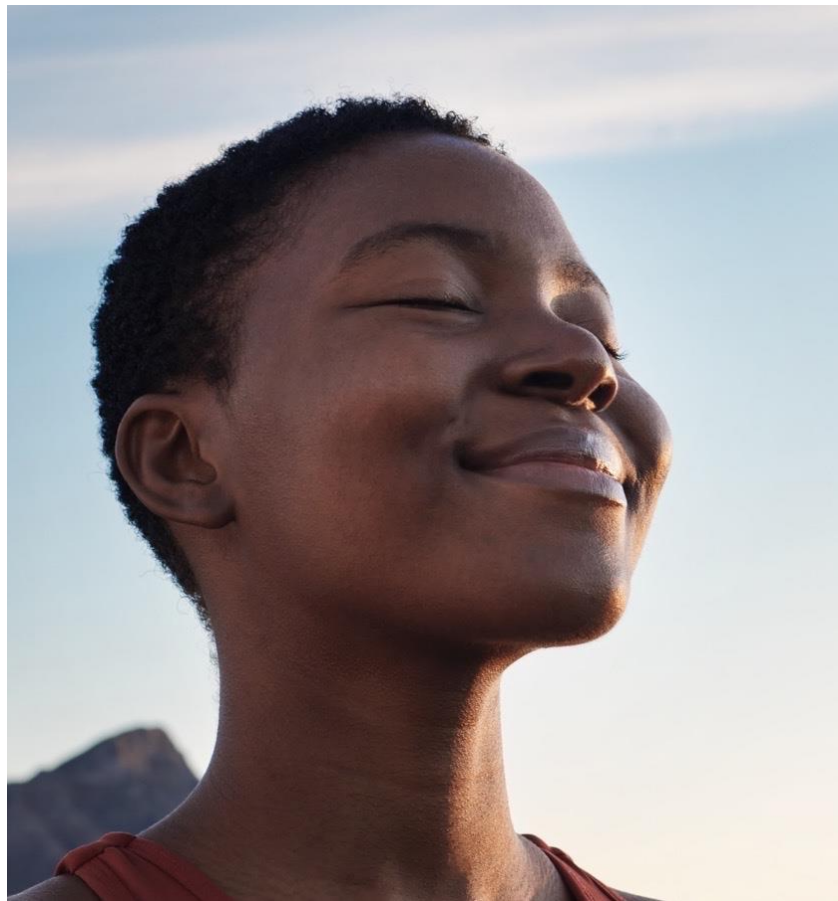


BREAKTHROUGH MEDICINES

WITH LONG-ACTING INJECTABLES (LAI)



IMPORTANT NOTICE - YOU MUST READ THE FOLLOWING BEFORE CONTINUING

This presentation contains 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 presentation 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.

These statements may include, but are not limited to, any statement beginning with, followed by or including words or phrases such as "objective", "believe", "anticipate", "expect", "foresee", "aim", "intend", "may", "anticipate", "estimate", "plan", "project", "will", "may", "probably", "potential", "should", "could" and other words and phrases of the same meaning or used in negative form. Forward-looking statements are subject to inherent risks and uncertainties beyond the Company's control that may, if any, cause actual results, performance, or achievements to differ materially from those anticipated or expressed explicitly or implicitly by such forward-looking statements. A list and description of these risks, contingencies and uncertainties can be found in the documents filed by the Company with the Autorité des Marchés Financiers (the "AMF") pursuant to its regulatory obligations, including the Company's universal registration document, filed with the AMF on July 28, 2022, (the "Universal Registration Document"), as well as in the documents and reports to be published subsequently by the Company. In particular, readers' attention is drawn to the section entitled "Facteurs de Risques" on page 24 of the Universal Registration Document.

Any forward-looking statements made by or on behalf of the Company speak only as of the date they are made. Except as required by law, the Company does not undertake any obligation to publicly update these forward-looking statements or to update the reasons why actual results could differ materially from those anticipated by the forward-looking statements, including in the event that new information becomes available. The Company's update of one or more forward-looking statements does not imply that the Company will make any further updates to such forward-looking statements or other forward-looking statements. Readers are cautioned not to place undue reliance on these forward-looking statements.

This presentation is for information purposes only. The information contained herein does not constitute an offer to sell or a solicitation of an offer to buy or subscribe for the Company's shares in any jurisdiction, in particular in France. Similarly, this presentation does not constitute investment advice and should not be treated as such. It is not related to the investment objectives, financial situation, or specific needs of any recipient. It should not deprive the recipients of the opportunity to exercise their own judgment. All opinions expressed in this document are subject to change without notice. The distribution of this presentation may be subject to legal restrictions in certain jurisdictions. Persons who come to know about this presentation are encouraged to inquire about, and required to comply with, these restrictions.

All information in the presentation speaks only as of (1) the date hereof, in the case of information about the Company and (2) the date of such information, in the case of information from persons other than the Company. The Company does not undertake any duty to update or revise the information contained herein, publicly or otherwise. The Company has not independently verified any third-party information and makes no representation as to the accuracy or completeness of any such information.

UZEDY® is a trademark of Teva Pharmaceuticals.

CONTENT

Corporate overview - 4

April 2024 - Strategic collaboration with AbbVie - 6

BEPO[®], Long-Acting Injectable cutting-edge technology platform - 7

Product on market - 10

- **UZEDY[®]** (*marketed by Teva Pharmaceuticals*)
Monthly and every 2 months subcutaneous risperidone for treatment of schizophrenia

R&D pipeline - 18

- **mdc-TJK**
Monthly subcutaneous olanzapine LAI for treatment of schizophrenia
Clinical Phase 3 results expected in H2 2024
- **mdc-CWM**
Intraarticular celecoxib for post-operative pain and inflammation management
Clinical Phase 3 results expected in Q1 2024 in Total Knee Replacement (TKR)
- **mdc-WWM**
Best-in-class contraceptive LAI (preclinical)
- **mdc-STM**
Global Health program to fight malaria (preclinical)

Financials & extra-financial performance - 35



CORPORATE OVERVIEW

BEPO®

Pioneering Long-Acting Injectable (LAI) innovator with breakthrough technology platform

UZEDY®

First product marketed by Teva Pharmaceuticals in the U.S. since May 2023

- UZEDY®, monthly and every 2 months subcutaneous risperidone for treatment of schizophrenia
- Approved by the FDA in April 2023
- First royalties received from Teva Pharmaceuticals in 2023
- Targeting primarily U.S. \$4.4 billion 12% CAGR market, up to \$105m milestones + royalties for MedinCell



Rich R&D pipeline including first-in-class therapies and potential blockbusters

- 2 products already in phase 3
- Growing number of products at formulation and preclinical stages

Tier one partners

- Teva Pharmaceuticals
- AbbVie
- Bill & Melinda Gates (*\$23m grant, Global Health*)
- Joint venture with Corbion (*GMP commercial Polymer*)



Return to profitability

- First commercial royalties from UZEDY® received in 2023, ramp-up in progress
- Strong commercial potential of mdc-TJK, next product expected to reach market in 2025-2026
- Active ongoing discussions for new strategic partnerships driven by booming interest for LAIs

MEDCL
LISTED
EURONEXT

PRODUCT PORTFOLIO AND R&D PIPELINE



● with Teva Pharmaceuticals ● with AIC ● with AbbVie ● with the Bill & Melinda Gates Foundation ● with Unitaid ● in-house program or undisclosed partner

April 2024

STRATEGIC COLLABORATION WITH ABBVIE

CO-DEVELOPMENT AND LICENSING AGREEMENT

Up to 6 Long-Acting Injectable therapies

- Multiple therapeutic areas and indications
- First program candidate selected; preclinical and CMC activities underway

Medincell to conduct formulation and preclinical activities

AbbVie to conduct clinical development

AbbVie responsible for commercialization globally

FINANCIAL METRICS

\$35 million upfront payment

up to \$1.9 billion in potential commercial and development milestones

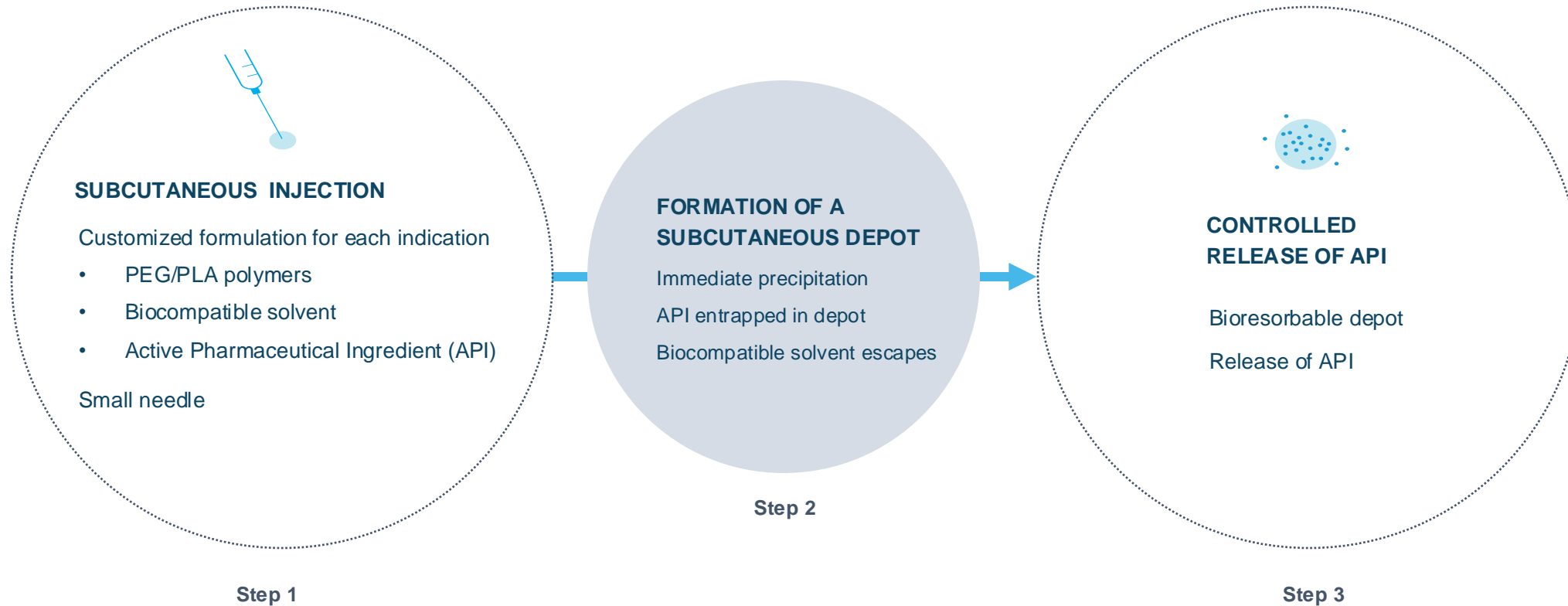
Tiered mid-single to low-double digit royalties



BEPO®

Long-acting injectables
cutting-edge technology platform

Long-Acting Injectable cutting-edge technology platform



BEPO® POLYMERS SECURED THROUGH INDUSTRIAL JOINT VENTURE WITH CORBION



Limited scale-up risk

Research and clinical batch polymers come from same production line as commercial polymers

Secure supply, ensure quality & preserve manufacturing IP

Dual GMP manufacturing facilities – Europe and U.S.

DMF filed in the US and Canada

50/50 Joint-Venture



Leading manufacturer of biomedical polymers worldwide

Pharma production standards (ICHQ & GMP)

Listed on Euronext Amsterdam (CRBN - market cap: ca. €1.1B as of January 1, 2023)



PRODUCT ON MARKET

UZEDY®

Monthly and every 2 months subcutaneous risperidone
for treatment of schizophrenia



UZEDY®

Market authorization by U.S. FDA on April 28, 2023

Commercialization by Teva Pharmaceuticals since May 2023

Targeting primarily US \$4.4 billion 12% CAGR market

2024 Teva's revenue outlook: ~\$80 million

MedinCell eligible for

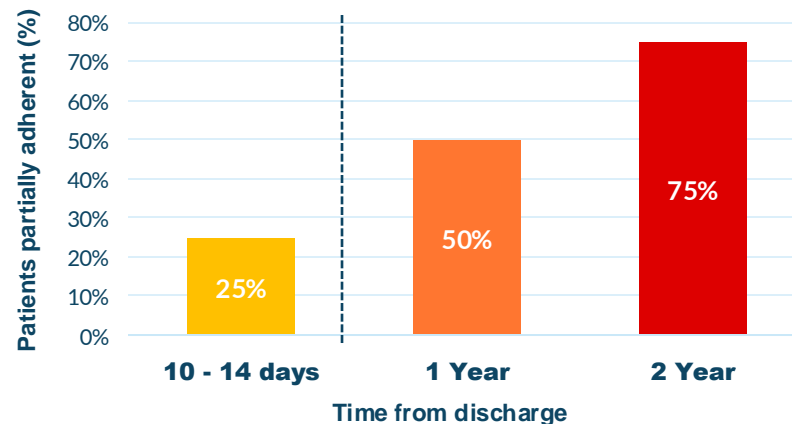
- mid-to high-single digit royalties on net sales
- up to \$105m in commercial milestones

ADHERENCE TO TREATMENT IS CRUCIAL IN SCHIZOPHRENIA

ca. 1% of the worldwide population will develop schizophrenia in their lifetime¹

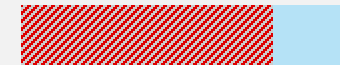
Approximately 80% of patients experience multiple relapses during the first five years of treatment², and each relapse carries a biological risk of loss of function, treatment refractoriness, and changes in brain morphology^{3,4}

Treatment compliance worsens over time⁵



¹ SSPAA, About Schizophrenia, Available at: sczaction.org/about-schizophrenia/ - Accessed June 2023; ² Emsley, R., & Kilian, S. (2018). Efficacy and safety profile of paliperidone palmitate injections in the management of patients with schizophrenia: an evidence-based review. *Neuropsychiatric disease and treatment*, 14, 205-223; ³ Emsley, R., Chhiza, B., Asmal, L., et al. (2013). The nature of relapse in schizophrenia. *BMC Psychiatry* 13, 50; ⁴ Andreasen, N. C., et al. (2013). Relapse duration, treatment intensity, and brain tissue loss in schizophrenia: a prospective longitudinal MRI study. *The American journal of psychiatry*, 170(6), 609-615; ⁵ Velligan, D. I., et al. *Psychiatric Serv.* 2003;54(5):655-667. Weinstein P. J., et al. Medication noncompliance in schizophrenia: I. assessment. *Journal of Practical Psychiatry and Behavioral Health*. 1997;3:106-110; ⁶ Comprehensive understanding of schizophrenia and its treatment, Maguire GA. *Am J Health Syst Pharm*. 2002; ⁷ Analysis Group, Otsuka, Lundbeck LLC - 2016

75%



of patients had discontinued medication within 2 years due to insufficient efficacy, intolerable side effects or for other reasons

In the U.S., schizophrenia accounts for 20% of all hospital bed-days and over 50% of all psychiatric beds⁶

Annual schizophrenia costs are estimated between \$134 and \$174 bn⁷

UZEDY[®], STRONG DIFFERENTIATION THANKS TO BEPO[®]



SUBCUTANEOUS INJECTION (vs. intramuscular)

- Smaller needle (16mm; 21 gauge)
- Multiple injection sites (upper arm and abdomen)
- Lower injection volume (0.1 –0.7 ml)

PREFILLED SYRINGE

- Ready-to-use (no reconstitution needed)
- Can be left out of the refrigerator for up to 90 days












IMMEDIATE ONSET OF ACTION

- Achieves therapeutic levels within 24 hours of first injection
- No loading dose or oral supplementation required

DESIRABLE PHARMACOKINETICS

- Multiple dosing options corresponding to oral risperidone
- Can be dosed every month or every two months

UZEDY[®], DIFFERENTIATED PROFILE FOR SCHIZOPHRENIA PATIENTS

	 UZEDY	Invega Sustenna [®]
Molecule	Risperidone	Paliperidone
Efficacy	 Efficacy profile consistent with risperidone	 Efficacy profile consistent with paliperidone
Safety	 Safety profile consistent with risperidone	 Safety profile consistent with paliperidone
Dose frequency	1-Month, 2-Month	1-Month
SC injection (and volume)	 (0.1-0.7 mL)	 ¹ (0.25-1.5 mL)
Therapeutic levels in 24h		 ²
No oral supplement / loading dose		 ²

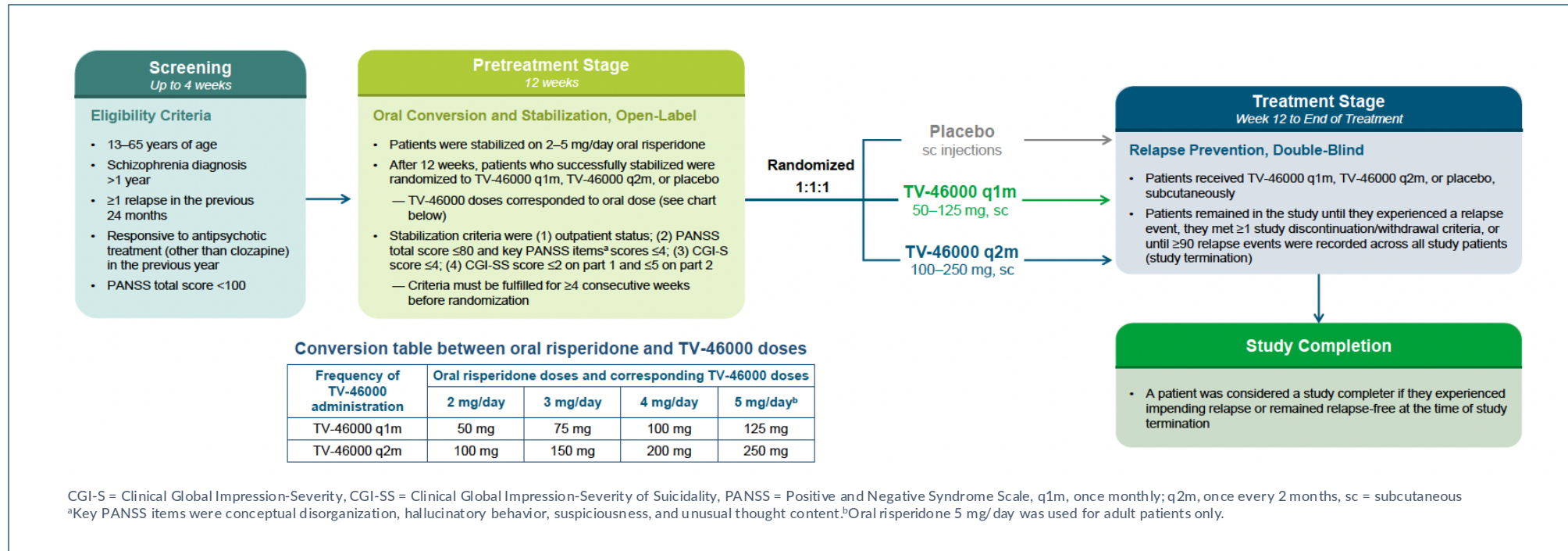
3M Invega Trinza[®] and 6M Invega Hafyera[®] formulations also available

← **70% of target LAI patients³ are on 1M formulation (*preferred by psychiatrists for patient monitoring*)**

1. Intramuscular injection 2. As per prescribing information, Invega Sustenna requires two initial deltoid IM injections of 234mg on day 1 and 156mg on day 8 to help attain therapeutic levels rapidly 3. U.S. patients on risperidone/paliperidone LAIs
 Note: No head-to-head studies have been conducted comparing UZEDY with any other therapy. The information on this slide should not be construed to imply any difference in safety, efficacy, or other clinical outcome. All trademarks referenced are properties of their respective owners
 Sources: UZEDY RISE Phase III pivotal study and prescribing information; Invega Sustenna Phase III pivotal study and prescribing information

UZEDY[®], EFFICACY AND SAFETY IN SCHIZOPHRENIA

Phase 3, Randomized, Double-Blind, Relapse Prevention Study (RISE Study)



In total, 1 267 patients were screened, 863 were enrolled, and 544 were randomized

The primary endpoint was time to impending relapse and secondary endpoints included proportions of patients with impending relapse at week 24 and proportion of patients who maintained stability at week 24

TV-46000 is the investigational product codename used by Teva during regulatory development of mdc-IRM

Source: Subcutaneous Risperidone (TV-46000) Efficacy and Safety in Schizophrenia: a Phase 3, Randomized, Double-Blind, Relapse Prevention Study (RISE Study)

John M. Kane,^{1,3} Eran Harary,⁴ Oma Tohami,⁴ Roy Eshet,⁴ Avia Merenlender-Wagner,⁴ Nir Sharon,⁵ Mark Sueti,⁶ Kelli R. Franzenburg,⁶ Christoph U. Correll^{1,3,6}

¹Zucker Hillside Hospital, Northwell Health, Department of Psychiatry, Glen Oaks, NY, United States; ²Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, Department of Psychiatry and Molecular Medicine, Hempstead, NY, United States; ³Feinstein Institutes for Medical Research, Institute of Behavioral Science, Manhasset, NY, United States; ⁴Teva Pharmaceutical Industries, Global Specialty Research & Development, Netanya, Israel; ⁵Teva Pharmaceutical Industries, Global Medical Affairs, West Chester, PA, United States; ⁶Charité–Universitätsmedizin Berlin, Department of Child and Adolescent Psychiatry, Berlin, Germany

Presented at Psych Congress 2021; October 29–November 1, 2021

UZEDY[®], KEY OUTCOMES FROM THE PIVOTAL PHASE 3 STUDY

EFFICACY

mdc-IRM significantly prolonged time to impending relapse compared to placebo¹

- 80.0% and 62.5% reduction in risk of relapse vs placebo for monthly and every two-month UZEDY[®], respectively
- x5 and x2.7 increase in time to impending relapse with monthly and every two-month UZEDY[™], respectively
- 7% and 13% of patients using monthly and every two-month UZEDY[®], respectively, relapsed within 24 months vs 28% of placebo patients

mdc-IRM provided continued symptom improvement in patients with schizophrenia²

SAFETY

No new safety signals versus accumulated safety data for oral risperidone and other long-acting risperidone formulations³



TV46000 is the investigational product codename used by Teva during regulatory development of mdc-IRM

¹ Subcutaneous Risperidone (TV-46000) Efficacy and Safety in Schizophrenia: a Phase 3, Randomized, Double-Blind, Relapse Prevention Study (RISE Study) - John M. Kane, Eran Harary, Oma Tohami, Roy Eshet, Avia Merenlender-Wagner, Nir Sharon, Mark Suett, Kelli R. Franzenburg, Christoph U. Correll ; ² TV-46000 Provided Continued Symptom Improvement in Patients With Schizophrenia in the Phase 3, Randomized, Double-Blind Relapse Prevention RISE Study - John M. Kane, Christoph U. Correll, Oma Tohami, Roy Eshet, Avia Merenlender-Wagner, Nir Sharon, Mark Suett, Kelli R. Franzenburg, Eran Harary ; ³ Behavioral-, Metabolic-, Endocrine-, and Cardiovascular-Related Adverse Events in Patients With Schizophrenia Treated With TV-46000 - Christoph U. Correll, Helena Knebel, Eran Harary, Roy Eshet, Oma Tohami, Mark Suett, Nir Sharon, Kelli R. Franzenburg, John M. Kane ; Presented at Psych Congress 2021; October 29–November 1, 2021



R&D PIPELINE

Long-acting injectables based on BEPO

R&D PIPELINE

Long-acting injectables based on BEPO®

CLINICAL PHASE 3

mdc-TJK

Olanzapine 1-Month
Schizophrenia

mdc-CWM

Celecoxib -
Intraarticular
Postoperative pain

PRECLINICAL

mdc-WWM

Progestin 6-Month
Contraception

mdc-STM

Ivermectin 3-Month
Malaria

AbbVie (1/6)

FORMULATION

Confidential

Confidential

Confidential

Confidential

Confidential

Confidential

Confidential

Confidential

Confidential

Confidential

● with Teva Pharmaceuticals ● with AIC ● with AbbVie ● with the Bill & Melinda Gates Foundation ● with Unitaid ● in-house program or undisclosed partner



CLINICAL PHASE 3 | mdc-TJK

Olanzapine 1-Month

Once-monthly subcutaneous long-acting injection of the atypical antipsychotic olanzapine

May 2024: Positive Phase 3 efficacy results in adult patients with schizophrenia

Phase 3 safety data topline readout expected in H2 2024

May be the first long-acting olanzapine with a favorable safety profile

mdc-TJK, MEDICAL NEED AND PRODUCT RATIONALE

Impact of relapse and psychosis in schizophrenia

High non-adherence rates with oral medication, eg 64% of patients assigned to olanzapine discontinued treatment within 18 months¹

Approved Olanzapine IM LAI

Existing olanzapine LAI has limited use:

- Black box warning for PDSS as a result of dose dumping hypothesized to be caused by a combination of IM route of administration and formulation characteristics²
- Only available through restricted distribution (REMS) program
- IM injection, requires a loading dose for low and middle doses

Envisaged mdc-TJK

Monthly long-acting subcutaneous injectable :

- SC administration & formulation characteristics of mdc-TJK may mitigate the hypothesized causes of PDSS
- No complex initiation program with no need for loading

LAI = long-acting injectable,

PDSS = post-injection delirium/sedation syndrome,

REMS = Risk Evaluation and Mitigation Strategy

Product characteristics are aspirational, and the product is still in development

References: 1. Lieberman JA, et al. N Engl J Med. 2005;353(12):1209-1223. 2. McDonnell, D.P., Delke, H.C., Bergstrom, R.F. et al. BMC Psychiatry. 10, 45 (2010). <https://doi.org/10.1186/1471-244X-10-45>. 3. Correll CU, et al. Am J Psychiatry. 2020;177(12):1168-1178. doi:10.1176/appi.ajp.2020.19121279; 4. Citrome L. CNS Spectr. 2021;26(2):118-129. doi:10.1017/S1092852921000243; 5. Roberge C, et al. Journal of Controlled Release. 2020; 319: 416-427.

mdc-TJK - POTENTIAL TO BE THE FIRST LAI OLANZAPINE WITH FAVORABLE SAFETY PROFILE

	1990's ▼		Today ▼
	Oral olanzapine	Zyprexa Relprevv® (LAI)	mdc-TJK Target profile
Efficacy	✓	✓	✓ Expect efficacy consistent with olanzapine
Safety	Well characterized safety profile ¹	Well-characterized safety profile ¹ with PDSS occurrence	Expected in line with oral olanzapine ² BEPO ^{®3} technology controls the steady release of API, as demonstrated with UZEDY [®]
Convenience	✗ Once daily	≈ Once every 2 weeks	✓ Once monthly

PDSS: Post-Injection Delirium/Sedation Syndrome PK: Pharmacokinetics

¹. With boxed warning for increased mortality in elderly patients with dementia-related psychosis ². Expected boxed warning for increased mortality in elderly patients with dementia-related psychosis

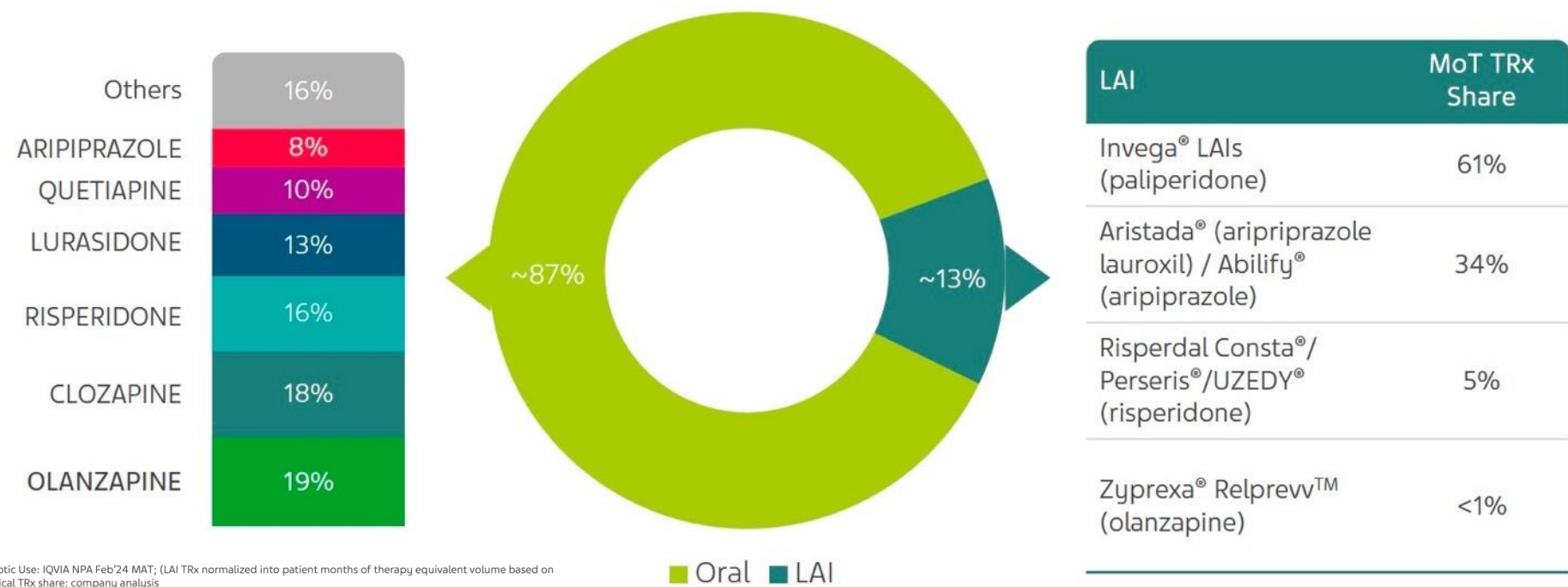
Note: No head-to-head studies have been conducted comparing olanzapine (749) with any other therapy. The information on this slide should not be construed to imply any difference in safety, efficacy, or other clinical outcome. Olanzapine (749) is an asset under investigation, not approved by regulators. SteadyTeq[®] is a registered trademark of Teva Pharmaceuticals USA, Inc.

³. Licensed under the name SteadyTeq[™] to Teva

OLANZAPINE LAI SIGNIFICANT POTENTIAL

Olanzapine is the most prescribed antipsychotic for schizophrenia in the U.S.

U.S. Schizophrenia Rx market, current atypical antipsychotic use

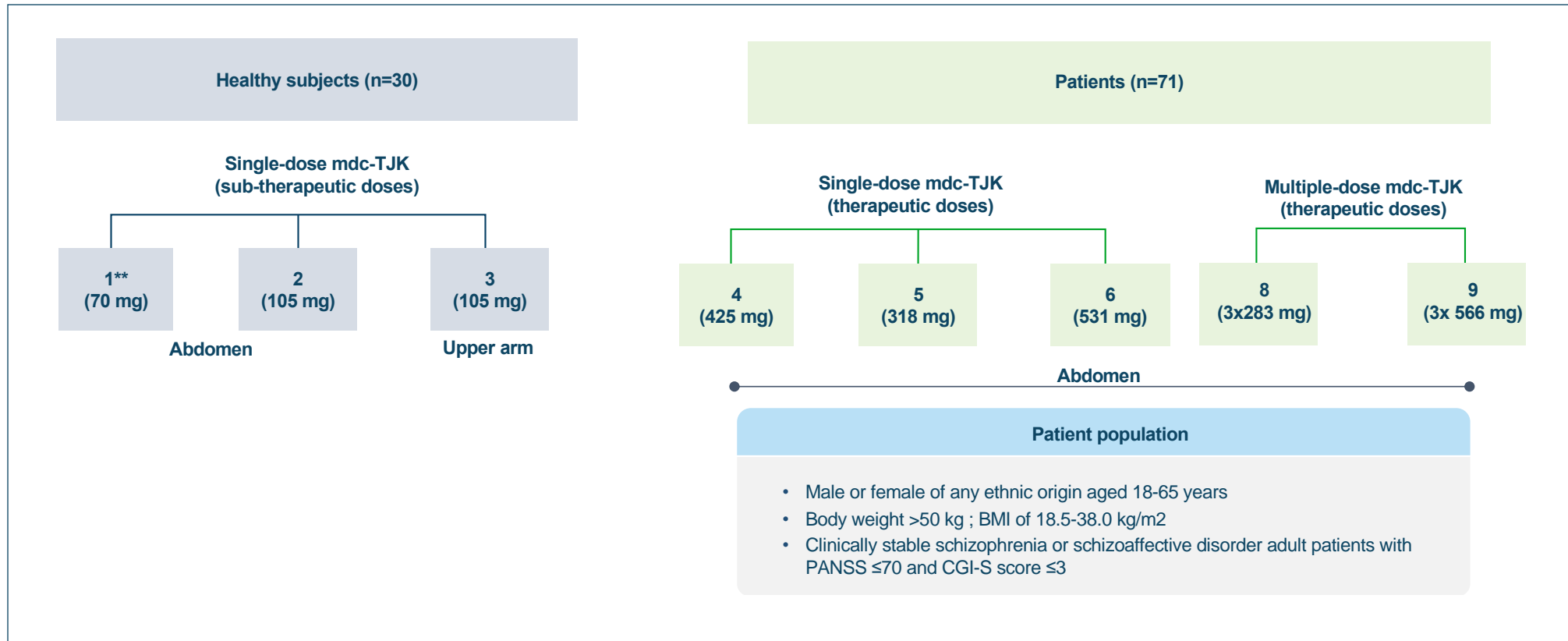


Current Atypical Antipsychotic Use: IQVIA NPA Feb'24 MAT; (LAI TRx normalized into patient months of therapy equivalent volume based on dosing regimen), Oral atypical TRx share: company analysis
Global peak sales source: Evaluate pharma complete data extract for LAI antipsychotics
Trademarks mentioned are the property of their respective owners

Source: Teva earnings call presentation - May 8, 2024

mdc-TJK PHASE 1 SAD/MAD STUDY DESIGN

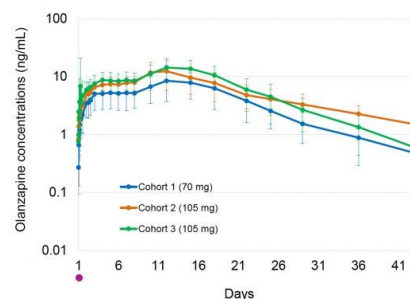
- Overall, 127 participants enrolled
- 101 participants were administered mdc-TJK



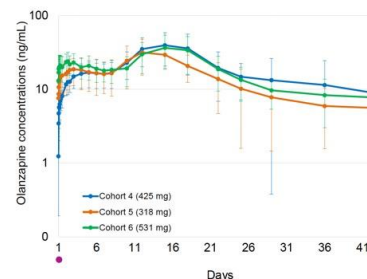
mdc-TJK PHASE 1 SAD/MAD

Pharmacokinetics in healthy volunteers and patients with schizophrenia or schizoaffective disorder

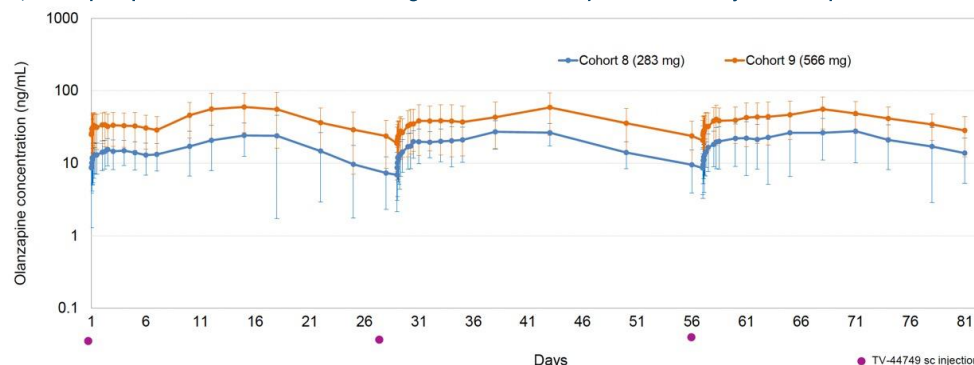
Mean (\pm SD) olanzapine plasma concentrations following single dose TV-44749 sc injections in healthy subjects



Mean (\pm SD) olanzapine plasma concentrations following single dose TV-44749 sc injections in patients with schizophrenia



Mean (\pm SD) olanzapine plasma concentrations following three once monthly TV-44749 sc injections in patients with schizophrenia



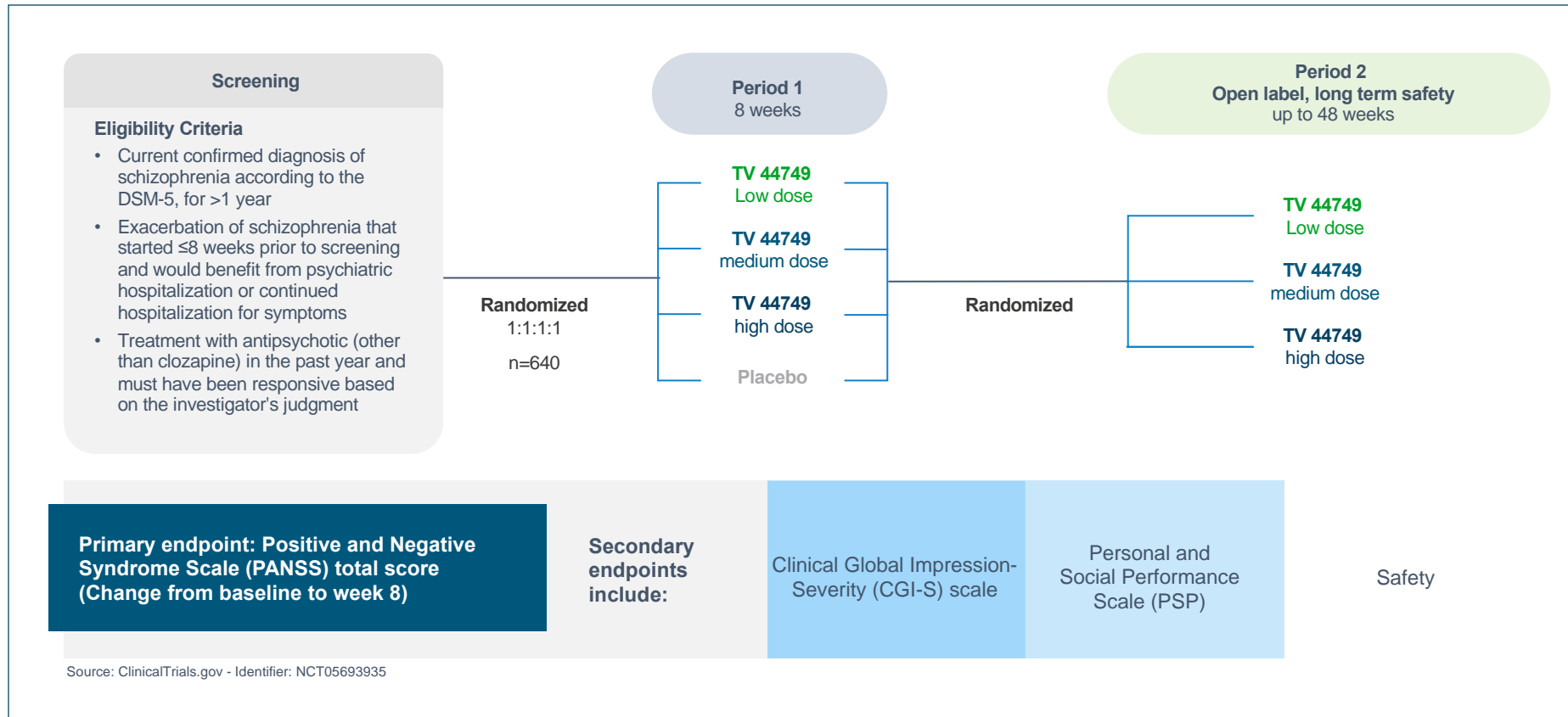
mdc-TJK exhibited favorable characteristics of extended-release profile:

- By reaching clinically relevant therapeutic olanzapine plasma concentrations (≥ 10 ng/mL) within a 1 to 2 day and maintaining them during the 28-day dosing interval
- At steady-state conditions over 28 dosing interval, the systemic exposure of mdc-TJK at doses 318, 425 and 531 mg were comparable to oral daily corresponding doses 10 mg, 15 mg, and 20 mg respectively
- No burst or uncontrolled rise in olanzapine plasma concentrations following mdc-TJK subcutaneous administration was observed

The results of this study, supported dose selection of mdc-TJK in ongoing Phase 3

mdc-TJK, EFFICACY AND SAFETY IN SCHIZOPHRENIA

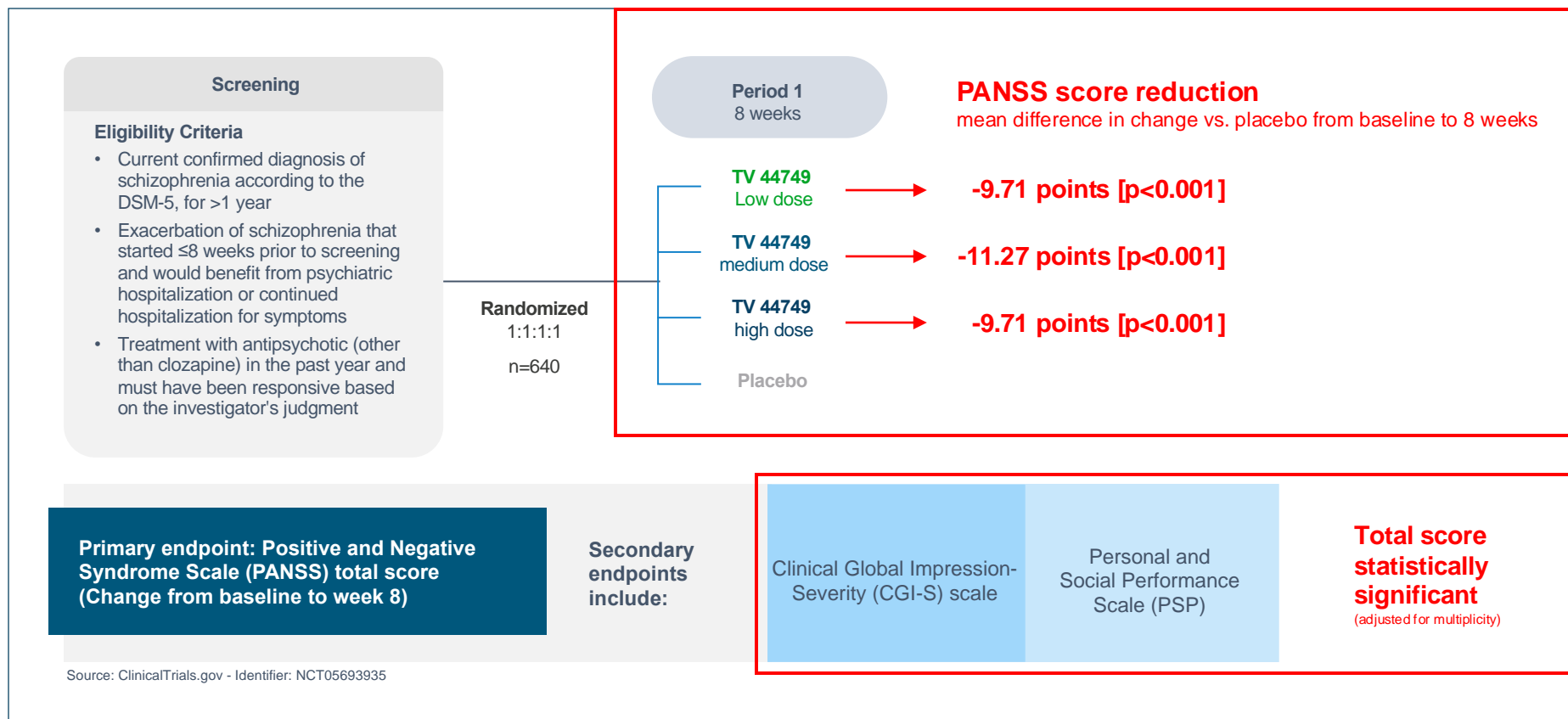
- A Phase 3 Randomized, Double-Blind, Placebo-Controlled Study (SOLARIS)



Study is designed to identify both safety and efficacy, including to identify PDSS event occurrence. However, MedinCell and Teva believe that BEPO® technology and subcutaneous administration will allow olanzapine LAI to have the favorable safety profile.

mdc-TJK, EFFICACY AND SAFETY IN SCHIZOPHRENIA

- May 2024: Positive Phase 3 efficacy results in adult patients with schizophrenia
- Phase 3 safety data topline readout expected in H2 2024

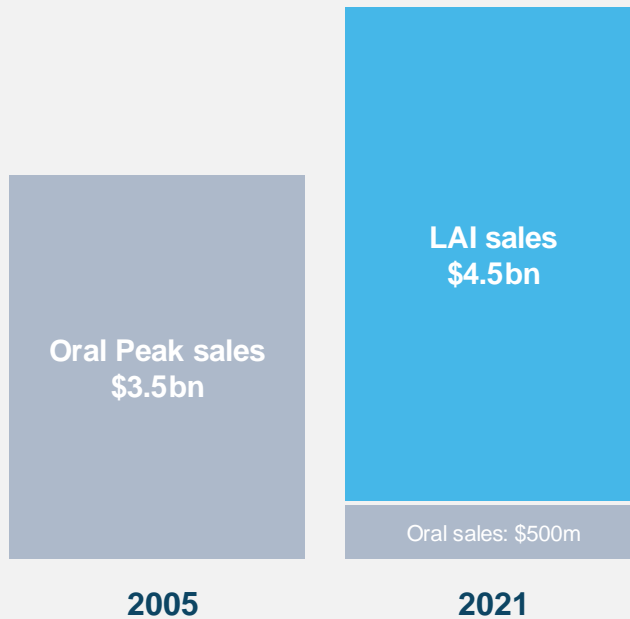


Study is designed to identify both safety and efficacy, including to identify PDSS event occurrence. However, Medincell and Teva believe that BEPO® technology and subcutaneous administration will allow olanzapine LAI to have the favorable safety profile.

STRONG OPPORTUNITY FOR OLANZAPINE LAI WITH FAVORABLE SAFETY PROFILE

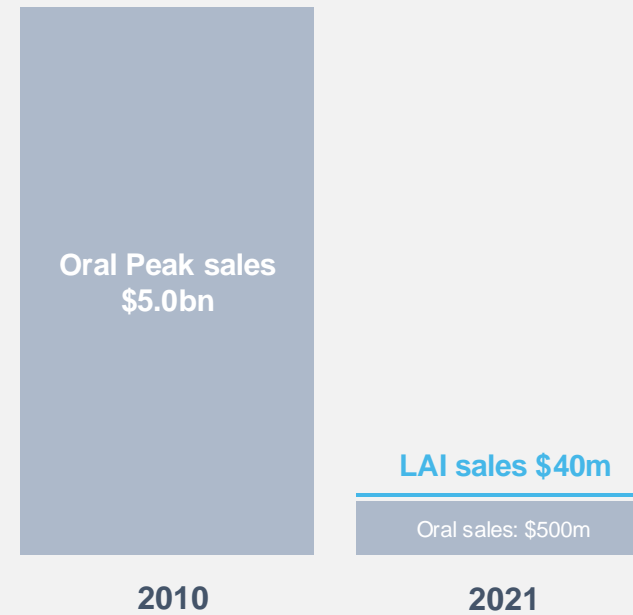
LAI franchise was successfully built with risperidone + paliperidone¹...

- Compensates sales fall after patent expiry of oral form of risperidone (2008)



... when current Olanzapine LAI does not reach potential

- Commercial failure of existing Olanzapine LAIs
- Black box warning from FDA



321 000 of U.S. treated patients out of 1,6m use Olanzapine (2022)¹, mostly with oral administration

Black box warning on existing LAI of olanzapine

- Must be injected in certified centers
- Requires continuous observation of patients by healthcare professional for at least 3 hours after each injection
- Patient must be accompanied to their destination from the health care facility

Sources: 7 Major Markets - Companies reported sales, IQVIA

1. Teva investor day presentation– May 2024



CLINICAL PHASE 3 | mdc-CWM

Intraarticular celecoxib

Intraarticular celecoxib for post-operative pain and inflammation management

Ongoing clinical Phase 3 in Total Knee Replacement (TKR) efficacy results expected in Q1 2024

May be the first product to provide pain relief over several weeks post-surgery

15% of TKR patients become long-term opioid users

Source: 2018 Choices Matter Survey - Exposing a silent gateway to persistent opioid use

mdc-CWM, MEDICAL NEED AND PRODUCT RATIONALE

One-time local delivery during surgery aiming at facilitating patient recovery by:

- Providing post-operative pain relief for weeks (vs. days for existing products)
- Accelerating improvement in knee function
- Potentially decreasing the need for addictive opioids

Little to no systemic exposure reduces risk of adverse issues associated with NSAIDs

Celecoxib was approved by the FDA for pain treatment in 1998. It is often used in the treatment of acute pain, rheumatoid arthritis, ankylosing spondylitis etc.

COLLABORATION WITH ARTHRITIS INNOVATION CORPORATION (AIC)

50-50 profit sharing agreement

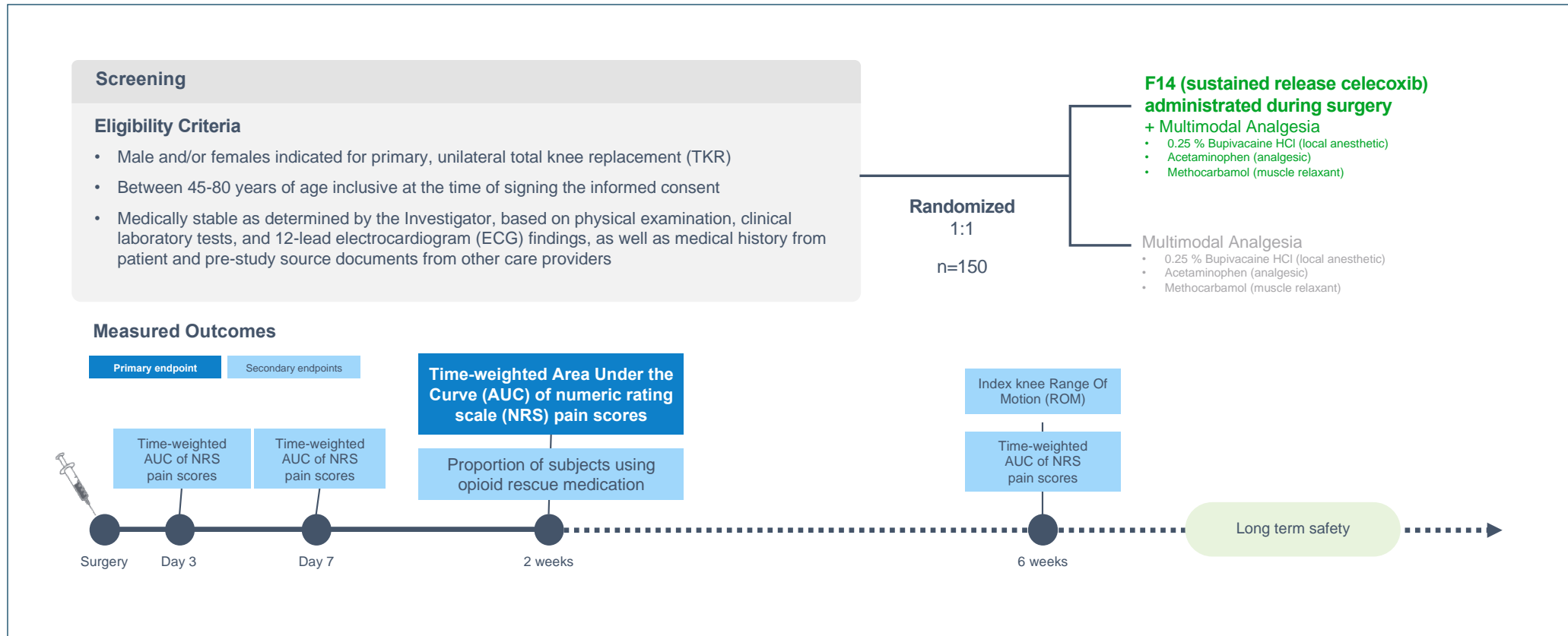
Clinical development in the U.S. led and financed by AIC

Company founded by North American orthopedic surgeons & former biotech CEO

Last private equity financing: CAD\$23 million in February 2021

mdc-CWM, EFFICACY AND SAFETY IN TOTAL KNEE REPLACEMENT

- A Phase 3 Randomized, Double-Blind, Placebo-Controlled Study



Recruitment completion and efficacy data are expected in Q1 2024 and will determine next development steps. Depending on results, at least one additional study will be needed for registration. Regulatory process in pain management typically comprises several efficacy and safety trials to provide convincing evidence of benefit for regulatory agencies.

mdc-CWM, EFFICACY AND SAFETY IN TOTAL KNEE REPLACEMENT

- Phase 3 results (May 2024)

Primary endpoint of time-weighted AUC of pain intensity over 14 days not met

Numerical improvement favoring the treated group observed for

- The primary endpoint
- Secondary endpoints of time-weighted AUC of pain over 3 and 7 days

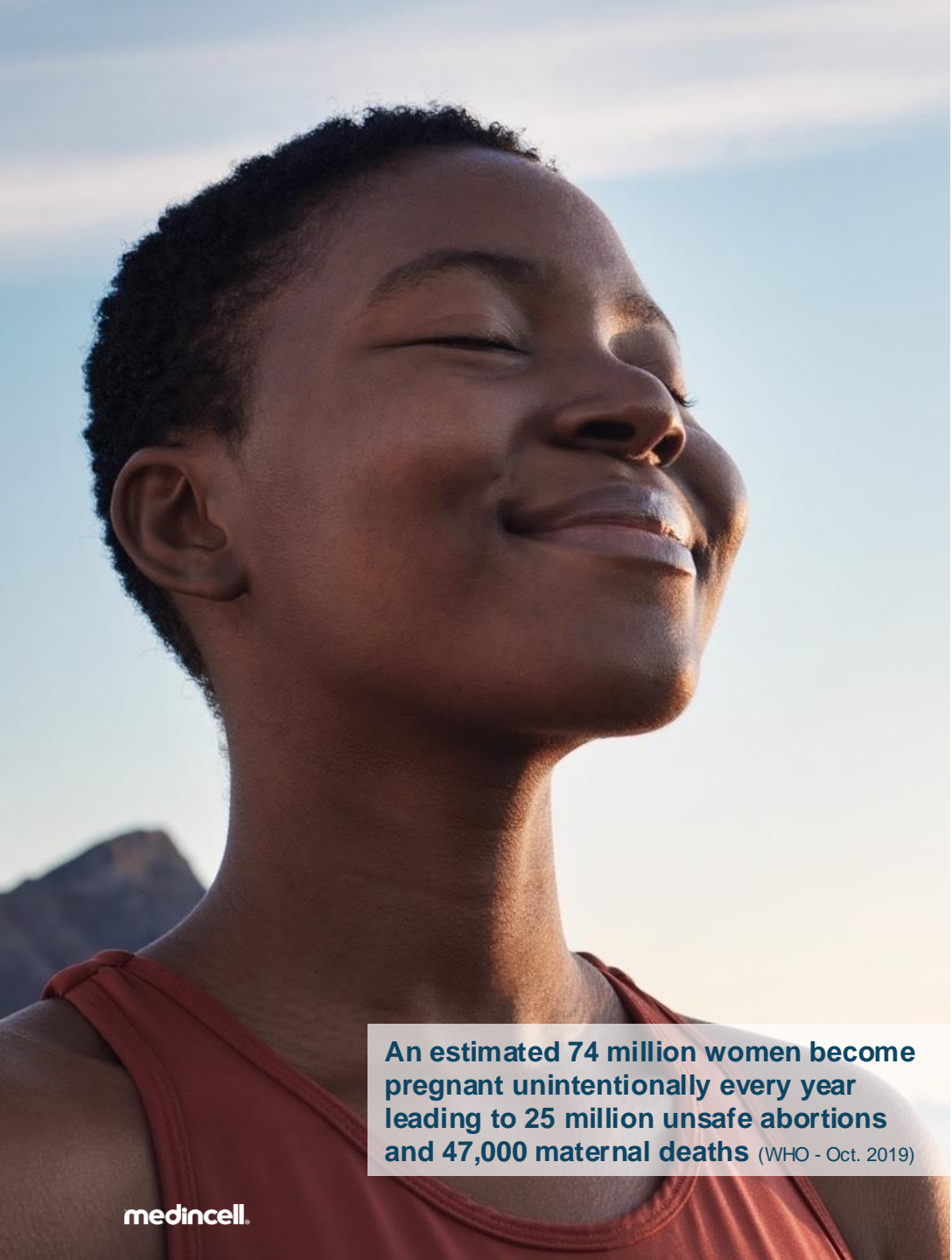
Other positive outcomes related to inflammation

- Improvement for knee range of motion (ROM) at 6 weeks ($p < 0.005$), as well as at 3 months ($p < 0.0005$)
- Improvement for swelling at 6 weeks ($p < 0.005$) and 3 months ($p < 0.05$)
- Improvement of the Timed-Up-and-Go (TUG) test at 6 weeks

Far greater improvement in a sub-group of 108 patients

- Patients had not previously undergone TKR in their contralateral knee
- Improvement in endpoints of time-weighted AUC of pain, opioid consumption, ROM, effusion, and TUG

No new safety signals were identified, and no SAEs were reported as related to F14 treatment



An estimated 74 million women become pregnant unintentionally every year leading to 25 million unsafe abortions and 47,000 maternal deaths (WHO - Oct. 2019)

PRECLINICAL | mdc-WWM

6-Month contraception

mdc-WWM could be the first contraceptive to combine essential features to make it a best-in-class product worldwide

- Progestin molecule (non-MPA)
- 6-month duration
- Subcutaneous injection
- Auto injectable
- Full bio resorption
- Affordability

All commercial rights owned by MedinCell with a significant potential

- Contraception is a \$5bn market in the U.S.
- LARC (Long-Acting Reversible Contraceptives, primarily solid implants and intrauterine devices) represent 28% of US market, i.e., \$1.4bn with 5- CAGR at 7.8% (Source: IQVIA)

Financial support from
BILL & MELINDA
GATES foundation

\$22.5m financing grant by the Bill & Melinda Gates Foundation for Global Access rights in low- and middle-income countries



Malaria in 2020:

- 627,000 deaths
- 95% in Africa,
- 80% children under 5

(WHO)

PRECLINICAL I mdc-STM

Ivermectin / Malaria

Objective: a new tool to fight malaria transmission

- mdc-STM enables sustained release of ivermectin following a single subcutaneous injection
- Administered at beginning of transmission season to people living in malaria-endemic areas
- Mosquitoes feeding on people who have received ivermectin will be killed or made less capable of transmitting malaria parasites further
- Goal is to decrease mosquito numbers, thus benefiting the whole community by lowering the risk of malaria transmission, particularly in children
- Community-based intervention – individuals receiving the injection would not be protected against malaria directly

\$12m financing by the international Health Agency, Unitaid

License agreement with Medicines Patent Pool

Covers all low- and middle-income countries and is royalty free in the public sector. Reasonable royalty in line with industry standards to be agreed in case there would be a private market for the licensed product in low and middle- income countries.



FINANCIALS & EXTRA-FINANCIAL PERFORMANCE

SELECTED FINANCIALS

as of March 31, 2024

€ million	Year end March 31, 2024	Year end March 31, 2023
Operating result	(20.9)	(24.0)
Revenues and other income	11.9	13.7
Operating expenses	(32.9)	(37.7)
Net result	(25.0)	(32.0)
Cash consumption from operating activities	11.9	21.0
Cash position	19.5⁽¹⁾	6.5⁽²⁾

⁽¹⁾ including 5.2 M€ in the form of non-risky financial assets

⁽²⁾ not including 2.8 M€ in short-term investments and 1.1 M€ in non-current financial assets

Balance sheet	Year end	Year end
€ million	March 31, 2024	March 31, 2023
Equity of the consolidated group	(40 824)	(42 294)
Total non-current liabilities	61 304	14 608
Total current liabilities	16 466	57 025
Total non-current assets	9 690	9 772
Of which financial assets and other non-current assets	1 792	1 460
Total current assets	27 258	19 568
Of which cash and cash equivalents	19 460	6 467

Main cash payments received after the closing

- \$35 million from AbbVie (May 2024)



Market Cap: ca. \$500m
as of September 1st, 2024

Outstanding Shares: 29.1M

Analyst coverage (average TP: €20,36)

Jefferies

Brian BALCHIN



Raghuram SELVARAJU



Alex COGUT



Nicolas PAUILLAC



Martial DESCOUTURES



Claire DERAY



Mohamed KAABOUNI

ESG PERFORMANCE

ISS ESG

C+
51.05/100



S&P Global

CSA score: 43 (92nd percentile)
ESG Score: 51 (Average panel pharma 20/100)



Medium Risk (high): 25,9
(rank 68/430°)



C
(Pharma/Biotech benchmark: B-)



80/100



SUSTAINABLE DEVELOPMENT GOALS



"Our mission is to contribute to the improvement and protection of the health of populations across the world.

The fair sharing of the value created with all our employees is the foundation of our business model.

The sustainability of MedinCell is an essential condition for achieving our objectives."

Raison d'être" of MedinCell voted by the General Assembly in September 2019