

**medincell.**

# Shift to Growth

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April 2026

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This presentation and the Fiscal Year 2024-25 results conference (French and English sessions) contain forward-looking statements, including statements regarding Company's expectations for (i) the timing, progress and outcome of its clinical trials; (ii) the clinical benefits and competitive positioning of its product candidates; (iii) the ability of its products to obtain regulatory approvals, commence commercial production and achieve market penetration and sales; (iv) its future product portfolio; (v) its future partnering arrangements; (vi) its future capital needs, capital expenditure plans and ability to obtain funding; and (vii) prospective financial matters regarding our business. Although the Company believes that its expectations are based on reasonable assumptions, any statements other than statements of historical facts that may be contained in this This presentation and the Fiscal Year 2024-25 results conference (French and English sessions) relating to future events are forward-looking statements and subject to change without notice, factors beyond the Company's control and the Company's financial capabilities.

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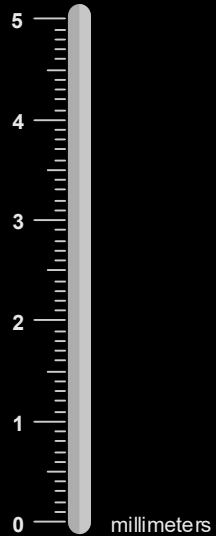
ZEDY® is a trademark of Teva Pharmaceuticals.

**The next two years will be the most  
transformative of Medincell's history**

Medincell - June 2025

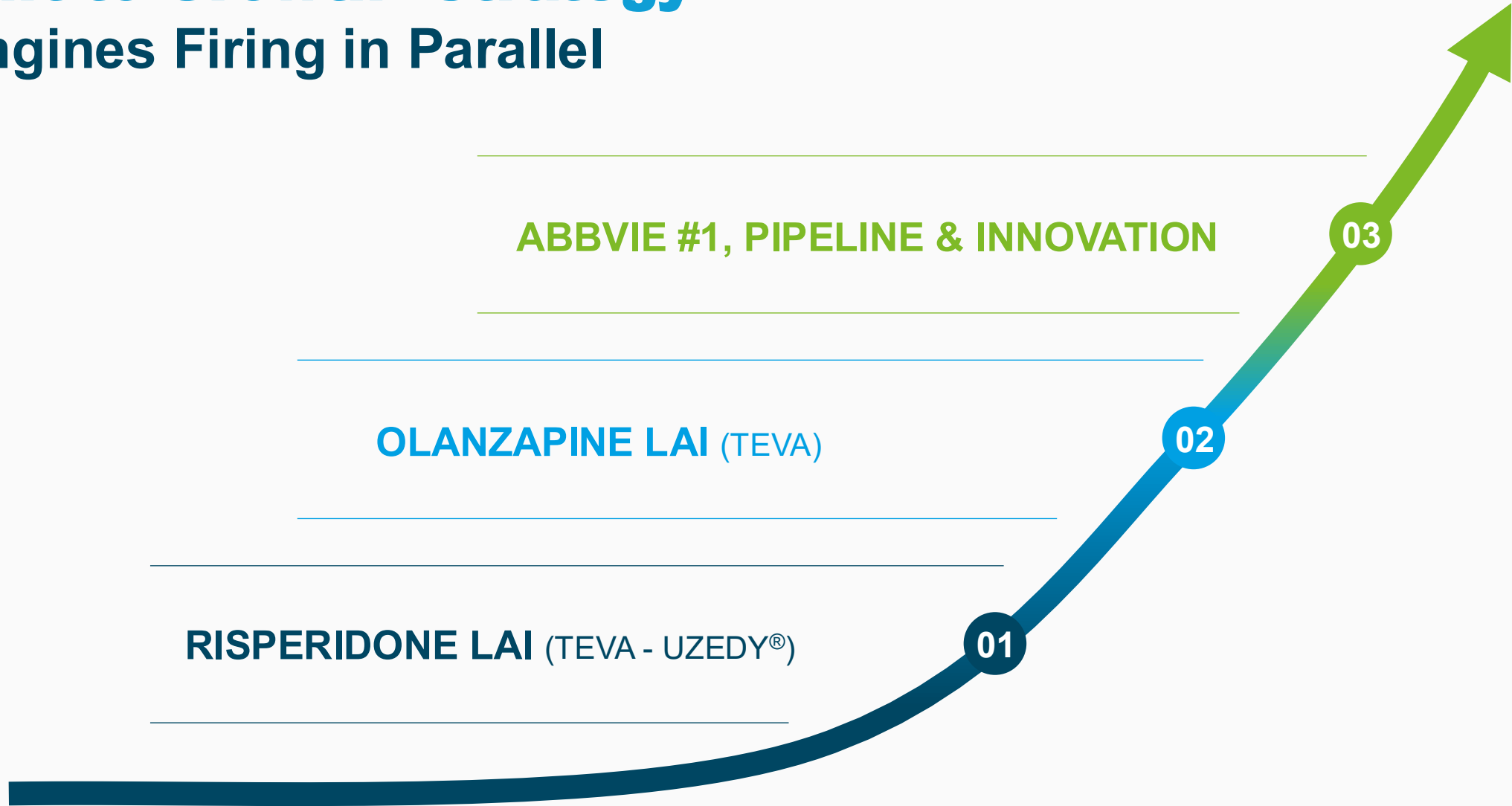
# BEPO®

Cutting-edge  
Long-Acting Injectable (LAI)  
Technology



# “Shift to Growth” Strategy

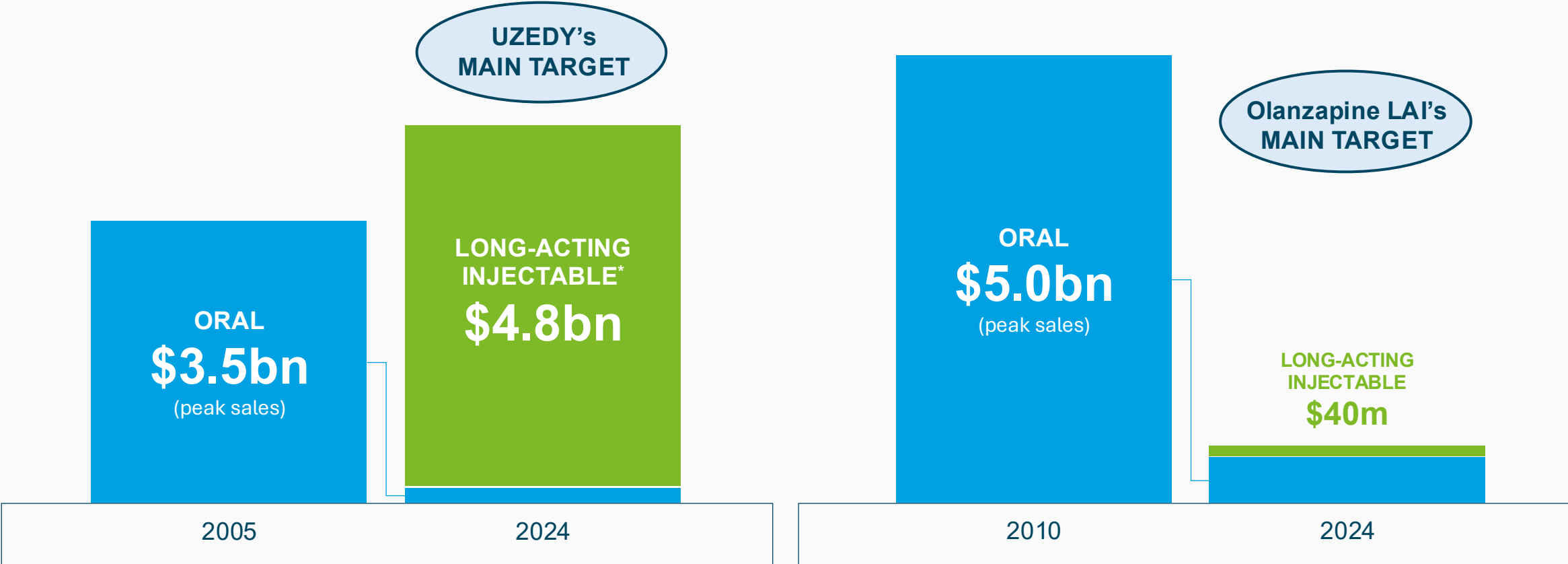
## 3 Engines Firing in Parallel



# Schizophrenia Strategy

**RISPERIDONE**  
Johnson & Johnson

**OLANZAPINE**  
Eli Lilly



Sources: 7 Major Markets - Companies reported sales, IQVIA  
\* Risperidone and its metabolite paliperidone

# Very Strong Momentum in Schizophrenia LAI



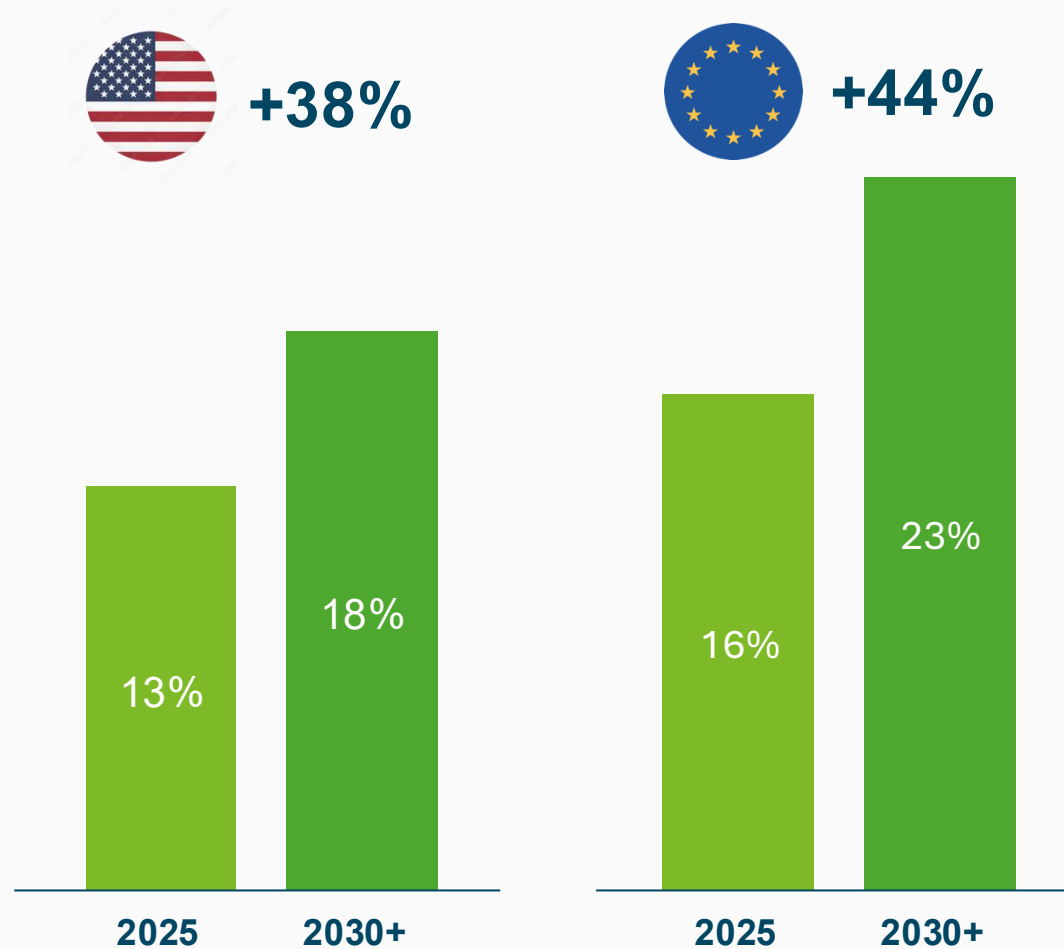
+38%



+44%

## LAI penetration in total treated schizophrenia population, in %

Source: LAI penetration based on IQVIA sales in Month of Therapy volume, with MIDAS (sales) dataset for EU and NDA Trx dataset for US / Teva's Innovation Strategy Day, May 2025



# **Teva is an Ideal Partner**

**UZEDY<sup>®</sup> and Olanzapine LAI are key assets in Teva's  
*Pivot to Growth* strategy**

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**Teva's psychiatry sales force has shown very strong recent  
commercial performance in the psychiatry field**

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**Worldwide reach**

# RISPERIDONE LAI (UZEDY<sup>®</sup>)

**Once-monthly subcutaneous injection of olanzapine**

**Approved for the maintenance treatment of Schizophrenia in adults**

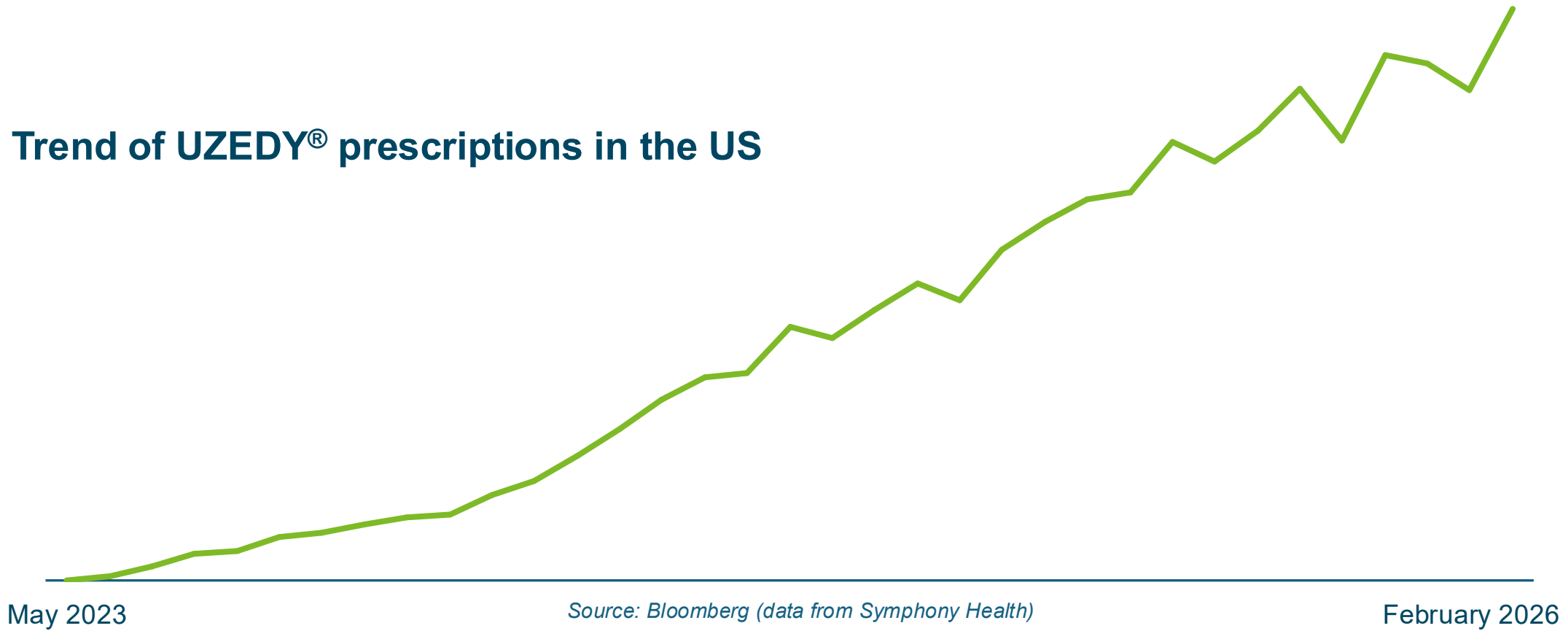
- 2023 United States
- 2025 South Korea (유제디), Canada (LONGAVO<sup>®</sup>)

**Approved for the maintenance treatment of bipolar I disorder in adults**

- 2025 United States

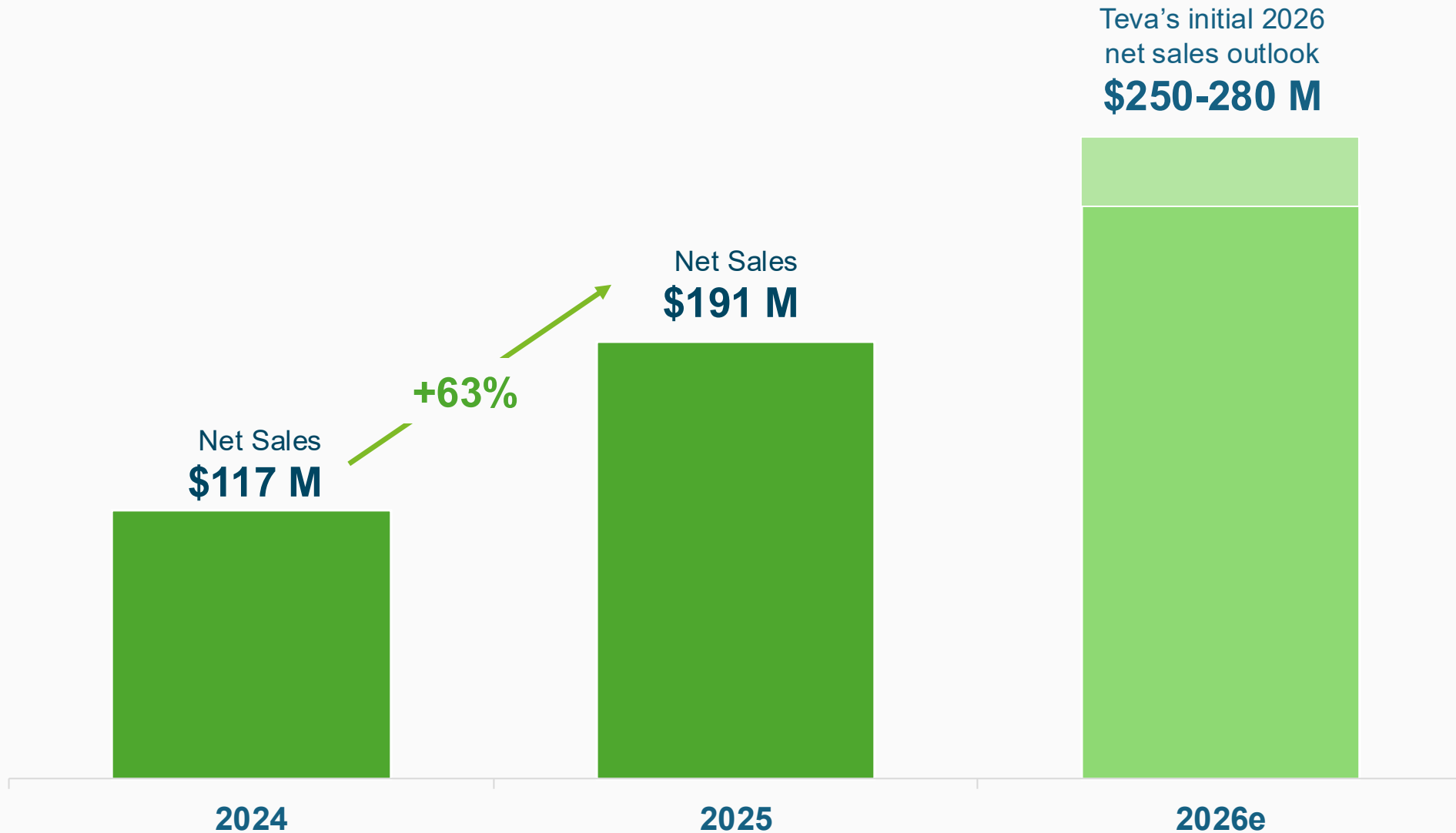
# UZEDY® - Already >15,000 Patients

Trend of UZEDY® prescriptions in the US

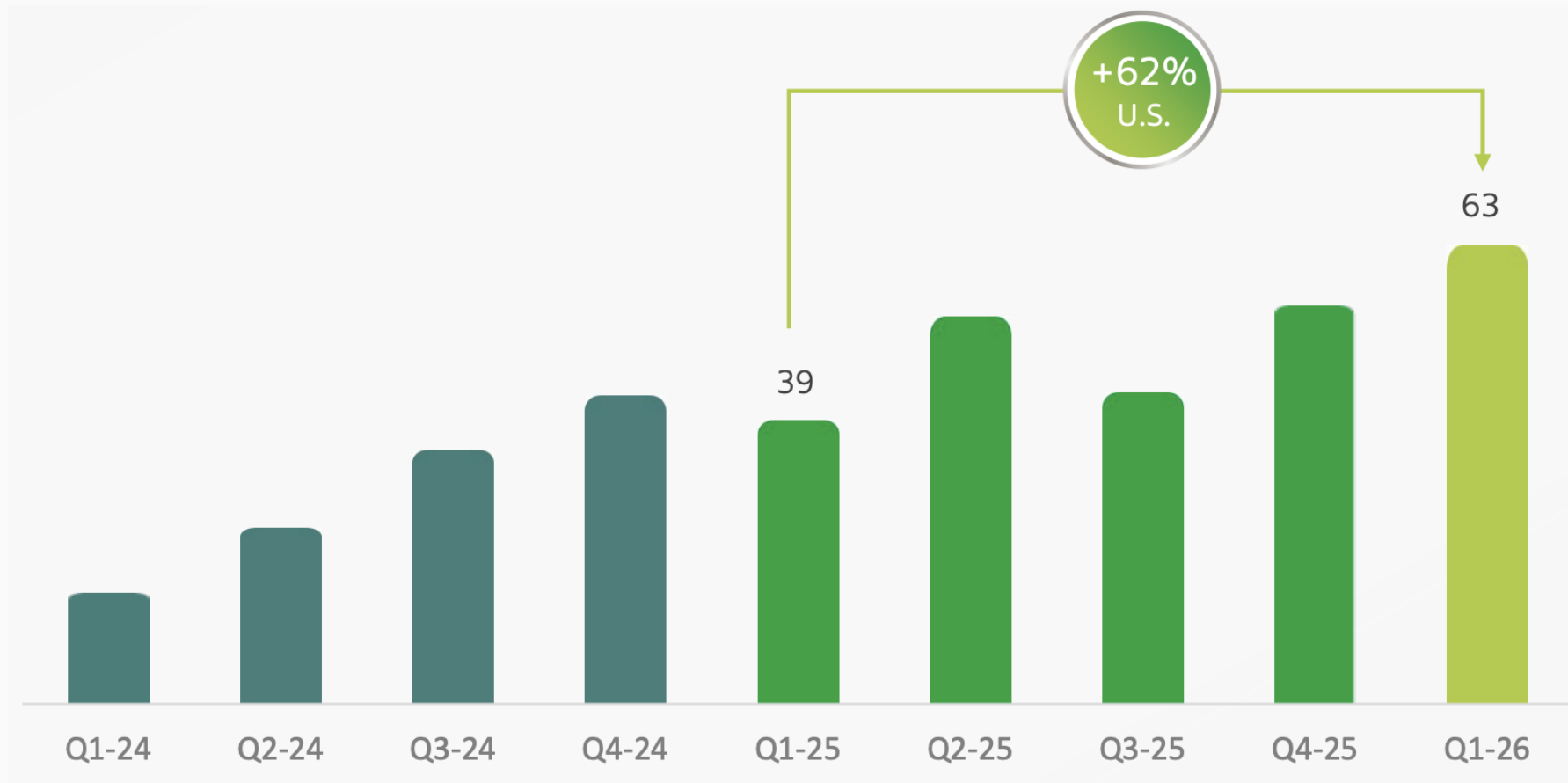


Source: Bloomberg (data from Symphony Health)

# UZEDY® - 2025 Net Sales Growth: +63%



# UZEDY<sup>®</sup> - Q1 2026 Net Sales Growth: +62%



Extract from Teva's Q1'26 Results Presentation - April 29, 2026

# UZEDY®: Analysts' Peak Sales Forecasts<sup>1</sup>

**Analysts' Consensus: \$1.2B**

**Low: \$0.6B - High: \$2.2B**















## Revenue potential for MedinCell

<p><b>Royalties</b></p>	<p><b>Mid- to high-single digit on all net sales</b></p>
<p><b>Milestones</b></p>	<p><b>Commercial: \$105 M</b></p>

<sup>1</sup>Based on most recent information received from analysts: Evercore, Truist, HC Wainwright, Jefferies, Stifel, Oddo BHF, Portzamparc, Kepler Cheuvreux, TP-icap. Any opinions, estimates, or forecasts regarding MedinCell's performance made by analysts are theirs alone and do not represent opinions, forecasts, or predictions of MedinCell or its management.

# UZEDY®: Medincell's Technology Enables Best-in-Class Features



Subcutaneous vs intramuscular

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No loading dose or oral supplementation\*

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Ready-to-use

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Same effect regardless of location of injection

\*Therapeutic levels are reached within 6 to 24 hours after administration

# UZEDY® Shows Strong Real-World Benefits

## UZEDY® vs second-generation oral antipsychotics (SGOAs) – May 2025

### RELAPSE RATE: -42%



### INPATIENT RATE: -47%



### MEAN TIME TO RELAPSE: +54%



### MEAN LENGTH OF HOSPITAL STAY: -50%



### MEAN ALL-CAUSE HCRU COSTS: -29%



Source: Poster presented by Teva Pharmaceuticals at the Annual Psych Congress Elevate, May 28–31, 2025, Las Vegas, NV: Treatment Patterns and Healthcare Resource Utilization Among Patients Receiving the Long-Acting Injectable Antipsychotic TV-46000 Versus Second-Generation Oral Antipsychotics. HCRU: healthcare resource utilization

# Multiple Patents Granted on UZEDY®

**RISPERIDONE LAI (UZEDY®)**  
US patent protection

**2042**

# OLANZAPINE LAI

## Once-monthly subcutaneous injection of olanzapine

**December 9, 2025**

Submission to U.S. FDA for the maintenance treatment of Schizophrenia in adults

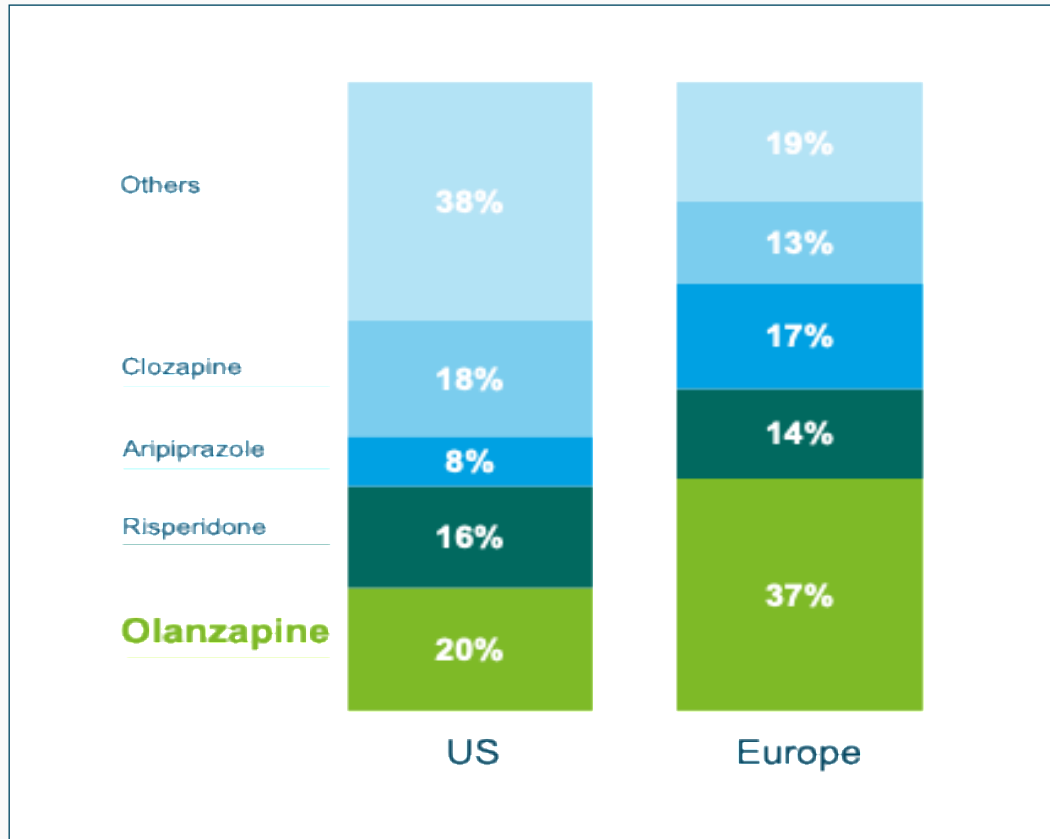
> *FDA approval expected in Q4 2026*

**April 28, 2025**

European submission to for the maintenance treatment of Schizophrenia in adults

> *Acceptance expected in Q2 2026*

# Olanzapine LAI is Designed to Address a High Unmet Medical Need



→ **Most-used oral antipsychotic in schizophrenia**

**Used for most severe and refractory patients**

**First-in-class potential, no valid competitor**

# Olanzapine LAI: Analysts' Peak Sales Forecasts<sup>1</sup>

**Analysts' Consensus: \$2.0B**

**Low: \$1.5B - High: \$2.9B**



**EVERCORE**  
**LEERINK PARTNERS**

**TRUIST**   
**Jefferies**

 **HCW**  
H.C.WAINWRIGHT&CO  
**STIFEL**



 **ODDO BHF**  
 **TP ICAP**

 **Degroof Petercam**

 **PORTZAMPARC**  
BNP PARIBAS GROUP  
 **Kepler Cheuvreux**

## Revenue potential for MedinCell

<b>Royalties</b>	<b>Mid- to high-single digit on all net sales</b>
<b>Milestones</b>	<b>US Approval: \$4 M</b> <b>EU Approval: \$3 M</b> <b>Commercial: \$105 M</b>

<sup>1</sup>Based on most recent information received from analysts: Evercore, Truist, HC Wainwright, Jefferies, Stifel, Oddo BHF, Portzamparc, Kepler Cheuvreux, TP-Icap. Any opinions, estimates, or forecasts regarding MedinCell's performance made by analysts are theirs alone and do not represent opinions, forecasts, or predictions of MedinCell or its management.

# Olanzapine LAI was Designed to Resolve Eli Lilly's PDSS-Linked Market Failure



## Risk of PDSS\* associated with Eli Lilly product's

- Overdose-like reaction risk at each injection
- Caused by unintended intravascular burst exposure during injection

## Regulatory restrictions on use

- Mandatory 3-hour monitoring on site after each injection
- Restricted distribution (REMS)

▶ **Commercial failure**

\* Post-Injection Delirium and Sedation Syndrome

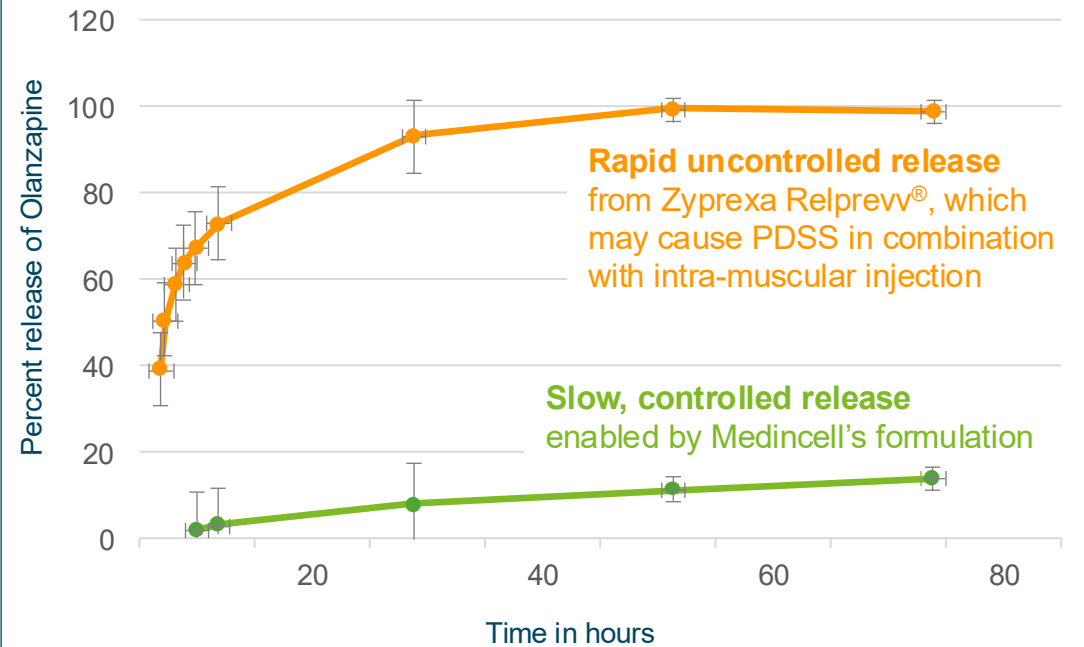
# Strong Evidence that Medincell Olanzapine LAI Will Not Cause PDSS

## Clinical evidence

**3600 injections**  
were required by FDA to demonstrate  
absence of risk of PDSS

**~4,000 injections**  
performed across multiple clinical studies,  
with no PDSS observed

## *In vitro* evidence Release tested in human plasma

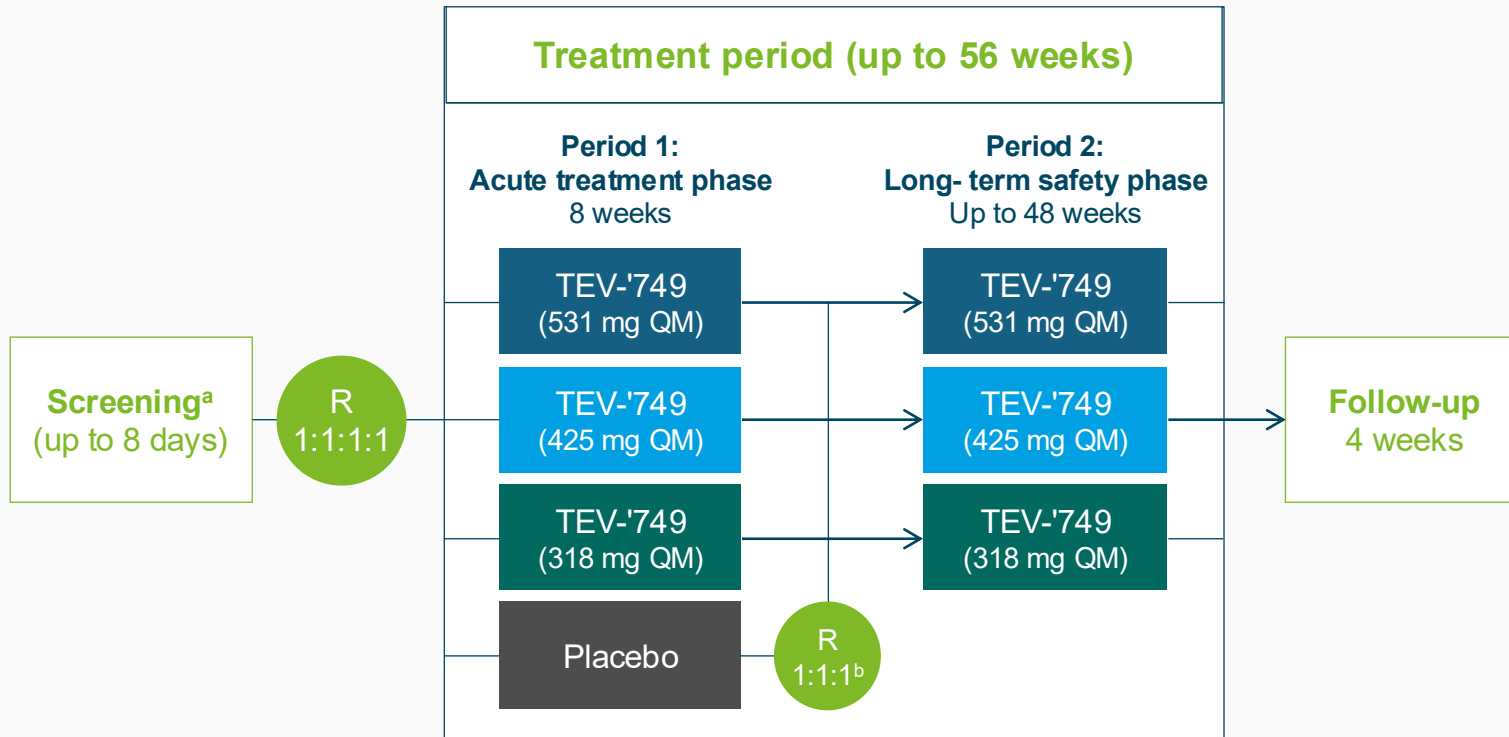


***“No required monitoring for Medincell Olanzapine LAI is expected to drive Olanzapine LAI usage growth.”***

Teva Q3 2025 Results presentation, Nov. 5, 2025

# SOLARIS - Pivotal Phase 3

Phase 3 randomized, double-blind, placebo-controlled (Period 1), open-label long-term safety (Period 2) trial to evaluate efficacy, safety in adult patients with acute exacerbation of schizophrenia



**Period 1** (8 weeks, n=675) aimed to assess the efficacy and safety, of TEV-'749 schizophrenia.

**In Period 2** (n=423), Period 1 TEV-'749 participants retained their treatment, placebo patients were re-randomized 1:1:1 to TEV-'749 (318 mg, 425 mg, or 531 mg).

TEV-'749 doses were comparable to daily oral olanzapine doses of 10 mg, 15 mg, and 20 mg.

a. Participants entering the trial who had not previously received oral olanzapine within the last year received 2 oral doses of olanzapine for 2 consecutive days at the screening period to assess tolerability. The investigator verified the previous use, tolerability, and duration of olanzapine treatment to assure prior tolerability.  
b. To maintain the blinding in Period 1, all participants were re-randomized between Periods 1 and 2; participants previously assigned to the active treatment groups retained their Period 1 treatment assignment (re-randomization was done to maintain blinding of Period 1, and de facto is a deterministic assignment and not randomization), and participants previously assigned to placebo were randomized to one of the active treatment groups in a 1:1:1 ratio. Participants were hospitalized for >28 days after receiving the first injection. QM, once monthly; R, randomization.

Correll CU, et al. Poster #97 presented at the 38th Psych Congress 2025; December 17-21, 2025; San Diego, CA, USA

# **SOLARIS - Positive Results**

## **Demonstrated long-term clinical effectiveness in all groups**

- **Improvements in PANSS Total Score**
  - **Sustained improvement in CGI-S and PSP scale scores**
- 

## **Long-term safety profile consistent with other olanzapine formulations**

- **No new systemic safety signals**
- **Mild/moderate injection site reactions decreasing with continued dosing**
- **No PDSS Events**

Sources: Teva Pharmaceuticals – Poster presented at the 38<sup>th</sup> Psych Congress 2025; September 17-21, 2025; San Diego, CA, USA

# Olanzapine LAI, Expected Timeline



9 Dec. 2025



**NDA submission to U.S. FDA**  
(10-Month review)

Feb. 2026



**NDA acceptance by U.S. FDA**

Q4 2026

**Approval by U.S. FDA**  
**U.S. commercial launch**



Q2 2026



**Submission in Europe**

# Multiple Patents Granted on Olanzapine LAI

**OLANZAPINE LAI**

US patent protection

**2044**

# AbbVie #1

**API and indication not disclosed**

**April 2024**

Strategic co-development and licensing agreement with AbbVie (up to 6 programs), \$35 million upfront

**September 2024**

Initiation of preclinical and supportive CMC work to advance AbbVie #1 into clinical development

***Expected to Be Ready for Launch of Clinical Development in 2026***

# **AbbVie #1 Expected to Be Ready for Launch of Clinical Development in 2026**

**All pre-IND activities conducted by Medincell**

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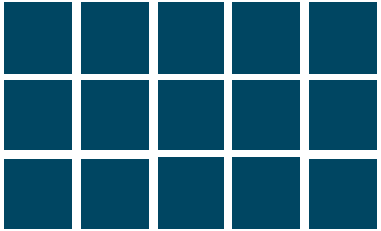




**AbbVie to conduct all clinical development**


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**Mid-single- to low-double-digit royalties  
\$315 million in potential milestones**

# Third Growth Engine Pipeline & Innovation

# Advancing Innovation Across the Pipeline

FORMULATION	PRECLINICAL	CLINICAL PHASE 3	NDA	MARKET
<p><b>~10-15</b> in-house and partnered programs</p> 	<p><b>AbbVie #1</b> Confidential API and indication</p> 	<p><b>Celecoxib Intraarticular</b> (mdc-CWM) Postoperative pain</p> 	<p><b>Olanzapine LAI</b> (mdc-TJK) Schizophrenia</p> 	<p><b>Risperidone LAI</b> <b>UZEDY®</b> Schizophrenia / BP-I</p> 

<p><b>Global Health programs</b></p>	<p><b>Contraception</b> (mdc-WWM) Progestin 6-Month Gates Foundation</p>
<p><b>Tuberculosis</b> Macozinone</p> 	<p><b>Malaria</b> (mdc-STM) Ivermectin 3-Month Gates Foundation</p>

# mdc-CWM: Regulatory Path to Approval



The first Phase 3 study missed the primary pain endpoint at Week 2, while showing positive results at Day 7 and on long-term functional outcomes

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Clear differentiation from approved and investigational therapies

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FDA meeting confirmed clear guidance for a second Phase 3 design focused on TKR-naïve subgroup

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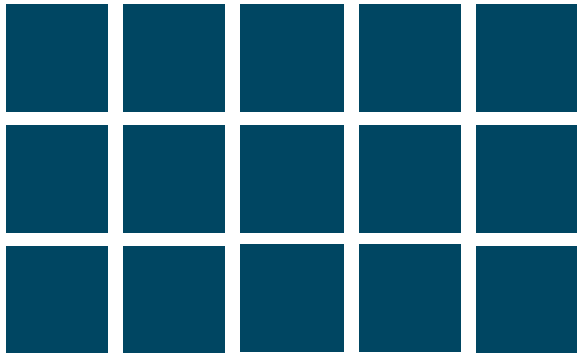
AIC is actively preparing the next Phase 3 study

# Early-Stage Pipeline

## FORMULATION

**~10-15**

in-house and  
partnered programs



## In-house programs & partnerships

(Big Pharma, Late-Stage Biotech, Global Health)

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## Commercial stage & development stage drugs

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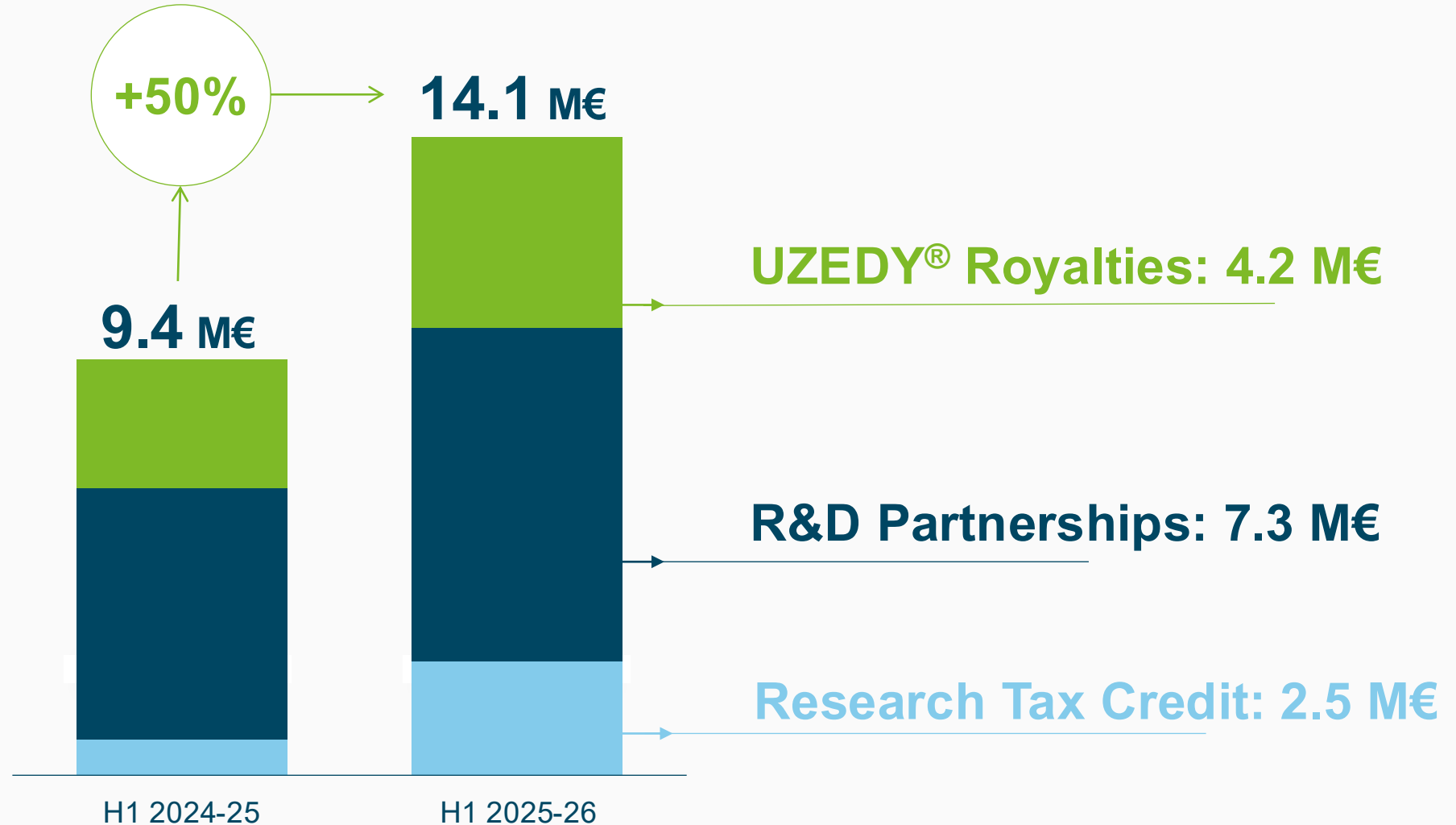
5+ therapeutic areas

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Several candidates with blockbuster potential

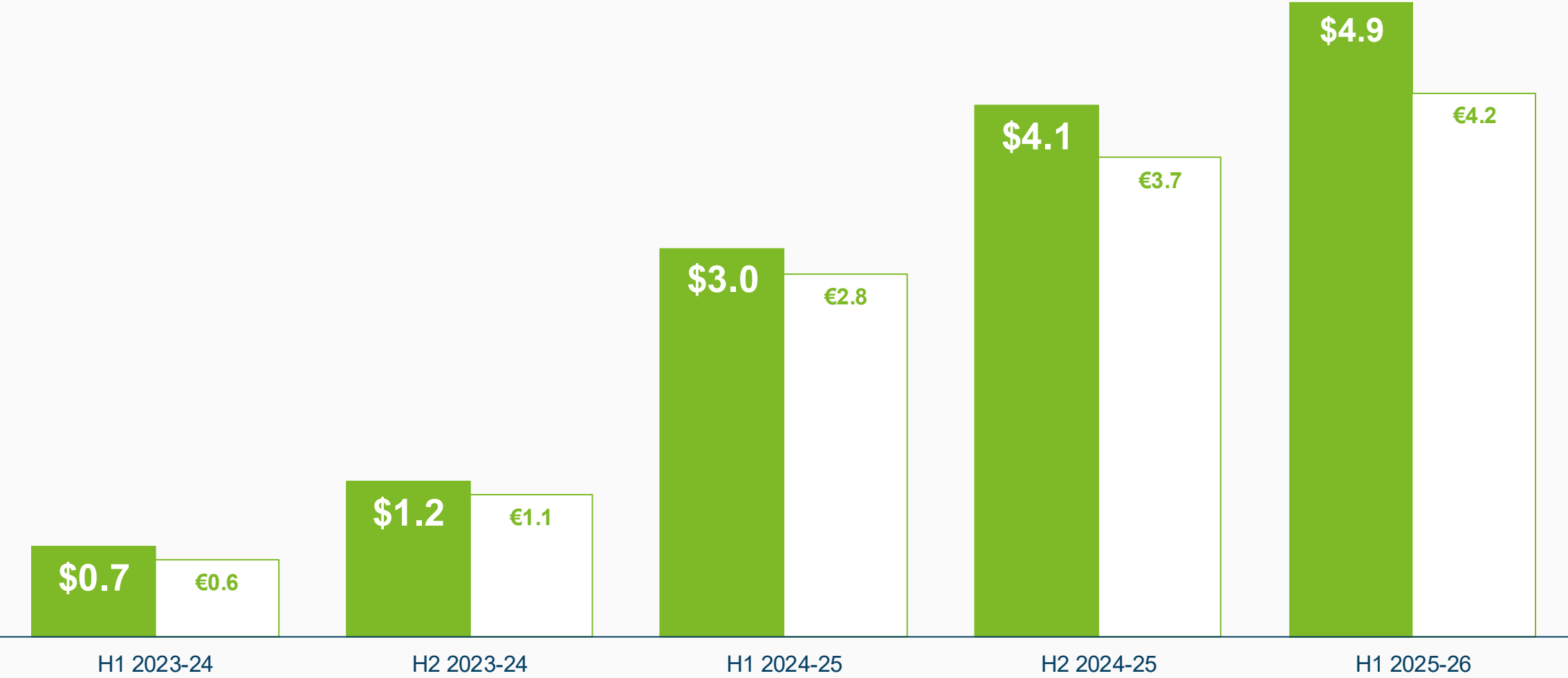
# Financials

# +50% Year-on-Year Total H1 Revenue Growth



# +65% Increase in UZEDY<sup>®</sup> Royalties

(H1 - M\$ - YoY)

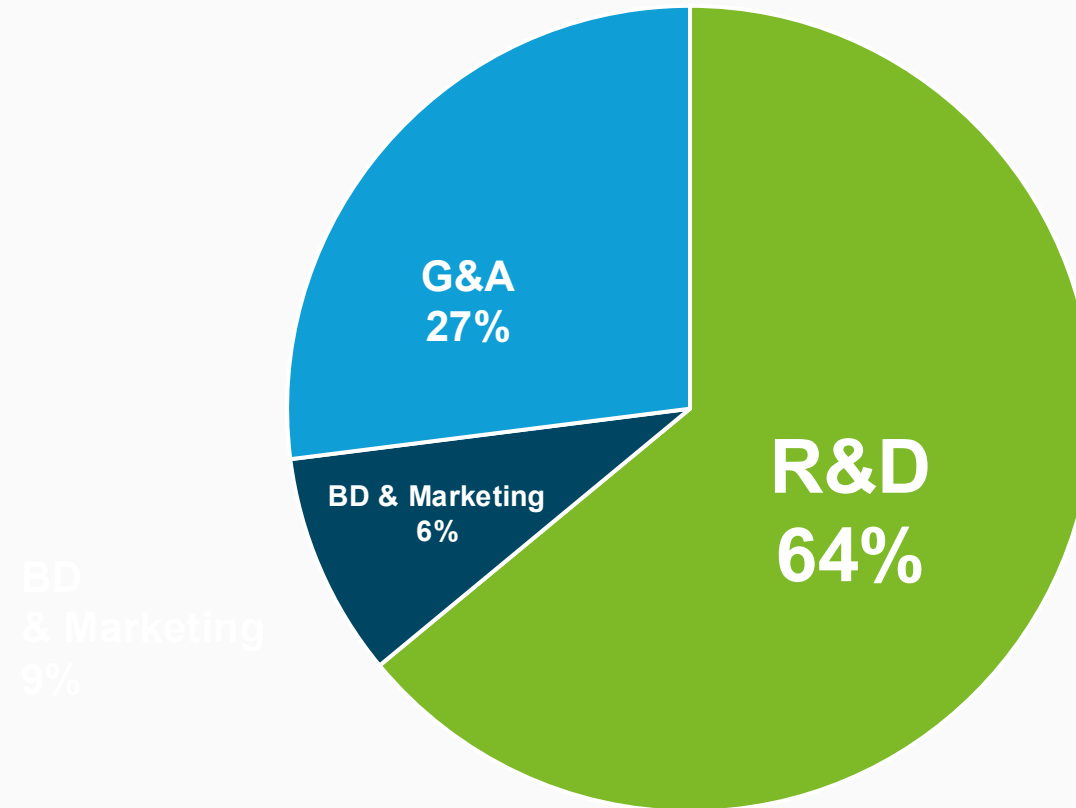


# H1 FY2025-26 Income Statement

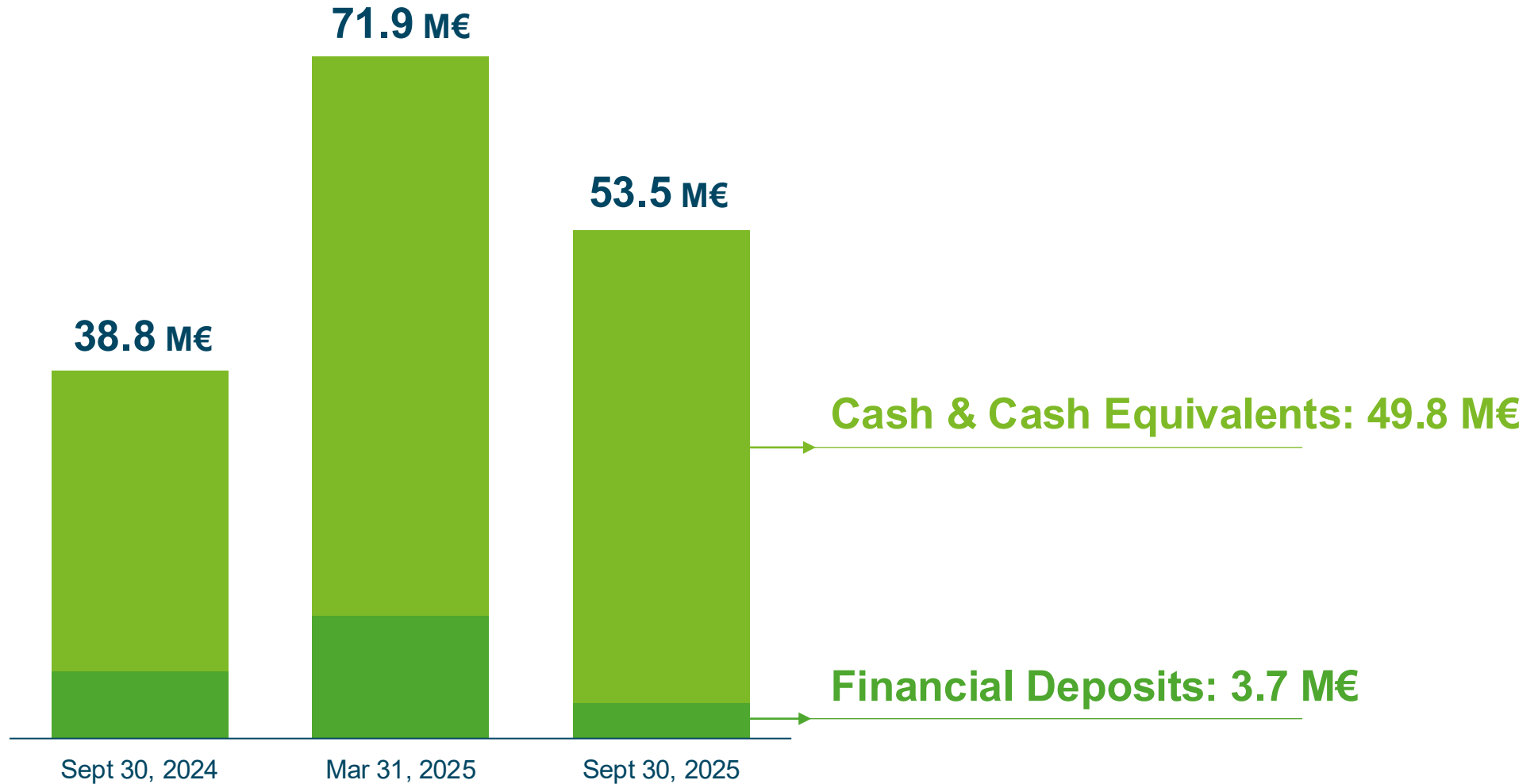
<b>Total income</b>	<b>14.1</b>	<b>+50% YoY</b>
of which revenue	11.6	+35% YoY
<b>Operating expenses</b>	<b>20.8</b>	<b>+22% YoY</b>
<b>Operating result</b>	<b>-6.6</b>	<b>13% improvement YoY</b>
<b>Financial result</b>	<b>-9.5</b>	
of which non-cash Impact of the change in the fair value of EIB Warrants	-6.8	Discussion progressing with EIB to eliminate warrants impact
<b>Net result</b>	<b>-16.1</b>	<b>10% decrease YoY</b>

# H1 Operating Expenses: 22.8 M€

(FY 2025-26)

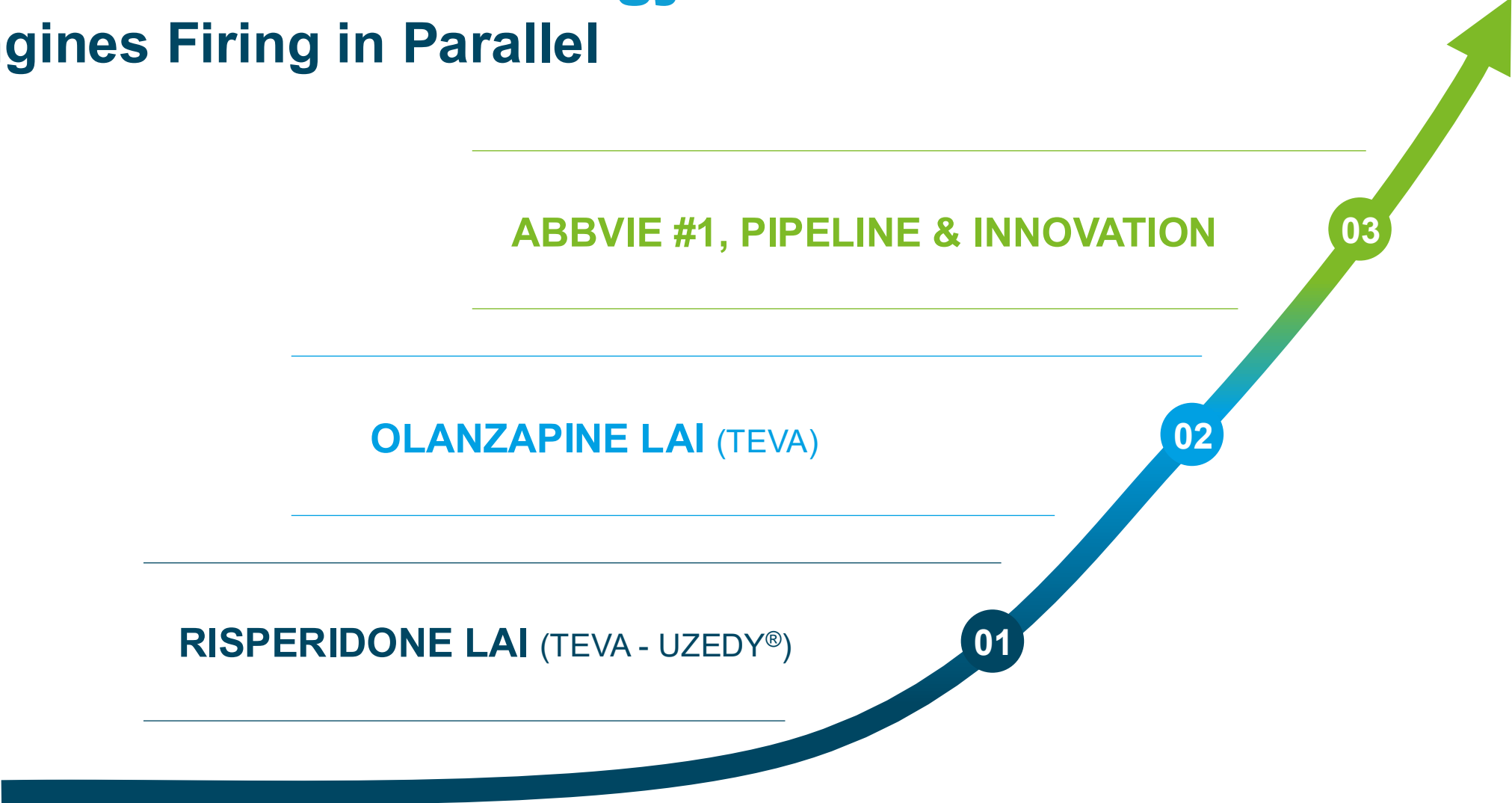


# Cash Position



# “Shift to Growth” Strategy

## 3 Engines Firing in Parallel



**medincell.**

**Thank you**