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): August 10, 2022

MIMEDX GROUP, INC. (Exact name of registrant as specified in charter)

Florida (State or other jurisdiction of incorporation)

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

Check the appropriate	box below if the Form 8-K filing is intended to sime	ultaneously satisfy the filing obligation of the registrant under any of	the following provisions (see General Instruction A.2. below):
□ Soliciting mater□ Pre-commencer	nications pursuant to Rule 425 under the Securities A ial pursuant to Rule 14a-12 under the Exchange Act nent communications pursuant to Rule 14d-2(b) unden nent communications pursuant to Rule 13e-4(c) under	(17 CFR 240.14a-12) er the Exchange Act (17 CFR 240.14d-2(b))	
Securities registered p	oursuant to Section 12(b) of the Act:		
	Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Со	mmon Stock, \$0.001 par value per share	MDXG	The Nasdaq Stock Market LLC
chapter). Emerging growth con	npany □	. ,	of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter) or Rule 12b-2 of this chapter (Securities Exchange Act of 1934 (
he Exchange Act.	ir company, indicate by check mark it the registrant i	has elected not to use the extended transition period for complying w	thi any new of revised infancial accomining standards provided phistianic to Section 13(a)

Regulation FD Item 7.01

Peter M. Carlson, MiMedx Chief Financial Officer and Jack Howarth, Senior Vice President of Investor Relations, are expected to attend the Canaccord Genuity Group, Inc. 42nd Annual Growth Conference on behalf of MiMedx Group, Inc. (the "Company"), on August 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. 99.1

Description of Exhibit Slide Presentation dated August 10, 2022

104 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.

August 10, 2022

By: /s/ Peter M. Carlson Peter M. Carlson, Chief Financial Officer



DISCLAIMER & CAUTIONARY STATEMENTS

This presentation includes forward-looking statements.

Statements regarding:

- (i) future sales or sales growth;
- (ii) our 2022 financial outlook and expectations for future financial results, including net sales and levels of selling, general and administrative expense;
- (iii) our expectations regarding the timing of clinical programs and trials;
- (iv) our expectations regarding the timing of new product launches; and
- (v) the effectiveness of amniotic tissue as a therapy for any particular indication or condition.

Additional forward-looking statements may be identified by words such as "believe," "expect," "may," "plan," "goal," "outlook," "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.





DISCLAIMER & CAUTIONARY 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:

- (i) future sales are uncertain and are affected by competition, access to customers, patient access to healthcare providers, and many other factors:
- (ii) the status, timing, results and expected results of the Company's clinical trials and planned regulatory submissions, and our expectations regarding our ability to potentially accelerate the timing of any trial or regulatory submission, depend on a number of factors including favorable trial results, patient access, and our ability to manufacture in accordance with Current Good Manufacturing Practices (CGMP) and appropriate chemistry and manufacturing controls;
- (iii) the Company may change its plans due to unforeseen circumstances, or delays in analyzing and auditing results, and may delay or alter the timeline for future trials, analyses, or public announcements;
- (iv) our access to hospitals and health care provider facilities could be restricted as a result of the ongoing COVID-19 pandemic or other factors;
- (v) the results of scientific research are uncertain and may have little or no value;
- (vi) our ability to sell our products in other countries depends on a number of factors including adequate levels of reimbursement, market acceptance of novel therapies, and our ability to build and manage a direct sales force or third party distribution relationship;
- (vii) the effectiveness of amniotic tissue as a therapy for particular indications or conditions is the subject of further scientific and clinical studies;
- (viii) we may alter the timing and amount of planned expenditures for research and development based on the results of clinical trials and other regulatory developments. 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



\$256.3M TTM Net Sales

83.4%

(\$21.5M) TTM Gross Margin TTM Net Loss

\$8.0M

TTM Adjusted EBITDA¹

11.6%

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

800+

\$582M Market Cap⁴ Employees³

\$72.5M

Cash at 6/30/22

2,000,000+

Allografts Distributed⁵

Purion.

AMNIOFIX° **EPIFIX**° EPICORD° AMNIOCORD **50**+

Clinical & Scientific **Publications** 100%

National Payor Coverage for DFUs6

300M+

people worldwide suffering from hip and knee OA⁷ **30M** (U.S.) with diabetes8

2.9M chronic wounds9 In a recent peer-reviewed study, the average cost/episode with EPIFIX was

~\$3000 less

versus other advanced treatments10

42%

of the low risk-of-bias studies in AHRQ assessment were on MIMEDX products¹¹

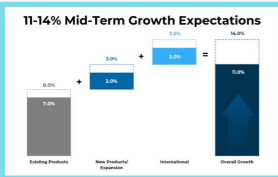


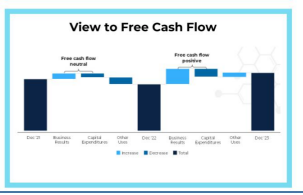


COMPELLING INVESTMENT THESIS













MIMEDX IS ON STRATEGY WITH STRONG COMMERCIAL MOMENTUM

Fourth Consecutive Quarter of Double Digit Revenue Growth in Continuing Portfolio

- · Achieved strong year-over-year increase in key focus area of Surgical Recovery
- Two new product launches on track for September
- Preparing for Japan launch of PURION® engineered EPIFIX® later this year, as early as September; Reimbursement acceptance imminent
- Scheduled Type B RMAT meeting with FDA in the third quarter; On track to enroll the first patient in our Knee Osteoarthritis clinical trial by year end
- · Business is generating the cash needed to fuel future investments

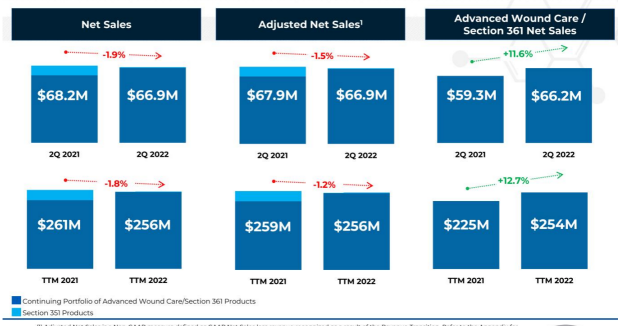






MIMEDX IS ON STRATEGY WITH STRONG COMMERCIAL MOMENTUM

Maintaining expectations of 11% to 14% growth in continuing portfolio in 2022



(1) Adjusted Net Sales is a Non-GAAP measure defined as GAAP Net Sales less revenue recognized as a result of the Revenue Transition. Refer to the Appendix for reconciliation of GAAP Net Sales to Adjusted Net Sales.



2Q 2022 FINANCIAL HIGHLIGHTS

Net Sales

\$66.9M

11.6% growth in continuing portfolio

Gross Margin

82.3%

Net Sales in TTM



Continuing Portfolio of AWC Products
Section 351 Products

2022 TTM demonstrates four consecutive quarters of double-digit revenue growth in continuing portfolio

Cash at 6/30/2022

\$72.5M

Adjusted EBITDA

\$(1.0)M

Includes:

- \$2.1M expense for 2022 annual meeting
- \$2.2M bad debt expense

Net Loss

\$(10.9)M

Includes

 \$3.2M charge for Investigation, Restatement and Related Expenses Estimated 2022 Revenue Growth

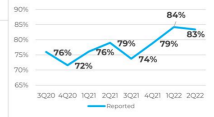
11-14%

Free Cash Flow

\$(1.4)M

Expect to be Free Cash Flow neutral in 2022

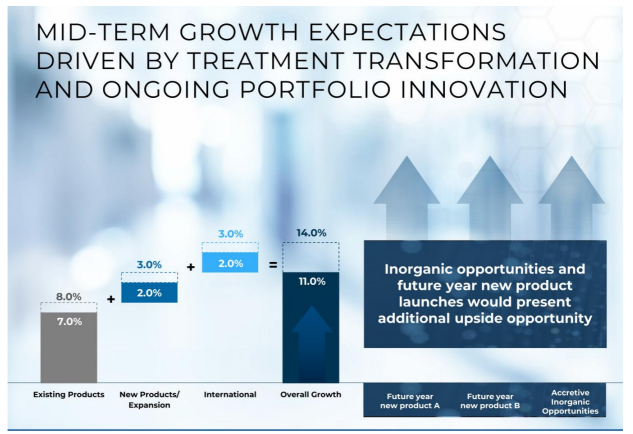
SG&A as % of Adjusted Net Sales



Expect the level of SG&A as percent of adjusted net sales to decline

AWC = Advanced Wound Care





Management estimates of annual revenue growth rate.







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



BODY OF EVIDENCE ACROSS SPECIALTIES AND PROCEDURES

Tissue augmentation

Barrier properties

Surgical closure

Posterior Lumbar Instrumentation⁷

4 of 5 had easily detachable tissue during epidural re-exploration

ACL Reconstruction¹¹

Early Maturation of hamstring autograft seen on MRI at 3 months and 6 months

Mohs⁸

Without EPIFIX:

19X rate poor cosmesis/revision

12X rate infection/reintervention

Prostatectomy¹²

Faster Recovery:

1.5X return to continence **2.5X** return to potency

Urethral Strictures5

67% Success

despite multiple prior recurrences

DFU: 5 RCTs14-23

90%+ closure rates

Burn⁶

Faster Resolution & Lower HTS & Contracture

vs. STSG in pediatrics

Anastomotic Leak¹⁰

74%+ reduction in leak rate

Endometriosis9

14 of 15 No adhesions

in 2nd look patients where AMNIOFIX was placed

VLU: RCT13

70%+ closure rates

I. Gellborn AC, Issn A, The Use of Dehydrated Human Ammion Chorison Membrane Allogail Injection for the Treatment of Translocapatity or Artivitis. A Case Series Insolving 4D Patricts. PM R. 2017;5(12):1236-1243; 2. Alder IX, Hums S, Hubbs S, Kor X, Himsen RB, Masson Charles Chemistra Series of Patricts and Charles Chemistra Series (Patricts Chemistra Series Chemist





2022 LAUNCHES EXPAND PLACENTAL PORTFOLIO



Wide range of sizes up to 9 cm x 20 cm

Improved handling for surgical procedures

Launch September 2022





Particulate offers versatile form factor for use as paste or powder

Retains key extracellular matrix components

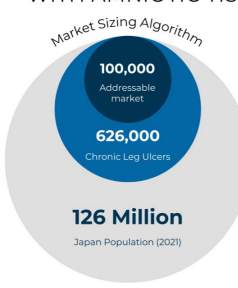
Launch September 2022

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





LARGE POTENTIAL AS FIRST TO MARKET IN JAPAN WITH AMNIOTIC TISSUE FOR WOUND TREATMENT



Total Addressable Market



Key Milestones to Launch

- First patient application
- Finalize & train distributor partner
- Leverage KOL network to facilitate market adoption

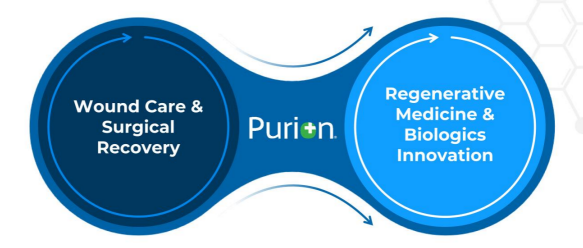
Anticipate launch as early as September; Reimbursement acceptance imminent



GlobalData Tissue Engineered-Skin Sub Data Model Wound Management Year 2020 – retrieved Sept 2021; Management estimates



PIONEER IN PLACENTAL BIOLOGICS



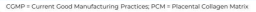
Distinct drivers of significant shareholder value with current and future growth potential





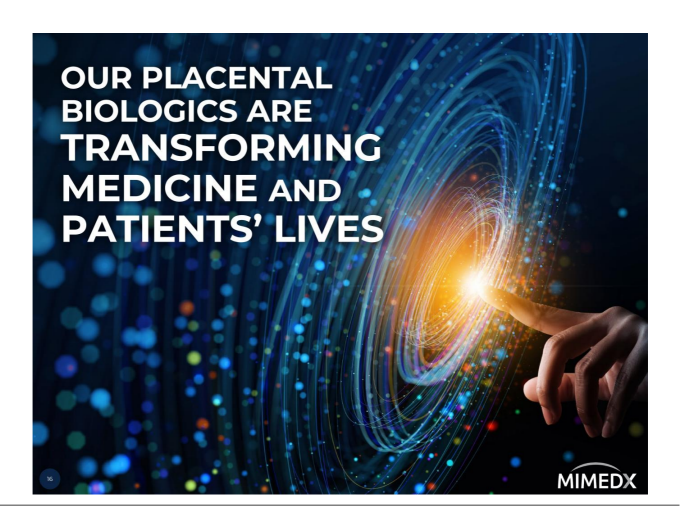
2022 OBJECTIVES SUPPORT CURRENT AND FUTURE GROWTH POTENTIAL

R&D	 □ Initiate KOA Clinical Trial Program □ 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 AXIOFILL™











SUMMARY BALANCE SHEETS

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q22
Assets									
Cash and Cash Equivalents	48.2	109.6	95.8	84.7	85.0	90.6	87.1	75.7	72.5
Accounts Receivable, net	30.1	33.0	35.4	35.4	37.2	36.5	40.4	37.7	37.7
Inventory	10.6	11.0	10.4	11.6	10.1	11.2	11.4	13.2	13.4
Other Current Assets	18.7	17.9	19.0	18.3	15.4	3.6	9.6	9.3	7.4
Total Current Assets	107.6	171.5	160.6	150.0	147.7	141.9	148.5	135.9	131.0
Property and Equipment, net	10.8	10.3	11.4	11.0	10.3	9.9	9.2	8.8	8.3
Other Assets	32.5	31.5	30.0	29.8	29.1	28.7	30.2	29.7	29.4
Total Assets	150.9	213.3	202.0	190.8	187.1	180.5	187.9	174.4	168.7
Liabilities and Stockholders' Equity (Deficit)									
Current Liabilities	63.7	57.3	59.2	55.4	50.6	41.7	42.4	36.6	37.1
Long Term Debt, net	61.5	47.6	47.7	47.8	47.9	48.0	48.1	48.2	48.4
Other Liabilities	2.9	4.4	3.7	3.6	3.3	4.1	4.9	4.6	4.3
Total Liabilities	128.1	109.3	110.6	106.8	101.8	93.8	95.4	89.4	89.8
Convertible Preferred Stock	0.0	91.1	91.6	92.0	92.5	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)	(13.6)
Total Liabilities and Stockholders' Equity (Deficit)	150.9	213.3	202.0	190.8	187.1	180.5	187.9	174.4	168.7

Note: Some figures may not add to subtotals due to immaterial rounding differences.





SUMMARY INCOME STATEMENTS

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q22
Net Sales	53.6	64.3	68.6	60.0	68.2	63.1	67.4	58.9	66.9
Cost of Sales	8.2	10.3	10.8	9.7	12.8	10.1	10.8	9.9	11.8
Gross Profit	45.4	54.0	57.8	50.3	55.4	53.0	56.6	49.0	55.1
Research & Development	2.3	3.4	3.4	4.3	4.1	4.3	4.6	6.0	5.5
Selling, General, and Administrative	37.3	48.0	48.8	45.4	53.6	46.3	53.1	49.6	55.8
Investigation, Restatement, and Related	11.4	12.0	20.4	7.2	(2.1)	3.2	(4.5)	2.6	3.2
Amortization of Intangible Assets	0.3	0.3	0.3	0.2	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	0.0
Operating (Loss) Income	(5.9)	(9.7)	(16.1)	(6.8)	(0.4)	(1.0)	3.3	(9.3)	(9.6)
Loss on Extinguishment of Debt	0.0	(8.2)	0.0	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)	(1.2)
Pretax (Loss) Income	(8.4)	(19.4)	(17.6)	(8.3)	(1.8)	(2.0)	2.1	(10.4)	(10.8)
Income Tax Provision Benefit (Expense)	0.0	0.0	1.0	(O.1)	0.0	(0.3)	0.1	(O.1)	(0.1)
Net (Loss) Income	(8.5)	(19.4)	(16.6)	(8.4)	(1.8)	(2.3)	2.2	(10.5)	(10.9)







SUMMARY CASH FLOW STATEMENTS

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q22
Net (Loss) Income	(8.5)	(19.4)	(16.6)	(8.4)	(1.8)	(2.3)	2.2	(10.5)	(10.9)
Share-Based Compensation	4.4	3.7	3.9	3.2	4.1	3.8	3.6	4.0	4.4
Depreciation	1.4	1.5	1.3	1.2	1.3	0.9	1.0	0.9	0.9
Other Non-Cash Effects	1.3	9.5	1.7	1.1	0.9	0.6	0.7	0.6	3.0
Changes in Assets	2.9	(1.8)	(6.2)	0.1	1.9	11.0	(9.5)	0.7	(0.7)
Changes in Liabilities	(4.7)	1.9	5.5	(3.9)	(4.8)	(7.6)	(1.3)	(5.9)	0.3
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)	(3.0)
Purchases of Property and Equipment	(0.4)	(0.7)	(2.2)	(1.9)	(0.4)	(0.6)	(O.3)	(0.1)	(0.4)
Patent Application Costs	(O.1)	0.0	(O.1)	(0.2)	(0.0)	(O.1)	(0.0)	(O.1)	(0.0)
Other	0.0	0.0	0.0	0.0	0.0	0.1	0.0	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)	(0.4)
Preferred Stock Net Proceeds	0.0	93.4	(0.8)	0.0	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	0.0
Repayment of Term Loan	(10.9)	(72.0)	0.0	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	0.0
Deferred Financing Cost	0.0	(2.8)	(0.3)	0.0	0.0	0.0	0.0	0.0	0.0
Stock Repurchased for Tax Withholdings on Vesting of Restricted Stock	(0.8)	(O.1)	0.0	(3.2)	(1.4)	(0.2)	0.0	(1.2)	0.0
Proceeds from Exercise of Stock Options	0.0	0.1	0.0	0.9	0.5	0.0	0.0	0.2	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)	0.2
Beginning Cash Balance	53.5	48.2	109.6	95.8	84.7	85.0	90.6	87.1	75.7
Change in Cash	(5.3)	61.4	(13.8)	(11.1)	0.3	5.6	(3.5)	(11.4)	(3.2)
Ending Cash Balance	48.2	109.6	95.8	84.7	85.0	90.6	87.1	75.7	72.5

Note: Some figures may not add to subtotals due to immaterial rounding differences.





REVENUE DETAIL

Quarter

Trailing 12 Months

Net Sales	\$ 53.6		\	\$ 60.0			\$ 67.4		\$ 66.9	\$261.1	\$259.9	\$258.6	\$257.5	\$256.3
Other ²	1.7	1.0	0.5	0.3	0.3	0.3	0.1	0.1	0.1	2.1	1.4	1.0	0.8	0.5
Section 351 ¹	6.1	8.2	8.7	8.2	8.6	0.5	0.3	0.4	0.6	33.7	26.0	17.6	9.8	1.9
Advanced Wound Care / Section 361 ¹	45.8	55.1	59.4	51.5	59.3	62.3	66.9	58.4	66.2	225.3	232.5	240.0	246.9	253.8
(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q22	2Q21	3Q21	4Q21	1Q22	2Q22

(I) Section 56 Includes Tissue + Cord sales. Section 35 Includes Micronized + Particulate sales. Advanced Wound Care/Section 38 and Section 35 Sales are Non-CAAP mentions. These two metrics allow understand the trend in sales between the two different product groups. [2] Adjusted not a fasle excludes impact of Revenuer Transport on mounts. Adjusted net sales is a non-CAAP measurement uses, Adjusted on the sales and the sales are sales and the sales are sales and an advanced by the sales are sales and an advanced by the sales are sales and an advanced by the sales are sales are sales are sales and an advanced by the sales are sal



NON-GAAP METRICS RECONCILIATION

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q22
Net Sales – Reported	53.6	64.3	68.6	60.0	68.2	63.1	67.4	58.9	66.9
Less: Revenue Transition Impact ¹	(1.7)	(1.0)	(0.5)	(0.3)	(0.3)	(0.3)	(0.1)		
Adjusted Net Sales	51.9	63.3	68.1	59.7	67.9	62.8	67.3	58.9	66.9
Gross Profit	45.4	54.0	57.8	50.3	55.4	53.0	56.6	49.0	55.1
Less: Revenue Transition Impact ¹	(1.5)	(0.9)	(0.4)	(0.2)	(0.3)	(0.3)	(O.1)		
Adjusted Gross Profit	44.0	53.1	57.4	50.1	55.1	52.7	56.6	49.0	55.1
Adjusted Gross Margin	84.7%	84.0%	84.2%	83.9%	81.3%	83.9%	84.1%	83.2%	82.3%
Adjusted EBITDA	11.7	7.8	10.8	5.0	3.1	7.0	3.6	(1.7)	(1.0)
Less: Capital Expenditures	(0.4)	(0.7)	(2.2)	(1.9)	(0.4)	(0.6)	(0.3)	(0.1)	(0.4)
Less: Patent Application Costs	(0.1)	0.0	(O.1)	(0.2)	(0.0)	(O.1)	(0.0)	(O.1)	(0.0)
Free Cash Flow	11.2	7.1	8.5	2.9	2.7	6.3	3.3	(1.9)	(1.4)



I Impact of revenue transition includes cash collected related to the remaining contracts. For a discussion of the revenue transition and the defined terms, refer to Item 8, Notes to the anosolidated Financial Statements in the MiMedx Group, Inc. Form 10-K for the years ended December 31, 2019 and 2020, and the respective Form 10-Qs for the noted quarterly periods lote: Some figures may not add to subtotals due to immaterial rounding differences.



ADJUSTED EBITDA RECONCILIATION

(\$ millions)	2Q20	3Q20	4Q20	1Q21	2Q21	3Q21	4Q21	1Q22	2Q22
Net (Loss) Income	(8.5)	(19.4)	(16.6)	(8.4)	(1.8)	(2.3)	2.2	(10.5)	(10.9)
Depreciation & Amortization	1.7	1.8	1.6	1.4	1.5	1.1	1.1	1.0	1.0
Interest Expense	2.6	1.5	1.5	1.5	1.4	1.0	1.2	1.1	1.2
Loss on Extinguishment of Debt	0.0	8.2	0.0	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	0.1
EBITDA	(4.2)	(7.9)	(14.5)	(5.5)	1.1	0.0	4.4	(8.3)	(8.6)
Investigation, Restatement & Related	11.4	12.0	20.4	7.2	(2.1)	3.2	(4.5)	2.6	3.2
Impairment of Intangible Assets	0.0	0.0	1.0	0.0	0.0	0.0	0.1	0.0	0.0
Share-Based Compensation	4.4	3.7	3.9	3.2	4.1	3.8	3.6	4.0	4.4
Adjusted EBITDA ¹	11.7	7.8	10.8	5.0	3.1	7.0	3.6	(1.7)	(1.0)

Investigation, Restatement & Related:

- Audit Committee Investigation completed in 2Q19

 Restatement activities completed in 2Q20

 Going forward, remainder is legal costs for Company matters, resolution costs for Company matters, recoveries from insurance providers, and indemnification costs under agreements with former officers and directors



(i) Adjusted EBITDA consists of GAAP net loss excluding: (i) depreciation, (ii) amortization of intangibles, (iii) interest expense, (iiv) loss on extinguishment of debt, (v) income tax provision, (vi) costs incurred in connection with Audit Committee Investigation, Restatement, and Related; (vii) Impairment of Intangible assets, and (vii) share-based compensation. Note: Some figures may not add to subtotate due to immaterial rounding differences.

