

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 226M¹⁾

Market Cap: SEK 624M²⁾

FTE: 9

- 1) Q3 report
- 2) 2021-12-30



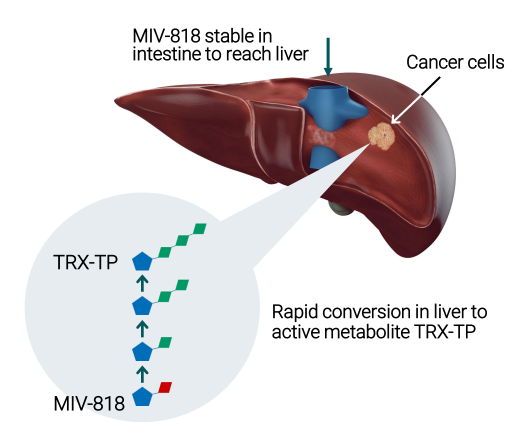
Lead asset – MIV-818 treating liver cancer

- 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 DNAreplication 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

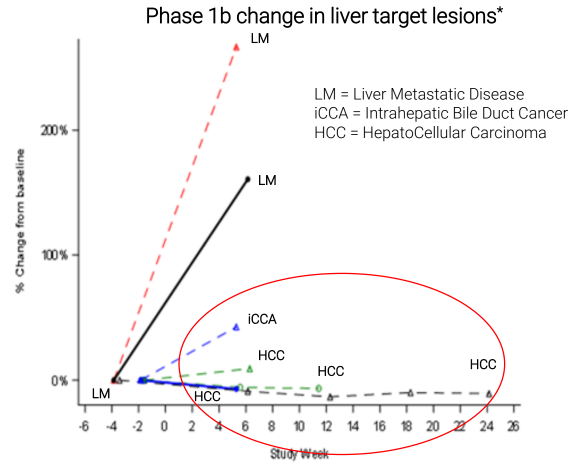
to assess pharmacokinetics and pharmacodynamic effects of MIV-818



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

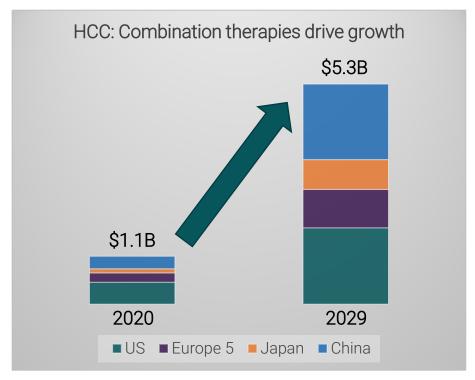
Focus on hepatocellular carcinoma (HCC)

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

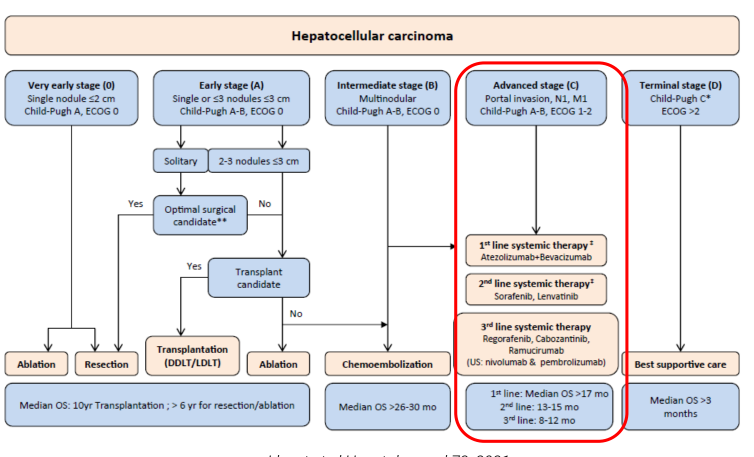
- Continued very high unmet medical need in HCC
 - Despite recent advances in treatment of HCC, there is still a large group of patients that do not respond to or are intolerant to current treatments
- The HCC market growth is driven by;
 - Combination therapies (especially immuno-oncology combinations)
 - 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

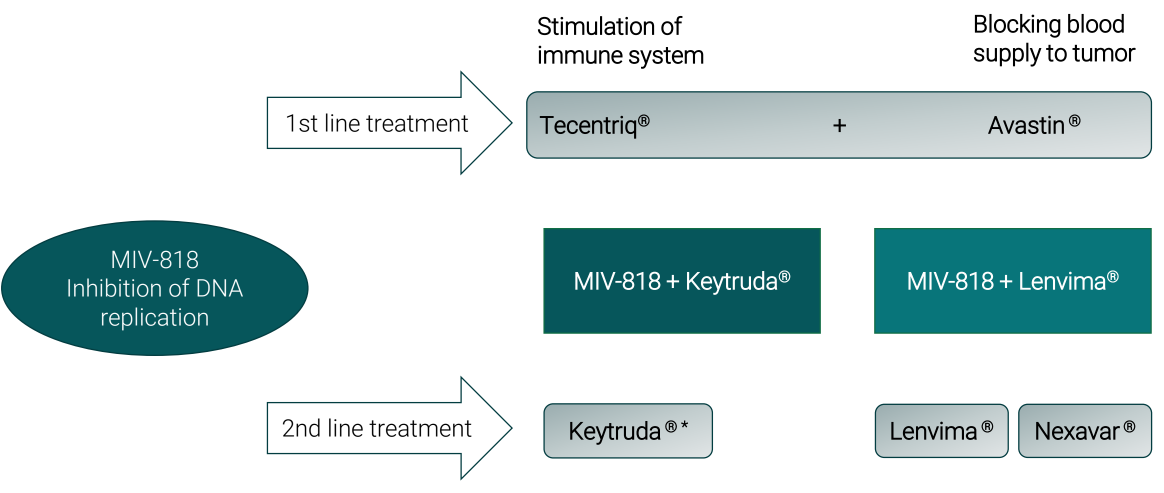
- 90% are hepatocellular carcinoma (HCC)
- 3rd leading cause of cancer-related death, with 600,000 deaths wordwide



Llovet et al Hepatology vol 73, 2021



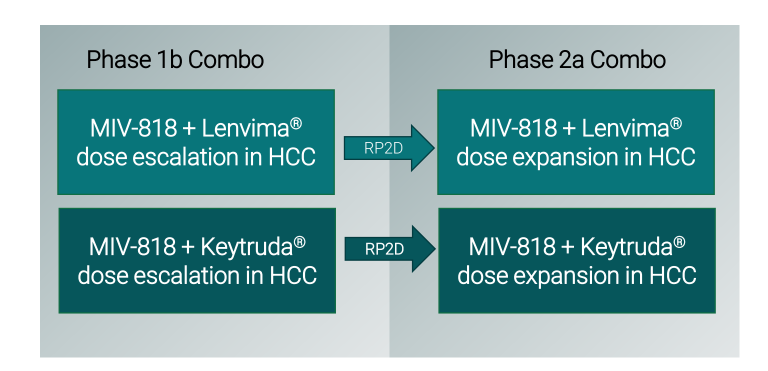
MIV-818 - aiming to be the new improved second line treatment



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

Once daily oral dosing

Targeting the liver

Bypasses resistance through the pro-drug approach

Unique Mechanism of Action









Patient convenience

Tumor selective for liver cancer

Increased efficacy

Attractive for combinations



Clinical portfolio and partnerships

MEDIVIR

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			©IGM biosciences™	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

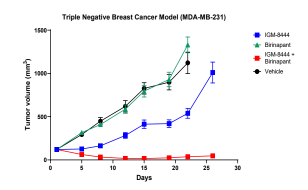
Asset	Date	Partner(s)	Type of deal	Potential future revenues
Xerclear ¹⁾	Feb 2020	SYB	Outlicensing	Royalties
Malt1	Feb 2020	Rheos Medicines	Option	Option fee
USP-1	March 2020	Tango Therapeutics	Outlicensing	Milestones and royalties
Birinapant	Dec 2020	Tetralogic	Re-negotiated to enable an outlicensing 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)



