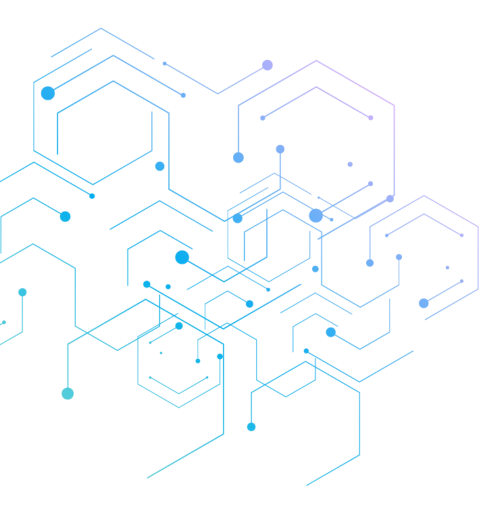


Transforming Lives

Glycobiology-based Therapeutics

May 2024 | NASDAQ: GLYC



Forward-Looking Statements

- To the extent that statements contained in this presentation are not descriptions of historical facts, they are forward-looking statements reflecting the current beliefs and expectations of the management of GlycoMimetics, Inc. ("GlycoMimetics," "we," "us," or "our"). Forward-looking statements contained in this presentation may include, but are not limited to: (i) the expected or projected timing of events, data readout and data analysis from clinical trials; (ii) the planned or potential clinical development and potential indications, benefits and impact of our drug candidates, including uproleselan and GMI-1687; (iii) the timing of receipt of clinical data; (iv) the potential safety, efficacy or clinical utility of our drug candidates; (v) the size of patient populations targeted by drug candidates we or our collaborators develop; (vi) the likelihood and timing of regulatory filings, and plans for interactions with regulatory authorities; (vii) our business and product development strategies, including our cash needs and expected cash runway; and (viii) any other statement containing terminology such as "may," "will," "should," "expects," "plans," "anticipates," "believes," "estimates," "predicts," "potential," "intends," or "continue," or the negative of these terms or other comparable terminology.
- Forward-looking statements are subject to known and unknown risks, uncertainties, and other factors that may cause our or our industry's actual results, levels of activity, performance, or achievements to be materially different from those discussed, implied or otherwise anticipated by such statements. You are cautioned not to place undue reliance on such forward-looking statements, which are current only as of the date of this presentation. Examples of risks, uncertainties and factors that may cause differences between our expectations and actual results include unexpected safety or efficacy data, unexpected side effects observed during preclinical studies or in clinical trials, whether results of early clinical trials will be indicative of results from later clinical trials, changes in expected or existing competition or additional market research that may cause our expectations about market opportunity to change, changes in the regulatory environment for our drug candidates, failure of our collaborators to support or advance our collaborations or drug candidates, our need for future capital, the inability to adequately protect our intellectual property, and becoming a party to litigation or other disputes. For a further description of the risks associated with forward-looking statements, as well as other risks facing GlycoMimetics, please see the risk factors described in the Company's Annual Report on Form 10-K filed with the U.S. Securities and Exchange Commission on March 27, 2024, as well as other reports we file with the U.S. Securities and Exchange Commission from time to time, including those factors discussed under the caption "Risk Factors" in such filings. Forward-looking statements speak only as of the date of this presentation, and GlycoMimetics undertakes no obligation to update or revise these statements, except as may be required by law.



Near-Term Catalysts and Promising, Glycobiology-based Pipeline

Uproleselan: Multiple Late-Stage Clinical Trials

- **Phase 3 trial** in R/R AML (n=388), topline results announced in Q2 2024
- Fully enrolled Phase 2 trial in front-line AML (n=267) ongoing, NCI-sponsored
- Ongoing IITs in other AML populations.
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 2022/2023
- Novel MOA/first-in-class → potential broad utility with Breakthrough Therapy, Fast Track, and Orphan designations

Promising Early-Stage Pipeline

- Novel small molecules inhibit carbohydrate signaling
- Potential application in multiple inflammatory diseases
- GMI-1687
 - Phase 1a trial in healthy volunteers completed
 - Initial indication: treatment of sickle cell disease (SCD) vaso-occlusive crisis (VOC)
 - Being developed for self-administration at time of VOC
- Galectins
 - Targeting fibrotic diseases
 - First oral Galectin-3 antagonist

Targeted Operational Execution

• Updating uproleselan plans and evaluating financial guidance



A Portfolio of Promising Product Candidates

Program	Therapeutic Area	Discovery	Preclinical	Phase 1	Phase 2	Phase 3	Market
SELECTINS							
	Relapsed / Refractory AML	Topline results	announced in Q2 2	024			
UPROLESELAN (GMI-1271)*	Newly Diagnosed "Fit" AML	Fully enrolled 2	67 patients Dec 20	21			
	Relapsed / Refractory Pediatric AML	Ph1 by NCI dos	sed 1 st patient				
GMI-1687 *	SCD Vaso-occlusive Events and Inflammatory diseases	Ph1a complete	d				
GALECTINS							
GMI-2093	Fibrosis and Oncology	Lead declared N	larch 2022				
	und with Annulancing in Operator Obier						

Breakthrough Therapy Designation in AML

Uproleselan (GMI-1271)

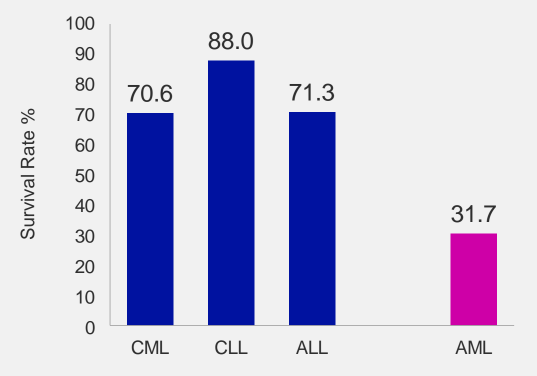




Significant Unmet Medical Need In AML¹

ESTIMATED NEW CASES (2023) 20,380 New AML Cases All Other 21,450 New Leukemias Cases

5-YEAR RELATIVE SURVIVAL (2013 – 2019)¹



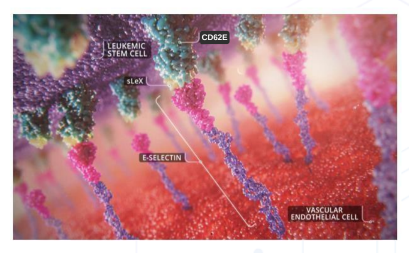
American Cancer Society. Cancer Facts and Figures 2023. Atlanta: American Cancer Society; 2023. Accessed May 10, 2023. https://www.cancer.org/content/dam/cancer-org/research/cancer-facts-and-statistics/annual-cancer-facts-and-figures/2023/2023-cancer-facts-and-figures.pdf.

Uproleselan: First-in-Class E-Selectin Antagonist for AML



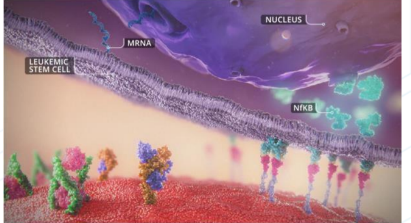
E-selectin:

- Leukocyte adhesion molecule constitutively expressed on marrow endothelial cells, also inducibly expressed throughout vasculature by innate inflammatory mediators
- ✓ Up-regulated by AML blasts via secreted inflammatory mediators, such as TNF-alpha and IL1-beta



E-selectin/E-selectin Ligand Interaction:

- Enables AML blast and leukemia stem cell sequestration in bone marrow
- ✓ Activates pro-survival NF-kB pathways
- E-selectin ligand sLex up-regulated on AML cells via multiple distinct drug resistance mechanisms



Uproleselan, a First-in-class E-Selectin Antagonist:

- Releases AML blasts and leukemic stem cells from vascular sequestration, agnostic to AML mutational status
- ✓ Disrupts NF-kB mediated chemoresistance pathways
- ✓ Potential broad utility across AML



Phase 1/2 Results in R/R and Newly Diagnosed AML Patients

AML population	CR	CR/CRi	Median O/S	MRD- negative
Relapsed / Refractory (n = 54)	35%	41%	8.8 mos	69%
Newly Diagnosed (n = 25) >=60yrs	52%	72%	12.6 mos	55%

E-selectin ligand expression

- Detectable in every patient tested
- Higher levels in R/R patients achieving CR/CRi, MRD- and prolonged median OS

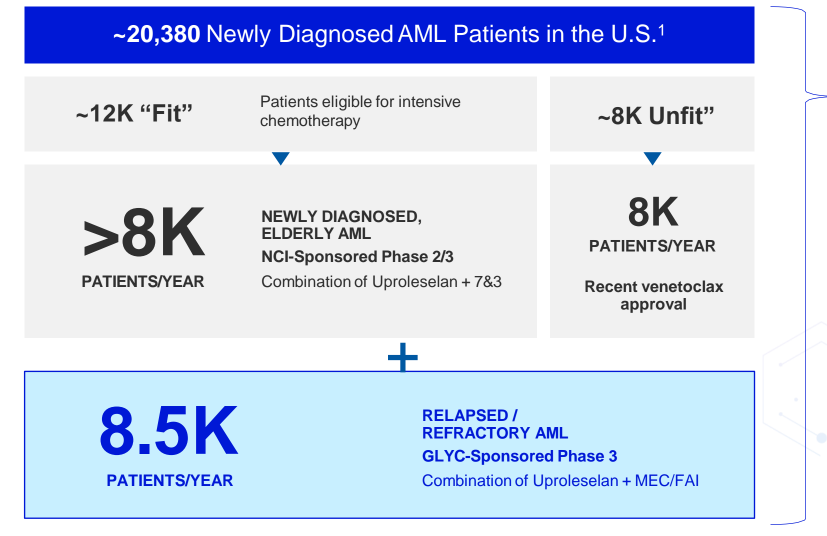


80% 70% 60% 50% 40% 30% 20% 10% 0% R/R AML (N=16) Newly Dx AML (N=9)

Percent MRD Negative



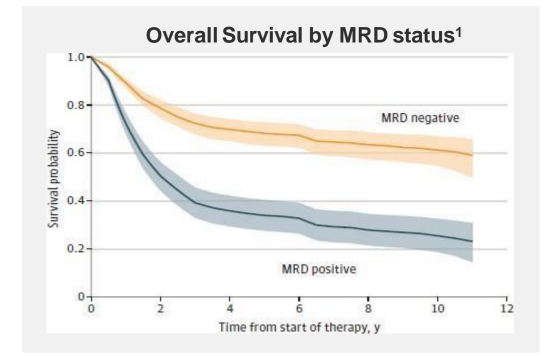
Potential Foundational Backbone Across Spectrum in AML



Uproleselan Value Proposition

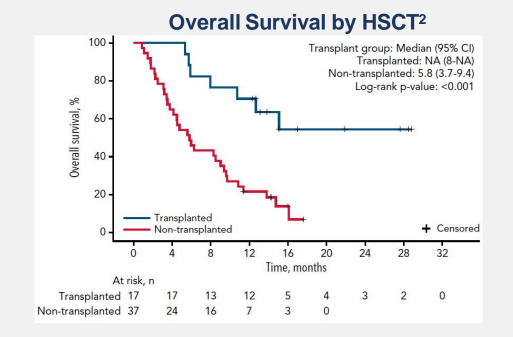
- Improve achievement / depth of remission
- Extend overall survival
- Mitigate chemotherapyrelated toxicity

MRD Negativity and HSCT Both Favorably Prognostic



Meta-analysis of 81 studies (N >11,000)

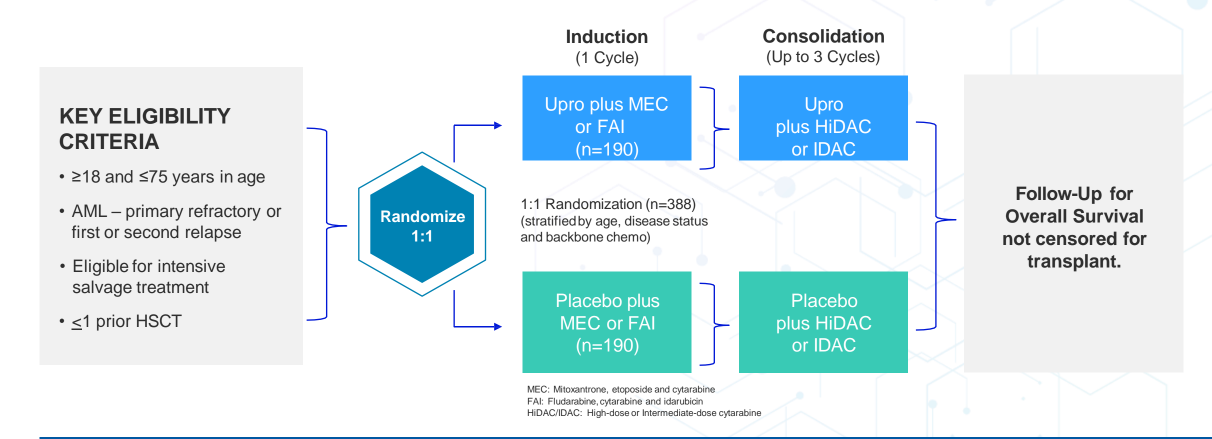
- MRD negativity favorably prognostic for survival
- Effect independent of age, subtype, timing, method



Uproleselan Phase 1/2 overall survival by HSCT

- N=54 R/R AML patients at 10 mg/kg RP2D
- Overall MRD-negative: 56% 1L, 69% R/R
- 10 longest survivors all MRD-negative

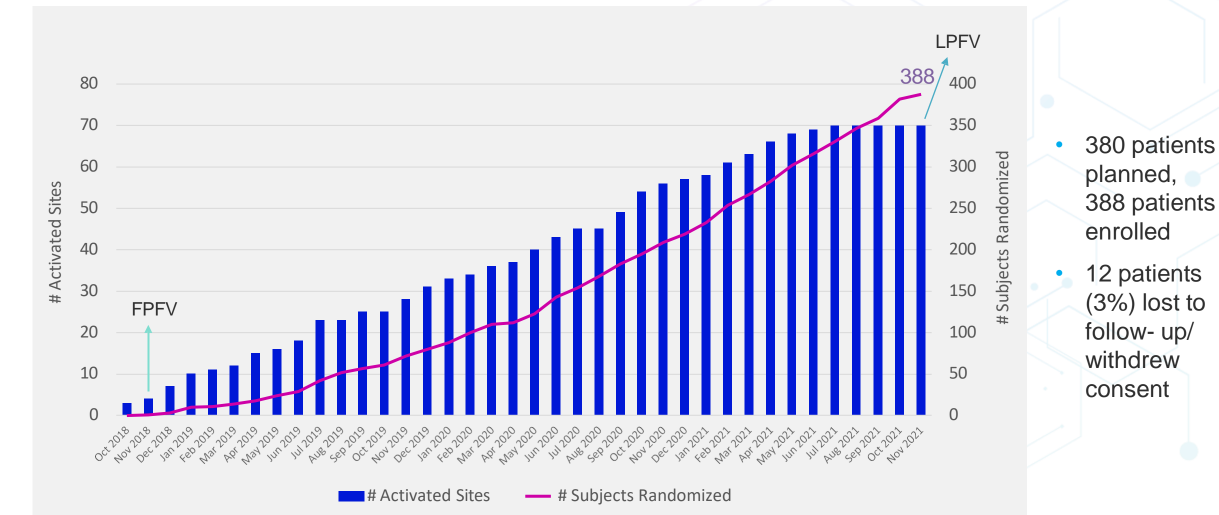
Phase 3 Global Pivotal Study of Uproleselan in R/R AML



Primary endpoint of overall survival was not achieved Median overall survival: 13 months (uproleselan) vs. 12.3 months (placebo arm) Adverse events consistent with known side effect profiles of chemotherapy used in the study Comprehensive analysis ongoing; plan to submit for presentation at an upcoming medical meeting

GlycoMimetics

Trial GMI-1271-301 Enrollment





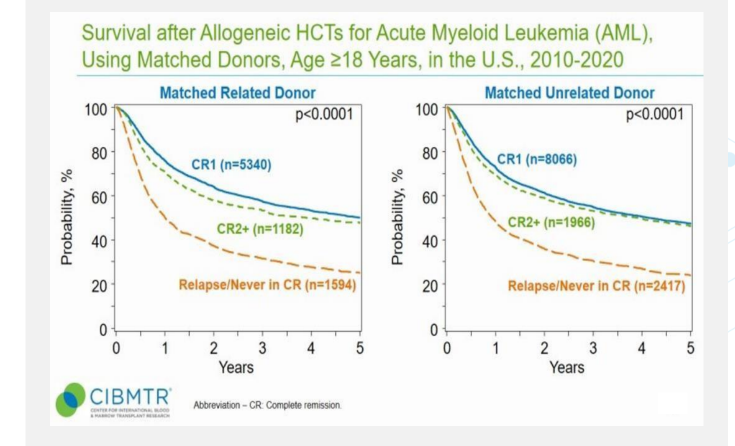
Phase 3 Patient Characteristics Broadly Similar to Phase 2

	301 Study N=388	201 Study N=66			
Relapsed/Refractory Patient Demographics					
Age, median (range)	58 (20-75)	59 (26-84)			
Refractory, n (%)	129 (33%)	22 (33%)			
Relapsed, n (%)	259 (67%)	44 (67%)			
Duration of prior remission ≤6 mos	56 (22%)	18 (41%)			
Prior Therapies					
HSCT	70 (18%)	12 (18%)			
≥2 Induction Regimens	63 (16%)	22 (33%)			
ELN Risk Category					
Adverse	42%	50%			
Intermediate	23%	17%			
Favorable	21%	11%			
Unknown	14%	22%			

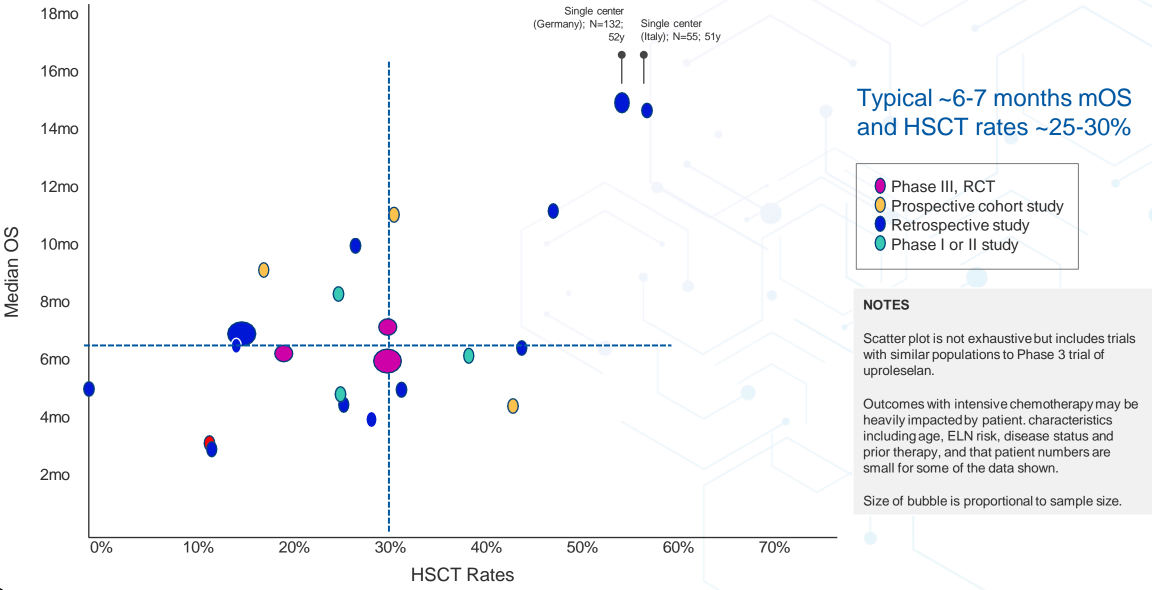


Time-Based Analysis Triggered with a March 31, 2024 Data Cutoff

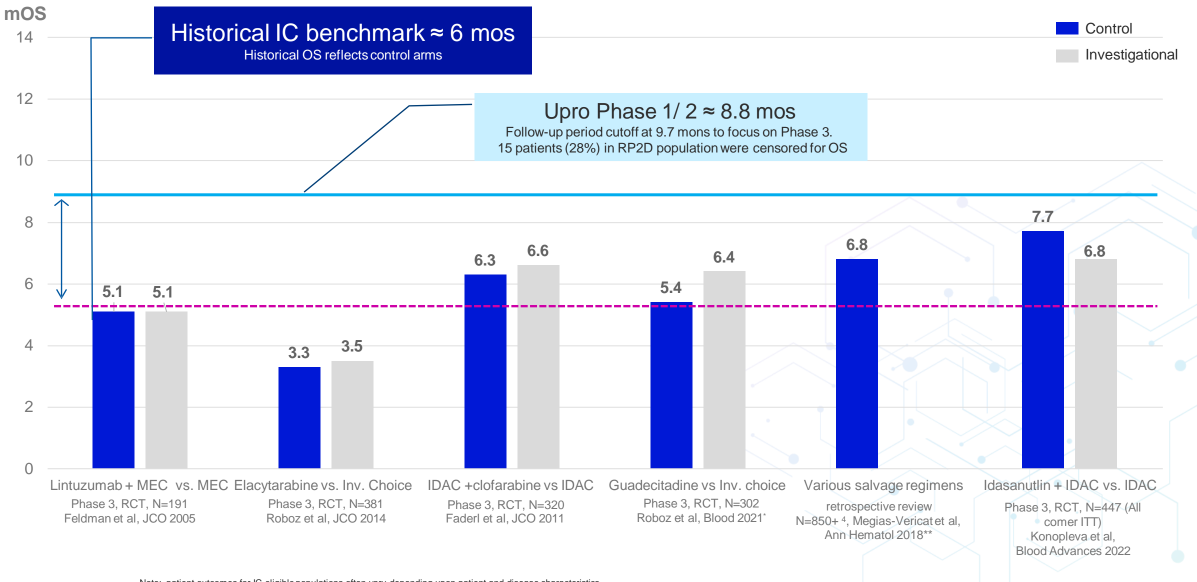
- June 2023 FDA cleared Phase 3 timebased OS analysis after defined cutoff if 295 events not reached by that date
- Clinically mature data in Q2 2024 reflects > 3 years median follow-up and > 2 years post-transplant followup for the substantial majority of remaining patients that received stem cell transplants
- After 2 years post-transplant, AML relapse becomes infrequent



Intensive Chemotherapy (IC) in R/R AML

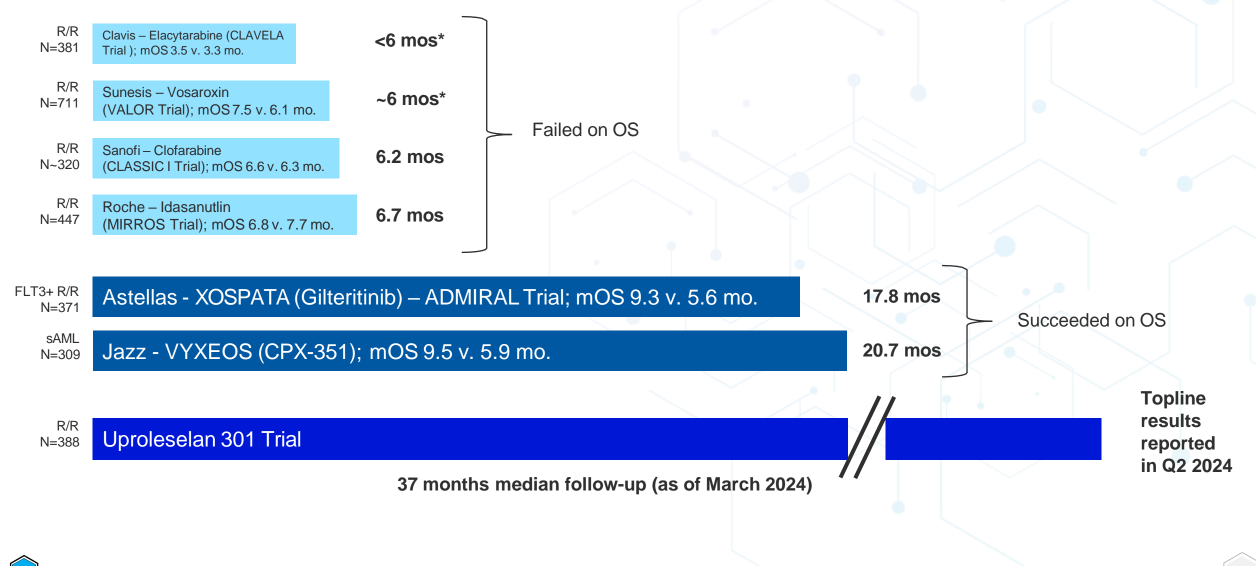


Historical Intensive Chemotherapy benchmarks for mOS are ~6 months





Duration of Follow-Up and Outcomes in Key AML Trials

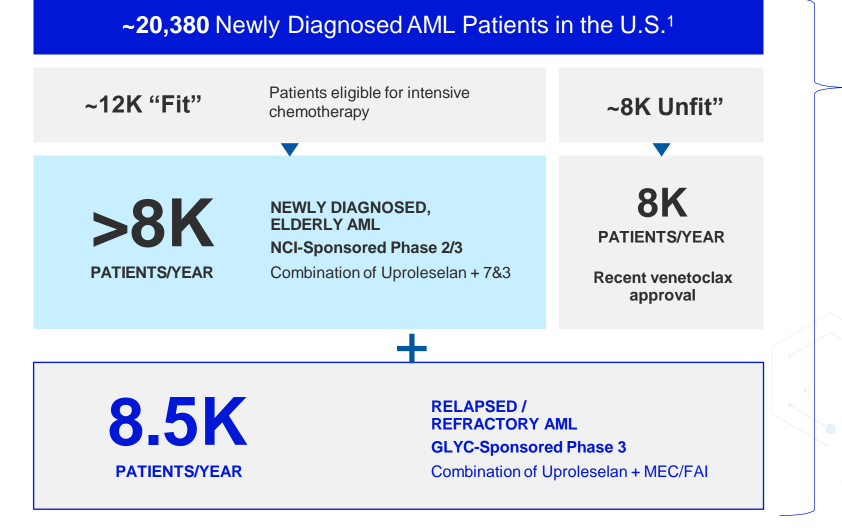


Follow-Up Versus Outcome in Select AML Trials

Trial	Median Survival (mos)	Median Follow-up (mos)	Enrolled (N)	Planned Events	OS HR	P-value
CLAVELA	3.5 vs 3.3 mos	< 6*	381	302	0.97	0.96
VALOR	7.5 vs 6.1 mos	~ 6*	711	562	0.87	0.0610
CLASSIC I	6.6 vs 6.3 mos	6.2	320	258	1.00	1.00
MIRROS	6.8 vs 7.7 mos	6.7	436	296	1.09	0.52
VIALE-A	15 vs 10 mos	20.5	433	270	0.66	< 0.001
VYXEOS	9.6 vs 6.0 mos	20.7	309	236	0.69	0.003
ADMIRAL	9.3 vs 5.6 mos	17.8	371	258	0.64	< 0.001
Uproleselan	13 vs.12.3 mos	37 (Mar '24)	388	295	TBD	TBD

Glyco Mimetics * Median follow-up at time of event trigger for CLAVELA and VALOR estimated from protocol and/or final results as it was not included in the publication

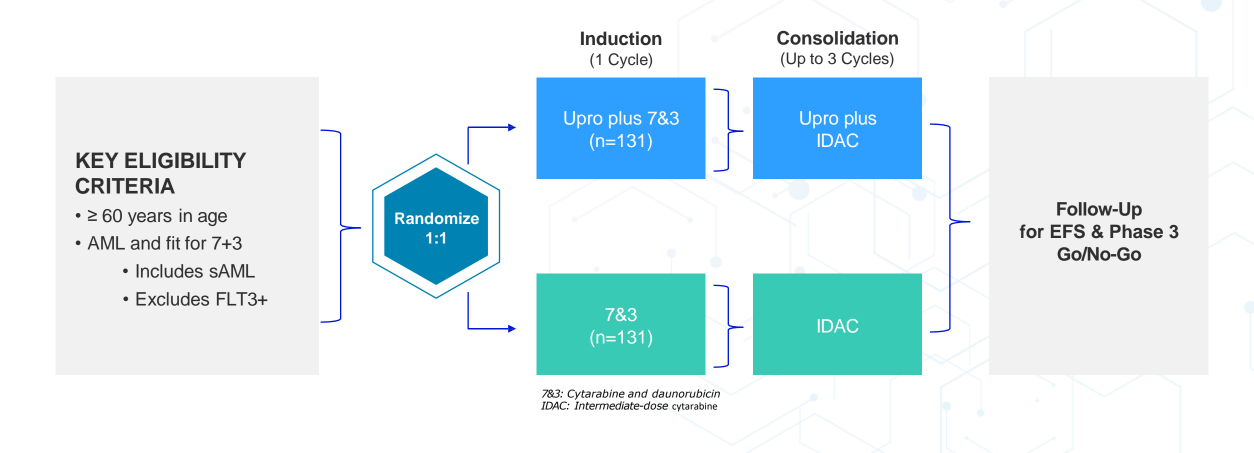
Potential Foundational Backbone Across Spectrum in AML



Uproleselan Value Proposition

- Improve achievement / depth of remission
- Extend overall survival
- Mitigate chemotherapyrelated toxicity

NCI / Alliance Frontline "Fit" AML Phase 2/3 Trial Design

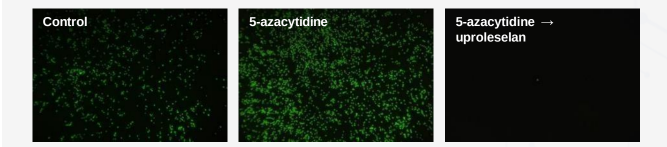


Enrollment of 267 Patients in Phase 2 Portion Completed in December 2021



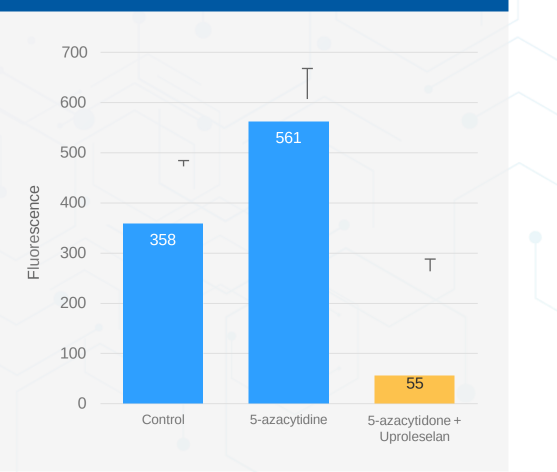
HMA Resistance is Driven by E-selectin, Broken by Uproleselan

UPROLESELAN INHIBITS BINDING OF BLASTS



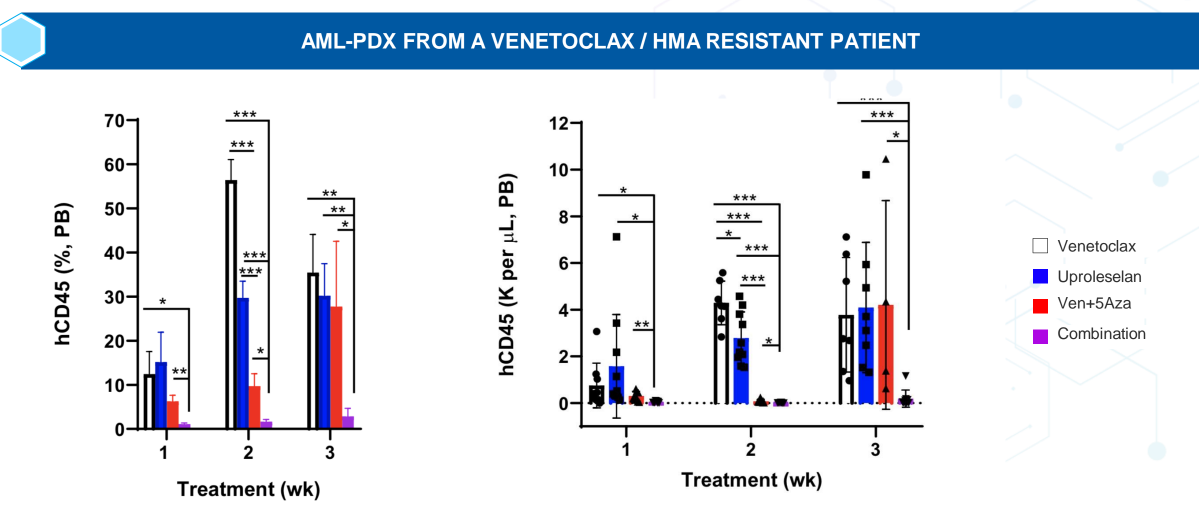
KG1 AML cells were incubated for 96 hours in the absence or presence of 100 nM 5-azacytidine, labeled with calcein and allowed to adhere to E-selectin coated plates (control and 5-azacytidine above). After 45 minutes of adhesion, Uproleselan was added to the wells and fluorescence determined after 30 minutes

(5-azacytidine \rightarrow Uproleselan above).





Targeting E-selection with GMI-1271 Overcomes Microenvironment-mediated Resistance to Venetoclax/HMA Therapy K.H. Chang, M. Muftuoglu, W.Zhang, M. Basyal, L. Ostermann, W.E. Fogler, J.L. Magnani, M. Andreeff, 2020 Uproleselan/ Venetoclax/ HMA Combination Significantly Reduces Leukemia Burden, Compared to Ven+5Aza Alone¹



*p< 0.05; **p<0.01; ***p<0.001, Student's t-test for experiments that compare two groups.

ASH 2022/2023: First Clinical Uproleselan Data Generated Outside of GLYC-Sponsored Trials

Uproleselan data from two investigator-initiated trials presented at ASH in December 2022/2023

A Phase I Study of Uproleselan Combined with Azacitidine and Venetoclax for the Treatment of Older or Unfit Patients

with Treatment Naïve Myeloid Leukemia B.A. Jonas, J.L. Welborn, N.S. Esteghamat, R.T. Hoeg, A.S. Rosenberg, L. Molnar, A. Linh Dang-Chu, S.L. steward, and J.M. Tuscano, 2022

Publication Number: 2764

Encouraging safety and evidence of disease activity

- 8 evaluable patients with poor prognosis
 - 6/8 (75%) were ELN 2017 adverse risk disease
 - 3/8 (38%) had complex cytogenetics
- Data outcomes
 - 6/8 (75%) CR/CRi
 - 5/8 (63%) full CR
 - 1/8 (13%) CRi
 - 5/8 (63%) CR/CRi responses occurred with cycle 1
 - 4 CR/CRi MFC MRD negative
 - 50% overall MRD negative rate
 - 67% among CR/CRi responders

Uproleselan added to Cladribine Plus Low Dose Cytarabine (LDAC) in Patients with Treated Secondary Myeloid

Leukemia (TS-AML) E.A. Huante, H. Kantarjian, K.S. Chien, C.D. DiNardo, N. Short, A. Maiti, G. Montalban, N. Daver, J.D. Kawedia, K. Bowie, S.A. Pierce, F. Ravandi, M. Konopleva, G. Garcia Manero, and T. M. Kadia, 2023

Publication Number: 2992

39% ORR in very high-risk patient population

- 18 evaluable patients
 - All patients had unfavorable cytogenetics and had previously received treatment with a hypomethylating agent.
 - 11 patients (55%) had received prior treatment with venetoclax, and five (25%) had undergone stem cell transplantation.
- Data outcomes
 - Combination of Cladribine + LDAC with uproleselan overall well tolerated with few treatment-related AEs
 - Combination reduced bone marrow blasts in 13 (72%) patients
 - Three patients went on to receive a potentially curative hematopoietic cell transplantation (HCT)
 - Study investigators concluded data support this low-risk approach to marrow blast reduction and disease control in preparation for HCT



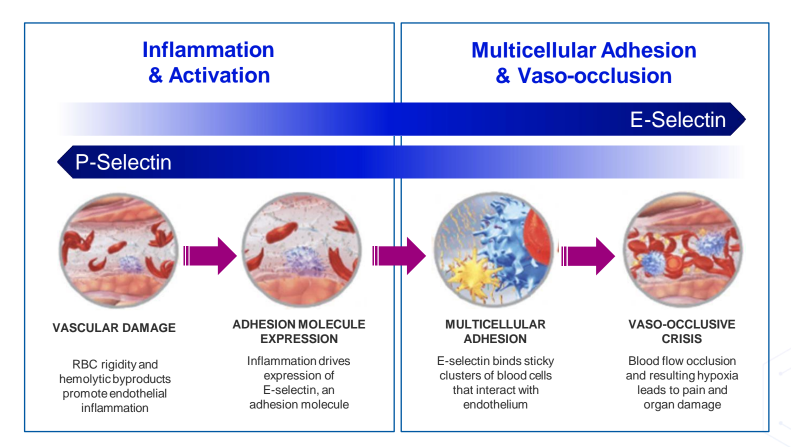
Treatment of Vaso-occlusive Crisis (VOC) in Patients with Sickle Cell Disease

GMI-1687





E-Selectin Mediates Multicellular Adhesion and Vaso-Occlusion



Data Supporting E-Selectin Role in Cellular Adhesion and Clotting

Preclinical

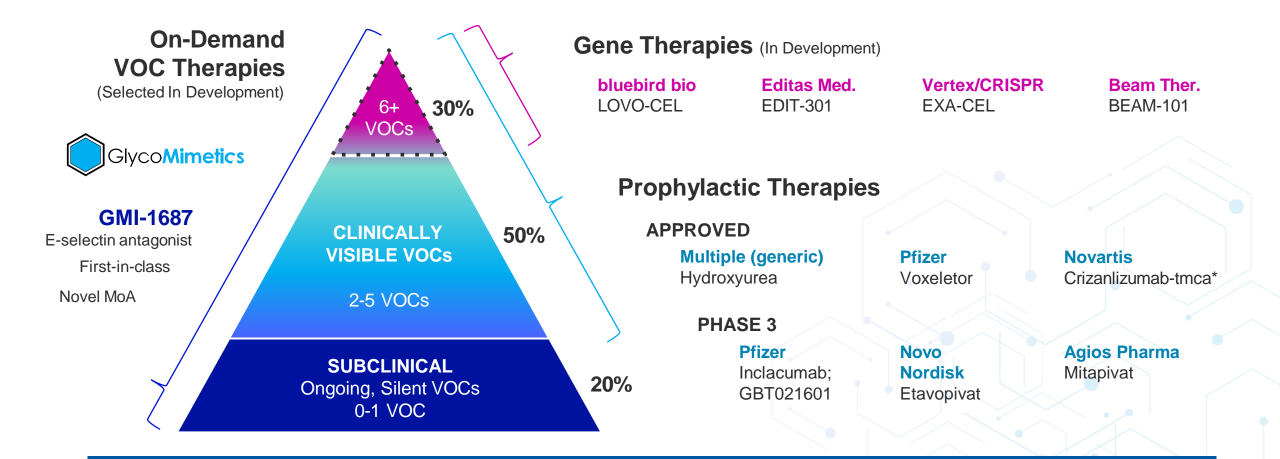
- · E-selectin leads to rolling and cell arrest
- Blocking E-selectin inhibits leukocyte
 adhesion
- Blocking E-selectin restores blood flow in animal models of vessel occlusion in sickle cell disease

Clinical

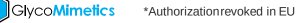
- sE-selectin correlates with frequency of VOC
- · sE-selectin correlates with poor survival
- Reduced sE-selectin correlated with clinical benefit in RESET trial (time to discharge)

E-selectin Antagonism Provides a Unique Therapeutic Target to Interrupt VOC in SCD patients

Even with Prophylactic and Gene Therapy Approaches, VOC Will Remain A Significant Unmet Medical Need

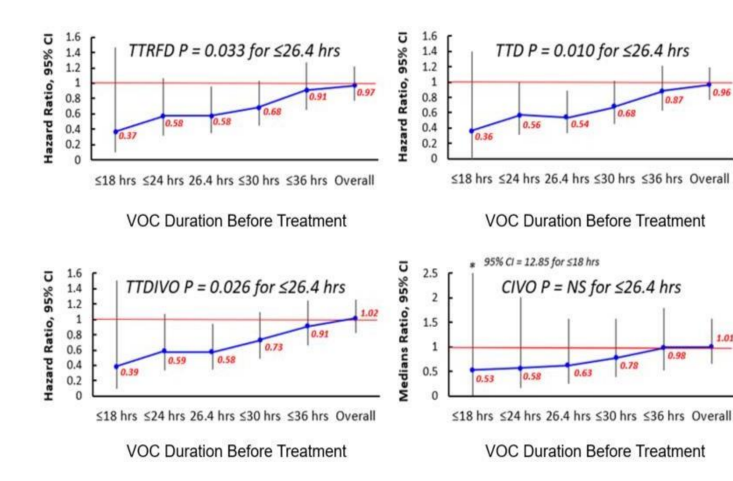


220,000 – 450,000 VOCs/year (in the era of prophylactic therapies)

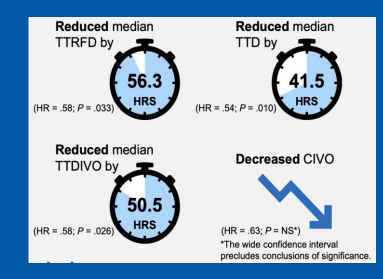


Dampier et al. 2017 American Society of Hematology Annual Meeting. Abstract# 4660 N Engl J Med 2019; 381:509-51; N Engl J Med 2017; 376:429-439

Early Treatment Resulted in Clinical Benefit



For patients treated within first quartile of treatment (<26.4hrs), a meaningful, statistically significant benefit was seen across study endpoints

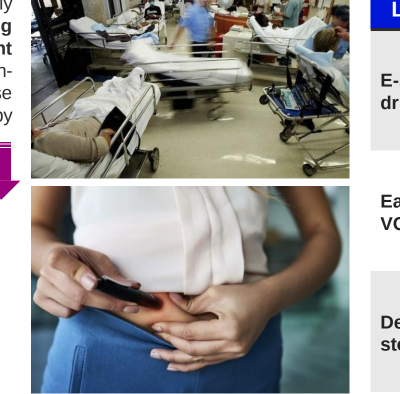


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GMI-1687 Seeks to Empower Patients to Take Control

Potentially revolutionizing the treatment paradigm to ondemand disease modifying therapy



Lessons Learned	GMI-1687
E-selectin drives VOC ¹	• Fast-acting , small molecule E-selectin antagonist to eliminate vaso-occlusion
Early treatment in VOC is critical	 Potential self-administration of GMI-1687 after patient recognizes VOC episode 100% bioavailable in preclinical models following subcutaneous administration
Deliver full dose to stop VOC	 Optimize dose and regimen based on reductions in sE-selectin Agreed to as part of FDA Pre-IND Meeting

Phase 1a Study Completed

Potential Treatments in Oncology, Inflammation and Fibrosis

GALECTIN-3 INHIBITORS





The Promise of Targeting Galectins

Potential to modulate the immune and inflammatory response to cancer and fibrosis



Target Galectin-3 carbohydratebinding protein



Chemistry Rationally designed with proprietary platform



Differentiation Compounds have high binding affinity and specificity for Galectin-3



Central role in fibrosis and cancer

- Inflammation, aberrant cell activation/proliferation, fibrogenesis
- Blockade may prevent/reverse fibrosis following organ damage
- Antifibrotic/antitumor activity in various disease models



Orally Bioavailable



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Thank You

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