

TESARO, Inc.  
Form 10-Q  
May 03, 2018  
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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 March 31, 2018

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 001-35587

TESARO, INC.

(Exact Name of Registrant as Specified in Its Charter)

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|                                                                               |                                                       |
|-------------------------------------------------------------------------------|-------------------------------------------------------|
| Delaware<br>(State or Other Jurisdiction of<br>Incorporation or Organization) | 27-2249687<br>(I.R.S. Employer<br>Identification No.) |
|-------------------------------------------------------------------------------|-------------------------------------------------------|

|                                                                                          |                     |
|------------------------------------------------------------------------------------------|---------------------|
| 1000 Winter Street<br>Waltham, Massachusetts<br>(Address of Principal Executive Offices) | 02451<br>(Zip Code) |
|------------------------------------------------------------------------------------------|---------------------|

(339) 970-0900

(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 that 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 Web site, if any, every Interactive Data File required to be submitted and posted pursuant to Rule 405 of Regulation S-T 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, a 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 smaller reporting company)

Smaller 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.

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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 April 30, 2018, there were 54,823,563 shares of the registrant's Common Stock, par value \$0.0001 per share, outstanding.

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TESARO, INC.

FORM 10-Q

FOR THE THREE MONTHS ENDED MARCH 31, 2018

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CERTIFICATIONS

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## PART I FINANCIAL INFORMATION

## Item 1. Financial Statements.

## TESARO, INC.

## Condensed Consolidated Balance Sheets

(all amounts in 000's, except share and per share data)

(Unaudited)

|                                      | December 31,<br>2017 | March 31,<br>2018 |
|--------------------------------------|----------------------|-------------------|
| Assets                               |                      |                   |
| Current assets:                      |                      |                   |
| Cash and cash equivalents            | \$ 643,095           | \$ 498,980        |
| Accounts receivable                  | 53,416               | 44,182            |
| Inventories                          | 57,939               | 74,444            |
| Other current assets                 | 33,511               | 38,184            |
| Total current assets                 | 787,961              | 655,790           |
| Intangible assets, net               | 56,384               | 54,947            |
| Property and equipment, net          | 9,652                | 10,272            |
| Restricted cash                      | 2,552                | 2,556             |
| Other assets                         | 5,636                | 6,127             |
| Total assets                         | \$ 862,185           | \$ 729,692        |
| Liabilities and stockholders' equity |                      |                   |
| Current liabilities:                 |                      |                   |
| Accounts payable                     | \$ 4,172             | \$ 5,079          |
| Accrued expenses                     | 154,808              | 150,589           |
| Deferred revenue, current            | 324                  | 117               |
| Other current liabilities            | 6,902                | 7,451             |
| Total current liabilities            | 166,206              | 163,236           |
| Convertible notes, net               | 143,446              | 146,529           |
| Long-term debt, net                  | 293,659              | 293,888           |
| Deferred revenue, non-current        | 211                  | 188               |
| Other non-current liabilities        | 9,577                | 8,123             |
| Total liabilities                    | 613,099              | 611,964           |

## Commitments and contingencies (Note 10)

## Stockholders' equity:

|                                                                                                                                                                                                                             |             |             |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------|-------------|
| Preferred stock, \$0.0001 par value; 10,000,000 shares authorized at both December 31, 2017 and March 31, 2018; no shares issued or outstanding at both December 31, 2017 and March 31, 2018                                | —           | —           |
| Common stock, \$0.0001 par value; 100,000,000 shares authorized at both December 31, 2017 and March 31, 2018; 54,464,039 and 54,801,636 shares issued and outstanding at December 31, 2017 and March 31, 2018, respectively | 5           | 5           |
| Additional paid-in capital                                                                                                                                                                                                  | 1,724,850   | 1,755,783   |
| Accumulated other comprehensive loss                                                                                                                                                                                        | (5,882)     | (5,357)     |
| Accumulated deficit                                                                                                                                                                                                         | (1,469,887) | (1,632,703) |
| Total stockholders' equity                                                                                                                                                                                                  | 249,086     | 117,728     |
| Total liabilities and stockholders' equity                                                                                                                                                                                  | \$ 862,185  | \$ 729,692  |

See accompanying notes to condensed consolidated financial statements.

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TESARO, INC.

Condensed Consolidated Statements of Operations and Comprehensive Loss

(all amounts in 000's, except per share data)

(Unaudited)

|                                                                                                                           | Three Months Ended<br>March 31, |              |
|---------------------------------------------------------------------------------------------------------------------------|---------------------------------|--------------|
|                                                                                                                           | 2017                            | 2018         |
| Revenues:                                                                                                                 |                                 |              |
| Product revenue, net                                                                                                      | \$ 2,139                        | \$ 50,172    |
| License, collaboration and other revenues                                                                                 | 934                             | (430)        |
| Total revenues                                                                                                            | 3,073                           | 49,742       |
| Expenses:                                                                                                                 |                                 |              |
| Cost of sales – product                                                                                                   | 444                             | 9,997        |
| Cost of sales – intangible asset amortization                                                                             | 490                             | 1,437        |
| Research and development                                                                                                  | 66,122                          | 96,755       |
| Selling, general and administrative                                                                                       | 69,262                          | 93,607       |
| Acquired in-process research and development                                                                              | —                               | —            |
| Total expenses                                                                                                            | 136,318                         | 201,796      |
| Loss from operations                                                                                                      | (133,245)                       | (152,054)    |
| Interest expense                                                                                                          | (4,267)                         | (12,092)     |
| Interest income                                                                                                           | 841                             | 1,665        |
| Other income                                                                                                              | —                               | 81           |
| Loss before income taxes                                                                                                  | (136,671)                       | (162,400)    |
| Provision for income taxes                                                                                                | 54                              | 416          |
| Net loss                                                                                                                  | \$ (136,725)                    | \$ (162,816) |
| Net loss per share applicable to common stockholders - basic and diluted                                                  | \$ (2.55)                       | \$ (2.98)    |
| Weighted-average number of common shares used in net loss per share applicable to common stockholders - basic and diluted | 53,685                          | 54,615       |
| Comprehensive income:                                                                                                     |                                 |              |
| Net loss                                                                                                                  | \$ (136,725)                    | \$ (162,816) |
| Other comprehensive income:                                                                                               |                                 |              |
| Unrealized gain on pension obligation                                                                                     | 45                              | 63           |
| Foreign currency translation adjustments                                                                                  | 35                              | 462          |
| Other comprehensive income                                                                                                | 80                              | 525          |
| Comprehensive loss                                                                                                        | \$ (136,645)                    | \$ (162,291) |

See accompanying notes to condensed consolidated financial statements.

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TESARO, INC.

Condensed Consolidated Statements of Cash Flows

(all amounts in 000's)

(Unaudited)

|                                                                                | Three Months Ended<br>March 31, |              |
|--------------------------------------------------------------------------------|---------------------------------|--------------|
|                                                                                | 2017                            | 2018         |
|                                                                                | (as revised)                    |              |
| Operating activities                                                           |                                 |              |
| Net loss                                                                       | \$ (136,725)                    | \$ (162,816) |
| Adjustments to reconcile net loss to net cash used in operating activities:    |                                 |              |
| Depreciation and amortization expense                                          | 954                             | 2,455        |
| Stock-based compensation expense                                               | 18,401                          | 26,128       |
| Non-cash interest expense                                                      | 2,757                           | 3,312        |
| Changes in operating assets and liabilities:                                   |                                 |              |
| Accounts receivable                                                            | 134                             | 9,287        |
| Inventories                                                                    | (179)                           | (16,200)     |
| Other assets                                                                   | (3,345)                         | (4,731)      |
| Accounts payable                                                               | 5,119                           | 895          |
| Accrued expenses                                                               | (1,847)                         | (4,834)      |
| Deferred revenues                                                              | (22)                            | (230)        |
| Other liabilities                                                              | 172                             | (898)        |
| Net cash used in operating activities                                          | (114,581)                       | (147,632)    |
| Investing activities                                                           |                                 |              |
| Purchase of property and equipment                                             | (1,790)                         | (1,619)      |
| Net cash used in investing activities                                          | (1,790)                         | (1,619)      |
| Financing activities                                                           |                                 |              |
| Proceeds from sale of common stock, net of issuance costs                      | (8)                             | —            |
| Proceeds from exercise of stock options and Employee Stock Purchase Plan       | 3,323                           | 4,497        |
| Net cash provided by financing activities                                      | 3,315                           | 4,497        |
| Effect of exchange rate changes on cash, cash equivalents, and restricted cash | 45                              | 651          |
| Decrease in cash, cash equivalents, and restricted cash                        | (113,011)                       | (144,103)    |
| Cash, cash equivalents, and restricted cash at beginning of period             | 787,866                         | 645,954      |
| Cash, cash equivalents, and restricted cash at end of period                   | \$ 674,855                      | \$ 501,851   |
| Non-cash investing and financing activities                                    |                                 |              |
| Stock option exercise proceeds receivable as of period end                     | \$ 103                          | \$ —         |
| Leasehold improvement assets funded by lessor                                  | \$ 585                          | \$ —         |
| Purchase of property and equipment - cash not paid as of period end            | \$ 301                          | \$ 56        |

Supplemental disclosure of cash flow information

|                                                |           |           |
|------------------------------------------------|-----------|-----------|
| Cash paid for interest                         | \$ 3,019  | \$ 12,274 |
| Milestone obligation not paid as of period end | \$ 24,790 | \$ —      |
| Income taxes paid                              | \$ 176    | \$ 80     |

The following table presents the line items and amounts of cash, cash equivalents and restricted cash reported within the condensed consolidated balance sheets:

|                                                  | December<br>31,<br>2017 | March 31,<br>2018 |
|--------------------------------------------------|-------------------------|-------------------|
| Cash and cash equivalents                        | \$ 643,095              | \$ 498,980        |
| Restricted cash included in other current assets | 307                     | 315               |
| Restricted cash, noncurrent                      | 2,552                   | 2,556             |
| Total cash, cash equivalents and restricted cash | \$ 645,954              | \$ 501,851        |

See accompanying notes to condensed consolidated financial statements.

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TESARO, INC.

Notes to Condensed Consolidated Financial Statements

(Unaudited)

1. Description of Business

TESARO, Inc., or the Company or TESARO, was incorporated in Delaware and commenced operations in 2010. Headquartered in Waltham, Massachusetts, TESARO is a commercial-stage biopharmaceutical company devoted to providing transformative therapies to people bravely facing cancer. TESARO's primary focus is to develop treatments for solid tumors using various approaches, including small molecules and immuno-oncology antibodies, as monotherapies and in combinations. The Company has in-licensed and is developing several oncology-related product candidates, and has entered into several research collaborations with third parties for the discovery of new candidates. The Company operates in one segment. The Company is subject to a number of risks, including dependence on key individuals, regulatory and manufacturing risks, the need to develop additional commercially viable products, risks associated with competitors, many of which are larger and better capitalized, risks related to intellectual property, and the need to obtain adequate additional financing to fund the development and potential commercialization of its product candidates and further its in-licensing and acquisition activities.

The Company's two currently marketed products, ZEJULA® and VARUBI®/VARUBY®, are approved in both the U.S. and the European Union, or EU. ZEJULA is approved as a maintenance treatment of adults with recurrent epithelial ovarian, fallopian tube, or primary peritoneal cancer who are in a complete or partial response to platinum-based chemotherapy. VARUBI/VARUBY is approved for use in combination with other antiemetic agents in adults for the prevention of delayed nausea and vomiting associated with initial and repeat courses of emetogenic cancer chemotherapy, including, but not limited to, highly emetogenic chemotherapy. The Company has determined that it will cease marketing and distribution of the intravenous, or IV, formulation of VARUBI and intends to pursue strategic alternatives for the VARUBI brand.

The Company has incurred significant operating losses since inception and has relied on its ability to fund its operations through private and public equity and debt financings and to a lesser extent through product sales and license and collaboration arrangements. Management expects operating losses and negative operating cash flows to continue for the foreseeable future. As the Company continues to incur losses, the transition to profitability is dependent upon the successful development, approval, and commercialization of its products and product candidates and the achievement of a level of revenues adequate to support its cost structure. The Company believes that its currently available funds, including an additional draw under the Company's term loan agreement, in addition to cash generated from sales of its products will be sufficient to fund the Company's operations through at least the next 12 months from the issuance of this Quarterly Report on Form 10-Q. Management's belief with respect to its ability to fund operations is based on estimates that are subject to risks and uncertainties. If actual results are different from

management's estimates, the Company may need to seek additional funding.

## 2. Basis of Presentation and Significant Accounting Policies

### Basis of Presentation

The accompanying condensed consolidated financial statements are unaudited and have been prepared by TESARO in conformity with accounting principles generally accepted in the United States of America, or GAAP.

The Company's condensed consolidated financial statements reflect the operations of the Company and its wholly-owned subsidiaries. All significant intercompany balances and transactions have been eliminated in consolidation. The Company currently operates in one business segment, which is the identification, acquisition, development and commercialization of oncology-related therapeutics, and has a single reporting and operating unit structure.

Certain information and footnote disclosures normally included in the Company's annual financial statements have been condensed or omitted. These interim financial statements, in the opinion of management, reflect all normal recurring adjustments necessary for a fair presentation of the Company's financial position and results of operations for the interim periods ended March 31, 2017 and 2018.

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The results of operations for the interim periods are not necessarily indicative of the results of operations to be expected for the full fiscal year. These interim financial statements should be read in conjunction with the audited financial statements as of and for the year ended December 31, 2017 and the notes thereto, which are included in the Company's Annual Report on Form 10-K for the year ended December 31, 2017. The significant accounting policies used in preparation of these condensed consolidated financial statements for the three months ended March 31, 2018 are consistent with those discussed in Note 2 to the consolidated financial statements in the Company's 2017 Annual Report on Form 10-K and are updated below as necessary.

### Use of Estimates

The preparation of condensed consolidated financial statements in conformity with GAAP requires management to make certain estimates and assumptions that affect the reported amounts of assets, liabilities, revenues and expenses, other comprehensive income (loss) and the related disclosures. Significant estimates in these condensed consolidated financial statements include estimates made in connection with accrued research and development expenses, stock-based compensation expense, revenue, valuation of convertible notes, inventory, intangible assets and related amortization. The Company bases its estimates on historical experience, known trends and other market-specific or other relevant factors that it believes to be reasonable under the circumstances. Actual results may differ from those estimates or assumptions.

### Fair Value of Financial Instruments

The Company is required to disclose information on all assets and liabilities reported at fair value that enables an assessment of the inputs used in determining the reported fair values. The fair value hierarchy prioritizes valuation inputs based on the observable nature of those inputs. The fair value hierarchy applies only to the valuation inputs used in determining the reported fair value of the investments and is not a measure of investment credit quality. The hierarchy defines three levels of valuation inputs:

Level 1 inputs      Quoted prices in active markets for identical assets or liabilities

Level 2 inputs      Observable inputs other than Level 1 inputs, including quoted prices in active markets for similar assets or liabilities and quoted prices for identical assets or liabilities in markets that are not active

Level 3 inputs    Unobservable inputs that reflect the Company's own assumptions about the assumptions market participants would use in pricing the asset or liability

The following table presents information about the Company's financial assets and liabilities that have been measured at fair value as of December 31, 2017 and March 31, 2018 and indicates the fair value hierarchy of the valuation inputs utilized to determine such fair value (in thousands):

| Description        | Balance Sheet Classification | December 31, 2017 |            |         |         |
|--------------------|------------------------------|-------------------|------------|---------|---------|
|                    |                              | Total             | Level 1    | Level 2 | Level 3 |
| Assets:            |                              |                   |            |         |         |
| Money market funds | Cash and cash equivalents    | \$ 593,955        | \$ 593,955 | \$ —    | \$ —    |
| Total assets       |                              | \$ 593,955        | \$ 593,955 | \$ —    | \$ —    |

| Description        | Balance Sheet Classification | March 31, 2018 |            |         |         |
|--------------------|------------------------------|----------------|------------|---------|---------|
|                    |                              | Total          | Level 1    | Level 2 | Level 3 |
| Assets:            |                              |                |            |         |         |
| Money market funds | Cash and cash equivalents    | \$ 403,121     | \$ 403,121 | \$ —    | \$ —    |
| Total assets       |                              | \$ 403,121     | \$ 403,121 | \$ —    | \$ —    |

The carrying amounts of accounts payable and accrued expenses approximate their fair values due to their short-term maturities.

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In September 2014, the Company issued \$201.3 million aggregate principal amount of 3.00% convertible senior notes due October 1, 2021, or the Convertible Notes. Interest is payable semi-annually in arrears on April 1 and October 1 of each year. As of March 31, 2018, the carrying value of the Convertible Notes, net of unamortized discount and debt issuance costs, was \$146.5 million and the estimated fair value of the principal amount was \$369.0 million. As of March 31, 2018, the carrying value of the Company's borrowing under its term loan agreement approximated its fair value. The Convertible Notes and the term loan agreement are discussed in more detail in Note 5, "Debt".

## Revenue Recognition

The Company recognizes revenue when its customer obtains control of promised goods or services, in an amount that reflects the consideration which the Company expects to receive in exchange for those goods or services. To determine revenue recognition, the Company performs the following five steps: (i) identify the contract(s) with a customer; (ii) identify the performance obligations in the contract; (iii) determine the transaction price; (iv) allocate the transaction price to the performance obligations in the contract; and (v) recognize revenue when (or as) the Company satisfies a performance obligation. The Company only applies the five-step model to contracts when it is probable that the Company will collect the consideration it is entitled to in exchange for the goods or services it transfers to the customer. For a further discussion of accounting for net product revenue and license, collaboration and other revenues, see Note 11, "Revenue Recognition".

## Intangible Assets

The Company maintains definite-lived intangible assets related to milestone payments made to third parties subsequent to regulatory approval for acquired and in-licensed product candidates. These assets are amortized over their remaining useful lives, which are generally estimated to be the remaining patent life. If the Company's estimate of the product's useful life is shorter than the remaining patent life, then the shorter period is used. Intangible assets are amortized using the economic consumption method if anticipated future revenues can be reasonably estimated. The straight-line method is used when future revenues cannot be reasonably estimated, with a cumulative catch-up of amortization expense for milestone payments that do not result in additional intellectual property rights and/or incremental cash flows. Amortization expense is recorded as a component of cost of sales in the condensed consolidated statements of operations and comprehensive loss.

The Company assesses its intangible assets for impairment if indicators are present or changes in circumstance suggest that impairment may exist. Events that could result in an impairment, or trigger an interim impairment assessment, include the receipt of additional clinical or nonclinical data regarding one of the Company's drug candidates or a potentially competitive drug candidate, changes in the clinical development program for a drug candidate or new information regarding potential sales for the drug. If impairment indicators are present or changes in circumstance suggest that impairment may exist, the Company performs a recoverability test by comparing the sum of

the estimated undiscounted cash flows of each intangible asset to its carrying value on the condensed consolidated balance sheet. If the undiscounted cash flows used in the recoverability test are less than the carrying value, the Company would determine the fair value of the intangible asset and recognize an impairment loss if the carrying value of the intangible asset exceeds its fair value.

#### Comprehensive Loss

Comprehensive loss is defined as the change in equity of a business enterprise during a period from transactions and other events and circumstances from non-owner sources. Comprehensive loss consists of net loss and other comprehensive loss. Other comprehensive loss includes foreign currency translation adjustments and unrealized gains

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and losses on pension obligations. The following table presents changes in the components of accumulated other comprehensive loss (in thousands):

|                                      | Foreign currency translation<br>adjustments | Unrealized loss on pension<br>liability | Total      |
|--------------------------------------|---------------------------------------------|-----------------------------------------|------------|
| Balance at December 31, 2016         | \$ (142)                                    | \$ (2,782)                              | \$ (2,924) |
| Other comprehensive (loss)<br>income | 35                                          | 45                                      | 80         |
| Balance at March 31, 2017            | \$ (107)                                    | \$ (2,737)                              | \$ (2,844) |
| Balance at December 31, 2017         | \$ 259                                      | \$ (6,141)                              | \$ (5,882) |
| Other comprehensive (loss)<br>income | 462                                         | 63                                      | 525        |
| Balance at March 31, 2018            | \$ 721                                      | \$ (6,078)                              | \$ (5,357) |

## New Accounting Pronouncements - Recently Adopted

In August 2016, the Financial Accounting Standards Board, or FASB, issued Accounting Standards Update, or ASU, No. 2016-15, which is intended to simplify and clarify how certain transactions are classified in the statement of cash flows, and to reduce diversity in practice for such transactions. This ASU addresses eight specific issues regarding classification of cash flows. The standard is effective for financial statements issued for fiscal years beginning after December 15, 2017, and interim periods within those fiscal years. The Company adopted this ASU effective January 1, 2018. The adoption of this guidance did not have an impact on the Company's condensed consolidated financial statements and related disclosures.

In October 2016, the FASB issued ASU No. 2016-16, which removes the prohibition in ASC Topic 740 against the immediate recognition of the current and deferred income tax effects of intra-entity transfers of assets other than inventory. As a result, the income tax consequences from the intra-entity transfer of an asset, other than inventory, and associated changes to deferred taxes will be recognized when the transfer occurs. The Company adopted this new standard effective January 1, 2018 using the modified retrospective method. Upon adoption, the Company recorded a deferred tax asset and corresponding valuation allowance of \$52.7 million. There was no cumulative effect adjustment to accumulated deficit as of the beginning of the period of adoption.

In November 2016, the FASB issued ASU No. 2016-18, which requires amounts generally described as restricted cash and restricted cash equivalents to be included with cash and cash equivalents when reconciling the total beginning and ending amounts for the periods shown on the statement of cash flows. ASU No. 2016-18 is effective for fiscal years beginning after December 15, 2017 (including interim periods within those periods) using a retrospective transition method to each period presented. The Company adopted this ASU effective January 1, 2018. The adoption of this guidance required the following changes and disclosures to the presentation of the condensed consolidated financial

statements:

- Cash, cash equivalents and restricted cash and cash equivalents reported on the condensed consolidated statements of cash flows now includes restricted cash and cash equivalents and totals \$646.0 million and \$501.9 million as of December 31, 2017 and March 31, 2018, respectively.
- Restricted cash generally consists of cash balances held as collateral for the Company's employee credit card programs.

In May 2017, the FASB issued ASU No. 2017-09, which clarifies when a change to the terms or conditions of a share-based payment award must be accounted for as a modification. The new guidance requires modification accounting if the fair value, vesting condition or classification of the award is not the same immediately before and after a change to the terms and conditions of the award. This ASU is effective on a prospective basis beginning on January 1, 2018, with early adoption permitted. The Company adopted this ASU effective January 1, 2018, and the adoption did not have an impact on the Company's condensed consolidated financial statements and related disclosures.

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## New Accounting Pronouncements – Recently Issued

In February 2016, the FASB issued ASU No. 2016-02, a comprehensive new lease accounting standard, which provides revised guidance on accounting for lease arrangements by both lessors and lessees and requires lessees to recognize a lease liability and a right-of-use asset for most leases. This ASU also requires additional disclosures. The new guidance is effective for fiscal years beginning after December 15, 2018, and interim periods within those fiscal years, with early adoption permitted. The new standard must be applied using a modified retrospective transition approach that requires application of the new guidance for all periods presented. Although its assessment is not complete, the Company currently expects the adoption of this guidance to result in the addition of material balances of leased assets and corresponding lease liabilities to its consolidated balance sheets, primarily relating to leases of office space.

## 3. Net Loss per Share

Basic and diluted net loss per common share is calculated by dividing net loss applicable to common stockholders by the weighted-average number of common shares outstanding during the period, without consideration of common stock equivalents. The Company's potentially dilutive shares, which include outstanding stock options, Employee Stock Purchase Plan awards, unvested restricted stock units, or RSUs, and shares issuable upon conversion of the Convertible Notes, are considered to be common stock equivalents and are only included in the calculation of diluted net loss per share when their effect is dilutive.

The following table presents amounts that were excluded from the calculation of diluted net loss per share, due to their anti-dilutive effect (in thousands):

|                                                            | Three Months<br>Ended March 31, |        |
|------------------------------------------------------------|---------------------------------|--------|
|                                                            | 2017                            | 2018   |
| Outstanding stock options and Employee Stock Purchase Plan | 7,366                           | 8,067  |
| Unvested restricted stock units                            | 1,093                           | 2,197  |
| Shares issuable upon conversion of Convertible Notes       | 4,186                           | 1,441  |
|                                                            | 12,645                          | 11,705 |

In September 2014, the Company issued Convertible Notes, which provide in certain situations for the conversion of the outstanding principal amount of the Convertible Notes into shares of the Company's common stock at a predefined conversion rate. See Note 5, "Debt", for additional information. In conjunction with the issuance of the Convertible Notes, the Company entered into capped call option transactions, or Capped Calls, with certain counterparties. The Capped Calls are expected generally to reduce the potential dilution, and/or offset, to an extent, the cash payments the Company may choose to make in excess of the principal amount, upon conversion of the Convertible Notes.

As provided by the terms of the indenture underlying the Convertible Notes, the Company has a choice to settle the conversion obligation for the Convertible Notes in cash, shares or any combination of the two. The Company currently intends to settle the par value of the Convertible Notes in cash and any excess conversion premium in shares. Accordingly, the par value of the Convertible Notes will not be included in the calculation of diluted net income per share, but the dilutive effect of the conversion premium will be considered in the calculation of diluted net income per share using the treasury stock method. The share figures in the table above represent the estimated incremental shares that would be issued, after consideration of the Capped Calls, assuming conversion of all of the outstanding Convertible Notes as of March 31, 2017 and 2018.

#### 4. Inventories

The following table presents inventories as of December 31, 2017 and March 31, 2018 (in thousands):

|                   | December<br>31,<br>2017 | March 31,<br>2018 |
|-------------------|-------------------------|-------------------|
| Raw materials     | \$ 17,876               | \$ 22,428         |
| Work in process   | 38,629                  | 50,231            |
| Finished goods    | 1,434                   | 1,785             |
| Total inventories | \$ 57,939               | \$ 74,444         |

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Inventories are related to the Company's approved products, primarily ZEJULA. If future sales of ZEJULA or VARUBI are less than expected, the Company may be required to write down the value of such inventories.

5. Debt

Our outstanding debt obligations consisted of the following (in thousands):

|                            | December<br>31,<br>2017 | March 31,<br>2018 |
|----------------------------|-------------------------|-------------------|
| Convertible notes, net     | \$ 143,446              | \$ 146,529        |
| Term Loan (Tranche A), net | 293,659                 | 293,888           |
| Total long-term debt       | \$ 437,105              | \$ 440,417        |

Convertible Notes

On September 29, 2014, in a registered underwritten public offering, the Company completed the issuance of \$201.3 million aggregate principal amount of Convertible Notes. In conjunction with the sale of the Convertible Notes, the Company used \$20.8 million of the net proceeds to enter into separate Capped Calls.

The Convertible Notes bear interest at a rate of 3.00% per annum, payable semi-annually on April 1 and October 1, and will be convertible into cash, shares of the Company's common stock or a combination of cash and shares of the Company's common stock, at the Company's election. The Convertible Notes will mature on October 1, 2021, unless earlier converted or repurchased in accordance with their terms. Prior to the close of business on the business day immediately preceding April 1, 2021, the Convertible Notes will be convertible only upon the occurrence of certain events and during certain periods as discussed below, and thereafter, at any time until the close of business on the second scheduled trading day immediately preceding the maturity date. The initial conversion price of the Convertible Notes is approximately \$35.13 per share of common stock at an initial conversion rate of 28.4627 shares of the Company's common stock per \$1,000 principal amount of Convertible Notes.

The conversion rate is subject to adjustment from time to time upon the occurrence of certain events, including, but not limited to, the issuance of stock dividends and payment of cash dividends. At any time prior to the close of business on the business day immediately preceding April 1, 2021, holders may convert their Convertible Notes at their option only under the following circumstances:

- (1) during any calendar quarter commencing after the calendar quarter ending on December 31, 2014 (and only during such calendar quarter), if the closing sale price of the Company's common stock for at least 20 trading days (whether or not consecutive) during a period of 30 consecutive trading days ending on the last trading day of the immediately preceding calendar quarter in which the conversion occurs is greater than 130% of the conversion price on each applicable trading day;
- (2) during the five business day period after any ten consecutive trading day period, or the measurement period, in which the trading price per \$1,000 principal amount of the Convertible Notes for each trading day of the measurement period was less than 98% of the product of the closing sale price of the Company's common stock and the conversion rate on each such trading day; or
- (3) upon the occurrence of specified corporate events.

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As of March 31, 2018, the carrying value of the Convertible Notes, net of unamortized discount and debt issuance costs, was \$146.5 million and the estimated fair value of the principal amount was \$369.0 million. As provided by the terms of the indenture underlying the Convertible Notes, the Company has a choice to settle the conversion obligation for the Convertible Notes in cash, shares or any combination of the two. The Company currently intends to settle the par value of the Convertible Notes in cash and any excess conversion premium in shares.

The following table presents total interest expense recognized related to the Convertible Notes during the three months ended March 31, 2017 and 2018 (in thousands):

|                                     | Three Months Ended<br>March 31, |          |
|-------------------------------------|---------------------------------|----------|
|                                     | 2017                            | 2018     |
| Contractual interest expense        | \$ 1,509                        | \$ 1,509 |
| Amortization of debt discount       | 2,615                           | 2,953    |
| Amortization of debt issuance costs | 143                             | 130      |
| Total interest expense              | \$ 4,267                        | \$ 4,592 |

## 2017 Term Loan Agreement

In November 2017, the Company entered into a loan agreement, or the Loan Agreement, with BioPharma Credit PLC and affiliates, or the Lenders. The Lenders agreed to provide up to an aggregate principal amount of \$500.0 million in two tranches, with the first tranche equal to \$300.0 million, or Tranche A, and the second in an amount between \$50.0 million and \$200.0 million at the Company's discretion, or Tranche B. The Company drew Tranche A on December 6, 2017 with a maturity date of December 6, 2024. Tranche B is available for draw at the Company's option, on 90 days' notice, from June 30, 2018 until December 20, 2018 and if drawn by the Company, will also have a maturity date of December 6, 2024. Borrowings under the Tranche A and Tranche B loans bear interest at rates equal to the London Interbank Offered Rate, or LIBOR, plus an applicable margin of 8% per annum and 7.5% per annum, respectively (with the LIBOR rate subject to a floor of 1% and cap equal to the LIBOR rate as of the Tranche A closing date plus 1.5%). The loans have an up-front fee of 2% on the funded amount of each tranche, payable at the applicable closing date.

The Tranche A loan was recorded on the condensed consolidated balance sheets, net of a debt discount of \$6.0 million in upfront fees assessed by the Lenders at the time of borrowing. The debt discount and deferred financing costs of \$0.4 million are being amortized to interest expense using the effective interest method over the same term. The effective annual interest rate of the outstanding debt under Tranche A loan is approximately 10.0%. For the three months ended March 31, 2018, the Company recognized \$7.5 million of interest expense related to the Tranche A loan, including \$0.2 million related to the accretion of debt discount and amortization of deferred financing costs.

## 6. Accrued Expenses

The following table presents the components of accrued expenses (in thousands):

|                                          | December<br>31,<br>2017 | March 31,<br>2018 |
|------------------------------------------|-------------------------|-------------------|
| Research and development                 | \$ 55,949               | \$ 57,285         |
| Salaries, bonuses and other compensation | 33,717                  | 23,625            |
| Product revenue allowances               | 22,847                  | 18,590            |
| Inventory                                | 16,469                  | 28,633            |
| Sales and marketing                      | 6,701                   | 7,347             |
| Royalties                                | 6,552                   | 4,982             |
| Professional services                    | 3,944                   | 3,101             |
| Other                                    | 8,629                   | 7,026             |
| Total accrued expenses                   | \$ 154,808              | \$ 150,589        |

## 7. Stock-Based Compensation

The Company maintains several equity compensation plans, including the TESARO, Inc. 2012 Omnibus Incentive Plan, or the 2012 Incentive Plan, the TESARO, Inc. 2010 Stock Incentive Plan, or the 2010 Incentive Plan, the TESARO, Inc. 2015 Non-Employee Director Stock Incentive Plan, or the 2015 Director Plan, and the TESARO, Inc. 2012 Employee Stock Purchase Plan, or the 2012 ESPP.

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On April 27, 2012, the stockholders of the Company approved the 2012 Incentive Plan, which had been previously adopted by the board of directors. Upon effectiveness of the 2012 Incentive Plan, the Company ceased making awards under the 2010 Incentive Plan. The 2012 Incentive Plan initially allowed the Company to grant awards for up to 1,428,571 shares of common stock plus the number of shares of common stock available for grant under the 2010 Incentive Plan as of the effectiveness of the 2012 Incentive Plan (an additional 6,857 shares) plus the number of shares of common stock related to awards outstanding under the 2010 Incentive Plan that terminate by expiration, forfeiture, cancellation, cash settlement or otherwise. The number of shares available for grants of awards under the 2012 Incentive Plan is increased automatically each January 1 by a number of shares of common stock equal to the lesser of 4% of the shares of common stock then outstanding or the number of shares determined by the Company's board of directors. Most recently, on January 1, 2017 and 2018, the number of shares authorized for issuance under the 2012 Incentive Plan was increased by 2,144,867 shares and 2,178,561 shares, respectively. Awards under the 2012 Incentive Plan may include the following award types: stock options, which may be either incentive stock options or nonqualified stock options; stock appreciation rights; restricted stock; RSUs; dividend equivalent rights; performance shares; performance units; cash-based awards; other stock-based awards, including unrestricted shares; or any combination of the foregoing. The exercise price of stock options granted under the 2012 Incentive Plan is equal to the closing price of a share of the Company's common stock on the grant date.

The 2015 Director Plan allows the Company to grant awards for up to 500,000 shares of common stock. Awards under the 2015 Director Plan may include the following award types: stock options; stock appreciation rights; restricted stock; RSUs; unrestricted stock; or any combination of the foregoing. The exercise price of stock options granted under the 2015 Director Plan is equal to the closing price of a share of the Company's common stock on the grant date.

The following table presents stock-based compensation expense as reflected in the Company's condensed consolidated statements of operations and comprehensive loss (in thousands):

|                                                             | Three Months Ended<br>March 31, |           |
|-------------------------------------------------------------|---------------------------------|-----------|
|                                                             | 2017                            | 2018      |
| Research and development                                    | \$ 7,125                        | \$ 7,940  |
| Selling, general and administrative                         | 11,276                          | 18,493    |
| Subtotal                                                    | 18,401                          | 26,433    |
| Capitalized stock-based compensation costs                  | —                               | (305)     |
| Stock-based compensation expense included in total expenses | \$ 18,401                       | \$ 26,128 |

## Stock Options

The following table presents a summary of the Company's stock option activity and related information:

|                                  | Shares    | Weighted-average<br>exercise price per<br>share |
|----------------------------------|-----------|-------------------------------------------------|
| Outstanding at December 31, 2017 | 6,908,313 | \$ 52.00                                        |
| Granted                          | 1,366,909 | 56.12                                           |
| Exercised                        | (121,278) | 37.08                                           |
| Cancelled                        | (141,904) | 90.70                                           |
| Outstanding at March 31, 2018    | 8,012,040 | \$ 52.24                                        |
| Vested at March 31, 2018         | 4,558,825 | \$ 36.97                                        |

At March 31, 2018, there was approximately \$132.6 million of unrecognized compensation cost related to unvested stock options, which the Company expects to recognize over a remaining weighted-average period of 2.67 years.

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## Restricted Stock Units

The following table presents a summary of the Company's RSU activity and related information:

|                                                      | Shares    | Weighted-average<br>grant date fair<br>value per share |
|------------------------------------------------------|-----------|--------------------------------------------------------|
| Unvested restricted stock units at December 31, 2017 | 1,159,118 | \$ 115.01                                              |
| Granted                                              | 1,315,916 | 59.07                                                  |
| Vested                                               | (216,319) | 108.51                                                 |
| Forfeited                                            | (62,001)  | 114.08                                                 |
| Unvested restricted stock units at March 31, 2018    | 2,196,714 | \$ 82.17                                               |

At March 31, 2018, there was approximately \$166.8 million of unrecognized compensation cost related to unvested RSUs, which the Company expects to recognize over a remaining weighted-average period of 3.19 years.

## ESPP

Under the Company's 2012 ESPP, an aggregate of 275,000 shares of common stock have been reserved for issuance pursuant to purchase rights granted to the Company's employees or to employees of the Company's designated subsidiaries. As of March 31, 2018, 151,859 shares remained available for issuance. During the three months ended March 31, 2017 and 2018, the Company did not issue any shares under the 2012 ESPP, and recognized approximately \$0.5 million and \$0.4 million in related stock-based compensation expense, respectively.

## 8. Income Taxes

Deferred tax assets and deferred tax liabilities are determined based on temporary differences between the financial reporting and tax bases of assets and liabilities and are measured using the enacted tax rates and laws that will be in effect when the differences are expected to reverse. A valuation allowance is recorded against deferred tax assets if it is more likely than not that some portion or all of the deferred tax assets will not be realized.

The Company does not recognize a tax benefit for uncertain tax positions unless it is more likely than not that the position will be sustained upon examination by tax authorities, including resolution of any related appeals or litigation

processes, based on the technical merits of the position. The tax benefit that is recorded for these positions is measured at the largest amount of cumulative benefit that has greater than a 50 percent likelihood of being realized upon ultimate settlement. Deferred tax assets that do not meet these recognition criteria are not recorded and the Company recognizes a liability for uncertain tax positions that may result in tax payments. If such unrecognized tax benefits were realized and not subject to valuation allowances, the entire amount would impact the tax provision. As of March 31, 2018, the Company's uncertain tax positions were subject to valuation allowances.

As of March 31, 2018, the Company continues to consider interpretations of the application of SEC Staff Accounting Bulletin No. 118, and has not finalized incremental accounting adjustments related to the Tax Cuts and Jobs Act of 2017.

The Company recorded provisions for income taxes for the three months ended March 31, 2017 and 2018 of \$0.1 million and \$0.4 million, respectively. The provision for income taxes consists of current tax expense, which relates primarily to the Company's subsidiary operations in non-U.S. tax jurisdictions.

## 9. Intangible Assets

The following table presents intangible assets as of December 31, 2017 and March 31, 2018 (in thousands):

|                                 | December 31,<br>2017 | March 31,<br>2018 | Estimated useful life |
|---------------------------------|----------------------|-------------------|-----------------------|
| Acquired and in-licensed rights | \$ 64,665            | \$ 64,665         | 8-15 Years            |
| Less accumulated amortization   | (8,281)              | (9,718)           |                       |
| Total intangible assets, net    | \$ 56,384            | \$ 54,947         |                       |

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The Company recorded \$0.5 million and \$1.4 million in amortization expense related to intangible assets during the three months ended March 31, 2017 and 2018, respectively. Estimated future amortization expense for intangible assets as of March 31, 2018 is \$4.5 million for the remainder of 2018, \$6.0 million per year for 2019, 2020, 2021, and 2022, and \$26.5 million thereafter.

### 10. Commitments and Contingencies

The Company leases approximately 275,000 square feet of office space in Waltham, Massachusetts under a non-cancelable operating lease agreement. The Company also leases office space in several locations throughout Europe. The Company recognizes rental expense on a straight-line basis over the respective lease term including any free rent periods and tenant allowances.

Future minimum rental commitments under the Company's leased properties as of March 31, 2018 were \$8.4 million for the remainder of the year ending December 31, 2018 and \$12.1 million, \$8.0 million, \$3.8 million, \$3.6 million and \$2.3 million for the years ending December 31, 2019, 2020, 2021, 2022 and thereafter, respectively.

The Company has entered into agreements with certain vendors for the provision of services, including services related to commercial manufacturing, data management and clinical operation support, that the Company is not able to terminate for convenience under its contracts, and thus avoid any and all future obligations to the vendors. Under such agreements, the Company is contractually obligated to make certain minimum payments to the vendors, with the exact amounts in the event of termination to be based on the timing of the termination and the exact terms of the agreement.

The Company has certain obligations under licensing agreements with third parties that are contingent upon achieving various development, regulatory and commercial milestones. Pursuant to these license agreements, the Company is required to make milestone payments if certain development, regulatory and commercial sales milestones are achieved, and may have certain additional research funding obligations. Also, pursuant to the terms of each of these license agreements, when and if commercial sales of a product commence, the Company will pay royalties to its licensors on net sales of the respective products.

### Litigation and Other Proceedings

The Company may periodically become subject to legal proceedings and claims arising in connection with ongoing business activities, including claims or disputes related to patents that have been issued or that are pending in the field of research on which the Company is focused.

A putative class action complaint was filed on January 17, 2018 in the United States District Court for the District of Massachusetts, entitled Roger Bowers v. TESARO Incorporated (sic), et. al., Case No. 18-10086. The complaint alleges that the Company and its Chief Executive Officer and its Chief Financial Officer violated certain federal securities laws, specifically under Sections 10(b) and 20(a) of the Securities Exchange Act of 1934, as amended, and Rule 10b-5 thereunder. The plaintiff seeks unspecified damages on behalf of a purported class of purchasers of the Company's common stock between March 14, 2016 and January 12, 2018. On March 19, 2018, six separate applicants filed motions seeking appointment as lead plaintiff. Four of these applicants subsequently withdrew their motions or indicated that they did not oppose a competing motion filed by another applicant. The Court has not yet ruled on the remaining motions or selected a lead plaintiff. The Court has also not set a trial date for this matter. The Company believes that the allegations contained in the complaint are without merit and intends to defend the case vigorously. The Company has not recorded an estimated liability associated with this legal proceeding as it does not believe that such a liability is probable.

## 11. Revenue Recognition

### Product Revenue, Net

The Company sells its products principally to a limited number of specialty distributors and specialty pharmacy providers in the U.S., and directly to hospitals and clinics as well as to certain wholesale distributors in Europe, or collectively, its Customers. These Customers subsequently resell the Company's products to health care providers and patients. In addition to distribution agreements with Customers, the Company enters into arrangements with health care

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providers and payors that provide for government-mandated and/or privately-negotiated rebates, chargebacks and discounts with respect to the purchase of the Company's products.

Revenues from product sales are recognized when the Customer obtains control of the Company's product, which occurs at a point in time, typically upon delivery to the Customer. When the Company performs shipping and handling activities after the transfer of control to the Customer (e.g., when control transfers prior to delivery), they are considered as fulfillment activities, and accordingly, the costs are accrued for when the related revenue is recognized. Taxes collected from Customers relating to product sales and remitted to governmental authorities are excluded from revenues.

**Product Revenue.** Net product revenue relates to sales of ZEJULA and VARUBI/VARUBY. The Company commenced sales of ZEJULA in the U.S. in April 2017 and in Europe in December 2017. The following table presents net product revenues by product for the three months ended March 31, 2017 and 2018, respectively (in thousands).

|                      | Three months ended<br>March 31, |           | Change    |            |
|----------------------|---------------------------------|-----------|-----------|------------|
|                      | 2017                            | 2018      | Amount    | Percentage |
| ZEJULA               | \$ —                            | \$ 48,869 | \$ 48,869 | n/m        |
| VARUBI/VARUBY        | 2,139                           | 1,303     | (836)     | (39)%      |
| Product revenue, net | \$ 2,139                        | \$ 50,172 | \$ 48,033 | 2,246%     |

n/m = not meaningful

The following table summarizes activity in each of the product revenue allowance and reserve categories for the three months ended March 31, 2017 and 2018 (in thousands):

|                                           | Chargebacks,<br>discounts and<br>fees | Government<br>and other<br>rebates | Returns | Total    |
|-------------------------------------------|---------------------------------------|------------------------------------|---------|----------|
| Balance at December 31, 2016              | \$ 177                                | \$ 1,312                           | \$ 18   | \$ 1,507 |
| Provision related to current period sales | 736                                   | 562                                | 8       | 1,306    |
| Adjustment related to prior period sales  | —                                     | —                                  | —       | —        |
| Credit or payments made during the period | (756)                                 | (1,157)                            | —       | (1,913)  |
| Balance at March 31, 2017                 | \$ 157                                | \$ 717                             | \$ 26   | \$ 900   |

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|                                           |          |          |           |           |
|-------------------------------------------|----------|----------|-----------|-----------|
| Balance at December 31, 2017              | \$ 2,088 | \$ 6,450 | \$ 16,350 | \$ 24,888 |
| Provision related to current period sales | 3,310    | 3,115    | 194       | 6,619     |
| Adjustment related to prior period sales  | —        | —        | —         | —         |
| Credit or payments made during the period | (4,208)  | (3,801)  | (3,697)   | (11,706)  |
| Balance at March 31, 2018                 | \$ 1,190 | \$ 5,764 | \$ 12,847 | \$ 19,801 |

License, Collaboration and Other Revenues

The Company enters into out-licensing agreements which are within the scope of Topic 606, under which it licenses certain rights to its product candidates to third parties. The terms of these arrangements typically include payment to the Company of one or more of the following: non-refundable, up-front license fees; development, regulatory and commercial milestone payments; payments for manufacturing supply services the Company provides through its contract manufacturers; and royalties on net sales of licensed products. Each of these payments results in license, collaboration and other revenues, except for revenues from royalties on net sales of licensed products, which are classified as royalty revenues.

In determining the appropriate amount of revenue to be recognized as it fulfills its obligations under each of its agreements, the Company performs the following steps: (i) identification of the promised goods or services in the contract; (ii) determination of whether the promised goods or services are performance obligations including whether they are distinct in the context of the contract; (iii) measurement of the transaction price, including the constraint on variable consideration; (iv) allocation of the transaction price to the performance obligations; and (v) recognition of revenue when (or as) the Company satisfies each performance obligation.

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The following table presents changes in the Company's contract assets and liabilities during the three months ended March 31, 2017 and 2018 (in thousands):

|                                   |          | Balance at<br>Beginning<br>of Period | Additions  | Deductions | Balance at<br>End of Period |
|-----------------------------------|----------|--------------------------------------|------------|------------|-----------------------------|
| Three months ended March 31, 2017 |          |                                      |            |            |                             |
| Contract assets                   |          | \$ 1,000                             | \$ —       | \$ —       | \$ 1,000                    |
| Contract liabilities:             |          |                                      |            |            |                             |
| Deferred revenue                  |          | \$ 399                               | \$ —       | \$ (23)    | \$ 376                      |
| Three months ended March 31, 2018 |          |                                      |            |            |                             |
| Contract assets                   | \$ 1,000 | \$ —                                 | \$ (1,000) | \$ —       |                             |
| Contract liabilities:             |          |                                      |            |            |                             |
| Deferred revenue                  | \$ 306   | \$ —                                 | \$ (24)    | \$ 282     |                             |

During the three months ended March 31, 2017 and 2018, the Company recognized the following revenues as a result of changes in the contract asset and the contract liability balances in the respective periods (in thousands):

|                                                                                              | Three Months<br>Ended<br>March 31,<br>2017      2018 |            |
|----------------------------------------------------------------------------------------------|------------------------------------------------------|------------|
| Revenue recognized in the period from:                                                       |                                                      |            |
| Amounts included in the contract liability or contract assets at the beginning of the period | \$ 23                                                | \$ 24      |
| Performance obligations satisfied in previous periods                                        | \$ —                                                 | \$ (1,000) |

The \$(1.0) million noted above for the three months ended March 31, 2018 was a reversal of license revenue based on a re-evaluation of the probability of a milestone being achieved, as more fully described in Note 12, "Collaboration Arrangements", under "Jiangsu Hengrui Medicine Co., Ltd."

## 12. Collaboration Arrangements

## Merck Collaboration

In May 2015, the Company entered into a research agreement with Merck Sharp & Dohme B.V., a subsidiary of Merck, to perform a trial to evaluate the preliminary safety and efficacy of niraparib plus KEYTRUDA® in patients with triple negative breast cancer and patients with ovarian cancer. Under the terms of this agreement, the Company is responsible for providing niraparib study materials and for carrying out clinical research activities. The Company and Merck share in the external costs of the study equally, with certain exceptions. The Company records cost-sharing payments due from Merck as reductions of research and development expense. During the three months ended March 31, 2017 and 2018, the Company incurred \$2.1 million and \$1.8 million in external costs related to this study, of which \$1.0 million and \$0.9 million is reimbursable by Merck, respectively. At March 31, 2018, \$2.2 million of cost-sharing receivable from Merck has been recorded in other current assets on the condensed consolidated balance sheets.

## Out-Licenses

### Takeda Pharmaceutical Co., Ltd.

On July 27, 2017, the Company entered into an exclusive license agreement, or the Takeda Agreement, with Millennium Pharmaceuticals, Inc., a wholly-owned subsidiary of Takeda Pharmaceutical Company Limited, or Takeda. Pursuant to the Takeda Agreement, the Company granted Takeda licenses under certain patent rights and know-how relating to niraparib to develop and commercialize niraparib for the treatment of all tumor types in Japan, and all tumor types excluding prostate cancer in South Korea, Taiwan, Russia and Australia.

In connection with the Takeda Agreement, the Company received a \$100.0 million up-front payment and is eligible to receive additional payments of up to \$140.0 million related to the achievement of certain clinical development and regulatory milestones as well as up to \$100.0 million related to the achievement of additional sales milestones. The Company will also be eligible to receive tiered royalties from Takeda based on percentages of net product sales ranging from the high teens to low thirties. Takeda is responsible for conducting and funding all development and

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commercialization of niraparib in the licensed territories, including research, development, regulatory and commercialization activities. Unless earlier terminated, the Takeda Agreement will continue in effect until the date on which the royalty term and all payment obligations with respect to all products in all countries have expired.

The Company identified the following performance obligations at the inception of the Takeda Agreement: (1) exclusive license with rights to develop and commercialize niraparib to Takeda in the licensed territories for the associated tumor types, and (2) initial supply to Takeda of certain materials for the manufacture of niraparib. In addition, the Company may also become responsible for manufacturing certain niraparib products for clinical and commercial supply and providing technical assistance related to the transfer of know-how, at Takeda's option, for the manufacture of niraparib for which the Company will receive reimbursement that approximates stand-alone selling prices.

Revenue associated with the transfer of the license was fully recognized during the third quarter of 2017, the performance obligation is fully satisfied, and no changes have occurred in the transaction price during the three months ended March 31, 2018.

Revenue associated with the initial supply of niraparib materials will be recognized when delivered to Takeda. During the three months ended March 31, 2018, the Company recognized \$0.1 million as other revenues within license, collaboration and other revenues in the Company's condensed consolidated statements of operations and comprehensive loss related to materials delivered to Takeda. No revenue was recognized during the three months ended March 31, 2017. No changes have occurred in the transaction price of previously delivered goods during the three months ended March 31, 2018.

Janssen Biotech, Inc.

Under the terms of the Company's collaboration agreement with Janssen Biotech, Inc., or Janssen, the Company granted Janssen licenses under certain patent rights and know-how relating to niraparib for prostate cancer worldwide, except for Japan. Janssen will conduct all development and commercialization of niraparib in the field of prostate cancer worldwide (excluding Japan).

Pursuant to the collaboration agreement, within 30 days after the date of the collaboration agreement, the Company provided Janssen with electronic copies of certain know-how relating to development of niraparib. In addition, at Janssen's request and in return for certain reimbursement, the Company is also responsible for manufacturing and supplying to Janssen all of Janssen's requirements of active pharmaceutical ingredient, or API, and finished drug product, for niraparib and niraparib products to be used by Janssen for its development activities in prostate cancer indications. Also at Janssen's request, the Company is responsible for manufacturing of certain niraparib products and API for commercial sale in the field of prostate cancer. In both cases, if Janssen exercises its right to receive the

manufacturing services, the Company will receive reimbursement that will at least cover its cost of providing such services.

The Company received a \$35.0 million up-front, non-refundable license fee from Janssen. Assuming successful development and commercialization of niraparib products for prostate cancer, the Company could receive up to an additional \$43.0 million in clinical milestones and \$372.0 million in regulatory and sales milestones as well as tiered, double-digit royalties on aggregate net sales of products in the field of prostate cancer. Janssen is responsible for funding all development and commercialization of niraparib in prostate cancer worldwide (excluding Japan), including research, development, manufacturing, regulatory and commercialization activities. Janssen may terminate the collaboration agreement at any time upon 90 days' written notice, upon termination of the Company's license agreement with Merck or in the event of certain safety concerns. Either party may terminate the collaboration agreement for uncured material breach or bankruptcy. Unless earlier terminated, the collaboration agreement will continue in effect until the date on which the royalty term and all payment obligations with respect to all products in all countries have expired.

The Company assessed this arrangement in accordance with Topic 606 and concluded that the contract counterparty, Janssen, is a customer. The Company identified the following material promises under the contract: (1) the licenses under certain patent rights relating to niraparib for prostate cancer worldwide, except for Japan, and transfer of certain development and regulatory information; and (2) the obligation to participate in Joint Committees. In addition, the Company identified the following customer options that will create manufacturing obligations for the Company upon

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exercise by Janssen: (1) the supply of API and niraparib products for Janssen's development and commercial needs; and (2) the supply of niraparib for Janssen's clinical trial needs. The Company considered the manufacturing capabilities of Janssen, Janssen's right to sublicense and manufacture API, and the fact that the manufacturing services are not proprietary and can be provided by other vendors, to conclude that the license has stand-alone functionality and is distinct. The Company's obligation to participate in the Joint Committees and provide development, regulatory and commercialization information to Janssen does not significantly impact or modify the licenses' granted functionality. Further, the customer options for manufacturing services were evaluated as a material right, but were concluded to be immaterial to the Company's financial statements. Based on these assessments, the Company identified the license and the participation in Joint Committees as the only performance obligations at the inception the arrangement, which were both deemed to be distinct.

Revenue associated with the transfer of the license was fully recognized during the second quarter of 2016, the performance obligation is fully satisfied, and no changes have occurred in the transaction price during the three months ended March 31, 2018.

Revenue associated with the Joint Committees performance obligation, \$0.5 million, is being recognized on a straight-line basis over a period of five years, which, in management's judgment, is the best measure of progress toward satisfying the performance obligation and represents the Company's best estimate of the period of the obligation to participate in the Joint Committees. The remaining transaction price of \$0.3 million is recorded in deferred revenue as of March 31, 2018 on the condensed consolidated balance sheets and will be recognized as revenue over the remaining period of 36 months.

Revenue associated with the materials supply service is recognized when the material is delivered to Janssen. During the three months ended March 31, 2017 and 2018, the Company recognized \$0.9 million and \$0.5 million, respectively, as other revenues within license, collaboration and other revenues in the Company's condensed consolidated statements of operations and comprehensive loss related to materials delivered to Janssen. No changes have occurred in the transaction price of previously delivered goods during the three months ended March 31, 2018.

Zai Lab (Shanghai) Co., Ltd.

On September 28, 2016, the Company entered into a Collaboration, Development and License Agreement, or the Zai Agreement, with Zai Lab. Under the terms of the Zai Agreement, the Company exclusively licensed the rights to develop and commercialize niraparib to Zai Lab for China, Hong Kong and Macao, or the China Territories. Zai Lab will conduct all development, manufacturing and commercialization of niraparib in the China Territories, except for prostate cancer.

Under the terms of the Zai Agreement, the Company received a \$15.0 million up-front, non-refundable license fee from Zai Lab in the fourth quarter of 2016. Assuming successful development and commercialization of niraparib products in the China Territories, the Company could receive additional regulatory and sales milestones as well as tiered, double-digit royalties on aggregate net sales of products in the China Territories. Zai Lab is responsible for funding all development and commercialization of niraparib in the China Territories, including research, development, manufacturing, regulatory and commercialization activities. The term of the Zai Agreement continues, on a country-by-country basis, until the later of expiration of the last patent in the China Territories covering the niraparib product, or ten years from the first commercial sale in such country. The Zai Agreement may also be terminated by Zai Lab at any time upon prior written notice, or by either party for material breach or insolvency.

The Company identified three performance obligations under the contract. Revenue associated with all three performance obligations was fully recognized during 2016, the performance obligations are fully satisfied, and no changes have occurred in the transaction price during the three months ended March 31, 2018. No revenue was recognized during the three months ended March 31, 2017 and 2018.

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Jiangsu Hengrui Medicine Co., Ltd.

In July 2015, the Company entered into a license agreement with Jiangsu Hengrui Medicine Co., Ltd., or Hengrui, pursuant to which Hengrui has licensed the rights to develop, manufacture and commercialize rolapitant in the China Territories. The Company received a \$1.0 million up-front, non-refundable license fee from Hengrui in the fourth quarter of 2015. The Company has evaluated the terms of this arrangement under Topic 606 and has determined that there are two performance obligations: (1) exclusive license with rights to develop, manufacture and commercialize rolapitant in the China Territories; and (2) provision of technical assistance related to the know-how transfer for the development of the rolapitant formulations. The Company further determined that the transaction price for this arrangement included the \$1.0 million up-front consideration received and a future regulatory development milestone of \$1.0 million. This future milestone payment relates to the submission of the clinical trial application with the China Food and Drug Administration. The Company re-evaluates the transaction price in each reporting period and as uncertain events are resolved or other changes in circumstances occur.

During the first quarter of 2018, Hengrui ceased development activities related to the licensed product. All performance obligations were recognized in 2015 and 2016, including a \$1.0 million future regulatory development milestone that the Company assessed as probable. The Company re-evaluated the probability of the milestone, including its impact on the transaction price, and recognized a \$1.0 million reversal of license revenue during the three months ended March 31, 2018. No revenue was recognized during the three months ended March 31, 2017.

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Item 2. Management's Discussion and Analysis of Financial Condition and Results of Operations.

The following information should be read in conjunction with the unaudited financial information and the notes thereto included in this Quarterly Report on Form 10-Q and the audited financial information and the notes thereto included in our Annual Report on Form 10-K for the year ended December 31, 2017.

Except for the historical information contained herein, the matters discussed in this Quarterly Report on Form 10-Q may be deemed to be forward-looking statements that involve risks and uncertainties. We make such forward-looking statements pursuant to the safe harbor provisions of the Private Securities Litigation Reform Act of 1995 and other federal securities laws. In this Quarterly Report on Form 10-Q, words such as "may," "will," "expect," "anticipate," "estimate," "intend," and similar expressions (as well as other words or expressions referencing future events, conditions or circumstances) are intended to identify forward-looking statements.

Examples of forward-looking statements contained in this report include statements regarding the following: our commercialization plans for niraparib and rolapitant, including the progress of the commercial launches of ZEJULA® (niraparib) in the U.S. and Europe and VARUBI®/VARUBY® (rolapitant) in the U.S. and Europe; our intent to in-license or acquire additional product candidates; our expectations regarding product revenues and license, collaboration and other revenues; our expectation that research and development and selling, general and administrative expenses will increase in the future; our expectations regarding the timing and design of our development plans, the timing of regulatory filings, and the timing of data from clinical trials, including with respect to each of our niraparib, TSR-042, TSR-022 and TSR-033 programs; our expectations regarding new clinical trials of our product candidates, including the commencement and timing thereof; our expectations regarding our discovery and development plans for immunotherapy antibodies, including the timing thereof; our anticipated milestone and royalty payment obligations; our expectations that our operating losses and negative operating cash flows will continue, and possibly increase, for the foreseeable future; efforts to pursue strategic alternatives for the VARUBI/VARUBY brand, including potentially out-licensing the product line; our intent to settle the par value of the Convertible Notes in cash and any excess conversion premium in shares of common stock; and our needs for additional capital and the forecast of the period of time through which our financial resources will be adequate to support our operations.

Forward-looking statements are not guarantees of future performance. Actual future results, performance, achievements or the timing of certain events may differ significantly from those expressed or implied by the forward-looking statements. Risks and uncertainties involved in the forward-looking statements include, among others: uncertainties inherent in the development or commercialization of any new pharmaceutical product and the execution and completion of clinical trials; risks related to competition; the timing and availability of data from clinical trials; uncertainties regarding ongoing discussions with and actions by regulatory authorities; patient accrual rates for clinical trials; manufacturing and supply risks; risks related to intellectual property; and other matters that could affect the timing of data or the potential regulatory approval or commercial availability or success of our products. Forward-looking statements contained in this Quarterly Report on Form 10-Q should be considered in light of these factors and the factors discussed elsewhere in this Quarterly Report on Form 10-Q, and in light of factors discussed in our Annual Report on Form 10-K for the year ended December 31, 2017, including under the heading

“Risk Factors”. You should read carefully the factors described in the “Risk Factors” section to better understand the risks and uncertainties inherent in our business and underlying any forward-looking statements. You are also advised to consult any further disclosures we make on related subjects in our Quarterly Reports on Form 10-Q, Current Reports on Form 8-K and our website.

We caution readers not to place undue reliance on any forward-looking statements made by us, which speak only as of the date they are made. We disclaim any obligation, except as specifically required by law and the rules of the U.S. Securities and Exchange Commission, or the SEC, to publicly update or revise any such statements to reflect any change in our expectations or in events, conditions or circumstances on which any such statements may be based, or that may affect the likelihood that actual results will differ from those set forth in the forward-looking statements.

TESARO, the TESARO logo, VARUBI, VARUBY and ZEJULA are trademarks of TESARO, Inc. in the United States and in other selected countries. All other brand names or trademarks appearing in this report are the property of their respective holders. Unless the context requires otherwise, references in this report to “TESARO”, the “Company,” “we,” “us,” and “our” refer to TESARO, Inc.

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### Overview

We are a commercial-stage biopharmaceutical company devoted to providing transformative therapies to people bravely facing cancer. Our primary focus is to develop treatments for solid tumors using various approaches, including small molecules and immuno-oncology antibodies, as monotherapies and in combinations. We have in-licensed and are developing several oncology-related product candidates, and we have entered into several research collaborations with third parties for the discovery of new candidates. We have also entered into arrangements with other companies for the development and commercialization of certain of our product candidates in specific indications and/or geographies.

To date, we have received regulatory approvals for the following products, which are currently marketed and sold in the U.S. and in certain countries in Europe:

- The U.S. Food and Drug Administration, or FDA, approved ZEJULA® (niraparib) in March 2017 for the maintenance treatment of women with recurrent epithelial ovarian, fallopian tube, or primary peritoneal cancer who are in a complete or partial response to platinum-based chemotherapy. The European Commission, or EC, approved ZEJULA in November 2017 as a monotherapy for the maintenance treatment of adult patients with platinum-sensitive relapsed high grade serous epithelial ovarian, fallopian tube, or primary peritoneal cancer who are in complete or partial response to platinum-based chemotherapy.
- VARUBI® (rolapitant) is a potent and long-acting neurokinin-1, or NK-1, receptor antagonist for the prevention of chemotherapy induced nausea and vomiting, or CINV. The FDA approved the oral formulation of VARUBI in September 2015 for use in combination with other antiemetic agents in adults for the prevention of delayed (24 to 120 hours after chemotherapy administration) nausea and vomiting associated with initial and repeat courses of emetogenic cancer chemotherapy, including, but not limited to, highly emetogenic chemotherapy. The EC approved VARUBI®, the brand name of oral rolapitant in Europe, in April 2017 for the prevention of delayed nausea and vomiting associated with highly and moderately emetogenic chemotherapy in adults. The FDA approved the intravenous, or IV, formulation of VARUBI in October 2017. In January 2018, after post-marketing reports of side effects experienced following the commercial introduction of VARUBI IV, we updated the VARUBI IV package insert, including modifications to the contraindications, warnings and precautions, and adverse reactions sections, and issued a “Dear Healthcare Professional Letter” to healthcare providers to highlight the updates. In February 2018, we determined that we would cease marketing and distribution of VARUBI IV and pursue strategic alternatives for the VARUBI brand, including potentially out-licensing the VARUBI product line.

Recently, we have also reported data from various ongoing clinical trials of niraparib and TSR-042, as summarized below:

- In March 2018, we announced the presentation of preliminary data from the TOPACIO trial of niraparib in combination with an anti-PD-1 monoclonal antibody, KEYTRUDA®, at the 2018 Society for Gynecologic

Oncology Annual Meeting on Women's Cancer. Preliminary results suggest the combination of niraparib and an anti-PD-1 antibody could provide meaningful clinical benefit to patients with platinum-resistant or platinum-refractory ovarian cancer, regardless of biomarker status. Planning of a registration study is underway to support approval of ZEJULA and TSR-042 (our anti-PD-1 antibody) combination therapy.

- In April 2018, we presented summary initial data from the Phase 1 GARNET trial of TSR-042 in patients with microsatellite instability high, or MSI-H, endometrial cancer and non-small cell lung cancer, during the American Association for Cancer Research Annual Meeting. We expect to complete enrollment in the MSI-H endometrial cohort of the GARNET trial by the end of 2018. The GARNET trial is intended to support a biologics license application submission to the FDA in 2019 for patients with MSI-H cancers.
- In April 2018, we announced positive top-line results from the QUADRA trial, which was designed to assess clinical benefit of ZEJULA treatment in heavily pre-treated patients with ovarian cancer. Results successfully achieved the pre-specified primary endpoint and demonstrated ZEJULA monotherapy activity in a biomarker selected patient population. We intend to discuss a biomarker-focused regulatory submission with the FDA for a potential supplemental new drug application in the second half of 2018.

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As of March 31, 2018, we had an accumulated deficit of \$1.6 billion. Our net losses were \$162.8 million, \$496.1 million, \$374.2 million, and \$247.7 million for the three months ended March 31, 2018 and the years ended December 31, 2017, 2016, and 2015, respectively. We expect to incur significant expenses and operating losses for the foreseeable future. Overall, we expect operating expenses to increase over current levels as we continue to incur costs related to: (i) our ongoing U.S. and international commercialization and pre-commercial activities including executing related marketing and promotional programs for the launches and commercialization of ZEJULA; (ii) the advancement of clinical trial and other development and regulatory activities under our current development programs for niraparib, TSR-042, TSR-033 and TSR-022, and our collaborations; (iii) expanding our international operations; and (iv) other research and development activities and potential future collaborative or in-licensed development programs. Operating expenses (research and development and selling, general and administrative expenses) could fluctuate up or down from quarter to quarter, based on the timing and magnitude of activities taking place in a given quarter. In addition, future license payments or milestone payments could cause our total operating expenses and cash usage to fluctuate. If we obtain regulatory approval for any of our other product candidates, or if we anticipate the near term possibility of obtaining regulatory approval, we expect that we will incur significant additional commercialization expenses related to product sales, marketing, manufacturing and distribution. We will also continue to incur substantial interest expense related to our outstanding convertible debt and term loan. The actual amount of many of the expenditures described above will depend on numerous factors, including the timing of expenses and the timing, progress and results of our clinical trials and other development and regulatory activities, and commercialization efforts for ZEJULA. Accordingly, until we can generate a sufficient amount of revenue from our products, if ever, we expect to finance our operations in part through additional public or private equity or debt offerings, and we may seek additional capital through arrangements with strategic partners or from other sources. Adequate additional financing may not be available to us on acceptable terms, or at all. Our failure to raise capital as and when needed would have a negative impact on our financial condition and our ability to pursue our business strategy. We will need to generate significant revenues to achieve profitability, and we may never do so.

## Financial Operations Overview

### Revenues

Product revenue is derived from sales of ZEJULA and VARUBI in the United States, and ZEJULA and VARUBY in Europe.

License, collaboration and other revenues relate to our license agreements with Takeda, Janssen, Zai Lab and Hengrui. Takeda has licensed the rights to develop and commercialize niraparib for the treatment of all tumor types in Japan, and all tumor types excluding prostate cancer in certain other specified countries. Janssen has licensed the rights to develop, manufacture and commercialize niraparib worldwide (except for Japan) for the treatment of prostate cancer. Zai Lab has licensed the rights to develop and commercialize niraparib for China, Hong Kong and Macao, or the China Territories, except for prostate cancer. Hengrui has licensed the rights to develop, manufacture and commercialize rolapitant in the China Territories. During the first quarter of 2018, Hengrui ceased development activities related to rolapitant.

## Research and Development Expenses

Research and development expenses consist primarily of costs incurred for the development of our product candidates, which include:

- employee-related expenses, including salaries, bonuses, benefits, travel and stock-based compensation expense;
- fees and expenses incurred under agreements with contract research organizations, investigative sites, research consortia and other entities in connection with the conduct of clinical trials and preclinical studies and related services, such as administrative, data management, laboratory and biostatistics services;
- the cost of acquiring, developing and manufacturing active pharmaceutical ingredients for product candidates that have not received regulatory approval, clinical trial materials and other research and development materials;

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- pre-commercial license fees and milestone payments related to the acquisition of in-licensed product candidates, which are reported on our statements of operations as acquired in-process research and development;
- fees and costs related to regulatory filings and activities;
- facilities, depreciation and other expenses, which include direct and allocated expenses for rent, utilities, maintenance of facilities, insurance and other supplies; and
- other costs associated with clinical, preclinical, discovery and other research activities.

Research and development costs are expensed as incurred. License fees and development milestone payments related to in-licensed products and technology are expensed as acquired in-process research and development if it is determined at that point that they have no established alternative future use. Costs for certain development activities, such as clinical trials, are recognized based on an evaluation of the progress to completion of specific tasks using data such as patient enrollment, clinical site activations and information provided to us by our vendors.

Research and development activities are central to our business model. Product candidates in later stages of clinical development generally have higher development costs than those in earlier stages of clinical development, primarily due to the increased size and duration of later-stage clinical trials and manufacturing costs. We expect that our total future research and development costs will continue to increase over current levels, depending on the progress of our clinical development programs. We also anticipate increasing costs associated with our collaborations, manufacturing activities, and potential development milestone payments. More specifically, we expect costs to increase, including as we: continue our currently ongoing clinical trials, continue our manufacturing development and validation, and initiate additional investigative and collaborative studies related to niraparib; incur research and development related milestones; incur increased discovery, development and manufacturing related expenses associated with our immuno-oncology platform and related collaborations; lease additional facility space; and hire additional development and scientific personnel.

We cannot determine with certainty the duration and completion costs of the current or future clinical trials of our product candidates or if, when, or to what extent we will generate revenues from the commercialization and sale of any of our product candidates that obtain regulatory approval. We may never succeed in achieving regulatory approval for any of our currently unapproved product candidates. The duration, costs and timing of clinical trials and development of our product candidates will depend on a variety of factors, including the uncertainties of future clinical and preclinical studies, uncertainties in clinical trial enrollment rates and significant and changing government regulation. In addition, the probability of success for each product candidate will depend on numerous factors, including competition, manufacturing capability and commercial viability. We will determine which programs to pursue and how much to fund each program in response to the scientific and clinical success of each product candidate, as well as based upon an assessment of each product candidate's commercial potential. If we experience delays in the completion of, or the termination of, any clinical trial of our product candidates, the commercial prospects of our product candidates will be harmed, and our future ability to generate product revenues from any of these product candidates will be delayed or jeopardized. These occurrences would harm our business, financial

condition and prospects, perhaps significantly, which would require us to alter our current operating plan and potentially delay, scale back, or discontinue the development or commercialization of one or more programs and/or other areas of the business in order to reduce our future expenses and continue to fund our remaining operations.

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The following table presents research and development expenses and acquired in-process research and development expenses on a program-specific basis for our in-licensed products and product candidates for the three months ended March 31, 2017 and 2018 (in thousands):

|                                              | Three Months Ended<br>March 31, |           |
|----------------------------------------------|---------------------------------|-----------|
|                                              | 2017                            | 2018      |
| Niraparib Expenses                           |                                 |           |
| Acquired in-process research and development | \$ —                            | \$ —      |
| Research and development                     | 23,344                          | 30,688    |
| Niraparib total                              | 23,344                          | 30,688    |
| Immuno-Oncology Expenses                     |                                 |           |
| Acquired in-process research and development | —                               | —         |
| Research and development:                    |                                 |           |
| TSR-042                                      | 3,542                           | 11,611    |
| TSR-022                                      | 2,502                           | 6,658     |
| TSR-033                                      | 2,147                           | 2,401     |
| Combinations and other                       | 2,332                           | 2,371     |
| Immuno-Oncology total                        | 10,523                          | 23,041    |
| Rolapitant Expenses                          |                                 |           |
| Acquired in-process research and development | —                               | —         |
| Research and development                     | 3,558                           | 1,017     |
| Rolapitant total                             | 3,558                           | 1,017     |
| Personnel and Other Expenses                 | 28,697                          | 42,009    |
| Total                                        | \$ 66,122                       | \$ 96,755 |

For further discussion of the changes in our research and development expenses with respect to the three months ended March 31, 2018 and the corresponding period of 2017, see “Results of Operations — Comparison of the Three Months Ended March 31, 2017 and 2018 — Research and Development Expenses” below.

Personnel-related costs, depreciation and stock-based compensation are not allocated to any programs, as they are deployed across multiple projects under development and, as such, are separately classified as personnel and other expenses in the table above.

## Selling, General and Administrative Expenses

Selling, general and administrative expenses consist principally of salaries and related costs, including stock-based compensation, for our commercial personnel, including our field sales force, certain medical education professionals and other commercial support personnel, as well as personnel in executive and other administrative or non-research and development functions. Other selling, general and administrative expenses include certain facility-related costs, information technology costs, pre-commercial and commercial consulting, advertising, market research and other activities necessary to prepare for and support product launches, and professional fees for legal, patent review, consulting and accounting services.

We anticipate that our selling, general and administrative expenses will continue to increase in the future in support of our commercial and pre-commercial activities related to ZEJULA and potential other products, continued research and development activities, and the continued costs of operating as a public multinational company. These increases will likely include increased costs related to the hiring of additional personnel, executing marketing and promotional programs, hiring consultants, leasing of additional facility space, enhancing information technology systems, and legal and other professional fees, among other expenses.

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## Other Income and Expense

Other income and expense consists primarily of interest expense related to our debt instruments and interest income earned on cash and cash equivalents. A portion of the interest expense is non-cash expense relating to accretion of debt discounts and amortization of issuance costs.

## Results of Operations

## Comparison of the Three Months Ended March 31, 2017 and 2018

| (data in thousands)                           | Three Months Ended<br>March 31, |              | Change      |            |
|-----------------------------------------------|---------------------------------|--------------|-------------|------------|
|                                               | 2017                            | 2018         | Amount      | Percentage |
| Revenues:                                     |                                 |              |             |            |
| Product revenue, net                          | \$ 2,139                        | \$ 50,172    | \$ 48,033   | 2,246%     |
| License, collaboration and other revenues     | 934                             | (430)        | (1,364)     | n/m        |
| Total revenues                                | 3,073                           | 49,742       | 46,669      | 1,519%     |
| Expenses:                                     |                                 |              |             |            |
| Cost of sales – product                       | 444                             | 9,997        | 9,553       | 2,152%     |
| Cost of sales – intangible asset amortization | 490                             | 1,437        | 947         | 193%       |
| Research and development                      | 66,122                          | 96,755       | 30,633      | 46%        |
| Selling, general and administrative           | 69,262                          | 93,607       | 24,345      | 35%        |
| Acquired in-process research and development  | —                               | —            | —           | —          |
| Total expenses                                | 136,318                         | 201,796      | 65,478      | 48%        |
| Loss from operations                          | (133,245)                       | (152,054)    | (18,809)    | 14%        |
| Other income (expense), net                   | (3,426)                         | (10,346)     | (6,920)     | 202%       |
| Loss before income taxes                      | (136,671)                       | (162,400)    | (25,729)    | 19%        |
| Provision for income taxes                    | 54                              | 416          | 362         | 670%       |
| Net loss                                      | \$ (136,725)                    | \$ (162,816) | \$ (26,091) | 19%        |

n/m = not meaningful

Product Revenue. Net product revenue relates to sales of ZEJULA and VARUBI/VARUBY in the U.S. and Europe. The following table presents net product revenues by product for the three months ended March 31, 2017 and 2018, respectively (in thousands).

|                      | Three months ended<br>March 31, |           | Change    |            |
|----------------------|---------------------------------|-----------|-----------|------------|
|                      | 2017                            | 2018      | Amount    | Percentage |
| ZEJULA               | \$ —                            | \$ 48,869 | \$ 48,869 | n/m        |
| VARUBI/VARUBY        | 2,139                           | 1,303     | (836)     | (39)%      |
| Product revenue, net | \$ 2,139                        | \$ 50,172 | \$ 48,033 | 2,246%     |

n/m = not meaningful

We began to recognize revenues on sales of ZEJULA in the U.S. starting in the second quarter of 2017, and in Europe late in the fourth quarter of 2017. With respect to ZEJULA net product revenues for the three months ended March 31, 2018, the average net sales price per unit to us was approximately 90% of the Wholesale Acquisition Cost, or WAC, which is the gross list price at which our direct customers purchase each unit.

We began to recognize revenues on sales of VARUBI in the U.S. in the fourth quarter of 2015 and in Europe starting in the second quarter of 2017. For the three months ended March 31, 2018 as compared to the same period in 2017, net VARUBI product revenues decreased due to lower unit sales volumes and higher discounts, rebates and chargebacks. The average net sales price per unit to us was approximately 54% of WAC for the three months ended March 31, 2018 as compared to 62% of WAC for the same period in 2017. The percentage for the three months ended March 31, 2018 was impacted by a significant returns reserve for VARUBI IV units shipped prior to our determination in February 2018 to cease distribution of VARUBI IV.

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**License, Collaboration and Other Revenues.** License, collaboration and other revenues of \$0.9 million for the three months ended March 31, 2017 relates to our license agreement with Janssen. License, collaboration and other revenues of \$(0.4) million for the three months ended March 31, 2018 includes a \$(1.0) million reversal of milestone revenue recognized during 2015 and 2016 in connection with the Hengrui license agreement, due to our reassessment that the milestone is no longer probable of achievement, offset by \$0.6 million in other revenues recognized from shipments of materials to Janssen and Takeda.

**Cost of Sales - Product.** Cost of sales of \$0.4 million and \$10.0 million for the three months ended March 31, 2017 and 2018, respectively, consists of costs associated with the manufacturing of ZEJULA and VARUBI and royalties owed to our licensors for such sales, as well as costs of product provided under our sampling and other commercial programs and certain period costs. The increase was primarily related to the U.S. launch of ZEJULA in April 2017. Based on our policy to expense costs associated with the manufacture of our products prior to regulatory approval, certain of the costs of units recognized as revenue during the three months ended March 31, 2017 and 2018 were expensed prior to the respective initial regulatory approval dates, and therefore are not included in cost of sales during these periods. We expect cost of sales to increase in relation to net product revenues as we deplete these inventories, which is expected by the end of 2018 for ZEJULA.

**Cost of Sales - Intangible Asset Amortization.** The amounts of \$0.5 million and \$1.4 million for the three months ended March 31, 2017 and 2018, respectively, consist of amortization of intangible assets recorded as a result of post-approval milestones paid to licensors. The increase was primarily related to new milestone assets recorded during the year ended December 31, 2017 related to ZEJULA and VARUBY regulatory approvals.

**Research and Development Expenses.** Research and development expenses were \$66.1 million for the three months ended March 31, 2017, compared to \$96.8 million for the three months ended March 31, 2018, an increase of \$30.7 million. Significant changes resulting in this increase included:

- an increase of \$13.3 million in personnel and other costs (excluding stock-based compensation), primarily related to increased research and development headcount supporting the growth of our development activities, and costs related to research collaborations;
- an increase of \$12.5 million in costs associated with our immuno-oncology platform due to increased costs related to the TSR-042 and TSR-022 clinical trials, biologics manufacturing and non-clinical and other immuno-oncology program research activities; and
- an increase of \$7.3 million in costs associated with our niraparib program, primarily related to development of a new tablet formulation of niraparib and increased costs of ongoing clinical trials.

In addition, stock-based compensation expense included in research and development expenses increased by \$0.7 million, primarily due to increased awards of employee stock options and restricted stock units.

**Selling, General and Administrative Expenses.** Selling, general and administrative expenses were \$69.3 million for the three months ended March 31, 2017, compared to \$93.6 million for the three months ended March 31, 2018, an increase of \$24.3 million. The increase was primarily due to increases of: \$11.5 million in salaries, benefits and other personnel-related costs (excluding stock-based compensation), primarily due to the hiring of sales, marketing, medical affairs and other support personnel associated with the commercialization of ZEJULA, plus hiring to support our international operations; \$5.7 million in professional and consulting fees and other expenses related to commercial and pre-commercial activities such as global marketing and promotional programs and market research as well as other corporate operational activities; and \$7.0 million in stock-based compensation expense.

**Acquired In-Process Research and Development.** There were no acquired in-process research and development expenses for the three months ended March 31, 2017 or 2018.

**Other Income (Expense), Net.** Other income (expense) is primarily comprised of interest expense related to our Convertible Notes and our term loan, and interest income earned on cash and cash equivalents. Interest expense increased by \$7.8 million for the three months ended March 31, 2018 as compared to same period in 2017, primarily due to interest on the first tranche of the term loan, which we drew in December 2017. Interest income increased by \$0.8 million during the three months ended March 31, 2018 as compared to the same period in 2017, primarily due to higher yields on interest-bearing cash equivalents.

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## Liquidity and Capital Resources

## Sources of Liquidity

As of March 31, 2018, our principal source of liquidity was cash and cash equivalents, which totaled \$499.0 million. Since our inception in 2010, we have funded our operations primarily through public offerings of our common stock, the private placement of our equity securities, the issuance of convertible notes and a term loan financing. From inception through March 31, 2018, we received \$1.9 billion in total net cash proceeds from these sources.

## Cash Flows

The following table presents the primary sources and uses of cash for each of the periods noted (in thousands):

|                                                                               | Three Months Ended<br>March 31, |              |
|-------------------------------------------------------------------------------|---------------------------------|--------------|
|                                                                               | 2017                            | 2018         |
| Net cash provided by (used in):                                               | (as revised)                    |              |
| Operating activities                                                          | \$ (114,581)                    | \$ (147,632) |
| Investing activities                                                          | (1,790)                         | (1,619)      |
| Financing activities                                                          | 3,315                           | 4,497        |
| Effect of exchange rate changes on cash, cash equivalents and restricted cash | 45                              | 651          |
| Decrease in cash, cash equivalents and restricted cash                        | \$ (113,011)                    | \$ (144,103) |

## Cash Flows from Operating Activities

The use of cash in operating activities during both the three months ended March 31, 2017 and 2018 resulted primarily from our net losses adjusted for non-cash charges and changes in components of working capital. Net cash used in operating activities increased by \$33.1 million for the three months ended March 31, 2018 as compared to the same period in 2017, primarily due to higher costs associated with increased employee headcount and increased external expenses related to commercialization activities for ZEJULA. Increases in headcount in research and development and general and administrative functions, as well as increased external research and development expenses for niraparib and our immuno-oncology portfolio, also contributed to the increase in cash used in operating activities. Purchases of ZEJULA inventories during the three months ended March 31, 2018 also contributed to the increase.

## Cash Flows from Investing Activities

Net cash used in investing activities for the three months ended March 31, 2017 and 2018 of \$1.8 million and \$1.6 million, respectively, consisted of purchases of property and equipment, primarily related to the expansions our leased premises at our corporate headquarters in Waltham, Massachusetts.

## Cash Flows from Financing Activities

Cash proceeds from exercises of stock options increased from \$3.3 million for the three months ended March 31, 2017 to \$4.5 million for the three months ended March 31, 2018.

## Operating Capital Requirements

We expect to incur significant expenses and operating losses for the foreseeable future. Overall, we expect operating expenses to continue to increase over current levels as we incur increased costs related to: (i) our ongoing U.S. and international commercialization and pre-commercial activities including executing related marketing and promotional programs for the commercialization of ZEJULA; (ii) the advancement of clinical trial and other development and regulatory activities under our current development programs including niraparib, TSR-042, TSR-022 and TSR-033; (iii) activities under our research collaborations; (iv) expanding our international operations; and (v) other research and development activities and potential future collaborative or in-licensed development programs. If we obtain regulatory approval for any of our product candidates in addition to our current product approvals, or in anticipation of obtaining regulatory approval, we expect that we will incur significant additional commercialization expenses related to product sales, marketing, manufacturing and distribution. We also expect to incur increasing selling, general and administrative costs associated with our anticipated growth and continuing operation as a multinational public company, and we will continue to incur substantial interest expense related to our outstanding debt. The actual

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amount of many of these expenditures will depend on numerous factors, including the timing of expenses and the timing and progress of our clinical trial activity and commercialization efforts for our products and product candidates.

We may require additional capital for the continuing commercialization of our products, further development and potential commercialization of our product candidates, and cash interest obligations related to our outstanding debt. In addition, future license payments or milestone payments could cause our total operating expenses and cash usage to fluctuate. We may also need additional funds to pursue our strategy of in-licensing or acquiring additional product candidates and to meet our obligation to repay the principal of our term loan when due and to repay the Convertible Notes at maturity or, at our election, upon conversion. We are subject to the risks incident to the development of new biopharmaceutical products, and to global expansion, and we may encounter unforeseen expenses, difficulties, complications, delays and other unknown factors that may adversely affect our business and cause increased uses of cash.

We will need to generate significant revenues to achieve profitability, and we may never do so. Until we can generate a sufficient amount of revenue from our products, if ever, we expect to finance future cash needs through public or private equity or debt offerings and facilities, including an additional draw under our loan agreement, and we may seek additional capital through arrangements with strategic partners or from other sources. Additional capital may not be available to us on reasonable terms, if at all. If we are unable to raise additional needed capital in sufficient amounts or on terms acceptable to us, we would have to significantly delay, scale back or discontinue the development or commercialization of one or more of our product candidates and/or other areas of our business. Raising additional funds through the issuance of equity or debt securities could result in dilution to our existing stockholders, increased fixed payment obligations, or both. Furthermore, these securities may have rights senior to those of our common stock and could contain covenants that would restrict our operations and potentially impair our competitiveness, such as limitations on our ability to incur additional debt, limitations on our ability to acquire, sell or license intellectual property rights and other operating restrictions that could adversely impact our ability to conduct our business. Any of these events could significantly harm our business, financial condition and prospects.

We believe our currently available funds, including an additional draw under our loan agreement, and the cash we expect to generate from product sales will be sufficient to fund our existing cash flow requirements and our operations at their currently planned levels through at least the 12 months following the filing of this Quarterly Report on Form 10-Q. Our forecast of the period of time through which our financial resources will be adequate to support our operations is a forward-looking statement and involves risks and uncertainties, and actual results could vary as a result of a number of factors. We have based this estimate on assumptions that may prove to be wrong, and we could utilize our available capital resources sooner than we currently expect. Our future funding requirements, both short and long-term, will depend on many factors, including:

- our ability to generate revenues from sales of ZEJULA and, if approved, our other product candidates;

- the cost of continuing to expand our development and commercial capabilities for our products and our product candidates, both in the U.S. and in certain foreign markets, including Europe;
- the outcome, timing and cost of regulatory approvals by the FDA and comparable non-U.S. regulatory authorities and the potential that the FDA or comparable non-U.S. regulatory authorities may require that we perform more studies than those that we currently expect;
- the initiation, progress, timing, costs and results of clinical trials for our current product candidates and any future product candidates we may in-license;
- the cost and timing of completion of commercial-scale outsourced manufacturing activities for niraparib and our immuno-oncology antibody product candidates;
- the cost and timing of clinical development activities for niraparib;
- the cost and timing of preclinical and clinical development and manufacturing activities associated with our immuno-oncology antibody product candidates;
- the cost of filing, prosecuting, defending and enforcing any patent claims and other intellectual property rights that we own or control;
- the cost of acquiring or in-licensing global rights for additional product candidates;

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- the amount and timing of potential conversion requests, if any, and interest expense associated with our Convertible Notes; and
- our need to repay amounts due under our loan agreement.

If we lack sufficient capital to expand our operations or otherwise capitalize on our business opportunities, our business, financial condition and results of operations could be materially adversely affected.

## Contractual Obligations and Commitments

There have been no material changes to our contractual obligations and commitments included in our Annual Report on Form 10-K for the year ended December 31, 2017.

## Off-Balance Sheet Arrangements

As of March 31, 2018, we did not have any off-balance sheet arrangements as defined in Item 303(a)(4)(ii) of Regulation S-K.

## Critical Accounting Policies

Our management's discussion and analysis of our financial condition and results of operations are based on our financial statements, which have been prepared in accordance with U.S. generally accepted accounting principles. The preparation of these financial statements requires us to make estimates and judgments that affect the reported amounts of assets, liabilities, revenues and expenses and the disclosure of contingent assets and liabilities in our financial statements. On an ongoing basis, we evaluate our estimates and judgments, including those related to accrued research and development expenses, net product revenue, stock-based compensation expense and intangible assets. We base our estimates on historical experience, known trends and events and various other factors that are believed to be reasonable under the circumstances, the results of which form the basis for making judgments about the carrying values of assets and liabilities that are not readily apparent from other sources. Actual results may differ from these estimates under different assumptions or conditions. In making estimates and judgments, management employs critical accounting policies.

For a description of our other critical accounting policies, please see “Management’s Discussion and Analysis of Financial Condition and Results of Operations” included in our Annual Report on Form 10-K for the year ended December 31, 2017. There have not been any material changes to our critical accounting policies since December 31, 2017.

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Item 3. Quantitative and Qualitative Disclosures About Market Risk.

There have been no material changes with respect to the information appearing in Part II, Item 7A, “Quantitative and Qualitative Disclosures About Market Risk,” in our Annual Report on Form 10-K for the year ended December 31, 2017.

Item 4. Controls and Procedures.

Management’s Evaluation of our Disclosure Controls and Procedures

Our principal executive officer and our principal financial officer, after evaluating the effectiveness of our “disclosure controls and procedures” (as defined in the Securities Exchange Act of 1934, as amended, or the Exchange Act, Rule 13a-15(e) or Rule 15d-15(e)), with the participation of our management, has concluded that, as of the end of the period covered by this Quarterly Report on Form 10-Q, our disclosure controls and procedures are effective and are designed to ensure that information we are required to disclose in the reports that we file or submit under the Exchange Act is accumulated and communicated to management, including our principal executive officer and principal financial officer, as appropriate, to allow timely decisions regarding required disclosure, and is recorded, processed, summarized and reported within the time periods specified in the SEC’s rules and forms. It should be noted that any system of controls is designed to provide reasonable, but not absolute, assurances that the system will achieve its stated goals under all reasonably foreseeable circumstances. Our principal executive officer and principal financial officer have concluded that our disclosure controls and procedures are effective at a level that provides such reasonable assurances.

Changes in Internal Control Over Financial Reporting

There were no changes in our internal control over financial reporting that occurred during the fiscal quarter covered by this report that materially affected, or are reasonably likely to materially affect, our internal control over financial reporting.



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PART II OTHER INFORMATION

Item 1. Legal Proceedings.

A putative class action complaint was filed on January 17, 2018 in the United States District Court for the District of Massachusetts, entitled Roger Bowers v. TESARO Incorporated (sic), et. al., Case No. 18-10086. The complaint alleges that we and our Chief Executive Officer and our Chief Financial Officer violated certain federal securities laws, specifically under Sections 10(b) and 20(a) of the Securities Exchange Act of 1934, as amended, and Rule 10b-5 thereunder. The plaintiff seeks unspecified damages on behalf of a purported class of purchasers of our common stock between March 14, 2016 and January 12, 2018. On March 19, 2018, six separate applicants filed motions seeking appointment as lead plaintiff. Four of these applicants subsequently withdrew their motions or indicated that they did not oppose a competing motion filed by another applicant. The Court has not yet ruled on the remaining motions or selected a lead plaintiff. The Court has also not set a trial date for this matter. We believe that the allegations contained in the complaint are without merit and intend to defend the case vigorously.

Item 1A. Risk Factors.

An investment in our stock involves a high degree of risk. You should carefully consider the following discussion of risk factors, in its entirety, in addition to the other information contained in this Quarterly Report on Form 10-Q, our Annual Report on Form 10-K for the year ended December 31, 2017 and the other filings we make with the SEC. We cannot assure you that any of the events discussed in the risk factors below or in our other filings will not occur. These risks, or other events that we do not currently anticipate or that we currently deem immaterial, may have a material adverse effect on our business, prospects, financial condition and results of operations.

Risks Related to Our Business and Industry

We face substantial competition for our marketed products, ZEJULA and VARUBI/VARUBY, which could limit our ability to generate significant product sales.

The development and commercialization of new drug products is highly competitive. Many of our competitors are more established companies and may therefore have competitive advantages due to their size, cash flows, and institutional experience. We may be unable to compete successfully against these more established companies.

There are a number of large pharmaceutical and biotechnology companies that market and sell products or are pursuing the development of products that compete or we expect will compete with ZEJULA. There are currently two commercially available PARP inhibitors other than ZEJULA. AstraZeneca Plc's LYNPARZATM (olaparib) was initially approved by the FDA for use by ovarian cancer patients with a germline BRCA mutation, and was granted a new approval in August 2017 by the FDA for use as a maintenance treatment for recurrent, epithelial ovarian, fallopian tube or primary peritoneal adult cancer who are in response to platinum-based chemotherapy, regardless of BRCA status. In February 2018, LYNPARZA received a positive opinion from the Committee for Medicinal Products for Human Use, or the CHMP, of the European Medicines Agency, or EMA, for a new tablet formulation and a broad maintenance label, which is expected to lead to an EC marketing authorization in the near future. In addition, Clovis Oncology, Inc.'s RUBRACATM (rucaparib) was approved in December 2016 by the FDA for use as a monotherapy for the treatment of patients with deleterious BRCA mutation (germline and/or somatic) associated advanced ovarian cancer who have been treated with two or more chemotherapies. In April 2018, the FDA granted an expanded approval of RUBRACA for the maintenance treatment of adult women with recurrent ovarian cancer who have responded to their latest treatment with platinum chemotherapy, regardless of BRCA status, and in March 2018, the CHMP issued a positive opinion recommending the granting of a conditional approval for RUBRACA as a third line treatment of adult women with platinum sensitive, relapsed or progressive ovarian cancer with a BRCA-mutation and who are unable to tolerate further platinum chemotherapy. We believe there are also several additional products in clinical development targeting the PARP pathway, as detailed in Part I, Item 1, "Business – Competition" of our Annual Report on Form 10-K for the year ended December 31, 2017. Both LYNPARZA and rucaparib have received "orphan drug designation" from the EMA, which provides certain benefits including market exclusivity for up to ten years in the approved indication post-approval. In addition to other PARP inhibitors, ZEJULA also competes with AVASTINTM (bevacizumab), Roche's angiogenesis inhibitor. AVASTIN is FDA- and EMA-approved in combination with chemotherapy for the treatment of recurrent ovarian cancer following platinum-containing chemotherapy (platinum-sensitive and platinum-resistant) and has been approved by the EMA, in combination with chemotherapy, for front-line treatment. In the U.S., it is currently under FDA review for the front-line setting.

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We also face substantial competition with respect to oral and IV formulations of VARUBI and with respect to VARUBY. VARUBI/VARUBY competes with EMEND, an NK-1 receptor antagonist marketed by Merck, as well as AKYNZEO, an oral combination NK-1 receptor antagonist and 5-HT3 receptor antagonist (netupitant plus ALOXI (palonosetron HCl)) that is marketed by Helsinn Healthcare, and Sandoz's generic version of aprepitant. Additionally, Heron Therapeutics recently received FDA approval, and launched marketing of, its aprepitant IV formulation product, CINVANTITM. VARUBI/VARUBY would face additional competition if additional generics are introduced to the market, or other products are developed and approved, for the treatment and prevention of CINV, or if an IV formulation of AKYNZEO is developed.

We are aware of several companies that have antibody-based products on the market or in clinical development that are directed at the same biological targets as some of our immuno-oncology programs. There are currently two anti-PD-1 antibody products and three anti-PD-L1 antibody products being marketed. OPDIVO® (nivolumab, marketed by Bristol-Myers Squibb) is approved in a number of indications as a monotherapy or in combination with other products; KEYTRUDA® (pembrolizumab, marketed by Merck) is approved in a number of indications; TECENTRIQ® (atezolizumab, marketed by Roche) is approved in patients with various forms of locally advanced or metastatic urothelial carcinoma or metastatic NSCLC; IMFINZI® (durvalumab, marketed by AstraZeneca) was approved in 2017 for patients with various forms of locally advanced or metastatic urothelial carcinoma; and BAVENCIO® (avelumab, co-marketed by Merck KGa and Pfizer) was approved in 2017 for adults and pediatric patients 12 years and older with metastatic Merkel cell carcinoma and for patients with various forms of locally advanced or metastatic urothelial carcinoma. Although there are currently no anti-TIM-3 antibody products or anti-LAG-3 antibody products being marketed, we are aware that two companies, Novartis and Eli Lilly, have anti-TIM-3 modulator antibodies in Phase 1/2 clinical development for various indications. We are also aware of several companies that have anti-LAG-3 modulators in development for various indications, including Bristol-Myers Squibb, which has an anti-LAG-3 antibody in Phase 2 clinical development; Novartis, which has an anti-LAG-3 antibody in Phase 1/2 and Phase 2 clinical development; and Merck, Boehringer Ingelheim and Regeneron Pharmaceuticals, each of which has an anti-LAG-3 antibody in Phase 1 clinical development.

For further detail on the specific competition that VARUBI/VARUBY, ZEJULA and our immuno-oncology antibody product candidates face, see Part I, Item 1, "Business – Competition" of our Annual Report on Form 10-K for the year ended December 31, 2017.

Many of the approved drugs with which our products or product candidates may compete are well-established therapies or products and are widely accepted by physicians, patients and third-party payors. Insurers and other third-party payors may also encourage the use of generic products. Any of our product candidates that are approved may be priced at a significant premium over competitive generic products.

Our competitors may obtain regulatory approval of their products more rapidly than we are able to or may obtain patent protection or other intellectual property rights that limit our ability to develop or commercialize our product candidates. Our competitors may also develop drugs that are more effective, more widely used and less costly than

ours, and may also be more successful than us in manufacturing and marketing their products.

Significant changes in U.S. and international trade policies that restrict imports or increase tariffs could have a material adverse effect on our results of operations.

We depend on third-party manufacturers and suppliers outside of the United States, including China, in connection with the manufacture of certain of our products and product candidates. Accordingly, our business is subject to risks associated with international manufacturing. For example, the Trump Administration has called for substantial changes to U.S. foreign trade policy, including the possibility of imposing greater restrictions on international trade and significant increases in tariffs on goods imported into the United States, and has increased tariffs on certain goods imported into the United States from China. The institution of additional protectionist trade measures could adversely affect our manufacturing costs, and in turn our business, financial condition, operating results and cash flows.

#### Risks Related to Our Dependence on Third Parties

We have no manufacturing facilities, and we are dependent on a limited number of third-party manufacturers for the manufacture and supply of ZEJULA, VARUBI, VARUBY and our product candidates, including our immuno-

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oncology assets. If we experience problems with any of these third parties, the manufacturing of our products or our product candidates could be delayed, which could cause significant harm to our competitive position and our ability to generate and grow revenues from our approved products, to our ability to complete clinical trials and obtain regulatory approval for our product candidates, and to our future results of operations.

We do not own or operate facilities for the manufacture of our products or product candidates. Our ability to successfully develop and commercialize our products and our product candidates will require us to establish large scale manufacturing capabilities through our contract manufacturing organizations, or CMOs. We currently have no plans to build our own clinical or commercial scale manufacturing capabilities. We currently work with two CMOs for the production of ZEJULA drug substance, and one other CMO, for ZEJULA drug product supply, for our clinical and commercial needs. We currently work with one CMO for the production of rolapitant drug substance used for VARUBI and VARUBY, and one CMO for commercial production of VARUBI and VARUBY.

As we continue to commercialize ZEJULA and VARUBI/VARUBY and develop our other product candidates, including our immuno-oncology assets, we will have a greater need for both clinical study supply and commercial manufacturing capacity. We have limited experience manufacturing pharmaceutical products on a commercial scale, and some of our suppliers, including our supplier of ZEJULA drug product, and our suppliers of both drug substance and drug product for our immuno-oncology assets, have limited capacity and will need to continue to increase their scale of production to meet our projected needs for both clinical and commercial manufacturing. Manufacturing commercial quantities of our products, including ZEJULA, to meet our projections will require our third-party manufacturers to invest substantial additional funds to increase capacity and to hire and retain additional personnel who have applicable large-scale commercial manufacturing experience. Our third-party manufacturers may not successfully complete any required increase to existing manufacturing capacity in a timely manner, or at all. Because of the complex nature of our compounds, our manufacturers may not be able to manufacture our compounds at an acceptable cost or in sufficient quantities or in a timely manner necessary to make commercially successful products, or may require us to pay significant costs, including for capital improvements to their facilities.

If our contract manufacturers or other third parties fail to deliver our products, including ZEJULA, for commercial sale on a timely basis, with sufficient quality, and at commercially reasonable prices, and we fail to find replacement manufacturers or to develop our own manufacturing capabilities, we may be required to delay or suspend commercialization of our current products or other potential future products, which would likely cause significant harm to our competitive position in the marketplace and our ability to generate and grow revenues from our approved products, and to our future results of operations.

For development of our immuno-oncology antibody product candidates, we currently work with one CMO for the production of biologics. For each of our product candidates, we may elect to pursue arrangements with other CMOs for manufacturing clinical supplies for later-stage trials and for commercialization. We have not yet qualified alternate suppliers in the event the current CMOs we utilize are unable to scale production, or if we otherwise experience any problems with them. If we are unable to arrange for alternative third-party manufacturing sources, or to do so on commercially reasonable terms or in a timely manner, we may not be able to complete development of our product candidates, or market or distribute them.

Reliance on third-party manufacturers entails risks to which we would not be subject if we manufactured product candidates or products ourselves, including reliance on the third party for regulatory compliance and quality assurance, the possibility of breach of the manufacturing agreement by the third party because of factors beyond our control (including a failure to synthesize and manufacture our product candidates or any products we may eventually commercialize in accordance with our specifications) and the possibility of termination or nonrenewal of the agreement by the third party, based on its own business priorities, at a time that is costly or damaging to us. In addition, the FDA and similar foreign authorities require that our product candidates and approved products be manufactured according to cGMP and similar foreign standards. Any failure by our third-party manufacturers to comply with cGMP or failure to scale up manufacturing processes, including any failure to deliver sufficient quantities of product candidates in a timely manner, could lead to a delay in, or failure to obtain, regulatory approval of any of our product candidates. In addition, such failure could be the basis for the FDA or an equivalent foreign regulatory authority to issue a warning or untitled letter, withdraw approvals previously granted to us for our products or product candidates, or take other regulatory or legal action, including recall or seizure, total or partial suspension of production, suspension of ongoing clinical trials, refusal to approve pending applications or supplemental applications, detention of product, refusal to permit the import or export of products, injunction, or imposition of civil and criminal penalties.

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Any significant disruption in our supplier relationships could severely harm our business. We source key materials from third parties, either directly through agreements with suppliers or indirectly through our manufacturers who have agreements with suppliers. For example, we source key raw materials for ZEJULA drug substance from one supplier. Such suppliers may not sell these key materials to us or our manufacturers at the times we need them or on commercially reasonable terms. In certain cases, we do not have any control over the process or timing of the acquisition of these key materials by our manufacturers. Any significant delay in the supply of a product or product candidate or its key materials for an ongoing clinical study could considerably delay completion of our clinical studies, product testing and potential regulatory approval of our product candidates. If our manufacturers or we are unable to purchase these key materials for products or product candidates after regulatory approval, the commercial launch of our product candidates could be delayed or there could be a shortage in supply, which would impair our ability to generate revenues from the sale of our products and product candidates.

Because we have relied and plan to continue to rely on third parties for the foregoing preclinical and clinical functions, our internal capacity to perform these functions is limited. Switching or adding additional CMOs involves additional cost, requires management time and focus, and could result in substantial delays in our development programs. Identifying, qualifying and managing the performance of third-party service providers can be difficult, time consuming and cause delays in our development programs. In addition, there is a natural transition period when a new CMO commences work and the new CMO may not provide the same type or level of services as the original provider. If any of our relationships with our third-party CMOs terminates, we may not be able to enter into arrangements with alternative CMOs or do so on commercially reasonable terms.

## Item 6. Exhibits.

The exhibits listed below are filed or furnished as part of this Quarterly Report on Form 10-Q.

## EXHIBIT INDEX

| Exhibit<br>Number | Exhibit Description                                                                                                                          |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------|
| 10.1*+            | <u>Seventh Amendment to 1000 Winter Street Lease Agreement, dated October 10, 2017, by and between the Company and BP Bay Colony LLC.</u>    |
| 31.1              | <u>Certification of principal executive officer pursuant to Rule 13a-14(a)/15d-14(a) of the Securities Exchange Act of 1934, as amended.</u> |
| 31.2              | <u>Certification of principal financial officer pursuant to Rule 13a-14(a)/15d-14(a) of the Securities Exchange Act of 1934, as amended.</u> |
| 32.1              |                                                                                                                                              |

32.2

Certification of principal executive officer pursuant to 18 U.S.C. §1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002.

Certification of principal financial officer pursuant to 18 U.S.C. §1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002.

|            |                                                        |
|------------|--------------------------------------------------------|
| EX-101.INS | XBRL Instance Document                                 |
| EX-101.SCH | XBRL Taxonomy Extension Schema Document                |
| EX-101.CAL | XBRL Taxonomy Extension Calculation Linkbase Document  |
| EX-101.DEF | XBRL Taxonomy Extension Definition Linkbase Document   |
| EX-101.LAB | XBRL Taxonomy Extension Label Linkbase Document        |
| EX-101.PRE | XBRL Taxonomy Extension Presentation Linkbase Document |

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\* Confidential Treatment has been requested with respect to certain portions of this exhibit. Omitted portions have been filed separately with the U.S. Securities and Exchange Commission.

+This exhibit is a corrected version of the exhibit that was previously filed as Exhibit 10.46 to the Company's Annual Report on Form 10-K for the year ended December 31, 2017.

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SIGNATURES

Pursuant to the requirements of Section 13 or 15(d) 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.

TESARO, INC.

By: /s/ Leon O. Moulder, Jr.  
Leon O. Moulder, Jr.  
Chief Executive Officer  
(principal executive officer)

Date: May 3, 2018

By: /s/ Timothy R. Pearson  
Timothy R. Pearson  
Executive Vice President and Chief Financial Officer  
(principal financial officer)

Date: May 3, 2018

