

NUVASIVE INC
Form 10-Q
July 27, 2017

UNITED STATES

SECURITIES AND EXCHANGE COMMISSION

Washington, D.C. 20549

Form 10-Q

(Mark One)

QUARTERLY REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

For the quarterly period ended June 30, 2017

OR

TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934

For the transition period from _____ to _____

Commission File Number: 000-50744

NUVASIVE, INC.

(Exact name of registrant as specified in its charter)

Delaware 33-0768598
(State or other jurisdiction of (I.R.S. Employer

incorporation or organization) Identification No.)

7475 Lusk Boulevard

San Diego, CA 92121

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(Address of principal executive offices)

(858) 909-1800

(Registrant's telephone number, including area code)

Indicate by check mark whether the registrant (1) has filed all reports required to be filed by Section 13 or 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period than the registrant was required to file such reports), and (2) has been subject to such filing requirements for the past 90 days. Yes No

Indicate by check mark whether the registrant has submitted electronically and posted on its corporate website, if any, every Interactive Data File required to be submitted and posted pursuant to Rule 405 of Regulation S-T (§232.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit and post such files). Yes No

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, a non-accelerated filer, smaller reporting company, or an emerging growth company. See the definitions of "large accelerated filer," "accelerated filer," "smaller reporting company," and "emerging growth company" in Rule 12b-2 of the Exchange Act.

Large accelerated filer

Accelerated filer

Non-accelerated filer

(Do not check if a small reporting company)

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.

Indicate by check mark whether the registrant is a shell company (as defined in Rule 12b-2 of the Exchange Act). Yes No

As of July 26, 2017 there were 50,803,820 shares of the registrant's common stock (par value \$0.001 per share) outstanding.

NuVasive, Inc.

Quarterly Report on Form 10-Q

June 30, 2017

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PART I. FINANCIAL INFORMATION

Item 1. Financial Statements

NUVASIVE, INC.

CONSOLIDATED BALANCE SHEETS

(in thousands, except par values and share amounts)

	June 30, 2017	December 31, 2016
ASSETS	(Unaudited)	
Current assets:		
Cash and cash equivalents	\$ 130,932	\$ 153,643
Restricted cash and investments	2,402	—
Accounts receivable, net of allowances of \$9,399 and \$8,912, respectively	190,169	171,595
Inventory, net	236,839	208,249
Prepaid income taxes	19,576	31,926
Prepaid expenses and other current assets	12,310	10,030
Total current assets	592,228	575,443
Property and equipment, net	214,601	181,524
Intangible assets, net	268,466	291,143
Goodwill	486,439	485,685
Deferred tax assets	5,961	5,810
Restricted cash and investments	4,945	7,405
Other assets	33,744	23,794
Total assets	\$ 1,606,384	\$ 1,570,804
LIABILITIES AND EQUITY		
Current liabilities:		
Accounts payable and accrued liabilities	\$ 82,933	\$ 77,585
Contingent consideration liabilities	19,271	49,742
Accrued payroll and related expenses	49,323	51,000
Income tax liabilities	11,995	2,469
Short-term borrowings	20,000	—
Senior convertible notes	63,302	61,701
Total current liabilities	246,824	242,497
Long-term senior convertible notes	573,532	564,412
Deferred and income tax liabilities, non-current	16,110	18,607
Other long-term liabilities	46,312	44,764
Commitments and contingencies		
Stockholders' equity:		
Preferred stock, \$0.001 par value; 5,000,000 shares authorized, none outstanding	—	—
Common stock, \$0.001 par value; 120,000,000 shares authorized at June 30, 2017 and December 31, 2016, 58,081,702 and 55,184,660 issued and outstanding at June 30, 2017	58	55

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and December 31, 2016, respectively

Additional paid-in capital	1,033,546	1,010,238
Accumulated other comprehensive loss	(8,131)	(10,631)
Accumulated deficit	(53,077)	(66,859)
Treasury stock at cost; 4,974,534 shares and 4,758,828 shares at June 30, 2017 and December 31, 2016, respectively	(253,503)	(237,867)
Total NuVasive, Inc. stockholders' equity	718,893	694,936
Non-controlling interest	4,713	5,588
Total equity	723,606	700,524
Total liabilities and equity	\$ 1,606,384	\$ 1,570,804

See accompanying Notes to Unaudited Consolidated Financial Statements.

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NUVASIVE, INC.

CONSOLIDATED STATEMENTS OF OPERATIONS

(in thousands, except per share amounts)

(unaudited)	Three Months Ended		Six Months Ended	
	June 30,	2016	June 30,	2016
Revenue	\$260,573	\$236,210	\$510,437	\$451,314
Cost of goods sold (excluding below amortization of intangible assets)	66,421	59,745	128,034	113,971
Gross profit	194,152	176,465	382,403	337,343
Operating expenses:				
Sales, marketing and administrative	139,109	134,487	279,611	259,325
Research and development	12,572	11,871	24,986	22,500
Amortization of intangible assets	11,349	10,603	23,410	18,474
Litigation liability (gain)	—	(43,310)	—	(43,310)
Business transition costs	1,369	2,756	1,424	8,063
Total operating expenses	164,399	116,407	329,431	265,052
Interest and other expense, net:				
Interest income	139	406	276	734
Interest expense	(10,083)	(10,537)	(19,882)	(19,009)
Loss on repurchases of convertible notes	—	—	—	(17,444)
Other expense, net	(501)	(246)	(243)	(196)
Total interest and other expense, net	(10,445)	(10,377)	(19,849)	(35,915)
Income before income taxes	19,308	49,681	33,123	36,376
Income tax expense	(7,079)	(19,891)	(8,569)	(10,411)
Consolidated net income	\$12,229	\$29,790	\$24,554	\$25,965
Add back net loss attributable to non-controlling interest	\$(432)	\$(423)	\$(875)	\$(880)
Net income attributable to NuVasive, Inc.	\$12,661	\$30,213	\$25,429	\$26,845
Net income per share attributable to NuVasive, Inc.:				
Basic	\$0.25	\$0.60	\$0.50	\$0.54
Diluted	\$0.22	\$0.57	\$0.44	\$0.51
Weighted average shares outstanding:				
Basic	51,082	50,027	50,825	49,822
Diluted	58,330	53,159	58,059	52,354

See accompanying Notes to Unaudited Consolidated Financial Statements.

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NUVASIVE, INC.

CONSOLIDATED STATEMENTS OF COMPREHENSIVE INCOME

(in thousands)

(unaudited)	Three Months		Six Months Ended	
	Ended June 30,		June 30,	
	2017	2016	2017	2016
Consolidated net income	\$12,229	\$29,790	\$24,554	\$25,965
Other comprehensive income:				
Unrealized gain (loss) on marketable securities, net of tax	1	(6)	(1)	342
Translation adjustments, net of tax	642	2,734	2,501	5,419
Other comprehensive income	643	2,728	2,500	5,761
Total consolidated comprehensive income	12,872	32,518	27,054	31,726
Net loss attributable to non-controlling interest	(432)	(423)	(875)	(880)
Comprehensive income attributable to NuVasive, Inc.	\$13,304	\$32,941	\$27,929	\$32,606

See accompanying Notes to Unaudited Consolidated Financial Statements.

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NUVASIVE, INC.

CONSOLIDATED STATEMENTS OF CASH FLOWS

(in thousands)

(unaudited)	Six Months Ended June 30,	
	2017	2016
Operating activities:		
Consolidated net income	\$24,554	\$25,965
Adjustments to reconcile net income to net cash provided by operating activities:		
Depreciation and amortization	58,688	46,329
Loss on repurchases of convertible notes	—	17,444
Amortization of non-cash interest	10,882	10,943
Stock-based compensation	15,411	12,357
Reserves on current assets	(95)	6,751
Other non-cash adjustments	7,380	8,387
Deferred income taxes	(2,570)	14,691
Changes in operating assets and liabilities, net of effects from acquisitions:		
Accounts receivable	(17,586)	(8,615)
Inventory	(29,012)	(12,019)
Prepaid expenses and other current assets	(2,485)	728
Contingent consideration liabilities	(11,200)	—
Accounts payable and accrued liabilities	4,987	14,384
Litigation liability	—	(43,310)
Accrued payroll and related expenses	(2,004)	(4,356)
Income taxes	10,172	10,534
Net cash provided by operating activities	67,122	100,213
Investing activities:		
Acquisition of Ellipse Technologies, net of cash acquired	—	(380,080)
Other acquisitions and investments	(14,417)	(8,079)
Purchases of intangible assets	(1,695)	(5,918)
Purchases of property and equipment	(68,690)	(52,566)
Purchases of marketable securities	—	(128,956)
Proceeds from sales of marketable securities	—	339,320
Net cash used in investing activities	(84,802)	(236,279)
Financing activities:		
Proceeds from the issuance of common stock	5,369	6,150
Purchase of treasury stock	(10,844)	(22,549)
Payment of contingent consideration	(18,800)	—
Proceeds from issuance of convertible debt, net of issuance costs	—	634,140
Proceeds from sale of warrants	—	44,850
Purchase of convertible note hedge	—	(111,150)
Repurchases of convertible notes	—	(343,835)

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Proceeds from revolving line of credit	20,000	50,000
Repayments on revolving line of credit	—	(50,000)
Other financing activities	(2,205)	(1,545)
Net cash (used in) provided by financing activities	(6,480)	206,061
Effect of exchange rate changes on cash	1,449	748
(Decrease) increase in cash and cash equivalents	(22,711)	70,743
Cash and cash equivalents at beginning of period	153,643	192,339
Cash and cash equivalents at end of period	\$ 130,932	\$ 263,082

See accompanying Notes to Unaudited Consolidated Financial Statements.

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NUVASIVE, INC.

NOTES TO UNAUDITED CONSOLIDATED FINANCIAL STATEMENTS

1. Description of Business and Basis of Presentation

Description of Business

NuVasive, Inc. (the “Company” or “NuVasive”) was incorporated in Delaware on July 21, 1997, and began commercializing its products in 2001. The Company’s principal product offering includes a minimally-disruptive surgical platform called Maximum Access Surgery, or MAS. The MAS platform combines three categories of solutions that collectively minimize soft tissue disruption during spine fusion surgery, provide maximum visualization and are designed to enable safe and reproducible outcomes for the surgeon and the patient. The platform includes the Company’s proprietary software-driven nerve detection and avoidance systems and Intraoperative Monitoring (“IOM”) services and support; MaXcess, an integrated split-blade retractor system; and a wide variety of specialized implants and biologics. In May 2015, the Company launched Integrated Global Alignment (“iGA”), in which products and computer assisted technology under the MAS platform help achieve more precise spinal alignment. The individual components of the MAS platform, and many of the Company’s products, can also be used in open or traditional spine surgery. The Company continues to focus research and development efforts to expand its MAS product platform and advance the applications of its unique technology into procedurally-integrated surgical solutions. The Company dedicates significant resources toward training spine surgeons on its unique technology and products.

The Company’s primary business model is to loan its MAS systems to surgeons and hospitals that purchase implants, biologics and disposables for use in individual procedures. In addition, for larger customers, the Company’s proprietary nerve monitoring systems, MaXcess and surgical instrument sets are placed with hospitals for an extended period at no up-front cost to them. The Company also offers a range of bone allograft in patented saline packaging, disposables and spine implants, which include its branded CoRoent products and fixation devices such as rods, plates and screws. The Company sells MAS instrument sets, MaXcess and nerve monitoring systems to hospitals, however, such sales are immaterial to the Company’s results of operations.

The Company also designs and sells expandable growing rod implant systems that can be non-invasively lengthened following implantation with precise, incremental adjustments via an external remote controller using magnetic technology called MAGnetic External Control, or MAGEC, which allows for the minimally invasive treatment of early-onset and adolescent scoliosis. This technology is also the basis for the Company’s PRECICE limb lengthening system, which allows for the correction of long bone limb length discrepancy, as well as enhanced bone healing in patients that have experienced traumatic injury.

The Company intends to continue development on a wide variety of projects intended to broaden surgical applications for greater procedural integration of its MAS techniques and additional applications of the MAGEC technology. Such applications include tumor, trauma, and deformity, as well as increased fixation options, sagittal alignment products, imaging and navigation. The Company also expects to continue expanding its other product and services offerings as it executes on its strategy to offer customers an end-to-end, integrated procedural solution for spine surgery. The Company intends to continue to pursue business and technology acquisition targets and strategic partnerships.

Basis of Presentation and Principles of Consolidation

The accompanying Unaudited Consolidated Financial Statements include the accounts of the Company and its majority-owned or controlled subsidiaries, collectively referred to as either NuVasive or the Company. The Company translates the financial statements of its foreign subsidiaries using end-of-period exchange rates for assets and liabilities and average exchange rates during each reporting period for results of operations. When there is a portion of equity in an acquired subsidiary not attributable, directly or indirectly, to the respective parent entity, the Company records the fair value of the non-controlling interest at the acquisition date and classifies the amounts attributable to non-controlling interest separately in equity in the Company's Consolidated Financial Statements. Any subsequent changes in a parent's ownership interest while the parent retains its controlling financial interest in its subsidiary are accounted for as equity transactions. All significant intercompany balances and transactions have been eliminated in consolidation.

The accompanying Unaudited Consolidated Financial Statements have been prepared pursuant to the rules and regulations of the Securities and Exchange Commission (the "SEC"). Pursuant to these rules and regulations, the Company has condensed or omitted certain information and footnote disclosures it normally includes in its annual Consolidated Financial Statements prepared in accordance with generally accepted accounting principles in the United States ("GAAP"). Operating results for the three and six months ended June 30, 2017 are not necessarily indicative of the results that may be expected for any other interim period or for the full year. These Unaudited Consolidated Financial Statements should be read in conjunction with the audited Consolidated Financial Statements and notes thereto for the year ended December 31, 2016 included in the Company's Annual Report on Form 10-K filed with the SEC. In the opinion of management, the Unaudited Consolidated Financial Statements include all adjustments that are of a normal and recurring nature that are necessary for the fair presentation of the Company's financial position and of the results of operations and cash flows for the periods presented.

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The Company has reclassified certain operating expenses into business transition costs. The reclassification had no impact on previously reported results of operations or financial position. Refer to “Recently Adopted Accounting Standards” below for information regarding historical financial information adjusted for a change in accounting policy.

Use of Estimates

To prepare financial statements in conformity with GAAP, management must make estimates and assumptions that affect the amounts reported in the financial statements and accompanying notes. Actual results could differ from those estimates.

Recent Accounting Pronouncements Not Yet Adopted

In May 2014, the Financial Accounting Standards Board (“FASB”) issued Accounting Standard Update No. 2014-09, Revenue from Contracts with Customers (“ASU 2014-09”), an updated standard on revenue recognition. ASU 2014-09 provides enhancements to the quality and consistency of how revenue is reported by companies while also improving comparability in the financial statements of companies reporting using International Financial Reporting Standards or GAAP. The main purpose of the new standard is for companies to recognize revenue to depict the transfer of goods or services to customers in amounts that reflect the consideration to which a company expects to be entitled in exchange for those goods or services. The new standard also will result in enhanced disclosures about revenue, provide guidance for transactions that were not previously addressed comprehensively and improve guidance for multiple-element arrangements. In August 2015, the FASB issued ASU No. 2015-14, Revenue from Contracts with Customers: Deferral of the Effective Date, which deferred the effective date of the new revenue standard for periods beginning after December 15, 2016 to December 15, 2017, with early adoption permitted but not earlier than the original effective date. Accordingly, the updated standard is effective for the Company in the first quarter of fiscal 2018. The Company performed a preliminary assessment of the impact of ASU 2014-09 on the Consolidated Financial Statements, and considered all items outlined in the standard. In assessing the impact, the Company has outlined all revenue generating activities, mapped those activities to deliverables and traced those deliverables to the standard. The Company is now assessing what impact the change in standard will have on those deliverables. The Company will continue to evaluate the future impact and method of adoption of ASU 2014-09 and related amendments on the Consolidated Financial Statements and related disclosures throughout 2017. The Company believes the adoption will modify the way the Company analyzes contracts. The Company will adopt the new standard beginning January 2018.

In January 2016, the FASB issued Accounting Standards Update No. 2016-01, Financial Instruments-Overall: Recognition and Measurement of Financial Assets and Financial Liabilities (“ASU 2016-01”), which requires that (i) all equity investments, other than equity-method investments, in unconsolidated entities generally be measured at fair value through earnings and (ii) when the fair value option has been elected for financial liabilities, changes in fair value due to instrument-specific credit risk will be recognized separately in other comprehensive income. Additionally, the ASU 2016-01 changes the disclosure requirements for financial instruments. The new standard will be effective for the Company starting in the first quarter of fiscal 2019. Early adoption is permitted for certain provisions. The Company is in the process of determining the effects the adoption will have on its Consolidated Financial Statements as well as whether to early adopt certain provisions.

In February 2016, the FASB issued Accounting Standards Update No. 2016-02, Leases, which outlines a comprehensive lease accounting model and supersedes the current lease guidance. The new accounting standard

requires lessees to recognize lease liabilities and corresponding right-of-use assets for all leases with lease terms of greater than twelve months. It also changes the definition of a lease and expands the disclosure requirements of lease arrangements. The new accounting standard must be adopted using the modified retrospective approach and will be effective for the Company starting in the first quarter of fiscal 2019. Early adoption is permitted. The Company believes the adoption will modify its analyses and disclosures of lease agreements considering operating leases are a significant portion of the Company's total lease commitments. The Company is in the process of determining the effects the adoption will have on its Consolidated Financial Statements as well as whether to early adopt the new guidance.

In June 2016, the FASB issued Accounting Standards Update No. 2016-13, Financial Instruments – Credit Losses, which changes the accounting for recognizing impairments of financial assets. Under the new guidance, credit losses for certain types of financial instruments will be estimated based on expected losses. The new guidance also modifies the impairment models for available-for-sale debt securities and for purchased financial assets with credit deterioration since their origination. The new guidance will be effective for the Company starting in the first quarter of fiscal 2021. Early adoption is permitted starting in the first quarter of fiscal 2020. The Company believes the adoption will modify the way the Company analyzes financial instruments, but it does not anticipate a material impact on results of operations. The Company is in the process of determining the effects the adoption will have on its Consolidated Financial Statements as well as whether to early adopt the new guidance.

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In August 2016, the FASB issued Accounting Standards Update No. 2016-15, Classification of Certain Cash Receipts and Cash Payments (“ASU 2016-15”), which eliminates the diversity in practice related to the classification of certain cash receipts and payments for debt prepayment or extinguishment costs, the maturing of a zero coupon bond, the settlement of contingent liabilities arising from a business combination, proceeds from insurance settlements, distributions from certain equity method investees and beneficial interests obtained in a financial asset securitization. ASU 2016-15 designates the appropriate cash flow classification, including requirements to allocate certain components of these cash receipts and payments among operating, investing and financing activities. The retrospective transition method, requiring adjustment to all comparative periods presented, is required unless it is impracticable for some of the amendments, in which case those amendments would be prospectively as of the earliest date practicable. This update is effective for annual periods beginning after December 15, 2017, and interim periods within those fiscal years, with early adoption permitted, including adoption in an interim period. The Company does not expect the adoption to have any significant impact on its Consolidated Financial Statements.

In November 2016, the FASB issued Accounting Standards Update No. 2016-18, Restricted Cash, which requires entities to show the changes in the total of cash, cash equivalents, restricted cash and restricted cash equivalents in the statement of cash flows. As a result, entities will no longer present transfers between cash and cash equivalents and restricted cash and restricted cash equivalents in the statement of cash flows. The amendments in this update should be applied using a retrospective transition method to each period presented. This update is effective for annual periods beginning after December 15, 2017, and interim periods within those fiscal years with early adoption permitted, including adoption in an interim period. The Company does not expect the adoption to have any significant impact on its Consolidated Financial Statements.

In January 2017, the FASB issued Accounting Standards Update No. 2017-01, Clarifying the Definition of a Business, which clarifies and provides a more robust framework to use in determining when a set of assets and activities is a business. The amendments in this update should be applied prospectively on or after the effective date. This update is effective for annual periods beginning after December 15, 2017, and interim periods within those periods. Early adoption is permitted for acquisition or deconsolidation transactions occurring before the issuance date or effective date and only when the transactions have not been reported in issued or made available for issuance financial statements. The Company is in the process of determining the effects the adoption will have on its Consolidated Financial Statements as well as whether to early adopt the new guidance.

In January 2017, the FASB issued Accounting Standards Update No. 2017-04, Intangibles – Goodwill and Other, which eliminates the requirement to calculate the implied fair value of goodwill to measure a goodwill impairment charge. Instead, entities will record an impairment charge based on the excess of a reporting unit’s carrying amount over its fair value. The standard has tiered effective dates, starting in 2020 for calendar-year public business entities that meet the definition of an SEC filer. Early adoption is permitted for annual and interim goodwill impairment testing dates after January 1, 2017. The Company is in the process of determining the effects the adoption will have on its Consolidated Financial Statements as well as whether to early adopt the new guidance.

In February 2017, the FASB issued Accounting Standards Update No. 2017-05, Other Income – Gains and Losses from the Derecognition of Nonfinancial Assets, which clarifies the scope of asset derecognition and adds guidance for partial sales and nonfinancial assets. An entity is required to apply the amendments in this update at the same time that it applies the amendments in ASU 2014-09. For public entities, this update is effective for annual periods beginning after December 15, 2017, and interim periods within those periods. Public entities may apply the guidance earlier but only as of annual reporting periods beginning after December 15, 2016, including interim reporting periods within that

reporting period. The Company will adopt the new standard beginning January 2018.

In May 2017, the FASB issued Accounting Standards Update No. 2017-09, Compensation – Stock Compensation, which clarifies when changes to the terms or conditions of a share-based payment award must be accounted for as a modification. Entities will apply the modification accounting guidance if the value, vesting conditions or classification of the award changes. This update is effective for annual periods beginning after December 15, 2017, and interim periods within those periods. The Company does not expect the adoption to have any significant impact on its Consolidated Financial Statements.

In July 2017, the FASB issued Accounting Standards Update No. 2017-19, Earnings Per Share, Distinguishing Liabilities from Equity, Derivatives and Hedging, which changes the accounting and earnings per share for certain instruments with round down features. The amendments in this update should be applied using a cumulative-effect adjustment as of the beginning of the fiscal year or retrospective adjustment to each period presented. This update is effective for annual periods beginning after December 15, 2018, and interim periods within those periods. The Company is in the process of determining the effects the adoption will have on its Consolidated Financial Statements.

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Recently Adopted Accounting Standards

In March 2016, the FASB issued Accounting Standards Update 2016-09, Improvements to Employee Share-Based Payment Accounting (“ASU 2016-09”), which simplifies the accounting for employee share-based payments. The new standard requires the immediate recognition of all excess tax benefits and deficiencies in the income statement, and requires classification of excess tax benefits as an operating activity as opposed to a financing activity in the statements of cash flows. The provisions of the new standard are effective for the Company beginning January 1, 2017, with early adoption permitted. The Company elected to early adopt ASU 2016-09 in the second quarter 2016, which requires any adjustments to be recorded as of the beginning of fiscal 2016. As a result, the Company recorded a modified retrospective adjustment of \$16.6 million to deferred tax assets and accumulated deficit as of January 1, 2016, and a retrospective adjustment to the previously reported first quarter 2016 provision for income taxes of approximately \$5.5 million for the recognition of excess tax benefits in the provision for income taxes rather than additional paid-in capital. This resulted in a decrease in net loss per share of \$0.11 for the three months ended March 31, 2016. The Company elected to apply the change in classification for excess tax benefits in the statement of cash flows on a prospective basis, and elected to continue estimating stock-based compensation award forfeitures in determining the amount of compensation cost to be recognized each period.

In October 2016, the FASB issued Accounting Standards Update No. 2016-16, Intra-Entity Transfers of Assets Other Than Inventory (“ASU 2016-16”), which aims to improve the accounting for the income tax consequences of intra-entity transfers of assets other than inventory. This amendment requires an entity to recognize the income tax consequences of an intra-entity transfer of an asset other than inventory when the transfer occurs. The amendments in this update should be applied on a modified retrospective basis through a cumulative-effect adjustment directly to retained earnings as of the beginning of the period of adoption. This update is effective for annual periods beginning after December 15, 2017, and interim periods within those fiscal years with early adoption permitted, including adoption in an interim period. The Company elected to early adopt ASU 2016-16 in the first quarter 2017, which requires any adjustments to be recorded as of the beginning of fiscal 2017. As a result, the Company recorded a modified retrospective adjustment of \$11.6 million to deferred tax assets and accumulated deficit as of January 1, 2017. The early adoption resulted in a decrease of \$0.9 million and \$1.5 million in income tax expense that would have amortized out of prepaid income taxes during the three and six months ended June 30, 2017, respectively, and an increase in both basic and diluted earnings per share of \$0.02 and \$0.03 for the three and six months ended June 30, 2017.

In January 2017, the FASB issued Accounting Standards Update No. 2017-03, Accounting Changes and Error Corrections and Investments – Equity Method and Joint Ventures (“ASU 2017-03”), which will require registrants to disclose the effect that recently issued accounting standards will have on their financial statements when adopted in a future period. This update is effective immediately. The Company is in the process of determining the effects of recently issued accounting standards on its Consolidated Financial Statements. The Company will revise its disclosures for the standards not yet adopted as required by ASU 2017-03 as the Company progresses through its impact assessments.

Revenue Recognition

In accordance with the SEC guidance, the Company recognizes revenue when all four of the following criteria are met: (i) persuasive evidence that an arrangement exists; (ii) delivery of the products and/or services has occurred; (iii) the selling price is fixed or determinable; and (iv) collectability is reasonably assured. Specifically, revenue from

the sale of implants, biologics and disposables is generally recognized upon a purchase order from the hospital or acknowledgment from the hospital indicating product use or implantation, or upon shipment to third-party customers who immediately accept title. Revenue from IOM services is recognized in the period the service is performed for the amount of payment expected to be received. Revenue from the sale of instrument sets and nerve monitoring systems is recognized upon receipt of a purchase order and the subsequent shipment to customers who immediately accept title.

Comprehensive Income

Comprehensive income is defined as the change in equity during a period from transactions and other events and circumstances from non-owner sources. Comprehensive income includes unrealized gains or losses, net of tax, on the Company's marketable securities and foreign currency translation adjustments. The cumulative translation adjustments included in accumulated other comprehensive income were \$8.1 million and \$10.6 million at June 30, 2017 and December 31, 2016, respectively.

Product Shipment Costs

Product shipment costs, included in sales, marketing and administrative expense in the accompanying Consolidated Statements of Operations, were \$5.7 million and \$11.6 million for the three and six months ended June 30, 2017, respectively, and \$6.6 million and \$12.8 million for the three and six months ended June 30, 2016, respectively. The majority of the Company's shipping costs are related to the loaning of instrument sets, which are not typically sold as part of the Company's core sales offering. Amounts billed to customers for shipping and handling of products are reflected in revenues and are not significant for any period presented.

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Business Transition Costs

The Company incurs certain costs related to acquisition, integration and business transition activities which include severance, relocation, consulting, leasehold exit costs, third party merger and acquisition costs and other costs directly associated with such activities. The Company incurred \$1.4 million of business transition costs during the three and six months ended June 30, 2017, respectively, and \$2.8 million and \$8.1 million during the three and six months ended June 30, 2016, respectively, primarily related to acquisition and integration activities.

Litigation Liability Gain

During the three and six months ended June 30, 2016, the Company agreed to settle its ongoing litigation with Warsaw Orthopedic, Inc., Medtronic Sofamor Danek USA, Inc. and other Medtronic related entities (collectively, "Medtronic"). As a result of the settlement, the Company agreed to pay \$45.0 million to Medtronic and accordingly recorded a gain of \$43.3 million related to the settlement by reducing its previous accrual of \$88.3 million related to the matter.

See Note 11 to the Unaudited Consolidated Financial Statements for further discussion.

2. Net Income Per Share

The following table sets forth the computation of basic and diluted net income per share attributable to the Company:

(in thousands, except per share data)	Three Months Ended June 30,		Six Months Ended June 30,	
	2017	2016	2017	2016
Numerator:				
Net income attributable to NuVasive, Inc.	\$12,661	\$30,213	\$25,429	\$26,845
Denominator for basic and diluted net income per share:				
Weighted average common shares outstanding for basic	51,082	50,027	50,825	49,822
Dilutive potential common stock outstanding:				
Stock options and employee stock purchase plan	138	374	180	409
Restricted stock units	1,356	1,281	1,386	1,090
Warrants	2,929	819	2,987	409
Senior Convertible Notes	2,825	658	2,681	624
Weighted average common shares outstanding for diluted	58,330	53,159	58,059	52,354
Basic net income per share attributable to NuVasive, Inc.	\$0.25	\$0.60	\$0.50	\$0.54
Diluted net income per share attributable to NuVasive, Inc.	\$0.22	\$0.57	\$0.44	\$0.51

The following weighted-average outstanding common stock equivalents were not included in the calculation of net income per diluted share because their effects were anti-dilutive:

(in thousands)	Three Months Ended June 30,		Six Months Ended June 30,	
	2017	2016	2017	2016
	41	3	71	1,817

Stock options, employee stock purchase plan, and restricted stock units				
Warrants	10,865	10,865	10,865	15,642
Senior Convertible Notes	—	10,865	—	15,100
Total	10,906	21,733	10,936	32,559

As discussed in Note 1 to the Unaudited Consolidated Financial Statements, the Company elected to early adopt ASU 2016-09 in the second quarter 2016, which requires any adjustments to be recorded as of the beginning of the fiscal year. The retrospective adjustments to the Company's financial results for the three months ended March 31, 2016 included a decrease in net loss attributable to the Company of \$5.5 million, which resulted in a decrease in net loss per share of \$0.11. The financial information in the table above for the six months ended June 30, 2016 reflects this retrospective adjustment to the Company's financial results for the three months ended March 31, 2016.

3. Financial Instruments and Fair Value Measurements

As of June 30, 2017, the Company held investments in securities classified as cash equivalents. During the periods presented, the Company did not hold any investments that were in a significant unrealized loss position and no impairment charges were recorded. Realized gains and losses and interest income related to marketable securities were immaterial during all periods presented.

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Foreign Currency and Derivative Financial Instruments

The Company translates the financial statements of its foreign subsidiaries using end-of-period exchange rates for assets and liabilities and average exchange rates during each reporting period for results of operations.

Some of the Company's reporting entities conduct a portion of their business in currencies other than the entity's functional currency. These transactions give rise to receivables and payables that are denominated in currencies other than the entity's functional currency. The value of these receivables and payables is subject to changes in currency exchange rates from the point at which the transactions are originated until the settlement in cash. Both realized and unrealized gains and losses in the value of these receivables and payables are included in the determination of net income. Net currency exchange gains (losses), which include gains and losses from derivative instruments, were \$(0.5) million and \$(0.3) million for the three and six months ended June 30, 2017, respectively, and \$(0.3) and \$(0.2) million for the three and six months ended June 30, 2016, respectively, and are included in other expense, net in the Consolidated Statements of Operations.

To manage foreign currency exposure risks, the Company uses derivatives for activities in entities that have short-term intercompany receivables and payables denominated in a currency other than the entity's functional currency. The fair value is based on a quoted market price (Level 1). As of June 30, 2017 and December 31, 2016 a notional principal amount of \$16.1 million and \$15.1 million, respectively, in foreign currency forward contracts was outstanding to hedge currency risk relative to the Company's foreign receivables and payables. Derivative instrument net losses on the Company's forward exchange contracts were \$0.9 million and \$1.3 million for the three and six months ended June 30, 2017, respectively, and was immaterial and \$0.2 million for the three and six months ended June 30, 2016, respectively, and are included in other expense, net in the Consolidated Statements of Operations. The fair value of the forward contract exchange derivative instrument asset (liability) was de minimis as of June 30, 2017 and \$(0.2) million as of December 31, 2016. The derivative instruments are recorded in other current assets or other current liabilities in the Consolidated Balance Sheets commensurate with the nature of the instrument at period end.

Fair Value Measurements

The Company measures certain assets and liabilities in accordance with authoritative guidance which requires fair value measurements be classified and disclosed in one of the following three categories:

Level 1: Quoted prices (unadjusted) in active markets that are accessible at the measurement date for assets or liabilities.

Level 2: Observable prices that are based on inputs not quoted on active markets, but corroborated by market data.

Level 3: Unobservable inputs are used when little or no market data is available.

Assets and liabilities are classified based on the lowest level of input that is significant to the fair value measurements. The Company reviews the fair value hierarchy classification on a quarterly basis. Changes in the ability to observe valuation inputs may result in a reclassification of levels for certain assets or liabilities within the fair value hierarchy. The Company did not have any transfers of assets and liabilities between the levels of the fair value measurement hierarchy during the three months ended June 30, 2017.

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The fair values of the Company's assets and liabilities, including cash equivalents, marketable securities, and restricted investments are measured at fair value on a recurring basis, and are determined under the fair value categories as follows:

(in thousands)	Total	Quoted Price in Active Market (Level 1)	Significant Other Observable Inputs (Level 2)	Significant Unobservable Inputs (Level 3)
June 30, 2017:				
Cash equivalents:				
Money market funds	\$17,000	\$ 17,000	\$ —	\$ —
Total cash equivalents	\$17,000	\$ 17,000	\$ —	\$ —
December 31, 2016:				
Cash equivalents:				
Money market funds	\$72,866	\$ 72,866	\$ —	\$ —
Corporate notes	4,551	—	4,551	—
Commercial paper	21,471	—	21,471	—
Securities of government-sponsored entities	5,995	—	5,995	—
Total cash equivalents	\$104,883	\$ 72,866	\$ 32,017	\$ —

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The fair value of certain financial instruments was measured and classified within Level 1 of the fair value hierarchy based on quoted prices. Certain financial instruments classified within Level 2 of the fair value hierarchy include the types of instruments that trade in markets that are not considered to be active, but are valued based on quoted market prices, broker or dealer quotations, or alternative pricing sources with reasonable levels of price transparency.

The carrying amounts of certain financial instruments such as cash equivalents, accounts receivable, prepaid expenses, other current assets, accounts payable, accrued expenses, and other current liabilities as of June 30, 2017 and December 31, 2016 approximate their related fair values due to the short-term maturities of these instruments.

The fair value, based on a quoted market price (Level 1), of the Company's outstanding Senior Convertible Notes due 2017 at June 30, 2017 and December 31, 2016 were approximately \$116.2 million and \$102.7 million, respectively. The fair value, based on a quoted market price (Level 1), of the Company's outstanding Senior Convertible Notes due 2021 at June 30, 2017 and December 31, 2016 was \$901.8 million and \$827.6 million, respectively. See Note 13 to the Unaudited Consolidated Financial Statements for further discussion on the carrying value and settlement of the Company's Senior Convertible Notes due 2017.

Contingent Consideration Liabilities

The fair value of contingent consideration liabilities assumed in business combinations is recorded as part of the purchase price consideration of the acquisition, and is determined using a discounted cash flow model or probability simulation model. The significant inputs of such models are not observable in the market, such as certain financial metric growth rates, volatility rates, projections associated with the applicable milestone, the interest rate, and the related probabilities and payment structure in the contingent consideration arrangement. Fair value adjustments to contingent consideration liabilities are recorded through operating expenses in the Consolidated Statement of Operations. Contingent consideration arrangements assumed by an asset purchase will be measured and accrued when such contingency is resolved.

Contingent consideration liabilities were \$37.4 million and \$67.5 million as of June 30, 2017 and December 31, 2016, respectively, and were recorded in the Consolidated Balance Sheet commensurate with the respective payment terms. In April 2017, the Company paid the \$30.0 million outstanding milestone obligation associated with the Ellipse Technologies acquisition. In accordance with the guidance outlined in ASU 2016-15, \$18.8 million of the \$30.0 million represented the initial purchase price allocation and is presented as a cash outflow for financing activities on the Consolidated Statement of Cash Flows and the remaining \$11.2 million related to increased fair value adjustments and is presented in operating activities. See Note 5 to the Unaudited Consolidated Financial Statements for further discussion on contingent consideration liabilities assumed in business combinations.

The following table sets forth the changes in the estimated fair value of the Company's liabilities measured on a recurring basis using significant unobservable inputs (Level 3):

(in thousands)	Six Months Ended	
	June 30,	
	2017	2016
Fair value measurement at beginning of period	\$67,501	\$—
Contingent consideration liability recorded upon acquisition	533	21,439
Change in fair value measurement	(657)	339

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Changes resulting from foreign currency fluctuations	44	32
Contingent consideration paid or settled	(30,000)	—
Fair value measurement at end of period	\$37,421	\$21,810

Non-financial assets and liabilities measured on a nonrecurring basis

Certain non-financial assets and liabilities are measured at fair value, usually with Level 3 inputs including the discounted cash flow method or cost method, on a nonrecurring basis in accordance with authoritative guidance. These include items such as non-financial assets and liabilities initially measured at fair value in a business combination and non-financial long-lived assets measured at fair value for an impairment assessment. In general, non-financial assets, including goodwill, intangible assets and property and equipment, are measured at fair value when there is an indication of impairment and are recorded at fair value only when any impairment is recognized. The carrying values of the Company's capital lease obligations approximated their estimated fair value as of June 30, 2017 and December 31, 2016.

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4. Goodwill and Intangible Assets

Goodwill and intangible assets consisted of the following:

(in thousands, except years)	Weighted- Average Amortization Period	Gross	Accumulated	Intangible
	(in years)	Amount	Amortization	Assets, net
June 30, 2017:				
Intangible assets subject to amortization:				
Developed technology	8	\$247,148	\$ (82,352)	\$ 164,796
Manufacturing know-how and trade secrets	12	21,139	(14,432)	6,707
Trade name and trademarks	9	25,200	(9,022)	16,178
Customer relationships	9	118,275	(37,490)	80,785
Total intangible assets subject to amortization	9	\$411,762	\$ (143,296)	\$ 268,466
Intangible assets not subject to amortization:				
Goodwill				\$ 486,439
Total goodwill and intangible assets, net				\$ 754,905
December 31, 2016:				
Intangible assets subject to amortization:				
Developed technology	8	\$247,148	\$ (66,833)	\$ 180,315
Manufacturing know-how and trade secrets	13	20,572	(13,604)	6,968
Trade name and trademarks	9	25,200	(7,478)	17,722
Customer relationships	9	117,018	(30,880)	86,138
Total intangible assets subject to amortization	9	\$409,938	\$ (118,795)	\$ 291,143
Intangible assets not subject to amortization:				
Goodwill				\$ 485,685
Total goodwill and intangible assets, net				\$ 776,828

The following table summarizes the changes in the carrying value of the Company's goodwill:

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(in thousands)

December 31, 2016	
Gross goodwill	\$493,985
Accumulated impairment loss	(8,300)
	485,685
Changes to gross goodwill	
Increases recorded in business combinations	374
Changes in purchase price allocation	386
Changes resulting from foreign currency fluctuations	(6)
	754
June 30, 2017	
Gross goodwill	494,739
Accumulated impairment loss	(8,300)
	\$486,439

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Total expense related to the amortization of intangible assets, which is recorded in both cost of goods sold and operating expenses in the Consolidated Statements of Operations depending on the functional nature of the intangible asset, was \$12.2 million and \$11.3 million for the three months ended June 30, 2017 and June 30, 2016, respectively, and \$25.2 million and \$20.1 million for the six months ended June 30, 2017 and June 30, 2016, respectively.

Total future amortization expense related to intangible assets subject to amortization at June 30, 2017 is set forth in the table below:

(in thousands)	
Remaining 2017	\$24,489
2018	46,917
2019	45,235
2020	44,780
2021	42,861
Thereafter through 2026	64,184
Total future amortization expense	\$268,466

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5. Business Combinations

The Company recognizes the assets acquired, liabilities assumed, and any non-controlling interest at fair value at the date of acquisition. Certain acquisitions contained contingent consideration arrangements that required the Company to assess the acquisition date fair value of the contingent consideration liabilities, which was recorded as part of the purchase price allocation of the acquisition, with subsequent fair value adjustments to the contingent consideration recorded in the Consolidated Statements of Operations. See Note 3 to the Unaudited Consolidated Financial Statements for further discussion on contingent consideration liabilities.

Acquisition of Ellipse Technologies, Inc.

On February 11, 2016, the Company acquired all of the stock interest in Ellipse Technologies, Inc., which now operates as a wholly owned subsidiary of the Company under the renamed legal entity NuVasive Specialized Orthopedics, Inc. (“NSO”), for a purchase price of \$380.0 million (including holdbacks for retained employment of Ellipse Technologies leadership that is to be expensed and is not considered part of the final purchase price) and a milestone payment of \$30.0 million payable in cash in 2017 related to the achievement of a specific revenue target. A cash payment of \$382.2 million, which included additional amounts for cash on hand and traditional working capital adjustments, was transferred at the closing. Subsequent to the closing payment, the Company received \$0.6 million from the escrow for traditional working capital adjustments finalized after the closing.

NSO designs and sells expandable growing rod implant systems that can be non-invasively lengthened following implantation with precise, incremental adjustments via an external remote controller using magnetic technology called MAGnetic External Control, or MAGEC. The technology platform provides the basis of NSO’s core product offerings, including MAGEC-EOS, which allows for the minimally invasive treatment of early-onset and adolescent scoliosis, as well as the PRECICE limb lengthening system, which allows for the correction of long bone limb length discrepancy, as well as enhanced bone healing in patients that have experienced traumatic injury.

The Company applied certain assumptions and findings in the valuation outcome for the assets acquired and liabilities assumed, for which the allocation of the purchase price is based on their fair values, as follows:

(in thousands)	
Cash paid for purchase	\$381,579
Accounts receivable	7,148
Inventory	22,451
Other current assets	1,855
Property, plant and equipment, net	6,725
Definite-lived intangible assets:	
Developed technology	133,900
Customer relationships	33,200
Trade names	16,200
Goodwill	241,905
Deferred tax assets	18,471
Other assets	1,868
Contingent consideration liability	18,800

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Deferred tax liabilities	75,160
Other liabilities assumed	8,184
	\$381,579

Goodwill recognized in this transaction is not deductible for income tax purposes. Goodwill largely consists of expected revenue synergies resulting from the combination of product portfolios, cost synergies related to elimination of redundant facilities, functions and staffing; use of the Company's existing commercial infrastructure to expand sales of NSO's products; and the assembled workforce. The intangible assets acquired will be amortized on a straight-line basis over weighted-average useful lives of seven years, nine years and seven years for technology-based, customer-related intangible assets, and trade name related intangible assets, respectively. The estimated fair values of the intangible assets acquired were primarily determined using the income approach based on significant inputs that were not observable market data.

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In connection with the acquisition, a contingent liability of \$18.8 million was recorded as of the acquisition date for the potential revenue-based milestone payment. The liability was fair valued using the Monte Carlo simulation based on specific revenue achievement scenarios and discount factors. Changes in fair value of the liability over the measurement period were recorded in the results of operations in the Consolidated Statements of Operations. The revenue-based milestone was achieved as of December 31, 2016, and the Company adjusted the milestone liability to \$30.0 million, which represented the full amount of the milestone obligation under the merger agreement. The Company paid the milestone in April 2017, and no additional consideration is owed related to the acquisition.

Acquisition costs of \$4.0 million were recognized in business transition costs as incurred. The Company's results of operations included the operating results of NSO, since the date of acquisition, of \$15.1 million and \$21.0 million of revenue for the three and six months ended June 30, 2016, respectively, and net income (loss) of \$1.1 million and \$(0.7) million for the three and six months ended June 30, 2016, respectively, in the Unaudited Consolidated Statement of Operations.

The following table presents the unaudited pro forma results for the three and six months ended June 30, 2017 and June 30, 2016. The unaudited pro forma financial information combines the results of operations of NuVasive and Ellipse Technologies as though the companies had been combined as of January 1, 2015 and therefore many of the non-recurring business combination adjustments would have been included in the year ended December 31, 2015 by nature of such adjustments instead of the periods presented. The pro forma information is presented for informational purposes only and is not indicative of the results of operations that would have been achieved if the acquisition had taken place at such times. The comparable periods for the three and six months ended June 30, 2016, include adjustments directly attributable to the business combination, including adjustments for increased fair value of acquired inventory of \$(7.4) million and \$(12.3) million, respectively, immaterial adjustments to revenue for deferred revenue adjustments, and related tax effects. The six month period ended June 30, 2016 also includes an adjustment of \$4.0 million for acquisition related expenses. The pre-acquisition accounting policies of Ellipse Technologies were materially similar to the Company, with the differences adjusted to reflect the accounting policies of the Company in the unaudited pro forma results presented.

(in thousands, except per share amounts)	Three Months Ended		Six Months Ended	
	June 30, 2017	2016	June 30, 2017	2016
	(unaudited)	(unaudited)	(unaudited)	(unaudited)
Revenues	\$260,573	\$ 236,270	\$510,437	\$ 457,282
Net income (loss) attributable to NuVasive, Inc.	12,661	35,966	25,429	28,438
Net income per share attributable to NuVasive, Inc.:				
Basic	\$0.25	\$ 0.72	\$0.50	\$ 0.57
Diluted	\$0.22	\$ 0.68	\$0.44	\$ 0.54

Other Acquisitions

The Company has completed other acquisitions that were not considered material to the overall Unaudited Consolidated Financial Statements during the periods presented. These acquisitions have been included in the Unaudited Consolidated Financial Statements from the respective dates of acquisition. The Company does not believe that collectively the acquisitions made during the periods presented, excluding NSO, are material to the overall

financial statements.

For certain acquisitions completed during the periods presented, excluding NSO, the Company is still in the process of finalizing the purchase price allocation given the timing of the acquisition and the size and scope of the assets and liabilities subject to valuation. While the Company does not expect material changes in the valuation outcome, certain assumptions and findings that were in place at the date of acquisition could result in changes in the purchase price allocation.

Variable Interest Entities

Progentix Orthobiology B.V.

In 2009, the Company completed the purchase of forty percent (40%) of the capital stock of Progentix Orthobiology B.V. (“Progentix”), a company organized under the laws of the Netherlands, from existing shareholders pursuant to a Preferred Stock Purchase Agreement for \$10.0 million in cash (the “Initial Investment”). As of June 30, 2017, the Company has loaned Progentix cumulatively \$5.3 million at an interest rate of 6% per year. The Company is not obligated to provide additional funding. Concurrently, with the Initial Investment, the Company and Progentix entered into a Distribution Agreement (as amended, the “Distribution Agreement”), whereby Progentix appointed the Company as its exclusive distributor for certain Progentix products. The Distribution Agreement is in effect for a term of ten years unless terminated earlier in accordance with its terms.

In accordance with authoritative guidance, the Company has determined that Progentix is a variable interest entity (“VIE”), as it does not have the ability to finance its activities without additional subordinated financial support and its equity investors will not absorb their proportionate share of expected losses and will be limited in the receipt of the potential residual returns of Progentix.

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Total assets and liabilities of Progentix included in the accompanying Consolidated Balance Sheets are as follows:

(in thousands)	June 30, 2017	December 31, 2016
Total current assets	\$781	\$ 334
Identifiable intangible assets, net	9,826	10,900
Goodwill	12,654	12,654
Accounts payable and accrued expenses	496	551
Deferred tax liabilities, net	689	880
Non-controlling interest	4,713	5,588

The following is a reconciliation of equity (net assets) attributable to the non-controlling interest:

(in thousands)	Six Months Ended June 30,	
	2017	2016
Non-controlling interest at beginning of period	\$5,588	\$7,309
Less: Net loss attributable to the non-controlling interest	(875)	(880)
Non-controlling interest at end of period	\$4,713	\$6,429

NuVasive Clinical Services and Physician Practices

The Company's NuVasive Clinical Services division, which provides IOM services to surgeons and healthcare facilities across the U.S., maintains contractual relationships with several physician practices ("PCs") which were inherited through the 2011 acquisition of Impulse Monitoring, Inc. and the 2016 acquisition of BNN Holdings Corp. In accordance with authoritative guidance, the Company has determined that the PCs are VIEs and therefore, the accompanying Unaudited Consolidated Financial Statements include the accounts of the PCs from the date of acquisition. During the periods presented, the results of the PCs were immaterial to the Company's financials. The creditors of the PCs have claims only on the assets of the PCs, which are not material, and the assets of the PCs are not available to the Company.

6. Indebtedness

The carrying values of the Company's Senior Convertible Notes are as follows:

(in thousands)	June 30, 2017	December 31, 2016
2.75% Senior Convertible Notes due 2017:		
Principal amount	\$63,302	\$63,317
Unamortized debt discount	—	(1,417)
Unamortized debt issuance costs	—	(199)

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	63,302	61,701
2.25% Senior Convertible Notes due 2021:		
Principal amount	650,000	650,000
Unamortized debt discount	(64,880)	(72,713)
Unamortized debt issuance costs	(11,588)	(12,875)
	573,532	564,412
Total Senior Convertible Notes	\$636,834	\$626,113
Less: Current portion	(63,302)	(61,701)
Long-term Senior Convertible Notes	\$573,532	\$564,412

2.25% Senior Convertible Notes due 2021

In March 2016, the Company issued \$650.0 million principal amount of unsecured Senior Convertible Notes with a stated interest rate of 2.25% and a maturity date of March 15, 2021 (the "2021 Notes"). The net proceeds from the offering, after deducting initial purchasers' discounts and costs directly related to the offering, were approximately \$634.1 million. The 2021 Notes may be settled in cash, stock, or a combination thereof, solely at the Company's discretion. It is the Company's current intent and policy to settle all conversions through combination settlement, which involves satisfying the principal amount outstanding with cash and any note conversion value over the principal amount in shares of the Company's common stock. The initial conversion rate of the 2021 Notes is 16.7158 shares per \$1,000 principal amount, which is equivalent to a conversion price of approximately \$59.82 per share, subject to adjustments. The Company uses the treasury share method for assumed conversion of the 2021 Notes to compute the weighted average shares of common stock outstanding for diluted earnings per share. The Company also entered into transactions for convertible note hedge (the "2021 Hedge") and warrants (the "2021 Warrants") concurrently with the issuance of the 2021 Notes.

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The cash conversion feature of the 2021 Notes required bifurcation from the Notes and was initially accounted for as an equity instrument classified to stockholders' equity, which resulted in recognizing \$84.8 million in additional paid-in-capital during 2016.

The interest expense recognized on the 2021 Notes during the three months ended June 30, 2017 includes \$3.7 million, \$3.9 million and \$0.7 million for the contractual coupon interest, the accretion of the debt discount and the amortization of the debt issuance costs, respectively. The interest expense recognized on the 2021 Notes during the six months ended June 30, 2017 includes \$7.3 million, \$7.8 million and \$1.3 million for the contractual coupon interest, the accretion of the debt discount and the amortization of the debt issuance costs, respectively. The interest expense recognized on the 2021 Notes during the three months ended June 30, 2016 includes \$3.7 million, \$3.7 million and \$0.6 million for the contractual coupon interest, the accretion of the debt discount and the amortization of the debt issuance costs, respectively. The interest expense recognized on the 2021 Notes during the six months ended June 30, 2016 includes \$4.3 million, \$4.4 million and \$0.7 million for the contractual coupon interest, the accretion of the debt discount and the amortization of the debt issuance costs, respectively. The effective interest rate on the 2021 Notes is 5.8%, which includes the interest on the notes, amortization of the debt discount and debt issuance costs. Interest on the 2021 Notes began accruing upon issuance and is payable semi-annually.

Prior to September 15, 2020, holders may convert their 2021 Notes only under the following conditions: (a) during any calendar quarter beginning June 30, 2016, if the reported sale price of the Company's common stock for at least 20 days out of 30 consecutive trading days ending on the last trading day of the immediately preceding calendar quarter is greater than 130% of the conversion price on each applicable trading day; (b) during the five business day period in which the trading price of the 2021 Notes falls below 98% of the product of (i) the last reported sale price of the Company's common stock and (ii) the conversion rate on that date; and (c) upon the occurrence of specified corporate events, as defined in the 2021 Notes. From September 15, 2020 and until the close of business on the second scheduled trading day immediately preceding March 15, 2021, holders may convert their 2021 Notes at any time (regardless of the foregoing circumstances). The Company may not redeem the 2021 Notes prior to March 20, 2019. The Company may redeem the 2021 Notes, at its option, in whole or in part on or after March 20, 2019 until the close of business on the business day immediately preceding September 15, 2020 if the last reported sale price of the Company's common stock has been at least 130% of the conversion price then in effect for at least 20 trading days during any 30 consecutive trading day period ending on, and including, the trading day immediately preceding the date on which the Company delivers written notice of a redemption. The redemption price will be equal to 100% of the principal amount of such 2021 Notes to be redeemed plus accrued and unpaid interest to, but excluding, the redemption date. No principal payments are due on the 2021 Notes prior to maturity. Other than restrictions relating to certain fundamental changes and consolidations, mergers or asset sales and customary anti-dilution adjustments, the 2021 Notes do not contain any financial covenants and do not restrict the Company from paying dividends or issuing or repurchasing any of its other securities. The Company is unaware of any current events or market conditions that would allow holders to convert the 2021 Notes.

2021 Hedge

In connection with the offering of the 2021 Notes, the Company entered into the hedge transaction with the initial purchasers and/or their affiliates (the "2021 Counterparties") entitling the Company to purchase up to 10,865,270 shares of the Company's common stock at an initial stock price of \$59.82 per share, each of which is subject to adjustment. The cost of the 2021 Hedge was \$111.2 million and accounted for as an equity instrument by recognizing \$111.2 million in additional paid-in-capital during 2016. The 2021 Hedge will expire on March 15, 2021.

The 2021 Hedge is expected to reduce the potential equity dilution upon conversion of the 2021 Notes if the daily volume-weighted average price per share of the Company's common stock exceeds the strike price of the 2021 Hedge. An assumed exercise of the 2021 Hedge by the Company is considered anti-dilutive since the effect of the inclusion would always be anti-dilutive with respect to the calculation of diluted earnings per share.

2021 Warrants

The Company sold warrants to the 2021 Counterparties to acquire up to 10,865,270 shares of the Company's common stock. The 2021 Warrants will expire on various dates from June 2021 through December 2021 and may be settled in cash or net shares. It is the Company's current intent and policy to settle all conversions in shares of the Company's common stock. The Company received \$44.9 million in cash proceeds from the sale of the 2021 Warrants, which was recorded in additional paid-in-capital. The 2021 Warrants could have a dilutive effect on the Company's earnings per share to the extent that the price of the Company's common stock during a given measurement period exceeds the strike price of the 2021 Warrants, which is \$80.00 per share. The Company uses the treasury share method for assumed conversion of its 2021 Warrants to compute the weighted average common shares outstanding for diluted earnings per share.

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Repurchases of Senior Convertible Notes due 2017

In March 2016, the Company used approximately \$345.2 million of the net proceeds from the 2017 Notes offering to repurchase approximately \$276.8 million principal amount outstanding of the Senior Convertible Notes due 2017 (the “2017 Notes”), the associated conversion feature of the repurchased notes (which is recorded in additional paid-in capital), and the accrued interest on the repurchased notes. Subsequently, in the fourth quarter of 2016, the Company used approximately \$96.3 million of cash on hand to repurchase an additional \$62.3 million in principal amount outstanding of 2017 Notes, the associated conversion feature of the repurchased notes (which is recorded in additional paid-in capital), and the accrued interest on the repurchased notes. The repurchases of 2017 Notes in 2016 resulted in a cumulative loss of approximately \$19.1 million, including \$17.4 million recorded during the six months ended June 30, 2016. The Company recorded the loss on the repurchases of 2017 Notes in other expense on the accompanying Consolidated Statements of Operations. The loss on the repurchases included the related debt issuance costs that were previously capitalized in connection with the issuance of the 2017 Notes. The remaining balances resulting from the aggregate repurchase of a portion of the 2017 Notes were \$63.3 million, \$1.4 million, and \$0.2 million of principal outstanding, debt discount, and debt issuance costs, respectively, immediately following the repurchase.

2.75% Senior Convertible Notes due 2017

In June 2011, the Company issued \$402.5 million principal amount of 2017 Notes with a stated interest rate of 2.75% and a maturity date of July 1, 2017. The net proceeds from the offering, after deducting initial purchasers’ discounts and costs directly related to the offering, were approximately \$359.2 million. The 2017 Notes may be settled in cash, stock, or a combination thereof, solely at the Company’s discretion. It is the Company’s current intent and policy to settle all conversions through combination settlement, which involves satisfying the principal amount outstanding with cash and any note conversion value over the principal amount in shares of the Company’s common stock. The initial conversion rate of the 2017 Notes is 23.7344 shares per \$1,000 principal amount, which is equivalent to a conversion price of approximately \$42.13 per share, subject to adjustments. The Company uses the treasury share method for assumed conversion of the 2017 Notes to compute the weighted average shares of common stock outstanding for diluted earnings per share. The Company also entered into transactions for convertible note hedge (the “2017 Hedge”) and warrants (the “2017 Warrants”) concurrently with the issuance of the 2017 Notes.

The interest expense recognized on the 2017 Notes during the three months ended June 30, 2017 includes \$0.4 million, \$0.7 million and \$0.1 million for the contractual coupon interest, the accretion of the debt discount and the amortization of debt issuance costs, respectively. The interest expense recognized on the 2017 Notes during the six months ended June 30, 2017 includes \$0.9 million, \$1.4 million and \$0.2 million for the contractual coupon interest, the accretion of the debt discount and the amortization of the debt issuance costs, respectively. The interest expense recognized on the 2017 Notes during the three months ended June 30, 2016 includes \$0.9 million, \$1.3 million and \$0.2 million for the contractual coupon interest, the accretion of the debt discount and the amortization of debt issuance costs, respectively. The interest expense recognized on the 2017 Notes during the six months ended June 30, 2016 includes \$3.3 million, \$5.0 million and \$0.7 million for the contractual coupon interest, the accretion of the debt discount and the amortization of the debt issuance costs, respectively. The effective interest rate on the 2017 Notes is 8.0%, which includes the interest on the notes, amortization of the debt discount and debt issuance costs. Interest on the 2017 Notes began accruing upon issuance and is payable semi-annually.

Prior to January 1, 2017, holders could convert their 2017 Notes only under the following conditions: (a) during any calendar quarter beginning October 1, 2011, if the reported sale price of the Company’s common stock for at least 20

days out of 30 consecutive trading days ending on the last trading day of the immediately preceding calendar quarter is greater than 130% of the conversion price on each applicable trading day; (b) during the five business day period in which the trading price of the 2017 Notes falls below 98% of the product of (i) the last reported sale price of the Company's common stock and (ii) the conversion rate on that date; and (c) upon the occurrence of specified corporate events, as defined in the 2017 Notes. From January 1, 2017 and until the close of business on the second scheduled trading day immediately preceding July 1, 2017, holders could convert their 2017 Notes at any time (regardless of the foregoing circumstances). The Company could not redeem the 2017 Notes prior to maturity. Other than restrictions relating to certain fundamental changes and consolidations, mergers or asset sales and customary anti-dilution adjustments, the 2017 Notes do not contain any financial covenants and do not restrict the Company from paying dividends or issuing or repurchasing any of its other securities. As of June 30, 2017, a minimal amount of holders of the 2017 Notes had elected to convert their notes. The Company settled such conversions through the combination settlement described above. The 2017 Notes are recorded as current liabilities on the June 30, 2017 Consolidated Balance Sheet.

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2017 Hedge

In connection with the offering of the 2017 Notes, the Company entered into the 2017 Hedge with the initial purchasers and/or their affiliates (the “2017 Counterparties”) entitling the Company to purchase up to 9,553,096 shares of the Company’s common stock at an initial stock price of \$42.13 per share, each of which is subject to adjustment. The cost of the 2017 Hedge was \$80.1 million and accounted for as derivative assets upon issuance of the 2017 Notes. Upon obtaining stockholder approval for the additional authorized shares of the Company’s common stock, the derivative asset was reclassified to stockholders’ equity, which resulted in recognizing cumulatively \$37.1 million in other expense for the change in fair value measurement and \$43.0 million in additional paid-in-capital during 2011. The 2017 Hedge has an expiration date of July 1, 2017. The 2017 Hedge is expected to reduce the potential equity dilution upon conversion of the 2017 Notes if the daily volume-weighted average price per share of the Company’s common stock exceeds the strike price of the 2017 Hedge. An assumed exercise of the 2017 Hedge by the Company is considered anti-dilutive since the effect of inclusion would always be anti-dilutive with respect to the calculation of diluted earnings per share.

2017 Warrants

The Company sold warrants to the 2017 Counterparties to acquire up to 477,654 shares of the Company’s Series A Participating Preferred Stock at an initial strike price of \$988.51 per share, subject to adjustment. Each share of Series A Participating Preferred Stock is convertible into 20 shares of the Company’s common stock, or up to 9,553,080 common shares in total. The 2017 Warrants will expire on various dates from September 2017 through January 2018 and may be settled in cash or net shares. It is the Company’s current intent and policy to settle all conversions in shares of the Company’s common stock. The Company received \$47.9 million in cash proceeds from the sale of the 2017 Warrants, which was recorded in additional paid-in-capital. The 2017 Warrants could have a dilutive effect on the Company’s earnings per share to the extent that the price of the Company’s common stock during a given measurement period exceeds the strike price of the 2017 Warrants. The Company uses the treasury share method for assumed conversion of its 2017 Warrants to compute the weighted average common shares outstanding for diluted earnings per share.

Settlement of 2017 Notes, 2017 Hedge and 2017 Warrants

On July 1, 2017, the 2017 Notes reached maturity and a majority of the holders elected to convert their outstanding notes. The Company paid \$64.0 million in cash for the settlement of the outstanding principal amount including accrued interest and issued 650,070 shares for settlement of the conversion value over the principal amount of the notes. On the same date, the Company exercised the 2017 Hedge and received 4,160,789 shares of its own common stock on a net share basis from the 2017 Counterparties.

On May 24, 2017, the Company entered into warrant termination agreements with the 2017 Counterparties to settle the outstanding 2017 Warrants by accelerating the expiration period to varying settlement dates from June 2017 through July 2017, which terminated the existing 2017 Warrants settlement period. The settlement will be delivered in shares of Company common stock, based on a fixed formula using the daily volume weighted average price as the settlement measure. As of June 30, 2017, 2017 Warrants with respect to an aggregate of 6,100,000 shares were settled on a net share basis, resulting in the issuance of 2,328,351 shares of the Company’s common stock to the 2017 Counterparties. The remaining 2017 Warrants, with respect to an aggregate of 3,453,096 shares, will be settled on a net share basis throughout July 2017.

See Note 13 to the Unaudited Consolidated Financial Statements for further discussion on the settlement of the 2017 Notes, 2017 Hedge and 2017 Warrants.

Revolving Senior Credit Facility

In April 2017, the Company entered into an Amended and Restated Credit Agreement (the “2017 Credit Agreement”) for a revolving senior credit facility (the “2017 Facility”), which replaced the previous Credit Agreement the Company had entered into in February 2016. The 2017 Credit Agreement provides for secured revolving loans, multicurrency loan options and letters of credit in an aggregate amount of up to \$500.0 million. The 2017 Credit Agreement also contains an expansion feature, which allows the Company to increase the aggregate principal amount of the 2017 Facility provided the Company remains in compliance with the underlying financial covenants, including but not limited to, compliance with the consolidated interest coverage ratio and certain consolidated leverage ratios. The 2017 Facility matures in April 2022 (subject to an earlier springing maturity date), and includes a sublimit of \$100.0 million for multicurrency borrowings, a sublimit of \$50.0 million for the issuance of standby letters of credit, and a sublimit of \$5.0 million for swingline loans. All assets of the Company and its material domestic subsidiaries are pledged as collateral under the 2017 Facility (subject to customary exceptions) pursuant to the term set forth in the Amended and Restated Security and Pledge Agreement (the “2017 Security Agreement”) executed in favor of the administrative agent by the Company. Each of the Company’s material domestic subsidiaries guarantees the 2017 Facility. In connection with the 2017 Facility, the Company incurred issuance costs which will be amortized over the term of the 2017 Facility. As of June 30, 2017, the Company had \$20.0 million outstanding under the 2017 Facility, at an interest rate of 2.97% (one month LIBOR plus 1.75%).

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Borrowings under the 2017 Facility are used by the Company to provide financing for working capital and other general corporate purposes, including potential mergers and acquisitions. Borrowings under the 2017 Facility bear interest, at the Company's option, at a rate equal to an applicable margin plus: (a) the applicable Eurocurrency Rate (as defined in the 2017 Credit Agreement), or (b) a base rate determined by reference to the highest of (1) the federal funds effective rate plus 0.50%, (2) the Bank of America prime rate, and (3) LIBOR for an interest period of one month plus 1.00%. The margin for the 2017 Facility ranges, based on the Company's consolidated leverage ratio, from 0.00% to 1.00% in the case of base rate loans and from 1.00% to 2.00% in the case of Eurocurrency Rate loans. The 2017 Facility includes an unused line fee ranging, based on the Company's consolidated leverage ratio, from 0.20% to 0.35% per annum on the revolving commitment.

The 2017 Credit Agreement contains affirmative, negative, permitted acquisition and financial covenants, and events of default customary for financings of this type. The financial covenants require the Company to maintain ratios of consolidated earnings before interest, taxes, depreciation and amortization (EBITDA) in relation to consolidated interest expense and consolidated debt, respectively, as defined in the 2017 Credit Agreement. The 2017 Facility grants the lenders preferred first priority liens and security interests in capital stock, intercompany debt and all of the present and future property and assets of the Company and each guarantor. The Company is currently in compliance with the 2017 Credit Agreement covenants.

7. Stock-Based Compensation

The compensation cost that has been included in the Consolidated Statements of Operations for all stock-based compensation arrangements was as follows:

(in thousands)	Three Months		Six Months Ended	
	Ended June 30,		June 30,	
	2017	2016	2017	2016
Sales, marketing and administrative expense	\$7,891	\$7,415	\$14,686	\$11,846
Research and development expense	428	397	567	406
Cost of goods sold	75	53	158	105
Stock-based compensation expense before taxes	8,394	7,865	15,411	12,357
Related income tax benefits	(3,190)	(3,146)	(5,856)	(4,943)
Stock-based compensation expense, net of taxes	\$5,204	\$4,719	\$9,555	\$7,414

At June 30, 2017, there was \$62.7 million of unamortized compensation expense for restricted stock units ("RSUs") and performance-based restricted stock units ("PRSUs") to be recognized over a weighted average period of 2.3 years.

Restricted Stock Units

The Company issued approximately 16,000 and 316,000 shares of common stock, before net share settlement, upon vesting of RSUs (including PRSUs) during the three and six months ended June 30, 2017, respectively, and issued approximately 772,000 shares of common stock in settlement of RSUs (including PRSUs) upon their vesting during the year ended December 31, 2016.

Stock Options and Purchase Rights

The weighted average assumptions used to estimate the fair value of stock purchase rights under the employee stock purchase plan (“ESPP”) are as follows:

	Three Months Ended June 30, 2017		Six Months Ended June 30, 2016	
ESPP				
Volatility	21 %	31 %	23 %	31 %
Expected term (years)	0.5	0.5	0.5	0.6
Risk free interest rate	0.8 %	0.4 %	0.6 %	0.3 %
Expected dividend yield	— %	— %	— %	— %

Under the terms of the ESPP, shareowners can elect to have up to 15% of their annual compensation, up to a maximum of \$21,250 per year, withheld to purchase shares of the Company’s common stock for a purchase price equal to 85% of the lower of the fair market value per share (at closing) of the Company’s common stock on (i) the commencement date of the two-year or six-month offering period (depending on the purchase period enrolled), or (ii) the respective purchase date.

The Company has not granted any options since 2011. The Company issued approximately 23,000 and 162,000 shares of common stock, before net share settlement, upon the exercise of outstanding stock options during the three and six months ended June 30, 2017, respectively, and issued approximately 1,556,000 shares of common stock upon the exercise of outstanding stock options during the year ended December 31, 2016.

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8. Income Taxes

Income taxes are determined using an estimated annual effective tax rate applied against income, and then adjusted for the tax impacts of certain significant and discrete items. For the six months ended June 30, 2017, the Company treated the tax impact of the following as discrete events for which the tax effect was recognized separately from the application of the annual effective tax rate: tax benefits related to excess share-based payments and certain losses for which the Company receives no tax benefit. The Company's effective tax rate recorded for the six months ended June 30, 2017 was 26%.

In accordance with the disclosure requirements as described in ASC Topic 740, Income Taxes, the Company has classified unrecognized tax benefits as non-current income tax liabilities, or a reduction in deferred tax assets, unless expected to be paid within one year. The Company's continuing practice is to recognize interest and/or penalties related to income tax matters in income tax expense. The Company had an increase in gross unrecognized tax benefits of approximately \$0.5 million during the six months ended June 30, 2017, primarily related to research and development credits and domestic production activities deductions. The Company does not anticipate that there will be a significant change in unrecognized tax benefits within the next 12 months.

The Company is subject to routine compliance reviews on various tax matters around the world in the ordinary course of business. Currently, income tax audits are being conducted in the state of New York and the state of Louisiana. U.S. and most foreign jurisdictions remain subject to examination in all years due to prior year net operating losses and R&D credits.

9. Business Segment, Product and Geographic Information

The Company operates in one segment based upon the Company's organizational structure, the way in which the operations and investments are managed and evaluated by the chief operating decision maker ("CODM") as well as the lack of availability of discrete financial information at a lower level. The Company's CODM reviews revenue at the product line offering level, and manufacturing, operating income and expenses, and net income at the Company wide level to allocate resources and assess the Company's overall performance. The Company shares common, centralized support functions, including finance, human resources, legal, information technology, and corporate marketing, all of which report directly to the CODM. Accordingly, decision-making regarding the Company's overall operating performance and allocation of Company resources is assessed on a consolidated basis. As such, the Company operates as one reporting segment. The Company has disclosed the revenues for each of its product line offerings to provide the reader of the financial statements transparency into the operations of the Company.

The Company reports under two distinct product lines; spinal hardware and surgical support. The Company's spinal hardware product line offerings include implants and fixation products. The Company's surgical support product offerings include IOM services, disposables and biologics, all of which are used to aid spinal surgery.

Revenue by product line was as follows:

Three Months Ended	Six Months Ended
June 30,	June 30,

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(in thousands)	2017	2016	2017	2016
Spinal Hardware	\$183,589	\$171,242	\$357,293	\$323,199
Surgical Support	76,984	64,968	153,144	128,115
Total Revenue	\$260,573	\$236,210	\$510,437	\$451,314

Revenue and property and equipment, net, by geographic area were as follows:

(in thousands)	Revenue				Property and Equipment, Net	
	Three Months Ended June 30,		Six Months Ended June 30,		June 30,	December 31,
	2017	2016	2017	2016	2017	2016
United States	\$217,147	\$200,599	\$431,354	\$388,949	\$177,657	\$148,227
International (excludes Puerto Rico)	43,426	35,611	79,083	62,365	36,944	33,297
Total	\$260,573	\$236,210	\$510,437	\$451,314	\$214,601	\$181,524

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10. Commitments

Licensing and Purchasing Agreements

As of June 30, 2017 the Company has obligations under certain consulting arrangements to pay up to approximately \$20.4 million in the aggregate in the event that specified revenue-based milestones are achieved prior to 2024. Any such payment will be made in a combination of cash and the Company's common shares as provided in the agreements. Any payments in satisfaction of these contingent obligations are considered a cost of goods sold and are recognized ratably as and if milestones are achieved. These agreements expire on various dates through 2024.

Executive Severance Plans

The Company has employment contracts with key executives and maintains severance plans that provide for the payment of severance and other benefits if terminated for reasons other than cause, as defined in those agreements and plans. Certain agreements call for payments that are based on historical compensation, accordingly, the amount of the contractual commitment will change over time commensurate with the executive's applicable earnings. At June 30, 2017, future commitments for such key executives were approximately \$32.6 million. In certain circumstances, the agreements call for the acceleration of equity vesting. Those figures are not reflected in the above information.

11. Contingencies

The Company is subject to potential liabilities under government regulations and various claims and legal actions that are pending or may be asserted from time-to-time. These matters arise in the ordinary course and conduct of the Company's business and include, for example, commercial, intellectual property, environmental, securities and employment matters. The Company intends to continue to defend itself vigorously in such matters. Furthermore, the Company regularly assesses contingencies to determine the degree of probability and range of possible loss for potential accrual in its financial statements.

An estimated loss contingency is accrued in the Company's financial statements if it is probable that a liability has been incurred and the amount of the loss can be reasonably estimated. Based on the Company's assessment, it has adequately accrued an amount for contingent liabilities currently in existence. The Company does not accrue amounts for liabilities that it does not believe are probable or that it considers immaterial to its overall financial position. Litigation is inherently unpredictable, and unfavorable resolutions could occur. As a result, assessing contingencies is highly subjective and requires judgment about future events. The amount of ultimate loss may exceed the Company's current accruals, and it is possible that its cash flows or results of operations could be materially affected in any particular period by the unfavorable resolution of one or more of these contingencies.

Legal Proceedings

Medtronic Sofamor Danek USA, Inc. Litigation

In August 2008, Medtronic filed a patent infringement lawsuit against the Company (the "Medtronic Litigation"), alleging that certain of the Company's products or methods, including the XLIF procedure, infringe, or contribute to the infringement of, various U.S. patents assigned or licensed to Medtronic. The Company brought counterclaims against Medtronic alleging infringement of certain of the Company's patents. On July 13, 2016, the Company entered

into a settlement and patent license agreement (the “2016 Settlement Agreement”) with Medtronic to settle the Medtronic Litigation. The Company no longer has any remaining liability or restricted cash related to this matter.

The Medtronic Litigation was administratively broken into three phases. The initial trial on the first phase of the case concluded in September 2011 in the U.S. District Court for the Southern District of California (the “District Court”), and a jury delivered an unfavorable verdict against the Company with respect to certain Medtronic patents and a favorable verdict with respect to one Company patent, including a monetary damages award of approximately \$101.2 million to Medtronic.

Both parties appealed the verdict, and the Company entered into an escrow arrangement and transferred \$113.3 million of cash into a restricted escrow account in March 2012 to secure the amount of judgment, plus prejudgment interest, during pendency of the appeal. In March 2015, the U.S. Court of Appeals for the Federal Circuit issued a decision upholding the jury’s findings of liability as to all patents, but overturning the damage award against the Company as improper (the “Court of Appeals Decision”). The case was remanded back to the District Court for further proceedings and a retrial to determine a proper damages award. As a result of the Court of Appeals Decision, the parties agreed to release all of the escrow funds related to this matter back to the Company. During the year ended December 31, 2015, the Company transferred all of the funds in escrow related to this matter, approximately \$114.1 million, from long-term restricted cash and investments into its unrestricted investment accounts. In March 2015, the Company sought reexamination of certain claims of one of the Medtronic patents at issue and for which the Company was found to have infringed. On June 15, 2016, the District Court stayed remand proceedings and retrial of this first phase of the case pending the reexamination.

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The second phase of the case involved one Medtronic cervical plate patent. In April 2013, the Company and Medtronic entered into a settlement agreement fully resolving the second phase of the case. As part of the settlement, the Company received a license to practice various patent families that collectively represent a majority of Medtronic's patent rights related to cervical plate technology. In exchange for these license rights, the Company made a one-time payment to Medtronic of \$7.5 million in May 2013. In addition, Medtronic will receive a royalty on certain cervical plate products sold by the Company, including the Helix and Gradient lines of products.

The third phase of the case involved Medtronic filing additional patent claims in the U.S. District Court for the Northern District of Indiana in August 2012 alleging that certain Company spinal implants (including its CoRoent XL family of spinal implants), the Company's Osteocel Plus bone graft product, and the Company's XLIF procedure and use of MaXcess IV retractor during the XLIF procedure infringe several Medtronic patents.

Under the terms of the 2016 Settlement Agreement, the Company paid Medtronic \$45.0 million, and the parties released each other from, inter alia, any and all past patent infringement arising from the Medtronic Litigation. As a result, the Company adjusted its litigation accrual from \$88.3 million to \$45.0 million and recorded a \$43.3 million gain in the Consolidated Statement of Operations during the three months ended June 30, 2016. Pursuant to the 2016 Settlement Agreement, the parties granted each other irrevocable, worldwide, nonexclusive, paid-up, royalty-free licenses to practice certain of their respective patents as to certain of their respective existing product lines, subject to specified exceptions and limitations. The 2016 Settlement Agreement also provides that, subject to certain limitations and exceptions, and for a period of seven years, neither party will assert against the other certain claims for patent infringement (generally claims related to spinal implants and related instruments, biologics and neuromonitoring) other than through a specified dispute resolution process, with the right to thereafter pursue claims outside that process subject to certain limitations and exceptions. Further, Medtronic has agreed that, for a period of five years, and subject to limitations and exceptions, it will not assert against the Company certain other claims for patent infringement other than through a specified dispute resolution process, with the right to thereafter pursue claims outside that process subject to certain limitations and exceptions.

Securities Litigation

On August 28, 2013, a purported securities class action lawsuit was filed in the U.S. District Court for the Southern District of California naming the Company and certain of its current and former executive officers for allegedly making false and materially misleading statements regarding the Company's business and financial results, specifically relating to the purported improper submission of false claims to Medicare and Medicaid. The operative complaint asserts a putative class period stemming from October 22, 2008 to July 30, 2013. The complaint alleges violations of Sections 10(b) and 20(a) of the Securities Exchange Act of 1934, as amended, and Rule 10b-5 promulgated thereunder and seeks unspecified monetary relief, interest, and attorneys' fees. On February 13, 2014, Brad Mauss, the lead plaintiff in the case, filed an Amended Class Action Complaint for Violations of the Federal Securities Laws. The Company answered the complaint on August 25, 2016, and discovery is proceeding. The plaintiffs filed motions for class certification on October 28, 2016 and the Company's opposition papers were filed on January 9, 2017. On March 22, 2017, the court issued an order granting class certification. The Company filed a petition to appeal the order granting class certification with the U.S. Court of Appeals for the Ninth Circuit on April 5, 2017 and the plaintiffs have filed an opposition to the petition. Trial has been set for December 18, 2017. At June 30, 2017, the probable outcome of this litigation cannot be determined, nor can the Company estimate a range of potential loss. In accordance with authoritative guidance on the evaluation of loss contingencies, the Company has not recorded an accrual related to this litigation.

Shareholder Derivative Litigation

On September 28, 2016, a shareholder derivative complaint was filed by James Borta in the Superior Court of California for the County of San Diego naming certain of the Company's current and former executive officers and directors for allegedly breaching their fiduciary duties by, among other things, making allegedly false and misleading statements about the Company's business, operations, and prospects. The derivative complaint is based upon the same factual allegations as the securities class action litigation and names the Company as a nominal defendant. The plaintiff filed an Amended Complaint on March 1, 2017. The Company demurred to the Amended Complaint on April 7, 2017 and the court sustained the Company's demurrer and provided the plaintiff thirty days to file an amended complaint. On June 30, 2017 the plaintiff filed a Second Amended Derivative Complaint. At June 30, 2017, the probable outcome of this litigation cannot be determined, nor can the Company estimate a range of potential loss. In accordance with authoritative guidance on the evaluation of loss contingencies, the Company has not recorded an accrual related to this litigation.

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Madsen Medical, Inc. Litigation

On February 19, 2016, an unfavorable jury verdict was delivered against the Company in its litigation in the U.S. District Court for the Southern District of California against Madsen Medical, Inc. (“MMI”), a former sales agent. Specifically, the jury awarded MMI \$7.5 million in lost profits for tortious interference, \$14.0 million for unjust enrichment, \$20.0 million in punitive damages, and approximately \$0.3 million in damages for breach of contract. On March 18, 2016, the trial court entered judgment in favor of MMI in the amount of \$27.8 million, which amount excluded the \$14.0 million disgorgement awarded by the jury. On July 5, 2016, the trial court also awarded MMI attorney’s fees and costs of approximately \$1.1 million. The Company’s post-trial motions for judgment as a matter of law and/or for a new trial were denied, and the Company has appealed both the verdict and the court’s subsequent award of attorney’s fees and costs. However, the Company did not appeal the judgment with respect to breach of contract and accordingly accrued the \$0.3 million in damages during the six months ended June 30, 2017. During pendency of any appeals, the Company has secured a bond to cover the amount of the judgment and attorneys’ fees and costs.

Historically the Company had believed the likelihood of a loss in this case was remote given the underlying facts of the case, however, during the quarter ended March 31, 2016, the judgment entered caused the Company to reassess its position. The Company, based on its own assessment as well as that of outside counsel, believes that upon either post-trial motions or appeal the judgment will be vacated and have deemed it probable that is the outcome for all appealed judgments. The Company continues to believe for all judgments under appeal that such judgments will be vacated, and accordingly, at June 30, 2017, the Company believes that the outcome of the case does not constitute a probable nor an estimable loss associated with the litigation but rather a reasonably possible loss rather than a remote loss as historically contemplated. Therefore, for all judgments under appeal the Company has not recorded a loss contingency but has assessed a reasonable range of potential loss, which would be from zero to the current amount entered as a judgment, as well as attorney’s fees and interest, in accordance with the accounting guidance required by ASC 450, Contingencies.

12. Regulatory Matters

On August 31, 2015, the Company received a civil investigative demand (“CID”) issued by the Department of Justice (“DOJ”) pursuant to the federal False Claims Act. The CID requires the delivery of a wide range of documents and information related to an investigation by the DOJ concerning allegations that the Company assisted a physician group customer in submitting improper claims for reimbursement and made improper payments to the physician group in violation of the Anti-Kickback Statute. The Company is cooperating with the DOJ. No assurance can be given as to the timing or outcome of this investigation. At June 30, 2017, the probable outcome of this matter cannot be determined, nor can the Company estimate a range of potential loss. In accordance with authoritative guidance on the evaluation of loss contingencies, the Company has not recorded an accrual related to this matter.

On June 9, 2017, the Company received a subpoena from the Office of the Inspector General of the U.S. Department of Health and Human Services (“OIG”) in connection with an investigation into possible false or otherwise improper claims submitted to Medicare and Medicaid. The subpoena seeks discovery of documents for the period January 2014 through June 2017, primarily associated with sales to a particular customer and relationships related to that customer account. The Company is working with the OIG to understand the scope of the subpoena and its request for documents, and the Company intends to fully cooperate with the OIG’s request. No assurance can be given as to the timing or outcome of this investigation. At June 30, 2017, the probable outcome of this matter cannot be determined,

nor can the Company estimate a range of potential loss. In accordance with authoritative guidance on the evaluation of loss contingencies, the Company has not recorded an accrual related to this matter.

13. Subsequent Events

Settlement of 2017 Notes and 2017 Hedge

On July 1, 2017, the 2017 Notes reached maturity and a majority of the holders elected to convert their outstanding notes. The Company paid \$64.0 million in cash for the settlement of the outstanding principal amount including accrued interest and issued 650,070 shares for settlement of the conversion value over the principal amount of the notes. On the same date, the Company exercised the 2017 Hedge and received 4,160,789 shares of its own common stock on a net share basis from the 2017 Counterparties.

On May 24, 2017, the Company entered into warrant termination agreements with the 2017 Counterparties to settle the outstanding 2017 Warrants by accelerating the expiration period to varying settlement dates from June 2017 through July 2017, which terminated the existing 2017 Warrants settlement period. The settlement will be delivered in shares of Company common stock, based on a fixed formula using the daily volume weighted average price as the settlement measure. As of June 30, 2017, 2017 Warrants with respect to an aggregate of 6,100,000 shares were settled on a net share basis, resulting in the issuance of 2,328,351 shares of the Company's common stock to the 2017 Counterparties. The remaining 2017 Warrants, with respect to an aggregate of 3,453,096 shares, will be settled on a net share basis throughout July 2017.

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Item 2. Management's Discussion and Analysis of Financial Condition and Results of Operations
Forward-Looking Statements May Prove Inaccurate

This quarterly report on Form 10-Q ("Quarterly Report"), including the following discussion and analysis, may contain forward-looking statements that involve risks, uncertainties, assumptions and other factors which, if they do not materialize or prove correct, could cause our results to differ from historical results or those expressed or implied by such forward-looking statements. In some cases, you can identify these forward-looking statements by words like "may", "will", "should", "could", "expects", "plans", "anticipates", "believes", "estimates", "predicts", "potential", "intends" (the negative of those words and other comparable words). Forward-looking statements include, but are not limited to, statements about:

- our intentions, beliefs and expectations regarding our expenses, sales, operations and future financial performance;
- our operating results;
- our plans for future products and enhancements of existing products;
- anticipated growth and trends in our business;
- the timing of and our ability to maintain and obtain regulatory clearances or approvals;
- our belief that our cash and cash equivalents and investments will be sufficient to satisfy our anticipated cash requirements;
- our expectations regarding our revenues, customers and distributors;
- our beliefs and expectations regarding our market penetration and expansion efforts;
- our expectations regarding the benefits and integration of recently-acquired businesses and our ability to make future acquisitions and successfully integrate any such future-acquired businesses;
- our anticipated trends and challenges in the markets in which we operate; and
- our expectations and beliefs regarding and the impact of investigations, claims and litigation.

These statements are not guarantees of future performance or events. Our actual results may differ materially from those discussed here. The potential risks and uncertainties that could cause actual results to differ materially include, but are not limited to those set forth under the heading "Risk Factors" in our Annual Report on Form 10-K for the year ended December 31, 2016 and this Quarterly Report on Form 10-Q, and similar discussions in our other Securities and Exchange Commission filings. We assume no obligation to update any forward looking statements to reflect new information, future events or circumstances or otherwise.

This information should be read in conjunction with the consolidated financial statements and the notes thereto included in Part I, Item 1 of this Quarterly Report and with Management's Discussion and Analysis of Financial Condition and Results of Operations for the year ended December 31, 2016 contained in our 2016 Annual Report on Form 10-K.

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Overview

We are a leading medical device company in the global spine surgery market, focused on developing minimally-disruptive surgical products and procedurally-integrated solutions for spine surgery. Our currently-marketed product portfolio is focused on applications for spine fusion surgery, including ancillary products used to aid in the surgical procedure.

Our principal product offering includes a minimally-disruptive surgical platform called Maximum Access Surgery, or MAS. The MAS platform combines three categories of solutions that collectively minimize soft tissue disruption during spine fusion surgery, provide maximum visualization and are designed to enable safe and reproducible outcomes for the surgeon and the patient. The platform includes our proprietary software-driven nerve detection and avoidance systems, NVM5, and Intraoperative Monitoring, or IOM, services and support; MaXcess, an integrated split-blade retractor system; and a wide variety of specialized implants and biologics. Many of our products, including the individual components of our MAS platform can also be used in open or traditional spine surgery. Our spine surgery product line offerings, which include products for the thoracolumbar and the cervical spine, are primarily used to enable surgeon access to the spine to perform restorative and fusion procedures in a minimally-disruptive fashion. In May 2015, we launched Integrated Global Alignment, or iGA, in which products and computer assisted technology under our MAS platform help achieve more precise spinal alignment. Our biologics products, which are used to aid in the spinal fusion process or bone healing process, include allograft (donated human tissue) and synthetic offerings.

We believe our MAS platform and its related offerings provide a unique and comprehensive solution for the safe and reproducible minimally-disruptive surgical treatment of spine disorders by enabling surgeons to access the spine in a manner that affords both direct visualization and detection and avoidance of critical nerves. The fundamental difference between our MAS platform, which is sometimes referred to in the industry as “minimally invasive surgery” is the ability to customize safe and reproducible access to the spine while allowing surgeons to continue to use instruments that are familiar to them and effective during surgery. Accordingly, the MAS platform does not force surgeons to reinvent or learn new approaches that add complexity and undermine safety, ease of use and/or efficacy. We have dedicated and continue to dedicate significant resources toward training spine surgeons around the world; both those who are new to our MAS and other product platforms, as well as ongoing education for MAS-trained surgeons attending advanced courses. An important ongoing objective of ours has been to maintain a leading position in access and nerve avoidance, as well as to pioneer and remain the ongoing leader in minimally invasive spine surgery. Our MAS platform, with the unique advantages provided by our nerve monitoring systems, enables an innovative lateral procedure known as eXtreme Lateral Interbody Fusion, or XLIF, in which surgeons access the spine for a fusion procedure from the side of the patient’s body, rather than from the front or back. It has been demonstrated clinically that XLIF and other procedures facilitated by our MAS platform decrease trauma and blood loss, and lead to faster overall patient recovery times compared to open spine surgery.

We also design and sell expandable growing rod implant systems that can be non-invasively lengthened following implantation with precise, incremental adjustments via an external remote controller using magnetic technology called MAGnetic External Control, or MAGEC, which allows for the minimally invasive treatment of early-onset and adolescent scoliosis. This technology is also the basis for our PRECICE limb lengthening system, which allows for the correction of long bone limb length discrepancy, as well as enhanced bone healing in patients that have experienced traumatic injury.

We intend to continue development on a wide variety of projects intended to broaden surgical applications for greater procedural integration of our MAS techniques and additional applications of the MAGEC technology. Such applications include tumor, trauma, and deformity, as well as increased fixation options, sagittal alignment products, imaging and navigation. We also expect to continue expanding our other product and services offerings as we execute on our strategy to offer customers an end-to-end, integrated procedural solution for spine surgery. We intend to continue to pursue business and technology acquisition targets and strategic partnerships.

Revenues and Operations

To date, the majority of our revenues are derived from the sale of implants, biologics and disposables and we expect this trend to continue for the foreseeable future. Additionally, with our acquisition of BNN Holdings on July 1, 2016, we expect our IOM service and support revenue to increase compared to previous periods. We loan our proprietary software-driven nerve monitoring systems and surgical instrument sets at no cost to surgeons and hospitals that purchase disposables and implants for use in individual procedures. In addition, we often place our proprietary software-driven nerve monitoring systems, MaXcess and other MAS instrument sets with hospitals for an extended period at no up-front cost to them. Our implants, biologics and disposables are currently sold and shipped from our distribution and warehousing operations. We generally recognize revenue for implants, biologics and disposables upon receiving a purchase order from the hospital, or acknowledgment from the hospital indicating product use or implantation, or upon shipment to third-party customers who immediately accept title. Revenue from IOM services is recognized in the period the service is performed for the amount of payment we expect to receive. We sell MAS instrument sets, MaXcess devices, and our proprietary software-driven nerve monitoring systems, however this does not make up a material part of our business.

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The majority of our operations are located and the majority of our sales have been generated in the United States. We sell our products in the United States through a sales force comprised primarily of exclusive independent sales agents and directly-employed sales representatives, both engaged to sell only NuVasive products. Our sales force provides a delivery and consultative service to our surgeon and hospital customers and is compensated based on sales and product placements in their territories. Sales force commissions are reflected in the sales, marketing and administrative operating expense line item within our Statement of Operations. We continue to invest in international expansion with the focus on European, Asia-Pacific and Latin American markets. Our international sales force is comprised of directly-employed sales personnel, independent sales agents, as well as exclusive and non-exclusive independent third-party distributors.

Results of Operations

Revenue

	June 30,		\$		
(in thousands, except %)	2017	2016	Change	% Change	
Three Months Ended					
Revenue					
Spinal Hardware	\$183,589	\$171,242	\$12,347	7	%
Surgical Support	76,984	64,968	12,016	18	%
Total revenue	\$260,573	\$236,210	\$24,363	10	%
Six Months Ended					
Revenue					
Spinal Hardware	\$357,293	\$323,199	\$34,094	11	%
Surgical Support	153,144	128,115	25,029	20	%
Total revenue	\$510,437	\$451,314	\$59,123	13	%

Our spinal hardware product line offerings include our implants and fixation products. Our surgical support product line offerings include IOM services, disposables and biologics, all of which are used to aid spinal surgery.

The continued adoption of minimally invasive procedures for spine has led to the expansion of our procedure volume. In addition, increased market acceptance in our international markets contributed to the increase in revenues for the periods presented. We expect continued adoption of our innovative minimally invasive procedures and deeper penetration into existing accounts and international markets as our sales force executes on our strategy of selling the full mix of our products and services. However, the continued consolidation and increased purchasing power of our hospital customers and group purchasing organizations, the continued existence of physician-owned distributorships, recent changes in the public and private insurance markets regarding reimbursement, and ongoing policy and legislative changes in the United States have created less predictability in the lumbar portion of the spine market and have limited the domestic spine market's procedural growth rate. Accordingly, we believe that our growth in revenue in 2017 will come primarily from share gains in the shift toward less invasive spinal surgery, revenue from new products and services, and international growth.

Revenue from our spinal hardware product line offerings increased \$12.3 million and \$34.1 million, or 7% and 11%, during the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016. Revenue associated with our 2016 acquisitions accounted for approximately 2% of the increase in spinal hardware revenue for

the six months ended June 30, 2017, as compared to the same period in 2016. Product volume in spinal hardware, excluding 2016 acquisitions, increased our revenue by approximately 11%, offset by unfavorable pricing impacts of approximately 2% for both the three months and six months ended June 30, 2017, respectively, as compared to the same periods in 2016. Foreign currency fluctuation had an insignificant impact on revenue from spinal hardware for the periods presented.

Revenue from our surgical support product line offerings increased \$12.0 million and \$25.0 million, or 18% and 20%, during the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016. Revenue associated with our 2016 acquisitions accounted for approximately 23% of the increase in surgical support revenue for both the three and six months ended June 30, 2017, respectively, as compared to the same periods in 2016. This was offset by unfavorable volume and pricing impacts in surgical support, excluding 2016 acquisitions, of approximately 3% and 1%, respectively, for both the three months and six months ended June 30, 2017 as compared to the same periods in 2016. Foreign currency fluctuation had an insignificant impact on revenue from surgical support for the periods presented.

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Cost of Goods Sold, Excluding Below Amortization of Intangible Assets

(in thousands, except %)	June 30,		\$	% Change	
	2017	2016			
Three Months Ended					
Cost of goods sold (excluding below amortization of intangible assets)	\$66,421	\$59,745	\$6,676	11	%
% of total revenue	25	% 25	%		
Six Months Ended					
Cost of goods sold (excluding below amortization of intangible assets)	\$128,034	\$113,971	\$14,063	12	%
% of total revenue	25	% 25	%		

Cost of goods sold consists primarily of purchased goods, raw materials, labor and overhead associated with product manufacturing, inventory-related costs and royalty expenses, as well as the cost of providing IOM services, which includes personnel and physician oversight costs. We primarily procure and manufacture our goods in the United States, and accordingly, foreign currency fluctuations have not materially impacted our cost of goods sold.

Cost of goods sold increased \$6.7 million and \$14.1 million, or 11% and 12%, during the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016. The cost of goods sold associated with the operations of our 2016 acquisitions accounted for approximately 15% and 18% of the total increase during the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016. Cost of goods sold for our business, excluding our 2016 acquisitions, increased primarily due to growth in volume, but also includes shifts in purchase price and product mix, for an overall increase of approximately 10% and 11% for the three and six months ended June 30, 2017, respectively. These increases in cost of goods sold were partially offset by reduced expenses for approximately 14% and 17%, for the three and six months ended June 30, 2017, respectively, as compared to the same periods in 2016 which included inventory expense associated with the purchase accounting for our acquisition of Ellipse Technologies, non-recurring inventory expenses, and obsolescence reserves for a 2016 new product launch in the cervical market. Cost of goods sold as a percentage of revenue remained relatively consistent for the three and six months ended June 30, 2017 compared to the same periods in 2016 for the reasons described above.

On a long-term basis, we expect cost of goods sold, as a percentage of revenue, to decrease moderately.

Operating Expenses

(in thousands, except %)	Three Months Ended		\$	% Change	
	2017	2016			
June 30,					
Sales, marketing and administrative	\$139,109	134,487	\$4,622	3	%
% of total revenue	53	% 57	%		
Research and development	12,572	11,871	701	6	%
% of total revenue	5	% 5	%		
Amortization of intangible assets	11,349	10,603	746	7	%
Litigation liability (gain)	—	(43,310)	43,310	(100)	%

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	2017	2016	\$ Change	% Change
Business transition costs	1,369	2,756	(1,387)	(50)%
Six Months Ended June 30,				
(in thousands, except %)	2017	2016	\$ Change	% Change
Sales, marketing and administrative	\$279,611	259,325	\$20,286	8%
% of total revenue	55%	57%		
Research and development	24,986	22,500	2,486	11%
% of total revenue	5%	5%		
Amortization of intangible assets	23,410	18,474	4,936	27%
Litigation liability (gain)	—	(43,310)	43,310	(100)%
Business transition costs	1,424	8,063	(6,639)	(82)%

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Sales, Marketing and Administrative

Sales, marketing and administrative expenses consist primarily of compensation costs, commissions and training costs for shareowners engaged in sales, marketing and customer support functions. The expense also includes commissions to sales representatives, freight expenses, surgeon training costs, depreciation expense for property and equipment such as surgical instrument sets, and administrative expenses for both shareowners and third party service providers.

Sales, marketing and administrative expenses increased by \$4.6 million and \$20.3 million, or 3% and 8%, during the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016, primarily related to increases in shareowner compensation and related expenses due to increased headcount. Other costs which increased as a function of the increase in revenue and expansion included consulting, facilities, travel and equipment, all offset by a decrease in legal expense due to the settlement of the Medtronic litigation in 2016. Sales, marketing and administrative expenses associated with our 2016 acquisitions, which is included in the results discussed herein, accounted for approximately 2% and 4% of the increase in sales, marketing and administrative expenses for the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016.

Sales, marketing and administrative expenses as a percentage of revenue slightly decreased during the three and six months ended June 30, 2017 compared to the same periods in 2016. On a long-term basis, we expect total sales, marketing and administrative costs, as a percentage of revenue, to decrease moderately. To date, foreign currency fluctuations have not materially impacted our sales, marketing, and administrative expense.

Research and Development

Research and development expense consists primarily of product research and development, clinical trial and study costs, regulatory and clinical functions, and compensation and other shareowner related expenses. In the last several years, we have introduced numerous new products and product enhancements that have significantly expanded our MAS platform, including iGA, and our comprehensive product portfolio. We have also acquired complementary and strategic assets and technology, particularly in the area of spinal hardware products. We continue to invest in research and development programs.

Research and development expense increased by \$0.7 million and \$2.5 million, or 6% and 11%, during the three and six months ended June 30, 2017, respectively, compared to the same periods in 2016. The increase in spending is primarily due to increased spending for our integrated operative solutions technologies and expenses associated with our 2016 acquisitions.

Research and development costs as a percentage of revenue have remained relatively consistent during the three and six months ended June 30, 2017 compared to the same periods in 2016. On a long-term basis, we expect total research and development costs as a percentage of revenue to increase moderately in support of our ongoing development and regulatory approval efforts.

Litigation Liability Gain

During the three and six months ended June 30, 2016, we agreed to settle our ongoing litigation with Medtronic. As a result of the settlement, we agreed to pay \$45.0 million to Medtronic and accordingly recorded a gain of \$43.3 million related to the settlement by reducing our previous accrual of \$88.3 million related to the matter.

Interest and Other Expense, Net

(in thousands, except %)	June 30,		\$	% Change	
	2017	2016			
Three Months Ended					
Interest income	\$ 139	\$ 406	(267)	(66)	%
Interest expense	(10,083)	(10,537)	454	(4)	%
Other expense, net	(501)	(246)	(255)	104	%
Total interest and other expense, net	\$(10,445)	\$(10,377)	\$(68)	1	%
Six Months Ended					
Interest income	\$ 276	\$ 734	(458)	(62)	%
Interest expense	(19,882)	(19,009)	(873)	5	%
Loss on repurchases of convertible notes	—	(17,444)	17,444	(100)	%
Other expense, net	(243)	(196)	(47)	24	%
Total interest and other expense, net	\$(19,849)	\$(35,915)	\$ 16,066	(45)	%

Total interest expense was consistent during the three and six months ended June 30, 2017 compared to the same periods in 2016. A loss of \$17.4 million was recognized during the six months ended June 30, 2016 related to the repurchases of a portion of the Senior Convertible Notes due 2017. The total interest and other expense, net, for all periods presented included marginal income earned on marketable securities.

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Income Tax Expense

	June 30,			
(in thousands, except %)	2017	2016		
Three Months Ended				
Income tax expense	\$7,079	\$19,891		
Effective income tax rate	37 %	40 %		
Six Months Ended				
Income tax expense	\$8,569	\$10,411		
Effective income tax rate	26 %	29 %		

The provision for income tax expense as a percentage of pre-tax income from continuing operations was 37% for the three months ended June 30, 2017 compared with 40% for the three months ended June 30, 2016. The rate was lower in 2017 due primarily to stronger global earnings, increased benefits related to domestic production activities deductions and the elimination of certain tax expense items as a result of the early adoption of ASU 2016-16 during the first quarter of 2017, offset by lower tax benefits in 2017 related to excess share-based compensation payments.

The provision for income tax expense as a percentage of pre-tax income from continuing operations was 26% for the six months ended June 30, 2017 compared with 29% for the six months ended June 30, 2016. The rate was lower in 2017 due primarily to stronger global earnings, increased benefits related to domestic production activities deductions and the elimination of certain tax expense items as a result of the early adoption of ASU 2016-16 during the first quarter of 2017, offset by lower tax benefits in 2017 related to excess share-based compensation payments.

Liquidity, Cash Flows and Capital Resources

Liquidity and Capital Resources

Our principal sources of liquidity are our existing cash, cash equivalents and marketable securities, cash generated from operations, proceeds from our convertible notes issuances, and access to our revolving line of credit. We expect that cash provided by operating activities may fluctuate in future periods as a result of a number of factors, including fluctuations in our operating results, working capital requirements and capital deployment decisions. We have historically invested our cash primarily in the U.S. treasuries and government agencies, corporate debt, and money market funds. Certain of these investments are subject to general credit, liquidity and other market risks. The general condition of the financial markets and the economy may increase those risks and may affect the value and liquidity of investments and restrict ability to access the capital markets.

Our future capital requirements will depend on many factors including our rate of revenue growth, the timing and extent of spending to support development efforts, the expansion of sales, marketing and administrative activities, the timing of introductions of new products and enhancements to existing products, successful vertical integration of our manufacturing process, the continuing market acceptance of our products, the expenditures associated with possible future acquisitions or other business combination transactions, the outcome of current and future litigation, the evolution of our globalization initiative, and continuous international expansions of our business. We believe that our cash flow from operations and growing operations will continue to fund the ongoing core business. As current borrowing sources become due, we may be required to access the capital markets for additional funding. As we assess inorganic growth strategies, we may need to supplement our internally generated cash flow with outside sources. In the event that we are required to access the debt market, we believe we can do so at reasonable borrowing rates. As

part of our liquidity strategy, we will continue to monitor our current level of earnings and cash flow generation as well as our ability to access the market in light of those earning levels.

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A substantial portion of our operations are located in the United States, and the majority of our sales and cash generation since inception have been made in the United States. Accordingly, we do not have material net cash flow exposures to foreign currency rate fluctuations. However, as our business in markets outside of the United States continues to increase, we will be exposed to foreign currency exchange risk related to our foreign operations. Fluctuations in the rate of exchange between the United States dollar and foreign currencies, primarily in the pound sterling, the euro, the Australian dollar, the Singapore dollar, and the yen, could adversely affect our financial results, including our revenues, revenue growth rates, gross margins, income and losses as well as assets and liabilities. We enter into forward currency contracts to partially offset the impact from fluctuations of the foreign currency rates on our third party and short-term intercompany receivables and payables between our domestic and international operations. We currently do not hedge future forecasted transactions but will continue to assess whether that strategy is appropriate. At June 30, 2017, the cash balance held by our foreign subsidiaries with currencies other than the United States dollar was approximately \$29.9 million and it is our intention to indefinitely reinvest all of current foreign earnings in order to partially support foreign working capital and to expand our existing operations outside the United States. As of June 30, 2017, our account receivable balance held by our foreign subsidiaries with currencies other than the United States dollar was approximately \$31.0 million. We have operations in markets in which there is governmental financial instability which could impact funds that flow into the medical reimbursement system. In addition, loss of financial stability within these markets could lead to delays in reimbursement or inability to remit payment due to currency controls. Specifically, we have operations and/or sales in Puerto Rico, Brazil, Argentina and Venezuela. We do not have any material financial exposure to one customer or one country that would significantly hinder our liquidity.

On August 31, 2015, we received a civil investigative demand, or CID, issued by the Department of Justice, or DOJ, pursuant to the federal False Claims Act. The CID requires the delivery of a wide range of documents and information related to an investigation by the DOJ concerning allegations that we assisted a physician group customer in submitting improper claims for reimbursement and made improper payments to the physician group in violation of the Anti-Kickback Statute. We are cooperating with the DOJ. No assurance can be given as to the timing or outcome of this investigation, and the probable outcome of this matter cannot be determined.

On June 9, 2017, we received a subpoena from the Office of the Inspector General of the U.S. Department of Health and Human Services, or OIG, in connection with an investigation into possible false or otherwise improper claims submitted to Medicare and Medicaid. The subpoena seeks discovery of documents for the period January 2014 through June 2017, primarily associated with sales to a particular customer and relationships related to that customer account. We are working with the OIG to understand the scope of the subpoena and its request for documents, and we intend to fully cooperate with the OIG's request. No assurance can be given as to the timing or outcome of this investigation, and the probable outcome of this matter cannot be determined.

We are involved in a number of legal actions and investigations arising out of the normal course of our business as discussed in Note 11 of the Unaudited Consolidated Financial Statements. Due to the inherent uncertainties associated with pending legal actions and investigations, we cannot predict the outcome, and, with respect to certain pending litigation or claims where no liability has been accrued, to make a meaningful estimate of the reasonably possible loss or range of loss that could result from an unfavorable outcome, other than those matters disclosed in this Quarterly Report. We have no material accruals for pending litigation or claims for which accrual amounts are not disclosed in our Unaudited Consolidated Financial Statements. It is reasonably possible, however, that an unfavorable outcome that exceeds our current accrual estimate, if any, for one or more of the matters described in our Unaudited Consolidated Financial Statements could have a material adverse effect on our liquidity and access to capital

resources. Additionally, it is possible that as part of the ongoing legal appeals process, regardless of our assessment of the probability of a loss, we could be required to set aside funds in an escrow or purchase a performance bond. These requirements to escrow funding could have an adverse impact on our ability to access our current liquidity or impact our access to additional capital resources.

On July 1, 2017, the Senior Convertible Notes due 2017, which we refer to as the 2017 Notes, reached maturity and a majority of the holders elected to convert their outstanding notes. We paid \$64.0 million in cash for the settlement of the outstanding principal amount including accrued interest and issued 650,070 shares for settlement of the conversion value over the principal amount of the notes. Refer to the below section subtitled “2.75% Senior Convertible Notes due 2017” for further details.

On September 12, 2016, we completed an acquisition of an imaging software and technology platform known as LessRay. In connection with the acquisition we recorded a purchase accounting fair value estimate of \$34.1 million for contingent consideration liabilities related to the achievement of certain regulatory and commercial milestones. We anticipate these milestones will become payable at varying times between 2017 and 2020. We expect the imaging software and technology platform to be incorporated into our MAS platform to form a foundational element in our imaging, navigation and automation platform development strategy.

On February 11, 2016, we acquired Ellipse Technologies for an upfront payment of \$380.0 million (including holdbacks for retained employment of Ellipse Technologies leadership that is to be expensed and is not considered part of the final purchase price) and a potential milestone payment of \$30.0 million payable in 2017 related to the achievement of a specific revenue target. The revenue-based milestone was achieved as of December 31, 2016. We paid the milestone in April 2017, and no additional consideration is owed related to the acquisition.

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In furtherance of our initiative to increase the amount of products that we self-manufacture, in 2015, we added an approximately 180,000 square foot manufacturing facility in West Carrollton, Ohio; we have built out and equipped the new facility and initial production is underway.

Cash, cash equivalents and marketable securities were \$130.9 million and \$153.6 million at June 30, 2017 and December 31, 2016, respectively. We believe that our existing cash, cash equivalents, marketable securities and available liquidity will be sufficient to meet our anticipated cash needs for the next twelve months. We could have varying needs for cash as a result of the achievement of certain acquisition related milestones. We anticipate funding these milestones from cash on hand and operations, however, we also have the ability to fund these from our existing line of credit if necessary. The change in liquidity during the six months ended June 30, 2017 of \$22.7 million was mainly driven by \$68.7 million in cash used for purchases of property and equipment, \$30.0 million in cash used for a contingent consideration payment to Ellipse Technologies, \$14.4 million in cash used for strategic investments, and \$10.8 million in cash used on treasury stock purchases, offset by a \$20.0 million draw on the line of credit and \$67.1 million from cash inflow from operations. At June 30, 2017, we have cash totaling \$7.3 million in restricted accounts which are not available to us to meet any ongoing capital requirements if and when needed. Future litigation or requirements to escrow funds could materially impact our liquidity and our ability to invest in and run our business on an ongoing basis.

Cash Flows from Operating Activities

Cash provided by operating activities was \$67.1 million for the six months ended June 30, 2017, compared to \$100.2 million for the same period in 2016. The \$33.1 million decrease in cash provided by operating activities was primarily due to reduced operational cash flows in the current period related to timing of spending and cash receipts, offset by \$45.0 million in cash paid for the settlement of the Medtronic litigation matter in 2016. Additionally, we paid \$30.0 million in 2017 for contingent consideration related to the acquisition of Ellipse Technologies, of which \$11.2 million related to increased fair value adjustments and thus decreased cash flows from operating activities, with the remaining \$18.8 million representing the initial purchase price allocation, which flows through financing activities.

Cash Flows from Investing Activities

Cash used in investing activities was \$84.8 million for the six months ended June 30, 2017, compared to \$236.3 million used for the same period in 2016. The \$151.5 million decrease in cash used in investing activities was primarily due to the \$380.1 million cash payment (net of cash received) to fund the acquisition of Ellipse Technologies during the six months ended June 30, 2016. The funding of the Ellipse Technologies acquisition was partially offset by a net increase of \$210.4 million cash received related to activities within investment portfolios during the six months ended June 30, 2016. Additionally, there was an increase of \$16.1 million in cash used on purchases of property and equipment associated with our manufacturing initiative and general business during the six months ended June 30, 2017 as compared to the same period in 2016.

Cash Flows from Financing Activities

Cash used in financing activities was \$6.5 million for the six months ended June 30, 2017, compared to \$206.1 million cash provided for the same period in 2016. The \$212.6 million decrease in cash provided by financing activities was primarily due to the net issuance of the Senior Convertible Notes due 2021 of \$634.1 million, offset by the net \$66.3 million purchase of a call spread related to that issuance during the six months ended June 30, 2016. Additionally, we

used approximately \$343.8 million of the net proceeds to repurchase a portion of the Senior Convertible Notes due 2017 during the six months ended June 30, 2016.

Treasury stock purchases related to equity award vesting and stock option exercises totaled \$10.8 million during the six months ended June 30, 2017. We use net share settlement on stock issuances, which results in cash tax payments we make on behalf of shareowners and a decrease in the cash receipt from the issuance of common stock upon the exercising of stock options. Net share settlement is generally used in lieu of cash payments by shareowners for minimum tax withholding or exercise costs for equity awards. The net share settlement is accounted for as a treasury share repurchase transaction, with the cost of any deemed repurchased shares included in treasury stock and reported as a reduction in total equity at the time of settlement. Additionally, net share settlement for tax withholding requires us to fund a significant amount of cash for certain tax payment obligations from time-to-time with respect to the shareowner tax obligations for vested equity awards. We anticipate using cash generated from operating activities to fund such payments.

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Senior Convertible Notes

2.25% Senior Convertible Notes due 2021

In March 2016, we issued \$650.0 million principal amount of unsecured senior convertible notes with a stated interest rate of 2.25% and a maturity date of March 15, 2021, which we refer to as the 2021 Notes. The net proceeds from the offering, after deducting initial purchasers' discounts and costs directly related to the offering, were approximately \$634.1 million. Interest on the 2021 Notes began accruing upon issuance and is payable semi-annually. The 2021 Notes may be settled in cash, stock, or a combination thereof, solely at our discretion. It is our current intent and policy to settle all conversions through combination settlement, which involves satisfying the principal amount outstanding with cash and any note conversion value over the principal amount in shares of our common stock. The initial conversion rate of the 2021 Notes is 16.7158 shares per \$1,000 principal amount, which is equivalent to a conversion price of approximately \$59.82 per share, subject to adjustments. Prior to September 15, 2020, holders may convert their 2021 Notes only under the following conditions: (a) during any calendar quarter beginning June 30, 2016, if the reported sale price of our common stock for at least 20 days out of 30 consecutive trading days ending on the last trading day of the immediately preceding calendar quarter is greater than 130% of the conversion price on each applicable trading day; (b) during the five business day period in which the trading price of the 2021 Notes falls below 98% of the product of (i) the last reported sale price of our common stock and (ii) the conversion rate on that date; and (c) upon the occurrence of specified corporate events, as defined in the 2021 Notes. From September 15, 2020 and until the close of business on the second scheduled trading day immediately preceding March 15, 2021, holders may convert their 2021 Notes at any time (regardless of the foregoing circumstances). We may not redeem the 2021 Notes prior to March 20, 2019. We may redeem the 2021 Notes, at our option, in whole or in part on or after March 20, 2019 until the close of business on the business day immediately preceding September 15, 2020 if the last reported sale price of our common stock has been at least 130% of the conversion price then in effect for at least 20 trading days during any 30 consecutive trading day period ending on, and including, the trading day immediately preceding the date on which we deliver written notice of a redemption. The redemption price will be equal to 100% of the principal amount of such 2021 Notes to be redeemed plus accrued and unpaid interest to, but excluding, the redemption date. No principal payments are due on the 2021 Notes prior to maturity. Other than restrictions relating to certain fundamental changes and consolidations, mergers or asset sales and customary anti-dilution adjustments, the 2021 Notes do not contain any financial covenants and do not restrict us from paying dividends or issuing or repurchasing any of our other securities. We are unaware of any current events or market conditions that would allow holders to convert the 2021 Notes. The impact of the convertible feature will be dilutive to our earnings per share when our average stock price for the period is greater than the conversion price.

In connection with the offering of the 2021 Notes, we entered into transactions for convertible notes hedge, which we refer to as the 2021 Hedge and warrants, which we refer to as the 2021 Warrants. The 2021 Hedge was entered into with the initial purchasers and/or affiliates, which we refer to as the 2021 Counterparties, entitling us to purchase up to 10,865,270 shares of our own common stock at an initial stock price of \$59.82 per share, each of which is subject to adjustment. The cost of the 2021 Hedge was \$111.2 million. The 2021 Hedge will expire on March 15, 2021. The 2021 Hedge is expected to reduce the potential equity dilution upon conversion of the 2021 Notes if the daily volume-weighted average price per share of our common stock exceeds the strike price of the 2021 Hedge. Our assumed exercise of the 2021 Hedge is considered anti-dilutive since the effect of the inclusion would always be anti-dilutive with respect to the calculation of diluted earnings per share.

In addition, we sold the 2021 Warrants to the 2021 Counterparties to acquire up to 10,865,270 common shares of our stock. The 2021 Warrants will expire on various dates from June 2021 through December 2021 and may be settled in cash or net shares. It is our current intent and policy to settle all conversions in shares of our common stock. We received \$44.9 million in cash proceeds from the sale of the 2021 Warrants. The 2021 Warrants could have a dilutive effect on our earnings per share to the extent that the price of our common stock during a given measurement period exceeds the strike price of the 2021 Warrants, which is \$80.00 per share.

2.75% Senior Convertible Notes due 2017

On July 1, 2017, the 2017 Notes reached maturity and a majority of the holders elected to convert their outstanding notes. We paid \$64.0 million in cash for the settlement of the outstanding principal amount including accrued interest and issued 650,070 shares for settlement of the conversion value over the principal amount of the notes.

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In June 2011, we issued \$402.5 million principal amount of the 2017 Notes with a stated interest rate of 2.75% and a maturity date of July 1, 2017. The net proceeds from the offering, after deducting initial purchasers' discounts and costs directly related to the offering, were approximately \$359.2 million. The 2017 Notes may be settled in cash, stock, or a combination thereof, solely at our discretion. It is our current intent and policy to settle all conversions through combination settlement, which involves repayment of an amount of cash equal to the principal amount and any excess of the conversion value over the principal amount in shares of common stock. The initial conversion rate of the 2017 Notes is 23.7344 shares per \$1,000 principal amount, or equivalent to conversion price of approximately \$42.13 per share, which is subject to adjustment. Beginning January 1, 2017 and until the close of business on the second scheduled trading day immediately preceding July 1, 2017, holders may convert their 2017 Notes at any time. Prior to January 1, 2017, holders may convert their 2017 Notes only under the conditions as described in Note 6 to the Unaudited Consolidated Financial Statements, which includes our common stock trading at 130% of the conversion price for 20 out of 30 consecutive trading days. It is our current intent and policy to settle all conversions through combination settlement, which involves satisfying the principal amount outstanding with cash and any note conversion value over the principal amount in shares of our common stock. The impact of the convertible feature will be dilutive to our earnings per share when our stock price average for the period is greater than the conversion price. Interest on the 2017 Notes began accruing upon issuance and is payable semi-annually on January 1st and July 1st each year.

In connection with the offering of the 2017 Notes, we entered into convertible note hedge transactions, which we refer to as the 2017 Hedge, with the initial purchasers and/or their affiliates, which we refer to as the 2017 Counterparties, entitling us to purchase up to 9,553,096 shares of our common stock at an initial stock price of \$42.13 per share, each of which is subject to adjustment. The cost of the 2017 Hedge was \$80.1 million. The 2017 Hedge has an expiration date of July 1, 2017. The 2017 Hedge is expected to reduce the potential equity dilution upon conversion of the 2017 Notes if the daily volume-weighted average price per share of our common stock exceeds the strike price of the 2017 Hedge. Our assumed exercise of the 2017 Hedge is considered anti-dilutive since the effect of the inclusion would always be anti-dilutive with respect to the calculation of diluted earnings per share. On July 1, 2017, we exercised the 2017 Hedge and received 4,160,789 shares of our own common stock on a net share basis from the 2017 Counterparties.

In addition, we sold warrants, which we refer to as the 2017 Warrants, to the 2017 Counterparties to acquire up to 477,654 shares of our Series A Participating Preferred Stock, at an initial strike price of \$988.51 per share, subject to adjustment. Each share of Series A Participating Preferred Stock is initially convertible into 20 shares of our common stock, or up to 9,553,080 common shares in total. The 2017 Warrants expire on various dates from September 2017 through January 2018 and may be settled in cash or net shares. It is our current intent and policy to settle all conversions in shares of our common stock. We received \$47.9 million in cash proceeds from the sale of the 2017 Warrants. The 2017 Warrants could have a dilutive effect on our earnings per share to the extent that the price of our common stock during a given measurement period (the quarter or year-to-date period) exceeds the strike price of the 2017 Warrants, which is \$49.43 per share.

On May 24, 2017, we entered into warrant termination agreements with the 2017 Counterparties to settle the outstanding 2017 Warrants by accelerating the expiration period to varying settlement dates from June 2017 through July 2017, which terminated the existing 2017 Warrants settlement period. The settlement will be delivered in shares of our common stock, based on a fixed formula using the daily volume weighted average price as the settlement measure. As of June 30, 2017, 2017 Warrants with respect to an aggregate of 6,100,000 shares were settled on a net share basis, resulting in the issuance of 2,328,351 shares of our common stock to the 2017 Counterparties. The

remaining 2017 Warrants, with respect to an aggregate of 3,453,096 shares, will be settled on a net share basis throughout July 2017.

Revolving Senior Credit Facility

In April 2017, we entered into an Amended and Restated Credit Agreement (the “2017 Credit Agreement”) for a revolving senior credit facility (the “2017 Facility”), which replaced the previous credit agreement we had entered into in February 2016. The 2017 Credit Agreement provides for secured revolving loans, multicurrency loan options and letters of credit in an aggregate amount of up to \$500.0 million. The 2017 Credit Agreement also contains an accordion feature, which allows us to increase the aggregate principal amount of the 2017 Facility provided we remain in compliance with the underlying financial covenants, including but not limited to, compliance with the consolidated interest coverage ratio and certain consolidated leverage ratios. The 2017 Facility matures in April 2022 (subject to an earlier springing maturity date), and includes a sublimit of \$100.0 million for multicurrency borrowings, a sublimit of \$50.0 million for the issuance of standby letters of credit, and a sublimit of \$5.0 million for swingline loans. All of our assets including the assets of our material domestic subsidiaries are pledged as collateral under the 2017 Facility (subject to customary exceptions) pursuant to the term set forth in the Amended and Restated Security and Pledge Agreement (the “2017 Security Agreement”) executed in favor of the administrative agent. Each of our material domestic subsidiaries guarantees the 2017 Facility. In connection with the 2017 Facility, we incurred issuance costs which will be amortized over the term of the 2017 Facility. As of June 30, 2017, we had \$20.0 million outstanding under the 2017 Facility, at an interest rate of 2.97% (one month LIBOR plus 1.75%).

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Borrowings under the 2017 Facility bear interest, at our option, at a rate equal to an applicable margin plus: (a) the applicable Eurocurrency Rate (as defined in the 2017 Credit Agreement), or (b) a base rate determined by reference to the highest of (1) the federal funds effective rate plus 0.50%, (2) the Bank of America prime rate, and (3) LIBOR for an interest period of one month plus 1.00%. The margin for the 2017 Facility ranges, based on our consolidated leverage ratio, from 0.00% to 1.00% in the case of base rate loans and from 1.00% to 2.00% in the case of Eurocurrency Rate loans. The 2017 Facility includes an unused line fee ranging, based on our consolidated leverage ratio, from 0.20% to 0.35% per annum on the revolving commitment.

The 2017 Credit Agreement contains affirmative, negative, permitted acquisition and financial covenants, and events of default customary for financings of this type. The financial covenants require us to maintain ratios of consolidated earnings before interest, taxes, depreciation and amortization (EBITDA) in relation to consolidated interest expense and consolidated debt, respectively, as defined in the 2017 Credit Agreement. The 2017 Facility grants the lenders preferred first priority liens and security interests in capital stock, intercompany debt and all of our present and future property and assets including each guarantor. We are currently in compliance with the Credit Agreement covenants.

Critical Accounting Policies

Our discussion and analysis of our financial condition and results of operations is based upon our Unaudited Consolidated Financial Statements, which have been prepared in accordance with generally accepted accounting principles in the United States, or GAAP. The preparation of these financial statements requires us to make estimates and judgments that affect the reported amounts of assets, liabilities, revenues and expenses. On an ongoing basis, we evaluate our estimates including those related to bad debts, inventories, valuation of goodwill, intangibles, other long-term assets, stock-based compensation, income taxes, and legal proceedings. We base our estimates on historical experience and on various other assumptions we believe to be reasonable under the circumstances, the results of which form the basis for making judgments about the carrying values of assets and liabilities not readily apparent from other sources. Actual results may differ from these estimates. Our critical accounting policies and estimates are discussed in our Annual Report on Form 10-K for the fiscal year ended December 31, 2016 and there have been no material changes during the six months ended June 30, 2017.

Off-Balance Sheet Arrangements

As of June 30, 2017, we did not have any off-balance sheet arrangements.

Contractual Obligations and Commitments

As of June 30, 2017, there were no material changes, excluding the aforementioned lease commitment, outside of the ordinary course of business, in our outstanding contractual obligations from those disclosed within “Management’s Discussion and Analysis of Financial Condition and Results of Operations” in our Annual Report on Form 10-K for the fiscal year ended December 31, 2016.

Item 3. Quantitative and Qualitative Disclosures About Market Risk

As of June 30, 2017, there has been no material change in our assessment of our sensitivity to market risk since our presentation set forth in Item 7A, “Quantitative and Qualitative Disclosures About Market Risk”, in our Annual Report on Form 10-K for the fiscal year ended December 31, 2016.

Item 4. Controls and Procedures

Disclosure Controls and Procedures

We maintain disclosure controls and procedures that are designed to ensure that information required to be disclosed in our reports under the Securities Exchange Act of 1934, as amended, or the Exchange Act, is recorded, processed, summarized and reported within the time lines specified in the SEC's rules and forms, and that such information is accumulated and communicated to our management, including our Chief Executive Officer and Chief Financial Officer, as appropriate, to allow timely decisions regarding required disclosure. In designing and evaluating the disclosure controls and procedures, management recognized that any controls and procedures, no matter how well designed and operated, can only provide reasonable assurance of achieving the desired control objectives, and in reaching a reasonable level of assurance, management necessarily was required to apply its judgment in evaluating the cost-benefit relationship of possible controls and procedures.

Under the supervision and with the participation of our management, including our Chief Executive Officer and our Chief Financial Officer, we carried out an evaluation of the effectiveness of the Company's disclosure controls and procedures (as defined in SEC Rules 13a - 15(e) and 15d - 15(e)) as of June 30, 2017. Based on such evaluation, our management has concluded that as of June 30, 2017, the Company's disclosure controls and procedures are effective.

Changes in Internal Control Over Financial Reporting

Under the supervision and with the participation of our management, including our Chief Executive Officer and our Chief Financial Officer, we carried out an evaluation of any potential changes in our internal control over financial reporting during the fiscal quarter covered by this Quarterly Report.

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There has been no change to our internal control over financial reporting during our most recent fiscal quarter that our certifying officers concluded materially affected, or is reasonably likely to materially affect, our internal control over financial reporting.

PART II. OTHER INFORMATION

Item 1. Legal Proceedings

For a description of our material pending legal proceedings, refer to Note 11 “Contingencies” of the Notes to Unaudited Consolidated Financial Statements included in Part I, Item 1 of this Quarterly Report, which is incorporated herein by reference.

Item 1A. Risk Factors

The risk factors set forth below contain material changes to the risk factors previously disclosed and included in our Annual Report on Form 10-K for the fiscal year ended December 31, 2016. An investment in our common stock involves a high degree of risk. You should consider carefully the risks and uncertainties described under Item 1A of Part I of our Annual Report on Form 10-K, as updated in this Item 1A (collectively the “Risk Factors”) together with all other information contained or incorporated by reference in this report before you decide to invest in our common stock. If any of the Risk Factors were to actually occur, our business, financial condition, results of operations and our future growth prospects could be materially and adversely affected. Under the circumstances, the trading price of our common stock could decline, and you may lose all or part of your investment.

Risks Related to Regulatory and Compliance

We are subject to federal, state and foreign fraud and abuse laws and health information privacy and security laws, which, if violated, could subject us to substantial penalties.

There are numerous U.S. federal and state, as well as foreign, laws pertaining to healthcare fraud and abuse, including anti-kickback, false claims and physician transparency laws. Our relationships with physicians, providers and hospitals are subject to scrutiny under these laws. We may also be subject to patient privacy regulation by both the federal government and the states and foreign jurisdictions in which we conduct our business.

Healthcare fraud and abuse laws are broad in scope and are subject to evolving interpretation, which could require us to incur substantial costs to monitor compliance or to alter our practices if they are found not to be in compliance. Violations of these laws may be punishable by criminal or civil sanctions, including substantial fines, imprisonment and exclusion from participation in governmental healthcare programs. Despite implementation of a comprehensive global healthcare compliance program, we cannot provide assurance that any of the healthcare fraud and abuse laws will not change or be interpreted in the future in a manner which restricts or adversely affects our business activities or relationships with healthcare professionals, nor can we make any assurances that authorities will not challenge or investigate our current or future activities under these laws.

In July 2015, we entered into a settlement agreement with the U.S. Department of Justice, or DOJ, pursuant to which we paid \$13.5 million to resolve an investigation into possible false or otherwise improper claims submitted to Medicare and Medicaid. We admitted no wrongdoing as part of the settlement. In August 2015, we received a civil investigative demand, or CID, issued by the DOJ pursuant to the federal False Claims Act. The CID requires the delivery of a wide range of documents and information related to an investigation by the DOJ concerning allegations that we assisted a physician group customer in submitting improper claims for reimbursement and made improper payments to the physician group in violation of the Anti-Kickback Statute. We are cooperating with the DOJ. Additionally, on June 9, 2017, we received a subpoena from the Office of the Inspector General of the U.S. Department of Health and Human Services, or OIG, in connection with an investigation into possible false or otherwise improper claims submitted to Medicare and Medicaid. We are working with the OIG to understand the scope of the subpoena and its request for documents, and we intend to fully cooperate with the OIG's request.

No assurance can be given as to the timing or outcome of these investigations. Responding to government requests and investigations requires considerable resources, including the time and attention of management. If we were to become the subject of an enforcement action, including any action resulting from the investigation by the DOJ or the OIG, it could result in negative publicity, penalties, fines, the exclusion of our products from reimbursement under federally-funded programs and/or prohibitions on our ability to sell our products, which could have a material adverse effect on our results of operations, financial condition and liquidity.

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Risks Related to Our Financial Results and Need for Financing

If we fail to comply with the covenants and other obligations under our credit facility, the lenders may be able to accelerate amounts owed under the facility and may foreclose upon the assets securing our obligations.

In April 2017, we entered into an Amended and Restated Credit Agreement (the “2017 Credit Agreement”) that provides for a revolving senior credit facility (the “2017 Facility”), which replaced the previous Credit Agreement we had entered into in February 2016. The 2017 Credit Agreement provides for secured revolving loans, multicurrency loan options and letters of credit in an aggregate amount of up to \$500.0 million. The 2017 Credit Agreement also contains an expansion feature, which allows us to increase the aggregate principal amount of the 2017 Facility provided we remain in compliance with the underlying financial covenants, including but not limited to, compliance with the consolidated interest coverage ratio and certain consolidated leverage ratios. All of our assets and the assets of our material subsidiaries are pledged as collateral under the 2017 Facility (subject to customary exceptions) and each of our material domestic subsidiaries guarantee the 2017 Facility. The covenants set forth in the 2017 Credit Agreement restrict, among other things, our ability to: create liens on assets, incur additional indebtedness, make investments, make acquisitions and other fundamental changes, sell and dispose of property or assets, pay dividends and other distributions, change the business conducted, engage in certain transactions with affiliates, enter into burdensome agreements, limit certain use of proceeds, amend organizational documents, change accounting policies or reporting practices, modify or terminate documents related to certain indebtedness, enter into sale and leaseback transactions, fund any person or business that is the subject of sanctions, and use proceeds for any breach of anti-corruption laws. If we fail to comply with the covenants and our other obligations under the 2017 Facility, the lenders would be able to accelerate the required repayment of amounts due under the 2017 Credit Agreement and, if they are not repaid, could foreclose upon our assets securing our obligations under the 2017 Facility.

Item 2. Unregistered Sales of Equity Securities and Use of Proceeds

On May 24, 2017, we entered into warrant termination agreements (the “Termination Agreements”) with each of Goldman, Sachs & Co. LLC (f/k/a Goldman, Sachs & Co.) (“Goldman”) and Bank of America, N.A., an affiliate of Merrill Lynch, Pierce, Fenner & Smith Incorporated (“Bank of America”) to terminate the outstanding 2017 Warrants that were issued to Goldman and Bank of America pursuant to the letter agreements between us and each of Goldman and Bank of America, dated as of June 22, 2011 and June 24, 2011. Pursuant to the terms of the Termination Agreements, 2017 Warrants with respect to an aggregate of 9,553,096 shares of our common stock will be terminated. In consideration of the termination of the 2017 Warrants, we will deliver to Goldman and Bank of America shares of our common stock, which are being issued in reliance on the exemption from registration provided by Section 4(a)(2) of the Securities Act of 1933, as amended. As of June 30, 2017, 2017 Warrants with respect to an aggregate of 6,100,000 shares were settled on a net share basis, resulting in the issuance of 991,675 shares to Goldman and 1,336,676 shares to Bank of America. The remaining 2017 Warrants, with respect to an aggregate of 3,453,096 shares, will be settled on a net share basis throughout July 2017. The 2017 Warrants were sold by us to Goldman and Bank of America as part of the 2017 Notes issuance in which proceeds from that issuance were used to finance the Company for general purposes.

In connection with the issuance of the 2017 Notes, we entered into letter agreements with respect to the 2017 Hedge with Goldman and Bank of America, dated as of June 22, 2011 and June 24, 2011. The 2017 Hedge entitled us to purchase up to 9,553,096 shares of our common stock at an initial stock price of \$42.13 per share. On July 1, 2017, we exercised the 2017 Hedge and received 4,160,789 shares of our common stock from Goldman and Bank of America.

Item 3. Defaults Upon Senior Securities

None.

Item 4. Mine Safety Disclosures

None.

Item 5. Other Information

On July 26, 2017, we amended and restated the NuVasive, Inc. Executive Severance Plan (the “Executive Severance Plan”). The Executive Severance Plan provides eligible executives with severance benefits upon an involuntary termination of employment by the Company that is not for cause (as defined in the Executive Severance Plan). The primary purpose of the amendment and restatement of the Executive Severance Plan is to (i) provide that severance pay and benefits shall be subject to and conditioned upon continued compliance with post-termination restrictive covenants, clawback provisions, and a general release of claims and (ii) modify the payment terms for severance pay such that any severance pay shall be paid in installments rather than as a lump sum. The foregoing description of the Executive Severance Plan, as amended and restated, is qualified in its entirety by reference to the full text of the Amended and Restated Severance Plan, which is filed as Exhibit 10.3 to this Quarterly Report on Form 10-Q and is incorporated by reference herein.

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Item 6. Exhibits

Exhibit	Number Description
3.1	Restated Certificate of Incorporation (incorporated by reference to our Quarterly Report on Form 10-Q filed with the SEC on August 13, 2004)
3.2	Certificate of Amendment to the Restated Certificate of Incorporation (incorporated by reference to our Current Report on Form 8-K filed with the SEC on September 28, 2011)
3.3	Restated Bylaws (incorporated by reference to our Current Report on Form 8-K filed with the SEC on January 6, 2012)
3.4	Amendment No. 1 to the Restated Bylaws (incorporated by reference to our Current Report on Form 8-K filed with the SEC on May 19, 2014)
3.5	Amendment No. 2 to the Restated Bylaws (incorporated by reference to our Current Report on Form 8-K filed with the SEC on August 1, 2016)
10.1	Warrant Termination Agreement, dated as of May 24, 2017, between the Company and Bank of America, N.A.
10.2	Warrant Termination Agreement, dated as of May 24, 2017, between the Company and Goldman Sachs & Co. LLC
10.3#	NuVasive, Inc. Amended and Restated Executive Severance Plan
10.4	Amended and Restated Credit Agreement, dated April 25, 2017, by and among the Company, as the Borrower, certain material subsidiaries of the Company, as guarantors, Bank of America, N.A. and each of those additional Lenders party thereto (incorporated by reference to our Current Report on Form 8-K filed with the SEC on April 25, 2017)
10.5	Amended and Restated Security and Pledge Agreement, dated April 25, 2017, by and among the Company and certain material subsidiaries of the Company in favor of Bank of America, N.A. (incorporated by reference to our Current Report on Form 8-K filed with the SEC on April 25, 2017)
31.1*	Certification of the Chief Executive Officer pursuant to Rule 13a-14(b) of the Securities Exchange Act of 1934, as amended, and 18 U.S.C. section 1350
31.2*	Certification of the Chief Financial Officer pursuant to Rule 13a-14(b) of the Securities Exchange Act of 1934, as amended, and 18 U.S.C. section 1350
32.1*	Certifications of Chief Executive Officer and Chief Financial Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002
101.INS	XBRL Instance Document
101.SCH	XBRL Taxonomy Extension Schema
101.CAL	XBRL Taxonomy Extension Calculation Linkbase
101.DEF	XBRL Taxonomy Extension Definition Linkbase
101.LAB	XBRL Taxonomy Extension Label Linkbase
101.PRE	XBRL Taxonomy Extension Presentation Linkbase
#	Indicates management contract or compensatory plan.
*	

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These certifications are being furnished solely to accompany this annual report pursuant to 18 U.S.C. Section 1350, and are not being filed for purposes of Section 18 of the Securities Exchange Act of 1934 and are not to be incorporated by reference into any filing of NuVasive, Inc., whether made before or after the date hereof, regardless of any general incorporation language in such filing.

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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, thereunto duly authorized.

NUVASIVE, INC.

Date: July 27, 2017 By: /s/ Gregory T. Lucier
Gregory T. Lucier
Chairman and Chief Executive Officer
(Principal Executive Officer)

Date: July 27, 2017 By: /s/ Quentin S. Blackford
Quentin S. Blackford
Executive Vice President and Chief Financial Officer, Head of Strategy and Corporate
Integrity
(Principal Financial Officer)

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#	Indicates management contract or compensatory plan.
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