

### Oncocyte Announces Successful GraftAssure™ Beta Launch and Q2 2024 Results

Interest in kidney transplant assay represents approximately 25% of US transplant volume Company lays out land and expand strategy in transplant

IRVINE, Calif., August 8, 2024 (GLOBE NEWSWIRE) -- Oncocyte Corp. (Nasdaq: OCX), a diagnostics technology company, today published its second quarter 2024 results and the following letter to shareholders.

#### Dear shareholders,

We are thrilled to report that we have successfully launched GraftAssure<sup>TM</sup>, which is our research-use-only assay that can detect early evidence of graft organ damage in a patient's blood.

On July 11, 2024, our GraftAssure assay was run at a major metropolitan transplant center and research university in the northeast United States, representing the first time our test had been run outside of an Oncocyte lab. Days later, a lab at a leading transplant center in Southeast Asia began running the test.

Though we believe this milestone is just the beginning, it's an important proof point in an 18-month journey. To recap, in January 2023, we decided to double down on commercializing our transplant test IP, primarily by making a kitted test that relies upon a biomarker, donor-derived cell-free DNA (dd-cfDNA). Our first prototypes of GraftAssure came off the manufacturing line in December. Four months later, in April 2024, we welcomed Bio-Rad Laboratories (NYSE:BIO) as an investor and strategic partner, supporting the GraftAssure global launch.

And in June 2024, we began shipping to our launch customers. We shipped to two continents in our first week – representing major transplant centers and research universities in the U.S. and Southeast Asia. As of today, we have signed up several centers in the U.S, Europe and Southeast Asia.

We are also pleased with the momentum we are seeing coming out of the American Transplant Congress in Philadelphia in June. That conference was especially meaningful for Oncocyte this year because it coincided with the New England Journal of Medicine's <u>publication</u> of favorable data regarding our centralized lab product VitaGraft Kidney<sup>1</sup>. That clinical product uses the same background IP as our GraftAssure kitted test.

We are pleased to report that our US sales funnel of confirmed interest in our transplant assay is made up of transplant centers that represent approximately 25% of US transplant volumes<sup>2</sup>. We also are seeing significant interest in Germany, as well as in other European countries, and in Asia and the Middle East. The level of interest and enthusiasm in the market is exceeding our expectations.

As a reminder, under our partnership, Bio-Rad and Oncocyte are co-marketing GraftAssure inside the US and Germany, with Oncocyte acting as commercial lead. Bio-Rad is acting as commercial lead outside the US and

<sup>&</sup>lt;sup>1</sup> Oncocyte's Drs. Ekke Schuetz and Julia Beck, inventors of the technology, are among the authors of the study. VitaGraft Kidney was used to monitor graft injury in a phase 2 double-blind, placebo-controlled study (NCT05021484) of the investigational drug felzartamab, a fully human CD38 monoclonal antibody, for antibody-mediated rejection (AMR), a leading cause of kidney allograft failure.

<sup>&</sup>lt;sup>2</sup> The centers in our U.S. funnel who have confirmed interest in GraftAssure represent more than 11,000 transplants per year, which is about 24% of 46,000 annual transplants conducted, according to the United Network for Organ Sharing. (Source: https://unos.org/about/fast-facts/#:~:text=ln%202023%3A,continuing%20annual%20record%2Dsetting%20trend)

Germany, and correspondingly has been granted exclusive distribution and commercial rights in those countries.

Beyond growing our sales pipeline, momentum also continues through our industry relationships and partnerships. For example, in the second quarter, we signed an agreement with a European biotechnology company to be the provider of dd-cfDNA testing for a Phase II clinical trial for a therapeutic in antibody mediated rejection.

Our partnerships with the pharmaceutical industry are strategically important. First, the partnerships highlight and validate the robustness of our technology and our partnership approach to the market. These trials also advance and expand the body of science using our diagnostic technology to develop the potential for new clinical applications for our diagnostic tools.

Ultimately, the expansion of clinical utility increases the opportunity for future long-term recurring revenue. When we participate in a clinical trial, we perform the services at our clinical (CLIA) laboratory, receiving a nominal amount of revenue to cover the cost of performing the test. While these clinical trials are limited in short-term revenue generation, they lay the foundation for future revenue expansion and growth.

The dd-cfDNA test that we run at our CLIA laboratory on behalf of customers is VitaGraft Kidney. As a reminder, VitaGraft received Medicare reimbursement in 2023. We are increasing the throughput for VitaGraft in our CLIA lab by establishing higher automation with simple scaling options. This increased capacity supports ongoing clinical trials using our tests and the data generation required by our IVD program, which we discuss further below.

#### **Transplant: Land & Expand Strategy**

We believe that we are at an inflection point in commercializing our molecular diagnostic tests that can detect early evidence of graft organ damage in the blood. Bringing research centers up online with our GraftAssure assay is a key part of our land-and-expand strategy to drive commercial adoption of our tests and capture market share in an estimated \$1 billion total addressable market.

Simply, our strategy is to *land* major transplant centers and research universities with our research-use-only product. Doing so establishes our technology and increases its potential utility by enabling researchers to explore potential applications of dd-cfDNA. Then, once we have achieved FDA (Food and Drug Administration) clearance for our test kits to be used to make clinical decisions – that is, approved as an in-vitro diagnostic (IVD) – we believe that these institutions will begin using our tests to manage their patient populations, while continuing to use our GraftAssure test kit to perform research. Importantly, our GraftAssure research product may *not* be used to support clinical treatment decisions.

We believe that we can execute upon this strategy with a limited sales force deployed in key territories due to the favorable customer concentration of our industry. For example, fewer than 100 academic and research centers in the US account for about 80% of transplant volumes. And rest-of-world markets are similarly concentrated at high-end academic institutions. Bio-Rad's global infrastructure puts those centers well within reach, allowing for high-touch sales and service at the local level. Partnering into this commercial channel allows Oncocyte to keep our infrastructure costs low while we focus on scaling in manufacturing and developing new use cases. As a result, we believe we have the potential to be more profitable than many larger, better-known diagnostic companies.

We are seeing early indications of remarkable product-market fit and we are pleased with the market response to our assay. We believe that GraftAssure's appeal is due to workflows that are easy and user-friendly<sup>3</sup> and that market enthusiasm will translate to adoption of the IVD product once it is available. We discuss the IVD clearance process in more detail below.

Our initial beta customers sought to deploy our RUO assay as soon as we could ship it to them – which we find compelling in light of the fact that bringing our RUO assay up and running is a major undertaking for a transplant center's research lab, requiring significant investment of time and resources. It requires performing instrument validation, running an analytical validation study, and training laboratory employees.

To support the lab's validation process, we ship non-revenue test kits and send a team to support the lab in getting the test up and running, including troubleshooting any initial issues. The first lab to be up and running was able to complete the process in about two weeks, with some initial support from the Oncocyte team until they could run with the assay.

Our teams ensure the pre-existing equipment is calibrated to support the test and that the customer laboratory teams are trained to run it effectively. Once we've confirmed that the instrument and assay are performing as intended, the labs begin running their first research samples and preparing to launch new studies. Most of the customers in our funnel come to us with studies in mind that they've wanted to do for a long time and are excited that they'll get a chance to move forward on their own.

Notably, some of these centers are eager to support the development of our IVD product. Whether collecting samples or producing data, the clinical researchers are excited to work with us on this process. Our openly collaborative approach is winning hearts and minds across the industry. We take our commitments and responsibility to the research community very seriously – they use our tests to advance the body of science in transplant medicine, which may save many lives over time.

We aim to have more than 20 transplant centers running GraftAssure tests through the end of 2025, including 15 in the US. We estimate that each center represents an eventual annual high-margin revenue stream of several hundred thousand dollars to \$2 million, depending on the size of the center.

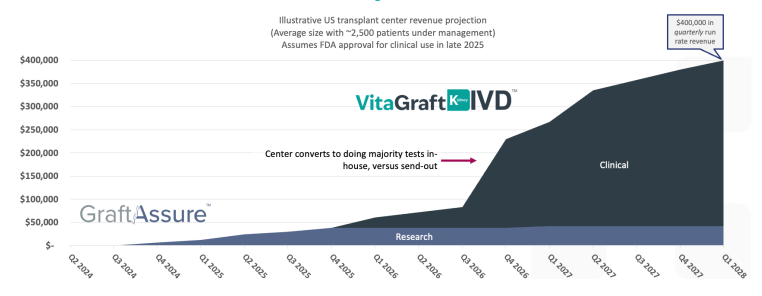
The following illustration shows a projected quarterly revenue path for one typical beta customer. The illustration shows the customer entering 2028 contributing \$400,000 in quarterly revenue, or \$1.6 million in annual run rate revenue. Importantly, this revenue path assumes our best-case timeline<sup>4</sup> for FDA clearance of the IVD product for clinical use in late 2025 and a MoIDX<sup>5</sup> reimbursement decision attached to the IVD product in early 2026.

<sup>&</sup>lt;sup>3</sup> Bio-Rad's QX600 Droplet Digital PCR (ddPCR) System – which is its new six color instrument – allows us to create a simple workflow for the lab technician, delivering a result in four to eight hours, compared with ≥30 hours for competitors that are using NGS technology. Further, testing a single sample is an affordable option, given that the batch size – in contrast to NGS – does not alter the price per result.

<sup>&</sup>lt;sup>4</sup> The revenue build is purposely limited in scope to the current clinical utility established for VitaGraft Kidney. Incremental revenue opportunities could come from additional claims, such as additional tissue types like heart and lung, anti-CD38 drug monitoring, and high-risk patient monitoring. Notably, our best-case FDA clearance timeline should not be confused with our best-case revenue opportunity.

<sup>&</sup>lt;sup>5</sup> MolDX is a program that identifies and establishes coverage and U.S. government reimbursement for molecular diagnostic tests.

### Beta customer revenue path



#### What's next:

- **Continued roll-out of research product:** We will continue to bring transplant center customers up online running our GraftAssure assay throughout 2024 and 2025.
- Impact of legislation: We believe our timing is serendipitous as the Increasing Organ Transplant Access Model, by the Centers for Medicare and Medicaid Services, rolls out on January 1, 2025. Transplant surgeons have told us that they expect that this legislation will lead to more marginal organs being transplanted into patients and potentially higher rates of adverse events, which could increase demand for dd-cfDNA testing. As a reminder, our IVD strategy aims to empower transplant centers to perform dd-cfDNA tests in-house and submit for reimbursement for their own labs.
- Q-Sub submission: The Q-Sub process is a formal procedure in which a company submits a validation
  plan to the FDA for review and receives their feedback. We hope to receive feedback from the FDA by
  the end of 2024. We need this feedback to move forward with confidence into our clinical validation
  processes prior to submitting for FDA IVD clearance next year.
- **FDA IVD-clearance process:** Medical devices are classified into three categories based on their risk our tests are considered Class II devices, meaning that they are considered to have a moderate level of risk<sup>6</sup> to the patient. We will submit under the 510(k) De Novo Classification process, which is premarket submission made to the FDA for a new "device," or in our case, test. We are aiming to achieve FDA clearance in late 2025, which is our best-case timeline. Below are some of the data points that give us confidence going into this process.
  - The successful launch of GraftAssure, and the resulting engagement that we are having with the community, supports our IVD product development efforts. Many of the sites that are in our

<sup>&</sup>lt;sup>6</sup> An example of a Class I, or low risk, device, would be a bandage and an example of a Class III, or high risk, device would be an implantable device like a cardiac pacemaker.

- funnel have expressed interest in collecting samples and potentially being a reproducibility site to support our ultimate submission to the FDA.
- We have found recent dialogue with the FDA to be constructive. We look forward to more interactions throughout the year in the formal Q-Sub process described above.
- Key members of our team have deep FDA regulated product experience.<sup>7</sup>
- We have deep technical and clinical expertise in transplant, with a greater-than-ten-year track record of developing innovative technology and publishing novel data. It's a history that compares favorably to any other player in the transplant rejection diagnostics space.
  - Examples include:
    - Patents granted in both the US and EU support the technical innovation core to the IVD development process, protecting our market opportunity.
    - MolDX<sup>8</sup> cited our publications twice when it established the LCD (Local Coverage Determination) for Medicare and Medicaid reimbursement coverage for cell free DNA testing.
    - In the New England Journal of Medicine publication referenced above, we are
      the first company to have published randomized interventional data showing the
      efficacy of a transplant rejection diagnostic test along with antibody mediated
      rejection (AMR) therapy.
    - We are also the only company to have randomized interventional clinical trial data using dd-cfDNA to detect AMR in *de novo* DSA+ (donor-specific antibody positive) kidney patients, finding AMR 10 months sooner<sup>9</sup> than other monitoring protocols.
- Our manufacturing partner has significant experience in the production of regulated products for both US and global distribution, and the ability to scale to meet global demand.

#### **New CFO Appointed**

In preparation for the next several years of sustained, rapid growth, we welcomed Andrea James as Chief Financial Officer in June. She has a proven track record of guiding financial strategy through multiple phases of growth, raising and stewarding capital, and building relationships with high quality institutional investors.

Prior to joining Oncocyte, Andrea served as chief communications officer and head of investor relations at Axon Enterprise, Inc. (Nasdaq: AXON). She joined Axon's finance team in 2017 to build the company's investor relations function and played an integral role through the company's growth from \$1 billion in market capitalization to a company valued at more than \$20 billion. She led Axon's repositioning as a top-tier technology company with the investment community and established Axon's corporate strategy function, overseeing capital stewardship, mergers and acquisitions, strategic partnerships and investments. Her work was instrumental in supporting more than \$550 million in equity capital offerings.

Prior to Axon, Andrea worked in a strategic investor relations role for Tesla, Inc., in 2016 and 2017. From 2009 to 2016 she worked as a sell-side analyst for Dougherty & Company (now Colliers Securities), becoming a vice president and senior research analyst. At Dougherty, she researched emerging technologies on behalf of institutional investors. Previously, she was a reporter at publications including Bloomberg News and the

<sup>&</sup>lt;sup>7</sup> As an example, our Chief Technology Officer, Johnson Chiang, has successfully led or participated in numerous IVD submissions to the US FDA and the EU under IVDD

<sup>8</sup> Source: https://www.cms.gov/medicare-coverage-database/view/lcd.aspx?lcdId=38671&ver=4

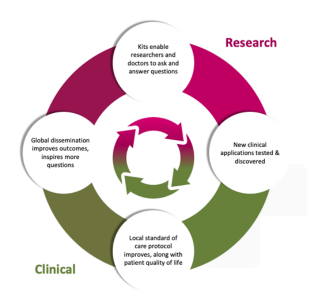
<sup>&</sup>lt;sup>9</sup> For more information: https://investors.oncocyte.com/news-releases/2023/09-18-2023

Seattle Post-Intelligencer covering a range of business and financial beats. She holds a Bachelor of Science, summa cum laude, in Computer Information Systems from American University. She also holds a Master of Science in Journalism from Northwestern University, graduating with Kappa Tau Alpha honors. While at Dougherty, her Financial Industry Regulatory Authority (FINRA) licenses included the Series 7, 86, 87 and 63.

#### **Q2 2024 Financial Overview**

- Revenue of \$104,000 in the second quarter was derived from pharma services performed in Nashville.
- A gross profit of \$50,000 reflected the relatively fixed costs of operating our Nashville CLIA laboratory. These costs include labor as well as infrastructure expenses such as the depreciation of laboratory equipment, allocated rent costs, leasehold improvements, and allocated information technology costs.
- Operating expenses of \$4.7 million included \$386,000 in non-cash stock-based compensation expenses, \$326,000 in non-cash depreciation and amortization expenses and a \$1.0 million non-cash benefit from the change in fair value of contingent consideration. Excluding these non-cash items in the current and prior periods, our Q2 operating expenses decreased approximately 5% sequentially and decreased 9% year over year.
- Research and development expense of \$2.5 million increased approximately 6% sequentially.
- Sales and marketing expense of \$853,000 was flat sequentially, reflecting cost discipline even as we began to ramp the commercialization of our transplant tests.
- General and administrative expense of \$2.4 million fell 10% sequentially, reflecting cost discipline and reductions in facilities and personnel related costs.
- Loss from continuing operations was \$4.5 million, or \$0.36 per share.
- Non-GAAP loss from operations was \$5.0 million and excludes certain non-cash items. Please refer to the table below, "Reconciliation of Non-GAAP Financial Measure," for additional information.
- Oncocyte's cash, cash equivalents, and restricted cash balance at the end of the second quarter was approximately \$11.0 million, up \$3.7 million sequentially. This balance includes proceeds from our capital raised in April as well as \$1.7 million in restricted cash.
  - On April 11, 2024, we entered into a private placement securities purchase agreement with certain accredited investors. The resulting net proceeds were approximately \$9.9 million, after deducting offering expenses of \$539,000 and deducting \$5.4 million for the redemption of all remaining shares of our Series A Redeemable Convertible Preferred Stock. These proceeds include the investment from Bio-Rad. We are pleased to be entering a growth inflection point with a streamlined capital structure.
  - Cash flows from operations reflected \$1.4 million in bonus and retention payments made to employees in the second quarter, mostly as part of our annual compensation cycle.

#### **New Investor Deck Published**



To view our quarterly update in the context of our larger strategy, you may access a refreshed investor deck at investors.oncocyte.com or access it at:

https://investors.oncocyte.com/~/media/Files/O/Oncocyte-IR/events-and-presentations/ocx-investor-deck-q2-2024.pdf.

In the deck, we outline our innovation flywheel and provide more details on our land and expand strategy, with key upcoming milestones and catalysts.

#### Conclusion

We are pleased to be delivering upon our commitments and are thrilled with our market momentum. After

months of effort, we are seeding the market with our RUO product – which is the first step in our land and expand strategy. And we believe that we already have clear signs of remarkable product-market fit.

Our mission is to democratize access to novel molecular diagnostic testing to improve patient outcomes. We believe that combining innovative science with a simple and disruptive business model can create enormous societal and shareholder value.

We look forward to welcoming you on our growth journey and updating you on our continued progress throughout the year.

The Oncocyte Management Team

#### **Webcast and Conference Call Information**

Conference Call and Webcast on Thursday, August 8, 2024 at 2:00 p.m. PT / 5:00 p.m. ET

You may access the live call via telephone by dialing toll free 800-715-9871 for domestic callers. Once dialed in, ask to be joined to the Oncocyte Corporation call.

The live webcast of the call may be accessed by visiting the "Events & Presentations" section of the Company's website at https://investors.oncocyte.com. A replay of the webcast will be available on the Company's website.

#### CONFERENCE CALL DETAILS SEP

Participant Toll-Free Dial-In Number: (800) 715-9871 Participant Toll Dial-In Number: +1 (646) 307-1963

Conference ID: 4153469

#### Webcast SEP

https://events.q4inc.com/attendee/478523544

#### **About Oncocyte**

Oncocyte is a molecular diagnostics technology company. The Company's tests are designed to help provide clarity and confidence to physicians and their patients. VitaGraft™ is a clinical blood-based solid organ transplantation monitoring test. GraftAssure™ is a research use only (RUO) blood-based solid organ transplantation monitoring test. DetermalO™ is a gene expression test that assesses the tumor microenvironment to predict response to immunotherapies.

DetermaCNI™ is a blood-based monitoring tool for monitoring therapeutic efficacy in cancer patients. For more information about Oncocyte, please visit https://oncocyte.com/. For more information about our products, please visit the following web pages:

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VitaGraft Kidney™ - https://oncocyte.com/vitagraft-kidney/⑤  
VitaGraft Liver™ - https://oncocyte.com/vitagraft-liver/⑥  
GraftAssure™ - https://oncocyte.com/graftassure/⑥  
DetermalO™ - https://oncocyte.com/determa-io/⑥  
DetermaCNI™ - https://oncocyte.com/determa-cni/
VitaGraft™, GraftAssure™, DetermalO™, and DetermaCNI™ are trademarks of Oncocyte Corporation.
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#### **Forward-Looking Statements**

Any statements that are not historical fact (including, but not limited to statements that contain words such as "will," "believes," "plans," "anticipates," "expects," "estimates," "may," and similar expressions) are forward-looking statements. These statements include those pertaining to, among other things, future expansion and growth, the Company's land-and-expand strategy to drive commercial adoption of its tests and capture market share, plans to have transplant centers running GraftAssure tests through the end of 2025, projected revenue path, IVD strategy, assumptions regarding regulatory approvals and clearances, timing and planned regulatory submissions, and other statements about the future expectations, beliefs, goals, plans, or prospects expressed by management. Forward-looking statements involve risks and uncertainties, including, without limitation, risks inherent in the development and/or commercialization of diagnostic tests or products, uncertainty in the results of clinical trials or regulatory approvals, the capacity of Oncocyte's third-party supplied blood sample analytic system to provide consistent and precise analytic results on a commercial scale, potential interruptions to supply chains, the need and ability to obtain future capital, maintenance of intellectual property rights in all applicable jurisdictions, obligations to third parties with respect to licensed or acquired technology and products, the need to obtain third party reimbursement for patients' use of any diagnostic tests Oncocyte or its subsidiaries commercialize in applicable jurisdictions, and risks inherent in strategic transactions such as the potential failure to realize anticipated benefits, legal, regulatory or political changes in the applicable jurisdictions, accounting and quality controls, potential greater than estimated allocations of resources to develop and commercialize technologies, or potential failure to maintain any laboratory accreditation or certification. Actual results may differ materially from the results anticipated in these forward-looking statements and accordingly such statements should be evaluated together with the many uncertainties that affect the business of Oncocyte, particularly those mentioned in the "Risk Factors" and other cautionary statements found in Oncocyte's Securities and Exchange Commission (SEC) filings, which are available from the SEC's website. You are cautioned not to place undue reliance on forward-looking statements, which speak only as of the date on which they were made. Oncocyte undertakes no obligation to update such statements to reflect events that occur or circumstances that exist after the date on which they were made, except as required by law.

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## ONCOCYTE CORPORATION CONDENSED CONSOLIDATED BALANCE SHEETS (In thousands, except per share data)

	June 30, 2024		December 31, 2023	
	(U:	naudited)		
ASSETS				
CURRENT ASSETS				
Cash and cash equivalents	\$	9,256	\$	9,432
Accounts receivable, net of allowance for credit losses of \$1 and \$5, respectively		85		484
Prepaid expenses and other current assets		595		643
Assets held for sale		32		139
Total current assets		9,968		10,698
NONCURRENT ASSETS				
Right-of-use and financing lease assets, net		2,591		1,637
Machinery and equipment, net, and construction in progress		3,347		3,799
Intangible assets, net		56,551		56,595
Restricted cash		1,700		1,700
Other noncurrent assets		563		463
TOTAL ASSETS	\$	74,720	\$	74,892
LIABILITIES AND SHAREHOLDERS' EQUITY				
CURRENT LIABILITIES				
Accounts payable	\$	1.051	\$	953
Accrued compensation	•	1,309	•	1,649
Accrued royalties		1.116		1.116
Accrued expenses and other current liabilities		379		452
Accrued severance from acquisition		2,314		2,314
Right-of-use and financing lease liabilities, current		1,029		665
Current li abilities of discontinued operations		-,		45
Total current liabilities		7,198		7,194
NONCURRENT LIABILITIES				
Right-of-use and financing lease liabilities, noncurrent		2,638		2,204
Contingent consideration liabilities		42,181		39,900
TOTAL LIABILITIES		52,017		49,298
Complement and continue in				
Commitments and contingencies				
Series A Redeemable Convertible Preferred Stock, no par value; stated value \$1,000 per share; 5 shares issued and				
outstanding at December 31, 2023; aggregate liquidation preference of \$5,296 as of December 31, 2023				5,126
SHAREHOLDERS' EQUITY				
Preferred stock, no par value, 5,000 shares authorized; no shares issued and outstanding		-		-
Common stock, no par value, 230,000 shares authorized; 13,368 and 8,261 shares issued and outstanding at				
June 30, 2024 and December 31, 2023, respectively		326,201		310,295
Accumulated other comprehensive income		37		49
Accumulated deficit		(303,535)		(289,876
Total shareholders' equity		22,703		20,468
TOTAL LIABILITIES AND SHAREHOLDERS' EQUITY	\$	74,720	\$	74,892

### ONCOCYTE CORPORATION UNAUDITED CONDENSED CONSOLIDATED STATEMENTS OF OPERATIONS (In thousands, except per share data)

	Three Months Ended June 30,			Six Months Ended June 30,				
		2024		2023		2024		2023
Net revenue	\$	104	\$	463	\$	280	\$	760
Cost of revenues		32		169		141		434
Cost of revenues - amortization of acquired intangibles		22		22		44		44
Gross profit		50_		272		95		282
Operating expenses:								
Research and development		2,453		2,435		4,765		4,562
Sales and marketing		853		805		1,699		1,500
General and administrative		2,407		3,531		5,080		6,943
Change in fair value of contingent consideration		(1,031)		1,795		2,281		(16,512)
Impairment loss		-		-		-		4,950
Impairment loss on held for sale assets		-		-		169		1,283
Total operating expenses		4,682		8,566		13,994		2,726
Loss from operations		(4,632)		(8,294)		(13,899)		(2,444)
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Other (expenses) income:								
Inter est expense		(8)		(14)		(23)		(25)
Unrealized (loss) gain on marketable equity securities		-		(24)		-		97
Other income (expenses), net		110		(1)		263		(2)
Total other income (expenses)		102		(39)		240		70
Loss from continuing operations		(4,530)		(8,333)		(13,659)		(2,374)
Loss from discontinued operations						<u>-</u> _		(2,926)
Net loss	\$	(4,530)	_\$	(8,333)	\$	(13,659)	\$	(5,300)
Net loss per share:								
Net loss from continuing operations - basic and diluted	\$	(4,587)	\$	(8,644)	\$	(13,922)	\$	(2,915)
Net loss from discontinued operations - basic and diluted	\$	-	\$	-	\$	-	\$	(2,926)
Net loss attributable to common stockholders - basic and diluted	\$	(4,587)	\$	(8,644)	\$	(13,922)	\$	(5,841)
Net loss from continuing operations per share - basic and diluted	\$	(0.36)	\$	(1.07)	\$	(1.32)	\$	(0.41)
Net loss from discontinued operations per share - basic and diluted	\$	-	\$	-	\$		\$	(0.42)
Net loss attributable to common stockholders per share - basic and	\$	(0.36)	\$	(1.07)	\$	(1.32)	\$	(0.83)
Weighted average shares outstanding - basic and diluted		12,870		8,090		10,567		7,030

# ONCOCYTE CORPORATION UNAUDITED CONDENSED CONSOLIDATED STATEMENTS OF CASH FLOWS (In thousands)

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	(52)			296
12	` '	(50)		
	(1 657)			567
,240)	(1,657)	(386)		(4,319
(27)	(85)	(123)		(118
,978)	(6,832)	(9,808)		(15,132
-	123	-		123
(191)	-	(215)		-
`-	-	`- ´		(1,372
(191)	123	(215)		(1,249
.807	13,848	15,807		13,848
(538)	(427)	(538)		(427
,389)	(1,118)	(5,389)		(1,118
(33)	(29)	(33)		(57
<del></del>	12,274	9,847		12,246
	5,565	(176)		(4,135
				23,203
,678	13,503	11,132		19,068
9	3,678		3,678 5,565 (176)	3,678 5,565 (176)

#### Oncocyte Corporation Reconciliation of Non-GAAP Financial Measure Consolidated Adjusted Loss from Operations

Note: In addition to financial results determined in accordance with U.S. generally accepted accounting principles ("GAAP"), this press release also includes a non-GAAP financial measure (as defined under SEC Regulation G). We believe the adjusted amounts are more representative of our ongoing performance. The following is a reconciliation of the non-GAAP measure to the most directly comparable GAAP measure:

	Three Months Ended						
	Ju	me 30,	March 31,		June 30,		
		2024		2024		2023	
	(un	audited)	(unaudited)		(unaudited)		
			(In th	nousands)			
Consolidated GAAP loss from operations	\$	(4,632)	\$	(9,267)	\$	(8,294)	
Stock-based compensation		386		418		816	
Depreciation and amortization expense		326		335		457	
Change in fair value of contingent consideration		(1,031)		3,312		1,795	
Impairment losses		-		-		-	
Impairment loss on held for sale assets		-		169		-	
Consolidated Non-GAAP loss from operations, as adjusted	\$	(4,951)	\$	(5,033)	\$	(5,226)	