



Immunic
THERAPEUTICS

Immunic Therapeutics

First Quarter 2024 Financial Results and Corporate Update

NASDAQ: IMUX | May 8, 2024

Cautionary Note Regarding Forward-Looking Statements

→ This presentation contains “forward-looking statements” that involve substantial risks and uncertainties for purposes of the safe harbor within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. These include statements regarding management’s intentions, plans, beliefs, expectations or forecasts for the future, and, therefore, you are cautioned not to place undue reliance on them. No forward-looking statement can be guaranteed, and actual results may differ materially from those projected. Immunic undertakes no obligation to publicly update any forward-looking statement, whether as a result of new information, future events or otherwise, except to the extent required by law. We use words such as “anticipates,” “believes,” “plans,” “expects,” “projects,” “future,” “intends,” “may,” “will,” “should,” “could,” “estimates,” “predicts,” “potential,” “continue,” “guidance,” and similar expressions to identify these forward-looking statements that are intended to be covered by the safe-harbor provisions of the Private Securities Litigation Reform Act of 1995.

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→ Forward-looking statements included in this presentation are based on information available to Immunic as of the date of this presentation. Immunic does not undertake any obligation to update such forward-looking statements except as required by applicable law.



Agenda

First Quarter 2024 Financial Results and Corporate Update

01 First Quarter 2024 and Subsequent Highlights

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01

First Quarter 2024 and Subsequent Highlights

January: Three-Tranche Private Placement of up to \$240M, Cash Runway Extended Into Q3/2025 Based on Initial \$80M Tranche

Private Investment in Public Equity (“PIPE”) financing

- **First tranche** was an upfront payment of **\$80 million** at \$1.43 per share
- **Second tranche** is a conditional mandatory purchase of an **additional \$80 million** at \$1.716 per share
 - Representing 120% of the first tranche purchase price
 - Conditioned on the announcement of phase 2b top-line data for the CALLIPER trial of vidofludimus calcium in PMS, volume weighted average share price levels, and minimum trading volumes
- **Third tranche** provides for the issuance of **\$80 million** of shares at the same price per share as the second tranche
 - To occur no later than three years after the second tranche
 - Permits investors to fund their purchase obligations on a “cashless” or net settlement basis
 - Conditioned on the same volume weighted average share price levels and minimum trading volumes as the second tranche
- Any of the conditions in the second or third tranches can be waived by holders of a majority of the outstanding securities, including the lead investor

Total Gross Proceeds

- **Up to \$240 million**

Participating Investors

- Led by BVF Partners
- Includes participation from **new and existing investors**, including Avidity Partners, Janus Henderson Investors, Soleus Capital, RTW Investments and Adage Capital Partners

Closing Date

- January 8, 2024 for initial \$80 million tranche

Lead Placement Agent / Placement Agent / Capital Markets Advisors

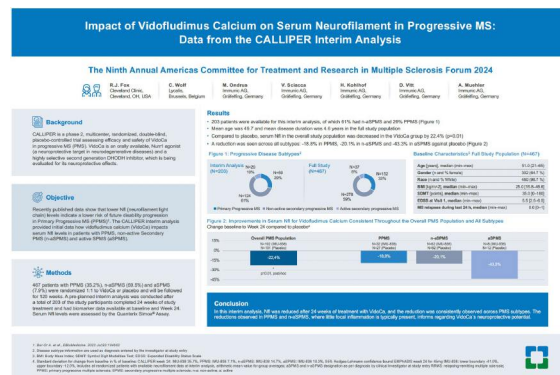
- Leerink Partners / Ladenburg Thalmann / Piper Sandler, B. Riley Securities, Brookline Capital Markets

February: Presented Data From Phase 2 CALLIPER and CALVID-1 Trials of Vidofludimus Calcium at the ACTRIMS Forum 2024 (February 29-March 2 in West Palm Beach, FL)



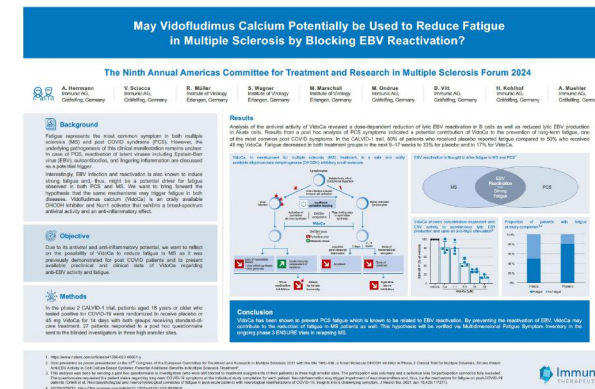
CALLIPER Interim Analysis: Clear Separation in NfL Levels Across All PMS Patients

- Oral Presentation: Robert J. Fox, MD, Staff Neurologist, Mellen Center for Multiple Sclerosis, Vice-Chair for Research, Neurological Institute, Cleveland Clinic, Cleveland, Ohio
- Title: Impact of Vidofludimus Calcium on Serum Neurofilament in Progressive MS: Data from the CALLIPER Interim Analysis

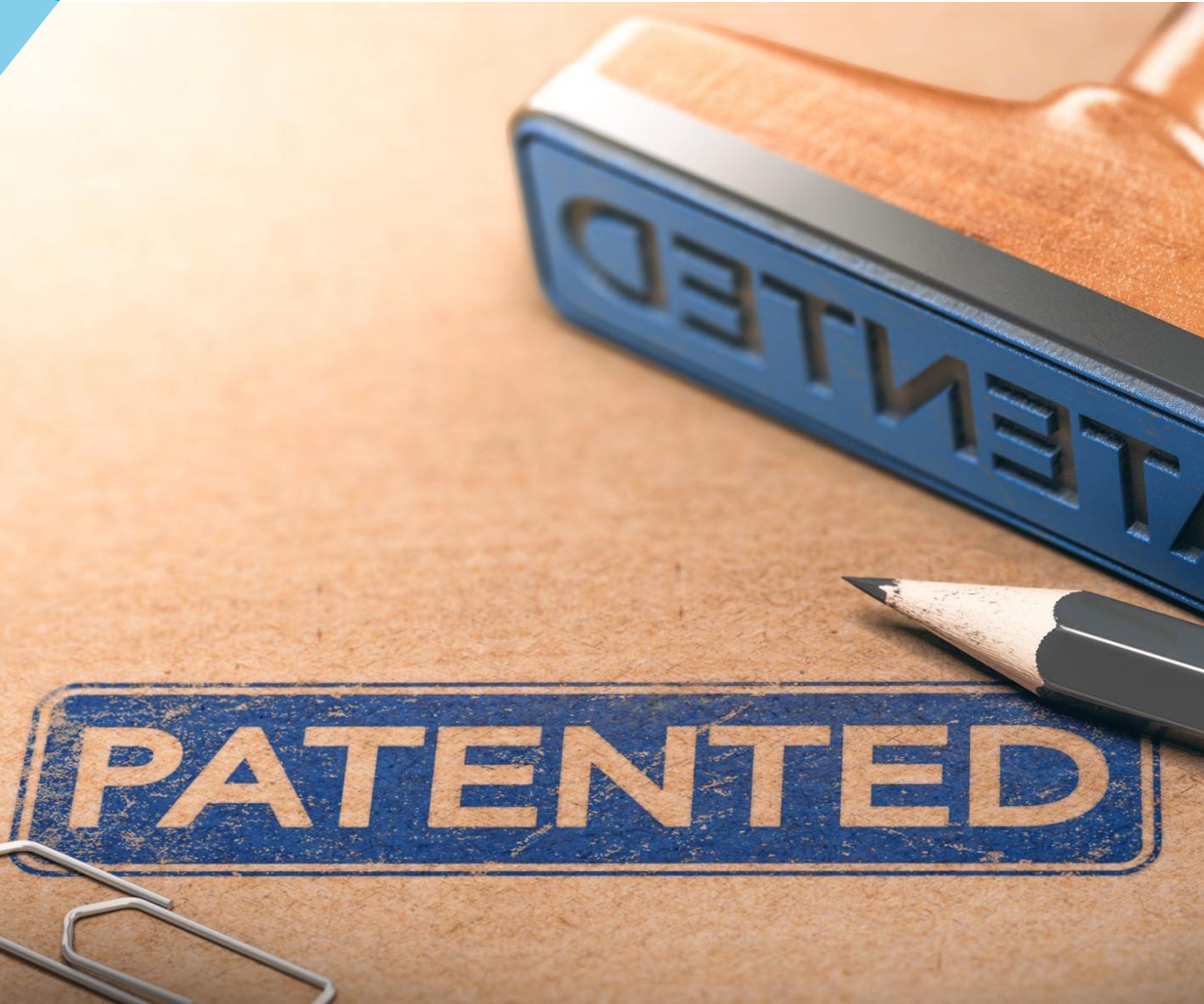


CALVID-1: Potential Contribution to the Reduction of Fatigue in MS Patients

- Oral Presentation: Dr. Alexandra Herrmann, Manager Translational Pharmacology, Immunic
- Title: May Vidofludimus Calcium Potentially be Used to Reduce Fatigue in Multiple Sclerosis by Blocking EBV Reactivation?



March: Received Fourth U.S. Patent Directed to Use of Vidofludimus Calcium in MS



Notice of Allowance from the USPTO for patent application 16/981,122, covering the composition-of-matter of a specific polymorph of vidofludimus calcium and a related method of production of the material



Claims are expected to provide protection into 2039 internationally, unless extended further; patent previously granted in Australia, Canada, Indonesia, Japan and Mexico



Multi-layered intellectual property strategy for vidofludimus calcium provides protection into 2041 in the US



April: Hosted In-Person Multiple Sclerosis R&D Day in San Francisco



Could Vidofludimus Calcium be the First Neuroprotective Treatment Option for Multiple Sclerosis?

Immunic speakers:

- Daniel Vitt, PhD, CEO & President
- Hella Kohlhof, PhD, CSO
- Andreas Muehler, MD, CMO

Attending expert:

- Zuoming Sun, Ph.D., Professor, Department of Molecular Imaging & Therapy City of Hope, Duarte, CA

Recording: <https://www.youtube.com/watch?v=pmrwoTVxEZo>



02

Financial and Operating Results

Condensed Consolidated Statements of Operations

(In thousands, except share and per share amounts, unaudited)

| | Three Months Ended March 31, | |
|--|------------------------------|--------------------|
| | 2024 | 2023 |
| Operating expenses: | | |
| Research and development | \$ 18,736 | \$ 22,963 |
| General and administrative | 5,145 | 4,288 |
| Total operating expenses | 23,881 | 27,251 |
| Loss from operations | (23,881) | (27,251) |
| Other income (expense): | | |
| Interest income | 1,187 | 800 |
| Change in fair value of the tranche rights | (4,796) | — |
| Other income (expense), net | (2,094) | 1,179 |
| Total other income (expense) | (5,703) | 1,979 |
| Net loss | \$ (29,584) | \$ (25,272) |
| | | |
| Net loss per share, basic and diluted | \$ (0.30) | \$ (0.58) |
| | | |
| Weighted-average common shares outstanding, basic and diluted | 97,299,955 | 43,664,783 |

\$97.3 million in cash and cash equivalents as of March 31, 2024 expected to fund operations into Q3/2025



03

Clinical Development Programs

Several Clinical Value Inflection Points Ahead



IMU-838 in PMS

- Top-line data from phase 2 CALLIPER trial expected in April 2025

IMU-838 in RMS

- Interim, non-binding futility analysis of phase 3 ENSURE program expected in late 2024
- Readout of first phase 3 ENSURE trial anticipated in Q2/2026, second in H2/2026

IMU-856

- Phase 2 clinical trial in preparation
- Potentially applicable to a multitude of gastrointestinal disorders



04

Q&A Session



05

Summary and Highlights

Advanced Clinical Pipeline

Well Differentiated Programs in Various Phases of Clinical Development

| Program | Preclinical | Phase 1 | Phase 2 | Phase 3 |
|--------------------------------|---|---------|---------|---------|
| Vidofludimus Calcium (IMU-838) | Relapsing Multiple Sclerosis (RMS) – ENSURE Trials | | | |
| | Progressive Multiple Sclerosis (PMS) – CALLIPER Trial | | | |
| | Ulcerative Colitis (UC) – CALDOSE-1 Trial | | | |
| IMU-856 | Celiac Disease | | | |
| | | | | |
| IMU-381 | Gastrointestinal Diseases | | | |
| | | | | |

■ Completed or ongoing ■ In preparation or planned

Thank You!



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