UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

FORM 8-K

CURRENT REPORT

Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): April 9, 2024

Genelux Corporation

(Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation) 001-41599 (Commission File Number) 77-0583529 (I.R.S. Employer Identification No.)

2625 Townsgate Road, Suite 230 Westlake Village, California (Address of principal executive offices) 91361

91361 (Zip Code)

Registrant's telephone number, including area code: (805) 267-9889

Not Applicable

(Former name or former address, if changed since last report.)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

□ Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)

□ Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)

D Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

□ Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered		
Common stock, par value \$0.001 per share	GNLX	The Nasdaq Stock Market LLC		

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§ 230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter).

Emerging growth company \boxtimes

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. \Box

Item 7.01 Regulation FD Disclosure.

On April 9, 2024, Genelux Corporation (the "Company") made available the corporate presentation attached hereto as Exhibit 99.1 (the "Corporate Presentation"). Information from the Corporate Presentation may also be used by the management of the Company in future meetings regarding the Company. For important information about forward-looking statements in the Corporate Presentation, see the slide titled "Forward-Looking Statements" in Exhibit 99.1 attached hereto.

The information contained or incorporated in this Item 7.01 of this Current Report on Form 8-K, including Exhibit 99.1, shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or otherwise subject to the liabilities of that section, nor shall it be deemed to be incorporated by reference into any filing under the Exchange Act or the Securities Act of 1933, as amended (the "Securities Act"), except as expressly set forth by specific reference in such filing to this Current Report on Form 8-K.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No.	Description
99.1	Corporate Presentation, dated April 9, 2024
104	Cover Page Interactive Data File (embedded within the Inline XBRL document).

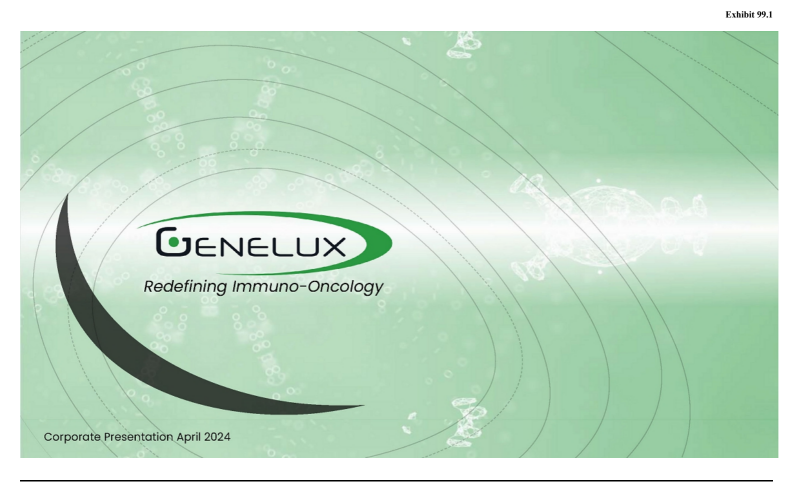
SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

Genelux Corporation

By: /s/ Thomas Zindrick, J.D. Thomas Zindrick, J.D. President and Chief Executive Officer

Date: April 9, 2024



Forward Looking Statements

This presentation contains forward-looking statements 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, which are subject to the "safe harbor" created by those sections, about Genelux Corporation ("Genelux," the "Company," "we," "us" or "our") that are based on the beliefs and assumptions of our management team, and on information currently available to such management team. These forward-looking statements include, but are not limited to, statements concerning: the expansion and advancement of our platform and pipeline and our approach and strategy related to the platform and pipeline; Olvi-Vec's potential utility and our plans and expectations for Olvi-Vec across various designs and indications; our expectations regarding the field of oncolytic viral immunotherapy; Olvi-Vec's potential to provide utility across multiple tumor types, and our expectations regarding our Phase 3 trial; the potential of our current and future pipeline to produce best-in-class drugs; our clinical trial strategy and design; our expectations regarding (i) the timing of our Phase 2 and Phase 3 clinical trials and (ii) our intellectual property rights under the Newsoara license agreement; our planned investments to meet worldwide clinical trial demand and facilitate our U.S. commercial launch; the commercial market opportunity for Olvi-Vec in the United States; our various commercial strategies for self-launching Olvi-Vec for ovarian cancer in the United States, including from Newsoara. These forward-looking statements are subject to numerous risks and uncertainties, many of which are beyond our control. All statements, other than statements of historical fact, contained in this presentation, including statements regarding future events, future financial performance, business strategy and plans, and objectives of our sfor future operations, are forward-looking statements.

Although we do not make forward-looking statements unless we believe we have a reasonable basis for doing so, we cannot guarantee their accuracy. These statements are only predictions and involve known and unknown risks, uncertainties and other factors, including the risks set forth under the heading "Risk Factors" in Annual Report on Form 10-K for the year ended December 31, 2023 and in our other filings with the SEC, which may cause our actual results, levels of activity, performance or achievements of and those of our industry to be materially different from any future results, levels of activity, performance or achievements so forward-looking statements. You should not place undue reliance on any forward-looking statement. Forward-looking statements are inherently subject to risks and uncertainties, some of which cannot be predicted or quantified. In some cases, you can identify forward- looking statements by terminology such as "anticipate," "believe," "contemplate," "continue," "could," "estimate," "expect," "intend," "may," "plan," "potential," "predict," "project," "should," "target," "will" or "would," or the negative of these terms or other comparable terminology, although not all forward-looking statements contain these identifying words. You should not put undue reliance on any forward-looking statements for ward. You should not put undue reliance on any forward-looking statements contain these identifying words. You should not put undue reliance on any forward-looking statements for words. You should not put undue reliance on any forward-looking statements contain these identifying words. You should not put undue reliance on any forward-looking statements should not be read as a guarantee of future performance or results, and will not necessarily be accurate indications of the times at, or by, which such performance or results will be achieved, if at all. Except as required by law, Genelux does not undertake any obligation to publicly update or revise any forward-looking statement, whether as a res

Trade names, trademarks and service marks of other companies appearing in this presentation are the property of their respective owners. Solely for convenience, the trademarks and tradenames referred to in this presentation appear without the $^{\circ}$ and $^{\text{\tiny M}}$ symbols, but those references are not intended to indicate, in any way, that we will not assert, to the fullest extent under applicable law, our rights, or the right of the applicable licensor to these trademarks and tradenames.

This presentation discusses a product candidate that is under clinical study and which has not yet been approved for marketing by the U.S. Food and Drug Administration. No representation is made as to the safety or effectiveness of this product candidate for the use for which it is being studied.

Highlights



The Most Advanced Non-local Delivery Oncolytic Immunotherapy

Olvi-Vec: 7 Completed Clinical Trials (>150 Patients)



Physician-preferred routes of delivery

- Regional and Systemic Administration to preferentially locate, colonize and destroy tumor cells, including metastatic disease
- In Ovarian Cancer trial, catheter placement is prior to chemotherapy, with removal 2 days after initial placement
- IV therapy currently being used in small cell lung cancer Phase 1b/2 trial





Antitumor Effect and Well Tolerated

- Strong data in Phase 1b/2 trial in platinumresistant/refractory ovarian cancer
- No Maximum Tolerated Dose (MTD) observed
- Potential utility in multiple cancers (demonstrated in 20 pre-clinical liquid & solid tumor models, e.g., ovarian, lung, breast, colon, kidney, prostate)



Ideal Backbone of Combination Therapy

- Turns tumors "hot" by localized inflammation and induction of the influx of tumor infiltrating lymphocytes (TILs)
- Positively modulates anti-tumor pathways in tumor microenvironment
- Potential to use with various modalities including in patients who failed platinum-based chemotherapy in multiple tumor types

Program Builds on Completed Trials to Exploit Competitive Advantages

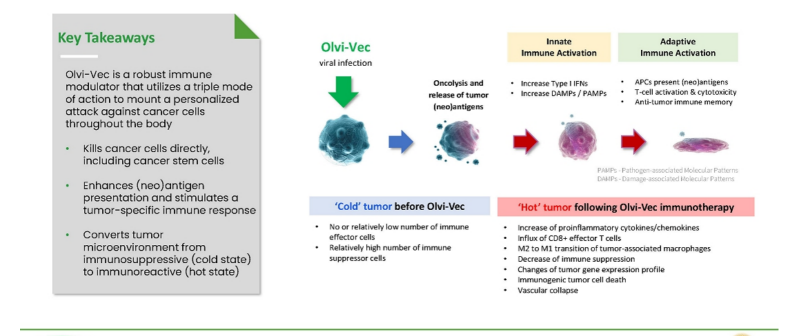
3 Upcoming Trial Readouts have Potential to Redefine:

Therapy (platinum resensitization in multiple indications)
 Modality (systemic administration of an oncolytic virus)

Olvi-Vec	Indication	Design	Preclinical	Phase 1	Phase 2	Phase 3	Anticipated Milestones	Collaborators
Regional Route	Ovarian Cancer (platinum-resistant/ refractory)	Olvi-Vec (i.pe) +Platinum-based regimen	Ph3 OnPrir	me/GOG-3076 Stu Received F	dy Actively Enrolli DA Fast Track Design		Topline results expected in 2H, 2025	Cooperative Group
	Non-Small Cell Lung Cancer (recurrent/platinum-ICI failure)	Olvi-Vec (IV) +Platinum/Checkpoint inhibitor-based regimen	Ph2 Regulat	tory Submission			Expected to initiate in 1H, 2024	
	Small Cell Lung Cancer (recurrent/platinum failure)	Olvi-Vec (IV) +Platinum-based regimen	Ph1b/2 Enro	lling			Expected interim readout in 2H, 2024	
Systemic Route	Ovarian Cancer (recurrent/platinum failure)	Olvi-Vec (IV) + Platinum-based regimen	Ph1b/2 Regulator Submission	V				NEWSGARA (Greater China)
Koule	Non-Small Cell Lung Cancer (recurrent/platinum-ICI failure)	Olvi-Vec (IV) +Platinum/Checkpoint inhibitor-based regimen	Planned					
	Pancreatic Cancer (recurrent)	Olvi-Vec (IV) +Adoptive Cell Therapy	Regulatory Submission					(Worldwide Rights Ex- Greater China)

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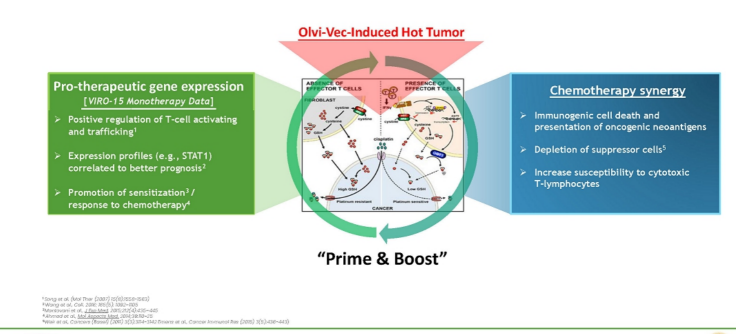
Selective Replication In Tumors Unleashes Immune System Against Cancer



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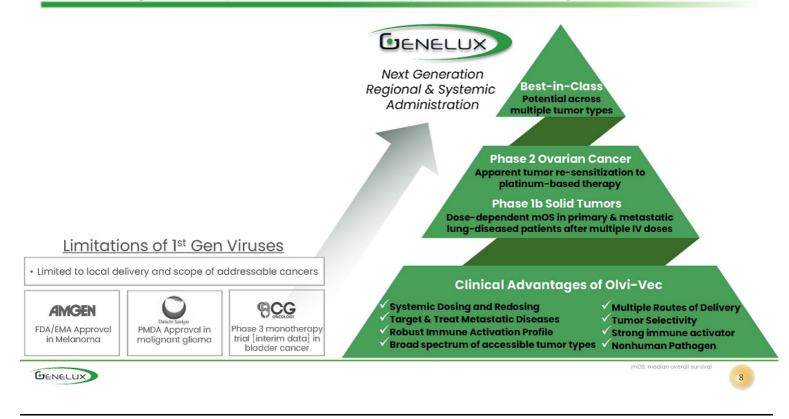
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Olvi-Vec-Primed Immunochemotherapy: Reversing Platinum Resistance





A Maturing Modality with Phase 3 Companies Validating OV Potential





Regional Administration Program

Ovarian Cancer

Ovarian Cancer Program: Regional (Intraperitoneal) Delivery

Key Takeaways

- Phase 1 tested condensed dosing schedule and demonstrated tolerability with evidence of antitumor activity
- Phase 2 demonstrated promising Objective Response Rate (ORR) and Progression Free Survival (PFS), and clinical reversal of platinum resistance and refractoriness
- Phase 3 registrational trial ongoing with topline results expected in 2H, 2025

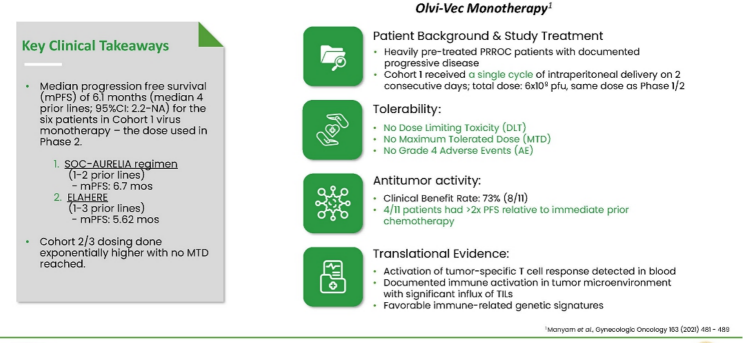
Completed and ongoing clinical trials in heavily pre-treated platinum resistant/refractory patients

Trial Sites Location / (#)	Clinical Stage	Design	Patients	Randomization	Status
US / (1)	Phase 1	Monotherapy (Dose Escalation)	11	Single Arm	Completed ¹
US / (2)	Phase 2	Combination (platinum-based regimen)	27	Single Arm	Completed ²
US / (~30)	Phase 3	Combination (platinum-based regimen)	186	2:1	Enrolling ³

¹ Manyam et al., <u>Synecol Oncol</u>. 2021;163(3);481–489. ² Holloway et al., <u>JAMA Oncol</u>. 2023 Jul 1;9(7):903–908. ³ Holloway et al., <u>Int J Gynecol Cancer</u>. 2023 Sep 4;33(9):1458–1463.



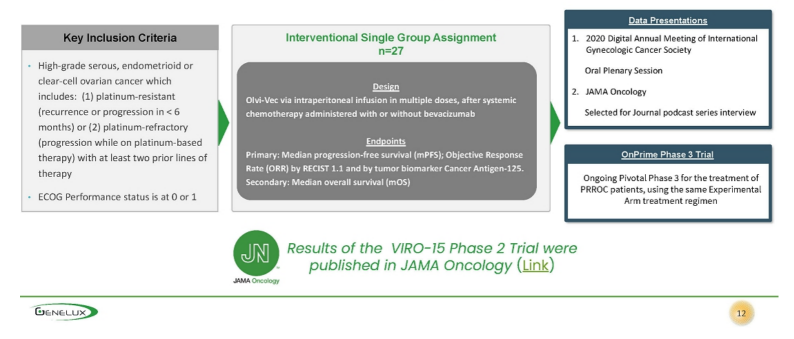
Phase lb: Anti-tumor Activity as Monotherapy Leading into Combination





Completed Phase 2 Tested Olvi-Vec-primed Immunochemotherapy

Heavily Pretreated Patients with Platinum-Resistant or Platinum-Refractory Ovarian Cancer



Key Clinical Takeaways

Promising ORR and PFS, and clinical reversal of platinum resistance and refractoriness among patients with PRROC

- All patients had documented ٠ progressive disease at enrollment
- The mPFS of the patients' . immediately preceding line of therapy was ~4.5 months
- Based on historical data, the ٠ mPFS would be expected to decrease in the subsequent line of therapy

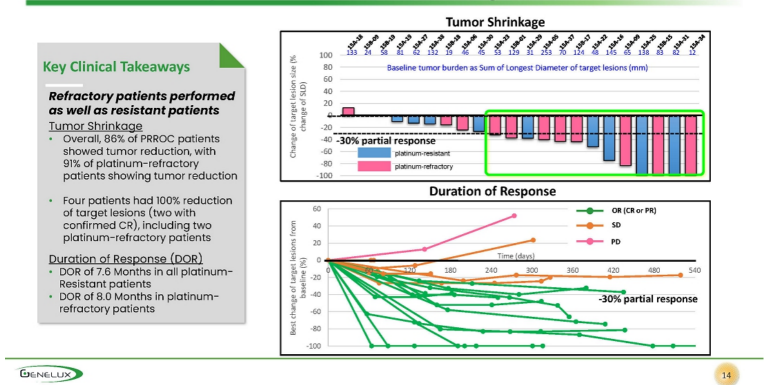
Overall Response Rate (ORR) & Progression-Free Survival (PFS)*

	ORR by RECIST1.1 [™]	Duration of Response	ORR by CA-125	Median PFS	Median Overall Survival (OS)
All patients (n= 27) (95% Cl)	54% (13⁰/24⁰) (33 - 74)	7.6 mos (3.7 - 9.6)	85% (22/26 ^{∞∞}) (65 - 96)	11.0 mos (6.7 - 13.0)	15.7 mos (12.3 - 23.8)
Platinum-resistant (n=14) (95% Cl)	55% (6/11) (26 - 84)	7.6 mos (3.7 - NA)	85% (11/13) (55 - 98)	10.0 mos (6.4 - NA)	18.5 mos (11.3 – 23.8)
Platinum- refractory (n=13) (95% CI)	54% (7/13) (27 - 81)	8.0 mos (3.7 - NA)	85 % (11/13) (55 - 98)	11.4 mos (4.3 -13.2)	14.7 mos (10.8 – 33.6)

*Baseline for ORR & PFS evaluation is the timepoint immediately prior to starting post-Olvi-Vec carboplatin doublet +/-bevacizumab to allow direct comparison to historical data or patients' own previous line of chemotherapy **Eligible for evaluation: with at least 1 measurable target lesion at baseline; including 2 patients without post-chemo scan after virotherapy, and therefore are assigned to the 'inevaluable for response' category per RECIST1.1 *Including 3 unconfirmed; 2 in resistant and 1 in refractory groups *Three of 27 patients were not evaluable as defined by RECIST 1.1 criteria due to no measurable disease. However, these 3 patients were evaluable by the Gynecological Cancer InterGroup (GCIG) CA-125 criteria, showing 2 partial responses and 1 complete response as best response.



Demonstrated Deep and Durable Tumor Shrinkage

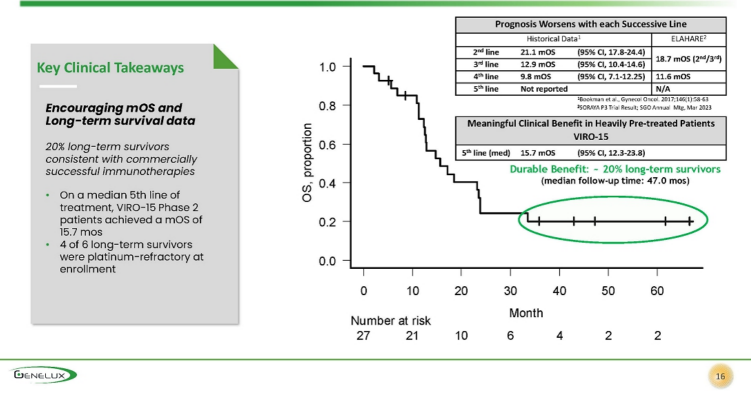


Olvi-Vec-Primed Immunochemotherapy Overcomes "Refractoriness"

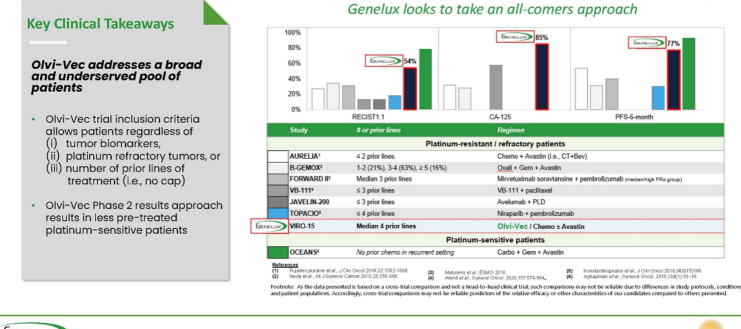
Exemplary platinum-refractory patients, after platinum re-challenge, achieved PFS exceeding any prior lines



Durable Survival Benefit



Seeking to Reset Life Clock of Heavily Pre-treated Patients



While clinical remissions are obtainable, a majority of patients will relapse.



Phase 3 Pivotal Trial Design Founded on Phase 2 Trial Design & Results

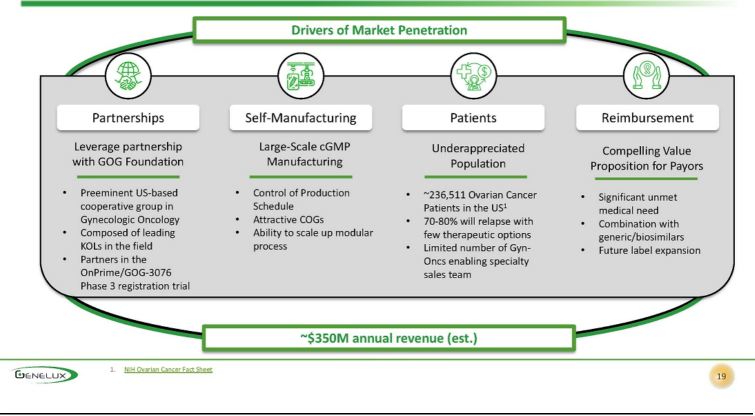
Trial design intends to replicate previous data showing anti-tumor activity of Olvi-Vec and reversal of platinum resistance.



- ¹ International Journal of Gynecological Cancer, Holloway RW, et al. 2023;33:1458–1463.
 ² Protocol amended to make platinum optional in the Active Comparator Arm with Intent to Implement upon receipt of IRB approvals.
 ³ Journal of Investigative Medicine High Impact Case Reports, Volume 6: 1–3, 2018 DOI: 10.1177/2324709618760080 Journals.sagepub.com/home/hic



Self Launch Olvi-Vec for Ovarian Cancer in the US





Systemic Administration Programs

Lung Cancers

Key Takeaways

- Funding commitment by Newsoara of the US-based Genelux Phase 2 trial in NSCLC
- Genelux has worldwide commercial rights (ex-Greater China) to all clinical data generated in China¹
- All patients in these trials will be treated systemically and have previously failed platinum-based therapy
 - Expected Milestones
- Ph2 NSCLC: Initiate 1H, 2024
- Phib SCLC: Interim readout 2H, 2024

¹Newsoara has development and commercialization rights in Greater China



Ongoing and Planned Clinical Trials

Sponsor	Trial Sites	Indication	Clinical Stage	Patients (est.)	Randomization	Status
G	US	Recurrent/platinum- ICI failure NSCLC	Phase 2	~142	1:1	Regulatory Submission
		Recurrent/platinum failure SCLC	Phase 1b/2	~110	Single Arm	Enrolling
NEWSGARA	China	Recurrent/platinum failure OC	Phase 1b/2	~150	2:1	Regulatory Submission
		Recurrent/platinum- ICI failure NSCLC	Phase 1b/2	~150	2:1	Planned

Systemic Administration Demonstrated Dose-dependent OS Benefit

Key Clinical Takeaways

Demonstrated feasibility and clinical benefit of multiple IV cycles

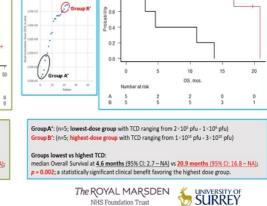
- Median 5 prior lines of therapy
- Regimen: various dosing levels and schedules (typically over 4-6 months)
- Well tolerated: no-MTD reached with one DLT
- Duration of Treatment (DoT): Higher cumulative-dose patients assigned to cohorts with DoT shorter than (condensed schedule) or equal to the DoT of patients assigned to lower cumulative-dose cohorts
- <u>Clinical Benefit</u>: statistically significant virus dose-dependent OS benefit in primary and metastatic lung diseases

1.0 Group B 0.8 20.6 6 6 0.4 0.2 0.0 50 10 40 20 30 OS mor AB 11 0 0 3 0

Dose Escalation Phase 1b Monotherapy Study in Solid Tumors Progressed from Last Prior Therapy

Group A : (n=11; lower-dose group with TCD ranging from 2×10^{9} pfu - 2×10^{9} pfu) Group B : (n=11; higher-dose group with TCD ranging from 3×10^{9} pfu - 3×10^{10} pfu)

Groups lower vs higher TCD: median Overall Survival at 4.6 months (95% CI: 1.3 - 11.0) vs 16.8 months (95% CI: 5.9 - NA); $\rho = 0.026$; a statistically significant clinical benefit favoring the higher dose group.



NHS Foundation Trust

ICR The Institute of Cancer Research

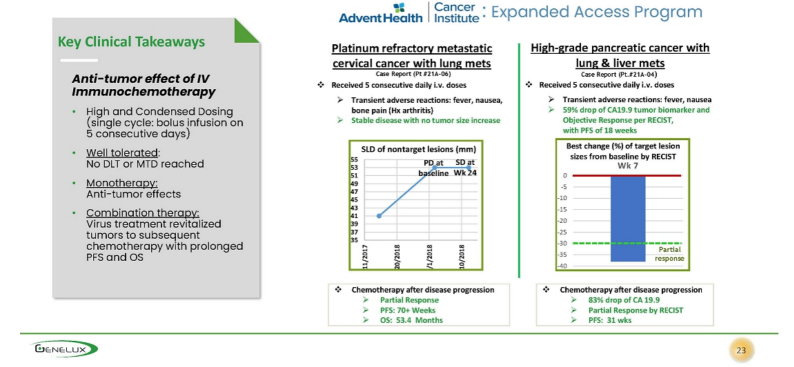
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0.8

Group A'



Systemic Administration + Chemo Generated Encouraging Data



Phase 2 Trial in Recurrent Non-small Cell Lung Cancer

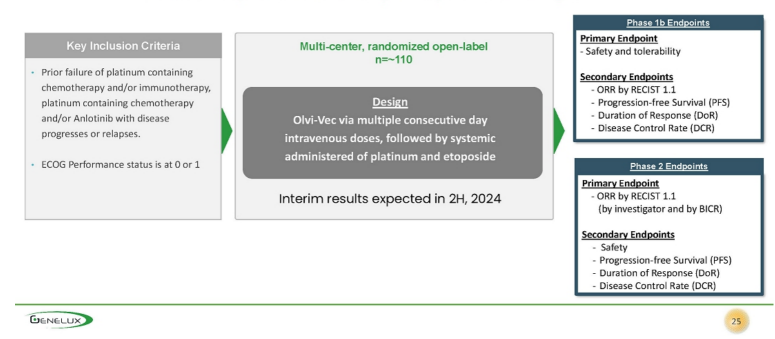
Patients with Non-small Cell Lung Cancer after First Progression while on Front-Line Immune Checkpoint Inhibitor-based Maintenance





Phase 1b/2 Trial in Recurrent Small Cell Lung Cancer

Heavily Pretreated Patients with Platinum-Relapse or Platinum-Refractory Small Cell Lung Cancer



Validating Industry Collaboration with Newsoara BioPharma Co., Ltd





V2ACT Therapeutics LLC: Joint Venture between GNLX and TVAX BioMedical



Key Trial Takeaways

V2ACT Immunotherapy, combines an oncolytic immunotherapy and adoptive cell therapy

- Induces an acute inflammatory response in the tumor and converts tumor microenvironment from immunosuppressive to immunostimulatory;
- Anticipated to enhance effect of neoantigen specific effector T cells

V2ACT Therapeutics is a joint venture between Genelux Corporation and TVAX Biomedical, Inc. established to develop and test V2ACT.

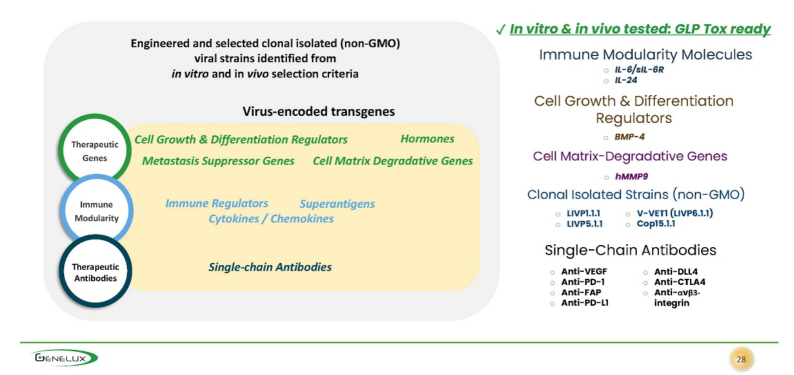
Vaccination increases the numbers of neoantigen-specific T cells in the body and Olvi-Vec kills cancer cells and potentiates T cells by increasing cancer tissue receptivity to adoptively transferred neoantigen-specific effector T cells.

Technology	TVI Adoptive Cell Therapy	Olvi-Vec Oncolytic Immunotherapy
Patients Dosed	~ 130	~ 150
Regulatory	Fast Track Designation / FDA Grant - glioblastoma	Phase 3 enrolling - ovarian

Novel IO modality: United States Patent No. 11,633,442, issued in April 2023



Choice Platform Library: 500+ Vectors with 110+ Transgenes



Intellectual Property: Market Exclusivity & Freedom to Operate



Patent Portfolio: 33 issued patents & 7 pending; Olvi-Vec covered by Composition of Matter (2031*) and Manufacturing (2038)



Olvi-Vec: Worldwide operating freedom; No third-party royalties due



Long Duration of Regulatory / Marketing Exclusivity



*Reflects Patent Term Extension





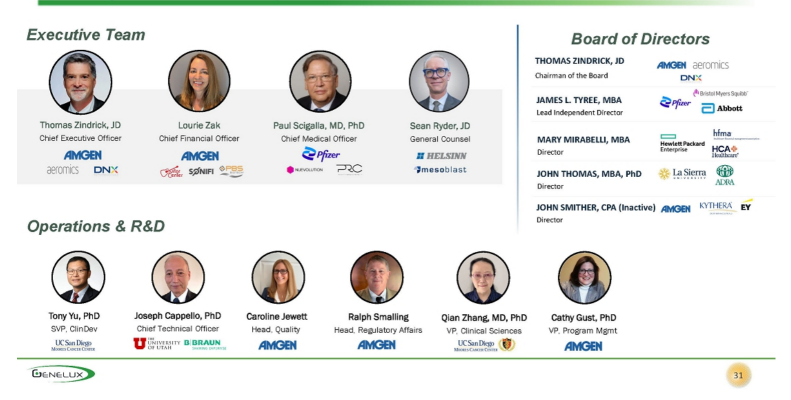
Integrated R&D and Manufacturing Capabilities For Phase 3 And Launch





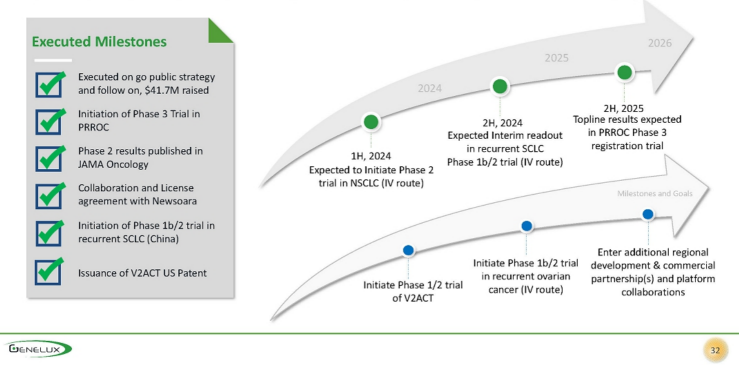


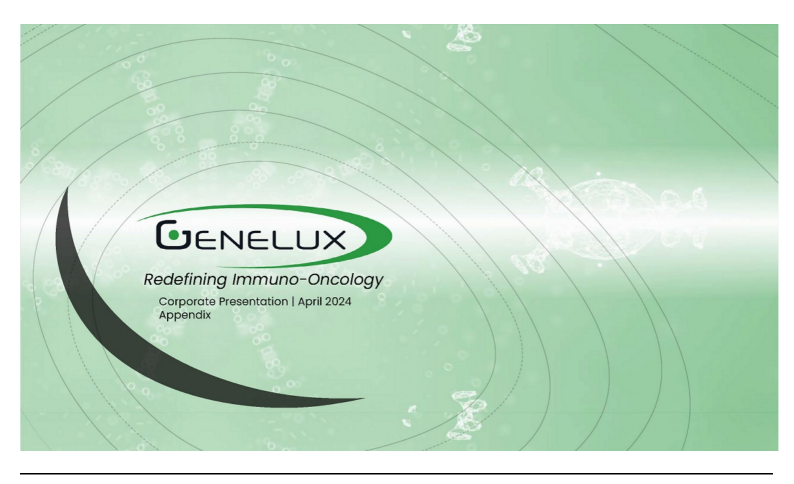
Accomplished Leadership Team



Genelux Has Executed on Multiple Milestones and is Positioned for the Future

Expected Operating Runway into 2Q 2025 - Regular Cadence of Important Program Milestones starting 2H, 2024



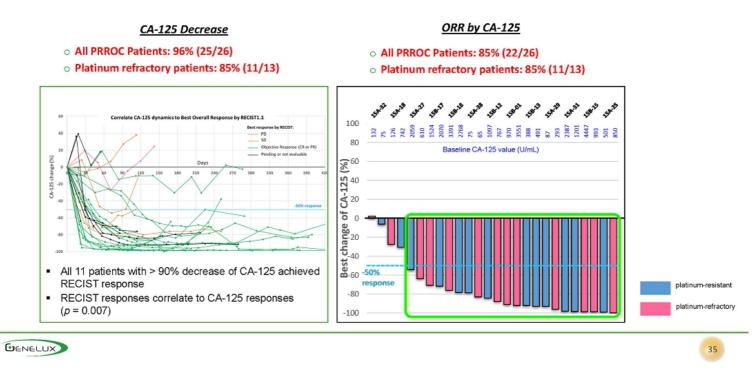


Accomplished Clinical Advisory Board

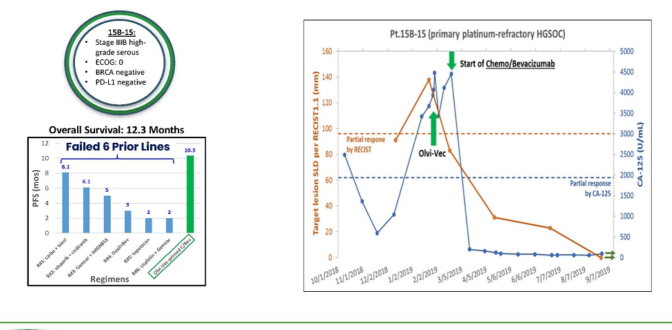
Medical Director, Gynecologic Cncology, AdventHealth Cancer Institute	Robert Holloway, MD CHAIRMAN	Dr. Holloway is the principal investigator for VIRO-15 and has served on several committees of the Society of Gynecologic Oncology (SGO), including its Board of Directors.
Chief Medical Officer, Vanium Croup	Robert Coleman, MD Member	Dr. Coleman currently serves on the Board of Directors of Gynecologic Oncology Group and is co-Director of GOG-Partners. In addition, he is immediate Past-President of the International Gynecologic Cancer Society.
Co-Director, Gynecologic Oncology, Hoog Memorici Hospital Presbyterian	Albert A. Mendivil, MD Member	Dr. Mendivil, site principal investigator for VIRO-15, serves as Co-Director of Gyn- Onc and Complex Pelvic Surgery, Hoag Hospital. He has been the principal investigator or site sub-investigator on 20+ clinical trials.
Deputy Director of the University of Cincinnati Cancer Institute	Thomas J. Herzog, MD Chief Executive Officer	Dr. Herzog is President of the GOG Foundation. He has served on the leadership board or council of SGO, the Foundation for Women's Cancer, and ACOG.
Professor and Division Director, Chio State University Comprehensive Cancer Center	David M. O'Malley, MD Chief Medical Officer	Dr. O'Malley is the clinical trial advisor/lead for ovarian cancer within GOG Partners, a committee member for the NCI Gynecologic Cancer Steering Committee's Ovarian Task Force and the NRG Oncology.
Forsythe & Bear,	Alan Forsythe, PhD Chief Financial Officer	Dr. Forsythe has had a distinguished career in pharmaceutical drug development. As Vice President of Corporate Biomedical Information at Amgen, Alan led the Biostatistics, Epidemiology and HOER depts.
GENELUX		34

Olvi-Vec-primed Immunochemotherapy Anti-tumor Activity: CA-125 Biomarker

Rapid, Common and Durable Responses



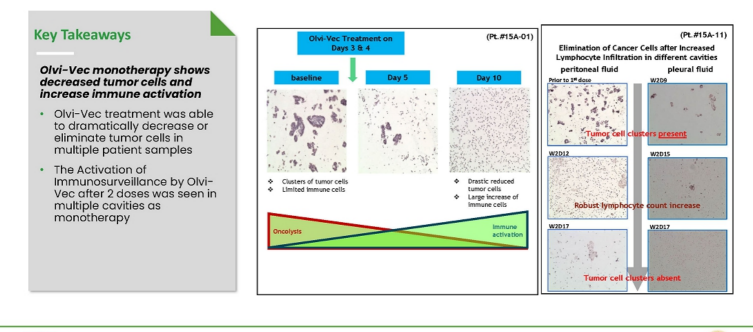
Exemplary platinum-refractory patients, after platinum re-challenge, achieved PFS exceeding any prior lines



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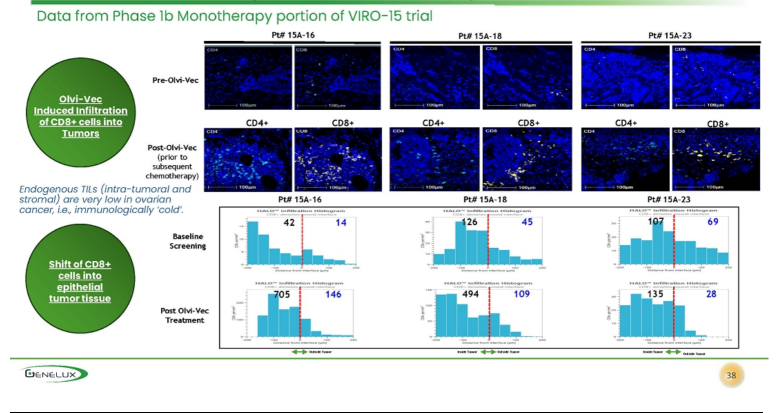
Olvi-Vec Monotherapy Demonstrates Oncolysis and Immune Activation

Data from Phase 1b Monotherapy portion of VIRO-15 trial



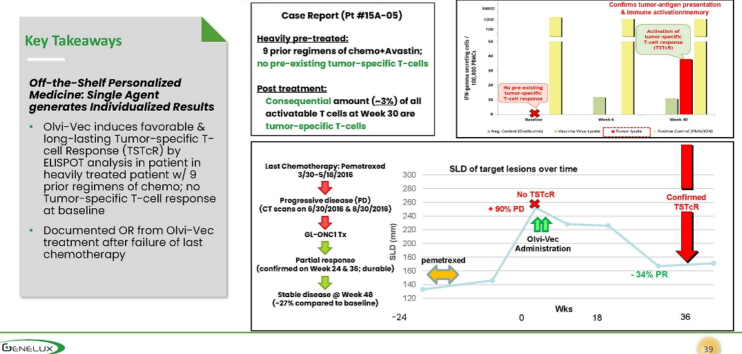
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CD8+ T-cells: Infiltrating Lymphocytes are Prognostic for Response/Survival



Long-lasting, Tumor-specific T cell Response Corresponds to Tumor Reduction

Data from Phase 1b Monotherapy portion of VIRO-15 trial



Olvi-Vec: Ideal Backbone for Combination Therapy

