Issuer Free Writing Prospectus Filed Pursuant to Rule 433 Registration Statement No. 333-269606 Dated April 18, 2023 (To Preliminary Prospectus dated April 4, 2023)

Free Writing Prospectus MAIA Biotechnology, Inc.

This free writing prospectus relates to the proposed public offering of shares of common stock, par value \$0.0001 of MAIA Biotechnology, Inc. (the "Company"), which are being registered on a Registration Statement on Form S-1, as amended (No. 333-269606) (the "Registration Statement"). This free writing prospectus should be read together with the preliminary prospectus dated April 4, 2023 included in that Registration Statement, which can be accessed through the following link:

https://www.sec.gov/ix?doc=/Archives/edgar/data/1878313/000156459023005432/maia-s1a.htm

We have filed the Registration Statement with the Securities and Exchange Commission (the "SEC") for the offering to which this communication relates. Before you invest, you should read the preliminary prospectus in the Registration Statement (including the risk factors described therein) and other documents we have filed with the SEC for more complete information about our Company and this offering. You may access these documents for free by visiting EDGAR on the SEC Web site at http://www.sec.gov. Alternatively, we or any underwriter participating in the offering will arrange to send you the prospectus if you contact ThinkEquity, Prospectus Department, 17 State Street, 41st Floor, New York, New York 10004, telephone: (877) 436-3673 or e-mail: prospectus@think-equity.com.



FREE WRITING PROSPECTUS

This presentation highlights basic information about us and the proposed offering. Because it is a summary, it does not contain all of the information that you should consider before investing. We have filed a registration statement (including a prospectus) with the SEC for the offering to which this presentation relates. The registration statement has not yet become effective. Before you invest, you should read the prospectus in the registration statement (including the risk factors described therein) and other documents we have filed with the SEC for more complete information about us and the offering.

You may access these documents for free by visiting EDGAR on the SEC Web site at http://www.sec.gov. Alternatively, we or any underwriter participating in the offering will arrange to send you the prospectus if you contact ThinkEquity, Prospectus Department, 17 State Street, 41st Floor, New York, New York 10004, telephone: (877) 436-3673 or e-mail: prospectus@think-equity.com.

This presentation shall not constitute an offer to sell, or the solicitation of an offer to buy, nor will there be any sale of these securities in any state or other jurisdiction in which such offer, solicitation or sale would be unlawful prior to the registration or qualification under the securities laws of such state or jurisdiction. The offering will only be made by means of a prospectus pursuant to a registration statement that is filed with the SEC after such registration statement becomes effective.

FORWARD-LOOKING STATEMENTS

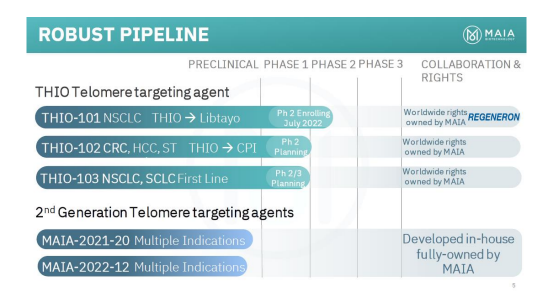
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All statements in this presentation, other than those relating to historical facts, are "roward-looking statements." These forward-looking statements may include, but are not limited to, statements waiting to our objective, and strategies, statements that contain, projections of neurolis of operations, or developments that is inford, espect, project, believe, or adolostat will or may rocur in the fotus. Forward-looking statements or developments that we inford, espect, project, believe, or adolostat will or may rocur in the fotus. Forward-looking statements are strategies, statements that contain projections of an assulptic to this and uncertainties. We have been developments, and other factors they believe to be appropriate. Important factors that could cause actual results, developments, and business decidents of differ materiality from these anticipated in these forward-looking statements induce, among direct hings, the overall global economic environment: general marks, polices, and economic conditions in the unities of the marks and contains envice learnes, and lightion and ergulatory proceedings. The Company has field a registration statement in model (negatation the: 332-259466). Environments for the systemismic to better understand the hisks and uncertainties inherent in our business and industry and for more complete information about as and the offend, "You may get the exploration of the Registration statement. Including the factors are operated or operations. The addition, which we depreter project all comparison operations in the forward-looking statements are objective and industry and for more complete information about as and thereing. You may get there by using EOAR on the Commission to betweeney withing attements are objectives and industry, and second marks and uncertainties in there in any limitation in which we development in the systement are objectives of a subject to operations. There is a uncertainties in the results are adoloted and the systement and busintes of the development of the i

INVESTMENT OVERVIEW

- Telomere-Targeting Agents:
 - \circ THIO in clinic
 - Advancing pipeline
- Efficacy
- Safety
- FDA: 2 Orphan Drug Designations
- REGN: Clinical Supply Agreement
- Phase 2 THIO-101 trial in NSCLC underway
 - Enrolling in AUS and EU
 - $\circ~$ On track to open sites in US in 2023
 - Upcoming Milestones: Safety, ORR, DoR
- Phase 2 THIO-102 basket/umbrella trial in 2023
- THIO-103 basket trial in 2023





SCIENCE OVERVIEW





THIO (6-thio-dG)

Telomere Targeting Agent

- Small molecule (penetrates blood-brain barrier)
- Eligible for NCE marketing exclusivity
- Dual MoA: telomere targeting + immunogenic
- Complete Response with No Recurrence in vivo in Lung, Colorectal, Liver, Melanoma, Brain Cancer (GBM, DIPG, MB), etc
- FDA Orphan Drug Designations: HCC and SCLC

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Next Generation

Telomere Targeting Candidates

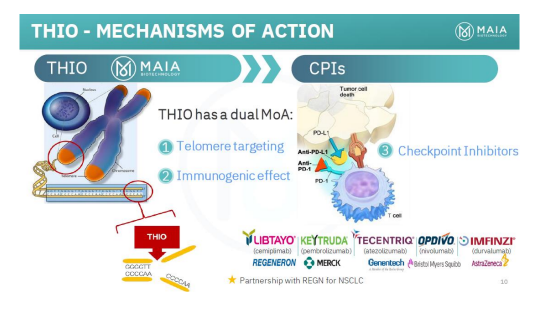
- Similar MoA
- Structures: evolution of THIO; other new structures
- Objective: advance to pre-IND development one agent every 12 months





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THIO is the only direct telomere targeting agent currently in clinical development



REGENERON AGREEMENT

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MAIA Biotechnology, Inc. Announces Clinical Supply Agreement with Regeneron for Phase 1/2 Clinical Trial Evaluating THIO in Sequential Administration with Libtayo[®] (cemiplimab) in Advanced Non-Small Cell Lung Cancer



CLINICAL DEVELOPMENT OVERVIEW

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THIO-101

Ph 2 trial THIO → LIBTAYO®

- Go-to-Market
- NSCLC Second Line
- REGN supply agreement
- Enrolling at multiple sites in AUS and EU (2022)
- File US IND and commence enrolling in US in 2023
- Select optimal dose and expand
- File for accelerated approval (2025)

THIO-1027 PA

Ph 2 trial THIO → CPIs

- Go-to-Market
- CRC, HCC, ST
- Select most efficacious combination with 3 CPIs
- 9+ possible market entry indications
- US, EU, Asia (2023)
- File for accelerated approval (2026)

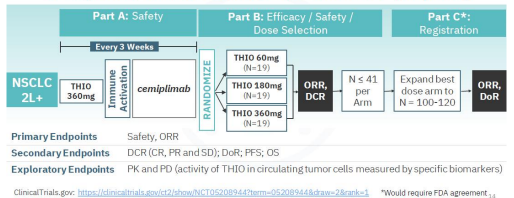
THIO-103

Ph 2/3 trial of THIO → CPIs

- Confirmatory study for accelerated approvals
- First Line NSCLC, SCLC to start
- Market Expansion
- 9+ tumor types
- First approvals in additional tumor types / global markets

THIO-101 TRIAL (ONGOING)

A Multicenter, Open-Label, Dose-Finding Phase 2 Trial Evaluating the Safety and Efficacy of THIO Administered in Sequence with LIBTAYO[®] (*cemiplimab*)



FAVORABLE SAFETY PROFILE

- · Safety events reported during dose-limiting toxicity window
- 360 mg/cycle THIO highest dose
- Data from 6 patients who completed the dose-limiting toxicity (DLT) period in Cycle 1 (3 weeks)
- No Serious Adverse Events (SAE) or Serious Unexpected Suspected Adverse Reactions (SUSAR)
- Safety profile substantially better than current standard of care
- Chemotherapy has 70-80% incidence of grade 3-4 very severe side effects
- Started Part B (efficacy/dose selection) of the trial upon recommendation by the Safety Review Committee

FAVORABLE SAFETY PROFILE

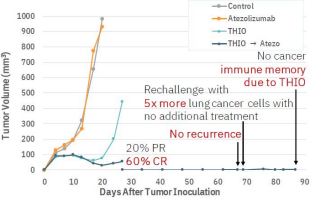
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Adverse events (AE) reported – DLT window (3 weeks)	Grade
Fatigue	1
Decreased appetite	1
Blood pressure fluctuation	1
Dyspnea	1
Nausea	1
Interleukin-6 (IL-6) level increased*	1
Rash erythematous	1
Constipation	1
Myalgia	1
Vomiting	2
Nausea**	3

*T-cell activation **Resolved within 72 hours with treatment

THIO-101 - RATIONALE

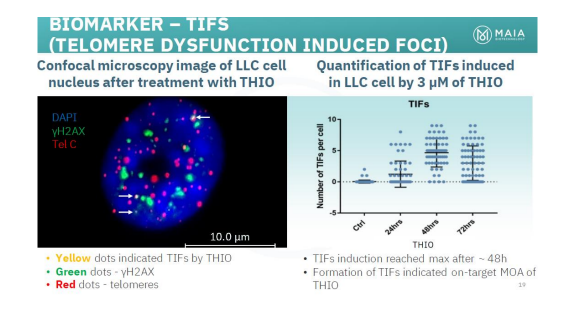
- THIO followed by CPI results in 60% complete response
- No recurrence after long-term
- No recurrence after long-term follow-up
 Anticancer immune memory has been induced: no cancer after rechallenge with 5x more lung cancer (LLC) cells with no additional therapy



Mender et al, Cancer Cell, 2020; THIO followed by Tecentriq (atezolizumab; Roche/Genentech) tested first; repeated later with THIO followed by Keytruda (pembrolizumab; Merck); and Libtayo (cemiplimab; Regeneron) 17

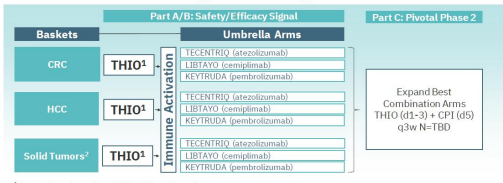
THIO-101 – NON-SMALL CELL LUNG CANCER 🔞 MALA

• Target: 20% improvement on Standard of Care (SOC)	Catalyst	Timing	Current SoC (Chemo)
	Toxicity	Q1 2023	72-79% Grd 3-4
	ORR (Overall Response Rates)	2023	11-23%
	CR (Complete Response)	2023	0%
	DoR (Duration of Response), PFS (Progression Free Survival)	2024	4-4.5m
	OS (Overall Survival)	2025	8.1-10.5m
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THIO-102 TRIAL (PLANNED)

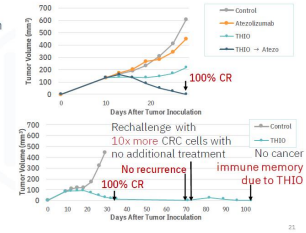
A Multicenter, Open-label, Phase 2 Trial Evaluating the Safety and Efficacy of THIO Administered in Sequence with Anti-PD-1 or Anti-PD-L1



¹ Dose to be selected from THIO-101 study results ² E.g. Breast, Prostate, Gastric, Pancreatic, Ovarian, etc

THIO-102 TRIAL - COLORECTAL RATIONALE 🔞 MALA

- THIO followed by CPI results in 100% complete response
- No recurrence after long-term follow-up
- Anticancer immune memory has been induced: no cancer after rechallenge with 10x more CRC cells with no additional therapy



Mender et al, Cancer Cell, 2020

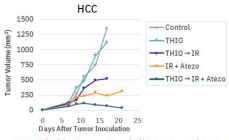
THIO-102 TRIAL - COLORECTAL

• Target: 20% improvement on Standard of Care (SOC)	Catalyst	Timing	Current SoC (Chemo)
	Toxicity	2024	50-60% Grd ≥ 3
	ORR	2024	1-1.6%
	DoR, PFS	2025	1.9-2.0m
	OS	2026	6.4-7.2m

SCLC & HCC - ORPHAN DRUG DESIGNATION MALA

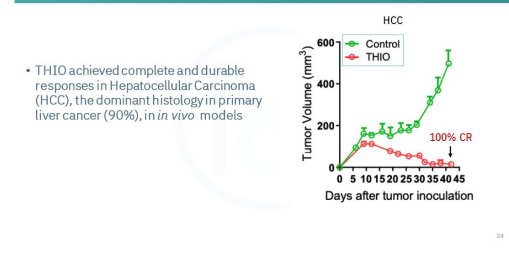


- THIO is synergistic with anti-PD-1 agent Pembrolizumab in Small Cell Lung Carcinoma (H2081) in vivo in humanized murine cancer model.
- Treatment with THIO followed by Pembrolizumab results in highly potent anticancer effect, as compared to Pembrolizumab alone.
- THIO converts immunologically "cold non-responsive" SCLC tumor into "hot and responsive" to Pembrolizumab.



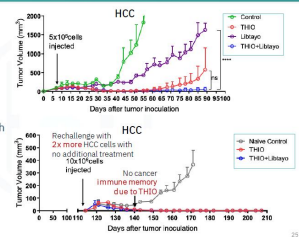
- THIO is highly synergistic and effective in combination with anti-PD-L1 agent Atezolizumab and Ionizing Radiation (IR 10Gy) in HCC53N Hepatocellular Carcinoma.
- Treatment with THIO in combination with IR and Atezolizumab results in a complete regression of aggressive HCC tumors. The combination of IR and Atezolizumab is just partially efficacious.
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EXCELLENT EFFICACY IN HCC MODELS



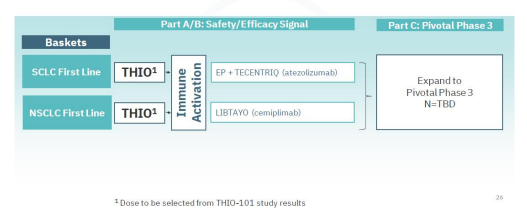
HCC ANTI-CANCER IMMUNE MEMORY

- When combined with immunotherapy checkpoint inhibitor Libtayo[®], duration of response was further potentiated
- Upon rechallenge with two times more cancer cells and no additional treatment, tumor growth was completely prevented
- Administration of THIO alone and in combination with Libtayo[®] generated anti-cancer immune memory



THIO-103 TRIAL (PLANNED)

A Multicenter, Open-label, Phase 2 Trial Evaluating the Safety and Efficacy of THIO Administered in Sequence with Anti-PD-1 or Anti-PD-L1



EXCLUSIVITY AND INTELLECTUAL PROPERTY M MALA

Goal: New Chemical Entity (NCE) Marketing Exclusivity

- THIO has never been previously approved by the FDA for commercialization
- Robust exclusivity
- US: 7 years; EU, Japan, other markets: 10 years

Robust and Growing Patent Portfolio for THIO

- 1 issued US patent
- 4 issued foreign patents
- 5 pending US patent applications
- 7 pending foreign patent applications

Current patents/provisional applications broadly cover the following key areas:

- Telomere targeting compounds (2034+)
- THIO's immunogenic treatment strategy: sequential combination with CPIs (2041)



EXPERIENCED MANAGEMENT TEAM



CAPITALIZATION TABLE & CASH BALANCE MAIA BIOTECHNOLOGY

Capitalization Table (as of 12/31/2022)					
Common stock	10,955,904				
Options (WAEP: \$2.55) ¹	6,545,628				
Warrants (WAEP: \$6.04)	796,985				
Fully Diluted Shares Outstanding	18,298,517				
Cash Balance of \$10.95 million (as of 12/31/2022)					

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¹ 4,282,309 options held by directors and officers **Note:** Directors and officers, and their affiliates, own 44% of the 18,298,517 fully diluted shares outstanding



SIGNIFICANT MARKET OPPORTUNITY

MAIA



COMPARABLE COMPANIES

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• On June 3, 2022, Bristol Myers Squibb announced the acquisition of Turning Point Therapeutics in an all-cash transaction for <u>\$4.1B</u> in equity value.



Market Caps as of April 18, 2023 (source: S&P CapitalIQ)

MULTIPLE VALUE-DRIVING MILESTONES



