

"Patients are at the heart of what we do"

David Veitch CEO

Kepler Cheuvreux Digital Life Science Days June 21, 2021



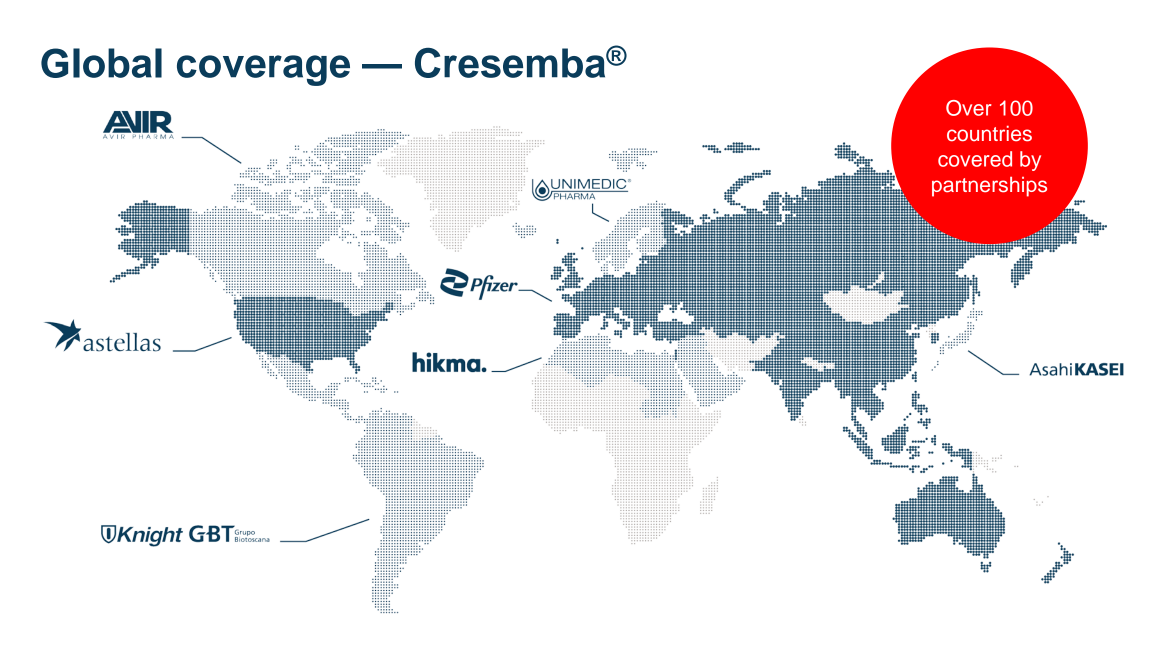
At a glance

- Well funded, commercial-stage biopharmaceutical 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



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 a				
Antibiotics	Zevtera® (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				
Oncology	Derazantinib FGFR kinase inhibitor Intrahepatic cholangiocarcinoma (iCCA) – monotherapy Urothelial cancer – monotherapy and combination with atezolizumab Gastric cancer - monotherapy and combination with ramucirumab/paclitaxel or atezolizumab Lisavanbulin (BAL101553) tumor checkpoint controller Glioblastoma – monotherapy, targeted, biomarker-driven patient selection Glioblastoma – combination with radiotherapy	oral oral oral oral				
	Internal & external innovation	Research	Development			



The company we keep — established strong partnerships

License partners





Europe (excl. Nordics), China Asia-Pacific, Russia, Turkey and Israel (Cresemba®)

U.S. (Cresemba®)

Asahi **KASEI**

Japan (Cresemba®)



Distribution partners



Europe (excl. Nordics), Israel (Zevtera®)



Nordics (Cresemba® and Zevtera®)

hikma.

MENA region (Cresemba® and Zevtera®)

(Cresemba® and Zevtera®)

UKnight GBTGrupo Biotoscana

(Cresemba® and Zevtera®)

LatAm





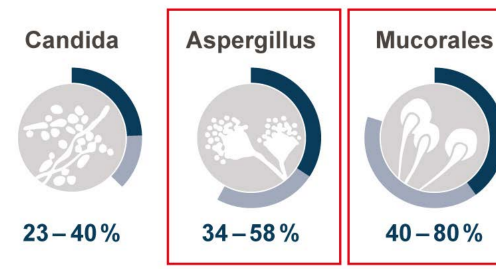
Canada



The market — Invasive fungal infections

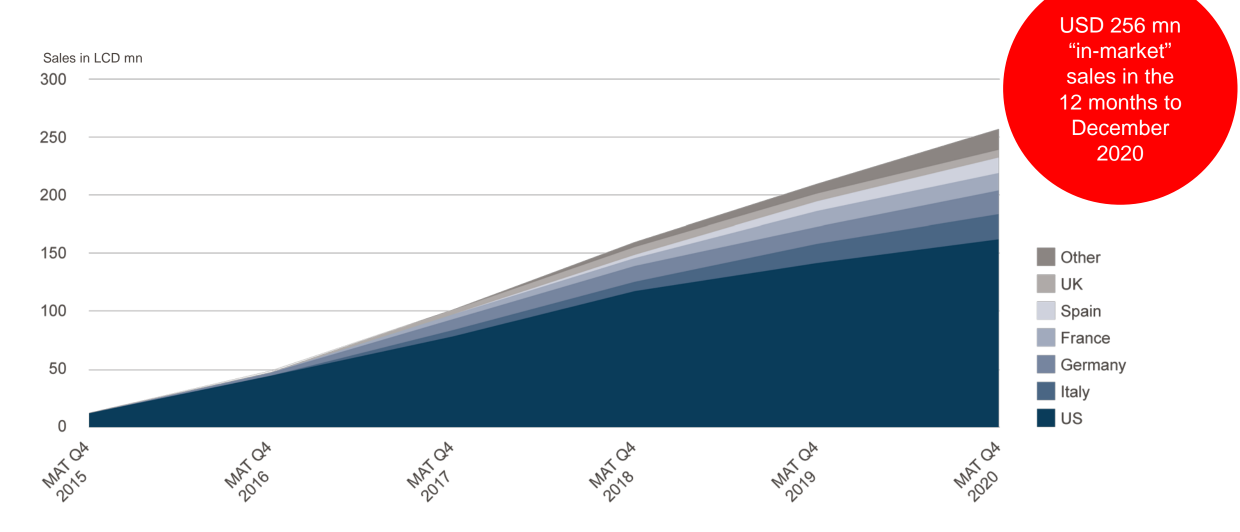
- Severe, potentially life-threatening infections mainly affecting immunocompromised patients
- An important cause of morbidity and mortality in cancer patients undergoing intensive chemotherapy regimens
- Rising number of immunocompromised patients
 (cancer and transplantations) driving therapeutic 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, December 2020

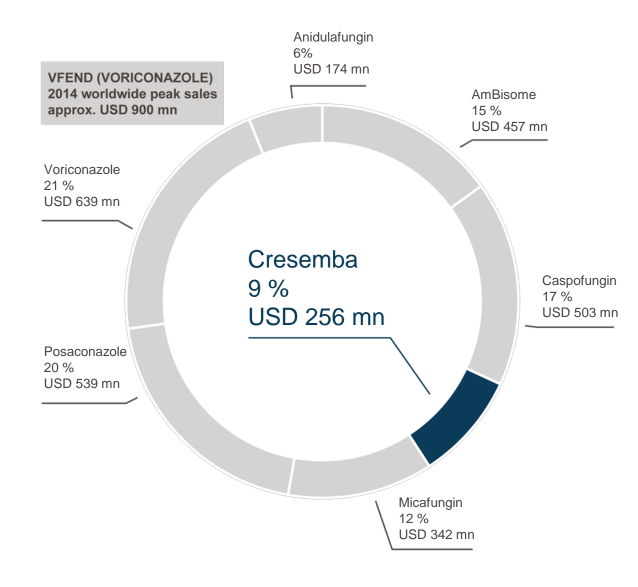


Sales of best-in-class antifungals* by product

USD 3.0 bn sales (MAT Q4 2020)

- Potential to increase Cresemba[®]
 (isavuconazole) market share
 - 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, December 2020



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 hospital-acquired pneumonia (HAP), excluding ventilator-associated pneumonia (VAP), and community-acquired pneumonia (CAP). Not approved in the U.S.

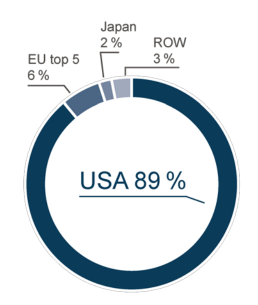
MENA: Middle East and North Africa



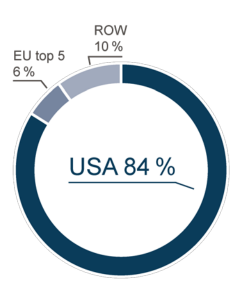


The hospital anti-MRSA antibiotic market — A USD 2.6 bn market* with the U.S. being the most important region

Daptomycin sales by region (2015, before LOE)



Ceftaroline sales by region (MAT Q4 2020)



^{*} 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, December 2020



Strategy for accessing the U.S. market

- Two cross-supportive phase 3 studies under FDA Special Protocol Assessment (SPA)
 - 1. 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 H1 2022



Phase 3 program largely funded by BARDA (up to USD ~130 mn, ~70% of total program costs)



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

² Hamed K et al. Future Microbiol. 2020;15:35-48. (NCT03138733)



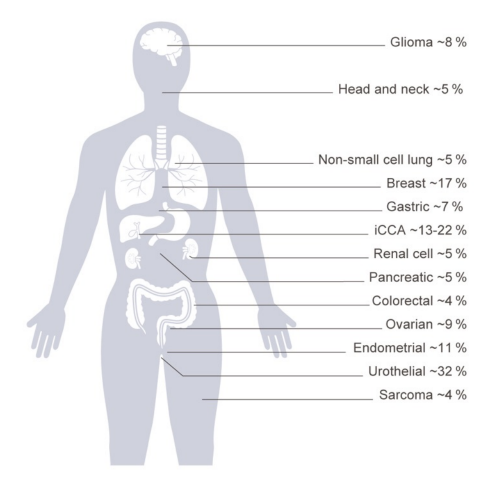
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¹Overcash JS et al. ECCMID 2020, abstract 1594. (NCT03137173)



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
- Three clinical studies ongoing
 - FIDES-01 (Ph 2) in intrahepatic cholangiocarcinoma (iCCA)
 - FIDES-02 (Ph 1/2) in urothelial cancer
 - FIDES-03 (Ph 1/2) in gastric cancer



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

Phase 2 study in iCCA* – established clinical proof of concept in FIDES-01

FIDES-01 Cohort 1

(N=103)

FGFR2 fusions

(~15% of iCCA)

Topline results:

ORR: 21.4% DCR: 74.8%

Median PFS: 7.8 months

- Consistent with earlier Phase1/2 data¹ and with interim analysis from FIDES-01
- Clinical proof of concept supporting anticancer efficacy and a favorable benefit to risk profile

FIDES-01 Cohort 2

(N=43) - ongoing

FGFR2 mutations/amplifications

(~5% of iCCA)

Interim results (n=14):

DCR: 79% (1 confirmed CR, 1 unconfirmed PR, 9 SD)

Pooled data from 23 patients

(clinical studies/EAP)²
Median PFS 7.2 months

- Encouraging PFS in pooled analysis consistent with outcome in patients with FGFR2 gene fusions
- Interim analysis successfully completed based on at least 8 patients with PFS >3 months (PFS not yet mature)
- Topline results expected H1 2022

Manageable safety profile with low incidence of nail toxicity, retinal events, hand-foot syndrome and stomatitis

FIDES-01: NCT03230318

¹Mazzaferro et al. Br J Cancer. 2019

²Droz Dit Busset et al. Annals of Oncology (2020) 31 (suppl_5): abstract 45P (NCT01752920, NCT03230318)

*in patients who progressed after at least one prior systemic chemotherapy regimen



Clinical program in urothelial cancer – FIDES-02

Multi-cohort phase 1b/2 study of derazantinib monotherapy or in combination with atezolizumab in patients with advanced urothelial cancer harboring FGFR genetic aberrations

- Substudies (N≈200) in various treatment settings, including:
 - Post-chemotherapy/immunotherapy recurrence (second-line and post second-line)
 - First-line platinum-ineligible
 - Resistance to prior FGFR-inhibitor treatment
- Clinical supply agreement with Roche for atezolizumab
- Interim results in monotherapy and combination therapy with atezolizumab in patients refractory to prior FGFR-inhibitor treatment expected H2 2021*

- Exploring an intensified dose regimen of derazantinib in two cohorts of the study:
 - Focus on maximizing efficacy by using an intensified dose regimen of 400 mg per day
 - as monotherapy in a second-or post second-line setting in FGFR-inhibitor naïve patients
 - as monotherapy or in combination with atezolizumab in first-line cisplatin-ineligible patients
 - Supported by the observed safety and tolerability profile of derazantinib and by pharmacology data
- Initial results from cohorts utilizing 400 mg per day dose regimen expected H1 2022



Clinical program in gastric cancer – FIDES-03

Multi-cohort Phase 1b/2 study of derazantinib as monotherapy or in combination therapy with standard of care (ramucirumab/paclitaxel) 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 with ramucirumab/paclitaxel
 - Combination of derazantinib with atezolizumab
- Clinical supply agreement with Roche for atezolizumab
- Clinical trial collaboration and supply agreement with Lilly for ramucirumab

- Exploring an intensified dose regimen of derazantinib 400 mg per day in monotherapy and in combination therapy
- Interim results in derazantinib monotherapy and recommended phase 2 dose of derazantinib in combination with ramucirumab/paclitaxel expected H1 2022

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	Phosphorus얍 Nausea Vomiting	Phosphorus û Fatigue Stomatitis	Phosphorus*얍 Diarrhea* Dry mouth*	Phosphorus û Alopecia Diarrhoea	Diarrhoea Alopecia Constipation	Phosphorus û Stomatitis Fatigue	
Blood phosphorus ⊕†	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,

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



⁶ Balversa[™] 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 Pemazyre™ U.S. Prescribing Information (April 2020) and Balversa™ 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 Pemazyre[™] across clinical trials;

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



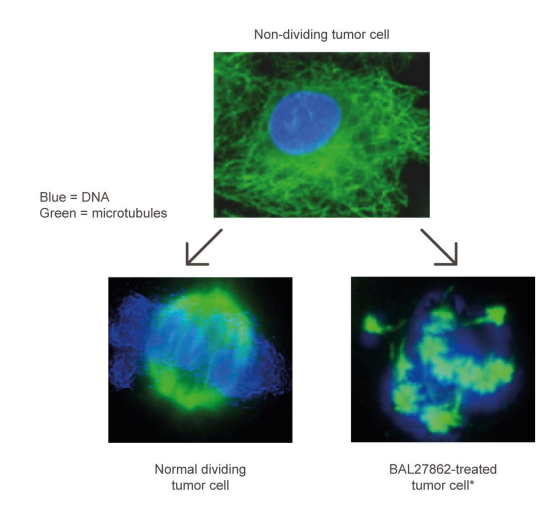
Lisavanbulin (BAL101553)

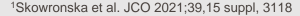
Glioblastoma and other solid tumors



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
- Crosses the blood-brain barrier with potent activity in brain tumor models alone and in combination
- Comprehensive biomarker program to optimize patient selection, e.g. EB1 (end-binding protein 1)
 - Prevalence of EB1-positivity in glioblastoma approximately 5%
 - EB1-positivity also reported in other tumor types (most prominent in medulloblastoma, neuroblastoma and metastatic melanoma)¹







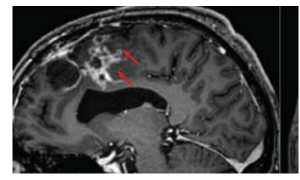
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^{*} Lisavanbulin (BAL101553) is a prodrug of BAL27862

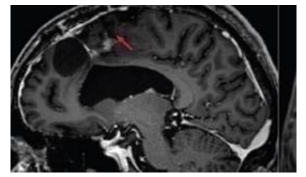
Biomarker-driven phase 2 study ongoing in recurrent glioblastoma

- EB1 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
- Results from phase 1 study with daily oral lisavanbulin in patients with recurrent glioblastoma (n= 20):^{1, 2}
 - Three patients with EB1-positive glioblastoma
 - Two of the EB1-positive patients with long-lasting clinical benefit, ongoing for more than 2 years
 - One exceptional response with >80% reduction in glioblastoma tumor size
 - No clear clinical benefit for EB1-negative patients
- Phase 2 interim results expected H2 2021

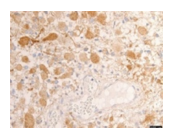
Glioblastoma tumor size reduction in an exceptional responder and EB1 staining of glioblastoma tissue compared to non-responding patients



Baseline (May 2018)



Post Cycle 12 (April 2019)



Responder



Non-responder

² Tiu et al. JCO 2021;39,15 suppl, TPS2068 (NCT02490800)



¹ Lopez et al. JCO 2019;37,15 suppl, 2025 (NCT02490800)



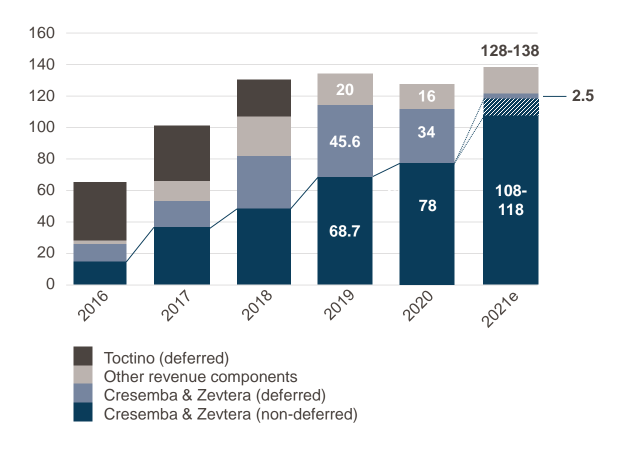
Financials & Outlook



Financial guidance

In CHF mn	FY 2020	FY 2021 guidance
Total revenue	127.6	128 - 138
thereof: Contributions Cresemba® & Zevtera® non-deferred deferred	78.2 33.8	108 - 118 2.5
Operating loss	8.2	13 - 23
Cash and investments*	167.3	155 – 160**

Continued strong double-digit growth in Cresemba & Zevtera non-deferred revenue contributions Y-o-Y, CHF mn



^{*}Cash, cash equivalents, restricted cash and investments / **Excluding any potential impact from a reduction of the outstanding convertible bonds



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Key Milestones & Outlook 2021 / 2022

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

		H1 2021	H2 2021	H1 2022	H2 2022
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	Topline results from SAB phase 3 study	
	FIDES-01	√ Topline results (FGFR2 gene fusions)			
	(iCCA)	✓ Interim results (other FGFR2 gene aberrations)		Topline results (other FGFR2 gene aberrations)	
Derazantinib	FIDES-02 (urothelial cancer)		Interim results in monotherapy and combination therapy with atezolizumab in patients refractory to prior FGFR inhibitors	Interim results in monotherapy (400 mg/day) in 2nd-line FGFR-inhibitor naïve patients and atezolizumab combination in 1st-line cisplatinineligible patients	
	FIDES-03 (gastric cancer)			Interim results in monotherapy (400 mg/day) and recommended phase 2 dose with ramucirumab/paclitaxel	Interim efficacy results in combination with ramucirumab/paclitaxel
Lisavanbulin			Interim results from phase 2 biomarker- driven glioblastoma study	Topline results from phase 2 biomarker-driven glioblastoma study	
			Recommended phase 2 dose in phase 1 study in newly-diagnosed glioblastoma in combination with radiotherapy		

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Focused on Growth and Innovation

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