Q3 report
2) 2021-12-30

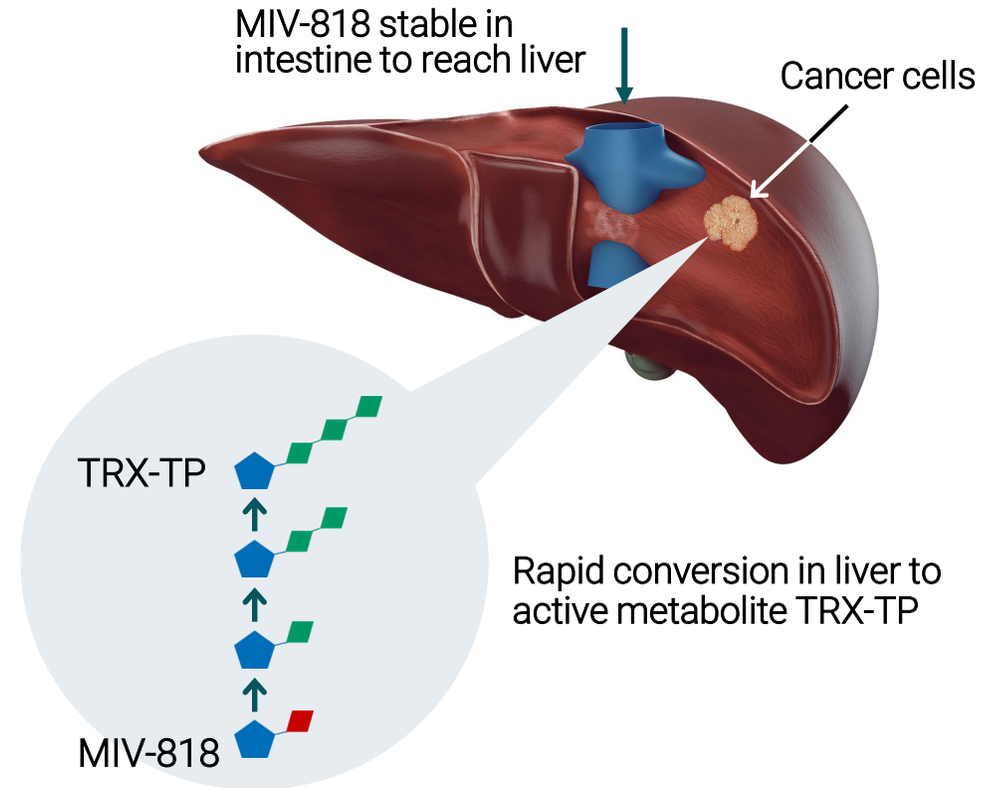
Lead asset – MIV-818 treating liver cancer

- Orphan drug designation by EMA and FDA for the treatment of hepatocellular carcinoma (HCC)
- Despite recent developments most patients with advanced liver cancer have a very poor prognosis
- Medivir have developed MIV-818, a prodrug that selectively targets cancer in the liver
- Positive data from phase 1b monotherapy, demonstrating Proof-Of-Concept, presented at ESMO in September
- First patient dosed in the MIV-818 phase 1b/2a combination study

MIV-818 – for the treatment of liver cancer

MIV-818 – Introduction

- Novel nucleotide prodrug inhibiting DNA-replication of tumor cells, targeting cancer cells in the liver
- Designed to deliver high levels of active metabolite to the liver while minimizing systemic exposure



MIV-818 – Study design phase 1 monotherapy

Population studied

- advanced inoperable HCC, intrahepatic bile duct cancer and liver metastatic disease from solid tumors
- 9 patients (phase 1a) doses of 3-70 mg for 3-5 days in 21-day cycles, 10 patients (phase 1b) dose escalation starting at 40 mg for 5 days in 21-day cycles
- adult patients that had exhausted all approved therapies

Primary objective

- to assess safety and tolerability of MIV-818 as monotherapy
- to determine the recommended phase 2 dose for monotherapy

Secondary objective

- to evaluate tumor response rate based on RECIST v1.1

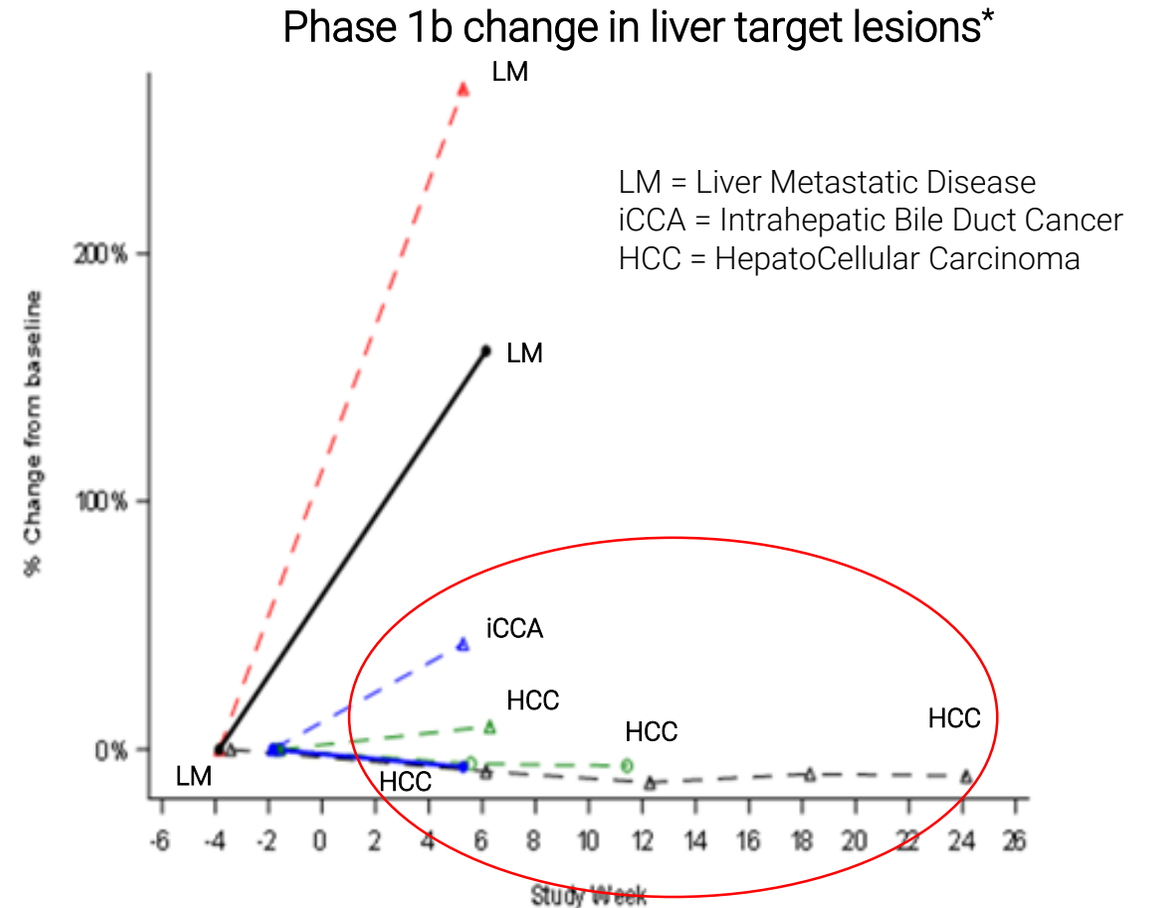
Exploratory objective

- to assess pharmacokinetics and pharmacodynamic effects of MIV-818

Phase 1b monotherapy results presented at ESMO

Supports continued development of MIV-818 in HCC

- Decreases in blood cell counts were the most common side effects, these resolved quickly
- In phase 1b four patients out of seven with primary liver cancer (e.g. HCC, iCCA) had stable disease as best overall response; one stayed on treatment for eight months
- Liver biopsy data has demonstrated delivery of MIV-818 to the liver, and a selective effect of MIV-818 on cancer cells vs normal liver tissue, across different types of cancer



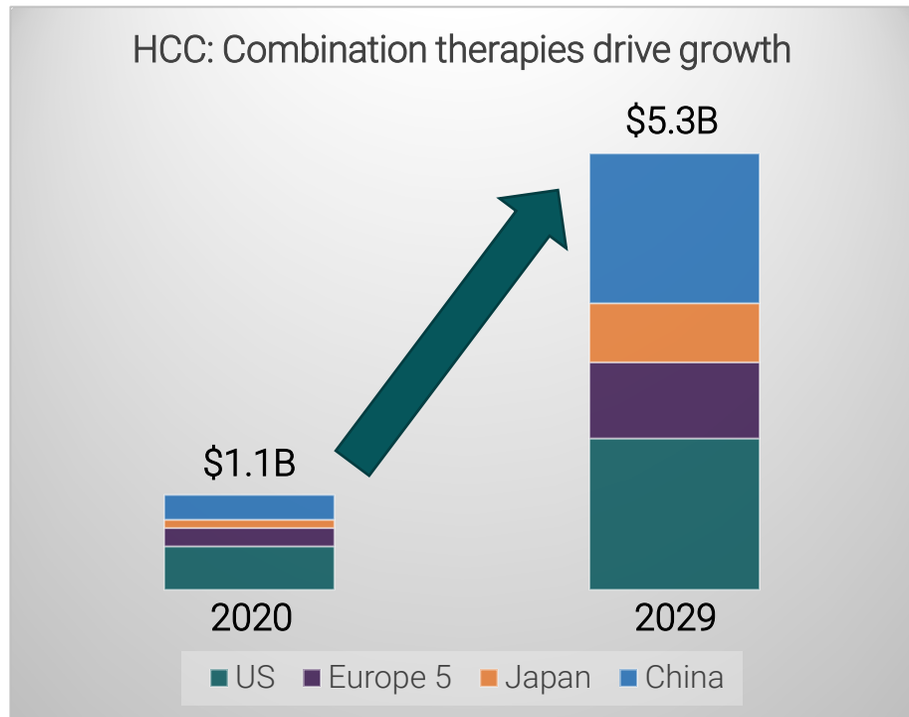
*Out of 10 enrolled patients, one did not complete safety follow up and one lacked independent radiologist assessment

Focus on hepatocellular carcinoma (HCC)

Based on its liver-targeting design and broad mechanism of action MIV-818 is a potential treatment for several tumors in the liver, and in combination with other therapies

- Clinical development program is initially focused on HCC
- Future opportunities for MIV-818 to be used in other settings and/or other cancer indications in liver (e.g. liver metastases or intrahepatic cholangiocarcinoma)

Hepatocellular carcinoma (HCC) is a growing market



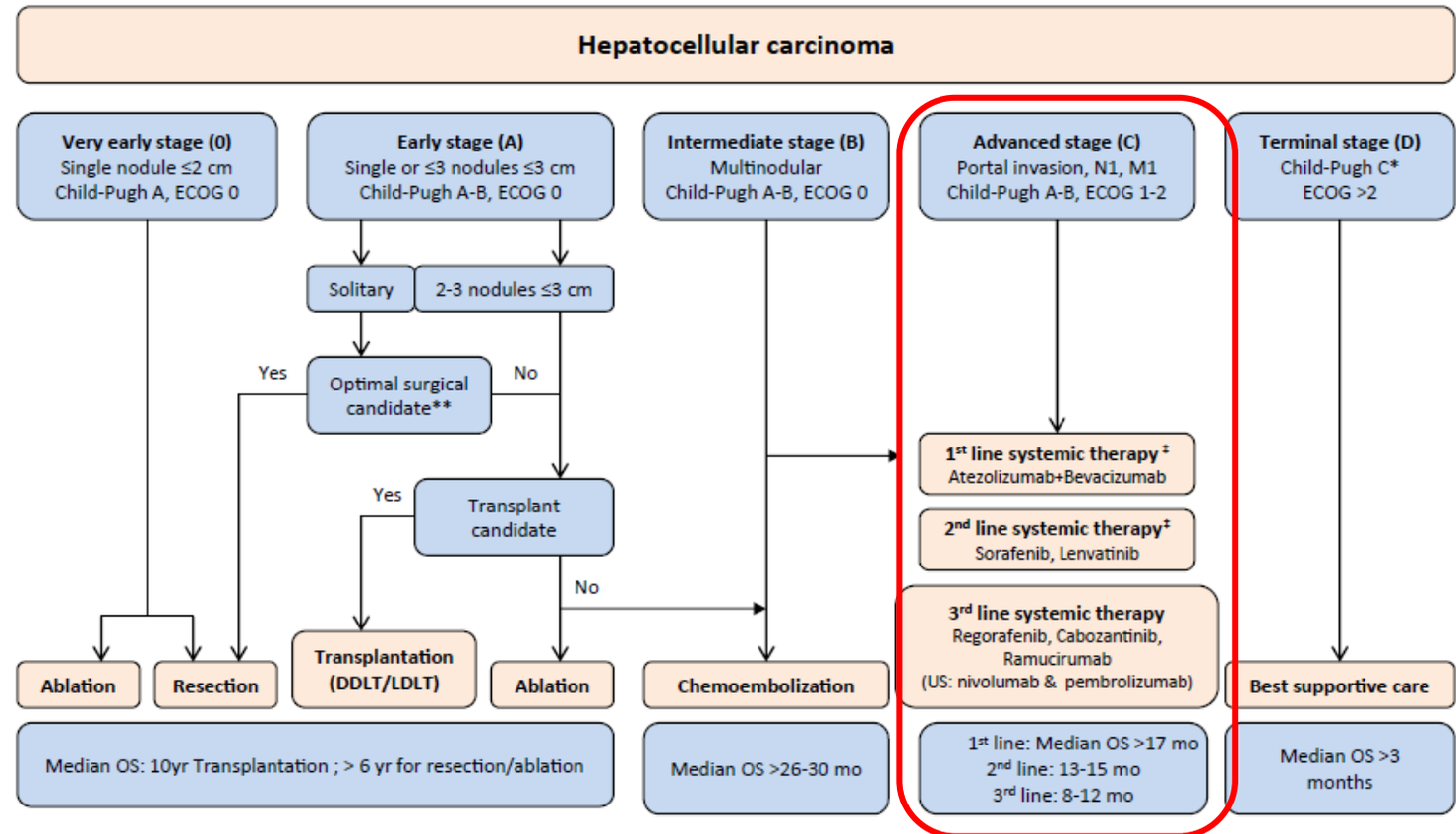
Source: GlobalData 2021

- Continued very high unmet medical need in HCC
 - Despite recent advances in treatment of HCC, there is still a large group of patients that do not respond to or are intolerant to current treatments
- The HCC market growth is driven by;
 - Combination therapies (especially immuno-oncology combinations)
 - More patients receiving therapy when patients are treated in earlier disease stages
- Liver cancer incidence and mortality are increasing and 5-year survival for those with advanced disease is less than 3% (<https://seer.cancer.gov/statfacts/html/livibd.htm>)

HCC Epidemiology and current treatments

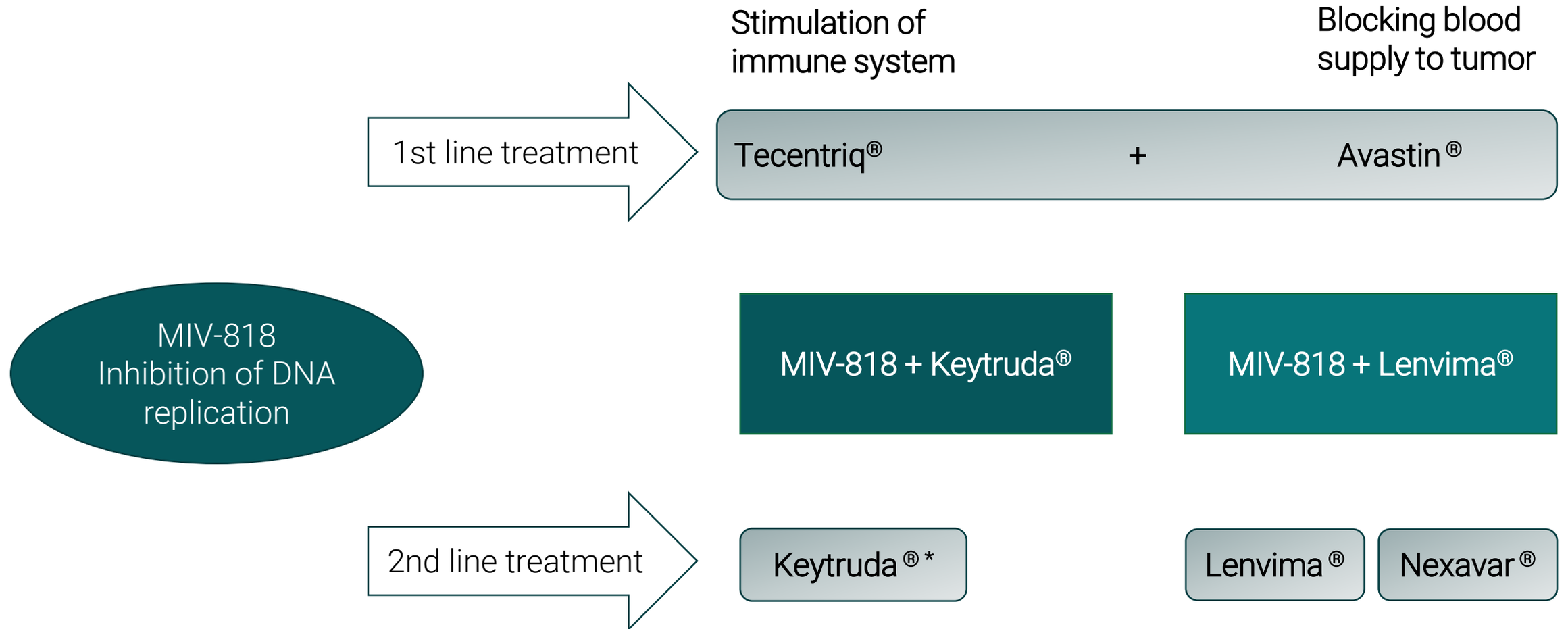
Primary liver cancers: 850,000 cases worldwide annually

- 90% are hepatocellular carcinoma (HCC)
- 3rd leading cause of cancer-related death, with 600,000 deaths worldwide



Llovet et al Hepatology vol 73, 2021

MIV-818 - aiming to be the new improved second line treatment

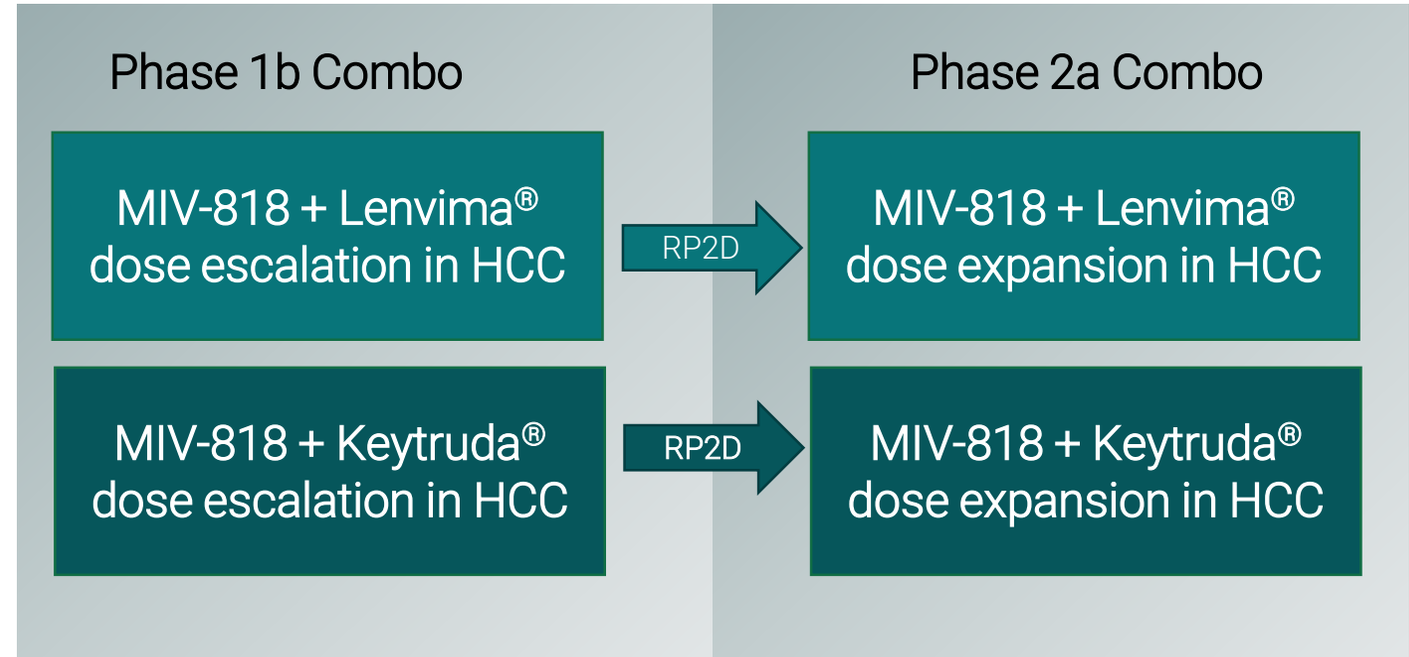


* Keytruda® only approved as monotherapy in HCC in US

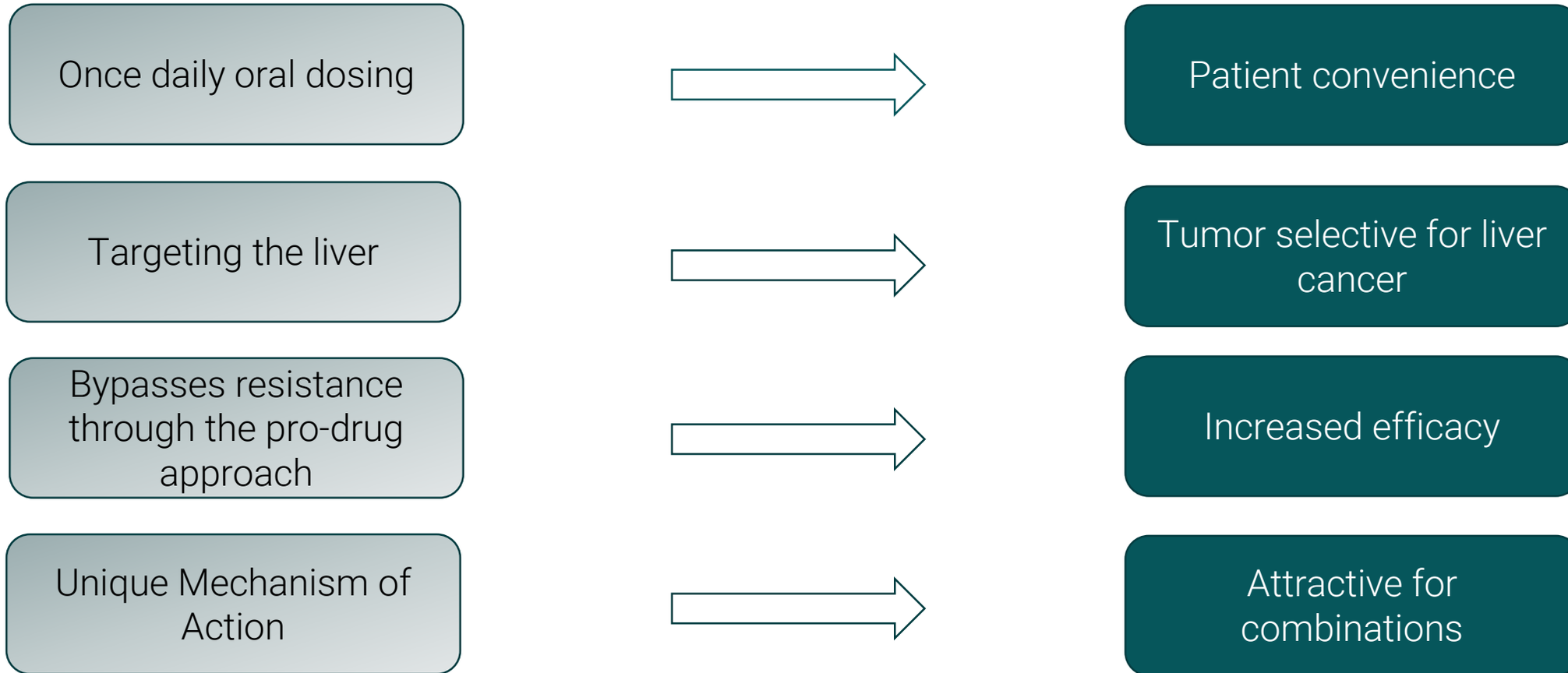
Phase 1b/2a combination study in 2nd line HCC

Patient population to be studied

- advanced inoperable HCC
- progressed on or intolerant of first line standard therapy for HCC
- candidates for Keytruda or Lenvima treatment



MIV-818 – Key advantages



Clinical portfolio and partnerships

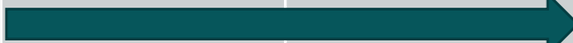
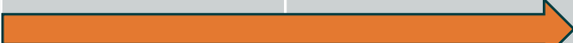
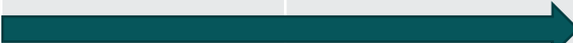
Focused clinical program

Nucleotide prodrug	Indication	Preclinical	Phase I	Phase II	Exclusivity
MIV-818	Liver cancer				IP : 2035

Partnered assets in clinical development

Compound	Mechanism	Indication	Phase I	Phase II	Partner	Exclusivity
Birinapant (IGM-9427)	SMAC mimetic	Solid tumors				IP : 2034

Multiple clinical programs for partnering/out-licensing

Compound	Mechanism	Indication	Phase I	Phase II	Phase III	Exclusivity
Remetinostat	Topical HDAC	MF-CTCL ¹⁾ BCC, SCC				IP : 2034
MIV-711	Cathepsin K inhibitor	Osteoarthritis				IP : 2034

1) Indications: basal cell carcinoma, squamous cell carcinoma, mycosis fungoides cutaneous T-cell lymphoma (phase III ready)

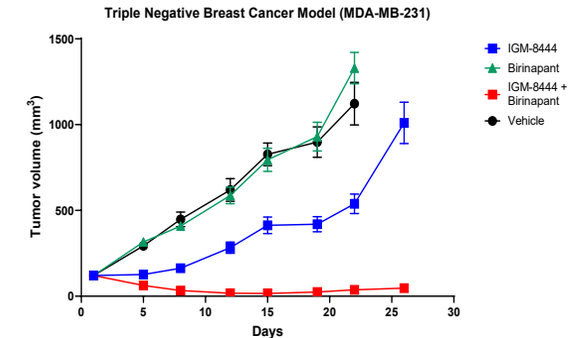
Delivering on our partnering strategy

Asset	Date	Partner(s)	Type of deal	Potential future revenues
Xerclear ¹⁾	Feb 2020	SYB	Outlicensing	Royalties
Malt1	Feb 2020	Rheos Medicines	Option	Option fee
USP-1	March 2020	Tango Therapeutics	Outlicensing	Milestones and royalties
Birinapant	Dec 2020	Tetralogic	Re-negotiated to enable an outlicensing deal	
Birinapant	Jan 2021	IGM Biosciences	Outlicensing	Milestones and royalties
USP-7	Feb 2021	Ubiquigent		Revenue share
Remetinostat	August 2021	Several stakeholders	Re-negotiated to enable an outlicensing deal	

1) Medivir receives royalties on Xerclear[®]/(Zoviduo[®]) European sales from Glaxosmithkline

Birinapant - Licensing agreement with IGM Biosciences

- IGM is a clinical-stage biotechnology company focused on creating and developing engineered IgM antibodies
- Birinapant will initially be combined with IGM-8444, a Death Receptor 5 (DR5) agonist being developed by IGM, which has demonstrated synergistic anti-tumor activity without added toxicity in several preclinical models
- Clinical studies with birinapant (IGM-9427) in combination with IGM-8444 has started
- Should birinapant be successfully developed and approved, Medivir is entitled to receive development, regulatory and sales milestone payments up to a total of approximately USD 350 million plus tiered royalties from the mid-single digits up to mid-teens on net sales



Open-label, Multicenter, Phase I Study with IGM-8444 in combination with Birinapant (IGM-9427) in patients with solid tumors will be in two stages: a dose-escalation stage and an expansion stage (NCT04553692)



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