

MaaT Pharma Microbiota as a Therapy

July 2023

Disclaimer

This document has been prepared by MaaT Pharma (the "Company") and is for information and background purposes only.

While the information contained herein has been prepared in good faith, neither the Company, nor its shareholders, directors, officers, agents, employees, or advisors give, have given or have authority to give, any representations or warranties (express or implied) as to, or in relation to, the fairness, accuracy, reliability or completeness of the information in this document, or any revision thereof, or of any other written or oral information made or to be made available to any interested party or its advisers, including financial information (all such information being referred to as "Information"), and liability therefor is expressly disclaimed. Accordingly, neither the Company nor any of its shareholders, directors, officers, agents, employees, affiliates, representatives or advisers take any responsibility for, or will accept any liability whether direct or indirect express or implied, contractual, tortuous, statutory or otherwise, in respect of the accuracy or completeness of the Information or for any of the opinions contained herein or for any errors, omissions or misstatements or for any loss, howsoever arising from this document.

The information and opinions contained in this document are provided as of the date of this document only and may be updated, supplemented, revised, verified or amended, and thus such information may be subject to significant changes. The Company is not under any obligation to update the information or opinions contained herein which are subject to change without prior notice.

The information contained in this document has not been subject to independent verification and are qualified in their entirety by the business, financial and other information that the Company is required to publish in accordance with the rules, regulations and practices applicable to companies listed on the regulated market of Euronext in Paris, including in particular the risk factors and other information in the Company's Document d'enregistrement (Registration Document) registered by the French *Autorité des marches financiers* (Financial Markets Authority) (the "AMF") on October 1st, 2021 under no. I.21-0057 and its supplement on October 14, 2021 under no. I.21-0061 and in any other periodic report, which are available free of charge on the websites of the Company (https://www.maatpharma.com/) and the AMF (www.amf-france.org).

No representation, warranty or undertaking, express or implied, is made as to the accuracy, completeness or appropriateness of the information and opinions contained in this document. The Company, its subsidiaries, its advisors and representatives accept no responsibility for and shall not be held liable for any loss or damage that may arise from the use of this document or the information or opinions contained herein.

This document contains information on the Company's markets and competitive position, and more specifically, on the size of its markets. This information has been drawn from various sources or from the Company's own estimates which may not be accurate and thus no reliance should be placed on such information. Any prospective investors must make their own investigation and assessments and consult with their own advisers concerning any evaluation of the Company and its prospects, and this document, or any part of it, may not form the basis of or be relied on in connection with any investment decision.

This document contains certain forward-looking statements. These statements are not guarantees of the Company's future performance. These forward-looking statements relate to the Company's future prospects, developments and marketing strategy and are based on analyses of earnings forecasts and estimates of amounts not yet determinable.

Forward-looking statements are subject to a variety of risks and uncertainties as they relate to future events and are dependent on circumstances that may or may not materialize in the future. Forward-looking statements cannot, under any circumstance, be construed as a guarantee of the Company's future performance and the Company's actual financial position, results and cash flow, as well as the trends in the sector in which the Company operates, may differ materially from those proposed or reflected in the forward-looking statements contained in this document. Even if the Company's financial position, results, cash-flows and developments in the sector in which the Company operates were to conform to the forward-looking statements contained in this document, such results or developments cannot be construed as a reliable indication of the Company's future results or developments. The Company does not undertake any obligation to update or to confirm projections or estimates made by analysts or to make public any correction to any prospective information in order to reflect an event or circumstance that may occur after the date of this document.

All persons accessing this document are deemed to agree to all the limitations and restrictions set out above.

A Uniquely-Positioned Microbiome Company





Multi-asset clinical and preclinical pipeline with near-term, value-creating catalysts

Proprietary gutPrint® metagenomics technology platform driving product candidate generation

European cGMP production facilities supporting versatile product range and optimized positioning

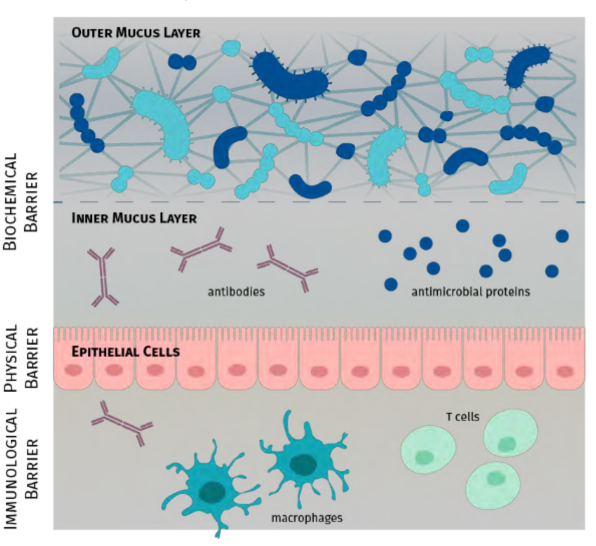
Strong IP portfolio of 14 patent families that **provides protection until 2036-2042 in all major markets**

Strong leadership team with a proven track record and supported by a **scientific advisory board of global experts** and **top tier specialist investors**





Host - Microbiota Interactions are Critical for a Functional Immune System



A rich and diversified gut ecosystem actively modulates the immune system functionality

- A diversified microbiome contributes to the education and modulation of our immune system throughout life
- Bacterial richness and mucus layer prevent colonization by pathogens and improve gut barrier

80%

cellular host defense localized in the gut

Cross-section of a healthy gut



IMMUNOLOGICAL

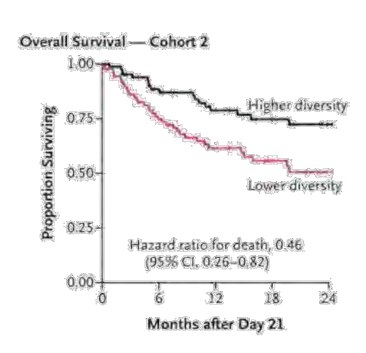
Diversity matters! Higher gut microbiome diversity is associated with ...

Liquid Tumors

Lower incidence and lower mortality from aGvHD*,2

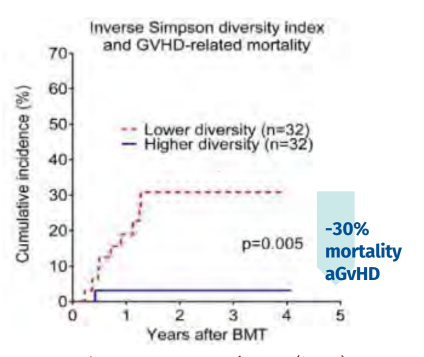
Solid Tumors

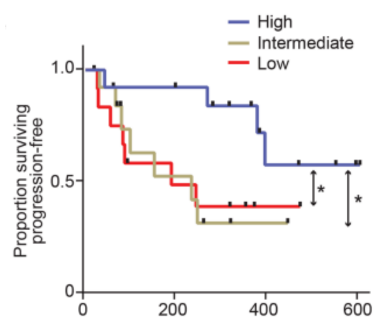
Higher response rate to ICI* in patients with metastatic melanoma³



Higher survival rate in patients

receiving allo-HSCT *,1





MaaT Pharma MET Inverse Simpson (mean): 24

*allo-HSCT: allogeneic hematopoietic stem cell transplantation; aGvHD: acute Graft-vs-host-Disease; ICI: Immune Checkpoint Inhibitors ¹Peled, J.U. & al N Engl J Med 2020;382:822-34; ²Ghani, 2021; Jenq RR. et al, Biol Blood Marrow Transplant 21 (2015) 1373e1383; Pamer, Blood, 2014; ³Gopalakrishnan et al., Science, 2017, see also Routy et al, Science, 2018; Vetizou et al Science 2015;



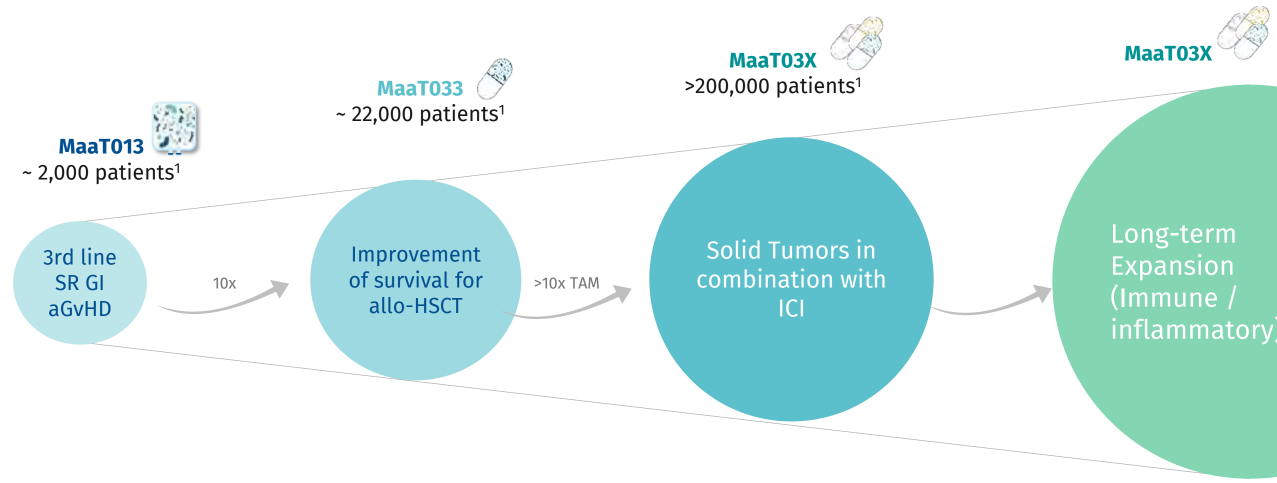
MaaT Pharma's Microbiome Ecosystem Therapy (MET) platform has generated a diverse line of product candidates

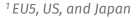
Platform	Program	Indication	Preclinical	Phase I	Phase II	Phase III	Upcoming milestone
	MaaT013	aGvHD	ARES				DSMB H2 2023
		IO Melanoma	PICASSO (IST)				Results H2 2024
MET-N	ı	1					
	MaaT033	HSCT	PHOEBUS				Initiation H2 2023
		ALS	IASO				FPI 2023
MET-C	MaaT03X	Multiple	R&D				Targeting FIH 2024

IST: Investigator Sponsored Trial; aGvHD: acute Graft-versus-Host Disease; IO: Immuno-oncology; HSCT: Hematopoietic Stem Cell Transplantation; ALS: Amyotrophic Lateral Sclerosis



Looking ahead: addressing growing market opportunities with severe medical need



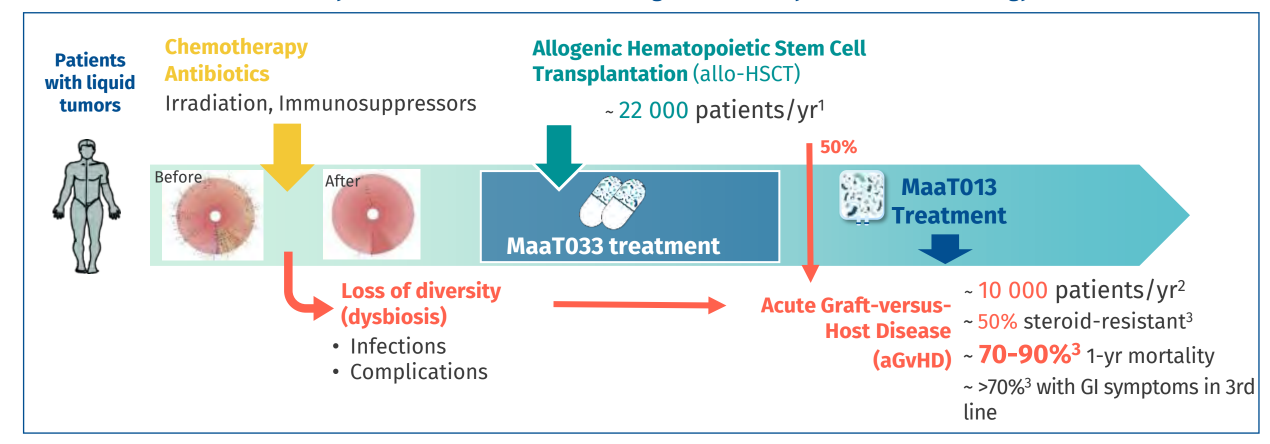






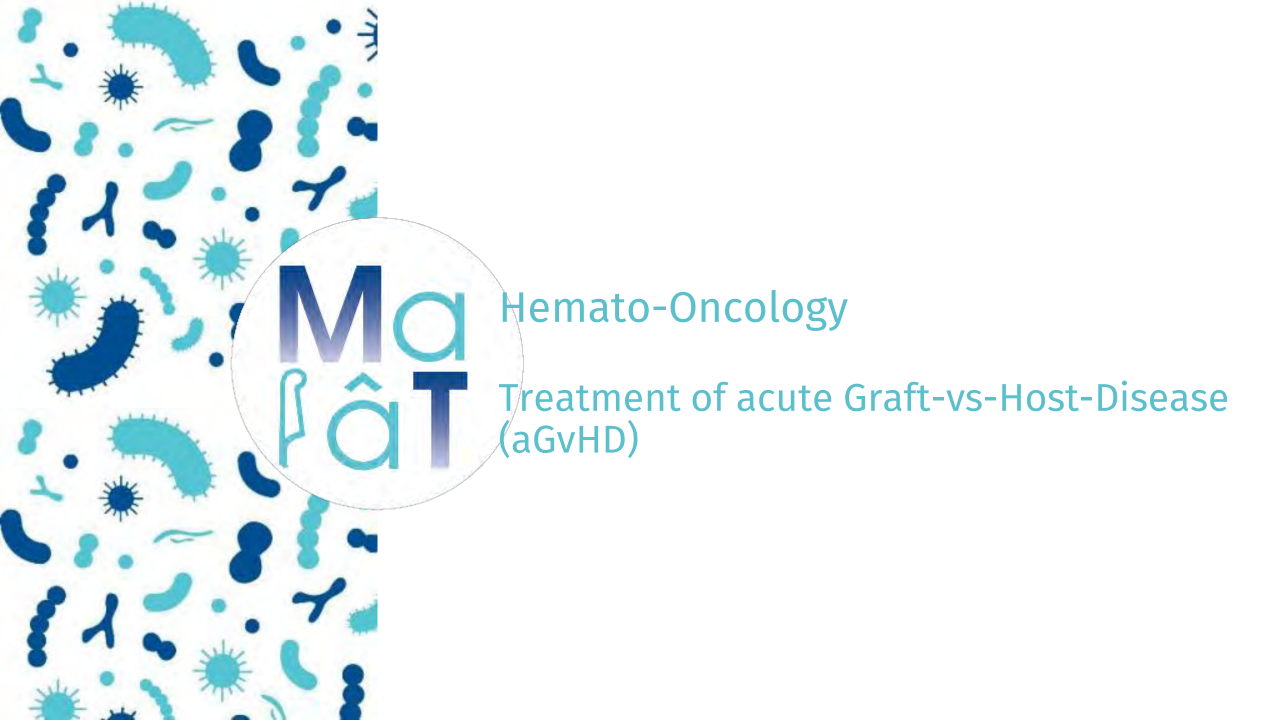
MaaT013 and MaaT033 aim to restore the gut microbiota to improve survival in patients with liquid tumors

Intestinal dysbiosis is associated with higher mortality in hemato-oncology



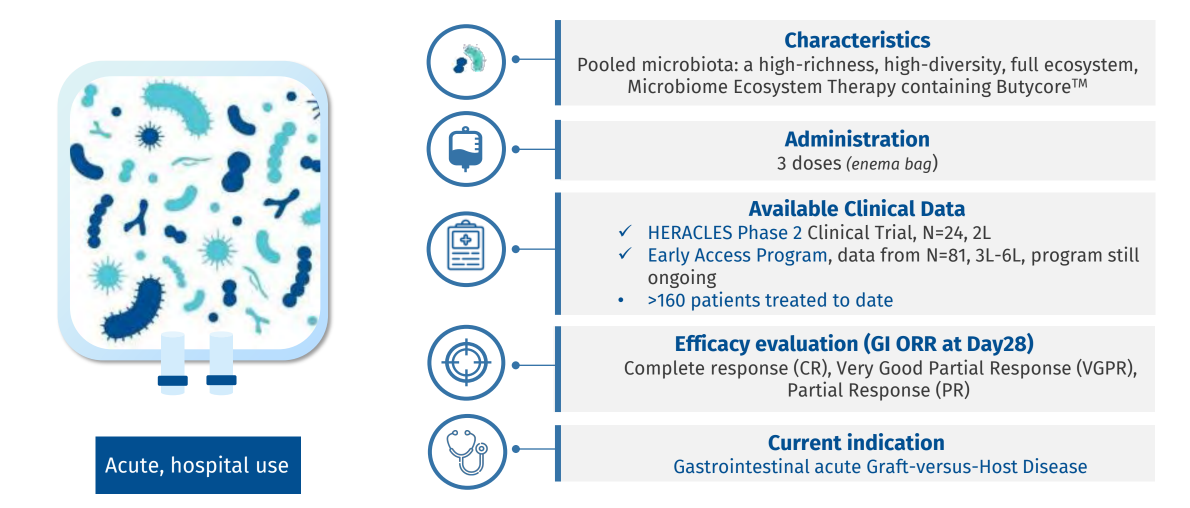
1. EU5 + US : (~ 20 500 primary procedures with an additional 7%-10% recurring), 2. EU5 + US, ³ According to MAGIC database







MaaT013: restore the microbiome to *cure* gastrointestinal acute Graft vs. Host Disease



MaaT013 has received Orphan Drug Designation from FDA and EMA



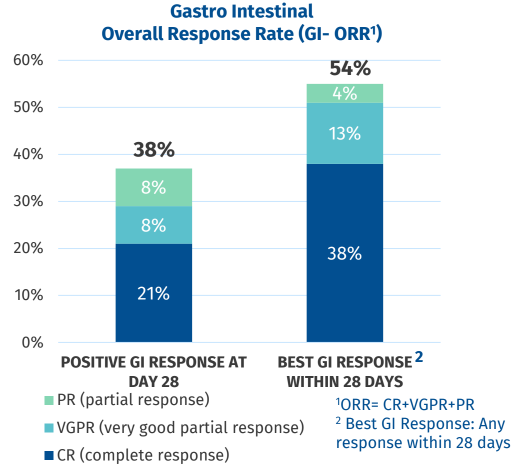


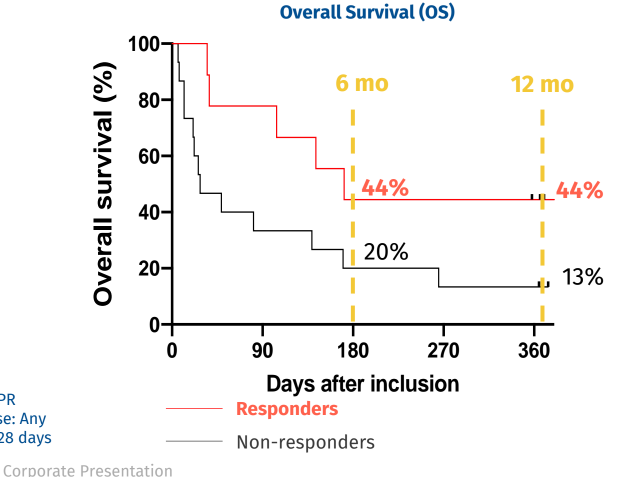




HERACLES Phase 2 Clinical Trial Promising results in a very severe (III-IV) GI aGvHD population

- N=24 patients, 96% grade III (4% grade IV), 2nd line (Steroid-resistant)
- Very good safety and tolerability profile
- MaaT013 increases responders' gut microbiome diversity







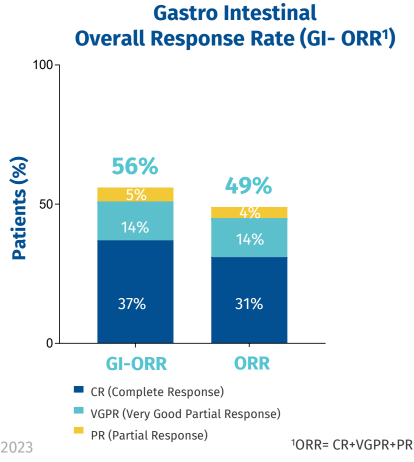


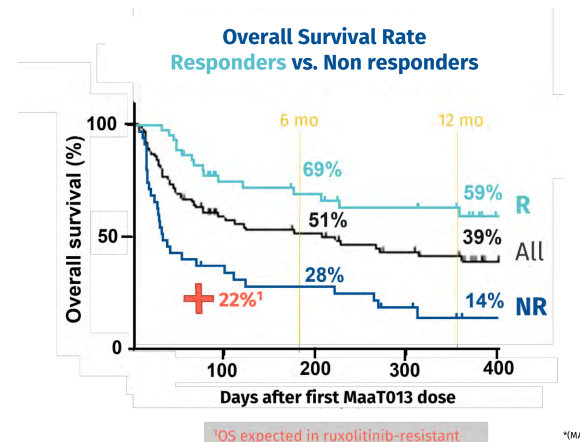




Early Access Program (EAP) is corroborating positive data in an advanced, severe and more diverse GI aGvHD population

- N=81 84% SR; Grade III (51%) or Grade IV (38%) aGvHD*, Up to 6 lines of prior treatment (median 2; 66/81 received ruxolitinib)
- Good tolerability and safety profile in a fragile population
- Data presented in December 2022 at the 64th Annual Meeting of the American Society of Hematology (ASH)





patients at 2 months (REACH1 study)



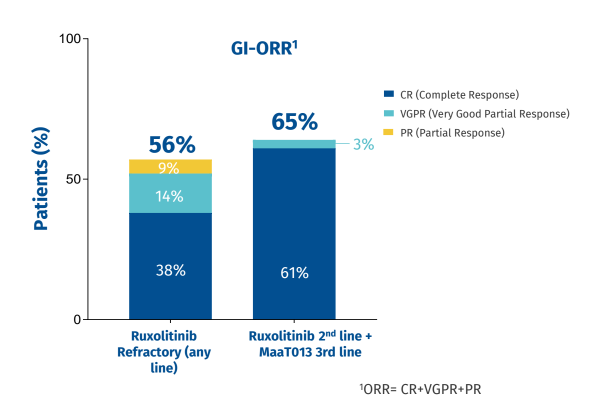




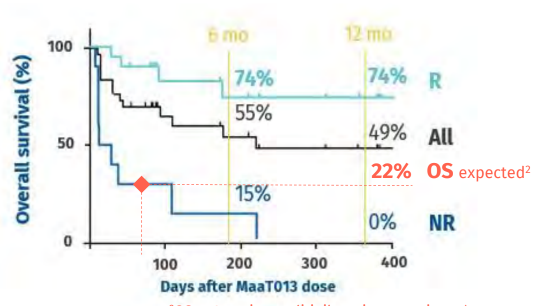
OCIETA OF WITH ON

Among 81 patients in EAP, excellent response to MaaT013 was shown in n=31 corticoid and ruxolitinib-refractory patients

- N=31 Ruxolitinib-refractory in 2nd line, MaaT013 given in 3rd line
- Clinical response to MaaT013 translates to an important increased overall survival
- Data presented in December 2022 at the 64th Annual Meeting of the American Society of Hematology (ASH)



Overall Survival Rate in ruxolitinib-refractory patients Responders vs. Non responders



²OS at 2 mo in ruxolitinib-resistant patients (REACH1 study)

This patient population resembles the ongoing Phase 3 ARES clinical trial (NCT04769895) being conducted in Europe.



aGvHD Phase 3

The ARES Phase 3 study is designed to establish MaaT013 as the 3rd line agent in GI aGvHD treatment

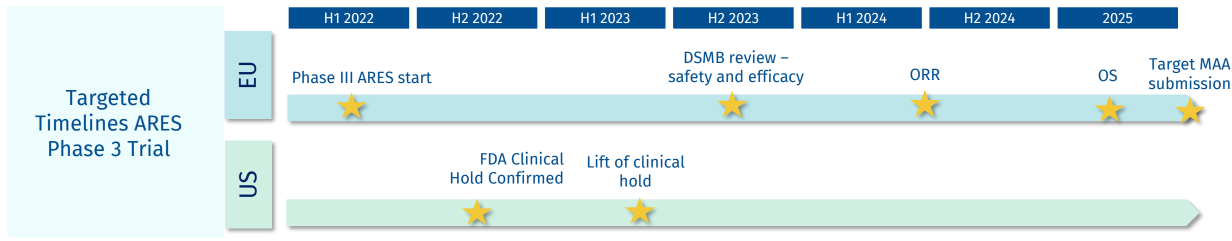
- Further investigation currently ongoing in a pivotal single arm Phase 3 trial of MaaT013 as 3rd line
- 75 patients with GI aGvHD who are refractory to both steroids and ruxolitinib
- Primary endpoint: GI-ORR at Day28

EUROPE: ongoing clinical trial

- ✓ First patient dosed in Q1 2022
- CTA approved in 6 European countries: Austria, Belgium, France, Germany, Italy, Spain.

USA: FDA IND cleared to enable clinical activity in the U.S.

- ✓ Lift of clinical hold in April 2023
- MaaT Pharma intends to consult with the FDA on the next steps of the regulatory process to bring MaaT013 to US patients in the most expeditious way possible.



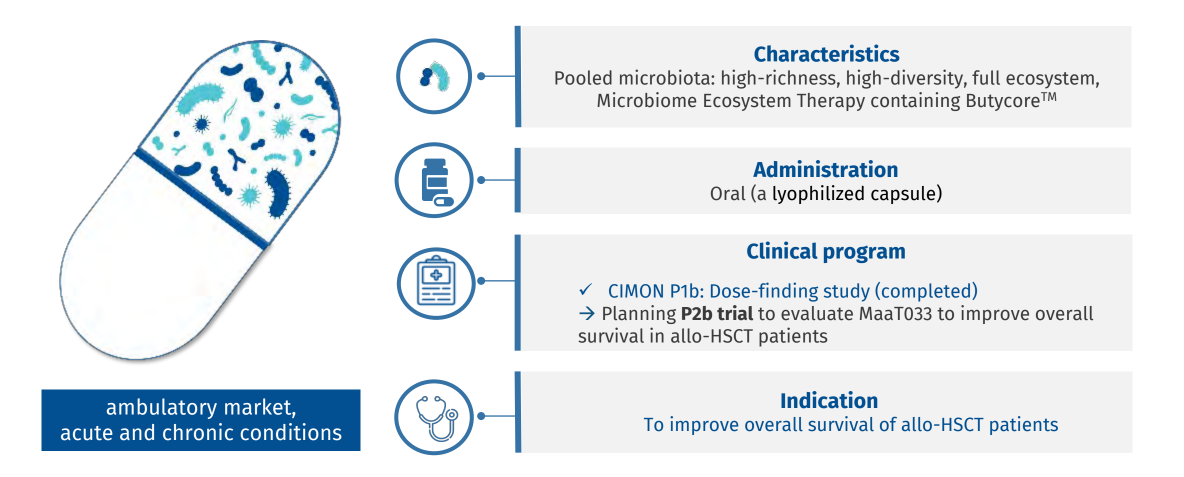


Clinical trials.gov: NCT04769895





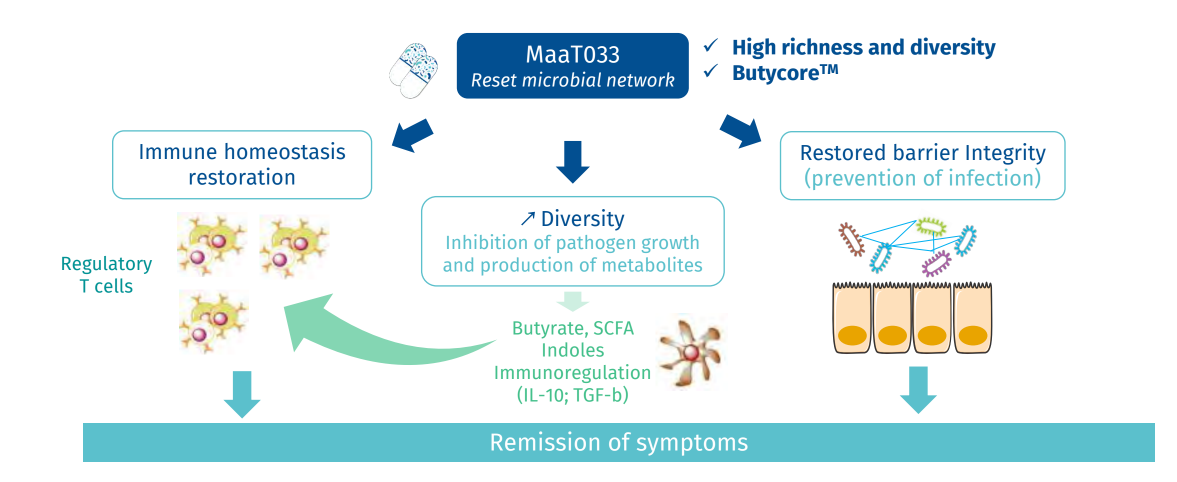
MaaT033: An oral capsule to be used as an *adjunctive therapy* for patients with hematological malignancies receiving HSCT







MaaT033's MOA aims to restore and protect the gut microbiota, to improve overall survival in allo-HSCT patients



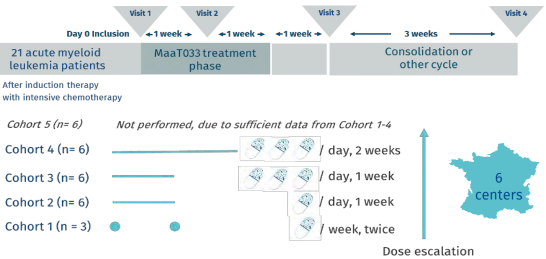


Phase Ib CIMON study: Positive dose ranging study with promising engraftment and safety data

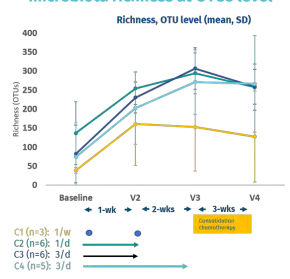




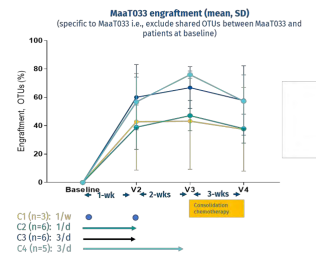




MaaT033 induces an increased microbiota richness at OTUs level



MaaT033 bacterial engraftment is inversely correlated with patients' baseline microbiota richness



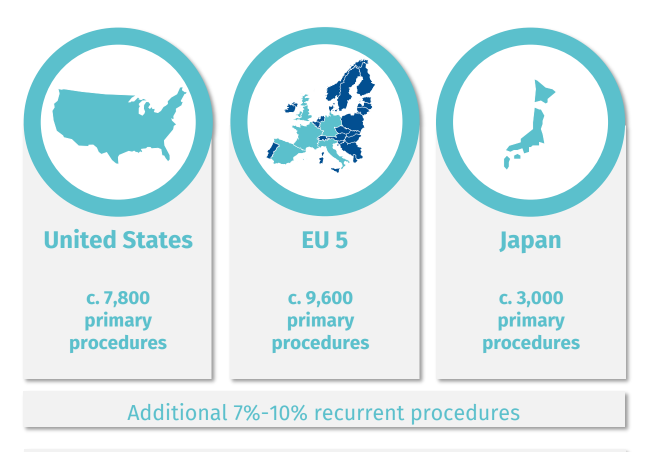
First clinical POC of MaaT033 oral formulation

- Robust and persistent engraftment
- ✓ Good safety profile:
 - 21 patients exposed, 20 completed.
 - 100% drug compliance.
 - 4/4 positive DSMB meetings
- ✓ Engraftment following MaaT033 treatment correlated with increased anti-inflammatory markers.



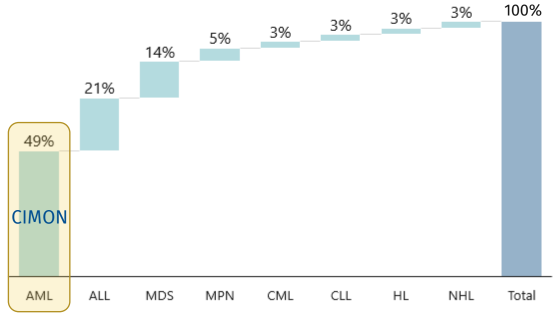


CIMON results open an attractive market opportunity: Improving survival in patients receiving allo-HSCT



Approximately 22,500 procedures/year

Hematological Malignancy Patients Receiving Allo-HSCT¹



AML: acute myeloid leukemia; ALL: acute lymphoblastic leukemia; MDS: myelodysplastic syndrome; MPN: myeloproliferative neoplasms; CML: chronic myeloid leukemia; CLL: chronic lymphocytic leukemia; HL: Hodgkin's Lymphoma; NHL: Non Hodgkin Lymphoma

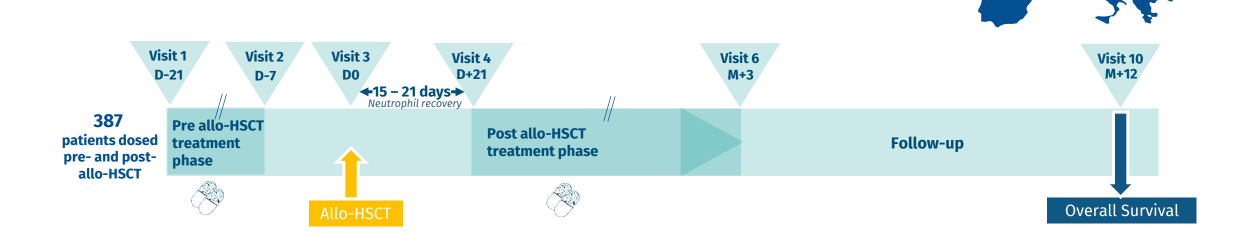


¹EBMT aHSCT Survey, 2017 (published in Bone Marrow Transplantation (2019) 54:1575–1585), Global Data 2020

56 sites targeted1

The Phase 2b is designed to establish MaaT033 as an adjunctive treatment for patients with hematological malignancies receiving HSCT

- 387 patients in a randomized, double-blind, placebo-controlled international study
- Primary endpoint: efficacy of MaaT033 in improving overall survival at 12 months
- Study is expected to start in H2 2023, results are expected in H1 2026





¹Expansion to US sites subject to ongoing discussion with the FDA for MaaT013 IND;



A diverse gut microbiome increases survival in patients receiving immune checkpoint inhibitors (ICI)

FMT from ICI responders to ICI non-responding patients with metastatic melanoma

√ 6/15

√ 3/10

Non-responders

→ Responders

(Davar et al, 2021)

Non-responders

→ Responders
(Baruch et al, 2021)



- Immune check-point inhibitors (ICI) therapies have established themselves as key therapeutic options in solid tumors, but ORR may be as low as 20% in some indications.
- Richness, Diversity and composition of gut microbiome drive survival and ICI toxicity in patients receiving ICI^{1,2,3,4}
- FMT from ICI responders (R) could induce response in metastatic melanoma non-responders (NR)^{5,6}

→ Leveraging the gut microbiome richness, diversity and its key functional networks may be a game-changer in immuno-oncology in the coming years

^{4.} Mc Culloch et al, Nat Med 2022; ^{5.} Baruch et al, Science 2021; ^{6.} Davar et al, Science 2021



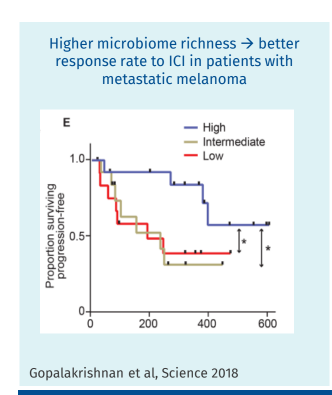
^{1.} Gopalakrishnan et al, Science 2018, ^{2.} Matson, et al Science 2018; ^{3.} Routy et al, Science 2017;

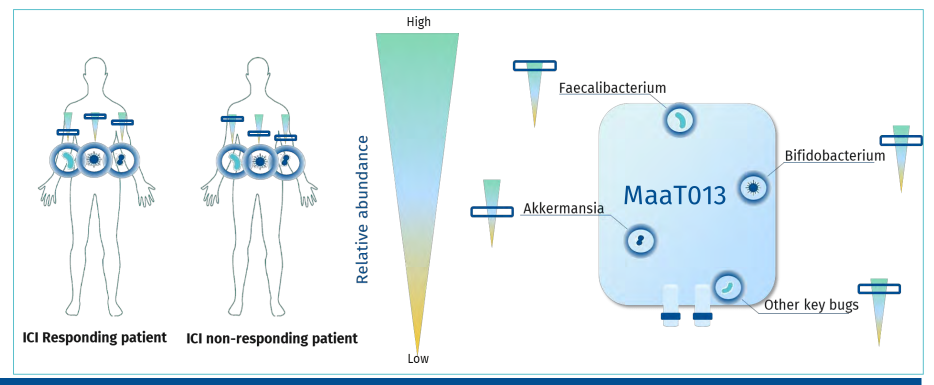


MaaT013 ensures high diversity and contains specific bacterial strains that have been identified to improve ICI response









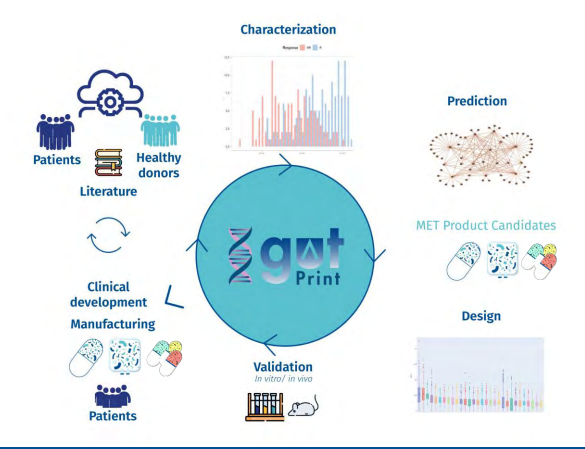
Ongoing Phase IIa PICASSO trial¹, in collaboration with Assistance Publique - Hôpitaux de Paris (sponsor).

- ✓ RCT [MaaT013 + ICI] vs. [Placebo + ICI] in 60 metastatic melanoma patients
- ✓ **Key study endpoints** after 23 weeks of treatment:
 - MaaT013 safety profile vs placebo as add-on treatment to Ipilimumab + Nivolumab
 - MaaT013 best-overall response rate vs placebo as add-on treatment to Ipilimumab + Nivolumab





Proprietary gutPrint® platform synergizes multi-source data to generate innovative and indication-specific microbiome ecosystem therapies



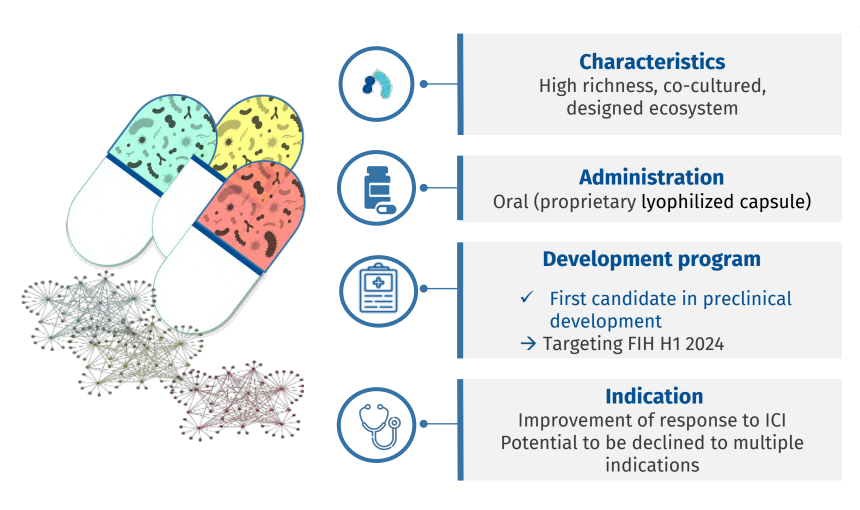
gutPrint® is the engine that drives MaaT Pharma's MET product candidate generation capabilities

to broaden and strengthen the pipeline

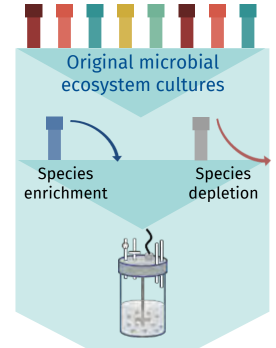
✓ Full cycle in 15 months to enter clinical phase

MaaT03X: Modulate the gut microbiome to *improve response* to Immune Checkpoint Inhibitors treatment in solid tumors





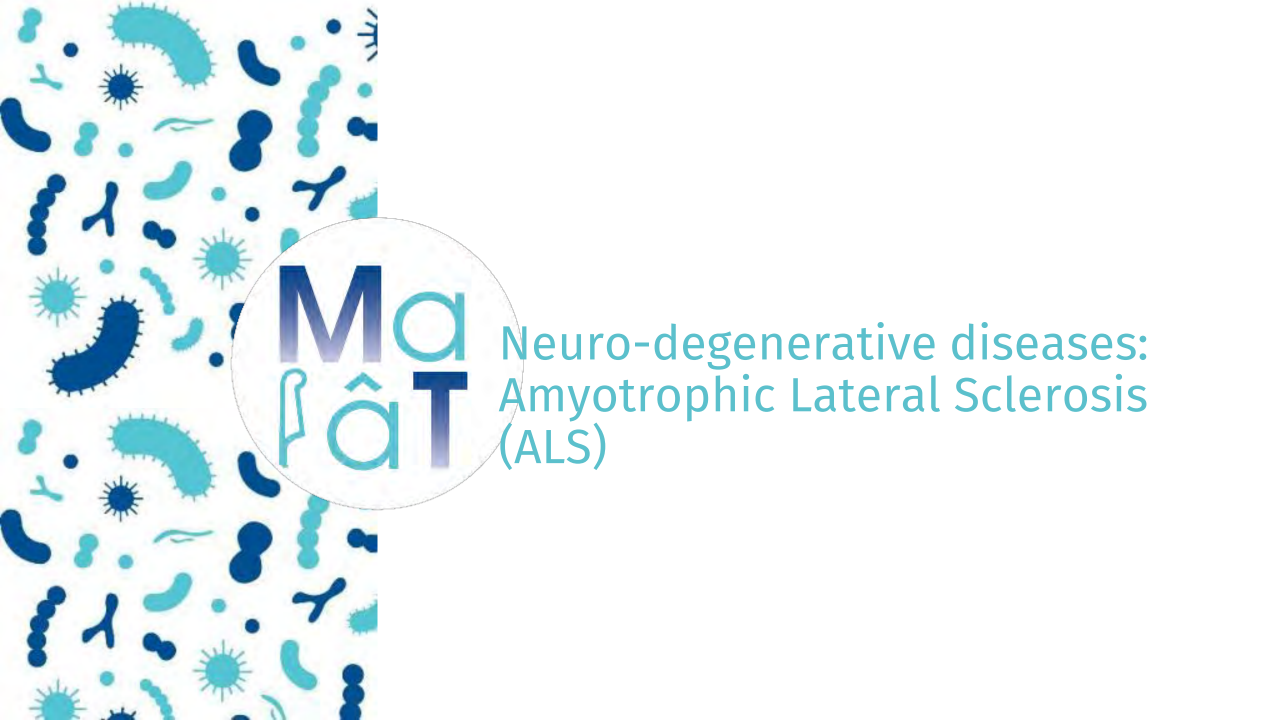
Customizable, donor-independent, scalable co-culture process



CO-CULTURED A FULL ECOSYSTEM

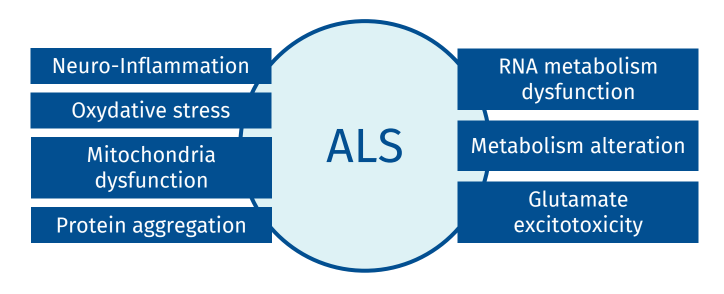








Amyotrophic Lateral Sclerosis: a uncurable disease leading to death within 3-5 years after diagnosis



~ 60,000 patients in US & EU by 2040 (2016)¹ Paralysis
Death 3-5 years after
diagnosis²

No curative treatment Few symptomatic treatments

¹Arthur, K., Calvo, A., Price, T. *et al.* Projected increase in amyotrophic lateral sclerosis from 2015 to 2040. *Nat Commun* **7**, 12408 (2016). https://doi.org/10.1038/ncomms12408



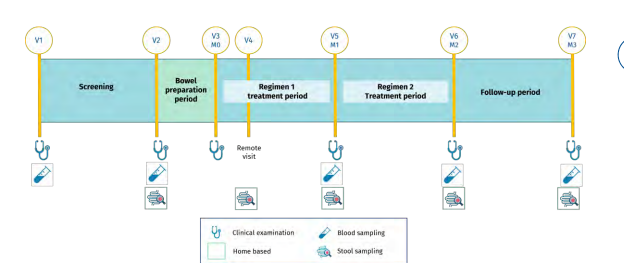
²https://tousensellescontrelasla.fr/la-sla-cest-quoi/





IASO trial is designed to develop the potential first oral microbiotherapy in ALS*

- Up to 15 patients in a pilot, open-label, Phase 1b study in France
- Study is expected to start in 2023, results are expected in H1 2024









With the support of



Key study endpoints:

- Assess safety and tolerability of multiple doses of MaaT033
- Assess gut microbiota composition evolution
- Identify biomarkers sensitive to treatment before considering a larger randomized controlled efficacy study

Potential to extend further to other chronic CNS diseases/ immuno-inflammatory diseases as MaaT Pharma collects data and in-depth understanding of MOA.

^{*} One academic study testing native gut microbiome in ALS patients using an invasive administration procedure on going.



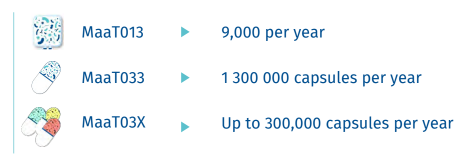


Building Europe's largest specialized cGMP manufacturing facility for Microbiome Ecosystem Therapies





Building a dedicated 1,600m² site (which could be doubled) to support up to 2034 needs of clinical and then commercial production of native MET (MaaT013 & MaaT033) and R&D and clinical batches of cultured products MaaT03X (est. first step):





Site provides for a fully integrated Manufacturing and development platform to allow for a quick and efficient product development, scaleup and GMP process.



Ongoing CSR global strategy: participating in a reforestation program in France (opting for more ecological items (GoGreen) and joining the Cap Vert pour la forêt program and furnishing the plant with sustainable & used materials







Delivery expected in mid-2023



July 2023 Corporate Presentation



Meaningful milestones in both the near and long term

MaaT013 (pooled enema)

- GvHD | ARES P3 DSMB H2
- IO in Melanoma | PICASSO P2a Internal Review

MaaT033 (pooled capsule)

- HSCT | PHOEBUS P2b Launch H2
- ALS | IASO P1b FPI

MaaT013 (pooled enema)

- GvHD | ARES P3 ORR mid-year
- IO in Melanoma | PICASSO P2a Results H2

MaaT033 (pooled capsule)

- HSCT | PHOEBUS P2b Safety
 Interim H2
- ALS | IASO P1b Results H2

MaaT03X (co-cultured capsule)

Undisclosed | Targeting FIH

MaaT013 (pooled enema)

- GvHD | Final Results (OS)
- GvHD | Commercial launch

MaaT033 (pooled capsule)

HSCT | PHOEBUS Results

MaaT03X (co-cultured capsule)

Undisclosed | P1b Results

2023

2024

2025/2026+



Key differentiators of MaaT Pharma from other microbiome competitors

Leveraging the complexity of the microbiome

Pioneering a full ecosystem approach to restore host/microbiome immune symbiosis, based on proprietary Al and manufacturing capacities

Manufacturing versatility

In-house cGMP manufacturing scalability for both native and co-cultured products and end-to-end control of its supply chain

Marian

Oncology focus

Addressing **high unmet needs** in the hemato-oncology and immuno-oncology therapeutic areas

Established proof of concept

First company to reach Phase 3 testing for a microbiome product in oncology globally



A highly experienced team



Hervé Affagard Founder & CEO



Siân Crouzet Chief Financial Officer









Carole Schwintner, Ph.D Chief Technology Officer



Philippe Moyen, Ph.D **Chief Operating Officer**













Pr. Gervais Tougas Chief Medical Officer



Nathalie Corvaia, Ph.D Chief Scientific Officer





BOARD OF DIRECTORS



Karim Dabbagh ** Chairman of the Board **President - CEO Second Genome**



Nadia Kamal* Non-Executive Director

Director of the Technology, Health, and Innovation Divisions – Harmonie Mutuelle



lean-Marie Lefèvre Non-Executive Director **President - Biocodex**



Isabelle de Crémoux Non-Executive Director **CEO & Managing Partner - Seventure**



Claude Bertrand * Non-Executive Director General Director R&D - Servier



Jean Volatier ★ Non-Executive Director **CFO** - Inventiva



Dorothée Burkel * Non-Executive Director Former Chief Corporate and People Operations Officer - PartnerRe



Muriel Prudent Censor **VC Investment Manager – Fonds PSIM - Bpifrance**



Hervé Affagard **Executive Director MaaT Pharma**

Corporate Social Responsibility

MaaT Pharma aims to become the source of Microbiome excellence providing patients with safe and innovative medicines. The Company develops products from sustainable biological matters, driving optimal impact of Microbiome.

Patients are our priority. We are committed to our patients and to the protection of human health by respecting environmental protection, respecting our employees and ensuring good governance practices. Our way of working every day is driven by the 4 guidelines below:







- Innovate and raise awareness to deliver better care,
- Contribute to employees-growth within a people-oriented ecosystem,
- Place ethics and transparency at the core of the Company's strategy,
- Control and measure our impact on the environment.







2022 CSR indicators

SOCIAL

Gender Equality Index Employment of young people (under 30 and less than 5 years experience)

QWL: Job satisfaction

ENVIRONMENT

Carbon footprint

Energy consumption per employees on site

SOCIETAL

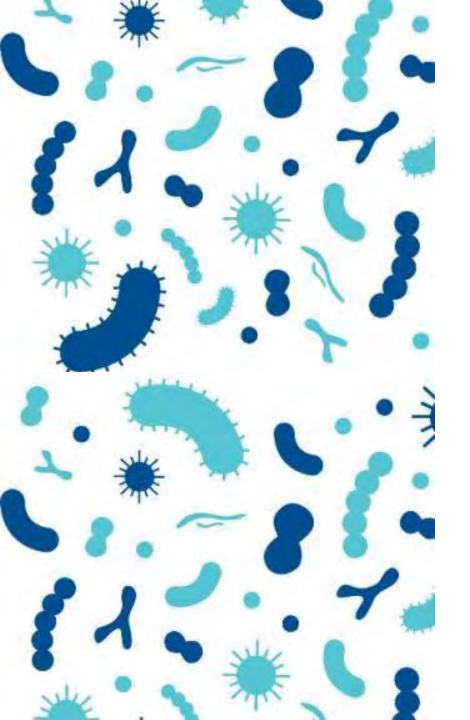
Responsibility to patients and practitioners
Increase awareness of Microbiome therapies
R&D at the focus of our

investments

GOVERNANCE

% of women in the board of directors and management team

% of women in the top 10 earners



THANK YOU

contact@maat-pharma.com



33 4 28 29 14 00