

Focused on Growth and Innovation

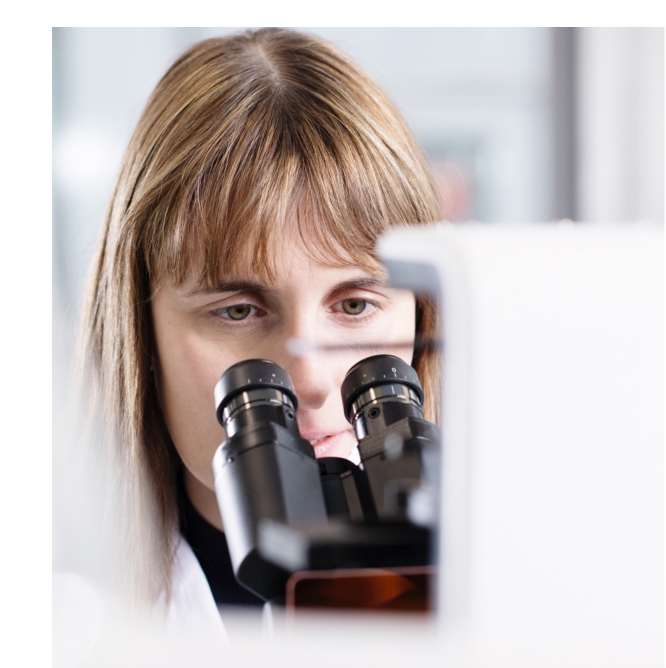
"Patients are at the heart of what we do"

Investor presentation August 2020



Table of contents

- Executive summary
- Five reasons to invest
- Portfolio
 - Antifungal
 - Cresemba® (isavuconazole)
 - Antibiotic
 - Zevtera[®] / Mabelio[®] (ceftobiprole)
 - Oncology
 - Derazantinib
 - Lisavanbulin (BAL101553)
- Financials
- Appendix



2



Executive summary



Experienced leadership team



(basilea)

At a glance

- Well funded, commercial-stage biotech company with significantly growing cash flows from commercialized products
- Focused in the areas of oncology and infectious diseases
- Potential for sustainable growth and value creation based on commercialized brands and an innovative pipeline
- Experienced people with the proven expertise to take compounds from research to market
- Two revenue generating hospital anti-infective brands, Cresemba[®] and Zevtera[®] and two clinical oncology drug candidates
- Recognized ability to establish and manage partnerships in both the development and commercial phase, providing access to international markets
- Listed on SIX Swiss Stock Exchange, SIX: BSLN
- Based in life sciences hub, Basel, Switzerland





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Potential for sustainable growth and value creation based on commercialized brands and innovative pipeline

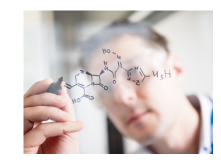
	Products / Product candidates / Indication	Preclinical	Phase 1	Phase 2	Phase 3	Market	
Antifungals	Cresemba® (isavuconazole) Invasive aspergillosis and mucormycosis (U.S. and EU and several other countries) Invasive fungal infections (Japan)	intravenous and oral intravenous and oral					
Antibiotics	Zevtera[®]/Mabelio[®] (ceftobiprole) Hospital- and community-acquired pneumonia (HAP, CAP) (major European and several non-European countries) Acute bacterial skin and skin structure infections (ABSSSI) Staphylococcus aureus (MSSA/MRSA) bacteremia (bloodstream infections)	intravenous intravenous intravenous					
Oncology	Derazantinib FGFR kinase inhibitor Intrahepatic cholangiocarcinoma (iCCA) – registrational study Urothelial cancer – monotherapy and combination with atezolizumab (Tecentriq®)* Gastric cancer (planned) Lisavanbulin (BAL101553) tumor checkpoint controller Glioblastoma – targeted, biomarker-driven phase 2 study (planned) Glioblastoma – combination with radiotherapy	oral oral oral oral oral					
	Internal & external innovation	Research	Development				

* Tecentriq® is a registered trademark of Hoffmann-La Roche Ltd.

Our strategy



Foster Foster an agile organisation based on a dynamic and open culture



Focus Focus on continuously increasing cash flow from our two commercial-stage hospital anti-infective brands, Cresemba[®] and Zevtera[®]



Leverage Leverage our expertise in bringing drugs from research to market by utilising appropriate partnerships with established organisations

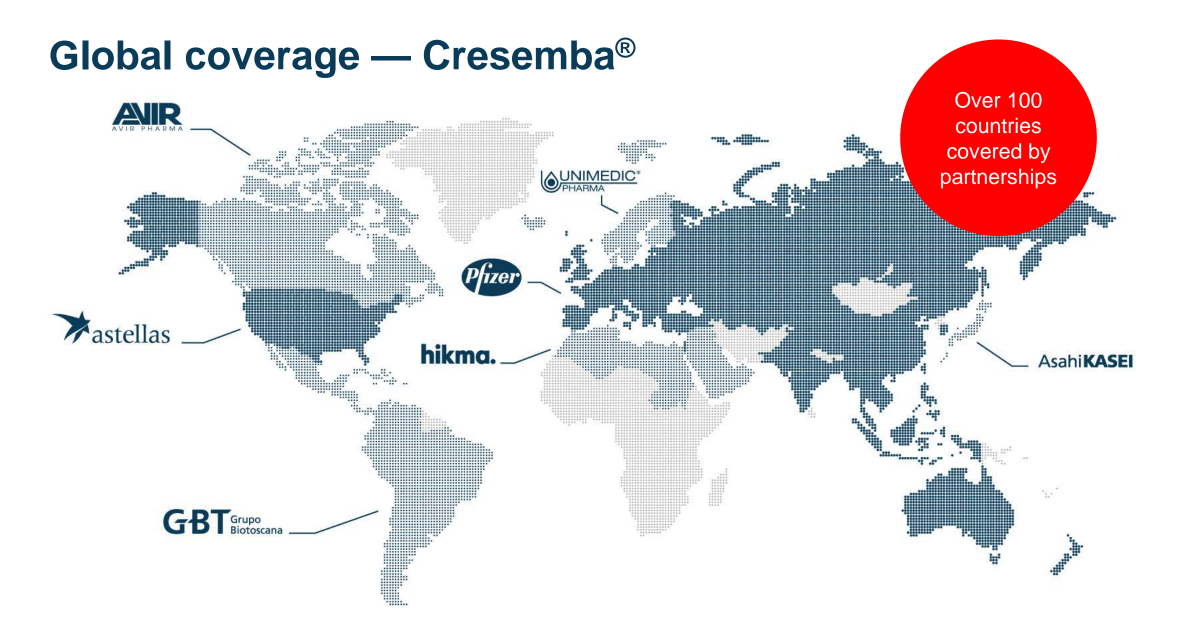


Invest Invest in our clinical portfolio of targeted, small molecule, oncology drug candidates and the phase 3 ceftobiprole program



Innovate Continue to broaden our R&D pipeline through both internal and external innovation





The company we keep — established strong partnerships





Five reasons to invest



Five reasons to invest



Growth Well funded with increasing and sustainable cash flow through commercialized brands



Prospects Opportunity to share in pipeline value creation and proven approach to the successful commercialization of products around the world



Leadership Experienced team working in an agile culture able to turn pipeline projects into revenue generating brands



Partnerships Proven ability to build successful partnerships in research, development and commercialization with leading academic, governmental and industrial organisations



Focus One of the few biopharmaceutical companies in the world focused on the development and commercialization of targeted oncology small molecules and new antibiotics and antifungals



Antifungal Cresemba® (isavuconazole)

Invasive mold infections

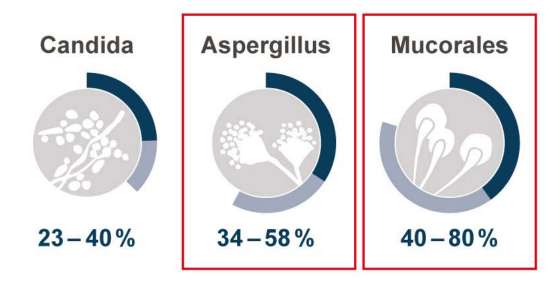
The market — Invasive fungal infections

- Severe, potentially life-threatening infections mainly affecting immunocompromised patients
- An important cause of morbidity and mortality in cancer patients undergoing intensive chemotherapy regimens
- Rising number of immunocompromised patients (cancer and transplantations) driving

therapeutic demand

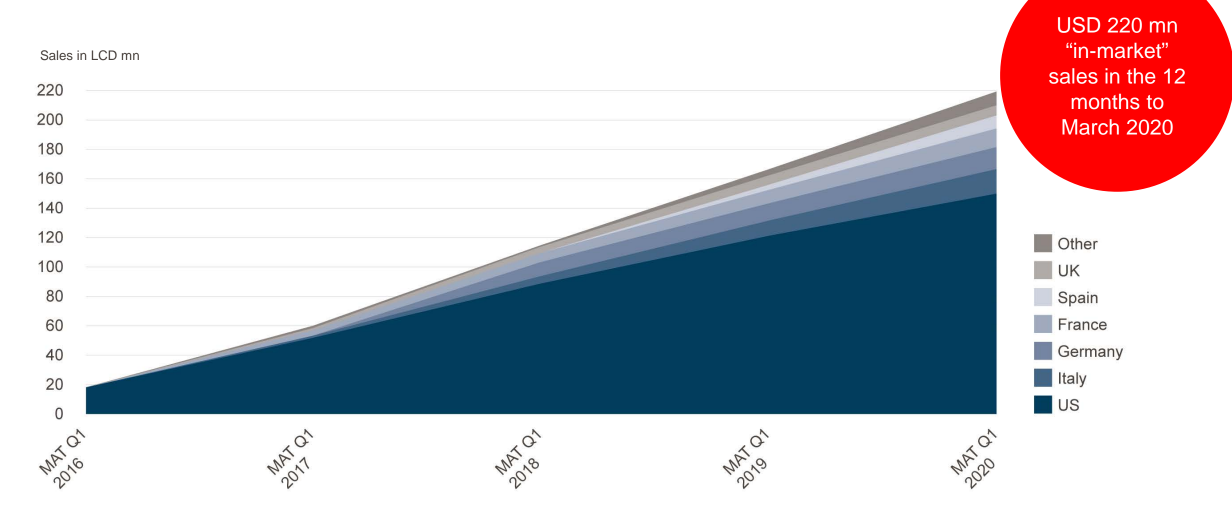
- Mucorales infections on the rise doubled from 2000 to 2013
- Limitations of current therapies (spectrum of activity, toxicity, effective plasma levels) drive the need for new agents

Mortality rates for invasive fungal infections**



**Kullberg/Arendrup *N Engl J Med* 2015, Baddley *Clin Infect Dis* 2010, Roden *Clin Infect Dis* 2005, Greenberg *Curr Opin Infect Dis* 2004

Cresemba continues strong in-market sales uptake



LCD: USD corrected for currency fluctuations; MAT: Moving annual total; Source: IQVIA, March 2020



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Sales of best-in-class antifungals* by product

USD 3.1 bn sales (MAT Q1 2020)

 Potential to increase Cresemba[®] (isavuconazole) market share

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- Anticipate to be launched in 60 countries by end-2021
- Exclusivity through 2027 in the U.S. and potential pediatric exclusivity extension to 2027 (from 2025) in the EU



* Best-in-class antifungals: isavuconazole, posaconazole, voriconazole, AmBisome, anidulafungin, caspofungin, micafungin

MAT: Moving annual total; Sales figures in USD, corrected for currency fluctuations; Source: IQVIA, March 2020

Cresemba[®] — Differentiated by spectrum, safety and tolerability

- Broad spectrum of activity against molds, including emerging molds (mucorales)
- Consistent plasma levels
- Statistically fewer drug-related adverse events and treatment-emergent adverse events (liver, skin, eye) in invasive aspergillosis patients vs. voriconazole in SECURE phase 3 study
- Can be administered without restriction in patients with renal impairment

- Manageable drug-drug interaction profile
- Once daily maintenance dose, i.v./oral treatment
- ECIL-6 guideline: Cresemba[®] recommended for the first-line treatment of invasive aspergillosis in leukemia and hematopoietic stem cell transplant patients. ECIL states that isavuconazole is as effective as voriconazole with a better safety profile.

Antibacterial Zevtera[®] / Mabelio[®] (ceftobiprole)

Severe bacterial infections

Zevtera[®] — An introduction

- Broad-spectrum anti-MRSA cephalosporin (including Gram-negative bacteria)
- Rapid bactericidal activity
- Potential to replace antibiotic combinations
- Early improvement in HAP, particularly in patients with MRSA, and CAP, including highrisk patients
- Cephalosporin class safety profile
- Marketed in selected countries in Europe, Latin America and the MENA-region as well as in Canada

Approved in major European countries & several non-European countries for both hospitalacquired pneumonia (HAP), excluding ventilator-associated pneumonia (VAP), and community-acquired pneumonia (CAP). Not approved in the U.S.

MENA: Middle East and North Africa



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for solution for infusion

Ceftobiprole (as ceftobiprole medocaril sodium).

equivalent to 666.6 mg of ceftobiprole medocaril sodium.

Each vial contains 500 mg of ceftobiprole,

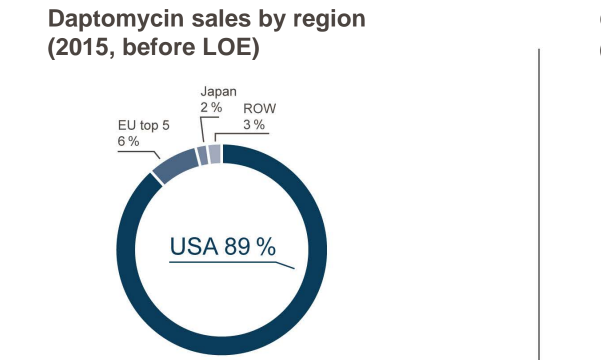
For intravenous use after reconstitution and dilution.

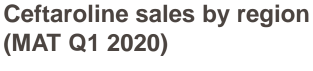
Read the package leaflet before use.

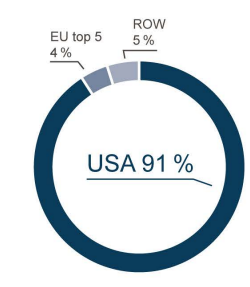
Chariles

10 vials

The hospital anti-MRSA antibiotic market — A USD 3 bn market* with the U.S. being the most important region







* Vancomycin, linezolid, teicoplanin, daptomycin, tigecycline, telavancin, ceftaroline, dalbavancin, oritavancin, and tedizolid

MRSA: Methicillin-resistant *Staphylococcus aureus*; LOE: Loss of exclusivity; ROW: Rest of world MAT: Moving annual total; Sales figures in USD, corrected for currency fluctuations; Source: IQVIA, March 2020

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Strategy for accessing the U.S. market

- Two cross-supportive phase 3 studies under FDA Special Protocol Assessment (SPA)
 - Acute Bacterial Skin and Skin Structure Infections (ABSSSI)¹ successfully completed



2. Staphylococcus aureus bacteremia (SAB)² ongoing, topline results from phase 3 study expected in Q1 2022



 Phase 3 program largely funded by BARDA (up to USD 128 mn, ~70% of total program costs)



 Qualified Infectious Disease Product (QIDP) designation extends U.S. market exclusivity to 10 years from approval

¹ Overcash JS et al. ECCMID 2020, abstract 1594. (NCT03137173) ² Hamed K et al. Future Microbiol. 2020;15:35-48. (NCT03138733)

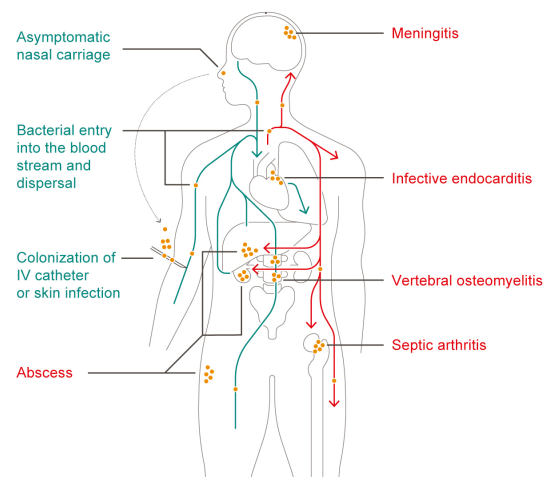


SAB – an area with high medical need

- Nearly 120,000 S. aureus bloodstream infections in the US (in 2017)¹
- ERADICATE targets complicated SAB, characterized by concomitant or metastatic infections such as bone, joint or heart valve infections; persistent bacteremia; or bacteremia in patients on dialysis
- Substantial morbidity and approximately 20%
 30-day mortality²
- Limited antibiotic treatment options with only two approved treatments for SAB in the U.S. that cover both MSSA and MRSA, i.e. vancomycin and daptomycin

¹ MMWR, 2019;68:214–219.

² Hamed K et al. Future Microbiol. 2020;15:35-48. MRSA: methicillin-resistant *Staphylococcus aureus* MSSA: methicillin-susceptible *Staphylococcus aureus*



Causes and consequences of SAB

Adapted from Edwards AM et al. Trends Microbiol. 2011;19:184-190.



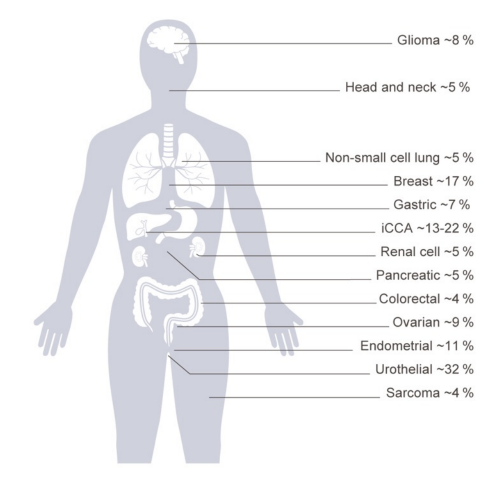
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Oncology Derazantinib

FGFR-driven tumors

Targeting FGFR-driven tumors as single agent and in combination with immunotherapy

- Small molecule, oral inhibitor of FGFR family of kinases
- Development strategy focused on achieving differentiation by leveraging unique properties of derazantinib
 - Kinase inhibition profile: exploring therapeutic potential of additional targets of derazantinib such as CSF1R and VEGFR2 kinase
 - Safety profile: exploring relevance for potential combination therapies
- Two clinical studies ongoing (FIDES-01 in iCCA & FIDES-02 in urothelial cancer)
- Plan to start a multi-cohort phase 1/2 study (FIDES-03) in patients with advanced gastric cancer in Q3 2020



Sources: Helsten et al., Clin Cancer Res 2016 (22), 257-267; FGFR2 fusions in iCCA: Graham et al. Hum Pathol 2014 (45), 1630-1638; Jain et al. JCO Precis Oncol 2018 (2) 1-12

Registrational phase 2 study in iCCA (FIDES-01)¹

Cohort 1: Patients with FGFR2 gene-fusion expressing iCCA (2nd line)

- Encouraging interim results, consistent with earlier phase 1/2 data²
- 21% ORR with six confirmed partial responses from 29 evaluable patients, 83% disease control rate
- Completed patient enrolment in July 2020
- Topline results expected H2 2020

Cohort 2: Patients with FGFR2 gene mutations or amplifications

- Assessing the activity of derazantinib in a broader range of FGFR2-driven tumors
- Clinical benefit observed in a subset of iCCA patients in the phase 1/2 study²
- Aim to confirm phase 1/2 study results in a larger cohort of iCCA patients¹
- Define the full therapeutic potential of derazantinib in iCCA with potential for differentiation
- Interim results expected H2 2020

²Droz Dit Busset et al. Annals of Oncology (2019) 30 (suppl_5): abstract 3879 (NCT01752920)

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Clinical program in urothelial and gastric cancer

FIDES-02¹ | Urothelial Cancer

Multi-cohort Phase 1b/2 study of derazantinib monotherapy or in combination with atezolizumab (Tecentriq®) in patients with urothelial cancer expressing activating molecular FGFR aberrations

- Substudies (N≈300) in various treatment settings, including:
 - Post-chemotherapy/immunotherapy recurrence (second-line and post second-line)
 - First-line platinum-ineligible, PD-L1-low
 - Resistance to prior FGFR-inhibitor treatment
- First interim results expected in H2 2020

FIDES-03 | Gastric Cancer

Multi-cohort Phase 1b/2 study of derazantinib as monotherapy or in combination therapy with standard of care or atezolizumab in patients with advanced HER2-negative gastric adenocarcinoma harboring FGFR genetic aberrations

- Substudies using derazantinib monotherapy or combination treatment, including:
 - Derazantinib monotherapy in various molecular subtypes
 - Combination of derazantinib and standard of care
 - Combination of derazantinib with atezolizumab (Tecentriq[®])
- Expected start of enrolment in Q3 2020

¹NCT04045613



FGFR-inhibitors show differences in safety profiles

	Cholangiocarcinoma				Urothelial cancer		
	DZB ¹ (N=44)	INF ² (N=71)	FUT ³ (N=67)	PEM ⁴ (N=146)	PEM ⁵ (N=108)	ERD ⁶ (N=87)	
Dosing regimen	300mg QD	125mg Q4W QD for 3w	20 mg QD	13.5mg Q3W QD for 2w	13.5mg Q3W QD for 2w	8 mg QD (titration to 9mg)	
Most frequent safety events	Phosphorus û Nausea Vomiting	Phosphorus û Fatigue Stomatitis	Phosphorus*û Diarrhea* Dry mouth*	Phosphorus û Alopecia Diarrhoea	Diarrhoea Alopecia Constipation	Phosphorus û Stomatitis Fatigue	
Blood phosphorus û†	59%	73%	88%	60%	31%	76%	
Fatigue [†]	43%	49%	NR	42%	32%	54% [#]	
Alopecia [†]	20%	38%	NR	49%	40%	26%	
Dry eye/xerophthalmia [†]	16%	32%	NR	35%#	NR	28% [#]	
Retinopathy [¶]	0%	NR	9%	6% ‡	NR	25%	
Alanine aminotransferase (ALT) 企	30%**	NR	NR	43%**	NR	41%**	
Hand-foot syndrome/PPE	0%	27%	18%	15%	NR	26%	
Nail toxicities	<5%	NR	42%	43% [#]	NR	41% [#]	
Stomatitis	11%	45%	NR	35%	34%	56%	

¹ Droz Dit Busset et al., ESMO 2019 and Basilea data on file, ² Javle et al., ESMO 2018, ³ Goyal et al., ASCO 2020, ⁴ PemazyreTM U.S. Prescribing Information (April 2020), ⁵ Necchi, et al., ESMO 2018, ⁶ BalversaTM U.S. prescribing information (April 2019)

† assumed FGFR inhibitor class-effect; *futibatinib treatment-related adverse events

* includes various and different adverse reactions; for details see PemazyreTM U.S. Prescribing Information (April 2020) and BalversaTM U.S. prescribing information (April 2019);

[¶]Refers to reported adverse events of Retinal Pigment Epithelial Detachment (RPED) for pemigatinib, Central Serous Retinopathy (CSR)/RPED for erdafitinib and CSR for futibatinib

[‡] reported incidence is from 466 patients who received PemazyreTM across clinical trials;

** based on reported adverse events for DZB; based on reported laboratory abnormalities, regardless of causality for PEM and ERD.

Abbreviations: DZB: derazantinib, INF: infigratinib (BGJ398), FUT: futibatinib (TAS-120), PEM: pemigatinib (INCB54828), ERD: erdafitinib; PPES: Palmar-plantar erythrodysesthesia; NR: not reported; QD: daily; Q3W/Q4W: every 3/4 weeks; w: weeks

Oncology Lisavanbulin (BAL101553)

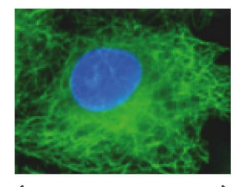
Glioblastoma and other solid tumors

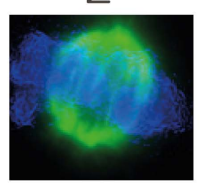


Novel tumor checkpoint controller crossing the blood-brain barrier

- Novel compound inducing tumor cell death through spindle assembly checkpoint activation
- Targeting diverse tumor types resistant to standard therapeutic approaches
- Flexible dosing potential, including daily oral dosing
- Comprehensive biomarker program to optimize patient selection
- Crosses the blood-brain barrier with potent activity in brain tumor models alone and in combination
- Clinical program focused on glioblastoma (GBM) using a biomarker-driven approach

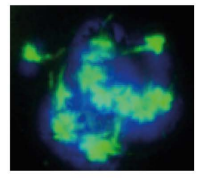
Non-dividing tumor cell





Blue = DNA

Green = microtubules



Normal dividing tumor cell

BAL27862-treated tumor cell*

* Lisavanbulin (BAL101553) is a prodrug of BAL27862



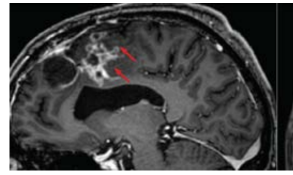
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EB1 — A potential response-predictive clinical biomarker for lisavanbulin

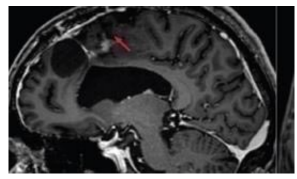
- EB1 (plus-end binding protein) is located on the microtubules and involved in microtubule dynamics and has been shown to be a response predictive marker for lisavanbulin in preclinical studies
- Strong EB1 staining was observed in a patient with an exceptional response to daily oral lisavanbulin in the phase 1 dose-escalation study in recurrent GBM¹
 - Patient ongoing for more than two years
 - >80% reduction in GBM tumor size
- Biomarker-driven phase 2 study in patients with recurrent GBM using EB1-positivity as patient selection criterion, anticipated to start in the next few months

¹ Lopez et al. Phase 1/2a study of once daily oral BAL101553, a novel tumor checkpoint controller, in adult patients with progressive or recurrent glioblastoma or high-grade glioma. JCO 2019;37:15 suppl, 2025 (NCT02490800)

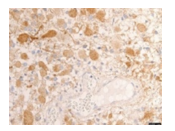
GBM tumor size reduction in an exceptional responder and EB1 staining of GBM tissue compared to non-responding patients



Baseline (May 2018)



Post Cycle 12 (April 2019)



Responder



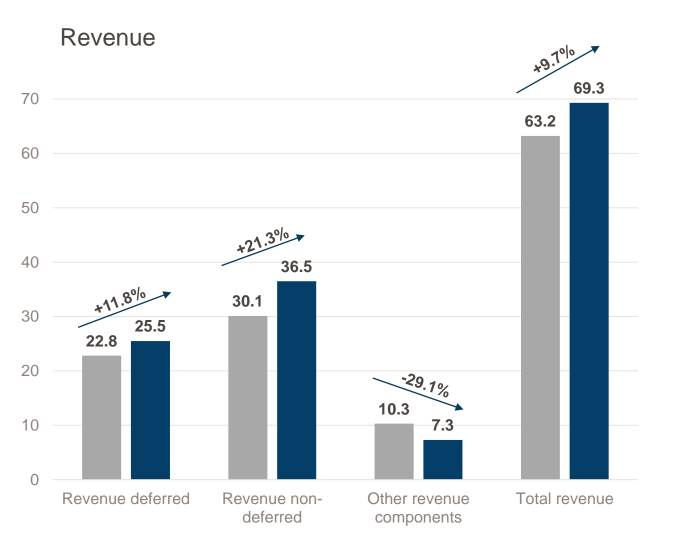
Non-responder

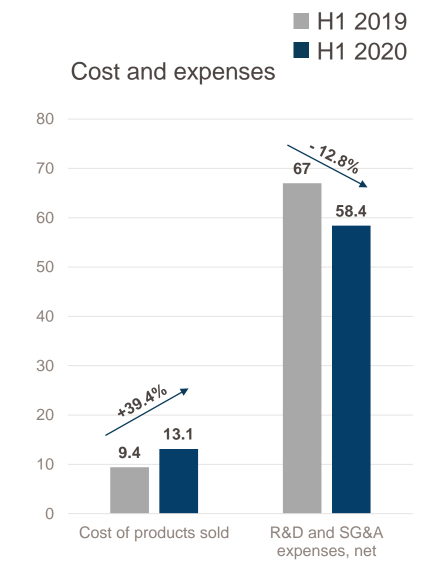


Financials



Financial summary, in CHF mn (1/2)

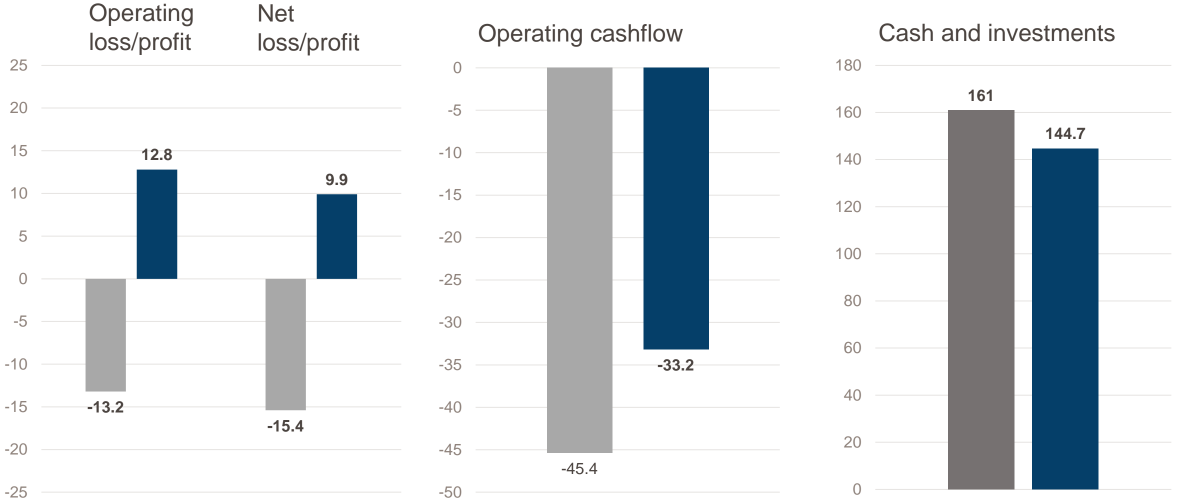




Note: Consolidated figures in conformity with U.S. GAAP; rounding applied consistently

Financial summary, in CHF mn (2/2)





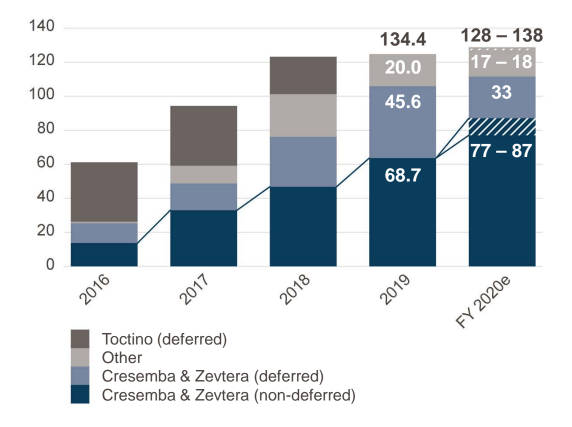
Note: Consolidated figures in conformity with U.S. GAAP; rounding applied consistently

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Financial guidance

In CHF mn	FY 2020e	FY 2019
Total revenue	128 – 138	134.4
thereof: Contributions Cresemba [®] & Zevtera [®] non-deferred deferred	77–87 33	68.7 45.6
Operating loss	5-15	17.2
Cash and investments	150	161.0

Strong increase in non-deferred revenue contributions Y-o-Y, CHF mn



Outlook 2020 / 2021

Cresemba[®] & Zevtera[®] — Increasing cash flows By the end of 2021, Cresemba to be on the market in 60 countries

			H1 2020	H2 2020	H1 2021	H2 2021
Isavuconazole				Complete patient enrolment in phase 3 study in Japan		Topline results from phase 3 study in Japan
Ceftobiprole						Complete patient enrolment in SAB phase 3 study
	FIDES-01 (iCCA)	\checkmark	Complete patient enrolment in phase 2 registrational study (FGFR2 fusions)	Topline results (FGFR2 fusions)		
				Interim results (other FGFR2 gene aberrations)		Topline results (other FGFR2 gene aberrations)
Derazantinib	FIDES-02 (urothelial cancer)			Safety data and recommended phase 2 dose (RP2D) for derazantinib/Tecentriq combination and expansion into phase 2	Interim results in derazantinib monotherapy	Interim results in combination therapy with Tecentriq
	FIDES-03 (gastric cancer)	\checkmark	Clinical supply agreement with Roche in gastric cancer	Start of phase 1/2 study		Interim results
Lisavanbulin		\checkmark	Full results of phase 1 study in glioblastoma*	Start phase 2 biomarker-driven glioblastoma study	Interim results from phase 2 biomarker-driven glioblastoma study	Topline results from phase 2 biomarker-driven glioblastoma study
(Oral)					Complete patient enrolment in phase 1 study in newly diagnosed glioblastoma	

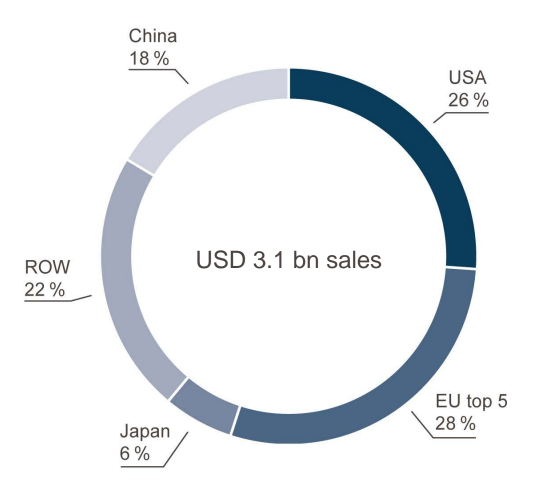
* Accepted for ESMO poster presentation (Sept. 2020)

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Appendix

Significant sales of bestin-class antifungals in all major regions — Covered by our partnerships

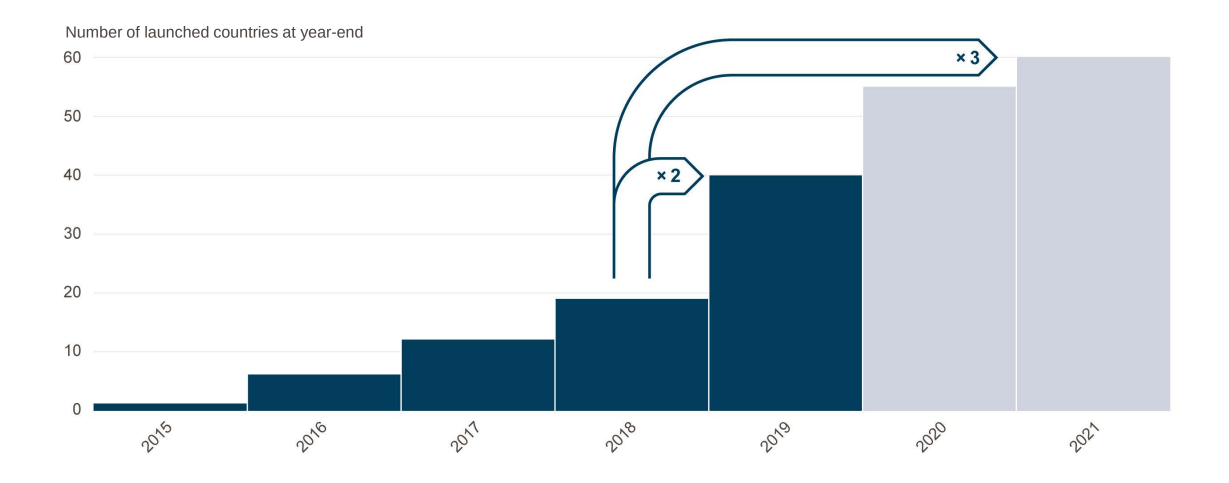
USD 3.1 bn sales of best-in-class antifungals* (MAT Q1 2020)



* Best-in-class antifungals: isavuconazole, posaconazole, voriconazole, AmBisome, anidulafungin, caspofungin, micafungin

MAT: Moving annual total; Sales figures in USD, corrected for currency fluctuations; Source: IQVIA, March 2020

Cresemba[®] — Strong global roll out



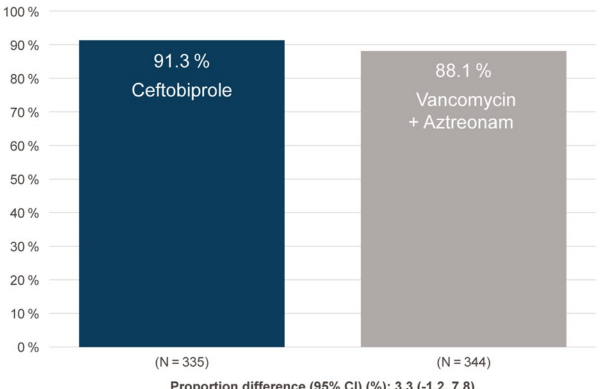
Ceftobiprole — **Positive** topline phase 3 results reported in ABSSSI

Key topline study¹ results showing non-inferiority of ceftobiprole to vancomycin plus aztreonam for the primary and secondary endpoints



Early clinical response at 48–72h after start of treatment (ITT population)

Patients with early clinical success at 48-72 hours (%)



Proportion difference (95% CI) (%): 3.3 (-1.2, 7.8)

¹NCT03137173 ABSSSI: Acute bacterial skin and skin structure infections

(basilea) Focused on Growth and Innovation ITT: intent-to-treat Pre-defined limit of non-inferiority = lower limit of 95 % CI for difference > -10 %

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Ceftobiprole — **Positive** topline phase 3 results reported in ABSSSI

Key topline study¹ results showing non-inferiority of ceftobiprole to vancomycin plus aztreonam for the primary and secondary endpoints

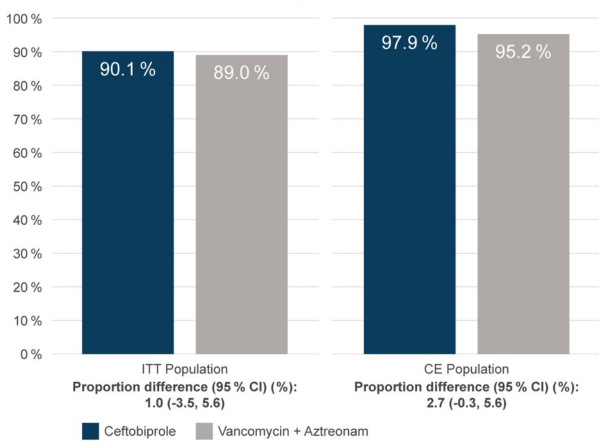


¹NCT03137173 ABSSSI: Acute bacterial skin and skin structure infections

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Investigator-assessed clinical success at test-of-cure (TOC) 15-22 days after randomization (ITT, CE populations)

Patients with clinical success at the TOC visit (%)



CE: clinically evaluable; ITT: intent-to-treat

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39

Ceftobiprole key attributes for SAB treatment

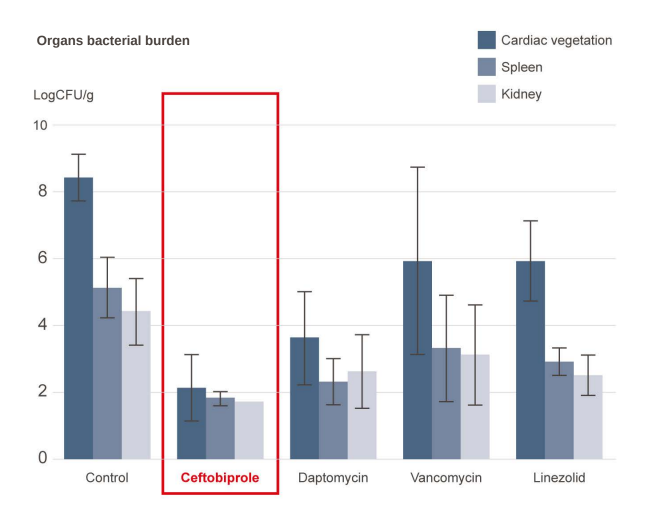
- Beta-lactam antibiotic with rapid bactericidal activity against MSSA and MRSA¹
- Superior activity profile in preclinical models of endocarditis compared to vancomycin and daptomycin²
- Low propensity for resistance development¹
- Gram-negative coverage¹ in cases with polymicrobial infections
- Efficacy demonstrated in Phase 3 clinical trials in pneumonia and complicated skin and soft tissue infections^{1,3,4}
- Established safety profile consistent with the cephalosporin class^{1,3}

¹Syed YY. Drugs. 2014;74:1523-1542.

²Tattevin P et al. Antimicrob Agents Chemother. 2010;54:610-613.
³Giacobbe DR et al. Expert Rev Anti Infect Ther. 2019;17:689-698.
⁴Overcash JS et al. ECCMID 2020, abstract 1594

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Comparative efficacy in a rabbit model of endocarditis



Organism titers in cardiac vegetations, spleens and kidneys of untreated and antibiotic treated rabbits infected with MRSA²

40

Phase 3 study with ceftobiprole in the treatment of patients with SAB



- Design: randomized, double-blind, multicenter
- Enrolment: approximately 390 adult patients (male and female)
- Indications: Staphylococcus aureus bacteremia (SAB), including endocarditis (IE) and other forms of complicated SAB
- Main inclusion criteria: Positive S. aureus
 blood culture and signs & symptoms for SAB
- Intervention: ceftobiprole medocaril i.v.; comparator daptomycin i.v. or daptomycin plus aztreonam to cover Gramnegative bacteria

- Primary endpoint: overall success as assessed by an independent Data Review Committee (DRC) in the treatment of SAB, including IE, at the post-treatment evaluation (PTE) visit (70 days after randomization) in the modified intent-to-treat (mITT) population.
- Secondary endpoints: includes all-cause mortality at Day 28 and Day 70 (PTE visit) in the intent-to-treat (ITT) and mITT populations; and time to *S. aureus* bloodstream clearance

NCT03138733



FGFR-inhibitors show differences in kinase-inhibition profiles¹

FGFR-inhibitor compound (Sponsor)	Parameter	FGFR1	FGFR2	FGFR3	FGFR4	CSF1R	VEGFR2
Derazantinib (Basilea)	Ratio to FGFR2 activity	4	1	4	77	3	6
Pemigatinib (Incyte)	Ratio to FGFR2 activity	3	1	4	39	231	62
Erdafitinib (Janssen)	Ratio to FGFR2 activity	2	1	2	13	95	6
Rogaratinib (Bayer)	Ratio to FGFR2 activity	5	1	6	18	116	48
Infigratinib (QED)	Ratio to FGFR2 activity	2	1	2	47	86	55
Futibatinib (Taiho)	Ratio to FGFR2 activity	2	1	2	18	NA	NA

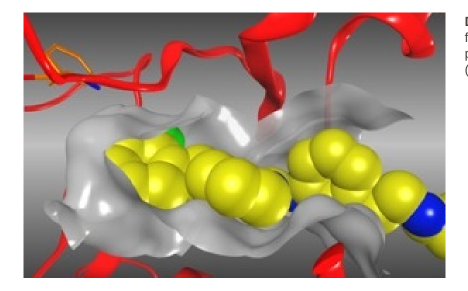
¹ McSheehy et al. Derazantinib (DZB): A dual FGFR/CSF1R-inhibitor active in PDX-models of urothelial cancer. Mol Cancer Ther. 2019 (18) (12 Supplement) LB-C12

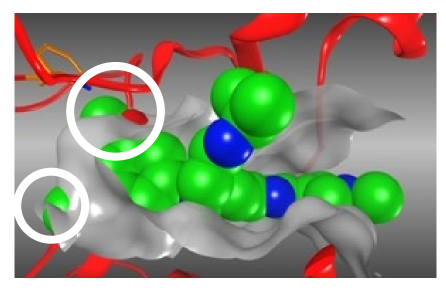
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In-silico analysis of derazantinib binding to CSF1R

- Crystal structures indicate differences in inhibitor binding sites of FGFR and CSF1R kinases
- Improved kinase inhibition activity of derazantinib against CSF1R versus other FGFR-inhibitors can be explained by the unique chemical structure of derazantinib¹





Derazantinib (yellow) fits to smaller active site pocket of CSF1R (grey/red)

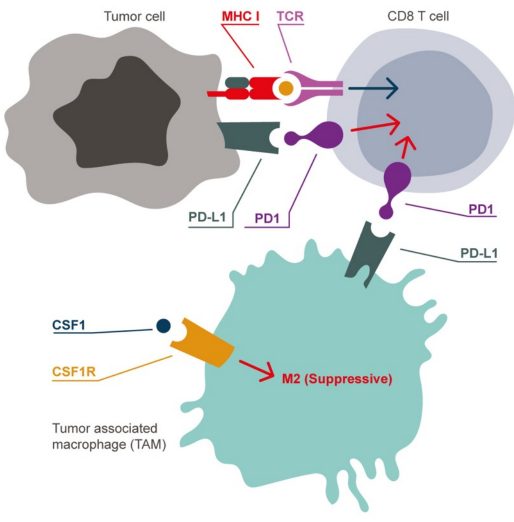
Erdafitinib (green) is too large (white circles) for the active site pocket of CSF1R (grey/red)

¹ McSheehy et al. Derazantinib (DZB): A dual FGFR/CSF1R-inhibitor active in PDX-models of urothelial cancer. Mol Cancer Ther. 2019 (18) (12 Supplement) LB-C12

Potential therapeutic relevance of CSF1Rinhibition

- Derazantinib is active in inhibiting FGFR kinases and CSF1R (Colony-stimulating factor-1 receptor)
- CSF1R-inhibition may reprogram immunosuppressive tumor-infiltrating macrophages, restore T-cell activity and thereby improve the susceptibility to PD1/PD-L1 inhibitors¹
- Derazantinib may address several oncogenic mechanisms at the same time, i.e. inhibiting FGFR and making the tumor more susceptible to immunotherapy
- Basilea entered into a clinical supply agreement with Roche to study a combination of derazantinib and Roche's PD-L1blocking immune-checkpoint inhibitor atezolizumab (Tecentriq[®]) in patients with urothelial cancer

Tumor microenvironment

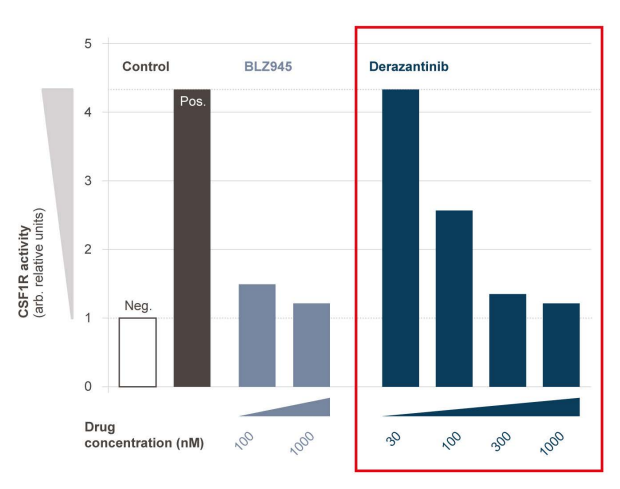


Graph adapted from: A. Ghasemzadeh et al. New Strategies in Bladder Cancer: A Second Coming for Immunotherapy. Clin Cancer Res. 2016;22(4):793-801

Derazantinib inhibits mouse macrophage CSF1R activity

- Derazantinib treatment reduced CSF1stimulated CSF1R activation (pCSF1R) in a concentration-dependent manner
- The maximum effect is similar to the specific CSF1R inhibitor BLZ945
- Derazantinib active-concentration is achievable in patients

Inhibition of CSF1R activity



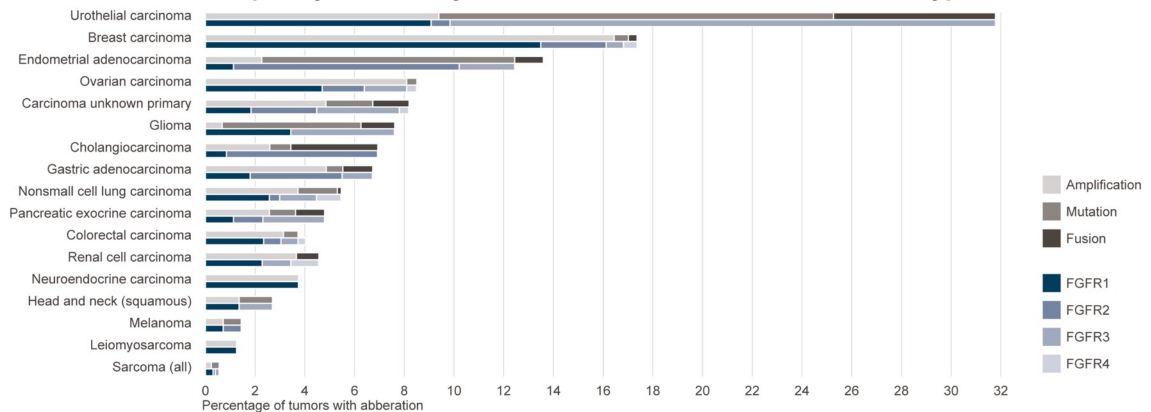
Method: bone-marrow-derived mouse macrophages were starved overnight, treated with CSF1 for 3 min, with or without pre-incubation with BLZ945 or DZB, and then extracted for subsequent immunoblot. The graph shows the quantification of the experiment based on densiometric analysis of the immunoblots

¹ McSheehy et al. Derazantinib (DZB): A dual FGFR/CSF1R-inhibitor active in PDX-models of urothelial cancer. Mol Cancer Ther. 2019 (18) (12 Supplement) LB-C12

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Derazantinib — Significant potential beyond iCCA



Frequency of currently known FGFR aberrations across tumor types

Source: Helsten et al., Clin Cancer Res. 2016;22:259-67

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Derazantinib — Multi-cohort phase 1/2 study in advanced urothelial cancer (FIDES-02)¹

- Derazantinib as single agent and in combination with atezolizumab (Tecentriq[®]) in patients with advanced urothelial cancer testing positive for mutations or fusions of FGFR1, FGFR2 or FGFR3 genes
- The subgroup of patients with low PD-L1 expression have limited clinical benefit from the treatment with PD1/PD-L1 inhibitors. This subgroup, however, shows frequent FGFR genomic abnormalities (mainly FGFR3 fusions)
- Derazantinib combined with PD1/PD-L1 inhibitors may provide benefits related to multiple mechanisms (FGFR-inhibition, macrophage modulation, enhanced response to immunotherapy), in particular in the low PD-L1 expression subgroup

- Across a total of four sub-studies, FIDES-02 potentially can enroll up to approximately 300 patients
- Patient cohorts in various treatment settings, including:
 - Post-chemotherapy/immunotherapy recurrence (second-and post second-line)
 - First-line platinum-ineligible
 - Resistance to prior FGFR-inhibitor treatment
- Study conducted in multiple centers in Asia-Pacific, Europe and North America
- Clinical supply agreement with Roche for the immune-checkpoint inhibitor atezolizumab (Tecentriq[®])

¹NCT04045613

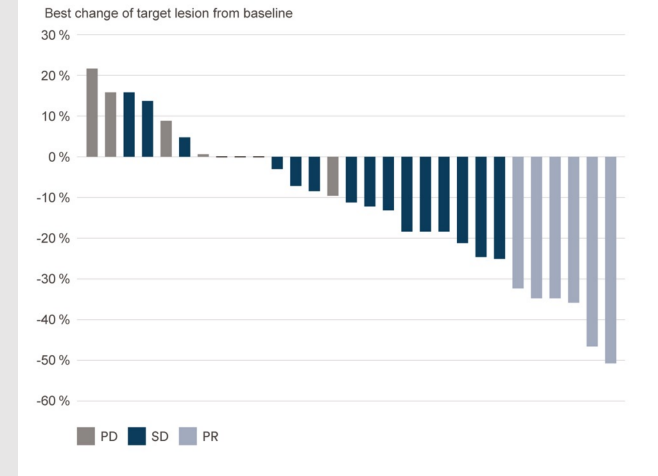


Derazantinib — Established proof-of-concept in iCCA in phase 1/2a study

- Subgroup analysis of 29 patients with FGFR2-fusion positive iCCA:
 - Objective response rate of 21%
 - In 72% of patients, tumor response or disease stabilization for ≥16 weeks was achieved*
- Compares favorably to Standard-of-Care (SoC) chemotherapy (cross-trial comparison)
 - Objective Response Rate (ORR) 21% for derazantinib¹ versus <10% for SoC^{2, 3}
 - Progression-Free Survival (PFS) approx. 6 months¹
 versus 3 months for SoC^{2, 3}
- Manageable safety profile^{1, 4}

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¹ V. Mazzaferro et al. Derazantinib (ARQ 087) in advanced or inoperable FGFR2 gene fusion-positive intrahepatic cholangiocarcinoma. British Journal of Cancer 2018 ² A. Lamarca et al. Second-line chemotherapy in advanced biliary cancer: a systematic review. Annals of Oncology 2014 (25), 2328-2338; ³ L. Fornaro et al. Second-line chemotherapy in advanced biliary cancer progressed to first-line platinum-gemcitabine combination: a multicenter survey and pooled analysis with published data. Journal of Experimental & Clinical Cancer Research 2015 (34), 156 ⁴ K. P. Papadopoulos et al. A phase 1 study of ARQ 087, an oral pan-FGFR inhibitor in patients with advanced solid tumors. British Journal of Cancer 2017, 1-8



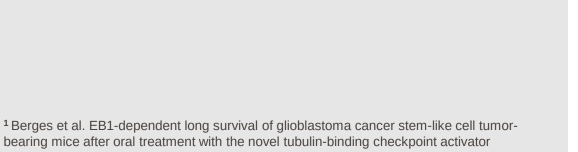
Sources: Mazzaferro et al. British Journal of Cancer 2018; * Mazzaferro et al. J Clin Oncol 2017;35 suppl: abstract 4017

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48

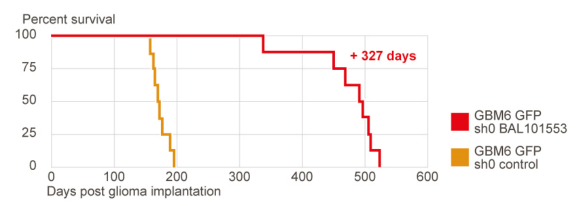
EB1 — A potential response-predictive clinical biomarker for lisavanbulin

- EB1 (plus-end binding protein)¹ is located on the microtubules and involved in microtubule dynamics
- Predictive of response to lisavanbulin in mouse models¹

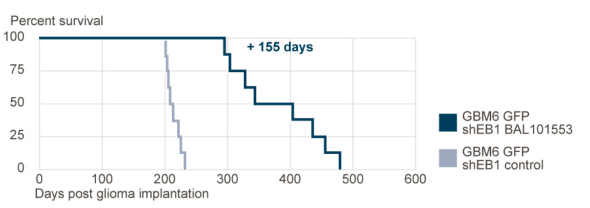


Effect of lisavanbulin (BAL101553) on survival in mice with EB1-expressing or EB1 downregulated GBM

EB1-expressing GBM







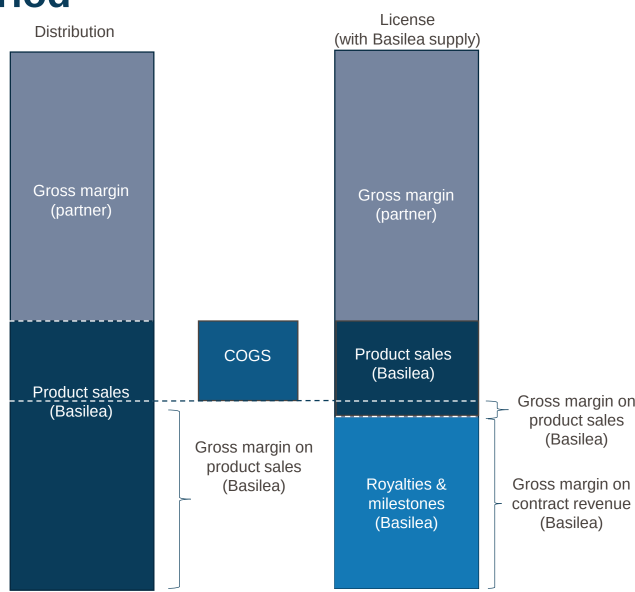
bearing mice after oral treatment with the novel tubulin-binding checkpoint activator BAL101553. Eur. J. Cancer 2018, 103, E61-62, A166

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Extension of Pfizer supply period

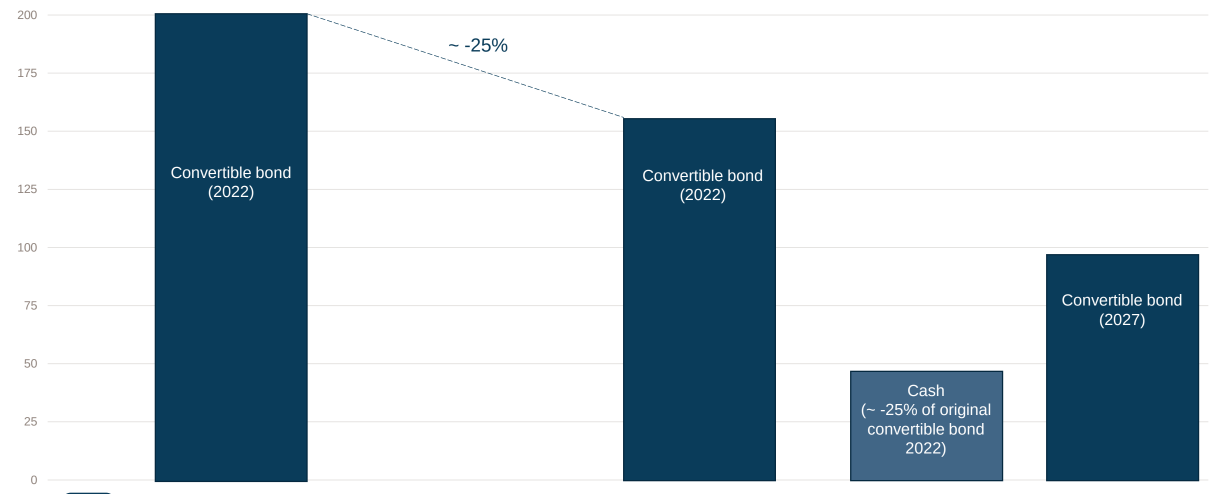
- Supply API and bulk Cresemba vials 2020/2021
 - Increase in product sales (in CHF)
 - Increase in cost of products sold (in CHF); economies-of-scale in supply to other partners
 - Lower gross margin (in % of product sales)
 - Temporary increase in working capital
- => Net positive cash flow over 2020/2021



Convertible bond transactions — successfully improved debt maturity profile (in CHF mn)

Pre-transactions

Post-transactions



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Glossary

- ABSSSI: Acute bacterial skin and skin structure infections
- CSF1R: Colony-stimulating Factor 1 Receptor
- FGFR: Fibroblast Growth Factor Receptor
- GBM: Glioblastoma multiforme
- iCCA: Intrahepatic cholangiocarcinoma
- MRSA: methicillin-resistant *Staphylococcus aureus*
- MSSA: methicillin-susceptible *Staphylococcus aureus*
- SAB: *Staphylococcus aureus* bacteremia
- VEGFR2: Vascular Endothelial Growth Factor Receptor 2

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Grenzacherstrasse 487 PO Box 4005 Basel Switzerland

investor_relations@basilea.com www.basilea.com

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