OSE IMMUNO THERAPEUTICS

Breaking Through the Therapeutic Ceiling with First-In-Class Immunotherapies

March 2024

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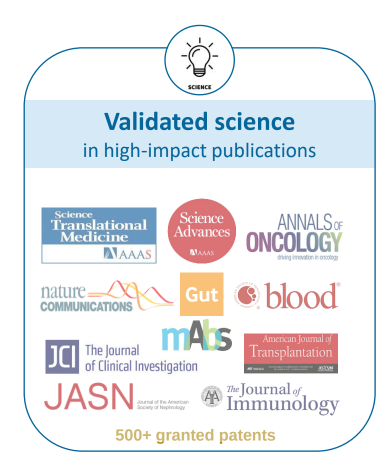
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OSE's strong foundation & recurrent track record of success

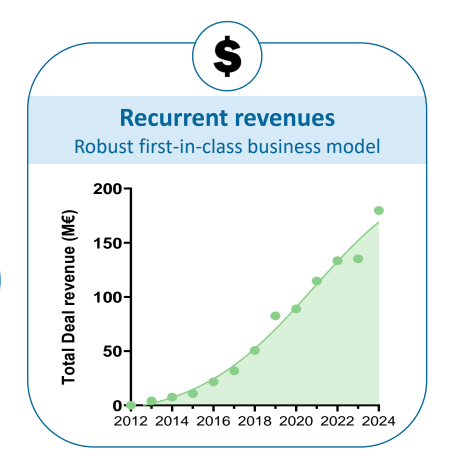
10 years of validated innovation in immunology thanks to an Extra[not]Ordinary R&D engine











Delivering First-in-Class immunotherapies from Target to Clinic

Key strategic pharma partnerships driving long-term value

- Founded in 2012
- IPO/Euronext in 2015
- 60+ FTEs
- **500+** granted patents
- **52 M€** : Equity
- 180 M€: Partnerships*

+75% non-dilutive funding





Phase 3 asset in Oncology

Tedopi[®] most advanced cancer vaccine NSCLC 2L post-CPI market: +5b\$/year



Phase 2 asset in Inflammation

Lusvertikimimab anti-IL7R mAb Ulcerative colitis market: +10b\$/year

Strategic Pharma Partners

+2.1b€ potential milestones ObVie







Clinical stage assets

- 3 **Fully** owned (Phase 1, 2, 3)
- 2 Partnered (Phase 1, 2)

Pre-clinical platforms Assets approaching development

- Innovative MoA & Targets to address critical unmet need
- International Research Collaboration













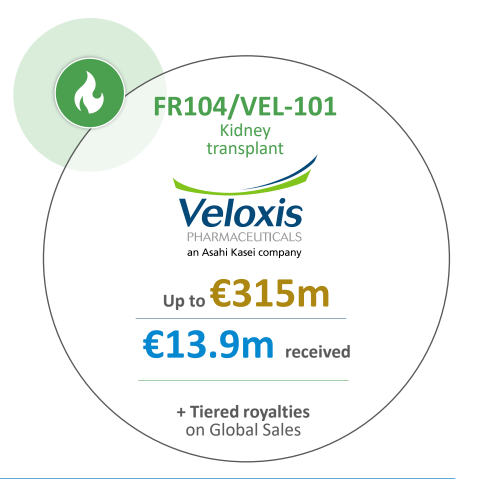
Strategic partners provide industry-leading clinical support & strong financial foundations

Over €2.1bn in potential milestones; €180m* already received



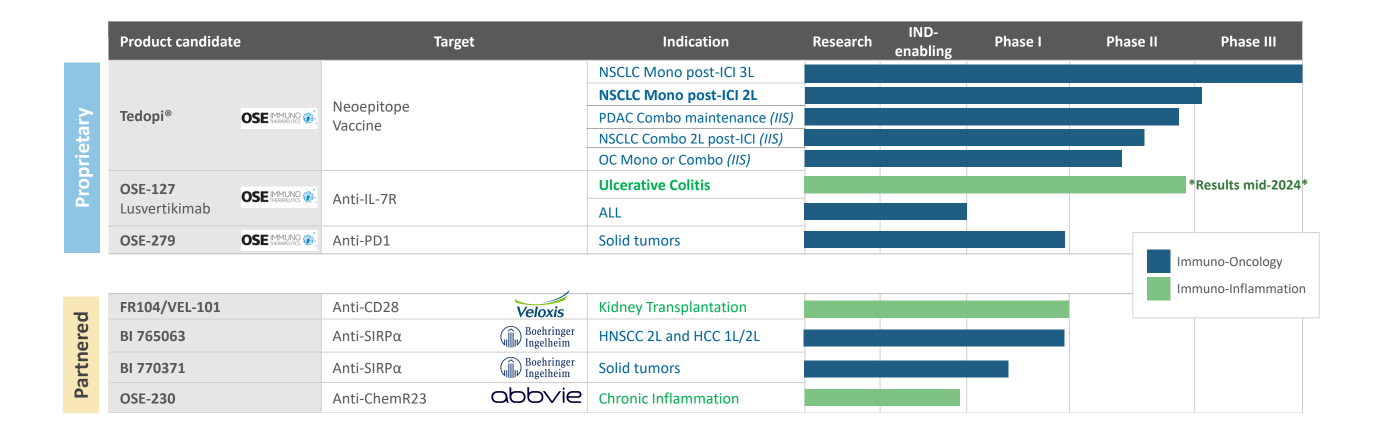






OSE's Clinical pipeline

Combining a clinical portfolio of first-in-class immunotherapies and diversified assets in IO and I&I



OSE's Research platforms

Extra[not]Ordinary Research PowerHouse



Pro-resolutive mAb

Partnered Asset:
Anti-ChemR23*

Ongoing programs
Undisclosed new
pro-resolutive GPCRs

Cis-Targeted
Augmented Cytokine

Main Asset : Anti-PD1-IL7v

Ongoing programs

Cis-Demasking

new technologies

Myeloid Checkpoint

Partnered Asset:
Anti-SIRPa#

Ongoing programs

Anti-CLEC-1 mAbs

preclinical program





Key catalysts



ℬ Readouts

- Lusvertikimab
 Phase 2 results in UC
- OSE-279
 Phase 1 results
- BI 765063/BI 770371 (partnered)
 Phase 1/2 results in solid tumors
- FR104/VEL-101 (partnered)
 Phase 1/2 results in Kidney
 Transplantation



Progress

- Tedopi®
 Phase 3 start in NSCLC 2L
- FR104/VEL-101 (partnered)
 Phase 2 start in Kidney
 Transplantation
- OSE-230 (partnered) IND/Phase 1
- R&D programs & Lusvertikimab
 New partnering opportunities



Readouts

- Tedopi® Phase 3 <u>results</u> in NSCLC 2L
- BI 765063/BI 770371 (partnered)
 Phase 2 results
- FR104/VEL-101 (partnered)
 Phase 2 results in Kidney
 Transplantation
- OSE-230 (partnered)
 Phase 1 results + Phase 2 results



Progress

- Lusvertikimab (to partner)
 Phase 3 start
- BiCKI®-IL7v IND/Phase 1
- CLEC-1 IND/Phase 1
- New R&D programs/platforms

2024

2025-2027



Investment highlights

Late-stage compelling product

Promising clinical data from the lead asset Tedopi®

- Met primary overall survival endpoint in monotherapy in Pol pivotal NSCLC post-ICI study
- Significant better Safety profile & Quality of Life with positive Net Treatment Benefit versus SOC

Large market opportunities

Focus on multi-billion \$ markets

- I/O: NSCLC (2L, 3L), HCC (1L, 2L), HNSCC (2L), Leukemia
- I&I: IBD (Ulcerative Colitis), Kidney Transplantation

Strong pharma partnerships

Sustainable business through multi-partnership strategy

>€2.1bn milestones: AbbVie, Boehringer Ingelheim, Veloxis

Long duration IP portfolio

IP extends to 2040's

I/O: Tedopi® (>2038), OSE-172 (>2037), OSE-279 (>2039), CLEC-1 (>2040) I&I: OSE-127 (>2037), FR104 (>2035), OSE-230 (>2040)

Multiple upcoming catalysts

Multiple key clinical and regulatory milestones expected in next 12 months

- Tedopi®: Confirmatory pivotal phase 3 NSCLC 2L start
- Lusvertikimab (OSE-127): Top-line results Ulcerative Colitis Phase 2
- BI 765063/BI 770371: Phase 1b results in solid tumors
- FR104/VEL-101: Phase 2 start in Kidney Transplantation
- **OSE-230:** Phase 1

Financial position

Cash visibility until 2026

€15m available cash as of December 30, 2023, + \$48m payments on recent pharma partnership

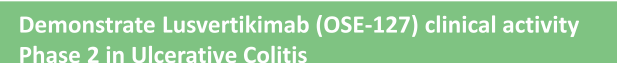


Our plan to build a leading immunotherapy company

Position Tedopi® as the best treatment option after ICI-failure in cancer patients



Leverage the clinical advantage of anti-SIRP α in Solid Tumors



Confirm FR104/VEL-101 benefit as maintenance therapy in kidney transplantation



Explore the pro-resolutive mAb potential in chronic & severe inflammation



Advanced proprietary early-stage assets from OSE's research platforms + New Partnering Opportunities





First-in-class strategy



Proprietary clinical programs

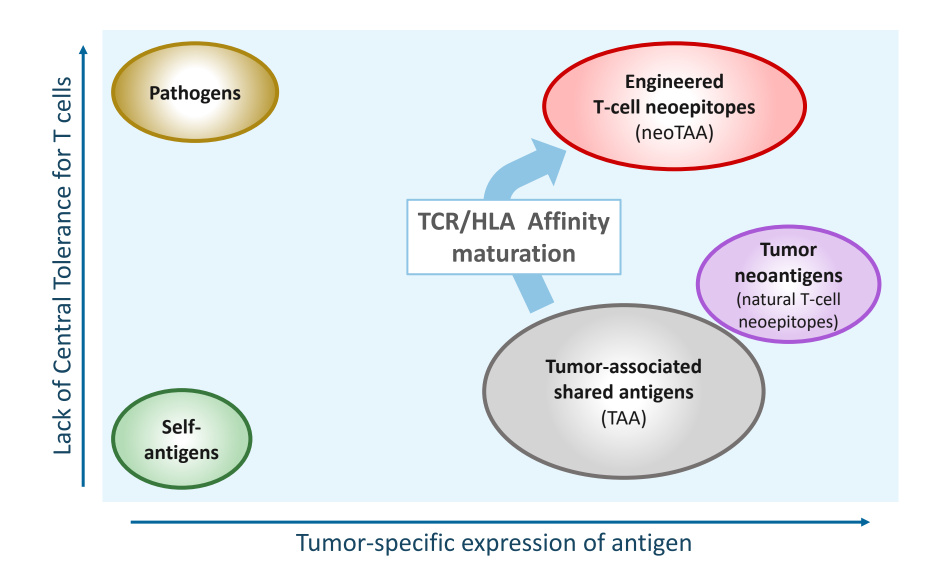
TEDOPI®

Most Advanced Therapeutic Cancer Vaccine

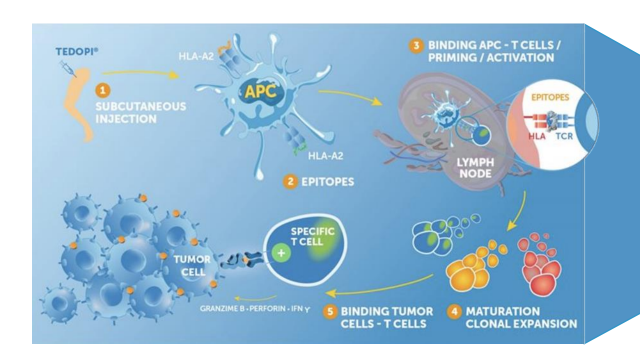
Bringing new hope to patients in the fight against ICI resistant NSCLC



Cancer Antigens Immunogenicity



An immunotherapy activating specific T-cells to revive anti-tumor response



Most advanced Cancer Vaccine in clinical development

- Unique combination of neoepitopes: small peptides deriving from tumor specific antigens* expressed in various cancers
- Strong binding to HLA-A2 receptor (45% population)
- Direct activation of tumor specific T-cells differs from checkpoint inhibitors releasing the break of immune response

Proprietary combination (9 **optimized neoepitopes** + 1 epitope giving universal

T helper response)

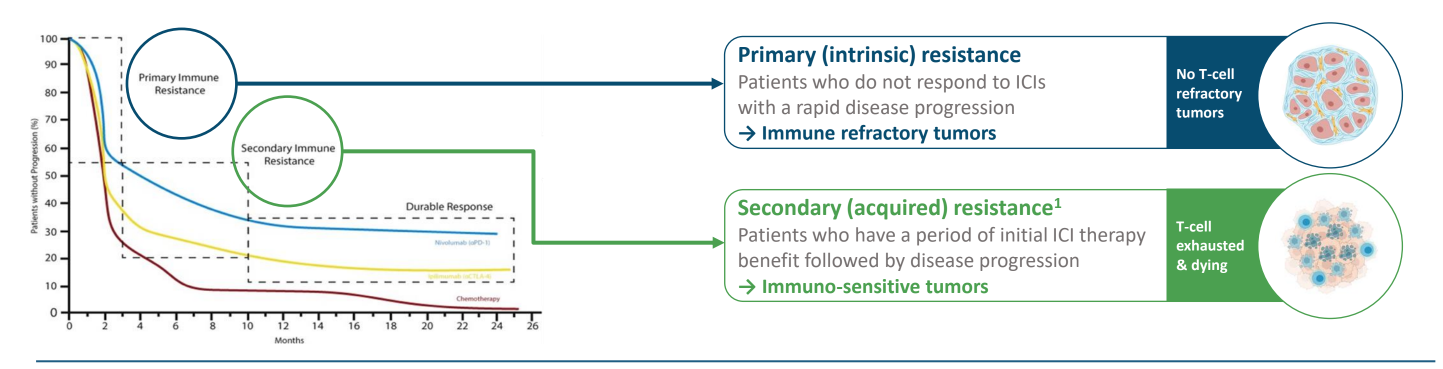
Induces early T cell
memory responses
+
Migration in tissues

Ready to Use subcutaneous formulation with Q3W injection Orphan Drug
Designation (FDA)
>1,000 injection
in clinical trials

Strong IP position until **2038**¹ (US / EU / Asia)

Tedopi® is a novel cancer vaccine with a strong biological rational in post-ICI secondary resistance

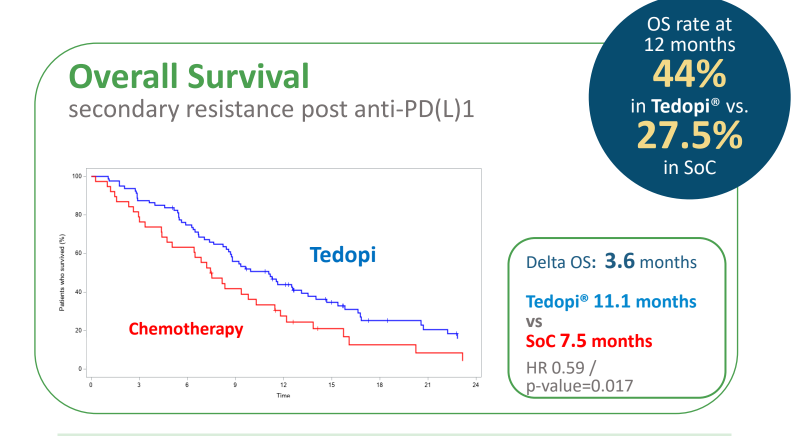
Shifting paradigms with cancer vaccine immunotherapy



Tedopi[®] has the **potential to rejuvenate & refresh specific TILs** in immuno-sensitive tumors. Neoepitope-specific T cells have tumor killing potential and limited side effects.

Clinically meaningful benefit of Tedopi®

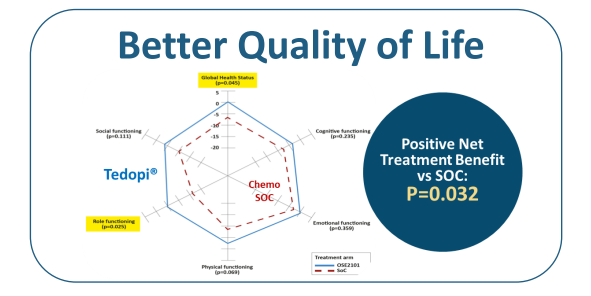
First randomized Phase 3 with positive results vs. standard of care (SOC)



Risk of Death reduced by 41% versus chemo.

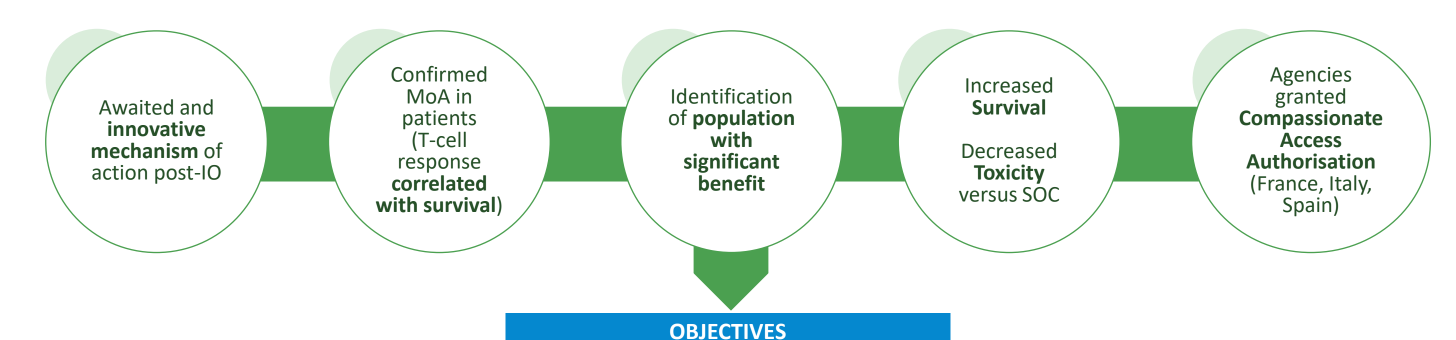
Significantly safer than Chemo.

11% vs **35%** grade 3-5 AEs





Position Tedopi® as the best treatment option after ICI-failure in cancer patients





Compassionate use in 3L NSCLC



FDA/EMA optimal regulatory paths for the new confirmatory pivotal phase 3 trial and CDx for potential approval in 2L NSCLC after ICI-failure



Additional Phase 2 clinical trials in combination (NSCLC, Pancreatic, Ovarian)

Target population estimated at 100k patients/year in NSCLC post-ICI (2nd line)

Expending the PD-(L)1 NSCLC market is growing (US\$bn)1 potential in 2L post-ICI in G7 years 14.7 13.6 **VUBTAYO** 12.3 58% 2.3 **IMFINZI** 1.8 10.0 1.1 TECENTRIQ® 7.5 mNSCLC 2L OPDIVO. 1.8 2.2 1.7 - drug treated 5.3 0.3 4.2 3.7 **KEYTRUDA** 7.6 7.5 7.3 (pembrolizumab) Injection 100 mg 45% 3.2 1.2 3.0 1.2 2016 2017 2018 2019 2020 2021

- Lung cancer is the leading cause of cancer mortality worldwide, accounting for about 1.8m deaths each year.²
- NSCLC is the most common type of lung cancer, accounting for 85% of all lung cancers.³
- ~60% of 1L patients progress within 18 months.
- HLA-A2 phenotype in about 45% of the population.
- Target NSCLC population: ~10%

HLA-A2+

Tedopi® delivers important clinical benefits vs competition

Better Safety profile and QoL in current landscape of late-stage drug development post CT-IO

Company	OSE IMMUNO (MIRATI THERAPEUTICS	Roche SIPSEN	MERCK Eisai	gsk	BIONTECH OncoC4	AstraZeneca Duido-Sankyo	GILEAD	SANOFI	abbvie
Tougot	Naviti opitopos vessins	TKIs (anti-angiogenic)		Checkpoint Inhibitors		ADCs				
Target	Multi-epitopes vaccine			TIM-3	CTLA-4	TROP2	TROP2	CEACAM5	c-MET	
Current Study	ATALANTE-1	SAPPHIRE	CONTACT-01	LEAP-008	COSTAR Lung	PRESERVE-003	Tropion-LUNG1	EVOKE-01	CARMEN-LC03	NCT04928846
n	219 118 (secondary resistant)	500	350	405	750	600	604	580	554	698
Therapy	Tedopi® vs docetaxel	Sitra + Opdivo vs. docetaxel	Cabo+Tecentriq vs. docetaxel	Lenvi + Keytruda vs. docetaxel	Cobolimab + Jemperli vs. docetaxel	Gostistobart vs. docetaxel	datopotamab deruxtecan vs docetaxel	Sacituzumab Govitecan-hziy vs docetaxel	SAR408701 vs. docetaxel	Telisotuzumab Vedotin vs. Docetaxel
Primary endpoints	os	OS	OS	PFS and OS	OS	OS	PFS and OS	OS	PFS and OS	PFS and OS
Initiation	2017	Q3 2019	Q3 2020	Q2 2019	Dec 2020	Q2 2023	Q4 2020	Q4 2021	Q1 2020	Q1 2022
Read-out	2022	Failed	Failed	Failed	2024+	2027+	Failed OS (interim analysis)	Failed	Failed	2025+
		Safety data from early-stage trials in NSCLC post-ICI								
- TEAEs G3/4	11%	60%	39%	78%	n.a.	43%	25-30%	> 30%	36%	36%
Source	Besse et al. 2023	Leal, et al ESMO 2021	Neal et al, ASCO 2022	Taylor et al, J. Clin. Oncol. 38, 1154–1163.	Davar et al, SITC 2018	He et al, ASCO 2023	Lisberg et al, ESMO 2023	Suk Heist et al. JCO 2017	Gazzah et al, ASCO 2020	Camidge DR, et al. WCLC 2021



Further additional potential clinical value in combination NSCLC, PDAC and OC

Phase 2 ISS trials in combination with immunotherapy or chemotherapy treatments

2nd line post 1st line chemo IO

CombiTED - NSCLC In combination with nivolumab



Tedopi[®] Plus Docetaxel or Tedopi Plus Nivolumab as 2nd line Therapy in Metastatic NSCLC failing standard 1st line Chemo-immunotherapy¹

Sponsored by FoRT
PI: Federico CAPPUZZO
(Roma Cancer Institute)
Italy /Spain/ France



Readout expected 2025

Maintenance setting post standard of care

TEDOVA - Ovarian Cancer In combination with pembrolizumab



ARCAGY - GINECO

Tedopi[®] Alone or in Combination With Pembrolizumab vs Best Supportive Care as Maintenance in Patients with Platinum-Sensitive Recurrent Ovarian Cancer²

Sponsored by ARCAGY-GINECO
PI: Alexandra LEARY
(Gustave Roussy Institute)
France/ Germany/ Belgium

Readout expected in 2025

TEDOPaM - Pancreatic CancerIn combination with FOLFIRI



Tedopi® plus FOLFIRI vs FOLFIRI as Maintenance Treatment in Controlled Advanced or Metastatic Pancreatic Ductal Adenocarcinoma after 8 Cycles of Folfirinox³

Sponsored by GERCOR PRODIGE
PI: Cindy NEUZILLET
(Curie Institute)
France



Recruitment completed Q2 2023

Readout expected in 2024

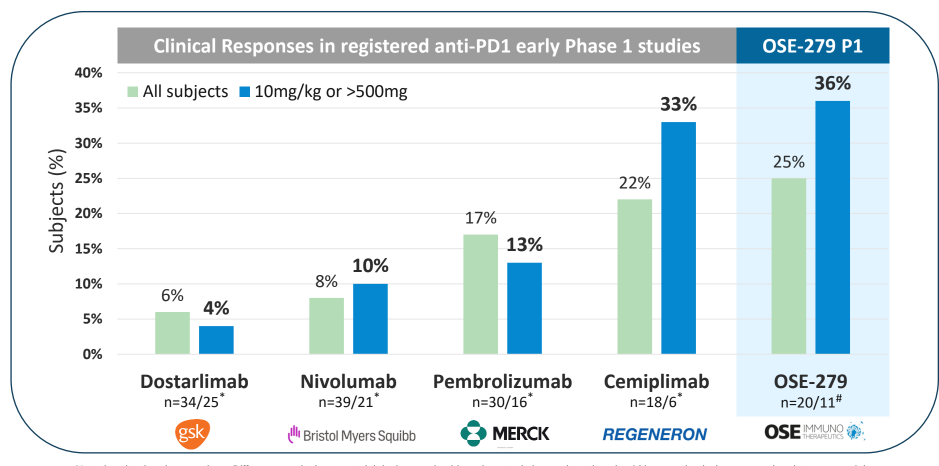


- 1 NCT04884282 105 Patients planned
- 2 NCT04713514 180 Patients
- 3 NCT03806309 136 patients -recruitment completed

OSE-279: Proprietary anti-PD1 mAb

High affinity PD-1 antibody, recent patent granted in US, Europe, China, Japan

- Potential of combo with internal asset
- Potential for partnership with biotech/biopharma in combo with external assets
- Potential future marketing approvals in orphan indications with strong unmet medical needs



Not a head-to-head comparison. Differences exist between trial designs and subject characteristics, and caution should be exercised when comparing data across trials.

For illustrative purposes only.



^{*} Patnaik et al. Cancer Chem & Pharm 2021; Brahmer et al. JCO 2010; Patnaik et al. Clin Cancer Res 2015; Papadopoulos et al. Clin Cancer Res 2020

[#]Robert et al. ESMO-TAT 2024

Lusvertikimab

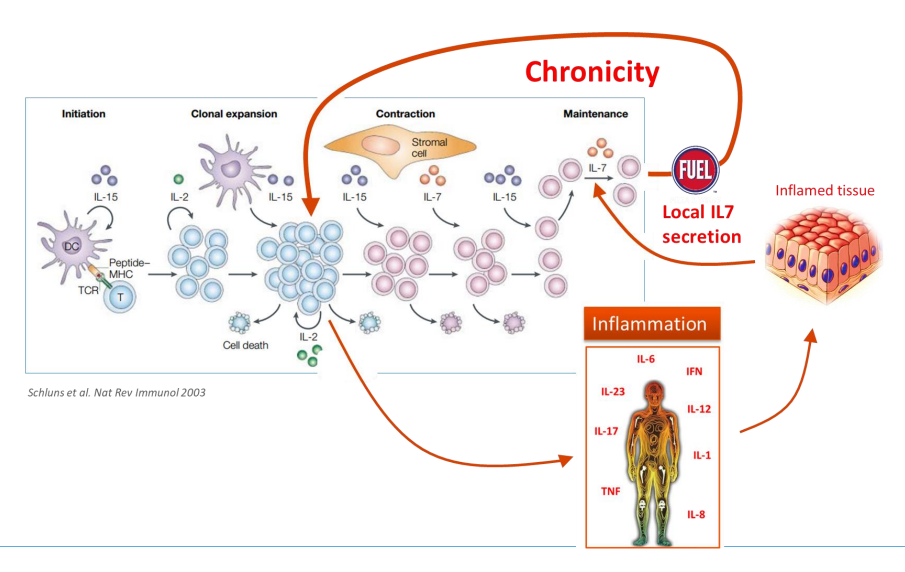
Most advanced anti-IL-7R mAb

Strong biological rational in refractory IBD patients



IL-7 fuels chronic inflammation in tissues

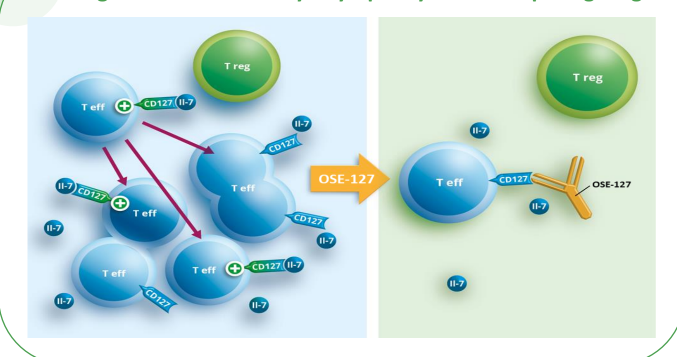
Lusvertikimab controls pathogenic memory T-cell persistence





Lusvertikimab/OSE-127 - Differentiated MoA as full IL-7 receptor antagonist

Tackling the fuel of memory T-lymphocytes while sparing Tregs



A differentiated and highly qualified candidate

- Lusvertikimab, first non-internalizing (fully antagonist) anti-IL-7R mAb¹ and **most advanced** IL-7R antagonist in clinic
- IL7 produced by inflamed tissues sustain T-cell survival and chronicity
- IL-7R pathway overexpression in anti-TNF IBD non-responders²
- Good safety, PK/PD profile in Phase 1³, no cytokine release, confirmed target-engagement
- High preclinical activity in acute leukemia (T and B-ALL)⁴ **ASH Merit Award**



On-going Phase 2 study in UC with clinical readout mid-2024

Lusvertikimab most advanced First-in-Class anti-IL-7R mAb

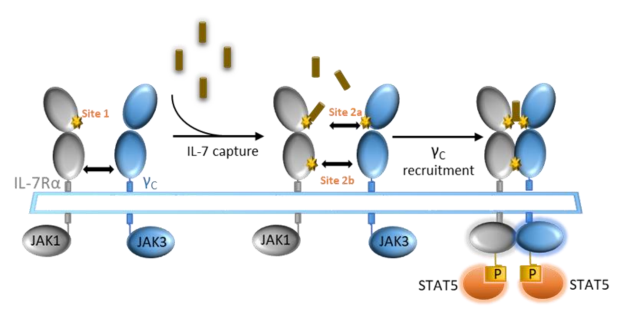
Differentiated by its Mechanism of Action

	OSE IMMUNO (S)	zurabio	©32 BIO	gsk	
Isotype	lgG4	lgG1	lgG1	lgG1	
MoA	- Non-Internalizing ¹ - Full Antagonist IL7R	 Internalizing Antago + Partial Agonist IL7R TSLP Antago T-cell Depletion² 	- TSLP Antago	- Internalizing - Antago + Partial Agonist IL7R	
Phase	2	1b	2 a	1	
Indication	Ulcerative Colitis (IBD) (Completion Enrollment Q4 2023)	Alopecia Areata (not initiated)	Atopic Dermatitis (Initiated Q4 2022) Alopecia Areata (Initiated Q3 2023)	Multiple Sclerosis (Discontinued, High Immunogenicity ^{3,4})	

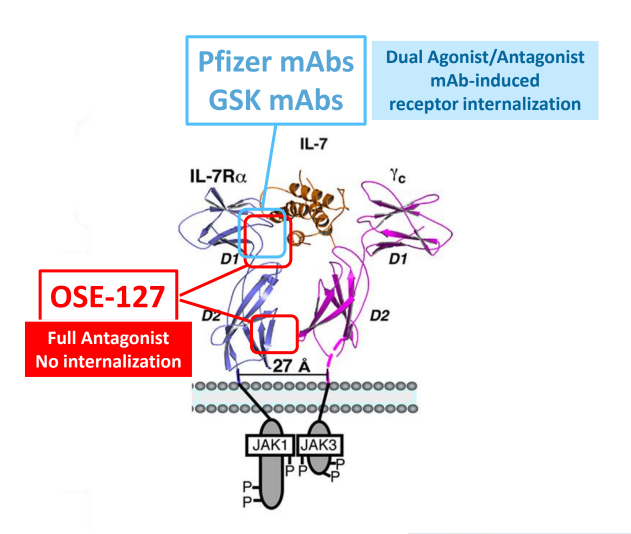
Lusvertikimab - Targets a specific "site 1/2b" Epitope

Full antagonist, preventing receptor internalization & signaling

Cytokine-induced receptor heterodimerization signaling mechanism



Walsh ST et al Immunol. Rev. 2012

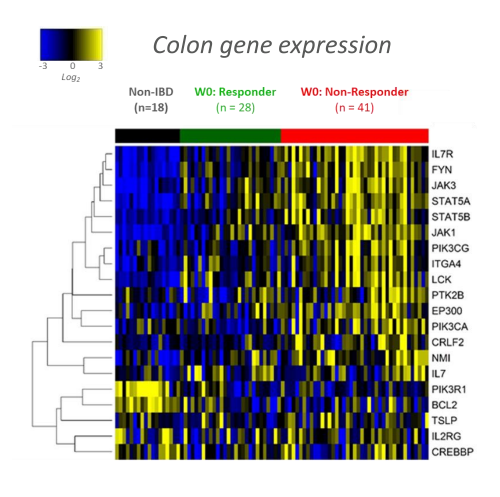


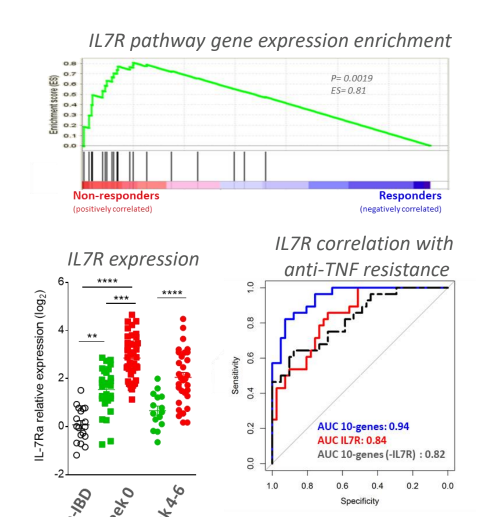




Mucosal IL-7R pathway over-expression in IBD tissues

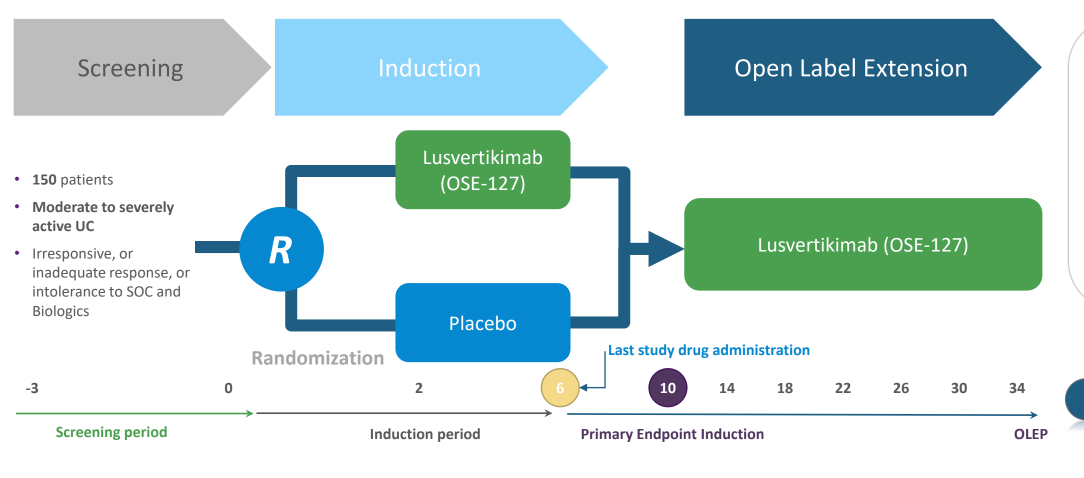
High IL-7R expression in anti-TNF refractory patients





Anti-TNF Responder patients Anti-TNF Refractory patients

Lusvertikimab in moderate-to-severe ulcerative colitis



Positive Recent Futility Analysis¹

- Futility analysis conducted on 33% of the total patient enrolment (n=150)
- Primary endpoint is the efficacy assessment of Lusvertikimab vs. placebo on the reduction of the modified Mayo Score at W10
- 24 weeks open-label extension study planned (NCT04605978)

results expected mid-2024



Colitis after the Interim Futility Analysis

Significant opportunity in Ulcerative Colitis & Acute Lymphoblastic Leukemia targeted markets

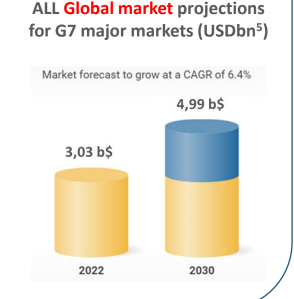
Ulcerative Colitis (UC)

- UC affects 3.3 million patients in US, Europe and Japan
- ~50% UC patients "moderate to severe", requiring methotrexate, corticosteroids, anti-TNFa, JAK etc.
- Despite broad options, remission rates are of only 25-30% leaving most patients without satisfactory treatment



Acute Lymphoblastic Leukemia (ALL)

- ALL is a rare disease with a diagnosed incident cases in EU, US, China, Japan estimated to achieve 26,482 in 2029².
- 40% cases of ALL diagnosed are in adults and among them about 50% present refractory disease or undergo relapse under current conventional therapies³.
- IL-7R expression in >84% of B-ALL and T-ALL samples⁴



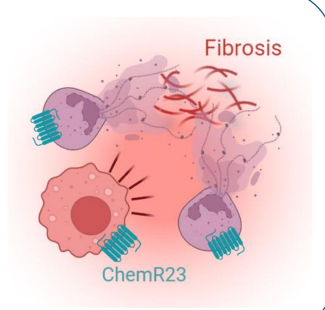
Partnered clinical programs

OSE-230 - Resolving inflammation is an active immune process

abbyie

During chronic inflammation

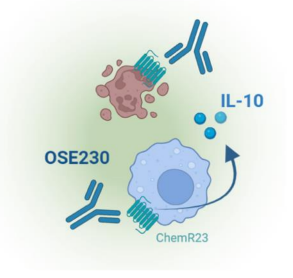
Dying neutrophils **send out inflammatory signals (e.g. NETosis)** that are important in maintaining chronic inflammation & fibrosis



With ChemR23 agonistic mAbs

OSE-230 limits recruitment, survival & NETosis of inflammatory neutrophils & reprograms macrophages, removing further chronic inflammatory signals

Restoration of homeostasis



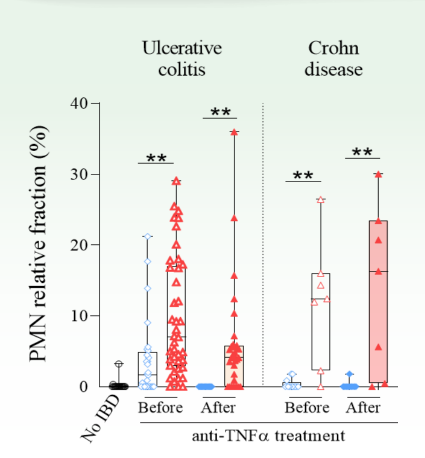
Potential First-in-class pre-IND candidate



OSE-230 - Strong rationale in IBD

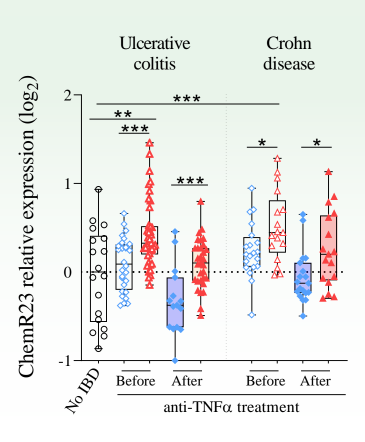


High Neutrophil infiltrates in anti-TNFα refractory patients

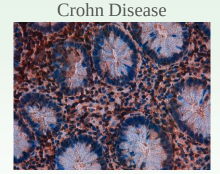


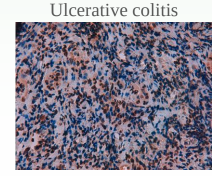
Non Responders Responders

High ChemR23 expression in anti-TNFα refractory patients



ChemR23 staining

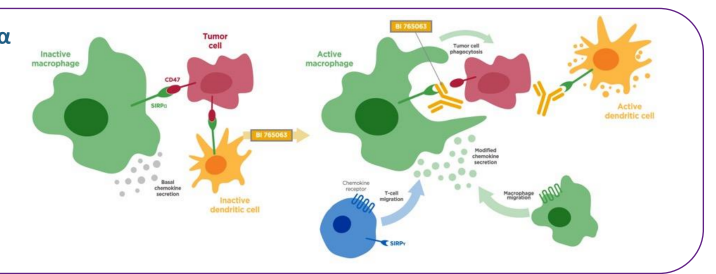




SIRPα inhibition may have a synergistic antitumour effect when combined with ICIs

- Infiltrating myeloid cells promotes immune evasion, and this has generated interest in myeloid-immune targets^{1,2}
 - \circ The CD47–SIRP α interaction transduces inhibitory signals on macrophages and other myeloid cells 2
- Preclinical studies have indicated that CD47 or SIRPα blockade in combination with ICIs may have a synergistic antitumour effect³

The use of SIRPa antagonists to enhance antitumour immunity is currently being explored⁴



	Anti-CD47	Anti-SIRP $lpha$	
Broad/restricted expression	Broad	Restricted to cells of the myeloid lineage] L
Safety signals	Acute anemia, Thrombocytopenia	No hematotoxicity	ŀ
Interaction CD47/SIRPγ	Inhibit human T cells	OSE-172 is SIRP $lpha$ specific	F

Limited side effects expected and less frequent dosing

Boehringer Ingelheim

Higher therapeutic window expected

Favors T cell responses in solid tumors

CD: cluster of differentiation; ICI: immune checkpoint inhibitor; SIRPa: signal regulatory protein-a.



Clinical development overview

Most advanced clinically-tested SIRPα



	Dose Escalation 8	& Expansion studies	ONGOING Studies		
Trial number	NCT03990233	NCT04653142	NCT05249426	NCT05327946	
Phase	la	la	Ib	la	
N	108	36	150	42	
Treatment	BI 765063 +/- Ezabenlimab	BI 765063 +/- Ezabenlimab	BI 765063 + Ezabenlimab ± chemotherapy, cetuximab or VEGF/Ang2 inhibitor	BI 770371 +/- Ezabenlimab	
Patient population	Solid tumors	Solid tumors	HNSCC HCC	Solid tumors	
Region					

Key takeaways from dose escalation

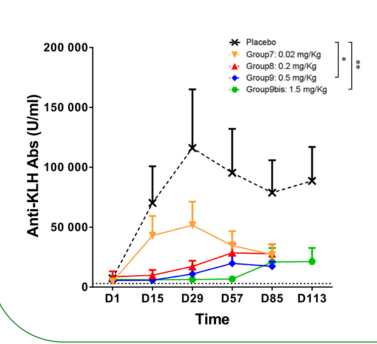
- Safety
 No hematotoxicity reported, no DLTs, MTD not reached^{1,2}
- Efficacy BI765063 in P1a
- 1 PR in HCC, 45% clinical benefit rate as a single agent¹
- 3 PRs in MSS endometrial cancer and CRC in combination with a checkpoint inhibitor²



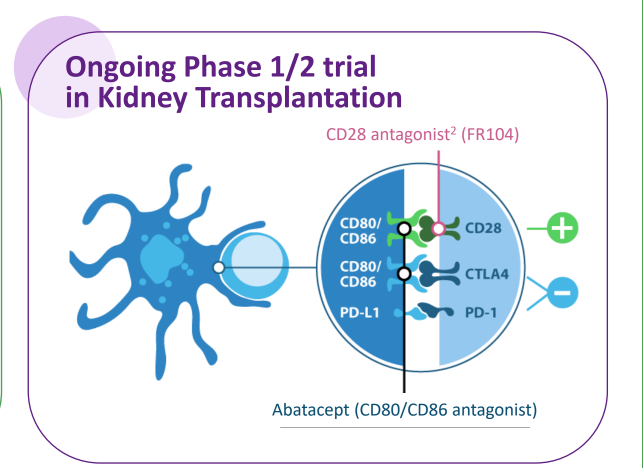
FR104/VEL-101 CD28 antagonist in transplantation



Phase 1 results: Selective CD28 antagonist FR104 persistently reduces antibody responses



- Good safety¹ demonstrated
- Absence of clinical or biological events
- No change in total lymphocyte counts
- No cytokine elevation
- Controls model IgG (anti-KLH) response for up to 57 days
- Controls T follicular helper and IgG responses
- Tfh cells correlated with autoimmune diseases activity



FR104/VEL-101 - Transforming kidney transplant management



Ambitious Partnership with Veloxis

- Deal value: EUR 315m¹ and tiered royalties on sales
- Veloxis is a global leader in transplantation with leading product Envarsus XR (tacrolimus) realizing c. USD 140m² turnover
- o Joined **Asahi Kasei** in FY2019³, a **USD 17bn** annual turnover conglomerate with healthcare representing 17% of sales
- First patient dosed by Veloxis⁴
- Phase 1/2 in kidney transplantation, sponsored and conducted by the Nantes University Hospital, patient enrolment completed

Kidney Transplant Market Opportunity

- 40k+ new kidney transplant annually for an estimated 500k+ people living with a functioning kidney graft in G7 countries
- Chronic exposure to CNIs is associated with renal toxicity, cardiometabolic complications, insufficient graft protection as well as cancer and infections
- FR104/VEL-101 seeks to address challenges associated with current immunosuppressive transplantation regimens using CNIbased therapies
- Potential to provide "One Transplant for Life" with improved patient and graft survival and become the new SoC in transplant



⁻ OSE Immunotherapeutics and Veloxis Pharmaceuticals Enter Into Global License Agreement to Develop, Manufacture, and Commercialize FR104, a CD28 Antagonist, in the Organ Transplantation Market

https://www.asahi-kasei.com/ir/library/presentation/pdf/211005.pdf

^{3 –} https://www.asahikasei.com/ir/library/presentation/pdf/191125eng.pdf

^{4 -} OSE Immunotherapeutics Announces Dosing of the First Participant in a Phase 1 Study of VEL-101/FR104, a Novel Investigational Drug for Kidney Transplant Immunosuppressio

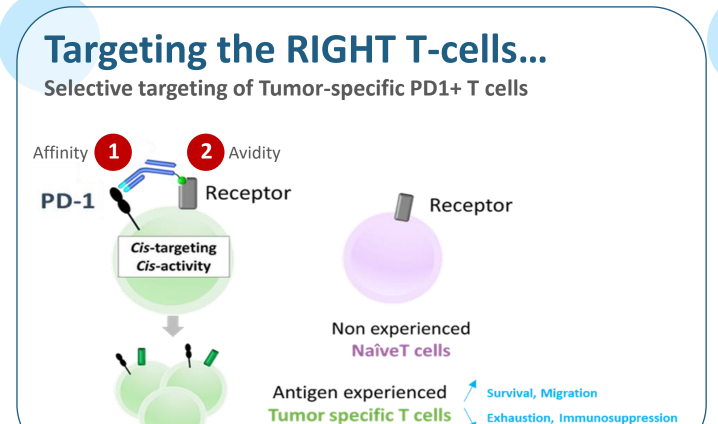
Our Innovative Discovery Engines

Designed to deliver next generation first-in-class immunotherapies

Next-generation anti-PD1 bispecifics

Improving the quality of tumor-specific T-cell responses both in TME & Lymph Nodes





...at the right place Selective Biodistribution in TME + Lymphoid tissues cells to tumors Infiltration of T cells **Lymphoid Tissues** (PD1 expression) Recognition of cancer **Tumor MicroEnvironment** 8 Killing of cancer cells



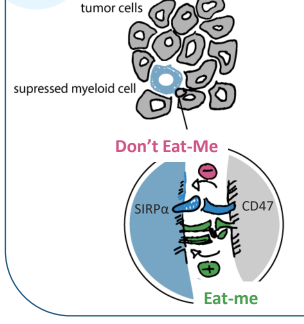
BiCKI®-IL7v* candidate highlighted at AACR 2022*

CLEC-1 - Another way to not get eaten

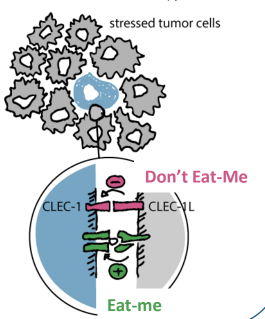
Blocking myeloid immune checkpoint from delivering another "Don't-eat-me" signal



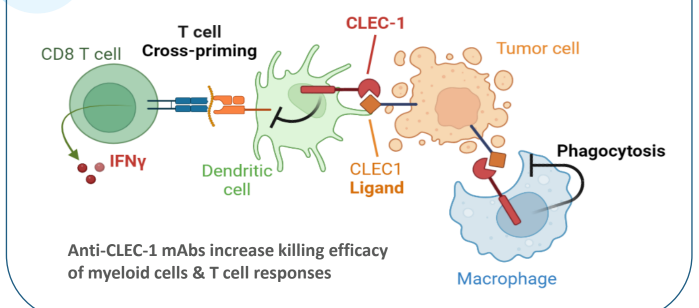
Tumor homeostasis



+ damage-inducing interventions (e.g. chemo-, radio-, immunotherapy)



CLEC-1 mAbs disrupt tumor homeostasis²



First-in-class preclinical LEAD validation¹







An experienced Executive leadership team



Nicolas Poirier, PhD CEO, CSO

- 18+ year experience in biotech/immunotherapy
- Advanced 5 novel immunotherapies to clinic
- Leading to 4 pharma deals
- Global Management & Finance (INSEAD, HEC)



Anne-Laure Autret-Cornet Chief Financial Officer

- 15+ year experience in Finance / Biotech
- Graduated from ESSCA Management school
- Corporate Finance, HEC



Dominique Costantini, MD Chief Development & Strategy

- 30+ years in product development/ marketing
- Chairwoman, Co-founder
- IPO completion in 2015



Jean-Jacques Mention, PhD
Chief Business Officer

- 15+ years of Research in Immunology at King's College London, Institut Pasteur
- 7+ year experience in Business Development



Aurore Morello, PhD Head of Research

- 13+ year experience in Immunotherapy
- International Postdoctoral Fellowship (MSKCC, NYC)



Silvia Comis, MD
Head of Clinical

- 30+ year experience in Pharma
- Previously Senior Medical Director IQVIA, and European Head of Early Products Medical Affairs in oncology at Novartis



Valérie Gabarre, PharmD Medico-Marketing Director

- 25+ years of experience in Pharma/Biotech, in Medico-Marketing & Sales - EU & Global, Immunotherapy & Oncology
- Global Network of Leaders & Corporative Groups in Oncology
- PharmD

A Board of Directors combining international expertise in drug development, industry & finance & experience in listed biotech companies



Dominique Costantini, MD Chairwoman, Chief Development & Strategy

- 30+ years in product development/ marketing
- Chairwoman, Co-founder
- IPO completion in 2015



Maryvonne Hiance Vice Chairwoman

- Founder and CEO of Effimune
- General Manager SangStat Atlantic, DrugAbuse Sciences
- Former President & Vice
 President of France Biotech



Nicolas Poirier, PhD
Director, Chief
Executive Officer &
Chief Scientific Officer

- 15+ year experience in biotech/immunotherapy
- Advanced 5 novel therapies to clinic
- 4 pharma deals
- Global Management, INSEAD



Anne-Laure Autret-Cornet
Director representing the
employee shareholders,
Chief Financial Officer

- 15+ years in Finance & Biotech
- ESSCA Management School
- Finance Corporate, HEC



Brigitte Dréno, MD Independent Director

- Head Depart of Dermatology Nantes
- Director of Biotherapy Clinical Investigation Centre
- Operational functions and research responsibilities



Didier Hoch, MD Independent Director

- 25+ years in pharma and vaccine industry
- Several functions incl. commercial, marketing, general management



Eric Leire, MD Independent Director

- Genflow Bioscience CEO
- Previously chairman & CEO of several biotech listed in US
- Previous Marketing Director position in Pharma US & EU

International SAB - Renowned experts in IO and I&I





Wolf-Hervé Fridman, MD Chairman of the SAB, Professor Emeritus of Immunology at the Université de Paris, France





Myriam Merad, MD, PhD
Director of the Precision
Immunology Institute at Mount
Sinai School of Medicine in New
York and the Director of the
Mount Sinai Human Immune
Monitoring Center (HIMC)







Charles N. Serhan, PhD, DSc Professor of Anaesthesia (Biochemistry and Molecular Pharmacology) at Harvard Medical School, Professor of Oral medicine, Infection and Immunity at Harvard School of Dental Medicine





M.M.Sc Professor of Genomic Medicine & Surgical Oncology, UT MD Anderson Cancer Center

Jennifer Wargo, MD,





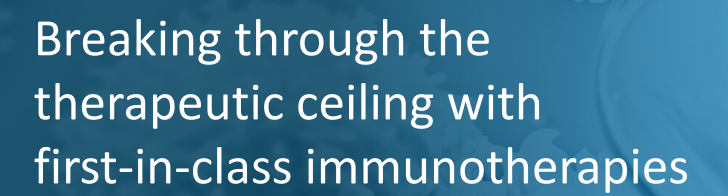
Bernard Malissen, PhD
Group Leader at Centre
d'Immunologie de MarseilleLuminy and Founding-Director of
Center for Immunophenomics,
Marseille, France





Sophie Brouard, PhD
Immunologist and Director in
Veterinary Sciences, Director of
Research at the Institut National
de la Santé et Recherche
Médicale (Inserm, National
Institute for Health and Medical
Research) in Nantes

OSE IMMUNO THERAPEUTICS



Immuno-Oncology & Immuno-Inflammation

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Company Information: http://ose-immuno.com/en/