



MEDIVIR

Improving life for cancer patients
through transformative drugs

June 2018

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Medivir in Brief

Improving life for cancer patients through transformative drugs

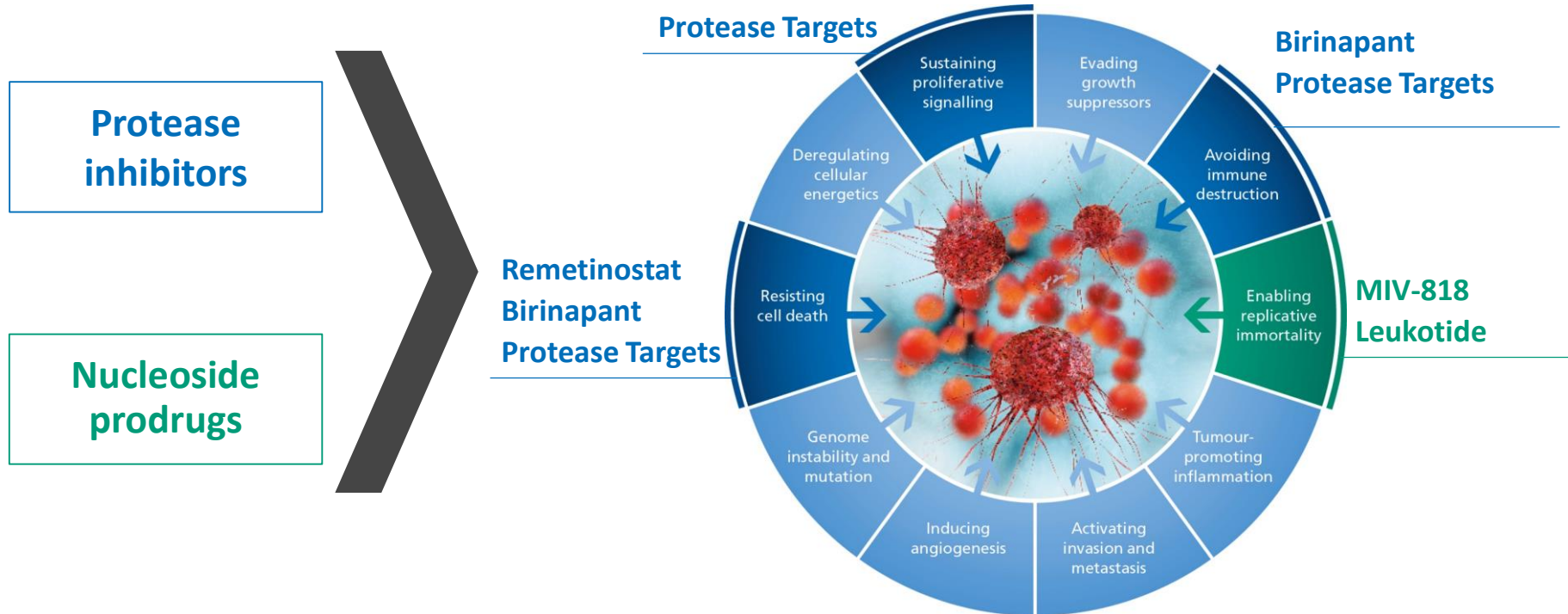
- Using **world-class scientific expertise** to bring new therapies to cancer patients
- **Clinical pipeline** composed of projects with **multi-billion dollar sales** potential as well as **orphan cancer drug candidates**
- Strong commercial focus – delivered more than **20 global partnerships** and **2 products from idea to market**

Basic facts

- Headquarters in Huddinge, Sweden
- 77 employees, 43 with PhDs
- Listed on the Nasdaq Stockholm, ticker: MVIR
- Current market capitalization: SEK 1bn (~USD 115m)¹
- Website: www.medivir.com



Leveraging scientific expertise to build pipeline in oncology



Adapted from: The Hallmarks of Cancer: The Next Generation.
Hanahan and Weinberg, Cell (2011), 144, 646-674

Oncology drug development in areas of high unmet need

Strong and balanced development pipeline based around areas of scientific expertise and focused on cancer

	Project, Mechanism	Disease area	Clinical phase				Market	Next step
			Preclinical	Phase I	Phase II	Phase III		
Cancer	Remetinostat Topical HDAC inhibitor	Early-stage cutaneous T-cell lymphoma	[Blue bar spanning Preclinical, Phase I, and Phase II]				~\$1b US only	P3 start 2018
	Birinapant SMAC mimetic	Solid tumors (combo with Keytruda®)	[Blue bar spanning Preclinical and Phase I]				Blockbuster	P2 start 2H2018
	MIV-818 , Nucleotide DNA polymerase inhibitor	Hepatocellular carcinoma	[Green bar in Preclinical]				Orphan US/EU Significant Asia	P1 start 2H2018
	MIV-711 Cathepsin K inhibitor	Osteoarthritis	[Blue bar spanning Preclinical, Phase I, and Phase II]				Blockbuster	Partner

■ Protease related
■ Nucleot(s)ide related

Collaborations enhance the value of programs

Academic



Industrial

Product/Project

Zovido®/Xerclear
(labial herpes)
acyclovir + hydrocortisone

MIV-802 (HCV)
*Nucleotide NS5B
polymerase inhibitor*

Platform Link

Nucleoside
analogue

Nucleotide

Partners



Status

Marketed

Phase I ready

Medivir Interests

- Royalties from sales
- Approval milestones for additional OTC switches
- Development milestones
- Royalties from sales

Competences from discovery through regulatory approvals

Management team with extensive experience and proven track record of successful development



RICHARD BETHELL, Chief Scientific Officer

- 28 years drug discovery and development in oncology and infectious disease
- VP, Biology/DMPK (Boehringer Ingelheim (Canada))
- VP, Therapeutic Research (Shire)
- Pfizer and GlaxoSmithKline R&D
- Doctor of Philosophy (D. Phil.) in chemistry from Oxford University



JOHN ÖHD, Chief Medical Officer

- Senior director of Experimental Medicine, Shire
- Early development group director, cognitive and neurodegenerative disorders at AstraZeneca
- Cancer research at Lund University and at Karolinska Institute
- Clinical training at Karolinska University Hospital
- MD, Linköping University, PhD in Experimental Pathology, Lund University



ÅSA HOLMGREN, EVP Strategic Regulatory Affairs

- Head of Regulatory Affairs at Orexo AB
- Various large pharmaceutical companies, including 12 years as Senior Global Regulatory Affairs Director at AstraZeneca, and at AstraZeneca in Canada and Japan
- M. Sc. in Pharmacy, trained Uppsala University



CHRISTINE LIND, President and CEO

- EVP, Business Development at Medivir
- VP, Business Development, LifeCell Corporation
- Biotech and pharma strategic advisory and capital raising at Merrill Lynch & Co. and GKM & Co.
- B. Sc. Finance and Info Systems, NYU and MBA, Columbia Business School



ERIK BJÖRK, Chief Financial Officer

- CFO for AstraZeneca Sweden Operations
- 11 years with Procter & Gamble, in global finance leadership positions in Switzerland, UK and Sweden
- MSc in Finance and LLM from Lund University



CHRISTINA HERDER, EVP Strategic Business Development

- CEO of Modus Therapeutics
- Director, Corporate Development at Sobi
- Project & Portfolio Management at Biovitrum
- Current member of the boards of PCI Biotech and Idogen
- Ph. D. in physical chemistry from Royal Institute of Technology and MBA from Stockholm University



DANIEL ERIKSSON, Chief Information Officer

- Various IT roles relating to security, decision support, innovation, and digitalization
- Most recently, Technical Director for G4S Risk Management
- PhD Coventry University and BSc in Systems Science, Linköping University

Cancer biology, chemistry, intellectual property, DMPK, CMC, toxicology, clinical development, regulatory strategy, business development

77 employees, 43 with PhDs,
18 nationalities, balanced gender split

MIV-711 for Osteoarthritis

MIV-711: ORAL ONCE DAILY CATHEPSIN K INHIBITOR WITH FDA FAST TRACK STATUS FOR OA DISEASE MODIFICATION

No existing disease-modifying drug for osteoarthritis

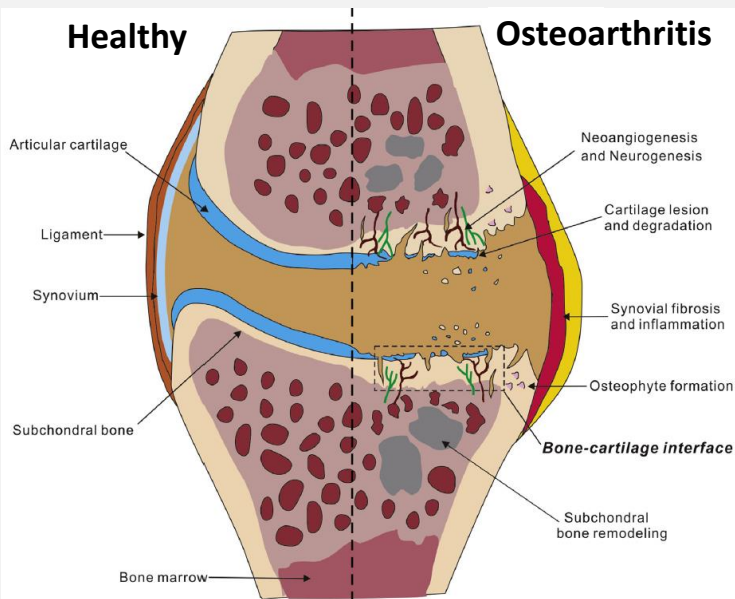
Blockbuster revenue opportunity for a disease-modifying osteoarthritis drug

- Affects >30m adults in the US, and ~240m worldwide
- Prevalence increasing due to aging population and obesity epidemic
- Current treatments are insufficient focusing only on symptom relief



Disease involves both bone and cartilage

- Cathepsin K protease involved in the breakdown of collagen in both bone and cartilage



Yuan et al. Osteoarthritis and Cartilage (2014), 22, 1077

MIV-711: ORAL ONCE DAILY CATHEPSIN K INHIBITOR WITH FDA FAST TRACK STATUS FOR OA DISEASE MODIFICATION

Phase IIa data show unprecedented OA disease modification after 6 months

Potential disease-modifying convenient osteoarthritis drug

- Only cathepsin K-inhibitor in development for osteoarthritis
- Once daily oral administration

Strong patent position

- Expected patent life to ~2034, including extensions

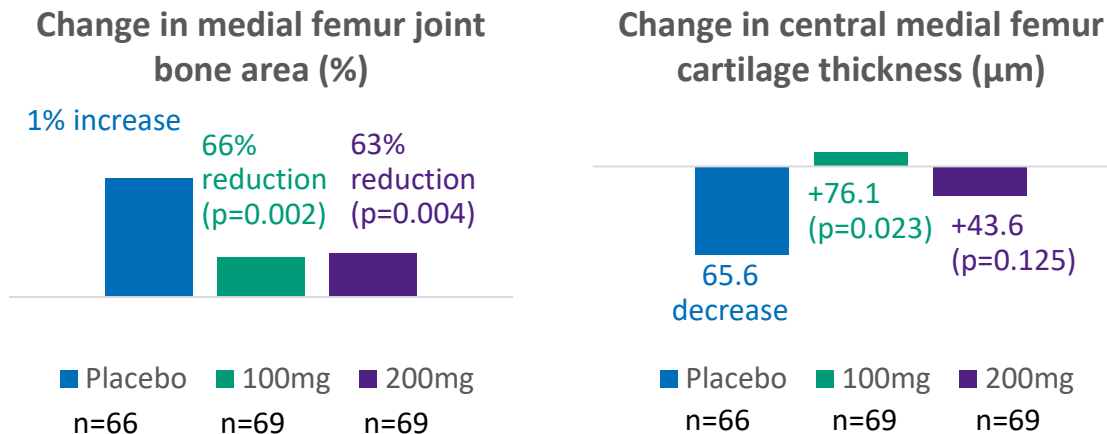
US FDA Fast Track designation

- Granted by FDA October 2017

“The finding that MIV-711 slows the degenerative changes on both bone and cartilage in knees affected by OA is an enormously exciting finding”

Professor Philip Conaghan, Professor of Musculoskeletal Medicine at the University of Leeds, UK, and lead investigator on the MIV-711-201 study

Benefit on both bone and cartilage in Phase IIa study

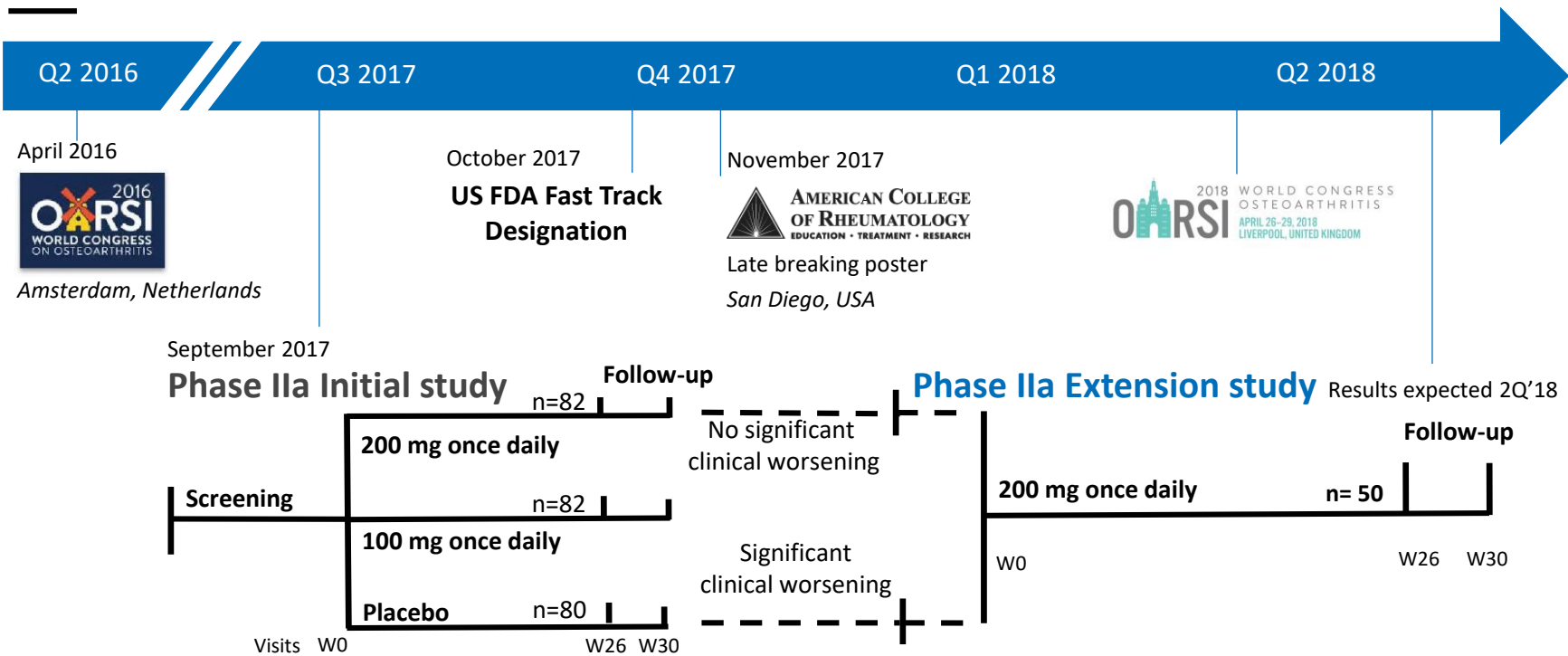


- MIV-711 consistent tendency to improve outcomes across all pain and other patient reported symptoms vs. placebo
- Acceptable safety and tolerability profile

<http://acrabstracts.org/> Abstract 14L

MIV-711: ORAL ONCE DAILY CATHEPSIN K INHIBITOR WITH FDA FAST TRACK STATUS FOR OA DISEASE MODIFICATION

MIV-711: Building towards partnership

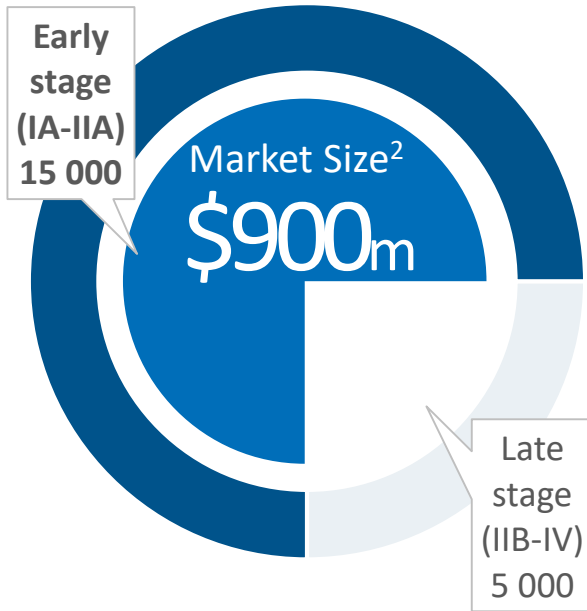


- Partnering discussions ongoing

Remetinostat for early-stage CTCL

CTCL: orphan blood cancer with significant market opportunity

US CTCL patients¹: orphan disease



Early Stage CTCL: Disease background

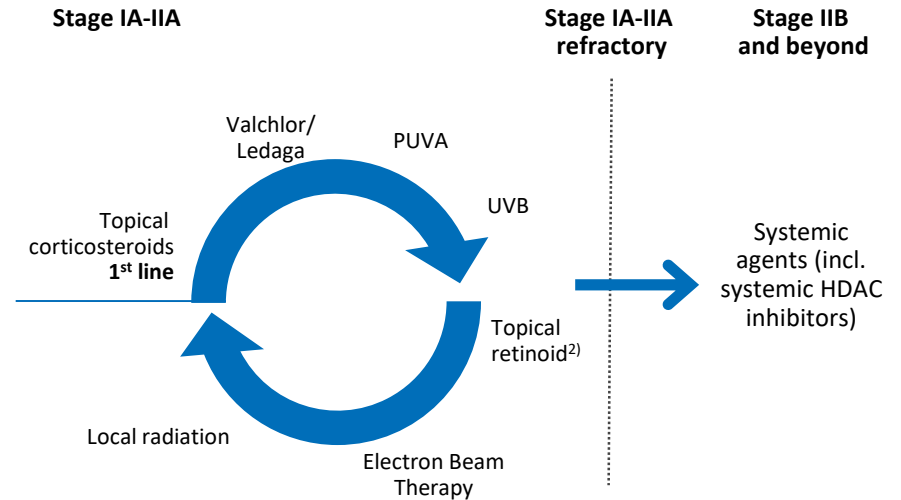
- Confined to the skin
- High 5-year survival rates (~85%)
- Patients remain at this stage for extended periods and require long-term treatment
- Significant quality of life issues, especially pruritus (itch)

Key unmet need:
balance of efficacy and long-term tolerability

Patients & physicians need new treatments for early-stage CTCL

- Currently approved early-stage CTCL drugs lack sustained efficacy and/or tolerability and are highly irritating
- Valchlor/Ledaga is a topical alkylating agent (the active agent in mustard gas)
- No single treatment for long-term use
- At this early stage, physicians avoid using systemic drugs, including other HDAC inhibitors, due to the side effect profile

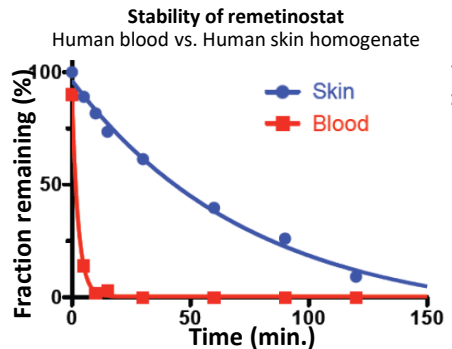
Currently approved therapies by disease stage



Remetinostat potential to meet patients' key unmet need

Designed to act only where needed

- HDAC inhibitors¹ approved in late-stage CTCL patients but not in early-stage patients due to systemic toxicities
- Reteminostat's unique design and topical application provides activity in skin, but rapid degradation in blood



- Expected patent life to ~2034 (including extensions)
- US orphan drug designation

“As a topical, skin-specific HDAC inhibitor, reteminostat has the potential to be efficacious and have an improved safety profile compared to other available treatments.”

Youn Kim M.D.
Stanford University Medical Center, USA

Addresses key unmet need with positive Phase II data

Effect on lesions & reduction of pruritus (itch)

Dose	1% 1x/day n=20	0.5% 2x/day n=20	1% 2x/day n=20
Lesion responses ¹	20%	25%	40%
Patients with clinically significant pruritus ²	8/20 (40%)	6/20 (30%)	10/20 (50%)
Pruritus responses	37.5%	50%	80%

Highly tolerable with no systemic side effects

- Even dose distribution of AEs, mostly grade 1 or 2
- No HDAC inhibitor-associated systemic adverse events
- Median time on treatment: 332 days (1% 2x/day dose)

M Duvic *et al.*, EORTC Cutaneous Lymphoma Task Force Meeting (2017), Abstract O55

Planned Phase III clinical development for early-stage CTCL

Design

- One Phase III study expected to be sufficient for NDA
- Past approvals in CTCL were based on pivotal clinical studies involving ≤ 260 patients
- Focus on treatment-experienced patients where the medical need is high

Program Timing

- Phase II final data reported (EORTC CLTF Meeting, 2017)
- Ongoing discussions with the FDA (End of Phase II) to enable Phase III

Costs

- ~\$50m (SEK 400m) expected clinical costs to NDA submission over a ~3 year period (incl. Phase III study and third party milestones)

“The introduction of remetinostat to the market as a novel topical agent for the treatment of CTCL is likely to find broad application and lead to novel combinatorial approaches in CTCL.”

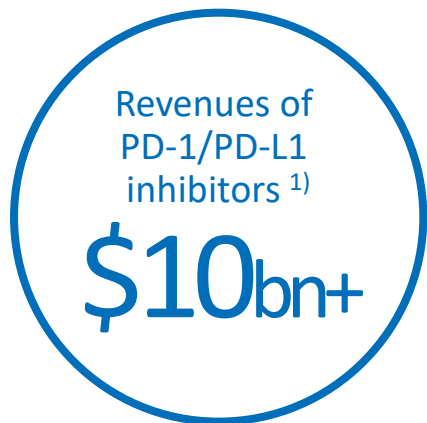
*Pierluigi Porcu, M.D.
Jefferson University Hospital, USA*



Birinapant for solid tumors

Despite immuno-oncology breakthroughs patients have unmet needs

Multi-billion dollar market,
and growing for immuno-
oncology agents



< 1/2

of patients derive
meaningful clinical
benefit in approved
indications

0-5%

ORR in other indications
such as MSS colorectal
cancer

Combination
regimens to
enhance benefit
in underserved
patients

Linking targeted therapy with immuno-oncology

Uniquely potent molecule against a novel target

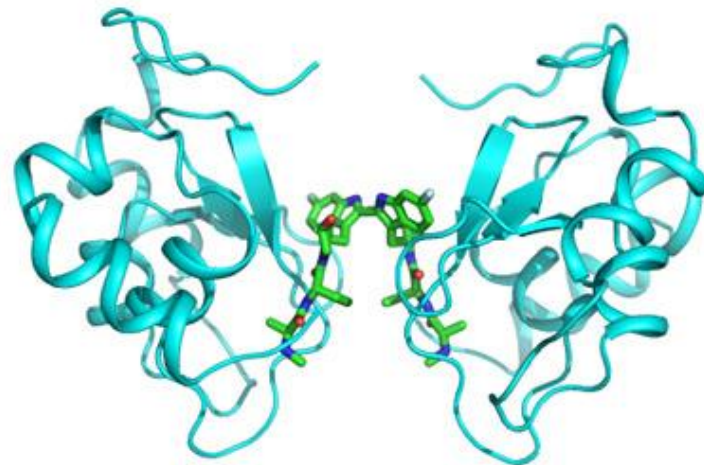
- Bivalent SMAC (second mitochondrial-derived activator of caspases) mimetic
- Targeting of cIAPs results in dual action on T-cells and tumor cells

Strong rationale for use in combinations

- Synergistic with anti-PD1 and likely other IO pathways
- Efficacy observed in both hematological and solid tumor models
- Phase I/II study in combination with Merck's Keytruda® underway

Blockbuster potential and strong patent position

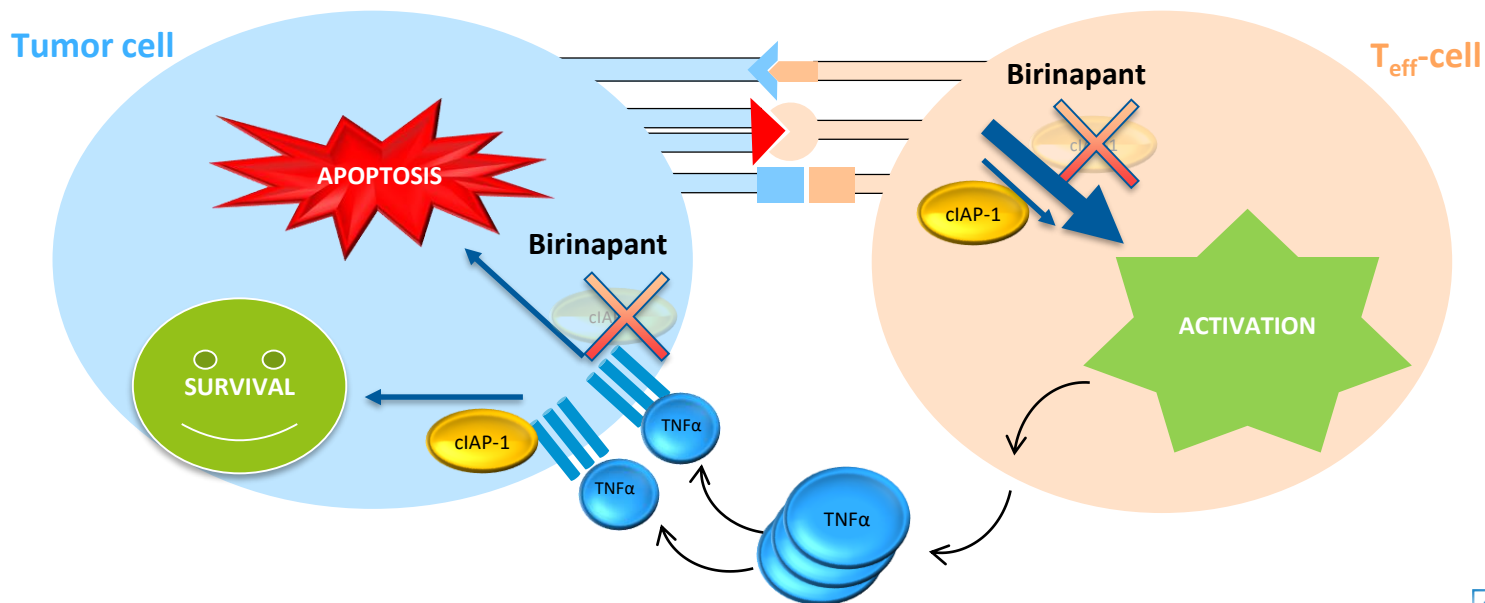
- Expected patent life to ~2034, including extensions



Dual action enhances cancer cell death

Targeting of cIAPs results in dual action on T-cells and tumor cells

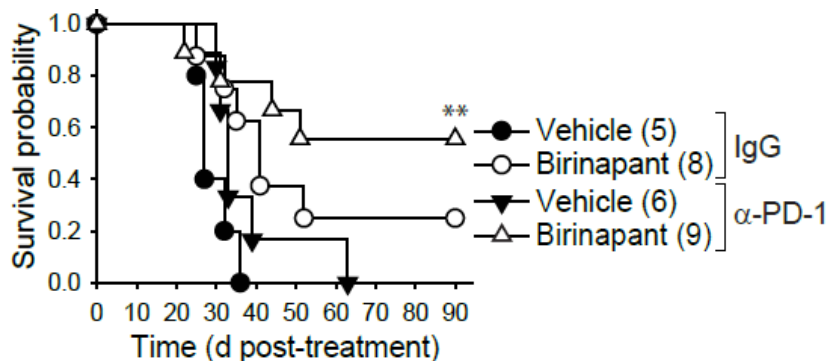
- Enhances pro-apoptotic, and suppresses pro-survival, signaling downstream of TNF- α
- Augments human T cell responses to physiologically relevant stimuli¹



Potential to enhance patient response with immune-oncology therapies

Strong rationale for combination with Keytruda®

- Birinapant/anti-PD1 mAb combo showed enhanced activity in preclinical models¹ compared to either agent alone



¹) Solid tumor model: Beug et al., Nature Communications (2017) 8:14278

Multiple myeloma model: Chesi et al., Nature Med. (2016) 22, 1411–1420

Cooperation of IAPs and PD-L1 to protect tumor cells: Kearney et al., Cell Death and Diff. (2017), 24, 1705–1716

Phase I/II study underway in collaboration with MERCK

- Development collaboration for the Phase I/II study in solid tumors
- Keytruda® provided at no cost
- Joint Development Committee to oversee the study, bringing Merck's immuno-oncology expertise
- Medivir retains full global rights to birinapant and data

Birinapant/Keytruda® combination: Phase I/II Study underway



June 2018
ASCO American Society of Clinical Oncology
Chicago, USA



Phase I: Sequential group dose-escalation to determine the dose-limiting toxicity and recommended Phase II dose, in combination with Keytruda®

Phase II: Safety and tolerability of the recommended dose of birinapant, in combination with Keytruda® in defined disease cohorts:

- MSS Colorectal cancer
- Ovarian cancer
- Cervical cancer
- Other cancer types exploratory cohort



MIV-818 for liver cancers

Liver cancer is 2nd leading cause of cancer related death worldwide

Liver cancer¹

- Orphan disease in Western markets, but much more common in Asia
- One of fastest growing and most deadly cancers in US
- Genetically heterogeneous leading to limited effect of molecularly targeted therapies

Patients with advanced liver cancer in need of new treatments

Sorafenib
(kinase inhibitor)

- ~3 month survival benefit



Regorafenib
(kinase inhibitor)

- ~3 month survival benefit

Nivolumab
(PD-1 antagonist)

- 15-20% ORR
- Survival benefit yet to be fully defined

1) Howlader et al. (eds). SEER Cancer Statistics Review, 1975-2011, National Cancer Institute. Bethesda, MD, http://seer.cancer.gov/csr/1975_2011/

Potential to improve efficacy and safety for patients with liver cancers

Improve a nucleoside with Medivir prodrug technology

Troxacitabine

(nucleoside)

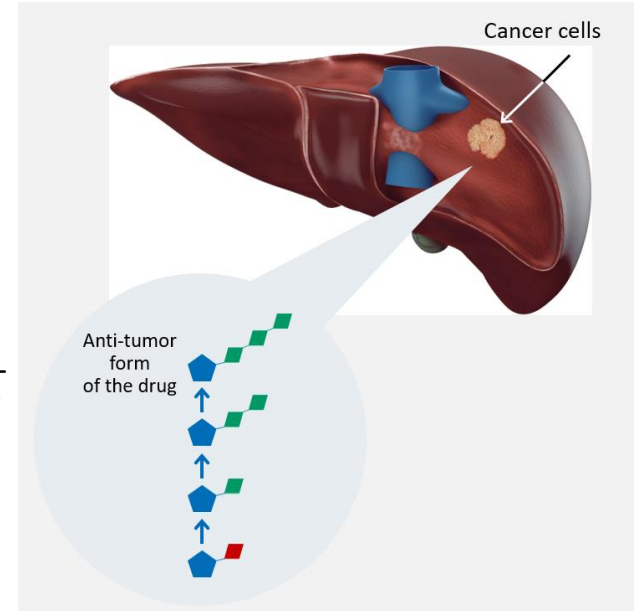
- **Active** in preclinical cancer models and in clinic
- Failed in clinic due to systemic dose-limiting **toxicities**

Medivir
prodrug
technology

MIV-818

(liver-targeted nucleotide prodrug)

- **Enhanced activity** 10x more potent against HCC cell lines than parent troxacitabine
- **Selectivity for cancer** Active on HCC cells while sparing non-cancerous hepatocytes
- **Improved delivery to the liver** >100-fold relative to systemic exposure of troxacitabine
- **Market exclusivity** with full new chemical entity patent protection



MIV-818: Gearing up for Phase I study start

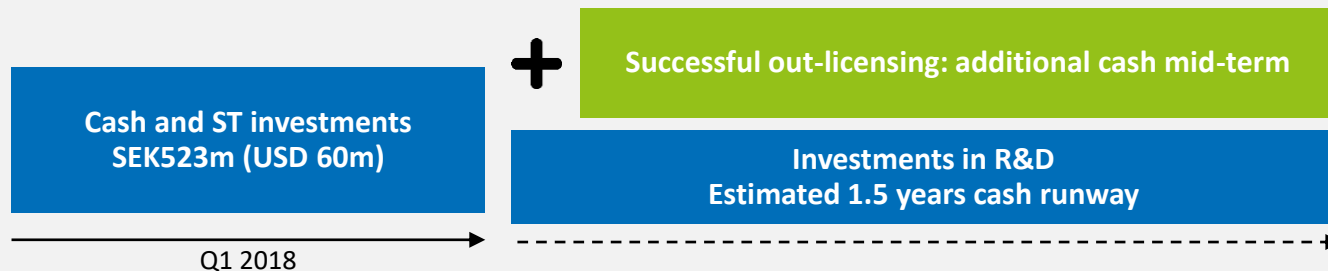




Outlook

Cash position and shareholder base

CASH POSITION



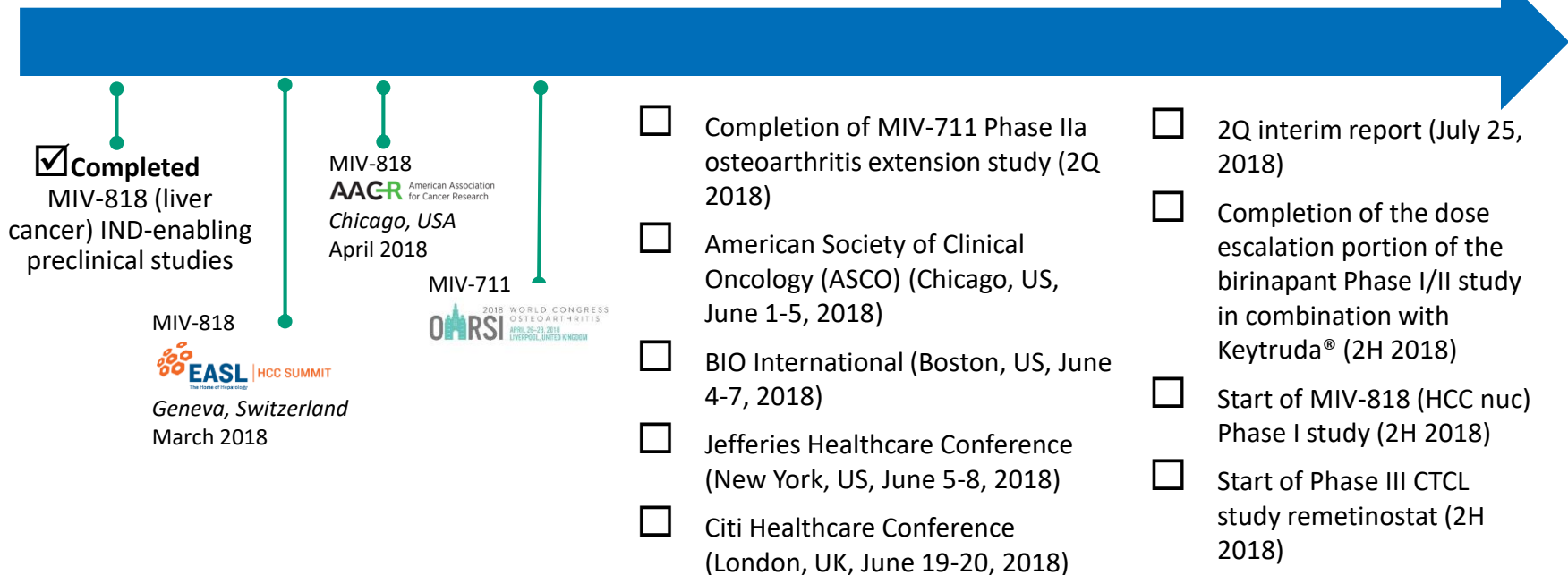
THE SHARE



What to look for in 2018

Continuous track record of delivery

Coming events



Why Medivir?

For more information:

- Nasdaq Stockholm, ticker: MVIR
- www.medivir.com

- Track record of delivery

3 new drugs from research
into development in 2 years

2 products from
idea to market

>20 global partnerships,
multiple repeat partners

- Strong pipeline from discovery through clinical stages with upcoming catalysts
- Competences from discovery through regulatory approvals
- Near-term opportunities for revenues from partnerships



Appendix

Positive Phase II data: Confirmed efficacy on skin lesions and reduced itching

Study design

- 60 patients with stage IA-IIA MF were randomized into three dose arms and treated for up to 12 months
- Index lesions were identified at baseline and assessed throughout the study
- The primary end-point was the proportion of patients with a complete or partial confirmed response assessed using the Composite Assessment of Index Lesion Severity (CAILS)

Results

- Dose response: CAILS ORR & pruritus VAS responses
- Patients in the highest dose group had the highest proportion of confirmed responses (40%), including 1 complete response
- A positive effect was also seen on the severity of pruritus, a secondary objective in the trial

Dose	Once Daily		Twice Daily	
	1% (n=20)	0.5% (n=20)	1% (n=20)	1% (n=20)
Lesion Outcomes				
CAILS Confirmed Overall Response Rate (ORR)	4 (20%)	5 (25%)	8 (40%)	
Median Duration of CAILS Confirmed Response ¹	2 months	3 months	7 months	
Pruritus Outcome				
Patients with clinically significant pruritus at baseline (VAS ≥30 mm at baseline)	8/20 (40%)	6/20 (30%)	10/20 (50%)	
Confirmed response in patients with clinically significant pruritus at baseline	3/8 (37.5%)	3/6 (50%)	8/10 (80%)	

Well tolerated without signs of systemic adverse events

Results

- Across all the dose groups, remetinostat was well-tolerated without signs of systemic adverse effects, including those associated with systemic HDAC inhibitors
- A balanced safety profile across all three doses
- Minimal systemic exposure, even at the top dose
- Most patients remained on study for the maximum possible duration
 - Median time on treatment: 332 days (1% 2x/day dose)

Treatment-related Adverse Events seen in ≥1 Patients ¹	Once daily		Twice daily	
	1%	0.5%	1%	1%
Any Adverse Event	11	10	11	11
Pruritus	5	3	1	1
Any Other Skin ²	9	10	11	11
Infections	3	1	0	0
Skin papilloma	0	0	1	1

Positive trends across all Pain and other Patient Reported Outcomes

MIV-711 showed consistent tendency to improve patient-reported symptoms, including pain, however did not reach statistical significance

- A tendency was observed favoring both the 100mg and 200mg groups for patient-reported pain on the NRS scale (the primary endpoint)
- This tendency was observed consistently across other patient-reported symptoms such as:
 - Daily reporting of pain in E-diaries
 - Measures of pain associated with the daily activities
 - Satisfaction with the function of the diseased knee
- The findings on pain and other clinical symptoms from this study will enable the design of future, longer, studies aimed at demonstrating the effects of MIV-711 on both joint structure degeneration and pain and other clinical symptoms



Drug discovery expertise: Nucleoside Prodrugs & Protease Inhibitors

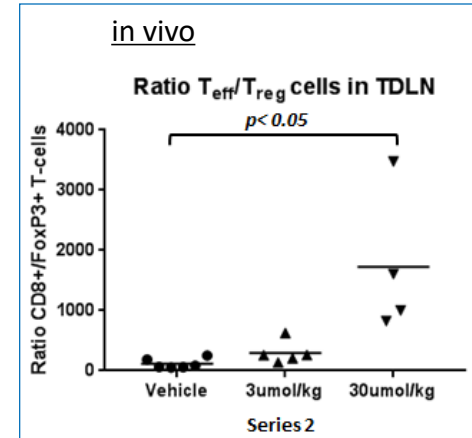
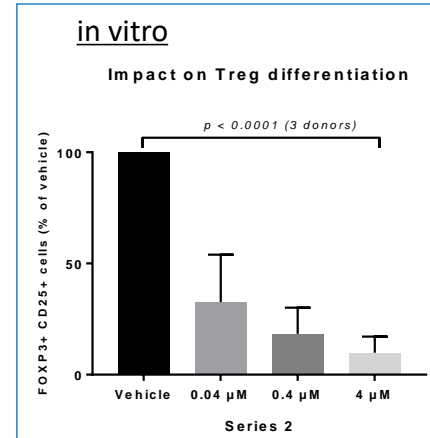
Leukotide nucleotide prodrug for AML

- Aim to develop better tolerated and more effective agent to improve outcomes for patients with acute myeloid leukemia (AML) and other hematological cancers
- AML is a relatively rare cancer, with around 21,000 cases expected in the US in 2017. The prognosis is poor for many AML patients, especially those who are elderly, because they are often unable to tolerate the intensive treatments currently used to cure the disease.
- Five-year survival of patients in the US diagnosed with AML was 27% in the period 2007–2013

Market exclusivity with full NCE patent protection

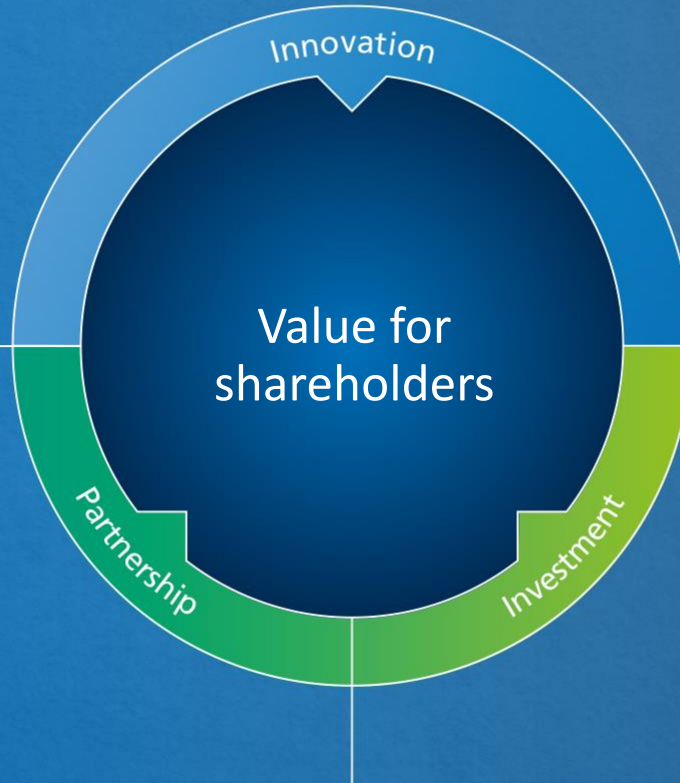
TRIP: T_{reg} inhibitor project for immuno-oncology

- Novel biological target enabling selective suppression of T_{reg} cells
- IP filed on the target itself and 2 classes of small molecules
- Highly potent compounds (K_i values <15 nM against the molecular target)
- Increase of T_{eff}/T_{reg} cell ratio demonstrated:



Track record of delivery

- Medivir was founded 1988
- Public company since 1996



3 candidate drugs into development in 2 years

- MIV-818 for liver cancer
- MIV-323 for RSV
- MIV-802 for HCV

>20 global partnerships,
multiple repeat partners

>\$400m¹

2 products from idea to market

- Xerclear/Zoviduo for labial herpes
- Olysio (simeprevir) for HCV, peak sales \$2.3b worldwide

Transformation into an oncology-focused biopharma company

<2017 – THE NEW MEDIVIR EMERGES

- Sold BioPhausia (established portfolio of branded generics)
- Transitioned commercial products in the Nordics to partners
- Delivered first oncology development project from in-house research
- Reorganized to increase focus and agility in R&D
- Acquired two clinical stage oncology projects:
 - Remetinostat (Phase II for CTCL)
 - Birinapant (Phase I/II solid tumors)



>2017 - DELIVERING ON ONCOLOGY PIPELINE

- Remetinostat (early-stage CTCL): delivered positive Phase II topline efficacy & safety from 12 month study
- Birinapant (solid tumors): started Phase I/II combination study with Keytruda
- MIV-818 (liver cancer): completed IND-enabling GLP safety studies
- New cancer project Leukotide based on in-house discovery



2017 Accomplishments

- ✓ MIV-711 groundbreaking phase IIa data announced and presented as late breaker at ACR
- ✓ MIV-711 Fast Track designation from US FDA
- ✓ MIV-711 extension study fully enrolled
- ✓ Remetinostat Phase II data announced and presented at EORTC Cutaneous Lymphoma Task Force meeting
- ✓ Birinapant/Keytruda® Phase I/II study started
- ✓ MIV-802 for HCV in Greater China licensed to Ascleptis
- ✓ Beta lactamase research program licensed to AMRC
- ✓ Distribution of 858 MSEK to shareholders from sale of BioPhausia
- ✓ Completion of reorganization to reduce costs
- ✓ New management team in place, including new CEO



Financial Summary

Summary of the Group's figures

(SEK m)

	Q1		Full Year	
	2018	2017	2017	2016
Net turnover	4.5	17.8	36,6	93,0
EBITDA	-73.1	-80.9	-342,6	-300,6
Basic earnings per share, SEK	-3.17	-3.59	-16,40	-10,94
Net worth per share, SEK	24.14	38.93	25,31	64,38
Cash flow from operating activities	-87.1	-123.9	-358,5	-182,3
Cash and cash equivalents at period end	522.7	708.9	467,8	1 698,5

- Net turnover from royalty revenue alone, vs. own sales in earlier periods
- Cost and cash spend is driven by the strong development in our clinical projects
- Lower cost base due to full realization of the reorganization announced in fall 2016
- Quarter end 2018 cash position positively impacted by the proceeds from the directed share issuance completed in February 2018

Board of directors



ANNA MALM BERNSTEN, Chairman, director since 2006

- **Education:** M.Sc. in Engineering
- **Background:** M.Sc. in Engineering with an extensive knowledge of the life sciences sector. Runs own consultancy firm operating in the fields of leadership strategy and business development. Experience of senior positions with, amongst others, GE Healthcare Life Sciences, Pharmacia, Assa Abloy, Medvir and Baxter Medical. Former President & CEO of Carmeda AB
- **Other assignments:** Chairman of the Boards of CEBA AB and Oatly AB. Member of the Boards of Cellavision, Pägengruppen and Probi



ULI HACKSELL Director since 2018

- **Education:** PhD at Uppsala University
- **Background:** Senior positions in major pharmaceutical and biotech companies for over 25 years and more than 10 years' experience as the CEO of publicly owned companies. As the CEO of ACADIA Pharmaceuticals from 2000–2015, he led its development from a private start-up to a public, multibillion dollar company. In the 1990s, he held senior positions at Astra AB, prior to which he was a Professor of Organic Chemistry at Uppsala University.
- **Other assignments:** Chairman of the Board of Cerecor Inc., and Member of the Boards of InDex Pharmaceuticals AB, Beactica AB and Uppsala University.



ANDERS R HALLBERG, Director since 2012

- **Education:** Professor of Medicinal Chemistry at Uppsala University's Faculty of Pharmacy
- **Background:** Held a number of positions as a scientific advisor at AstraZeneca and smaller pharmaceutical companies between 1990 and 2006. Prior to this, he was the Head of the Medicinal Chemistry Department at Astra in Lund. Vice-Chancellor of Uppsala University between 2006 and 2011. He has published over 270 scientific articles, a large number of which are on the subject of pharmaceuticals for the treatment of infectious diseases, and co-inventor of a large number of granted patents. Member of the Royal Swedish Academy of Sciences, and the Royal Swedish Academy of Engineering Sciences. Awarded honorary doctorates in Sweden and other countries
- **Other assignments:** Member of the Boards of foundations and universities.



LENNART HANSSON Director since 2018

- **Education:** PhD in Genetics from Umeå University
- **Background:** Extensive experience of pharmaceutical and commercial development in senior positions with both biotech and pharmaceutical companies, such as KabiGen AB, Symbicom AB, AstraZeneca, and Biovitrum AB, and as CEO of Arexis AB. Responsible for Industrifonden's life science operations from 2008 to 2016, and currently working as a senior advisor to the fund on a consultancy basis. He has held seats on the Boards of over 30 companies and is also a co-founder of two pharmaceutical development companies.
- **Other assignments:** Member of the Boards of InDex Pharmaceuticals AB, Calliditas Therapeutics AB and Cinclus Pharma Holding AB. Chairman of the Boards of Ignitus AB and Sixera Pharma AB.



BENGT JULANDER, Director since 2017

- **Education:** Pharmacist. Has worked in the pharmaceutical industry since 1978
- **Background:** CEO of Linc AB, which invests in life sciences. Since 1990, primarily active as an investor in and a Member of the Boards of pharmaceutical development companies. Experience of developing and commercialising products
- **Other assignments:** Member of the Boards of Linc AB, Livland Skog AB, Calliditas Therapeutics AB, Proequo AB, Sedana Medical AB, Stille AB and Swevet AB, and a number of smaller companies.



HELENA LEVANDER, Director since 2015

- **Education:** B.Sc. in Economics and Business Administration from the Stockholm School of Economics
- **Background:** Extensive experience of the financial and equity markets and of corporate governance issues. Previously employed by Neonet, Odin Förvaltning, Nordea Asset Management and SEB Asset Management, amongst others
- **Other assignments:** Founder and now Chairman of the Board of Nordic Investor Services AB. Member of the Boards of Concordia Maritime AB, Recipharm AB and Stendörren Fastigheter.



BENGT WESTERMARK, Director since 2017

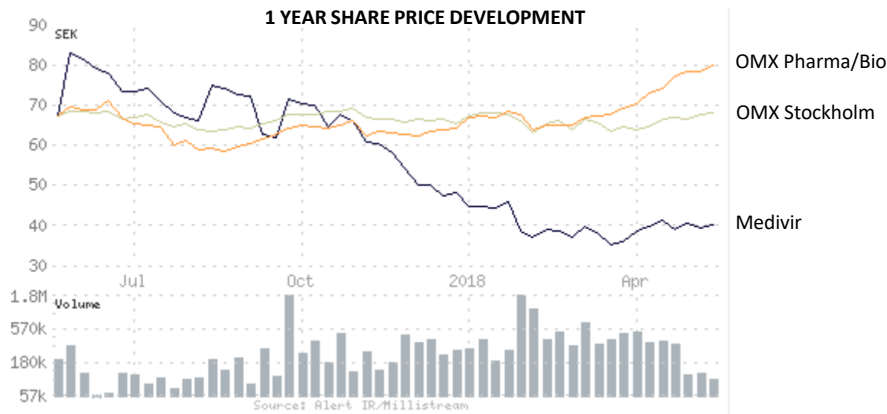
- **Education:** Professor of Tumour Biology at Uppsala University, the Faculty of Medicine, since 1986
- **Background:** Dean of the Faculty of Medicine at Uppsala University, 1996-2002, and Vice-Rector of Medicine and Pharmacy, 1999-2002. Chairman of the research board of the Swedish Cancer Society, 2003-2013. He has published over 300 papers in scientific journals, primarily on the mechanisms governing the uncontrolled growth of cancer cells. Member of the Royal Swedish Academy of Sciences, the European Molecular Biology Organisation, and the European Academy of Cancer Sciences. He has received a number of prizes and awards for his research and has been cited over 25,000 times by other researchers
- **Other assignments:** Member of the Board of Hamlet Pharma AB and several advisory groups for medical research financing

Board of directors with extensive scientific and operational experience of developing and leading businesses within biotech and pharma

The share and shareholder base

The Medivir share

- Single class of shares (B shares) outstanding totaling 24,287,818
- Traded on Nasdaq Stockholm Mid Cap, under the ticker MVIR B (ISIN: SE0000273294)
- Current market capitalization of SEK 1bn (USD 115m)¹



Shareholder base as of 30 April 2018

Owners	Class B shares	% of Capital
Nordea Investment Funds	1,996,367	8.22
MSIL IPB client account	1,731,060	7.13
Avanza Pension	1,168,730	4.81
Gladiator	1,050,000	4.32
Linc AB	958,283	3.95
UNIONEN	897,970	3.70
Hans Sköld	754,584	3.11
Credit Suisse SA	715,175	2.94
Ålandsbanken	609,231	2.51
Danica Pension	602,788	2.48
Nordnet Pensionsförsäkring AB	428,182	1.76
Bo Öberg	347,744	1.43
SEB life international Assurance	320,000	1.32
Rolf Kraft	245,325	1.01
Nils Gunnar Johansson	235,424	0.97
Total 15 largest shareholders	12,060,863	49.66
Total other shareholders	12,226,955	50.34
Total	24,287,818	100