PHAX AM

PHAXAM

Building a Global Leader in Severe Infections Therapies

Agenda

Global Context
 PHAXIAM Differentiation
 Development Strategy
 Communication & Financing

Experienced & Complementary Leadership Team

Thibaut du Fayet CEO









Pascal Birman, MD

CMO



Cindy Fevre CSO Frédéric Mathat CFO



Pherecydes and Erytech merged to build **PHAX A**M

Leveraging on Complementary Capabilities from both Executive Teams

December 2024

PHAXAM

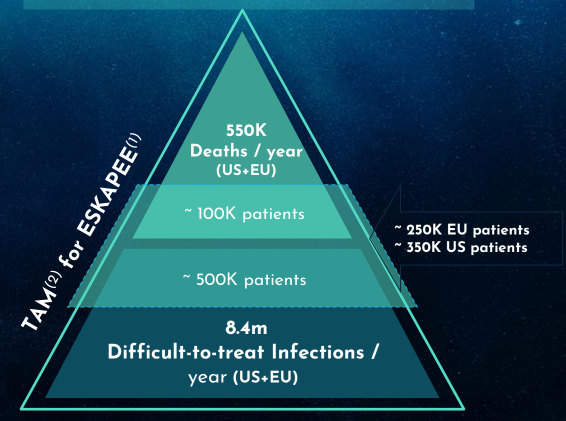
Phages Therapy: A Fast-emerging Large Targeted Addressable Market in EU / US

Significant Unmet Medical Needs

 Deaths from resistant bacterial INFECTIONS reaching in 2024 ~ 550K patients in EU + US: ~250K (EU) and ~300K⁽⁴⁾ (US) (Lancet⁽²⁾, sept 2024)

 Upwards of 8.4 million patients experience every year unresolved resistant bacterial/ difficult-to-treat INFECTIONS in high income countries: EU ~3.7m / US ~4.7m (Lancet⁽²⁾, sept 2024)

 Phages therapy applications will range from treating "difficult-to-treat" patients to "last-resort / death" patients
 → TAM⁽²⁾ in EU and US ~ 600K patients TAM⁽²⁾ Estimate for Phages ESKAPEE⁽¹⁾ therapy in EU+US \rightarrow ~ 600K patients / year



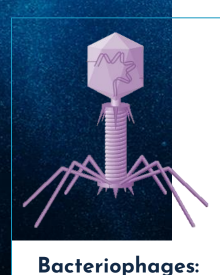
ESKAPEE = most important pathogens covering ~90% of severe resistant infections: E. coli, S. aureus, K. Pneumonia, A. baumannii, P. aeruginosa, E. faecium, Enterobacter
 Global burden of bacterial antimicrobial resistance 1990–2021: a systematic analysis with forecasts to 2050 (Lancet, Sept 2024)

December 2024

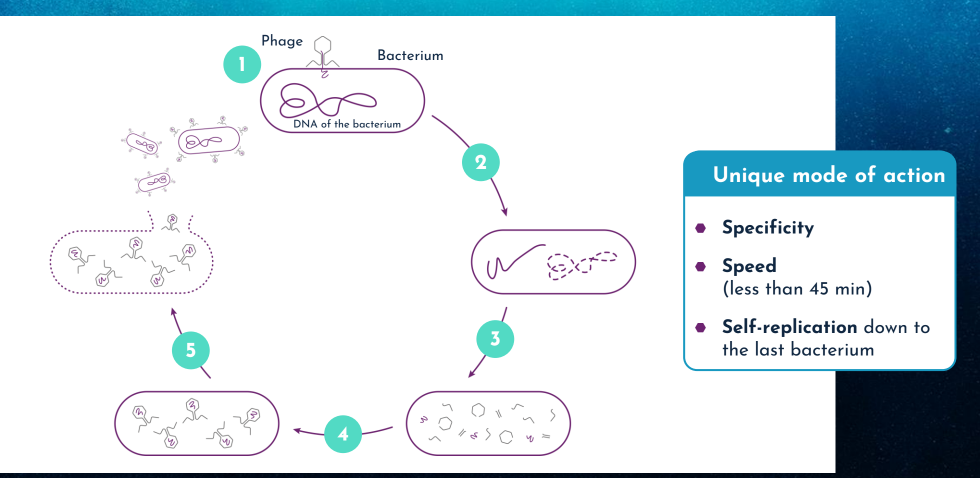
Targeted Addressable Market Estimated – internal estimation



Phage Therapy, a Solution for Resistant Infections



Bacteriophages: viruses, natural predators of bacteria



6

Phage Therapy allows SIMPLE, EFFECTIVE and WELL-TOLERATED treatments

PHAXIAM

The Strong Momentum of Phage Therapy

INCREASING IMPACT of difficult-to-treat resistant infections Rising concern among public authorities and medical community

High safety and promising clinical benefits from REAL-LIFE treatments Increased probability of success from early clinical evidence

Active collaboration with REGULATORY AGENCIES Strong support / Clear development guidelines / Accelerated paths

> GMP STANDARDIZATION & CMC developments Robust GMP processes / Well characterized phages

Better understanding of resistance mechanisms NEW TOOLS to address potential emergence of phage resistance



CRITICAL NEEDS FOR ALTERNATIVE TECHNOLOGIES

PROMISING REAL-LIFE CLINICAL DATA

ATTRACTIVE REGULATORY CONTEXT

MATURITY OF THE TECHNOLOGY

SOLUTIONS TO POTENTIAL RESISTANCE

Phage Therapy is a TOP-10 INNOVATION to be developed according to the 2023 World Economic Forum



2- PHAXIAM Differentiation

Leading Phage Therapy Platform

Leading Edge in Clinical Development Regular Interactions with Regulatory Agencies Strong internal R&D, CMC & GMP Capabilities Proprietary PHAGOGRAM IVD Solution Large Phage Bank for the most critical bacterial Infections Strong IP with 87 patents filed



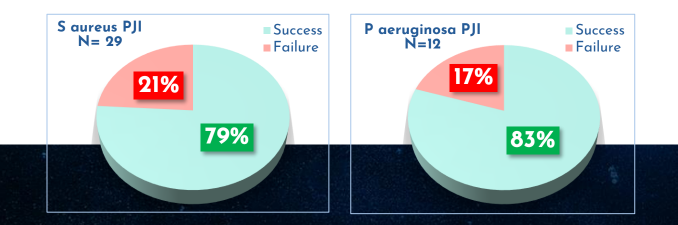
December 2024

PHAXIAM

2- PHAXIAM Differentiation

Real-life Clinical Data From ~120 Treated Patients

- STRONG SUPPORT from Regulatory Authorities & >15 hospitals
- SEVERAL ROUTES OF ADMINISTRATION TESTED, including local, intravenous, nebulisation, ...
- 7 DIFFERENT INDICATIONS TREATED with a majority of PJI



PROMISING CLINICAL ACTIVITY RESULTS

(First 77 patients evaluated) 2020-2024

- EXCELLENT RESULTS observed in reported cases: safety + clinical benefit
- Several PUBLICATIONS

~ 80% CONTROL RATE of infection @3months for PJI Patients (n=41) VERY PROMISING Data in « hard to treat » population (very severe infections - 2nd/3rd line antibiotics)

December 2024

ΡΗΑΧΙΑΜ

AAC* Regulatory Status from French Authority (June 2022)



SOURCE OF REVENUES

COMPASSIONATE ACCESS AUTHORIZATION (AAC) To be extended to OTHER AAC* INDICATIONS

Process towards an EARLY ACCESS AUTHORIZATION

To be extended to ORPHAN STATUS

An Important First Step for MARKET ACCESS in EUROPE

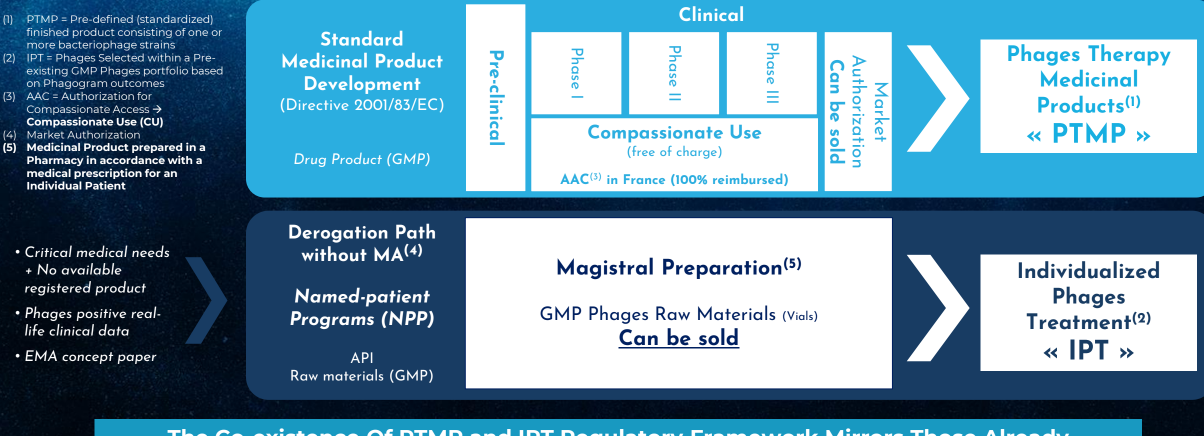
December 2024

PHAXIAM

*Compassionate Access Authorization

(5)

Strong Demand From Physicians Opens A Complementary Market Access Business Model



The Co-existence Of PTMP and IPT Regulatory Framework Mirrors Those Already Implemented in The Allergens Therapeutic Domain

December 2024

PHAXIAM



1- Phage Therapy Medicinal Product (PTMP⁽²⁾) Accelerated Regulatory Authorization 2- Individualized Phages Therapy (IPT⁽¹⁾) Magistral preparation / Compounding

Consequently, PHAXIAM seeks to position itself to:

- 1. Capitalise on its first mover advantage in the strategically-important emerging IPT model, where market access can start as soon as 2026,
- 2. Leverage on its existing capabilities in PTMP to secure Conditional Marketing Approval (CMA) by H2 2027.



3- Development Strategy PHAXIAM Has Strong Clinical Development and Market Access Capabilities in the PTMP⁽¹⁾ Model



Phages Therapy Medicinal Product (PTMP⁽¹⁾)

3 Main Pathogens Covered in Clinic

- S. Aureus → PJI Pilot Trial, GLORIA PJI Phase II, Endocarditis Phase I PK, PhagoPIED Phase II (IST)
- P. aeruginosa → PyoPhaneb Phase
- E. coli → Phase | PK to be initiated

- Strong push from FDA / EMA to seek robust clinical POC in RCTs + Based on a EU Pharmacopeia monograph (March 24)
- Recent FDA IND clearance for GLORIA Phase II
- Large hospitals & KOLs network in EU/US, leveraging on ongoing clinical trials: PhagoDAIR (15 EU sites), GLORIA (35 EU sites + 10 US sites)
- PHAXIAM AAC⁽²⁾ (Compassionate Use) Revenues to be extended to other indications

Clinical Strategy Value Drivers

- Relevant choice of Clinical indications (PJI) & clinical design to maximize Clinical POC for Phages
- Accelerated registration through CMA / Early Access in EU & US leveraging on Real-life compassionate clinical data

Pre-defined (standardized) finished product consisting of one or more bacteriophage strains
 AAC = Authorization for Compassionate Access -> Compassionate Use (CU)





14

An Ambitious Clinical Development Strategy

Target High-Value Indications

Severe Resistant Infections with High unmet medical needs high mortality rate / high budget impacts -> claims high prices

Accelerate the Path to Global Registration

Launch the 1st global randomized Phase 2 study at international scale in PJI* Leverage on potential Early access pathway (after Phase 2)

Diversify Portfolio

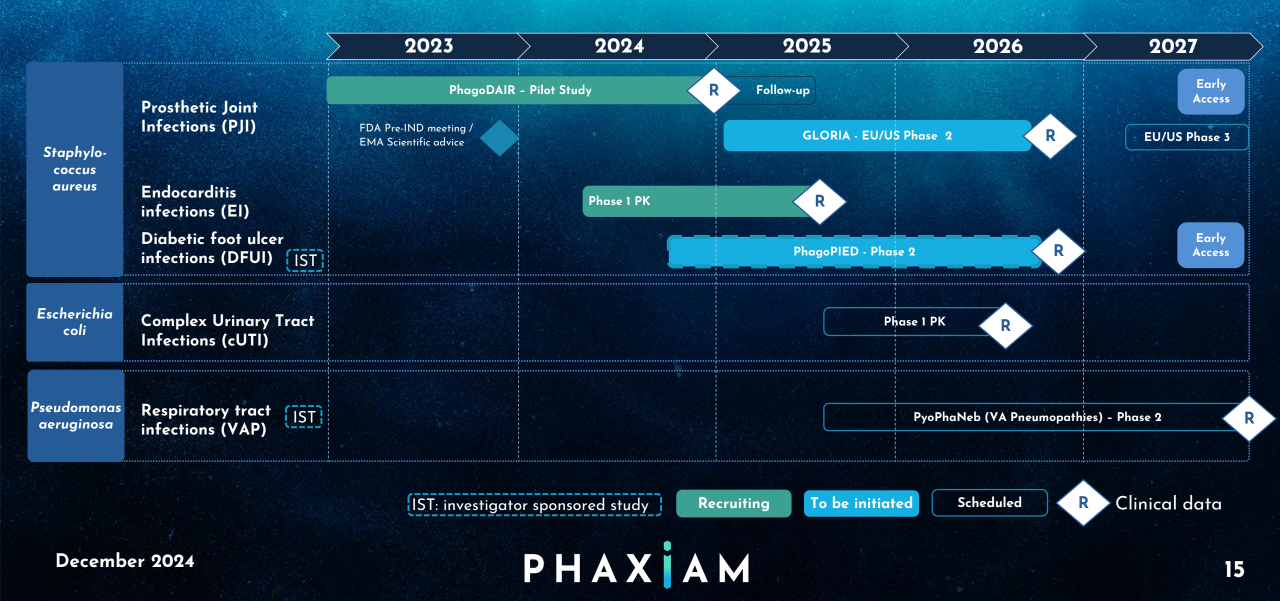
Target several Life-Threatening Infections (Endocarditis, VAP**, ...) Target several Virulent & Resistant Bacteria (E. coli, P. aeruginosa, ...)

PHAXIAM

* Prosthetic Joint Infections
 **Ventilated-acquired Pneumopathies

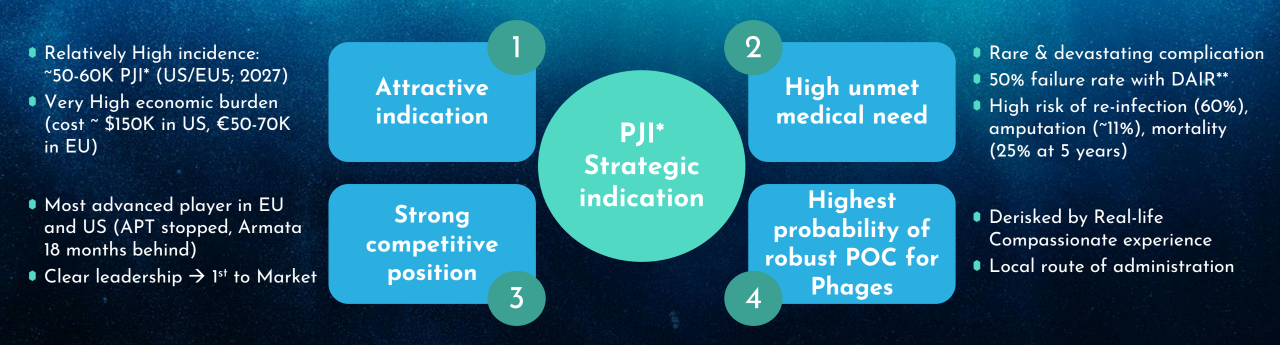
A Balanced Clinical Portfolio







PJI, A Strategic Indication For PHAXIAM



Unique and leadership position in the strategic PJI indication

December 2024

PHAXAM

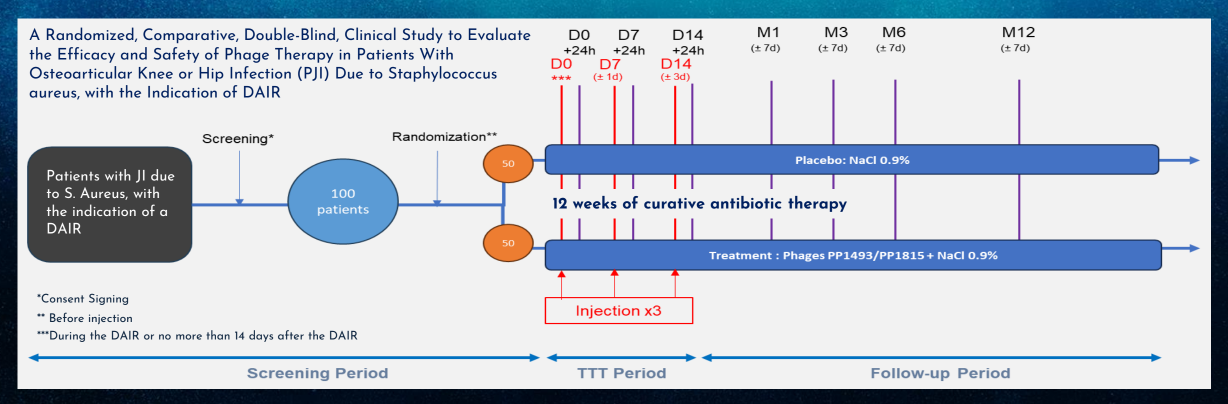
* Prosthetic Joint Infection (PJI)
** SOC = Debridement, Antibiotic, Implant Retention



* Proof of Concept

17

GLORIA -first global Phase 2 POC* clinical study of phages in PJI



PHAXIAM

Primary endpoint: Safety; Efficacy (% of patients with clinical cure at M3)
The GLORIA study is the most strategic clinical trial for PHAXIAM

Septembre 2024



Preparation of The GLORIA Study

Positive and consistent feedback from the FDA (pre-IND meeting) and EMA (scientific advice)

- Sites selection ongoing: ~45 sites and countries (FR, ESP, GER, NLD, ITA, UK, US)
- Launch of the clinical study in Q1 2025, clinical results expected in Q3 2026

- Progress in line with our objectives
- GLORIA Is Our Most Important Asset, Having The Highest Priority



Expected Major Short term Clinical Catalysts



December 2024

PHAXAM

PTMP

Near Term Commercial Opportunities Arise From the Emergence of the Individualized Phages Therapy (IPT⁽¹⁾) Market Access Model

IPT

Strong Demand from Physicians

«Push» from Regulatory Stakeholders (EMA) & National Bodies

High unsatisfied clinical needs → new solutions to address morbidity / mortality induced by severe resistant infections, when the PTMP⁽²⁾ model cannot address clinical needs, as no Medical Products are available

Promising clinical evidence from hundreds of successful compassionate treatments in key reference hospitals across EU

EMA Concept paper Guidelines

- → Regulatory landscape evolving, making a special case on bacteriophages "Named-Patient Use Program" (NPP)
- Phages Therapy Medicinal Product: predefined (standardised) finished product consisting of one or more bacteriophage strains→ Major requirements for wellconducted RCTs
- **Individualized Phages Therapy (IPT)** → Delineate context of clinical use where Medical Products cannot be used / are not available

Individualized Phages Therapy (IPT)

- Magistral Preparation / Compounding with <u>GMP</u> Phages
- Pre-requisite for Regulatory
 Validation per country (e.g. Exemption granted in Belgium)
- **Based on a Diagnostic Test** (Phagogram - CE marked)
- Selection of Phages drawn from a pre-existing GMP Phages portfolio

- IPT model = Phages Selected within a Pre-existing GMP Phages portfolio based on Phagogram outcomes
- (2) Pre-defined (standardized) finished product consisting of one or more bacteriophage strains

ΡΗΑΧΙΑΜ

For IPT⁽¹⁾, PHAXIAM Is Targeting As A 1st Priority European Countries Familiar with Magistral Preparation



21

UK

Strong interest from Physicians for Phages Magistral preparation concept / Compounding very much developed

Regulatory constraint today → to be discussed and validated by MHRA for general use Solution funded by national / regional / hospital

FRANCE

envelops

- Strong interest from Physicians for Phages Magistral preparations less used than before / reduced number of compounding organisations
- AAC process in place as Compassionate Use

SPAIN

- Strong interest from Physicians for Phages
- Magistral preparations progressively disappeared / not a common practice

NORDIC

Strong interest from Physicians for Phages Magistral preparation / Compounding common practice



BELGIUM

 Strong interest from Physicians for Phages

Magistral preparation common practice benefiting from an exemption

NETHERLANDS

Strong interest from Physicians for Phages Magistral preparation / Compounding developed

GERMANY

Strong interest from Physicians for Phages Magistral preparation/compounding feasible and standard practices

ITALY

(1) IPT model = Phages Selected within a Pre-existing GMP

Phages portfolio based on Phagogram outcomes Pre-defined (standardized) finished product consisting of

one or more bacteriophage strains

Physicians want to use Phages Magistral preparation commonly used / large compounding community existing

Increasing Interest From Physicians For Phages All Over Europe Magistral Preparation / Compounding Are Common Practices In Northern Europe

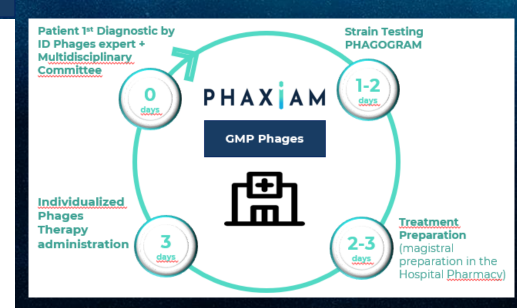
AM

PHAX

3- Development Strategy PHAXIAM Will Deliver IPT to Patients in Key Reference Hospitals As Soon As 2026

Actions Plan

- Obtain Local regulatory approval (national / regional) for magistral preparation (leveraging on existing exemption)
- Develop the EU network within reference hospitals (physicians, pharmacists) to be extended to smaller hospitals
- Set-up locally Supply chain and Diagnostic capabilities → Phagogram to be transfered



PHAXIAM

Value Drivers

IPT

22

- Average Selling Price of ~€20-25K (EU)
- Treatment invoiced at a local/hospital level, Regional level in some countries
- HTA/Payor negotiation in a 2nd step leveraging on KOL's concerted efforts to support creation of reimbursement codes, at local / national levels

3- Development Strategy PTMP And IPT Models Are Synergistic Market Channels With Largely Overlapping Requirements



Paneuropean Specialty Pharma in Critical Care

Individualized Phages Therapy (IPT)

- TAM⁽¹⁾ in EU / US ~ 600K patients, of which
 ~250 K patients in EU
- A Fast-emerging Large Market of > ~ €10bn (EU+US)
- Conservative Average Selling Price of ~€20-25K (EU)
- Not subject to any clinical development risk
- EU market access Scale-up to reach
 - ~1K pts / 30 hospitals in 2027; Potential Sales →
 €20m
 - ~7-8K pts / 100 hospitals in 2030; Potential Sales
 → ~€150m
- Conservative 2030 Commercial Target is limited to ~3% of TAM⁽¹⁾

Fully integrated platform including R&D, GMP manufacturing

Strong relationships with KOLs & Physicians

Presence in major reference hospitals in EU

Strong relationships with Regulatory authorities across EU / US

Diagnostic robust solution in place & potential future reference method

Market Access capabilities

Phage Therapy Medicinal Product (PTMP)

AAC (Compassionate Use) Potential Sales → ~
 €4m in 2027 (PJI + other indications)

• PJI (lead clinical indication)

- TAM(1) in EU/US ~ 40K pts / year
- Average Selling Price of ~ €20-25K (EU) and
 ~€25-30K (US)
- Seeking Conditional Market Approval (CMA)
 / Early access pathways in H2 2027
- PJI CMA Potential Sales → ~ €8m in 2027;
 €90-100m in 2030

Synergies Are Reducing The Investments Requirements To Scale-up The IPT Model

December 2024

ΡΗΑΧΙΑΜ

4. Communication & Financing Emerging Paneuropean Specialty Pharma in Critical Care Targeting Positive FCF⁽¹⁾ AND Profitability From 2027



Paneuropean Specialty Pharma in Critical Care

Individualized Phages Therapy (IPT)

- TAM⁽¹⁾ in EU / US ~ 600K patients, of which
 ~250 K patients in EU
- A Fast-emerging Large Market of > ~ €10bn (EU+US)
- Conservative Average Selling Price of ~€20-25K (EU)
- Not subject to any clinical development risk
- EU market access Scale-up to reach
 - ~1K pts / 30 hospitals in 2027; Potential Sales →
 €20m
 - ~7-8K pts / 100 hospitals in 2030; Potential Sales
 → ~€150m
- Conservative 2030 Commercial Target is limited to ~3% of TAM⁽¹⁾

Fully integrated platform including R&D, GMP manufacturing

Strong relationships with KOLs & Physicians

Presence in major reference hospitals in EU

Strong relationships with Regulatory authorities across EU / US

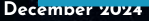
Diagnostic robust solution in place & potential future reference method Phage Therapy Medicinal Product (PTMP)

- AAC (Compassionate Use) Potential Sales → ~
 €4m in 2027 (PJI + other indications)
- PJI (lead clinical indication)
- TAM(1) in EU/US ~ 40K pts / year
- Average Selling Price of ~ €20-25K (EU) and
 ~€25-30K (US)
- Seeking Conditional Market Approval (CMA)
 / Early access pathways in H2 2027
- PJI CMA Potential Sales → ~ €8m in 2027;
 €90-100m in 2030

PHAXIAM'S Market Access Strategy Is To Rely On Two Concurrent Commercial Pillars: IPT (magistral preparation) and PTMP (ACC + PJI CAM)

PHAXAM

Market Access capabilities



(1) Free Cash Flow

4- Communication & Financing

PHAXIAM, An Emerging Specialty Pharma In Critical Care

IPT_

PTMP

- Millions of patients experience every year unresolved resistant bacterial/ difficult-to-treat infections.
- Facing this critical medical need, there is strong demand from KOLs /Physicians for Phages therapy.
- Concurrently with its classical clinical development pathway (Phages Therapy Medical Product=PTMP), EMA has
 opened the market for the commercialization of Individualized Phages Therapy (IPT) magistral preparations.
- PTMP and IPT Models are complementary and synergistic approaches that can be managed by PHAXIAM with limited additional resources.
- PHAXIAM will benefits greatly from these two complementary commercial pillars:
 - IPT with very large potential that will secure short term revenues (2026-2030),
 - PTMP with already existing AAC Revenues, then potential conditional market approval (CMA) from H2 2027.
- PTMP and IPT form together a self-reinforcing virtuous cycle for Phages therapy:
 - Commercial success in IPT will help finance PTMP programs and facilitate rapid uptake of future approved medicinal products,
 - Success (clinical validation) in PTMP will help broaden clinical and market acceptance of IPT in areas not yet addressable by PTMP.
 - This is strategically important because PHAXIAM is the only competitor with the capabilities to succeed in both markets

PHAXIAM targets positive Free Cash Flow and Profitability from 2027

PHAXAM

4- Communication & Financing

Financial Position & Objectives

- Major shareholders (equity share): BPI (25%), Auriga / Elaia (8%), Go Capital (5%)
- Listed @Euronext: valuation of €18m
- Capital increase of €7.8m in June 2024
- Cash Runway into March 2025
- Financing
 - Non-dilutive funding (EU, France, Region)
 - Dilutive funding: Private placement in January 2025, targeting €15m; use of proceeds: GLORIA study conduct (1st global EU/US study in PJI* and Phage therapy) + other clinical studies

Till The Market PHAXIAM Still Needs €15-20m To Be Raised In Equity

PHAXIAM

*Prosthetic Joint Infections

Thanks PHAXIAM

Reinforced interest in Phage Therapy Major Commitment from Top Tier US investors in March 2024



Purchase agreement with a \$50m financing led by Top tier investor base, incl. Deerfield and Orbimed



Armata Pharmaceuticals has recently announced a \$35m refinancing with Innoviva

Growing attractiveness of the phage-therapy field to leading investors

December 2024

PHAXAM

Key Technology Assets for PHAXIAM Treatments

SELECT phages to maximize breadth of repertoire Internal PHAXIAM Technology and Expertise

PRODUCE large scale GMP batches of high purity Internal PHAXIAM process development Capabilities and Expertise

Industrial partnership to produce "off-the-shelf" GMP-grade phages

TEST PHAGOGRAM for a precision therapy PHAXIAM proprietary IVD Test

DISTRIBUTE personalized therapeutics to patients' bed Supply chain in place with a few days leadtime

Discovery, Screening, Characterization, GMP production, Testing, Distributing

3









Manufacturing & Logistics strategic capabilities

Major achievements

Request the active Phage
 From GMP Phage Bank

30



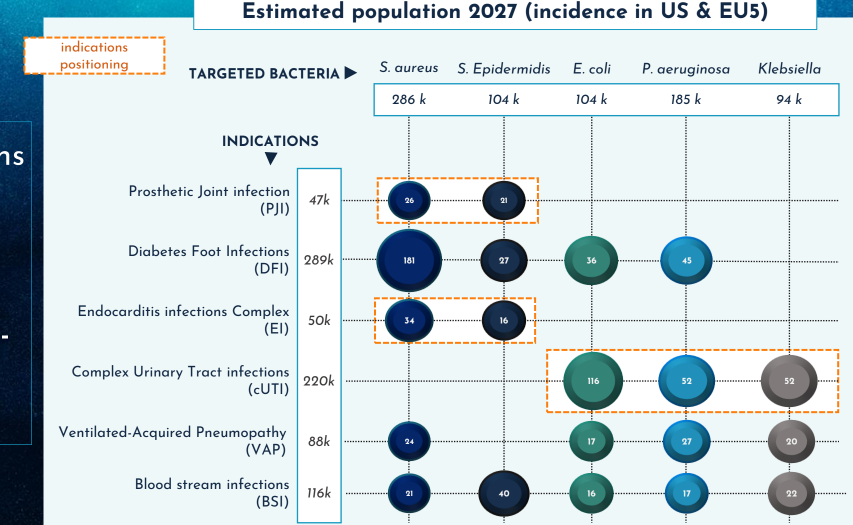
Manufacturing & Logistics Capacities Fit to Address future Clinical Demand

PHAXIAM

Appendixes: Development Strategy

Targeting High-value Resistant Infections

PHAXIAM



31

Life-Threatening Conditions

Potential Orphan Status Indications

The most severe Hospital-Acquired and Resistant Infections

Appendixes: Development Strategy

Additional Clinical Studies

TRIALS	STATUS AND PROGRESS
Endocarditis Infections (EI) Staph. aureus Phase I PK	 Demonstration for IV indications before Registration Study Resistant infections in cardiac chambers and valves IV-administered Phages <u>Key milestones</u> : First Patient-In, April 2024
Complex Urinary Tract Infections (cUTI) E. coli Phase I PK	 Demonstration for intra-bladder administration before Registration Study cUTI with resistant E. Coli infections in the bladder Phages administered locally into the bladder <u>Key milestones</u> : ANSM study validation in April 2024

PHAXIAM