

Pharming Group N.V.

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JP Morgan Conference

San Francisco

11-16 January 2020

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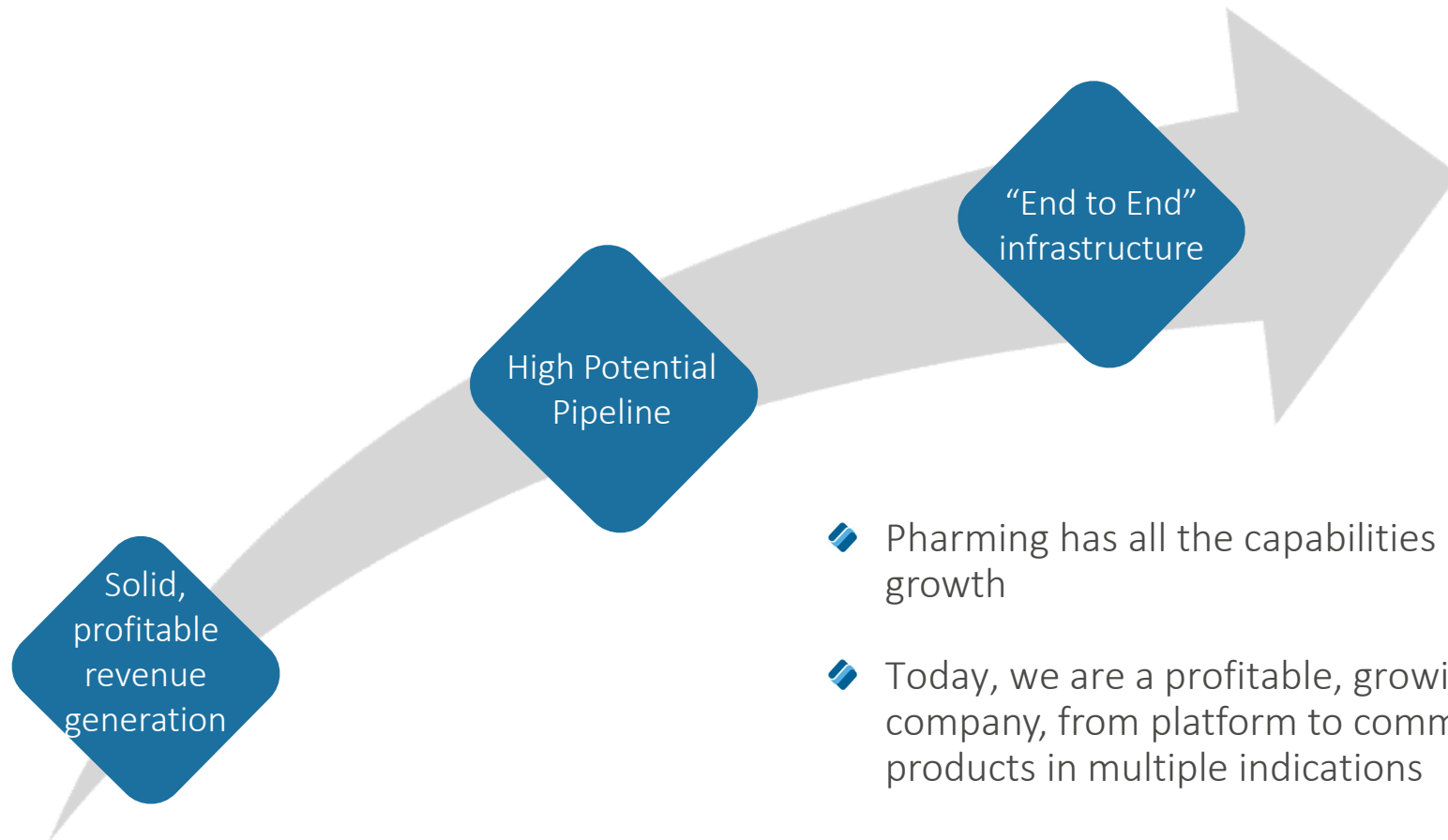
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- ◆ **Public Company:** Euronext: PHARM: ~€965 million (~\$1.07 billion)
- ◆ **Located:** the Netherlands, ~220 employees globally
- ◆ **Current Focus:** Rare and Ultra-rare disease development and commercialization
 - Marketed product: **RUCONEST®**
 - Recombinant human C1-esterase inhibitor (enzyme replacement therapy)
 - For acute angioedema attacks in patients with hereditary angioedema (HAE)
 - Marketed in USA, EU, LatAm, Korea and Israel with other territories coming
- ◆ **Profitable and cash flow positive with 9M2019 net sales of €123M, and expecting continued growth in sales**





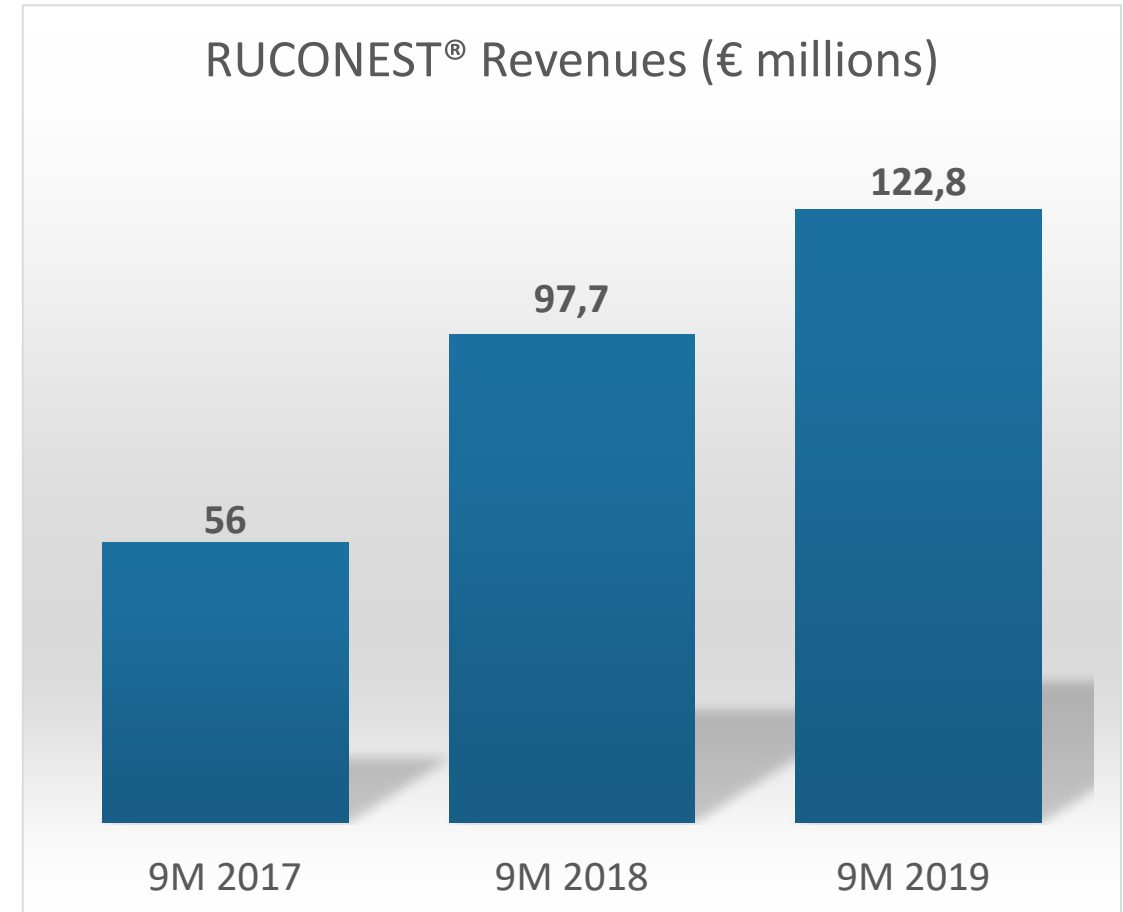
- ◆ Pharming has all the capabilities needed for sustainable, high growth
- ◆ Today, we are a profitable, growing and fully integrated biopharma company, from platform to commercialization, with multiple products in multiple indications
- ◆ From this we can drive lasting additional growth by extending the uses of our existing drug into larger indications and developing innovative solutions in select rare, ultra-rare and specialty diseases

Solid,
profitable
revenue
generation



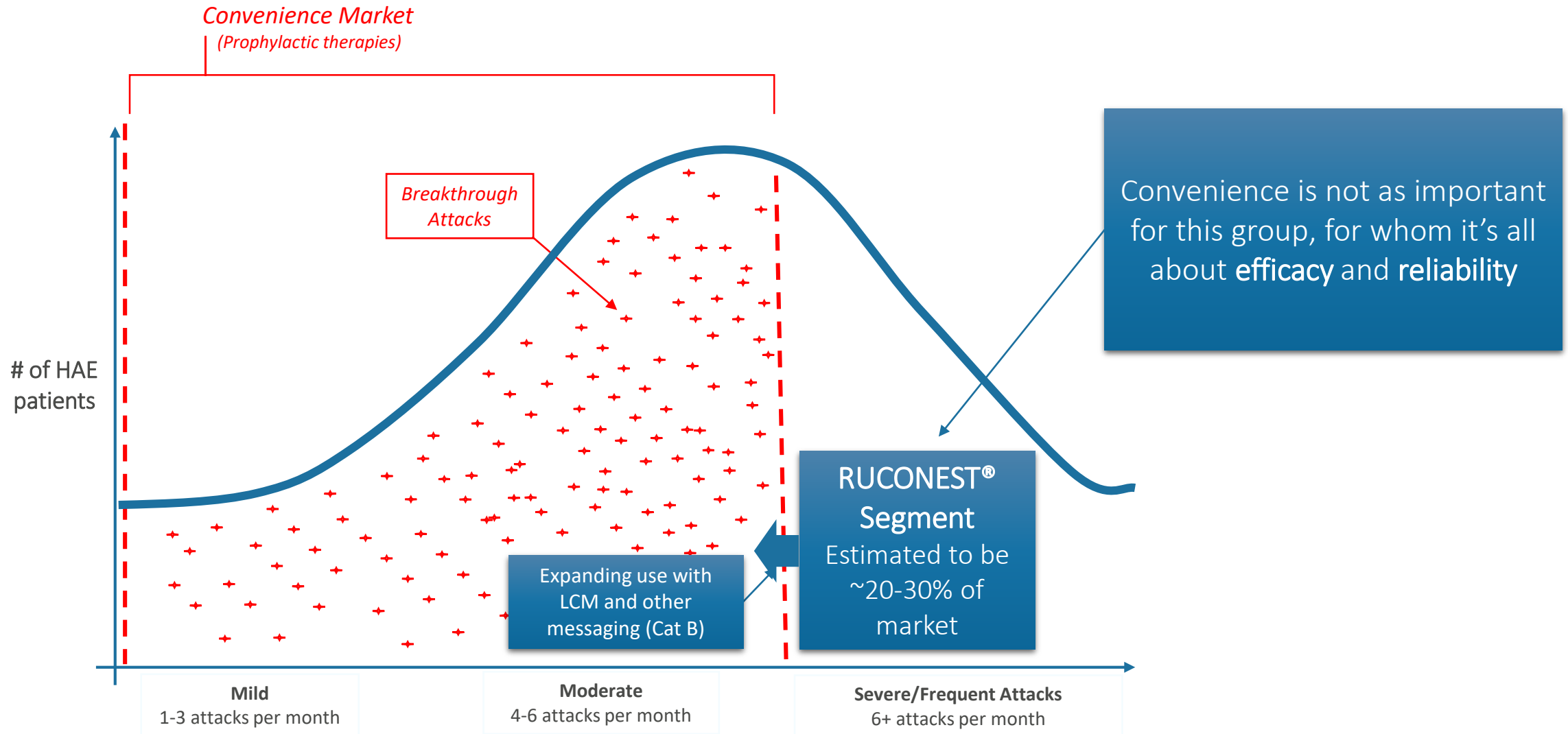
RUCONEST® : Strong Execution of Commercial Strategy

- ❖ HAE is a complex, serious disease with many idiosyncrasies and a varied market.
- ❖ The current approved therapies all address certain specific segments/phenotypes of HAE.
- ❖ RUCONEST® is the only recombinant PRT, and serves a segment the other therapies are unable to serve in an adequate way, due to its dosing and method of administration
- ❖ As a result of the solid RUCONEST® business, Pharming has a strong balance sheet with growing cash position
- ❖ New re-acquisition of territories licensed to Sobi in December 2019 allows expansion of EU and RoW sales



Pharming is in a very strong position to continue to grow

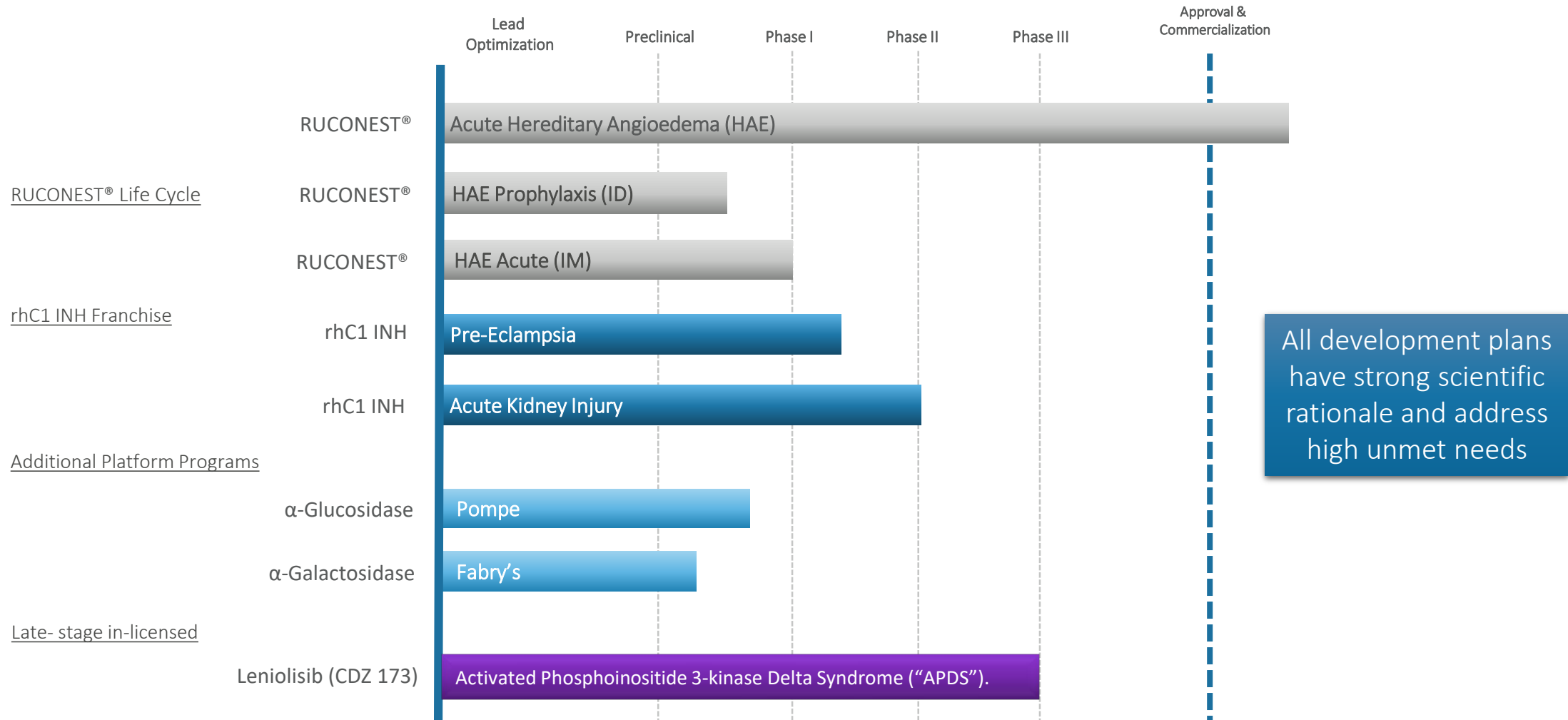
RUCONEST®: Patient Segmentation in HAE



High
Potential
Pipeline



High Potential Pipeline





Acute Kidney Injury (AKI)

- ❖ First described in the 1950s
- ❖ Radiographic contrast medium are responsible for 11% of cases of hospital-acquired renal insufficiency, the third most common cause of renal failure after impaired renal perfusion and the use of nephrotoxic medications.
- ❖ 40 million contrast-enhanced scans per year in the US alone, with around 20% on high-risk subgroup patients
- ❖ AKI from contrast media is responsible for a third of all hospital-acquired acute kidney injury (AKI)
- ❖ AKI affects between 1% and 2% of the general population, and up to 50% of high-risk subgroups following coronary angiography (CA) or percutaneous coronary intervention (PCI).¹
- ❖ No specific therapy available at present: outcomes vary from reversible AKI which requires 7-21 day ICU treatment, through permanent dialysis, renal transplantation and death.

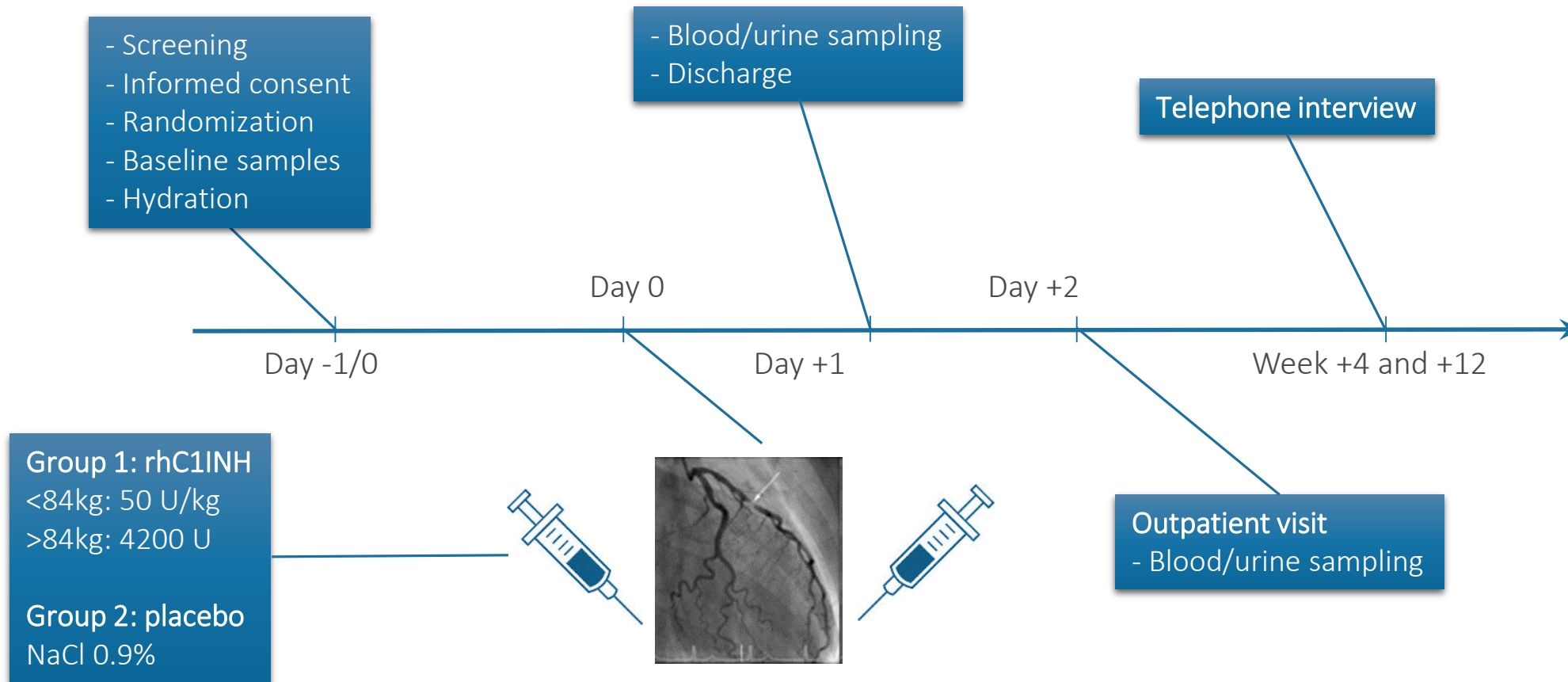
Table 1 | Risk factors for the development of CIN

Fixed (non-modifiable) risk factors	Modifiable risk factors
Older age	Volume of CM
Diabetes mellitus	Hypotension
Pre-existing renal failure	Anemia and blood loss
Advanced CHF	Dehydration
Low LVEF	Low serum albumin level (< 35 g/l)
Acute myocardial infarction	ACE inhibitors
Cardiogenic shock	Diuretics
Renal transplant	Non-steroidal anti-inflammatory drugs
	Nephrotoxic antibiotics
	IABP

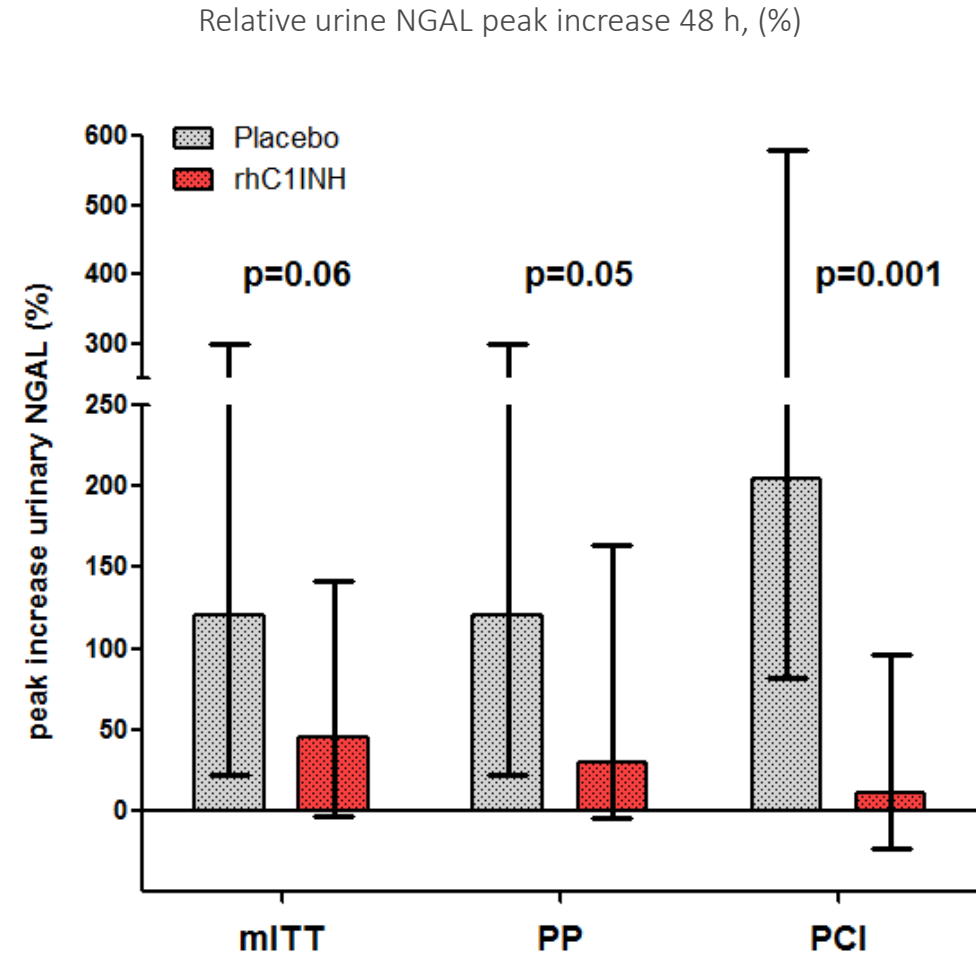
Abbreviations: ACE, angiotensin-converting enzyme; CHF, congestive heart failure; CIN, contrast-induced nephropathy; CM, contrast media; IABP, intra-aortic balloon pump; LVEF, left ventricular ejection fraction.

¹ Mehran R, Nikolsky E. Contrast-induced nephropathy: definition, epidemiology, and patients at risk. *Kidney Int Suppl* 2006:S11–15
CIN= Contrast induced nephropathy

Study: AKI Resulting from Contrast Medium (CM)



- ◆ Study of contrast-induced nephropathy at University Hospital Basel reported Oct 2018
- ◆ 75 eligible patients given either drug or placebo before and after treatment
- ◆ Study used existing HAE dose, and showed clinical and statistical significance
- ◆ Results were especially strong in percutaneous coronary intervention patients
- ◆ Data has been used to prepare for a study in this specific group led by the same team in Switzerland, starting very shortly





Pre-Eclampsia (PE)

- ❖ Pre-eclampsia (PE) has a prevalence of 1-17% throughout the world. Estimated yearly cases of PE in the US alone: 120,000+.
(Steeegers et al., 2010; Osungbade and Ige, 2011)
- ❖ Delivery is presently the only therapy of PE, but this is not an option for early PE (from week 20 of gestation).
- ❖ The main goal of symptomatic therapy is to prolong gestation of PE patients as far as possible

Panel 1: Maternal and fetal complications in severe pre-eclampsia

Maternal complications

- Abruptio placentae (1–4%)
- Disseminated coagulopathy/HELLP syndrome (10–20%)
- Pulmonary oedema/aspiration (2–5%)
- Acute renal failure (1–5%)
- Eclampsia (<1%)
- Liver failure or haemorrhage (<1%)
- Stroke (rare)
- Death (rare)
- Long-term cardiovascular morbidity

Neonatal complications

- Preterm delivery (15–67%)
- Fetal growth restriction (10–25%)
- Hypoxia-neurologic injury (<1%)
- Perinatal death (1–2%)
- Long-term cardiovascular morbidity associated with low birthweight (fetal origin of adult disease)

Table 1 Analytical data (mean \pm 1 SD) in normal pregnancy, preeclampsia and in non-pregnant women

	(A) Normal pregnancy (n = 20)	(B) Mild preeclampsia (n = 17)	(C) Moderate preeclampsia (n = 10)	(D) Non-pregnant women (n = 20)
C1-INH activity (%)	74.3 \pm 15.5	64.4 \pm 14.0	55.5 \pm 15.8	95.1 \pm 10.8
C1-INH antigen (%)	68.2 \pm 10.4	62.7 \pm 13.3	53.1 \pm 8.8	86.5 \pm 12.2

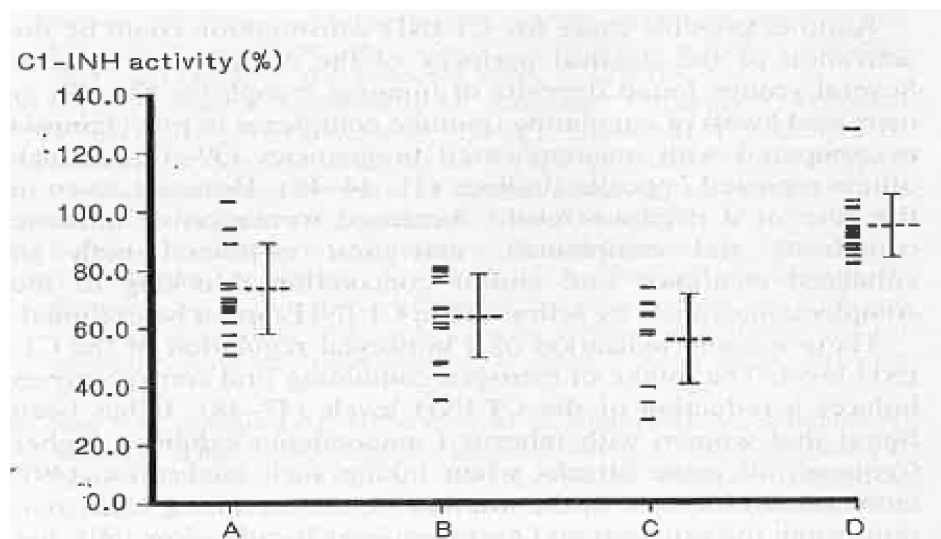


Fig. 2 Scattergram of C1-INH activities. Uncomplicated pregnancies (A), mild preeclampsia (B), moderate preeclampsia (C) and non-pregnant controls (D)

- ◆ High unmet need with no current treatment
- ◆ Significant cost to healthcare system and families
- ◆ Challenging disease to study; demands thoughtful, ethical approach
- ◆ Initial clinical study initiated (Netherlands and Australia)



Activated PI3K- δ Syndrome (APDS)

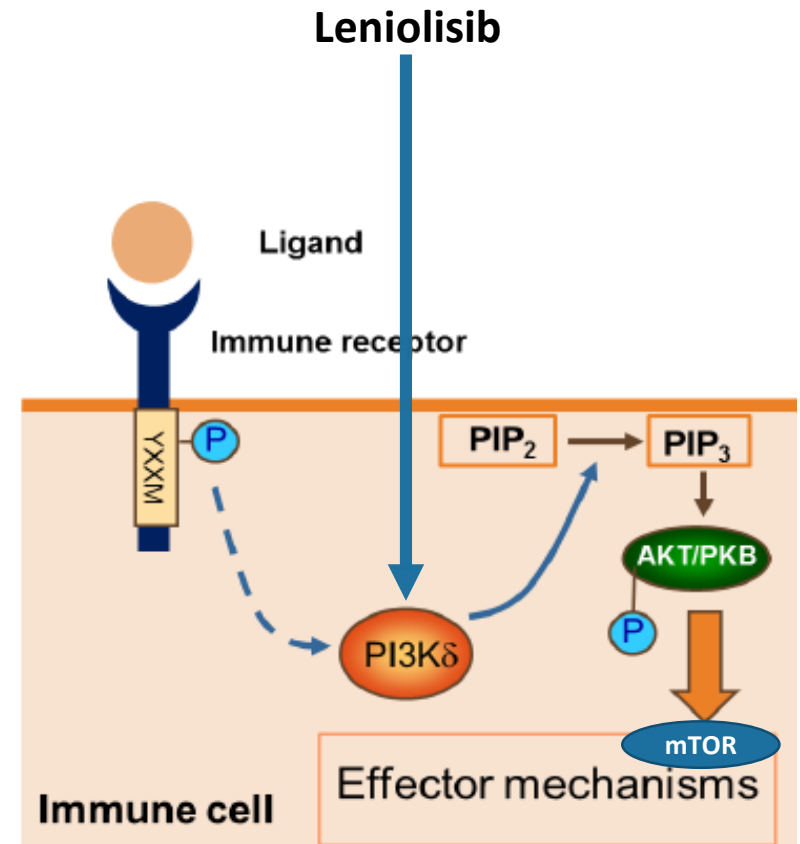
- ◆ Primary immunodeficiencies (PID) lead to immune system dysregulation with numerous resulting complications
 - Prevalence 1 in 1200
 - More than 300 gene mutations known to cause different PIDs
 - Highly variable clinical presentation, but increased susceptibility to infection is common to most PIDs

- ◆ Activated PI3 kinase delta syndrome (APDS) is a primary immunodeficiency
 - Caused by autosomal dominant mutations
 - Increased activity of phosphoinositide-3-kinase δ (PI3K δ)
 - Estimated prevalence 1-2/million
 - More than 240 reported in literature
 - Screening in subset of PID patients has found rates: 5/669 (1%) and 17/184 (9%)
 - Commercially available genetic test

Angulo I, et al. Science. 15;342. 2013. Lucas CL. Nature Immunology. 15, 88-97, 2014.

Michalovich D, et al. Frontiers Immuno. 2018. Jamee M, et al. Clin Rev Allergy Immunology. 2019.

- ◆ Current treatment options for APDS:
 - Symptomatic treatment e.g., antibiotics
 - Immune globulin replacement therapy (IVIG/SCIG)
 - Stem cell transplantation
 - Case reports of mTOR inhibitor rapamycin
- ◆ Leniolisib
 - Potent, selective PI3K δ inhibitor
 - Treats the root cause of APDS
 - Orally bioavailable – tablet/capsule
 - Direct PK/PD relationship observed
 - Currently in registration-enabling pivotal study
 - If approved, the drug is expected to reach the market in mid-2022



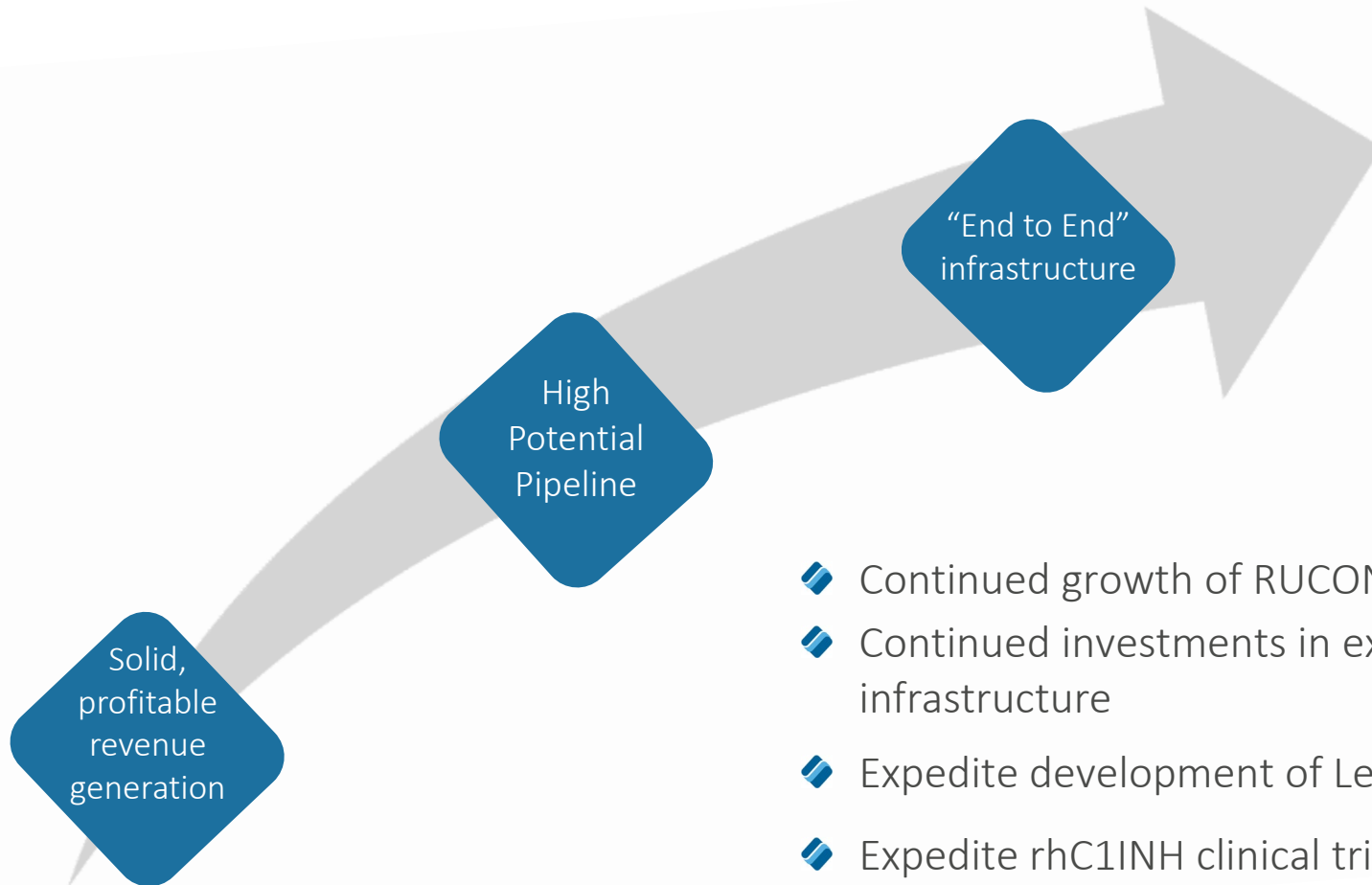
Financial Performance & 2019 Outlook



Third Quarter 2019: Financial Results

9 months to 30 September

<i>Amounts in €m except per share data</i>	<i>2019 3rd Quarter</i>	<i>2019 1st 9 months</i>	<i>2018 1st 9 months</i>	<i>% Change</i>
<i>Income Statement</i>				
Revenue from product sales	45.3	122.8	97.7	26%
Other revenue	0.2	0.6	0.6	
Total revenue	45.5	123.4	98.3	26%
Gross profit	40.1	107.1	82.4	30%
Operating result	18.1	42.7	31.0	38%
Net result	10.5	24.1	13.9	73%
<i>Balance Sheet</i>				
Cash & marketable securities	64.4	64.4	72.2	(11%)
<i>Share Information</i>				
Earnings per share (€): - Undiluted	0.017	0.038	0.022	73%
- Fully diluted	0.015	0.036	0.021	71%



- ◆ Continued growth of RUCONEST[®] sales, profitability and +ve cashflow
- ◆ Continued investments in expansion of commercial & manufacturing infrastructure
- ◆ Expedite development of Leniolisib to FDA and EMA approval
- ◆ Expedite rhC1INH clinical trials for PE and AKI
- ◆ Continued development PRT for Pompe and Fabry
- ◆ Re-evaluate most advantageous administration route for RUCONEST[®]

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ENXTAM: PHARM

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