UNITED STATES SECURITIES AND EXCHANGE COMMISSION WASHINGTON, DC 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): May 10, 2022

MIMEDX GROUP, INC.

(Exact name of registrant as specified in charter)

Florida (State or other jurisdiction of incorporation)

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

001-35887 (Commission File Number) 26-2792552 (IRS Employer Identification No.)

1775 West Oak Commons Ct., NE, Marietta GA 30062 (Address of principal executive offices) (Zip Code) Registrant's telephone number, including area code: (770) 651-9100

Chec	ck 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 (see General Instruction A.2. below):
	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)
	Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

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

		Name of each exchange
Title of each class	Trading Symbol(s)	on which registered
Common Stock, \$0.001 par value per share	MDXG	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 □

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. \square

Item 7.01 Regulation FD

Timothy R. Wright, MiMedx Chief Executive Officer, and Peter M. Carlson, MiMedx Chief Financial Officer, are expected to attend the Bank of America Securities 2022 Healthcare Conference on behalf of MiMedx Group, Inc. (the "Company"), on May 10, 2022. A copy of the presentation materials made available by the Company in connection with the conference is furnished as Exhibit 99.1 to this Current Report on Form 8-K (this "Current Report") and is incorporated herein by reference.

The information in this Current Report, 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 such information be deemed incorporated by reference in any filing under the Securities Act of 1933, as amended, except as expressly set forth by specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No. Description of Exhibit

Slide Presentation dated May 10, 2022

The cover page from this Current Report on Form 8-K, formatted in Inline XBRL.

SIGNATURES

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

MIMEDX GROUP, INC.

Date: May 10, 2022 By: /s/ Peter M. Carlson

Peter M. Carlson, Chief Financial Officer



DISCLAIMER & CAUTIONARY STATEMENTS

This presentation includes forward-looking statements. Forward-looking statements are subject to risks and uncertainties, and the Company cautions investors against placing undue reliance on such statements. Actual results may differ materially from those set forth in the forward-looking statements. Such forward-looking statements include statements regarding:

- · future sales or sales growth;
- the Company's expectations regarding its mdHACM product's potential use as a safe and effective treatment option, and that it may be an effective treatment for persons battling inflammatory conditions; the Company's plans for meetings with the U.S. Food & Drug Administration (FDA), and planned biologics license application (BLA) submissions to the FDA, and their timing; plans for future clinical trials, including the Company's decision to pursue or not pursue, and their timing;
- the effectiveness of amniotic tissue as a therapy for any particular indication or condition;
- · estimates of potential market size for the Company's current and future products;
- · plans for expansion outside of the U.S.;
- · expected spending on clinical trials and research and development;
- the Company's long-term strategy for value creation, the status of its pipeline products, expectations for future products, and expectations for future growth;





DISCLAIMER & CAUTIONARY STATEMENTS

Additional forward-looking statements may be identified by words such as "believe," "expect," "may," "plan," "potential," "will," "preliminary," and similar expressions, and are based on management's current beliefs and expectations. Forward-looking statements are subject to risks and uncertainties, and the Company cautions investors against placing undue reliance on such statements. Actual results may differ materially from those set forth in the forward-looking statements. Factors that could cause actual results to differ from expectations include:

- future sales are uncertain and are affected by competition, access to customers, patient access to healthcare providers, and many other factors;
- the results of a clinical trial or trials may not demonstrate that the product is safe or effective, or may have little or no statistical value; the Company may change its plans due to unforeseen circumstances, and delay or alter the timeline for future trials, analyses, or public announcements; the timing of any meeting with the FDA depends on many factors and is outside of the Company's control, and the results from any meeting are uncertain; a BLA submission requires a number of prerequisites, including favorable study results and statistical support, and completion of a satisfactory FDA inspection of the Company's manufacturing facility or facilities; plans for future clinical trials depend on the results of pending clinical trials, discussion with the FDA, and other factors; and conducting clinical trials is a time-consuming, expensive, and uncertain process;
- the future market for the Company's products can depend on regulatory approval of such products, which might not occur at all
 or when expected, and is based in part on assumptions regarding the number of patients who elect less acute and more acute
 treatment than the Company's products, market acceptance of the Company's products, and adequate reimbursement for such
 therapies;
- the process of obtaining regulatory clearances or approvals to market a biological product or medical device from the FDA or similar regulatory authorities outside of the U.S. is costly and time consuming, and such clearances or approvals may not be granted on a timely basis, or at all, and the ability to obtain the rights to market additional, suitable products depends on negotiations with third parties which may not be forthcoming;
- whether there is full access to hospitals and healthcare provider facilities, as a continuation or escalation of access restrictions or lockdown orders resulting from the ongoing COVID-19 pandemic; and
- · expected spending can depend in part on the results of pending clinical trials.

The Company describes additional risks and uncertainties in the Risk Factors section of its most recent annual report and quarterly reports filed with the Securities and Exchange Commission. Any forward-looking statements speak only as of the date of this presentation and the Company assumes no obligation to update any forward-looking statement.





LEADING PRODUCT PORTFOLIO POSITIONED FOR GROWTH

MDXG

\$257.5M

TTM Net Sales

83.1%

TTM Gross Margin

(\$12.4M)TTM Net Loss

\$12.0M TTM Adjusted EBITDA¹

13.4%

Year-over-year Revenue growth in Wound Care & Surgical business²

800+

Employees³

\$481M

Market Cap⁴

\$75.7M

Cash at 3/31/22

2,000,000+

Allografts Distributed⁵

Purion.

EPIFIX°

AMNIOFIX® EPICORD® AMNIOCORD® Reimbursement coverage, U.S.:

300M+

lives

300M+

people worldwide suffering from hip and knee OA⁶ **30M** (U.S.) with diabetes7

2.9M chronic wounds8 In a recent peer-reviewed study, the average cost/episode with EPIFIX was

~\$3000 less

versus other advanced treatments9

42%

of the low risk-of-bias studies in AHRQ assessment were on MIMEDX products10

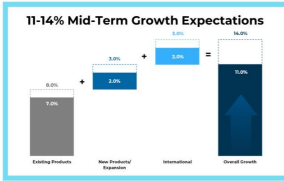




COMPELLING INVESTMENT THESIS



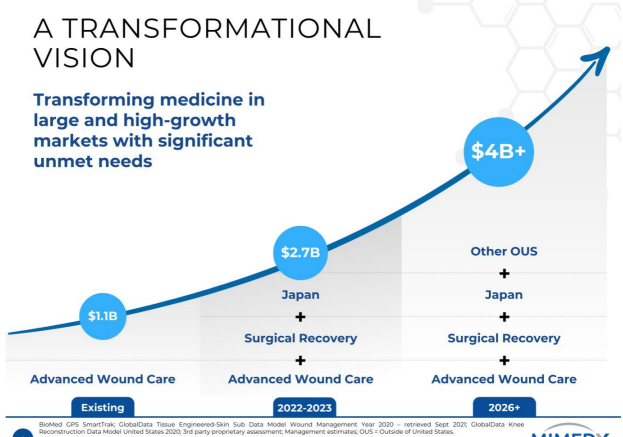




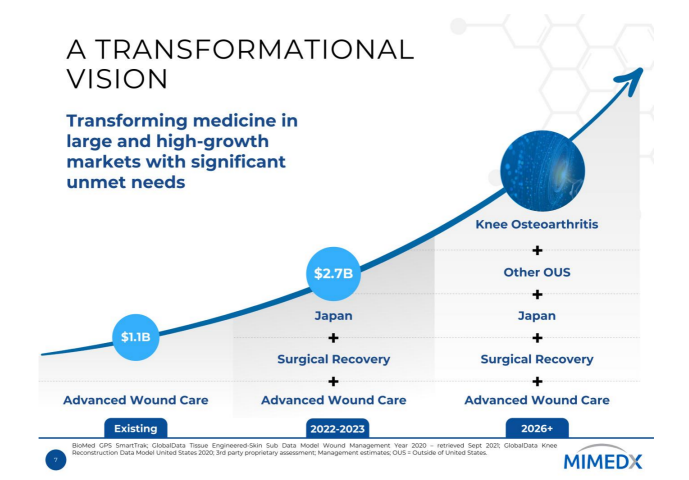


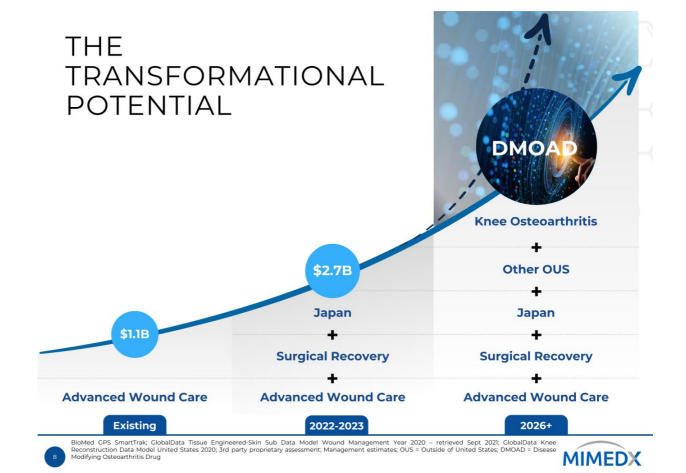




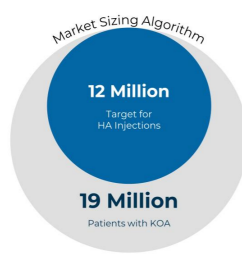








SIGNIFICANT UNMET CLINICAL NEED IN KNEE OSTEOARTHRITIS



Multiple factors drive overall transformation

Value Multipliers

- Product Label
- Dosing Regimen
- Bilateral Application
- Prophylactic Use
- Place in Treatment Algorithm
- Clinical Trial Results
- DMOAD





DMOAD substantially amplifies market opportunity



GlobalData: 2020 Orthopedic Devices Knee Reconstruction US (2015-2030); GlobalData: Viscosupplementation Model (HA) U.S. (2015-2030); KOA = Knee Osteoarthritis; HA = Hyaluronic Acid



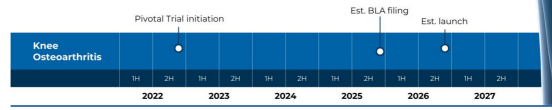
mdHACM HOLDS POTENTIAL TO REDUCE PAIN AND INCREASE FUNCTION IN KOA

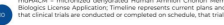
Phase 2B study did not meet primary endpoints, but demonstrated statistically significant and clinically meaningful improvement within Pre-Interim analysis cohort

190-patient Cohort	3-months	6-months
WOMAC Pain	p=0.032	p=0.009
WOMAC Function	p=0.046	p=0.009
WOMAC Total	p=0.038	p=0.008

Plan to commence KOA Clinical Trial Program in 2022

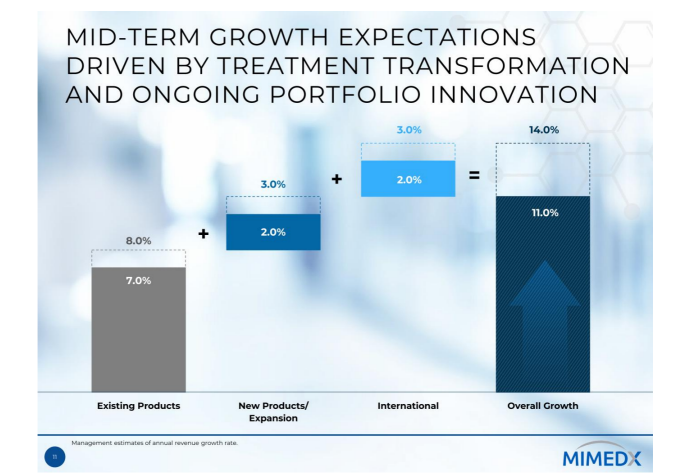
Anticipate BLA filing in late-2025 with greater probability of success





mdHACM = micronized dehydrated Human Amnion Chorion Membrane; WOMAC = Western Ontario and McMaster Universities Osteoarthritis Index; BLA = Biologics License Application; Timeline represents current plans and estimates only. Actual results and timing may differ materially. There can be no assurance that clinical trials are conducted or completed on schedule, that trial results are favorable, or that we obtain regulatory approval for our products and indications.





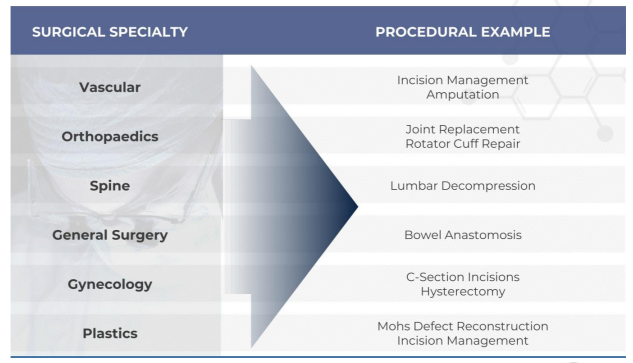




 ${\tt BioMed\ GPS\ SmartTrak;\ 3rd\ party\ proprietary\ assessment;\ Management\ estimates}$



MULTIPLE OPPORTUNITIES TO EXPAND EXISTING PROCEDURE BASE









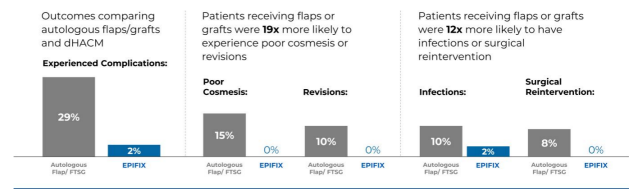
- Expand Reach in O.R.
- Procedural Training
- KOL Development by Specialty
- New Product Launches
- Clinical & Economic Evidence



OPPORTUNITIES TO EXTEND LEADERSHIP IN DIFFERENTIATED CLINICAL EVIDENCE



Mohs Defect Repair with Dehydrated Human Amnion/Chorion Membrane





(1) Toman J, Michael GM, Wisco OJ, Adams JR, Hubbs BS. Mohs Defect Repair with Dehydrated Human Amnion/Chorion Membrane. Facial Plast Surg Aesthet Med. 2021 Oct 29. doi: 10.1089/fpsam.2021.0167. Epub ahead of print. PMID: 34714143. FTSG = Full Thickness Skin Grafts



2022 LAUNCHES EXPAND PLACENTAL PORTFOLIO



AMNIOEFFECT™

Wide range of sizes up to 9 cm x 20 cm

Improved handling for minimally invasive procedures



Placental Collagen Matrix

Particulate format fulfills key portfolio gap

Retains key extracellular matrix components

Anticipate two new, organic products launched per year; future year new product launches would present additional upside opportunity

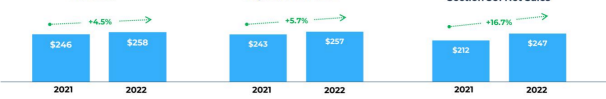




ADVANCED WOUND CARE CONTINUES TO EXHIBIT STRONG DOUBLE-DIGIT GROWTH

Results for the Three Months Ended March 31 (\$M)





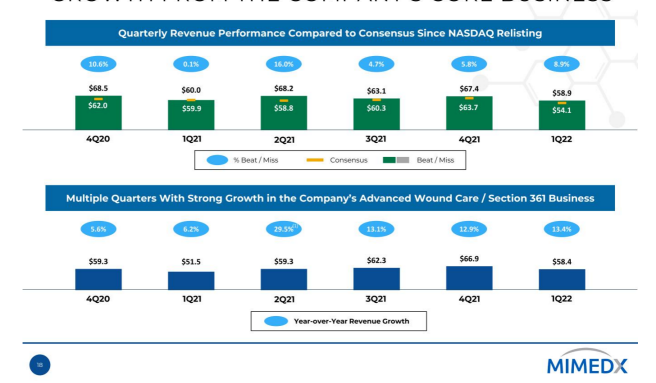
MIMEDX Expects Sales of Its Advanced Wound Care / Section 361 Products to Grow 11% to 14% in 2022, with Growth Expected to Accelerate as the Year Progresses



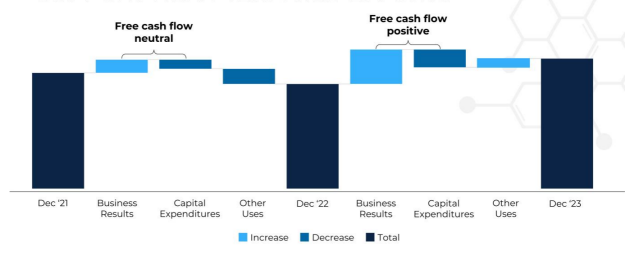
(1) Adjusted net sales excludes revenue recognized from cash collections on remaining contracts. Adjusted net sales is a non-GAAP measurement. (2) Section 361 includes Tissue + Cord sales. Section 351 includes Micronized + Particulate sales. (3) TTM refers to the trailing twelve months ended March 31, 2022, and is calculated for any measure by adding the results for the full year ended December 31, 2021 to the results for the quarter ended March 31, 2022 and subtracting the results for the quarter ended March 31, 2021.



CONSISTENT OUTPERFORMANCE COMPARED TO CONSENSUS WITH STRONG, SUSTAINED GROWTH FROM THE COMPANY'S CORE BUSINESS



EXISTING CASH LEVELS ARE SUFFICIENT TO SUPPORT NEAR-TERM R&D EFFORTS



Cash and cash equivalents at December 31, 2021 = \$87.1 million

Expect two clinical trials for Knee OA indication to cost less than \$30 million; incurred over three years Over the 12 – 15 months ending December 2022, we continue to expect:

- · Base business to be cash flow neutral
- Overall revenue to return to levels consistent with those prior to end of Enforcement Discretion



Business Results represents expected Adjusted EBITDA. Other Uses include debt service, and investigation, restatement and related expenses.



PIONEER IN PLACENTAL BIOLOGICS



Distinct drivers of significant shareholder value with current and future growth potential





2022 OBJECTIVES SUPPORT CURRENT AND FUTURE GROWTH POTENTIAL

R&D	 □ Initiate Phase 3 KOA Clinical Studies □ Increase Product Vitality Index □ Advance body of scientific evidence
Operations	 □ Implement CGMP throughout supply chain □ Leverage cost base through production efficiencies □ Optimize quality, processes and scale
Commercial	 Achieve sustainable double-digit growth target Expand international footprint, with initial launch in Japan Launch two new products – AMNIOEFFECT™ and PCM











SUMMARY BALANCE SHEETS

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22
Assets								
Cash and Cash Equivalents	48.2	109.6	95.8	84.7	85.0	90.6	87.1	75.7
Accounts Receivable, net	30.1	33.0	35.4	35.4	37.2	36.5	40.4	37.7
Inventory, net	10.6	11.0	10.4	11.6	10.1	11.2	11.4	13.2
Other Current Assets	18.7	17.9	19.0	18.3	15.4	3.6	9.6	9.3
Total Current Assets	107.6	171.5	160.6	150.0	147.7	141.9	148.5	135.9
Property and Equipment	10.8	10.3	11.4	11.0	10.3	9.9	9.2	8.8
Other Assets	32.5	31.5	30.0	29.8	29.1	28.7	30.2	29.7
Total Assets	150.9	213.3	202.0	190.8	187.1	180.5	187.9	174.4
Liabilities and Stockholders' Equity (Deficit)								
Current Liabilities	63.7	57.3	59.2	55.4	50.6	41.7	42.4	36.6
Long Term Debt, net	61.5	47.6	47.7	47.8	47.9	48.0	48.1	48.2
Other Liabilities	2.9	4.4	3.7	3.6	3.3	4.1	4.9	4.6
Total Liabilities	128.1	109.3	110.6	106.8	101.8	93.8	95.4	89.4
Convertible Preferred Stock	0.0	91.1	91.6	92.0	92.5	92.5	92.5	92.5
Stockholders' Equity (Deficit)	22.9	12.9	(0.2)	(8.0)	(7.2)	(5.8)	0.1	(7.4)
Total Liabilities and Stockholders' Equity (Deficit)	150.9	213.3	202.0	190.8	187.1	180.5	187.9	174.4







SUMMARY INCOME STATEMENTS

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22
Net Sales	53.6	64.3	68.6	60.0	68.2	63.1	67.4	58.9
Cost of Sales	8.2	10.3	10.8	9.7	12.8	10.1	10.7	9.9
Gross Profit	45.4	54.0	57.8	50.3	55.4	53.0	56.7	49.0
Research & Development	2.3	3.4	3.4	4.3	4.1	4.3	4.6	6.0
Selling, General, and Administrative	37.3	48.0	48.8	45.4	53.6	46.3	53.1	49.6
Investigation, Restatement, and Related	11.4	12.0	20.4	7.2	(2.1)	3.2	(4.5)	2.6
Amortization of Intangible Assets	0.3	0.3	0.3	0.2	0.2	0.2	0.2	0.2
Impairment of Intangible Assets	0.0	0.0	1.0	0.0	0.0	0.0	0.1	0.0
Operating (Loss) Income	(5.9)	(9.7)	(16.1)	(6.8)	(0.4)	(1.0)	3.3	(9.3)
Loss on Extinguishment of Debt	0.0	(8.2)	0.0	0.0	0.0	0.0	0.0	0.0
Interest Expense, net	(2.6)	(1.5)	(1.5)	(1.5)	(1.4)	(1.0)	(1.2)	(1.1)
Pretax (Loss) Income	(8.4)	(19.4)	(17.6)	(8.3)	(1.8)	(2.0)	2.1	(10.4)
Income Tax Provision Benefit (Expense)	0.0	0.0	1.0	(0.1)	0.0	(0.3)	0.1	(O.1)
Net (Loss) Income	(8.5)	(19.4)	(16.6)	(8.4)	(1.8)	(2.3)	2.2	(10.5)

Note: figures don't add to subtotals due to immaterial rounding differences





SUMMARY CASH FLOW STATEMENTS

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22
Net (Loss) Income	(8.5)	(19.4)	(16.6)	(8.4)	(1.8)	(2.3)	2.2	(10.5)
Share-Based Compensation	4.4	3.7	3.9	3.2	4.1	3.8	3.6	4.0
Depreciation	1.4	1.5	1.3	1.2	1.3	0.9	1.0	0.9
Other Non-Cash Effects	1.3	9.5	1.7	1.1	0.9	0.6	0.7	0.6
Changes in Assets	2.9	(1.8)	(6.2)	0.1	1.9	11.0	(9.5)	0.7
Changes in Liabilities	(4.7)	1.9	5.5	(3.9)	(4.8)	(7.6)	(1.3)	(5.9)
Net Cash Flows (Used in) Provided By Operating Activities	(3.1)	(4.6)	(10.4)	(6.7)	1.6	6.4	(3.3)	(10.2)
Purchases of Property and Equipment	(0.4)	(0.7)	(2.2)	(1.9)	(0.4)	(0.6)	(0.3)	(0.1)
Patent Application Costs	(0.1)	0.0	(O.1)	(0.2)	(0.0)	(0.1)	(0.0)	(0.1)
Other	0.0	0.0	0.0	0.0	0.0	0.1	0.0	0.0
Net Cash Flows Used in Investing Activities	(0.5)	(0.7)	(2.3)	(2.1)	(0.4)	(0.6)	(0.3)	(0.1)
Preferred Stock Net Proceeds	0.0	93.4	(0.8)	0.0	0.0	0.0	0.0	0.0
Proceeds from Term Loan	10.0	49.5	0.0	0.0	0.0	0.0	0.0	0.0
Repayment of Term Loan	(10.9)	(72.0)	0.0	0.0	0.0	0.0	0.0	0.0
Prepayment Premium on Term Loan	0.0	(1.4)	0.0	0.0	0.0	0.0	0.0	0.0
Deferred Financing Cost	0.0	(2.8)	(0.3)	0.0	0.0	0.0	0.0	0.0
Stock Repurchased for Tax Withholdings on Vesting of Restricted Stock	(0.8)	(0.1)	0.0	(3.2)	(1.4)	(0.2)	0.0	(1.2)
Proceeds from Exercise of Stock Options	0.0	0.1	0.0	0.9	0.5	0.0	0.0	0.2
Net Cash Flows (Used in) Provided By Financing Activities	(1.8)	66.7	(1.1)	(2.3)	(0.9)	(0.2)	0.0	(1.0)
Beginning Cash Balance	53.5	48.2	109.6	95.8	84.7	85.0	90.6	87.1
Change in Cash	(5.3)	61.4	(13.8)	(11.1)	0.3	5.6	(3.5)	(11.4)
Ending Cash Balance	48.2	109.6	95.8	84.7	85.0	90.6	87.1	75.7

lote: certain figures may not foot due to rounding.





REVENUE DETAIL

Quarter								Trailing 12	2 Months			
(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q21	3Q21	4Q21	1Q22
Advanced Wound Care / Section 361 ¹	45.8	55.1	59.4	51.5	59.3	62.3	66.9	58.5	225.3	232.5	240.0	247.0
Section 351 ¹	6.1	8.2	8.7	8.2	8.6	0.5	0.3	0.4	33.7	26.0	17.6	9.8
Other ²	1.7	1.0	0.5	0.3	0.3	0.3	0.1	0.0	2.1	1.4	1.0	0.7
Net Sales	\$ 53.6	\$ 64.3	\$ 68.5	\$ 60.0	\$ 68.2	\$ 63.1	\$ 67.4	\$ 58.9	\$261.1	\$259.9	\$258.6	\$257.5





NON-GAAP METRICS RECONCILIATION

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22
Net Sales – Reported	53.6	64.3	68.6	60.0	68.2	63.1	67.3	58.9
Less: Revenue Transition Impact ¹	(1.7)	(1.0)	(0.5)	(0.3)	(0.3)	(0.3)	(0.1)	0.0
Adjusted Net Sales	51.9	63.3	68.1	59.7	67.9	62.8	67.2	58.9
1								\\ /
Gross Profit	45.4	54.0	57.8	50.3	55.4	53.0	56.7	49.0
Less: Revenue Transition Impact ¹	(1.5)	(0.9)	(0.4)	(0.2)	(0.3)	(0.3)	(O.1)	0.0
Adjusted Gross Profit	44.0	53.1	57.4	50.1	55.1	52.7	56.6	49.0
Adjusted Gross Margin	84.8%	83.9%	84.3%	83.9%	81.3%	83.9%	84.2%	83.1%
Adjusted EBITDA	11.7	7.8	10.8	5.0	3.1	7.0	3.6	(1.7)
Less: Capital Expenditures	(0.4)	(0.7)	(2.2)	(1.9)	(0.4)	(0.6)	(0.3)	(0.1)
Less: Patent Application Costs	(O.1)	0.0	(O.1)	(0.2)	(0.0)	(O.1)	(0.0)	(O.1)
Adjusted Free Cash Flow	11.2	7.1	8.5	2.9	2.7	6.3	3.3	(1.9)



npact of revenue transition includes cash collected related to the remaining contracts and cost of sales recognized on those collections, as applicable. For a discussion of the revenue transition and the defined terms refer me. 8, Notes to the Consolidated Financial Statements in the MiMedx Group, Inc. Form 10-K for the years ended December 31, 2021 and 2020, and the respective Form 10-Qs for the noted quarterly periods. Note: certain figure asy not foot due to rounding.



ADJUSTED EBITDA RECONCILIATION

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22
Net (Loss) Income	(8.5)	(19.4)	(16.6)	(8.4)	(1.8)	(2.3)	2.2	(10.5)
Depreciation & Amortization	1.7	1.8	1.6	1.4	1.5	1.1	1.1	1.0
Interest Expense	2.6	1.5	1.5	1.5	1.4	1.0	1.2	1.1
Loss on Extinguishment of Debt	0.0	8.2	0.0	0.0	0.0	0.0	0.0	0.0
Income Tax	0.0	0.0	(1.0)	0.1	(0.0)	0.3	(0.1)	0.1
EBITDA	(4.2)	(7.9)	(14.5)	(5.5)	1.1	0.0	4.4	(8.3)
Investigation, Restatement & Related	11.4	12.0	20.4	7.2	(2.1)	3.2	(4.5)	2.6
Impairment of Intangible Assets	0.0	0.0	1.0	0.0	0.0	0.0	0.1	0.0
Share-Based Compensation	4.4	3.7	3.9	3.2	4.1	3.8	3.6	4.0
Adjusted EBITDA ¹	11.7	7.8	10.8	5.0	3.1	7.0	3.6	(1.7)



) Adjusted EBITDA is a non-GAAP measure consisting of GAAP net loss excluding; (i) depreciation, (ii) amortization of intangibles, (iii) interest expense, (iv) income tax provision, (v) costicured in connection with the Audit Committee Investigation and Restatement, (vi) impairment of intangible assets, and (vii) share-based compensation. Refer to Item 7 of our Annua eport on Form 10-K for the year ended December 31, 2021, filed with the SEC. Note: certain figures my not foot due to rounding.

